Four manuals on implementing the Partners Respite Model, which provides respite care for children with disabilities or chronic illnesses, comprise this document. The Community Planning Manual offers a step-by-step guide to replication of the Partners Respite Model and is divided into sections on developing the Partners program, implementing the Partners program, measuring program success, and offering resources. The Trainer's Workshop Manual is intended to be used for a 6-hour group training in implementing the model's 5 steps: (1) recruit families and caregivers; (2) conduct partners workshop; (3) help families choose caregivers; (4) guide families as they train their own caregivers; and (5) provide continuing support. The Family Manual guides families in choosing and training their own respite caregivers by using their children's daily routines to develop training plans. It also provides worksheets, resources, and forms. Finally, the Caregiver Manual offers basic information on caring for young children with disabilities and supporting families. It includes information about respite, communication, building partnerships, disabilities awareness, child development, children with special needs, and personalized training with families. (DB)
Partners Plus

Families and Caregivers in Partnerships

A Family-Centered Guide to Respite Care
Child Development Resources

BEST COPY AVAILABLE

DRAFT 10.99
Partners Plus

Families and Caregivers in Partnerships: A Family-Centered Guide to Respite Care

By
Lisa L. Ownby, M.S.W.
Amanda C. Hooke, M.S.W.
Dee Wylie Moore, B.S.
Corinne W. Garland, M.Ed.
Acknowledgments

The authors would like to thank Jayne Naughton for her technical assistance; Amy Watson for editorial assistance; families and caregivers who wrote the perspectives stories in section four of the Family and Caregiver Manuals; and our community advisory committee for multiple tasks throughout the development of training materials. Their suggestions and comments provided a significant contribution to the quality of this manual.

Amy Consiglio, of CM Consulting, designed and produced this manual.

A Word About Funding

Support for the Partners Plus project was provided under grant number H024B40032 from the Early Education Program for Children with Disabilities, U.S. Department of Education. Points of view or opinions expressed herein do not, however, necessarily represent official views or opinions of the Department of Education.

Funding to revise the Partners Plus manuals to help families use existing community programs as respite for their school-age children with mental retardation came from the Joseph P. Kennedy, Jr. Foundation. Revisions to the manuals include practical “how to” information for personnel working with children in before and after school programs, recreational, camp, and other community programs.

Copyright © 1999. All rights reserved. Trainers may reproduce this manual, without written permission, and in whole for use only with families and caregivers. For permission to reprint single pages or to purchase additional copies of the manual, please contact:

Child Development Resources
Post Office Box 280
Norge, VA 23127-0280
Phone (757) 566-3300 / Fax (757) 566-8977
Contents

• Community Manual
• Trainer's Workshop Manual
• Family Manual
• Caregiver Manual
Partners Plus

Foreward

A Guide to Family-Centered Respite
Using this Guide

Format of this Guide

The Partners Plus Guide is a comprehensive resource that includes practical easy-to-use information to help communities establish a family-centered respite program. The Guide contains four individual manuals developed by Partners Plus, a program of Child Development Resources (CDR). Partners staff designed each manual for specific audiences. Partners designed the Community Manual to help community planning groups develop a respite program. The Trainer's Workshop Manual gives trainers the information they need to conduct workshops for families and caregivers. We designed the Family Manual and Caregiver Manual to meet the training needs of families and caregivers as they work together in respite partnerships.

Communities, families, caregivers, and trainers will find each manual useful as an independent resource or as a complete package for carrying out a community respite program. CDR Partners staff offer support and technical assistance to communities interested in replicating the Partners respite program. Please contact CDR to learn how we can help you establish a respite program in your community.

Child Development Resources
Phone: 747-220-1168
Fax: 757-253-1779
E-Mail: ppcdr@tni.net
Mail: 1490 Government Road
Williamsburg, VA 23185
The Community Planning Manual

The Community Planning Manual, the first part of this Guide, describes the process for successful replication of the Partners model of family-centered respite care. Partners Plus and several community planning groups in Virginia used this manual as a resource while working toward replication of the Partners model. Coordinating leadership in an initial community planning group is the beginning process of replication that leads to conducting the Partners' workshop for families and caregivers. Continual support to families and caregivers follows a workshop as families and their caregivers develop long-term respite partnerships. In following the replication process, community leaders can work collaboratively with respite stakeholders to develop a respite care program. Partners divided the Community Planning Manual is divided into four sections:

- Developing your Partners program
- Implementing your Partners program
- Measuring your success
- Offering resources

When community planning groups are working together toward establishing their own Partners program they can easily pull the Community Planning Manual section out of the Guide to make copies of pages appropriate to group members' tasks.
Using this Guide

The Trainer's Workshop Manual

Staff of the Partners program designed the Trainer's Workshop Manual to help Partners trainers in planning and conducting the six-hour workshop for families and caregivers. Pages of the manual have holes to fit a three-ring binder so that trainers can add or remove contents to fit their presentation style. Some sections of the Trainer's Workshop Manual include the use of overhead projector transparencies. We provide master copies for making these transparencies at the end of each corresponding section. Partners trainers may choose to add some or all of the other Guide components to their personal Trainer's Workshop Manual.

The Family Manual

Staff of the Partners program designed the Family Manual for families to use in choosing and training their own respite caregivers. Communities may make additional copies for families or order them directly from CDR. At the beginning of a Partners workshop, trainers give each family a manual. During the workshop families learn to use the manual as a resource for finding and training their own caregivers. This manual guides families in using their children's daily routines to develop personalized training plans for respite caregivers. It includes information about respite, communication, building partnerships, selecting caregivers, and developing personalized training. The Family Manual contains worksheets, resources, and forms that can be duplicated for future use.
The Caregiver Manual

Staff of the Partners program designed the Caregiver Manual to give caregivers basic information about caring for young children with disabilities and supporting families. Again, communities may reprint the whole manual or purchase them from CDR. At Partners workshops, trainers give caregivers a manual to use during the workshop that they can take with them afterwards for use as a resource when partnering with a family. This manual includes information about respite, communication, building partnerships, disabilities awareness, child development, children with special needs, and personalized caregiver training.
Partners Plus
Families and Caregivers in Partnerships:
Trainer's Workshop Manual

A Guide to Conducting Partners Plus Workshops
Child Development Resources

DRAFT 10.99
Partners Plus

Trainer's Workshop Manual

Developed By
Amanda C. Hooke, M.S.W.
Lisa L. Ownby, M.S.W.
Dee Wylie Moore, B.S.
Corinne W. Garland, M.Ed.
Adrienne Frank, O.T.R.

Acknowledgments

The authors would like to thank Jayne Naughton for her technical assistance; Amy Watson for editorial assistance; families and caregivers who wrote the perspectives stories in section four of the Family and Caregiver Manuals; and our community advisory committee for multiple tasks throughout the development of training materials. Their suggestions and comments provided a significant contribution to the quality of this manual.

Amy Consiglio, of CM Consulting, designed and produced this manual.

A Word About Funding

Support for the Partners Plus project was provided under grant number HO24B40032 from the Early Education Program for Children with Disabilities, U.S. Department of Education. Points of view or opinions expressed herein do not, however, necessarily represent official views or opinions of the Department of Education.

Funding to revise the Partners Plus manuals to help families use existing community programs as respite for their school-age children with mental retardation came from the Joseph P. Kennedy, Jr. Foundation. Revisions to the manuals include practical “how to” information for personnel working with children in before and after school programs, recreational, camp, and other community programs.

Copyright © 1999. All rights reserved. Trainers may reproduce this manual, without written permission, and in whole for use only with families and caregivers. For permission to reprint single pages or to purchase additional copies of the manual, please contact:

Child Development Resources
Post Office Box 280
Norge, VA 23127-0280
Phone (757) 566-3300 / Fax (757) 566-8977
Contents

The Partners Model of Respite Care iii
Partners Plus Replication Process vii
Partners Plus Manuals ix
About the Partners Workshop x
Partners Workshop Objectives xii
Use of the Trainer's Workshop Manual xiv
Preparation for Training xvi
Workshop Introduction—Welcome to the Partners Workshop Intro–1
Section 1: About Respite 1-1
Section 2: Building Partnerships: Communication 2-1
Section 3: Building Partnerships: Working Together 3-1
Section 4: All Kids Like Cookies 4-1
Section 5–F (Families): Finding Respite Caregivers 5 F-1
Section 6–F (Families): Interviewing Caregivers 6 F-1
Section 7–F (Families): Developing Your Personalized Training Plan 7 F-1
Section 5–C (Caregivers): Disabilities Awareness 5 C-1
Section 6–C (Caregivers): Understanding Child Development 6 C-1
Section 7–C (Caregivers): Children with Special Needs 7 C-1
Section 8–C (Caregivers): Becoming a Caregiver 8 C-1
Final Partners Workshop Activity
Resources
Trainer's Workshop Manual

Introduction
The Partners Model of Respite Care...

*Helps families of children with special needs find and train their own respite caregivers.*

*Through training and support, families and caregivers build long-lasting, successful respite relationships.*

**Introduction**

Like all families, families of children with special needs want and need time away from the demands of parenting. Respite—or temporary child care for children with disabilities or chronic illness—gives families time to enjoy an evening out by themselves, with other family members, or with friends. However, for these families, finding respite care is not as simple as calling a neighborhood babysitter. These families need caregivers who understand their children's special needs.

When the staff at Child Development Resources (CDR) spoke with families about what they wanted in a respite program, the message was clear. Families wanted to choose and train their own respite caregivers. They wanted affordable care that was as simple to schedule as calling the neighborhood babysitter. And, families wanted caregivers who were trained to provide care.

In response to families and with support from the Virginia Institute for Developmental Disabilities (VIDD), a University Affiliated Program at Virginia Commonwealth University in Richmond, Virginia, CDR developed a pilot project that was low-cost, family-directed respite care called PARTners: Parents As Respite Trainers.

CDR was awarded, in 1994, a three-year grant from the U.S. Department of Education, Office of Special Education Programs (OSEP), Early Education Program for Children with Disabilities (EEPCD), to build on the pilot and to develop a model of respite that could be replicated in other communities. The project, **Partners Plus: Families and Caregivers in Partnerships**, provides a model for families to use in finding and training respite caregivers. This model encourages families to use available, natural family and community supports to meet their respite needs.
Within this model, Partners broadly defines the terms respite and caregiver. For example, respite can take place in a family's home, a caregiver's home, or in before and after school, recreation, camp, or other children's program. A caregiver can be any individual who provides respite care to a child with special needs or includes a child with special needs within an existing community program.

Partners Plus provided training to more than 200 families and caregivers in a five-city, three-county area of eastern Virginia. Partners usually conducted the six-hour Partners Plus workshop for families and caregivers in one day. Sometimes when it was more convenient for a group, Partners conducted the workshop in two parts on separate days. This flexibility ensured that Partners adapted training to fit the schedules of families and caregivers who are generally most receptive to training during nontraditional hours such as evenings and Saturdays. Families attending the workshop learned to provide their own caregivers with personalized training based on their children's daily routines. Families either came to training with their own caregivers, met caregivers through the workshop, or through other project activities.

The Partners Project resulted in the development of four manuals designed to help communities, trainers, families, and caregivers use the Partners model. The Trainer's Workshop Manual is one of four in a series of manuals that Partners has conveniently packaged together in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care. Besides the set of manuals contained in the guide, you can purchase the Family and Caregiver Manuals separately from the Partners Plus program through CDR.

Child Development Resources
Post Office Box 280
Norge, VA 23127-0280
Phone (757) 566-3300
Fax (757) 566-8977
Email ppcdr@tni.net
The Partners Model: A New Model of Respite Care

Partners programs provide group training and continuing support to families and potential caregivers. The Partners model contains five steps:

The Five Steps of the Partners Model

1. Recruit Families and Caregivers
2. Conduct Partners Workshop
3. Help Families Choose Caregivers
4. Guide Families as they Train their Own Caregivers
5. Provide Continuing Support

Step 1: Recruit Families

First, programs recruit families and potential caregivers to participate in Partners. Community awareness activities invite potential participants to a Partners workshop. Information about the workshop and the Partners program is distributed through parent groups, newspapers, radio, flyers, word of mouth, etc.

Step 2: Conduct Partners Workshop

Next, programs conduct the Partners workshop. During the six-hour workshop, families learn how to find, screen, and interview caregivers for their children. In addition, they learn to structure training for their own caregivers based on the specific needs and daily routines of their individual children. Caregivers learn basic information about caring for children with special needs. During the workshop, caregivers are preparing to be active in their work with families, reflecting on their motivations for becoming respite caregivers and identifying their own learning needs for personalized training. Partners Family and Caregiver Manuals
provide participants with useful resources to actively engage in during the workshop and to use as resources as they move through the next steps of the model.

**Step 3: Families Choose Caregivers**

After the workshop, Partners programs provide continuing support as families choose caregivers. Families frequently identify respite caregivers within their own circles of natural support. If families are unable to identify their own caregivers, Partners will assist by providing social and educational events that will help families and caregivers meet. During interviews, families and caregivers negotiate how respite occurs. Respite may be in-home or out-of-home and reimbursement for care can be flexible. Care may be provided for a fee, voluntarily, or in exchange for another service, such as yard work.

**Step 4: Families Train Caregivers**

Once families and caregivers decide to work together, families train their own caregivers. This “personalized training,” discussed during the workshop, is organized by families with help from Partners staff if needed. Families decide what routines caregivers need to learn and work with caregivers to determine how training will occur, thus forming true partnerships necessary for successful respite.

**Step 5: Provide Continuing Support**

As families and caregivers work together to develop a personalized respite training, the community Partners program offers continuing support, information and guidance. The Partners program is available to support families and caregivers in various ways as they begin their respite relationship. Partners staff members respond to requests for help and support from both families and caregivers. For example, Partners may help families and caregivers with screening, interviewing, developing personalized training plans, and providing educational opportunities.
Partners Plus Replication Process

The Partners Plus Community Planning Manual, contained in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care, helps communities use the Partners model of respite care. Each community interested in model replication develops a plan for collaboration among agencies, organizations, and individuals in order to implement the model. Community resources and supports are needed to sustain the model, to create a home for the program, and to ensure continuation.

The community replication process includes four major steps:

1. Identifying the site for replication
2. Planning for replication
3. Implementing the model
4. Evaluating the model

Each community needs to develop a plan for replication. Awareness activities aimed at a broad array of agencies, organizations, and individuals help strengthen community support. The creation of an advisory committee brings expertise and resources that will ensure a quality program. The replication...
Partners Community Implementation Process

plan includes finding a home for the program and securing staff responsible for implementing the program. Formal or informal interagency agreements and perhaps grant proposals may be necessary for community replication.

The Community Planning Manual provides strategies and materials to help communities develop and implement the respite care program. The Trainer's Workshop Manual, accompanying the Community Planning Manual, includes agendas, overheads, and suggestions for what to say during the Partners workshop. There are also suggestions for the kinds of resources that are needed by families and caregivers in order to foster and maintain long-term relationships.

Communities that replicate the Partners model gather information from families and caregivers to evaluate the quality and usefulness of training and support.

Partners in your Community

Plan for Replication of the Partners Model
- Create Community Support
- Determine Program Advisors
- Determine Administrative Structure
- Secure Financial Resources

Implement the Partners Model
- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families As They Train Their Own Caregivers
- Provide Continuing Support

Evaluate the Partners Program
Partners Plus Manuals

There are four manuals in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care that support the Partners Plus replication process:

1. The Family Manual
2. The Caregiver Manual
3. The Community Planning Manual, and

The Family and Caregiver Manuals are used in the Partners workshop for families and caregivers. These manuals include resources that families and caregivers can continue to use as they develop partnerships for respite care for children with special needs. The Community Planning Manual helps communities develop a community-based, family-centered respite system. The Trainer's Workshop Manual contains overheads, forms, and "helpful hints" for conducting Partners workshops.

The Family Manual is easy to follow and guides families in choosing and training their own respite caregivers. Families use their children's daily routines to develop training plans for caregivers. The manual includes information about respite, communication, building partnerships, selecting caregivers, and developing personalized training. There are worksheets, resources, and forms that can be duplicated for future use.

The Caregiver Manual provides caregivers with basic information on caring for young children with disabilities and supporting families. This manual includes information about respite, communication, building partnerships, disabilities awareness, child development, children with special needs, and personalized training with families.

The Community Planning Manual provides communities with a step-by-step guide to replication of the Partners model. By following each step, communities and families can work together to develop respite care programs. The manual contains forms and materials to assist with replication of the model.

The Trainer's Workshop Manual, used in conjunction with the Community Planning Manual, helps community trainers to conduct the six-hour Partners workshop for families and caregivers.
About the Partners Workshop

The six-hour Partners workshop helps families and caregivers gain knowledge and skills to build respite partnerships. The workshop can take place in one day-long session or two half-day sessions. Flexibility in scheduling workshops makes it easier for families and caregivers to attend the workshop on Saturdays or evenings.

In the workshop, families and caregivers participate in lectures, discussion and practice activities designed to meet the needs of adult learners. Activities and materials engage and inform participants in ways that support different styles of learning.

Partners divide the workshop into sections. Following this page is a flow chart to help trainers recognize how Partners organized information in this manual. To begin the workshop for participants, the trainer shares information from the Welcome to the Partners Workshop section. Families and caregivers stay together through the first few sections. Later, they split into family and caregiver groups for “Break-Out Sessions” designed to meet their different needs. Finally, families and caregivers reunite for a practice activity employing the skills developed through the workshop.
Flow Chart to the Trainer’s Workshop Manual

Introduction
The first section of the Trainer’s Workshop, Family, and Caregiver Manuals have the same Introduction.

How To Use the Trainer’s Workshop Manual
The second section of the Trainer’s Workshop Manual tells the trainer how to use the manual.

Welcome to the Partners Workshop
Workshop information for all participants begins in the Trainer’s Workshop Manual at this section.

Sections 1-4: Participants Work Together
All workshop participants work together during sections 1, 2, 3, and 4, where the information is the same for both families and caregivers:

Section 1: About Respite
Section 2: Building Partnerships: Communication
Section 3: Building Partnerships: Families and Caregivers Working Together
Section 4: All Kids Like Cookies Activity

Sections 5-10: Participants Work Separately
After completing section 4, the group splits into families and caregivers, each working with a trainer on topics that are different:

Family Break-Out Session
Section 5-F: Finding Respite Caregivers
Section 6-F: Interviewing Caregivers
Section 7-F: Developing a Personalized Training Plan
Resources
Forms for Sharing Information

Caregiver Break-Out Session
Section 5-C: Disabilities Awareness
Section 6-C: Understanding Child Development
Section 7-C: Children with Special Needs
Section 8-C: Becoming a Caregiver
Resources
Forms for Sharing Information

Final Activity: Participants Work Together
After completing their sections, the two groups return for the Final Activity, involving daily routines to help develop a personalized training plan.
Partners Workshop Objectives

As a result of this workshop, families and caregivers will:

- Be able to define respite care and identify the benefits to families, children and caregivers,
- Know how families drive family-centered services, and what rights and responsibilities they have as participants in family-centered systems,
- Understand the Partners model of respite care,
- Know the importance of clear and open communication in a respite relationship,
- Know the variety of ways that people send and receive messages,
- Be able to recognize and mend communication breakdowns,
- Practice communication skills,
- Have firsthand experience of some of the challenges faced by children with special needs,
- Be able to identify the ways in which children with special needs adapt to their environments, and
- Practice developing a personalized training plan using a child's daily routines.

As a result of this workshop, families will:

- Become more aware of caregivers' perspectives on caring for children with special needs,
- Understand why many caregivers experience “burn out” and stop providing respite care,
- Know strategies to prevent caregiver burn out,
- List qualities/criteria for selecting caregivers,
- Know other strategies for finding respite caregivers,
- Be able to identify potential caregivers in their natural support networks,
- Know what questions to ask caregivers in interviews,
- Understand different options for conducting interviews,
- Know other strategies for gathering information about caregivers,
- Understand the four-step process for teaching daily routines to caregivers,
- Know some principles for teaching adults, and
- Understand the importance of sharing emergency procedures and household rules with caregivers.
As a result of this workshop, caregivers will:

- Become more aware of the perspectives of families with children with special needs,
- Recognize families' need for respite and the value of the service that respite caregivers provide,
- Understand the importance of learning about a family's experience from that family,
- Understand common terms, services and issues related to children with special needs,
- Understand the basics of child development,
- Be aware of the importance of play and enriching environments for child development,
- Be aware of potential hazards in an environment and appropriate safety precautions,
- Be aware of the basic developmental and medical effects of common special needs on children.

- Be aware of what they may be asked to do when caring for children with common special needs,
- Know how to match with families for respite, and
- Understand how caregivers can contribute to personalized training.
Use of the Trainer’s Workshop Manual

The Trainer’s Workshop Manual gives trainers information on respite and helping families and caregivers develop skills to build strong respite partnerships. Experienced trainers enhance the workshop, however, Partners designed the manuals to help even inexperienced trainers to conduct successful workshops. It is especially beneficial when trainers can bring their personal or professional perspectives in disabilities to the workshop. While comfort and skills in teaching adults are also helpful, the Trainer’s Workshop Manual helps trainers understand how to facilitate training and make it effective for participants with different learning styles.

Partners divides the workshop into Sections. The Trainer’s Workshop Manual outlines training activities and includes overheads and supplemental materials for use in the workshop. Each Section begins with a list of the learning objectives, time and materials needed, and specific instructions that apply to the section content. Workshop notes to the trainer are on the left of each page. On the right of each page is a list of overheads or materials needed. In the center of each page in a bold font are suggested ways to present essential information using specific words or phrases. However, trainers are encouraged to use their own words and illustrations rather than reading the phrases from the manual.

In developing the Trainer’s Workshop Manual, Partners took great care to avoid language that creates or reinforces negative stereotypes of people with special needs. When paraphrasing these notes or making other comments, trainers should use language that conveys positive and respectful images of people with special needs. Trainers are encouraged to always choose words that emphasize the person, not his or her condition, disability, or special health needs.

The Trainer’s Workshop Manual includes master copies of overheads and supplemental materials in the back of each section. Trainers can easily copy overheads onto transparencies. An effective training method used in several years of field testing is the Suggested Overheads. To help trainers use the overheads appropriately, Partners coded them. On the bottom right of every page is a code beginning with PP. Following the PP is a number that identifies the section where the trainer uses the overhead. The next number is the sequential number of the overheads in that section. Thus, the code “PP 2-3” is the third overhead of the second section covered in the workshop.
Partners provide instructions on the use of the *Supplemental Materials* in each section. Trainers make copies of these materials for each participant in the workshop. Again, at the bottom of each page of the *Supplemental Materials* is a code to help trainers use these materials appropriately. The code begins with PP and follows with SM, meaning *Supplemental Materials*. The next letter or number is the section where the trainers use the material. The final number is the sequential number of *Supplemental Materials* in that section.
Preparation for Training

Preparation is key to an effective Partners workshop. Trainers should become acquainted with the Partners' workshop materials, including this manual and the Family and Caregiver Manuals. An accessible training facility with adequate size rooms, tables, and chairs for participant activities is necessary for a successful workshop.

If your Partners' workshop includes child care for families, then trainers must plan for this. Child care spaces should be in the same facility as the Partners' workshop for adults. Caring for children in a range of ages, with varying developmental needs, will require appropriate spaces, toys and supplies.

Trainers familiar with their communities' Partners program will help ensure smooth linkages for families, caregivers, and community contacts for support to those participating in the workshops.

Trainers should gather all materials and resources, including handouts and overheads, before the workshop. Familiarity with the Trainers' Workshop Manual and the Family and Caregiver Manuals will help trainers learn the content and flow of the workshop, and will prepare trainers to help participants with questions and concerns.

Workshop materials, including overheads handouts, and training manuals should be prepared before the workshop. Partners included master copies of overheads and handouts in this manual.


Participants need an agenda that guides them through the workshop and increases their comfort by letting them know what to expect. At the end of the Welcome Section, among the Supplemental Materials are a one day and two-day sample agenda. Agendas should be prepared before the workshop.

Partners include sample evaluation measures as Supplemental Materials at the end of the Welcome Section. Trainers working with Partners staff, for replication of the Partners model use these evaluations. Instruments specific to the workshop should be prepared before training. Data from these measures give trainers information on how well participants reached the workshop objectives.

Partners developed a Workshop Checklist to help trainers prepare for workshops. The checklist is with the Supplemental Materials at the end of the Welcome Section.
About Overheads

The overheads contained in the Trainer's Workshop Manual were developed using proven methods to promote visual learning. Trainers may copy these overhead pages onto transparencies and may develop their own overheads for use in the workshop. Cartoons or drawings, in particular, can help make points clear or reinforce the presented information.

Effective visual aids enhance learning and reinforce essential information. Trainers who do not have access to an overhead projector may choose to copy the content of the included overheads onto flip chart pages. When trainers create their own overheads or flip chart pages, we hope they will follow these tips for effective visual aids:

- follow the participant manuals or handouts,
- reinforce or clarify one idea,
- stimulate thinking, discussion, and
- are simple and easy to read.

When using overheads, remember to:

- check the projector before training,
- keep an extra light bulb and extension cord on hand,
- position the projector so that all participants have a clear view of the screen,
- organize overheads for easy access during training,
- stand facing participants, reading from the machine or notes and not from the screen,
- use a pointer or cover portions of an overhead so that participants can easily follow the presentation, and
- give participants time to read each overhead before removing it from the screen.

Source:
The Great Lakes RAP. (1993). Train the Trainers. Foxon
About Handouts

Partners have included handouts in the *Supplemental Materials* in some sections of the manual. Trainers who choose to develop additional handouts may find the following tips helpful. Effective trainers:

- color code handouts for easy reference,
- distribute handouts essential to discussion before presenting,
- save supplemental handouts for distribution after training,
- get permission to reproduce copyrighted material, and credit original authors, and
- ensure that copies are clear and readable.

*Adapted from Wolfe, 1996.*
Welcome to the Partners Workshop

Objectives

- Register participants, introduce trainers, explain Partners Plus role
- Introduce participants, encourage interaction
- Outline workshop agenda

Preparations

Read the Introduction section of the Family and Caregiver Manuals carefully. Be familiar with this content.

Determine the number of participants. Prepare enough workshop materials for the group, including Family Manuals, Caregiver Manuals, workshop agendas, and photocopies of:

- Registration / Sign-In Sheet
- Initial Family Survey
- Caregiver Comfort Scale
- Caregiver Knowledge Scale

- What Did You Think of Training
- Get-To-Know-You Bingo (for groups of 10 or more only)

These forms are provided within the Supplemental Materials at the end of this section.

Prepare overhead transparencies of:

- Overhead Partners Plus PP W-1
- Overhead Partners Welcome (Part 1) PP W-2

Corresponding Sections

The content of this section corresponds with sections in the Family Manual or the Caregiver Manual.
Registration

Distribute the following materials to the group:

- Registration / Sign-In Sheet
- Initial Family Survey
- Caregiver Comfort Scale
- Caregiver Knowledge Scale

Allow participants sufficient time to register and complete forms. The information families and caregivers supply in their pre-evaluations (Initial Family Survey, Caregiver Comfort and Knowledge Scales) will provide you with helpful information about their experience, knowledge, and comfort levels prior to the workshop.

Once participants have completed their paperwork, distribute the following course materials:

- Workshop Agendas
- Workshop Manuals

This activity is not scripted. This activity does not require overheads.
Introductions

Introduce yourself. Tell your connection with the agency or organization sponsoring the Partners workshop.

Hello, and welcome. My name is ___________, and I’m a trainer for Partners Plus. Today’s workshop is supported by . . .

This activity does not require overheads.

Give participants a brief history of Partners Plus.

The original Partners Plus project, developed at Child Development Resources in Norge, Virginia, was initially funded by a grant from the U.S. Department of Education (#H024B40032). That Partners Plus project designed a model to help families find and train respite caregivers for their child(ren) with special needs. Four manuals were created to support families, caregivers, trainers, and community planners in implementing this model. With subsequent outreach funding from the U.S. Department of Education (#H324R990006) and using a train-the-trainer approach, the Partners project now helps communities across the country replicate this
Introductions continued

Display overhead and summarize the information it contains.

The original program was called Partners Plus: Families and Caregivers in Partnerships, and was developed using interviews and feedback from families.

Summarize the goals of the workshop.

We will go over the workshop agenda in a few minutes. When the workshop is over, individual families will have information and strategies to find and train caregivers for their children. Potential caregivers will have basic information and skills to work with families in respite.

model for young children with disabilities. Funding from the Joseph P. Kennedy, Jr. Foundation has supported the adaptation of the four Partners manuals to meet the needs of communities serving older children with mental retardation and related developmental disabilities.

Overhead Partners Plus 1-1

Partners Plus:
Families and Caregivers in Partnerships

- Developed with feedback from families at Child Development Resources
- Families needed a break but did not feel comfortable with available options
- Families wanted to find and train their own respite caregivers

Partners Plus Trainer's Workshop Manual
Child Development Resources, P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Introduction-4
Introductions continued

Give participants a brief history of the agency or organization supporting the workshop. If more than one group is sponsoring the program, speak about each group.

In your description, be sure to answer the following questions:

- How does Partners fit the mission of your group?
- Who does your agency or organization serve?
- What other relevant services does your agency or organization provide?

Today's workshop is supported by ____________ (agency or organization). Our purpose is to . . .

The Partners Plus program fits the mission of our group because . . .

We primarily serve . . .

We provide a number of services, such as . . .

This activity does not require overheads.
Introductions continued

Allow each of the trainers to introduce themselves. Have them explain their connection with the sponsoring organization.

Next, have those trainers that feel comfortable doing so share any personal or professional experiences with children with special needs, their families, and their caregivers.

Review housekeeping items, such as where restrooms are located, to help participants feel more comfortable with their training environment.

Reinforce your desire to allow open, active participation.

Hello. My name is ____________, and I'm also a trainer for Partners Plus. I work with the sponsoring group by ...

Having worked with families with children with special needs for many years, I am very in tune with the issues families face in caring for their child ...

Restrooms are located ...
Water fountains and soda machines are located ...
Feel free to help yourselves to some light refreshments, located ...

Feel free to ask questions or let us know your thoughts at any point during the discussion. We do have a lot to cover, however, and may have to limit conversation in the interest of time.

This activity does not require overheads.
Icebreaker Activity

Choose your icebreaker: Round Robin or Get-To-Know-You Bingo (used with a workshop audience of 10 or more).

**If Round Robin:**
Introduce the activity and explain the rules.

Now that we have introduced ourselves, we want to learn a little bit about you. We would like to go around the room and have you introduce yourselves. Tell us your name and a little bit about why you are here.

**If Get-To-Know-You Bingo:**
Introduce the activity and explain the rules.

Now that we have introduced ourselves, we want to learn a little bit about you. We’re going to play a game to get everyone moving and talking to each other. It’s called Get-To-Know-You Bingo. It’s played just like Bingo, but with a twist.

Distribute copies of Get-To-Know-You Bingo. Have participants wait to begin until all instructions are given.

In Bingo, you mark boxes on your card as each letter is called. Here, you mark a box on your card when a person tells you that she or he fits the description in that box. Write the person’s name in the box so that you remember what you know about him or her. When you have a row of names, call out “Bingo!”
Icebreaker Activity continued

If Get-To-Know-You Bingo:

When everyone understands the rules, have them begin. Encourage participants to speak to more than one person.

When someone calls “Bingo,” announce that there is one winner. If time permits, play a second round for another winner.

At the end of the game, have everyone return to their seats. Ask the winners to introduce the people they marked on their cards.

If possible, present small prizes to winners, such as pencils, candy, or certificates.

This activity is not scripted. This activity does not require overheads.
Review the Workshop Agenda

Refer participants to their agendas.

Briefly describe the sections on the agenda for Part 1 and summarize the objectives of each section.

Review the plan for the break between Part 1 and Part 2. Determine how you will handle the break—eat together, move into a separate space for meals, etc.

Note: If Part 1 and Part 2 are held on different days, review the information about Part 2 at both the end of the first day (end of Part 1) and the beginning of the second day.

This activity is not scripted.

Overhead 1-2 Partners Workshop (part 1)

Partners Workshop (Part 1)
Welcome and Introductions
About the Partners Model
Communication
Families and Caregivers Working Together
All Kids Like Cookies Activity
Review the Workshop Agenda continued

Briefly describe the sections on the agenda for Part 2 and summarize the objectives of each section.

This activity is not scripted.

Overhead 1-3 Partners Workshop (Part 2)

<table>
<thead>
<tr>
<th>Partners Workshop (Part 2)</th>
<th>Caregiver Break-Out Session</th>
</tr>
</thead>
<tbody>
<tr>
<td>Family Break-Out Session</td>
<td>Disabilities</td>
</tr>
<tr>
<td>Finding Respite Caregivers</td>
<td>Sensitivity</td>
</tr>
<tr>
<td>Interviewing Caregivers</td>
<td>Child Development</td>
</tr>
<tr>
<td>Developing a Personalized Training Plan</td>
<td>Children with Special Needs</td>
</tr>
<tr>
<td>Using Daily Routines to Develop a Training Plan</td>
<td>Becoming a Caregiver</td>
</tr>
</tbody>
</table>
Review the Workshop Materials

Refer participants to their manuals.

We will be referring to your manuals throughout the workshop. Your manuals follow the flow of the workshop.

We will highlight the essential information in your manuals during the workshop. However, we will not be able to cover everything. Your manuals will be yours to take with you after the workshop and to refer to as you build respite partnerships together.

This activity does not require overheads.
Answer Questions and Wrap Up

Conclude the section by answering questions.

We are looking forward to our workshop today! Do not hesitate to ask questions throughout the workshop or to let us know if you have additional needs.

Let's get started . . .

This activity does not require overheads.
Welcome to the Partners Workshop

Suggested Overheads
Partners Plus: Families and Caregivers in Partnerships

- Developed with feedback from families at Child Development Resources
- Families needed a break but did not feel comfortable with available options
- Families wanted to find and train their own caregivers
Partners Plus: Families and Caregivers in Partnerships

Unit 1–Welcome and Introductions
Unit 2–About the Partners Model
Unit 3–Building Partnerships: Communication
Unit 4–Families and Caregivers Working Together
Unit 5–All Kids Like Cookies Activity

Family Break-Out Session

Unit 6 Families–Finding Respite Caregivers
Unit 7 Families–Interviewing Caregivers
Unit 8 Families–Developing a Personalized Training Plan

Caregiver Break-Out Session

Unit 6 Caregivers–Disabilities Sensitivity
Unit 7 Caregivers–Understanding Child Development
Unit 8 Caregivers–Children with Special Needs
Unit 9 Caregivers–Becoming a Caregiver

Unit 10–Using Daily Routines To Develop a Personalized Training Plan

DRAFT 10.99
Welcome to the Partners Workshop

Supplemental Materials
Checklist of Materials and Equipment for Partners Workshop

Date: __________ Time: __________ Location: __________

Purpose: __________________________________________________________________

Number Expected to Attend: ________ Presenter(s): __________________________________________________________________

Room and Facility Preparation

___ Table for Registration
___ Table for Display Materials
___ Table Space for All Participants
___ Tables Arranged in a Horseshoe Shape
___ Chairs for All Participants
___ Extra Tables for "All Kids Like Cookies" Activity
___ Tables for Lunch or Refreshments (optional)
___ Overall Agenda for Training
___ Rooms for Childcare According to Numbers of Children Attending
___ Childcare Rooms with Accessibility to Lavatories (Preferably in Room)

Childcare Preparation

___ Caregiver Sign-In Sheet
___ Appropriate Toys and Activities
___ Activity Supplies
___ Snacks
___ First Aid Kit
___ "Partners Plus Workshop Childcare Information"

Getting Started

___ Refreshment Supplies (optional)
___ Name Tags or Tents
___ Sign-In Sheets
___ Initial Family Survey

Getting Started, continued

___ Comfort Measure-Pre
___ Knowledge Measure-Pre
___ Participant's Manuals
___ Overall Agenda for the Training
___ Getting Acquainted Activity
___ Pencils for Participants
___ Sample Press Release and Feature Articles form (optional)
___ Permission for Videotape with Sound / Photo Use (Enough for All Participants)

Section 1

___ Handouts
___ Prepared Overhead Transparencies / Flip Charts
___ Overhead Projector and Screen (optional)
___ Markers for Either Flip Charts or Overhead Transparencies
___ Flip Chart Stand
___ Incentives for Participation in Methods of Communication Activity

Section 2

___ Prepared Overhead Transparencies / Flip Charts
___ Overhead Projector and Screen (optional)
Workshop Materials, continued

Section 2, continued

- Markers for Either Flip Charts or Overhead Transparencies
- Flip Chart Stand
- Incentives for Participation in Methods of Communication Activity

Section 3

- Prepared Overhead Transparencies / Flip Charts
- Overhead Projector and Screen (optional)
- Markers for Either Flip Charts or Overhead Transparencies
- Flip Chart Stand

Section 4

- Handouts
- Enough of the Easy and Hard Version of the XYZ Tests (Equal Number of Each) for All Participants
- Method for Dividing Participants into Groups of Equal Number (for the "All Kids Like Cookies" Activity)
- Directions for Each Station
- Props for the "All Kids Like Cookies" Activity
  Vision Station: several pairs of prepared sunglasses, blindfolds, a page out of a coloring book or a crossword puzzle, crayons, children’s books and magazines, plastic cup and pitcher with water, paper towels
  Speech Station: marshmallows, gauze, trash can, paper towels
  Fine Motor Station: several pairs of thick gloves, masking tape, puzzles, small manipulative toys, thick string and beads or pasta
  Gross Motor Station: two large balls, two retrievable toys, masking tape, use masking tape to make two ten to fifteen foot walkways

Section 4, continued

- Prepared Overhead Transparencies / Flip Charts
- Overhead Projector and Screen (optional)
- Markers for Either Flip Charts or Overhead Transparencies
- Flip Chart Stand

Section 5-F

- Overhead Transparencies / Flip Charts
  Developed by Trainers (optional)
- Overhead Projector (if needed)
- Flip Chart Stand (if needed)

Section 5-C

- Prepared Overhead Transparencies / Flip Charts
- Overhead Projector
- Flip Chart Stand

Section 6-F

- Overhead Transparencies / Flip Charts
  Developed by Trainers (optional)
- Overhead Projector (if needed)
- Flip Chart Stand (if needed)

Section 6-C

- Prepared Overhead Transparencies / Flip Charts
- Overhead Projector
- Flip Chart Stand

Section 7-F

- Overhead Transparencies / Flip Charts
  Developed by Trainers (optional)
- Overhead Projector (if needed)
- Flip Chart Stand (if needed)
Partners Workshop Agenda

Date
Location
Time

Part II

Time  Registration and Light Refreshments

Time  Activity: All Kids Like Cookies

Time  Break Out Session for Families
Finding Respite Caregivers
Interviewing Caregivers
Developing an Personalized Training Plan

Time  Break Out Session for Caregivers
Understanding Child Development
Disabilities Awareness
Children with Special Needs
Becoming a Respite Caregiver

Time  Break - 5 minutes
Practice Activity: Using Daily Routines
Wrap up and Evaluations

Time  Congratulations! You are on your way!
Partners Workshop Agenda

Date
Location
Time

Part I

Time
Registration and Light Refreshments

Welcome and Introductions

About Partners

Communication

Time
Break - 5 minutes

Building Partnerships: Families and Caregivers Working Together

Preview of Coming Attractions

Time
See Ya for Part 2!
Partners Workshop Agenda

<table>
<thead>
<tr>
<th>Date</th>
<th>Location</th>
<th>Time</th>
</tr>
</thead>
</table>

Part II

**Time**

<table>
<thead>
<tr>
<th>Activity</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Registration and Light Refreshments</td>
<td><strong>Time</strong></td>
</tr>
<tr>
<td>Activity: All Kids Like Cookies</td>
<td><strong>Time</strong></td>
</tr>
<tr>
<td>Break Out Session for Families</td>
<td><strong>Time</strong></td>
</tr>
<tr>
<td>Finding Respite Caregivers</td>
<td><strong>Time</strong></td>
</tr>
<tr>
<td>Interviewing Caregivers</td>
<td><strong>Time</strong></td>
</tr>
<tr>
<td>Developing an Personalized Training Plan</td>
<td><strong>Time</strong></td>
</tr>
</tbody>
</table>

Break Out Session for Caregivers

<table>
<thead>
<tr>
<th>Understanding Child Development</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disabilities Awareness</td>
<td><strong>Time</strong></td>
</tr>
<tr>
<td>Children with Special Needs</td>
<td><strong>Time</strong></td>
</tr>
<tr>
<td>Becoming a Respite Caregiver</td>
<td><strong>Time</strong></td>
</tr>
</tbody>
</table>

Break - 5 minutes

<table>
<thead>
<tr>
<th>Practice Activity: Using Daily Routines</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wrap up and Evaluations</td>
<td><strong>Time</strong></td>
</tr>
</tbody>
</table>

Congratulations! You are on your way!
# Partners Plus Workshop Sign-In Sheet

**Location:** ______________________________  **Date:** __________________

**Trainers:** ______________________________

<table>
<thead>
<tr>
<th>Name</th>
<th>Address</th>
<th>Phone</th>
<th>How did you hear about Partners Plus?</th>
<th>Are you a Family or Caregiver?</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*DRAFT 10.99*
Partners Plus Initial Family Survey

Please describe any special needs your child might have. If you have more than one child, you may use the space on the back.

Child's Name: ___________________________ Age: ______

1. Does your child have any special needs such as: developmental delay, multiple disabilities, vision, hearing, feeding, behavior, heart problems, breathing problems, other? If so, please explain.

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

2. Has your child received early intervention, special education, or therapeutic services? If so, please explain.

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

3. Does your child use any assistive technology such as: electric wheelchair, medical equipment, communications board, other? Please describe.

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________
Initial Family Survey continued

Answer the following questions if you have ever left your child in another person’s care. Please circle the best answer.

4. In general, how hard has it been to arrange respite (temporary child care) for your child(ren)?
   - Not Difficult
   - Somewhat Difficult
   - Very Difficult
   
   1 2 3 4 5

5. How comfortable did you feel leaving your child(ren) with the caregiver?
   - Not Difficult
   - Somewhat Difficult
   - Very Difficult
   
   1 2 3 4 5

6. Have you ever taught a caregiver to care for your child?
   - Yes
   - No

7. How much time did you spend teaching the person how to care for your child?
   - Less than 30 mins.
   - About 3 hours
   - Over 3 hours

   0 1-3 4-6 6 and up

8. In the last three months, how often have you used respite care?
   0 1-3 4-6 6 and up

9. In the last three months, how often did you want respite, but could not find a caregiver?
   0 1-3 4-6 6 and up

10. In the last three months, how many times did caregivers cancel?
    0 1-3 4-6 6 and up

Please answer the following questions about your family members. This information is optional.

11. Child’s Mother’s Age ________  Child’s Father’s Age ________

12. What is the child’s mother’s highest level of education?
   - Middle School
   - High School
   - Some College
   - College Graduate (4 years)
   - Some Graduate School
   - Graduate Degree

13. What is the child’s father’s highest level of education?
   - Middle School
   - High School
   - Some College
   - College Graduate (4 years)
   - Some Graduate School
   - Graduate Degree
Partners Plus Caregiver Comfort Scale Pre-Evaluation

Please circle the number that best describes how comfortable you feel in working with children with special needs.

<table>
<thead>
<tr>
<th>How comfortable are you:</th>
<th>Very Uncomfortable</th>
<th>Very Comfortable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Providing care for a child with special needs.</td>
<td>1 2 3</td>
<td>4 5</td>
</tr>
<tr>
<td>Planning activities that a child with special needs can enjoy.</td>
<td>1 2 3</td>
<td>4 5</td>
</tr>
<tr>
<td>Meeting the routine health care needs of a child with special needs.</td>
<td>1 2 3</td>
<td>4 5</td>
</tr>
<tr>
<td>Feeding a child with special needs.</td>
<td>1 2 3</td>
<td>4 5</td>
</tr>
<tr>
<td>Talking with families of children with disabilities about their child and their child’s strengths and needs.</td>
<td>1 2 3</td>
<td>4 5</td>
</tr>
<tr>
<td>Talking to other people about any concerns they may have about children with special needs in your care.</td>
<td>1 2 3</td>
<td>4 5</td>
</tr>
<tr>
<td>Identifying resources and getting help you may need in caring for children with special needs.</td>
<td>1 2 3</td>
<td>4 5</td>
</tr>
</tbody>
</table>

Thank You!
Partners Plus Caregiver Knowledge Pre Evaluation

Name: __________________________ Date: __________________________

This survey is designed to gather information about your experience and knowledge of caring for children with disabilities or special needs.

Read each statement and circle the single best answer.

1. Early intervention services are available to children with disabilities who are
   a. Birth to three years old
   b. In public schools
   c. Three to five years old
   d. I don’t know

2. Children who receive early intervention special education services must
   a. Get therapy
   b. Go to a classroom program
   c. Have an individual plan
   d. I don’t know

3. All children with special needs receive education through
   a. MCH
   b. CHIP
   c. IDEA
   d. I don’t know

4. Which of these diagnoses tells you that a child has a special health care need?
   a. Autism
   b. Cystic Fibrosis
   c. Mental retardation
   d. I don’t know

5. Providing good respite care for a child with a disability depends heavily on
   a. Desire/attitude of caregivers
   b. Educational level of the caregiver
   c. A child’s disability
   d. I don’t know

6. When caring for children with disabilities in their homes, it is necessary to
   a. Learn about their daily household routines
   b. Change things to suit your way of providing care
   c. Ignore the way the family trained you to take care of the child
   d. I don’t know
7. A key to a successful respite care program for families is
   a. Providing each family with many caregivers
   b. Allowing each family to train their own caregivers
   c. To start more respite agencies
   d. I don’t know

8. It is most important for respite caregivers to
   a. Understand child development c. Have a degree or certification
   b. Know the needs of the child or children in their care d. I don't know

9. The term “cognitive development” is used to describe how a child
   a. Relates to others c. Thinks and solves problems
   b. Sits and grasps toys d. I don’t know

10. If caregivers have questions or concerns about a child’s special needs, they should
    a. Discuss their concerns with the child’s parent(s) or guardian
    b. Call the local special education program
    c. Not do anything because if they are wrong it would upset the child’s parents
    d. I don’t know

Thank You!
What Did You Think About the Workshop?

Name: __________________________ Date: ___________________

Presenter(s): ____________________ ______________________

1. What I liked about the workshop...

________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________

2. How Did We Do? The information presented was:

____ easy to understand

____ difficult to understand because...
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________

3. The information presented was:

____ useful

____ not useful because...
________________________________________________________________________
________________________________________________________________________
4. The overall quality of the workshop was:  
   1  2  3  4  5  
   Very Poor  Not at All  Very Good

5. Were the workshop materials helpful?  
   1  2  3  4  5  
   Not at All  Very

6. Was the information presented appropriate for your needs?  
   1  2  3  4  5  
   Not at All  Very

7. Was there enough chance for questions and discussion?  
   1  2  3  4  5  
   Not at All  Very

8. I gained knowledge about...

   ____________________________________________________________
   ____________________________________________________________
   ____________________________________________________________

9. Suggestions I would like to offer for improving the workshop...

   ____________________________________________________________
   ____________________________________________________________
   ____________________________________________________________

10. I think what I’ll remember most was...

    __________________________________________________________
    __________________________________________________________
    __________________________________________________________

11. Please share any additional comments:

    __________________________________________________________
    __________________________________________________________
## Get-To-Know-You Bingo!

<table>
<thead>
<tr>
<th>B</th>
<th>I</th>
<th>N</th>
<th>G</th>
<th>O</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volunteers at least once a week</td>
<td>Sings in a choir</td>
<td>Exercises to cope with stress</td>
<td>Can define respite</td>
<td>Works with persons with disabilities</td>
</tr>
<tr>
<td>Grew up in another state</td>
<td>Knows what IDEA stands for</td>
<td>Would rather draw a picture than write a story</td>
<td>Makes it a policy to laugh at least once a day</td>
<td>Loves to play with play dough!</td>
</tr>
<tr>
<td>Has a family member with special needs</td>
<td>Loves it when it snows!</td>
<td>FREE</td>
<td>Has younger brothers/sisters</td>
<td>Is carrying pictures of his/her family</td>
</tr>
<tr>
<td>Hates to sit still</td>
<td>Has no extended family in the area</td>
<td>Hasn’t seen a movie in a theater for over two years</td>
<td>Wears contacts or eye glasses</td>
<td>Is a member of a nontraditional family</td>
</tr>
<tr>
<td>Considers her/himself creative</td>
<td>Has never worked with someone with special needs</td>
<td>Enjoyed elementary school</td>
<td>Is certified in CPR</td>
<td>Has a pet</td>
</tr>
</tbody>
</table>
Partners Plus Order Form for Printed Materials

Name: ____________________________________________

Title, Company: ____________________________________

Street Address: ____________________________________

City, State, Zip: ____________________________________

Telephone: ______________________ Fax: _____________

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Item Number/Description</th>
<th>Unit Cost</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Subtotal ___________

Shipping and Handling ___________

Total ___________

Method of Payment:

Check number: __________________ P.O. number: _____________

Visa or Mastercard number ($25 Minimum Charge): __________________

Expires: __________________ Signature: _______________________

Shipping and Handling Charges:

Orders totalling $1.00 to $49.99 $7.00 Additional Fees required for international and priority delivery service

Orders totalling $50.00 to $99.00 $10.00

Orders totalling $100.00 or more $12.00

Return Completed Form To:

Child Development Resources, 1490 Government Road, Williamsburg, VA 23185

Phone (757) 220-1168 · Fax (757) 253-1779 · Email: ppcdr@tni.net

DRAFT 10.99
Section 1—About Respite

Objectives

- Introduce participants to respite and describe the Partners model of respite care
- Define respite care and identify the benefits to families, caregivers and children
- Demonstrate how families drive family-centered services
- Explain the rights and responsibilities individuals have as participants in family-centered systems
- Ensure participants understand the Partners model of respite care

Preparations

Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 1, “About Respite,” in the Family and Caregiver Manuals.

Prepare overhead transparencies of:

- Overhead Partners Plus What is Respite Care? 1-1
- Overhead Partners Plus Benefits of Respite to Families 1-2
- Overhead Partners Plus Benefits of Respite to Children 1-3
- Overhead Partners Plus Benefits of Respite to Caregivers 1-4
- Overhead Partners Plus Why Don’t Families Use Respite Care? 1-5
- Overhead Partners Plus Partners . . . 1-6
- Overhead Partners Plus Partners Model Steps 1-7
- Overhead Partners Plus What Respite Means 1-8

Corresponding Sections

Please refer participants to Section 1, “About Respite,” in the Partners Plus Family and Caregiver Manuals.
Respite

The goal of this activity is to involve participants in discussion early in the program and to get them interested and excited about respite and the Partners model.

Introduce the concepts of the section. Refer to overhead Partners Plus What is Respite Care? 1-1.

Let's begin by discussing what we mean when we are talking about respite and respite care. The term “respite” means relief, or to take a break. “Respite care,” then, is occasional child care that gives families respite, or a break from the daily demands of raising children with special needs.

All families need a break, but families who have children with special needs may have a more difficult time finding reliable, responsible caregivers.

Overhead Partners Plus 1-1

What is Respite Care?
- Respite care is occasional child care that gives families a break (that gives families respite) from the daily demands of raising children with special needs.
Respite continued

Benefits of Respite to Families

Introduce the benefits of respite for each participant.

Ask the group to talk about how family members benefit from time away. Give families and caregivers a few minutes to generate answers.

As people share comments, they are educating each other. Affirm their answers: respite can be everything for families that they want it to be.

You may need to stimulate the conversation with a few examples. Sharing your own personal experience is best.

There are three participants in respite relationships: family members, the children with special needs who are receiving care, and respite caregivers. Let's take a minute to consider how each member of the respite relationship benefits from this experience.

How might individual family members benefit from time away from caring for a child with special needs?

How might they spend their time, their respite?

What about resting, entertaining themselves, shopping, pampering each other? What else?
Respite continued

Refer to overhead Partners Plus Benefits of Respite to Families 1-2. Reveal information one line at a time in order to stimulate discussion.

Briefly highlight the benefits of respite that participants do not list. Refer them to Section 1 in the Partners Plus Family and Caregiver Manuals.

Benefits of Respite to Children

Introduce a few ways in which children with special needs also benefit from respite care.

Ask the group to list other ways children with special needs might benefit from respite. Again, take a few minutes to generate responses.

Excellent. Those are all benefits of respite. What are some others? Pursuing interests, spending time with other family members and friends?

Great. Let's go over a few more ways that families can benefit from respite, like . . .

It's not just families who benefit from respite; children benefit also.

Respite gives the child the opportunity to interact with new people and develop social skills. Respite also introduces children to different caregivers, who bring different experiences. Children feel a sense of relief when their families get a break.
Respite continued

Refer to overhead Partners Plus Benefits of Respite to Children 1-3. Reveal information one line at a time in order to stimulate discussion.

Briefly highlight the benefits of respite that participants do not list. Refer them to Section 1 in the Partners Plus Family and Caregiver Manuals.

Benefits of Respite to Caregivers

Introduce the benefits of respite to caregivers as a starting point for discussion.

Did you know that after providing respite, caregivers often say that they learn far more from the children they work with than those children learn from them?

Respite gives children with special needs the opportunity to teach others about their experience. It also helps send the message that children with special needs are not so different from other children.

The third group to benefit from respite care are the caregivers themselves. Caregivers meet some amazing children and families, and make an important difference in their lives.

Respite care encourages caregivers to learn new skills, such as sign language. Some caregivers view respite as a job, and value the working experience.

Overhead Partners Plus 1-3

Benefits of Respite to Children

- Relief
- Exploration
- Special friends
- Parents feel rejuvenated
- Interaction
- Time for independence
- Educate others
Respite continued

Ask participants to list benefits of the respite experience.

Refer to overhead Partners Plus Benefits of Respite to Caregivers 1-4. Refer participants to Section 1 in the Partners Plus Family and Caregiver Manuals.

Wrap up the discussion.

What are some other ways caregivers might benefit from providing respite?

So, all three of the members of a respite relationship can and should benefit from the respite experience. Respite is important to the well-being of children and families, and caregivers can feel confident that what they are doing is important, life-enriching work!

Overhead Partners Plus 1-4

Benefits of Respite to Caregivers

- Really great children and families
- Especially rewarding
- Self-improvement
- Positive attitudes
- Involvement
- Truly appreciated
- Everything they want
Barriers for Families Who Want Respite Care

Introduce this section by asking this question:

Discuss potential barriers to respite.

Refer to overhead Partners Plus Why Don't Families Use Respite Care? 1-5 during the discussion. Refer participants to Section 1 in the Partners Plus Family and Caregiver Manuals.

So, if respite is so wonderful, why aren't families using respite care?

It can be very hard to find a trusted, familiar person who is willing to provide care, particularly with the training and the commitment families are looking for.

Sometimes, caregivers cancel. Planning for respite care can be far more energy- and time-consuming than planning for other child care. Imagine how it would feel to do all of that planning and get excited about having some time away, only to have a caregiver cancel at the last minute.

Also, families expect a lot from themselves, and may feel guilty that they need a break. And, finding community respite programs can be difficult.

Overhead Partners Plus 1-5

Why Don't Families Use Respite Care? Families . . .

- Struggle to find caregivers
- Fear leaving children with strangers
- Think caregivers don't have enough training
- Feel guilty
- Can't find community programs
- Feel too disappointed when caregivers cancel
The Partners Model

Introduce the section by explaining the Partners concept. Review the overhead to outline the components.

Refer participants to Section 1 in the Partners Plus Family and Caregiver Manuals for a definition of “family-centered.”

Refer to overhead Partners Plus 1-6 Partners... Participants may follow along in Section 2 of their manuals.

The Partners model of respite training was developed in response to the barriers and concerns identified by families. Families also identified components that are essential to a successful respite program.

First, Partners is a family-centered model. When we say family-centered, we mean that we recognize the family as the constant in a child’s life. We believe that services like ours must support the child by supporting, respecting, and encouraging the family.

We do this by giving families options for how they participate, who their caregivers are, how they are trained, and how respite is provided. We support families in finding caregivers and in training them. By doing this, we hope to expand respite options for families.

How does Partners achieve these goals?

Overhead Partners Plus 1-6

Partners...

- is a family-centered model
- offers families choices
- supports families and caregivers
- helps families and caregivers build skills to communicate
- helps families and caregivers build long-term relationships
Refer participants to overhead Partners Plus 1-7 Partners Model Steps.

There are five steps in the Partners model. In Step 1, the project recruits potential caregivers and families for participation in the program.

In Step 2, families and caregivers attend a workshop, which is the first part of the training process. At the workshop, families learn to find, screen and train their own caregivers. Meanwhile, caregivers get basic information on working with children with special needs and their families.

In Step 3, the program helps families choose caregivers who they want to provide respite care for their children. Families are best able to choose caregivers for their children. Likewise, caregivers are best able to decide who they will be successful with. The Partners program provides families and caregivers with information and resources to interview each other and make choices about working together.

Note: As you discuss Step 3, be sure to describe briefly how your community's Partners Plus program supports families and caregivers as they try to make matches.

<table>
<thead>
<tr>
<th>Overhead Partners Plus 1-7</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Partners Model Steps</strong></td>
</tr>
<tr>
<td>Step 1</td>
</tr>
<tr>
<td>Step 2</td>
</tr>
<tr>
<td>Step 3</td>
</tr>
<tr>
<td>Step 4</td>
</tr>
<tr>
<td>Step 5</td>
</tr>
</tbody>
</table>
Note: As you discuss Step 4, be sure to describe briefly how your community's Partners Plus program supports families and caregivers in personalized training.

In Step 4, the Partners program assists families as they train their own caregivers. In workshops, we give caregivers basic information on working with children with special needs and their families. After group training, families, who know their children best, teach caregivers about their particular children. At the same time, we give families a structure for giving caregivers training specific to their children. We call this training "personalized training."

Finally, in Step 5, Partners provides continuing support in two ways. First, Partners is available to families and caregivers as they work together in respite. Second, Partners staff ensure that this respite program is continued in the community.
Summary

Conclude the section by asking if there are any questions or comments about respite or the Partners model. You may need to curtail discussion politely in order to keep within the time allotted.

Any questions or final thoughts on respite or the Partners model that you'd like to share with the group?

Great. In closing this section, I want to bring it back to the importance of respite for families. This quote is from a father who participated in Partners: “Think of a candle that slowly burns, the flame flickers and goes out, and just as a candle, we need to be relit too.”

Respite is a lifesaver for many families, and respite care is an important job for caregivers who want to make a difference!

Overhead Partners Plus 1-8

What Respite Means

“Think of a candle that slowly burns, the flame flickers and goes out, and just as a candle, we need to be relit too.”

— A Parent
Section 1—About Respite
Suggested Overheads
What Is Respite Care?

Respite care is occasional child care that gives families a break from the daily demands of raising children with special needs.
Benefits of Respite to Families

Rest
Entertainment
Shopping
Pampering
Interests
Time
Everything you want it to be!
Benefits of Respite to Children

Relief

Exploration

Special friends

Parents feel rejuvenated

Interaction with other people

Time to be independent

Educating others
Benefits of Respite to Caregivers

Really great children and families

Especially rewarding

Self-improvement

Positive attitudes

Involvement

Thoroughly appreciated

Everything you want it to be!
Why Don't Families Use Respite Care?

Families . . .

- struggle to find caregivers
- fear leaving children with strangers
- think caregivers don't have enough training
- feel guilty
- can't find community programs
- feel too disappointed when caregivers cancel
Partners . . .

- is a family-centered model
- offers families choices
- supports families and caregivers
- helps families and caregivers build skills to communicate
- helps families and caregivers build long-term relationships
Partners Model Steps

Step 1  Recruit families and caregivers

Step 2  Conduct initial workshop

Step 3  Help families choose caregivers

Step 4  Support families as they train their own caregivers

Step 5  Provide continuing support
What Respite Means

"Think of a candle that slowly burns . . . the flame flickers and goes out. Just as a candle, we need to be re-lit, too."

— A Dad
Section 2—Building Partnerships: Communication

Objectives
- Build communication skills
- Define the importance of clear and open communication in a respite relationship
- Identify the variety of ways that people send and receive messages
- Learn how to recognize and mend communication breakdowns

Preparations
Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 2, “Building Partnerships: Communication,” in the Family and Caregiver Manuals.

Prepare overhead transparencies of:
- Overhead Partners Plus Communication Is... 2-1
- Overhead Partners Plus How Do You Feel? 2-2
- Overhead Partners Plus The Record 2-3
- Overhead Partners Plus Six Messages of Communication 2-4
- Overhead Partners Plus Non-Verbal Communication and Lifeways 2-5
- Overhead Partners Plus Building Partnerships Through Communication 2-6

Corresponding Sections
Please refer participants to Section 2, “Building Partnerships: Communication,” in the Partners Plus Family and Caregiver Manuals.
Introduction

Introduce this section by talking about how communication helps families and caregivers build lasting relationships with each other.

Our goal with Partners is to help families and caregivers build long-lasting respite relationships. We believe that communication and understanding are the most important tools to build these partnerships. In this section, we will talk about how communication works, and how it can be used in respite.

Refer to overhead Partners Plus Communication Is . . . 2-1. Participants may follow along in Section 2 in the Partners Plus Family and Caregiver Manuals.

When we talk about communication, we are talking about the way that information is passed from one person to another.

We communicate in a lot of different ways. Research tells us that we communicate 7% of our messages through the words that we use. We choose our words very carefully.

Overhead Partners Plus 2-1

Communication Is . . .

- 7% Words
- 38% Voice
- 55% Facial and Body Language
In the field of disability, we use many words that we do not use anywhere else. When people enter this field and try to communicate, they might feel like they are learning a new language. When caregivers and families work together, it is very important that they feel comfortable asking what words mean.

Even more important than the words that people choose is the voice that they use. You can use the very same words and communicate two different messages just by changing your voice. When we talk to people, we need to be aware of how we use our voices.

The most powerful way that we communicate is through our facial and body language. Though none of you are speaking to me right now, all of you are telling me something with your bodies and faces. For example, your posture is telling me . . .
Non-Verbal Communication Activity

Introduce the activity.
Refer to overhead Partners Plus How Do You Feel? 2-2.

Explain the activity.

Let's try to illustrate this point further. Please turn to page 2-4 of your manuals. We are going to do a short activity to help us better understand how we communicate with facial expressions and body language.

On your page, there are some faces and some blank circles. Think about an emotion that you were feeling before you came to training today—an emotion different from those shown here. Now, draw what you think your face looked like when it was expressing that emotion. Don't label your picture, just draw it.

Now let's divide into groups of two. Try to guess what emotion your partner drew.

Give the group a minute to complete the activity. Then, divide the group into pairs. Ask the participants to guess the emotion that their partners drew. Spend two minutes on this.
Non-Verbal Communication Activity continued

Ask participants to share what they learned. You might need to prompt with some questions to facilitate discussion.

Draw conclusions for the group.

How did you do? Did you find it easy to guess what emotion was drawn?

How many guessed correctly? For those who did: So the faces communicated clear messages to you. Sometimes, people have trouble interpreting the looks on people’s faces. They might need to use words to clarify. They might say, “Oh, you look angry. Did I say something to upset you?” This is a good way to check to make sure that their perceptions are right.
Building Communication Skills

Refer to overhead Partners Plus The Record 2-3.

Like faces, the same words can mean different things to different people. Here, the same sentence has different meanings for two people with two different definitions of the same word. Again, using words can help clarify. Families and caregivers can use the words "describe," "show me," and "tell me," to clarify misunderstandings.

Refer to overhead Partners Plus Six Messages of Communication 2-4. Participants may follow along in Section 2 in the Partners Plus Family and Caregiver Manuals.

When we think about communication, it makes sense to think of it as a process. There are many places that communication can break down. Sometimes, what people say is not what they mean. At other times, someone hears something different from what was said or misinterprets a message. Families and caregivers must pay attention to communication, and watch for misunderstandings. Feel free to ask again if you are confused about what the other person is communicating.

Overhead Partners Plus 2-3

The Record
- She has a great record! (music record)
- She has a great record! (data record)

Overhead Partners Plus 2-4

Six Messages of Communication
- What you mean to say
- What you actually say
- What the other person hears
- What the other person thinks he/she hears
- What the other person says
- What you think the other person says
We know that communication is influenced by what we call our "life ways." People in different cultures, and even different families, learn their own ways of communicating.

When you are working with each other, remember that it may take some time for you to feel comfortable with each other's communication.

A lot of life ways differences in communication are nonverbal. For some people, too much silence can be uncomfortable. Other people need some pauses in conversation.

Different people think about body space differently too. In America, we usually feel comfortable talking to someone who is about a foot away from us. In other cultures, they prefer people to be closer when they are talking.
Make the following points.

For some people, eye contact conveys trust, like the person is listening and interested. In some cultures, however, eye contact is considered disrespectful.

Body postures can tell us a lot too. In America, an open body posture, where the arms are uncrossed, is more inviting than a closed body posture, with the arms crossed to block you.

People's eyes can be very expressive. Families often use glances to convey meaning. For example, some couples can look at each other from across a crowded room at a party and let each other know that it is time to go. Parents often have looks that tell their children that they need to behave.

Finally, presumed understanding can be very powerful. Similar experiences can really help speed up communication.
Building Communication Skills continued

The following activity is designed to help participants talk more specifically about how communication facilitates the respite relationship.

For example, when families of children with disabilities get together, it can be a relief for them to know that the person next to them knows how difficult it is to find good care for their children. It is helpful for caregivers to share with families the experiences that they have had working with children with special needs. This gives families a better idea of what they need to tell you and what you already know.

We have talked about communication and how it works. Now, let's look at communication in action in respite.

There are several scripts laid out in this section. The first script is of a conversation between a family and a caregiver who are just getting to know each other.

Ask participants to turn to Section 2: "What You Need to Know About Communication" in their manuals. Introduce the first script.

Overhead Partners Plus 2-5

Non-Verbal Communication and Culture Life Ways (Our Customs that Reflect Our Culture and Values)

- Reactions to silence
- Amount of space between bodies
- Eye contact
- Body postures
- Eyes that speak
- Presumed understanding that comes from similar experiences
Building Communication Skills continued

Ask for two volunteers: one to read the Family part, and one to read the Caregiver part. When you have assigned roles, ask the participants to read the entire first script aloud.

Ask participants to talk about the kinds of communication at work in this script.

Participants will probably generate a lot of answers. It is particularly important to highlight the way that the family encouraged the caregiver to share more information about what she liked to do with children.

Move on to the next script on the next page.

What kinds of communication do we see used here? Verbal? Nonverbal?

What about the family. How did they encourage the caregiver to share more information? What language did they use?

This second script is a discussion that takes place during personalized training. The caregiver is learning about the child's feeding routine.
Ask for more volunteers to read the roles in the second script. When they have done so, ask participants to share what they heard.

Again, there may be many answers. Here, highlight the way that the family perseveres, despite the caregiver’s fears, and the words that the caregiver uses.

Move to the final script at the bottom of the page.

What did you hear going on in that script?

Notice how the family perseveres when the caregiver expressed fear in a negative way. These words were not intended to be hurtful, but they could have sounded hurtful to the family. But the family did not shut down, instead they continued to probe for additional information until the caregiver felt comfortable.

This final script is a conversation between a family and a caregiver after care has taken place. It is critical that families and caregivers continue to communicate throughout their relationships. This family has taken the time to check in with their caregiver after respite.
Building Communication Skills continued

Ask for two more volunteers to read through the script. After they have finished reading, ask for comments on communication.

Here, you want to make sure that participants recognize the need to check in after respite.

What did you hear going on in that script?

This is a great example of the need to check in after respite. This caregiver was open to sharing her experience, which is very helpful, too. Both the family and the caregiver use words to gather more information and to clarify what happened.
Summary

Ask if there are any questions or comments about this section. Refer to overhead Partners Plus 2-6 Building Partnerships Through Communication. Refer participants to Section 2 in the Partners Plus Family and Caregiver Manuals.

Summarize the key points of the section.

Any questions on what we've covered so far?

As families and caregivers work together, they need to know how to communicate with one another. This communication will strengthen the partnership. Remember these simple points as you work together:

- Sharing thoughts and feelings helps you understand each other.
- It is as important to listen to each other as it is to talk to each other.
- Recognize and appreciate common experiences, similarities, and differences.
- Honesty and openness allow you to build cooperative, long-lasting relationships.

Overhead Partners Plus 2-6

Building Partnerships through Communication

- Sharing information increases understanding
- Listen carefully to each other!
- Share common experiences
- Appreciate differences
- Honesty and openness are key
Section 2—Building Partnerships:

Communication

Suggested Overheads
Communication Is . . .

7 % Words

38 % Voice

55 % Facial and Body Language

(UCLA, 1967)
How Do You Feel?

Anxious  Angry  Bashful  Bored  Confident

Thinking  Scared  Innocent  Relieved  Confused

Adapted from feelings posters by:
The Record

She has a great record!
Six Messages of Communication

• What you mean to say
• What you actually say
• What the other person hears
• What the other person thinks he or she hears
• What the other person says
• What you think the other person says
Nonverbal Communication and Life Ways

- Reactions to silence
- Amount of space between bodies
- Eye contact
- Body postures
- Expressive eyes
- Presumed understanding that comes from similar experiences
Building Partnerships through Communication

- Share information to increase understanding
- Listen carefully to each other
- Share common experiences
- Appreciate differences
- Be open and honest
Section 3—Building Partnerships: Families and Caregivers Working Together

Section Purpose

• To give participants an overview of and introduction to building successful relationships with each other.

Objectives for Families

• Become more aware of caregivers’ perspectives
• Understand why many caregivers experience “burn out” and stop providing respite care
• Know strategies to prevent caregiver burn out

Objectives for Caregivers

• Become more aware of families’ perspectives
• Recognize families’ need for respite and the value of the service that respite caregivers provide
• Understand the importance of learning about a family’s experience from that family

Preparations

Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 3, “Building Partnerships: Working Together,” in the Partners Plus Family and Caregiver Manuals.

Read each family story so that you feel comfortable reading any of them aloud word for word.

Select a training option to discuss family stories. Be prepared to either facilitate discussion, share personal experiences, or discuss “A Personal Story,” outlined in Section 3 of the Family and Caregiver Manuals.

Prepare overhead transparencies of:

• Overhead Partners Plus The Challenges of Respite 3-1
• Overhead Partners Plus Meeting the Challenge 3-2
• Overhead Partners Plus Successful Partnerships 3-3
Preparing for Training continued

Note to Trainers
Often during workshops, participants meet objectives for this section in previous sections. If so, make appropriate modifications in your presentation.

Corresponding Sections

Notes about Presenters
The first part of this section deals specifically with building relationships with families who have children with special needs. As a result, we strongly recommend that a trainer with personal experience as a family member facilitate or co-facilitate this session. Depending on their comfort level, family members can provide a different perspective through personal stories and anecdotes.

Similarly, the second part of this section deals with building relationships with caregivers. We suggest that a trainer who has experience as a caregiver for children with special needs facilitate or co-facilitate this session. Again, trainers with any kind of experience working with children with special needs can add to the training by sharing their experiences if they are comfortable doing so. Family members who have worked with respite caregivers can also provide valuable insight as trainers in this part of the section.

During this section, participants may have information to share. Group sharing can be unpredictable and challenging to facilitate. However, families and caregivers who participated in the field testing of this model consistently said the stories they heard in this group discussion provided some of the most valuable information that they learned.

We strongly suggest that you take the time to allow participants to share during this section. Extra time is built into the agenda to allow you to do that.

If you must curtail discussion in the interest of time, use positive language, such as: “I hope our discussion can continue during a break in the training, but unfortunately, we need to move on in order to address all of the information and stay on schedule. Thank you for sharing your experiences with us.”
Introduction to Sharing Perspectives

Introduce this section by talking about how understanding helps families and caregivers build lasting relationships with each other.

We have talked about communication, which we feel is one cornerstone of a strong respite relationship. Understanding is another cornerstone that grows only out of communication.

People who understand each other's perspectives can interact more respectfully and sensitively. In respite, once you match up with caregivers and families, it is important that you spend time sharing perspectives, and talking honestly about how the relationship is working for you. In this section, we will share stories from families and caregivers about what it is like to be in their shoes.

As we talk, we hope that you will feel free to add your perspective. We may say things that ring true with you, and other things that are different from your experience. Everyone has slightly different experiences. We will start with family perspectives and move on to caregiver perspectives later.

There are no overheads for this section.
Family Perspectives

"Welcome to Holland"

As an introduction to family perspectives, read this story to participants exactly as it is written. This story is one family's perspective, and the message must be delivered as intended by that family.

In order to give the most powerful representation of this family's experience, read expressively with sincere emotion for their concerns.

Participants may want to follow along in Section 3 of the Partners Plus Family and Caregiver Manuals.

I would like to read you a story written by the parent of a child with Down Syndrome. It expresses one parent's view of what it is like to have a child born with special needs. The story is called "Welcome to Holland."

I am often asked to describe the experience of raising a child with a disability—to try to help people who have not shared that unique experience to understand it, to imagine how it would feel. It's like this . . .
Read “Welcome to Holland” for the group.

When you’re going to have a baby, it’s like planning a fabulous vacation trip—to Italy. You buy a bunch of guidebooks and make your wonderful plans—the Coliseum, the Michelangelo’s David, the gondolas in Venice. You may learn some handy phrases in Italian. It’s all exciting.

After months of eager anticipation, the day finally arrives. You pack your bags and off you go. Several hours later the plane lands. The stewardess comes in and says, “Welcome to Holland!”

“Holland?!?” you say. “What do you mean, ‘Holland’? I signed up for Italy! I’m supposed to be in Italy. All my life I’ve dreamed of going to Italy!” But there’s been a change in the flight plan. They’ve landed in Holland and there you must stay.

The important thing is that they haven’t taken you to a horrible, disgusting, filthy place, full of pestilence, famine and disease. It’s just a different place.

So you must go out and buy new guidebooks. And you must learn a whole new language. And you will meet a whole new group of people that you would have never met.
Family Perspectives continued

Read “Welcome to Holland” for the group.

It’s just a different place. It’s slower-paced than Italy, less flashy than Italy. But after you’ve been there for awhile you look around and you begin to notice Holland has windmills, Holland has tulips, Holland even has Rembrandts.

But everyone you know is busy coming and going from Italy, and they’re bragging about what a wonderful time they had there. And for the rest of your life, you will say, “Yes, that’s where I was supposed to go. That’s what I had planned.” The pain of that will never, ever, go away because the loss of that dream is a very significant loss.

But if you spend your life mourning the fact that you didn’t go to Italy, you may never be free to enjoy the very special, the very lovely things in Holland.

(Emily Perl Kingsley, 1981)
Summarize the key messages of the story in your own words.

Few families plan for or expect to have a child with special needs. To develop a partnership with that family, caregivers should think about what that experience might be like.

In this story, Ms. Kingsley, shares her thoughts and feelings about learning that she was the parent of a child with special needs. She talks about the huge impact that child's disability had on her life and other family members. She talks about the pain that will never go away that came because of her child's special need.

But Ms. Kingsley also tells us that her child is special, not just in his special needs but also in the joy he brings. It is important to remember that there are joys and triumphs in raising and loving a child with special needs.
Family Perspectives continued

Note to the Family Presenter:

There are several different ways to meet the objectives of this section: either by asking the group to share experiences, or by sharing your own experiences, or by following the discussion outlined in Section 3 of the Partners Plus Family and Caregiver Manuals. Choose the method most comfortable for you.

If you select Options 1 or 2 (below), you may find that it takes less than 20 minutes. Remind participants that they can continue sharing information during a break or after training. If you select Option 1 or 2 and after you have completed the discussion, proceed to page 3.14 of the Trainer's Workshop Manual to continue with trainer notes, content, and section objectives.

If Option 1:
Sharing Group Reactions

Invite families to share their reactions to this story with the group.

There are a number of families here who I'm sure can relate to this story. Would anyone like to share their personal experience with caring for a child with special needs? What memories or feelings did this story raise for you?
Family Perspectives continued

If Option 2:
Sharing Personal Experiences
This is a good time for family presenters to share related personal stories if they wish to do so.

I can really relate to this story. I have a personal history of caring for a child with special needs . . .

If Option 3:
Following Section Discussion
Refer participants to “A Personal Story” in Section 3 of the Partners Plus Family and Caregiver Manuals.
Give participants some time to read through the story.

Now let’s take a few moments to read the story by Ms. Moore to ourselves. Once everyone has finished we can share our reactions.

There are no overheads for this section.
If Option 3 (continued):  
Following Section Discussion

Follow the discussion outlined at right.

Ms. Moore touches on a lot of important topics in her story. She begins by talking about the impact Matthew's birth had on her family. Like Ms. Kingsley's family, her family lived a very different kind of life when her child was born. Matthew required round-the-clock care, and certainly his birth changed their life. Even children who require less intense care affect families. Sometimes, families no longer feel comfortable in the communities where they live.

People who do not take the time to learn about children and their special needs sometimes react in ways that are hurtful to the child and the family. Sometimes, people stare or ask insensitive questions or say insensitive things. Sometimes, close friends of the family are so unsure of what to say they never come by to say anything at all.
If Option 3 (continued):
Following Section Discussion

Ms. Moore goes on to describe more about the impact Matthew had on her family. She talks about what they learned from Matthew. Often, people in the community only see the negative side to raising a child with a special need. The reality is that there are hard times. However, families tell us that there are many positive sides to raising their children too. Matthew taught Ms. Moore's family a lot about their talents as well. They learned that there were people in their community who would support them.

Other people in Ms. Moore's community were blinded by Matthew's disability. They could not, or would not, see the child behind the diagnosis. One of the most important things that we share with you today is this: all children have special gifts and special needs. When you are working with children with special needs, remember to look for the children's strengths. Your respite relationship will be made stronger!
Family Perspectives continued

If Option 3 (continued):
Following Section Discussion

Ms. Moore also talks about the professionals who worked with her family and Matthew. Very often, families with children with special needs have many people coming in and out of their homes. The services that these professionals provide can be very helpful to families and children. Often, their work is valued highly by families. However, it is difficult to have so many people in their homes. Their lives are less private. Needing help to learn how to care for their children can make parents feel inadequate at times. Caregivers can be sensitive to those issues.

Ms. Moore also talks about her family's goals for Matthew. Her family wanted two things for her son. All families have dreams for their children. Ms. Moore’s goals for Matthew are not so different from the goals other families have for their children. In working with families, it might be important for caregivers to take some time to

There are no overheads for this section.
Family Perspectives continued

If Option 3 (continued):
Following Section Discussion

Follow the discussion outlined at right.

learn the goals that families have for children. Listen to what families are doing to help children toward their goals. Caregivers might also have some ideas about how to help them reach their goals.

Ms. Moore closes by talking about respite and what it meant to her family. The respite care that Ms. Moore received was important to her whole family. As we said earlier, families need time to relax, recuperate, focus on other things, and maintain other relationships.

Ask for input from the group.

Would any of the families in the room like to share what respite might mean for their family?

There are no overheads for this section.
Family Perspectives continued

Following Section Discussion

Continue this discussion by talking about the stresses of raising children.

Throughout this chapter in the manual, there are stories written by family members. The stories are about children with a range of ages and special needs. I encourage you to take the time to read through these perspectives later as they will give you a deeper understanding of families, and how important respite can be.

Raising any child can be challenging. Research tells us that changes in families' lives pose the greatest challenge. Families of children with special needs face the same worries and concerns as other parents.

Children with special needs may need additional support. Some children with special needs require extra help with everyday tasks like eating, moving from place to place, or communicating. In addition, when families of children with special needs are making larger life changes, their choices may be more limited. For example, when they select schools for their children, they need to consider a whole list of service providers in addition to a teacher.
Family Perspectives continued

Following Section Discussion

Continue this discussion by talking about the stresses of raising children.

It is not surprising that these families experience more stress than families with typically developing children. The key to dealing with stress depends on how you perceive a situation. The same situation can be stressful to one person but not stressful for another.

If you perceive a situation as stressful, you can reduce your stress by using a variety of coping strategies.

Respite can provide stress relief alone. It also gives families the opportunity to do some of these activities that relieve stress.

There are no overheads for this section.
Family Perspectives continued

Following Section Discussion

Refer participants to “Strategies for Dealing with Stress” in Section 3 of the Partners Plus Family and Caregiver Manuals.

It is important to remember that caregiving can be stressful for caregivers as well. We encourage families to share information with caregivers about how to work with their children. We also encourage open communication. Talk with caregivers about how they deal with stress. Can you help them relieve the stress they may have when caregiving?

There are no overheads for this section.
Challenges of Respite

Introduce the concepts of this section.

Refer to overhead Partners Plus The Challenges of Respite 3-1. Participants may wish to follow along on in Section 3 in the Partners Plus Family and Caregiver Manuals.

When the Partners Plus model was first developed, families said that they had trouble keeping caregivers.

Caregivers gave these reasons for stopping respite care:

- Respite is harder than they expected. As families are training their caregivers, we encourage you to be honest with them. Tell caregivers the challenges of working with your children as well as the joys.

- Time or physical demands are too great. Caregivers can dedicate too much of their time and energy to respite. Too much, too fast, can lead to caregiver burnout. Families and caregivers need to spend time talking realistically about the caregivers' schedules.

Overhead Partners Plus 3-1

The Challenges of Respite
Caregivers may stop giving respite if:
- Respite is harder than they expected
- Time demands or physical demands are too great
- They don't feel appreciated
- Training is not adequate
- “Burn out” occurs
Challenges of Respite continued

Explore the reasons why caregivers may stop giving respite care.

- Caregivers also need to be adequately trained. They need to have the skills to give the care children need. Without this training, caregivers will be uncomfortable and feel stressed. Later in this training we will talk more with families and caregivers about personalized training.

- Sometimes, caregivers' patience just wears out. If caregivers seem stressed, families must take the time to find out why and how they can help.

- Finally, caregivers leave if they don't feel appreciated. Families can show their appreciation for caregivers in a variety of ways.

Some strategies are listed in Section 3 of the Family Manual.
Meeting the Challenge

Refer to overhead Partners Plus Meeting the Challenge 3-2. Participants can follow along in Section 3 in the Partners Plus Family and Caregiver Manuals.

Families and caregivers can work together to maintain their relationships. Caregivers can learn families' purpose for respite. This gives them a sense of how much their work is needed and valued.

Caregivers can be active in developing personalized training plans. In this way, they can make sure that they get all of the information they need to feel comfortable.

Be clear about your terms for providing respite. Take the time to consider your schedule. Overcommitting leads to burnout.

Learn more about working with children and families. Today, we will give caregivers information to use while providing respite care. We encourage caregivers to continue collecting information that will be helpful to them in their work.

Overhead Partners Plus 3-2

- Understand the purpose for respite
- Be active in helping to develop training
- Be clear about your terms
- Learn about working with children and families
- Get information and support from other caregivers
Explore ways to help caregivers better provide respite care.

Also, make the most of your personalized training time. Get to know the families and children you will be working with. As we said earlier, understanding is a powerful tool.

Finally, get information and support from other caregivers. It is helpful to have someone who you can discuss problems and share ideas with!
Summary

Ask if there are any questions or comments about this section.

Conclude the section by restating the key points.

Refer to overhead Partners Plus Successful Family and Caregiver Partnerships 3-3. Refer participants to Section 3 in the Partners Plus Family and Caregiver Manuals.

Does anyone have any questions or comments at this stage?

Families and caregivers can build successful respite relationships that grow into long-lasting partnerships. Use understanding and communication and these guidelines will become second nature in time:

- Appreciate differences.
- Be willing to learn from each other.
- Call on each other for help.
- Allow time to talk and share experiences.
- Finally, share a commitment to a child.

Overhead Partners Plus 3-3

Successful Partnerships
Successful partnerships are those in which families and caregivers:

- Appreciate differences
- Are willing to learn from each other
- Ask each other for help
- Allow time to talk and share experiences
- Share commitment to children
Section 3—Building Partnerships: Families and Caregivers Working Together
Suggested Overheads
The Challenges of Respite

Caregivers may stop giving respite if:

- respite is harder than they expected
- time demands or physical demands are too great
- they don't feel appreciated
- training is not adequate
- "burn out" occurs
Meeting the Challenge

- Understand the purpose for respite
- Be active in helping develop training
- Be clear about your terms
- Learn about working with children and families
- Get information and support from other caregivers
Successful Partnerships

are those in which families and caregivers:

• appreciate differences

• are willing to learn from each other

• ask each other for help

• allow time to talk and share experiences

• share commitment to children
Section 4—All Kids Like Cookies

Objectives

- Experience firsthand some of the challenges faced by children with special needs, and
- Identify the ways in which children with special needs adapt to their environments

Preparations

Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 4, “All Kids Like Cookies,” of the Family and Caregiver Manuals.

Prepare for the All Kids Like Cookies Activity, Parts 1 and 2. While it requires advance preparation, the All Kids Like Cookies Activity vividly demonstrates the difficulties that children with disabilities can experience. Families and caregivers who participated in the field testing of this curriculum overwhelmingly concluded that this was one of the best and most memorable training activities in which they had ever participated.

Determine the number of participants. Prepare enough materials for the group so that half will receive handout number 1, and the rest will receive handout number 2.

1. All Kids Like Cookies Handout
   Partners Plus Supplement 1-E (easy version)

2. All Kids Like Cookies Handout
   Partners Plus Supplement 1-H (hard version)

Prepare four stations for the All Kids Like Cookies Activity, Part 2. Gather the following materials for each station:

Station 1: Vision

- Directions (Photocopy the Partners Plus Supplement 2: Directions for Stations for All Kids Like Cookies Activity, Part 2)
- A table and a few chairs
- Safety glasses (available at hardware stores). Wrap the lenses of one pair in foil with a pinhole in each lens. Wrap the lenses of the second pair in several layers of saran wrap. Wrap the lenses of the third pair in wax paper.
Preparing for Training continued

Station 1: Vision (continued)
- Blindfolds
- Crayons
- Several children's books and magazines
- A cup
- A pitcher with water

Station 2: Speech
- Directions (Photocopy the Partners Plus Supplement 2: Directions for Stations for All Kids Like Cookies Activity, Part 2)
- A table and a few chairs
- A bag of marshmallows
- A package of gauze
- Paper towels or napkins
- A trash can

Station 3: Fine Motor
- Directions (Photocopy the Partners Plus Supplement 2: Directions for Stations for All Kids Like Cookies Activity, Part 2)
- A table and a few chairs
- Several pairs of thick gloves (gardening or work gloves)
- Masking tape
- Two puzzles for young children with big pieces and knobs
- Large beads or pasta that can be threaded onto a string
- Thick string to thread the beads or macaroni with holes
- Two or three small, manipulative toys, like a wind up truck

Station 4: Gross Motor
- Directions (Photocopy the Partners Plus Supplement 2: Directions for Stations for All Kids Like Cookies Activity, Part 2)
- 10 to 15 feet of cleared space where participants can walk
- Two large balls. Place the balls at one end of the walkway.
- Three-foot long pieces of thick string
- Kick-ball, large toy (stuffed animal or doll)
- Video tape

Prepare overhead transparencies of:
- Overhead Partners Plus All Kids Like Cookies 4-1
Introduction to All Kids Like Cookies

Introduce the section. The purpose of this section is for trainers to help participants understand some of the challenges for children with special needs, and how caregivers can help children with special needs in both their daily activities and group care settings.

Refer to overhead Partners Plus All Kids Like Cookies 4-1.

Section 4: All Kids Like Cookies, was adapted from:


In the last section, we spent time talking about the perspectives of families and caregivers. Children are the third members of respite relationships. “All Kids Like Cookies” is an activity designed to give you the opportunity to experience the challenges that children with special needs face.

Before we begin this activity, let’s read this message from the Council for Exceptional Children that sums up why this activity is important for families and caregivers alike:

“We cannot help until we understand. We cannot understand until we feel. We cannot feel until we become.”

—Council for Exceptional Children

Overhead Partners Plus 4-1

All Kids Like Cookies

“We cannot help until we understand.
We cannot understand until we feel.
We cannot feel until we become.”
All Kids Like Cookies Activity

Set the tone for the activity.

This activity is divided into two parts, and is a little more challenging than some of the others we've tried. The first part of the activity is done individually. You'll take an XYZ test. During this test, I cannot allow any talking and all eyes must stay on your own paper.

Distribute the XYZ Test alternately, giving half the participants the "easy" version and half the "hard" version. Use Partners Plus Supplementary Material PP 4 SM. Do not let the group know that you are distributing two different tests.

Place the handouts face down in front of the participants. Do not allow any talking during this time.

Do not turn over your papers until I tell you to, and remember, do not talk or ask questions.

Each of you has received a test. There are 16 items on the test. When you finish the test, put your pencil down and turn your paper over. You will have three minutes to complete the test. Begin.

There are no overheads for this section.
All Kids Like Cookies continued

Time participants for three minutes. Announce when time is up.

Have participants grade their own tests.

Again, do not allow questions or talking at this stage. Later, when you discuss the activity, participants will have an opportunity to discuss what they experienced during the test.

(The test was adapted from the Early Integration Training Project, 1991.)

Do not start a discussion.

Time is up, put your pencils down.

That wasn't so hard, right? I'll call out the correct answers and each of you can score your work.

1. D 9. D
2. C 10. C
3. A 11. C
4. C 12. D
5. C 13. A
7. D 15. B
8. A 16. D

How'd you do? Raise your hand if you got all 16 answers correct; 10 or more answers correct; 5 or more answers correct; fewer than 5 answers correct. We'll have an opportunity to discuss the test after the next activity.

There are no overheads for this section.
All Kids Like Cookies continued

Divide the participants into four equal groups. Use creative ways to divide the group, such as four different kinds of candy, gum, color pencils, etc.

You will work in small groups for Part 2. We're going to form small groups now . . .

There are no overheads for this section.

Explain Part 2 of the “All Kids Like Cookies” activity.

In a minute, we will ask you to go to an assigned station to begin Part 2 of the “All Kids Like Cookies” activity. Directions are provided at each station. Read and follow the instructions at each of the four activity stations.

You will have about 4 minutes at each station. At our signal, move to the next station.

Begin timing the first rotation of stations. Circulate among the groups to make sure that everyone understands what to do and has enough time to try all the activities.

After you have had a chance to visit all four stations, we will regroup to talk about what you experienced. Begin.

After 4 minutes, give a signal and have each group move to another station. Continue rotating until all of the groups have visited all four stations.
Discussion

Debrief both Part 1 and Part 2 of the “All Kids Like Cookies” activity.

Talk about the individual activity and then the small groups activities.

Allow a minute for participants to examine and react to the differences in their tests.

Okay! Let's gather together and talk about what we just experienced.

Let's start with the test. What were you thinking during the test? How did it make you feel? How about for those of you who did not do as well as you expected—how did you feel? Did anyone just give up?

Half the group was given a significantly harder test than the other half. Take a minute to look at the test of the person next to you.

Children with various disabilities affecting the way they think or solve problems often have trouble in school. They can feel frustrated and incompetent. Many children with these disabilities give up in school and need encouragement and support to continue.
All Kids Like Cookies continued

We know that children with mental retardation do learn. It may be at a different level or rate than others. They may need more repetition to remember information.

Children with learning disabilities have differences in how they learn. They may need to take tests orally. A variety of teaching techniques are used to teach all children. For example: some children learn basic math by feeling the shape of numbers made out of sandpaper or felt, other children may need to sort out objects to learn math ideas.

Let's talk about Part 2: the stations. At station 1, you were given activities that were done with a visual impairment. Were you able to see anything at all? How did you accomplish the activities? Did you have to modify your behavior in order to accomplish each task?

Move on to Part 2. Discuss how each station posed challenges for the group.

There are no overheads for this section.
All Kids Like Cookies continued

Children with visual impairments often hold their heads at an unusual angle to keep points of interest in their vision. Were all the tasks at the vision station appropriate for children with visual impairments? How might they have been adapted?

How about using plain paper with scented markers, rather than paper with pictures and lines to stay in? What about using large print books and magazines?

Remember, you may have to think about adapting activities for children with special needs. Use activity materials that are enjoyable enough through smell or feel, such as cooking or clay molding. Make sure activities give children a sense of accomplishment through a balance of challenge and success. Adapted activities are often fun for all children!

If participants do not offer enough illustrations for adopting the station activities, the trainer should give them.

There are no overheads for this section.
Discuss how station 2 posed challenges for the group.

Think about the speech activities at station 2. What was this experience like for you? How did you make yourselves understood?

In this activity many of you chose to keep your messages brief. Children with speech impairments often become frustrated when others do not understand them and will say as little as possible. It is easy to misinterpret silence as not having anything to say.

Did anyone use gestures? Children with speech impairments often use other parts of their bodies to communicate. One illustration of this is when a child with no speech and limited control of their body movements, uses his or her eyes intently to communicate.

Experiencing the activities at the speech and language station often makes us laugh at ourselves and others. Children with speech impairments do not laugh at the differences in the way they sound. Sometimes they stop talking rather than cope with the difficulty of getting people to understand them.
Discuss how station 3 posed challenges for the group.

In group care settings, children may ask about another child whose speech is different, "why does she talk funny?" Caregivers should model responses that are matter of fact and appropriate to a child's level of understanding. One response may be, "you are having trouble understanding Anna." Caregivers can also teach children specific ways to communicate that increase understanding and promote social interactions.

Next, let us talk about the fine motor activities at station 3. Did you feel awkward performing the group tasks with limited use of your fingers and hands? How did you adapt to accomplish the tasks? Were all the activities appropriate?

In order for children to fully participate and learn about the things in their environment they need to handle objects. A child with fine motor impairment may need to have a knob glued to puzzle pieces. When an older child's ability to solve complex puzzles is affected by fine motor impairments adaptations are needed.

At station 3, some of you may have been too frustrated to have enjoyed stringing objects because the holes were too small and the string too limp. An appropriately planned activity would have included objects with larger holes and stiffer string.
Discuss how station 4 posed challenges for the group.

At station 4, the tasks you did require using the large muscles of your body. What was your experience during this activity? How did you adapt?

Children with motor impairments that affect their legs, feet, and arms may look different when they move their bodies. During the activity at station 4, participants often remain in an upright position while walking toward the toys. Sometimes participants will roll on the floor or move on their knees to get to the toys.

A child with motor impairments affecting their large muscles, may need to use more energy and effort to get around. Some children may use special equipment to help them, such as walkers and wheelchairs. Clear pathways are important to mobility. Generally, eliminating physical barriers that limit children's participation is as simple as rearranging furnishings or play areas.

There are no overheads for this section.
Summary

Conclude the section by summarizing key points.

Are there any other questions or comments you’d like to add about All Kids Like Cookies?

Today, we have had the opportunity to peer inside the lives of children with special needs. We must remember, however, that the disabilities we experienced were temporary. Children with special needs and their families deal with these challenges on a daily basis. During the activities in ALL KIDS some of you looked different in the ways you moved your bodies, used your voices, or behaved.

It may be necessary for caregivers in group settings to help children feel comfortable with the way we look, sound, or behave when involved in the same activities. Caregivers can model accepting attitudes that celebrate differences.

There are no overheads for this section.
All Kids Like Cookies continued

Sometimes directors and staff of recreational programs think that they will have to make extensive changes to include children with disabilities. Often, they only need to make a few accommodations to meet the needs of children with special needs. Valuable respite to families who have children with disabilities can be offered by providing the same kinds of appropriate experiences that characterize quality programs for all children. Leaders, directors, and counselors of community children's and youth programs can provide valuable respite to families who have children with disabilities.

It is important to note how children adapt to what is asked of them. While many of you were frustrated with the activities today, you managed to accomplish what we asked you to do. Children with special needs have incredible strengths. As you work with them, notice their strengths and build on them.

At this point, we're going to take a break for lunch. We'll meet back here at . . .

If One Day Workshop:

The end of this session provides a natural breaking point for a one-day workshop. If you are presenting a full day workshop, we recommend breaking for lunch after this activity.

There are no overheads for this section.
Section 4—All Kids Like Cookies
Suggested Overheads
All Kids Like Cookies

"We cannot help until we understand.
We cannot understand until we feel.
We cannot feel until we become."

— Council for Exceptional Children
Section 4—All Kids Like Cookies
Supplemental Materials
All Kids Like Cookies Activity

Part 1: XYZ Test

Directions: Circle the best answer for each question.

1. The following is a color:
   a. fish
   b. car
   c. stove
   d. blue

2. 6 x 2 =
   a. 27
   b. 97
   c. 12
   d. 67

3. How many inches are in a foot:
   a. 12
   b. 3
   c. 36
   d. 18

4. The following is not a planet:
   a. Mars
   b. Jupiter
   c. Hamilton
   d. Saturn

5. When riding a bicycle at night, you should
   a. wear dark clothing and sunglasses.
   b. ride on the other side of the road, opposite the side for day riding.
   c. wear a white shirt or coat, and turn on head lamp and tail lamp.
   d. not ride at night.

6. The following state is not located in the East:
   a. Maine
   b. Florida
   c. Virginia
   d. California

7. A refrigerator is used to
   a. wash clothes.
   b. cook food.
   c. cut food.
   d. cool food.

8. The following is a bird:
   a. robin
   b. deer
   c. cat
   d. monkey

9. The following is not a form of transportation:
   a. bus
   b. train
   c. automobile
   d. football

10. The following is a winter month:
    a. May
    b. July
    c. December
    d. September
11. The following is an even number:
   a. 1
   b. 11
   c. 6
   d. 15

12. A dentist would
   a. deliver a baby.
   b. prescribe glasses.
   c. perform a heart transplant.
   d. pull a tooth.

13. Milk could be purchased at
   a. a grocery store.
   b. a hardware store.
   c. a florist.
   d. a clothing store.

14. The following would be used to mend clothes:
   a. a nail
   b. a staple
   c. a tack
   d. thread

15. The following would be worn for snow skiing:
   a. wing tips
   b. boots
   c. sneakers
   d. sandals

16. The following is used to brush teeth:
   a. a shovel
   b. a hose
   c. a wrench
   d. a brush

Adapted from:

All Kids Like Cookies Activity

Part 1: XYZ Test

Directions: Circle the best answer for each question.

1. The following is a color:
   a. fish
   b. car
   c. stove
   d. blue

2. $6 \times 2 =$
   a. 27
   b. 97
   c. 12
   d. 67

3. How many inches are in a foot:
   a. 12
   b. 3
   c. 36
   d. 18

4. The following is not a planet:
   a. Mars
   b. Jupiter
   c. Hamilton
   d. Saturn

5. When riding a bicycle at night, you should
   a. wear dark clothing and sunglasses.
   b. ride on the other side of the road, opposite the side for day riding.
   c. wear a white shirt or coat, and turn on head lamp and tail lamp.
   d. not ride at night.

6. The term Dorian refers to
   a. a Greek column style.
   b. a shade of gray.
   c. a psychoanalytic procedure.
   d. a musical style prevalent in the Middle Ages.

7. A chain has 5 links. The first link can hold 7 pounds. The second can hold 9 pounds. The third can hold 4 pounds. The fourth can hold 8 pounds. The last can hold 7 pounds. How many pounds can the whole chain hold?
   a. 9 pounds
   b. 7 pounds
   c. 8 pounds
   d. 4 pounds
   e. 35 pounds

8. The henry is
   a. a unit of measurement of inductance.
   b. a unit of resistance.
   c. a volt equivalent.
   d. a chemical substance.

9. Alnico is
   a. a skin disease.
   b. a type of mental illness.
   c. an engineering term.
   d. a precious metal.
   e. an alloy used for magnets.
10. The term joule refers to
   a. a musical instrument.
   b. tempo.
   c. a unit of energy.
   d. a carpenter's tool.
   e. a type of electrical wire.

15. A dart is
   a. a kitchen design.
   b. a flat tool used in plastering.
   c. a type of saw.
   d. a type of linen.
   e. a fishing lure.

11. Kirchoff is well known for his contributions to the field of
   a. music.
   b. psychology.
   c. electronics.
   d. economics.
   e. education.

16. The word pentatonic refers to
   a. five states of emotional behavior.
   b. five types of antidepressant medicine.
   c. a play by Sophocles.
   d. a five-tone scale of music.

12. A French seam is
   a. a machine stitch.
   b. a seam within a seam.
   c. a type of wood lamination.
   d. a seam sewn with a double needle.

13. Muntin is used in
   a. The production of a window frame.
   b. food production (cooking).
   c. farming.
   d. tuning a piano.

14. The term puzzolano refers to
   a. music tempo.
   b. a difficult problem.
   c. a food ingredient.
   d. volcanic dust.

Adapted from:
All Kids Like Cookies Activity

Directions for
Station #1—Vision

Put on a pair of special glasses or blindfold and try to do the following activities. Take turns using different pairs of glasses.

1. Color a picture and try to stay in the lines.

2. Look at the book.

3. Pour a cup of water.

4. Complete the crossword puzzle.
All Kids Like Cookies Activity

Directions for Station #2—Speech

1. Have one person play the role of the child.

2. The child puts 2-3 marshmallows or some gauze in his/her mouth, on his/her tongue.

3. The child asks other group members a question, or asks them to get something.

4. Switch roles so that everybody has the chance to be the child.
All Kids Like Cookies Activity

Directions for
Station #3—Fine Motor

Put on a pair of gloves or tape your fingers together. Try to do the following activities.

1. Put a puzzle together.

2. String beads.

3. Play with a toy.

4. Take off and put back on your watch, earrings, bracelet, etc.
All Kids Like Cookies Activity

Directions for
Station #4—Gross Motor

1. Put a ball between your ankles.

2. Clasp your hands and keep them together during the activity.

3. Use any method (except hopping) to reach the toy.

4. With hands still clasped, pick up the toy or video and give it to another group member.
Introduction to Separate Family and Caregiver Sessions

Begin by explaining what families and caregivers will be doing.

During the next several sections, families and caregivers will be learning different things. Families will be learning more about finding, screening, interviewing and training caregivers for their children. Caregivers will be learning about special needs, child development, common disabilities, health needs, and becoming Partners caregivers.

Identify which trainer will work with each group and where they will meet.

Families and caregivers will work as separate groups with a trainer who will cover specific topics in your manuals. At the end of these separate sessions, you will reunite for a final activity and some closing thoughts.

Any questions?
Section 5 Families—Finding Respite Caregivers

Objectives
- Develop strategies for finding appropriate respite caregivers
- List qualities/criteria for selecting caregivers
- Identify potential caregivers in their natural support networks
- Know other strategies for finding respite caregivers

Preparations
Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 5, “Finding Respite Caregivers,” in the Family Manual.

In this section, families get information on matching with caregivers through the Partners program. Each community decides how or if they distribute information on trained caregivers to families in the community. Please prepare for this section by becoming familiar with these procedures and developing necessary materials to communicate them to families.

Families in training often are a great resource for one another. This section includes extra time to allow for discussion and information sharing.

Overheads are not necessary for Section 5 in the Family Break-Out Session. If you prefer, you may create overheads for your own use.

Corresponding Sections
Introduction

Families need to think ahead when they are finding and training caregivers. In order to make sure that everyone in the partnership is comfortable, families and caregivers must commit time and energy. This means that you want to start looking for a respite caregiver NOW, before you reach a point where respite is an emergency need.

The Partners model is based on the idea that families want to find and train their own caregivers. It can be difficult to find appropriate people to care for your children. Today, we will give you a few strategies to help you find caregivers who will be helpful to you.
Important Qualities for Your Respite Caregiver

Before you begin looking for a caregiver, it is helpful to think about what qualities you feel are important in a caregiver for your child.

Consider this question: What do I want in a respite caregiver? If you were to describe the perfect caregiver for your child, what qualities would that person need? Would she need to be physically strong to lift your child? Would she need a car to get to your home? Would he need to be gentle? Is it more important that she be creative and fun-loving?

This question requires some thought and consideration. Right now, let's take a minute to think about these important qualities.
Important Qualities for Your Respite Caregiver continued

Give family members one minute to consider this question. At the end of the minute, ask:

You will want to take more time with this later. For right now, what did you come up with?

Give everyone the opportunity to share at least one quality. After everyone has done this, conclude this section.

It is very helpful to think about these qualities before you begin your search for a caregiver for two reasons.

First, these qualities are quick checks for potential caregivers. They help you screen caregivers quickly and easily. If I need someone with a car, who can travel to my area, I can save time by asking potential caregivers whether they have use of a car in my first conversation with them. If they do not have a car, then I do not need to interview them.

Another way to find out if a caregiver has some of the qualities you are looking for is to use the Caregiver Profile in your Family Manual or ask a caregiver for their resume.
Important Qualities for Your Respite Caregiver continued

Make sure that families understand this concept of screening. You may need to give some more examples of screening questions, referring to the qualities families listed in their activity.

This process of quickly excluding unlikely matches is called “screening.” It saves a lot of time for everyone. For example, if a family said that it was important that their caregiver had prior experience with children with autism, they should be sure to ask whether caregivers have that experience in their first conversation with them.

This list of qualities is also important in thinking about where to find appropriate caregivers for your child.
Circles of Family and Friends Activity

Introduce this activity by referring families to Circles of Family and Friends in Section 5 of the Family Manual.

Explain the information listed.

While other options are available to you, it is often best to match with a caregiver who already knows your family. These are people who already have a relationship with you and your child. You may already know someone who is a good match for you. Sometimes, these people are not obvious. But as you think about caregivers, think about people you know who have offered to help, or who have expressed an interest in your child. Maybe there is someone you have considered asking but did not because they did not have experience or special training to care for children with special needs. After you complete today's workshop, you will know how to train caregivers for your child.

Sometimes, it is difficult to think about who those people in your circle of family and friends might be. The circle chart in your Family Manual is designed to help you.

There are no overheads for this section.
Circles of Family and Friends Activity continued

Give families two minutes to brainstorm ideas from their circles of family and friends.

Encourage them to keep an open mind and think particularly about people who have asked how they can help.

At the end of the two minutes, ask them how they did. Encourage participants to share ideas to conclude the activity.

Think about yourself in the center of the center circle, where it says “Family.” All around that circle are other circles, circles of family and friends. Fill in the lines by thinking about people in those circles who might be interested in providing respite.

As you do this, stretch your imagination. Who can you think of who has been friendly to you and your child? Who has your child already developed a relationship with?

Don’t reject any idea until you are sure it will not work for you. I will give you a few minutes to fill in some of the lines in the circles.

Let’s see how we did. Who would like to share what they learned?

There are no overheads for this section.
Strategies for Finding Respite Caregivers

**Recruiting from the Community**

Refer families to Resources in Your Community in the Family Manual. Explain the information listed.

There are many agencies in this community that employ people who might be wonderful caregivers for your children. You can call these agencies and ask if they know of anyone on staff or in the community who might be interested in providing respite. They might take your name and have potential caregivers call you back.

Another way to access caregivers through these agencies is to ask if you can post flyers there. Make a colorful poster that describes your child's strengths and your family's desire to find a capable caregiver for your child. Remember, it is important to get permission to post flyers!

Think about your list of important qualities as you decide who to call and where to post information. If you need an energetic caregiver, college campuses are a good place to start. If, on the other hand, you need someone with a lot of experience in child care, you might start by asking at some local child care centers.
Strategies for Finding Respite Caregivers continued

The Partners Directory, If Available

Note: Many communities choose to create a directory of Partners caregivers to distribute to families looking for caregivers. If this is true of your community, use this time to describe this directory to families.

The Partners program has developed a directory of potential caregivers for families to access. But please note that Partners is not an employing agency. It does not license caregivers. In order to be included in the directory, caregivers need to...

Families can get the following kinds of information from the directory...

If the local Partners program fosters family/caregiver partnerships by planning matching events, be sure to mention it.

One of the best opportunities for finding a respite caregiver is by talking to some of the other participants in the session today. Families can exchange information and phone numbers if they find someone they think would be a suitable match.

There are no overheads for this section.
Summary

Conclude the section by asking for group feedback.

Once you have a few ideas, you have the beginning of a potential list of caregivers. The next step is to talk to them about respite and to ask them if they would be interested in becoming a caregiver.

Are there any questions or comments on what we've discussed in this section? Would anyone like to share some additional ideas for finding caregivers that we haven't covered?
Section 6 Families—Interviewing Caregivers

Objectives
- Prepare families to interview potential caregivers
- Identify what questions to ask caregivers in interviews
- Understand different options for conducting interviews
- Know other strategies for gathering information on caregivers

Preparations
Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 6, “Interviewing Caregivers,” in the Family Manual.

Overheads are not necessary for Section 6 in the Family Break-Out Session. If you prefer, you may create overheads for your own use.

Corresponding Sections
Please refer participants to Section 6, “Interviewing Caregivers,” in the Family Manual.
Introduction to Interviewing Respite Caregivers

Begin by explaining the interview process.

After finding a potential caregiver, and learning some preliminary information about her, the next step is the interview. You want to collect as much information as possible about a caregiver before you begin respite with her. In this section, we will talk about the different ways to interview a caregiver, provide you with some sample interview questions, and give you a few tips on getting the most information from a caregiver in an interview.

There are no overheads for this section.
Conducting Interviews

Describe how to conduct interviews.

It is important to make interviews comfortable for both you and the caregiver. You can conduct an interview in a number of different ways. Some families choose to interview in their own homes. Some prefer to interview in the homes of the caregivers they are working with. Some like to work over the telephone. It is often helpful to see the caregiver in the environment where he will be working in respite.

We suggest that families interview more than one person and try and find as many caregivers as possible. In this way, you will have more people to call on when you need respite.

There are no overheads for this section.
Conducting Interviews continued

Describe how to conduct interviews. Also decide whether you want your children to be present for these interviews. Often, families find it useful to watch caregivers interact with their children. If your child's presence would be distracting, however, you may choose to hold a two-part interview. In the first part, you can sit down with a potential caregiver without your child present. At that time, the caregiver can answer any questions you have for her. If you feel the interview is going well, you can ask this caregiver to return so that she can meet your child. At that time, watch how she interacts with your child. Remember, she will not have the personalized training that she will need to work independently in respite. But this should give you an idea of how she works with your child on a natural level.

There are no overheads for this section.
Remind families that caregivers may have some questions as well.

Interviews are an important time for caregivers to get information on your family as well. Be prepared to answer questions that they might have about your children. Be very honest with caregivers about what will be expected of them. Emphasize your children's talents and strengths while, at the same time, being open about their special needs. This is the time when you and the caregiver decide whether you want to work together. By being honest, you will be sure that you both are committed to this respite relationship.

Define a successful interview. Refer families to "Interviewing Potential Caregivers" in their manuals.

Successful interviews are those where families and caregivers learn what they need to know from each other in order to decide whether or not they are ready for a respite relationship.
Conducting Interviews continued

Tips for Successful Interviewing

Refer participants to the Caregiver Profile in Section 9, “Forms for Sharing Information” in the Family Manual.

Partners has created a form for gathering relevant information about a potential caregiver, called the Caregiver Profile. If you find it useful, make copies for potential caregivers to complete before conducting an interview.

Here are some tips for making sure that your interviews are successful.

- Be prepared.
  You may ask your applicants to give you a resume of their experiences with children or a Caregiver Profile before they come for an interview.

  Make notes as you review this information.
  What else do you need to know?

  You may choose to use some or all of these questions in your interviews with caregivers.
  You may want to add some questions that are more specific to your child and family.

Refer families to “Sample Interview Questions” in their manuals.

There are no overheads for this section.
Conducting Interviews continued

Continue discussing successful interview tips.

- Decide how to conduct the interview.
  Again, talk with your caregiver about where and how to conduct the interview. Make sure that you schedule it at a time and place that are convenient for everyone. Decide before the interview who will be there. Some families like for both parents and the caregiver to have time alone to get to know each other. Sometimes the whole family is present for all or part of the interview. It is a courtesy to the caregiver to let them know how long the interview may take and who will be there.

- Break the ice.
  Sometimes, it is difficult to know where to begin. It is useful to begin by asking the applicant to tell you about his experience with children.

There are no overheads for this section.
Conducting Interviews continued

Continue discussing successful interview tips.

• Ask open-ended questions.

An open-ended question is a question that is not answered with a yes or a no. They ask the applicant to provide more information. For example, “Do you like children?” is not a question that allows the applicant to give you a lot of information. “What do you like to do with children?” allows caregivers to expand on their skills.

• Take notes.

We suggest that families take notes or tape their conversations with caregivers. In this way, you can go back and ask follow-up questions to some of their statements. In addition, you can use those notes as you consider the applicants at a later time.

There are no overheads for this section.
Conducting Interviews continued

Continue discussing successful interview tips.

- Reference checks.
  It is a good idea to ask potential caregivers for references. These are people who know the applicant, and can comment on their character or their work with children.
  When you call references, tell them who you are, and that you are considering hiring the applicant to provide care for your child. Ask if they have a few moments to speak with you. If they do, ask these questions, and any additional questions that you have about the applicant.

Refer families to Section 9, "Forms for Sharing Information." A form for checking caregiver references is included in this section.
Conducting Interviews continued

Continue discussing successful interview tips.

- Criminal record checks.

We would like to think that every applicant for child care positions are good people who love children. Unfortunately, some people want to work with children for the wrong reasons. There are some things that families can do to protect their children. The state police should be able to check the criminal records of potential caregivers. You will need to get forms from the police and ask applicants to sign them. Usually, there is a fee for this service, but a clean criminal record check may give you peace of mind.

Note: Some communities may be able to help families by sponsoring criminal record checks on their caregivers. If your program is able to do this, share that information with families now.

There are no overheads for this section.
Conducting Interviews continued


This section in your manual gives you information on some things to look for when you are interviewing caregivers. Included on these pages are some warning signs that may indicate a caregiver is not safe or appropriate for your child. Information about what to ask and look for in home and community-based caregivers is included. I encourage you to read this section carefully, and to use some of the questions listed here as you interview caregivers.

The most important tools that you have for protecting your children are your instincts. If you are uncomfortable with an applicant for any reason, that person is not the right caregiver for your family.

Explain how to conclude the interview.

After you have finished the interview, be sure to follow-up. Thank applicants for their time, and let them know how and when you will make your decisions. Do not leave an
Conducting Interviews continued

It's important to be able to say "no" to caregivers who are not a good fit for your family.

It can be very difficult to tell an applicant "no." However, it is important to be honest and direct.

This can be as simple as saying, "We interviewed several people, and we found someone else who we think best meets our needs."
Summary

Ask for group feedback.

Are there any questions or comments on what we've discussed in this section? I encourage you to take the time to read through all of the information included in this section in the manual.

Conclude the section by restating key points.

Feel free to write a note if you find saying “no” on paper is easier than saying it over the phone.

Interviewing is an important step in the Partners process. When you find a caregiver you feel comfortable with, offer her the job. After you agree to work together, get a clear idea of your caregiver's schedule and when and how she can and cannot provide care. You started to build your relationship in the interview.

Now, build on that relationship by making sure that she has the opportunity to set limits and schedule her time. The next step is training. When interviewing community program staff be sure to find out if they are willing to work with you in learning how to include your child in their program. The next step is to make a plan for how you will work together.
Section 7 Families—Developing a Personalized Training Plan

Objectives
- Be prepared for personalized training with caregivers.
- Understand the four-step process for teaching daily routines to caregivers.
- Know some principles for teaching adults.
- Understand the importance of sharing emergency procedures and household rules with caregivers.

Preparations
Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 7, “Developing Your Personalized Training Plan,” in the Family Manual.

Overheads are not necessary for Section 7 in the Family Break-Out Session. If you prefer, you may create overheads for your own use.

Each community decides if they want to keep a resource library for caregivers and families. In this section, families get information on the Partners resource library. If your community has one, prepare to share information and material on accessing those resources.

Corresponding Sections
Introduction to Developing a Personalized Training Plan

The next step in the respite process is to teach your caregiver about your child and your family and your daily routines. In the next section, we will discuss a four-step process for developing a personalized training plan for your caregiver. We will also go over some tips on training adults.

Introduce the personalized training plan concept.
Four Step Process for Developing a personalized Training Plan

Introduce the four step process.

You can begin developing your plan by thinking about the time of day or time during the week you want care for your children.

Some families like to have respite in the evenings. Some families particularly want a Friday or Saturday night to go out. Some families need someone during the day during the week or on the weekends. When you are thinking about training a caregiver, be as specific as you can about when you want respite.

There are no overheads for this section.
Process for Developing a Personalized Training Plan continued

Ask for an example of a specific time from the group, and help participants think about all of the routines necessary for that time.

For any period of time, there are a number of routines that occur. Think about what happens in your family during the time you are planning respite. If respite takes place in a before- or after-school program, think about what happens during that time.

Will a caregiver need to learn a feeding routine? A play routine? Will a caregiver need to learn how to help your child interact with other children? Will a caregiver need to know how to adapt games in order for your child to fully participate? What about a special bedtime routine? A wake-up routine? A bathroom routine? A bath routine? A cleanup routine? A medication routine? Think about everything a caregiver will need to know for this time with your child.

We suggest that you use this four-step process to train your caregiver. We will go through all of the steps.

Refer families to “Step by Step: A Guide to Developing Your Personalized Training Plan” in their manuals.
Step 1: Identify Training Goals

The first step is to identify training goals. What do you want your caregiver to learn to be able to provide care during this time? What are the routines that they need to learn?

One goal might be that you want your caregiver to know how to feed your child. Another goal might be that you want your caregiver to know how to help your child participate in an activity with other children. You can have as many goals as you need for personalized training.

There is a worksheet in Section 7 of the Family Manual for you to write down your goals for personalized training for a particular time of day.

Partners Plus Trainer’s Workshop Manual
Child Development Resources, P.O. Box 280, Norge, Virginia 23127

There are no overheads for this section.
Process for Developing a Personalized Training Plan continued

**Step 2: List Training Methods**

Refer families to “Step 2: List Training Methods” in their manuals.

Define each of the methods listed.

The next step is to list training methods for each goal. What activities will you use to help the caregiver achieve the goal? There are many options.

**Modeling:** A family shows a caregiver what to do.

**Videotaping:** A family videotapes a routine, enabling the caregiver to watch it again and again.

**Shadowing:** A caregiver helps the family work with their child. In this way, she gains hands-on experience but is not alone.

**Talking, Telling, Describing:** These are all ways that a family gives a caregiver information verbally.

There are no overheads for this section.
Define each of the methods listed. | Writing: A family shares information with caregivers on paper.
|---|

Be sure to emphasize the following points. | Answering Caregiver Questions: A family responds to a caregiver's questions.
|---|

When teaching a caregiver your routines, try and remember how it was that you learned them. You may want to use those same methods to train your caregiver.

Remember that different people learn best in different ways. In the Caregiver Break-out Session, we ask caregivers to think about the ways that they learn best. Is it easier for them to hear information or to read it? Do they understand quicker if they watch someone perform a task or if it is described to them? Ask caregivers to share their learning styles with you, and use that information as you plan your training.
Process for Developing a Personalized Training Plan continued

Be sure to emphasize the following points.

Giving people information in a lot of different ways is usually more effective than giving them information in only one way. As you develop your plan, think about including more than one training method for each goal.

There is a worksheet in Section 7 of the Family Manual to help you think about what methods would be effective in teaching caregivers each goal.

Refer families to “Step 3: Use Resources to Help Train Your Caregiver” in their manuals.

The third step is to use resources to help train your caregiver. Resources are things that your caregiver can read, watch or look at that will give her extra information on working with your child and family. There are many different resources that you can use.

If your child is in a before- or after-school program that takes place in the same building where he attends classes, a teacher may be a helpful resource to program staff.

There are no overheads for this section.
Process for Developing a Personalized Training Plan continued

Refer families to "Sharing Information with Caregivers" in their manuals. Flip through the section with families as you make the following points.

You can create resources for your caregiver. The forms in this section are designed to help you think about the information your caregiver might want or need during respite. They address health, safety and medication issues for your child, information about your child's personality and skills, and other family information.

It is up to you whether you decide to use these forms. However, remember that caregivers will need all of this information.

The health and safety information is particularly important to caregivers. They need to know how to handle an emergency in your home. The list of phone numbers in your manuals should be as complete as possible. Cover all of the topics on the health and safety checklist. Do not place your child in the care of a caregiver who has not been prepared with this information!

There are no overheads for this section.
Process for Developing a Personalized Training Plan continued

Flip through “Sharing Information with Caregivers” as you make the following points.

There is a medical treatment release form in your manuals. This gives your caregiver permission to seek medical attention for your child in an emergency. Some medical care facilities will not treat children without the permission of a parent or legal guardian. This form is crucial to your child in an emergency. Keep it with other emergency information in a place where caregivers know to find it.

Feel free to create your own forms and documents as well. You may want to give your caregiver a diagram of how to position your child, for example. Or it might be important for them to have a map of the neighborhood. You might give them a videotape of your child’s bedtime routine. Any of these resources can be valuable training tools.

There are no overheads for this section.
Process for Developing a Personalized Training Plan continued

Flip through “Sharing Information with Caregivers” as you make the following points.

There are many other resources that others have developed that can help you train your caregivers. There may even be some that others have developed specifically for your child. Think about sharing diagnostic reports, personalized plans, or suggested activities from your child’s therapists with caregivers.

You might also want to share general information on your child’s condition. You may have books, magazines, or brochures in your home that could help a caregiver have a better understanding of your child.

If you feel like some of this general information would be useful to caregivers, but you do not have it in your home, there is a list of places where you or your caregiver can find resources on this page.

Step 3: Use Resources To Help Train Your Caregiver

Refer families to “Step 3: Use Resources to Help Train Your Caregiver.”
Process for Developing a Personalized Training Plan continued

Note: In some communities, the Partners program has a library of resources that are available to families and caregivers. If your community has such a library, describe it now.

Describe the resources worksheet.

**Step 4: Evaluation**

Refer families to “Step 4: Evaluation” in their manuals.

Our program has a library of resources available for your use. To take advantage of this service . . .

In your manuals there is a worksheet to help you think about the resources you will use in your personalized training plan.

The final step of your training plan is evaluation. How will you and your caregiver know that the training goal has been accomplished and that the parents, caregivers, and children feel comfortable? There are many different ways to evaluate your caregiver’s skills and knowledge. You might choose to ask your caregiver questions and make sure that she answers correctly. Another way is to ask a caregiver to show you a skill that you have taught him. You might just watch your child and caregiver for a time. Do they look comfortable with each other?

There are no overheads for this section.
Describe Step 4.

Be sure to ask your caregiver to evaluate his knowledge and skill. He might want you to watch him do a task a few more times, or he might have some more questions to ask you. Make sure that he is comfortable with all of his responsibilities before you leave for respite.

In your manuals there is a worksheet to help you plan your evaluation for your training.

Refer families to “Sample Personalized Training Plan” in their manuals.

On the following page there is a sample personalized training plan for lunchtime respite. Take a moment to read it now.

Give families time to read through the plan. Then ask for feedback.

Are there any questions about the four step process?

Later today, we will practice developing a personalized training plan using these four steps with the caregivers in the group.
Process for Developing a Personalized Training Plan continued

Ask if there is a person in the group who would be willing to volunteer to use one of their child's routines as an example for that activity.

If there are no volunteers, you can use an experience with a child you know as an example.

Is there a person in the group who would be willing to volunteer to use one of their child's routines as an example for that activity? This is a great way for an individual family to begin to develop a plan. Some of the caregivers in the group might want to work with your child once they know a little about him or her.

There are no overheads for this section.
Training Tips

Introduce this section.

Training adults is a slightly different experience than teaching children. This section in your manual also includes some tips for training adults.

Remind families that caregivers should have input in personalized training plans, as well.

Caregivers may want to be involved in planning personalized training. They may have specific ideas about what activities or resources will be helpful to them. Invite them to share this information and their other ideas.

Refer families to “Training Tips” in their manuals.

Turn to Section 7 in your manuals for some training tips. The tips on this page will help make your training effective.

There are no overheads for this section.
Training Tips continued

Review training tips with families.

- Be clear with your instructions.
  Be certain that a caregiver understands the words that you use to describe your child, her condition, and household equipment. If necessary, show caregivers things that might cause them confusion.

- Be very specific.
  When you are training, think about the details of your care of your child. It may be that your child wears a bib for meals. Which bib is her favorite? Which bib does she never wear? Give this information to caregivers. Also, be careful that you do not assume that a caregiver knows where you keep things. Show her exactly where equipment and supplies are.

There are no overheads for this section.
Training Tips continued

Review training tips with families.

- Use your knowledge and resources to train.

  Again, think about how you learned about your child. Adapt these methods to your style of teaching.

- Give your caregiver written information.

  Written information is helpful because it can be taken home and studied and referred to at a later time.

- Talk to your caregivers about the information that you give them.

  A caregiver may have additional questions and comments about what he has read or heard from you.

- Make sure that he has the opportunity to learn more from you. Check to make sure that he is feeling comfortable with new skills and tasks.

There are no overheads for this section.
Review training tips with families.

- Use diagrams or pictures.
  Some people learn better from pictures. Include these in your training resources if they are appropriate.

- Be sure to get feedback from your caregiver.
  Find out what is helpful and not helpful to caregivers in training.

- Write down important facts or information.
  Again, this information should be easy for caregivers to study and come back to at a later time.

- Share information about your culture, or life ways (how your customs reflect your culture and your values), that is important to your family.
  This information will give a caregiver guidelines for working with your child and planning activities.

There are no overheads for this section.
Summary

Ask the group for feedback.

Are there any questions or comments about this section?

Prepare the group for the next step.

The next step is to use your personalized training plan to teach your caregiver about your child and your family. With its use, your caregiver should be well prepared to care for your child.

Remember that the kind of communication that occurs during training should continue throughout your relationship. Share new information with your caregiver. Caregivers may even have information to share with you! Personalized training should never end. Part of sharing a partnership is continually learning more about one another and your child.
Section 5 Caregivers—Disabilities Awareness

Objectives
- Identify terms, services, and issues related to children with special needs
- Understand common terms, services, and major issues related to children with special needs.

Preparations
Read the Introduction to this manual, the “Overview of the Section” and the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 5, “Disabilities Awareness,” of the Caregiver Manual.

The information on services and legislation in this section is accurate as of the day of printing. Please be aware that legislation and the provision of services for children with special needs are ever-changing. Please keep your materials up-to-date for families and caregivers.

Prepare overhead transparencies of:
- Overhead Partners Plus Caregiver IDEA (Individuals with Disabilities Education Act) 5-1
- Overhead Partners Plus Caregiver Plan 5-2
- Overhead Partners Plus Caregiver IFSP / IEP 5-3
- Overhead Partners Plus Caregiver Early Intervention and Public School Services 5-4
- Overhead Partners Plus Caregiver Common Disability Issues 5-5

Corresponding Sections
Please refer participants to Section 5, “Disabilities Awareness,” in the Caregiver Manual.
Introduction to Disabilities Awareness

Introduce the section. Refer caregivers to Section 5, “Disabilities Awareness” in the Caregiver Manual.

As you work with children and families, it can be very helpful to have an understanding of some of the terms, issues, and services for people with special needs.

Every day with a child with special needs is a day of learning: about the child, about how to help her in her development, and about how to get the services available to her. Often, when you hear a family of a child with a disability speak about these experiences, it is like you are hearing a whole new language. Families had to learn this language, too, and it is a tool that they use to communicate about their children.

You can think about this language as a kind of alphabet soup. Before you learn what the terms stand for, they look and sound like a jumble of letters. As we go through some of these terms, think about what it feels like for families new to the experience of raising children with special needs to be faced with all of this confusing language! They have to learn fast!

There are no overheads for this section.
Services for Children with Special Needs

Introduce this section.
Refer to overhead Partners Plus Caregiver IDEA 5-1. Participants may wish to follow along in Section 5 of the Caregiver Manual.

Many of these terms are related to intervention and education for children with special needs.

The first term, "IDEA" is an acronym that stands for the Individuals with Disabilities Education Act. IDEA is a federal law that guarantees that children with special needs receive services. It is broken down into several parts, which guarantee special services for children at different ages.

One part, Part C, says that infants and toddlers from birth to age two with disabilities or developmental delays may receive early intervention services. I will describe these in more detail in a moment.
Services for Children with Special Needs continued

Another part, Part B, guarantees eligible children from three to 21 years free and appropriate public educations (FAPE). Children from three to five years old receive early childhood special education (ECSE) services. The services provided to older children are simply called special education services.

It is important to remember that each state has different rules about eligibility for services.

IDEA also says that children who are eligible for services must have a written plan that is reviewed at least once a year.

There are no overheads for this section.
Services for Children with Special Needs continued

Refer to overhead Partners Plus Caregiver Plan 5-2. This information is in Section 5 of the Caregiver Manual.

The plan is written by a team that includes family members and other people working with a child. Teams can include teachers and special educators, physical, occupational and/or speech therapists, doctors or nurses, and counselors or social workers. The plan that the team writes includes information about the child's development, specific goals or outcomes for the child, and the activities that the team members will use to help the child reach the goals. Family members are crucial members of these teams, and must agree to the plan written for their child.

Overhead Partners Plus Caregiver Plan 5-2

Plan

Plan includes:
- Information about the child's development
- Goals for each child
- Activities to help the child achieve the goals
Services for Children with Special Needs continued

Refer to overhead Partners Plus Caregiver IFSP/IEP 5-3.

In early intervention services for young children, the plan is called an IFSP: an Individualized Family Services Plan. In services for children beyond age three, the plan is called the IEP, or Individualized Education Program.

Reviewing an IFSP or IEP can be very helpful in planning activities for respite. Keep in mind, however, that the information included in these plans is personal and confidential, and families might prefer that you not read them.

While early intervention and special education are all parts of IDEA, and they provide many of the same services, the programs often look very different.

Appropriate services for each child are determined by the members of their planning teams. Services for all IDEA programs can include home visits, classroom programs or play groups, and therapies.

Overhead Partners Plus Caregiver IFSP/IEP 5-3

IFSP
Individualized Family Service Plan:
• For early intervention
• Serving the child within the context of the family

IEP
Individualized Education Plan:
• For preschool service and beyond
• Serving the child within the context of the school system
Services for Children with Special Needs continued

Refer to overhead Partners Plus Caregiver Early Intervention and Public School Services 5-4.

Early intervention is a family-centered program, like Partners. As a result, IFSPs, the plans written for children in early intervention, can include many different kinds of services. Family support groups are one of the services offered by many early intervention programs.

In ECSE and special education programs, however, the focus of the plan and services changes. These programs focus on education, and the services provided are those that will help children benefit from school. A child may get physical therapy to help her move from place to place in a classroom or hold a pencil; but services to support families in a larger sense are more limited.

Remember, too, that the planning team decides services under local, federal and state guidelines. As children age, their services often look very different, even if their primary concerns remain the same. In addition, when families move from place to place, services for their children can be determined under very different guidelines, as each state and locality is different.

Overhead Partners Plus Caregiver 5-4

Early Intervention and Public School Services Can Include:

- Home visits
- Classroom programs
- Play groups
- Therapies
- Groups for Families
Common Disability Issues

Introduce the concept of transition.

Refer to overhead Partners Plus Caregiver Common Disability Issues 5-5. The first issue listed, Transition, is described in Section 5 of the Caregiver Manual.

When we think about issues that affect children with disabilities and their families, then, we want to include moving from program to program, or “transition,” as a common issue for families.

While families are often very happy to see their children grow and graduate from early intervention services, they are asked to learn about very different programs with very different rules and different staff members all at the same time. While they may be pleased to move, they might be nervous about what services for their children will be like in unfamiliar communities.

Transition can be very difficult for families, and caregivers may need to be supportive of families as they learn about their new situations.

Overhead Partners Plus Caregiver 5-5

Common Disability Issues

- Transition
- Accessibility
- Language
Introduce the concept of accessibility. Accessibility is discussed in Section 5 of the Caregiver Manual.

People with special needs also face discrimination. In 1990, the American with Disabilities Act (or the ADA) was signed into law. For the first time, that legislation outlawed discrimination of people with disabilities. One of the issues that it addresses is accessibility. Under this law, places that the public uses must be accessible. That is, people with disabilities must be able to use them. People who use wheelchairs, for example, cannot use buildings where there are steps to the front door and no wheelchair ramps. Signs that do not include braille writing are unreadable to people with severe visual impairments. ADA was enacted to encourage people to make their facilities accessible.

Despite the passage of the ADA, however, many businesses and agencies have yet to make their facilities accessible because of federal waivers, exemptions, the burden of cost, time lines, etc. Accessibility continues to be an issue for many children and families.
Common Disability Issues continued

The final issue, Language, is addressed in Section 5 of the Caregiver Manual.

Using “people first” language when you are working with children with disabilities is helpful and respectful.

A third issue is language. As you work with families, listen to the words that they use to describe their children and conditions. Sometimes, people choose those words for very important reasons.

At one time, people with disabilities were called “handicapped” or “disabled.” In recent years, people with disabilities have told us that they feel that these terms are very limiting. They would like people to use language that describes them as people first before describing their condition.

Make sure that participants understand this point. Use the examples listed in Section 5 of the Caregiver Manual for practice. Go over these with participants to make sure everyone understands.

When using people-first language, instead of saying, “the disabled man,” we say “the man with a disability.” Let’s go over a few examples from your manuals. Turn to Section 5.
Common Disability Issues continued

Follow the family's lead regarding the use of people-first language.

Some families will not use people-first language, and you can feel free to adopt their way of describing their children. It is crucial to be sensitive to the people you work with. Listen carefully to what they say, and if they ask you to use different language, be sure to do that.

Overhead Partners Plus Caregiver 5-5

Common Disability Issues

- Transition
- Accessibility
- Language
Summary

Ask the group for feedback.

Does anyone have any questions or comments about the information presented in this section? I encourage you to read through the information in the manual for more information on disabilities awareness.

Conclude the section by reinforcing key concepts.

Families and children with disabilities have far more options today than they did 20 years ago. As caregivers, you are in a strong position to advocate for the children you work with and their families. Let families know that they have rights under the law.

Encourage businesses in your community to make their facilities accessible to people with disabilities. IDEA and the ADA were made into law because people with disabilities and their families and friends took a stand. Your voice is important!
Section 5 Caregivers—Disabilities Awareness
Suggested Overheads
IDEA
(Individuals with Disabilities Education Act)

Birth - 2:
• Early Intervention Services (EI)

Ages 3-21:
• Free and Appropriate Public Education (FAPE)

Ages 3-5:
• Early Childhood Special Education (ECSE)

Ages 5-21:
• Special Education (SpEd)
Plan

Includes:

- information about the child's development
- goals or outcomes for the child
- activities to help the child achieve the goals
IFSP

Individualized Family Service Plan
(for early intervention)

Serving the child within the context of the family.

IEP

Individualized Education Plan
(for preschool service and beyond)

Serving the child within the context of the school system.

DRAFT 10.99
Early Intervention and Public School Services

can include:

• home visits
• classroom programs
• play groups
• therapies
• groups of families
Common Disability Issues

Transition

Early Intervention

Accessibility

Language

Partners Plus Workshop
Caregiver Manual Section 5
Child Development Resources, P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99

BEST COPY AVAILABLE
Section 6 Caregivers—Understanding Child Development

Objectives
- Learn information on child development and planning appropriate activities for children
- Understand the basics of child development
- Use knowledge about child development to work effectively with children
- Be aware of the importance of play and enriching environments for child development
- Be aware of potential hazards in an environment and appropriate safety precautions

Preparations
Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 6, “Understanding Child Development,” of the Caregiver Manual.

Prepare overhead transparencies of:
- Overhead Partners Plus Caregiver Areas of Development 6-1
- Overhead Partners Plus Caregiver Areas of Development 6-2
- Overhead Partners Plus Caregiver Guidelines for Caregivers’ Involvement in Play 6-3

Corresponding Sections
Please refer participants to Section 6, “Understanding Child Development,” in the Caregiver Manual.
Introduction to Child Development

Introduce the concepts of the section.

Providing respite care for children with special needs is not so different from caring for children who are typically developing. Respite caregivers who work with children who have special needs do require some special information and skills to do their job effectively. However, the most important knowledge in caring for children is understanding child development. It is important to know how children grow and learn new skills in order to be able to plan enjoyable and challenging activities for them.

At each age, we expect children to reach certain developmental milestones. For example, we expect to see children take their first steps at around a year old.
Children with special needs may reach certain milestones later than other children. For example, a child with cerebral palsy might walk later, and a child with a cleft palate might say her first words later. Sometimes, children with special needs do not reach milestones at all.

It is important to remember that development follows a logical order for all children. A person of any age must be able to hold her body up before she can stand.
Planning Appropriate Experiences

We need to understand the order in which we can expect to see different skills develop in children so that we know what children can be expected to learn.

For example, a very young child first plays alone. Next we expect a child to play with toys close to other children, before being ready to join in play with one or more children.

When we know the range and sequence of typical development, we are able to know what skills we can help a child learn with success and what activities to plan.

There are no overheads for this section.
Planning Appropriate Experiences continued

Explain how to plan activities that give children with disabilities a chance to use their developmental strengths.

Many children with disabilities have some areas of development in which their skills are similar to those of their peers. Emphasizing those strengths is important to each child's confidence and overall development.

Offer an example to illustrate this point.

For example, a 4-year-old child with mental retardation may not be able to play with others or take part in social activities typical for children her age. However, the way she uses her body may be typical. She can climb on play equipment, ride a trike, and run and jump like other children her age. Giving her opportunities to practice using her muscles will help her to continue developing her abilities, help her to feel good about herself, and give her ways to play with other children. You can help her expand her social skills by building on her strengths, playing near or with other children on playground equipment.
Describe how a child with a visual impairment can take the lead in a game like pinata.

In games where groups of school-aged children participate, caregivers can find opportunities for children with disabilities to take the lead and demonstrate their abilities among peers. A child with visual impairment whose motor skills are like that of their peers could take the lead in a game of pinata. In this game, a brightly colored ornament is suspended above children's heads, and blindfolded players take turns hitting it with a stick to break it open. Taking the lead in this game helps the child gain confidence and build self-esteem.

Make sure that participants understand these principles before going on to the Basic Principles of Child Development.

Are there any questions on this material?

Basic Principles of Child Development

Introduce the basic principles of child development.

Child development is an orderly sequence of physical, intellectual, and emotional change that occurs throughout childhood. Although a child develops as a whole person, different words and categories are used to describe different areas of development.

For a child with a disability, understanding the different areas of child development will help us understand the child's development. However, a disability in one or more areas of development frequently affects other areas of development as well.

There are no overheads for this section.
Discuss the five areas of development. Refer to overhead Partners Plus Caregiver Areas of Development 6-1. Refer participants to Section 6 in the Caregiver Manual.

As children grow older, they change in all kinds of ways. There are five different areas of development. Caregivers can plan activities that will support children in all of these areas of development.

The first area of development is physical development. Physical development refers to the way the child's body grows and functions.

Gross motor development is a type of physical development. When we talk about gross motor development, we are referring to the ways that children use their big muscles to support and move their bodies.

Overhead Partners Plus Caregiver 6-1

<table>
<thead>
<tr>
<th>Areas of Development</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physical Development:</td>
</tr>
<tr>
<td>• Gross Motor</td>
</tr>
<tr>
<td>• Fine Motor</td>
</tr>
<tr>
<td>• Vision</td>
</tr>
<tr>
<td>• Hearing</td>
</tr>
<tr>
<td>Communication Development:</td>
</tr>
<tr>
<td>• Receptive Language</td>
</tr>
<tr>
<td>• Expressive Language</td>
</tr>
</tbody>
</table>
Discuss the five areas of development.

Fine motor development refers to how children use their small muscles (their fingers and their tongues) for activities such as grasping small objects, coloring, turning knobs, and chewing and swallowing.

Vision refers to how children see.

Hearing refers to how children hear.

Communication development refers to the way that children understand messages and let us know what they want, need, or understand. There are two parts of communication development: receptive language and expressive language.

Receptive language describes the way that children understand messages from others.

Expressive language describes how children get their messages across - through sounds, gestures, words, signing, or the use of communication devices.
Basic Principles of Child Development continued

Discuss the five areas of development.

Children learn receptive language before they learn expressive language. A young child will understand the sentence “Go get the ball” long before he will be able to say that sentence or even the word “ball.”

Refer to overhead Partners Plus Caregiver Areas of Development 6-2.

Cognitive development describes the way that children think and solve problems.

As you work with children, think about how they use their brains to play with the toys that they enjoy.

Many activities that caregivers do with children help support cognitive development. For example, pop-up boxes, toys where children push buttons or twist knobs to make animals pop out, teach children that their actions cause other things to happen. Understanding cause and effect is one important cognitive milestone.
Basic Principles of Child Development continued

Discuss the five areas of development.

Adaptive development refers to the way that children take care of their personal needs. As you work with children, ask yourself: How do these children get dressed, feed themselves, brush their teeth? Learn from families what their children can do for themselves in this area. Encourage their independence as much as possible!

Social or emotional development refers to the way that children think and feel about themselves and the people around them. It also refers to the way children form relationships with the people in their lives.

Caregivers can promote social and emotional development by offering children encouragement and care.
Basic Principles of Child Development continued

Emphasize the following points.

Children with special needs sometimes have "delays" in development. They may not reach developmental milestones at the same time other children typically do. Caregivers can work with families to help support children as they try to reach developmental milestones.

Delays in one area can affect development in another area. For example, a child with significant physical delays may not be able to play on the slide without extra support. What happens when everyone else is playing on the slide and this child is not? He is missing out on the opportunity to work on his social skills in play with these other children. A caregiver can physically support the child so that he, too, can enjoy the slide and the social play.

Children with special needs also have strengths. Attend to children's strengths and build on them. A child who is showing...
Basic Principles of Child Development continued

Discuss the five areas of development. Language delays might have strong muscles. Suggest to her family that you take her to the playground. As she is playing, label objects for her to help her language development. Encourage her to play with other children, so she can hear their language and develop social skills. Praise her efforts! All of these activities build on her strong muscles and support her development in other areas as well.

Discuss the importance of play. One of the best ways to support the development of children is to play with them. Play is a child's work, and it is what we use to help children achieve developmental milestones. There are a few simple guidelines for playing with children.

Overhead Partners Plus Caregiver 6-3

Guidelines for Caregivers' Involvement in Play

- Observe
- Follow
- Be Creative
- Be Safe
- Have Fun!

First, it is important to observe the child as he plays. Note what skills he is using, and what activities he enjoys. This information will be helpful later, as you plan activities.
Discuss the importance of play. With him. In play, you can help him build on his skills using activities you know he enjoys.

Join the play when the child invites you. As you play, make sure that you are playing at the child's level. Let the child be in charge! Follow her lead!

Be creative in your play! There are lots of different ways to play and learn. Think about a shoe box.

As caregivers answer, ask them to think about what areas of development are being used in the activities they describe.

How many different ways can you play with a shoe box?

What areas of development are being used when we play with the box this way?

**Overhead Partners Plus Caregiver 6-3**

**Guidelines for Caregivers' Involvement in Play**

- Observe
- Follow
- Be Creative
- Be Safe
- Have Fun!
Basic Principles of Child Development continued

Discuss how play uses various areas of development.

Usually, play involves more than one area of development. Hiding an object under a shoe box, for example, encourages physical development (as the child lifts the box and grasps the object) and cognitive skills (as the child tries to find the object).

Show caregivers how they can use this activity to work on other areas as well.

You can use play to work on other areas, as well. By labeling the object and the box, caregivers encourage communication development. By clapping when the child finds the object, the caregiver encourages emotional development.

So, with a little creativity, you can work on all of the areas of development with just a shoe box!

There are no overheads for this section.
Discuss guidelines for play.

The next guideline is to be safe. Before you begin respite, discuss safety issues with families. What might put their children in danger? What should you watch for?

Examine the space you will be using to provide care. Are knives and dangerous tools safely stored? Are electrical outlets covered? Are glass trinkets on low tables where children can break them? Are there sharp corners to watch for? Where will you go in case of fire? Some caregivers find it helpful to get down on their hands and knees to examine the space from the child's perspective. There is also a safety checklist included in the Caregiver Manual. When you are planning care with a family, read over this checklist and discuss it with the family.

The last guideline is to have fun!
Summary

Ask the group for feedback.

Are any questions or comments on this section? I encourage you to read through Section 6 in the Caregiver Manual for more information on child development.

Conclude the section with a summary statement.

Respite is a wonderful experience when you are meeting a child at his own level, challenging him, playing with him, and watching him build skills. Awareness of child development is the key to this kind of experience!
Section 6—Understanding Child Development Suggested Overheads
Areas of Development

Physical Development

- Gross Motor
- Fine Motor
- Vision
- Hearing

Communication Development

- Receptive Language
- Expressive Language
Areas of Development

• Cognitive Development
• Adaptive Development
• Social or Emotional Development
Guidelines for Caregivers' Involvement in Play

Observe
Follow
Be Creative
Be Safe
Have FUN!

Partners Plus Workshop
Caregiver Manual Section 6
Child Development Resources, P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Section 7 Caregivers—Children with Special Needs

Objectives

- Understand basic information about common special needs
- Be aware of the basic developmental and medical effects of common special needs on children
- Understand what caregivers may be asked to do when they care for children with common special needs

Preparations

Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 7, “Children with Special Needs,” in the Caregiver Manual.

Prepare overhead transparencies of:

- Overhead Partners Plus Caregiver Common Disabilities 7-1
- Overhead Partners Plus Caregiver Common Disabilities 7-2
- Overhead Partners Plus Caregiver Common Special Health Care Needs 7-3
- Overhead Partners Plus Caregiver Common Special Health Care Needs 7-4

Corresponding Sections

Please refer participants to Section 7, “Children with Special Needs,” in the Caregiver Manual.
Introduction to Children with Special Needs

Another useful tool for caregivers in respite is a basic understanding of common special needs. Refer caregivers to Section 7 of their manuals: Children with Special Needs. As we do this, remember these three things:

1. The best source of information about a particular child and his diagnosis is that child's family.
2. Through the Partners model, you have choices about working with particular children and families.
3. There is a wide range of special needs.

Let's talk about each of these items separately.
Describe the three main points to remember when working with children with special needs.

First, the best source of information about a particular child and his diagnosis is that child's family. The same condition can affect individual children differently. While I will give you some general information today, pieces of it may not apply to certain children. Take your cues from families about how to work with their children.

Second, through the Partners model, you have choices about working with particular children and families. If you feel uncomfortable about working with a child with a particular diagnosis for any reason, discuss it with her family. All we ask is that you keep an open mind and see what you learn!

Third, there is a wide range of special needs. While we will talk briefly about common special needs today, we will not be able to discuss everything. Please take the time to read this information and the information in the Resources section of your manual thoroughly. Your manual lists places where you can gather more information if you want to do so.
Common Disabilities

Refer to the overhead Partners Plus Caregiver Common Disabilities 7-1. The information in this section corresponds with Section 7 of the Caregiver Manual.

Work your way through the disabilities, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

This is a list of the disabilities that we commonly hear about. In the next few minutes, we will briefly describe each diagnosis and talk a little bit about special things caregivers might need to do for children with those conditions.

Attention Deficit Disorder (ADD) or Attention Deficit Hyperactivity Disorder (ADHD):

Children with ADD or ADHD have very short attention spans. They often have difficulty focusing on tasks, and may act on impulse. Children with ADHD may be physically active as well.

Caregivers may need to work with families to make sure that children with ADD or ADHD have consistent, structured routines. In addition, in some cases, caregivers may need to learn medication routines for these children.

Overhead Partners Plus Caregiver 7-1

<table>
<thead>
<tr>
<th>Common Disabilities</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Attention Deficit Disorders</td>
</tr>
<tr>
<td>- Autism</td>
</tr>
<tr>
<td>- Blindness / Visual Impairment</td>
</tr>
<tr>
<td>- Deafness / Hearing Impairment</td>
</tr>
<tr>
<td>- Emotional Disturbance</td>
</tr>
</tbody>
</table>
Common Disabilities continued

Work your way through the disabilities, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child’s family.

Autism:
Autism is a condition that affects a child’s brain. Children with autism often have trouble with social and communication skills. Caregivers may need to learn to communicate with these children and closely watch their interaction with other people. Children with autism also tend to respond well when caregivers structure their routines.

Blindness/Visual Impairment:
These diagnoses tell you that the children you are working with have trouble seeing. Visual impairments are less severe than blindness, and more common. Children with visual impairments have some level of vision. As a caregiver, find out what a child can and cannot see. Adapt activities to make sense for them. If you are working with...
Common Disabilities continued

Work your way through the disabilities, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

children with visual impairments in a space that is unfamiliar to them, you want to stay close to them as they learn about this new space. Try to keep furniture and objects in the same places each time.

Deafness/Hearing Impairment:
Deafness is the inability to hear. Again, hearing impairments are less severe than deafness.

Caregivers for children with these diagnoses should learn how to communicate with them. How do they communicate with others? It may be that you will have the opportunity to learn a whole new language!

Emotional Disturbance:
Children with emotional disturbance usually have trouble with social skills or

Overhead Partners Plus Caregiver 7-1

Common Disabilities
- Attention Deficit Disorders
- Autism
- Blindness / Visual Impairment
- Deafness / Hearing Impairment
- Emotional Disturbance
Common Disabilities continued

Work your way through the disabilities, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child’s family.

Refer to overhead Partners Plus Caregiver Common Disabilities 7-2.

Learning Disability:
Learning disabilities cause children to have trouble with reading, writing, math, listening, speaking or using logic. These children generally do poorly in school, but they may be very smart. Their brains simply have trouble understanding or sending information. There are many different kinds of learning disabilities. Caregivers should learn more specifically what kinds of learning disabilities the children in their care have. Find out from families how these children learn best, and how you can best support them.

Overhead Partners Plus Caregiver 7-2

Common Disabilities
- Learning Disability
- Mental Retardation
- Severe and Profound Disabilities
- Speech Impairment / Language Disorder
- Traumatic Brain Injury
Common Disabilities continued

Work your way through the disabilities, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

Mental Retardation (MR):
Children with MR show delays in cognitive and adaptive development. They often learn new skills at a slower rate. Caregivers may need to take things at a slower pace with these children. Find out the age level at which children with MR are operating. For example, an eleven-year-old may have the cognitive skills of an eight-year-old. This information will help you plan activities for the children you are working with.

Orthopedic Impairment:
An orthopedic impairment affects the bones, muscles, and/or the way the body moves. There are many diagnoses included in this category, including cerebral palsy, spina bifida, and muscular dystrophy. Children with orthopedic impairments generally show motor delays.

Overhead Partners Plus Caregiver 7-2

Common Disabilities
- Learning Disability
- Mental Retardation
- Severe and Profound Disabilities
- Speech Impairment / Language Disorder
- Traumatic Brain Injury
Common Disabilities continued

Work your way through the disabilities, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

Depending on which parts of the body are affected, caregivers may need to help children with orthopedic impairments sit, stand, move from place to place, eat, communicate, and/or toilet.

Severe and Profound Disabilities:
Children with severe mental retardation are generally given the diagnosis of severe and profound disabilities. These children may have additional disabilities as well.

Children with this diagnosis generally need extensive help with self-care. Caregivers can learn from families about the best ways to work with them.

Speech Impairment/Language Disorders:
Children with speech impairments have difficulty producing sounds or have trouble with the way that their voices

Overhead Partners Plus Caregiver 7-2

Common Disabilities
- Learning Disability
- Mental Retardation
- Severe and Profound Disabilities
- Speech Impairment / Language Disorder
- Traumatic Brain Injury
Work your way through the disabilities, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

A language disorder is diagnosed when a person has trouble using words appropriately or understanding what is said to him or her.

Caregivers will need to learn to understand and communicate with these children.

Traumatic Brain Injury (TBI):

TBI occurs when a person experiences a sharp blow to the head or the brain is damaged from an injury. Because different parts of the brain control different pieces of functioning, symptoms of TBI vary according to what part of the brain was injured.

Caregivers need to learn how the children they are working with are affected by TBI, and how they can help.
Common Special Health Care Needs

Refer to overhead Partners Plus Caregiver Common Special Health Care Needs 7-3. The information in this section corresponds with Section 7 of the Caregiver Manual.

Working with children with special health care needs can be a different experience from working with children with other disabilities. Their concerns are generally more medical. Here again, we will briefly go through this list of special health concerns and talk about how caregivers might work with these children.

Allergies:
Some children need to avoid certain substances because they are allergic to them.

Caregivers will need to learn what those substances are, how they react when they come in contact with them, and what to do if they have a reaction.
Common Special Health Care Needs continued

Work your way through the diagnoses, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

Asthma:
Children who have asthma have difficulty breathing at times. Allergens generally trigger attacks.
Caregivers will need to learn how to know when a child is having an asthma attack and how to help him. In addition, caregivers should know what triggers attacks so they can plan activities that do not increase the risk of attack.

Cancer:
Cancer is a group of abnormal cells that multiplies uncontrollably. Sometimes, these cells can travel through the body and affect different body organs. Like TBI, cancer affects children differently depending on where these cells are growing.
Caregivers should plan for extra rest periods for children with cancer, who

Overhead Partners Plus Caregiver 7-3

Common Special Health Care Needs
• Allergies
• Asthma
• Cancer
• Congenital Heart Disease
• Cystic Fibrosis
Work your way through the diagnoses, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

...may need to take medication or eat special diets. Ask families how to keep their children comfortable during care.

**Congenital Heart Disease:**
Children with congenital heart disease have trouble with their hearts from birth. Their hearts may not beat regularly or they are not structured the way they should be to do their jobs.

Caregivers of children with congenital heart disease should learn what activities are too stressful for children and what they enjoy doing. Caregivers may learn how to use a heart monitor, and how to respond to an emergency.

**Cystic Fibrosis:**
Cystic fibrosis is a condition that causes the lungs and other body organs to produce a lot of mucus. This mucus can build up in the child's body and affect the way that body organs function.

**Overhead Partners Plus Caregiver 7-3**

<table>
<thead>
<tr>
<th>Common Special Health Care Needs</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Allergies</td>
</tr>
<tr>
<td>• Asthma</td>
</tr>
<tr>
<td>• Cancer</td>
</tr>
<tr>
<td>• Congenital Heart Disease</td>
</tr>
<tr>
<td>• Cystic Fibrosis</td>
</tr>
</tbody>
</table>
Common Special Health Care Needs continued

Work your way through the diagnoses, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

Refer to overhead Partners Plus Caregiver 7-4.

Caregivers for children with cystic fibrosis may feed them special diets and give them medication. They may also need to learn how to help these children clear their lungs of mucus.

Diabetes:

Children with diabetes do not process the sugars in food the way that other people do.

Caregivers may help children follow special diets, and give them medicine. In addition, caregivers may help children take blood tests to monitor their blood sugar. Finally, caregivers should watch children with diabetes for signs of high blood sugar or low blood sugar, and learn how to help them in these events.

Overhead Partners Plus Caregiver 7-4

Common Special Health Care Needs

- Diabetes
- HIV / AIDS
- Kidney Disease
- Seizure Disorders
Common Special Health Care Needs continued

Work your way through the diagnoses, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

HIV/AIDS:
HIV, the virus that causes AIDS, attacks the immune system. As a result, children with AIDS are vulnerable to infections. They may have other health conditions for which they will need special care. Caregivers should learn from families how HIV/AIDS affects their children and how they can help.

Kidney Disease:
The kidneys clean waste from the blood. Children who have kidney disease have trouble with the way that their kidneys function.
Caregivers may need to take children with kidney disease to the hospital periodically to get the waste cleaned out of their blood. In addition, they may help children with special diets and medication routines.

Overhead Partners Plus Caregiver 7-4

Common Special Health Care Needs
- Diabetes
- HIV / AIDS
- Kidney Disease
- Seizure Disorders
Work your way through the diagnoses, emphasizing the points listed and using questions from the Caregiver Manual to illustrate how caregivers can learn to care for a child from the child's family.

Seizure Disorders:
Seizures occur when there is a sudden change in the way that the brain functions. As a result, the person with the seizure temporarily loses control over all or parts of her body. Epilepsy is one type of seizure disorder. Caregivers will need to learn about the specific kind of seizure disorders affecting the children in their care. Many children with this diagnosis eat special diets and/or take medication. Caregivers will also learn how to monitor and keep children safe during seizures.
Summary

Ask the group for feedback.

Does anyone have any questions or comments about the information we just discussed? I encourage you to read through the information in this section in the Caregiver Manual very carefully for more information on special needs, how they affect development, questions to ask families about their children's conditions, and where to look for more information.

Summarize the section.

This information should be useful to you as you begin working with families with children with special needs. However, remember that families will give you the most information about their children and their children's conditions.

If you have questions, ask them to learn more. You need to feel comfortable with the children in your care.
Section 7—Children with Special Needs

Suggested Overheads
Common Disabilities

- Attention Deficit Disorder
- Autism
- Blindness / Visual Impairment
- Deafness / Hearing Impairment
- Emotional Disturbance
Common Disabilities

- Learning Disability
- Mental Retardation
- Orthopedic Impairment
- Severe and Profound Disabilities
- Speech Impairment / Language Disorder
- Traumatic Brain Injury
Common Special Health Care Needs

- Allergies
- Asthma
- Cancer
- Congenital Heart Disease
- Cystic Fibrosis
Common Special Health Care Needs

- Diabetes
- HIV/AIDS
- Kidney Disease
- Seizure Disorders
**Section 8 Caregivers—Becoming a Caregiver**

**Objectives**
- Understand how caregivers can become active members of a respite partnership in the Partners program
- Understand how caregivers can contribute to personalized training
- Know how to match with families for respite

**Preparations**
Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of Section 8, “Becoming a Caregiver,” in the Caregiver Manual.

In this section, caregivers get information on becoming a Partners caregiver. Each community can choose how or if they distribute information on trained caregivers to families in the community. Please prepare for this section by becoming familiar with these procedures and developing necessary materials to communicate this to caregivers.

Prepare overhead transparencies of:
- Overhead Partners Plus Caregiver How I Learn Best 8-1
- Overhead Partners Plus Caregiver Training Ideas 8-2

**Corresponding Sections**
Please refer participants to Section 8, “Becoming a Caregiver” and “Final Activity” in the Caregiver Manual.
Introduction to Becoming a Caregiver

Introduce the concepts of the section.

Congratulations! You have almost finished Partners workshop! You have learned some basic information for providing respite care to children with special needs.

There are no overheads for this section.
The Caregiver Role in the Partners Plus Model

Refer caregivers to Section 8, “Becoming a Caregiver,” in the Caregiver Manual.

After the workshop, the next step for interested caregivers is to match up with families for respite, and to go through personalized training with those families.

Partners is a family-centered model, and families have choices about what respite will look like for them. Likewise, caregivers also have choices about how they want to participate. The materials in this section will help you think about how you want to participate in interviewing, personalized training, and the respite experience.

Introduce this section.

Ask participants to turn to “What Can I Contribute as a Respite Caregiver?” in Section 8.

Turn to Section 8, “What Can I Contribute as a Respite Caregiver?” in your manuals. The information on this page will help you think about why you want to be a respite caregiver. Families need committed caregivers. Think about whether this is work you will enjoy.

There are no overheads for this section.
The Caregiver Role in the Partners Plus Model continued

Discuss the sample interview questions. The “Sample Interview Questions” on the next few pages of your Caregiver Manuals are the questions we suggest that families use in interviewing caregivers. Before an interview, take some time to consider these questions and your answers.

In this process of preparing yourself for the interview, think about what you want in this partnership, and why you think you will be a good respite caregiver.

Remember, the interview process is a time when you gather information from families just as they are gathering information about you. Take the time to think about some questions that you have for families before you interview with them.

There are no overheads for this section.
The Caregiver Role in the Partners Plus Model continued

Refer caregivers to “Developing a Personalized Training Plan” in their manuals.

If you and a family decide to work together, you will begin personalized training. As we have been talking, families have been learning a structure for training caregivers about their children. On the next pages, we have included that information for you, so you will have an idea of what training might look like.

Give an example to illustrate a “Personalized Training Plan.”

We have suggested a four-step process for training. The family plans around the time of day their child will need care. In the first step, families identify their training goals. What will they need you to know for that time period? Then, they decide what training methods to use. How will they teach you what you need to know? In Step 3, families use resources to help you learn about their child. And finally, in Step 4, caregivers and families decide how they can evaluate the plan. How will you and families know when you have learned the goals, and that children, parents and caregivers are comfortable?
The Caregiver Role in the Partners Plus Model continued

Refer caregivers to "Tips for Learning" in their manuals.

Caregivers should be active in their personalized training. There are many ways that you can be involved in planning and evaluating what and how you will learn.

Refer to overhead Partners Plus Caregiver How I Learn Best 8-1. This information is in Section 8 in the Caregiver Manual.

Different people learn best in different ways. Some people learn best when they read information. Others learn best when they hear it. Some people like pictures and diagrams. Families want to make sure that you are learning information in the best way possible. Think about how you learn best and share that information with families.

Overhead Partners Plus Caregiver 8-1

How I Learn Best
- Written Materials
- Audiotapes
- Videotapes
- Oral Instruction
- Demonstration, Feedback, Practice
- Other
The Caregiver Role in the Partners Plus Model continued

Refer to overhead Partners Plus Caregiver Training Ideas 8-2. This information is in Section 8 in the Caregiver Manual.

As you are trained by families, three things to keep in mind are:

- Do I have the knowledge to care for this child?
- Do I have the skills to care for this child?
- Do I have the comfort to care for this child?

If you are feeling weak in any area, let the family know so they can provide additional training.

Throughout your respite relationship, share your thoughts, input, and concerns with families. If you are not comfortable in respite, you can be sure that the family you are working with wants to help you become comfortable with their child.

Conclude this section.

Overhead Partners Plus Caregiver 8-2

Training Ideas
- Knowledge
- Skills
- Comfort
The Caregiver Role in the Partners Plus Model continued

Note: Use this time to tell caregivers how your program fosters caregiver/family partnerships. Many communities choose to create a directory of Partners caregivers to distribute to families looking for respite caregivers. If this is true of your community, use this time to tell caregivers how they can be included on that directory.

If any forms need to be completed before becoming a Partners caregiver, introduce them at this time for participants to complete.

Our community has created a directory of Partners caregivers to distribute to families looking for respite caregivers. If you would like to be included on that directory, please . . .

There are no overheads for this section.
Summary

Ask the group for feedback. We've covered a lot of ground in a relatively short period of time. Does anyone have any questions on this section, or on anything we've covered in this breakout session?

Would anyone like to share some comments with the group about what we've discussed?

Conclude the section by stressing key ideas. Remember that you are a partner in this respite relationship. Your opinion counts! Be sure to speak to families about your needs for training and support. Families do not expect that you will know a lot about their children without ever having met them and worked with them. Families tell us that, more than education and experience, the most important qualities in a respite caregiver are a desire and willingness to learn and a good attitude.

There are no overheads for this section.
Section 8—Becoming a Caregiver
Suggested Overheads
How I Learn Best

- Written Materials
- Audiotapes
- Oral Instruction
- Demonstration,
- Practice
- Other
Training Ideas

- Knowledge
- Skills
- Comfort
Final Partners Workshop Activity—Using Daily Routines To Develop a Personalized Training Plan

Overview
Families and caregivers, who were separated for breakout sessions, should be reunited for this activity.

Objectives
- Learn more about developing personalized training plans
- Practice developing a personalized training plan using a child’s daily routines

Preparations
Read the “Training Session Notes” carefully. Be sure you are familiar with the content, materials and design of the Final Activity.

In Section 8 Families, the trainer asked for a family volunteer to share information on her child’s daily routine for this activity. Be prepared to use a child from your own experiences as an example in case no one volunteers. You should know this child well, feel comfortable answering questions about the child, and be able to describe a routine in the child’s day.

Prepare overhead transparencies of:
- Overhead Partners Plus Step 1 FA-1
- Overhead Partners Plus Step 2 FA-2
- Overhead Partners Plus Step 3 FA-3
- Overhead Partners Plus Step 4 FA-4
- Overhead Partners Plus Good Luck FA-5

Gather the following supplemental materials:
- If using overheads, markers for overheads
- If using flip charts, markers for flip charts
Preparing for the Final Activity continued

Prepare overhead transparencies of:
- Overhead Partners Plus 10-1 Step 1
- Overhead Partners Plus 10-2 Step 2
- Overhead Partners Plus 10-3 Step 3
- Overhead Partners Plus 10-4 Step 4

Gather the following supplemental materials:
- If using overheads, markers for overheads
- If using flip charts, markers for flip charts

**Corresponding Sections**

Please refer participants to Section 7 of the Family Manual, "Developing Your Personalized Training Plan," and Section 8 of the Caregiver Manual, "Becoming a Caregiver."
Introduction to Using Daily Routines  
To Develop a Personalized Training Plan

Introduce the concepts of the Final Activity. Refer participants to the Personalized Training Plan in their manuals.

We are glad to bring families and caregivers back together for this final activity. We will practice developing a personalized training plan. As we are going through this activity, think about what we talked about earlier in training: the importance of open and honest communication and the sharing of perspectives.

We are going to begin to build a personalized training plan for ‘s child, (name of the child).

We are going to use the steps outlined in your manuals to do that.

Note: If there were no volunteers from the Family Breakout Session, use a child you know as an example. Be certain to protect confidentiality!
Using Daily Routines To Develop a Personalized Training Plan Activity

Ask the volunteer to tell the group about his or her child.

Before we begin, why don’t you tell us a little about your child.

Great. What time do you want respite for your child? Be specific about the time of day and the day of the week. Having many times that you would like respite is fine, but for our discussion today just choose one.

Excellent. The first step to developing a personalized training plan is to identify your goals for training. What do you need the caregiver to know for this time with your child?

Refer to overhead Partners Plus Step 1 FA-1.

Overhead Partners Plus FA-1

Step 1
• Identify your training goals
Ask the volunteer to list the different goals for personalized training.

Why don't you list the different goals you have for personalized training for a caregiver providing respite at o'clock [the time chosen].

List all of the goals that she mentions on the overhead. Be sure that they are stated as specific goals. For example, instead of writing "feeding routine," write, "The caregiver will know how to feed (the child) lunch."

You may want to prompt the family member to ensure all goals are included.

Does a caregiver need to know about feeding your child? Giving medication? Playing with your child?

There are generally many different training goals for one period of time. Not all of these goals will be taught to caregivers using the same activities. Let's choose one to discuss in a little more detail.

**Overhead Partners Plus FA-1**

- Step 1
  - Identify your training goals
    - 
    - 
    - 
    - 
    - 
    - 
    - 
    - 
Using Daily Routines To Develop a Personalized Training Plan Activity continued

Have the volunteer choose a goal that she feels comfortable describing to the group.

Refer to the overhead Partners Plus Step 2 FA-2.

Encourage the volunteer to share her ideas. Again, write them on the overhead. You may need to prompt her.

Ask other members of the group to get involved. Add additional ideas to the overhead.

Which goal would you feel most comfortable exploring in detail?

Fine. The second step is to decide what methods you could use to achieve your goal. How are you going to teach a caregiver this goal?

How did you learn that routine? What about the teaching activities in the manual?

What about the rest of you? How would you want to learn? Would any of the listed ideas not work for you? Do you learn better in a different way? This is the perfect time to share their ideas and opinions openly.

Overhead Partners Plus FA-2

Step 2
- List your training methods


Partners Plus Trainer's Workshop Manual
Child Development Resources, P.O. Box 280, Norge, Virginia 23127
At this point in the discussion, families and caregivers may ask the family specific questions about the child and the routine. Question and answer is a valuable learning activity.

Now I'd like each of you to pretend that you are the caregiver for this family: do you need more information or details from the family?

Many different methods can be used. Families and caregivers can decide which would work best for them, and in what order they should be used.

Refer to overhead Partners Plus Step 3 FA-3. Ask the volunteer for specific instructions for the caregiver. List resources on the overhead.

If she does not mention anything, describe some possible resources families may use for caregivers.

Let's move on to step 3: The third step is to use resources to help train your caregiver. Is there anything that you can give a caregiver to help him learn about your child or this routine?

Families do not always have resources for each goal. Some possible resources may be . . .

**Overhead Partners Plus FA-3**

<table>
<thead>
<tr>
<th>Step 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use resources to help train your caregiver</td>
</tr>
</tbody>
</table>

---

* Partners Plus Trainer's Workshop Manual
  Child Development Resources, P.O. Box 280, Norge, Virginia 23127

**DRAFT 10.89**

Final Activity-7
Using Daily Routines To Develop a Personalized Training Plan Activity continued

Could you draw or write anything for the caregiver that might be useful? You can do it right on the overhead sheet.

Refer to overhead Partners Plus Step 4 FA-4. Write responses on the overhead.

The final step is evaluation. How will you know when the caregiver has reached the goal and the parents, caregiver, and child all feel comfortable enough to be in respite?

Let's let our volunteer think about that question for a minute. Do you have any ideas for evaluation?

If the volunteer does not have evaluation ideas, prompt her.

Will you know that the caregiver has learned the feeding routine when you watch him go through it? Will you ask the caregiver questions to see if he knows the correct answers?

Overhead Partners Plus FA-4

Step 4
- Evaluate your training plan

- 

- 

- 

- 

- 

- 

Ask caregivers for their input. Add their ideas to the list as well.

What about the caregivers? How will you know when you have reached the family's goals? It is important that caregivers feel comfortable in respite, just as it is important that families feel comfortable leaving their children. When will caregivers feel prepared to do a routine on their own? Do you want families to watch you go through the routine? Do you want to be observed more than once?

Overhead Partners Plus FA-4

Step 4

- Evaluate your training plan
Summary

Ask the group for feedback. Thank you for volunteering to share your experiences with the rest of the group. Does anyone have any questions about this section?

Would anyone like to share some comments with the group about what we've discussed?

Conclude the final activity. After planning for the goal we discussed is completed, the family would go back and repeat this planning process for every goal listed in Step 1.

Families and caregivers who work together through this process will have plans not just personalized to the child and family, but to all three members of the partnership—the family, the caregiver, and the child. These plans are most effective when each personalized training plan is specific to a family-caregiver match. Building these plans together is another way to strengthen the family-caregiver relationship.
Conclude the workshop.

Special Note: Pass out the evaluation measures to families and caregivers. Master copies of these are included with the Supplemental Materials in the Welcome to the Partners Workshop section.

Pass out evaluation measures and tell the participants how helpful their comments are. Distribute copies of "What Did You Think About the Training?" to the group. Distribute copies of "Caregiver Knowledge Measure" and "Caregiver Comfort Measure" to the caregivers in the group.

We hope that you will use the information you have learned here as you build respite relationships in the future. Children only have adults to support them, care for them, and advocate for them. When adults—families and caregivers—work together closely and develop strong relationships, children reap the rewards.

Thank you for coming today! Before you leave, we'd like you to take a moment to complete some evaluation measures for the workshop. These measures will let us know whether we are meeting objectives in training, what we are doing well, and what needs additional work. Your comments are very helpful in our efforts to improve training.
Summary continued

Evaluation measures to be distributed at the workshop are included as supplemental materials at the end of this section. Refer to the Community Planning Manual for a complete set of evaluation measures.

Administer evaluation measures after concluding the session. Leave 30 minutes for people to complete their evaluations.

Overhead Partners Plus FA-5

Good Luck in Your Respite Partnerships!
Final Activity—Using Daily Routines To Develop a Personalized Training Plan

Suggested Overheads
Step 1

- Identify your training goals.
Step 2

• List your training methods.
Step 3

- Use resources to help train your caregiver.
Step 4

• Evaluate your training plan.
Good Luck in Your Respite Partnerships!
Final Partners Workshop Activity—Using Daily Routines To Develop a Personalized Training Plan Supplemental Materials
What Did You Think About the Workshop?

Name: ______________________ Date: ________________
Presenter(s): ______________________

1. What I liked about the workshop...
   ______________________________________________________
   ______________________________________________________
   ______________________________________________________
   ______________________________________________________

2. How Did We Do? The information presented was:
   ____ easy to understand
   ____ difficult to understand because...
   ______________________________________________________
   ______________________________________________________
   ______________________________________________________

3. The information presented was:
   ____ useful
   ____ not useful because...
   ______________________________________________________
   ______________________________________________________
   ______________________________________________________
4. The overall quality of the workshop was: 1 2 3 4 5
   Very Poor Not at All Very Good

5. Were the workshop materials helpful? 1 2 3 4 5
   Not at All Very

6. Was the information presented appropriate for your needs? 1 2 3 4 5
   Not at All Very

7. Was there enough chance for questions and discussion? 1 2 3 4 5
   Not at All Very

8. I gained knowledge about...

   ___________________________
   ___________________________
   ___________________________

9. Suggestions I would like to offer for improving the workshop...

   ___________________________
   ___________________________
   ___________________________

10. I think what I’ll remember most was...

    ___________________________
    ___________________________
    ___________________________

11. Please share any additional comments:

    ___________________________
    ___________________________
    ___________________________
Partners Plus Caregiver Comfort Scale Post Evaluation

Caregiver’s Name: __________________________   Date: __________________________

Please circle the number that best describes how comfortable you feel in working with children with special needs.

<table>
<thead>
<tr>
<th>How comfortable are you:</th>
<th>Uncomfortable</th>
<th>Somewhat Uncomfortable</th>
<th>Very Comfortable</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Providing care for a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>2. Planning activities that a child with special needs can enjoy.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>3. Meeting the routine health care needs of a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>4. Feeding a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>5. Talking with families of children with disabilities about their child and their child’s strengths and needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>6. Talking to other people about any concerns they may have about children with special needs in your care.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>7. Identifying resources and getting help you may need in caring for children with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

Thank You!
Partners Plus Caregiver
Knowledge Post Evaluation

Name: ___________________________ Date: ___________________________

This survey is designed to gather information about your experience and
knowledge of caring for children with disabilities or special needs.

*Read each statement and circle the single best answer.*

1. Early intervention services are available to children with disabilities who are
   a. Birth to three years old  
   b. In public schools  
   c. Three to five years old  
   d. I don't know

2. Children who receive early intervention special education services must
   a. Get therapy  
   b. Go to a classroom program  
   c. Have an individual plan  
   d. I don't know

3. All children with special needs receive education through
   a. MCH  
   b. CHIP  
   c. IDEA  
   d. I don't know

4. Which of these diagnoses tells you that a child has a special health care need?
   a. Autism  
   b. Cystic Fibrosis  
   c. Mental retardation  
   d. I don't know

5. Providing good respite care for a child with a disability depends heavily on
   a. Desire/attitude of caregivers  
   b. Educational level of the caregiver  
   c. A child's disability  
   d. I don't know

6. When caring for children with disabilities in their homes, it is necessary to
   a. Learn about their daily household routines  
   b. Change things to suit your way of providing care  
   c. Ignore the way the family trained you to take care of the child  
   d. I don't know
7. A key to a successful respite care program for families is
   a. Providing each family with many caregivers
   b. Allowing each family to train their own caregivers
   c. To start more respite agencies
   d. I don’t know

8. It is most important for respite caregivers to
   a. Understand child development
   b. Know the needs of the child or children in their care
   c. Have a degree or certification
   d. I don’t know

9. The term “cognitive development” is used to describe how a child
   a. Relates to others
   b. Sits and grasps toys
   c. Thinks and solves problems
   d. I don’t know

10. If caregivers have questions or concerns about a child’s special needs, they should
    a. Discuss their concerns with the child’s parent(s) or guardian
    b. Call the local special education program
    c. Not do anything because if they are wrong it would upset the child's parents
    d. I don’t know

Thank You!
Resources

Overview

This section includes just a few of the many resources available to training programs. Check your local library, internet, and video stores for resources to help trainers become more effective.
References on Adult Learning


Training Resources

Creative Training Techniques Newsletter
Lakewood Publications
50 S. 9th St.
Minneapolis, MN 55402

(800) 707-7749

http://www.trainingsupersite.com/publications
Forms for Sharing Information

Information Worksheets

There are a variety of forms available in this section for families to use in sharing important information with caregivers. Families can use these forms as needed. Some forms provide information specific to the child with special needs and other forms are specific to the needs of siblings also in care. Some forms provide space to include other family and household information. Emergency, health, and safety information sheets and checklists are helpful for caregivers to have.

Master copies of all forms and worksheets are included in this section. Families can make copies of forms that several caregivers may need. Families may also keep blank copies and use them when changes occur or new information becomes available.
Forms for Sharing Information

Information Worksheets

There are a variety of forms available in this section for families to use in sharing important information with caregivers. Families can use these forms as needed. Some forms provide information specific to the child with special needs and other forms are specific to the needs of siblings also in care. Some forms provide space to include other family and household information. Emergency, health, and safety information sheets and checklists are helpful for caregivers to have.

Master copies of all forms and worksheets are included in this section. Families can make copies of forms that several caregivers may need. Families may also keep blank copies and use them when changes occur or new information becomes available.
Essential Information

General Information

Child's Name: __________________________ Nickname: __________________________

Address: ______________________________________________________________________

Phone: __________________________ Allergies: ______________________________________________________________________

Age: _______ D.O.B. _______ Height: _______ Weight: _______

Child diagnosis or description of special needs: ______________________________________________________________________

__________________________________________________________________________

Health Insurance: __________________________ I.D. Number: __________________________

You can reach me at ______________________________________________________________________

If you cannot contact me, please call: _____________________________________________ Phone: __________________________

Medication

Name of medication: ______________________________________________________________________

Purpose for medication: ______________________________________________________________________

Time to be given: __________________________ Dosage: __________________________

Pharmacy: __________________________ Phone: __________________________

Comments: _____________________________________________________________________________

__________________________________________________________________________
Essential Information (continued)

Food

Allergies: 

Cannot have: 

Acceptable snacks: 

Meals: 

Daily Schedule

AM: 

PM: 

Nap Time:
Health and Medical Information

Child's Name: ____________________________ Date of Birth: ____________________________

Current Health Problems / Needs: ______________________________________________________

__________________________________________________________________________________

Allergies (if any): ________________________________________________________________

__________________________________________________________________________________

Child's Physician: ________________________________________________________________

Address: _______________________________________________________________________

Phone: _______________________________________________________________________

Name and Number of Medical Insurance: ____________________________________________

Authorization for Emergency Medical Care

______________________________________ has permission to obtain immediate medical
care for my child if any emergency occurs when the parent or guardian
cannot be located. I understand the child's physician will be contacted
if available.

Parent or Guardian: ____________________________ Date: ____________________________
Additional information about my child:
## Important Phone Numbers

<table>
<thead>
<tr>
<th>Category</th>
<th>Phone Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mother's Work</td>
<td></td>
</tr>
<tr>
<td>Father's Work</td>
<td></td>
</tr>
<tr>
<td>Police/Emergency</td>
<td></td>
</tr>
<tr>
<td>Fire/Emergency</td>
<td></td>
</tr>
<tr>
<td>Poison Control</td>
<td></td>
</tr>
<tr>
<td>Doctor's Office</td>
<td></td>
</tr>
<tr>
<td>Pharmacy</td>
<td></td>
</tr>
<tr>
<td>Hospital</td>
<td></td>
</tr>
<tr>
<td>Dentist</td>
<td></td>
</tr>
<tr>
<td>Neighbor/Friend</td>
<td></td>
</tr>
<tr>
<td>Neighbor/Friend</td>
<td></td>
</tr>
<tr>
<td>Relative</td>
<td></td>
</tr>
<tr>
<td>Relative</td>
<td></td>
</tr>
<tr>
<td>Gas Company</td>
<td></td>
</tr>
<tr>
<td>Electric Company</td>
<td></td>
</tr>
<tr>
<td>Phone Company</td>
<td></td>
</tr>
<tr>
<td>Apartment Manager</td>
<td></td>
</tr>
<tr>
<td>Veterinarian</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
</tr>
</tbody>
</table>
About My Child

Child's Name: ___________________________ Date: __________

What I would like to tell you about my child: ___________________________

__________________________________________________________

Favorite toys or playthings: ___________________________

__________________________________________________________

Special routines (a good-bye hug, snack after school, story for bedtime, etc.)

my child enjoys: ___________________________

__________________________________________________________

Special things (blanket, a pacifier, stuffed animal, etc.) that my child enjoys:

__________________________________________________________

Things that scare my child: ___________________________

__________________________________________________________
Things that quiet or comfort my child:


Special things my child does:


Activities my child likes:


Activities my child dislikes:


Personality or behavior style (such as active, quiet, silly, serious):


Things that are easy for my child:


Things that are hard for my child:


Special names or words for people or things that my child understands/uses:


Other special information or concerns:


531
About Our Family

The people in our family are: _________________________________________________
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________

Some things our family does together: __________________________________________
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________

Special things each member of my family does alone or with someone else:
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________

Things we would like to do if we had respite: _________________________________
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________
Dreams for our child with disabilities: __________________________

__________________________

__________________________

__________________________

__________________________

__________________________

Dreams for our family: __________________________

__________________________

__________________________

__________________________

__________________________

__________________________
About Sisters and Brothers

General Information

Brother/Sister's Name: ___________________________ Nickname: ___________________________

Age: _______ D.O.B.: _______ Height: _______ Weight: _______

Allergies: ________________________________________________

Important health or developmental history: ________________________________

_____________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

Medications

Name of medication: ____________________________________________

Purpose for medication: _________________________________________

Time to be given: ___________________________ Dosage: ______________

Pharmacy: ___________________________ Phone: _______________________

Prescribing Doctor: ___________________________

Doctor's Phone: ___________________________

Let me tell you about _______________________________________

_____________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________


Let me tell you about ____________________________

__________________________

__________________________

__________________________

__________________________

__________________________

__________________________

__________________________

__________________________

__________________________

__________________________

__________________________

__________________________
Health and Safety Checklist

Our caregiver is certified in:

- [ ] infant/child CPR
- [ ] first aid

Our caregiver knows where we keep our:

- [ ] medical treatment release forms
- [ ] medical insurance information
- [ ] first-aid kit and book
- [ ] sunscreen

We have provided our caregiver individualized training on:

- [ ] emergency plans
  - [ ] what we would consider an emergency for our child
  - [ ] what we would like our caregiver to do in case of an emergency
  - [ ] where we keep emergency phone numbers
- [ ] fire safety
  - [ ] location of and how to use the fire extinguisher
  - [ ] our fire evacuation plan
  - [ ] location of smoke detector
- [ ] poison safety
  - [ ] where we keep the Poison Control Center phone number
  - [ ] where we keep the Syrup of Ipecac
- [ ] kitchen safety (i.e., stove, appliances, pots on the stove, etc.)
  - [ ] first aid for burns
We have provided our caregiver individualized training on:

___ hand washing for caregivers and children after toileting and outside time and before food preparation

___ medication administration

___ our smoking policy

___ diaper/toileting routine and diaper disposal

___ toy safety (e.g., balloons might suffocate small children, young children might choke on small toys or toys with small parts)

___ use of medical equipment

___ crib/bed safety
   ___ young children should sleep on their backs
   ___ the side rail of a crib should be latched when the child is asleep
   ___ caregivers should check on sleeping children
   ___ bottles should be held and not propped for young children

___ positioning, exercises, and therapeutic activities

___ car safety
   ___ car seats are available and adjusted for our child
   ___ correct use of a car seat
   ___ child should always be in a car seat or fastened seat belt when the car moving

___ my child's allergies
   ___ medical care if my child is exposed to allergens

___ safety in the home
   ___ electrical outlets are covered
   ___ hot water heater adjusted to 120 degrees or lower
   ___ guns are unloaded and locked in a cabinet
   ___ bullets are locked in cabinets separate from guns

___ medications, poisonous chemicals, and liquor are kept out of the reach of children

___ rubber mats or nonslip surfaces are in showers or tubs children will use
Step 1: Identify training goals.

What do you want your caregiver to learn about your child and family routines?

I want my caregiver to learn about these routines:

Goal 1: __________________________________________________________
______________________________________________________________
______________________________________________________________

Goal 2: __________________________________________________________
______________________________________________________________
______________________________________________________________

Goal 3: __________________________________________________________
______________________________________________________________
______________________________________________________________

Goal 4: __________________________________________________________
______________________________________________________________
______________________________________________________________

Goal 5: __________________________________________________________
______________________________________________________________
______________________________________________________________
Step 2: List training methods.

What methods will you use to teach your caregivers?

In order to reach these goals I will use the following methods:

Goal 1: ____________________________________________

Methods:

1. ____________________________________________

2. ____________________________________________

3. ____________________________________________

Goal 2: ____________________________________________

Methods:

1. ____________________________________________

2. ____________________________________________

3. ____________________________________________
Goal 3: ________________________________

Methods:
1. ________________________________
2. ________________________________
3. ________________________________

Goal 4: ________________________________

Methods:
1. ________________________________
2. ________________________________
3. ________________________________

Goal 5: ________________________________

Methods:
1. ________________________________
2. ________________________________
3. ________________________________
Step 3: Use resources to help train your caregiver.

What information will you use to help your caregiver learn?

These are the resources I would like to use:

- books
- handouts
- videotapes
- audiotapes
- computer search
- I develop resources
- other (please describe)
Step 4: Evaluation

How will you and your caregiver know when a specific skill or routine has been learned?

I will know a specific skill or routine has been learned when we accomplish the following:

Goal 1: ____________________________

______________________________

______________________________

Goal 2: ____________________________

______________________________

______________________________

Goal 3: ____________________________

______________________________

______________________________

Goal 4: ____________________________

______________________________

______________________________

Goal 5: ____________________________

______________________________

______________________________
Personalized Training Plan

Child's Name: ___________________________ Date: __________

Routine: ____________________________________________

____________________________________________________

Time of Day: _________________________________________

Step 1: Training Goal __________________________________

____________________________________________________

Step 2: Training Methods ________________________________

____________________________________________________

Step 3: Resources ______________________________________

____________________________________________________

Step 4: Evaluation ______________________________________

____________________________________________________
Respite Caregiver Profile

Date: ____________

Personal:

Name: ___________________________ SS#: ___________________________

Address: ______________________________________________________________________________________

Home Telephone: ________________ Work Telephone: ________________

In case of emergency, notify: ______________________________________________________________________

Relationship: _____________________________________________________________________________________

Home Telephone: ________________ Work Telephone: ________________

Experience:

Describe experience you have had with children (including your own). _________________________
_______________________________________________________________________________________________
_______________________________________________________________________________________________
_______________________________________________________________________________________________

Have you ever provided care for children with disabilities?

___ Yes ___ No

If yes, please list your experiences: _________________________
_______________________________________________________________________________________________
_______________________________________________________________________________________________
_______________________________________________________________________________________________
Respite Caregiver Profile, continued

Have you had any courses in early childhood, child development, or child health?
  ____ Yes  ____ No

Are you certified in Red Cross First Aid?
  ____ Yes  ____ No  Date of Expiration ______________________

Are you certified in Infant/Child CPR?
  ____ Yes  ____ No  Date of Expiration ______________________

Do you have a valid driver’s license?
  ____ Yes  ____ No  Date of Expiration ______________________

Education:
I have completed:

High School _____________________  Some College _____________________

College ________________________  Graduate School ___________________

Degree or Major (specify): ________________________________

Other training/education: ________________________________

Employment:

Current Employer: ________________________________

Position: ________________________________
Respite Caregiver Profile, continued

Interests: (Check all that apply)

I am willing to care for children ages

birth · 2 ________ 2 · 5 ________ 5 · 8 ________ over 8 ________

I am willing to provide care (check all that apply):

_____ at my home
_____ at child’s home
_____ during the day
_____ during evenings

_____ weekdays
_____ weekends
_____ overnight
_____ anytime

When I provide care, I will care for:

_____ only the child with disabilities
_____ the child with disabilities plus siblings
_____ the child with disabilities and other children in my care
_____ the child with disabilities and my own child/children

I would prefer to provide care:

_____ as a volunteer
_____ in exchange for a service (i.e., transportation, lawn care, meals, etc.)
_____ for an hourly rate of $__________
_____ for a negotiable hourly rate

How far are you willing to travel to provide care? (Check all that apply.)

_____ Williamsburg/James City County
_____ York County
_____ Newport News

_____ Hampton
_____ Poquoson
_____ Gloucester
Respite Caregiver Profile, continued

References:

Please provide the names, addresses, and phone numbers of two persons we may contact who have known you for more than 1 year (excluding relatives or roommates).

Name: ___________________________ Phone: ___________________________
Address: ___________________________

Name: ___________________________ Phone: ___________________________
Address: ___________________________

I am most interested in providing respite care because _______________________
____________________________________________________________________
____________________________________________________________________

Thank You!
Reference Check

Name of Potential Caregiver: ________________________________

Name of Reference: ___________ Phone: _______________________

1. How do you know _________________________________?
   (put name of potential caregiver here.)

2. How long have you known ________________________________?

3. What words best describe ________________________________?

4. How would you best describe how __________________________ relates to children?

5. Describe what you think are _____________________________'s strong points in working with children with special needs and their families?

6. Are there any other comments about __________________________ you would like to make?
Partners Plus
Community Planning Manual

Families and Caregivers in Partnerships
Child Development Resources

DRAFT 10.99
Partners Plus

Families and Caregivers in Partnerships:
Community Planning Manual

Developed By
Amanda C. Hooke, M.S.W.
Lisa L. Ownby, M.S.W.
Dee Wylie Moore, B.S.
Corinne W. Garland, M.Ed.
Adrienne Frank, O.T.R.

The Community Planning Manual is part of a series of resources in the Partners Plus Families and Caregivers in Partnerships: A Family-Centered Guide to Respite Care. The rest of the series includes the Trainer’s Workshop, Family, and Caregiver Manuals, created by Child Development Resources.
Revised from the original
PARTners Plus: Parents as Respite Trainers Community Planning Manual
Authors: Corinne W. Garland, Barbara A. Kniest, Karen S. McKinney, Dee W. Moore, and Nancy Tyree (1995)

Acknowledgments
The authors would like to thank Jayne Naughton for her technical assistance; Amy Watson for editorial assistance; families and caregivers who wrote the perspectives stories in section four of the Family and Caregiver Manuals; and our community advisory committee for multiple tasks throughout the development of training materials. Their suggestions and comments provided a significant contribution to the quality of this manual.

Amy Consiglio, of CM Consulting, designed and produced this manual.

A Word About Funding
Support for the Partners Plus project was provided under grant number HO24B40032 from the Early Education Program for Children with Disabilities, U.S. Department of Education. Points of view or opinions expressed herein do not, however, necessarily represent official views or opinions of the Department of Education.

Funding to revise the Partners Plus manuals to help families use existing community programs as respite for their school-age children with mental retardation came from the Joseph P. Kennedy, Jr. Foundation. Revisions to the manuals include practical “how to” information for personnel working with children in before and after school programs, recreational, camp, and other community programs.

Copyright © 1999. All rights reserved. Trainers may reproduce this manual, without written permission, and in whole for use only with families and caregivers. For permission to reprint single pages or to purchase additional copies of the manual, please contact:

Child Development Resources
Post Office Box 280
Norge, VA 23127-0280
Phone (757) 566-3300 / Fax (757) 566-8977
# Contents

## Introduction
- The Partners Model: A New Model of Respite Care vi
- A Brief Partners History vii
- Partners Model: 5 Steps ix
- Partners Plus Replication Process xii
- Partners Community Implementation Process xiv
- Partners Plus Manuals xv

## Section 1: Planning Your Partners Program
- The Replication Process 1-2
- Planning a Partners Program 1-5
- Creating Community Support 1-6
- Naming Program Advisors 1-9
- Creating Program Structure 1-11
- Securing Financial Resources 1-15

## Section 2: Sample Forms and Information for Planning Your Program
- Developing Your Partners Program: Replication Checklist 2-3
- Partners Project Inquiry Form 2-5
- Individual Contact Log 2-7
- Partners Communication Log 2-9
- Sample Introductory Letter to Families 2-11
- Sample Partners Advisory Committee: Roles and Responsibilities 2-13
Section 2: Sample Forms and Information for Planning Your Program, continued

Sample Partners Advisory Committee Organization Meeting Agenda 2-15
Partners Advisory Committee Workplan 2-17
Supplementary Training Topics 2-19
Sample Partners Budget 2-21

Section 3: Implementing Your Partners Program

Recruiting Families and Caregivers 3-2
Conducting Partners Workshops 3-6
Choosing Caregivers 3-18
Supporting Caregiver Training 3-25
Providing Continuing Support 3-28
Supplementary Training 3-29
Social Support Strategies 3-30
Family Support Groups 3-31

Section 4: Sample Forms and Information for Implementing Your Program

Sample brochure 4-3
Sample newspaper article 4-5
Sample press release and feature articles 4-7
Sample flyer 4-9
Implementing Partners: Planning for Workshop 4-11
Partners Workshop Agendas 4-13
Checklist for Materials and Equipment for Partners Workshop 4-15
Partners Plus Workshop Childcare Information 4-19
Permission for Videotape with Sound/Photograph Use 4-21
Section 4: Sample Forms and Information for Implementing Your Program, continued

Permission to Release Caregiver Name 4-23
Permission to Include Information in Family Directory 4-25
Partners Plus Order Form for Printed Materials 4-27

Section 5: Measuring Your Success
Measuring Your Success 5-2

Section 6: Sample Evaluation Instruments
What Did You Think About the Workshop? 6-3
Partners Plus Initial Family Survey 6-5
Partners Plus Follow-up Family Survey 6-7
Partners Plus Follow-up Caregiver Survey 6-9
Partners Plus Caregiver Knowledge Pre 6-13
Partners Plus Caregiver Knowledge Post 6-15
Partners Plus Caregiver Comfort Scale Pre 6-17
Partners Plus Caregiver Comfort Scale Post 6-19

Section 7: Resources
Resources for Community Programs 7-2
Local and National Resources 7-8
References on Adult Learning 7-9
Notes on this Section
The Partners Model of Respite Care...

*Helps families of children with special needs find and train their own respite caregivers.*

*Through training and support, families and caregivers build long-lasting, successful respite relationships.*

**Introduction**

Like all families, families of children with special needs want and need time away from the demands of parenting. *Respite—or temporary child care for children with disabilities or chronic illness*—gives families time to enjoy an evening out by themselves, with other family members, or with friends. However, for these families, finding respite care is not as simple as calling a neighborhood babysitter. These families need caregivers who understand their children's special needs.

When the staff at Child Development Resources (CDR) spoke with families about what they wanted in a respite program, the message was clear. Families wanted to choose and train their own respite caregivers. They wanted affordable care that was as simple to schedule as calling the neighborhood babysitter. And, families wanted caregivers who were trained to provide care.
In response to families and with support from the Virginia Institute for Developmental Disabilities (VIDD), a University Affiliated Program at Virginia Commonwealth University in Richmond, Virginia, CDR developed a pilot project that was low-cost, family-directed respite care called PARTners: Parents As Respite Trainers.

CDR was awarded, in 1994, a three-year grant from the U.S. Department of Education, Office of Special Education Programs (OSEP), Early Education Program for Children with Disabilities (EEPCD), to build on the pilot and to develop a model of respite that could be replicated in other communities. The project, Partners Plus: Families and Caregivers in Partnerships, provides a model for families to use in finding and training respite caregivers. This model encourages families to use available, natural family and community supports to meet their respite needs.

Within this model, Partners broadly defines the terms respite and caregiver. For example, respite can take place in a family's home, a caregiver's home, or in before and after school, recreation, camp, or other children's program. A caregiver can be any individual who provides respite care to a child with special needs or includes a child with special needs within an existing community program.

Partners Plus provided training to more than 200 families and caregivers in a five-city, three-county area of eastern Virginia. Partners usually conducted the six hour Partners Plus workshop for families and caregivers in one day. Sometimes when it was more convenient for a group, Partners conducted the workshop in two parts on separate days. This flexibility ensured that Partners adapted training...
to fit the schedules of families and caregivers who are generally most receptive to training during nontraditional hours such as evenings and Saturdays. Families attending the workshop learned to provide their own caregivers with personalized training based on their children’s daily routines. Families either came to the workshop with their own caregivers, met caregivers through the workshop, or through other project activities.

The Partners Project resulted in the development of four manuals designed to help communities, trainers, families, and caregivers use the Partners model. The Community Planning Manual is one of four in a series of manuals that Partners has conveniently packaged together in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care. Besides the set of manuals contained in the guide, you can purchase the Family and Caregiver Manuals separately from the Partners Plus program through CDR.

Child Development Resources
Post Office Box 280
Norge, VA 23127-0280
Phone (757) 566-3300
Fax (757) 566-8977
Email ppcdr@tni.net
The Partners Model: A New Model of Respite Care

Partners programs provide group training and continuing support to families and potential caregivers. The Partners model contains five steps:

**The Five Steps of the Partners Model**

1. **Recruit Families and Caregivers**
2. **Conduct Partners Workshop**
3. **Help Families Choose Caregivers**
4. **Support Families as they Train their Own Caregivers**
5. **Provide Continuing Support**

**Step 1: Recruit Families**

First, programs recruit families and potential caregivers to participate in Partners. Community awareness activities invite potential participants to a Partners workshop. Information about the workshop and the Partners program is distributed through parent groups, newspapers, radio, flyers, word of mouth, etc.
Partners Model: 5 Steps

Step 2: Conduct Partners Workshop

Next, programs conduct the Partners workshop. During the six-hour workshop, families learn how to find, screen, and interview caregivers for their children. In addition, they learn to structure training for their own caregivers based on the specific needs and daily routines of their individual children. Caregivers learn basic information about caring for children with special needs. During the workshop, caregivers are preparing to be active in their work with families, reflecting on their motivations for becoming respite caregivers and identifying their own learning needs for personalized training. Partners Family and Caregiver Manuals provide participants with useful resources to actively engage in during the workshop and to use as resources as they move through the next steps of the model.

Step 3: Families Choose Caregivers

After the workshop, Partners programs provide continuing support as families choose caregivers. Families frequently identify respite caregivers within their own circles of natural support. If families are unable to identify their own caregivers, Partners will assist by providing social and educational events that will help families and caregivers meet. During interviews, families and caregivers negotiate how respite occurs. Respite may be in-home or out-of-home and reimbursement for care can be flexible. Care may be provided for a fee, voluntarily, or in exchange for another service, such as yard work.
Step 4: Families Train Caregivers

Once families and caregivers decide to work together, *families train their own caregivers*. This “personalized training,” discussed during the workshop, is organized by families with help from Partners staff if needed. Families decide what routines caregivers need to learn and work with caregivers to determine how training will occur, thus forming true partnerships necessary for successful respite.

Step 5: Provide Continuing Support

As families and caregivers work together to develop a personalized respite training, the community Partners program *offers continuing support*. Information and guidance. The Partners program is available to support families and caregivers in various ways as they begin their respite relationship. Partners staff members respond to requests for help and support from both families and caregivers. For example, Partners may help families and caregivers with screening, interviewing, developing personalized training plans, and providing educational opportunities.
Partners Plus Replication Process

The Partners Plus Community Planning Manual, contained in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care, helps communities use the Partners model of respite care. Each community interested in model replication develops a plan for collaboration among agencies, organizations, and individuals in order to implement the model. Community resources and supports are needed to sustain the model, to create a home for the program, and to ensure continuation.

The community replication process includes four major steps:

1. Identifying the site for replication
2. Planning for replication
3. Implementing the model
4. Evaluating the model


Your community will be a good candidate for replication if you have the following:

- support from agencies that provide, or are involved with, respite services
Partners Replication Process

- knowledge of current respite funding to support families without financial resources
- support and commitment from local early intervention programs, mental health or mental retardation programs, and/or family support programs

Each community needs to develop a plan for replication. Awareness activities aimed at a broad array of agencies, organizations, and individuals help strengthen community support. The creation of an advisory committee brings expertise and resources that will ensure a quality program. The replication plan includes finding a home for the program and securing staff responsible for implementing the program. Formal or informal interagency agreements and perhaps grant proposals may be necessary for community replication.

The Community Planning Manual provides strategies and materials to help communities develop and implement the respite care program. The Trainer’s Workshop Manual, accompanying the Community Planning Manual, includes agendas, overheads, and suggestions for what to say during the Partners workshop. There are also suggestions for the kinds of resources that are needed by families and caregivers in order to foster and maintain long-term relationships.

Communities that replicate the Partners model gather information from families and caregivers to evaluate the quality and usefulness of training and support.
Partners in your Community

Partners Community Implementation Process

Identify Community Site

Plan for Replication of the Partners Model
- Create Community Support
- Determine Program Advisors
- Determine Administrative Structure
- Secure Financial Resources

Implement the Partners Model
- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families As They Train Their Own Caregivers
- Provide Continuing Support

Evaluate the Partners Program
There are four manuals in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care that support the Partners Plus replication process:

1. The Family Manual
2. The Caregiver Manual
3. The Community Planning Manual, and

The Family and Caregiver Manuals are used in the Partners workshop for families and caregivers. These manuals include resources that families and caregivers can continue to use as they develop partnerships for respite care for children with special needs. The Community Planning Manual helps communities develop a community-based, family-centered respite system. The Trainer’s Workshop Manual contains overheads, forms, and “helpful hints” for conducting Partners workshops.

The Family Manual is easy to follow and guides families in choosing and training their own respite caregivers. Families use their children’s daily routines to develop training plans for caregivers. The manual includes information about respite, communication, building partnerships, selecting caregivers, and developing personalized training. There are worksheets, resources, and forms that can be duplicated for future use.
The Caregiver Manual provides caregivers with basic information on caring for young children with disabilities and supporting families. This manual includes information about respite, communication, building partnerships, disabilities awareness, child development, children with special needs, and personalized training with families.

The Community Planning Manual provides communities with a step-by-step guide to replication of the Partners model. By following each step, communities and families can work together to develop respite care programs. The manual contains forms and materials to assist with replication of the model.

The Trainer's Workshop Manual, used in conjunction with the Community Planning Manual, helps community trainers to conduct the six-hour Partners workshop for families and caregivers.
Section 1
Planning your Partners program
The Replication Process

Planning Your Partners Program:
The Replication Process

This manual, the Community Planning Manual, helps communities use the Partners model. It Guides communities through the replication process outlined in the figure on page 1-3. Each community interested in model replication develops a plan for collaboration among agencies, organizations, and individuals in order to implement the model. Community resources and supports are needed to sustain the model and to create a lasting home for the program.

The replication process includes: identifying communities that will be able to replicate the Partners model, planning to use the model, implementing the model, and measuring success.

Your community will be a good candidate for replication if you have the following:

- support from agencies that provide, or are involved with, respite services,
- knowledge of current respite funding to support families without financial resources, and
- support and commitment from local early intervention programs, the school system, mental health or mental retardation programs, and/or family support programs

Partners Plus Community Planning Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127
The Replication Process

Partners Community Implementation Process

Identify Community Site

Plan for Replication of the Partners Model
- Create Community Support
- Determine Program Advisors
- Determine Administrative Structure
- Secure Financial Resources

Implement the Partners Model
- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families As They Train Their Own Caregivers
- Provide Continuing Support

Evaluate the Partners Program
The Replication Process

Each community needs to develop a plan to use the Partners model. Activities aimed at creating an awareness of family needs and the Partners model within a broad array of agencies, organizations, and individuals helps build a variety of community supports. The creation of an advisory committee brings expertise and resources that will ensure a quality program. Planning includes finding a home for the Partners program and knowing who is going to be responsible for implementing the program. Sometimes formal or informal interagency agreements will provide sufficient support, while in other communities new financial or personnel resources may be necessary for community replication.

This manual provides strategies and materials to help communities plan for and implement the Partners family-centered respite care program. CDR's Partners Plus has developed three additional manuals to be used in conjunction with this one. The Trainer's Workshop Manual enables community trainers to conduct the 6-hour Partners workshop for families and caregivers (the foundation of the Partners model of respite care). The Trainer's Workshop Manual includes the training content, sample agendas, overheads, and additional information that enables trainers to conduct an effective workshop for families and caregivers.

The Family and Caregiver Manuals are provided to each workshop participant, and are the basis of the workshop training content. Both manuals are designed specifically for families and caregivers to be used as resources during, after, and in absence of attending a Partners workshop. These manuals provide their readers with helpful
information about respite, caring for children with special needs, and building long-term respite relationships.

Communities that replicate the Partners model gather information from families and caregivers to measure the success, the quality and usefulness of training and support.

The Partners replication process will guide you in developing a family-centered respite program and in providing Partners training for families and caregivers. In planning for your Partners program, there are four major tasks: creating community support; determining program advisors; determining the administrative structure; and securing financial resources. A checklist is included in Section 2 to help your community plan your Partners program.

Planning Your Partners Program

- Create Community Support
- Determine Program Advisors
- Determine Administrative Structure
- Secure Financial Resources
Creating Community Support

The first task in planning for your Partners program is to create community support. A broad base of community support will ensure success for your Partners program. The design of your program will depend upon your community and its resources. A number of community agencies, organizations, and individuals may be involved in planning and implementing Partners.

Ideally, more than one agency will take part in planning and will contribute financial and other resources to the development of the program. Agencies and organizations may provide personnel, office space, and materials, workshop trainers, sponsor social events, or underwrite child care costs. Individuals may volunteer time, serving as advisors to the program or as trainers.

Because maintaining good community relationships is so important, it is an excellent idea to keep track of all communication. Provided in Section 2 is a sample form that may help your community maintain a record of communication.

The following pages offer some helpful hints in creating community support for your Partners program.

Consider the following questions when searching for community resources:

- Who in the community serves families of children with special needs?
- Which agencies or groups provide financial assistance for respite services?
Creating Community Support

- What parent organizations or support groups exist?
- Are there civic or religious groups that support special projects?
- Are there sororities or fraternities that support special projects?
- Who are the influential members of your community?

Ask some of the following groups/organizations to provide resources or supports:

- Educational institutions or organizations
- Civic groups or religious organizations
- Parent organizations or support groups
- Disability advocacy groups
- Early intervention or public school programs
- State and local service agencies
- Hospitals and health organizations
- Local physicians or other health care professionals
- Local interagency coordinating councils
- Other interested groups or organizations.
Creating Community Support

Ask community partners to make any of the following contributions:

- Office space for personnel and materials
- Secretarial support
- Personnel/individuals to conduct training
- Space for training workshops and activities
- Funding to assist families in obtaining respite (e.g., stipends, vouchers)
- Stipends to caregivers for individual or group training
- Refreshments for group training
- Duplication of training materials
- Educational opportunities and resources
- Social opportunities for families and caregivers
- Promotional/awareness activities
- Transportation to training or social events
Determining Program Advisors

Community members and consumers of your services can participate as advisors to your respite program. The roles and responsibilities of your advisors can be to help plan your program, to market your services, to review materials, or to evaluate the program's success. Advisors can be individuals, focus groups, an independent committee, or a board of directors.

Advisors should have knowledge, experience, resources, and/or be influential community members. As advisors, the expertise of families and caregivers is essential. It will also be helpful to include individuals with skills in financial planning, community relations, or strategic planning. You may need representatives from local or state agencies, or individuals from the business community who have expertise in budgeting, finance, and funding. Your advisors can serve as a talent pool from which to draw future resources.

Focus groups can help you plan or review your program. Focus groups are frequently made up of consumers. A sample letter inviting parent participation in a focus group is included in Section 2.

Advisory committees may help to garner community support for your program and help to
Naming Program Advisors

make decisions about how to plan and carry out your services. To ensure a family-centered program, families should comprise the majority of committee members. Other community stakeholders to be included may be from the fields of health, recreation, education, social services, early intervention, or from faith groups, civic groups, etc.

Advisory committees/groups are different from the board of directors for organizations. A board of directors is legally responsible for the agency or program, its management, and finances. A governing board sets policies. Advisory committees are not legally responsible for an agency, but can assist governing bodies in all aspects of management, except policy development. Advisory committees can provide objective program evaluation, develop strategies to overcome problems, and provide resources and expertise to support the program.

Included in Section 2 are sample roles and responsibilities of program advisors, an advisory committee agenda, and an advisory committee work plan.

Planning Your Partners Program

- Create Community Support
- Determine Program Advisors
- Determine Administrative Structure
- Secure Financial Resources
Determining Administrative Structure

Whom will your program serve? How will your program be staffed? These are some critical questions about the administrative structure of your Partners program that will need to be answered in your planning.

**Whom will the program serve?**

Defining whom your program will serve may be critical in planning and establishing your Partners program. Defining your training audience will help in determining what community agencies and individuals will be involved and to what extent each will commit resources.

There are several characteristics of consumers of respite to consider, such as age, disability, the family income, or place of residence. For example, if you are going to serve children with mental retardation, look for support from agencies or organizations with special interest in supporting this population such as the Arc. If you are going to serve individuals with special health care needs, look to hospitals or health care agencies for support.

**Caregiver characteristics such as knowledge, skills, and abilities** may also influence decision making about the program. For example, if you choose to target trained child care providers, you may need to pay for their participation in group and/or personalized training.

Eligibility of respite consumers or caregiver characteristics may influence your ability to access state and local monies,
Creating Program Structure

grants, or donations. Funding sources may limit the number, or age, of clients served. Early planning about whom you will serve will ensure a clear focus in awareness materials and promotional activities. Clear eligibility parameters will help to attract appropriate consumers of your services and providers of respite care.

What services will the program provide?

Your Partners program will need to identify the types of services that will be available to support families and caregivers. Using the Partners model will mean that your program will at a minimum provide: Partners workshops for families and caregivers; support to families as they train their own caregivers; and continuing support to both families and caregivers. Continuing support may consist of additional services, such as:

- phone consultations;
- home visits;
- meetings with families, caregivers, and/or community groups;
- a resource library;
- social activities;
- support groups and/or a parent cooperative for families and/or caregivers.

When deciding which support services your Partners program will provide, you will need to consider your financial and personnel resources, and supports currently available in your community. In Section 2, there is a discussion about providing continuing educational and social supports and services.
How will your program be staffed?

Once you know the agency or group that will administer your Partners program, decide whether you need to hire a program coordinator or whether that role can be added to the responsibilities of existing personnel. Having one person to monitor and coordinate Partners activities will help ensure an efficient and effective respite program. Program advisors often have the expertise to serve as a resource to the coordinator, especially during the initial phase of program development.

The coordinator's first responsibilities may be to identify key agencies, groups, or influential people in your community to serve on the board of directors or, more typically, on the advisory committee. The coordinator may need to seek additional community support, write funding proposals, if the program will exist as a new or separate agency, or begin a public awareness campaign, in addition to seeking the help in developing the initial work-scope goals and in determining roles and responsibilities.

Coordinator roles and responsibilities may include developing awareness materials, coordinating group workshops and personalized training, purchasing materials, and collecting data for evaluation of services. The coordinator is often responsible for hiring and supervising Partners workshop training staff. He or she will supervise volunteer and/or paid personnel and ensure that they are educated in the method and content of training.

It is helpful to have at least two trainers to conduct Partners workshops. A parent-professional team of trainers is highly
**Creating Program Structure**

**Recommended.** A family member’s stories and perspectives can be extremely helpful for caregivers to hear. A parent or sibling, in the trainer’s role, can contribute when family participants are reluctant to talk or can facilitate valuable discussion between families and caregivers.

Professionals from a variety of fields such as social work, education, health, counseling, etc., may be involved in Partners. It is helpful for your pool of professionals to have experience in training and knowledge of adult learning. Resources on adult learning are provided in Section 7. Additional information on learning styles and training tips can be found in the *Trainer’s Workshop Manual*.

Professionals and family members can also provide additional training to families and caregivers in related topics to enhance knowledge and skills. Topics that may supplement a Partners workshop include specific disabilities, adult learning, advocacy skills, or activities for young children. You may want to encourage family members and/or caregivers to be trained in First Aid and CPR. An additional list of topics that may compliment your basic Partners training is included in Section 2.

**Securing**

<table>
<thead>
<tr>
<th>Planning Your Partners Program</th>
</tr>
</thead>
<tbody>
<tr>
<td>• <strong>Create Community Support</strong></td>
</tr>
<tr>
<td>• <strong>Determine Program Advisors</strong></td>
</tr>
<tr>
<td>• <strong>Determine Administrative Structure</strong></td>
</tr>
<tr>
<td>• <strong>Secure Financial Resources</strong></td>
</tr>
</tbody>
</table>

*Partners Plus Community Planning Manual*  
Child Development Resources  
P.O. Box 280, Norge, Virginia 23127  
DRAFT 10.99
Securing Financial Resources

Financial Resources

Although needs will vary based on your community and planned work scope, your Partners program will need personnel, facilities, materials, and other resources. Although some of these may be donated by agencies and organizations it is important to plan adequate funding to support your budget.

Funding for your Partners program may come from a variety of public and/or private sources. Public agencies such as local family support services or early intervention programs may supply charitable giving or funds for the provision of respite care. In addition to agency support, you may access corporate and other grants, personnel time, facilities, materials and even funds for respite care. Local groups such as interagency coordinating councils, civic and religious groups, and interested individuals may be possible contributors or volunteers for your program.

Besides personnel costs, facilities often require the greatest financial commitment. Your Partners program needs one place to call home -- a central place where families and caregivers can call or visit to obtain information or assistance. It is preferable to have one location for Partners staff to hold meetings and keep materials. You will want to consider several factors when choosing a home for your program including the mission, the reputation, and the logistics of the agency or organization, i.e., is this building location accessible to public transportation etc.?

Materials and resources needed may include flyers to announce Partners workshops, child care during workshops, and refreshments for participants. Caregivers and families
Securing Financial Resources

may need educational resources and stipends. Criminal or background checks on caregivers will help to ensure that children and families feel safe in developing new relationships with caregivers. If your community has the financial support necessary, a toll-free 800 phone number could also be a valuable resource to families and caregivers. Plan your budget in advance so you will have enough financial resources to cover your costs. A sample budget for your Partners program is provided in Section 2.

Consider the following sources of funding for your program:

- Grants from public agencies at the state and local levels
- Private foundation grants
- Donations from civic, religious, or other groups
- Donations from individuals, memorials
- Fund raising events
- Family support funds through state and community agencies
- State or local interagency coordinating councils
- Funds available from early intervention or other statewide services
Consider the following questions when selecting a home for the program:

- Does the agency or group provide respite or other services for children and families?
- Does the agency or group have a family-friendly image in the community?
- Is the facility easily accessible to families and caregivers?
- Are there supports or personnel available to answer inquiries and provide information?
- Can this agency or group serve as a central point of contact for the program?
- Is there a natural fit between the Partners program and the potential home?
Notes on this Section
Section 2

Sample forms and information for planning your program
Sample Forms and Information

Sample Forms and Supplementary Information

- Developing Your Partners Program: Replication Checklist
- Project Inquiry Form
- Individual Contact Log
- Communication Log
- Letter of Invitation
- Advisory Committee Roles and Responsibilities
- Advisory Committee Organization Meeting Agenda
- Advisory Committee Workplan
- Supplementary Training Topics
- Budget
Developing Your Partners Program: Replication Checklist

<table>
<thead>
<tr>
<th>Activity</th>
<th>Person Responsible/Comments</th>
<th>Date Complete</th>
</tr>
</thead>
<tbody>
<tr>
<td>Select local planning group (LPG) members</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Review Partners model</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Review Partners replication process</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Secure financial resources for:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• the Partners program</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• a home</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• materials</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Identify program advisors (advisory committee/group)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Define whom your program will serve</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hire and/or select:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• program staff</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• coordinator</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• trainers</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Partners Project Inquiry Form

Use this Sample Partners Project Inquiry Form to track families, caregivers, and others interested in participating in your Partners project.

Caller Information:

Date: __________________________ Message taken by: __________________________
Name: __________________________

___ Family  ___ Caregiver  ___ Agency

Age of child: _______________________ Agency (if applicable): ______________________
Address: __________________________

______________________________
______________________________
City/County State ________________ Zip: __________________________
Phone (Work): ____________________ Fax: __________________________

Reason for Inquiry / Call:

___ Workshop information
___ Respite funding
___ Other (please specify):

___ General respite information
___ Request for speaker/presenter
___ ______________________
___ ______________________

How did you hear about Partners?

___ From a family
___ From a caregiver
___ Project staff
___ Television

___ Newspaper article/press release
___ Community agency
___ Public awareness presentation
___ Other (please specify): __________________________
___ ______________________

Partners Plus Community Planning Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99 2-5
Follow-Up:

- Send Partners identity packet
- Send agency identity packet
- Send other materials
- Refer to other community resources
- Register for Partners workshops
- Add to mailing list
- Send copy to: ________________________
- File

Comments:

________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
## Partners Communication Log

*Use this Sample Partners Communication Log to track communication with families, caregivers, respite stakeholders, etc.*

<table>
<thead>
<tr>
<th>Name</th>
<th>Date</th>
<th>Mode*</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Mode Code: P=In Person; M=Mail; T=Telephone; E=E-mail; F=Fax*
Sample Introductory Letter to Families Informing Them of Partners

How to use this sample letter:

Prior to hosting your first Partners workshop, you may want to meet with families to inform them of your community's new Partners program. Meeting with families will give you a sense of their interest as well as times and locations for workshops that are best for them. Adapt this with your communities information, filling in the blanks, or altering as needed.

Dear:

_________________________ has a new program—Partners.

Partners is designed to increase respite (temporary childcare) options for families of children with special needs. The program helps families identify and train their own respite caregivers and develop a system in the community to ensure continuation of respite services. Partners is a family-centered program that involves families in all phases of development.

Families will have the opportunity to be involved in many ways. One way is to participate in a focus group where you can share your ideas with members of the program staff. Since our first workshop for families and caregivers will begin soon, we are inviting you and several other families to participate in a focus group to share ideas on how we can best conduct and design trainings for families and caregivers.

We have selected the following date, time, and location for a meeting:

_________________________

A member of the program will call you in a few days to see if you are interested in participating. Please feel free to call us as well to let us know if you will be able to attend.

The Partners staff are very excited about this new program and looks forward to meeting with each of you as we work together to expand respite options for families.

Sincerely,
Sample Partners Advisory Committee: Roles and Responsibilities

Purpose:

To provide the Partners program with special guidance and counsel.

Member’s Term: One Year

Composition:

The committee members consist of 51% family and caregiver representatives, and remaining 49% community representatives.

Meeting Attendance:

The Advisory Committee meets six times a year in alternating months.

Responsibilities:

• Become informed about the program’s goals and activities
• Promote Partners within your personal associations and to the community
• Assist in special problem-solving for the project
• Provide ideas and review program and materials development
• Assist in recruiting families and caregivers for respite
• Contribute knowledge and share experience
• Develop a plan for continued community support for continuing respite services
• Assist with project evaluation activities
Sample Partners Advisory Committee Organizational Meeting Agenda

Date
Time

Agenda

I. Welcome and Introductions

II. About Partners
  • What is respite?
  • Project goals
  • Project activities

III. About the Advisory Committee
  • Roles and responsibilities
  • Workscope review
  • Specific tasks and assignments
  • Getting started
  • Meeting schedule
**Partners Advisory Committee Workplan**

### Objectives

1. **To become informed about Partners goals and activities.**
   - 1.1 Attend Advisory Committee orientation meeting.
   - 1.2 Attend scheduled meetings on the 4th Monday of alternating months.
   - 1.3 Review Partners materials and advisory committee minutes in advance of meetings.

2. **To work toward building a strong base of support and stakeholders within the community.**
   - 2.1 Promote Partners within personal contacts and community groups.
   - 2.2 Recruit 4-6 additional members to represent families and community.
     - a. Contact churches and pastoral care associations.
     - b. Invite participation by families who have received Partners training.
     - c. Contact civic groups that are focused on children's issues.
   - 2.3 Participate in Partners training workshops and presentations.

### Activities

<table>
<thead>
<tr>
<th>Activities</th>
<th>Initiation</th>
<th>Party Responsible</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1 Attend Advisory Committee orientation meeting.</td>
<td>Date</td>
<td>Coordinator, Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>1.2 Attend scheduled meetings on the 4th Monday of alternating months.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>1.3 Review Partners materials and advisory committee minutes in advance of meetings.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>2.1 Promote Partners within personal contacts and community groups.</td>
<td>Date</td>
<td>Partners Staff, Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>2.2 Recruit 4-6 additional members to represent families and community.</td>
<td>Date</td>
<td>Coordinator, Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>a. Contact churches and pastoral care associations.</td>
<td>Date</td>
<td>Partners Staff</td>
<td></td>
</tr>
<tr>
<td>b. Invite participation by families who have received Partners training.</td>
<td>Date</td>
<td>Partners Staff</td>
<td></td>
</tr>
<tr>
<td>c. Contact civic groups that are focused on children's issues.</td>
<td>Date</td>
<td>Coordinator, Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>2.3 Participate in Partners training workshops and presentations.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
</tbody>
</table>
### Partners Advisory Committee Workplan continued

**Objectives**

3. To assist program staff in refining the model and revising training and public awareness materials.

4. To recruit interested families and caregivers to Partners training.

5. To secure community resources for project continuation.

#### Activities

<table>
<thead>
<tr>
<th>Objective</th>
<th>Activities</th>
<th>Initiation</th>
<th>Party Responsible</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.1</td>
<td>Review and comment on Partners workshop flyers.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>3.2</td>
<td>Review and comment on Partners project brochure.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>3.3</td>
<td>Review and comment on Partners training manuals.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>3.4</td>
<td>Problem solve at meetings with staff to discuss model implementation issues.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>4.1</td>
<td>Generate listing of potential target groups.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>4.2</td>
<td>Identify contact persons.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>4.3</td>
<td>Assist project in making presentations.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td></td>
</tr>
<tr>
<td>5.1</td>
<td>Support parent network for exchanging/bartering respite.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td>Partners Staff</td>
</tr>
<tr>
<td>5.2</td>
<td>Assist in developing respite co-op.</td>
<td>Date</td>
<td>Advisory/Partners Staff</td>
<td></td>
</tr>
<tr>
<td>5.3</td>
<td>Explore home for Partners program.</td>
<td>Date</td>
<td>Advisory Committee</td>
<td>Partners Staff</td>
</tr>
</tbody>
</table>
Examples of Supplementary Training Topics

- First Aid
- CPR
- Disability specific workshops
- Communication/sign language
- Behavior management
- Positioning and handling techniques for caregivers caring for children with special needs
- Feeding issues for children with special needs
- Rehabilitation equipment (e.g. wheel chair, braces, etc.)
- Adapting toys for children with disabilities
- Talking to others about special needs
- Advocacy/Legislation awareness
- Activities to do with children
- Child development
- Relationships
- Principles of adult learning
- Training tips
Sample Partners Budget

The following is based on four training workshops per year. Additional budget considerations are listed on the back.

<table>
<thead>
<tr>
<th>Expenses</th>
<th>Funds Available</th>
<th>Donated / In-Kind</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trainers' Salaries</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parent (40 hours @$15.50/hr)</td>
<td>$100.00</td>
<td>$520.00</td>
</tr>
<tr>
<td>Professional (40 hours @$20/hr)</td>
<td></td>
<td>$800.00</td>
</tr>
<tr>
<td>Coordinator (40 hours @$15/hr)</td>
<td></td>
<td>$600.00</td>
</tr>
<tr>
<td>Payment to Caregivers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 hours respite child care</td>
<td></td>
<td></td>
</tr>
<tr>
<td>during personalized training</td>
<td>$772.50</td>
<td></td>
</tr>
<tr>
<td>(15 @$5.15/hr)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 hours child care for families</td>
<td></td>
<td></td>
</tr>
<tr>
<td>during workshop (6 caregivers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>@$5.15/hr for 6 hours)</td>
<td>$741.60</td>
<td></td>
</tr>
<tr>
<td>Materials/Supplies</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Manuals purchased from CDR</td>
<td>$600.00</td>
<td></td>
</tr>
<tr>
<td>for 4 workshops (20 family</td>
<td></td>
<td></td>
</tr>
<tr>
<td>and caregiver @$15)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Refreshments at workshops</td>
<td>$75.00</td>
<td></td>
</tr>
<tr>
<td>Postage, phone, utilities</td>
<td></td>
<td>$400.00</td>
</tr>
<tr>
<td>&quot;All Kids Like Cookies&quot; props</td>
<td>$50.00</td>
<td></td>
</tr>
<tr>
<td>Training</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CPR/First.Aid (10 caregivers)</td>
<td></td>
<td>$300.00</td>
</tr>
<tr>
<td>Facility</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Home office space (@$600/month)</td>
<td></td>
<td>$7,200.00</td>
</tr>
<tr>
<td>Office supplies (@$250/month)</td>
<td></td>
<td>$3,000.00</td>
</tr>
<tr>
<td>Workshop training space</td>
<td></td>
<td>$800.00</td>
</tr>
<tr>
<td>(2 rooms @$100 each)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>$2,339.10</strong></td>
<td><strong>$13,620.00</strong></td>
</tr>
</tbody>
</table>
Other budget considerations:

Office equipment/furniture, advertisements, Xeroxing of training materials (agenda, ice-breakers), maintenance of the caregiver directory (personnel time, postage to mail to families), resource library, stipends for supplemental training for families and/or caregivers (optional), money to support actual respite (optional), facility use for social events, and any background/criminal history checks (optional).
Section 3
implementing your Partners program
Recruiting Families and Caregivers

In order to recruit families and caregivers to participate in a workshop, your Partners program will need to use a variety of marketing strategies. You will want to have written awareness materials, news releases, and community announcements designed to reach specific community members.

In addition to helping with recruitment, marketing strategies will enhance community support for your program. Influential community members who could provide financial and personnel resources should be considered when developing promotional materials. Keep in mind who will be hearing, or reading about your program. Your approach and materials will vary depending on whether your audience is families, advisors, community agencies, or funding sources. Gear your marketing strategies to one or more specific target audiences.

Since one of the best marketing strategies is meeting and talking with people, provide information about Partners and its training during regular meetings and telephone conversations with individuals in the community.

Implementing the Partners Model

- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families as they Train their Own Caregivers
- Provide Continuing Support
Recruiting Families and Caregivers

Public speaking informs a large number of people at one time and can immediately address their questions about the program. Present an overview of your program to community groups including religious, civic, advocacy and parent groups. Speak to agencies, colleges and universities, and health clinics. Invite community members to an orientation meeting to introduce your Partners program.

Direct mailing is a good way to reach people. Send brochures, flyers announcing workshops, and other promotional materials through the mail again targeting specific audiences such as families and referral sources. Include a telephone number to call and/or a registration form to mail back so that individuals can easily sign up for a workshop. If your budget allows, you may choose to use a graphic artist and professional printer or desktop publishing software for promotional materials.

Many agencies and groups produce newsletters. Your program may choose to produce a newsletter or submit announcements or articles to community newsletters. Some communities have parent newsletters or magazines that may publish feature articles.

Use the media to promote your program. Newspapers are likely to publish an interesting photograph with a caption. You may want to collect photographs and/or videos of events for future media coverage and awareness activities. Remember, when you take photographs and submit them to the media, you will need permission from the people in your pictures or videos. Sample release forms are included at the end of this section. A scrapbook for your program's history and events can be taken to future conferences and other public speaking engagements.
Recruiting Families and Caregivers

Families can recruit caregivers to participate in Partners workshops. Encourage families to tap into their informal supports (extended family members, friends, neighbors, and community programs) to train as their respite caregivers. People from within their own natural support network may be reliable and accessible over the long term. However, families with limited natural support may need your help to identify a caregiver. A directory of caregivers, and social events where families and caregivers meet, are ways in which the program can facilitate family and caregiver matches.

There are a number of strategies you can use to encourage caregivers to participate. Try using incentives, such as paying caregivers for their time to attend Partners workshops. An hourly rate, such as minimum wage, may attract child care professionals. Offering child care to caregivers as well as to families may also be an incentive for caregivers to attend Saturday or evening workshops.

Another powerful incentive is to pay caregivers for the time they spend with families in personalized training. In other jobs, employees are paid for their time spent in training. It should be no different for respite caregivers. Although the time spent in training varies some, families and caregivers spend up to ten hours learning together before respite begins. However, paying for training time may be a financial burden for some families. It is helpful if a community Partners program can pay caregivers for this time or help families gain access to available public funds to support training.
Strategies to promote awareness of your program:

- Brochures describing the program
- Flyers announcing training
- Paid advertisements on radio, TV, or in the newspaper
- Press releases for events
- Feature articles about families or caregivers
- Direct mailings to specific audiences
- A newsletter or newsletter articles
- Presentations at local agencies, meetings, or events

Strategies to recruit caregivers and families:

- Ask families to bring caregivers to training
- Place classified ads in local newspapers
- Distribute brochures and flyers to families through public schools
- Distribute brochures and flyers to agencies, organizations, and groups
- Post bulletins in public locations such as grocery stores, malls, churches, etc.
- Present at early childhood classes at colleges and universities
Conducting Partners Workshops

The purpose of the six-hour Partners workshop is to help families and caregivers understand the Partners model and family-centered respite. During the workshop, families develop skills in screening, interviewing, and training caregivers. Caregivers develop a basic understanding of caring for children with special needs. For families who do not bring their own caregiver to the workshop, this can be an opportunity for families to conduct an initial caregiver interview and begin their respite relationship.

Good planning will ensure that your six-hour workshop is a success. Planning for the workshop involves identifying and arranging space; compiling or obtaining Family and Caregiver Manuals; providing refreshments; and coordinating child care for participants. A sample checklist, included in Section 4, helps workshop trainers organize all the necessary information and materials for Partners workshops.

The Partners Trainer’s Workshop Manual (found in Families and Caregivers as Partners: A Guide to Family-Centered Respite Care) is an additional resource to help...
trainers conduct the six-hour workshop. It contains helpful hints for how to conduct a Partners workshop, a script of what trainers can say, overheads, and examples of activities to break the ice and conclude the workshop. A flowchart to the Trainer’s Workshop Manual is located on the following page.

In preparation for the six-hour workshop, consider the following:

- preparation,
- materials,
- training location,
- refreshments,
- child care,
- training tips, and
- evaluation measures.

Preparation

If your marketing strategies were effective, you should have a group of families and caregivers eager to attend a Partners workshop. As individuals express interest in your Partners program and/or in attending a Partners workshop, keep a record of their name, phone number, address and other pertinent demographic information. This will enable you to complete a mailing or in some way notify individuals about workshop dates and other Partners related information and/or activities.
Conducting Partners Workshops

Flow Chart to the Trainer’s Workshop Manual

Introduction
The first section of the Trainer’s Workshop, Family, and Caregiver Manuals have the same Introduction.

How To Use the Trainer’s Workshop Manual
The second section of the Trainer’s Workshop Manual tells the trainer how to use the manual.

Welcome to the Partners Workshop
Workshop information for all participants begins in the Trainer’s Workshop Manual at this section.

Sections 1-4: Participants Work Together
All workshop participants work together during sections 1, 2, 3, and 4, where the information is the same for both families and caregivers:

Section 1: About Respite
Section 2: Building Partnerships: Communication
Section 3: Building Partnerships: Families and Caregivers Working Together
Section 4: All Kids Like Cookies Activity
Sections 5-10: Participants Work Separately
After completing section 4, the group splits into families and caregivers, each working with a trainer on topics that are different:

**Family Break-Out Session**
Section 5-F: Finding Respite Caregivers
Section 6-F: Interviewing Caregivers
Section 7-F: Developing a Personalized Training Plan
Section 8-F: Resources
Section 9-F: Forms for Sharing Information

**Caregiver Break-Out Session**
Section 5-C: Disabilities Awareness
Section 6-C: Understanding Child Development
Section 7-C: Children with Special Needs
Section 8-C: Becoming a Caregiver
Section 9-C: Resources
Section 10-C: Forms for Sharing Information

**Final Activity: Participants Work Together**
After completing their sections, the two groups return for the Final Activity, involving daily routines to help develop a personalized training plan.
Conducting Partners Workshops

Maintaining a registration list for the workshop can help you plan for the event. You will be able to limit the number of no-shows if you use some simple strategies, such as calling or sending a postcard as a reminder to those who register. A small fee can be used to ensure registrants actually attend. However, many programs choose not to charge families to enroll in workshops. An alternative strategy to ensure good attendance is the use of a refundable registration fee.

To prepare participants for the workshop you may want to remind them about workshop content, times, locations, and about items they should bring for children who will be in child care during the workshop. Families who bring children should plan to come at least twenty minutes early to meet with their caregivers and become familiar with a new environment. A sample workshop agenda included in Section 4 can be distributed prior to your Partners workshops.

Materials

CDR has developed Family and Caregiver Manuals that participants will be able to use during and after the workshop as a resource. Families will depend on their manuals to help them develop personalized training plans for their caregivers. Manuals for each workshop participant may be purchased from CDR, or one master set can be purchased and duplicated by your Partners.
program for each participant. You may wish to compare
the costs in personnel and resources to determine which
method of obtaining manual materials is best for your
program.

You’ll need **Additional supplies** such as name tags, pencils,
pens, and stickers. Name tags help trainers learn
participants’ names and facilitate discussion between
families and caregivers. Using different colors for caregiver
and family name tags will help families identify caregivers
and help trainers separate participants into small groups
for activities during the workshop. It is helpful to supply
participants with pens or pencils and highlighters to use
during the workshop. You may want to purchase pens or
pencils with your logo, phone number, and address on
them as a promotional tool for your Partners program.
Pencils or other promotional items can be used as
incentives for workshop participants who volunteer to
participate in activities. You can give highlighters to all
participants or just place a few at each table to be shared.
Participants also like getting colorful tags or sticky dots to
use as markers in sections of the manual, and extra paper
or sticky pads to jot down ideas, comments, and/or
questions.

*Bringing books, videos, and other resources* or an
annotated bibliography to the workshop shows
participants what materials they can borrow or purchase.
If you plan to make resources available on loan, be sure to
establish a catalog system that allows you to track them so
materials will be returned.
Conducting Partners Workshops

Workshop Location

Think about the location where your workshops will be held. If your program serves a large geographic area, you may want to use several locations to make it easier for participants to attend workshops. Make sure the buildings you conduct your workshops in are centrally located and physically accessible for participants with disabilities.

There may be workshop space available in your building or other nearby locations, at no cost to you. However, if space is not readily available, you may want to explore the use of local churches, synagogues, schools, or other facilities. Often, space can be found in the community free of charge or for a small donation.

It is necessary to have access to at least two rooms for adult use and a place for child care if you are offering that service. One of these rooms should be large enough to accommodate the entire group. The other, for use in small group activities. Churches or synagogues work well if you provide child care because many have nurseries and/or a kitchen for preparing refreshments. Early intervention programs, schools, child care centers, or recreation centers often have rooms and materials appropriate for child care.

The space you use should be comfortable (e.g., air conditioned/heated) and have access to telephones and rest rooms. When arranging the space for the workshop, consider size of the group and arrange the tables and chairs so that participants can easily see the trainers and...
also talk freely with others. For a group of approximately 20 participants, tables can be arranged in a U-shape. For larger groups, circular tables can be used. Groups larger than 30 are not recommended for Partners workshops. Due to the nature of small group activities in the workshop and break-out sessions designed to foster participant sharing, smaller groups (under 30) are more conducive to meeting Partners workshop objectives.

If your Partners program plans on holding workshops in a variety of locations to help reach as many people as possible it is helpful to have suitcases with wheels for transporting resources, manuals, and other materials easily. When you travel, it is important to plan ahead and make sure you have packed all the materials and items you will need.

If you use borrowed space, you may not know what resources are available within the building, or how much time and effort it will take to arrange the rooms to meet your needs. For this reason, it is helpful to be able to set up the day before the workshop. If that is not possible, plan to arrive at the location in time to arrange the room and set up the workshop, activities, resources, and refreshments.

Be sure to give participants clear directions to workshop locations. A copy of a map or specific written directions are helpful in ensuring that participants can find their way.
Conducting Partners Workshops

Refreshments

Depending on the time or length of your workshop, you may decide to offer snacks and/or lunch for participants. To reduce costs, your program may ask participants to bring their own lunch. Other options are to have participants leave for lunch and then return, or to schedule workshops over two evenings or two mornings to avoid a meal altogether.

Having families, caregivers, and their children eat a meal together has proven to be successful in helping families and caregivers match. Not only do participants have an opportunity to meet each other, but families and potential caregivers have an opportunity to watch caregivers interact with children. For some potential caregivers, this may be the first time they meet children with special needs and talk with their families.

Child Care

Without child care, many families will be unable to participate in your Partners workshop. One way to help families attend is to provide quality child care. Child care is also an opportunity for recently trained caregivers to obtain hands-on experience caring for children with special needs. The group care experience gives families an opportunity to meet and observe potential caregivers, thus facilitating family and caregiver matches. By providing child care at workshop sites, you are providing caregivers with a chance to practice their skills with the support of having families and other caregivers present. When
arranging child care it is helpful to select at least one seasoned caregiver to serve as a mentor for new and/or inexperienced caregivers in group care.

Workshop participants should preregister their children so you will know how many children will be attending and how many caregivers you will need to provide care. You will need to have one or two extra caregivers available for unexpected arrivals. Or be prepared to tell participants who have not preregistered that you cannot provide care for more children than anticipated.

In order to ensure successful group care experiences for children, families, and caregivers, you must use appropriate child care settings. Make sure the rooms will accommodate the number of children's activities appropriate for their ages, and any special accommodations that may be needed. For example, children with wheelchairs will need space to maneuver.

Families should be encouraged to come early to help familiarize their children with the setting and to share necessary information about caring for their children. It is beneficial to both families and caregivers to be nearby during the workshops, if help or guidance is needed. This is especially true for caregivers who are newly trained to care for children with special needs.

You may be able to find caregivers who are willing to volunteer their time in order to gain a valuable and satisfying experience. If not, you will need to pay caregivers through your program funds or seek support from other sources.
Conducting Partners Workshops

Workshop Tips

Good training combines lecture, audiovisuals, and individual and group activities. In the Trainer's Workshop Manual, there are hints for conducting the workshop and information about adult learning. Reviewing this material will help ensure that your workshop is appropriate to varied adult learning styles and preferences.

During the workshop, real-life illustrations convey content in meaningful ways. The Trainer's Workshop Manual provides stories from real families. Trainers who have family members with disabilities can tell their own stories.

Here are some tips on making training more meaningful and enjoyable:

- Have knowledgeable trainers who are well prepared
- Develop and follow an agenda
- Foster a respectful, open, and informal atmosphere
- Use a little humor and cartoons to illustrate points
- Encourage discussion and an open environment for interaction
- Include fun activities and energizers

Partners Plus Community Planning Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Conducting Partners Workshops

Measuring Your Success

In order to ensure the quality of the workshop, Partners workshop trainers need feedback from participants about their presentation styles and their use of the Partners materials. Participants may rate or comment about the organization of the workshop, the helpfulness of presenters, the usefulness of materials, or what they learned during the workshop. A sample workshop training evaluation is included in Section 6.

It is useful to let workshop participants know how important and valued their comments are and to thank them in advance for taking the time to provide you with feedback. Participant comments will inform you about needed changes in your presentation.

In addition to evaluation of the six-hour workshop, it is extremely useful for Partners programs to receive feedback regarding the success of your respite program. You may wish to collect pre and post knowledge and comfort measures to determine what caregivers have gained from participating in the workshop. Program evaluation data may be used to educate potential funding sources, advisory committee members, the media, or other people whom you would like to receive more information about the program. Program or model evaluation measures are discussed and sample instruments are provided in Section 6.
Choosing Caregivers

Helping Families Choose Caregivers

After the workshop, families who did not bring their own caregivers to training match up with trained Partners caregivers to begin personalized training. During the workshop, families think about what they will want from a caregiver and learn how to screen potential caregivers. Now, families interview caregivers to find out if their caregiving preferences and styles are similar.

Families may meet caregivers in the workshop and may begin interviewing in preparation for personalized training. Other families may try some additional strategies to recruit caregivers or decide to postpone personalized training until a later date.

Families who were unable to either bring their own family or friends to the workshop as caregivers or meet a caregiver at the workshop may want help facilitating matches. The Family Manual contains a list of community resources available to families to help them get in touch with potential respite caregivers. You may be aware of additional resources in your community that families can contact to help them find caregivers.

Implementing the Partners Model

- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families as they Train their Own Caregivers
- Provide Continuing Support
A Directory of Potential Caregivers

Your Partners program may want to supply families with a directory of trained caregivers. Your directory of potential caregivers should be regularly updated so that families can find potential caregivers to interview and train.

Computer Databases are easy to maintain and are extremely helpful for a variety of reasons. A database can store your list of potential caregivers and help you track workshop participants. You will be able to retrieve caregiver or family names based on their caregiving preferences or geographic locations. Databases can store large quantities of information, can sort the information in many ways, and can create reports and mailing labels. There are many database programs available, from simple and “user friendly” to elaborate and complex. Whether you choose to keep this information as hard copies in individual files or on a computerized database, we highly recommend gathering and storing this information.

Once you spread the word about Partners, people will be calling for additional information. It is important to record names, addresses, phone numbers, interests, etc. for your database. You may also want to record what type of help you gave the caller (i.e., questions answered, added to the mailing list, registered for a workshop, referred to another resource, etc.).

Your database will help you organize the information you collect from callers into a system that will allow you to identify families and potential caregivers who have called for information so you can notify them of upcoming
Choosing Caregivers

workshops. You may want to use your database to identify agencies who can distribute information about Partners to their customers or clients.

Some of the other information you might want to keep on file includes:

- Date training was completed
- Age of child(ren) in the family
- Disability of the child(ren)
- Caregiver certifications
- Agency affiliations
- Status (i.e., if they are a caregiver, family, or with an agency)

Background Checks

If your Partners program plans to provide a caregiver directory for families, you will want to make some decisions about screening and background checks.

Screening of caregivers may include criminal/sex offender background checks, reference checks, department of motor vehicles background checks, and/or child abuse registry checks.
Choosing Caregivers

Background check forms can usually be obtained from your local police department and the department of human or social services, typically for a small fee.

Your Partners program may routinely do criminal or other official background checks on all caregivers who participate in the workshop or who are registered with your program. You may prefer to do checks on caregivers only when requested by a family, or to provide families with information on how to do their own background checks.

Partners encourages all families to complete reference checks calling at least two former employers or other people who know the caregiver, excluding relatives. The family or staff from your program, should ask reference questions such as: How long have you known the person? How would you describe them? What are their strong points in working with children with special needs? A sample reference check form is included in the Family Manual component of Families and Caregivers as Partners: A Guide to Family-Centered Respite Care.

Decisions your program will need to make about screening caregivers:

- Will there be selection criteria for Partners caregivers?

- Will your program require any background information checks or reference checks? Who will pay for them?
Choosing Caregivers

- How much help will your program give families in checking criminal records or doing reference checks for caregivers?

- Will the program perform checks on caregivers who have been recruited by families or just on caregivers who have been recruited by the program?

- Will the program endorse caregivers or simply provide a directory of caregivers' names and phone numbers specifying that although they have attended training, families are responsible for ensuring a good match and/or performing background checks?

Matching Opportunities

_Educational or social events_ can help facilitate family and caregiver matches. Your program can host a variety of training events with topics such as first aid, stress and coping, or handling difficult behavior. Social events such as picnics, holiday parties, or children's festivals are more informal events that not only help families choose caregivers, but also help families meet each other. Family support groups promote networking among families.

Families who are uncomfortable calling strangers on a list may be interested in participating in Partners social events that promote family/caregiver matches. _Parents' Day Out_ is a respite opportunity for parents that, besides offering respite, brings potential caregivers, families, and their children together to get to know each other better.
Parents' Day Out can also be a learning experience for caregivers. Newly trained caregivers can watch and learn more about caring for children with special needs from experienced caregivers. Caregivers tell us that having the support of other caregivers in group care helps them to feel more confident in their own abilities to care for children with special needs. The group caregiving experience can also lead to increased frequency of caregivers responding to the respite needs of families who call them.

Group care works well when at least one experienced caregiver acts as a mentor or lead caregiver for the rest of the group. Mentor caregivers can help less experienced caregivers by showing them what to do and encouraging them to practice new skills.

Another option for matching caregivers is to choose to form a family cooperative. In a coop, families can provide respite for each other without exchanging money. Family support groups in which families get to know each other well are likely to foster coop arrangements.
Choosing Caregivers

Essential elements for hosting a "Parents' Day Out":

- Schedule date, time, and length of event
- Decide who should attend and if a fee will be required
- Inform families of the event and register families as they confirm
- Determine the amount of children attending
- Obtain adequate caregiver coverage, preferably of both new and experienced caregivers
- Find adequate, appropriate space for child care
- Have plenty of toys and supplies appropriate for each child's age and development available
- Encourage families to meet with caregivers before and after to share information and to get to know caregivers
- Ensure that families have an opportunity to observe caregiver and child interactions
- Schedule time for families, caregivers and/or Partners staff to debrief together afterwards
Supporting Families as They Train Their Own Caregivers

Families will vary in the extent to which they rely on your Partners program to help them through this next phase of training. Your program can provide guidance, technical assistance, and emotional support to both families and caregivers.

During the six-hour Partners workshop, families receive opportunities to learn and practice new skills. In addition, their manual will be a resource in personalized training of caregivers. The Family Manual contains a step-by-step process for personalized training and various forms that guide the family through the experience. Caregivers will also rely on their Caregiver Manual as a resource after group training. It will be helpful to your program's staff and trainers to be familiar with the Family and Caregiver Manuals in order to support families and caregivers as they begin personalized training.

Implementing the Partners Model

- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families as they Train their Own Caregivers
- Provide Continuing Support
Successful partnerships between families and caregivers result when both work toward building a relationship. As in any relationship, good communication between families and caregivers is important in building successful respite relationships.

Families need to spend enough time in preparing caregivers, so they feel comfortable in caregiving. Caregivers need to know that the family is willing to give them the support they need to be successful in their partnership.

A caregiver must have a genuine desire to help a family meet their respite needs and a willingness to learn from families. Caregivers who listen, and learn a family’s way of caring for their child, help that family feel comfortable in leaving their child. Caregivers who ask appropriate questions and follow through on instructions earn families’ confidence.

A good respite relationship is one of mutual respect. Respect is shown in appreciating others’ knowledge and experience, and being considerate of others’ time and energy. Using a little humor can help caregivers relax while learning the skills necessary to care for a family’s child. Allowing for flexibility in caring for a child is a way for families to show they value a caregiver’s experience.

Families must also show appreciation to their caregivers. A good first step is letting caregivers know how much having respite means to them. Remembering that caregivers have a need to spend time with their own families and friends shows respect for a caregiver’s personal life and will help families to keep their caregivers.
Providing Continuing Support

Your Partners program can support families and caregivers in a variety of ways. Often families and caregivers need only minimal support beyond the Partners workshop. Support can come as recognition for a job well done or as help in making a phone call.

Strategies for support can include telephone consultation, home visits, or even a postcard with a few words of encouragement. On occasion, you can support families by making a referral or creating linkages between the community. Referrals to other services or financial resources may help ensure respite. To achieve a successful and lasting program, consider what resources your community has available to ensure adequate support for families and caregivers.

It is critical to recognize your program's role in fostering successful respite relationships and ensuring long term partnerships. Flexibility in how your program supports families and caregivers is essential. Listen to families and caregivers and respond to their individual needs. Using families and caregivers as advisors will help make decisions about the kinds of support your program should offer.

*Your program's support may be educational, using strategies such as:*

- offering resource materials (i.e. a lending library), or
- providing additional training.
Providing Continuing Support

Your program’s support may be social using strategies such as:

- holding special events, or
- having family support group meeting.

Your program’s support may combine many strategies.

Materials Loan

Access to a resource library and the Internet is very beneficial for families and caregivers. Up-to-date technical information on disabilities and special health care needs is frequently requested. Families often request strategies for handling difficult behavior and coping with stress. Local libraries, public schools and early intervention programs may be able to provide both reading and loan materials and Internet access. If resources are not accessible through community agencies or programs, you will need to access other resources.

Implementing the Partners Model

- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families as they Train their Own Caregivers
- Provide Continuing Support
want to collect written materials for loan and keep them readily available for workshop trainers to take with them to display at workshops.

**Supplemental Training**

Educational opportunities related to caring for children with special needs can increase families' and caregivers' knowledge and skills and many families and caregivers find it helpful to attend training beyond the Partners initial six-hour workshop. While skill-building workshops are valuable to families and caregivers, costs may be prohibitive. Your Partners program may choose to provide funds for these activities, as a way of offering additional support to families and caregivers.

Caregivers are often interested in additional training related to the special needs of children with disabilities. When caregivers learn more about caring for children with special needs, they become even more comfortable in caregiving. Additional training may also involve certifying family members or caregivers in First Aid and CPR to increase their awareness of safety and lifesaving techniques.

Additional training benefits the program by facilitating possible matches between families and caregivers. Families and caregivers have found that additional training offers them more contact with each other, making it easier to get to know and to find ways to support each other during respite care.

Your program will need to decide whether and how much of its budget can support supplemental training. A Partners program can offer scholarships to families and caregivers for workshops.
Social Support Strategies

Families of children with disabilities or special health care needs share common interests and concerns, and often seek social support from one another. Families tell us how helpful it is to hear from another family who has a child with a similar disability or life experience. Your Partners program can help families to meet each other.

A Family Directory can help promote parent-to-parent support. A Caregiver Directory can promote both matches between families and caregivers and caregiver-to-caregiver support. These directories can be kept by the program or distributed to families or caregivers who want them. An individual who wants to be included in the directory should give permission to share his or her name, address, phone number, and perhaps a brief description of their family or caregiving experience. Sample release forms are included in Section 4.

Social events that include caregivers help facilitate matches with families and can build social supports for both families and caregivers. Families and caregivers can develop friendships in social contexts such as parent and child groups, pot luck dinners, and recreational events. Your Partner's advisory committee may designate a social committee that could obtain support from the community for social activities. For example, the committee could ask...
for donations or reduced-price tickets to a family event or theme park.

Many caregivers are interested in receiving support from other caregivers with similar interests and concerns for providing respite care. Regularly scheduled, monthly meetings for caregivers can offer additional training and learning through shared experiences. To defray costs, caregivers can take turns hosting meetings and take turns providing refreshments. Caregivers have said that having this kind of regular, continuing support helps them remain interested in providing respite to families.

Family Support Groups

Support groups help families unite to deal with a range of issues unique to families who have children with special needs. It is very helpful for families to have an updated listing of local support groups and available resources in the community. If local support groups do not exist, or are inadequate to meet your families' needs, then your Partners program can help families form a group.

Families who are likely to participate in a support group may be those who want:

• Social opportunities for themselves and their children

• Information about specific disabilities
Family Support Groups

- Information about how to plan for the years ahead
- Information about educational, health care or other community services
- Information about independent living and work opportunities for their children
- To know how to get respite

It has been great for me to come to training today and hear what it is like for other families. I don't know other families to talk to.

- Debbie Warren, 1996, Parent
Section 4

sample forms

and information

for implementing

your program

Community Planning Manual
Sample Forms and Supplementary Information

- Sample brochure
- Sample article
- Sample press release
- Sample flyer
- Implementing Partners Planning for Workshop Checklist
- Workshop Agendas
- Permission for Videotape with Sound/Photograph Use
- Presentation/Workshop Checklist
- Permission to Release Caregiver Name
- Permission to Include Information in Family Directory
You may want to develop a brochure to help people learn about your Partners program. You can easily develop a brochure by using desktop publishing software or by using the expertise of a graphic advisor and printer. Your budget will guide your choice to use one or the other.

Child Development Resources

For more information, contact Partners Plus:
Child Development Resources
1490 Government Road
Williamsburg, VA 23185
Phone: (757) 220-1168
Fax: (757) 253-1779
e-mail: cdrtraining@tez.net

Support for the project is provided under grant number H325X060096 from the U.S. Department of Education. Office of Special Education Programs. Points of view or opinions do not necessarily represent official views or opinions of the Department of Education.

"Think of a candle that slowly burns, the flame flickers and goes out. Just as a candle, we need to be re-lit, too."

A Parent

---

**THE PARTNERS PLUS OUTREACH PROJECT...**
- works with communities to establish a Partners Plus respite program through replication of a proven model
- helps families learn how to find and train respite caregivers
- supports individuals in becoming caregivers

**WHAT IS RESPITE?**
Respite is temporary child care that relieves families from the daily responsibilities of caring for their children with special needs. Together families and caregivers decide when, where, and under what terms respite will take place.

**WHO CAN PARTICIPATE?**
- Communities interested in creating a Partners Plus respite program
- Families of children with special needs who might need or want respite
- Individuals interested in learning about providing care for children with special needs

---

**PARTNERS PLUS OUTREACH OFFERS...**
- Support and training to communities, families, and caregivers
- Resource manuals for communities, families, and caregivers
- A proven model for family-centered respite

The project helps communities develop a respite program using a train-the-trainer approach. The project helps families recruit caregivers and plan training specific to their children's needs. In training, families and caregivers:
- meet one another
- learn about caring for children with special needs
- discuss the challenges children with special needs experience
- build successful partnerships with each other

---

**CHILD DEVELOPMENT RESOURCES...**
A private nonprofit agency, has won national recognition for its services for young children and their families and technical assistance to the professionals who serve them. Partners Plus, initially serving Virginia and now offering its respite model to other states, is one of several programs offered by Child Development Resources.

---

FOR MORE INFORMATION
Please call Partners Plus staff or complete and return this form.
Phone: (757) 220-1168 Fax: (757) 253-1779 e-mail: cdrtraining@tez.net

---

**PARTNERS PLUS**
Families and Caregivers in Partnerships
Caring for Children with Special Needs

---

**CHILD DEVELOPMENT RESOURCES**

---

**CHILD DEVELOPMENT RESOURCES**

---

**BEST COPY AVAILABLE**

---

**Partners Plus Community Planning Manual**
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

---

**DRAFT 10.99**
CDR offers parents help with respite care

by Lisa L. Ownby

Where do parents turn when they need a break from the everyday demands of caring for their children? Most families can find a friend or neighborhood babysitter to get a night out but for families of children with disabilities and/or special health care needs it's not that easy. Finding a caregiver who is available, comfortable, and knowledgeable about caring for a child with special needs can be difficult.

These parents also need a break and can benefit from respite (temporary child care). Respite is described by one family in this way: "think of a candle that slowly burns, the flame flickers and goes out and just as a candle, we need to be re-lit, too."

PARTners Plus, a program of Child Development Resources (CDR) is designed to assist families of children (birth-8) with special needs in selecting and training their own respite caregivers. PARTners also offers training and support to individuals interested in serving as caregivers for families.

PARTners holds workshops regularly for both families and potential caregivers. At the workshops participants will learn about caring for children with special needs; meet other families and caregivers; discover some of the challenges children with special needs experience, and talk about how families and caregivers can build successful partnerships.

These workshops are free of charge and all participants receive a resource notebook. Staff are available after the workshop to help families train the caregiver they select. PARTners will pay caregivers up to 10 hours of time spent in training. For more information ask for PARTners Plus staff at (757) 566-3300.

After completing a PARTners workshop a parent said, "it's sometimes difficult to make the time to find a caregiver when you have a child with special needs... but having the support of CDR gives us peace of mind to enjoy time away. For our family, respite is a life saver!"

Lisa L. Ownby, M.S.W., is the project coordinator for PARTners Plus.
Sample Press Release and Feature Articles

How to use:

Adapt this to include your community’s information and submit to local media.

FOR IMMEDIATE RELEASE

CONTACT:
Name
Phone Number

Caring for Children with Special Needs

(City/Town) Date -- ___________’s Partners program is conducting a workshop on caring for children with special needs. Families of children with special needs will learn about respite (short-term care for their children) and how to find and train caregivers. Individuals interested in becoming caregivers will learn about working with children with special needs and their families.

The workshop will be held on ___________ from ___________ to ___________ at ___________.

Accommodations for participants with special needs, including sign language interpreters, will be made available upon request. Child care will be available for those who pre-register. For more information, or to register, please call ___________.

###

Partners Plus Community Planning Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99 4-7
The following is a sample Partners Plus flyer.

Partners Plus
Free Workshop

Caring for children with special needs -
Finding a babysitter is usually a routine chore for most families with young children. But parents of children with special needs often have a hard time finding caregivers who are knowledgeable, comfortable, and available to care for their child with special needs.

Partners Plus helps families learn how to find and train caregivers to provide respite (temporary child care). This project also offers support and training to individuals interested in helping families by serving as a caregiver for their children.

Special Accommodations
...can be made with sufficient time of request. Participants will receive a free notebook and child care is available for those who pre-register. Light refreshments will be provided, however, participants will need to bring a bag lunch.

When?
Saturday, November 20, 1999
9:00 a.m. until 3:00 p.m.

Where?
Child Development Resources
150 Point O' Woods Road
Norge, Virginia
757-566-3300

How do I sign up?
Call 220-1168 to register and receive more information.

AT THE WORKSHOP FAMILIES AND CAREGIVERS WILL:

- learn about caring for children with special needs
- meet other families and caregivers
- discover some of the challenges children with special needs experience
- talk about how families and caregivers can build successful partnerships

It's free!
# Implementing Partners: Planning for Workshop

<table>
<thead>
<tr>
<th>Activity</th>
<th>Person Responsible/Comments</th>
<th>Date Complete</th>
</tr>
</thead>
<tbody>
<tr>
<td>Promote community awareness</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Recruit families and caregivers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Schedule Partners group training</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• trainers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• date</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• location</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• time</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Develop training announcement and distribute</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Register training participants and create/maintain database</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plan for child care during training</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Register children</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Recruit caregivers</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Planning for Workshop continued

<table>
<thead>
<tr>
<th>Activity</th>
<th>Person Responsible/Comments</th>
<th>Date Complete</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prepare training materials for participants</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• manuals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• agenda</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• ice breaker</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• evaluation forms</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Call registered participants to remind them to attend training</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Purchase or provide refreshments</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prepare/set up rooms at training location</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Evaluate training</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Provide follow-up technical assistance such as:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Help families choose caregivers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Support families as they train their own caregivers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Provide continuing support</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Partners Workshop Agenda

Date

Location

Time

Part I

Time

Registration and Light Refreshments

Welcome and Introductions

About Partners

Communication

Time

Break - 5 minutes

Building Partnerships: Families and Caregivers Working Together

Preview of Coming Attractions

Time

See Ya for Part 2!
Partners Workshop Agenda

Date
Location
Time

Part II

Time  Registration and Light Refreshments

Time  Activity: All Kids Like Cookies

Time  Break Out Session for Families
   Finding Respite Caregivers
   Interviewing Caregivers
   Developing an Personalized Training Plan

Time  Break Out Session for Caregivers
   Understanding Child Development
   Disabilities Awareness
   Children with Special Needs
   Becoming a Respite Caregiver

Time  Break - 5 minutes
   Practice Activity: Using Daily Routines
   Wrap up and Evaluations

Time  Congratulations! You are on your way!
Checklist of Materials and Equipment for Partners Workshop

Date: ___________ Time: ___________ Location: _______________________

Purpose: ___________________________________________________________

Number Expected to Attend: _____ Presenter(s): _______________________

Room and Facility Preparation

___ Table for Registration
___ Table for Display Materials
___ Table Space for All Participants
___ Tables Arranged in a Horseshoe Shape
___ Chairs for All Participants
___ Extra Tables for "All Kids Like Cookies" Activity
___ Tables for Lunch or Refreshments (optional)
___ Overall Agenda for Training
___ Rooms for Childcare According to Numbers of Children Attending
___ Childcare Rooms with Accessibility to Lavatories (Preferably in Room)

Childcare Preparation

___ Caregiver SignIn Sheet
___ Appropriate Toys and Activities
___ Activity Supplies
___ Snacks
___ First Aid Kit
___ "Partners Plus Workshop Childcare Information"

Getting Started

___ Refreshment Supplies (optional)
___ Name Tags or Tents
___ Sign-In Sheets
___ Initial Family Survey

Getting Started, continued

___ Comfort Measure-Pre
___ Knowledge Measure-Pre
___ Participant’s Manuals
___ Overall Agenda for the Training
___ Getting Acquainted Activity
___ Pencils for Participants
___ Sample Press Release and Feature Articles form (optional)
___ Permission for Videotape with Sound / Photo Use (Enough for All Participants)

Section 1

___ Handouts
___ Prepared Overhead Transparencies / Flip Charts
___ Overhead Projector and Screen (optional)
___ Markers for Either Flip Charts or Overhead Transparencies
___ Flip Chart Stand
___ Incentives for Participation in Methods of Communication Activity

Section 2

___ Prepared Overhead Transparencies / Flip Charts
___ Overhead Projector and Screen (optional)
Workshop Materials, continued

Section 2, continued
- Markers for Either Flip Charts or Overhead Transparencies
- Flip Chart Stand
- Incentives for Participation in Methods of Communication Activity

Section 3
- Prepared Overhead Transparencies / Flip Charts
- Overhead Projector and Screen (optional)
- Markers for Either Flip Charts or Overhead Transparencies
- Flip Chart Stand

Section 4
- Handouts
- Enough of the Easy and Hard Version of the XYZ Tests (Equal Number of Each) for All Participants
- Method for Dividing Participants into Groups of Equal Number (for the "All Kids Like Cookies" Activity)
- Directions for Each Station
- Props for the "All Kids Like Cookies" Activity
  - Vision Station: several pairs of prepared sunglasses, blindfolds, a page out of a coloring book or a crossword puzzle, crayons, children’s books and magazines, plastic cup and pitcher with water, paper towels
  - Speech Station: marshmallows, gauze, trash can, paper towels
  - Fine Motor Station: several pairs of thick gloves, masking tape, puzzles, small manipulative toys, thick string and beads or pasta
  - Gross Motor Station: two large balls, two retrievable toys, masking tape, use masking tape to make two ten to fifteen foot walkways

Section 4, continued
- Prepared Overhead Transparencies / Flip Charts
- Overhead Projector and Screen (optional)
- Markers for Either Flip Charts or Overhead Transparencies
- Flip Chart Stand

Section 5-F
- Overhead Transparencies / Flip Charts
  - Developed by Trainers (optional)
- Overhead Projector (if needed)
- Flip Chart Stand (if needed)

Section 5-C
- Prepared Overhead Transparencies / Flip Charts
- Overhead Projector
- Flip Chart Stand

Section 6-F
- Overhead Transparencies / Flip Charts
  - Developed by Trainers (optional)
- Overhead Projector (if needed)
- Flip Chart Stand (if needed)

Section 6-C
- Prepared Overhead Transparencies / Flip Charts
- Overhead Projector
- Flip Chart Stand

Section 7-F
- Overhead Transparencies / Flip Charts
  - Developed by Trainers (optional)
- Overhead Projector (if needed)
- Flip Chart Stand (if needed)
Workshop Materials, continued

Section 7-C
____ Prepared Overhead Transparencies / Flip Charts
____ Overhead Projector
____ Flip Chart Stand

Section 8-F
____ Overhead Transparencies / Flip Charts
____ Developed by Trainers (optional)
____ Overhead Projector (if needed)
____ Flip Chart Stand (if needed)

Section 8-C
____ Prepared Overhead Transparencies / Flip Charts
____ Overhead Projector
____ Flip Chart Stand
____ Criminal History Release (optional)
____ Department of Social Services Check (optional)
____ Stamped Return Envelopes (optional)

Section 9-F
____ "Add My Name" Family Directory

Section 9-C
____ Overhead Transparencies / Flip Charts
____ Developed by Trainers (optional)
____ Overhead Projector (if needed)
____ Flip Chart Stand (if needed)

Section 10
____ Caregiver Profile Form for Each Participant

Final Activity
____ Prepared Overhead Transparencies / Flip Charts
____ Overhead Projector
____ Flip Chart Stand

Wrap Up
____ What Did You Think?
____ Comfort (Post)
____ Knowledge (Post)
____ Certificates

General Supplies to Bring
____ Masking Tape
____ Scissors
____ Markers for Name Tents
____ Play Doh (optional)
____ Sticky Pads (optional)
____ Other Table Supplies for Participants
____ Candy or Other Energizer (optional)
Partners Plus Workshop
Childcare Information

We will be caring for your child at the Partners Plus Workshop scheduled for: __________. Please help us know more about your child:

Child's Name: ___________________________ Date of Birth: __________

Routine at Home—Wake Time: __________ Meal Times: _________________
  Naptime: ___________________________ Bedtime: _________________

His / Her Special Abilities: ___________________________

_________________________________________________________________________________

_________________________________________________________________________________

Favorite Toys or Playthings: ___________________________

Favorite Foods / Any Allergies? _________________________

Favorite Games, Activities, Songs: _______________________

What Quiets Him / Her? _________________________________

What Scares Him / Her? _________________________________

What Excites Him / Her? _________________________________

Names and Ages of Brothers or Sisters, if any: _______________________

Familiar Names (Friends, Family, Pets): _______________________

_________________________________________________________________________________

Any Concerns About Your Child? _________________________

_________________________________________________________________________________

How Can We Help? ___________________________________

_________________________________________________________________________________
Permission for Videotape with Sound/Photograph Use

I _____ Do _____ Do not (check one) give permission for me and/or my child, to be filmed on videotape with sound to be used for public information and publicity about the Partners project.

I _____ Do _____ Do not (check one) give my permission for me and/or my child, to be photographed for informal pictures to be used for dissemination materials for the Partners project.

Permission is applicable from the date signed to: ____________________________

Signature: ____________________________

Relationship to child: ____________________________

Date: ____________________________
Permission to Release Caregiver Name

How to use this sample form:

Prior to listing a caregiver's name in any publication (e.g. the Caregiver Directory), they will need to give permission for the release of their personal information.

I _____ Do _____ Do not (check one) give my permission for Partners to share my name and phone number with families interested in finding respite care. I understand that Partners will perform a criminal record check before adding my name to the caregiver directory.

I would like the following information to be added to the Caregiver Directory:

Name: ____________________________________________
Address: ____________________________________________
Telephone: __________________________ Date: ____________
Signature: ____________________________________________
Permission to Include Information in Family Directory

How to use this sample form:

Prior to listing a family’s name in any publication (e.g. the Family Directory), they will need to give permission for the release of their personal information.

Families wishing to meet other families who have children with special needs may want to be included in the Partners Family Directory. The Directory can serve as a networking tool which allows families to meet other families and possibly exchange respite services with other interested families.

I would like the following information to be added to the Partners Family Directory:

Name: ______________________________________________________________
Address: ___________________________________________________________
            ___________________________________________________________
            ___________________________________________________________
Telephone: ________________________________
Child’s Age and Special Needs: _________________________________________
            ___________________________________________________________
Sibling’s Age(s): ___________________________________________________
Signature: ___________________________ Date: ___________________________
**Partners Plus Order Form for Printed Materials**

Name:  
Title, Company:  
Street Address:  
City, State, Zip:  
Telephone:  Fax:  

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Item Number/ Description</th>
<th>Unit Cost</th>
<th>Total</th>
</tr>
</thead>
</table>

Subtotal  
Shipping and Handling  
Total  

Method of Payment:
Check number:  P.O. number:  
Visa or Mastercard number ($25 Minimum Charge):  
Expires:  Signature:  

Shipping and Handling Charges:
Orders totalling $1.00 to $49.99  $7.00  Additional Fees required for international and priority delivery service
Orders totalling $50.00 to $99.00  $10.00
Orders totalling $100.00 or more  $12.00

Return Completed Form To:
Child Development Resources, 1490 Government Road, Williamsburg, VA 23185
Phone (757) 220-1168 · Fax (757) 253-1779 · Email: ppcdr@tni.net
Section 5
measuring your success
Measuring Your Success

The purpose of evaluating your Partners program is twofold: documentation of activities and measurement of program impact. Program evaluation is a continual process of monitoring the quality of services and accomplishments. Efficacy data can be compiled and presented to program staff, families, caregivers, funding sources, and other community groups to build support for your program.

Partners Plus developed an evaluation system to determine the effectiveness of the model by collecting some simple data on the numbers of families and caregivers participating and the quality of the six-hour workshop. In addition, measures were developed to determine the efficacy of the project in increasing respite options.

If your Partners program plans to collect evaluation data, then you must first decide what you want to measure and who the audience is for your evaluation results.

Your program may want to consider three central evaluation questions.

- What did we do? (e.g. Who did we serve? How many workshops were held etc.?)
- What were the outcomes? (i.e., Did respite options increase for families? Did caregiver knowledge and comfort increase?)
- How satisfied were the participants? (i.e. What did families and caregivers think about training and technical support?)
Once you have decided what you want to measure, you'll need to consider how that information will be collected. Consider using:

- written surveys
- telephone interviews

Your program may want to collect descriptive information about the families and caregivers you support i.e., the age of the child, family/caregiver educational background, etc.

Sample instruments are included in Section 6. You may choose to use and/or modify these evaluation instruments to meet the needs of your program.

- **What Did You Think About the Training?** This instrument is used at the end of the six-hour training to gather numerical data and narrative comments about the quality of the workshop content, process, and trainers.

- The **Initial Family Survey** is designed to measure a family's respite options prior to Partners training and also collects demographic information. This survey can be conducted over the telephone, through the mail, or at the beginning of the workshop. When given more than once, it can measure a change in respite options over a period of time.

- The **Follow-up Family Survey** is designed to determine if families respite options increased as a result of Partners training. It is completed via mail or telephone after participating in the Partners workshop.
Out-of-Home Respite

- The Caregiver Knowledge and Caregiver Comfort Scale Pre and Post evaluations are used to measure caregiver knowledge and comfort levels as a result of Partners workshops. These are completed before and after the six-hour workshop.

- The Follow-up Caregiver Survey is used to measure if the personalized training has increased the caregivers' comfort level. These are completed via mail or telephone after attending the Partners workshop.
Section 6

Sample evaluation instruments
Sample Evaluation Instruments

Sample Evaluation Instruments

- What Did You Think About the Training?
- Initial Family Survey
- Follow-up Family Survey
- Follow-up Caregiver Survey
- Caregiver Knowledge Pre/Post
- Caregiver Comfort Scale Pre/Post
What Did You Think About the Workshop?

Name: ___________________________ Date: ___________________________

Presenter(s): ___________________________

1. What I liked about the workshop...

_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________

2. How Did We Do? The information presented was:

___ easy to understand
___ difficult to understand because...
_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________

3. The information presented was:

___ useful
___ not useful because...
_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________
4. The overall quality of the workshop was:  
   1  2  3  4  5  
   Very Poor  Very Good

5. Were the workshop materials helpful?  
   1  2  3  4  5  
   Not at All  Very

6. Was the information presented appropriate for your needs?  
   1  2  3  4  5  
   Not at All  Very

7. Was there enough chance for questions and discussion?  
   1  2  3  4  5  
   Not at All  Very

8. I gained knowledge about...

   ___________________________________________________
   ___________________________________________________
   ___________________________________________________

9. Suggestions I would like to offer for improving the workshop...

   ___________________________________________________
   ___________________________________________________
   ___________________________________________________

10. I think what I'll remember most was...

    ___________________________________________________
    ___________________________________________________
    ___________________________________________________

11. Please share any additional comments:

    ___________________________________________________
    ___________________________________________________
    ___________________________________________________
Partners Plus Initial Family Survey

Please describe any special needs your child might have. If you have more than one child, you may use the space on the back.

Child's Name: ___________________________ Age: ______

1. Does your child have any special needs such as: developmental delay, multiple disabilities, vision, hearing, feeding, behavior, heart problems, breathing problems, other? If so, please explain.

__________________________________________________________________________

__________________________________________________________________________

__________________________________________________________________________

2. Has your child received early intervention, special education, or therapeutic services? If so, please explain.

__________________________________________________________________________

__________________________________________________________________________

__________________________________________________________________________

3. Does your child use any assistive technology such as: electric wheelchair, medical equipment, communications board, other? Please describe.

__________________________________________________________________________

__________________________________________________________________________

__________________________________________________________________________
Initial Family Survey continued

Answer the following questions if you have ever left your child in another person's care. Please circle the best answer.

4. In general, how hard has it been to arrange respite (temporary child care) for your child(ren)?

<table>
<thead>
<tr>
<th>Not Difficult</th>
<th>Somewhat Difficult</th>
<th>Very Difficult</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5</td>
</tr>
</tbody>
</table>

5. How comfortable did you feel leaving your child(ren) with the caregiver?

<table>
<thead>
<tr>
<th>Not Difficult</th>
<th>Somewhat Difficult</th>
<th>Very Difficult</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5</td>
</tr>
</tbody>
</table>

6. Have you ever taught a caregiver to care for your child?

<table>
<thead>
<tr>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
</table>

7. How much time did you spend teaching the person how to care for your child?

<table>
<thead>
<tr>
<th>Less than 30 mins.</th>
<th>About 3 hours</th>
<th>Over 3 hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1-3</td>
<td>4-6</td>
</tr>
<tr>
<td></td>
<td>6 and up</td>
<td></td>
</tr>
</tbody>
</table>

8. In the last three months, how often have you used respite care?

<table>
<thead>
<tr>
<th>0</th>
<th>1-3</th>
<th>4-6</th>
<th>6 and up</th>
</tr>
</thead>
</table>

9. In the last three months, how often did you want respite, but could not find a caregiver?

<table>
<thead>
<tr>
<th>0</th>
<th>1-3</th>
<th>4-6</th>
<th>6 and up</th>
</tr>
</thead>
</table>

10. In the last three months, how many times did caregivers cancel?

<table>
<thead>
<tr>
<th>0</th>
<th>1-3</th>
<th>4-6</th>
<th>6 and up</th>
</tr>
</thead>
</table>

Please answer the following questions about your family members. This information is optional.

11. Child's Mother's Age ____________ Child’s Father’s Age ____________

12. What is the child’s mother’s highest level of education?

<table>
<thead>
<tr>
<th>Middle School</th>
<th>College Graduate (4 years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>High School</td>
<td>Some Graduate School</td>
</tr>
<tr>
<td>Some College</td>
<td>Graduate Degree</td>
</tr>
</tbody>
</table>

13. What is the child’s father’s highest level of education?

<table>
<thead>
<tr>
<th>Middle School</th>
<th>College Graduate (4 years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>High School</td>
<td>Some Graduate School</td>
</tr>
<tr>
<td>Some College</td>
<td>Graduate Degree</td>
</tr>
</tbody>
</table>
Partners Plus Follow-up Family Survey

Name: ______________________  Date: ______________________

If you have participated in the Partners workshop, please answer the following questions:

1. How long ago did you participate in the Partners workshop? _____ Months _____ Years

2. How helpful was the Partners workshop for finding caregivers?  
   Not Helpful  Somewhat Helpful  Very Helpful  
   1  2  3  4  5

3. How helpful was the Partners workshop for teaching caregivers?  
   Not Helpful  Somewhat Helpful  Very Helpful  
   1  2  3  4  5

4. How comfortable do you now feel leaving your child with someone else (a caregiver)?  
   Uncomfortable  Somewhat Comfortable  Very Comfortable  
   1  2  3  4  5

5. Since the Partners workshop, have you found a caregiver?  
   Yes  No

6. In the last three months, how often have you used respite care?  
   0  1-3  4-6  6 and up

7. In the last three months, did you want respite, but could not find someone?  
   Yes  No

8. In the last three months, how many times did caregivers cancel?  
   0  1-3  4-6  6 and up
9. Are you having any of the following problems finding good care for your child(ren)?

- Caregiver would not care for your child(ren)
- Caregiver would not follow your instructions
- Caregiver had insufficient skills
- Caregiver would not handle emergencies
- Other (please specify):

10. How helpful was the Family Manual?  
Not Helpful  Somewhat Helpful  Very Helpful

1  2  3  4  5

Thank You!

Please return this survey to:

Partners Plus Community Planning Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Partners Plus Follow-up
Caregiver Survey

Caregiver Name: ____________________ Date: ____________________

Part 1. Answer the following questions if you participated in the Partners workshop.

1. How long ago did you participate in the Partners workshop? _____ Months _____ Years

2. How well did the Partners workshop prepare you to work with children with special needs and their families?
   Poorly Prepared Somewhat Prepared Very Prepared
   1 2 3 4 5

3. Have you been matched with a family needing respite for their child?
   Yes No
   Number: ____________

4. Did the family you worked with participate in the Partners workshop?
   Yes No
   Number: ____________

5. Did the family provide you with individual training specific to their child?
   Yes No
   Number: ____________

Part 2. Answer the following questions if you cared for a child with special needs since the Partners workshop. If not, please continue to Part 3.

1. How many times have you provided respite care for a child with special needs since the Partners workshop? _____ 1-3 Times _____ 4-6 Times _____ 7 or More

2. How many different children with special needs have you provided care for since the Partners workshop? _______________ number of children
Please describe the child(ren) you provide care for and his/her special needs.

<table>
<thead>
<tr>
<th>Child</th>
<th>Child</th>
<th>Child</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Child’s Age</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Services Received</strong>&lt;br&gt;such as:&lt;br&gt;- early intervention&lt;br&gt;- special education&lt;br&gt;- therapies</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Special Needs</strong>&lt;br&gt;such as:&lt;br&gt;- development delay&lt;br&gt;- multiple disabilities&lt;br&gt;- vision&lt;br&gt;- hearing&lt;br&gt;- feeding&lt;br&gt;- behavior&lt;br&gt;- heart&lt;br&gt;- breathing&lt;br&gt;- other</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Assistive Technology</strong>&lt;br&gt;such as:&lt;br&gt;- electric wheelchair&lt;br&gt;- medical equipment&lt;br&gt;- communication board&lt;br&gt;- other</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Follow-up Caregiver Survey continued

Part 3. Please circle the number that best describes how comfortable you feel in working with children with special needs.

<table>
<thead>
<tr>
<th>How comfortable are you:</th>
<th>Uncomfortable</th>
<th>Somewhat Uncomfortable</th>
<th>Very Comfortable</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Providing care for a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>2. Planning activities that a child with special needs can enjoy.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>3. Meeting the routine health care needs of a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>4. Feeding a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>5. Talking with families of children with disabilities about their child and their child's strengths and needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>6. Talking to other people about any concerns they may have about children with special needs in your care.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>7. Identifying resources and getting help you may need in caring for children with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

Thank You!

Partners Plus Community Planning Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Partners Plus Caregiver
Knowledge Pre Evaluation

Name: __________________________ Date: __________________________

This survey is designed to gather information about your experience and knowledge of caring for children with disabilities or special needs.

Read each statement and circle the single best answer.

1. Early intervention services are available to children with disabilities who are
   a. Birth to three years old
   b. In public schools
   c. Three to five years old
   d. I don’t know

2. Children who receive early intervention special education services must
   a. Get therapy
   b. Go to a classroom program
   c. Have an individual plan
   d. I don’t know

3. All children with special needs receive education through
   a. MCH
   b. CHIP
   c. IDEA
   d. I don’t know

4. Which of these diagnoses tells you that a child has a special health care need?
   a. Autism
   b. Cystic Fibrosis
   c. Mental retardation
   d. I don’t know

5. Providing good respite care for a child with a disability depends heavily on
   a. Desire/attitude of caregivers
   b. Educational level of the caregiver
   c. A child’s disability
   d. I don’t know

6. When caring for children with disabilities in their homes, it is necessary to
   a. Learn about their daily household routines
   b. Change things to suit your way of providing care
   c. Ignore the way the family trained you to take care of the child
   d. I don’t know
7. A key to a successful respite care program for families is
   a. Providing each family with many caregivers
   b. Allowing each family to train their own caregivers
   c. To start more respite agencies
   d. I don’t know

8. It is most important for respite caregivers to
   a. Understand child development
   b. Know the needs of the child or children in their care
   c. Have a degree or certification
   d. I don’t know

9. The term “cognitive development” is used to describe how a child
   a. Relates to others
   b. Sits and grasps toys
   c. Thinks and solves problems
   d. I don’t know

10. If caregivers have questions or concerns about a child’s special needs, they should
    a. Discuss their concerns with the child’s parent(s) or guardian
    b. Call the local special education program
    c. Not do anything because if they are wrong it would upset the child’s parents
    d. I don’t know

Thank You!
Partners Plus Caregiver Knowledge Post Evaluation

Name: ___________________________ Date: ___________________________

This survey is designed to gather information about your experience and knowledge of caring for children with disabilities or special needs.

Read each statement and circle the single best answer.

1. Early intervention services are available to children with disabilities who are
   a. Birth to three years old
   b. In public schools
   c. Three to five years old
   d. I don't know

2. Children who receive early intervention special education services must
   a. Get therapy
   b. Go to a classroom program
   c. Have an individual plan
   d. I don't know

3. All children with special needs receive education through
   a. MCH
   b. CHIP
   c. IDEA
   d. I don't know

4. Which of these diagnoses tells you that a child has a special health care need?
   a. Autism
   b. Cystic Fibrosis
   c. Mental retardation
   d. I don't know

5. Providing good respite care for a child with a disability depends heavily on
   a. Desire/attitude of caregivers
   b. Educational level of the caregiver
   c. A child's disability
   d. I don't know

6. When caring for children with disabilities in their homes, it is necessary to
   a. Learn about their daily household routines
   b. Change things to suit your way of providing care
   c. Ignore the way the family trained you to take care of the child
   d. I don't know
7. A key to a successful respite care program for families is
   a. Providing each family with many caregivers
   b. Allowing each family to train their own caregivers
   c. To start more respite agencies
   d. I don’t know

8. It is most important for respite caregivers to
   a. Understand child development
   b. Know the needs of the child or children in their care
   c. Have a degree or certification
   d. I don’t know

9. The term “cognitive development” is used to describe how a child
   a. Relates to others
   b. Sits and grasps toys
   c. Thinks and solves problems
   d. I don’t know

10. If caregivers have questions or concerns about a child’s special needs, they should
   a. Discuss their concerns with the child’s parent(s) or guardian
   b. Call the local special education program
   c. Not do anything because if they are wrong it would upset the child’s parents
   d. I don’t know

Thank You!
Partners Plus Caregiver Comfort Scale Pre Evaluation

Please circle the number that best describes how comfortable you feel in working with children with special needs.

<table>
<thead>
<tr>
<th>How comfortable are you:</th>
<th>Uncomfortable</th>
<th>Somewhat Uncomfortable</th>
<th>Very Comfortable</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Providing care for a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>2. Planning activities that a child with special needs can enjoy.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>3. Meeting the routine health care needs of a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>4. Feeding a child with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>5. Talking with families of children with disabilities about their child and their child's strengths and needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>6. Talking to other people about any concerns they may have about children with special needs in your care.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>7. Identifying resources and getting help you may need in caring for children with special needs.</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

Thank You!

Partners Plus Community Planning Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127
Partners Plus Caregiver Comfort Scale Post Evaluation

Caregiver's Name: ____________________ Date: ____________________

Please circle the number that best describes how comfortable you feel in working with children with special needs.

How comfortable are you:

Uncomfortable Somewhat Uncomfortable Very Comfortable

1. Providing care for a child with special needs. 1 2 3 4 5

2. Planning activities that a child with special needs can enjoy. 1 2 3 4 5

3. Meeting the routine health care needs of a child with special needs. 1 2 3 4 5

4. Feeding a child with special needs. 1 2 3 4 5

5. Talking with families of children with disabilities about their child and their child's strengths and needs. 1 2 3 4 5

6. Talking to other people about any concerns they may have about children with special needs in your care. 1 2 3 4 5

7. Identifying resources and getting help you may need in caring for children with special needs. 1 2 3 4 5

Thank You!
Section 7

resources
Resources for Community Programs

Resources for Creating a Community Respite Program

The following are helpful resources that may be used in conjunction with planning and developing your Partners program.

Respite care for children with developmental and/or physical disabilities: A parent's perspective. Chapel Hill, NC: Author.


References for Board and Committee Development

If your Partners program is governed or advised by a board or committee, the following resources may support those board and committee members in carrying out their work.

While aimed specifically at Big Brothers/Big Sisters, this two-volume notebook is adaptable to many other voluntary associations. It includes a leader's guide for persons who have responsibility for staff and board training relative to the work of the board.

Areas covered are leadership styles, committee structure, agenda setting, conducting meetings, taking and reporting minutes, the nominating process, duties of officers, and board development.


Andringa and Engstrom, who collectively have more than 70 years of experience as chief executives, board members, and facilitators in the training of more than 20,000 board members in 20 countries, take a closer look at the practical aspects of nonprofit governance, including formulating board structure and process, developing a strong partnership between board and staff, structuring effective board and committee meetings, and other critical issues.
Resources for Community Programs


Twelve experts in various aspects of boardsmanship share their expertise on the board and the life of the organization in three major sections: overview of the board, organizing the board, and tasks and essential concerns of the board.


Conrad and Glenn present clear answers, backed by graphics, to questions such as a definition of mission, policy formation, functions of boards and staff, roles of board members and the Chief Executive Officer, why people join boards, and evaluation of board members. Also presented is a basic management concept for voluntary organizations.


A monograph from the same series as “Developing Effective Boards, Councils and Commissions.” Detailed explanation of committee purpose and function and specific responsibilities of both the appointing body and committee members.

Persons who serve on voluntary governing or advisory boards will find this book important reading. It offers a clear management concept and system appropriate to the unique challenges of voluntary organizations together with much practical application.


A compendium of research studies on board make-up and fund-raising success, board-member expectations, board member participation, women and boardsmanship, board/staff roles in management, and the planning and implementation of programs.


This pamphlet was written to give you a better understanding of the effectiveness of nonprofit organizations and how you can strengthen the capacities of their leadership.
Resources for Community Programs


This booklet is designed to help you improve the quality of board meetings you attend either as a leader or a participant by offering guidelines for planning, participating, and leading.


Houle outlines in detail the responsibilities of the board, the chief executive, and staff, and shows how these three units can divide important tasks and work together effectively to help organizations fulfill their missions. He describes strategies for structuring the board and improving overall board operation, showing how to bring out the best in board members, maximize each person's contribution, and help the board identify and achieve important goals.


Based on extensive experience working with volunteer boards and serving on them, O'Connell has developed a practical guide on such topics as legal responsibilities; finding good board members; planning, budget and financial accountability; the role of the board president; board/staff relations; and fund-raising.

This guidebook was prepared to aid the director of a nonprofit corporation in performing his or her duties. It was written for anyone presently serving as such a director, as well as anyone contemplating such service.


A loose-leaf notebook format with over 400 pages of text, work forms, checklists, etc., covering the areas of assessment, boardmanship process, the board as managers of planning and evaluation, committees and meetings, the board and the community, and leadership and team building. (Permission to copy is given.)


This compendium of over 60 articles originally appeared in newsletters or newspaper columns. Swanson usually covers one topic per article. They are easy to read and to remember.
Local and National Resources

Local and National Resource—Boards, Councils, and Committees

The following are resources that may be helpful in creating and maintaining boards, councils, and committees. In addition, Better Boards programs are located in many communities and organized collaboratively by volunteer centers, United Ways, county or city cooperative extension offices, and community colleges. These groups offer workshops and conferences on boardmanship tailored to the needs of local organizations.

Center for Volunteer Development
Virginia Tech
Blacksburg, VA 24061-0150
(703) 231-7966

Department of Volunteerism
Commonwealth of Virginia
223 Governor Street
Richmond, VA 23219
(804) 786-1431

The National Center for Nonprofit Boards
1225 19th Street, N.W., Suite 340
Washington, DC 20036
(202) 452-6262
References on Adult Learning

The following resources may be helpful to trainers conducting Partners workshops. Specific training tips are included in the Workshop Facilitator's Manual contained in Families and Caregivers as Partners: A Guide to Family-Centered Respite Care.


References on Adult Learning


Partners Plus
A Family Manual
Families and Caregivers in Partnerships
Child Development Resources
DRAFT 10.99
Partners Plus

Families and Caregivers in Partnerships:

A Family Manual

By
Lisa L. Ownby, M.S.W.
Amanda C. Hooke, M.S.W.
Dee Wylie Moore, B.S.
Corinne W. Garland, M.Ed.

The Family Manual is part of a series of resources in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care. The rest of the series includes the Community Planning, Trainer's Workshop, and Caregiver Manuals, created by Child Development Resources.
Acknowledgments

The authors would like to thank Jayne Naughton for her technical assistance; Amy Watson for editorial assistance; families and caregivers who wrote the perspectives stories in section four of the Family and Caregiver Manuals; and our community advisory committee for multiple tasks throughout the development of training materials. Their suggestions and comments provided a significant contribution to the quality of this manual.

Amy Consiglio, of CM Consulting, designed and produced this manual.

A Word About Funding

Support for the Partners Plus project was provided under grant number HO24B40032 from the Early Education Program for Children with Disabilities, U.S. Department of Education. Points of view or opinions expressed herein do not, however, necessarily represent official views or opinions of the Department of Education.

Funding to revise the Partners Plus guide to help families use existing community programs as respite for their school-age children with mental retardation came from the Joseph P. Kennedy, Jr. Foundation. Revisions to the manuals include practical “how to” information for personnel working with children in before- and after-school programs, recreational, camp, and other community programs.

Copyright © 1999. All rights reserved. Trainers may reproduce this manual, without written permission, and in whole for use only with families and caregivers. For permission to reprint single pages or to purchase additional copies of the manual, please contact:

Child Development Resources
Post Office Box 280
Norge, VA  23127-0280
Phone (757) 566-3300 / Fax (757) 566-8977
Contents

Introduction
   The Partners Model: A New Model of Respite Care viii
   A Brief Partners History ix
   Partners Model: 5 Steps xi
   Partners Plus Replication Process xiv
   Partners Community Implementation Process xvi
   Partners Plus Manuals xvii

Section 1—About Respite
   Respite . . . Rest, Relax, Renew 1-2
   Families Who Need Respite 1-3
   Children Who Need Respite Care 1-4
   Benefits of Respite to Families 1-5
   Benefits of Respite to Children 1-6
   Benefits of Respite to Caregivers 1-6
   Why Don’t Families Use Respite Care? 1-7
   Keys to a Successful Respite Program 1-8
   Description of Family 1-9
   Description of Family-Centered Philosophy 1-9
   Family-Centered Care 1-10
   A New Respite Model: Partners 1-11
   Partners Model of Respite Care 1-12
Contents, continued

Section 2—Building Partnerships: Communication

Building Partnerships 2-2
Building Partnerships: Communication 2-2
How Do You Feel? 2-4
Six Messages of Communication 2-5
What You Need to Know About Communication 2-5
Methods of Communication 2-8
Sharing Information 2-9
Tips for Communicating with Your Caregiver 2-10

Section 3—Building Partnerships: Working Together

Building Partnerships: Working Together 3-2
"Welcome to Holland" 3-3
A Personal Story 3-5
Being a Father of a Child with Special Needs 3-7
A Parent’s Perspective 3-8
Malachi 3-10
A Sibling Perspective 3-13
One Caregiver’s Perspective 3-17
Stress 3-20
Coping with Stress 3-21
Strategies for Dealing with Stress 3-22
The Challenges of Respite 3-23
Meeting the Challenge 3-24
Successful Family and Caregiver Partnerships 3-25

Section 4—All Kids Like Cookies

All Kids Like Cookies Activity Sheet 4-2
Visual Impairment 4-3
Speech and Language Impairment 4-4
Physical Disability 4-5
Contents, continued

Section 5–Finding Respite Caregivers

Preparation 5-2
In-Home Respite Care 5-3
Out-of-Home Respite Care 5-4
Family Cooperative Respite Care 5-5
Finding Respite Caregivers: Circles of Family and Friends 5-5
Circles of Family and Friends 5-6
Other Ways to Find Respite Caregivers 5-8
Resources in Your Community 5-8
Choosing Potential Caregivers 5-9

Section 6–Interviewing Caregivers

Interviewing Potential Caregivers 6-2
Sample Interview Questions 6-5
Answering Caregivers Questions 6-7
What To Do After the Interview 6-8
Other Factors To Consider 6-9
Other Questions To Ask 6-10
Noticing Warning Signs 6-12
Visit the Caregiver’s Home 6-13
Visit the Community Program 6-14
Watch Your Child for Signs 6-15
Evaluating Your Respite Decision 6-16
## Contents, continued

### Section 7—Developing Your Personalized Training Plan

<table>
<thead>
<tr>
<th>Developing a Personalized Training Plan</th>
<th>7-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step by Step: A Guide to Developing Your Personalized Training Plan</td>
<td>7-3</td>
</tr>
<tr>
<td>Step 1: Identify training goals</td>
<td>7-4</td>
</tr>
<tr>
<td>Step 2: List training methods</td>
<td>7-5</td>
</tr>
<tr>
<td>Step 3: Use resources to help train your caregiver</td>
<td>7-8</td>
</tr>
<tr>
<td>Step 4: Evaluation</td>
<td>7-10</td>
</tr>
<tr>
<td>Sample Personalized Training Plan</td>
<td>7-12</td>
</tr>
<tr>
<td>Training Tips</td>
<td>7-13</td>
</tr>
</tbody>
</table>

### Section 8—Resources

<table>
<thead>
<tr>
<th>Universal Precautions Guidelines</th>
<th>8-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hand Washing</td>
<td>8-6</td>
</tr>
<tr>
<td>References on Adult Learning</td>
<td>8-9</td>
</tr>
<tr>
<td>Training Resources</td>
<td>8-10</td>
</tr>
<tr>
<td>ARCH: Respite Care</td>
<td></td>
</tr>
<tr>
<td>ARCH: A Parent's Perspective</td>
<td></td>
</tr>
<tr>
<td>ARCH: Caregivers Grieve, Too!</td>
<td></td>
</tr>
<tr>
<td>NICHCY: General Information About Disabilities</td>
<td></td>
</tr>
<tr>
<td>NICHCY: National Toll-Free Numbers</td>
<td></td>
</tr>
<tr>
<td>NICHCY: National Resources</td>
<td></td>
</tr>
</tbody>
</table>

### Section 9—Forms for Sharing Information

<table>
<thead>
<tr>
<th>Information Worksheets</th>
<th>9-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Essential Information</td>
<td>9-3</td>
</tr>
<tr>
<td>Health and Medical Information</td>
<td>9-5</td>
</tr>
<tr>
<td>Important Phone Numbers</td>
<td>9-7</td>
</tr>
<tr>
<td>About My Child</td>
<td>9-9</td>
</tr>
</tbody>
</table>
Contents, continued

Section 9—Forms for Sharing Information, continued

About Our Family 9-11
About Sisters and Brothers 9-13
Health and Safety Checklist 9-15
Step 1: Identify training goals 9-17
Step 2: List training methods 9-19
Step 3: Use resources to help train your caregiver 9-21
Step 4: Evaluation 9-23
Personalized Training Plan 9-25
Respite Caregiver Profile 9-27
Reference Check Form 9-31

Final Activity

Step 1: Identify training goals
Step 2: List training methods
Step 3: Use resources to help train your caregiver
Step 4: Evaluation
Family Manual

introduction
The Partners Model of Respite Care...

Helps families of children with special needs find and train their own respite caregivers.

Through training and support, families and caregivers build long-lasting, successful respite relationships.

Introduction

Like all families, families of children with special needs want and need time away from the demands of parenting. Respite—or temporary child care for children with disabilities or chronic illness—gives families time to enjoy an evening out by themselves, with other family members, or with friends. However, for these families, finding respite care is not as simple as calling a neighborhood babysitter. These families need caregivers who understand their children's special needs.

When the staff at Child Development Resources (CDR) spoke with families about what they wanted in a respite program, the message was clear. Families wanted to choose and train their own respite caregivers. They wanted affordable care that was as simple to schedule as calling the neighborhood babysitter. And, families wanted caregivers who were trained to provide care.
In response to families and with support from the Virginia Institute for Developmental Disabilities (VIDD), a University Affiliated Program at Virginia Commonwealth University in Richmond, Virginia, CDR developed a pilot project that was low-cost, family-directed respite care called PARTners: Parents As Respite Trainers.

CDR was awarded, in 1994, a three-year grant from the U.S. Department of Education, Office of Special Education Programs (OSEP), Early Education Program for Children with Disabilities (EEPCD), to build on the pilot and to develop a model of respite that could be replicated in other communities. The project, Partners Plus: Families and Caregivers in Partnerships, provides a model for families to use in finding and training respite caregivers. This model encourages families to use available, natural family and community supports to meet their respite needs.

Within this model, Partners broadly defines the terms respite and caregiver. For example, respite can take place in a family's home, a caregiver's home, or in before and after school, recreation, camp, or other children's program. A caregiver can be any individual who provides respite care to a child with special needs or includes a child with special needs within an existing community program.

Partners Plus provided training to more than 200 families and caregivers in a five-city, three-county area of eastern Virginia. Partners usually conducted the six hour Partners Plus workshop for families and caregivers in one day. Sometimes when it was more convenient for a group, Partners conducted the workshop in two parts on separate days. This flexibility ensured that Partners adapted training
A Brief Partners History

to fit the schedules of families and caregivers who are generally most receptive to training during nontraditional hours such as evenings and Saturdays. Families attending the workshop learned to provide their own caregivers with personalized training based on their children’s daily routines. Families either came to training with their own caregivers, met caregivers through the workshop, or through other project activities.

The Partners Project resulted in the development of four manuals designed to help communities, trainers, families, and caregivers use the Partners model. The Family Manual is one of four in a series of manuals that Partners has conveniently packaged together in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care. Besides the set of manuals contained in the guide, you can purchase the Family and Caregiver Manuals separately from the Partners Plus program through CDR.

Child Development Resources
Post Office Box 280
Norge, VA 23127-0280
Phone (757) 566-3300
Fax (757) 566-8977
Email ppcdr@tni.net
The Partners Model: A New Model of Respite Care

Partners programs provide group training and continuing support to families and potential caregivers. The Partners model contains five steps:

The Five Steps of the Partners Model

1. Recruit Families and Caregivers
2. Conduct Partners Workshop
3. Help Families Choose Caregivers
4. Guide Families as they Train their Own Caregivers
5. Provide Continuing Support

Step 1: Recruit Families

First, programs recruit families and potential caregivers to participate in Partners. Community awareness activities invite potential participants to a Partners workshop. Information about the workshop and the Partners program is distributed through parent groups, newspapers, radio, flyers, word of mouth, etc.
Partners Model: 5 Steps

Step 2: Conduct Partners Workshop

Next, programs conduct the Partners workshop. During the six-hour workshop, families learn how to find, screen, and interview caregivers for their children. In addition, they learn to structure training for their own caregivers based on the specific needs and daily routines of their individual children. Caregivers learn basic information about caring for children with special needs. During the workshop, caregivers are preparing to be active in their work with families, reflecting on their motivations for becoming respite caregivers and identifying their own learning needs for personalized training. Partners Family and Caregiver Manuals provide participants with useful resources to actively engage in during the workshop and to use as resources as they move through the next steps of the model.

Step 3: Families Choose Caregivers

After the workshop, Partners programs provide continuing support as families choose caregivers. Families frequently identify respite caregivers within their own circles of natural support. If families are unable to identify their own caregivers, Partners will assist by providing social and educational events that will help families and caregivers meet. During interviews, families and caregivers negotiate how respite occurs. Respite may be in-home or out-of-home and reimbursement for care can be flexible. Care may be provided for a fee, voluntarily, or in exchange for another service, such as yard work.
Step 4: Families Train Caregivers

Once families and caregivers decide to work together, *families train their own caregivers*. This “personalized training,” discussed during the workshop, is organized by families with help from Partners staff if needed. Families decide what routines caregivers need to learn and work with caregivers to determine how training will occur, thus forming true partnerships necessary for successful respite.

Step 5: Provide Continuing Support

As families and caregivers work together to develop a personalized respite training, the community Partners program *offers continuing support*, information and guidance. The Partners program is available to support families and caregivers in various ways as they begin their respite relationship. Partners staff members respond to requests for help and support from both families and caregivers. For example, Partners may help families and caregivers with screening, interviewing, developing personalized training plans, and providing educational opportunities.
Partners Replication Process

The Partners Plus Community Planning Manual, contained in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care, helps communities use the Partners model of respite care. Each community interested in model replication develops a plan for collaboration among agencies, organizations, and individuals in order to implement the model. Community resources and supports are needed to sustain the model, to create a home for the program, and to ensure continuation.

The community replication process includes four major steps:

1. Identifying the site for replication
2. Planning for replication
3. Implementing the model
4. Evaluating the model


Your community will be a good candidate for replication if you have the following:

- support from agencies that provide, or are involved with, respite services
knowledge of current respite funding to support families without financial resources

support and commitment from local early intervention programs, mental health or mental retardation programs, and/or family support programs

Each community needs to develop a plan for replication. Awareness activities aimed at a broad array of agencies, organizations, and individuals help strengthen community support. The creation of an advisory committee brings expertise and resources that will ensure a quality program. The replication plan includes finding a home for the program and securing staff responsible for implementing the program. Formal or informal interagency agreements and perhaps grant proposals may be necessary for community replication.

The Community Planning Manual provides strategies and materials to help communities develop and implement the respite care program. The Trainer's Workshop Manual, accompanying the Community Planning Manual, includes agendas, overheads, and suggestions for what to say during the Partners workshop. There are also suggestions for the kinds of resources that are needed by families and caregivers in order to foster and maintain long-term relationships.

Communities that replicate the Partners model gather information from families and caregivers to evaluate the quality and usefulness of training and support.
Partners in your Community

Partners Community Implementation Process

Identify Community Site

Plan for Replication of the Partners Model
- Create Community Support
- Determine Program Advisors
- Determine Administrative Structure
- Secure Financial Resources

Implement the Partners Model
- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families As They Train Their Own Caregivers
- Provide Continuing Support

Evaluate the Partners Program
There are four manuals in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care that support the Partners Plus replication process:

1. The Family Manual
2. The Caregiver Manual
3. The Community Planning Manual, and

The Family and Caregiver Manuals are used in the Partners workshop for families and caregivers. These manuals include resources that families and caregivers can continue to use as they develop partnerships for respite care for children with special needs. The Community Planning Manual helps communities develop a community-based, family-centered respite system. The Trainer's Workshop Manual contains overheads, forms, and "helpful hints" for conducting Partners workshops.

The Family Manual is easy to follow and guides families in choosing and training their own respite caregivers. Families use their children's daily routines to develop training plans for caregivers. The manual includes information about respite, communication, building partnerships, selecting caregivers, and developing personalized training. There are worksheets, resources, and forms that can be duplicated for future use.
Partners Plus Manuals

The Caregiver Manual provides caregivers with basic information on caring for young children with disabilities and supporting families. This manual includes information about respite, communication, building partnerships, disabilities awareness, child development, children with special needs, and personalized training with families.

The Community Planning Manual provides communities with a step-by-step guide to replication of the Partners model. By following each step, communities and families can work together to develop respite care programs. The manual contains forms and materials to assist with replication of the model.

The Trainer’s Workshop Manual, used in conjunction with the Community Planning Manual, helps community trainers to conduct the six-hour Partners workshop for families and caregivers.
Section 1

About respite
About Respite Care

Respite . . . Rest, Relax, Renew

Often we are asked, What is respite care? Our first response is simple: respite care is temporary care for a family member with special needs.

Families have told us in many different ways what respite means to them:

- Time to readjust.
- A break from the routine.
- Rest in spite of demanding circumstances.
- I feel refreshed and renewed when I come back from respite.
- Think of a candle that slowly burns: the flame flickers and goes out, and, just as a candle, we need to be relit, too.

— Partners Plus Families

More About Respite Care

Respite care is a term used to describe occasional care for an individual with special needs. Respite care programs began in the 1960s when people with disabilities began receiving care in their homes and communities instead of in institutions. Acknowledging the importance of respite care to families providing home care, Congress passed federal legislation providing funds to help communities develop respite care programs.
The 1986 federal legislation that provided funds to set up model programs of respite care defines respite care as in-home or out-of-home temporary, nonmedical care for families who have children (young and/or adults) with disabilities or chronic or terminal illnesses. More simply, respite care is any type of temporary care provided to a person with a disability or special health care need.

There are a variety of respite models, each designed to meet the unique needs of families. Descriptions of these models are included in Section 9 of this manual (see ARCH Fact Sheet Number 2).

**Families Who Need Respite**

Families who have children with special needs are very much like other families. Like all families, they have values, beliefs, interests, concerns, needs, and dreams. However, families of children with special needs also have special challenges that affect the way they live. Here are some things we have learned from families.

- Like other parents, parents of children with special needs are concerned about their children's development, health, education, training, and social life.
- Parents of children with special needs must put forth extra effort to help their children.
Who Needs Respite?

- Parents of children with special needs are often very knowledgeable about their children's disabilities and have learned some of the best ways to help them.
- Like other parents, parents of children with special needs are concerned about the needs of all of their family members, the education of all of their children, the activities of all of their family members, and their jobs.
- Daily routines for families of children with special needs are similar to those of other families.
- Families of children with special needs have hopes and dreams that are like those of most other families.
- Children with special needs and their brothers and sisters can grow up healthy and lead productive lives.

Children Who Need Respite Care

A child who needs a respite caregiver is one who has special physical, emotional, or health needs that require care beyond that necessary for other children. The kinds of special needs or disabilities that a child may have can vary greatly from hearing impairments that require a child to communicate through sign language to mental retardation to terminal illness.
There is no one approach to caring for children with special needs but there are some basic guidelines:

- Children with special needs are more like other children than they are different.
- Children with special needs can and do learn.
- Children with special needs should be encouraged to do as much for themselves as they can.
- Children with special needs thrive when given love, acceptance, and necessary support.
- Like other children, children with special needs benefit from healthy play and recreation, friendships, being outdoors, and going places.

Benefits of Respite to Families

Respite care gives parents a break from the daily demands of parenting a child with special needs, and offers much more.

- Rest to recuperate, rejuvenate, replenish, rejoice
- Enjoying movies, dinner, fishing
- Shopping for groceries, school, self, kids, family
- Pampering at a hair salon or gymnasium or having lunch with a friend
- Interests to pursue such as hobbies, sports, community activities, and education
- Time to spend with siblings, other family members, and friends
- Everything or anything families want to do!
Benefits of Respite Care

Benefits of Respite to Children

Respite is also beneficial to children. Some of these benefits include the following:

- Relief that their parents finally got out
- Exploring new faces and activities
- Special friends of their own
- Parents who feel rejuvenated
- Interacting with other adults and children
- Time to be independent from their family
- Educating each other about differences and similarities

Benefits of Respite to Caregivers

Respite also benefits caregivers in many ways.

- Really great children and families to get to know
- Especially rewarding to know that they are making a positive difference for a family
- Self-improvement and growth through learning new skills and ideas
- Positive attitudes about themselves and about people who are different from themselves
- Involvement with children, families, and communities
- Thoroughly appreciated by families who need them
- Everything caregivers want respite to be!
Who Can Be Respite Caregivers?

- Neighbors
- Friends
- Relatives
- Community Program Staff
- Church Members
- Students
- Bus Aides
- Anyone with a desire

Why Don’t Families Use Respite Care?

Despite the benefits of respite for everyone, many families do not use respite care. Families have identified various reasons:

- It is difficult to find someone to provide temporary care.
- We are afraid to leave our children with strangers.
- We fear that respite caregivers lack adequate training.
- We sometimes feel guilty about leaving our children.
- It is difficult to find community programs for respite care.
- It is too disappointing when caregivers cancel frequently.
Keys to a Successful Respite Program

Families say a successful respite program must have these elements:

- Helps families feel comfortable
- Gives families options for how they want to participate
- Gives families options for times to be involved
- Is family centered
- Supports families in developing new linkages
- Supports families in developing individualized caregiver training
- Asks families if they want to be involved in program planning
- Gives families access to dependable, reliable caregivers

Source:
Reprinted with permission of ARCH National Respite Network and Resource Center, 800 Eastowne Drive, Suite 105, Chapel Hill, NC 27514:
Description of Family

What Do We Mean by Family? We all come from families. Families are big, small, extended, nuclear, multigenerational, with one parent, two parents, and grandparents. We live under one roof or many. A family can be as temporary as a few weeks, or as permanent as forever. We become part of a family by birth, adoption, marriage, or from desire for mutual support. As family members, we nurture, protect, and influence each other. Families have strengths that flow from individual members and from the family as a whole. Each family is a culture unto itself, with unique values and its own way of realizing dreams. Together, our families are the source of our rich cultural heritage and spiritual diversity. Our families create neighborhoods, communities, states, and nations.

Source:

Description of Family-Centered Philosophy

Family centered means the recognition that the family is the constant in a child’s life and that service systems and the personnel must support, respect, encourage, and enhance the strengths and competence of the family.

Source:
Family-Centered Care

What does providing family-centered care mean?

- Recognizing that a family will always be in a child's life while services and professionals will change
- Making it easier for families, volunteers, and professionals to work together in hospitals, homes, and community care
- Sharing information, ideas, and concerns among families, volunteers, and professionals as a team
- Honoring the diversity of families, knowing that we each bring richness to the world we live in through our cultures and the ways we live our lives
- Recognizing and respecting that families meet their own needs in different ways
- Designing services that are flexible, that are easy for families to use, and that meet each family's individual needs
- Appreciating families as families and children as children and recognizing that they have a wide range of strengths, concerns, emotions, and dreams beyond their needs for specialized health and developmental services and support

Source:
Reprinted with permission of the Association for the Care of Children's Health, 19 Mantua Road, Mt. Royal, NJ 08061:
A New Respite Model: Partners

The Partners model of respite care is designed to increase respite options for families of children with special needs by helping families find and train their own respite caregivers.

Respite can be provided in:
- Families' homes
- Caregivers' homes
- Community respite programs
- Before and after school programs
- Recreation programs
- Camps

Respite can take place anytime, any place, with anyone in order to support families in meeting their needs.

Partners is a natural and family-centered model that involves families in the design, implementation, and evaluation of respite care. The model encourages the development of community collaboration among agencies and parent, religious, and/or support groups to support this model and to create a home for the program.

Together, families and caregivers attend group training. Families learn how to recruit, screen, interview, and train caregivers. Caregivers are provided with basic education in caring for children with special needs. After group training, the next step is for families and caregivers to match up and begin individualized training. The Family and Caregiver Manuals are provided to families and caregivers to use as a resource to help them through this process.
Partners Respite Model

The Partners program provides support to families and caregivers to help them begin building respite relationships. When a family cannot find a respite caregiver within their own circle of natural supports, they can choose from the pool of caregivers recruited by Partners Plus or can attend social and educational events coordinated by the program to help families and caregivers meet and match up.

With the Partners model, families and caregivers have choices about how respite care is provided. Respite may be provided in-home or out-of-home. Care may be provided for a fee, in exchange for another service, or by a volunteer. These decisions are made between families and caregivers. Families may also choose to develop a parent (or family) cooperative in which they care for each other’s children. Families decide what kind of training caregivers need and work with caregivers to determine how training will proceed, forming a true partnership necessary for successful respite.

Partners Model of Respite Care

**Step 1** Recruit families and caregivers

**Step 2** Conduct Partners workshop

**Step 3** Support families in choosing caregivers

**Step 4** Guide families as they train their own caregivers

**Step 5** Provide continuing support
Section 2

Building partnerships: Communication
Building Partnerships

*Respite care is not just about giving out medicine. It is a whole dialogue and relationship with a family.*

— Partners Plus Caregiver

Partners Plus supports strong relationships between families who have children with special needs and their caregivers. In partnerships, families and caregivers respect one another and commit to working together.

Partnerships actively prevent caregiver burnout and family disappointment. Partnerships are built with open and honest communication and understanding. Sections 2 and 3 of this manual explore building partnerships through communication and collaboration.

Building Partnerships Through Communication

Good communication strengthens the relationship between family and caregiver.

*Honesty and openness in communication help build cooperative relationships over time!*

What does communication mean?

- Sharing information, ideas, and feelings that increase understanding about yourself and others
- Listening carefully to each other in ways that develop partnerships and promote the best care for children
- Sharing common values and traditions and talking about similar experiences to strengthen your relationship with each other
- Appreciating differences to strengthen mutual respect
Building Partnerships: Communication

Clear communication is one key ingredient in building partnerships. We send messages all the time, even when we are not speaking.

Communication is . . . 7% Words

Research shows the words we use or choose are important in gaining understanding. In communicating with another person, we need to use language that is common to both the speaker and the listener.

38% Voice

The voice we hear is more important than the words we use. We need to be aware of our expression and tone of voice. We use our voices to help express things like sincerity, conviction, requests, and concern.

55% Facial and Body Language

Our face and body are the most powerful means of communication. Smiles, frowns, raised eyebrows, widened eyes, tears, laughter, scrunched-up faces, and wrinkled noses all tell us very different things.

Source:
Facial expressions can communicate a lot about our thoughts and feelings. Use the blank circles to draw faces that send messages to others.

Anxious  Angry  Bashful  Bored
Confident  Thinking  Confused  Scared
Innocent  Relieved

Adapted from feelings posters by: How Do You Feel Today? Productions, Box 1085, Agoura Hills, CA 91376. 818-706-2288. Email: p&p@howdoyoufeeltoday.com. Web site: www.howdoyoufeeltoday.com. International Copyright MCMLXXIX A.K. Graca © All rights reserved.
Six Messages of Communication

Communication is a cycle. As you communicate with others, think about the many places that communication can break down. Does the other person really understand what you mean? Each communication contains the possibility of six separate messages:

- What you mean to say
- What you actually say
- What the other person hears
- What the other person thinks he hears
- What the other person says
- What you think the other person says

Adapted from:

What You Need to Know About Communication

- Communication is the way we send and receive messages.

Most people use a combination of verbal and nonverbal communication. Being aware of and sensitive to the different ways people communicate helps to build mutual respect and understanding between families and caregivers.
Learning To Communicate

- Our ways of communicating are influenced by our culture.

  Our culture determines much about how we think, feel, act, and communicate with others.

- The term life ways refers to the many ways we are influenced by others and by our environment throughout our lives.

- Gestures, silence, body movements, distance, and emotional expressiveness are other ways of communicating that are influenced by our cultures and life ways.

For Example:
- Too much silence makes some people uncomfortable.
- Too few pauses while talking may disturb others.
- To be comfortable, some people need some distance between themselves and others.
- Lots of eye contact is important in establishing trust for some.
- For others, too much eye contact is considered disrespectful.
- A relaxed open body posture, with arms not crossed in front, invites others to talk.
- Leaning slightly forward shows interest.
- Sitting up straight helps you look more alert.
- Much of our communication is done without words.
- Our eyes are used to communicate.
For Example:
A look can be exchanged between parents that says, *It's time to leave*, or a look from a mother's eyes to her child can communicate the message, *You need to behave*.

- When people have similar experiences, we sometimes presume a shared understanding of feelings or thoughts.

For Example:
When a parent at an oncology clinic says to another parent in the waiting area, *My child is receiving chemotherapy for leukemia too*, an exchange that takes place by looking at each other in the eyes can send a message of shared understanding, even though each family's experience is unique.

Depending on their life ways, families use more spoken or written words to communicate. Words help us to be direct, exact, and logical and encourage others to say more. The following is an example of a family and caregiver interview in which words are used to encourage a response.

Caregiver: I want to provide respite because I enjoy being with children.

Family: That's great! You like being with children.
Methods of Communication

Caregiver: Yes, I miss being a parent and my grandchildren live in another state. When my children were young, I enjoyed planning activities for them.

Family: What type of activities do you like to plan for children?

- Spoken and written words are helpful in finding out about a person's feelings and attitudes.

Questions that cannot be answered with a yes or no encourage others to talk about what is on their mind. Use words like how, why, what, and tell me.

For Example:

Family: On your first visit to my home you watched me feed Crystal and I gave you some information to take home to read. Why don't you try feeding Crystal today?

Caregiver: I think I should watch again to make sure I remember how.

Family: You seem uncomfortable trying this now. Tell me about your concerns.

Caregiver: I remember how much Crystal coughed when you fed her and how much she gagged. I am afraid I will choke her. I think you need someone with nursing skills to take care of your child.
Family: Perhaps we need to talk again about why Crystal has difficulty eating and why she might cough or gag. I want you to understand Crystal's special needs and to feel comfortable in caring for her. I am here to help while you feed her. Let me show you again. Then you try.

- Using words like describe, show me, and tell me can help determine if a caregiver has learned a skill or has understood the instructions.

For Example:

Family: How did your first day go?

Caregiver: I had trouble using Jared’s wheelchair when we went outside.

Family: Tell me what gave you trouble.

Sharing Information

To ensure a successful respite relationship, families and caregivers need to openly share information. This is necessary in initial training and throughout the relationship.

Information can be shared in many ways:

- Keep and share a respite journal.
- Plan information you want to share when you get together.
Tips for Caregiver Interaction

- Allow time to meet with each other before respite care takes place.
- Allow time to talk after respite care.
- Make necessary decisions promptly.
- Resolve conflicts in a timely way.
- Solve problems together.

Tips for Communicating with Your Caregiver

- Communication is essential and is an important part of maintaining a successful relationship.
- Give clear instructions and make sure your caregiver understands what you have said.
- Provide immediate feedback to your caregiver during all training activities.
- Put yourself in the caregiver’s place; remember, you once had to learn new ways to care for your child.
- Let your caregiver know when respite care goes well and when you are pleased.
- Be positive. When changes are necessary, tell your caregiver in a positive way.
Section 3

building partnerships:
working together
Partnerships: Working Together

Anyone with a desire to help families of children with special needs can become a capable respite caregiver. At first, many parents do not know how to care for their children with special needs, but gradually they learn. Through willingness to learn from families, respite caregivers can do the same.

Many respite caregivers, although not parents of children with special needs, can appreciate the benefits of getting a break from caring for children. Some people have life experiences that help them become more comfortable as respite caregivers. They may know someone who has special needs—a friend, a student, a classmate, someone at church, or a family member. The most important quality a respite caregiver has is a positive attitude.

We know children and their families depend on each other. Children depend on their families for love, support, and guidance. The family is a continuing influence in their child’s life, whereas friends, neighbors, and caregivers change. Parents know their own children best. Families and caregivers need to know how to work together for successful respite care.

While caring for children with special needs can be a wonderful experience for both families and caregivers, it can also be stressful. It is helpful for families to understand the challenges that caregivers face when caring for children with special needs. Caregivers may feel unimportant or underappreciated. They may be uncomfortable with the tasks they are given. The material on the following pages can help you learn from each other.
Family and Caregiver Perspectives

Families and caregivers can better understand each other by listening to personal experiences. Several perspectives from families and caregivers are included in this section.

The first story is “Welcome to Holland” by Emily Pearl Kingsley. From that story, you will see how one parent, whose child was born with a disability, saw her life changing. Other stories from families and caregivers are also included. Your story may look very different from these. To get a better understanding of one another we encourage families and caregivers in partnerships to share their unique stories.

“Welcome to Holland”

I am often asked to describe the experience of raising a child with a disability—to try to help people who have not shared that unique experience to understand it, to imagine how it would feel. It’s like this . . .

When you’re going to have a baby, it’s like planning a fabulous vacation trip—to Italy. You buy a bunch of guidebooks and make your wonderful plans—the Coliseum, the Michelangelo David, the gondolas in Venice. You may learn some handy phrases in Italian. It’s all exciting.

After months of eager anticipation, the day finally arrives. You pack your bags and off you go. Several hours later the plane lands. The stewardess comes in and says, “Welcome to Holland!”
"Holland?!" you say. "What do you mean, Holland? I signed up for Italy! I'm supposed to be in Italy. All my life I've dreamed of going to Italy!"

But there's been a change in the flight plan. They've landed in Holland and there you must stay.

The important thing is that they haven't taken you to a horrible, disgusting, filthy place, full of pestilence, famine, and disease. It's just a different place.

So you must go out and buy new guidebooks. And you must learn a whole new language. And you will meet a whole new group of people that you would have never met.

It's just a different place. It's slower paced than Italy, less flashy than Italy. But after you've been there for awhile you look around and you begin to notice Holland has windmills, Holland has tulips, Holland even has Rembrandts.

But everyone you know is busy coming and going from Italy, and they're bragging about what a wonderful time they had there. And for the rest of your life, you will say, "Yes, that's where I was supposed to go. That's what I had planned." The pain of that will never, ever, go away because the loss of that dream is a very significant loss.

But if you spend your life mourning the fact that you didn't go to Italy, you may never be free to enjoy the very special, the very lovely things in Holland.

— Emily Perl Kingsley, 1981, Parent
A Personal Story

Although my son Matthew died, he is still a part of my life. Matthew made a difference in who I have become. At first I shared the same thoughts and feelings that Ms. Kingsley expresses in "Welcome to Holland." After Matthew was born, my family began to live a different kind of life. It was often very difficult for us. We learned a lot about ourselves and we learned a lot from Matthew. We discovered that our love for Matthew and each other was very powerful. We found that we had many hidden strengths. Our family grew stronger ties with our community. Friends, neighbors, professionals, and even strangers helped us in many different ways. There were two very important things we wanted most from others. One was for others to accept our son. The other was respect for our family’s decisions in how to live with our child’s chronic illness and disabilities.

After getting to know Matthew, his disabilities did not seem like the most important part of his life. My four-year old daughter taught me to love Matthew just because he was a baby. To my daughter, Matthew was always her baby brother, not a baby with disabilities. She never noticed much difference until she got older. Other families of children with disabilities have shared some similar thoughts with me. Many say they wish others would stop seeing only their child’s wheelchair, their tubes, the way they talk differently or look different. At times, families want professionals to stop testing their
child long enough to get to know him or her as a child, a child with a name and family.

When I describe my son, I tell people that Matthew was small and easy to carry places. Daily care for Matthew was like taking care of an infant. He had a crooked little smile. His sister could make him smile most often, but others could sometimes if they gave him a gentle squeeze around his shoulders. His favorite game was to repeat sounds that we made to him. Matthew's favorite toy was a baby's beaded teething ring. He liked to fall asleep on his father's shoulder. Matthew's greatest strength was that he could teach others about the power of the human spirit.

In describing Matthew's disabilities I would say that Matthew had a genetic impairment called trisomy 22. Trisomy 22 is an extra chromosome that affects every part of the body. He never grew to be more than five pounds and twenty-three inches. There was a hole in the roof of Matthew's mouth that made sucking and feeding very difficult. We fed Matthew with a tube that we put down his throat and into his stomach. His heart and all other systems in his body did not work well. Matthew was delayed in all areas of his development. With help from professionals we learned to deal with all of his special needs.

Getting information about our son's disability through an early intervention program and from medical practitioners helped us understand how to care for our child. A service that helped our family a
"Father of Child with Special Needs"

lot was respite care. My family and I needed time separate from being with Matthew. I wanted to give all my attention to my daughter at her soccer games. My husband and I needed time together to laugh and have fun. I needed time to rest physically from the frequent round the clock feedings required for Matthew. I needed emotional rest from the constant concerns I had for Matthew. A few friends, neighbors, and community agencies helped us get respite from time to time. One agency linked us with a community family who was willing to give me a few hours of respite each week. Just those few hours gave me the relief I needed to renew my energy and continue to care for Matthew.

— Dee Moore, 1995, Parent

Being a Father of a Child with Special Needs

Being a father of a special needs child takes a special person. There are times when you just want to give up but you know that you can't. The doctors gave my son 3 days to live and now my son is 3 years old. It's a blessing from God that he's still with me today. One thing that I try to remember is to treat him like a normal child. I take him to amusement parks on vacation and to the mall, just to name a few places.

There are times when it's very difficult being a father of a special needs child. I have to watch my
child go through intensive surgeries just to be able to eat and breathe like you or I. There are times when all you can do is just listen to your child cry uncontrollably, but overall being the father of a special needs child is a blessing from God. If you're a dad and wonder what it's like to be a dad to a special needs child, it is like taking care of three children.

— Lamar Ramsey, 1997, Parent

A Parent's Perspective

From the moment Kieren was born, I knew something wasn't quite right. He used to cry all the time. He was totally fearless, and I could leave him with anyone. He didn't seem to care if I was there or not. As he grew older, he used to run away all the time. We had to lock all the doors and never let go of his hand when we were out or he was gone and he didn't come back. If he saw something he liked, he would jump up and down, flap his hands, and shriek. Looking back, there were a lot of warning signs but I didn't know what I was looking for. I just knew that something was not right. I think one of the worst things about not knowing he had high-functioning autism was that no one else seemed to think there was a problem (or so they led me to believe). I would say to family and friends that I was concerned, and they would tell me, "Oh, all children do that," and dismiss it, and in the end I was beginning to doubt my sanity.
When he was 2, we were stationed in Germany, where 9 months later my daughter was born. I think it was a blessing in disguise that we were in that country as they test routinely for cystic fibrosis in all newborns. We found out when she was 6 weeks old that she had it. She was having some problems but the doctors told me this is normal, all babies do these things. Where have I heard that before?!

So we were sent back to the states so that we could get the health care she needed. Once she was relatively stable, I turned my attention back to Kieren. I made an appointment with the school division to get Kieren assessed to see if he was eligible for the special education program. During all the appointments I remember thinking he is doing really well today and they are going to think I'm crazy. As it turned out, he was eligible with significant delays in language, cognitive and fine motor skills.

He started the special education program, and things went relatively well for the first 6 months as Kieren's language started to emerge and he became a little more in control of his actions. I will never forget the first home visit I had with the special education teacher. She told me how Kieren didn't do this or that, wouldn't pay attention, kept flapping his hands, and wouldn't sit down and eat (tell me something I don't know!). During that visit I could hardly talk for fear of crying. I felt she was telling me what a lousy job I had done in bringing up my son. I blamed myself for a long time, thinking I should
have done something and he would be normal. I thought that the "experts" knew what they were talking about, and I tended to listen more to them than to my inner feelings.

I am glad to say that now I know better. I listen to myself first, and then incorporate what the "experts" say into my plan. I've learned the hard way that parents are the experts on their child, and what's good for one child is not necessarily so for all children.

— Itala Eller, 1997, Parent

Malachi

I have a special sibling. My brother is intelligent and trusting and 2 1/2 years older than me. When we were little, we used to have bitter sock wars on laundry day, and he taught me all about dinosaurs (he knew a lot about dinosaurs). He drew these amazing, elaborate, creative pictures. I remember one called "War of the Grapes." It was a crayon picture of hundreds of little grapes in all kinds of armor and camouflage, and the red grapes were battling the green grapes. You looked at this picture and were amazed at the 7-year-old mind that constructed it.

Malachi also has a disability. For many years, my parents have been trying to figure out what disability it is. His symptoms have grown progressively worse since he was young, although,
at times, some medications have given us hope for improvement. Malachi is painfully shy. While he learned to make eye contact, you wish he would stop, it seems so painful for him. To ask him a question, you need to get his attention, then repeat the question until he understands, and wait patiently for his answer. Malachi never asks the follow-up questions to statements, and he rarely involves himself in a conversation involving more than one person. He has trouble with directions. While he nods and indicates that he understands, he may not recall the conversation when it is time to carry the directions out. Malachi was slow to speak, and used a speech therapist for a time after he learned his first words.

Later, Malachi started talking to himself, repeating the same words over and over while he worked them out in his complicated brain. When he was a teenager, he started talking more roughly to himself, pounding his fist in the wall, having even more trouble focusing. His counselors don’t know how to label him. At different times, he has been diagnosed with learning disabilities, speech delays, a perceptual disorder, pervasive developmental disorder, brain injury, mild mental retardation, depression, schizophrenia, and disassociative disorder. He doesn’t fit into a neat little box. But who does? At this point, I give up. My parents, still trying to find services to help him and guidance for their work with him, need a diagnosis. Otherwise, how will he be eligible? How will they know what interventions to try next?
"Malachi"

When we were little, there was special education for Malachi, but he was intelligent, with no physical disabilities. The services offered to him were very limited. There was certainly no respite. When I turned 6, I entered public school with Malachi. My parents had recently gotten divorced, and my mother had to work, but she could not afford any of the child care options available to us.

We became "latchkey children." Malachi and I always came home right after school. Malachi always met me at home, and he went in first to check the closets and the basement for monsters and kidnappers. Together, Malachi and I did our homework and the chores that Mom laid out for us the night before. I am sure that, when Malachi thinks about that time, he remembers me as bossy. We bickered A LOT. (Imagine that! Siblings bickering?) We wanted to make sure that Mom would be in a good mood when she came home, and I knew that Malachi needed to be told to do things more than once in order to make sure that they got done.

When Malachi started experiencing new symptoms, I was starting high school, and I was given the opportunity to tune out. I got VERY involved in school activities and spent little time at home. My mother, exhausted by Malachi’s constant needs and frustrated because she still did not know what was going on with him, just could no longer have him in the home. Malachi moved to my Dad’s. For many years afterwards, I did not really know him or the
person that he became. Maybe if my life with him had not been so intense when we were younger, I would have stayed connected with him during that time. My respite in high school was WONDERFUL for me, but it came too late in my relationship with Malachi. As I work to reestablish my relationship with him, I have a lot of regret. I can look back and say that respite would have been very important to us all at a crucial time in our lives.

So, in Malachi, I have a special sibling. I never had a brother who screened my dates or scared off my bullies or who my friends had crushes on when I was younger. When my parents divorced, I never talked to Malachi about my anger, and he will not celebrate my wedding day the way that I always felt a brother should. We might never have a deep meaningful conversation about our life paths. I will always miss all those things with him. But Malachi is one of the greatest teachers I ever had, and, these days, I am learning to enjoy all of the things we are able to do together and appreciate all of the things that he is. He is a very special sibling.

— Anonymous, 1997, Sibling

A Sibling Perspective

My brother David Eugene Ownby was born with multiple disabilities—profound mental retardation, epilepsy, cerebral palsy, and microcephaly. I was 3 years old when my parents brought him home from
the hospital. They gave me a life-size doll that day -- but I wanted the real baby. I always took my responsibility of being the big sister seriously. I made sure that I "protected" David. When other kids made fun of him, they quickly got an education in disability awareness. I made sure they knew that David had feelings just like the rest of us. Although he could not talk, he still knew when someone was making fun of him. Little did I know I was becoming an advocate.

David required a significant level of care. Both my parents worked, and, while most any babysitter would watch me, few wanted to watch David. My parents tried many variations of baby-sitters, from live-in nannies to next-door neighbors. Most often sitters would watch me and David for a week or two and then David would have a grand mal seizure. That coupled with the rush to the emergency room would usually result in the sitter agreeing to watch only me and not David.

Respite was not an option for my family in the early seventies. While my family worked with many therapists and social workers, the only options given to them were permanent foster care or institutionalization for David. Neither sounded very appealing, but my parents' ability to function as a family was severely compromised by the time David was 5 years old. My parents needed a break from the demands of caring for David. They approached the social workers about institutionalization.
"state of the art" institution had just been built 8 miles from our home. This service was touted as the best option available. Even still, the waiting lists were long, and David was denied admission time and time again. My father knew Senator Hubert Humphrey and, with his political clout, my parents did what they thought was the best thing for David and my family -- institutionalized David.

While having David cared for on a daily basis by caregivers in the institution was a relief, my family quickly learned that institutional life was not conducive to normalization -- it was not a home. My mom quit her job at the Library of Congress to manage the institution's library and be close to David. I quickly became involved with Special Olympics and volunteered there most Saturdays. My family and I stayed actively involved in David's life, bringing him home on the weekends and visiting often during the week. While this seemed to work for our family, I always felt that David was cheated. I did not think it was fair that he had to live there. I vowed at age 7 to "grow up" and change things for people like my brother.

I had to wait until I was 18 years old to become employed at the institution where David lived, and, when I did, I got a job there. I wanted to know what went on behind closed doors. While my mom had some sense of institutional life, she was far removed from the living units and day-to-day activities. I was employed as a direct care aide. I quickly learned
what I had always felt — institutional life should not be an option for people like David. This environment was volatile, violent, uncaring, and stripped residents of their basic human dignity. As a young adult, I vowed to find David a better living situation. I wanted to become a disability rights advocate and work to change the system, allowing real choice for families and people with disabilities.

In seeking services for David, I faced what so many families still face — closed doors, long waiting lists, and lack of services. The care David received at the institution was never good enough for me and had gotten considerably worse. More frequently, my parents and I met David at the emergency room with unexplained broken bones, bumps, and bruises.

In May 1992 David was left unattended in his institution bedroom after a series of grand mal seizures and aspirated. When institution aides found him, he was unconscious, with no respiration or pulse, and was cyanotic. He was rushed to the hospital and lived on life support for 2 months.

David had taught me long ago to never give up. He was certainly a fighter. He always bounced back from an injury or illness, and there were so many — we all thought for sure he’d live through this too. While David struggled to live, this fight was too hard. In the time he had been left unconscious, he suffered significant brain damage. He was now blind, paralyzed, unable to swallow, and all the large
organs in his body had begun to atrophy. My little brother was dying and my advocacy was too late.

David did die, but not in vain. While my efforts to help him move into a group home failed him, his death, my advocacy, and many others’ resulted in 80 other residents having the real choice to live in a home. While I advocated to close this institution, after 5 years it remains open, but considerably smaller, better staffed, and closely watched by agencies providing oversight.

I learned from David to laugh, to have fun with the little things in life, and to persevere. I held David’s hand as he lay dying and promised him I would work to ensure that no one else would have to suffer this way and that I would commit my life to making real choices for families and people with disabilities. I promised to share his story and make sure as many people as possible knew him. While David and I can no longer do McDonalds on a Saturday afternoon, and I’ll never again have a big bear hug from him, he is alive in the work that I do. His life has given me my perspective.

— Lisa Ownby, 1997, Sibling

One Caregiver’s Perspective

Respite caregiving for children with disabilities and their families truly presents an opportunity for a "give and take" relationship. This has been my experience from the time I decided to attend group
training, through working as group caregiver, and finally when a family "match" occurred.

My initial interest in respite was both personal and professional in nature. Being a single person without children left me feeling the need for contact with children, and my work in the social/health care field made working with children with disabilities a natural. I truly felt that I had personal traits of an affinity for children, patience, and a desire to learn and help a family. At that point, I signed up for training without knowing the specifics. What an informative, supportive, friendly, and nonthreatening experience!

Following initial training, small group exposure to children and the opportunity to discuss the children’s needs and personalities with their families gradually increased my confidence and my joy in understanding that children with disabilities appreciate interest, hugs, snacks, toys, walks, playing ball, swings and slides as well! Challenges are there too! They may come in the way of feeding, behavior challenges, toileting, or special wheelchair accommodations, which may initially be unfamiliar to caregivers.

This is where the "give and take" occurs on the part of caregivers and families. Caregivers need training, feedback, and encouragement from the family "experts" in order to best understand the child’s limits and special needs. Caregivers may appear to ask many questions, seem anxious, and
anticipate what to do in an emergency. I believe families can aid caregivers by remembering their own initial thoughts and feelings about helping their child with his/her limitations. On the other hand, caregivers may want to appear knowledgeable and confident and not ask too many questions! Families, please anticipate this and VOLUNTEER INFORMATION!

My caregiving experience with a 9 year old and his 6-year-old brother has truly been rewarding and fun! This match occurred following group respite in which I had the chance to interact, observe, and build a relationship with both brothers and parents. A need for individual respite was there, and I had the time and geographic availability. Truly a match made in heaven! I really feel that this family provided information, training, and demonstration about their son’s needs. I had the opportunity to question, observe, and build a relationship with the parents as well. I believe that caregivers need to maintain contact periodically in between respite dates to develop rapport. This also helps families feel freer to call upon the caregiver, particularly families that typically “do it all” or have difficulty in asking for help. Families and caregivers need to communicate and appreciate the gifts each has to bring to this partnership. The Partners Plus program is truly partners in action!

— Linda Malone, 1997, Caregiver
Stress

The potential for stress exists in all of our lives. Stress may be good or positive stress (eustress), or bad or negative (distress). The key to coping with stress depends on how you perceive a situation or life event. Stress is also closely related to your view of how hard or easy it may be to manage the event. For example, going on vacation can be perceived as both good and not so good. Many of us welcome vacations! Yet, vacations may be a major source of stress if you consider the packing, the drive, making reservations, or scheduling time to be difficult. Many strategies are available to both families and caregivers to help cope with daily stress. Included in the following pages are things families and caregivers can do to reduce stress, prevent burn out, and ensure successful respite relationships.

The Special Challenges of Parenting

Being a parent involves many changes, which may be perceived as stressful. Families who have children with special needs share the stresses of all families facing the challenges of parenting. In addition, they may have other challenges related to their children’s special health, education, and care needs. Many resources are available to help families cope with these special challenges.
The Special Challenges of Caregiving

Being a caregiver for children with special needs can be stressful too. Caregivers are responsible for the children in their care. If those children need extra attention or medical care, caregivers are more challenged. In addition, it takes time to become comfortable with children and familiar with their needs and strengths. Not knowing what to expect from children can be stressful. Caregivers who are still learning about the children in their care may experience stress.

Coping with Stress

If you perceive a situation as stressful or hard to manage, you can reduce your stress by using a variety of coping strategies. Consider some ways you deal with stressful situations or special challenges. You might already know strategies that work well for you. For example, it might help to talk with family and friends in hard times.

Families often use respite to cope with stress. Taking a break from daily demands not only is relaxing, it allows families time to try the strategies listed on the following page. Families may want to talk with caregivers about how they use respite time to cope with stress.

Caregivers may need specific ways to cope with the challenges of working with children with special needs. Caregivers and families need to take time to talk after respite takes place. Was there anything that was stressful? Is there more information families can share? Are there...
new skills the caregiver can learn? Certain caregivers may want soothing music playing as they work with your child(ren) or use other methods to create a calm atmosphere. Families can encourage caregivers to talk about how they handle stress. Together, caregivers and families can work out ways for caregivers to relieve stress during respite. The following fact sheet lists strategies you may find useful in reducing your stress.

**Strategies for Dealing with Stress**

People deal with stress in many ways. Here are some examples:

- Seeking information
- Sharing information
- Learning new skills
- Doing something that makes them laugh
- Listening to something they find soothing
- Planning ahead
- Finding a favorite cartoon that brings a smile to their face
- Walking, jogging, hiking, swimming, playing
- Communicating openly
- Taking a break
The Challenges of Respite

Building effective partnerships takes time and skill. Families and caregivers need to know how to work together for successful respite care.

One problem families often face is keeping caregivers. Families tell us many respite caregivers cancel appointments. We also know that many caregivers stop providing respite. Respite caregivers cancel or quit for many reasons. Why do you think they quit? The information in this section will help you understand why and help you develop strategies to build successful respite partnerships.

Caregivers may stop giving respite for the following reasons:

- Respite is harder than they expect.
- Time demands are too great.
- Physical demands are too great.
- Burnout occurs.
- Training was not adequate.
- Patience wears out.
- They don't feel appreciated.
- They are not paid enough.
Meeting the Challenge

Meeting the Challenge

Taking these steps can help you meet the challenge of respite:

- Help caregivers to know your family.
- Help caregivers to know your child.
- Communicate your purpose for respite.
- Develop and carry through with a thorough training plan.
- Avoid overuse of a caregiver.
- Show your caregiver respect.
- Show your caregiver appreciation.
- Be clear about your caregiver's terms for providing care.
- Be considerate with schedules.

Little Things Can Make a Difference

It is important to recognize and appreciate your caregivers. Here are some ways you can show them what their caregiving means to your family.

- Send notes or tell your respite caregivers what a difference they are making in your family's life.
- Tell them how well they are doing a specific task.
- Tell them how you appreciate the extra effort they make.
- Remember your caregiver on special days.
Send or give them pictures made by your child.
Do things for your caregivers that would be meaningful to them.
Write about your caregiver and put the article in a newspaper or newsletter.

Successful Family and Caregiver Partnerships

Many components are helpful in establishing and maintaining successful partnerships. Families and caregivers should strive to accomplish these goals:

- Appreciate differences in skills and approach.
- Be willing to learn from each other.
- Ask each other for help.
- Allow time to talk with each other.
- Share experiences.
- Be willing to work together, even when differences occur.
- Be flexible in trying new ways.
- Have a sense of humor and fun.
- Be patient when things are not clear or exact.
- Share commitment to children with special needs.
Notes on this Section
Section 4

All Kids Like Cookies

activity
All Kids Like Cookies

All Kids Like Cookies

Children with special needs are important members of respite partnerships. The following activity is designed to allow you the opportunity to experience some of the daily challenges children with special needs face. This may be useful in understanding a child's perspective and keeping that in mind when developing respite partnerships.

We know children with special needs are more similar to other children than different. "All Kids Like Cookies" is a series of activities designed to help persons without special needs experience challenges children with special needs face every day. You can do the activities on the next pages by yourself or with a group. Answer the questions at the end of each page as you finish each activity. This will help you become more aware of what it is like to have special needs. As you will see, from the perspective of the child with special needs, the world is a more challenging place.

We cannot help until we understand.
We cannot understand until we feel.
We cannot feel until we become.
(Council for Exceptional Children)

Adapted from:
All Kids Like Cookies Activity Sheet

The following activities can be done alone or with a group. During the activities you remain yourself, but a special challenge is added. Think about how you feel when doing each of these activities and be aware of any adaptation you make.

To experience a visual impairment, you will need:

- a blindfold
- a pitcher of water
- an empty glass
- a crayon or other marker
- a page out of a coloring book or a crossword puzzle
- a child’s storybook or magazine

- Put a strip of cloth over your eyes and tie it around your head to make a blindfold.
- Pour water from the pitcher into the glass.
- Color the coloring book page or complete the crossword puzzle.
- Read the storybook or magazine.

Questions

- How did you hold your head while pouring water, coloring, and reading?
- If doing this activity with a group, did anyone talk you through the tasks?
Activity Sheet

- What other senses did you use to help yourself? (touch, sound, smell)
- Did it matter that you were coloring out of the lines or what color you were using?
- Were you able to complete the crossword puzzle?

Vision helps a child learn about the objects and people in the world around them. Few children are totally blind. A child is considered to have a visual impairment if the child's ability to see after correction limits his ability to learn through vision.

To experience a speech and language impairment, you will need:
- a bag of large marshmallows and gauze
- one other person or a small group

- Put enough marshmallows or gauze in your mouth to fill it; two or three usually work.
- Say your name and give your address or phone number.
- Ask your partner or others in the group to bring you something.
- Tell your partner or others something you did today.
Questions

- How were your words spoken differently?
- How did you feel when others did not understand you?
- If you spoke this way all the time and others laughed at you, would you want to talk very much?

Sometimes children with speech and language impairments stop talking rather than cope with the difficulty of getting people to understand them.

To experience a physical disability that affects using the smaller muscles in your body, like the ones in your fingers, you will need:

- a pair of large garden or work gloves
- a puzzle
- a string and macaroni with holes or beads
- very small toys that have moveable parts

- Put the gloves on.
- String the macaroni or beads.
- Put the puzzle together.
- Play with the toy.
- If you are wearing a watch, take it off and put it back on.
- If you are wearing earrings, take them out and put them back in.
Activity Sheet

Questions
- Were you able to pick things up easily?
- Did you need more time to put the puzzle together?
- Were you able to put on your watch and/or earrings?

To experience a physical impairment that affects the large muscles of the body (like the ones you use for walking) you will need:
- a kick ball
- a large toy like a stuffed animal or truck
- a video

- Place the ball between your ankles and keep your hands folded together.
- Walk 10 feet with the ball between your ankles and pick up the toy or video.
- If doing this with a group, hand the toy or video to someone else.

Questions
- Did you take a long time to get to the toy or video?
- Did you feel awkward?
- Did you drop anything?
Children first learn to explore their surroundings by using their large muscles to move around. Children learn how things work by using the small muscles of their hands to explore toys.

Children delayed in their motor development have difficulty learning to become physically independent because they cannot move well. Children with small muscle development delays may have difficulty feeding and dressing themselves or learning how things work because they cannot use their hands and fingers well.

Source:
Notes on this Section
Section 5

finding respite
caregivers
Finding Respite Caregivers

Preparation

Finding a good caregiver is often difficult. Many people decide to go out for dinner, take a quiet walk, run errands, or attend a sports activity, then consider their choices and plan for child care. But what if you have no plan?

Planning ahead and putting in the time and effort to train individuals to care for your child's specific needs will give you regular respite. Over time, you will come to trust and rely on your caregiver's help. On the other hand, if you wait until an emergency or crisis develops, you will feel helpless, isolated, and overwhelmed.

For Example:

Miguela and Tony want to go shopping for the evening. They would like to go without their daughter Angela, who has cerebral palsy. They cannot simply call the 20-year-old college student, Pat, who lives next door because a caregiver needs special training in helping Angela move and feed herself. Or can they?

Yes, Miguela and Tony can call Pat to provide child care, if they plan ahead.

In developing their plan for respite, Miguela and Tony need to recruit several people, including Pat, who are interested in providing care for Angela. Then they need to provide training to these caregivers. The advantage of having several trained caregivers is that caregivers feel more comfortable saying no when they need to, knowing that a family has someone else to call.
When you have a child with special needs, it is important to develop a plan so that you can have time away from the demands of parenting. Partners Plus can help you develop a plan for recruiting and training respite caregivers.

Before looking for caregivers, think about the type of respite your family wants. Families have many choices for what respite will be like. Evenings might be the best time for you to get respite, or perhaps a morning out. Maybe you and your family need to spend a weekend away. Your family might want a combination of all three.

You might want respite in your home, or you may prefer to find someone to care for your child out of your home. You may want to enroll your child in a before- or after-school or recreation program in your community. You may want to build a respite cooperative with other families who have children with special needs. In a cooperative, families trade care.

The benefits and disadvantages of in- and out-of-home care and of cooperatives are listed on the next few pages.

**In-Home Respite Care**

**Advantages**

- You do not have to arrange transportation for your child or for any special equipment needed.
- Your child does not have to adjust to a new environment.
- Other children in your family can also be cared for at the same time.
Out-of-Home Respite

Disadvantages

- Having a respite care provider in your home can feel like a loss of privacy.
- You may have to arrange transportation for your respite care providers.
- Some respite care providers may already care for other children in their homes but may not be available to come to yours.

Out-of-Home Respite Care

Advantages

- Your child may have an opportunity to interact with other children and adults.
- When your child is in out-of-home respite, other family members can take a break but still be at home.
- Your child has an opportunity to be included in community programs.

Disadvantages

- Out-of-home respite may feel very different from usual baby-sitting.
- It may be hard to leave a child in an unfamiliar setting.
- Children may have a difficult time adjusting to a new place.
- You may have to provide transportation.
- Medication and special equipment have to be packed and taken along.

Adapted from:
Family Cooperative Respite Care

Advantages

- Families provide respite care for each other instead of paying out of pocket.
- Parents may be more comfortable leaving their child with a person who is experienced in caring for a person with special needs.
- Cooperatives build support networks for families.

Disadvantages

- It may be an additional demand on parents.
- It may be difficult to work around family schedules.

Finding Respite Caregivers: Circles of Family and Friends

One way to help you think about your respite care recruiting plan is to consider your natural support network. You might call this network your circles of family, friends, and community programs. Consider yourself at the center of this network. Surrounding you are people from various groups. These groups interact with your family in some way to provide different levels of support. Think about your circles of support and the people in these circles.

For example, one circle is the people in your neighborhood. Who are your next-door neighbors? Who lives down the street? Do your neighbors know anyone who might be
Circles of Family and Friends

interested? What about the circles of support in your church or synagogue? Your child’s school? Your place of work? Are there extended family members who would be more likely to provide child care if they were adequately trained? What community programs are available?

Another way to find people from your circles of family and friends is to invite several parents of children with special needs to your home to share ideas about what worked for them. Begin by drawing the circles and labeling them. Then, think of people you might ask. You may also consider asking your service coordinator, case manager, or child’s teacher to help you in this activity.

By identifying even one person from each circle, you have started a list of possible respite caregivers. Now, find some time to talk with them about Partners Plus and ask them if they would be interested in learning more.
Circles of Family and Friends

Our Family

Extended Family
Before/After School Programs
Neighbors
Other Parents
Recreation Programs
Work
Camps
School
Place of Worship

Partners Plus Family Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Other Ways to Find Respite Caregivers

Partners Plus helps families think about groups of different people who may be interested in providing respite care. Recruiting other parents to start a cooperative may also become a part of your plan. As mentioned earlier, cooperatives offer families another option for child care.

As you move beyond your circle of friends, neighbors, family, and community programs, consider other people in your community (e.g., workers from community agencies, local universities, or child care centers). Community agencies may maintain a list of people interested in providing child care or know of organizations or programs that do. A list of community groups that may be able to help you find a respite caregiver follows.

Resources in Your Community

The following organizations in your community may help you get in touch with caregivers:

- Child care resource and referral agencies
- Home child care providers
- Child care centers
- Schools
- Mental health/mental retardation agencies
- Home health agencies
- Hospitals
- Civic or community groups
- Senior centers
Choosing Potential Caregivers

In the Partners model, families and caregivers choose each other for a respite care partnership. This takes place by a natural selection in much the same way as we choose our doctors, dentists, preschools, neighborhoods, places of worship, and friends. In making these selections, we are looking for a good match with our own values, beliefs, and ways of doing things. Families often make their choices based on special qualities they want in an individual respite caregiver, or a community program that offers respite through group care, such as before and after school programs, camps, or recreational activities.

Partners Plus offers opportunities for families to choose potential caregivers. In group training, families and caregivers meet, talk, and listen to each other. Partners Plus can help families and caregivers get together through social events such as potluck dinners or picnics, giving families a chance to see how their children and potential caregivers interact together.
Notes on this Section
Section 6

interviewing caregivers
Interviewing Potential Caregivers

caregivers meet, talk, and listen to each other. Partners Plus can help families and caregivers get together through social events such as potluck dinners or picnics, giving families a chance to see how their children and potential caregivers interact together.

For communities that choose to maintain a caregiver directory, each Partners Plus caregiver fills out a caregiver profile. These communities would designate a place to keep each profile on file for families. Communities may also maintain a list of programs where respite can take place.

The profile is much like a resume or job application. It gives a family information they can use in deciding if they want to get to know a caregiver better through an interview. An interview is a chance for both family and caregiver to ask questions and learn what each expects from the other.

A sample caregiver profile is included in Section 8. Families can duplicate this and ask caregivers to complete prior to an interview as part of the interview process.

Interviewing Potential Caregivers and Caregiving Programs

Interviewing is an important step in choosing a caregiver or caregiving program. Take some time to think about your respite schedule and respite needs before you begin to interview caregivers or community programs.
Many families have told us that selecting a good caregiver or child care program is sometimes hard for them to do. We have included some simple tips on interviewing to help you make a good match for your family.

1. **Be prepared.**

   One key to a more relaxed interview is to be prepared. Prepare yourself by reviewing the caregiver profile or program description. Questions on this form ask the potential caregiver or program director to provide information on education or training in child development or child health, first aid, CPR, or other certifications they may hold. Potential caregivers and programs are also asked to describe why they are interested in providing respite care or including children with special needs into their program.

   Make notes as you review the caregiver profile or program description. What else would you like to know about the person or program? Do you want to know more about a caregiver’s or program staff’s education and training? Do you want to learn more about their experiences in working with children? You might also want to ask when a caregiver or program would be available to provide respite.

2. **Decide how to conduct the interview.**

   Some parents are more comfortable doing a telephone interview first to screen potential caregivers or programs. If, after the initial contact, you decide you want to consider a person or program to provide care, you can schedule a meeting time for a
Interviewing Potential Caregivers

face-to-face interview. You might also consider whether you would like to have your child present at the interview.

3. Break the ice.

A good way to begin the interview is to ask potential caregivers or program directors about their interests and experiences. For example:

I see from your caregiver profile that you are interested in respite because... or,
I see from your caregiver profile that you already have some experience caring for children with special needs. Tell me what that was like for you.

If you are interviewing a program director, you will want to know more about the knowledge and skills of existing staff in including children with special needs. You may want to ask how new staff are trained to help children with special needs participate in all activities.

You will also want to ask questions about the program's accessibility as it relates to the individual needs of your child.

4. Ask open-ended questions.

Another good way to gain more information is to ask open-ended questions. For example, if you ask a person Do you believe in punishing a child?, you are asking for a response of yes or no. But, if you ask How do you handle difficult behavior?, the person can give you much more information on their feelings and specific examples. Some sample interview questions are included on the following pages.
5. Take notes.

As a potential caregiver or program director begins to share information, jot down some notes. Taking notes will help you to ask follow-up questions. It will be helpful to review your notes when you consider potential caregivers or programs later. You might decide ahead of time that your spouse or partner will take notes while you ask questions or vice versa. You may want to tape the interview instead. With a tape, you can listen to the caregiver's answers more than once. If you choose to tape, ask the potential caregiver for permission to tape-record the interview.

Sample Interview Questions

You may wish to ask several of the following questions when you interview potential caregivers. Adapt these questions when interviewing a community program director.

- Tell us about your education and/or experience in working with children.
- Discuss your experience working with children with disabilities.
- Why are you interested in providing respite care services?
- Describe your strengths. How would you describe yourself?
- Why are you the right person for the job?
Sample Interview Questions

- What are your thoughts about families who have children with disabilities?
- What would you do if we disagreed about something?
- If my child had to be taken to the hospital for an emergency, what steps would you take?
- How do you believe children’s difficult behaviors should be managed?
- What would you do if a child deliberately hit another child?
- What would you do if you found a child with an open bottle you suspect is poison?
- Are you willing to take my child out on activities?
- Describe a difficult problem you’ve had to handle with a child. How did you handle it?
- Tell me about any special training or experience you might have.
- How would you know if my child likes a particular food or activity if he cannot speak?
- Are you willing to have a criminal background check done?
- Do you have CPR, first aid, and/or medication administration training?
- Do you have a valid driver’s license?
Some Additional Questions To Ask
Directors of Community Programs

- How do you encourage children to play together?
- How do you teach children to ask other children to play with them, share toys, take turns, express affection, and help other children?
- How do you help children to control aggressive behaviors?
- What kinds of small groups activities do you have that require cooperation and sharing?
- What agency or organization provides oversight or certification for this program?
- Do all staff working in the program have criminal history background checks?

Adapted from:

Questions Caregivers Ask

Caregivers may ask these questions about what will be expected of them. Families should be prepared to answer them during interviews.

- Will I be expected to care for siblings as well?
- Will the parent be at home or away while I am providing care?
What To Do After the Interview

- If I receive a fee for my caregiver service, how will I be paid?
- How much will I be paid?
- What kind of training will I receive?
- How often will respite be needed?

Program directors often request families fill out an application that provides them with the kind of information they need for all children. There may be additional questions they ask that are specific to the needs of your child.

After the Interview

1. Reference checks

   It is a good idea to check the references of a potential caregiver. The caregiver profile asks potential caregivers to provide the names of two people. You may choose to ask for more references. A Reference Check form is included in this manual as a guide to help you know what questions to ask.

2. Criminal record checks

   To obtain a criminal record check on a potential caregiver, you can contact your state police. There may be a charge for the service. Criminal record checks are limited in the information they provide.
3. **Follow-up**

Be sure to thank potential caregivers for their time and explain how and when you will decide. You might say,

- We have decided to interview three people and do reference checks for each.
- We hope to have a decision in one week and will let each person know of our decision by phone or by a note in the mail.

4. **How to Say No**

Sometimes it is difficult to say no to a person. If you decide a potential caregiver is not the best person, be honest and direct. You might say,

- I was fortunate to have interviewed several qualified caregivers such as yourself and we have selected another person that we think best meets our needs.
- Thank you for your time and interest, but I don't think this would be the best match for us.

You may feel more comfortable telling a caregiver in a note, rather than over the phone.

**Additional Things to Consider**

Most people who want to care for children are responsible and love this work. Others may not be good choices to work with children. **Trust your instincts!** If you are uncomfortable
Other Questions To Ask

with a caregiver, he or she is not the right caregiver for your child. Please use the following to help guide you in selecting a caregiver. There are a few things to look for when talking to caregivers that might alert you to some concerns. Adapt the following questions for appropriate use for community program staff.

A Few Questions to Ask

- **Why do you want to care for my child? What do you hope to gain from this?**

  Many people will answer that they love children. Some will say that they want experience working with children with special needs. Others will tell you that respite was important for their families. Some people want to make a difference in their communities by providing respite.

  These are all the right words to use. Be sure to watch body language closely. Listen for answers that make you uneasy.

- **How do you deal with stress/anger?**

  (Give the caregiver an example. What can be stressful about caring for your child? For example, does your child show anger by hitting? Ask the caregiver how they would deal with that stress.)

  There is some stress involved in any child care setting. There are many different ways to handle stress, some positive, some not. Be sure you are comfortable with the answers you get.
• Tell me about any lessons you learned from the way that you were raised that influence the way you interact with children.

Often, people care for children the same way that they were raised. You may want to ask other questions that get the same information: Tell me a little about yourself. What is the thing that you remember most about your childhood? Sometimes, people can work through bad experiences as children to become loving, capable caregivers. If you have concerns about something a caregiver tells you about his or her childhood, ask about how it affects the ways they might take care of your child. Again, be sure to watch their body language.

• Will you be taking any medicine that will affect your ability to care for my child? Do you have any physical limitations that might affect your ability to take care of my child?

These questions might give you some information about the medical and emotional history of a caregiver. Feel free to ask a caregiver who will be taking medicine what those medications are and how they will affect him or her. Find out if there are any side effects to the medicines and what they are. Also, talk about where medicines will be kept so that the child cannot get to them. Remember, a medical history might not mean that a caregiver is unsuitable.

• Tell me about how you discipline your children. How would you discipline my child if he were not behaving?
Noticing Warning Signs

Watch for answers that do not agree with the way that you want your child to be disciplined, especially anything that involves physical contact. This might be a good time to talk with a caregiver about how and when you discipline your child.

- **How long have you lived in Virginia?**
  
  Remember, criminal history checks only give information about convictions in this state. If a person has moved around a lot, ask why. Again, watch body language!

### Some More Things to be Alert to When Interviewing and Screening

- **Unusual interest in your child’s medication(s)**
  
  Remember, a person may need to know a lot about the medication a child takes. We encourage caregivers to ask as many questions as they need to make them comfortable in working with children. If a caregiver’s questions do not seem to relate to understanding your child, you may have some concerns.

- **Curiosity about where liquor or firearms are kept in the house**
  
  Again, a caregiver may be asking these questions because they want to keep your child safe. However, if caregivers want to know where the keys to the liquor/gun cabinets are kept, you may need to ask why they need that information.
References cannot give positive information about a caregiver

Caregiver wants to care for a child only of a specific age or gender
If a caregiver states such a preference, ask why. Watch body language!

Caregiver does not want you to visit while she is providing care
Suggest to a caregiver that you might drop by while she is providing care. If a caregiver is not comfortable with your dropping by, find out why. Remember that caregivers must plan for their own lives. They will need to know when you plan to be back so they can plan the rest of their day.

Things to Look for When Visiting a Caregiver’s Home

• A lot of video equipment/cameras
• Books or pictures of children that make you uncomfortable
• Medicine, alcohol, or drugs out in the open
• Weapons not stored safely in the home
• Room arranged in a way that does not allow children to play freely
Visit the Community Program

Look at the space where your child will spend time in respite. Make sure that the environment is safe and comfortable for your child.

Is there room for your child and your child's equipment? Are there a lot of sharp corners in the room? Can your child play in this room? Are there fragile decorations that your child might bump into? How will the caregiver protect items so your child will not break her things? Will your child be kept in a small room or have freedom to move from room to room?

Things to Look for When Visiting the Community Program

- Too much attention from adults that may interfere with the children's interactions
- Are children with disabilities situated next to children without disabilities so they may interact easily
- Do all children get to lead activities and be successful in front of each other
- Are the equipment and play materials safe and in good working order
- Are the toys and supplies appropriate for the age group and their abilities
- Are the number of toys available limited intentionally, in order to encourage children to share and engage in the same activity

Adapted from:
Things to Look for After Care Has Started

There are few signs to watch for in your child after care has started that may be signs that he is poorly treated in respite.

- Eating a lot less or a lot more
- Sleeping a lot less or a lot more
- Unusual fear of adults
  Children from around 5 months to a year typically go through a period of stranger anxiety. At this time, children are learning to separate people they know well from those they do not. At this stage, it may take time for children to adjust to new caregivers.
- Sexually acting out with younger children or toys
- Increased violence or anger
- Rashes and bruises that are not explained
- Unexplained stained and/or torn clothes or underwear

A Few Final Words

It is very important that you feel comfortable with your caregiver. If the answer to any question makes you uneasy (even if it is the answer to only one question), then that
Evaluating Your Respite Decision

person is probably not the right caregiver for your child. Your instincts might give you the strongest clue about how well a caregiver will work for you and your family. Trust yourself.

Summary

Now that you have finished recruiting, screening, and interviewing, you are ready to select your caregiver.

Consider these questions about caregiver(s) you have selected:

• Will she feel comfortable caring for your child with special needs?

• Will she be willing to spend time learning how to care for your child?

• Will she be willing to learn about your family?

• Will she be warm, affectionate, and enjoy children?

• Will she have respect for your beliefs and values?

• Does she have good character references?
Section 7

developing your
personalized
training plan
Developing a Personalized Training Plan

The Partners model is based on the belief that families know how to care for their own children and that they have the skill and knowledge needed to train their own respite caregivers. The family and caregiver together decide the specific amount of personalized training needed to care for a child. Training time will vary according to child, family, and caregiver needs.

Remember, caregivers learn in many ways. The Caregiver Manual asks caregivers to think about how they learn best. Before you begin training, talk with your caregivers about how they think they will learn your child’s routine best. Work with your caregivers to develop a training plan that blends your teaching style and their learning styles.

Some caregivers may not be sure about how they learn best and may want you to go ahead with your training plan. During this process, caregivers may find certain training techniques that work best for them. It is important to be ready to change your training plan to meet everyone’s needs.

There are four simple steps to help a family develop a personalized training plan. This training plan is helpful for all caregivers, those providing individual and group care, in families homes or community programs.

These steps are explained on the following pages.
Step by Step: A Guide to Developing Your Personalized Training Plan

To develop your training plan, follow this step-by-step guide. The worksheets and examples in this section will help keep you organized. Clean copies of all worksheets are included in Section 8.

Step 1: Identify training goals.
What should your caregiver learn about your child and family routines?

Step 2: List training methods.
What teaching methods will you use?

Step 3: Use resources to help train caregivers.
What information will you use to help your caregiver learn?

Step 4: Evaluation
How will you and your caregiver know when your caregiver has learned a specific skill or routine?
**Identify Training Goals**

**Step 1: Identify training goals.**

Partners Plus is based on the belief that families can train their own respite caregivers using their child’s daily routines. First consider what part of the day you want respite. If you think you will schedule respite in the morning, then the morning breakfast routine would be helpful to use as a basis for training. If you think you are more likely to schedule respite in the evening, then dinner and bedtime routines should be considered in developing your training.

Your training goals will be based on the daily routine that you selected for training. Goals are broad statements of what you want to achieve or do. For example, you decide that you most likely would have respite care take place during lunchtime between noon and 1:15 p.m. Your goal, therefore, may be for your caregiver to be able to feed your child lunch.

If your child will be participating in a community program, you will need a clear understanding of the schedule of activities for children enrolled in the program. You and the staff will need to communicate well in developing a plan for training program caregivers to include your child in the daily routines.

The number of training goals each family has will vary and will be determined by the needs of the child, his or her routine, and the length and time of day involved.

What do you want your caregiver to learn about your child and family routines?
I want my caregiver to learn about these routines:

Goal 1: (Example) My caregiver will be able to feed Maria lunch.

Goal 2: ____________________________

Goal 3: ____________________________

Goal 4: ____________________________

Goal 5: ____________________________

Step 2: List training methods.

At this point you should have at least one training goal. Now it is time to consider how you will train your caregiver.

One family tells us that they train their respite caregiver by modeling and shadowing. The family asks a caregiver to spend several hours with them during a period of the day such as lunchtime. This is a time when many routines can be observed. This gives the caregiver the opportunity to watch...
how the parent interacts with the child during these routines. We call this *modeling* the ways you want someone to care for your child. By spending several hours with the family, the caregiver can also watch play activities and a nap routine. On the next visit the caregiver helps the parent, or *shadows*, in the caregiving routines. This gives the caregiver another learning experience.

Another family tells us that they train their respite caregivers by making a videotape of their child's daily routines. This gives the parent the opportunity to select specific routines to tape for training. It also gives the caregiver the opportunity to view the videotape, write questions, view the routines with the parent, and discuss.

It is important to think about all the different ways you might teach your respite caregivers so they can learn how to provide quality care for your child. Will you have the caregiver learn to feed your child by watching you? Will you want the caregiver to assist in feeding your child? You may also want to ask the caregiver questions about his or her skills and comfort level in feeding your child. There are many ways to train and you have many options available to you. Some suggestions include the following:

- Modeling (showing)
- Videotaping (another way of showing)
- Shadowing (helping)
- Talking, telling, describing
- Writing (another way to tell or describe)
• Answering caregiver questions

• Sharing ideas

Even if you choose other ways of training, it is a good idea to have important information and instructions in writing, in case your caregiver needs help remembering what you have discussed.

What methods will you use to teach your caregivers?

In order to reach these goals I will use the following methods:

Goal 1: My caregiver will be able to feed Maria.

Methods:

1. (Example) My caregiver will watch me feed Maria at lunchtime.

2. 

3.
Goal 2: ______________________________

Methods: ______________________________

1. ______________________________
   ______________________________
   ______________________________

2. ______________________________
   ______________________________
   ______________________________

3. ______________________________
   ______________________________
   ______________________________

Step 3: Use resources to help train your caregiver.

It may be helpful to gather resources you can use when training your caregiver. There are many, many information resources available to us. You also may want to network or get in touch with others, so you will have links to new information.
Locating Training Resources

You may want to give a caregiver written information about your child's diagnosis, other special health care needs, medications, or sibling issues as a part of your training.

Will you tape a home video of a lunchtime routine? Would you like to give your caregiver any handouts on special feeding issues similar to your child's? Do you have information that you want to share on special feeding techniques from your child's occupational therapist?

The following is a list of organizations and services that you may want to contact for resources that you can use in training.

Where to Find Resources:

- Local early intervention programs
- Doctors, nurses, therapists, social workers, and teachers
- Local, college, and university libraries
- Parent Resource Centers
- Parent to Parent Networks
- Local hospitals
- Parent Training Information Centers
- National Information Center for Children and Youth with Disabilities (NICHCY), call toll-free 1 (800) 695-0285
- Exceptional Parent: Parenting Your Child or Young Adult with a Disability (magazine)
- National Health Information Center, call toll-free 1 (800) 336-4797
- Internet (accessible through many “on-line” services)
Locating Training Resources

What information will you use to help your caregiver learn?

These are the resources I would like to use:

- Books
- Handouts
- Videotapes
- Audiotapes
- Computer search
- Resources I develop
- Other (please describe)

Step 4: Evaluation

You now have some goals, have planned how you will teach a caregiver, and have some resources to help you in your training. But how do you know if your training plan works? The final step in developing your training plan is evaluation.

Evaluation helps you to know how well your training goals have been met. Through evaluation you can tell how well you presented the training, what your caregivers learned, and if they can do the skills or tasks identified in your training plan.

You can evaluate in several ways. One way is to ask your caregiver to answer some questions in writing or in a discussion. You may also ask your caregiver to show you a caregiving routine or skill you have taught.
Just as it is important to include caregivers in the development of the personalized training plan, it's also important to seek and include their ideas in evaluation. Families will have some ideas for deciding when their caregivers have mastered skills that will allow them to provide respite care. But caregivers may also have some ideas about how they can show that they are ready to care for the child(ren) without supervision.

**How will you and your caregiver know when a specific skill or routine has been learned?**

I will know a specific skill or routine has been learned when we accomplish the following:

**Goal 1:** (Example) My caregiver successfully feeds Maria with no assistance.

**Goal 2:**

______________________________

______________________________

______________________________

**Goal 3:**

______________________________

______________________________

______________________________
Sample Personalized Training Plan

Child's Name: Maria  Date: May 10, 1997
Routine: Lunchtime  Time of Day: Noon - 1:15 p.m.

Step 1: Training Goal

My caregiver will be able to feed Maria lunch.

Step 2: Training Methods

My caregiver will watch me feed Maria at lunchtime.
My caregiver will assist in feeding Maria.
My caregiver will feed Maria with some assistance.
I will watch my caregiver feed Maria.

Step 3: Resources

I can give my caregiver

- a handout on feeding techniques developed by Maria's occupational therapist.
- a handout on special feeding problems in children with cerebral palsy.
- a video of lunchtime routine taped by us at home.

Step 4: Evaluation

I will ask my caregiver questions about skills and comfort in feeding Maria.
I will watch my caregiver feed Maria two bites.
I will watch my caregiver prepare and feed lunch to Maria.
Training Tips

- **Be clear with your instructions.**

  Make sure your caregiver understands what you have said. For example, if your child's care involves medical equipment such as an apnea monitor, be sure to describe what the equipment does and how it is used.

- **Be very specific.**

  Instead of telling your caregiver(s) to feed Kim, tell your caregiver to feed Kim using the purple plastic spoon. Then show the spoon to your caregiver and where it is kept.

- **Use your knowledge and resources to train.**

  Think about how you learned from others. Think about how the professionals you work with showed you how to position or feed your child or to handle different behaviors. Then adapt these methods to your style of teaching.

- **Give your caregiver written information.**

  Providing written information is another way to help caregivers learn. Share articles, diagrams, or books with your caregiver. Visit the parent resource centers in your public school district to discover new resources. Give important instructions or information in writing even if you have shared it in other ways.
Training Tips

- **Talk to your caregivers about the information you give them.**

  Your caregivers may have lots of questions about words or unfamiliar terms that you use. Check to see if your caregiver has the same understanding as you do about your child with special needs, etc.

- **Use diagrams or pictures.**

  These are often helpful to show caregivers how to use sign language or to play with your child.

- **Be sure to get your caregiver’s opinion.**

  If your caregiver did not benefit from a certain type of information, you will know to try something different.

- **Write down important facts or information.**

  Use the forms in your manual. Make extra copies of the blank forms to use with several caregivers or when information changes.

- **Share information about your culture, or life ways, that is important to your family.**

  For example, in your family it may be important for your children to play with and care for each other. In another family, children may be encouraged to be independent and play with friends in the neighborhood.
Section 8

resources
Defining Universal Precautions

Resources

The following pages contain valuable information and may be used as a resource for families and caregivers.

Universal Precautions Guidelines

In order to prevent disease, it is important to practice good personal hygiene and to take some precautions, especially when working with children. This is sometimes referred to as universal precautions.

The Occupational Safety and Health Administration (OSHA) has established methods people can use to avoid the transmission of infections and disease. By practicing the following methods, you can help yourself, your family, and the child you provide respite for stay well.

Infectious diseases can be prevented with planning and preparation. Always practice good personal hygiene and wash your hands often. Washing your hands is the most important defense to avoid contracting or transmitting an infectious disease. Good health and up-to-date immunizations are important. Always be careful and act as if all people you come into contact with are potentially infected. There is no way to determine if a person is infected just by looking at them.
In general, it is good to follow these basic precautions when you provide care:

- Wear disposable gloves when it is possible that you will contact blood or body fluids. This contact could occur directly from the person or indirectly through contact with soiled clothing, diapers, or other articles.

- Remove gloves by turning them inside out beginning at the wrist and pulling them off.

- Throw away damaged gloves.

- Do not clean or reuse disposable gloves.

- Avoid touching any items while you are wearing soiled gloves.

- Change gloves when you touch different people.

- Wear protective coverings, such as a mask, eyewear, and/or a gown, whenever you may contact body fluids that might splash.

- Cover any cuts, scrapes, or skin irritations you may have with clothing or bandages.

- Use breathing devices, such as disposable resuscitation masks and airway devices.

- Avoid needle sticks.

- Perform all procedures in such a way as to avoid splashing, spraying, splattering of body fluids.
Defining Universal Precautions

- Avoid touching your face after exposure to body fluids before you wash your hands.
- Remove soiled clothing as soon as possible.
- Clean and disinfect all areas soiled by blood or other body fluids.
- Use approximately 1/4 cup bleach to 1 gallon of water to clean and disinfect soiled surfaces and other items.
- Clean up spills immediately.
- If spills are mixed with sharp items such as broken glass, use tongs or two pieces of cardboard to pick them up, not your hands.
- Use an absorbent material to clean up a spill. Discard the used absorbent material.

If you think that you have been exposed to an infectious disease, wash the area of contact immediately. Exposure usually involves contact with potentially infected blood or fluids through a needle stick or the mucous membranes (eyes, nose, mouth) or inhaling potentially infected airborne droplets.

Infectious diseases can typically be transmitted in four ways: 1) by direct contact with an infected person, through body fluids or through contaminated needles; 2) by contact with a soiled object; 3) by inhaling air exhaled by an infected person; 4) from a bite from an animal, an insect, or an infected person.
If you are working with a child who has an infectious disease, ask the family how it is transmitted and what you should do to protect yourself. Some infectious diseases that you should be aware of are herpes, meningitis, tuberculosis, hepatitis, and HIV/AIDS. Please refer to the following chart for more information.

### How Diseases Are Transmitted

<table>
<thead>
<tr>
<th>Disease</th>
<th>Signs and Symptoms</th>
<th>Mode of Transmission</th>
<th>Infective Material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herpes</td>
<td>Lesions, general ill feeling, sore throat</td>
<td>Direct contact</td>
<td>Broken skin, mucous membranes</td>
</tr>
<tr>
<td>Meningitis</td>
<td>Respiratory illness, sore throat, nausea, vomiting</td>
<td>Airborne, direct and indirect contact</td>
<td>Food, water, mucus</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td>Weight loss, night sweats, occasional fever, general ill feeling</td>
<td>Airborne, direct and indirect contact</td>
<td>Mucus, broken skin</td>
</tr>
<tr>
<td>Hepatitis</td>
<td>Flu-like, jaundice</td>
<td>Direct and indirect contact</td>
<td>Blood, saliva, semen, feces, food, water, other products</td>
</tr>
<tr>
<td>HIV/AIDS</td>
<td>Fever, night sweats, weight loss, chronic diarrhea, severe fatigue, shortness of breath, swollen lymph nodes, lesions</td>
<td>Direct and indirect contact</td>
<td>Blood, semen, vaginal fluid</td>
</tr>
</tbody>
</table>

*Adapted from:*
Preventing the Spread of Disease

Hand Washing

The most important defense against spreading infectious diseases is thorough hand washing with soap and water. Gloves may not always be necessary or even possible to use when reacting to an emergency, but it is extremely important to wash your hands immediately afterward!

There are particular times that you should always wash your hands. It should be done frequently:

- Before putting on a smock in preparation for working with children
- Before drinking, eating, or smoking
- Before handling clean utensils or equipment
- Before and after handling children’s food
- Before and after going to the bathroom
- After contact with body secretions, such as blood, urine, feces, mucus, saliva, or drainage from wounds
- After handling soiled diapers, pads, garments, or equipment
- After caring for any child, especially those with nose, mouth, or ear discharge
- After removing disposable gloves
- After removing smock or shirt when leaving the work area
Protocol for Hand Washing

Adapted from:
California State Department of Education. (1983).

Protocol for Hand Washing

Essential Steps

1. Remove all jewelry.
2. Wet hands with warm, running water.
3. Apply liquid soap and lather well.
4. Wash hands, using a circular motion and friction, for 15 to 30 seconds.
5. Rinse hands well under warm, running water.

Key Points and Precautions

Jewelry should not be worn when working with students who require repeated physical contact and care. Microorganisms can become lodged in settings or stones of rings.

Warm water, combined with soap, makes better suds than does cold water. Hot water removes protective oils and will dry skin. Running water is necessary to carry away dirt and debris.

Liquid soap is preferred to bar soap. Bacteria grow on bar soap and in soap dishes.

Include front and back surfaces of hands, between fingers and knuckles, around nails, and the entire wrist area. Avoid harsh scrubbing to prevent skin breaks.

Hold hands under the water so that water drains from wrist area to
Protocol for Hand Washing

6. Repeat steps 3 through 5.

7. Wipe surfaces surrounding sink with a clean paper towel and discard.

8. Dry hands well with paper towels and discard towels immediately.

9. Apply lotion as desired.

Source:
References on Adult Learning


**Training Resources**

Creative Training Techniques Newsletter
Lakewood Publications
50 S. 9th St.
Minneapolis, MN 55402

(800) 707-7749

http://www.trainingsupersite.com/publications
Section 9

forms for sharing information
Information Worksheets

There are a variety of forms available in this section for families to use in sharing important information with caregivers. Families can use these forms as needed. Some forms provide information specific to the child with special needs and other forms are specific to the needs of siblings also in care. Some forms provide space to include other family and household information. Emergency, health, and safety information sheets and checklists are helpful for caregivers to have.

Master copies of all forms and worksheets are included in this section. Families can make copies of forms that several caregivers may need. Families may also keep blank copies and use them when changes occur or new information becomes available.
Final Activity
Step 1: Identify training goals.

What do you want your caregiver to learn about your child and family routines?

I want my caregiver to learn about these routines:

Goal 1: ________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

Goal 2: ________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

Goal 3: ________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

Goal 4: ________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________

Goal 5: ________________________________________________________________

_____________________________________________________________________

_____________________________________________________________________
Step 2: List training methods.

What methods will you use to teach your caregivers?

In order to reach these goals I will use the following methods:

Goal 1: ____________________________________________

Methods:

1. ____________________________________________

2. ____________________________________________

3. ____________________________________________

Goal 2: ____________________________________________

Methods:

1. ____________________________________________

2. ____________________________________________

3. ____________________________________________
Goal 3: ____________________________________________

Methods:
1. ____________________________________________
2. ____________________________________________
3. ____________________________________________

Goal 4: ____________________________________________

Methods:
1. ____________________________________________
2. ____________________________________________
3. ____________________________________________

Goal 5: ____________________________________________

Methods:
1. ____________________________________________
2. ____________________________________________
3. ____________________________________________

To Learn More about Partners Plus Contact
Child Development Resources at 757-220-1168
Step 3: Use resources to help train your caregiver.

What information will you use to help your caregiver learn?

These are the resources I would like to use:

____ books
____ handouts
____ videotapes
____ audiotapes
____ computer search
____ I develop resources
____ other (please describe)

________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
Step 4: Evaluation

How will you and your caregiver know when a specific skill or routine has been learned?

I will know a specific skill or routine has been learned when we accomplish the following:

Goal 1: ____________________________

______________________________

______________________________

Goal 2: ____________________________

______________________________

______________________________

Goal 3: ____________________________

______________________________

______________________________

Goal 4: ____________________________

______________________________

______________________________

Goal 5: ____________________________

______________________________

______________________________
Essential Information

General Information

Child's Name: ________________________  Nickname: ________________________
Address: ______________________________
Phone: ________________________  Allergies: ______________________________
Age: ____  D.O.B. ____  Height: ____  Weight: ____
Child diagnosis or description of special needs: ______________________________
______________________________
______________________________
Health Insurance: ________________  I.D. Number: ________________________
You can reach me at ________________________
If you cannot contact me, please call:
______________________________  Phone: ________________________

Medication

Name of medication: ______________________________
Purpose for medication: ______________________________
Time to be given: ________________________  Dosage: ________________________
Pharmacy: ________________________  Phone: ________________________
Comments: ______________________________
______________________________
______________________________
Essential Information (continued)

Food

Allergies:  

Cannot have:  

Acceptable snacks:  

Meals:  

Daily Schedule

AM:  

PM:  

Nap Time:
Health and Medical Information

Child's Name: ____________________ Date of Birth: ____________________

Current Health Problems / Needs: __________________________________________

Allergies (if any): ________________________________________________________

Child's Physician: ________________________________________________________

Address: ____________________________ Phone: _____________________________

Name and Number of Medical Insurance: ________________________________

Authorization for Emergency Medical Care

_________________________ has permission to obtain immediate medical care for my child if any emergency occurs when the parent or guardian cannot be located. I understand the child's physician will be contacted if available.

Parent or Guardian: ________________ Date: ________________________________
Additional information about my child:

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________
## Important Phone Numbers

<table>
<thead>
<tr>
<th>Category</th>
<th>Phone Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mother's Work</td>
<td></td>
</tr>
<tr>
<td>Father's Work</td>
<td></td>
</tr>
<tr>
<td>Police/Emergency</td>
<td></td>
</tr>
<tr>
<td>Fire/Emergency</td>
<td></td>
</tr>
<tr>
<td>Poison Control</td>
<td></td>
</tr>
<tr>
<td>Doctor's Office</td>
<td></td>
</tr>
<tr>
<td>Pharmacy</td>
<td></td>
</tr>
<tr>
<td>Hospital</td>
<td></td>
</tr>
<tr>
<td>Dentist</td>
<td></td>
</tr>
<tr>
<td>Neighbor/Friend</td>
<td></td>
</tr>
<tr>
<td>Relative</td>
<td></td>
</tr>
<tr>
<td>Relative</td>
<td></td>
</tr>
<tr>
<td>Gas Company</td>
<td></td>
</tr>
<tr>
<td>Electric Company</td>
<td></td>
</tr>
<tr>
<td>Phone Company</td>
<td></td>
</tr>
<tr>
<td>Apartment Manager</td>
<td></td>
</tr>
<tr>
<td>Veterinarian</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
</tr>
</tbody>
</table>
Other information I want you to know:

__________________________________________________________________
__________________________________________________________________
__________________________________________________________________
__________________________________________________________________
__________________________________________________________________
__________________________________________________________________
__________________________________________________________________
About My Child

Child's Name: ____________________________  Date: ________

What I would like to tell you about my child: ____________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

Favorite toys or playthings: ____________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

Special routines (a good-bye hug, snack after school, story for bedtime, etc.) my child enjoys: ____________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

Special things (blanket, a pacifier, stuffed animal, etc.) that my child enjoys: ____________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

Things that scare my child: ____________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________
Things that quiet or comfort my child:


Special things my child does:


Activities my child likes:


Activities my child dislikes:


Personality or behavior style (such as active, quiet, silly, serious):


Things that are easy for my child:


Things that are hard for my child:


Special names or words for people or things that my child understands/uses:


Other special information or concerns:
About Our Family

The people in our family are: ________________________________
______________________________
______________________________
______________________________

Some things our family does together: ________________________________
______________________________
______________________________
______________________________

Special things each member of my family does alone or with someone else:

______________________________
______________________________
______________________________
______________________________

Things we would like to do if we had respite: ________________________________
______________________________
______________________________
______________________________
About Sisters and Brothers

General Information

Brother/Sister’s Name: ___________  Nickname: ___________

Age: _______  D.O.B.: _______  Height: _____  Weight: _____

Allergies: ____________________________________________

Important health or developmental history: ____________________________________________

__________________________________________________________________________

Medications

Name of medication: ____________________________________________

Purpose for medication: ____________________________________________

Time to be given: ___________  Dosage: ___________

Pharmacy: ___________  Phone: ___________

Prescribing Doctor: ____________________________________________

Doctor’s Phone: ____________________________________________

Let me tell you about ____________________________________________

__________________________________________________________________________
Let me tell you about

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________

_________________________
Health and Safety Checklist

Our caregiver is certified in:

____ infant/child CPR
____ first aid

Our caregiver knows where we keep our:

____ medical treatment release forms
____ medical insurance information
____ first-aid kit and book
____ sunscreen

We have provided our caregiver individualized training on:

____ emergency plans
      ____ what we would consider an emergency for our child
      ____ what we would like our caregiver to do in case of an emergency
      ____ where we keep emergency phone numbers

____ fire safety
      ____ location of and how to use the fire extinguisher
      ____ our fire evacuation plan
      ____ location of smoke detector

____ poison safety
      ____ where we keep the Poison Control Center phone number
      ____ where we keep the Syrup of Ipecac

____ kitchen safety (i.e., stove, appliances, pots on the stove, etc.)
      ____ first aid for burns
We have provided our caregiver individualized training on:

- hand washing for caregivers and children after toileting and outside time and before food preparation
- medication administration
- our smoking policy
- diaper/toileting routine and diaper disposal
- toy safety (e.g., balloons might suffocate small children, young children might choke on small toys or toys with small parts)
- use of medical equipment
- crib/bed safety
  - young children should sleep on their backs
  - the side rail of a crib should be latched when the child is asleep
  - caregivers should check on sleeping children
  - bottles should be held and not propped for young children

- positioning, exercises, and therapeutic activities
- car safety
  - car seats are available and adjusted for our child
  - correct use of a car seat
  - child should always be in a car seat or fastened seat belt when the car moving

- my child's allergies
  - medical care if my child is exposed to allergens

- safety in the home
  - electrical outlets are covered
  - hot water heater adjusted to 120 degrees or lower
  - guns are unloaded and locked in a cabinet
  - bullets are locked in cabinets separate from guns
  - medications, poisonous chemicals, and liquor are kept out of the reach of children

- rubber mats or nonslip surfaces are in showers or tubs children will use
Step 1: Identify training goals.

What do you want your caregiver to learn about your child and family routines?

I want my caregiver to learn about these routines:

Goal 1: ____________________________________________

_________________________________________________

_________________________________________________

Goal 2: ____________________________________________

_________________________________________________

_________________________________________________

Goal 3: ____________________________________________

_________________________________________________

_________________________________________________

Goal 4: ____________________________________________

_________________________________________________

_________________________________________________

Goal 5: ____________________________________________

_________________________________________________

_________________________________________________
Additional Thoughts
Step 2: List training methods.

What methods will you use to teach your caregivers?
In order to reach these goals I will use the following methods:

Goal 1: ________________________________

Methods:
1. ________________________________
2. ________________________________
3. ________________________________

Goal 2: ________________________________

Methods:
1. ________________________________
2. ________________________________
3. ________________________________
Goal 3: ________________________________

Methods:
1. ________________________________
2. ________________________________
3. ________________________________

Goal 4: ________________________________

Methods:
1. ________________________________
2. ________________________________
3. ________________________________

Goal 5: ________________________________

Methods:
1. ________________________________
2. ________________________________
3. ________________________________
Step 3: Use resources to help train your caregiver.

What information will you use to help your caregiver learn?

These are the resources I would like to use:

- [ ] books
- [ ] handouts
- [ ] videotapes
- [ ] audiotapes
- [ ] computer search
- [ ] I develop resources
- [ ] other (please describe)

________________________________________

________________________________________

________________________________________
Additional Thoughts
Step 4: Evaluation

How will you and your caregiver know when a specific skill or routine has been learned?

I will know a specific skill or routine has been learned when we accomplish the following:

Goal 1: ____________________________________________

________________________________________________________________________

Goal 2: ____________________________________________

________________________________________________________________________

Goal 3: ____________________________________________

________________________________________________________________________

Goal 4: ____________________________________________

________________________________________________________________________

Goal 5: ____________________________________________

________________________________________________________________________
Additional Thoughts
Personalized Training Plan

Child’s Name: _______________________________ Date: ________

Routine: ________________________________

______________________________

Time of Day: ______________________

Step 1: Training Goal ______________________

______________________________

Step 2: Training Methods ______________________

______________________________

Step 3: Resources ______________________

______________________________

Step 4: Evaluation ______________________

______________________________
Additional Thoughts
Respite Caregiver Profile

Date: ____________

Personal:
Name: ____________________  SS#: ____________________
Address: ____________________
Home Telephone: ____________  Work Telephone: ____________
In case of emergency, notify: ____________________
Relationship: ____________________
Home Telephone: ____________  Work Telephone: ____________

Experience:
Describe experience you have had with children (including your own). ____________
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________

Have you ever provided care for children with disabilities?
__ Yes  __ No

If yes, please list your experiences:
__________________________________________________________________________
__________________________________________________________________________
__________________________________________________________________________

Partners Plus Family Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99  8/27
Respite Caregiver Profile, continued

Have you had any courses in early childhood, child development, or child health?
   ___ Yes   ___ No

Are you certified in Red Cross First Aid?
   ___ Yes   ___ No   Date of Expiration ________________

Are you certified in Infant/Child CPR?
   ___ Yes   ___ No   Date of Expiration ________________

Do you have a valid driver's license?
   ___ Yes   ___ No   Date of Expiration ________________

Education:

I have completed:

High School ______________________ Some College ______________________

College ______________________ Graduate School ______________________

Degree or Major (specify): ______________________

Other training/education: ______________________

Employment:

Current Employer: ______________________

Position: ______________________

Partners Plus Family Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Interests: (Check all that apply)

I am willing to care for children ages

birth - 2    2 - 5    5 - 8    over 8

I am willing to provide care (check all that apply):

____ at my home     ____ weekdays
____ at child's home____ weekends
____ during the day  ____ overnight
____ during evenings ____ anytime

When I provide care, I will care for:

____ only the child with disabilities
____ the child with disabilities plus siblings
____ the child with disabilities and other children in my care
____ the child with disabilities and my own child/children

I would prefer to provide care:

____ as a volunteer
____ in exchange for a service (i.e., transportation, lawn care, meals, etc.)
____ for an hourly rate of $________
____ for a negotiable hourly rate

How far are you willing to travel to provide care? (Check all that apply.)

____ Williamsburg/James City County    _____ Hampton
____ York County             _____ Poquoson
____ Newport News          _____ Gloucester
Respite Caregiver Profile, continued

References:

Please provide the names, addresses, and phone numbers of two persons we may contact who have known you for more than 1 year (excluding relatives or roommates).

Name: ___________________________ Phone: ___________________________
Address: ____________________________________________________________

Name: ___________________________ Phone: ___________________________
Address: ____________________________________________________________

I am most interested in providing respite care because _______________________

__________________________________________________________

Thank You!
Reference Check

Date: ____________

Name of Potential Caregiver: ________________________________

Name of Reference: ______________ Phone: ____________________

1. How do you know ________________________________?
   (put name of potential caregiver here.)

2. How long have you known ________________________________

3. What words best describe ________________________________

4. How would you best describe how ________________________________
   relates to children?

5. Describe what you think are ________________________________'s strong
   points in working with children with special needs and their families?

6. Are there any other comments about ________________________________
   you would like to make?
Additional Thoughts
Partners Plus

Families and Caregivers in Partnerships:
Caregiver Manual

By
Lisa L. Ownby, M.S.W.
Amanda C. Hooke, M.S.W.
Dee Wylie Moore, B.S.
Corinne W. Garland, M.Ed.

The Caregiver Manual is part of a series of resources in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care. The rest of the series includes the Community Planning, Trainer's Workshop, and Family Manuals, created by Child Development Resources.
Acknowledgments

The authors would like to thank Jayne Naughton for her technical assistance; Amy Watson for editorial assistance; families and caregivers who wrote the perspectives stories in section four of the Family and Caregiver Manuals; and our community advisory committee for multiple tasks throughout the development of training materials. Their suggestions and comments provided a significant contribution to the quality of this manual.

Amy Consiglio, of CM Consulting, designed and produced this manual.

A Word About Funding

Support for the Partners Plus project was provided under grant number HO24B40032 from the Early Education Program for Children with Disabilities, U.S. Department of Education. Points of view or opinions expressed herein do not, however, necessarily represent official views or opinions of the Department of Education.

Funding to revise the Partners Plus manuals to help families use existing community programs as respite for their school-age children with mental retardation came from the Joseph P. Kennedy, Jr. Foundation. Revisions to the manuals include practical “how to” information for personnel working with children in before and after school programs, recreational, camp, and other community programs.

Copyright © 1999. All rights reserved. Trainers may reproduce this manual, without written permission, and in whole for use only with families and caregivers. For permission to reprint single pages or to purchase additional copies of the manual, please contact:

Child Development Resources
Post Office Box 280
Norge, VA 23127-0280
Phone (757) 566-3300 / Fax (757) 566-8977

DRAFT 10.99
# Contents

**Introduction**

- The Partners Model: A New Model of Respite Care viii
- A Brief Partners History ix
- Partners Model: 5 Steps xi
- Partners Plus Replication Process xiv
- Partners Community Implementation Process xvi
- Partners Plus Manuals xvii

**Section 1—About Respite**

- Respite . . . Rest, Relax, Renew 1-2
- Families Who Need Respite 1-3
- Children Who Need Respite Care 1-4
- Benefits of Respite to Families 1-5
- Benefits of Respite to Children 1-6
- Benefits of Respite to Caregivers 1-6
- Why Don’t Families Use Respite Care? 1-7
- Keys to a Successful Respite Program 1-8
- Description of Family 1-9
- Description of Family-Centered Philosophy 1-9
- Family-Centered Care 1-10
- A New Respite Model: Partners 1-11
- Partners Model of Respite Care 1-12
## Contents, continued

### Section 2—Building Partnerships: Communication

- Building Partnerships: Communication 2-2
- How Do You Feel? 2-4
- Six Messages of Communication 2-5
- What You Need to Know About Communication 2-5
- Methods of Communication 2-8
- Sharing Information 2-9
- Tips for Communicating with Your Caregiver 2-10

### Section 3—Building Partnerships: Working Together

- Building Partnerships: Working Together 3-2
- "Welcome to Holland" 3-3
- A Personal Story 3-5
- Being a Father of a Child with Special Needs 3-7
- A Parent's Perspective 3-8
- Malachi 3-10
- A Sibling Perspective 3-13
- One Caregiver's Perspective 3-17
- Stress 3-20
- Coping with Stress 3-21
- Strategies for Dealing with Stress 3-22
- The Challenges of Respite 3-23
- Meeting the Challenge 3-24
- Successful Family and Caregiver Partnerships 3-26

### Section 4—All Kids Like Cookies

- All Kids Like Cookies Activity Sheet 4-2
- Visual Impairment 4-3
- Speech and Language Impairment 4-4
- Physical Disability 4-5
## Contents, continued

### Section 5–Disabilities Awareness

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Understanding Disabilities</td>
<td>5-2</td>
</tr>
<tr>
<td>Finding Educational Services</td>
<td>5-3</td>
</tr>
<tr>
<td>Understanding Early Intervention</td>
<td>5-5</td>
</tr>
<tr>
<td>Our Family Journey in Early Intervention</td>
<td>5-7</td>
</tr>
<tr>
<td>Jack's Story</td>
<td>5-8</td>
</tr>
<tr>
<td>A Parent's Thoughts on Special Education</td>
<td>5-10</td>
</tr>
<tr>
<td>Common Issues for Families</td>
<td>5-11</td>
</tr>
<tr>
<td>A Parent's Perspective on Transition</td>
<td>5-12</td>
</tr>
<tr>
<td>Accessibility</td>
<td>5-15</td>
</tr>
<tr>
<td>Language</td>
<td>5-17</td>
</tr>
<tr>
<td>How Parents Describe Their Child</td>
<td>5-18</td>
</tr>
<tr>
<td>What Words Activity</td>
<td>5-20</td>
</tr>
<tr>
<td>Alphabet Soup</td>
<td>5-21</td>
</tr>
<tr>
<td>Summary</td>
<td>5-22</td>
</tr>
</tbody>
</table>

### Section 6–Understanding Child Development

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Understanding Child Development</td>
<td>6-2</td>
</tr>
<tr>
<td>Areas of Development</td>
<td>6-4</td>
</tr>
<tr>
<td>Basics of Child Development</td>
<td>6-6</td>
</tr>
<tr>
<td>The Importance of Play</td>
<td>6-10</td>
</tr>
</tbody>
</table>

### Section 7–Children with Special Needs

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Understand Each Child's Needs</td>
<td>7-2</td>
</tr>
<tr>
<td>How Special Needs Affect Development</td>
<td>7-18</td>
</tr>
<tr>
<td>High-Risk Signs</td>
<td>7-21</td>
</tr>
<tr>
<td>Where to Find More Information on Special Needs</td>
<td>7-24</td>
</tr>
</tbody>
</table>
## Contents, continued

### Section 8—Becoming a Caregiver
- Congratulations 8-2
- What Can I Contribute As a Respite Caregiver? 8-2
- Getting Ready for Training 8-3
- Sample Interview Questions 8-4
- Personalized Training 8-5
- Developing a Personalized Training Plan 8-6
- Sample Personalized Training Plan 8-7
- How I Learn Best 8-8
- Tips for Learning 8-9
- Essential Information for Caregivers 8-11

### Section 9—Resources
- Defining Universal Precautions 9-2
- Hand Washing 9-6
- References on Adult Learning 9-9
- Training Resources 9-10
- ARCH: Respite Care
- ARCH: A Parent's Perspective
- NICHCY: National Toll-Free Numbers
- NICHCY: National Resources
- NICHCY: General Information About Disabilities

### Section 10—Forms for Sharing Information
- Information Worksheets 10-2
- Essential Information 10-3
- Health and Medical Information 10-5
- Important Phone Numbers 10-7
- About My Child 10-9
- About Our Family 10-11
- About Sisters and Brothers 10-13
- Health and Safety Checklist 10-15
Contents, continued

Section 10-Forms for Sharing Information, continued

<table>
<thead>
<tr>
<th>Step</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step 1: Identify training goals</td>
<td>10-17</td>
</tr>
<tr>
<td>Step 2: List training methods</td>
<td>10-19</td>
</tr>
<tr>
<td>Step 3: Use resources to help train your caregiver</td>
<td>10-21</td>
</tr>
<tr>
<td>Step 4: Evaluation</td>
<td>10-23</td>
</tr>
<tr>
<td>Personalized Training Plan</td>
<td>10-25</td>
</tr>
<tr>
<td>Respite Caregiver Profile</td>
<td>10-27</td>
</tr>
<tr>
<td>Reference Check Form</td>
<td>10-31</td>
</tr>
</tbody>
</table>

Final Activity

- Step 1: Identify training goals
- Step 2: List training methods
- Step 3: Use resources to help train your caregiver
- Step 4: Evaluation
Notes on this Section
Caregiver Manual

introduction
The Partners Model of Respite Care...

*Helps families of children with special needs find and train their own respite caregivers.*

*Through training and support, families and caregivers build long-lasting, successful respite relationships.*

**Introduction**

Like all families, families of children with special needs want and need time away from the demands of parenting. *Respite—or temporary child care for children with disabilities or chronic illness—*gives families time to enjoy an evening out by themselves, with other family members, or with friends. However, for these families, finding respite care is not as simple as calling a neighborhood babysitter. These families need caregivers who understand their children's special needs.

When the staff at Child Development Resources (CDR) spoke with families about what they wanted in a respite program, the message was clear. Families wanted to choose and train their own respite caregivers. They wanted affordable care that was as simple to schedule as calling the neighborhood babysitter. And, families wanted caregivers who were trained to provide care.
In response to families and with support from the Virginia Institute for Developmental Disabilities (VIDD), a University Affiliated Program at Virginia Commonwealth University in Richmond, Virginia, CDR developed a pilot project that was low-cost, family-directed respite care called PARTners: Parents As Respite Trainers.

CDR was awarded, in 1994, a three-year grant from the U.S. Department of Education, Office of Special Education Programs (OSEP), Early Education Program for Children with Disabilities (EEPCD), to build on the pilot and to develop a model of respite that could be replicated in other communities. The project, Partners Plus: Families and Caregivers in Partnerships, provides a model for families to use in finding and training respite caregivers. This model encourages families to use available, natural family and community supports to meet their respite needs.

Within this model, Partners broadly defines the terms respite and caregiver. For example, respite can take place in a family’s home, a caregiver’s home, or in before and after school, recreation, camp, or other children’s program. A caregiver can be any individual who provides respite care to a child with special needs or includes a child with special needs within an existing community program.

Partners Plus provided training to more than 200 families and caregivers in a five-city, three-county area of eastern Virginia. Partners usually conducted the six hour Partners Plus workshop for families and caregivers in one day. Sometimes when it was more convenient for a group, Partners conducted the workshop in two parts on separate days. This flexibility ensured that Partners adapted training
to fit the schedules of families and caregivers who are generally most receptive to training during nontraditional hours such as evenings and Saturdays. Families attending the workshop learned to provide their own caregivers with personalized training based on their children's daily routines. Families either came to training with their own caregivers, met caregivers through the workshop, or through other project activities.

The Partners Project resulted in the development of four manuals designed to help communities, trainers, families, and caregivers use the Partners model. The Caregiver Manual is one of four in a series of manuals that Partners has conveniently packaged together in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care. Besides the set of manuals contained in the guide, you can purchase the Family and Caregiver Manuals separately from the Partners Plus program through CDR.

Child Development Resources
Post Office Box 280
Norge, VA 23127-0280
Phone (757) 566-3300
Fax (757) 566-8977
Email ppcdr@tni.net
The Partners Model: A New Model of Respite Care

Partners programs provide group training and continuing support to families and potential caregivers. The Partners model contains five steps:

The Five Steps of the Partners Model

1. Recruit Families and Caregivers
2. Conduct Partners Workshop
3. Help Families Choose Caregivers
4. Guide Families as they Train their Own Caregivers
5. Provide Continuing Support

Step 1: Recruit Families

First, programs recruit families and potential caregivers to participate in Partners. Community awareness activities invite potential participants to a Partners workshop. Information about the workshop and the Partners program is distributed through parent groups, newspapers, radio, flyers, word of mouth, etc.
Step 2: Conduct Partners Workshop

Next, programs conduct the Partners workshop. During the six-hour workshop, families learn how to find, screen, and interview caregivers for their children. In addition, they learn to structure training for their own caregivers based on the specific needs and daily routines of their individual children. Caregivers learn basic information about caring for children with special needs. During the workshop, caregivers are preparing to be active in their work with families, reflecting on their motivations for becoming respite caregivers and identifying their own learning needs for personalized training. Partners Family and Caregiver Manuals provide participants with useful resources to actively engage in during the workshop and to use as resources as they move through the next steps of the model.

Step 3: Families Choose Caregivers

After the workshop, Partners programs provide continuing support as families choose caregivers. Families frequently identify respite caregivers within their own circles of natural support. If families are unable to identify their own caregivers, Partners will assist by providing social and educational events that will help families and caregivers meet. During interviews, families and caregivers negotiate how respite occurs. Respite may be in-home or out-of-home and reimbursement for care can be flexible. Care may be provided for a fee, voluntarily, or in exchange for another service, such as yard work.
Step 4: Families Train Caregivers

Once families and caregivers decide to work together, families train their own caregivers. This “personalized training,” discussed during the workshop, is organized by families with help from Partners staff if needed. Families decide what routines caregivers need to learn and work with caregivers to determine how training will occur, thus forming true partnerships necessary for successful respite.

Step 5: Provide Continuing Support

As families and caregivers work together to develop a personalized respite training, the community Partners program offers continuing support, information and guidance. The Partners program is available to support families and caregivers in various ways as they begin their respite relationship. Partners staff members respond to requests for help and support from both families and caregivers. For example, Partners may help families and caregivers with screening, interviewing, developing personalized training plans, and providing educational opportunities.
Partners Plus Replication Process

The Partners Plus Community Planning Manual, contained in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care, helps communities use the Partners model of respite care. Each community interested in model replication develops a plan for collaboration among agencies, organizations, and individuals in order to implement the model. Community resources and supports are needed to sustain the model, to create a home for the program, and to ensure continuation.

The community replication process includes four major steps:

1. Identifying the site for replication
2. Planning for replication
3. Implementing the model
4. Evaluating the model


Your community will be a good candidate for replication if you have the following:

- support from agencies that provide, or are involved with, respite services
Partners Replication Process

- knowledge of current respite funding to support families without financial resources
- support and commitment from local early intervention programs, mental health or mental retardation programs, and/or family support programs

Each community needs to develop a plan for replication. Awareness activities aimed at a broad array of agencies, organizations, and individuals help strengthen community support. The creation of an advisory committee brings expertise and resources that will ensure a quality program. The replication plan includes finding a home for the program and securing staff responsible for implementing the program. Formal or informal interagency agreements and perhaps grant proposals may be necessary for community replication.

The Community Planning Manual provides strategies and materials to help communities develop and implement the respite care program. The Trainer's Workshop Manual, accompanying the Community Planning Manual, includes agendas, overheads, and suggestions for what to say during the Partners workshop. There are also suggestions for the kinds of resources that are needed by families and caregivers in order to foster and maintain long-term relationships.

Communities that replicate the Partners model gather information from families and caregivers to evaluate the quality and usefulness of training and support.
Partners in your Community

Partners Community Implementation Process

Identify Community Site

Plan for Replication of the Partners Model
- Create Community Support
- Determine Program Advisors
- Determine Administrative Structure
- Secure Financial Resources

Implement the Partners Model
- Recruit Families and Caregivers
- Conduct Partners Workshop
- Help Families Choose Caregivers
- Support Families As They Train Their Own Caregivers
- Provide Continuing Support

Evaluate the Partners Program

Partners Plus Caregiver Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

867 DRAFT 10.99
Partners Plus Manuals

There are four manuals in the Partners Plus Families and Caregivers in Partnerships, A Family-Centered Guide to Respite Care that support the Partners Plus replication process:

1. The Family Manual
2. The Caregiver Manual
3. The Community Planning Manual, and

The Family and Caregiver Manuals are used in the Partners workshop for families and caregivers. These manuals include resources that families and caregivers can continue to use as they develop partnerships for respite care for children with special needs. The Community Planning Manual helps communities develop a community-based, family-centered respite system. The Trainer's Workshop Manual contains overheads, forms, and "helpful hints" for conducting Partners workshops.

The Family Manual is easy to follow and guides families in choosing and training their own respite caregivers. Families use their children's daily routines to develop training plans for caregivers. The manual includes information about respite, communication, building partnerships, selecting caregivers, and developing personalized training. There are worksheets, resources, and forms that can be duplicated for future use.
The Caregiver Manual provides caregivers with basic information on caring for young children with disabilities and supporting families. This manual includes information about respite, communication, building partnerships, disabilities awareness, child development, children with special needs, and personalized training with families.

The Community Planning Manual provides communities with a step-by-step guide to replication of the Partners model. By following each step, communities and families can work together to develop respite care programs. The manual contains forms and materials to assist with replication of the model.

The Trainer’s Workshop Manual, used in conjunction with the Community Planning Manual, helps community trainers to conduct the six-hour Partners workshop for families and caregivers.
Section 1

About respite
About Respite Care

Respite . . . Rest, Relax, Renew

Often we are asked, What is respite care? Our first response is simple: respite care is temporary care for a family member with special needs.

Families have told us in many different ways what respite means to them:

- Time to readjust.
- A break from the routine.
- Rest in spite of demanding circumstances.
- I feel refreshed and renewed when I come back from respite.
- Think of a candle that slowly burns: the flame flickers and goes out, and, just as a candle, we need to be relit, too.

— Partners Plus Families

More About Respite Care

Respite care is a term used to describe occasional care for an individual with special needs. Respite care programs began in the 1960s when people with disabilities began receiving care in their homes and communities instead of in institutions. Acknowledging the importance of respite care to families providing home care, Congress passed federal legislation providing funds to help communities develop respite care programs.
The 1986 federal legislation that provided funds to set up model programs of respite care defines respite care as in-home or out-of-home temporary, nonmedical care for families who have children (young and/or adults) with disabilities or chronic or terminal illnesses. More simply, respite care is any type of temporary care provided to a person with a disability or special health care need.

There are a variety of respite models, each designed to meet the unique needs of families. Descriptions of these models are included in Section 9 of this manual (see ARCH Fact Sheet Number 2).

**Families Who Need Respite**

Families who have children with special needs are very much like other families. Like all families, they have values, beliefs, interests, concerns, needs, and dreams. However, families of children with special needs also have special challenges that affect the way they live. Here are some things we have learned from families.

- Like other parents, parents of children with special needs are concerned about their children's development, health, education, training, and social life.
- Parents of children with special needs must put forth extra effort to help their children.
Who Needs Respite?

- Parents of children with special needs are often very knowledgeable about their children's disabilities and have learned some of the best ways to help them.
- Like other parents, parents of children with special needs are concerned about:
  - the needs of all of their family members,
  - the education of all of their children,
  - the activities of all of their family members, and
  - their jobs.
- Daily routines for families of children with special needs are similar to those of other families.
- Families of children with special needs have hopes and dreams that are like those of most other families.
- Children with special needs and their brothers and sisters can grow up healthy and lead productive lives.

Children Who Need Respite Care

A child who needs a respite caregiver is one who has special physical, emotional, or health needs that require care beyond that necessary for other children. The kinds of special needs or disabilities that a child may have can vary greatly from hearing impairments that require a child to communicate through sign language to mental retardation to terminal illness.
There is no one approach to caring for children with special needs but there are some basic guidelines:

- Children with special needs are more like other children than they are different.
- Children with special needs can and do learn.
- Children with special needs should be encouraged to do as much for themselves as they can.
- Children with special needs thrive when given love, acceptance, and necessary support.
- Like other children, children with special needs benefit from healthy play and recreation, friendships, being outdoors, and going places.

Benefits of Respite to Families

Respite care gives parents a break from the daily demands of parenting a child with special needs, and offers much more.

- Rest to recuperate, rejuvenate, replenish, rejoice
- Enjoying movies, dinner, fishing
- Shopping for groceries, school, self, kids, family
- Pampering at a hair salon or gymnasium or having lunch with a friend
- Interests to pursue such as hobbies, sports, community activities, and education
- Time to spend with siblings, other family members, and friends
- Everything or anything families want to do!
Benefits of Respite Care

Benefits of Respite to Children

Respite is also beneficial to children. Some of these benefits include the following:

- Relief that their parents finally got out
- Exploring new faces and activities
- Special friends of their own
- Parents who feel rejuvenated
- Interacting with other adults and children
- Time to be independent from their family
- Educating each other about differences and similarities

Benefits of Respite to Caregivers

Respite also benefits caregivers in many ways.

- Really great children and families to get to know
- Especially rewarding to know that they are making a positive difference for a family
- Self-improvement and growth through learning new skills and ideas
- Positive attitudes about themselves and about people who are different from themselves
- Involvement with children, families, and communities
- Thoroughly appreciated by families who need them
- Everything caregivers want respite to be!
Who Can Be Respite Caregivers?

- Neighbors
- Friends
- Relatives
- Community Program Staff
- Church Members
- Students
- Bus Aides
- Anyone with a desire

Why Don’t Families Use Respite Care?

Despite the benefits of respite for everyone, many families do not use respite care. Families have identified various reasons:

- It is difficult to find someone to provide temporary care.
- We are afraid to leave our children with strangers.
- We fear that respite caregivers lack adequate training.
- We sometimes feel guilty about leaving our children.
- It is difficult to find community programs for respite care.
- It is too disappointing when caregivers cancel frequently.
Keys to Successful Respite

Keys to a Successful Respite Program

Families say a successful respite program must have these elements:

- Helps families feel comfortable
- Gives families options for how they want to participate
- Gives families options for times to be involved
- Is family centered
- Supports families in developing new linkages
- Supports families in developing individualized caregiver training
- Asks families if they want to be involved in program planning
- Gives families access to dependable, reliable caregivers

Source:
Reprinted with permission of ARCH National Respite Network and Resource Center, 800 Eastowne Drive, Suite 105, Chapel Hill, NC 27514:
Description of Family

What Do We Mean by Family? We all come from families. Families are big, small, extended, nuclear, multigenerational, with one parent, two parents, and grandparents. We live under one roof or many. A family can be as temporary as a few weeks, or as permanent as forever. We become part of a family by birth, adoption, marriage, or from desire for mutual support. As family members, we nurture, protect, and influence each other. Families have strengths that flow from individual members and from the family as a whole. Each family is a culture unto itself, with unique values and its own way of realizing dreams. Together, our families are the source of our rich cultural heritage and spiritual diversity. Our families create neighborhoods, communities, states, and nations.

Source:

Description of Family-Centered Philosophy

Family centered means the recognition that the family is the constant in a child’s life and that service systems and the personnel must support, respect, encourage, and enhance the strengths and competence of the family.

Source:
Family-Centered Care

What does providing family-centered care mean?

- Recognizing that a family will always be in a child's life while services and professionals will change
- Making it easier for families, volunteers, and professionals to work together in hospitals, homes, and community care
- Sharing information, ideas, and concerns among families, volunteers, and professionals as a team
- Honoring the diversity of families, knowing that we each bring richness to the world we live in through our cultures and the ways we live our lives
- Recognizing and respecting that families meet their own needs in different ways
- Designing services that are flexible, that are easy for families to use, and that meet each family's individual needs
- Appreciating families as families and children as children and recognizing that they have a wide range of strengths, concerns, emotions, and dreams beyond their needs for specialized health and developmental services and support

Source:
Reprinted with permission of the Association for the Care of Children's Health, 19 Mantua Road, Mt. Royal, NJ 08061:
A New Respite Model: Partners

The Partners model of respite care is designed to increase respite options for families of children with special needs by helping families find and train their own respite caregivers.

Respite can be provided in:

- Families' homes
- Caregivers' homes
- Community respite programs
- Before and after school programs
- Recreation programs
- Camps

Respite can take place anytime, any place, with anyone in order to support families in meeting their needs.

Partners is a natural and family-centered model that involves families in the design, implementation, and evaluation of respite care. The model encourages the development of community collaboration among agencies and parent, religious, and/or support groups to support this model and to create a home for the program.

Together, families and caregivers attend group training. Families learn how to recruit, screen, interview, and train caregivers. Caregivers are provided with basic education in caring for children with special needs. After group training, the next step is for families and caregivers to match up and begin individualized training. The Family and Caregiver Manuals are provided to families and caregivers to use as a resource to help them through this process.

Partners... is a family-centered model.

- offers families many choices.
- allows families options for how they can participate.
- supports families in finding caregivers.
- supports families in training their own caregivers.
- helps families build necessary skills to communicate respite needs.
- helps families build long-term caregiver relationships.
- expands respite options for families.
Partners Respite Model

The Partners program provides support to families and caregivers to help them begin building respite relationships. When a family cannot find a respite caregiver within their own circle of natural supports, they can choose from the pool of caregivers recruited by Partners Plus or can attend social and educational events coordinated by the program to help families and caregivers meet and match up.

With the Partners model, families and caregivers have choices about how respite care is provided. Respite may be provided in-home or out-of-home. Care may be provided for a fee, in exchange for another service, or by a volunteer. These decisions are made between families and caregivers. Families may also choose to develop a parent (or family) cooperative in which they care for each other's children. Families decide what kind of training caregivers need and work with caregivers to determine how training will proceed, forming a true partnership necessary for successful respite.

Partners Model of Respite Care

**Step 1**  Recruit families and caregivers

**Step 2**  Conduct Partners workshop

**Step 3**  Support families in choosing caregivers

**Step 4**  Guide families as they train their own caregivers

**Step 5**  Provide continuing support
Section 2

building partnerships: communication
Building Partnerships: Communication

Building Partnerships

*Respite care is not just about giving out medicine. It is a whole dialogue and relationship with a family.*

— Partners Plus Caregiver

Partners Plus supports strong relationships between families who have children with special needs and their caregivers. In partnerships, families and caregivers respect one another and commit to working together.

Partnerships actively prevent caregiver burnout and family disappointment. Partnerships are built with open and honest communication and understanding. Sections 2 and 3 of this manual explore building partnerships through communication and collaboration.

Building Partnerships Through Communication

Good communication strengthens the relationship between family and caregiver.

*Honesty and openness in communication help build cooperative relationships over time!*

What does communication mean?

- Sharing information, ideas, and feelings that increase understanding about yourself and others
- Listening carefully to each other in ways that develop partnerships and promote the best care for children
- Sharing common values and traditions and talking about similar experiences to strengthen your relationship with each other
- Appreciating differences to strengthen mutual respect
Building Partnerships: Communication

Clear communication is one key ingredient in building partnerships. We send messages all the time, even when we are not speaking.

Communication is . . . 7% Words

Research shows the words we use or choose are important in gaining understanding. In communicating with another person, we need to use language that is common to both the speaker and the listener.

38% Voice

The voice we hear is more important than the words we use. We need to be aware of our expression and tone of voice. We use our voices to help express things like sincerity, conviction, requests, and concern.

55% Facial and Body Language

Our face and body are the most powerful means of communication. Smiles, frowns, raised eyebrows, widened eyes, tears, laughter, scrunched-up faces, and wrinkled noses all tell us very different things.

Source:
Facial expressions can communicate a lot about our thoughts and feelings. Use the blank circles to draw faces that send messages to others.

Anxious  Angry  Bashful  Bored
Confident  Thinking  Confused  Scared
Innocent  Relieved

Adapted from feelings posters by:
How Do You Feel Today? Productions, Box 1085, Agoura Hills, CA 91376. 818-706-2288. E-mail: p&p@howdoyoufeel today.com. Web site: www.howdoyoufeel today.com. International Copyright MCMLXXIX A.K. Graca © All rights reserved.
Six Messages of Communication

Communication is a cycle. As you communicate with others, think about the many places that communication can break down. Does the other person really understand what you mean? Each communication contains the possibility of six separate messages:

- What you mean to say
- What you actually say
- What the other person hears
- What the other person thinks he hears
- What the other person says
- What you think the other person says

Adapted from:

What You Need to Know About Communication

- Communication is the way we send and receive messages.

Most people use a combination of verbal and nonverbal communication. Being aware of and sensitive to the different ways people communicate helps to build mutual respect and understanding between families and caregivers.
Our ways of communicating are influenced by our culture. Our culture determines much about how we think, feel, act, and communicate with others.

The term *life ways* refers to the many ways we are influenced by others and by our environment throughout our lives.

Gestures, silence, body movements, distance, and emotional expressiveness are other ways of communicating that are influenced by our cultures and life ways.

For Example:
- Too much silence makes some people uncomfortable.
- Too few pauses while talking may disturb others.
- To be comfortable, some people need some distance between themselves and others.
- Lots of eye contact is important in establishing trust for some.
- For others, too much eye contact is considered disrespectful.
- A relaxed open body posture, with arms not crossed in front, invites others to talk.
- Leaning slightly forward shows interest.
- Sitting up straight helps you look more alert.
- Much of our communication is done without words.
- Our eyes are used to communicate.
For Example:
A look can be exchanged between parents that says, *It's time to leave*, or a look from a mother's eyes to her child can communicate the message, *You need to behave*.

- When people have similar experiences, we sometimes presume a shared understanding of feelings or thoughts.

For Example:
When a parent at an oncology clinic says to another parent in the waiting area, *My child is receiving chemotherapy for leukemia too*, an exchange that takes place by looking at each other in the eyes can send a message of shared understanding, even though each family's experience is unique.

Depending on their life ways, families use more spoken or written words to communicate. Words help us to be direct, exact, and logical and encourage others to say more. The following is an example of a family and caregiver interview in which words are used to encourage a response.

**Caregiver:** I want to provide respite because I enjoy being with children.

**Family:** That's great! You like being with children.
Methods of Communication

Caregiver: Yes, I miss being a parent and my grandchildren live in another state. When my children were young, I enjoyed planning activities for them.

Family: What type of activities do you like to plan for children?

- Spoken and written words are helpful in finding out about a person's feelings and attitudes.

Questions that cannot be answered with a yes or no encourage others to talk about what is on their mind. Use words like how, why, what, and tell me.

For Example:

Family: On your first visit to my home you watched me feed Crystal and I gave you some information to take home to read. Why don't you try feeding Crystal today?

Caregiver: I think I should watch again to make sure I remember how.

Family: You seem uncomfortable trying this now. Tell me about your concerns.

Caregiver: I remember how much Crystal coughed when you fed her and how much she gagged. I am afraid I will choke her. I think you need someone with nursing skills to take care of your child.
Family: Perhaps we need to talk again about why Crystal has difficulty eating and why she might cough or gag. I want you to understand Crystal’s special needs and to feel comfortable in caring for her. I am here to help while you feed her. Let me show you again. Then you try.

- Using words like describe, show me, and tell me can help determine if a caregiver has learned a skill or has understood the instructions.

For Example:

Family: How did your first day go?

Caregiver: I had trouble using Jared’s wheelchair when we went outside.

Family: Tell me what gave you trouble.

Sharing Information

To ensure a successful respite relationship, families and caregivers need to openly share information. This is necessary in initial training and throughout the relationship.

Information can be shared in many ways:
- Keep and share a respite journal.
- Plan information you want to share when you get together.
Tips for Caregiver Interaction

- Allow time to meet with each other before respite care takes place.
- Allow time to talk after respite care.
- Make necessary decisions promptly.
- Resolve conflicts in a timely way.
- Solve problems together.

Tips for Communicating with Your Caregiver

- Communication is essential and is an important part of maintaining a successful relationship.
- Give clear instructions and make sure your caregiver understands what you have said.
- Provide immediate feedback to your caregiver during all training activities.
- Put yourself in the caregiver’s place; remember, you once had to learn new ways to care for your child.
- Let your caregiver know when respite care goes well and when you are pleased.
- Be positive. When changes are necessary, tell your caregiver in a positive way.
Section 3

building partnerships:
working together
Partnerships: Working Together

Anyone with a desire to help families of children with special needs can become a capable respite caregiver. At first, many parents do not know how to care for their children with special needs, but gradually they learn. Through willingness to learn from families, respite caregivers can do the same.

Many respite caregivers, although not parents of children with special needs, can appreciate the benefits of getting a break from caring for children. Some people have life experiences that help them become more comfortable as respite caregivers. They may know someone who has special needs—a friend, a student, a classmate, someone at church, or a family member. The most important quality a respite caregiver has is a positive attitude.

We know children and their families depend on each other. Children depend on their families for love, support, and guidance. The family is a continuing influence in their child's life, whereas friends, neighbors, and caregivers change. Parents know their own children best. Families and caregivers need to know how to work together for successful respite care.

While caring for children with special needs can be a wonderful experience for both families and caregivers, it can also be stressful. It is helpful for families to understand the challenges that caregivers face when caring for children with special needs. Caregivers may feel unimportant or underappreciated. They may be uncomfortable with the tasks they are given. The material on the following pages can help you learn from each other.
Family and Caregiver Perspectives

Families and caregivers can better understand each other by listening to personal experiences. Several perspectives from families and caregivers are included in this section.

The first story is "Welcome to Holland" by Emily Pearl Kingsley. From that story, you will see how one parent, whose child was born with a disability, saw her life changing. Other stories from families and caregivers are also included. Your story may look very different from these. To get a better understanding of one another we encourage families and caregivers in partnerships to share their unique stories.

"Welcome to Holland"

I am often asked to describe the experience of raising a child with a disability—to try to help people who have not shared that unique experience to understand it, to imagine how it would feel. It's like this . . .

When you're going to have a baby, it's like planning a fabulous vacation trip—to Italy. You buy a bunch of guidebooks and make your wonderful plans—the Coliseum, the Michelangelo David, the gondolas in Venice. You may learn some handy phrases in Italian. It's all exciting.

After months of eager anticipation, the day finally arrives. You pack your bags and off you go. Several hours later the plane lands. The stewardess comes in and says, "Welcome to Holland!"
"Welcome to Holland"

"Holland?!" you say. "What do you mean, Holland? I signed up for Italy! I’m supposed to be in Italy. All my life I’ve dreamed of going to Italy!"

But there’s been a change in the flight plan. They’ve landed in Holland and there you must stay.

The important thing is that they haven’t taken you to a horrible, disgusting, filthy place, full of pestilence, famine, and disease. It’s just a different place.

So you must go out and buy new guidebooks. And you must learn a whole new language. And you will meet a whole new group of people that you would have never met.

It’s just a different place. It’s slower paced than Italy, less flashy than Italy. But after you’ve been there for awhile you look around and you begin to notice Holland has windmills, Holland has tulips, Holland even has Rembrandts.

But everyone you know is busy coming and going from Italy, and they’re bragging about what a wonderful time they had there. And for the rest of your life, you will say, “Yes, that’s where I was supposed to go. That’s what I had planned.” The pain of that will never, ever, go away because the loss of that dream is a very significant loss.

But if you spend your life mourning the fact that you didn’t go to Italy, you may never be free to enjoy the very special, the very lovely things in Holland.

— Emily Perl Kingsley, 1981, Parent
A Personal Story

Although my son Matthew died, he is still a part of my life. Matthew made a difference in who I have become. At first I shared the same thoughts and feelings that Ms. Kingsley expresses in “Welcome to Holland.” After Matthew was born, my family began to live a different kind of life. It was often very difficult for us. We learned a lot about ourselves and we learned a lot from Matthew. We discovered that our love for Matthew and each other was very powerful. We found that we had many hidden strengths. Our family grew stronger ties with our community. Friends, neighbors, professionals, and even strangers helped us in many different ways. There were two very important things we wanted most from others. One was for others to accept our son. The other was respect for our family’s decisions in how to live with our child’s chronic illness and disabilities.

After getting to know Matthew, his disabilities did not seem like the most important part of his life. My four-year old daughter taught me to love Matthew just because he was a baby. To my daughter, Matthew was always her baby brother, not a baby with disabilities. She never noticed much difference until she got older. Other families of children with disabilities have shared some similar thoughts with me. Many say they wish others would stop seeing only their child’s wheelchair, their tubes, the way they talk differently or look different. At times, families want professionals to stop testing their
child long enough to get to know him or her as a child, a child with a name and family.

When I describe my son, I tell people that Matthew was small and easy to carry places. Daily care for Matthew was like taking care of an infant. He had a crooked little smile. His sister could make him smile most often, but others could sometimes if they gave him a gentle squeeze around his shoulders. His favorite game was to repeat sounds that we made to him. Matthew’s favorite toy was a baby’s beaded teething ring. He liked to fall asleep on his father’s shoulder. Matthew’s greatest strength was that he could teach others about the power of the human spirit.

In describing Matthew’s disabilities I would say that Matthew had a genetic impairment called trisomy 22. Trisomy 22 is an extra chromosome that affects every part of the body. He never grew to be more than five pounds and twenty-three inches. There was a hole in the roof of Matthew’s mouth that made sucking and feeding very difficult. We fed Matthew with a tube that we put down his throat and into his stomach. His heart and all other systems in his body did not work well. Matthew was delayed in all areas of his development. With help from professionals we learned to deal with all of his special needs.

Getting information about our son’s disability through an early intervention program and from medical practitioners helped us understand how to care for our child. A service that helped our family a
lot was respite care. My family and I needed time separate from being with Matthew. I wanted to give all my attention to my daughter at her soccer games. My husband and I needed time together to laugh and have fun. I needed time to rest physically from the frequent round the clock feedings required for Matthew. I needed emotional rest from the constant concerns I had for Matthew. A few friends, neighbors, and community agencies helped us get respite from time to time. One agency linked us with a community family who was willing to give me a few hours of respite each week. Just those few hours gave me the relief I needed to renew my energy and continue to care for Matthew.

— Dee Moore, 1995, Parent

Being a Father of a Child with Special Needs

Being a father of a special needs child takes a special person. There are times when you just want to give up but you know that you can’t. The doctors gave my son 3 days to live and now my son is 3 years old. It’s a blessing from God that he’s still with me today. One thing that I try to remember is to treat him like a normal child. I take him to amusement parks on vacation and to the mall, just to name a few places.

There are times when it’s very difficult being a father of a special needs child. I have to watch my
child go through intensive surgeries just to be able to eat and breathe like you or I. There are times when all you can do is just listen to your child cry uncontrollably, but overall being the father of a special needs child is a blessing from God. If you're a dad and wonder what it's like to be a dad to a special needs child, it is like taking care of three children.

— Lamar Ramsey, 1997, Parent

A Parent’s Perspective

From the moment Kieren was born, I knew something wasn’t quite right. He used to cry all the time. He was totally fearless, and I could leave him with anyone. He didn’t seem to care if I was there or not. As he grew older, he used to run away all the time. We had to lock all the doors and never let go of his hand when we were out or he was gone and he didn’t come back. If he saw something he liked, he would jump up and down, flap his hands, and shriek. Looking back, there were a lot of warning signs but I didn’t know what I was looking for. I just knew that something was not right. I think one of the worst things about not knowing he had high-functioning autism was that no one else seemed to think there was a problem (or so they led me to believe). I would say to family and friends that I was concerned, and they would tell me, “Oh, all children do that,” and dismiss it, and in the end I was beginning to doubt my sanity.
When he was 2, we were stationed in Germany, where 9 months later my daughter was born. I think it was a blessing in disguise that we were in that country as they test routinely for cystic fibrosis in all newborns. We found out when she was 6 weeks old that she had it. She was having some problems but the doctors told me this is normal, all babies do these things. Where have I heard that before?!

So we were sent back to the states so that we could get the health care she needed. Once she was relatively stable, I turned my attention back to Kieren. I made an appointment with the school division to get Kieren assessed to see if he was eligible for the special education program. During all the appointments I remember thinking he is doing really well today and they are going to think I'm crazy. As it turned out, he was eligible with significant delays in language, cognitive and fine motor skills.

He started the special education program, and things went relatively well for the first 6 months as Kieren's language started to emerge and he became a little more in control of his actions. I will never forget the first home visit I had with the special education teacher. She told me how Kieren didn't do this or that, wouldn't pay attention, kept flapping his hands, and wouldn't sit down and eat (tell me something I don't know!). During that visit I could hardly talk for fear of crying. I felt she was telling me what a lousy job I had done in bringing up my son. I blamed myself for a long time, thinking I should
"A Parent's Perspective"

I have done something and he would be normal. I thought that the "experts" knew what they were talking about, and I tended to listen more to them than to my inner feelings.

I am glad to say that now I know better. I listen to myself first, and then incorporate what the "experts" say into my plan. I've learned the hard way that parents are the experts on their child, and what's good for one child is not necessarily so for all children.

— Itala Eller, 1997, Parent

Malachi

I have a special sibling. My brother is intelligent and trusting and 2 1/2 years older than me. When we were little, we used to have bitter sock wars on laundry day, and he taught me all about dinosaurs (he knew a lot about dinosaurs). He drew these amazing, elaborate, creative pictures. I remember one called "War of the Grapes." It was a crayon picture of hundreds of little grapes in all kinds of armor and camouflage, and the red grapes were battling the green grapes. You looked at this picture and were amazed at the 7-year-old mind that constructed it.

Malachi also has a disability. For many years, my parents have been trying to figure out what disability it is. His symptoms have grown progressively worse since he was young, although,
at times, some medications have given us hope for improvement. Malachi is painfully shy. While he learned to make eye contact, you wish he would stop, it seems so painful for him. To ask him a question, you need to get his attention, then repeat the question until he understands, and wait patiently for his answer. Malachi never asks the follow-up questions to statements, and he rarely involves himself in a conversation involving more than one person. He has trouble with directions. While he nods and indicates that he understands, he may not recall the conversation when it is time to carry the directions out. Malachi was slow to speak, and used a speech therapist for a time after he learned his first words.

Later, Malachi started talking to himself, repeating the same words over and over while he worked them out in his complicated brain. When he was a teenager, he started talking more roughly to himself, pounding his fist in the wall, having even more trouble focusing. His counselors don't know how to label him. At different times, he has been diagnosed with learning disabilities, speech delays, a perceptual disorder, pervasive developmental disorder, brain injury, mild mental retardation, depression, schizophrenia, and disassociative disorder. He doesn't fit into a neat little box. But who does? At this point, I give up. My parents, still trying to find services to help him and guidance for their work with him, need a diagnosis. Otherwise, how will he be eligible? How will they know what interventions to try next?
When we were little, there was special education for Malachi, but he was intelligent, with no physical disabilities. The services offered to him were very limited. There was certainly no respite. When I turned 6, I entered public school with Malachi. My parents had recently gotten divorced, and my mother had to work, but she could not afford any of the child care options available to us.

We became "latchkey children." Malachi and I always came home right after school. Malachi always met me at home, and he went in first to check the closets and the basement for monsters and kidnappers. Together, Malachi and I did our homework and the chores that Mom laid out for us the night before. I am sure that, when Malachi thinks about that time, he remembers me as bossy. We bickered A LOT. (Imagine that! Siblings bickering?) We wanted to make sure that Mom would be in a good mood when she came home, and I knew that Malachi needed to be told to do things more than once in order to make sure that they got done.

When Malachi started experiencing new symptoms, I was starting high school, and I was given the opportunity to tune out. I got VERY involved in school activities and spent little time at home. My mother, exhausted by Malachi’s constant needs and frustrated because she still did not know what was going on with him, just could no longer have him in the home. Malachi moved to my Dad’s. For many years afterwards, I did not really know him or the
person that he became. Maybe if my life with him had not been so intense when we were younger, I would have stayed connected with him during that time. My respite in high school was WONDERFUL for me, but it came too late in my relationship with Malachi. As I work to reestablish my relationship with him, I have a lot of regret. I can look back and say that respite would have been very important to us all at a crucial time in our lives.

So, in Malachi, I have a special sibling. I never had a brother who screened my dates or scared off my bullies or who my friends had crushes on when I was younger. When my parents divorced, I never talked to Malachi about my anger, and he will not celebrate my wedding day the way that I always felt a brother should. We might never have a deep meaningful conversation about our life paths. I will always miss all those things with him. But Malachi is one of the greatest teachers I ever had, and, these days, I am learning to enjoy all of the things we are able to do together and appreciate all of the things that he is. He is a very special sibling.

— Anonymous, 1997, Sibling

A Sibling Perspective

My brother David Eugene Ownby was born with multiple disabilities—profound mental retardation, epilepsy, cerebral palsy, and microcephaly. I was 3 years old when my parents brought him home from
"A Sibling's Perspective"

the hospital. They gave me a life-size doll that day -- but I wanted the real baby. I always took my responsibility of being the big sister seriously. I made sure that I "protected" David. When other kids made fun of him, they quickly got an education in disability awareness. I made sure they knew that David had feelings just like the rest of us. Although he could not talk, he still knew when someone was making fun of him. Little did I know I was becoming an advocate.

David required a significant level of care. Both my parents worked, and, while most any babysitter would watch me, few wanted to watch David. My parents tried many variations of baby-sitters, from live-in nannies to next-door neighbors. Most often sitters would watch me and David for a week or two and then David would have a grand mal seizure. That coupled with the rush to the emergency room would usually result in the sitter agreeing to watch only me and not David.

Respite was not an option for my family in the early seventies. While my family worked with many therapists and social workers, the only options given to them were permanent foster care or institutionalization for David. Neither sounded very appealing, but my parents' ability to function as a family was severely compromised by the time David was 5 years old. My parents needed a break from the demands of caring for David. They approached the social workers about institutionalization. A
state of the art" institution had just been built 8 miles from our home. This service was touted as the best option available. Even still, the waiting lists were long, and David was denied admission time and time again. My father knew Senator Hubert Humphrey and, with his political clout, my parents did what they thought was the best thing for David and my family -- institutionalized David.

While having David cared for on a daily basis by caregivers in the institution was a relief, my family quickly learned that institutional life was not conducive to normalization -- it was not a home. My mom quit her job at the Library of Congress to manage the institution's library and be close to David. I quickly became involved with Special Olympics and volunteered there most Saturdays. My family and I stayed actively involved in David's life, bringing him home on the weekends and visiting often during the week. While this seemed to work for our family, I always felt that David was cheated. I did not think it was fair that he had to live there. I vowed at age 7 to "grow up" and change things for people like my brother.

I had to wait until I was 18 years old to become employed at the institution where David lived, and, when I did, I got a job there. I wanted to know what went on behind closed doors. While my mom had some sense of institutional life, she was far removed from the living units and day-to-day activities. I was employed as a direct care aide. I quickly learned
what I had always felt -- institutional life should not be an option for people like David. This environment was volatile, violent, uncaring, and stripped residents of their basic human dignity. As a young adult, I vowed to find David a better living situation. I wanted to become a disability rights advocate and work to change the system, allowing real choice for families and people with disabilities.

In seeking services for David, I faced what so many families still face -- closed doors, long waiting lists, and lack of services. The care David received at the institution was never good enough for me and had gotten considerably worse. More frequently, my parents and I met David at the emergency room with unexplained broken bones, bumps, and bruises.

In May 1992 David was left unattended in his institution bedroom after a series of grand mal seizures and aspirated. When institution aides found him, he was unconscious, with no respiration or pulse, and was cyanotic. He was rushed to the hospital and lived on life support for 2 months.

David had taught me long ago to never give up. He was certainly a fighter. He always bounced back from an injury or illness, and there were so many -- we all thought for sure he'd live through this too. While David struggled to live, this fight was too hard. In the time he had been left unconscious, he suffered significant brain damage. He was now blind, paralyzed, unable to swallow, and all the large...
organs in his body had begun to atrophy. My little brother was dying and my advocacy was too late.

David did die, but not in vain. While my efforts to help him move into a group home failed him, his death, my advocacy, and many others' resulted in 80 other residents having the real choice to live in a home. While I advocated to close this institution, after 5 years it remains open, but considerably smaller, better staffed, and closely watched by agencies providing oversight.

I learned from David to laugh, to have fun with the little things in life, and to persevere. I held David's hand as he lay dying and promised him I would work to ensure that no one else would have to suffer this way and that I would commit my life to making real choices for families and people with disabilities. I promised to share his story and make sure as many people as possible knew him. While David and I can no longer do McDonalds on a Saturday afternoon, and I'll never again have a big bear hug from him, he is alive in the work that I do. His life has given me my perspective.

— Lisa Ownby, 1997, Sibling

One Caregiver's Perspective

Respite caregiving for children with disabilities and their families truly presents an opportunity for a "give and take" relationship. This has been my experience from the time I decided to attend group
training, through working as group caregiver, and finally when a family “match” occurred.

My initial interest in respite was both personal and professional in nature. Being a single person without children left me feeling the need for contact with children, and my work in the social/health care field made working with children with disabilities a natural. I truly felt that I had personal traits of an affinity for children, patience, and a desire to learn and help a family. At that point, I signed up for training without knowing the specifics. What an informative, supportive, friendly, and nonthreatening experience!

Following initial training, small group exposure to children and the opportunity to discuss the children's needs and personalities with their families gradually increased my confidence and my joy in understanding that children with disabilities appreciate interest, hugs, snacks, toys, walks, playing ball, swings and slides as well! Challenges are there too! They may come in the way of feeding, behavior challenges, toileting, or special wheelchair accommodations, which may initially be unfamiliar to caregivers.

This is where the “give and take” occurs on the part of caregivers and families. Caregivers need training, feedback, and encouragement from the family “experts” in order to best understand the child’s limits and special needs. Caregivers may appear to ask many questions, seem anxious, and
anticipate what to do in an emergency. I believe families can aid caregivers by remembering their own initial thoughts and feelings about helping their child with his/her limitations. On the other hand, caregivers may want to appear knowledgeable and confident and not ask too many questions! Families, please anticipate this and VOLUNTEER INFORMATION!

My caregiving experience with a 9 year old and his 6-year-old brother has truly been rewarding and fun! This match occurred following group respite in which I had the chance to interact, observe, and build a relationship with both brothers and parents. A need for individual respite was there, and I had the time and geographic availability. Truly a match made in heaven! I really feel that this family provided information, training, and demonstration about their son's needs. I had the opportunity to question, observe, and build a relationship with the parents as well. I believe that caregivers need to maintain contact periodically in between respite dates to develop rapport. This also helps families feel freer to call upon the caregiver, particularly families that typically "do it all" or have difficulty in asking for help. Families and caregivers need to communicate and appreciate the gifts each has to bring to this partnership. The Partners Plus program is truly partners in action!

— Linda Malone, 1997, Caregiver
Stress

The potential for stress exists in all of our lives. Stress may be good or positive stress (eustress), or bad or negative (distress). The key to coping with stress depends on how you perceive a situation or life event. Stress is also closely related to your view of how hard or easy it may be to manage the event. For example, going on vacation can be perceived as both good and not so good. Many of us welcome vacations! Yet, vacations may be a major source of stress if you consider the packing, the drive, making reservations, or scheduling time to be difficult. Many strategies are available to both families and caregivers to help cope with daily stress. Included in the following pages are things families and caregivers can do to reduce stress, prevent burn out, and ensure successful respite relationships.

The Special Challenges of Parenting

Being a parent involves many changes, which may be perceived as stressful. Families who have children with special needs share the stresses of all families facing the challenges of parenting. In addition, they may have other challenges related to their children's special health, education, and care needs. Many resources are available to help families cope with these special challenges.
The Special Challenges of Caregiving

Being a caregiver for children with special needs can be stressful too. Caregivers are responsible for the children in their care. If those children need extra attention or medical care, caregivers are more challenged. In addition, it takes time to become comfortable with children and familiar with their needs and strengths. Not knowing what to expect from children can be stressful. Caregivers who are still learning about the children in their care may experience stress.

Coping with Stress

If you perceive a situation as stressful or hard to manage, you can reduce your stress by using a variety of coping strategies. Consider some ways you deal with stressful situations or special challenges. You might already know strategies that work well for you. For example, it might help to talk with family and friends in hard times.

Families often use respite to cope with stress. Taking a break from daily demands not only is relaxing, it allows families time to try the strategies listed on the following page. Families may want to talk with caregivers about how they use respite time to cope with stress.

Caregivers may need specific ways to cope with the challenges of working with children with special needs. Caregivers and families need to take time to talk after respite takes place. Was there anything that was stressful? Is there more information families can share? Are there
new skills the caregiver can learn? Certain caregivers may want soothing music playing as they work with your child(ren) or use other methods to create a calm atmosphere. Families can encourage caregivers to talk about how they handle stress. Together, caregivers and families can work out ways for caregivers to relieve stress during respite. The following fact sheet lists strategies you may find useful in reducing your stress.

Strategies for Dealing with Stress

People deal with stress in many ways. Here are some examples:

- Seeking information
- Sharing information
- Learning new skills
- Doing something that makes them laugh
- Listening to something they find soothing
- Planning ahead
- Finding a favorite cartoon that brings a smile to their face
- Walking, jogging, hiking, swimming, playing
- Communicating openly
- Taking a break
The Challenges of Respite

Building effective partnerships takes time and skill. Families and caregivers need to know how to work together for successful respite care.

One problem families often face is keeping caregivers. Families tell us many respite caregivers cancel appointments. We also know that many caregivers stop providing respite. Respite caregivers cancel or quit for many reasons. Why do you think they quit? The information in this section will help you understand why and help you develop strategies to build successful respite partnerships.

Caregivers may stop giving respite for the following reasons:

- Respite is harder than they expect.
- Time demands are too great.
- Physical demands are too great.
- Burnout occurs.
- Training was not adequate.
- Patience wears out.
- They don’t feel appreciated.
- They are not paid enough.
Meeting the Challenge

Taking these steps can help you meet the challenge of respite:

- Understand the family's purpose for respite.
- Work with the family to develop a thorough training plan.
- Discuss a schedule for providing care.
- Avoid overcommitting your time and energy.
- Be clear about your terms for providing respite.
- Learn more about working with families.
- Talk to other caregivers to share ideas and support.

More Things to Know About Families with Children with Special Needs

- Individual families have their own values, beliefs, customs, and courtesies, and what is most important to one family may not be to another.
- Sometimes families have fears and bad feelings about their children's disabilities.
- Families with children with special needs generally have the same kinds of daily routines as other families.
- Families with children with special needs have to learn more about how to help their children.
Children with special needs like to play and have fun.

Children with special needs and their families can help others learn new things, like sign language.

Children with special needs grow and develop with love and support from their families and others.

Children with special needs and their families want to be accepted by others.

Families wonder if their children with disabilities can do things that other children do.

Families hope there will be services to help them and their children at school and in the community.

Families have information about their children's special needs to share with caregivers.

Families need others to know that it is okay to have children with disabilities.

How You Can Help Families

Remember that families and caregivers bring unique backgrounds and experiences to the respite caregiving partnership. Some ways to show respect to a family are being on time, keeping an appointment, and keeping scheduled times and dates to provide care.
There are also other ways that you can help families:

- Be supportive.
- Communicate honestly and openly.
- Ask questions.
- Be an advocate for them.
- Help recruit other caregivers.
- Be dependable.

**Successful Family and Caregiver Partnerships**

Families and caregivers can build successful partnerships by doing these things:

- Appreciate differences in skills and approach.
- Be willing to learn from each other.
- Ask each other for help.
- Allow time to talk with each other.
- Share experiences.
- Be willing to work together, even when differences occur.
- Be flexible in trying new ways.
- Have a sense of humor and fun.
- Be patient when things are not clear or exact.
- Share commitment to children with special needs.
Section 4

All Kids Like Cookies

activity
All Kids Like Cookies

Children with special needs are important members of respite partnerships. The following activity is designed to allow you the opportunity to experience some of the daily challenges children with special needs face. This may be useful in understanding a child's perspective and keeping that in mind when developing respite partnerships.

We know children with special needs are more similar to other children than different. "All Kids Like Cookies" is a series of activities designed to help persons without special needs experience challenges children with special needs face every day. You can do the activities on the next pages by yourself or with a group. Answer the questions at the end of each page as you finish each activity. This will help you become more aware of what it is like to have special needs. As you will see, from the perspective of the child with special needs, the world is a more challenging place.

We cannot help until we understand.
We cannot understand until we feel.
We cannot feel until we become.
(Council for Exceptional Children)

Adapted from:
All Kids Like Cookies Activity Sheet

The following activities can be done alone or with a group. During the activities you remain yourself, but a special challenge is added. Think about how you feel when doing each of these activities and be aware of any adaptation you make.

To experience a visual impairment, you will need:

- a blindfold
- a pitcher of water
- an empty glass
- a crayon or other marker
- a page out of a coloring book or a crossword puzzle
- a child’s storybook or magazine

- Put a strip of cloth over your eyes and tie it around your head to make a blindfold.
- Pour water from the pitcher into the glass.
- Color the coloring book page or complete the crossword puzzle.
- Read the storybook or magazine.

Questions

- How did you hold your head while pouring water, coloring, and reading?
- If doing this activity with a group, did anyone talk you through the tasks?
Activity Sheet

- What other senses did you use to help yourself? (touch, sound, smell)
- Did it matter that you were coloring out of the lines or what color you were using?
- Were you able to complete the crossword puzzle?

Vision helps a child learn about the objects and people in the world around them. Few children are totally blind. A child is considered to have a visual impairment if the child’s ability to see after correction limits his ability to learn through vision.

To experience a speech and language impairment, you will need:
- a bag of large marshmallows and gauze
- one other person or a small group

- Put enough marshmallows or gauze in your mouth to fill it; two or three usually work.
- Say your name and give your address or phone number.
- Ask your partner or others in the group to bring you something.
- Tell your partner or others something you did today.
Questions

- How were your words spoken differently?
- How did you feel when others did not understand you?
- If you spoke this way all the time and others laughed at you, would you want to talk very much?

Sometimes children with speech and language impairments stop talking rather than cope with the difficulty of getting people to understand them.

To experience a physical disability that affects using the smaller muscles in your body, like the ones in your fingers, you will need:

- a pair of large garden or work gloves
- a puzzle
- a string and macaroni with holes or beads
- very small toys that have moveable parts

- Put the gloves on.
- String the macaroni or beads.
- Put the puzzle together.
- Play with the toy.
- If you are wearing a watch, take it off and put it back on.
- If you are wearing earrings, take them out and put them back in.
**Activity Sheet**

**Questions**
- Were you able to pick things up easily?
- Did you need more time to put the puzzle together?
- Were you able to put on your watch and/or earrings?

**To experience a physical impairment that affects the large muscles of the body (like the ones you use for walking) you will need:**
- a kick ball
- a large toy like a stuffed animal or truck
- a video

- Place the ball between your ankles and keep your hands folded together.
- Walk 10 feet with the ball between your ankles and pick up the toy or video.
- If doing this with a group, hand the toy or video to someone else.

**Questions**
- Did you take a long time to get to the toy or video?
- Did you feel awkward?
- Did you drop anything?
Children first learn to explore their surroundings by using their large muscles to move around. Children learn how things work by using the small muscles of their hands to explore toys.

Children delayed in their motor development have difficulty learning to become physically independent because they cannot move well. Children with small muscle development delays may have difficulty feeding and dressing themselves or learning how things work because they cannot use their hands and fingers well.

Source:
Notes on this Section
Section 5

disabilities

awareness
Understanding Disabilities

The purpose of this section is to increase your knowledge and awareness of the terms, services, and issues related to disability services. This section contains many ideas, words, and issues that may be new to you, although these are everyday concepts for families of children with special needs.

For parents of a child with special needs, every day becomes a day of learning. Parents may learn how to adapt a cup so their child can learn to take a drink. A parent may learn about a new treatment procedure from a family doctor. Parents may also learn more about the educational system and its services and about the many words and acronyms used by professionals who work with their children. As a caregiver, you will be able to learn from Partners plus training and from the families of children in your care.

Included is a summary of the educational services that are available to children with special needs and the terms and abbreviations that are often used. As you learn more from group and personalized training, you will begin to understand some challenges that families face. Through understanding, we can all work together to help others in our community become aware of the issues affecting children with special needs and their families.

Children first learn to explore their surroundings by using their large muscles to move around. Children learn how things work by using the small muscles of their hands to explore toys. Children delayed in their motor development...
have difficulty learning to become physically independent because they cannot move well. Children with small muscle development delays may have difficulty feeding and dressing themselves or learning how things work because they cannot use their hands and fingers well.

**Source:**

**Educational Services**

- Federal legislation called IDEA (Individuals with Disabilities Education Act) is now in place to make sure that children with disabilities receive special services. Infants and toddlers with disabilities (ages birth through 2) may receive early intervention services. The public schools provide children from 3 to 21 a free and appropriate public education. For preschool children (ages 3 to 5), services are called early childhood special education (ECSE) services. Older children receive special education services. Beyond school, families and individuals with disabilities rely on residential and vocational support services from both public and private providers.
Finding Educational Services

- Children who receive special services will have an individual plan written for them each year. For children under age 3 in early intervention services, the plan is usually called the Individualized Family Service Plan, or IFSP. In special education services, the plan is usually called the Individualized Education Program, or IEP. In residential and vocational services, it is usually called the Individualized Service Plan (ISP) or Individualized Transition Plan (ITP).

- The IFSP, the IEP, the ISP, and the ITP all include goals for each child, information about the child's development, and activities to be used to help the child reach his individual goals. Ask the family if they will share their child’s plan with you so that you will know more about the child’s development and how you can help. Because an IFSP includes a plan for the family, it may be personal and you may want to ask families to share the child portion of their IFSP only.

- A plan must be developed by a team that includes members of a child’s family. Team membership varies and may include people such as a special educator, a speech pathologist, an occupational therapist, a physical therapist, a nurse, or others.

- Your observations and ideas about the child’s activities during respite are important for the family to have when they help develop their child’s plan.
Understanding Early Intervention

What Is Early Intervention?

Early intervention is a system of services for young children, birth through 2 years of age, and their families. It is designed for children who have disabilities or a delay in development and, in some states, for children who are at-risk for developmental delay. In early intervention, parents and professionals work together as partners. Each brings important information and skills to the partnership.

The agency responsible for early intervention is different in each state, and early intervention services are provided by many different agencies, programs, and professionals in each community. If a child is eligible for early intervention in your state, federal and state policies help to ensure that each family gets the services they want and need.

Adapted from:

Why Are There Programs and Services for Infants, Toddlers, and Their Families?

- Working with some babies early in life helps to prevent later problems in their development.

“Our dream has become a simple one . . . for Becky to experience life as other kids do, and for her to be happy.”
Understanding Early Intervention

- Experience and research has shown that working with children with disabilities early in life helps reduce the effects of their disabilities.

- Early intervention programs and services help families with concerns they have for their children with special needs or disabilities.

- Although many children learn to crawl, talk, and play without much help, some children need special help. Early intervention can help families become more skilled at encouraging the development of children with special needs.

- Early intervention programs and services can help families learn more about child growth and development and how to manage children's behavior.

Adapted from:

The following are firsthand stories from families to help you learn a little more about disability services.
Matthew was born with chromosomal anomalies that affected every cell in his body. The doctors prescribed his prognosis as quite grim. His medical and developmental problems were so pervasive that his life span was predicted to be short and his quality of life poor. During the 3 years of our son's life, he did not grow beyond 5 pounds and did not develop beyond infancy.

Our son's needs and our family's needs were not well matched to the more traditional kinds of early intervention services. Since our family had decided early on to care for Matthew in our home and to pursue a family lifestyle as close to the way we saw other typical families, we needed an early intervention service that would help us meet these goals. Early intervention helped our family by providing services like respite care through a center-based play group.

Another service I found helpful was the parents' group where I received a lot of support from other families with children with disabilities. Other parents and the staff of the early intervention program helped me find ways to cope with the everyday stresses of caring for a chronically-ill child and still be a "mom" to my other child. Through their help and special friends in the community, I could continue to enjoy life for myself and with others. Professionals helped me and my husband understand the medical implications of caring for our son so that we could keep Matthew at home. Physical and occupational therapists helped show us
ways to keep Matthew as healthy and comfortable as possible. The greatest help given to us by early interventionists was in sharing with us information about our son's disability and in encouraging us as a family. We gained the confidence we needed to live with and care for our medically fragile child.

— Dee Moore, 1997, Parent

"Jack's Story"

Jack was born 8 to 10 weeks premature, and it was over a year before doctors would agree to tests to confirm my suspicions that Jack was more than just premature. After a battery of tests, he was diagnosed with spastic quadriplegic cerebral palsy with a chance of mild to severe mental retardation. Those words changed our lives forever, first accepting the diagnosis and then striving to do the best for our son.

One of the best things we learned as parents is that what we feel and have been advised for Jack is best is not necessarily what the school system deems to be required. But having said that, I will say that the early years in the school system were much easier than the later years have been. Jack's needs were met in ways that were beneficial to me as a parent as well as Jack. I learned a great deal from his teacher and his physical, occupational, and speech therapists, ideas that were carried over to the home to provide continuity.

Jack's elementary school years were spent in a segregated classroom. There were no typically-
developing kids in this classroom; only kids with special needs. By the time Jack entered school, he was classified not only as orthopedically handicapped, but also as visually impaired and learning disabled. He had a severe startle reflex and did his best work in a one-on-one setting. With Jack's teacher and therapists, we tried many ways of teaching Jack and found that he could orally comprehend at a much higher level than he could read. By properly using his IEP, we could plan for Jack individual needs and how best to meet them.

Later, Jack was placed into some regular classes (with typically-developing kids) with a full-time aide. The aide allowed Jack to work on his independent goals without taking away instructional time from other students. Being in regular classes allowed Jack to be with a wider variety of people, increase his social skills, better his self-esteem, and generally feel more a part of the school. His IEP has allowed refinements over the years to accommodate Jack's abilities such as auditory learning skills and providing for his tests to be taken orally with the help of his aide. However, IEPs often look good on paper but are not always followed in practice.

We have learned there are several important things that can be done to benefit Jack in the school system. First, know his rights and our rights as parents in the educational process. Families are a child's best advocate and at times may be the only advocate. Second, parents have to be knowledgeable of their child's needs and what must be done to meet those needs. Third, there are public and private organizations and advocates who are willing to help, often at no charge to the family. They provide support, knowledge,
"Thoughts on Special Education"

and backing needed to make a child’s rights known. School systems have, in many cases, become more concerned with the monetary bottom line and often do not provide information about a child’s rights and entitlements.

Our most negative experiences have been with those teachers and other personnel who see our son not as a student but as “handicapped.” In those cases where teachers and others care and have the desire and ability to accept Jack as a student, and then figure out ways to accommodate his needs, a great deal more progress has been made with a lot less effort.

— Susan Bohannon, 1997, Parent

"A Parent’s Thoughts on Special Education"

In addition to special education, my son also receives physical therapy from a certified therapist, and he receives vision services since he is also visually impaired. Unfortunately, my son, and often other children as well, does not receive the amount of therapy recommended by a therapist because schools have limited budgets and must serve many children. However, I have found my son’s teacher to be extraordinary. She is very helpful in getting information to parents. This is true of the parent resource person also. Parents share information and resources with each other as well. Unfortunately, I have found that often I must be an active advocate to ensure my son receives the services he needs. Equipment is often hard to get and acceptance by some regular education teachers and other students can be hard to come by. The IEP process can help
families get the services their child needs, but in the beginning it can be intimidating. I have learned that the IEP is very important to my son's education.

— Teresa Lewis, 1997, Parent

Common Issues Families Face

The following are some common issues facing people with disabilities and their families.

Transition

For families who have children with special needs, life is full of transitions. From the time these families bring their children home from the hospital, many are involved in a service system designed to help support the family and meet the needs of the child. Families are initially involved in the early intervention system. Then they may make a transition to early childhood education and then on to special education, and ultimately they may tap into vocational and residential supports for the child.

Once in the school system, children will move from elementary,
"A Parent’s Perspective on Transition"

to middle, and then to high school. While all families who have children will experience these transitions, for families who have children with special needs, this means not only new schools but new therapists, new social workers, new nurses, a new Individualized Education Program (IEP), etc. These families must constantly learn new systems of support. In some respects it makes getting services for their children very difficult, because each system is so different. So, often when a family feels like they finally understand one system of support, their child moves on to another, and the process of learning the system starts all over again.

Service systems can be very frustrating for families and their children. Not only do systems change as the child ages, but they differ from locality to locality, from state to state, and around the country. This can be especially difficult for military families and for families who must move a lot due to their jobs.

"A Parent’s Perspective on Transition from Preschool to Kindergarten"

When Patrick was born, he seemed normal in every aspect. By the time he was about 18 months old, we realized that something did not seem quite right. He did not talk much, walked a little late, and had almost no interest in people. All his pediatrician said to us was “Do not worry—all children develop a little differently.” When Patrick was 2 1/2 years old, we moved to Virginia from New Hampshire. Shortly after we moved, Patrick stopped talking altogether and his temper tantrums were horrible. We kept thinking it had something to do
with the move, not having other children to play with, or we were doing something wrong. At his 3-year checkup, his pediatrician sent us to specialists. He was tested and observed, and on a rainy day in November, the diagnosis came. "Autism."

We called our local school system to see about early intervention services, and Patrick again went through more testing and observation. After all was done, we were delighted to find out he would be put in a special education preschool class in a local elementary school. We were thrilled! There was somebody out there to help us at a point when we felt as though we were losing ground. Patrick was put in a small class of seven children with a good teacher and a wonderful aide. He adjusted fairly quickly, and we could tell he loved school. Patrick got speech therapy and occupation therapy as part of his special services. In addition to language and other preschool activities, Patrick also worked on things such as potty training, dressing and undressing, how to play with other children, and gross and fine motor skills. We also got home visits from the teacher to discuss problems and concerns and to help us whenever needed.

Recently we had our first real IEP meeting. Patrick is going into kindergarten. Getting ready for the IEP meeting was a little scary. Patrick was going to a new school in an inclusion program. We were dealing with an all new staff. Would they be willing to work with us to get everything our son needed? In preschool, we had gotten to know the school staff and pretty much knew
"A Parent's Perspective on Transition"

what to expect. We were also not sure if we would get all the services we felt Patrick needed. Patrick had also been riding on a small bus with only his peers; next year he will be on a bigger bus with children of different ages.

The IEP meeting went well, although we are somewhat still concerned for his next school year. In preschool he was taught with all developmentally-delayed children, and this would be his first time being taught with nondisabled peers. How will he match up with them? How will they treat him? Will he make friends?

Patrick will have the same speech and occupational therapists, which is at least one constant for him and something he needs. The goals for next year changed also because he will deal more with academics, writing skills, and language, although social skills will still be very important. That was a big concern for us. We were afraid that if Patrick did not have an aide that he would be lost in the shuffle. We are still nervous and have many questions in our heads, but, after meeting the special education teacher and vice principal and going through the IEP, we feel better about the new experience ahead of him. With the help of teachers, aides, and everyone else involved with him, we feel sure that he will do fine.

—Deborah Warren, 1997, Parent
Accessibility

With the passage of the Americans with Disabilities Act (ADA) in 1990, America has made many changes to help make things more accessible for people with disabilities. The ADA requires that public buildings be accessible for people with disabilities. This means there must be curb cuts, elevators, easily usable door handles, timed door openers, braille signage, TDD/TTY (access to phones for the hearing impaired), etc. Likewise, ADA prohibits businesses from forcing people with disabilities to use back entrances. For years, the only way into some buildings was through kitchen or delivery doors. This was demoralizing to people with disabilities. With the ADA, people with disabilities have access through front doors like everyone else.

While the ADA represents progress, many facilities remain inaccessible to people with disabilities. Some buildings, such as religious or historic buildings, are exempt from adapting their facilities. In addition, all buildings have a certain amount of time in which to comply with the ADA. Finally, agencies can get exemptions from the ADA when the costs of adapting facilities are too high. As a result, despite this legislation, many places are still inaccessible.
Common Issues for Families

It is important to appreciate how much forethought persons with a disability or families of children with special needs must make each time they leave their house. They must plan ahead in case they are unable to enter a building, read a sign, hear a presenter, see an overhead, roll off the curb, open a door, etc.

As mentioned earlier, accessibility means more than wider doorways and ramps. Accessibility is also about stigma. Even with the ADA, our society is not always comfortable with disability and this is often reflected in the words people use, in uncomfortable silences, and in unnerving stares. So, often, even when a building is accessible, the people inside are not welcoming. This is very discouraging for people with disabilities and their families.

One thing that a parent of a child with disabilities desires most is for their child to develop friendships with other children of the same age. It is difficult for this to happen unless children have an opportunity to be together. Enrollment in a program is not enough. Once enrolled, the child must have access to the activities going on in the child care setting. Interaction occurs when caregivers have arranged an environment that sends the message to children that says “this is a place for me.”

A good way to think about accessibility is to arrange the setting so that anyone can see that it is a place that includes all children in play, learning, and recreational activities. Accessible environments should have play materials and equipment that are seeable, reachable, and usable for all children. A setting that offers a warm welcome to all who wish to use it is accessible.

Language

People with disabilities and their families are sensitive to the language we use to refer to disabilities. It is helpful and respectful to use people-first language. This is a way of speaking that mentions the person before the disability. A person may be referred to as the blind child or epileptic man. Using people-first language, you would say the child who is blind or the man who has epilepsy. This way of speaking makes it clear that the disability does not define who the person is but is one important characteristic.

People with disabilities across the country have expressed their preferences about the words that we use to describe their special needs. Besides using people-first language, the word disability is preferred over the word handicapped. The term special needs is often used to describe children with a range of disabilities, developmental delays, and/or special health care needs. Over the years, people with and without disabilities have changed the words and the way in which we talk about disability.

Years ago you may have heard someone described as wheelchair bound, crippled, challenged, stricken with... These words are no longer preferred by the disability community. Language and words we use change with time.
How Parents Describe Their Child

Because language changes, it is important as a caregiver to always use people-first language and to listen to the words that families use. Those are the words that you should use when talking with them about their child. For example, parents of older children or adults with disabilities may refer to their child as handicapped, because that was the word used many years ago and they may still be comfortable using it. Don't feel like you must correct a family in the words they use. Rather, model people-first language and use the words they use to describe their child.

Describing Children

Parents of children with special needs may choose to describe their children very differently from the ways in which doctors or other professionals would describe their children. Some parents have described their children in the following ways:

- A great kid who loves Barney, Wheel of Fortune, and girls.
- Quiet, handsome, and a bit of a challenge.
- Very happy and expressive. Physical abilities between 6 and 9 months but highly cognitive. Spoiled rotten.
- Calm, cuddly, and easygoing.
- Lovable and easy to care for. Responds to touch and sound. Loves attention, but plays well independently.
How Parents Describe Their Child

- The sweetest, cutest, most adorable child on the face of this earth.
- Good natured and talkative.

It is important to recognize that disability is a natural part of the human experience. Many of us will experience disability in some way, either personally or in our immediate family, in our lifetime. People with disabilities are actively involved in the development of public policy that shapes services and how we think about disability. It’s important to be aware of the trends and to listen to the people we support through respite and other services.

Professionals who provide services to people with disabilities and their families create and use many technical terms and abbreviations. This is often a challenge for families to learn. As each child grows and matures, the language changes and a family must learn new words and abbreviations. You can experience the challenge of language through the activities What Words and Alphabet Soup on the following pages.
What Words Activity

Language is very powerful, and we as caregivers must be sensitive to families. Words like *retard* and *pitiful crippled boy* are very hurtful and condescending.

Instead of ...  Use ...

Handicapped girl
Retard
Crip
Wheelchair bound
Blind boy
Autistic child
Alphabet Soup

As you continue your training and begin working with families, you may hear new words or letter combinations regarding children with special needs, services, and/or other support for children and their families.

Use this worksheet to write down the new words and acronyms that you learn. Remember that we are all learning together, so share your new knowledge with others.

<table>
<thead>
<tr>
<th>IFSP</th>
<th>Individualized Family Service Plan</th>
</tr>
</thead>
<tbody>
<tr>
<td>IEP</td>
<td>Individualized Education Program</td>
</tr>
</tbody>
</table>

---

Partners Plus Caregiver Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

946 DRAFT 10.99
Some Closing Thoughts

Summary

As a respite caregiver you will have an opportunity to meet and work with some wonderful families and their children. In so doing, you will learn about a child, his or her family, and disability awareness. You may even experience part of the stigma associated with disability. You can help with disability awareness by sharing with your friends and family what you have learned and continue to learn from families. It is most important for society to know and recognize that children with special needs are children first. You can be an advocate for families and children everywhere by increasing others’ awareness of the issues affecting people with disabilities.

Remember that, while children with special needs are given labels and diagnoses in order to obtain needed services, they are children first who are loved very much by their families.
Section 6

Understanding Child Development
Planning Appropriate Experiences

Understanding Child Development

Providing respite care for children with special needs is not so unlike providing care for typically developing children. Respite caregivers who work with children who have special needs do require some special skills and information to do their jobs effectively. However, the most important knowledge in caring for children is understanding child development.

Plan Appropriate Experiences for Children

We need to understand the order in which we can expect to see different skills develop in children so that we know what children can be expected to learn.

Example:

A very young child first plays alone. Next we expect a child to play with toys close to other children, before being ready to join in play with one or more children.

When we know the range and sequence of typical development, we are able to know what skills we can help a child learn with success and what activities to plan.

Adapted from:
Plan Activities That Give Children with Disabilities a Chance to Use Their Developmental Strengths.

Many children with disabilities have some areas of development in which their skills are similar to those of their peers. Emphasizing those strengths is important to each child’s confidence and overall development.

Example:

A 4-year-old child with mental retardation may not be able to play with others or take part in social activities typical for children her age. However, the way she uses her body may be typical. She can climb on play equipment, ride a trike, and run and jump like other children her age. Giving her opportunities to practice using her muscles will help her to continue developing her abilities, help her to feel good about herself, and give her ways to play with other children. You can help her expand her social skills by building on her strengths, playing near or with other children on playground equipment.

Another Example:

In games where groups of school-aged children participate, caregivers can find opportunities for children with disabilities to take the lead and demonstrate their abilities among peers.
Areas of Development

In games where groups of school aged children participate, caregivers can find opportunities for children with disabilities to take the lead and show their abilities among peers. A child with visual impairment whose motor skills are like that of their peers could take the lead in a game of pinata. In this game, a brightly colored ornament is suspended above children's heads, and blindfolded players take turns hitting it with a stick to break it open. Taking the lead in this game helps the child gain confidence and build self-esteem.

A Knowledge of Child Development Will Help You Care for All Children.

Areas of Development

Child development is an orderly sequence of physical, intellectual, and emotional change that occurs throughout childhood. Although a child develops as a whole person, different words and categories are used to describe different areas of development.

For a child with a disability, understanding the different areas of child development will help us understand the child's development. However, a disability in one or more areas of development frequently affects other areas of development as well.

Adapted from:
Areas of Development

Physical Development

Physical development includes gross motor, fine motor, vision, and hearing.

- **Gross motor** refers to how children use their large muscles for activities like sitting, crawling, walking, and climbing.
- **Fine motor** refers to how children use their small muscles for activities such as grasping objects, coloring, turning knobs, chewing, and swallowing.
- **Vision** refers to how children are able to see. A child may have a narrow view of things or may see blurred shapes.
- **Hearing** refers to children's ability to perceive sounds. Do they hear muffled sounds or high or low pitches?

Communication Development

Communication development describes how children let us know what they want, need, or understand.

- **Expressive language** describes how children communicate by using sounds, words, gestures, or communication devices.
- **Receptive language** describes how children understand messages from others.
- **Cognitive development** refers to the ways in which children think, understand the objects

**Adapted from:**
and experiences in their environments, and solve problems.

- Adaptive development refers to the ways children take care of their personal needs such as dressing, brushing teeth, and drinking from a cup or bottle.
- Social or emotional development refers to the feelings and thoughts that a child has about herself and the people around her.

Young children who do not yet show skills expected at their age in one or more areas of development may be considered to have developmental delay.

Basics of Child Development

There are several principles that describe how development occurs. There are patterns, characteristics, or trends that seem to relate to the growth of most children. These principles apply to the growth of children with and without disabilities.

Adapted from:
Lightfoot, VA: Child Development Resources.
Development is sequential

Development is sequential. There are steps in each area of development that occur in a certain order. Sometimes children do skip steps in the sequence, but steps do not get reversed or mixed up.

Sequential steps in the development of motor skills:

1. A newborn develops from head to toe.
2. A baby strengthens her neck muscles to control her head.
3. We see baby on her tummy pushing against the bed strengthening her arms, shoulders, and trunk to lift her chest.
4. On her tummy she pulls her knee up and turns her shoulder and hips to roll over.
5. Using her whole arm, she pulls forward and pushes with her feet to creep along.
6. She will pull her knees and feet up under her, push with her arms, and then move her arms and legs in an alternating motion to crawl.
7. She reaches to hold onto something while pulling up to standing.
8. Standing and holding onto something for balance, she cruises around furniture.
9. Without holding onto anything, she takes her first steps.
One Area of Development Affects Another

One area of development affects another. Growth in one area of development is not separate from that in other areas. A child's natural play and everyday routines can make growth happen in all areas of development at the same time. A child may have a disability in one area of development that also affects other areas of development.

*Visual impairment may affect motor development. If a child cannot see well enough to move about alone and to practice crawling, walking, or running safely, the child may become delayed in motor development.*

Children Have Individual Rates of Development

Children have individual rates of development. Some children seem to be ahead of others in their overall development. Other children may be behind but will catch up later or will catch up in one area of development but not in another.

*One child may be using many words while the second child is not; but the second child can ride a tricycle while the first child cannot.*

Adapted from:
Individual Development Depends Both on Inborn Characteristics and Environment

The path of individual development depends both on inborn characteristics and on a child's environment. Children seem to be born with a range of possible abilities in different areas, such as intelligence. An environment that gives a child lots of different experiences will help a child to strengthen his natural abilities.

A child born with a serious hearing impairment will have problems developing spoken language. Pronunciation, vocabulary, and speaking in sentences will all be affected. An environment rich in communication experiences, started early in the child's life, will help the child develop communication skills.

Development Is a Combination of Maturation and Learning

Development is a combination of maturation and learning. Children cannot learn certain things until they are physically ready.

We cannot teach a child to ride a tricycle just by telling her how. The child's legs must be long enough to reach the pedals and strong enough to push them. The child must have the strength to grip the handlebar. The child must have a sense of balance. The child must be able to alternate her legs and feet to cycle.

Adapted from:
The Importance of Play

Play is essential to children's development. It is the way in which children grow and learn. Through play, children learn to solve problems, get along with other people, and use their bodies to explore the world around them.

A child can develop creativity and imagination when playing alone with a toy. When children play with others, they grow emotionally and learn social skills. Toys and games should be selected to allow children to use them in many ways, based on their development and interests.

Social Interaction

In group respite settings, it is important for caregivers to help children with disabilities and children without disabilities to interact socially.

Sometimes children with disabilities have difficulty learning social behaviors. At first, they may need a caregiver's assistance to interact with others.

Children with disabilities may have more subtle or unusual ways to communicate or signal for attention. They may have a slower response time, be delayed in language, or have physical limitations that affect their interactions.
Once a caregiver is aware of how a child's disability affects his or her ability to interact with other children, you can help by modeling appropriate techniques to engage the child and by teaching those techniques to other children. In order to promote social interactions, caregivers should:

- Encourage children to play together and acknowledge them when they do
- Remember that too much attention from adults may interfere with the children's interactions
- Teach children specific ways to ask other children to play, share toys, take turns, express affection, and help other children

Sometimes staff of community programs think that they will have to make extensive changes to their programs to include children with disabilities. Often times only a few accommodations are needed to meet the needs of children with disabilities. Valuable respite to families with children with disabilities can be offered by providing the same kinds of appropriate experiences that characterize quality programs for all children.

Adapted from:
Getting Involved in Play

Guidelines for Caregivers’ Involvement in Play

Caregivers should play with a child by following the child’s lead. This may take some practice. Here are some guidelines to help you.

- **Observe**
  Watch children to see what skills they already have and what they like to do.

- **Follow**
  Join in when a child invites you. Let the child be in control and play at his level.

- **Be creative**
  Discover or rediscover the many different ways to play.

- **Be safe**
  Be sure toys, activities, and play space are safe and developmentally appropriate for the child. See page 10-15 for the health and safety checklist we give families. Consider these things as you work with families and children.

- **Have fun**
  Relax and enjoy each other’s company.
Section 7

children with special needs
Understand Each Child’s Needs

Children with Special Needs

There is a wide range of disabilities and special health needs. We will use this section to give you some information on some of the more frequently occurring disabilities. As you read, remember that the same disabilities and health conditions can look very different in individual children. Families are the experts on their own children! They will give you specific information about their children’s special gifts and special needs.

Attention Deficit Disorder (ADD) / Attention Deficit Hyperactivity Disorder (ADHD)

Children with these diagnoses have very short attention spans. They often have difficulty focusing on tasks. They may be impulsive, and, if they are diagnosed with ADHD, they are physically active as well. Children with ADD or ADHD often need very structured routines in order to feel comfortable. They may take medication to improve their attention spans. Families may also have clear rules for these children and specific consequences for following or not following the rules.

Questions to Ask:

- Does Wendell have a routine during the day? How does that work?
- Does Wendell take medication? When does he take it? Does he have a special medication routine? Does the medication have any side effects I should know about?
Understand Each Child’s Needs

- Do you have household rules that Wendell must follow? What are the consequences if he doesn’t follow the rules? What about rewards if he does follow them?

Autism

Children with autism often have trouble with social and communication skills. Some children with autism never speak at all; others achieve academic success but may have less than fully mature social skills. Children with autism may need to have the same routine every day to feel comfortable.

Questions to Ask:

- How does Joey communicate? How do you make sure that his needs are met?
- Does Joey have a set schedule? What is his routine?
- What is the best way to help Joey make transitions?

Blindness/Visual Impairment

Visual impairment is less severe than blindness. Most people who have a visual impairment have some level of vision. They may see light, or they may see fuzzy images. They may have vision like they are looking through a tunnel. They may have spots in their vision. Children who have a visual impairment may have trouble seeing, even when wearing contact lenses or...
Understand Each Child's Needs

eyeglasses. Because children often learn to use their bodies by reaching for things that they see, children with visual impairments may show delays in gross motor and fine motor skills.

Questions to Ask:

- What type of visual impairment does DiShannon have? What can she see?
- How is DiShannon doing with her motor skills?
- How does DiShannon move around? What is the best way to lead her when we are walking together?
- Is DiShannon familiar with the rooms in the house? (If you are working with DiShannon in your home, it may take some time for her to learn the layout of your rooms. She will need to explore to find out where the furniture is or where the rug stops. Be particularly aware of safety issues. Are there glass objects she could knock over? Are there steps that she could fall down? Stay close with DiShannon as she learns about your home.)
- When I set the table for dinner (or set up another activity), where should I put DiShannon’s things so she can find them?

Deafness/Hearing Impairment

Deafness is the inability to hear. Deaf children may need to watch lips or to learn a language of gestures, like sign language. Hearing impairments are less severe than total
Understand Each Child’s Needs

dearness. Children with hearing impairments may hear only loud noises. Sometimes, they may only hear people when they speak at very high or very low pitches. They may be fitted, as early as infancy, with hearing aids.

Questions to Ask:
- What type of hearing impairment does Juan have? What can he hear?
- How does Juan communicate?
- How do you communicate with Juan?
- How do you get Juan’s attention?
- Does Juan use a hearing aid? How does it work?

Emotional Disturbance

Children with emotional disturbances usually have trouble with social skills or making friends. There are many different kinds of emotional disturbances. Some children are very sad or react in ways that you would not expect. For example, they might laugh when they are hurt. Some children with emotional disturbances are very anxious and difficult to calm. Others may be very aggressive. They may fight a lot, or they may be very defiant. As a caregiver, you might need to watch children closely on the playground and with other children. Children with emotional disturbances frequently need a lot of encouragement and praise.
Understand Each Child's Needs

Questions to Ask:

- What type of emotional disturbance does George have? What does he do that is unusual?
- How do I calm George when he is upset?
- Does George take medication? Does he have a medication routine?
- How do you try to control George's behaviors? If he fights with a child, or doesn't listen to me, what should I do?
- What does George enjoy doing? What kinds of things is he good at?

Learning Disability

A learning disability affects the way that a child understands or gives information. A child with a learning disability may have trouble with reading, writing, math skills, listening, speaking, or using logic. Dyslexia is an example of a learning disability. The person with dyslexia has trouble reading and writing, though he may be smart and does not have any hearing or visual impairments. Some children with learning disabilities might have trouble understanding what you are saying to them. Some might have trouble remembering what you told them just a few minutes before. Caregivers working with children with learning disabilities might see motor delays or language delays as well.
Understand Each Child’s Needs

Questions to Ask:

- How does Lahn learn best?
- What is the best way to present information to her? Is it better if I tell her something, draw a picture, or write it down?
- What activities frustrate Lahn?

Mental Retardation

Children with mental retardation may have difficulty learning and remembering what they have learned. They often learn new skills at a slower rate. Because mental retardation is usually diagnosed through testing, a family may be able to tell you at what age level their child is operating. For example, a child may be 8 years old, but a family may tell you that he functions at a 4 year old’s level. This is valuable information! Think about the development of a child not his size or age. He may be big but may still have trouble playing games with lots of rules.

Questions to Ask:

- What does Mark do well? What should I encourage him to do on his own?
- In what areas does Mark need help? How can I help him?
Understand Each Child’s Needs

Orthopedic Impairment

An orthopedic impairment affects the bones, muscles, and/or the way the body moves. Many diagnoses are included in this category. A child who has a leg amputated is considered to have an orthopedic impairment. Children with muscular dystrophy, which causes muscles to deteriorate over time, develop orthopedic impairments. Children with spina bifida, which limits the brain from sending messages to healthy muscles because the spinal cord was not closed at birth, or cerebral palsy, which also limits the brain’s ability to control muscles, are considered to have orthopedic impairments. Children with orthopedic impairments generally show motor delays. Depending on which of their muscles are affected, children may also have trouble communicating, eating, or toileting.

Questions to Ask:

- What type of orthopedic impairment does Tawann have? What muscles are affected? What activities (eating, moving, picking up small objects, communicating, etc.) are affected? How can I support Tawann in those activities?
- What is the best way to hold Tawann so he is supported and comfortable?
- What is the best way to move him from place to place?
- What is the best way to position Tawann?
- Can I help with any physical therapy routines?
Understand Each Child’s Needs

Severe and Profound Disabilities

Children with severe and profound disabilities usually need extensive help with self-care. They may have more than one disability, such as having mental retardation together with seizures, cerebral palsy, or troubling behaviors.

Questions to Ask:
- What kinds of activities does Arthur enjoy? How do I make my time with him fun?
- How do I help Arthur with feeding? Toileting? Washing?

Speech Impairment/Language Disorder

Children with speech impairments have difficulties producing some sounds or have trouble with the way that their voices sound. Some people with speech impairments get stuck on sounds and say them over and over. Some people may replace sounds with other sounds. For example, they might say thay instead of say. Some speak too loudly or keep their voices at a very high pitch. A language disorder may be diagnosed when a person has trouble using words to communicate clearly or understanding what is said to him, although he can hear.

Questions to Ask:
- How does Julia communicate?
- What is the best way to communicate with Julia?
Understand Each Child’s Needs

- Does Julia see a speech therapist? Can I help with any speech therapy routines?

Traumatic Brain Injury (TBI)

TBI occurs when a person gets a sharp blow to the head or an object goes into the brain. Symptoms of people with TBI will vary according to what parts of the brain were injured. Children with TBI show a variety of cognitive, physical, social, emotional, and adaptive delays. When you work with children with TBI, it will be important for you to learn from individual families how their children have been affected and how you can help.

Questions to Ask:
- How is Darius affected by his TBI?
- How can I help?

Special Health Care Needs and Chronic Illness

Like children with other special needs, children with special health care needs are a diverse group. Even children with the same diagnosis can be very different. Working with children with special health care needs can be a very different experience from working with children with other disabilities. Some children might require special diets or feeding procedures. Other children may need machines to maintain or to monitor their heartbeat and their breathing.
Allergies

Some children need to avoid certain substances because they are allergic to them. The substances that children are allergic to are called allergens. Children might be allergic to specific foods, insect bites or stings, or animals. They might be allergic to certain types of material. For example, some children are allergic to the latex in some plastic gloves and in rubber balloons. Some children are allergic to pollen in the air or smoke. Children who have allergies react differently when they come in contact with their allergens.

Questions to Ask:

- What is Josie allergic to?
- What do I need to do to prevent exposing her to that allergen?
- How does Josie react when she is exposed to the allergen?
- How can I help her if she is exposed?

Asthma

Children who have asthma have difficulty breathing at times. An asthma attack may include gasping for air, shallow breathing, wheezing, dry coughing, and/or vomiting. Allergens generally trigger attacks. They are more frequent in cold weather and may occur after exercise.
Understand Each Child's Needs

Questions to Ask:
- What causes Jouaquin's asthma attacks? Can I prevent him from being exposed to that? How?
- What do Jouaquin's asthma attacks look like?
- How do I help Jouaquin when he is having an asthma attack? Does he take medicine? How does he take it?
- How long do Jouaquin's attacks usually last? Should I get concerned if they last beyond a certain time? What should I do if they last that long?

Cancer

Cancer is a group of abnormal cells that multiply uncontrollably. Sometimes, these cells can travel through the body and affect different body organs. Cancer affects children differently depending on the parts of the body where the cancer has grown. Speak with families about their children's symptoms and how you can help them have a good time in respite. Be aware that children with cancer may experience a lot of pain. They may be weak and tire easily. Plan plenty of rest periods in your play. Children with cancer may feel nauseous at mealtimes. Talk with families about special diets for their children and special medication routines.

Questions to Ask:
- How can I make Micah comfortable during respite?
- When you are playing with Micah, does he tell you that he is tired? If not, how will I know when Micah needs to rest?
Understand Each Child’s Needs

- Does Micah take medications? What is his medication routine? What side effects should I be aware of?
- Does Micah eat a special diet? What foods should he not have? Do some foods sit better in his stomach than others? What are they?
- What activities does Micah enjoy the most?

Congenital Heart Disease

Children with congenital heart disease have trouble with their hearts from birth. Their hearts may not beat regularly, they may stop beating at times, or they are not structured the way they need to be to do their jobs. Children with congenital heart disease may have had surgery or may need surgery in the future. Sometimes, they use heart monitors to track their heartbeats. As a caregiver for these children, you might need to learn how to use these machines and what to do if the alarm goes off.

Questions to Ask:
- How does Sridhar’s heart disease affect him? What activities does he enjoy? What should we avoid doing?
- Does Sridhar use a heart monitor? How does it work? How do I know if there is a problem?
- How should I respond if there is a problem?
Understand Each Child’s Needs

Cystic Fibrosis

Cystic fibrosis is a condition that causes the lungs and other organs in the body to produce a lot of mucus. This mucus can build up in the child’s system and affect the way that the organs function. Children with cystic fibrosis may cough a lot as they try to clear their lungs of mucus. They generally need special diets and medications to help them process their foods. They may also need their caregivers to give them physical therapy at mealtimes to help them clear their lungs.

Questions to Ask:

- What is Joanna’s medication routine? What is the most comfortable way for me to give her medicine?
- Does Joanna ever refuse her medicine? What do I do if she refuses the medicine from me?
- Do I need to give Joanna chest PT (physical therapy)? How do I do that? When do I do that?

Diabetes

Children with diabetes do not process sugar the way that other people do. Children with diabetes follow special diets and may have exercise built into their routines. They may need to have their blood sugar tested during care and often use insulin injections to regulate blood sugar levels. Caregivers will need to monitor children for signs of high or low blood sugar.
Understand Each Child’s Needs

Questions to Ask:

• How does Matt’s diet work? Does Matt ever refuse the food that he needs to eat? What should I do if he refuses food?
• How do you monitor Matt’s blood sugar levels? Does he take blood tests? How do I help him do that? When should he test his blood?
• Does Matt take insulin? How do I help him take his medication? When does he take it?
• What does it look like when Matt has low blood sugar? What do I need to do if he does? What does it look like when he has high blood sugar? What do I need to do if he does?

HIV/AIDS

HIV, the virus that causes AIDS, attacks the immune system. Children with AIDS are vulnerable to infections. They may have other health conditions for which they will need special care. For example, children with AIDS may have respiratory or other infections, chronic diarrhea, and growth and other developmental delays. Many people are afraid to work with children with HIV/AIDS although it is transmitted only by an exchange of bodily fluids. It is important for caregivers to protect and educate themselves by learning the facts about this disease and how it is transmitted. Section 9 includes Universal Precautions Guidelines that will help protect you from infections and disease.
Understand Each Child’s Needs

Questions to Ask:

- Do I need to take any special precautions when caring for Martha?
- How is HIV/AIDS affecting Martha?
- How can I best support her in her development?
- Does Martha take medication? What is her medication routine? Are there side effects that I need to know?
- Does Martha eat a special diet? What foods should I avoid feeding her?

Kidney Disease

The kidneys clean waste out of blood. Children who have kidney disease have trouble with the way that their kidneys function. They might have trouble with urination. Children with kidney disease may need to watch their diets and drink plenty of water. Children with kidney disease may go to a hospital on a regular basis for dialysis, a treatment through which waste is cleaned from the blood by a machine.

Questions to Ask:

- How does kidney disease affect Michael? What activities does he enjoy? What activities should I avoid doing with him?
- Does Michael take medication? What is his medication routine?
Understand Each Child's Needs

- What foods should Michael avoid eating? What else should I know about his diet?
- How often should Michael drink fluid? What fluids do you want me to give him?

Seizure Disorders

A person who has a seizure temporarily loses control over parts of his or her body. Seizures happen when there is a sudden change in the way that the brain is functioning. There are several types of seizures. Some children who have seizures have a change of consciousness for a short period. Others show brief, marked changes in movement or activity. Children with epilepsy have repeated seizures. Children with seizure disorders often take medication to control their seizures and may eat special diets. In general, people can help a person who is having a seizure by helping him lie down, preferably on his side. Clear the area of anything that he might get hurt on during the seizure.

**DO NOT TRY TO HOLD ONTO THE PERSON OR TRY TO CONTAIN HIM DURING THE SEIZURE!**

Questions to Ask:

- How do I know when Jerry is having a seizure?
- Is there anything that you know of that triggers seizures for Jerry? Are there any activities that we should avoid?
- How do I help Jerry when he is having a seizure?
How Special Needs Affect Development

- How long do Jerry's seizures typically last? When should I call for more help?
- How does Jerry feel when the seizure is over? How can I help him recover?
- Does Jerry take medication? What is his medication routine?
- Does Jerry eat a special diet? What foods should he avoid?

How Special Needs Affect Development

An awareness of how special needs affect development is a good starting place for caring for a child with a special need. Different conditions affect development in different ways. When working with children with special needs, you can use this information to plan activities for them. Remember that some children with special needs, especially children with health concerns, have spent some of their childhoods sick, in hospitals, or otherwise confined. These children may not have had many of the experiences that encourage development. Work with families to plan activities that encourage development in their children.

- Children with speech or language impairments might
  - have greater receptive (understanding) than expressive (speaking) language skills
  - substitute one sound for another such as wabbit for rabbit
  - omit sounds, such as saying han for hand
How Special Needs Affect Development

- distort sounds such as saying thay for say
- not use many words
- speak in short sentences
- have trouble understanding what others are saying

- **Children with mental retardation might**
  - have a hard time following directions
  - not pay attention for a very long time
  - need extra help with transitions
  - have a hard time remembering things
  - need help applying what they learned

- **Children with learning disabilities might**
  - have difficulty in processing information
  - have difficulty in recalling information
  - have difficulty with coordination
  - Children with motor disabilities might
  - walk or move in an unusual way
  - be very clumsy
  - fall or stumble very easily, especially on uneven ground
  - have trouble controlling certain body parts
  - need special equipment like crutches, braces, or a wheelchair
  - not be able to grasp toys securely

- **Children with hearing impairments might**
  - not talk very much
  - leave out certain sounds when they speak
  - talk in a flat sounding voice without expression
  - misunderstand what you say
  - have a short attention span and be easily distracted
How Special Needs Affect Development

- not turn toward the sound of voices
- watch your mouth very closely when you talk

- **Children with visual impairments might**
  - blink, squint, or rub their eyes quite often
  - hold books or other objects very close to their faces
  - cover one of their eyes or tilt their heads in an unusual way
  - complain of headaches, dizziness, itchy eyes, or fuzzy vision
  - trip over small objects

- **Children who have behavioral, social, or emotional disabilities might**
  - be aggressive
  - withdraw from other children and not talk or play with the group
  - have extreme outbursts of crying, tantrums, or yelling
  - have unusual behaviors that they often display, such as waving their hands in front of their eyes or making noises
  - be impulsive
  - have a short attention span

**Adapted from:**
High-Risk Signs

The following list includes a sample of behaviors that may indicate that a child is having problems. You might be the first to notice an unusual behavior. By telling families about the things you notice in a supportive way, you give them valuable information about their children. Remember that any child may demonstrate some of these behaviors and not be having difficulty. However, becoming familiar with these high-risk signs will help you be more aware of the special needs of children.

**Vision**
- Squints, tilts, or turns head to look at angles
- Closes one eye
- Often bumps into things or falls on the playground
- Frequently rubs eyes or complains that eyes hurt
- Has red, watering, or encrusted eyelids
- Sometimes or always crosses one or both eyes
- Frequently holds books or other objects very close to eyes

**Hearing**
- Does not turn toward you when you call his or her name softly
- Shows marked reaction to sudden noises
- Attends to music or stories for less time than other children
- Has unusually unclear speech compared to other children
High Risk Signs

- Uses gestures rather than words to communicate after age 2
- Has frequent earaches or fluid running from ears
- Talks in a very loud or very soft voice
- Always turns the same ear toward a sound
- Does not respond when you call from another room

- Motor
  - One side of the body is more coordinated than the other
  - Hands are shaky or clumsy when doing fine motor tasks
  - Cannot balance on one foot for a short time by age 4
  - Cannot cut with scissors by age 4
  - Cannot throw a ball overhand or catch a large ball by age 5

- Social-Emotional

  Interactions with adults:
  - Has an unusual number of attention-getting behaviors

  Interactions with other children:
  - Avoids certain children
  - Latches onto one particular child
  - Disrupts other children's play often
  - Does not play in group games or pretend play by age 4
  - Does not share or take turns by age 5
Individual behaviors:
- Displays self-abusive behaviors (hurts herself)
- Displays self-stimulatory behaviors (rocking, spinning, picking)
- Overreacts to unexpected stimuli (loud noises, sirens)
- Reacts unusually strongly to the feel of certain substances or textures
- Seems uncomfortable being touched by others
- Has toileting problems at least once a week by age 4

Language
- Does not show interest in books or listening to stories
- Does not follow simple verbal directions
- Is not talking at all by age 2
- Echoes questions or other phrases after age 3
- Is not talking in short sentences by age 4
- Is not understood by people outside the family by age 4

Cognitive
- Cannot answer simple questions about a story by age 3
- Does not give reasonable answers to such questions as What do you do when you are sleepy? or What do you do when you are hungry? by age 4
- Does not demonstrate simple problem-solving skills such as completing puzzles or following patterns for bead stringing by age 4
Finding Additional Information

- Does not seem to understand the meaning of the words *today*, *tomorrow*, and *yesterday* by age 5
- Does not demonstrate memory skills such as repeating short lists of numbers or telling what's missing from a group of objects by age 5

*Adapted from:*

Where to Find More Information on Special Needs

There is an incredible range of disabilities and special health needs. Families and caregivers can use the following agencies in their communities to gather resources on specific special needs.

- Local early intervention programs
- Public school, preschool, or special education programs
- Doctors, nurses, therapists, social workers, and teachers
- Local, college, and university libraries
- Parent Resource Centers
- Parent to Parent Networks
- Local hospitals
Finding Additional Information

- Parent Training Information Centers
- National Information Center for Children and Youth with Disabilities (NICHCY), call toll-free 1 (800) 695-0285
- *Exceptional Parent: Parenting Your Child or Young Adult with a Disability* (magazine)
- National Health Information Center, call toll-free 1 (800) 336-4797
- Internet (accessible through many “on-line” services)
Notes on this Section
Section 8

becoming a
caregiver
Congratulations!

You have learned some basic information for providing respite care to children with special needs. The next step is to participate in personalized training with a family. In this section, we have included sample interview questions families may use to help them choose a caregiver. Maybe you have already made a match with a family and are ready to begin specific training!

As you talk and train with families, be active and open. Ask questions! Find out if this will be a good match for you. If you are not comfortable about working with a child or family, address that early. Families want caregivers to let them know if they cannot work with their children.

What Can I Contribute As a Respite Caregiver?

Your life experiences, education, training, and openness to learning can be valuable to families looking for a respite caregiver. Think about your strengths and use this worksheet to write down what you have to offer. You might want to tell families about these strengths when interviewing with them.
I want to provide respite care because...


I have these skills to offer...


I have these life experiences that will help me care for children with disabilities and build successful relationships with their families...
Sample Interview Questions

You may be asked several of these questions when you interview with families:

- Tell us about your education and/or experience in working with children.
- Discuss your experience working with children with special needs.
- Why are you interested in providing respite care services?
- Describe your strengths. How would you describe yourself?
- Why are you the right person for the job?
- What are your thoughts about families who have children with special needs?
- What would you do if we disagreed about something?
- If my child had to be taken to the hospital for an emergency, what steps would you take?
- How do you believe children's difficult behaviors should be managed?
- What would you do if a child deliberately hit another child?
- What would you do if you found a child with an open bottle you suspect is poison?
Training Designed for You

- Are you willing to take my child out on activities?
- Describe a difficult problem you've had to handle with a child. How did you handle it?
- Tell me about any special training or experience you might have.
- How would you know if my child likes a particular food or activity if he or she is not yet verbal?
- Are you willing to have a criminal background check?
- Do you have CPR, First Aid, and medication assistance training?
- Do you have a valid driver's license?

Personalized Training

In the Partners workshop, families learn about and are prepared for personalized training. Families are given a structure to help them train caregivers. In the next few pages, we have included information on that structure.

Think about ways that you can help plan your personalized training. Different people learn differently and will have different questions for families. The information on the following pages will help you share your preferences, thoughts, and feelings about training with families. Through sharing information, the training will be a positive learning experience for everyone.
Developing a Personalized Training Plan

The Partners Plus model is based on the belief that families know best how to care for their own children and that they have the skill and knowledge needed to train their own respite caregivers. The family and caregiver together decide the specific amount of individual training needed to provide for a child. Training time will vary according to child, family, and caregiver needs.

There are four simple steps to help a family develop a personalized training plan.

**Step 1: Identify training goals.**

*What will a family want a caregiver to learn about their child and family routines?*

**Step 2: List training methods.**

*What methods will a family use to teach their caregiver?*

**Step 3: Use resources to help train your caregiver.**

*What information will a family use to help their caregiver learn?*

**Step 4: Evaluation**

*How will a family and caregiver know when the caregiver has learned a specific skill or routine?*
Sample Personalized Training Plan

Child’s Name: Maria  Date: May 10, 1997
Routine: Lunchtime  Time of Day: Noon - 1:15 p.m.

Step 1: Training Goal
My respite caregiver will be able to feed Maria lunch.

Step 2: Training Methods
My caregiver will watch lunchtime feeding.
My caregiver will assist in feeding Maria.
My caregiver will feed Maria with some assistance.
I will watch my caregiver feed Maria.

Step 3: Resources
I can give my caregiver:
  — a handout on feeding techniques developed by Maria’s occupational therapist.
  — a handout on special feeding problems in children with cerebral palsy.
  — a video of lunchtime routine taped by us at home.

Step 4: Evaluation
I will ask my caregiver questions about skills and comfort in feeding Maria.
I will watch my caregiver feed Maria two bites.
I will watch my caregiver prepare and feed lunch to Maria.
How I Learn Best

One way to help families plan your training is to think about how you learn best. Families can choose a variety of ways to train. These can include written materials and instructions; print materials; audiotapes or videotapes; oral instructions; or demonstration, practice, and feedback.

How do you learn best? Use this worksheet to check the ways that you learn best. Share this with families when you are developing a training plan.

- Written materials
- Audiotapes
- Videotapes
- Oral instruction
- Demonstration, opinions, and practice
- Other

Partnering with Your Family

Families learn in the Partners workshop that caregivers learn in many different ways. It is helpful to talk with families about how you learn. Before you begin training, talk with families about how you think you will learn their child's routine best. Work with them to develop a training plan that incorporates their teaching style and your learning style.
You may not be sure about how you learn best and may want families to proceed with their training plan. During this process, you may discover that you learn best with certain training techniques or that other techniques are not as effective for you. Families learn that it is important to be flexible and adapt their training plan to best meet everyone's needs.

The following are some techniques families have been taught in group training to help a caregiver learn their child's routine:

- Provide written information.
- Ask caregivers to repeat what they thought they heard.
- Ask caregivers to assist in a routine or activity.
- Encourage caregivers to ask questions.
- Ask caregivers to demonstrate the activity or skill.
- Ask caregivers questions.
- Share personal experiences.
- Encourage caregivers to share their ideas.

**Tips for Learning**

- **Discuss training ideas with families.**
  
  If you have identified areas of training that you think will help you in providing care, then share your ideas.
Tips for Learning

- **Give feedback.**
  
  If a teaching method did not help you learn, be sure to tell the family so they can try another way.

- **Write down questions or important facts.**
  
  Bring a notepad with you when you work with a family and their children. You may note something and want to refer back to it or share it later with the family.

- **Ask questions.**
  
  If you do not understand something, then ask for more information.

- **Share new information that you discover with families.**
  
  Families will appreciate knowing you are learning more on your own.

Training Ideas

A major part of successful training is the ability to identify what you want to include in a training plan. Once you match with a family, you may still have concerns about being adequately trained. Use this worksheet to identify others areas of training you might need. These areas may be related to your need for additional knowledge, skills, or comfort in caring for the child or working with the family. Share this information when you meet with a family to help them know other areas to include in your training.
Essential Information for Caregivers

Knowledge
What more do you need to know about working with this child?

Skills
What do you need to learn to do to work with this child?

Comfort
What do you need to increase your comfort level in working with this child?

Essential Information for Caregivers

There is essential information that a caregiver will want to feel comfortable in providing respite care. Be sure to talk with a family about these issues before completion of your individual training. Use this as a checklist to make sure you have the necessary information before providing respite care.

- Emergency phone numbers
- Emergency plan
- Medication schedule
- Home safety issues
- How to use equipment
- Household routines
- Household rules
- Information about siblings
Section 9
resources
Defining Universal Precautions

Resources

The following pages contain valuable information and may be used as a resource for families and caregivers.

Universal Precautions Guidelines

In order to prevent disease, it is important to practice good personal hygiene and to take some precautions, especially when working with children. This is sometimes referred to as universal precautions.

The Occupational Safety and Health Administration (OSHA) has established methods people can use to avoid the transmission of infections and disease. By practicing the following methods, you can help yourself, your family, and the child you provide respite for stay well.

Infectious diseases can be prevented with planning and preparation. Always practice good personal hygiene and wash your hands often. Washing your hands is the most important defense to avoid contracting or transmitting an infectious disease. Good health and up-to-date immunizations are important. Always be careful and act as if all people you come into contact with are potentially infected. There is no way to determine if a person is infected just by looking at them.
In general, it is good to follow these basic precautions when you provide care:

- Wear disposable gloves when it is possible that you will contact blood or body fluids. This contact could occur directly from the person or indirectly through contact with soiled clothing, diapers, or other articles.
- Remove gloves by turning them inside out beginning at the wrist and pulling them off.
- Throw away damaged gloves.
- Do not clean or reuse disposable gloves.
- Avoid touching any items while you are wearing soiled gloves.
- Change gloves when you touch different people.
- Wear protective coverings, such as a mask, eyewear, and/or a gown, whenever you may contact body fluids that might splash.
- Cover any cuts, scrapes, or skin irritations you may have with clothing or bandages.
- Use breathing devices, such as disposable resuscitation masks and airway devices.
- Avoid needle sticks.
- Perform all procedures in such a way as to avoid splashing, spraying, splattering of body fluids.
Defining Universal Precautions

- Avoid touching your face after exposure to body fluids before you wash your hands.
- Remove soiled clothing as soon as possible.
- Clean and disinfect all areas soiled by blood or other body fluids.
- Use approximately 1/4 cup bleach to 1 gallon of water to clean and disinfect soiled surfaces and other items.
- Clean up spills immediately.
- If spills are mixed with sharp items such as broken glass, use tongs or two pieces of cardboard to pick them up, not your hands.
- Use an absorbent material to clean up a spill. Discard the used absorbent material.

If you think that you have been exposed to an infectious disease, wash the area of contact immediately. Exposure usually involves contact with potentially infected blood or fluids through a needle stick or the mucous membranes (eyes, nose, mouth) or inhaling potentially infected airborne droplets.

Infectious diseases can typically be transmitted in four ways: 1) by direct contact with an infected person, through body fluids or through contaminated needles; 2) by contact with a soiled object; 3) by inhaling air exhaled by an infected person; 4) from a bite from an animal, an insect, or an infected person.
If you are working with a child who has an infectious disease, ask the family how it is transmitted and what you should do to protect yourself. Some infectious diseases that you should be aware of are herpes, meningitis, tuberculosis, hepatitis, and HIV/AIDS. Please refer to the following chart for more information.

### How Diseases Are Transmitted

<table>
<thead>
<tr>
<th>Disease</th>
<th>Signs and Symptoms</th>
<th>Mode of Transmission</th>
<th>Infective Material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herpes</td>
<td>Lesions, general ill feeling, sore throat</td>
<td>Direct contact</td>
<td>Broken skin, mucous membranes</td>
</tr>
<tr>
<td>Meningitis</td>
<td>Respiratory illness, sore throat, nausea, vomiting</td>
<td>Airborne, direct and indirect contact</td>
<td>Food, water, mucus</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td>Weight loss, night sweats, occasional fever, general ill feeling</td>
<td>Airborne, direct and indirect contact</td>
<td>Mucus, broken skin</td>
</tr>
<tr>
<td>Hepatitis</td>
<td>Flu-like, jaundice</td>
<td>Direct and indirect contact</td>
<td>Blood, saliva, semen, feces, food, water, other products</td>
</tr>
<tr>
<td>HIV/AIDS</td>
<td>Fever, night sweats, weight loss, chronic diarrhea, severe fatigue, shortness of breath, swollen lymph nodes, lesions</td>
<td>Direct and indirect contact</td>
<td>Blood, semen, vaginal fluid</td>
</tr>
</tbody>
</table>

*Adapted from:*
Preventing the Spread of Disease

Hand Washing

The most important defense against spreading infectious diseases is thorough hand washing with soap and water. Gloves may not always be necessary or even possible to use when reacting to an emergency, but it is extremely important to wash your hands immediately afterward!

There are particular times that you should always wash your hands. It should be done frequently:

- Before putting on a smock in preparation for working with children
- Before drinking, eating, or smoking
- Before handling clean utensils or equipment
- Before and after handling children's food
- Before and after going to the bathroom
- After contact with body secretions, such as blood, urine, feces, mucus, saliva, or drainage from wounds
- After handling soiled diapers, pads, garments, or equipment
- After caring for any child, especially those with nose, mouth, or ear discharge
- After removing disposable gloves
- After removing smock or shirt when leaving the work area

Adapted from:
California State Department of Education.
## Protocol for Hand Washing

### Essential Steps

1. Remove all jewelry.

2. Wet hands with warm, running water.

3. Apply liquid soap and lather well.

4. Wash hands, using a circular motion and friction, for 15 to 30 seconds.

5. Rinse hands well under warm, running water.

6. Repeat steps 3 through 5.

### Key Points and Precautions

- Jewelry should not be worn when working with students who require repeated physical contact and care. Microorganisms can become lodged in settings or stones of rings.

- Warm water, combined with soap, makes better suds than does cold water. Hot water removes protective oils and will dry skin. Running water is necessary to carry away dirt and debris.

- Liquid soap is preferred to bar soap. Bacteria grow on bar soap and in soap dishes.

- Include front and back surfaces of hands, between fingers and knuckles, around nails, and the entire wrist area. Avoid harsh scrubbing to prevent skin breaks.

- Hold hands under the water so that water drains from wrist area to fingertip.

- All remaining bacteria and soil should now be removed.
Protocol for Hand Washing

7. Wipe surfaces surrounding sink with a clean paper towel and discard.

Damp surfaces promote the growth of bacteria.

8. Dry hands well with paper towels and discard towels immediately.

Because of frequent hand washing, it is important to dry gently and thoroughly to avoid chapping. Chapped skin breaks open, thus permitting bacteria to enter one's system.

9. Apply lotion as desired.

Lotion helps keep skin soft and reduces chapping.

Source:
References on Adult Learning


**Training Resources**

Creative Training Techniques Newsletter
Lakewood Publications
50 S. 9th St.
Minneapolis, MN 55402

(800) 707-7749

http://www.trainingsupersite.com/publications
General Information About
Respite Care

Background

Respite Care programs emerged in the late 1960's as a result of the deinstitutionalization movement with the belief that the best place to care for a child with disabilities was in the child's home and community. Respite care became an essential component in the overall support services that families need to provide home care. Recognizing the effectiveness of respite care services, in 1986, Congress passed the Temporary Child Care for Children with Disabilities and Crisis Nurseries Act (as amended). This Act established federal funding for respite care demonstration projects. Administered through the U.S. Department of Health and Human Services, Children's Bureau, competitive grants have been awarded to States since 1988 to assist private and public agencies in developing model respite care services across the United States.

Purpose

Respite care programs provide temporary relief for families or caregivers. Additional family benefits include: allowing the family to engage in daily activities thus decreasing their feelings of isolation; providing the family with rest and relaxation; improving the family's ability to cope with daily responsibilities; maintaining the family's stability during crisis situations; helping preserve the family unit by lessening the pressures that might lead to divorce, institutionalization, neglect and/or child abuse; and, making it possible for individuals with disabilities to establish individual identities and enrich their own growth and development.

What is Respite Care?

The federal legislation defines respite care as in-home or out-of-home temporary, non-medical child care for families who have children with disabilities, chronic, or terminal illnesses. Most respite care programs offer services to families on a sliding fee scale with hourly and/or daily rates. Respite care services can range from a few hours of care up to three months of care depending on the needs of the families and the type of respite care program model available in a community. The ages served by respite programs range from infancy to adulthood. Often programs serve a particular disability or illness (e.g., children who have HIV-related conditions, children who are medically fragile, children with mental retardation).

Many respite care services are incorporated within larger social service agencies both at the community and/or state levels which provide an array of support services to families. Some of these support services can include: family counseling; family support groups; parent training; service coordination; assistive equipment services; and, access to medical services.

Respite Care Program Models

Respite care program models for each local service area may differ according to the needs of families/caregivers within the community. For example, some respite care programs may utilize an available bed in a health care facility or state institution for families who require extended respite options and whose child requires skilled care; whereas, other respite care programs may only offer time-limited (a few hours) services in the family's home. In addition, respite care services may be available to families through formal programs (i.e., trained staff) or may be available to families through informal networks (e.g., parent cooperatives, or cash subsidies from states to purchase respite through relatives and friends). The following descriptions are examples of local respite care program models.

Model 1: In-Home Respite Program Model

In-home respite programs offer services in the family's home for certain periods of time by trained, paid respite providers or volunteers. Respite programs train providers or volunteers in basic child care areas. Most respite programs interview families and providers and match the two in the provision of respite services; whereas, other respite programs act as "brokers" in which the families choose their own providers from a designated list. In-home respite services provide relief with minimal disruption of routine child care activities. State regulations regarding particular activities (e.g., administration of medications) are followed (over)
by the respite program. In this type of model, families are usually charged on a sliding fee scale with the respite program paying the provider directly. In many states, families receive a cash subsidy or voucher to pay for in-home respite services.

Model 2: Out-of-Home: Host Family Model

Many respite programs offer services through a host family model in which respite care is provided in a surrogate family home similar to a foster care model. This model allows the child with disabilities to be cared for in a setting outside of his/her natural home. Host family models allow families to receive extended periods of respite. State regulations regarding fire and safety codes, licensing and other requirements must be followed. Most host families are trained by the respite program and paid a daily rate of care. Families are charged on a sliding fee scale for this service.

Model 3: Out-of-Home: Facility-based Model

This type of respite care program is similar to a Mother’s Day Out program in which a particular facility (e.g., day care center, church, community center) offers respite on certain days at certain periods of time. This model allows families to take their children to a supervised environment. Facility-based respite models are staffed with trained respite providers, paraprofessionals, or volunteers who are able to care for children with certain disabilities. Families are charged for services based on a sliding fee scale. This model is very effective in rural areas.

Model 4: Out-of-Home: Residential Respite Model

In the residential respite model, respite services are provided in a residential or nursing center (e.g., group home, state institution, nursing home, hospital) within a community for individuals with disabilities or terminal illnesses who require skilled care services. Respite, in this model, is conducted by professional licensed personnel, usually for periods longer than 24 hours. Payment for respite services is usually a combination of family fees, state funding (i.e., Medicaid or waiver services), and/or private insurance.

Model 5: Out-of-Home: Respite House Model

In this type of model, a house, specifically designed for the provision of respite care, is available in the community with 24-hour care. Extended respite periods can occur up to 30 days. The respite house model is usually sponsored by a community or state agency and staffed with professional and paraprofessional staff. Families are charged daily rates based on a sliding fee scale. State requirements for food preparation, staff-child ratio, health and safety, and other licensing requirements must be followed.

Model 6: Parent Cooperative Model

Parent cooperatives have been developed in communities, especially rural areas, where respite services are very limited. In this type of model, families of children with disabilities and/or chronic illnesses develop an informal association and "trade" respite services with each other. This exchange program allows families to receive respite on scheduled dates. In most parent cooperatives, fees are not assessed. This model has proven to be effective for families whose children have similar disabilities. In this regard, families feel comfortable caring for someone else’s child who has a disability or illness similar to their own child’s.

Model 7: Respitality Model

Respitality is an innovative concept for providing respite care. It provides a cost-effective partnership between the private sector and respite care agencies. During Respitality, participating hotels provided the family with a room, a pleasant dining experience, and perhaps entertainment while a local respite program provides respite either in the family’s home or in an out-of-home respite situation. The Respitality concept was developed by United Cerebral Palsy Associations, Inc., in 1985.

References


Resource Centers

Texas Respite Resource Network' (TRRN), Santa Rosa Children's Hospital, P.O. Box 7330, San Antonio, Texas 78207-3198; (512) 228-2794

National Information Center for Children and Youth with Handicaps (NICHCY), P.O. Box 1492, Washington, DC 20013: 1-800-999-5599

This fact sheet was developed by the ARCH National Resource Center for Child Development and Parent Support, and funded by the U.S. Department of Health and Human Services, Administration for Children and Families. This publication does not necessarily reflect the views or policies of the funders, nor does mention of trade names, commercial products or organizations imply endorsement by the U.S. Department of Health and Human Services.
Background
Over the past fifteen years, parents have increasingly made the choice to keep their families together. Children with developmental or physical disabilities are no longer routinely placed in institutional care. As a result, the need for respite care (temporary child care) services has increased. Until recently, respite care has been either unavailable or unaffordable in many states. Personal care through the Title XIX Medicaid program and grants through the "Temporary Child Care for Children with Disabilities and Crisis Nurseries Act of 1986" (as amended), have enabled families to receive respite care for free, or at a reduced, reasonable cost. Forty-six states have some form of respite programs for children with disabilities (Knoll, et al., 1990). States and/or agencies that administer respite care projects have realized the need to provide other supports to families which are flexible and meet the needs of the whole family, not just the person with a disability. Respite care services form the basis for many state family support programs.

Purpose
For years families caring for a child with medical or physical involvements have been reluctant to allow others to provide care for their child in their own home. The two most common reasons stated by parents are: 1) concerns about the caregiver's training; and 2) feelings that no one can take care of their child as well as they can. Babysitters, in general, do not have the skills needed to provide the kind of support required to care for children with medical or physical disabilities. Respite care programs for families who have children with developmental and/or physical disabilities provide trained respite care providers for in-home or out-of-home "free" time (respite) for families.

Respite Programs
Highlighting the strengths and needs of the entire family, and not just the needs of the individual with a disability, has been a common thread in exemplary programs. Providing respite care for children with developmental or physical disabilities is not unlike providing respite for any other child. The respite provider needs to know the kinds of behaviors the child is likely to exhibit and be prepared to deal with any unusual needs of the child. This is true of any child. The behaviors and needs of children with disabilities may be more acute, but ascertaining them should be done in the same manner.

For the sake of those programs considering starting a respite program for families with children with developmental or physical disabilities, the following outline is designed to give an idea of the kinds of training and attitudes that are required to provide a quality program.

The Keys to a Successful Respite Program
- Help families feel comfortable and build trust. Spend enough time with the family to develop a level of trust that will allow a free flow of information.
- Provide a flexible structure. The agency should provide as many options as possible to meet the needs of the family, rather than fitting the family into the service available.
- Be family-focused. Empower families to make decisions for themselves. Discuss the strengths and needs of the whole family.
- Provide support. The administering agency and respite care providers must be available to field questions and concerns of parents that may be outside the realm of respite care. This might include linkage and referrals to other agencies or services.
- Allow families to train providers. Be sure to include families in the process of designing the child-specific portion of the provider’s training.

Training
Because of the concern that families have about the specialized needs of their children, respite providers should be trained in first aid, CPR, disability awareness issues, specific medical conditions, behavior management, and individual "hands-on" training with the child for whom
they will provide care. Many states and local programs have developed respite training manuals which are available to train local providers. (The ARCH Information Center has copies of many of these manuals.)

Building trust is probably the key ingredient in a successful respite program. Training respite workers is only one step in this process. Establishing close working relationships with the family and ensuring close supervision are other important parts of a successful program.

The Basics — These areas can be taught in a classroom setting using professional staff.

- CPR and First Aid
- Philosophy of family empowerment
- “How to speak DD” — introduction to developmental disability terminology
- “Don’t Panic” — introduction to medical and environmental emergencies

The Details — These units should be presented in at least two different forums. The first, as a parent panel with discussion of some personal stories in each of the areas mentioned. The second, as a clinical approach to care, to include a speech therapist, an occupational therapist, and a physical therapist.

- Communication sensitivity — how to communicate with someone who does not talk very well, if at all
- Eating etiquette — good table manners (skills) when working with someone who needs assistance
- The comfort factor, or “how would you feel sitting in the same position all day?”

“Hands On” — This section should take place in the families’ homes, and/or, the setting where the respite will actually occur.

- Meet the family and get to know the child
- Trainee asks parents about specific assistance he or she may need to provide
- Parents demonstrate how they assist their child to move, eat, and go to the bathroom
- Trainee works with the child on each of the specific skills parents request until parents are comfortable in how the trainee performs

Conclusion

Successful programs from around the country report that, once these key requirements are met, families will enthusiastically participate in respite services. Programs also report that this process takes some time. The trust that families need does not occur overnight. Allow a considerable amount of time for the program to be fully operational.

This can be one of the most rewarding programs to operate. Families and staff report many heartwarming tales of the benefits that respite has provided.

References


Resources

Association for the Care of Children’s Health, 3615 Wisconsin Avenue, N.W., Washington, DC 20016. (202) 244-1801. The Association provides information and support on family-centered care for children with special health care needs.

Human Services Research Institute, 2336 Massachusetts Avenue, Cambridge, MA 02140. (617) 876-0426. The Institute has compiled extensive information on respite and other family support services throughout the United States and has numerous documents available.


Exceptional Parent Magazine, P.O. Box 3000, Dept. EP, Denville, NJ 07834. This is an excellent resource for families and professionals on a wide range of issues concerning children with disabilities.

This fact sheet was developed by Scott Miller, Family Support Services Coordinator for West Virginia. Scott is also the parent of Josh, who has Down’s Syndrome.
DISABILITIES WHICH QUALIFY CHILDREN AND YOUTH FOR SPECIAL EDUCATION SERVICES UNDER THE INDIVIDUALS WITH DISABILITIES EDUCATION ACT (IDEA)

The Education of the Handicapped Act, Public Law (P.L.) 94-142, was passed by Congress in 1975 and amended by P.L. 99-457 in 1986 to ensure that children with disabilities would have a free, appropriate public education available to them which would meet their unique needs. It was again amended in 1990, and the name was changed to the Individuals with Disabilities Education Act (P.L. 101-476), or IDEA.

IDEA defines “children with disabilities” as having any of following types of disabilities: autism, deaf-blindness, hearing impairments (including deafness), mental retardation, multiple disabilities, orthopedic impairments, other health impairments, serious emotional disturbance, specific learning disabilities, speech or language impairments, traumatic brain injury, and visual impairments (including blindness). These terms are defined in the regulations for IDEA, as described below.

1. AUTISM
A developmental disability significantly affecting verbal and non-verbal communication and social interaction, generally evident before age three, that adversely affects educational performance.

2. DEAFNESS
A hearing impairment which is so severe that a child is impaired in processing linguistic information through hearing, with or without amplification, which adversely affects educational performance.

3. DEAF-BLINDNESS
Simultaneous hearing and visual impairments, the combination of which causes such severe communication and other developmental and educational problems that a child cannot be accommodated in special education programs solely for children with deafness or children with blindness.

4. HEARING IMPAIRMENT
An impairment in hearing, whether permanent or fluctuating, which adversely affects a child’s educational performance but which is not included under the definition of “deafness.”

5. MENTAL RETARDATION
Significantly subaverage general intellectual functioning existing concurrently with deficits in adaptive behavior and manifested during the developmental period, which adversely affects a child’s educational performance.

6. MULTIPLE DISABILITIES
Simultaneous impairments (such as mental retardation/blindness, mental retardation/orthopedic impairment, etc.), the combination of which causes such severe educational problems that the child cannot be accommodated in a special education program solely for one of the impairments. The term does not include children with deaf-blindness.

7. ORTHOPEDIC IMPAIRMENT
A severe orthopedic impairment which adversely affects a child’s educational performance. The term includes impairments caused by a congenital anomaly (e.g., clubfoot, absence of some limb, etc.), impairments caused by disease (e.g. poliomyelitis, bone tuberculosis, etc.), and impairments from other causes (e.g., cerebral palsy, amputations, and fractures or burns which cause contractures).

8. OTHER HEALTH IMPAIRMENT
Having limited strength, vitality, or alertness, due to chronic or acute health problems such as a heart condition, tuberculosis, rheumatic fever, nephritis, asthma, sickle cell anemia, hemophilia, epilepsy, lead poisoning, leukemia, or diabetes, which adversely affects a child’s educational performance. According to the Office of Special Education and Rehabilitative Services’ clarification statement of September 16, 1991, eligible children with ADD may also be classified under “other health impairment.”

9. SERIOUS EMOTIONAL DISTURBANCE
(1.) A condition exhibiting one or more of the following characteristics over a long period of time and to a marked degree, which adversely affects educational performance:
(A) an inability to learn which cannot be explained by intellectual, sensory, or health factors; (B) an inability to build or maintain satisfactory interpersonal relationships with peers and teachers; (C) inappropriate types of behavior or feelings under normal circumstances; (D) a general pervasive mood of unhappiness or depression; or (E) a tendency to develop physical symptoms or fears associated with personal or school problems. (II) The term includes children who have schizophrenia. The term does not include children who are socially maladjusted, unless it is determined that they have a serious emotional disturbance.

10. SPECIFIC LEARNING DISABILITY
A disorder in one or more of the basic psychological processes involved in understanding or in using language, spoken or written, which may manifest itself in an imperfect ability to listen, think, speak, read, write, spell, or to do mathematical calculations. The term includes such conditions as perceptual disabilities, brain injury, minimal brain dysfunction, dyslexia, and developmental aphasia. The term does not include children who have learning problems which are primarily the result of visual, hearing, or motor disabilities, of mental retardation, of emotional disturbance, or of environmental, cultural, or economic disadvantage.

11. SPEECH OR LANGUAGE IMPAIRMENT
A communication disorder such as stuttering, impaired articulation, a language impairment, or a voice impairment, which adversely affects a child's educational performance.

12. TRAUMATIC BRAIN INJURY
An acquired injury to the brain caused by an external physical force, resulting in total or partial functional disability or psychosocial impairment, or both, which adversely affects educational performance. The term does not include brain injuries that are congenital or degenerative, or brain injuries induced by birth trauma.

13. VISUAL IMPAIRMENT INCLUDING BLINDNESS
A visual impairment which, even with correction, adversely affects a child's educational performance. The term includes both children with partial sight and those with blindness.

SERVICES FOR INFANTS, TODDLERS, AND PRESCHOOLERS WITH DISABILITIES
P.L. 99-457, the Education of the Handicapped Act Amendments of 1986, created a new mandate for all state education agencies to serve all children with disabilities from age three by 1991-1992. The Preschool Program's purpose is to extend the P.L. 94-142 rights to children from age three, including all definitions and requirements. However, Congress made an important distinction for preschoolers: States are not required to label 3-5 year-olds in order to serve these children.

P.L. 99-457 also established the Part H program, now known as the Early Intervention Program for Infants and Toddlers with Disabilities. This program is directed to the needs of children, from birth to their third birthday, who need early intervention services because they: (1) are experiencing developmental delays in one or more of the following areas: cognitive, physical, language and speech, psychosocial, or self-help skills; (2) have a physical or mental condition that has a high probability of resulting in delay, such as Down Syndrome, cerebral palsy, etc.; or (3) at the state's discretion, are at risk medically or environmentally for substantial developmental delays if early intervention is not provided. In addition, under this program the infant or toddler's family may receive services that are needed to help them assist in the development of their child. State definitions of eligibility under this program vary; many states are still in the process of developing their Part H programs. Therefore, depending on the state, services may be fully available or still in the process of developing.

FOR ADDITIONAL INFORMATION
If you feel that any of the above statements accurately describes your child, we encourage you to find out more about special education and related services available in your child's public school district. Many parents have found the NICHCY publication entitled "Questions Parents Often Ask About Special Education Services" helpful. For children birth through 5 years, ask for the publication "A Parents' Guide to Accessing Programs for Infants, Toddlers, and Preschoolers with Disabilities." All NICHCY publications are free of charge.

The Special Education Director for your child's school district, Child Find Coordinator, or the principal of your child's school should be able to answer specific questions you may have about obtaining special education and related services for your child. In addition, the Federally funded Parent Training and Information Programs across the country are excellent sources of information. For a listing of information sources in your state, NICHCY has a State Resource Sheet for each state and U.S. Territory; this sheet includes the address of the Parent Training and Information Program.

This fact sheet is made possible through Cooperative Agreement #H320A1A30003 between the Academy for Educational Development and the Office of Special Education Programs, U.S. Department of Education. The contents of this publication do not necessarily reflect the views of the Department of Education, nor does mention of trade names, commercial products or organizations imply endorsement by the U.S. Government.

This information is copyright free. Readers are encouraged to copy and share it, but please credit the National Information Center for Children and Youth with Disabilities (NICHCY).
The agencies described below provide assistance to people with disabilities and their families. The State Resource Sheet enclosed with this packet will tell you the names and addresses of these agencies in your state. One of the best resources for finding help for your child with a disability is your local school district (sometimes called Local Education Agency). If your child has already begun school and you think your child needs special services, we suggest that you begin by discussing your concerns with your child’s teacher or school principal. If your child is an infant, we suggest that you contact the office listed for Programs for Children with Disabilities: Ages Birth through 2 Years. If your child is of preschool age, we suggest that you contact the office listed for Programs for Children with Disabilities: Ages 3 Through 5 Years. These are listed on the State Resource Sheet for your state. Ask for the Child Find Coordinator in your community.

STATE EDUCATION DEPARTMENT

The State Department staff can answer questions about special education and related services in your state. Many states have special manuals explaining the steps to take. Check to see if one is available. State Department officials are responsible for special education and related services programs in their state for preschool, elementary and secondary age children.

OFFICE OF STATE COORDINATOR OF VOCATIONAL EDUCATION FOR STUDENTS WITH DISABILITIES

States receiving Federal funds used for vocational education must assure that funding is used in programs which include students with disabilities. This office can tell you how your state funds are being used and provide you with information on current programs.

STATE VOCATIONAL REHABILITATION AGENCY

The state vocational rehabilitation agency provides medical, therapeutic, counseling, education, training and other services needed to prepare people with disabilities for work. This state agency will provide you with the address of the nearest rehabilitation office where you can discuss issues of eligibility and services with a counselor. The state vocational rehabilitation agency can also refer you to an independent living program in your state. Independent living programs provide services which enable adults with disabilities to live productively as members of their communities. The services might include, but are not limited to, information and referral, peer counseling, workshops, attendant care, and technical assistance.

STATE MENTAL RETARDATION/DEVELOPMENTAL DISABILITIES AGENCIES

The functions of state mental retardation/developmental disabilities agencies vary from state to state. The general purpose of this office is to plan, administer and develop standards for state/local mental retardation/developmental disabilities programs provided in state-operated facilities and community-based programs. This office provides information about available services to families, consumers, educators and other professionals.
STATE DEVELOPMENTAL DISABILITIES COUNCIL

Assisted by the U.S. Department of Health and Human Services' Administration on Developmental Disabilities, state councils plan and advocate for improvement in services for people with developmental disabilities. In addition, funding is made available for time-limited demonstration and stimulatory grant projects.

STATE MENTAL HEALTH AGENCIES

The functions of state mental health agencies vary from state to state. The general purposes of these offices are to plan, administer, and develop standards for state and local mental health programs such as state hospitals and community health centers. They can provide information to the consumer about mental illness and a resource list of contacts where you can go for help.

PROTECTION AND ADVOCACY AGENCY AND CLIENT ASSISTANCE PROGRAM

Protection and Advocacy systems are responsible for pursuing legal, administrative and other remedies to protect the rights of people who are developmentally disabled or mentally ill, regardless of their age. Protection and Advocacy agencies may provide information about health, residential, and social service in your area. Legal assistance is also available.

The Client Assistance Program provides assistance to individuals seeking and receiving vocational rehabilitation services. These services, provided under the Rehabilitation Act of 1973, include assisting in the pursuit of legal, administrative, and other appropriate remedies to insure the protection of the rights of individuals with developmental disabilities.

PROGRAMS FOR CHILDREN WITH SPECIAL HEALTH CARE NEEDS

The U.S. Department of Health and Human Services’ Office of Maternal and Child Health and Resource Development provides grants to states for direct medical and related services to children with handicapping conditions. Although services will vary from state to state, additional programs may be funded for training, research, special projects, genetic disease testing, and counseling services. For additional information about current grants and programs in your state, contact the National Center for Education in Maternal and Child Health, 8201 Greensboro Dr., Suite 600, McLean, VA 22102. The telephone number is (703) 821-8955.

UNIVERSITY AFFILIATED PROGRAMS

A national network of programs affiliated with universities and teaching hospitals, UAPs provide interdisciplinary training for professionals and paraprofessionals and offer programs and services for children with disabilities and their families. Some UAPs provide direct services for children and families. Individual UAPs have staff with expertise in a variety of areas and can provide information, technical assistance, and inservice training to agencies, service providers, parent groups, and others.

You can obtain a listing of all University Affiliated Programs by contacting The Maternal and Child Health Clearinghouse, 8201 Greensboro Dr., Suite 600, McLean, VA 22102. The telephone number is (703) 821-8955. Additional information about UAPs may be obtained by contacting:

American Association of University Affiliated Programs for Persons with Developmental Disabilities (AAUAP)
8630 Fenton Street, Suite 410
Silver Spring, MD 20910
(301) 588-8252

For more information contact NICHCY
**NATIONAL RESOURCES**

<table>
<thead>
<tr>
<th>CLEARINGHOUSES</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Clearinghouse on Disability Information</strong></td>
<td>National Health Information Center</td>
</tr>
<tr>
<td>Office of Special Education and Rehabilitative Services</td>
<td>P.O. Box 1133</td>
</tr>
<tr>
<td>Room 3132, Switzer Building</td>
<td>Washington, D.C. 20013-1133</td>
</tr>
<tr>
<td>330 C Street S.W.</td>
<td>(301) 565-4167</td>
</tr>
<tr>
<td>Washington, DC 20202-2524</td>
<td>(800) 336-4797</td>
</tr>
<tr>
<td>(202) 205-8241 (Voice/TT)</td>
<td></td>
</tr>
<tr>
<td><strong>DB-LINK</strong></td>
<td>National Information Center on Deafness (NICD)</td>
</tr>
<tr>
<td>National Information Clearinghouse on Children Who Are Deaf-Blind</td>
<td>800 Florida Avenue, N.E.</td>
</tr>
<tr>
<td>345 N. Monmouth Avenue</td>
<td>Washington, D.C. 20002</td>
</tr>
<tr>
<td>Monmouth, OR 97361</td>
<td>(202) 651-5051 (Voice)</td>
</tr>
<tr>
<td>(800) 438-9376</td>
<td>(202) 651-5052 (TT)</td>
</tr>
<tr>
<td>(800) 854-7013 (TTY)</td>
<td>Publications available in Spanish</td>
</tr>
<tr>
<td><strong>ERIC Clearinghouse on Disabilities and Gifted Education</strong></td>
<td></td>
</tr>
<tr>
<td>Council for Exceptional Children (CEC)</td>
<td>National Information Clearinghouse (NIC) for Infants with Disabilities and Life-Threatening Conditions</td>
</tr>
<tr>
<td>1920 Association Drive</td>
<td>Center for Developmental Disabilities</td>
</tr>
<tr>
<td>Reston, VA 22091-1589</td>
<td>University of South Carolina</td>
</tr>
<tr>
<td>(703) 620-3660</td>
<td>Columbia, SC 29208</td>
</tr>
<tr>
<td>(800) 328-0272</td>
<td>(800) 922-9234, ext. 201 (Voice/TT)</td>
</tr>
<tr>
<td><strong>HEATH Resource Center (National Clearinghouse on Postsecondary Education</strong></td>
<td>National Information System for Vietnam Veterans and Their Families</td>
</tr>
<tr>
<td>for Individuals with Disabilities)</td>
<td>Center for Developmental Disabilities</td>
</tr>
<tr>
<td>One Dupont Circle, N.W., Suite 800</td>
<td>University of South Carolina</td>
</tr>
<tr>
<td>Washington, D.C. 20036-1193</td>
<td>Columbia, SC 29208</td>
</tr>
<tr>
<td>(202) 939-9320; (800) 544-3284 (Voice/TT)</td>
<td>(800) 922-9234, ext. 401 (Voice/TT)</td>
</tr>
<tr>
<td><strong>National Clearinghouse for Professions in Special Education</strong></td>
<td></td>
</tr>
<tr>
<td>Council for Exceptional Children</td>
<td>National Maternal and Child Health Clearinghouse</td>
</tr>
<tr>
<td>1920 Association Drive</td>
<td>8201 Greensboro Dr., Suite 600</td>
</tr>
<tr>
<td>Reston, VA 22091-1589</td>
<td>McLean, VA 22102</td>
</tr>
<tr>
<td>(703) 264-9474</td>
<td>(703) 821-8955 Ext. 254 or 265</td>
</tr>
<tr>
<td>(203) 746-6518</td>
<td>Publications available in Spanish</td>
</tr>
<tr>
<td>(203) 746-6927 (TT)</td>
<td>Spanish speaker on staff</td>
</tr>
<tr>
<td><strong>National Clearinghouse on Family Support and Children's Mental Health</strong></td>
<td></td>
</tr>
<tr>
<td>Portland State University</td>
<td>National Organization for Rare Disorders (NORD)</td>
</tr>
<tr>
<td>P.O. Box 751</td>
<td>100 Route-37</td>
</tr>
<tr>
<td>Portland, OR 97207-0751</td>
<td>P.O. Box 8923</td>
</tr>
<tr>
<td>(503) 725-4040</td>
<td>New Fairfield, CT 06812-1783</td>
</tr>
<tr>
<td>(503) 725-4165 (TT)</td>
<td>(800) 999-6673</td>
</tr>
<tr>
<td><strong>Spanish speaker on staff</strong></td>
<td>(203) 746-6518</td>
</tr>
<tr>
<td><strong>Spanish speaker on staff</strong></td>
<td>(203) 746-6927 (TT)</td>
</tr>
</tbody>
</table>
### ORGANIZATIONS

<table>
<thead>
<tr>
<th>Organization</th>
<th>Address</th>
<th>City, State, Zip</th>
<th>Phone Numbers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alliance for Technology Access</td>
<td>2173 East Francisco Boulevard, Suite L</td>
<td>San Rafael, CA 94901</td>
<td>(415) 455-4575</td>
</tr>
<tr>
<td>Alliance of Genetic Support Groups</td>
<td>35 Wisconsin Circle, Suite 440</td>
<td>Chevy Chase, MD 20815</td>
<td>(800) 336-4363, (301) 652-5553</td>
</tr>
<tr>
<td>American Council of Rural Special Education (ACRES)</td>
<td>Department of Special Education</td>
<td>University of Utah</td>
<td>Milton Bennion Hall, Room 221</td>
</tr>
<tr>
<td>American Foundation for the Blind (AFB)</td>
<td>11 Penn Plaza, Suite 300</td>
<td>New York, NY 10001</td>
<td>(800) 232-5463 (Voice), (212) 502-7600 (Voice), (212) 502-7662 (TT)</td>
</tr>
<tr>
<td>American Occupational Therapy Association (AOTA)</td>
<td>4720 Montgomery Lane</td>
<td>Bethesda, MD 20824-1220</td>
<td>(301) 652-2682, (800) 377-8555 (TT)</td>
</tr>
<tr>
<td>American Physical Therapy Association (APTA)</td>
<td>1111 North Fairfax Street</td>
<td>Alexandria, VA 22314</td>
<td>(703) 684-2782, (800) 999-2782</td>
</tr>
<tr>
<td>American Speech-Language-Hearing Association (ASHA)</td>
<td>10801 Rockville Pike</td>
<td>Rockville, MD 20852</td>
<td>(301) 897-5700 (Voice/TT), (800) 638-8255</td>
</tr>
<tr>
<td>Association for the Advancement of Rehabilitation Technology (RESNA)</td>
<td>1700 N. Moore Street, Suite 1540</td>
<td>Arlington, VA 22209-1903</td>
<td>(703) 524-6686 (Voice), (703) 524-6639 (TT)</td>
</tr>
<tr>
<td>Association for the Care of Children’s Health (ACCH)</td>
<td>7910 Woodmont Avenue, Suite 300</td>
<td>Bethesda, MD 20814-3015</td>
<td>(301) 654-6549</td>
</tr>
<tr>
<td>Association for Persons with Severe Handicaps (TASH)</td>
<td>29 W. Susquehanna Ave., Suite 210</td>
<td>Baltimore, MD 21204</td>
<td>(410) 828-8274, Spanish speaker on staff</td>
</tr>
<tr>
<td>The Arc (formerly the Association for Retarded Citizens of the U.S.)</td>
<td>500 E. Border Street, Suite 300</td>
<td>Arlington, TX 76010</td>
<td>(817) 261-6003, (817) 277-0553 (TT), (800) 433-5255</td>
</tr>
<tr>
<td>Autism Society of America (formerly NSAC)</td>
<td>7910 Woodmont Avenue, Suite 650</td>
<td>Bethesda, MD 20814</td>
<td>(301) 657-0881, (800) 3-AUTISM</td>
</tr>
<tr>
<td>Fact sheet available in Spanish</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Council for Exceptional Children (CEC)</td>
<td>1920 Association Drive</td>
<td>Reston, VA 22091</td>
<td>(703) 620-3660</td>
</tr>
<tr>
<td>Disability Statistics Research and Training Center Institute for Health and Aging</td>
<td>University of California, San Francisco</td>
<td>San Francisco, CA 94143-0646</td>
<td>(415) 502-5210 (Voice/TT)</td>
</tr>
<tr>
<td>Epilepsy Foundation of America (EFA)</td>
<td>4351 Garden City Drive, 5th Floor</td>
<td>Landover, MD 20785</td>
<td>(301) 459-3700, (800) 332-1000</td>
</tr>
<tr>
<td>Publications available in Spanish, Spanish speaker on staff</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Family Resource Center on Disabilities</td>
<td>20 East Jackson Boulevard, Room 900</td>
<td>Chicago, IL 60604</td>
<td>(800) 952-4199 (in IL only), (312) 939-3513, (312) 939-3519 (TT)</td>
</tr>
</tbody>
</table>
Head Start Bureau
Administration on Children, Youth and Families
U.S. Department of Health & Human Services
P.O. Box 1182
Washington, D.C. 20013
(202) 205-8347

Independent Living Research Utilization Project (ILRU)
The Institute for Rehabilitation and Research
2323 South Sheppard, Suite 1000
Houston, TX 77019
(713) 520-0232
(713) 520-5136 (TT)

International Rett Syndrome Association
9121 Piscataway Rd., Suite 2B
Clinton, MD 20735
(301) 856-3334
Publication "What is Rhett Syndrome?" available in Spanish

Learning Disability Association of America (LDA)
(formerly ACLD)
4156 Library Road
Pittsburgh, PA 15234
(412) 341-1515
(412) 341-8077
Publications available in Spanish

March of Dimes Birth Defects Foundation
1275 Mamaroneck Avenue
White Plains, NY 10605
(914) 428-7100
Publications available in Spanish
Spanish speaker on staff

Muscular Dystrophy Association (MDA)
3300 East Sunrise Drive
Tucson, AZ 85718
(602) 529-2000
Publications available in Spanish
Spanish speaker on staff

National Alliance for the Mentally Ill (NAMI)
200 N. Glebe Rd., Suite 1015
Arlington, VA 22203-3754
(703) 524-7600
(800) 950-NAMI

National Association of Private Schools for Exceptional Children (NAPSEC)
1522 K Street N.W., Suite 1032
Washington, DC 20005
(202) 408-3338

National Association of Protection and Advocacy Systems (NAPAS)
900 Second Street N.E., Suite 211
Washington, DC 20002
(202) 408-9514 (Voice)
(202) 408-9521 (TT)

National Down Syndrome Congress
1605 Chantilly Drive, Suite 250
Atlanta, GA 30324
(800) 232-6372 (Toll-free)
(404) 633-1555
Pamphlet available in Spanish
Spanish speaker on staff

National Down Syndrome Society
666 Broadway
New York, NY 10012
(212) 460-9330
(800) 221-4602
Publications available in Spanish
Spanish speaker on staff

National Easter Seal Society, Inc.
230 West Monroe Street, Suite 1800
Chicago, IL 60606
(312) 726-6200
(312) 726-4258 (TT)
(800) 221-6827
Publications available in Spanish
Spanish speaker on staff

National Fragile X Foundation
1441 York Street, Suite 303
Denver, CO 80206
(303) 333-6155
(800) 688-8765

National Head Injury Foundation, Inc.
1776 Massachusetts Ave., N.W., Suite 100
Washington, D.C. 20036
(202) 296-6443
Publications available in Spanish

National Library Services for the Blind & Physically Handicapped
The Library of Congress
Washington, D.C. 20542
(202) 707-5100
(800) 424-8567
(800) 424-9100 (TT, English)
(800) 345-8901 (TT, Spanish)
### NATIONAL TOFF FREE NUMBERS

The following is a selected list of toll-free numbers for national organizations concerned with disability and children's issues. Inclusion on this list does not imply endorsement by NICHCY or the Office of Special Education Programs. There are also many national disability organizations providing services and information which do not have toll-free numbers. If you would like additional help in locating assistance contact NICHCY at 1-800-695-0285 (Voice/TT).

*Note: Telephone numbers are designated either Voice (V) or Text Telephone (TT), indicating their accessibility to TT users. Spanish-language resources are also indicated.*

#### AIDS
- **CDC National AIDS Clearinghouse**
  - 1(800) 458-5231 (V; English/Spanish)
  - 1(800) 243-7012 (TT)
- **CDC National AIDS Hotline**
  - 1(800) 342-2437 (V)
  - 1(800) 344-7432 (V; Spanish)
  - 1(800) 243-7889 (TT)
- **Hemophilia and AIDS/HIV Network for Dissemination of Information**
  - 1(800) 424-2634 (V)

#### ALCOHOL AND OTHER DRUG ABUSE
- **National Institute on Drug Abuse Helpline**
  - 1(800) 662-4357 (V)
- **OSAP National Clearinghouse for Alcohol and Drug Information**
  - 1(800) 729-6686 (V)
  - 1(800) 487-4889 (TT)

#### AMERICANS WITH DISABILITIES ACT (ADA)
- **Disability Rights Education and Defense Fund**
  - ADA Technical Assistance Information Line
  - 1(800) 466-4232 (V/TT)
- **Equal Employment Opportunity Commission**
  - 1(800) 669-3362 (V)
  - 1(800) 800-3302 (TT)

#### ASSISTIVE TECHNOLOGY/DEVICES
- **AbleNet**
  - 1(800) 322-0956 (V)
- **Apple Office for Special Education Material**
  - 1(800) 732-3131, ext. 950 (V)
- **AT&T Accessible Communications Product Center**
  - 1(800) 233-1222 (V)
  - 1(800) 833-3232 (TT)
- **IBM Special Needs Information Referral Center**
  - 1(800) 426-4832 (V)
  - 1(800) 284-4833 (TT)
- **TECHKNOWLEDGE**
  - 1(800) 726-9119
  - (404) 894-4960 (V; Atlanta Metro Area)

---

**NATIONAL INFORMATION CENTER FOR CHILDREN AND YOUTH WITH DISABILITIES (NICHCY)**

P.O. Box 1492, Washington, D.C. 20013-1492
1-800-695-0285 (Toll Free, Voice/TT)

SpecialNet User Name: NICHCY ** Internet: nichcy@capcon.net**
BLINDNESS/VISUAL IMPAIRMENTS
American Council of the Blind
1(800) 424-8666 (V/TT)

American Foundation for the Blind
1(800) 232-5463 (V)

Blind Children's Center
1(800) 222-3566 (V)
1(800) 222-3567 (V; in CA)

Hadley School for the Blind
1(800) 323-4238 (V)

Job Opportunities for the Blind
1(800) 638-7518 (V)
(410) 659-9314 (V; in MD)

Lighthouse National Center for Vision and Child Development
1(800) 334-5497 (V)
(212) 808-5544 (TT)

National Association of Parents of the Visually Impaired
1(800) 562-6265 (V)

National Society to Prevent Blindness
1(800) 331-2020 (V)

Recording for the Blind
1(800) 221-4792 (V)

Retinitis Pigmentosa Foundation
1(800) 683-5555 (V)
(410) 225-9400 (V; in MD)
(410) 225-9409 (TT)

BURNS
Phoenix Society
1(800) 888-2876 (V)

CANCER
Cancer Information and Counseling Line
1(800) 525-3777 (V)

Candlelighters Childhood Cancer Foundation
1(800) 366-2223 (V)

National Cancer Information Service
1(800) 422-6237 (V; English/ Spanish)

CHILD ABUSE
Clearinghouse on Child Abuse and Neglect/Family Violence Information
1(800) 394-3366 (V)

National Resource Center on Child Sexual Abuse
1(800) 543-7006 (V)

COMMUNICATION DISORDERS
National Institute on Deafness and Other Communication Disorders Clearinghouse
1(800) 241-1044 (V)
1(800) 241-1055 (TT)

CRANIOFACIAL SYNDROMES
Children's Craniofacial Association
1(800) 535-3643 (V)

FACES — National Association for the Craniofacially Handicapped
1(800) 332-2373 (V)

National Foundation for Facial Reconstruction
1(800) 422-3223

DEAFNESS/HEARING IMPAIRMENTS
American Society for Deaf Children
1(800) 942-2732 (V/TT)

Better Hearing Institute
1(800) 327-9355 (V/TT)

Deafness Research Foundation
1(800) 535-3323 (V/TT)
(212) 684-6559 (V/TT; in NY)

Hear Now
1(800) 648-4327 (V/TT)

John Tracy Clinic
1(800) 522-4582 (V/TT)
(213) 748-5481 (V; in 213 area)
(213) 747-2924 (TT; in 213 area)

National Hearing Aid Society
1(800) 521-5247 (V)

National Institute on Deafness and Other Communication Disorders Clearinghouse
1(800) 241-1044 (V)
1(800) 241-1055 (TT)
| Medical/Health Disorders | | | |
|--------------------------|--------|------------------|
| American Association of Kidney Patients | 1(800) 749-2257 (V) | National Clearinghouse on Family Support and Children's Mental Health | 1(800) 628-1696 (V) |
| American Brain Tumor Association | 1(800) 886-2282 (V) | National Mental Health Association | 1(800) 969-6642 (V) |
| American Diabetes Association | 1(800) 582-8323 (V) | MENTAL RETARDATION | |
| American Kidney Fund | 1(800) 638-8299 (V) | American Association on Mental Retardation | 1(800) 424-3688 (V) |
| American Liver Foundation | 1(800) 223-0179 (V) | The Arc | 1(800) 433-5255 (V) |
| American Lupus Society | 1(800) 331-1802 (V) | NUTRITION | |
| Asthma and Allergy Foundation of America | 1(800) 727-8462 (V) | Beech-Nut Nutrition Hotline | 1(800) 523-6633 (V) |
| Chronic Fatigue and Immune Dysfunction Syndrome Association | 1(800) 442-3437 (V) | Gerber Consumer Information | 1(800) 443-7237 (V) |
| Federal Hill-Burton Free Hospital Care Program | 1(800) 638-0742 (V) | PHYSICAL DISABILITIES | |
| | 1(800) 492-0359 (V; in MD) | American Paralysis Association | 1(800) 526-3456 (V) |
| | | Human Growth Foundation | 1(800) 451-6434 (V) |
| Leukemia Society of America | 1(800) 955-4572 (V) | Physically Challenged Resource Center | 1(800) 255-9877 (V) |
| Lupus Foundation of America | 1(800) 558-0121 (V) | RARE SYNDROMES | |
| | 1(800) 558-0231 (V; Spanish) | Alliance of Genetic Support Groups | 1(800) 336-4363 (V) |
| National Association for Sickle Cell Disease | 1(800) 421-8453 (V) | National Information Center on Orphan Drugs and Rare Diseases | 1(800) 456-3505 (V) |
| Shriners Hospital for Crippled Children | 1(800) 237-5055 (V) | National Organization for Rare Disorders | 1(800) 999-6673 (V/TT) |
| | 1(800) 282-9161 (V; in FL) | RECREATION | |
| United Ostomy Association | 1(800) 826-0826 (V) | Adventures in Movement for the Handicapped, Inc. | 1(800) 332-8210 (V) |
| MENTAL HEALTH | | Magic Foundation | 1(800) 362-4423 (V) |
TRIPOD (Information for parents of deaf children)
1(800) 352-8888 (V/TT)
1(800) 346-8888 (V/TT; in CA)

DISABILITY AWARENESS
Kids on the Block
1(800) 368-5437

EDUCATION
American Association for Vocational Instructional Materials
1(800) 228-4689 (V)

Association for Childhood Education International
1(800) 423-3563 (V)

HEALTH Resource Center
1(800) 544-3284 (V/TT)
(202) 939-9320 (V/TT; in DC)

National Center for Research in Vocational Education
1(800) 762-4093 (V)

National Center for School Leadership
1(800) 643-3205 (V)

National Committee for Citizens in Education Clearinghouse
1(800) 638-9675 (V)
1(800) 532-9832 (V; Spanish)

U.S. Office of Educational Research and Improvement
1(800) 424-1616 (V)

EMPLOYMENT
Equal Employment Opportunity Commission
1(800) 669-3362 (V)
1(800) 800-3302 (TT)

Job Accommodation Network
1(800) 526-7234 (V/TT)
1(800) 232-9675 (V/TT; ADA Information)

Job Opportunities for the Blind
1(800) 638-7518 (V)
(410) 659-9314 (V; in MD)

FINANCIAL COUNSELING
National Foundation for Consumer Credit
1(800) 388-2227 (V)

HOSPICE
Children's Hospice International
1(800) 242-4453 (V/TT)

HOSPICELINK
1(800) 331-1620
(203) 767-1620 (V; in CT)

INFORMATION SERVICES
ABLEDATA/National Rehabilitation Information Clearinghouse
1(800) 346-2742 (V/TT)

ACCESS ERIC
1(800) 538-3742 (V)

BRS Information Technologies
1(800) 289-4277 (V)

National Center for Youth with Disabilities
1(800) 333-6293 (V)
(612) 624-3939 (TT)

National Easter Seal Society
1(800) 221-6827 (V)
(312) 726-4258 (TT)

National Information Clearinghouse for Infants with Disabilities and Life Threatening Conditions
1(800) 922-9234, ext. 201 (V/TT)
1(800) 922-1107, ext. 201 (V/TT; in SC)

ODPHP National Health Information Center
1(800) 336-4797 (V)

Office of Minority Health Resource Center
1(800) 444-6472 (V)

LITERACY
National Literacy Hotline
1(800) 228-8813 (V)
1(800) 552-9097 (TT)
North American Riding for the Handicapped, Inc.  
1(800) 369-7433 (V)

Sunshine Foundation  
1(800) 767-1976 (V)

REHABILITATION  
Clearinghouse for Rehabilitation and Technology Information  
1(800) 638-8864 (V)  
1(800) 852-2892 (TT)

National Clearinghouse of Rehabilitation Training Materials  
1(800) 223-5219 (V/TT)

National Rehabilitation Information Clearinghouse/ABLEDATA  
1(800) 346-2742 (V/TT)

RESPIRATORY DISORDERS  
National Jewish Center for Immunology and Respiratory Medicine – LUNGLINE  
1(800) 222-5864 (V)

RESPITE CARE  
Access to Respite Care and Help (ARCH) National Resource Center  
1(800) 473-1727 (V)

RURAL  
ERIC Clearinghouse on Rural Education and Small Schools  
1(800) 624-9120 (V)  
1(800) 344-6646 (V; in WV)

Rural Institute on Disabilities (Montana University Affiliated Program)  
1(800) 732-0323 (V/TT)

SPECIFIC DISABILITIES  
Attention Deficit Disorder Association  
1(800) 487-2282 (V)

Cleft Palate Foundation  
1(800) 242-5338 (V)

Cooley's Anemia Foundation  
1(800) 221-3571 (V)  
1(800) 522-7222 (V; in NY)

Cornelia de Lange Syndrome Foundation  
1(800) 223-8355 (V)  
1(800) 753-2357 (V; in CT)

Cystic Fibrosis Foundation  
1(800) 344-4823 (V)

Epilepsy Foundation of America  
1(800) 332-1000 (V)

Little People of America  
1(800) 243-9273 (V)

National Down Syndrome Congress  
1(800) 232-6372 (V)

National Down Syndrome Society  
1(800) 221-4602 (V)

National Fragile X Foundation  
1(800) 688-8765 (V)

National Multiple Sclerosis Society  
1(800) 532-7667 (V)

National Organization for Albinism and Hypopigmentation  
1(800) 473-2310 (V)

National Reye's Syndrome Foundation  
1(800) 233-7393 (V)

National Tuberous Sclerosis Association  
1(800) 225-6872 (V)

Orton Dyslexia Society  
1(800) 222-3123 (V)

Prader-Willi Syndrome Association  
1(800) 926-4797 (V)

Spina Bifida Associations of America  
1(800) 621-3141 (V)

Stuttering Foundation of America  
1(800) 992-9392 (V)

Sudden Infant Death Syndrome Alliance  
1(800) 221-7437 (V)
Tourette Syndrome Association
1(800) 237-0717 (V)

United Cerebral Palsy Associations
1(800) 872-5827 (V/TT)

United Leukodystrophy Foundation
1(800) 728-5483 (V)

United Scleroderma Foundation
1(800) 722-4673 (V)

SUPPLEMENTAL SECURITY INCOME (SSI)
Social Security Administration
1(800) 772-1213 (V)
1(800) 325-0778 (Tr)
1(800) 392-0812 (TT; in MO)

Zebley Implementation Project
1(800) 523-0000 (V)
(215) 893-5356 (V; Philadelphia Metro Area)

TRAUMA
American Trauma Society
1(800) 556-7890 (V)

National Head Injury Foundation
1(800) 444-6443 (V)

National Spinal Cord Injury Association
1(800) 962-9629 (V)

VIETNAM VETERANS/AGENT ORANGE
Access Group for Children of Vietnam Veterans
1(800) 821-8580 (V)

National Information System for Vietnam Veterans and
Their Families
1(800) 922-9234, ext. 401 (V/TT)
1(800) 922-1107, ext. 401 (V/TT; in SC)

BEST COPY AVAILABLE

This list is made possible through Cooperative Agreement #H030A30003
between the Academy for Educational Development and the Office of
Special Education Programs, U.S. Department of Education. The contents
of this publication do not necessarily reflect the views or policies of the
Department of Education, nor does mention of trade names, commercial
products, or organizations imply endorsement by the U.S. Government.

This information is copyright free. Readers are encouraged to copy and
share it, but please credit the National Information Center for Children and
Youth with Disabilities (NICHCY).

For more information contact NICHCY.

UPDATE 6/94
**PUBLICATIONS LIST 1995**

Single copies of NICHCY materials are FREE. To enable us to provide materials to as many people as possible, please request in writing only those materials you must need at this time. Allow 3-4 weeks for delivery.

Permission to duplicate NICHCY materials is not required. In fact, we encourage duplication; however, please credit NICHCY as the source of the material.

<table>
<thead>
<tr>
<th>Send information in:</th>
<th>English</th>
<th>Spanish</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>GENERAL RESOURCE</strong></td>
<td>FS6</td>
<td>Epilepsy *</td>
</tr>
<tr>
<td>GR1 Brochure *</td>
<td>FS7</td>
<td>Learning Disabilities *</td>
</tr>
<tr>
<td>GR2 National Resources</td>
<td>FS8</td>
<td>Mental Retardation *</td>
</tr>
<tr>
<td>GR3 General Information about Disabilities *</td>
<td>FS10</td>
<td>Severe and/or Multiple Disabilities</td>
</tr>
<tr>
<td>GR4 Public Agencies Fact Sheet</td>
<td>FS11</td>
<td>Speech and Language Impairments</td>
</tr>
<tr>
<td>GR5 National Toll Free Numbers</td>
<td>FS12</td>
<td>Spina Bifida *</td>
</tr>
<tr>
<td>GR6 State Resource Sheet: State</td>
<td>FS13</td>
<td>Visual Impairments *</td>
</tr>
<tr>
<td>GR8 Publications List</td>
<td>FS14</td>
<td>Attention Deficit Disorder (Briefing Paper)</td>
</tr>
<tr>
<td></td>
<td>FS17</td>
<td>Reading and Learning Disabilities Resource Guide</td>
</tr>
<tr>
<td></td>
<td>FS18</td>
<td>Traumatic Brain Injury *</td>
</tr>
</tbody>
</table>

**NEWS DIGEST**

| ND11 | Children with Disabilities: Understanding Sibling Issues |
| ND12 | Respite Care: A Gift of Time |
| ND13 | Assistive Technology: Becoming an Informed Consumer |
| ND14 | Having a Daughter with a Disability: Is it Different for Girls? |
| ND18 | Estate Planning (Vol. II, No. 1, 1992) |
| ND21 | Questions & Answers about the Individuals with Disabilities Education Act (Vol. III, No. 2, 1993)* |
| ND23 | *Assessing Children for the Presence of a Disability* (Vol. IV, No. 1, 1994) |

**TRANSITION SUMMARY**

| TS8 | Transition Services in the IEP |

**DISABILITY INFORMATION**

| FS1 | Autism * |
| FS2 | Cerebral Palsy * |
| FS3 | Deafness * |
| FS4 | Down Syndrome * |
| FS5 | Emotional Disturbance * |

**MATERIALS FOR PARENTS**

| PA2 | Parent's Guide to Accessing Programs for Infants, Toddlers, Preschoolers w/Disabilities (ages 0-5) * |
| PA6 | Parent's Guide: Accessing the ERIC Resource Collection |
| PA7 | Parent's Guide to Doctors, Disabilities, and the Family |
| PA8 | Parent's Guide: Planning a Move, Mapping Your Strategy |
| PA10 | Parent's Guide: Accessing Parent Groups* |

**BIBLIOGRAPHIES**

| B1 | Assessing Children for the Presence of a Disability: Schools |
| B2 | Assessing Children for the Presence of a Disability: Families |
| B3 | Behavior Management: Schools |
| B4 | Behavior Management: Families |
| B5 | Children's Literature |
| B6 | Mental Health/Mental Illness: Schools |
| B7 | Mental Health/Mental Illness: Families |

**OTHER**

| PAY | Paying the Medical Bills |
| ADT | Resources for Adults with Disabilities |
| LG1 | Questions Often Asked About Special Education Services (ages 3-21) * |
| LG2 | Individualized Education Programs (IEPs) |
| TA1 | Technical Assistance Guide: Operating a Local Information and Referral Center |
Please complete the following:

What age ranges are you interested in? (Please circle all ages that apply)

   0-2  3-5  6-12  13-18
   19-22  23-over  All ages

Have you contacted NICHCY before?
__ Yes __ No

Check the one that best describes you:
__ Parent/Family Member
__ Educational Professional
__ Other Professional
__ Advocate
__ Adult with Disabilities
__ Student with Disabilities
__ Student without Disabilities
__ Parent Organization/Group
__ Library/Information Center
__ Other (Please specify ___________)

PLEASE TYPE OR PRINT CLEARLY.

First Name _______________ Last Name _______________

Organization (for mailing list purposes only)

Address ________________________________

City ___________ State ___________ Zip Code _______

Daytime Telephone (Area Code + Number) ________________

The National Information Center for Children and Youth with Disabilities (NICHCY) is a national information and referral clearinghouse. We operate through the Clearinghouse Program authorized by Section 633 of Part D of the Individuals with Disabilities Education Act (20 U.S.C. 1433), as amended by Public Law 101-476.

AED

Academy for Educational Development

NICHCY is operated by the Academy for Educational Development, pursuant to Cooperative Agreement #H030A30003 with the Office of Special Education Programs. The contents of this publication do not necessarily reflect the views or policies of the Department of Education, nor does mention of trade names, commercial products, or organizations imply endorsement by the U.S. Government.
Section 10

forms for sharing information
Sharing Information with Caregivers

Information Worksheets

There are a variety of forms available in this section for families to use in sharing important information with caregivers. Families can use these forms as needed. Some forms provide information specific to the child with special needs and other forms are specific to the needs of siblings also in care. Some forms provide space to include other family and household information. Emergency, health, and safety information sheets and checklists are helpful for caregivers to have.

Master copies of all forms and worksheets are included in this section. Families can make copies of forms that several caregivers may need. Families may also keep blank copies and use them when changes occur or new information becomes available.
Essential Information

General Information

Child's Name: ____________________  Nickname: ____________________

Address: _______________________________________________________

Phone: ____________________  Allergies: ____________________

Age: ______  D.O.B. ______  Height: ______  Weight: ______

Child diagnosis or description of special needs: ____________________

______________________________________________________________

Health Insurance: ____________________  I.D. Number: ____________________

You can reach me at ______________________________________________

If you cannot contact me, please call:

______________________________________________________________  Phone: ____________________

Medication

Name of medication: ____________________

Purpose for medication: ____________________

Time to be given: ______  Dosage: ____________________

Pharmacy: ____________________  Phone: ____________________

Comments: ___________________________________________________

______________________________________________________________

______________________________________________________________

______________________________________________________________
Essential Information (continued)

Food

Allergies: ________________________________

Cannot have: ______________________________

Acceptable snacks: _________________________

Meals: ________________________________

Daily Schedule

AM: ________________________________

PM: ________________________________

Nap Time: ________________________________
Health and Medical Information

Child's Name: ___________________ Date of Birth: ___________________

Current Health Problems / Needs: ______________________________________

______________________________________________________________

Allergies (if any): ________________________________________________

______________________________________________________________

Child's Physician: _________________________________________________

Address: _______________________________________________________

Phone: __________________________________________________________

Name and Number of Medical Insurance: _____________________________

Authorization for Emergency Medical Care

_________________________ has permission to obtain immediate medical

care for my child if any emergency occurs when the parent or guardian

cannot be located. I understand the child's physician will be contacted

if available.

Parent or Guardian: ________________ Date: ___________________
Additional information about my child:

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________
## Important Phone Numbers

<table>
<thead>
<tr>
<th>Category</th>
<th>Phone Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mother's Work</td>
<td></td>
</tr>
<tr>
<td>Father's Work</td>
<td></td>
</tr>
<tr>
<td>Police/Emergency</td>
<td></td>
</tr>
<tr>
<td>Fire/Emergency</td>
<td></td>
</tr>
<tr>
<td>Poison Control</td>
<td></td>
</tr>
<tr>
<td>Doctor's Office</td>
<td></td>
</tr>
<tr>
<td>Pharmacy</td>
<td></td>
</tr>
<tr>
<td>Hospital</td>
<td></td>
</tr>
<tr>
<td>Dentist</td>
<td></td>
</tr>
<tr>
<td>Neighbor/Friend</td>
<td></td>
</tr>
<tr>
<td>Neighbor/Friend</td>
<td></td>
</tr>
<tr>
<td>Relative</td>
<td></td>
</tr>
<tr>
<td>Relative</td>
<td></td>
</tr>
<tr>
<td>Gas Company</td>
<td></td>
</tr>
<tr>
<td>Electric Company</td>
<td></td>
</tr>
<tr>
<td>Phone Company</td>
<td></td>
</tr>
<tr>
<td>Apartment Manager</td>
<td></td>
</tr>
<tr>
<td>Veterinarian</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
</tr>
</tbody>
</table>
Other information I want you to know:

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________

________________________________________________________________________
About My Child

Child's Name: _____________________________ Date: ________

What I would like to tell you about my child: _____________________________

_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________

Favorite toys or playthings: _____________________________

_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________

Special routines (a good-bye hug, snack after school, story for bedtime, etc.)

my child enjoys: _____________________________

_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________

Special things (blanket, a pacifier, stuffed animal, etc.) that my child enjoys:

_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________

Things that scare my child: _____________________________

_____________________________________________________________________
_____________________________________________________________________
_____________________________________________________________________
Things that quiet or comfort my child: ____________________________________________

__________________________________________________________________________

Special things my child does: ________________________________________________

__________________________________________________________________________

Activities my child likes: ____________________________________________________

__________________________________________________________________________

Activities my child dislikes: _________________________________________________

__________________________________________________________________________

Personality or behavior style (such as active, quiet, silly, serious): __________

__________________________________________________________________________

Things that are easy for my child: ____________________________________________

__________________________________________________________________________

Things that are hard for my child: ____________________________________________

__________________________________________________________________________

Special names or words for people or things that my child understands/uses:

__________________________________________________________________________

__________________________________________________________________________

Other special information or concerns: ________________________________________

__________________________________________________________________________
About Our Family

The people in our family are: ____________________________________________
______________________________________________________________________
______________________________________________________________________
______________________________________________________________________

Some things our family does together: _____________________________________
______________________________________________________________________
______________________________________________________________________
______________________________________________________________________

Special things each member of my family does alone or with someone else:
______________________________________________________________________
______________________________________________________________________
______________________________________________________________________
______________________________________________________________________

Things we would like to do if we had respite: _______________________________
______________________________________________________________________
______________________________________________________________________
______________________________________________________________________
Dreams for our child with disabilities: ____________________

__________________________

__________________________

__________________________

__________________________

__________________________

Dreams for our family: ____________________

__________________________

__________________________

__________________________

__________________________
About Sisters and Brothers

General Information

Brother/Sister's Name: ____________________________ Nickname: ____________________________

Age: _______ D.O.B.: _______ Height: _______ Weight: _______

Allergies: ____________________________________________________________________________

Important health or developmental history: ____________________________________________________________________________

_____________________________________________________________________________________

_____________________________________________________________________________________

_____________________________________________________________________________________

_____________________________________________________________________________________

Medications

Name of medication: __________________________________________________________________

Purpose for medication: __________________________________________________________________

Time to be given: ____________________________ Dosage: ____________________________

Pharmacy: ____________________________ Phone: ____________________________

Prescribing Doctor: ___________________________________________________________________

Doctor's Phone: ______________________________________________________________________

Let me tell you about __________________________________________________________________

_____________________________________________________________________________________

_____________________________________________________________________________________

_____________________________________________________________________________________

_____________________________________________________________________________________

Partners Plus Caregiver Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Let me tell you about


Health and Safety Checklist

Our caregiver is certified in:

___ infant/child CPR
___ first aid

Our caregiver knows where we keep our:

___ medical treatment release forms
___ medical insurance information
___ first-aid kit and book
___ sunscreen

We have provided our caregiver individualized training on:

___ emergency plans
    ___ what we would consider an emergency for our child
    ___ what we would like our caregiver to do in case of an emergency
    ___ where we keep emergency phone numbers

___ fire safety
    ___ location of and how to use the fire extinguisher
    ___ our fire evacuation plan
    ___ location of smoke detector

___ poison safety
    ___ where we keep the Poison Control Center phone number
    ___ where we keep the Syrup of Ipecac

___ kitchen safety (i.e., stove, appliances, pots on the stove, etc.)
    ___ first aid for burns
We have provided our caregiver individualized training on:

- hand washing for caregivers and children after toileting and outside time and before food preparation
- medication administration
- our smoking policy
- diaper/toileting routine and diaper disposal
- toy safety (e.g., balloons might suffocate small children, young children might choke on small toys or toys with small parts)
- use of medical equipment
- crib/bed safety
  - young children should sleep on their backs
  - the side rail of a crib should be latched when the child is asleep
  - caregivers should check on sleeping children
  - bottles should be held and not propped for young children
- positioning, exercises, and therapeutic activities
- car safety
  - car seats are available and adjusted for our child
  - correct use of a car seat
  - child should always be in a car seat or fastened seat belt when the car moving
- my child’s allergies
  - medical care if my child is exposed to allergens
- safety in the home
  - electrical outlets are covered
  - hot water heater adjusted to 120 degrees or lower
  - guns are unloaded and locked in a cabinet
  - bullets are locked in cabinets separate from guns
  - medications, poisonous chemicals, and liquor are kept out of the reach of children
- rubber mats or nonslip surfaces are in showers or tubs children will use
Step 1: Identify training goals.

What do you want your caregiver to learn about your child and family routines?

I want my caregiver to learn about these routines:

Goal 1: __________________________________________________________

_________________________________________________________________

_________________________________________________________________

Goal 2: __________________________________________________________

_________________________________________________________________

_________________________________________________________________

Goal 3: __________________________________________________________

_________________________________________________________________

_________________________________________________________________

Goal 4: __________________________________________________________

_________________________________________________________________

_________________________________________________________________

Goal 5: __________________________________________________________

_________________________________________________________________
Step 2: List training methods.

What methods will you use to teach your caregivers?
In order to reach these goals I will use the following methods:

Goal 1:

Methods:
1. 
2. 
3. 

Goal 2:

Methods:
1. 
2. 
3. 
Goal 3: _______________________________________________________________________

Methods:
1. _______________________________________________________________________
2. _______________________________________________________________________
3. _______________________________________________________________________

Goal 4: _______________________________________________________________________

Methods:
1. _______________________________________________________________________
2. _______________________________________________________________________
3. _______________________________________________________________________

Goal 5: _______________________________________________________________________

Methods:
1. _______________________________________________________________________
2. _______________________________________________________________________
3. _______________________________________________________________________

Partners Plus Caregiver Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Step 3: Use resources to help train your caregiver.

What information will you use to help your caregiver learn?

These are the resources I would like to use:

- [ ] books
- [ ] handouts
- [ ] videotapes
- [ ] audiotapes
- [ ] computer search
- [ ] I develop resources
- [ ] other (please describe)

-----------------------------

-----------------------------

-----------------------------
Step 4: Evaluation

How will you and your caregiver know when a specific skill or routine has been learned?

I will know a specific skill or routine has been learned when we accomplish the following:

Goal 1: __________________________________________________________

______________________________________________________________

Goal 2: __________________________________________________________

______________________________________________________________

Goal 3: __________________________________________________________

______________________________________________________________

Goal 4: __________________________________________________________

______________________________________________________________

Goal 5: __________________________________________________________

______________________________________________________________
Personalized Training Plan

Child's Name: ___________________________ Date: __________

Routine: ____________________________________________________________

____________________________________________________________________

Time of Day: ______________

Step 1: Training Goal

____________________________________________________________________

____________________________________________________________________

Step 2: Training Methods

____________________________________________________________________

____________________________________________________________________

Step 3: Resources

____________________________________________________________________

____________________________________________________________________

Step 4: Evaluation

____________________________________________________________________

____________________________________________________________________
Respite Caregiver Profile

Date: __________

Personal:

Name: ___________________________ SS#: ___________________________

Address: __________________________________________________________

Home Telephone: _______________ Work Telephone: _______________

In case of emergency, notify: _________________________________________

Relationship: ______________________________________________________

Home Telephone: _______________ Work Telephone: _______________

Experience:

Describe experience you have had with children (including your own). __________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

Have you ever provided care for children with disabilities?

___ Yes  ___ No

If yes, please list your experiences: ________________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________

____________________________________________________________________
Respite Caregiver Profile, continued

Have you had any courses in early childhood, child development, or child health?
___ Yes ___ No

Are you certified in Red Cross First Aid?
___ Yes ___ No Date of Expiration

Are you certified in Infant/Child CPR?
___ Yes ___ No Date of Expiration

Do you have a valid driver’s license?
___ Yes ___ No Date of Expiration

Education:
I have completed:
High School __________________ Some College __________________
College __________________ Graduate School __________________
Degree or Major (specify): ______________________________________
Other training/education: ______________________________________

Employment:
Current Employer: __________________
Position: __________________
Respite Caregiver Profile, continued

Interests: (Check all that apply)

I am willing to care for children ages

birth - 2 ________ 2 - 5 ________ 5 - 8 ________ over 8 ________

I am willing to provide care (check all that apply):

____ at my home ______ weekdays
____ at child's home ______ weekends
____ during the day ______ overnight
____ during evenings ______ anytime

When I provide care, I will care for:

____ only the child with disabilities
____ the child with disabilities plus siblings
____ the child with disabilities and other children in my care
____ the child with disabilities and my own child/children

I would prefer to provide care:

____ as a volunteer
____ in exchange for a service (i.e., transportation, lawn care, meals, etc.)
____ for an hourly rate of $ _________
____ for a negotiable hourly rate

How far are you willing to travel to provide care? (Check all that apply.)

____ Williamsburg/James City County ______ Hampton
____ York County ______ Poquoson
____ Newport News ______ Gloucester
Respite Caregiver Profile, continued

References:

Please provide the names, addresses, and phone numbers of two persons we may contact who have known you for more than 1 year (excluding relatives or roommates).

Name: ___________________________ Phone: ___________________________
Address: ___________________________

Name: ___________________________ Phone: ___________________________
Address: ___________________________

I am most interested in providing respite care because ___________________________

__________________________________________________________________________

Thank You!
Reference Check

Name of Potential Caregiver: ________________________________

Name of Reference: _______________  Phone: ______________________

1. How do you know ________________________________?  
   (put name of potential caregiver here.)

2. How long have you known ________________________________? 

3. What words best describe ________________________________? 

4. How would you best describe how __________________________ relates to children?

5. Describe what you think are ____________________________’s strong points in working with children with special needs and their families?

6. Are there any other comments about __________________________ you would like to make?
Final Activity
Step 1: Identify training goals.

What do you want your caregiver to learn about your child and family routines?

I want my caregiver to learn about these routines:

Goal 1: 

Goal 2: 

Goal 3: 

Goal 4: 

Goal 5: 

Partners Plus Caregiver Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

1056 DRAFT 10.99
Step 2: List training methods.

What methods will you use to teach your caregivers?

In order to reach these goals I will use the following methods:

Goal 1: ____________________________________________________________

Methods:

1. ________________________________________________________________
2. ________________________________________________________________
3. ________________________________________________________________

Goal 2: ____________________________________________________________

Methods:

1. ________________________________________________________________
2. ________________________________________________________________
3. ________________________________________________________________
Goal 3: ________________________________

Methods:

1. ________________________________

2. ________________________________

3. ________________________________

Goal 4: ________________________________

Methods:

1. ________________________________

2. ________________________________

3. ________________________________

Goal 5: ________________________________

Methods:

1. ________________________________

2. ________________________________

3. ________________________________
Step 3: Use resources to help train your caregiver.

What information will you use to help your caregiver learn?

These are the resources I would like to use:

- books
- handouts
- videotapes
- audiotapes
- computer search
- I develop resources
- other (please describe)

---

Partners Plus Caregiver Manual
Child Development Resources
P.O. Box 280, Norge, Virginia 23127

DRAFT 10.99
Step 4: Evaluation

How will you and your caregiver know when a specific skill or routine has been learned?

I will know a specific skill or routine has been learned when we accomplish the following:

Goal 1:

Goal 2:

Goal 3:

Goal 4:

Goal 5:
NOTICE

Reproduction Basis

This document is covered by a signed "Reproduction Release (Blanket)" form (on file within the ERIC system), encompassing all or classes of documents from its source organization and, therefore, does not require a "Specific Document" Release form.

This document is Federally-funded, or carries its own permission to reproduce, or is otherwise in the public domain and, therefore, may be reproduced by ERIC without a signed Reproduction Release form (either "Specific Document" or "Blanket").