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RESEARCH ARTICLE

PROPOSAL LEARNING STRATEGY INTERFACIAL POLYMERIZATION OF NYLON 6,10

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ARTICLE INFO ABSTRACT Polymer Chemistry is a subject in the Chemistry Programs, student have a natural curiosity about Article History: polymeric materials, that we use in our everyday activities. Our proposal is to apply cooperative Received 05th September, 2021 learning in the Interfacial polymerization of Nylon 6,10 experimentation. The specific objectives of Received in revised form our project can be summarized as: To identify the origin of the raw materials used for industrial 19th October, 2021 Accepted 30th November, 2021 Published online 29th December, 2021 Keywords Learning Strategy, Interfacial Polymerization, Nylon, Complementary Activities.

*Corresponding author: Adolfo Eduardo Obaya Valdivia synthesis, to become acquainted with the most important processes used for the synthesis, To identify the most important variables related with the control of the polymerization process, To correlate the physical chemical properties with the final applications of this material. The reader will find in the text a series of questions that will make him reflect -so-called complementary activitiespromoting the development of competences complementary ones based on research exercises. Through to synthesize unsupported membranes of Nylon 6,10 by unstirred interfacial step polymerization of hexamethylene diamine (HMDA) and sebacoyl chloride. To observe the quality of the nylon film produced as a function of the rate of removal and of the concentrations of diamine and diacid chloride in their respective phases. The reader will find in the text a series of questions that will make him reflect ---so-called complementary activities--- promoting the development of competences complementary ones based on research exercises

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INTRODUCTION

Polymer Chemistry is a subject in the Chemistry Programs, student have a natural curiosity about polymeric materials, that we use in our everyday activities, such as poly(ethylene), PE, poly(styrene), PS, poly(amides) are well known by the group, and they are highly interested in their identification, production, characterization, and peculiar properties. Cooperative group strategy may be used to teaching methods (Obaya, 1997) It involves students in small group learning activities that promote positive interaction resulting in improved learning. Small heterogeneous groups of students work together to achieve individual and group objectives. The objectives may include both content and process skill outcomes. Additionally, students can develop social interaction skills. Our proposal is to apply cooperative learning in the Interfacial polymerization of Nylon 6,10 experimentation. The specific objectives of our project, can be summarized as:

To identify the origin of the raw materials used for industrial synthesis

To become acquainted with the most important processes used for the synthesis

To identify the most important variables related with the control of the polymerization process To correlate the physical chemical properties with the final applications of this material To reconognize the importance of the polymer's marked characteristics, the supply and demand To evaluate the importance of polymers recycling

The reader will find in the text a series of questions that will make him reflect ---so-called complementary activities-promoting the development of competences complementary ones based on research exercises. It is recommended that the student answers the questions before continuing the reading, as this will be very useful in the learning of the subject. When the purification of the raw materials, and quality characteristics are well understood, the student is in a good position to understand the production process. Once the monomer was purified and identified by means of spectroscopy test (IR, NMR, RI) The group can synthesize the polymer Activities such as literature search, monomer purification, polymer synthesis, characterization and identification, samples collection, final report and poster preparation must be programmed along the laboratory class Questions concerned with the type of industrial reactor and equipment requires to separate and purify the polymer must be also discussed

The 2015 Guidelines approved by the American Chemical Society Professional Training Committee require the curriculum for certified bachelor's degree to include the principles governing macromolecular, supramolecular, mesoscale and nanoscale. The justification for this new requirement recognizes that the synthesis, analysis, and physical properties of small molecules give an incomplete picture of the higher-order interactions that occur in these systems. Programs can meet this requirement through an indepth course on these systems or coverage distributed across two or more required courses. It is anticipated that most undergraduate programs will include at least some synthetic polymer coverage, but there may be concerns in many departments about how best to implement such coverage (Kosbar and Wenzel, 2017), suggested topics illustrating important aspects of the chemistry and properties of synthetic polymers. Teachers should enrich activities by investing the necessary time in the class by making students feel their experiences (Obaya et al, 2019). Thus, teaching related to aspects of technology would be encouraged as a way of complementing science, and the use of scientific processes and the linkage of teaching would be encouraged to the interests of students and to aspects that are significant to society.

INTERFACIAL POLYMERIZATION OF NYLON 6,10

Objectives

- To synthesize unsupported membranes of Nylon 6,10 by unstirred interfacial step polymerization of hexamethylene diamine (HMDA) and sebacoyl chloride. To observe the quality of the nylon film produced as a function of the rate of removal and of the concentrations of diamine and diacid chloride in their respective phases.
- To study the tensile properties of the synthesized collapsed film and compare it to the tensile properties of Nylon tire cord.
- To synthesize supported membranes of Nylon 6,10 on hydrophilic Millipore membranes.

Theoretical framework

Step polymerization: The two main types of polymerizations are chain and step polymerization (Rempp and Merrill, 1992). In chain polymerization, only the monomer can attach to the growing chain. However, in step polymerization the chain can grow by addition of monomers, oligomers, and other polymer chains. For example, a chain consisting of four units could be formed by two routes - a) the addition of a monomer to a trimer (chain with three units) or b) the reaction of two dimers (chain with two units). Note that in chain polymerization only the first route would occur. Step polymerization involves the reaction of bifunctional monomers (i.e., the two ends of the monomer each have a reactive functional group). Often in notes and texts, the functional groups are generically written as –A and –B; these two groups react toform a new group, -C-. For polyesters, this reaction might be:



Step Growth Polymerization – Polyamides (Nylons)

Polyamides are polymers where the monomer units are connected by amide groups (-C(=O)-NH-). Two generic schemes for polyamide synthesis include condensation of diacid and diamine monomers in the bulk state or at high concentrations. The first illustrated in Equation 1.

$$H_{2N} + (CH_{2})_{6}^{NH_{2}} + Ho^{2} + (CH_{2})_{8}^{2} \text{Coh} \longrightarrow$$

$$\left[NH - (CH_{2})_{6}^{NH} - NH - (CH_{2})_{8}^{2} \text{Coh} \right]_{n} \qquad \text{Equation (1)}$$

The above reaction creates H_2O as a byproduct, producing (n-1) moles of water for n molesof repeat units. It turns out that the reaction above is at or near equilibrium at low to moderate temperatures (Keq is around one to ten in value). To drive toward the product, hightemperature and very effective water byproduct removal must be achieved for high polymer conversion. A second approach is to use the much more reactive acid chloride group in place of the acid. The reaction of diamines with diacid chlorides have equilibrium constants that are about twoorders of magnitude higher than those of diamines with diacids. The reaction greatly favorsthe production of amide product, and the reaction kinetics is extremely fast:

Here (n-1) moles of HCl are created as a byproduct. HCl can be neutralized by the presence of weak base in the reaction solvent. This second reaction is extremely rapid, and rather difficult to control in moderate to high monomer concentrations. It is possible to accomplish a polymerization of diamine and diacid chloride in a good solvent for both monomers. On the other hand, the reaction is very fast, irreversible, and highly exothermic. The concentrations must be kept low, and the temperature must also be controlled, often below room temperature, to prevent overheating and side reactions. The solvent must be a good one not only for the monomers, but for the resulting polymer, to achieve high molecular weight.

Interfacial polymerization: A rapid, but more controlled approach is to take advantage of the fast kinetics of the polymerization of diamines and diacid chlorides at the interface between two immiscible phases (Odian, 2004). Interfacial polymerization involves the reaction of monomer A dissolved in an organic phase with a monomer B dissolved in an inorganic phase at the interface between the two nonmiscible phases. Interfacial polymerization can be of two types. In one case, there are two separate macroscopic layers in contact with each other. This is called unstirred interfacial polymerization. This is what we are going to study in this experiment. Unstirred interfacial polymerization is used to produce membranes and to create polymer by continuous removal in a single rope. The other case is one where one phase is dispersed astiny droplets in the other (continuous) phase by using high-speed stirring. This is called stirred interfacial polymerization.

Stirred interfacial polymerization is used to produce tiny microcapsules (hollow inside) and microspheres (not hollow) for various applications like controlled release of drugs and pesticides. Unstirred interfacial polymerization of Nylon 6,10: We shall study the reaction of 1,6 diamino hexane (HMDA) with diacid chloride (sebacoyl dichloride-SC) to form Nylon 6,10 as shown in reaction (1). The polymer will form at the interface between a layer of diaminohexane dissolved in water (aqueous phase) and a layer of diacid chloride dissolved in dichloromethane-DCM (organic phase). The aqueous phase

also contains sodium bicarbonate as a getter for HCl (see reaction 2).

$$2HCl+Na_2CO_3 \longrightarrow 2NaCl+H_2O+CO_2$$
 Equation (2)

The denser organic phase forms the lower phase (chlorinated solvents are used, which are denser than water) while the aqueous phase will be carefully poured on top of the organic phase. Interfacial polymerization occurs instantly forming a thin film of solid nylon 6,10 at the interface. The film at the interface stops further reaction by preventing the monomers from meeting each other (some monomers do reach other by diffusion through the film, but this is so slow that the film appears to have stopped growing).

The hydrochloric acid is rapidly neutralized by the carbonate in the aqueous phase. If the nylon film is pulled up, out of the interface, new film is formed continuously. In contrast to reaction in the bulk(discussed before while discussing reaction 8), there is no need to control of the stoichiometric ratio in this experiment.

The stoichiometry at the interface is controlled perfectly by the mass transfer-controlled rate of diffusion and consumption of each monomer species. In other words, in this diffusioncontrolled situation, the monomer a to bratio is essentially one to one. Notice that interfacial polymerization is a very nice problem to observe the interplay of core chemical engineering concepts like diffusion and reaction.

Complementary activities: Can you identify all the diffusion and reaction processes?

If these processes are important, then what experimental variables are important?

Fibers: Natural (e.g., cotton, silk etc.) and synthetic fibers (Nylon, Rayon, Polyester etc.) are all polymers. They are visibly different in being "thread-like" -- elongated in one direction and small in cross-section. Further, they distinguish themselves from other polymers in being anisotropic in their properties, i.e., their properties along their long axis are different from that in the perpendicular direction. Fibers are usually very strong in directions.

Synthetic fibers are typically drawn from undrawn polymers by stretching them with a specific force. Membranes: Membranes are films that are used to separate out a mixture of species into its constituents. Often, a membrane will contain nano to micro-scale pores, and may thus have some void space within its structure. This void space is useful for separations operations. It also has a large effect on the final mechanical properties of the material. Tensile testing:

Tensile testing is perhaps the most used form of mechanical testing of polymeric materials. As shown in the figure below, a polymer sample of initial area of cross-section A_0 and length Lo is subjected to a variable tensile (as opposed to say, compression or shear) force F such that the sample is elongated at a constant rate of dL/dtwhere L is the length of the sample at time t.



Figure 1. Schematic illustrating the tensile test and features on a stress-strain plot

The force F and the length L is noted at periodic intervals. Following this the following quantities are calculated:

Draw ratio,
$$\lambda = \frac{L}{L_o}$$
 Equation (4)

Engineering strain, $e = \frac{L-L_0}{L_0}$ Equation (5) Engineering stress, $s = -\frac{F}{A_0}$ Equation (6)

The plot of σ vs. ϵ is called the stress-strain plot. The stressstrain plot gives properties of the material like the elastic modulus and the yield stress which are independent of the shape and size of the test sample but depends on the rate of strain (or elongation rate). The elastic modulus is the slope of the stressstrain plot at very low strains.

METHODOLOGY

Prepare the two phases and introduce the aqueous phase (Guillermo, 2018) on top of the organic phase HMDA (hexamethylenediamine) in the hood (Table 1). For manipulation use vinyl gloves. The students must complete the different stages for the experiment

Procedures: Synthesis of unsupported membranes of Nylon 6,10

- First, set the variable speed shaft (a lab mixer shaft) to its lowest STABLE rotating speed. Estimate this RPM by measuring how much string is wound up in, say, 60 seconds, or by counting revolutions. Record this value. What you really want is the linear rate of withdrawal, cm/s, so the string test will give it directly. Otherwise, it will be (RPM/60) (pD)where D = shaft diameter, cm.
- For each of the recipes you are to do, follow the procedure below: Using a 250 ml beaker in a hood, introduce 50 ml of the organic phase. Then tip the beaker and slowly let the 50 ml of aqueous solutions slide down the glass wall over the organic phase. Restore thebeaker upright.
- Remove from hood and set under the lab-mixer shaft (the shaft held horizontally). With spatula or forceps, "catch" the membrane that has formed at the interface, draw it upand out, and attached the top end to the shaft of your variable speed set (Lima et al, 1997). Now run the set until you have produced 3 to 4 meters of collapsed membrane (which will be wound aroundthe shaft).
- Unwind the collapsed membrane into a large (1 to 2 liter) vessel of water, so as to rinse away the soluble

salt. 5. Separate into two portions. Leave one portion wet for tensiletesting. Dry other portion in the 70°C air oven for weight measurement

• Using the dial micrometer (calibrated in 0.001 inches), estimate the wet thickness of the film. Determine the approximate weight per meter of length of dry film.

Tensile test properties of the collapsed film

- Take a length of about 50 cm of your wet film. Make two lines 10 cm apart. Fastenone end of the length to a post, the other end to the force gauge. If the gauge reaches its limit, disconnect, and pull manually until the film breaks.
- Record the final length Lf as a ratio to the original length Lo. This is the draw ratio.
- You will plot tensile force (read from the gauge) as a function of L/L₀. Do this for two or three 50 cm lengths from each run α and β or α and γ , to establish how closely the data agree.
- Repeat the same experiment with Nylon undrawn tire cord furnished to you from a spool. You will probably be able to take a single filament of this material and measure its tensile load vs. L/L₀ relation all the way to break. Measure the weight of a known length of the nylon cord.
- Determine what happens when you do similar measurements on a dried film.
- Take a length of about 50 cm of your wet film. Make two lines 10 cm apart. Fastenone end of the length to a post, the other end to the force gauge. If the gauge reaches its limit, disconnect, and pull manually until the film breaks.
- Record the final length Lf as a ratio to the original length Lo. This is the draw ratio.
- You will plot tensile force (read from the gauge) as a function of L/Lo. Do this for two or three 50 cm lengths from each run α and β or α and γ, to establish how closely the data agree.
- Repeat the same experiment with Nylon undrawn tire cord furnished to you from a spool. You will probably be able to take a single filament of this material and measure its tensile load vs. L/Lo relation all the way to break. Measure the weight of a known length of the nylon cord.

Complementary activities: Using this and the literature value for the density of commercial nylon, you can determine the cross-sectional area of the cord.

Complementary activities: Postulate as to the differences between the wet and dry film.

Complementary activities: Using this and the literature value for the density of commercial nylon, you can determine the cross-sectional area of the cord. Determine what happens when you do similar measurements on a dried film.

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Synthesis of supported membranes

- Use the solutions of recipe □ only. Use a hydrophilic Millipore 25 mm 0.45-micron membrane to absorb from a petri dish the aqueous solution. Then float this on a layer of the organic phase in another petri dish, for 10 minutes. Remove and wash by submersion inwater.
- Using a second 25 mm 0.45-micron hydrophilic membrane, let it absorb from a petri dish the organic solution. Then float this on a thin layer of the aqueous solution. After 120minutes, remove and wash.

For each of the supported membranes, test for wettability: Will a water drop spread on the nylon film or not? Test the supported membranes for permeability to yellow food dye.

Complementary activities

Will the dye pass through the membranes?

RESULTS AND CALCULATIONS

- Record all the experimental conditions /variables and the measurements mentioned in the lab notebook.
- Unsupported membranes:

Table 1. Steps for development the experiment

Recipe O	rganicPhase Aqueous Ph	aseLaboratory Grou	ps doingit
α	1.5ml SC + 50 ml DCM	2.20g HMDA +50 ml water +4 g Na2CO3	A, B, C, D, E, F
β	0.75ml SC + 50 ml DCM	1.10g HMDA +50 ml water +4 g Na2CO3	A, C, E
γ	3.0ml SC + 50 ml DCM	4.40g HMDA +50 ml water +4 g Na _{2CO3}	B, D, F

Compare the weight per meter of length of the two procedures. Choose your methodof presentation --- table or bar chart etc.? Plot the stress-strain plots for the different recipes. The plot should illustrate the difference, if any, between the two recipes. Be sure to indicate the recipe used as a legend. You can estimate the elastic moduli of each recipe from the slopes of the stressstrain plots. Plot stress-strain plots of Nylon tire cord and determine its elastic modulus. Explain the differences in b and c results.

Complementary activities: What is different between the cord filament and the nylon rope that you made?

Supported membranes: Tabulate the required observations. Comment on whether your nylon films supported on Millipore membranes are porous and permeable to dye, or impermeable.

Complementary activities: Considering that the membrane is hydrophilic, what do you think happened in each case.

CONCLUSION

A more positive affective relationship among students and between students and faculty has been observed in the group during the polymer laboratory class. In this experiment the students compare the weight per meter of length of the two procedures, determine the approximate weight per meter of length of dry film, postulate as to the differences between the wet and dry film, test the supported membranes for permeability to yellow food dye. Complementary activities promoting the development of competences complementary ones based on research exercises. The concepts were shared, as a result, a more positive psychological well-being, greater self-esteem, self-sufficiency, social skills was developed in the class. The students' comments were very favorable to the Project. This was a lifetime experience that we want to share with others

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