Thermosetting polyurethane foams by supercritical CO₂ as physical blowing agent

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ABSTRACT

The aim of the present study was the achievement of thermosetting polyurethane foams by a batch foaming process that uses CO₂ as a physical blowing agent (PBA). In particular, to address the recent interest in combining the gas (physical) foaming with the conventional (chemical) polyurethane foaming, a novel instrumented pressure vessel was designed for investigating: i) sorption under high gas pressure on the two, separate, components of the polyurethane foams, ii) synthesis under high gas pressure after the two components mixing, and iii) foaming upon release of the pressure. Results revealed a significant effect of sorbed CO₂ on the polyurethane synthesis and the need for the design of a new chemistry to exploit the use of physical foaming on thermosetting polyurethanes.

INTRODUCTION

In the field of thermosetting polyurethane foams (PUs) obtained by the curing reaction between poliol and isocyanate, the selection and the amount of the blowing agents is very important to the final cellular structure and properties of the foam. Since the beginning of the history of PUs, CO₂, derived from the blowing reaction of isocyanate with water, is used as blowing agent. In the late 1950s, large PUs production also utilized chlorofluorocarbons (CFCs) and hydrochlorofluorocarbons (HCFCs) as PBAs, then replaced by hydrocarbons (HCs) (e.g. pentane) because of their negative environmental impact. However, although cheap and with zero ozone depletion potential (ODP), HCs use was limited by flammability concerns. In this context, CO₂ is eco-friendly and safe, with zero ODP and the lowest GWP among known blowing agents [1]. In order to use CO₂ as a physical blowing agent (in a totally different manner to what has been done previously, with CO₂ resulting from the reaction of isocyanate with water) knowledge of the behavior of the whole system in the presence of high pressure CO₂ is fundamental. In particular, the following are needed: 1) the physical properties of polymer/gas solutions, namely solubility, diffusivity, specific volume and interfacial tension of the poliol- and isocyanate-CO₂ solutions in contact with the external, high pressure CO₂ phase; 2) the effect of the CO₂ on the reactions involving the different chemicals in the polyurethane synthesis; 3) the way the reactive system behaves when CO₂ is rapidly released to induce foaming.
We herein report our approach to the problem, with the design of new experimental apparata. We report, in particular, measurements at 35°C of solubility, mutual diffusivity, specific volume and interfacial tension of polyol/CO₂ solution at CO₂ pressures up to 80 bar and isocyanate/CO₂ solution up to 55 bar, obtained by a coupled gravimetry-Asyncmometric Drop Shape Analysis (ADSA) [2-4]. Foaming experiments, conducted at 35°C and various CO₂ pressures, with a novel instrumented pressure vessel capable of retrieving data on sorption, synthesis and foaming are described. This pressure vessel makes use of a sample holder with a flexible impeller that allows to keep polyol and PMDI separated during the sorption stage and to mix the components to start the PU synthesis. Furthermore, it makes use of a sapphire window and FT-NIR spectroscopy to monitor sorption and the synthesis. The effect of CO₂ on the reaction has been evaluated at 35°C and 40bar.

MATERIALS AND METHODS

Materials

A formulated polyol and a polymeric methylene diphenyl di-isocyanate (PMDI) were supplied by DOW Italia S.r.l. (Correggio, RE, Italy) and used “as received”. High purity grade CO₂ was supplied by SOL (Naples, Italy).

Sorption measurements of CO₂ in polyol and PMDI

Coupled sorption-ADSA experiments were conducted by using magnetic suspension balance (MSB) (Rubotherm Prazisionsmesstechnik GmbH, Germany), equipped with a high temperature/high pressure (HT-HP) view cell. The simultaneous measurement was made possible by adopting a custom-designed measuring device, which consists of a crucible containing the polymer/CO₂ solution (for weight monitoring) and a Teflon rod to which the polymer/CO₂ solution drop is stuck (for ADSA) (see Figure 1a). Drop changes in volume and shape were observed through two optical quality windows, by using an adjustable high resolution CCD camera (BV-7105H, Appro), equipped with a modular zoom lens system (Zoom6000, Navitar). The CCD camera is connected to a computer, and a commercial software (FTA32 Video 2.0, First Ten Angstroms) is used to analyze drop profile [2-4]. Furthermore, in order to achieving the optimal threshold background for digitizing the drop image, a uniform bright background was be provided by light emitting diodes.

The data flow adopted for the elaboration of the data acquired during the coupled sorption-ADSA measurement is illustrated in Figure 1b.
Sorption and ADSA experiments were carried out by isothermal pressure increments up to 80 bar for polyol while up to 55 bar for PMDI, at 35°C, after having placed both the crucible containing the polymer and the rod with the polymer drop in the HT–HP view cell. In detail, sorption measurements were performed by step-wise increments of the gas pressure (500 kPa steps ca.), after the attainment of equilibrium sorption in the previous step. Concurrently, during each pressure step, image acquisition of the pendant drop was performed every 10 min. and drop volume was calculated by using the commercial software.

**PU foaming with CO₂**

In Figure 2, the steps of foaming experiment are shown. Inside the pressure vessel, polyol and PMDI, separated by a flexible impeller, are placed (1), pressurized with CO₂ at 35°C until saturation (2) and mixed (3). Finally, at any stage of the curing reaction physical foaming occurs due to pressure release (4).

Different foaming tests will be carried out, by changing process parameters as sorption time in order to evaluate its influence on the PU morphology.
Spectroscopic measurements

The curing reaction, at 35°C and atmospheric pressure, was monitoring by using a fiber optic reflectance probe, placed beneath the sampler holder contained the two reactants, connected to Frontier™ NIR spectrometer (Perkin Elmer Inc., Waltham, MA, USA) equipped with a NIR source, a CaF₂ beam splitter and a INGAAS detector. During the cure, spectra were recorded automatically at regular time interval of 3s for 10 minutes using Perkin Elmer TimeBase software, in the spectral range 4000-10000 cm⁻¹ with a resolution of 16 cm⁻¹ and 4 scans just after mixing.

RESULTS

Sorption measurements of CO₂ in polyol and PMDI

Sorption measurements for the polyol/CO₂ solution show quite a large CO₂ solubilization (17% wt ca. at 80 bar and 35°C), in turn responsible for a moderate swelling of the polymer and an extensive effect on the interfacial tension, which reaches vanishing values at the highest investigated pressure. Mutual diffusivity goes from 1e⁻⁶ to 1e⁻⁵ cm²/s in the whole experimental range. The same study conducted on the PMDI at 35°C and CO₂ pressures up to 50 bar has shown for the sorption isotherm a maximum of about 6% wt, with mutual diffusivity of 2 10⁻⁶ cm²/s ca. and a minor swelling of the polymer/gas solution with respect to the neat polymer, while the interfacial tension reaches the value of 15 mN/m ca. In both sorption measurements, for higher pressure it has been observed that low molecular weight components of polyol and PMDI are solubilized in CO₂, as confirmed by Gel Permeation Chromatography analysis.

PU foaming with CO₂

In Figure 3 PUs obtained at 35°C and 40 bar of CO₂ pressure by changing sorption time (or CO₂ amount) are shown. Curing time was fixed at 1 minute and mixing at 250 rpm for 10s. It is possible to note that by increasing sorption time, PU change its morphology from an inhomogeneous to more homogeneous one until to become a pulverized product for high sorption time indicating a CO₂ effect on the system.

![Figure 3. PUs obtained at different sorption times (or CO₂ amount).](image-url)
Spectroscopic measurements

NIR spectra of polyol and PMDI at room temperature are shown in Figure 4 a and b, with their characteristic absorption peaks.

![Figure 4. NIR spectra of polyol (a) and PMDI (b) at room temperature.](image)

In Figure 5 spectra collected at the beginning (blue line) and the end (red line) of the curing reaction, at room temperature and atmospheric pressure, are reported. From these spectra is possible to follow the NCO band decrease and the NH band (of urethane group) formation and increase [5,6].

![Figure 5. NIR spectra collected at the beginning t₀ (blue line) and the end tₚₐₜ (red line) of the cure.](image)
CONCLUSIONS
The use of CO\textsubscript{2} as PBA to obtain PUs on lab scale was investigated. Solubility, mutual diffusivity, specific volume and interfacial tension of polyol/CO\textsubscript{2} solution at CO\textsubscript{2} pressures up to 80 bar and PMDI/CO\textsubscript{2} solution up to 55 bar, were measured by using a fully-experimental, coupled gravimetry- Axisymmetric Drop Shape Analysis at 35°C. These data were used to produce PUs in a pressure vessel, at 35°C and 40bar of CO\textsubscript{2} pressure, by changing the sorption time. It was noticed an influence of CO\textsubscript{2} on the final product, in fact by increasing sorption time PU morphology improves until an optimum and then it changes resulting in a pulverized product. FT-NIR spectra were recorded to monitor the curing reaction after mixing of the two reactants at room temperature and atmospheric pressure.

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REFERENCES


