



Proceedings Recyclable Insulating Foams for High Temperature Applications ⁺

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Abstract: The recently approved restriction on diisocyanates highlights the health and safety issues concerning polyurethane manufacturing and the relevance of developing sustainable insulating polymeric foams. This is particularly challenging for applications where the foam is subjected to high temperatures (>80 °C) and bear loads, such as insulating and bonding material for district heating pipes. As part of a PhD project concerning pre-insulated district heating pipes for the circular economy, polybutylene (PB-1) has been identified as a promising candidate for the application, due to its low thermal conductivity, high temperature mechanical properties retention, excellent environmental stress cracking resistance (ESCR) and outstanding creep resistance. It is a recyclable thermoplastic and of non-toxic nature, pre-requisites for circular product development. On the contrary to other polyolefins, PB-1 is reported to strain-harden and has high melt strength, required properties for foaming. The purpose of the study is to assess the foamability of PB-1 through extrusion foaming experiments. A twin-screw extruder was used with varying concentrations of a chemical blowing agent. The obtained samples have been characterised for density, expansion ratio and microstructure. Foams with a volume expansion ratio of 1.8 were achieved. The results confirm the foamability of this polymer. The increase of the die pressure and its contribution to strain hardening were identified as key parameters for successful foaming. Further research will include improving the expansion ratio with a physical blowing agent and mechanical characterization of the foam.

Keywords: polymeric foam; polybutylene; district heating; insulation; thermoplastic; circular economy

1. Introduction

District heating (DH) is called to be a cornerstone element towards decarbonizing the heating sector. Thanks to a district heating network, the use of decentralized renewable and waste heat sources is enabled. For its technical and economic feasibility, correct insulation of district heating pipes is critical. As for state of the art, all district heating pipes are insulated with polyurethane foam (PU) [1].

PU foams present the lowest thermal conductivity among foamed polymers used commercially [2]. While PU has excellent thermal and mechanical properties, it has important environmental and toxicity drawbacks, which trigger the need for research in new and alternative insulating foams.

PU is produced from a ca. 50/50 mix of a polyol and either TDI (toluene diisocyanate) or MDI (methylene diphenyl diisocyanate). TDI and MDI are classified as suspected of causing cancer, as dermal and respiratory sensitizers, for acute toxicity following inhalation, as well as eye, skin and

respiratory irritants under the EU's CLP Regulation [3]. The European Commission has aproved to restrict the use of diisocyanates to a concentration limit of 0.1% by weight under REACH on 4 August 2020 [4].

There is a need for developing environmentally friendly insulating foams following a holistic approach: conceptualized from the beginning out of non-toxic and recyclable materials and a safe manufacturing process. In this context, the foamability of PB-1 is explored. Promissing properties for the application are its low thermal conductivity of 0.114 W/mK [5], high temperature mechanical properties retention, excellent environmental stress cracking resistance (ESCR) and outstanding creep resistance [6]. It is a recyclable thermoplastic and of non-toxic nature, pre-requisites for circular product development. On the contrary to other polyolefins, PB-1 is reported to strain-harden and has high melt strength [6], required properties for foaming [7]. However, its foaming behaviour has not been found reported in the open literature. In this study the foamability of PB-1 is studied through extrusion foaming experiments with a chemical blowing agent. The obtained samples have been characterised for density, expansion ratio and microstructure. Foamability is assessed in terms of achieved volume expansion ratio, microstrucure and processing window using available extrusion equipment. Optimization of the foaming process is out of the scope of this study. The presented results are part of a larger screening of commertial PB-1 grades.

2. Experiments

2.1. Materials

Toppyl PB0110M PB-1 from LyondellBasell was extrusion foamed using Hydrocerol CT 550 chemical blowing agent (CBA), kindly provided by Clariant.

2.2. Extrusion Foaming Process

The extrusion process was undertaken with a twin screw ZSE 27 MAXX extruder (Leistritz Extrusionstechnik GmbH, Nürnberg, Germany), with D = 28.3 mm, L/D = 48, and 12 modular barrels with 2.1 kW heating power each and water cooling. A strand die with 3 strands of 4 mm diameter each was used. The feeding temperature was set as low as possible as to create a melt seal and avoid premature degassing of the CBO, which was set at 140 °C. The temperature was progressively increased up to 175 °C after the CBO dosing point, which as to allow its complete decomposition, and then progressively lowered. The amount of dosed CBO was varied from 4.2 to 10% wt., for die exit temperatures of 110 and 115 °C. Screw speed rate and feed rate were kept constant to 100 rpm and 4 kg/h respectively.

2.3. Foam Characterization

The foam samples were randomly collected at each processing condition after achieving the stedy state and characterized for density, volume expansion, cell size and cell density.

The density was determined with a 100 mL glass pycnometer, destilled water and a Sartorius AC 211 S (Goettingen, Germany) balance.

Foam volume expansion ratio is often determined as the ratio between the density of the polymer and the density of the foam. However, since a CBO is used, for which 70% wt. are effective foaming components, and up to 10% wt. CBO was dosed, this results in the addition of 3.3% wt. of the decomposition products. It was found more accurate to determine the volume expansion ratio V_{exp} as

$$V_{exp} = \frac{\text{diameter foam extrudate}}{\text{diameter polymer extrudate}}$$
(1)

in an analogous method as used by [8].

The diameter of the obtained foam samples and neat polymer extrudates where obtained from photographs taken using a Nikon D700 camera and a tripod. Photographs were recorded from immediately after collection (time between collection and initial photo max 20 s) up to over 2 h after

extrusion, in 20 s time-laps intervals. The diameter of each strand was measured in 5 different locations of each strand once its dimension was stable using Fiji [9]. The average of the five measurements is presented in each case.

The morphology of the foams was examined in an optical microscope (Leica DMLP, Wetzlar, Germany). Cell size was measured from the obtained micrographs using Fiji [9]. Cell population density (N₀), defined as the number of cells per unit volume of the original unfoamed polymer, was calculated as [10,11]:

$$N_0 = \left(\frac{n}{A}\right)^{3/2} \cdot V_{exp} \tag{2}$$

where (N₀) is the cell population density (cell/cm³), n is the number of cells in the micrograph and A the area of the micrograph (cm²).

3. Results and Discussion

The density of the obtained samples are presented in Table 1:

die T (°C)	CBO % wt.	Density (g/mL)
110	4.2	0.740 ± 0.032 *
110	7.1	0.5 ± 0.017 *
110	8.50	0.598 ± 0.031 *
110	10	0.53 ± 0.034 *
115	4.2	0.50 8± 0.122 *
115	7.1	0.508 ± 0.036 *
115	8.50	0.488 ± 0.178 *
115	10	0.601 ± 0.089 *

Table 1. Density for samples under different process conditions.

* Standard deviation.

The determined volume expansion ratio, cell population density and cell size and their relation with the process conditions can be found in Figure 1:



Figure 1. (**a**) expansion ratio vs introduced amount of CBO. (**b**) Cell population density vs introduced amount of CBO. (**c**) Cell size vs. introduced amount of CBO.

Representative micrographs for each process condition can be found in Figure 2.



Figure 2. Micrographs of obtained foams under different process conditions: die $T^\circ = 110 \text{ °C}$ and 4.2% CBO (**a**); 7.1% CBO (**b**); 8.5% CBO (**c**) and 10% CBO (**d**). die $T^\circ = 115 \text{ °C}$ and 4.2% CBO (**e**); 7.1% CBO (**f**); 8.5% CBO (**g**) and 10% CBO (**h**).

3.1. Microstructural Characterization and the Relationship with the Process Conditions

As can be seen in Figures 1c and 2, foams present large pores with significant cell size variability. The two die temperatures used did not produce any difference in the obtained cell size for CBO concentrations of 4.2 and 7.1% wt. Cell size decreased for 8.5% wt. as cell population density increased. This inverse trend is often encountered [12]. This effect is particularly significant for the 110 °C die T° trial, where the average cell size was reduced by more than 50%. After this point N₀ decreased again and cell size increased. This can be explained as the increase of CBO provides more

available gas for cell nucleaion, until a maximum is reached. With further gas increase and cell groh, cells collide against each other, reducing the n° of cells and increasing their size [13] as can be observed by the trends in Figure 1b,c.

3.2. Effect of Process Condictions on the Volume Expansion Ratio

The two die temperatures tested were selected as a result of a preliminar process parameters screening. With die T° over 120 °C, limited expansion was achieved. This is an indication for low melt strength, which impeches the polymer melt to withstand the stretching forces of the bubble growth, leading to cell collapse, and of high gas diffusion out of the polymer matrix. The lower processing temperature was conditioned by viscosity, being 110 °C the lowest temperature at which the melt would flow out of the extruder.

As it can be seen in the Figure 1a, with the die T° of 110 °C, the expansion ratio increased until a maximum and then decreased. This indicated that at this temperature the expansion behaviour is governed by the polymer melt's stiffness. Expansion increases with increasing CBO concentration due to the plastitication effect of the gas [12]. After reaching a maximum, the gas starts diffusing out of the polymer's hot skin. For the 115 °C die temperature serie, the maximum expansion was achieved with the lowest concentration of CBO tested. Since the cell size and cell population density of the samples obtained at the lowerst concentration (4.2%) was not higher than those obtained with increasing CBO dosages, we can conclude that from that dosage the increased level of gas just increased the diffusion out of the hot skin of the extrudates, thus the volume expansion remained at constant levels.

During the execution of the trials, a transient situation was encountered where the die pressure was increassed from the 45 bars of the reported trials up to 75 bar. Under these conditions a higher volume expansion was obtained, and strain hardening was observed on the extrudates. The relationship between strain hardening and succesful foaming is known in the field [7], as well as the relation between pressure drop and cell population density [14]. The data of this obtained foam is not included due to the non-reproductible process conditions (transient), but shows the potential of improved foaming of this resin under optimal process conditions. As mentioned in the introduction, process optimization is out of the scope of this study, which aims and screening foamability.

4. Conclusions

An experimental study on the extrusion foaming behaviour of a comertial PB-1 resin was presented. Foamability of the resin was confirmed. High density foams where produced, with a volume expansion of 1.8. Processing window was screened and the influence of process conditions on the microstructure and volume expansion of the obtained foams was studied and indicated that PB-1 foam of medium density can be produced with the optimization of the foaming process.

Author Contributions: L.D. was responsible for the study design and conception, performing the experiments, analyzing the data and chiefly responsible for preparing the manuscript. I.W. provided scientific supervision of the study. Both authors contributed with the critical discusion of the results and editing of the manuscript. All authors have read and agreed to the published version of the manuscript.

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Conflicts of Interest: The authors declare no conflict of interest.

Abbreviations

The following abbreviations are used in this manuscript:

PB-1	polybutylene
ESCR	environmental stress cracking resistance
DH	District heating

PU	Polyurethane
TDI	toluene diisocyanate
MDI	methylene diphenyl diisocyanate
EU	European Union
CLP Regulation	classification, labelling and packaging of substances and mixtures regulation
REACH	Registration, Evaluation, Authorisation and Restriction of Chemicals
CBA	Chemical Blowing Agent
wt.	weight
T°	temperature

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