Biosourced polymer foam production using a (SC CO2)assisted extrusion process

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A process based on extrusion coupled with supercritical carbon dioxide (scCO₂) was implemented. $scCO_2$ modifies the rheological properties of the material in the barrel of the extruder and acts as a blowing agent during the relaxation at the passage through the die. An experimental device based on a single-screw extruder allows the injection of $scCO_2$ into the melt, the mixing of both components and the creation of porosity into the extruded polymer. In this work, it was used to produce foams of two bio sourced, semi-crystalline polymers The effects on material porosity of different parameters such as, temperature before and after the die, pressure drop or CO_2 concentration, were studied. Different configurations on the extruder were tested.

INTRODUCTION

Polymers are widely used in several areas. However, due to their slow degradation and the predicted exhaustion of the world petroleum reserves, significant environmental problems have arisen. Therefore, it is necessary to replace them with biosourced polymers with equivalent properties. It is also important to find new way to process them, which are more environmental friendly.

Extrusion is a process converting a raw material into a product of uniform shape and density by forcing it through a die under controlled conditions [1]. It has extensively been applied in the plastic and rubber industries, where it is the most important manufacturing process. A particular application concerns the generation of polymeric foams. Polymeric foams are expanded materials with large applications in the packaging, insulating, pharmaceutical and car industries because of their high strength/weight ratio or their controlled release properties.

Conventional foams are produced using either chemical or physical blowing agents. Various chemical blowing agents, which are generally low molecular weight organic compounds, are mixed with a polymer matrix and decompose when heated beyond a threshold temperature. This results in the release of a gas, and thus the nucleation of bubbles. This implies however the presence of residues in the porous material and the need for an additional stage to eliminate them.

Injection of $scCO_2$ in extrusion process modifies the rheological properties of the polymer in the barrel of the extruder and $scCO_2$ acts as a blowing agent during the relaxation when flowing through the die [2]. The pressure drop induces a thermodynamic instability in the polymer matrix, generating a large number of bubbles. The growth of cells continues until the foam is rigidified (when T<T_g or T_f). Moreover, its relatively high solubilisation in the polymer results in extensive expansion at the die. The reduction of viscosity decreases the mechanical constraints and the operating temperature within the extruder. Thus, coupling

extrusion and $scCO_2$ would allow the use of fragile or thermolabile molecules, like pharmaceutical molecules. The absence of residues in the final material is also an advantage for a pharmaceutical application.

Our lab has developed a scCO₂-assisted extrusion process that leads to the manufacturing of microcellular polymeric foams and already elaborated microcellular foams using a biocompatible amorphous polymer [3, 4]. In this work, this process has been applied to two semi-crystalline biosourced polymers. Foam production of semi-crystalline polymer is less frequent in the literature. Crystallinity hinders the solubility and diffusion of CO_2 into the polymer and leads consequently to less uniform porous structure [5]. Moreover, it has been shown that a large volume expansion ratio could be achieved by freezing the extrudate surface of the polymer melt at a reasonably low temperature [6].

Thus, in this work, we investigate the feasibility of this process with biosourced semicrystalline polymers. We first try to control and improve the porous structure, studying the influence of melt and die temperatures on PHBV, which a copolymer of poly3hydroxybutyrate and hydroxyvalerate. Then, we modify the experimental device in order to experiment the mixing of carbon dioxide into the polymer. We used a different polymer for this study, called BP here, and we looked at the influence of the mixing device and of the carbon dioxide content.

MATERIALS AND METHODS

PHBV (M_w=600 kDa), with a HV content of 13 % and plasticized with 10 % of a copolyester was purchased from Biomer (Germany). Melting onset temperature was measured at 159°C by DSC (ATG DSC 111, Setaram) and no glass transition temperature was detected at temperatures as low as -20°C. The solid density ρ_P , determined by helium pycnometry (Micromeretics, AccuPYC 1330) was found to be 1216 kg.m⁻³.

Melting onset temperature of the BP polymer was measured at 184°C by DSC (ATG DSC 111, Setaram) and the glass transition temperature was detected at 44°C. The solid density ρ_P , determined by helium pycnometry (Micromeretics, AccuPYC 1330) was found to be 1027± 5 kg.m⁻³.

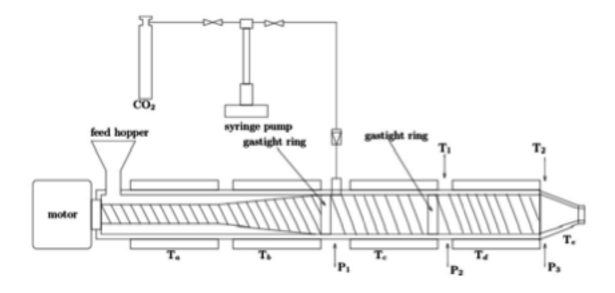
Figure 1 shows the experimental set up, which has previously been detailed elsewhere [3, 4]. The single-screw extruder has a 30 mm-screw diameter and a length to diameter ratio (L/D) of 35 (Rheoscam, SCAMEX). A great L/D ratio generally indicates a good capacity of mixing and melting but important energy consumption. The screw is divided into three parts. The first one has a length to diameter ratio of 20 and the two others have a length to diameter ratio of 7.5. Between each part, a restriction ring has been fitted out in order to obtain a dynamic gastight which prevents scCO₂ from backflowing. The first conical part allows the transport of solid polymers and then, their melting and plasticizing. Then, the screw has a cylindrical geometry from the first gastight ring to the die. This die has a diameter of 1 mm and a length of 11.5 mm. The temperature inside the barrel is regulated at five locations: T_a and T_b before the CO₂ injection, T_c and T_d after the injection and T_e in the die.

There are three pressure and two temperature sensors: P_1 after the CO_2 injector, P_2 and T_1 before the second gastight ring and P_3 and T_2 by the die. This allows measuring the

temperature and the pressure of the polymer inside the extruder. Errors associated to pressure and temperature measurements were about 2 bars and 3.3°C respectively.

CO₂ (N45, Air liquide) is pumped from a cylinder by a syringe pump (260D, ISCO) and then introduced at constant volumetric flow rate. The pressure in the CO₂ pump is kept slightly higher than the pressure P₁. The CO₂ injector is positioned at a length to diameter ratio of 20 from the feed hopper. It corresponds to the beginning of the metering zone, that is to say the part where the channel depth is constant and equal to 1.5 mm. The pressure, the temperature and the volumetric CO₂ flow rate are measured within the syringe pump. CO₂ density, obtained on NIST website by Span and Wagner equation of state [7], is used to calculate mass flow rate and thus the CO₂ mass fraction w_{CO_2} .



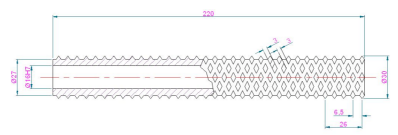


In the second part of the work, this device was modified, replacing the last part of the screw (7.5 L/D) by a mixing element, that is a pineapple mixer with diamond-shaped pins. This one can be seen on Figure 2.

Dynamic mixers are often used to improve mixing in single screw extruder. Distributive mixing and dispersing one are commonly distinguished even if they are not physically separated. The component to be mixed here is CO_2 and the first mixing to ensure is the distributive one. Hence, the aim here is to go further in this aspect. We chose a distributive mixer, like the pineapple one, which distributes spatially the phase to be mixed acting on the shear flow. The pineapple has advantages in that the flow turns to be streamlined, the barrel is well wiped by the pins and rather frequent flow splitting occurs [8].

The other type of mixers, the dispersive ones, generate high enough shear to allow breaking agglomerates. There are rather used to disperse a solid phase into a polymer melt.

Figure 2 Mixing device (pineapple)



Experimental conditions have to ensure that the polymer flows through the barrel without being thermally degraded. Temperatures have been adapted to each polymer to obtain rapid enough solidification after the die to avoid coalescence of the bubbles and a collapse of the foam structure but also to avoid the blocking of the screw.

As for PHBV, influence of the die temperature was tested. For each experiment, only the temperature of the metering zone T_d and of the die T_e were changed. The three other temperatures T_a , T_b and T_c were kept constant at 160°C. CO₂ mass fraction w_{CO_2} was also kept constant at 1.3 %, which might be less than solubility: in [9], for a similar polymer with a HV content of 12%, at 313K and 15 MPa, the solubility of CO₂ was found to be of 55% in weight. The rotation speed of the screw was fixed at 30 rpm. Three series of experiments were carried out. T_d was fixed at 140°C, 135°C and 130°C respectively and T_e varied from 140 down to 110°C. At lower values of T_e , the extruder stopped due to over-limited pressure P₃ according to the established alarm value.

Then we tested the distributive mixing level in the metering zone using another biosourced polymer (BP). The introduction of a mixing element was tested in order to know its influence on the mixing efficiency. In each configuration, CO_2 flowrate was varied and we studied its influence on the porosity of the foams.

Once steady state conditions were reached, extrudates were collected and flowrates were measured. Several samples were collected during the experiment in order to check the homogeneous structure of the extrudates. To calculate the apparent porosity ρ_{app} , samples were weighed and their volumes were evaluated by a water pycnometer. Porosity, defined as the ratio of the pore volume to total volume is calculated by equation 1:

$$\varepsilon = 1 - \frac{\rho_{app}}{\rho_P} \tag{1}$$

 ρ_P is the polymer density and ρ_{app} the apparent density of the extrudates.

The theoretical maximum porosity ε_{max} is obtained considering that all the dissolved CO₂ becomes gaseous inside the extrudate at ambient conditions and thus create porosity. It is then calculated by the following equation:

$$\varepsilon_{\max} = \frac{w_{CO_2}\rho_P}{w_{CO_2}\rho_P + \rho_{CO_2}(atm)}$$
(2)

 w_{CO_2} is the CO₂ mass fraction and $\rho_{CO_2}(atm)$ is the CO₂ density at ambient conditions.

Samples were also analysed with an helium pycnometer (Micrometerics AccuPyc 1330). Helium can reach open pores, so the density measurements allow to evaluate the closed porosity with the same equation 1.

To complete the characterization of the porosity structure, samples were examined by scanning electron microscopy (ESEM, FEG, Philips).

RESULTS

Results obtained with PHBV when testing the die and melt temperatures.

T h e	m a i	n ı	e s u l t	S	a r e	pre	esented	d i n	
Td (°C)	Te (°C)	porosity (%)	T2 (°C)	P1 (bar)	P2 (bar)	P3 (bar)	C (N.m)	Qpolym(g/mn)	
before die	die								
140	110	48.04	128	94	103	120	43	37.4	
140	120	47.36	129	91	96	106	43		
140	125	48.71	130	93	98	104	42		
140	135	45.32	131	91	94	97	43.9		
140	140	47.34	132	95	90	93	42		
135	110								
135	120	57.22	125	106	131	122	44		
135	125	55.48	126	102	122	118	42		
135	135	52.62	129	104	124	102	39	41.3	
135	140	47.21	130	96	109	98	41	44.3	
130	110	69.45	128	107	142	130	43		
130	120	61.46	125	106	131	114	44		
130	125	59.08	126	107	122	107	42		
130	135	63.35	123	107	129	115	40		
130	140	52.11	129.9	106	112	108	41	45.4	

. The varying operating conditions are given, which are temperature T_e in the die and T_d before the die, together with the resulting pressure evolution, torque, temperature T_2 measured before the die, and polymer flowrate measured at the die outlet. Rotation speed of the screw is fixed together with CO₂ concentration.

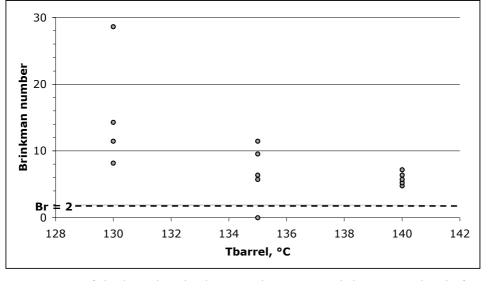
Table 1: Operating conditions and porosity results with PHBV experiments

Td (°C)	Te (°C)	porosity (%)	T2 (°C)	P1 (bar)	P2 (bar)	P3 (bar)	C (N.m)	Qpolym(g/mn)
before die	die							
140	110	48.04	128	94	103	120	43	37.4
140	120	47.36	129	91	96	106	43	
140	125	48.71	130	93	98	104	42	
140	135	45.32	131	91	94	97	43.9	
140	140	47.34	132	95	90	93	42	
135	110							
135	120	57.22	125	106	131	122	44	
135	125	55.48	126	102	122	118	42	
135	135	52.62	129	104	124	102	39	41.3
135	140	47.21	130	96	109	98	41	44.3
130	110	69.45	128	107	142	130	43	
130	120	61.46	125	106	131	114	44	
130	125	59.08	126	107	122	107	42	
130	135	63.35	123	107	129	115	40	
130	140	52.11	129.9	106	112	108	41	45.4

In order to investigate the temperature profiles, the Brinkman number, Br, was evaluated: it compares the extent of viscous heat generation relative to the heat conduction resulting from the imposed temperature difference (T_{Barrel} - $T_{product}$), $T_{product}$ being the temperature of the product measured before the die, that is T_2 . It is calculated according to

$$Br = \frac{\eta v_B^2}{\lambda_P \left| T_B - T_P \right|} \tag{3}$$

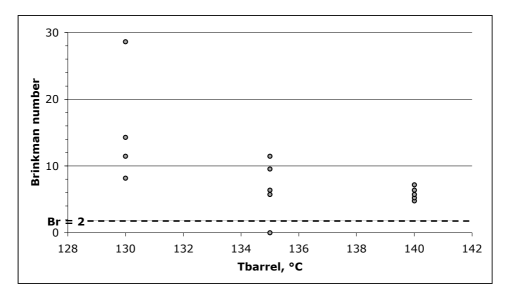
where v_B is the relative velocity of the barrel, η the molten polymer viscosity, λ_P , the molten polymer conductivity. A rheological study, which was previously investigated using a rheometer MARS, Thermo Scientific, showed a pseudo plastic behaviour and the viscosity was taken to be 1630 Pa.s for an average shear rate of 31 s⁻¹ in the pumping zone. λ_P of the solid polymer was measured at 150°C and found to be 0.205 W/m/K. The temperature difference given by was $|T_{d}-T_{2}|$. The values of Br presented are on



in function of

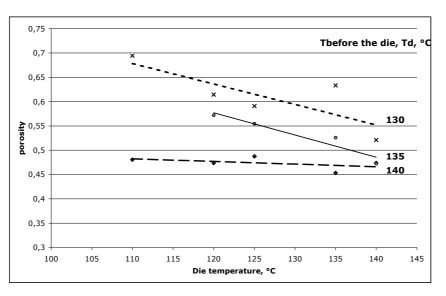
temperature of the barrel T_d in the pumping zone. Brinkman number is found to be larger than 2, which means that there is a maximum temperature, over the barrel one, at a position interm ediate between the barrel wall and the screw surface. Viscous dissipation is then important in our case.





Considering the porous structure, porosity is presented in function of the die temperature T_e at fixed melt temperature T_d on figure 4.

Figure 4: Porosity in function of the die temperature at fixed metering zone temperature (diamonds at T_{barrel}=140°C, circles at T_{barrel}=135°C, crosses at T_{barrel}=130°C)



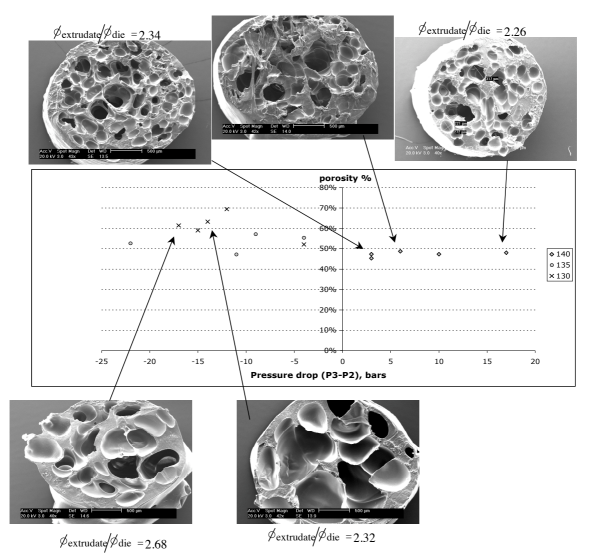
Considering temperature effects at fixed die temperature, porosity decreases when melt temperature increases. This last one influences the CO_2 solubility, which decreases as melt temperature increases. It was previously observed for polystyrene that, at a reasonably low temperature of polymer melt, there exists an optimal die temperature for which large volume expansion ratio are achieved by freezing the extrudate surface [6]. This effect is explained because more gas is retained in the foam at lower temperature and used for cell nucleation and growth. However, when the nozzle temperature was further decreased, the volume expansion ratio decreased because of the increased stiffness of the frozen skin layer. In our case, we didn't observe this phenomenon since we couldn't decrease more the temperature because of

torque limitation.

At fixed melt temperature and as die temperature increases, porosity decreases or even remains constant at melt temperature of 140° C. Note that the highest die temperatures correspond to the lowest pressures before die P₃, which are around 100 ± 7 bars. Indeed, when depressurization range decreases, nucleation rate decreases [10]. These explanations could be satisfying considering the porosity as a global characteristic but they may be incomplete considering the pore size and number.

Going further into the pore structure analysis, it appears that the pore structure differs greatly according to the pressure drop through the metering zone, P_3 - P_2 , as it is shown on figure 5.

Figure 5: SEM pictures together with porosity results in function of pressure drop between P3 and P2.



As P3 is greater than P2 and (P_3-P_2) is positive, CO₂ remains dissolved into the melt until the flow reaches the die where nucleation occurs. This corresponds to the case when melt temperature is 140°C, which is then equal or higher than the die temperature. In addition, no significative effect of the die temperature is then observed. As P3 is lower than P2 and (P_3-P_2) is negative, the mixture may desaturate, nucleation begin in the melt zone and then only growth take place in the die. This induces less pores but bigger ones, leading to a greater

porosity. This last one decreases with die temperature when transfer of $\rm CO_2$ out of the extrudate is enhanced.

Endly, it can be noticed that, in any case, porosities measured remain below the maximal one, which was estimated at 90%.

<u>Results obtained with BP when studying the influence of the distributive mixing and the</u> <u>CO₂ concentration on porous structure.</u>

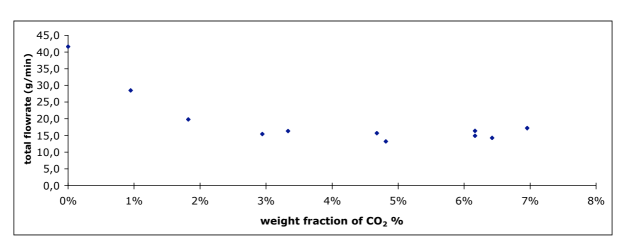
Experiments investigating the influence of carbon dioxide content and the influence of the mixing device were performed. Table 2 presents the operating conditions of each experiment.

Experiment number	Mixing device	Rotation Speed	Ta/b	Tc/d	Te	T1	T2	P1	Р2	Р3
		rpm	°C	°C	°C	°C	°C	bar	bar	bar
1	Classical screw	40	220	180	175	174	174	65-85	65-85	50
2	Pineapple mixer	40	240	180	180	183	176	50-70	110-140	60-70

 Table 2: Operating conditions in BP experiments

The first experiment was done with the former unmodified screw. We can see on Figure 6 that the total flowrate decreases with the increase of the CO_2 content and then reaches a constant value above 3% of CO_2 . This transition is also observed considering the structure aspect of the foams. Below 3% of CO_2 , the pores are rather small near the surface of the extrudate and are bigger in the center of the extrudate cross section. This can be explained by coalescence and growth enhanced by a slower cooling in the center than near the surface at the oulet of the die





Beyond the same amount of CO_2 (3%), we can see on Figure 7 a change in the porous structure. As the global porosity decreases, the closed one increases significantly and the mean pore diameter decreases. At the highest CO_2 concentration, the whole porosity seems to

be closed. Those porosity measurements are in agreement with SEM observations. The increase of CO_2 amount seems to increase the number of nucleation sites. In addition, the increasing cooling effect of the depressurisation through the die freezes the structure as soon as the outlet of the die is reached or even before. The evolution of the porosity with carbon dioxide concentration is consistent with other reported works.

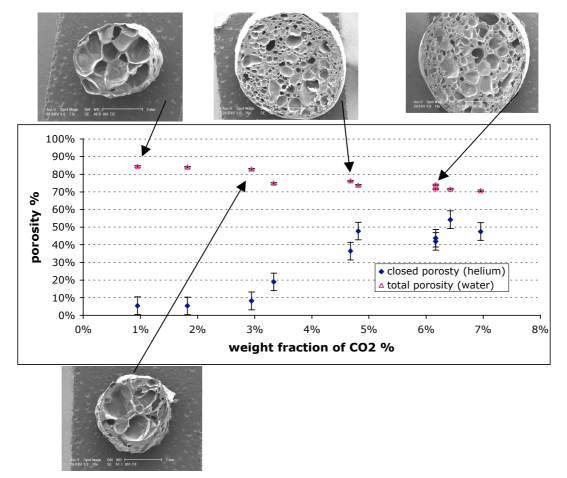


Figure 7 : Evolution of the porosity with CO₂ fraction. Exp1.

Comparing to PHBV results, at low concentration of CO_2 , foams obtained look rather the same. The negative pressure drop (P3-P2) might be the governing process leading to big pores. However, at high CO_2 concentration, porosity levels remain rather high and the porous structures seem to follow the classical statements from the literature: the governing mechanisms are those implied during the depressurization through the die and the cooling rate at the outlet of the die [10]. The negative pressure drop (P3-P2) through the metering zone doesn't seem to influence anymore the porosity results: mass and heat transfer kinetics may be significantly different due to the big amount of carbon dioxide.

The second experiment was conducted using the pineapple mixing element. This device greatly changes the pressure profile in the extruder. It increases the pressure gradient from P_2 to P_3 enhancing the shear flow towards the die. It seems to improve the mixing since higher contents of CO_2 can be incorporated into the melt without noticing any failure while processing the mixture. As we can see on Figure 8, the total flowrate is not disturbed by the addition of great amounts of carbon dioxide. Besides, the high porosity level is maintained until 15% of CO_2 . Above this amount, the total porosity is drastically decreased, as it is

shown on Figure 9 and the whole porosity is closed. There is far too much CO_2 and the induced freezing at the exit of the die prevents the pores from growing.

The pores obtained with the pineapple mixer are less homogeneous in size and are bigger than those obtained with the unmodified screw, at similar CO_2 contents. This could be explained by the pressure profile induced by the mixer compared to the unchanged screw. Pressure drop (P₃-P₂) is also negative but P₂ is much higher, which may induce a significative desaturation of the mixture. Hence, nucleation may take place in the metering zone of the extruder, as it is suspected for PHBV.

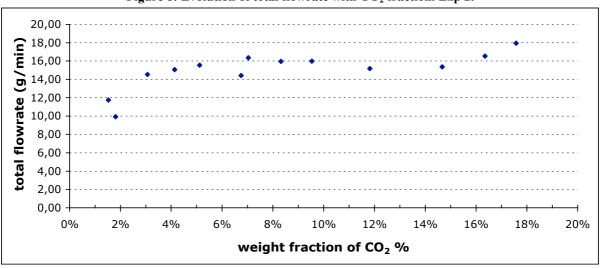
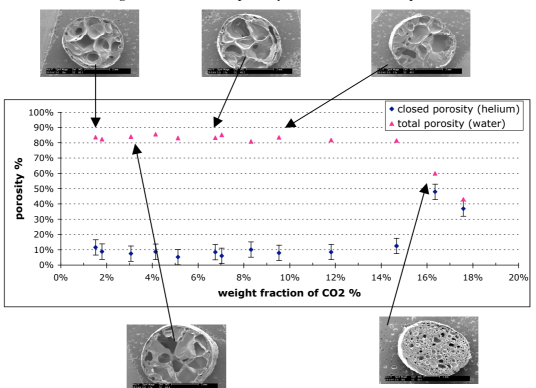


Figure 8: Evolution of total flowrate with CO₂ fraction. Exp 2.

Figure 9: Evolution of porosity with CO₂ fraction. Exp2.



Therefore, the use of this device may not be adapted to the production of foams with small and numerous pores. Obviously, it allows to incorporate greater amounts of CO_2 but the

porous structure obtained is not homogeneous and pores remain rather large. Besides, the high pressure drop induced in the mixer element zone enhances the shear flow towards the die together with a desaturation of the mixture. It also increases the counterpressure flow before this mixing zone forcing some carbon dioxide towards the hopper, which could explain some CO_2 leaks that we occasionally observed. Considering then the higher amounts of CO_2 and the raise in pressure P_2 at the upstream part of the mixing device, not all the carbon dioxide may be processed through the die. Therefore, if great amounts of CO_2 are required in the mixture depending on the process aim, possibly above the solubility value, an additional static mixer could be placed between the screw and the die. The CO_2 introduction could then be splitted between the actual one and another one placed just before the static mixer. This last one might improve again the mixing and allow incorporating carbon dioxide above the solubility value without flowing back through the hopper. Experiments are currently conducted to understand the impact of those mixing devices on the porosity and on the homogeneity of the extrudates.

CONCLUSION

PHBV foaming by extrusion assisted by supercritical fluid is thus feasible according to the temperature profile established. Porosity up to 70% was obtained. However, beyond the temperature profile effect, this foaming seems greatly influenced by the pressure profile, especially by the pressure drop through the metering zone just before the die. Shear flow may be enhanced together with desaturation of the mixture leading to early nucleation before the die. Comparing to the case when counterpressure flow is favoured and the mixture kept saturated, it leads to less pores but bigger ones and to a greater porosity. Acting on distributive mixing level by the introduction of a pineapple element and testing another biosourced polymer led to similar observations concerning the effect of the pressure profile. Foams with porosities up to 85% were obtained but the porous structure is not homogeneous and pores remain rather large. The high pressure drop induced in the mixer element zone enhances the shear flow towards the die together with a desaturation of the mixture. It also increases the counterpressure flow before this mixing zone forcing some carbon dioxide towards the hopper. Hence, this mixing element is not well adapted if dense and small pores are required. Endly, experiments with this biosourced polymer with the unmodified screw, so without the pineapple element, led to results consistent with literature ones. When saturation of the mixture is maintained all along the screw until the die is reached by maintaining a sufficient pressure level, the governing mechanisms are those induced by the depressurisation rate through the die and the cooling rate at the die outlet.

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