3D in situ observations of the compressibility and pore transport in Sheet Moulding Compounds during the early stages of compression moulding

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The properties of Sheet Moulding Compounds (SMCs) are altered by their porosity induced during the manufacturing or moulding of these prepregs. To characterise the pore scale mechanisms occurring during SMC flow, 3D synchrotron X-ray images were acquired during the compression of two uncured SMCs. For the high fibre content SMC (50 wt%), with a high porosity, pores mainly decreased in size and disappeared during SMC consolidation. These mechanisms were related to the elasticity of the fibrous networks and to the permeability of the porous phase, estimated using pore scale CFD simulation. For the standard fibre content SMC (29 wt%), with a lower porosity, the compressibility was limited and closed pores were transported towards the external surface of samples with non-affine motion, i.e., faster than the SMC flow and with tortuous trajectories. Besides, pores coalesced and decreased in size during compression. The size decrease was mainly related to the dissolution of pore gases.

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1. Introduction

Thanks to their excellent properties and their cost effectiveness, Sheet Moulding Compounds (SMCs) are thermoset prepregs that are broadly used in the aeronautical, automotive and electrical industries to manufacture body and structural parts. They are usually composed of a thermoset matrix (polyester, vinylester, epoxy) reinforced with 15 to 60 wt% of chopped fibres (glass or carbon fibre bundles, biosourced fibres) and are prepared in the form of 1–3 mm sheets [1]. The sheet processing consists of three steps: paste mixing, impregnation of the fibre network by the thermoset polymer paste, calendering and paste thickening. As for some other prepregs, SMCs exhibit a non-negligible amount of pores (≥1–20%) after their fabrication [2–6]. In addition, during their forming phase, i.e., compression moulding, initial pores can be transported, their size decreases and they can disappear [4,5,7–8]. Other pores can also nucleate or be introduced into the SMCs, e.g. by solvent boiling or squish flow phenomena at the flow front, respectively [5,9]. These intricate mechanisms are accompanied by the deformation of the fibrous reinforcement with complex volume variations. They can result in the appearance of pores which usually lower the physical and mechanical properties of moulded parts [10–12]. Among the challenges to overcome to optimise compression moulding of SMCs and the end-use properties of produced parts, understanding and modelling the aforementioned phenomena is a critical issue. To this end, various experimental studies were conducted to find how to reduce SMC porosity during compression moulding, e.g. by using vacuum assisted moulding, tuning the mould closing pressure or velocity and the mould temperature or by preheating SMCs before compression [5,12–14]. However, pore scale mechanisms are still not very well understood so that, to the authors’ knowledge, there is no proper description and model of pore kinetics and kinematics during compression moulding of SMCs. Consequently, to date, the compression moulding stage is generally numerically simulated assuming that SMCs are incompressible [2,3,15–20], which is a strong a priori assumption when considering the SMC initial pore contents.

It is worth mentioning that pore kinetics and kinematics was extensively studied for other prepregs, see for examples [21–24] among the numerous studies. These studies provide interesting guidelines to understand and model pore mechanisms that occur in prepregs as well as observations of the pore morphology. However, most of the studied prepregs do not exhibit in-plane flow during their forming stage, whereas SMCs are usually subjected to substantial in-plane flow during compression moulding [1]. This

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probably yields other pore transport mechanisms. In addition, in most of the aforementioned studies, 3D quantitative analyses of the pore morphology and evolution, e.g. using X-ray microtomography [25], are scarce. Recently, Centea and Hubert [23] used this technique to analyse the evolution of pores in out-of-autoclave prepregs. Thanks to X-ray microtomography, consolidation mechanisms were better understood. However, as the considered prepreg exhibited continuous fibre bundles, the in-plane flow of the prepregs was limited, whereas this flowing mode can be important in SMCs as fibre bundles are discontinuous. In addition, samples were imaged ex situ so that the issue of the in situ evolution of pores was not tackled. This is now possible using synchrotron X-ray sources, which enable the acquisition of 3D images at very fast scanning times and to perform real-time 3D in situ observations of the structure of composites during their deformation [26,27].

Thus, the objective of this study was to complete these former experimental studies to better characterise the pore kinetics and kinematics during compression moulding of SMCs. For that purpose, we studied the compressibility, the evolution of the content and morphology of pores as well as their transport in two uncured SMC formulations having standard and high fibre contents, respectively. Hence, a compression rheometer was specially designed and mounted in the synchrotron X-ray microtomograph of the ID19 beamline (ESRF, Grenoble, France). Then, rheometry experiments were performed while scanning prepregs at different compression stages. This method also enabled 3D in situ images of the evolving mesostructures to be reconstructed. These original data were combined with macroscale strains measurements that were simultaneously recorded during the compression experiments to give a relevant analysis of the early stages of compression moulding of SMC.

2. Materials and methods

2.1. Materials

Two industrial formulations of SMC with two very different fibre contents were used and prepared by MCR – Plastic Omnium (Tournon-sur-Rhône, France). The first SMC formulation, which was denoted by the letter S, is typically used to produce light-weight automotive semi-structural parts. It consisted of a polyester-based paste reinforced with 29 wt% of flat glass fibre bundles with a 25-mm length, approximately 0.05 mm in height and 0.5 mm in width. The rheological behaviour of this formulation was similar to that reported for other SMCs in previous studies [28,29]. The second SMC formulation is used to produce structural parts. It was denoted by H in the following sections and consisted of a vinyl ester-based paste reinforced with 50 wt% of glass fibre bundles having similar dimensions than those of the S formulation. For compression tests, five cylindrical samples were cut out of both SMCs. These samples were made of two stacked sheets, leading to initial sample heights \( h_0 \) approximately equal to 4 and 6 mm for the \( H \) and \( S \) SMCs, respectively. The initial diameter \( D_0 \) of the samples was chosen to acquire 3D images with the best spatial resolution, i.e., to extract relevant information on both the pore mesostructures and macroscale sample flow mechanisms. Thus, the diameter \( D_0 \) was set to 30 mm for the \( H \) samples and for one \( S \) sample (Table 1). The two other studied \( S \) samples had a lower initial diameter of 20 mm (Table 1) in order to analyse the onset of their in-plane flow at higher compressive strains. Further, it was observed that the sample macroscale surface and volume strains (next subsection) were very close to those measured during lubricated compression experiments (not shown here) using the same transparent rheometer as reported by Guiraud et al. [30] for samples with larger 55-mm diameter. In addition, it was also verified that the samples with 20-mm and 30-mm diameters exhibited similar pore content and morphology (Fig. 4(c)), in the initial state but also during compression: their typical size was sufficiently small with respect to the investigated sample diameters so that the results could be considered to be representative [31].

2.2. Compression tests with 3D in situ X-ray imaging

The compression experiments were designed to mimic what is observed in many flow situations during SMC compression moulding: out-of-plane consolidation and in-plane plug flow of the core of SMCs at moderate temperatures (≈40–100 °C), combined with intensive shear of hot (≈150 °C) paste-rich outer layers acting as low viscosity lubricants [8,9,32,33]. Thus, as a first step towards more realistic configurations, the SMC samples were subjected to lubricated simple compression loading at room temperature (25 °C) using a specially designed compression rheometer similar to that used in previous studies [34,35] but equipped with a load cell of higher capacity (2kN). The SMC samples were positioned on the lower moving compression platen (Fig. 1). The vertical motion of the lower platen was achieved using an electromechanical actuator and gearing (maximal translation velocity of 300 mm s\(^{-1}\)). The SMC samples were subjected to a very small pre-compression (0.017 MPa) to ensure the contact with the platen. After stress relaxation (≈3 min), the mechanical loading was applied from the actuator attached to the lower compression platen so that the compression experiments were performed at two constant velocities \( h \), i.e., 5 and 50 μm s\(^{-1}\), corresponding to two initial compressive strain rates \( \dot{\varepsilon}_c = h/h_0 \) of 0.001 and 0.01 s\(^{-1}\) along the direction \( e_z \). These strain rates correspond to those encountered by SMC during moulding of parts with a mid-plane that is not perpendicular to the principal compression direction (taking into account typical compression moulding velocities that range between 0.01 and 10 mm s\(^{-1}\)). From an industrial standpoint, these flow zones are considered to be critical since pores are more difficult to extract, even at high mould closing pressure. The propagation distance between the sample and the X-ray detector was set to 4 m, enabling 3D-image reconstruction (the so-called Paganin reconstruction mode) based on enhanced phase contrast [36]. The following acquisition parameters were also used: 1999 X-ray projections (beam energy of 45 keV) acquired with a 360° rotation of the micro-rheometer. Then, thanks to high speed PCO edge camera, a very short scanning time of 2 s for each 3D image was obtained. For the highest strain rate, only interrupted loading experiments were performed and 8 scans were taken. When a targeted sample height \( h \) was reached, the test was stopped (for 2 s), a scan was taken and the loading was continued until the next stage. For the experiments performed at the lowest strain rate, interrupted but also continuous scans were performed, and 6–9 scans were obtained. In the particular case of the S4 sample, 30 images were acquired without stopping the loading, enabling real-time in situ 3D observations. Three-dimensional images of the internal structure of the sample, with volumes of size 1698 × 1698 × 464 voxels and with a voxel size of 23.5 \( \mu \)m\(^3\) (Fig. 2(a) and (b)), were obtained using suitable reconstruction algorithms.

2.3. Image analysis and data post-treatment

Most of the image analysis was achieved using the softwares Fiji [37] and Matlab. Standard filtering operations (median and Gaussian filter) were used to obtain 3D grey scale images that enabled the macroscale strains of the compressed samples to be measured. The height \( h \) of each sample was precisely measured to estimate the Hencky compressive strain \( \varepsilon_{\text{h}} = \ln(h/h_0) \) (Fig. 2(a)). In addition, we also estimated the mean area \( \bar{A} \) of the sample in-plane
surface, *i.e.*, the mean sample surface in the \((e_1; e_2)\) plane. For that purpose, the sample areas within a dozen of in-plane cross sections were estimated, \(A\) being their mean value. Notice that for each slice the SMC area was measured using the function “Wand” of Fiji. Filling operations were also used to remove pores. Then, the sample surface strain \(e_S = \ln\left(\frac{A}{A_0}\right)\) and the sample volumetric strain \(e_V = e_S + e_{33}\) were calculated. Other image analysis operations were performed using the binarised images. As shown in Fig. 2 (c)–(f), the good contrast between the pore phase and the solid phases of the sample (enhanced by the Paganin imaging reconstruction) enabled easy manual thresholding and measurements of several descriptors for the pore phase. These analyses were achieved in cylindrical or cubic Regions Of Interest (ROIa) and (ROIb), respectively, depending on the used algorithms, as shown in Fig. 2(b):

- The volume fraction of pores \(\phi_p\) in the sample was calculated by counting the number of voxels of the pore phase and dividing it by the number of voxels of the ROIs.
- We also computed 2D mean porosity \(\bar{\phi}_{2D}\) maps along the plane \((e_1, e_2)\) of the sheets to observe qualitatively pore transport and pore size decrease during the sample compression, as illustrated in Fig. 3. Each pixel \(p(i,j)\) of the 2D colormaps corresponded to a porosity defined as the porosity of the subvolume (thickness of \(N\) voxels) given by the set of voxels \(v(i,j,k)\) of the corresponding initial 3D images \((k \in [1,N])\). The plugin “Pointpicker” of Fiji was also used to follow quantitatively the in-plane motion of the centre of mass in these images with respect to the centre of mass of the samples.
- The plugin “BoneJ” of Fiji [38] was used to perform connectivity analysis in the porous phase of ROIb. This plugin was used to detect closed pores in the images, *i.e.*, pores which were not connected with at least one edge of the ROIs. Then, the mean volume \(V_{cp}\) of the closed pores was calculated using the plugin “Particle Analyser” of Fiji. In addition, the volume fractions of closed \(\phi_{cp}\) and open pores \(\phi_{op} = \phi_p - \phi_{cp}\) in the samples were calculated.
- The anisotropy of the pore phase was estimated in ROIb by computing the mean chord lengths \(l_1, l_2,\) and \(l_3\) of the pore phase along the \(e_1, e_2,\) and \(e_3\) directions, respectively [39]. Then, the anisotropy ratios \(r_{12} = l_1/l_2, r_{13} = l_1/l_3\) and \(r_{23} = l_2/l_3\) were calculated to quantify the mean shape of pores.

The motion of several selected closed pores \(i\) along the \((e_1, e_2)\) was also analysed in one of the \(S\) samples. For that purpose, the 2D positions \(x_{cp}^{i0}\) of their centres of mass \(G_i\) were tracked from Fig. 3. Then, these positions were compared with the positions \(x_{cp}^{\text{exp}}\) that these pores would have followed assuming an affine motion with the SMC:

\[
x_{cp}^{\text{dir}} = x_{cp}^{0} + \int_{0}^{t} \dot{e}_2 \otimes e_2 dt ,
\]

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\]
where \( x_{c_i}^{\text{exp}} \) were the initial in-plane positions of the considered pores. For comparison, the deviations \( \Delta x_{c_i} = x_{c_i}^{\text{exp}} - x_{c_i}^{\text{aff}} / h_0 \) were also estimated.

### 2.4. Permeability estimation

To analyse and discuss the experimental results, the permeability tensor \( K \) of the porous media constituted by the fibre bundles and the paste matrix was calculated (Section 3) using (i) the 3D binarised images (ROIb) obtained during the compression of the two SMC formulations, (ii) the theoretical framework of the homogenisation method with multiple asymptotic expansions \([40]\), (iii) and pore scale fluid flow numerical simulation using the 3D images \([41,42]\). Practically, the Finite Volume software GeoDict was used to compute the permeability tensor \( K \) by solving the Stokes localisation problem derived from the homogenisation using 3D segmented images as calculation volumes. Details about this procedure are reported by Chalencon et al. \([42]\).

### 3. Results

#### 3.1. Initial mesostructures

Fig. 2. Initial grey scale slices of one \( S \) (a) and one \( H \) (b) samples. In (b), ROIa and ROIb are represented with dotted and continuous lines, respectively. (c, d) Histograms of the images (ROIb) of the upper slices shown in (a, b). (e, f) Examples of thresholded slices. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

The vertical and horizontal slices shown in Fig. 2, the two first 2D maps shown in Fig. 3 and the 3D views shown in Fig. 4 (a) and (b) show that both SMC formulations exhibited pores before compression.

For the \( S \) formulation, the initial average porosity \( \phi_{\text{p0}} \) was approximately 0.025 (Fig. 5(a)), with large variations between samples (from 0.015 to 0.055). These values were close to those previously reported for this type of SMCs \([4,5]\). In addition, this formulation exhibited a significant amount of closed and isolated pores since the initial closed pore content \( \phi_{cp0} \) was approximately 0.015 (Fig. 6(a)). The pores that were initially closed had rounded and ellipsoidal shapes with a moderate transverse isotropy and larger in-plane dimensions (Fig. 7(a)). It was verified that the transverse isotropy measured for each sample, made of several layers, was identical to that given when analysing only one SMC layer. The arrow in Fig. 4(a) and (b) shows that open pores were very flat.
Fig. 3. 2D maps showing the porosity averaged along the thickness of samples $\Phi_{p2D}$ at various compressive strains $\epsilon_{33}$: (a) 0, (b) 0.14, (c) 0.3, (d) 0.5 for the sample S4 and (e) 0, (f) 0.09, (g) 0.22, (h) 0.26 for the sample H1. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. 4. 3D views showing the evolution of pores at various compressive strains $\epsilon_{33}$: (a-b) 0, (c-d) 0.11, (e-f) 0.14, (g-h) 0.17 for the S4 sample. The equivalent diameter of the pores is colour-coded. The arrow points to an open pore between two SMC layers. The open diamond, square and circle put in the upper views illustrate pores that either disappeared or were expelled from the sample or coalesced, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Their potential origin could be the gases entrapped between the stacked layers of the specimen. In contrast, both the morphology and the content of the initial pore phase of the H formulation were completely different (Figs. 2(b) and 3(e)). The initial porosity was much higher than that of the S formulation, i.e., $\phi_0 \approx 0.23$ (Fig. 5(b)), and most of pores were open since $\phi_{cp} \approx 0.02$ (Fig. 6(b)). Fig. 2(b) shows that the matrix mainly impregnated the fibre bundles. It also formed mesoscale menisci at the vicinity of contact zones between fibre bundles, leading to complex morphologies of the porous phase, with networks made of connected and flat pores with intricate shapes.

3.2. Flow-induced strains and pores

3.2.1. Macroscale strains and porosity

During compression, the S samples flowed in the sheet plane (Fig. 3(a)-(d)). The flow was accompanied by a progressive and limited consolidation which tended to stabilise at high compressive
strains (Fig. 5(c)). This volume strain, i.e., \( e_{V} = \frac{1}{C_{0}} \), exhibited a small magnitude compared with the value of \( e_{33} \) reached by the compressive strain and was accompanied by a decrease in the sample porosity (Fig. 5(a) and insets in Fig. 6(a)). Also note that samples S4 and S5, which had almost the same initial porosity, also exhibited similar evolutions of their surface and volumetric strains (Fig. 5(c)), showing that the strain rate and its history had minor effects on these kinematic variables.

The compressions of the two H specimens were very different. Indeed, during the experiments, no in-plane flow was observed up to the 2-kN force that the micro-rheometer enabled (Fig. 3(e)–(h)) and the surface strain \( e_{S} \) remained close to 0 (Fig. 5(d)). Instead, noteworthy consolidation occurred as the sample volumetric strain \( e_{V} \) decreased down to \( C_{0}/25 \), thus leading to a progressive decrease of the sample porosity (Fig. 5(b) and inset in Fig. 6(b)). Fig. 5(b) and (d) clearly show that the consolidation mechanisms of the H specimens were not affected by the strain rate: surface strain, volumetric strain as well as the specimen porosity were nearly identical although these samples were deformed at two different initial compressive strain rates (Table 1).

### 3.2.2. Overall pore morphology

Fig. 6(a) shows that the open pores of the S formulation practically disappeared in the early stages of the compression, i.e., for \( |e_{33}| < 0.05 \). Fig. 4(b) and (d) shows that the related mechanisms corresponded to the closure of the thin space between stacked layers. Above this value, only closed pores remained in the sample and progressively disappeared while the compressive strain decreased. During this process, the mean pore volume \( V_{p} \) also gradually decreased (Fig. 8). At the same time, the transverse isotropy of pores exhibited a complex evolution (Fig. 7(a)). It firstly increased (from point ① to point ③) and then decreased (from point ⑥ to point ⑦).

During compression, the pore morphology of the H formulation evolved in a different way. Open porosity \( \phi_{op} \) remained in the samples up to very high values of \( |e_{33}| \) and followed the trend observed for the total porosity (Fig. 6(b)). In parallel, the closed porosity \( \phi_{cp} \) remained constant up to high values of \( |e_{33}| \) and then decreased. Finally closed pores disappeared in the last stages of the tests (Fig. 6(b)). At the same time, the mean volume of closed pores \( V_{cp} \) showed a sharp and constant decrease (Fig. 8). Combined with the two previous remarks, this demonstrates that during compression open pores progressively disappeared while an increasing number of smaller and smaller closed pores were forming. Lastly, it is also interesting to note that the initial transverse isotropy of the porous phase progressively decreased and tended to be isotropic (graph and micrograph in Fig. 7(b)).

### 3.2.3. Pore scale flow mechanisms

The 3D images in Fig. 4 and the surrounding 3D views in Fig. 7 reveal that the aforementioned trends were induced by more complicated pore scale mechanisms. For instance, the transverse isotropy that both formulations exhibited resulted in averaging the complex shape of pores due to their flow/motion and compression through the network of fibre bundles. For example, Fig. 7 shows that the considered closed pore which had an initial rounded transversely ellipsoidal shape was progressively stretched, confined and sheared. It progressively gained a complicated anisotropic shape during compression. In addition, the porosity decrease shown in Figs. 5(a) and (b) and 6 resulted from various mechanisms that were identified thanks to a close inspection of the 3D images in Fig. 8.
Fig. 4. For example, during compression, closed pores exhibited compaction but also 3D motions with, in the case of the S formulation, noticeable in-plane flow which led some pores (i) to coalesce and/or (ii) to be expelled from the compressed samples, mainly through their lateral surfaces (Fig. 4(a), (c), (e), (g)).

4. Discussion

It is worth recalling that the observations carried out in this study, as well as the following discussion, are limited to objects or heterogeneities with a size that remains above the spatial resolution of the 3D images, i.e., above approximately twice the voxel size (23 3 μm3). Keeping this important point in mind, the aforementioned results show that combining compression tests with 3D images enabled the compressibility of SMCs and their related pore scale mechanisms to be suitably characterised.

4.1. Origins of pores

As already reported in previous studies [4–6], our observations confirm that prepregs such as SMCs usually exhibit pores after their fabrication. Pores in SMCs can be induced during the polymer paste preparation and mixing, the paste impregnation through the network of fibre bundles via calendering operations, and the ageing/maturating the produced sheets before compression moulding. In this study, it was additionally shown that both the content and the morphology of pores depended on the volume fraction of fibres: the higher the volume fraction of fibres, the higher the initial volume fraction of pores and the more connected the porous phase. Two major reasons could explain this trend. First, as for other porous media, increasing the fibre content in fibrous networks usually leads to reducing their permeability [43,44]. Thus, this phenomenon could limit the impregnation of SMCs by the polymer paste [45]. Unfortunately, due to the low contrast between the matrix and the fibre bundles, we could not estimate the permeability tensor of the fibrous phase for the H samples (Fig. 2(b)). Thus, it was not possible to compare it to that of the S samples. Possible solutions to circumvent these effects would consist in either lowering the paste viscosity during the impregnation, e.g. by increasing its temperature, or modifying the surface properties of the fibre bundles (sizing) or the matrix to optimise the wetting of the fibrous network by the matrix. The second important reason is related to the fibre content in the prepregs. Increasing the fibre content by compacting the fibrous networks, e.g. during the calendering of SMCs, is known to increase their elastic energy [46,47]. Hence, the elastic energy stored during the calendering could be at the origin of springback effects, i.e., deconsolidation of the fibrous networks and consequently nucleation of pores in SMCs. The higher the fibre content after the calendering phase, and the higher the stored elastic energy, the springback effects, and the pore contents before compression. Finally, the 3D images also revealed the presence before compression of a significant amount of pores between SMC layers (arrow in Fig. 4). This type of porosity was mainly induced by the wavy surfaces of prepregs that were induced by the crenelated shapes of calendering rolls [1,6]. It was shown that this type of porosity mainly disappeared in the early stages of compression, but some residual pores remained during compression. A possible solution to minimise them would consist of a last calendering operation using flat rolls.

4.2. Compressibility of SMCs

In accordance with the trends that have already been reported in previous studies [8], it was shown that the initial porosity of SMCs systematically resulted in macroscale compressible flows of SMC specimens during compression, regardless of the fibre content. The magnitude of the SMC compressibility depended on the initial pore content and was very high for the high fibre content H formulation. In addition, the experimental data suggested that macroscale and mesoscale compressibility mechanisms could be considered to be independent of the strain rate. Finally, depending on the type of pores, i.e., closed vs. connected pores, the experimental results also showed that the SMC flow was different (see also the next subsection). As mentioned in the introduction, the compressibility is not taken into account in rheological models for prepregs such as SMCs (or GMTs) or other similar unsaturated concentrated fibre suspensions to the best of our knowledge, although it was shown in this study that compressibility could drastically alter the flow of these materials (Fig. 5(c) and (d)). Thus, these results constitute an interesting database linking macroscale flow phenomena to pore scale compressibility mechanisms that can be used to build relevant constitutive theories which could be implemented in simulation software for compression moulding processes.

4.3. Related mesoscale flow mechanisms

Two types of flow mechanisms were identified and correlated with the types of pores in SMCs, namely open or closed pores.

4.3.1. Open pores

The first flow mode mainly occurred for tests performed with the H specimens, at least up to compressive strains [ε33] below 0.2. It was also observed to a less extent during the first compression stages of the S specimens, i.e., for compressive strains [ε33] below 0.05. In these situations, the main SMC flow mode was transverse consolidation with pronounced compressibility effects. Since the studied SMCs contained a sufficient amount of connected pores (Fig. 6(a)), the permeability of the porous phase was large enough so that air/gases in pores could be expelled from the SMCs during compression, leading to a progressive closure of pores. However, during compression, the extraction of air/gases from the open pores possibly became more and more difficult and complex flow mechanisms took place. This is illustrated in Fig. 9 that shows the evolution of the diagonal components of the permeability tensor K of the porous phase for the H samples during compression (the non-diagonal components were negligible and were thus

Fig. 9. Evolution of the components $k_{ii} (\text{circles}), k_{22} (\text{triangles}), k_{33} (\text{square})$ of the pore phase permeability tensor K as a function of the pore content $\phi_p$ for the two tested H samples (black and with marks).
omitted for clarity). The permeability components underlined once again the transverse isotropy of pores (the out-of-plane permeability component being different from the in-plane ones) and exhibited a sharp decrease during compression, but were not zero up to the two last compression stages. It is also interesting to note that during the first stages of compression, the flow in pores should rather occur in the sheet plane as the in-plane permeability components were higher than the out-of-plane components (Fig. 9). This scenario should be reversed at the end of compression (Fig. 9). From a practical standpoint, these results are be interesting in order to optimise vacuum assisted compression mouldings, e.g., by optimising the placement and the activation of the void extractors.

### 4.3.2. Closed pores

The second flow mechanism occurred when the SMCs, such as the S formulation, principally contained closed pores. In this case, the compressibility and the related consolidation mechanisms of SMCs were much weaker than those obtained for the first flow mode, leading to substantial in-plane flow, i.e., perpendicular to the compression axis $e_j$. Closed pores exhibited complex kinetics and kinematics with coalescence, shrinkage up to full closure (see next paragraph) and motion up to the external surfaces of the samples, as shown qualitatively in Fig. 4.

To better understand these intricate mechanisms, Fig. 10(a) shows the deviation $\Delta \xi_{ij}$ of the pore placement $\xi_{ij}^{exp}$ with respect to the affine prediction $\xi_{ij}^{aff}$: obviously, $\Delta \xi_{ij}$ reached very high values. In addition, as emphasised in Fig. 10(b), the in-plane trajectories $\xi_{ij}^{exp}$ and $\xi_{ij}^{aff}$ exhibited two major differences: (i) the experimental in-plane velocities of closed pores were much faster than those predicted by the affine assumption, and (ii) most of the experimental trajectories exhibited noticeable angular deviations from the affine radial trajectories. The first difference is similar to that observed during the transport of bubbles in capillaries \[48,49\], where bubbles are prone to flow faster than their suspending fluid, due to shear stress and pressure gradient effects and to related bubble deformation mechanisms. Westerberg et al. \[50\] also reported the same trend by squeezing between two parallel platens grease with air bubbles which exhibited radial velocities faster than the squeezed grease. In the present lubricated compression experiments, the phenomena were similar but much more complex due the presence of fibre bundles; our results suggest that closed pores were transported in tortuous capillaries induced by the fibrous networks.

It is also interesting to notice that closed pores were also subjected to severe shrinkage and sometimes disappeared (Figs. 4 and 8). This shrinkage was linked with the pore pressure $p_j$ that was estimated according to the following procedure. We first estimated the macroscopic compression stress $\sigma_{33} = F/A$ recorded during the experiments, $F$ and $A$ being the compression force and the macroscopic in-plane surface of the samples, respectively. Then, the following assumptions were stated: (i) the surface tension effects at the pore-SMC interfaces were negligible because of the very high shear viscosity of SMCs ($\approx 10000$ Pa s), (ii) dense SMCs (without pores) behaved as incompressible materials, and (iii) the SMC extra-stress tensor was purely deviatoric (rigorously, this assumption may be questionable taking into account the high anisotropy of SMCs \[29\], but was expected to provide a good order of magnitude). Hence, when open pores disappeared, i.e., for $\varepsilon_{33} < -0.05$ for the specimen S4 (Table 1), the mean pressure $p_j$ in closed pores of mean volume $V_{ij}$ was estimated as $p_j \approx p_a - \sigma_{33}/3$, where $p_a$ was the atmospheric pressure.

The resulting evolution of $p_j$ during the compression of the specimen S4 is shown in Fig. 11 as a function of $\varepsilon_{ij}$ for $\varepsilon_{33} < -0.05$: only a small increase in the pore pressure $p_j$ occurred during compression, i.e., from 0.15 to 0.17 MPa. In parallel, another estimate $p_j^{exp}$ of the closed pore pressure was provided using the values of $p_a$ and $V_{ij}$ for $\varepsilon_{33} = -0.1$ for the considered sample, and the mean number of moles $\Pi$ of gases (probably a mixture of air and styrene) contained in the pores and calculated from the ideal gas law ($g_l$). Then, assuming that $\Pi$ remained constant during compression, i.e., there was no molecule adsorption/diffusion at the interface between the pores and SMC, $p_j^{exp}$ was computed both from the ideal gas law and the evolution of $V_{ij}$ plotted in Fig. 8. Obviously, Fig. 11 shows that $p_j$ and $p_j^{exp}$ exhibited completely different evolution so that $p_j^{exp}$ was one hundred times higher than $p_j$ at the end of the compression. Consequently, this result tends to prove that approximately 99% of the gas molecules initially inside closed pores disappeared during the compression-induced pore shrinkage, most of them being probably dissolved in the SMC matrix. Note that rather similar phenomena have been reported by Leterrier and G’Sell \[22\] for the dissolution of gases during the compression of glass-reinforced thermoset composites.

![Fig. 10](image-url) (a) In-plane motion of the centres of mass of several closed pores manually identified during the compression of the sample S4 (from Fig. 3) and compared to the affine motion given by Eq. (2). (b) Related deviation $\Delta \xi_{ij}$ as a function of the compressive strain $|\varepsilon_{33}|$. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
The compressibility and the porosity of two SMCs having high and moderate fibre contents, respectively, were investigated using synchrotron X-ray microtomography. This technique enabled a better understanding of the mechanisms that are related to the macroscopic scale as well as the kinetics and the kinematics of pores in these prepregs during their compression. Thus, for the first time, in situ and 3D images of the flow of several SMC samples as well as their evolving porous mesostructures were acquired. In the initial state, SMC samples were generally porous and contained transversely isotropic pores. The volume fraction and the morphology of the pores depended on the fibre content. The porosity was elevated and the pores were mainly open at high fibre content. On the contrary, at moderate fibre content, the porosity was moderate and the pores were mainly closed. Optimising the sizing of the fibre bundles to improve their wetting properties, together with minimising the elastic energy stored in the fibrous reinforcement during the calendaring phase could provide two proper solutions to minimise the SMC initial porosity. In addition, this study showed that the compressibility of both studied SMCs was associated with a decrease in the sample porosity, both being practically independent of the strain rate. As these phenomena could be particularly important for high fibre content SMC formulations, they should be taken into account in future rheological models for the numerical simulation of the compression moulding process. Finally, the 3D in situ images also provided useful information to better understand pore compressibility and transport mechanisms. The first primary mechanisms occurred for connected pores, i.e., mainly for high fibre content SMC. During compression, these pores reduced in size and, simultaneously, the gases that they contained could flow through the open pore network and be expelled from the samples. The second mechanisms were related to closed pores and were mainly observed for moderate fibre contents. During compression, closed pores decreased in size and sometimes coalesced. Combining the 3D images and the compression stress recorded during the experiments, it was shown that the pore shrinkage was related to the dissolution of pore gases (air, styrene) into the SMC polymer paste. Besides, these closed pores were also expelled from the centre of samples to the sample external surfaces, following tortuous trajectories and exhibiting displacements that were faster than that of the bulk SMC flow. This phenomenon was attributed to the squeezing effect exerted by the complex fibre bundle network on the SMC paste and the closed pores. The proper modelling of these complex mechanisms would require further investigation of other representative mechanical loadings, e.g., plane strain compression, tensile or shear loadings, and higher testing temperatures.

References

[16] Hill; 1996


