Experimental investigation of porosities evolution in a bonded assembly by means of X-ray tomography

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ABSTRACT
Structural bonding is a very advantageous technique for lots of application fields such as aeronautics or marine industry, which require both advanced performances and lightweight structures. Nonetheless, adhesive joints are often subject to bonding defects: kissing bonds, uneven polymerisation, or pores within the material, for instance. These pores, depending on their sizes and distributions, could jeopardise the mechanical strength of the assembly. Moreover, it is legitimate to hypothesise that these voids in the medium could be influenced by the application of a mechanical stress. In order to investigate this assertion, bonded samples are loaded by various tensile stress levels, and the pores within the joint are visualised and characterised using in-situ X-ray microtomography. This paper deals with the evolutions of various quantities such as the number of pores or their volumetric ratio along with the increasing load and with the diverse phenomena (nucleation, growth, coalescence, etc.) experienced during the testings. These results are extracted from the microtomographic data using a custom processing tool, whose parameters and performances are discussed.

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1. Introduction

Structural bonded joints[1] represent today a promising technique due to its main advantages such as: obtaining lightweight and complex structures, its simplicity of implementation, the possibility of unification of elements of different nature (in particular composite and metal), and providing a more homogenous distribution of the loading state in the two bonded elements. In recent years, a large number of studies[2–6] were carried out in the bonded joints field, both in terms of experimental characterisation and numerical modeling of the mechanical behaviour of adhesive.

Experimental characterisation of adhesive requires the use of different types of tests dedicated to this area. Among the most common, one may
include: (i) single lap joint test (SLJ), (ii) standardised thick adherend shear test (TAST) or (iii) modified Arcan test.\textsuperscript{[5,7]}

SLJ is characterised by the simplicity of its samples and of the test, which only requires a simple test machine. However, this test is accompanied by significant stress concentrations at the ends of the overlap length, which may cause a significant scattering of the experimental results.

An alternative to this problem is the modified Arcan test, which diminishes drastically the stress concentrations close to the free edges of the adhesive. Also this test can generate different loading states in the joint such as: pure tensile, tensile-shear, pure shear or compressive-shear. However putting into practice the modified Arcan test involves a certain degree of difficulty, which makes it less usable in industrial environments. An alternative for the modified Arcan test is the modified Scarf test.\textsuperscript{[8]} This test setup allows to generate tensile or tensile-shear loads on an adhesive joint. The sample tested in this case is simple and can be used on standard tensile test machines without any additional devices. Moreover, this test is perfectly adapted for the fatigue tests investigation in the industrial environment.

Full field measurement of the displacement or strain by optical non-contact measurement systems, are increasingly being used in order to obtain the kinematic information necessary to characterise the mechanical behaviour.\textsuperscript{[9]} In most cases, the information obtained using non-contact measurement systems are obtained on the surface of the sample. Notwithstanding, it is necessary to get information inside the sample for the understanding of failure mechanisms and of their evolutions during the test.

In the last decade, several studies have been conducted, which highlight the major advantages of observations using X-ray tomography (XRT).\textsuperscript{[10–12]} This tool is very advantageous because it allows to access the details of the microstructure of a material without its destruction. An original methodology is proposed\textsuperscript{[13]} to estimate 3D displacement fields from pairs of images from X-ray computed micromicroscopy. Contrary to local approaches, a global approach is followed herein that evaluates continuous displacement fields. Fast tomography combined with local crack driving force analysis has been employed to analyse crack-tip stress/strain singularities in an aluminium alloy.\textsuperscript{[11]} The application of fast microtomography has made possible to observe real crack initiation and propagation behaviours without intermediate unloading. The anisotropy of fracture toughness in AA2139 (AlCuMg) alloy sheet has been investigated via synchrotron radiation computed tomography of arrested cracks in Kahn tear test pieces for different loading cases.\textsuperscript{[14]} More recent work\textsuperscript{[15]} in the field of bonded joints have investigated the quantitative penetration of three coldset wood adhesives under hydraulic pressure into different types of modified wood using fluorescence microscopy and the results were compared to these of a previous study without pressure on adjacent wood samples. The three-dimensional visualisation of
the penetration of the adhesive into heat-treated Scots pine was also examined by X-ray tomography. X-ray microtomography is used in\textsuperscript{[16]} to visualise the distribution of melamine-urea-formaldehyde adhesive in the wood composite, particle board, and to examine changes in adhesive distribution on wood particles (flakes) before and after pressing. Also X-ray computed microtomography (XCT) was used to analyse the 3D adhesive penetration behaviour of different wood-adhesive bondlines.\textsuperscript{[17]}

However, there are only few papers that analyse quantitatively the information that can be achieved using X-ray tomography in the field of structural bonded assemblies. It should be noted that the damage mechanisms are linked mainly by the presence of defects such as: pores, lack of glue, the presence of several phases in the adhesive structure, \textit{etc.}

The present paper focuses on the quantitative analysis at the micrometric scale of bonded joints (epoxy adhesive and aluminium alloy). The main objective is to obtain the 3D microstructure of the adhesive after polymerisation process (\textit{i.e.} to identify the phases in the glue and their distribution) and to monitor the evolution of these phases when the sample is subjected to a tensile loading. In the first section, the experimental procedure will be presented, followed by the development and the validation of the adequate processing tool. Finally, the results will be outlined and discussed in the last section.

2. Experimental procedure

2.1. X-ray tomography setup

Since its first applications in the medical field,\textsuperscript{[18]} X-ray tomography has been the subject of many studies both from a theoretical and applied point of view. A brief description is therefore given here in order to introduce the concepts, which will be used further in the text. Various experimental setups can be used to perform X-ray tomography, but the basic principles of the technique remain the same (see Figure 1). Tomography measurements are based on the variation of the linear X-ray attenuation coefficient, hereafter written $\mu$, through the volume of a material. For a perfectly isotropic material this $\mu$ coefficient is a constant with respect to the spatial coordinates $(x, y, z)$. However, for non-homogenous material, such as porous environments, $\mu$ is a function of these coordinates. It is possible to obtain the spatial distribution of this $\mu$ coefficient by sending a source X-rays beam through the bulk of the studied sample, from different angles, and by collecting the transmitted beam. Hence, the sample is mounted on a rotator included in the tomograph chamber, and for each angular step, a X-ray beam is emitted, attenuated by the internal structure of the sample, and gathered by a detector. The attenuation phenomenon is described by a Beer – Lambert law depending on $\mu(x, y, z)$:
where $I_0$ represents the source intensity (emitted), $I$ is the detected intensity (transmitted and collected by the X-ray detector) and $x$ represents distance along the transmission path $[x_0; x_{\text{max}}]$.

A series of $N$ radiograms, or radios, is obtained through the complete acquisition process (Figure 1). A reconstruction algorithm is then used to build the 3D internal structure of the sample (i.e. the spatial distribution of $\mu(x, y, z)$) from the radiographs series.

Those algorithms may be based on two different approaches: the software may either solve a set of linear equations to compute $\mu(x, y, z)$ (algebraic approach) or use a backprojection of the detected intensity by a Fourier transformation (analytical approach). Although the analytical approach is faster to compute, it requires a full dataset with no missing views from the radiographs series. This method is referred to as filtered backprojection reconstruction. For more details about these techniques and about their theoretical backgrounds, see.\cite{10,19} The results presented in this paper were obtained by a filtered backprojection approach. Those algorithms depend on the X-ray source used to perform the measurements, being either a synchrotron or a laboratory tomograph, for the emitted beam itself depends on the underlying technology. For laboratory devices, the emitted beam is produced into a cone shape, rather than being constituted by parallel rays (which is the case for synchrotron X-ray sources). This has to be taken into account during the Fourier transformation backprojection.\cite{10} Although the formalism of the algorithm shall not be
discussed here, the reader could find more information about this particular matter in various reference works.\textsuperscript{[10,19,20]} The reconstruction step provides the 3D structure of the observed sample (Figure 1), defined with a given voxel size, in a similar fashion as pixels constitute a picture. It is worth noticing that the voxel size is related to the spatial resolution of the measurement, but is not equivalent. An empirical relationship between those quantities may be established, the spatial resolution being then twice the voxel size.\textsuperscript{[10]} It is obvious that the spatial resolution of the experimental set-up depends not only on the voxel size, but also on various external parameters relative to the whole experimental set-up, and that the resolution must be in compliance with the characteristic sizes of the microstructure defects to observe. Some laboratory tomographs allow the operator to perform \textit{in-situ} measurements during mechanical and/or thermal loadings.\textsuperscript{[21]} These kinds of experiments are though complex to handle due to the measurement artifacts it may produce. It is hence possible to access the microstructure of the sample, which is now a function of time, as opposed to classical, one-shot experiments delivering a unique radios series. The acquisition time for each radiographs series comes into play and defines the time resolution of the measurement, which has to be negligible when compared to the characteristic time of the observed phenomena. The following results in this paper are gathered using such an experimental set-up.

2.2. Sample preparation

The samples consist of two Aluminium (2017A) substrates adhesively bonded by an epoxy resin layer of a given thickness. A simple, butt joint-like, geometry is designed as presented in Figure 2a. The area of the bonded section is set to \(6 \times 6 = 36\, \text{mm}^2\). A non-uniform stress state is then expected within the bulk of the adhesive joint, along with stress concentration effects due to the influence of the edges of the substrates during the mechanical loading.\textsuperscript{[1]}

![Figure 2. Samples and bonding apparatus.](image-url)
The assemblies are obtained by bonding the substrates described above with the epoxy adhesive Huntsman\textsuperscript{TM} Araldite 420 A/B. The adhesive thickness is set to a value of 400\(\mu\)m and is controlled by a system of spacers. This system, and the specially designed corresponding apparatus, are presented in Figure 2a. This system enables both the control of the thickness of the adhesive joint and the correct alignment of the substrates. A clear space is also preserved in order to clean the adhesive joint once the substrates are assembled, and thus to obtain a neatly defined sample.

In order to guarantee the good adhesion between the adhesive layer and the substrates, a classical (chemical and mechanical) surface treatment is performed on the bonding surfaces to remove any greasy byproducts that may remain after the machining of the substrates and to create an adequate surface roughness for the desired mechanical interlocking between the substrates and the adhesive. In a first step, the substrates are placed in an acetone bath for several hours to degrease the surfaces. They are then treated with grade 180 sandpaper to remove the remaining impurities and to eliminate any oxide layer that could have formed due to the exposition of the Aluminium to the ambient environment. Moreover, this mechanical treatment creates the aforementioned surface roughness required for a good adhesion. Lastly, the surfaces thusly treated are cleaned with acetone to remove the impurities and Aluminium particles created with the mechanical treatment. The adhesive joint is then prepared by applying the mixed epoxy resin and hardener on both substrates, which are assembled and installed in the apparatus (Figure 2b) carefully adjusted to obtain the desired joint thickness. The samples are finally placed in a Secasi Technologies 100/60 thermal enclosure at 115\(^\circ\)C for 1 hour to ensure a fully polymerised material.

Due to the very limited volume of the adhesive joints, it is assumed that the thermal field in the material is homogeneous, and that there is no significant influence of possible curing degree gradients in the adhesive. It can be seen in Figure 2a that threaded holes are machined at each end of the samples to allow the positioning of the rods necessary to the installation of the samples in the clamping jaws of the tensile machine to be used. Those holes are also used during the gluing process to control the relative positioning of the substrates.

2.3. Test procedure

The samples are placed inside a Phoenix\textsuperscript{TM} VtomeX tomograph equipped with a Varian\textsuperscript{TM} Paxscan X-Ray detector featuring a resolution of 1920 \(\times\) 1536 pixels. This detector outputs a 14-bits coded grayscale picture of the attenuation. The chamber of the tomograph includes a 3kN tensile machine, which allows to perform X-ray tomography measurements while applying a mechanical load to the sample (Figure 3).

The voxel size obtained using this experimental set-up is 6 \(\mu\)m \(\times\) 6 \(\mu\)m \(\times\) 6 \(\mu\)m. The corresponding spatial resolution is expected to be in compliance with the
characteristic size of the pores to be observed. Also, the adhesive contains 0.16% (weight ratio, according to manufacturer data) of glass beads, whose diameters are about 80 µm, i.e. more than 10 times greater than the resolution of the device. The measurements are assumed to occur for a quasi-static state of the samples, and therefore the corresponding characteristic time overshadows the acquisition time for a radiographs series. Hence, it is concluded that the experimental set-up is well-suited to the measurements to be performed. In order to reconstruct the full observed volume, the acquisition is performed along with a rotational motion of the sample. A total of 912 pictures are taken throughout the rotation, for a complete acquisition time of 12 min. These series are then used as previously mentioned by an algorithm based on the results presented in[^4], in the perimeter of an analytical approach of the computation of the spatial field \(\mu(x, y, z)\). The full volume of the samples, visualised by means of the variation of the values of the attenuation coefficient \(\mu(x, y, z)\), is then broken into a series of slices (Figure 5).

An amount of 96 slices is obtained for the adhesive joint only, from the tomographs series. As already mentioned, the tensile test machine included in the tomograph chamber allows us to perform these measurements while applying a mechanical loading to the samples. The decision has been made to incrementally increase the applied load with a 200 N step (Figure 8). The ramps between the steps are displacement-controlled (0.5 mm/min). A stress-strain curve for the considered material is given in Figure 4b along with the stress states corresponding to each aquisition step (Figure 4a). In addition, the elastic properties of the adhesive are presented in Table 1.

The process goes on until the failure of the sample. It is clear from Figure 4a that stress relaxation phenomena are occurring during these steps, especially for the high loads. For each step, the corresponding displacement is therefore maintained for a certain time, in order to let these phenomena occur. Otherwise, the measurement would be disturbed by the on-going relaxation, due to the similarity between the acquisition time and the relaxation characteristic time. By considering significantly longer load steps, it is possible for the load to reach an asymptotical value and hence to obtain a relatively steady state.
Figure 4. Mechanical loading and properties of the adhesive.

Figure 5. Reconstructed volume of the sample obtained from tomography investigation.

Table 1. Elastic parameters of the adhesive\(^{[22]}\).

| Parameter | Value       |
|-----------|-------------|
| $E$ [MPa] | $2000 \pm 50$ |
| $\nu$ [-] | $0.41 \pm 0.01$ |
During the 12 minutes necessary to complete a full rotation of the sample. The aforementioned asymptotical value is harder to reach in a reasonable time the higher the load is, and may even not be reachable in some cases (loads greater than 800 N, Figure 4a). This may lead to blurry acquisition data, as those measures are critically sensitive to any disturbances, which cause significant artifacts. It is then possible to acquire a snapshot of the 3D microstructure of the adhesive joint for several loads. In other words, given the adequate post-processing tools, it is possible to follow the evolution, along with the applied load, of the microstructure, in a non-destructive in-situ fashion.

3. Postprocessing tool

A time series of radiographs sets is delivered by the experiments detailed above. The reconstructed volume of the central part of the sample for the initial state of load is shown in the Figure 6a. Those raw grayscale images, display 3 visible phases (see Figure 5b–d): the adhesive, the air pores and the glass beads introduced by the manufacturer.

The raw data is initially encoded in 16 bits. In order to reduce the volume of data and thus facilitate post-processing steps, the raw 16-bits data were converted to 8-bits grayscale levels. This step results in a lighter dataset and in faster processing steps without damaging excessively the quality of the data to observe. The data obtained with X-ray tomography were then post-processed in order to extract the following information: the spatial distribution and rate of each phase as well as their evolution during the different load applied. The complete study of the microstructure with respect to the applied load requires the development of a processing tool, which will: (i) eliminate unwanted artifacts from the

Figure 6. Slice of raw data.
reconstructed slices of adhesive and extract the mean intensity; (ii) filter the
signal to remove the noise and (iii) detect the microstructural entities observed
(here, pores and glass beads) from a grayscale intensity threshold.

This processing tool should allow to automatically detect the requested
microstructural entities from the raw signal presented in Figure 6b.

3.1. Creation of synthetic microtomographic data

In order to design the processing tool, a preliminary study is performed using
synthetic microtomographic data. The main advantage of this technique is
that every property of the microstructure is accurately known, since it is
imposed. The grayscale levels of the voxels for a 2D slice taken in this
synthetic volume are given in Eq.2.

\[
S_{zj}(x, y) = I_{0,zj}(x, y) + y_i \cdot p_i(x, y) + b_{zj}(x, y)
\]  (2)

where \( S_{zj}(x, y) \) are the grayscale levels of the slice \( j \) in the \( Z \) direction with
\( j \in [1, 65] \), \( I_{0,zj}(x, y) \) is the mean grayscale level, \( y_i \) is the contrast for each
phase \( i \), \( p_i(x, y) = N_i \) with \( N_i \in [0, 2^8 - 1] \) are 8-bits grayscale levels for each
phase \( i \), and \( b_{zj}(x, y) \) is a 8-bits grayscale level correspondong to Gaussian
measurement noise.

The intensity of the measurement noise is assumed to be independant of
the position. Therefore, the standard deviation of the Gaussian distribution
used to generate the measurement noise is the same in the entire volume.
The value of the standard deviation \( \sigma \) is experimentally obtained, by sub-
tracting two microtomographic acquisitions of the same volume. The proper-
tries of the generated volume are displayed in Table 2.

The spatial distribution of the generated pores can be visualised in Figure 7a.
Moreover, the distribution of their corresponding diameters is available in
Figure 7b. One may notice that the diameters distribution is not perfectly
uniform. This is due to the non-overlap condition for the pores used during
the generation of the volume. Therefore, a few of the randomly generated
diameters (mainly the largest ones) were programmatically modified or dis-
carded in order to comply with this constraint.

| Table 2. Synthetic data properties. |
|-----------------------------------|
| Property                          | Value                                           |
| Size                              | 960x960x65 voxels                               |
| Pores number \( n_{ph} \)         | 3114 (non-overlapping)                          |
| Pores ratio \( p_{th} \)          | 9.69%                                           |
| Pores diameters                   | Uniformly distributed between 1 and 35 voxels   |
| Noise intensity                   | Standard deviation \( \sigma = 9.62 \)          |
3.2. Postprocessing tool architecture

In order to characterise the pores located in a volume, it is necessary to segment the tomographic data. To do so, a processing tool is designed and programmed using Matlab™ software. The scheme of treatment of the raw data is shown in Figure 8.

3.2.1. Preprocessing

As it may be seen in Figure 5b–d, the radiograms include a large part of the substrates and some blank space around the samples. Hence, the reconstructed volume also contains these elements, resulting in some useless volume slices (with only Aluminium showing). The adhesive being absent from these pictures, they are removed from the dataset. By doing so, it is possible to reduce the number of slices to be analysed from several hundreds to 65.

3.2.2. Subtraction of $I_0$

The mean grayscale intensity is assumed to follow a polynomial expression, as in Eq.3. The degree of this polynomial surface is left undetermined in Eq.3. In practice, $m = 3$ was enough for the study presented in this paper. The value of $m$ may be adjusted accordingly with the shape of $I_0$.

Figure 7. Artificially generated volume for known pores amount, pore ratio and pores locations.

Figure 8. Processing tool architecture.
The coefficients $c_{i,j-i}$ in Eq.3 are identified using the grayscale levels of each reconstructed volume slice (Figure 9a). The obtained values of $I_0$ are then subtracted to the grayscale levels of the volume (Figure 9b), in order to obtain a flattened signal (Figure 9c and d).

### 3.2.3. Filtering and thresholding

The filtering and the thresholding steps are designed to smooth the signal (i.e. remove as much noise as possible from the flattened signal) and to identify precisely the transition between the adhesive and the pores in terms of...
grayscale levels (Figure 10). To correctly tune the corresponding parameters ($t_f$ and $n_s$, see Figure 8), an optimisation loop is performed using Matlab$^\text{TM}$. The filter used in the process is a classical 2D median filter,$^{[23]}$ characterised by its size $t_f$. The main advantage of this formulation is that it is edge-preserving$^{[24]}$ (i.e. it will preserve the edges between the adhesive and the pores), which is paramount to accurately detect the pores included in the adhesive.

The threshold $n_s$ is trivially defined as the minimal grayscale level shift to consider to detect a microstructural phase switchover. Those values are then adjusted in order to minimise an error function, characterising the performance of the tool. It is possible to consider two different shapes for this error function: $Err_{M1}$ which takes into account only the difference between the pore ratios $p_i$ (method M1, Eq.4) and $Err_{M2}$ which takes into account the difference between the pore ratios $p_i$ and the difference between the number of pores detected $n_{p_i}$ (method M2, Eq.5).

\[
Err_{M1}(t_f, n_s) = \frac{p_{th} - p_{obt}(t_f, n_s)}{p_{th}}^2 \tag{4}
\]
\[
Err_{M2}(t_f, n_s) = \alpha_1 \left[\frac{p_{th} - p_{obt}(t_f, n_s)}{p_{th}}\right]^2 + \alpha_2 \left[\frac{n_{p_{th}} - n_{p_{obt}}(t_f, n_s)}{n_{p_{th}}}\right]^2 \tag{5}
\]

where $p_{th}$ and $p_{obt}$ are respectively the imposed pore ratio and the detected pore ratio for a given $(t_f, n_s)$ couple, and $n_{p_{th}}$ and $n_{p_{obt}}$ are respectively the imposed pores number and the detected pores number for a given $(t_f, n_s)$ couple.

The method M1 defines the error between the generated pores field and the detected pores field as a function of the pore ratios whereas the method M2 defines a more extensive error function as a function of both the pore ratios and the pores amounts (Eq.4 and 5). In the M2 formulation, a couple $(\alpha_1, \alpha_2)$ is

**Figure 10.** Filtering of the synthetic data.
introduced so it is possible to assign different weights to each one of the two terms if need be. By default, these values are set to 0.5, so as to obtain an equivalent weight for each term in Eq.5. The optimisation loop is performed in a first stage for the M1 approach, and delivers a first set $(t_f, n_s)_{M1}$ identified as the parameters corresponding to the lowest error value ($t_{fM1} = 3$ and $n_{sM1} = -15$) easily pinpointed on the error map (Figure 11a). The loop is then repeated for the M2 approach, which delivers two error maps (Figure 11b and c) instead of one, as two terms are used to compute the error. By combining the minimum valleys obtained through the M2 method, it is possible to compute the global error for this approach (Figure 11d). Due to the removal of $I_0$ from the baseline signal (Figure 9c and Figure 9d), a negative value is obtained for $n_s$. It is therefore not a 8-bits grayscale level anymore, but a grayscale intensity.

(a) Error map obtained for the M1 method as a function of $t_f$ and $n_s$

(b) Error map relative to the number of pores for the M2 method as a function of $t_f$ and $n_s$

(c) Error map relative to the pore ratio for the M2 method as a function of $t_f$ and $n_s$

(d) Optimal parameters for the M2 method

**Figure 11.** Identification of the $(n_s, t_f)_{M2}$ set.
From the intersection of these domains (Figure 11d), it is possible to extract a second parameters set \((t_f, n_s)_{M_2}\), which outputs not only the correct pores volume fraction, but also the correct pores amount. From these observations, it can be concluded that the processing tool designed in this paper allows for a correct evaluation of the pore distribution in an adhesive layer, in terms of both spatial distribution and radius distribution.

It is clear from Figure 12a that the M1 method is inefficient to accurately detect the correct number of pores, especially for the lowest diameters. This inefficiency tends to fade along with the increase of the equivalent diameters of the pores to be detected. The M2 method displays results of overall higher quality, and is less prone to be disturbed by very small pores. This method

![Graphs showing comparison between methods M1 and M2 in terms of pore detection, effect of noise on pore ratio, and number of pores.](image)

**Figure 12.** Comparison between the results delivered by the M1 and M2 approaches for 5 different noise draws.
allows the user to accurately detect the voids in the adhesive, with a fairly high accuracy in terms of both number and equivalent diameter.

The tool, successfully calibrated here, was used on a generic adhesive volume including a pores field whose diameters are randomly distributed in a given range representative of the actual configuration. To be perfectly in compliance with a real experimental case, these diameters should be normally distributed. However, this slight inadequacy is expected to be of negligible influence, as it seems from Figure 12a that the designed tool outputs the correct number of pores for nearly every equivalent diameter. The only difference between a normal and a random distribution being the number of entities for each equivalent diameter, one should expect that the processing tool would correctly estimate the voids field for a different distribution.

### 3.3. Influence of measurement noise on pores detection

Although it appears that the two approaches presented give similar results, they can be evaluated in a more extensive fashion by studying the influence of the picture noise level on the pore field detected. To perform this comparison, the two resulting tools are applied to five cases with five noise draws (Figure 12b and 12c).

From Figure 12b it is obvious that even though the two methods presented previously give similar results in terms of pores volume fractions (which is slightly underestimated due to the voxelisation of a continuous media), only the M2 approach is able to deliver a correct estimation of the pores amount (Figure 12c) in a given volume. It is then concluded that the M2 method should be used, for \( t_f = 4 \) and \( n_s = -14 \). The obtained results are summarised in Tables 3 and 4, where \( \bar{i} \) and \( \sigma_i \) are respectively the mean value and the standard deviation of the quantity \( i \).

**Table 3.** M1 and M2 methods results compared to the theoretical pore ratio.

|            | \( p_{th} \) [%] | \( p_{obt} \) [%] | \( \sigma_{p_{obt}} \) [%] |
|------------|------------------|-------------------|---------------------------|
| Theoretical| 9.59             | –                 | –                         |
| M1         | –                | 9.522             | 1.05 \( \times 10^{-3} \) |
| M2         | –                | 9.516             | 1.80 \( \times 10^{-3} \) |

**Table 4.** M1 and M2 methods results compared to the theoretical number of pores.

|            | \( n_{th} \) [-] | \( n_{obt} \) [-] | \( \sigma_{n_{obt}} \) [-] |
|------------|------------------|-------------------|---------------------------|
| Theoretical| 3114             | –                 | –                         |
| M1         | –                | 8427.2             | 107.7                     |
| M2         | –                | 2930.1             | 12.7                      |
Even though fairly similar results are obtained in terms of pore ratio $p_{obt}$ with both methods (Table 3), it is clear from Table 4 that the M1 method is not able to accurately detect the correct number of pores in the artificial datasets. Quantitatively, the error on the detected number of pores is above 170% using M1, whereas it is only of 6% using M2. This is explained by the fact that M1 tends to detect erroneous pores for small diameters (Figure 12a).

4. Results and discussion

Given the extensive use of X-ray tomography in materials science since the early 1990s, and due to the large amount of published works based on this method,\[10\] it is accepted that this experimental technique provides fairly good results. Consequently, the measurements performed using this technique are considered to be valid, and do not require any further characterisation.

4.1. Evolution of the pores number and of the pore ratio with respect to the applied load

From the observations made regarding the two methods presented above, it was chosen to use the M2 methodology, with an optimised $(t_f, n_s)_{M2}$ couple equal to $(4, -12)$.

An immediate output of the processing tool presented in the previous section is the pores-to-adhesive volume ratio. It is then possible to track the evolution of this quantity along with the increase in the applied load (Figure 13a).

In addition, by means of the M2 method defined by the Eq.5, it is possible to obtain a fairly accurate estimation of the number of pores included in the adhesive layer, which may be plotted versus the increasing load as well (Figure 13b).

It appears from these data that the mechanical solicitation applied to the sample may lead to significant changes in the microstructural properties of the adhesive. Both these quantities seem to follow a trend defined by two distinct domains: $F \leq 800$ N, for which the evolution seems fairly linear with respect to the applied load; and $F \geq 800$ N, for which the evolution becomes non-linear as a function of the applied load, with a sudden increase in pore volume fraction, and with a sudden decrease in pores number.

It is possible to validate the results presented above by verifying that the developed method outputs the correct glass beads ratio, which is given by the manufacturer.

It appears from Figure 13c that this quantity remains almost constant, as it should be since there is obviously no external glass beads input. It is possible to compute the mean $\overline{p_{beads}}$ of these values, to be compared to the manufacturer data:
These numerical values are fairly consistent with the numbers provided by the manufacturer regarding this adhesive, since the adhesive theoretically contains 0.16% of beads (in terms of volume).

Finally, the distribution of the diameters of the detected glass beads may be computed. This distribution is shown in Figure 14.

### 4.2. Evolution of the diameters distribution with respect to the applied load

Most of the pores are quasi-spherical, and thus may be characterised by their equivalent diameter. More specifically, it is possible to compute the...
distribution of those equivalent diameters in the adhesive layer, and to track the evolution of this distribution along with the increase in the applied load.

It appears from Figure 15a–c that the equivalent diameters distribution variations are somehow negligible for a load below 800 N. Starting from this load state (Figure 15b–c), the evolution becomes much clearer, as the distribution, which roughly follows a Gaussian shape, tends to widen.

To better visualise this trend, it is possible, for each distribution displayed in Figure 15, to fit a Gaussian distribution function (Eq.6) to compute the mean equivalent diameter \( \mu_d \) and the standard deviation \( \sigma_d \).

\[
f(x) = \frac{1}{\sigma_d \sqrt{2\pi}} \exp \left[ -\left( \frac{x - \mu_d}{\sigma_d} \right)^2 \right]
\]

(6)

This approach results in the data presented in Figure 15d. A clearer trend can be seen in Figure 15d, which points out that even for the relatively low loads \( F < 800 \) N, the microstructure of the adhesive undergoes slight, but nonetheless visible, transformations. Both the mean and the standard deviation of the distributions seem to follow the same trend. As it was expected from the data in Figure 13 and Figures 15a-c, from \( F = 800 \) N and higher, the distributions start to change significantly. The increase in size of the detected pores may be explained by two factors: (i) the increasing load tends to open and expand the pores (Figure 16d) and (ii) the increasing load may trigger coalescence phenomena, which cause neighboring pores to merge (Figure 16a–c). These hypotheses are directly linked with the observations formulated regarding Figure 13 and Figures 15a-c, as they seem to be the most plausible explanation for these phenomena.
It is fairly easy to notice that some entities merged due to the application of the mechanical load, especially where the initial pores were already close to each other. This is perfectly supported by the diminishing number of pores detected by the processing tool, as it is shown in Figure 13b. Such merger phenomena were expected, as they have already been reported for other materials.\cite{21,25}

The visualisation of these simultaneous phenomena may be performed on the reconstructed volume using \textit{FiJi} software,\cite{26} as presented in Figure 17. The global increase in volume of the pores in the adhesive is easy to visualise in Figure 17. This suggests that it is a phenomena which occur in the entirety of the joint, and not only in specific, predisposed locations. Moreover, it is possible to see in Figure 17 that there are a few locations where the pores seem to be more concentrated. The adhesive bond is therefore weaker in these spots, and one may hypothesise that such concentrations

\textbf{Figure 15.} Effect on an increasing mechanical load on the equivalent diameters distribution.
of pores could trigger mechanical failure mechanisms (microcracking, stress concentrations, localised plasticisation, etc.) when stressed, and lead to the premature ruin of the assembly. Finally, the pores in Figure 17 may be divided into two groups, depending on their size and incidence: (i) the most common pores are rather small (a few tens of micrometers in terms of equivalent diameter) and they are created during the mixing of the adhesive, (ii) and the less usual pores (in particular 2 specimens in Figure 17, easily identifiable), one order of magnitude bigger than the others.
(a few hundreds of micrometers in terms equivalent diameter), which are
created during the spreading of the adhesive with the spatula on the adher-
ends. This last type of pores is the most critical threat to the mechanical
strength of the joint, due to the size they feature. One might expect that if
several of these large, but uncommon, pores are located within a certain
sphere of influence of each other (still to be determined), this could lead to
an important and localised decrease in mechanical properties of the joint and
once again be the cause of the unexpected mechanical ruin of the assembly.

4.3. Sphericity of the pores

The pores detected by the tomography measurements display for the most
part a sphere-like shape, therefore a description based on the sphericity
seems to be the best approach to characterise the evolution of their general
shape. The sphericity $\Psi$ of an object is, as introduced by Wadell in 1935$^{[27]}$,
defined by the expression in Eq.7:

$$\Psi = \frac{\pi^{\frac{1}{3}}(6V_p)^{\frac{2}{3}}}{A_p}$$  \hspace{1cm} (7)

where $V_p$ stands for the volume of the entity and $A_p$ for its area.

It is obvious from Eq.7 that the sphericity varies theoretically between 0
(the studied entity being a plane) and 1 (the studied entity being a perfect
sphere).

The volume and the areas of the pores being quantities easily accessible
from the tomography data, the sphericity of each pore may be computed, for
various load states. The resulting data are plotted in Figure 18.

The technique employed in this paper is then able to detect the variation
of the shape of each pore, caused by the mechanical solicitation applied to the
sample. It seems that the overall shape of the points clouds remains fairly the
same, the most significant change in Figure 18 being the increase in equivalent
diameters of the pores, which tends to shift these points clouds to the
right. Most of the computed sphericities are located around 1, which corro-
borates the observation according to which most of the pores are quasi-
spherical.

One can see in Figure 18 that some points are located above the theoretical
limit of 1, especially for the lowest equivalent diameters. This may be due to
the effect of the voxelisation during the measurements, as the voxel size is of
6 m. For very small objects, this may induce seemingly incorrect sphericity
values, due to areas miscalculations.$^{[28]}$
5. Conclusion

Pores are commonplace defects in adhesive joints, formed during the mixing of the adhesive and during the bonding step. As such, adhesively bonded samples were made in order to investigate the pore state within the bulk of their joints. The influence of tensile loadings on this pore state was characterised using in-situ X-ray microtomography, until the mechanical failure of the samples.

In order to process the tomographic datasets, a processing tool was designed and calibrated on artificial datasets. This calibration was performed using optimisation algorithms to tune the parameters of this processing tool: the size of kernel of the median filter, and the grayscale threshold used to identify the pores in the reconstructed volumes. Two methods were proposed for this parameters optimisation, and their performances were discussed. It was found that the best approach was to optimise the parameters of the tool using both the pores number and the pore ratio.

Finally, it was experimentally proven that the application of a mechanical stress on the adhesive bonds has a significant influence on the pores located in the medium. Once a certain stress is reached, coalescence phenomena are triggered, resulting in an abrupt decrease in pores number. Moreover, due to the effect of the mechanical stress, the pores tend to expand, resulting in an increasing pore volumetric ratio. These changes in microstructure could be viewed as indicators that damage is occurring in the material. This is confirmed by the fact that the rate of growth significantly increase shortly before the fracture of the samples. Moreover, coalescence is occurring roughly for the

Figure 18. Sphericity of each detected pore, for different load levels.
same stress levels, leading to the possible relationship between these observed phenomena and mechanical damage.

These results could give new ways of improvements for models developed to predict the damage and the failure of the adhesively bonded joints. Indeed, it seems that the damage is due to a cavitation process (increase in the size of existing pores or creation of new voids) and the failure is due to the coalescence of these pores/voids. Thus, the effects of the triaxiality should be included in the damage model instead of only the out-of-plane stresses as commonly considered.

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