Adherend thickness effect on the tensile fracture toughness of a structural adhesive using an optical data acquisition method

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Abstract

Adhesive bonding is nowadays a serious candidate to replace methods such as fastening or riveting, because of attractive mechanical properties. As a result, adhesives are being increasingly used in industries such as the automotive, aerospace and construction. Thus, it is highly important to predict the strength of bonded joints to assess the feasibility of joining during the fabrication process of components (e.g. due to complex geometries) or for repairing purposes. This work studies the tensile behaviour of adhesive joints between aluminium adherends considering different values of adherend thickness (h) and the double-cantilever beam (DCB) test. The experimental work consists of the definition of the tensile fracture toughness (G_{IC}) for the different joint configurations. A conventional fracture characterization method was used, together with a J-integral approach, that take into account the plasticity effects occurring in the adhesive layer. An optical measurement method is used for the evaluation of crack tip opening and adherends rotation at the crack tip during the test, supported by a Matlab® sub-routine for the automated extraction of these quantities. As output of this work, a comparative evaluation between bonded systems with different values of adherend thickness is carried out and complete fracture data is provided in tension for the subsequent strength prediction of joints with identical conditions.

Keywords:
Aluminium and alloys
Experimental testing
Joint design
Fracture

1. Introduction

In order to increase the efficiency of structures and reduce their weight, adhesively-bonded joints can be used over fastening, riveting or other traditional joining methods. Different joint configurations can be applied in structures, such as single-lap, double-lap, stepped and scarf. Many other configurations exist, with specific advantages, like the joggle-lap joints, applied for instance to join fuselage parts in aircraft, reinforcement doublers, peel joints and L-section joints. Adhesives are progressively being applied in structures of several branches of engineering, which also makes certification-related issues more important, especially in industries such as aeronautics [1]. Because of this, it is highly important the availability of robust predictive techniques that can reliably be used as design tools, to allow minimization of experimentation during the design stages of products and structures without compromising the design process [2,3]. This will allow assessing the feasibility of joining during the fabrication process of components (e.g. due to complex geometries) or joining as a repair method. A large number of predictive techniques are currently available, ranging from analytical to numerical, using different criteria to infer the onset of material degradation, damage or even complete failure. Initially, the prediction was performed by theoretical studies as those of Volkersen [4] or Goland and Reissner [5], which had a lot of embedded simplifying assumptions, by comparing current stresses with the allowable material strengths. Many improvements were then introduced, such as the assumption of elasto-perfectly plastic adhesive proposed by Hart-Smith [6], but these analyses usually suffered from the non-consideration of the material ductility, which is highly relevant because of stress gradients. Fracture mechanics-based methods took the fracture toughness of materials as the leading parameter for material selection. These methods included more simple energetic or stress-intensity fracture techniques that required the existence of an initial flaw in the materials [7]. More recent numerical techniques, such as cohesive zone models (CZM), combine stress criteria to account for damage initiation with energetic, e.g. fracture toughness, data to estimate damage propagation [8]. This allows to consider the distinct ductility of adhesives and to gain

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accuracy in the predictions. All of these fracture toughness-dependent analyses rely on an accurate measurement of $G_{IC}$ and $G_{ICc}$ (shear toughness). CZM in particular can accurately predict damage growth in structures if the fracture laws are correctly estimated [9]. These fracture laws are based on the values of cohesive strength in tension and shear, $t_s$ and $t_f$, respectively, and also $G_{IC}$ and $G_{ICc}$ [10]. These parameters that cannot be directly related with the material properties measured as bulk, since they account for constraint effects (in the case of adhesive joints, the constraints are caused by the adherends). Although these parameters do not have a clear physical significance, they are able to accurately reproduce the behaviour of the materials in a macro scale point of view that is quite accurate [2]. The estimation of these fracture parameters is generally accomplished by performing pure tension or shear tests. Regarding $G_{IC}$, the DCB test is the most suitable, due to the test simplicity and accuracy [11]. The typical $G_{IC}$ estimation methods are based on linear-elastic fracture mechanics (LEFM) and require the continuous measurement of the crack length ($a$) during the test. However, $G_{IC}$ of adhesives with large scale plasticity is not accurately characterized with LEFM methods since the assumed stress fields at the crack tip vicinity are not accurate [12]. More recently, methods that do not require the measurement of $a$ were developed, based on equivalent cracks and including the plasticity effects around the crack tip [13]. As it was described by Suo et al. [14], in the presence of large-scale plasticity, $J$-integral solutions can also be employed for accurate results. The $J$-integral is a relatively straight-forward technique, provided that the analytical solution for a given test specimen exists for the determination of $G_{IC}$ or $G_{ICc}$. The most prominent example is the DCB specimen, for which $J$-integral solutions are available, either for loading by pure bending moments [15] or the standardized and moment-free tensile loading [16]. It is also possible to estimate the tensile CZM law. Carberger and Stigh [17] computed the CZM laws of adhesive layers in tension and shear using the DCB and end-notched flexure (ENF) tests, respectively, considering $0.1 \leq t_A \leq 1.6$ mm ($t_A$ is the adhesive thickness). The rotation of the adherends was measured by an incremental shaft encoder and the crack tip opening by two linear variable differential transducers (LVDT). Ji et al. [18] studied the influence of $t_A$ in DCB joints on $t_s$ and $G_{IC}$ for a brittle epoxy adhesive. $G_{IC}$ was measured by a direct technique. For the measurement of the adherends rotation, two digital inclinometers with a 0.01° precision were attached at the free end of each adherend. The normal displacement at the crack tip was measured by a charge-coupled device (CCD) camera.

Regarding the fracture parameters of adhesives, different studies showed that these are similar as bulk and in a joint for brittle adhesives, since the yield zone ahead the crack tip is practically nonexistent [19]. Contrarily, when speaking about moderately-to-highly ductile adhesives, the fracture parameters are not invariant to the joint geometry [20,21]. This occurs because of the different degree of restriction to the development of this yield zone within the adhesive layer and also premature adhesive fracture because of excessive adherend yielding near the adhesive layer. This brings issues about the transfer of small specimen (test) results to real-life and complex structures. Actually, the majority of industrial structures bonded by adhesives consist of thin sheets between 1 and 3 mm thick, whilst the fracture behaviour of adhesives is mostly characterized within considerably thicker joints [22]. Pardoen et al. [23] discussed the two types of constraint effects that affect the fracture toughness of adhesive layers in bonded assemblies: external and internal constraint effects. External effects deal with the varying states of deformation of the adherends, which alter the stress and strain distributions of the adhesive layer. For instance, Wang et al. [24] found that $G_{IC}$ of adhesives measured in peel tests varies upon the thickness of the peeling arms. Internal constraint effects are $t_A$ and specimen width ($B$), which also influence the size of the yield zone and stress and strain distributions in the adhesive. As an example, the increase of $t_A$ can change the yielding conditions from small-scale to full plastic [25]. In the work of Giannis et al. [26], two different sealants for aircraft fuel tanks were tested by a modified peel test, and it was shown for both adhesives that $t_A$ largely increased the peel resistance between 2 and 4 mm. Because of these issues, taking into account of these constraints requires a precise simulation of the states of deformation of both the adhesive and surrounding structure. One of the methods that allow this to be modelled is CZM, by considering the adhesive layer modelled by cohesive elements [27].

Most of the published work addressing the influence of $h$ examines directly its influence on the failure loads, either static [28] or fatigue [29] and, in general, increasing $h$ has shown to improve the strength. Some studies associated this behaviour to the reduction of peel and shear peak stress at the overlap edges of bonded structures [30]. This actually occurs, but other phenomena are also on the basis of these differences in strength. Actually, a few studied showed variations of $G_{IC}$ by modification of the structures thickness. In the work of Mangalirai et al. [31], symmetric and unsymmetric DCB specimens were experimentally tested with different values of $h$ (by considering 8, 16 or 24 plies of carbon–fibre adherends). The static tests showed a large improvement of $G_{IC}$ between composites with 8 and 16 plies. Devitt et al. [32] equally used the DCB test to investigate this effect, and found a 9% increase in the value of $G_{IC}$ of bonded joints made of glass–epoxy composites by duplicating the number of plies of the adherends. From these studies, it is clear that the differences take place at relatively low $h$ values. Since most bonded joints are made between thin adherends/sheets, the understanding of how $h$ affects the fracture toughness is highly relevant.

In this work, the value of $G_{IC}$ of adhesive joints between aluminium adherends is studied, considering different values of $h$ and the DCB test. A conventional fracture characterization method was used, in comparison with a $J$-integral approach, that take into account for the plasticity effects occurring in the adhesive layer. An optical measurement method is used for the evaluation of crack tip opening and adherends rotation at the crack tip during the test, supported by a Matlab\textsuperscript{\textregistered} sub-routine for the automated extraction of these quantities.

2. Experimental work

2.1. Characterization of the materials

The adherends were cut from a high strength aluminium alloy sheet (AA6082 T651) by precision disc cutting. This material was characterized in bulk tension in previous works by the authors [33,34] using dogbone specimens and the following mechanical properties were obtained: Young’s modulus ($E$) of 70.07 ± 0.83 GPa, tensile yield stress ($\sigma_y$) of 261.67 ± 7.65 MPa, tensile failure stress ($\sigma_f$) of $324 \pm 0.16$ MPa and tensile failure strain ($\varepsilon_f$) of $21.70 \pm 4.24\%$. The two-component polyurethane adhesive SikaForce\textsuperscript{\textregistered} 7888, selected for this work, was formerly tested in the work of Neto et al. [35]. The bulk specimens were tested in a servo-hydraulic machine to obtain $E$, $\sigma_y$ and $\varepsilon_f$. The DCB test was selected to obtain $G_{IC}$ and the ENF test was used for $G_{ICc}$. The collected data of the adhesive is summarized in Table 1. To be noted that the obtained values of $G_{IC}$ and $G_{ICc}$ were obtained in partial adhesive failure conditions, which is important to notice when comparing $G_{IC}$ with the results obtained in this work.
2.2. Joint geometries

Fig. 1 represents the geometry of the DCB specimens. The dimensions of the specimens are: total length \( L = 160 \text{ mm} \), initial crack length \( a_0 \approx 40 \text{ mm} \), \( h = 1, 2, 3 \) or \( 4 \text{ mm} \), \( B = 25 \text{ mm} \) and \( t_a = 1 \text{ mm} \). The bonding process consisted in grit blasting with corundum sand, debris cleaning with acetone and assembly in a steel mould for the correct alignment between the adherends. To guarantee a uniform \( t_a \) value, calibrated spacers were inserted between the adherends. The required sharp pre-crack was assured using a 0.1 mm thick razor blade between calibrated bars at the crack tip. After applying the adhesive and assembling the specimens, these were given a full cure before testing. The spacers were removed and the adherends sides were sprayed with white brittle paint, to allow an easy identification of \( a_0 \), and a printed scale was glued in both adherends to aid the \( a \) measurement or input data for the digital correlation technique. Twenty-four specimens were tested (six for each configuration) at room temperature and 1 mm/min in an electro-mechanical testing machine (Shimadzu AG-X100) with a load cell of 100 kN. Each test was fully documented using a 18 MPixel digital camera with no zoom and fixed focal distance to approximately 100 mm. This procedure allowed obtaining the crack tip opening (\( \delta_{ct} \)) and rotation (\( \theta_a \)), necessary for the \( J \)-integral method. The correlation of the mentioned parameters with the load–displacement (\( P–\delta \)) data was done by the time elapsed since the beginning of each test.

3. Estimation of \( G_{IC} \)

It is known that, when the adhesives are ductile, as it occurs with the Sikaflex® 7888, LEFM methods are inaccurate [36], even though considering techniques with correction factors to account for plasticity (e.g. ASTM D3433-99:2005 and BS 7991:2001). Thus, the compliance-based beam method (CBBM), which accounts for the damage zone ahead of the crack tip, and the \( J \)-integral, were considered for the present study.

3.1. Compliance-based beam method

The CBBM was initially selected to measure \( G_{IC} \) of the adhesive [13]. It is a relatively straightforward but robust method, based on an equivalent crack, and it only depends on the specimen’s compliance during the test. Applied to the DCB test specimen, it gives

\[
G_{IC} = \frac{6P^2}{B^2h} \left( \frac{2a_0 \delta_{ct}^2}{h^3 E_f} + \frac{1}{3C} \right) \tag{1}
\]

While detailed explanations are presented elsewhere [13], a brief explanation of the parameters is given: \( a_0 \) is an equivalent crack length estimated from the current specimen compliance and taking into consideration the damage zone, \( E_f \) is a corrected flexural modulus to account for phenomena affecting the \( P–\delta \) curve, such as stress concentrations at the crack tip and stiffness variability between specimens, and \( C \) is the shear modulus of the adherends [37].

3.2. Direct method to define \( G_{IC} \) and the CZM law

The method followed in this work allows obtaining the cohesive law of the adhesive by the simultaneous measurement of the \( J \)-integral and \( \delta_{ct} \) [16]. The \( J \)-integral applies to the non-linear elastic behaviour of materials, but it remains valid in the presence of a plastic but monotonically-applied loading, as it is the case of the cohesive separation and plastic dissipation in adhesive layers [18]. Based on the fundamental expression for \( J \) defined by Rice [38], it is possible to present a closed-form solution for the tensile energy release rate (\( G_t \)) from the concept of energetic force and also the beam theory for the DCB specimen, as [39]:

\[
G_t = \frac{12I_2(a_0)^2}{E_h h^3} + P_o \theta_a \tag{2}
\]

or

\[
G_t = P_o \theta_p, \tag{3}
\]

where \( P_o \) represents the applied load per unit width at the adherends edges, \( E_h \) the Young’s modulus of the adherends and \( \theta_p \) the relative rotation of the adherends at the loading line (Fig. 2). Expression (2) was used in the present work, considering \( \theta_a \) instead of \( \theta_p \) to gain accuracy in the measurements, since the optical method is inherently more precise. The \( J \)-integral is defined along an arbitrary path encircling the start of the adhesive layer, giving [18]:

\[
G_t = \int_{0}^{\delta_{ct}} t_o(\delta_{ct})d\delta_{ct}. \tag{4}
\]

The value of \( \delta_{ct} \) corresponds to the crack-tip end-opening at failure of the cohesive law, while \( t_o \) is the current normal traction. \( G_t \) is the value of \( G_t \) when the crack initiates propagation, and is given by the steady-state value of \( G_t \) in the \( G_o-\delta_{ct} \) plot [18]. The \( t_o(\delta_{ct}) \) plot is obtained by differentiation of Eqs. (2) and (3) with respect to \( \delta_{ct} \):

\[
t_o(\delta_{ct}) = \frac{dG_t}{d\delta_{ct}}. \tag{5}
\]

3.2.1. Optical method for the parameter measurement

The process detailed here is an improvement of the work developed by the authors in reference [40]. For calculating \( \delta_{ct} \) and \( \theta_a \) for a given image, the optical method requires the identification of eight points (Fig. 3): two points (\( p_1 \) and \( p_2 \)) for measuring the current \( t_o \) value at the crack tip (\( t_o^{ct} \)) during loading in image units (pixels), two points (\( p_2 \) and \( p_6 \)) identifying a line segment in the image for which the length (\( d \)) is known in real world units (mm), two points (\( p_1 \) and \( p_3 \)) on the top specimen and two points (\( p_2 \) and \( p_6 \)) on the bottom specimen for computing \( \theta_o \).

3.2.1.1. Points identification. All eight points are manually identified in the first picture of a trial using an in-house software tool. The
identification of the points is aided by the ruler attached to the specimens, which helps finding their correct locations. In addition, each point between $p_1$ and $p_6$ is printed with a distinct colour (although this is not perceptible in Fig. 3). Using the location of the points in the first picture, the points of the following pictures are automatically identified using a computing algorithm implemented in Matlab. Basically, for each point $p_i$, a rectangular region centred in $p_i$ is extracted from the first image forming a template ($t_i$). This template describes the image pattern that surrounds the point and is used for locating the point in the next image. This is done by finding the position $(u,v)$ in the next image ($I$) that has the highest normalized cross-correlation with the template. The normalized cross-correlation is a measure of similarity between two images that is invariant to linear changes in the pixel intensities and that quantifies the correlation between two images/regions [41]. This measure of similarity was chosen due to its low computational requirements, which is a critical factor given the high resolution of the images, and because changes in rotation and scale of the specimens are expected to be small between two consecutive acquisitions (gapped by 5 s). To take advantage of the colour information, the colour space of the images (and consequently, of the templates) was transformed to the CIELAB colour space. The CIELAB system represents the value of a pixel through three components, $L$, $a$, and $b$ where $L$ is the luminosity and $a$ and $b$ define colour. Since points $p_1$ to $p_6$ are differentiated by their colour, only the $a$ and $b$ components are used when detecting points. The normalized cross-correlation \( \gamma \) of template $t$ with image $I$ at the position $(u,v)$ of image $I$ for the colour component $c$ is defined as:

\[
\gamma(u,v,c) = \frac{\sum_{xy}[(I(x,y,c) - \bar{I}_{u,v,c}) \times (I(x-u,y-v,c) - \bar{I}_c)]}{\left( \sum_{xy}[(I(x,y,c) - \bar{I}_{u,v,c})^2 \times \sum_{xy}[(I(x-u,y-v,c) - \bar{I}_c)^2] \right)^{1/2}}
\]

where $I(x,y,c)$ is the intensity of the colour component $c$ of the pixel $(x,y)$ of image $I$; $I(x,y,c)$ is the intensity of the colour component $c$ of the pixel $(x,y)$ of the template $t$; $\bar{I}_{u,v,c}$ is the average intensity of the colour component $c$ of the region of image $I$ centred at pixel $(u,v)$ and with the same size as $t$, and $\bar{I}_c$ is the average intensity of the colour component $c$ for the template $t$. Finally, the normalized cross-correlation for a single pixel taking into account the colour components $a$ and $b$ is defined as:

\[
\gamma(u,v) = \sqrt{\gamma(u,v,a)^2 + \gamma(u,v,b)^2}
\]

Calculating $\gamma$ for all the pixels of $I$ results in a matrix where the maximum absolute value yields the location of the region of $I$ that has the highest correlation with $t$ and, thus, the most likely location of $p_i$ in the next image. This is done for every one of the eight points identified in the first image. After successfully identifying all the points of the second image, new templates are computed from the second image to search for the eight points in the third image, and so on until processing all images.

3.2.1.2. Computation of $\delta_n$. The value of $t_A^{CT}$ in real world units (mm) is calculated as follows

\[
t_A^{CT} = \frac{d(p_3 - p_4)}{|p_7 - p_8|}
\]

Assuming that the lens distortion is negligible, which is valid for the central area of pictures acquired with modern CCD cameras [42], a length of $d=15$ mm was used for all trials (illustrated in Fig. 3). The pixel size was on average 0.021 mm and, thus, the estimated maximum error of the image acquisition process is $\pm$ 0.011 mm. Finally, $\delta_n$ can be defined as

\[
\delta_n = t_A^{CT} - t_A
\]

where $t_A$ is the theoretical design value of 1 mm. Since $t_A$ can show small variations due to the fabrication process, an adjustment to $\delta_n$ is also applied to make $\delta_n=0$ at the beginning of the test (detailed in Section 4).

3.2.1.3. Computation of $\theta_n$. $\theta_n$ is calculated as the angle between the tangents to the horizontal curves of the 2 scales closest to the adhesive, measured at the crack tip (Fig. 4). The curvature of the top adherend is first computed by fitting a quadratic function to points $p_1$, $p_2$ and $p_6$. The first derivative of the quadratic function at $p_2$ yields the slope of the top curve ($m_{top}$) at the crack tip, which is then used to define a direction vector $\vec{v}_{top}=(1,m_{top})$. The same process is repeated for points $p_2$, $p_4$ and $p_{6}$, yielding the slope of the tangent to the bottom curve at the crack tip ($m_{bottom}$) and its direction vector $\vec{v}_{bottom}=(1,m_{bottom})$. Finally, $\theta_n$ is obtained by measuring the angle between the two vectors:

\[
\theta_n = \arccos \left( \frac{\vec{v}_{top} \times \vec{v}_{bottom}}{|\vec{v}_{top}| \times |\vec{v}_{bottom}|} \right)
\]
4. Results

After having performed the tests on the DCB specimens with varying values of $t_A$, no signs of plasticity were found in the adherends, although for the specimens with $t_A=1$ mm the tensile displacements of the loading points during crack propagation was significant.

4.1. Evaluation of $G_{IC}$ by the CBBM

$G_{IC}$ was initially estimated by the CBBM, which is a well known method for this purpose, and allows accounting for the plasticity of materials. By using this method, it was possible to obtain the $R$-curve for each specimen, relating $G_I$ with $a_{eq}$, as the crack progressed. Fig. 5 gives an example of an experimental $R$-curve obtained by this technique for a specimen with $h=3$ mm, clearly showing the attainment of a steady-state value of $G_I$. For this specimen, $a_0$ was measured at 31.50 mm. The corresponding value of $a_{eq}$ by the CBBM was 38.57 mm, calculated directly from the $P$–$\delta$ data by the first drop of $P$ in the $P$–$\delta$ curve. This difference occurred since $a_{eq}$ accounts for the damage zone [13]. Fig. 6 compares the $G_{IC}$ (N/mm) values of the specimens with varying values of $h$, including the standard deviation for each batch of specimens. An increasing trend for $G_{IC}$ was found as a function of $h$, starting from the specimens with $h=1$ mm, with a steeper increase up to 3 mm and stabilization above this value. Fig. 6 also shows some deviation between specimens of identical conditions, but perfectly within reported deviations under identical testing conditions [40]. Actually, this scatter is related to experimental phenomena such as fabrication issues or small measurement errors or geometry deviations. However, the reliability of the observed tendency with $h$ obviously cannot be questioned, or solely attributed to experimental scatter, on account of the large improvement visible in Fig. 5. In fact, for $h=1$ mm, the value of $G_{IC}=0.688 \pm 0.153$ N/mm was found. The improvement to the specimens with $h=2$ mm (considering average values) was of 13.8%, 48.3% to $h=3$ mm and 56.7% to $h=4$ mm.

4.2. Evaluation of $G_{IC}$ by the $J$-integral

$G_{IC}$ was calculated by Eq. (2), in which $\theta_0$ was estimated as described in Section 3.2.1.3 and plotted with the time elapsed since the test initiation, with one data point every 5 s. Fig. 7 gives an example of the evolution of $\theta_0$ for a selected test specimen (with $h=4$ mm). This specimen is also used in the following figures as being representative of the tests. Shown in the graphic are the raw curve, the 4th degree fitting curve and the corrected polynomial and final curve, adjusted to make $\theta_0$(testing time=0)=0. It should be mentioned that the raw curve of each specimen was adjusted by the most suited polynomial function between 3rd and 6th degrees, by choosing the best correlation factor, $R$ (this also applies to the forthcoming fitting data). This polynomial adjustment is required to smooth the raw data and remove experimental measurement scatter, but also to cancel any eventual misalignment between glued scales in both adherends. The following step consisted on estimating the curve relating $\delta_0$ with the testing time, to determine the cohesive law by Eq. (5).
Polynomial (Experimental law)

Polynomial (Raw curve)

Fig. 8. Plot of $\delta_n$-testing time for a specimen with $h=4$ mm: raw curve, polynomial approximation and adjusted polynomial curve.

Fig. 9. Experimental $G_{IC}$-$\delta_n$ law for one specimen and polynomial fitting.

Fig. 10. Average values and deviation of $G_{IC}$ as a function of $h$ by the J-integral.

in reference [40], where an average value of 1.182 N/mm was found in DCB specimens of natural fibre composites with $h = 5$ mm. The $\epsilon_r-\delta_n$ law for the test specimen followed in this section is presented in Fig. 11, together with a possible trapezoidal approximation, particularly suited to model CZM laws for ductile adhesives [43]. The pertinent parameters for this specimen were $\epsilon_r = 25.1$ MPa and $\delta_{IC} = 0.0692$ mm.

4.3. Discussion

The comparison between both of the proposed methods, the CBBM and $J$-integral, showed that under the present testing conditions both methods agree quite well, although the $J$-integral results are slightly higher. Actually, the difference between average values was of 13.5%, 12.3%, 5.3% and 1.6% for increasing $h$ values between 1 and 4 mm. Despite this small variation, which is probably due to statistical reasons, the observed trend was in high correlation, and in both cases a stabilization of $G_{IC}$ can be expected for joints with higher degrees of restraint (i.e., stiffer or thicker adherends). On the other hand, also, thicker adherends used a larger amount of the input energy in peeling to cleavage, and giving a larger length for the damage zone. These findings are corroborated in the work of Azari et al. [46], regarding the adherent stiffness influence on the fatigue failure of bonded joints, which proved by finite elements that the plastic zone in adhesive joints between steel adherends was consistently higher than identical joints between aluminium adherends during the entire damage uptake process up to crack initiation. Pardoen et al. [23] developed an analytical model to study the constraint effects in adhesive joint fracture, considering steel adherends and two epoxy adhesives, and reported a significant increase of the adhesive plastic dissipation in the fully plastic regime by increasing the adherends thickness. The reported dependence of $G_{IC}$ with $h$ was also addressed in joints with composite adherends, in which $h$ is defined by the number of plies, and results were found to be consistent with those of this work. Mangalgiri et al. [31] justified this tendency with the plastic zone and stress distributions ahead of the debond tip. Actually, the plastic zone was bigger in length across the adhesive layer with increasing number of composite plies (and thus, increasing $h$). Also, thicker adherends used a larger amount of the input energy to the specimen to develop a lengthier plastic zone, thus leaving less available energy for damage growth [47]. On account of this, higher values of $G_{IC}$ can be expected for joints with higher degrees of restraint (i.e., stiffer or thicker adherends). On the other hand, because of the damage zone length limitation by the adhesive ductility, it is also noted that tough adhesives are particularly
prone to these effects, while brittle adhesives are not. The consideration of adherends with different stiffness provides a similar effect to the variation of $h$, as stiffer adherends give a higher restriction to the adhesive layer deformation. Under this scope, Bell and Kinloch [44] tested DCB specimens with aluminium, steel and carbon-fibre adherends, and obtained an increasing trend of $G_{IC}$ with the adherend stiffness. Contradicting results were found by Choupiani [48], but in this case the differences were given by a modification of the failure mechanism for the joints with steel adherends (steel compared to aluminium). With increasing values of $h$, eventually the plastic zone reaches a maximum, i.e., a value above which it does not increase any more, justifying the stabilization of $G_{IC}$. The plastic zone assumption implies that brittle adhesives are marginally affected, if so, by this parameter, and this was confirmed in the work of Fernlund and Spelt [49], whose results on DCB with aluminium adherends up to 12.7 mm thick and a brittle adhesive did not show any variations beyond the statistically related scatter.

5. Conclusions

This work aimed to analyse the influence of $h$ on the measured value of $G_{IC}$ of a ductile adhesive within a pure tensile test as it is the DCB test. Different values of $h$ were considered, between 1 and 4 mm. Two techniques were used for $G_{IC}$, the CBBM and the $J$-integral. For the $J$-integral methodology, a relatively time-consuming approach was required, which involved evaluating by an optical method $\theta$ and $\delta_s$ at the crack tip during the test, followed by polynomial fitting and differentiation. The trend between both methods was consistent, although the $G_{IC}$ values obtained by the $J$-integral were slightly bigger than for the CBBM (between 1.6 and 13.5%, depending on $h$). Disregarding the data reduction method, an increasing trend of $G_{IC}$ with $h$ was found, suggesting that $G_{IC}$ is not a material parameter, but a geometry-dependent quantity instead. The increase of $G_{IC}$ was bigger for the smaller $h$ values, eventually attaining a steady-state value for a given $h$ value, as it sounded by comparing results for specimens with $h=3$ and 4 mm. This result is highly relevant since structural bonding usually falls within small values of $h$, and was considered to be due to an increasing degree of adherend restraining for bigger $h$ values, as a larger region is loaded ahead of the crack tip. The $J$-integral enabled obtaining the tensile CZM law of the adhesive. The CZM curves showed the large plasticity of the polyurethane adhesive. As output of this work, $G_{IC}$ data was given for the strength prediction of bonded joints.

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References

[41] Lewis JP. Fast template matching, vision interface 95, Canadian Image Processing and Pattern Recognition Society, Quebec City, Canada, May 15-19, 1995, p. 120–123.


