Powder Epoxy for One-Shot Cure, Out-of-Autoclave Applications: Lap Shear Strength and Z-Pinning Study

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Abstract: Large composite structures manufactured out-of-autoclave require the assembly and bonding of multiple parts. A one-shot cure manufacturing method is demonstrated using powder epoxy. Lap shear plates were manufactured from powder epoxy and glass fiber-reinforced plastic with four different bonding cases were assessed: secondary bonding using standard adhesive film, secondary bonding using powder epoxy, co-curing, and co-curing plus a novel Z-pinning method. This work investigates the lap shear strength of the four cases in accordance with ISO 4587:2003. Damage mechanisms and fracture behavior were explored using digital image correlation (DIC) and scanning electron microscopy (SEM), respectively. VTFA400 adhesive had a load at break 24.8% lower than secondary bonding using powder epoxy. Co-curing increased the load at break by 7.8% compared to powder epoxy secondary bonding, with the co-cured and pinned joint resulting in a 45.4% increase. In the co-cured and co-cured plus pinned cases, DIC indicated premature failure due to resin spew. SEM indicated shear failure of resin areas and a large amount of fiber pullout in both these cases, with pinning delaying fracture phenomena resulting in increased lap joint strength. This highlights the potential of powder epoxy for the co-curing of large composite structures out-of-autoclave.

Keywords: co-curing; lap shear strength; powder epoxy; Z-pinning; out-of-autoclave manufacturing

1. Introduction

The autoclave process remains a benchmark for creating high-quality composites parts; however, it involves significant acquisition, operating, and tooling costs, especially for large parts [1]. Out-of-autoclave (OOA) manufacturing methods for composites are therefore becoming an increasingly popular alternative to the use of autoclaves, allowing part quality to be maintained whilst reducing cost and manufacturing time. The acceptance of OOA manufacturing is due to their ability to produce autoclave-quality parts without the use of an autoclave [2]. The use of OOA methods provides an opportunity for composites to be used more widely in larger structures such as wind turbine blades.

OOA-manufactured large composite structures currently require the assembly of a number of parts using an additional adhesion step due to oven size restrictions [3]. The joining of these parts generally uses one of three techniques: co-bonding, secondary bonding, and co-curing. A full review of adhesive bonding techniques has previously been completed by Budhe et al. [4]. Co-bonding involves the adhesion of an uncured composite part to a cured composite part, where the uncured adherend plus adhesive are cured together. Secondary bonding involves the adhesion of two cured composites parts. This adhesion step for secondary bonding and co-bonding generates further issues, including
the inability to easily have uniform adhesive contact on complex bonding lines, the inherent additional time, material and labor costs associated, and the additional complication of introducing assembly stresses which affect the overall life of the composite structure [5].

Co-curing is an alternative bonding method which removes the need for an adhesive layer. Co-curing uses resin from the pre-assembled parts and produces no discernable interface due to the simultaneous cure of the assembly (parts + joint) [6]. In comparison, secondary and co-bonded joints have at least one adhesive interface, which allows the introduction of unwanted contaminants such as silicones or moisture [6]. Co-curing has a number of advantages, such as chemical bonding and increased cross-linking, improved mechanical properties, and an inherent lower cost due to reduced labor and material requirement [6,7]. Co-curing is seen as an attractive option due to the reduction in the number of parts, fixings, and curing cycles required [4,8]. This technique is used to make aircraft components; however, information on co-curing techniques is not commonly disclosed by companies [9]. Applications of co-curing in the late 1970s include the McDonnell Douglas AV-8B Strike Fighter Aircraft, where co-curing on the forward fuselage reduced part count by 63%, fixings count by 62%, and the weight by 25%. Similarly, the use of co-curing on the landing gear fairing reduced labor hours by 20%, along with a 65% reduction in part count, a 45% reduction in fastener count, and a 9% weight reduction [10]. Further examples include the effective use of co-curing in the speed brake on the F-15 fighter aircraft, and a co-cured wing created using research from the Technical Research and Development Institute of the Japan Defense Agency in the 1990s for the XF-2 fighter aircraft [11,12].

While co-curing provides the benefits mentioned previously, there are some limitations which are preventing the method from becoming more widely used in composites. Namely, as the size of the co-cured composite structure increases, it is likely that the complexity of the part may increase. This results in increased complexity of the tooling approach which has an inherent additional cost [8,11]. Therefore, the benefits from reduced assembly costs must be balanced against this additional cost.

In this context, one-shot (or single) cure manufacturing processing methods (RTM, Prepreg) have recently been gaining popularity in industry due to their lower cost and higher processing speed [13]. In this exploratory study, a powder epoxy system with separate melting and curing temperatures was used to highlight the co-curing benefits. The powder epoxy has several advantages which are covered in the next section, with this study intending to be an initial investigation into the effectiveness of this resin for bonding purposes, with the idea of future scalability. The lap shear cohesive strength of co-cured samples is investigated in comparison with traditional two-step cures for bonding purposes. A novel pinning technique, developed at Ulster University, has additionally been combined with the co-curing bonding method.

2. Materials and Methods
2.1. Materials
2.1.1. Powder Epoxy

The powder epoxy (PE6405) used in this study was developed by Swiss CMT AG (Siebnen, Switzerland) and produced by Freilacke (Bräunlingen-Döggingen, Germany). The powder system has multiple advantages, such as a low curing exotherm [14,15], which reduces the risk of thermal runaway during manufacturing of thick sections, and a low minimal viscosity [14,15] for optimal wetting and minimal porosity, leading to superior mechanical properties [16–19].

Another main asset of the powder is the ability to separate the powder melting from the cure (Figure 1), enabling composite parts to be melted, pre-shaped, and consolidated without curing. This ability to melt, pre-shape, and consolidate, combined with co-curing is envisaged to help reduce the complexity of tooling required.
Figure 1. Differential scanning calorimetry showing a temperature sweep of (20 °C/min) of PE6405. The temperature gap between the end of the melt and beginning of the cure is represented by purple lines [16].

The resin can be stored at room temperature and is suitable for out-of-autoclave industrial processes [16,17], as well as one-shot-cure processing, which will be demonstrated in this paper.

The powder epoxy has also been shown in previous work to have improved interlaminar fracture toughness [19] compared to untoughened liquid epoxy resins, which is very interesting as there is a linear correlation between fracture toughness and lap shear strength [20].

Finally, the manufacture of large OOA composite structures was performed previously, proving the viability of the system for this specific purpose. Indeed, in the case of the POWDERBLADE project, a 6-metre-long carbon/glass fiber hybrid wind turbine blade was manufactured by the University of Edinburgh (Edinburgh, UK) with this specific powder epoxy system [21].

2.1.2. Glass Fibers

As part of this study, a 591-g/m² UD stitched glass fabric from SAERTEX was used for sample manufacturing. The fabric contains 520 gsm of 0° E-glass fibers as well as 17 gsm of polyester stitching and 54 gsm of 90° E-glass fibers [22]. UD stitched glass fabric was chosen for this initial study with a view of potential applications in standard OOA processes such as wind turbine blade manufacturing [23].

2.1.3. Adhesive Prepreg

The adhesive prepreg used in this study is VTFA400 from SHD Composites (Sleaford, UK). This is a toughened epoxy adhesive film within a very thin glass fiber mesh to ensure uniform thickness and minimal porosity. Adhesive prepreg is commonly used as a large area structural adhesive, for example in the aerospace industry [24].
2.2. Lap Shear Samples Manufacturing and Testing
2.2.1. Manufacturing and Testing Overview

A summary of the manufacturing process is provided in Figure 2.

Firstly, glass fiber-reinforced plastic and powder epoxy plates were produced using a previously developed manufacturing process [19].

Due to the low minimum viscosity of powder epoxy prior to gelation, the resin tends to bleed out of the composite during the sintering/curing stage. This can cause some variability in the fiber volume fraction (FVF) of the manufactured plates [19]. To solve
this, a previously manufactured steel frame was used (8 mm thick, inner dimensions of 300 mm × 280 mm) along with two steel caul plates. The frame rests on the larger lower caul plate, while the upper caul plate inserts tightly into the frame to ensure uniform pressure is exerted by the applied vacuum, and to limit resin bleed-out. For ease of removal of the cured/sintered plates, the steel caul plates were coated in TFG 250 PTFE coated glass fiber, with the frame being coated in thin TF 050AH PTFE tape.

The GFRP plies were cut to fit a steel mold (lower caul plate and frame) covered in PTFE. The powder was weighed and homogeneously sprinkled on interlayers to reach 50% FVF, ensuring good fiber wetting and an even resin distribution (step 1). The PTFE-covered steel upper caul plate was then placed on top (step 2) and the whole system was vacuum bagged and cured or only melted/sintered, depending on the sample set investigated (step 3). The sintering and curing cycles for the powder epoxy GFRP plates are provided in Figure 3.

![Figure 3. Curing and sintering cycles for powder epoxy and VTFA400 adhesive.](image)

The plates were then cut in half at 90° to the longitudinal direction and the edges were trimmed (step 4). At this stage, Z-pinning of two sintered half plates was completed, outlined in detail in Section 2.2.3. Plates were then placed in a PTFE covered steel mold fitting the ISO 4587 dimensions (step 5a + 5b). Figure 4 shows the sintered half plate inside the PTFE covered steel mold.
The half plates were placed under vacuum again and bonding lines were then cured accordingly. The purpose of the side steel plates (also shown in step 5a, Figure 2) was to maintain uniform pressure on the plates and bonding line while under vacuum.

The co-cured plate is shown in Figure 5 below.

Figure 4. Sintered GFRP + powder epoxy half-plate inside steel frame.

Figure 5. Photo of co-cured plate (case 3).
Finally, the lap shear plates were cut to dimension according to ISO 4587 (step 6). During the cutting process, end tabs were applied temporarily over the bonding region with double-sided tape to reinforce and prevent premature damage to the plate and samples during cutting (Figure 6), then removed prior to testing. The double end tab configuration was used to obtain a perfect alignment during gripping, therefore preventing undesired bending of the sample during testing.

![End tab configuration during cutting of the lap shear plates.](image)

The co-curing cohesive strength was investigated according to ISO 4587: Adhesives—Determination of tensile lap-shear strength of rigid-to-rigid bonded assemblies. The lap shear strength test setup is provided in Figure 7. An INSTRON 3369 10 kN universal testing system was used, and samples were tested at 2 mm/min. The temperature was maintained at a stable 22 °C in the lab throughout the tests.

### 2.2.2. Sample Set Description

Four different sets of samples were manufactured to compare their cohesive properties (Figure 8).

- **Case 1:** The powder epoxy-based GFRP half-plates were initially cured and an adhesive prepreg (VTFA400, SHD composites), representative of industrial standard practice, was placed on the bonding line and cured.

- **Case 2:** The powder epoxy-based GFRP half-plates were cured, and then powder epoxy was sprinkled on the bonding line and used as an adhesive.

- **Case 3:** The powder epoxy-based sintered GFRP plate was melted and then cooled without curing, producing a vitreous-like, sintered plate which was then cut. The structural integrity of the powder epoxy sintered plate is another advantage when compared to standard uncured prepreg, which would present issues if cutting was attempted. The half-plates were then co-cured in a one-shot-cure process (Figure 5).

- **Case 4:** The half plates were melted then pinned and finally co-cured to assess the adhesion reinforcement potential of pinned one-shot-cure samples as compared to case 3. The sintered plates have the ability and ease to be pinned without much damage, and then the plates are melted again during their co-curing phase, allowing for a very good consolidation at the pinned locations. This consolidation is arguably better than the one a thermoplastic would be able to achieve due to a very low viscosity. The pinning process is described in detail in Section 2.2.3.

The lap shear sample cross sections were checked before testing to check for macroscopic inhomogeneities, of which none were found. Resin overflows were observed in cases 3 and 4, a standard behavior associated with co-curing.
Figure 7. Lap shear strength test setup on an INSTRON 10 kN universal testing system.

Figure 8. Schematics of the four sets of lap shear samples manufactured.
2.2.3. Pinning

A novel technique for providing through-thickness reinforcement was developed at Ulster University through the EPSRC Future Composites Manufacturing Research Hub. The method involved using a nylon/carbon filament which was placed through the thickness of the preform using a veterinary needle and trimmed to length. Once the desired array of pins was placed, they were treated via compression with hot plates, causing the extended ends of the pins to lie flat against the surface of the fabric, which, once cool, binds it together [25].

In this study, the filament used was a commercial PA6-based 3D printing filament by Markforged®. There are several options available, differing primarily in the material used for the fiber reinforcement. In this work, the filament used was KB-AB-50 which comprises a Kevlar based reinforcement.

The pinning process can be broadly described as follows, with the steps illustrated in Figure 9:

1. The fiber lay-up to be pinned was placed into a clamping frame which provides a fixed back-stop distance which can be adjusted by the addition of plates under the parts.
2. The filament was threaded through an 18 Birmingham Gauge veterinary needle that was mounted in a 3D printed handle for operator convenience.
3. The needle was pressed through the fiber in the desired locations.
4. The filament was then gently threaded through the needle until it hits the backstop, this provides a consistent excess length on the underside of the part.
5. The needle was then withdrawn whilst holding the filament in place. The closing of the fibers around the needle provides a light frictional force holding the filament in place.
6. The filament was then cut above the part to match the backstop distance.

![Figure 9. Representative image of the main steps of the pinning process.](image-url)
Once the desired array of pins was placed, the fiber was then subjected to a hot-press process. The tool is first heated to 150 °C in an oven. The fiber was then placed into the hot tool and the tool clamped up whilst at temperature. This has the dual effect on both bending and flattening the ends of the pins against the part, but also compressing the part to the correct thickness for infusion.

In this study however, the technique was modified to satisfy the need to maintain alignment of the overlapping sections for the lap-shear samples. This was achieved by leaving the fiber in the clamping frame and using a 2-stage process. This modified process can be described as follows:

1. In Press 1, a support plate is placed underneath the part (a). This prevents push-through of the pins.
2. A press-plate is heated in an oven to 150 °C, placed on the upper surface of the part, and left to cool to ambient temperature overnight (b & c). The mass of this plate provides the necessary pressure to deform and flatten the pin ends. The flattening of the pins is illustrated as steps a, b & c in Figure 10.
3. Once the first side has cooled, the fixture is inverted with the support plate now placed against the flattened pins.
4. The process described for Press 1 is repeated for the underside of the part (Press 2: a–c).

![Figure 10. Graphical representation of the 2-step press process.](image-url)

After this process was completed, the plate was removed from the clamping frame and subjected to the same curing cycle, as described in the manufacturing overview.

2.2.4. Pinning Review

The exploratory pinning process for these parts presented some challenges compared to the previous work on dry fibers that was conducted at Ulster University.

The first challenge was the toughness of the sintered epoxy preventing the penetration of the fibers by the needle. This was solved by gently introducing cracks within the epoxy in the region to be pinned. Some of the epoxy was lost with this process; however, this was primarily in the form of dust on the outer surface with the fibers preventing egress of epoxy within the part.
The second challenge was the overlap of the pinning area. Previous work had centered primarily around pinning the full area of the plaque without the concerns of overlap presented by the geometry of the ISO 4587 specimens.

To overcome this, the fibers were placed into the clamping frame with spacer plates. These plates prevented pinning in the area outside of the overlap region whilst allowing proper pass-through in that region and supported the fibers, as shown in Figure 11.

![Figure 11. Case-4 samples prior to undergoing hot-press.](image)

Once the array of pins was placed, the next challenge was conducting the hot press process. In prior works, the larger pinned region allowed sufficient integrity for the pinned part to be removed from the clamping frame with only the light friction from the fibers holding the pins in position. However, with the small overlap area, there was concern that this would be insufficient for these samples. It proved not to be.

The solution was to perform the hot-press process without removing the material from the frame. To do this, the part was pressed twice using a plate heated to 150 °C with pressure applied by means of a 5 kg weight placed on top. The first press retained the backstop used in the pinning process to prevent the pins from merely being pushed through the part. The second press followed the same setup; however, the underside of the part was supported again to prevent the pins from being pushed through the part. In both cases, a layer of Richmond Aerovac ETFE release film was used to prevent the powder epoxy adhering to the plate.

### 3. Results/Discussion—Mechanical Properties

#### 3.1. Lap Shear Tensile Testing

The lap shear results are presented in Figure 12. A large lap shear strength reduction for VTFA400 commercial adhesive bonding was measured (−24.8%) compared to the cured samples that are bonded with the powder epoxy. This behavior highlights the superior adhesive properties of the powder epoxy system for out-of-autoclave (OOA) manufacturing purposes. Indeed, the powder epoxy system was previously shown to have high fracture toughness [19] compared to standard epoxy prepregs. Additionally, the very low melt viscosity allowed for a homogeneous adhesion consolidation.
Co-curing the plates has introduced both the abilities of covalent bonding and a better consolidation at the inter-plate region (adhesion surface), leading to a +7.8% lap shear strength increase for co-cured compared to cured and bonded samples. This shear strength increase is attributed to both a better consolidation and crosslinking.

The through-thickness reinforcement pinning of uncured powder epoxy laminates, followed by co-curing, showed further substantial improvements, with a +45.4% lap shear strength increase for pinned and co-cured compared to cured and bonded samples. This result highlights the ability to pin through the sintered/vitreous plates, and to then remelt and co-cure the powder epoxy around the pins, resulting in a superior consolidation. The mechanical viability of the pinning method was therefore demonstrated in this study.

3.2. Digital Image Correlation (DIC)

A comparison of typical damage mechanisms for each joint configuration can be inferred from digital image correlation (DIC). Specimen cross-sections were speckled with white spray paint spanning the full gauge length, to provide evenly distributed and high contrast tracking objects. A 12-megapixel Sony IMX477 sensor and Sill Optics Correctal T/0.2D telecentric lens assembly was used to capture high-resolution orthographic images of the specimens during testing. For each sample type, three tests were recorded (H.264 video) at a 1080 × 1920 resolution and with a frame rate of 25 frames per second (fps). Video files were subsequently exported into image (.png) stacks with a 1:5 conversion ratio, resulting in an effective frame rate of 5 fps. Displacement and strain fields were calculated by importing image stacks into DICe open-source software by Sandia National Laboratories (SNL) [26]. For each analysis, the selected shape functions were translation, rotation, normal stretch, and shear stretch. An SSSIG threshold of 150, a 31-pixel subset size, and a 15-pixel step size were used for all analyses. The numerical error for this (2D-DICe) analysis is approximately ±5% of the recorded value for elastic deformations [27].

Visualizations of the DIC results were produced using Paraview-5.9.1 open-source software. For each specimen, the DIC generated and imported an exodus (.e) file, and a Delaunay 2D-triangulation filter was applied. An alpha value of 22 was selected to populate the produced mesh. A Cauchy xy-shear strain and xx-longitudinal strain distributions of a representative sample for each manufacturing case are plotted in Figures 13 and 14, respectively.
Figure 13. Pre-failure Cauchy strain$_{xy}$ fields of representative (a) VTFA400 specimens (b) cured and bonded (c) co-cured; and (d) Z-pinned.

Figure 14. Cont.
For each of the four representative specimens, observations were plotted at the frame prior to brittle failure, where the specimen strains were at peak magnitude. To facilitate straightforward comparisons between fields in Figures 13 and 14, distribution limits were defined as $-0.1$ to $0.0$ and $0.0$ to $0.1$ mm/mm, respectively. For comparison, post-failure photographs of the same specimens are provided in Figure 15. In each of these images, the background was enhanced to improve contrast, and annotations were added where appropriate.
Figure 15. Post-failure images of representative (a) VTFA400 specimens (b) cured and bonded (c) co-cured; and (d) Z-pinned.

In observing the strain fields provided in Figures 13 and 14, local peaks can be associated with damage locations via the fractured specimen photographs in Figure 15. For all specimens, a shear strain peak occurs along the central bond line region with values generally largest at the upper and lower extremities. For both the cured and VTFA400 specimen types, these large shear and longitudinal strains correspond directly to points of damage initiation. As seen in Figure 15a,b, fracture in these bonded specimens has occurred entirely along the bond line via cohesive failure. The larger shear strain in the cured and bonded specimen is also anticipated, since the VTFA400 failure load is 24.8% lower. The co-cured specimen shows an oval shaped shear and longitudinal strain peak at the upper left corner of the lower adherend. This strain peak corresponds to a macro-void within the resin spew region and is present in certain co-cured samples. In considering Figure 15c, it can be discerned that a crack is initiated at the stress concentration around this void, extending horizontally until reaching the bond line and subsequently propagating downwards. Given the relatively low strains present across the remaining specimen cross section, it is possible that this mechanism may have caused minor premature failure in some co-cured samples. Similarly, the longitudinal strain distribution in Figure 14d indicates a high strain location at the upper left of the lower adherend in the Z-pinned representative specimen. Comparing the specimen strain distribution with the post-failure image in Figure 15d, it is apparent that damage has initiated at the location of a Z-pin and propagated along a similar path to the co-cured specimen. Although localized stress concentrations from Z-pinning are inevitable, larger load bearing capacities can still be achieved after further manufacturing process refinement [28].

3.3. Fracture Behaviour

Following testing, the lap shear samples were observed under a TM4000Plus scanning electron microscope (SEM) manufactured by Hitachi (Tokyo, Japan). Figure 16 shows the fracture surfaces under a magnification of 80 for all four sets of samples. The reinforcing mesh of the VTFA400 adhesive is clearly visible in Figure 16a as well as the glass fibre
polyester stitching. In some areas, the raw fibers from the adherend are visible, suggesting an adhesive failure. However, in the bottom left of the figure, resin damage is clearly visible as well as the holes contained within the VTFA400 reinforcing mesh. This suggests a mixture of cohesive and adhesive failure for these specimens. Similar features are visible in Figure 16b for the cured and powder epoxy bonded samples with a mixture of adhesive and cohesive failure clearly visible in the fracture surface. However, there is a higher degree of resin damage on the surface of these samples which suggests a slightly better compatibility between the adhesive and adherend than for the VTFA400 samples, and could explain the higher measured lap shear strength.

Figure 16. SEM Pictures of Fracture Surfaces with ×80 Zoom for (a) VTFA400 Specimens (b) Cured and Bonded with Powder Epoxy (c) Co-cured and (d) Co-cured and Z-pinned.

The fracture surface of the co-cured and Z-Pinned samples are different from that of the first two specimens. Indeed, hardly any resin is visible on the fracture surfaces. The 0°, 90°, and polyester stitching from the glass fiber fabrics are visible from both Figure 16c,d. Resin was present in only small areas of the fracture surface. Observation of the resin areas
under higher magnification revealed hackle patterns characteristic of shear failure [29].

The fracture surfaces of the co-cured and Z-pinned samples looked very similar apart from
the presence of the Z-pins, as shown in Figure 16d. Co-curing of the resin has led to a very
thin bond line and fiber migration within the bond. This explains why hardly any resin
is present on the fracture surface while a large amount of fiber pull-out is visible. This
explains the higher strength measured for these samples compared to the previous two
sets. The presence of the Z-pins provides further reinforcement of the lap joint, but this is
not characterized by significant differences in the fracture surface. Instead, the Z-pins delay
the fracture phenomena occurring in the co-cured samples by resisting crack propagation,
leading to a higher lap joint strength.

4. Conclusions

Glass fiber-reinforced powder epoxy plates were manufactured using a process de-
veloped in a previous study [19] and subsequently used to produce lap shear plates. The
tensile lap-shear strength of the glass fiber reinforced powder epoxy lap shear plates was
investigated, with results showing an increased load at break for co-curing methods com-
pared to secondary bonding. Secondary bonding using VTFA400 had a load at break 24.8%
lower than secondary bonding using powder epoxy. Co-curing increased the load at break
by 7.8% compared to powder epoxy secondary bonding, with the co-cured and pinned
joint resulting in a 45.4% increase in the load at break.

The typical damage mechanisms were investigated using digital image correlation.
For secondary bonding using VTFA400 and powder epoxy, fracture occurred entirely along
the bonding line via cohesive failure. Bonding using co-curing displayed premature failure
in some samples related to macro voids in the resin spew area of the joint. The Z-pinned
and co-cured joint showed damage around the Z-pin locations, which was to be expected;
however, it is anticipated that, with further refinement of the manufacturing process, larger
load bearing capacities could be achieved.

SEM indicated a mixture of cohesive and adhesive failure for the VTFA400 and powder
epoxy bonded lap shear samples, with higher resin damage on the powder epoxy bond
suggesting better compatibility of the adhesive and adherend. The co-cured and Z-pinned
samples showed very similar fracture surfaces with very little resin, and indicated shear
failure in resin areas, and that a large amount of fiber pullout had occurred. The Z-pinning
provides further reinforcement to the lap joint, delaying the fracture phenomena and
therefore resulting in the higher lap joint strength. Although a combination of DIC and
SEM imaging have effectively indicated dominant fracture mechanisms from mechanical
testing, a multiscale evaluation, such as that undertaken by Stukhlyak et al. [30], could
provide a more in-depth understanding of damage mechanism in future investigations.

This initial study was a promising proof of concept to demonstrate the advantages of
the powder epoxy resin when combined with co-curing and pinned joining methods. The
very specific processing properties of the powder epoxy system allowed to show the best
of both thermoset (low viscosity, better co-curing mechanical properties) and thermoplastic
(non-brittle pinning on complex structures, higher toughness), allowing co-curing and
pinning all at once, leading to vastly superior mechanical properties when combined.

Overall, the powder epoxy resin has a range of advantages, making it a suitable can-
didate for the co-curing of large composite structures. Future work will involve employing
the co-curing and pinning bonding methods on more complex structures, for example on
thick section composites such as tidal turbine blades. Automated applications, such as
additive manufacturing, will also be considered. There will be the ability to locally change
the polymeric properties to meet specific requirements, e.g., erosion protection [31].

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