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Novel epoxy powder for manufacturing thick-section composite parts under vacuum-bag-only conditions. Part I: Through-thickness process modelling

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12 Abstract

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13 Thick-section composite parts are difficult to manufacture using thermosetting resins due to their exothermic curing reaction. If processing is not carefully controlled, the build-up of heat 14 15 can lead to warpage or material degradation. This risk can be reduced or removed with the use 16 of a low-exotherm resin system. Material and process models are presented which describe 17 vacuum-bag-only processing of thick-section composites using a novel, low-exotherm epoxy 18 powder. One-dimensional resin flow and heat transfer models are presented which govern the 19 fabric impregnation and temperature evolution, respectively. A semi-empirical equation is 20 presented which describes the sintering of the epoxy powder. The models are coupled via 21 laminate thickness change, which is determined for a simplified ply microstructure. The 22 resulting system of equations are discretised and solved numerically using a finite difference 23 code. A case study is performed on a 100-ply laminate, and the advantages and disadvantages 24 of using epoxy powders are discussed.

25 Keywords: A. Epoxy powder, C. Process modelling, E. Consolidation, E. Cure

26

27 1. Introduction

28 The concept of using polymer powders for composite manufacturing has been explored since 29 the 1970s [1]. In the late 1980s and early 1990s, NASA and supporting academics conducted 30 several investigations into 'towpregging' and tape production using both thermoplastic and 31 thermosetting powders [2–5]. More recently, significant research has been focused on the use 32 of epoxy powders for the production of low-cost, vacuum-bag-only (VBO) prepregs, which

1 can be used to manufacture wind and tidal turbine blades (e.g. European projects 2 MARINCOMP and POWDERBLADE). These projects investigated the processing of epoxy 3 powder composites [6-8] and their mechanical performance [9-11]. The results of the projects 4 showed that epoxy powder composites were mechanically comparable to conventional epoxy 5 composites used by the wind energy industry. They also revealed that the epoxy powder has 6 several processing advantages which make it well suited to the VBO prepreg format. Firstly, 7 the epoxy powder could maintain a viscosity of 10 - 100 Pa s for up to 3 hr at 120°C, which is 8 low enough to infuse glass-fibre (GF) and carbon-fibre (CF) fabrics when using a VBO prepreg 9 format [8]. Secondly, the epoxy powder has excellent storage stability which means that the 10 epoxy powder doesn't suffer from 'out-time' curing effects to the same extent as conventional epoxy-based VBO prepregs (i.e. they could be left at ambient conditions for several months 11 12 without any significant change in the degree of cure (DoC)). Its chemical stability at 13 temperatures up to 120°C also provides the potential to infuse composite parts separately (e.g. 14 wind turbine blade skins, spars, shear webs, etc.), and then co-cure them to form a "one-shot" 15 structure without the need for adhesive bonding [6]. In terms of its curing behaviour above 16 120°C, the epoxy powder produces relatively little exothermic heat (< 200 J/g) compared to 17 conventional epoxies (e.g. 400 - 600 J/g), which is significant for the manufacture of thick-18 section components due to the risk of "thermal runaway" [8].

19 While the aforementioned results showed the potential of the epoxy powder, the material 20 characterisation experiments focussed on processing small samples using programmed 21 temperature cycles. In practical terms, however, manufacturing thick-section composite parts 22 is significantly more complex. It has been shown that the initial state of the powder-based 23 composite can affect how the material is processed; Ramasamy et al. [12] noted that braiding 24 with powder-coated towpregs was challenging due to the increased bulk of the tows and 25 preform. Moreover, for thick-sections, temperature divergence from the programmed 26 temperature cycle is unavoidable due to the lag in heat transfer, particularly in the centre of the 27 laminate [13]. It is impossible to know whether the overall manufacturing process will benefit 28 from the seemingly advantageous characteristics of the epoxy powder without taking a more 29 holistic view of the system. This system includes key processes such as resin flow and heat 30 transfer, which can be modelled mathematically.

The modelling and simulation of flow processes in composite materials has undergone a great deal of development in the last three decades, owing largely to the advent of liquid composite moulding (LCM) processes [14]. In most cases, thermoset resin flow through a fibre-bed has

1 been modelled using Darcy's Law as the governing equation [15]. Numerous resin flow models 2 have been developed to describe manufacturing methods which utilise through-thickness flow 3 (e.g. Resin Film Infusion (RFI), Compression Resin Transfer Moulding (CRTM), VBO 4 prepregs, etc.) [16–18]. These models vary in their complexity depending on the processing 5 phenomena which they describe. For VBO prepregs (a.k.a. out-of-autoclave (OoA) prepregs), 6 plies are partially impregnated by resin, which means the resin is only required to flow into the 7 adjacent plies. This a less complex flow problem than RFI or CRTM where resin must flow 8 through an entire preform. As such, some resin flow models for VBO prepregs have been 9 comparatively simple. Centea and Hubert [18] used Darcy's Law to model the process as one-10 dimensional (1D) radial resin flow into the fibre tows, using an equivalent hydraulic radius R_t for the elliptical geometry of the tows. Alternatively, the resin flow in VBO prepregs can be 11 12 considered dual-scale in nature [19]; macroscopic flow around the fibre tows (inter-tow flow), 13 and microscopic flow within the fibre tows (intra-tow flow), with up to four orders of 14 magnitude difference between the permeabilities of these regimes [20,21]. Cender et al. [19] 15 modelled the dual-scale flow phenomena as 1D resin flow through two porous media in series; 16 assuming that the resin would fill the inter-tow region entirely before beginning to impregnate 17 the intra-tow region due to the difference in permeabilities.

18 It should be noted that these VBO prepreg resin flow models were used to simulate the 19 processing of thin laminates only (< 4 mm thick), with simplified assumptions made for the 20 temperature conditions. For example, Cender et al. [19] assumed isothermal conditions for their 21 experiments and were able to solve the resin flow problem analytically, while Centea and 22 Hubert [18] input the time-temperature data directly into the model and assumed uniform 23 temperature throughout the laminate. For thick-section composites, a uniform temperature 24 distribution cannot be assumed because large temperature gradients are known to develop 25 within the laminates due to the inherent insulating properties of the fibres and polymer matrix. 26 This is compounded by using exothermic thermosetting resin systems which can cause 27 "thermal runaway" of the composite part during manufacture, as previously mentioned. As a 28 result, a considerable amount of research has been carried out on heat transfer modelling of 29 thick-section composite parts manufactured using thermosetting prepregs. Several heat transfer 30 models for thick-section composites were developed in the 1980s and early 1990s [13,22-25]. 31 For these models, heat transfer within the laminate was described using the well-known heat 32 equation with an additional term for heat generation from the exothermic curing reaction.

1 In general, the aforementioned publications vary in terms of whether or not they account for 2 consolidation of the thick-section laminate. Bogetti and Gillespie [13] opted not to model 3 consolidation, due to the added complexity of modelling in 2D, however, they acknowledged 4 that thickness effects would strongly influence the processing of thick sections. With 5 simulations, they demonstrated that increasing laminate thickness created a thermal lag 6 between the prescribed temperature cycle and the centre of the laminate. Building on this, 7 Twardowski et al. [25] implemented approximations for consolidation (based on either uniform 8 linear or exponential change in FVF over a set period of time) and found that it had a significant 9 effect on heat transfer in the laminate during processing. This trend was corroborated by Shi 10 [26] via analytical models, numerical simulations, and experiments. Oh and Lee [27] modelled 11 both heat transfer and consolidation, however, the simulations were performed separately so 12 that the models were not fully coupled i.e. data from the heat transfer model was input into the 13 consolidation model, but not vice versa. Shin and Hahn [28], on the other hand, developed a 14 finite difference code which fully coupled resin flow and heat transfer of a thick prepreg 15 laminate processed in a heated press.

16 Initial work on the process modelling of thick-section epoxy powder-based composites began 17 with a focus using on coupling resin flow to heat transfer via cure kinetics and chemorheology, 18 and coupling heat transfer to resin flow via thickness change [29,30], however, these models 19 did not account for the thickness change due to powder sintering. The bulk volume of epoxy 20 powder is up to twice the volume of fully sintered powder depending on the packing of 21 particles. Greco and Maffezzoli [31] presented models to describe the melting and sintering of 22 polymer powder. While polymer melting and sintering are linked, densification of the powder 23 due to sintering does not occur instantaneously upon the polymer reaching its melting point. 24 Instead, it is dependent on the viscosity and surface energy of the polymer [32]. Kandis and 25 Bergman [33] developed an Arrhenius-type semi-empirical relation to describe the kinetics of 26 the sintering process. Greco and Maffezzoli [31] fit this sintering model to thickness 27 measurements from thermal mechanical analysis (TMA) and showed that it better described 28 the densification of the powder than their original melting model.

Part I of this study focuses on the development of numerical models for simulating the manufacture of thick-section structures using epoxy powder technology. Process models are presented for coupled resin flow and heat transfer, and a semi-empirical model is introduced which describes the sintering of the powder. These models are combined with existing material models for the epoxy powder [8] to describe how temperature, viscosity, DoC, degree of impregnation (DoI), and laminate thickness evolve during the manufacturing process.
 Experimental validation of the model and additional simulations are the subject of Part II of
 this study [34].

4

5 2. Methodology

6 2.1 Semi-empirical sintering model for epoxy powder

An epoxy powder system (GRN 918), supplied by ÉireComposites Teo., is considered in this
study [35]. This system was previously characterised using thermogravimetric analysis (TGA),
differential scanning calorimetry (DSC) and parallel-plate rheometry (PPR), and semiempirical models were developed to describe its cure kinetics and chemorheological behaviour
with respect to temperature [8].

12 For this study, a HAAKE MARS II rheometer (Thermo Scientific) was used to perform an 13 additional set of PPR tests which measured the compaction of the powder due to sintering. 14 Khoun et al. [36,37] showed that it was possible to measure the thickness change of an epoxy 15 sample by monitoring the gap height between the parallel plates for a given normal force. For 16 the MARS II rheometer, a Peltier module was used to heat the lower plate, and a PEEK sample 17 hood was used for insulation. For all tests, the top plate was lowered onto the powder sample 18 until it resisted compaction (the rheometer had a limiting normal force of 50 N). A normal force 19 of 50 N was approximately equivalent to 52 kPa of pressure for the 35 mm plates that were used. Isothermal tests at 50°C, 55°C, and 60°C were carried out because these temperatures 20 21 were near the glass transition temperature (T_a) of the epoxy powder; the temperature at which 22 the powder would begin to sinter. The rheometer began recording the sample thickness as soon 23 as the test temperature was reached. No resin squeeze-out was observed during testing due to 24 the resin's high viscosity at the test temperatures (> 20,000 Pa s [8])

The results of the tests were fitted to a semi-empirical model developed specifically for GRN 918. Kandis and Bergman [33] previously modelled the polymer powder sintering process using an Arrhenius-type expression due to the exponential temperature dependence of the polymer viscosity [32] (i.e. as the viscosity reduces, the powder collapses due to surface energy or some external driving force). From chemorheological modelling of GRN 918, it was known that its viscosity reduction was best modelled using the Williams-Landel-Ferry (WLF) equation [8]. As such, the results of the PPR testing were fitted to the following WLF-type
 model,

$$\frac{d\chi}{dt} = -\chi_E \exp\left(\frac{C_{\chi 1}[T - T_{\theta}]}{C_{\chi 2} + T - T_{\theta}}\right) (\chi - \chi_{\infty})^B \tag{1}$$

3 Where χ_E is a pre-exponential rate constant, χ_{∞} is the powder void fraction at $t = \infty$, *T* is 4 temperature [K] at time *t* [s], and T_{θ} is the onset temperature for melting [K]. $C_{\chi 1}$, $C_{\chi 2}$ [K], 5 and *B* are fitting constants. The powder void fraction χ at time *t* is described by,

$$\chi = 1 - \frac{\rho_r}{\rho_P} \tag{2}$$

6 Where ρ_P is the bulk density of the polymer, and the density ρ_r is determined using,

$$\rho_r = \rho_0 \frac{L_0}{L} \tag{3}$$

7 Where ρ_0 is the initial density of the powder, L_0 is the initial thickness of the powder samples, 8 and L is the thickness at time t, which was measured using PPR.

9 The sintering model parameters for GRN 918 are given in Table 1. From the PPR tests, the 10 initial powder void fraction was estimated as being approximately 0.503. This was confirmed 11 by manually compacting the powder in a 5 ml graduated cylinder and weighing it on a 12 microbalance.

14	- I	fable	1.	Parameters	for	the	sintering	model.
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Parameter [units]	Value
Хе	3×10^{-5}
$C_{\chi 1}$	11.5
$C_{\chi 2}$ [K]	24.5
$T_{ heta}$ [K]	313.48
χ_{∞}	0.0
В	0.5
	$\frac{d\chi}{dt} = -\chi_E \exp\left(\frac{C_{\chi 1}[T - T_{\theta}]}{C_{\chi 2} + T - T_{\theta}}\right) (\chi - \chi_{\infty})^B$

1 **2.2 Resin flow**

Vacuum-bag-only (VBO) prepregs are plies of fabric reinforcement partially impregnated with resin on one or both sides. When manufacturing thick laminates, a prescribed number of plies are stacked on top of each other, as shown in Figure 1. At the beginning of the manufacturing process, the lay-up of plies is effectively a system of alternating resin and fabric layers. As heat and pressure are applied, the resin viscosity drops, and the resin is forced to flow into the fabric layer, leading to a change in laminate thickness.

8 Figure 2 illustrates the resin flow pattern which is expected for an epoxy powder based VBO 9 prepreg, i.e. resin flow around the fibre tows (inter-tow flow), followed by resin flow into the 10 fibre tows (intra-tow flow). Note that, while Figure 2 serves to illustrate the dual-scale flow 11 phenomena in VBO prepregs, VBO prepregs are usually partially impregnated with resin; 12 meaning some of the inter-tow and intra-tow regions may already be filled. While the model developed by Centea and Hubert [18] is perhaps more accurate in modelling the porous media 13 14 as an ellipse, it assumes that the initial degree of impregnation (DoI) has been advanced to the 15 point that the inter-tow region is completely filled. This is not always the case as a lower DoI 16 can increase the air permeability of the VBO prepreg [38], leading to improved gas evacuation 17 and low porosity [39]. For epoxy powder VBO prepregs, the powder is dispersed on only one 18 side of the VBO prepreg, meaning that the "dry side" of the VBO prepreg is not completely 19 filled with powder, hence there is inter-tow space which the epoxy must fill. Given the high 20 initial viscosity of the epoxy powder (measured as approx. 20,000 Pa s at 70°C [8]), relatively 21 slow viscous resin flow can be expected in the inter-tow region when processing thick-section structures, despite the its high permeability (circa 10⁻⁹ m²). Moreover, ÉireComposites Teo.'s 22 23 recommended drying temperature for the epoxy powder ranges from 40°C to 55°C; at 40°C, the epoxy is solid with a viscosity approaching infinity, while, at 55°C, the powder sinters and 24 has a viscosity of 5×10^6 Pa s (predicted using the chemorheological model from [8] – see 25 26 Equation A.1). As such, both inter-tow and intra-tow resin flow are considered herein.



14 compared to process models which consider the fibre bed compaction of VBO prepregs [40–

15 42] or process models which consider resin flow through multiple layers of fabric [16,17].

Under these assumed conditions, there is a fixed volume of inter-tow space and intra-tow space that the resin could flow into and fill. The dual scale resin flow models developed by Cender et al. [19] are chosen as a suitable basis for modelling the epoxy powder VBO prepreg system. Some modifications are required, however, to adapt the equations for the highly transient temperature conditions of thick-section composite processing.

6



7

Figure 2. Schematic of the dual-scale flow pattern in VBO prepregs. (Left) If the DoI is zero, as shown no
 resin has flowed into the fabric layer (which is bounded by the red dashed line). (Middle) Resin flows first
 into the inter-tow space due to its higher permeability. (Right) Resin then flows into the low permeability
 tow.

12 To solve for transient temperature conditions (i.e. accounting for viscosity change due to 13 temperature and DoC), Darcy's Law is written as,

$$\frac{dl}{dt} = -\frac{K}{\varphi\eta} \cdot \frac{dP}{dx} \tag{4}$$

- 14 Where *l* is the flow front position [m] at time *t* [s], *K* is the permeability $[m^2]$, φ is the porosity,
- 15 η is the resin viscosity [Pa s], which is temperature and cure dependent, and P is pressure [Pa].
- 16 Equation 4 is integrated over the pressure boundary conditions for inter-tow flow, $P|_{x=0} =$ 17 P_{app} and $P|_{x=l} = 0$,

$$\int_{0}^{l} \frac{dl}{dt} dx = \int_{P_{app}}^{0} -\frac{K}{\varphi \eta} \cdot \frac{dP}{dx} dx \quad , \quad l < L_{1}$$
(5)

$$\frac{dl}{dt} = \frac{K_1}{\varphi_1 \eta} \frac{P_{app}}{l} \quad , \quad l < L_1 \tag{6}$$

18 Where L_1 is the characteristic length of the inter-tow region [m], K_1 is the inter-tow 19 permeability [m²], φ_1 is the inter-tow porosity, and P_{app} is the pressure applied by the vacuum 20 bag [Pa]. The viscosity along with the cure-dependent glass transition temperature (T_g) and the

- 1 DoC (α) are described using Equations A.1, A.2, and A.3, respectively, in Appendix A.
- 2 Supplementary material.
- For intra-tow flow, Equation 4 is integrated over pressure boundary conditions of $P|_{x=0} = P_{L1}$ and $P|_{x=l} = 0$,

$$\frac{dl}{dt} = \frac{K_2}{\varphi_2 \eta} \cdot \frac{K_1 P_{app}}{K_2 L_1 + K_1 (l - L_1)} \quad , \quad l \ge L_1 \tag{7}$$

5 Where the pressure at the boundary of the intra-tow region is,

$$P_{L1} = \frac{K_1 P_{app}(l - L_1)}{K_2 L_1 + K_1 (l - L_1)}$$
(8)

- 6 Where K_2 is the intra-tow permeability $[m^2]$ and φ_2 is the intra-tow porosity. The derivation of
- 7 Equation 8 is given in Appendix A. Supplementary material.

8



9

Figure 3. Physical representation of the 1D resin flow model at nodal position j and time step i. Resin flows into the fabric layer from above and below; in this case, the resin flow front position is defined as the distance from the inlet (dashed red line i.e. x = 0) to the flow front. The fabric layer is separated into inter-tow and intra-tow regions which are in series. The layer thicknesses are summed to calculate the total laminate thickness change.

15 Cender et al. [19] showed that the fill time is proportional to the square of the flow distance i.e. 16 halving the flow distance results in a factor of four reduction in the time taken to fill the fabric. 17 For this reason, it is important to account for resin flow into the fabric layer from above and

- 1 below, as illustrated in Figure 3. The total impregnation depth for each ply, $l_{tot(i,j)}$ [m] is found
- 2 by combining the resin flow front position from the top $(l_{top(i,j)})$ and bottom $(l_{bot(i,j)})$,

$$l_{tot(i,j)} = l_{top(i,j)} + l_{bot(i,j)}$$
(9)

4 2.3 Thickness change

5 It is known that resin flow in VBO prepregs corresponds with a reduction in laminate thickness 6 as the resin flows into the fabric layers [40,43]. For the finite difference code developed herein, 7 a simplified ply microstructure is defined so that this thickness change can be computed. As 8 mentioned in the previous section, it is assumed that compaction due to air evacuation has 9 already occurred, and the fibre-bed is incompressible. The simplified ply microstructure is 10 segmented into: a resin layer thickness, h_r [m]; a fabric layer thickness, h_{fab} [m]; and the total impregnation depth l_{tot} (see Figure 3). By summing these quantities, it is possible to determine 11 the total laminate thickness, h_{lam} [m] for each time step i, 12

$$h_{lam,i} = \sum_{j=1}^{N_p} l_{tot(i,j)} + \sum_{j=1}^{N_p} (h_{fab} - l_{tot(i,j)}) + \sum_{j=1}^{N_p} h_{r(i,j)}$$
(10)

13 Where N_p is the number of plies in the laminate.

14 As impregnation progresses, the thickness of the resin layer diminishes from its initial value 15 $h_{r(0,j)}$:

$$h_{r(i,j)} = h_{r(0,j)} - \beta \left(\varphi_{fab} h_{fab} \right) \tag{11}$$

16 Where φ_{fab} is the porosity of the fabric layer, and β is the degree of impregnation (DoI), which 17 can be written as,

$$\beta = \frac{l_{tot}}{\varphi_{fab}h_{fab}} \quad , \quad l_{tot} < L_1 \tag{12}$$

$$\beta = \frac{\varphi_1 h_{fab} + (l_{tot} - \varphi_1 h_{fab})\varphi_2}{\varphi_{fab} h_{fab}} \quad , \quad l_{tot} \ge L_1 \tag{13}$$

18 The characteristic length, L_1 is assumed to be equivalent to the inter-tow thickness i.e. $\varphi_1 h_{fab}$. 19 Assuming that there are no significant gaps between tows in the same fabric layer (i.e. the 20 vertices of each tow touch the vertices of the adjacent tows), the inter-tow porosity, φ_1 is constant for an elliptical tow geometry, bound at its vertices and co-vertices by a rectangular
 cross-section (red dashed line in Figure 2),

$$\varphi_1 = \frac{A_{rec} - A_{ell}}{A_{rec}} = \text{constant} \cong 0.2146 \tag{14}$$

3 Where A_{rec} is the total area of the rectangular cross-section [m²] outlined in Figure 2, and A_{ell} 4 is the area of the elliptical tow [m²].

5 For a known intra-tow porosity, φ_2 , the porosity of the fabric layer, φ_{fab} is determined as 6 follows [44,45],

$$\varphi_{fab} = \varphi_1 + \varphi_2 - \varphi_1 \varphi_2 \tag{15}$$

For a fully impregnated ply, l_{tot} is equivalent to h_{fab} such that the second term on the righthand side of Equation 10 cancels. Naturally, the calculation of ply thickness, h_{ply} reduces to the sum of the fabric thickness and any excess resin layer that remains:

$$h_{ply} = h_{fab} + h_r \tag{16}$$

10 The resin volume fraction, φ , determines the quantity of resin in the ply and can be written as,

$$\varphi = \frac{\varphi_{fab}h_{fab} + h_r}{h_{fab} + h_r} \tag{17}$$

11 Rearranging,

$$h_r = h_{fab} \left(\frac{\varphi - \varphi_{fab}}{1 - \varphi} \right) \tag{18}$$

12 Note: in the case that φ is less than φ_{fab} , this term becomes negative and the code outputs an

- 13 error messages warning that the plies will not achieve full impregnation.
- 14 Combining Equations 16 and 18, h_{fab} is approximated for a given φ ,

$$h_{fab} = h_{ply} \left(\frac{1 - \varphi}{1 - \varphi_{fab}} \right) \tag{19}$$

As can be seen with the above set of equations, only the cured ply thickness (h_{ply}) , the resin volume fraction (φ) , and the intra-tow porosity (φ_2) are required to fully describe the ply microstructure shown in Figure 3. The cured ply thickness and the resin volume fraction can be measured experimentally, while values of intra-tow porosity can be found in the literature. 1 Another important consideration for modelling the thickness change is the initial form of the 2 epoxy resin i.e. powdered or solid resin layer. For a powder layer, significant thickness 3 reduction is expected during the sintering process. As such, the thickness of the resin layer, 4 $h_{r(i,j)}$ is recalculated using,

$$h_{r(i,j)} = \frac{h_{r(i,j)}^*}{1 - \chi_{(i,j)}} \tag{20}$$

5 Where h^{*}_{r(i,j)} is the thickness of the fully sintered resin layer, and the powder void fraction,
6 χ_(i,j), is modelled using Equation 1.

7

8 2.4 Heat transfer

9 Transient one-dimensional (1D) heat conduction in a plane wall is used as the basis for the heat 10 transfer model under the assumption that convective heat transfer due to resin flow is negligible 11 [46,47]. The heat transfer equation is written using an explicit forward time, centred space 12 finite difference formulation (illustrated in Figure 4). For the model developed here, Δz can 13 vary as a function of resin flow in each fabric layer (i.e. heat transfer is coupled to the resin 14 flow model via thickness change).

15 The structure of the heat transfer model is also approximated as individual layers of resin (h_r) , 16 fabric (h_{fab}) , and impregnated fabric (l_{tot}) in series (see Figure 3). Each layer is treated as a 17 separate material with a unique thermal conductivity, which can be modelled as thermal 18 resistances in series. These resistances change based on the state of the resin (powder or liquid) 19 and the progression of resin flow. The contact resistance between these layers is assumed to be 19 negligible.

21 As such, the heat equation is approximated as,

$$\left(\frac{T_{i+1} - T_i}{\Delta t}\right)_j = \left(\frac{1}{\Delta z \rho c_P}\right)_{i,j} \left(\frac{T_{j-1} - T_j}{R_{j,j-1}} + \frac{T_{j+1} - T_j}{R_{j+1,j}}\right)_i + \left(\frac{m_r H_T}{c_P} \frac{d\alpha}{dt}\right)_{i,j}$$
(21)

Where subscripts *i* and *j* represent the time step and nodal position, respectively, *T* is temperature [K], Δt is the time step size [s], Δz is the element thickness of node *j* [m], ρ is density [kg/m³], c_P is specific heat capacity [J/kg K], $R_{j,j-1}$ is the thermal resistance of the material [m²K/W] between nodes *j* and *j* – 1, m_r is the resin mass fraction, and H_T is the total enthalpy of the curing reaction [J/g], which is 184 J/g [8]. The final term in Equation 21 couples

- 1 heat transfer to the cure kinetics model, which is given by Equation A.3 in Appendix A.
- 2 Supplementary material.
- 3



Figure 4. Illustration of the forward time, centred space scheme used for the heat transfer model. The
temperature for the time step, *i* + 1, is calculated explicitly based on the conditions at time step, *i*. The
various layers of resin, dry fabric and impregnated fabric are approximated as thermal resistances in
series.

9 The element thickness is approximated by,

$$\Delta z_{(i,j)} = h_{r(i,j)} + h_{fab} \tag{22}$$

10 The density of the material at each node is approximated by,

$$\rho_{(i,j)} = \rho_c \frac{h_{ply}}{h_{fab} + h_{r(i,j)}}$$
(23)

11 Where ρ_c is the composite density [kg/m³], which is determined via the rule of mixtures. The

- 12 specific heat capacity, c_P is also determined via the rule of mixtures method (see Appendix A.
- 13 Supplementary material for the corresponding equations; Equations A.7 and A.8, respectively).
- 14 The thermal resistances $R_{(i,(j,j-1))}$ and $R_{(i,(j+1,j))}$ are written as,

$$R_{(i,(j,j-1))} = \frac{h_{r(i,j-1)}}{2\kappa_{r(i,j)}} + \frac{l_{tot(i,j)}}{\kappa_{c,T(i,j)}} + \frac{h_{fab} - l_{tot(i,j)}}{\kappa_{fab}} + \frac{h_{r(i,j)}}{2\kappa_{r(i,j)}}$$
(24)

$$R_{(i,(j+1,j))} = \frac{h_{r(i,j)}}{2\kappa_{r(i,j)}} + \frac{l_{tot(i,j+1)}}{\kappa_{c,T(i,j+1)}} + \frac{h_{fab} - l_{tot(i,j+1)}}{\kappa_{fab}} + \frac{h_{r(i,j+1)}}{2\kappa_{r(i,j+1)}}$$
(25)

1 Where κ_r is the thermal conductivity of the epoxy [W/m K], $\kappa_{c,T}$ is the transverse thermal 2 conductivity of the impregnated composite [W/m K], and κ_{fab} is the thermal conductivity of 3 the dry fabric [W/m K].

4 The resin mass fraction, m_r is:

$$m_{r(i,j)} = \frac{\rho_{r(i,j)}(1 - V_f)}{\rho_f V_f + \rho_{r(i,j)}(1 - V_f)}$$
(26)

5 Where V_f is the volume fraction.

6

7 2.5 Thermal boundary conditions

8 Two sets of boundary conditions (BCs) are considered for this work: specified temperature 9 BCs for heated tooling; and forced convection BCs for oven heating. Heat transfer due to 10 radiation is not modelled explicitly.

11 As described by Bogetti and Gillespie [13], the specified temperature BC is implemented by 12 setting the surface node temperature at time step $i, T_{(i,s)}$ [K] equal to the programmed 13 temperature cycle, $T_{P(i)}$ [K]:

$$T_{(i,s)} = T_{P(i)} \tag{27}$$

For describing heat transfer by forced convection, the following expression from Çengel [48]is used,

$$\rho c_P \frac{\Delta z}{2} \frac{T_{(i+1,s)} - T_{(i,s)}}{\Delta t} = h \big(T_{P(i)} - T_{(i,s)} \big) + \kappa \frac{T_{(i,s\pm 1)} - T_{(i,s)}}{\Delta z}$$
(28)

16 Where *h* is the heat transfer coefficient (HTC) [W/m²K], and $T_{(i,s\pm 1)}$ is equal to $T_{(i,s+1)}$ for the 17 bottom BC and $T_{(i,s-1)}$ for the top BC.

For forced convection BCs, Zimmermann and Van Den Broucke [49] recommended that a convective HTC of $10 - 100 \text{ W/m}^2\text{K}$ be used for gases in an autoclave. Similarly, average convective HTCs have been experimentally measured in the range of $10 - 40 \text{ W/m}^2\text{K}$ for autoclaves [50,51]. A convective HTC of 40 W/m²K was chosen for all simulations in this study. The remaining material parameters in Equation 28 depend on the surface composition. For VBO prepreg manufacturing it is expected that the top boundary would consist of bagging materials, while the bottom boundary would be the B-side of the mould tool. Oh and Lee [27] assumed negligible thermal contact resistance between the bagging, laminate, and tooling; the same assumption is made in this work. Conduction through any additional layers (i.e. tooling, bagging, insulation, etc.) is modelled for all BCs. These additional layers are also modelled as thermal resistances in series. The corresponding equations are given in Appendix A. Supplementary material.

8

9 2.6 Material properties

10 This work focusses on manufacturing simulations for GRN 918 epoxy powder with glass-fibre 11 fabrics. Some important material characteristics of GRN 918 epoxy powder have already been 12 discussed, however, additional data for thermal conductivity and specific heat capacity is 13 required for heat transfer modelling. It is assumed that the specific heat capacity of GRN 918, 14 as a function of temperature and DoC, follows the empirical relationship given by Shin and 15 Hahn [28], and that it is valid for epoxy in both its powder and liquid form, as it is a quantity 16 which is independent of volume. Similarly, it is assumed that the thermal conductivity of the 17 fully sintered epoxy follows the empirical relationship given by Shin and Hahn [28] to describe 18 the thermal conductivity as a function of temperature and DoC.

To the author's knowledge, no data is available for the thermal conductivity of epoxy powders, however, the thermal conductivity of polyamide powder, $\kappa_{r,pow}$ was reported to be 0.05 – 0.0913 W/m K [52,53]. Given that the bulk thermal conductivity of polyamide and epoxy is similar (0.15 – 0.25 W/m K), data for polyamide powder is judged to be a reasonable approximation. The transition from powder to liquid resin is described for thermal conductivity using the following expression,

$$\kappa_r = \kappa_{r,pow} \, \frac{\chi}{\chi_0} + \kappa_{r,liq} \left(1 - \frac{\chi}{\chi_0}\right) \tag{29}$$

25 Where χ is the void fraction of the powder as determined using Equation 1, and χ_0 is the initial 26 void fraction of the powder.

An alternative to the above may be to use analytical models to determine the effective thermal conductivity of the epoxy powder. Several such models have been developed for porous solids [54]. These models are a function of void fraction and the thermal conductivities of the two 1 phases (i.e. epoxy powder and vacuum). These models return a range of thermal conductivities 2 for epoxy powder from 0.041 W/m K to 0.083 W/m K, which overlaps with the range reported 3 for polyamide powder. Note, this range was based on a void fraction of 0.503 and Shin and 4 Hahn's empirical relationship for the thermal conductivity of epoxy [28] at 55°C.

5 The thermal properties used for epoxy in this study are summarised in Table A.4 in Appendix 6 A. Supplementary material.

7 Characterisation of the fibre-bed was not within the scope of this project. For this reason, most 8 of the fibre-bed properties have been taken from the literature. Properties for stitched uni-9 directional (UD) fabrics and stitched triaxial fabrics were used in this study. The triaxial fabrics 10 are treated as 3 UD plies in series. Assuming that the thermal conductivity of a composite lamina is transversely isotropic, the model developed by Clayton [55] is used to describe the 11 through-thickness thermal conductivity, $\kappa_{c,T}$ of the UD plies. Some authors have 12 13 experimentally measured the through-thickness conductivity of dry fabric under vacuum 14 [56,57]. For dry GF fabric, they measured through-thickness thermal conductivities of 0.1 -0.2 W/m K, depending on the fibre architecture (i.e. stitched UD, plain weave, twill weave, 15

16 etc.).

17 Some properties and parameters are required to describe the fibre-bed for the flow models

18 also. The permeability of a fibre-bed can be modelled or measured experimentally, however,

19 reported values can vary significantly. The Gebart model for a quadratic packing structure

20 [58] is used to calculate the intra-tow permeability for all simulations presented here.

21 Experimental values from the literature were used for the inter-tow permeabilities of stitched 22 UD fabrics.

- 23
- 24
- 25
- 26
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29 Table A.5 in Appendix A. Supplementary material compiles the fibre-bed properties, 30 parameters, and models that are used for the process simulations.

Additional material properties are required for the bagging materials and the mould tooling.
 For simplicity, the bagging is treated as one layer and the values reported by Joshi et al. [59]
 are used (see Table A.6 in Appendix A. Supplementary material). The bagging layer thickness
 is assumed to be 1.5 mm.

5

6 2.7 Initial conditions

7 Table 2 summarises the initial conditions that are used for the process simulations. A 100-ply 8 glass-fibre/epoxy-powder laminate is used as a case study for the simulations. The cured ply 9 thickness (h_{ply}) of a thin laminate was measured experimentally. The initial FVF of the VBO prepreg is controlled during the prepregging process using automated powder deposition. In-10 plane resin flow is expected to be negligible during processing, so it is assumed that the final 11 12 FVF is approximately the same as the initial FVF i.e. no significant resin bleeding. An initial 13 DoI of 0.113 is chosen based on experimental fitting detailed in Part II of this study [34]; this 14 equates to a quarter of the available inter-tow space being filled during the powder deposition 15 process. It is assumed that the epoxy powder is brought above its T_q during the prepregging process so that it adheres to the fibre-bed, but still retains its powder form. It is assumed that 16 17 this prepregging process has a negligible effect on the DoC and there are negligible out-time 18 effects between prepregging and laminate processing. For initialising the heat transfer model, 19 this study assumed a uniform DoC (α) and temperature throughout the thick laminate. Based on real-world observations of vacuum bagged parts, an applied pressure of 90×10^3 Pa (90 kPa) 20 21 is used for simulating laminate manufacture; it has been shown that deficient pressure conditions commonly occur and can be an important factor in VBO processing [43]. 22

23

24 Table 2. Initial conditions used for the process simulations.

Parameter [units]	Value
No. of plies, N_p	100
Cured ply thickness, h_{ply} [mm]	1.0
Fibre volume fraction, V_f	0.5
Degree of impregnation, β	0.113

Powder void fraction, χ_0	0.485
Degree of cure, α	0.01
Applied pressure, P _{app} [Pa]	90×10^{3}
Laminate/bagging/tool temperature [°C]	23

2 2.8 Numerical computation

The finite difference code is run in MATLAB R2015a. To simulate the processing of a thicksection laminate, the laminate is discretized through the thickness, and the process models are numerically solved at the nodes. Many explicit and implicit finite difference methods exist for solving the heat equation, however, as previously indicated, an explicit forward time, central space scheme is used in this work. This is a first-order scheme, so it has limited accuracy (due to a large temporal truncation error), however, it is very simple to implement and is stable if the following criterion is met [48,60],

$$\gamma \frac{\Delta t}{\Delta z^2} < \frac{1}{2} \tag{30}$$

10 Where γ is the thermal diffusivity [m²/s],

$$\gamma = \frac{\kappa}{\rho c_P} \tag{31}$$

Similarly, the Euler method is implemented for solving any ordinary differential equations (ODEs) in the simulation (e.g. cure kinetics, resin flow, etc.). Again, this is an explicit first order method which is simple to implement but has limited accuracy.

14 The number of nodes, N_n is based on the number of plies i.e,

$$N_n = N_p + 1 \tag{32}$$

The thermal diffusivity of the laminate, as well as that of the bagging and tooling, must be considered when choosing the maximum allowable time step size for a simulation. In cases where aluminium or steel tooling is used, the thermal diffusivity of the tooling is much higher than the composite material itself and limits the time step size. The maximum allowable time step size for a 1 mm thick ply of GF/Epoxy-powder is approximately 2.80 s (see Figure A.1 in Appendix A. Supplementary material), while it is 0.57 s for a 10 mm aluminium tool, and 3.55 s for a 10 mm steel tool. For a small enough time step, first order methods provide sufficient 1 accuracy due to the low rates of change associated with manufacturing thick-section 2 composites i.e. slow ramp rates are used. As such, a time step of 0.5 s is chosen for all 3 simulations using an aluminium tool, while a time step of 2.5 s is chosen for all simulations 4 using a steel tool. Simulations show that there is no loss of accuracy when increasing from 0.5 5 s to 2.5 s increments for a steel tool (see Figure A.2 in Appendix A. Supplementary material).

6

7



8 Figure 5. Comparison of the results from RAVEN and from the finite difference code developed in 9 MATLAB. The simulations were carried out for a 96-ply GF/Epoxy-powder laminate heated on a 10 mm 10 steel plate in an oven.

As a final verification of the numerical methods, the solution of the finite difference code is compared with results from RAVEN, a commercially available software for performing 1D heat transfer and cure kinetics simulations on composite laminates. Figure 5 shows that there is little difference between the solutions (< 5°C at any time). The simulations did not include compaction due to resin flow because it was not a featured capability of RAVEN. Furthermore, cure kinetics were excluded as it was not possible to implement a user-defined cure kinetics model in RAVEN which could simulate the cure kinetics of the GRN 918 epoxy powder.

1 **3. Results and discussion**

2 **3.1** Powder sintering model

Figure 6 shows the results of the powder sintering model compared to the experimental data; converted from sample thickness to powder void fraction via Equations 2 and 3. As can be seen, Equation 1 is relatively accurate in describing the sintering behaviour of the GRN 918 powder. The sintering process corresponded to a 50% reduction of the powder sample's thickness. This highlighted that using epoxy powder would significantly increase the initial thickness of a composite preform compared to liquid epoxy.

9



10



Figure 6. Results of the sintering model and experimental data for GRN 918.

Discrepancies between the data and model could be a result of the experimental set-up, which 12 relied upon one-sided heating and an insulating sample hood. Use of a full environmental 13 14 chamber may have given better temperature control, but it was unavailable at the time of 15 testing. Another limitation of the test results was that the applied pressure was 52 kPa rather 16 than 101.35 kPa (i.e. it was roughly half the pressure that would be expected for VBO 17 processing). As such, the relationship of applied pressure and rate of sintering is not fully 18 understood, and an improvement would be to repeat tests under various levels of applied 19 pressure and determine if there is any significant effect.

1 **3.2** Case study – Oven processing of a 100-ply laminate

As a starting point for investigating thick-section processing, the standard manufacturing process for a 100-ply laminate was considered. A thick-section laminate of GF/Epoxy-powder may be processed on a heated tool or in an oven using an extended version of the standard temperature cycle used for thin laminates i.e. with longer isothermal dwells to allow heat to transfer through all the plies. For this case study, the simulations focussed on a laminate being processed in an oven, on a steel plate with one layer of vacuum bagging around it.

8 The following temperature cycle was used:

- 9 Drying stage: Ramp to 55°C and hold for 900 min
- 10 Impregnation stage: Ramp to 120°C at 1.5°C/min and hold for 360 min
- 11 Cure stage: Ramp to 180°C at 1.5°C/min and hold for 200 min

12 - Cool down

13

14 **3.3 Temperature profile**

15 Figure 7 shows how the temperature is predicted to evolve within the laminate with respect to 16 time. Despite having symmetric thermal BCs, the temperatures at the bottom and top of the laminate (Ply 1 and Ply 100, respectively) are asymmetric because of the presence of the tool 17 18 and bagging. During the drying stage (0 - 15 hr), the temperature at the centre of the laminate 19 lags behind the outside of the laminate significantly. Heat transfer improves as the epoxy 20 powder sinters, and the temperature of the laminate is predicted to approach the programmed 21 temperature due to a small amount of exothermic heat being generated by the curing reaction 22 (i.e. the DoC increases to approximately 0.2 during the drying stage; see Figure 8). The rate of heat transfer improves during the impregnation stage (15 - 22 hr), and again during the cure 23 24 stage (22 - 26 hr). This is due to the increasing thermal diffusivity of the material as the 25 laminate compacts and the thermal properties of the resin changes w.r.t. temperature and DoC; 26 see Figure A.3, Figure A.4, and Figure A.5 in Appendix A. Supplementary material for plots 27 of specific heat capacity, density and thermal conductivity, respectively. Naturally, the density 28 of the laminate increases as the powder sinters and the fabric is impregnated. The thermal 29 conductivity is influenced by these processes also due to the difference in thermal 30 conductivities for powder vs fully sintered resin (ratio of approximately 1:2) and dry GF fabric 31 vs fully impregnated GF fabric (ratio of approximately 1:1.3) - note, these ratios are

representative and assume ambient temperature and no curing. There is no thermal overshoot predicted for the laminate during the cure stage (i.e. above 120°C) due to GRN 918's low enthalpy of reaction. Conversely, the heat that is generated during the curing reaction aids heat transfer and allows the laminate to approach the programmed temperature, as previously mentioned for the drying cycle.

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7

8

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Figure 7. Plot of simulated ply temperatures within a 100-ply laminate. There is no temperature overshoot predicted, however, large thermal gradients are still present.

10

11 **3.4 Cure profile**

It has been demonstrated previously that GRN 918 epoxy powder can maintain a very low rate 12 of cure up to 120°C due to the use of latent curing agents; allowing sufficient time for fabric 13 14 impregnation to occur [8]. Despite this, the drying temperature (55°C) for the standard processing cycle is higher than the T_g of GRN 918 powder (approximately 40°C), resulting in 15 increased molecular mobility for the epoxy. As such, Figure 8 shows that the simulations 16 17 predict a slow increase in DoC during the drying stage. The rate of curing increases during the 18 impregnation stage as the temperature increases, however, none of the plies are predicted to 19 reach the gel point of the epoxy (i.e. DoC = 0.56) prior to the cure stage. Upon heating past 20 120°C, the DoC increases rapidly as the latent curing agent is activated. The entire laminate is 21 predicted to reach a DoC > 0.9 within the first 3 hr of the cure stage, which is approximately

- 1 15% of the total processing cycle time (the impregnation stage accounts for approximately
- 2 27%, and the drying stage accounts for approximately 58%).
- 3



4

5

Figure 8. Plot of the simulated DoC evolution within the laminate.

7 **3.5** Powder sintering, resin flow, and thickness change

8 An additional consequence of drying the powder above its T_a is that the powder is predicted to 9 melt and sinter as the viscosity of the epoxy resin decreases (see Figure 9). This is beneficial 10 for heat transfer in the laminate because fully sintered polymers have a higher thermal 11 conductivity than polymer powders. On the other hand, dynamic vapour sorption (DVS) 12 analysis has shown that powder sintering negatively affected the desorption properties of the epoxy [61]. By sintering the powder, moisture can no longer desorb from the surface of the 13 14 powder particle and must diffuse through the bulk polymer. Furthermore, the powder acts as 15 permeable layer through which gases can be evacuated, whereas the sintered epoxy forms a barrier and the gases must pass through the dry fibre tows; similar to conventional VBO 16 17 prepregs [38,62].





Figure 9. Powder sintering during the drying stage.

The sintering process also results in a significant decrease in the laminate thickness; approximately 26% as shown in Figure 10. Ideally, a shortened drying stage would be carried out below the T_g of the epoxy powder (due to the advantageous sorption properties of the powder), and then an additional stage at an intermediate temperature could be introduced to promote powder sintering and improve heat transfer within the laminate. To optimise the drying stage in this way, more research is required which is outside the scope of this work.

9 After the drying stage, the remaining thickness reduction predicted (19%) in Figure 10 is due 10 to resin flow; in total the laminate thickness decreases by approximately 45%. As shown in Figure 11, the simulation predicts that the inter-tow region (DoI < 0.45) would fill during the 11 12 drying stage when powder has sintered and reaches a sufficiently low viscosity to flow between 13 the tows. For the outer plies, it is predicted that the inter-tow region would be completely filled 14 within the first few hours of the temperature cycle, however, the inner plies lag by several hours 15 due to slow heat transfer. The timescale for this process confirms the need to model inter-tow 16 flow for the current material system. In contrast, intra-tow filling (DoI > 0.45) is relatively fast 17 when the resin viscosity is sufficiently low (see Figure 12 for the viscosities).



Figure 10. The thickness change plot of the GF/Epoxy-powder laminate during the temperature cycle. A
large reduction in thickness is predicted during the drying stage due to the sintering of the powder. After
the cessation of sintering, the resin is predicted to slowly flow into and fill the inter-tow region of the
fibre-bed. During the impregnation stage, the viscosity of the resin reduces, the resin fills the fibre tows,
and the laminate thickness decreases again.





8

Figure 11. Plot of the degree of impregnation during the simulated temperature cycle. The inter-tow
 region (DoI < 0.45) fills during the drying cycle, but lower viscosities are required for filling the fibre tows
 (DoI > 0.45).

12 It is noted that the simulated viscosities in Figure 12 are relatively high for GF fabric 13 impregnation, however, additional calculations using the analytical solution described by 14 Cender et al. [19] confirms that, even for a resin viscosity of 1000 Pa s, a ply could be fully 15 impregnated within 80 mins. Whether this is true remains unclear, as other factors can inhibit

- 1 the completion of filling (e.g. pressure sharing of the fibre-bed, pressure deficient conditions,
- 2 etc.).
- 3



5 Figure 12. Plot of the resin viscosity during the impregnation and cure stages of the temperature cycle 6 (note the reduced time span on the horizontal axis). The viscosities are predicted to drop significantly 7 when the temperature is increased to 120°C. Upon heating above 120°C, the epoxy reaches its gel point, 8 and the viscosity tends towards infinity.

9

10 **4.** Conclusions

11 The development of 1D numerical models for manufacturing thick-section composites using 12 powder-based VBO prepregs has been described. This includes process models for coupled 13 resin flow and heat transfer within VBO prepregs, and material models specifically developed 14 for an epoxy powder system. Using first order numerical methods, it was possible to solve these 15 models for a typical composites temperature cycle. The accuracy of the heat transfer model 16 was verified by comparing the results of the code to results from the commercial composite 17 processing software, RAVEN. Due to limitations with RAVEN, it was not possible to verify 18 or validate the cure kinetics model, resin flow model, or powder sintering model. Validation 19 experiments for these models are the subject of concurrent work (Part II [34]).

A case study was carried out on a 100-ply laminate which was processed in an oven on steel tooling. The study was used to demonstrate the capability of the numerical models and to analyse a typical temperature cycle used by industry for manufacturing thick-section composite parts. For the drying stage, it was found that the current drying temperature was high enough to melt and sinter the powder throughout the laminate. This plays an important role in improving heat transfer within the laminate but may not be efficient in terms of drying the epoxy. With more analysis and modelling of moisture desorption and gas evacuation, it would possible to optimise the drying stage of the temperature cycle, however, this was outside the scope of this work.

8 An existing dual-scale resin flow model for VBO prepreg was used as the basis of the resin 9 flow in this system. This dual-scale model simulated resin flow around the fibre tows (inter-10 tow) and into the fibre tows (intra-tow). It was found that the resin viscosity profile varied 11 significantly depending on the resin's position within the laminate. Due to the advancement of 12 cure, the resin viscosity did not reach below 100 Pa s, however, the resin flow model still 13 predicted that the fibre-bed would reach full impregnation during the 120°C dwell. The fabric 14 impregnation, along with powder sintering, contributed to a 45% reduction in the thickness of 15 the laminate. This thickness change was computed based on a simplified microstructural model 16 which required values for the cured ply thickness, the intra-tow porosity, and the fibre volume 17 fraction of an individual ply. This microstructural model also played a role in coupling resin 18 flow to the heat transfer model by updating each ply's thickness, and thermal properties.

Simulated temperature profiles for the 100-ply laminate showed that heat transfer improved over the duration of the processing cycle. Powder sintering and fabric impregnation contributed to increases in thermal conductivity. Temperature-dependent and cure-dependent material properties were taken from the literature; these contributed to improved heat transfer also, especially during the cure stage. The low enthalpy of reaction for GRN 918 epoxy powder meant that there was no significant build-up of heat predicted in the laminate and subsequently no overshoot in temperature, thus eliminating the risk of thermal runaway.

26

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6

Nomenclature [units] 7

8	α	Degree of cure	38	A _{rec}	Area of a rectangle [m ²]
9	β	Degree of impregnation	39	В	Fitting constant
10	γ	Thermal diffusivity [m ² /s]	40	C_P	Specific heat capacity [J/kg K]
11	η	Viscosity [Pa s]	41	$C_{\chi 1}$	Fitting constant for sintering model
12	κ	Thermal conductivity [W/m K]	42	$C_{\chi 2}$	Fitting constant for sintering model [K]
13	$\kappa_{c,T}$	Transverse thermal conductivity of the	43	h	Heat transfer coefficient [W/m ² K]
14		impregnated composite [W/m K]	44	h_{fab}	Thickness of the fabric layer [m]
15	κ _{fab}	Thermal conductivity of the dry fabric	45	h _{lam}	Total laminate thickness [m]
16		[W/m K]	46	h_{ply}	Thickness of a fully impregnated ply (cured
17	κ _r	Thermal conductivity of the epoxy	47		ply thickness) [m]
18		[W/m K]	48	h_r	Thickness of resin layer [m]
19	$\kappa_{r,liq}$	Thermal conductivity of the liquid epoxy	49	H_T	Total enthalpy of reaction [J/g]
20		[W/m K]	50	i	Time step for finite difference code
21	$\kappa_{r,pow}$	Thermal conductivity of the epoxy powder	51	j	Through-thickness nodal position for finite
22		[W/m K]	52		difference code
23	ρ	Density [kg/m ³]	53	Κ	Permeability [m ²]
24	$ ho_0$	Initial density of the powder [kg/m ³]	54	<i>K</i> ₁	Inter-tow permeability [m ²]
25	$ ho_c$	Composite density [kg/m ³]	55	<i>K</i> ₂	Intra-tow permeability [m ²]
26	$ ho_f$	Fibre density [kg/m ³]	56	L	Thickness of sintered powder samples [m]
27	$ ho_P$	Bulk polymer density [kg/m ³]	57	L_0	Initial thickness of powder samples [m]
28	$ ho_r$	Resin density [kg/m ³]	58	L_1	Characteristic length of the inter-tow
29	φ	Total porosity/resin volume fraction	59		porous medium [m]
30	$arphi_1$	Inter-tow porosity	60	l	Resin flow front distance from the inlet [m]
31	φ_2	Intra-tow porosity	61	l_{bot}	Resin flow front distance from the bottom
32	φ_{fab}	Porosity of the fabric layer	62		of the fabric layer [m]
33	χ	Powder void fraction during sintering	63	l_{top}	Resin flow front distance from the top of
34	χo	Initial powder void fraction of the powder	64		the fabric layer [m]
35	χ_E	Pre-exponential rate constant for sintering	65	l _{tot}	Total impregnation depth [m]
36	χ_∞	Final powder void fraction during sintering	66	m_r	Resin mass fraction
37	A_{ell}	Area of an ellipse [m ²]	67	N_n	Number of nodes

1	N_p	Number of plies	9	$T_{i,s}$	Surface temperature at time step <i>i</i> [K]
2	Р	Pressure [Pa]	10	$T_P(i)$	Programmed temperature at time step i [K]
3	P_{app}	Applied pressure [Pa]	11	T_{θ}	Onset temperature for melting [K]
4	$R_{j,j-1}$	Thermal resistance of material between	12	t	Time [s]
5		nodes j and $j - 1 [m^2 K/W]$	13	Δt	Time step size for finite difference code [s]
6	$R_{j+1,j}$	Thermal resistance of material between	14	V_f	Fibre volume fraction
7		nodes $j + 1$ and $j [m^2 K/W]$	15	x	Flow direction
8	Т	Temperature [K]	16	Δz	Element thickness of node <i>j</i> [m]

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1 Appendix A. Supplementary material

2 Additional material models from Maguire et al. [8]

- 3 The viscosity, η , is described using the following WLF-type chemorheological model which
- 4 was developed by Kenny & Opalicki [63] for toughened epoxy systems,

$$\eta = \eta_{g0} \, \exp\left(\frac{-C_{\eta1}[T - T_g]}{C_{\eta2} + T - T_g}\right) \left(\frac{\alpha_g}{\alpha_g - \alpha}\right)^A \tag{A.1}$$

- 5 Where η_{g0} is the viscosity of the uncured resin [Pa s] at the initial glass transition temperature,
- 6 T_{g0} [K], α_g is the DoC at gelation, $C_{\eta 1}$, $C_{\eta 2}$ [K], and A are fitting constants. The cure-dependent
- 7 glass transition temperature, T_g [K], was modelled using the DiBenedetto equation [64],

$$\frac{T_g - T_{g0}}{T_{g\infty} - T_{g0}} = \frac{\lambda\alpha}{1 - (1 - \lambda)\alpha}$$
(A.2)

8 Where T_{g0} is the initial glass transition temperature of the uncured resin [K], $T_{g\infty}$ is the glass 9 transition temperature of the fully cured resin [K], and λ is a fitting constant.

The viscosity couples the resin flow model to the heat transfer and cure kinetics models via the
temperature dependency and DoC dependency, respectively. The following cure kinetics
model was developed for GRN 918 [8],

$$\frac{d\alpha}{dt} = \frac{(k_{\alpha 1} + k_{\alpha 2} + k_{\alpha 3} \alpha^m)(1 - \alpha)^n}{1 + \exp[C(\alpha - \alpha_c)]}$$
(A.3)

13 Where $k_{\alpha 1}$, $k_{\alpha 2}$, and $k_{\alpha 3}$ are cure rate constants [s⁻¹], *m* and *n* are the reaction orders, *C* is a 14 diffusion constant, and α_c is the temperature-dependent critical DoC, above which the reaction 15 becomes diffusion-controlled.

16 The fitting parameters for Equations A.1, A.2, and A.3 are given in Table A.1, Table A.2, and

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3	Table A.3 respectively.
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8 Table A.1. Chemorheological model parameters for GRN 918 epoxy powder.

Parameter [units]	Value	
η_{g0} [Pa s]	2×10^{11}	
$C_{\eta 1}$	32.25	
<i>C</i> _{<i>η</i>²} [K]	30	
$lpha_g$	0.56	
Α	1.6	
$\eta = \eta_{g0} \exp\left(\frac{-C_{\eta1}[T - T_g]}{C_{\eta2} + T - T_g}\right) \left(\frac{\alpha_g}{\alpha_g - \alpha}\right)^A$		

10 Table A.2. Parameters for the DiBenedetto model.

Parameter [units]	Value
<i>T_{g0}</i> [°C]	40
$T_{g\infty}$ [°C]	106
λ	0.53
	$\frac{T_g - T_{g0}}{T_{g\infty} - T_{g0}} = \frac{\lambda \alpha}{1 - (1 - \lambda)\alpha}$

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7 Table A.3. Fitted cure kinetics parameters for GRN 918 epoxy powder.

Parameter [u	nits] Val	ue	Parameter [units]	Value
$A_{\alpha 1} [s^{-1}]$	4.07	73×10^{-4}	m	1.24
$E_{\alpha 1}$ [J/mol]	120	06	n	1.8
$A_{\alpha 2} [s^{-1}]$	10.	112×10^{9}	С	50
$E_{\alpha 2}$ [J/mol]	111	792	α _c	0.006 <i>T</i> - 1.748
$A_{\alpha 3} [s^{-1}]$	1.63	36×10^{13}		
$E_{\alpha 3}$ [J/mol]	131	240		
	$\frac{d\alpha}{dt} = \frac{(k_{\alpha 1} + k_{\alpha 2})}{1 + e^{2}}$	$\frac{+k_{\alpha 3}\alpha^m)(1-\alpha)^n}{\exp[C(\alpha-\alpha_c)]} ,$	$k_{\alpha i} = A_{\alpha i} \exp\left(\frac{-E_{\alpha i}}{RT}\right)$,	<i>i</i> = 1,2,3

8

9 Derivation of the pressure boundary condition at the interface of two porous media

10 For intra-tow flow, the pressure boundary condition at the interface of the two porous media

11 P_{L1} must be derived. Assuming mass conservation for resin flow through the porous media,

$$\frac{du}{dx} = 0 \tag{A.4}$$

12 Then,

$$\frac{-K_1(P_{app} - P_{L1})}{\eta L_1} = \frac{-K_2(P_{L1} - P_f)}{\eta (l - L_1)}$$
(A.5)

13 Where P_f is the pressure at the flow front (assumed to be zero under vacuum conditions). As 14 such, the equation reduces to:

$$P_{L1} = \frac{K_1 P_{app}(l - L_1)}{K_2 L_1 + K_1 (l - L_1)}$$
(A.6)

2 <u>Rule of mixtures equations for the heat transfer model</u>

A rule of mixtures approach was implemented by several authors for describing the composite
density and specific heat capacity within the heat equation [22,23,49,59,65,66]:

$$\rho_c = \rho_f V_f + \rho_r (1 - V_f) \tag{A.7}$$

$$c_{P,c} = \frac{c_{P,f}\rho_f V_f + c_{P,r}\rho_r (1 - V_f)}{\rho_f V_f + \rho_r (1 - V_f)}$$
(A.8)

5 Where V is the volume fraction, and subscripts f and r denote fibre and resin, respectively.

6

7 Equations for heat transfer through the bagging or tooling

8 Heat transfer at the boundary of the laminate and bagging was written as,

$$\left(\Delta z \rho c_P \frac{dT}{dt}\right)_{i,B} = \left(\frac{T_{j-1} - T_B}{R_{B,j-1}} + \frac{T_B - T_{B+1}}{R_{B+1,B}}\right)_i + \left(z_r \rho_r (1 - V_f) H_T \frac{d\alpha}{dt}\right)_{i,B}$$
(A.9)

9 Where subscript *B* represents the node at the boundary and is equivalent to node *j* in the 10 laminate. Note that the above can be easily written for the boundary of the laminate and tooling 11 by changing the subscripts appropriately.

12 The average spatial difference, Δz , was approximated by,

$$\Delta z_{(i,B)} = \frac{2h_{r(i,j)} + h_{fab} + 2z_{bt}}{2} \tag{A.10}$$

13 Where z_{bt} is the spatial increment size for the bagging/tooling, represented by,

$$z_{bt} = \frac{h_{bt}}{N-1} \tag{A.11}$$

- 14 Where h_{bt} is the total thickness of the bagging/tooling, and *N* is the total number of nodes for 15 the discretised bagging/tooling.
- 16 The combined density and specific heat capacity at the boundary node were approximated
- 17 using a rule of mixtures approach,

$$\rho_{(i,B)} = \frac{\rho_{bt} z_{bt} + \rho_j (h_{r(i,j)} + h_{fab}/2)}{\Delta z_{(i,B)}}$$
(A.12)

$$c_{P,B} = \frac{c_{P,bt}\rho_{bt}z_{bt} + c_{P(i,j)}\rho_{(i,j)}(h_{r(i,j)} + h_{fab}/2)}{\rho_{bt}z_{bt} + \rho_{(i,j)}(h_{r(i,j)} + h_{fab}/2)}$$
(A.13)

1 Where the subscript *bt* refers to bagging/tooling.

2 The thermal resistance, $R_{B,j-1}$, was taken to be equivalent to $R_{j,j-1}$ (Equation 24). If the

3 thermal conductivity and thickness of each bagging layer is known, a full thermal resistance

4 term can be deduced. For this work, however, $R_{B+1,B}$ was simply written as,

$$R_{B+1,B} = \frac{z_{bt}}{\kappa_{bt}} \tag{A.14}$$

5 Where κ_{bt} was a lumped thermal conductivity for the whole bagging layer.

6

9

7 Properties and parameters used in the process models

8 Table A.4. Thermal properties used to model heat transfer in GRN 918.

Property [units]	Value	Source	
Specific heat capacity of epoxy, $c_{P,r}$ [J/kg °C]*	Equation A.15	[28]	
Thermal conductivity of liquid epoxy, $\kappa_{r,liq} [W/m ^{\circ}C]^*$	Equation A.16	[28]	
Thermal conductivity of powder epoxy, $\kappa_{r,pow}$ [W/m K]	0.075**	-	
$c_{P,r} = 4184(0.468 + 5.975 \times 10^{-4}T - 0.141\alpha)$			(A.15)
$\kappa_{r,liq} = 0.04184 [3.85 + (0.035 T - 0.41) \alpha]$			(A.16)

* Predicted temperatures within the simulation are converted to °C so that the empirical relationships can be
used.
** Average value taken from [52,53]

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Property/Parameter [units]	Value	Source
Transverse thermal conductivity of impregnated GF fabric, $\kappa_{c,T} [W/m K]^*$	Equation A.17	[55]
Thermal conductivity of glass fibre, κ_f [W/m K]	0.417	[67]
Thermal conductivity of dry GF fabric, κ_{fab} [W/m K]	0.2	[57]
Density of E-glass fibre, ρ_f [kg/m ³]	2560	[68]
Specific heat capacity of E-glass fibre, $c_{P,f}$ [J/kg K]	810	[26]
Inter-tow permeability of stitched GF fabric, $K_1 \ [m^2]^{**}$	13.675×10^{-10}	[20]
Intra-tow permeability of GF K_2 [m ²]	Equation A.18	[58]
Glass fibre radius, R_{fib} [m]	$8.0 imes 10^{-6}$	[69]
Fibre volume fraction of glass fibre tow, $V_{f,tow}$	0.67	[45]

4 Table A.5. Properties and parameters used for glass fibre fabric.

$$\frac{\kappa_{c,T}}{\kappa_r} = \frac{1}{4} \left[\sqrt{\left(1 - V_f\right)^2 \left(\frac{\kappa_f}{\kappa_r} - 1\right)^2 + \frac{4\kappa_f}{\kappa_r}} - \left(1 - V_f\right) \left(\frac{\kappa_f}{\kappa_r} - 1\right) \right]^2 \tag{A.17}$$

$$K_{2} = K_{\perp quad} = \frac{16}{9\pi\sqrt{2}} \left[\sqrt{\frac{\pi}{4V_{f,tow}}} - 1 \right]^{5/2} R_{fib}^{2}$$
(A.18)

6 * The Springer-Tsai model [70] can also be used with little change to the overall result.

*** Average value for stitched GF fabric.*

4 Table A.6. Properties used for bagging and tooling materials.

Property [units]	Value	Source
Thermal conductivity of bagging [W/m K]	0.069	[59]
Thermal conductivity of aluminium [W/m K]	216.3	[59]
Thermal conductivity of tool steel [W/m K]	53.35	[51]
Density of bagging [kg/m ³]	355.6	[59]
Density of aluminium [kg/m ³]	2692.1	[59]
Density of tool steel [kg/m ³]	7822.8	[51]
Specific heat capacity of bagging [J/kg K]	1256.0	[59]
Specific heat capacity of aluminium [J/kg K]	916.9	[59]
Specific heat capacity of tool steel [J/kg K]	485.0	[51]



Figure A.1. The maximum allowable time step to achieve a stable solution for a GF/Epoxy-powder ply.
 The curing and compaction during the cycle causes the thermal diffusivity, γ, and the spatial increment,
 Δz, to change. This results in the max time step being reduced.



Figure A.2. Test of simulation accuracy for two time step sizes: 0.5 s, and 2.5 s. Heat transfer through a
 96-ply GF/Epoxy-powder laminate is simulated with a 10 mm steel tool.



Figure A.3. Plot of the specific heat capacity variation during the temperature cycle. It increases with
 temperature, and it decreases with the advancement of cure.



Figure A.4. Plot of the density variation of the plies during the temperature cycle. The ply density
 increases as the epoxy powder sinters and infuses into the fibre-bed.





Figure A.5. Plot of the effective thermal conductivity variation during the temperature cycle. It increases
 with temperature, and is also affected by sintering of the powder and impregnation of the fibre-bed. The
 values are back-calculated from the thermal resistances of each ply.