Ceramic Binder Jetting Additive Manufacturing: A Literature Review on Density

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Abstract

The objective of this review paper is to summarize the current status and identify the knowledge gaps in ceramic binder jetting additive manufacturing, in a particular focus on density. This paper begins with an overview of the process, material considerations, and process parameters. It then discusses different aspects of density, including various terminologies, measurement methods, and achieved values. Afterwards, it reviews two categories of techniques to increase the part density: material preparation techniques (powder granulation, mixing powders of different sizes, using slurry feedstock, and mixing different materials) and post-processing techniques (sintering, chemical reaction, infiltration, and isostatic pressing). Finally, it presents the knowledge gaps in the literature.

1 Introduction

Ceramic materials can have outstanding properties, such as extraordinary hardness, excellent resistance to wear, heat, and corrosion, and exceptional biocompatibility. Therefore, ceramic materials have a wide range of applications, from orthopaedic and dental implants in the biomedical industry to engine components in the aerospace and automotive industries. However, it is very costly to fabricate ceramic parts of complex shapes using conventional manufacturing techniques. For complex ceramic parts, tooling can contribute up to 80% of the overall cost if conventional techniques are used [1]. Conventional techniques have other disadvantages including excessive cost in prototyping and difficulty to make design changes. All these disadvantages have impeded the widespread applications of advanced ceramic materials (e.g., alumina, zirconia, and silicon carbide). For example, there are millions of joint replacement surgeries every year [2]. Although ceramic materials are an excellent choice for joint implants because of their excellent wear resistance and exceptional biocompatibility [3], they are not widely utilized because it is not cost-effective to fabricate them using conventional manufacturing technologies [4].

According to the ASTM standard [5], most additive manufacturing (AM) technologies can be classified into seven categories: binder jetting, vat photopolymerization, powder bed fusion, directed energy deposition, material extrusion, material jetting, and sheet lamination. Binder jetting is defined by ASTM as "additive manufacturing processes in which a liquid bonding agent is selectively deposited to join powder materials" [5]. Binder jetting was initially developed at Massachusetts Institute of Technology in the late 1980s [6,7] and commercialized by Soligen in 1993 [8], Z Corporation [9] and Therics [10] in 1997, ExOne in 2001 [11], Voxeljet in 2002 [12], Microjet in 2016 [13], and Desktop Metal [14] and Digital Metal [15] in 2017. After presenting their innovation in various conference [7,16,17] and journal [18,19] papers, the inventors investigated various perspectives of this technology, including the powder bed density improvement [20,21], powder-binder interaction [22,23], new feedstock form (i.e., slurry instead of powder) [24–31], and potential application areas (e.g., casting cores and shells [32], optical lenses [29], cutting tools [33], and biomedical devices [34]).

Table 1 summarizes advantages and disadvantages of the seven AM categories for printing ceramic materials. Compared with others, binder jetting has some unique features. First, the surrounding powder supports the printed part during the building process. Therefore, there is no need for explicit support structures for overhangs and undercuts [35]. Second, the amount of binder in the green body is low, and thus debinding is much easier for large parts than some other processes such as vat photopolymerization [36,37]. Third, as print heads can consist of up to thousands of jets working in parallel, binder jetting is easily scalable for fabricating large parts [38]. Moreover, binder jetting is suitable for biomedical applications due to its capability of printing functionally graded materials [39]. For example, bioinks with drugs can be added to the binder [40].

Category	Advantage	Disadvantage
Binder jetting	No need for support [35], minimal amount of sacrificial materials [37], high scalability [38]	High porosity [41]
Vat photopolymerization	Excellent resolution and accuracy [42], smooth surface [42], high density [41]	Limited part size
Powder bed fusion (indirect)	No need for support	High porosity
Material extrusion	Low cost, high density, multiple materials within a part [43]	Low resolution [41], interfacial porosity
Material Jetting	Excellent surface quality [42], multiple materials within a part	Limited part size
Directed energy deposition	High density [44]	Thermal cracks [45]
Sheet lamination	High speed, no need for support	Delamination, interfacial porosity [41]

Table 1. Advantages and disadvantages of various AM categories for ceramics

Several review papers on ceramic AM have discussed binder jetting. The first one [46], which was published in 2003, discussed its process, dimensional control, and applications. Other four

ceramic AM review papers, published in 2014 [42,47], 2015 [41], and 2017 [48], also included binder jetting. Travitzky et al. [42] reviewed the dimensional control methods and several applications in the orthopaedic field. Deckers et al. [47] reviewed the low-density issue of ceramic binder jetting. Zocca et al. [41] reviewed the material preparation methods to improve the powder bed density, while Yang et al. [48] discussed the advantages and material limitations of ceramic binder jetting. A recent review paper about binder jetting summarized various powder deposition methods, reviewed binder selection criteria, and discussed key printing parameters such as binder saturation and layer thickness [49]. In addition to these review papers, a book chapter also reviewed some ceramic materials that have been used in binder jetting, including several applications related to these materials [50].

The literature has reported an increasing number of studies on binder jetting. Various techniques have been reported to improve the density (and mechanical properties in some cases) of printed parts. However, there are no review papers devoted to ceramic binder jetting that comprehensively compile the available reports in the literature on density improvement. This literature review is to fill the gap.

This paper first provides an overview of ceramic binder jetting, including the process, materials, and resultant densities. The focus of this review is to summarize and analyze reported techniques for density improvement, which is considered as the most challenging issue in this field. Finally, existing knowledge gaps are identified.

2 Description of Ceramic Binder Jetting

2.1 Process overview

Figure 1 illustrates major components of a binder jetting 3D printer, including powder stock, build platform, spreader, binder cartridge, and print head. The printing process is as follows. First, the spreader deposits a thin layer of powder from the powder stock onto the build platform, forming the powder bed. Then the print head jets binder onto selected areas defined by a 3D model to bond the powder particles in these areas. After one layer is finished, the build platform is lowered and the powder stock is raised, after which a new powder layer is spread onto the finished layer. These steps are repeated until the whole part is printed. The printed part is then separated from the loose powder after curing [6,37].



Figure 1. Major components of a binder jetting 3D printer

A generic cycle of ceramic binder jetting is shown in Figure 2. The feedstock is ceramic powder. The powder needs to meet certain requirements (e.g., high flowability and sinterability to be discussed in Section 2.3) for making high-quality parts. The printed part is called "green part". The green part is then heated to a specific temperature (usually around 200°C), during which the

binder is thermally activated by solvent evaporation, polymerization, cross-linking, or other mechanisms to strengthen the green part [42,47]. It should be noted that curing can be applied layer by layer [51]. Curing temperature could be higher (up to 800°C) if inorganic binder, usually colloidal silica [18], is used.

Afterward, post-processing steps (such as debinding and sintering) are usually applied to the green parts to improve their material properties. In the debinding step, the binder is thermally decomposed or burned out by heat treatment between 400 °C and 800 °C, producing a "brown part". The heating rate is controlled to ensure the escape of gaseous reaction products. In the sintering step, the part is heated to a high temperature (usually just below the melting point of the ceramic material), followed by dwelling and furnace cooling. The sintering step increases the density of the part by mass transport across the boundaries of ceramic particles [52]. Other densification techniques (e.g., isostatic pressing, infiltration, etc.) can also be used to further increase the density and thus mechanical properties. However, they may compromise other properties such as geometrical accuracy, biocompatibility, and heat resistance. Details of some post-processing steps are reviewed in Section 5.



Figure 2. Steps of ceramic binder jetting

2.2 Materials and applications

Figure 3 shows the proportions of application areas of ceramic binder jetting in reported studies. The application with the highest number of papers is in the orthopaedic field to make bone scaffolds and implants. The structural application has the second highest number of papers. A small number of dentures and crowns are reported as well. For electronic applications, fabrication of dielectric radio frequency filters and ferroelectric dielectric capacitors is reported. There is also one paper about the fabrication of a gradient-index lens for optical application.



Figure 3. Proportions of various application areas of ceramic binder jetting reported in the literature

Table 2 summarizes the reported materials grouped by their application areas. Sometimes, the feedstock and resultant materials (if identified) are different and thus listed in different columns. Additives and infiltrants are also included.

Application	Feedstock material	Additive (A) or infiltrant (I)	Resultant material	Reference
Structural	Al	/	Al ₂ O ₃	[53]
Structural	Al+Al ₂ O ₃	/	Al ₂ O ₃	[54]
Structural	AlN	/	AlN	[55,56]

Table 2. Feedstock and resultant materials in reported studies on ceramic binder jetting

Structural	Al ₂ O ₃	/	Al ₂ O ₃	[7,18,19,23 ,24,38,57–
Structural	A1.O.	(\mathbf{I}) $\mathbf{A1}$.O.	A1-O-	68] [60]
Structural		$(I) AI_2O_3$ $(I) Cu+Cu_2O_3$	Al_2O_3 $Al_2O_3 + Cu + Cu_2O_3$	[67 70]
Structural		(I) $Cu + Cu_2O$	$Al_2O_3 + Cu + Class$	[07,70]
Structural	Al_2O_3 Al_2O_2+ ZrO_2	(1) 01ass	$Al_2O_3 + Olass$	[72]
Structural	$R_1 \ge 0_3 + \Sigma_1 O_2$	/	$R_1 \ge 0_3 + \Sigma_1 = 0_2$ $R_2 = C + S_1 = C$	[72]
Structural	$D_4C + SIC$	$(\Lambda) 7r\Omega_{r}$	$D_4 C + S C$	[73]
Structural		$(A) ZIO_2$	$C_{a}SO_{10}SH_{10}$	[74]
Structural	$C_{a}SO_{4} 0.5H_{2}O$	$(I) C \cdot H \cdot C O$	$C_{a}SO_{4} = 0.511_{2}O_{2}$	[75]
Structural	Porcelain	(1) C211125C1O5	Porcelain	[70]
Structural	Si	/	SiC+Si ₂ N ₄ +SiON	[//] [78 79]
Structural	Si+SiC	(I) Silicone	sision signation	[70,79]
Structural	SIC			[81.82]
Structural	Silicona		SISIC	[82]
Structural	Sint	/	Sint	[03]
Structural	S131N4 SiO-	(I) A1	$\Delta 1 \pm \Delta 1_{\odot} O_{\odot}$	[24]
Structural		$(I) \mathbf{A} \mathbf{I}$	$\mathbf{A}_{1} + \mathbf{A}_{12} \mathbf{O}_{3}$ $\mathbf{T}_{12} \mathbf{S}_{12} \mathbf{C}_{1} + \mathbf{T}_{12} \mathbf{C}_{12} \mathbf{S}_{12} $	[04]
Structural	TiC+TiO	(I) SI (I) A1	$A_1 + A_1 + A_2 + T_1 $	[86]
Siructural	$\Pi C + \Pi O_2$	(I) AI	$A_1 + A_1 O + T; A_1 + T; A_1 C + T;$	[80]
Structural	TiC+TiO ₂	(I) Al	$A_1 + A_{12}O_3 + 11A_{13} + 11_3A_1C_2 + 11$ C	[87]
Structural	TiC+TiO ₂	(I) Al	Al ₂ O ₃ +TiAl ₃ +Ti ₃ AlC ₂	[88]
Structural	TiO ₂	(I) [CH ₃ CH(O-)CO ₂ NH ₄] ₂ Ti(OH) ₂	TiO ₂	[89]
Structural	Ti ₃ SiC ₂	/	Ti ₃ SiC ₂	[90,91]
Structural	WC+Co	/	WC+Co	[92–94]
Structural	ZrO_2	/	ZrO ₂	[95]
Orthopaedic	Bioactive glass	/	Bioactive glass	[96–98]
Orthopaedic	Bioactive glass+Ca ₅ (PO ₄) ₃ (OH)	/	Bioactive glass+Ca ₅ (PO ₄) ₃ (OH)	[99]
Orthopaedic	Bioactive glass+Ca ₃ (PO ₄) ₂	/	/	[100,101]
Orthopaedic	Bioactive glass+Ca ₅ (PO ₄) ₃ (OH)	/	Ca ₃ (PO ₄) ₂ +Ca ₅ (PO ₄) ₃ (OH)+Ca ₂ SiO ₄	[102]
Orthopaedic	Bioactive glass+CaCO ₃ +Sil icone	/	$\begin{array}{c} Ca_5(PO_4)_3F+Ca_5(PO_4)_3(OH)+C\\ aSiO_3+Ca_2SiO_4+SiO_2 \end{array}$	[103]
Orthopaedic	Ca ₈ H ₂ (PO ₄) ₆ ·5H ₂ O	/	Ca(HPO ₄)·2H ₂ O+Ca ₃ (PO ₄) ₂	[104]
Orthopaedic	Ca(HPO ₄)+Ca(O H) ₂	/	$Ca(HPO_4)+Ca(OH)_2+Ca_5(PO_4)$ 3(OH) or Ca(HPO_4)+Ca_3(PO_4)_2	[105]
Orthopaedic	$Ca_2KNa(PO_4)_2$	/		[106]
Orthopaedic	$CaO+Na_2O+P_2O_5$ +SiO ₂	/	CaO+Na ₂ O+P ₂ O ₅ +SiO ₂	[107,108]
Orthopaedic	$[Ca(PO_3)_2]_n$	/	$[Ca(PO_3)_2]_n$	[109.110]
Orthopaedic	$Ca_3(PO_4)_2$	/	/	[111–115]

Orthopaedic	$Ca_3(PO_4)_2$	/	Ca(HPO ₄)	[116–119]
Orthopaedic	$Ca_3(PO_4)_2$	/	Ca(HPO ₄)+Ca(HPO ₄)·2H ₂ O	[120]
Orthopaedic	$Ca_3(PO_4)_2$	/	$Ca(HPO_4)+Ca(HPO_4)\cdot 2H_2O+$ $Ca_3(PO_4)_2$	[121–123]
Orthopaedic	$Ca_3(PO_4)_2$	/	$Ca(HPO_4) \cdot 2H_2O$	[124]
Orthopaedic	$Ca_3(PO_4)_2$	/	$Ca(HPO_4) \cdot 2H_2O + Ca_3(PO_4)_2$	[125,126]
Orthopaedic	$Ca_3(PO_4)_2$	/	$Ca_3(PO_4)_2$	[115,127– 137]
Orthopaedic	$Ca_3(PO_4)_2$	(A) MgO+SiO ₂	Ca ₃ (PO ₄) ₂ +MgO+SiO ₂	[137]
Orthopaedic	$Ca_3(PO_4)_2$	(A) MgO+SrO	$Ca_3(PO_4)_2 + MgO + SrO$	[135,136]
Orthopaedic	$Ca_3(PO_4)_2$	(A) $S_1O_2 + ZnO$	$Ca_{3}(PO_{4})_{2}+S_{1}O_{2}+Z_{n}O$	[128]
Orthopaedic	$Ca_3(PO_4)_2$ +CaCO	/	Ca ₃ (PO ₄) ₂ +Ca ₅ (PO ₄) ₃ (OH)	[138]
Orthopaedic	$Ca_3(PO_4)_2+Ca_4(PO_4)_2O$	/	Ca4(PO4)2O+Ca5(PO4)3(OH)	[139]
Orthopaedic	$\begin{array}{c} Ca_3(PO_4)_2 + Ca_5(P \\ O_4)_3(OH) \end{array}$	/	Ca(HPO ₄)+Ca ₃ (PO ₄) ₂ +Ca ₅ (PO 4) ₃ (OH)	[140]
Orthopaedic	$Ca_{3}(PO_{4})_{2}+Ca_{5}(PO_{4})_{3}(OH)$	/	/	[141,142]
Orthopaedic	$Ca_3(PO_4)_2+Ca_5(PO_4)_3(OH)$	/	Ca ₃ (PO ₄) ₂ +Ca ₅ (PO ₄) ₃ (OH)	[143–146]
Orthopaedic	$Ca_3(PO_4)_2+Ca_5(PO_4)_3(OH)+CaSO_4$	/	Ca ₅ (PO ₄) ₃ (OH)+CaSO ₄ ·2H ₂ O	[147]
Orthopaedic	$Ca_4(PO_4)_2O$	/	$Ca(HPO_4)+Ca(HPO_4)\cdot 2H_2O+$ $Ca_3(PO_4)_2$	[123]
Orthopaedic	Ca5(PO4)3(OH)	/	/	[115,148– 151]
Orthopaedic	Ca ₅ (PO ₄) ₃ (OH)	/	Ca ₅ (PO ₄) ₃ (OH)	[35,115,12 9,152–157]
Orthopaedic	Ca5(PO4)3(OH)	(I) Bioactive polymer	Bioactive polymer+Ca ₅ (PO ₄) ₃ (OH)	[158]
Orthopaedic	CaSO ₄	ĵ -	$CaSO_4 \cdot 0.5H_2O$	[159–161]
Orthopaedic	$CaSO_4 \cdot 0.5H_2O$	/	$Ca_5(PO_4)_3(OH)$	[162]
Orthopaedic	$CaSO_4 \cdot 0.5H_2O$	/	$CaSO_4 \cdot 2H_2O$	[163]
Orthopaedic	$CaSO_4 \cdot 0.5H_2O$	/	$Ca_{5}(PO_{4})_{3}(OH)+CaSO_{4}\cdot0.5H_{2}$ O	[164]
Orthopaedic	$CaSO_4 \cdot 0.5H_2O$	/	Ca ₅ (PO ₄) ₃ (OH)+CaSO ₄ ·2H ₂ O	[162,163]
Orthopaedic	CaSO ₄ ·0.5H ₂ O	/	$Ca(HPO_4)+Ca_5(PO_4)_3(OH)+Ca \\SO_4\cdot 2H_2O$	[163]
Orthopaedic	$CaSO_4 \cdot 0.5H_2O$	/	$CaCO_3 + CaSO_4 + CaSO_4 \cdot 2H_2O$	[165]
Orthopaedic	CaSO ₄ ·0.5H ₂ O	(A) AgNO ₃	Ag ₃ PO ₄ +Ca ₅ (PO ₄) ₃ (OH)	[166]
Orthopaedic	$CaSO_4 \cdot 0.5H_2O$	(I) $(C_6H_{10}O_2)n$	$(C_6H_{10}O_2)n+Ca_5(PO_4)_3(OH)$	[162]
Orthopaedic	$Mg_3(PO_4)_2$	/	$Mg_3(PO_4)_2$	[167]
Orthopaedic	Mg ₃ (PO ₄) ₂ +Mg ₅ S r(PO ₄) ₄ +Mg ₂ Sr(P O ₄) ₂	/	MgHPO ₄ ·3H ₂ O+Mg(OH) ₂ +Sr CO ₃	[167]
Orthopaedic	Mg ₃ (PO ₄) ₂ +(NH ₄) ₂ HPO ₄	/	NH4MgPO4·6H2O	[168]
Electronic	Al ₂ O ₃	/	Al ₂ O ₃	[25,27,169]

Electronic	BaTiO ₃	/	BaTiO ₃	[27,170,17 1]
Electronic	Si ₃ N ₄	/	Si ₃ N ₄	[25]
Dental	Al_2O_3	(I) Dental glass	Al ₂ O ₃ +Dental glass	[172]
Dental	Dental porcelain	/	Dental porcelain	[173,174]
Dental	Dental porcelain	(A) Al_2O_3	Al ₂ O ₃ +Dental porcelain	[175]
Optical	SiO ₂	/	SiO ₂	[176]

Materials investigated for structural applications include oxides (Al₂O₃, TiO₂, ZrO₂), carbides (WC, TiC, SiC), and nitrides (AlN, Si₃N₄). For orthopaedic applications, materials in the calcium phosphate family, such as hydroxyapatite (HA, Ca₅(PO4)₃(OH)) and tricalcium phosphate (TCP, Ca₃(PO₄)₂), are the most common choice due to their compositional similarity to human bones and thus excellent biocompatibility [177]. Other calcium phosphate materials include calcium polyphosphate (CPP, [Ca(PO₃)₂]_n), tetracalcium phosphate (TTCP, Ca₄(PO₄)₂O), and dicalcium phosphate (DCP, CaHPO₄). Alumina is the first material studied by the inventors of binder jetting [6,7]. In the past decades, it is also one of the mostly studied materials in different applications including structural, electronic, and dental areas.

In addition to the pure compound materials, a large portion of studies used composites to enhance various properties. For example, 12.5 vol.% zirconia was mixed with 40 vol.% alumina slurry to attain optimum strength for the printed and sintered parts [72]. Although composite materials can offer improved properties, they sometimes sacrifice other properties, such as biocompatibility [37].

2.3 Feedstock powder

Feedstock powder and its deposition method determine various physical and chemical processes during fabrication, such as powder spreading, binder-powder interaction, and sintering

densification [178]. This section discusses important feedstock powder parameters and deposition methods.

2.3.1 Powder flowability

Flowability, the ability of a powder to freely flow, is crucial for uniform powder spreading [179] and thus homogeneous green and sintered part structure [48]. Flowability [103,170] is also called depositability [180], mobility [59], pourability [42], and spreadability [69,75,112,172].

Flowability can be assessed using different metrics, including flow factor (f_c) [111,112], Hausner Ratio (HR) [11, 43, 79–81], Carr Index (CI) [151], and flow rate [149,153,155], among others. Flow factor is defined by the following equation,

$$ff_c = \frac{\sigma_1}{\sigma_c} \tag{1}$$

where σ_1 is the consolidation stress and σ_c is the compression strength, both of which can be measured with a ring shear tester [111,112].

Hausner Ratio [11, 43, 79-81] is defined by the following equation,

$$HR = \frac{\rho_t}{\rho_a} \tag{2}$$

where ρ_t and ρ_a are the tap density (the density of powder after a certain number of tapping cycles [64,98,181,182]) and apparent density (the density of freely settled powder), respectively. *HR* value is always larger than or equal to one. A smaller *HR* value, i.e., closer to one, indicates better flowability. Carr Index [151] is similar to Hausner Ratio. A smaller index value indicates better flowability. Its definition and relationship with Hausner Ratio are shown as follows,

$$CI = 100 \left(1 - \frac{\rho_a}{\rho_t} \right) \tag{3}$$

$$CI = 100 \left(1 - \frac{1}{HR} \right) \tag{4}$$

Flow rate is usually measured by a Hall flowmeter [149,153,155], in which a defined volume of powder passes through a small opening of a metal funnel. The flow rate can be represented by the total time required for a certain amount of powder to pass [92,153] or the mass of the powder passing through in a unit time [149,155].

2.3.2 Powder sinterability

Sintering is "a thermal treatment for bonding particles into a coherent, predominantly solid structure via mass transport events that often occur on the atomic scale" [183]. It is affected by powder properties and packing state. Powder sinterability is used to compare the sintering performance of different powders under a similar packing condition, and it is important for the selection of the optimal feedstock material before printing. Sintered bulk density is commonly used to describe powder sinterability since high sinterability leads to high sintered density under same conditions [52]. Volumetric shrinkage is another sinterability metric because high sinterability leads to large volumetric shrinkage under same conditions [184]. Powder sinterability can also be assessed by the densification ratio,

$$\varphi = \frac{\rho_s - \rho_g}{\rho_{th} - \rho_g} \tag{5}$$

where φ is the densification ratio and ρ_g , ρ_s , and ρ_{th} are the green density, sintered density, and theoretical density, respectively [185]. Theoretical density is calculated based on corresponding crystal structure [186].

2.3.3 Powder particle shape

Various particle shapes have been reported in the literature, as shown in Figure 4. Spherical particles usually have better flowability. For example, the flow times of spherical and irregular calcium alkaline phosphate powders (45–90 μ m, 50 g) are 121 and 166 s under the same conditions, respectively [106], which indicates the better flowability of spherical particles.

There are no reported studies about effects of powder particle shape in ceramic binder jetting. However, effects of particle shape on powder bed density were reported from other fields, such as geology [187]. Compared with non-spherical powders, sand powder with spherical particle shape could lead to a higher value of powder bed density [187].



Figure 4. Different ceramic particle shapes used in binder jetting: (a) spherical, (b) rounded [179], (c) angular, (d) polygonal [172], (e) irregular [102], and (f) aggregate.

2.3.4 Powder particle size

Reported particle size of feedstock powder used in ceramic binder jetting ranges from 0.3 (the minimum particle size in [173]) to 355 μ m (the maximum particle size in [167]). Particle size of feedstock powder in binder jetting affects powder flowing and sintering behaviors.

Coarse powder usually has good flowability while fine powder often has good sinterability [111]. When particle size is smaller than a certain value, the interparticle cohesion becomes more dominant than the inertia [188]. Therefore, fine particles tend to agglomerate, usually resulting in bimodal pore size distribution in the spread powder layer [147,165,189] and a low powder bed density. Particle size also plays a vital role in powder sinterability. The specific surface area of fine powder is larger than that of coarse powder, leading to a larger sintering driving force and consequently a higher densification after sintering [98]. It is noted that larger particles are desirable for better flowability while smaller particles are preferred for better sinterability. These two requirements on particle size are contradictory.

Sun et al. reported that the poor flowability of a mixed glass-ceramic powder of a relatively small particle size (< 25 μ m) caused powder adhesion onto the roller and thus the movement of the printed layer when spreading a new layer, eventually leading to a significant misalignment between printed layers [98]. In Zocca et al.'s study [97], lithium alumino-silicate glass powder with a median diameter of 75 μ m (fine powder) had an *HR* value of 1.29 and the powder with a median diameter of 223 μ m (coarse powder) had an *HR* value of 1.11, indicating the better flowability for the coarse powder.

2.3.5 Powder deposition methods

In binder jetting, the powder can be deposited with various methods. Figure 5 shows four powder deposition methods, where *h* is desired layer thickness and h_d is pre-deposited thickness. A parameter called compaction ratio, which is the ratio of h_d to *h*, is used to quantify the compaction level, and a value of two is a common choice [75,170]. The effect of an amount increase of the pre-deposited powder, which induces a higher compaction ratio, has been

experimentally studied [57]. It shows that green parts made with more pre-deposited powder have a higher bulk density.

Among all deposition methods, doctor blade spreading is the simplest, with little powder compaction occurring. Other three methods use a roller instead of a blade, and the roller can be fixed [151], forward-rotating [75], or counter-rotating [151]. The counter-rotating roller is the most commonly used. The traversing movement of the roller deposits and compacts the powder, while the rotation smooths the powder bed. Compared with counter-rotating roller, fixed roller and forward-rotating roller have not drawn much interest because of the higher surface roughness of the spread powder bed [151] and powder bed disturbance by the powder adhering onto the roller surface [75], respectively.

In a numerical study, counter-rotating roller outperforms the doctor blade in achieving high powder bed density and low surface roughness [190]. Combination of different methods, e.g. forward-rotating roller and doctor blade [75] has been reported to increase powder bed density. Parameters of each deposition method can also be tuned to improve powder bed density. For example, Shanjani et al. [191] introduced an analytical model and calculated the powder bed density formed by a counter-rotating roller system. It was found that larger roller diameter leads to denser powder bed. It should be noted that there is no research directly investigating the effect of different powder deposition methods on printed part properties.



Figure 5. Various powder deposition methods: (a) doctor blade, (b) fixed roller, (c) forward-rotating roller, and (d) counter-rotating roller. h is desired layer thickness and h_d is pre-deposited thickness.

Though sub-micron and nanometer powders can be easily sintered, it is not easy to spread them due to their high surface energy and thus agglomeration issue [192]. One possible method to address this problem is vibration-assisted spreading. For example, Sachs employed three different methods to vibrate different parts during the printing process, including the build platform, the powder bed surface, and the scraper [20], to spread ceramic powders with a particle size of about 20 μ m. A vibrating counter-rotating roller [192], though not used for ceramic materials, is another potential method to break down agglomerates in fine powder with a particle size as small as 100 nm.

2.4 Binder

2.4.1 Binder material and concentration

Table 3 lists binder materials used in reported studies. Organic materials, including polymers (e.g., polyvinyl alcohol) and carbohydrates (e.g., dextrin), are the most common choice for the binder. They have versatility to almost any powder and capability of thermal decomposition with little residue. Phosphoric acid is another common choice, especially for the scaffold parts from the calcium phosphate family. Colloidal silica is used in some cases where the binder is to be incorporated into the final component [18].

Binder material	Reference
Carbohydrates (dextrin, maltodextrin, starch, etc.)	[38,54,68,71,80,82,85,86,88,96,99,102,107,108,1 15,144,145,148,149,155,156,158,162,163,166,16 7]
Phosphoric acid	[85,101,104,111–113,116,118– 123,125,126,133,138,140,141,168]
Polymers (polyvinyl alcohol, polyethylene glycol, polyvinylpyrrolidone, etc.)	[23– 25,27,38,57,58,64,65,67,69,73,80,94,95,109,110, 114,134,142,143,153,157]
Colloidal silica	[6,7,18,20,72]
Acrylic acid	[23,27,79,146]

Table 3. Binder materials in reported studies on ceramic binder jetting

Binder concentration quantifies the amount of the adhesive material in the binder solution. For a powder-binder system in which chemical reaction happens between them, binder concentration could affect the green density. In Gbureck et al.'s study [120], tricalcium phosphate (Ca₃(PO₄)₂) part was printed with various phosphoric acid binder concentrations (5, 10, 20, and 30 wt.%) and the green density was measured. Results show that the increase of binder concentration led to an improvement of the binder-powder reaction between tricalcium phosphate and the phosphoric acid forming dicalcium hydrogen phosphate (CaHPO₄·2H₂O) and dicalcium pyrophosphate (Ca₂P₂O₇) as cement materials for the part [101,126,193]. They observed a downward trend of the green porosity. It should be noted that for a powder-binder system in which binder only bonds the particles together and does not involve any chemical reaction, binder concentration does not significantly affect the part density as the binder will be burnt off during debinding.

2.4.2 Binder application methods

Figure 6 illustrates two different methods of binder application. The first method, called binder jetting in-place, is to add the binder material to the printing solution and then jet the binder solution onto the powder bed [6,111,160,173]. The second is to pre-mix powder and binder by either a dry [109,110,163] or wet [85,142,158] approach and use the powder-binder mixture as the feedstock material. For dry mixing, the ceramic and binder materials are pre-mixed using a grinder, ball mill, or pestle and mortar. For wet mixing, the binder material is dissolved in a solvent (e.g., water), and the ceramic material is added to obtain mixture slurry. Then the slurry is spray-dried [35,68,100,101,115,127,129,138,142,146,148,149,152–155,158] or freeze-dried [63,70,71,80,85,87,88] and sieved. For both dry and wet mixing, the binder within the powder feedstock joins the ceramic particles wherever the printing liquid (usually a water-based solution) is jetted.



Figure 6. Two binder application methods: (a) binder jetting in-place and (b) powder-binder premixing

Figure 7 shows shares of each of the binder application methods in the reported studies. Binder jetting in-place method is less complicated than the other two due to fewer feedstock preparation steps, and has the most shares in the reported studies. The powder-binder pre-mixing method allows for the use of a low-viscosity printing liquid because the binder is supplied in the powder

feedstock, which decreases the possibility of nozzle clogging. Wet pre-mixing can be more advantageous than dry pre-mixing because the binder is more evenly mixed with the ceramic material.



Figure 7. Shares of different binder application methods in reported studies

2.5 Process parameters

2.5.1 Layer thickness

Layer thickness directly affects the time needed to print a part and the surface roughness of the part. A smaller layer thickness can lead to smoother surface but a longer printing time, and vice versa [101]. The particle size is an important parameter to consider for selecting layer thickness. For binder jetting, there are no universally accepted rules for selecting layer thickness. Rules reported in the literature include at least greater than the largest particle size [75,165,194], twice the particle size [195], and at least three times the particle size [112,180].

Some studies have been reported about the effects of layer thickness on various material properties. Meier et al. employed a discrete element method (DEM) model that considered particle-to-particle and particle-to-wall interactions involving frictional contact, rolling resistance, and

adhesive forces [196]. It was found that the powder bed density increased with increasing layer thickness. Some other studies found different effects of layer thickness on printed part density. For example, Shanjani et al. introduced an analytical model to predict the powder bed density under various layer thickness values [191]. The model was based on the mechanics of plastic strains of a volume-compressible continuum [197]. They found that an increase of layer thickness led to a decrease of the powder bed density.

2.5.2 Binder saturation

Binder saturation, S_b , quantifies the amount of the binder solution applied during the printing process. It is defined as the percentage of air space (in the powder bed) that is filled with binder solution, given by the following equations,

$$S_b = \frac{V_b}{V_a} \tag{6}$$

$$V_a = \left(1 - \rho_p'\right) \times V_e \tag{7}$$

where V_b is the volume of the binder applied to a pre-defined envelope, V_a is the air space volume of the powder bed within the same envelope before jetting binder, ρ'_p is the relative packing density of the powder bed, and V_e the volume of the pre-defined envelope. It should be noted that the value of binder saturation may exceed one hundred percent because the binder in the previous layers can dry out and therefore more binder than the presumed total free volume can be jetted to the powder [97].

A low binder saturation level leads to limited contact between powder and binder, and results in fragile green parts. In contrast, a high binder saturation level causes the binder to spread out of the selected area, which is called bleeding, compromising the dimensional accuracy and surface smoothness of printed parts [179]. Effect of binder saturation on the sintered density was studied in Sun et al.'s study. A continuously decreasing trend of the sintered bulk density from 66.5% to 55.9% on titanium silicon carbide was reported with the increase in the saturation level of a commercial binder from 10% to 30% [90], which may be due to the increased surface roughness resulted from binder bleeding and thus larger apparent porosity. Melcher et al. found that the bulk density of sintered alumina parts from granulated powder (granule size of <150 μ m) increased from 56% to 67% with the increase of the binder saturation of a water-based commercial binder from 0.14 g/cm³ to 0.35 g/cm³ [63]. It might be due to stronger particle bonding and thus denser green part after printing with the higher binder saturation.

3 Terminologies, Measurement Methods, and Achieved Values for Density and Porosity

3.1 Definition of various terminologies

Density and porosity are the most commonly assessed material properties in ceramic binder jetting. There are many terminologies related to density and porosity, dependent on what packing state the powder is in, what processing stage the part has gone through, and what pores are included in the calculation. The definitions are not always made clear in published reports. These terminologies and their definitions are summarized in Table 4.

Category	Terminology	Definition	Reference
	Apparent density	Density of freely settled powder	[97,98,111,147]
Powder density in different packing	Tap density	Density of powder after standard tapping process	[97,98,147,155]
states	Powder bed (packing) density	Density of powder that is spread on the build platform	[23– 25,63,80,94,99,147,172 ,174]
	Green density	Density after printing	[23,25,80,88,99,110,12 0,143,155]
Part density after different process	Brown density	Density after debinding	[198]
stages	Sintered density	Density after sintering	[63,70,71,78,85,87,92,9 6,97,102,103,108,135– 138,144,146,174]
Part density and porosity when different pores are included	Apparent solid density	Mass per unit apparent solid volume (total volume of solid material and closed pores)	[68,149,169,170]
	Bulk density	Mass per unit bulk volume (total volume of the solid material, open pores, and closed pores)	[98,130,135,136,158,16 2]
	Apparent porosity	Volume percentage of open pores in the bulk material	[63,84,97,98,105,110,1 20,124,125,134,143,14 9,155,157,161,164,167, 175]
	Closed porosity	Volume percentage of closed pores in the bulk material	[128,130,135,136]
	Bulk porosity	The sum of apparent and closed porosity	[83,87,160,96,100,103, 117–119,127,144]

Table 4. Terminologies for density and porosity

3.1.1 Powder density in different packing states

There are three terminologies to define the powder density depending on different packing states: apparent density, tap density, and powder bed density. Apparent density is the density of

freely settled powder [199–202]. Tap density is the density of a powder that has been tapped, to settle contents, in a container under specified conditions [181,182]. Powder bed density is the density of the powder that is spread on the build platform. Unlike apparent density and tap density that can be measured by following standards [181,182,199–202], powder bed density can be measured by various methods. A common method is to spread a number (e.g., 50) of layers of powder and then measure the bulk volume and the mass of the powder [63,75,170]. Another method is to print multiple cylindrical cups and then measure the cup size and weight change after depowdering [203]. For a given powder, powder bed density is usually between those of apparent and tap densities.

3.1.2 Part density after different process stages

Green density, brown density, and sintered density are the part densities that just after printing, debinding, and sintering, respectively. Green and sintered densities have been vastly investigated while brown density has been rarely reported for the friability of the debound part and thus the difficulty of density measurement. Other densities can be obtained if additional post-processing steps are applied. For example, infiltrated density can be measured if infiltration is performed [69,82,85,88,158,162] and pressed density can be obtained if isostatic pressing is implemented [64,91,94].

Relative density (ρ') is the ratio of absolute density (ρ) to theoretical density (ρ_{th}):

$$\rho' = \frac{\rho}{\rho_{th}} \tag{8}$$

where theoretical density (ρ_{th}) is calculated based on crystal structure and unit cell dimensions. In this paper, relative densities are used since they can be readily compared across various materials.

3.1.3 Part density and porosity including different pores

For a part with both open and closed pores, its volume measurement is complicated. Figure 8 schematically shows these pores in a part. Open pores are permeable from a surface while closed pores are not. The bulk volume of the part includes volumes of solid as well as both open and closed pores, while the apparent solid volume includes volumes of solid as well as closed pores. The only difference between these two volumes is that the bulk volume includes open pores but the apparent solid volume does not. Accordingly, the bulk density is the mass per unit bulk volume while the apparent solid density is the mass per unit apparent solid volume. Similarly, apparent porosity [204], closed porosity, and bulk porosity are the fractions of the volumes of open pores, closed pores, and both of them, respectively, in the bulk volume. The bulk porosity (π_a), and closed porosity (π_c) can be estimated from the relative apparent solid density (ρ'_{as}) and relative bulk density (ρ'_b) based on the following equations

$$\pi_b = 1 - \rho_b' \tag{9}$$

$$\pi_a = 1 - \frac{\rho_b'}{\rho_{as}'} \tag{10}$$

$$\pi_{c} = \frac{\rho_{b}'(1 - \rho_{as}')}{\rho_{as}'}$$
(11)



Figure 8. Open and closed pores in a cross section of a part

The methods for part density and porosity measurement are listed in Table 5. The geometry method works for a part with a simple shape (e.g., cuboid and cylinder). The simple Archimedes' method measures two kinds of sample masses: dry mass and immersed mass in water (to calculate the immersed volume). Usually, water is infiltrated into the open pores. The immersed volume is the apparent solid volume. Therefore, the simple Archimedes' method only measures the apparent solid density. Compared with that, a full water absorption into the open pores is induced in the modified Archimedes' method by boiling [204,205] or vacuuming [204,206,207], after which immersed mass and soaked mass (in addition to dry mass) are measured to evaluate the open pore volume [204]. Therefore, this method is capable to simultaneously measure apparent solid and bulk densities.

In the mercury intrusion method, mercury as a non-wetting liquid is used, and does not enter open pores unless under pressure. The volume of the pores is determined from the mercury volume intruded at each pressure increment [208]. Gas pycnometer has a similar principle with that of simple Archimedes' method but with a displacement medium of gas (e.g., helium), which is suitable for physically or chemically sensitive materials [209] or for small pores. Computed tomography is a non-destructive method which uses X-ray and mathematical algorithm to generate cross section images of the part [210]. From these cross section images, the volume can be determined. Image analysis method uses 2D images of the sample cross section and quantifies the porosity [170].

Method	Bulk density	Apparent solid density	Bulk porosity	Apparent porosity	Closed porosity	References
Geometry	Yes	No	Yes	No	No	[95,113,130,135, 136,139,146,158 ,162]
Simple Archimedes'	No	Yes	No	No	No	[68,95,98,168– 170]
Modified Archimedes'	Yes	Yes	Yes	Yes	Yes	[38,69,83,86– 88,96,97,102,10 3,109,110,128,1 32,148,149,155, 157,171]
Mercury intrusion	Yes	Yes	Yes	Yes	Yes	[23,63,70,71,108 ,143,156,163]
Gas pycnometer	No	Yes	No	No	No	[82,105,139,146]
Computed tomography	Yes	Yes	Yes	Yes	Yes	[141,144,160,16 1]
Image analysis	Yes	Yes	Yes	Yes	Yes	[170]

Table 5. Capabilities of different density and porosity measurement methods

3.2 Achieved part density in reported studies

As relative densities and porosities can be readily compared across different materials, all absolute values have been transformed to relative values based on Equation (8). Theoretical density values are shown in Table 6. It should be noted that only materials whose density values were reported in absolute values are shown in this table.

Material	Theoretical density (g/cm ³)	Reference
Al ₂ O ₃	3.97	[211]
AlN	3.25	[211]
BaTiO ₃	6.02	[211]
CaSO ₄	2.32	[212]
SiC	3.22	[211]
TiC	4.91	[211]
TiAl ₃	3.40	[213]

Table 6. Theoretical density values of different materials

Figures 9 and 10 show the achieved relative densities and porosities of the green and sintered parts. Different densities and porosities are interpreted by the reported terminologies and measurement methods in the corresponding papers. Special treatment techniques are used to categorize all values. Specific data values of Figures 9 and 10 and the corresponding references are shown in Appendixes A and B, respectively. It should be noted that besides these 83 density and porosity values from 65 publications, there are some unclear values because of the lack of specific description of the measurement method, or non-standard metrics (e.g., porosity derived from apparent solid density).



Figure 9. Achieved values of relative green densities and porosities by various special treatment techniques (N stands for no special treatment, G for powder granulation, L for using slurry feedstock, and S for mixing powders of different sizes) [23–25,27,38,54,55,57,58,63,64,67,68,73,74,81,82,86–89,92,110,156,161]

Green bulk density determines the sintered bulk density and thus other material properties. A slurry feedstock usually leads to a green bulk density of higher than 50%. Using a slurry with a 35 vol.% alumina (0.5 μ m), a green bulk density of 67% was achieved in the study by Grau et al. [24], which is the highest in the literature. Mixing powders of different size is another commonly used method to improve green density. Kunchala et al. [58] introduced alumina nanoparticles (less than 50 nm) to the binder and printed alumina powder feedstock with an average size of 40 μ m. A green bulk density of 65.7% was obtained when the nanoparticle concentration in the binder was 15 wt.%.



Figure 10. Achieved values of relative sintered densities and porosities by various special treatment techniques (N stands for no special treatment, C for chemical reaction, G for powder granulation, I for infiltration, L for using slurry feedstock, M for mixing different materials, P for pressing, and S for mixing powders of different sizes) [24,27,38,53,55,63–65,68–71,77,83,85,86,88–90,92,94–98,100,102,103,108–110,127,128,130,132,135,136,144,146,149,155–158,169,170,174]

Sintered bulk density directly determines other material properties. Therefore, it is mostly reported in the literature. A high sintered bulk density (>90%) was achieved in some studies by applying slurry feedstock [24,27,38,94], infiltration [86], or isostatic pressing [64,90,94]. Slurry feedstock is among the most effective techniques. In Cima et al's study [27], parts printed with alumina slurry achieved a green bulk density of 58% and a sintered bulk density of 99.9%. Combinations of special treatments can lead to high sintered bulk density as well. Specifically,

Kernan et al. [94]_studied the printing of tungsten carbide with an average particle size of 0.8 μ m, in which a combination of mixing different materials (tungsten carbide and cobalt), slurry feedstock (25 vol.% water-based slurry) and hot isostatic pressing (5.5 MPa during sintering) was applied and the achieved sintered bulk density was 100%. A bulk density of 99.2% was achieved by using an alumina slurry feedstock (0.5 μ m, 34 vol.%) in combination with applying sintering additives in Zocca et al's study [38]. Bulk density of the printed alumina part increased from 34% to 61% by warm isostatic pressing in Yoo et al's study [64], leading to a sintered bulk density of 99.2% (with the additional help of sintering additive and powder granulation).

4 Material Preparation Techniques for Density Improvement

4.1 Powder granulation

The contradiction between the flowability and the sinterability of feedstock powder is among the main challenges in the field of ceramic binder jetting. Generally, the particle size used in binder jetting is in the range of 10–100 μ m to ensure a good flowability and avoid defects in powder bed layers. At the same time, the particle size should be less than 1 μ m to ensure a high sinterability and thus a dense part after sintering [64]. However, the flowability of fine powder (less than 1 μ m) is usually not high enough to form a uniform and smooth layer of powder bed due to its high interparticle cohesion [151,214]. Fine powder spreading usually creates pores and cracks on powder bed, which are consequently inherited by the green and sintered parts. Therefore, increasing the flowability of fine powder is highly beneficial to fabricate dense parts with a minimal number of pores.

One common technique to increase powder flowability is granulation [215]. Granulated powder can significantly improve the flowability of feedstock powder. The fine raw particles ensures the required sinterability and the coarse resultant granules ensures the required flowability.

Its principle is schematized in Figure 11. Basically, raw (fine) particles are bonded by a binder to form larger aggregates (or granules). Steps for granulation include mixing, drying, comminuting, and sieving, as shown in Figure 12 and listed in Table 7. A ball mill, grinder, or pestle and mortar can be used to mix raw particles with a binder solution to form a slurry, which can be dried by a spray dryer, freeze dryer, or oven. Comminuting step is not needed for the powder from a spray dryer because the size is determined by sprayed droplets. For the cakes dried by a freeze dryer or an oven, comminuting step is needed to decrease the size by ball milling, grinding, or manual crushing. Then a sieving step screens the powder into several groups with different size ranges. Generally, the manual granulation method, comprising of manual mixing in a crucible, oven drying, manual crushing in a crucible, and finally sieving, is simple, although the granule shape is irregular. Commercial granulation machines based on spray drying technologies, though costly, can produce spherical and large-batch granulated powder.



Figure 11. Principle of granulation



Figure 12. Basic granulation steps

Granulation process	Reference
Dry mixed, dry ball milled	[86]
Dry mixed, sieved	[80]
Wet mixed, (wet ball milled), spray dried	[68,100,101,127,128,142,146,148,149,153,155,158]
Wet mixed, (wet ball milled), oven dried, dry ball milled	[78,130,135–137]
Wet mixed, (wet ball milled), freeze dried, dry ball milled, sieved	[63,70,71,85,87,88,95]
Commercial spray-dried powder	[35,129,152,154]

Table 7. Granulation process practiced in reported studies

Chumnanklang et al. [149] investigated the effect of binder fraction on the flowability of spray-dried hydroxyapatite powder. The powder flow rate increased from 14.6 to 21.2 g/s when binder fraction decreased from 48.3 to 15.7 vol.%. Gildenhaar et al. prepared spray-dried calcium alkaline phosphate powder with various raw particle sizes [106]. They found that printed parts made with granules from fine raw powder ($3.1 \mu m$) had a higher compressive strength (2.5 MPa) than those from coarse raw powder ($7.5 \mu m$, 1.80 MPa).

With the same raw powder, different granulation technologies can result in different material properties. Suwanprateeb et al. [155] studied two granulation technologies, manual granulation with mortar and pestle, and spray drying. After printing and sintering at 1300 °C, samples made with manually granulated hydroxyapatite powder showed ~20% higher bulk density and approximately two times higher flexural modulus and strength than those made from spray-dried powder. One of the reasons is the larger number of inter-granule pores from the spray-dried powder than the manually granulated powder, leading to a lower green density. Another possible reason is the large pores within the spray-dried granules, which are formed due to the solvent evaporation during the drying process [216].

These two issues of spray-dried granules can potentially be resolved by another granulation technique called spray freeze drying [217–219]. The granules from spray freeze drying can be soft, making the particles loosely bonded and easily breakable. The granules can be crushed by externally exerted forces such as the roller compaction force. Therefore, the granules retain a microscale size before spreading and facture to a nanoscale size after compaction. The other feature of the granules from spray freeze drying is its homogeneous structure. No large pores are formed inside a granule after drying as the sprayed droplet is frozen first and the drying process is solvent sublimation instead of evaporation [217].

4.2 Mixing powders of different sizes

Particle size distribution plays a key role in powder packing, which can be either monomodal or multimodal. Since fine particles can fill voids between coarse particles, powder with a multimodal size distribution has less interparticle voids. From this point of view, mixing powders of different sizes can be an effective technique to increase powder bed density and consequently green density of printed parts [65]. Moreover, this technique could also reduce sintered shrinkage as there are less voids in the green part. This is beneficial for improving the dimensional accuracy after sintering.

Table 8 lists studies on mixing powders of different sizes for feedstock preparation. For example, Sun et al. studied the effects of powder mixing on powder flowability, part sintered density, and part bending strength [98]. Glass-ceramic powders with three size ranges were mixed in various fractions. The mixture with 60 wt.% 45–100 μ m and 40 wt.% 0–25 μ m particles achieved the highest sintered bulk density of 1.6 g/cm³ and bending strength of 13.8 MPa. In Spath et al.'s experiment, parts made from mixed hydroxyapatite powder had a higher compressive strength than those from the two constituent powders [148].

Instead of finding the optimal ratio by trial and error, a promising direction is model-guided selection of particle size and mixing ratio of constituent powders. In the study by Du et al. [65], packing densities with various mixing ratios were predicted using the linear packing model [220]. Optimal mixing ratios to achieve the highest packing density were selected for both binary and ternary mixtures. The printed parts from optimal ternary ratios achieved the largest sintered bulk density. Although the improvement is small due to the decreased flowability, the study showed that the linear packing model has a potential to guide the selection of the particle size and mixing ratio.

Material	Size (µm)	Mixing Ratio	Reference
Al ₂ O ₃	2, 10, 70	17.6:21.1:61.3	[65]
Al ₂ O ₃	30, 45, 53	1:1:1	[169]
Al ₂ O ₃	20, 3.4, 0.4	63:27:10	[67]
Al_2O_3 and Al	0.6, 3	25:100, 30:100, 35:100, 40:100, 50:100	[54]
$Ca_3(PO_4)_2$ and bio-glass	8, 38	1:3, 1:1	[100]
Ca ₅ (PO ₄) ₃ (OH)	38, 125	15:85, 25:75, 35:65	[148]
Ca ₅ (PO ₄) ₃ (OH)	4, 50	15:85, 25:75, 40:60, 60:40, 75:25, 100:0	[156]
Glass-ceramic compound	0–25, 25– 45, 45– 100	0:100, 10:90, 20:80, 30:70, 40:60, 100:0	[98]

Table 8. Reported studies on mixing powders of different sizes

Another method to mix powders of different sizes is to introduce nanoparticles of the same material during printing [58,95]. These nanoparticles can enhance the sintering performance due to their high activity. For example, Zhang et al. demonstrated the feasibility of using 10 wt.%

zirconia nanoparticle suspension as the printing liquid while printing zirconia parts [95]. The printed and sintered bulk density increased from 75.2% to 86.8% as the binder saturation increased from 50% to 120%.

4.3 Slurry feedstock

Structural ceramic parts usually require fine-grained powder (submicron) and high green density (>50%) to achieve a full density after sintering [24]. However, it is difficult to meet these requirements simultaneously using dry powder feedstock. Due to its low packing density and agglomeration issues, dry fine powder feedstock generally results in parts with low green density and defects.

Slurry feedstock was used to meet both requirements of fine powder and high green density [24–31,38,77,81,94]. In this technique, a layer of ceramic slurry is deposited and then dried to form the powder bed [26]. Two slurry deposition methods have been reported in the literature. The first one is nozzle jetting [23–25,27,94,176] that was studied by the inventors of binder jetting. In this case, a thin slurry layer is deposited by a single nozzle scanning over a porous substrate [25]. The second deposition method is doctor blade spreading that was studied by Zocca et al. [38,81] and Lima et al. [77]. In both deposition methods, the layer top is dried by a heating lamp, while the bottom was dried due to capillary forces from the porous substrate or previously deposited and dried layers [24,221]. Delicate temperature and ventilation control is needed to achieve crack-free powder bed layers.

Compared with the dry powder deposition method, the slurry-based method can handle submicron particle size (down to 0.5 μ m). It can achieve high powder bed density (up to 53% in Kernan et al.'s study [94]) and high green part bulk density (up to 67% in Grau et al.'s study [24]).

Different materials (i.e., alumina [24,25,27,38,77], silicon nitride [24,25], silicon carbide [81], silica [176], and tungsten carbide with cobalt [94]) have been investigated with this method. In Zocca et al's study [39], printed parts from an alumina slurry (solid loading of 34 vol.% and average particle size of 0.5 μ m) achieved near theoretical density (i.e., bulk density of 99.2%) after sintering on 1600 °C with the help of sintering additives.

Maximum layer thickness, i.e. critical saturation thickness (CST) [30], exists for slurry feedstock technique. A layer thickness larger than the CST leads to cracking or warping of the powder bed due to capillary pressure of solvent evaporation [24,26,30]. Furthermore, thick parts are prone to crack and difficult to be fully dried [49].

It should be noted that slurry feedstock has been extensively used in other ceramic AM technologies such as vat photopolymerization and material extrusion. For example, Hu et al. prepared a 60 vol.% alumina slurry (average size of 138 nm) [222] for vat photopolymerization. After printing and sintering at 1650 °C for 2 h, the part achieved a bulk density of 3.96 g/cm³ (99.7% in relative). Mamatha et al. [223] used a slurry with a solid loading of 64.75 wt.% of alumina powder (average size of 331 nm) in material extrusion. After printing and sintering at 1650 °C for 1 h, the part achieved a bulk density of 97.7%. Therefore, application of slurry feedstock in binder jetting can be advanced by learning from other AM technologies.

4.4 Mixing different materials

Mixing different materials is a common technique to increase printed and sintered density [77,92,96–98,100,102,105,108,120,123,124,128,138,139,141,143,144,146,174]. The mechanisms for density enhancement could be boosting the mass transport [38,74,128,135–137] or enabling liquid phase sintering [224]. Yoo et al. doped alumina (mean particle size of 0.8 μm)

with MgO and granulated both undoped and doped powders to a granule size range of 70–150 μ m [64]. The printed and sintered parts from doped alumina achieved a higher relative sintered bulk density (99.2%) and flexural strength (324 MPa) than the undoped powder (97.8% and 231.6 MPa) after printing, isostatic pressing, and sintering.

Although this mixing technique could be effective, other material properties (e.g., biocompatibility) may be impaired. The limited choice of additive materials is another of the drawbacks of this technique. Moreover, the additive particles may not be homogenously distributed within the ceramic powder, leading to structure heterogeneity (i.e., local variation) in the sintered part [52,225]. In addition, the selection of additives has remained largely empirical as the mechanism of the sintering additive has not been fully understood [225]. Therefore, this technique only works for specific materials.

5 Post-processing Techniques for Density Improvement

5.1 Sintering

Sintering is the process by which a powder compact is transformed to a dense body by heating [226]. It's the most commonly used method to increase the density. Sometimes it is considered an essential part of the process and not considered a separate post-processing method. The macroscopic driving force in sintering is the reduction in surface energy. The densification during sintering happens by the elimination of solid/vapor interfaces [226]. However, sintering is not required in some cases, for example, when porous parts are preferred or other post-processing techniques are applied.

One of the important sintering parameters is the sintering temperature. Sintering temperature significantly affects the material properties as it determines the mass transport of the ceramic

particles. Typically, a higher sintering temperature can facilitate mass transport and subsequently increase part density and decrease porosity [53]. Another parameter significantly affects the sintered density is the green density. To present the effect of green density on sintered density, the bulk densities before and after sintering are shown in Figure 13. Powder bed density is also included as it is close to green density. As some parts were isostatically pressed before sintering [64], their pressed green densities are presented instead of printed green densities. Specific data values of Figure 13 and the corresponding references are shown in Appendix C. Generally, high green density leads to high sintered density. A high green density (i.e., >50%) is required to achieve a high sintered density (i.e., >90%).



Figure 13. Bulk densities before and after sintering [24,27,94,156,38,55,63–65,68,83,89]

Shrinkage is a critical index to determine the density change by sintering. For parts with the same green density, a larger shrinkage leads to a higher sintered density. A primary factor that affects shrinkage is sinterability, which is directly related to particle size [227]. For example, Du et al. [65] used alumina powder with an average particle size of 10 µm and achieved a powder bed density of 57.1%. Using 0.5 µm alumina powder, Zocca et al. [38] achieved a green density of

57.2%. As there is no significant density change for the powder bed after printing, it can be assumed that green parts from Du et al.'s and Zocca et al.'s studies have approximately the same green densities. After sintering at 1600 °C, parts made from coarse powder (10 μ m alumina in Du et al.'s study) achieved only 0.7% volumetric shrinkage, leading to a bulk density of 57.7%. However, parts from fine alumina powder (0.5 μ m in Zocca et al.'s study) achieved a volumetric shrinkage of 73.4% and bulk density of 99.2%.

It should be noted that sintering could also result in shape distortion due to gravity effect, nonuniform temperature gradient, liquid phase formation due to high sintering temperature, etc. [228]. For example, Grant et al. [89] studied the distortion of printed titanium dioxide cantilever beams at different sintering temperatures up to 1420 °C. It was found that most of the distortion happened above 1180 °C. To mitigate the distortion, they infiltrated the printed parts with an aqueous precursor solution and then heated the parts at 300 °C to decompose the precursor to form titanium dioxide nanoparticles through hydrothermal reactions. The distortion of infiltrated beams (1 mm) was 75% less than that of un-infiltrated beams (4 mm) after sintering at 1420 °C for 10 h.

5.2 Chemical reaction

Chemical reaction is another common method to increase green and sintered densities in ceramic binder jetting, which includes metal oxidation [53,54,79,84], phosphoric acid immersion for calcium phosphate materials [113,117,163,119,120,122,123,125,126,141,162], and pyrolysis of preceramic polymers [83,103]. Metal oxidation uses a metal powder as the feedstock material and convert a printed metal part to a ceramic one by oxidation at a high temperature. Phosphoric acid immersion uses this acid solution to immerse calcium phosphate parts, which are printed with a binder of phosphoric acid solution. The immersion leads to a further reaction of unreacted material and thus a cemented structure. Pyrolysis of preceramic polymer uses polymeric precursor

as the feedstock and ceramize the printed part at a high temperature. For example, Zocca et al. [83] investigated the possibility of using silicone (polymethylsilsesquioxane) as a preceramic polymer. A bulk density of 87.1% for the ceramized SiOC part was achieved after sintering at 1200 °C for 1 h. It should be noted that chemical reaction technique only works for certain material systems. Specifically, the ceramization reaction in the preceramic polymer pyrolysis technique involves a significant mass transport of the reagents and/or the by-products. Therefore, part size is limited with this technique.

5.3 Infiltration

Infiltration is another common method to increase the part density. Both a different material [71,85,88,89,162] and the same material [69] have been investigated to infiltrate parts from binder jetting. The most common infiltration method is melt infiltration [39,63,70,71,80,82,85–88,162,172]. In the study of Nan et al. [85], the printed and sintered TiC parts were infiltrated in melted silicon at 1600–1700 °C for 1 h and annealed at 1400 °C for 2 h, which produced Ti₃SiC₂ (about 45 vol.%), TiSi₂ (about 21 vol.%), and SiC. Infiltration with a different material changes the composition of a part and the resultant phases might be difficult to control sometimes.

Solution infiltration can be used for infiltrating a part with the same material, in which fine powder is mixed with a solvent and the part is immersed in it. The most critical parameter for this infiltration method is solid loading. For example, Maleksaeedi et al. [69] printed and pre-sintered alumina parts at 1000 °C for 2 h, infiltrated them in alumina slurries of different solid loadings, and post-sintered them at 1650 °C for 2 h. As the solid loading increased from 0 (no infiltration) to 50%, the bulk density of the final parts increased from 38% to 85%. Cross sections of the infiltrated parts after post-sintering are schematically shown in Figure 14. The infiltration depth for parts with the same shape and size decreased along the increase of solid loading. A noninfiltrated core remained in the parts treated with a slurry of a high solid loading.



Figure 14. Schematic illustration of the cross sections of parts infiltrated at various solid loadings: (a) 35%, (b) 40%, (c) 45%, and (d) 50% [69]

Another method is precursor solution infiltration followed by hydrothermal reactions of the precursor to generate the ceramic material. For example, Grant et al. printed parts with titanium dioxide powder with a particle size of $<63 \mu m$ [89]. Then the printed parts were infiltrated by a 50 wt.% aqueous solution of titanium (IV) bis(ammonium lactato) dihydroxide, followed by heating the parts at 300 °C to turn the precursor into titanium dioxide. The infiltrated parts achieved a density of 55% after sintering at 1420 for 10 h, compared with 48% of un-infiltrated parts. Moreover, the distortion of infiltrated parts (1 mm) was 75% less than that of un-infiltrated parts (4 mm) after sintering.

5.4 Isostatic pressing

Isostatic pressing is a traditional powder metallurgy process that applies equal pressure in all directions on a powder compact, thus achieving uniform density and microstructure [211,229],

The pressing medium that exerts equal pressure can be water, oil, and gas. Based on the operation temperature, this technique is classified as cold isostatic pressing (CIP), warm isostatic pressing (WIP), and hot isostatic pressing (HIP) [229,230]. CIP can consolidate green parts under room temperature to obtain higher green density ready for sintering. WIP operates at intermediate temperatures from 80 °C to 450 °C [230], which is suitable for parts requiring a heat-induced chemical reaction [229]. HIP heats and presses the part under gas medium (e.g., argon) with an accurate control of both temperature (up to 2000 °C) and pressure (50–200 MPa) [229]. It usually requires longer processing time than CIP and WIP.

All three kinds of isostatic pressing techniques have been utilized in ceramic binder jetting. For example, Sun et al. used cold isostatic pressing (CIP) method to consolidate green Ti₃SiC₂ bodies [90], which led to an increased sintered bulk density of 94.3% compared with 65.5% without CIP. In Yoo et al's study, both CIP and WIP were utilized to densify alumina green parts. The bulk density increased from 36% to 54% after CIP and 34% to 61% after WIP, which resulted in sintered bulk densities of 95.9% and 99.2%, respectively [64]. Printed AlN parts were heated to 2000 °C and pressed at 310 MPa for 8 h by HIP, leading to a bulk density of 60.1% [55]. Isostatic pressing can increase green and thus sintered densities significantly, but it is not applicable to parts of complex geometries such as internal cavities.

6 Knowledge Gaps

To produce dense ceramic parts by binder jetting followed by pressureless sintering, fine powder (e.g., less than 1 μ m) is needed because it has a large specific surface energy and thus allows for a high densification level through sintering [52]. However, the spreading, compaction, and densification behaviors are still not well understood.

6.1 Spreading behavior

Fine powder is known for its spreading difficulty: it is challenging to form a dense, smooth, and uniform powder bed. It is generally accepted that the spreading difficulty is a result of interparticle cohesion. However, the nature of the interparticle cohesion is not clear. When discussing the interparticle cohesion, researchers usually refer to van der Waals interaction and electrostatic interaction. Hydrogen bonding and capillary bridging are often overlooked. They could play an important role in powder spreading. In addition, the spreader design, spreading strategy and parameters, and environmental humidity can also affect the spreading behavior. However, these effects are still not well understood.

6.2 Compaction behavior

A high powder bed density is needed to have high green and consequently sintered densities. An effective way to increase powder bed density is compaction, during which the externally applied stress exceeds the yield strength of the powder and leads to particle rearrangement [231,232]. Several compaction approaches, including a forward-rotating roller [64,233] and compaction plate [78], have been reported. However, important issues about this layer-by-layer compaction, have not been studied. For example, effects of compaction parameters on the powder bed and its uniformity with the same layer or across different layers are critical but unknown.

6.3 Densification behavior

The densification behavior of fine powder has been studied for traditional ceramic pressing and sintering. However, binder jetting poses unique challenges for densification. For example, the green density from binder jetting is usually not as high as that from pressing. It is necessary to understand the densification behavior of loosed packed particles. In addition, binder jetting is developed for producing parts with complex geometries. The relationship between geometry and densification is not well understood.

7 Concluding Remarks

Ceramic binder jetting additive manufacturing has many advantages. The main limitation preventing its widespread industrial applications seems to be related to the low density (and thus certain inferior mechanical properties) of printed parts. While various density terminologies have been used for parts by binder jetting, bulk density is more suitable in technical communications for structural applications as it directly determines mechanical properties. It has been reported that special treatment techniques could improve densities. However, most of these techniques have drawbacks, making them inappropriate for a wide range of applications. Granulation is a promising technology, but the literature does not contain sufficient studies on how to eliminate intra- and inter-granule porosities. Fine powder is needed to obtain dense ceramic parts by binder jetting, but there are no reported studies on its spreading, compaction, and densification behaviors.

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Special treatment	Density or porosity type	Value (%)	Standard deviation (%)	Reference
/	Bulk density	32	/	[55]
/	Bulk density	39	/	[89]
/	Bulk porosity	60.5	1.4	[110]
/	Bulk density	40	/	[82]
/	Bulk density	46.8	1.9	[57]
/	Bulk density	50.7	/	[58]
/	Bulk density	55	/	[73]
/	Apparent porosity	35.28	/	[161]
G	Bulk density	31	/	[64]
G	Bulk density	35.3	/	[63]
G	Bulk density	36	/	[64]
G	Bulk density	42	/	[92]
G	Apparent solid density	45	/	[68]
G	Bulk density	45.9	/	[86]
L	Bulk density	54	/	[25]
L	Bulk density	57.2	/	[38]
L	Bulk density	58	/	[27]
L	Bulk density	63.4	/	[23]
L	Bulk density	66	/	[77]
L	Bulk density	67	/	[24]
S	Bulk density	36.05	0.57	[54]
S	Bulk density	39.6	0.6	[67]
S	Bulk density	45	/	[74]
S	Bulk porosity	55	/	[156]
S	Bulk density	65.7	/	[58]
S, G	Apparent porosity	55	/	[88]
S, G	Bulk porosity	55	/	[87]

Appendix A. Supplementary data for Figure 9.

Special	Density or porosity	Value	Standard	Reference
	Bully density	25		[55]
/	Bulk density	38.4	/	[55]
/	Dulk density	20.4 /8	/	[09]
/	A moment monosity	46.07	8 52	[09]
/	Apparent porosity	40.07	0.52	[132]
/	A moment a modified	<i>JJ.09</i> <i>A1 A2</i>	1 25	[90]
/	Apparent porosity	41.42	4.55	[137]
/	Apparent sond density	22.1	26	[170]
/	Bulk porosity	32.1 22	2.0	[110]
C C	Dulk porosity	52 55 2	/	[109]
C C	Dulk density	12.0	0.5	[33]
C C	Auropeant solid density	12.9	0.5	[85]
G	Apparent solid density	32.2 29.79	2 (1	[85]
G	Bulk density	38.78 50.42	3.64	[135]
G	Apparent porosity	59.43 42.05	/	[149]
G	Bulk density	42.95	1.6	[130]
G	Bulk density	43.2	/	[158]
G	Bulk density	48	/	[127]
G	Apparent solid density	48	/	[68]
G	Apparent porosity	51	0.9	[155]
G	Bulk density	62.5	/	[64]
G	Apparent porosity	36	/	[70]
G	Bulk density	67	/	[63]
G	Apparent solid density	91.4	2.8	[128]
l	Bulk density	85	/	[69]
L	Bulk density	99	/	[24]
L	Bulk density	99.9	/	[27]
М	Bulk porosity	65.3	/	[144]
M	Bulk porosity	44	/	[108]
М	Bulk density	66	/	[97]
M	Bulk density	72	2	[96]
М	Bulk density	94.5	/	[174]
Р	Bulk density	60.1	/	[55]
Р	Bulk density	94.33	/	[90]
S	Bulk density	65.5	2.4	[65]
S	Bulk porosity	30	/	[156]
S	Apparent solid density	96	/	[169]
G, I	Bulk porosity	19	0.5	[71]
G, I	Bulk density	91	/	[86]
G, I	Apparent porosity	2.4	/	[85]
G, P	Bulk density	97.8	/	[64]

Appendix B. Supplementary data for Figure 10.

G, S	Bulk density	86.8	/	[95]
I, C	Bulk density	55	/	[89]
M, G	Bulk density	45.06	3.05	[135]
M, G	Bulk density	52.84	2.76	[136]
M, G	Apparent solid density	95.1	4.72	[128]
M, G	Bulk density	95.74	/	[92]
M, G	Apparent porosity	2.5	0.12	[102]
M, G	Apparent solid density	99	0.1	[146]
M, L	Bulk porosity	8.79	1.44	[77]
M, L	Bulk density	99.2	/	[38]
M, S	Bulk density	54.8	/	[98]
M, G, P	Bulk density	99.2	/	[64]
M, G, C	Bulk porosity	48	2	[103]
M, G, I	Apparent porosity	1.7	/	[88]
M, L, P	Bulk density	100	/	[94]
M, S, G	Bulk density	53.01	/	[100]

Density before sintering		Domaitry often			
Powder bed density (%)	bed Green Press (%) density (%) dens		sintering (%)	Reference	
39.7	/	/	46.8	[65]	
45.2	/	/	50.7	[65]	
51	/	/	87.1	[83]	
53	/	/	100	[94]	
57.1	/	/	57.5	[65]	
58.2	/	/	60.5	[65]	
59.5	/	/	64.3	[65]	
53.6	/	/	64.8	[65]	
61.7	/	/	65.5	[65]	
67	/	/	99	[24]	
/	31	/	62.5	[64]	
/	32	/	35	[55]	
/	39	/	48	[89]	
/	43.6	/	61.5	[63]	
/	45	/	48	[68]	
/	45	/	70	[156]	
/	46.9	/	64.1	[63]	
/	48	/	67	[63]	
/	57.2	/	99.2	[38]	
/	58	/	99.9	[27]	
/	/	47	87.9	[64]	
/	/	54	95.9	[64]	
/	/	56	97.8	[64]	
/	/	61	99.2	[64]	

Appendix C	C. Suppleme	entary data	for Figure	13.
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Table Caption List

Table 1. Advantages and disadvantages of various AM categories for ceramics

Table 2. Feedstock and resultant materials in reported studies on ceramic binder jetting

Table 3. Binder materials in reported studies on ceramic binder jetting

Table 4. Terminologies for density and porosity

Table 5. Capabilities of different density and porosity measurement methods

Table 6. Theoretical density values of different materials

Table 7. Granulation process practiced in reported studies

Table 8. Reported studies on mixing powders of different sizes

Figure Caption List

Figure 1. Major components of a binder jetting 3D printer

Figure 2. Steps of ceramic binder jetting

Figure 3. Proportions of various application areas of ceramic binder jetting reported in the literature

Figure 4. Different ceramic particle shapes used in binder jetting: (a) spherical, (b) rounded [179], (c) angular, (d) polygonal [172], (e) irregular [102], and (f) aggregate.

Figure 5. Various powder deposition methods: (a) doctor blade, (b) fixed roller, (c) forward-rotating roller, and (d) counter-rotating roller. h is desired layer thickness and h_d is pre-deposited thickness.

Figure 6. Two binder application methods: (a) binder jetting in-place and (b) powder-binder premixing

Figure 7. Proportions of different binder application methods in reported studies

Figure 8. Open and closed pores in a cross section of a part

Figure 9. Achieved values of relative green densities and porosities by various special treatment techniques (N stands for no special treatment, G for powder granulation, L for using slurry feedstock, and S for mixing powders of different sizes) [23–

25,27,38,54,55,57,58,63,64,67,68,73,74,81,82,86-89,92,110,156,161]

Figure 10. Achieved values of relative sintered densities and porosities by various special treatment techniques (N stands for no special treatment, C for chemical reaction, G for powder granulation, I for infiltration, L for using slurry feedstock, M for mixing different materials, P for pressing, and S for mixing powders of different sizes) [24,27,38,53,55,63–65,68–

71,77,83,85,86,88–90,92,94–98,100,102,103,108–

110, 127, 128, 130, 132, 135, 136, 144, 146, 149, 155-158, 169, 170, 174]

Figure 11. Principle of granulation

Figure 12. Basic granulation steps

Figure 13. Bulk densities before and after sintering [24,27,94,156,38,55,63–65,68,83,89]

Figure 14. Schematic illustration of the cross sections of parts infiltrated at various solid loadings: (a) 35%, (b) 40%, (c) 45%, and (d) 50% [69]