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Trends in bioactivity: inducing and detecting mineralization of regenerative polymeric scaffolds

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Due to limitations of biological and alloplastic grafts, regenerative engineering has emerged as a promising alternative to treat bone defects. Bioactive polymeric scaffolds are an integral part of such an approach. Bioactivity importantly induces hydroxyapatite mineralization that promotes osteoinductivity and osseointegration with surrounding bone tissue. Strategies to confer bioactivity to polymeric scaffolds utilize bioceramic fillers, coatings and surface treatments, and additives. These approaches can also favorably impact mechanical and degradation properties. A variety of fabrication methods are utilized to prepare scaffolds with requisite morphological features. The bioactivity of scaffolds may be evaluated with a broad set of techniques, including *in vitro* (acellular and cellular) and *in vivo* methods. Herein, we highlight contemporary and emerging approaches to prepare and assess scaffold bioactivity, as well as existing challenges.

1. Introduction

1.1. Current methods to treat bone defects

Bone tissue is critical for mechanical functionality, protection, and hematopoiesis. 1,2 These traits stem from its unique combination of carbonated hydroxyapatite (HAp) $[Ca_{10}(PO_4)_6(OH)_2]$ embedded in an extracellular matrix (ECM) comprised of collagen (primarily type I), proteoglycans, and glycoproteins. While capable of regeneration, bone tissue healing is limited for defects beyond a critical size that stem from traumatic injury, surgical excision, or congenital anomalies. Bone tissue healing is also hindered by advanced age, osteoarthritis, and radiological treatment. Numerous products have been developed to treat and heal critical-sized bone defects, and are exemplified in Table 1.

Biological grafting approaches are frequently employed, wherein the living tissue graft becomes incorporated into the surrounding tissue. Autografting remains the 'gold standard' with over two million bone autografts performed annually. However, autografting is associated with complex surgical harvesting (e.g., from tibia or iliac crest), donor site morbidity, and limited availability, as well as premature resorption stemming from poor contact with adjacent tissue. Cadaveric

mitigate brittleness and afford replacement by neotissue,

numerous degradable synthetic and natural polymers have been employed.^{19–22} Synthetic polyesters and copolymers

allografts are also associated with limited availability and

premature resorption, as well as immune rejection. 4 Xeno-

grafts, particular bovine grafts, have been explored but pose a

risk for disease transmission, a greater chance of host immune

response, highly variable resorption rates, and reduction in

osteoinductive properties due to strict manufacturing and

processing requirements.^{5,6} Alloplastic bone substitutes have been leveraged as an alternative to biological grafts. For example, demineralized bone matrix (DBM) is prepared via decalcification (i.e., removal of HAp) of cortical bone allografts with an acidic solution that leaves behind a composite of collagens, non-collagenous proteins, growth factors, residual calcium phosphate mineral (1–6%), and trace cellular debris. Alloplastic grafts have also been prepared based on one or more synthetic bioactive "bioceramics", including: bioactive glasses (BGs), 8,9 HAp^{10–12} β -tricalcium phosphate [β -Ca₃(PO₄)₂; β -TCP], 13,14 and calcium sulfate. 15 Silicate BGs are widely used for their capacity to bond to bone, and represent certain compositions of SiO₂-Na₂O-CaO-P₂O₅ (e.g., 45S5-BG). In the granular form, DBM and bioceramics provide advantageous microporosity and complete resorption. Still, these are also often combined with a polymer coating or matrix to afford injectability or moldability within irregular shapes, as well as to circumvent brittleness in certain cases. 16 The use of poly(methyl methacrylate) (PMMA) is associated with exothermic cures, post-cure shrinkage, lack of porosity, and nondegradability, as well as brittle mechanical properties. 17,18 To

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Perspective

Product name	Bioactive component	Matrix/carrier component
AlloFuse [®] (AlloSource)	DBM	Reverse phase medium gel
Calcigen® S (Zimmer Biomet)	Calcium sulfate	Not reported
Cerament® (BONESUPPORT)	HAp & calcium sulfate	Not reported
CranioSculpt [™] (KLS Martin)	Calcium phosphate	Not reported
Cortoss [®] (Stryker)	Bioactive glass ceramic	Acrylate copolymer
Grafton™ DBM (Medtronic)	DBM	Glycerol
Kinex [®] (Globus Medical)	BG	Collagen, hyaluronic acid
MinerOss [®] (BioHorizons)	Cortical & cancellous bone chips	None
NanoFUSE [®] (Amend Surgical)	DBM & BG-45S5 coated with gelatin	None
Optium [®] (LifeNet Health)	DBM	Glycerol
OSferion (Arthrex)	β-ТСР	None
Osteosponge [®] (XTANT Medical)	DBM	None
Puros® (ZimVie)	DBM	Reverse phase medium gel
Vestakeep [®] Fusion (Evonik)	Biphasic calcium phosphate	PEEK

Table 1 Examples of commercial biological and alloplastic materials to treat bone defects

thereof have been widely utilized given the tunability of properties.²⁰ More recently, polyether ether ketone (PEEK) 3D printed alloplastic devices have been created in complex, patient-specific geometries, 23 including as composites with bioactive fillers.24,25

1.2. Regenerative engineering approaches with bioactive

Owing to the limitations of biological and alloplastic grafts, regenerative engineering approaches have emerged to heal bone defects. 26 The scaffold compositions play an instrumental role, and must fulfill a demanding set of criteria to maximize bone tissue healing. 27,28 Bioactivity is of significant importance as it leads to the formation of HAp that promotes osteogenic differentiation (i.e., osteoinductivity), as well as osseointegration with surrounding bone tissue (Fig. 1). 29-31

While bioactive HAp is innately present in biological grafts, for alloplastic grafts, bioactivity is traditionally afforded by the inclusion of DBM or bioceramics. Scaffolds that are potently bioactive may reduce or even eliminate the necessity of exogeneous growth factors (e.g., bone morphogenic protein 2, BMP-2), which risk off-target responses.³² Beyond bioactivity, the scaffold must also be osteoconductive (i.e., permitting cell migration and neotissue infiltration).33,34 This is achieved through porosity that may be afforded through a variety of fabrication methods (e.g., 3D printing, solvent cast particulate leaching [SCPL], gas foaming, freeze drying, and electrospinning) with polymers forming the regenerative bone scaffold.³⁵

Degradation of the scaffold also facilitates osteoconductivity, making the rate of scaffold resorption important to healing. To enable osseointegration, the scaffold must form close contact with adjacent bone tissue.²⁹ This has been particularly addressed with injectable scaffolds, 36 3D-printed scaffolds, 37,38 and shape memory polymer (SMP) scaffolds. 39

Imparting bioactivity to scaffolds

Given the importance of bioactivity to bone regeneration, bioactive polymeric scaffolds continue to be developed (Fig. 2). Approaches include bioactive composite scaffolds based on combining polymers with bioactive fillers (e.g., DBM and bioceramics), bioactive coatings and surface treatments, as well as surface modification of scaffolds. Recent reports are exemplified herein.

2.1. Bioactive composite scaffolds

Bioactive composite scaffolds, comprised of one or more bioceramics embedded in a polymer matrix, 40-42 remain prolific in regenerative bone engineering. Versus bioceramic-only scaffolds, these composites can improve processibility, increase rigidity and strength, and mitigate brittleness that contribute to post-surgical fracture. Furthermore, owing to the hydrophilicity and susceptibility to hydrolysis, bioceramic-polymer composites also degrade at favorably faster rates compared to polymer-only scaffolds. A variety of bioactive composite scaffolds have recently been reported (Table 2). Both biodegradable

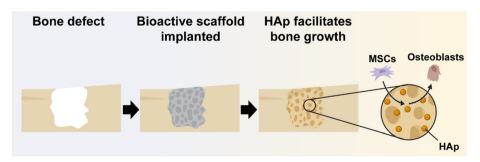


Fig. 1 Bioactive scaffolds lead to HAp mineralization, and subsequently promote osteogenesis and osseointegration.

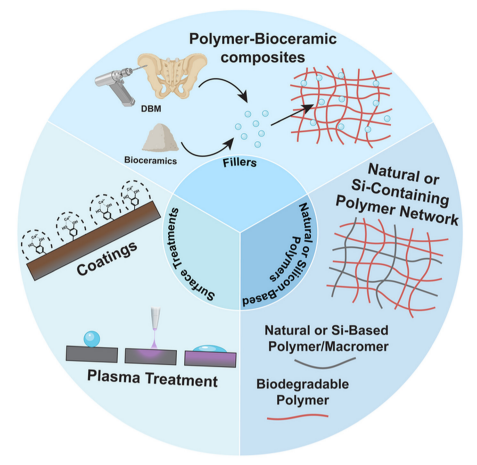


Fig. 2 Methods to prepare bioactive polymeric scaffolds for bone regeneration.

synthetic and natural polymers have been utilized. A variety of bioceramics and glasses have been leveraged, ranging from historically used types (e.g., BG, HAp, and β -TCP), to newer types (e.g., nanosilicates, and eggshell particles). Their mineralization activity and mechanism vary, with some capable of acellular mineralization in physiological environments including simulated body fluid (SBF) (Fig. 3). For instance, 45S5-BG [45 wt% SiO₂, 24.5 wt% Na₂O, 24.5 wt% CaO, and 6 wt% P₂O₅] is known to promote rapid HAp formation on its surface within hours. 43 The mechanism involves a series of sequential steps: (1) exchange of Na⁺ ions with solution H⁺, (2) hydrolysis of Si-O-Si bonds to form SiOH bonds, and the release of Si(OH)₄, (3) polycondensation to form a hydrated silica gel, (4) formation of an amorphous calcium phosphate phase via absorption of Ca²⁺, PO₄³⁻, and CO₃²⁻ ions, and (5) crystallization to carbonated HAp. Synthetic HAp is also capable of inducing the formation of a HAp layer in SBF, and at greater levels versus BG-45S5.44 HAp forms through several steps: (1) adsorption of Ca²⁺ ions to the surface, (2) formation of Ca-rich amorphous calcium phosphate at surface, (3) transition to a Ca-poor surface due to adsorption of solution PO₄³⁻ as well as CO₃²⁻ ions, and (4) crystallization into carbonated HAp.45 In addition to bone bonding, the formed HAp layer facilitates bone formation via the stimulation of mesenchymal stem cell (MSC) osteogenesis,

especially via increased expression of growth factors (e.g., BMP) and enhanced alkaline phosphatase (ALP) activity. In contrast, β -TCP does not mineralize with SBF exposure, but rather is osteoconductive and osteoinductive. 14 β -TCP leads to osteoclast-mediated resorption and osteoconduction that is associated with rapid bone formation and high bone bonding strengths.

For bioactive composite scaffolds, the level of incorporated bioactive bioceramics is highly variable, ranging from less than 1 wt% to over 50 wt%. Composites prepared with two or more distinct types of bioceramics have also been reported. A number of fabrication methods have been employed (e.g., SCPL, electrospinning). Notably, various forms of 3D printing have been leveraged extensively to impart finer control of microarchitecture as well as to produce patient-specific scaffolds.⁴⁶ For instance, selective laser sintering (SLS) 3D printing employs a laser to sinter a powder (e.g., a mixture of polymer and bioceramic) and fuse particles together, while unfused particles support the structure. Comparison of composite scaffold bioactivity efficacy is difficult, as in vitro and in vivo evaluations of such scaffolds are highly variable in the literature. For example, time-points selected to confirm HAp mineralization following exposure to SBF vary appreciably (from 1 day to several weeks). Still, some studies directly compare scaffolds prepared with two different bioceramics.

 Table 2
 Examples of bioactive composite scaffolds for bone regeneration

	-	-	n		
Study	Bioactive component	Matrix component	Matrix component Wt% of Filler	Fabrication method	Key findings
Shuai, 2021 ⁴⁷	НАр	PLLA, or PGA	10%	3D printing (SLS)	HAp/PLLA/PGA scaffolds displayed tunable degradation and mechanical properties.
Xu, 2017 ⁴⁸	Xu, 2017 ⁴⁸ HAp or β-TCP	PLGA	50%	SCPL	HAp/PLGA scaffolds exhibited higher strength and faster degradation; TCP/PLGA scaffolds improved bone regeneration in a rabbit model.
Cheng, 2021 ⁴⁹	β -TCP (with cucurbitacin B)	PLGA	25%	3D printing (low temperature rapid prototyping)	TCP/PLGA/Cur-B scaffolds increased bone regeneration and angiogenesis in a rat model.
Nyberg, 2017^{50}	HAp, β-TCP, DBM, or BO	PCL	30%	3D printing (extrusion)	DBM/PCL and BO/PCL scaffolds exhibited greater osteoinduction; PCL/HAp displayed higher compressive modulus.
Shuai, 2022 ⁵¹	PDLA-grafted HAp (g-HAp)	PDLA	0.5, 1, 2 or 4%	3D printing (SLS)	g-HAp/PDLA scaffolds exhibited increased modulus and strength.
Sultan, 2022 ⁵²	45S5-BG	PLA	5%	TIPS and 3D printing (extrusion)	45S5-BG/PLA scaffolds exhibited increased strength.
Nitschke, 2023 ⁵³	45S5-BG	PCL, PLLA	PCL, PLLA 2.5 to 30%	Modified SCPL	45S5-BG/SMP scaffolds exhibited faster degradation, and retained shape memory behavior.
Distler, 2020^{54}	45S5-BG	PLA	1–10 wt%	FDM	45S5-BG/PLA scaffolds exhibited decreased strength with increased BG wt%; triggered cellular osteogenesis.
Monfared, 2022^{55}	45S5-BG and β -TCP	Gelatin, and PVA	35–40% (equal parts each)	3D printing (extrusion)	45S5-BG/TCP/Gelatin/PVA scaffolds exhibited increased compressive strength and faster degradation.
Han, 2023 ⁵⁶	BBG	PCL	5-40%	3D printing (SLS)	BBG/PCL scaffolds with 20 wt% BGG exhibited optimal mechanical properties, and promoting regeneration & integration in a rabbit model.
Hatton, 2019 ⁵⁷	ICIE16M-BG	Alginate	50%	Freeze-drying	ICI16M-BG/alginate scaffolds exhibited increased strength.
Du, 2019 ⁵⁸ MBG	MBG	SF, or PCL 80%	%08	3D printing (extrusion)	MBG/SF exhibited superior compressive strength, and greater heterotopic bone formation in a mouse model.
Qi, 2017 ⁵⁹		PCL	70%	3D printing (extrusion)	MBG/CS scaffolds exhibited superior osteogenesis of cultured human bone marrow mesenchymal stem cells (hBMSCs) and greater bone formation in a rat model.
Carrow, 2019^{60}	LAPONITE®	PEOT/PBT	5, 10, or 15%	3D printing (extrusion)	LAPONITE $^{(g)}$ PEOT/PBT scaffolds exhibited decreased degradation rates <i>versus</i> copolymer-only scaffolds, as well as increased osteogenesis of hMSCs.
Wu, 2021 ⁶ .	Wu, 2021 ⁶¹ Eggshell micro- particles (ESP)	Gel-MA	ESP combined with 5% prepolymer solution	Casting/UV cure	ESP/Gel-MA scaffolds exhibited superior osteogenesis of cultured MC3T3-E1 pre-osteoblasts versus Gel-MA-only scaffold, and greater bone formation in a rat model.
Huang, 2020 ⁶²	MWCNTs &/or nHAp	PCL	0.75% MWCNT & 20% nHAp	3D printing (screw assisted extrusion)	PCL/MWCNT/nHAp scaffolds exhibited increased modulus versus PCL-only scaffolds, and greater osteogensis of cultured hADSCs.

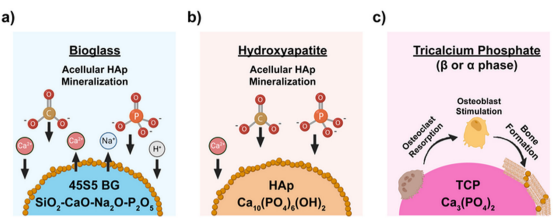


Fig. 3 Simplified mechanisms of mineralization for common bioceramics: (a) BG-45S5, (b) synthetic HAp, and (c) β-TCP.

Both HAp and β-TCP continue to be utilized to prepare bioactive composite scaffolds, either alone or in combinations. Shuai et al. utilized SLS 3D printing to fabricate composite scaffolds based on HAp (10 wt%) with poly(L-lactic acid) (PLLA) and poly(glycolic acid) (PGA).47 Versus HAp/PLLA composite scaffolds, the faster degradation of HAp/PLLA/PGA composites (50:50 wt% PLLA: PGA) enhanced the exposure of HAp, leading to superior mineralization and regeneration in a rabbit segmental defect model. Xu et al. reported both HAp- and β-TCPcontaining poly(lactic-co-glycolic acid) (PLGA) composite scaffolds via SCPL. 48 Compressive tests revealed that HAp-scaffolds had higher strengths and rates of in vitro degradation compared to β-TCP-scaffolds. In a rabbit calvarial defect, while early-stage bone growth was faster for HAp-scaffolds, β-TCPscaffolds exhibited greater bone mineral densities and higher compressive strengths of repaired bone at 20 weeks. Cheng et al. introduced cucurbitacin B, a plant-derived terpene, to β-TCP/PLGA scaffolds leading to enhanced osteogenesis and neovascularization.49 In an example by Nyberg et al., a series of 3D printed composite poly(e-caprolactone) (PCL) scaffolds were prepared by altering the bioactive filler: a bioceramic (HAp or β-TCP) or a biologic (decellularized bone matrix [DBM] or Bio-Oss[®] [BO], a clinically available form of DBM).⁵⁰ BO/PCL and DBM/PCL scaffolds exhibited enhanced osteoinduction versus HAp/PCL and TCP/PCL scaffolds, while the compressive modulus was highest for HAp/PCL scaffolds. Shuai et al. reported poly(D,L-lactic acid) (PDLLA) grafted onto HAp (g-HAp), combined with PLLA to prepare composite scaffolds via SLS 3D printing.⁵¹ As a result of improved interfacial bonding via stereo-complexation, g-HAp/PLLA composite scaffolds displayed significant enhancements in stiffness and strength.

BGs continue to be utilized to prepare bioactive composite scaffolds. Sultan *et al.* prepared composite scaffolds based on 45S5-BG (5 wt%) and poly(lactic) acid (PLA).⁵² Thermally induced phase separation (TIPS) was used to create a homogenous distribution of BG, and the resulting composite spheres were subsequently formed into scaffolds *via* extrusion 3D printing. Due to the homogeneous distribution of BG, scaffold compressive strength increased for printed scaffolds.

We have previously reported 'self-fitting' SMP scaffolds based on PCL,63 including PCL/PLLA semi-interpenetrating networks (semi-IPNs).⁶⁴ Recently, Nitschke et al. reported analogous composite scaffolds that included 45S5-BG.53 A modified SCPL protocol was utilized wherein the fused template was formed from a mixture of BG and salt, resulting in localization of BG on the scaffold pore walls. At just 5 and 10 wt% BG, 45S5-BG/PCL scaffolds induced HAp mineralization after 1 day in SBF (1X), and degraded faster versus corresponding polymeronly scaffolds, all while maintaining shape memory behavior. Distler et al. described 45S5-BG/PLA scaffolds with 1-10 wt% BG formed by fused deposition modeling (FDM) of 45S5-BG/ PLA filaments.⁵⁴ Monfared et al. combined 45S5-BG with β-TCP (50:50 wt% ratio; 35-45 wt% total) to prepare composite scaffolds based on gelatin, poly(vinyl alcohol) (PVA), and Tween[®] 60 using extrusion-based 3D printing.⁵⁵ The printed scaffolds achieved higher moduli versus analogous scaffolds prepared by foam casting.

Newer types of bioceramics have also been developed and used to prepare bioactive composite scaffolds. For example, borate-containing BGs (BBGs), wherein borate (B2O3) is partially or completely substituted for silica (SiO₂), are associated with faster rates of HAp formation compared to silicate bioactive glasses (e.g., 45S5).65 Thus, BBG composite scaffolds have been formed, as in Han et al. wherein BBG/PCL scaffolds were prepared by SLS 3D printing.⁵⁶ Furthermore, ICIE16-BG, a potassium-containing BG (48.0 SiO₂, 6.6 Na₂O, 32.9 CaO, 2.5 P₂O₅, 10.0 K₂O [wt%]), was developed as an alternative to 45S5-BG.57 For instance, Hatton et al. combined ICIE16-BG with alginate to form composite scaffolds via freeze-drying.66 Still, different types of bioactive silicate bioceramics have been leveraged to form bioactive composite scaffolds. Mesoporous bioactive glasses (MBG) based on silicates have well-defined pores with diameters around 5 to 20 nm, presenting a large surface area.⁶⁷ Du et al. 3D printed composite scaffolds from MBG (80 wt%) combined with silk fibroin (SF) and PCL, with MBG/SF scaffolds exhibiting superior strength and bioactivity versus MBG/PCL scaffolds.⁵⁸ While relatively low cost, calcium sulfate exhibits particularly rapid resorption as well as limited bioactivity.⁶⁸ Thus, Qi et al. prepared 3D printed scaffolds wherein MBG and calcium sulfate were combined with PCL.⁵⁹ In addition to MBG, other nanosized bioactive fillers have also been utilized to form bioactive scaffolds,69 including HAp nanoparticles, 70 and nanosilicates (e.g., LAPONITE®). 71,72 Carrow et al. reported 3D printed scaffolds using LAPONITE® in combination with a poly(ethylene oxide terephthalate) (PEOT)/ poly(butylene terephthalate) (PBT) (PEOT/PBT) copolymer. 60 Wu et al. reported the use of chicken eggshell microparticles (ESP), possessing high levels of calcium and representing a sustainable alternative to BGs, to form bioactive composite scaffolds in combination with gelatin methacrylate (Gel-MA). 61 To achieve biomimetic scaffolds in terms of not only bioactivity but structure, Huang et al. prepared PCL scaffolds with multi-walled carbon nanotubes (MWCNTs) and nano-HAp (nHAp), using screw-assisted extrusion based 3D printing to align the MWCNTs. 62 Piezoelectric perovskites such as barium titanate (BTO) have also shown to be capable of HAp mineralization.⁷³ In the case of BTO, when exposed to an electric field or mechanical stress, the titanium and oxygen ions switch locations, leading to a concentration of negatively charged O²⁻ on the material surface. In turn, positively charged Ca²⁺ ions from the physiologic fluid are attracted to the surface, leading to the formation of an apatite layer.⁷⁴

2.2. Bioactive coatings and surface treatments

Various bioactive coatings have historically been applied to metal implants. 75 To achieve bioactivity of polymeric scaffolds, coatings have been likewise applied.76,77 Such coatings can enhance the bioactivity of composite scaffolds, or be used in lieu of fillers to avoid brittleness. Bioceramic coatings may also be directly applied to scaffolds. Recently, several bioactive coatings and surface treatments have been applied to scaffolds to promote bioactivity (Table 3). For instance, Fazeli et al. reported the deposition of HAp and BG onto 3D printed PCL scaffolds via an immersion method with a HAp/BG solution.⁷⁸ Zhang et al. reported PCL scaffolds coated with HAp via pulsed laser deposition (PLD).⁷⁹ Li et al. described HAp deposition onto PVA/PLA scaffolds via electrodeposition. 80 Coatings based on a combination of bioceramics and polymers have also been reported. For instance, based on the bioactivity of chitosan, 81 Shaltooki et al. applied a chitosan/BG coating to PCL/BG composite scaffolds by exposing scaffolds to homogenized solutions.82 Alternatively, a bioactive polymeric coating may be applied to scaffolds. Collagen type I, a natural component of bone tissue, is a frequently used material for bone regeneration.⁸³ Thus, Tabatabaei et al. reported PCL/β-TCP scaffolds coated by collagen using an immersion method that included homogenization.84 While developed for the purpose of assessment of biomaterial bioactivity, SBF exposure has been used to deposit bioactive mineral coatings onto polymeric scaffolds. 76,85 Polydopamine (PDA) was established to readily form adherent coatings onto substrates via base-catalyzed autooxidation of dopamine. 86,87 PDA was confirmed to induce HAp mineralization in SBF, 88 leading to its use as a coating to create bioactive scaffolds for bone regeneration. Numerous

studies have been noted in a recent review by Tolabi et al., with the scaffold polymer component frequently being a biodegradable polyester. 89 We likewise applied a PDA coating to polyester SMP scaffolds, which resulted in enhanced osteogenic differentiation of hMSCs as well as HAp mineralization after SBF exposure.90

Direct surface treatment of polymeric scaffolds has also been utilized to invoke bioactivity. 76,91 Plasma treatment, wherein an electric current passes through a gas (e.g., oxygen, argon, and ammonia), is a popular method. 92 This process can be used to produce surface functionalization, etching, or film deposition, while maintaining scaffold bulk properties. 93 Oxygen plasma treatment has been extensively utilized, including for PCL-based scaffolds, to enhance surface hydrophilicity and surface energy for improved cellular adhesion. 94 Depending on plasma parameters and polymer type, oxygen plasma treatment can result in both surface functionalization and etching. 93 For instance, Kim et al. reported oxygen plasma treatment of PCL scaffolds resulting in enhanced hydrophilicity and surface roughening.95 These scaffolds exhibited in vitro mineralization by cultured cells. To confirm that such surfaces also give rise to acellular mineralization, Murab et al. used oxygen plasma treated PCL/TCP scaffolds exposed to SBF to demonstrate that resulting -COOH groups act as nucleation sites for amorphous calcium phosphate to form HAp crystals. 96 A significant challenge of plasma generated polymeric surfaces is their ageinstability, as surfaces hydrophobically recover to the untreated state within as little as hours. 109 However, some examples display improved stability. For instance, Yamada et al. reported oxygen plasma treated PLA-co-trimethylene carbonate (PLA-co-TMC) scaffolds whose surfaces were stable for over 2 weeks.⁹⁷ Plasma-enhanced chemical vapor deposition (PECVD) may be utilized to apply thin films, including inorganic-organic composite films. 110 However, recent reports on its use to enhance polymeric bioactivity appear scarce. Terriza et al. used deposition by PECVD of SiO2 onto the surfaces of PLGA membranes, resulting in morphological changes to osteoblasts.98 Another form of CVD, initiated CVD (iCVD), employs a combination of a volatile initiator as well as monomer(s) to produce thin films. 110 This process avoids fragmentation of organic precursors as with PECVD. Song et al. reported formation of a polyelectrolyte coating via iCVD onto HAp scaffolds that was then exposed to supersaturated HAp, resulting in mineralized scaffolds that promoted osteogenesis.99

2.3. Bioactive polymers as additives

As previously noted, a variety of synthetic and natural polymers are utilized to form bone tissue scaffolds. Most of these are considered "nearly inert", or lacking in bioactivity to promote bone regeneration.111 Yet, several natural polymers (e.g., collagen, 112 gelatin, 113 chitosan, 81 alginate, 114 and hyaluronic acid¹¹⁵) display bioactivity, and have thus been formed into regenerative bone scaffolds (Table 3). In addition to being used as coatings, these bioactive polymers may be used as an additive in combination with a nearly inert polymer to form the scaffold bulk or a discrete structure within the inert

Table 3 Recent studies in bioactive coatings, surface treatment, and polymers

Bioac	Bioactive Ma coating co i, 2023 ⁷⁸ HAP PC g, HAP PC 119 ⁸⁰ HAP PC 119 ⁸⁰ HAP PC 12023 ⁷⁸ HAP PC 12020 ⁸⁶ Collagen PC Rim, 2014 ⁹⁵ O ₂ plasma 2020 ⁹⁶ PECVD PC Rim, 2014 ⁹⁵ O ₂ plasma 2021 ⁹⁷ PECVD PC Bioactive su treatment Rim, 2014 ⁹⁵ O ₂ plasma 2021 ⁹⁷ PECVD Song, Song, ICVD Bioactive additive additive se, 2009 ¹⁰⁰ Collagen ang, 2021 ¹⁰¹ Gelatin nityaghoubi, Chitosan 2021 ¹⁰² Alginate my, 2021 ¹⁰³ Fibrin/ alginate my, 2021 ¹⁰³ Hyaluroni assica, PDMS _{sur} PDMS _{sur} PPO Alginate Bioactive additive additive additive additive Alginate PS PS PS PS PS PS PS PS PS P		Coating meth Immersion in PLD Electrodepos Immersion o osan/BG solu Immersion in lagen solutio Submersion in RATIX COMPONENT PCL PCL PCL PCL PCL PCL PCC PLGA HAP Electrospi MEW of F COAting w Spray-prec microsph Solvent in	region (See findings) HAp-coated scaffolds displayed increased osteogenesis in rat calvarial defect models. Increased osteogenic potential of HAp-coated scaffolds cultured with rat BMSCs, and superior bone regeneration in critical rat calvarial defects. Longer electrodeposition times resulted in increased surface roughness and higher crystallinity of HAp. Chitosan/BC-coated scaffolds displayed increased hydrophilicity, degradation, and osteogenic properties. In PDA solution PDA solution PDA solution and surface roughness, acellular mineralization in SBF, and higher osteogenesis with cultured h-MSCs. Increased hydrophilicity and surface roughness, acellular mineralization in SBF, and increased osteogenesis with cultured h-MSCs. Increased hydrophilicity and surface roughness and hydrophilicity, and improved osteogenesis of cultured rat BMSCs. Is nm StO ₂ film deposition onto PLGA films produced changes to osteoblast morphology. IcVD treatment coated scaffolds with a polyelectrolyte induced mineralization and enhanced osteogenesis of cultured MC3T3-E1s. Increased hydrophilicity and osteogenic potential in vitro compared to PCL-only manofibers. PCL/HOkintosan seroffolds subject or an animal defects. Increased differentiation of hBMSCs. PCL/HA microspheres exhibited increased bydrophilicity, surface roughness, and promoted osteogenic differentiation of hBMSCs. PCL/HA microspheres exhibited increased osteogenic potential in vitro and bone regeneration in rate cranial defects. In art at model. PCL/HA microspheres exhibited increased osteogenic potential in vitro and osteogenesis of hBMSCs. PCL/HA microspheres exhibited increased osteogenic potential in vitro and bone regeneration in ethors of pages. PDMS _{sur} PEG lydrophilacy acellular mineralization on SEP. PDMS _{sur} PEG lydrophilacy acellular mineralization in SBF, and osteogenesis of hBMSCs.
2019 ¹⁰⁵ Frassica, 2020 ¹⁰⁶ Beltran, 2021 ¹⁰⁷ Beltran, 2021 ¹⁰⁷		DA PEG-DA - PCL-DA - PCL-DA	Separation (SIPS) SIPS SCPL	PPMS/PCL scaffolds exhibited superior osteogenic potential <i>in vitro</i> compared to PEG and PEG/PDMS scaffolds. PDMS-containing scaffolds mineralized in 1X SBF and presented accelerated degradation rates. PMHS-containing scaffolds had a faster onset of acellular mineralization in SBF, and expedited acceleration profiles compared to analogous PDMS scaffolds.

polymer. In some cases, bioactivity is demonstrated in the absence of bioactive fillers. For instance, collagen has been blended with polyesters to form electrospun scaffolds, 116 including for bone regeneration per Jose et al. 100 Wang et al. combined gelatin with PCL to produce scaffolds with hierarchal morphological structures using melt electrospinning writing (MEW) and solution electrospinning (SE). 101 Amiryaghoubi et al. introduced chitosan to PCL/polyurethane (PU) to form scaffolds via freeze-drying. 102 Ren et al. prepared PCL scaffolds via MEW and was subsequently impregnated with fibrin/alginate (FA).103 Jang et al. reported PCL/hyaluronic acid microspheres that were embedding into a tissue defect using an *in situ* gelling alginate hydrogel. ¹⁰⁴

Inspired by bioactive silicates, we have utilized silicon-based synthetic polymers as bioactive additives to form bone tissue scaffolds. While poly(ethylene glycol) (PEG) hydrogels have been evaluated for bone regeneration, they lack innate bioactivity. 117 Thus, star-polydimethylsiloxane methacrylate (PDMS_{star}-MA) and PEG-diacrylate (PEG-DA) macromers were combined to form templated PDMS-PEG hydrogels that exhibited acellular mineralization when exposed to SBF, as well as enhanced osteogenesis of cultured hBMSCs. 105 Greater bioactivity was observed for phosphonated-siloxane PEG hydrogels prepared with a poly(diethyl(2-(propylthio)-ethyl)phosphonate methylsiloxane)-diacrylate (PPMS-DA) macromer. 106 Finally, we sought to induce bioactivity to our previously reported PCL SMP scaffolds.³⁹ In the first study, PDMS-dimethacrylate (DMA) was combined with PCL-DA at varying wt% ratios (90:10, 75:25, and 60:40), giving rise to PCL-PDMS scaffolds. 107 These maintained shape memory behavior, but displayed acellular mineralization with SBF exposure, as well as enhanced degradation rates due to phase separation effects. In a subsequent study, towards the goal of enhancing bioactivity, polymethylhydrosiloxane-dimethacrylate (PMHS-DMA) was utilized to form analogous PCL-PMHS scaffolds. 108 The increased hydrophilicity of PMHS versus PDMS, stemming from the capacity of silane (Si-H) groups to form dihydrogen bonding with hydroxyl (-OH) groups of water, was expected to better parallel hydrophilic bioactive silicates. Indeed, PCL-PMHS scaffolds exhibited enhanced rates of HAp mineralization, as well as in vitro degradation rates. However, incorporation of a PEG-tethered cell adhesive peptide to PCL-PMHS scaffolds resulted in less presentation at the surface, reducing cellular adhesion.

For the future development of bioactive materials, the combination of high throughput screening (HTS)^{118,119} along with artificial intelligence (AI), 120 specifically machine learning (ML), 121 has great potential. ML used to analyze data from HTS assays can be critical in predicting properties of regenerative scaffolds, with different material combinations. For HTS and ML to be effective toward generating and predicting attributes of bioactive scaffolds, there must be standardized protocols for testing, which is currently lacking. 122 In addition to selecting optimal material combinations, AI can be helpful in determining suitable 3D printing fabrication techniques and scaffold structure depending on the patient and location of the bone

defect. Integrating AI with computer-aided design (CAD) to 3D print patient-specific scaffolds, especially with complex geometries, can reduce the time to generate scaffold structures and facilitate the selection of scaffold parameters (e.g., pore size, percent porosity, strut size) to mimic the adjacent trabecular or cortical bone. For example, after an imaging modality (e.g., CT scan) is used to identify the exact geometry of the defect, the AI integrated CAD system could generate a 3D scaffold that perfectly fits into the bone defect to enable proper osseointegration and angiogenesis through the scaffold. 123 Furthermore, the use of AI techniques to determine optimal scaffold parameters during fabrication can pave the way for more thorough in vitro and in vivo assessments of the most promising scaffold compositions.

Assessment of scaffold of bioactivity

A number of analyses have been used to assess scaffold bioactivity, namely via acellular HAp formation, in vitro cellular behavior, and in vivo bone formation (Fig. 4). Assessment of acellular bioactivity should also consider scaffold sterilization and any pre-treatments (e.g., pre-wetting with ethanol graded baths) to be used for subsequent in vitro cell culture or in vivo assessment.

3.1. In vitro (acellular) assessment of scaffold bioactivity

Immersion of scaffolds to induce mineralization. SBF has been widely adopted to measure bioactivity of materials, including per ISO 23317.124 Exposure to SBF, which is acellular and protein-free, is frequently utilized to confirm scaffold bioactivity in terms of HAp formation. 14,44,125 Developed by Kokubo in 1991, 126 SBF mimics the inorganic composition of human plasma (e.g., Mg²⁺, Ca²⁺, Na⁺, and K⁺) with a physiological pH (\sim 7.4), and is used at body temperature (\sim 37 °C). Kokubo *et al.* suggested the SBF volume (in mL) be greater than 1/10 the surface area of a porous material (in mm²).⁴⁴ As a means to accelerate mineralization, SBF of higher concentrations have been utilized.85 However, highly concentrated SBF solutions can produce uneven, localized precipitation onto surfaces, and also exhibit spontaneous precipitation. This may be somewhat mitigated by increasing the temperature 127 or decreasing pH of concentrated SBF solution. Due to the labor-intensive preparation of SBF, alternatives have been explored. Dulbecco's modified Eagle medium (DMEM), a commercially available cell culture medium, possesses ion concentrations similar to that of human blood plasma.¹²⁹ α-TCP, β-TCP, and HAp were each incubated in DMEM (37 °C, 5% CO₂) for 4 days, leading to CaP precipitate on these surfaces. However, it was noted that the presence of serum can decrease the rate of precipitation.

Scanning electron microscopy (SEM)/energy-dispersive X-ray spectroscopy (EDS). SEM/EDS may be used in conjunction to evaluate the mineralized surface of bioactive scaffolds, including after immersion in SBF. SEM is frequently utilized to provide images of scaffold morphology and microstructure. 130 Coupling with EDS affords determination of elemental

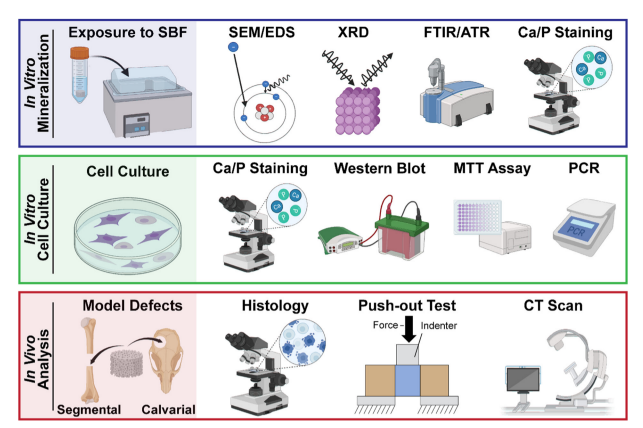


Fig. 4 Generalized methods for assessing scaffold bioactivity: (top) acellular mineralization via exposure to SBF, characterization of HAp mineralization, (middle) cell culture, and (bottom) in vivo analyses using bone defect models.

composition of the surface. ¹³¹ Briefly, when the surface is penetrated by the electron beam, element-specific X-rays are emitted and can be quantified by the spectrometer. SEM/EDS can thus determine the Ca to P molar ratio of a surface, which is ~ 1.67 in the case of HAp. ¹³²

X-ray diffraction (XRD). Another tool used to confirm HAp mineralization on scaffolds is XRD based on the characteristic diffraction signature. Briefly, incident X-rays irradiate the surface and the intensities and scattering angles of the emitted X-rays are measured. XRD can determine HAp phase composition, degree of crystallinity, and crystallite size. 134,138,139

Vibrational spectroscopy. Attenuated total reflectance-Fourier transform infrared (ATR-FTIR) spectroscopy is also commonly used to evaluate HAp mineralization. 140,141 Briefly. molecular functional groups are revealed by nature of their distinct IR absorption band. 142 The ATR mode configuration acquires molecular vibrations through a reduced path length of the probing IR beam, allowing evaluation of the surface to a depth of a few micrometers. ATR-FTIR can be used to identify PO_4^{3-} [~560 and 600 cm⁻¹ and ~1000-1100 cm⁻¹], CO_3^{2-} [between ~ 1460 and $\sim 1530~\text{cm}^{-1}$], and OH^- [$\sim 3570~\text{cm}^{-1}$] groups present in HAp. 138,143 HAp formation, size, and distribution may be assessed with FTIR imaging via micro-ATR-IR. 144 Raman spectroscopy has also been used to evaluate HAp deposits onto scaffolds via the inelastic scattering of light, and water produces less interference versus IR spectroscopy. 145,146

Staining. Staining techniques can be used to evaluate acellular HAp mineralization of bioactive scaffolds. These methods generally involve incubation of the mineralized specimen in an aqueous staining solution, followed by fluorescent imaging and analysis (*e.g.*, with ImageJ software) to yield mineral intensity or mineralized area (% coverage). While advantageously rapid, most stains lack specificity to HAp mineral deposits *versus* calcium- and phosphate-containing deposits and so cannot be used alone to identify HAp.

Alizarin red S, [3,4-dihydroxy-9,10-dioxo-9,10-dihydroanth-racene-2-sulfonic acid; $C_{14}H_7NaO_7S$; "AzHNa"] is a water-soluble sodium salt of Alizarin sulfonic acid that undergoes a reaction with HAp, described as follows: ¹⁴⁹

$$\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2 + 10\text{AzHNa} \rightarrow 10\text{CaAz}\downarrow + \text{HPO}_4^{\ 2-} + 2\text{H}_2\text{PO}_4^{\ -} + 10\text{Na}^+ + 2\text{H}_2\text{O}$$

The red/orange colored precipitate can be quantified, permitting the relative extent of HAp mineralization to be compared across specimens. Per Gregory *et al.*, the precipitate can be removed *via* acetic acid extraction, neutralized with ammonium hydroxide, and absorbance intensity evaluated by a scanning spectrometer at 405 nm. ¹⁵⁰ Alizarin red S staining is not specific to HAp, and will likewise produce such deposits from other sources of Ca²⁺ ions. ¹⁵¹

Von Kossa staining may also be used to evaluate HAp mineralization on scaffolds. Von Kossa staining utilizes a silver

nitrate solution to transform calcium phosphate salts to silver phosphate salts, described as follows: 152

$$Ca_3(PO_4)_2 + 6AgNO_3 \rightarrow 3Ca(NO_3)_2 + \downarrow 2Ag_3PO_4$$

The grey/black precipitate can be quantified to afford comparison of scaffold specimen HAp mineralization. However, this stain is not specific to phosphates of HAp, and cannot be used to provide absolute identification of HAp. 153

Other dyes that can stain HAp mineral deposits on scaffolds include xylenol orange and calcein blue. 154,155 Xylenol orange [3,3'bis[N,N-bis(carboxymethyl)amino-methyl]-o-cresolsulf-onephthalein tetrasodium salt; C31H28N2Na4O13S] forms orange, fluorescent complexes with divalent metal ions (e.g., Ca²⁺). ¹⁵⁶ Calcein blue [4-methylumbelliferone-8-methyliminodiacetic acid] also binds to calcium to afford blue staining of the mineral. 157

The OsteoImage[™] mineralization assay (Lonza) is based on a fluorescent stain that is advantageously specific to HAp mineralization. Thus, OsteoImage™ has been used to stain HAp deposits formed via acellular mineralization 158,159 as well as following cell culture. 160-162

3.2. In vitro (cellular) culture to assess scaffold bioactivity

The bioactivity of scaffolds may also be assessed via cell culture. 163 A variety of cell sources have been implemented, namely stem cells such as bone marrow mesenchymal stems cells (BMSCs), adipose-derived MSCs (ASCs), perivascular stem cells, and induced pluripotent stem cells (iPSCs). 164 Such osteoprogenitor cells undergo osteogenic differentiation into osteoblasts, osteoclasts, and osteocytes, all cells found in native bone tissue. 165 Originating from the bone marrow, BMSCs have been particularly utilized in bone regeneration strategies. 166 Biomolecules, such as the fibronectin-derived peptide sequence Arg-Gly-Asp-Ser (RGDS), are often incorporated into the scaffold to direct or support desired stem cell adhesion, spreading, and differentiation.91 Exogenous (external) growth factors, particularly BMP-2, have also been incorporated into scaffolds to accelerate osteogenesis, although these strategies often risk off-target effects in vivo. 32 A wide range of methods have been used for biomolecule incorporation, as a single method of biomolecule incorporation is not necessarily universally effective for all scaffold types. For instance, phase separation within the scaffold can alter incorporated biomolecule surface presentation.¹⁰⁸ Fluorescent imaging can be used to confirm the quality of cellular adhesion and spreading, whereby fixed cells are stained with phalloidin (cellular actin cytoskeleton) and DAPI [4',6-diamidino-2-phenylindole] (cell nuclei). 167 Prior to the aforementioned analyses, the cytocompatibility of the scaffold should be confirmed using a variety of assays (e.g., MTT assay, ¹⁶⁸ or lactate dehydrogenase [LDH] assay ¹⁶⁹).

Scaffolds that demonstrate cytocompatibility and the capacity to support cell adhesion and spreading, are often then evaluated for their capacity to support the osteogenic differentiation of adherent stem cells. In these studies, osteogenic medium - typically prepared by supplementing a conventional medium with some combination of L-ascorbic acid, β-glycerophosphate, and dexamethasone - may be utilized to mimic the osteogenic milieu present in bone. 170,171 During osteogenesis, stem cells cultured on bioactive scaffolds will produce HAp deposits, the extent to which can be evaluated using histological methods with some of the previously noted stains (e.g., Alizarin red S and von Kossa^{150,172}). The expression of the numerous osteogenic markers involved in osteogenesis, 171 and methods for assessment have been recently reviewed by Le et al. 173 Briefly, following a defined period of culture, the scaffold homogenates may be subjected to a variety of analyses to detect the expression of mRNA levels of genes (via polymerase chain reaction [PCR]) or expression of proteins (via immunofluorescence staining, western blot, and ELISA assays). Multiple osteogenic markers are typically evaluated as each gives insight into specific aspects of osteogenesis, including stages of progression. These include transcription factors (e.g., RUNX2, Osterix, Msx2), extracellular matrix proteins¹⁷⁴ (e.g. secreted protein acidic and rich cysteine [SPARC], osteopontin [OPN], osteocalcin, collagen 1 α1 chain [COL1A1]), and secreted growth factors¹⁷⁵ (e.g., vascular endothelial growth factor [VEGF], 176 BMP-2, 177 and BMP-4178). ALP, an early marker of osteoblast differentiation, is often quantified as an indicator of scaffold bioactivity. 179,180 Expression of "off-target" markers can also be assessed to delineate the specificity of scaffold bioactivity for osteogenesis. Off-target evaluation often includes assessment of markers for chondrogenic (e.g., SRY-box transcription factor 9 [SOX9] and collagen 2 α1 chain [COL2A1]) and adipogenic (e.g., CCAAT/enhancer binding protein $[\alpha$ -C/EBP- α], and adipocyte fatty acid binding protein [AFABP]) differentiation.

3.3. In vivo and ex vivo methods to assess scaffold bioactivity

A number of in vivo and ex vivo models of bone repair are available for the assessment of bioactive scaffolds. 181 Animal species that have been utilized include rodent, rabbit, dog, sheep, goat, and pig, with each presenting unique advantages and disadvantages. 182 Bone defects are frequently created in calvariae [as confined defects], or in ulnae, tibiae, and femurs [as segmental defects]. The minimum size for a critical defect, wherein spontaneous healing does not occur over a long duration, must be considered. 183,184 However, non-critically sized calvarial defects, permitting two rather than one defect per animal, have been used to assess scaffold osseointegration and neotissue infiltration at the perimeter.³⁹ This approach exemplifies a potential way to commit to the 3Rs (reduce, replace, refine) principle of humane animal research. 185 Recently, models have also been created to assess osteoporotic defect healing, 186,187 and attention has been given to sex-based differences in bone defect healing. 188 Several methods to detect in vivo mineralization of scaffold treated defects have been commonly employed as highlighted below. Tissue preservation is required for most ex vivo analyses, and includes methods such as slow freezing, vitrification, hypothermic preservation, and cryopreservation. 189

In vivo analyses. Numerous methods exist for noninvasive, longitudinal monitoring of scaffold-induced bone

regeneration, including bone mineral density (BMD). 190,191 Micro-computed tomography (micro-CT) is perhaps the most widely utilized, given its relative low cost and efficacy. 192,193 Having a spatial resolution of 50-1 µm, ¹⁹⁴ micro-CT affords 3D evaluation of bone ingrowth and volumetric changes. Micro-CT can also be used to determine BMD (mg HAp cm⁻³). 195 Positron emission tomography (PET) can be used to evaluate longitudinal bone formation using γ -ray emitting tracers, such as a sodium fluoride ([18F]-NaF) that forms fluoroapatite with HAp of new bone tissue. 196 Bone regeneration can also be monitored by single-photon emission computed tomography (SPECT) employing positron-emitting tracers, such as 99m technetium [99mTc-labelled diphosphonates] that are absorbed by HAp. 197 Dual-modality, integrated micro-CT/PET^{198,199} and micro-CT/ SPECT¹⁹⁹ images have been used to assess scaffold-induced bone regeneration. Dual-energy X-ray absorptiometry (DEXA) may also be used to assess BMD. 200-202 Other methods of noninvasive monitoring have been employed to avoid potential tissue damage associated with X-rays. Magnetic resonance imaging (MRI) based on semi-quantitative methods may be employed to overcome low sensitivity to bone. 203 For instance, Ribot et al. developed a 3D anatomic and perfusion MRI protocol to observe scaffold-induced healing of femoral defects in rats, including mineralization and neovascularization.²⁰⁴ Ultrasound imaging, while limited by depth of penetration, may be used to quantify bone regeneration. 205 Optical fluorescence imaging (e.g., IVIS) can be performed on fluorescentlylabelled scaffolds to monitor resorption in vivo. 201

Ex vivo analyses. Endpoint tissue specimens harvested from experimental models, as well as tissue culture specimens, are frequently evaluated with the aforementioned *in vivo* methods. Micro-CT is widely used to view morphological features and HAp mineral deposits, wherein longer scan times are permitted for improved spatial resolution. Environmental SEM is also useful as it retains the natural state of the specimen by excluding the need for high vacuum conditions, as well as specimens that are clean, dry, and electrically conductive. Other examples demonstrate the use of *ex vivo* MRI, DEXA, and ultrasound in the evaluation of endpoint tissue specimens. Raman spectroscopy is also useful to evaluate the chemical properties of regenerated bone tissue at the nanoscale, including the degree of mineralization.

Histological and histomorphometric analyses. Histological and histomorphometric analyses of regenerated, mineralized bone tissue is crucial in assessing the bioactivity of scaffolds. Bone histomorphometry provides quantitative evaluation through the use of digitized histological images, ^{212,213} using various image analysis platforms. ^{214,215} Typically, harvested specimens are sequentially fixed, decalcified, dehydrated, embedded (*e.g.*, in paraffin or PMMA), and sectioned. A variety of stains are available for these analyses, including some mentioned previously to detect acellular mineralization of scaffolds. Hematoxylin & eosin (H&E) staining – which stains cell nuclei a dark blue/purple color and basic proteins in the ECM a pink/orange color^{216,217} – is useful to identify woven bone, an early stage of bone development characterized by

random collagen matrix deposition.^{39,218} Masson's trichrome staining employs acid-base chemistry by using 3 dyes to selectively stain tissue components.²¹⁹ When staining for bone regeneration, new bone, collagen, and osteoids are stained blue, while mature bone is stained red.²²⁰ Von Kossa staining can also be used to differentiate mineralized [stained black] *versus* unmineralized [stained red] bone matrix produced by bioactive scaffolds.²¹³

Mechanical testing. A variety of biomechanical tests are utilized to evaluate the efficacy of scaffolds to promote bone regeneration in experimental models. 221,222 In macroscopic assessments, harvested constructs are frequently subjected to quasi-static tests wherein stress or strain is applied in different modes (e.g., compression, tension, bending, torsion, and shear). Bulk modulus, strength, and toughness values can then be determined. In some cases, standards are applied such as ISO 604²²³⁻²²⁵ and ISO 5833.^{226,227} Push-out tests are also frequently employed to give insight into scaffold osteointegration with surrounding tissues, and efforts continue to be made to refine best practices.²²⁸ Microscopic biomechanical analyses are also utilized to give insight into nanoscale mechanical properties.²²² Nanoindentation can be utilized to measure enhanced local hardness imparted by mineralized bone tissue.²¹¹ Atomic force microscopy (AFM), using contact or tapping modes, can be used to reveal nanoscale features (e.g., collagen fibrils and HAp crystals), while the nanoindentation mode is useful for nanomechanical modulus mapping. 229-231 Sub-resonance AFM (e.g., PeakForce Tapping mode) has also been developed for nanomechanical mapping²³² and was used by Zhou et al. to evaluate bone tissue submerged in an aqueous environment.233 It was also shown that micro-CT has also been coupled with mechanical testing to measure contact area and 3D full-field strain in bone/dental implant constructs; 234 such a method could likewise be highly informative to the assessment of bioactive scaffolds.

4. Conclusions

Bioactive scaffolds remain a contemporary approach to bone regenerative engineering. Emerging in recent years are an array of methods to induce bioactivity to polymeric scaffolds that utilize bioceramic fillers, coatings and surface treatments, and additives. Bioactive composite scaffolds continue to be formed with traditional bioceramic fillers (e.g., DBM and BGs), including combinations of two or more types. Newer bioceramics have also emerged (e.g., LAPONITE® and eggshell microparticles). Bioactive coatings applied to scaffolds include deposited bioceramics, bioceramics embedded in a polymer matrix, and polymer-only types. Surface treatments such as plasma treatment, PECVD, and iCVD have also been leveraged to induce bioactivity. Finally, bioactive additives have been combined with 'bioinert' polymers to form bioactive scaffolds. Such additives include primarily certain natural polymers (e.g., chitosan and collagen), as well as silicon- and phosphonated/ silicon-based synthetic polymers. Overall, these methods yield

bioactivity throughout the scaffold (via fillers and additives) or at the surface of the scaffold (via coatings and surface treatments). For the former types, bulk properties (e.g., stiffness and degradation) are impacted which may or may not be desirable. With surface modification, bulk properties are retained, but bioactivity can be expected to be diminished when the surface is lost. High throughput screening and machine learning will be critical for efficient and successful future development of bioactive scaffolds, but necessitates standardized characterization methods. AI integrated CAD also has potential to play a pivotal role in the treatment of patient-specific complex bone defects. A plethora of in vitro (acellular and cellular), in vivo, and ex vivo methods exist to evaluate scaffold mineralization and other aspects of bone regeneration. Perhaps the most common initial assessment of scaffold bioactivity is the capacity to undergo HAp mineralization when exposed to SBF. HAp can be subsequently assessed with numerous methods (e.g., imaging, spectroscopy, and staining). Mineralization by cultured cells, particularly MSCs, as well as evaluation of other markers of osteogenesis is also commonly used to assess scaffold bioactivity. Such analyses typically rely on staining techniques and other assays. In vivo models of bone repair afford an opportunity to assess scaffold bioactivity in a physiological environment. Several assessment methods (e.g., PET, SPECT, and MRI) afford longitudinal monitoring of mineralization and tissue regeneration, with micro-CT being the most frequently employed. Harvested endpoint tissue specimens typically undergo histological and histomorphometric analyses, and biomechanical assessments are also valuable. While recent studies highlight the breadth of methods to prepare and assess bioactive scaffolds, several primary challenges remain for bioactive scaffold-based approaches to displace the clinical use of standard biological and alloplastic grafting. Overall, comparison of scaffold bioactivity in the literature is difficult. Studies employ different analyses (e.g., methods, test conditions, and selected time points) to assess bioactivity both in vitro and in vivo. Thus, standardized methods would be extremely useful to the field. Controls that could be uniformly included along with experimental scaffolds would also be beneficial to studies, but are currently lacking.²³⁵ Explant cultures (a.k.a. organ or ex vivo cultures), wherein explanted tissue-scaffold constructs are maintained in vitro and often with applied mechanical loading, may provide a valuable intermediate step between in vitro cell culture and in vivo experimental models.²³⁶ Despite these challenges, bioactive scaffolds hold tremendous promise in the treatment of bone defects.

Author contributions

Brandon Nitschke: writing - conceptualization, visualization, writing - original draft, writing - review & editing; Felipe Beltran: conceptualization, writing - original draft, writing review & editing; Mariah Hahn: writing - review & editing; Melissa Grunlan: writing - original draft, writing - review & editing.

Conflicts of interest

There are no conflicts to declare.

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