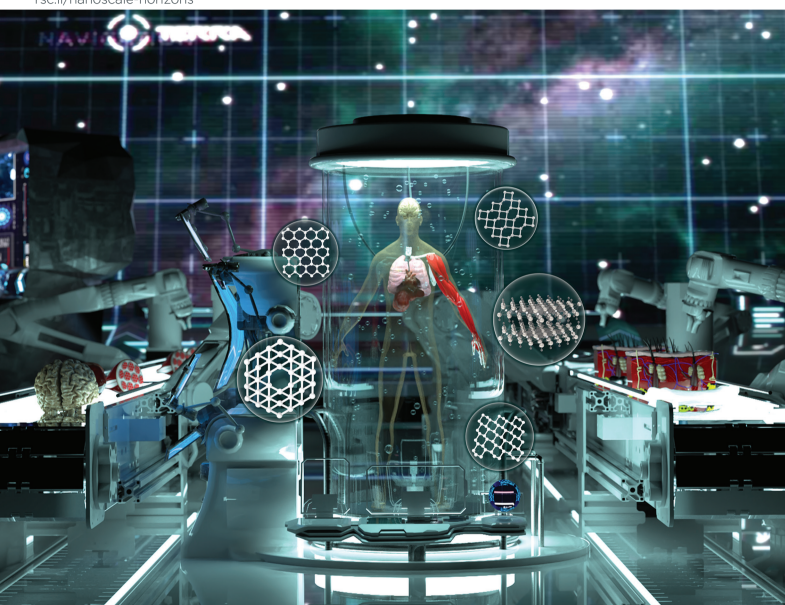
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MXene and Xene: promising frontier beyond graphene in tissue engineering and regenerative medicine

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The emergence of 2D nanomaterials (2D NMs), which was initiated by the isolation of graphene (G) in 2004, revolutionized various biomedical applications, including bioimaging and -sensing, drug delivery, and tissue engineering, owing to their unique physicochemical and biological properties. Building on the success of G, a novel class of monoelemental 2D NMs, known as Xenes, has recently emerged, offering distinct advantages in the fields of tissue engineering and regenerative medicine. In this review, we focus on the comparison of G and Xene materials for use in fabricating tissue engineering scaffolds. After a brief introduction to the basic physicochemical properties of these materials, recent representative studies are classified in terms of the engineered tissue, i.e., bone, cartilage, neural, muscle, and skin tissues. We analyze several methods of improving the clinical potential of Xene-laden scaffolds using state-of-the-art fabrication technologies and innovative biomaterials. Despite the considerable advantages of Xene materials, critical concerns, such as biocompatibility, biodistribution and regulatory challenges, should be considered. This review and collaborative efforts should advance the field of Xene-based tissue engineering and enable innovative, effective solutions for use in future tissue regeneration.

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Introduction

Recently, interest in 2D nanomaterials (2D NMs) has increased owing to their potential applicability in a wide range of biomedical fields.1 It began when André Geim and Kostya Novoselov successfully isolated graphene (G) in 2004,² and subsequently, derivatives of G, such as G oxide (GO) and reduced GO (rGO), with numerous functional groups and lattice defects, were introduced.3-5 The G derivatives have garnered significant attention in the biomedical field, particularly in bioimaging,^{6,7} -sensing,^{8,9} and -robotics,¹⁴⁻¹⁶ theranostics,^{10,11} drug delivery, 12 and tissue engineering, 13 owing to their special physicochemical and biological properties. The application of G in the fields of tissue engineering and regenerative medicine has enabled the tailoring of the biological and mechanical properties of native materials by introducing binding sites for further bio-functionalization with biological molecules.

Notably, extensive research on G enabled the development of other functional 2D NMs. The remarkable progress reported with respect to G over the past few decades has led to novel categories of functional 2D NMs, i.e., transition metal dichalcogenides, transition metal oxides, layered double hydroxides, hexagonal BN, and metal-organic frameworks.²⁰ MXenes, which are 2D NMs comprising transition metal carbides and nitrides, were first discovered by Gogotsi et al. in 2011.²¹ They exhibit remarkable features in converting photothermal energy, rendering them well-suited for applications in photonic hyperthermia treatment within the second near-infrared (NIR) biowindow (NIR-II biowindow), facilitating deep tissue penetration.^{22,23} The range of metals available for use within MXenes renders them useful contrast agents in computed tomography (CT) and magnetic resonance imaging. The remarkable electrical conductivity and numerous functional groups of MXenes, which are highly promising for use in the fields of tissue engineering and regenerative medicine, merit attention in this regard.24-27

Subsequently, a novel category of monoelemental 2D materials emerged, known as Xenes. The component elements

Additional properties, such as conductivity for regulating cell behavior, proliferation, and differentiation, which promote specific tissue regeneration, may also be introduced. 17-19

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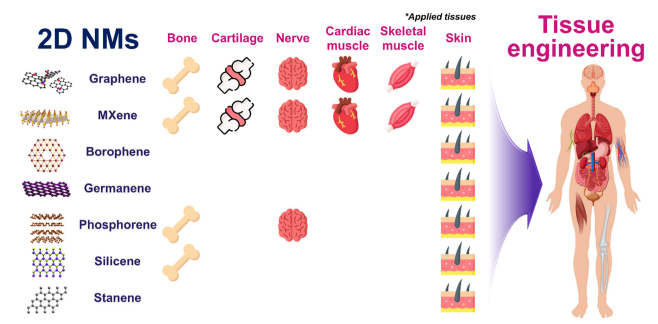


Fig. 1 Schematic of Xene materials for use in various tissue engineering and regeneration

primarily occur in the main groups of the periodic table, i.e., IIIA (B, Al, Ga, and In), IVA (Si, Ge, Sn, and Pd), VA (P, As, Sb, and Bi), and VIA (Se and Te).28 Following a nomenclature akin to that of G, these monoelemental 2D materials are typically named by combining the elemental names with the suffix "-ene," yielding names such as borophene, germanene, and tellurene. Xenes exhibit rapid responses to external stimuli (e.g., NIR laser irradiation or pH changes) owing to their ultrathin 2D structures.²⁹ This enables the triggered and/or controlled release of loaded molecules, rendering them suitable for use in the targeted delivery of multi-responsive therapeutics to desired sites.30 Additionally, the chemical properties of Xenes, including their tunable, versatile surface chemistries, enable the adsorption of various biomolecules, proteins, and cyto-/chemokines, facilitating regenerative medicinal applications.^{31–33}

In this review, we explore the emerging era of MXenes and other types of Xene materials that display potential for use in the fields of tissue engineering and regenerative medicine (Fig. 1). We analyze the distinctive attributes of Xenes, which surpass those of G, and their potential applications in engineering and restoring various tissues, including bone, cartilage, nerves, muscles, and skin. Moreover, we address the opportunities and challenges associated with the utilization of 2D NMs beyond G in the context of the future clinical and industrial applications of Xenes.

Classification and physicochemical characteristics of Xenes

This section provides a concise overview of commonly employed Xenes, including their synthetic techniques and typical features. Based on the positions of their elements in the periodic table, Xenes may be classified as follows: Group III (borophene and gallenene), Group IV (silicene, germanene, and stanene), Group V (phosphorene, arsenene, antimonene, and bismuthene), and Group VI (selenene and tellurene).²⁹

G derivatives

G serves as a representative 2D NM and displays numerous structural features characteristic of 2D NMs. Its structure comprises a single layer of C atoms covalently bonded in a flat, regular hexagonal pattern via sp² bonds.³⁴ Conversely, GO exhibits the same hexagonal C atom arrangement but contains sp³ C atoms bonded to functional groups above or below the plane of the NM, resulting in a rougher structure, accompanied by pronounced local polarity. GO is a non-stoichiometric compound of C, O, and H in a variable ratio, which is influenced by the processing methods used. It contains abundant O-containing functional groups, including epoxide, carbonyl, hydroxyl, and phenol groups, which are introduced during chemical exfoliation.35 These defects alter the inert G structure, resulting in unique properties that enable diverse applications as sensors, photovoltaics, membranes, purification materials, etc. 36 rGO lies in between G and GO in terms of structure. It may be produced by reducing GO via electrochemical, microwave, and photo-assisted thermal methods, which eliminate most functional groups and partially restore sp2 hybridization.37,38 rGO exhibits excellent light absorption characteristics across the entire spectrum, with even a single layer displaying the capacity to absorb a substantial amount of light at visible and NIR wavelengths.39

MXenes

MXenes are synthesized by selectively removing specific atomic layers from parent MAX materials using HF or a combination of

a potent acid and fluoride salt. 40 Their structural formula is $M_{(n+1)}X_nT_x$, with M representing early transition metals, such as Ti, V, Zr, and Nb, X representing C and/or N, and T representing surface functional groups, such as -O, -OH, and -F. Varying the atomic layer number in the unit cell yields the typical structures $M_{2X}T_x$, $M_3X_2T_x$, and $M_4X_3T_x$. Initially, MXenes were investigated for use in energy conversion and storage systems due to their high theoretical capacity and electrical conductivity, but recently they have received significant attention in the field of photocatalysis. 42 Their exceptional physicochemical characteristics may be ascribed to several factors. Wet chemical etching generates an abundance of functional groups, facilitating close contact between the MXene and biological system or material. 43 Adjustments in surface chemistry enable the tuning of the bandgap alignment of the MXene. The conductive metal cores within the layered structure endow the MXene with excellent metallic conductivity and electron-accepting capacity. Consequently, MXenes emerged as strong candidates among 2D materials, and they have been thoroughly explored for use in various photocatalytic applications, including H2O splitting, CO₂ reduction, pollutant degradation, and N₂ fixation. 42 Additionally, the simple surface modification of an MXene improves the in vivo performance by reducing toxicity, enhancing colloidal stability, and prolonging circulation within the body.⁴⁴

Group III Xene (borophene)

B is one of the most chemically complex elements due to its trivalent electronic configuration.⁴⁵ This complexity disrupts the octet rule, resulting in an unusual electron-poor bonding pattern, where the electrons of B and its compounds are shared among ≥ 3 atoms. ⁴⁶ In its bulk form, pure B exhibits significant structural diversity, with 5-16 different polymorphs with highly complex unit cells. 47 Borophene, which is a unique 2D B sheet, displays remarkable properties, including a tunable anisotropic structure, metallic behavior, optical transparency, and potential high-temperature superconductivity (10-20 K). 48-50 Various synthetic methods, such as physical vapor deposition (PVD), mechanical cleavage, etching, and liquidphase exfoliation, have been successfully employed in producing borophene. 51,52

Group IV Xenes (silicene, germanene, and stanene)

Silicene displays a non-planar buckled honeycomb configuration in its monolayer form, which is distinct from the sp²hybridized C atoms within bulk graphite.53 It should exhibit G-like Dirac fermions and semi-metallic properties, along with enhanced spin-orbit coupling effects due to its lower symmetry. 54,55 Theoretical insights suggest that silicene may exhibit topologically nontrivial electronic states, gate-tunable bandgaps, and spin-polarized edge states, rendering it suitable for use in devices such as tunable transistors and photodetectors.⁵⁶ Experimental synthetic methods, such as PVD and chemical exfoliation, have been successfully employed in producing silicene. 57 Germanene, akin to silicene, features a buckled honeycomb structure,⁵⁸ which is characterized by Ge atoms forming a corrugated 2D layer structure, driven by its

enhanced spin-orbit coupling due to its heavier atomic nature compared to that of Si.59 This attribute endows germanene with stronger topological insulator properties. Common methods of synthesizing germanene include mechanical cleavage and PVD.60 Similar to silicene and germanene, stanene displays a buckled honeycomb lattice structure with stable $\pi \cdots \pi$ bonding within the atomic plane. 61 The distinct feature of stanene is its strong spin-orbit coupling, which deviates from the behavior of graphene and results in a bandgap opening of approximately 0.1 eV, with topologically nontrivial states at the edges of the material. 62 Stanene, which is a promising topological insulator, has been experimentally synthesized using PVD.63

Group V Xene (phosphorene)

P exhibits various allotropes, including gaseous P, black P (BP), blue P, white P, violet P, and red P. BP, which is the most thermodynamically stable allotrope, has been studied extensively, particularly its 2D variant, referred to as phosphorene. Phosphorene exhibits an orthorhombic structure characterized by parallel atomic layers that are puckered in a double-floor arrangement. 64 Every P atom within an atomic layer bears 5 valence electrons and engages in covalent bonding with adjacent P atoms. 65 At an elevated pressure, BP displays the capacity to undergo transition to a semi-metallic β phase, which is characterized by a double-layered rhombohedral structure comprising ruffled, interlocked, six-membered rings. 64,66 Due to its versatility, BP is one of the most studied Xenes after G, and various experimental methods have been developed for use in phosphorene synthesis, including mechanical cleavage, 67 liquid-phase exfoliation, 68 etching, 69 chemical vapor deposition, 70 PVD, 71 and wet-chemistry techniques. 72

Tissue regeneration using MXenes and Xenes: comparison with G derivatives

Bone and cartilage tissue engineering using G, MXenes, and Xenes

Bone tissue engineering. In recent decades, G and its derivatives (GO and rGO) have been extensively employed in bone and cartilage tissue engineering (Table 1). rGO, in particular, has residual functional groups that render it more biocompatible than pristine G.73,74 Furthermore, its characteristic structural defects and O-containing functional groups provide sites for potential bioconjugation and interaction with biological molecules, rendering rGO a superior choice for interfacing with biological systems.⁷⁵ Kang et al. studied the fabrication of rGO-coated Ti (rGO-Ti) substrates for use in orthopedic and dental tissue regeneration (Fig. 2a-e).⁷⁶ These substrates were fabricated using meniscus-dragging deposition to introduce a robust, uniform rGO coating layer. The R_0 values of the rGO-Ti substrates were lower than that of intact Ti, indicating significantly flattened surfaces with microscale grooves. An in vitro assay using human mesenchymal stem cells (hMSCs) showed that the rGO-Ti substrates significantly increased cell proliferation after 7 days (d) of incubation.

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Ref.	92 pu	97	Î	8/		86		79 na		of 80		st 81				85			98 uc		87
Outlooks	Osteogenic properties of rGO can be applied for dental and orthopedic bone implants	Potential ex vivo model for critical-sized defects		Viable orthopaedic materiais for clinical application		Customizable structured GBR implant for dental regeneration		Photothermal properties in the NIR-II biowindow offer effective treatment of bone malignancy and osteosarcoma		Biocompatibility, osteoinductivity, and osseointegration of 80 MXene NPs are highlighted for GBR therapy		Piezoresistive and electroactive properties promote direc	osteogenesis by activating the Ca ²⁺ /CALM signaling	patnway		Enhanced mechanical strength, electrical conductivity, and continuous phosphate ion release for bone resenera-	tion therapy		Supplying extra phosphorus for effective bone regeneration 86		Multifunctional therapeutic BP nanocomposite offering photothermal antibacterial capability and bone regeneration
Enhanced osteogenic markers	ALP Mineralization nodule formation	COL1A2, SOST, ALP, BGLAP, and PHEX	Mineralization nodule formation Stress fiber expression	ALF, OFN, and smads	BV/TV, TbN, TbSp	Bone mineral density BV/TV, Tb.Th, Tb.Sp, and Tb.N		NIR-assisted tumor suppression Neovascularization	Col1, OCN, RUNX2, and OPN BV/TV, BMD, and lowered porosity	ALP, OCN, and OPN	Bone volume fracture, Tb.Th, and Tb.Sp	New bone volume RUNX-2, Col-1, OCN, COL-1, and	ALP	ca /cALM signaling pathway		ALP and OCN	nesion and proliferation CSF-1, GDF10, VEGF-A, SMAD1, CSF3, NOG, and INTB1	Osseointegration in bone defects	Col1, BMP4, and RUNX2	Accelerated new bone formation ALP and mineralization nodule	RUMALL, OCN, and OPN ALP and mineralization nodule formation BMD, BV/TV, and BV New bone formation
Biocompatibility and therapeutic applications	High viability at 4 mg mL ⁻¹ rGO-coated substrates	90% viability at 2 mg mL $^{-1}$		95-100% viabinty at 0.5 W/w%			spreading, and proliferation for 5 d High membrane integrity with no intracellular ROS generation	90% cell viability at 1 mg mL ⁻¹ NIR-radiated photonic hypothermia		Enhanced adhesion, actin filament spreading, and proliferation on free- standing MXene films	No noticeable inflammatory cell infiltration and necrosis	$MXene < 8 \text{ mg mL}^{-1} \text{ was}$	cytocompatible	Machine Supported angiogenesis, upregulation of immune regulatory markers with M2 polarization	ES enhanced neovascularization and macrophage polarization	No cell viability at 0.4 mg mL^{-1} BP	Enhanced adhesion and proliferatior	ES supported osteogenesis and cell growth both <i>in vitro</i> and <i>in vivo</i>	No significant cytotoxicity up to 300 ms mJ.	Supported cell adhesion, spreading, and migration	No cytotoxicity up to 0.1 w/v% BP NIR-assisted anti-bacterial effects
Test species	hMSCs	hMSCs		SD rat pri- mary osteoblasts	New Zeal- and rabbit	MC3T3-E1 cells		Saos-2 cells		MC3T3-E1 cells		BMSCs		KAW 264.7 cells	HUVECs Sprague– Dawley rats	MC3T3-E1	Rabbits		hDPSCs	New Zeal- and	hMSCs Saos-2 cells SD rats
Xene Applications materials Formulations	rGO-Ti substrate	GO-Alg/Gel		PGBC SD re nanocomposites mary		GO-coated PLA film		3DP NBGS lattices		MXene films		MXene/RSF	hydrogels			BP-CNTpega injectable gel	6		BP-GelMA/U- Arg-PFA	hydrogel	BP/GelMA-CaP hydrogel
Xene tions material	rGO	OĐ						${ m Nb_2C}$ MXene		${ m Ti}_3{ m C}_2{ m T}_x$ MXene						BP					
Applica	Bone																				

Fable 1 (continued)

Ref.	93			94					96		
Outlooks	Thermodynamic properties under NIR-II irradiation promoting sequential and multistage bone regeneration			Orchestrating scaffold degradation and bone regeneration 94	in a spatiotemporal manner through osteoimmunomodu- lation for treating large bone defects				Fracture healing by regulating the acidic microenvironment suppressing oxidative stress and facilitating osteo-	genic differentiation	
Enhanced osteogenic markers	ALP and mineralization nodule formation	n OPN, BSP, RUNX2, OXS Lama5, Tgfb3, Coll5a1, Fgf10, Ramp1, Bmp2, Smad3, Bmp6,	wheel, semasa, and smadlenew bone formation with enhanced BMD, BV/TV, and Tb.N	SP7, BMP2, SPP1, and ALP	TGFβ and BMP signaling pathway	n Mineralization nodule formation	New bone formation with less	fibrous encapsulation and more osseous tissue accumulation	SOD1, OCN, OPN, and OSX	ALP activity and mineralization	Osseointegration with enhanced BMD and BV/TV
Biocompatibility and therapeutic applications	No cytotoxicity up to 29 μ mol mL ⁻¹	NIR radiation promoted proliferation OPN, BSP, RUNX2, OXS and angiogenesis Ramp1, Bmp2, Smad3, Martin (1998), Smad3, Martin (1998), Smad3, Sman (1998), Smad3, Sman (1998), Smad3, Sman (1998), Smad3, Sman (1998),		ROS scavenging effects	No cytotoxicity up to 50 $\mu g m L^{-1}$	Supported adhesion and proliferation Mineralization nodule formation of BMSCs	Immunoregulatory effects on	Raw264.7 and M2 polarization	ROS scavenging and autophagy activation	MC3T3-E1 No hemotoxicity cells	Supported cell adhesion and proliferation up to 1.2 mg mL^{-1}
Test species	BMSCs	HUVECs SD rats		BMSCs	Raw264.7 cells	SD rats			BMSCs	MC3T3-E1 cells	SD rats
Xene ications materials Formulations	Silicene SNSs@AIPH/ CPC scaffold			H-Si TCP	scaffold				H-Si@HAp-Ti		

Furthermore, the rGO-Ti substrates significantly promoted alkaline phosphatase (ALP) activity and matrix mineralization, which are respective markers of early- and late-stage differentiation. Therefore, these substrates may be effectively utilized as dental and orthopedic bone substitutes due to their superior bioactivities and osteogenic potential and potent effects in stimulating the osteogenic differentiation of hMSCs. Meanwhile, 3D printing methods enable the fabrication of scaffolds with more precise designs, enabling the emulation of the 3D microenvironments of natural extracellular matrices (ECMs). Multiple recent studies have utilized 3D printing to produce 3D scaffolds that encourage encapsulated cells to exhibit morphologies and characteristics closer to those of their *in vivo* states.⁷⁷ Zhang et al. introduced a GO-incorporated alginate (Alg) and gelatin (Gel) composite bioink (GO-Alg/Gel) for use in the 3D bioprinting of bone-mimicking scaffolds. The prepared bioinks with higher GO concentrations (0.5, 1, and 2 mg mL⁻¹) exhibited improved bioprintability, scaffold fidelity, compressive moduli, and cell viability. The bioink supported the osteogenic differentiation of hMSCs, with the upregulation of osteogenic-related gene expression, including ALPL, BGLAP, and PHEX. After 42 d of culture in a bioreactor, the hMSC laden constructs displayed high mineral volumes with enhanced proliferation and osteogenic differentiation. This highlights the potential of the GO-Alg/Gel composite bioink for use in fabricating engineered bone tissues and developing in vitro tissue/organ models for application in ex vivo studies. Tan et al. developed a bioactive GO-functionalized self-expandable hydrophilic osteogenic nanocomposite for use in orthopedic applications.⁷⁸ They added GO to self-expandable P(MMA-AA-St) polymers (PGBC nanocomposites) to reinforce their biomechanics and -activities. The PGBCs exhibited outstanding compressive strengths (>70 MPa) and levels of H₂O absorption and volume expansion, in addition to extended handling times and reduced setting temperatures. Micro-CT was used to analyze the porosity of the PGBCs; the results indicated enhanced levels of bone-PGBC contact and osteogenic capacity. In vitro studies indicated that the cytocompatibility of the PGBCs was superior to that of the commonly used poly(methyl methacrylate) bone cement, based on the cell counting kit-8 assay, live/dead cell staining, and flow cytometry. Osteoblasts extracted from Sprague-Dawley (SD) rats exhibited improved levels of adhesion to the PGBCs, and the levels of expression of ALP, osteopontin (OPN), and Smad5 were upregulated. In vivo studies using rabbits revealed the promising intraoperative handling properties, convenient implantation, and satisfactory biosafety of the PGBCs. Kang et al. used 3D printing to fabricate GO-incorporated polylactic acid (PLA) films for use in guided bone regeneration (GBR). The incorporation of GO into the PLA films significantly increased the film's hydrophilicity and levels of protein adsorption. Moreover, the GO-PLA films supported the viability, migration, and proliferation of MC3T3-E1 preosteoblasts, whereas no significant membrane damage or expression of intracellular reactive oxygen species (ROS) was observed. Micro-CT revealed that, after transplantation into a rat calvarial bone defect model, GO

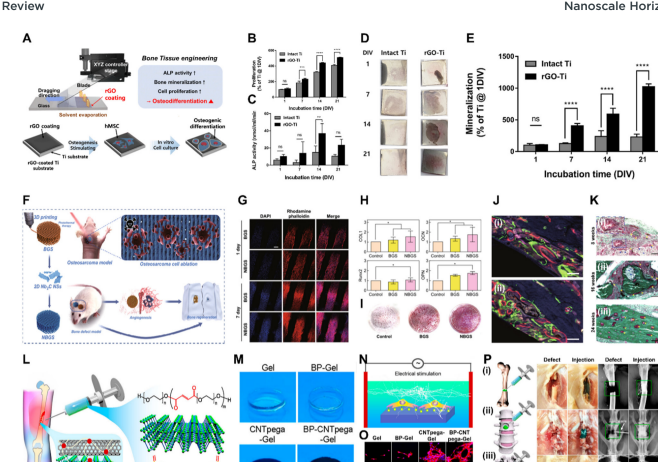


Fig. 2 Xene-based tissue engineering scaffolds for use in bone regeneration. (A)-(E) rGO-coated Ti substrates for use in osteogenic differentiation of hMSCs. (A) Fabrication of the rGO-Ti substrates. (B) Proliferation and (C) ALP activity of hMSCs within 21 d. (D) Digital images and (E) quantified results of the ARS staining of the hMSCs cultured on the rGO-Ti substrate within 21 d. Data reproduced from ref. 76. Copyright Springer Nature 2021. (F)–(K) 3DP NBGS lattices for use in osteosarcoma phototherapy and osteogenesis. (F) Fabrication and study scheme of the 3DP NBGS lattices. (G) Immunofluorescence-stained images of the hBMSCs on BGS/NBGS (blue: nucleus and red: F-actin). (H) Relative expression of osteogenic genes. (I) ARS staining of hBMSCs cultured on different groups. (J) New woven bone around the scaffold at wk 8 at spots (i) 1 and (ii) 2. (K) Goldner trichrome staining of the regenerated tissue in the NBGS group at wks (i) 8, (ii) 16, and (iii) 24. Data reproduced from ref. 79. Copyright Springer Nature 2021. (L)–(P) BP-CNTpega injectable gel for use in bone tissue engineering. (L) Schematic and (M) digital microscopy image of the fabricated BP-CNTpega gel. (N) Schematic and (O) immunofluorescence-stained images of ES-promoted cell proliferation. (P) Capacity of the injectable BP-CNTpega-gels to fill various bone defects. Photograph and X-ray images of rabbit (i) femur and (ii) vertebral body and (iii) posterolateral spinal fusion. Data reproduced from ref. 85. Copyright American Chemical Society 2020. The scale bars represent (J) 250, (G) and (K) 200, and (O) 100 µm, and the asterisks denote statistical significance (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001, and ns: not significant).

significantly improved the bone mineral density and volume fraction and trabecular thickness (Tb.Th), separation, and number (Tb.N), suggesting that the GO-PLA films act as barrier membranes in GBR.

Owing to their biofunctionalities, which are due to their numerous functional groups and elemental nature, MXene nanoparticles (NPs) were recently utilized in bone tissue engineering. Yin et al. developed Nb2C MXene-functionalized scaffolds, which were combined with NIR-radiation osteosarcoma phototherapy in restoring bone defects via angio- and osteogenesis (Fig. 2F-K).⁷⁹ Bioactive glass (BGS) was reinforced with Nb₂C MXenes to fabricate NBGS bioinks. These Nb₂C-MXene nanosheets (NSs) exhibit unique photonic responses in the NIR-II biowindow, rendering them effective in killing bone cancer cells, with deep tissue penetration. Additionally, as the Nb-based components of the Nb₂C MXenes are degraded, they stimulate the growth and migration of blood vessels in the vicinity of the bone defect. This enhances the delivery of O₂, nutrients, and immune cells, accelerating scaffold degradation and providing space for bone remodeling. Furthermore, Ca and PO₄ ³⁻ released during scaffold degradation support the mineralization of new bone tissue. Therefore, the multimodal properties of the NBGS scaffolds may facilitate bone regeneration with anticancer effects, rendering them promising biomaterials for use in treating bone tumors. Zhang et al. assessed the biocompatibility and osteogenic potential of a Ti₃C₂T_x MXene in

in vitro and in vivo settings to elucidate its suitability for use in bone tissue engineering.80 They produced flexible MXene films using the MILD method and conducted in vitro and in vivo studies. In vitro studies using MC3T3-E1 cells revealed that the MXene films significantly enhanced early-stage osteogenic differentiation, as indicated by the increased ALP activity and expression of osteogenic genes. The subcutaneous implantation of the MXene films in rats demonstrated their excellent biocompatibility, with mild inflammatory responses and tissue integration. In a rat calvarial defect model, the MXene films adhered well to the bone tissue, whereas the positive control Ti membranes exhibited reduced mechanical compatibility over time. Micro-CT revealed that the MXene films promoted significant, uniform new bone formation in the defect area, which was guided by the films. Histological analysis confirmed mature bone regeneration on the MXene films without inflammatory reactions. Overall, this study provided valuable insights into the biocompatibility and osteoinductive properties of Ti₃C₂T_r MXenes, thereby supporting their application in bone regeneration. In a study conducted by Hu et al., a Ti₃C₂ MXeneincorporated regenerated silk fibroin (RSF) hydrogel was prepared and used in elucidating the role of electrical stimulation (ES) in bone regeneration.81 ES accelerates the development, specialization, and multiplication of various cell types, including stem cells (SCs).82 RSF, which is sourced from Bombyx mori silkworms, is an extensively studied biopolymer known for its exceptional mechanical strength, biocompatibility, and controllable biodegradability.83 The MXene and ES enhanced osteogenic differentiation, based on increased ALP activity and Alizarin red S (ARS) staining, in addition to the upregulation of osteogenic markers (i.e., runt-related transcription factor 2 (RUNX2), Col-1, osteocalcin (OCN), COL-1, and ALP) at the protein and messenger ribonucleic acid (mRNA) levels. ES also promotes M2 macrophage polarization and indirectly enhances the osteogenic differentiation of BMSCs. The MXene/ RSF hydrogels combined with ES promoted neovascularization in vitro, enhancing the migration and tube formation capacity of human umbilical vein endothelial cells (HUVECs). Additionally, the use of MXene/RSF hydrogels resulted in significant levels of bone regeneration, mineralization, and angiogenesis in cranial defect models, with enhanced levels of M2 macrophage polarization, good biocompatibility, and gradual levels of degradation. Ribonucleic acid (RNA) sequencing indicated that ES upregulated genes related to biomineral tissue development and the Ca signaling pathway, particularly the CALM gene, suggesting that ES-induced osteogenic effects are associated with the activation of Ca²⁺/CALM signaling within BMSCs.

BP is one of the most advantageous biomaterials for use in tissue engineering, because P is involved in various cell signal cascades, in addition to metabolism, membrane and nucleic acid structures, mineralized matrix formation, and enzyme catalysis.84 Liu et al. engineered a BP-infused C nanotubepolyethylene glycol-acrylate (BP-CNTpega) injectable hydrogel for use in repairing irregular tissue defects by enhancing the mechanical strength, electrical conductivity, and PO₄³⁻ release

to support bone tissue regeneration (Fig. 2L-P).85 The BP-CNTpega hydrogel exhibited a PO₄³⁻ release profile with an initial burst release, followed by a slower release due to BP oxidation. Cell viability and proliferation were significantly enhanced within the nanocomposite gel, likely due to the endogenous PO₄³⁻ release and improved mechanical properties. ES was applied to enhance cellular growth and osteogenesis in MC3T3 preosteoblasts, leading to significantly increased levels of cell proliferation, ALP activity, OCN content, and expression of multiple osteogenic genes. The BP-CNTpega hydrogel effectively filled the defect sites in rabbit models, according to X-ray visualization, highlighting its potential for use in diverse bone defect repair applications. In a study conducted by Huang et al., BP was incorporated into a photopolymerizable hydrogel comprising gelatin methacrylamide (GelMA) and cationic arginine-based unsaturated poly(ester amide)s (U-Arg-PEAs), denoted the BP-GelMA/U-Arg-PEA hydrogel.86 The prepared hydrogels exhibited photoresponsive levels of PO₄³⁻ release upon exposure to 808 nm NIR irradiation and enhanced levels of in vitro mineralization and retained superior mechanical properties, even after 15 d of immersion in simulated body fluid. The BP-GelMA/U-Arg-PEA hydrogel stimulated the enhanced mineralization of human dental pulp SCs (hDPSCs). Western blot analysis and the enzyme-linked immunosorbent assay and reverse transcription quantitative polymerase chain reaction (rt-qPCR) confirmed elevated levels of osteogenic markers, such as Col-1, bone morphogenetic protein 4 (BMP4), and RUNX2, within the hDPSCs treated with BPN-containing hydrogels. This suggests that the presence of Ca-free P within these hydrogels was crucial in promoting osteogenic differentiation via the BMP-RUNX2 pathway. In vivo evaluation of rabbit calvarial defects demonstrated enhanced bone regeneration, leading to the formation of mature bone within 12 weeks (wk), as indicated by histological analysis, vascular formation, and the expression of osteogenic markers. Meanwhile, Miao et al. fabricated a BP/GelMA-CaP hydrogel via 3D printing and assessed its in vitro and in vivo osteogenic properties.87 CaP reinforces crosslinked networks and enhances multiple bioactivities of hydrogels.88,89 The BP/GelMA-CaP hydrogel displayed significant photothermal properties under NIR irradiation, leading to a substantial increase in temperature that diminished the activities and viability of Saos-2 osteosarcoma cells and Staphylococcus aureus. In vitro assays using hMSCs revealed significantly upregulated expression of osteogenesis-related genes, including OCN and OPN, and increased ALP activity. Immunofluorescence assays revealed increased expression of the RUNX2, ALP, OCN, and OPN proteins within the BP/GelMA-CaP hydrogel, and increased Ca deposition was observed using ARS staining. In vivo compatibility and therapeutic performance were assessed using a rat cranial defect model, which indicated excellent biocompatibility for subcutaneous implantation and significantly enhanced new bone formation, as revealed by micro-CT, histomorphometric analysis, and hematoxylin and eosin (H&E) staining.

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Si, as the second most abundant element on Earth, is highly regarded for its biocompatibility. It is a key component in various nanomedicines and biomaterials due to its roles in collagen and elastin syntheses and its presence in essential bodily components, such as bone and hair. 90 Silicene exhibits exceptional characteristics, such as a large surface area and high biocompatibility, photothermal conversion in the NIR biowindow, and biodegradability, rendering it an attractive candidate for use in biomedical engineering. 91,92 In this context, Ni et al. engineered Ca₃(PO₄)₂ cement (CPC) containing SiO₂-silicene@2,2'-azobis(2-(2-imidazolin-2-yl) propane) (SNSs@AIPH/CPC), with inherent thermodynamic properties and osteoinductive activity. 93 In vitro studies using HUVECs revealed that SNSs@AIPH/CPC treatment with NIR-II irradiation led to increased cell proliferation, migration, and tube formation and the upregulation of vascular endothelial growth factor (VEGF). In vivo studies revealed denser vascular networks and enhanced VEGF expression around the bone defect sites after NIR-II irradiation. Additionally, SNSs@AIPH/ CPC significantly enhanced the early-stage osteogenic differentiation and late-stage mineralization of BMSCs. Highthroughput RNA sequencing and pathway analysis revealed that SNSs@AIPH upregulated genes associated with the tumor growth factor β and BMP signaling pathways, which are critical in osteogenesis. In vivo studies using a rat cranial defect model indicated that SNSs@AIPH/CPC-NIR-II exhibited outstanding osteoinductive properties, significantly improving the bone mineral density and volume and Tb.N compared to the other groups. Polychrome fluorescent labeling confirmed the strong osteogenic capacity, particularly with NIR-II activation, and immunohistochemical staining supported the enhancement of new bone formation. Lin et al. introduced a hydrogenated-Si NSs (H-Si NSs)-functionalized β-tricalcium phosphate (H-Si TCP) scaffold for use in healing bone defects.⁹⁴ Due to their capacity to scavenge excess ROS, the H-Si TCP scaffolds promoted M2 macrophage polarization, with up- and downregulation of the M2 and M1 markers, respectively. The H-Si TCP scaffolds upregulated the expression of key osteogenic markers, including BMP2, ALP, secreted phosphoprotein 1, and SP7, in the early and late stages of osteogenesis and promoted ALP activity and Ca deposition. This may be attributed to the sustained release of ionic Si, which activates osteogenic signaling pathways and supports biomineralization. In vivo studies revealed that the H-Si TCP scaffolds induced well-coordinated immune responses with reduced fibrous encapsulation, monocyte infiltration, and pathological fibrosis aided by foreignbody giant cells. In vivo, H-Si TCP significantly accelerated osteogenesis by promoting osteoblastic differentiation and activation, as indicated by the increased osteogenic activity and histological evidence of more activated osteoblasts, leading to the enhancement of new bone formation. It also significantly promoted bone remodeling by providing Ca, P, and Si, leading to improved bone repair and mineralization, as indicated by micro-CT and histological analysis of the improved bone volume and density and Tb.Th. Hydroxyapatite (HAp) exhibits the advantages of excellent biocompatibility and the capacity to

integrate with natural bone tissue, rendering it a preferred material for use in bone implants and coatings.95 Yuan et al. anchored 2D H-Si NSs onto HA-coated Ti substrates to develop H-Si@HAp-Ti implants.96 The study investigated the influence of the H-Si@HA coating on BMSCs and MC3T3 cells, highlighting the role of Si ion release from the H-Si NSs in enhancing cell proliferation, osteogenic differentiation, and the expression of osteogenic markers. The results suggest that Si ions are critical in promoting osteogenic processes, including enhanced Ca deposition, activation of Wnt signaling, and ROS scavenging. The study investigated the in vivo osteogenesis and bone fracture-healing properties of H-Si@HA@Ti in a rat tibial fracture model. H-Si@HA@Ti significantly enhanced fracture healing, bone formation and mineral density, and the maximum bending load compared to the other groups. Additionally, the composite coating was biosafe and inhibited osteoclast differentiation and activity with a low attrition rate during implantation, indicating strong binding to the metal substrate. Gene set enrichment analysis showed enrichment in pathways related to Ca channel regulation, hormone binding, and signaling, suggesting the role of H-Si in promoting osteogenesis. Furthermore, a Venn plot identified 90 upregulated mRNAs associated exclusively with H-Si, with hub mRNAs, such as ADAD1 and AQP2, potentially influencing autophagy-related osteogenesis.

Cartilage tissue engineering. The use of G-based materials in tissue engineering is increasing because of their potential as culture substrates for SC and chondrocyte differentiation, in addition to their capacity to activate biomolecules and support chondrogenic differentiation and growth via chemical bonding (Table 2).99 Lee et al. introduced G-BMSC biocomposites to induce chondrogenic differentiation in vitro. 100 Compared to pristine G and GO, pGO displayed a superior long-term viability, which was attributed to its porosity. Additionally, composites containing 25 µg of G and pGO exhibited optimal chondrogenic differentiation. Conversely, higher concentrations led to limitations in differentiation due to decreased cell communication and nutrient exchange, with pGO enhancing differentiation at higher concentrations due to its porous structure. Gene expression analysis corroborated these findings, indicating enhanced chondrogenesis (Col-2, SOX9, and aggrecan (ACAN)) in the pGO and GO composites. Meanwhile, cartilage tissue is required to provide structural support and enable smooth joint motion, and crucial issues in fabricating cartilage tissue engineering scaffolds include biomechanical compatibility, cell integration, and the promotion of chondrogenesis. 101 Zho et al. introduced a cartilage printing bioink containing GelMA, polyethylene glycol diacrylate, and GO. 102 The GO-incorporated scaffolds exhibited significantly higher protein adsorption capacity, promoting cell adhesion and growth. Furthermore, a three-wk chondrogenic differentiation study revealed that GO incorporation led to the increased synthesis of the ECM, including glycosaminoglycans (GAGs) and Col-2, which are crucial in cartilage formation. rt-qPCR analysis showed a significantly higher expression of chondrogenic markers, including Col-2, SOX9, and ACAN, within the

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Table 2 G, MXene, and Xene-based tissue engineering approaches for use in cartilage regeneration

Applications	Xene materials	Formulations	Test species	Biocompatibility and therapeutic applications	Enhanced chondrogenic markers	Outlooks	Ref.		
Cartilage	G and pGO	G-BMSCs biocomposite	BMSCs	50 μg mL ⁻¹ G and pGO did not induce cytotoxicity while 25 μg mL ⁻¹ GO induced long-term toxicity	Col2, Alcian blue, Sox9, and ACAN	Construction of cartilage tissues optimizing G concentrations and porosity	100		
	GO	GelMA- PEGDA-GO hydrogel	BMSCs	0.25 mg mL ⁻¹ GO induced high cell spreading and proliferation	Alcian blue stain Col2, GAG, SOX9, ACAN	Chondrogenic differentiation with abundant ECM production for cartilage regeneration	102		
		ACG/GO scaffold	hADMSCs	1 mg mL ⁻¹ GO enhanced cell viability and proliferation	ACAN, SOX9, Col2	Bioconjugated nanocomposite bioink showcasing excellent printability, cytocompatibility, and chondroinductive capability			
		GO-ACM hydrogel	Rabbit BMSCs SD rats New Zealand	2 mg mL ⁻¹ GO promoted cell adhesion and proliferation Suppressed macrophage activation <i>in vivo</i>		3D acellular cartilage scaffold promising treatment of articular cartilage injuries	106		
	$ ext{Ti}_3 ext{C}_2 ext{T}_x$ MXene	CFOM/PLLA scaffold	white rabbits Mouse BMSCs	NIR-irradiated localized hyperthermia of MXene exhibited antibacterial effects on <i>P. Aeruginosa</i> and <i>S. Aureus</i> MXene did not hinder cell viability and proliferation	in vivo Alcian blue stain ACAN, Col2, SOX9	NIR-triggered antimicrobial properties with photothermal and photodynamic effects for clinical treatment of tracheal injuries	107		

GO-incorporated scaffolds, indicating the enhanced chondrogenic differentiation of the SCs. These findings suggest that GO incorporation improves protein affinity and fosters a favorable environment for chondrogenic differentiation, rendering GO a promising material for use in tissue engineering. Alg, which is a marine-derived biomaterial comprising guluronic acid and mannuronic acid, is a biocompatible, cost-effective, hydrogelforming material frequently utilized in designing bioinks for use in 3D bioprinting, offering versatility in tissue engineering. 103 Olate-Moya et al. synthesized a composite bioink comprising Alg, chitosan (CS), and gel (ACG) reinforced with GO (ACG/GO) (Fig. 3).104 GO significantly improved the proliferation, distribution, and viability of human adiposederived mesenchymal SCs (hADMSCs) on the scaffolds. The study also assessed the chondrogenic differentiation of hADMSCs via the immunofluorescence staining of chondrogenic markers and detected upregulated expression of Col-2, ACAN, and SOX9, indicating the successful induction of chondrogenic differentiation. Notably, the ACG/GO scaffold with a low GO concentration (0.1 mg mL⁻¹) promoted a homogeneous cell distribution and ECM deposition, whereas the one without GO and with a high GO concentration (1 mg mL⁻¹) exhibited high-density cell aggregates. Hence, the presence of GO at low concentrations enhanced long-term cell distribution and ECM formation on the 3D printed scaffold. Meanwhile, a decellularized ECM is a preferred scaffold material because it may replicate the complex composition and topology of a natural ECM, which is distinct for each tissue, due to the intricate interactions between the resident cells microenvironments.105 Gong et al. synthesized a GO-modified 3D acellular cartilage ECM (GO-ACM) as a printable bioink for use in cartilage tissue engineering. 106 In vitro chondrogenesis studies with BMSCs showed that the GO-ACM scaffolds supported cell growth and ECM production, with higher deoxyribonucleic acid and GAG contents, indicating BMSC differentiation to chondrocytes and ECM synthesis. Upon subcutaneous implantation in rats, the GO-ACM scaffolds exhibited good biocompatibility, with mild levels of inflammation at 1 wk post-surgery, which improved by wk 2, with milder inflammatory responses. In the rabbit cartilage defect model, the GO-ACM group exhibited partial scaffold degradation without significant tissue repair at 6 wk post-implantation. At 12 wk post-implantation, the GO-ACM scaffold displayed complete tissue repair with a smooth surface and mature hyaline cartilage, indicating its superior cartilage regeneration potential following in vitro chondrogenic induction.

To date, despite the significant potential of Xene materials in the field of cartilage tissue engineering, only a single relevant study has been presented. Qian et al. developed an NIRtriggered antimicrobial CuFe₂O₄-MXene heterojunction via the formation of CuFe₂O₄ on a Ti₃C₂ MXene, which was then uniformly incorporated into poly(L-lactic acid) (PLLA) and utilized in generating a tracheal scaffold (CFOM/PLLA).107 The NIR-irradiated CFOM/PLLA tracheal scaffolds displayed powerful antibacterial performances, with respective antibacterial rates of 96.49% and 95.33% for S. aureus and Pseudomonas aeruginosa. Additionally, these scaffolds exhibited excellent antibiofilm activities and levels of disruption of bacterial membranes, consumption of glutathione, and generation of ROS, leading to bacterial death. Moreover, the CFOM/PLLA scaffolds released low concentrations of Cu and Fe ions to promote chondrogenic differentiation in BMSCs, as indicated by the increased levels of GAG deposition and the upregulated expression of chondrogenic genes, including ACAN, Col-2, and

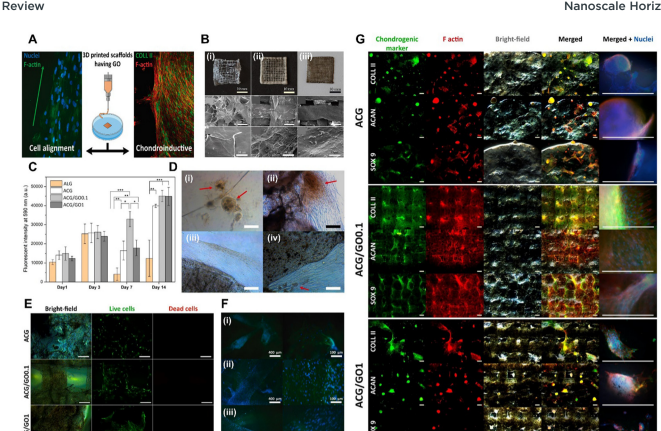


Fig. 3 Xene-based tissue engineering scaffolds for use in cartilage regeneration. (A) Schematic of the chondroinductive ACG/GO scaffold. (B) Digital and SEM images of the lyophilized 3D printed (i) ACG, (ii) ACG/GO0.1, and (iii) ACG/GO1 inks. (C) Proliferation of the hADMSCs within 14 d and (D) optical microscopy images at 7 d on (i) Alg, (ii) ACG, (iii) ACG/GO0.1, and (iv) ACG/GO1. (E) Live/dead assay of the hADMSCs seeded on each scaffold at 7 d. (F) Fluorescence microscopy images of the hADMSCs at 7 d seeded on the (i) ACG, (ii) ACG/GO0.1, and (iii) ACG/GO1 scaffolds (green: F-actin and blue: nucleus). (G) Fluorescence microscopy images of the chondrogenic markers (green), including Col-2, ACAN, and SOX9, within the 3D printed scaffolds: ACG (top), ACG/GO0.1 (middle), and ACG/GO1 (bottom), after 28 d of culture (red: F-actin and blue: nucleus). Data reproduced from ref. 104. Copyright American Chemical Society 2020. The scale bars represent 10 mm, 300 µm and 50 µm for (B) upper, middle, and lower layers, respectively; and (D) 100 μ m; (E) 200 μ m; (F) 400 μ m (left) and 100 μ m (right); and (G) 500 μ m. The asterisks denote statistical significance (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001, and ns: not significant).

SOX9. Thus, the fabricated CFOM/PLLA tracheal scaffolds display potential for use in clinical applications because of their excellent antibacterial and regenerative properties.

Neuromuscular tissue engineering using G, MXenes, and Xenes

Neural tissue engineering. Despite decades of progress in understanding the biological mechanisms of the nervous system, the regeneration and repair of nerve tissues remain challenging. This is primarily because of the complex anatomy and physiology of the nervous system, which distinguishes it from other tissues, with injuries often resulting in lifelong disability. 108 However, various approaches, including autoand allografts and pharmacological agents, have been explored, leading to increased interest in developing novel strategies, such as neural tissue engineering and nanotechnology-based approaches, for use in neural tissue regeneration (Table 3). 109 G materials exhibit hydrophilic properties and morphological characteristics that influence their capacity to interfere with neuronal and astrocyte behaviors. Small GO flakes may

enhance inhibitory synapses in neuronal networks, while modifications of the O species of GO may be used to produce biocompatible substrates to promote cell adhesion, differentiation, and neurite growth (Fig. 4A-E). 110 rGO with a higher C/O ratio displays an enhanced electrical conductivity and hydrophobicity, affecting its interactions with cells and modulating biological processes, such as inflammation and neural cell differentiation.111,112 Typically, nanofiber matrices are widely used in engineering linear tissues, such as nerves and muscles, due to their orientational guidance and capacity to endow cells with topical stimuli-responsiveness. 113 Girao et al. introduced size- and O group-controllable GO-laden gel and polycaprolactone (PCL) composite nanofiber matrices (rGO-PCL/gel) to induce the neurogenic differentiation of embryonic neural progenitor cells (ENMCs). 110 The rGO-PCL/gel scaffolds maintained their structural integrity and displayed distinct mechanical properties, with excellent compressive moduli. Both scaffolds promoted the formation of interconnected neural networks, with cells attaching, growing, and extending neurites

G, MXene, and Xene-based tissue engineering approaches for use in neural tissue regeneration Table 3

Ref.	110	14			.15	17		.18		19				.20			22	
Outlooks	Customized GO to modulate the properties of electrospun nano- fibers for neural tissue engineering	Promising peripheral nerve repair and regeneration by promoting 114 neural cell behaviors <i>in vitro</i> and facilitating nerve repair <i>in vivo</i>			The adECM functionalized with PDA-rGO guiding neural stem cell 115 adhesion, migration, and spontaneous neuronal differentiation	Promoting nerve regeneration, angiogenesis, and the transmission 117 of physiological neural electric signals for peripheral nerve injury	repair	Promoting neural network formation when combined with a low- 118	ing loss through cochlear implantation	Electrical conductivity. surface functionality, and biocompatibility. 119	promoting NSC neurogenesis, and neurite outgrowth, making it a promising substrate for NSC engineering			Interface for regulating NSCs with electrically conductive interface 120 for biosystems and future clinical annications			Neuritogenesis in hippocampal neurons for potential neural tissue 122	engineering and regeneration
Enhanced neurogenic markers	- MAP2, vimentin	- NGF, PMP22, and Krox20	 Neurite length and neurogenic differentiation In vivo nerve regeneration with enhanced nerve conduc- 	tion velocity – Schwann cell population, myelination, and axon diameter – S100, NF200, and GFAP	- Tuj1, GFAP, and MAP2	- Functional and histological regeneration of sciatic nerve	- Gastrochemius muscle regeneration - Myelinated axon - S100, MBP	- Neurite length with growth	- Synapsin-1, PSD95 - Calcium oscillations	 NGS on neurogenic markers Neurite guidance with 	enhanced branch number,	- Nestin, Ki67, GFAP, and Tuj1 - Synaptophysin and synaptic	puncta – PI3K-Akt and Ca-signal	pathway – Nestin and Tuj1 – Neuronal differentiation.	neurite length, branches, and dendrite protrusion	- Synapsin-1, PSD95	- Neurofilament heavy	Polypepude - Neurite length and guidance - OCK, NeuN, MAP2, and
Biocompatibility and therapeutic applications	- 10 w/v% rGO enhanced cell viability	 - 2 mg mL TGO-incorporated matrices with ES enhanced cell viability and proliferation 	1 30 1 2		- 1.5 wt% rGO did not hinder cell viability and proliferation	ting did not roliferation	- No main organ damage	- 300 µg mL ⁻¹ MXene did not hinder -		- 250 ug mL ⁻¹ MXene NPs were		 - Anti-inflammatory reaction with less - Nestin, Ki67, GFAP, and Tuj1 foreign body responses - Synaptophysin and synaptic 		 - MXene-coating did not hinder cell viability. proliferation , and membrane			' was decreased at ≥	c3 kg inc. - 0.4 g BP-laden matrices enhanced - cell proliferation
Test species	- Rat ENPCs	- ES, SCS, and PC12 cells	– SD rats		– NE-4C cells – NSCs	- RSCs	- SD rats	- SGNs	- Wild- type mice	- Mouse	NSCs			- Mouse NSCs			- HT22	193
Xene Test Applications materials Formulations species	PCL/Gel-rGO nanofiber	Ap/PLCL-rGO nanofiber			adECM-PLA/ PCL-rGO nanofiber	MXene-PCL NGCs		MXene-	hydrogel	MXene-PLLA				Lam-MXene film			PLCL/Lam/ - H	Dr manomber
Xene ns materials	rGO					${ m Ti}_3{ m C}_2{ m T}_x$ MXene											BP	
Application	Neural tissues																	

Table 3 (continued)

V	iew		
	Enhanced neurogenic markers Outlooks Ref.	 - 0.3 mg mL⁻¹ BP@PDA with ES did - Nestin, Tuj1, GFAP, and MAP2 Electrical conductivity and neural differentiation of MSCs offering 123 not induce cytotoxicity - Cell spreading and proliferation tissues - ES supported neurogenic differentiation 	 BBB score Spinal cord injury repair by reducing inflammation and promoting 124 neuronal regeneration through the activation of the AKT signaling pathway CSPG, NeuN, MAP2, and GFAP Spinal cord tissue regeneration PDK1/AKT/GSK3 and PDK1/AKT/AKT/GSK3 and PDK1/AKT/AKT/GSK3 and PDK1/AKT/AKT/GSK3 and PDK1/AKT/AKT/GSK3 and PDK1/AKT/AKT/AKT/AKT/AKT/AKT/AKT/AKT/AKT/AKT
	Enhanced 1	- Nestin, Tu	– BBB score – CSPG, NeuN, MAP GFAP – Spinal cord tissue regeneration – PDKI/AKT/GSK3 a AKT/14-3-3/BAD patl
	Biocompatibility and therapeutic applications	 - Rat - 0.3 mg mL⁻¹ BP@PDA with ES did BMSCs not induce cytotoxicity - SD rats - Cell spreading and proliferation were enhanced - ES supported neurogenic differentiation 	Rat pri- – 100 µg mL ⁻¹ BP alleviated inflam- – BBB score nary matory effects, reduced apoptosis, neuron and promoted proliferation SCI rats – 100 µg mL ⁻¹ BP showed no toxicity – CSPG, NeuN, MAP2, and on blood and main organs – Spinal cord tissue regeneration – PDK1/14-3-3/BAD pathways AKT/14-3-3/BAD pathways
	Test ns species	– Rat BMSCs – SD rats	- Rat pri- mary neuron - SCI rats
	Xene Test Biocompatibi Applications materials Formulations species applications	GelMA- BP@PDA nanofiber	E@BP hydrogel

along the fibrous porous structures. The scaffolds resulted in good cell viability, with minimal areas covered by dead cells, indicating no significant release of toxic products. Furthermore, rGO-PCL/gel supported the differentiation of ENMCs with neural branching and the upregulation of microtubuleassociated protein 2 (MAP2) and vimentin. Wang et al. utilized Antheraea pernyi-derived silk fibroin (ApF) to enhance the cellmatrix interactions of the rGO-laden PLCL nanofiber matrices (Ap/PLCL-rGO) for use in peripheral nerve regeneration. 114 The use of Ap/PLCL-rGO, particularly when subjected to ES, led to increased gene expression of neurotrophic factors, including nerve growth factor, peripheral myelin protein 22, and early growth response 2 (Krox20), which are associated with Schwann cell myelination. Simultaneously, the expression of the neural cell adhesion molecule, which is a marker of immature SCs, was decreased. The use of Schwann cell-conditioned media cultured on Ap/PLCL-rGO with ES significantly enhanced PC12 cell differentiation with neurite outgrowth, which was attributed to the neurotrophins secreted by the Schwann cells. Subsequently, nerve guidance conduits (NGCs) were transplanted for peripheral nerve regeneration in a 10 mm sciatic nerve defect rat model, bridging the nerve defects with no signs of inflammation. Tissue weight analysis, Masson's trichrome staining, and electrophysiological studies indicated that functional recovery improved nerve conduction. Immunohistochemical and -fluorescence analyses of the regenerated nerve tissue revealed an increased Schwann cell density, myelin thickness, and axon diameter, along with increased expression of neural markers (S100 and NF200) and reduced expression of glial fibrillary acidic protein (GFAP), which is a glial marker. Meanwhile, da Silva et al. used adipose tissue-derived ECM (adECM) and rGO-reinforced PLA/PCL nanofiber matrices (adECM-PLA/ PCL-rGO) to differentiate NE-4C neuronal cells and neural SCs (NSCs). 115 The adECM-PLA/PCL-rGO matrices provided anchoring sites for the NSCs, leading to the formation of large cell clusters, particularly in 3D constructs. The presence of rGO increased NSC metabolic activity, whereas the 3D environments with interconnected micropores promoted cell migration and spontaneous differentiation into neurons, with noticeable neurite outgrowth. Immunoblot analyses revealed significant increases in the neuronal markers Tuj1, GFAP, and MAP2a/b, indicating enhanced neuronal differentiation and neuritogenesis in the 3D adECM/PLA polydopamine (PDA)-rGO constructs.

Owing to their high electrical conductivity derived from their 2D structures and elemental nature, MXenes have also gained considerable attention in the field of neural tissue engineering.116 Nan et al. developed Ti₃C₂T_x MXene-coated electrospun PCL NGCs to enhance neurite regeneration and angiogenesis (Fig. 4F-L).117 The MXene-PCL NGCs exhibited excellent biocompatibility, with no significant toxicity toward Schwann cells, good levels of cell attachment, and no abnormalities in major organs when postoperatively evaluated in rats at wk 12. The NGCs remained structurally intact but displayed some surface degradation, which did not affect their biocompatibility or levels of nerve regeneration. At 12 wk postimplantation, the MXene-PCL NGCs significantly enhanced

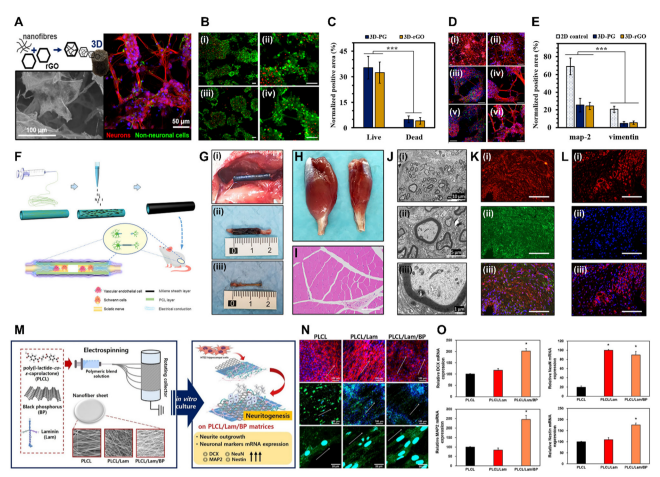


Fig. 4 Xene-based tissue engineering scaffolds for use in neural tissue regeneration. (A)-(E) PCL/Gel-rGO nanofiber scaffolds. (A) Schematic and representative fluorescence images of neurons differentiated on the PCL/Gel-rGO nanofiber scaffolds. (B) Live/dead assay and (C) quantified results of the ENMCs cultured on (i) PCL/Gel and (ii) PCL/Gel-rGO. (D) Immunofluorescence staining and (E) guantified results of the ENMCs cultured on (i) PCL/Gel and (ii) PCL/Gel-rGO (red: MAP2, green: vimentin, and blue: cell nuclei). Data reproduced from ref. 110. Copyright American Chemical Society 2020. (F)-(L) MXene-PCL NGCs. (F) Schematic of the fabrication of MXene-PCL NGCs for use in neural regeneration. (G) Morphologies of the MXene-PCL NGCs and regenerated nerve. (i) Digital image captured at 12 wk after implantation. (ii) MXene-PCL NCGs and (iii) regenerated nerves extracted from SD rats. Representative (H) optical and (I) H&E staining images of the gastrocnemius muscle at 12 wk post-operation. (J) TEM images of the cross-sections of the nerves regenerated using the MXene-PCL conduit: (i) Low-, (ii) medium-, and (iii) high-magnification images. (K) Immunofluorescence staining of S100 (red) and MBP (green) of the MXene-PCL group. (L) Immunofluorescence staining of CD34 (red) and 4',6-diamidino-2-phenylindole (blue) of the MXene-PCL group. Data reproduced from ref. 117. Copyright Frontiers 2022. (M)-(O) PLCL/Lam/BP nanofiber matrices for use in neural tissue engineering. (M) Schematic of the electrospinning of the PLCL/Lam/BP matrices and the enhanced neuritogenesis of the HT22 cells. (N) Immunofluorescence staining of the HT22 cells cultured within each matrix (red: F-actin, blue: nucleus, and green: neurofilament). (O) Relative levels of mRNA expression of DCX, NeuN, MAP2, and nestin. Data reproduced from ref. 122. Copyright Elsevier 2020. The scale bars represent (I) and (K) 200 µm, (A-left, N and L) 100 µm, (Aright, D, B and C) 50 μ m, (J-i) 10 μ m, (J-ii) 5 μ m, and (J-iii) 1 μ m, and the asterisks denote statistical significance (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001, and ns: not significant).

gastrocnemius muscle and muscle mass restoration and increased the muscle fiber diameters. Functional recovery of the sciatic nerve, as assessed using the sciatic functional index and electrophysiological analysis, was excellent when using the MXene-PCL NGCs, and immunohistological and -fluorescence analyses revealed that these NGCs contributed to nerve fiber regeneration and myelination. Additionally, the MXene-PCL group displayed increased neovascularization, as indicated by the higher microvessel density compared to that of the PCL group, with comparable results to those of the autograft group. These findings indicate the potential of the MXene-PCL NGCs for use in promoting nerve regeneration and functional recovery. Meanwhile, Liao et al. introduced an ES setup prepared by integrating a cochlear implant with a conductive Ti₃C₂T_x MXene-matrigel hydrogel, which was utilized to culture spiral ganglion neurons (SGNs) and expose them to ES delivered via the cochlear implant. 118 At concentrations of 40-100 μg mL⁻¹, the MXene-matrigel hydrogel promoted neurite outgrowth and induced significant increases in the growth cone areas and filopodia numbers of the SGNs. Additionally, the cochlear implant-MXene-matrigel hydrogel-electroacoustic stimulation (EAS) system enhanced the neurite development of the SGNs, resulting in longer neurites, larger growth cone areas, and increased filopodia numbers, without inducing

neurotoxicity. The system also promoted the formation of neural networks among the SGNs without affecting the expression or co-localization of synapsin 1 and postsynaptic density protein 95, indicating mature potential synapses. This system also increases the number of SGN synapses and enhances the Ca oscillations within the SGNs, potentially accelerating signal transmission and promoting neural network formation. RNA sequencing analysis revealed differential gene expression between the control and EAS groups, with genes related to ion transmembrane transport, synaptic transmission and plasticity, and cell adhesion significantly affected, suggesting that the system regulates various aspects of SGN behavior and function. Zhu et al. introduced Ti₃C₂T_x MXene-coated PLLA nanofiber (MXene-PLLA) matrices for use in NSC applications. 119 The prepared MXene-PLLA matrices increased the mRNA expression levels of Tuj1 and GFAP in NSCs with the maturation of the NSC-derived neurons and astrocytes (i.e., neurite branching, extended branching length, and increased dendritic complexity indices and synaptic densities). Furthermore, Kyoto Encyclopedia of Genes and Genomes analysis highlighted involvement in neuroactive ligand-receptor interactions and signaling pathways related to NSC self-renewal and Ca modulation. In vivo analysis indicated that the MXene-PLLA matrices exhibited excellent in vivo biocompatibility as they reduced foreign body responses, inflammation, and fibrotic scar formation, while enhancing matrix stability and slowing degradation, rendering them promising for use in nerve tissue engineering. Guo et al. investigated the neurogenic effects of a laminin-coated Ti₃C₂T_x MXene (Lam-MXene) film using HT22 hippocampal neuronal cells. 120 The Lam-MXene film significantly enhanced the differentiation of NSCs to neurons, as indicated by an increased neuron content, extended neurite length, higher numbers of branch points and tips, and an improved expression of neurogenic markers (PSD95 and nestin). However, no discernible effect on synaptic development in the NSC-derived neurons was observed. In Ca imaging studies, the NSCs cultured on the Lam-MXene film exhibited spontaneous Ca spikes with high frequencies and

Meanwhile, BP aids the intercellular interactions and recovery of peripheral nerves due to its high electrical conductivity. 121 Kang et al. developed Lam- and BP-coated PLCL (PLCL/Lam/BP) nanofiber matrices to enhance the neuritogenesis of HT22 hippocampal cells (Fig. 4M-O). 122 Due to the hydrophilic nature of BP and Lam, the fabricated PLCL/Lam/ BP matrices displayed enhanced hydrophilicity and decreased nanofiber diameters, which are favorable for neural cell growth. The levels of cell proliferation on 7 d were significantly increased in the PLCL/Lam/BP matrices, whereas no difference in initial cell adhesion was observed. HT22 cells cultured within the PLCL/Lam/BP matrices exhibited increased neurite length, with clear expression of neurofilament heavy chains. Based on mRNA analysis, doublecortin X (DCX), NeuN, MAP2, and nestin were significantly upregulated within the

synchronization. Moreover, ES supported the Lam-MXene film

by enhancing NSC adhesion, distribution, terminal extension,

PLCL/Lam/BP matrices, indicating the neurogenic potential of the BP-incorporated nanofiber matrix. Xu et al. introduced a PDA-modified and BP-laden GelMA (GelMA-BP@PDA) hydrogel to promote the neural differentiation of rat BMSCs. 123 ES at 100 mV cm⁻¹ was applied to mesenchymal SCs seeded on the GelMA-BP@PDA hydrogels, resulting in increased expression of the early neural markers nestin, Tuj1, MAP2, and GFAP, suggesting that the rat BMSCs were differentiated into neurogenic lineage. Four wks after subcutaneous implantation of the GelMA-BP@PDA hydrogel into SD rats, accelerated in vivo degradation was observed owing to enhanced cell infiltration and the presence of biological enzymes. Additionally, histological staining confirmed the excellent cytocompatibility of the hydrogel, with cells infiltrating and surrounding it over time, suggesting that the prepared GelMA-BP@PDA hydrogel combined with ES was favorable for use in in vitro neural tissue engineering and in vivo applications. Xie et al. fabricated BP quantum dot containing epigallocatechin-3-gallate (E@BP) hydrogels that target the protein kinase B (Akt) signaling pathway in spinal cord injury repair. 124 E@BP significantly improved motor function recovery in SCI rats, as indicated by their higher Basso-Beattie-Bresnahan scores and improved performances in inclined plate studies. Furthermore, E@BP treatment reduced apoptosis and promoted neural regeneration in injured neurons, as indicated by the decrease in the number of caspase-3-positive and propidium iodide/annexin double-positive neurons and the improved mitochondrial morphology. Additionally, E@BP facilitation of neural regeneration was indicated by the increase in the number of 5-ethynyl-2'deoxyuridine-positive neurons and cell cycle progression. E@BP repaired the spinal cord tissue by reducing inflammation and promoting neuronal survival. Finally, the study investigated the molecular mechanisms underlying the effects of E@BP, revealing that it regulates the phosphoinositidedependent kinase-1 (PDK1)/Akt/glycogen synthase kinase 3 and PDK1/Akt/14-3-3/Bcl-2-associated death promoter pathways, promotes cell cycle progression, and reduces apoptosis in SCI rats.

Muscle tissue engineering. Skeletal muscles, which constitute approximately 45% of the human body, are crucial in functions such as support, motion, and metabolic regulation. Skeletal muscle tissue engineering aims to repair or replace damaged skeletal muscles via various fabrication techniques, with applications spanning regenerative medicine, biorobotics and -sensing, energy harvesting, and drug screening. 125 In terms of cardiac muscle tissue engineering, the primary goal is to replicate the natural 3D tissue structure comprising cardiomyocytes (CMs), capillary endothelial and vascular smooth muscle cells, fibroblasts, and macrophages to mimic the intricate intercellular and cell-matrix interactions within the heart.126 G materials have been extensively utilized in skeletal and cardiac muscle tissue engineering in recent decades (Table 4). 127-129 Jo et al. developed nano-GO-incorporated polyurethane (PU) nanofiber matrices (PU-nGO), which were mechanically flexible and myogenesis-inducing, for use in skeletal muscle tissue engineering. 130,131 The PU-nGO

and metabolism.

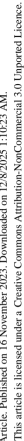
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Table 4 G, MXene, and Xene-based tissue engineering approaches for use in muscle tissue regeneration

Applications	Xene materials	Formulations	Test species	Biocompatibility and therapeutic applications	Enhanced myogenic markers	Outlooks	Ref.
Skeletal muscle	GO	PU-nGO nanofiber	- C2C12 cells	8 w/w% GO promoted cell adhesion, spreading, and proliferation Dynamic tensional stimuli facilitated myogenesis	– MHC, α-actinin, MyoG, and MyoD	Flexibility, mechanical properties, and capacity for stimulating myogenic differentiation for ske- letal muscle engineering	130
		GO/GHPA hydrogel	- C2C12 cells	$-$ 10 μg mL $^{-1}$ GO promoted cell growth and proliferation	MHC and aligned myotube formationMyoD and MyoG	3D printable GO-laden bioink for customizable skeletal muscle- mimetic constructs	132
Cardiac muscle	rGO	rGO-Col scaffold		– 90 $\mu g \; mL^{-1} \; rGO \; enhanced$ viability, spreading, and proliferation	- <i>In vivo</i> host cell migration	Enhanced mechanical properties and ability to upregulate cardiac gene expression	133
			- Rat CMs	- In vitro and in vivo angiogenesis	Cx43, Actn4, andTrpT-2Cardiac muscle regeneration		
	GO	GO-PEI scaffold	– HL-1 cells	– GO coating maintained good viability and adhesion	– Cx43, Nkx 2.5, and intercellular interaction	Biocompatibility and cardiac cell growth with electrical signal con- duction for cardiac tissue engi- neering and <i>in vitro</i> cardiac tissue models	
Skeletal muscle	Ti ₃ C ₂ T _x MXene	– MXene/ AuNP/GelMA hydrogel	- C2C12 cells	$-$ 0.05 mg mL $^{-1}$ MXene induced high cell viability and proliferation	– MHC – Myotube matura- tion indexes	AuNPs and MXene improved printability, conductivity, and cel- lular differentiation, showing potential for muscle tissue engi- neering applications	136
		– MXene-F127 hydrogel		– 80 $\mu \mathrm{g\ mL}^{-1}$ MXene induced high cell viability	- MHC	Antioxidation, anti-inflammation, and angiogenesis properties for skeletal muscle repair by regulat-	140
			- C2C12 cells - HUVECs	Antioxidant and anti- inflammatory effectsAngiogenic effects	Myotube maturation indexesMyoO, MyoG, and MHC	ing cell niche	
			- SD rats		- <i>In vivo</i> regeneration of myofibers and capillary		
Cardiac muscle	Ti ₃ C ₂ T _x MXene	– MXene-PEG hydrogel	- iCMs	– High viability and proliferation on MXene constructs		Clinically relevant cardiac patches for myocardial infarction treat- ment, addressing the need for ordered structure and electro- conductivity in cardiac patches	141
		– MXene-Col film	- C3H10 cells	 60 w/w% MXene induced cell metabolic activity and proliferation ES stimulated maturation 	- CMs count and spreading	Conductive biohybrid platform for cardiac tissue engineering, showing improved electrical con- ductivity and cell growth	142
			- iCMs	and elongation of iCMs - Antibacterial properties on S. aureus	actinin, and Cx43 – action potential	ductivity and ten growth	

nanofibrous membrane displayed distinct myotubular structures and well-arranged actin filaments, along with increased major histocompatibility complex (MHC) expression and upregulated mRNA levels of myogenic markers (i.e., myogenin (MyoG), α-actinin, and myoblast determination protein 1 (MyoD)). Therefore, the PU-nGO nanofibers promoted myogenic differentiation owing to improved cell adhesion and their capacity to mechanically stretch cells under dynamic tensional stimuli. Kang et al. fabricated a 3D bioprintable GO-incorporated functionalized Gel hydrogel (GO@GHPA) for use as a skeletal muscle printing bioink (Fig. 5A-H). The phenol-rich GHPA hydrogel could be crosslinked using glucose oxidase and horseradish peroxide to yield a cytocompatible crosslinked hydrogel, which was then laden with cells. The use of the 3D bioprinted GO@GHPA hydrogel resulted in

enhanced cell viability, distribution, and proliferation over 7 d of culture. Over a prolonged culture period, the laden C2C12 myoblasts exhibited enhanced myotube formation and MHC expression, with the upregulation of MyoD and MyoG, suggesting the myogenic potential of GO and the GO@GHPA hydrogel. Meanwhile, Norahan et al. introduced rGOincorporated Col scaffolds (Col-rGO) for use in accelerating angio- and myogenesis in cardiac tissues. 133 Cytotoxicity studies using the extracts from the Col-rGO scaffolds revealed excellent cytocompatibility, indicating that the degradation products were not harmful to the cells. Scanning electron microscopy (SEM) revealed that all scaffolds supported cell attachment and proliferation. Gene expression analysis at 7 d after seeding rat neonatal CMs on the scaffolds indicated increased levels of expression of cardiac genes, such as Cx43,



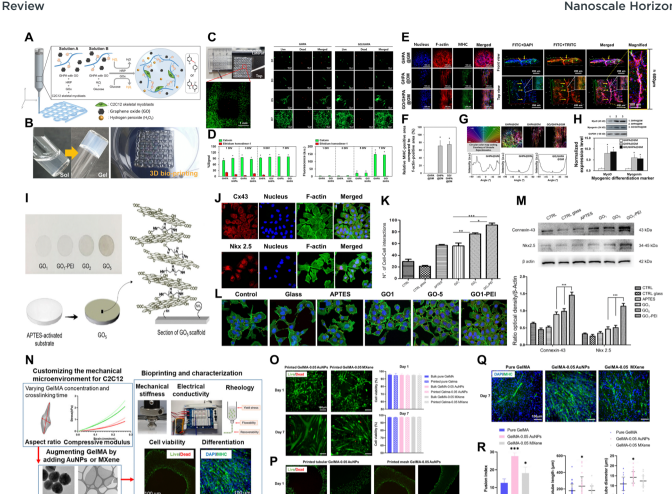


Fig. 5 Xene-based tissue engineering approaches for use in muscle tissue regeneration. (A)-(H) GO@GHPA printable bioink for use in skeletal muscle tissue regeneration. (A) Schematic of the in situ crosslinkable GO@GHPA hydrogel. (B) Digital images of in situ crosslinking and the 3D printed construct. (C) Live/dead assay within 7 d and (D) quantification. (E) Immunofluorescence staining of the C2C12 myoblasts cultured within the GO@GHPA bioinks (blue: nucleus, red: F-actin, and green: MHC). (F) Relative expression of the MHC-positive area. (G) Cell alignment quantification. (H) Relative levels of mRNA expression of the myogenic differentiation markers MyoD and MyoG. Data reproduced from ref. 132. Copyright American Chemical Society 2021. (I)-(M) GO-PEI scaffolds for use in cardiac muscle engineering. (I) Schematic of GO-PEI scaffold fabrication. (J) Immunofluorescence staining images of Cx43 (upper panel) and the expression of Nkx 2.5 (lower panel) in HL-1 cells (blue: nucleus and green: F-actin), with (K) quantification. (L) Immunofluorescence staining of Cx43 (green) in the different groups (blue: nucleus). (M) Levels of protein expression of Cx43 and Nkx 2.5 in the different groups. Data reproduced from ref. 134. Copyright American Chemical Society 2023. (N)-(R) MXene/AuNP/GelMA hydrogel for use in skeletal muscle regeneration. (N) Schematic of MXene/AuNP/GelMA hydrogel formation and characterization of the bioprinted constructs. (O) Live/dead assay and quantification of the C2C12 myoblasts cultured within the printed constructs. (P) Low-magnification results of the live/dead assays of the constructs, which were printed differently. (Q) Immunofluorescence staining of the C2C12 myoblasts cultured within the printed constructs (blue: nucleus and green: MHC). (R) Myotube maturation parameters, including fusion index and myotube length and diameter. Data reproduced from ref. 136. Copyright American Chemical Society 2021. The scale bars represent (C-left) 1 mm and (E-right) 200 μm and (C-right, E-left, N, O and Q) 100 μm, and the asterisks denote statistical significance (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001, and ns: not significant)

Actn4, and TrpT-2. In vivo angiogenesis was evaluated by examining subcutaneously implanted scaffolds after 2 and 4 wk, which revealed enhanced levels of cell migration and increased levels of vascularization, as indicated by enhanced capillary formation and higher rates of cell migration. Pilato et al. developed composite scaffolds by combining GO with polyethylenimines (PEIs) to form highly porous, biocompatible scaffolds (GO-PEI) for use in repairing damaged heart tissues following myocardial infarction and heart failure (Fig. 5I-M). 134 In cell culture studies, HL-1 cardiac muscle cells exhibited good viability and morphological stability and numerous focal

adhesions and intercellular networks, indicating strong cellsubstrate interactions. Moreover, the levels of expression of the cardiac markers connexin 43 (Cx43) and Nkx 2.5 were notably higher in HL-1 cells cultured on the GO-PEI substrates. This may be attributed to the hydrophilic, polycationic nature of PEIdecorated GO, which promotes cell attachment and proliferation. Additionally, the nanotopographical cues provided by the GO-PEI surfaces contributed to improved cell behaviors. Quantitative analysis further revealed a significant increase in gap junctions, indicating functional syncytia characteristic of cardiac organization, in cells cultured on GO-PEI, highlighting its

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potential for use in cardiac tissue engineering. With the utilization of MXene NPs, Au NPs (AuNPs) have been widely applied in tissue engineering owing to their versatility and tunability in terms of size, morphology, surface chemistry, and other biological properties. 135 Boularaoui et al. introduced an AuNP- and MXene NS-laden GelMA (MXene/AuNP/GelMA) bioink for use in fabricating 3D bioprinted skeletal muscle-mimetic constructs (Fig. 5N-R). 136 The MXene/AuNP/GelMA bioinks exhibited excellent shear-thinning properties, reducing the shear stress that could harm encapsulated cells. It should be noted that GelMA crosslinking was impeded when MXene concentrations exceeded 0.1 mg mL⁻¹. The weakened crosslinking at higher MXene concentrations may be attributed to the spatial occupancy of the NPs and the interference between MXene's oxygencontaining functional groups and GelMA's amine and hydroxyl groups. This interference, in turn, hampers the free-radical photopolymerization process that GelMA undergoes when exposed to UV radiation. 137 Cell viability analysis showed no significant difference between the bulk and printed bioinks, indicating that printing did not negatively affect cell viability. Moreover, the addition of MXene and AuNPs to the bioinks enhanced skeletal muscle differentiation, as indicated by the increased fusion index and myotube length and diameter and upregulated MHC expression. Researchers attributed this improvement to the electrical conductivity of the bioinks, which may promote electrical communication between muscle cells and induce myogenic differentiation, rendering the bioinks promising for use in skeletal muscle tissue engineering. The beneficial biological properties of MXenes include free radical scavenging characteristics and anti-inflammatory activities due to M2 macrophage polarization. 138,139 Based on these properties, Ma et al. fabricated an MXene-laden F127 hydrogel (MXene-F127) to promote M2 macrophage polarization for skeletal muscle regeneration. 140 In vitro studies demonstrated that the MXene displayed strong antioxidant properties, effectively scavenged free radicals even at low concentrations, and reduced the levels of ROS in macrophages, which are induced by external stimuli. Furthermore, the MXene exhibited antiinflammatory properties due to the downregulation of proinflammatory markers (i.e., interleukin-1 β and tumor necrosis factor α) and promotion of the transformation of M1-type macrophages into M2-type macrophages, yielding a favorable microenvironment for myogenic differentiation and angiogenesis during skeletal muscle repair. Furthermore, the MXene enhanced the myogenic differentiation of C2C12 cells, promoting the formation of myotubes and the upregulation of myogenic genes, such as MyoD, MyoG, and MHC. The migration and angiogenesis of HUVECs were significantly promoted, facilitating the formation of complex vascular networks and the upregulation of angiogenic genes. In vivo studies using a rat tibialis anterior muscle defect model showed that the MXene effectively promoted skeletal muscle regeneration and the differentiation of macrophages to the M2 phenotype and reduced inflammation, further supporting its potential as a multifunctional bioactive material for use in skeletal muscle

MXene-polyethylene glycol (MXene-PEG) composite hydrogel constructs to provide conductive and topographical cues for human induced pluripotent SC (iPSC)-derived CMs (iCMs). 141 The iCMs cultured in the MXene-PEG hydrogel exhibited improved levels of structural and sarcomere alignment and increased levels of expression of cardiac maturity-related genes (i.e., MYH7, TNNT2, SERCA2, and GJA1). Electrophysiological analysis revealed that the MXene-PEG hydrogel cardiac patches exhibited spontaneous beating activities, with variations in the Ca²⁺ transient kinetics across different regions of the patterns. The conduction velocities between adjacent groups indicated synchronized beating in several regions, and detectable but slower conduction in others. Therefore, the MXene-PEG hydrogel displays potential for use in enhancing the structural and functional characteristics of iCMs in cardiac tissue engineering. A study conducted by Asaro et al. combined iCMs and MXene-functionalized Col (MXene-Col) films for use in cardiac tissue regeneration. 142 The MXene-Col films exhibited antibacterial properties, particularly due to the inhibition of the attachment and proliferation of S. aureus. These films also supported the growth of CMs and enhanced cell attachment, distribution, and maturation, which are essential in cardiac tissue engineering. When the iCMs were cultured on the MXene-Col films and exposed to an external electric field, they exhibited more mature phenotypes with improved levels of cell elongation and functional markers (i.e., Cx43, cardiac troponin T, sarcomeric α -actinin, and action potential properties). These studies suggest that MXenes and ES-combined MXenes show promise for application in muscle tissue engineering.

Skin tissue engineering using G, MXenes, and Xenes

Skin tissue engineering involves preparing artificial substitutes that replicate human skin, primarily driven by the crucial role of the skin as a protective barrier against various environmental threats. This field employs advanced tissue engineering techniques to develop skin substitutes for use as sophisticated models in research and in applications such as wound healing. A key challenge is the *ex vivo* expansion of cells, while preserving their normal properties, for use in transplantation or *in vitro* studies. Owing to its remarkable characteristics, such as its substantial surface area-to-volume ratio, mechanical durability, antimicrobial properties, and capacity to facilitate collagen crosslinking, G is used in the field of skin tissue engineering (Table 5). 145,146

Chu *et al.* fabricated a PEGylated, GO-mediated, quercetin (Que)-modified acellular dermal matrix (ADM; ADM-GO-PEG/Que) hybrid scaffold for use in skin tissue engineering and diabetic wound healing. The GO-PEG composite displays the capacity to efficiently deliver and release various substances, including proteins, gene therapeutics, imaging agents, chemotherapy drugs, and anti-cancer medications, while enabling precise control over release. The use of the ADM-GO-PEG/Que hybrid scaffold resulted in accelerated wound healing, with a faster rate of wound closure compared to those of other groups, reaching 87.34% closure by 21 d. Histological analysis revealed Col deposition, neovascularization, and the

tissue regeneration. Basara et al. introduced 3D printed Ti₃C₂T_x

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G, MXene, and Xene-based tissue engineering approaches for use in skin tissue regeneration Table 5

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Ref.	147		151		154			155			157		158				161	
Outlooks	 Accelerated regenera- Improved diabetic wound healing for potential drug tion on diabetic wounds delivery, stem cell therapies, and tissue engineering applications 		- In vivo wound closure Efficient antibacterial properties and positive impact on cutaneous wound healing		Drug delivery and temperature-responsive wound-			Reversible phase transformation under electrical sti- mulus offering promising applications in skin tissue	engineering with adjustable mechanical properties, controlled drug release, and electrical antibacterial	activity		energy harvesting, medical assistive systems, gait phase sensing, and wound monitoring and therapy,		uons and wound nearing			Germanene nanocrystals for photothermal anti- bacterial properties and wound healing capability	
Enhanced wound regeneration markers		 - Re-epithelialization - Dermal regeneration - Collagen deposition and neovascularization - Skin appendage regeneration - Col1, Col3, and α-SMA 	– <i>In vivo</i> wound closure	 Re-epithelialization Dermal regeneration Granulation tissue and collagen fiber formation Neovascularization 	- CD31, VEGF, and	FCINA – In vivo wound closure – Re-epithelialization	- Dermal regeneration	– <i>In vivo</i> wound closure	Re-epithelializationDermal regeneration		– In vivo wound closure	- Re-epithelialization	 Dermal regeneration In vivo wound closure 	: – Dermal regeneration	 Granulation tissue with collagen deposition 	- In vivo cell migration	– <i>In vivo</i> wound closure	- Re-epithelialization - Dermal regeneration
Test species Biocompatibility and therapeutic applications	 - 50 µM GO-PEG/Que supported high cell viability, proliferation, and spreading with reduced anomosis 	- Reduced <i>in vivo</i> inflammation	– Antibacterial activity against <i>S. aureus</i> and <i>Pseudomonas</i>	– Support proliferation	- T-RMF nanofiber matrices supported cell adhe-	sion, spreading, and promeration – No toxicity on main organs – NIR-irradiated wound healing		 ES-supported cell behaviors and antibacterial properties 	 Enhanced cell viability and wound healing No hemolytic activity 	 No erythema or edema after transplantation In vitro antibacterial property against E. coli Prevention of in vivo bacterial inflammation 	- Enhanced cell spreading, proliferation, and	<i>in vitro</i> wound healing – ES support wound healing	– 150 μg mL ⁻¹ Sn@PEI hydrogel maintained high	cen viabunty – US induced antibacterial activity against <i>S. aureus</i> – Dermal regeneration and MRSA	– Hemocompatibility	 - In vivo activation of inflammatory cells - ES support wound healing 	– No toxicity up to 2 mg mL ⁻¹	– NIR-irradiated antibacterial activity against <i>E. coli</i> – Re-epithelialization and <i>S. aureus</i> – Hemocompatibility – No toxicity on main organs <i>In vivo</i> suppression of bacterial colony
	' – MSCs	– Diabetic ICR mice	– MEF cells	- Rat	- BMSCs	- Kunming mice		- GES-1 cells	- Kunming mice		- L929 cells	r – SD rats	- 3T3 and	LUZ CEIIS - HUVECS	- BALB/c mice		- HeLa and NIH3T3 cells	- Kunming mice
Formulations	ADM-GO-PEG/ Que scaffold		UC-GSC scaffold		T-RMF-	vitalilli nanofiber		HA-DA@BP hydrogel			B-FENG	nanogenerator	Sn@PEI	nydrogei			Germanene CS/GeNCs0.8 hydrogel	
Xene Applications materials	09		rGO		$Ti_3C_2T_x$	MAGIIG		ВР			Borophene B-FENG		Stanene				Germanen	
Applicatic	Skin																	

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regeneration of skin appendages, including mature vessels and hair follicles, in the ADM-GO-PEG/Que scaffold-treated group on 14 d. The ADM-GO-PEG/Que scaffold enhanced Col synthesis and capillary formation and supported the formation of granulation tissue, rendering it effective in promoting the healing of diabetic wounds. In a study conducted by Esmaeili et al., nanofibrous scaffolds, denoted UC-GSC scaffolds, which were prepared using PU and cellulose acetate and contained rGO/Ag nanocomposites, were used to exploit the strong antibacterial properties of rGO/Ag. 151 The prepared UC-GSC scaffolds displayed similar rates of H₂O vapor transmission compared to those observed in other studies and biocompatibility with cells, indicating their viability for use in tissue engineering. Additionally, the scaffolds exhibited significant antibacterial activities against Gram-negative and -positive bacteria. The histological findings revealed that the UC-GSC scaffold exhibited significant improvements in wound healing,

including epidermal hyperplasia, granulation tissue formation, collagen fiber deposition, and neovascularization, indicating its potential for use in promoting wound healing.

MXenes may undergo degradation via their interactions with H₂O and O₂, indicating that their distinctive photothermal and biodegradable attributes may be exploited practically in wound healing. 152,153 Jin et al. developed temperature-responsive MXene nanobelt fibers (T-RMFs) loaded with vitamin E (T-RMF-vitamin E) for use in wound healing (Fig. 6). 154 These T-RMFs exhibited photothermal properties that could be controlled using NIR irradiation, which facilitated the release of vitamin E in wound healing. The experimental results indicated that the photothermal properties of the T-RMFs were controlled under NIR irradiation, thereby ensuring safe increases in temperature. In vitro and in vivo studies confirmed the excellent biocompatibility of the T-RMFs, which promoted cell attachment and proliferation. Furthermore, the T-RMFs, which could

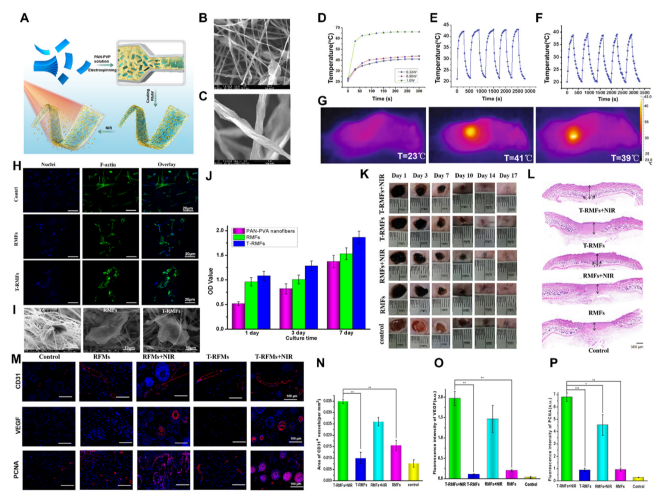


Fig. 6 Xene-based tissue engineering approaches for use in skin tissue regeneration. (A) Schematic of the fabrication of the T-RMF-vitamin E nanofiber matrices. (B) Low- and (C) high-magnification SEM images of the RMFs. (D) Heating profiles of the RMFs under 0.33, 0.50, and 1.0 W NIR light. Temperature changes of the (E) RMFs and (F) T-RMFs over 5 on/off cycles. (G) Thermal images of each group. (H) Immunofluorescence staining (blue: nucleus and green: F-actin), (I) SEM images, and (J) proliferation of the BMSCs cultured on the different nanofiber groups. (K) Representative photographs and (L) H&E staining images of the skin wounds of the various groups. (M) Immunofluorescence staining of CD31, VEGF, and PCNA of the BMSCs cultured on the different groups with or without NIR irradiation. Quantitative levels of expression of (N) CD31, (O) VEGF, and (P) PCNA. Data reproduced from ref. 154. Copyright Springer Nature 2021. The scale bars represent (M) 100 μm, (A-upper, H) 20 μm, (I) 10 μm, and (A-lower) 5 μm, and the asterisks denote statistical significance (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001, and ns: not significant).

release vitamin E under NIR exposure, significantly upregulated cluster of differentiation 31 (CD31), VEGF, and proliferating cell nuclear antigen (PCNA), which are markers of stimulated wound healing and angiogenesis. Notably, these therapeutic effects were realized with no adverse effects on the main organs, highlighting the potential of T-RMFs for use in safe, effective wound healing. BP may be utilized in skin tissue engineering by incorporating it into conductive hydrogels. Liu et al. prepared a conductive hydrogel, denoted HA-DA@BP, using hyaluronic acid (HA) and dopamine (DA), which was designed to produce antibacterial effects on wounds with the aid of ES. 155 The experimental results confirmed that the BP nanosheets within the hydrogel retained their original structures during degradation, and BP incorporation significantly enhanced the conductivity of the hydrogel, enabling effective electrical antibacterial activity against Escherichia coli when exposed to ES. Further in vitro studies were conducted using normal human gastric epithelial cells (GES-1), which revealed an excellent biocompatibility and the promotion of cell proliferation and migration under ES. In a mouse wound model, the HA-DA@BP hydrogel combined with ES significantly accelerated wound healing, reduced bacterial counts, and mitigated inflammation, indicating its potential for in vivo application as a wound dressing, with disin-

fection and tissue regeneration capacities.

Meanwhile, interest in borophene has been significant owing to its intricate layered structure, pronounced anisotropy, and exceptional electron transport and mechanical characteristics, but the number of studies exploring its potential in the field of tissue engineering is limited. 156 Chen et al. reported the utilization of a borophene/ecoflex nanocomposite-based triboelectric nanogenerator (B-TENG) in energy harvesting, medical devices, and wound healing. 157 The flexible B-TENG functioned based on triboelectricity, producing an electrical output via charge transfer upon contact with polyester, and it maintained its stability and durability, even after extensive evaluation and mechanical strain. The B-TENG was incorporated into a smart keyboard and robotic system for use in upper-limb medical assistance for disabled users; it was also applied in a lower-limb gait phase visualization platform to detect and display various gait phases. In terms of biological assessment, in vitro studies with L929 fibroblasts showed that the B-TENG could enhance cell proliferation and migration. Animal models also demonstrated the effectiveness of the B-TENG in promoting wound healing, with reduced wound sizes and improved levels of tissue regeneration. Stanene NSs (SnSNs) are semiconductors that may exhibit thickness-dependent adjustable bandgaps which render them favorable for use in cancer photothermal therapy. 63 Tao et al. introduced SnSNs-assembled thermosensitive poly(D,L-lactide)-PEG-poly(D,L-lactide) for use in preparing Sn@hydrogels to exploit their sonodynamic antibacterial activities. 158 Sn@PEI exhibited excellent sonodynamic properties, generating ROS effectively under ultrasound (US) irradiation, leading to antibacterial activity. Sn@PEI displayed high antibacterial efficiencies against S. aureus and methicillinresistant S. aureus when exposed to US, whereas no significant cytotoxicity was observed toward normal cells. In vivo analysis

revealed accelerated wound healing (full regeneration on 14 d) in mice after the transplantation of the ES-supported Sn@PEI hydrogel. The hydrogel also exhibited a potent antibacterial activity, drastically reducing the levels of bacterial contamination of infected skin wounds. Germanene displays considerable potential as a photothermal agent for use in treating infected wounds due to its efficient photothermal conversion, while its biocompatibility and low toxicity render it promising for use in clinical applications. 159,160 Therefore, Wang et al. developed a CS hydrogel embedded with hydroxy-functionalized germanene nanocrystals (GeNCs) via the crosslinking of CS and zinc acetate and self-assembly with the GeNCs to yield the photothermal wound-healing hydrogel CS/GeNCs0.8.161 In vitro studies revealed that the CS/GeNCs0.8 hydrogels effectively adsorbed and killed Gram-positive S. aureus and Gramnegative E. coli when combined with NIR laser irradiation. SEM revealed that the NIR-irradiated CS/GeNCs hydrogels caused significant protein and nucleic acid leakage by disrupting membrane integrity. Additionally, the CS/GeNCs0.8 hydrogels exhibited excellent hemostatic properties and negligible hemolytic effects on red blood cells. In vivo studies in mice indicated that the CS/GeNCs0.8 hydrogel effectively inhibited bacterial infection and promoted wound healing, with no significant toxicity or adverse effects on body mass or the major organs, indicating its potential as a photothermal antibacterial platform for use in clinical applications.

Conclusion and future perspectives of MXene and Xene materials in tissue engineering

The achievements of G have demonstrated the feasibility of producing stable, ultra-thin layers of van der Waals materials, consisting of just one or a few atoms, using diverse techniques for fabricating single-layer, few-layer, and multi-layer assemblies in solution, on surfaces, and at a wafer scale. 162,163 In recent decades, there has been great achievement led by pioneers in the synthesis, characterization, and applications of 2D nanomaterials.164 For instance, P. M. Ajayan led the way in the large-scale production and analysis of single layers of hexagonal boron nitride, while A. Kis and his research team achieved the synthesis of single-layer MoS₂ for use in optoelectronics and energy harvesting applications. 165,166 These advancements in 2D nanomaterials have led to increased utilization of these materials in bioengineering research, exploiting their distinctive physicochemical and biological properties.167

This review extensively compares the research progress of Xene-material-based tissue engineering with that of G-based materials, which have been extensively studied over the last decade. Section 2 provides an overview of the fundamental physicochemical characteristics of G, MXenes, and group III-V Xene materials. In the subsequent section, we review recent experimental advancements in Xene-based tissue engineering and regenerative medicine for different tissue types, *i.e.*, bone, cartilage, neural, muscle, and skin tissues. The versatility of

fabrication techniques (e.g., 3D bioprinting and electrospinning) and cell sources (e.g., SCs and primary cells) enabled researchers to produce numerous scaffolds and tissue mimetics with various structures and biofunctionalities. Each study was reviewed to explore the innovative combinations of Xene materials with other biomaterials and their capacity to promote tissue maturation from a mechanistic perspective.

The goal of tissue engineering and regenerative medicine is to prepare biomimetic constructs or matrices that provide biochemical and structural support to cells, induce the expression of intrinsic phenotypes, and facilitate differentiation to mature tissues. G-based research laid the foundation for 2D NM-based tissue engineering and serves as a valuable starting point for the subsequent development of MXene and Xene materials. Despite the promising progress, several critical challenges remain unresolved. Safety and biocompatibility are primary concerns, as the biocompatibility of most Xene materials has not been fully elucidated, necessitating further studies and long-term clinical investigations under various conditions. Understanding the interactions between Xene materials and the body (i.e., biodistribution, organ accumulation, clearance pathways, inflammatory reactions, and potential genotoxicity) is crucial in exploring their future in vivo applications. 168 Moreover, engineered scaffolds exhibit unique complexities compared to conventional grafts, rendering the following of regulatory guidelines and processes challenging. Additionally, the multitude of active components and fabrication parameters may unpredictably affect reactions within the human body.

To address these limitations, further interdisciplinary studies and clinical trials are required to elucidate long-term efficacy and potential side effects. Standardization of culture conditions and quality control are essential to ensure the safety, efficacy, and reproducibility of Xene-based engineered scaffolds. For future clinical applications of Xene materials, it is crucial to understand the unique properties of various elemental composites and surface reactivity with the biological system in relation to their fabrication processes and structures. The ability to tailor their exterior surface functional groups enhances their adaptability for therapeutic and imaging purposes. Successful translation of preclinical findings into clinical settings, demonstrated by ongoing clinical trials, will pave the way for the acceptance and implementation of these nanomaterials in personalized medicine. Integrating nanotechnology with artificial intelligence (AI)-based methods for drug combination and dosing optimization holds great potential for enhancing the efficacy and clinical feasibility of Xene-based biomedical applications. In conclusion, researchers with diverse backgrounds should advance the field of Xene-based tissue engineering by addressing these challenges and considering the proposed suggestions.

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Conflicts of interest

There are no conflicts to declare.

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