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REVOLUTIONIZING HEALTHCARE: THE ROLE OF NANOMATERIALS IN TISSUE ENGINEERING AND MEDICAL THERAPIES

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Abstract:- Nanomaterials are revolutionizing tissue engineering and medical practices through their unique attributes, such as high surface area, diverse physical and chemical properties, and their ability to engage with biological systems at the molecular level. In tissue engineering, these materials significantly improve scaffold designs, which supports enhanced cell attachment, proliferation, and differentiation necessary for effective tissue regeneration. They are also crucial for the advancement of controlled drug delivery systems, which are integral to developing functional tissue models. In medical applications, nanomaterials are increasingly being used for diagnostic procedures, targeted drug delivery, and various treatments, leading to improved therapeutic outcomes and reduced side effects. Their use in imaging and diagnostics facilitates early disease detection and supports tailored treatment strategies, transforming patient care. Ongoing research into nanomaterials is expected to tackle complex healthcare issues, resulting in the development of safer and more innovative medical therapies.

Key words: Nanomaterials; Tissue Engineering; Scaffold Design; Controlled Drug Delivery; Medical Diagnostics; Targeted Therapy; Regenerative Medicine

I. INTRODUCTION

Tissue engineering and regenerative medicine have revolutionised the methods for repairing and replacing the damaged tissues and organs by employing engineered tissues, biomaterials, and cellular therapies to restore normal functions. As the need for effective treatments increases, nanomaterials have become essential in addressing the limitations of traditional biomaterials. Their unique physicochemical properties such as a high surface area-to-volume ratio, enhanced mechanical strength, and adjustable interactions with biological systems make them vital for advancing these fields. Nanomaterials, with sizes ranging from 1 to 100 nanometers, represent a major advancement in scaffold design for tissue engineering. These nanostructured

scaffolds can better replicate the natural extracellular matrix, promoting improved cell attachment, growth, and development. Additionally, drug delivery systems based on nanoparticles enable precise and controlled release of medications, enhancing their effectiveness and minimising side effects. Nanomaterials also play a key role in diagnostic imaging and biosensing, allowing for earlier detection of diseases and real-time monitoring of treatment outcomes. Despite these advancements, there are still significant challenges in moving nanomaterials from research into clinical practice. Ensuring biocompatibility, managing potential toxicity, and meeting regulatory standards remain critical. Developing comprehensive methods to assess the long-term effects of nanomaterials on health and the environment is essential for their successful clinical application.(Fig 1)

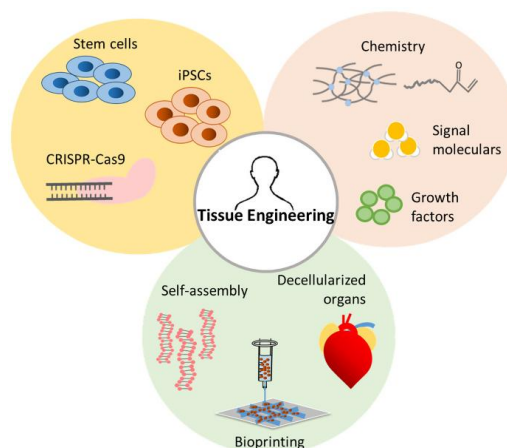


Fig 1 : Key elements in Tissue engineering , Source : Tianyu Yao, Baker, M. B., & Moroni, L. (2020b). Tissue engineering process. Strategies to Improve Nanofibrous Scaffolds for Vascular Tissue Engineering. MDPI. Retrieved April 24, 2020

INNOVATIONS IN TISSUE ENGINEERING

1. Combating Antibiotic Resistance with Nanoparticles

The Centre for Disease Control (CDC) had predicted that deaths from antibiotic-resistant bacteria could surpass those from all cancers by 2050, thus highlighting the urgent need for



non-antibiotic treatments. Nanoparticles show promise in this area, especially silver nanoparticles, which are known for their antibacterial properties and wound healing capabilities. For example, poly(3-hydroxybutyrate-co-3-hydroxyvalerate) nanofibrous scaffolds with silver exhibit good cell compatibility and strong antibacterial activity, making them potential candidates for joint arthroplasty[44]. Bio composite scaffolds with nano silver also show promise in preventing infections during reconstructive bone surgeries. Additionally, selenium nanoparticles are effective against various bacterial strains without resistance and iron oxide nanoparticles, especially when functionalized with a magnetic field, show potential in breaking down biofilms in tissue engineering. These advancements demonstrate the broad potential of nanoparticle-based therapies in addressing bacterial infections and antibiotic resistance.[43]

2. Stimulation of cells for Mechano transduction

The human body's cell function is known to be regulated by a variety of bioactive chemicals and growth factors, but mechanical forces also have a big impact via affecting mechano transduction pathways[45]. Although shear stress has been applied and cell activities have been mechanically controlled using bioreactors and substrate stiffness, magnetic nanoparticles (MNPs) provide unique benefits. MNPs are excellent at controlling cell behaviour because they can be remotely regulated both spatially and temporally using a magnetic field. A magnetic field applied to MNPs coated with certain targeting antibodies causes cells to align, which affects receptor-mediated cell functions. Research conducted by Mannix et al. and Gopinath et al. revealed that MNPs have the ability to regulate intracellular calcium levels and trigger apoptotic pathways, respectively. These findings highlight the potential of MNPs in comprehending and modifying cell activities in regenerative medicine. On the other hand, extracellular stressors induced by $11.0 \mu\text{g}\cdot\text{mL}^{-1}$ of silver nanoparticles have been used to induce apoptosis in BHK21 and HT29 cells, which results in the induction of p53 and subsequent pathways leading to cell death. On the other hand, because these methods include internalization processes or interactions between nanoparticles and cell membranes, care must be taken to prevent unintentional changes in cellular functioning.[43]

3. Gene delivery in Tissue engineering

Gene delivery technology in tissue engineering (TE) targets both matured cells and stem cells like human mesenchymal stem cells (hMSCs), which can differentiate into various cell types. Non viral methods, such as magnetofection using magnetic nanoparticles (MNPs), offer simplicity and reduced immune response compared to viral vectors [46]. MNPs complexed with DNA are guided by magnetic fields to enhance gene transfection efficiency, especially in challenging cell types like endothelial and embryonic stem cells. Iron oxide particles in polymer matrices, mesoporous silica nanoparticles,

and carbon nanotubes (CNTs) are effective non viral carriers due to their structural properties and biocompatibility. In viral transduction, magnetite cationic liposomes (MCLs) significantly enhance the delivery efficiency of retroviral vectors. Adenoviral vectors coated with MNPs have also successfully transduced various cell lines for therapeutic applications, including delivering antisense oligonucleotides and siRNA.[47] Overall, magnetically guided gene delivery systems show promise for advancing TE and treating diseases by improving cellular uptake and enhancing the therapeutic efficacy of genetic materials.[43]

4. Magnetic Cell patterning

Successful tissue engineering (TE) requires mimicking in vivo tissue architecture, often achieved by controlling cell adhesion patterns. Traditional methods are limited by complex equipment or chemically modified surfaces, but magnetic nanoparticles (MNPs) offer a practical alternative. MNPs in magnetite cationic liposomes (MCLs) facilitate precise cell patterning using magnetic fields [48]. For example, human umbilical vein endothelial cells formed capillary-like structures when patterned using MCLs on acryl resin plates. In vascular graft construction, magnetic seeding with MCLs improved cell attachment efficiency on decellularized arteries compared to conventional methods. Another approach involves MNPs coated with aminosilane and poly(ethylene glycol) (PEG-Mags) to create cell-resistant regions on tissue culture dishes [49]. HaCaT cells selectively adhered to untreated surfaces, demonstrating successful heterotypic cell patterning with mouse myoblast C2C12 cells. This method showed minimal cytotoxicity, highlighting its biocompatibility and potential for complex cell arrangements in tissue engineering applications.[43]

5. Constructing 3D tissues

MCL-labeled cells have been effectively used to create keratinocyte sheets for skin applications on ultra low-attachment plates, with a neodymium magnet underneath to promote cell adhesion and development. These sheets formed stronger, 10-layer epidermal constructions when cultivated in a high-calcium medium. Magnetic force-based scaffolds have enabled the assembly of multilayered sheets of myoblast cells in skeletal muscle tissue engineering, resulting in dense and homogeneous tissue creation [53]. By adjusting the magnetic field gradient, this method has also been modified to produce 3D muscle fiber bundles in the form of strings. Magnetic force-based tissue engineering (Mag-TE) techniques have been employed in liver tissue engineering to create cell sheets for cocultures of endothelial and hepatocyte cells [50]. Magnetic cell patterning facilitated strong cellular connections, improved the deposition of extracellular matrix (ECM), and enhanced cytokine production between cell layers [51]. These advancements in Mag-TE have demonstrated the ability to enhance liver function by providing stable cellular

constructions for extended periods and increasing albumin secretion.[43]

APPLICATIONS IN TISSUE ENGINEERING

1. Dental Tissue Engineering

Metal nanomaterials like silver (Ag), gold (Au), titanium dioxide (TiO₂), and zinc oxide (ZnO) are highly valued for their strong antibacterial abilities [2], making them important in dental tissue engineering. These materials can be modified in various ways to enhance their antibacterial effectiveness. Research indicates that nanoparticles (NPs) smaller than 10 nm exhibit stronger bactericidal effects, with specific shapes like triangular NPs showing superior antibacterial properties compared to spherical or needle-shaped counterparts [4]. In dental applications, Ag/Au alloy bimetallic nanoparticles produced through electric current displacement reactions have shown good biocompatibility and strong antibacterial effects against pathogens such as *Porphyromonas gingivalis* W83, which is associated with periodontal disease [3]. Environmental factors like hydrogen peroxide can improve their antibacterial effects by mimicking the oxidative stress found in chronic periodontal inflammation. A new method uses light-activated nano-antibacterial scaffolds with gold nanocages, designed to release antibiotics while combining the benefits of phototherapy and chemotherapy. This technology has shown promise in lab and animal studies, indicating its potential for effective antibacterial treatment in dental care.[1]

2. Neural Tissue Engineering

This employs various materials, such as scaffolds, hydrogels, nanoparticles, and nerve conduits, to repair nerve damage and promote regeneration. Since the peripheral nervous system (PNS) and central nervous system (CNS) have limited ability to heal on their own, these materials are essential for treating functional problems caused by diseases and injuries.

- **Collagen:** This is a natural biopolymer found in connective tissues, is used for regeneration of nerves because of its structural support and biocompatibility, with products like NeuraGen® and Neuromaix® showing success in animal studies[5].
- **Gelatin:** Derived from collagen, nanoparticles and electrospun scaffolds which are gelatin based nano materials provide biodegradability, biocompatibility, and improved cell adhesion for nerve regeneration [6].
- **Protein-Based Materials:** Elastin, keratin, and silk help in nerve tissue engineering by offering biocompatibility, mechanical support, and unique biological functions, with silk-based hydrogels aiding in neuronal differentiation and tissue regeneration[7].
- **Carbon-Based Nanomaterials:** Carbon nanotubes, fullerenes and graphene have great electrical conductivity and mechanical properties, making them ideal for interacting with neural tissues. They help promote neuron

cell growth and differentiation, which aids in regeneration of nerves [8].

Materials like chitosan, alginate, PLGA, PLA, and PEG are extensively studied for tissue engineering of nerves due to their biodegradability, biocompatibility and controlled release features. They are key in developing drug delivery systems and advanced nerve conduits with ongoing research aiming to improve these materials and discover new applications.[1]

3. Bone Tissue Engineering

Bone tissue engineering seeks to repair severe injuries with advanced nanomaterial implants. Current strategies use scaffolds with physical and biological methods to support bone regeneration[11]. 3D scaffolds are essential for structural support and bone growth, with key factors being mechanical strength, surface features, and biocompatibility.

- **Chitosan:** It is an FDA approved biodegradable polymer and is crucial in bone tissue engineering for its biocompatibility and support for bone growth, as demonstrated by Chesnutt et al. with their chitosan/nanocrystalline calcium phosphate framework [9].
- **Hydroxyapatite nanoparticles (nHAp):** As a vital inorganic bone component, nHAp is extensively studied for enhancing bone regeneration. Liu et al. explored a chitosan- hydroxyapatite biomimetic nanocomposite scaffold promoting bone marrow mesenchymal stem cell proliferation and activating bone growth pathways.[10]
- **Bioactive glass-ceramic nanoparticles (nBGC):** These mimic bone composition and significantly contribute to bone tissue engineering. Singh et al. demonstrated improved bone marrow MSC proliferation and differentiation using bioactive glass/polyvinyl alcohol and silk fibroin scaffolds.[12]
- **Carbon-based nanomaterials:** Graphene and carbon nanotubes offer tunable surface functionalities and excellent mechanical properties. Composite scaffolds like PEI/GO promote human bone marrow MSC proliferation and enhance osteoblast differentiation, crucial for bone repair.[13]
- **Metal nanoparticles:** Silver, gold, and titanium dioxide are gaining attention for their mechanical strength and osteogenic properties in bone tissue engineering. Silver nanoparticles enhance cell absorption capacity without adverse effects, showing promise for bone regeneration applications.[14]

These advanced materials offer new solutions for bone tissue engineering, improving clinical outcomes and reducing patient discomfort with natural degradation and biocompatibility, while ongoing research explores their use in orthopedic and reconstructive surgeries.[1]



4. Skin Tissue Engineering

Wound healing involves several stages: hemostasis, inflammation, proliferation, and remodelling. Acute wounds usually heal quickly, but chronic wounds from conditions like obesity and diabetes are much more challenging[15]. Angiogenesis which is the formation of new blood vessels and is crucial for supplying nutrients and oxygen to wounds and forming granulation tissue [17]. Effective wound treatment includes therapies such as local oxygen, hyperbaric oxygen, ozone, and negative-pressure wound therapy. Although autologous skin transplantation is limited by availability, cell-based therapies can aid healing, though their success rates, treatment cycles, and costs can vary.

Nanotechnology improves wound healing by creating barriers for keratinocyte regeneration, securely adhering to the dermis, remodeling blood vessels, and providing structural support [16]. Key materials include:

- **Nanocellulose:** Derived from cellulose, nanocellulose absorbs wound fluids, makes dressing removal easier, and helps tissue regeneration while reducing inflammation. Nanocellulose-based dressings keep wounds moist and support healing, and are effective for treating chronic ulcers and severe burns[17].
- **Nanocarriers for Gene Transfection:** These carriers protect molecules from breaking down, control gene release, and improve effectiveness in skin tissue engineering. PLGA nanoparticles with anti-VEGF plasmids help manage vascular growth factor levels, which is useful for treating blood vessel disorders and improving wound healing. Engineered stem cells that produce growth factors like VEGF boost blood vessel formation and aid wound healing with nanostructured scaffolds.[18]

ADVANCEMENTS IN TISSUE REGENERATION AND REPAIR

1. Bone Tissue Regeneration

Bone tissue is a complex nanocomposite and is made of collagen and hydroxyapatite (HA), which contains calcium phosphate. It includes various cells such as osteocytes, osteoblasts, osteoclasts, and other bone-related proteins. Bone loss can result from trauma, diseases or surgeries, posing significant challenges for reconstruction. Traditional methods for bone regeneration, such as autografts and allografts, have limitations like availability, infection risk, and donor site issues. Ideal bone grafts should be pathogen-free, osteoinductive, stable, and have minimal antigenicity. Bioinert materials have been used for mechanical support, but there's a shift towards bioactive materials that enhance implant-bone integration.[52]

- Nanostructured scaffolds are advanced tools in bone tissue engineering. Materials like carbon nanotubes

(CNTs), electrospun hydroxyapatite (HA), anodized titanium, and nano titanium improve osteoblast attachment and bone regeneration. Bioinspired nanoscale synthetic HA mimics natural bone and helps regenerate bone cells. Calcium phosphate materials boost biocompatibility and support implant integration, while adding metallic nanoparticles to polymers like chitosan and HA enhances the scaffolds' biological properties and antimicrobial benefits.[55]

- Nanotechnology shows great potential for improving mesenchymal stem cell (MSC) differentiation in bone regeneration. Nanoscale matrices, like nanofibrous scaffolds and nanoparticles, enhance MSC growth and promote bone formation. These nanomaterials can be as effective as traditional growth factors and offer a flexible alternative in tissue engineering. However, it's crucial to balance their effectiveness with potential toxicity risks, requiring careful safety assessments and characterization to optimise MSC differentiation for bone regeneration.[56]

2. Cartilage Tissue Regeneration

Cartilage, primarily composed of type II collagen, plays a crucial role in facilitating smooth joint movement. However, its limited natural repair ability, attributed to the absence of blood vessels and stem cells, presents challenges in healing. Recent strides in nanotechnology have offered promising avenues for enhancing cartilage regeneration.

- **Nanofiber Scaffolds:** Electrospun nanofibers, integrated into microfibrillar scaffolds, closely mimic the cartilage extracellular matrix (ECM). These scaffolds support cell growth and glycosaminoglycan (GAG) deposition crucial for cartilage regeneration, even in serum-free conditions.[52]
- **Peptide Nanofibers:** By incorporating proteins and GAGs into peptide nanofibers, researchers have developed scaffolds that promote the formation of cartilage tissue nodules. This approach achieves both chemical and structural resemblance to the ECM, facilitating effective cartilage repair.
- **Pentosan Polysulfate (PPS):** PPS, a sulfated polysaccharide, enhances the differentiation of mesenchymal stem cells (MSCs) into chondrocytes when combined with a polyethylene glycol (PEG)-hyaluronic acid (HA) medium. The bound form of PPS in PEG-HA shows promising results for regenerating intervertebral discs.[58]
- **Stem Cell-Coated Scaffolds:** Scaffold coatings with MSCs have demonstrated superior cell growth and differentiation properties compared to uncoated scaffolds. In vitro and in vivo studies highlight the efficacy of MSC-coated scaffolds in enhancing cartilage repair processes.[57]



- **Functional Biomimetic Scaffolds:** Modification of scaffolds with materials like multiwalled carbon nanotubes (MWCNTs) and poly-lysine improves their mechanical properties and supports MSC differentiation into chondrocytes. These modifications also stimulate GAG synthesis, crucial for effective cartilage repair and regeneration.[52]

3. Nerve Tissue Regeneration

Nerve tissue regeneration is challenging and often requires grafts or guiding materials. Nanotechnology offers solutions through aligned electrospun nanofibers that guide nerve cell growth and enhance neurite extension. Nanofibrous guides coated with proteins and growth factors improve nerve repair in animal models. Nanoporous scaffolds and conductive nanopolymers support nerve fiber growth and cell maturation. [54] For the central nervous system (CNS), nanotechnology aids drug delivery across the blood-brain barrier (BBB) using modified nanoparticles like chitosan nanogels. While the peripheral nervous system (PNS) benefits more from nanobiotechnology due to Schwann cells, the CNS struggles with nerve cell remyelination. Ongoing research aims to improve nanotechnology for nerve regeneration and CNS treatments.[52]

4. Myocardial Tissue Regeneration

Cardiac myocytes have limited growth potential and often form scars after ischemic injury, which can lead to complications like cardiac rupture. Preventing myocyte death and promoting regeneration are crucial. [63] Combining IGF-1 with PLGA nanoparticles helps reduce cell death by activating Akt phosphorylation in ischemic heart tissue. Smaller PLGA nanoparticles can deliver more IGF-1, enhancing this effect. For encouraging human mesenchymal stem cells (hMSCs) to become cardiomyocytes, using scaffolds like cross-linked electrospun hemoglobin/gelatin/fibrinogen nanofibers with high oxygen-carrying capacity has proven effective. [64] Additionally, poly(glycerol sebacate) (PGS) biomaterials with a gelatin shell support cell adhesion and proliferation. Injecting hMSCs with short PGS nanofibers into heart tissue improves cell retention and increases the production of key cardiac proteins compared to standard methods.[52]

5. Skin Tissue Regeneration

The skin, serving as the body's largest organ, fulfills critical functions in protection, temperature regulation, and vitamin D synthesis but remains vulnerable to injuries and infections due to its external position. While current bioengineered skin models have advanced, they still fall short of replicating the intricate characteristics of natural skin. [52] Nanotechnology has emerged as a transformative field in wound healing by employing several innovative approaches such as :

- **Nanostructured Scaffolds:** [59] Designed to mimic the extracellular matrix, these scaffolds promote cell growth and migration. Electrospun nanofibers, for example, enhance cell proliferation and adhesion on wound surfaces. Plasma-treated electrospun scaffolds, enriched with integrin-binding sites in gelatin, significantly improve cell adhesion and proliferation.
- **Nanoparticle-Based Delivery Systems:** [52] Nanoparticles deliver growth factors directly to wound sites, enhancing local availability and accelerating healing. Nanoparticle-delivered growth factors like KGF 2 have demonstrated effectiveness in promoting cell proliferation and improving wound closure rates.
- **Hydrogels:** These provide a supportive matrix for skin regeneration. Incorporating rosette nanotubes (RNTs) into hydrogels such as poly(2-hydroxyethyl methacrylate) enhances keratinocyte and fibroblast proliferation, leveraging their natural skin-mimicking properties[60] .
- **Stem Cell-Based Therapy:** Utilizing various stem cell types, particularly adipose-derived stem cells (ASCs), offers a promising avenue for skin regeneration. ASCs are abundant, easy to harvest, and effective in promoting healing. Electrospun nanofibrous scaffolds further optimize ASC differentiation and proliferation, facilitating more efficient skin tissue regeneration.[52]

6. Dental Tissue Regeneration

The human tooth has two main parts: the visible crown and the root embedded in the bone. It consists of four primary tissues: enamel, dentin, pulp, and cementum. Enamel covers the crown and is the hardest substance in the body, while dentin forms most of the tooth structure. Pulp contains nerves and blood vessels crucial for tooth health, and cementum anchors the root. Modern diets and longer lifespans have increased dental issues like cavities, which weaken teeth by demineralizing enamel and dentin. [61] Nanotechnology is transforming dental care with innovations such as biocompatible implants that improve durability and integrate well with the jawbone, and novel materials like bioactive glasses and nano-filled resins to address tooth wear and infections. These advances promise to enhance the effectiveness, longevity, and appearance of dental treatments, improving oral health and patient well-being.[52]

7. Hepatic Regeneration

Nanotechnology is transforming liver regeneration by creating precise models of the hepatic extracellular matrix (ECM) using nanoparticles and nanostructured scaffolds. These innovations help progenitor cells become functional hepatocytes. A major challenge is keeping hepatocytes alive outside the body in conditions that mimic their natural environment. Giri et al. developed a nanomaterial with self-assembling peptides and growth factors that can sustain hepatocytes for up to 3 months without toxicity. Silymarin nanoparticles, made through nano precipitation, protect the

liver and boost glutathione levels to counteract damage from substances like paracetamol. Additionally, mesenchymal stem cells (MSCs) from bone marrow, fat, and myeloid tissue can turn into hepatocyte-like cells. Studies using chitosan-alginate scaffolds with MSCs have shown liver tissue growth and hepatocyte marker expression, suggesting these scaffolds can effectively deliver and maintain MSC differentiation into hepatocytes[52].

8. Ocular Regeneration

Nanotechnology is advancing ophthalmology, especially in regenerative nanomedicine for vision restoration. Research by Teo et al. found that nano pillars support bovine corneal endothelium regeneration better than micro wells/pillars by boosting Na⁺/K⁺ ATPase activity and mRNA levels crucial for corneal health. Another study used superparamagnetic nanoparticles with tropomyosin-related kinase B agonist antibodies to activate pathways that promote neurite growth, offering potential for treating central nervous system injuries. [62] Uzunalli et al. developed bioactive nanofibers from laminin and fibronectin that improved human keratocyte growth and morphology in the lab. In animal models, these laminin-mimetic nanofibers enhanced corneal healing and regeneration, indicating potential for treating corneal diseases.[52]

nanoparticles, like those based on molybdenum and selenium, offer potential for treating acute kidney injury with their antioxidant properties. Carbon nanotubes (CNTs) enhance scaffold properties for tissue regeneration but pose toxicity risks, requiring further research [20]. Functionalized CNTs show promise for renal tissue regeneration and drug delivery. Exosomes, small vesicles from mesenchymal stem cells, aid tissue repair and improve kidney function by enhancing cell growth and reducing cell death [21]. Biomimetic nanofibers, mimicking the extracellular matrix, play a crucial role in tissue regeneration and bio-artificial kidneys. Recent advances include zeolite-enhanced polyacrylonitrile (PAN) nanofiber membranes for dialysis and wearable blood purification systems using nanofiber meshes [22]. Biofabrication techniques help create miniaturised hemodialysis models and wearable artificial kidneys. Despite ongoing challenges, nanotechnologies offer exciting prospects for improving kidney treatments and organ development.[19]

2. Advanced cancer therapies

Cancer continues to be a major cause of death, and chemotherapy is a prevalent treatment method. However, conventional chemotherapeutic drugs face issues like poor solubility in water, dose-dependent toxicity, and lack of precise targeting to tumors. Additionally, multidrug resistance complicates treatment by enhancing efflux pumps that expel drugs from cells. Recent advancements in nanoparticle-based delivery systems offer potential solutions. These systems can target cancer cells directly, control drug release rates, and enhance therapeutic effectiveness. Two primary strategies are employed: passive and active targeting. Passive targeting exploits the unique characteristics of tumor vasculature, which allows nanoparticles to accumulate in tumors due to their leaky blood vessels and poor lymphatic drainage, a phenomenon known as the Enhanced Permeability and Retention (EPR) effect. However, challenges such as mucosal barriers and non-specific particle uptake can limit effectiveness. Active targeting, on the other hand, uses nanoparticles coated with ligands that specifically bind to cancer cell receptors, improving targeting precision. Numerous FDA-approved nanoparticles are reformulations of existing chemotherapeutic agents combined with polymeric carriers. For example, [24] Doxil® uses PEGylated liposomes to extend drug circulation and reduce heart-related side effects, leveraging the EPR effect for better tumor accumulation. Marqibo® utilizes liposomes to enhance the effectiveness and circulation time of vincristine sulfate compared to traditional formulations [23]. Lipoplatin®, which involves cisplatin in a liposomal formulation, shows reduced kidney toxicity and is approved for treating several cancers [26]. Drug conjugates also play a role, such as Kadcyla®, which links the drug DM1 to the antibody trastuzumab for targeted delivery to HER2+ breast cancer cells, or Abraxane®[25], where paclitaxel is combined with albumin to improve delivery and reduce side effects. These developments

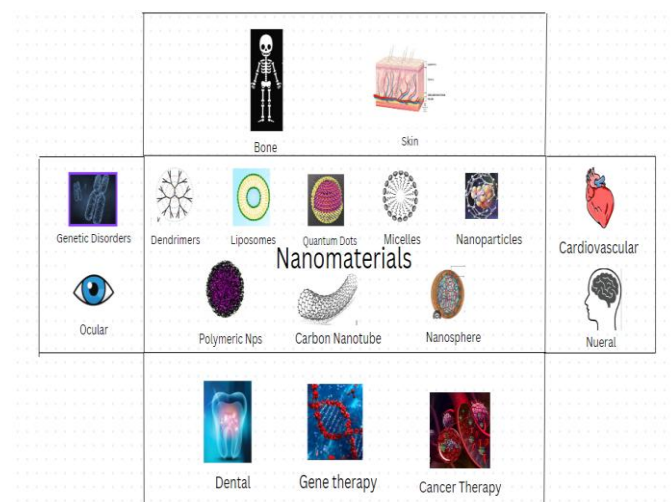


Fig 2 : Applications of nanomaterials in healthcare

NANOMATERIALS IN DRUG DELIVERY AND THERAPEUTICS

1. Kidney targeted drug delivery systems

Nanoparticles are increasingly vital in kidney-related treatments and the creation of artificial kidneys due to their customizable size, shape, and surface properties. They enable precise drug delivery, such as actinomycin D-loaded formulations, and improve renal therapies with nanoporous membranes that filter solutes based on size. However, their biocompatibility needs careful evaluation. Inorganic



illustrate how nanoparticle-based therapies are advancing cancer treatment by improving drug delivery and reducing adverse effects.

NANOMATERIALS IN DIAGNOSTICS AND IMAGING

Throughout history, pandemics have demonstrated the critical need for effective and scalable diagnostic tools. The emergence of highly contagious diseases such as the Black Death and COVID-19 has highlighted this need, prompting advancements in medical technology. Nanotechnology has played a transformative role in developing diagnostic tools that are both rapid and scalable. Carbon-based nanomaterials, including carbon nanotubes and graphene, have been employed to create highly sensitive biosensors for detecting pathogens. For example, carbon nanotube sensors can identify SARS-CoV-2 with exceptional sensitivity, detecting viral proteins at concentrations as low as 2.4 picograms per milliliter [28]. Gold nanoparticles, facilitate the detection of viral proteins at femtomolar levels, enhancing diagnostic accuracy. The advent of lipid nanoparticles (LNPs) has revolutionized vaccine delivery, particularly with mRNA vaccines for COVID-19. LNPs safeguard mRNA from degradation and enable its efficient delivery into cells [29]. This technology has been crucial in the swift development of highly effective vaccines like those from Pfizer-BioNTech and Moderna, which have shown around 95% efficacy. The adaptability of LNPs opens the door to rapid vaccine development for other diseases and future pandemics.[27]

In cancer diagnostics, nanomaterials have enabled significant advances. Nanoparticles such as gold and graphene oxide enhance imaging and detection. For instance, engineered bacteriophages have improved fluorescence for better tissue imaging, while graphene oxide-based devices offer efficient cell capture from blood samples, enhancing diagnostic precision [30]. "Smart" nanomaterials that respond to specific biological signals are pushing the frontiers of diagnostic technology. These materials can detect disease biomarkers, providing timely and non-invasive diagnostic options. For example, some nanomaterials react to changes in pH or other indicators in tumors, aiding in early cancer detection. Overall, nanotechnology is transforming medical diagnostics and treatment by providing more precise, rapid, and cost-effective solutions. These innovations are crucial for improving public health and offering personalized care options.[27]

BIOCOMPATIBILITY OF NANOMATERIALS

1. Hemocompatibility

Nanoparticles (NPs) play a significant role in drug delivery, gene therapy, and biosensing, necessitating thorough evaluation of their compatibility with blood. Key assessments include hemolysis and coagulation tests. Chouhan et al. and Bajpai et al. had evaluated the blood compatibility of PHEMA NPs using the hemolysis assay [32]. They found the least hemolysis with a formulation of 12.37 mM HEMA and 1.06 mM EGDMA, indicating a moderate level of

biocompatibility. In another study, Sanoj Rejinold et al. examined curcumin-loaded thermo responsive chitosan NPs (TRC-NPs) for cancer treatment [33]. Their hemolysis test showed that the hemolytic ratio was below 5%, which is considered safe according to ISO/TR 7406 standards. He tested MSNs-RhB for blood compatibility by analyzing hemolysis and coagulation over a concentration range of 50–500 mg/mL. The results, along with SEM and TEM images showing uniform, approximately 400 nm particles, indicated that MSNs and MSNs-RhB did not affect blood coagulation. These particles maintained normal coagulation functions and showed no significant aggregation of blood cells. Further, citrate-capped and gold NPs were also tested, showing acceptable levels of hemolysis [34]. These findings underscore that the blood compatibility of NPs is influenced by their surface properties, such as area, charge, and hydrophobicity, with size and structure being particularly important.[31]

2. Histocompatibility

Targeted drug delivery using nanoparticles (NPs) has made significant advances, particularly in terms of biocompatibility and diagnostic utility. Superparamagnetic iron oxides (SPIONs) are widely regarded as biocompatible, with dextran-coated variants showing minimal toxicity in human macrophages[35]. However, they can modulate cytokine production based on the type of macrophage used. Dendrimers, which are branched polymers, show generation-dependent toxicity, with higher generations and cationic surfaces being more toxic. This toxicity can be mitigated by surface modification with polyethylene glycol (PEG). Silica NPs, including those functionalized with quantum dots, have demonstrated excellent biocompatibility, making them suitable for drug delivery and biosensing applications due to their ability to enter cells without inducing cell death[36]. Gold NPs, known for their stability and high scattering power, are used in a range of applications from biosensing to drug delivery. They typically show good biocompatibility, though they can provoke immune responses when conjugated with peptides. These studies underscore the importance of NP surface modifications and functionalization in tailoring their biocompatibility and effectiveness in biomedical applications.[31]

TOXICITY OF NANOMATERIALS

The interaction mechanisms between nanoparticles (NPs) and living systems remain complex and not fully understood. NPs can bind and interact with biological matter, altering their surface characteristics based on their environment. Despite growing knowledge about NP-cell interactions, understanding intracellular mechanisms is challenging due to variability in NP behavior based on factors like surface coating, charge, and size. This complexity differentiates nanotoxicology from classical toxicology, where methodologies and dose metrics are more standardized.[31]



Toxicity varies by administration route and deposition site. A system-based approach examines lung, dermal, liver, and nervous system targets:

1. Lung Cells- NPs can affect alveolar macrophages, fibroblasts, and bronchial epithelial cells. Polymeric NPs (e.g., PLGA) have showed their potential for treating pulmonary diseases but require thorough cytotoxicity evaluations. Silica (SiO₂) NPs, though generally low in toxicity, can cause oxidative stress and protein aggregation. Silver NPs (AgNP) may cause lung inflammation and reduced function under high-dose chronic exposure[37].

2. Dermal Cells: The skin acts as a barrier, but topically applied NPs can penetrate and affect systemic circulation. Titanium dioxide (TiO₂) NPs in sunscreens show variable penetration and toxicity, with some studies suggesting limited dermal infiltration. Gold NPs (AuNP) demonstrate size-dependent skin permeation, with smaller particles penetrating deeper, but findings on their cytotoxicity are inconsistent.[38][39]

3. Liver Cells: The liver is crucial for evaluating NP toxicity due to its role in metabolism and accumulation. Quantum dots (QDs) have been studied for their hepatocellular toxicity, which may arise from metal ion release and oxidative damage. Surface coatings are critical in mitigating toxicity; insufficient coverage can lead to increased ion leakage and cytotoxicity. [40]

4. Neurotoxicity: NPs can cross the blood-brain barrier (BBB) and potentially cause neurotoxicity. Superparamagnetic iron oxide nanoparticles (SPIONs) are used in imaging but may induce cytotoxic effects such as oxidative stress and mitochondrial dysfunction. The size and surface coating of SPIONs impact their cellular uptake and toxicity. Research on SPIONs for brain imaging reveals potential neurotoxic risks that need further exploration.[41][42]

CHALLENGES

The use of nanomaterials in tissue engineering and medicine faces several significant challenges, including ensuring biocompatibility and long-term safety, as the interactions between nanomaterials and biological systems can vary greatly among individuals. This variability, coupled with insufficient long-term data, complicates the assessment of potential chronic toxic effects. Additionally, production costs for nanomaterials are often high, and achieving reproducibility in their synthesis remains difficult, leading to inconsistencies in material quality and performance. Regulatory hurdles further complicate matters; current guidelines may not fully address the unique properties and potential risks of nanomaterials, creating uncertainty in their clinical approval process. Moreover, integrating nanomaterials with complex biological systems presents challenges such as managing

unintended immune responses and ensuring that multifunctional materials perform effectively without adverse effects. Addressing these issues requires continued research, standardized testing protocols, and evolving regulatory frameworks to support the safe and effective use of nanomaterials in medical applications.

FUTURE PERSPECTIVES

Nanomaterials in tissue engineering and medicine will continue to make significant advances in the future. Smart nanomaterials have the potential to offer more targeted and regulated medicines because of their ability to respond to changes in their surroundings. It is anticipated that the creation of biodegradable nanomaterials will improve safety by lowering long-term accumulation within the body. Nanotechnology will probably help personalized medicine by enabling therapies based on unique biological and genetic profiles. Furthermore, combining nanotechnology with new domains like AI might completely change the way these materials are developed and used, enhancing both their efficacy and security.

II. CONCLUSION

Nanomaterials have become essential in driving advancements in tissue engineering and medicine, by offering new innovative therapeutic approaches. Their unique features, such as a high surface area, customizable physicochemical properties, and the ability to interact on a molecular level with biological systems, position them as excellent candidates for the development of new treatments and the improvement of existing medical practices. In tissue engineering, nanomaterials contribute significantly to scaffold design, promoting better cell adhesion, growth, and differentiation, which in turn leads to more effective tissue regeneration. Their ability to replicate the extracellular matrix and their role in controlled drug delivery further highlight their significance in developing functional tissue constructs. In the medical field, nanomaterials are applied in various areas, including diagnostics, drug delivery, and therapeutic treatments. Their use in targeted drug delivery systems helps to reduce side effects while enhancing therapeutic effectiveness. Furthermore, their role in imaging and diagnostics provides the potential for early disease detection and personalized treatment plans, revolutionizing patient care. The incorporation of nanomaterials into tissue engineering and medicine offers great potential for addressing some of today's most pressing healthcare challenges. As research continues to advance, the development of safer, more effective, and cost-efficient nanomaterials is likely to lead to groundbreaking treatments and improved healthcare outcomes overall.

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