History, progress and future challenges of artificial blood vessels: a narrative review

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From the Contents
Introduction 81
History of the Generation of Artificial Blood Vessels 82
Current Status 83
Discussion and Future Challenges 90

ABSTRACT
Cardiovascular disease serves as the leading cause of death worldwide, with stenosis, occlusion, or severe dysfunction of blood vessels being its pathophysiological mechanism. Vascular replacement is the preferred surgical option for treating obstructed vascular structures. Due to the limited availability of healthy autologous vessels as well as the incidence of postoperative complications, there is an increasing demand for artificial blood vessels. From synthetic to natural, or a mixture of these components, numerous materials have been used to prepare artificial vascular grafts. Although synthetic grafts are more appropriate for use in medium to large-diameter vessels, they fail when replacing small-diameter vessels. Tissue-engineered vascular grafts are very likely to be an ideal alternative to autologous grafts in small-diameter vessels and are worthy of further investigation. However, a multitude of problems remain that must be resolved before they can be used in biomedical applications. Accordingly, this review attempts to describe these problems and provide a discussion of the generation of artificial blood vessels. In addition, we deliberate on current state-of-the-art technologies for creating artificial blood vessels, including advances in materials, fabrication techniques, various methods of surface modification, as well as preclinical and clinical applications. Furthermore, the evaluation of grafts both in vivo and in vitro, mechanical properties, challenges, and directions for further research are also discussed.

Introduction
Cardiovascular disease is the leading cause of morbidity and mortality in the world, of which coronary and peripheral artery vascular diseases comprise the largest proportion.1, 2 Vascular grafts are mainly used for the surgical treatment of vascular diseases that require new long-term revascularisation, including abdominal aortic aneurysms, coarctation of the aorta and chronic haemodialysis access. For this purpose, a single vascular graft is taken as an autologous graft from the patient, which is commonly the saphenous vein from the leg or the internal thoracic artery from the chest wall.3 However, autologous blood vessel availability is limited and requires invasive harvesting techniques. Moreover, with passage of time, the quality of autologous blood vessels can be difficult to guarantee, and the incidence of postoperative complications is higher.4, 5 Therefore, it is of great clinical significance to obtain vascular prosthetics that can function as blood vessels in order to restore blood flow around a blockage or replace damaged blood vessels.6 Currently, various types of materials are used to prepare artificial vascular grafts, including synthetic polymers, natural materials, or a mixture of types. In addition, there is increasing interest in regard to the development of tissue-engineered vascular grafts (TEVG) among various other approaches, which are currently under investigation.7 A tissue-engineered vessel can be grown, remodelled, and repaired in vivo without the necessity for autograft surgery, producing demonstrable benefits. Vascular graft failures are generally related to thrombosis, intimal hyperplasia, atherosclerosis, or infection.8, 9
Moreover, ideal artificial vascular grafts have good biocompatibility and blood compatibility which will last until the completion of endothelialisation. Any degradation in vivo is balanced by new tissue formation, replacing lost material with native proteins. The most promising materials provide mechanical support equal to autogenous blood vessels to sustain blood pressure load and promote tissue regeneration in vivo with long-term patency. Furthermore, materials possessing the capacity for scaled production, sterilisation, and storage are preferred, enabling them to conform to existing clinical applications. Different anatomic locations for artificial blood vessels have different primary design requirements: for small-calibre or low-flow vessels, a non-thrombogenic luminal surface may be more important than for large-diameter vessels, which, in contrast, call for greater wall tensile properties to improve mechanical durability. Accordingly, this review attempts to outline the problems associated with the use of artificial blood vessels in vivo and describe past research efforts. In addition, current state-of-the-art technologies involved in the creation of artificial blood vessels, including advances in materials, fabrication techniques, various methods of surface modification, as well as preclinical and clinical applications, are discussed. Furthermore, the evaluation of grafts both in vivo and in vitro, mechanical properties, challenges, and directions for further research are also elaborated. Here, we searched related articles in the electronic databases of PubMed and Web of Science from inception to January 2022, using the terms “artificial blood vessels”, “vascular graft”, “vascular prosthesis”, and related keywords.

History of the Generation of Artificial Blood Vessels
Researchers have attempted to ameliorate the need for artificial vascular grafts in the past, as shown in Figure 1. Specifically, Austrian-German surgeon Payr experimented around 1900 with a thin tube made from absorbable magnesium, which marked the first time that a vascular prosthesis was used in humans. However, the patient died of pneumonia 3 days later due to the formation of fibrotic tissue induced by the magnesium. Dubost et al. utilised a cadaveric aortic allograft to perform the first successful aneurysm resection and graft implantation on March 29, 1951. In 1954, Blakemore and Voorhees were the first to treat 10 patients with a synthetic arterial substitute, laying the foundation for the rapid development of subsequent artificial blood vessels. Then, in 1958, De Bakey et al. presented the Dacron vascular prosthesis as an alternative to cadaveric allografts, which are prone to late complications. Most researchers have subsequently focused on various other synthetic materials, of which Dacron, polyurethane, and expanded polytetrafluoroethylene (ePTFE) have proven to be the most viable in vascular surgery. However, such materials are not suitable for use as implants when the diameter of the vessel is smaller than 6 mm due to the high risk of thrombus formation and compliance mismatch. A type of hybrid graft, prepared from polyurethane and poly-L-lactic acid, was developed in 1985 and has been demonstrated to ensure the fast development of a complete new arterial wall, possessing strength, compliance, and thromboresistance equal to normal arterial wall tissue. Eventually, techniques have become more efficient and have led to novel developments. In 1986, the first tissue-engineered blood vessel construct was produced by Weinberg and Bell. Bovine endothelial cells (ECs), fibroblasts, and smooth muscle cells (SMCs) were cultured in a collagen matrix and shaped into tubes, realising tissue architectures analogous to natural blood vessels. However, the mechanical properties of the constructs were poor and required the support of a Dacron mesh. Around this time, Argentinean surgeon Parodi, in collaboration with the Argentinian radiologist Palmaz, experimented with stents made of stainless steel and Dacron tube grafts in canine models and developed an endovascular technique for abdominal aorta repair. In 1990, the first successful human endovascular aneurysm repair was performed. Additionally, the first completely autologous engineered arteries were implanted into porcine recipients in 1999. These vessels, which were 3 mm in diameter and 5–6 cm in length, demonstrated patency and satisfactory mechanical durability for 4 weeks.

Over the last two decades, significant progress has been made in developing various types of tissue-engineered vascular substitutes. Tissue engineering technology combines cells, tissue scaffold, and engineering in order to generate vascular grafts. Particularly, creation of small-diameter artificial blood vessels has progressed well in terms of employing tissue engineering techniques through different approaches, such as artificial blood vessels using biodegradable polymers as scaffolds or consisting of decellularised vascular tissue. Meanwhile, novel approaches have been used to construct TEVGs for clinical use and these have been demonstrated to have many advantages, such as three-dimensional (3D) bioprinting, which allows for the use of autologous cells alone without any scaffolding in the preparation of different sized small-diameter artificial blood vessels. Nevertheless, few comprehensive studies have been conducted pertaining to artificial small-calibre vascular grafts that possess long-term patency in vivo. In addition, an ideal tissue-engineered vascular graft has yet to be made clinically available. Evaluating the next generation of grafts encompasses resolving complex and multifaceted issues via a combination of mechanical engineering, vascular biology, and immunoregulation, which continue to pose a great challenge.
**Figure 1.** The generation of artificial blood vessels. A few significant time points in the generation of artificial blood vessels and the main research areas are shown. ePTFE: expanded polytetrafluoroethylene; PCL: polycaprolactone; PGA: poly(glycolic acid); PLA: polylactic acid; PLCL: poly(L-lactide-co-caprolactone). Created with BioRender.com.

### Current Status

#### Main complication and failure mechanisms

Vascular blood vessels made of synthetic polymers have been widely applied in cardiovascular disease treatment. Nevertheless, failures continue to occur, and failure to replicate key elements of the natural vasculature (Figure 2) along with the mismatch of biomechanical properties may further cause multiple complications.28, 29 Accordingly, it is essential to understand the most common failure modes in order to facilitate improvements in the materials' design (Figure 3).

Vessel graft failures are most commonly caused by restenosis and thrombus formation and always occur secondary to the progression of atherosclerotic disease or the development of intimal hyperplasia.30 In addition, infection also serves as the principal cause of late failure in grafts, in which the inflammatory response to infection can promote restenosis.31 In one case, an aneurysm developed in a Dacron prosthesis implanted in the femoral artery, with this complication occurring about five and a half years after insertion.32

Preventing vascular graft occlusion due to endothelial hyperplasia would enable grafts to maintain their patency for a long period. There are three main causes of intimal hyperplasia, namely, intravascular SMC migration from the media to the intima, hyperproliferation of fibroblasts and SMCs, and extracellular matrix (ECM) deposition. A strong relationship is known to exist between haemodynamic factors and the development of intimal hyperplasia, which may develop in the graft vessel or the native vessel around the anastomosis.13 In addition, multiple other factors, such as mismatch of vessel diameter, trauma during surgery, and low blood flow may also serve as causes.34–36 Damage or lack of ECs lining the graft lumen results in thrombosis, which can cause the adherence of blood proteins as well as the activation of clotting mechanisms.37 The progression of atherosclerosis limits the long-term utility of grafts.38 Atheroma formation in artificial grafts is comparable to that in native arteries. Specifically, the development of atherosclerotic plaques is caused by the invasion of monocytes into the vascular neointima to form macrophages and eventually foam cells.39 Graft infections are more common in synthetic grafts, and foreign bodies are susceptible to bacterial colonisation. Such infections induce chronic inflammation, where bacteria release toxins that interfere with graft healing and may eventually result in sepsis, anastomotic failure or even rupture.40–42

Biodegradable materials are regarded as an auspicious choice for use as artificial blood vessels as the graft will be replaced by tissue ingrowth through the pores of the graft. However, this kind of reconstruction depends on the counterbalance between the degradation of the materials, ingrowth of host cells, and replacement of the materials by host tissue.43, 44 Moreover, aneurysms or even vascular rupture may develop if the degradation rate of the degradable material does not match the rate of vascular tissue regeneration. Another such issue is that SMCs may over-proliferate, causing restenosis in the later period after implantation while the endodermis of the artificial blood vessel grafts is not complete.45 These problems make it difficult for grafts to completely carry out model vascular tissue regeneration and adequately preserve biocompatibility in vivo.

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| Year | Event |
|------|-------|
| 1950 | Natural materials: Silk, ePTFE (Gore-Tex) |
| 1955 | Synthetic materials: Polyethylene terephthalate (Dacron) |
| 1960 | Natural materials: Silk, Synthetic materials: Polyethylene terephthalate (Dacron) |
| 1965 | Hybrid materials: PU + poly-L-lactic acid (PLLA) |
| 1970 | The first TEVG |
| 1975 | Polyethylene glycol (PEG) |
| 2000 | Bovine SMCs-collagen |

**Figure 2.** Artificial blood vessel: history, progress, and challenges
Figure 2. Typical structure of normal human arteries and veins. Created with BioRender.com.

Figure 3. The challenges after vascular graft implantation. After implantation, insufficient endothelialisation and aggressive proliferation of SMCs over time can lead to IH and thrombosis. Inflammatory cells play an important role in regulating the functions of ECs and SMCs. EC: endothelial cell; EPC: endothelial progenitor cell; IH: intimal hyperplasia; SMC: smooth muscle cells. Created with BioRender.com.

Common materials

Synthetic polymers

Synthetic biomaterials are produced from polymers derived by chemical synthesis. They have exact chemical structures and allow controlled adjustment to produce excellent mechanical properties. Owing to the simplicity of carrying out polymer synthesis, its mass reproducibility, and rapid availability, polymer-based synthetic vascular grafts have become an attractive option.46, 47 Most of these grafts have been commercially available since the 1970s and are often made of ePTFE, polyethylene terephthalate (e.g., Dacron), or polyurethane, which are all non-degradable synthetic materials. Particularly, Dacron and ePTFE have been involved in manufacture of large-diameter artificial vascular grafts for over 50 years.48 Vascular grafts based on biodegradable materials may be more appropriate for use in clinical applications, especially for the generation of small-diameter blood vessels. Regenerated vascular tissue guarantees the same function as autologous blood vessels when the grafts degrade.49, 50 Polycaprolactone
Artificial blood vessel: history, progress, and challenges

(PCL) is an US Food and Drug Administration (FDA)-approved biodegradable polymer that has been extensively applied to manufacture tissue engineering scaffolds, particularly its nanofibre form via electrospinning. Synthetic polymers endow strength to a biomaterial. However, they generally have poor cellular compatibility, with the majority of the degradation products having been shown to elicit adverse immune responses in vivo. They also have limited resistance to thrombus formation on the surface, which leads to obstruction of the grafts. In this regard, in order to further enhance the potency of synthetic polymers, a tactical approach must be adopted to modify their surface.

Natural biomaterials

In addition to the above-mentioned synthetic materials for vascular grafts, natural biomaterials are also widely used. They are usually extracted from plants or animals and include proteins, polysaccharides and proteoglycans. In general, classical components of mammalian ECM, such as collagen and elastin, as well as non-mammalian macromolecules, such as silk, cellulose, and chitosan, have all been shown to be of great value for use as materials for artificial blood vessels. The aforementioned biomaterials have better biocompatibility compared to that of synthetic materials, while some have been found to have beneficial properties for vascular applications when engineered using appropriate manufacturing techniques. The physical properties of natural polymers have improved with the advancement of new separation, purification and manufacturing processes so that it is feasible and increasingly attractive to use them alone.

Silk fibroin extracted from natural silk is extensively used in the field of biomedicine because of its special biodegradability and biocompatibility. Moreover, it has been demonstrated to promote faster endothelialisation of the vascular graft compared to ePTFE. Collagen is the most bountiful protein in higher animals and may be easily obtained in large quantities from pigs, sheep, and fish. In addition, gelatine can also be obtained cheaply, as it can be sourced from hydrolysed collagen. Likewise, elastin, a crucial constituent of the vascular ECM, is also derived from animals, though in smaller quantities.

One shortcoming of natural polymers is their easy degradation. Therefore, finding new materials that are not easily degraded in natural systems may act as an effective solution. Bacterial cellulose (BC) is a polysaccharide produced by microorganisms, primarily Komagataeibacter xylinus, which is resistant to degradation in the human body due to the absence of cellulases. The potential of using BC materials alone to fabricate vascular grafts has been demonstrated because of their unique properties including high crystallinity, purity, good water retention, adequate mechanical properties, and enhanced biocompatibility. Long-term (9 months) preclinical studies in sheep have verified that BC confers high stability and low thrombogenicity in vivo and, according to the degree of cellular infiltration and ECM deposition, the strength of grafts may increase over time. Li et al. showed that utilising the shape-memory property of BC membranes can serve as an approach to promptly fabricate multi-layered artificial blood vessels. Notably, small-calibre bacterial nanocellulose (BNC) conduits composed of a nanofibril architecture possess greater potential in the creation of small-calibre vascular grafts. This is especially true of BNC hydrogel and air-dried BNC due to their three-dimensional network structure, which is similar to that of natural ECM, as well as their capacity to regulate cell growth, proliferation, and differentiation. It has been reported that air-dried BNC conduits remain patent after replacement of the carotid artery in rabbits 46 days later, exhibit superior properties, and possess manifestations convenient for surgical applications as small-diameter blood vessels. However, further development of naturally-derived graft substitutes in the future has continued to be limited by their weak mechanical strength, the complexity of design, difficulties in out-of-lab translation, and the presence of complications following implantation. Further efforts in regard to in vitro and in vivo studies are therefore indispensable so as to improve grafts made from natural biomaterials.

Hybrid materials

As previously discussed, synthetic polymers have disadvantages in terms of their use in grafts as they often cause thrombosis, intimal hyperplasia, calcification, and have poor haemocompatibility and regenerative potential, although they exhibit excellent mechanical properties. Few natural materials have adequate mechanical properties for use as stand-alone materials. Natural materials have excellent biocompatibility; however, they are prone to causing vascular graft dilation and aneurysms (Table 1). In such cases, interest in hybrid materials has increased, since they constitute a reliable combination of natural and synthetic polymers and possess greater strength. Several pre-clinical studies have mixed natural materials with synthetic polymer materials in order to prepare artificial small-calibre vascular grafts. The properties of such composite materials have always been shown to be improved and balanced. Grafts made from PCL, a widely-used synthetic polymer in vascular graft research, typically degrade by 70–80% of their original weight by 18 months after implantation, which has been demonstrated in an abdominal aorta rat model. However, the quality of the regenerated tissue is insufficient in the long term and the tissue formed by collagen, myofibroblasts, and macrophages at earlier time points does not persist, which is detrimental to the long-term performance of a vascular graft. The addition of chitosan to PCL grafts has been shown to improve remodelling, resulting in a lumen fully covered with endothelium and organised contractile SMCs, which simultaneously increase the resistance to thrombosis in sheep models. Moreover, only 9.1 ± 5.4% of the original PCL/chitosan grafts have been shown to remain after the 6-month evaluation period, indicating that chitosan addition greatly accelerates degradation. Hybrid PCL collagen grafts have also been demonstrated to reduce the accumulation of oxidised lipid species and increase contractile SMCs when implanted in vivo.
Review

Tissue-engineered vascular grafts (TEVGs)

Notably, large-diameter (inner diameter > 8 mm) artificial blood vessels made of ePTFE, Dacron, silk, polyurethane, or other polymers have been successfully used in clinics for conditions such as coarctation of the aorta, aneurysm of the thoracic aorta, and aortic dissection.\(^{78}\) To date, treatment of abdominal aortic aneurysms primarily comprises two different surgical procedures: endovascular placement of an aortic stent graft and open surgical repair of abdominal aortic aneurysms. Most current stent-grafts are based on self-expandable nitinol stents made of Dacron or ePTFE fabric.\(^{77,78}\) However, in most cases they fail to replace vessels of a small diameter (inner diameter < 6 mm) due to thrombosis, atherosclerosis, infections, or neoimal hyperplasia\(^{62}\) (Table 2). In view of the limitations of current vascular substitutes for small-diameter arteries, TEVGs may be an attractive potential solution.

A TEVG is made using appropriate materials and processes that possess high tissue compatibility, leading to a lack of rejection and maintaining long-term patency following transplantation.\(^{79}\) Due to these advantages, tissue engineering is frequently used for the construction of artificial blood vessels. Both synthetic and natural biodegradable materials have been tried in vascular tissue engineering. The scaffolds are considered to possess high porosity as well as a microstructure that can deliver, align, and maintain cell connections in favour of angiogenesis.\(^{80}\) The scaffold material required for TEVG creation should be biodegradable and nonimmunogenic. Furthermore, material processing techniques often affect the function of 3D scaffolds, as the same material composition with distinct micro/nano-scale structures can eventually perform different functions. To date, a number of fabrication techniques have been adopted to create clinically-viable TEVGs which vary widely in terms of materials, manufacturing methods, cell sources, and culture protocols. In a word, these strategies can be broadly classified into three common approaches: decellularised natural matrix methods; scaffold-based approaches using synthetic or natural materials; and scaffold-free processes.

Decellularised vessels

One type of currently-used artificial vascular vessel that has demonstrated satisfactory preclinical and clinical results is made by decellularising allograft and xenograft vessels, such as arteries from swine or bovine models or the human umbilical artery.\(^{67,81}\) Decellularisation is a tool that can be used in the generation of acellular scaffolds for tissue engineering, which completely removes all cellular and nuclear material from tissues with positive outcomes.\(^{82,83}\) This outcome can be achieved through physical, chemical, or enzymatic methods.\(^{84}\)

### Table 2. Advantages and disadvantages of artificial blood vessels in different materials

| Vessel type     | Materials                     | Advantages                                               | Disadvantages                                                   | References |
|-----------------|-------------------------------|----------------------------------------------------------|-----------------------------------------------------------------|------------|
| Synthetic polymers | ePTFE, Gore-Tex, PET (Dacron), PU, PCL, PLCL, PLA, PGS | Excellent mechanical properties. Easy availability. Mass production. Easy surgical suturing. Preventing vascular burst. Can be stored for off-the-shelf use. | Causes thrombosis, intimal hyperplasia, calcification, and chronic inflammation. No growth potential. Poor haemocompatibility. Compliance mismatch. |
| Natural biomaterials | Silk fibroin, collagen, elastin, chitosan, bacterial cellulose | Excellent biocompatibility. Enhanced biological signalling. Tunable mechanical properties. | Weak mechanical strength. Causes vascular graft dilation and aneurysms. Easy degradation. Overly complex designs. Difficulty in translation. |
| Decellularised vessels | Animal artery, umbilical artery, umbilical vein | Low immunogenicity. Preserved extracellular matrix, meso- and microvasculature. | Increased thrombogenicity. Host immune response. Difficulty in precise recellularisation. Calcification. |

Note: ePTFE: expanded polytetrafluoroethylene; PCL: polycaprolactone; PET: poly(ethylene terephthalate); PGS: poly(glycerol sebacate); PLA: poly(lactic acid); PLCL: poly(L-lactide-co-caprolactone); PU: polyurethane.

The scaffold material required for TEVG creation should be biodegradable and nonimmunogenic. Furthermore, material processing techniques often affect the function of 3D scaffolds, as the same material composition with distinct micro/nano-scale structures can eventually perform different functions. To date, a number of fabrication techniques have been adopted to create clinically-viable TEVGs which vary widely in terms of materials, manufacturing methods, cell sources, and culture protocols. In a word, these strategies can be broadly classified into three common approaches: decellularised natural matrix methods; scaffold-based approaches using synthetic or natural materials; and scaffold-free processes.

### Table 2. Artificial blood vessels of different diameters

| Artery diameter | Usage sites | Indications | Main challenges | Commercial materials |
|-----------------|-------------|-------------|-----------------|----------------------|
| Large > 8 mm    | Aortoiliac arteries. | Open aortic aneurysm repair. Coarctation of the aorta. | Weak mechanical durability. Compliance mismatch. Aneurysmal-like dilation. | Non-degradable materials: PET, ePTFE |
| Medium 6–8 mm   | Carotid artery, Femoral artery. | Similar to large | | | Similar to large |
| Small < 6 mm    | Coronary arteries. Infraparinguinal arteries (below the inguinal ligament). Infrageniculate arteries (below the knee). Coronary artery bypass. | Arteriovenous shunts. Coronary artery bypass. Peripheral arterial occlusive disease. | Stenosis/occlusion caused by thrombosis or intimal hyperplasia. Low haemocompatibility. | Autologous vessels: ITA, SV |

Note: ePTFE: expanded polytetrafluoroethylene; ITA: internal thoracic artery; PET: poly(ethylene terephthalate); SV: saphenous vein.
The resultant acellular scaffolds represent a great substrate that can promote cell adhesion, differentiation, and proliferation since specific ECM components and structures are key to tissue regeneration and remodelling. Moreover, decellularised scaffolds must have certain mechanical properties to avoid a mismatch with those of native tissues, which is key for blood vessels in tissue engineering. This can reduce immunogenicity but may also result in lower patency, which has been attributed to increased thrombogenicity, host immune response, and increased calcification. Such grafts may have a clinical role, though they have not yet been widely used. Complete removal of tissue antigens remains a major issue, and the diversity of source materials makes it difficult to achieve manufacturing reproducibility.

Scaffold-based grafts
Scaffold-based techniques use macromolecular structures to support cells during the regeneration process and are prefabrication strategies that are relatively simple to implement. Except for decellularised vessels, many methods fail to mimic the structure of the natural ECM. Electrospinning technology is a promising technique that may be used in the production of TEVGs, which can address this problem. It has been reported that nanofibres produced via electrospinning technology can simulate the composition of the ECM by loading various drugs and growth factors. Moreover, it is also the only method used to prepare nano-scale polymer material fibres directly and continuously by applying a high voltage to materials dissolved in volatile solvents so as to create dry fibres that can be collected to form conduits. Therefore, it has been extensively applied in tissue engineering, especially to create small-diameter blood vessel grafts. Additionally, the surfaces of artificial blood vessel grafts prepared by electrospinning have good specific surface areas and high porosities. This method facilitates fine control of fibre size and deposition and allows for blending of materials. New methods for simultaneous spinning of multiple components and derivative techniques, such as gel spinning and conjugate electrospinning technology, have been developed in order to optimise the technique. Small-calibre blood vessel grafts have been constructed by combining both electrospinning technology and freeze-drying technology, providing insight into the fusion of multiple processes and further improvement of artificial blood vessel fabrication.

Scaffold-free grafts
Cell-assembled grafts have been a dominant research topic over the past decade, where efforts following translational work indicate that TEVGs could be assembled entirely from human cells. Cell-assembly strategies involving culturing a range of human cell types (SMCs, fibroblasts, and induced pluripotent stem cells) for long periods of time to generate grafts are complex but the resulting grafts have the greatest degree of biomimicry. According to a study by Dahl et al., employing human allogeneic or canine SMCs grown on a degradable polymer scaffold has been shown to be a feasible method of engineering a TEVG, in which grafts in baboons were demonstrated to retain their patency for up to 6 months. These grafts have shown great potential in preclinical evaluation. However, in a Phase II study, the final results were suboptimal as only 28% of the grafts remained open at 12 months.

Essentially, TEVGs have great potential in the development of successful artificial blood vessels using several approaches. However, more research is required in order to advance the field and ultimately develop a TEVG with properties similar to that of gold standard autologous vessels.

Fabrication techniques
With developments in science and technology, the materials and structures of artificial blood vessels have been continuously improved, and the preparation processes of artificial blood vessels have also been gradually enhanced. Preparation technologies including stretch forming, electrospinning, thermal phase separation, weaving technology, 3D printing and cross-linking of different types of materials have been applied to different types of artificial blood vessels to improve the mechanical properties of grafts, optimise structure and improve product stability and repeatability.

Stretch moulding is a traditional manufacturing technique for preparing ePTFE artificial blood vessels, which has achieved wide commercial application. The process involves extruding the raw material polymer polytetrafluoroethylene resin to form a preform that is longitudinally arranged into a fibrous shape, stretched at a high speed and then cooled in order to obtain a formed product. The process of each link has a great influence on the structure and mechanical properties of ePTFE. Porosity can affect the mechanical strength and biocompatibility of ePTFE artificial blood vessels and is key to affecting the performance of ePTFE.

Electrospinning is a process in which a polymer solution or melt is used to form a small jet of polymer under the action of a high-voltage electrostatic field for spinning. Electrospinning enables the direct and continuous preparation of polymer nanofibres. Electrospinning has the following technical advantages: (1) it can increase the porosity of blood vessels to meet the requirements of cell growth; (2) it has a large specific surface area, which is conducive to cell adhesion and growth; (3) the diameter of the fibres produced is similar to the size of natural extracellular matrix or cells, hence, it can mimic the structure of human extracellular matrix to the greatest extent; and (4) it can load growth factors and induce cell adhesion, proliferation and differentiation.

Braiding is the process of weaving multiple filaments in a specific way to prepare artificial blood vessels. Preparation of artificial blood vessels by braiding technology improves elasticity and compression resistance, and enhances the compliance of the blood vessels. Due to the easy displacement and damage of the edge of a braided graft, its clinical application has been greatly hindered. Therefore, the braiding technique is often used to improve the strength of anastomosis while ensuring the flexibility of the inner layer material through the design of a multi-layer artificial blood vessel.

3D printing technology appeared in the mid-1990s and is based on digital model files, using powdered metal or plastic materials.
to construct objects through a layer-by-layer printing method. Melchiorri et al. fabricated a vascular graft adapted to the curvature of the patient’s anatomy by using poly(propylene fumarate). Although it has an adjustable shaping feature, few studies have focused on the direct preparation of artificial blood vessels by 3D printing. Specifically, most research has focused on the preparation of in vitro vascular abrasives to assist in the production of inner layer materials of artificial blood vessels and the fabrication of organ chips in order to provide support for the study of material patency.

In addition to the above-mentioned preparation methods, the production of artificial blood vessels also includes technologies such as gas foaming, hydrogel technology and freezing-casting. However, it is often difficult for a single processing method to meet the various performance requirements of artificial blood vessels, and different processing methods are often used in combination. The most common example is the combination of electrospinning and weaving techniques. Electrospinning produces filaments that adapt to cell growth and migration, while weaving improves overall vascular elasticity and compliance. In addition, Lee et al. prepared a nanofibre hybrid artificial blood vessel material by electrospinning and 3D printing techniques, which exhibited excellent mechanical properties and good patency.

Modified methods

Thus far, commercialised vascular grafts are generally only used as structural substitutes and rarely have bioactivity or biofunctions. Hence, various strategies have been used to optimise the structure of engineered tissues. In recent years, novel fabrication techniques of preparing artificial blood vessel grafts have been combined with chemical grafting, surface modification, and coating technologies to improve the patency rate of artificial blood vessels in early transplantation, especially in small-calibre grafts. Most studies have focused on surface modification of materials so as to enhance haemocompatibility and facilitate cell adhesion and growth. Therefore, the lifespan and effectiveness of a graft can be greatly extended by improving these two properties.

Reducing thrombogenicity

Thrombosis is a non-negligible cause of restenosis in vascular graft applications, especially in small-calibre or low-flow arterial bypass, and endothelial damage or foreign body anastomosis triggers an inflammatory response that activates platelets and leads to thrombotic graft occlusion. Therefore, a suitable non-thrombogenic luminal surface may be required. The surface can be either cellular or biochemical, however, it should inhibit protein and cell adsorption, prevent blood coagulation contact activation, inhibit platelet adhesion and activation, and thus avoid thrombosis in the arterial system. A study has shown that platelet adhesion is reduced when ePTFE tubes are coated with a defined mix of extracellular matrix components. In addition, the amount of fibronectin adhering to it was also observed to be reduced after blood flow assay, which indicates a decrease in its thrombogenic nature. One researcher used albumin as a coating and found that it can significantly decrease platelet adhesion compared to other plasma proteins and achieve a thromboresistant effect. As the most effective and routine anticoagulant for clinical cardiovascular surgery, heparin inhibits the occurrence of acute thrombosis following vascular transplantation. Heparin helps to promote the release of tissue factor inhibitors, synergistic anticoagulants, and plasminogen activators from ECs. Moreover, heparin has been shown to promote the growth of ECs and inhibit smooth muscle proliferation after vascular injury. In this regard, immobilised heparin can significantly influence the adherence and activation of platelets and enhance fibrinolysis. Heparinised silk fibroin can significantly enhance anticoagulation at the initial stage of implantation. Furthermore, Kuang et al. introduced a biodegradable graft made of core-poly(L-lactide-co-caprolactone)-based nanofibres coated with heparin/silk gel as a shell layer, and demonstrated that the graft was unobstructed for over 8 months via in vivo animal experiments, providing a useful model for remodelling vascular structures.

Improving endothelialisation

After graft transplantation, inflammatory and mesenchymal cells entering the artificial blood vessel wall and endothelial cells covering the endo-surface of artificial blood vessels can promote graft healing. Early endothelialisation of artificial vessels is crucial for the suppression of thrombosis and maintenance of patency. ECs can provide a suitable interface between blood and surrounding tissue while maintaining a balance between haemostasis and thrombosis. It is essential to endothelialise the inner surface of artificial blood vessels to prevent thrombosis and intimal hyperplasia. There are three ways to improve endothelialisation of the grafts’ inner surface: trans-anastomotic ingrowth of the innate arteries’ endothelia, transmural ingrowth of the capillaries at the implantation sites, and haematogenous seeding of endothelia from the circulatory system. The presence of natural materials is known to improve biological function according to consensus; therefore, various coatings based on adding natural materials can be designed. The addition of an elastin coating to grafts has been shown to reduce platelet adhesion and increase EC attachment. Interestingly, the proportion of SMCs in a contractile phenotype was also increased in elastin-containing grafts, suggesting their usefulness for limiting neointimal hyperplasia. Moreover, collagen has been shown to improve EC viability, attachment, and migration when incorporated into poly(lactic acid) knitted and poly(vinyl alcohol) vascular scaffolds in vitro. Similarly, a polymer blend of recombinant human collagen peptides and PCL gelatine have been shown to enhance the adhesion of human umbilical vein ECs (HUVECs) to the lumen, which adds further support for favourable vascular cell signalling provided by collagen sequences. Notably, increasingly complex surface coatings have been studied for their ability to promote endothelialisation through significant biological signalling while simultaneously having low thrombogenicity. Wang et al. showed that functional vascular grafts with in-situ catalysis of nitric oxide release
based on enzyme prodrug therapy are helpful for localised and on-demand nitric oxide release. This approach illustrates the effective inhibition of thrombus formation in vivo as well as the enhancement of vascular tissue regeneration and remodelling on the grafts, which may be useful for the development of novel cell-free vascular grafts in treatment of vascular disease. Surface modification of polyurethane with hydrophilic molecules improves haemocompatibility; however, this may limit initial vascular EC adhesion.110 The protein Dickkopf-3 plays key roles in inducing the directional migration of vascular progenitor cells and promoting tissue regeneration. Issa Bhaloo et al.116 proposed the construction of artificial blood vessels with the function of delivering Dickkopf-3. In addition, it has been demonstrated that histone deacetylase 7 mRNA in vascular progenitor cells can be selectively translated to produce a polypeptide containing seven amino acids. This polypeptide induces the migration and differentiation of vascular progenitor cells into vascular ECs, thereby effectively promoting the endothelialisation of artificial blood vessels.117 To date, various small animal model experiments have been conducted to improve endothelialisation on artificial vascular grafts with specific peptides and biomolecules, which has shown some potential; however, further development in larger preclinical models is lacking.

Evaluation of vascular grafts

**In vitro tests**

Two main aspects, including mechanical properties and biocompatibility, are analysed in the evaluation of vascular grafts in vitro.

The mechanical performance of artificial blood vessels is an important design requirement to ensure that they can withstand a certain pressure generated by circulating blood flowing in the blood vessel. The most fundamental performance criteria include adequate mechanical strength to retain structural integrity and resist permanent deformation. The compliance of the graft and the manner in which it deforms under pressure are also important, as adverse biological responses are associated with a compliance mismatch between native and prosthetic vascular grafts. In terms of mechanical considerations, the core issue lies in achieving high strength while retaining compliance. Tensile strength results indicated that a nanofibre-oriented structure offers a higher elastic modulus than a non-oriented structure. Likewise, other properties including tensile strength, elongation at breakage, and burst pressure were also found to be significantly enhanced by an oriented alignment of nanofibres. Consequently, the oriented structure of vascular grafts may very much enhance their mechanical properties.116 An ideal artificial blood vessel should also have appropriate micropores, which can be represented by water permeability. The size and porosity of the micropores have an effect on cell culture.118 **Table 3** shows several mechanical properties of natural arteries and various common artificial blood vessels.119

Another consideration is individual variability, signifying the variation in the mechanical properties of blood vessels from vessel to vessel and between individuals. In future, vascular tissue engineering may ultimately involve tailoring the specific mechanical properties of a graft to the intended implantation site. A far greater understanding of vascular biomechanics, interactions between the graft and the native vessels, and how to engineer the mechanical properties of a vascular graft would be required.

Biocompatibility of biomaterials is also the most basic property that should be considered when designing viable grafts, ensuring that the material has good haemocompatibility and a low likelihood of immune rejection due to problems such as allergic reactions. Several studies have shown that properly degummed and sterilised silk products have better biocompatibility than virgin silk.120 They support cell attachment, migration, proliferation, differentiation, cell-cell interactions and have the ability to promote tissue repair. ECs are derived from their progenitor cells, which are characterised by the expression of CD133.121 Biocompatibility can be evaluated by culturing HUVECs in contact with biomaterials for a period of time and observing platelet adhesion under a scanning electron microscope. Moreover, when evaluating haemocompatibility, plasma re-calcification can be used as a measure of contact phase activation and the efficacy of heparinisation following blood contact with biomaterials.122 The adhesion morphology of HUVECs can be observed by electron microscopy and performing qualitative detection of HUVECs.123 Evaluation of biocompatibility involves determining whether EPCs can proliferate well in vitro when designing tissue-engineered blood vessels. Contemporary approaches rely on in situ endothelialisation of vascular grafts, which provides an optimal native haemocompatible surface and reduces rapid failure from blood clotting; however, ensuring the biocompatibility of biomaterials remains a primary concern for tissue engineers.

| Values for artificial blood vessels12, 91 | Burst pressure (mmHg) | Compliance (%/100 mmHg) |
|------------------------------------------|-----------------------|-------------------------|
| Dog femoral artery118                     | 2895 ± 263            | 10.3 ± 2.3              |
| Human internal mammary artery119          | 3196 ± 1264           | 11.5 ± 3.9              |
| Biodegradable chitosan vascular grafts (with 4 mm inner diameter)118 | 1688 ± 236 | 5.7 ± 1.3 |
| Tissue-engineered blood vessels114        | 3490 ± 892            | 3.4 ± 1.6               |
| Expanded polytetrafluoroethylene grafts119| –                     | 0.51                    |
| Silk fibroin grafts119                     | –                     | 1.90                    |
**In vivo animal models**

In order to meet the demand to reasonably test the performance of various types of artificial blood vessels, it is necessary to establish a viable and stable animal model. Transplanting and facilitating postoperative observations may be easily performed if the appropriate animal models exist, which are able to simulate applications in the environment of artificial blood vessels. Currently, commonly-used vascular *in vivo* animal models are pigs, sheep, dogs, rabbits, and rats.124-126 The carotid, abdominal, and femoral arteries in these animal models are the most chosen surgical sites. Moreover, different options are available based on the animals’ size, according to the calibre of the blood vessel. Rat abdominal aortic replacement and rabbit carotid replacement are viable animal models for *in vivo* validation of small-calibre vessels. In addition, specific criteria, such as implantability, mechanical performance, biocompatibility, thrombogenicity, and haemodynamics, must be considered.127, 128 Achieving balance with model-specific factors should also be taken into consideration, such as animal availability, ease of handling, ease of performing the implantation surgery, study duration, compatible methods of graft evaluation, and cost. The characteristics of common animal models are summarised in Table 4.129-137

Rats and mice are widely used because they are inexpensive to acquire, easy to maintain, and can be used to evaluate larger sample sizes.135-137 Furthermore, the availability of various transgenic strains makes it possible to examine the molecular mechanisms involved in graft remodelling.138 However, differences in their circulatory system with respect to humans limit their utility in short-term studies to only very small grafts (< 2 mm diameter). Rabbits are also suitable for artificial blood vessel assessments although they are limited to short-term studies.132-134 Their physiology is similar to that of humans in terms of endothelialisation rates and thrombogenicity, and they can accept clinically-relevant graft diameters (1–4 mm). Dogs differ from humans in terms of thrombogenicity and represent a more stringent model for the investigations because of their lack of spontaneous endothelialisation.119, 131 Moreover, they can provide a range of implantation site options with varied vessel sizes, although their immune response can restrict study durations. Both pigs and sheep demonstrate similarities to human physiology and they are appropriate for studying thrombogenicity, calcification, and long-term graft patency.129, 130

The cardiovascular physiology of non-human primates, such as baboons, is most similar to that of humans compared to dogs, pigs or sheep. Their structural similarity allows for various imaging techniques and testing methods that are typically applied in humans to be used on them. Despite such advantages, they are rarely used for vascular graft evaluation due to their high cost and ethical concerns.81, 91 The optimal model for studying all of the performance criteria associated with artificial vascular grafts has yet to be elucidated. An immunodeficient pig model, the operational immunodeficient pig, has been reported to be a large animal model that allows for long-term accommodation of artificial human vascular grafts to properly evaluate the clinical efficacy of artificial vascular grafts.139

High-resolution ultrasound may be used in order to evaluate the efficacy of a graft, such as the flatness of the mouth, the vascular lumen, velocity of the blood flow in the lumen, and spectral waveform of the artificial blood vessel grafts after implantation, which can also eventually be used to assess vascular patency following implantation by contrasting with the ultrasound image taken before surgery.140 Additionally, micro-computed tomographic angiography may also be used to evaluate the patency of grafts and the specific situation of artificial blood vessels *in vivo*. Immunofluorescence staining can be used to verify biomimetic remodelling of endothelial and smooth muscle layers.123 Common evaluation methods, including tissue section staining, are shown in Figure 4.

Thus far, *in vitro* and short-term small animal studies continue to dominate, making it difficult to predict the long-term patency and appropriateness of artificial blood vessels for clinical translation. Longitudinal assessments in large animal models are necessary, but are costly and have high barriers to entry. Tables 5 and 6 summarise the artificial blood vessels in clinical trials and the various artificial blood vessels that are already on the market.

**Discussion and Future Challenges**

As previously discussed, the rise in patients with cardiovascular disease requiring replacement therapies has created a large demand for suitable vascular grafts. From an initial simple synthetic blood vessel substitute to tissue-engineered vascular grafts, much progress has recently been made, and tissue engineering techniques offer potential ideas for overcoming current vascular prosthesis replacement challenges and addressing future organ regeneration.141 Many scientists have focused on various fields related to artificial blood vessels, such as structural simulation, material improvement, performance optimisation, and tissue engineering.

Normal human arteries consist of three structural layers. The intima, media and adventitia closely fit together to carry blood flow, and this structure is closely related to the physiological function of blood vessels. In order to simulate the multilayered structure of natural blood vessels as closely as possible, researchers have improved the manufacturing process and used advanced technologies such as electrospinning, thermal-induced phase separation, microfluidics, stretch forming, and weaving technology.142, 143 Searching for the best fabrication techniques that can be effectively used to ameliorate issues related to the failure of artificial blood vessels has garnered much interest in research. 3D bioprinting technology is able to rotate and print along a rod template to obtain multiple orders of magnitude of complex three-dimensional structures.144, 145 By including a variety of modifications of the microenvironment, the overall structure of an artificial blood vessel can be made closer to the structure of the human body, which may result in better biological properties.

Materials are the basis of artificial blood vessels and also directly affect their performance. The best way to mix synthetic polymers and natural materials so as to maximise...
Artificial blood vessel: history, progress, and challenges

Figure 4. Evaluations of artificial blood vessels in vivo. (A) Macroscopic views of grafts upon implantation. (B) Macroscopic views of grafts post-operation. (C) Ultrasound image of graft implanted into rat common carotid artery. (D) Representative image of recorded angiogram showing graft patency. (E) Haematoxylin & eosin cross-sectional image. (F) Immunofluorescence image of the middle section of a vascular graft. (G) Common animal models for evaluation in vivo. EVG: elastic Van Gieson (Verhoeff’s Van Gieson); GRAFT: graft. Created with BioRender.com.

Table 4. Common animal models for evaluation of artificial blood vessels in vivo.

| Animal model | Characteristics | Application inner diameter (mm) | Common length (mm) | Longest implantation period | Implantation site | References |
|--------------|-----------------|---------------------------------|-------------------|-----------------------------|------------------|------------|
| Sheep        | Similar cardiovascular physiology, endothelialisation mechanisms and thrombogenicity mechanisms to humans. Suitable size and long-term studies possible. Higher incidence of hypercoagulability. | 4–6 | 80–100 | 9 months | Carotid artery | 48, 74 |
| Pig          | Similar vascular physiology and anatomy to humans. Well established as a model for assessing vascular grafts. Mount an extensive immune response to implanted tissues. | 3–6 | 30–100 | 6 months | Iliac artery | 130 |
| Dog          | Lack of spontaneous endothelialisation and immune response restricts study lengths. Ease of accessing vessel due to thin skin. Thrombogenicity mechanisms and vessel viscoelastic properties differ from humans. | 3–6 | 30–50 | 6 months | Abdominal artery | 131 |
| Baboon       | Physiology and cardiovascular anatomy are the most similar to humans. Suitable for a wide range of non-invasive imaging techniques adapted from humans. High cost and ethical concerns associated with using primates in medical research. | 3–6 | 30–50 | 6 months | Arteriovenous graft | 26 |
| Rabbit       | Similar endothelialisation rates and thrombogenicity mechanisms to humans. Demanding higher anticoagulant function. Limited to short-term studies. | 1–4 | 5–30 | 12 months | Carotid artery | 94, 132 |
| Rat          | Large sample size. Wide variety of transgenic lines. Allows exploration of genetic/molecular mechanisms. | 1–3 | 5–30 | 18 months | Abdominal artery | 73, 135 |
| Mouse        | Ideal for biocompatibility and cell infiltration studies. | 0.5–1 | 3–10 | 6 months | Carotid artery | 137 |
their advantages should be highlighted in further study.\textsuperscript{123, 146, 147} The selection of appropriate materials will help to improve the requisite tissue mechanics, providing good biocompatibility, anti-thrombogenicity and long-term patency rate of blood vessels. A number of people have developed methods to improve materials, mainly by synthesising new materials, combining multiple materials together, improving the properties of materials through chemical modification and using decellularised heterogeneous blood vessels. Currently, many promising structures are multicomponent systems that have been created through the addition of natural surface coatings to basic synthetic materials. Grafts with three or more components are also increasingly common (e.g., heparin/silk/poly(L-lactide-co-caprolactone). Future development trends

| Intervention/treatment | Number of patients | Condition/disease | Years | Trial ID/phase | Testing status |
|------------------------|--------------------|------------------|-------|---------------|---------------|
| Device: Synthetic vascular grafts | 207 | Peripheral arterial occlusive disease | 2010–2013 | NCT01113892/NA | Completed |
| Device: HAVG graft implantation | 40 | End-stage renal disease Kidney failure, chronic | 2012–2016 | NCT01744418/NA | Active, not recruiting |
| Device: FUSION Vascular Graft | 117 | Peripheral arterial occlusive disease (PAOD) | 2009–2013 | NCT01601496/NA | Terminated |
| Device: POSS-PCU vascular graft | 30 | Renal insufficiency | 2021–2025 | NCT02301312/NA | Not yet recruiting |
| Combination Product: Tissue Engineered Vascular Grafts | 4 | Single ventricle cardiac anomaly | 2009–2017 | NCT01034007/Phase 1 | Completed |
| Device: ProEndoTecc Vascular Graft | 33 | Peripheral arterial disease Peripheral vascular disease | 2010–2012 | NCT01095237/NA | Terminated |
| Device: ASC coated ePTFE vascular graft Device: Propaten graft | 60 | Lower limb ischemia | 2011–2022 | NCT01305863/NA | Active, not recruiting |
| Biological: natural human collagen arteriovenous graft for haemodialysis access | 10 | End stage renal disease | 2021–2022 | NCT04905511/Phase 1 | Recruiting |
| Device: POSS-PCU vascular graft | 30 | Renal insufficiency | 2021–2025 | NCT02301312/NA | Not yet recruiting |
| Procedure: Revascularisation using a BIOPROTEC graft | 45 | Peripheral artery disease | 2018–2023 | NCT04018846/NA | Recruiting |
| Device: Expanded polytetrafluoroethylene graft Device: Bovine carotid artery graft | 10 | End stage renal disease Haemolysis Arteriovenous graft | 2015–2018 | NCT03300024/NA | Terminated (Funding ended) |
| Device: Expedial vascular access graft | 172 | End stage renal disease | 2004–2006 | NCT00313872/Phase 2 | Terminated |
| Device: Covera vascular covered stent | 100 | Arteriovenous fistula | 2020–2023 | NCT04261686/NA | Enrolling by invitation |
| Device: Endovascular revascularisation of peripheral arteries Device: InnAVasc arteriovenous graft surgical implant | 150 | Vascular diseases, peripheral Kidney failure, chronic Renal dialysis | 2021–2022 | NCT04765556/NA | Active, not recruiting |
| Procedure: blood sampling procedure the vascular prosthesis manufactured by electrospinning | 120 | Arterial occlusive disease | 2014–2015 | NCT02255188/NA | Completed |
| Procedure: revascularisation Device: Propaten® Device: Crude PTFE Device: Paclitaxel-eluting graft | 228 | Ischemia lesions | 2018–2023 | NCT03430076/NA | Recruiting |
| Device: Gore PROPATEN vascular graft Procedure: disadvantaged autologous vein graft | 31 | Peripheral arterial occlusive disease | 2007–2010 | NCT00617279/Phase 4 | Terminated (study terminated due to low enrolment) |
| Device: EvoCit Device: EvoHep | 38 | Kidney diseases Haemodialysis complication End stage renal disease | 2018–2019 | NCT03887468/NA | Completed |
| Combination product: Tissue engineered vascular grafts | 24 | Cardiovascular diseases | 2020–2025 | NCT04467671/Phase 2 | Recruiting |

Note: The information is obtained from clinicaltrials.gov. NA: not available.
may include changing from synthetic materials to natural materials, or even hybrid materials, in order to produce better biocompatibility and improve the long-term patency rate.

In fact, many artificial blood vessels have already been shown to possess adequate performance, however, they are still unable to meet usage requirements in terms of long-term patency rate and thus require optimisation. Current commonly-used optimisation methods are mainly related to certain biological technologies, such as gene editing and exosomes. The optimal degradation rate of vascular grafts composed of degradable materials remains under investigation, since this process encompasses rapid degradation and a slower breakdown. Methods that help promote endothelialisation are also being gradually applied in manufacturing artificial blood vessels.

TEVG includes 3D printing, dynamic culture, and cell-assembled grafts, and has been extensively studied and served a particular role. Future development should go in the direction in which cells and materials work together to build artificial blood vessels. Essentially, in order to develop a successful, long-lasting artificial blood vessel that satisfies the growing demand, a strong collaborative effort among various forms of expertise in fields involving material science, engineering, mechanics, and biology is needed. We remain hopeful that the development of a successful artificial blood vessel that meets the requirements of an ideal graft may be introduced with the emergence of innovative technologies.
Author contributions
WW and KH conceptualised and designed the review. KH, YL, YG, CL and ZK acquired and analysed the data; KH, YL, YG prepared and finished the manuscript; WW and HY edited the manuscript; WW, KH, YL, YG, ZK, YL and HY reviewed the manuscript; WW and HY supervised manuscript drafting and determined the final draft. All authors reviewed and approved the final version of the manuscript.

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Conflicts of interest statement
The authors declare no conflict of interest.

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