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3D printing of PEEK and its composite to increase biointerfaces as a biomedical material- A review

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Abstract

Poly ether ether ketone (PEEK) is a polymer with better lignin biocompatibility than other polymers and useful biomedical engineering applications. This research summarises the outcomes of an evaluation conducted on PEEK material composites such as cellular calcium hydroxyapatite (CHAp) for medical applications. PEEK's prospects for medical implant are highlighted, and a critical analysis and review is presented of 3D printing of PEEK and CHAp and their biological macromolecular behaviours. An electronic search is carried out of Scopus database, Google search and peer reviewed papers published in the last ten years. Because of the extraordinary strength and biological behaviour of PEEK and its composite CHAp, 3D printed PEEK has several biomedical applications, and its biological macromolecular behaviour leads to health sustainability. This work highlights its biological macromolecular behaviour as bone implant material and the optimum 3D printing process for PEEK and CHAp for medical applications. The current problems with printing PEEK and CHAp are investigated along with their possible uses. Possible solutions to improve the 3D printability of PEEK and CHAp, based on scientific mechanisms, are explained. This detailed report is of benefit to both the scientific community and medical industry seeking to enhance 3D printing concepts for PEEK and CHAp.

Keyword: PEEK; 3D printing; Cellular CHAp; Biomaterial; Bone implant

1. Introduction

An implant is any medical device developed by humans to replace or assist the regeneration of the human body's biological macromolecular structure. The use of implants began with autologous implants, made of natural materials taken from other parts of the same body. Through the evolution of material engineering and the development of so-called biomaterials, it became possible to create manufactured implants. The use of biomaterials for implant manufacture saw significant growth in the 1960s [1-3]. There are several classes of implant, based on the medical speciality to which the implants relate, including orthopaedic, cardiac and dental cochlear implants. Some properties are required for all implant classes, such as biocompatibility. As defined in the Williams Dictionary of Biomaterials [4,5], biocompatibility is the ability of a material to present an appropriate response in its intended application. In addition to the general characteristics common to all implantable biomaterials, each implant class has specific requirements [6,7].

Any biomaterial in contact with the human body generates a series of responses, which should not be harmful to the implanted person's health. These responses include: cytotoxicity, if the material is cytotoxic it causes the death of cells around it; genotoxicity, if the material causes chromosomal alteration of the cells around it; and

hemotoxicity, if the material causes blood cell death [8,9]. The bio-functionality of a material is its ability to perform its function within a specified period. Orthopaedic implants, due to their primary functional property of long-term mechanical support, need to have excellent mechanical properties of both short and long durations, in addition to the characteristics previously described. Such a need for good mechanical properties initially led to the use of materials with too high an elastic modulus, such as stainless steel, to support the mechanical loads associated with the various types of orthopaedic implant such as hips, knees and spines. The use of materials with a high elastic modulus had harmful consequences, such as the stress-shielding effect [10,11]. The use of highly flexible materials leads to a loss of bone density in the regions around the implant, due to Wolf's law, published in the monograph the Law of Bone Transformation [12,13]. Wolf observes that bone density in regions of high mechanical stress is elevated more than areas with less mechanical stress. With implants with a high elastic modulus, mechanical loading is transferred from the bone structure to the implant.

Several authors report that bone remodelling is highly sensitive to dynamic loads [14,15]. Due to stress-shielding effects, it became necessary to research and develop materials with a lower elastic moduli than metallic materials, combined with high short- and long-term mechanical resistance for orthopaedic implants. Orthopaedic implants are among the most complicated implant classes because they involve almost all parts of the human body [16,17], and nearly all play a role in supporting a mechanical load. The area of spinal implants is one of the most recently developed, and can be divided practically into two sub-classes: spine prostheses, which are implants intended to preserve the functional properties of the original spine structure, that is, to maintain functionality; and implants for spinal arthrodesis, which are used for bone fusion between spine segments to correct instabilities, used if prosthesis is not feasible [18,19].

The area of spine prostheses is still in an embryonic stage of development, and the volume of use of this type of treatment is minimal. Arthrodesis treatment is the more common approach for spinal instabilities. In these procedures, intervertebral height needs to be maintained. The role of carbon nanotubes and cellular CHAp as scaffold composites in bone tissue production and regeneration is shown in Fig. 1. Height is maintained by implants known as intervertebral cages [20,21]. These devices work as mechanical shims, supporting the intervertebral space and transferring the mechanical load from one vertebra to another. The biomaterial used to produce the vertebral cage must meet a number of functional requirements. Mechanical resistance is the materials ability to undergo mechanical effort, mainly in compression but also bending and shearing. Such loads are exercised cyclically in the walking motion of the human skeleton. Therefore, the mechanical resistance of a material needs to be evaluated in both static and dynamic fatigue, and it must be able to withstand cyclic mechanical effort for at least the period over which the arthrodesis occurs. After this period, the functional properties of the product cease [22,23].

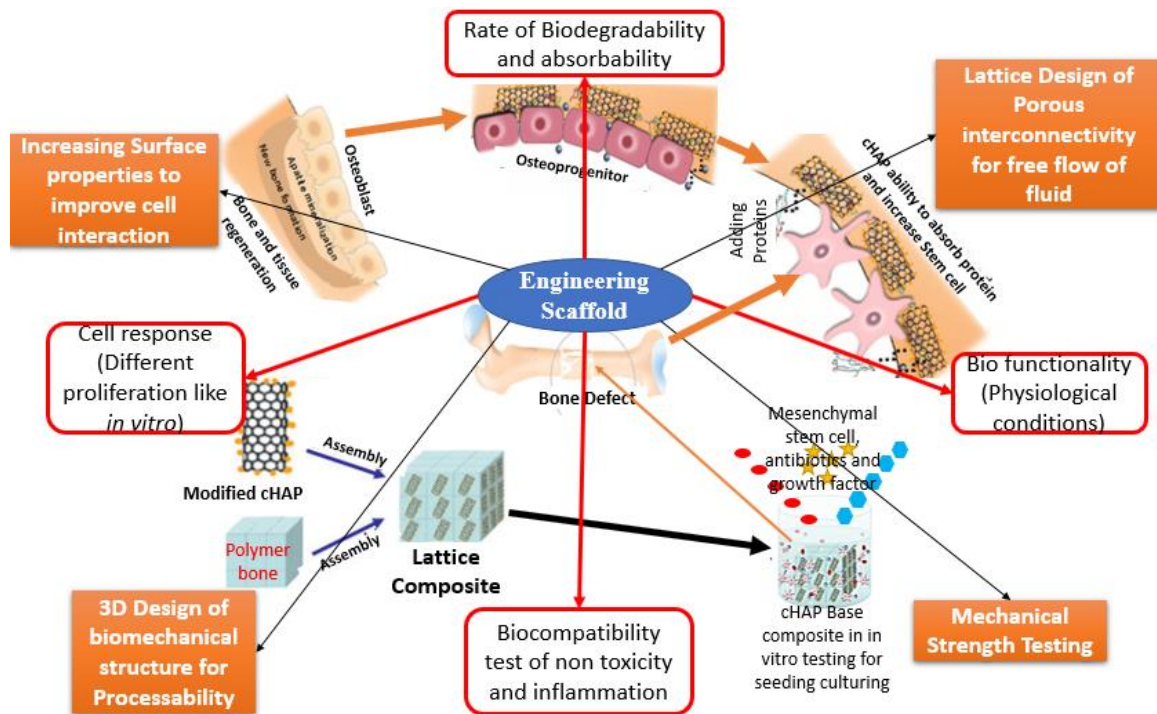


Figure 1. Cell cultures, lattices structure, and seeding for bone tissue engineering and regeneration [20-22]

Osteointegration separates two vertebrae, and maintains the intervertebral space until they are joined by bone growth. The biomaterial used must allow for bone growth around it and adapt to its new adjacent structure. Arthrodesis occurs quickly with PEEK as it is a semicrystalline thermoplastic, and therefore it is widely used for mechanical support applications such as spinal, thoracic, lumbar cervical, orthopaedic and trauma implants. It is, therefore, vital to assess the mechanical activity of PEEK in these applications in both the short and long term [24,25]. The mechanical behaviours of semicrystalline thermoplastics are strongly influenced by the degree of polymer crystallinity, the process used and the conditions of the final product [26,27]. This research is standard for the manufacture of technical parts made of semicrystalline thermoplastics used for engineering applications, including surgical implants, in order to optimise the various transformation processes, such as injection moulding, blow moulding or extrusion. The aim is to increase the degree of crystallinity and minimise internal frozen stresses in the final product, for superior performance and mechanical durability. Therefore, this research proposes combining CHAp for bone tissue engineering with state-of-the-art analysis of the 3D printing of PEEK and its composites. The work highlights the optimum 3D printing processes for PEEK and CHAp in various forms, such as powder and filament, the current printing problems and potential applications of 3D printing of PEEK for medical implants.

2. PEEK as a biomaterial

Column implants made of PEEK have relatively recently come into use. The biocompatibility and potential for biomedical applications of PEEK were first reported by Williams, McNamara and Turner [28, 29]. Fig. 2 shows the structure of the PEEK molecule. In the late 1990s, PEEK emerged as a high-performance thermoplastic to replace metal implants, and has been commercially supplied for this purpose since April 1998, pioneered by the British manufacturer Invibio [30, 31]. The first application of PEEK as a material for spinal implants was as an intervertebral cage. The first PEEK intervertebral cage, of the type posterior lumbar interbody fusion (PLIF) was

developed in the 1990s by AcroMed. Intervertebral cages made of PEEK overcome two problems caused by conventional metal intervertebral cages: the stress-shielding caused by the differential elastic modulus between human bone; and the implants being radioluminescent, which means they present artefacts under imaging techniques such as X-rays or magnetic resonance, making it impossible to visualise the bone growth in the inner region of the cage. The invention of intervertebral cells made of PEEK serves as the foundation for its current use in spinal implants [32,33]. Fig. 2 also shows the calcium-based nanomaterials such as CHAp, which are often used to build new bone scaffolds on a PEEK biological macromolecule structure with spinal electrostimulation through a longitudinal axial channel [34, 35].

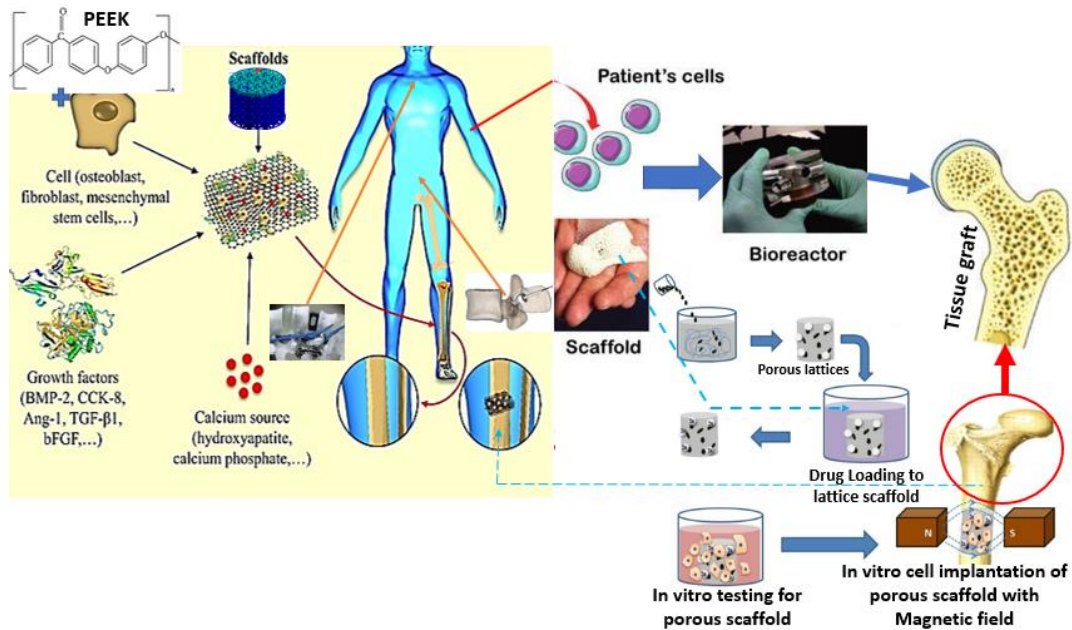


Fig. 2. PEEK biological cellular structure with Brantingan cage, electrostimulation of the spinal with a longitudinal axial channel and lumbar vertebrae[35-37]

The first clinical studies on intervertebral cages of PEEK were carried out on a group of 26 patients who were monitored over two years. Of these, 21 surgeries were successful in terms of the consolidation of arthrodesis, and the five unsuccessful surgeries failed for reasons unrelated to the cages. It was possible to follow up the surgeries using X-ray imaging, due to PEEK's radiolucency. This research demonstrates PEEK's biomechanical efficiency as a material suitable for intervertebral cages, and the ease of post-surgical follow-up. The osteointegration capacity of PEEK has been compared to titanium, a metal used in traditional metal cages. Researchers [35-37] have compared PEEK to titanium substrates using various manufacturing processes, such as bar machining and injection moulding. The authors verify, in vitro, that the adhesion of osteoblasts of bone cells is almost the same, with titanium having a slight advantage over PEEK. Injection moulding of PEEK shows results slightly superior to machining [38,39].

2.1. Biological macromolecules of cellular CHAp and its application

Unlike synthetic CHAp, biological cells of CHAp have several impurities. These impurities come about because the apatite structure allows for the replacing of a series of ions present in the human body, such as sodium and potassium. A comparative table of the composition of biological macromolecules and synthetic CHAp [40, 41] is presented in Table 1. Several biocompatibility studies have been conducted to show the biocompatibility of synthetic biological macromolecules of CHAp [42- 44]. CHAp powder used in implants is standardised by the

International Organization for Standardization (ISO), under ISO 13779-1 of implants for surgery - CHAp - Part 1: Ceramic CHAp. The standard defines the minimum percentage of powder crystallinity, at least 95%, the Ca/P ratio from 1.65 to 1.82 and the maximum trace element percentage, as shown in Table 1 [2-48]. The coating method has special rules for the essential powder properties, such as Ca/P ratio and crystallinity.

Table 1. Constituent chemical elements between biological macromolecules and synthetic, and limited heavy metals [42, 48].

Constituents (fraction by weight) %	Biological CHAp	Synthetic CHAp	Element	Maximum limit (mg/kg)
Ca	24.50	39.6	Arsenic	3
P	11.50	18.5	Cadmium	5
Na	0.70	Trace	Mercury	5
K	0.03	Trace	Lead	30
Mg	0.55	Trace		
CO ₃ ²⁻	5.80	-		

2.2. Thermomechanical behaviour of CHAp

Surface engineering modification processes usually involve high temperatures. The plasma deposition process, for example, requires 600°C in the torch [49-51]. At such temperatures, thermal decomposition can occur, altering the phase balance of the CHAp particles, changing parameters such as the crystalline structure, composition and phase morphology. It is widely accepted [51,52] that heating CHAp has three consequences: water evaporation, dehydroxylation and decomposition. CHAp is highly hygroscopic, so the evaporation of water occurs when heating the CHAp powder due to the adsorbed water. Dehydroxylation occurs in a part of the apatite structure as CHAp gradually loses the hydroxyl (-OH) group. Decomposition only occurs beyond a certain temperature, as CHAp maintains its crystalline structure during dehydroxylation and rehydrates itself on cooling [53,54]. However, upon reaching a critical temperature, irreversible decomposition occurs, leading to the formation of other calcium phosphates, such as beta-tricalcium phosphate (β -TCP) and calcium tetrakisphosphate (TTCP), both of which turn into calcium oxide. The reactions are given in Eqs. (1--3) [55, 56]. The temperature effect was presented by Livingstone [57,58], according to Table 2.

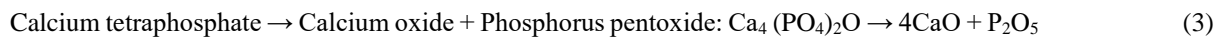
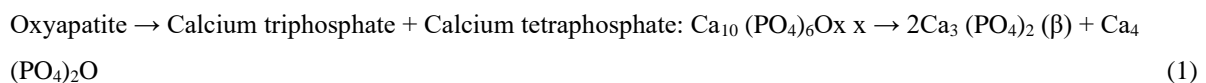


Table 2. Thermal effects of CHAp plasma [57,58].

Effect	Temperature (°C)
Evaporation of absorbed water	25 - 600
Decarbonisation	600 - 800
CHAp dehydroxylation, forming	800 - 900
Partially dehydroxylated or completely dehydroxylated	1050 - 1400
AH decomposition, forming β -TCP	< 1120
TTCP	1120 - 1470
β -TCP is stable	1550
β -TCP is converted to α -TCP	1630
CHAp fusion temperature	1730

2.3. Thermal processing of CHAP

Thermal spray processes are processes in which particles are accelerated and deposited in a molten or viscous state on a previously prepared substrate. Several thermal spray techniques are currently available. The process involves a pistol which fires melted particles at high temperatures, up to 600°C, in a plasma state, at high speed towards the substrate [56-58]. The development of this technique began in the 1960s, with further improvements, such as robotic systems, introduced in the 1980s. The thermal energy of the process is provided by high-energy plasma formed inside a gun which consists of a cathode electrode and an anode nozzle separated by a small void. Through the application of direct current to the cathode, an electric arc is formed between the anode and the cathode, with argon, helium, hydrogen or nitrogen gas ionised at high energy levels to create a plasma flame. This flame is highly unstable and recombines to form a gas, releasing large amounts of thermal energy [59, 60]. The speeds obtained can reach up to 2300m/s.

During the thermal process, particles are fed into the gun flame by the ionising gas, and accelerated towards the substrate. The high temperatures between the gun and the substrate exposed to the particles causes them to fuse. The melting point depends on a series of parameters, such as the flame temperature, the particles' location in the flame, the velocity and the particle size. The impact of CHAP particles on polymeric substrates, such as PEEK, can negatively impact the material's fatigue properties. The state of the particles, fused, semi-fused or solid, can generate micro-cracks on the PEEK surface. This could be critical, due to its high sensitivity to stress-concentrating effects. High process temperatures and the particle/substrate impact temperature can also be harmful to the material's resistance to mechanical stress in fatigue. Very little scientific information about this influence on the long-term mechanical properties is available [61-64].

2.4. Other Polymers

Due to their structure, other types of synthetic polymer can be biodegraded in vivo, and degradation by-products can be reabsorbed by metabolic pathways. This property, together with their processability using various techniques such as additive manufacturing, and their relatively good mechanical properties are the main advantages of synthetic polymers. The main disadvantage is their low cellular affinity, which makes it challenging to colonise structures, especially in the early stages after implantation. Polylactic acid (PLA) is a biodegradable synthetic aliphatic polyester obtained by the polymerisation of lactic acid. It has high mechanical resistance, biocompatibility and adequate processability. There are many studies of its potential for structures with bone regeneration applications [60, 61]. A schematic diagram of the cell growth in lattice PEEK and CHAP scaffolds is presented in Fig. 3.

One key restriction of this material is the accumulation of its degradation by-products, which are acidic in nature and can cause inflammation of the surrounding tissue. Polyglycolic acid has worse mechanical properties than the PLA, so its use in bone regeneration is limited. However, its copolymer with polylactic acid (PLGA) is a material with an accelerated degradation rate compared to pure PLA, useful for specific applications such as cranial implants in paediatric cases.

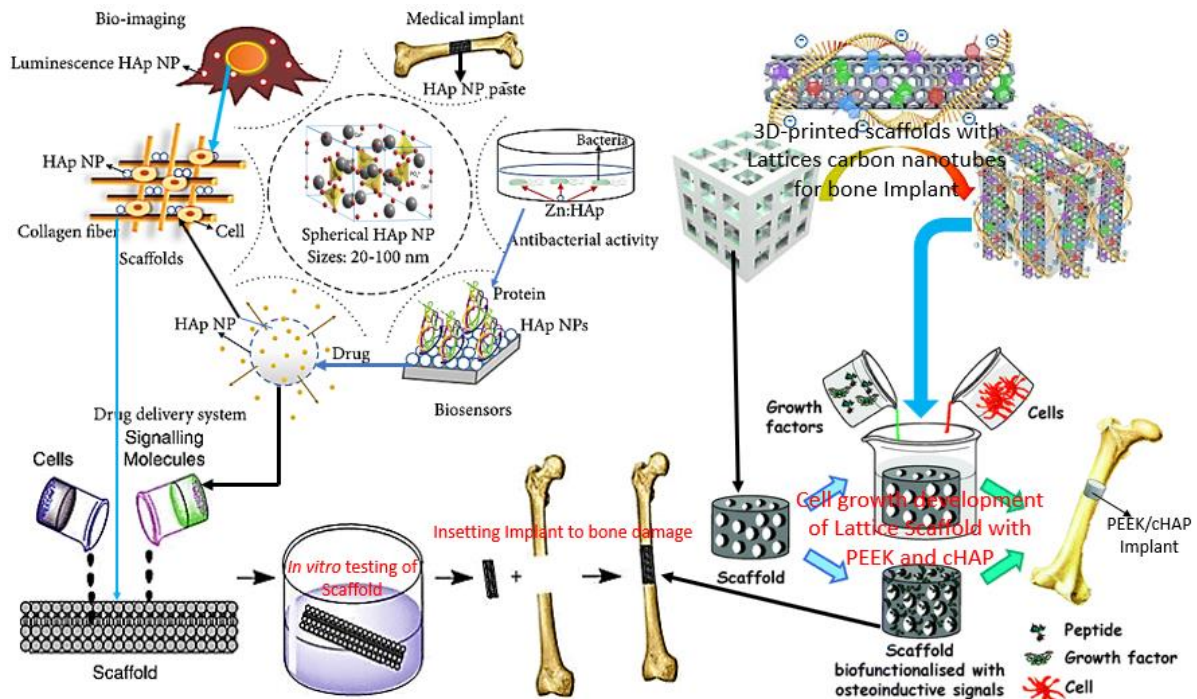


Fig. 3. Cell growth in lattice scaffold with PEEK and CHAP nanoparticle coating to create effective bone-implant Biointera [42-44]

3. PEEK polymer

Despite its macromolecular rigidity, PEEK has considerable flexibility and can withstand high plastic deformation levels in traction and compression. Its stress-strain behaviour exhibits a clear flow transition, as shown in Figs. 4(a) and 4(b), from both tensile and compression tests [65-68]. PEEK exhibits yield stress in compression about 30 to 40% higher than in tension. PEEK is an aromatic thermoplastic that has a high glass transition temperature (T_g) of 143°C, due to the high rigidity of its macromolecular skeleton. However, its planar zig-zag molecular conformation, despite its high macromolecular rigidity, allows the polymer chains to be arranged in crystalline and amorphous domains. Consequently, PEEK has a crystalline melting point (T_m) of 343°C. This implies that processing temperatures can vary between 370 and 400°C, depending on the polymer's molar mass and the type and processing conditions used in manufacturing [69-72], and PEEK's short- and long-term mechanical properties are significantly influenced by its transition temperature and melting point. The properties of stiffness, strength and toughness of the polymer at in-service temperature have increased the relevance of PEEK in surgical implant applications. Figs. 4(a) and 4(b) show the mechanical behaviour of PEEK under creep in slow deformation as a function of test temperature, in the form of short duration isochronous curves of 100s and curves of creep modulus. The properties of the elastic modulus of short (Fig. 4a) and long (Fig. 4b) duration of PEEK change minimally in the temperature range between 20 and 80°C [73-75].

As shown in Fig. 4, the use of a higher mould temperature of 200°C and high injection speed leads to a higher degree of crystallinity of 35% and excellent uniformity throughout the PEEK mould's thickness. These same injection conditions with an increased mould temperature contribute to a higher degree of crystallinity. This gives excellent crystallinity uniformity in various regions in the mould, as shown in Fig. 4. In summary, the use of

slower cooling rates, with higher mould temperature of 200°C, allows PEEK injection moulds have a higher degree of crystallinity of 30-35%, producing excellent uniformity of crystallinity with less variation of crystallinity in both the thickness and length. PEEK has a crystallisation interval well above room temperature (T_g of 143°C and T_m of 343°C), and the higher mould temperature is above the polymer T_g. The longer the polymeric melt resides in this interval, the greater the degree of polymer crystallinity [76,79].

As shown in Fig. 4, several authors have investigated the effect of injection speeds between 5.2 and 23.2 cm²/s on the maximum degree of crystallinity and crystallinity variation across thickness and duration of PEEK. Again, as shown in Fig. 4(a), using the highest mould temperature of 200°C, the maximum degree of crystallinity can be achieved, and the uniformity across the thickness is practically identical for high and low injection speeds. There is, however, a small advantage to higher injection speeds. Low injection speeds combined with low mould temperatures do not allow the mould cavity to be filled at the highest mould temperature of 200°C, as shown in Fig. 4, when the degree of variation of crystallinity along the length of the mould is evaluated. Therefore, a combination of a higher mould temperature of 200°C and a higher injection speed of 23.2 cm³/s is the moulding condition that allows the maximum degree of crystallinity. This brings excellent uniformity of crystallinity, both through the thickness and length of the PEEK mould.

The same authors evaluate the parts retention time in the mould cavity. They demonstrate that a mould temperature of 150°C, which is slightly above The PEEK T_g, achieves both full crystallinity and crystallinity uniformity. The thickness and length of the mould tend to increase with the retention time, as shown in Figs. 4(a) and 4(b). Therefore, for a higher mould temperature, above the T_g of semicrystalline thermoplastic, the shorter the time needed to reach a certain degree of crystallinity and uniformity throughout the thickness and length of the moulded PEEK part [86-88].

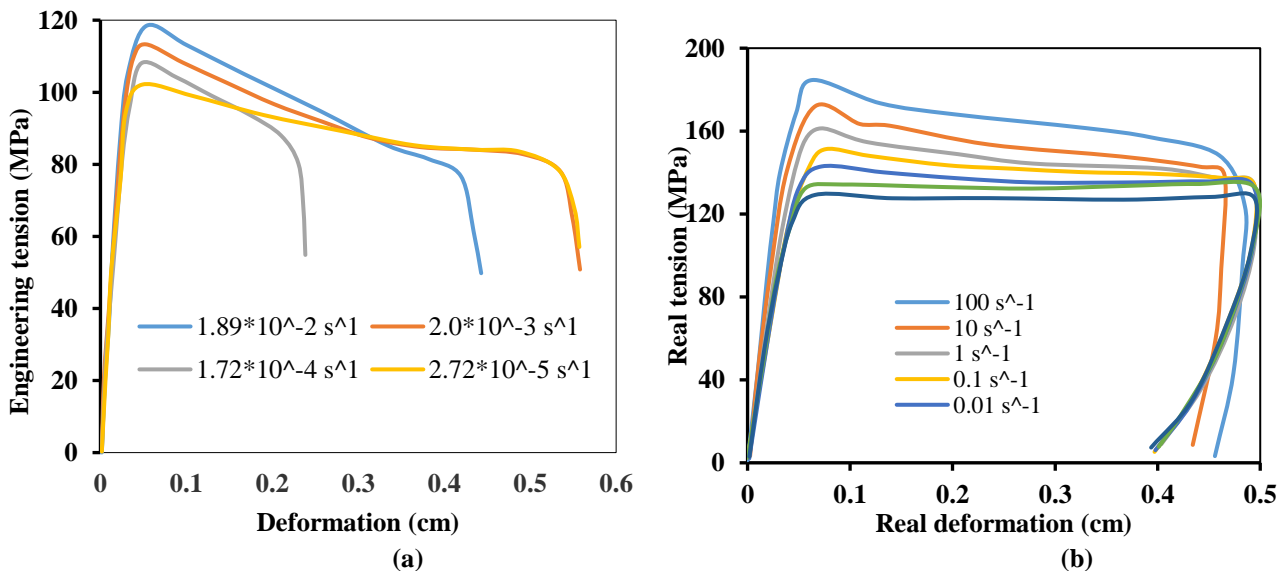


Fig. 4. Stress-strain curve of PEEK in (a) tension and (b) compression for different strain rates

The long-term mechanical stiffness of PEEK is minimally influenced by temperatures that do not exceed 400°C. The use of PEEK in surgical implants does not exceed this limit, since human body temperature is 37°C, and the long-term mechanical stiffness behaviour of PEEK remains unchanged in these applications in the direction perpendicular to the injection-moulded flow at 90°C [77-79]. The data in Figs. 4(a) and 4(b) correspond

to the creep module property under static mechanical loading, but the long-term mechanical force of PEEK is influenced by the type of mechanical loading which its products are subject to under specific service conditions. Comparing the two fatigue curves under static and dynamic loads shows that the resistance under dynamic fatigue is always lower than the resistance under static fatigue. The fabrication methods for producing scaffolds using 3D printing and the hierarchical structure of bone indicate that various elements are nanostructured CHAp embedded in collagen fibrils produced from osteoprogenitors, osteoblasts, osteoclasts and osteocytes. These are the major cellular constituents of bone contained in the combination of CHAp and PEEK (fig. 5).

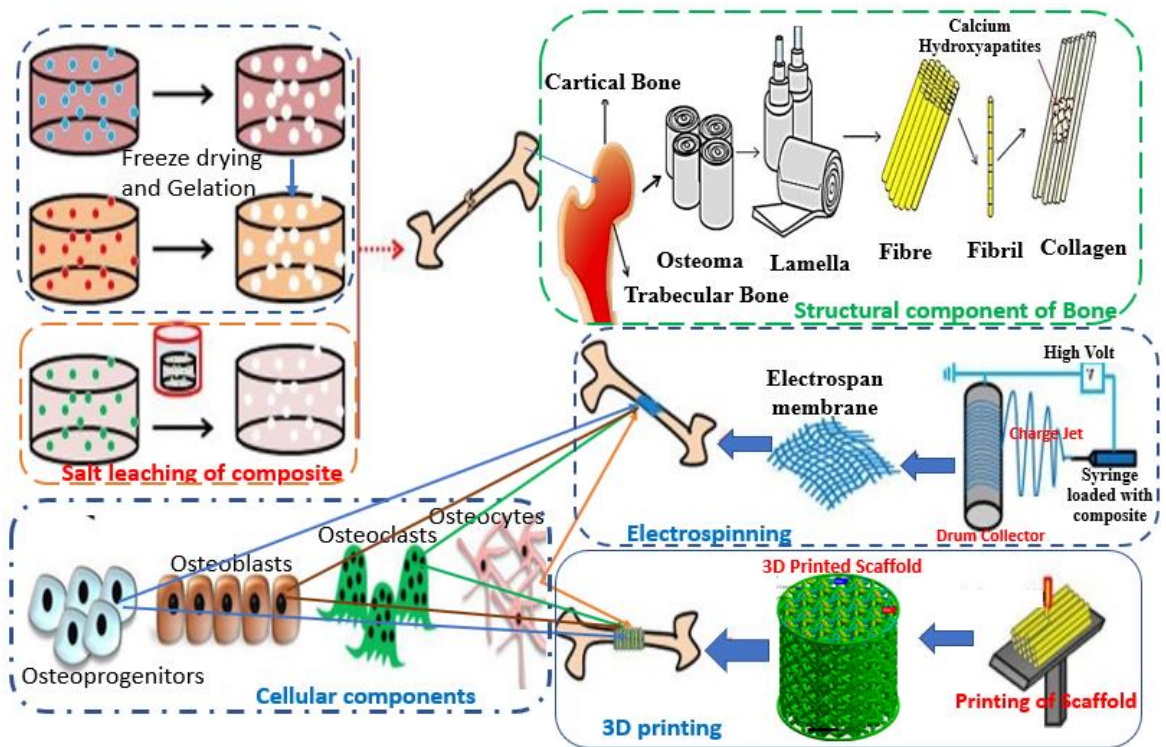


Fig. 5. Hierarchical structure of bone and 3D printed scaffolds [50-52]

PEEK applications are subject to mechanical loading under a combination of static and cyclic efforts. In surgical implants for the human spine, the dynamic fatigue resistance curve should always establish the stress limits in structural design calculations. The endurance limit recorded for a specific type of PEEK is approximately 65MPa, depending on the non-linear viscoelastic nature of the polymeric material and its rigidity susceptibility to mechanical hysteresis [81-83]. Other conditions of the fatigue test, such as type and frequency of the request wave, request geometry, test temperature and other relevant parameters, contribute to the expectation that the fatigue strength may change. Considering the failure of plastic materials under mechanical cycling almost always occurs due to crack propagation or fragile failure, it is common to stipulate the design stress value which incorporates a safety factor above the fatigue resistance limit [84, 85]. As demonstrated in the collected figures and tables, the assessment of PEEK and CHAp's mechanical and thermal-mechanical efficiency is inferred to be closely connected with its crystallinity and uniformity throughout the thickness and length of structural parts.

3.1. PEEK influence and processing conditions

PEEK implants are mainly manufactured in two types: machine-extruded bars or direct injection moulding of

implants. Implant manufacturers widely use parts machined from extruded bars, because they use the same machinery as the manufacture of traditional metal implants, such as machining and turning centres. However, the high cost of implant grade raw materials leads companies to use injection moulding in order to reduce raw material consumption. The injection moulding process results in greater productivity in the manufacture of complex shaped implants, but, as explained, the mechanical and other physical properties of PEEK parts are influenced by the level of crystallinity and crystalline morphology.

3.2. PEEK implants compared to others

PEEK has better corrosion resistance than stainless steel, but the mechanical features of PEEK are not optimal for the production of implants. Steel is cheaper, but less resistant to variations in heat, meaning that during daily use, the fact that the implant efficiently heats and cools negatively affects the patient's life. When the patient is subjected to X-rays, the steel diffuses the radiation, making treatment more complex in terms of doses and targets, when, for example, attacking a cancerous tumour. An implant made of 3D printed PEEK has properties very close to those of human bone in terms of mechanical rigidity and elasticity [87, 88]. PEEK implants synchronise better with the internal movements of bones. Unlike stainless steel, PEEK has insulating properties and therefore undergoes less heat variation. One of the strong points of printed PEEK implants is that they are transparent to X-rays. Table 3 shows methods which can benefit PEEK-based biomedical components and shows some examples of additives referenced in the literature.

Table 3. Methods of improving PEEK's quality characteristics

Materials	Major Property	Methods	Results and findings	Ref.
PEEK/Alumina	Mechanical	Composite fabrication	PEEK/Alumina improved the dynamic strength by 78%.	20-22
PEEK/HAp	Mechanical		Due to good interactivity between HAp and PEEK, the tensile strength of the composite was increased.	22-24
PEEK/Carbon nano-tubes	Mechanical and crystalline		The reinforcement improved the mechanical characteristics and decreased PEEK crystallisation rate.	24-27
PEEK/ β -TCP	Biological cell		The growth rates in β -TCP/PEEK osteoblast cells were below PEEK.	27-30
Titanium	Biological	Electron beam deposition	In-vivo studies have shown the enhanced contact ratio between bones and PEEK implants due to titanium coating.	30-32
PEEK/ Polytetrafluoroethylene	Tribological	Blending	The friction coefficient was reduced with the use of polytetrafluoroethylene in PEEK	32-35
	Biological	Plasma Immersion ion deposition	The tested technique helps improvement of orthopaedic implants in the next decade	36-38
PEEK, PEKK	Chemical	Plasma treatment	The PEEK atmospheric pressure plasma treatment performed remarkably towards improving the stickiness of the polymer	39-42

According to the thermal cooling cycle of the injection moulding process, PEEK crystallinity can vary between almost zero and 40%, also depending on the injection mould temperature and thickness of the moulded part. The crystallisation kinetics are slow in crystalline thermoplastic polyethylene terephthalate. A crystallisation interval between T_m and T_g , well above room temperature, is associated with its macromolecular skeleton's high rigidity. Therefore, in the regular cooling cycles of the injection moulding process, the polymer remains for a short time within its crystallisation interval. Table 4 gives the biomedical applications of hydroxyapatite nanocomposites [87-89].

Table 4. Comparison of high-performance PEEK composite-based biomedical devices

PEEK composite	Processing method	Mechanical Properties	Service Area	Description
Nano-TiO₂/PEEK [78-80]	Mixing of powder and compression at 400°C	3.8 GPa bending module and 93 MPa bending power	Bone substitution	The bioactivity of the end product with a nano-TiO ₂ filling improved significantly.
Nano-HAp/PEEK [80-82]	380°C compression and powder mixing	4.6 MPa elastic module and 160 MPa compression power	Orthopaedic implants	The composites nano-HAp/PEEK have been discovered to facilitate cell attachment, propagation, proliferation and osteogenic.
BioHPP [82-84]	Wax moulding at 400°C	–	Dental prosthesis	The replacement for traditional metallic materials should not be considered BioHPP
HAP embedded PEEK [84-86]	Moulding and chemical liquidation of compression	–	Interbody spinal fusion	For the backbone applications, the resulting material is promising
PEEK blanks [86-88]	Acid etching and silica coating are the techniques used for milling.	14.5 ± 2.6 MPa	Dental prosthesis	Silica PEEK coated has enhanced mechanical properties
PEEK [83-85]	Mould and soak with 400°C phosphates buffered saline	The average maximum load is 1790 N	Orthopaedic	Soak does not affect PEEK
Knitted carbon/PEEK [86-88]	Micro-braiding yarn and hot moulding at 380°C	–	Fixations Orthopedic	There was more excellent deformability in knitted composition bone plates.
Chitosan/PEEK [87-89]	Air plasma modification	–	Regenerative medicine	The antibacterial properties of Chitosan led to polymer
PEEK PSIs [90-92]	–	–	Facial Maxillo-Operation	There were no risks to the materials.
PEEK films [93-95]	Moulding compression and treatment of NaOH at 180°C	–	Ophthalmology	The creation of the PEEK apatite layer has been extended
PEEK [95-97]	CNC machine	–	Treatment of Skull Bone	The findings supported the use of PEEK

Therefore, with low mould temperatures, a rapid cooling cycle can lead to the manufacture of entirely amorphous and transparent moulded parts of small thicknesses. As the thickness of the cavity increases, pieces are formed with a translucent skin and an opaque core of crystalline material under the same rapid cooling conditions. In the case of slow cooling, with increasing mould temperatures, with T_g up to 200°C, PEEK moulds exhibit good uniformity in the degree of crystallinity throughout the thickness, with minimal variation crystallinity in the skin regions and crumb along the piece's length. The thermal process of coating CHAp onto PEEK in biological macromolecular cells is illustrated in Fig. 6 for a 3D printing process of PEEK and CHAP, with a constant melt processing temperatures of 400°C, a mould temperature ranging between 20 and 200°C, and injection speeds of 5.2, 13.1 and 23.2 cm³/s as processing variables [98-100].

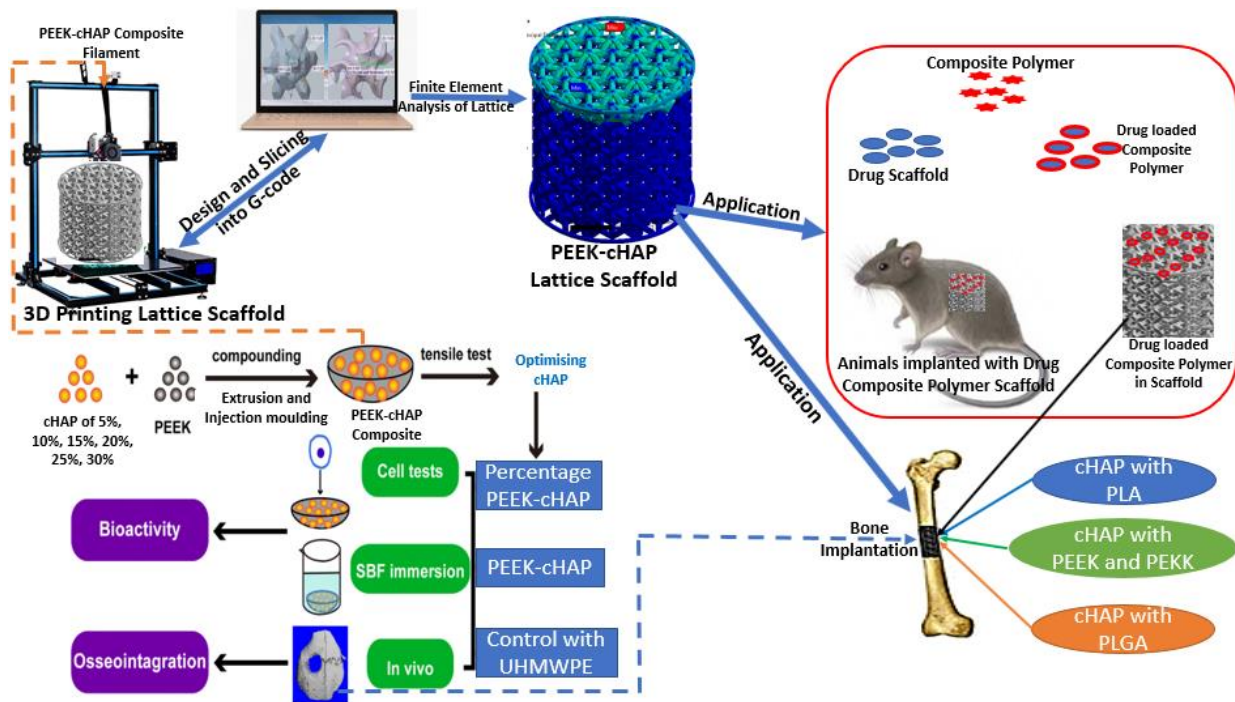


Fig. 6. Thermal process of CHAp on a coating of PEEK and its biological cellular evaluation and application [56-58]

4. Concluding remarks

In conclusion, a combination of PEEK and CHAp can be constructed either through composite material coating or extrusion, or through multilayer structures. Another alternative to improve the properties of structures used as supports to promote bone regeneration is surface treatment. The thermal shock impact from the thermal crystallinity process of CHAp on PEEK and the thermomechanical properties are discussed. The manufacturing parameters are compared for injection moulding and coating from the literature. This work reviews the mechanical characteristics of static and dynamic tests and their impacts on PEEK filament's mechanical properties after the CHAp coating process. Also, the characteristics of the coating and particle dispersion quality, morphological analysis of the granulometry, depth of penetration, CHAp layer state after static and dynamic mechanical stresses and their general applications are discussed. The X-ray diffraction characteristics of CHAp determine whether the thermal process effectively forms a CHAp layer with properties suitable for osteointegration, as revealed by this critical review with parameters preserved in technical and international standards.

In the future, interfacial biological macromolecular cellular studies of CHAp embedded in PEEK and other polymers would benefit biomedical application research, as PEEK and CHAp can be used to repair and replace bone in hard tissues for better support strength. Cellular CHAp with a soft polymer such as PLA fabric maintains the functional properties of skin tissues which need flexibility. In tendon repair and cartilage substitution, long-term blood vessels or catheters are required. To meet the current scientific challenges, interactions at nano-biomaterial interfaces must be explored. To develop safe new techniques for detecting biomolecular interactions, an accurate understanding of cellular behaviour during exposure to implanted PEEK and CHAp scaffolds is essential. In particular, tissue engineering techniques must consider cell adhesion properties, either for surface enhancement through absorption or the implantation of specific binding factors. The study of biological cellular interfaces of hybrid autonomous cell and bone material has great potential to support biomaterial development.

Ethical statement

The authors declare no ethical issue; the study was conducted in full agreement with ethical standards. Also, the manuscript is neither under review nor published elsewhere.

Declaration of competing interest

We, at this moment, affirm that there is no conflict of interest.

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