Review

Silicon nitride, a close to ideal ceramic material for medical application

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Abstract: This topical review describes salient results of recent research on silicon nitride, a ceramic material with unique properties. The outcome of this ongoing research strongly encourages the use of monolithic silicon nitride and coatings as contemporary and future biomaterial for a variety of medical applications. Crystallographic structure, synthesis and processing of monolithic structures and coatings, and examples of their medical applications are covered that relate to spinal, orthopedic and dental implants, bone grafts and scaffolds, platforms for intelligent synthetic neural circuits, antibacterial and antiviral particles and coatings, optical biosensors, and nano-photonic waveguides for sophisticated medical diagnostic devices. The examples provided show convincingly that silicon nitride is destined to become a leader to replace titanium and other entrenched biomaterials in many fields of medicine.

Keywords: Silicon nitride; structure; properties; processing; coatings; spinal implants; arthroplastic implants; bone scaffolds; dental implants; neural circuits; biosensors; medical diagnostics.
1. Introduction

Research and development of metallic, ceramic, polymeric and composite biomaterials have progressed to a level of involvement and sophistication similar to modern electronic and photonic materials. This is related to the fact that an ever increasing part of the world population is in need of repair or replacement of dysfunctional or damaged parts of the body such as dental roots and teeth, alveolar ridge augmentation, intraocular lenses, heart pacemakers, cochlear implants, and hip and knee endoprostheses [1]. As these biomaterials are supposed to be functioning in close contact with biological tissues, they have to be biologically safe, mechanically resilient, economically viable, and physiologically and esthetically acceptable. Biomaterials are poised for continuing to provide a substantial economic impact on modern medicine as parts of prosthetic devices in orthopedic, dental, cardiovascular, ophthalmological, and reconstructive surgeries, including their use as scaffolds for tissue engineering, surgical sutures, and drug and gene delivery vehicles [1].

Bioceramics come in two flavors: bioinert and bioactive. Typical bioinert ceramics in clinical use today are aluminum oxide, partially-stabilized zirconium oxide and pyrolytic carbon, whereas bioactive ceramics include a range of calcium phosphates and surface-active bioglasses. Bioinert ceramics are copiously used in applications that require mechanically strong and resilient wear- and corrosion-resistant materials with excellent tribological properties, for example as femur heads and acetabular cups of hip endoprostheses but also as intervertebral parts, in knee arthroplasty, and as dental root implants. These properties have made aluminum oxide, zirconium oxide, titanium dioxide, silicon nitride, pyrolytic graphite and diamond-like carbon important contenders for parts of load-carrying implants [3]. However, the inherent brittleness of ceramics and their finite capability to integrate with tissues restrict their clinical application at present.

Bioactive ceramics including hydroxylapatite, tri- and tetracalcium phosphates and bioglasses are utilized predominantly on account of their osseointegrative and, in combination with tissue growth-supporting non-collagenous proteins and glycosaminoglycans, osseoinductive properties that involves the recruitment of immature stem cells and the stimulation of these cells to develop into pre-osteoblasts. While hydroxylapatite is mainly used as osseconductive, i.e. bone-growth supporting coating [2] for hip and dental root implants that promote attachment of osteoblasts to the surface and throughout the interior of scaffolds or implants, bioresorbable tricalcium phosphate finds application as filler materials for bone cavities and bone replacement material, whereas tetracalcium phosphate is used in dental and bone cements. Today, bioactive ceramics are applied increasingly together with polymers as hybrid nanocomposites designed to mimic salient properties of natural biomaterials.

Recently, increasing interest has been devoted to silicon nitride, an advanced ceramic material that excels by high compressive and flexural strengths, comparatively high fracture toughness, very low friction coefficient, strong corrosion resistance, improved medical imaging ability based on radioluency in the visible and near-infrared region, enhanced biocompatibility and osseoconductivity, antimicrobial activity, and microengineering capability conducive to integrate electronics and mechanics together with computational, chemical and optical elements. Owing to these properties, silicon nitride is now emerging as novel and promising bioceramic material for a multitude of medical applications. These applications include wear-resistant bearings for hip- and knee endoprosthetic implants, novel dental implants, spinal intervertebral spacers, tissue engineering scaffolds, antibacterial and antiviral coatings, wave guides for medical diagnostics, microtubes for intelligent neural circuits, micro-spectroscopic imaging devices, photonic ICs, optical biosensors, and others. As an additional bonus, most recently silicon nitride has been found to be a powerful solid-state bioceramic deactivator of single-stranded RNA (ssRNA) viruses including the SARS-CoV-2 virus [4]. Developments are underway to commercialize antimicrobial ‘catch-and-kill’ face masks comprising silicon nitride-coated polypropylene fibers to protect against the COVID-19 pandemic [5]. In the light of all these beneficial properties, silicon nitride may be correctly considered a close to ideal bioceramics.

2. Crystalloigraphic structure of silicon nitride
Silicon nitride occurs in three polymorphic structures, two hexagonal $\alpha$- and $\beta$-$\text{Si}_3\text{N}_4$ modifications and a cubic ($\gamma$-$\text{Si}_3\text{N}_4$) high pressure/high temperature modification with spinel structure. The $\alpha$-$\text{Si}_3\text{N}_4$ polymorph crystallizes in the hexagonal space group $P31c$ (159) with lattice dimensions $a = 774.8(1)$ pm and $c = 561.7(1)$ pm, whereas $\beta$-$\text{Si}_3\text{N}_4$ crystallizes in the hexagonal space group $P6_3/m$ (176) with $a = 760.8$ pm and $c = 291.1$ pm.

As shown in Figs. 1 and 2, the structures of hexagonal silicon nitride consist of somewhat distorted SiN$_4$ tetrahedra that, by sharing corners, form distorted hexagonal rings arranged in layers with ABCD… stacking sequence ($\alpha$-$\text{Si}_3\text{N}_4$) and ABAB… stacking sequence ($\beta$-$\text{Si}_3\text{N}_4$).

As shown in Fig. 1, the basic AB layer is identical in both $\alpha$- and $\beta$- phases. However, in the $\alpha$-phase the second (CD) layer is related to the first (AB) layer by a c-glide plane. Since the double layer in $\alpha$-$\text{Si}_3\text{N}_4$ can be considered a superposition of a $\beta$-$\text{Si}_3\text{N}_4$ layer and its opposite inverted by 180°, there are twice as many atoms per unit cell in $\alpha$-$\text{Si}_3\text{N}_4$ ($Z = 4$) compared to $\beta$-$\text{Si}_3\text{N}_4$ ($Z = 2$) (Fig. 2). The interconnected SiN$_4$ tetrahedra in $\beta$-$\text{Si}_3\text{N}_4$ form wide channels within the AC plane. The c-glide plane causes the structure of $\alpha$-$\text{Si}_3\text{N}_4$ to yield two (isolated) narrow interstitial sites per unit cell instead of the wide channels present in $\beta$-$\text{Si}_3\text{N}_4$. These wide channels allow for enhanced ion diffusion throughout the lattice that assist in easy sintering at elevated temperatures. Consequently, fabrication of dense biomedical parts should advantageously commence with $\beta$-$\text{Si}_3\text{N}_4$ precursor powders [6].

3. Synthesis and processing

3.1 Synthesis and properties of monolithic silicon nitride ceramics

The mechanical, elastic, and thermal properties of structural variants of silicon nitride differ remarkably as shown in Table 1. Different synthesis routes of silicon nitride structures are shown in Fig. 3.
Processing routes to manufacture monolithic silicon nitride compacts [6, 8]. © With permission by Wiley-VCH.

Table 1. Selected mechanical and thermophysical properties of RBSN and SSN [7, 8]

| Property                        | RBSN    | SSN     |
|---------------------------------|---------|---------|
| Density (Mg·m⁻³) (% of theoretical density) | 70-88   | 95-100  |
| Compressive strength (MPa)      | 600     | 2950    |
| Flexural strength (4-point, 25°C) (MPa) | 150-350 | 500-1,000 |
| Fracture toughness (25°C) (MPa·√m) | 1.5-3   | 5-8     |
| Fracture energy (J·m⁻²)         | 4-10    | ~60     |
| Modulus of elasticity (25°C) (GPa) | 120-220 | 300-330 |
| Thermal conductivity (25°C) (W/m·K) | 4-30    | 15-50   |
| Thermal shock resistance R (K)   | 220-580 | 300-780 |
| Thermal shock fracture toughness R' (W·m⁻¹) | 500-10,000 | 7,000-32,000 |
| Coefficient of thermal expansion (10⁶ K⁻¹) | 2.5     | 3.2     |
| Specific heat capacity (J/kg·K)  | 720     | 800     |
| Poisson ratio                   | 0.24    | 0.27    |

In engineering applications, the most frequently used structural variants of silicon nitride are porous reaction-bonded silicon nitride (RBSN) and dense sintered silicon nitride (SSN). Processing of RBSN starts from fine compacted silicon powders that during prolonged heating in nitrogen or ammonia atmospheres between 1000 and 1450°C results in a mixture of α- and β-Si₃N₄ polymorphs. No shrinkage during nitriding occurs because the volume increase of about 22% will be fully accommodated by the void spaces between the silicon powder particles. Hence, the outside dimensions of the green ceramics will be connoturally retained. This is the reason why RBSN is predominantly used to manufacture complex shaped components without the need for expensive post-sintering machining with diamond tools, making RBSN parts economically highly competitive. Although the porosity of RBSN is between 20 and 30%, its mechanical performance is remarkable as flexural strengths values up to 350 MPa can readily be attained (Table 1) [6].

To fabricate fully dense sintered silicon nitride (SSN) monolithic devices, hot (HP) or hot isostatic pressing (HIP) is performed starting from powders to which oxides of magnesium, aluminum or yttrium as binder materials are added. At temperatures beyond 1550°C, these oxides form, together with a contaminant silica film surrounding individual silicon nitride particles, a thin molten quinary Si-Al-Y-O-N glass film able to readily dissolve silicon nitride, resulting in efficient densification of the ceramic body. However, there is a catch. Both HP and HIP sintering techniques are rather expensive and also limited to simple
geometric shapes that, however, are rarely sufficient for biomedical devices and applications. Critical parameters of SSN processing are (i) powder particle sizes in the 100 to 200 nm range, (ii) the capability of binder oxides to form a liquid phase around silicon nitride grains, (iii) a high-pressure nitrogen atmosphere to prevent thermal dissociation, and (iv) sinter temperatures up to 1850°C. During processing in the presence of aluminum oxide and yttrium oxide reactants, Al and O ions are able to enter the Si₃N₄ structure to form various SiAlON alloy components. Cooling of the intergranular liquid grain boundary film results in glassy and/or crystalline (YAG) phases, depending on the cooling rate [6].

To manufacture strong and mechanically resilient silicon nitride monolithic shapes, an advanced texturing technique has been applied [9]. During high temperature densification of powder compacts, the low temperature α-Si₃N₄ phase transforms to β-Si₃N₄, and by so-called texturing a rod-like morphology is generated with a specific microstructure that drastically reinforces the mechanical and thermal performance of the final monolithic end product. To speed up the conversion rate, seeds of β-Si₃N₄ are being added to the α-Si₃N₄ precursor powder. The rod-like β-Si₃N₄ crystallites will be forced to align by hot pressing, hot forging or sinter forging, or by template grain growth (TGG) followed by cold pressing, slurry extrusion, tape casting, or static or rotating magnetic fields [6, 8].

3.2 Synthesis and properties of silicon nitride thin films and coatings

In many industrial applications, it is required to strengthen, protect and/or functionalize the surface of low-grade materials by adding a suitable high-grade coating. The decisive advantage of modern coating technology is that it marries two dissimilar materials to improve, in a synergistic way, the mechanical, chemical, tribological, corrosive, optical or biological performance of the ‘tandem’ structure substrate/coating. Frequently, the mechanical strength and fracture toughness is provided by a low-cost bulk metallic substrate while a high-cost thin coating applied to its surface provides sufficient protection against environmental degradation by corrosion, wear, erosion, and by thermal and biological attack. There exists a plethora of coating techniques [2], ranging from non-thermal coatings deposited by sol-gel deposition, dip and spin coating, chemical vapor deposition (CVD), electrochemical deposition (ECD), electrophoretic deposition (EPD), hydro- and hydrothermal processing, electron- and ion beam-assisted deposition (EBAD, IBAD), radio-frequency magnetron sputtering, and plasma electrolytic oxidation (PEO) to thermal coatings deposited by atmospheric plasma spraying (APS), low-pressure plasma spraying (LPPS), suspension plasma spraying (SPS), high-velocity flame spraying (HVFS), solution precursor plasma spraying (SPPS), cold gas dynamic spraying (CGDS), pulsed laser deposition (PLD), high velocity oxyfuel spraying (HVOF) and several others, less frequently applied techniques.

Formation of pure silicon nitride coatings using conventional thermal deposition techniques including plasma spraying has been deemed impossible since Si₃N₄ dissociates beyond 1800 °C in the hot plasma plume or combustion flame. Silicon nitride also oxidizes in the presence of oxygen at high temperature. In contrast, thin amorphous silicon nitride films are routinely deposited by chemical vapor deposition (CVD) and applied as masking layers for profile etching during production of semiconductor ICs, diffusion barriers in very-large-scale integrated (VLSI) production lines to prevent cross penetration of dopants, to protect optical fibers against mechanical damage, as gate dielectrics with low leakage and high resistance for metal-insulator semiconductor (MIS) memory devices, for antireflection coatings of solar cells, and as moisture barrier for organic light emitting diode (OLED) displays [10].

Sadly, any attempt to deposit mechanically stable, thick and well-adhering silicon nitride coatings for biomedical application by thermal spraying using metallic [11, 12] or silicate glass binders [13], conversion in the as-deposited state through a reactive spray process [14], or in situ nitridation in flight [15] was not met with resounding success; such coatings contained only rather small amounts of silicon nitride but instead, substantial amounts of embrittling metal silicides, binder ceramic materials, and unreacted precursor materials. Slightly more successful were endeavors to prepare high-Si₃N₄ coatings by starting from β’-Si₃N₄-ALO₃-NS₄ compositions of the SiAlON family with different degrees of substitution, z [16] or clad-type powder consolidation using alloy bond coats [17]. Also, high particle velocities generated by techniques as
diverse as detonation spraying (DS), Top Gun technology, high-frequency pulse detonation (HFPD), and atmospheric plasma spraying (APS) with axial powder injection were conducive to deposit dense and reasonably well-adhering silicon nitride-containing coatings [18-22]. In addition, optimization of heat transfer into the powder particles was found to be one of the most decisive factors to deposit well-adhering coatings [23]. This, however, necessitates special powder preparation procedures [24, 25].

All attempts to deposit silicon nitride coatings on metallic substrates are marred so far by the fact that addition of binder metal oxides such as Al₂O₃ + Y₂O₃ is required. Hence, no phase-pure homogeneous silicon nitride coatings could be produced to date. This problem was, at least partially, solved by spraying particles with high velocities up to 3 km/s that were generated by an electromagnetically accelerated plasma (EMAP). Under these extreme conditions, dense, homogeneous silicon nitride coatings with reduced amounts of binder oxides could be deposited with suitable mechanical properties, including tight adhesion to the surface of polished stainless steel [26, 27].

Figure 4 shows such dense and well-adhering, mechanically stable silicon nitride-based coatings on austenitic stainless steel substrates deposited by conventionally detonation spraying (left) and ultrahigh-velocity pulsed plasma spraying, EMAPS (right). However, at the time of this writing, the novel technology has not progressed to a state where the pulsed mode could be replaced by a continuous delivery mode. Consequently, the deposition process is still economically untenable while tedious since after each ‘shot’ the detonation chamber must be refilled with powder to be ready for the next shot. Nevertheless, this advanced deposition method may be at the cutting edge of future medical coating technologies designed for silicon nitride coatings as protective bearing surface for novel arthroplastic devices or ceramic hip resurfacing (HR) prostheses (see below).

In a similar development, a magneto-plasma-dynamic (MPD) arc jet generator was utilized to deposit silicon nitride coatings by reactively spraying material from a crystalline silicon rod in a nitrogen atmosphere [28]. After 200 shots at a frequency of 0.03 Hz with a discharge current of 9 kA, 30 µm thick, dense and homogeneous β-Si₃N₄ coatings could be deposited.

4. Biomedical applications

Although non-oxide ceramics such as silicon nitride are predominantly being used in high-temperature applications including parts of advanced gas turbines [29], turbocharger rotors for high-performance automotive engines [30] and rocket thrusters [31], more recent research results have strongly encouraged the use of silicon nitride in a variety of biomedical applications [32, 33]. Silicon nitride excels by a unique combination of bulk mechanical and surface chemical properties thus, making it a close to ideal biomaterial for orthopedic implants. It is already being used clinically for interbody spinal fusion cages and is being developed for total joint arthroplasty. Its surface texture and chemistry are both highly tunable, yielding physicochemical combinations of properties that may lead to enhanced osseointegration and bacterial and viral resistance.
On top of superior mechanical, thermal and chemical properties, silicon nitride ceramics have demonstrated high cell adhesion and enhanced osteoconductivity similar to non-stoichiometric (Ca-deficient) biological hydroxylapatite. This advantageous feature is thought to be based on the zwitterionic-like behavior of silicon nitride that is a function of the tradeoff between positively charged vacancies in the partial N lattice of suitable doped silicon nitride and negatively charged silanol groups at its surface [32].

a. Intervertebral spacers

Spinal implants are used to alleviate many forms of back pain and deformity including trauma-based injuries and disorders such as scoliosis, kyphosis, degenerative disc disease, and fracture. Their primary function is to help to fuse two vertebrae together and to replace natural disc material. The four types of surgery that utilize spinal implants are anterior, posterior, and transforaminal lumbar interbody (TLIF) and axial fusions.

Silicon nitride has been in clinical use for porous intervertebral spacers in spinal fusion surgery for over thirty years without showing any undesirable long-term effects [34-38]. Silicon nitride possesses high strength, long-term resilient, mechanically reliable, and osseointegrative properties, partly based on a high proportion of covalent bonds. To achieve interbody fusion, the ability of silicon nitride to yield superior new bone ingrowth and osseointegration, along with proven bacteriostatic properties and enhanced medical imaging, outperforms both conventional poly(ether ether ketone, PEEK) and titanium implants [39, 40]. This is largely based on its pronounced hydrophilicity that attracts extracellular fluid rich in bone growth-mediating non-collagenous proteins, turning on osteoblasts and suppressing osteoclasts. Consequently, silicon nitride speeds up bone healing, bone fusion and thus, implant integration. The efficacy of silicon nitride for spinal reconstruction is supported by its strong bacterial resistance caused by the synergistic effects of surface chemistry, surface pH, texture, and electric charge [37].

Anterior cervical discectomy with fusion is a commonly applied surgical procedure to treat radicular arm pain. A study compared silicon nitride implants with PEEK cages filled with autograft harvested from osteophytes [41]. Patients treated with either silicon nitride or PEEK implants reported comparable recovery rates during follow-up interviews. No significant differences in clinical outcome were noticed at 24 months observation time. Fusion rates improved over time and were comparable between both groups. Figure 5 shows the Valeo C®C® silicon nitride cervical interbody fusion device used in the CASCADE trial with radiographic characteristics. The outer part of the device consists of dense silicon nitride whereas the inset comprises cancellous structured ceramic (CSC) of the same composition, providing a scaffold for easy bone ingrowth [42].

Figure 5. Silicon nitride cervical interbody fusion device (Valeo C®C®, left) with radiographic image of the implanted device (right) [41]. © Creative Commons Attribution 4.0 International License.

4.2 Knee- and hip endoprosthetic implants
Silicon nitride compacts can be polished to a high surface finish to provide an exceptionally smooth, wear resistant, and tribological superior surface. This renders the material ideal for articulating applications including bearings for knee and hip endoprostheses [43-45] as well as acetabular cups. Salient materials properties conducive to supporting bearing applications are phase stability [46], wear resistance [46, 47], strength and fracture toughness [48], hydrophilicity [49], favorable medical imaging capability [37], and unsurpassed bacterial resistance [39, 50]. Figure 6 shows a selection of silicon nitride compacts produced by SINTX Technologies of Salt Lake City, Utah, USA, the only vertically integrated silicon nitride medical device manufacturer worldwide [51].

Silicon nitride ceramics were found to be highly biocompatible and non-cytotoxic as revealed by culture tests with mouse fibroblasts (L-929 cell line) and human mesenchymal stem cells (hMSCs) [52]. Excellent cytocompatibility was confirmed by the outcome of live/dead cell staining and pronounced differentiation towards osteoblasts. Biocompatibility was also confirmed by the tissue response around cylindrical silicon nitride implants installed in the tibiae of rabbits [53]. Histological analysis showed that bone growth occurred preferentially in the cortical area, and different types of tissue were identified on the implant surfaces such as lamellar bone tissue containing osteocytes and osteons, non-calcified matrix containing osteoblasts as well as collagen type III that might in time be converted to collagen type I.

Tribological hip simulator tests showed a very low friction coefficient that decreased to still lower values when the articulating time was increased. This is likely caused by the hydrodynamic action of a self-lubricating hydrogen-terminated silicon oxide boundary film that develops during prolonged experimental articulation in an aqueous environment [54].

![Figure 6](https://example.com/figure6.jpg)

**Figure 6.** Various silicon nitride compacts produced by SINTX Technologies, including ball bearings (left), an acetabular cup (center), an intervertebral spacer with a CSC core (foreground, center), and spinal interbody fusion devices (right) [51]. © Creative Commons Attribution 4.0 International License.

Silicon nitride with a high degree of covalent bonding was found to possess significantly higher mechanical strength compared to oxide ceramics such as Al₂O₃ or ZrO₂ with predominantly ionic bonds. Finite element analysis (FEA) revealed a stress distribution within a femur, implanted with a silicon nitride prosthesis, comparable to that of an intact and healthy one [55]. Moreover, lifetime prediction studies showed conclusively that implants made from silicon nitride are mechanically reliable and consequently, may be suitable for use in arthroplastic endoprostheses. These results strongly suggest that non-oxide ceramics such as silicon nitride with high mechanical strength may be useful future biomaterial candidates for highly loaded ceramic hip resurfacing (HR) endoprostheses based on their low wear rate, low friction coefficient, and pronounced biocompatibility and antibacterial properties [56].

However, there is a drawback. The high modulus of elasticity of HPSN and SSN in excess of 300 GPa prohibits the use of these ceramic materials for load-bearing stems of hip implants, based on the effect of strong stress shielding that is caused by the large gradient of the modulus of elasticity between implant material and cortical bone. Although reaction-bonded silicon nitride (RBSN) with a noticeably lower elastic modulus around 150 GPa has been suggested as a viable alternative option for this particular application,
the modulus is still an order of magnitude higher than that of cortical bone (around 20 GPa). Moreover, as a consequence of the high porosity of RBSN both flexural strength (150-350 MPa) and fracture toughness (1.5-3 MPa√m) are substantially below those of HPSN and SSN (see Table 1) [6, 8].

In contrast, acetabular cups based on silicon nitride may find their way into the orthopedic operation theater [57]. In the majority of hip endoprosthetic implants, the femoral head articulates against an acetabular cup liner composed of tough polymers such as ultra-high molecular weight polyethylene (UHMWPE) or X-linked polyethylene. These polymers degrade over time by chain scission and thus, strategies must be applied to slow done this deterioration, including radiative cross-linking [58] or doping with antioxidants such as vitamin E [59]. Owing to the unique surface chemistry of silicon nitride, this material is able to absorb oxygen released from degrading polyethylene [60], thus limiting acetabular liner degradation in hip endoprostheses. This also helps to avoid polymer wear debris that is known to cause the so-called ‘particle disease’ that may lead to aseptic loosening of the implant within the femur by periprosthetic osteolysis [61] and, when distributed by the lymphatic system throughout the body, may cause hepatic degeneration.

Frequently, CoCrMo alloys such as ASTM F75-12 or ASTM F1537 are utilized as hip endoprosthetic stem materials [1], with high implant survival rates. However, certain risks including accelerated wear rates and release of cytotoxic Co and Cr ions limit their usefulness. In a recent study [62], 4 µm-thick silicon nitride (SiNₓ) (N/Si = 1.10 to 1.25) coatings were deposited by reactive d.c. magnetron sputtering onto CoCrMo surfaces, with a 0.5 to 0.8 µm CrN bond coat applied to increase adhesion. Wear tests performed against SiNₓ in a ball-on-disc tribometer revealed specific wear rates essentially below those of CoCrMo. Similar results were obtained by Maria Petterson as reported in her Ph.D. thesis [63]. To address the question of cytotoxicity, the role of particle size, surface composition and donor variations of silicon nitride on its biological impact were investigated [64]. Cytotoxicity, inflammatory cytokine release, oxidative stress, and genotoxicity potential of silicon nitride and CoCr particles were tested and compared using human peripheral blood mononuclear cells (hPBMNCs). Minimum biological impact of silicon nitride on human PBMCNs was recorded as opposed to CoCr particles that showed significantly higher levels of TNF-α cytokine release, oxidative stress, and DNA damage, confirming that silicon nitride, when properly engineered, is an attractive orthopedic biomaterial for joint replacement devices.

4.3 Bone grafts and scaffolds

Bone grafting is a surgical procedure that uses transplanted bone to repair and rebuild diseased or damaged bones. In the clinical practice, bone grafts present limitations that include harvesting morbidity, insufficient bioactivity, and concern about transmission of diseases [65]. Silicon nitride bone scaffolds and bone fusion devices [36] excel by high and reliable mechanical strength, biocompatibility, and antibiotic capability, resulting in a bone healing sequence comparable to hydroxylapatite [32]. MC3T3-E1 cells were used to study the osteoblastic differentiation and mineralization on sterile samples of silicon nitride, and compared them to samples of titanium and PEEK, standard materials for bone scaffolds [66]. The study reported more profound and faster ECM deposition and mineralization on SiNₓ surface as compared to titanium and PEEK. Results further indicated the upregulation of osteogenic transcription factors such as RUNX2 and osterix (SP7), as well as collagen type I and osteocalcin. Hence, silicon nitride rapidly conducts mineralized tissue formation via extracellular matrix deposition and biomarker expression in murine calvarial pre-osteoblast cells.

4.4 Dental implants

Silicon nitride is an emerging material for dental restoration and dental implant application. Besides excellent structural, mechanical, thermal, tribological and biocompatible properties, silicon nitride is a potent antibacterial material. Dental implants made from silicon nitride have been shown to exhibit mechanical properties including modulus of elasticity approaching those of bone tissue that promise to reduce stress shielding when compared to classic titanium alloy implants [67]. Hence, in view of available pre-
clinical data, silicon nitride appears to have the essential characteristics to be a strong candidate for dental implants material. This novel ceramic has a surface with potentially antimicrobial properties, and if this is confirmed in future research, it could be of great interest for oral use [68, 69].

A recent study [70] investigated the potential use of porous silicon nitride for all-ceramic dental restorations as a core material and results were compared with those of a commercial ZrO2 ceramic. The color of silicon nitride could be tailored by the porosity introduced and a color shade suitable for restorative applications was obtained. The flexural strength of Si3N4 was found to be 418 MPa despite an open porosity of nearly 10.5% (cp. Table 1). This rather high strength may be related to strong neck formation between β-Si3N4 grains, the intertwined distribution of these grains, and the crack deflection potential of rod-like β-Si3N4. The Vickers hardness of Si3N4 was found to be 10.9 MPa whereas ZrO2 had 13.7 MPa, which reduces the risk for wear of opposing natural teeth. Shear bond strength to dentin was tested and indicated low values for silicon nitride without a coupling agent (2.24±1.15 MPa) that increased strongly by adding a silane coupling agent to 8.44±2.98 MPa, suggesting that the usage of coupling agents for Si3N4 is essential.

The radiolucent behavior of Si3N4 will enable for both the restorations and the surrounding tissues to be imaged using plain radiography (Fig. 7). Hence, this study shows that with tailored manufacturing methods, silicon nitride can be considered an effective dental restorative material.

White silicon nitride can be prepared by pressureless sintering at 1650°C in a nitrogen atmosphere [69]. In a recent study [72], partially sintered Si3N4 ceramic blocks were prepared by heating to 1400°C in a N2 gas environment. After full sintering at 1650°C for 2 h, the linear shrinkage value was recorded at 19.9±0.6%. The flexural strength and fracture toughness were measured, the results were 891.2±37.2 MPa and 6.3±0.30 MPa·√m, respectively. In conclusion, white Si3N4 can be favorably used to fabricate multi-unit dental restorations conforming to ISO 6872 designation [73].

Although zirconia is still the most commonly used ceramic for dental implants, its intrinsic bioinert nature often results in a lack of biological integration. A silicon nitride-based laser-cladding process has been developed to improve the biological response to biomedical zirconia by forming a composite coating with Si3N4 particles dispersed in a nano-crystalline/amorphous silicon matrix. Raman spectroscopy revealed that the bone tissue quality parameters were comparable to healthy human bone tissue, suggesting that laser-cladding with silicon nitride might be able to improve the stability of zirconia dental implants [74].
In conclusion, the metal-free nature of silicon nitride makes this material a promising candidate for dental implantology as release of cytotoxic metallic by-products able to invade any part of the body can be avoided. Higher esthetics, better osseointegration and biocompatibility, antibacterial property along with high wear resistance and metal-free corrosion products render silicon nitride a powerful contender for dental implant application.

4.5 Intelligent synthetic neural circuits

Synthetic regulatory circuits encoded in RNA rather than DNA could provide a means to control cell behavior while avoiding potentially harmful genomic integration in therapeutic applications. Post-transcriptional circuits have been developed using RNA-binding proteins that can be wired in a plug-and-play fashion to create networks of higher complexity [75]. Application of this research in the realm of future information technology include signal transmission platforms for artificial intelligence (AI) algorithms with unparalleled computing speed based on novel synthetic neural circuits.

A neuron cell culturing platform consisting of arrays of ordered silicon nitride microtubes (2.7–4.4 μm in diameter) was formed by strain-induced self-rolled-up nano-membrane (s-RUM) technology using ultrathin (<40 nm) silicon nitride films deposited on transparent substrates such as glass [76]. These microtubes presented strong guidance for fast growing primary cortical neurons, with a coaxially confined configuration resembling that of myelin sheaths. Neurons revealed a dramatically enhanced growth rate inside the microtube compared to regions outside the microtubes. The acceleration and precise guidance provided by the silicon nitride microtube array was attributed to 3D-adhesion and electrostatic interaction with the silicon nitride microtubes, respectively. The microtubes not only provide structure for the neural network and guiding connections, but also accelerate the growth of axons, an important feature as time is crucial for restoring severed connections in the case of spinal cord injury or limb reattachment. A time-lapse sequence of a cortical neuron rapidly growing from inside a microtube through neighboring microtubes can be watched on the Internet [77].

This research has important implications for designing and building of intelligent synthetic neural circuits for potential treatment of neurological disorders. In addition, the novel self-rolled-up silicon nitride microtubes could provide an ideal route to integrate electrodes into neural cultures, coupled with photonics, to directly monitor dynamic neural activities and record action potentials with enhanced resolution and fidelity [78].

In conclusion, self-rolled-up silicon nitride microtubes are attractive for photonic, electronic, and biological applications based on their transparency under visible light, tunable refractive index, non-toxicity, and compatibility with Si-integrated circuits [79]. Thus, they may become important additions to the toolbox of 4th generation biomaterials that will be integrated with the human body to provide powerful diagnostic and therapeutic capability able to manipulate, amplify and monitor bioelectric cellular signals that are important for tissue regeneration and eventually allowing for communication with the host tissues [1, 80].

4.6 Antibacterial and antiviral particles and coatings

The efficacy of silicon nitride as a ‘killer ceramics’ for Gram-negative bacteria such as Porphyromonas gingivalis was found to be the result of chemically driven mechanisms, related to the peculiar pH-dependent surface chemistry of silicon nitride [50]. In an alkaline pH environment, a buffering effect controlled by the release of ammonium (NH4+) ions causes lysis of bacterial proteins as confirmed by conventional fluorescence and in situ Raman microprobe spectroscopies. Formation of peroxinitrite within the bacteria leads to degradation of nuclei acid, and reduction in phenylalanine content and liquid concentration were observed. Experiments confirmed that modification of the surface chemistry of silicon nitride by chemical etching or thermal oxidation influenced the peroxinitrite formation and thus, affected bacteria metabolism.
Polyethylene coated by pulsed laser technique with silicon nitride powder in a nitrogen gas atmosphere showed antibacterial properties in vitro against S. epidermidis. The coating significantly reduced the amount of living bacteria when compared to the uncoated polymer. Osteoconductivity was also tested in vitro using SaOS-2 osteosarcoma cells. The presence of the silicon nitride coating resulted in an increased amount of hydroxyapatite deposited from extracellular fluid (ECF). Hence, coating of polyethylene with silicon nitride may lead to improved performance of orthopedic endoprosthesis medical devices [81].

To confront the challenge of bacterial resistance to antibiotic treatment has become one of the most critical issues of present medical research. Methicillin-resistant strains of S. aureus (MRSA) are major killers of patients in hospitals or other health care settings, such as nursing homes and dialysis centers. In these settings, the harmful bacteria are known as health care-associated MRSA (ha-MRSA). A study has found that antimicrobial behavior of machined Si$_3$N$_4$ surfaces was due to repulsion of the bacteria, a phenomenon that also limited osteogenesis, as host cells were also unable to adhere to the machined surface. This suggests that the in vivo biological behavior of Si$_3$N$_4$ orthopedic implants is driven by critical features of their surface nanotopography [82].

Silicon nitride has been shown to inactivate several bacterial species and viral strains including SARS-CoV-2, while mammalian cells would remain unaffected [83]. SARS-CoV-2 virions (2×10$^4$ PFU/mL diluted in growth media) were exposed to 5, 10, 15, and 20% (w/v) of an aqueous suspension of sintered Si$_3$N$_4$ particles for durations of 1, 5, and 10 minutes, respectively. Following each exposure to Si$_3$N$_4$, the remaining infectious virus was quantitated by plaque assay. It was found that the reduction in SARS-CoV-2 titers were markedly reduced when exposed to all concentrations of Si$_3$N$_4$, showing reductions between 85% - 99.6%, depending on the dose and duration of exposure. In conclusion, surface treatment strategies based on silicon nitride particles or coatings may offer novel methods to discourage SARS-CoV-2 persistence and infectivity on surfaces and thus, control the spread of the COVID-19 pandemic. Additional research has provided insight into the mechanism of the antiviral properties of silicon nitride [4]. Conceivably, it derives from a hydrolytic reaction of the silicon nitride surface [50] and the related formation of reactive nitrogen species (RNS) that could be easily metabolized by mammalian cells but that are lethal to pathogenic bacteria and ssRNA viruses. It was confirmed by PCR (polymerase chain reaction) tests that RNS directly react with viral proteins and RNA. Consequently, silicon nitride may play an important role in controlling the spread of human epi- and pandemics caused by ssRNA mutant viruses and other pathogens.

4.7 Medical diagnostics

Cellular sampling and characterization include culture, differentiation and fixing of cell on silicon nitride substrates for imaging and mapping across the electromagnetic spectrum, from X-ray fluorescence (XRF) to X-ray absorption near edge spectroscopy (XANES) to visible and infrared microspectroscopies [84]. Cells were found to adhere strongly to silicon nitride surfaces, allowing for investigation of visually displayed proliferative and phenotypic growth.

Lasers for medical application operate in a wide range of the electromagnetic spectrum, from X-ray up to UV, continue in the visible and near-infrared (NIR) to mid-IR. These electromagnetic waves need to be transmitted from the laser source to the target tissue by a flexible device, called a waveguide that will enable easy manipulation of the laser beam in a medical diagnostic setting [85]. Novel integrated nanophotonic sensing devices operating in the visible and near-infrared regions with drastically reduced propagation losses were developed based on silicon nitride waveguides [86]. Silicon nitride is frequently chosen as the passive waveguide material due to its several advantages compared to silicon, including higher transparency at wavelengths below 1.1 μm, ultra-low two-photon absorption effect at telecommunication wavelengths, and ultimately, smaller propagation losses. These properties have made silicon nitride a widely used platform in silicon-integrated photonic applications [87].

In addition, silicon nitride waveguide-based nano-photonic integration platforms have potential for the miniaturization of optical coherence tomography (OCT) systems [88, 89]. Novel silicon nitride-based
nano-photonic waveguides find increasingly application in Raman spectroscopy to enhance signal strength [90] and to miniaturize chips for SERS (surface-enhanced Raman spectroscopy) [91].

Chemical activation of silicon nitride surfaces with either aqueous or organic solutions of 3-aminopropyl triethoxysilane (APTES) led to the development of an optical biosensor allowing for immobilization of biomolecules such as proteins by covalent bonding [92]. Protein immobilization was followed by incubation with murine γ-globulin and reaction with fluorescently labelled goat antimouse IgG antibody. Recording the surface fluorescence revealed improved signal intensity and reproducibility compared to other protocols.

5. Conclusion

Research and development effort on silicon nitride has impressively shown that this unique ceramic biomaterial provides a host of advantageous properties conducive to application in modern medicine as part of prosthetic devices in orthopedic, dental, cardiovascular, ophthalmological, and reconstructive surgeries. These applications are based on high compressive and flexural strengths, high fracture toughness, low friction coefficient, high wear and corrosion resistance, improved medical imaging ability as a radiolucent material, and enhanced biocompatibility, osseointegration and antimicrobial activity against bacteria and viruses. These unique properties suggest that silicon nitride is destined to become a leader to replace titanium and other entrenched biomaterials in many contemporary and future medical applications. However, as advantageous the mechanical, tribological, chemical and biological properties of silicon nitride are, its comparatively high modulus of elasticity poses a challenge for application for load-bearing stems for arthroplastic implants. Even though modulus reduction can be achieved by using porous reaction-bonded silicon nitride (RBSN) instead of dense pressureless sintered silicon nitride (SSN), the modulus gradient between implant and cortical bone is still too large to suppress effectively stress shielding of hip endoprosthetic implants, in contrast to intervertebral spacers that have enjoyed lasting success for more than thirty years.

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