15. Titanium and Titanium Alloy Applications in Medicine

15.1 Metallurgical Aspects

15.1.1 Introduction

Titanium is a transition metal. It occurs in several minerals including rutile and ilmenite, which are well dispersed over the Earth’s crust. Even though titanium is as strong as some steels, its density is only half of that of steel. Titanium is broadly used in a number of fields, including aerospace, power generation, automotive, chemical and petrochemical, sporting goods, dental and medical industries, [1–3]. The large variety of applications is due to its desirable properties, mainly the relative high strength combined with low density and enhanced corrosion resistance, [4]. Among metallic materials, titanium and its alloys are considered the most suitable materials in medical applications because they satisfy the property requirements better than any other competing materials, like stainless steels, Cr-Co alloys, commercially pure (CP) Nb and CP Ta, [5–6]. In terms of biomedical applications, the properties of interest are biocompatibility, corrosion behavior, mechanical behavior, processability and availability, [7–9].

Titanium may be considered as being a relatively new engineering material. It was discovered much later than the other commonly used metals, its commercial application starting in the late 40’s, mainly as structural material. Its usage as implant material began in the 60’s, [10]. Despite the fact that titanium exhibits superior corrosion resistance and tissue acceptance when compared with stainless steels and Cr-Co-based alloys, its mechanical properties and tribological behavior restrain its use as biomaterial in some cases. This is particularly true when high mechanical strength is necessary, like in hard tissue replacement or under intensive wear use, [11]. To overcome such restrictions, CP titanium was substituted by titanium alloys, particularly, the classic grade 5, i.e. Ti-6Al-4V alloy. The Ti-6Al-4V $\alpha+\beta$ type alloy, the most
worldwide utilized titanium alloy, was initially developed for aerospace applications, [12, 13]. Although this type of alloy is considered a good material for surgically implanted parts, recent studies have found that vanadium may react with the tissue of the human body, [2]. In addition, aluminum may be related with neurological disorders and Alzheimer’s disease, [2]. To overcome the potential vanadium toxicity, two new vanadium free α+β type alloys were developed in the 1980’s. Vanadium, a β-stabilizer element, was replaced by niobium and iron, leading to Ti-6Al-7Nb and Ti-5Al-2.5Fe α+β type alloys, [4, 6, 14]. While both alloys show mechanical and metallurgical behavior comparable to those of Ti-6Al-4V, a disadvantage is that they all contain aluminum in their compositions.

In recent years, several studies have shown that the elastic behavior of α+β type alloys is not fully suitable for orthopedic applications, [15–18]. A number of studies suggest that unsatisfactory load transfer from the implant device to the neighboring bone may result in its degradation, [9]. Also, numerical analyses of hip implants using finite element method indicate that the use of biomaterials with elastic behavior similar to cortical bones improves the distribution of stress around the implanted bone, [19]. While the elastic modulus of a cortical bone is close to 18 GPa, [7], the modulus of Ti–6Al–4V alloy is 110 GPa, [7]. In such a case, the high elastic modulus of the implant material may lead to bone resorption and possible unsuccessful implantation procedure. The elastic behavior mismatch between the implant and the adjacent bone is named ‘stress shielding effect’, [19].

Since CP titanium and some specific α+β type titanium alloys do not completely meet the demands of medical applications, especially concerning mechanical behavior and toxicity to human body, a new class of alloys has been investigated for biomedical applications in the last decade, the β type alloys. After proper heat treatments this type of alloys may exhibit low elastic modulus, very good corrosion resistance, suitable mechanical properties and good biocompatible behavior, as they may be obtained by adding biocompatible alloying elements like Nb, Ta and Zr to titanium, [20–24].

15.1.2 Basic Aspects of Titanium Metallurgy

The microstructure diversity of titanium alloys is a result of an allotropic phenomenon. Titanium undergoes an allotropic transformation
at 882°C. Below this temperature, it exhibits a hexagonal close-packed (HCP) crystal structure, known as α phase, while at higher temperature it has a body-centered cubic (BCC) structure, β phase. The latter remains stable up to the melting point at 1,670°C, [5]. As titanium is a transition metal, with an incomplete d shell, it may form solid solutions with a number of elements and hence, α and β phase equilibrium temperature may be modified by allowing titanium with interstitial and substitutional elements.

Titanium alloying elements fall into three class: α-stabilizers, β-stabilizers and neutral. While elements defined as α-stabilizers lead to an increase in the allotropic transformation temperature, other elements, described as β-stabilizers provoke a decrease in such a temperature, [25]. When a eutectoid transformation takes place, this β-stabilizer is termed eutectoid β-stabilizer, otherwise, it is called isomorphous β-stabilizer. If no significant change in the allotropic transformation temperature is observed, the alloying element is defined as neutral element. Figure 15.1 shows a schematic representation of types of phase diagram between titanium and its alloys elements, [5, 25].

As a result, titanium alloys with an enormous diversity of compositions are possible. Among α-stabilizer elements are the metals of IIIA and IVA groups (Al and Ga) and the interstitials C, N and O. On the contrary, β-stabilizer elements include the transition elements (V, Ta, Nb, Mo, Mg, Cu, Cr and Fe) and the noble metals.

**Fig. 15.1.** Schematic representation of types of phase diagram between titanium and its alloying elements [5, 25]
Addition of $\alpha$ and $\beta$-stabilizer elements to titanium gives rise to a field in the corresponding phase diagram where both $\alpha$ and $\beta$ phase may coexist. Titanium alloys exhibit a variety of properties, which are connected to chemical composition and metallurgical processing, [26–28]. According to the nature of their microstructure, titanium alloys may be divided as either $\alpha$ alloys, $\beta$ alloys and $\alpha+\beta$ alloys, [29]. Beta alloys may be further classified into near $\beta$ and metastable $\beta$ alloys.

$\textit{Alpha titanium alloys}$ are especially formed by CP titanium and alloys with $\alpha$-stabilizer elements, which present only $\alpha$ phase at room temperature. Such alloys show high creep resistance and are thus suitable for high temperature service. Since no metastable phase remains after cooling from high temperature, no major modification in terms of microstructure and mechanical properties are possible using heat treatments. Finally, as $\alpha$ phase is not subjected to ductile-brittle transition, these alloys are proper for very low temperature applications. Regarding mechanical and metallurgical properties, $\alpha$ alloys present a reasonable level of mechanical strength, high elastic modulus, good fracture toughness and low forgeability, which is due to the HCP crystal structure.

$\textit{Beta titanium alloys}$ are obtained when a high amount of $\beta$-stabilizer elements are added to titanium, which decreases the temperature of the allotropic transformation ($\alpha/\beta$ transition) of titanium, [30]. If the $\beta$-stabilizer content is high enough to reduce the martensitic start temperature (Ms) to temperatures below the room temperature, nucleation and growth of $\alpha$ phase will be very restricted, and hence, metastable $\beta$ is retained at room temperature under rapid cooling, as depicted in Figure 15.2. This type of titanium alloy may be hardened by using heat treatment procedures, [31]. In some cases, depending upon composition and heat treatment parameters, precipitation of $\omega$ phase is possible. However, $\omega$ phase may cause embrittlement of a titanium alloy and, in general its precipitation must be avoided, [32]. $\beta$ titanium alloys are very brittle at cryogenic temperatures and are not meant to be applied at high temperatures, as they show low creep resistance.

Finally, $\alpha+\beta$ alloys include alloys with enough $\alpha$ and $\beta$-stabilizers to expand the $\alpha+\beta$ field to room temperature, [5, 25]. The $\alpha$ and $\beta$
Applying proper heat treatments and thermo-mechanical processing, a significant assortment of microstructures may be obtained when compared to $\alpha$ type alloys. The Ti-6Al-4V alloy is an example of $\alpha + \beta$ type alloy. Due to its large availability, very good workability and enhanced mechanical behavior at low temperatures, such an alloy is the most common composition among the titanium alloys and based on these characteristics it is still largely applied as a biomaterial, mainly in orthopedic implant devices. Figure 15.3 depicts the microstructures of $\beta$ and $\alpha + \beta$ titanium alloys.

As in the case of iron (steels), allotropic transformation is the main reason for the enormous variety of microstructure in titanium alloys. Titanium alloy microstructures are formed by stable and metastable phases, [33, 34]. In general, for limited $\beta$-stabilizer content and depending on cooling conditions, titanium alloys show only $\alpha$ and $\beta$ phases.

However, if the thermodynamic equilibrium is not reached, metastable phases may be retained at room temperature, mainly, martensitic and $\omega$ phases. According to several authors, [35–37], titanium alloys with $\beta$-stabilizer elements like Mo, Nb, Ta and V, may form two types of martensitic structures. If the $\beta$-stabilizer content is considered low, rapid cooling leads to formation of hexagonal martensite, termed $\alpha'$.
When this content is high, $\alpha'$ martensite undergoes a distortion, loses its symmetry and is substituted by orthorhombic martensite, defined as $\alpha''$. [37].

When titanium alloys with $\beta$-stabilizer elements are submitted to rapid cooling from high temperature, the $\beta$ phase may transform either in martensitic structures or eventually, in metastable $\omega$ phase, [35]. Figure 15.4 presents a microstructure of the Ti-25Nb (wt%) after cooling in water and in air showing $\alpha''$ and $\omega$ formation.

Precipitation of $\omega$ phase occurs only in a limited range of the alloy element and may arise during the quenching from high temperature ($\beta$ phase), forming a thermal $\omega$ phase. However, $\omega$ phase may also
form after ageing of a rapid by quenched structure at medium temperatures, resulting in isothermal $\omega$ phase, as indicated in Figure 15.5.

Fig. 15.4. Microstructure of the Ti-25Nb (wt%) alloy: (a) water cooled sample showing martensitic structure (OM analysis) and (b) air cooled sample showing $\omega$ phase dispersed in $\beta$ matrix and respective SADP showing $\omega$ and $\beta$ phases microstructure (TEM analysis)
Concerning mechanical behavior, biomedical titanium alloys applied as biomaterial mainly in hard tissue replacement, must exhibit a low elastic modulus combined with enhanced strength, good fatigue resistance and good workability. Mechanical behavior of titanium alloys is directly related to composition and mainly, thermo-mechanical processing. Some mechanical properties of selected titanium-based materials applied as biomaterials are shown in Table 15.1 [38].

Mechanical strength may be increased by adding alloying elements, which may lead to solid-solution strengthening or even, precipitation

Fig. 15.5. A schematic TTT diagram for $\beta$ phase transformation in titanium alloys with $\beta$-stabilizer elements [35]

15.1.3 Mechanical Behavior

Concerning mechanical behavior, biomedical titanium alloys applied as biomaterial mainly in hard tissue replacement, must exhibit a low elastic modulus combined with enhanced strength, good fatigue resistance and good workability. Mechanical behavior of titanium alloys is directly related to composition and mainly, thermo-mechanical processing. Some mechanical properties of selected titanium-based materials applied as biomaterials are shown in Table 15.1 [38].

Mechanical strength may be increased by adding alloying elements, which may lead to solid-solution strengthening or even, precipitation
Titanium alloys present a high strength-to-weight ratio, which is higher than with most steels. While CP titanium has yield strength between 170 (grade 1) and 485 MPa (grade 4), titanium alloys may present values higher than 1500 MPa [25].

The elastic modulus or Young modulus corresponds to the stiffness of a material and is associated with the way inter-atomic forces vary with distance between atoms in the crystal structure. A comparison of second phases. Also, by using ageing processes, metastable structures obtained by rapid quenching from \( \beta \) field may give rise to fine precipitates, which considerably increases mechanical strength.

Titanium and Titanium Alloy Applications in Medicine     541

| Table 15.1. Selected Ti-base materials developed for medical applications [38] |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Material        | Tensile Strength (MPa) | Yield Strength (MPa) | Elongation (%) | Elastic Modulus (GPa) |
| Pure Ti grade 1 | 240              | 170              | 24             | 102.7            |
| Pure Ti grade 2 | 345              | 275              | 20             | 102.7            |
| Pure Ti grade 3 | 450              | 380              | 18             | 103.4            |
| Pure Ti grade 4 | 550              | 485              | 15             | 104.1            |
| Ti-6Al-4V       | 895–930          | 825–869          | 6–10           | 110–114          |
| Ti-6Al-4V ELI   | 860–965          | 795–875          | 10–15          | 101–110          |
| Ti-6Al-7Nb      | 900–1050         | 880–950          | 8.1–15         | 114              |
| Ti-5Al-2.5Fe    | 1020             | 895              | 15             | 112              |
| Ti-13Nb-13Zr    | 973–1037         | 836–908          | 10–16          | 79–84            |
| Ti-12Mo-6Zr-2Fe | 1060–1100        | 1000–1060        | 18–22          | 74–85            |
| Ti-15Mo         | 874              | 544              | 21             | 78               |
| Ti-15Mo-5Zr-3Al | 852–1100         | 838–1060         | 18–25          | 80               |
| Ti-15Mo-2.8Nb-0.2Si | 979–999          | 945–987          | 16–18          | 83               |
| Ti-35.3Nb-5.1Ta-7.1Zr | 596.7        | 547.1            | 19             | 55               |
| Ti-29Nb-13Ta-4.6Zr | 911            | 864              | 13.2           | 80               |

of second phases. Also, by using ageing processes, metastable structures obtained by rapid quenching from \( \beta \) field may give rise to fine precipitates, which considerably increases mechanical strength.

Titanium alloys present a high strength-to-weight ratio, which is higher than with most steels. While CP titanium has yield strength between 170 (grade 1) and 485 MPa (grade 4), titanium alloys may present values higher than 1500 MPa [25].

The elastic modulus or Young modulus corresponds to the stiffness of a material and is associated to the way inter-atomic forces vary with distance between atoms in the crystal structure. A comparison between both crystal structures of titanium has led to the conclusion
that HCP structure presents higher values of elastic modulus than the BCC structure. Hence, addition of β-stabilizer elements allows β phase stabilization and hence, low elastic modulus alloys. While CP titanium shows elastic modulus values close to 105 GPa, Ti-6Al-4V type α+β alloy presents values between 101 and 110 GPa, type β titanium alloys may present values as low as 55 GPa, [38]. When compared with common alloys used as biomaterials, such 316 L stainless steel (190 GPa) and Co-Cr alloys (210–253), low elastic modulus titanium alloys display a more compatible elastic behavior to that of the human bone [39]. In general, as the elastic modulus decreases, so does the mechanical strength and vice versa.

Analysis of slip systems in different crystal structures reveals that plastic deformation is easier in BCC crystal structure than in HCP structure. It explains the enhanced ductility of β phase when compared to α phase. In a HCP structure the number of slip systems is only three, while this number increases to 12 in the case of BCC structure. In addition, the ease of plastic deformation facility is directly connected to the minimum slip distance, \( b_{\text{min}} \) [25], which is given by the interatomic distance divided by the respective lattice parameter. Since, HCP structure exhibits a higher slip distance than BCC structure, it is possible to conclude that the atomic planes slip or the plastic deformation is easier in BCC structure than HCP. Hence, β type alloys present the best formability among the titanium alloys.

15.1.4 Corrosion Behavior

Corrosion resistance is one of the main properties of a metallic material applied in the human body environment and the success of an implant depends on the careful examination of this phenomenon. The performance of an implant is directly related to its ability in functioning in the aggressive body fluids. In general, these fluids consist of a series of acids and certain amount of NaCl. In normal conditions, its pH is 7, however, it may be altered due to immune system response, like in the case of a infection or inflammation. In an event of a corrosion process, the implant component may lose its integrity, leading to a failure. In addition, release of corrosion products may lead to undesirable biological reactions. Certainly, this will depend on the nature
of chemical reactions on the implant surface in view of the fact that corrosion is essentially a chemical process.

Titanium shows an excellent corrosion resistance, which is directly related to the formation of a stable and protective oxide layer, essentially TiO₂. The reactivity of titanium can be measured by its standard electrode potential (standard electromotive force (EMF) series), which is –1.63 V [5]. Such a value indicates that titanium has a high chemical reactivity and is easily oxidized, giving rise to a very adherent and thin oxide layer on the titanium surface. This oxide layer passivates the titanium, which results in a protection against further corrosion process as long as this layer is maintained. Actually, formation of passivation films on titanium does not mean cessation of corrosion processes. It means that the corrosion rate will be significantly reduced. Therefore, titanium is corrosion resistant in oxidizing environments but not resistant in reducing medium [5].

In general, anodic polarization testing is an efficient method of analyzing corrosion behavior of metallic material in a corrosive environment. Figure 15.6 depicts the anodic polarization curve for CP titanium and Ti-6Al-4V alloys, showing the applied potential as a function of resulting currenty density (versus saturated calomel electrode (SCE)), obtained with 5g/l NaCl, pH 4 solution as an electrolytic medium at 310 K. The potential was scanned at 0.1 mVs⁻¹ [11].

The anodic portion of the polarization curve allows one to evaluate the corrosion behavior of a metallic material in an electrolytic medium. Evaluation is obtained by determining the range of potentials in which passivation films are stable, and also by finding the current density. As usual, polarization tests started at a negative potential of –1.0 V vs. SCE, reaching more positive values.

In such a process, the initial sector of the anodic polarization curve refers to the beginning of a corrosion phenomenon, where the metallic material reacts with the supporting electrolyte, leading to active corrosion.

The following segment is related to the formation of an oxide passivating film, when the electric current stabilization takes place. As the potential increases, the current density also increases and eventually, the rupture of passivation film occurs. At this point, the protective layer loses its efficiency causing pitting corrosion. However, this hypothesis is not confirmed during the reverse scanning of potential.
Fig. 15.6. Polarization curves for CP titanium and Ti-6Al-4V alloy (Scan rate of 0.1 mV.s$^{-1}$)

Polarization curves obtained during forward and backward scans of potential are superimposed and no pitting potential is observed, which
allows one to conclude that both materials show outstanding resistance to corrosion.

15.2 Principal Requirements of Medical Implants

15.2.1 Introduction

Medical implants are products that have to satisfy functionality demands defined by the working environment-human body. They could be used in almost every organ of the human body. Ideally, they should have biomechanical properties comparable to those of autogenous tissues without any adverse effects. The principal requirements of all medical implants are corrosion resistance, biocompatibility, bioadhesion, biofunctionality, processability and availability. To fulfil these requirements most of the tests are directed into the study extracts from the material, offering screens for genotoxicity, carcinogenicity, reproductive toxicity, cytotoxicity, irritation, sensitivity and sterilization agent residues [40]. The consequences of corrosion are the disintegration of the implant material per se, which weakens the implant, and the harmful effect of corrosion on the surrounding tissues and organs is produced.

Medical implants are regulated and classified in order to ensure safety and effectiveness to the patient. One of the main goals of implant research and development is to predict long-term, in-vivo performance of implants. Lack of useful computer-modeling data about in-vivo performance characteristics makes the evaluation of synergistic contributions of materials, design features and therapeutic drug regimens difficult. The present trends in modern implant surgery are networking various skilled and gifted specialists such as traumatologists, orthopedists, mechanical engineers, pharmacists and others in order to get better results in research, development and implementation into practice.

15.2.2 Metallic Biomaterials

The first metal alloy developed specifically for the human body environment was the ‘vanadium steel’ that was used to manufacture bone
fracture plates (Sherman plates) and screws. Most metals that are used to make alloys for manufacturing implants such as iron (Fe), chromium (Cr), cobalt (Co), nickel (Ni), titanium (Ti), tantalum (Ta), niobium (Nb), molybdenum (Mo), and tungsten (W), can only be tolerated by the body in minute amounts. Sometimes these metallic elements, in naturally occurring forms, are essential in red blood cell functions (Fe) or synthesis of a vitamin B12 (Co), etc. but cannot be tolerated in large amounts in the body [41].

Metallic biomaterials can be divided into four subgroups: stainless steels, the cobalt-based alloys, titanium metals, and miscellaneous others (including tantalum, gold, dental amalgams and other special metals). They are very effective in binding the fractured bone, do not corrode and do not release harmful toxins when exposed to body fluids and therefore can be left inside the body for a long period of time. Their disadvantage is a much larger hardness and stiffness compared to the bone and possibility of interfering with the re-growth of the bone.

15.2.3 The Surface-Tissue Interaction

Good corrosion resistance of titanium depends upon the formation of a solid oxide layer (TiO\textsubscript{2}) to a depth of 10 nm. After the implant is inserted, it immediately reacts with body liquids that consist of water molecules, dissolved ions, and proteins as shown in Figure 15.7.

Geometry, roughness and other characteristics of the implant surface also importantly influence the surface-tissue interaction, which is considered to be dynamic. Due to these phenomena, over time new stages of biochemical formations can be developed. In the first few

---

**Fig. 15.7.** Interaction between titanium and body liquids [42]
seconds after the contact has been made, there is only water, dissolved ions, and free biomolecules in the closest proximity of the surface, but no cells. The composition of the body liquid changes continuously as inflammatory and healing processes proceed, causing changes in the composition of the adsorbed layer of biomolecules on the implant surface until it is balanced. Cells and tissues eventually contact the surface and, depending on the nature of the adsorbed layer, they respond in specific ways that may further modify the adsorbed biomolecules, [43].

Surface roughness also plays an important role in osteointegration. Osteoblast cells are more likely to get attached to rough-sand blasted surfaces, meaning lower cell numbers on the rougher surfaces, decreased rate of cellular proliferation, and increased matrix production compared to smooth surface. Experiments made by Feighan [44] showed that an average roughness increased from 0.5 to 5.9 µm also increase the interfacial shear strength from 0.48 to 3.5 MPa.

15.2.4 Machining of Titanium Alloys

Titanium alloys are among the most widely used and promising materials for medical implants. Selection of titanium alloy for implementation is determined by a combination of the most desirable characteristics including immunity to corrosion, biocompatibility, shear strength, density and osteointegration, [3]. The excellent chemical and corrosion resistance of titanium is to a large extent due to the chemical stability of its solid oxide surface layer to a depth of 10 nm, [45]. Under in vivo conditions the titanium oxide (TiO₂) is the only stable reaction product whose surface acts as catalyst for a number of chemical reactions. However, micromotion at the cement-prosthesis and cement-bone are inevitable and consequently, titanium oxide and titanium alloy particles are released in the cemented joint prosthesis. Sometimes this wear debris accumulates as periprosthetic fluid collections and triggers giant cell response around the implants, [46]. TiO₂ film, such as the ones anodically formed in aqueous electrolytes, consists mainly of anatase and is an n-type semiconductor with a low electronic charge conductivity and a high resistance to anodic current [47].

Processes of machining titanium alloys involve conventional machining operations (turning, face milling, high-speed cutting (HSC),
milling, drilling), forming operations (cold and hot forming; hydroforming, forging) and alternative machining operations (laser cutting, waterjet cutting, direct metal laser sintering). Machining operations of titanium alloys are considered to be difficult, due to its relatively high tensile strength, low ductile yield, 50% lower modulus of elasticity (104 GPa) and approximately 80% lower thermal conductivity than that of steel [48]. The lower modulus of elasticity may cause greater ‘spring back’ and deflection effect of the workpiece. Therefore, more rigid setups and greater clearances for tools are required. In the tool contact zones high pressures and temperatures occur (the tool-to-workpiece interface). The amount of heat removed by laminar chips is only approximately 25%, the rest is removed via the tool. Due to this phenomenon titanium alloys can be machined at comparatively low cutting speeds. At higher temperatures caused by friction the titanium becomes more chemically reactive and there is a tendency for titanium to »weld« to tool bits during machining operations. Overheating of the surface can result in interstitial pickup of oxygen and nitrogen, which will produce a hard and brittle alpha case. Carbides with high WC-Co content (K-grades) and high-speed steels with high cobalt content are suitable for use as cutting materials in titanium machining operations [48]. Turning operations of titanium alloys should have cutting depths as large as possible, cutting speeds \( V_C \) from 12 to 80 m/min and approximately 50% lower when high-speed steel (HSS) tools are used. The heat generated should be removed via large volumes of cooling lubricant. Chlorinated cutting fluids are not recommended because titanium can be susceptible to stress corrosion failures in the presence of chlorine. Any type of hot working or forging operation should be carried out below 925°C due to high level of titanium reactivity at high temperature.

Some medical implants are produced modularly, using different materials and processing techniques. For example, the femoral stem as part of the hip endoprostheses is produced in a combination of casting, forging and milling. The final machining operation is performed on CNC machine using CAD-CAM principle.

Good alternatives to conventional machining techniques are alternative techniques such as waterjet cutting, sintering, or direct metal laser sintering. The latter is a rapid prototyping technique enabling prompt modeling of metal parts with high bulk density on the basis
of individual three-dimensional data, including computer tomography models of anatomical structures, [49]. The concept of layer by layer building rather than removing waste material to achieve the desired geometry of a component, opens up endless possibilities of alternative manufacture of medical devices and is more environment friendly.

15.2.5 Surface Treatments and Coatings

Mechanical methods for surface treatment can be divided into methods involving removal of surface material by cutting (machining of the surface), abrasive action (grinding and polishing) and those where the treated material surface is deformed by particle blasting. Chemical methods are based mainly on chemical reactions occurring at the interface between titanium and a solution (solvent cleaning, wet chemical etching, passivation treatments and other chemical surface treatments such as Hydrogen peroxide treatment). Electrochemical surface methods are based on different chemical reactions occurring at an electrically energized surface (electrode) placed in an electrolyte (electropolishing and anodic oxidation or anodizing).

Improving the method for both wear and corrosion resistance of titanium implant surfaces in cases where the protection by natural surface oxide films is insufficient can be done through the deposition of thin films. These coatings should have a sufficiently high adherence to the substrate throughout the range of conditions to which the implant is exposed in service. They must tolerate the stress and strain variations that any particular part of the implant normally impose on the coating. The coating process must not damage the substrate and must not induce failure in the substrate or introduce impurities on the surface, which may change interface properties [47]. Coatings should be wear resistant, barrier layers preventive of substrate metal-ion release, to low-friction haemocompatible, non-thrombogenic surfaces [47]. Such surface modification could be done by various processes such as precipitations from the chemical vapor phase, Sol-Gel coatings, chemical vapor deposition (CVD) or physical vapor deposition (PVD). The properties of PVD coatings are good thickness, roughness, hardness, strength and adhesion as well as structure, morphology, stoichiometry and internal stresses. PVD processes include evaporation, sputtering, ion plating and ion implantation. They are carried out in vacuum,
at backpressures of less than 1 Pa, [47]. Speaking of the CVD methods, they involve the reaction of volatile components at the substrate surface to form a solid product. Typical CVD coatings are depositions of TiN, TiC and TiC\textsubscript{x}N\textsubscript{1-x}. The early coatings were deposited onto hard metal tools such as WC-Co. Good coating uniformity is an advantage of the CVD method, lower operating temperatures of PVD method that can be combined in the plasma-assisted CVD process. Biomaterial produced by low temperature CVD and PVD is diamond-like carbon (DLC). DLC coatings can address the main biomechanical problems with the implants currently used, e.g. friction, corrosion and biocompatibility [50].

The concept of bioactive coatings uses a principle of enabling an interfacial chemical bond between the implant and the bone tissue due to a specific biological response, [51]. Surface modifications should provide distinct properties of interaction with cell molecules, which promote the adaptation or in-growth of cells or tissue onto the surface of fixation elements of a medical implant or prevent cell interaction with the implant surface.

Suitable bioactive surface modifications are comparable to those known from the stoichiometrically passivated titanium surface, in terms of high mechanical stability against shearing forces, long-term chemical stability and corrosion resistance in a biocompatible manner. One of the most popular bioactive coatings is hydroxyapatite (HA), which is similar to the mineral phase of natural hard tissue, i.e. about 70% of the mineral fraction of a bone has a HA-like structure. HA can also be regarded as non-resorbable in a physiological environment, as long as it remains crystalline and is of high purity. It is the most stable calcium phosphate phase in aqueous solutions [52]. It has weaker mechanical properties and low resistance to fatigue failure. Surface treatments techniques for HA are plasma spraying (Vacuum Plasma Spraying-VPS) electrophoretic deposition of HA and micro-arc oxidation.

### 15.2.6 Applications in Practice

Different types of fracture repair mechanisms are known in medical practice. Incomplete fractures such as cracks, which only allow micro-motion between the fracture fragments, heal with a small amount of
fracture-line callus, known as primary healing. In contrast, complete fractures that are unstable, and therefore generate macromotion, heal with a voluminous callus stemming from the sides of the bone, known as secondary healing. The treatments can be non-surgical or surgical. Non-surgical treatments are immobilization with plaster or resin casting and bracing with a plastic apparatus. The surgical treatments of bone fractures (osteosynthesis) are divided into external fracture fixation, which does not require opening the fracture site, or internal fracture fixation, which requires opening the fracture. With external fracture fixation, the bone fragments are held in alignment by pins placed through the skin onto the skeleton, structurally supported by external bars. With internal fracture fixation, the bone fragments are held by wires, screws, plates, and/or intramedullary devices [53].

*Surgical wires* are used to reattach large fragments of bone. They are also used to provide additional stability in long-oblique or spiral fractures of long bones which have already been stabilized by other means. Straight wires are called Steinmann pins. In the case of a pin diameter less than 2.38 mm, they are named Kirschner wires. They are widely used primarily to hold fragments of bones together provisionally or permanently and to guide large screws during insertion. *Screws* are the most widely used devices for fixation of bone fragments, Figure 15.8(a). There are two types of bone screws cortical bone screws, which have small threads, and cancellous screws, which have large threads to get more thread-to-bone contact. They may have either V or buttress threads. According to their ability to penetrate the cortical screws are sub-classified further, into self-tapping and non-self-tapping. The self-tapping screws have cutting flutes that thread the pilot drill-hole during insertion. In contrast, the non-self-tapping screws require a tapped pilot drill-hole for insertion. The bone immediately adjacent to the screw often undergoes necrosis initially, but if the screw is firmly fixed when the bone revascularizes, permanent secure fixation may be achieved, [54]. This is particularly true for titanium alloy screws or screws with a roughened thread surface, with which bone ongrowth results in an increase in removal torque [54]. *Plates* are available in a wide variety of shapes and are intended to facilitate fixation of bone fragments, Figure 15.8(b). They range from the very rigid, intended to produce primary bone healing, to the relatively flexible, intended to facilitate physiological loading of bone. The rigidity and
strength of a plate in bending depends on the cross-sectional thickness and material properties of which it is made. Consequently, the weakest region in the plate is the screw hole, especially if the screw hole is left empty, due to a reduction of the cross-sectional area in this region. The effect of the material on the rigidity of the plate is defined by the elastic modulus of the material for bending, and by the shear modulus for twisting, [55]. Thus, given the same dimensions, a titanium alloy plate will be less rigid than a stainless steel plate, since the elastic modulus of each alloy is 110 GPa and 200 GPa, respectively. Intramedullary devices (IM nails or rods) are used as internal struts to stabilize long bone fractures, Figure 15.8(c). IM nails are also used for fixation of femoral neck or intertrochanteric bone fractures; however, this application requires the addition of long screws. A whole range of designs is available, going from solid to cylindrical, with shapes such as cloverleaf, diamond, and slotted cylinders. Compared to plates, IM nails are better positioned to resist multidirectional bending than a plate or an external fixator, since they are located in the center of the bone. However, their torsional resistance is less than that of the plate, [55].

Fig. 15.8. Titanium trauma medical implants
The design of an implant for joint replacement should be based on the kinematics and dynamic load transfer characteristic of the joint. The material properties, shape, and methods used for fixation of the implant to the patient determine the load transfer characteristics. This is one of the most important elements that determines long-term survival of the implant, since bone responds to changes in load transfer with a remodeling process, known as Wolff’s law. Overloading the implant-bone interface or shielding it from load transfer may result in bone resorption and subsequent loosening of the implant, [56].

The endoprosthesis for total hip replacement consists of a femoral component and an acetabular component, Figure 15.9(b). The femoral stem is divided into head, neck, and shaft. The femoral stem is made of Ti alloy or Co-Cr alloy and is fixed into a reamed medullary canal by cementation or press fitting. The femoral head is made of Co-Cr alloy, aluminum, or zirconium. Although Ti alloy heads function well under clean articulating conditions, they have fallen into disuse because of their low wear resistance to third bodies, e.g., bone or cement particles. The acetabular component is generally made of ultra-high molecular weight polyethylene (UHMWPE). The prosthesis for total knee joint replacement consists of femoral, tibial, and/or patellar components, Figure 15.9(a). Compared to the hip joint, the knee joint has a more complicated geometry and movement mechanics, and it is not intrinsically stable. In a normal knee, the center of movement

Fig. 15.9. Titanium orthopedics medical devices [58]
is controlled by the geometry of the ligaments. As the knee moves, the ligaments rotate on their bony attachments and the center of movement also moves. The eccentric movement of the knee helps distribute the load throughout the entire joint surface, [57]. Total knee replacements can be implanted with or without cement, the latter relying on porous coating for fixation. The femoral components are typically made of Co-Cr alloy and the monolithic tibial components are made of UHMWPE. In modular components, the tibial polyethylene component assembles onto a titanium alloy tibial tray. The patellar component is made of UHMWPE, and a titanium alloy back is added to components designed for uncemented use.

For maxillofacial osteosynthesis in the cranio-facial and mandibular areas titanium plate and screw systems are preferred. In order to make them pliable, many of the plates are made from CP titanium sheet that is in the soft-recrystallized condition. The corresponding screws are either made from CP titanium or alloy and can be as small as 1 mm in diameter, [59].

15.3 Shape Memory Alloys

15.3.1 Introduction

Smart materials have been given a lot of attention mainly for their innovative use in practical applications. One example of such materials is also the family of shape memory alloys (SMA) which are arguably the first well known and used smart material. Shape memory alloys possess a unique property according to which, after being deformed at one temperature, they can recover to their original shape upon being heated to a higher temperature. The effect was first discussed in the 1930s by Ölander [60] and Greninger and Mooradian [61]. The basic phenomenon of the shape memory effect was widely reported a decade later by Russian metallurgist G. Kurdjumov and also by Chang and Read [62]. However, presentation of this property to the wider public came only after the development of the nickel-titanium alloy (nitinol) by Buehler and Wang [63]. Since then, research activity in this field has been intense, and a number of alloys have been investigated, including Ag-Cd, Au-Cd, Cu-Zn, Cu-Zn-Al, Cu-Al-Ni, Cu-Sn,
Cu-Au-Zn, Ni-Al, Ti-Ni, Ti-Ni-Cu, Ni-Ti-Nb, Ti-Pd-Ni, In-Ti, In-Cd and others. Crystallography of shape memory alloys have been studied for the last four decades. Only a fraction of the available literature is listed here [64–73]. Because these materials are relatively new, some of the engineering aspects of the material are still not well understood. Many of the typical engineering descriptors, such as Young’s modulus and yield strength, do not apply to shape memory alloys since they are very strongly temperature dependent. On the other hand, a new set of descriptors must be introduced, such as stress rate and amnesia. That is why numerous constitutive models have been proposed over the last 20 years to predict thermomechanical behavior [74–87].

15.3.1.1 Thermomechanical Behavior

These materials have been shown to exhibit extremely large, recoverable strains (on the order of 10%), and it is these properties as functions of temperature and stress that allow SMAs to be utilized in many exciting and innovative applications. From a macroscopic point of view, the mechanical behavior of SMAs can be separated into two categories: the shape memory effect (SME), where large residual (apparently plastic) strain can be fully recovered upon raising the temperature after loading and unloading cycle; and the pseudoelasticity or superelasticity, where a very large (apparently plastic) strain is fully recovered after loading and unloading at constant temperature. Both effects are results of a martensite phase transformation. In a stress-free state, an SMA material at high temperatures exists in the parent phase (usually a body-centered cubic crystal structure, also referred as the austenite phase). Upon decreasing the material temperature, the crystal structure undergoes a self-accommodating crystal transformation into martensite phase (usually a face-centered cubic structure). The phase change in the unstressed formation of martensite from austenite is referred to as ‘self-accommodating’ due to the formation of multiple martensitic variants and twins that prohibits the incurrence of a transformation strain. The martensite variants, evenly distributed throughout material, are all crystallographically equivalent, differing only by habit plane. The process of self-accommodation by twinning allows an SMA material to exhibit large reversible strains with stress.
However, the process of self-accommodation in ordinary materials like stainless steel does not take place by twinning but via a mechanism called slip. Since slip is a permanent or irreversible process, the shape memory effect cannot occur in these materials. The difference between the twinning and slip process is shown in Figure 15.10.

In the stress-free state, an SMA material has four transition temperatures, designated as \( M_f, M_S, A_S, A_f \), i.e. Martensite Finish, Martensite Start, Austenite Start, and Austenite Finish, respectively. In the case of ‘Type I’ materials, temperatures are arranged in the following manner: \( M_f < M_S < A_S < A_f \). A change of temperature within the range \( M_S < T < A_S \) induces no phase changes and both phases can coexist within \( M_f < T < A_f \). With these four transformation temperatures and the concepts of self-accommodation, the shape memory effect can be adequately explained.

As an example let us consider a martensite formed from the parent phase, Figure 15.11(a), cooled under stress-free conditions through \( M_S \) and \( M_f \). This material has multiple variants and twins present, Figure 15.11(b), all crystallographically equivalent, but with different orientation (different habit plane indices). When a load applied to this material reaches a certain critical stress, the pairs of martensite twins begin ‘de-twinning’ to the stress-preferred twins, Figure 15.11(c). It means that the multiple martensite variants begin to convert to a single variant determined by alignment of the habit planes with the direction of loading, Figure 15.11(d). During this process of reorientation, the stress rises very slightly in comparison to the strain. As the single variant of martensite is thermodynamically stable at \( T < A_S \), upon unloading there is no conversion to multiple variants and only a small elastic strain is recovered, leaving the material with a large residual strain, Figure 15.11(e). The de-twinned martensite material can recover the entire residual strain by
simply heating above $A_f$; the material then transforms to the parent phase, which has no variants, and recovers to its original size and shape, Figure 15.11(f), thus creating the shape memory effect.

The pseudoelastic effect can be explained, if an SMA material is considered to be entirely in the parent phase (with $T > A_f$), Figure 15.12(a). When stress is applied to this material, there is a critical stress at which the crystal phase transformation from austenite to martensite can be induced, Figure 15.12(b). Due to the presence of stress during the transformation, specific martensite variants will be formed preferentially and at the end of transformation, the stress-induced martensite will consist of a single variant of detwinned martensite, Figure 15.12(c). During unloading, a reverse transformation to austenite occurs because of the instability of martensite at $T > A_f$ in the absence of stress, Figure 15.12(e). This recovery of high strain values upon unloading yields a characteristic hysteresis loop, diagram in Figure 15.12, which is known as pseudoelasticity or superelasticity.

Many of the possible medical applications of SMA materials in the 1980’s were attempting to use the thermally activated memory effect. However, temperature regions tolerated by the human body are
Fig. 15.12. Pseudoelasticity or superelasticity

very limited. Small compositional changes around the 50–50% of Ti-Ni ratio can make dramatic changes in the operating characteristics of the alloy. Therefore very precise control of phase transition temperatures is required. On the other hand, pseudoelasticity is ideally suited to medical applications since the temperature region of optimum effect can easily be located to encompass ambient temperature through body temperature.

15.3.2 Biocompatibility

It is important to understand the direct effects of an individual component of the alloy since it can dissolve in the body due to corrosion and it may cause local and systemic toxicity, carcinogenic effects and immune response. The cytotoxicity of elementary nickel and titanium has been widely researched, especially in the case of nickel,
which is a toxic agent and allergen, [88, 89, 90]. Nickel is known to have toxic effects on soft tissue structures at high concentrations and also appears to be harmful to bone structures, but substantially less than cobalt or vanadium, which are also routinely used in implant alloys. Experiments with toxic metal salts in cell cultures have shown decreasing toxicity in the following order: Co > V > Ni > Cr > Ti > Fe, [91]. The dietary exposure to nickel is 160–600 mg/day. Fortunately most of it is eliminated in the feces, urine and sweat. Pure nickel implanted intramuscularly or inside bone has been found to cause severe local tissue irritation and necrosis and high carcinogenic and toxic potencies. Due to corrosion of medical implants, a small amount of these metal ions is also released into distant organs. Toxic poisoning is later caused by the accumulation, processing and subsequent reaction of the host to the corrosion of the Ni-containing implant. Nickel is also one of the structural components of the metalloproteins and can enter the cell via various mechanisms. Most common Ni\(^{2+}\) ions can enter the cell utilizing the divalent cation receptor or via the support with Mg\(^{2+}\), which are both present in the plasma membrane. Nickel particles in cells can be phagocytosed, which is enhanced by their crystalline nature, negative surface energy, appropriate particle size (2–4 µm) and low solubility. Other nickel compounds formed in the body are most likely to be NiCl and NiO, and fortunately there is only a small chance that the most toxic and carcinogenic compounds like Ni\(_3\)S\(_2\), are to be formed. Nickel in soluble form, such as Ni\(^{2+}\) ions, enters through receptors or ion channels and binds to cytoplasmic proteins and does not accumulate in the cell nucleus at concentrations high enough to cause genetic consequences. These soluble Ni\(^{2+}\) ions and are rapidly cleaned from the body. However, the insoluble nickel particles containing phagocytotic vesicles fuse with lysosomes, followed by a decrease of phagocytic intravesicular pH, which releases Ni\(^{2+}\) ions from nickel containing carrier molecules. The formation of oxygen radicals, DNA damage and thereby inactivation of tumor suppressor genes is contributed by that.

On the other hand, titanium is recognized to be one of the most biocompatible materials due to the ability to form a stable titanium oxide layer on its surface. In an optimal situation, it is capable of excellent osteointegration with the bone and it is able to form a calcium phosphate-rich layer on its surface, Figure 15.13, very similar to
hydroxyapatite which also prevents corrosion. Another advantageous property is that in case of damaging the protective layer the titanium oxides and Ca-P layer regenerate.

The properties and biocompatibility of nitinol have their own characteristics which are different from those of nickel or titanium alone. In vitro NiTi biocompatibility studies on the effects of cellular tolerance and its cytotoxicity have been performed on various cell culture models [93, 94]. Human monocytes and microvascular endothelial cells were exposed to pure nickel, pure titanium, stainless steel and nitinol. Nitinol has been shown to release higher concentrations of Ni\(^{2+}\) ions in human fibroblast and osteoblast cultures, which did not affect cell growth [95–97]. Metal ion release study also revealed very low concentrations of nickel and titanium that were released from nitinol. Researchers therefore concluded that nitinol is not genotoxic.

For in vivo biocompatibility studies of nitinol effect, different experiments have been done on animals. Several in vivo nitinol biocompatibility studies that were done in the last decade disclosed no allergic reactions, no traces of alloy constituents in the surrounding tissue and no corrosion of implants. Studies of rat tibiae response to NiTi, compared with Ti-6Al-4V and stainless steel, showed that the number and area of bone contacts was low around NiTi implants, but the thickness of contact was equal to that of other implants. Normal new bone formation was seen in rats after 26 weeks after implantation. Good biocompatibility results of NiTi are attributed to the fact that implants are covered by a titanium oxide layer, where only small traces of nickel are being exposed.
15.3.2.1 Corrosion Behavior

The body is a complicated electrochemical system that constitutes an aggressive corrosion environment for implants which are surrounded by bodily fluids of an aerated solution containing 0.9\% NaCl, with minor amounts of other salts and organic compounds, serum ions, proteins and cells which all may modify the local corrosion effect. High acidity of certain bodily fluids is especially hostile for metallic implants. Acidity can increase locally in the area adjacent to an implant due to inflammatory response of surrounding tissues mediating hydrogen peroxide and reactive oxygen and nitrogen compounds. The local pH changes for infected tissues or near haematomas are relatively small, however these changes can alter biological processes and thereby the chemistry around the implant. It is known that small point corrosion or pitting prevails on surfaces of metallic implants. Another important feature is roughness of the surface which increases the reacting area of the implant and thereby add to total amount of corrosion. Therefore surface finishing is a major factor in improving corrosion resistance and consequently biocompatibility of medical devices, [98, 99].

Corrosion resistance of SMA has also been studied \textit{in vivo} on animals. Plates and stents have been implanted in dogs and sheeps for several months. Corrosion has been examined under microscope and pitting was established as predominant after the implants were removed. Thus surface treatments and coatings were introduced. The improvement of corrosion resistance was considerable, since pitting decreased in some cases from 100 microns to only 10 microns in diameter.

15.3.3 Surface of Implant

The human response to implanted materials is a property closely related to the implant surface conditions. The major problems associated with the implants currently used are inadequate implant-tissue interface properties. Parameters that characterize surface property are chemical composition, crystallinity and heterogeneity, roughness and wettability or surface free energy that is a parameter important for cell adhesion. Each parameter is of great importance to biological
response of the tissue. Another problem is implant sterilization that can remarkably modify desired parameters. Steam and dry sterilization are nowadays replaced by more advanced techniques like hydrogen peroxide plasma, ethylene oxide, and electron and γ-ray irradiation.

The surface of NiTi SMA has revealed a tendency towards preferential oxidation of titanium. This behavior is in agreement with the fact that the free enthalpy of formation of titanium oxides is negative and exceeds in absolute value the enthalpy of formation of nickel oxides by at least two to three times. The result of oxidation is an oxide layer of a thickness between 2–20 nm, which consists mainly of titanium oxides TiO₂, smaller amounts of elemental nickel Ni²⁺, and low concentrations of nickel oxides NiO. The surface chemistry and the amount of Ni may vary over a wide range, depending on the preparation method. The ratio of Ti/Ni on polished surface is around 5.5, while boiled or autoclaved items in water show decreased concentration of Ni on the surface and the Ti/Ni ratio increases up to 23 to 33, [100]. Different in vitro studies have shown how the physical, chemical and biocompatible properties of the implant surface can be improved, [101–105].

15.3.3.1 Surface Improvements

Some of the most important techniques for improving the properties of Ni-Ti alloy surfaces are: (1) Surface modification by using energy sources and chemical vapors like hydroxyapatite, laser and plasma treatment, ion implantation, TiN and TiCN chemical vapor deposits. Hydroxyapatite coatings result in the best known biocompatibility and reveal a tendency to dissolution due to its relative miscibility with body fluids. Ion implantation and laser treatments usually result in surface amorphization that improves corrosion resistance, but the obtained amorphous surface layers are often not uniform. Laser surface melting leads to an increased oxide layer, decrease of Ni dissolution and improvement of the cytocompatibility up to classical Ti level. There is also a possibility that laser melted surfaces may be enriched in nickel, and become harder than bulk and swell. TiN and TiCN coatings are known to improve corrosion resistance but large deformations caused by the shape memory effect may cause cracking of the coating. Therefore, for plates and staples a plasma-polymerized
tetrafluoroethylene has been introduced. (2) Development of bioactive surfaces is another approach to improve biocompatibility of the SMA. Human plasma fibronectin covalently immobilised to NiTi surface improved the attachment of cells while corrosion rates were reduced drastically. Studies showed NiTi surface improved with this method caused a development of Ca-P layers, which in fact eliminate the need for hydroxyapatite coatings, [102, 106]. (3) Electrochemical processing for oxidation in air/oxygen is a most common way of metal surface treatment. The technique combines electrochemical processes and oxidation in various media. Growth of native passive films that are highly adhesive and do not crack or break due to dynamic properties of SMA is promoted with this method. Oxide films obtained in air have different colors, thickness, and adhesive properties, with TiO$_2$ as a predominant oxide type. (4) Oxidation of SMA medical devices in water and steam is also one of the surface improvement techniques. Implants are preliminary chemically etched and boiled in water. The result is a surface with a very low Ni concentration, while etching removes surface material that was exposed to processing procedures and acquired various surface defects and heterogeneity. It also selectively removes nickel and oxidizes titanium. Surfaces obtained after oxidation in steam show better properties than those oxidized in water. (5) Electrochemical techniques are commonly used to passivate NiTi surfaces. Surface passivation using electropolishing is often considered as a treatment of first choice just because this technique is used for surface conditioning of stainless steel, Co-Cr alloys, etc. However, the universal techniques developed for surface passivation of various alloys used for medical purposes are not necessary efficient for NiTi.

It should also be noted that the implant surface coatings are not always beneficial. The major problem of titanium based alloys is that the formation of TiO$_2$, according to the chemical equation Ti + 2H$_2$O $\rightarrow$ TiO$_2$ + 4H$^+$ + 4e$^-$, reduces the pH level at the titanium/coating interface. This means that if the coating is composed of hydroxyapatite, it can dissolve, which gradually leads to detachment of the coating.

15.3.4 Medical Applications

The trends in modern medicine are to use less invasive surgery methods that are performed through small, leak tight portals into the body
called trocars. Medical devices made from SMAs use a different physical approach and can pull together, dilate, constrict, push apart and have made difficult or problematic tasks in surgery quite feasible. Therefore unique properties of SMAs are utilized in a wide range of medical applications. Some of the devices used in various medical applications are listed below.

*Stents* are most rapidly growing cardiovascular SMA cylindrical mesh tubes that are inserted into blood vessels to maintain the inner diameter of a blood vessel. The product has been developed in response to limitations of balloon angioplasty, which resulted in repeated blockages of the vessel in the same area. Ni-Ti alloys have also become the material of choice for superelastic *self-expanding (SE) stents* that are used for a treatment of the superficial femoral artery disease, Figure 15.14(a). The SE nitinol stents are produced in the open state mainly with laser cut tubing and later compressed and inserted into the catheter. They can also be produced from wire and laser welded or coiled striped etched sheet. Before the compression stage, the surface of the stent is electrochemically polished and passivated to prescribed quality. Deployment of the SE stent is made with the catheter. During the operation procedure, when the catheter is in the correct position in the vessel, the SE stent is pushed out and then it expands against the inner diameter of the vessel due to a rise in temperature (thermally triggered device). This opens the iliac artery to aid in the normal flow of blood. The delivery catheter is then removed, leaving the stent within the patient’s artery. Recent research has shown that implantation of a self-expanding stent provides better outcomes, for the time being, than balloon angioplasty, [107–109].

The *Simon Inferior Vena Cava (IVC) filter* was the first SMA cardiovascular device. It is used for blood vessel interruption for preventing pulmonary embolism via placement in the vena cava. The Simon filter is filtering clots that travel inside bloodstream, [110]. The device is made of SMA wire curved similarly to an umbrella that traps the clots that are better dissolved in time by the bloodstream. For insertion, the device is exploiting the shape memory effect, i.e. the original form in the martensitic state is deformed and mounted into a catheter. When the device is released, the body’s heat causes the filter to return to its predetermined shape. The *Septal Occlusion System* is indicated for use in patients with complex ventricular septal defects (VSD) of
significant size to warrant closures that are considered to be at high risk for standard transatrial or transarterial surgical closure based on anatomical conditions and/or based on overall medical condition. The system consists of two primary components; a permanent implant, which is constructed of an SMA wire framework to which polyester fabric is attached, and a coaxial polyurethane catheter designed specifically to facilitate attachment, loading, delivery and deployment to the defect, [111]. The implant is placed by advancing the delivery catheter through blood vessels to the site of the defect inside the heart. The implant remains in the heart and the delivery catheter is removed. Instruments for minimally invasive surgery used in endoscopic surgery could not be feasible without implementation of SMA materials. The most representative instruments such as guidewires, dilatators and retrieval baskets exploit good kink resistance of SMAs, [112]. Open-heart stabilizers are instruments similar to a steerable joint endoscopic camera. In order to perform bypass operations on the open heart stabilizers are used to prevent regional heart movements while performing surgery. Another employment of the unique properties of SMAs such as constant force and superelasticity in heart surgery is a tissue spreader used to spread fatty tissue of the heart, Figure 15.14(b).

In general, conventional orthopedic implants by far exceed any other SMA implant in weight or volume. They are used as fracture fixation devices, which may or may not be removed and as joint replacement devices. Bone and nitinol have similar stress-strain characteristics, which makes nitinol a perfect material for production of bone fixation plates, nails and other trauma implants, [113]. In traditional trauma surgery bone plates and nails fixated with screws are used for fixation of broken bones. Shape memory fixators are one step forward applying a necessary constant force to faster fracture healing. The SMA embracing fixator consists of a body and sawtooth arms, [114]. It embraces the bone about 2/3 of the circumference, Figure 15.14(c). The free ends of the arms that exceed the semi-circle are bent more medially to match the requirement fixation of a long tubular body whose cross section is not a regular circle. The applied axial compression stress is beneficial for enhancing healing and reducing segmental osteoporosis caused by a stress shielding effect. Its martensitic transformation temperature is 4–7°C
and shape recovery temperature is around the body’s normal temperature, 37°C. Similar to the embracing fixator is the so called Swan-Like Memory-Compressive Connector (SMC) for treatment of fracture and nonunion of upper limb diaphysis. The working principle of the device is similar with one important improvement. The SMC trauma implant is able to put constant axial stress to a fractured bone, [115]. For fixation of tibial and femoral fractures nails fixated with screws are normally used. New SMA inter-locking intramedullary nails have many advantages compared to traditional ones. For example, when cooled SMA inter-locking nails are inserted into a cavity, guiding nails are extracted and body heat causes bending of nails into a preset shape applying constant pressure in the axial direction of the fractured bone, [116]. The SMA effect is also used in surgical fixators made from wire. Certain devices that have been developed to fix vertebra in spine fractures are similar to an ordinary staple. Staple shaped compression medical devices are also used for internal bone fixation, [117]. The compression staple is one of most simple and broadly used SMA devices in medicine, Figure 15.14(d). Since its introduction in 1981, over a thousand patients have been all successfully treated using this device. The SMA Patellar Concentrator was designed to treat patellar fractures, Figure 15.14(e). The device exerts continuous compression for the fixation of patella fracture. The shape of patellar concentrator consists of two basic patellae claws, conjunctive waist and three apex patellae claws. The thickness of the device may vary between 1.8 and 2.2 mm depending on different sizes of the concentrator. In clinical surgery, the claws are unfolded and put over fractured patella. Exposed to body temperature, the device tends to recover to its original state resulting in a recovery compressive force, [118].

Dentists are using devices made from SMA for different purposes. NiTi based SMA material performs exceptionally at high strains in strain-controlled environments, such as exemplified with dental drills for root canal procedures. The advantage of these drills is that they can be bent to rather large strains and still accommodate the high cyclic rotations, [111]. Superelastic SMA wires have found wide use as orthodontic wires as well, Figure 15(a). NiTi orthodontic archwire was first produced in batches and clinically used in China in the beginning of 1980’s, [119]. Due to its unique property-superelasticity, the
wire exerts gentle and retentive force to teeth, which is superior to stainless steel wire. Shape memory bracelet do not require as frequent visits to the dentist as the classical ones because of their ability to self adjust. The therapeutic period is therefore cut down by 50%.

![Fig. 15.14. Examples of nitinol medical devices](image)

![Fig. 15.15. Dental applications of nitinol](image)
Lately a special fixator for mounting bridgework has been developed, Figure 15.15(b). A small piece of SMA metal is notch on both sides and placed between teeth and bridgework. As the temperature rises the notched area of metal is expanded on both sides causing a permanent hold of bridgework. The tooth fixator can also be used to prevent a loose tooth from falling out.

15.4 Conclusions

The use of titanium alloys as biomaterials has been growing due to their reduced elastic modulus, superior biocompatibility, high strength to weight ratio and enhanced corrosion resistance when compared to more conventional stainless steel and Co-Cr alloys. Ti-6Al-4V (and Ti-6Al-4V ELI), the most common titanium alloy is still the most extensively used titanium alloy for medical applications. However, V and Al have been found to be toxic to the human body. In this context, β titanium alloys have been studied and developed and due to their high strength, biocompatible behavior, low elastic modulus, superior corrosion resistance and good formability, they are likely to replace the classic α+β type Ti-6Al-4V for medical applications in the near future.

SMA implants and medical devices have been successful because they offer a possibility of performing less invasive surgeries. Nitinol wires in medical instruments are more kink resistant and have smaller diameters compared to stainless steel 316L or polymer devices. Research to develop composite materials, containing SMA that will prove cost efficient and porous SMAs that will enable the transport of body fluids from the outside to inside of the bone is currently underway.

References

1. H. Sibum, Titanium and titanium alloys – from raw material to semi-finished products, Advanced Engineering Materials 5(6) (2003) 393.
2. K. Wang, The use of titanium for medical applications in the USA, Materials Science and Engineering A 213 (1996) 134.
3. H.J. Rack and J.I. Qazi, Titanium alloys for biomedical applications, Materials Science and Engineering C 26 (2006) 1269.
4. M. Niinomi, Recent metallic materials for biomedical applications, Metallurgical and Materials Transactions 33A (2002) 477.
5. G. Lütjering and J.C. Williams, Titanium, Springer-Verlag, Berlin, 2003.
6. M. Long, H.J. Rack, Titanium alloys in total joint replacement – a materials science perspective, Biomaterials, 19 (1998) 1621.
7. K.S. Katti, Biomaterials in total joint replacement, Colloids and Surfaces B: Biointerfaces 39 (2004) 133.
8. J.A. Disegi, Titanium alloys for fracture fixation implants, Injury, International Journal of The Care of the Injured 31 (2000) S-D14.
9. G. He and M. Hagiwara, Ti alloy design strategy for biomedical applications, Materials Science and Engineering C 26 (2006) 14.
10. B.P. Bannon and E.E. Mild, Titanium Alloys for Biomaterial Application: An Overview, Titanium Alloys in Surgical Implants, ASTM STP 796, H.A. Luckey and F. Kubli, Jr, Eds., American Society for Testing and materials, 1983, pp. 7–15.
11. V. Oliveira, R.R. Chaves, R. Bertazzoli and R. Caram, Preparation and characterization of Ti-Al-Nb orthopedic implants, Brazilian Journal of Chemical Engineering 17 (1998) 326.
12. R.R. Boyer, Ana overview on the use of titanium in the aerospace industry, Materials Science and Engineering A 213 (1996) 103.
13. J.G. Ferrero, Candidate materials for high-strength fastener applications in both the aerospace and automotive industries, Journal of Materials Engineering and Performance 14 (2005) 691.
14. M. Semlitsch, F. Staub and H. Weber, Titanium-aluminum-niobium alloy, development for biocompatible, high-strength surgical implants, Biomedizinische Technik 30 (1985) 334.
15. T.P. Vail, R.R. Glisson, T.D. Koukoubis and F. Guilak, The effect of hip stem material modulus on surface strain in human femora, Journal of Biomechanics 31 (1998) 619.
16. M. Niinomi, T. Akahori, T. Takeuchi, S. Katsura, H. Fukui and H. Toda, Mechanical properties and cytotoxicity of new beta type titanium alloy with low melting points for dental applications, Materials Science and Engineering C 25 (2005) 417.
17. M. Kikuchi, M. Takahashi and O. Okuno, Elastic moduli of cast Ti-Au, Ti-Ag, and Ti-Cu alloys, Dental Materials 22 (2006) 641.
18. H.-S. kim, W.-Y. Kim and S.-H. Lim, Microstructure and elastic modulus of Ti-Nb-Si ternary alloys for biomedical applications, Scripta Materialia 54 (2006) 887.
19. S. Gross and E.W. Abel, A finite element analysis of hollow stemmed hip prostheses as a means of reducing stress shielding of the femur, Journal of Biomechanics 34 (2001) 995.
20. Y.L. Hao, M. Niinomi, D. Kuroda, K. Fukunaga, Y.L. Zhou and R. Yang, Aging response of the Young’s modulus and mechanical properties of Ti-29Nb-13Ta-4.6Zr, Metallurgical and Materials Transactions 34A (2003) 1007.
21. Y.L. Hao, M. Niinomi, D. Kuroda, K. Fukunaga, Y.L. Zhou, R. Yang and A. Suzuki, Young’s modulus and mechanical properties of Ti-29Nb-13Ta-4.6Zr in relation to $\alpha''$ martensite, Metallurgical and Materials Transactions 33A (2002) 3137.

22. B. Gunawarman, M. Niinomi, T. Akahori, T. Souma, M. Ikeda and H. Toda, Mechanical properties and microstructures of low cost $\beta$ titanium alloys for healthcare applications, Materials Science and Engineering C 25 (2005) 304.

23. N. Sakaguchi, M. Niinomi, T. Akahori, J. Takeda and H. Toda, Relationship between tensile deformation behavior and microstructure in Ti-Nb-Ta-Zr, Materials Science and Engineering C 25 (2005) 363.

24. D. Kuroda, M. Niinomi, M. Morinaga, Y. Kato and T. Yashiro, Design and mechanical properties of new $\beta$ type titanium alloys for implant materials, Materials Science and Engineering A 243 (1998) 244.

25. M. Peters, H. Hemptenmacher, J. Kumpfert and C. Leyens, in: C. Leyens and P. Ari-Gur and S.L. Semiatin, Evolution of microstructure, macrotexture and microtexture during hot rolling of Ti-6Al-4V, Materials Science and Engineering A 257 (1998) 118.

26. G. Lütjering, Property optimization through microstructural control in titanium and aluminum alloys, Materials Science and Engineering A 263 (1999) 117.

27. Y.V.R.K., Prasad and T. Seshacharyulu, Processing maps for hot working of titanium alloys, Materials Science and Engineering A 243 (1998) 82.

28. H.L. Freese, M.G. Volas and J.R. Wood, in: D.M. Brunette, P. Tengvall, M. Textor and P. Thomsen (Eds.), Titanium in Medicine, Springer, 2001, p. 25–51.

29. F.H. Froes and H.B. Bomberger, The beta titanium alloys, Journal of Metals 37 (1985) 28.

30. O.P. Karasevskaya, O.M. Ivasishin, S.L. Semiatin and V. Yu. Matviychuk, Deformation behavior of beta-titanium alloys, Materials Science and Engineering A 354 (2003) 121.

31. D.J. Lin, J.H. Chern and C.P. Ju, Effect of omega phase on deformation behavior of Ti-7.5Mo-xFe alloys, Materials Chemistry and Physics 76 (2002) 191.

32. D.L. Moffat and D.C. Larbalestier, The competition between the alpha and omega phases in aged Ti-Nb alloys, Metallurgical Transactions 19A (1988) 1687.

33. H.M. Flower, S.D. Henry and D.R.F. West, The $\beta \Rightarrow \alpha$ transformation in dilute Ti-Mo alloys, Journal of Materials Science 9 (1974) 57.

34. X. Tang, T. Ahmed and H.J. Rack, Phase transformations in Ti-Nb-Ta and Ti-Nb-Ta-Zr alloys, Journal of Materials Science 35 (2000) 1805.

35. A.V. Dobromyslov and V.A. Elkin, Martensitic transformation and metastable b-phase in binary titanium alloys with d-metals of 4–6 periods, Materials Science and Engineering A 354 (2003) 121.
37. A.V. Dobromyslov and V.A. Elkin, The orthorhombic $\alpha''$-phase in binary titanium base alloys with d-metals of V-VIII groups, Materials Science and Engineering A (in press).
38. M. Niinomi, Mechanical properties of biomedical titanium alloys, Materials Science and Engineering A 243 (1998) 231.
39. J.B. Brunski, in: B.D. Ratner, A.S. Hoffman, F.J. Schoen and J.E. Lemons, (Eds.), Biomaterials Science – An Introduction to Materials in Medicine, Elsevier Academic Press, San Diego, 2004, p. 137–153.
40. F. Wataria, A. Yokoyamaa, M. Omorib, T. Hiraic, H. Kondoa, M. Uoa, T. Kawasakia. Biocompatibility of materials and development to functionally graded implant for bio-medical application, Composites Science and Technology, Vol. 64 (2004) 893–908.
41. J. Black. Biological Performance of Materials, 2nd ed., New York: M. Dekker, Inc., (1992).
42. T. Kokubo, H.M. Kim, M. Kawashita. Novel bioactive materials with different mechanical properties, Biomaterials 24/13 (2003) 2161–2175.
43. J.B. Park, Y.K. Kim. Metallic Biomaterials, Chapter 37. In The Biomedical Engineering HandBook, 2nd ed., J.D. Bronzino, B. Raton (eds.), CRC Press LLC, (2000).
44. J.E. Feighan, V.M. Goldberg, D. Davy, J.A. Parr, S. Stevenson. The influence of surface blasting on the incorporation of titanium-alloy implants in a rabbit intramedullary model, 77A, (1995) 1380–1395.
45. P. Tengvall, I. Lundstrom. Physico-chemical considerations of titanium as a biomaterial, Clin. Materials 9, (1992) 115–134.
46. V.E. Henrich, P.A. Cox. The Surface Science of Metal Oxides. Cambridge University Press, Cambridge (1994).
47. R. Thull, D. Grant. Physical and Chemical Vapor Deposition and Plasma-assisted Techniques for Coating Titanium. In Titanium in Medicine, D.M. Brunette, P. Tengvall, M. Textor, P. Thomsen (eds.), Springer-Verlang Berlin Heidelberg (2001) 284–335.
48. F. Klocke. Manufacturing Technology I, WZL-RWTH, Aachen (2001).
49. D.A. Hollander, M. von Walter, T. Wirtz, R. Sellei, B. S. Rohlfing, O. Paar, H.J. Erli. Structural, mechanical and in vitro characterization of individually structured Ti–6Al–4V produced by direct laser forming, Biomaterials, Vol. 27 (2006) 955–963.
50. M.M. Morshed, B.P. McNamara, D.C. Cameron, M.S.J. Hashmi. Stress and adhesion in DLC coatings on 316L stainless steel deposited by a neutral beam source, J. of Materials Processing Technology, Vol. 143 (2003) 922–926.
51. L.L. Hench, R.J. Splittr, W.C. Allen, T.K. Greenlec. Bonding mechanisms at the interface of ceramic prosthetic materials, J. Biomed. Mater. Res. Symp. 2, (1971) 117–141.
52. K. de Groot, C.P.A.T. Klein, J.G.C. Wolke, J.M.A. de Blieck-Hogervorst. Plasma-sprayed coatings of calcium phosphate, CRC Handbook of Bioactive Ceramics, CRC Press, Boston, 2, (1990) 133–142.
53. A. Hulth. Current concepts of fracture healing. Clin Orthop (1989) 249–265.
54. P. Hutzschenreuter, H. Brümmer. Screw design and stability. In Current concepts of Internal Fixation, H. Uhthoff (ed.), Springer-Verlag, Berlin (1980) 244–250.
55. G.V.B. Cochran. Biomechanics of orthopaedic Structures. In Primer in Orthopaedic Biomechanics, Churchill Livingstone, New York (1982) 143–215.
56. A. Sarmiento, E. Ebramzadeh, W.J. Gogan. Cup containment and orientation in cemented total hip arthroplasties. J. Bone Joint Surg 72B(6), (1990) 996.
57. A.H. Burstein, T.H. Wright. Biomechanics. In J. Insall, R Windsor, W Scott (eds.), Surgery of the knee, 2nd ed., Churchill Livingstone, New York (1993) Vol. 7, 43–62.
58. Lima-Lto SpA, Medical Systems, Via Nazionale 52, 33030 Villanova di San Daniele del Friuli (Udine), Italy. URL: http://www.lima.it/english/medical_syst.html
59. M.S. Perren, O.E.M. Pohler, E. Schneider. Titanium as Implant Material for Osteosynthesis Applications. In Titanium in Medicine, D.M. Brunette, P. Tengvall, M. Textor, P. Thomsen (eds.), Springer-Verlag Berlin Heidelberg (2001) 772–823.
60. A. Olander. An electrochemical investigation of solid cadmium-gold alloys, J. Am. Chem. Soc., 54 (1932) 3819–3833.
61. A.B. Greninger, V.G. Mooradian. Strain transformation in metastable beta copper-zinc and beta copper-tin alloys, AIME 128 (1938) 337–368.
62. L.C. Chang, T.A. Read. Plastic deformation and diffusionless phase changes in metals-the gold-cadmium beta phase, Am. Inst. Min. Metall. Eng., J. Met. 191/1 (1951) 47–52.
63. W.J. Buehler, F.E. Wang. A summary of recent research on the Nitinol alloys and their potential application in ocean engineering, Journal of Ocean Engineering 1 (1967) 105–108.
64. C.M. Wayman. Introduction to the crystallography of martensitic transformations, The Macmillan Company, 1964.
65. K. Otsuka, C.M. Wayman. Shape memory materials, Cambridge University Press, 1998.
66. M.S. Wechsler, D.S. Liberman, T.A. Read. On the theory of the formation of martensite. Trans. AIME 197 (1953) 1503–1515.
67. J.S. Bowles, J.K. Mackenzie. The crystallography of martensite transformations I, Acta Metallurgica 2 (1954) 129–137
68. T. Saburi, C.M. Wayman. Crystallographic similarities in shape memory martensites, Acta Metallurgica 27/6 (1979) 979–995.
69. K. Adachi, J. Perkins, C.M. Wayman. Type II twins in self-accommodating martensite plate variants in a Cu-Zn-Al shape memory alloy, Acta Metallurgica 34/12 (1986) 2471–2485.
70. R.D. James, K.F. Hane. Martensitic transformations and shape-memory materials, Acta Materialia 48/1 (2000) 197–222.
71. Madangopal Krishnan. The self accommodating martensitic microstructure of Ni-Ti shape memory alloys, Acta Materialia 46/4 (1998) 1439–1457.
72. T. Inamura, Y. Kinoshita, J.I. Kim, H.Y. Kim, H. Hosoda, K. Wakashima, S. Miyazaki. Effect of {0 0 1}<1 1 0> texture on superelastic strain of Ti-Nb-Al biomedical shape memory alloys, Materials Science and Engineering A (2006) In Press.
73. K. Bhattacharya. Microstructure of martensite: why it forms and how it gives rise to the shape-memory effect, Oxford Series on Materials Modelling, 1st Ed., Oxford University Press, Oxford 2003.
74. R. Stalmans, L. Delaey, J. Van Humbeeck. Generation of recovery stresses: thermodynamic modelling and experimental verification, J. de Phys. IV 7 (1997) 47–52.
75. G.R. Barsch, J.A. Krumhansl. Twin boundaries in ferroelastic media without interface dislocations, Phys. Rev. Lett. 53/11 (1984) 1069–1072.
76. F. Falk. Model free energy, mechanics, and thermodynamics of shape memory alloys, Acta Metall. 28 (1980) 1773-1780.
77. G.A. Maugin, S. Cadet. Existence of solitary waves in martensitic alloys, Int. J. Eng. Sci. 29/2 (1991) 243–258.
78. L.C. Brinson, R. Lammering. Finite element analysis of the behavior of shape memory alloys and their applications, Int. J. Solids and Struct., 30/23 (1993) 3261–3280.
79. Y. Ivshin, T.J. Pence. A thermomechanical model for a one variant shape memory material, J. Intell. Mat. Syst. and Struct., 5/7 (1993) 455–473.
80. C. Liang, C.A. Rogers. One-dimensional thermomechanical constitutive relations for shape memory materials, J. Intell. Mater. Syst. and Struct. 1/2 (1990) 207–234.
81. J.G. Boyd, D.C. Lagoudas. Thermomechanical response of shape memory composites, J. Intell. Mater. Syst. and Struct. 5 (1994) 333-346.
82. K. Tanaka. A thermomechanical sketch of shape memory effect: One-dimensional tensile behavior, Res Mech. 18 (1986) 251-263.
83. L.C. Brinson. One-dimensional constitutive behavior of shape memory alloys: thermomechanical derivation with non-constant material functions and redefined martensite internal variable, J. Intell. Mater. Syst. and Struct. 4 (1993) 229–242.
84. J. Lubliner, F. Auricchio. Generalized plasticity and shape-memory alloys, Int. J. Solids and Structures, 33/7 (1996) 991–1003.
85. V.P. Panoskaltsis, S. Bahuguna, D. Soldatos. On the thermomechanical modeling of shape memory alloys, Int. J. Non-Linear Mech. 39/5 (2004) 709–722
86. Q.P. Sun, K.C. Hwang. Micromechanics constitutive description of thermoeelastic martensitic transformations, Advances in Applied Mechanics 31 (1994) 249–298.
87. F. Kosel, T. Videnic. Generalized plasticity and uniaxial constrained recovery in shape memory alloys, Mech. Adv. Mater. Struc., 14/1 (2007) 3–12.
88. E. Denkhaus, K. Salnikow; Nickel essentiality, toxicity, and carcinogenicity, Critical Reviews in Oncology/Hematology 42 (2002) 35–56.
89. E. Nieboer RT. Tom, WE. Sanford. Nickel metabolism in man and animals. In: Nickel and Its Role in Biology: Metal Ions in Biological Systems, Vol 23 (Sigel H, ed). New York: Marcel Dekker, 1988, 91–121.
90. G. Glenn E. Fletcher, Franco D. Rossetto, John Turnbull, and Evert Nieboer; Toxicity, Uptake, and Mutagenicity of Particulate and Soluble Nickel Compounds, 1994, Environ Health Perspect 102(Suppl 3), 69–79.
91. A. Yamamoto, R. Honma, M. Sumita. Cytotoxicity evaluation of 43 metal salts using murine fibroblasts and osteoblastic cells, Journal of Biomedical Materials Research, 39 (1998) 331–340.
92. C. Combes, C. Rey, M. Freche. XPS and IR study of dicalcium phosphate dihydrate nucleation on titanium surfaces, Colloids and Surfaces B: Biointerfaces, 11/1–2 (1998) 15–27.
93. C. Shih, S. Lin, K. Chung, Y. Chen, Y. Su, S. Lai, G.Wu, C. Kwok and K. Chung. The cytotoxicity of corrosion products of Nitinol stent wires on cultured smooth muscle cells. J. Biomed. Mater. Res. 52 (2000) 395–403
94. D.J. Wever, A.G. Veldhuizen, M.M. Sanders, J.M Schakenraad, J.R. Horn. Cytotoxic, allergic and genotoxic activity of a nickel-titanium alloy, Biomaterials, 18 (1997) 1115–1120.
95. I.C. Wataha, P.E. Lockwood, M. Marek, M. Ghazi. Ability of Ni-containing biomedical alloys to activate monocytes and endothelial cells in vitro, Journal of Biomedical Materials Research, 45 (1999) 251–257.
96. J. Ryhänen, E. Niemi, W. Serlo, E. Niemelä, P. Sandvik, H. Pernu, T. Salo. Biocompatibility of nickel-titanium shape memory metal and its corrosion behavior in human cell cultures. J Biomed Mat Res 35 (1997) 451–457.
97. C. Wirth, V. Comte, C. Lagneau, P. Exbrayat, M. Lissac, N. Jaffrezic-Renault, L. Ponsonnet. Nitinol surface roughness modulates in vitro cell response: a comparison between fibroblasts and osteoblasts, Materials Science and Engineering: C, 25 (2005) 51–60.
98. C. Trepanier, T. Leung, M. Tabrizian, L’H. Yahia, J. Bienvenu, J. Tanguay, D. Piron and L. Bilodeau. Preliminary investigation of the effect of surface treatment on biological response to shape memory NiTi stents. J. Biomed. Mater. Res. 48 (1999) 165–171.
99. S.A. Shabalovskaya. Surface, corrosion and biocompatibility aspects of Nitinol as an implant material. Bio-Medical Materials and Engineering 12 (2002) 69–109.
100. S.A. Shabalovskaya, On the nature of the biocompatibility and on medical applications of NiTi shape memory and superelastic alloys. Biomed.Mater.Eng. 6 (1996) 267–289.
101. V.M. Frauchiger, F. Schlottig, B. Gasser, M. Textor. Anodic plasma-chemical treatment of CP titanium surfaces for biomedical applications, Biomaterials, 25 (2004) 593–606.
102. X. Lu, Z. Zhao, Y. Leng. Biomimetic calcium phosphate coatings on nitric-acid-treated titanium surfaces, Materials Science and Engineering: C, In Press (2006).
103. J. Park, D.J. Kim, Y.K. Kim, K.H. Lee, K.H. Lee, H. Lee, S. Ahn. Improvement of the biocompatibility and mechanical properties of surgical tools with TiN coating by PACVD, Thin Solid Films, Volume 435 Issues 1–2 (2003) 102–107.
104. N. Shevchenko, M.T. Pham, M.F. Maitz. Studies of surface modified NiTi alloy, Applied Surface Science, 235 (2004) 126–131.
105. K. Endo. Chemical modification of metallic implant surfaces with biofunctional proteins (Part 1). Molecular structure and biological activity of a modified NiTi alloy surface. Dent. Mater. J. 14 (1995) 185–198.
106. F. Liu, F. Wang, T. Shimizu, K. Igarashi, L. Zhao. Hydroxyapatite formation on oxide films containing Ca and P by hydrothermal treatment, Ceramics International, Volume 32/5 (2006) 527–531.
107. M. Schillinger, S. Sabeti, C. Loewe. Balloon angioplasty versus implantation of nitinol stents in the superficial femoral artery, Journal of Vascular Surgery, Volume 44 Issue 3 (2006) 684.
108. B. Rapp. Nitinol for stents, Materials Today, Vol. 7 Issue 5 (2004) 13.
109. S. Tyagi, S. Singh, S. Mukhopadhyay, U. A. Kaul. Self- and balloon-expandable stent implantation for severe native coarctation of aorta in adults, American Heart Journal, Volume 146 Issue 5 (2003) 920–928.
110. M. Simon, R. Kaplow, E. Salzman, D. Freiman. A vena cava filter using thermal shape memory alloy Experimental aspects, Radiology, Volume 125 (1977) 87–94.
111. T. Duerig, A. Pelton, D. Stöckel. An overview of nitinol medical applications, Materials Science and Engineering, A273–275 (1999) 149–160.
112. H. Fischer, B. Vogel, A. Grünhagen, K.P. Brhel, M. Kaiser. Applications of Shape-Memory Alloys in Medical Instruments, Materials Science Forum, V. 394-395 (2002) 9–16.
113. A.R. Pelton, D. Stöckel, T.W. Duerig. Medical Uses of Nitinol, Materials Science Forum. V. 327-328 (2000) 63–70.
114. K. Dai, X. Wu, X. Zu. An Investigation of the Selective Stress-Shielding Effect of Shape-Memory Sawtooth-Arm Embracing Fixator, Materials Science Forum, V. 394-395 (2002) 17–24.
115. C. Zhang, S. Xu, J. Wang, B. Yu, Q. Zhang. Design and Clinical Applications of Swan-Like Memory-Compressive Connector for Upper-Limb Diaphysis, Materials Science Forum, V. 394-395 (2002) 33–36.
116. G. Da, T. Wang, Y. Liu, C. Wang. Surgical Treatment of Tibial and Femoral Factures with TiNi Shape-Memory Alloy Interlocking Intramedullary Nails, Materials Science Forum, V. 394-395 (2002) 37–40.
117. C. Song, T.G. Frank, P.A. Campbell, A. Cuschieri. Thermal Modelling of Shape-Memory Alloy Fixator for Minimal-Access Surgery, Materials Science Forum, V. 394-395 (2002) 53–56.
118. S. Xu, C. Zhang, S. Li, J. Su, J. Wang. Three-Dimensional Finite Element Analysis of Nitinol Patellar Concentrator, Materials Science Forum, V. 394–395 (2002) 45–48.

119. Y. Chu, K. Dai, M. Zhu, X. Mi. Medical Application of NiTi Shape Memory Alloy in China, Materials Science Forum, V. 327-328 (2000) 55–62.