Review

Anodic TiO$_2$ Nanotubes: Tailoring Osteoinduction via Drug Delivery

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Abstract: TiO$_2$ nanostructures and more specifically nanotubes have gained significant attention in biomedical applications, due to their controlled nanoscale topography in the sub-100 nm range, high surface area, chemical resistance, and biocompatibility. Here we review the crucial aspects related to morphology and properties of TiO$_2$ nanotubes obtained by electrochemical anodization of titanium for the biomedical field. Following the discussion of TiO$_2$ nanotopographical characterization, the advantages of anodic TiO$_2$ nanotubes will be introduced, such as their high surface area controlled by the morphological parameters (diameter and length), which provides better adsorption/linkage of bioactive molecules. We further discuss the key interactions with bone-related cells including osteoblast and stem cells in in vitro cell culture conditions, thus evaluating the cell response on various nanotubular structures. In addition, the synergistic effects of electrical stimulation on cells for enhancing bone formation combining with the nanoscale environmental cues from nanotopography will be further discussed. The present review also overviews the current state of drug delivery applications using TiO$_2$ nanotubes for increased osseointegration and discusses the advantages, drawbacks, and prospects of drug delivery applications via these anodic TiO$_2$ nanotubes.

Keywords: anodic TiO$_2$; TiO$_2$ nanotubes; osteoinduction; electric field; drug delivery

1. Introduction

Titanium (Ti) and titanium-based alloys are one of the most widely used metallic materials in biomedical applications, e.g., in implants, as they possess by far the most perfect mix of properties, including high biocompatibility and corrosion resistance, good tensile strength, as well as flexibility [1,2]. The high biocompatibility of Ti arises from its inertness and chemical resistance due to the low electrical conductivity that contributes to the electrochemical oxidation of Ti, forming a thin passive oxide barrier layer. The latter, in turn, leads to the high resistance of Ti to corrosion [1,3], as well as being responsible for its high surface energy characteristics [4].

The choice of metallic biomaterials between pure Ti or other Ti alloys depends on the targeted application, e.g., Ti or Ti6Al7Nb, Ti6Al4V, Ti13Cu4.5Ni, Ti25Pd5Cr, or the more recent TiNi, TiNiAg, TiZr for dental implants [1,2] and Ti6Al4V or more recently Ti6Al7Nb or Ti5Al2.5Fe for orthopedic applications [2] are preferred. The recent paradigm shift in the biomedical field from the microscale to nanoscale topography [5,6] was also applied to nanostructures on Ti and its alloys [7–10].

Simultaneously with the shift to miniaturization of surface topography, the spotlight of the biomaterial field moved also to more complex systems based on interdisciplinary works, such as materials design, surface functionalization, nanomanufacturing of devices, and tissue engineering [11]. Among these, the modification of the surface properties
of biomaterials (surface nanoscale topography, physical properties, and surface chemistry) [9,12,13] is a critical factor affecting the biocompatibility and efficiency of drug delivery and other biomedical applications [10,11,13–19]. Especially for biomaterials targeting dental/orthopedic bone regeneration, a good osteoinduction and optimal osseointegration with necessary mechanical properties are key aspects to consider. Recent studies showed that the surface properties of implantable biomaterials play a vital role in both (i) establishing a stable fixation of the implant and the osseointegration by preventing fibrous tissue engagement, and (ii) are accountable for local immune responses during wound healing and regeneration processes [11,16,17,20,21].

The relevant surface modification methods include so far: (a) mechanical methods (grinding, machining, etc.) [13,22], (b) acidic treatments (e.g., sulfuric or hydrochloric acid for cleaning, inducing some roughness and a more efficient deposition of additional bioactive layers) [13,23,24], (c) hydrogen peroxide [25,26], (d) hydroxyapatite coatings on Ti either by micro-arc oxidation, sol-gel methods or plasma spraying [27–30], (e) silver (Ag) coatings on Ti by plasma sputtering of other Ag containing diamond-like carbon coatings on Ti [31–33], (f) electrochemical anodization allowing a nanostructured layer (nanopores, nanotubes or mesosponge) to grow directly on the metallic biomaterial (Ti or Ti alloys) [9,34]. These mechanical, physical, or chemical methods enable morphological surface modification and can be combined with the addition of a coating layer on the Ti or Ti alloys surface.

Among the available surface modification techniques, electrochemical anodization is one of the most widely used approaches for Ti, already used for nanostructuring of the surface for a wide range of applications (photocatalytic, photoelectrochemical, batteries, biomedical, etc.). This is due to its controlled nanoscale morphology, high aspect ratio self-organized nanostructures, facile use, and good integrity of layers grown directly on the substrate [34]. Figure 1 demonstrates the growth in research and applications based on anodic titanium dioxide (TiO\textsubscript{2}) layers encountered in the last 20 years.

![Figure 1.](image)

**Figure 1.** Publication record (year, publication number) in the 2000–August 2021 timeframe (Scopus–Elsevier) considering TiO\textsubscript{2} compact and anodic TiO\textsubscript{2} nanotubes or nanostructures.

Nanostructures grown under various electrochemical anodization conditions can be further modified by adjusting the crystallinity (amorphous, annealed to anatase, or anatase/rutile) or modified with active molecules for targeted applications [35–39].
Surface modification of Ti or Ti alloys provides several aspects critically affecting the successful osseointegration and bone regeneration, e.g., the response of the neighboring cells to the implant (migration, adhesion, proliferation, and mineralization of cells), a foreign body reaction/osteoclastic activity, and antibacterial activity. Various methods to modify micro and/or nano-structural topography have been developed to influence cell adhesion and proliferation, [40] and osteogenic activities [41,42].

The key benefits of anodic nanotubular structures are the nanotopographical advantages that are useful in osseointegration, as well as drug delivery applications due to the high surface area of nanostructures [7,9,34]. Recently, surface modifications based on electrochemical anodization followed by further decoration by active molecules or drugs were widely introduced for improving and tailoring osseointegration accompanied with a minimal immune response [20,35,43].

For the present review, we will confine our discussion only to nanotubular structures obtained via electrochemical anodization of Ti or its alloys and their uses for drug delivery applications tailoring osteoinduction, though several remarkable nanostructure technologies for biomedical application besides anodization have been developed. The present review aims to be a clear overview of the state-of-the-art of anodic TiO$_2$ nanotubes and the control over their formation. Next, the properties of nanotubes will be discussed, in particular in terms of biomedical applications, including the influence of micro- and nanoscale topography, surface roughness, and wetting behavior. We further introduce cell responses to such nanotubular substrates including growth and differentiation of bone-related cell types (mesenchymal stem cells, osteoblasts, and osteoclasts) on anodic nanotubes and electric-field induced osteogenic differentiation on TiO$_2$ nanotubes. In the last section, we present recent trends of TiO$_2$ nanotubes modification in drug delivery targeting osteoinduction via several application strategies to maximize the loading of anodic TiO$_2$ nanotubes and to prolong the release rate or induce beneficial biological effects.

2. Anodic TiO$_2$ Nanotubes

2.1. Nanomorphology and Critical Aspects of Anodic TiO$_2$ Nanotubes

2.1.1. Electrochemical Anodization

The shift of the materials from micro- to nanoscale topography has resulted in a large number of published works dealing with the nanostructuring of biomedical metals or alloys. In particular, nanostructuring of Ti and Ti alloys via electrochemical anodization was extensively explored. The principle of anodization and the resulting self-organization is quite simple, i.e., under optimal conditions, a steady state is achieved between oxide formation and chemical dissolution, while this balance can be influenced by several factors including anodizing potential, temperature, electrolyte composition, to name but a few [34].

Briefly, electrochemical anodization of Ti and its alloys is performed in an electrochemical cell, with the metal/alloy of interest used as an anode and a metal counter electrode such as Pt used as a cathode, as shown in Figure 2a. If a high enough voltage is applied and without the presence of fluoride ions in the electrolyte, the metal (M, usually valve metals such as Ti, Ta, Zr, Nb, etc., and their alloys) is oxidized (Equation (1)) and can than undergo the formation of a metal oxide (MO$_{x/2}$, Equation (2)), or is solvated followed by its dissolution in the electrolyte (M$^{x+}_{solv}$, Equation (3)) [34,44]. At the same time, at the cathode hydrogen gas is produced (Equation (4)) [34].

\[
M \rightarrow M^{x+} + xe^- \tag{1}
\]
\[
M^{x+} + \frac{x}{2} H_2O \rightarrow MO_{x/2} + xH^+ + xe^- \tag{2}
\]
\[
M^{x+} + \text{solv} \rightarrow M^{x+}_{solv} \tag{3}
\]
\[
xH^+ + xe^- \rightarrow \frac{x}{2} H_2 \tag{4}
\]
It should be noted that thermodynamic aspects such as oxide stability and solubility, as well as the corresponding reaction rates determine the equilibrium between oxide formation and dissolution (namely between Equations (2) and (3)). However, in systems where the oxide is not soluble and no additional side reactions take place, the oxide formation dominates, leading to very high efficiencies in the formation of the oxide [34,44]. Nevertheless, an equilibrium between the film formation and dissolution can be obtained, if there is some solubility of the oxide, which will result in a steady-state situation. In this respect, crucial is that, in the presence of fluoride ions, Ti\(^{4+}\) ions can be solvatized by forming fluoro-complexes (either by chemical dissolution of the oxide—Equation (5)—or by direct complexation of the metal cations at the oxide/electrolyte interface—Equation (6)) [34,44].

\[
MO_2 + 6F^- + H^+ \rightarrow [MF_6]^{2-} + 2H_2O \quad (M = Ti) \tag{5}
\]

\[
M^{4+} + 6F^- \rightarrow [MF_6]^{2-} \quad (M = Ti) \tag{6}
\]

When there is a competition between oxide formation and solvatization, where the latter is aided by the presence of fluoride ions in the case of titanium dioxide, the obtained steady-state condition results in the growth of a porous oxide; moreover, if there are optimum formation and dissolution rates, pores or nanotubes are formed [34,44]. The concentration of fluoride ions is key, as if too high, no oxide formation is observed (the controlling step of the reaction is the diffusion of the [TiF_6]^{2-} complex to the surface, as the complex formation is very fast), or if to low, a stable compact oxide layer is formed [34,44]. Porous or tubular layers are obtained in an intermediate fluoride ion concentration range, where the oxide formation/Ti\(^{4+}\) solvatization competition can be easily observed. For example, in the nanotube formation process, three stages are usually present: (i) the initial state, when a compact oxide formation; (ii) the second stage, nanoscale pores are irregularly formed and penetrate the compact oxide layer formed in (i); (iii) the third stage, when regular nanotubes formation occurs. A simplified schematic of the tube growth is also shown in Figure 2b, following the above-mentioned stages, where initially a compact oxide layer is formed, followed by irregular nanopores development which grow into self-organizing nanotubular structures finally reaching a steady-state formation rate. Typical of the anodic nanopores or nanotube layer is also the formation of the fluoride-rich layer throughout the whole nanotube length (as indicated also in Figure 2b), due to the fast migration of fluoride ions through the growing oxide layer.

Figure 2. Schematic of the electrochemical anodization process: (a) the electrochemical anodization setup and possible resulting nanostructures, (b) formation of the nanotubes (red—fluoride-rich layer).
For additional details related to more specific mechanistic aspects and more experimental investigations regarding the formation of such specific tubular shapes, the reader is referred to the following references [34,44,45].

The nanostructure morphology depends on the anodization conditions and the electrolytes, while nanostructures such as nanopores (with no individual separation in between the pores) and nanotubes (distinct separation of the cell boundaries into individual tubes) were obtained [34,44]. Anodic nanochannels or mesoporous structures can also be obtained when anodization is performed in hot glycerol phosphate electrolytes [16,46,47]. In the following section, we will discuss morphological aspects of anodic nanotubes and how to control the tube morphology as a function of the anodization parameters.

2.1.2. Morphology Aspects of Anodic TiO$_2$ Nanotubes

The different porous and tubular morphologies that can be obtained by electrochemical anodization of Ti are summarized in Figure 3. These include nanotubes obtained in glycerol: water (H$_2$O) and ammonium fluoride (NH$_4$F) electrolytes with a higher water content (30–50 vol.%), which show a uniform morphology (Figure 3a). The length, however, is limited to around 2 µm (see for example ref. [17,48,49]). In particular, when organic-based electrolytes with a lower amount of water (e.g., in between 1–20%) were used, nanostructures with much high aspect ratios were obtained (up to tens of µm for the lower water content electrolytes) [50,51] and can reach even faster growth rates by the use of additives (such as lactic acid) [52,53] or a more uniform morphology by surface pre-treatments (electropolishing [54], deposition of a photoresist [51]).

![Figure 3. Scanning Electron Microscopy (SEM) images of different TiO$_2$ nanostructures that can be grown by electrochemical anodization: (a) nanotubes grown in Glycerol:H$_2$O (60:40 vol.%) + NH$_4$F (reprinted with permission from ref. [20]. Copyright 2014 Elsevier), (b) open-top 100 nm diameter nanotubes grown in ethylene glycole (EG) with a low amount of water (4 M) + HF (reprinted with permission from ref. [55] Copyright 2016 Elsevier), (c) nanopores grown in EG with the low amount of water (6 M) + HF, (reprinted with permission from ref. [55] Copyright 2016 Elsevier), (d) nanotubes grown in EG, a lower amount of water (2%) + NH$_4$F in a 2 step anodization approach showing a typical initiation layer, (e) nanotubes in o-H$_3$PO$_4$ + NH$_4$F (reprinted with permission from ref. [56]. Copyright 2020 American Chemical Society), and (f) spaced nanotubes grown in diethylene glycole (DEG), low amount of water (1 wt.%) + HF (reprinted with permission from ref. [57]. Copyright 2016 Elsevier).](image-url)

An open-top morphology can be obtained under specific conditions, i.e., when hydrofluoric acid (HF) is used as the fluoride source [55] (see Figure 3b), or if an optimal mild ultrasonication is performed—either in water [9], ethanol/isopropanol [58], or in water with a very low amount of HF [59]). However, tubes are commonly covered by an initiation layer, either more compact or nanograss (when using NH$_4$F as fluoride source, or
a double anodization approach) [60]—see also Figure 3d showing a typical initiation layer for a double anodization approach. It was previously shown that for nanotube formation in low water content electrolytes (up to 0.5 vol.% water), the nanotubes are formed from an ordered porous oxide by following a pore-wall-splitting mechanism [61], i.e., by the selective dissolution of the fluoride-rich layer that exists at the hexagonal cell boundary of the nanoporous oxide layer. This can be significantly influenced by the water content in the electrolyte as well as the fluoride amount [61,62]. See also a typical example of nanopores in Figure 3c (EG + 0.2 M HF, 1 h at 10 V) [63].

Very ordered, regular, and highly-uniform TiO$_2$ nanotubes (initially referred to as nanostumps) can be obtained in concentrated H$_3$PO$_4$/NH$_4$F electrolytes at high temperatures (100 °C) [64,65], while the intertube distance decreases by increasing the temperature in the 75–140 °C range [56]—such a nanotubular structure is shown in Figure 3e. More interestingly, a nanotubular morphology with a two-scale organization and a distinct space in between each nanotube from the top to the bottom of the tube length can be obtained in certain organic electrolytes, such as diethylene glycol (DEG) [57,66], triethylene glycol [67], or dimethyl sulfoxide [66,68]. As shown in Figure 3f, such nanostructures (e.g., grown in DEG-based electrolytes) are highly uniform and provide additional functionality due to the spaced morphology.

The key factors to control the morphology of the nanostructures are well-established in the literature and can briefly be summarized as follows: the inner diameter is directly influenced by the applied voltage and the tube length by the anodization time, and both are strongly influenced by the electrolyte composition (water content and fluoride amount) [34,44,69]. A very good overview of the influence of water content on the nanostructure morphology (i.e., nanopore, nanotube, or sponge, grown in EG-based electrolytes) is shown by Albu et al. [69]—see Figure 4 for the regions of existence and representative SEM (Scanning Electron Microscopy) images of the nanopore and sponge morphologies.

![Figure 4](image-url)

**Figure 4.** (a) Voltage–water concentration diagram showing the different growing regions for nanostructures (TiO$_2$ nanopores, nanotubes, or sponge) for ethylene glycol-based electrolytes (reprinted with permission from ref. [69]. Copyright 2010 John Wiley and Sons). Representative SEM top and cross-section images for (b,c) nanopores—low voltage, low water content electrolyte, (d,e) sponge–high water content (b–e: reprinted with permission from ref. [69]. Copyright 2010 John Wiley and Sons).

Namely, if the applied voltage becomes too high, tube breakdown occurs and this for both a lower (<13 wt.%) or higher content of water, but in the case of the latter it leads to the formation of a sponge-like morphology. The water content influences the potential range into which nanopores can be grown [69]. For instance, anodization in an EG-based electrolyte with HF at 10 V, in an electrolyte with a 2–6 M H$_2$O content will result in a nanopore morphology, similar to the nanopore morphology in Figure 3c. However, for
the higher H₂O content of 6 M, the pore-wall-splitting transition is shifted to much lower
anodization times, e.g., of 1 h from 2.5 h, compared to lower water content electrolytes [63].

Nowadays, to take full advantage of the high aspect ratio of anodic nanostructures,
anodic nanotubular structures obtained predominantly in organic-based electrolytes are
used. As a result, the “V-shape” of the nanotubes, more pronounced for nanotubes grown
in organic electrolytes such as ethylene glycol (EG), has to be considered for applications
where the available surface area is crucial. This shape means that the inner diameter is
larger at the top and smaller at the bottom (as shown in Figure 5a–c), due to the chemical
etching occurring at the top of the nanotubes. The V-shape is easier to view for larger
diameter nanotubes with a high aspect ratio, but it is also present for nanotubes with
diameters of 15 nm and a length of 0.37 μm. Similarly, a characteristic of the nanotubes
grown in most organic-based fluoride-containing electrolytes is the double-wall structure,
with an outer shell (OS) and a rich carbon inner shell (IS) [34]—as shown in Figure 5d.
The latter is more easily observed after the annealing of the nanotubular layers. For many
applications, the IS can be removed by decorating (i.e., a chemical etching treatment that leads
to the removal of the C-rich inner layer) [70], or by using specifically designed anodization
conditions where only a single-wall nanotube structure is obtained (e.g., mixed EG and
dimethylsulfoxide electrolytes DMSO [71] or in DMSO electrolytes [68]).

![Figure 5. Schematic of the V-shape of nanotubes (b) and corresponding SEM images for nanotubes (D—diameter, L—length) with (a) 100 nm D nanotubes with 7 μm L (reprinted with permission from [55] Copyright 2016 Elsevier), and (c) 15 nm D nanotubes with 0.37 μm L (reprinted with permission from [55] Copyright 2016 Elsevier). (d) Separation into an inner shell (IS) and outer shell (OS) visible by annealing and a representative SEM image showing the double-wall structure.](image-url)

More importantly, from the various Ti alloys used as dental or orthopedic alloys, such as Ti6Al4V, Ti6Al7Nb, Ti13Cu4.5Ni, Ti25Pd5Cr, Ti20Cr0.2Si, Ti13Nb13Zr, Ti12Mo6Zr, TiMo alloys, Ti22Nb2Cr, TiZr alloy, etc. [72,73], only some can be anodized successfully to a uniform nanostructure by electrochemical anodization. For instance, anodic nanostructures in the form of nanotubes can be grown on Ti6Al4V [74,75], Ti6Al7Nb [74,76], TiZr alloys with Zr amount in the 5–50 wt.% [77,78], Ti24Zr10Nb2Sn [79], Ti13Zr13Nb [80,81], Ti28Zr8Nb [82], TiMo alloys (6–7 wt.% Mo [83,84], 15 wt.% [84]), TiNb alloys [85,86]. Additionally, specifically designed alloys can also be anodized, such as TiNbZr/Hf (Ti25Nb3Zr and Ti25Nb3Hf with x = 0.7 and 15 wt.% alloying element [87], Ti29Nb3Zr with x = 3, 15 wt.%
Zr [88,89], Ti35Nb5Zr with x = 3–10 wt.% Zr [90], Ti35Nb5Ta7Zr [91], Ti24Nb4Zr8Sn [92], Ti3xNb2Ag2Pt with x = 10, 30 and 50 wt.% [93], TiTa alloys [94], and other ternary alloys as Ti30TaxZr (with x = 3, 15 wt.% Zr) [89] or NiTi shape memory alloy [95,96]. Depending on the amount and type of alloying element, the anodization conditions or the tube growth range can differ to some extent in comparison to Ti anodized in the same electrolyte and under similar conditions. For more details, the reader is referred to the above-mentioned references and the dedicated reviews for anodization of biomedical alloys [97–100].

2.2. Key Properties and Their Improvement for Biomedical Applications

Up to this point, the different morphology of TiO$_2$ nanotube layers obtained on Ti and Ti alloys by electrochemical anodization have been introduced, in the next section, an overview of the crystallinity, surface roughness, corrosion protection, and wettability will be discussed. Properties such as topography, nanoscale roughness, chemical composition, wettability, and surface charge distribution, are key in influencing and controlling the interaction of the biomaterial’s surface with osteogenic cells and early bone response enabling the implant integration, decreasing the inflammatory response, as well as the bacterial adhesion or the foreign body response [1,10,22,101,102].

2.2.1. Crystallinity of Anodic TiO$_2$ Nanotubes

As-anodized TiO$_2$ nanotubes are amorphous; hence, to improve their properties and functionality, crystallinity is induced by thermal annealing treatments performed generally in air, at temperatures in the range of 250–750 °C, which convert the layers to an anatase or anatase/rutile crystalline structure [34,103–105]. However, the anatase phase starts to appear from temperatures as low as 250 °C and rutile from temperatures of around 450–500 °C (to note that the rutile growth starts from the Ti metallic interface) [34,105]. It was demonstrated that annealing TiO$_2$ nanotube membranes, which were removed from the Ti substrate, resulted in an anatase crystalline phase only, at temperatures of up to 950 °C [53].

The annealing time has a key influence on the crystallinity as longer annealing time at lower temperatures also results in crystallization to anatase [103]. There is a large number of works evaluating the crystallinity of TiO$_2$ nanotubes, achieved by annealing in air. In addition, further evaluation of the morphological changes occurring in the nanotubular structure due to the appearance of rutile (due to sintering) or due to morphological aspects of the double-wall structure of the nanotubes are discussed—see comprehensive reviews [34,106] and experimental works [53,103–105,107–109]. It should be noted, that in the case of mesosponge or mesoporous nanostructures obtained by anodization in hot-glycerol electrolytes the obtained nanostructures are already partially crystalline [47,110].

As in the case of biomedical applications targeting osseointegration, the nanostructures have to be grown directly on the surface of the implant, a clear example of the X-ray diffraction (XRD) patterns of 1 µm long TiO$_2$ nanotubes with 80 nm diameter annealed at temperatures up to 750 °C is shown in Figure 6. No significant differences were reported in the tube morphology for annealing up to 450 °C [111], but for higher temperatures of 550–750 °C, an increase in the rutile thermal oxide layer starting at the oxide/metal interface was observed (see also the cross-sectional SEM images in Figure 6b,c showing the thermal rutile layer). More so, for thermal treatments at temperatures higher than 650 °C, the crystallization process was accompanied by sintering (collapse) of the nanotubular layer [111].
2.2.1. Crystallinity of Anodic TiO$_2$ Nanotubes

As-anodized TiO$_2$ nanotubes...reprinted with permission from ref. [111]. Copyright 2016 Elsevier).

![Figure 6](image)

Figure 6. $\approx 1$ µm long TiO$_2$ nanotubes annealed for 2 h at temperatures in the 350–750 °C range: (a) X-Ray diffraction (XRD) patterns and selected SEM images showing the rutile thermal oxide layer growth (reprinted with permission from ref. [111]. Copyright 2016 Elsevier), and (b,c) nanotube morphology as a function of the annealing treatment, i.e., 550 °C and 650 °C, respectively (reprinted with permission from ref. [111]. Copyright 2016 Elsevier).

It is worth mentioning that after thermal crystallization of the TiO$_2$ nanotubes into anatase or anatase/rutile mixtures, a significant loss of water, fluoride, or carbon compounds frequently present in the nanostructures occurs (carbon comes from the inner shell of the classic double-wall nanotubes grown usually in organic electrolytes such as ethylene glycol), which also influences their wetting characteristics [34,66,71].

2.2.2. Surface Roughness of Anodic TiO$_2$ Nanotubes

A general trait of biomaterials is that biologically active implant materials possess increased surface roughness, which is one of the vital characteristics improving cell response to the implantable materials. A correlation between the surface roughness and osteoblast cell attachment was demonstrated [22,42], together with an effect on other aspects of a successful implant osteointegration (e.g., a selective protein adhesion, chondrocyte maturation) [112]. Nevertheless, though in vivo studies have shown that surfaces with a microstructure enable an improved contact between surface and bone and after implantation, increased mechanical retention [113,114], overall, more information is required, particularly for the recent developments in nanostructured surfaces [115].

The roughness of the biomaterial also influences additional properties such as higher local electrostatic charge density on the surface, [101] and in a combination with hydrophilicity can result in improvements in the early stages of bone healing or osteoporosis [116–118].

Figure 7a,b shows typical AFM (Atomic Force Microscopy) 2D topography and corresponding SEM images of NTs obtained by anodization of Ti in HF-containing aqueous electrolyte at an applied potential of 20 V for 20 min [119]. Similar NTs and AFM results were also obtained in 1 M sodium dihydrogen phosphate (NaH$_2$PO$_4$) + 0.3 wt.% HF, 20 V, 2 h [120], or as listed in Figure 7c,d for NTs grown on TiZr alloy by an optimized double anodization approach in a glycerol-based electrolyte with 15 vol.% H$_2$O and 0.2 M NH$_4$F (75 V, 1 h in the 2nd step) [121]. An overview of the average roughness (Ra) values, which is arithmetic means of the deviation in height from the profile mean value, encountered in literature for NTs grown on Ti or Ti alloys, are compiled in Figure 7e. While Ti has reported Ra values in the range of 10.7–22.7 nm, NTs have roughness values in the range of 17–112 nm depending on morphology, i.e., diameter size, where usually higher applied voltages lead to a certain point to higher diameters [78,119–126], or, to a certain extent,
to the annealing treatments (a slight increase in roughness is observed, with increasing the annealing temperature and the appearance of the rutile crystalline phase) [127]. In addition, also the inner diameter value is included in this overview where available (either not mentioned in the respective article or values were in a wide range of values due to the initiation layer covering the nanotubes). Peng et al. [128] has reported much higher Ra values for both Ti and TiO$_2$ NTs (i.e., 1.02 ± 0.06 µm for Ti, and 0.95 ± 0.02 µm for NTs). Based on the data presented above, there can be no conclusion concerning a possible correlation between tube diameter and average roughness; however, there is some variance in the values due to tube morphology, obtaining conditions, and top morphology of the NTs (open top, initiation layer, ultrasonication performed to remove the nanograss, etc.).

Figure 7. Top-view SEM image of nanotubes and corresponding 2D AFM (Atomic Force Microscopy) topography image for nanotubes grown (a,b) in an aqueous electrolyte on Ti foil (reprinted with permission from ref. [119]. Copyright 2010 Elsevier), or (c,d) in an organic electrolyte on TiZr alloy (reprinted with permission from ref. [121]. Copyright 2019 Elsevier). (e) Overview of Ra values on Ti, TiO$_2$ nanotubes or nanotubes on TiZr alloy (Ra and diameter values from refs. [78,120–127]; *—value of Rq).

2.2.3. Wetting Characteristics of Anodic TiO$_2$ Nanotubes

As mentioned previously, the early bone response is influenced greatly by surface properties (e.g., topography, roughness, chemical composition, and surface energy) [1,10,22,101,102], which in turn influences the wettability of solid surfaces. Wetting is the ability of a liquid to maintain contact with a solid surface, resulting from their intermolecular interactions, while the degree of wetting is determined by a force balance between adhesive and cohesive forces called the contact angle (CA) [129]. As the tendency of a drop to spread out over a flat solid surface increases, the CA decreases. A CA less than 90°, i.e., hydrophilic surfaces, usually indicates that wetting of the surface is favorable, and the fluid will spread over a large area of the surface. In contrast, CAS greater than 90°, i.e., hydrophobic surfaces, indicating that the wetting of the surface is unfavorable, and the fluid will minimize its contact with the surface by forming a liquid droplet. There are also two extreme wettability ranges, in which the CA either exceeds 150°, namely superhydrophobic state (SHS), or goes below 10°, namely superhydrophilic state. Such extreme cases can be only achieved on rough surfaces [130]. Besides the general studies on hydrophilic/hydrophobic surfaces, there are both fundamental and practical interests to extend the investigation of the interaction between proteins or cells and surfaces to the extreme wettability ranges.

Since biomaterials used in medical devices are intended to come into intimate contact with living cells and biological fluids, their surface wettability should be prospectively designed [131]. Firstly, the coating of the implant materials with proteins from blood and interstitial fluids will occur [10,132,133]. Therefore, the regulation of wettability and protein
adsorption on implanted surfaces is a key aspect in the field of regenerative biomedicine and tissue engineering.

Immediately after implantation, water molecules bind to the surface forming a water mono- or bilayer. Here, their arrangement depends on the implant surface properties at the atomic level accompanied by hydrated ions, such as Cl\(^-\), Na\(^+\), and Ca\(^{2+}\), followed by blood and tissue-specific proteins that adsorb/desorb on the surface [134]. It is common knowledge nowadays that the adsorption of proteins, as well as cell adhesion on implantable surfaces, depend strongly on its structure and topography. In this sense, the electrochemical anodization of implantable Ti substrates induces the following surface modification: (i) it forms a layer of TiO\(_2\) material, which is intrinsically hydrophilic, and (ii) introduces micro and/or nano roughness to strengthen the hydrophilicity of the implantable surface [9,111,135,136]—a typical example of contact angles for Ti, as-grown NTs and NTs annealed at different temperature is shown in Figure 8a [111]. Here, the fine-tuning of surface wettability can be achieved by either development of particular surface roughness or surface modification with low-energy materials. This is because one step is normally required to alter the hydrophilic surface to a hydrophobic one, i.e., by the deposition of low surface energy material (as exemplified in Figure 8b,c, whereby the deposition of polydimethylsiloxane, PDMS, using an ultraviolet (UV) light grafting technique turns hydrophilic TiO\(_2\) NTs into hydrophobic [136]). The latter was obtained by various surface modification techniques as discussed in recent reviews or the literature [136–138].

Figure 8. (a) Typical contact angle values for Ti foil, as-grown nanotubes (NTs), and NTs annealed at different temperatures in the temperature range from 350 to 750 °C (data taken from reference [111]). Typical contact angle for (b) bare highly ordered TiO\(_2\) NTs and (c) coated by polydimethylsiloxane (PDMS) molecules using ultraviolet (UV) light-grafting technique.

2.2.4. Corrosion Resistance of Anodic TiO\(_2\) Nanotubes

Concerning the biocompatibility of implant biomaterials, in vivo corrosion resistance has a crucial contribution to the implant’s lifetime, and for metallic biomaterials (excluding the biodegradable ones) this is “the more corrosion resistant, the more biocompatible” [139]. This, as physiological solutions, are viewed as extremely corrosive towards metallic biomaterials and if corrosion occurs it can lead to diminishing biocompatibility, i.e., a release of metallic ions or particles as a result of corrosion can lead to inflammation and, finally, to
tissue loss. Though usually, compared to compact biomaterials, porous materials are more prone to corrosion [140], in the case of nanostructures, both the morphology and the metal oxide itself (pure Ti or alloys) contribute to the corrosion performance [93].

Investigations into the corrosion performance of amorphous/annealed NT structures, either grown on Ti [111,141–149] or its alloys (Ti6Al7Nb [76,150,151], TiZr [152]) indicated that for all NT layers the electrochemical parameters show very stable characteristics—as shown in Figure 9 for various NT structures. E.g., in the well-established Fusayama artificial saliva solution (consider also that the NTs had a corrosion current density, \( j_{\text{corr}} \) of 0.12 \( \mu \text{A cm}^{-2} \), while bare Ti had 7.12 \( \mu \text{A cm}^{-2} \)) [141], another artificial saliva (showing the influence of tube morphology) [142], and Hank’s solution (evaluating the influence of the annealing treatment) [111]. For annealed NTs, lower corrosion current density (at least one order of magnitude) was recorded compared to the amorphous NTs [111,147], and this could also be linked to the appearance of rutile at the NT/metal interface. For instance, in Hank’s solution Ti is reported with a corrosion current density of 1.29 \( \mu \text{A cm}^{-2} \) [153] to 2.3 \( \mu \text{A cm}^{-2} \) [154] (and 0.116 \( \mu \text{A cm}^{-2} \) for smooth Ti [144]), while for the annealed NTs this is in the range of 2.51 to 0.07 \( \mu \text{A cm}^{-2} \) decreasing with annealing temperature up to 750 °C (see also the Tafel plots in Figure 9c) [111], or 0.014 \( \mu \text{A cm}^{-2} \) for anatase NTs [144]. Overall, a similar trend is maintained for the corrosion current values in the literature for the various saliva or solutions used, namely with annealing the corrosion protection of the nanotube layers increases. To note that (i) linear polarization measurements/Tafel plots are usually measured when steady-state conditions are achieved at the interface between metal/solution (e.g., the change of the corrosion potential is less than 5 mV in 10 min) [155], and (ii) at least one of the branches has to exhibit linearity on a semi-logarithmic scale over at least one decade of current density to allow the extrapolation of the Tafel slope [156–158] (see also the Tafel plots in Figure 9c showing linearity only in the cathodic region).

![Figure 9](image_url)

*Figure 9.* Typical Tafel plot for nanostructures on Ti measured in (a) Fusayama solution for TiO\(_2\) nanotubes (reprinted with permission from ref. [141]. Copyright 2010 Elsevier), (b) in artificial saliva for mechanically polished Ti (MPT) and TiO\(_2\) nanotubes obtained with different morphology as a result of anodization at different voltages (5, 10, 15, and 20 V) (reprinted with permission from ref. [142]. Copyright 2011 Elsevier, (c) in Hank’s solution for as-formed TiO\(_2\) nanotubes and annealed in air at temperatures of 350, 450, 550, 650, and 750 °C (reprinted with permission from ref. [111]. Copyright 2016 Elsevier).

For NT structures, the tube diameter and the thickness of the barrier layer are also crucial, significantly affecting their electrochemical corrosion performance [142,145,148,159]. The linear polarization curves in Figure 9b demonstrate that the passivation currents of the different morphology NTs, i.e., increasing diameter and tube length with increasing the applied voltage, were mostly lower than for the bare Ti. It is worth mentioning that the morphology type, pore diameter, and thickness of the barrier layer are controlled by the anodization conditions [34]. Concerning morphology, NTs grown on NiTi alloys lead to lower corrosion resistance and more Ni release compared to the bare NiTi alloys, due to the increased surface area [95]. Whereas a 5.3 \( \mu \text{m} \) thick nanoporous layers (93 nm pore diameter) led to an improvement of the corrosion resistance [160]. In general, for NTs
grown on alloys, the alloying elements and their amounts contribute substantially to the electrochemical stability of NTs [89,152]. For example, as in the case of Ti30NbXZr or Ti30TaxZr, where the corrosion resistance was improved with the increase in the Zr content from 3 to 15 wt.% [89].

Similar performance, i.e., resulting in an improvement of the corrosion resistance in artificial saliva by the formation of NT structures (amorphous), on either ultrafine-grained or coarse-grained Ti13Nb13Zr alloys was observed [161]. Saji et al. [91] reported that in the case of nanostructures on Ti35Nb5Ta7Zr alloy, namely nanopores or nanotubes with diameters of 30–50 nm or 30–80 nm, respectively, and annealed to anatase in argon atmosphere (550 °C), nanopores showed a corrosion current density of 0.76 µA cm⁻², lower than the 3.12 µA cm⁻² obtained for NTs (in aerated Ringer’s solution). This difference is due to the distinctly separated barrier oxide/tube bottom interface. Furthermore, also for a nanochannel morphology, a decrease in the corrosion current density was observed compared to compact oxide (e.g., for nanochannels on TiZr alloy that show partially crystalline structure due to the anodization in hot-glycerol phosphate electrolytes) [16].

3. Tailoring Osteoinduction with Anodic TiO₂ Nanotubes

3.1. Advantages of Anodic TiO₂ Nanotubes for Osteoinduction

3.1.1. Nanotopographical Cues of Anodic TiO₂ Nanotubes

Most known mammalian cells exhibit the instinct to adhere onto a surface to carry out normal metabolism, proliferation, and differentiation [10,162,163]. Cells anchor onto a surface and sense the extra-environment through ion channels and receptors presenting at their membranes, then integrate the chemical and physical signals from the extra-environment and give the response that some transmembrane receptors form clusters known as integrins, and associate intracellularly with groups of proteins which link them to the cytoskeleton. Subsequently, focal adhesions take place through the binding between the cluster integrin receptor and the ligand of the extracellular matrix (ECM) [133,163,164]. The cells will undergo apoptosis if are not able to synthesize and deposit their own ECM molecules in a short time [164,165].

Nowadays, much more attention has been paid to nanoscale microenvironments surrounding cells, due to the highly sensitive cell responses including cell adhesions and cell decisions for cell growth and differentiation to nanotopographic cues. Various techniques on different biomaterial systems including TiO₂ nanostructures have been introduced to achieve surface structures in the sub-100 nm region for the investigation of cell-stimulating effects and biomimetic activation [166]. For example, geometrically defined, adhesive and stable surface protrusions were made based on polymer demixing [167], ordered gold cluster arrays [168,169], nanophase ceramics, biomedical alloys [42,170], or self-organized nanoporous aluminum surfaces. Among the biocompatible implant materials suitable for bone repair, titanium has been widely accepted as the most favorable for osteogenic differentiation in vitro and bone regeneration in vivo. One of the most clinically relevant advantages of anodic nanostructures for osteoinduction and osteogenesis is well-defined, self-organized, nanoscale topography coatings on TiO₂ implant materials, which support the growth and osteogenic differentiation of mesenchymal stem cells (MSCs) in vitro and in vivo bone formation [7,171]. Moreover, TiO₂ substrates with well-defined nanoscale topography allowed highly reproducible cell behavior responding to the topographic cues from the oxide surface.

Fundamental findings reported more than a decade ago, demonstrated that the diameter of anodic TiO₂ NTs vitally influences the cell (mesenchymal stem cells—MSCs, osteoblasts, osteoclasts, etc. [34]) activities leading to increased cell adhesion and growth as compared to compact oxide or bare Ti [7,43,172]. These findings led to further research focused on diameter-controlled nanostructures for implants/tissue engineering and drug delivery design.

For the influence of NT diameter on the different cell type adhesion, proliferation, and differentiation, please see (i) the works of Brammer et al. [173] and Chamberlain
et al. [21] and (ii) the works of Park et al. [7,8,35,43] and Khaw et al. [174]. Brammer et al. and Chamberlain et al. showed that small diameter NTs (30 nm) promoted the highest degree of osteoblast activity and diameters of 70 nm resulted in the diminished inflammatory response, respectively [21,173]. Park et al. [7,8,35,43] proposed that 15 nm NTs in diameter is an optimal tube diameter for increased cell adhesion and proliferation for MSCs and endothelial cells. Further, Bauer et al. [175] confirmed the same trend for ZrO$_2$ NTs, indicating that the NT size effect dominates over the crystal and fluoride content [8]. In addition, Khaw et al. [174] showed that there is a discrepancy concerning the affinity of hMSCs and human osteoblasts (HOB) for TiO$_2$ NT of various pore diameters in the sense that hMSCs prefer NT with a 20 nm diameter in terms of osteogenic differentiation while 50 nm nanotubular surfaces potentiate osteoblastic maturation of HOBs.

At the cell-substrate contact area, nanotopography mainly affects the cell responses and following behaviors including cell decisions, for example, whether cell divides or starts to be differentiated. Particularly in the nanoscale cell-substrate contact area, the size of the lateral spacing can determine cell fate depending on whether the lateral gap size allows the integrin clustering on the NTs or not. Integrins have been known as one of the most important microenvironmental signal receptors in the mammalian cell system, and integrin clustering can be developed when integrins are activated by environmental signals out of extracellular matrix/substrates. The study of Park et al. [7] showed that the gap size of around 10–20 nm in NT diameter perfectly fits the integrin clusters leading to focal adhesion complexes and downstream signaling to the nucleus. Likewise, recent studies [176,177] have brought to focus the contribution to the decipherment of the molecular mechanisms of how integrin-mediated cell interaction with TiO$_2$ NTs may direct cell fate, i.e., cell adhesion and proliferation, cytoskeleton reorganization, motility, cell shape, and osteogenic differentiation.

These results led to the establishment of the osteogenic differentiation mechanism. First, smaller tube diameters of 10–20 nm provide the optimal length scale for integrin clustering and the formation of focal contacts on the nanotube surface which further result in higher cell proliferation, migration, and differentiation to osteogenic lineages of the MSCs compared to bare Ti implants or compact TiO$_2$ layers. Considering the estimated occupancy size of the head of the integrin heterodimer, the 15–20 diameter of the nanotubes result in the clustering of integrins in the closest possible packing which leads to an optimal integrin activation [7,43]. In contrast, larger tube diameters (e.g., >50 nm) impaired cell spreading, adhesion and even prevented integrin clustering and focal adhesion complex formation—we this led to a strikingly reduced cell behavior (proliferation, migration, differentiation) and in the end induced the adhesion-dependent form of apoptosis [7]. Secondly, on the small diameter nanotubes, phosphorylation of the focal adhesion kinase (FAK) and extracellular regulated kinase (ERK), target of the FAK signaling pathway was highest compared to on the much larger 100 nm diameter nanotubes [7].

Reports in literature confirm that different nanotopography cues (micro-scale surface modifications, nanopits, nanosheets, nanotubes, etc.) influences osteogenic differentiation via various downstream molecular pathways following integrin signaling, reorganization of the actin cytoskeleton, and nuclear translocation/transcription [10,178,179]. These include several canonical pathways such as FAK [7], ERK [7,179], phosphoinositide 3-kinase/protein kinase B (PI3K/AKT) [179], Rho-associated kinase (Rho-ROCK) [179,180], autophagy-mediated signaling between Yes-associated protein (YAP) and β-catenin [181], mammalian target of rapamycin complex with Rictor (mTORC2) [182], and Wnt/β-catenin [183] signaling pathways. Recently, Lv et al. [184] revealed the epigenetic mechanism of nanotube-guided osteogenic differentiation of MSCs and that changes in cell adhesion and cytoskeleton reorganization are linked with epigenetic alterations. The mechanism would be even more complex as the above-mentioned pathways can crosstalk and have a crucial role in the cell adhesion, migration, proliferation, and differentiation directed by the biomaterial surface.
Not only topography itself but also a minute mechanical strain on nanotube layers seems to affect the stem cells, resulting in their osteogenic differentiation [176,180,185–187]. This mechanical strain has been reported to promote osteogenic differentiation of MSCs on TiO$_2$ nanotubes via the FAK [185], ERK1/ERK2 [186], Rho-ROCK [180], Yes-associated protein/Tafazzin (YAP/TAZ) [187] FAK-ERK1/ERK2-RunX2 [176] pathways. The exact mechanism through which nanotopography modulates mechanotransduction for osteogenic differentiation of MSCs still remains to be further confirmed through additional investigations.

Recently, we compared the generally-grown (close-packed) NTs with spaced NTs, where both NTs have similar diameters of $\approx 80$ nm, but spaced NTs have a 80 nm tube-to-tube individual spacing. The spaced NT morphology did not show any detrimental effect on osteoblast functions in vitro, rather having a beneficial influence on the osteogenic differentiation of pre-osteoblasts [188], indicating that gap distances irrespective of inner or outer nanotube rims may affect cells that come into contact with these nanoscale gaps, further delivering contact signals to the nucleus to decide cell differentiation.

Many previous works demonstrated that various diameter TiO$_2$ NT structures can also induce the anti-inflammatory response of hosts (Chamberlain et al. [21], Smith et al. [189], Neacsu et al. [20], Yao et al. [190], Bai et al. [191]). The surface modification induced by the nanotube structures combined with annealing can change the hemocompatibility of TiO$_2$ NTs, by alleviating platelet activation (Mazare et al. [111,150], Junkar et al. [15], Huang et al. [192], Gong et al. [193], Bai et al. [191,194]). Several in vivo studies have shown that TiO$_2$ nanotubular layers could induce successful peri-implant bone formation/osseointegration in various animal models using different diameter NTs: von Wilmowsky et al. (30 nm diameter nanotubes) [195], Wang et al. (30, 70, and 100 nm diameter NTs) in minipigs [174], Alves-Rezende et al. ($\approx 74$ nm diameter NTs) [196], Baker et al. (TiO$_2$ NTs—in vitro and in vivo intramedullary fixation) in rats [197], or the review of Wang et al. [198] for the effect of TiO$_2$ NTs grown on the implants’ surface on osseointegration in animal models.

Another advantage of anodic nanostructures for osteoinduction lies in the potential of a synergistic combination of nanotopographic cues with drug delivery via NT surface or inner space decoration, as a cargo. By modifying the NTs with active molecules or growth factors (such as bone morphogenetic protein-2—BMP2, epidermal growth factor—EGF, osteogenic growth peptide—OGP), improved cell adhesion or differentiation can be obtained. For example, using a BMP2 decoration, Balasundaram et al. [199] showed an increased osteoblast adhesion on TiO$_2$ NTs, and Park et al. [35] reported accelerated differentiation of MSCs towards osteogenic or chondrogenic lineage on BMP2-coated NT layers. Lai et al. [200] further confirmed MSCs differentiation to osteoblasts on BMP2-decorated TiO$_2$ NTs (via polydopamine). Moreover, several growth factors/cytokines can be coupled with different functional layers on TiO$_2$ NTs surface [201]. For example, BMP6-loaded TiO$_2$ NTs can be coated with platelet-derived growth factor containing silk fibroin in a composite for increased osseointegration [202]. Several other modifications include EGF on NT structures for increased adhesion of MSCs, OGP functionalization of the NTs for an improved cell spreading and differentiation of osteoblasts [203], and calcitonin gene-related peptide (CGRP) on TiO$_2$ NTs for osteoporotic bone implants [125]. While the previous studies mostly focused on decorating growth factors or drugs on the top of NT surfaces, recent studies tend to expand drug delivery applications through the inner nanotube wall/space, as well as decorating the top surface of NTs. We will discuss this recent trend in NT drug delivery more in detail in one of the next sections.

3.1.2. Electric Field Stimulation of Anodic TiO$_2$ Nanotubes

In bone tissue engineering, a classic biological triad represents three critical components leading to successful bone regeneration using biomaterial implantation: (1) biomaterial/scaffolds-derived micro-, nanoenvironmental cues, (2) electrochemical, electrical, physical regulatory signals including bioreactive molecules, and (3) cells including
cell-cell contact and functional matrix signaling. Among these three major components, we already discussed in a former section the TiO$_2$ nanotube-derived nanoenvironmental cues affecting cell behaviors. We also shortly mentioned bioactive molecules on TiO$_2$ nanotubes, that stimulate osteogenic differentiation and osteoinduction. Considering the scope and limitation of the present review, discussion about cell components (related to point 3) including the interplay of different bone and precursor cells interacting via cell-cell contacts, is beyond the topic of the review. In this section, electrical stimulation among regulatory signals in the triad, especially on TiO$_2$ nanotubes for osteoinduction will be introduced and reviewed.

In the field of bone tissue engineering using biocompatible implants, cellular behavior and cell fate of stem cells have been known to be determined not only by microenvironmental signals, such as substrate topography, soluble growth factors and cytokines, and cell–cell and cell–extracellular matrix interactions, but also by electrochemical signals [164]. Electric fields have been applied in tissue engineering for many different purposes. The main applications of electric fields are in the characterization of artificial tissues and their component cells, and the formation of artificial tissue-like materials, either by assisting in the formation of the artificial extracellular matrix (e.g., the formation of scaffolds by electrospinning), or the micromanipulation of the cells themselves using electric fields [204]. Of further potential interest in tissue engineering are also the biological effects of the electric fields. In fact, the clinical application of an electric field (EF)/an electromagnetic field for bone regeneration has a long history. A meta-analysis of the data in the last 60 years shows clinical EF trials to have an overall favorable influence on bone healing [205,206]. However, such clinical trials have also considerable variations depending on the treatment regimen and study design, thus hampering a direct comparison and a critical evaluation of results [205]. Further, despite a long history of clinical trials, the mechanism linking electric stimuli and its sensing on the cell surface is still elusive. Therefore, during recent decades, the mechanism of EF (e.g., how cells sense and react to EFs) affecting cell adhesion, migration, proliferation, and differentiation has been extensively investigated in various primary cell/cell lines [206–211] including epithelial, mesenchymal, and neural cells [212–215]. Exposure of the cells to an electric current, if a certain current value threshold is exceeded, leads to cell death, while currents lower than 11 As/m$^2$ have shown no decrease in cell viability [216]. Asides from the overwhelming impact of the electric current value, there are additional aspects that can negatively influence the viability of the cells, such as local acidic pH [217], metal ions release from the electrodes or cell contact directly on the electrodes [216]. Commonly, in a physiologically tolerable EF strength (<5 V cm$^{-1}$), many of the different cell types can immediately react to EF stimuli, which include EF directional axis-dependent cell migration (galvanotaxis), neuronal activation and regular muscle cell alignments. These EF-corresponding specific cell responses have been suggested for use in wound healing, neurons stimulation, and tissue engineering on various scaffolds [206,218].

In the bone tissue engineering field, evidence has accumulated that clearly points out that electric (direct or alternating current) or electromagnetic fields can induce osteogenic differentiation of mesenchymal stem cells or osteoblasts [210,213,215]. Recent studies have shown that various scaffolds can be used in concert with electric or electromagnetic fields for bone cell stimulation, such as (i) collagen scaffolds (with piezoelectric properties) for human osteoblasts under magnetic and additional alternating electric field [219], (ii) non-piezoelectric three-dimensional matrix for osteoblasts under electromagnetic stimulation [220], (iii) polycaprolactone based scaffolds under alternating current electric fields for osteoblast-like MG63 cells [221], or (iv) conductive polypyrrole/polycaprolactone scaffolds under electrical stimulation [222].

Similarly, TiO$_2$ substrates can also be used to stimulate bone cell differentiation under an applied EF. Already in 2016, we have reported an efficient EF-induced osteogenic differentiation of MSCs grown on 15 nm diameter TiO$_2$ NT layers, without any osteogenic chemical supplements stimulating the differentiation [223]. MSCs discern the EFs and re-
spond to both x-y planar and z-axis EFs (Figure 10a) at smaller EF strength than 0.4 V cm\(^{-1}\), i.e., at a comparable current level of endogenous provoking current in the body. Interestingly, under EF stimulation MSCs grow fast as multilayer on TiO\(_2\) nanotube layers while without EF they remained in a monolayer (as shown in Figure 10b).

Figure 10. Effect of electric fields (EFs) on mesenchymal stem cells (MSCs) cultured on TiO\(_2\) nanotubes: (a) illustration of MSC under x-y planar (left), z-axis (right)-directed EF device, respectively (reprinted with permission from ref. [223]. Copyright 2016 Mary Ann Liebert, Inc.). (b) Osteogenic differentiation of green fluorescent protein (GFP) labeled MSCs under z-axis-directed EF (200 mV cm\(^{-1}\)) for 8 days. Bar: 200 mm (reprinted with permission from ref. [223]. Copyright 2016 Mary Ann Liebert, Inc.). (c) Gene expressions of osteocalcin and osterix following 3 days of x-y or z-axis directed EF application by quantitative Polymerase Chain Reaction, qPCR (N = 3) (reprinted with permission from ref. [223]. Copyright 2016 Mary Ann Liebert, Inc.). Reduced NTs show higher efficiency for EF stimulation of MSCs: (d) SEM images of reduced 100 nm diameter nanotubes, inset: high magnification image of the tube wall structure) (reprinted with permission from ref. [224]. Copyright 2019 Elsevier), (e) resistance values for as formed and reduced nanotubes obtained from solid-state conductivity measurements (data taken from ref. [224]), (f) schematic showing the EF-triggered calcium influx reprinted with (permission from ref. [224]. Copyright 2019 Elsevier), (g) the voltage-
dependence of the intracellular calcium elevation in as formed and reduced TiO$_2$ at the end of a
10 min-EF stimulation, data were obtained from 100 randomly chosen cells/experiment and three
independent experiments were performed for each group (reprinted with permission from ref. [224].
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The study further revealed that enhanced intracellular calcium signaling and the
spreading of the intracellular Ca$^{2+}$ to the adjacent cells under EF are the main mechanisms
of EF-induced osteogenic differentiation. It is worth mentioning that in bone differentiation
a z-axis EF using an MSC-plated TiO$_2$ substrate as an anodic electrode can induce a
comparable stimulating effect with x-y planar EF when MSCs were separated from EF
electrodes as shown in Figure 10c (a comparison of bone differentiation-specific marker
expressions). The finding indicates that MSCs can be plated or moved to adhere directly
on the EF-generating TiO$_2$ electrode, which might be useful for EF-engaged bone tissue
engineering.

Other works also supported the stimulation of bone cells on TiO$_2$ or TiO$_2$ composite
layers by EF, showing that electric field stimulation of osteoblast on anodic nanotubes leads
to an increase in their specific biomarkers [225], and more recently Sahm et al. [226] also
reported the influence of alternating electric fields on human osteoblasts growing on and
in the surrounding of Ti6Al4V electrodes.

Further, a shortened EF-engaging time and an effective but minimal EF strength are
critical issues in EF-applied clinical bone tissue engineering. We have reported signifi-
cantly shortened EF exposure time by improving the conductivity of the TiO$_2$ NTs [224].
“Black” TiO$_2$ NTs, subjected to an optimal reduction treatment in argon hydrogen (Ar/H$_2$)
environment can lead to a significant increase in tube conductivity and decrease of the
electron transport time, as a result of forming Ti$^{3+}$ and oxygen vacancies in the NTs (see
the morphology of reduced tubes in Figure 10d and conductivity values in Figure 10e).
Hence, much shorter EF-exposure time (from previous days to 10 min) and a decrease
in the applied EF strength (from 400 mV cm$^{-1}$ to 100 mV cm$^{-1}$) sufficiently enabled the
EF provocation of an early response of the stem cells (a schematic drawing of increasing
intracellular calcium influx under Z-axis EF in Figure 10f and a superior intracellular
calcium activation on reduced TiO$_2$ NTs compared to as formed NTs in Figure 10g).

Overall, in the past decade, great achievements were accomplished through many
valuable studies that verified the EF mechanism (how cells recognize the EF stimuli and
which signaling pathway dictates EF-induced bone differentiation). The new understand-
ing of the mechanism may allow the accomplishment of potential clinical EF application in
bone tissue engineering in the near future, following the careful optimization of EF parame-
ters. Electric/electromagnetic field application for bone regeneration using bio-implantable
materials could be a useful option when combined with the other novel technologies
including TiO$_2$ surface modifications and drug delivery systems which will be discussed
in the following section.

### 3.2. Drug Delivery Applications Based on Anodic TiO$_2$ Nanotubes

We have previously discussed the advantages of anodic TiO$_2$ nanotubes on osteoin-
duction and osteogenesis and their validity for a wide range of Ti alloy substrates. The
nanotubes’ high surface area and their distinct topography can provide their full advantage
in drug/active agents delivery applications [14,227,228]. Extensive studies are investi-
gating the release profiles of drugs, including antibiotics, peptides, metal ions (Ag, zinc -
Zn, copper - Cu), or various biopolymer coatings. Overall, the studies investigating the
mechanism and release rate from anodic TiO$_2$ nanotube layers confirm the higher loading
capabilities of the NTs (that are, as expected, linked with their high aspect ratio). Previous
studies have also revealed that vacuum impregnation techniques usually lead to a higher
elution time, compared to soaking techniques [229]. A variety of drug/active compound
release approaches based on anodic TiO$_2$ NT structures are summarized in Figure 11.
3.2.1. Release Rate from Anodic TiO$_2$ Nanotubes

In the first step, we will discuss key aspects concerning the release rate of drug/active components. Generally, the drug release rate is determined by the design of the delivering platform (varied from the incorporation of metal ions/nanoparticles on the surface to a direct incorporation into the TiO$_2$ NT space) and biochemical characteristics of drugs: see, for example, Ag [230,231], strontium - Sr$^{2+}$ [149], magnesium (Mg) and silicon (Si) [93], vancomycin [230], ibuprofen [127] gentamicin [127,232], doxorubicin [229,233], anti-inflammatory drugs (indomethacin [234], sodium naproxen [235]) co-delivery of drugs (gentamicin and ibuprofen) [236], or various active molecules (e.g., quercetin [237]). Usually, one of the main challenges in the design of drug delivery systems based on either TiO$_2$ NTs or other nanostructured materials is a controlled and sustained release of drug in contrast to a burst release that can rapidly lead to the accumulation of the active drug/active agent to the toxic levels at the targeted site [18]. Table 1 represents an overview of drug/active agents with specific loading and release strategies using anodic nanostructures.
Table 1. Overview of drug/active compounds loading and releasing from anodic nanotubes on Ti or Ti alloys (D—diameter, L—length).

| Drug/Compound Type | Drug/Agent Deposition | Release Rate | Reference |
|--------------------|-----------------------|--------------|-----------|
| Sr²⁺ | Sr²⁺—dip coating | A viable alternative in orthopedics to provide improved corrosion resistance and enhanced biocompatibility | [149] |
| Ag or Vancomycin (VAN) | Aqueous: (a) D 70 nm, L 0.87 µm (b) D 100 nm, L 1.45 µm. Organic: L 6.5 µm | VAN release was significantly retarded from NTs in organic electrolytes (compared to aqueous). Ag release was retarded from aqueous nanotubes compared to Ti surfaces. | [230] |
| Gentamicin (GEN) | Immersion of samples in phosphate-buffered saline solution containing GEN | Partly delayed release of gentamicin, for targeting bacterial inflammation around the implant. | [232] |
| Ibuprofen (IBU) or Gentamicin (GEN) | 10 wt.% solution of IBU in methanol, and 10 wt.% GEN in water. 5× of 1 mL pipetting and drying (drying in air, room temperature or 75 °C) | The release process is governed by the desorption of the drug from the top surface, followed by a combination of desorption and slow diffusion of the drug from the inside of the nanostructure. | [127] |
| Doxorubicin (DOX) | 15 cycles of deposition and drying under vacuum in DOX solution. Polyethylene glycol (PEG) layer for capping (1 to 50% PEG) | Release of doxorubicin can be controlled (slowed down) only during the first 3 h by the PEG layer. TiO₂ NTs are competitive for drug release of low polarity drugs compared to other boron or carbon-based materials. | [233] |
| Ibuprofen (IBU) and Gentamicin (GEN) | 10 wt.% solution of IBU in methanol, or GEN in water were prepared. 1 mL of each solution: a) IBU and GEN at the same time (de-noted IG), b) GEN and then IBU (GI), c) IBU and then GEN (I&G) | The length, crystallinity, and loading procedure of NTs influence the drug loading and release processes. Drug release can be modified by the loading procedure (GI approach led to the longest period or release time for GEN as the initial burst release was inhibited). | [236] |

Jarosz et al. [127] clarified the drug release kinetics and mechanism using a desorption-desorption-diffusion (DDD) model of the drug release. In their study for ibuprofen/gentamicin release from TiO₂ NTs, two different drug release kinetics were shown: drug desorption from the top of the nanostructure, and then desorption and diffusion of the drug from the inner nanostructure. A first-order kinetics is responsible for the initial fast drug delivery (from the surface of the nanostructure), while the slow release of the drug from within the nanostructure combined the first-order kinetics with a Higuchi model [236]. Further, the work of Pawlik et al. [236] has shown that release rate can be differentially controlled according to the characteristics of drug combinations. The co-delivery of ibuprofen and gentamicin, moderate water-insoluble and water-soluble, respectively, enables controlling the release time from TiO₂ NTs when the loading procedure...
consisted of first gentamicin and then ibuprofen serially, leading to a much slower release of gentamicin thus avoiding an initial overdose burst.

Coating methods can also contribute to prolong the drug release rate, for example (i) capping techniques, such as doxorubicin loaded in NTs with a polyethylene glycol layer coating as a barrier [233], chitosan coatings of NTs loaded with indomethacin [234], gentamicin [238], selenium [239], quercetin [237] or mixed coatings of gelatin/chitosan [240], (ii) deposition of the ions on top of polymer layers (for example, Ag on polydopamine decorated TiO\textsubscript{2} NTs [231]), or (iii) for encapsulation of the drugs in micelles and their loading into the NT structures (indomethacin [234]).

3.2.2. Drug Delivery for Antibacterial and Osteoinductive Activities

For drug delivery on TiO\textsubscript{2} nanotubular bio-implants, so far a high number of investigations has mainly focused on antibacterial effect using (i) metal ions or nanoparticles (Ag [241–243], silver oxide - Ag\textsubscript{2}O [244,245], gold - Au [122], selenium - Se [239], Zn [246], zinc oxide - ZnO [247], Cu [248], Sr [249,250], tantalum (Ta) and Cu [124], Zn-Ag [251], Sr-Ag [250], or calcium and phosphorus - Ca-P [252,253]), (ii) drugs (vancomycin [36], gentamicin [254], metformin [255], alendronate [256], simvastatin [257]) or (iii) polymers (polyaniline [258], chitosan [210,238]) to provide a contamination/infection-reduced condition beneficial for osseointegration (see Figure 11 and Table 2).

| Drug/Compound | Nanostructure | Drug/Compound Deposition | Biological Effects | Reference |
|---------------|---------------|--------------------------|-------------------|-----------|
| Sr\textsuperscript{2+}, Ag | TiO\textsubscript{2} NTs (anatase) | Sr\textsuperscript{2+}—hydrothermal treatment; Ag\textsuperscript{+} by photodeposition to Ag NPs | Enhancement of the osteobonding capability of the nanotubes, as well as of their antibacterial activities by combining the pro-osteogenic effects of Sr\textsuperscript{2+} and strong antibacterial effect of Ag NPs. | [250] |
| Ag\textsubscript{2}O | TiO\textsubscript{2} NTs (amorphous) | Ag\textsubscript{2}O nanoparticles are embedded into the nanotubes. Substrates are TiAg layers (magnetron sputtering) | Sustained antibacterial activity due to the controlled low dose Ag\textsuperscript{+} release, improved cell attachment and spreading, no deleterious effects on pre-osteoblast cell viability, proliferation, and differentiation. | [244] |
| Zn | TiO\textsubscript{2} NTs (30 nm: anatase, rutile; 80 nm: anatase) | Zn—deposition onto the NTs by hydrothermal treatment | Antibacterial effects depending on the amount of loaded and released Zn in NTs. 80 nm NTs (3 h Zn deposition) enhance MSC osteogenic differentiation (enhanced protein deposition, enabling cell functionalities and Zn release). | [246] |
| Polyaniline (PANI) | TiO\textsubscript{2} NTs (anatase/rutile) | PANI deposition by electropolymerization | PANI/TiO\textsubscript{2} NTs supported the viability/proliferation of MG-63 osteoblasts and showed good anti-biofilm activity. | [258] |
| Se-Chitosan | TiO\textsubscript{2} NTs (amorphous) | Se is deposited by electrodeposition and Chitosan by spin coating | NTs-Se-Chi samples showed excellent antibacterial activity and promoted the proliferation and biological functions of healthy osteoblasts while inhibiting the growth of cancerous osteoblasts. | [239] |
Table 2. Cont.

| Drug/Compound | Nanostructure | Drug/Compound Deposition | Biological Effects | Reference |
|---------------|---------------|--------------------------|-------------------|-----------|
| Metformin (MET)-Chitosan | TiO₂ NTs | D: 160 nm L: ≈9–10 µm | 5 cycles of deposition-drying in air of MET solution in fetal bovine serum. Chitosan was deposited by spin coating | [255] |

Briefly, for Ag nanoparticle (NP) decoration, deposition is achieved via: (i) a combination of solution deposition and subsequent photodeposition or UV-induced photoreduction [242,250], which can also be available for depositing the Ag⁺ ions on polydopamine decorated TiO₂ NTs [231], or (ii) solution deposition and reduction by gluconolactone (reducing the silver ammonia to Ag NPs) [243], and more recent approaches such as (iii) anodizing alloys (TiNbAg [242] and TiNbAgPt alloys [93]), or (iv) ion implantation (graded Ag incorporation into TiO₂ NTs by silver plasma immersion ion implantation [126] or the mixed Zn-Ag ion co-implantation [251]). While most of the works focus on the antibacterial activity of Ag decorated NTs (see also Coman et al. [259]), some researchers investigated also the interactions between Ag and host cells in view of biocompatibility. While an Ag overdose strongly influences biocompatibility due to increased toxicity against the host cells and surrounding tissue [126,241], if the Ag amount is optimal, the Ag-decorated NTs have been reported to show a similar cellular response as the undecorated NTs with epithelial cells and fibroblasts in vitro and a minimal inflammatory response in vivo [126]. Moreover, Taipinia et al. [242] reported that anodic NTs grown on TiNb alloys containing Ag have antibacterial activity without a detrimental effect on MC3T3-E1 pre-osteoblasts rather promoting cell proliferation.

Gao et al. [244] investigated Ag₂O NPs-embedded NT structures (size 5 to 20 nm, and 0–15 at.% Ag) by a combination of sputtering titanium and silver on Ti and subsequent anodization, showing adequate antibacterial properties without cytotoxicity. The crystallized Ag₂O NPs were embedded in an amorphous TiO₂ NT wall enabling the sustained and slow Ag⁺ release, thus minimizing the cytotoxicity and ensuring a long-lasting antibacterial activity. More importantly, they showed no appreciable influence on the osteoblast viability, proliferation, and differentiation compared to the Ag-free bare nanotubes. Similar results of a bactericidal effect against Escherichia coli without detrimental effects on human osteoblast proliferation were reported when the Ag₂O NPs were deposited by physical vapor deposition on NTs grown on Ti6Al4V [245].

Strontium represents another widely used element for functionalization of biomedical materials due to its dual mode of action on bone cells, namely, stimulation of osteoblast proliferation and differentiation, and inhibition of osteoclast function [260,261]. Sr²⁺ decoration of the NTs via a hydrothermal treatment has been reported to increase the osteobonding ability of the materials based on in vitro experiments with Saos-2 osteosarcoma cells and MC3T3-E1 pre-osteoblasts, respectively [249,250]. It is worth noting that in the latter study, Pan et al. [250] additionally incorporated Ag NPs onto Sr-loaded NTs by adsorption of Ag⁺ from 0.02 AgNO₃ solution and subsequent UV irradiation. The simultaneous presence of Sr and Ag endowed the materials with excellent antibacterial properties and osteogenic capability in terms of pre-osteoblast adhesion, proliferation, and mineralization, as well as gene expressions of osteoblast-specific markers.

A similar hydrothermal treatment in Zn acetate solution results in Zn decoration of the TiO₂ NTs with very good antibacterial activity and biocompatibility—while an optimum sample in respect of tube diameter (80 nm) and the time of the hydrothermal
treatment (3 h) with optimal Zn release significantly enhances the osteogenic differentiation of the MSCs due to increased matrix protein deposition [246]. In addition, Jin et al. [251] reported that a simultaneous Zn and Ag co-decoration of the NTs (by plasma immersion ion implantation) results in Ag NP decoration on a Zn layer covering the NTs and this co-decoration increases initial adhesion and spreading, proliferation, differentiation, and osteogenesis of MSCs coupled with a long-term antibacterial effect (compared to Ti, or Ag or Zn single deposition), in both in vitro and in vivo experiments (due to formation of Ag-Zn micro-galvanic couples [262]). ZnO NPs decorated on NTs have also an antibacterial effect, and once the optimal loading is exceeded its efficacy can be tackled with further co-doping of Ag [247]. In a most recent study, Chen et al. [263] reported on the fabrication of Zn-incorporated TiO$_2$ surfaces and their influence on the osteogenic microenvironment and bone formation. In their culture model, a MC3T3-E1 pre-osteoblasts cell line was grown in the conditioned media (CM) derived from a RAW 264.7 macrophages cell lines cultured in standard or pro-inflammatory (stimulation with lipopolysaccharide—LPS) conditions on these Zn-incorporated TiO$_2$ surfaces. The results showed that macrophages cultured on Zn-incorporated TiO$_2$ NTs display a M2 phenotype, while M1 markers were moderately inhibited, as compared to the LPS group. The pre-osteoblasts grown on Zn-incorporated NTs incubated in CM showed increased cell adhesion and proliferation, as well as osteogenic differentiation in comparison to their TiO$_2$ NTs counterparts and the Ti group. The authors hypothesized that superimposing Zn onto a titania NT surface could increase the osteogenic potential of osteoblasts through the improved immunomodulatory function of macrophages. The study thus brought to attention the crucial roles of Zn in both bone homeostasis and regeneration [264] and in innate and adaptive immune systems [265].

Cu is a well-known antibacterial agent that is widely used in biomedical applications with other biomaterials and has anti-inflammatory, anti-microbial, and anti-proliferative properties [266]. More recently, the focus was on developing TiCu alloys, and their in vitro and in vivo evaluation corroborated their good biocompatibility and osteogenesis ability (e.g., 5 wt.% Cu content [267,268], or 10 wt.% Cu [248]). TiCu alloys enhanced the expressions of osteogenesis-related genes (including alkaline phosphatase - ALP, Collagen I, osteopontin - OPN, and osteocalcin - OCN) in vitro [268] and promoted the surrounding bone-bonding (bone-to-implant contact) and the osteogenesis in vivo [267]. In addition, Wang et al. [248] has reported the anodization of TiCu alloys (containing 90-x % Ti, 10% Cu and x Al, with x = 0.45) to have excellent antibacterial activity and minimal cytotoxicity on osteoblast cells. Currently, Wu et al. [124] has shown that a multifunctional TaCu coating on anodic TiO$_2$ by magnetron sputtering (Ta:Cu 1:1 at.% ratio) enables effective bacteriostatic properties with distinct angiogenesis compared to bare NTs or a Ta coating only.

During the electrochemical anodization, ions such as Ca and P can be incorporated into the anodic nanostructures by anodizing in an electrolyte containing simulated body fluid (SBF) and subsequent cathodic deposition of Ag. This resulted in a functionalized NT layer with calcium, phosphorus and silver (Ca-P-Ag), which can enhance bone-like apatite formation in SBF and stimulate cell adhesion and proliferation of murine pre-osteoblast cells accompanying inhibition of the bacterial growth [252]. Similar results can be achieved by thin Ca-P layer deposition on NTs via immersion in Hank’s solution, followed by nano-Ag deposition by magnetron sputtering [253].

For Ag NPs decoration of NTs, a nano-hydroxyapatite co-decoration can be employed too [243], which has been reported to increase the biocompatibility and improve the control of the release rate [269,270]. A similar approach can also be used for the incorporation of drugs [36,256]. As depicted in Figure 11, polymers can be employed for filling/capping of the NTs to enhance the biocompatibility or to control the drug release (polyaniline [258], polyethylene glycol [233], chitosan [234,238] or gelatin and chitosan [240,257] as a capping agent).

For targeting antibacterial effect combined with an improved osteoinduction, the release mechanism and release rate of gentamicin [254,271], metformin [255], ibuprofen [37] from TiO$_2$ NTs have been extensively studied [127,229,236]. Further, a variety of com-
Combination sets of drugs and NT decorations has been designed and suggested to get the synergistic effects on antibacterial effect and improved osteoinduction. Draghi et al. [254] confirmed that loading gentamicin into the smaller diameter (31 nm) NTs resulted in a protracted release and antibacterial action together with improved cell adhesion and proliferation of an osteosarcoma cell line. Lai et al. [257] tailored the release of simvastatin from chitosan/gelatin-coated TiO$_2$ NTs improving osteoblast differentiation and inhibition of osteoclastic differentiation (compared to free-drug bare NTs).

BMP2-loaded TiO$_2$ NTs with a hyaluronidase-sensitive-multilayer coating consisting of chitosan, sodium hyaluronate-lauric acid, chitosan, and gelatin have been reported to have good biocompatibility, higher cell viability, mineralization capability, and antibacterial effect [272]. Metformin-loaded NTs with a 15 layers chitosan deposition to control the drug release led to a long-term release rate of up to 21 days and significantly decreased the burst release while promoting cell attachment and proliferation of MSCs [255].

3.2.3. In Vivo Drug Delivery Approaches Using Anodic TiO$_2$ Nanotube Implants

In contrast to abundant in vitro research works investigating the advantages of TiO$_2$ NTs as drug delivery systems, only a limited number of studies progressed to in vivo drug delivery experiments on animal models using nanostructured implants based on TiO$_2$ NTs (please see Table 3).

Table 3. Overview of bioactive compounds delivery platforms using anodic TiO$_2$ nanostructures in the animal in vivo models (D—diameter, L—length).

| Bioactive Compound | Implant Characteristics | Drug Loading Method | Animal in Vivo Model/Biological Effects | Reference |
|--------------------|-------------------------|---------------------|----------------------------------------|-----------|
| rhBMP2             | TiO$_2$ NTs D: ~70 nm, ~110 nm; Implant: D 3.5 mm; L 8.5 mm | Dip-coating in 1.5 mg rhBMP-2/mL (in a vacuum chamber) | Pilot in vivo study: New Zealand white rabbits, 4 types of implants (proximal tibia); rhBMP2-loaded implants: the highest BIC and enhanced bone remodeling. | [38] |
| rhBMP2/ Lenti-BMP2 | TiO$_2$ NTs: D ~100; L: 400 nm; Ti rods (D: 2 mm; L: 8 mm) | Lyophilization of Lenti-BMP2 in the presence of trehalose | Femur defect model in Fisher 344 rats; TiO$_2$-Lyo-Tre-BMP2 implant: most effective in terms of BMP2 stability, sustained release, bioactivity, bone regeneration. | [273,274] |
| rhBMP2 and Ibuprofen (IBU) | TiO$_2$ NTs: D ~70 nm; L: 5 μm; Ti rods (D: 2 mm; L: 8 mm) | IBU (1.5 mg/mL) and rhBMP-2 (10 mg/mL) loading by dip coating (3×), lyophilized, freeze and vacuum dried. | IBU-NTs behaved as an anti-inflammatory drug and improved the osseointegration of orthodontic miniscrews in vivo. However, the effect of rhBMP2-loaded NTs on the osseointegration was slightly lower. | [37] |
| rhPDGF-BB          | Ti rods (D: 2 mm; L: 8 mm) NTs: D—70 nm | Immersion in 100 μg/mL rhPDGF-BB at RT (PDGF group) or put in the vacuum pump (PDGF + Vacuum group) for 10 min | O VX rats with bilateral femurs were used for the implantation; the newly designed coating contributed to the new bone formation surrounding the implant and enhanced bone fixation in O VX rats showing great promise for clinical applications in osteoporotic patients. | [275] |
| Bioactive Compound | Implant Characteristics | Drug Loading Method | Animal in Vivo Model/Biological Effects | Reference |
|---------------------|-------------------------|---------------------|----------------------------------------|-----------|
| **Alendronate (ALN)** | Ti rods (D: 3 mm; L: 13 mm); NTs D: 70 nm; L: 0.7–1.0 µm (anodic oxidation), HA layers: alternate immersion method on TiO₂ NTs surface. | NTs-HA-ALN implant: immersion into ALN 20 mg/mL solution at RT (12 h). NT-ALN implant: physical absorption of ALN on TiO₂ NTs | Implants into the femoral epiphysis of OVX female New Zealand white rabbits. NTs-HA-ALN implants showed great potential for increasing osseointegration as compared to Ti, NTs, and NTs-HA implants, with the highest anti-osteoporosis potential | [256] |
| **Icariin (ICA)** | Cylindrical implants (D: 1.5 mm, L: 2 mm); Anodic TiO₂ NT (D: 80 ± 10 nm) Immersion in ICA solution (2 days), drying at 37 °C (1 day): PLGA coating (twice in a dropwise manner) | Sprague Dawley rats received implants in the femora's mid-diaphysis; TiO₂ NT structure and ICA synergistically promote osteoblasts' function and PLGA coating endowed the implant surface with better osteogenic/osseointegration ability. | [276] |
| **Silicon (Si)** | Ti screws, inner/outer D: 1.7/2 mm, L: 10 mm; Anodic NTs: inner/outer D 60/80 nm. Si plasma immersion ion implantation (PIII) method | Sprague Dawley rats received implants in the distal femur in the horizontal direction; Si-TiO₂-NTs induced enhanced early osseointegration positive effect on implant osseointegration and trabecular microarchitecture formation. | [277] |
| **Polyhexa-methylene guanidine (PHMG)** | cp-Ti rods (D: 3.175 mm; L: 1.5 ± 0.1 cm; Anodic TiO₂ NTs: D—46.4 ± 5.9 nm, L—650–800 nm. Addition of 100 µL of 25% PHMG aqueous solution onto rods dropwise and drying by a vacuum oven at RT for 1 h (×10 times). | Rabbits implanted with S. aureus-contaminated rods in the femoral medullary cavity; PHMG-NTs showed an excellent capacity to prevent bacterial infections, as well as to promote osteogenic differentiation by increased expression of osteogenic-related genes in the femur tissues around the implants. | [278] |
| **Propolis (PL)** | Ti rods (D: 0.85 mm; L: 4.5 mm) screw-processed at a thread angle of 20 degrees; anodic TiO₂ NT (D: 60–90 nm). Immersion in propolis solution for 24 h at 25 °C followed by vacuum-drying at 25 °C for 24 h | Sprague Dawley rat mandibular model; increased new bone formation and mineral density around the PL-NT-Ti implant; enhanced osteogenic differentiation and increased expression of collagen fibers while pro-inflammatory markers decreased | [279] |

The most recent studies have been focused on growth factors applications that can endow the NT surfaces with additional benefits in terms of implant osseointegration. BMP2, which was proven as the most potent stimulator in inducing bone regeneration [280,281], has been widely used to functionalize TiO₂ NTs. One of the well-designed in vivo studies performed by Lee et al. [38] established anodic TiO₂ NTs (with diameters of ~70 nm and ~110 nm, respectively) decorated with rhBMP2 by dip coating. The in vivo study was performed on the following implant groups in New Zealand white rabbits: (Group 1) a machined surface; (Group 2) a sandblasted large-grit and acid-etched SLA implant (as a positive control group); (Group 3) TiO₂ NT; and Group 4) TiO₂ NTs with rhBMP-2. Histomorphometric/micro-computed tomographic analysis at 8 weeks post-implantation showed that Group 4 obtained the highest BIC and bone volume ratio. As a conclusion, the authors suggested the designed NT drug delivery platform as a promising reservoir that allows a slow and sustained rhBMP2 release to reinforce implant osseointegration.
In line with this, Zhang et al. [273,274] proposed a new strategy to functionalize TiO$_2$ NTs by loading a lentiviral vector encoding BMP2 (Lenti-BMP2) by lyophilization following trehalose addition. The obtained TiO$_2$-Lyo-Tre-BMP2 nanoplatform ensured a slow and prolonged BMP2 release and promoted the highest extent osteogenic differentiation of bone marrow stromal cells as well as anti-inflammatory activity. The studies suggested that the delivery of BMP2 using the Lyo-Tre-based system is an effective method avoiding the adverse effects induced by the administration of high doses of BMP2.

In other studies, the osseointegration capability of BMP2 was compared with other bioactive agents such as the anti-inflammatory drug ibuprofen [37] (see also Table 2). The in vivo study [37] composed of four implant groups (conventional, NT coated, rhBMP2 loaded-, and ibuprofen loaded-TiO$_2$ NTs) showed that the histological analysis at eight weeks after implantation surprisingly revealed the highest BIC ratio (71.6%) in the ibuprofen-loaded group, while rhBMP2 loaded implants showed a significantly decreased BIC (24.6%). The authors explained this effect by the osteoclastic bone-resorbing activity of rhBMP2, indicating that careful fine-tuning may be required to get the optimal dosage of rhBMP2 for successful osseointegration.

Another growth factor widely investigated for bone regeneration is platelet-derived growth factor-BB (PDGF-BB) [275], as it can stimulate the recruitment, proliferation, and osteogenic differentiation of osteoprogenitor cells or MSCs as well as the angiogenesis process [282]. Zhang et al. [275] applied rhPDGF-BB on 70 nm diameter TiO$_2$ NT surfaces arrays by vacuum extraction for in vivo osseointegration in ovariectomized (OVX) rats as an osteoporosis-induced animal model. Protein particles aggregated on the surface and inside NTs could be slowly released for at least 14 days without losing their bioactivity. A higher rhPDGF-BB immobilization to the underlying TiO$_2$ NTs substrate by vacuum extraction method led to an enhanced cell adhesion, proliferation, and osteogenic differentiation in vitro. Further, an in vivo study of osseointegration (oxalic acid-etched Ti group; TiO$_2$ NTs modified Ti group; PDGF-coated TiO$_2$ NTs group; and PDGF coated TiO$_2$ NTs + Vacuum group) showed that the NTs loaded with rhPDGF-BB under vacuum improve the implant fixation ability and the rapid new bone formation in OVX rats, suggesting a novel implant coating strategy in the treatment of bone defects associated with osteoporosis.

To endow the NT surface with a local anti-osteoporosis property, Shen et al. [256] proposed a loading of alendronate (ALN), a powerful anti-osteoporosis compound that is largely used in clinics due to its antiresorptive capacity [283], onto hydroxyapatite-TiO$_2$ NT substrates (NTs diameter 70 nm, length 0.7–1.0 µm). The resulted material (NTs-HA-ALN) has shown great potential in improving the proliferation and osteogenic differentiation of osteoblasts isolated from neonatal rat calvaria and inhibiting differentiation of RAW 264.7 in mature osteoclasts, compared to bare Ti, TiO$_2$ NTs, and TiO$_2$ NTs deposited with nano-HA layers (NTs-HA). Furthermore, in vivo tests with osteoporotic rabbits, attested the highest potential of NTs-HA-ALN implants to enhance the local osseointegration with excellent pro-osteogenic and anti-osteoporosis properties at three months after implantation, considered as the synergistic effects of the release of Ca$^{2+}$ and ALN.

Natural compounds such as propolis [279] and icariin [276,284,285] have also been introduced as another strategy of surface functionalization to increase the in vivo performance of TiO$_2$ NTs. Propolis is a natural compound collected by honeybees from various plants whose biological activity mainly depends on the flavonoids from the polyphenolic fractions, followed by aromatic acids, phenolic acid esters, etc. [286]. Its beneficial effects on bone healing are well known and attributed to its anti-inflammatory, antioxidant, and anti-osteoclastic activities [287]. Somsanith et al. [279] loaded propolis (PL) on anodic TiO$_2$ NTs (PL-NTs-Ti) and showcased their ability to sustain the viability and differentiation of MC3T3-E1 pre-osteoblasts and in vivo osseointegration in a rat mandibular model. Besides enhanced cell viability and alkaline phosphatase activity in cell culture experiments, the PL-NTs-Ti resulted in enhanced formation of new bone and increased mineral density in the region surrounding the implant as well as higher expressions of collagen fibers, and BMP2/BMP7 as compared to the drug-free TiO$_2$. Moreover, the peri-implant expressions
of the pro-inflammatory cytokines (IL-1ß, and TNF-α) were significantly reduced, suggesting that the propolis-functionalized nanostructure also has the potential to inhibit early inflammation and block peri-implantitis.

Ma et al. [276] functionalized TiO₂ NT surfaces with icariin (ICA, a Herba Epimedii derived flavonoid with osteogenic and anti-osteoclastogenic effects [288]) and poly lactic-co-glycolic acid (PLGA). The resulting NTs-ICA-PLGA substrate was able to ensure a sustained drug release up to two weeks with in vitro best osteogenic differentiation and in vivo higher bone formation area percentage compared to other groups during the early-stage of osseointegration. Another approach to improve the osseointegration of TiO₂ NTs was the incorporation of Sr and icariin onto the anodized Ti surface through hydrothermal treatment and vacuum freeze drying, respectively [284]. In this study, the authors demonstrated that icariin loading onto Sr-containing TiO₂ NT surfaces exerted additional positive effects on the pre-osteoblast behavior in terms of proliferation and osteogenic differentiation, as well as on the bone formation around screw-shaped Ti-based implants. These effects can be ascribed to the icariin ability to guide bone regeneration and promote osteogenesis [289] combined with the dual effects of Sr on bone tissue through stimulating the proliferation and differentiation of osteoblasts and inhibiting the osteoclast activity [290,291], implying that the ICA-Sr-TiO₂ coating may be a potentially useful option in osteoporotic patients.

Si, another bioactive trace element, possesses a bone affinity that has been shown to enhance osteoblast proliferation and differentiation, stimulate collagen synthesis and bone mineralization [292,293], and to further inhibit the osteoclastogenesis and bone resorption processes [294]. Zhao et al. [277] fabricated Ti substrates (discs and screws) modified with silicon doped TiO₂ NTs (Si-TiO₂-NTs) by in situ anodization and Si plasma immersion ion implantation (PIII) technique and compared their activities with those for Ti and TiO₂ NTs. The in vivo results, in line with the in vitro findings, showed that the Si-TiO₂-NTs surface increased pre-osteoblast cell proliferation and differentiation, indicated extensive bone apposition between the screw threads at the spongy level on this surface and the formation of more new trabecular bone as compared to bare Ti and TiO₂-NTs screws.

Considering that chronic implant-associated bone infections represent a major problem in orthopedic and trauma-related surgery due to the severe complications in the affected patients, increasing interest has been given to design surfaces with bacteriostatic and bactericidal properties. With this purpose, many antibacterial agents such as antibiotics, metal ions, anti-microbial peptides, and biopolymer coatings [14,227,295,296] were incorporated into TiO₂ NTs and investigated for their therapeutic efficiency, mostly by in vitro antibacterial and cell culture-based studies. A very recent paper by Wu et al. [278] has revealed the antibacterial potential of polyhexamethylene guanidine (PHMG) coated TiO₂ NTs (PHMG-NTs) in an animal model implanted with Staphylococcus aureus-contaminated rods in the femoral medullary cavity. This eco-friendly polymer was shown to exhibit a high and broad-spectrum antibacterial effect, which can efficiently inhibit bacterial biofilm formation [297,298] with improved bone-forming capacity.

4. Conclusions and Outlook

Nanoscale surface modification of Ti or Ti-based alloys via electrochemical anodization, resulting in TiO₂ nanostructures including nanotubes (NTs), has gained significant recognition and insight in biomedical applications. Herein we discussed the current state-of-the-art in NT morphology (nanotube, nanopore, mesoporous) and their synthesis (closed-packed, open-top, and spaced NTs or long-range ordered NTs), with the critical aspects affecting the drug delivery. Further, we reviewed the properties of anodic NTs including crystallinity, surface roughness, wetting characteristics, and corrosion resistance.

Next, we evaluated the key interactions of anodic TiO₂ nanostructures with bone-related cells, such as osteoblast cells and mesenchymal stem cells, emphasizing the influence of the nanoscale topography on the interactions with these cells. In addition, a more recent approach, highlighting the synergistic effects of electrical stimulation on cells combined
with the nanoscale environmental cues from TiO$_2$ NTs for enhancing bone induction was further discussed. Such a synergistic approach would be promising for tissue engineering applications using nanostructured scaffolds materials based on anodic TiO$_2$ NTs with concomitant drug delivery and/or electrical/electromagnetic stimulation.

For dental or orthopedic implants, anodic TiO$_2$ NTs and nanostructures with surface modifications delivering specific targeting drugs are shown to be clinically useful approaches for superior osteoinduction and successful osseointegration, especially in medically compromised patients including osteoporosis, chronic inflammatory, and metabolic diseases in the future. For successful osseointegration, nanoscale topographical cues tuning by nanoscale lateral spacing can be a powerful stimulator when combined with the specific nanotubular/nanoporous shape and high surface area as a delivery platform, allowing decoration/incorporations of antibacterial agents and/or other drugs/active agents with time-scheduled drug release rate. The release rate can be further improved avoiding an initial burst release, either by the use of polymer capping or by tuning the morphology of the nanostructures. We discussed in detail the various modifications of anodic NTs with metal ions, nanoparticles, drugs, growth peptides for controlling the drug release rate and enhancing antibacterial and osteoinductive properties, both in vitro and in vivo.

Overall, in recent decades, we fully recognized the nanoscale significance of bio-implantable substrate surface topography on a local wound healing and bone regeneration, and now more clinicians/researchers in tissue engineering fields are excited to find out which combinations of stimuli including nanoscale topography, electrical, and/or biomolecular approaches could provide a best-fit synergistic effect on bone tissue engineering.

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