Bioactive Carbon-Based Hybrid 3D Scaffolds for Osteoblast Growth

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Supporting Information

ABSTRACT: Bone, nerve, and heart tissue engineering place high demands on the conductivity of three-dimensional (3D) scaffolds. Fibrous carbon-based scaffolds are excellent material candidates to fulfill these requirements. Here, we show that highly porous (up to 94%) hybrid 3D framework structures with hierarchical architecture, consisting of microfiber composites of self-entangled carbon nanotubes (CNTs) and bioactive nanoparticles are highly suitable for growing cells. The hybrid 3D structures are fabricated by infiltrating a combination of CNTs and bioactive materials into a porous (∼94%) zinc oxide (ZnO) sacrificial template, followed by the removal of the ZnO backbone via a H2 thermal reduction process. Simultaneously, the bioactive nanoparticles are sintered. In this way, conductive and mechanically stable 3D composite scaffolds are created. The adopted strategy demonstrates great potential for implementing low-dimensional bioactive materials, such as hydroxyapatite (HA) and bioactive glass nanoparticles (BGN), into 3D carbon-based microfibrous networks. It is demonstrated that the incorporation of HA nanoparticles and BGN promotes the biomineralization ability and the protein adsorption capacity of the scaffolds significantly, as well as fibroblast and osteoblast adhesion. These results demonstrate that the developed carbon-based bioactive scaffolds are promising materials for bone tissue engineering and related applications.

KEYWORDS: carbon nanotubes, 3D scaffolds, bioactive glass, hydroxyapatite, osteoblasts

1. INTRODUCTION

Bioactive materials are highly important for inducing specific cellular responses, particularly in tissue engineering. A highly promising bioactive material in bone tissue engineering is silicate bioactive glass (BG). In particular, the presence of CaO in silicate BG is decisive in the formation of a bone-like calcium phosphate layer on the material. This surface layer results in a strong connection between the biomaterial and both bone and soft tissue. To expand the interface between BG and tissue, BG can be prepared as scaffolds or incorporated into scaffolds. Depending on the microstructure and fabrication method of the BG, the Young’s modulus of porous scaffolds made of BG can be varied between 1 and 22 MPa. In addition, BG can be degraded by releasing ions into the environment so that a main component of bone, hydroxyapatite (HA), is formed. BG is also known to support the adhesion and proliferation of osteoblast cells, such as MC3T3-E1 and MLO-A5. In addition, many studies have shown that the ions released from BG, including Si, Ca, P, and Cu, play an important role in enhancing osteoblast proliferation and bone formation. Another important bioactive material is hydroxyapatite, as it is a main component of bone within the human body. It has frequently been used as a coating to promote bone ingrowth or as a bone graft substitute. BG is often implemented in biomaterials in the form of composites. Out of such composite systems, BG and carbon nanotubes (CNTs) have shown to be most promising due to a mutually beneficial modification of the features and functionality of each other. On the one hand, CNTs enhance the features of BG such as stiffness, electrical conductivity, and surface roughness in several ways: (i) the mechanical properties, such as stiffness and fracture toughness, are improved by the CNTs; (ii) the local and overall electrical conductivity is increased by CNTs; (iii) CNTs lead to a rougher surface on the nanoscale and consequently to an improvement of cell attachment and proliferation. On the other hand, BG makes CNTs more appropriate for biomedical applications by accelerating biomineralization and increasing the concentration of physiologically relevant ions, thus resulting in higher cell proliferation rates and faster regeneration of soft tissue.
To exploit the unique properties of composites made of CNTs and bioactive materials (such as BG and HA) for tissue engineering, three-dimensional (3D) scaffold fabrication methods are needed that provide a mechanically stable 3D environment that supports cell proliferation. In such 3D environments, the pore architecture is one of the most decisive parameters as it affects cell growth by determining nutrient and oxygen diffusion, waste removal, as well as the growth rate of the cells. Open porous and interconnected pore architectures are strongly necessary to facilitate these parameters. Commonly used techniques to generate pores in materials, such as salt-leaching, freeze-drying, gas foaming, sol–gel crosslinking, and phase separation only provide very limited control of the internal pore architecture of the scaffold material for tissue engineering. Therefore, rapid prototyping techniques have been employed recently, which enable the fabrication of scaffolds with control over pore size and architecture together with good design reproducibility. Nevertheless, rapid prototyping techniques are limited with respect to the biomaterials that can be printed and they cannot be used to fabricate 3D structures from nanomaterials only.

Recently, an infiltration-based synthesis method for the fabrication of porous, 3D scaffolds consisting of interconnected CNT-based microtubes has been reported. With this method, open porous scaffolds with total dimensions on the cm³ scale can be fabricated, resulting in a 3D hierarchical CNT tube (CNTT) structure with improved mechanical and electrical properties compared to other 3D assemblies prepared by wet chemistry. As the reported fabrication method relies on the infiltration of nanoparticle dispersions into sacrificial ZnO templates, in principle, also other nanoparticle types besides CNTs can be embedded. Therefore, this method is a great and versatile starting point for fabricating hybrid 3D scaffold materials with properties that can be tailored towards specific, electronic, catalytic, or biomedical applications, e.g., by the incorporation of bioactive nanoparticles.

We show here that low-dimensional bioactive materials in the form of bioactive glass and hydroxyapatite nanoparticles can be integrated into CNTT materials, to make it bioactive and...
supportive for cell growth. The bioactive nanoparticles incorporated into the scaffolds are sintered during a single step etching/sintering process. The structure of the resulting material was characterized with scanning electron microscopy (SEM), transmission electron microscopy (TEM), and X-ray diffraction (XRD). To prove the bioactivity of the scaffolds, we investigated biomineralization as well as their protein adsorption capacity. In vitro studies demonstrate that fibroblasts and osteoblasts can adhere to the fibrous structures of the hybrid CNT network, demonstrating the feasibility of this 3D composite material for bone tissue engineering.

2. RESULTS AND DISCUSSION

2.1. Bioactive Carbon-Based Scaffolds. The successful fabrication of free-standing, fibrous 3D composite scaffolds, consisting of CNTs and a nanoscale bioactive ceramic biomaterial has been achieved via a very simple strategy based on a ceramic template. Figure 1A shows a schematic representation of the fabrication process, in which compressed and sintered ZnO tetrapods (Figure 1B) serve as sacrificial materials. To generate free-standing carbon-based and bioactive composite structures these 3D and porous (porosity ~94%) ZnO templates were infiltrated with an aqueous dispersion of CNTs and either BGN, HA nanoparticles or both of them. Due to the strong capillary forces and the superhydrophilic properties of the ZnO template,27 the dispersion is rapidly soaked into the template during the infiltration process. Figure 1C shows an SEM image of a ZnO template after infiltration with CNTs and BGN. Figure 1E,F show SEM images of the templates after infiltration with CNTs and HA nanoparticles. In the experiments, it is in principle also possible to tailor the amount of CNTs as well as the amount of BGN and HA nanoparticles added on the template by controlling the volume of the injected nanoparticle dispersion into the ZnO network.

To generate free-standing hybrid structures, we needed to remove the ZnO. This is also necessary to make the materials biocompatible, as a high amount of ZnO is toxic for cells.28 Most methods to remove ZnO are based on wet chemistry, e.g., using diluted HCl. However, since an acidic treatment would also dissolve the incorporated bioactive materials, a H2 etching process was used to remove the ZnO.29 At 900 °C, ZnO is completely reduced by H2, evaporated to Zn (gas) without the presence of a carbon source, and removed by the employed carrier gas (argon). In Figure 1D−J, SEM images of the resulting structures are shown. Figure 1D shows CNT and BGN scaffolds after etching of the ZnO template. Figure 1H,1J show that broken microtubes are hollow, proving that the entire ZnO template has been removed during the etching process. The free-
The bioactivity of the scaffolds is induced by the formation of new bone tissue based on the principle of tissue engineering in this context. The idea of tissue engineering is to generate scaffolds that replace bone in load-bearing areas, as their compressive strength is smaller than that of trabecular bone. Instead, the mechanical properties will strengthen upon implantation so that a new bone is formed. The suitability of soft materials for bone tissue engineering has recently been shown by Huebsch et al. ([Table 1](#tab1)). These ceramic nanoparticles are believed to contribute to a further reinforcement of the mechanical properties of the scaffold. Moreover, the geometry of the networks provides highly accessible interconnected pores from all sides, whereas the preferential alignment of CNTs in other 3D assemblies confines the accessibility of cavities.

In addition, the porous structure of our scaffolds leads to a large free volume, which should in turn result in higher bioactivity. The scaffolds possess up to 94% porosity ([Table S1](#tabS1)), thus being in the range of that of highly porous BG-containing scaffolds. The porosity of the scaffolds presented is higher than that of 3D printed biomaterials that contain HA nanoparticles, including HA nanoparticle incorporated poly(lactide-co-glycolide) with up to 63.33% of porosity. Furthermore, due to the interwoven arrangement of CNTs in the microtubes, we assume that the scaffolds possess a similar conductivity ([Table 1](#tab1)) as the recently reported self-entangled CNT assemblies, but with the additional feature that the incorporated ceramic nanoparticles make the material bioactive.

It is important to note that our scaffolds are not intended to replace bone in load-bearing areas, as their compressive strength ([Table 1](#tab1)) is smaller than that of trabecular bone. Instead, the idea of tissue engineering in this context is to generate scaffolds, which induce the formation of new bone tissue based on the bioactivity of the scaffold. If the scaffold exhibits bioactivity, the mechanical properties will strengthen upon implantation so that a new bone is formed. The suitability of soft materials for bone tissue engineering has recently been shown by Huebsch et al., who used soft hydrogels to induce bone formation. As our scaffolds are bioactive (see Section 2.5) and support osteoblast growth (see Section 2.6), we are convinced that they are suitable for bone tissue engineering.

### 2.2. Morphology and Composition of Ceramic Nanoparticles on CNT Scaffold

**Table 1. Conductivity and Compressive Strength of CNTT−HA and CNTT−BGN Scaffolds**

| scaffold     | specific conductivity (S/m) | compressive strength (kPa) | compressive strength/density (kPa cm^3/g) | Young’s modulus (MPa) |
|--------------|----------------------------|----------------------------|-------------------------------------------|-----------------------|
| CNTT−HA      | ∼0.88                      | ∼7.2                       | ∼27                                       | ∼0.4                  |
| CNTT−BGN     | ∼12.7                      | ∼32                        | ∼118                                      | ∼1                    |

“The large discrepancy in conductivity between CNTT−HA and CNTT−BGN is most probably a result of clamping difficulties, as the surface of the material is covered with HA and BGN, respectively. Still, the conductivity is in the range of that of CNTTs without incorporated ceramic nanoparticles.”

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**Figure 2.** TEM images of CNTT−BGN, CNTT−HA, and CNTT−BGN/HA scaffolds. The striking result from these images is that all generated structures are free-standing and the ZnO template has been completely removed. Furthermore, the images show that the ceramic nanoparticles were fused to each other in the etching/sintering procedure. In addition, HA and BGN were fused to each other in the etching/sintering procedure. In addition, HA and BGN were fused to each other in the etching/sintering procedure.

**Figure 3.** XRD patterns of CNTT−BGN, CNTT−HA, and CNTT−BGN/HA scaffolds. CNTT−BGN/HA composites. Interestingly, BGN mainly retain their spherical shape and are only slightly fused together. In contrast, the initially also spherical HA nanoparticles were deformed to flakes or uneven shaped particles during the etching/sintering procedure. This allowed us to distinguish the particle types even in CNTT−BGN/HA scaffolds ([Figure 2E,F]). Furthermore, these images suggest that the HA particles bridged the gaps between the BGN in CNTT−BGN/HA samples ([Figure 2E,F]).

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The conductivity of the CNTs in our scaffolds form interwoven layers of self-entangled CNTs with high mechanical stability similar to the ones shown in previous work. Most importantly, the high temperature during the ZnO template removal fuses the ceramic nanoparticles to each other and leads to a physical interaction with the CNTs, resulting in a hollow free-standing entangled CNT architecture combined with sintered ceramic nanoparticles ([Figure 2]). These ceramic nanoparticles are believed to contribute to a further reinforcement of the mechanical properties of the scaffold. Moreover, the geometry of the networks provides highly accessible interconnected pores from all sides, whereas the preferential alignment of CNTs in other 3D assemblies confines the accessibility of cavities.

In addition, the porous structure of our scaffolds leads to a large free volume, which should in turn result in higher bioactivity. The scaffolds possess up to 94% porosity ([Table S1]), thus being in the range of that of highly porous BG-containing scaffolds. The porosity of the scaffolds presented is higher than that of 3D printed biomaterials that contain HA nanoparticles, including HA nanoparticle incorporated poly(lactide-co-glycolide) with up to 63.33% of porosity. Furthermore, due to the interwoven arrangement of CNTs in the microtubes, we assume that the scaffolds possess a similar conductivity ([Table 1]) as the recently reported self-entangled CNT assemblies, but with the additional feature that the incorporated ceramic nanoparticles make the material bioactive.

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To check the crystallinity of the ceramic nanoparticles after the fabrication process, we carried out an XRD study. The XRD pattern ([Figure 3]) of a CNTT−HA sample revealed diffraction peaks of HA. In contrast, the XRD pattern of a CNTT−BGN sample revealed no diffraction peaks related to BGN, but diffraction peaks related to α-quartz are recognized. Pristine BGN are amorphous and only possess broad peaks in the range of 2θ = 20–35°. Since the incorporated BGN had the
composition 92SiO$_2$–8CaO (in mol %), their crystallization behavior is similar to that of pure silica nanoparticles. However, in this study the processing temperature (900 °C) used for the fabrication of the scaffolds was high enough to cause the crystallization of silica-based nanoparticles, which explains the appearance of the diffraction peaks of α-quartz in CNTT–BGN. For the CNTT–BGN/HA structure, the diffraction pattern only correlates to that of crystallized HA. In addition, SiO$_2$–CaO-based BGN sintered at 900 °C should be partially crystallized according to previous work. Importantly, Figure 3 also shows the effect of different amounts of infiltrated CNTs: the intensity of the ceramic peaks decreased if more CNTs had been infiltrated.

The XRD results of the HA-containing scaffolds show no change of the crystalline phase of HA. As the first phase transformation of HA occurs at 1000–1100 °C, we assume that our HA particles are not decomposed at our sintering temperature (900 °C). Furthermore, in a previous study, no reaction of multiwalled CNTs (MWCNTs) with glass matrices to form SiC or other reaction phases in response to sintering between 850 and 1000 °C was detected by powder X-ray diffraction.

### 2.3. Protein Adsorption on Scaffolds

The adsorption of proteins on bioceramics is essential because it influences cellular adhesion and can facilitate scaffold integration into tissues. To investigate the protein adsorption capacity of CNTT–BGN and CNTT–HA scaffolds, we used bovine serum albumin (BSA) as a model protein. The adsorption capacity of the scaffolds was quantified for 4, 8, 12, 24, 48, and 72 h of incubation with protein solution. The bicinchoninic acid (BCA) assay shows that the protein adsorption is higher on CNTT–BGN scaffolds than on CNTT–HA scaffolds.

This difference in adsorbing proteins is highest during the first 4 h of incubation with proteins and levels out after 8 h of incubation. Despite the fact that there was a slight difference regarding the protein adsorption ratio between the two scaffold types, both exhibited a similar temporal progression of protein adsorption. This can be explained by the fact that CNTs presumably play the main role in protein adsorption due to the high amount of CNTs in the matrix in both the CNTT–HA and CNTT–BGN scaffolds (Figure 1). It is also important to mention here that the protein adsorption capacity of scaffolds is a decisive parameter for osteoblast attachment. As the CNT matrix can serve as an attachment site for a variety of extracellular matrix molecules, biomolecules, proteins, and growth factors, it can further mediate cell proliferation and adhesion. Interestingly, in the present study CNTT–BGN has a higher adsorption capacity than CNTT–HA (Figure 4).

This result could be due to an electrostatic interaction between the highly polar BSA and the BGN surface, which might be a result of the etching/sintering process, as explained in the following: at the sintering temperature of 900 °C, H$_2$ reacts with silica to form SiOx on the surface of BGN. Therefore, due to a change in surface charge, the BGN surface might have the potential to bind more BSA proteins. In addition, previous studies indicated that surface-modified bioactive glass adsorbs a higher amount of serum protein than hydroxyapatite.

### 2.4. Ion Release from Scaffolds in Phosphate Buffered Saline (PBS)

To explore the ion release capability of the fabricated hybrid scaffolds within biologically relevant media, we measured the concentration of Ca, Si, and Zn ions in phosphate buffered saline (PBS) after 4, 8, 12, 24, 96, 230, and 302 h of incubation with the scaffolds using inductively coupled plasma-mass spectrometry (ICP-MS) (Figure 5). Clearly, the amount of ions released from the scaffolds increased with incubation time. The release of Zn ions is almost zero (∼5 μg after 400 h), even after 16 days of immersing CNTT–BGN or CNTT–HA scaffolds in PBS. This proves again that the sacrificial ZnO templates had been entirely removed during the H$_2$ etching process. Hence, the hydrogen etching process at elevated temperatures provides a clear improvement compared to previously reported methods used for ZnO removal. As shown in Figure 5, a high amount of Ca$^{2+}$ ions was released into the PBS from CNTT–HA and CNTT–BGN structures during the first 24 h of immersion (Figure 5). Afterwards, the release rate of Ca$^{2+}$ ions was decreased. The high accessibility of Ca$^{2+}$ ions on the surface of BG and HA nanoparticles leads to a high Ca$^{2+}$ ion concentration gradient between PBS and the ceramic nanoparticles, presumably resulting in a high initial release rate and a smaller and more stable release rate for longer immersion times. As expected, we did not measure any Si ion release from the CNTT–HA scaffolds (Figure 5A). In contrast, CNTT–BGN scaffolds released Si ions (Figure 5B) and the release rate has a similar temporal progression as the one reported on pure BGN by Zheng et al., though with a slightly lower release rate than our scaffolds. Zheng et al. carried out their experiments in Dulbecco’s modified Eagle’s medium (DMEM), whereas we used PBS. The ions in DMEM could cause the formation of an amorphous Ca-P layer on the surface of BGN, thus leading to a decreased release rate.

### 2.5. In Vitro Bioactivity

A highly important indicator for the ability of scaffolds to integrate with bone is the formation of hydroxyapatite on the scaffolds. On our samples, needle-shaped crystals are formed on CNTT–BGN scaffolds after immersion in a simulated body fluid (SBF) for 7 days (Figure 6A), whereas such crystals are neither observed on CNTT–HA nor on CNTT–BGN/HA scaffolds after immersion in the SBF (Figure 6B,C). To explain this, we checked the results from energy dispersive X-ray spectroscopy (EDS). Our EDS results

![Figure 4. Bovine serum albumin adsorption (mean values) on CNTT–BGN and CNTT–HA scaffolds, measured with a BCA colorimetric assay. BGN containing structures have a slightly higher protein adsorption capacity compared to CNTT–HA scaffolds. (Each experiment was carried out on three samples and three replicates each. Error bars: standard deviation.)](image-url)
Supporting Information, Figure S1) show that phosphorus is present on the surface of CNTT−BGN filaments after immersion in the SBF for 7 days, which indicates the formation of calcium phosphate species. Additionally, silicon is still present according to the EDS spectrum of CNTT−BGN indicating that BGN have not completely dissolved yet after 7 days of immersion in SBF. After immersion in SBF for 14 days, cauliflower-like crystals are present on CNTT−BGN scaffolds (Figure 6D). This is a typical morphology of hydroxyapatite crystals formed on BG scaffolds after immersion in SBF. In addition, spherical BGN are still present on these scaffolds, showing that the time until full dissolution of the BGN is longer than 14 days. In general, BG is more surface reactive and has a higher dissolvability than other bioceramics (e.g., HA). Therefore, the formation of bone-like apatite crystals, which is a result of surface dissolution and ion exchange, occurs faster on BG surfaces than on HA surfaces in body fluids. In the study presented here, the apatite forming ability of the scaffolds was improved after coating of BGN in comparison with HA nanoparticles, considering that the apatite crystals were observed on CNTT−BGN scaffolds after 7 days of soaking in SBF, whereas no crystals were present on CNTT−HA. As shown in Figure 6C, the situation for CNTT−BGN/HA is similar as for CNTT−HA. Inhibited apatite formation occurs when BGN are mixed with non-reactive or less reactive materials. Our composite coating is like a glass-ceramic phase with less reactive surfaces, thus reducing the dissolution and ion exchange. The presence of HA in the coating also reduces the number of sites for apatite nucleation. Hence, apatite formation was inhibited in CNTT−BGN/HA scaffolds compared to CNTT−BGN. Interestingly, the formation of apatite crystals on our CNTT−BGN composite scaffolds took longer than when using pure BGN in SBF, where crystals had formed after 3 days of immersion. This result can be explained...
Figure 7. Fluorescence images of pre-osteoblast cells (MC3T3-E1) cultured for 21 days on CNTT–HA, CNTT–BGN, and CNTT–BGN/HA structures. The nucleus of the cells was stained with 4,6-diamidino-2-phenylindole (DAPI) (Hoechst), live cells with calcein AM (green), and dead cells with propidium iodide (red). All the scaffolds support osteoblast adhesion and proliferation. There is only a small number of dead cells after 21 days of cultivation (scale bars: 0.1 mm).

by the etching/sintering process that is necessary to remove the ZnO templates and caused the partial crystallization of BGN. The XRD result (Figure 3) shows the formation of α-quartz phase in BGN. Such a partial crystallization should not eliminate the biodegradability and bioactivity of BGN.54 BGN maintain their bioactivity and biodegradability, though these properties could be inhibited.55 Nevertheless, the BGN coating promotes the bioactivity and can therefore also be expected to improve the osteogenesis and osseointegration of our scaffolds.

2.6. Osteoblast and Fibroblast Growth on the Composite Scaffolds. The proliferation and adhesion of rat embryonic fibroblast and pre-osteoblastic cells (MC3T3-E1) on CNTT–HA, CNTT–BGN, and CNTT–BGN/HA were assessed individually using fluorescence microscopy. As the CNTT scaffolds absorb light almost completely,56 optical imaging was restricted to approximately the first 200 μm from the surface. A long-term (21 days) in vitro study of MC3T3-E1 cells with our fabricated scaffolds revealed a surprisingly low number of dead cells in comparison to live cells (Figure 7) and that the scaffolds support osteoblast growth and adhesion. In addition, pre-osteoblasts appear to be well supported by all the scaffolds as well.

To investigate cell adhesion further, we also checked the adhesion of REFS2 cells stably transfected with YFP-paxillin.57 Paxillin is an essential part of the fibroblast adhesion machinery.58 After 4 days of cultivation, fibroblast cells had well-adhered on all scaffold types (CNTT–BGN, CNTT–HA, CNTT–BGN/HA) (Figure 8). Imaging the paxillin fluorescence from adhesion clusters shows tiny and blurry spots (Figure 8A3–F3). Actin fibers are arranged in a meshed-form fibroblast cytoskeleton rather than stress fibers on all of our scaffolds (Figure 8A1–F1). Such a mesh-like arrangement of actin is typical for fibroblast adhesion in three dimensions.50,59–61

To investigate cell morphology in our 3D scaffold materials, we studied cell shape and morphology via SEM (Figure 9). SEM micrographs reveal that the cells were stretched and spanned between filaments on all scaffolds. Interestingly, most of the fibroblast cells are stretched between two or three filaments, leading to large cell membrane extensions. A comparison of the SEM images with fluorescence images shows that the cells within the scaffolds are associated with scaffold microtubes and developed physical contact with scaffold filaments (Figure 9, right column), especially with the CNT matrix. In addition, the actin fiber meshes are elongated between microtubes of the scaffolds (Figure 8). This is in agreement with SEM images (Figure 9) of fibroblasts on all scaffold types, as membrane projections attach to the microtubes of the scaffold. Despite the ability of the fibroblasts to adhere to the scaffolds, an methylthiazolyldiphenyl-tetrazolium bromide (MTT) assay (Figure 10) revealed a 5–20% lower growth rate for fibroblasts compared to the negative control after 1 day of incubation. Fibroblasts and fibroblast growth factors have an ambivalent impact on bone growth and osteogenesis. On the one hand, fibroblast growth factors increase the proliferation of immature osteoblasts,62 whereas fibroblast growth factors cause apoptosis in differentiated osteoblasts and block mineralization,63–64 and fibroblasts inhibit biomineraled bone nodule formation.65 As fibroblasts adhere very well to our materials (Figures 8 and 9), we assume that our material is feasible for tissue engineering applications.

3. CONCLUSIONS

In summary, we have demonstrated a simple and efficient strategy to fabricate highly porous composite scaffolds made of self-entangled CNTs in microtube structures with incorporated bioactive ceramic nanoparticles (BGN and HA). These
nanoparticles have been chosen to promote bone tissue ingrowth. To prepare the scaffolds, we have employed a H₂ thermal reduction process to etch ZnO and sinter BG or HA nanoparticles in a single step. The preferential removal of the ZnO template by the H₂ etching/sintering process offers the opportunity of implementing a variety of ceramic nanoparticles into the highly porous 3D networks for the fabrication of diverse hybrid composite structures. Moreover, we have proven that the incorporation of BGN and HA nanoparticles leads to biomineralization of the scaffolds. Apart from its bioactivity, the great advantage of our scaffolds compared to other 3D carbon-based porous structures is that its stiffness and porosity...
can in principle be tuned. The hollow microtubes lead to a low density of the scaffolds, and thus to an increase in ion release and electrical conductivity per scaffold weight. In addition, the hollow microtubes can serve as channels for nutrient transport and in future studies might also be filled with drugs or growth factors. The spatial architecture of fibers in the fabricated structures provides large free space for cell adhesion and both osteoblasts and fibroblasts were capable of adhesion to the scaffolds and stretched out along the fibers. The bioactivity of the scaffolds together with the electrical conductivity of the CNTT backbone makes them promising candidates for applications where porous 3D architectures are essential for cell growth, stimulation, function, or biomineralization, e.g., in bone tissue engineering.

4. MATERIALS AND METHODS

4.1. Fabrication of Bioactive Scaffolds. The fabrication of the bioactive cell scaffold materials is based on infiltrating the bioactive material into a highly porous (~94%) ceramic sacrificial template. For fabricating this template, tetrapodal-shaped ZnO microparticles (t-ZnO) were prepared using a flame transport approach as described by Mishra et al.66 To interconnect the microparticles, the loose powder was pressed into a cylindrical cast (diameter: 12 mm; height: 2 mm) and sintered at around 1150 °C for 5 h to form a 3D template.27

BGN with a nominal composition of 70SiO₂−30CaO (in mol %) and a particle diameter of ~400 nm were synthesized by a sol−gel method as described by Zheng et al.40 Hydroxyapatite (HA) nanoparticles were bought from Sigma (nanopowder, <200 nm). CNTs were purchased from Carbobyk (CARBOBYK-9810). The zinc oxide (ZnO) templates were coated with such nanoparticles using an infiltration process recently described by Schütt et al. The ceramic nanoparticles (BGN and HA) with a concentration of 214.28 ± 11.9 mg/cm³ were dispersed in absolute ethanol (Sigma, Germany). Due to the high porosity (~94%) of the template and its superhydrophilic properties, nanoparticle dispersions are rapidly sucked into the template and a homogenous coverage is achieved after evaporation of the solvent. This process can be repeated several times to increase the amount of infiltrated nanoparticles. It is also possible to infiltrate different nanoparticle types in sequence so that a composite layer is formed around the template structure. Here, the templates were alternately infiltrated with a 1 wt % aqueous carbon nanotube (CNT) dispersion and a 4 wt % ceramic nanomaterial (HA or BGN) dispersion (in ethanol). The concentration of the ceramic nanoparticles was the same both for HA and BGN 214.28 ± 11.9 mg/cm³, whereas different CNT concentrations between 32.14 ± 1.78 and 75.00 ± 4.16 mg/cm³ were tested. After finishing the infiltration procedure, the ZnO template was removed by H₂ etching. To do so, the samples were placed in a sealed quartz tube furnace and the air was replaced by pure argon. The pressure was adjusted to 200 mbar. Subsequently, the temperature was increased to 900 °C. Then, an evaporator (170 °C) was used to decompose urea into ammonia (NH₃).67 At 900 °C ammonia decomposes to N and H₂, thereby etching the ZnO template, resulting in free-standing, fibrous composite structures of CNTs with bioactive nanoparticles. Depending on the content of the composite scaffolds, they are called CNTT−BGN (free-standing CNT networks with BGN), CNTT−HA (free-standing CNT networks with hydroxyapatite...
nanoparticles), CNTT−BG/HA (free-standing CNT networks with both BG and hydroxyapatite nanoparticles).

4.2. Characterization. The porosity of the scaffolds is obtained as follows: a defined amount of tetrapodal ZnO powder (here: 0.068 g) is pressed into a cylindrical shape (12 mm diameter; 6 mm height). The resulting volume is ∼0.226 cm³ and leads to a density of around 0.3 g/cm³. Since the bulk density of ZnO is 5.61 g/cm³, the porosity is ∼94% for the ZnO template. For the preparation of the bioactive scaffolds, a CNT dispersion of 1 wt % and a BG or HA dispersion of 4 wt % were used and infiltrated into the ZnO templates. The amount of infiltrated dispersion was adjusted to the amount, which was needed to completely fill the free volume of the template (∼230 μL).

The porosities of the final scaffold materials were calculated as follows: by adding 230 μL of a 1 wt % CNT dispersion, around 2.3 mg of CNTs are added to the network. Considering the density for CNTs to be 1.4 g/cm³ (which is already quite a high value for MWCNTs) the additional volume is around 0.0016 cm³ for each infiltration. By the addition of 230 μL HA or BG dispersion 9.2 mg are added. Considering the densities of BG and HA to be in the range of 2−6 g/cm³, we obtain an additional volume of 0.0015−0.0046 cm³. Therefore, in the case of a template that was infiltrated five times with a 1 wt % CNT dispersion and five times with a 4 wt % HA or BG dispersion, we obtain a porosity between 86 and 93% (depending on the density of HA, as given by the manufacturer). More information is shown in the Supporting Information (Table S1).

Electrical characterization was performed using a Keithley 6400 source-meter, which is controlled by a self-written LabView program, capable of measuring IV curves. Therefore, a sample holder was used in which the flat sides of the cylindrical samples (d = 6 mm, h = 3 mm) were connected to copper plates. To ensure a good contact between the copper plates and the sample surface, conductive silver paste was used at the interface. To avoid any electrical contact between the sample holder and the copper plates, Kapton tape was used as an insulator (Figure S2). The current was measured as a function of applied voltage (from −1 to +1 V using a step size of 0.1 V).

Mechanical characterization was performed with a self-built setup consisting of a Mähräuser Wetzlar HS 6.3 micromanipulator, which is driven by a stepper motor and a load cell (Burster präzisionsmes-technik GmbH & Co KG, type 8523-5050). A self-written LabView program is used to control all components and measure the force. To avoid any vibration damping, the whole setup is located on a very rigid aluminum plate in a box filled with sand, which is mounted on a vibration isolated table. For the compression tests, the samples were placed in between the micromanipulator and the load cell. The samples were compressed by 2000 μm with a rate of ∼66.67 μm/s and the force was measured using the load cell. Finally, the stress−strain curves are evaluated and the compressive strength was determined (Figure S3).

4.3. Protein Adsorption Rate. Bovine serum albumin (Pierce; Thermo Fisher, Germany) was used to test the protein adsorption capacity of the scaffolds. Quantification was carried out using bicinchoninic acid (BCA; Thermo Fisher, Germany) in calorimetric detection assay. For this, each scaffold type was immersed and incubated with 200 μL of bovine serum albumin (BSA) solution (1 mg/mL) for 4, 8, 12, 24, 48, and 72 h. Next, the supernatant was removed and mixed with 200 μL working reagent (Pierce; Thermo Fisher, Germany 23225). After 30 min of incubation, the protein concentration in the supernatant was quantified by a colorimetric microplate reader (BioTek uQuant) at 570 nm. Adsorption values were calculated by subtracting the measured protein concentration in the supernatant from the initial protein concentration in the protein solution before adding to the samples (1 mg/mL).

4.4. In Vitro Apatite Formation. The ability of the scaffolds to form apatite in vitro was evaluated by immersing the scaffolds into the simulated body fluid (SBF) according to the protocol proposed by Kokubo et al.43 Briefly, the scaffolds were incubated in the SBF at a ratio of 1 mg/mL at 37 °C and stirred at 90 rpm for up to 14 days. The SBF was replaced once every 3 days during the incubation period. At every predetermined time point, the scaffolds were removed from the SBF, gently rinsed with deionized water, dehydrated with acetone, and dried at 60 °C for 12 h. The apatite formation on the scaffolds was assessed by field emission scanning electron microscopy (FE-SEM; Auriga, Carl Zeiss) and energy dispersive X-ray spectroscopy (EDS; X-MaxN Oxford Instruments).

4.5. Quantification of Ion Release. The concentration of ions released from the scaffolds during incubation in PBS (Biochrom, Germany) at different time points was analyzed with inductively coupled plasma mass spectrometry (ICP-MS). In brief, 15 mg of scaffold material was placed in 1× PBS for 4, 8, 12, and 24 h at 37 °C as a short-term experiment. In a long-term experiment, the amount of released ions was measured every 3 days under the same conditions. For measuring ion release, PBS was extracted and the samples were immersed in fresh PBS. The extracts were acidified by 20 μL of nitric acid (Sigma, Germany). Acidified pure PBS was used as the control and the ion concentrations in pure PBS were subtracted from the ion concentrations measured in extractions to receive the ions released from the scaffolds.

4.6. Sample Preparation for Transmission Electron Microscopy (TEM). A tiny piece of a scaffold was crushed and dispersed in butanol prior to dropping small crumbs on TEM grids. Then, the micrographs of structures were taken by TEM (JEOL JEM-2100) at 200 kV.

4.7. Cell Adhesion Assays. 4.7.1. Fibroblasts. The scaffolds (CNTT−BG, CNTT−HA, CNTT−BG/HA) were autoclaved at 121 °C and afterwards soaked in DMEM supplemented (Biochrom, Germany) with 10% fetal bovine serum (FBS, Biochrom, Germany) and 1% penicillin/streptomycin (Sigma, Germany). Approximately 10 000 rat embryonic fibroblast cells were seeded on each sample. The cells were incubated on the scaffold for 4 days at 37 °C and 5% CO₂. Afterwards, they were fixated with paraformaldehyde (Thermo Fisher, Germany) and dried using critical point drying (EMS 3000). Prior to scanning electron microscopy (Ultra Plus Zeiss SEM, 5 kV), the cells were coated with a thin sputtered gold layer (Bal-Tec SCD 050, 30 mA, 30 s).

Cell morphology, adhesion, and cytoskeleton were investigated by fluorescently staining cell nuclei and stress fibers. Cell nuclei were stained with DAPI (Thermo Fisher, Germany), which binds to DNA, and stress fibers were stained with Phalloidin (Alexa Fluor 647 Phalloidin, Thermo Fisher, Germany), which binds to the F-actin of the
cytoskeleton. Images of the stained cells were recorded using a fluorescence microscope (Olympus IX81, camera: Hamamatsu, UV lamp: Lumencor).

4.7.2. Osteoblasts. The adhesion of osteoblast-like MC3T3-E1 to the scaffolds was visualized by detecting the filamentous actin of the cytoskeleton of cells on the scaffolds. Live/dead staining was carried out to assess the cytotoxicity of the scaffolds. In brief, the scaffolds were immersed into MEM (Biochrom, Germany) to stabilize the pH value prior to the seeding of cells on the scaffolds. After the pre-treatment, 0.2 mL of MC3T3-E1 cell suspension (2 × 10⁶ cells/mL) was added on the scaffolds (in 24-well plates). After 3 h of incubation, an additional 1.8 mL of MEM was added. The culture medium was changed every 2 days. To minimize the influence of cells adhering to the bottom surface of the well during cultivation, the scaffolds were placed into new wells of a 24-well plate when exchanging the medium. After 21 days of culture, cell adhesion on the scaffolds was visualized by staining. Cell nuclei were stained by 4,6-diamidino-2-phenylindole (DAPI, dilactate, Invitrogen), prepared by incubating the scaffolds at 37 °C for 30 min. Live/dead staining was carried out using calcein (Thermo Fisher, Germany) and propidium iodide (Thermo Fisher, Germany), according to the manufacturer’s protocol. Images of the fluorescently stained MC3T3-E1 cells were taken by a fluorescence microscope (Axio Scope A.1, Carl Zeiss Microimaging GmbH, Germany).

4.8. MTT Assay. The viability of fibroblast cells on the scaffolds was quantified according to the ISO 10993 norm. In brief, extractions were prepared by incubating the scaffolds at 37 °C in 1 mL culture medium (DMEM supplemented with 10% FBS and 1% penicillin/streptomycin) for 72 h. 10,000 REFS2wt cells were cultured with 100 μL of medium for 24 h. The supernatant was replaced with 100 μL of extraction medium, then the cells were incubated for 24 h at 37 °C again. The number of vital cells incubated with the extraction medium was measured by adding 50 μL of methylthiazolyltetrazolium bromide (MTT; Sigma-Aldrich, Germany) solution. After 4 h of incubation the absorbance was measured at 570 and 620 nm as a reference. Fresh and untreated culture medium was used as a negative control and medium containing 20% dimethyl sulfoxide as a positive control. The results were normalized to the absorbance measured in the controls. Five technical repeats were carried out in three independent experiments.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.8b13631. EDS results of the scaffolds after immersion in simulated body fluid, further data on the scaffolds (e.g., density, porosity) (PDF)

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*Notes*

The authors declare no competing financial interest.

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**ABBREVIATIONS**

- 3D, three-dimensional
- CNT, carbon nanotube
- ZnO, zinc oxide
- HA, hydroxyapatite
- BGN, bioactive glass nanoparticles
- BG, bioactive glass
- CaO, calcium oxide
- CNTT, hierarchical carbon nanotube tube networks
- SEM, scanning electron microscopy
- TEM, transmission electron microscopy
- XRD, X-ray diffraction
- CNTT–BGN, CNT-based tubes containing bioactive glass nanoparticles
- CNTT–HA, CNT-based tubes containing hydroxyapatite nanoparticles
- CNTT–BGN/HA, CNT-based tubes containing both bioactive glass and hydroxyapatite nanoparticles
- t-ZnO, tetrapodal zinc oxide
- BSA, bovine serum albumin
- SBF, simulated body fluid
- ICP-MS, inductively coupled plasma mass spectrometry
- PBS, phosphate buffered saline
- REFS2wt, rat embryonic fibroblasts S2 wild type

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