Preparation and Characterization of Nanofibrous Scaffolds of Ag/Vanadate Hydroxyapatite Encapsulated into Polycaprolactone: Morphology, Mechanical, and In Vitro Cells Adhesion

Hany El-Hamshary 1,2,*, Mehrez E. EL-Naggar 3,*, Ayman El-Faham 4,5, M. A. Abu-Saied 5,6, M. K. Ahmed 6,7 and Mosaed Al-Sahly 1

Abstract: Series of nanofibrous composites of polycaprolactone (PCL) were fabricated in different compositions of modified hydroxyapatite (HAP). The encapsulated HAP was co-doped with Ag/vanadate ions at different Ag contributions. XRD and FTIR techniques confirmed the powder and fibrous phase formation. Further, the morphological and mechanical behaviors of the electrospun nanofibrous scaffolds containing hydroxyapatite were investigated. The nanofibrous phases were biologically evaluated via studying contact angle, antibacterial, cell viability, and in vitro growth of human fibroblasts cell line (HFB4). It is obvious that silver ions cause gradual deviation in powder grains from wafer-like to cloudy grains. The maximum height of the roughness (R\text{t}) ranged from 308.3 to 442.8 nm, for the lowest and the highest additional Ag ions for powdered phases. Moreover, the highest contribution of silver through the nanofibrous phases leads to the formation of lowest filaments size ranged from 0.07 to 0.53 \mu m. Further, the fracture strength was increased exponentially from 2.51 ± 0.35 MPa at zero concentration of silver ions up to 4.23 ± 0.64 MPa at 0.6 Ag/V-HAP@PCL. The fibrous phases were biologically evaluated in terms of antibacterial, cell viability, and in vitro growth of human fibroblasts cell line (HFB4). The nanofibrous composition of 0.8 Ag/V-HAP@PCL reached the maximum potential against E. coli and S. aureus and recorded 20.3 ± 1.1 and 19.8 ± 1.2 mm, respectively. This significant performance of the antibacterial activity and cell viability of co-doped HAP distributed through PCL could recommend these compositions for more research in biological applications, including wound healing.

Keywords: hydroxyapatite; polycaprolactone; silver; antibacterial; wound healing

1. Introduction

Skin is a vital organ that protects the human body against outer pathogens [1–3], and thus it is a crucial part of the immune system [4–6]. Additionally, skin displays a key role in the human body metabolic cycle [7,8]. Skin tissues possess an ability to be self-integrated [9–12], however, a bacterial attack may retard this process [13–16]. Thus, the germicidal potential of biocomposites utilized in wound healing applications is a
must [17–19]. Biomaterials should possess surface morphology that offers both chemical and physical adhesion with surrounding tissues, as well as ionic species release to accelerate pathogen mortality [9,12,20]. Furthermore, mechanical properties and a screening of composites’ biodegradability are crucial to support tissue until healing occurs and avoid inflammation. There has been rapid progress in medical research interests in the last decades, such as designing appropriate wound dressing ingredients [6,21]. Multifunctional innovative nanomaterials exhibit great interest due to exhibiting antibacterial activity and nontoxic behavior, as well as tissue repairing potential. Biologically, hydroxyapatite (Ca_{10}(PO_{4})_{6}(OH)_{2}/HAP) is one of the most studied biomaterials, although it lacks antibacterial potential. HAP possesses excellent biocompatibility, high activity, in addition to significant wettability and biodegradability characteristics. HAP structure exhibits a high capability to perform ionic substitutions in different crystallographic sites such as phosphate and calcium sites [13]. This feature displays diverse functions within the metabolic processing and antibacterial potential. For instance, vanadium compounds have been suggested for special medical usages owing to their beneficial exploitation against dyslipidemia and diabetes. Additionally, the antibiotic potential of vanadium coordinated compounds [22,23], while vanadium should not exceed 0.01 × 10^{-3} M [24]. The V5+ ions are absorbed four times obtained with V4+ along the intestine canal [25,26]. Therefore, Oxo-vanadate anions’ insertion instead of phosphate ions through HAP crystal is recommended for biological uses due to its ability to act with body proteins [27].

Silver compounds are considered germicidal agents from ancient ages [1,28,29]. Also, Ag might be capable of conquest bacterial resistance phenomena [30]. Abdulsada et al. [31] have examined the antibacterial activity of the HAP Ag nanoparticles (AgNPs) coating against Gram-positive bacteria Escherichia coli (E. coli) and Staphylococcus aureus (S. aureus). Indicating HAP suspension without AgNPs in the case of E. coli reduced the growth of bacteria to a limited percentage. Thus, there is the adhesion of bacterial cells to HAP. While in the case of S. aureus bacteria, HAP suspension shows higher antibacterial potential than that of the E. coli bacteria. On the other hand, HAP/AgNPs suspension displays a higher germicidal effect owing to AgNPs active role in killing bacterial strains, while S. aureus growth was inhibited, reaching 99%. This could be explained by the weakness of Gram (+) bacteria cellular membranes. Thus, AgNPs easily penetrate the cellular walls and increase the bacterial death. These results matched with antimicrobial activity results of AgNPs by Mirzaee et al. [32].

Poly(ε-caprolactone) (PCL) polymer has been exploited in tissue engineering due to its degradability, biocompatibility [9,10], nontoxicity, great mechanical behavior, and its bioresorbability [33,34]. PCL offers simple processing at low temperature (60 °C), as well as a wide range of suitable solvents and viscoelastic features that simplify the fabrication procedure [9–12]. Biologically, a hydrophobic property of PCL leads to its lack of adhesion capability through living tissues, which can be modified partially via additional inorganic substances [35,36].

The electrospun technique offers porous designs to allow blood and nutrients to pass in case of biological usage [37–39]. The properties of the obtained nanofibers can be adjusted upon the variation of preparation conditions, including viscosity, concentrations, voltage, and spray rate. Furthermore, the increasing porosity ratio can encourage the biological applicability, however mechanical resistance might be deteriorated. Therefore, optimization of the fabricated composition should take into consideration both design and components [40]. Milovac et al. have investigated the effect of HAP coating with a PCL via the vacuum impregnation technique. The results showed that the mechanical behavior of (HAP/PCL) scaffolds was improved, as well as keeping porous structure, which is crucial for tissue growth and vascularization. The compressive strength reached around 0.88 MPa, and the elastic modulus was achieved (15.5 MPa) [41].

Regarding the previously mentioned data, combining these constituents could introduce an appropriate biomaterial for accelerating the wound healing process. This work aims to fabricate nanofibrous scaffolds of PCL incorporated with co-doped modified HAP
at different contributions of Ag ions. The structural, morphological, and mechanical behaviors of the obtained scaffolds will be inspected, in addition to the cellular response to the human fibroblast cell line in vitro.

2. Experimental Section

2.1. Materials

Silver nitrate (AgNO₃), calcium chloride dihydrate [CaCl₂·2H₂O], vanadium oxide (V₂O₅), diammonium hydrogen phosphate [(NH₄)₂HPO₄], and solvents including chloroform (purity = 99.5%) and methanol (purity > 99%) were bought from LOBA, Mumbai, India. Polycaprolactone (PCL, Mₘ = 80,000 g/mol) was purchased from Sigma-Aldrich Co. (St. Louis, MO, USA).

2.2. Series of (Ag/VO₃⁻)-HAP@PCL Preparation

Co-doped Ag/V into HAP compositions were prepared by the precipitation technique, starting with the preparation of calculated concentrations of raw materials, then mixing. The solutions of CaCl₂·2H₂O (0.5 M), V₂O₅ (0.02 M), and (NH₄)₂HPO₄ (0.28 M) were prepared separately by using double-distilled water containers (50 mL).

Then, V₂O₅ solution (0.02 M) was added gradually to the phosphate precursor beaker (around 1 mL/s). Then, the solution of Ca was added into the container of (P + V) while keeping the pH at 11, with stirring for 2 h.

Next, HAP co-substituted at different contents of Ag was synthesized as per the previous protocol, while the concentration of the added silver (Ag) was kept at 0.0 ≤ x ≤ 0.8 with a step of 0.02 on the account of Ca contents. Then, the solutions of (P + V) were added in dropwise manner to the container of (Ca + Ag). For each sample, the stirring step was held at 1200 rpm for 2 h, and afterwards, the solutions were aged for 24 h until precipitation. Following that, the gels were washed several times with double-distilled water. Finally, the gels were purified and dried for 6 h at 50–60 °C.

The viscous PCL gel was then obtained by adding 10 g of PCL pellets in a solvent mixture of chloroform (66.7 mL) and methanol (100 mL) (33.3 mL). The PCL nanofibers were produced from 110 mg of the dry product obtained. Afterwards, the composite samples were placed in a syringe pump and electrospun using a custom-made electrospinning system with input values for both samples.

2.3. Characterization

2.3.1. X-ray Diffraction (XRD) Measurements for the Prepared Samples

An X-ray diffractometer (XRD, analytical x’pertpro, The Netherlands) was used to conduct the analysis. The data was obtained between 4° ≤ 2θ ≤ 60°.

2.3.2. Fourier Transformed Infrared (FTIR) for the Detection of Newly Created Bands

An FTIR spectrometer (PerkinElmer 2000, Waltham, MA, USA) was utilized for the measurements in the scanned range of 4000–400 cm⁻¹.

2.3.3. Surface Topography Examination by Making Use of Field Emission Scanning Electron (FESEM)

The surface structure was detected via FESEM (model: QUANTA-FEG250; The Netherlands). The roughness behavior was determined using Gwydion 2.45 software [42].

2.3.4. Tensile Stress Behavior

The mechanical properties were obtained for all samples. Each sample was cut with a length of 8 cm to be pulled up to the fracture, whereas the strain was recorded using an apparatus of Omnitest-25, 5 kN (Mecmesin Co., West Sussex, UK), with a rate of 30 mm/min, upon a standard code ASTM D882.
2.3.5. Thermogravimetric Analysis (TGA)

In a DTG-60 SHIMADZU analyzer, TGA was performed at room temperature (RT) up to 600 °C with a heating rate of 10 °C/min.

2.3.6. In Vitro Ionic Release

To assess the release of Ag ions by simulated body fluid (SBF), a 0.25 g piece of 0.8 Ag/V-HAP@PCL nanofiber was immersed in 100 mL of SBF for 24 h and held at 37 °C, and 5 mL of SBF was taken to be analyzed using an inductively coupled plasma (ICP) apparatus after 4, 6, 8, 16, and 24 h (720 ICP-OES, Agilent Technologies, New York, NY, USA).

2.3.7. Cell Viability

The human fibroblasts cell line (HFB4) was cultured in Dulbecco’s modified Eagle’s medium (DMEM, Gibco) at 37 °C and 5% CO\textsubscript{2} to determine in vitro cell viability. The cells were seeded on the fibers at a density of 5 × 10\textsuperscript{3} cells/cm\textsuperscript{2} using 12-well plates. The medium was then extracted and MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) was injected into the wells after 3 days of incubation. The ratio of viable cells to control cells was used to determine cell viability using an optical analyzer [43,44]:

\[
\text{Viability (\%)} = \frac{\text{Mean optical density of test samples}}{\text{Mean optical density of the control}} \times 100
\]

A well injected with cells but no additional fibers serves as the control sample. To measure standard deviation, the procedure was repeated three times.

2.3.8. Cell Growth In Vitro

Cell growth on the scaffold was used to assess the novel nanofibers’ applicability, while FESEM was used to observe the growth activity of HFB4 cells seeded on the nanofibers. As a result, each sample was reaped into two 0.5 × 0.5 cm bits, which were then sterilized using UV. Every well was then filled with 1.5 mL of HFB4 cells. Finally, the plate was sealed and incubated for three days at 37 °C. Thereafter, the fibers were washed in phosphate buffered saline (PBS) and processed for FESEM analysis.

2.3.9. Antibacterial Effectiveness

The antibacterial experiments were performed against *Staphylococcus aureus* (*S. aureus*, American Type Culture Collection (ATCC) number 29213), and *Escherichia coli* (*E. coli*, ATCC number 25922). The experiments were carried out for 24 h; then, the measurements were taken via the inhibition zone. The inhibition zone indicates the sensitivity degree of bacteria towards the fibrous sample. The starting concentration was around 50 mg/mL.

3. Results and Discussion

3.1. XRD of Powder and Fiber Phases

As is obvious in Figure 1a, XRD analysis indicated that HAP was formed in a single-phase and hexagonal structure with a P\textsubscript{6}\textsubscript{3}/m space group, according to the ICDD card no. 01-0.73-0293. Figure 1a represents the powder phase of co-doped HAP with the different associations of Ag ions. The characteristic peaks of HAP appear clearly upon the five diffractograms. This confirms the formation of HAP powder phase formation in the pure phase. No secondary phases were detected with silver incorporation into the HAP structure matrix, which refers to the successful incorporation of silver ions into HAP crystals. This is expected for the low content of silver dopants. The crystallinity seems to be low upon the additional ions. The co-reprecipitation method is a simple route to fabricate structurally modified HAP. The synthesis was performed at room temperature; thus, the crystallinity of the obtained powders was low. It can be mentioned that the natural HAP possesses low crystallinity, which is accompanied by ionic dopants. Figure 1b exhibits the patterns of fiber phases of Ag/V-HAP@PCL with changing silver contributions. PCL
signifies the highest intensity, whilst Ag/V-HAP reflects the short peaks. Moreover, the peaks of semi-crystalline PCL polymer appear at 21.49° and 23.67°, which confirms the PCL existence.

Figure 1. (a,b) X-ray diffraction (XRD) pattern of HAP modified with co-doped Ag/V and incorporated into polycaprolactone nanofibers: (a) Ag/V-HAP in powder phase and (b) Ag/V-HAP@PCL in nanofibrous composition (*: HAP).
3.2. FTIR of Powder and Fiber Phases

FTIR spectra do not reveal clear deviations in IR spectra of HAP with increasing silver contents through HAP powder (Ag/V-HAP), as shown in Figure 1a. The bands centered at 560 and 590 cm\(^{-1}\) in Ag/V-HAP powder could be equivalent to (\(\nu_4\)) of the phosphate group \([45]\), as reported in Table 1. Moreover, the vibrational mode (\(\nu_3\)) of the phosphate group band appears at 870 cm\(^{-1}\) \([46,47]\). Further, the band at 1039 cm\(^{-1}\) is corresponding to a stretching mode \(\nu_3\) (belonging to the phosphate group) \([48]\). It is obvious that the intensity of bands almost approximately reflected its contribution. Lastly, the presence of water and hydroxyl group is confirmed by the broadband at 3441 cm\(^{-1}\). Figure 2a confirmed the formation of HAP. On the other hand, spectra of Ag/V-HAP@PCL show almost steady band intensities with changing silver concentration. The bands at 2858 and 2935 cm\(^{-1}\) described the stretching mode of C-H bond, whereas the band at 1734 cm\(^{-1}\) is appropriate to carbonyl group (C=O) \([49]\). Furthermore, characteristic bands of PCL are found at 1239 and 1279 cm\(^{-1}\), attributed to the stretching of C-O and the asymmetric stretching of C-O-C, respectively \([46,49]\). Additionally, powder phase bands were located in lower frequencies compared to fiber phase bands, as reported in Table 1. FTIR spectra have confirmed Ag/V-HAP and Ag/V-HAP@PCL formation. For the FTIR of powder spectra, silver containing compositions show broader bands than those of HA, thus silver dopant causes a decline in crystallinity \([31]\). Thus, FTIR results match with XRD, as shown in Figure 1a.

![FTIR Spectra of HAP modified with co-doped Ag/V and incorporated into polycaprolactone](image-url)

**Figure 2.** (a,b) Fourier Infra Red (FTIR) Spectra of HAP modified with co-doped Ag/V and incorporated into polycaprolactone: (a) Ag/V-HAP in powder phase and (b) Ag/V-HAP@PCL in nanofibrous composition.
Table 1. Characteristic FTIR bands of Ag/V-HAP and Ag/V-HAP@PCL at different silver ions concentrations.

| 0.0 Ag/V-HAP PCL | 0.0 Ag/V-HAP | 0.2 Ag/V-HAP@PCL | 0.4 Ag/V-HAP@PCL | 0.6 Ag/V-HAP@PCL | 0.8 Ag/V-HAP@PCL | Assignments | Ref |
|-------------------|--------------|------------------|------------------|------------------|------------------|-------------|-----|
| 560 | 563 | 560 | 563 | 560 | 560 | 560 | 560 | 560 | 560 | 560 | 560 | 560 | 560 | 560 | ν₄ PO₃⁻ | [45] |
| 580 | 727 | 580 | 727 | 580 | 727 | 580 | 727 | 580 | 727 | 580 | 727 | 580 | 727 | 580 | ν₄ PO₃⁻ | [45] |
| 870 | 949 | 870 | 949 | 870 | 949 | 870 | 949 | 870 | 949 | 870 | 949 | 870 | 949 | ν₃ PO₃⁻ | [46,47] |
| 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | 1033 | ν₃ PO₃⁻ | [46] |
| — | 1149 | — | 1149 | — | 1149 | — | 1149 | — | 1149 | — | 1149 | — | 1149 | C-O and C-C | [45] |
| — | 1181 | — | 1181 | — | 1181 | — | 1181 | — | 1181 | — | 1181 | — | 1181 | C-O-C | [45,49] |
| — | 1239 | — | 1239 | — | 1239 | — | 1239 | — | 1239 | — | 1239 | — | 1239 | C-O-C | [45] |
| — | 1279 | — | 1279 | — | 1279 | — | 1279 | — | 1279 | — | 1279 | — | 1279 | C=O-C | [45] |
| — | 1734 | — | 1734 | — | 1734 | — | 1734 | — | 1734 | — | 1734 | — | 1734 | C=O | [45] |
| — | 2858 | — | 2858 | — | 2858 | — | 2858 | — | 2858 | — | 2858 | — | 2858 | C-H | [50] |
| — | 2935 | — | 2935 | — | 2935 | — | 2935 | — | 2935 | — | 2935 | — | 2935 | C-H | [46] |
| 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | 3441 | O-H | [51,52] |

3.3. Surface Morphology of Powder Phase

Figure 3a–e offers detailed micrographs of the Ag/V-HAP surface structure. Figure 3a represents V-HAP composition with zero silver content. The surface is shown with aggregated nonporous behavior. The formed grains look like thin wafers arranged upon each other, with two classes according to their size: the small grains ranged from 0.11 to 0.27 μm, while the large ones started from 0.66 to 1.1 μm. Figure 3b shows deviations in the surface structure, which might indicate additional silver on the composition. The 0.2 Ag/V-HAP composition displays a porous surface with shallow and minor pores. The grain size is dropped, to start from 0.11 to 0.21 μm. It is obvious that with increasing silver percentage through the HAP structure, the surface morphology is changed gradually from a well-defined wafer appearance to a cloudy surface. Further, the cloudy surface offers a high surface area to the formed powder. Figure 3c shows the composition of 0.4 Ag/V-HAP with an intermediate appearance between the wafer and cloudy surface. The grain size ranged from 0.22 to 0.33 μm, while deep pores appeared in-between aggregated grains. The composition of 0.6 Ag/V-HAP is illustrated in Figure 3d with minor cloudy grains ranging from 0.05 to 0.27 μm. The composition with the highest additional Ag ions, which was formed with a cloudy surface, shallow pores, and average grain size of 0.41 μm.

Figure 3. FESEM micrographs of HAP modified with co-doped Ag/V in powder phase: (a) 0.0 Ag/V-HAP, (b) 0.2 Ag/V-HAP, (c) 0.4 Ag/V-HAP, (d) 0.6 Ag/V-HAP, (e) 0.8 Ag/V-HAP.
Figure 4a–e displays three-dimensional micrographs of Ag/V-HAP surface, demonstrating surface roughness deviation due to gradually increasing silver content. Several roughness parameters are reported for each composition and are mentioned in Table 2. Regarding the roughness average (Ra), it is obvious that the resultant data displays an increasing pattern, starting from 93.9 nm at free silver composition and reaching 129.1 nm in 0.8 Ag/V-HAP. The maximum height of the surface (Rt) and maximum roughness valley depth (Rv) represent an increasing pattern, whereas the silver ions’ insertion induced both crests and notches formation. The Rt ranged from 902.0 to 956.9 nm, while Rv ranged from 308.3 to 442.8 nm for the lowest and the highest Ag ions. Moreover, the surface roughness variation pointed to the change in powder composition, which leads to different responses to host media. Thus, silver ions promote biological performance where crests act as hooks that promote resultant adhesion force [53,54]. This clear increase of roughness parameters with Ag insertion points to the crystal defects’ formation. These defects enhance the interlocking process, which is supported by adhesion of the surface, while chemical adhesion often works by boosting the bonding between the two surfaces via increasing crystallographic defects [45]. The roughness property offers higher potential in biological applications in addition to promoting the ionic release property [53,54].

Table 2. Surface roughness parameters of Ag-V-HAP powder with different silver content. Parameters include the roughness average (Ra), root mean square (Rq), maximum height (Rt), maximum valley depth (Rv), maximum peak height (Rp), and average maximum height (Rtm).

| Composition     | Ra    | Rq    | Rt    | Rv    | Rp    | Rtm  |
|-----------------|-------|-------|-------|-------|-------|------|
| 0.0 Ag-V-HAP    | 78.49 | 146.1 | 902.0 | 308.3 | 529.7 | 349.0|
| 0.2 Ag-V-HAP    | 93.9  | 119.2 | 902.0 | 323.8 | 510.7 | 474.5|
| 0.4 Ag-V-HAP    | 120.5 | 94.31 | 941.2 | 391.2 | 557.2 | 443.1|
| 0.6 Ag-V-HAP    | 122.5 | 149.4 | 843.5 | 427.1 | 557.2 | 454.9|
| 0.8 Ag-V-HAP    | 129.1 | 160.2 | 956.9 | 442.8 | 617.3 | 513.7|

Figure 4. (a–e) Behavior of surface roughness of HAP doped with Ag/V in powder phase: (a) 0.0 Ag/V-HAP, (b) 0.2 Ag/V-HAP, (c) 0.4 Ag/V-HAP, (d) 0.6 Ag/V-HAP, (e) 0.8 Ag/V-HAP.
3.4. Transmission Electron Microscopy (TEM) of Nanofiber Composites

Figure 5a,b shows the microstructural behavior of the nanofibrous scaffolds of 0.4 Ag/V-HAP@PCL and 0.8 Ag/V-HAP@PCL, respectively. The micrographs show the good distribution of co-doped HAP composites upon the compositions with particle sizes around 3 to 8 nm. Figure 5a exhibits a one-grain aggregate with a diameter reaching 25 nm. Figure 5b shows less aggregation with increasing silver content; thus, higher surface area is offered.

Figure 5. (a,b) Transmission Electron Microscopy (TEM) micrographs of nanofibrous scaffolds of PCL containing HAP doped with co-ions (Ag/V): (a) 0.4 Ag/V-HAP@PCL, (b) 0.8 Ag/V-HAP@PCL.

3.5. Surface Morphology of Nanofiber Composites

FESEM micrographs for surface illustration of Ag/V-HAP@PCL nanofibrous scaffolds with changing Ag ions content are shown in Figure 6a–e. Figure 6a displays 0.0 Ag/V-HAP@PCL. The nanofibers could be described as wide cross-linked fibers divided into two groups, with thinner filaments having a wide range of pores size in-between. The size of larger filaments is from 1.14 to 1.42 µm, while the thinner ones are from 0.21 to 0.42 µm. The formed macrospores’ size begins from 1.42 to 2.81 µm. The thin fiber filaments are distributed deeply, while large filaments appeared strongly on the surface. Figure 6b exhibits a nanofibrous scaffold of 0.2 Ag/V-HAP@PCL with reduced width compared to the former. The thin filaments ranged from 0.21 to 0.33 µm, while the larger filaments recorded a range from 1.1 to 1.4 µm. The porous surface exhibits pores with a range of 1.61–3.57 µm. Figure 6c displays the homogeneous distribution of networked nanofibers with diameters in ranges of 0.1 to 0.21 and 0.21 to 0.31 µm. Further, the size of the pores ranged from 1.2 to 3.1 µm. Moreover, it is obvious that with increasing silver content, the fibers’ diameters become thinner and well-defined, while the filaments become cloudy, as shown in Figure 6d. The cloudy networked filaments’ diameters are categorized as large and thin, with ranges of 0.97–1.4 µm and 0.28–0.56 µm, respectively. The pores were sized from 1.6 to 1.3 µm. Figure 6e shows the highest silver contribution surface, which was formed with ranges of (0.07–0.53) and (0.71–0.97) µm, while minor pores are sized from 0.28 to 1.14 µm.

The surface roughness of Ag/V-HAP@PCL fiber is screened with increasing silver content, as shown in Figure 7a–e. The Ra demonstrates an upward growing pattern, from 131.8 to 228.6 nm in the composition of the highest silver contribution. Both parameters of Rt and Rv display data confirming the positive influence of silver addition on surface roughness. The Rt parameter of 0.0 Ag/V-HAP@PCL composition recorded 947.4 nm, while it reached 987.3 nm for 0.8 Ag/V-HAP@PCL. The Rv increased from 414.9 to 452.0 nm. Therefore, additional Ag ions tend to promote both notches and crests.
Figure 6. (a–e) FESEM micrographs of PCL scaffolds encapsulated with co-doped Ag/V-HAP in nanofibrous phase: (a) 0.0 Ag/V-HAP@PCL, (b) 0.2 Ag/V-HAP@PCL, (c) 0.4 Ag/V-HAP@PCL, (d) 0.6 Ag/V-HAP@PCL, (e) 0.8 Ag/V-HAP@PCL, (f) fibers’ diameter distribution of 0.8 Ag/V-HAP@PCL.

Figure 7. (a–e) Surface Roughness of PCL scaffolds encapsulated with Ag/V-HAP at different additional Ag ions: (a) 0.0 Ag/V-HAP@PCL, (b) 0.2 Ag/V-HAP@PCL, (c) 0.4 Ag/V-HAP@PCL, (d) 0.6 Ag/V-HAP@PCL, (e) 0.8 Ag/V-HAP@PCL.
Furthermore, a rough surface is more applicable than smooth in the biological field. The rough surfaces support polymeric surface capability to adhere to the surrounding environment in two prospective ways: physical and chemical adhesion [52]. Consequently, controlling silver dopants through PCL nanofibers could introduce an effective film to attach with biological environments [55]. Further, many parameters could recompense for the weak roughness, for instance, porosity that enhances nurturing of cells, as well as chemical and mechanical adherence between graft composition and the host tissue [56].

3.6. Mechanical Properties

Nanofibers’ capability to stand for a suitable time in the biological environment hinge on their mechanical behavior and fiber morphology, including fiber alignment and porosity. The effect of ionic modification and its concentration is demonstrated through plotting strain against tensile stress. Silver insertion supported tensile strength to be developed from 3.74 ± 0.32 MPa to 5.10 ± 0.37, 5.65 ± 0.41, 6.51 ± 0.49, and 4.00 ± 0.23 MPa for a gradual increase in silver content, respectively. Moreover, the growth in tensile strength is controlled with the reduction in porosity. Consequently, the chemical composition is a crucial factor to estimate mechanical behavior [57–60]. The free silver content composition exhibits the lowest toughness, which is 2.09 ± 0.23 MJ/m$^3$, while 0.6 Ag/V-HAP@PCL scaffold required the highest energy to break by 3.43 ± 0.42 MJ/m$^3$. On the other hand, the fracture strength increased exponentially from 2.51 ± 0.35 MPa at zero concentration of silver ions up to 4.23 ± 0.64 MPa at 0.8 Ag/V-HAP@PCL (Figure 8).

![Figure 8](image_url)  
**Figure 8.** Mechanical properties of PCL scaffolds HAP-modified with co-doped Ag/V and incorporated into polycaprolactone.

Moreover, the maximum strain at the break started from 68.70% ± 4.12% in 0.0 Ag/V-HAP@PCL composition and dropped with silver ions’ insertion as shown in Figure 8 and Table 3. Further, the mechanical properties depend on fiber alignment and distribution. Thus, 0.6 Ag/V-HAP@PCL nanofiber represents the highest tensile strength, which refers to the fiber’s good alignment compared to the other scaffolds. Additionally, the deterioration of the tensile value of 0.8 Ag/V-HAP@PCL could be referred to as the higher ratio of porosity in comparison to 0.6 Ag/V-HAP@PCL. Furthermore, Shkarina et al., has compared tensile strength of HAP containing silicate/PCL nanofibers with parallel and random alignment of fiber filaments, and the result was 5.83 ± 1.34 MPa for well-aligned samples and 1.67 ± 0.1 MPa for random alignment [61]. Additionally, the existence of HAP within
PCL nanofibers induces mechanical properties, as well as growth enhancement of cells owing to its chemical structure that boosts the formation of direct chemical bonds with surrounding tissues.

Table 3. The mechanical properties of Ag/V-HAP@PCL at different contents of Ag, including Young’s modulus, tensile strength (MPa), fracture toughness, fracture strength, and maximum strain before the break.

|                | Young’s Modulus (MPa) | Tensile Strength (MPa) | Fracture Strength (MPa) | Maximum Strain at the Break (%) | Toughness (MJ/m³) |
|----------------|-----------------------|------------------------|-------------------------|----------------------------------|-------------------|
| 0.0 Ag-V-HAP@PCL | 0.37 ± 0.12           | 3.74 ± 0.32            | 2.51 ± 0.35             | 68.70 ± 4.12                    | 2.09 ± 0.23       |
| 0.2 Ag-V-HAP@PCL | 0.60 ± 0.17           | 5.10 ± 0.37            | 0.77 ± 0.12             | 67.60 ± 2.12                    | 2.18 ± 0.54       |
| 0.4 Ag-V-HAP@PCL | 0.97 ± 0.22           | 5.65 ± 0.41            | 0.00 ± 0.11             | 67.34 ± 1.92                    | 3.19 ± 0.17       |
| 0.6 Ag-V-HAP@PCL | 0.54 ± 0.10           | 6.51 ± 0.49            | 1.70 ± 0.21             | 67.69 ± 3.32                    | 3.43 ± 0.42       |
| 0.8 Ag-V-HAP@PCL | 0.44 ± 0.19           | 4.00 ± 0.23            | 4.23 ± 0.64             | 67.34 ± 3.72                    | 2.39 ± 0.11       |

3.7. Contact Angle

The contact angle reflects the fiber’s capability to adhere physically or chemically to living tissues. The 0.8 Ag/V-HAP@PCL scaffold displays the narrowest angle, with 83°, while contact angles show an increasing pattern with decreasing silver content, as shown in Figure 9. The rest of the scaffolds display 105°, 100°, 93°, and 87° from free silver composition to 0.6 Ag/V-HAP@PCL, respectively. Thus, the nanofibrous scaffold of 0.8 Ag/V-HAP@PCL represents the higher applicability. Ahmed et al. have studied the hydrophobic nature of pristine PCL, trying to modify with the additive Ag-MNPs. The contact angle exhibited a declining trend, from 105.4 ± 4.5° to 96.3 ± 5.2° to 88.5 ± 4.1° for 0.0 Ag-MNPs@PCL, 0.1 Ag-MNPs@PCL, and 0.2 Ag-MNPs@PCL, respectively. They reported that this change in the contact angle with the Ag content is in accordance with the increased surface roughness with Ag concentration [62]. Thus, the contact angle of 0.8 Ag/V-HAP@PCL nanofibrous compositions is in accordance with previous roughness data in Table 4.

Figure 9. Contact angle of nanofibrous scaffolds of PCL encapsulated with Ag/V-HAP at different additions of Ag ions; (a) 0.0 Ag/V-HAP@PCL, (b) 0.2 Ag/V-HAP@PCL, (c) 0.4 Ag/V-HAP@PCL, (d) 0.6 Ag/V-HAP@PCL, (e) 0.8 Ag/V-HAP@PCL.
Table 4. Surface roughness parameters of Ag-V-HAP@PCL nanofiber with different silver content.

| Composition       | $R_a$ (nm) | $R_q$ (nm) | $R_t$ (nm) | $R_v$ (nm) | $R_p$ (nm) | $R_{tm}$ (nm) |
|-------------------|------------|------------|------------|------------|------------|---------------|
| 0.0 Ag-V-HAP@PCL  | 131.8      | 162.2      | 947.4      | 414.9      | 548.0      | 451.0         |
| 0.2 Ag-V-HAP@PCL  | 153.5      | 187.8      | 964.3      | 433.4      | 562.4      | 443.1         |
| 0.4 Ag-V-HAP@PCL  | 168.5      | 203.2      | 969.1      | 441.2      | 558.8      | 454.9         |
| 0.6 Ag-V-HAP@PCL  | 216.2      | 259.7      | 975.6      | 437.6      | 566.6      | 466.7         |
| 0.8 Ag-V-HAP@PCL  | 228.6      | 267.9      | 987.3      | 452.0      | 585.1      | 423.5         |

3.8. TGA Analysis

Thermal stability plays a crucial role in estimating the optimum range of temperature at which the scaffolds can be effective. Thus, the thermogravimetric analysis has been done for the scaffolds starting from room temperature to 600 °C. As is obvious in Figure 10a,b, the behavior of the weight loss can be divided into three stages: room temperature to 320 °C, 320–510 °C, and ≥510 °C. In the first stage, the fibers seem to be thermally stable, while in the second stage, the fibers start to degrade exponentially. The higher concentrations of co-doped Ag/V tend to be faster to degenerate, which indicates the high disorder of their compositions compared to the lower ones. Above 510 °C, the samples seem to be fully degenerated.

![TGA Analysis](image1)

Figure 10. Thermogravimetric analysis of the nanofibers of HAP@PCL at different concentrations of doped ions of Ag/V, including (a) TGA and (b) DTA.

3.9. Ionic Release from Nanofibers

The degradation of nanofibers through the biological environment is an essential property to evaluate the applicability of the fabricated scaffolds. The biodegradation rate of the compositions depends strongly on the topographical features, as well as the structural stacking. The low regularity of stacked crystals indicates high crystal defects and a high affinity to interact with molecules from the ambient atmosphere. Therefore, the release of
Ag ions through SBF has been examined. As is obvious in Figure 11, the release reached around 5 ± 0.8 ppm after 4 h of immersion and increased exponentially, reaching about 17 ± 1.2 ppm after 24 h. This quantity might be sufficient to inhibit bacterial growth.

![Graph of Ag+ release from nanofibrous scaffolds of PCL containing modified HAP at 0.8 Ag/V-HAP@PCL concentration.](image)

**Figure 11.** Ionic release of Ag⁺ from the nanofibrous scaffolds of PCL containing modified HAP at the concentration of 0.8 Ag/V-HAP@PCL.

### 3.10. Cell Viability

Regarding the cytotoxicity of Ag/V-HAP@PCL fiber, the free silver composition reached excellent performance with 89.4% ± 3%. The 0.2 Ag/V-HAP@PCL composition recorded 91.4% ± 3%, while 0.4 Ag/V-HAP@PCL and 0.8 Ag/V-HAP@PCL composites recorded 94.6% ± 4% and 96.5% ± 5% respectively, as seen in Figure 12. The 0.6 Ag/V-HAP@PCL composition represents the highest cell viability potential, with 97.2% ± 2%. The non-cytotoxicity property persuades cell proliferation and enhances wound healing. Generally, surface roughness, type, and amount of composition constituents are crucial in bio-applicability potential; therefore, they are the tools to manipulate physical and chemical characteristics of the prepared nanofibrous composite [63]. Consequently, silver insertion boosts the healing process indirectly by enhancing the adhesion capability of the formed nanofibrous composition upon the surface of living tissues.

![Graph of cell viability of scaffolds of PCL embedded with Ag/V-HAP through human fibroblast cell lines at different Ag contributions.](image)

**Figure 12.** Cell viability of scaffolds of PCL embedded with Ag/V-HAP through human fibroblast cell lines at different Ag contributions.
3.11. Antibacterial Properties

Microbiological studies that evaluate the antibacterial potential, assessing germicidal effectiveness against *E. coli* and *S. aureus*, were performed. The first scaffold is for zero silver content composition. The recorded inhibition area against *E. coli* is \(12.7 \pm 1.1\) mm, while against *S. aureus* is \(12.9 \pm 0.9\) mm. The antibacterial screening shows an upward growing pattern with silver content for both screened organisms, as is obvious in Figure 13. Regarding germicidal effectiveness against *E. coli*, inhibition zones’ diameters are \(14.6 \pm 1.5\), \(16.1 \pm 0.9\), \(19.2 \pm 1.3\), and \(20.3 \pm 1.1\) mm for 0.2 Ag/V-HAP@PCL, 0.4 Ag/V-HAP@PCL, 0.6 Ag/V-HAP@PCL, and 0.8 Ag/V-HAP@PCL composites, respectively. Moreover, the 0.8 Ag/V-HAP@PCL scaffold of nanofibrous composition reached the maximum potential against both organisms, against *S. aureus* recording about \(19.8 \pm 1.2\), while the rest of the scaffolds, from 0.0 Ag/V-HAP@PCL to 0.6 Ag/V-HAP@PCL, recorded \(12.9 \pm 0.9\), \(13.9 \pm 1.2\), \(16 \pm 0.9\), and \(18.9 \pm 1.1\) mm, respectively. Roughness property, composite constituents (type and amount), and porosity control the bio-applicability of the prepared material; thus, the previous physical and chemical characteristics are in accordance with the resultant cell viability. The previous results have explained silver effectiveness by its capability to inhibit bacterial replicating capacity, causing bacterial mortality due to the deep penetration of nanoparticles into the cell wall, causing failure to interact with compounds containing phosphorus and sulfur, such as protein DNA. The increasing pattern could be explained by silver contribution, which boosts intracellular reactive oxygen species (ROS) release, thus enhancing bacterial cell wall penetration [63]. Moreover, silver ions bound to the cell wall, causing cell wall permeability malfunction and failure in bacterial cell respiration [33]. Thus, silver insertion boosts the healing process indirectly by eliminating the bacterial attack retarding effect.

![Figure 13. Antibacterial activity of HAP modified with co-doped Ag/V and incorporated into poly-caprolactone.](image)

3.12. Cell Attachment

Figure 14a–e has screened the surface adhesion and proliferation potential of prepared nanofibrous scaffolds for human fibroblasts cell lines (HFB4) in vitro. The evaluation was executed after three days of cultivation in vitro. Free silver scaffolds represent significant proliferation potential, whereas a wide Island of cells appeared with a diameter of around 6.2 \(\mu\)m. The cell proliferation is spread upon the fiber surface, leaving the pores. Silver addition thickened the cell island and improved proliferation upon the surface, while the cellular islands appeared wider with dimensions around 7.3 \(\mu\)m, as shown in Figure 14b.
Figure 14. Cell growth of PCL nanofibrous scaffolds encapsulated with HAP co-doped with Ag/V after three days of cultivation media of human fibroblasts cell lines in vitro: (a) 0.0 Ag/V-HAP@PCL, (b) 0.2 Ag/V-HAP@PCL, (c) 0.4 Ag/V-HAP@PCL, (d) 0.6 Ag/V-HAP@PCL, (e) 0.8 Ag/V-HAP@PCL.

Additionally, Figure 14c represents the 0.4 Ag/V-HAP@PCL scaffold, whereas the cell proliferations deeply fill a high percentage of pores, and the nanofibers were almost covered. Moreover, the nanofibers of 0.6 Ag/V-HAP@PCL seem to be filled with growing cells, while the 0.8 Ag/V-HAP@PCL leads to coverage of about 80% of fiber by spreading of HFB4 cells upon it. Thus, the resultant micrographs reported a clear adhesion potential owing to a hydrophilic property of inorganic silver ions, as well as the fibrous scaffold, which exhibited the capacity of protein adsorption [64]. Therefore, adjusting ingredients’ amounts and design could offer a scaffold with appropriate performances for the biological applications.

4. Conclusions

Nanofibrous scaffolds based on PCL were embedded with co-doped HAP using the electrospinning technique. The HAP was modified with different concentrations of silver/vanadate. The scaffold of 0.8 Ag/V-HAP@PCL nanofibers represented the lowest fiber diameters, from 0.07 to 0.53 µm. Additionally, 0.0 Ag/V-HAP@PCL nanofibers exhibited the lowest toughness, with 2.09 ± 0.23 MJ/m³, while the 0.6 Ag/V-HAP@PCL scaffold recorded 3.43 ± 0.42 MJ/m³. The antibacterial effectiveness indicated that increasing silver contribution enlarged the inhibition zone, reaching around 20.3 ± 1.1 and 19.8 ± 1.2 mm against E. coli and S. aureus, respectively. The cell viability of silver-containing nanofibrous material exceeded 90%, while the 0.6 Ag/V-HAP@PCL scaffold offered the highest non-cytotoxicity potential, with 97.2 ± 2. Results of adjusting ingredients’ amounts offered a nanofibrous substance with applicable performances for wound healing purposes. The 0.6 Ag/V-HAP@PCL nanofibers offered cell proliferation potential that leads to almost full coverage of fiber, while the 0.8 Ag/V-HAP@PCL composite leads to coverage of about 80% of fiber. The optimizing response of fibroblasts cells throughout the scaffolds encourages clinical investigation to be followed-up for these types of composition.
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