Therapeutic Applications of Antimicrobial Silver-Based Biomaterials in Dentistry

Qiyu Wang1, Yu Zhang1, Qiang Li2,*, Li Chen1, Hui Liu1, Meng Ding2, Heng Dong1*, Yongbin Mou1

1Department of Oral Implantology, Nanjing Stomatological Hospital, Medical School of Nanjing University, Nanjing, People’s Republic of China; 2Central Laboratory, Nanjing Stomatological Hospital, Medical School of Nanjing University, Nanjing, People’s Republic of China

*These authors contributed equally to this work

Correspondence: Yongbin Mou; Heng Dong, Department of Oral Implantology, Nanjing Stomatological Hospital, Medical School of Nanjing University, #30 Zhongyang Road, Nanjing, 210008, People’s Republic of China, Tel +86 25 83620236, Fax +86 25 83620202, Email yongbinmou@163.com; dongheng90@sinau.edu.cn

Abstract: Microbial infection accounts for many dental diseases and treatment failure. Therefore, the antibacterial properties of dental biomaterials are of great importance to the long-term results of treatment. Silver-based biomaterials (AgBMs) have been widely researched as antimicrobial materials with high efficiency and relatively low toxicity. AgBMs have a broad spectrum of antimicrobial properties, including penetration of microbial cell membranes, damage to genetic material, contact killing, and dysfunction of bacterial proteins and enzymes. In particular, advances in nanotechnology have improved the application value of AgBMs. Hence, in many subspecialties of dentistry, AgBMs have been researched and employed, such as caries arresting or prevention, root canal sterilization, periodontal plaque inhibition, additives in dentures, coating of implants and anti-inflammatory material in oral and maxillofacial surgery. This paper aims to provide an overview of the application approaches of AgBMs in dentistry and present better guidance for oral antimicrobial therapy via the development of AgBMs.

Keywords: silver, antimicrobial, dentistry, oral biofilm formation, biomaterials, nanotechnology

Introduction

Tissue surfaces of the oral cavity provide an ideal environment for attachment and formation of plaque biofilms, which are a crucial cause of dental caries, gingivitis or periodontitis, denture stomatitis, and other dental diseases. These bacterial or fungal communities cause infections by protecting pathogenic microbes from medical treatment and evading host defense responses. Dental plaque biofilms provide a reservoir for the growth and spread of multidrug-resistant bacteria and pose threats to immunocompromised patients. Despite the clinical use of antibiotics and other antimicrobial drugs, multidrug resistance remains an intractable problem.

The antibacterial properties of silver have been known since ancient times. Silver-based biomaterials (AgBMs) have been employed in several areas of dentistry due to their broad-spectrum antimicrobial properties (Figure 1). AgBMs include various forms of materials, such as traditional amalgam silver, silver-citrate, silver diamine fluoride, colloidal silver, and silver-based nanoparticles (AgNPs). AgBMs have been proven to be effective against most oral pathogens, including Streptococcus mutans (S. mutans), Streptococcus mitis (S. mitis), Streptococcus oralis (S. oralis), Streptococcus sanguis (S. sanguis), Porphyromonas gingivalis (P. gingivalis), Staphylococcus aureus (S. aureus), Candida albicans (C. albicans), Escherichia faecalis (E. faecalis) and Escherichia coli (E. coli). The effective bactericidal property of AgBMs arrests the attachment and formation of biofilms, which is essential for the application of dental materials.

The advancement of nanotechnology brings tremendous amounts of new materials with unique properties. Among all kinds of AgBMs, AgNPs have received the most attention for their outstanding antimicrobial properties against bacteria, fungi, and enveloped viruses. AgNPs are defined as elemental silver particles in size at a nanometer scale (1 nm to 100 nm). Compared to the particles in traditional colloidal silver, AgNPs are more consistent in size and shape, which means...
consistency in properties. The inner core of most AgNPs is metallic silver and the outer shell is surface oxide or sulfide layers. Silver ions are released in the process of the oxidation and dissolution of the surface layer of Ag$_2$O. The existing synthesis methods can be divided into “top-down” and “bottom-up” approaches, namely, the “cutting” process from bulk silver to nanosize or synthesis from molecular, atomic, and ionic components.

At the nanoscale, most of the atoms of nanomaterials are surface atoms, which dramatically alters their properties. As a kind of nanomaterial, AgNPs exhibit unique properties. Featuring their ultrasmall size, large surface area to mass ratio, and enhanced chemical reactions, AgNPs have broad prospects in antimicrobial therapy. Thus, AgNPs have been proven to be powerful weapons against multidrug-resistant bacteria. Additionally, decreased size improves wear resistance, which is valuable in dentistry. Furthermore, an increased surface/volume ratio results in higher surface reactions and stronger bond strength of mechanical interlock with the polymer matrix. Besides, AgNPs could improve the aesthetics of dental materials. The diameter of nanoparticles is smaller than the wavelength of visible light (400 nm to 800 nm), which means that more transparent dental materials can be made.

In this review, we systematically discuss the advancement and application of antimicrobial AgBMs in dentistry with emphasis on the mechanism, biosecurity and clinical application.

### Characteristics and Performance of AgBMs

#### Category of AgBMs

As a promising material in the antibacterial field, AgNPs have been extensively studied. According to chemical properties, AgBMs can be divided into metallic silver, silver salt (eg, AgNO$_3$ and silver sulfadiazine), silver oxides (eg, Ag$_2$O) and composite materials (eg, silver zeolite). According to the size scale, AgBMs can be divided into bulk materials, traditional colloids and AgNPs. More specifically, AgNPs can be divided into metallic AgNPs and their ionic...
Ionic AgNPs are nanoparticles of insoluble silver compounds, such as silver bromide (AgBr NPs), silver vanadate (AgVO₃ NPs) and silver oxide (Ag₂O NPs).¹⁵–¹⁷

**Antimicrobial Mechanisms of AgBMs**

The antibacterial mechanisms of silver, especially AgNPs, has been a major issue in numerous scientific publications. AgBMs have similar killing mechanisms on bacteria and fungi.¹³,¹⁸ However, the exact molecular mechanisms of action remain unclear. The potential mechanisms of the antimicrobial effect of AgBMs are described below (Figure 2).

AgBMs exert suppression on bacterial metabolism. The common mechanism of AgBM is attributed to the gradual release of silver ions (Ag⁺, Ag²⁺, Ag³⁺) in the majority of publications.¹⁹ Furthermore, it has been reported that silver ions have an affinity for multiple functional groups, such as phosphate, carboxyl, amino, and thiol groups, which disturbs those biological functions. More specifically, silver ions first affect the function of proteins and enzymes. Yamanaka et al confirmed that the 30S subunit and succinyl-CoA synthase of the ribosome were particularly inhibited by silver in prokaryotic cells such as *E. coli*. The former participates in the decoding of mRNA, while the latter is an important part of the glucose pathway and the tricarboxylic acid cycle (TCA cycle), which is crucial for the survival of cells.²⁰ Thus, the interruption of adenosine triphosphate (ATP) finally results in bacterial death. Gordon et al suggested that thiol groups (e.g., glutathione and cysteine) are the main targets that determine the antimicrobial properties of AgBMs. Their theory was confirmed because the toxicity of AgNO₃ to succinate dehydrogenase (from the respiratory chain) and aconitase (from the TCA cycle) could be alleviated by the addition of glutathione.²¹ Therefore, the bactericidal mechanism of AgNPs is regarded as a “ion carrier” through the cytomembrane. Accordingly, the size of AgNPs is designed to be as small as possible because a larger surface-to-volume ratio means more efficient release of silver ions, namely, greater antimicrobial activity.²²–²⁴ In addition, silver ions adhere to ribosomes and inhibit protein synthesis by disturbing their conformation after entering the cytoplasm.²⁵

---

**Figure 2** Antimicrobial and Dysfunction Mechanisms of AgBMs. Antimicrobial mechanisms of AgBMs (yellow frame) include destruction of the cell wall and cytoplasmic membrane, interruption of ATP production, production of destructive chemicals, interference with DNA replication and denaturation of ribosomes. The dysfunction mechanisms of AgBMs (blue frame) include efflux pump draining Ag⁺, flagellin aggregating AgNPs, coordinating the binding of histidine residues to Ag⁺, peptidoglycan membrane barrier, and biofilm barrier.

**Abbreviations:** ATP, adenosine triphosphate; DNA, deoxyribonucleic acid; ROS, reactive oxygen species; PDT, photodynamic therapy.
AgBMs cause the accumulation of chemicals that damage cellular components, such as proteins, DNA and lipids. The production of excessive reactive oxygen species (ROS) induced by silver ions has long been observed and regarded as an important bactericidal mechanism. The specific mechanism of how silver ions ignite the Fenton reaction remains controversial. It has been proposed that the exhaustion of thiol groups by silver ions leads to the accumulation of ROS because thiol groups in glutathione and cysteine are essential in the homeostasis antioxidant system. Interestingly, the release of silver ions is not the only method by which AgNPs ignite the production of ROS. AgNPs and silver nanoclusters (AgNCs) are also employed in photodynamic therapy (PDT) as promising photosensitizers. Photosensitizers produce ROS, such as hydroxyl radicals, superoxides, or singlet oxygen ($^1O_2$), when they are irradiated by light whose wavelength matches them. In the area of sterilization, common photosensitizers are limited by their cytotoxicity to mammalian cells. With the advancement of nanotechnology, AgNPs serve as a novel photosensitizer for the following reasons: (1) little cytotoxicity to mammalian cells, (2) intracellularly induced photosensitivity, and (3) silver ions released simultaneously. Therefore, AgNPs have incomparable advantages over traditional photosensitizers.

AgBMs damage the integrity of the cell membrane due to their physical properties. First, silver ions can cause depolarization of the cellular membrane due to electrostatic interactions with membrane phospholipids and proteins, resulting in destabilization of the cytomembrane. Notably, AgNPs have unique antibacterial mechanisms that are totally different from those of other AgBMs. Direct contact with uncharged nanoparticles can also rupture the bacterial cell membrane as a result of the physiochemical features of nanoparticles, such as surface topology, size, geometry and hydrophobicity, regardless of germ type. This “contact killing” mechanism has been employed to prolong antibacterial effects. For example, AgNPs immobilized on a material surface resist the formation of biofilms by rupturing the bacterial cell membrane and releasing silver ions simultaneously.

**Dysfunction Mechanism of AgBMs**

Unlike ordinary antibiotics, silver activates multiple mechanisms and hits different targets within bacterial cells. Therefore, it is difficult for these microorganisms to develop adaptive resistance. The only fully silver-resistant strain known at present is *P. stutzeri* which is found in silver mines. Interestingly, some antisilver mechanisms of bacteria have been described and summarized in Figure 2: (1) the energy-dependent efflux pump is the most efficient antisilver method of bacteria; the sil gene on the Salmonella pMG101 plasmid was identified to encode the silver ejection system; (2) flagellin, a globular protein in the flagellum, is considered to be an anti-silver agent of gram-negative bacteria. It is speculated that flagellin causes AgNPs to aggregate as an adhesive protein, thereby eliminating the antimicrobial effect of AgNPs. This mechanism involves phenotypic changes and can be eliminated by flagellin inhibitors easily. (3) bacterial can coordinate the binding of histidine residues to silver ions. Microbes are usually susceptible to AgBM because of the lack of thiol groups. However, *Salmonella* has been observed to coordinate the binding of histidine residues to silver ions, as an alternative to thiol groups; (4) gram-positive bacteria sequester silver in their peptidoglycan membrane, which explains why they are less sensitive than gram-negative bacteria; and (5) biofilms act as physical barriers. Its components, including nucleic acids, proteins, polysaccharides, and lipids, interact with AgBM through electrostatic interactions, hydrophobic interactions, hydrogen bond interactions, and van der Waals interactions, which reduces the antimicrobial activity of AgBM.

Additionally, the precipitation reaction of silver ions with halogen is also an important cause of inactivation. Given that saliva and gingival crevicular fluid in the mouth are rich in Cl$^-$, more researches are needed on the long-term antimicrobial effects of silver.

**Biosafety Considerations of AgBMs**

It has been reported that the toxicity of silver ions to bacteria is comparable to that of human cells. During application, AgBMs are exposed to digestive mucosa and bone tissue. Meanwhile, there is also a risk of inhalation during chairside extracorporeal polishing operations. Considering that dental AgBMs may continue functioning in the oral environment for decades, it is essential to lower their toxicity to human cells.

In regard to the specific types of AgBMs, soluble compounds are always more toxic than insoluble ones. For example, the LD$_{50}$ of AgNO$_3$ is estimated to be 28 mg/kg, while that of AgCl is over 10 g/kg. Notably, it is not convenient to
assess the toxicology of nanomaterials such as AgNPs with the traditional mass-only dose metric model. The cytotoxicity of AgNPs in mammalian cells depends on the nanoparticle size, shape, surface charge, dosage, oxidation state, agglomeration condition, and cell type. Generally, smaller AgNPs are more toxic due to the higher surface-to-volume ratio. Obviously, toxicity limits the application of certain types of AgNPs. For example, Ag$_2$O nanoparticles (Ag$_2$O NPs) exhibit high toxicity to eukaryotic cells. Therefore, they are employed as antitumor materials instead of bactericidal agents despite their high efficiency of killing bacteria and protozoa.

Generally, AgBMs are described as barely perceptible irritants because of the low concentration used in oral treatment. However, some atopic dermatitis can still be observed. There is no convincing research describing the absorption of silver in oral mucosa, while the absorption of silver into the blood has been demonstrated via ocular, uterine and nasal mucous membranes. Furthermore, silver can disrupt mitochondrial electron transport and induce apoptosis through an oxidative stress pathway, which has been demonstrated in liver and bronchial epithelial cells. Recordati et al observed slow clearance of silver and ultrastructural changes of the blood-brain barrier after oral administration of 1 mg/kg AgNPs and silver acetate in mice, without any treatment-related signs or effects on body and organ weights.

However, the application of AgBMs still shows some risks, such as argyria. Argyria is an infrequent, silver-related disease, that presents as blue-gray discoloration of the skin due to silver deposition. According to recent research, Argyria is attributed to infusible precipitates. It was reported that silver serum levels below 300 μg/L were never observed to be related to Argyria. In addition, occupation exposure could occur in clinics while manipulating materials such as polishing composite restoration with no water coolant, which results in aerosols—with concentrations above $1 \times 10^6$ particles/cm$^3$ inhaled by dentists and patients. It has been reported that rodents exposed to inhaled silver particle aerosols have pulmonary inflammation and even asthma at concentrations of 0.11–0.75 mg/m$^3$.

Compounds containing reduced thiol groups could significantly reduce the toxicity of silver ions. Given their limited size, bacterial cells contain far fewer low-molecular weight thiols than human cells, in addition to other antioxidant systems in human cells, such as cysteine-rich metallothionein. In vitro tests of silver toxicity are often closed systems where thiol content is much lower than that in the human body. Therefore, the toxicity of silver to human cells and its antimicrobial effect in vivo have been somewhat overestimated.

**Antimicrobial AgBMs Incorporated in Dental Materials**

**Preventing and Arresting Caries**

Silver materials have been used for combating caries in various forms. For instance, silver could be added to restorative materials in case secondary caries occurs. AgNPs can also be added to adhesives and acrylic resins coated onto the surfaces of orthodontic appliances to prevent dental demineralization caused by bacteria or accumulation. In addition, daubing- or gargling-related medicine in the oral cavity is another method utilizing the antimicrobial properties of silver ions.

**Dental Filling Material**

Early in 1985, the antimicrobial properties of silver amalgam were proven, as it gives off silver ions. An in vitro study carried out by Hegde et al showed that silver amalgam had a marked inhibitory effect better than glass ionomer cement (GIC) and composites. Later studies introduced a novel category of resins incorporated with AgBMs (Figure 3), which could exert extra antimicrobial effects, particularly against *S. mutans*. Jia et al synthesized Zn-Ag-loaded nano-SiO$_2$ specimens, which can be added into dental resin-based materials to enhance their antimicrobial properties. Kasraei et al reported that composite resins containing AgNPs exhibited antimicrobial activity against *S. mutans* and *Lactobacillus*. Suggested mass fractions vary based on the specific auxiliary material used, while consensus is that mass fractions larger than 1 wt% is not recommended due to the variation of mechanical properties and biocompatibility.

The addition of AgNPs (250 ppm) to primers could improve the antimicrobial effect, with no effect on bond strength or the biocompatibility of adhesives. Furthermore, the combination of AgNP and primer preserves the interface between tooth and adhesive, extending the utility time of dental restorations.
Orthodontic Appliance

Etching is an essential process before adhering orthodontic brackets onto the teeth, which results in potent caries and white spot lesions (WSLs). WSLs may develop since the beginning of the treatment. Silver is utilized as an additive in orthodontic adhesive and coating of orthodontic brackets. It has been reported that caries around orthodontic retainers could be prevented if AgNPs were incorporated into composite resin. Considering enamel remineralization, some researchers confirmed that orthodontic resin modified by a certain concentration of AgNPs (1 wt%) prevents WSL formation more efficiently (Figure 4). In addition, orthodontic brackets coated with titanium oxide mixed with Ag could significantly reduce the adhesion of S. mutans and P. gingivalis, and biofilm formation.

Caries Treatment

Inspired by the marked antimicrobial property of AgNPs, Haghgoo et al suggested that dental varnishes supplemented with AgNPs show good cariostatic efficacy. Silver diamine fluoride (SDF), approved by the US Food and Drug Administration (FDA), is a solution used for arresting caries or preventing potential lesions, and contains ionic silver, fluoride, and ammonia. It has been widely used in the prevention of early childhood caries (ECC) due to its effective, safe, painless, inexpensive and timely properties. SDF exhibited significantly higher caries arrest rates than fluoride varnish, GIC and cross tooth-brushing technique (CTT). It has been claimed that 38% SDF solution was the most effective dose for arresting advanced cavitated caries, ranging from moderate to high certainty, if applied biannually. However, the precipitation of silver byproducts in dental tissues stains the lesions and soft tissue black.

Although many studies have suggested that the stain on lesions and gingival tissue could fade gradually at the start of 6 months after treatment, the stain trace is a limitation for the use of SDF in the anterior aesthetic area. A double-blind clinical trial evaluated the cariostatic efficacy of 5% AgNP-incorporated sodium fluoride (NSSF) and 38% SDF in primary teeth, suggesting that they are equivalent in inhibiting dentine caries progression. NSSF is more economical and leaves no dark stains. Thus, NSSF has the potential to be an alternative to SDF owing to its antimicrobial effect and better aesthetic effect.
Bioactive glasses incorporated into dentifrices have proven the efficacy of treating demineralization and dental sensitivity because they can release therapeutic ions after dissolution, such as strontium (Sr$^{2+}$) and fluoride (F$^{-}$), to enhance remineralization.$^{84}$

It is suggested that the combination of AgNPs and PDT has good cariostatic efficacy. It could be a method that minimizes microbial residue after mechanical removal of caries. Unlike traditional antimicrobial agents, PDT based on photosensitizers activated by light sources is noninvasive, repeatable, nonresistant, and penetrable. When provided with the right energy density, it may kill microorganisms in a matter of minutes.$^{85}$

Applications of AgBM$s$ in Endodontic Treatment
In the area of endodontics, AgBM$s$ are applied to irrigation, medication, and addiction in sealer materials to improve their antimicrobial functions.$^{86}$

Irrigation
Conventional endodontic irrigants include chlorhexidine (CHX), ethylenediaminetetraacetic acid (EDTA) and sodium hypochlorite (NaClO). Their limitations are cytotoxicity to periapical tissue or inadequate removal of bacterial biofilms and smear layers.$^{87}$ It was reported that AgNPs can serve as active ingredients in root canal irrigation solutions.$^{88-90}$ Application of root canal irrigation solution with AgNPs resulted in superior antimicrobial effects (Figure 5A), reduced cytotoxicity and even enhanced fracture resistance of endodontically treated roots, compared to 2.5% sodium hypochlorite (Figure 5B).$^{91}$ In addition, a new silver-citrate root canal irrigation solution (BioAKT®) was introduced as a smear layer removal agent, which allows reliable sealer penetration at the apical portion of the root canal system and offers outstanding antimicrobial effects.$^{92,93}$ SDF was also reported as an adequate irrigation solution.$^{94,95}$ Additionally,
polyvinyl alcohol-coated AgNPs and farnesol are more conducive to tissue repair and less cytotoxic than NaClO. However, it is worth noting that AgNPs and SDF both have an effect on the bond strength of cemented fiber posts, which depends on the types of resin used.

Intracanal Medication and Gutta Percha Disinfectant
The most common intracanal medication is calcium hydroxide paste, which releases hydroxyl ions to increase the pH and kill microbes in the canal. Afkhami et al mixed AgNPs (size 20 nm) into calcium hydroxide paste, increasing their antimicrobial effect.

Gutta percha is the most common root canal filling material and needs to be disinfected before use. Conventional detergent sodium hypochlorite can cause surface topography deterioration while a low concentration of AgNPs can avoid this problem. In addition, mineral trioxide aggregate (MTA) mixed with AgNPs and new experimental cement (CEM) significantly increased the antimicrobial activity, along with good biocompatibility and excellent mechanical strength.

Endodontic Sealers
The requirements of ideal root canal sealants are: (1) excellent sealing properties during solidification; (2) sufficient solidification time to ensure the working time; (3) dimensional stability; (4) insoluble tissue fluid; (5) good adhesion to the tube wall; (6) suitable antimicrobial properties; and (7) biocompatibility. Enterococcus faecalis is the main microorganism associated with the failure of endodontic treatments. Utilizing antimicrobial agents, such as...
AgNPs, to mix with endodontic sealers may yield higher antimicrobial efficacy. Researchers have incorporated silver vanadate (AgVO$_3$) into various sealers, such as methacrylate resin-based sealers (Real Seal$^\text{®}$/EndoRez$^\text{®}$), zinc oxide eugenol (ZOE), epoxy resin-based sealers (AH Plus$^\text{®}$) and calcium hydroxide-based sealers (Apexit$^\text{®}$). The addition of AgBO$_3$ improves the antimicrobial performance of these commercial sealers against $E. faecalis$, $P. aeruginosa$ and $E. coli$ without reducing their mechanical properties.$^{104-107}$ Baras et al reported a therapeutic root canal sealer including 5% DMAHDM and 0.15% AgNPs to inhibit biofilms and release silver ions into the complex and difficult-to-reach root canal environment.$^{107}$ The sealer inhibited endodontic biofilms and prevented recurrent endodontic infections via potent antibiofilm ability, and ensured sufficient sealer flow and sealing ability (Figure 6). Toxic products of sealers may hamper the periapical healing process by inhibiting the proliferative capability of the periradicular cell population.$^{108}$ Although methacrylate resin-based sealers show the strongest potential to inhibit endodontic biofilms and prevent recurrent endodontic infections, they exhibit the greatest cytotoxicity and inflammatory potential against pulp cells compared to others.$^{109}$ Marica et al prepared multiwalled carbon nanotubes (MWCNTs) coated with chlorhexidine (CHX) and colloidal AgNPs and incorporated them into commercial sealants, which improved the antimicrobial activity of the modified pulp sealants.$^{110}$

**Applications of AgBMs in Periodontics**

Periodontitis is an inflammatory disease caused by multiple bacteria, resulting in the loss of alveolar.$^{111}$ Traditional periodontal treatment involves subgingival scaling and root planning to remove plaque. Antibiotics are used in periodontal pockets to achieve good anti-infection results.$^{112}$ AgNPs combined with antibiotics have a synergistic effect, which

![Figure 6 Application of AgBMs in Endodontic sealers. (A). Stereomicroscopic images showing apical dye penetration, and all experimental sealers showed similar sealing properties. (B). Live (green)/dead (red) images of 2-day biofilms adherent to resins. (C) Polysaccharide production by biofilms adherent to the sealer disks. (D) Colony-forming unit (CFU) counts of 2-day biofilms adherent to sealer disks. Reproduced with permission from Baras BH, Melo MAS, Sun J, et al. Novel endodontic sealer with dual strategies of dimethylaminohexadecyl methacrylate and nanoparticles of silver to inhibit root canal biofilms. Dent Mater. 2019;35(8):1117–1129. Copyright©2019 The Academy of Dental Materials. Published by Elsevier Inc.$^{107}$](https://doi.org/10.2147/IJN.S349238)
showed enhanced antimicrobial activity at concentrations far below the minimum inhibitory concentration (MIC) of individual antibiotics and AgNPs, with no cytotoxic effect toward mammalian cells.\textsuperscript{113}

AgBM\^{s} have the potential to promote periodontal tissue regeneration. AgNPs could be utilized to enhance new collagen synthesis and neovascularization, resulting in gingival wound healing.\textsuperscript{114} Furthermore, a type of mesoporous silica-containing AgNPs was introduced by Li et al as a novel antimicrobial and osteogenic medicine that eliminates periodontal pathogenic microorganisms and stimulates bone tissue regeneration simultaneously (Figure 7).\textsuperscript{115}

As a broad-spectrum antimicrobial material, silver can be used as an antibiotic additive to reduce the effects of bacterial resistance and can also modify scaffolds to induce alveolar bone regeneration. Conventional treatment could not regenerate alveolar bone lost in periodontitis. Guided tissue regeneration (GTR) is a surgical procedure to regain periodontal tissues by isolating periodontal defects with a membrane so that periodontal tissue will not be preempted by connective tissue. In this case, a silver-combined GTR membrane serves as an effective method to arrest bacterial colonization.\textsuperscript{116} Lee found that small concentrations of AgNPs can be used as a safe and reliable method when applied in GTR.\textsuperscript{117} Fibrin membranes rich in white blood cells and platelets can be used in a variety of regeneration treatments, and the addition of AgNPs improves their mechanical properties and antimicrobial activity.\textsuperscript{114} Electrospun chitosan/poly(ethylene oxide) membranes with different amounts of AgNPs were evaluated for antimicrobial properties and cytotoxicity in vitro and for tissue response.\textsuperscript{135} In addition, Qian et al developed a new type of silver-modified and collagen-coated electrosprin polylactic acid-glycolic acid/polycaprolactone (PLGA/PCL) scaffold with improved antimicrobial and osteogenic properties.\textsuperscript{118}

Applications of AgBM\^{s} in Prosthodontics

Prosthodontic materials require good mechanical properties to maintain shape during oral picking or chewing, so the impact on mechanical properties is a challenge before using AgBM\^{s}.

Denture stomatitis, mostly caused by \textit{Candida albicans} biofilms, occurs in two-thirds of denture wearers.\textsuperscript{42,116} Polymethyl methacrylate (PMMA) is the most conventional material used as a denture base. However, it possesses poor antimicrobial properties.\textsuperscript{119} AgNPs and certain ionic AgNPs, such as Ag\textsubscript{3}O\textsubscript{3} NPs and AgBr NPs, could be incorporated into PMMA,
improving the antimicrobial properties and mechanical strength of the denture base. Additionally, AgBMs can also be loaded and integrated into a resin base, such as silver sulfadiazine-loaded mesoporous silica nanocarriers and AgNP-loaded cellulose nanocrystals. PMMA with cellulose nanocrystal-AgNPs of 0.1 wt% possessed the most desirable mechanical properties and can be used as a functional dental restoration material (Figure 8).

Swathy et al reported that the antimicrobial effects of silver ions are weakened because they will bind to chloride in a saliva solution. To overcome this dilemma, Yoshihara et al loaded acid-silver clathrate (Met or His-Ag) into montmorillonite, a porous structure capable of controlled release, which showed a higher Ag⁺ concentration in the presence of Cl⁻. Similarly, the mouthguard (MG) used to protect athletes’ teeth requires modification of antimicrobial materials because the porous structure is

**Figure 8** 3D-Printable Reinforced Composite Resin-Based PMMA Modified with AgNPs. (A) Synthetic route of the nanocrystalline cellulose-silver (CNCs-Ag) composite. (B) 3D printed denture base using composite PMMA containing 1 wt% CNCs-Ag. Flexural properties and impact resistance of PMMA resin containing CNCs-Ag with different mass concentrations: (C) Flexural strength and (D) rupture work. Antibacterial activity of PMMA resin containing CNCs-Ag with different mass concentrations. (E) S. aureus; (F) E. coli. *p < 0.05 was considered statistically significant. Reproduced with permission from Chen S, Yang J, Jia YG, Lu B, Ren L. A Study of 3D-Printable Reinforced Composite Resin: PMMA Modified with Silver Nanoparticles Loaded Cellulose Nanocrystal. Materials (Basel). 2018;11(12):2444. Copyright © 2018 by the Authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).
a reservoir for bacterial growth. MG containing silver can significantly improve its antimicrobial activity without affecting its mechanical properties of absorption kinetic energy.\textsuperscript{129}

Oral impression is a traditional method to reproduce dentition shape in vitro, which requires antimicrobial disinfection during transportation. Conventional sprays and immersion disinfection of irreversible colloidal impression materials can cause size changes, while the addition of AgNPs can produce excellent antimicrobial activity without adversely affecting its properties.\textsuperscript{130} The fracture toughness and antimicrobial properties of dental crowns are important in prosthodontics. It was confirmed that AgNPs significantly improved the Young’s modulus and fracture toughness of dental porcelain and reduced the risk of fracture of porcelain restorations.\textsuperscript{131,132}

Applications of AgBM\textsubscript{s} in Implant Dentistry

Dental implant surgery is performed in a contaminated oral cavity, in which case, bacterial invasion (\textit{S. mutans}, \textit{P. gingivalis}) is inevitable. Polymicrobial biofilms containing Candida species—which may compromise periimplantitis—causes the loss of supporting bone and the failure of implant surgery.\textsuperscript{133}

The antibacterial performance of the implant surface can be improved by coating modification.\textsuperscript{134} To overcome this challenge, silver-containing implant coatings have attracted increasing attention because they effectively retard microbial growth.\textsuperscript{135,136} The presence of AgNPs on titanium provides antimicrobial activity related to the bacteria involved in periimplantitis.\textsuperscript{137} However, Ag biodegradation may limit its antibacterial function. Considering that early-stage (6 hours after surgery) infection is the crucial factor of implantation failure, controlled release within the postsurgery period is the key resolution. Long-term antimicrobial effects can be achieved by various coating methods, including titanium dioxide nanotubes, polydopamine coatings, microarc oxidation crafts, calcium-treated Ti metal, and poly-l-lysine/sodium alginate coatings.\textsuperscript{136,138–141} AgNPs modification can not only significantly enhance the antimicrobial activity of titanium implants

![Figure 9](https://doi.org/10.2147/IJN.S349238)

**Figure 9** AgBM\textsubscript{modified implants promote antimicrobial property and bone formation. (A) Schematic illustration of antibacterial surface modification of Ti implants for preventing implant-associated infection. (B) Fluorescence micrographs of \textit{S. aureus} and \textit{S. mutans} and (C) bacterial colony-forming units on different implant surfaces after incubation for 24 h. (A–C) Reproduced with permission from Guo C, Cu W, Wang X, et al. Poly-l-lysine/sodium alginate coating loading nanosilver for improving the antibacterial effect and inducing mineralization of dental implants. ACS Omega. 2020;5(18):10562–10571. Copyright©2020 American Chemical Society. (D) Clinical photograph and the corresponding radiograph at 12 weeks postsurgery in labrador dogs. (E) Radiography and micro-CT evaluation of the bone tissue around dental implants. (F) Histological images demonstrating the landmarks that were used for the assessment of linear distance for the soft tissue. (G) Histological sections stained with van Gieson’s picro fuchsin. (D–G) Reproduced with permission from Qiao S, Cao H, Zhao X, et al. Ag-plasma modification enhances bone apposition around titanium dental implants: an animal study in Labrador dogs. Int J Nanomedicine. 2015;10:653–64. Copyright©2015 Dove Medical Press Limited.
but also promote osteogenesis around titanium dental implants (Figure 9). Contact killing of immobilized AgNPs is another method to maintain the antimicrobial effect, offering good defense against *F. nucleatum* and *S. aureus*, with no apparent toxic influence on rat bone marrow mesenchymal stem cells. Immobilized AgNPs were almost confined to cell surfaces compared to free AgNPs, which were distributed throughout cells. Sol-gel and chloride-induced self-assembly and electrochemical deposition are the main methods for making silver coatings on titanium surfaces. The photocatalysis-mediated antimicrobial activity of AgNPs may offer a promising solution for the improvement of the antimicrobial properties of dental implants. Furthermore, silver-coated abutments could prevent leakage of microorganisms via the implant-abutment interfaces. The attachment and proliferation of *Candida albicans* on implants could be reduced by AgNPs. Interestingly, silver-induced electrochemical effects could promote antimicrobial and osteogenic effects simultaneously. Zhang et al first created a nanoscale galvanic redox system between AgNPs and 316L stainless steel alloy, which promoted peri-implant osteogenesis and antimicrobial activity.

In addition to titanium, silver coatings on other materials also show excellent antimicrobial ability and biocompatibility. Polyether ether ketone (PEEK), a thermoplastic polymer closer to the elastic modulus of the bone cortex than titanium, is mainly used in the manufacture of temporary abutments, implant abutments, and healing caps. PEEK coated with AgNPs has been proven to exhibit significantly improved antimicrobial properties and biocompatibility. In regard to implant-supported fixed partial dentures, the contact killing effect of silver is utilized to complement the weakened Ag⁺. For instance, AgNP-coated yttria-stabilized zirconia controls dental caries and periodontal disease without releasing Ag⁺.

Similar to GTR periodontal treatment, guided bone generation (GBR) aims at regenerating alveolar bone by isolating the defect with an occlusive membrane. AgNP-coated collagen membranes showed antibacterial effects against *Staphylococcus aureus* and *Pseudomonas aeruginosa*. It also reduces the release of inflammatory cytokines including IL-6 and TNF-α. Furthermore, AgNP-coated collagen membranes were able to induce osteogenic differentiation of mesenchymal stem cells that guide bone regeneration.

**AgBM in Oral and Maxillofacial Surgery**

In the area of oral and maxillofacial surgery, AgBM is employed in wound dressings and sutures due to their biocompatibility, reduction of tissue reaction and sterilization.

Colloidal silver gelatin sponges are generally utilized in tooth extraction wounds, speeding up the bone healing process in the prevention of dry sockets. Oral infectious diseases may lead to bone loss, and the rise of multidrug-resistant bacteria limits the use of antibiotics. Methicillin-resistant *Staphylococcus aureus* (MRSA) is one of the most common pathogens of oral infection. Dong et al reported that gelatin sponges containing colloidal silver can effectively reduce infections caused by MRSA in skull defects (Figure 10). Additionally, AgBM is widely employed as dressings for burn wounds. For example, silver sulfadiazine (AgSD) has been applied as a topical drug in the treatment of burn wounds since the 1960s, while its effectiveness was questioned by a Cochrane database systematic review. With the advancement of new technologies, AgSD shows strong sterilization in combined use with other dressing materials or scaffolding materials, such as microsponges and silk fibroin.

Silver wires were used as wound sutures early in 1800. In recent years, several deposition processes have been attempted to make AgNP-coated sutures a novel approach against surgical infections. These AgNP-coated sutures exhibit strong bactericidal effects on common wound pathogens, such as *S. aureus*, *E. coli*, *K. pneumoniae* and *P. aeruginosa*, with no influence on the viability of human keratinocyte cells.

Interestingly, the antibacterial properties of AgNPs can be used as a tool to regulate the microbiota to interfere with the host’s immune system. Zheng et al discovered that in oral squamous cell carcinoma (OSCC), AgNPs could regulate the oral microbiota and make *P. anaerobius* the dominant bacteria, which induces an anticancer immune response and inhibits tumor growth. This study provides an enlightening research direction for antibacterial biomaterials in which antimicrobial biomaterials could be utilized in host microbiota reactions.

**The Future Prospective**

The widespread presence of pathogenic bacteria has posed threat to healthcare, especially in the dental field. Plaque biofilms of oral cavity are complex community which can causes infection by protecting pathogenic microorganisms.
from external drug agents and escaping host defense mechanisms. Thus, plaque biofilm is a major factor resulting in caries, periodontitis, perimplantitis, and other dental diseases. AgBMs are promising in antibacterial therapies owing to their unique physicochemical properties. For example, the nanoparticle form of AgBMs have ultra-small sizes, large surface-area-to-mass ratios and increased chemical reactivity, and the large surface area of AgNPs with high charge density enables them to interact with the negatively-charged surface of bacterial cells to a greater extent resulting in enhanced antimicrobial activity. In addition, AgBMs were found to exhibit superior antimicrobial properties in the oral cavity, when combined with polymers or coated onto dental material surfaces. Thus, AgBMs enhance the antimicrobial properties of dental materials and are widely used in dental caries control, root canal therapy, prostheses, periodontal treatment, dental implants or maxillofacial surgery.

AgBMs have unique antibacterial advantages based on the variety of mechanisms. The major antimicrobial mechanism of AgBMs is now recognized as the release of silver ions. The combination of silver ions and thiol groups causes ribosome and respiratory chain dysfunction, which determines the survival of cells. The exhaustion of thiol groups in turn leads to the overproduction of ROS, destroying proteins, DNA and lipids. Moreover, AgBMs can suppress bacterial metabolism, accumulate chemicals that damage cellular components, or damage the integrity of the cell membrane. AgBMs can also be employed in PDT as promising photosensitizers for promoting and prolonging antibacterial effects. Therefore, compared with traditional antibiotics, AgBMs could improve the antibacterial effect while reducing the drug resistance.

Several disadvantages limit the clinical application of AgBMs. First, the antimicrobial effectiveness of AgBMs is limited in the oral environment due to silver resistance of bacteria. Chloride ions extensively exist in saliva and gingival crevicular fluid, leading to the precipitation reaction with silver ions. Bacteria can weaken the antibacterial properties of AgBMs through efflux ion pumps, flagellin and other means. Thus, the optimization of controlled-release mechanism and contact-killing property are potential strategies to extend the valid time of AgBMs. Combined utilization with other antibacterial biomaterials or molecules may have a better effect. For instance, the complementary use of flagellin production inhibitors also offers viable measures against gram-negative bacteria. Second, the biosafety of many AgBMs remains controversial, particularly burgeoning AgNPs. Thus, as dental materials or ingredients of topical drugs, AgBM carriers need to be designed to reduce systemic toxicity. Multiple influential toxicity factors of AgNPs...
need to be considered, such as nanoparticle size, shape, surface charge, dosage, oxidation state, agglomeration condition, and cell type. Third, AgBMs raise concern for oral aesthetics. The precipitation of some silver byproducts, such as silver amalgam and SDF, could stain dental or soft issue black. Therefore, aesthetic considerations should be taken before AgBMs are used in the oral environment. Overcoming the dyeing disadvantage of AgBMs is an important factor to promote their application in the clinic.

AgBMs have been widely researched and applied in cariostatic treatment, coatings of various dental appliances, periodontal treatment, and wound dressings because of their antimicrobial properties. Based on predictable clinical effects, AgBMs have broad development prospects in the future. AgBMs could even interfere in the immune system due to the host-microbiota reaction. Interdisciplinary antibacterial and immunomodulation may be the future development direction of AgBMs. With the advancement of research on engineering technology and composite materials, more applications of AgBMs will be developed. Safer and more efficient antimicrobial AgBMs in dentistry are expected to be obtained soon.

Acknowledgments
The authors would like to thank the original authors of the figures and all references for their significant research contributing to this manuscript.

Author Contributions
All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

Funding
This review was funded by the National Natural Sciences Foundation of China (81371680), and Jiangsu Provincial Medical Talent (ZDRC2016016), Key Project of Research and Development Plan of Jiangsu Province (BE2020629), and Development of Science and Technology of Nanjing (YKK19094).

Disclosure
The authors declare no conflicts of interest.

References


97. Mishra P, Tyagi S. Surface analysis of gutta percha after disinfecting with sodium hypochlorite and silver nanoparticles by atomic force
102. Teixeira ABV, de Castro DT, Schiavon MA, et al. Cytotoxicity and release ions of endodontic sealers incorporated with a silver and vanadium
106. Teixeira ABV, de Castro DT, Schiavon MA, et al. Cytotoxicity and release ions of endodontic sealers incorporated with a silver and vanadium
endod.2013.08.029


