Abstract: Wide-bandgap semiconductors modified with nanostructures of noble metals for photocatalytic activity under vis irradiation due to localized surface plasmon resonance (LSPR), known as plasmonic photocatalysts, have been intensively investigated over the last decade. Most literature reports discuss the properties and activities of plasmonic photocatalysts for the decomposition of organic compounds and solar energy conversion. Although noble metals, especially silver and copper, have been known since ancient times as excellent antimicrobial agents, there are only limited studies on plasmonic photocatalysts for the inactivation of microorganisms (considering vis-excitation). Accordingly, this review has discussed the available literature reports on microbiological applications of plasmonic photocatalysis, including antibacterial, antiviral and antifungal properties, and also a novel study on other microbiological purposes, such as cancer treatment and drug delivery. Although some reports indicate high antimicrobial properties of these photocatalysts and their potential for medical/pharmaceutical applications, there is still a lack of comprehensive studies on the mechanism of their interactions with microbiological samples. Moreover, contradictory data have also been published, and thus more study is necessary for the final conclusions on the key-factor properties and the mechanisms of inactivation of microorganisms and the treatment of cancer cells.

Keywords: plasmonic photocatalyst; vis-responsive material; antimicrobial effect; antifungal properties; antiviral effect; disinfection; bacteriocyte; noble metal; LSPR; environmental purification

1. Introduction

Clean water and sanitation, one of Sustainable Development Goals (adopted in United Nations summit in 2015), are highly necessary to achieve a better and sustainable society. Accordingly, the efficient technologies of water and wastewater treatment, also being accessible to everyone, have been considered as an urgent issue. It has been assumed that, in the world, one in three people do not have access to clean water for drinking and two out of five cannot wash hands properly, due to a lack of basic washing facilities [1]. Moreover, nearly 1000 children die every day due to diarrheal diseases, connected with contaminated water [1]. The coronavirus disease 2019/2020 (COVID-19) has unfortunately clarified the importance of sanitation, hygiene and adequate access to clean water for preventing the spear of infectious disease. In developing countries with insufficient water purification facilities, people must use water contaminated with feces, heavy metals, inorganic and organic pollutants, and pathogenic microorganisms. Waterborne pathogens might cause acute (e.g., diarrhea) and chronic (e.g., infectious hepatitis, cancer) health effects. Furthermore, not only slight symptoms (e.g., diarrhea), but also serious ones have been reported, such as cholera, hemolytic-uremic syndrome by Escherichia coli (E. coli) O157, and even cancer [2]. Whereas, even in developed countries, despite the fact that chemical contamination is well monitored and controlled [3–5], unpredicted outbreaks of pathogenic microorganisms have also occurred. It should be pointed out that, not only imported
infectious diseases (e.g., typhus caused by \textit{Salmonella enterica} serovar Typhi, \textit{Salmonella enterica} serovar Paratyphi A or \textit{Rickettsia prowazekii}) and polio (Poliovirus), but also illnesses caused by indigenous microorganisms via fecal-oral route (e.g., contaminated drinking water, or pools and public spas), have been observed, including a cryptosporidiosis by \textit{Cryptosporidium} species (sp.), which is resistant to chlorination, and a Legionnaires’ disease by \textit{Legionella} sp.

In association with the development of society, water demand is increasing for both domestic and industrial applications, but unfortunately water sources are limited. Therefore, it is necessary to reuse water without additional loads for the environment. In developed countries, the treatment of (waste) water is carried out by combinations of various methods, including precipitation, filtration (e.g., membrane technologies), adsorption, microbial purification (by activated sludge), oxidation (ozonization, chlorination, advanced oxidation technologies (AOPs)), depending on the kind and levels of contamination (water and municipal or industrial wastewater). Chlorination is commonly used for various types of wastewater and water, mainly as a final step of water disinfection, because chlorine can completely inactivate some microorganisms. However, it might also negatively influence aquatic organisms, due to the ability to bind to nitrogen and organic compounds, resulting in the formation of highly toxic by-products (e.g., trihalomethane). In addition, chlorine disinfection has a low effect on some pathogenic protozoa (e.g., \textit{Cryptosporidium} sp.) and viruses (e.g., norovirus). Similarly, UV-irradiation and ozonization both possess advantages and disadvantages (i.e., high efficiency against microorganisms, including viruses (UV and ozone), but low sensitivity to some viruses, such as adenovirus, no residual effect (UV), own toxicity, and the formation of toxic by-products (ozone)). Therefore, clean, inexpensive, and environmentally friendly methods for water purification and wastewater treatment are highly needed. Moreover, point-of-use water treatment methods are strongly desired in many areas that are inaccessible to municipal water treatment technologies.

Among oxidation methods, AOPs have been indicated as highly efficient and are recommended, due to the in situ generation of powerful reactive oxygen species (ROS), including hydroxyl radicals (HO\textsuperscript{*}). However, many of them involve high investments and operating costs. Heterogenous photocatalysis on semiconductor oxides also belong to AOPs, since under irradiation the semiconductor is excited, and generates charge carriers (i.e., electrons in the conduction band (CB) and holes in the valence band (VB)) might form ROS in the presence of oxygen and water. There is only one main limitation of heterogenous photocatalysis as AOPs, i.e., low activity under solar radiation because the most active photocatalysts have wide bandgap, and thus must be excited with UV, being only ca. 3–4% of solar spectrum. Accordingly, vis-responsive materials have been intensively investigated for efficient photocatalysis under real solar conditions, including plasmonic photocatalysts (i.e., wide-bandgap semiconductor modified with plasmonic NPs, such as gold, silver, platinum, and copper). Although, plasmonic photocatalysis is a new topic, various reports on plasmonic photocatalysts have already been published, including some on antimicrobial activity. Accordingly, this review summarizes and discusses the available literature on plasmonic photocatalysis for microbial inactivation. Additionally, some other biological applications of plasmonic photocatalysts are presented.

2. Plasmonic Photocatalysis

Titanium(IV) oxide (titania, TiO\textsubscript{2}) is one of the most studied semiconductor photocatalysts, due to various advantages, such as high photocatalytic activity, stability, abundance, inexpensiveness and low toxicity (except for the toxicity of nanomaterials [6]). It should be pointed out that photocatalysis has been considered as one of the best methods for environmental purification, since additional chemical compounds, such as strong oxidants (ozone, hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}), chlorine [7–14]) are not introduced into the environment [15,16], and the energy consumption is much lower than that in other AOPs (e.g., wet air oxidation [17], supercritical water oxidation [18], or H\textsubscript{2}O\textsubscript{2}/UV-C [19]). However, titania has wide bandgap of ca. 3.0–3.2 eV (depending on the crystalline form), and thus it must be excited by UV-light irradiation (absorption edge at ca. 385–410 nm). The principle of heterogeneous photocatalysis might be presented by the band-structure model. In brief, irradiation
with higher or equal energy than the bandgap excites electrons from VB to CB, generating positive holes in VB. Generated electrons \((e^-)\) and holes \((h^+)\) reduce and oxidize adsorbed substances (e.g., organic compounds, water), respectively. When electrons and holes react with water and oxygen, ROS are formed (e.g., hydroxyl radical \((\text{HO}^\cdot)\), superoxide anion radical \((\text{O}_2^{'-})\) and \(\text{H}_2\text{O}_2\)). On the other hand, charge carriers \((e^-/h^+)\) might recombine either in the bulk or on the surface of titania, which is typical for all semiconductors, resulting in a lower than expected efficiency of photocatalytic reactions. Indeed, the quantum yields of photocatalytic reactions are usually much lower than 100% (e.g., 4% has been reported for the generation of hydroxyl radicals [20]). In addition, titania cannot absorb a large fraction of solar light, which contains only about 3–5% of UV, due to its wide bandgap. Therefore, in order to improve the photocatalytic activity of titania, a number of studies have been performed, focusing on the utilization of visible light and the suppression of charge carrier recombination (e.g., surface modification, doping (nitrogen, sulfur, carbon and self-doping) and preparation of coupled nanostructures [21–30]). Although titania doping has resulted in the appearance of vis activity, dopants could be recombination centers, and thus decreasing photocatalytic activity under UV irradiation.

Modification with noble metal nanoparticles (NMNPs) (e.g., Ag, Au, Pt, Pd) seems to be the most promising, since it is well known that NM (noble metal) works as an electron sink under UV irradiation, inhibiting the \(e^-/h^+\) recombination [31–33]. When NMNPs are in contact with semiconductors, a Schottky barrier is established, hindering the charge carriers’ recombinations [34].

On the other hand, under vis irradiation, titania is activated by the plasmonic properties of NM (plasmonic photocatalysis) [35,36]. NM-modified titania photocatalysts show localized surface plasmon resonance (LSPR), and obtained vis response depends on the kind of metal (e.g., LSPR for small spherical NPs of Ag, Cu, and Au at ca. 420, 600, and 550 nm, respectively), their morphology, size, and environment (refractive index of medium). For example, Au nanospheres of several nanometers show LSPR at ca. 520 nm and bathochromic shift has been observed with an increase in the particle size [37]. A similar shift has been obtained for nanorods with an increase in their aspect ratio [37]. The most important findings by action spectrum analysis (action spectra resembling absorption spectra) confirm that plasmon resonance is responsible for vis-activity of wide-bandgap semiconductors [35,38,39].

Three main mechanisms of plasmonic photocatalysis under vis irradiation have been proposed (i.e., (i) charge transfer; (ii) energy transfer; (iii) plasmonic heating), as follows:

(i) Under irradiation with LSPR wavelength, NMNPs absorb the photons and “hot” electrons are transferred to the CB of the semiconductor. The oxygen on the semiconductor surface is reduced by the electron and the resultant electron-deficient NMNP might oxidize substances to recover to its original metallic state. For example, Tian et al. have shown an electron transfer from Au to titania and from electron donor to Au by the observation of anodic and cathodic photocurrents in Au/TiO_2/indium tin oxide (ITO) and TiO_2/Au/ITO, respectively [35,40,41].

(ii) Plasmon resonance energy transfer (PRET) might occur for energy overlapping between plasmonic metals and semiconductors (LSPR band of metals and the band gap absorption of semiconductors) (e.g., Au NPs (LSPR of ca. 2.2 eV) and Cu_2O (ca. 2.2 eV) [42]). Accordingly, although energy transfer is not expected for Au-modified titania, due to the wide band gap of titania (ca. 3–3.2 eV), PRET has been claimed for Au-modified nitrogen-doped titania, due to the generation of new energy levels (inside bandgap) [43]. Additionally, PRET has already been proposed for application in biomolecular imaging, e.g., cytochrome c in living cells [44].

(iii) Plasmonic heating has also been considered as one of the key factors for photocatalytic activities (e.g., oxidation of HCHO [45], degradation of methylene blue [46], reduction of 4-nitrobenzenethiol [47] and CO_2 reduction [48]). According to Chen et al., the plasmon-induced heating on Au NPs might activate organic molecules to induce their oxidation [45]. Although other studies have claimed that energy release by plasmonic heating is insufficient for chemical bond cleavage, it seems that local
heating must be important for microorganism killing, as microorganisms are highly sensitive to any environmental changes.

As described above, plasmonic photocatalysts induce redox reactions of organic compounds on the surface of NMNP and titania. However, those photocatalysts do not possess as strong oxidation abilities as that of titania under UV irradiation (2–3 orders in magnitude lower photocatalytic activity under vis than that under UV have been reported [33,38,39]), due to the low energy of visible light and fast charge carrier recombination (i.e., back electron transfer to NMNPs (e.g., NM→TiO₂→NM) [49]). Therefore, various studies on the improvement of the photocatalytic activities of plasmonic photocatalysts have been proposed (e.g., modification of NMNPs’ morphology [50], heterojunction with other semiconductors/complexes [33,51,52], deposition of second NMNPs [53–55], the addition of adsorbents [56–58], and morphology modifications of semiconductor [49,59–61]).

3. Heterogeneous Photocatalysis for Microbiological Purposes

The first report on microbial inactivation by heterogeneous photocatalysis was published by Matsunaga et al. in 1985 [62]. It was found that irradiated titania and platinum (Pt)-loaded titania could sterilize yeast, Gram-positive and Gram-negative bacteria, and algae, due to the photoelectrochemical oxidation of coenzyme A (CoA). Accordingly, it has been proposed that the oxidation of CoA has inhibited respiratory activity, inducing the death of cells [62–64]. The bacterial killing mechanisms by titania photocatalysts has been summarized comprehensively by Markowska-Szczupak et al. [65], pointing to three main pathways (i.e., peroxidation of cell membrane phospholipids [66,67], direct DNA damage [64], and the oxidation of CoA [62]). All these mechanisms are attributed to ROS formation from oxygen and water by reduction/oxidation reactions on irradiated titania. It has been reported that mainly HO• radicals, the most active ROS, participate in bactericidal effect [4,68], despite their short half-lifetime. Moreover, other ROS (e.g., O₂•− and H₂O₂) and direct redox reactions by charge carriers (surface e⁻/h⁺) have also been proposed for bacterial inactivation [61,69]. In addition, Du et al. have found that proteins are an initial target of radicals (HO•), preceding lipids and DNA [70]. On the other hand, some bacteria possess their protection system against oxidation stress (i.e., forming ROS scavenger enzymes, such as superoxide dismutase (SOD) and catalase (CAT)). SOD enzymes might decompose ROS to H₂O₂ and O₂, and not only in the cytoplasm, but also on the surface of cells [71,72]. Generated H₂O₂ is further decomposed to O₂ and H₂O by CAT. Importantly, some bacteria secreting CAT might be released outside of cells [73,74], which also might contribute to their resistance against extracellular ROS. For example, it has been found that the levels of SOD and CAT increase during first 30 min of titania irradiation with UV/vis, and then decrease (after 60 min) [75]. According to these changes, the two-step mechanism of bacteria response (Gram-negative E. coli and Gram-positive Staphylococcus epidermidis) has been proposed, where the changes in the enzyme activity during the photocatalytic disinfection might indicate that the defense capacity has been overwhelmed by the rapidly created ROS at the initial stage. The correlation between high content of generated HO• radicals during 90 min of irradiation and the loss of enzymatic activity suggests that oxidative stress might act as an important step, in which the photocatalyst induces bacterial death.

Moreover, heat shock proteins protect cells under an oxidative stressed environment [76], and DNA repair enzymes might repair damaged DNA [77]. In addition to enzymes, cell morphology also influences microbial inactivation by photocatalysis. It has been proposed that an inactivation effect of titania photocatalyst against microorganisms might be put in the following order: Gram-negative bacteria > Gram-positive bacteria >> fungi (yeast) > fungi (mold) [24], possibly due to the complexity of cells. Gram-negative bacteria have a thin (ca. 10 nm) peptidoglycan layer, whereas the cell walls in Gram-positive bacteria are much thicker (ca. 10–100 nm). Moreover, fungal cells (i.e., eukaryotic organisms), have rigid cell walls, containing chitin, proteins, polysaccharide polymers and lipids [78]. Furthermore, they have nuclear membranes and multicellular structures (except for unicellular organisms).
It should be pointed out that the intrinsic and surface properties of photocatalysts (e.g., bandgap energy, specific surface area, crystallite/particle size, aggregation, surface charge, defects distribution and impurities content), are also crucial for the overall antimicrobial effect [49, 54, 75, 79]. For example, rutile (polymorphic form of titania) has shown much higher activity than anatase (usually the most active titania form) under natural indoor light against mold fungi, probably because of its narrower bandgap, and thus ability to absorb more photons [79]. Moreover, it has been found that sporulation and mycotoxin generation have been highly inhibited by photocatalytic reactions on titania photocatalysts, as exemplarily shown in Figure 1. Interestingly, some fungi could be stimulated by titania presence (i.e., although the growth of Pseudallescheria boydii and Aspergillus versicolor has been highly inhibited by titania and light, the growth of Stachybotrys chartarum has been accelerated by titania). It should be remembered that different fungi species have different water and nutrient demand. Accordingly, adsorbed water and some impurities on the titania surface might either stimulate or inhibit the fungal growth [79].

![Figure 1](https://res.mdpi.com/data/covers/catalysts/big_cover)  
Figure 1. Images of three-day growth of Aspergillus melleus: (a) in the dark and (b) under irradiation in the presence of commercial titania photocatalyst P25 (after homogenization); adapted with permission (after formatting) from [79]. Copyright (2015) Elsevier.

Various applications of titania photocatalysts for microbial inactivation have already been proposed, including commercialized ones (e.g., the surfaces of equipment/walls covered with titania for efficient sterilization in hospitals). Titania modified with glucose has also been proposed for water disinfection (e.g., by placing concrete plates covered with titania in aqueous reservoirs or disinfecting water in TiO\textsubscript{2}/concrete-covered containers used for storage of drinking water or fish farms), as exemplified for a home aquarium in Figure 2a [75]. Other microbiological applications of titania photocatalysts have also been proposed, including for biomedical purposes (e.g., for decomposition of human breast cancer cells (Figure 2b) [80]).

3.1. Plasmonic Photocatalysts for Inactivation of Microorganisms

Plasmonic photocatalysts are surely promising candidates as antimicrobial agents. Although there are only several studies on microbial inactivation on plasmonic photocatalysts (i.e., under vis irradiation (not considering activities of NMNP-modified titania tested under UV and in the dark)), high activity is expected, coming from: (i) intrinsic activity of noble metals; (ii) enhanced activity under UV irradiation (inhibition of e\textsuperscript{−}/h\textsuperscript{+} recombination); (iii) activity under vis irradiation, resulting from the plasmonic activation of wide-bandgap semiconductors. The exemplary effects of the intrinsic properties of NM are described below.

Although the safety of silver for human and animals has been confirmed, it is also known that a slight amount of silver shows remarkable antimicrobial efficiency. Generally, the antibacterial activity of silver is more effective for Gram-negative bacteria than Gram-positive ones, presumably due to the differences in their biological structures. Although, the antimicrobial effect of silver has been
well-studied, the mechanism of its action is still under discussion. Therefore, different antimicrobial
pathways of Ag NPs have been proposed in the literature, as follows:

1. Adsorption of Ag cations on a negatively charged bacterial cell wall, followed by the collapse
of the cell wall and the plasma membrane, and a release of substances outside of the cell, leading to
cell lysis and death [81,82];

2. Interaction with thiol groups of transport and respiratory enzymes that are fatal to the survival
of cells, causing the uncoupling of respiration from the synthesis of ATP [83];

3. Collapse of membrane potential and a leakage of protons, as a result of the destabilization of
plasma membrane and de-energization of bacterial cells [84].

Accordingly, it has been considered that the release of Ag cations from carrier (free Ag cations in
solution/suspension) is important for bactericidal activity [85–89]. For example, it has been reported
that the activity of Ag/TiO$_2$ in the dark is higher than that in the presence of light, due to the instability
of Ag on TiO$_2$ without irradiation (i.e., Ag release) [85]. In order to obtain the continuous release of Ag
ions from Ag NPs (to keep long-term antimicrobial activities), Ag NPs have been deposited on some
supporting materials (e.g., silica [90], clay [91], and zeolite [57]). Silver NPs also possess antimicrobial
activities against fungi, algae, and viruses. The proposed mechanism of antivirus action is based on
the interaction between Ag NPs and glycoproteins from the virus surface, which induces an inhibition
of virus binding to target cells, as well as the possible inhibition of virus replication when viral cells are
penetrated by Ag NPs [92,93].

Copper (Cu) is also considered as an excellent purifier, due to its low price and high activity.
The proposed bactericidal mechanisms are based on the adsorption of Cu ions on the surface of bacteria
and, subsequently: (1) the structure of surface proteins is denatured [94], and/or (2) adsorbed Cu
ions induce oxidative stress in bactericidal process [95]. The accumulation of Cu ions inside bacteria
has been reported as the main mechanism of bacteria inactivation by Cu-modified blotting paper
(i.e., accumulation of Cu ions from direct contact with Cu NPs in the paper [96]). Furthermore, in the
case of copper oxides, it has been proposed that Cu$_2$O has higher bactericidal activity than CuO and
Ag [94]. On the other hand, Cu NPs exhibit lower toxicity than Ag NPs against some species of fungi [97].
Recently, the bactericidal activity of gold NPs has been proposed, however, it is still under discussion. There are many contradictory reports on the activity of Au NPs (i.e., Au NPs exhibit negligible activity for both Gram-negative and Gram-positive bacteria [98–100], possibly due to a lack of ability to surround bacterial cells [101], and oppositely, high activity [100,102–104]). For size dependence, Zheng et al. have shown that only Au nanoclusters (sub-nanometer size) exhibit bactericidal activity, but NPs (>2 nm) do not [100]. On the contrary, Badwaik et al. have demonstrated that, although 25-nm Au NPs show low activity, larger NPs (60 and 120-nm Au NPs) are highly active [103]. It has been proposed that the bactericidal mechanisms of Au NPs is based on: (i) the ability to change the membrane potential; (ii) inhibition of ATP synthase activities to decrease the ATP level; (iii) inhibition of the subunit of ribosome for tRNA binding and, importantly, all these mechanisms are independent on ROS generation [104]. Moreover, Au NPs possess antifungal activity, depending on the particle size of Au NPs (i.e., activity increase with a decrease in the particle size, due to an increase in the specific surface area of Au NPs, enhancing the interaction between Au and the binding sites of the plasma membrane proteins, and thus resulting in the inhibition of H+—ATPase-mediated proton pumping [105]). Au NPs also show antiviral activity (e.g., inhibition of viral attachment, entry, and cell-to-cell spread [106]).

Therefore, due to the high antimicrobial activity of NMNPs, as described above, various studies on silver- [57,69,85,89,107–109] and copper (and copper oxide)-modified titania have been reported [52,110–112]. Moreover, Au-modified titania has also been investigated, due to high photocatalytic activity (ROS generation), and the ability to disrupt the electron transport in the cells [59]. The proposed mechanisms of microbial inactivation and exemplary applications of plasmonic photocatalysts with antimicrobial properties (summarized in Table 1 for property-governed activity) are briefly discussed in the following paragraphs.

It is widely known that silver-modified titania shows high antimicrobial activity, due to the intrinsic activity of silver in the dark and enhanced activity of titania under irradiation. The release of Ag+, silver adsorption onto bacteria, and ROS generation are important factors for bactericidal action. For example, Castro et al. have suggested that increased Ag+ content on the surface of TiO2 by vis pre-irradiation (oxidation of Ag0 to Ag+ and Ag2+) enhances bacteriostatic activity in the dark [109]. Importantly, not only the inactivation of bacteria, but also the decomposition of bacterial cells is essential as the complete removal of possible allergen [113]. Accordingly, the destruction of bacteria cells (e.g., protoplast formation; Figure 3e) and initiation of their complete decomposition (mineralization), estimated by continuous CO2 evolution (Figure 4), has been observed on Ag/TiO2 photocatalysts only under irradiation (negligible CO2 evolution in the dark due to bacteria breathing), as exemplarily shown in Figures 4 and 5 [114,115].

The importance of ROS generation for the inactivation and decomposition of bacterial cells has been proposed by many researchers (e.g., vis irradiation (λ > 420 nm) of Ag/AgBr/TiO2 nanotube array causes the oxidative attack of E. coli by HO·, O2−, h+, and Br0 from the exterior to the interior, resulting in cell death as the primary mechanism of photo-electrocatalytic (PEC) inactivation (Figure 5) [116]).

Silver-modified TiO2 samples show high antibacterial activity under visible light irradiation (much higher than that in the dark) with complete decomposition of E. coli cells [114,115]. The surface properties of photocatalysts are crucial for the antimicrobial effect and, generally, a decrease in particle sizes results in an increase in activity, due to a larger interface between microorganisms and photocatalysts. Moreover, morphology-controlled titania NPs (e.g., faceted anatase with octahedral shape-octahedral anatase particles (OAP), which are highly active for the decomposition of organic compounds, also show high antimicrobial activity against bacteria (E. coli) and fungi (Candida albicans) [117]. Interestingly, it has been found that an Ag-modified OAP sample has superior bactericidal activity under visible light than that under UV–vis irradiation (Figure 6), possibly because of Ag oxidation with simultaneous release of Ag cations (an electron transfer from Ag to the CB of TiO2 under LSPR excitation; in contrast to a reverse electron transfer under UV irradiation (i.e., from TiO2 to Ag)), and thus free Ag cations might penetrate the cell membrane, resulting in the death of bacteria [117].
Similarly, Ye et al. have shown that the flower-like hierarchical TiO$_2$/Ag composites with slightly higher activity than spherical ones decompose bacterial cells (both *E. coli* and *Staphylococcus aureus*), due to the synergistic effect of the generated ROS and release of Ag ions [118]. On the other hand, the stability of Ag/TiON (on polyester) photocatalyst has been shown in repetitive experiments for bacterial inactivation under indoor light, suggesting that leaching of Ag into the environment does not happen [119]. Interestingly, van Grieken et al. have shown that substantial lixiviation of Ag (Ag$^+$) occurs in the dark, increasing the bactericidal activity, but UV irradiation stabilizes Ag deposits [85]. Liga et al. have described that increased virus (bacteriophage MS2) adsorption onto silver sites and leaching of Ag$^+$ contribute to virus inactivation [89]. In addition, enhanced ROS generation by Ag-modified titania photocatalysts has correlated with the microbial inactivation under UV–vis irradiation [69,87,120]. Swetha et al. have proposed that ROS in large quantity, generated under UV irradiation of nano titania in aqueous medium, might attack the cell membrane, peroxidized the lipid layer with the release of proteins, K$^+$ ions, nucleic acids and β-D-galactosidase [121]. However, it should be pointed out that different mechanisms of photocatalyst activation, and thus different mechanisms of microorganism inactivation should be considered under vis and UV irradiation, i.e., LSPR excitation with possible “hot” electron transfer from Ag to TiO$_2$ (resulting in more positively charged silver with high possibility of its leakage) and TiO$_2$ excitation with electron transfer to Ag (resulting in less positively charged Ag with lower possibility of leakage), respectively. Interesting data have been found by action spectra analyses (activity depending on the irradiation wavelengths) for titania modified with mono- and bi-metallic Ag and Au NPs, where high stability of Au@Ag/TiO$_2$ photocatalysts under vis irradiation resulted from the possible stabilization of silver by gold (electron transfer from Au to electron-deficient Ag) [122,123].

**Figure 3.** SEM images of the decomposition of *E. coli* under vis ($\lambda > 420$ nm) irradiation of Ag/TiO$_2$ photocatalyst: (a) healthy bacteria cells, (b) bacteria covered with titania before photocatalytic reaction, (c,f) destroyed bacteria cells after: (c) 1 h, (d,e) 3 hand (f) 24 h of irradiation; adapted with permission from [115]. Copyright 2018, Creative Commons Attribution.
Although fungal cells are more complex than bacterial cells and quite resistant to antimicrobial agents, plasmonic photocatalysts have been proposed as effective fungicides \[117,124-126\]. For example,

**Figure 4.** *E. coli* decomposition presented as a decrease in bacteria number (closed symbols) and CO\(_2\) evolution (open symbols), tested in the suspension of titania under vis irradiation (\(\lambda > 450\) nm) and in the dark; adapted with permission (after formatting) from \[114\]. Copyright 2015 Elsevier.

**Figure 5.** *E. coli* inactivation by PEC on Ag/AgBr/TiO\(_2\)-nanotube electrode: (a,b) ESEM images of *E. coli*: (a) untreated and (b) after PEC inactivation for 80 min; (c,e) TEM images of *E. coli*: (c) untreated and after PEC inactivation for: (d) 40 min and (e) 80 min; (f) inhibition of *E. coli* inactivation by scavengers. Reprinted with permission from \[116\]. Copyright (2012) American Chemical Society.

Recently, the fungicidal activity of Ag-modified titania has also been studied intensively. Although fungal cells are more complex than bacterial cells and quite resistant to antimicrobial agents, plasmonic photocatalysts have been proposed as effective fungicides \[117,124-126\]. For example,
the antifungal activity (C. albicans) of Ag-modified faceted titania (OAP) is much higher than that by OAP modified with other NMMPs (Cu, Au, and Pt), both under visible light and in the dark (ca. 50% higher activity under vis during 1 h) [117]. Moreover, it has been reported that the fungicidal activity of Ag–TiO₂ depends on the fungal strain and Ag content [124], and thus the higher content of Ag–TiO₂ or prolonged exposure time might be necessary for specific fungal species [125]. Interestingly, although mono Ag or Cu-modified titania does not show enhanced activity under visible light, bi-metallic titania (Ag/Pt and Cu/Ag) promote vis activity, due to the oxidation of organic compounds by superoxide anions [126]. Similarly, a synergistic effect has been observed for titania modified with Ag@CuO in respect to mono-modified titania (Ag/TiO₂ and CuO/TiO₂) against mold fungi (A. melleus and P. chrysogenum) [127]. Interestingly, only the sample with molar ratio of Ag to CuO of 1:3 shows synergism, and only bi-modified samples exhibit higher activity in the dark than that under vis irradiation.

![Figure 6](image_url)  
**Figure 6.** Antibacterial activity of OAP samples modified with NMMPs under: (a) UV/vis irradiation and (b) vis irradiation; adapted with permission (after formatting) from [117]. Copyright 2017 Creative Commons Attribution.

| Sample | Preparation Method | Important Properties | Experimental Procedure | Best Performance | Ref. |
|--------|--------------------|----------------------|------------------------|-----------------|-----|
| Ag/TiO₂ | hydrothermal synthesis | Ag oxidation | UV or vis-preirradiated samples under vis irradiation with *Escherichia coli* | non-treated and vis-preirradiated samples under vis | [109] |
| AgOAP | hydrothermal synthesis (OAP) and Ag or Cu by photodeposition | Ag oxidation, Ag release | catalysis suspensions with bacteriophage MS2 under vis, UV, and dark | small Ag in dark and large Ag under vis | [114] |
| CuO/TiO₂ | impregnation method | Cu₂O/CuO ratio | suspension of E. coli, S. aureus or Qβ bacteriophage on CuO/TiO₂ under vis | CuO clusters, Cu₂O | [52] |
| Au/TiO₂ | magnetron sputtering | size and distribution of Au | E. coli or S. aureus attached on Au/TiO₂ in dark | annealed Au/TiO₂ (large and isolated Au) | [59] |

**Table 1.** Key-factors of antimicrobial properties of NM-modified titania photocatalysts.
Copper, especially copper (I) oxide (although it could not be considered as plasmonic, usually a mixed-oxidation state of copper has been reported), has been well known as an antimicrobial agent since ancient times. It has been applied to improve the photo-induced antimicrobial activity of titania. The proposed mechanisms include: (i) the structure of surface proteins being denatured [94]; (ii) the adsorbed copper ions inducing oxidative stress in the bactericidal processes [95], and the accumulation of copper ions inside bacteria [128]. It has been found that the optimal balance between Cu$_2$O and CuO in the Cu$_{x}$O/TiO$_2$ composite photocatalyst is important to achieve good antibacterial performance under visible light irradiation and dark conditions, and furthermore that Cu$_2$O/TiO$_2$ is more active than CuO/TiO$_2$ and CuNPs/TiO$_2$ [52]. The mechanistic study by Rtimi et al. In the presence of scavengers (dimethyl sulfoxide and superoxide dismutase) has shown that the VB holes in TiO$_2$ and the toxicity of the Cu ion are responsible for E. coli inactivation under actinic light [111]. In the case of two kinds of faceted anatase photocatalysts (i.e., OAP and DAP (decahedral anatase particle)), the modification with NM (Ag, Cu, Au and Pt) resulted in enhanced antibacterial activity (in comparison to the activity of bare samples) under vis irradiation ($\lambda > 420$ nm) against E. coli only for Ag- and Cu-modified samples [129]. It has been proposed that the surface oxidation states of NM deposits have been responsible for this behavior (i.e., mainly $+1$ for Ag and Cu, and 0 for Au and Pt), resulting in the facile adsorption of Ag/Cu-TiO$_2$ on the bacteria surface. Moreover, these results indirectly support the mechanism of plasmonic activation of titania by the charge transfer mechanism, resulting in the formation of more positively charged Ag/Cu under vis irradiation, and thus an increase in the overall bactericidal activity [122,123,129]. The change in surface oxidation states of NM has been confirmed in another study by XPS analysis for mono- and bi-metal (Au/Ag)-modified titania under monochromatic irradiation [122,123].

In addition to Ag and Cu, Au-modified titania has also been considered as an antimicrobial agent. Moreover, not only the plasmonic photocatalysis (ROS generation) under irradiation [61,115,130,131], but also electron transfer between Au and bacteria (extracellular electron transfer) [59,132] might cause bacterial death. On the other hand, another report insists that Au-modified titania does not possess any bactericidal property [120]. Although silver-modified samples usually show much higher activity than other plasmonic photocatalysts, gold-modified samples prove to be the most active against mold fungi, especially for the inhibition of the sporulation, as shown in Figure 7 [54,115,122].

### 3.2. Biomedical Applications of Plasmonic Materials

Considering the use of titania for other biological applications, cancer therapy should be noticed, as already presented for bare titania in Figure 2b. Titania has various advantages (e.g., effective ROS generation and non-toxicity), but also disadvantages (e.g., usage of UV light (low penetration through skin, thus, limited to superficial cancer) and no-specification to the target. To overcome these disadvantages, titania has been modified in various ways to obtain visible–near infrared (NIR) response and specification for cancer cells. For example, hydrogenated black titania [133–135], green titania [136], titania modified with NM nanocomposites [137–139], and core/shell photocatalysts with NaYF$_4$: Yb, Tm up-conversion NPs as a core and titania as a shell [140], show NIR absorption and have been proposed for cancer therapy. Among them, plasmonic photocatalysts seem to be the most promising antitumor agents, due to the enhanced photocatalytic property and vis/NIR plasmonic absorption. For example, in the case of Au nanorods and nanoshells, LSPR might be tuned to the NIR region, enabling us to perform in vivo imaging and therapy through the selective localized photothermal heating of cancer cells [141].

Xu et al. [142] have found that Au/TiO$_2$ nanocomposites, synthesized by the deposition–precipitation (DP) method, might kill the carcinoma cells efficiently, and the amount of Au on the surface of TiO$_2$ strongly affects the photocatalytic inactivation efficiency (i.e., 2 wt% of Au being the most active). Seo et al. have demonstrated that Ag/AgBr/TiO$_2$ NPs induce the killing of mammalian cancer cell lines in vitro under visible light illumination (>450 nm) and, moreover, reduced tumor volume in vivo (Figure 8) [139].
Therefore, similar to drug delivery systems, titania has been modified with various photocatalysts that can be activated by light. For instance, TiO₂ NPs can be used as a photocatalyst for the photothermal effect. When irradiated with a laser, these NPs generate heat, which can be localized to the cancer site, thus causing cell death. This localized heating is highly effective because it minimizes damage to surrounding healthy cells.

One example of a metal oxide-based photocatalyst is Au/TiO₂, which shows promise in photodynamic therapy. The Au NPs facilitate the transfer of energy to the TiO₂, which then generates singlet oxygen, a highly reactive species that can damage cancer cells. The Au NPs enhance the photothermal effect, while the TiO₂ provides stability and durability.

Another interesting aspect of photocatalysts is their ability to target cancer cells. For example, gold nanorods (Au NRs) can be functionalized with ligands that bind specifically to cancer cell surface receptors. This targeted approach can improve the efficiency and specificity of the treatment.

In conclusion, while plasmonic photocatalysts may cause anticancer effects, they are deficient in specificity. Therefore, similar to drug delivery systems, titania has been modified with various strategies to improve their targeting and specificity.
components. For example, some receptors are overexpressed on the surface of cancer cells (e.g., epidermal growth factor receptor (EGFR) [143], interleukin-13R2 receptor domain (IL13R2) [144] and folic acid-folate receptor [145]). Accordingly, in order to add the binding ability to cancer cells to titania (or other materials), an 11 amino acid peptide fragment of EGF [143], anti-IL13R2R antibody [144], and folic acid [145] are first conjugated with titania. Accordingly, the bound titania to receptor is uptaken by endocytosis, incorporating into cytoplasm, and then probably into nucleus. Next, ROS are generated under irradiation and attack proteins, lipids, nucleic acid, and so on, inducing cell death by apoptosis or necrosis. Recently, it has been found that nuclear-targeted AuNPs (AuNPs with three ligands, methoxypolyethylene glycol thiol (PEG), RGD (RGDGRDRGDRGDPGC) peptides, and nuclear localization signal (NLS, CCGPKKKRKVK GG) peptides), inhibit cancer cell migration by increasing their nuclear stiffness, which greatly reduces the AuNP dosage, resulting in the suppression of the metastasis [146]. Moreover, Ali et al. have proposed that integrin-targeted Au nanorods (Au nanorods with Arg–Gly–Asp (RGD) peptide), activated by NIR light, causes changes in cell morphology and a decrease in cell migration [138], since integrin plays critical roles in controlling the organization of cytoskeletons (strongly related to cell migration).

On the other hand, plasmonic photocatalysts might be used as drug-delivery carriers for anti-tumor drugs [137]. For example, gold nanorods (GNR)/TiO$_2$ has been used as the carrier of gambogic acid (GA), a potent anticancer agent with poor solubility in aqueous solutions, providing the stable dispersion and enhanced intracellular GA delivery. Moreover, GNR/TiO$_2$ shows high photothermal conversion efficiency, and thus irradiation with low-intense laser at 808 nm has enhanced the anticancer effect of the GA-loaded GNR/TiO$_2$, resulting in the improvement of the therapeutic efficacy of GA (Figure 9).

![Figure 9](image_url)

**Figure 9.** Enhanced cytotoxic effect of GA by using GNR/TiO$_2$ nanostructure-mediated photothermal therapy. U-87 MG cells were incubated with GNR/TiO$_2$ or GA-loaded GNR/TiO$_2$ nanostructures for 24 h, followed by 808 nm NIR irradiation (5.3 W cm$^{-2}$) for 2 min. After incubation for 24 h, cell viability was determined using an ATP assay (a) and calcein AM staining; (b) Live cells were stained with green fluorescence by calcein AM. The data shown represent the mean ± S.E.M., ***p < 0.001. Reprinted with permission (after formatting) from [137]. Copyright (2017) The Royal Society of Chemistry.

4. Conclusions

Plasmonic photocatalysts have proved to be efficient agents against various microorganisms. Their activity is usually much higher under vis irradiation than in the dark (intrinsic properties of noble metals), resulting from the generation of ROS and the direct redox reactions between microorganisms and photocatalysts. The properties of both photocatalysts and target microorganisms are crucial for the overall antimicrobial effect. Generally, smaller NPs (and thus larger specific surface area) and more simple organisms have shown fast and efficient microbial inactivation.
Unfortunately, some contradictory results have been published. It might be expected that some experiments have been performed in the presence of UV irradiation (e.g., indoor light, artificial or natural solar light), and thus different mechanisms of microorganism inactivation should be considered, i.e., direct excitation of wide-bandgap semiconductor (under UV) and plasmonic excitation (under vis). Considering “hot” electron transfer under plasmonic excitation, the opposite electron transfer under vis and UV is expected, i.e., from NM to semiconductor and from semiconductor to NM, respectively. Therefore, the change in the oxidation state of less noble metals (e.g., Ag and Cu) might result in the facile adsorption of the photocatalyst on the surface of microorganisms, but also a release of metal cations into solution/suspension. For the mechanisms study, it seems that the tests under sole vis (also under sole UV for the comparison) and in the dark are the most recommended (e.g., by action spectra analysis).

The other microbiological applications of plasmonic photocatalysts are also quite promising, especially for cancer treatment and anticancer therapy, but more studies on the improved selectivity against only danger cells are highly necessary.

**Author Contributions:** Conceptualization, E.K.; writing—original draft preparation, M.E.-K.; writing—review and editing, E.K. and M.E.-K.; supervision, E.K. All authors have read and agreed to the published version of the manuscript.

**Funding:** This research was funded by a Grand Challenges Explorations Grant (GCE RB, OPP1060234) from Bill & Melinda Gates Foundation, and “Yugo-Sohatsu Kenkyu” for an Integrated Research Consortium on Chemical Sciences (IRCCS) project from the Ministry of Education and Culture, Sport, Science and Technology-Japan (MEXT). The APC was funded by E.K.

**Acknowledgments:** Authors thank Agata Markowska-Szczupak from Western Pomeranian University of Technology in Szczecin, for fruitful discussion, motivation, and support.

**Conflicts of Interest:** The authors declare no conflict of interest.

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