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ORIGINAL RESEARCH

Biogenic pentagonal silver nanoparticles for safer and more effective antibacterial therapeutics

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Background: Biological synthesis of nanomaterials possesses unprecedented potential in the production of nanomaterials due to their ability to produce nanomaterials with improved biocompatibility in addition to eco-friendly synthetic procedures.

Methods: This article reports the isolation of an air-borne fungus from the campus of Integral University, Lucknow, with an exceptional ability to withstand very high concentrations of silver salt. The fungus was found to produce pentagonal silver nanoparticles (AgPgNps) when silver ions were reduced from silver nitrate. Molecular analysis and biochemical characterization techniques based on 18-seconds rRNA identified the fungus to belong to the *Aspergillus* sp. with the NCBI accession no KF913249. Material characterization techniques including ultraviolet (UV)–visible spectroscopy, transmission electron microscopy, and zeta potential analysis were used to satisfactorily characterize the as-synthesized AgPgNps.

Results: The AgPgNps synthesized by the fungus *Aspergillus* sp. exhibit an absorption that is maximum centered at about 416 nm, with a standard particle size of 23.22 \pm 2 nm. These AgPgNps exhibited broad-spectrum antimicrobial activities against an array of bacterial pathogens with remarkable minimum inhibitory concentration (MIC₅₀) values: *Staphylococcus aureus* (ATCC 25923) – 9.230 µg/mL, *Bacillus* sp. (ATCC 14593) – 12.781 µg/mL, *Escherichia coli* (ATCC 25922) – 5.063 µg/mL, and *Klebsiella pneumoniae* (ATCC 13883) – 5.426 µg/mL. In vitro cytotoxicity analysis of biosynthesized AgPgNps showed a dose–response activity against human cervical cancer cell line (HeLa) and adenocarcinoma cells (A549) with MIC₅₀ values of 0.038 µg/mL and 0.044 µg/mL, respectively.

Conclusion: These findings are very crucial to evaluate the biosynthetic process for the synthesis of nanoparticles (NPs) with unique properties. These NPs may find potential applications in sensing, medicine, and antimicrobial and anticancer therapies.

Keywords: pentagonal Ag NPs, fungus, biosynthesis, antibacterial agent, anticancer, in vitro cytotoxicity

Introduction

Different forms of silver such as silver metal, nitrates salts of silver, and silver sulfadiazine have been used to treat injuries due to burns and bacterial infections since ancient times.¹ Silver metal also shows antimicrobial activity against a wide range of both Gram-negative and Gram-positive bacteria, fungi, protozoa, and certain viruses. The continuing fight against the emergence and robust nature of antibiotic-resistant bacteria has built high hopes over the efficacy of silver and silver-based compounds in tackling pathogens, including silver nanoparticles (NPs).² Metallic silver in the form of silver NPs has shown lots of promise as a potential antimicrobial agent. Thus, a wide assortment of uses has discovered use of silver NPs going from silver-based dressings and silver-covered medicinal gadgets, eg, nanogels and nano lotions.¹

International Journal of Nanomedicine 2018:13 7789-7799

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The characteristics of microbicidal actions of silver NPs (from fungi to viruses) is an active field of research.³ Silverbased compounds such as silver-doped polymer fabrics, catheters, polyurethane,⁴ colloidal silver,⁵ fabrics laced with silver NPs,⁶ nano silver metal oxide granules,⁷ and ceramic materials coated with silver NPs⁸ are well known for their bactericidal functionality, while burn care uses silver compounds in an array of different forms.⁹ The bactericidal and veridical properties of silver and its derivatives have been altogether settled.^{10–12} However, in small concentrations, silver shows no toxicity toward animal cells, whereas it shows high toxicity toward microbes.

Silver has been widely applied in the medical industry since ancient times due to its inherent nontoxic nature.^{13,14} Silver possesses unique properties in that it is less toxic at certain concentrations to animal cells and certain bacteria while being effective against certain other pathogenic strains of bacteria at that concentration.¹⁵ It is widely established that the bactericidal properties of silver are ascribed to the catalytic oxidation by metallic silver and response with broke up univalent silver ions.¹⁶ The development of resistance by microbes against silver is not feasible since the microorganisms would have to undergo an array of mutations to overcome the damages to the targets attacked by the metals.¹³ In small concentrations, silver is unique in that it is safe for human cells at less concentrations while being very effective against different types of microbes and also has found application in medicines and in the maintenance of water and food qualities.¹⁷ In general, the toxic nature of metal NPs toward cells is attributed to their ability to produce reactive oxygen species (ROS).¹⁸ The ability of silver NPs to get oxidized slowly to release Ag⁺ ions renders them a potential biocide. In addition, these small-sized particles are ideal to move across the cell membrane to attack intracellularly. Additionally, the antibacterial properties of these NPs can also be attributed to their enhanced surface area to volume ratio characteristic.¹⁹ However, the effects of NPs on the inflammatory responses in humans have not been extensively investigated. Elechiguerra et al³ reported size-dependent interaction of silver NPs with HIV type 1 by binding to gp120 glycoprotein knobs to bring about an effect. In a similar note, they reported the size-dependent action of silver NPs against Gram-negative bacteria Escherichia coli.

In this study, a novel fungus was isolated from the campus of Integral University, Lucknow, and used to synthesize pentagonal silver nanoparticles (AgPgNps), which showed bactericidal properties against *Bacillus* spp. (ATCC 14593), *Staphylococcus aureus* (ATCC 25923), *E. coli* (ATCC 25922), and *Klebsiella pneumoniae* (ATCC 13883). These NPs showed no toxicity against primary osteoblast cells up to a moderately high concentration.

It might be estimated that different shaped (having almost equal surface area) silver NPs might be having distinctive features interacting with the surroundings. In this report, the relationship between the surface area of the NPs and their antimicrobial properties is not discussed. Even though there is a dearth of literature at present on the bactericidal efficacy of silver NPs related to their active surface areas, this study might serve as an important foundation to emphasize the shape-dependent bactericidal properties of silver NPs. Nonetheless, it is basic that the innovation in the methods of NP synthesis and their effect on different microbial strains should be investigated.¹³ Another study reported on the use of exopolysaccharides produced by Aeromonas punctata as the stabilizing agent by capping the NPs.²¹ A relatively less toxicity was shown by exopolysaccharide-capped particles toward S. aureus, E. coli, and Micrococcus luteus compared to that by non-capped particles, which shows that the encapsulation of exopolysaccharides acts as a physiological defense tool.²² Khan et al have done considerable research for the utilization of microbial-inferred biomolecules in the surface functionalization of nanomaterials. As indicated by their examination, zeta capability of NPs influences the surface adsorption of exopolysaccharides on NPs.²

Materials and methods Materials

All the chemicals and solvents were bought from HiMedia, Merck, and Sigma-Aldrich Co. (St Louis, MO, USA).

Isolation of fungi

The fungus used in this study was isolated from the air in the campus of Integral University, Lucknow, Uttar Pradesh, India. Malt extract Glucose Yeast extract Peptone (MGYP) media containing 5 mM silver nitrate solution was exposed to the open air to entrap fungus capable of withstanding high concentrations of metal salt. The entrapped fungal strains were isolated to purity using the serial dilution method. The pure fungal strain was stored at -20° C in 30% glycerol.

Taxonomic identifications

The DNA was extracted from 48 hours old culture of fungus grown in 100 mL MGYP broth for phylogenetic analysis by analyzing 18S rRNA gene. Universal oligonucleotide primers 18SF (5'-CGW CGR AAN CCT TGT NAC GAS TTT TAC TN-3') and 18 SR (5'-AWG CTA CST GGT TGA TCC TSC CAG N-3') from Amnion Biosciences Pvt. Ltd. (Hyderabad, India) were used for amplification. The given polymerase chain reaction (PCR) program was followed for amplification of 18S rRNA gene (Bio-Rad thermal cycler; Bio-Rad Laboratories Inc., Hercules, CA, USA): one cycle of preheating for 5 minutes at 94°C and 30 cycles at 94°C for 30 seconds, 55°C for 30 seconds, and 72°C for 3 minutes. These cycles were followed by incubation at 72°C for 5 minutes. Eventually, the reaction mixture was cooled to 4°C. The amplified PCR product was checked using 1% agarose gel with 5 kb molecular weight markers. The amplified gene was sequenced by the Sanger Dideoxy Sequencing/Chemistry-BDT version 3.1 method according to the company's protocols using ABI 3730xl genetic analyzer (Amnion Biosciences Pvt. Ltd.). NCBI "BLASTn" program was used for alignment and assembly of the sequences. The neighbor-joining method was used to establish evolutionary history from 1,000 replicates.²³

Microorganism and growth

The Potato Dextrose Agar slants with composition 20% w/v potato, 2% w/v dextrose, and 2% w/v agar were used to maintain new isolate of *Aspergillus* sp. (accession no KF913249) at 25°C. The 7-day-old Potato Dextrose Agar slant was used to inoculate a fungal mat (1 cm in diameter) into 100 mL liquid MGYP medium (malt extract [0.3% w/v], glucose [1.0% w/v], yeast extract [0.3% w/v], and peptone [0.5% w/v]) with incubation at 26°C±1°C on a rotary shaker (200 rpm) for 96 hours. Eventually, centrifugation (4,500× g, 20 minutes at 10°C) was used to collect mycelia under aseptic conditions with distilled water and used accordingly.

Extracellular synthesis of AgPgNps

The extracellular synthesis of AgPgNps was achieved by incubating mycelia (20 g wet mass) with 1 mM AgNO₃ in 100 mL sterile distilled water under shaking conditions (200 rpm) for 120 hours at room temperature. The progress of synthesis of AgPgNps was checked regularly using ultraviolet (UV)–visible spectroscopy. The unutilized proteins were precipitated by using absolute ethanol (2 volume), and isolated AgPgNps were collected for further analysis.

Characterization of AgPgNps

Several physical techniques were used to characterize AgPgNps such as UV–visible spectroscopy, zeta potential analysis, dynamic light scattering, and transmission electron microscopy (TEM). Shimadzu dual-beam UV–visible spectrophotometer (model UV-1601 PC) operated on a declaration

of 1 nm was used to authenticate the preliminary synthesis of AgPgNps. Zetasizer Nano-ZS model ZEN3600 (Malvern Instruments, Malvern, UK) was used to measure zeta potential. A drop of AgPgNps (suspension) was dried on a carboncoated TEM copper grid to analyze the size of the inorganic core using transmission electron microscope (TEM) TecnaiTM G^2 Spirit Bio-TWIN (FEI, Hillsboro, OR, USA) at a step-up voltage of 80 kV.

Fourier transform infrared (FT-IR) spectroscopy of biosynthesized AgPgNps

The presence of different functional groups at the surface of AgPgNps was confirmed by Perkin-Elmer Spectrum 2 FT-IR (PerkinElmer Inc., Waltham, MA, USA) furnished through a worldwide attenuated full reflectance sampling tool and scanned at room temperature using transmission approach above the wave number range of 4,000–650 cm⁻¹ at a resolution of 4 cm⁻¹.

Antibacterial activity

Determination of minimum inhibitory concentration (MIC_{50})

The preliminary antibacterial potential of AgPgNps was checked by the disk diffusion method.²⁴ The MIC₅₀ of confirmed antibacterial AgPgNps was evaluated by the microdilution method.²⁵ Briefly, S. aureus (ATCC 25923), Bacillus sp. (ATCC 14593), K. pneumoniae (ATCC 13883), and E. coli (ATCC 25922) were obtained from NCIM, National Chemical Laboratory (Pune, India), developed to mid-logarithmic stage, gathered by centrifugation, rinsed with 10 mM sodium phosphate support (SPB) at pH 7.4, and diluted to 2×10⁵ colony forming units (CFUs)/mL in SPB including 0.03% Luria-Bertani (LB) broth. The required concentrations $(0.1-15 \ \mu g/mL)$ of AgPgNps with the bacterial inoculum $(5 \times 10^4 \text{ CFU per well})$ were successively diluted in 50 µL of LB medium within 96-well microtitre dishes and incubated overnight at 37°C. The lowest concentration of AgPgNps that inhibited 50% bacterial growth was considered as MIC₅₀. The agar plate count method was adapted to further confirm the results. In this method, 25 µL aliquots from the freshly prepared 96-well microtitre plates (as abovementioned) were plated on LB plates and kept at 37°C for a night. Eventually, grown colonies of bacteria were counted on the next day.

Determination of minimum bactericidal concentration S. aureus (ATCC 25923) – MIC₅₀ 9.230 μ g/mL, Bacillus sp. (ATCC 14593)–MIC₅₀ 12.781 μ g/mL, E. coli (ATCC 25922)– MIC₅₀ 5.063 μ g/mL, and K. pneumoniae (ATCC 13883)–MIC₅₀

5.426 µg/mL were used to evaluate minimum bacterial concentration (MBC). The culture (2 µL) from the well of a microtitre plate (which was used to determine MIC_{50}) with no visible growth was transferred into another microtitre plate having 100 µL of broth for serial subcultivation and incubated for next 24 hours. The minimum concentration with no noticeable growth was characterized as MBC, showing a 99.5% loss of viability of the first inoculum. The bacterial growth was estimated at 620 nm wavelength using Microtitre Plate Manager 4.0 (Bio-Rad Laboratories Inc.) and evaluated with a blank. The same was also repeated on Petri plates for validity using dimethyl sulfoxide (DMSO) as the negative control. The experiment was repeated thrice for each compound with three replicates every time.

Cell culture

The human adenocarcinoma (A549) and human cervical growth cell line (HeLa) were received from National Center for Cell Science (NCCS; Pune, India). The primary osteoblasts were segregated from calvaria of the neonatal rat through enzymatic processing. Primary osteoblasts cells, A549 and HeLa, were developed as a monolayer in MEM, DMEM–F12, and EMEM media, individually, supplemented with 10% fetal bovine serum and 1% antibiotics containing 10,000 units of penicillin, 10 mg of streptomycin, and 25 μ g of amphotericin B in a humidified CO₂ incubator at 37°C. All the cell stocks were managed in 75 cm² tissue culture flasks.

Assessment of cytotoxicity

The cytotoxicity of AgPgNps, A549, and HeLa cells was assayed according to the protocol described elsewhere²⁶ where various concentrations (0.373, 0.186, 0.0932, 0.046, 0.0233, and 0.0116 μ g/mL) of AgPgNps were used to determine the IC₅₀ against each cell line.

Measurement of cytomorphological changes in A549 and HeLa cells

A549 and HeLa cells were pretreated with various concentrations of AgPgNps and kept warm for 48 hours at 37° C in 5% CO₂ atmosphere. Toward the end of incubation, net morphological transforms in the cells were seen using an inverted phase contrast microscope (Nikon Eclipse Ti-S; Nikon Corporation, Tokyo, Japan).

Analysis of changes in nuclear morphology

Nuclear morphological changes in AgPgNp-treated and -untreated (control) primary osteoblast A549 and HeLa cells were observed under an Olympus fluorescence microscope associated with an Olympus Cool Snap Camera (Olympus Corporation, Tokyo, Japan) after staining with 4',6-diamidino-2-phenylindole (DAPI). Observation of minimum 200 cells under the microscope revealed that live cells were having normal nuclei and blue chromatin with organized structure; stressed cells were found to have bright-blue chromatin, which was highly condensed, marginated, or fragmented. Three different softwares, namely, Micro-Image, ImageTool, and Matlab 7.0, were used for quantitative estimations. On the other hand, control DAPI staining revealed a single, bright, spherical nucleus and peripheral cell spots corresponding to stained mitochondria. The investigation of apoptosis on AgPgNp-treated A549 and HeLa cells were finished by staining with fluorescent nuclear dye DAPI. The cells were seeded and treated as mentioned earlier. Then, the cells were rinsed with PBS and kept in 4% paraformaldehyde for 10 minutes. Consequently, the cells were permeabilized through the permeabilizing buffer (3% paraformaldehyde and 0.5% Triton X-100) and stained with DAPI dye. A fluorescence microscope (Nikon Eclipse Ti-S) was used to acquire images of treated cells after staining. The cells with fragmented and condensed nuclei were measured as apoptotic cells.

Measurement of intracellular ROS generation

The Image-iT LIVE Green Reactive Oxygen Species Detection Kit (Invitrogen) was used to detect the production of ROS in AgPgNp-treated HeLa and A549 cells. The ROS production in viable cells was directly estimated by using a fluorogenic marker, 5-(and-6)-carboxy-2',7'-dichlorodihydrofluorescein diacetate (carboxy-H₂DCFDA). The 96-well culture plates were used to seed the cells (1×10^4 per well) for 24 hours in a CO₂ incubator at 37°C. Diverse concentrations (0.373, 0.186, 0.0932, 0.046, 0.0233, and 0.0116 µg/mL) of AgPgNps were used to incubate HeLa and A549 cells for 24 hours. Eventually, H₂DCFDA (10 mM) was used to incubate cells for 30 minutes at 37°C. The reaction mixture was aspirated and changed by 200 µL of PBS in all the wells. The plates were kept back on a shaker for 10 minutes at room temperature within the dark. The intracellular fluorescence of cells was examined under an inverted fluorescence microscope (Nikon Eclipse Ti-S).

Quantitative estimation of ROS generation was done by re-seeding cells $(1.5 \times 10^4 \text{ per well})$ in 96-well black bottom culture plate with incubation for 24 hours at 37°C. Subsequently, cells were treated with various concentrations of AgPgNps (0.373, 0.186, 0.0932, 0.046, 0.0233, and 0.0116 µg/mL) for 12 hours and additionally incubated with 10 µM of DCFH-DA for 30 minutes at 37°C. Fluorescence

Biogenic Pentagonal AgNPs effective against bacteria

intensity was estimated by the multiwall microplate reader (Synergy H1 Hybrid Multi-Mode Microplate Reader; BioTek, Winooski, VT, USA) at an excitation wavelength of 485 nm and the emission wavelength of 528 nm. Values were articulated as power percent overlay of fluorescence intensity with respect to control.

Results and discussion

Isolation, molecular characterization, and phylogenetic analysis of Aspergillus sp.

The fungus was isolated from the air at the campus of Integral University, Lucknow, India. In the quest for an uncommon fungus with uncommon features using an uncommon approach, MGYP media with 5 mM silver salt solution was exposed to the open air. In this particular media, only those fungi were able to grow that could withstand such high concentrations of silver ions. Furthermore, these fungi were found to possess the ability to produce AgPgNps by the reduction of silver ions.

The PCR-amplified 18S rRNA gene was found to be 1,718 bp long (Figure 1). This sequence was submitted to the NCBI with the unique accession number KF913249. The evolutionary history of the above gene was constructed

based on the Jukes–Cantor model using the maximum likelihood method using Molecular Evolutionary Genetics Analysis 6 (Figure 2).^{27–29} A polyphasic approach combining phenotype and molecular techniques was used to classify taxonomy of the isolate. The sequence similarity of the 18S rRNA gene (1,718 bp) showed that this fungal isolate is a close relative of *Aspergillus* sp. This isolate encompasses a unique phylogenetic position with respect to *Aspergillus fumigates* and other species of *Aspergillus* sp. Thus, on the basis of physico-biochemical tests and molecular characteristics, this isolate appears to occupy a distinct phylogenetic position by being one of the unique species of *Aspergillus* and was named *Aspergillus* KF913249.

Synthesis of pentagonal NPs

The AgPgNps were synthesized by incubating 100 g of fungal mycelia (in the log phase), *Aspergillus* sp. KF913249, in a solution of 1 mM AgNO₃ in 100 mL distilled water at 27°C. It is proposed that *Aspergillus* sp. produces AgPgNps due to its ability to produce reducing enzymes with secondary metabolites, which could synergistically generate enough reducing potential to reduce AgNO₃ (+1 oxidation state) into

AAAAATAGTGTTATCATGATTGATACCCTACCTGTCTAAGTATAAGCAATTTATACG GTGAAACTGCGAATGGCTCATTAAATCAGTTATCGTTTATTTGATAGTACCTTACTAC ATGGATACCTGTGGTAATTCTAGAGCTAATACATGCTAAAAACCTCGACTTCGGAAG GGGTGTATTTATTAGATAAAAAAACCAATGCCCTTCGGGGGCTCCTTGGTGAATCATAA ATCAACTTTCGATGGTAGGATAGTGGCCTACCATGGTGGCAACGGGTAACGGGGAA GATACGGGGCTCTTTTGGGTCTCGTAATTGGAATGAGTACAATCTAAATCCCTTAAC GAGGAACAATTGGAGGGCAAGTCTGGTGCCAGCAGCCGCGGTAATTCCAGCTCCAA TAGCGTATATTAAAGTTGTTGCAGTTAAAAAGCTCGTAGTTGAACCTTGGGTCTGGC TGGCCGGTCCGCCTCACCGCGAGTACTGGTCCGGCTGGACCTTTCCTTCTGGGGAAC CTCATGGCCTTCACTGGCTGTGGGGGGGAACCAGGACTTTTACTGTGAAAAAATTAGA GTGTTCAAAGCACGCCTTTGCTCGAATACATTAGCATGGAATAATAGAATAGGACGT GCGGTTCTATTTTGTTGGTTTCTAGACCGCCGTAATGATTAATAGGGATAGTCGGGG AAAGCATTCGCCAAGGATGTTTTCATTAATCAGGGCACGAAAGTTAGGGGATCGAA GACGATCAGATACCGTCGTAGTCTTAACCATAAACTATGCCGACTAGGGATCGGGC GGTGTTTCTATGATGACCCGCTCGGCACCTTACGAGAAATCAAAGTTCTTGGGTTCT GGGGGGAGTATGGTCGCAAGGCTGAAACTTAAAGAAATTGACGGAAGGGCACCAC AAGGCGTGGAGCCTGCGGCTTAATTTGACTCAACACGGGGAAACTCACCAGGTCCA GACAAAATAAGGATTGACAGATTGAGAGCTCTTTCTTGATCTTTTGGATGGTGGTGC A TGGCCGTTCTTA GTTGGTGGAGTGATTTGTCTGCTTA A TTGCGATAACGA A CGAGA CCTCGGCCCTTA A ATA GCCCGGTCCGCATTTGCGGGCCGCTGGCTTCTTA GGGGGAC TATCGGCTCAAGCCGATGGAAGTGCGCGGCGAATAACAGGTCTGTGATGCCCTTAGAT GTTCTGGGCCGCACGCGCGCTACACTGACAGGGCCAGCGAGTACATCACCTTGGCC GAGAGGTCTGGGTAATCTTGTTAAACCCTGTCGTGCTGGGGGATAGAGCATTGCAATT ATTGCTCTTCAACGAGGAACGCCTAGTAGGCACGAGTCATCAGCTCGTGCCGATTAC GTCCCTGCCCTTTGTACACACCGCCCGTCGCTACTACCGATTGAATGGCTCGGTGAT ATGTTACCCCGGGCTCCTGCCAGTATGCTACCCGGTTGATCCTCCCAGTATGCTATCC TGGTGTCCGTCACCGG

Figure 1 The Basic Local Alignment Search Tool analysis of the above sequence shows maximum similarity with Aspergillus terreus.

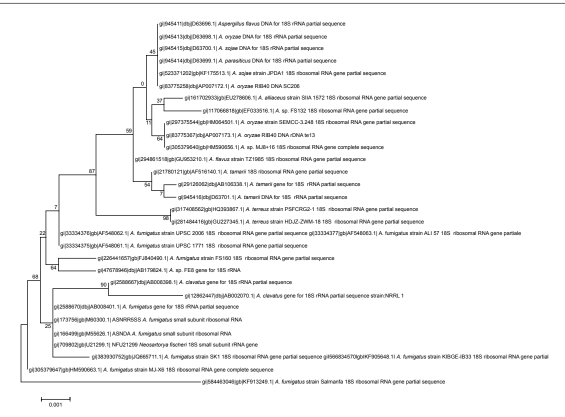


Figure 2 Phylogenetic tree showing interrelationship of Aspergillus luteus (KF913249) with closely related species of Aspergillus sp. and other related genera. Notes: The tree is generated by neighbor joining of top 27 sequences from BLAST using FigTree software. Bootstrap values expressed as percentage of 1,000 replications are indicated on nodes. The name of each type strain for comparative position with isolate is shown in parenthesis. The isolated strain represented by its accession number KF913249 and its distinct position in the radii of phylogenetic tree indicates a novel species of Aspergillus.

Ag (0 oxidation state) NPs in an aqueous medium. Reaction in the absence of salt confirmed the efficacy of mycelia in the synthesis of AgPgNps. Likewise, incubation of fungal mycelia alone in distilled water additionally did not create any assimilation peak(s) distinctive for AgPgNps. The characteristic absorption maximum (Figure 3A) centered at 416 nm attributed to the surface plasmon resonance (SPR) band of the AgPgNps. TEM analysis of the samples determined the average size of NPs to be 23 ± 2 nm (Figure 3B). Further analysis of the TEM micrograph using Gatan Digital-Micrograph showed the pentagonal shape of NPs. The TEM micrographs showed distinct well-separated particles of as-synthesized AgPgNps. Zeta potential of AgPgNps was found to be -12.6 ± 0.2 mV (Figure 3C), which made them stable for longer periods of time; the absence of aggregates is attributed to the electrostatic repulsive forces due to like forces presented by individual NPs.

Fourier transform infrared spectral analysis (Figure 3D) of AgPgNps presented peaks centered at 1,643.24 cm⁻¹ characteristics of C=O of amide groups of the amide I linkage and amide II band presents as a medium-wide shoulder at 1,537.02 cm⁻¹. The bands of amides I and II appear due

to carboxyl stretch and N–H deformation vibrations in the amide bond of the proteins, which encapsulated AgPgNps.³⁰ In addition, peak at 3,296.5 cm⁻¹ confirms the N–H stretching vibration. This mode of vibration does not depend on the backbone conformation but is very sensitive to the strength of a hydrogen bond. Furthermore, a peak at 1,081.8 cm⁻¹ shows (C–O–C/C–OH) C–O stretching³⁰ of alcohol and ether group with C-N (aliphatic amine) stretching vibration. Due to the presence of various secondary metabolites, a peak at 2,127.39 cm⁻¹ represents C≡C stretch of alkynes. Terminal-free hydroxyl (O–H) is represented by representative peaks 3,756.013 cm⁻¹ and 3,868.76 cm⁻¹.

Antibacterial activity

In this study, the antimicrobial activity of AgPgNps was evaluated. The AgPgNps displayed microbicidal activity toward a range of different pathogenic microorganisms (Figure 4). The MIC₅₀ values were determined against pathogenic bacteria, which included *Bacillus* sp. (12.781 µg/mL), *S. aureus* (9.230 µg/mL), *E. coli* (5.063 µg/mL), and *K. pneumoniae* (5.426 µg/mL), suggesting its broad-spectrum nature. This analysis further proved that AgPgNps displayed

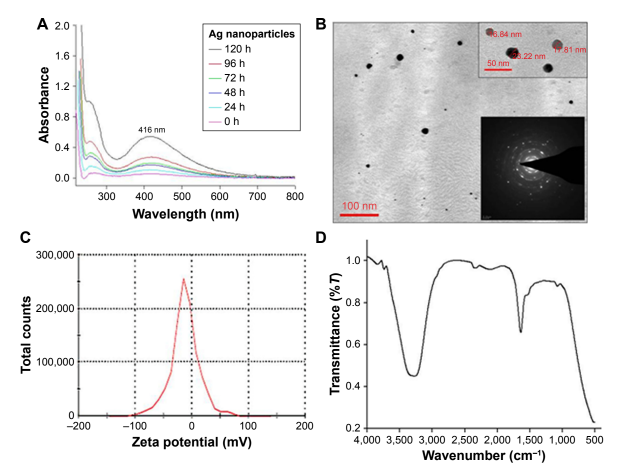


Figure 3 Characterization of AgPgNps by (A) UV-visible spectroscopy, (B) TEM and SAED (inset), (C) zeta potential analysis, and (D) FT-IR spectroscopy. Abbreviations: AgPgNps, pentagonal silver nanoparticles; UV, ultraviolet; TEM, transmission electron microscopy; FTIR, Fourier Transform infrared spectroscopy; SAED, selected area electron diffraction.

better efficacy toward Gram-positive strains as compared to the Gram-negative counterparts.

This analysis was performed by a method adapted from the method reported by Deda et al.³¹

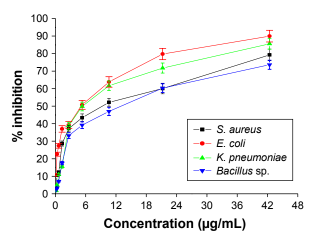


Figure 4 Graph showing antibacterial potential of AgPgNps against Bacillus sp., Klebsiella pneumonia, Escherichia coli, and Staphylococcus aureus. Abbreviation: AgPgNps, pentagonal silver nanoparticles.

Since ancient times, silver has enjoyed being the inorganic antibacterial agent of choice to fight against infections and spoilage. The antibacterial properties of silver NPs, which make it an ideal biocidal agent, are thought to be due to the slow oxidation exhibited by silver following the liberation of monovalent silver ion (Ag⁺) into the environment.¹⁹ Studies have well established the sizemediated antibacterial properties of silver NPs19,20 in detail, but very little is known about shape-dependent antimicrobial activity. However, shape-dependent mode of interaction of these NPs with the Gram-negative organism E. coli has been found to be dependent on the shape of NPs.¹³ In addition, some researchers have attempted to elucidate the mechanism responsible for the greater bactericidal property of NPs toward G⁻ than that toward G⁺ bacteria. In general, a widely studied and accepted hypothesis is that generation of ROS is responsible for cellular toxicity exhibited by metal NPs (ROS).¹⁸ The microbicidal property exhibited by silver NPs has been widely believed to be due to loss of subcellular materials caused by the creation of pits in the

cell membrane. This continues with the dehydrogenases involved in the respiratory chain getting inhibited, affecting the growth of cells altogether. Meanwhile, some proteins and phospholipids may work together to initiate the collapse of the membrane, eventually leading to cell decomposition and death.³² The Gram-positive bacterial cells respond differently following exposure to metal NPs.^{32–35} The most recent mechanism states that silver NPs target both the secondary (α -helix) and primary structures of the bacterial cell wall, by making connections on both the surface of the peptide and glycan ports of a bacterial cell wall, leading to the generation of "pits" on it. Silver NPs connect to β –1/4 bonds of *N*-acetylglucosamine and *N*-acetylmuramic acid of glycan strands, destroying their linkage and releasing them into the growing environment.³⁶

The cytotoxic effect of AgPgNps was tested on A549 and HeLa cell lines using MTT assay where primary osteoblasts were taken as control. The viability of these cells was measured at various concentrations (0.373, 0.186, 0.0932, 0.046, 0.0233, and 0.0116 μ g/mL) of AgPgNps (Figure 5). The results clearly illustrated that HeLa cells (Figure 5) rapidly lost their viability in comparison to A549 cells when incubated with AgPgNps in the concentration range of 0.0116–0.046 μ g/mL, which was trailed by an attenuated decrease when the NPs were used at higher concentrations from 0.046 to 0.373 μ g/mL. Interestingly, these NPs were found to be less cytotoxic to primary osteoblasts (Figure 5), resisting to a higher permissible biological limit.

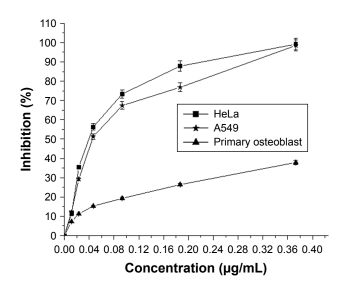


Figure 5 The cytotoxicity (dose dependent) study of AgPgNps against HeLa, A549, and primary osteoblasts.

Notes: AgPgNps inhibited cellular development considerably of HeLa and A549 with IC_{s0} values 0.038 μ g/mL and 0.044 μ g/mL correspondingly but did not influence primary osteoblasts cells extensively. All the information was represented in mean \pm SD of three examinations.

Abbreviation: AgPgNps, pentagonal silver nanoparticles.

In addition, when 70% confluent HeLa and A549 cells were incubated with AgPgNps for 48 hours, the phase-contrast microscopic images (Figure 6) showed changes in cellular morphology where majority of the cells of HeLa (Figure 6B) and A549 (Figure 6D) experienced changes in shape turning circular, with failure of membrane integrity, inhibition of cell growth, condensation of cytoplasm, and clumping of cells. On the other hand, unimportant changes in morphology were observed for primary osteoblasts (picture not shown) and untreated cells (Figure 6A and C). The quantification of

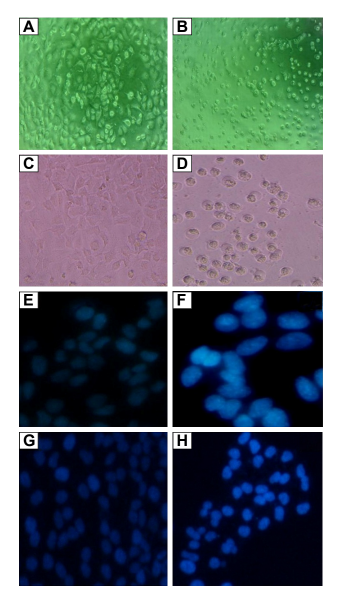


Figure 6 Images showing cytotoxic effect of AgPgNps on HeLa and A549 cell lines at $20 \times$ magnification.

Notes: HeLa cells: (**A**) control and (**B**) treated with AgPgNps. A549 cells: (**C**) control and (**D**) treated with AgPgNps. DAPI staining of HeLa cells: (**E**) control and (**F**) treated with AgPgNps. DAPI staining of A549 cells: (**G**) control and (**H**) treated with AgPgNps. Primary osteoblasts were used as control and were unaffected up to a substantial concentration of AgPgNps (picture not shown).

Abbreviations: AgPgNps, pentagonal silver nanoparticles; DAPI, 4',6-diamidino-2-phenylindole. these morphological changes was performed by means of ImageTool software. The extent of roundedness of the cell was quantified to measure the change in the morphology of the cells. Furthermore, the mechanism of cellular uptake and internalization of AgPgNps was analyzed using a fluorescent stain DAPI. Caveolae-mediated endocytosis is implicated in the internalization of anionic AgPgNps. These are microdomains produced in the caveolar vesicles that are generally made up of sphingomyelins (an amine group present on its polar domain), cationic lipids that intermingle and initiate the endocytosis of anionic NPs. Furthermore, NPs might also enter cells non-specifically through macropinocytosis, which is a vesicular transmonolayer transport that involves dynamin and microtubule network. AgPgNPs internalized through caveolae-mediated endocytosis enjoy improved bioavailability since they are prevented from getting degraded by avoiding interaction with lysosomes after their entry. This improved the efficacy of these NPs by enabling them to interact with subcellular structures further down. After entering into cells, AgPgNps containing vesicles fused with caveosomes or multivesicular body. Subsequently, caveosomes containing AgPgNps moved along with microtubules to the endoplasmic reticulum (ER),³⁵ eventually leading to their entry into the nucleus through nuclear pore complex.^{37,38} In addition, clathrin-caveolae-independent endocytosis

involved Arf6-, Cdc42-, and Rhoa-dependent pathways³⁹ and cholesterol with specific lipid compositions. Hence, AgPgNps causes apoptosis by condensing chromatin, which can be observed by DAPI analysis.

It is a well studied and accepted fact that pro-inflammatory and toxic activities of inorganic NPs including silver NPs are brought through the production of ROS. When human cells are exposed to oxidative stress, they result in an increase in the transcription of proinflammatory cytokines inducing apoptosis brought about by the c-Jun N-terminal kinase pathway.^{40,41} The silver NPs are known to be cytotoxic, genotoxic, and antiproliferative because ideally, DNA-damaging agents induce genome instability, which is a predisposing factor in carcinogenesis.

AgPgNps augmented intracellular ROS generation in HeLa and A549 cells

The amount of intracellular ROS generated actually dictates the caspase-dependent and caspase-independent cell death. The production of intracellular ROS has been implicated in cell death induced by both caspase-dependent and independent forms.^{42,43} In order to corroborate the mechanism of action of AgPgNps, the intracellular ROS levels of both AgPgNp-treated and -untreated HeLa (Figure 7A and B) and A549 cells (Figure 7C and D) were determined by the

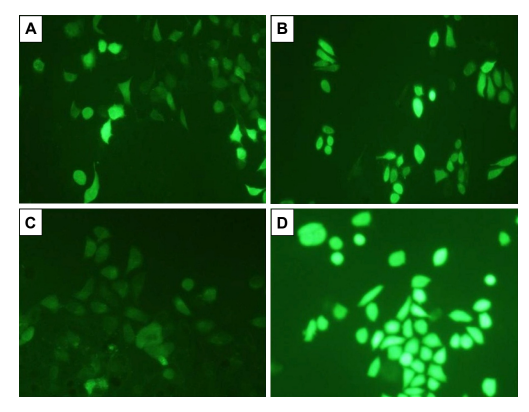


Figure 7 (Continued)

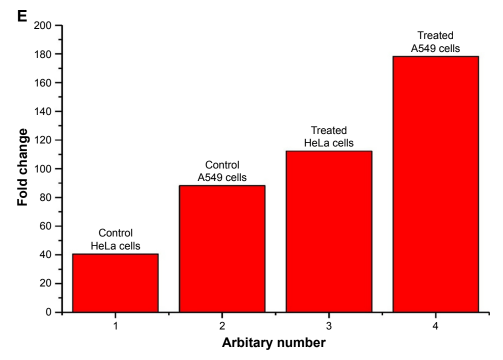


Figure 7 DCFDA staining for ROS generation on HeLa cells (20×), (**A**) control and (**B**) treated with AgPgNps, and A549 cells, (**C**) control and (**D**) treated with AgPgNps. Scale bar – 100 μm. Information demonstrated are illustrative of three independent experiments. (**E**) Quantification of ROS level in terms of intensity fold change of DCFDA fluorescence in HeLa- and A549-treated cells in comparison to untreated control cells. **Note:** Values are communicated as mean±SEM of three autonomous experiments.

Abbreviations: AgPgNps, pentagonal silver nanoparticles; SEM, standard error of the mean.

DCFH-DA method. Briefly, HeLa and A549 cells were monitored for changes in intracellular ROS levels by staining them with DCFH-DA after12 h of treatment. AgPgNp-treated cells (Figure 7A) demonstrated more grounded DCF-fluorescence intensity in both HeLa and A549 cells, which suggested an upgraded intracellular ROS level (Figure 7A). The above perceptions were further approved by the quantitative estimation of the ROS level, which demonstrated that intracellular ROS level was raised by approximately 2.76 folds (HeLa cells) and 2.01 folds (A549) when contrasted with control when both HeLa and A549 cells were treated with 0.046 μ g/mL of AgPgNps separately. Consequently, both the above outcomes recommended that AgPgNps enhanced ROS development in both the cell lines in a dose-dependent manner (Figure 7E).

Conclusion

We report the isolation of a fungus from Integral University campus and characterization using molecular and biochemical methods. This strain of fungus is capable of withstanding high concentrationd of metal ions. Hence, this fungus was used for the production of AgPgNps. These biogenic NPs showed effective antibacterial activity and anticancer activity in a dose-dependent manner. Additional researches are expected to evaluate the toxicity in detail and the mechanism implicated in the antibacterial and anticancer potential of the synthesized AgPgNps, which may propel these NPs as a potent nanomedicine.

Acknowledgments

The authors would like to extend their sincere appreciation to the Deanship of Scientific Research at King Saud University for its funding through research group number (RG-1438-069). The authors acknowledge Integral University and Department of Biosciences for providing facilities and support for the research.

Disclosure

The authors report no conflicts of interest in this work.

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