Optimization of the Synthesis of Fungus-Mediated Bi-Metallic Ag-Cu Nanoparticles

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Abstract: Bi-metallic nanoparticles (NPs) have appeared to be more efficient as antimicrobials than mono-metallic NPs. The fungus Aspergillus terreus-mediated synthesis of bi-metallic Ag-Cu NPs was optimized using response surface methodology (RSM) to reach the maximum yield of NPs. The optimal conditions were validated using ANOVA. The optimal conditions were 1.5 mM total metal (Ag + Cu) concentration, 1.25 mg fungal biomass, 350 W microwave power, and 15 min reaction time. The structure and shape of the synthesized NPs (mostly 20–30 nm) were characterized using several analytical tools. The biological activities of the synthesized NPs were assessed by studying their antioxidant, antibacterial, and cytotoxic activity in different NP concentrations. A dose-dependent response was observed in each test. Bi-metallic Ag-Cu NPs inhibited three clinically relevant human pathogens: Klebsiella pneumoniae, Enterobacter cloacae, and Pseudomonas aeruginosa. Escherichia coli, Enterococcus faecalis, and Staphylococcus aureus were inhibited less. The DPPH and hydrogen peroxide scavenging activities of the NPs were high, reaching 90% scavenging. Ag-Cu NPs could be studied as antimicrobials in different applications. The optimization procedure using statistical analyses was successful in improving the yield of nanoparticles.

Keywords: bi-metallic nanoparticles; RSM; optimization; Ag-Cu; microwave power

1. Introduction

Nanotechnological solutions are increasingly developed to be used as antimicrobial agents in biomedicine. Physical and chemical techniques to produce nanoparticles (NPs) have largely been inserted with eco-friendly biological techniques [1,2]. Nowadays, nanotechnology offers eco-friendly solutions not only for biomedicine but for agriculture and the remediation of polluted environments [3–5].

The use of fungi in the synthesis of NPs has gained much attention recently. Fungi have been found to be efficient in reducing metal salts to NPs by producing proteins and catalysts that enable rapid synthesis [6,7]. Fungi are especially suitable due to their metal resistance and ability to accumulate metals [8]. Moreover, fungi-mediated NPs are of controlled size and morphology [9–13]. As a whole, the fungi-mediated synthesis of NPs offers a technology that can be scaled up to industrial needs.

Nanoparticles can be synthesized using one or more metals. Recent research has recognized the positive synergistic effects of the metals in bi-metallic NPs [14,15]. The shapes of bi-metallic NPs are diverse and the ratio of surface to volume is high compared to mono-metallic NPs [16]. Moreover, bi-metallic NPs have higher stability and activity than mono-metallic NPs [17,18]. Several combinations of metals such as Ag-Cu, Au-Pt, Co-Ni, Ag-Pd, and Fe-Pt have been presented [17–21]. Ag-Cu NPs have been reported for structural and thermal stability [19–21].

Various techniques to synthesize NPs have been studied. Recently, the advantage of microwave reactors in the synthesizing process has been presented [22,23]. The efficient penetration of microwaves enables the homogeneous, rapid heating of the materials. The
use of microwaves produces a high yield of NPs and needs less time and chemicals. Thus, the microwave method fulfills the principles of green chemistry.

In the synthesizing process, many parameter values such as the microwave power, the amount of biomass used, the concentration of metals, and the reaction time must be chosen. It is crucial, yet seldom carried out previously, to optimize the conditions by modeling. One method to optimize the conditions is response surface methodology (RSM) [3], which has also been taken into use in NP synthesis recently [24,25]. The advantage of optimizing the conditions is studying several parameters and choosing the best combination of the parameter values to gain the maximum yield of NPs.

The objectives of this study were to combine two novel aspects in nanoparticle research: to optimize the process parameters using statistical methods and to produce bimetallic Ag-Cu NPs that have high biological activities. Treatments to optimize four parameters of the NP synthesizing process were carried out. Then, tests to study the antimicrobial, antioxidant, and cytotoxic activities of the synthesized Ag-Cu NPs were carried out.

2. Material and Methods

2.1. Synthesis of Ag/CuO NPs

The fungus Aspergillus terreus was isolated from the southern Red Sea coast, Saudi Arabia, as described by Ameen et al. [26]. The fungus was identified using molecular methods as described by [27].

To produce Ag/CuO NPs, metal salts (AgNO₃ and Cu(NO₃)₂) and fungal biomass were mixed and kept in a microwave (MW) reactor under stirring for a specific time, as described by Cuevas et al. [28]. The parameter values were optimized and are described above. The final turbid or precipitate was collected and washed thrice to remove unwanted metal salts and biomass. The final pellet was calcinated at 450 °C, and the purified NPs were stored in an airtight container.

2.2. Experimental Design and Statistical Analysis

An experiment using a single factor design was carried out. The experiment was carried out using the Box–Behnken experimental design [29]. A total of 2⁷ treatments were carried out with the different process parameter values presented in Table S1 (supplementary file). Response surface methodology (RSM) was used to optimize the parameter values, as described by [30]. The adequacy of the model was analyzed with ANOVA using Minitab (version 13) software. The significance level was set to p < 0.05. A graphical presentation based on quantile charts was used to describe the optimization.

An empirical second-order polynomial model with three parameters was used with the equation:

\[
\text{Yield} = \beta_0 + \beta_1 \times_1 + \beta_2 \times_2 + \beta_3 \times_3 + \beta_{11} \times_1^2 + \beta_{22} \times_2^2 + \beta_{33} \times_3^2 + \beta_{12} \times_1 \times_2 + \beta_{13} \times_1 \times_3 + \beta_{23} \times_2 \times_3
\]

where yield is NP yield (response variable); \(\beta_0\) is interception coefficient; \(\beta_{11}, \beta_{22}, \text{ and } \beta_{33}\) are quadratic terms; \(\beta_{12}, \beta_{13}, \text{ and } \beta_{23}\) are interaction coefficients; and \(\times_1, \times_2, \text{ and } \times_3\) are independent variables (fungal biomass (Extract), metal concentration (Conc), microwave power (Power), reaction time (Time)). The significance of the RSM model based on ANOVA is presented in Table S2 (supplementary file). The final equation is as follows:

\[
\text{Yield (mg)} = -73 + 50 \text{ Extract} + 40 \text{ Conc} + 0.09 \text{ Power} + 4.2 \text{ Time} - \ast \text{Extract} 16.50 - , \ast \text{Conc} - 0.0002 , \ast \text{Power} - 0.2 \\
\ast \text{Time (min)} - 5.3, 11.15 \text{ Extract} , , \text{Conc} - 0.004 , \text{Power} - 0.6, + 0.2 \text{ Time}
\]

2.3. Ag/CuO NPs Characterization

The formation of Ag/CuO NPs was monitored using a UV-Vis spectrophotometer (UH3500, Hitachi, Tokyo, Japan). Reducing and capping agents present in the fungal biomass were analyzed using a Fourier transform infrared spectrophotometer (FT-IR, Thermo Fisher Scientific, Massachusetts, United States). The powder X-ray diffraction
technique was used to analyze the structure and nature of the synthesized Ag/CuO NPs using CuKα radiation. The average size of the NPs was calculated using the Scherrer equation:

\[ D = \frac{k\lambda}{\beta \cos \theta} \]

where \( k \) is the shape factor (0.9), \( \beta \) the peak’s full width at half-maximum (FWHM), \( D \) the average size, \( \lambda \) the wavelength, and \( \theta \) is the diffraction angle.

The morphology of the synthesized NPs was observed using a scanning electron microscope (SEM, ZEISS EVO, Massachusetts, United States) coupled with energy dispersed X-ray spectroscopy (EDS) and a transmission electron microscope (TEM), Phillips CM200. The particle size histogram was calculated using ImageJ software and plotted using Origin Pro 8.5 software [31].

2.4. Biological Activities of the Ag/CuO NPs

All biological tests were carried out as three replicates. The antioxidant activity analysis using the free radical 2,2-diphenyl-1-picrylhydrazyl (DPPH) was carried out as described by [32] with slight modifications. The NPs solution was mixed with 1 mL of freshly prepared DPPH (1 mM) in methanol. The test solution was kept in the dark at room temperature for 30 min, and the absorbance was recorded using UV-Vis spectroscopy at 517 nm.

Hydrogen peroxide scavenging by the NPs was carried out as described by Fowsiya and Madhumitha [33]. Ascorbic acid was used as the standard. Scavenging was calculated using the following equation:

\[ \% \text{ Scavenging} = \frac{\text{Ac} - \text{As}}{\text{Ac}} \times 100 \]

where \text{Ac} and \text{As} indicate the absorbance of the control (phosphate buffer solution) and sample solution/standard solution. The lowered absorbance indicates the scavenging of free radicals by NPs.

The antibacterial activity of the NPs was analyzed using six bacterial species (Klebsiella pneumoniae, Enterobacter cloacae, Pseudomonas aeruginosa, Escherichia coli, Enterococcus faecalis, and Staphylococcus aureus) collected from the microbiology department of King Saud University and identified molecularly [34]. The inhibition of NPs was studied with the well-diffusion method [34]. The bacteria were first incubated at 37 °C for 24 h. NPs at 50 \( \mu \)g/mL concentration in dimethyl sulphoxide (DMSO) were added to wells in agar. The zone of inhibition (mean of width and length in mm) around the wells was measured using a ruler after the incubation at 37 °C for 24 h. The minimum inhibitory concentration (MIC) and the lethal minimum bactericidal concentration (MBC) were determined using various concentrations (1, 5, 10, 15, 20, 25, 35, 45, 50) of the NPs.

The cytotoxicity test was carried out as described previously [35]. The cells were kept in D-MEM medium enhanced with 10% fetal calf serum (FCS), 1 mM sodium pyruvate, 2 mM glutamine, 50 U/mL penicillin, 50 mg/mL streptomycin, and 100 mM trivial amino acids. For morphological examinations, the cells were cultivated in 6-well plates at a centralization of 3 × 105 cells/well in 2 mL of complete medium. The cells were brooded in a humidified hatchery at 33 °C and 5% CO2. The MTT test was utilized to quantify mitochondrial movement, as recently portrayed [34], with minor modifications. The cell culture medium was disposed of, and cells were hatched with 100 \( \mu \)L of MTT for 2 h at 37 °C, 5% CO2. The MTT arrangement was then, at that point, disposed of, and 100 \( \mu \)L of DMSO was added to each well. Optical density was perused on a microplate per user at 550 nm, with a reference at 655 nm (EL800, Bio-Tek Instruments, Inc., Vermont, USA). Cell death was determined as the proportion of the mean of OD obtained for each condition to that of the control (no molecule) condition:

\[ \% \text{Cell death} = \frac{\text{OD (s)}}{\text{OD (c)}} \times 100 \]

where \text{OD (s)} and \text{OD (c)} are the optical density of the sample and control.
3. Results

3.1. Optimization of Operating Variables

Three-dimensional response surfaces and two-dimensional contour plots were presented to describe the optimization: the diagrams illustrate the effects of two factors on NP yield at a time. Figure 1 shows the effect of the fungal biomass (extract) and the concentration of total metals on NP yield after 15 min with 350 W microwave power. Increasing the metal concentration from 0.5 mM to 1.5 mM doubled the NP yield. Increasing the fungal extract from 1 mg to 1.5 mg reduced the yield. When the concentration was 1 mM, the maximal yield was achieved between 300 and 500 W and between 0.8 and 1.25 mg extract (Figure 2).

![Figure 1. Fungal biomass (extract) and total metal concentration effect on NP yield.](image1)

A concentration of 1.5 mM caused a decline in the NP yield (Figure 3). Considering the reaction time, the optimal time was 15 min (Figure 4). When the MW power and the concentration were kept constant, first an increase and then a decrease in the NP yield were observed along with both time and extract (Figure 4a). Increasing the concentration from 0.5 mM to 1.5 mM increased the NP yield gradually when the extract and MW power were constant (Figure 4b). The optimal MW power was 350 W and time 15 min (Figure 4c). In summary, the yield was, at lowest, <15 mg and, at highest, 32 mg. The final conditions were 1.5 mM total metal concentration, 1.25 mg fungal biomass, 350 W, and 15 min.
Figure 3. Microwave power and total metal concentration effect on NP yield.

Figure 4. The effect of (a) time and fungal biomass (extract) (b) time and total metal concentration, and (c) time and microwave power on the yield of Ag-Cu NPs.

3.2. UV-Vis Spectroscopy

The UV-Vis spectra showed the maximum absorbance after 60 min (Figure 5). The formation of NPs started at 5 min and ended at 60 min.
3.3. FT-IR Analysis

The FT-IR spectrum showed vibrational stretching at 3432 cm\(^{-1}\) relating to -OH extending, 1638 cm\(^{-1}\) relating to C-C bond, and 663 relating to Cu ions (Figure 6.). The broad peak corresponding to –OH has been reduced in the NPs peak. Therefore, the -OH group acts as a good reducing, stabilizing, and capping agent of the NPs. The FT-IR spectrum was consistent with the UV-Vis spectrum.

Figure 5. UV-Vis spectra of Ag/CuO NPs formation.

Figure 6. FT-IR spectra of synthesized Ag/Cu NPs.
3.4. XRD Analysis

The X-beam diffraction pattern (Figure 7.) shows sharp crystalline peaks of Ag/CuO NPs. Six diffraction peaks at 37.2°, 45.45°, 48.3°, 65.21°, 75.1°, and 81.3° compare to the standard face-centered cubic designs. The intense peak at 48.36° affirmed the presence of AgO, and the peaks at 32.41° and 37.2° affirmed CuO.

![XRD pattern](image)

**Figure 7.** XRD pattern of fungal biomass-mediated Ag/Cu NPs.

3.5. Morphological Analysis

SEM images showed the agglomerated and spherical shapes of the bi-metallic NPs (Figure 8a). From the TEM image, spherical NPs and the dispersion of agglomerated particles were observed (Figure 8b). The elemental analysis showed the presence of Ag, Cu, and O without any impurities (Figure 8c). The Scherrer equation gave the average NP size of 22 nm. The particle size histogram shows that the particle size 20–22 mm was the most common, having 11 counts (Figure 8d). Small (12–18 nm) and large (>30 nm) were in the minority with fewer than 10 counts combined.

![SEM, TEM, EDS, and particle size histogram](image)

**Figure 8.** Images of SEM (a), TEM (b), and EDS (c) and the particle size histogram (d) of Ag/Cu NPs.
3.6. Antioxidant Activity

The DPPH scavenging activity increased with the NP concentration (Figure 9a). Hydrogen peroxide scavenging inhibition increased from 30% in 100 mg/L NP to 75% in 500 mg/L NP (Figure 9b).

![Figure 9. DPPH (a) and hydrogen peroxide (b) scavenging activity of Ag-Cu NPs (mean and SD, n = 3).](image)

3.7. Antibacterial Activity

The highest inhibition of the NPs was observed against *K. pneumonia*, *E. cloacae*, and *P. aeruginosa*. Their inhibition zones varied between 9 mm and 22 mm and were almost double those of *E. coli*, *E. faecalis*, and *S. aureus*, which varied between 3 mm and 12 mm (Table 1). The inhibition was about double for each bacterium at the highest NP concentration (5 μg/mL) to the lowest concentration (0.5 mg/mL). The MIC varied between 0.2 μg/mL against *E. cloacae* and *K. pneumoniae* and 1 μg/mL against *E. faecalis* (Table 2). For *S. aureus*, the MIC was 1 μg/mL and for *P. aeruginosa* 0.3 μg/mL. The MBC was lowest (0.5 μg/mL) to *P. aeruginosa* and highest (1.5 μg/mL) to *S. aureus* (Table 2).

![Table 1. Antibacterial activity (as the zone of inhibition, mean ± SD, n = 3) of Ag-Cu NPs against six bacterial species.](image)

| NPs Concentration (μg/mL) | Zone of Inhibition (mm) against Bacteria |
|---------------------------|----------------------------------------|
|                           | *K. pneumoniae* | *E. cloacae* | *P. aeruginosa* | *E. coli* | *E. faecalis* | *S. aureus* |
| 0.5                       | 10 ± 1.6       | 11 ± 1.5      | 9 ± 1.6         | 6.5 ± 2.4  | 3 ± 1.5       | 6 ± 1.4     |
| 1                         | 12 ± 1.5       | 13 ± 1.2      | 11 ± 1.8        | 7.6 ± 2.1  | 4 ± 1.6       | 7 ± 1.5     |
| 2.5                       | 15 ± 1.7       | 15 ± 1.6      | 14 ± 2.1        | 9 ± 2.2    | 6 ± 1.6       | 9 ± 1.4     |
| 4                         | 19 ± 2.1       | 18 ± 2.2      | 16 ± 2.5        | 10 ± 1.2   | 9 ± 2.3       | 10 ± 2.6    |
| 5                         | 21 ± 2.2       | 22 ± 2.5      | 20 ± 2.5        | 12 ± 2.4   | 10 ± 2.6      | 15 ± 2.0    |

![Table 2. Minimum inhibitory concentration (MIC) and minimum bactericidal concentration (MBC) of Ag-Cu NPs against different bacterial species.](image)

| Bacteria            | NPs Concentration (μg/mL) |  |
|---------------------|---------------------------|---|
|                     | MIC          | MBC |
| *K. pneumoniae*     | 0.2          | 1   |
| *E. cloacae*        | 0.2          | 1   |
| *P. aeruginosa*     | 0.3          | 0.5 |
| *E. coli*           | 0.5          | 0.6 |
| *E. faecalis*       | 1            | 0.8 |
| *S. aureus*         | 0.8          | 1.5 |
3.8. Cytotoxicity Test

Bi-metallic NPs caused the highest cell death at a 5 μg/mL NP concentration, where 95% of the cells died in 48 h (Figure 10). At the highest NP concentration (50 μg/mL), cell death was lower, at 90%.

![Figure 10. Cytotoxicity at different concentrations of Ag-Cu NPs (mean and SD, n = 3) after 24 and 48 h.](image)

4. Discussion

The novelty of this work was to use microwaves and fungal biomass to synthesize bi-metallic Ag-Cu NPs and optimize the process conditions using statistical methods. The synthesis process depended on the amount of fungal biomass used, the concentration of metals, irradiation power, and the reaction time, which we optimized for the final synthesis.

The synthesized bi-metallic NPs showed antimicrobial activities against three clinically relevant Gram-negative bacterial pathogens, namely *K. pneumoniae*, *E. cloacae*, and *P. aeruginosa*. Three of the bacteria, Gram-negative *E. coli* and Gram-positive *E. faecalis* and *S. aureus* were inhibited slightly. The higher inhibition of Gram-positive bacteria has been proposed, and for instance, *E. coli* and *P. aeruginosa* were inhibited less than *S. aureus* by Zn and Cu NPs [36,37]. Our study does not support the separation between Gram-positive and Gram-negative bacteria. It may be that the NP characteristics may be of great importance, in addition to the resistance of the particular bacterial strain [38]. Ag NPs have been shown to efficiently inhibit *P. aeruginosa*, *E. coli*, and *S. aureus*. *E. coli* was inhibited, for instance, by Zn NPs [39] and Ag NPs [40]. Moreover, meticillin-resistant *S. aureus* (MRSA) has recently been shown to be inhibited by NPs [41–43]. For some reason, our bi-metallic NPs did not inhibit *S. aureus* especially efficiently. More than double the minimum inhibitory concentration and thrice the minimum bactericidal concentration were needed to inhibit *S. aureus* compared to *P. aeruginosa*. All bacterial species that we tested have been inhibited by some metal NPs previously, at least to some extent [44–48].

Bi-metallic NPs are promising antimicrobial agents, especially against multidrug-tolerant pathogenic microorganisms [49]. The bacteria *S. pneumonia*, *P. aeruginosa*, *E. coli*, *Bacillus subtilis*, and *S. aureus* were inhibited efficiently by Ag-Cu NPs previously [50–52]. The ultra-high antibacterial activity of 2 nm size Ag-Cu NPs [53] was observed against *E. coli* and *S. aureus*. Ag-Cu NPs have been proposed to be useful to inhibit bacteria in food packages [54–56].
In general, NPs with a size of 1–100 nm have shown strong antimicrobial effects against both Gram-positive and Gram-negative bacteria [57]. Smaller particles are more efficient because the ratio of the surface region to the volume increases when the particle size diminishes. Our NPs were relatively small, varying between 12 and 30 nm. Although only the dynamic light scattering (DLS) method would give a precise size of the NPs, we suggest that the process we propose here is suitable for producing small and efficient bi-metallic NPs.

Microwave power and optimization of the synthesizing conditions using RSM have been carried out only recently. By optimizing the process, very small Se and Cu nanoparticles (1–10 nm) were synthesized using a microwave power of 790 W for 15 min [58,59]. In our tests, the highest power, 500 W, was not the most efficient in producing bi-metallic Ag-Cu NPs. Instead, the highest yield was reached in 15 min with 350 W. Elsewhere, a constant 800 W power, and the maximum tested time (6 min) produced Ag NPs of 46 nm size [60]. Relatively small Fe NPs (14 nm) were produced in 10 min [61]. We were able to synthesize smaller than 30 nm bi-metallic NPs. It is possible that testing a larger scale of parameter values would produce smaller NPs, which is a future challenge.

5. Conclusions

The synthesis of Ag-Cu NPs was optimized using statistical methods. The optimal conditions of four process parameters (fungal biomass, microwave power, metal concentration, and reaction time) doubled the yield of nanoparticles when compared to starting conditions. The optimal synthesis produced bi-metallic nanoparticles with a size of 20–30 nm. The nanoparticles had antibacterial, antioxidant, and cytotoxic properties. Three of the six tested bacterial pathogens were remarkably inhibited in our tests, while the cytotoxicity and antioxidant activities were high in general. Smaller particles with higher activities might be synthesized using a wider scale of process parameter values and by further optimizing the process. It seems evident that the optimization procedure should be included in each future nanoparticle synthesis study.

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