Gamma Ray-Induced Synthesis of Silver Nanoparticles using Bacterial Cellulose as a Multi-Functional Agent and its Application

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Abstract

Antibacterial coatings based on bacterial cellulose (BC) have been widely used in many fields including food packaging and wound dressing. In this study, we aimed to synthesis of colloidal AgNPs and BC/AgNP composite by using BC as a reducing and capping agent in one step reaction induced by gamma-ray. Bacterial strain *Komagataeibacter rhaeticus* N1 MW322708 was used for biosynthesis BC by inoculation on Hestrin and Schramm medium and incubated statically at 35 °C for 10 days. BC sheet was formed, harvested, purified, and dried, then used for the synthesis of AgNPs and BC/AgNP by soaked 0.05 g of dried BC in 10ml of 1mM aqueous AgNO$_3$ solution for 2h and then irradiated by gamma-ray under different doses. Color change from yellow to deep brown indicated the synthesis of AgNPs and BC/AgNP. The optical spectra of synthesized AgNPs revealed that the surface plasmon resonance was localized around 420 nm. DLS analysis showed that the mean diameter of AgNPs was 49.5 nm with a -19.36-mV value of zeta potential. TEM images revealed the spherical shape of synthesized AgNPs. The results of FESEM, FTIR, and XRD confirmed the formation of BC/AgNO$_3$ composite. The highly crystalline nature of the BC membrane and BC/AgNP composite was observed in XRD measurements with a crystal size of 5.416 and 5.409 nm, respectively. The antibacterial activity of BC and BC/AgNP against pathogenic bacterial isolated from Pastirma food samples revealed that BC does not show antibacterial activity, while BC/AgNP composite showed antibacterial potency against *Staphylococcus aureus*, *Enterococcus faecalis*, *Listeria monocytogenes*, *Proteus mirabilis*, and *Escherichia coli*, with an inhibition zone of (mm) $9\pm1$, $9\pm0.57$, $10\pm1.15$, $8\pm0.5$ and $7\pm0.28$, respectively. We concluded that this novel method presented in this paper offers a promising route for both AgNPs and BC/AgNP composites synthesis using a green, renewable biopolymer as a multifunctional agent and potential to be applied in the future development of food packing, biomedical instruments, and therapeutics.

1. Introduction

Bacterial cellulose (BC) is a versatile biopolymer produced from the metabolic activity of cellulose-producing bacteria such as *Komagataeibacter*, *Agrobacterium*, and *Rhizobium* (Toda et al., 1997; Iguchi et al., 2000; Lin et al., 2013; Rajwade et al., 2015; Petrova et al., 2020). Cellulose synthesized by bacteria has a chemical composition similar to plant cellulose; however, thanks to its structural arrangements but, BC has remarkable physical properties. A study by Lin and the group showed that BC possesses high tensile strength, high crystallinity, high hydrophilicity, and excellent biocompatibility (Lin et al., 2013). BC has very high purity so that making other purification processes (such as delignification), which is very energy-consuming for plant cellulose, is unnecessary (Santoso et al., 2020). BC production gained more attention over the last several decades and its potential uses cover very varied fields including food (Padrao et al., 2016; Azeredo et al., 2019) and medical application (Moniri et al., 2017; Pal et al., 2017 and Jiji et al., 2020). The ability to use BC in diverse application resulting from its stunning properties that include high purity, biodegradability, three-dimensional (3-D) nanofiber, high water holding capacity, good mechanical properties, high crystallinity, and biocompatibility (Shi et al., 2014; Shabanpour et al., 2018; Wang et al., 2019), besides, the availability of incorporating nanostructured materials into BC 3-D network
structure make it attractive and appropriate for more application (Torres et al., 2019). Nanostructured materials and their applications represent an active area at the scientific and traditional levels (Xue et al., 2019). Nanoparticles and nanostructured materials have found their way into different fields such as food (Enescu et al., 2019), medicine (Jiji, 2020), environment (Umar et al., 2016), and so on. It is well known that silver nanoparticle materials have been proven to be the most useful because they showed great antimicrobial performance against a wide range of microorganisms including various bacteria and fungi (Siddiqi et al., 2018 and Wang et al., 2018). AgNPs are extensively used in industrial applications than any other nanomaterial (Vance et al., 2015). However, the aggregation tendency of AgNPs may frustrate their unique properties at the nanoscale. We can prevent aggregation by the incorporation of AgNPs into a nanoporous polymer (Cai, 2009). The 3-D network nanofiber of BC can make it easy and ensure a well-distribution of AgNPs through BC nanofiber and forming BC /AgNP composites (Ifuku et al., 2009; Wu et al., 2014; Torres et al., 2019). Thus BC /AgNP nano-composites aroused great interest among researchers in food packaging (de Moura et al., 2012; Adepu and Khandelwal 2018; Salari et al., 2018 and Azeredo et al., 2019) and wound dressing applications (Manerung et al., 2008; Pal et al., 2017; Tabaii et al., 2018; Horue et al., 2020; Jiji et al., 2020; Wan et al., 2020).

A variety of synthesis methods that including different reducing agent have been investigated to obtain BC/AgNP nano-composites which include, sodium borohydride (NaBH₄) (Maneerung et al., 2008; Pinto et al., 2009; Barud et al., 2011; Wang et al., 2020), sodium tripolyphosphate (TPP) (Tabaii and Emtiazi, 2018) and triethanolamine (TEA) (Barud et al., 2011). Some of these methods are cost, ecologically unfriendly, and unsafe that need to use toxic chemicals. Thus, the development of green approaches for the synthesis of colloidal AgNO₃ and BC /AgNP composite is necessary. Recently gamma radiation gains more attention in AgNPs synthesis. Gamma radiation was used to induces the reduction of Ag⁺ into metallic Ag in different aqueous solutions; acetic water solution containing chitosan (Chen et al., 2007), aqueous silk fibroin (SF) solution (Madhukumar et al., 2017), and poly (N-vinylpyrrolidone) solution (Dhayagude et al., 2018). The main objective of this work is to synthesis of colloidal AgNPs and BC /AgNP composite by green, eco-friendly, free of toxic approach, characterization, and evolution of the antibacterial activity against some pathogenic bacteria.

2. Materials And Methods

The most potent bacterial isolate was characterized and identified based on morphological, physiological, biochemical characteristics, 16S rDNA sequence analysis, and recorded in GenBank under accession number (MW322708) strain K. rhaeticus N1 MW322708.

https://www.ncbi.nlm.nih.gov/nuccore/MW322708.

2.1 Production and purification of bacterial cellulose

The bacterial strain Komagataeibacter rhaeticus N1 MW322708 strain isolated from Kombucha was used for BC synthesis. K. rhaeticus N1 MW322708 was inoculated into a sterile plastic box that contains 400 ml of Hestrin and Schramm (HS) medium (% w/v): 2% glucose, 0.5% peptone, 0.5% yeast extract,
0.27% Na$_2$HPO$_4$, and 0.15% citric acid (Hestrin & Schramm, 1954) the medium was modified by adding 1.5% ethanol to final concentration, the final pH was adjusted to 7, then *K. rhaeticus* N1MW322708 statically incubated at 35°C. BC sheet formed in air-liquid phase after 10 days were harvested, treated with 0.1 M NaOH solution at 85°C for 2h to remove the bacterial cells, then rinsed several times with deionized water to achieve a neutral pH; finally BC sheet was dried at 85°C for 1h and then subjected to autoclave.

### 2.2. Synthesis of colloidal AgNPs and BC/AgNP composite under gamma radiation

Purified BC sheets (0.05 g, dry weight) were soaked in 10 ml of 1 mM aqueous AgNO$_3$ solution for 2h then irradiated by gamma-ray under different doses (0.2, 0.4, 0.8, 1, 5, 10, 20, 40, 80 and 100 KGy). BC sheets were taken, rinsed several times with distilled water, and oven-dried at 40°C. Finally, the deep brown color solution and BC/AgNP sheet were stored in dark conditions at 4°C for future use.

### 2.3. Gamma radiation Source

Gamma irradiation was applied using a cobalt 60 irradiation source (Gamma cell 4000-A-India), located at the Egyptian Atomic Energy Authority (EAEA), Cairo, Egypt. The irradiation dose rate was 1.0 kGy/h at the time of the experiment and the irradiation process was carried out at ambient temperature.

### 2.4. Characterization of colloidal AgNPs and BC/AgNP composite

#### 2.4.1. UV–visible absorption spectra

UV–visible absorption spectra were recorded on a Cole Parmer spectrophotometer with spectra over a range of 300–800 nm.

#### 2.4.2. Dynamic Light Scattering (DLS) and Zeta Potential

DLS measurement was performed on a PSS-NICOMP 380-ZLS particle sizing system (St. Barbara, California, USA). The same instrument was used to obtain the Zeta Potential value. The results were obtained at 25°C.

#### 2.4.3. Transmission electron microscopy (TEM)

The morphology of the obtained AgNPs was obtained with a JEOL JEM-100CX (Japan) transmission electron microscopy.

#### 2.4.4. Field emission scanning electron microscopy (FESEM)

Morphological characterizations of BC and BC/AgNP composite samples were performed on a Zeiss (Sigma 300 VP; Germany) field emission scanning electron microscopy (FESEM).
2.4.5. Fourier transform infrared (FTIR)

Fourier transforms infrared (FTIR) spectrum was performed on ATI Mattson (Genesis series, Unicom, England). BC/AgNP sheet was scanned at the frequency range of 400 to 4000 cm\(^{-1}\).

2.4.6. X-ray diffraction analysis (XRD)

X-ray diffraction spectra of the dried BC and BC/AgNP composite were obtained using the XRD-6000 Shimdazu device (Japan). A standard Theta/2Theta diffractometer (using a copper X-ray source) was used. Scans were performed at 2 degrees per min from the diffraction angle ranged from 4 to 90°. The apparent crystal size (ACS) of BC and BC/AgNP was calculated using Scherrer's equation (Klug and Alexander, 1954), as follows:

\[
ACS = \frac{k\lambda}{\beta \cos(\theta)}
\]

Where \(k\) is the unknown shape factor and usually considered as 0.9, \(\lambda\) is the X-ray wavelength, \(\beta\) is the full width at half maximum in radians and \(\theta\) is the diffraction angle.

2.5. Antibacterial activity of BC and BC/AgNP composites

The antibacterial activity of prepared BC, BC/AgNP composites was determined using the disk diffusion method against Gram-positive (\textit{Staphylococcus aureus}, \textit{Enterococcus faecalis}, and \textit{Listeria monocytogenes}) and Gram-negative (\textit{E. coli} and \textit{Proteus mirabilis}) bacteria species were isolated from Pastirma food samples collected from local markets in Cairo, Egypt, and identified by VITEK 2 compact automated system (Biomerieux Inc., Marcy l’Etoile, France). Tested bacteria were cultured into a tube containing 5 ml of Muller Hinton broth (CLSI, 2016) at 37°C for 24 h, then 0.1 ml of bacterial suspension with 0.5 McFarland turbidity was swapped on the surface of the Mueller Hinton agar medium, dried discs (6mm in diameter) of BC/AgNP composite and pure BC were placed on the agar medium along with amoxicillin/clavulanic acid (AMC) (20/10µg/ml), ceftazidime (CAZ) (30µg/ml) and streptomycin (S) (10µg/ml) against tested bacteria. The petri-dishes were incubated at 37°C for 24h. The antibacterial activities of the samples against tested organisms were monitored by observing the zone of inhibition formed surrounding the disks. The inhibition zone diameter was measured (Patelet et al., 2017; Buszewski et al., 2018).

3. Results And Discussion

Bacterial isolates

The bacterial strain used in this study was isolated from kombucha tea in Egypt and identified by morphological, and biochemical characteristics as well as by 16S rRNA gene sequence analysis and deposit in GenBank under accession number (MW322708) strain \textit{K. rhaeticus} N1 MW322708. https://www.ncbi.nlm.nih.gov/nuccore/MW322708
Synthesis of bacterial cellulose

The BC produced by *K. rhatecus* in static culture is initially extruded from the pores on the cell surface as microfibres and results in the growth of a dense, white BC pellicle at the air-liquid interface of modified HS medium after 10 days of incubation. Figure (1) shows BC produced by *K. rhatecus* N1 MW322708 strain after 10 days using static culture conditions (a), harvesting of BC (b), and BC after purification and drying (8 g/l) (c). *Komagataebacter rhaeticus* is the most efficient BC producer, as it has the capacity to assimilate several different sugars and yields high levels of cellulose in a liquid culture medium (Rajwade et al., 2015; Petrova et al., 2020).

Synthesis of AgNPs and BC/AgNP composite under gamma-ray irradiation:

In this work, gamma-ray was attempted to induce the reduction of Ag⁺ to Ag⁰ by surface OH groups on BC surface for synthesis of colloidal AgNPs and BC/AgNP composite, with no chemicals involved in the chemical reaction or no surface modification of BC, just pure BC without any surface modification soaked in AgNO₃ solution under gamma-ray which promote the reduction process and were used for creating a hydrated electron and primary radicals in many studies (Chen et al., 2007; Park et al., 2012; Van Phu et al., 2014; Madhukumar et al., 2017; Dhayagude et al., 2018). It was found that no apparent color change of the BC membrane and solution at (0.2, 0.4, 0.8, 1, and 5 KGy). At 10 kGy visual observations showed that as the reaction started, the color shifted from pale yellow to deep brown for higher doses (20–100 kGy), signifying the reduction of Ag⁺ to Ag⁰ and formation of colloidal AgNPs and BC/AgNP composite. Recently, using of BC as a template in BC /AgNP composite synthesis gaining a lot of attention among researchers. The composite synthesis process and the mechanism of reaction in which the metal is reduced and binds onto BC surface have not been studied in much detail in many reports (Kaushik and Moores, 2016). Many researchers showed that it’s necessary to make the surface modification of BC by some compounds i.e. TEMPO (2,2,6,6-tetramethylpiperidine-1-oxyl radical), that oxidize BC and introducing surface-active carboxyl groups for BC /AgNP composite synthesis (Lal and Mhaske 2018; Elayaraja et al., 2020). Recently, Musino et al., (2021) showed that the hydroxyl groups on the surface of BC act as nucleation points for AgNPs through ion-dipole interaction, and OH groups represent the effective nucleation point for AgNPs synthesis on a BC solid surface. As far as we know it’s the first study for the synthesis of colloidal AgNPs and BC/AgNP composite using only BC as a reducing and stabilizing agent under gamma radiation.

UV–visible spectrum

Figure (2) showed UV–visible spectrum of AgNPs synthesized by BC under different doses of gamma-ray. The reduction of Ag⁺ to Ag⁰ by OH groups on BC surface under gamma-ray was preliminarily identified by observing the change in color of the reaction medium from clear to yellowish-brown. The change in color to the pale yellow of the reaction mixture started at a dose of 10 KGy and gradually turned to deep brown-at higher doses (20, 40, 80, and 100KGy). It is well known that AgNPs show a yellowish-brown color in water or aqueous solution; these colors arise due to the excitation of surface plasmon vibrations in the
metal nanoparticles (Shankar et al., 2004). This color change was associated with the development of a strong absorption band at 420 in the UV-vis. spectrum. This UV-Visible absorption peak at around 420 nm was attributed to the surface plasmon resonance (SPR) absorption peak of AgNPs, which confirmed the AgNPs formation with small size and narrow size distribution. Appeared peak around 420 nm was previously reported as a spherical or quasi-spherical Ag NPs SPR band (Sivasankar et al., 2018; Tabaii et al., 2018).

**DLS analysis**

To accurately measure the mean diameter of synthesized AgNPs DLS analysis was used. The results of the DLS analysis (Fig. 2) showed that the mean diameter of AgNPs was 49.5 nm. The size distribution of colloidal AgNPs illustrated by details in a table (1).

**Zeta potential analysis**

Zeta potential measures the electric charge on the surface of nanoparticles. Zeta potential value delivers information about the stability of nanoparticles. When nanoparticles in suspension have a large negative zeta potential value, nanoparticles will tend to repel with each other, thus there will be no tendency of the nanoparticles to agglomerate together. In contrast, in the case of low zeta potential values, no force to prevent the nanoparticles from coming together, thus nanoparticles tend to agglomerate (Roy et al., 2013). The value of the zeta potential of colloidal AgNPs was − 19.36 mV (Fig. 4). Obtained results from zeta potential proved that synthesized AgNPs were poly-dispersed, due to the high negative zeta potential value. The electrostatic repulsive force between nanoparticles results in the prevention of flocculation of nanoparticles and has an important role in nanoparticles' long-term stability in the solution (Kotakadi et al., 2016).

**TEM examination**

TEM examination of synthesized AgNPs was used to obtain information about the morphology and size of metal nanostructures. The obtained result revealed that the shape of obtained AgNPs was spherical (Fig. 5).

**FE-SEM examination**

FE-SEM images of BC and BC/AgNP composite shows three-dimensional structures of BC nanofibers. After the BC sheet was soaked in AgNO₃ for 2h and then irradiated, silver ions reduced by OH surface groups of BC to AgNPs (white dots) which appeared to adhere to the surface of BC bers (Fig. 6).

**FTIR spectrum** was performed to detect the interaction between BC and Ag-NPs. Figures (7) and table (2) show the FTIR spectra of BC and BC/Ag nanocomposites. For BC (a) characteristic bands of cellulose that appeared at the 3200–3400 cm⁻¹ region are assigned to the OH stretching vibration of the hydroxyl groups present in the BC nanofiber (Zhu et al., 2014; Cacicedo et al., 2020). Peaks at 3000 – 2800 cm⁻¹ were assigned to the stretching vibrations of the CH₂ and CH₂-OH groups (Li et al., 2011; Cacicedo et al.,
The band at 1640 cm$^{-1}$ can be assigned to C = O (Wang et al., 2017). The band at 1426.38 cm$^{-1}$ is assigned to the stretching vibrations of CH$_2$ or OH in-plane bending (Barud et al., 2011). The peak at 1321.94 to 1369.67 can be assigned to O-H in-plane bending (Osorio et al., 2019). The band at 1022.36 cm$^{-1}$ for the C-O-C and C-O-H stretching vibrations of the sugar ring (Huang et al., 2015). The absorption at 1158.31 cm$^{-1}$ is coming from the C-O-C stretching vibration of the pure cellulose. A group of absorption peaks at the wavenumber region of 1200 – 900 cm$^{-1}$ arise due to the C-O and C-C stretching vibrations of the cellulose network (Cui et al., 2014). The FTIR spectrum of BC/AgNP composite (b) contained all the characteristic peaks of BC along with an additional new band for AgNPs at 1545.5 cm$^{-1}$ in the BC/AgNP composite resulting from the hydrogen bonding interaction between BC and AgNPs (Wang et al., 2017; Wan et al., 2020).

**XRD analysis**

XRD patterns provide information about the crystalline structure of BC and BC/AgNP composite. Figure (8) showed three diffraction peaks at about 14.32°, 16.82°, 22.57°, and 14.66°, 16.56°, 22.86° for BC and BC/AgNP composite respectively, the peaks corresponded to (110), (110), and (200) crystal planes of cellulose (French, 2014 and Volova et al., 2018). In many studies, the XRD of pure BC membrane showed three characteristic peaks at 14.60°, 16.82°, and 22.78° (Yan et al., 2008; Ul-Islam et al., 2013; Wu et al., 2014). The XRD graph of BC/AgNP composite exhibited newly three peaks 27.4°, 32.4°, and 46.5° attributed to the diffractions from the planes of Ag. Mageswari et al., (2015) reported diffraction peaks of 46°, 54°, and 68°, that attributed to 2 1 1, 2 2 0, and 2 2 2 lattice planes of the face-centered cubic crystal structure of silver, while diffraction peak at 26° and 32° was indexed to 1 1 0 and 1 1 1, planes of silver oxide. In many studies, the face-centered cubic crystal structure of crystalline Ag showed diffraction peaks at around 38 °, 46°, and 64° that was attributed to the planes of 1 1 1, 2 0 0 and 2 2 0, respectively (Prakash et al., 2013; Jyoti et al., 2016 and Anjum and Abbasi, 2016). The peak at about 27° and 32° was also reported for the diffractions from the planes of silver (Kumar et al., 2012 and Rose et al., 2019).

The apparent crystal size (nm) of BC and BC/AgNP composite are reported in Table (3). According to the Scherrer equation, the peak that was used for calculating the crystalline size of BC and BC/AgNP was 22.57° and 22.86° respectively. The average size of the BC membrane is determined to be 5.416 nm and 5.4091nm for BC/AgNP composite.

Table (4) show the antibacterial activity of BC/AgNP composite, purified BC, AMC, CAZ, and S against Gram-positive (*Staphylococcus aureus, Enterococcus faecalis, and Listeria monocytogenes*) and Gram-negative (*proteus mirabilis and E. coli*) bacteria. Purified BC and amoxicillin/clavulanic acid did not show antibacterial activity against any tested bacteria. *Listeria monocytogenes, Proteus mirabilis, and E. coli* were resistant to ceftazidime. BC/AgNP composite and streptomycin showed antibacterial potency against both Gram-positive and Gram-negative bacteria. It was observed that *Enterococcus faecalis, Staphylococcus aureus, and Listeria monocytogenes* were more sensitive and gave higher inhibition zone for BC/AgNP composite while Gram-negative *E. coli* and *Proteus mirabilis* were more resistant. Many studies reported that AgNPs have shown higher antimicrobial potency in Gram-positive than Gram-
negative bacteria (Mandal et al., 2016; El-Sherbiny et al., 2020, Jiji et al., 2020). The antibacterial activity of AgNPs depends on many factors including size, charge, the concentration of AgNPs (Van Phu et al., 2014, Jiji et al., 2020), and also the stabilizer used (Phu et al., 2014). Various mechanisms have been proposed for the antibacterial action of AgNPs, the most common one can be that free silver ions uptake may preventing DNA replication or may cause direct damage to the bacterial cell wall by forming pits in the cell wall that lead to an increase in permeability of cell wall and final cell death (Jones and Hoek, 2010; Bapat et al., 2018).

4. Conclusion

In this work, a facile green method for the preparation of colloidal AgNPs and BC/AgNP composite under the induction effect of gamma radiation. BC soaked in AgNO₃ was used as a reducing and capping agent. There is no need for using a chemical reducing agent or a supplementary catalyst in this way. The prepared composite exhibit antibacterial activity against Gram-positive and Gram-negative bacteria. Therefore, the BC/AgNP composite has the great potential to be applied in the future development of biomedical instruments, food packaging, and therapeutics such as wound dressing.

5. Declarations

Conflict of interest

The authors declare that they have no conflict of interest.

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Human Participants and/or Animals

N/A

Informed consent

N/A

6. References

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7. Tables

Table (1). Particle size (nm) of AgNPs synthesized under gamma-ray

| Nanoparticles         | Particle size (nm) of AgNPs |
|-----------------------|-----------------------------|
| 25% of distribution   | 28.0 nm                     |
| 25% of the distribution| 28.0-49.5 nm                |
| 25% of the distribution| 49.5 -73.0 nm               |
| 5% of the distribution | 73.0 – 80.9 nm              |
| 10% of the distribution| 80.9 -106.8 nm              |
| 10% of the distribution| 106.8 – 210.9 nm            |

Table (2): FTIR spectrum of BC and BC/AgNP composite
| Samples                | β      | I/I1 | 2Theta (deg) | θ (rad) | ACS(nm) |
|-----------------------|--------|------|--------------|---------|---------|
| BC                    | 1.7217 | 100  | 22.57        | 0.1963  | 5.416   |
| BC/AgNPs composite    | 1.4253 | 100  | 22.86        | 0.1993  | 5.409   |

Table (3). Physical Parameters obtained from the XRD

Table (4): Antibacterial effect of BC/ AgNP composite, pure BC, AMC, CAZ, and S against tested bacteria
| Bacterial strain          | Zone inhibition (mm) | BC/Ag composite | BC | AMC | CAZ | S    |
|--------------------------|----------------------|-----------------|----|-----|-----|------|
| *Enterococcus faecalis*  | 9 ±0.57              | 0.0             | 0.0| 7±0.57| 14±1.5|
| *Staphylococcus aureus*  | 9±1                  | 0.0             | 0.0| 6±0.57| 13±0.28|
| *Listeria monocytogenes* | 10±1.15              | 0.0             | 0.0| 0    | 20±1.7|
| *Proteus mirabilis*      | 8±0.5                | 0.0             | 0.0| 0    | 28±1  |
| *Escherichia coli*       | 7±0.28               | 0.0             | 0.0| 0    | 19±0.57|

BC= Bacterial Cellulose, AMC= amoxicillin/clavulanic acid, CAZ= ceftazidime,

S = streptomycin

**Figures**

![Figure 1](image1)

**Figure 1**

BC produced by Komagataeibacter rhaeticus N1 MW322708 strain on modified HS medium at 30oC for 10 days using static culture conditions (a), harvesting of BC (b) and BC after purification and drying (c)
Figure 2

UV–visible spectrum of silver nanoparticles synthesized by BC under different doses of gamma-ray.

| Mean Diameter       | = 49.5 nm          |
|---------------------|--------------------|
| Stnd. Deviation     | = 34.4 nm (69.5%)  |
| Norm. Stnd. Dev.    | = 0.695            |

Variance (P.I.) = 0.483
Chi Squared = 27.177
Baseline Adj. = 0.000 %
Z-Avg. Diff. Coeff. = 9.39E-008 cm²/s

Figure 3
DLS analysis of colloidal AgNPs synthesized under gamma-ray

Figure 4

zeta potential of colloidal AgNP synthesized under gamma irradiation
Figure 5

TEM micrographs and particles size of synthesized AgNP
Figure 6

FESEM images (A, C) purified BC showing 3D network of fibers, (B, D) BC/AgNP composite that shows impregnated AgNP (white dots) into BC nanofibers
Figure 7

FTIR diagram (a) BC, (b) BC/AgNP composite
Figure 8

X-ray diffraction (XRD) patterns of purified BC (a) and BC/AgNP composite (b)