Antibacterial activity of molybdenum disulfide and green silver nanoparticles reduced by tea tree essential oil compounds

Atividade antibacteriana de dissulfeto de molibdênio e nanopartículas de prata verde reduzida por compostos de óleo essencial de melaleuca

Actividad antibacteriana del disulfuro de molibdeno y nanopartículas de plata verde reducidas por compuestos de aceite esencial de melaleuca

Abstract
The increasing resistance of microorganisms against antibiotics puts at risk the human being. This event requires urgently developing strategies for producing alternative antibacterial agents. The combination of green silver nanoparticles (reduced by essential oil) and molybdenum disulfide can be synergistically explored given the interaction of components. Herein, it is reported the response of isolated and combined components with resulting outstanding kinetics of kill-time (complete elimination of Staphylococcus aureus and Escherichia coli after four hours of contact), effective action against biofilm formation (~99% of inhibition in the biofilm formation). These results confirm that the intercalation of silver nanoparticles between exfoliated sheets of MoS$_2$ represents a promising strategy to develop efficient antibacterial agents against Gram-positive and Gram-negative bacteria.

Keywords: Tea tree; Silver nanoparticles; Molybdenum disulfide; Antibacterial.

Resumo
A crescente resistência dos microrganismos aos antibióticos coloca em risco o ser humano. Este evento requer o desenvolvimento urgente de estratégias para a produção de agentes antibacterianos alternativos. A combinação de nanopartículas de prata verde (reduzidas por óleo essencial) e dissulfeto de molibdênio pode ser explorada sinergicamente dada a interação dos componentes. Aqui, é relatada a resposta de componentes isolados e combinados com excelente cinética de tempo de morte (eliminação completa de Staphylococcus aureus e Escherichia coli após quatro horas de contato), ação eficaz contra a formação de biofilme (~99% de inibição no biofilme formação). Estes resultados confirman que a intercalação de nanopartículas de prata entre folhas esfoliadas de MoS$_2$ representa uma estratégia promissora para desenvolver agentes antibacterianos eficientes contra bactérias Gram-positivas e Gram-negativas.

Palavras-chave: Melaleuca; Nanopartículas de prata; Dissulfeto de molibdênio; Antibacteriano.

Resumen
La creciente resistencia de los microorganismos frente a los antibióticos pone en riesgo al ser humano. Este evento requiere el desarrollo urgente de estrategias para la produccion de agentes antibacterianos alternativos. La combinación de nanopartículas de plata verde (reducidas por aceite esencial) y disulfuro de molibdeno se puede explorar de forma sinérgica dada la interacción de los componentes. En este documento, se informa la respuesta de los componentes aislados y combinados con la resultante cinética sobresaliente de tiempo de muerte (eliminación completa de
Staphylococcus aureus y Escherichia coli después de cuatro horas de contacto), acción efectiva contra la formación de biopelículas (~99% de inhibición en la biopelícula formación). Estos resultados confirman que la intercalación de nanopartículas de plata entre láminas exfoliadas de MoS\textsubscript{2} representa una estrategia prometedora para desarrollar agentes antibacterianos eficientes contra bacterias Gram positivas y Gram negativas.

**Palabras clave:** Melaleuca; Nanopartículas de plata; Disulfuro de molibdeno; Antibacteriano.

1. **Introducción**

El prevalencia de microorganismos en la presencia de medicamentos resultan de resistencia antimicrobiana (Dadgostar, 2019) en un proceso favorecido por el uso no controlado de antibióticos en la producción humana y animal (Levy, 2005; Levy, 1998). Estas microorganismos resistentes a antibióticos conducen a las enfermedades y los procesos de prolongada hospitalización, considerado un brote debido a las limitaciones de tratamiento posibles (Levy, 2005). Este escenario es crítico dados las escasas nuevas antibióticos (LEVY, 2005). Consecuentemente, se ha observado un intenso esfuerzo por desarrollar alternativas para contrarrestar la proliferación de bacterias superresistentes.

En particular, las nanopartículas de plata (Ag NPs) han sido aplicadas exitosamente como agentes antibacterianos en heridas dermás (Nam et al., 2015) y contra un amplio espectro de especies bacterianas. La gran relación área-volumen de las nanopartículas de metal mejora la actividad antibacteriana del material resultante (Yılmaz et al., 2023). El mecanismo general involucra un proceso multifactorial que se basa en la adherencia de AgNPs a la membrana celular y la fuerte interacción con sus componentes sulfuro y fosfato (More et al., 2023), dañando su integridad por la fuga de componentes intracelulares (Bruna et al., 2021; Shaheen, 2016). Además, la formación de especies de oxígeno reactive afecta la metabolismo y la duplicación del DNA (Shaheen, 2016).

Un aspecto importante a considerar en la síntesis de nanopartículas de plata es el impacto ambiental debido a la utilización de agentes químicos tóxicos y dañinos (Vinicius de Oliveira Brisola Maciel et al., 2020). Para evitar estas desventajas, el proceso de síntesis verde de nanopartículas de plata ha considerado un proceso amigable con el medio ambiente que incluye la utilización de materiales (como extractos de plantas y aceites esenciales) que contienen terpenos y flavonoides relevantes para la reducción de Ag\textsuperscript{+} a Ag\textsuperscript{0} (M.L. Guimarães et al., 2020; Milena Lima Guimarães et al., 2019, 2022a; Vinicius de Oliveira Brisola Maciel et al., 2020). Los aceites esenciales han sido reconocidos por sus actividades adicionales (antibacteriana, antiviral, y antifúngica) que se pueden incorporar por reducción en las nanopartículas (Milena Lima Guimarães & Amarante, 2021; Nehme et al., 2021).

En el otro lado, las nanoestructuras de MoS\textsubscript{2} han sido exitosamente aplicadas en la fototerapia terapéutica del cáncer, pero también para aplicaciones antibacterianas (Wentao Zhang et al., 2016). Con este fin, el uso de MoS\textsubscript{2} en disoluciones de multi-resistencia a antibióticos (Zhao et al., 2021) mientras la combinación de MoS\textsubscript{2} y vancomicina ha sido aplicada en el tratamiento fototerapéutico inhibiendo el crecimiento de *Staphylococcus aureus* (Weiwei Zhang et al., 2022).

Aquí, las nanopartículas de plata verdes reducidas por el aceite esencial de árbol de té (aceite esencial de melaleuca) se combinan con hojas exfoliadas de MoS\textsubscript{2} en compositos que se utilizan como agentes antibacterianos contra *S. aureus* y *E. coli* explorando la acción combinada de aceite esencial como una agente/agentes antibacteriano, nanopartículas de plata y hojas exfoliadas de molybdenum disulfide en un proceso synergístico que resulta en una actividad eficiente de inhibición bacteriana a bajas concentraciones de componentes. La respuesta de composados de Ag/MoS\textsubscript{2} (aislados y combinados) se evaluó en la difusión de agar, inhibición de biopelículas, tiempo de muerte, vida útil de biofilm, inhibición y toxicidad. El esquema general del uso de este artículo se dibuja en la Figura 1, que representa las hojas exfoliadas de MoS\textsubscript{2}, las nanopartículas de plata verdes (Ag NPs), y el compuesto (MoS\textsubscript{2}+Ag NPs) – (con la correspondiente imagen de MoS\textsubscript{2} y agrupaciones de AgNPs) aplicada para adherirse a la superficie bacteriana provocando la muerte de células por fuga de fluidos vitales.
Figure 1 - General scheme for exfoliated nanosheets of MoS\(_2\), silver nanoparticles, and MoS\(_2\)+Ag NPs nanostructures with the corresponding SEM image and the general process of nanoparticles adhesion and bacterial cell inactivation with DNA leakage.

Source: Authors

2. Methodology

Molybdenum disulfide (Radmax, Brazil), sodium hydroxide (Dinâmica, Brazil), ethyl alcohol (Dinâmica, Brazil), and silver nitrate (Aldrich, USA). Tryptic soybean broth (TSB) (Kasvi, Spain), phosphate buffered saline solution (PBS) (Kasvi, Spain), plate counter agar (PCA) AlphaTec (Brazil), and Brain Heart Infusion (BHI) (Kasvi, Spain) were used as received and all solutions were prepared using Milli-Q water. Tea tree essential oil (Melaleuca alternifolia) was purchased from Akã Oils Essential (Brazil).

Scanning Electron Microscope Vega 3XM, Tescan evaluated the morphology of materials. The absorbance in the UV-vis region was performed in a UV-vis Hach DR5000 using a single quartz cuvette in the range of 300 nm to 800 nm at a step of 1 nm. Zetasizer Nano ZS-90 Malvern provided data about the size of particles and zeta potential. FTIR Prestige-21 Shimadzu evaluated the structure of materials while the composition of MoS\(_2\) was provided by an X-ray fluorescence spectrometer CTX model 800.

2.1 Exfoliation of Molybdenum disulfide

For exfoliation of molybdenum disulfide (Zhou et al., 2011), 120 mg of powder was dispersed in 30 mL of a solution containing 45% of ethanol and 55% of water with sodium hydroxide (10 mg) (Ikram et al., 2020) with the resulting solution sonicated for 8 hours. Subsequently, the samples were centrifuged at 15,000 rpm for 10 minutes. The resulting solution was dried and washed until it reached pH 7 providing the dispersion of MoS\(_2\) nanosheets to be explored in the following synthesis of the MoS\(_2\)+AgNPs composite.
2.2 Synthesis of silver nanoparticles with exfoliated molybdenum disulfide

Silver nanoparticles were biosynthesized according to the method described by Maciel et al. (Maciel et al., 2019) with some modifications. As a general standard method, an aliquot of tea tree oil (100 μL) was diluted in 17 mL of acetone at pH 8, which was adjusted by incorporating an aqueous NaOH solution (0.1 M). A second aqueous solution of silver nitrate (1 mM) was also prepared (system Ag NPs). For the preparation of MoS<sub>2</sub>+Ag NPs, 30 mL of AgNO<sub>3</sub> received exfoliated MoS<sub>2</sub> nanosheets (from the reaction described in section 2.1), and it was kept under intense agitation and heating (95 °C). After this step, the tea tree oil solution was added dropwise into an aqueous silver nitrate solution with the MoS<sub>2</sub> nanosheets until it reached a total volume of 32 mL (2 mL of essential oil solution). The kinetics of silver nanoparticle formation associated with exfoliated molybdenum disulfide were monitored at fixed time intervals (15, 30, 45, and 60 min) with the measurement of the characteristic peak of absorbance of silver nanoparticles in the UV-Vis spectrum. All of the procedures were conducted in the dark to avoid the effect of light-induced degradation.

2.3 Broth microdilution assays

Broth microdilution assays were performed following the Institute for Clinical and Laboratory Standards (CLSI, 2019) for the determination of the minimum bactericidal concentration (MBC) of silver nanoparticle solutions prepared by green synthesis (Ag NPs system), MoS<sub>2</sub> system, and MoS<sub>2</sub>+AgNPs system. S. aureus (ATCC 25923) and E. coli (ATCC 25922) were kept in BHI at 4 ºC until the use. Initially, 100 μL of TSB was added to each microplate well. Then, 100 μL of the solution under test (MoS<sub>2</sub>, AgNPs, and MoS<sub>2</sub>+AgNPs) were added to the first well and, after homogenization, it was transferred to the second well, and so on, obtaining concentrations of 1:1, 1:2, 1:4, 1:8; 1:16; 1:32; 1:64; 1:128 compared to the initial concentration of the antibacterial agent and a bacterial suspension standardized to 0.5 McFarland scale was prepared. Aliquots of 10 μL were transferred to each well of the microplate containing the dilutions of the nanoparticle solutions, and incubated at 37 °C for 24 h. With the aid of a microplate replicator, an aliquot of the solution from the microplate wells was seeded on the surface of the PCA medium taken for 24 h at 37 ºC.

2.4 Agar diffusion assays

The agar diffusion assays were performed according to Guimarães et al., Gram-positive S. aureus (ATCC 25923) and gram-negative E. coli (ATCC 25922) bacterial inoculums were prepared from cultures maintained in agar at 4 ºC. Then, aliquots of bacterial viable cells dispersed in culture media were removed and inserted into saline solution (0.85%), reaching turbidity of 0.5 on the McFarland scale (10<sup>8</sup> CFU mL<sup>-1</sup>). The bacterial suspension was distributed in a Petri dish containing PCA medium with a swab. Discs formed by liquid samples of MoS<sub>2</sub>, AgNPs, and the MoS<sub>2</sub>+AgNPs association were incorporated into the plates to assess their activity. After this step, the plate was incubated at 37 ºC for 24 h (Milena Lima Guimarães et al., 2022b).

2.5 Kill time assays

The shortest time for the samples to eliminate the bacteria from the medium was evaluated from kill time assays, using the bacterial inoculum S. aureus (ATCC 25923) and E. coli (ATCC 25922), which were prepared from cultures maintained in agar at 4 ºC. Assays were performed as follows: 5 mL of 10<sup>8</sup> CFU mL<sup>-1</sup> bacterial solution were placed in a test tube, which was further diluted to 10<sup>6</sup> CFU mL<sup>-1</sup>. After homogenizing the bacterial solution, the three samples (MoS<sub>2</sub>, AgNPs, and MoS<sub>2</sub>+AgNPs) were disposed of in three tubes. Positive control was performed to compare the results. Then, 100 μL aliquots were removed from the tubes at time intervals (1 h, 2 h, 3 h, and 4 h) and disposed of in Petri dishes containing PCA. The plates were kept at 37 ºC for 24 hours (Milena Lima Guimarães et al., 2022b).
2.6 Antibiofilm assays

Using a bacteriological loop, inoculums of *S. aureus* (ATCC 25923) and *E. coli* (ATCC 25922) were dispersed in 10 mL of TSB in different tubes. After preparing the bacterial solutions, the different compounds (MoS$_2$, AgNPs, and MoS$_2$+AgNPs) were added to three different tubes with the positive control performed for comparison with the results. Then, the tubes were incubated at 37 °C for 24 hours. After incubation, the contents of the tubes were discarded, the tubes were washed with milli-Q water and 12 mL of saline solution was added to each tube. Then, the tubes were sonicated in a bath (f = 40 kHz) for 15 min to remove the trapped species. Aliquots of 100 μL were taken in triplicate from each system and seeded in a PCA medium. After incubation at 37°C for 24 h, colonies were counted to determine the number of remaining viable cells in the biofilms.

2.7 Cytotoxicity assays

Dehydrated cysts of Artemia salina were purchased from a local aquarium store (Artemio Pur JBL GmbH&Co., Germany). Then, the cysts were hydrated in artificial seawater (ASA), dissolving 52.5 g of NaCl in 1.5 L of Milli-Q water under stirring (maintaining a concentration of 35 g/L) under artificial lighting (1500 lux supplied continuously by a fluorescent lamp). The nauplii hatched in 48 hours and after that period, the tests with the samples took place. Stock solutions of MoS$_2$, AgNPs, and the MoS$_2$+AgNPs systems were prepared and diluted in ASA, using the MBC value for each sample. 30 nauplii were placed in contact with each material under test in triplicate with the positive control applied for comparison (nauplii in artificial seawater). Suspensions diluted in ASA were vortexed and subsequently sonicated for 15 minutes to completely disperse the material under study.

The solutions were added to the 12-well plate, in triplicate. After that, 10 Artemia Salina nauplii were added to each well and incubated at 26 °C +/- 2 °C for 24 h. The number of surviving nauplii in each well was counted after 24 and 48 hours with larvae not fed in the bioassays. A nauplius is considered dead if there is no movement of its antennae after a slight agitation of the water for 10 seconds. Mortality percentages were calculated by comparing the number of survivors in the test and the control experiments.

3. Results and Discussion

Scanning electron microscopy images were performed to evaluate the morphology of MoS$_2$, AgNPs, and the MoS$_2$+AgNPs systems. As shown in Figure 2a, MoS$_2$ exhibits nanosheets-like morphology. Figure 2b shows silver nanoparticle aggregates for sample AgNPs, while sample MoS$_2$+AgNPs shown in Figure 2c is characterized by a high density of sheets with intercalated and exfoliated structures with aggregates of silver nanoparticles on the molybdenum disulfide sheets. The components identification by the dispersive energy assays, by characteristic peaks of Mo and Ag (see Figure 2d) confirms the successful nucleation of silver nanoparticles in MoS$_2$ dispersion (Anderson et al., 2017).
The evaluation of the composition of the molybdenum disulfide powder sample was scrutinized by using X-ray fluorescence spectroscopy. The result of the analysis indicates a content of Mo = 53.56%, S = 42.39%., SiO$_2$ = 5.92% and Fe=1.49%. The characteristic groups of the samples MoS$_2$, Ag NPS, and MoS$_2$+AgNPs were studied by the FTIR spectrum, as summarized in Figure 3. As can be seen, the FTIR spectrum for molybdenum disulfide presents a strong absorption peak at 3435 cm$^{-1}$ due to the O–H stretching vibration (Awasthi et al., 2016) and a peak at 1132 cm$^{-1}$ attributed to the formation of sulfur complexes with the active sites of MoS$_2$ (Feng et al., 2014). The stretching vibrations of the C=O bond can be observed at 1228 cm$^{-1}$, which are groups resulting from the exfoliating solution (Bodík et al., 2019). The peak at 866 cm$^{-1}$ corresponds to the Mo–O elongation vibrations (Li et al., 2014), indicating that the exfoliated MoS$_2$ has partially oxidized edges (Ikram et al., 2020) while the band at 1019 cm$^{-1}$ is attributed to Mo–S vibrations (Ikram et al., 2020). The 1228 cm$^{-1}$ and 1130 cm$^{-1}$ bands correspond to SO$_2$ groups from interactions between sulfur from molybdenum disulfide and adsorbed oxygen (Bodík et al., 2019). The band around 878 cm$^{-1}$ represents the stretching vibration of the S – S bonds (Mohan et al., 2019).

The FTIR spectrum for the AgNPs shows a broad band at 3435 cm$^{-1}$ that is attributed to the O–H and N–H stretching vibrations. The band at 1383 cm$^{-1}$ arises from the C–O–H bending that can be attributed to polysaccharides commonly in plants (Bharadwaj et al., 2021; Huang et al., 2020). At 866 cm$^{-1}$, a band is attributed to the double bond between C=C carbon atoms, characteristic of essential oils. For the sample MoS$_2$+AgNPs, combined peaks are observed, confirming the presence of both components in the material.
Figure 3 - FTIR spectrum for MoS$_2$ (curve in black), AgNPs (curve in red), and the MoS$_2$+AgNPs (curve in blue) samples.

The UV-Vis spectrum of the MoS$_2$ sample is shown in Figure 4a being characterized by bands centered at 604 nm and 662 nm, respectively (Winchester et al., 2014). These responses are characteristic of the 2H-MoS$_2$ phase (Eda et al., 2011). Also, a broad absorption peak at 400 nm confirms the theoretical and experimental values reported in the literature for MoS$_2$ (Mak et al., 2010).
The optical response for nanoparticles synthesized from tea tree essential oil (Ag NPS system) is shown in Figure 4b. From this result, it is possible to observe an efficient activity of essential oil as reducing agents for silver ions from the observed plasmon band at 420 nm. The reduction of silver nanoparticles can be explained by the presence of monoterpenoids and biomolecules in an essential oil solution, described as potential silver ion reducers (Melo et al., 2020) in addition to α-pinene and γ-terpinene (Wei et al., 2020). The intensity at this peak position increases with reaction time, indicating progressive nucleation and formation of silver nanoparticles (Milena Lima Guimarães et al., 2021).

Figure 4c shows the UV-Vis absorption spectrum for the MoS$_2$+AgNPs system. As can be seen, there is a typical shift in the specific bands relative to MoS$_2$ and AgNPs samples, suggesting an additional step of aggregation level for these materials.

The size of particles is summarized in Figures 5a, 5b and 5c for MoS$_2$, Ag NPs and MoS$_2$+AgNPs compounds, respectively. As seen in Figure 5a, a single-size distribution is observed for exfoliated molybdenum disulfide, centered at 342 nm. For silver nanoparticles (see Figure 5b) two populations of size of particles are observed, at 15.69 nm and 122.4 nm. The presence of particles in the nanoscale is an important property to be considered in the applications as biomaterials for antibacterial application, improving their ability to attach to the bacterial cell wall (Agnihotri & Dhiman, 2017). The particle sizes for MoS$_2$ AgNPs compounds are characterized by particles at 24.36 nm and 164.2 nm, indicating that the exfoliation degree of MoS$_2$ is also affected by the intercalation of the AgNPs.
Figure 5 - Size distribution in (a) MoS$_2$, (b) Ag NPs, and (c) MoS$_2$+AgNPs systems.

The polydispersity index (PDI) is a dimensionless measurement of the particle size distribution, which ranges from 0 to 1. The smaller the PDI, the more homogeneous the sample. The sample MoS$_2$ presents a PDI of 0.255. The silver nanoparticles presented a PDI of 0.449, a less homogeneous distribution of particles. For sample MoS$_2$ + AgNPs, the PDI was 0.285. In addition, the system based on MoS$_2$ presented a zeta potential of -8.24 mV, the AgNPs presented a zeta potential of -2.48 mV and sample MoS$_2$+AgNPs presented a zeta potential of -7.80 mV, characterizing as experimental systems with low stability degree.

The broth microdilution test was applied in the MBC determination of different experimental systems against *S. aureus* (10$^6$ CFU mL$^{-1}$) and *E. coli* (10$^6$ CFU mL$^{-1}$). In a microplate, serial dilutions were made in the TSB broth of the c. Aliquots of 10 μL of bacterial suspension were added to each well, except for wells applied as negative control (TSB solution). The results against *S. aureus* for MoS$_2$ samples returned efficiency at a dilution of 1:1. On the other hand, it was observed that silver nanoparticles obtained inhibition at a dilution of 1:4 and the sample MoS$_2$+AgNPs inhibited until a dilution of 1:16. For the system MoS$_2$, the complete elimination of bacteria was observed at a dilution of 1:2 while silver nanoparticles obtained inhibition at a dilution of 1:8 and for the system, MoS$_2$+AgNPs the value was reduced to 1:32.

As can be observed, with all of the dilution relative to the originally synthesized systems, the formulation MoS$_2$, AgNPs, and MoS$_2$+AgNPs were successfully applied against Gram-positive and Gram-negative bacteria with efficiency following the order MoS$_2$ + AgNPs > Ag NPs > MoS$_2$ with dilutions in the order of 1/32 from the initial concentration preserving antibacterial activity, an important advantage for this experimental system.

The agar diffusion tests were carried out for samples MoS$_2$, AgNPs, and MoS$_2$+AgNPs. The culture medium was placed in the Petri dish and then perforated to accommodate the solution under study with subsequent observation and identification of the inhibition halos of each material. The solutions were tested against *S. aureus* and *E. coli*. Figures 6a and 6b show the result of the inhibition halos against *S. aureus* while results in Figures 6c and 6d are observed for *E. coli*, in which it is possible to
observe increasing haloes for MoS$_2$, AgNPs, and MoS$_2$+AgNPs, in the following order: MoS$_2$ ~ AgNPs < MoS$_2$+AgNPs. The antibacterial activity observed for exfoliated molybdenum disulfide is attributed to the interaction of the edges of nanosheets with the bacterial membrane, resulting in cell damage and death.

**Figure 6** - Inhibition halo assay (a) and measured value for halo (b) for samples MoS$_2$, AgNPs, and MoS$_2$+AgNPs against *S. aureus* and inhibition halo assay (c) and measured value for halo (d) for samples MoS$_2$, AgNPs, and MoS$_2$+AgNPs against *E. coli*.

The antibiofilm activity of the compounds was evaluated from the direct count of viable cells attached to the reactor in contact. Figure 7 shows the result of antibiofilm activity against *S. aureus* (a typical biofilm-forming system) for the three different compositions. In biofilms, bacterial cells are encapsulated by secreted polymeric extracellular proteins that strongly protect bacterial cells from the effects of antibiotics. These results confirm the strong antibiofilm activity of materials in comparison with the control sample. For these compositions, an antibiofilm activity AB ~ 95% is observed for molybdenum disulfide nanosheets, AB ~ 97% for silver nanoparticles, and AB ~ 99% for the MoS$_2$+AgNPs system.
Figure 7 - Relative biofilm inhibition for exfoliated molybdenum disulfide, silver nanoparticles, and for the association between the two materials against *S. aureus*.

These results confirm the ability of exfoliated molybdenum disulfide, silver nanoparticles, and the association of the two materials to cross the extracellular polymeric matrix accessing the *S. aureus* viable cells embedded in the biofilm, causing membrane damage and death of cells (Roy et al., 2019). Thus, the ability of these materials to eradicate the biofilm of *S. aureus* opens the possibility to further explore these nanomaterials as a disinfection agent, with an improvement in the overall response for systems with the interaction of AgNPs and MoS$_2$.

The assays of the kinetics of death of bacteria (*S. aureus* and *E. coli*) provided by the three experimental systems (MoS$_2$, AgNPs, and MoS$_2$+AgNPs) were determined from the direct measurement of viable cells after fixed intervals of time of interaction of the bacteria with these three antibacterial agents. As can be seen in Figure 8, from a comparison of three experimental systems (against *S. aureus* (a) and *E. coli* (b)), the observed trend is a progressive reduction of viable bacteria with increasing treatment time, while the negative control (as expected) returned negligible variation in the number of viable cells for both bacteria.
As observed, a strong reduction in viable bacterial cells is observed in the first hour of the interaction. Throughout the experiments, a gradual and noticeable reducing number of viable bacterial cells is observed between 2h and 4h of the experiment. By direct comparison of the results, it is possible to verify that there is a synergism in the association between exfoliated molybdenum disulfide and silver nanoparticles, a fact that becomes from the evaluation of slope in the MoS$_2$+AgNPs system against the two bacterial strains. Intermediate performance is observed for silver nanoparticles and exfoliated molybdenum disulfide. A low percentage of immobilization (~3.33%) was obtained for Artemia Salina nauplii treated with silver nanoparticles, 0% for molybdenum disulfide, and 0% for MoS$_2$+AgNPs system.

Accumulation of NPs on artemia was observed for all materials from a stereomicroscope equipped with a camera. Compared to the negative controls, it is possible to conclude that Artemia consumed the particles. The ingested particles appeared as a long dark streak within the bowels. This can be explained by the lack of food intake and absorption and the filling of the intestine with aggregates of nanoparticles. These results suggest that the silver nanoparticles synthesized by the green synthesis, the exfoliated molybdenum disulfide, and the association between these materials were not toxic to Artemia Salina.

### 4. Conclusion

Despite good performance against *S. aureus* and *E. coli* observed for individual components (AgNPs and MoS$_2$ samples), the combination of silver nanoparticles reduced by tea tree essential oil in the presence of molybdenum disulfide nanosheets demonstrated synergistic antibacterial performance with superior performance in terms of kinetics of inhibition, diffusive assays, and antibiofilm activity, in an indication that reduction of silver nanoparticles into exfoliated MoS$_2$ results in a material with intercalated nanosheets of MoS$_2$ decorated with silver nanoparticles that present economic advantages (reduction in the MBC), outstanding performance against biofilms and environmentally friendly component from the green synthesis of silver nanoparticles, being considered a promising new alternative for antibacterial system against Gram-positive and Gram-negative species. It is expected to develop novel prototypes based on adsorbing surfaces impregnated with MoS$_2$+AgNPs for application as smart and multifunctional cleaning fibers loaded with green-based compounds with mutual activity against planktonic and biofilm of single and multiple bacterial species.
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References


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