Modulating Effect of 4,6,6-trimethylbicyclo [3.1.1] hept-3-ene on Antimicrobials Used to Combat *Escherichia coli*

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**Graphical Abstract**

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**Abstract.** The growing number of resistant bacteria is considered a worldwide public health problem. In this context, several control initiatives have been proposed, such as the elaboration of medicines from vegetable raw material as an alternative to fight against microorganisms such as *Escherichia coli*, which is related to approximately 50% of hospital infections. α-Pinene is found in the oils of many coniferous tree species and has antimicrobial activity against some microorganisms. The positive enantiomer of this compound (+) - α - pinene (+ AP), also called 4,6,6-trimethylbicyclo [3.1.1] hept-3-ene, obtained from the company Sigma-Aldrich do Brasil Ltda. The solutions were dissolved in 1% Tween 80 and 5% DMSO and sterile distilled water was used to achieve desired concentrations. The tests were performed on the *Escherichia coli* strain ATTC 25922. To perform the modulation and adaptation tests, discs containing commercial antibiotics (ATM) were used. The modulating action of monoterpenes was determined by the disc diffusion method. The antimicrobials Ceftazidime, Amoxicillin, Cefepime and Cefoxitin demonstrated synergism through association with phytoconstituent. For the remaining ATM, there was no statistically significant difference, and the effect of the association was classified as indifferent. It is concluded that
Introduction

Currently, the class of antibiotics is at the top of the list of the most prescribed medications in the world, a fact that may be responsible for the development of bacterial resistance. Recent research shows that on average 50% of antimicrobials are not prescribed correctly by doctors, and the number of people who self-medicate without a professional indication is quite significant. Continuing these errors not only increases the selection of resistant strains, but also increases the costs of each patient and the effect of disease and death on a population (LIMA; BENJAMIM; SANTOS, 2017).

In theory, the practice of combining antimicrobial compounds would increase the chance of employing at least one effective agent to treat a given infection. In this case, if there were any resistance to one of the antimicrobials employed, but not to all, the microorganism would be destroyed by the agent (s) that maintained its activity in the scheme, with no resistance evolution (ZIMERMAN, 2010). In this context, the use of drugs of natural origin becomes an effective and economical alternative because they provide high chances of obtaining prototype molecules due to the diversity of their phytoconstituents with possibilities of increased antibacterial potential when associated with pathogen antibiotics (DE SOUZA ELLER et al., 2015).

In search of biological materials that interact in this sense, PEDROSA (2014) verified the association of three products, the phytoconstituent thymol, vancomycin and EDTA, which evaluated the sensitivity and the possibility of reducing the doses of the antibiotic cited against *E. coli* and may reduce the usual dose against this pathogen by up to sixteen times. In addition, other studies have shown that monoterpenes have detrimental effects on the bacterial cell membrane, showing themselves as alternatives to fighting pathogens such as *E. coli*, which is the subject of this study and is related to approximately 50% of hospital infections and 70-90% of episodes of urinary tract infections (KORB et al., 2013).

Among these monoterpenes is α-pinene, which has shown promise against bacterial biofilm formation and presented growth inhibition halos of *E. coli* strains. This fact demonstrates the possibility of using this phytoconstituent as an antimicrobial agent alone or in combination with other drugs (FARIAS et al., 2017).
Therefore, this study aims to evaluate the modulating effect of 4,6,6-trimethylbicyclo [3.1.1] hept-3-ene monoterpene, (+) - α – pinene, with antimicrobials used in clinical therapy to combat infections caused by *Escherichia coli* as a way to help fight resistant pathogens.

**Materials and Methods**

Phytoconstituent 4,6,6-trimethylbicyclo [3.1.1] hept-3-ene (+ AP), obtained from the company Sigma-Aldrich do Brasil Ltda. The solutions were dissolved in 1% Tween 80 and 5% DMSO and sterile distilled water was used to achieve desired concentrations. The tests were performed on the *Escherichia coli* strain ATTC 25922. The culture media used were Mueller-Hinton Agar (AMH), Mueller-Hinton Broth (CMH) and BHI Broth (brain and heart infusion broth). For the modulation and adaptation tests, discs containing the commercial antibiotics (ATM) were used, and those acting on the bacterial cell wall were as follows: Cephalothin, Ceftazidime, Amoxicillin, Ampicillin, Cefepime, Cefoxitin, Meropenem and Cefuroxime. Prior to use, the media were solubilized in distilled water and autoclaved at 121°C for 15 minutes.

After the incubation period, the inoculum was prepared by making a direct suspension in saline of selected isolated colonies. The modulating action of monoterpene was determined by the disc diffusion method. After incubation of the plates at 35 ± 2 °C for 24 hours, the diameter of the microbial growth inhibition halos was read. To investigate the effect of subinhibitory concentrations of 4,6,6-trimethylbicyclo [3.1.1] hept-3-ene on the sensitivity profile of the tested strain, the adapted experimental protocol of Ultee et al. (2000) and McMahon et al (2007).

**Results and Discussion**

The study results were verified by comparing growth inhibition halos before and after association with the phytoconstituent to demonstrate whether there was a statistically significant change or synergism in this use, as can be seen in Figure 1.
**Escherichia coli ATCC 25922**

![Diagram showing the diameter of halos for various antibiotics and phytoconstituents]

α-pineno – (+)-alfa-pineno CFL-Cefalotina, CAZ-Ceftazidime, AMX-Amoxicilina, AMI-Amicacina, AMP-Ampicilina, CPM-Cefepime, CFO-Cefoxitina, NIT-Nitrofurantoína, CLO-Cloranfenicol, SUT-Sulfazotrim, MER-Meropenem, GEN-Gentamicina, CRX-Cefuroxime, CIP-Ciprofloxacina. Data were submitted to analysis of variance (One-Way ANOVA) and expressed as mean ± m.s.e.

The following growth inhibitor halo diameters for antimicrobial isolated cell wall inhibitors and monoterpenic inhibitors were as follows: (+) - α-pinene: 13.33 mm (± 0.58 mm), Cephalothin: 14.00 mm (± 2.00 mm), Ceftazidime: 25.33 mm (± 1.16 mm), Amoxicillin: 23.33 mm (± 1.16 mm), Amikacin: 21.33 mm (± 1.16 mm), Ampicillin: 19.67 mm (± 0.58 mm) Cefepime: 31.33 mm (± 1.16 mm), Cefoxitin: 23.33 mm (± 1.16 mm), Nitrofurantoin: 28.67 mm (± 3.51 mm), Chloramphenicol: 27.33 mm (± 1.16 mm), Sulfazotrim: 25.33 mm (± 3.06 mm), Meropenem: 34.67 mm (± 3.06 mm), Gentamycin: 22.00 mm (± 4.00 mm), Cefuroxime: 20.67 mm (± 3.06 mm), Ciprofloxacin: 32, 67 mm (± 2.31 mm).

Applying the classification proposed by Cleeland and Squires (1991) for the effect of interaction between substances, we can classify the effect of the association of phytoconstituent to the antimicrobials Ceftazidime, Amoxicillin, Cefepime and Cefoxitin as a synergistic effect. However, for the other ATMs, there was no statistically significant difference, and the effect of the association was classified as indifferent. Thus, if there was antagonism or indifference in the results for some drugs, the possibilities open for further studies on the resistance and adaptation mechanism of the strain used and how the synergistic associations were able to increase the potential toxic effect against the pathogen.

**Conclusions**
Thus, it is concluded that phytoconstituent 4,6,6-trimethylbicyclo[3.1.1]hept-3-ene, having synergistic effects with some drugs tested in this work, can be used as an associated substance and potentiator against *E. coli* ATCC 25922, provided that it is submitted to tests to verify their allergenic potential against humans. Similarly, the association of this monoterpen with the antibiotics that had antagonistic or indifferent effects in this study should be avoided, not allowing the growth of resistant bacterial strains. Thus, opportunities are created for the reduction of morbidity and mortality associated with resistant microbes and for the reduction of toxicity related to certain commercially available drugs by reducing the doses used.

References

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