Toxicity of clove essential oil and its ester eugenyl acetate against Artemia salina

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Received: August 11, 2015 – Accepted: December 10, 2015 – Distributed: February 28, 2017
(With 3 figures)

Abstract
The production of compounds via enzymatic esterification has great scientific and technological interest due to the several inconveniences related to acid catalysis, mainly by these systems do not fit to the concept of “green chemistry”. Besides, natural products as clove oil present compounds with excellent biological potential. Bioactives compounds are often toxic at high doses. The evaluation of lethality in a less complex animal organism can be used to a monitoring simple and rapid, helping the identification of compounds with potential insecticide activity against larvae of insect vector of diseases. In this sense, the toxicity against Artemia salina of clove essential oil and its derivative eugenyl acetate obtained by enzymatic esterification using Novozym 435 as biocatalyst was evaluated. The conversion of eugenyl acetate synthesis was 95.6%. The results about the evaluation of toxicity against the microcrustacean Artemia salina demonstrated that both oil (LC₅₀ = 0.5993 µg.mL⁻¹) and ester (LC₅₀ = 0.1178 µg.mL⁻¹) presented high toxic potential, being the eugenyl acetate almost 5 times more toxic than clove essential oil. The results reported here shows the potential of employing clove oil and eugenyl acetate in insecticide formulations.

Keywords: Novozym 435, Caryophyllus aromaticus, enzymatic esterification.

Toxicidade do óleo essencial de cravo e seu éster acetato de eugenila contra Artemia salina

Resumo
A produção de compostos via esterificação enzimática possui grande interesse científico e tecnológico devido às inúmeras inconveniências relacionadas com a catálise ácida, principalmente por estes sitemas não se adequarem ao atual termo “tecnologias limpas”. Além disso, produtos naturais como o óleo de cravo, apresentam compostos com excelentes potenciais biológicos. Compostos bioativos são quase sempre tóxicos em altas doses. A avaliação da letalidade em um organismo animal menos complexo pode ser usada para um monitoramento simples e rápido, servindo também para a identificação de compostos com potencial atividade inseticida contra larvas de insetos vetores de doenças. Neste sentido, foi determinada a toxicidade frente a Artemia salina do óleo essencial de cravo e do seu derivado acetato de eugenila obtido por esterificação enzimática com lipase Novozym 435. A conversão da reação de síntese de acetato de eugenila foi de 95,6%. Os resultados referentes à avaliação da toxicidade frente ao microcrustáceo Artemia salina demonstraram que tanto o óleo (LC₅₀ = 0,5993 µg.mL⁻¹) quanto o éster (LC₅₀ = 0,1178 µg.mL⁻¹) apresentam elevado potencial toxicológico, sendo que o éster apresenta aproximadamente 5 vezes mais toxicidade em relação ao óleo. Estes resultados demonstram o potencial emprego do óleo de cravo e de acetato de eugenila em formulações de inseticidas.

Palavras-chave: Novozym 435, Caryophyllus aromaticus, esterificação enzimática.

1. Introduction
The species Caryophyllus aromaticus L. belongs to the family Myrtaceae, and is commonly known as clove. The essential oil obtained from the clove is constituted by a mixture of compounds, being the eugenol (4-alil-2-metoxyphenol) its major compound. The concentration of eugenol varies from 77-95% (Trajano et al., 2010;
The oil also presents other components of terpenes source as β-cariofilen, α-humulen and eugenyl acetate (Chaieb et al., 2007; Dzamic et al., 2009). Eugenol has different biological properties confirmed: bactericide (Moon et al., 2011), antifungal (Rana et al., 2011), larvicidal (Pandey et al., 2013), antioxidant (Chiaradia et al., 2012; Vanin et al., 2014), anti-inflammatory (Daniel et al., 2009), among others.

Thus, the eugenol esters can also present promising biological properties. Carrasco et al. (2008) observed that eugenyl acetate presented anti-carcinogen activity in cells of prostate and oral squamous cancers. The acaricide activity of eugenyl acetate was related by Pasay et al. (2010), since eugenol acetate presented high toxicity against human scabies mites. This compound was also described as a potent antioxidant agent by Vanin et al. (2014). The authors observed that the clove essential oil after esterification presented higher antioxidant activity compared to crude essential oil (IC\textsubscript{50} of 283.9 μg.mL\textsuperscript{-1} and 367.5 μg.mL\textsuperscript{-1}, respectively).

Although they are natural compounds, both eugenol and eugenyl acetate, they can present toxicity according to the form of use, prepare and concentration. So, the care with the use is of primordial importance in the control of possible collateral and adverse effects that the chronic and/or acute use can cause on the organism.

The use of bioassays for monitoring the bioactivity of extracts, fractions and plant isolated compounds have often been incorporated into the phytochemical research. Among these biological assays it is the toxicity test with Artemia salina (BST-Brine Shrimp Test) which was developed to detect bioactive compounds in plant extracts (Meyer et al., 1982; Noldin et al., 2003). This test is a simple method in the research of natural products, which has a good correlation with tests of acute oral toxicity in vivo (Parra et al., 2001).

Literature has been presented correlations between the general toxicity with the microcrustacean Artemia salina and the citotoxicity on strains of human cells of solid tumors (Mclaughlin, 1991; Mclaughlin et al., 1998) and activity anti-Trypanosoma cruzi (Zani et al., 1995). It has been shown that there is a very good correlation between the median lethal concentrations (LC\textsubscript{50}) of plant extracts to brine shrimp larvae and the median lethal doses (LD\textsubscript{50}) of the same extracts, administered orally in mice (Parra et al., 2001). Besides, the essential oils with high toxic potential have been target of investigation to the production of insecticides, related to control of larvae of vectors of diseases related to public health (Leite et al., 2009). Despite described properties of eugenol and eugenyl acetate, there is a lack in the literature regarding the eugenyl acetate toxicity in Artemia salina.

Based on these aspects, this work aimed the production of eugenyl acetate by enzymatic esterification of clove essential oil, and the investigation of toxic potential against the microcrustacean Artemia salina of clove essential oil (Caryophyllus aromaticus) before and after the enzymatic esterification, in order to provide toxicity information for future applications thereof.

2. Material and Methods

2.1. Enzymatic production of eugenyl acetate

2.1.1. Substrates and catalyst

Clove essential oil (Caryophyllus aromaticus) was purchased from Viafarma (São Paulo-Brazil) and used as substrate for enzymatic esterification. The main characteristics of the oil are presented in Table 1. Acetic anhydride (Vetec, 97% purity) was also used as substrate and the commercial lipase from Candida antarctica (Novozym 435) immobilized on a macroporous anionic resin was donated by Novozymes Brazil (Araucária, PR, Brazil) was used as catalyst. Eugenyl acetate from Sigma-Aldrich was used as standard for product confirmation.

2.1.2. Experimental procedure

The production of eugenyl acetate via enzymatic esterification was carried out in the experimental condition optimized by Vanin et al. (2014). The scheme of the reaction is presented in Figure 1.

The esterification reactions were performed by preparing a reaction mixture of acetic anhydride and clove essential oil (eugenol content of 85.5%) at molar ratio of 5:1 (90-18 mmol) in a 50 mL Erlenmeyer flask.

![Figure 1. Synthesis of eugenyl acetate by enzymatic esterification of clove essential oil.](image)

**Table 1. Characteristics of clove essential oil (Caryophyllus aromaticus).**

| Analysed Parameters | Specification | Result |
|---------------------|---------------|--------|
| Color               | Light and dark yellow | Yellow |
| Density             | 0.953 to 0.965 g/m\textsuperscript{2} (20 °C) | 0.957 g/m\textsuperscript{2} (20 °C) |
| Index of refraction | 1.486 to 1.498 (20 °C) | 1.489(20 °C) |
| Eugenol content*    | 85.0 to 88.0% | 85.43% |

*data from Viafarma laboratory.*
After complete dissolution of the substrates, 5.5 wt% of enzyme (based on the total mass of substrates) was added to the mixture. Experiments were carried out in an orbital shaker at constant agitation of 150 rpm and 50 °C. After 2 h of reaction, the biocatalyst was filtered and samples were kept at 5 °C for further analysis and determination of reaction conversion.

2.1.3. Determination of reaction conversion

Quantitative analyses of eugenyl acetate produced were carried out in a gas chromatography (Shimadzu GC-2010) equipped with data processor, using a capillary column of fused silica INOWAX (30 m length × 250 μm i.d. × 0.25 μm thickness), flame ionization detector, with the following temperature program: 40-180 °C (3 °C/min), 180-230 °C (20 °C/min), 230 °C (20 min), injector temperature 250 °C, detector at 275 °C, injection in the mode split, ratio of split 1:100, H 2 (56 KPa) as carrier gas, injected volume of 0.4 μL of sample diluted in n-hexane (1:10). Reaction conversion was calculated based on the reduction of area of limiting reagent on the basis of reaction stoichiometry (Paroul et al., 2011).

2.2. Determination of toxicity against Artemia salina

To evaluate the toxicity of both clove essential oil and eugenyl acetate against Artemia salina, after esterification, the reaction mixture was submitted to vacuum microdistillation, to remove any residue of acetic acid and acetic anhydride from the final sample, avoiding, in this way, an incorrect interpretation of the results. In order to ensure that residual esterification products did not produce the observed response, the mortality profile of eugenyl acetate ester was assessed in parallel with eugenyl acetate commercial standard (Sigma Aldrich).

Toxicity test was carried out using the methodology described by Meyer et al. (1982) with some modifications. The cysts of Artemia salina were placed in a plastic container with artificial saline solution (23 g of marine salt/1 liter of distilled/deionized water/0.7 g of sodium bicarbonate) with artificial illumination, under aeration, with control of temperature (20-30 °C) during 24 hours of incubation for the hatching. After this period, the organisms-test were exposed to different concentrations of both products to be tested (clove essential oil and eugenyl acetate) for 24 hours, using test tubes, each one containing at least 10 nauplii of Artemia salina, at 10 different concentration of products, in triplicate runs. In a first assay, the range of concentration to be tested was determined (10, 20, 30, 40, 50, 60, 70, 80, 90 and 100 μg.mL −1 ). After, for eugenyl acetate the assay was performed with smaller concentrations. A control experiment was also carried out to be sure that mortality observed of nauplii of Artemia salina was resulting of toxicity of the compounds and not due to the restriction of food (Carballo et al., 2002). After 24 h of exposure, the counting of alive and dead nauplii was carried out.

The LD 50 values were determined in triplicate employing non-linear regression model available in GraphPad Prism 6.0 software, and expressed as mean ± standard deviation. The means were compared using the t test, adopting 5% as significance level.

3. Results and Discussion

3.1. Enzymatic production of eugenyl acetate

Following the experimental condition described in the previous section (Vanin et al., 2014), the eugenyl acetate conversion was 95.6%. These results are similar to other studies from the literature. Chiaradia et al. (2012) achieved a conversion of about 99% in the esterification of eugenol and acetic anhydride to eugenol molar ratio of 3:1, enzyme concentration of 5.5 wt% (based on the total mass of substrates), 50 °C in 6 hours of reaction. Chaibakhsh et al. (2012) synthesized eugenyl caprylate using Lipozyme TL IM as catalyst in solvent-free system, and obtained a maximum reaction yield of 72.2% at 65 °C, 250 rpm, 100 mg of enzyme, caprylic acid do eugenol molar ratio of 1:2 after 4.5 hours of reaction.

Before evaluating the toxicity of eugenyl acetate against Artemia salina, this compound was purified and its purity was confirmed by gas chromatograph, as shown in Figure 2. As we can observe, the retention time (R t) of the major compound after esterification (Figure 2c) and purification (Figure 2d) (R t = 25.8 min) corresponds to the retention time of standard eugenyl acetate (Figure 2e) (R t = 25.9 min). The peaks of R t = 8.5 min and R t = 11.7 min correspond to the excess of acetic anhydride and acetic acid produced, respectively. A decrease of the peak of eugenol (R t = 24.5 min) after enzymatic esterification was also observed (Figure 2a and 2c).

3.2. Determination of toxicity against Artemia salina

The results related to the toxicity of clove essential oil before and after esterification are presented in Table 2. The percentage of mortality increases proportionally with the concentration, reaching the maxima value of

Table 2. Mean mortality of Artemia salina according to clove essential oil and eugenyl acetate ester concentrations at t = 24 h of exposure.

| Concentration (μg/mL) | Mean of dead nauplii (%) ± standard deviation |
|-----------------------|---------------------------------------------|
|                       | Clove oil                                    | Eugenyl acetate  |
| 0.0                   | 0.0 ± 0.0                                   | 0.0 ± 0.0         |
| 0.1                   | 0.0 ± 0.0                                   | 37.8 ± 3.8        |
| 0.2                   | 15.0 ± 3.1                                  | 80.2 ± 9.5        |
| 0.3                   | 14.1 ± 6.3                                  | 100 ± 0.0         |
| 0.4                   | 24.9 ± 16.3                                 | 100 ± 0.0         |
| 0.5                   | 25.7 ± 13.7                                 | 100 ± 0.0         |
| 0.6                   | 53.2 ± 14.1                                 | 100 ± 0.0         |
| 0.7                   | 45.6 ± 20.0                                 | 100 ± 0.0         |
| 0.8                   | 69.7 ± 19.8                                 | 100 ± 0.0         |
| 0.9                   | 93.3 ± 7.1                                  | 100 ± 0.0         |
| 1.0                   | 100 ± 0.0                                   | 100 ± 0.0         |

Braz. J. Biol., 2017, vol. 77, no. 1, pp. 155-161
mortality, 100%, for concentrations of 1.0 µg·mL⁻¹ and 0.3 µg·mL⁻¹, for clove essential oil and eugenyl acetate, respectively. In absence of test substance was not observed mortality, in both cases. The correlation between the mortality with the log of clove essential oil concentration, log of eugenyl acetate concentration and log of commercial eugenyl acetate allowed to obtain the models described in Figures 3a, 3b and 3c.

Figure 2. Chromatograms of clove essential oil (A), acetic anhydride (B), clove essential oil after enzymatic esterification (C), eugenyl acetate after purification that was used for testing with *A. salina* (D) and the standard of eugenyl acetate (E).
Eugenyl acetate toxicity

Related to clove essential oil, the correlation between the mortality (%) and the log of concentration of oil \( y = \frac{100}{1 + 10^{3.910(-0.2219-x)}} \) with \( R^2 = 0.8959 \) provided a value of \( LC_{50} \) of 0.1178 ± 0.0041 µg.mL\(^{-1}\). We have not found works from the literature related to the larvicidal effect of eugenyl acetate against \( A.\) salina. Eugenyl acetate commercial standard produced a \( LC_{50} \) of 0.1041 ± 0.0062 µg.mL\(^{-1}\) \( (y = 100/1 + 10^{2.499(-0.9826-x)}; R^2 = 0.9518) \). Considering this value, the toxicological effect produced by ester obtained from enzymatic esterification was not associated with residual reaction products.

A smaller concentration of ester was necessary for produce 50% of response, if compared with clove oil \((p<0.0001)\). The results analysis of \( LC_{50} \) before and after the enzymatic esterification, showed the increase of about 5 times after the reaction. Both concentrations found, for the oil and its ester, indicate the toxicity of the compounds. The World Health Organization reports that plant extracts showing \( LC_{50} \) < 0.4 µg.mL\(^{-1}\) has some potential to be applied as molluscicidal or larvicidal compound (WHO, 1993).

Different work have been published in the literature indicating the good larvicidal activity of essential oils on several species against \( A.\) salina, but with values of \( LC_{50} \) higher than those obtained in the present work. Oliveira et al. (2011) evaluated the effect of \( Pectis\) brevipedunculata essential oil in \( A.\) salina and obtained values of \( LC_{50} \) of 36 µg.mL\(^{-1}\) for the oil extracted from the plant \textit{in natura} and 19 µg.mL\(^{-1}\) for the oil extracted from plants dried at 40 °C.

Costa et al. (2010) evaluated the extracts of different species of medicinal plants under larvae of \( A.\) salina based on the percentage of mortality, after 24 h exposure to the treatments. All species tested showed good larvicidal activity as compared to a reference compound and literature data. The extract from \( Vanillosmopsis\) arborea was the most active with an \( LC_{50}\) of 3.9 µg.mL\(^{-1}\). Arcanjo et al. (2012) evaluated different medicinal plants and observed a higher larvicidal effect under \( A.\) salina for extracts of flowers of \( Acmella\) uliginosa \( (LC_{50}\) of 18.76 µg.mL\(^{-1}\)).

Regarding the interesting in vitro toxicity of the assayed essential oils against \( A.\) salina and the importance of the diseases with larval vector in the public health, our results are promising, however it is necessary that complementary researches are performed focusing the possibility of their practical and rational application to impair the survival of the different larval etiological agent.

The enzymatic esterification of clove essential oil to the production of eugenyl acetate using Novozym 435 as catalyst showed to be a promising route to the obtainment of esters, taking into account the mild reaction conditions and the low amount of catalyst needed to conduct to high process conversion. The results obtained in this work were similar to obtained by Vainin et al. (2014), where the results of esterification of clove essential oil are better discussed. The low lethal concentrations obtained for both clove essential oil and eugenyl acetate could also indicate toxicity to other organisms such as larvae of insects vector of diseases.

Figure 3. Toxicological effect of clove oil (a) its eugenyl acetate ester (b) and commercial eugenyl acetate (c) on \( A.\) salina.
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Acknowledgements

The authors thank CAPES, CNPq, FAPERGS and URI-Erechim by the financial support and scholarships.
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