Contents lists available at ScienceDirect

Biochemical Systematics and Ecology

journal homepage: www.elsevier.com/locate/biochemsyseco

Sequestration of tropane alkaloids from *Brugmansia suaveolens* (Solanaceae) by the treehopper *Alchisme grossa* (Hemiptera: Membracidae)

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A R T I C L E I N F O

Article history: Received 5 January 2016 Accepted 20 March 2016 Available online 19 May 2016

Keywords: Scopolamine Sequestration Membracids Host specialization Plant secondary metabolites Alkaloids

ABSTRACT

Treehoppers (Hemiptera: Membracidae) are sap-feeding insects distributed mainly in tropical regions. *Alchisme grossa* is a treehopper that has been reported in the Bolivian Yungas forests using mostly *Brugmansia suaveolens* (Solanaceae) as host-plant, where adult females oviposit and take care of their nymphs until they molt to adults. *Brugmansia* is a subtropical genus producing a variety of tropane alkaloids (TAs). We herein report the sequestration by adult males and females of *A. grossa* of TAs from *B. suaveolens*, examining separately the distinct body sections of insects. Purified extracts of *A. grossa* and *B. suaveolens* were analyzed by gas chromatography/mass spectrometry. TAs in *A. grossa* were the same as those in its host-plant; furthermore, they were equally distributed between sexes and they were differentially allocated within the body of adult individuals. An ecological role for sequestered TAs is discussed.

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1. Introduction

Treehoppers (Hemiptera: Membracidae) are a diverse group (approximately 3300 spp.) of phytophagous insects distributed mainly in tropical regions of the world (Wood, 1993; Dietrich et al., 2001). They are sap feeders and their hostplants constitute a site for reproduction, oviposition and development. A conspicuous and morphologically variable pronotum constitutes a characteristic trait of treehoppers; this structure has been assigned a role in crypsis (Roy et al., 2007) and defense (Wood, 1974). Species range from monophagous to polyphagous (Dietrich and Deitz, 1991; Wood, 1993; Lin, 2006); diet breadth has been related to latitude, where tropical species tend to be oligophagous or polyphagous and temperate species tend to be monophagous, and to altitude, where higher elevations seemingly promote host specialization (Wood and Olmstead, 1984; Wood, 1984).

http://dx.doi.org/10.1016/j.bse.2016.03.015 0305-1978/© 2016 Elsevier Ltd. All rights reserved.







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The genus *Alchisme* thrives in humid montane and submontane ecosystems from northern Central America to the Brazilian shield and northern Chile, and shows preference for solanaceous host-plants (McKamey and Deitz, 1996; Godoy et al., 2006). *Alchisme grossa* is a subsocial species which displays diverse and complex maternal care traits such as egg and nymphal guarding, active defense against predators, and feeding facilitation (Godoy et al., 2006; Camacho et al., 2014; Torrico-Bazoberry et al., 2014). Adult females of *A. grossa* remain on the host-plant where they oviposit and take care of their nymphs until they molt to adults (Torrico-Bazoberry et al., 2014). In the summer (wet) season in the Yungas biogeographical region of Bolivia, *A. grossa* females oviposit mostly on young leaves of *Brugmansia suaveolens* (Solanaceae) (Torrico-Bazoberry et al., 2014), in spite of this being a habitat characterized as a site of high diversity of solanaceous species (Nee et al., 2007).

Host plant specialization is a common characteristic observed in phytophagous insects; the process is attributed in many cases to plant secondary metabolites which are a potential factor in the evolution of specific associations (Nishida, 2002; Schoonhoven et al., 2005). The Solanaceae is a large family comprising around 3500 species (D'Arcy, 1986), many of which contain alkaloids (Evans, 1986; Hawkes et al., 2000); in particular, *Brugmansia* produces tropane alkaloids (TAs) (Doncheva et al., 2006) which exhibit different levels of toxicity towards a wide variety of insects (Krug and Proksch, 1993; Detzel and Wink, 1993; Kitamura et al., 2004; Jolivet et al., 2012). In this study, we enquire about the sequestration of TAs from *B. suaveolens* in different body sections by *A. grossa* and their possible ecological role in the context of its pattern of specialization on this particular solanaceous species.

2. Materials and methods

2.1. Alchisme grossa and B. suaveolens

Periodic field observations showed that cohorts of *A. grossa* were synchronized, allowing collection of ten males and ten females, one from each of 20 different cohorts, within two days of having reached the adult stage. Immediately after collection, insects were taken to the laboratory, transferred to tomato plants (*Lycopersicum esculentum*), enclosed in tulle bags and allowed to feed for 24 h. The aim of this process was to allow insects to replace *B. suaveolens*-related contents in its gut by tomato-related contents (in terms of alkaloids, glycosteroidal alkaloids – Friedman, 2002); under these circumstances, TAs found in *A. grossa* should correspond only to compounds sequestered in the insect body.

Four groups of young leaves (ca. 20 g fr. wt each) were collected from different uncolonized individuals of *B. suaveolens*, carried to the laboratory, dried at 35 °C (Heraeus UT6 oven) for three days, ground in a laboratory mill and analyzed for TAs.

2.2. Extraction of alkaloids from leaves of B. suaveolens

Each pulverized sample (2 g) of *B. suaveolens* leaves was extracted with 40 ml CH₃OH and exposed to ultrasound in a bath (Power Sonic 405) at 25 °C for 30 min. The methanolic extract was filtered through a frit funnel and the resulting extract evaporated under reduced pressure on a rotatory evaporator (Büchi RE 111). The syrupy residue was dissolved in 3 ml 5% HCl. The acidic solution was washed with CHCl₃ (2×3 ml). The aqueous phase was adjusted to pH 10 with NH₄OH and was extracted twice with 3 ml CHCl₃; under these circumstances the aqueous extracts gave negative Dragendorff reaction. Finally, the organic extracts were dried with anhydrous Na₂SO₄, filtered through cotton wool placed at the tip of a Pasteur pipette and taken to dryness by means of a flow of nitrogen. Total alkaloids as free bases constituted on average of 2.1% dry weight of the plant extracts. The dry residues were disolved in 10 μ l methanol before injecting 2 μ l into the GC column.

2.3. Extraction of alkaloids from insects

Insects were sacrificed by freezing and then dried at 35 °C (Heraeus UT6 oven) for three days. Each adult was weighed $(8.1 \pm 1.2 \text{ mg dry weight; mean} \pm \text{SD})$ and its body dissected under a stereoscopic magnifying lens (Olympus SZ61) into three sections: a) pronotum $(18.7 \pm 0.024\% \text{ d.w.})$, b) head + thorax + abdomen $(66.3 \pm 0.083\% \text{ d.w.})$, and c) wings + legs $(15.0 \pm 0.028\% \text{ d.w.})$. Each section was ground by introducing it into a 1.8 ml stainless steel microvial with a polyethylene flange cap containing five 1.5-mm-diameter steel spheres and agitating it for 10 min in a bead beater (Mini-Beadbeater-96; Biospec Inc., Bartlesville, OK, USA). The pulverized sample was extracted with 2 ml CH₃OH and exposed to ultrasound in a bath (Power Sonic 405) at 25 °C for 30 min. The methanolic extract was not submitted to acid-base purification to avoid losses; it was filtered through cotton wool placed at the tip of a Pasteur pipette, collected in a 1.8 ml amber vial and dried under nitrogen flow. The dry residue was successively redissolved in small aliquots of methanol (ca. 50 µl) which were transferred to a 100 µl glass insert within an amber vial and evaporated to dryness by means of a nitrogen flow; this operation minimized the quantity of residue retained in the original vial walls. The dry residues were dissolved in 10 µl methanol before injecting 2 µl into the GC column.

2.4. Analysis of extracts by gas chromatography/mass spectrometry (GC/MS)

Purified extracts of *A. grossa* and *B. suaveolens* were analyzed by GC/MS (Shimadzu, GCMS-QS, 2010 Ultra), equipped with an Rtx-5MS Crossbond 5% diphenyl - 95% dimethyl polysiloxane capillary column (30 m length, 0.25 μ m film thickness, 0.25 mm internal diameter). The GC was operated in the splitless injection mode; injection volume was 1 μ l for plant extracts

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