



Host susceptibility to *Gonipterus platensis* (Coleoptera: Curculionidae) of *Eucalyptus* species

Catarina I. Gonçalves¹ · Liliana Vilas-Boas² · Manuela Branco² · Gabriel D. Rezende¹ · Carlos Valente¹

Received: 3 December 2018 / Accepted: 9 June 2019 / Published online: 20 June 2019
© INRA and Springer-Verlag France SAS, part of Springer Nature 2019

Abstract

• **Key message** *Gonipterus platensis* is an important insect pest of eucalypt plantations. Despite biological control by the egg parasitoid *Anaphes nitens*, economic losses remain high in several areas outside its native range where susceptible eucalypt species are grown in commercial plantations. The susceptibility to *G. platensis* of 17 *Eucalyptus* species was evaluated and possible alternatives for reforestation in high pest incidence areas were identified.

• **Context** *Gonipterus platensis* is an important pest of *Eucalyptus* worldwide. Despite biological control, it causes significant losses to *Eucalyptus* plantations in several areas, requiring alternative management options.

• **Aims** We analysed host preference of *G. platensis* towards 17 *Eucalyptus* species to identify less susceptible plant materials that could be used in areas of high pest incidence.

• **Methods** Feeding damage was assessed in field trials in three consecutive years. No-choice and choice tests were conducted with *Eucalyptus* species of contrasting susceptibility.

• **Results** Within subgenus *Symphyomyrtus*, all species from section Maidenaria were used by *G. platensis* for feeding. Within this section, *E. globulus* was always the preferred species, while *E. nitens* was the least preferred. Differences in susceptibility were less pronounced at high attack intensity by *G. platensis*. *Eucalyptus saligna* (section Latoangulatae) was the least preferred species among *Symphyomyrtus*. All species from subgenus *Eucalyptus* had low susceptibility to *G. platensis*, particularly *E. regnans*, which was never attacked under field conditions. The results were confirmed by choice and no-choice laboratory and semi-field tests.

• **Conclusion** Significant differences in susceptibility to *G. platensis* were found between the 17 *Eucalyptus* species tested, which could be explored for reforestation with less susceptible plant materials.

Keywords Tree susceptibility · Defoliation · *Eucalyptus* snout beetle · Reforestation

Handling Editor: Aurélien Sallé

Contributions of the co-authors Conceptualization and experimental design: CV, GDR, CIG, MB, LVB; Field and laboratory work: LVB, CIG; Data analysis: LVB, MB, CIG; Manuscript writing: CIG, LVB; Manuscript reviewing and editing: MB, CV, GDR; Project Administration: GDR, MB, CV; Funding acquisition: GDR, MB, CV.

✉ Catarina I. Gonçalves
catarina.goncalves@thenavigatorcompany.com

¹ RAIZ - Instituto de Investigação da Floresta e Papel, Quinta de São Francisco, Apartado 15, 3801-501 Eixo-Aveiro, Portugal

² Centro de Estudos Florestais, Instituto Superior de Agronomia, Universidade de Lisboa, Tapada da Ajuda, 1349-017 Lisbon, Portugal

1 Introduction

Eucalypts (*Eucalyptus* L'Hér., Myrtaceae) are among the most used tree species in plantations worldwide, primarily because of their fast growth and adaptability to environmental conditions (Campinhos 1999; Wingfield et al. 2008). In Europe, eucalypts are mainly cultivated in Portugal and Spain for the pulp and paper industry, with an overall area of roughly 810,000 ha in Portugal (ICNF 2013) and 630,000 ha in Spain (SECF 2010). *Eucalyptus globulus* Labill. is the most planted species in these two countries because of its high quality for pulping (Costa e Silva et al. 2009; González-García et al. 2009). *Eucalyptus* snout beetles, *Gonipterus* spp. (Coleoptera: Curculionidae), are among the most severe pests of *Eucalyptus*. *Gonipterus platensis* Marelli in particular is the most widespread species found outside Australia

(Mapondera et al. 2012). This pest is present in New Zealand, eastern and western South America, southwestern North America, and southwestern Europe (Mapondera et al. 2012).

Soon after its detection in Spain in 1991, *G. platensis* became the most serious pest of *E. globulus* plantations in Galicia (Spain) and northern Portugal (Mansilla-Vázquez and Pérez-Otero 1996; Sousa and Ferreira 1996). The snout beetle feeds on eucalypt leaves, with marked preference for growing shoots with developing leaves, for both feeding and oviposition. Adults will preferably feed along the edges of leaves but they will also feed on the soft bark of fresh shoots during periods of heavy infestation. The larvae, which cause most damage, feed on the entire leaf lamina, leaving only the harder fibres (Loch 2006; Tooke 1955). Severe and repeated defoliation can lead to thinning of the upper crown or ‘broom-topping’, reduced growth, tree deformation, and tree decline (Echeverri-Molina and Santolamazza-Carbone 2010; Loch 2006; Loch and Matsuki 2010; Tooke 1955). Biological control was one of the first strategies used to control *G. platensis*. The Australian egg parasitoid *Anaphes nitens* Girault (Hymenoptera: Mymaridae) has been introduced into all continents where the snout beetle is present, with good results (Arzone 1985; Cordero-Rivera et al. 1999; Hanks et al. 2000; Lanfranco and Dungey 2001; Tribe 2005). However, the parasitoid has not been efficient in reducing the damage caused by *G. platensis* below economically sustainable levels in several regions, in particular in cooler regions of countries such as Portugal, Spain, or Chile (Cordero-Rivera et al. 1999; Lanfranco et al. 2011; Reis et al. 2012; Valente et al. 2018). In Portugal, reductions in wood volume are estimated to be as high as 86% in high altitude regions where parasitism rates by *A. nitens* are low during winter and early spring (Reis et al. 2012), resulting in estimated losses of 648 million euros over a 20-year period (Valente et al. 2018).

Faced with significant economic losses, stakeholders are searching for alternative management strategies, such as biological control with alternative natural enemies or using eucalypt species that are less susceptible to *G. platensis* (Jactel et al. 2009; Richardson and Meakins 1986; Valente et al. 2017). Although several authors have focussed on host susceptibility to *Gonipterus* spp., such studies often dealt with distinct species within the snout beetle complex (Mapondera et al. 2012), resulting in discrepancies in literature (Newete et al. 2011). In countries where *G. platensis* is present, *E. globulus* is consistently found to be a preferred host, even though several other species have also been identified as susceptible, such as *Eucalyptus camaldulensis* Dehnh., *E. grandis* W.Hill, *E. longifolia* Link, *E. obliqua* L'Hér., *E. propinqua* Deane & Maiden, *E. robusta* Sm., and *E. viminalis* Labill. (Cordero-Rivera and Santolamazza-Carbone 2000; Hanks et al. 2000; Huerta-Fuentes et al. 2008; Lanfranco and Dungey 2001).

In the present study, we analysed the susceptibility of 17 *Eucalyptus* species to *G. platensis*, including *E. globulus*, in order to identify potential alternative species for reforestation in the areas most affected by the pest. Because these areas are mostly located in cool high altitude regions, the *Eucalyptus* species were pre-selected for their ability to withstand low temperatures. Host plant susceptibility to *G. platensis* was tested under field conditions by evaluating naturally occurring defoliation and insect abundance in field trials for 3 years. In addition, no-choice and choice tests with selected *Eucalyptus* species of contrasting susceptibility were conducted under semi-field and laboratory conditions.

2 Material and methods

2.1 Study sites and plant materials

Sixteen alternate *Eucalyptus* species to *E. globulus* were pre-selected for their adaptation to low temperatures. The seeds were obtained from CSIRO (Commonwealth Scientific and Industrial Research Organization, Australia). One to three provenances of each species were used (Table 1) to assure that each species was well represented. One seedling provenance and four commercial clones of *E. globulus* were also used, totalling 47 distinct provenances. All plants were produced in RAIZ and Viveiros Aliança nurseries (Pegões, Portugal). Field studies were conducted in three trials in Portugal, located in Carregal do Sal, hereafter Carregal (40° 47' 00" N, 8° 04' 30" W), Pampilhosa da Serra, hereafter Pampilhosa (40° 06' 30" N, 7° 47' 38" W), and Arouca (40° 58' 04" N, 8° 07' 14" W), planted in May 2010, October 2010, and April 2011, respectively. Seventeen species were planted in Carregal (47 provenances) and Pampilhosa (45 provenances), and fifteen species were used in Arouca (39 provenances), due to lower plant availability for later trials. In each trial, eight randomized blocks consisting of five plants of each provenance were used (1880, 1800, and 1375 plants per trial in Carregal, Pampilhosa, and Arouca, respectively). The number of plants available for the evaluation of *G. platensis* defoliation was variable due to mortality in the months following planting. For the purpose of evaluating susceptibility to the pest, all provenances of a single species were used in the analyses.

2.2 Field trial evaluation

The level of defoliation caused by *G. platensis* feeding by both adults and larvae was evaluated after spring feeding (between July and November in 2012, 2013, and 2014) in the Arouca and Pampilhosa trials. The snout beetle was not detected in Carregal in any year. Defoliation was evaluated by visual estimate of the leaf area consumed by *G. platensis* larvae and adults in recent foliage, in the upper third of the crown

Table 1 Eucalypt species and provenances used in field trials (*ACT*, Australian Capital Territory; *AUS*, Australia; *BR*, Brazil; *NSW*, New South Wales; *PT*, Portugal; *QLD*, Queensland; *SA*, South Africa; *TAS*, Tasmania; *VIC*, Victoria)

Subgenus ¹	Section ²	Series ²	Species	Provenance				
<i>Eucalyptus</i>	Cineraceae	Fraxinales	<i>Eucalyptus delegatensis</i> R.T.Baker	Rubicon, VIC (AUS) Mount Ewen Dargo-Bright, VIC (AUS) Bulls Head, ACT (AUS)				
			<i>Eucalyptus fraxinoides</i> H.Deane & Maiden	Southeast of Canberra, NSW (AUS) Nimmitabel, NSW (AUS) Badja, NSW (AUS)				
			<i>Eucalyptus oreades</i> R.T.Baker	Newnes State Forest, Lithgow, NSW (AUS)				
	Eucalyptus	Regnantes	<i>Eucalyptus fastigata</i> H.Deane and Maiden	Tallaganda State Forest, NSW (AUS) Brown Mountain Nimmitabel, NSW (AUS) Errinundra Plateau, VIC (AUS)				
			<i>Eucalyptus regnans</i> F.Muell.	Lisle, TAS (AUS) Traralgon, VIC (AUS) Moogara, TAS (AUS)				
			Latoangulatae	Transversae	<i>Eucalyptus saligna</i> Sm.	Blackdown Tableland, QLD (AUS) Styx River SF339, NSW (AUS) Richmond Range, NSW (AUS)		
					Maidenaria	Benthamianae	<i>Eucalyptus benthamii</i> Maiden & Cambage	SSO Crossley, NSW (AUS) Embrapa Florestas, Paraná (BR)
							Bridgesianae	<i>Eucalyptus dunnii</i> Maiden
			Compactae	<i>Eucalyptus badjensis</i> Beuzev. & M.B.Welch	<i>Eucalyptus smithii</i> R.T.Baker	Glenbog, NSW (AUS) Brown Mountain, NSW (AUS) Deua National Park, NSW (AUS) Tallaganda State Forest, NSW (AUS) Wingello State Forest, NSW (AUS) Mount Dromedary, NSW (AUS)		
						Foveolatae	<i>Eucalyptus macarthurii</i> H.Deane & Maiden	Pietermaritzburg (SA) Long Swamp Creek, NSW (AUS) Paddys River, NSW (AUS)
Globulares	<i>Eucalyptus bicostata</i> Maiden, Blakely & Simmonds	Narrow Neck, NSW (AUS) Wee Jasper, NSW (AUS) Nullo Mountain State Forest, NSW (AUS) 4 clones and 1 seedlot, Viveiros Aliança (PT) Yambulla State Forest, NSW (AUS) Myrtle Mountain, NSW (AUS) Monga, NSW (AUS)						
		<i>Eucalyptus globulus</i> Labill. <i>Eucalyptus maidenii</i> F.Muell.						Myrtle Mountain, NSW (AUS) Monga, NSW (AUS)
Viminalales	<i>Eucalyptus nitens</i> Maiden	<i>Eucalyptus nitens</i> Maiden				Blue Range Road, VIC (AUS) Tallaganda State Forest, NSW (AUS) Ebor, NSW (AUS)		
			<i>Eucalyptus dalrympleana</i> Maiden <i>Eucalyptus viminalis</i> Labill.	Mount Canobolas Orange, NSW (AUS) Otway, VIC (AUS) Canobolas State Forest, NSW (AUS) Glenbog State Forest, NSW (AUS)				

¹ Sensu Hill and Johnson (1995)² Sensu Brooker (2000)

where insect feeding is strongly concentrated. Tree height (on average 6.7 m in 2014) was an impediment to the use of a more classical Crown Damage Index (e.g. Stone et al. 2003, developed for young eucalypt plantations), and therefore defoliation intensity was assessed using the following classes: 1 (no evidence of insect feeding); 2 (<25% of the leaves with evidence of insect feeding); 3 (25–50% of the leaves with evidence of insect feeding); 4 (50–75% of the leaves with evidence of insect feeding); 5 (>75% of the leaves with evidence of insect feeding and at least one fourth of the leaves having more than 50% of their area intact); 6 (>75% of the leaves with evidence of insect feeding and less than one fourth of the leaves having more than 50% of their area intact); and 7 (100% defoliation). Mean defoliation class values were then assigned to each sampled tree as follows: 1 (0%); 2 (12.5%); 3 (37.5%); 4 (62.5%); 5 (81.3%); 6 (93.5%); and 7 (100%). Because some of the eucalypt species tested are heteroblastic, i.e. they produce juvenile foliage early in their development, which is replaced by morphologically and physiologically distinct adult foliage (Gosney et al. 2014; Steinbauer 2002), the presence of juvenile foliage in the upper third of the crown was recorded for each tree in 2012 and 2013. Trees were classified as having fully transitioned to adult foliage or as retaining juvenile foliage (totally or partially).

The abundance of *G. platensis* was evaluated in May and June 2012, in Pampilhosa and Arouca trials, respectively. Five trees (1.5 to 2.0 m in height) of each *Eucalyptus* provenance were selected and the number of snout beetle adults, larvae, and egg capsules was visually estimated in the canopy according to the following categories: 0 (no insects); 1 (1–10 insects); 2 (11–20 insects); and 3 (more than 20 insects).

During field evaluations, the presence of biological agents other than the snout beetle was recorded, but neither was found to reach damaging levels.

2.3 No-choice and choice tests

Adults of *G. platensis* were collected from an infested *E. globulus* stand near São Pedro do Sul (40° 47' 00" N, 8° 04' 30" W) and taken to RAIZ laboratory where they were weighed and sorted by sex.

No-choice tests were performed at the Carregal field trial site, as no *G. platensis* adults or larvae were present in this location. Five eucalypt species were selected based on differences in susceptibility to *G. platensis* observed at the Arouca and Pampilhosa trials, namely *E. globulus*, *E. badjensis* Beuzev. & M.B.Welch, *E. smithii* R.T.Baker, *E. nitens* Maiden, and *E. regnans* F.Muell.. Branches with similar length and number of newly expanded adult leaves were enclosed in 1 × 1-mm mesh sleeves (55 cm in length × 18 cm in diameter). One sleeve was placed in each of ten trees per eucalypt species. For each eucalypt species, trees of similar size in at least two different blocks were selected. Eight

randomly selected and previously weighed *G. platensis* adults (four males and four females) were placed inside each sleeve in March 2013. Insect weight at the beginning of the trial was confirmed not to differ between eucalypt species (Wald $\chi^2 = 1.6$; df = 4; $p = 0.803$). One month later, the snout beetles were removed, counted, and transferred to a new branch of the same tree. At the end of each month, the sleeved branches were removed and taken to the laboratory where their leaves were scanned. The leaf area consumed by *G. platensis* adults was then determined using Image J version 1.48 software (National Institute of Health, Bethesda, MD, U.S.A.). Total leaf area lost due to feeding was estimated by adding the area consumed in both periods. The surviving adult snout beetles were counted and weighed.

Choice tests were performed in the forest entomology laboratory in Instituto Superior de Agronomia (University of Lisbon). Three *Eucalyptus* species with contrasting susceptibility were selected, based on field experiments, namely *E. globulus* (high susceptibility), *E. nitens* (moderate susceptibility), and *E. regnans* (low susceptibility). Three host combinations were used: *E. globulus* vs *E. nitens*, *E. globulus* vs *E. regnans*, and *E. nitens* vs *E. regnans*. Trials were performed in a cylindrical see-through cage (60 cm length × 25 cm diameter) closed with a net at the top to allow airflow. Ca. 15 cm long branches with newly expanded adult leaves were collected from trees in the Carregal trial. For each species combination, one branch of each host species was placed on opposite sides of the cage. Branches were inserted in water soaked floral foam to maintain leaf turgor. Two *G. platensis* adults (one male and one female) were placed in the middle of the cage, and the number of feeding and contact with leaves events were recorded at 5 min intervals for 30 min. Forty replicates were used for each host combination. Before each trial, adults were starved for 24 h. Experiments were carried out in room conditions, between 11 am and 4 pm, over several consecutive days. The cylindrical cage was rotated after each replicate to avoid external effects (e.g. light).

2.4 Statistical analysis

Differences in *G. platensis* defoliation in the field between *Eucalyptus* species and years (fixed factors) were tested with Linear Mixed Models (LMM), considering provenance and leaf stage as random factors and trees as subjects. Leaf stage was included as a categorical variable with two levels: trees that had fully transitioned to adult foliage and trees that totally or partially retained juvenile foliage. *Eucalyptus* species with no evidence of defoliation in the field were excluded from the analysis. Least significant differences (LSD) were used for multiple comparisons. Abundance classes of adults, larvae, and egg capsules were converted as follows: 0 = 0 insects; 1 = 5 insects; 2 = 15 insects; and 3 = 30 insects. Mean values were then used to compare between *Eucalyptus* species using

non-parametric Kruskal-Wallis tests for the Arouca and Pampilhosa trials independently followed by pairwise comparisons with Mann-Whitney tests. In no-choice tests, *G. platensis* initial and final weight was compared among *Eucalyptus* species by Generalized Linear Model (GLM), with Gaussian distribution. Individual trees were considered subsamples nested within species. GLM with gamma distribution was used to test differences in leaf area consumed, and GLM with binomial distribution (alive/dead) was used to test differences in mortality. GLM tests were followed by pairwise comparisons. In choice tests, insect preference was analysed with Wilcoxon signed-rank test. Homoscedasticity and normality were confirmed with Levene and Kolmogorov-Smirnov tests, respectively, for the LMM test (Zar 1996). All analyses were performed with SPSS statistics package 22.0 (SPSS 2013) with a 5% ($\alpha = 0.05$) significance level.

3 Results

3.1 Field trials

Levels of defoliation by the snout beetle differed significantly between *Eucalyptus* species at both Arouca ($F_{12,24} = 26.9$; $p < 0.001$) and Pampilhosa ($F_{15,29} = 63.3$; $p < 0.001$). The level of tree defoliation also differed between years at both Arouca ($F_{2,2823} = 599.5$; $p < 0.001$) and Pampilhosa ($F_{2,3525} = 759.7$; $p < 0.001$). In Arouca, defoliation levels decreased along the 3 years (Fig. 1), with overall mean values of $27.2 \pm 1.0\%$, $13.1 \pm 0.7\%$, and $0.7 \pm 0.1\%$ for 2012, 2013, and 2014, respectively. In Pampilhosa, an opposite increasing trend in defoliation levels was found, with mean values of $23.2 \pm 0.9\%$, $35.2 \pm 1.0\%$, and $66.4 \pm 1.1\%$ for 2012, 2013, and 2014, respectively. All species in which defoliation occurred at the Pampilhosa site displayed a similar increasing tendency (Fig. 1).

Overall, *Eucalyptus* species in section Maidenaria were more defoliated than species belonging to any other section. In both trials, *E. globulus* was the most defoliated species, followed by *E. viminalis* and *E. smithii*. When damage by *G. platensis* was at low-medium intensity, some species within Maidenaria section were less attacked, particularly *E. benthamii* Maiden & Cambage and *E. nitens*. However, at high densities, such differences were no longer evident (Fig. 1 and Table 2). All species from subgenus *Eucalyptus* as well as *E. saligna* Sm. (section Latoangulatae) displayed low susceptibility to *G. platensis*, even though *E. fraxinoides* H.Deane & Maiden and *E. oreades* R.T.Baker were moderately defoliated in Pampilhosa in 2014. *Eucalyptus regnans* was not attacked by *G. platensis*, whereas *E. delegatensis* R.T.Baker and *E. fastigata* H.Deane and Maiden were only slightly defoliated (less than 3% defoliation), even when snout beetle populations caused the highest defoliation values in 2014 in Pampilhosa (Fig. 1 and Table 2).

Insect abundance in the canopies was found to differ significantly between *Eucalyptus* species in both Arouca ($\chi^2 = 90.3$, $\chi^2 = 142.0$, and $\chi^2 = 146.7$ for adult, larva, and egg capsule abundance, respectively; $df = 14$; $p < 0.001$) and Pampilhosa trials ($\chi^2 = 109.4$, $\chi^2 = 145.9$, and $\chi^2 = 146.8$ for adult, larva, and egg capsule abundance, respectively; $df = 16$; $p < 0.001$). Five *Eucalyptus* species consistently hosted the highest numbers of insects of all life stages in both trials, namely *E. dunnii*, *E. smithii*, *E. macarthurii*, *E. globulus*, and *E. viminalis* (Fig. 2). Particularly in the Arouca trial, more than 20 egg capsules or 20 larvae were frequently found in canopies of these species. *Eucalyptus badjensis*, *E. bicostata*, *E. maidenii*, and *E. dalrympleana* were found to host intermediate numbers of *G. platensis* egg capsules and larvae in the Arouca trial, but not in the Pampilhosa trial. Small numbers of insects were found in *E. nitens* and *E. saligna*, whereas no insects were found in any of the tree species belonging to subgenus *Eucalyptus*.

3.2 No-choice and choice tests

In no-choice tests, *G. platensis* adult weight at the end of the 2-month experiment differed significantly between the five *Eucalyptus* species tested (Wald $\chi^2 = 122.6$; $df = 4$; $p < 0.001$; Table 3). Insect weight was highest for *G. platensis* fed on *E. smithii* and *E. badjensis* leaves, and was lowest for those feeding on *E. regnans*. Intermediate values were found for *E. globulus* and *E. nitens*. Leaf area consumed by snout beetle adults was also affected by host species (Wald $\chi^2 = 16.8$; $df = 4$; $p < 0.01$; Table 3). The amount of leaves consumed was significantly greater on *E. badjensis*, *E. globulus*, and *E. smithii* than on *E. regnans*, while *E. nitens* displayed intermediate levels of leaf consumption. Within the 2-month trial period, adult mortality of *G. platensis* was highest on *E. regnans* (40%), intermediate on *E. nitens* (24%), and lowest on *E. badjensis*, *E. smithii*, and *E. globulus* (6% or lower) (Wald $\chi^2 = 46.4$; $df = 4$; $p < 0.001$; Table 3).

In choice tests (Fig. 3), *G. platensis* preferred to feed on *E. globulus* rather than on *E. nitens* ($U = 7.3$; $p < 0.001$) or on *E. regnans* ($U = 10.6$; $p < 0.001$), while *E. nitens* was preferred to *E. regnans* ($U = 6.9$; $p < 0.001$). With regard to contact events between *G. platensis* and host plants, no significant difference was found between *E. globulus* and *E. nitens* ($U = 1.2$; $p = 0.22$), while the frequency of contacts with *E. regnans* was lower than with either *E. globulus* or *E. nitens* (respectively $U = 4.6$ and $U = 5.6$; $p < 0.001$).

4 Discussion

Gonipterus platensis fed on 16 *Eucalyptus* species out of the 17 tested, from both subgenera. Within subgenus *Eucalyptus*, low levels of feeding by *G. platensis* were typically found, particularly on host species from series Regnantes

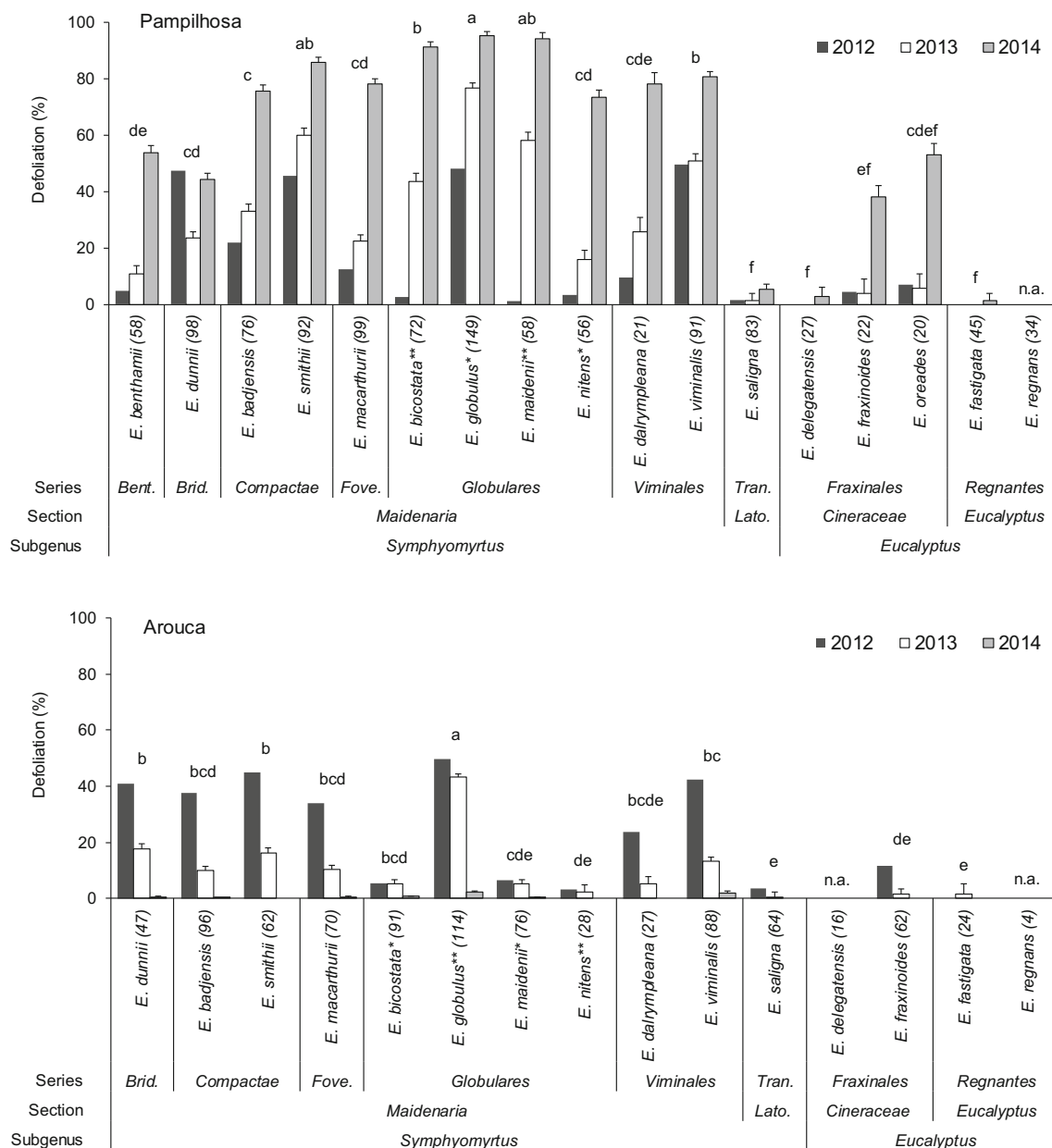


Fig. 1 Defoliation by *Gonipterus platensis* (mean \pm SE) on 17 *Eucalyptus* species in the field trials at Pampilhosa da Serra (top) and Arouca (bottom), between 2012 and 2014. The number of sampled trees is indicated in parenthesis. Different letters indicate significant differences between species for each trial (LMM and LSD multiple comparisons, $p < 0.05$) (Series/section abbreviations: Bent., Benthamianae; Brid., Bridgesianae; Fove., Foveolatae; Tran.,

Transversae; and Lato., Latoangulatae). Notes: * after species names indicates that, on average, 35.2% and 7.8% of trees had not fully transitioned to adult foliage in 2012 and 2013, respectively, underestimating defoliation; ** after species name indicates that, on average, 94.2% and 65.0% of the trees had not fully transitioned to adult foliage in 2012 and 2013, respectively, underestimating defoliation

(*E. regnans* and *E. fastigata*) and on *E. delegatensis*. In this study, *E. regnans* was found to be an unsuitable host for the snout beetle. Within subgenus *Symphyomyrtus*, species from section Maidenaria were consistently attacked by *G. platensis*, although variable levels of defoliation were recorded, while *E. saligna* (section Latoangulatae) displayed low levels of defoliation. A similar pattern was found in a previous study with *G. platensis* in Spain (Cordero-Rivera and Santolamazza-Carbone 2000), with most species belonging

to subgenus *Symphyomyrtus* being moderately to heavily attacked, with the exception of *E. saligna* and *E. rubida* Deane & Maiden (Table 2). Four species were addressed simultaneously in both the present study and the one by Cordero-Rivera and Santolamazza-Carbone (2000), namely *E. globulus*, *E. fastigata*, *E. saligna*, and *E. viminalis*, with consistent results (Table 2). Our study thus provides information on 13 additional *Eucalyptus* species regarding the host range and feeding preference of *G. platensis*.

Table 2 Damage level by *Gonipterus platensis* feeding on eucalypt species in two studies: (1) Cordero-Rivera and Santolamazza-Carbone (2000) and (2) present study. Damage level categorized as: 0: no feeding; *: low (first quartile); **: medium (second and third quartiles); and ***: high (fourth quartile)

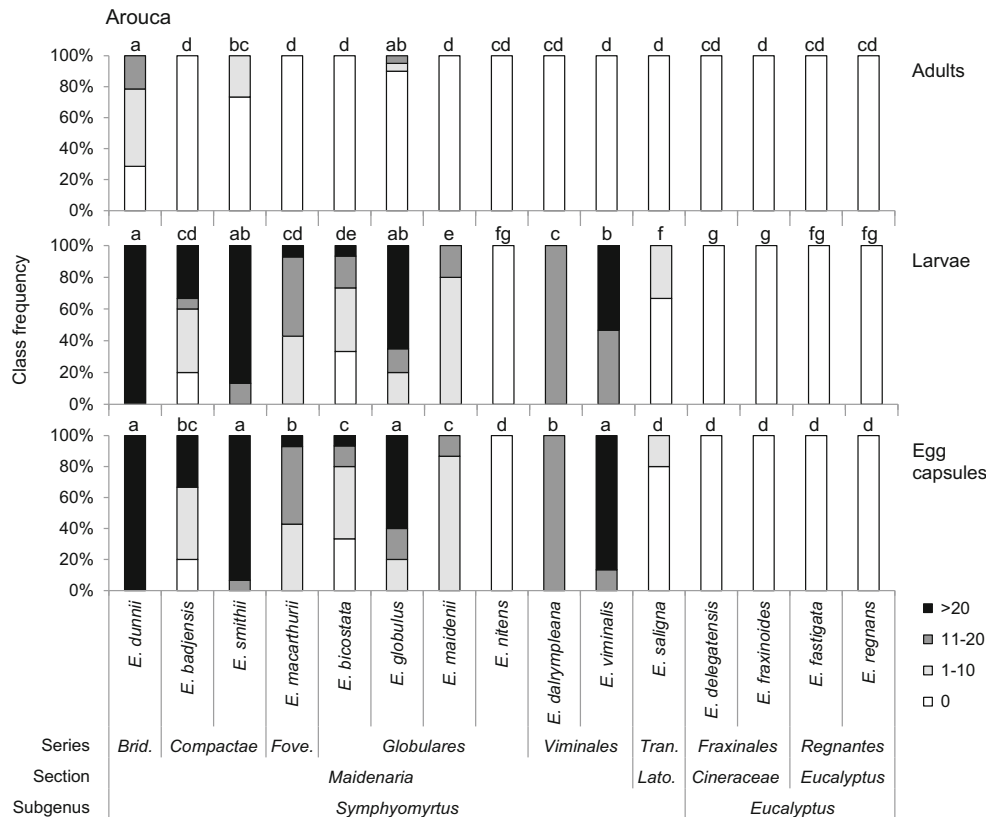
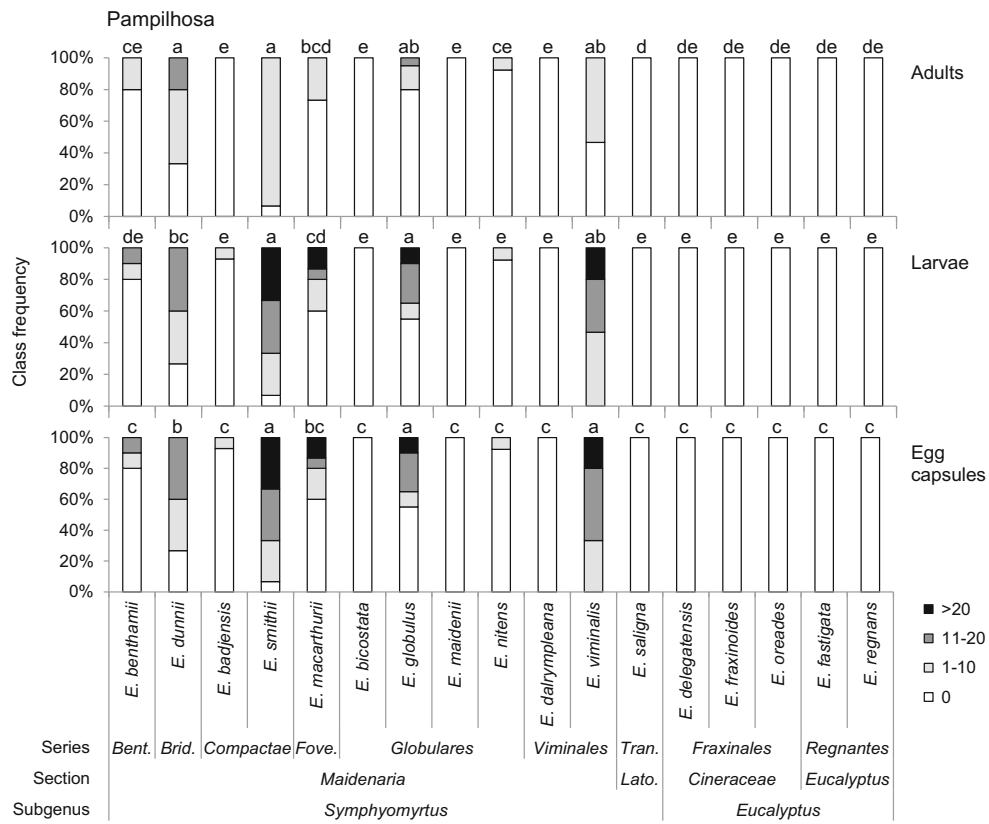
Genus (subgenus) ¹	Section ²	Series ²	Species	(1)	(2) Low pest intensity	(2) High pest intensity
<i>Corymbia</i>	Notiales	Disjunctae	<i>C. ficifolia</i> (F.Muell.) K.D.Hill & L.A.S.Johnson	0	–	–
	Septentrionales	Maculatae	<i>C. citriodora</i> (Hook.) K.D.Hill & L.A.S.Johnson	**	–	–
<i>Eucalyptus</i> (<i>Eucalyptus</i>)	Aromatica	Insulanae	<i>E. amygdalina</i> Labill.	0	–	–
		Radiatae	<i>E. dives</i> Schauer	*	–	–
	Cineraceae	Fraxinales	<i>E. delegatensis</i> R.T.Baker	–	*	*
			<i>E. fraxinoides</i> H.Deane & Maiden	–	*	**
			<i>E. oreades</i> R.T.Baker	–	*	**
	Eucalyptus	Pauciflorae	<i>E. pauciflora</i> Spreng.	**	–	–
		Eucalyptus	<i>E. obliqua</i> L'Her.	0	–	–
		Regnantes	<i>E. fastigata</i> H.Deane and Maiden	0	0	*
	Pseudophloius	–	<i>E. regnans</i> F.Muell.	–	0	0
			<i>E. pilularis</i> Sm.	*	–	–
<i>Eucalyptus</i> (<i>Nothocalyptus</i>)	–	–	<i>E. microcorys</i> F.Muell.	*	–	–
<i>Eucalyptus</i> (<i>Symphyomyrtus</i>)	Bisectae	Cornutae	<i>E. cornuta</i> Labill.	0	–	–
	Latoangulatae	Lepidotae-Fimbriatae	<i>E. propinqua</i> H.Deane & Maiden	***	–	–
		Transversae	<i>E. grandis</i> W.Hill	***	–	–
	Maidenaria	Benthamianae	<i>E. saligna</i> Sm.	*	*	*
			<i>E. benthamii</i> Maiden & Cabbage	–	*	**
			<i>E. dunnii</i> Maiden	–	**	**
		Bridgesianae	<i>E. badjensis</i> Beuzev. & M.B.Welch	–	**	**
			<i>E. smithii</i> R.T.Baker	–	**	***
		Compactae	<i>E. macarthurii</i> H.Deane & Maiden	–	**	**
			<i>E. ovata</i> Labill.	**	–	–
		Globulares	<i>E. bicostata</i> Maiden, Blakely & Simmonds	–	**	***
			<i>E. globulus</i> Labill.	***	***	***
			<i>E. maidenii</i> F.Muell.	–	**	***
	Viminales	<i>E. nitens</i> Maiden	–	*	**	
		<i>E. dabrypleana</i> Maiden	–	**	**	
		<i>E. rubida</i> H.Deane & Maiden	*	–	–	
		<i>E. viminalis</i> Labill.	**	***	***	
	Similares	–	<i>E. longifolia</i> Lindl.	***	–	–

¹ Sensu Hill and Johnson (1995)

² Sensu Brooker (2000)

The Tasmanian native *G. platensis* fed on several host species that are native to other parts of Australia, mostly within section Maidenaria. Closely related plants are likely to share traits that make them similarly acceptable to a particular phytophagous insect (Bertheau et al. 2010; Branco et al. 2014a; Östrand et al. 2008) and this seems to hold true for the snout beetle. In South Africa, Newete et al. (2011) found that *Gonipterus* sp. n. 2 (sensu Mapondera et al. 2012) also preferred to feed on eucalypts from section Maidenaria. In addition, while this *Gonipterus* species was found to accept some hosts within the Latoangulatae section,

namely *E. grandis* and *Eucalyptus propinqua* H.Deane & Maiden, it also displayed low levels of feeding on *E. saligna*. Similar results were described for *G. platensis* by Cordero-Rivera and Santolamazza-Carbone (2000) in Spain. *Gonipterus platensis* and *Gonipterus* sp. n. 2 therefore share considerable overlap in host range. Other *Gonipterus* species display quite a different host range, such as *Gonipterus notographus* Boisduval (sensu Mapondera et al. 2012), which was found to prefer hosts belonging to subgenus *Eucalyptus* rather than *Symphyomyrtus* (Clarke et al. 1998). This is not surprising, as *G. platensis* and



Gonipterus sp. n 2 are related species within the *Gonipterus scutellatus* Gyllenhal complex, while *G. notographus* is placed

in a sister-group (Mapondera et al. 2012) and related insect species often use related hosts (Morse and Farrell 2005; Winkler and

◀ **Fig. 2** Frequency of *Gonipterus platensis* abundance classes for egg capsules, larvae, and adults on 17 *Eucalyptus* species in the field trials at Pampilhosa da Serra (top) and Arouca (bottom), in 2012. Abundance classes used are 0 (no insects); 1–10 (1 to 10 insects); 11–20 (11 to 20 insects); and >20 (more than 20 insects). Different letters indicate significant differences between *Eucalyptus* species for each variable and trial (non-parametric Mann-Whitney *U* test, $p < 0.05$). (Series/section abbreviations: Bent., Benthaminae; Brid., Bridgesianae; Fove., Foveolatae; Tran., Transversae; and Lato., Latoangulatae)

Mitter 2008). Nevertheless, considerable differences between *G. platensis* and *Gonipterus* sp. n. 2 have been found regarding economically important *Eucalyptus* species. While *G. platensis* has been shown to prefer *E. globulus* over other host species (Cordero-Rivera and Santolamazza-Carbone 2000; Hanks et al. 2000; Lanfranco and Dungey 2001; Tooke 1955), this *Eucalyptus* species was only moderately attacked by *Gonipterus* sp. n. 2 (Newete et al. 2011). Inversely, *E. nitens* was moderately attacked by *G. platensis* while it was one of the species preferred by *Gonipterus* sp. n. 2.

Semi-field and laboratory tests mostly confirmed the results obtained under field conditions. Traits like pest survival, food consumption, or weight gain were consistently higher in host species preferred under field conditions, such as *E. globulus*, *E. smithii*, or *E. badjensis*, and lower in the least preferred species. However, under experimental conditions, when insects are forced to feed on a particular plant species, they may use hosts that they would normally not use in the field (Newete et al. 2011; Palmer and Goeden 1991). Indeed, in our study, *G. platensis* never fed on *E. regnans* whenever an alternative host was accessible, but some consumption occurred in the no-choice test. A similar effect was found under field conditions when pest pressure increased. In the Pampilhosa trial, *E. nitens*, *E. benthamii*, *E. fraxinoides*, and *E. oreades* displayed low levels of defoliation when pest pressure was low (2012 and 2013), but defoliation increased disproportionately when pest pressure was high (2014). An increase in the realized host range due to intraspecific competition at high population densities has been observed for other species (Branco et al. 2014b; Castagneyrol et al. 2016; Svanbäck and Bolnick 2007). Under high densities of *G. platensis*, it seems likely that decreased availability of foliage

of the most susceptible hosts due to intense defoliation caused the snout beetle to feed on less preferred species.

Laboratory choice tests against a preferred host such as *E. globulus* seemed adequate for screening susceptibility to *G. platensis* and can precede more laborious field testing. One interesting aspect was that the number of contacts with the host plant often did not indicate evidence of host discrimination, suggesting that feeding was induced by physical and/or chemical cues resulting from direct contact with leaves, as observed in other eucalypt pests (Ohmart et al. 1985; Steinbauer and Matsuki 2004).

Furthermore, during the course of field evaluations, *Eucalyptus* species in series Globulares were shifting to adult foliage. Differences in the response of herbivorous insects to juvenile and adult leaves within *Eucalyptus* species are widely recognized (Gosney et al. 2014; Steinbauer 2002). *Gonipterus* species in particular have been shown to display strong preference for expanding and newly expanded adult leaves (Branco et al. 2016; Loch 2006; Tooke 1955). Therefore, it seems likely that *G. platensis* defoliation of *Eucalyptus* species in series Globulares within the first years following plantation may not accurately characterize long-term susceptibility.

5 Conclusion

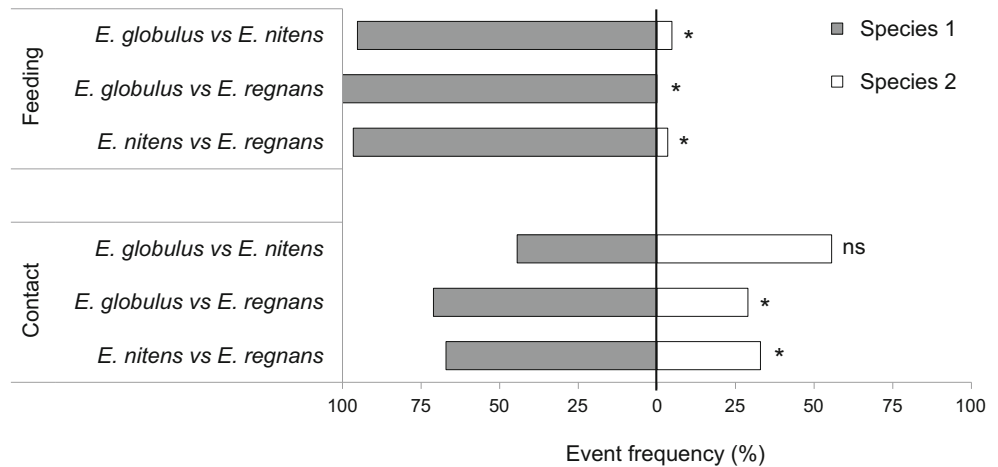
Although *G. platensis* is currently under biological control by the egg parasitoid *A. nitens*, it still causes severe damage in several regions in Portugal where large *E. globulus* plantations are located (Reis et al. 2012; Valente et al. 2018). In these regions, *Eucalyptus* genetic materials less susceptible to *G. platensis* can be a viable option. Here we tested 16 alternative *Eucalyptus* species for potential use in forest plantations in the areas most affected by *G. platensis*. Overall, feeding by the snout beetle was low on species from subgenus *Eucalyptus* and on *E. saligna*. Within subgenus *Symphomyrtus*, all of the tested species in section Maidenaria were susceptible to *G. platensis*, even though some may be good alternatives to *E. globulus*, provided that snout beetle populations are low to moderate.

Table 3 Dry weight, leaf area consumed, and adult mortality (mean \pm SE) by *Gonipterus platensis* adults kept on sleeved branches with five *Eucalyptus* species for 2 months (10 replicates per species) in a semi-field no-choice test. Different letters indicate significant differences between

host plants (GLM with normal distribution for dry weight; GLM with gamma distribution for leaf area consumed; and GLM with binomial distribution for adult mortality, $p < 0.05$; all tests followed by pairwise comparisons)

Species	Dry weight (mg)	Leaf area consumed (cm ²)	Adult mortality (frequency)
<i>E. badjensis</i>	27.1 \pm 0.8 a	62.3 \pm 16.9 a	0.03 \pm 0.02 c
<i>E. globulus</i>	23.9 \pm 0.6 b	48.4 \pm 6.4 ab	0.06 \pm 0.03 c
<i>E. nitens</i>	23.5 \pm 0.7 b	32.6 \pm 5.5 bc	0.24 \pm 0.05 b
<i>E. regnans</i>	19.2 \pm 0.5 c	19.5 \pm 3.3 c	0.40 \pm 0.06 a
<i>E. smithii</i>	27.8 \pm 0.7 a	36.3 \pm 6.2 ab	0.03 \pm 0.02 c

Fig. 3 Percentage of contact and feeding events by *Gonipterus platensis* adults exposed to combinations of three *Eucalyptus* species in choice tests. Species 1 and Species 2 refer to the first and second species in the combination, respectively (Wilcoxon signed-rank test, * $p < 0.001$, ns not significant)



However, before many of these species can be used in commercial plantations, additional information is required on growth and survival under local climatic conditions, and on wood properties for pulping. Moreover, further studies are required on the susceptibility to other pests and diseases, and on how they may affect eucalypt growth in single-species commercial plantations.

Acknowledgements The authors would like to thank Cláudia Ferraz, Fernanda Monteiro, Luís Queirós, José Cardoso, and Rui Gomes for laboratory and field assistance. We are indebted to António Ramos and José Alexandre Araújo for producing the seedlings used in field trials and four anonymous reviewers for valuable comments on the manuscript.

Funding This study was partially funded by projects QREN/5477/PT-LYPTUS, Interreg Forrisk SOE3/P2/F523, and by RAIZ. Centro de Estudos Florestais is a research unit funded by Fundação para a Ciência e a Tecnologia I.P. (FCT), Portugal (UID/AGR/00239/2013).

Data availability The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

References

- Arzone A (1985) Biological control of the *Eucalyptus* snout beetle in Italy. *Bull Sect Reg Ouest Palearct* 8:70–73
- Bertheau C, Brockerhoff EG, Roux-Morabito G, Lieutier F, Jactel H (2010) Novel insect-tree associations resulting from accidental and intentional biological ‘invasions’: a meta-analysis of effects on insect fitness. *Ecol Lett* 13:506–515. <https://doi.org/10.1111/j.1461-0248.2010.01445.x>
- Branco M, Brockerhoff EG, Castagneyrol B, Orazio C, Jactel H (2014a) Host range expansion of native insects to exotic trees increases with area of introduction and the presence of congeneric native trees. *J Appl Ecol* 52:69–77. <https://doi.org/10.1111/1365-2664.12362>
- Branco M, Dhahri S, Santos M, Jamaa MLB (2014b) Biological control reduces herbivore’s host range. *Biol Control* 69:59–64. <https://doi.org/10.1016/j.biocontrol.2013.11.001>
- Branco M, Battisti A, Mendel Z (2016) Foliage feeding invasive insects: defoliators and gall makers. In: Paine TD, Lieutier F (eds) *Insects and diseases of Mediterranean Forest Systems*. Springer, Basel, pp 211–238
- Brooker MIH (2000) A new classification of the genus *Eucalyptus* L’Hér. (Myrtaceae). *Aust Syst Bot* 13:79–148. <https://doi.org/10.1071/SB98008>
- Campinhos E (1999) Sustainable plantations of high-yield shape *Eucalyptus* trees for production of fiber: the Aracruz case. *New For* 17:129–143. <https://doi.org/10.1023/A:1006562225915>
- Castagneyrol B, Jactel H, Brockerhoff EG, Perrette N, Larter M, Delzon S, Piou D (2016) Host range expansion is density dependent. *Oecologia* 182:779–788. <https://doi.org/10.1007/s00442-016-3711-5>
- Clarke AR, Paterson S, Pennington P (1998) *Gonipterus scutellatus* Gyllenhal (Coleoptera: Curculionidae) oviposition on seven naturally co-occurring *Eucalyptus* species. *For Ecol Manag* 110:89–99. [https://doi.org/10.1016/S0378-1127\(98\)00277-1](https://doi.org/10.1016/S0378-1127(98)00277-1)
- Cordero-Rivera A, Santolamazza-Carbone S (2000) The effect of three species of *Eucalyptus* on growth and fecundity of the *Eucalyptus* snout beetle (*Gonipterus scutellatus*). *Forestry* 73:21–29. <https://doi.org/10.1093/forestry/73.1.21>
- Cordero-Rivera A, Santolamazza-Carbone S, Andrés JA (1999) Life cycle and biological control of the *Eucalyptus* snout beetle (Coleoptera, Curculionidae) by *Anaphes nitens* (Hymenoptera, Mymaridae) in north-west Spain. *Agric For Entomol* 1:103–109. <https://doi.org/10.1046/j.1461-9563.1999.00016.x>
- Costa e Silva J, Borralho NMG, Araújo JA, Vaillancourt RE, Potts BM (2009) Genetic parameters for growth, wood density and pulp yield in *Eucalyptus globulus*. *Tree Genet Genomes* 5:291–305. <https://doi.org/10.1007/s11295-008-0174-9>
- Echeverri-Molina D, Santolamazza-Carbone S (2010) Toxicity of synthetic and biological insecticides against adults of the *Eucalyptus* snout-beetle *Gonipterus scutellatus* Gyllenhal (Coleoptera: Curculionidae). *J Pest Sci* 83:297–305. <https://doi.org/10.1007/s10340-010-0298-1>
- González-García S, Hospido A, Moreira MT, Romero J, Feijoo G (2009) Environmental impact assessment of total chlorine free pulp from *Eucalyptus globulus* in Spain. *J Clean Prod* 17:1010–1016. <https://doi.org/10.1016/j.jclepro.2009.02.017>
- Gosney BJ, O’Reilly-Wapstra JM, Forster LG, Barbour RC, Iason GR, Potts BM (2014) Genetic and ontogenetic variation in an endangered tree structures dependent arthropod and fungal communities. *PLoS One* 9:e114132. <https://doi.org/10.1371/journal.pone.0114132>

- Hanks LM, Millar JG, Paine TD, Campbell CD (2000) Classical biological control of the Australian weevil *Gonipterus scutellatus* (Coleoptera: Curculionidae) in California. *Environ Entomol* 29: 369–375. <https://doi.org/10.1093/ee/29.2.369>
- Hill KD, Johnson LAS (1995) Systematic studies in the eucalypts. 7. A revision of the bloodwoods, genus *Corymbia* (Myrtaceae). *Telopea* 6:185–504. <https://doi.org/10.7751/telopea19953017>
- Huerta-Fuentes A, Chiffelle-Gómez I, Serrano-Garzón M, Vázquez-Silva T, Araya-Clericus J (2008) Susceptibility of eucalyptus species to *Gonipterus scutellatus* (Coleoptera: Curculionidae) and electrophoretic profiles of adult maker proteins. *Agrociencia* 42:27–334
- ICNF (2013) IFN6 - Áreas dos usos do solo e das espécies florestais de Portugal Continental em 1995, 2005 e 2010. Instituto da Conservação da Natureza e das Florestas. <http://www.icnf.pt/portal/florestas/ifn/resource/ficheiros/ifn/ifn6-res-prelimv1-1>. Accessed 12 June 2018
- Jactel H, Nicoll BC, Branco M, Gonzalez-Olabarria JR, Grodzki W, Långström B, Moreira F, Netherer S, Orazio C, Piou D, Santos H, Schelhaas MJ, Tojic K, Vode F (2009) The influences of forest stand management on biotic and abiotic risks of damage. *Ann For Sci* 66:1–18. <https://doi.org/10.1051/forest/2009054>
- Lanfranco D, Dungey H (2001) Insect damage in *Eucalyptus*: a review of plantations in Chile. *Austral Ecol* 26:477–481. <https://doi.org/10.1046/j.1442-9993.2001.01131.x>
- Lanfranco D, Peredo H, Ruiz C (2011) Forest insects and pathogens invasions in Chile: case studies in *Pinus* and *Eucalyptus* plantations. In: Pathogens, insects and their associations affecting forestry worldwide. Proceedings of the IUFRO Protection Joint Meeting, Colonia del Sacramento. <https://doi.org/10.13140/2.1.5020.0325>
- Loch AD (2006) Phenology of *Eucalyptus* weevil, *Gonipterus scutellatus* Gyllenhal (Coleoptera: Curculionidae), and chrysomelid beetles in *Eucalyptus globulus* plantations in south-western Australia. *Agric For Entomol* 8:155–165. <https://doi.org/10.1111/j.1461-9563.2006.00294.x>
- Loch AD, Matsuki M (2010) Effects of defoliation by *Eucalyptus* weevil, *Gonipterus scutellatus*, and chrysomelid beetles on growth of *Eucalyptus globulus* in southwestern Australia. *For Ecol Manag* 260:1324–1332. <https://doi.org/10.1016/j.foreco.2010.07.025>
- Mansilla-Vázquez JP, Pérez-Otero R (1996) El defoliador del eucalipto *Gonipterus scutellatus*. *Phytoma España* 81:36–42
- Mapondera T, Burgess T, Matsuki M, Oberprieler R (2012) Identification and molecular phylogenetics of the cryptic species of the *Gonipterus scutellatus* complex (Coleoptera: Curculionidae: Gonipterini). *Aust J Entomol* 51:175–188. <https://doi.org/10.1111/j.1440-6055.2011.00853.x>
- Morse GE, Farrell BD (2005) Ecological and evolutionary diversification of the seed beetle genus *Stator* (Coleoptera: Chrysomelidae: Bruchinae). *Evolution* 59:1315–1333. <https://doi.org/10.1111/j.0014-3820.2005.tb01782.x>
- Newete S, Oberprieler R, Byrne M (2011) The host range of the *Eucalyptus* weevil, *Gonipterus "scutellatus"* Gyllenhal (Coleoptera: Curculionidae), in South Africa. *Ann For Sci* 68: 1005–1013. <https://doi.org/10.1007/s13595-011-0108-9>
- Ohmart CP, Stewart LG, Thomas JR (1985) Effects of food quality, particularly nitrogen concentrations, of *Eucalyptus blakelyi* foliage on the growth of *Paropsis afomaria* larvae (Coleoptera: Chrysomelidae). *Oecologia* 65:543–549. <https://doi.org/10.1007/BF00379670>
- Östrand F, Wallis IR, Davies NW, Matsuki M, Steinbauer MJ (2008) Causes and consequences of host expansion by *Mnesampela privata*. *J Chem Ecol* 34:153–167. <https://doi.org/10.1007/s10886-007-9422-y>
- Palmer WA, Goeden RD (1991) The host range of *Ophraella communis* Lesage (Coleoptera: Chrysomelidae). *Coleopt Bull* 45:115–120
- Reis A, Ferreira L, Tomé M, Araujo C, Branco M (2012) Efficiency of biological control of *Gonipterus platensis* (Coleoptera: Curculionidae) by *Anaphes nitens* (Hymenoptera: Mymaridae) in cold areas of the Iberian Peninsula: implications for defoliation and wood production in *Eucalyptus globulus*. *For Ecol Manag* 270:216–222. <https://doi.org/10.1016/j.foreco.2012.01.038>
- Richardson K, Meakins R (1986) Inter and intra-specific variation in the susceptibility of *Eucalypts* to the snout beetle *Gonipterus scutellatus* Gyll. (Coleoptera: Curculionidae). *S Afr For J* 139:21–31. <https://doi.org/10.1080/00382167.1986.9630053>
- SECF (2010) Situación de los bosques y del sector forestal en España. Informe 2010. Sociedad Española de Ciencias Forestales, Palencia
- Sousa EMR, Ferreira LJC (1996) *Gonipterus scutellatus* Gyll., uma nova praga do eucalipto em Portugal. *Rev Florestal* 9:4–7
- Steinbauer MJ (2002) Oviposition preference and neonate performance of *Mnesampela privata* in relation to heterophyly in *Eucalyptus dunnii* and *E. globulus*. *Agric For Entomol* 4:245–253. <https://doi.org/10.1046/j.1461-9563.2002.00151.x>
- Steinbauer MJ, Matsuki M (2004) Suitability of *Eucalyptus* and *Corymbia* for *Mnesampela privata* (Guenée) (Lepidoptera: Geometridae) larvae. *Agric For Entomol* 6:323–332. <https://doi.org/10.1111/j.1461-9555.2004.00238.x>
- Stone C, Matsuki M, Carnegie A (2003) In: Parsons M (ed) Pest and disease assessment in young eucalypt plantations: field manual for using the Crown Damage Index. National Forest Inventory, Bureau of Rural Sciences, Canberra
- Svanbäck R, Bolnick DI (2007) Intraspecific competition drives increased resource use diversity within a natural population. *Proc R Soc B* 274:839–844. <https://doi.org/10.1098/rspb.2006.0198>
- Tooke F (1955) The *Eucalyptus* snout beetle, *Gonipterus scutellatus* Gyll. A study of its ecology and control by biological means. In: Entomological Memoirs. Department of Agriculture, Pretoria
- Tribe GD (2005) The present status of *Anaphes nitens* (Hymenoptera: Mymaridae), an egg parasitoid of the *Eucalyptus* snout beetle *Gonipterus scutellatus*, in the Western Cape Province of South Africa. *S Afr For J* 203:49–54. <https://doi.org/10.2989/10295920509505218>
- Valente C, Gonçalves CI, Reis A, Branco M (2017) Pre-selection and biological potential of the egg parasitoid *Anaphes inexpectatus* for the control of the *Eucalyptus* snout beetle, *Gonipterus platensis*. *J Pest Sci* 90:811–923. <https://doi.org/10.1007/s10340-017-0839-y>
- Valente C, Gonçalves CI, Monteiro F, Gaspar J, Silva M, Sottomayor M, Paiva MR, Branco M (2018) Economic outcome of classical biological control: a case study on the *Eucalyptus* snout beetle, *Gonipterus platensis*, and the parasitoid *Anaphes nitens*. *Ecol Econ* 149:40–47. <https://doi.org/10.1016/j.ecolecon.2018.03.001>
- Wingfield MJ, Slippers B, Hurley BP, Coutinho T, Wingfield BD, Roux J (2008) Eucalypt pests and diseases: growing threats to plantation productivity. *South For* 70:139–144. <https://doi.org/10.2989/SOUTH.FOR.2008.70.2.9.537>
- Winkler IS, Mitter C (2008) The phylogenetic dimension of insect-plant interactions: a review of recent evidence. In: Tilmon KJ (ed) Specialization, speciation, and radiation: the evolutionary biology of herbivorous insects. University of California Press, Berkeley, pp 240–263
- Zar JH (1996) Biostatistical analysis, 3rd edn. Prentice Hall, Upper Saddle River

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.