Responding positively to plant defences, a candidate key trait for invasion success in the New Zealand grass grub *Costelytra zealandica*

Marie-Caroline Lefort1*, Susan P. Worner1, Michael Rostas1, Jessica Vereijssen2 and Stephane Boyer1,3

1Bio-Protection Research Centre, PO Box 85084, Lincoln University, Lincoln 7647, Canterbury, New Zealand
2Plant & Food Research, Private Bag 4704, Christchurch 8140, New Zealand
3Department of Ecology, Faculty of Agriculture and Life Sciences, PO Box 85084, Lincoln University, Lincoln 7647, Canterbury, New Zealand
*Author for correspondence (Email: Marie-Caroline.Lefort@lincolnuni.ac.nz)

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**Abstract:** Occasionally, exotic plant introductions lead to the emergence of an invasive insect within its native geographical range. Such emergence could be explained by a pre-adaptation of the insect to break through the defences of the new encountered host. We investigated the fitness responses of two New Zealand endemic scarabs (*Costelytra brunneum* and *C. zealandica*) when given a diet of an exotic pasture species, *Trifolium repens*, whose defences were artificially triggered by the phytohormone jasmonic acid. We found differential fitness responses between the two species when they were exposed to a defence-induced diet. We observed a significant weight increase in the invasive species *C. zealandica* when it was fed with treated roots compared with untreated controls, whereas no significant weight increase was observed in the non-invasive *C. brunneum* compared with the control treatments. Our study suggests that *C. zealandica* has a pre-existing ability to tolerate the defence chemicals of its exotic host and, more interestingly, to benefit from them, which may explain why this species has become a serious pest of pasture throughout its native geographical range.

**Keywords:** below-ground induced defences; jasmonic acid; Melolonthinae; native invader; pre-adaptation; Scarabaeidae; *Trifolium repens*

**Introduction**

The reasons why some species become invasive, and other closely related species do not, remain unclear. The established explanations for why a species may become invasive either refer to its intrinsic characteristics, such as its dispersal ability (Rejmanek & Richardson 1996), tolerance limits (Jiménez-Valverde & Lobo 2011) or phenotypic plasticity (Price et al. 2003), or to the various external and/or environmental factors that can contribute to the species becoming invasive (Lefort 2013). The latter include theories about bottom-up and top-down controls (e.g. the novel-weapon hypothesis (Callaway & Ridenour 2004) and the enemy-release hypothesis (Elton 1958). When considering a phytophagous insect reaching the status of a successful invader within its own native range, a number of these established characteristics and theories of invasion success may not be so tenable. For instance, some theories appear to apply solely to plant species (e.g. novel-weapon hypothesis), while others were initially developed for plants and later extended to animal species (e.g. dispersal ability, enemy-release hypothesis). Additionally, the initial introduction stage that defines the process of biological invasion, where a species has to overcome major geographical barriers and/or distances (Valéry et al. 2008), does not necessarily apply to a species transitioning to becoming an invader in its native range following anthropogenic changes on its environment (Lefort 2013). These particular invaders were referred to as ‘native invader’ by Valéry et al. in 2009, and were thereafter defined as such by Simberloff and Rejmanek (2011). Hence, the dispersal ability of a species, defined by McDowall (2011) as the relocation of a number of individuals from a place of origin to another location, is of less interest for the study of native invaders. In a similar way, the concept of enemy release is of less interest because the invasive species is not always likely to completely escape its natural enemies (Lefort 2013).

Recently, Carey et al. (2012) urged scientists to investigate why some native/endemic species transition into invaders within their native range. The comparison of a relatively recent emergent pest and a closely related congeneric non-invasive species offers an ideal opportunity to explore the reasons for such transitions and to propose alternatives to established hypotheses about key traits and theories of invasion success. Such a possibility is given by *Costelytra zealandica* (White) and *C. brunneum* (Broun), two endemic New Zealand scarabs (Scarabaeidae: Melolonthinae) that have developed distinctive relationships with *Trifolium repens* (white clover), a temperate pasture species introduced into New Zealand because of its high economic value as livestock forage and a green manure crop (Gillet & Taylor 2001, Badr et al. 2012). The larval stage of *C. zealandica* quickly became a serious pest of white clover (Radcliffe 1971), while *C. brunneum*, although able to feed on the roots of this plant (Lefort 2013), did not. The present study tests the hypothesis that the invasion success of the New Zealand native pest *C. zealandica*, in contrast with *C. brunneum*, might be partially explained by a pre-adaptation to overcome the induced defences of its main introduced host, the
white clover. Induction of plant defences is a general response to herbivory and involves the elicitation of the jasmonic acid (JA) signalling pathway by chewing-biting insects (Karban & Baldwin 1997). An experiment was carried out where larval populations of *C. zealandica* and its congener, the non-pest *C. brunneum*, were fed with the roots of white clover, in which defence compounds were artificially triggered by the phytohormone JA.

### Methods

#### Insects and plants

Third-instar larvae of *C. zealandica* sampled from exotic pastures mainly composed of ryegrass (*Lolium* spp.) and white clover (*Trifolium repens*) (43°32′17″ S 171°57′16″ E) (Population 1, *n =* 60) and from native tussock grasslands (43°02′10″ S 171°45′40″E) (Population 2, *n =* 60), and *C. brunneum* sampled from native tussock grasslands (43°12′20″ S 171°42′16″ E) (Population 3, *n =* 20) were used for this experiment. Larvae were identified to species level on the basis of their rastor morphology (Lefort et al. 2013), and for a few difficult specimens, a rapid diagnostic confirmation was performed using the DNA recovered from their frass (Lefort et al. 2012).

White clover plants were grown from seeds in 200 ml of sowing mix comprising 60% peat and 40% sterilised pumice stones. Three weeks after sowing, pots were treated by direct soil injection of 5 ml of 10 µM JA (Sigma-Aldrich Chemical Co., St Louis, Missouri), while controls were treated with solvent (0.6 ml of EtOH in 4.4 ml of distilled water). Plants were used 48 h after treatment.

#### Evaluation of defence-induction in white clover roots

Semi-quantitative assessment of trypsin proteinase-inhibitor induction, used as an indicator of the JA pathway activity, was carried out by radial diffusion assay (Jongsma et al. 1993). Briefly, roots of JA-treated clover and controls (*n =* 24) were rinsed twice with distilled water. Sections (200 mg) of root material from individual pots were homogenised in liquid nitrogen and extracted with 175 µl buffer (100 mM Tris HCl and 10 mM CaCl₂). The extract was centrifuged for 2 min at 13 000 g and 4°C. The supernatant (25 µl) was transferred to wells made into a gel containing bovine trypsin as substrate. After overnight incubation and staining, trypsin proteinase-inhibitor activity was detected in all JA-treated clover root replicates while no trypsin proteinase-inhibitor activity, and therefore no defence induction, was detected in the untreated control samples (binomial test, *P* < 0.001). In treated roots, inhibition zone diameters were consistently equivalent to those produced by the standard soybean trypsin proteinase-inhibitor at concentrations between 35 and 70 pM.

#### Results

**Evaluation of defence-induction in white clover roots**

Trypsin-inhibitor activity was detected in all JA-treated clover root replicates while no trypsin proteinase-inhibitor activity, and therefore no defence induction, was detected in the untreated control samples (binomial test, *P* < 0.001). In treated roots, inhibition zone diameters were consistently equivalent to those produced by the standard soybean trypsin proteinase-inhibitor at concentrations between 35 and 70 pM.

**Costelytra spp. response to increased host defences**

After 6 weeks of feeding on JA-treated clover roots, individuals of both *C. zealandica* populations (exotic pasture and native grassland) had significantly larger biomasses than larvae that were fed with untreated roots (Fig. 1a,b). The treatment effect was significant (ANOVA, *F*₁,₅₇ = 5.744, *P* = 0.020 and *F*₁,₅₈ = 4.853, *P* = 0.032, respectively for Populations 1 and 2) with clear differences visible as early as the beginning of the third week (Fig. 1a,b). On the other hand, no treatment effect was detected for the non-pest *C. brunneum* (ANOVA, *F*₁,₈₈ = 0.002, *P* = 0.967) (Fig. 1c).

No significant treatment effect was detected on the survival of the larvae of both species (Fisher’s exact tests, *P* = 0.706, *P* = 0.353 and *P* = 1 for Populations 1, 2 and 3 respectively) (Fig. 2). However, at 10% level of significance, the survival of the pest *C. zealandica* was significantly higher than that of the non-pest *C. brunneum* when fed with either JA-treated clover (Fisher’s exact tests, *P* = 0.052 and *P* = 0.089 for Populations 1 and 3, and for Populations 2 and 3, respectively) or with the untreated clover (Fisher’s exact tests, *P* = 0.085 and *P* = 0.002 for Populations 1 and 3, and for Populations 2 and 3, respectively) (Fig. 2).

#### Discussion

Our study showed that, when induced, *Trifolium repens* has the capacity to increase its below-ground defences through the JA pathway, and both populations of the pest *C. zealandica* showed significantly better growth compared with the control treatment when fed with JA-treated white clover roots. In contrast, no treatment effect was observed in the non-pest species *C. brunneum*. The analyses of the survival rates of *Costelytra* spp. larvae revealed that both species were able to avoid the effect of an increase of their host defences. The fact that *C. brunneum* showed a lower
survival rate than its congener *C. zealandica*, both for treated and untreated clover roots, may reflect a certain sensitivity of the species to sampling manipulation, initial health assessment, experimental conditions or monitoring. This higher sensitivity in *C. brunneum* may also be demonstrated by the rapid weight increase during the first week of treatment, once the living conditions for the larvae were eventually stabilised. The lower survival rates were not ascribed to *C. brunneum* feeding preferences, which were similar when exposed to their native hosts or to white clover (Lefort 2013).

Increased fitness as measured by better growth on JA-treated clover was somewhat unexpected for *C. zealandica*; however, recent studies have reported similar responses of insect species to increased plant defences, whether they were triggered by JA application or by insect wounding. For example, Pierre et al. (2012) reported that JA application on turnip plants resulted in increased pupal size of the cabbage fly, *Delia radicam* (Linnaeus), and Robert et al. (2012a) reported that larvae of the Western corn rootworm, *Diabrotica virgifera* Leconte, gained over 30% more weight on maize plants damaged by conspecifics compared with healthy ones.

The detailed chemical nature of insect-induced plant defences remains largely unknown (Jansen et al. 2008) and little attention has been given to defence induction in roots (Erb et al. 2012; Pierre et al. 2012). It is clear, however, that JA triggers a regulatory cascade that results in the synthesis of numerous primary and secondary plant metabolites (Jansen et al. 2008). Phenolics for example, have been identified among the final products of defence-related genes mediated by the JA defence pathway and are well known for being detrimental to many plant feeding insects (Bernays & Woodhead 1982). Despite this, a significant number of studies have reported counterintuitive positive correlations between high plant phenolic content and high fitness performance in insects (e.g. Bernays & Woodhead 1982; Johnson et al. 2011; Pierre et al. 2012; Robert et al. 2012b). Some suggest that elevated contents of root phenolics rather than being detrimental can have beneficial antioxidant properties for certain phytophagous insects (e.g. Johnson & Felton 2001; Piskorski et al. 2011). Considering that some insects possess the ability to sequester various plant compounds for their own defence (see Jansen et al. (2008) for a review), maybe also, some species, such as *C. zealandica* in this study, have the ability to use certain host defences to their advantage. A possible alternative explanation may rely on the better use of altered primary root metabolite concentrations such as increases in C/N ratio that have been observed in herbivore-damaged clover roots (Murray et al. 1996), rather than on the direct use of secondary host plant compounds by the insect. In this sense, *C. zealandica* may possess an evolutionary advantage that allows it to cope

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**Figure 1.** Average larval weight gain (percentage) for *Costelytra zealandica* (a: from exotic pasture; b: from native grassland) and *C. brunneum* (c: from native grassland) in response to a diet consisting of clover roots treated with jasmonic acid (black) or untreated controls (grey). Vertical bars represent 5% Least Significant Difference (LSD).

**Figure 2.** Kaplan–Meier plot of larval survival of *Costelytra zealandica* and *C. brunneum* during 6 weeks of feeding treatment with untreated clover roots (treatment 1) in green or with jasmonic-acid-treated clover roots (treatment 2) in salmon. Right: final survival after 6 weeks of treatment. All pairwise comparisons were performed using Fisher’s exact tests after 6 weeks of treatment. Survival rates with different letters (a vs b) are significantly different (Fisher’s exact tests, $P < 0.05$).
better with induced host defences than *C. brunneum* and thus allows it to make better use of the high nutrient of these primary metabolites.

While the results here suggest that *C. zealandica* may benefit from host defence induction, it is notable that the population of *C. zealandica* collected from native grasslands also significantly increased weight when fed with the JA-induced ‘new’ host compared with the insects in control treatments. This population, collected from isolated native grassland, is unlikely to have ever been exposed to exotic white clover before the experiment as suggested by the evolution of distinct host-races in this species (Lefort et al. 2014). Such a result suggests that the ability to avoid the putative detrimental effects of plant defences and seemingly benefit from them has not resulted from an ‘arms race’ between the insect and its host, but is a pre-adaptation. The ability to avoid detrimental effects of host defence chemicals and to benefit from them has been consistently reported in particular reference to invasive species. For example, high concentrations of total plant phenols appeared favourable to grub development in the common cockchafer *Melolontha melolontha*, an invasive scarab (L. Sukovata, Polish Forest Research Institute, pers. comm.). Similar observations have been reported in other coleopteran pests, such as the vine weevil *Otiorhynchus sulcatus* (Johnson et al. 2011) and the western corn rootworm *Diabrotica virgifera* (Erb et al. 2012) as well as in invasive moths and grasshoppers (Bernays & Woodhead 1982).

Despite previous reports, to our knowledge, this study is the first to hypothesise a link between insect invasion success and a pre-existing ability to overcome and benefit from the defence metabolites of a new host. We expect the preliminary findings of this study to open new research perspectives in the field of invasion ecology and invite researchers to replicate the experiment described in this paper using other invasive/non-invasive species pairs. The confirmation of such a pre-adaptation as a key trait of invasive phytophagous insects, within but also outside their native range, could have important implications for improving pest risk assessment to prevent and predict the potential economic impact of newly introduced phytophagous insects into new areas.

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