Adaptation of flea beetles to Brassicaceae: host plant associations and geographic distribution of *Psylliodes* Latreille and *Phyllotreta* Chevrolat (Coleoptera, Chrysomelidae)*

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Abstract

The cosmopolitan flea beetle genera *Phyllotreta* and *Psylliodes* (Galerucinae, Alticini) are mainly associated with host plants in the family Brassicaceae and include economically important pests of crucifer crops. In this review, the host plant associations and geographical distributions of known species in these genera are summarised from the literature, and their proposed phylogenetic relationships to other Alticini analysed from published molecular phylogenetic studies of Galerucinae. Almost all *Phyllotreta* species are specialised on Brassicaceae and related plant families in the order Brassicales, whereas *Psylliodes* species are associated with host plants in approximately 24 different plant families, and 50% are specialised to feed on Brassicaceae. The current knowledge on how *Phyllotreta* and *Psylliodes* are adapted to the characteristic chemical defence in Brassicaceae is reviewed. Based on our findings we postulate that *Phyllotreta* and *Psylliodes* colonised Brassicaceae independently from each other.

Keywords

Alticini, chemical plant defence, detoxification, glucosinolates, plant-insect interaction, secondary plant metabolites, sequestration

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Introduction

Plant-feeding insects are often classified as specialists or generalists according to their food plant range. While generalist insect herbivores are able to feed on plants that belong to distantly related plant families, specialist insect herbivores feed selectively on one or a few closely related plant species (Schoonhoven et al. 2005). Many phytophagous insects, including numerous leaf beetle species, have a narrow food plant range (Jolivet and Hawkeswood 1995; Forister et al. 2015), which is at least partially determined by toxic and deterrent plant secondary metabolites. Plants produce more than 200,000 different secondary metabolites, and many of them are involved in defence against herbivores (Mithöfer and Boland 2012). The distribution of secondary metabolites in related plant species often correlates with the food plant range of specialised insect herbivores, which evolved strategies to avoid, tolerate, or detoxify these defence compounds (Heckel 2014). Such adaptations presumably played an important role in the species diversification of plant-feeding insects (Ehrlich and Raven 1964; Futuyma and Agrawal 2009), but the specific molecular mechanisms underlying host plant adaptation, and their role in insect ecology and speciation, are largely unknown.

Several genera in the family Chrysomelidae include species that are specialised to feed on plants in the family Brassicaceae (Table 1). In the subfamily Chrysomelinae, the genera *Colaphellus*, *Entomoscelis*, and *Microtheca* feed primarily on Brassicaceae (Jolivet and Petitpierre 1976b; Nielsen 1988), whereas the genus *Phaedon* is associated with several different plant families, e.g. Asteraceae, Brassicaceae, Scrophulariaceae, and Ranunculaceae (Table 1). In the subfamily Galerucinae, the flea beetle genera *Phyllotreta*, *Psylliodes*, *Leptophyes*, *Caeporis*, and *Hemiglyptus* utilise Brassicaceae as host plants (Furth 1979; Nielsen 1988; Jolivet 1991; Nadine 2010). In addition, many other polyphagous chrysomelid genera feed occasionally on this plant family. However, within Chrysomelidae, the genera *Psylliodes* and *Phyllotreta* comprise the highest number of crucifer specialists.

Glucosinolates are the characteristic secondary metabolites of Brassicaceae and other families in the order Brassicales (Agerbirk and Olsen 2012). Upon herbivory, glucosinolates are hydrolysed by β-thioglucosidase enzymes (myrosinases) to unstable aglucones, which can generate various hydrolysis products such as isothiocyanates, thiocyanates, and nitriles (Wittstock et al. 2016). Isothiocyanates, the most toxic glucosinolate hydrolysis products, are primarily reactive towards thiol (-SH) and amino(-NH$_2$) groups in peptides and proteins (Brown and Hampton 2011). Previous studies revealed that insects developed different strategies to overcome this plant defence (reviewed in Winde and Wittstock (2011) and Jeschke et al. (2016)). For example, *Plutella xylostella* larvae (Lepidoptera, Plutellidae) prevent glucosinolate breakdown by rapidly converting ingested glucosinolates to stable desulfo-glucosinolates (Ratzka et al. 2002), while *Pieris rapae* larvae (Lepidoptera, Pieridae) express a nitrile specifier protein (NSP) in their gut, which promotes the formation of less toxic nitriles instead of isothiocyanates (Wittstock et al. 2004). The evolution of NSP activity in Pierinae butterflies is regarded as an evolutionary key innovation that enabled a host shift from Fabales
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Plants to the glucosinolate-containing Brassicales. As predicted by the coevolutionary 'escape and radiate' hypothesis, speciation rates were higher in the clade that colonised Brassicales plants compared to their sister taxon (Wheat et al. 2007; Edger et al. 2015). In contrast, the host shift of Ceutorhynchini weevils from the plant family Lamiaceae to Brassicaceae was not associated with a speciation rate shift (Letsch et al. 2018).

Glucosinolates and their hydrolysis products are well known to affect the behavior of crucifer-feeding Chrysomelidae (reviewed in Mitchell (1988, 1994), and Nielsen (1988)). Volatile isothiocyanates, for example, attracted high numbers of *Phyllotreta* spp. and *Psylliodes chrysocephala* in field trapping experiments, indicating that isothiocyanates might play a role in host plant localisation (Görnitz 1956; Bartlet et al. 1992; Pivnick et al. 1992; Tóth et al. 2007). Glucosinolates, on the other hand, stimulated feeding of *Phyllotreta* spp., *Ps. chrysocephala*, *Phaedon cochleariae*, and *Entomoscelis americana* in laboratory experiments (Hicks 1974; Mitchell 1978; Nielsen 1978; Bartlet et al. 1994; Reifenrath and Müller 2008). Although these specialists are adapted to the glucosinolate-

### Table 1. Overview of Chrysomelidae genera that are associated with Brassicaceae hosts plants.

| Genus          | Approx. no. of species | Major host plant families | Known species feeding on Brassicaceae | References |
|----------------|------------------------|---------------------------|--------------------------------------|------------|
| **Subfamily Chrysomelinae** |                        |                           |                                      |            |
| *Chrysolina* Motschulsky, 1860 | 450                    | Lamiaceae                 | *C. cavigera, C. colati*              | Jolivet and Petitpierre 1976a, 1976b; Clark et al. 2004; Jurado-Rivera and Petitpierre 2015 |
| *Colaphellus* Weise, 1916 | 15                     | Brassicaceae              | *C. bowringii, C. hoefii, C. sophiae* | Döberl 2010; Gavrilović et al. 2014; Bieńkowski and Orłowa-Bieńkowska 2015; Rheinheimer and Hassler 2018 |
| *Entomoscelis* Chevrotat, 1836 | 14                     | Brassicaceae              | *E. adonidis, E. americana, E. berytensis, E. nigroventris, E. orientalis, E. pilula* | Mohr 1966; Gerber 1994; Ge et al. 2009 |
| *Microtheca* Dejean, 1835 | 15                     | Brassicaceae              | *M. ochroloma, M. picea, M. punctigera, M. semilaevii* | Jolivet 1951; Balsbaugh 1978; Jolivet and Hawkeswood 1995; Ameen 1996; Clark et al. 2004; Menezes et al. 2005; Balusu et al. 2017 |
| **Subfamily Galerucinae, Alticinae** |                        |                           |                                      |            |
| *Carioris* Dejean, 1837 | 1                      | Brassicaceae              | *C. stigmula*                         | Jolivet and Hawkeswood 1995; Cabrera and Rocca 2012; Nadein 2012 |
| *Hemiglyptus* Horn, 1889 | 1                      | Brassicaceae, Hydrophyllaceae | *H. basalis*                         | Clark et al. 2004; Nadein 2012 |
| *Leptophyza* Baly, 1877 | 15                     | Brassicaceae, Cleomaceae  | *L. batesi, L. bordoni, L. littoralis* | Jolivet 1991; Jolivet and Hawkeswood 1995; Bechyně 1997; Flowers and Janzen 1997 |
| *Phyllotreta* Chevrotat, 1836 | 242                   | Brassicaceae              | see Suppl. material 3                | This study; Heikertinger 1943; Furth 1979; Smith 1985; Clark et al. 2004 |
| *Psylliodes* Latreille, 1829 | 207                   | Brassicaceae, Poaceae     | see Suppl. material 1                | This study; Furth 1983; Cox 1998; Clark et al. 2004; Nadein 2010; Baviera and Biondi 2015 |

Table 1.
myrosinase defence system, both glucosinolate levels and myrosinase activity affected herbivory by *Phyllotreta cruciferae* in the field. The highest flea beetle damage was observed on *Brassica rapa* plants with intermediate glucosinolate levels (Siemens and Mitchell-Olds 1996), and *B. rapa* lines selected for high myrosinase activity displayed significantly less feeding damage (ca. 10%) than those with low enzyme activity (Mitchell-Olds et al. 1996). In contrast, studies with *Ps. chrysocephala* did not reveal a correlation between glucosinolate levels and feeding damage (Bartlet et al. 1996; Bartlet et al. 1999).

Here, we provide an overview on the host plants, diet breadth, and geographic distribution of known *Phyllotreta* and *Psylliodes* species, as well as their proposed relationships to other genera of Alticini. Diet breadth was classified according to Biondi (1996). Species feeding on one or two closely related botanical genera are considered as monophagous, species feeding on more plant genera of one or two closely related families are defined as oligophagous, and species feeding on many distantly related plant species are considered as polyphagous. For species with limited information on food plants, we did not specify the diet breadth. Data on the geographical distribution of the Palearctic *Psylliodes* and *Phyllotreta* species was primarily obtained from Döberl (2010) and is described according to Löbl and Smetana (2010). The zoogeographical regions are abbreviated as follows: Afrotropical Region (AFR), Australian Region (AUR), Nearctic Region (NAR), Neotropical Region (NTR), Oriental Region (ORR), Palearctic Region (PAR). In the second part of this review, we summarise the knowledge on the adaptations of *Phyllotreta* and *Psylliodes* spp. to the glucosinolate-myrosinase defence system and other defences in their host plants.

**Host plant associations of *Psylliodes* and *Phyllotreta* flea beetles**

The genus *Psylliodes* Latreille, 1829 comprises over 200 species (Suppl. material 1). Adult *Psylliodes* beetles are distinguished from other flea beetle genera based on their 10-segmented antennae and tarsi inserted pre-apically on the metatibia of the hind legs. Most other Alticini genera have 11-segmented antennae except for *Psylliodes*, *Decaria*, and *Monotalla* with ten segments and *Nonarthra* with nine segments (Konstantinov and Vandenbergh 1996; Nadein and Bezděk 2014). The genus comprises five subgenera: *Psylliodes* s. str. (194 species), *Semicnema* Weise (5 species), *Eupus* Wollaston (5 species), *Minicnema* Nadein (2 species) and *Psyllobactra* Lopatin (1 species) (Nadein 2007a, 2010). A subdivision of the subgenus *Psylliodes* s. str. based on morphological features was proposed by Leonardi (1970) and Nadein (2006, 2007a, 2007b) (Suppl. material 2).

According to the literature, host plants of 107 *Psylliodes* species have been reported, and these belong to 24 plant families (Suppl. material 1). Most *Psylliodes* species have a restricted host plant range (35% are monophagous and 51% are oligophagous), and only 14% are polyphagous. For instance, *Psylliodes toelgi* feeds only on *Biscutella laevigata* (Brassicaceae), whereas *Psylliodes luteola* has been recorded on Poaceae, Fagaceae, Salicaceae, Ulmaceae, and Solanaceae.
Of all *Psylliodes* species with known host plants, 50% are specialised on Brassicaceae, followed by 13% feeding on Poaceae, 10% on Solanaceae and 10% on Fagaceae (Fig. 1A). Previous surveys of host plant associations of *Psylliodes* spp. focused on specific countries or regions and thus included a much smaller total number of *Psylliodes* species (Furth 1983; Cox 1998; Döberl 2010; Baviera and Biondi 2015). Interestingly, host plant use often correlates with the proposed *Psylliodes* s. str. species groups, which indicates that presumably closely related *Psylliodes* species feed on closely related host plants (Suppl. material 2). For example, *Psylliodes* species in the *chrysocephala* and *pyritosa* groups are specialised to feed on Brassicaceae, while species in the *luteola* group are mainly associated with Fagaceae.

The genus *Phyllotreta* Chevrolat, 1836 comprises about 242 species and host plant information is available for 117 species (Suppl. material 3). Most *Phyllotreta* species are specialised on glucosinolate-containing plants in the order Brassicales (Fig. 1B). An analysis of the diet breadth of *Phyllotreta* species revealed that 31% are monophagous, 64% are oligophagous, and 5% are polyphagous. In *Phyllotreta*, 63% are specialised on Brassicaceae, whereas 18% feed on plants in more than one family in the order Brassicales (Fig. 1B). Very few *Phyllotreta* species feed on plant families, which do not contain glucosinolates, for instance, *Phyllotreta cruralis* is specialised on Amaranthaceae.

Several *Psylliodes* and *Phyllotreta* species are of economic importance. The cabbage stem flea beetle, *Ps. chrysocephala* is a serious pest of winter oilseed rape in Northern Europe (Zimmer et al. 2014), whereas *Phyllotreta striolata* and *Ph. cruciferae* are oilseed rape pests in Canada where their damage causes losses of tens of millions of US dollars annually (Lamb 1989; Hill 2008; Knodel 2017). On the other hand, the Palearctic species *Psylliodes chalcomera* (feeding on Asteraceae) was introduced to North America in 1997 as a control agent for the invasive weed *Carduus nutans* (musk thistle), but it likely did not establish in the Nearctic region (Antonini et al. 2008).

**Geographic distribution of *Psylliodes* and *Phyllotreta* flea beetles**

The genus *Psylliodes* has a worldwide distribution (Biondi and D’Alessandro 2018). The highest number of species occurs in the Palearctic region (160 species, 145 endemic species), followed by the Oriental region (27 species, 19 endemic species), the Nearctic region (13 species, 4 endemic species), the Afrotropical region (13 species, 9 endemic species), the Neotropical region (8 species, 4 endemic species), and the Australian region (8 species, 7 endemic species; Suppl. material 1). A graphical overview of the species distribution is shown in Figure 2A; the host plant associations of all species and endemic species in each zoogeographical region are shown in Figure 2B. Some species are wide-spread in more than one zoogeographical region such as *Ps. brettinghami* (feeding on Solanaceae), which is found in the Australian, Oriental, and Palearctic regions, while others are strictly endemic to very limited areas, e.g. *Ps. tarsata*, which is only found on Madeira (Portugal). *Psylliodes* species that are endemic to the Palearctic region account
Figure 1. Host plant associations of the genera *Psylliodes* (A) and *Phyllotreta* (B). The host plants of 107 *Psylliodes* species and 117 *Phyllotreta* species have been reported in the literature. The numbers of species which feed on plants in one plant family (monophagous and oligophagous), and the number of polyphagous species are given as percentages. 18% of the *Phyllotreta* species feed on more than one family in the order Brassicales (Brassic., Brassicaceae; Cappar., Capparaceae; Cleom., Cleomaceae; Resed., Resedaceae; Tropaeol., Tropaeolaceae). For detailed information, refer to Suppl. material 1 (*Psylliodes*) and 3 (*Phyllotreta*).

for 83% of those associated with Brassicaceae. All other Brassicaceae-feeding species are found in other zoogeographical regions except for Australia (Fig. 2B).

The geographic distribution of the genus *Phyllotreta* shows the highest number of species in the Palearctic region (137 species, 118 endemic species) followed by the Afrotropical region (49 species, 39 endemic species), the Nearctic region (49 species, 40 endemic species), the Oriental region (25 species, 18 endemic species), the Neotropical Region (5 species, 3 endemic species), and the Australian Region (4 species,
3 endemic species; Suppl. material 3). The species distribution is shown in Figure 3A, and the host plant associations of all species and endemic species in each zoogeographical region are shown in Figure 3B. In general, a high percentage of endemic *Phyllotreta* species is found in all geographical regions (≥ 60%) with highest values in the Palearctic, Afrotropical, and Nearctic regions (≥ 80%). In some areas, especially in the Nearctic region, several species of *Phyllotreta* are not native and have been introduced from other regions (Milliron 1953; Smith 1985). Most species feeding on Brassicaceae are found in the Palearctic and Nearctic regions. The host plants of a large proportion of the species endemic to the Afrotropical, Australian, and Neotropical regions are unknown (Fig. 3B; Suppl. material 3).

**Phylogenetic relationships of *Psylliodes* and *Phyllotreta* to other Alticini**

The most comprehensive phylogenetic analyses of the subfamily Galerucinae *sensu lato* are those of Ge et al. (2011, 2012) and Nie et al. (2018), which included about 80 and 70 genera of Alticini (including problematic genera), respectively. Ge et al. (2011, 2012) used two mitochondrial (16S rRNA and cytochrome oxidase (cox) 1) and two nuclear genes (18S and 28S rRNA) to infer phylogenetic relationships, while Nie et al. (2018) used the mitochondrial genome and nuclear rRNA genes. In these analyses, *Psylliodes* and *Phyllotreta* were never retrieved as sister genera, but instead clustered in distinct clades with other Alticini as summarised in Table 2. All three studies suggest a close phylogenetic relationship of *Psylliodes* to *Chaetocnema* and *Crepidodera* (see Table 2 for Bayesian posterior probability values and/or Maximum Likelihood bootstrap support values). Surprisingly, two different *Crepidodera* species included in
the analysis of Nie et al. (2018) were not monophyletic. *Crepidodera pluta* clustered in the *Chaetocnema* group with *Psylliodes*, while the second *Crepidodera* sp. clustered together with two *Phyllotreta* species in a distant clade. However, the proposed relationships of *Phyllotreta* to other Alticini differ among the studies, and are usually less supported than those suggested for *Psylliodes*. None of the genera with proposed close phylogenetic relationships to *Phyllotreta* and *Psylliodes* are associated with Brassicaceae plants (Table 2).

**Adaptations of crucifer-feeding flea beetles to chemical plant defences**

An unexpected observation revealed that *Ph. striolata* adults emit low amounts of toxic isothiocyanates, which are derived from glucosinolates that are stored at high concentrations of up to 50 µmol/g fresh weight (ca. 2% of the body weight) in adults (Beran 2011; Beran et al. 2014). When adults were transferred to different crucifer species, they selectively accumulated mainly aliphatic glucosinolates from their food plants, e.g. allyl glucosinolate from *Brassica juncea*, and 4-methylsulfinylbutyl (4MSOB) glucosinolate from *Arabidopsis thaliana*. In contrast, adults sequestered only low amounts of the benzenic 4-hydroxybenzyl glucosinolate from *Sinapis alba*. The glucosinolate accumulation pattern depended both on glucosinolate structure and on the host plant background, suggesting that the plants’ glucosinolate composition affects sequestration in *Ph. striolata*. The ability to accumulate high glucosinolate amounts demonstrates that *Ph. striolata* can at least partially prevent activation of ingested glucosinolates.
Table 2. Phylogenetic relationships of *Psylliodes* and *Phyllotreta* to other Alticini genera.

| Study                      | Psylliodes                        | Phyllotreta                               |
|----------------------------|-----------------------------------|------------------------------------------|
| Ge et al. (2011)           | Sister genus: *Chaetocnema* (Poaceae)¹ | Sister genus: *Batophila* (Rosaceae)     |
|                            | Phylogenetic support (B/ML): 0.84/67 | Phylogenetic support (B/ML): 0.79/50     |
|                            | Clade: *Crepidodera* (Salicaceae), | Clade: *Lipromela* (unknown),            |
|                            | *Epitrix* (Solanaeae)              | *Sphrea* (Euphorbiaceae),               |
|                            | Phylogenetic support (B/ML): 0.52/50 | *Altica* (Onagraceae, Lythraceae),       |
|                            | Taxonomic group: Unspecified        | *Macrohaltica* (Gunneraceae)            |
|                            |                                   | Phylogenetic support (B/ML): 0.98/50    |
| Ge et al. (2012)           | **Bayesian and Maximum-Likelihood phylogenies** |                                      |
|                            | Sister genus: *Chaetocnema* (Poaceae) | Sister genus: *Epitrix* (Solanaceae)    |
|                            | Phylogenetic support (B/ML): 0.95/67 | Phylogenetic support (B): 0.95          |
|                            | Clade: *Crepidodera* (Salicaceae), | Clade: *Diphaltica* (Aquifoliaceae),    |
|                            | *Epitrix* (Solanaceae), *Sphrea* (Euphorbiaceae), | *Agsicles* (Amaranthaceae),          |
|                            | *Altica* (Onagraceae, Lythraceae),  | *Disonycha* (Amaranthaceae)             |
|                            | *Macrohaltica* (Gunneraceae)       | Phylogenetic support (B): 0.81          |
|                            | Phylogenetic support (B/ML): 0.89/50 | Maximum-Likelihood phylogeny            |
|                            | Taxonomic group: *Chaetocnema*      |                                         |
| Nie et al. (2018)          | Sister genus: *Chaetocnema* (Poaceae), | Sister genus and clade:                 |
|                            | *Epitrix* (Solanaceae)              | *Crepidodera* (Salicaceae)              |
|                            | Phylogenetic support (B): 0.48      | Phylogenetic support (B): 0.83          |
|                            | Clade: *Crepidodera* (Salicaceae),  |                                          |
|                            | *Xuthea* (Urticaceae)               |                                          |
|                            | Phylogenetic support (B): 0.89      |                                          |
|                            | Taxonomic group: *Chaetocnema*      |                                          |

¹The major host-plant family for each genus according to Jolivet and Hawkeswood (1995) is given in parentheses.

ML – Maximum Likelihood bootstrap value; B – Bayesian posterior probability

However, quantitative feeding studies, for instance with radiolabeled glucosinolates, are needed to determine to which degree ingested glucosinolates are sequestered intact.

To activate sequestered glucosinolates, *Ph. striolata* possesses an insect myrosinase with high activity towards aliphatic glucosinolates, which evolved from insect β-O-glucosidases (Figure 4; Beran et al. 2014). To investigate how *Ph. striolata* activate sequestered glucosinolates and prevent autointoxication, dissected tissues from adults were analysed for the presence of glucosinolates and myrosinase activity, respectively. Interestingly, both glucosinolates and myrosinase were mainly localised in the hemolymph and elytra (Beran and Ahn, unpublished), but whether both components are stored separately in hemoplasma and hemocytes as previously reported for cyanogenic glycosides and the cyanogenic β-glucosidase in *Zygaena filipendulae* larvae (Lepidoptera, Zygaenidae; Pentzold et al. 2017), is not yet known.
In the genus *Psylliodes*, the cabbage stem flea beetle, *Ps. chrysocephala*, selectively sequesters glucosinolates as well, but compared to *Ph. striolata*, glucosinolate concentrations are much lower (ca. 4 µmol/g fresh weight; Beran et al. 2018). Although glucosinolates are present in all life stages of *Ps. chrysocephala*, a defensive function is unlikely, as neither larvae nor adults possess endogenous myrosinase activity (Beran et al. 2018). An analysis of the metabolic fate of ingested 4MSOB glucosinolate in *Ps. chrysocephala* adults revealed that adults utilise at least three strategies to prevent isothiocyanate formation and toxicity. *Ps. chrysocephala* sequester intact glucosinolates, detoxify glucosinolates by desulfation, and detoxify dietary isothiocyanates by conjugation to glutathione. The isothiocyanate-glutathione conjugate is metabolised via the conserved mercapturic acid pathway to three different cyclic cysteine conjugates, which are excreted. These three strategies accounted for the metabolic fate of 18.5%, 8%, and 17% of the total ingested glucosinolates, respectively. The amounts of other glucosinolate breakdown products (4MSOB-isothiocyanate, -cyanide, -amine, and -acetamide) corresponded to 17.5% of the total ingested glucosinolate (Figure 4; for details, refer to Beran et al. 2018). However, the metabolic fate of about 39% of the total ingested glucosinolate remained unknown in this study.

The detoxification of isothiocyanates in *Ps. chrysocephala* comes at the expense of the amino acid cysteine. Therefore, interference with protein digestion, for instance by plant proteinase inhibitors or other digestibility reducers, might affect the detoxification capacity for isothiocyanates by limiting the availability of cysteine for glutathione biosynthesis. Interestingly, there is evidence that *Ps. chrysocephala* can compensate for the ingestion of plant proteinase inhibitors. *Ps. chrysocephala* larvae reared on a transgenic *Brassica napus* line that overexpressed the cysteine proteinase inhibitor oryzacystatin I showed doubled proteolytic activity and were heavier than those reared on the corresponding *B. napus* wild type (Girard et al. 1998). This unexpected result shows that *Ps. chrysocephala* is not only adapted to glucosinolates but also to plant proteinase inhibitors.

Specialist chrysomelids are well known for discriminating between crucifer species (Feeny et al. 1970; Nielsen 1977; Bartlet and Williams 1991; Pachagounder and Lamb 1998; Pachagounder et al. 1998), but the factors that determine host suitability and preference are often not understood. Although leaf beetles recognise and differentially respond to individual glucosinolates when offered in isolation, there is little evidence that host plant preference relies on specific glucosinolate profiles (Nielsen 1988). Instead, the presence of other toxic secondary metabolites such as cucurbitacins and cardenolides was shown to affect host suitability for *Phyllotreta* spp. and *Phaedon cochleariae* (abbreviated *Phaedon*) (Nielsen 1978). Toxic cucurbitacins B, E, and I present in *Iberis* spp. deterred feeding of *Phyllotreta nemorum* but not of *Phaedon*, an effect that correlated with their feeding behavior towards *Iberis* plants. On the other hand, *Phaedon*, *Phyllotreta undulata*, and *Phyllotreta tetragaster* did not feed on cardenolide-containing *Cheiranthus* and *Erysimum* spp., which are accepted as food plants by *Ph. nemorum* (Nielsen 1978).
Figure 4. Metabolism of glucosinolates in *Psylliodes chrysocephala* and *Phyllotreta striolata*. Upon herbivory, glucosinolates are usually hydrolysed by the plant enzyme myrosinase to an unstable aglucone, which spontaneously rearranges to a toxic isothiocyanate. In the presence of plant specifier proteins, other hydrolysis products such as thiocyanates and nitriles are formed. Both flea beetle species sequester glucosinolates in their bodies, suggesting that not all glucosinolates are hydrolysed in feeding-damaged plant tissue. Sequestered glucosinolates may be activated for defensive purposes by an insect myrosinase in *Ph. striolata*, but not in *Ps. chrysocephala*. In addition, *Ps. chrysocephala* partially detoxifies glucosinolates by desulfation, whereas no glucosinolate sulfatase activity was found in *Ph. striolata*. According to a quantitative feeding study performed with *Ps. chrysocephala*, most ingested glucosinolates are activated, and isothiocyanates are detoxified by conjugation to glutathione. The isothiocyanate-glutathione conjugate is metabolized via the mercapturic acid pathway to several cyclic metabolites in *Ps. chrysocephala* adults (Beran et al. 2018). Examples of three structurally different glucosinolate side-chains are shown in the box. Beetle photos: Anna Schroll.

The oligophagous species *Ph. nemorum* is used as a model to study the genetic basis of host plant adaptation. The common wild crucifer, *Barbarea vulgaris* ssp. *arcuata* (abbreviated *B. vulgaris*), is an atypical host plant for *Ph. nemorum*. However, the discovery of two different flea beetle populations using *B. vulgaris* as natural host plant suggests that *Ph. nemorum* is extending its host plant range to include *B. vulgaris* in Denmark (Nielsen 1996; de Jong et al. 2000). There are two distinct types of *B.
The two *B. vulgaris* types differ not only morphologically but also regarding their chemical defences, i.e. glucosinolates and saponins. Feeding assays showed that susceptible *Ph. nemorum* larvae started to mine into the leaves of the G-type, but then either left and refused to feed or died in the mine, showing that the G-type is toxic for them (Nielsen 1997a, 1997b). Resistance of the G-type to *Ph. nemorum* is linked to the presence of the triterpenoid saponins hederagenin cellobioside, oleanolic acid cellobioside, gypsogenin cellobioside, and 4-epihederagenin cellobioside, and not to distinct glucosinolate profiles (Agerbirk et al. 2001; Kuzina et al. 2009; Nielsen et al. 2010). The toxicity of saponins is at least partially due to their interactions with cell membranes, which can cause cell death (Augustin et al. 2011). The activity of isolated hederagenin cellobioside and oleanolic acid cellobioside was tested separately in no-choice feeding assays with *Ph. nemorum* adults from five different near-isogenic lines (Nielsen et al. 2010). In these experiments, hederagenin cellobioside had a much stronger negative effect on adult feeding than oleanolic acid cellobioside, whereas the corresponding aglycones of both saponins were not active. An even stronger negative effect on some *Ph. nemorum* lines was observed for α-hederin, a saponin which is not present in *B. vulgaris*, and only differs from hederagenin cellobioside in its glycosylation pattern (Nielsen et al. 2010). These results show that aglycone structure as well as glycosylation pattern affect the biological activity of saponins towards *Ph. nemorum*.

Although the saponin-based defence of *B. vulgaris* is a dead-end for most *Ph. nemorum* genotypes, resistant individuals that performed well on the G-type were found at varying frequencies in all sampled populations (Nielsen and de Jong 2005; Nielsen 2012; Vermeer et al. 2012). The ability to use the G-type as a host plant clearly shows that resistant individuals can tolerate or detoxify saponins by an unknown mechanism. In genetic analyses, Nielsen and de Jong identified the presence of dominant resistance-conferring genes (R-genes) in all resistant individuals, but divergent modes of inheritance of these R-genes (autosomal and sex-linked) between populations (Nielsen 1997a; de Jong et al. 2000; de Jong and Nielsen 2002; Nielsen 2012). For example, in the resistant population from Ejby (Denmark), two major R-genes were linked to the sex chromosomes with additional autosomal R-genes. In a resistant population from Kværkeby (Denmark), most individuals were homozygous for a single autosomal R-gene (Nielsen 1997a; de Jong et al. 2000). In crossing experiments with resistant males from a Swiss population, an autosomal R-gene was inherited only to female offspring due to non-random segregation. The most likely explanation for this non-random segregation of the autosomal R-gene together with the X chromosome is the fusion of an autosome carrying the susceptible allele to the Y-chromosome in Swiss males (Nielsen 2012). When this R-gene was introduced into the genetic background
of the susceptible *Ph. nemorum* line, it showed a normal Mendelian inheritance pattern (Nielsen 2012). These results strongly suggest that the genetic architecture of *Ph. nemorum* males differs among flea beetle populations, and that this polymorphism affects the inheritance of R-genes that enable the offspring to use the otherwise toxic *B. vulgaris* G-type as a host plant. Interestingly, attempts to generate *Ph. nemorum* lines that are homozygous for an autosomal R-gene resulted in very low survival rated of the homozygous larvae (de Jong and Nielsen 2000; Breuker et al. 2007). This observation was surprising as the homozygous resistant genotype was common at least in the *B. vulgaris*-feeding population from Kværkeby, which suggests that co-adapted genes present in the field population counteract the fitness cost of R-genes (de Jong et al. 2000; de Jong and Nielsen 2002). The genetic diversity and population structure of *Ph. nemorum* makes this species an ideal model to study the genetic basis of host range expansion in an oligophagous herbivore.

**Conclusions and future directions**

The flea beetle genera *Psylliodes* and *Phyllotreta* are closely associated with glucosinolate-containing plants mainly in the family Brassicaceae. Nevertheless, they differ remarkably in their overall host plant use and their adaptations to glucosinolates, the characteristic defence metabolites in Brassicaceae. While *Ph. striolata* can utilise sequestered glucosinolates for its defence against predators, *Ps. chrysocephala* apparently does not possess endogenous myrosinase activity and accumulates much lower amounts of glucosinolates compared to *Ph. striolata*. In addition, both species differ regarding their ability to detoxify glucosinolates by desulfation (Beran et al. 2014, 2018).

Despite this progress, our knowledge on the adaptations of *Phyllotreta* and *Psylliodes* to the glucosinolate-myrosinase defence is far from complete. It is unknown, for instance, whether *Phyllotreta* rapidly sequester glucosinolates to prevent their breakdown to toxic isothiocyanates, and whether *Phyllotreta* gain protection from natural enemies by activating sequestered glucosinolates using their own myrosinase. In *Ps. chrysocephala*, the importance of the various detoxification strategies and their evolution needs to be investigated. To this end, a robust phylogenetic tree of the genus and comparative studies on how other *Psylliodes* species are processing dietary glucosinolates are necessary.

A future goal is to place adaptations of *Phyllotreta* and *Psylliodes* to their glucosinolate-containing host plants into a broader evolutionary context. While recent phylogenetic studies support the hypothesis that both genera adapted independently to Brassicaceae, their relationships to other genera of Alticini remain largely unresolved (Ge et al. 2011; Ge et al. 2012; Nie et al. 2018). At this background, a comprehensive and well-resolved phylogenetic tree of the tribe Alticini will enable studies on interactions with plants in general and adaptations to plant chemical defences, and how they contributed to the evolutionary success of this megadiverse lineage.
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**Supplementary material I**

**Table S1**

Authors: Matilda W. Gikonyo, Maurizio Biondi, Franziska Beran

Data type: (Species, host plants, diet breadth, geographic distribution)

Explanation note: List of Psylliodes species according to their subgenera, including their food plants, diet breadth and geographical distribution.

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Link: https://doi.org/10.3897/zookeys.856.33724.suppl1
Supplementary material 2

Table S2
Authors: Matilda W. Gikonyo, Maurizio Biondi, Franziska Beran
Data type: (Species, host plant families)
Explanation note: Species groups of *Psylliodes* s. str. and their associated host plant families.
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Supplementary material 3

Table S3
Authors: Matilda W. Gikonyo, Maurizio Biondi, Franziska Beran
Data type: (Species, host plants, diet breadth, geographic distribution)
Explanation note: List of *Phyllotreta* species including their food plants, diet breadth and geographical distribution.
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