

DOI: 10.1111/eea.12967

SPECIAL ISSUE: 6TH INTERNATIONAL ENTOMOPHAGOUS INSECTS CONFERENCE

Biological control of invasive stink bugs: review of global state and future prospects

Eric Conti¹* D, Gonzalo Avila^{2,3} D, Barbara Barratt^{3,4} D, Fernanda Cingolani⁵ D, Stefano Colazza⁶ D, Salvatore Guarino⁷ D, Kim Hoelmer⁸ D, Raul Alberto Laumann⁹ D, Lara Maistrello¹⁰ D, Guillaume Martel^{11,12} D, Ezio Peri⁶ D, Cesar Rodriguez-Saona¹³ D, Gabriele Rondoni¹ D, Michael Rostás¹⁴ D, Pio Federico Roversi¹⁵ D, René F.H. Sforza¹¹ D, Luciana Tavella¹⁶ D & Eric Wajnberg^{17,18} D

¹Department of Agricultural, Food and Environmental Sciences, University of Perugia, Borgo XX Giugno, Perugia 06121, Italy, ²The New Zealand Institute for Plant & Food Research Limited, Private Bag 92169, Auckland 1142, New Zealand, ³Better Border Biosecurity, Christchurch New Zealand, ⁴AgResearch, Invermay Research Centre, Private Bag 50034, Mosgiel 9053, New Zealand, ⁵CEPAVE (CONICET-UNLP), Boulevard 120 e/ 60 y 64 s/n, La Plata 1900, Argentina, ⁶Dipartimento di Scienze Agrarie, Alimentari e Forestali, Università degli Studi di Palermo, Viale delle Scienze, edificio 5, Palermo 90128, Italy, Institute of Biosciences and Bioresources (IBBR), National Research Council of Italy (CNR), Corso Calatafimi 414, Palermo 90129, Italy, ⁸USDA–ARS, Beneficial Insects Introduction Research Unit, 501 S. Chapel St, Newark DE, USA, ⁹Embrapa Recursos Genéticos e Biotecnologia. PqEB Avda W5 Norte (Final), CEP 70770-917, Brasília DF, Brazil, ¹⁰Dipartimento di Scienze della Vita, Centro BIOGEST-SITEIA, Università di Modena e Reggio Emilia, Via G. Amendola 2, Reggio-Emilia 42122, Italy, 11 USDA-ARS-European Biological Control Laboratory, Campus International de Baillarguet, 810, avenue du Campus Agropolis, Montferrier-sur-Lez 34980, France, ¹²Montpellier SupAgro, Place Pierre Viala, 2, Montpellier 34000, France, ¹³Department of Entomology, Rutgers University, New Brunswick NJ, 08901, USA, ¹⁴Agricultural Entomology, Department of Crop Sciences, University of Göttingen, Grisebachstr. 6, Göttingen 37077, Germany, ¹⁵CREA Consiglio per la Ricerca in Agricoltura e l'analisi dell'economia agraria, Via di Lanciola, 12/a, Cascine del Riccio, Firenze 50125, Italy, ¹⁶Dipartimento di Scienze Agrarie, Forestali e Alimentari (DISAFA), Università degli Studi di Torino, Largo Paolo Braccini 2, Grugliasco TO, 10095, Italy, ¹⁷INRA, 400 Route des Chappes, BP 167, Sophia Antipolis Cedex 06903, France, and ¹⁸INRIA, Sophia Antipolis, Project Hephaistos, 2004 Route des Lucioles, BP 93, Sophia AntipolisCedex 06902, France

Accepted: 11 June 2020

Key words: biocontrol, chemical ecology, Hemiptera, invasive species, landscape management, preemptive classical biological control, semiochemicals, parasitoid, Pentatomidae, risk assessment, Tachinidae, Scelionidae

Abstract

Invasive stink bugs (Hemiptera: Pentatomidae) are responsible for high economic losses to agriculture on a global scale. The most important species, dating from recent to old invasions, include Bagrada hilaris (Burmeister), Halyomorpha halys (Stål), Piezodorus guildinii (Westwood), Nezara viridula (L.), and Murgantia histrionica (Hahn). Bagrada hilaris, H. halys, and N. viridula are now almost globally distributed. Biological control of these pests faces a complex set of challenges that must be addressed to maintain pest populations below the economic injury level. Several case studies of classical and conservation biological control of invasive stink bugs are reported here. The most common parasitoids in their geographical area of origin are egg parasitoids (Hymenoptera: Scelionidae, Encyrtidae, and Eupelmidae). Additionally, native parasitoids of adult stink bugs (Diptera: Tachinidae) have in some cases adapted to the novel hosts in the invaded area and native predators are known to prey on the various instars. Improving the efficacy of biocontrol agents is possible through conservation biological control techniques and exploitation of their chemical ecology. Moreover, integration of biological control with other techniques, such as behavioural manipulation of adult stink bugs and plant resistance, may be a sustainable pest control method within organic farming and integrated pest management programs. However, additional field studies are needed to verify the efficacy of these novel methods and transfer them from research to application.

^{*}Correspondence: Eric Conti, Department of Agricultural, Food and Environmental Sciences, University of Perugia, Borgo XX Giugno, 06121, Perugia, Italy. E-mail: eric.conti@unipg.it

Introduction

Invasive insect herbivores are responsible for an annual loss of at least 70 billion USD globally (Bradshaw et al., 2016), with the biggest agricultural producers, USA and China, experiencing the highest costs and also representing the main potential sources of invasive pests (Paini et al., 2016). The economic loss caused by invasive insects is growing, mostly due to market globalization and climate change (Bradshaw et al., 2016). Herbivorous stink bugs (Hemiptera: Pentatomidae) serve as a good example of this, as many species are agricultural pests in their native range as well as serious invasive pests with high economic impact (Panizzi et al., 2000; McPherson, 2018). Stink bug damage is due to the feeding punctures of adults and nymphs on plant tissues, especially those of fruits and seeds and sometimes leaves and stems, resulting in large reductions in crop yield and/or quality (Panizzi et al., 2000; McPherson, 2018). Additionally, several species transmit plant pathogens (Mitchell et al., 2018).

At least two polyphagous pentatomid species, the brown marmorated stink bug, Halyomorpha halys (Stål) (Hamilton et al., 2018), and the southern green stink bug, Nezara viridula (L.) (Esquivel et al., 2018), are globally important pests. Halyomorpha halys is a temperate/subtropical species native to eastern Asia that has established in North America and Europe, where it damages many crops (reviewed by Leskey & Nielsen, 2018), has recently established in Chile (Faúndez & Rider, 2017), and has regularly been intercepted at New Zealand's border in recent years (Avila & Charles, 2018; Charles et al., 2019). Unlike H. halys, the native geographical range of N. viridula (Africa, southwest Asia, or Mediterranean region) is not yet clearly defined, but this species has spread into tropical, subtropical, and warm-temperate zones of five continents and might expand further (Panizzi & Slansky, 1985; Todd, 1989; Panizzi & Lucini, 2016; Esquivel et al., 2018).

The most recent widespread invasive pentatomid species, the painted bug or Bagrada bug, *Bagrada hilaris* (Burmeister), which has African and Asian origins, now has an almost worldwide distribution and has recently become a serious concern on brassicaceous crops in the USA (Bundy et al., 2018b). To date, two additional stink bugs, *Murgantia histrionica* (Hahn) and *Piezodorus guildinii* (Westwood), are less widely distributed than *B. hilaris*, having expanded their ranges only within the American continent. Historically, *M. histrionica*, native to Central America and Mexico, could be considered the first recorded invasive stink bug, having invaded the southern part of the USA during the 19th century where it is still causing economic losses on brassicaceous crops (McPherson et al., 2018). *Piezodorus guildinii*, originally described

from the Caribbean, is now a major pest of soybean and other Fabaceae in many parts of the American continent (Bundy et al., 2018a).

Stink bug expansion into new areas, especially when transported over long distances by global trade and tourism, can often be attributed to the overwintering behaviour of adults (Panizzi et al., 2000; Musolin et al., 2018). In autumn, several species aggregate in buildings and other artificial shelters to hibernate. Overwintering *H. halys* adults have been transported while hidden inside many types of material (Maistrello et al., 2018; Nixon et al., 2019). This has affected international trade due to the mandatory phytosanitary importation measures required by some countries (e.g., New Zealand and Australia). These regulations impose treatments on any type of commodities imported from countries where *H. halys* is confirmed as present (MPI, 2020).

Following the introduction of exotic species into new areas, native natural enemies sometimes adapt to the noncoevolved species. Fortuitous biological control of invasive stink bugs by native parasitoids and predators has been observed, but in most cases native parasitoids are not considered effective for biological control, especially when compared with coevolved species from the native area (McPherson, 2018). Because of a lack of effective natural enemies, combined with stink bug biology, locally favourable ecological conditions, and their capacity to cause direct damage to marketable produce, the impact of stink bugs in invaded agroecosystems is often significant, both in terms of crop loss and by disrupting established integrated pest management (IPM) practices (Panizzi et al., 2000; McPherson, 2018). It is therefore vital to develop new sustainable control methods that can be included within revised IPM strategies. In this paper, we review a century of biological control programs for invasive stink bugs and discuss the global biocontrol strategies developed against them through six case studies. These strategies include classical, augmentative, and conservation biological control, as well as an innovative pre-emptive classical biocontrol approach. Additionally, based on knowledge of stink bug and parasitoid behaviour, biology, and chemical ecology, we discuss the prospects of using semiochemical and other novel approaches to increase natural enemy efficacy and manage stink bug pests as part of sustainable, integrated control strategies.

Management of invasive stink bugs over a century of classical biological control

Recent stink bug invasions have renewed the interest in classical biological control as a potential pest control strategy. Evaluation of suitable candidates for classical biological control of B. hilaris is underway in North America, with research focusing on egg parasitoids, as the tachinid parasitoids of adult stink bugs (Diptera: Tachinidae) that have been recorded in Asia are poorly known and do not seem to have a significant impact on this host (Sforza et al., 2017; Bundy et al., 2018b). Similarly, the most important natural enemies of H. halys appear to be coevolved egg parasitoids in the native range of Eastern Asia (Zhang et al., 2017; Buffington et al., 2018; Hamilton et al., 2018; Leskey & Nielsen, 2018). The egg parasitoid Trissolcus japonicus (Ashmead) [Hymenoptera: Scelionidae (syn. Platygastridae s.l.; Sharkey, 2007, but see Popovici et al., 2017)] is the predominant natural enemy of H. halys in Asia and is therefore considered a promising biocontrol agent for this invasive pest (Zhang et al., 2017; Buffington et al., 2018; Leskey & Nielsen, 2018). However, environmental risk assessment of *T. japonicus* as candidate biocontrol agent, conducted in the USA and in Europe, raised ecological concerns due to the apparent low host specificity of T. japonicus and consequent risk that this species might attack non-target species in the areas of introduction, including beneficial predatory stink bugs (Hedstrom et al., 2017; Botch & Delfosse, 2018; Haye et al., 2020). Legislative restrictions in Europe and the USA severely limit the introduction of exotic species by focusing on perceived risks for non-targets rather than the benefits of pest reduction (van Lenteren et al., 2006; Rondoni et al., 2021). Now that the occurrence of T. japonicus has been reported in northern Italy (Sabbatini Peverieri et al., 2018), in early 2020 the Italian government authorized the mass production and release of this parasitoid in Italy, where H. halys has caused the highest economic losses in Europe since its arrival in 2012 (Maistrello et al., 2016, 2018). This is the first officially authorized release of the parasitoid against H. halys in Europe. As has occurred in Italy, constraints are probably becoming less relevant in the USA because of the recent finding of adventive populations of *T. japonicus* in the invaded areas (Talamas et al., 2015).

About 1 century ago, the first attempts with biological control of stink bugs started with *N. viridula* and they are still in progress globally. Classical biological control mostly involved the egg parasitoid *Trissolcus basalis* (Wollaston) (Hymenoptera: Scelionidae), which was introduced and released in nearly all geographical areas invaded by *N. viridula* (Esquivel et al., 2018). Additionally, the native parasitoids of adults, *Trichopoda pennipes* (Fabricius) and *Trichopoda giacomellii* (Blanchard) (Diptera: Tachinidae), were found parasitizing this species in North and South America, respectively, and were subsequently introduced into Hawaii (USA) and Australia for classical biological control of *N. viridula* (Liljesthröm & Rabinovich, 2004; Esquivel et al., 2018). *Trichopoda pennipes*, which attacks

adults and occasionally nymphs, was then accidentally introduced in Italy (Colazza et al., 1996) from where it spread to several countries in Europe and the Mediterranean (Tschorsnig, 2017). Despite the numerous classical biocontrol programs against *N. viridula*, this species is still an economically important pest worldwide except in South America (Panizzi & Lucini, 2016).

In contrast with other invasive stink bugs, no biocontrol efforts have been made against *M. histrionica* in the USA. However, several native natural enemies – the most common ones are the parasitoids *Trissolcus brochymenae* (Ashmead), *Trissolcus euschisti* (Ashmead) (Hymenoptera: Scelionidae), and *Ooencyrtus johnsoni* (Howard) (Hymenoptera: Encyrtidae) – have been recorded since the end of the 19th century but generally with low reported efficacy (McPherson et al., 2018).

Overall, naturally occurring and classical biological control of stink bugs, although useful in significantly reducing the exotic pest populations, are often inadequate in maintaining the pest below the economic injury level (McPherson, 2018). Therefore, additional approaches are being evaluated and developed regularly to improve the efficacy of parasitoids as biocontrol agents and to integrate biocontrol programs with other sustainable management methods targeting adult stink bugs. In the following sections, we report relevant case studies and discuss the potential application of conservation biological control and other sustainable control methods (Figure 1).

Case study 1: Selection of candidates for classical biological control of *Bagrada hilaris*

History, invasion, and pest status

The painted or Bagrada bug, B. hilaris, is a worldwide pest of brassicaceous crops (Bundy et al., 2018b). Like other stink bugs, it can also be a nuisance pest for humans by sheltering in homes (Faúndez, 2018). Bagrada hilaris is native to Africa, the Middle East, and Asia where it sporadically damages local crops (Gunn, 1918; Ahuja et al., 2008). Since its first introduction in California (USA) in 2008, it has become invasive in southern states of the USA (Reed et al., 2013) and in Hawaii (USA), Mexico, and Chile (Faúndez et al., 2016). In the Mediterranean basin, B. hilaris was accidentally introduced into Italy and Malta. Pinpointing the origin of invasive populations in the Americas will enable an understanding of the invasion routes of B. hilaris and ultimately help to find the parasitoid best suited to control this pest. A preliminary phylogeographical study using the DNA barcode region COI identified Pakistan as a source for the invasive B. hilaris in North America (Sforza et al., 2017).

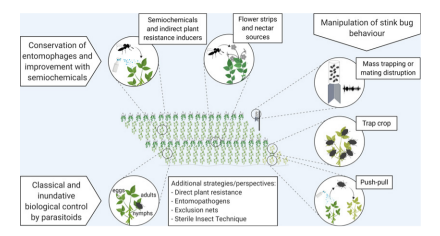


Figure 1 Possible strategies for successful stink bug management based on the integration of biological control (classical, inundative, or conservation) with manipulation of the natural enemy and/or herbivore behaviour, or with additional strategies (figure created with BioRender.com). [Colour figure can be viewed at wileyonlinelibrary.com]

This pest is multivoltine with gregarious behaviour (Reed et al., 2013; Bundy et al., 2018b). Unlike most of the 4 722 species of Pentatomidae (Rider, 2016) that glue eggs together in clusters onto plant material, B. hilaris deposits many of its eggs singly into the soil (Gunn, 1918; Taylor et al., 2014), which may affect egg parasitism (see below). Due to the economic impact of *B. hilaris* injury on crops in southwestern USA, its control currently relies on pesticide applications (Palumbo et al., 2016). Classical biological control could provide sustainable and long-term control, especially if implemented as part of an IPM program. To date, no classical biocontrol program is in place in any of the countries in the invaded range. However, the egg parasitoids Gryon myrmecophilum (Ashmead), Telenomus podisi (Ashmead), and Tr. basalis (all Hymenoptera: Scelionidae) were found parasitizing B. hilaris in Mexico (Felipe-Victoriano et al., 2019), suggesting a possible role for them in conservation biological control.

Exploration for natural enemies in the native range

Surveys for natural enemies began in countries where B. hilaris is native, including South Africa and India. As for other invasive stink bugs (see below), predators and resident North American parasitoids have been reported (Bundy et al., 2018b). However, we present only data on egg parasitoids from the native geographical range of the pest. Gunn (1918) first recorded the presence of an egg parasitoid emerging from artificially buried eggs of B. hilaris in cabbage and cauliflower fields. Subsequently, egg parasitoids in the scelionid genera Telenomus, Typhodytes (Mani, 1942; Samuel, 1942), and Trissolcus (Chacko & Katiyar, 1961) were collected from India. After the invasion of B. hilaris in the Americas, surveys were conducted in 2015 in Pakistan. Three species were collected, viz., Trissolcus hyalinipennis Rajmohana & Narendran (syn. Allophanurus indicus Subba Rao & Chacko), Grvon gonikopalense Sharma (Scelionidae), and an Ooencyrtus sp.

(Encyrtidae) (Mahmood et al., 2015). The latter was the first report of an *Ooencyrtus* species on *B. hilaris* eggs. These egg parasitoid candidates are currently under evaluation in the USA and European quarantine facilities. Since 2016, new surveys for *B. hilaris* biocontrol candidates have been conducted in Kenya and South Africa within the framework of Access and Benefit Sharing agreements (Mason et al., 2018). Various collecting protocols, including sentinel eggs and Malaise trapping, are being used to study the native biodiversity of egg parasitoids of *B. hilaris* (RFH Sforza, M Kasina, P Addison, MC Bon & E Talamas, unpubl.).

Promising candidates

Among the parasitoid species collected in Pakistan in 2015, only two have been studied in the laboratory. The basic biological traits of *Tr. hyalinipennis* were described by Subba Rao & Chacko (1961). Their studies reported a very high fecundity level of 158 progeny per female and a longevity of over 1 month. Studies of *G. gonikopalense* at the USDA-ARS European Biological Control Laboratory (Montpellier, France) suggest this species is a promising biocontrol candidate (Martel et al., 2019). Both the host and parasitoid share the same thermal requirements for their development and reproduction, suggesting that their development will be synchronous in the field.

An important criterion for an effective candidate biocontrol agent is its foraging capacity for *B. hilaris* eggs buried in the soil, as was highlighted by Bundy et al. (2018b). Recent investigations under laboratory conditions have shown that *Tr. hyalinipennis* is unable to parasitize buried eggs, whereas *G. gonikopalense* readily finds and parasitizes *B. hilaris* eggs in the soil (G Martel & RFH Sforza, unpubl.). This information provides a basis for investigating whether *G. gonikopalense* effectively parasitizes *B. hilaris* eggs under natural conditions, especially considering that *B. hilaris* is able to oviposit both in the soil and on

its host plant (Taylor et al., 2014). Another criterion is the host specificity of the selected egg parasitoids, which is currently under investigation in USA quarantine facilities for *B. hilaris* and several other stink bugs. If the above-mentioned criteria are met, the release of *G. gonikopalense* in North America can be considered. In the meantime, foreign exploration in Africa will continue to search for other coevolved parasitoids.

Case study 2: Biological control of *Halyomorpha halys* in North America

History, invasion, and pest status

The brown marmorated stink bug, *H. halys*, was first detected in the USA in the mid-1990s (Leskey & Nielsen, 2018) in Allentown, PA (Hoebeke & Carter, 2003), and has since spread to more than 42 states (Leskey & Nielsen, 2018). In Canada, it was first detected in Ontario and Quebec in 2010 (Fogain & Graff, 2011) and subsequently in British Columbia (Abram et al., 2017c). This species is a polyphagous pest with a host range of more than 300 plant species (Lee et al., 2013; Bergmann et al., 2016). In North America, this includes over 170 plants from more than a dozen families, and apples, peaches, nectarines, pears, grapes, sweet corn, soybeans, and hazelnuts are among the most susceptible cultivated hosts (Leskey & Nielsen, 2018).

Natural control by native parasitoids and predators

Early biological control research in North America focused on identifying the indigenous natural enemies of *H. halys*. These studies mainly concentrated on the egg stage, using sentinel (fresh and frozen) H. halys egg masses to assess predation and parasitism (Dieckhoff et al., 2017; Abram et al., 2017b; Leskey & Nielsen, 2018). In general, control by existing natural enemies has been considered ineffective (Ogburn et al., 2016; Cornelius et al., 2016a; Abram et al., 2017b). In agreement with the enemy release hypothesis (Keane & Crawley, 2002), H. halys seems to have escaped from its natural enemies in the invaded North American areas. For example, in organic agro-ecosystems across the eastern USA, maximum levels of natural biological control were estimated at 20%, primarily caused by chewing predators (Ogburn et al., 2016) whose efficacy is dependent on the H. halys life stage (Morrison et al., 2016; Pote & Nielsen, 2017). In the laboratory, late-instar Chrysoperla carnea (Stephens) (Neuroptera: Chrysopidae), katydids (Orthoptera: Tettigoniidae), earwigs (Dermaptera: Forficulidae), jumping spiders (Araneae: Salticidae), crickets (Orthoptera: Gryllidae), and ground beetles (Coleoptera: Scarabaeidae) were effective predators of H. halys eggs (Abram et al., 2015; Morrison et al., 2016). Grasshoppers (Orthoptera: Acrididae), Coccinella septempunctata (L.) (Coleoptera: Coccinellidae), and the spined soldier bug *Podisus maculiventris* (Say) (Pentatomidae) also preyed on eggs, whereas several hemipteran predators attacked young nymphs (Pote & Nielsen, 2017). Using surveillance cameras, Pote (2018) reported katydids feeding on *H. halys* eggs in the field. Possibly, indigenous generalist predators use pentatomid semiochemical cues to locate *H. halys* (Fraga et al., 2017).

Mortality from parasitism of H. halys eggs by indigenous parasitoids in North America tends to be lower than that from predation (Ogburn et al., 2016), accounting for less than 5% of parasitoid emergence from eggs in 87% of all surveys (Abram et al., 2017b). However, levels of parasitism and predation seem to vary depending on habitat (Cornelius et al., 2016a; Tillman et al., 2020), with parasitism greater than predation in woody ornamental nurseries (Jones et al., 2017), whereas the opposite is true for organic agroecosystems (Ogburn et al., 2016). Three parasitoid families are commonly found attacking H. halys eggs in the USA and Canada: Scelionidae (Telenomus, Trissolcus, and Gryon spp.), Eupelmidae (Anastatus spp.), and Encyrtidae (Ooencyrtus spp.) (Abram et al., 2017b; Leskey & Nielsen, 2018). According to Abram et al. (2017b), Anastatus and Trissolcus tend to dominate ornamental, semi-natural/urban, and forest habitats, whereas Te. podisi dominates field/vegetable crops and orchard habitats (see also Herlihy et al., 2016, and Tillman et al., 2020). One explanation for the low parasitism rates observed of H. halys eggs is that even when eggs are attacked, the native parasitoids frequently fail to develop to emergence, although they may still kill the host (Cornelius et al., 2016b). Abram et al. (2014) showed that H. halys eggs are attacked by Te. podisi at rates similar to indigenous P. maculiventris eggs. However, successful development occurred only in the latter species. They argued that H. halys acts as an evolutionary trap for this parasitoid, which could result in an increase in population levels of indigenous pentatomids. It is also worth noting that, although convenient, studies using sentinel eggs may underestimate the levels of parasitism (Jones et al., 2014; but see Herlihy et al., 2016, and Dieckhoff et al., 2017). This reduced parasitism could be due to the absence of cues used by indigenous parasitoids in host finding and recognition, i.e., host kairomones (Conti et al., 2003; Laumann et al., 2009; Tognon et al., 2016, 2017; Rondoni et al., 2017; Boyle et al., 2020) and oviposition-induced plant synomones (Colazza et al., 2004a; Conti & Colazza, 2012; Rondoni et al., 2017).

Exploration for natural enemies in the native range and biological control

To improve *H. halys* biological control, efforts were initiated in the late 2000s in the USA to introduce natural

enemies from its native range. Classical biocontrol efforts targeted egg parasitoids of the genus Trissolcus, including Tr. japonicus, the predominant egg parasitoid and most important natural enemy of H. halys in Asia capable of inflicting high rates of parasitism in landscapes and orchard crops (Zhang et al., 2017; Buffington et al., 2018; Leskey & Nielsen, 2018). In 2014, although still under evaluation in quarantine facilities in the USA, a population of Tr. japonicus was found during routine sentinel egg surveys in a wooded habitat in Beltsville, MD (Talamas et al., 2015; Buffington et al., 2018). Molecular analyses revealed that the Tr. japonicus specimens found were different than those maintained in quarantine, indicating that it was introduced into the USA by other means via an unknown pathway (Buffington et al., 2018). Since then, Tr. japonicus has been reported in at least 13 states across the USA (Milnes et al., 2016; Hedstrom et al., 2017; Morrison et al., 2018; Leskey & Nielsen, 2018; Jarrett et al., 2019) and two provinces in Canada (Abram et al., 2019; Gariepy & Talamas, 2019). They comprise three distinct haplotype populations, indicating that there have been several independent introductions whose source regions have not yet been clearly identified (MC Bon, pers. comm.). Although Tr. japonicus has been reported mainly from unmanaged landscape habitats, it has also been found in peach orchards (Kaser et al., 2018).

Laboratory testing of Tr. japonicus against North American pentatomids has shown a marked preference for H. halys but not complete host specificity (Hedstrom et al., 2017; Botch & Delfosse, 2018; Lara et al., 2019). Potential alternate hosts may be less physiologically suitable for development than H. halys if they are accepted for oviposition, producing smaller and less fit progeny (Botch & Delfosse, 2018). The first field study of a Tr. japonicus population in North America appears to confirm these laboratory studies, at least for several native pest stink bugs in the Pacific Northwest (Milnes & Beers, 2019). A degree of specificity is also conferred by chemicals associated with H. halys that are used by Tr. japonicus in host finding and recognition. In choice tests, Tr. japonicus remained longer on surfaces with H. halys contact kairomones (Hedstrom et al., 2017). Furthermore, they responded more strongly to H. halys kairomone traces deposited on leaves of apple, maple, and soybean than to traces of the predatory P. maculiventris (Boyle et al., 2020), a non-target species that is an important predator in agricultural habitats of several pests, including H. halys (Pote & Nielsen, 2017). Parasitoid females detected and responded in a similar manner to kairomone traces on leaves of host plants, increasing their search time in a Y-tube olfactometer, and female Tr. japonicus were attracted to n-tridecane but repelled by (E)-2-decenal, two defensive compounds emitted from H. halys

males and females (Zhong et al., 2017). In other experiments, *Tr. japonicus* females were attracted to volatiles from gravid females and mature males of *H. halys* and to volatiles induced in tomato plants by oviposition and feeding of this coevolved host, but females did not respond to similar cues associated with *P. maculiventris* (Bertoldi et al., 2019).

Although *Tr. japonicus* is also capable of parasitizing *P. maculiventris* (Hedstrom et al., 2017; Botch & Delfosse, 2018), the probability of *Tr. japonicus* locating and parasitizing *P. maculiventris* under field conditions is likely to be lower than for *H. halys* (Bertoldi et al., 2019; Boyle et al., 2020). *Trissolcus japonicus* did not appear to be attracted to the *H. halys* aggregation pheromone (Morrison et al., 2018), and it was never captured in stink bug traps baited with *H. halys* and *P. maculiventris* pheromones (Boyle, 2017). Whether populations of indigenous pentatomids in North America will be negatively impacted by *Tr. japonicus* and whether this foreign parasitoid will provide better biological control of *H. halys* remain open questions that must be answered with continued field studies.

Case study 3: Biological control of *Halyomorpha halys* in Europe

Invasion of Europe and pest status

The European invasion by H. halys started in Switzerland in 2004 (Haye et al., 2015b) followed by the colonization of neighbouring countries and interceptions at customs/ ports/airports (Claerebout et al., 2018). Genetic analyses demonstrate a high diversity of the invading H. halys populations, especially in Italy (Cesari et al., 2017) and Greece (Morrison et al., 2017), indicating multiple introductions that are still ongoing from their native range in Asia and other invaded countries. Serious damage in Italy was observed on peach, pear, and hazelnut, starting from 2013 (Maistrello et al., 2017; Bosco et al., 2018). Outbreaks in northern Italy in 2019 caused more than € 356 million damage on pear, peach, and nectarine, with up to 80-100% yield losses (CSO, 2019). Damage was reported also on apple, kiwi, and other fruit crops, as well as on horticultural and row crops (peppers, tomatoes, maize, and beans). Following H. halys invasion, previous IPM programs were seriously disrupted and official IPM guidelines had to be revised to allow for increased numbers of treatments with broad-spectrum insecticides.

Natural control by native parasitoids and predators

Due to restrictions on the practice of classical biological control in Europe, initial studies aimed to discover native natural enemies that were able to adapt to the new invasive species. Field surveys were performed in Switzerland, Italy, and Georgia, mainly to detect egg parasitoids and predators either using sentinel egg masses (frozen or fresh) or by collecting naturally laid egg masses. Overall, egg parasitism ranged from 3 to 39% in Switzerland (using frozen sentinel eggs), from 1 to 3% in Emilia Romagna, Italy (using fresh sentinel eggs), and from 12 to 21% in Piedmont, Italy (collecting naturally laid eggs). The generalist egg parasitoid Anastatus bifasciatus (Geoffroy) (Eupelmidae) was the predominant species in all surveys (Haye et al., 2015a; Costi et al., 2018; Kereselidze et al., 2018; Moraglio et al., 2020), confirming its ability to develop in both frozen and fresh eggs. Other species found included Trissolcus cultratus (Mayr) (Scelionidae) in Switzerland (Haye et al., 2015a), and Trissolcus kozlovi Rjachovsky, Trissolcus belenus (Walker) [syn. Tr. semistriatus (Nees von Esenbeck) sensu Talamas et al., 2017] (Tortorici et al., 2019), Tr. basalis, Telenomus turesis Walker (Scelionidae) and the hyperparasitoid Acroclisoides sinicus (Huang & Liao) (Pteromalidae) in Piedmont (Sabbatini Peverieri et al., 2019; Moraglio et al., 2020). In these surveys, a wide range of eggs were lost or showed symptoms of predation (chewing or sucking): up to 31% in Switzerland (Haye et al., 2015a), 2-5% in Emilia Romagna (Costi et al., 2018), and 0.4-9% in Piedmont (Moraglio et al., 2020).

Laboratory no-choice tests were carried out to evaluate the potential of several parasitoid species to attack eggs of European native pentatomids as well as some generalist predators. *Ooencyrtus telenomicida* (Vassiliev) (Encyrtidae) achieved 36% parasitism of *H. halys* eggs, followed by *Ooencyrtus pityocampae* (Mercet) (21%), *A. bifasciatus* (20%), and *Telenomus chloropus* Thomson (6%) (Roversi et al., 2016). In another study, seven scelionid species (six *Trissolcus* spp. and *Te. turesis*) were tested on *H. halys* eggs. All the *Trissolcus* spp. caused significantly higher egg abortion compared with unexposed eggs, but only *Tr. kozlovi* was able to produce offspring from fresh eggs (Moraglio et al., 2021), confirming what was observed in field surveys in Piedmont (Moraglio et al., 2020).

Several species of wild-collected native European generalist predators were tested for their ability to feed on eggs and nymphs of *H. halys* in no-choice tests (Bulgarini et al., 2019). Only the omnivorous *Pholidoptera littoralis* (Fieber) (Orthoptera: Tettigoniidae) was capable of preying on eggs and first and second instars. The other predators did not feed on the eggs. Young *H. halys* nymphs were preyed on by *Nagusta goedelii* (Kolenati), *Rhynocoris iracundus* (Poda) (both Hemiptera: Reduviidae), and two *Himacerus* species (Hemiptera: Nabidae), and *R. iracundus* also fed on the adults (Bulgarini et al., 2019). In a laboratory study, the arboreal ant *Crematogaster scutellaris* (Olivier) (Hymenoptera: Formicidae) did not prey on eggs but

proved to be effective at preying on all nymphal instars (Castracani et al., 2017). In addition, the predatory potential of bats (Chiroptera) was assessed by screening for the presence of *H. halys* DNA (Piemontese et al., 2019) in guano samples from nine Italian bat species collected in natural and agricultural environments. Two genera of bats, *Myotis* and *Nyctalus* (both Vespertilionidae), showed evidence of feeding on *H. halys* (Piemontese et al., 2019). In a similar approach, analysis of the gut contents of field-collected arthropods identified *H. halys* DNA in predatory insects and arachnids belonging to Dermaptera, Orthoptera, Hemiptera, Opiliones, and Araneae (G Bulgarini, L Piemontese, M Cesari, R Guidetti & L Maistrello, unpubl.).

Control by exotic parasitoids and possible impact on native nontarget stink bugs

Recently, field surveys in Switzerland and Italy discovered the presence of populations of Tr. japonicus and Trissolcus mitsukurii (Ashmead), which are considered the most effective egg parasitoids in China and Japan, respectively (Arakawa & Namura, 2002; Zhang et al., 2017), although the higher prevalence of Tr. mitsukurii over Tr. japonicus in Japan is debated (KA Hoelmer, unpubl.). Trissolcus japonicus was found first in Canton Ticino, Switzerland, in 2017 and in Piedmont and Lombardy, Italy, in 2018 (Sabbatini Peverieri et al., 2018; Stahl et al., 2019b; Moraglio et al., 2020). A genetic analysis using the barcode mitochondrial DNA determined the closest match of the 'Ticino populations' with Japanese populations, but the pathways of entry into Switzerland remain unknown (Stahl et al., 2019b). Trissolcus mitsukurii was first recorded in northeastern Italy starting from 2016 (Scaccini et al., 2020). After the discovery of populations of Tr. japonicus and Tr. mitsukurii in Italy, the situation has been changing, as highlighted by the increased parasitism rate in Piedmont where, together with A. bifasciatus, Tr. japonicus was detected in 2018 (Moraglio et al., 2020). Given the availability of their host, the distribution of these exotic egg parasitoids is expected to expand, even in the absence of human intervention.

The prospective host range of *Tr. japonicus* in Europe was evaluated in no-choice tests, followed by large-arena choice tests (Haye et al., 2020). The developmental suitability of European non-target host species for *Tr. japonicus* was demonstrated in no-choice tests by offspring emergence from 11 out of 13 non-target species tested (85%). In large-arena choice tests, the degree of non-target parasitism was substantially reduced (three out of four suitable species were parasitized), whereas parasitism of *Palomena prasina* L. (Pentatomidae) eggs was comparable to that of *H. halys* (Haye et al., 2020). Field data from the

invaded areas in Switzerland and Italy can contribute to a risk-benefit evaluation of releasing or re-locating Tr. *japonicus* populations into other parts of Europe.

Augmentative releases and efficacy of native parasitoids

Given that A. bifasciatus is the most prevalent native egg parasitoid of H. halys in field surveys (Haye et al., 2015a; Costi et al., 2018; Moraglio et al., 2020) and that it is capable of developing in viable H. halvs eggs (Roversi et al., 2016; Abram et al., 2017b), this species was selected as a candidate for augmentative releases. Anastatus bifasciatus can exploit cues associated with the non-coevolved host during its searching, as parasitoid females responded positively both to adult H. halys male volatiles and to H. halysinduced plant volatiles (Rondoni et al., 2017). Trials were performed in four apple orchards in Switzerland for 3 years and in an organic pear orchard in northern Italy for 1 year, releasing an equivalent of 11 000-26 000 A. bifasciatus females per ha (Stahl et al., 2019a). Parasitism averaged 6% (range: 2-16%) on frozen H. halys sentinel egg masses and 8% on frozen egg masses of lepidopteran non-target species. At the Italian site, parasitism on naturally laid eggs was considerably higher (49%) than on frozen sentinel eggs (16%). This suggests that although large quantities of frozen eggs are easier to obtain for experiments, data based upon their use in the field are likely to underestimate the actual impact of parasitism.

In 2019, further trials were performed in Emilia Romagna, Italy, where 10 000 A. bifasciatus females per ha were released over 5 weeks from the beginning of egg laying by H. halys (June-early July) in a small wooded area adjacent to a pear orchard. Parasitism averaged 9% on naturally laid eggs and 2.5% on frozen sentinel egg masses (Maistrello et al., 2020). However, these field trials underestimated the overall impact of A. bifasciatus, as they did not account for host eggs killed due to host feeding. According to previous laboratory trials, host feeding by A. bifasciatus may double the estimated host mortality (Stahl et al., 2019a).

Case study 4: Pre-emptive biological control of Halyomorpha halys in New Zealand – a world first

Risk of invasion, potential impact, and pre-emptive biocontrol approach

In New Zealand, H. halys presents a major risk not only to the sustainability and economics of food production but also to conservation due to its biology and very broad host range (Duthie, 2012; Lee et al., 2013; Bergmann et al., 2016). A recent assessment of the potential economic impact of H. halvs in New Zealand found that if it established, it could cost the horticulture and arable industries

4.2 billion NZD in 2038 (New Zealand Institute of Economic Research, 2017). In addition, some New Zealand native plant species may be at risk of attack by H. halys as well (Duthie, 2012).

Although H. halys is not yet known to be present in New Zealand, there is a high risk of entry and establishment, with more than 212 live interceptions of H. halys adults at New Zealand's border during the latest high-risk season (i.e., September 2018-April 2019), totalling more than 1 500 adults. Invasive pests can sometimes be eradicated on arrival, but if eradication attempts fail or are deemed technically or economically unfeasible, then there will be a need to undertake area-wide pest management. If this is the case, classical biological control is expected to become a key strategy for reducing H. halvs populations throughout New Zealand. The severity and imminent nature of H. halys incursion meant that there was an urgency for developing a pre-emptive classical biocontrol approach before the arrival of H. halys. Pre-emptive biological control is a novel approach that has the potential to enhance preparedness for a possible invasion of pest species and accelerate the response to invasive pests. With this approach, natural enemies can be selected, risk assessment studies carried out, and their potential release pre-approved by regulators for timely release after arrival of the pest. This would significantly reduce pest densities and rates of spread, ultimately reducing the economic or environmental damage associated with the pest (Hoddle et al., 2018). In preparation for a potential arrival/establishment of H. halys in New Zealand, a pre-emptive classical biocontrol program for this pest using the egg parasitoid Tr. japonicus, the most promising biocontrol agent of H. halys, was initiated in December 2015. This program aimed to determine the biosafety of Tr. japonicus to potential nontarget species in New Zealand and to pre-approve its release to be ready for the anticipated arrival of *H. halys*.

Risk assessment for biological control of Halyomorpha halys

The potential host range of Tr. japonicus in New Zealand was investigated in quarantine using imported parasitized H. halys eggs from USDA-ARS (Newark, DE, USA) between November 2015 and December 2016. Egg masses of seven species of New Zealand Pentatomidae, including one sub-species, were individually exposed to naive mated female Tr. japonicus in no-choice laboratory experiments. All tested species were non-endemic but naturalised species to New Zealand. The only Pentatomidae species endemic to New Zealand, Hypsithocus hudsonae Bergroth, was not available at the time and could not be tested (Charles et al., 2019). Results from laboratory host-specificity testing demonstrated that the predatory Pentatomidae Cermatulus nasalis (Westwood) nasalis, C. nasalis hudsoni,

and Oechalia schellenbergii (Guérin), and the phytophagous Monteithiella humeralis (Walker), Dictyotus caenosus (Westwood), Glaucias amyoti (Dallas), and Cuspicona simplex Walker were all physiological hosts for Tr. japonicus, although not all were equally susceptible to parasitism (Charles et al., 2019). No development or emergence of Tr. japonicus from eggs of N. viridula was observed (Charles et al., 2019).

In addition, a CLIMEX bioclimatic model was developed to estimate the potential global distribution of Tr. japonicus with particular reference to New Zealand and to investigate possible overlaps with the current distribution of potential non-target species (Avila & Charles, 2018). In the native range of Tr. japonicus, the model predicts its presence or potential expansion coinciding with the native range of H. halys (confirmed by specimen collection records that show it occurs throughout the entire native range of H. halvs) into most humid-subtropical and humid-continental areas (Avila & Charles, 2018). Globally, the model projected that many temperate, Mediterranean, and subtropical areas could be suitable for the establishment of Tr. japonicus. Laboratory studies also demonstrated that Tr. japonicus and Tr. cultratus could survive periods of winter temperatures as cold or colder as tolerated by their host, H. halys (Nystrom et al., 2017). In New Zealand, the north appears moderately to highly suitable for Tr. japonicus, whereas southern regions are mostly marginal. The risk posed by Tr. japonicus to non-target species in New Zealand is predicted to vary between different pentatomids (Avila & Charles, 2018). CLIMEX projections of the potential distribution of Tr. japonicus provide useful guidance for selecting release sites if importation/release of this parasitoid needs to be carried out in New Zealand or worldwide.

Approval to release Trissolcus japonicus in New Zealand

The importation, development, and release of new organisms into New Zealand are under strict regulation and must be approved by the Environmental Protection Authority (EPA), which implements the Hazardous Substances and New Organisms (HSNO) Act 1996 (Ehlers et al., 2019). An applicant seeking approval to release a candidate biocontrol agent must submit an application that includes a risk/benefit analysis providing evidence to support the proposal for release and demonstrating that it meets the minimum standards of the HSNO Act. The application submitted to the EPA must provide information about potential adverse effects and the expected economic, social, and cultural benefits of the introduction. The EPA then conducts a full evaluation and review of the benefits and risks associated with the proposed biocontrol agent (Barratt & Ehlers, 2017; Ehlers et al., 2019). If the perceived benefits outweigh the risks, then EPA may grant approval for release, subject (or not) to further conditions.

In March 2018, an application was submitted to the EPA by a representative group from the horticultural industries (BMSB Council) to seek pre-approval to import and release Tr. japonicus into New Zealand in the event of a H. halys incursion. The application included host range testing, bioclimatic modelling, and substantial additional information on economic and social benefits. As a result of this application, in August 2018, the EPA granted approval for a conditional release of Tr. japonicus in the event that a H. halys incursion is detected in New Zealand, and it is subject to the development of an appropriate release plan (EPA, 2018). This is the first approval granted to release a biocontrol agent into New Zealand prior to the arrival of its target pest and the first example worldwide to approve the future release of Tr. japonicus before the arrival of H. halys. The successful implementation of this approach will provide New Zealand with the opportunity for a quicker response against a H. halvs invasion and a greater chance of achieving an eradication or implementing an early area-wide pest management plan. This novel example of pre-emptive biological control might provide the impetus for biocontrol practitioners to adopt such an approach in the future for the early management of exotic pest incursions.

Case study 5: Biological control of stink bugs in South America

General overview of stink bugs as pests in South America

In South America, stink bugs are major pests primarily in arable crops, such as soybean, beans, and maize (Panizzi & Silva, 2012), which cover large agricultural areas. For example, the total area planted with soybean in Brazil, Paraguay, Uruguay, and Argentina is approximately 56 million ha (Trase, 2018). Since the start of soybean production and its expansion in South America, increasing problems with stink bugs have been reported. In general, these are species complexes with the dominant species varying among regions (Saluso et al., 2007; Ribeiro et al., 2009; Panizzi & Lucini, 2016; Aquino et al., 2019). The first serious pest mentioned in the literature was the cosmopolitan N. viridula (Panizzi & Lucini, 2016). Interestingly, its importance has changed over time. Currently, N. viridula is found at very low population levels in various regions of Brazil and Argentina (Panizzi & Lucini, 2016). This phenomenon could be due to adaptation to new cropping systems, expansion of soybean culture to hotter regions and broad adoption of non-tillage practices that favour native species. As a result, the dominant stink bugs in crops are currently native species such as Euschistus heros (Fabricius) (Pentatomidae) in hot regions (e.g., central and north Brazil) (Aquino et al., 2019) and *P. guildinii* in cold regions (e.g., southern Brazil, Paraguay, Uruguay, and Argentina) (Panizzi & Lucini, 2016). Other than *N. viridula*, no serious problems have been observed with invasive stink bugs in South America. However, because in recent years *H. halys* and *B. hilaris* have been reported in Chile, both species represent a serious threat to South American agriculture (Faúndez et al., 2016; Faúndez & Rider, 2017).

Classical and inundative biological control

The use of parasitoids for stink bug management in Brazil started in the 1990s with inundative biological control as part of an IPM program developed in the southern region of the country (Parana state). This biocontrol program was based on the use of the exotic Tr. basalis, which was already established in South America for the biological control of N. viridula, the main soybean pest at that time, and was shown to be especially useful for small to medium cultured areas surrounded by rivers. After demonstrating effective levels of control, the program was extended to more than 18 000 ha (Corrêa-Ferreira et al., 2000; Corrêa-Ferreira, 2002). In spite of its success, the program was eventually abandoned by growers who adopted new cropping systems with non-tillage practices, transgenic cultivars, and pest management based on cheap insecticides (Panizzi, 2013).

In Brazil, current efforts focus on E. heros control with Te. podisi as its main natural enemy. Because of the high dispersal capacity of this parasitoid, release of large numbers of individuals (ca. 5 000 individuals ha⁻¹) is necessary (Corrêa-Ferreira, 2002), requiring the rearing not only of many wasps but also their host. Mass rearing systems for stink bugs, including the use of artificial diets, have improved parasitoid production (Silva et al., 2008, 2018; Mendoza et al., 2016; Hayashida et al., 2018). This rearing system, together with new technologies - such as the preservation of host eggs and parasitoids at low temperatures until field release (Silva et al., 2018), protection of parasitized eggs in cardboard capsules, drone-based deployment systems, and accurate spatio-temporal field releases - could favour biocontrol programs in the near future. The high susceptibility of egg parasitoids to insecticides (Corrêa-Ferreira et al., 2010) is also important, so the identification and use of selective products are critical for successful inundative biocontrol strategies (Stecca et al., 2017). The commercial production of Te. podisi in Brazil is currently under consideration for license approval, which will allow a wider availability of parasitoids to growers.

In Argentina, a biocontrol program against *N. viridula* was implemented in 1981 with the introduction of *Tr. basalis* from Australia. A few years after the release of *Tr. basalis*, *N. viridula* was reduced to low densities (Crouzel & Saini, 1983). This was the only classical biocontrol effort for stink bugs in Argentina, and no subsequent evaluations were made regarding the success of the biocontrol program in the regions of release (Molinari et al., 2008).

In Uruguay, a multi-institution project was launched in 2005 that combined public and private interests for mass rearing and release of *Te. podisi*, with the aim of increasing the levels of parasitism of *P. guildinii* and reducing the use of insecticides. Releases were initially made of parasitized eggs as a source of parasitoids, but because of high levels of natural predation, adult wasps were released in the second stage of the project. Post-release parasitism of *P. guildinii* eggs by *Te. podisi* in two regions (Dolores and Paysandú) was 67 and 69%, respectively, which was higher than natural parasitism in areas without releases, i.e., 52 and 47%, respectively (Castiglioni et al., 2007).

Conservation biological control and semiochemical-based strategies to enhance parasitoid efficacy

Because of the high diversity of natural enemies (Cingolani, 2012), especially of egg parasitoids (Scelionidae), which can reach high natural parasitism rates of 60-80% (Corrêa-Ferreira & Moscardi, 1995; Cingolani et al., 2014a,b; Paz-Neto et al., 2015; Zerbino & Panizzi, 2019), conservation biological control appears to be an effective approach for stink bug management in South America. However, due to the expansion of agricultural fields, landscape and habitat management may be necessary to enhance the ecological services provided by parasitoids (Aquino et al., 2019). Other tools that could improve conservation biological control of stink bugs include the recruitment of parasitoids using semiochemicals (Vieira et al., 2013, 2014) and the use of soybean cultivars producing volatiles attractive to parasitoids (Michereff et al., 2014, 2016). Moreover, attract-and-reward (Simpson et al., 2011) and push-pull (Khan et al., 2016) strategies could increase parasitism rates of semiochemically or naturally attracted parasitoids in the field. The potential of flowering plants to improve Tr. basalis attraction and parasitism rates of N. viridula eggs has also been demonstrated (Foti et al., 2017, 2019). An attract-and-reward strategy for Te. podisi and other species of South American scelionids is under investigation (RA Laumann, MF Aquino, MC Blassioli Moraes & M Borges, unpubl.). Conservation biological control of N. viridula and semiochemical-based strategies against stink bugs are also discussed more in detail below.

38 Conti et al.

Case study 6: Conservation biological control of Nezara viridula

Global pest status of Nezara viridula

Perhaps as a result of climate warming, N. viridula seems to be rapidly expanding its range in both hemispheres to regions that were previously not warm enough to sustain the survival of the species. For instance, in Japan, N. viridula was confined to the southwestern part of the archipelago since its earliest record there in the late 19th century. However, the species has recently expanded its range into the warm-temperate zone and now reaches the central part of the archipelago (Tougou et al., 2009). Similarly, N. viridula is currently expanding its geographical range in Europe to the north (Salisbury et al., 2009; Marcu & Grozea, 2018). Reports of its occurrence have been documented in Slovakia (Hemala & Kment, 2017) and in The Netherlands, where an estimated 10-15% of sweet pepper greenhouses were infested in 2019 (H Hoogerbrugge, pers. comm.). Currently, N. viridula is considered a key pest in only a limited number of agricultural crops, mainly tomatoes and legumes (Esquivel et al., 2018). Changes in geographical distribution may profoundly affect its status as agricultural pest, particularly in the newly invaded areas. Given the need for sustainable management solutions, the possibility of combining the release of natural enemies with cost-effective approaches that involve habitat manipulation as a conservation biocontrol measure can be a valuable alternative to pesticides for controlling stink bug populations (Tillman, 2017).

Conservation biological control: habitat manipulation and cultural practices

Conservation biological control comprises a variety of methods, including habitat manipulation and cultural practices that aim at supporting natural enemy populations in the vicinity and within agricultural areas and thus reducing herbivory on crops (Gontijo, 2019). A particularly favoured measure is the planting of flower strips to provide parasitoids and predators with sugar resources, shelter, alternative prey, and a suitable microclimate (González-Chang et al., 2019). Although increasing habitat diversity per se may not lead to a consistent enhancement of pest control (Karp et al., 2018), it has been advocated that the 'right kind of diversity' is necessary, meaning that the traits of selected flowering plants must match the requirements of the targeted biocontrol agent (van Rijn & Wäckers, 2016). For example, corolla length and floral scent have been shown to be decisive features that confer a certain degree of specificity in the interactions between flowers and visiting insects (Tillman, 2017). In recent years, conserving and enhancing natural enemies for

controlling phytophagous stink bugs by providing floral resources have been receiving increased interest (Tillman, 2017). Several studies have demonstrated that a strategic arrangement of flowering plants can attract and enhance the efficacy of these parasitoids (Tillman, 2017). In the southeastern USA, where peanuts and cotton are widely grown, stink bugs such as N. viridula and Chinavia hilaris (Say) (Pentatomidae) have become important pests in cotton. Field experiments with potted milkweed, Asclepias curassavica L. (Apocynaceae), placed at peanut-cotton interfaces to serve as a nectar source for natural enemies, showed that parasitism of both adult stink bug species was significantly higher in the two years of the experiment. In this case, parasitism was mainly due to the tachinid fly Tric. pennipes (Tillman & Carpenter, 2014). Similarly, flowering buckwheat, Fagopyrum esculentum Moench (Polygonaceae), was able to increase the efficacy of Tric. pennipes parasitizing N. viridula in cotton (Tillman, 2017).

Studies conducted in Sicily, Italy, have shown differing impacts of buckwheat floral nectar on the abundance of Tr. basalis and O. telenomicida, two sympatric egg parasitoids of N. viridula (Foti et al., 2017, 2019). In the case of Tr. basalis, laboratory tests to screen for suitable non-crop plants showed that buckwheat and basil flowers have a positive effect on parasitoid longevity (Rahat et al., 2005; Foti et al., 2017). Buckwheat was more attractive to Tr. basalis than other companion plants because of characteristic compounds in the floral scent (Foti et al., 2017). When buckwheat margins were grown alongside tomato plots, Tr. basalis located and parasitized more N. viridula egg masses during the growing season (Foti et al., 2019). However, such beneficial effects cannot be generalized and may not apply to all N. viridula egg parasitoids. In fact, laboratory tests showed that buckwheat scent repels female O. telenomicida and flowering buckwheat margins fail to increase stink bug parasitism under field conditions (Foti et al., 2019). Although a net beneficial effect can be expected due to the dominance of Tr. basalis, the possibility that there may be contrasting effects of floral scent within the parasitoid guild highlights the importance of selecting appropriate companion plants on a case-by-case basis. Researchers might even need to go one step further, as it can be necessary to select not only a suitable species but also the right cultivar of a given companion plant. Field studies in Florida, USA, for instance, demonstrated large differences between three sweet alyssum, Lobularia maritima (L.) Desv. (Brassicaceae), varieties in attracting predators of N. viridula when grown within tomato crops (Haseeb et al., 2018). Another obstacle for conservation biological control of N. viridula is the fact that parasitoids such as Tr. basalis tend to stay within a limited range of the flower margin, rather than moving deeper into the crop.

Future work will need to focus on the question how this obstacle can be overcome. Parasitoid distribution could be enhanced, e.g., by planting flower strips within fields and using modern GPS technology to avoid accidental spraying. Large-scale trials testing this concept are currently underway in the UK (ASSIST programme; https://www.assist.ceh.ac.uk).

Future prospects for improving stink bug control

The role of biological control in reducing invasive pest populations is widely recognized, although only a small amount (around one-tenth) of past natural enemy introductions provided satisfactory control (Cock et al., 2016). Therefore, because of the generally low economic injury levels of stink bugs (McPherson, 2018), it is not surprising that the efficacy of egg parasitoids alone is often insufficient to keep their populations under control; although it could be improved through conservation biological control (see case study 6) and applied chemical ecology methods (see below). Additionally, the prospects for integrating biological control with other sustainable control methods could improve stink bug control efficacy (Figure 1). Below, we review prospective methods for improving the efficacy of egg parasitoids and, when available, parasitoids attacking adults by using a chemical ecology approach. Also, we evaluate their possible integration with other sustainable methods targeting stink bugs.

Chemical ecology for manipulating parasitoid behaviour

In recent years, the development of semiochemical-based tactics has increased greatly, as they are considered efficient tools for manipulating insect behaviour with the aim of enhancing the biological control of crop pests (Wajnberg & Colazza, 2013). Feeding and/or oviposition by herbivorous insects induce changes in a plant's ecophysiological traits and their emission of volatile organic compounds (VOCs), either as herbivore-induced plant volatiles (HIPVs) or oviposition-induced plant volatiles (OIPVs) (Hilker & Fatouros, 2015; Pashalidou et al., 2015). Changes in VOC emission can occur locally and/or systemically and at either quantitative or qualitative levels (Dicke & van Loon, 2000; Martinez et al., 2013). Emission of HIPVs or OIPVs might alert the neighbouring plants of the same or other species (Baldwin & Schultz, 1983; Arimura et al., 2000; Karban & Maron, 2002) and/or act as an indirect plant defence by recruiting natural enemies of the herbivorous insects (Meiners & Peri, 2013; Hilker & Fatouros, 2015; Pashalidou et al., 2015). Many parasitoids and predators are known to respond to HIPVs or OIPVs during their host/prey location behaviour and can discriminate between volatiles produced by undamaged or infested plants (D'Alessandro & Turlings, 2006; Hare, 2011).

Egg parasitoids in systems involving stink bugs optimize their foraging behaviour by efficiently exploiting OIPVs (Conti & Colazza, 2012). OIPVs reliably indicate the presence of host egg masses and are produced by plants in large quantities, making them easily detectable (Fatouros et al., 2008; Conti & Colazza, 2012). For example, *Tr. basalis* (Colazza et al., 2004a,b; Frati et al., 2017; Salerno et al., 2019) and *Te. podisi* (Blassioli Moraes et al., 2005, 2009; Michereff et al., 2011) are attracted to leguminous plants infested by *N. viridula* or *E. heros*, respectively. By providing reliable information to female egg parasitoids on the presence of suitable target hosts, OIPVs increase wasp recruitment on the host-infested plants, likely increasing parasitism rates.

Feeding and oviposition by the zoophytophagous predator *P. maculiventris* also induce the emission of VOCs in *Vicia faba* L. (Fabaceae) plants that attract *Te. podisi* females (Martorana et al., 2019). Interestingly, both *Tr. basalis* (Martorana et al., 2017; Rondoni et al., 2017) and *Te. podisi* (Martorana et al., 2019) show specificity in their response to OIPVs emitted by infested plants, as neither species are attracted by plants on which the alien *H. halys* had fed and oviposited. This lack of response, probably due to the absence of a history of coevolution between the interacting species, might allow egg parasitoids to optimize their time and energy budgets by exploiting cues only from suitable (coevolved) hosts (Martorana et al., 2017, 2019).

Stink bug activity may also leave short-range cues that are exploited by egg parasitoids once they have landed on plants. For instance, Tr. brochymenae shows an intense egg foraging behaviour on the leaf surface, exploiting substrate-borne chemical cues emitted by brassicaceous plants as a consequence of feeding and oviposition of M. histrionica (Conti et al., 2010). Several egg parasitoids are also able to detect chemical traces released by stink bugs and retained by the plant epicuticular waxes (Conti et al., 2003; Colazza et al., 2009; Frati et al., 2013; Boyle et al., 2020). Trissolcus basalis females were shown to discriminate traces left by N. viridula females from those left by males, due to the absence of *n*-nonadecane, a cuticular hydrocarbon present in N. viridula males but absent in females (Colazza et al., 2007). In doing so, wasps can restrict their searching behaviour to an area of the plant where host eggs are more likely to be found.

Laboratory studies that demonstrated the role of HIPVs, OIPVs, and contact cues in mediating host searching behaviour of egg parasitoids have opened up new opportunities for developing strategies for herbivorous stink bug control. However, the application of semiochemical-based

techniques in the field is often difficult and provides results that are sometimes inconsistent. Several potential limitations and risks have been reported in the literature, including a potential increase in intraguild predation on natural enemies that in turn induces a reduction in the top-down control of the pests (Poelman & Kos, 2016; Peri et al., 2018). These techniques are applied in complex environments in which plants are subjected to various biotic and abiotic stresses and in which several intra- and inter-specific interactions can occur simultaneously. For example, the chewing damage from Sitona lineatus (L.) (Coleoptera: Curculionidae), whose adults feed on leaves and larvae feed on roots, induces a change in the OIPV profile emitted by V. faba plants that are concurrently infested by N. viridula, which in turn reduces the recruitment of the latter's egg parasitoid Tr. basalis (Moujahed et al., 2014). Similar disruptive effects on host egg location by parasitoids were observed in the presence of concurrent infestations of both alien (H. halys) and local (N. viridula or P. maculiventris) stink bugs (Martorana et al., 2017, 2019). Moreover, changes in the volatile blend emitted by plants infested by stink bugs that resulted in modification of egg parasitoid searching behaviour were also observed as a result of abiotic factors, such as water stress (Salerno et al., 2017).

To manipulate parasitoid behaviour on crops, semiochemicals can be applied directly onto the plants by spraying or by using slow-release dispensers. Their emission can be induced by applying elicitors such as plant hormones (JA, methyl jasmonate, cis-jasmone, and SA) or plants themselves could be genetically engineered to increase their emission of HIPVs (Blassioli Moraes et al., 2013; Colazza et al., 2013; Simpson et al., 2013; Peri et al., 2018). However, few examples are reported in the literature on parasitoids of stink bugs, and the results are somewhat contradictory. Positive results were obtained in attracting the tachinid parasitoid Gymnosoma rotundatum (L.) to persimmon orchards using methyl (E,E,Z)-2,4,6-decatrienoate, the aggregation pheromone of the brownwinged green stink bug *Plautia stali* Scott (Pentatomidae) (Jang & Park, 2010; Jang et al., 2011). Positive results were also obtained in soybean crops using the aggregation pheromone of Riptortus pedestris (= clavatus) (Fabricius) (Hemiptera: Alydidae), which led to an increased abundance of its egg parasitoids, Ooencyrtus nezarae Ishii (Encyrtidae) and Gryon japonicum (Ashmead) (Scelionidae) (Lim & Mainali, 2013). Similarly, application of a racemic mixture of methyl 2,6,10-trimethyltridecanoate, a component of E. heros pheromone, attracted scelionid egg parasitoids (Borges et al., 1998). Conversely, although an increase in parasitoid recruitment in soybean fields was observed by using slow-release dispensers impregnated

with (E)-2-hexenal, a compound present in the metathoracic glands of E. heros (Laumann et al., 2007), a more indepth study demonstrated only early parasitoid recruitment but not a sustained increase in abundance (Vieira, 2010). Moreover, the application of cis-jasmone in soybean plants had positive effects on the foraging behaviour of scelionid stink bug egg parasitoids, leading to an increase in their abundance (Blassioli Moraes et al., 2009; Vieira, 2010), but this did not lead to an improvement of parasitism rate (Vieira et al., 2013). Parasitoid recruitment using semiochemicals can be affected by the experience of the foraging females, for example when the hosts are scarce or lacking. Without the reward of suitable host eggs, scelionid parasitoid females were shown to reduce their responses to semiochemicals due to habituation (Peri et al., 2006, 2016; Abram et al., 2017a). Additionally, most parasitoids are more dependent on other cues than those from the host-plant complex, such as floral odours from nectar plants. Therefore, manipulating the habitat by creating ecological infrastructures, e.g., using companion flowering plants such as buckwheat that serve as feeding sites for recruitment of stink bug egg parasitoids, might increase parasitism efficacy (see case study 6). The integration of semiochemical-based manipulation of parasitoid behaviour and habitat manipulation, e.g., the attract-andreward strategy (Khan et al., 2008; Simpson et al., 2011), might limit the possible negative effects of parasitoid recruitment in the absence of hosts.

Chemical ecology for manipulating stink bug behaviour

The use of semiochemicals against stink bugs might be combined with biological control to increase pest control efficacy within organic farming systems. The chemical ecology of stink bugs is characterized by a wide array of signals that drive their behaviour at both intra- and inter-specific levels. Although sex and aggregation pheromones of stink bugs are commonly used as lures for trapping (Borges et al., 1998, 2011; Leskey & Nielsen, 2018), few studies have reported the application of other semiochemicals.

Stink bugs exploit host plant volatiles to find their feeding and oviposition substrates (Martinez et al., 2013; Guarino et al., 2017a; Weber et al., 2018). The types of semiochemicals they use depend on their feeding habits and their level of polyphagy. Polyphagous species respond to blends of common plant volatiles in precise proportions, whereas monophagous and oligophagous species are more attracted by key plant-specific compounds (Guarino et al., 2017a). For example, *Eurydema pulchrum* Westwood (Pentatomidae) positively responds to volatiles from host plants belonging to different families, such as *Brassica oleracea* L., *Raphanus sativus* L. (Brassicaceae) and

Solanum lycopersicum L. (Solanaceae) (Rather et al., 2010). In contrast, *B. hilaris*, a specialist of Brassicaceae that exhibits a strong preference for certain species at the seedling stage (Huang et al., 2014; Guarino et al., 2017b), preferred *B. oleracea* and *B. napus* over *B. carinata*. This behaviour seems to be mediated by a novel diterpene hydrocarbon, still uncharacterized, emitted by *B. oleracea* and *B. napus* but missing in *B. carinata* blends (Guarino et al., 2018).

Stink bug preference for certain host plant species could be exploited for pest management using trap crops, which rely on an attractive host plant to arrest the pest and/or to concentrate it in certain zones of the field, where it can be eliminated (Hokkanen, 1991). Trap crops have shown the potential to effectively manage stink bugs in conventional and organic crop production systems (Todd & Schumann, 1988; Tillman, 2006). For example, the use of sorghum as a trap crop for N. viridula reduced insecticide applications in cotton, and black mustard used as a trap in sweet corn reduced pest damage by 22% (Rea et al., 2002; Tillman, 2006). Similarly, soybean and a mixture of triticale, sorghum, and sunflower were successfully used as trap crops in cotton fields for Euschistus servus (Say), C. hilaris, and N. viridula (Mizell et al., 2008; Tillman et al., 2015). The efficacy of the trap crop technique has also been demonstrated by combining it with pheromone-baited traps for suppressing E. servus attacking cotton. Such combinations of attractant stimuli increased the trap crop effectiveness in terms of numbers of attracted individuals and also consistently reduced the dispersal of E. servus in cotton fields (Tillman & Cottrell, 2012) and of P. stali in kaki persimmon orchards (Yamanaka et al., 2011). Recently, trap crops have been used for the management of invasive stink bug species. For example, H. halys has been controlled in pepper fields by using sunflower alone or in combination with sorghum as trap crops (Soergel et al., 2015; Mathews et al., 2017).

The use of trap crops for stink bug control may be particularly useful because there are few biologically based strategies that are available for the suppression of stink bugs in small organic farms or homeowner gardens (Mathews et al., 2017). Trap cropping might lead to (1) reduction in the use of pesticides, thus reducing the cost of pest control; (2) reduction in the selective pressure for development of pest resistance; and (3) preservation of natural enemies (Gordon et al., 2017). However, none of the cases reported have yet led to commercial implementation (Weber et al., 2018).

In order to manipulate pest behaviour more effectively, the trap crop technique can also be implemented by using a stimulus that is deterrent or repellent in nature, applied on the crop, leading to a push-pull strategy (Pickett et al., 2014). Information on the successful application of this strategy against stink bugs is scarce. However, a study carried out by Zhang et al. (2014) showed that several plant extracts obtained from clove, lemongrass, spearmint oil, and ylang-ylang plants acted as repellents toward *H. halys* and could be candidates as 'push stimuli' for this pest. The repellent stimuli could be used as extracts in dispensers or as intercropping plants to repel the pest that simultaneously could be attracted by a trap crop that serves as a 'pull stimulus'.

Intercropping with a non-host plant can directly reduce the damage to cash crops by masking the host from the pests, as some plants might have a repellent effect on herbivores (Smith & McSorley, 2000). In stink bug management, intercropping soybean with millet, cow pea, and groundnut reduced damage by *N. viridula* by 50% compared with a system containing soybean and millet only (Sastawa et al., 2004).

Future research on behavioural manipulation of stink bugs will be able to take advantage of new genomic and transcriptomic technology (Sparks et al., 2014) to identify the genes responsible for their pheromone production and behavioural responses to various odours. The objective will be, for example, to use plants or yeasts as 'factories' of stink bug attractants that can be used as 'dead end' trap crops (Møldrup et al., 2012).

Prospects for additional sustainable control methods

Additional sustainable control methods that could be considered in combination with parasitoids within organic farming systems include implementation of vibratory cues to disrupt stink bug sexual communication (Laumann et al., 2017; Čokl et al., 2019; Polajnar et al., 2019), direct plant resistance (Rondoni et al., 2018; Serteyn et al., 2020), exploitation of stink bug entomopathogens (Hajek et al., 2018), exclusion netting for fruit orchards (Candian et al., 2018), attract-and-kill methods (Morrison et al., 2019), and a novel mode of application of the sterile insect technique that combines the release of sterilized stink bugs with sterile parasitoids for pest eradication (Horrocks et al., 2020). Several of these possible methods are discussed below.

The reproductive behaviour in many stink bug species is mediated by multimodal communication, based on long-range attraction on a common substrate via chemical signals and the involvement of combinations of vibratory, chemical, and visual cues during calling and courtship behaviours (Čokl et al., 2019). Vibratory signals have been proposed as tools for mass trapping, using a bi-modal trap based on the aggregation pheromone and vibratory signal (Polajnar et al., 2019), and for mating disruption (Laumann et al., 2017). Considering that females of *Te. podisi* specifically respond to stink bug vibratory signals for

spatial orientation, such signals might also be exploited for parasitoid recruitment on infested crops (Laumann et al., 2017).

Host plant (*V. faba*) resistance to stink bugs has been investigated recently at the biological, molecular, and behavioural level (Rondoni et al., 2018; Serteyn et al., 2020). Rondoni et al. (2018) showed that plants recognize *H. halys* oviposition as a warning signal and respond by activating defences against *H. halys* nymphs, resulting in reduced size of juvenile offspring. This is possibly related to the priming of JA-dependent genes, including a cysteine proteinase inhibitor. The use of proteinase inhibitors in agriculture can be implemented through the development of transgenic plants, the use of recombinant microorganisms, or production of biopesticides (reviewed by Rodríguez-Sifuentes et al., 2020).

Conclusions

This review summarises more than a century of research on the biological control of invasive stink bugs. Invasive stink bugs are difficult to control, especially when they have already colonized entire regions and population outbreaks have occurred. A new pest often spreads rapidly after its introduction, and effective management is often subject to delays while available short-term tools are identified and a classical biological control program is initiated. Pre-emptive biological control is an innovative strategy for appropriate situations to reduce delays in research and implementation, by preparing in advance for the arrival of a pest. When a pest has already established in a new region, classical biological control can be an effective means of reducing the reservoir of populations in the wider landscape, thus reducing immigration to crops. However, classical biological control of stink bugs has limitations that have been addressed here and have been recently analysed by Abram et al. (2020). Using stage-structured matrix models, these authors showed that stink bug mortality at the adult stage would be expected to provide a greater reduction of population growth than mortality at the egg stage and that a multi-tactic approach targeting different stages would be much more effective in reducing populations below the economic injury level. Here, we show that egg parasitoids are the most common natural enemies of stink bugs globally, and for this reason they have been the most commonly studied parasitoids of stink bugs. However, parasitoids attacking adults also provide interesting perspectives, although these are only known for a few invasive stink bug species.

The combination of classical biological control followed by conservation methods and integrated with other sustainable control methods targeting the various instars should provide interesting prospects for the improvement of stink bug control efficacy (Figure 1). Conservation biological control, through a provision of suitable nectar-producing flowering plants, can attract and improve the efficacy of both parasitoids attacking adults (Tillman & Carpenter, 2014; Tillman, 2017) and egg parasitoids (Foti et al., 2017, 2019). The application of semiochemicals on crops or inducing them in plants by applying elicitors might also facilitate recruitment of parasitoid species attacking adults (Jang & Park, 2010; Jang et al., 2011) or eggs (Borges et al., 1998; Lim & Mainali, 2013), although a consequent improvement of parasitism has yet to be demonstrated with these applications (Blassioli Moraes et al., 2009; Vieira, 2010; Vieira et al., 2013). The attract-and-reward strategy, which combines behavioural manipulation and habitat manipulation, might further improve parasitoid efficacy (Khan et al., 2008; Simpson et al., 2011). Semiochemicals can be used to manipulate not only natural enemies but also stink bug behaviour by using either trap crops (Todd & Schumann, 1988; Tillman, 2006; Nielsen et al., 2016) or push-pull strategies (Pickett et al., 2014). Moreover, future tactics might exploit vibratory cues to disrupt stink bug sexual communication (Laumann et al., 2017; Čokl et al., 2019; Polajnar et al., 2019) or direct plant resistance (Rondoni et al., 2018). Ultimately, as most research has been conducted only under laboratory conditions, there is an urgent need for field validation of these data before effective implementation can become a reality.

Author Contribution Statement

All authors contributed to the writing of the paper. EC and GR conceived the idea of the article and designed the layout. EC coordinated the paper and wrote the general parts. GA and BB wrote about the pre-emptive classical biological control of *H. halys* in New Zealand. FC and RAL focused on the biocontrol of stink bugs in South America. SC and MR wrote about the conservation biocontrol of *N. viridula*. EP, SG, and EW contributed with the chemical ecology of stink bugs and egg parasitoids. RS and MG wrote about the classical biological control of *B. hilaris*. LM and LT focused on the biocontrol of *H. halys* in Europe. CRS and KH focused on the biocontrol of *H. halys* in North America.

References

Abram PK, Cusumano A, Abram K, Colazza S & Peri E (2017a) Testing the habituation assumption underlying models of parasitoid foraging behavior. PeerJ 5: e3097.

- Abram PK, Doyon J, Brodeur J, Gariépy TD & Boivin G (2015) Susceptibility of *Halyomorpha halys* (Hemiptera: Pentatomidae) eggs to different life stages of three generalist predators. Canadian Entomologist 147: 222–226.
- Abram PK, Gariepy TD, Boivin G & Brodeur J (2014) An invasive stink bug as an evolutionary trap for an indigenous egg parasitoid. Biological Invasions 16: 1387–1395.
- Abram PK, Hoelmer KA, Acebes-Doria A, Andrews H, Beers EH et al. (2017b) Indigenous arthropod natural enemies of the invasive brown marmorated stink bug in North America and Europe. Journal of Pest Science 90: 1009–1020.
- Abram PK, Hueppelsheuser T, Acheampong S, Clarke P, Douglas H & Gariepy TD (2017c) Evidence of established brown marmorated stink bug populations in British Columbia, Canada. Journal of the Entomological Society of British Columbia 114: 83–86.
- Abram PK, Mills NJ & Beers EH (2020) Review: classical biological control of invasive stink bugs with egg parasitoids what does success look like? Pest Management Science 76: 1980–1992.
- Abram PK, Talamas EJ, Acheampong S, Mason PG & Gariepy TD (2019) First detection of the samurai wasp, *Trissolcus japonicus* (Ashmead) (Hymenoptera, Scelionidae), in Canada. Journal of Hymenoptera Research 68: 29–36.
- Ahuja B, Kalyan R, Ahuja U, Singh S, Sundria M & Dhandapani A (2008) Integrated management strategy for painted bug, Bagrada hilaris (Burm.) inflicting injury at seedling stage of mustard (Brassica juncea) in arid western Rajasthan. Pesticide Research Journal 20: 48–51.
- Aquino MFS, Sujii ER, Borges M, Blassioli Moraes MC & Laumann RA (2019) Diversity of stink bug adults and their parasitoids in soybean crops in Brazil: influence of a latitudinal gradient and insecticide application intensity. Environmental Entomology 48: 105–113.
- Arakawa R & Namura Y (2002) Effects of temperature on development of three *Trissolcus* spp. (Hymenoptera: Scelionidae), egg parasitoids of the brown marmorated stink bug, *Halyomorpha halys* (Hemiptera: Pentatomidae). Entomological Science 5: 215–218.
- Arimura GI, Ozawa R, Shimoda T, Nishioka T, Boland W & Takabayashi J (2000) Herbivory-induced volatiles elicit defence genes in Lima bean leaves. Nature 406: 512–515.
- Avila GA & Charles JG (2018) Modelling the potential geographic distribution of *Trissolcus japonicus*: a biological control agent of the brown marmorated stink bug, *Halyomorpha halys*. BioControl 63: 505–518.
- Baldwin IT & Schultz JC (1983) Rapid changes in tree leaf chemistry induced by damage: evidence for communication between plants. Science 221: 277–279.
- Barratt BIP & Ehlers GAC (2017) Impacts of exotic biological control agents on non-target species and biodiversity: evidence, policy and implications. Environmental Pest Management: Challenges for Agronomists, Ecologists, Economists and Policymakers (ed. by M Coll & E Wajnberg), pp. 325–346. John Wiley & Sons, Oxford, UK.

- Bergmann E, Venugopal PD, Martinson HM, Raupp M & Shrewsbury P (2016) Host plant use by the invasive *Halyomorpha halys* (Stål) on woody ornamental trees and shrubs. PLoS ONE 11: e0149975.
- Bertoldi V, Rondoni G, Brodeur J & Conti E (2019) An egg parasitoid efficiently exploits cues from a coevolved host but not those from a novel host. Frontiers in Physiology 10: 746.
- Blassioli Moraes MC, Borges M & Laumann RA (2013) The application of chemical cues in arthropod pest management for arable crops. Chemical Ecology of Insect Parasitoids (ed. by E Wajnberg & S Colazza), pp. 225–244. Wiley-Blackwell, Oxford, UK.
- Blassioli Moraes MC, Laumann RA, Pareja M, Sereno FT, Michereff MF et al. (2009) Attraction of the stink bug egg parasitoid *Telenomus podisi* to defence signals from soybean activated by treatment with cis-jasmone. Entomologia Experimentalis et Applicata 131: 178–188.
- Blassioli Moraes MC, Laumann RA, Sujii ER, Pires C & Borges M (2005) Induced volatiles in soybean and pigeon pea plants artificially infested with the neotropical brown stink bug, *Euschistus heros*, and their effect on the egg parasitoid, *Telenomus podisi*. Entomologia Experimentalis et Applicata 115: 227–237.
- Borges M, Blassioli Moraes MC, Peixoto MF, Pires CSS, Sujii ER & Laumann RA (2011) Monitoring the Neotropical brown stink bug *Euschistus heros* (F.) (Hemiptera: Pentatomidae) with pheromone-baited traps in soybean fields. Journal of Applied Entomology 135: 68–80.
- Borges M, Schmidt FGV, Sujii ER, Medeiros MA, Mori K et al. (1998) Field responses of stink bugs to the natural and synthetic pheromone of the Neotropical brown stink bug, *Euschistus heros* (Heteroptera: Pentatomidae). Physiological Entomology 23: 202–207.
- Bosco L, Moraglio ST & Tavella L (2018) Halyomorpha halys, a serious threat for hazelnut in newly invaded areas. Journal of Pest Science 91: 661–670.
- Botch PS & Delfosse ES (2018) Host-acceptance behavior of *Trissolcus japonicus* (Hymenoptera: Scelionidae) reared on the invasive *Halyomorpha halys* (Heteroptera: Pentatomidae) and nontarget species. Environmental Entomology 47: 403–411.
- Boyle SM (2017) Novel Techniques for Evaluating the Potential Host Range of Candidate Biological Control Agent *Trissolcus japonicus* (Hymenoptera: Platygastridae). MSc Thesis, University of Delaware, Newark, DE, USA.
- Boyle S, Weber D, Hough-Goldstein J & Hoelmer KA (2020) Host kairomones influence searching behavior of *Trissolcus japonicus* (Hymenoptera: Scelionidae), a parasitoid of *Halyomorpha halys* (Heteroptera: Pentatomidae). Environmental Entomology 49: 15–20.
- Bradshaw CJ, Leroy B, Bellard C, Roiz D, Albert C et al. (2016) Massive yet grossly underestimated global costs of invasive insects. Nature Communication 7: 12986.
- Buffington ML, Talamas EJ & Hoelmer KA (2018) Team *Trissolcus*: integrating taxonomy and biological control to combat the brown marmorated stink bug. American Entomologist 64: 224–232.

44 Conti et al.

- Bulgarini G, Badra Z & Maistrello L (2019) Predatory ability of wild generalist predators against eggs and first instar nymphs of *Halyomorpha halys*. IOBC/WPRS Bulletin 146: 106–107.
- Bundy CS, Esquivel JF, Panizzi AR, Eger JE, Davis JA & Jones WA (2018a) *Piezodorus guildinii* (Westwood). Invasive Stink Bugs and Related Species (Pentatomoidea): Biology, Higher Systematics, Semiochemistry, and Management (ed. by JE McPherson), pp. 425–451. CRC Press, Boca Raton, FL, USA.
- Bundy CS, Perring TM, Reed DA, Palumbo JC, Grasswitz TR & Jones WA (2018b) *Bagrada hilaris* (Burmeister). Invasive Stink Bugs and Related Species (Pentatomoidea): Biology, Higher Systematics, Semiochemistry, and Management (ed. by JE McPherson), pp. 205–241. CRC Press, Boca Raton, FL, USA.
- Candian V, Pansa MG, Briano R, Peano C, Tedeschi R & Tavella L (2018) Exclusion nets: a promising tool to prevent *Haly-omorpha halys* from damaging nectarines and apples in NW Italy. Bulletin of Insectology 71: 21–30.
- Castiglioni E, Chiaravalle W, Cristino M, Silva H, Ribeiro A & Corrêa-Ferreira BS (2007) Implementação de um programa de controle biológico de *Piezodorus guildinii* (Hemiptera, Pentatomidae) com *Telenomus podisi* (Hymenoptera, Scelionidae), no Uruguay. Proceedings of the X SICONBIOL, p. 235. Sociedade Entomológica do Brasil, Brasília, DF, Brazil.
- Castracani C, Bulgarini G, Giannetti D, Spotti FA, Maistrello L et al. (2017) Predatory ability of the ant *Crematogaster scutellaris* on the brown marmorated stink bug *Halyomorpha halys*. Journal of Pest Science 90: 1181–1190.
- Cesari M, Maistrello L, Piemontese L, Bonini R, Dioli P et al. (2017) Genetic diversity of the brown marmorated stink bug *Halyomorpha halys* in the invaded territories of Europe and its patterns of diffusion in Italy. Biological Invasions 20: 1073–1092.
- Chacko M & Katiyar R (1961) *Hadrophanurus karnalensis* sp. n. (Hymenoptera: Scelionidae), a parasite of *Bagrada cruciferarum* Kirkaldy (Hemiptera: Pentatomidae). Systematic Entomology 30: 161–163.
- Charles JG, Avila GA, Hoelmer KA, Hunt S, Gardner-Gee R et al. (2019) Experimental assessment of the biosafety of *Trissolcus japonicus* in New Zealand, prior to the anticipated arrival of the invasive pest *Halyomorpha halys*. BioControl 64: 367–379.
- Cingolani MF (2012) Parasitismo de Huevos de *Piezodorus guildinii* (Hemiptera: Pentatomidae) por *Trissolcus basalis y Telenomus podisi* (Hymenoptera: Scelionidae) en el Noreste de la Provincia de Buenos Aires. PhD Dissertation, Universidad Nacional de La Plata, Buenos Aires, Argentina.
- Cingolani MF, Greco NM & Liljesthröm GG (2014a) Egg parasitism of *Piezodorus guildinii* and *Nezara viridula* (Hemiptera: Pentatomidae) in soybean, alfalfa and red clover. Revista FCA UNCUYO 46: 15–27.
- Cingolani MF, Greco NM & Liljesthröm GG (2014b) Effect of *Telenomus podisi, Trissolcus urichi*, and *Trissolcus basalis* (Hymenoptera: Platygastridae) age on attack of *Piezodorus guildinii* (Hemiptera: Pentatomidae) eggs. Environmental Entomology 43: 377–383.
- Claerebout S, Haye T, Ólafsson E, Pannier É & Bultot J (2018) Premières occurrences de *Halyomorpha halys* (Stål, 1855) pour

- la Belgique et actualisation de sa répartition en Europe (Hemiptera: Heteroptera: Pentatomidae). Bulletin de la Société Royale Belge d'Entomologie 154: 205–227.
- Cock MJW, Murphy ST, Kairo MTK, Thompson E, Murphy RJ & Francis AW (2016) Trends in the classical biological control of insect pests by insects: an update of the BIOCAT database. BioControl 61: 349–363.
- Čokl A, Blassioli-Moraes MC, Laumann RA, Žunič A & Borges M (2019) Stinkbugs: multisensory communication with chemical and vibratory signals transmitted through different media. Biotremology: Studying Vibrational Behavior (ed. by PSM Hill, R Lakes-Harlan, V Mazzoni, PM Narins, M Virant-Doberlet & A Wessel), pp. 91–122. Springer, Cham, Switzerland
- Colazza S, Aquila G, De Pasquale C, Peri E & Millar JG (2007) The egg parasitoid *Trissolcus basalis* uses n-nonadecane, a cuticular hydrocarbon from its stink bug host *Nezara viridula*, to discriminate between female and male hosts. Journal of Chemical Ecology 33: 1405–1420.
- Colazza S, Fucarino A, Peri E, Salerno G, Conti E & Bin F (2004a) Insect oviposition induces volatile emission in herbaceous plants that attracts egg parasitoids. Journal of Experimental Biology 207: 47–53.
- Colazza S, Giangiuliani G & Bin F (1996) Fortuitous introduction and successful establishment of *Trichopoda pennipes* F.: adult parasitoid of *Nezara viridula* (L.). Biological Control 6: 409–411.
- Colazza S, Lo Bue M, Lo Giudice D & Peri E (2009) The response of *Trissolcus basalis* to footprint contact kairomones from *Nezara viridula* females is mediated by leaf epicuticular waxes. Naturwissenschaften 96: 975–981.
- Colazza S, McElfresh JS & Millar JG (2004b) Identification of volatile synomones, induced by *Nezara viridula* feeding and oviposition on bean spp., that attract the egg parasitoid *Trissolcus basalis*. Journal of Chemical Ecology 30: 945–964.
- Colazza S, Peri E & Cusumano A (2013) Application of chemical cues in arthropod pest management for orchards and vineyards. Chemical Ecology of Insect Parasitoids (ed. by E Wajnberg & S Colazza), pp. 245–265. Wiley-Blackwell, Oxford, UK.
- Conti E & Colazza S (2012) Chemical ecology of egg parasitoids associated with true bugs. Psyche 2012: 651015.
- Conti E, Salerno G, Bin F, Williams HJ & Vinson SB (2003) Chemical cues from *Murgantia histrionica* eliciting host location and recognition in the egg parasitoid *Trissolcus brochymenae*. Journal of Chemical Ecology 29: 115–130.
- Conti E, Salerno G, Leombruni B, Frati F & Bin F (2010) Short-range allelochemicals from a plant–herbivore association: a singular case of oviposition-induced synomone for an egg parasitoid. Journal of Experimental Biology 213: 3911–3919.
- Cornelius ML, Dieckhoff C, Hoelmer KA, Olsen RT, Weber DC et al. (2016a) Biological control of sentinel egg masses of the exotic invasive stink bug *Halyomorpha halys* (Stål) in Mid-Atlantic USA ornamental landscapes. Biological Control 103: 11–20.
- Cornelius ML, Dieckhoff C, Vinyard BT & Hoelmer KA (2016b)

 Parasitism and predation on sentinel egg masses of the brown

- marmorated stink bug (Hemiptera: Pentatomidae) in three vegetable crops: importance of dissections for evaluating the impact of native parasitoids on an exotic pest. Environmental Entomology 45: 1536–1542.
- Corrêa-Ferreira BS (2002) Trissolcus basalis para o controle de percevejos da soja. Controle Biológico no Brasil: Parasitóides e Predadores (ed. by JRP Parra, PSM Botelho, BS Corrêa-Ferreira & JM Bento), pp. 449–476. Manole, São Paulo, Brazil.
- Corrêa-Ferreira BS, Alexandre TM, Pellizzaro EC, Moscard F & Bueno AF (2010) Práticas de manejo de pragas utilizadas na soja e seu impacto sobre a cultura. Embrapa Soja Circular Técnica 78: 16.
- Corrêa-Ferreira BS, Domit LA, Morales L & Guimarães RC (2000) Integrated soybean pest management in micro river basins in Brazil. Integrated Pest Management Review 5: 75–80.
- Corrêa-Ferreira BS & Moscardi F (1995) Seasonal occurrence and host spectrum of egg parasitoids associated with soybean stink bugs. Biological Control 5: 196–202.
- Costi E, Haye T & Maistrello L (2018) Surveying native egg parasitoids and predators of the invasive *Halyomorpha halys* in Northern Italy. Journal of Applied Entomology 143: 299–307.
- Crouzel IS & Saini ED (1983) Importación de *Trissolcus basalis* (Wollaston) (Hym. Scelionidae) en la Argentina para el control biológico de *Nezara viridula* (L.) (Hem. Pentatomidae). Revista de la Sociedad Entomológica Argentina 42: 257–260.
- CSO (2019) Estimation of Damage from Brown Marmorated Stink Bug and Pathologies Related to Climate Change. https:// www.csoservizi.com accessed 30-05-2020.
- D'Alessandro M & Turlings TCJ (2006) Advances and challenges in the identification of volatiles that mediate interactions among plants and arthropods. Analyst 131: 24–32.
- Dicke M & van Loon JJA (2000) Multitrophic effects of herbivore-induced plant volatiles in an evolutionary context. Entomologia Experimentalis et Applicata 97: 237–249.
- Dieckhoff C, Tatman K & Hoelmer KA (2017) Natural biological control of *Halyomorpha halys* (Hemiptera: Pentatomidae) by native egg parasitoids a multi-year survey in northern Delaware. Journal of Pest Science 90: 1143–1158.
- Duthie C (2012) Risk Analysis of *Halyomorpha halys* (Brown Marmorated Stinkbug) on All Pathways. Ministry for Primary Industries. https://www.mpi.govt.nz/dmsdocument/3943/send (accessed June 2020).
- Ehlers GAC, Caradus JR & Fowler SV (2019) The regulatory process and costs to seek approval for the development and release of new biological control agents in New Zealand. BioControl 65: 1–12.
- EPA (2018) Application and Approval to Release *Trissolcus japonicus* into New Zealand. https://www.epa.govt.nz/search/SearchForm?Search=Trissolcus+japonicus (accessed March 2019).
- Esquivel JF, Musolin DL, Jones WA, Rabitsch W, Greene JK et al. (2018) *Nezara viridula* (L.). Invasive Stink Bugs and Related Species (Pentatomoidea): Biology, Higher Systematics, Semiochemistry, and Management (ed. by JE McPherson), pp. 351–423. CRC Press, Boca Raton, FL, USA.

- Fatouros NE, Dicke M, Mumm R, Meiners T & Hilker M (2008) Foraging behaviour of egg parasitoids exploiting chemical information. Behavioural Ecology 19: 677–689.
- Faúndez EI, Lüer A, Cuevas AG, Rider DA & Valdebenito P (2016) First record of the painted bug *Bagrada hilaris* (Burmeister, 1835) (Heteroptera: Pentatomidae) in South America. Arquivos Entomolóxicos 16: 175–179.
- Faúndez EI (2018) From agricultural to household pest: the case of the painted bug *Bagrada hilaris* (Burmeister) (Heteroptera: Pentatomidae) in Chile. Journal of Medical Entomology 55: 1365–1368.
- Faúndez EI & Rider DA (2017) The brown marmorated stink bug Halyomorpha halys (Stål, 1855) (Heteroptera: Pentatomidae) in Chile. Arquivos Entomolóxicos 17: 305–307.
- Felipe-Victoriano M, Talamas E & Sánchez-Peña SR (2019) Scelionidae (Hymenoptera) parasitizing eggs of *Bagrada hilaris* (Hemiptera, Pentatomidae) in Mexico. Journal of Hymenoptera 73: 143–152.
- Fogain R & Graff S (2011) First records of the invasive pest, *Halyomorpha halys* (Hemiptera: Pentatomidae), in Ontario and Quebec. Journal of the Entomological Society of Ontario 142: 45–48.
- Foti MC, Peri E, Wajnberg E, Colazza S & Rostás M (2019) Contrasting olfactory responses of two egg parasitoids to buckwheat floral scent are reflected in field parasitism rates. Journal of Pest Science 92: 747–756.
- Foti MC, Rostás M, Peri E, Park FC, Slimani T et al. (2017) Chemical ecology meets conservation biological control: identifying plant volatiles as predictors of floral resource suitability for an egg parasitoid of stink bugs. Journal of Pest Science 90: 299–310.
- Fraga DF, Parker J, Busoli AC, Hamilton GC, Nielsen AL & Rodriguez-Saona C (2017) Behavioral responses of predaceous minute pirate bugs to tridecane, a volatile emitted by the brown marmorated stink bug. Journal of Pest Science 90: 1107–1118.
- Frati F, Cusumano A, Conti E, Colazza S, Peri E et al. (2017) Foraging behaviour of an egg parasitoid exploiting plant volatiles induced by pentatomids: the role of adaxial and abaxial leaf surfaces. PeerJ 5: e3326.
- Frati F, Salerno G & Conti E (2013) Cabbage waxes affect *Trissol-cus brochymenae* response to short-range synomones. Insect Science 20: 753–762.
- Gariepy TD & Talamas EJ (2019) Discovery of *Trissolcus japonicus* (Hymenoptera: Scelionidae) in Ontario, Canada. Canadian Entomologist 151: 824–826.
- Gontijo LM (2019) Engineering natural enemy shelters to enhance conservation biological control in field crops. Biological Control 130: 155–163.
- González-Chang M, Tiwari S, Sharma S & Wratten SD (2019) Habitat management for pest management: limitations and prospects. Annals of the Entomological Society of America 112: 302–317.
- Gordon TL, Haseeb M, Kanga LHB & Legaspi JC (2017) Potential of three trap crops in managing *Nezara viridula* (Hemiptera: Pentatomidae) on tomatoes in Florida. Journal of Economic Entomology 110: 2478–2482.

- Guarino S, Arif MA, Millar JG, Colazza S & Peri E (2018) Volatile unsaturated hydrocarbons emitted by seedlings of *Brassica* species provide host location cues to *Bagrada hilaris*. PLoS ONE 13: e0209870.
- Guarino S, Peri E & Colazza S (2017a) Plant and stink bug interactions at different trophic levels. Stinkbugs (ed. by A Cokl & M Borges), pp. 180–199. CRC Press, Boca Raton, FL, USA.
- Guarino S, Peri E, Colazza S, Luchi N, Michelozzi M & Loreto F (2017b) Impact of the invasive painted bug *Bagrada hilaris* on physiological traits of its host *Brassica oleracea* var *botrytis*. Arthropod-Plant Interactions 11: 649–658.
- Gunn D (1918) The bagrada bug (*Bagrada hilaris*). Bulletin of the Department of Agriculture in South Africa 9: 1–19.
- Hamilton GC, Ahn JJ, Bu W, Leskey TC, Nielsen AL et al. (2018) Halyomorpha halys (Stål). Invasive Stink Bugs and Related Species (Pentatomoidea): Biology, Higher Systematics, Semiochemistry, and Management (ed. by JE McPherson), pp. 243–292. CRC Press, Boca Raton, FL, USA.
- Hare JD (2011) Ecological role of volatiles produced by plants in response to damage by herbivorous insects. Annual Review of Entomology 56: 161–180.
- Haseeb M, Gordon TL, Kanga LH & Legaspi JC (2018) Abundance of natural enemies of *Nezara viridula* (Hemiptera: Pentatomidae) on three cultivars of sweet alyssum. Journal of Applied Entomology 142: 847–853.
- Hayashida R, Bueno AF, Hermel AO, Hirakuri MH, Silva FAC & Samuel R (2018) Euschistus heros (Hemiptera: Pentatomidae) fitness on artificial diets: an approach to optimize mass rearing of Telenomus podisi (Hymenoptera: Platygastridae) for augmentative biological control. Journal of Economic Entomology 111: 1605–1613.
- Haye T, Fischer S, Zhang J & Gariepy T (2015a) Can native egg parasitoids adopt the invasive brown marmorated stink bug, *Halyomorpha halys* (Heteroptera: Pentatomidae), in Europe? Journal of Pest Science 88: 693–705.
- Haye T, Gariepy T, Hoelmer K, Rossi J-P, Streito J-C et al. (2015b) Range expansion of the invasive brown marmorated stinkbug, *Halyomorpha halys*: an increasing threat to field, fruit and vegetable crops worldwide. Journal of Pest Science 88: 665–673.
- Haye T, Moraglio ST, Stahl J, Visentin S, Gregorio T & Tavella L (2020) Fundamental host range of *Trissolcus japonicus* in Europe. Journal of Pest Science 93: 171–182.
- Hajek AE, Solter LF, Maddox JV, Huang W-F, Estep AS et al. (2018) Nosema maddoxi sp. nov. (Microsporidia, Nosematidae), a widespread pathogen of the green stink bug Chinavia hilaris (Say) and the brown marmorated stink bug Halyomorpha halys (Stål). Journal of Eukaryotic Microbiology 65: 315–330.
- Hedstrom C, Lowenstein D, Andrews H, Bai B & Wiman N (2017) Pentatomid host suitability and the discovery of introduced populations of *Trissolcus japonicus* in Oregon. Journal of Pest Science 90: 1169–1179.
- Hemala V & Kment P (2017) First record of *Halyomorpha halys* and mass occurrence of *Nezara viridula* in Slovakia. Plant Protection Science 53: 247–253.

- Herlihy MV, Talamas EJ & Weber DC (2016) Attack and success of native and exotic parasitoids on eggs of *Halyomorpha halys* in three Maryland habitats. PLoS ONE 11: e0150275.
- Hilker M & Fatouros NE (2015) Plant responses to insect egg deposition. Annual Review of Entomology 60: 493–515.
- Hoddle MS, Mace K & Steggall J (2018) Proactive biological control: a cost-effective management option for invasive pests. California Agriculture 72: 1–3.
- Hoebeke ER & Carter ME (2003) Halyomorpha halys (Stål) (Heteroptera: Pentatomidae): a polyphagous plant pest from Asia newly detected in North America. Proceedings of the Entomological Society of Washington 105: 225–237.
- Hokkanen HM (1991) Trap cropping in pest management. Annual Review of Entomology 36: 119–138.
- Horrocks KJ, Avila GA, Holwell GI & Suckling DM (2020) Integrating sterile insect technique with the release of sterile classical biocontrol agents for eradication: is the Kamikaze Wasp Technique feasible? BioControl 65: 257–271.
- Huang TI, Reed DA, Perring TM & Palumbo JC (2014) Host selection behavior of *Bagrada hilaris* (Hemiptera: Pentatomidae) on commercial cruciferous host plants. Crop Protection 59: 7–13.
- Jang SA & Park CG (2010) Gymnosoma rotundatum (Diptera: Tachinidae) attracted to the aggregation pheromone of Plautia stali (Hemiptera: Pentatomidae). Journal of Asia-Pacific Entomology 13: 73–75.
- Jang SA, Cho JH, Park GM, Choo HY & Park CG (2011) Attraction of *Gymnosoma rotundatum* (Diptera: Tachinidae) to different amounts of *Plautia stali* (Hemiptera: Pentatomidae) aggregation pheromone and the effect of different pheromone dispensers. Journal of Asia-Pacific Entomology 14: 119–121.
- Jarrett BJM, Pote J, Talamas E, Gut L & Szucs M (2019) The discovery of *Trissolcus japonicus* (Hymenoptera: Scelionidae) in Michigan. Great Lakes Entomologist 52: 6–11.
- Jones AL, Jennings DE, Hooks CRR & Shrewsbury PM (2014) Sentinel eggs underestimate rates of parasitism of the exotic brown marmorated stink bug, *Halyomorpha halys*. Biological Control 78: 61–66.
- Jones AL, Jennings DE, Hooks CRR & Shrewsbury PM (2017) Field surveys of egg mortality and indigenous egg parasitoids of the brown marmorated stink bug, *Halyomorpha halys*, in ornamental nurseries in the mid-Atlantic region of the USA. Journal of Pest Science 90: 1159–1168.
- Karban R & Maron J (2002) The fitness consequences of interspecific eavesdropping between plants. Ecology 83: 1209–1213.
- Karp DS, Chaplin-Kramer R, Meehan TD, Martin EA, DeClerck F et al. (2018) Crop pests and predators exhibit inconsistent responses to surrounding landscape composition. Proceedings of the National Academy of Sciences of the USA 115: E7863–E7870.
- Kaser JM, Akotsen-Mensah C, Talamas EJ & Nielsen AL (2018) First report of *Trissolcus japonicus* parasitizing *Halyomorpha halys* in North American agriculture. Florida Entomologist 101: 680–683.

- Keane, RM & Crawley, MJ (2002) Exotic plant invasions and the enemy release hypothesis. Trends in Ecology & Evolution, 17: 164–170.
- Kereselidze M, Aleksidze G & Haye T (2018) First record native parasitoid attacking *Halyomorpha halys* (Heteroptera: Pentatomidae) in Georgia. Bulletin of Georgian Academy of Agricultural Sciences 39: 127–129.
- Khan ZR, James DG, Midega CA & Pickett JA (2008) Chemical ecology and conservation biological control. Biological Control 45: 210–224.
- Khan ZR, Midega CA, Hooper A & Pickett J (2016) Push-pull: chemical ecology-based integrated pest management technology. Journal of Chemical Ecology 42: 689–697.
- Lara JR, Pickett CH, Kamiyama MT, Figueroa S, Romo M et al. (2019) Physiological host range of *Trissolcus japonicus* in relation to *Halyomorpha halys* and other pentatomids from California. BioControl 64: 513–528.
- Laumann RA, Aquino MFS, Blassioli Moraes MC, Pareja M & Borges M (2009) Response of the egg parasitoids *Trissolcus basalis* and *Telenomus podisi* to compounds from defensive secretions of stink bugs. Journal of Chemical Ecology 35: 8–19.
- Laumann RA, Blassioli Moraes MC, Corrêa Ferreira MB, Caetano LD, Vieira ARA et al. (2007) Field applications of (*E*)-2-hexenal: a kairomone for egg parasitoids (Hymenoptera: Scelionidae). Book of Abstracts of the 23rd Annual Meeting of the International Society of Chemical Ecology, Jena, Germany, p. 177.
- Laumann RA, Bottura Maccagnan DH & Čokl A (2017) Use of vibratory signals for stink bug monitoring and control. Stink Bugs – Biorational Control Based on Communication Processes (ed. by A Čokl & M Borges), pp. 226–245. CRC Press, Boca Raton, FL, USA.
- Lee D-H, Short BD, Leskey TC, Joseph SV & Bergh JC (2013) Review of the biology, ecology, and management of *Halyomor-pha halys* (Hemiptera: Pentatomidae) in China, Japan, and the Republic of Korea. Environmental Entomology 42: 627–641.
- van Lenteren JC, Bale J, Bigler F, Hokkanen HMT & Loomans AJM (2006) Assessing risks of releasing exotic biological control agents of arthropod pests. Annual Review of Entomology 51: 609–634.
- Leskey TC & Nielsen AL (2018) Impact of the invasive brown marmorated stink bug in North America and Europe: history, biology, ecology, and management. Annual Review of Entomology 63: 599–618.
- Liljesthröm G & Rabinovich J (2004) Modeling biological control: the population regulation of *Nezara viridula* by *Trichopoda giacomellii*. Ecological Applications 14: 254–267.
- Lim UT & Mainali BP (2013) Effect of aggregation pheromone trap of *Riptortus pedestris* (Hemiptera: Alydidae) on the distribution and composition of its egg parasitoids. Journal of Economic Entomology 106: 1973–1978.
- Maistrello L, Costi E, Di Bella E, Vaccari G, Caruso S et al. (2020) Rilasci inondativi del parassitoide autoctono *Anastatus bifasciatus* per il controllo di *Halyomorpha halys* in Italia. Atti delle Giornate Fitopatologiche 2020, Extended Abstracts, CLUEB, Bologna, Italy (in press).

- Maistrello L, Dioli P, Bariselli M, Mazzoli GL & Giacalone-Forini I (2016) Citizen science and early detection of invasive species: phenology of first occurrences of *Halyomorpha halys* in Southern Europe. Biological Invasions 18: 3109–3116.
- Maistrello L, Dioli P, Dutto M, Volani S, Pasquali S & Gilioli G (2018) Tracking the spread of sneaking aliens by integrating crowdsourcing and spatial modeling: the Italian invasion of *Halyomorpha halys*. BioScience 68: 979–989.
- Maistrello L, Vaccari G, Caruso S, Costi E, Bortolini S et al. (2017) Monitoring of the invasive *Halyomorpha halys*, a new key pest of fruit orchards in northern Italy. Journal of Pest Science 90: 1231–1244.
- Mahmood R, Jones WJ, Bajwa BE & Rashid K (2015) Egg parasitoids from Pakistan as possible classical biological control agents of the invasive pest *Bagrada hilaris* (Heteroptera: Pentatomidae). Journal of Entomological Science 50: 147–149.
- Mani MS (1942) Studies on Indian parasitic Hymenoptera, II. Indian Journal of Entomology 4: 153–162.
- Marcu VC & Grozea I (2018) The potential impact of ecological and geographical parameters on the spread of the invasive pest *Nezara viridula*. Research Journal of Agricultural Science 50: 107–113.
- Martel G, Augé M, Talamas E, Roche M, Smith L & Sforza RFH (2019) First laboratory evaluation of *Gryon gonikopalense* (Hymenoptera: Scelionidae), as potential biological control agent of *Bagrada hilaris* (Hemiptera: Pentatomidae). Biological Control 135: 48–56.
- Martinez G, Soler R & Dicke M (2013) Behavioral ecology of oviposition-site selection in herbivorous true bugs. Advances in the Study of Behavior (ed. by HJ Brockmann, TJ Roper & M Naguib), pp. 175–207. Elsevier, Amsterdam, The Netherlands.
- Martorana L, Brodeur J, Foti MC, Agrò A, Colazza S & Peri E (2019) Egg parasitoid exploitation of plant volatiles induced by single or concurrent attack of a zoophytophagous predator and an invasive phytophagous pest. Scientific Reports 9: 18956
- Martorana L, Foti MC, Rondoni G, Conti E, Colazza S & Peri E (2017) An invasive insect herbivore disrupts plant volatile-mediated tritrophic signalling. Journal of Pest Science 90: 1079–1085
- Mason PG, Cock MJW, Barratt BIP, Klapwijk J, van Lenteren JC et al. (2018) Best practices for the use and exchange of inverte-brate biological control genetic resources relevant for food and agriculture. BioControl 63: 149–154.
- Mathews CR, Blaauw B, Dively G, Kotcon J, Moore J et al. (2017) Evaluating a polyculture trap crop for organic management of *Halyomorpha halys* and native stink bugs in peppers. Journal of Pest Science 90: 1245–1255.
- McPherson JE (2018) Invasive Stink Bugs and Related Species (Pentatomoidea): Biology, Higher Systematics, Semiochemistry, and Management. CRC Press. Boca Raton, FL, USA.
- McPherson JE, Bundy SC & Kuhar TP (2018) *Murgantia histrionica* (Hahn). Invasive Stink Bugs and Related Species (Pentatomoidea): Biology, Higher Systematics, Semiochemistry, and Management (ed. by JE McPherson), pp. 333–349. CRC Press, Boca Raton, FL, USA.

- Meiners T & Peri E (2013) Chemical ecology of insect parasitoids: essential elements for developing effective biological control programmes. Chemical Ecology of Insect Parasitoids (ed. by E Wajnberg & S Colazza), pp. 193–224. Wiley-Blackwell, Oxford, UK.
- Mendoza AC, Rocha ACP & Parra JRP (2016) Lyophilized artificial diet for rearing the Neotropical *Euschistus heros* (Hemiptera: Pentatomidae). Journal of Insect Science 16: 1–9.
- Michereff MFF, Borges M, Aquino MFS, Laumann RA, Mendes Gomes ACM & Blassioli Moraes MC (2016) The influence of volatile semiochemicals from stink bug eggs and oviposition-damaged plants on the foraging behaviour of the egg parasitoid *Telenomus podisi*. Bulletin of Entomological Research 106: 663–671.
- Michereff MFF, Laumann RA, Borges M, Michereff-Filho M, Diniz IR et al. (2011) Volatiles mediating a plant-herbivore-natural enemy interaction in resistant and susceptible soybean cultivars. Journal of Chemical Ecology 37: 273–285.
- Michereff MFF, Michereff Filho M, Blassioli Moraes MC, Laumann RA, Diniz IR & Borges M (2014) Effect of resistant and susceptible soybean cultivars on the attraction of egg parasitoids under field conditions. Journal of Applied Entomology 193: 207–216.
- Milnes JM, Wiman NG, Talamas EJ, Brunner JF, Hoelmer KA et al. (2016) Discovery of an exotic egg parasitoid of the brown marmorated stink bug, *Halyomorpha halys* (Stål) in the Pacific Northwest. Proceedings of the Entomological Society of Washington 118: 466–470.
- Milnes JM & Beers EH (2019) *Trissolcus japonicus* (Hymenoptera: Scelionidae) causes low levels of parasitism in three North American pentatomids under field conditions. Journal of Insect Science 19: 1–6.
- Mitchell PL, Zeilinger AR, Medrano EG & Esquivel JF (2018) Pentatomoids as vectors of plant pathogens. Invasive Stink Bugs and Related Species (Pentatomoidea) (ed. by JE McPherson), pp. 611–640. CRC Press, Boca Raton, FL, USA.
- Mizell RF III, Riddle TC & Blount AS (2008) Trap cropping system to suppress stink bugs in the southern coastal plain. Proceedings of the Florida State Horticultural Society 121: 377–382
- Møldrup ME, Geu-Flores F, de Vos M, Olsen CE, Sun J et al. (2012) Engineering of benzylglucosinolate in tobacco provides proof-of-concept for dead-end trap crops genetically modified to attract *Plutella xylostella* (diamondback moth). Plant Biotechnology Journal 10: 435–442.
- Molinari A, La Porta NC & Massoni F (2008) Parasitoides (Hymenoptera y Diptera) de hemípteros fitófagos. Chinches Fitófagas en Soja Revisión y Avances en el Estudio de su Ecología y Manejo (ed. by EV Trumper & JD Edelstein), pp. 107–128. INTA, Manfredi, Argentina.
- Moraglio ST, Tortorici F, Giromini D, Pansa MG, Visentin S & Tavella L (2021) Field collection of egg parasitoids of Pentatomidae and Scutelleridae in NW Italy and their evaluation on *Halyomorpha halys* in laboratory. Entomologia Experimentalis et Applicata 169: https://doi.org/10.1111/eea.12966 (this issue).

- Moraglio ST, Tortorici F, Pansa MG, Castelli G, Pontini M et al. (2020) A 3-year survey on parasitism of *Halyomorpha halys* by egg parasitoids in northern Italy. Journal of Pest Science 93: 183–194.
- Morrison WR, Blaauw BR, Nielsen AL, Talamas E & Leskey TC (2018) Predation and parasitism by native and exotic natural enemies of *Halyomorpha halys* (Stål) (Hemiptera: Pentatomidae) eggs augmented with semiochemicals and differing host stimuli. Biological Control 121: 140–150.
- Morrison WR, Blaauw BR, Short BD, Nielsen AL, Bergh JC et al. (2019) Successful management of *Halyomorpha halys* (Hemiptera: Pentatomidae) in commercial apple orchards with an attract-and-kill strategy. Pest Management Science 75: 104–114
- Morrison WR, Mathews CR & Leskey TC (2016) Frequency, efficiency, and physical characteristics of predation by generalist predators of brown marmorated stink bug (Hemiptera: Pentatomidae) eggs. Biological Control 97: 120–130.
- Morrison WR, Milonas P, Kapantaidaki DE, Cesari M, Bella ED et al. (2017) Attraction of *Halyomorpha halys* (Hemiptera: Pentatomidae) haplotypes in North America and Europe to baited traps. Scientific Reports 7: 1–11.
- Moujahed R, Frati F, Cusumano A, Salerno G, Conti E et al. (2014) Egg parasitoid attraction toward induced plant volatiles is disrupted by a non-host herbivore attacking above or belowground plant organs. Frontiers in Plant Science 5: 601.
- MPI (2020) Brown Marmorated Stink Bug Requirements. Ministry for Primary Industries, New Zealand. https://www.biosecurity.govt.nz/importing/vehicles-and-machinery/requirement-documents-for-importing-vehicles-machinery-or-parts/brown-marmorated-stink-bug-requirements/#bmsb-measures (accessed 30 July 2020).
- Musolin DL, Konjević A, Karpun NN, Protsenko VY, Ayba LY & Saulich AK (2018) Invasive brown marmorated stink bug *Halyomorpha halys* (Stål) (Heteroptera: Pentatomidae) in Russia, Abkhazia, and Serbia: history of invasion, range expansion, early stages of establishment, and first records of damage to local crops. Arthropod-Plant Interactions 12: 517–529.
- New Zealand Institute of Economic Research (2017) Quantifying the Economic Impacts of a Brown Marmorated Stink Bug Incursion in New Zealand. http://www.hortnz.co.nz/assets/UploadsNew/Quantifying-the-economic-impacts-of-a-Brown-Marmorated-Stink-Bug-Incursion.pdf (accessed 11 November 2019).
- Nielsen AL, Dively G, Pote JM, Zinati G & Mathews C (2016) Identifying a potential trap crop for a novel insect pest, *Halyomorpha halys* (Hemiptera: Pentatomidae), in organic farms. Environmental Entomology 45: 472–478.
- Nixon LJ, Tabb A, Morrison WR, Rice KB, Brockerhoff EG et al. (2019) Volatile release, mobility, and mortality of diapausing *Halyomorpha halys* during simulated shipping movements and temperature changes. Journal of Pest Science 92: 633–641.
- Nystrom E, Venette R, Dieckhoff C, Hoelmer K & Koch RL (2017) Cold tolerance of *Trissolcus japonicus* and *T. cultratus*, potential biological control agents of *Halyomorpha halys*, the brown marmorated stink bug. Biological Control 107: 11–20.

- Ogburn EC, Bessin R, Dieckhoff C, Dobson R, Grieshop M et al. (2016) Natural enemy impact on eggs of the invasive brown marmorated stink bug, *Halyomorpha halys* (Stål) (Hemiptera: Pentatomidae), in organic agroecosystems: a regional assessment. Biological Control 101: 39–51.
- Paini DR, Sheppard AW, Cook DC, Paul J, De Barro PJ et al. (2016) Global threat to agriculture from invasive species. Proceedings of the National Academy of Sciences of the USA 113: 7575–7579.
- Palumbo JC, Perring TM, Millar JG & Reed DA (2016) Biology, ecology and management of an invasive stink bug, *Bagrada hilaris*, in North America. Annual Review of Entomology 61: 453–473.
- Panizzi AR (2013) History and contemporary perspectives of the integrated pest management of soybean in Brazil. Neotropical Entomology 42: 119–127.
- Panizzi AR, McPherson JE, James DG, Javahery M & McPherson RM (2000) Stink bugs (Pentatomidae). Heteroptera of Economic Importance (ed. by CW Schaefer & AR Panizzi), pp. 421–474. CRC Press, Boca Raton, FL, USA.
- Panizzi AR & Silva FAC (2012) Seed-sucking bugs (Heteroptera). Insect Bioecology and Nutrition for Integrated Pest Management (ed. by AR Panizzi & JRP Parra), pp. 295-324. CRC Press, Boca Raton, FL, USA.
- Panizzi AR & Slansky F Jr (1985) Review of phytophagous pentatomids (Hemiptera: Pentatomidae) associated with soybean in the Americas. Florida Entomologist 68: 184–214.
- Panizzi AR & Lucini T (2016) What happened to *Nezara viridula* (L.) in the Americas? Possible reasons to explain populations decline. Neotropical Entomology 45: 619–628.
- Pashalidou FG, Frago E, Griese E, Poelman EH, van Loon JJA et al. (2015) Early herbivore alert matters: plant-mediated effects of egg deposition on higher trophic levels benefit plant fitness. Ecology Letters 18: 927–936.
- Paz-Neto AA, Querino RB & Margaría CB (2015) Egg parasitoids of stink bugs (Hemiptera: Coreidae and Pentatomidae) on soybean and cowpea in Brazil. Florida Entomologist 98: 929–932.
- Peri E, Moujahed R, Wajnberg E & Colazza S (2018) Applied chemical ecology to enhance insect parasitoid efficacy in the biological control of crop pests. Chemical Ecology of Insects (ed. by J Tabata), pp. 234–267. CRC Press, Boca Raton, FL, USA.
- Peri E, Salerno G, Slimani T, Frati F, Conti E et al. (2016) The response of an egg parasitoid to substrate-borne semiochemicals is affected by previous experience. Scientific Reports 6: 27098.
- Peri E, Sole MA, Wajnberg E & Colazza S (2006) Effect of host kairomones and oviposition experience on the arrestment behavior of an egg parasitoid. Journal of Experimental Biology 209: 3629–3635.
- Pickett JA, Woodcock CM, Midega CA & Khan ZR (2014) Push—pull farming systems. Current Opinion in Biotechnology 26: 125–132.
- Piemontese L, Bortolini S, Bulgarini G, Maistrello L, Rebecchi L et al. (2019) Identifying native predators of the alien pest *Haly-omorpha halys* (Hemiptera, Pentatomidae) with real-time PCR assay. Proceedings of the UZI 2019, Roma, Italy, p. 71

- Poelman EH & Kos M (2016) Complexity of plant volatile-mediated interactions beyond the third trophic level. Deciphering Chemical Language of Plant Communication (ed. by JD Blande & R Glinwood), pp. 211–225. Springer, Cham, Switzerland.
- Polajnar J, Maistrello L, Ibrahim A & Mazzoni V (2019) Can vibrational playback improve control of an invasive stink bug? Biotremology: Studying Vibrational Behavior (ed. by PSM Hill, R Lakes-Harlan, V Mazzoni, PM Narins, M Virant-Doberlet & A Wessel), pp. 375–398. Springer, Cham, Switzerland.
- Popovici OA, Vilhelmsen L, Masner L, Mikó I & Johnson N (2017) Maxillolabial complex in scelionids (Hymenoptera: Platygastroidea): morphology and phylogenetic implications. Insect Systematics & Evolution 48: 315–439.
- Pote JM (2018) Biological Control of *Halyomorpha halys* (Stål) (Hemiptera: Pentatomidae). PhD Dissertation, Rutgers University, New Brunswick, NJ, USA.
- Pote JM & Nielsen AL (2017) Life stage specific predation of Halyomorpha halys (Stål) by generalist predators. Biological Control 114: 1–7.
- Rahat S, Gurr GM, Wratten SD, Mo J & Neeson R (2005) Effect of plant nectars on adult longevity of the stinkbug parasitoid, *Trissolcus basalis*. International Journal of Pest Management 51: 321–324.
- Rather AH, Azim MN & Maqsood S (2010) Host plant selection in a pentatomid bug *Eurydema pulchrum* Westwood. Journal of Plant Protection Research 50: 229–232.
- Rea JH, Wratten SD, Sedcole R, Cameron PJ, Davis SI & Chapman RB (2002) Trap cropping to manage green vegetable bug *Nezara* viridula (L.) (Heteroptera: Pentatomidae) in sweet corn in New Zealand. Agricultural and Forest Entomology 4: 101–107.
- Reed D, Palumbo JC, Perring TM & May C (2013) *Bagrada hilaris* (Hemiptera: Pentatomidae), an invasive stink bug attacking cole crops in the southwestern United States. Journal of Integrated Pest Management 4: 1–7.
- Ribeiro A, Castiglioni E, Silva H & Bartaburu S (2009) Fluctuaciones de poblaciones de pentatómidos (Hemiptera: Pentatomidae) en soja (*Glycine max*) y lotus (*Lotus corniculatus*). Boletín de Sanidad Vegetal Plagas 35: 429–438.
- Rider DA (2016) Pentatomoidea Home Page. https://www.nd su.edu/pubweb/~rider/Pentatomoidea/ (accessed May 2020).
- van Rijn PCJ & Wäckers FL (2016) Nectar accessibility determines fitness, flower choice and abundance of hoverflies that provide natural pest control. Journal of Applied Ecology 53: 925–933.
- Rodríguez-Sifuentes L, Marszalek JE, Chuck-Hernández C & Serna-Saldívar SO (2020) Legumes protease inhibitors as biopesticides and their defense mechanisms against biotic factors. International Journal of Molecular Sciences 9: 3322.
- Rondoni G, Bertoldi V, Malek R, Djelouah K, Moretti C et al. (2018) *Vicia faba* plants respond to oviposition by invasive *Halyomorpha halys* activating direct defences against offspring. Journal of Pest Science 91: 671–679.
- Rondoni G, Bertoldi V, Malek R, Foti MC, Peri E et al. (2017) Native egg parasitoids recorded from the invasive

- Halyomorpha halys successfully exploit volatiles emitted by the plant–herbivore complex. Journal of Pest Science 90: 1087–1095.
- Rondoni G, Borges I, Collatz J, Conti E, Costamagna A et al. (2021) Exotic ladybirds for biological control of herbivorous insects a review. Entomologia Experimentalis et Applicata 169: https://doi.org/10.1111/eea.12963 (this issue).
- Roversi PF, Binazzi F, Marianelli L, Costi E, Maistrello L & Sabbatini Peverieri G (2016) Searching for native egg-parasitoids of the invasive alien species *Halyomorpha halys* Stål (Heteroptera, Pentatomidae) in Southern Europe. Redia 99: 63–70.
- Sabbatini Peverieri G, Talamas E, Bon MC, Marianelli L, Bernardinelli I et al. (2018) Two Asian egg parasitoids of *Halyomorpha halys* (Stål) (Hemiptera, Pentatomidae) emerge in northern Italy: *Trissolcus mitsukurii* (Ashmead) and *Trissolcus japonicus* (Ashmead) (Hymenoptera, Scelionidae). Journal of Hymenoptera Research 67: 37–53.
- Sabbatini Peverieri G, Mitroiu MD, Bon MC, Balusu R, Benvenuto L et al. (2019) Surveys of stink bug egg parasitism in Asia, Europe and North America, morphological taxonomy, and molecular analysis reveal the Holarctic distribution of Acroclisoides sinicus (Huang & Liao) (Hymenoptera: Pteromalidae). Journal of Hymenoptera Research 74: 123–151.
- Salerno G, Frati F, Conti E, Peri E, Colazza S & Cusumano A (2019) Mating status of an herbivorous stink bug female affects the emission of oviposition-induced plant volatiles exploited by an egg parasitoid. Frontiers in Physiology 10: 398.
- Salerno G, Frati F, Marino G, Ederli L, Pasqualini S et al. (2017) Effects of water stress on emission of volatile organic compounds by Vicia faba, and consequences for attraction of the egg parasitoid Trissolcus basalis. Journal of Pest Science 90: 635–647.
- Salisbury A, Barclay MVL, Reid S & Halstead A (2009) The current status of the southern green shield bug, *Nezara viridula* (Hemiptera: Pentatomidae), an introduced pest species recently established in South-East England. British Journal of Entomology and Natural History 22: 189–194.
- Saluso A, Ermácora O, Anglada M, Toledo C & Borguesan C (2007) Principales invertebrados plagas de la soja y técnicas utilizadas en la toma de decisiones (Campaña agrícola 2006– 2007). Revista de Ciencias Agropecuarias 11: 153–158.
- Samuel CK (1942) Biological notes on two new egg parasites of *Bagrada picta* Fabr., Pentatomidae. Indian Journal of Entomology 4: 92–93.
- Sastawa BM, Lawan M & Maina YT (2004) Management of insect pests of soybean: effects of sowing date and intercropping on damage and grain yield in the Nigerian Sudan savanna. Crop Protection 23: 155–161.
- Scaccini D, Falagiarda M, Tortorici F, Martinez-Sañudo I, Tirello P et al. (2020) An insight into the role of *Trissolcus mitsukurii* as biological control agent of *Halyomorpha halys* in northeastern Italy. Insects 11: 306.
- Serteyn L, Ponnet L, Saive M, Fauconnier M-L & Francis F (2020) Changes of feeding behavior and salivary proteome of brown marmorated stink bug when exposed to insect-induced plant defences. Arthropod-Plant Interactions 14: 101–112.

- Sforza RFH, Bon M-C, Martel G, Augé M, Roche M et al. (2017) Initial evaluation of two native egg parasitoids for the control of *Bagrada hilaris*, an invasive stink bug in western USA. Proceedings of the Vth International Symposium on Biological Control of Arthropods, 11–15 September 2017, Langkawi, Malaysia (ed. by PG Mason, DR Gillepsie & C Vincent), pp. 221-223. CABI, Wallingford, UK.
- Sharkey MJ (2007) Phylogeny and classification of Hymenoptera. Zootaxa 1668: 521–548.
- Silva CC, Laumann RA, Blassioli Moraes MC, Pareja M & Borges M (2008) Euschistus heros mass rearing technique for the multiplication of Telenomus podisi. Pesquisa Agropecuária Brasileira 43: 575–580.
- Silva GV, Bueno AF, Favetti BM & Neves PMOJ (2018) Use of low temperature storage to preserve host and parasitoid to improve the rearing of *Telenomus podisi* (Hymenoptera: Platygastridae) on *Euschistus heros* (Hemiptera: Pentatomidae) eggs. Neotropical Entomology 48: 126–135.
- Simpson M, Gurr GM, Simmons AT, Wratten SD, James DG et al. (2011) Attract and reward: combining chemical ecology and habitat manipulation to enhance biological control in field crops. Journal of Applied Ecology 48: 580–590.
- Simpson M, Read DM & Gurr GM (2013) Application of chemical cues in arthropod pest management for organic crops. Chemical Ecology of Insect Parasitoids (ed. by E Wajnberg & S Colazza), pp. 266–281. Wiley-Blackwell, Oxford, UK.
- Smith HA & McSorley R (2000) Intercropping and pest management: a review of major concepts. American Entomologist 46: 154–161.
- Soergel DC, Ostiguy N, Fleischer SJ, Troyer RR, Rajotte EG & Krawczyk G (2015) Sunflower as a potential trap crop of *Halyomorpha halys* (Hemiptera: Pentatomidae) in pepper fields. Environmental Entomology 44: 1581–1589.
- Sparks ME, Shelby KS, Kuhar D & Gundersen-Rindal DE (2014)
 Transcriptome of the invasive brown marmorated stink bug, *Halyomorpha halys* (Stål) (Heteroptera: Pentatomidae). PLoS
 ONE 9: e111646.
- Stahl JM, Babendreier D, Marazzi C, Caruso S, Costi E et al. (2019a) Can Anastatus bifasciatus be used for augmentative biological control of the brown marmorated stink bug in fruit orchards? Insects 10: 108.
- Stahl J, Tortorici F, Pontini M, Bon M-C, Hoelmer K et al. (2019b) First discovery of adventive populations of *Trissolcus japonicus* in Europe. Journal of Pest Science 92: 371–379.
- Stecca CS, Bueno AF, Pasini A, Silva DM, Andrade K & Zirondi Filho DM (2017) Impact of insecticides used in soybean crops to the egg parasitoid *Telenomus podisi* (Hymenoptera: Platygastridae). Neotropical Entomology 47: 281–291.
- Subba Rao BR & Chacko MJ (1961) Studies on Allophanurus indicus n. sp., an egg parasite of Bagrada cruciferarum Kirkaldy. Beiträge zur Entomologie 11: 812–824.
- Talamas EJ, Buffington ML & Hoelmer KA (2017) Revision of Palearctic *Trissolcus* Ashmead (Hymenoptera, Scelionidae). Journal of Hymenoptera Research 56: 3–185.
- Talamas EJ, Herlihy MV, Dieckhoff C, Hoelmer KA, Buffington M et al. (2015) Trissolcus japonicus (Ashmead) (Hymenoptera,

51

- Scelionidae) emerges in North America. Journal of Hymenoptera Research 43: 119-128.
- Trase (2018) Trase Yearbook 2018, Sustainability in forest-risk supply chains: Spotlight on Brazilian soy. Transparency for Sustainable Economies, Stockholm Environment Institute and Global Canopy. https://yearbook2018.trase.earth/
- Taylor ME, Bundy CS & McPherson JE (2014) Unusual ovipositional behavior of the stink bug Bagrada hilaris (Hemiptera: Heteroptera: Pentatomidae) with descriptions of immature stages. Annals of the Entomological Society of America 108: 536-551.
- Tillman PG (2006) Sorghum as a trap crop for Nezara viridula (L.) (Heteroptera: Pentatomidae) in cotton in the Southern United States. Environmental Entomology 35: 771-783.
- Tillman PG & Carpenter JE (2014) Milkweed (Gentianales: Apocynaceae): a farmscape resource for increasing parasitism of stink bugs (Hemiptera: Pentatomidae) and providing nectar to insect pollinators and monarch butterflies. Environmental Entomology 43: 370-376.
- Tillman PG & Cottrell TE (2012) Case study: trap crop with pheromone traps for suppressing Euschistus servus (Heteroptera: Pentatomidae) in cotton. Psyche 2012: 401703.
- Tillman PG, Khrimian A, Cottrell TE, Lou X, Mizell RF III & Johnson CJ (2015) Trap cropping systems and a physical barrier for suppression of stink bugs (Hemiptera: Pentatomidae) in cotton. Journal of Economic Entomology 108: 2324-2334.
- Tillman G (2017) Ecosystem-based incorporation of nectar-producing plants for stink bug parasitoids. Insects 8: 65.
- Tillman G, Toews M, Blaauw B, Sial A, Cottrell T et al. (2020) Parasitism and predation of sentinel eggs of the invasive brown marmorated stink bug, Halyomorpha halys (Stål) (Hemiptera: Pentatomidae), in the southeastern US. Biological Control 145: 104247.
- Todd JW & Schumann FW (1988) Combination of insecticide applications with trap crops of early maturing soybean and southern peas for population management of Nezara viridula in soybean (Hemiptera: Pentatomidae). Journal of Entomological Science 23: 192-199.
- Todd JW (1989) Ecology and behavior of Nezara viridula. Annual Review of Entomology 34: 273-292.
- Tognon R, Aldrich JR, Buffington ML, Talamas EJ, Sant'Ana J & Zalom FG (2017) Halyomorpha halys (Heteroptera: Pentatomidae) egg surface chemicals inhibit North American Telenomus and Trissolcus (Hymenoptera: Scelionidae) parasitism. Biological Control 114: 39-44.
- Tognon R, Sant'Ana J, Zhang Q-H, Millar JG, Aldrich JR & Zalom FG (2016) Volatiles mediating parasitism of Euschistus conspersus and Halyomorpha halys eggs by Telenomus podisi and Trissolcus erugatus. Journal of Chemical Ecology 42: 1016-1027.
- Tortorici F, Talamas EJ, Moraglio ST, Pansa MG, Asadi-Farfar M et al. (2019) A morphological, biological and molecular

- approach reveals four cryptic species of Trissolcus Ashmead (Hymenoptera Scelionidae), egg parasitoids of Pentatomidae (Hemiptera). Journal of Hymenoptera Research 73: 153–200.
- Tougou D, Musolin DL & Fujisaki K (2009) Some like it hot! Rapid climate change promotes shifts in distribution ranges of Nezara viridula and N. antennata in Japan. Entomologia Experimentalis et Applicata 30: 249-258.
- Tschorsnig HP (2017) Preliminary Host Catalogue of Palaearctic Tachinidae (Diptera). [online] http://www.nadsdiptera.org/ Tach/WorldTachs/CatPalHosts/Home.html (accessed August 2020).
- Vieira CR (2010) Ecologia Química de Insetos Parasitóides de Ovos (Hymenoptera: Scelionidae) e sua Aplicação no Controle Biológico de Pragas. Embrapa Recursos Genéticos e Biotecnologia. MEcol Thesis, Universidade de Brasília, Brasília,
- Vieira CR, Blassioli Moraes MC, Borges M, Pires CSS, Sujii ER & Laumann RA (2014) Field evaluation of (E)-2-hexenal efficacy for behavioral manipulation of egg parasitoids in soybean. BioControl 59: 525-537.
- Vieira CR, Blassioli Moraes MC, Borges M, Sujii ER & Laumann RA (2013) Cis-jasmone indirect action on egg parasitoids (Hymenoptera: Scelionidae) and its application in biological control of soybean stink bugs (Hemiptera: Pentatomidae). Biological Control 64: 75-82.
- Wajnberg E & Colazza S (2013) Chemical Ecology of Insect Parasitoids. Wiley-Blackwell, Oxford, UK.
- Weber DC, Khrimian A, Blassioli Moraes MC & Millar JG (2018) Semiochemistry of pentatomoidea. Invasive Stink Bugs and Related Species (Pentatomoidea): Biology, Higher Systematics, Semiochemistry, and Management (ed. by JE McPherson), pp. 677-725. CRC Press, Boca Raton, FL, USA.
- Yamanaka T, Teshiba M, Tuda M & Tsutsumi T (2011) Possible use of synthetic aggregation pheromones to control stinkbug Plautia stali in kaki persimmon orchards. Agricultural and Forest Entomology 13: 321–331.
- Zerbino MS & Panizzi AR (2019) The underestimated role of pest pentatomid parasitoids in Southern South America. Arthropod-Plant Interactions 13: 703-718.
- Zhang J, Zhang F, Gariepy T, Mason P, Gillespie D et al. (2017) Seasonal parasitism and host specificity of Trissolcus japonicus in northern China. Journal of Pest Science 90: 1127-1141.
- Zhang QH, Schneidmiller RG, Hoover DR, Zhou G, Margaryan A & Bryant P (2014) Essential oils as spatial repellents for the brown marmorated stink bug, Halyomorpha halys (Stål) (Hemiptera: Pentatomidae). Journal of Applied Entomology 138: 490-499.
- Zhong YZ, Zhang JP, Ren LL, Tang R, Zhan HX et al. (2017) Behavioral responses of the egg parasitoid Trissolcus japonicus to volatiles from adults of its stink bug host, Halyomorpha halys. Journal of Pest Science 90: 1097-1105.