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Monitoring the diversity of pest and nonpest noctuid moth (Lepidoptera: Noctuidae) species in Canadian prairie agroecosystems

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Abstract

The Noctuidae (Lepidoptera) comprise the most diverse and abundant lepidopteran families in the Canadian Prairie Ecozone. Within this group, some species are agricultural pests that require monitoring. Pheromone lures target specific species, whereas food-bait lures attract a broader range. This study reports the diversity and abundance of noctuid moths captured in traps baited with female sex pheromones of pest species and with food-bait lures consisting of acetic acid and 3-methyl-1-butanol (AAMB) with fermented byproduct or floral volatile compounds. Food-bait lures that attract pests and nonpest species can provide insight into moth populations and species richness in human-managed ecosystems. We trapped moths in wheat (Poaceae) and canola (Brassicaceae) fields in central Alberta, Canada. We captured and identified to species approximately 7900 noctuid moths. Community composition was similar in both crops. Sex pheromone-baited traps had variable specificity and low nontarget diversity. Traps baited with AAMB captured greater moth diversity than unbaited traps did. Noctuidae were the most diverse and abundant in AAMB-baited traps (62 species across 8 tribes). The AAMB lures captured more cutworm and armyworm pests than unbaited traps did. Fermented byproduct food-bait lures captured more noctuid pests than floral volatiles did. The AAMB lures can be implemented to monitor Noctuidae diversity and potentially assess local noctuid pest density in agroecosystems on the Canadian prairies.

Introduction

Long-term datasets indicate a global decline in insect abundance and diversity (Hallmann *et al.* 2017; Laussmann *et al.* 2021). One of the main drivers of insect decline generally is habitat loss due to extensive land modification to accommodate agriculture and urbanisation (Sánchez-Bayo and Wyckhuys 2019). Other factors contributing to insect decline are chemical pollutants from pesticides and fertilisers, biotic factors, including pathogens and invasive species, and climate change (Sánchez-Bayo and Wyckhuys 2019).

Lepidoptera are among the most affected insect taxa, with examples of severe moth decline observed in the Northern Hemisphere (Wagner *et al.* 2021). The decline in moth abundance and diversity in grassland ecosystems, in particular, demonstrates the sensitivity of this taxon to disturbances (Mangels *et al.* 2017; Sánchez-Bayo and Wyckhuys 2021). At least 75% of the grassland habitat in the Canadian Prairie Provinces has been altered for crop cultivation and livestock production (Shorthouse 2010). Landscape fragmentation to support intensive monocultures of annual crops and perennial forages is a major driver of the decline in

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arthropod diversity in agroecosystems (Meehan *et al.* 2013). Agronomic practices, such as crop rotations, soil disturbance through tillage and harvest, and chemical and organic inputs, disturb habitat to a high degree, which impacts insect community structure and population density of pest species (Shennan 2008; Evans and Sanderson 2018). Detection of changes in moth diversity and abundance requires systematic monitoring over time.

The order Lepidoptera is one of the most diverse insect taxa in the Prairie Ecozone of Canada, with 2232 species recorded in 61 families (Pohl *et al.* 2014). Moths from the families Geometridae, Erebiidae, and Noctuidae represent 78% of the total diversity of macromoth species, and noctuid moths alone, with 693 reported species, make up 28% of the entire lepidopteran fauna in the Canadian Prairies (Pohl *et al.* 2014). Noctuid moths perform important ecological roles in the biotic interactions in prairie habitats. Larvae and adults are food sources for higher trophic levels, including predatory beetles (Frank 1971; Cárcamo *et al.* 1995), spiders (Pearce *et al.* 2004), grassland birds (Maher 1979), and insectivorous bats (Vonhof and Hobson 2001). Immature stages of moths serve as hosts to a wide diversity of hymenopteran parasitoids, including wasps in the family Braconidae and Ichneumonidae, and dipteran parasitoids, including flies in the family Tachinidae (Mills 1993; Stireman and Singer 2003). Although often overlooked, noctuid moths also act as generalist nocturnal pollinators in several plant systems and are capable of dispersing pollen over longer distances than other insects typically do (Reynolds *et al.* 2009; Winfree *et al.* 2011). Furthermore, feeding damage by moth larvae can impact plant community structure, which is dependent on the spatial and temporal variation of moth populations (Crawley 1989).

A decrease in moth diversity and abundance can have cascading effects and lead to further losses of plant–animal interactions in grassland ecosystems that could threaten ecosystem services (Wagner *et al.* 2021). The vast majority of lepidopteran species recorded in the Canadian Prairie Provinces (~98%) have a neutral effect on agricultural production because crop plants either are not larval hosts nor provide valuable ecosystem services to managed systems (Pohl *et al.* 2014; Floate 2017). Only a few lepidopteran species (~25 species) are considered pests of some cereal, pulse, and rapeseed oil crops (Floate 2017; Vankosky *et al.* 2017). Cutworms and armyworms (Lepidoptera: Noctuidae) are a pest complex in North America that can cause occasional economic damage to several annual crops grown across the Canadian Prairies (Beirne 1971; Floate 2017). The larvae and adults are generalist herbivores that feed on a wide range of host species in several plant families. Cutworm damage occurs in early summer when late-instar larvae display characteristic feeding behaviour by cutting the stems of seedlings, which ultimately kills the plant (Beirne 1971). Armyworm injury occurs in mid to late summer. Late-instar larvae feed on foliage and disperse *en masse* across the landscape in search of host plants when food is depleted (Mason *et al.* 1998). At low population densities, larval-feeding damage results in crop thinning or bare patches, whereas sporadic outbreaks have caused the complete destruction of entire fields (Beirne 1971). For example, outbreaks of the Bertha armyworm, *Mamestra configurata* Walter (Lepidoptera: Noctuidae), in canola, *Brassica napus* Linnaeus (Brassicaceae), caused yield losses of \$Cdn 30 million in 1995 and spraying costs of approximately \$Cdn 16.5 million in western Canada in 1994 and 1995 (Mason *et al.* 1998). The redbacked cutworm, *Euxoa ochrogaster* (Guenée) (Lepidoptera: Noctuidae), and the pale western cutworm, *Agrotis orthogonia* (Morrison) (Lepidoptera: Noctuidae), are the most common cutworm species, with localised outbreaks across the Prairie Provinces reported by the Western Committee on Crop Pests (2015, 2016). The diversity and abundance of this pest complex are highly variable and are influenced by regional climate, agricultural practices, moth life history traits, and other factors (Floate and Hervet 2017). Systematic monitoring of cutworm and armyworm populations is necessary to detect and predict population increases and outbreak levels. Moth trap captures can be used as an early warning system to indicate the risk of larval damage and thereby inform producers and scouters of localised areas in which to emphasise larval sampling.

Traps baited with synthetic lures of species-specific female sex pheromones have been used to monitor populations of some lepidopteran pest species (Byers and Struble 1987;

Ayre and Lamb 1990). However, sex pheromone-baited traps attract only male moths, which may not accurately reflect populations of reproducing females (Byers *et al.* 1987). Although this approach is useful for monitoring individual species, it is not appropriate for assessing moth diversity. Traps baited with a generalised lure that attracts multiple cutworm and armyworm species, as well as nonpest species, can provide valuable information about the population dynamics of pest species and can document lepidopteran species richness in human-managed ecosystems in the prairie regions across temporal scales.

Light trapping with ultraviolet light sources has often been used to monitor lepidopteran diversity and abundance in agroecosystems (Ayre and Lamb 1990; Chey *et al.* 1997; Beck *et al.* 2002). Moth captures in light traps depend on environmental conditions (Yela and Holyoak 1997) and the wavelength of the light source (van Langevelde *et al.* 2011; Merckx and Slade 2014) and require an external power source. Extended day length in summer can reduce moth attraction to light traps in northern latitudes due to natural light reducing moth activity around light traps (Jonason *et al.* 2014). Due to these limitations, light trapping assessments generally occur periodically throughout the growing season and may not capture the temporal variation of the moth community assemblage (Lintott *et al.* 2014).

Food-based semiochemicals attract both male and female moths and have potential as lures to detect and monitor noctuid moth diversity and pest noctuid species (Joyce and Lingren 1998). In contrast to the limitations of light trapping, traps baited with food-bait lures remain in place throughout the growing season to survey moth populations and to gather information on seasonal flight patterns. Fermented sugars were some of the first food-bait lures used to monitor lepidopteran diversity. Utrio and Eriksson (1977) trapped several macrolepidopteran species with individual and mixtures of volatile compounds (simple alcohol, acids, esters, and aldehydes) released from fermented sources of multiple sugars. A mixture of two fermented sugar byproducts, acetic acid and 3-methyl-1-butanol (AAMB), has been used to monitor the diversity and abundance of moths in multiple cropping systems (Landolt *et al.* 2007, 2011). Several pest species from the cutworm and armyworm complex are attracted to AAMB lures, including the Bertha armyworm (Landolt 2000), true armyworm, *Mythimna unipuncta* (Haworth) (Landolt and Higbee 2002), and the redbacked cutworm (Landolt *et al.* 2007). Food-bait lures comprised of volatile compounds, including phenylacetaldehyde and benzaldehyde, released from flowers visited by noctuid moths have been used to monitor populations of the cabbage looper, *Trichoplusia ni* (Hübner) (Cantelo and Jacobson 1979), alfalfa looper, *Autographa californica* (Speyer) (Guédot *et al.* 2008), and the soybean looper, *Chrysodeixis ubcludens* (Walker) (Meagher 2001). Although fermented sugar baits and lures releasing floral volatiles attract a broad group of noctuid moths, responses to food-based semiochemicals by different lepidopteran taxa may differ.

Our previous work (Batallas and Evenden 2023) assessed the attractiveness of food-bait lures compared to sex pheromone-baited traps to noctuid pest species in the prairies. In the present study, we report on the diversity (number of species captured in traps) and abundance of all noctuid moths captured in traps baited with food-based semiochemicals derived from fermented sugar byproducts and floral volatiles compared to the diversity and abundance of noctuid moths captured in sex pheromone-baited traps. Specifically, we asked the following questions: (1) Does lepidopteran species composition vary in relation to the annual crop grown in the current season? and (2) Do food-based semiochemicals from fermented sugar baits or floral volatiles influence the attraction and capture of different lepidopteran taxa? We evaluate differences in species composition of moths sampled in two cropping systems – canola, *Brassica napus* Linnaeus (Brassicaceae), and wheat, *Triticum aestivum* Linnaeus (Poaceae) – in central Alberta, Canada. We examine differences in lepidopteran taxa – specifically moths within the subfamily Noctuidae – attracted to AAMB lures alone and to traps baited with additional food-based semiochemicals. We hypothesised that (1) noctuid moth species composition would be similar in canola and wheat fields, reflecting a generalised moth community in human-managed ecosystems due to the annual levels of

disturbance and the strong dispersal capabilities of moths, and (2) noctuid moths would be attracted to traps baited with volatiles from fermented sugar baits because noctuid moths use microbial volatile organic compounds as semiochemicals to locate food resources.

Despite the important role of noctuid moths in prairie ecosystems, information on the impact of agronomic practices on the status of moth diversity and abundance in agricultural ecosystems is lacking. Moth community composition can be used as a bioindicator to reflect the state of disturbance in agricultural ecosystems and to improve management strategies (Olfert *et al.* 2002).

Methods

Study area

Experiments were conducted in 2014 and 2015 in wheat and canola fields located in the Aspen Parkland Ecoregion of Alberta, Canada. The landscape is dominated by extensive agricultural plains with discontinuous clusters of trembling aspen, *Populus tremuloides* Michaux (Salicaceae), and balsam poplar, *P. balsamifera* Linnaeus (Salicaceae) (Shorthouse 2010). Seven sites separated by at least 20 km were selected over an area of approximately 7350 km² in central Alberta (Supplementary material, Fig. S1). Each site comprised a commercial canola field paired with a commercial wheat field, approximately 500 m distant from each other. Each field was an experimental unit, and the size of each field was 0.65 km². Experiments were conducted at the same sites in both years, but crops grown in the two years were rotated between fields.

Lures

All experiments compared moth diversity in traps baited with food-bait lures or synthetic female sex pheromone lures (Table 1). The sex pheromone lures of the cutworm and armyworm species targeted in the present study (Table 1) are commercially available as red rubber septa loaded by and purchased from Contech Enterprises Inc. (Delta, British Columbia, Canada). Food-bait lures were prepared in 15-mL narrow-mouth Nalgene HDPE bottles (Thermo Scientific, Rochester, New York, United States of America), following the methods of Landolt *et al.* (2007). The AAMB lure consisted of AAMB in a 50:50 by weight mixture (glacial acetic acid, 99.7% purity; Fisher Scientific, Fair Lawn, New Jersey, United States of America; and 3-methyl-1-butanol, 98.5% purity; Sigma Aldrich, St. Louis, Missouri, United States of America). Ten millilitres of the AAMB lure were loaded into each bottle, followed by two cotton balls to prevent spilling. A 3.0-mm-diameter hole in the centre of the bottle cap allowed for the release of volatile compounds.

Monitoring and moth identification

All experiments used nonsaturating green universal moth traps (Unitrap, Contech Enterprise Inc. Delta, British Columbia, Canada), baited with either a sex pheromone or a food-bait lure. At each site, unitraps were placed in a linear transect separated by 25 m and positioned 5 m into the field from the edge, at a height of 1.5 m above ground. One replicate of each of the treatments was randomly assigned to traps along the transect in each field at each site. Sex pheromone lures were placed inside baskets attached to the lid of the unitrap and replaced every four weeks. Food-bait lures were attached to the inside of the unitrap bucket with a twist tie and replaced every two weeks. An insecticidal strip of Hercon Vaportape II (10% dichlorvos; Hercon Environmental, Emigsville, Pennsylvania, United States of America) was placed in each trap bucket to kill captured insects and was replaced every four weeks.

Insect trap-catch was collected weekly and frozen at −20 °C until sorting and identification. Moths were separated by sex and pinned. Genitalic dissections were performed on specimens in poor condition (*i.e.*, no scales on wings or missing body parts), following Hardwick's (1950) methods (Batallas and Evenden 2023). Moth genitalia were spread and mounted on cardstock

Table 1. Lure composition and deployment schedule for noctuid moth monitoring experiments in canola and wheat fields in central Alberta, Canada, in 2014 and 2015. Experiments evaluated noctuid moth diversity and abundance captured in traps baited with food-bait lures from fermented sugar byproducts and floral volatiles compared to sex pheromone-baited traps

Year	Lure	Components	Ratio	Amount	Time deployed
2014	Redbacked cutworm (<i>Euxoa ochrogaster</i>)	Z5-12Ac, Z7-12Ac, Z9-12Ac, Z5-10Ac	200:2:1:1	1000 µg	23 June-10 October
	Bertha armyworm (<i>Mamestra configurata</i>)	Z11-16Ac, Z9-14Ac	95:5	500 µg	10 June-02 September
	True armyworm (<i>Mythimna unipuncta</i>)	Z11-16Ac	1	1000 µg	10 June-10 October
	Army cutworm (<i>Euxoa auxiliaris</i>)	Z5-14Ac, Z7-14Ac, Z9-14Ac	100:1:10	100 µg	02 September-10 October
	AAMB	Acetic acid, 3-methyl-1-butanol	1:1	10 mL	10 June-10 October
	Unbaited control	-	-	-	10 June-10 October
2015	Redbacked cutworm (<i>Euxoa ochrogaster</i>)	Z5-12Ac, Z7-12Ac, Z9-12Ac, Z5-10Ac	200:2:1:1	1000 µg	22 June-15 September
	Bertha armyworm (<i>Mamestra configurata</i>)	Z11-16Ac, Z9-14Ac	95:5	500 µg	22 June-04 August
	True armyworm (<i>Mythimna unipuncta</i>)	Z11-16Ac	1	1000 µg	22 June-15 September
	Pale western cutworm (<i>Agrotis orthogonia</i>)	Z7-12Ac, Z5-12Ac	2:1	500 µg	22 June-15 September
	AAMB	Acetic acid, 3-methyl-1-butanol	1:1	10 mL	22 June-15 September
	AAMB + MP	Acetic acid, 3-methyl-1-butanol, 2-methyl-1-propanol	1:1:1	10 mL	22 June-15 September
	AAMB + PAA	Acetic acid, 3-methyl-1-butanol, phenylacetaldehyde	1:1:1	10 mL	22 June-15 September
	AAMB + MP + PAA	Acetic acid, 3-methyl-1-butanol, 2-methyl-1-propanol, phenylacetaldehyde	1:1:1:1	10 mL	22 June-15 September
	Unbaited control	-	-	-	22 June-15 September

(2.0 × 0.5 cm) with an Euparal mounting medium (Bioquip Products Inc., Rancho Dominguez, California, United States of America). Moths were identified to species through wing maculation and morphological characters of genitalia, following taxonomic keys from Lafontaine (1987, 1998, 2004), Lafontaine and Robert (1991), and Mikkola *et al.* (2009). Identifications were verified using comparisons with reference collections at the E.H. Strickland Entomological Museum (University of Alberta, Edmonton, Alberta, Canada). Pinned moths in the best condition and mounted genitalia dissections from each identified species were selected as voucher specimens and deposited at the E.H. Strickland Entomological Museum.

Experiment 1: moth diversity in synthetic sex pheromone- and AAMB-baited traps

Experiment 1 assessed AAMB lures for monitoring noctuid moth diversity in canola and wheat fields. Sex pheromone-baited traps for cutworm and armyworm pest species were also erected at sites to determine if pest moths were present in the field at the time of the experiment. Unitraps were baited with redbacked cutworm pheromone (canola pest), Bertha armyworm pheromone (canola pest), true armyworm pheromone (cereal pest), army cutworm, *E. auxiliaris* (Grote), pheromone (canola and wheat pest), or AAMB lure or were left unbaited (Table 1). Six baited traps (one trap per treatment per field; Table 1) were positioned in a 125-m linear transect, as described above, in random order in both canola and wheat fields at each of the seven sites. The experiment was conducted from 10 June to 10 October 2014. Sex pheromone-baited traps were deployed in the field according to the flight period of the target moth species (Table 1), whereas the AAMB-baited and unbaited traps remained in the field throughout the 17-week sampling period to assess moth diversity in general. Moths captured in the differently baited traps were identified to species, and analyses were conducted on the total number of moths captured per lure type in the respective trap. Moth diversity and abundance in the sex pheromone-baited traps were used to evaluate the specificity of each synthetic sex pheromone lure. The diversity and abundance of moths captured in the AAMB-baited traps were compared to those captured in the unbaited control traps.

Experiment 2: moth diversity in traps baited with food-based semiochemicals

Experiment 2 evaluated the diversity and abundance of noctuid moths attracted to AAMB lures with and without the addition of other food-based compounds. Tóth *et al.* (2010) demonstrated that the combination of AAMB lures with 2-methyl-1-propanol or phenylacetaldehyde attracts multiple noctuid species in Europe. We tested these two compounds to enhance the attraction of AAMB lure to monitor cutworm and armyworm pests present in the central Alberta region. Compounds tested in conjunction with AAMB lure included an alcohol fermentation byproduct from sugar baits, 2-methyl-1-propanol (MP, > 99% purity; Acros Organics, Fair Lawn, New Jersey), and a floral volatile, phenylacetaldehyde (PAA, > 98% purity; Acros Organics). Treatments included AAMB alone, AAMB + MP, AAMB + PAA, and AAMB + MP + PAA, as well as an unbaited trap that served as the control. All lures were prepared in the laboratory in a mixture of equal proportions by weight (Table 1). Ten millilitres of the chemical mixtures were loaded in Nalgene HDPE bottles, as previously described (see Lures subsection). In addition to the different food-bait lures, sex pheromone-baited traps targeting redbacked cutworm, Bertha armyworm, pale western cutworm, and true armyworm were deployed to ensure target moths were present in the field at the time of the experiment (Table 1). Nine baited traps (one trap per treatment per field), including those with the pheromone and food-bait lures, and the unbaited trap were positioned in random order in a 200-m linear transect, as described above, in each canola field and each wheat field at each of the seven sites. The experiment was conducted from 22 June to 15 September 2015. Noctuid moth trap catch was identified to species, and analyses were

conducted on the total number of moths captured over the sampling season. Results focused on comparing the moth trap catch from the different food-bait lures and unbaited traps.

Statistical analyses

For experiment 1, we determined the specificity of the sex pheromone lures to monitor target moths. Moth trap catch was separated into two groups: target moths and nontarget moths. The percentage of target moths captured in sex pheromone-baited traps was calculated, based on the total moth trap catch ((total number of target moths/total moth trap catch) $\times$ 100) for each of the sex pheromone lures tested. Lure specificity was analysed in a binomial count model in which the response variable is a two-vector object comprised of the count of target moths as the first vector (success) and the count of nontarget moths as the second vector (failure). The two-vector response variable was analysed in a generalised linear mixed model with binomial family distribution with the “glmer” command in R package’s *lme4*, version 1.1-17 (Bates *et al.* 2015). Crop and sex pheromone lure were specified as explanatory fixed variables, and site was specified as a random block factor.

For both experiments, several analyses compared the total capture of moths in food-bait lures and unbaited traps. A nonmetric multidimensional scaling (“Bray–Curtis” distance) analysis was conducted (1) to determine the diversity of moth species attracted to the food-bait lures compared to the unbaited traps and (2) to evaluate differences in moth species composition sampled in canola fields and in wheat fields. A nonparametric permutation analysis of variance (ADONIS, “Bray” distance) was performed to define the variation in moth species composition explained (R^2) by crop type and food-bait lure treatment. Additionally, two separate analyses of similarities (“Bray” distance) were conducted to determine differences in moth diversity and abundance based on trap treatment and crop type. Similarities analysis compares the mean rank distances between and within the levels of a factor. If the levels of a factor differ significantly, then the dissimilarities between levels are greater than the dissimilarities within levels (analyses of similarities statistic: R -value = 1.0; $P < 0.05$). Similar analyses were conducted separately, focusing on species from the cutworm and armyworm pest complex to determine the effectiveness of AAMB-baited traps compared to that of unbaited traps (experiment 1) and to determine the diversity of trap capture in response to AAMB lures with additional food-based semiochemical compounds (experiment 2).

For both experiments, moth trap catch was grouped by family and subfamily to compare differences in attraction to food-bait lures by moth taxonomic group. Analyses were conducted on the total number of Noctuidae moths by tribe to compare moth capture in traps baited with the different food-based semiochemicals and that in unbaited traps. Noctuidae moth capture was analysed in a generalised linear mixed model (negative binomial family distribution) with the “glmer.nb” command in the R package, *lme4*, version 1.1-17 (Bates *et al.* 2015). Crop, food-bait lure treatment, and Noctuidae tribe were specified as explanatory fixed variables, and site was specified as a random block factor.

For all statistical analyses, models were first fit as full models, in which the fixed factors included the main effect of all relevant explanatory variables and all possible interactions. For all models, model simplification was performed in a step-wise *a posteriori* procedure by removing nonsignificant interaction terms and comparing nested models through likelihood ratio (LR) χ^2 tests with the “anova” command in the R package, *car*, version 3.0-0 (Fox and Weisberg 2011). The optimal model was selected using Akaike’s information criterion. Test statistic values, degrees of freedom, and P -values were obtained from the “anova” function in the R package, *car*, version, 3.0-0. The “anova” command produces analysis of variance tables from models created by “glmer” commands. Wald χ^2 tests are calculated for linear mixed models, and LR χ^2 are calculated for generalised linear models. A comparison of means for all experiments was performed using Tukey’s method ($\alpha = 0.05$) with the package *lsmeans*, version 2.17 (Lenth and Hervé 2015). All

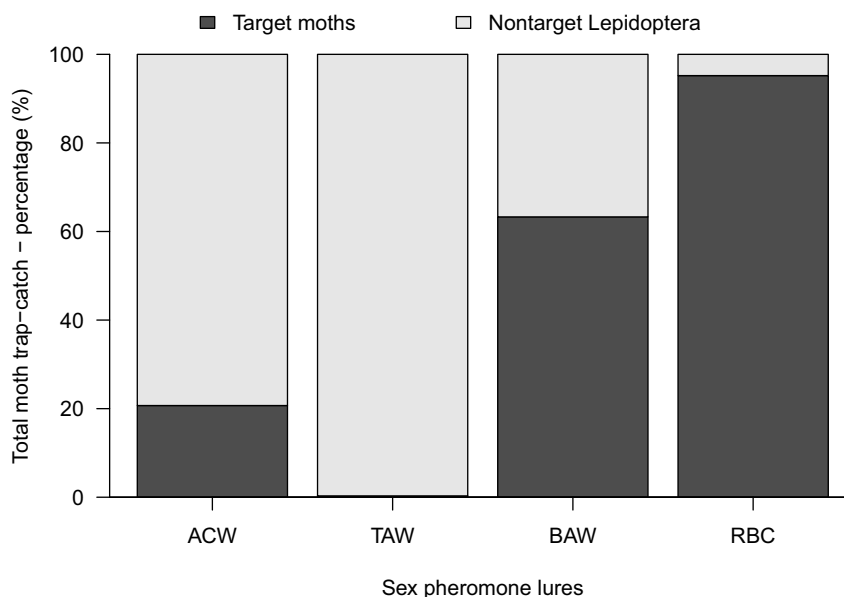


Figure 1. Sex pheromone lure specificity (experiment 1) expressed as a percentage (%) of target species captured in sex pheromone-baited traps from the total moth trap catch. RBC, redbacked cutworm, *Euxoa ochrogaster*; BAW, Bertha armyworm, *Mamestra configurata*; TAW, true armyworm, *Mythimna unipuncta*; ACW, army cutworm, *Euxoa auxiliaris*.

statistical tests were conducted using the freely available statistical package R, version 3.1.0 (R Core Team 2014), in RStudio, version 0.98 (<http://www.rstudio.com>).

Results

Experiment 1: moth diversity in synthetic sex pheromone- and AAMB-baited traps

Specificity of synthetic sex pheromone-baited traps. The redbacked cutworm, *E. ochrogaster*, was the most abundant captured species of the target pests in sex pheromone-baited traps. Redbacked cutworm sex pheromone-baited traps captured, on average, 1077.5 ± 407.3 (standard error) *E. ochrogaster* males per trap per site throughout the sampling period. The Bertha armyworm, *M. configurata*, was the second most abundant target species. Bertha armyworm sex pheromone-baited traps captured, on average, 89.7 ± 33.9 (standard error) *M. configurata* males per trap per site throughout the sampling period. The true armyworm, *Mythimna unipuncta*, and the armyworm cutworm, *E. auxiliaris*, were the least abundant target species. True armyworm sex pheromone-baited traps captured, on average, 1.4 ± 0.5 (standard error) *M. unipuncta* males per trap per site, whereas army cutworm sex pheromone-baited traps captured, on average, 0.4 ± 0.4 (standard error) *E. auxiliaris* males per trap per site.

No significant differences were found in the specificity of sex pheromone lure-baited traps between canola fields and wheat fields (Wald $\chi^2 = 0.29$, $df = 1$, $P = 0.591$). However, the specificity of the sex pheromone lures differed significantly among cutworm or armyworm species (Wald $\chi^2 = 242.47$, $df = 3$, $P < 0.001$; Fig. 1). Redbacked cutworm sex pheromone lures had the highest specificity, with which *E. ochrogaster* represented 95.2% of the total moth trap catch. *Plusia putnami* Grote (Lepidoptera: Noctuidae) was the most abundant nontarget species captured in redbacked cutworm sex pheromone-baited traps (3%; Supplementary material, Table S1). *Plusia putnami* male moths were captured in redbacked cutworm sex pheromone-baited traps in early summer, from 30 June to 5 August 2014, whereas *E. ochrogaster* male moths were captured later in the season, from 22 July to 10 October 2014. Bertha armyworm sex pheromone

lures were less specific, with only 63.3% of the total trap capture being *M. configurata*. *Apamea cogitata* (Smith) (Lepidoptera: Noctuidae) was the most abundant nontarget species in Bertha armyworm sex pheromone-baited traps (26%; Table S1). Both *M. configurata* and *A. cogitata* were captured in Bertha armyworm sex pheromone-baited traps from 24 June to 29 July.

The army cutworm sex pheromone lure had low specificity, with *E. auxiliaris* representing only 28.4% of the total moth trap catch. This low specificity, however, may be a result of the low population density of the target species across all fields. Army cutworm sex pheromone-baited traps captured, on average, 2.1 ± 1.8 (standard error) moths per trap across all sites throughout the season. True armyworm sex pheromone lures had the lowest specificity of the tested lures, with *M. unipuncta* representing only 0.3% of the total moth trap catch. True armyworm sex pheromone-baited traps captured high numbers of *E. ochrogaster* (67%), *Helatrophia reniformis* Grote (Lepidoptera: Noctuidae) (12%), *A. inficita* Walker (10%), and *Anarta trifolii* (Hugnagel) (Lepidoptera: Noctuidae) (4%; Supplementary material, Table S1). The low capture of *M. unipuncta* male moths in sex pheromone-baited traps may be a result of the low specificity of the synthetic pheromone lure and the low population density of the target species across all fields.

Diversity of moths captured in AAMB-baited traps. The AAMB-baited traps captured, on average, 70.6 ± 18.1 (standard error) moths per trap per site throughout the sampling period, whereas unbaited traps captured 18.4 ± 10.3 (standard error) moths per trap per site. In total, 54 macro-Lepidoptera species were captured in the AAMB-baited and the unbaited traps (Supplementary material, Table S2). Crop type explained only 5% of the variation in species composition (ADONIS $R^2 = 0.03$; $P = 0.34$), whereas lure type explained 15% of the variation in species composition (ADONIS $R^2 = 0.15$; $P = 0.001$). No significant difference was observed in the species composition of moths captured in traps deployed in canola fields compared to in wheat fields (analysis of similarities $R = 0.01$; $P = 0.267$). Only one species, *Mythimna oxygala* (Grote) (Lepidoptera: Noctuidae), was more abundant in traps positioned in wheat fields than in canola fields. The AAMB-baited traps captured a higher number of species (canola: 35 spp.; wheat: 42 spp.) than unbaited traps did (canola: 28 spp.; wheat: 21 spp.; analysis of similarities $R = 0.365$; $P = 0.001$; Fig. 2A).

Moths within the Noctuidae family were the most diverse and abundant group attracted to the AAMB-baited traps (47 spp.). Other moth families included Drepanidae (*Habrosyne scripta* Gosse), Erebididae (*Ctenucha virgata* Esper), and Sphingidae (*Darapsa choerlus* Cramer); however, moths from these families were captured in low numbers in AAMB-baited traps (Supplementary material, Table S2). Noctuinae moths were the most diverse and abundant subfamily attracted to the AAMB lure (44 spp.), whereas only a few specimens from Plusiinae (*Anagrapha falcifera* Kirby) and Acronictinae (*Acronicta americana* Harris and *Acronicta superans* Guenée) were captured. Moths from eight Noctuinae tribes were captured in the AAMB-baited traps (Fig. 3A and B). Apameini moths were the most diverse and abundant tribe captured, followed by Noctuini moths. Moths from the tribes Leucaniini, Eriopygini, Tholerini, and Hadenini were trapped in lower numbers, and Xylenini and Caradrini moths were the least represented tribes. The most abundant species were *A. cogitata* (15.4%), *Enargia decolor* (Walker) (14%), *Amphipoea interoceanica* Smith (12.4%), *Feltia jaculifera* (Guenée) (7.9%), *E. ochrogaster* (7.9%), *Helatrophia reniformis* (Grote) (5.3%), and *Apamea devastator* (Brace) (3.1%).

An independent analysis compared the total numbers of Noctuinae moths by tribe captured in the AAMB-baited traps and the unbaited traps. The AAMB-baited traps captured more Noctuinae moths in wheat fields (66.4 ± 17.3 (standard error) moths/trap/site) than in canola fields (37.6 ± 7.5 (standard error) moths/trap/site), whereas unbaited traps captured similarly low numbers of Noctuinae moths in both crops (crop $\times$ lure type, Wald $\chi^2 = 8.78$, $df = 1$, $P = 0.003$). The AAMB-baited traps captured a significantly higher abundance of Noctuinae moths than

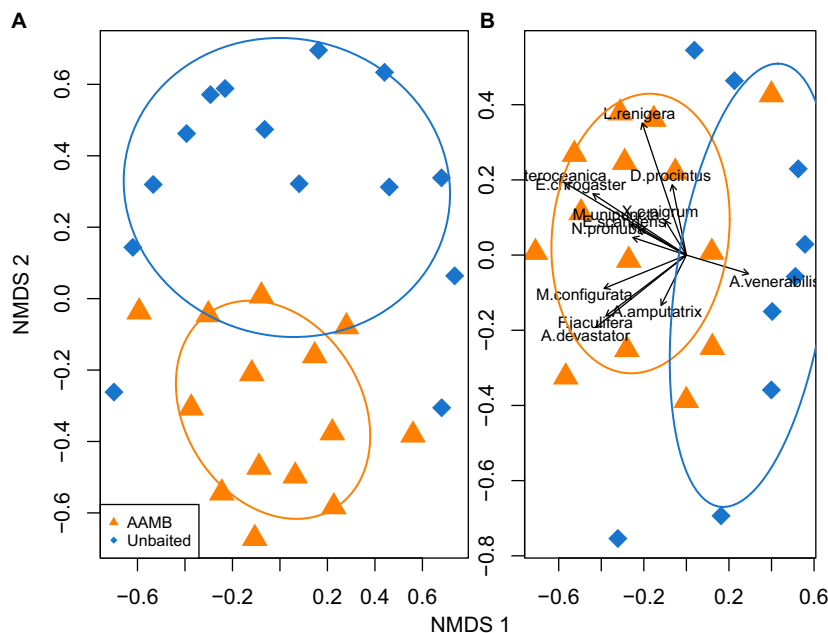


Figure 2. Nonmetric multidimensional scaling for the diversity of moths attracted to the acetic acid and 3-methyl-1-butanol (AAMB) lure compared to unbaited traps: **A**, all noctuid moth species (NMDS, stress = 0.265; $R^2 = 0.511$), and **B**, cutworm and armyworm species only (NMDS, stress = 0.209; $R^2 = 0.656$).

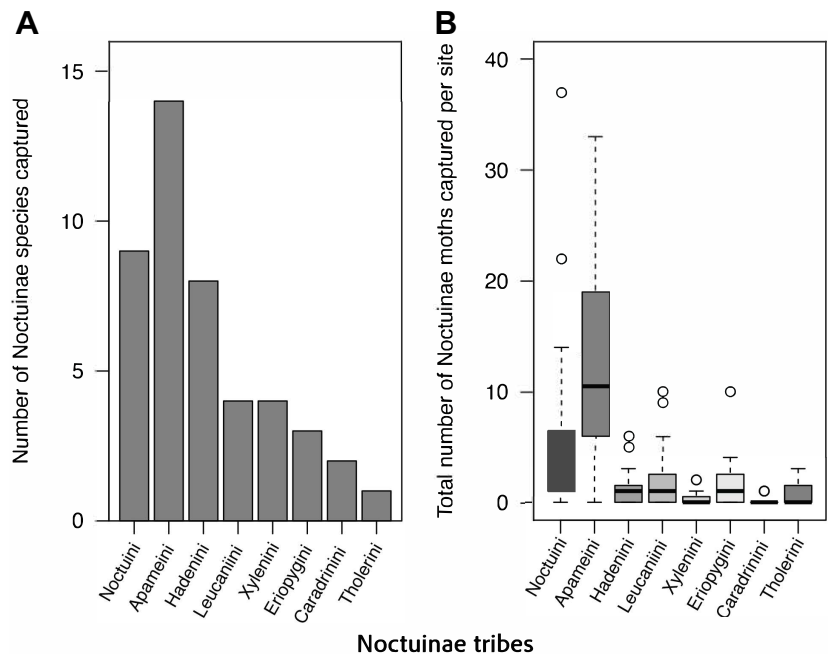


Figure 3. Diversity and abundance of Noctuidae moth tribes captured in traps baited with acetic acid and 3-methyl-1-butanol (AAMB): **A**, barplot of the total number of Noctuidae species by tribe, and **B**, boxplot of the total number of Noctuidae moths by tribe. The midline indicates the median; the top and bottom of the box indicate the first and third quartiles, respectively; the vertical line or whiskers represent the maximum value or 1.5 interquartile range of the data.

unbaited traps did, except for moths from the Caradrinini tribe, which were found in similarly low numbers in both trap types (lure type $\times$ tribe, Wald $\chi^2 = 73.23$, $df = 7$, $P > 0.001$).

The AAMB-baited traps captured a significantly higher diversity and abundance of species of the cutworm and armyworm pest complex than did the unbaited traps (analysis of similarities $R = 0.400$; $P = 0.001$; Fig. 2B). These pests represented on average 42.60% of the total moth trap catch in the AAMB-baited traps per site. The most abundant pest species included the strawberry cutworm, *A. interoceana* (13.3%), dingy cutworm, *F. jaculifera* (9.5%), redbacked cutworm (8.9%), glassy cutworm, *A. devastator* (2.8%), true armyworm (2.9%), and bronzed cutworm, *Nephelodes minians* Guenée (5.2%). Less abundant species ($< 2.0\%$) captured were the Bertha armyworm, bristly cutworm (*Lacinipolia renigera* (Stephens)), spotted cutworm (*Xestia c-nigrum* Linnaeus), the invasive pest winter cutworm (*Noctua pronuba* Linnaeus), white cutworm (*Euxoa scandens* (Riley)), yellow-headed cutworm (*Apamea amputatrix* (Fitch)), and olive-green cutworm (*Dargida procinctus* Grote). Dusky cutworm, *Agrotis venerabilis* Walker, was also captured in the AAMB-baited traps in low numbers; however, more dusky cutworm moths were found in unbaited than in AAMB-baited traps.

Experiment 2: moth diversity in traps baited with food-based semiochemicals

Moth trap catch significantly differed among the food-bait lure traps and unbaited traps (Wald $\chi^2 = 269.63$, $df = 4$, $P < 0.001$). Traps baited with AAMB alone and with AAMB + MP lures captured twice the total number of moths than did traps baited with AAMB + PAA lures or with AAMB + MP + PAA lures, and unbaited traps captured the fewest moths (Fig. 4A). Traps baited with AAMB alone and with AAMB + MP lures captured, on average, 173.4 ± 45.34 (standard error) and 163.25 ± 46.41 (standard error) moths per trap per site, respectively, whereas traps baited with AAMB + PAA lures and with AAMB + MP + PAA lures captured 82.5 ± 19.4 (standard error) and 83.1 ± 21.91 (standard error) moths per trap per site, respectively. The unbaited traps captured, on average, 12.4 ± 1.5 (standard error) moths per trap per site throughout the sampling period. In total, 76 macro-Lepidoptera species were captured in all traps across all sites (Supplementary material, Table S3). Crop type explained 2% of the variation in species composition (ADONIS $R^2 = 0.02$; $P = 0.059$), and lure type explained 16% of the variation in species composition (ADONIS $R^2 = 0.16$; $P = 0.001$). No significant difference was found between the species composition of moths captured in traps in canola fields and traps in wheat fields (analysis of similarities $R = 0.00$; $P = 0.282$). Food-bait lure traps captured a significantly higher number of species and abundance of moths compared to what unbaited traps captured (analysis of similarities $R = 0.20$; $P = 0.001$); however, species composition of moths did not vary among the traps baited with the different food-bait lures (analysis of similarities $R = 0.01$; $P = 0.287$; Fig. 4B).

Moths within the Noctuidae family were the most diverse and abundant group found to be attracted to food-bait lures (67 spp.). Other families included Erebidae (3 spp.), Sphingidae (2 spp.), Cambridae (1 sp.), Geometridae (1 sp.), and Hesperidae (1 sp.); however, few individuals from these families were captured in the traps with different food-bait lure combinations (Supplementary material, Table S3). Noctuinae moths were the most diverse subfamily attracted to the food-bait lures (62 spp.), whereas only a few specimens from Plusiinae (4 spp.) and Acronictinae (1 sp.) were captured in the baited traps (Supplementary material, Table S3). Moths from eight Noctuinae tribes were captured in the different food-bait lure traps. Apameini was the most diverse and abundant tribe, followed by Noctuini and Eriopygini. Several species from the tribes Leucaniini, Tholerini, and Hadenini were trapped, and very few species in the Xylenini and Caradrini tribes were captured (Supplementary material, Fig. S2).

An independent analysis compared the total numbers of Noctuinae moths by tribe captured in the different food-bait lure traps and unbaited traps placed in canola and wheat fields. A marginally significant interaction was revealed between crop and lure type (crop $\times$ lure type,

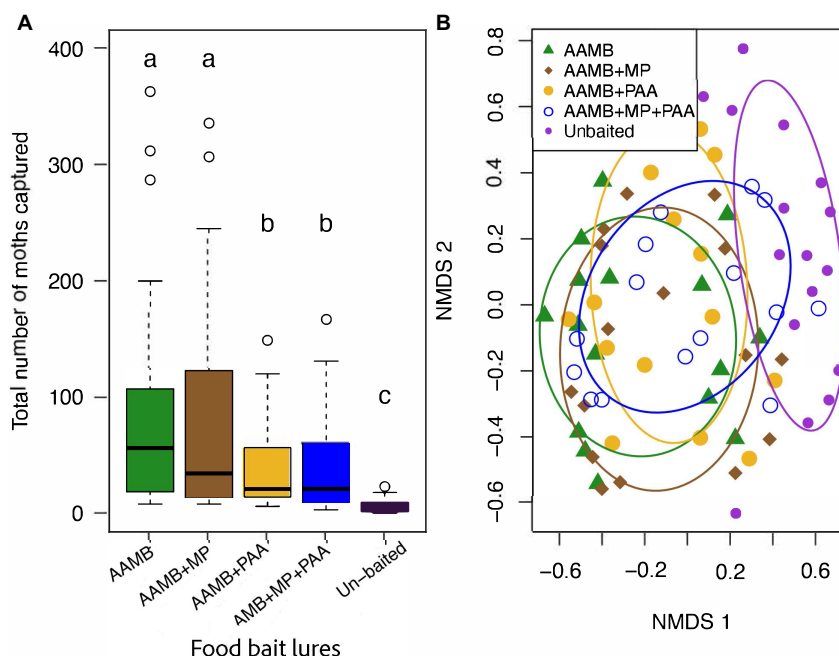


Figure 4. **A**, Boxplots of the total number of moths captured in acetic acid and 3-methyl-1-butanol (AAMB) lures with additional food-based semiochemical compounds. The midline indicates the median; the top and bottom of the box indicate the first and third quartiles, respectively; the vertical line or whiskers represent the maximum value or 1.5 interquartile range of the data. Boxplots marked with different letters are statistically different (Tukey method, $\alpha = 0.05$). **B**, Nonmetric multidimensional scaling (NMDS, stress = 0.214; $R^2 = 0.7182$) for the diversity of moths attracted to AAMB lure with and without additional food-based semiochemical compounds. Tested chemicals were an alcohol from fermented byproducts, 2-methyl-1-propanol (MP), and a floral volatile, phenylacetaldehyde (PAA). Treatments included AAMB alone, AAMB + MP, AAMB + PAA, AAMB + MP + PAA, and an unbaited trap that served as control.

Wald $\chi^2 = 9.40$, $df = 4$, $P = 0.052$) that affected the total Noctuid moth trap catch. In both canola and wheat fields, Noctuid captures in traps baited with the various food-bait lures were significantly higher than those in unbaited traps. Interestingly, the different food-bait lures captured similar numbers of Noctuid moths in canola fields, whereas, in wheat fields, more Noctuid moths were captured in traps baited with either AAMB or with AAMB + MP than in traps baited with AAMB + PAA or with AAMB + MP + PAA.

A significant interaction was found between the crop and Noctuid tribe (crop $\times$ tribe, Wald $\chi^2 = 17.48$, $df = 7$, $P = 0.016$) on moth capture. The number of moths captured from different Noctuid tribes depended on the type of crop in which the traps were positioned. More Apameini, Eriopygini, and Leucaniini moths were captured in traps located in wheat fields than in traps located in canola fields, whereas moths from the tribes Noctuid, Hadenini, Tholerini, Xylenini, and Caradrini occurred equally in traps positioned in both crops. Lastly, a significant response of Noctuid moths to food-bait lures was observed depending on the tribe (lure type $\times$ tribe, Wald $\chi^2 = 130.91$, $df = 28$, $P < 0.001$). The AAMB and the AAMB + MB lures attracted more Apameini, Hadenini, and Tholerini moths than did traps baited with fermented byproducts food-bait lures with floral volatiles (Supplementary material, Fig. S3A, E, and F). Noctuid, Eriopygini, and Leucaniini moths were similarly captured in all traps baited with food-bait lures (Supplementary material, Fig. S3B, C, and D). The number of Caradrini and Xylenini moths did not differ between the food-bait lures and the unbaited traps (Supplementary material, Fig. S3G and F).

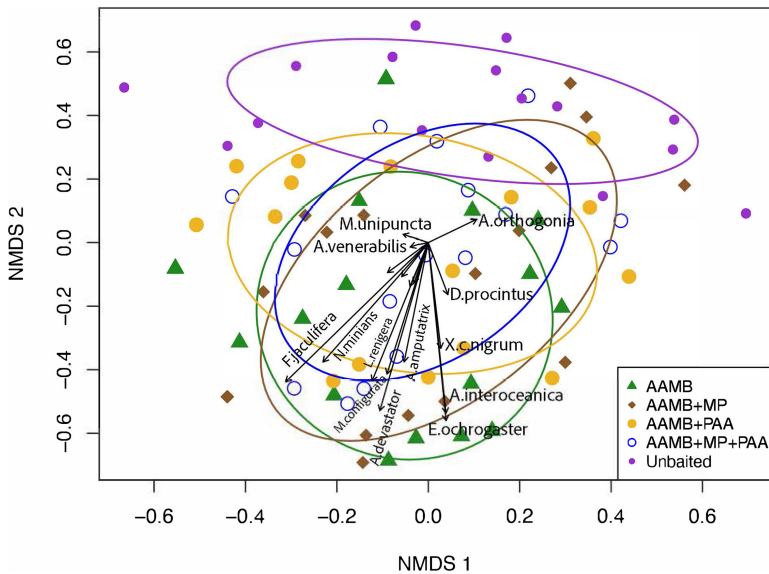


Figure 5. Nonmetric multidimensional scaling (NMDS, stress = 0.193; $R^2 = 0.765$) for the diversity of cutworm and armyworm species attracted to acetic acid and 3-methyl-1-butanol (AAMB) lure with and without additional chemical compounds. The tested chemicals were an alcohol from fermented byproducts, 2-methyl-1-propanol (MP), and a floral volatile, phenylacetaldehyde (PAA). Treatments included: AAMB alone, AAMB + MP, AAMB + PAA, AAMB + MP + PAA, and an unbaited trap that served as control.

Traps baited with AAMB lures with additional food-based chemicals captured a significantly higher number of species in the cutworm and armyworm pest complex than did unbaited traps (analysis of similarities $R = 0.16$; $P = 0.001$). Traps baited with AAMB and with AAMB + MP lures captured significantly higher proportions of pest species (43.1% and 46.1%, respectively) out of the total moth trap catch than did traps baited with AAMB + PAA or with AAMB + MP + PAA lures (41.7% and 37.2%, respectively; Wald $\chi^2 = 17.36$, $df = 3$, $P < 0.001$). Several cutworm species, however, were more attracted to AAMB and to AAMB + MP lures than they were to lures containing phenylacetaldehyde, including the dingy cutworm, redbacked cutworm, glassy cutworm, strawberry cutworm, and bronzed cutworm (Fig. 5).

Discussion

At least 2308 lepidopteran species occur in Alberta, representing 17% of the known lepidopteran diversity in North America (Pohl *et al.* 2010). There are 402 macromoths species recorded in the grassland in Alberta (Pohl *et al.* 2010). In the present study, traps baited with food-bait lures based on volatiles from byproducts of fermented sugar baits captured 67 lepidopteran species, which represent 16% of the recorded macromoth species in this ecoregion. In the Prairie Ecozone, parkland and grasslands are particularly rich in noctuid moth species, and many species do not occur elsewhere in Canada (Pohl *et al.* 2010). Only a small portion of the parkland and grassland ecoregions remain in a natural state due to modification of the landscape for agriculture and pastureland. It is therefore important to monitor lepidopteran diversity and abundance in this disturbed landscape.

The specificity of the commercially available synthetic sex pheromone lures varied among the tested pest species. The sex pheromone lures of the redbacked cutworm and Bertha armyworm were the most specific to capturing the targetted pests. As expected, the sex pheromone-baited traps captured larger numbers of target moths than did any of the food-bait lures; however,

farmers did not report damage the following crop season. The sex pheromone lures of the army cutworm and true armyworm had the lowest specificity in attracting the target species; however, this may be a result of the low population density across all sampled fields. Although female-produced sex pheromones are known for most pest species in the region (Steck *et al.* 1982), pheromone-based monitoring of several cutworm and armyworm pest species has been abandoned. Monitoring programmes with sex pheromone-baited traps were implemented across the Prairie Provinces in the 1980s; however, these programmes were discontinued because moth trap catch does not reflect the local population density or relate to crop damage (Byers and Struble 1987). Moths are attracted to sex pheromone-baited traps from farther distances (Ayre and Lamb 1990), and multiple trap-lure systems are required to monitor all pest species. The tested synthetic sex pheromone lures attracted little nontarget moth diversity. A generalised food-bait lure in a single trap that attracts multiple noctuid pest species can also provide important data on the diversity of nonpest moth species in the agroecosystem and on the local population density of pest species.

Moths in the family Noctuidae were the most diverse and abundant lepidopteran taxon captured in AAMB-baited traps in experiment 1. Traps baited with AAMB lures attracted only a few individuals from a few species in the families Erebididae, Geometrididae, Sphingidae, Cambridae, and Hesperiididae. The distribution of lepidopteran taxa sampled with AAMB-baited traps, however, is representative of the proportion of species in each family of Lepidoptera present in grassland habits within the Prairie ecozones. Noctuidae represent 80% of the total macro-Lepidoptera diversity in grasslands habitats on the Canadian Prairies, followed by Erebididae with 10% and Geometrididae with 9% (Pohl *et al.* 2014). Furthermore, the breadth of lepidopteran taxa captured in AAMB-baited traps in Canadian prairie agroecosystems is similar to that of the lepidopteran taxa captured in AAMB-baited traps in apple orchards in Washington state, United States of America (Landolt and Hammond 2001), and in horticultural gardens in Alaska, United States of America (Landolt and Hammond 2001). Moths from different families differ in responsiveness to specific cues in the environment and to specific fermented sugar bait byproducts. Traps baited with ethyl alcohol, acetoin, or β -phenyl alcohol attract geometrid moths in a mixed forested area, whereas acetic acid and 3-methyl-1-methanol lures attract noctuid moths in the same ecosystem (Utrio and Eriksson 1977). The response of lepidopteran families to different volatile compounds from fermented sugar baits is due to their olfactory sensitivity and their ability to use microbial volatile organic compounds as semiochemicals to locate food resources (Davis *et al.* 2013). Our experiments show that AAMB lures in the Prairie ecozone attract mostly noctuid moths.

Noctuinae moths were the most diverse and abundant noctuid subfamily attracted to AAMB lures in the present study, with the majority of species and the highest number of captured moths being from the tribes Apameini and Noctuini. *Apamea cogitata* was the most commonly captured species in AAMB-baited traps in 2014, and it was also the most frequently captured noctuid species in traps baited with AAMB lures in Alaska (Landolt *et al.* 2007). The AAMB-baited traps captured several cutworm and armyworm pest species in higher numbers than unbaited traps did in 2014 during the present study. Similarly, AAMB-baited traps captured redbacked cutworm, dingy cutworm, glassy cutworm, spotted cutworm, yellow-headed cutworm, and the olive-green cutworm in noctuid moth surveys in Washington state and Alaska (Landolt and Hammond 2001; Landolt *et al.* 2007). Those surveys, however, did not include unbaited control traps to determine if AAMB lures actually attract captured moths.

In the present study, AAMB lures, alone or with additional semiochemicals, captured more moths in wheat fields than in canola fields. This difference may be influenced by the host plant volatiles released by the crops in the background where the baited traps were positioned. Acetic acid is a dominant volatile organic compound emitted by canola plants at the flowering stage (Veromann *et al.* 2013), whereas acetic acid is present at low concentrations (Piesik *et al.* 2010, 2011). Traps baited with AAMB lures may be more apparent to moths in wheat

fields than in canola fields because the lure stands out from the background volatiles emitted from the crop. No difference was found, however, between the composition of moth species captured in the two crop types. This finding suggests a common noctuid moth community assemblage across agroecosystems in central Alberta and possibly significant dispersal throughout agricultural landscapes. Noctuid moths are strong flyers that disperse over long distances and can migrate into the Prairie Provinces in the summer. Sporadic infestations of the true armyworm and black cutworm, *Agrotis ipsilon* (Hufnagel) (Lepidoptera: Noctuidae), occur in the Prairie Provinces as a result of early season migration (Beirne 1971; Fields and McNeil 1984). Mark-recapture experiments illustrate that some noctuid species disperse over great distances. For instance, marked male and female *Spodoptera frugiperda* (Smith) (Lepidoptera: Noctuidae) were recaptured 806 m and 608 m from the release point in fields of corn, *Zea mays* Linnaeus (Poaceae), respectively (Vilarinho *et al.* 2011). *Heliothis virescens* (Fabricius) (Lepidoptera: Noctuidae) can disperse up to 30 km from the release point (Schneider 1999).

The results of adding 2-methyl-1-propanol and phenylacetaldehyde to AAMB lures in 2015 showed a pattern similar to the results of the survey conducted in 2014. A similar number of cutworm and armyworm species were found in traps baited with the different food-bait lure types; however, more noctuid pests were attracted to food-bait lures from fermented byproducts than to those with floral volatiles, specifically the dingy cutworm, redbacked cutworm, glassy cutworm, strawberry cutworm, bronzed cutworm, and yellow-headed cutworm. Similar patterns have been reported previously for the glassy cutworm and dingy cutworm (Landolt *et al.* 2011). Traps baited with AAMB alone or with the additional alcohol component captured more moths than did traps containing phenylacetaldehyde mainly because *A. cogitata* was not captured in traps baited with the floral volatile. Likewise, Landolt *et al.* (2011) reported more *A. cogitata* were captured in AAMB-baited traps than in floral volatile-baited traps in forested riparian sites adjacent to pastures. Although the information on the biology and adult foraging behaviour of *A. cogitata* is limited, its larvae are known to feed on grasses (Mikkola *et al.* 2009). Adult *A. cogitata* may not respond to floral volatiles because this type of semiochemical is not commonly released from their grass host inflorescence. In contrast, alcohol and acids are part of the volatile profile of several hosts in the Poaceae family. For example, winter wheat, *Triticum aestivum* Linnaeus, emits methanol and acetaldehydes throughout the growing season, whereas volatiles of acetic acid are mainly captured from the environment, and *Triticum* spp. lack phenylacetaldehyde (Bachy *et al.* 2020). *Apamea cogitata* appears to be more attracted to volatiles acetic acid and alcohols as cues to locate potential grass hosts.

Our results demonstrate a difference in attraction to food-bait lures by subfamilies within Noctuidae. This differential attraction to the different food-based semiochemicals reflects varying foraging strategies used by moths in the different tribes and their interactions with plants for nectar resources or host selection (Landolt *et al.* 2007). In some instances, moths associate floral scents with sexual attraction because some male sex pheromones include floral volatiles such as phenylacetaldehyde (Knudsen and Tollsten 1993). Food-bait lures based on phenylacetaldehyde are attractive to species from the subfamilies Plusiinae and Heliothinae, whereas food-bait lures based on isoamyl alcohols are more attractive to species in the subfamily Noctuinae and Hadeninae (Szanyi *et al.* 2017).

Plants with generalised or mixed pollination systems can attract pollinators when they are active with temporal and diurnal variation in floral scent production (Jürgens *et al.* 2014). Some moth-pollinated plant species release floral scents at dusk or in the early evening to coincide with moth flight (Knudsen and Tollsten 1993). Alternatively, volatile organic compounds produced by microbes associated with nectar (Davis *et al.* 2013) can attract generalist pollinators (Schiestl *et al.* 2006). Yeasts use sugars in flower nectar, and the volatile fermented byproducts of this interaction, which include acetic acid and alcohols, alter the floral volatile profile (Pozo *et al.* 2020). Noctuinae moths may employ microbial volatile organic compounds as a more reliable cue for foraging because floral volatiles may not be present when they are active. Food-bait lures from fermented byproducts therefore may be a more effective lure for monitoring a greater diversity of Noctuinae pests and nonpest species in prairie agroecosystems.

Overall, the present study shows the broad attractiveness of AAMB to a large number of noctuid moths and noctuid pest species. We hypothesised a similar noctuid moth species composition in canola and wheat fields. In both experiments, we observed no difference in the noctuid species composition in the two crop types. These findings reflect a generalised moth community in human-managed ecosystems due to the annual level of disturbance, crop rotation practices, and strong dispersal capabilities of moths. Traps baited with food-bait lures based on fermented byproducts can be used to monitor Noctuidae diversity in agroecosystems (Süsslenbach and Fiedler 1999), including those in the Canadian Prairies. We hypothesised that noctuid moths would be more attracted to traps baited with volatiles from fermented sugar baits than to traps baited with floral volatiles. The addition of phenylacetaldehyde did not enhance the attraction of noctuid pest species, and some cutworm species were less attracted to food-bait lures with floral volatiles or were potentially repelled by these cues. These findings indicate that noctuid moths use microbial volatile organic compounds as semiochemicals to locate food resources.

Insect biomass and diversity are in decline due to human-driven modification of habitats and climate change (Hallmann *et al.* 2017; Sánchez-Bayo and Wyckhuys 2019; Laussmann *et al.* 2021). Most parkland and grassland ecoregions have been modified to extensive agriculture and pasture, which could lead to homogenisation of the insect fauna. Unless biodiversity is continually surveyed, the impacts of human-caused disturbance cannot be reliably detected. Implementing baited traps with a generalised food-bait lure to monitor the presence, abundance, and flight activity of noctuid species can help to detect changes in lepidopteran diversity and abundance in grasslands and prairie agroecosystems.

Supplementary material. The supplementary material for this article can be found at <https://doi.org/10.4039/tce.2025.10014>.

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Competing interests. The authors declare that they have no competing interests.

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