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Potential causes and consequences of decreased body size in field populations of *Coccinella novemnotata*

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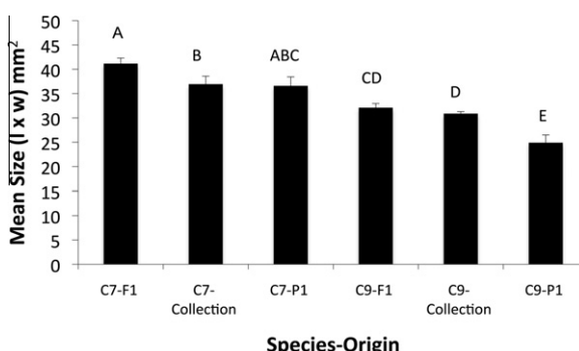
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HIGHLIGHTS

- ▶ Recent *Coccinella novemnotata* were significantly smaller than historical specimens.
- ▶ Offspring fed *ad libitum* prey were not smaller than historical specimens.
- ▶ Varying prey availability alone was sufficient to reproduce observed size variation.

GRAPHICAL ABSTRACT



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ABSTRACT

Coccinella novemnotata, the nine-spotted lady beetle, was historically one of the most common lady beetles across the US and southern Canada. In the 1980s it became extremely rare and has remained rare. In 2008 adult *C. novemnotata* were collected from field populations in Oregon and South Dakota and initial observations suggested that these individuals seemed smaller than the mean size of the species historically. These observations led to a series of experiments to determine if there had been significant decrease in size and if any decrease found was due to a genetic change or to environmental factors. In the first of these studies we quantified the size of *C. novemnotata* collected in the field and the size of *Coccinella septempunctata*, a congeneric introduced species that was collected in the same habitats and has been implicated as a cause for *C. novemnotata* decline. The size of these field-collected individuals of both species was compared with the size of historical specimens and individuals reared in the laboratory. Field-collected *C. novemnotata* adults were significantly (20%) smaller than specimens bred in captivity and specimens from collections. To determine if prey limitation alone could yield the range of sizes observed we reared larvae across a range of prey availability. There was a significant effect of prey availability and adult sizes across treatments bracketed the range we found in the field. Low fed larvae are significantly smaller than high fed larvae. While these results do not definitively point to any single explanation for the decline of this species they are consistent with expectations for competition between *C. novemnotata* and *C. septempunctata*.

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1. Introduction

1.1. Background

Body size is correlated with many determinants of fitness (Blueweiss et al., 1978) including fecundity (Honek, 1993), survival (Savage et al., 2004), and the ability to secure resources such as prey (Menge, 1972) and mates (Howard et al., 1997). The study of why some species' average body sizes have changed over time while others have not has the potential to shed light on the impact of several broad-scale long-term environmental changes including climate change (Roy, 2008; Smith et al., 1995) and interactions with invasive species (Brown and Maurer, 1986; Phillips and Shine, 2004). If the focal species is a biological control agent then changes in body size may impact its ability to suppress pest populations. If a species that was introduced as a biological control agent is partly or fully responsible for the size change in the focal species then there are potentially serious nontarget issues that need to be addressed (Phillips and Shine, 2004).

Once it has been determined that the mean size of individuals of a focal species has changed significantly over time beyond non-time dependent levels of variation (e.g. clines), discovering the reason for the change can offer insights into how the organism interacts with its changing environment. Comparing size changes of multiple species from similar habitats can allow additional insights by discerning why some species are impacted while others are not. Several factors could account for a shift in mean size over time, including (1) a change in resource availability, (2) infection with a pathogen, (3) size-dependent mortality, or (4) a genetic shift to a different size. Ecological factors (1–3 above) can be separated from genetic changes by manipulating ecological conditions while controlling for genetic factors (Naguib and Gil, 2005).

While the causes and consequences of body size have been studied for almost all animal taxa, one group where clear effects have been demonstrated is the insects. Both underlying genetic variation (Dingle, 1984) and environmental conditions during development (Gullan and Cranston, 1994; Hodek and Honek, 1996) have been shown to impact adult size in insects. Within this class of animals, prey availability during development, specifically, has been shown to impact body size in coccinellids (Blackman, 1965; Honek, 1993; Obrycki and Orr, 1990; Orr and Obrycki, 1990; Phoofolo et al., 2008).

One species of coccinellid of great current interest is the nine-spotted lady beetle *Coccinella novemnotata* Herbst. Up until the mid-1980s, this was the most prevalent species of lady beetle in the northeastern United States (Harmon et al., 2007). It served as an important biological control agent, moderating aphid populations in gardens and crops throughout the northeast. Yet, in 1993, an extensive USDA APHIS coccinellid survey found no evidence of the nine-spotted lady beetle in eleven Northeastern states. This cooperative study focused on 100 counties and was based on comprehensive fieldwork and data from personal collections. The most recent published records report that the nine-spotted lady beetle was last collected on the eastern seaboard in 1992 (Harmon et al., 2007). For more than a decade, entomologists have pondered the disappearance of *C. novemnotata*, but speculated that it might continue to persist in low densities across the United States (Wheeler and Hoebeke, 1995). The initial decline in the *C. novemnotata* population in eastern North America appears to align with the introduction of the European, seven-spotted lady beetle (*Coccinella septempunctata*) (Wheeler and Hoebeke, 1995). The seven-spotted lady beetle has since proven to be a powerful alternative to chemical management of agrarian pests (Hoffmann and Frodsham, 1993), but it may have had a negative impact on native species of coccinellids (Alyokhin and Sewell, 2004; Elliot

et al., 1996; Ellis et al., 1999; Simberloff and Stiling, 1996; Snyder et al., 2004; Staines et al., 1990; Wheeler and Hoebeke, 1995). Several alternative hypotheses have been proposed to explain the decline of *C. novemnotata* (e.g. land use change, climate change) that do not cite the introduction of *C. septempunctata* as the cause. These theories are logical in principle, but do not appear to align directly with the known time period for *C. novemnotata* disappearance (see Harmon et al., 2007).

The primary challenge in reaching a definitive conclusion regarding the effects of *C. septempunctata* on *C. novemnotata* is that very little data were systematically collected as *C. septempunctata* expanded and *C. novemnotata* contracted its range (Gordon, 1985; but see Evans, 2000). The Lost Ladybug Project (www.lostladybug.org), founded in 2000, utilizes citizen science to track the distribution and abundance of lady beetles across North America. It is the first initiative that attempts to fill the void left by the historical lack of methodological surveys. Since its inception, the Lost Ladybug Project has collected over 11,000 lady beetle specimens, and its investigators have established sustainable colonies of rare coccinellids. In 2006, after a 14-year lapse of sighting, a nine-spotted lady beetle (*C. novemnotata*) was discovered in Washington, DC (Losey et al., 2007). A grant from the National Science Foundation has allowed the Lost Ladybug Project to broaden its outreach and, since 2008, the initiative has logged images of over 40 individual *C. novemnotata* and collected over 30 live specimens, exclusively from the western states. Cornell University is home to one of just two colonies of *C. novemnotata* in the United States. Initial observations of the live *C. novemnotata* that were collected by the Lost Ladybug Project in 2008–2009 suggested that the individuals appeared to be notably smaller than historical specimens while the mean size of individuals of its congeneric potential competitor, *C. septempunctata*, did not appear changed.

1.2. Objectives

We initiated this study by assessing if the average size of *C. novemnotata* adults was indeed declining. To accomplish this we determined if the size of current *C. novemnotata* populations fell within the historical size range of individuals before the species declined. To investigate the causes and consequences of size change in *C. novemnotata* individuals in field populations we then compared the sizes of recently collected *C. novemnotata* and *C. septempunctata* specimens to the offspring of field-collected adults which have been reared in the laboratory with access to *ad libitum* prey. Finally, to ascertain if prey limitation could be the sole cause of size decline we set out to determine if the range in size of recently collected adult *C. novemnotata* could be duplicated by varying access to prey.

2. Methods

2.1. Rearing lady beetles

Adult *C. novemnotata* were placed in a separate colony and allowed to mate freely. All beetles, regardless of developmental phase, were placed on a 16 h light cycle, and lab temperature was set at a consistent 21 °C. Eggs were collected from each colony and observed for signs of fertilization. Coccinellid eggs are normally bright yellow to orange in color. Viable eggs retain the coloration, while nonviable eggs shrivel and turn dark brown or black within 2–3 days of oviposition. Clusters of viable eggs were placed in cylindrical, plastic containers, each with a volume of 44 ml. Each container stored a single egg cluster. Each cluster yielded approximately 3–7 individual first-instar larvae after approximately four

Table 1
The number of beetles measured from each origin.

Origin	Species	Sample size	Mean area ($l \times w$) \pm SEM	Location	Date collected
Field collected (P1)	<i>C. novemnotata</i>	8	24.82 \pm 2.29	OR	June 10–11, 2009
Field collected (P1)	<i>C. novemnotata</i>	13	24.95 \pm 1.79	SD/NE	June 18, 2008; early July 2009
Field collected (P1)	<i>C. novemnotata</i>	6	29.66 \pm 2.64	UT	Spring 2011
Field collected (P1)	<i>C. novemnotata</i>	11	24.36 \pm 1.95	CO	May 27, 2011
Raised in culture (F1)	<i>C. novemnotata</i>	41	37.66 \pm 1.62	NY (Cornell)	–
Raised in culture (F1)	<i>C. novemnotata</i>	33	25.17 \pm 1.13	SD (USDA)	–
Collection	<i>C. novemnotata</i>	180	30.71 \pm 0.48	NY (CUIC)	–
Collection	<i>C. novemnotata</i>	74	28.09 \pm 0.75	DC (Smith.)	–
Collection	<i>C. novemnotata</i>	50	35.56 \pm 0.91	UT	–
Field collected (P1)	<i>C. septempunctata</i>	16	36.59 \pm 1.62	OR	June 10–11, 2009
Raised in culture (F1)	<i>C. septempunctata</i>	41	41.15 \pm 1.01	NY (Cornell)	–
Collection	<i>C. septempunctata</i>	20	36.77 \pm 0.80	NY (CUIC)	–

days. To provide prey, the lab maintains a mixed colony of *Acyrtosiphon pisum* (pea aphid) and *Myzus persicae* (green peach aphid). The aphids were reared on *Vicia faba* (fava bean), and the aphid colony was kept in a moist growth chamber, at a constant temperature of 23 °C. Feeding practices were kept consistent until larvae reached the third instar. Developmental instar was confirmed by counting the number of moults. All individuals in a container were determined to have moulted when the number of exuviae equaled the number of larvae. No larvae moulted twice before all individuals had completed a single instar. At the third instar, larvae were separated into individual containers, each with a piece of paper towel to be used as an anchor for later pupation. After hatching, it took each larva approximately 4–5 days to reach third instar. Each third instar larva was fed roughly 15 aphids daily. Individuals were closely observed for fungal infection (*Microsporidia*) until pupation. From the onset of the third instar, it took each larva approximately 7 days to pupate. Each individual spends about 1 day in a pre-pupal stage where the larva ceases to eat and appears sluggish. It took approximately 4 days to emerge as adults. All newly emerged adult coccinellids were not fed for 24 h as their elytra hardened. Once they resumed normal feeding habits (roughly 15 aphids daily), adult coccinellids were returned to their respective colonies to mate.

2.2. Measuring lady beetles

Lady beetles for measurement were acquired from (1) field collections {P1} from eastern Oregon, northwestern Nebraska, and southwestern South Dakota, (2) the following insect collections: The Cornell University Insect Collection, The Smithsonian Institution, The Utah State University Insect Collection and, (3) laboratory colonies {F1 and beyond} maintained at Cornell University and at the USDA-ARS Laboratory in Brookings, South Dakota. The number of beetles measured from each origin is listed in Table 1. All of the individuals used for this study were dead. Coccinellids from collections were pinned specimens. All other specimens were stored in 70% isopropyl alcohol.

Measurements were taken using calipers or a microscope micrometer, both with accuracy to 0.01 mm. Individuals taken from storage in alcohol were allowed to dry before dimensions were taken. To enhance visual contrast, coccinellids were placed on white filter paper. The length measurement was taken from the tip of the head to the tip of the abdomen. The width measurement was taken across the widest portion of the insect. For some specimens, height was also measured to allow calculation of volume as, potentially, the most accurate measure of overall size. In order to acquire height measurements, each individual was placed on its side, and the dimension was again calculated across the widest portion of the abdomen. The volume was determined using a formula for hemi-ellipsoid volume: $(4/3)\pi[(\text{Length})/2 \times (\text{Width})/$

$2 \times (\text{Height})]$. Regression analysis confirmed that area ($l \times w$) is an excellent predictor of volume ($R^2 = 0.998$). Since coccinellids are generally ovate, area could be expressed as the area of an ellipse ($l \times w \times \pi$) but since the amount of curvature at both anterior and posterior varies among species and individuals we chose to analyze and present all data as simple rectangular area ($l \times w$). Hereafter, we will use “size” to denote rectangular area ($l \times w$).

2.3. Feeding study

In order to determine the effects of prey availability on ultimate adult size, 104 first-instar *C. novemnotata* larvae were selected from the laboratory colony, and divided into five subgroups to be reared at varying prey levels. Larvae were placed in individual containers within 24 h of hatching with a piece of paper towel as an anchor for eventual pupation. Containers were labeled to represent the appropriate experimental group. Larvae were fed a random mix of pea and green peach aphids at densities of 1, 3, 12, 21, or 30 aphids per 24-h period. All uneaten aphids were removed before new aphids were added, so uneaten aphids did not accumulate in the containers. Twenty replications were set up for each treatment except for prey densities of 1 and 12 for which 22 replications were initiated. The feeding regimens were selected based on preliminary studies to produce variation in adult sizes, while preserving a high level of survivorship at the higher levels. Aphids were replaced every 24-h regardless of the number consumed in the previous period. Developmental stage and all fatalities were noted daily for each individual until each of the original 104 individual beetles died. Upon emergence, adults were fed while their elytra hardened. Within 5 days of emergence, measurements of adult length, width, and height were taken as noted above.

2.4. Statistical analysis

Since sex ratio data was not available for all data sets and since the sex ratio was approximately 50:50 for all groups of adult beetles, sexes are lumped for calculating mean beetle sizes. The relationships between longitude, latitude, year, Julian date, elevation and all their interactions and the size of *C. novemnotata* from museum collections were analyzed individually with a regression analysis on JMP using backward selection starting with the saturated model. The mean size of *C. novemnotata* adults from different geographic locations was compared with *t*-tests adjusted by a *post priori* Bonferroni correction (critical $p = 0.05/\text{number of comparisons}$; Scheiner and Gurevitch, 2001). Since all locations were not significantly different they were grouped. Adult beetle sizes across the two species (*C. septempunctata*, *C. novemnotata*) and three origins (field collected, laboratory-reared, and museum collections) were analyzed using an ANOVA using JMP with experimental error

rates controlled by using Tukey's Honest Significance Test for means separation tests. The relationship between the number of aphids offered per day and adult beetle size was also analyzed with a regression analysis on JMP with treatment levels log transformed to preserve linearity. The relationship between prey level and the proportion of individuals that survive to adulthood was analyzed as a logistic regression between the number of aphids offered per day and the number in each of two binomial states; survival (to adulthood) or death. Because of the parabolic shape of the relationship and because initial regression confirmed the significance of a quadratic term the number of aphids offered was transformed as $X - 0.024X^2$ prior to the analysis. R^2 for the transformed regression was estimated using Efron's R^2 which is equivalent to squaring the correlation coefficient between predicted and observed Y values (Osborne, 2008).

3. Results

3.1. Determining the most appropriate historical comparison

Using a backward selection model-building process we determined the optimal model contained elevation, latitude, longitude and an elevation by latitude interaction. The equation for the optimal model was: $\text{Size} = 26.24 + 0.002(\text{Elevation}) + 0.25(\text{Latitude}) + 0.08(\text{Longitude}) + 0.0003(\text{Elevation} \times \text{Latitude})$. Although a significant relationship was found for this model ($p = 0.0001$) it explains very little of the variation in beetle size ($R^2 = 0.10$). Surprisingly, the relationship between all three factors (elevation, latitude, and longitude) and size was positive. This predicts that beetles will be larger as collection sites move towards higher elevations and in a northwesterly direction. Given that this positive relationship would predict larger rather than smaller beetles for our relatively high elevation samples from the northwestern US and given the low proportion of size variation explained we decided not to include any of the clines or their interactions in further analyses. We deemed it most appropriate to use a pooled mean measurement for all historical *C. novemnotata* specimens as the comparison against our recent samples.

3.2. Size comparisons

Factorial analysis of coccinellid species and origin revealed a significant effect of species ($p < 0.0001$) and origin ($p < 0.0001$) and a significant interaction between species and origin ($p = 0.03$). Across all origins, *C. septempunctata* adults were significantly larger than *C. novemnotata* adults with mean sizes of 38.17 and 29.29 mm^2 , respectively. Within each individual origin *C. septempunctata* adults were significantly larger than *C. novemnotata* adults. Across both species the size of recently field-collected adults (P1) were significantly smaller than the size of adults from historical collections that were significantly smaller than laboratory-reared adults (F1) with means of 30.75, 33.82, and 36.62 mm^2 , respectively. Since the interaction between species and origin was significant we compared individual species-origin combinations and found that this effect was driven primarily by *C. novemnotata*. Combined historical (collection) specimens were significantly larger than the recently collected *C. novemnotata* adults (P1) (Fig. 1). The offspring of the field-collected *C. novemnotata* adults reared in the laboratory without prey limitation (F1) were also significantly larger than their parental generation (Fig. 1). *C. septempunctata* adults collected from the field were not significantly different from either adults from historical collections or adults reared in the laboratory without prey limitation (Fig. 1).

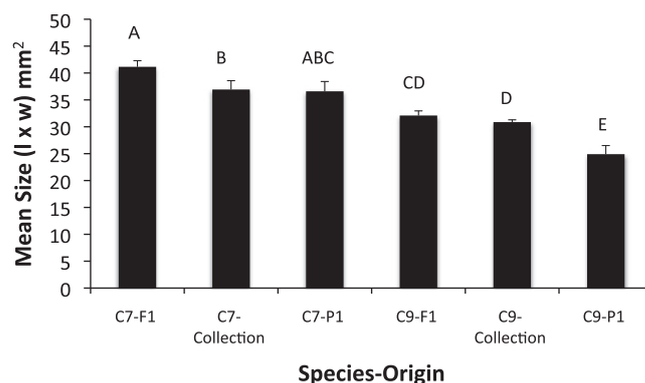


Fig. 1. Mean size (mm^2 with SEM as positive error bars) of *C. novemnotata* and *C. septempunctata* adults from recent field collections (P1), laboratory colonies (F1), and historical collections.

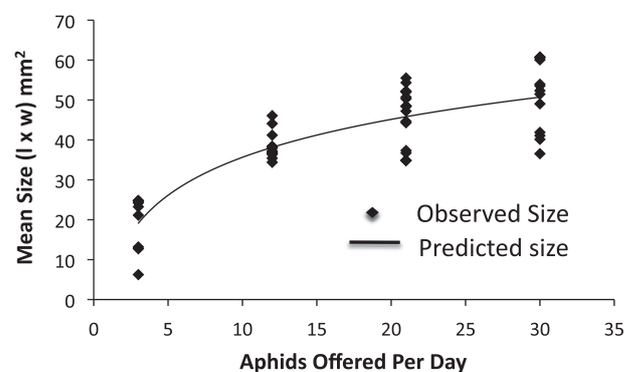


Fig. 2. Relationship between the number of aphids available per day to developing *C. novemnotata* larvae and resulting mean adult size (mm^2).

3.3. Feeding study

Prey availability during development was found to have a significant impact on adult *C. novemnotata* size and mortality. Prey availability had a significant positive relationship with adult beetle size ($p = 0.0001$) explaining over 70% of the variability ($R^2 = 0.72$; see Fig. 2). Adult size (measured as $l \times w$) ranged from 18.78 mm^2 at a prey level of 3 per day to 50.12 mm^2 at a prey level of 30 aphids per day. Prey availability also had a significant positive relationship with the proportion of larvae that survived to adulthood ($p < 0.00001$) explaining over 80% of the variability ($R^2 = 0.81$; see Fig. 3). The proportion of larvae that survived ranged from 0.00 when given one aphid per day to 0.75 at 21 aphids per day. Although the size of *C. novemnotata* individuals increased as prey availability increased from 21 to 30 aphids per day, the proportion of larvae that survived decreased to 0.60 at 30 aphids per day.

3.4. Extrapolating feeding study data to field populations

Based on a mean size of 24.90 for the recently field-collected adults this implies they may have had a diet of 4.57 aphids per day (95% confidence limits 3.39–5.70) in the field. Based on our logistic relationship this implies a survival rate of approximately 23%, which is less than a third of the mean maximum survival we found (75%) at 21 aphids per day.

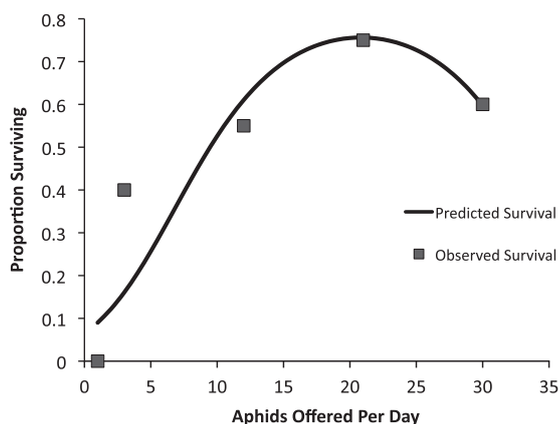


Fig. 3. Relationship between the number of aphids available per day to developing *C. novemnotata* larvae and resulting mean probability of survival to adulthood.

4. Discussion

When *C. novemnotata* adults were collected in the field, observations suggested that the mean size of these individuals was smaller than the mean historical size for this species. To test this hypothesis we first needed to determine if the adults we collected might be on the tail of an environmental cline. Pinned collections from around North America offered a wealth of data on historical size but it would have been inappropriate to compare the size of our adult beetles collected in the Northwest quadrant of the U.S. at relatively high altitudes to a simple mean size found in collections if the size of historical populations varied across either geographic location or altitude. Our results show that the size of adult *C. novemnotata* from historical populations did vary significantly across latitudinal, longitudinal, and elevational gradients. However, only a relatively small proportion of the variation was explained. It is interesting that size was positively correlated with elevation and latitude. This trend follows “Bergmann’s rule” which was developed for warm-blooded vertebrates (Bergmann, 1847) and states that body size increases with latitude and more generally lower temperatures. The trend we found is counter to the trend that other studies have found for other insects (Vamosi et al., 2007; Peck and Chapelle, 2003; Orr, 1996). Since the analysis from pinned specimen data predicts that adult *C. novemnotata* from high elevations in the Northwestern U.S. should be relatively larger than those from other areas and since very little of the size variation is actually explained, we concluded that the most appropriate comparison of sizes of our recently collected adults (which we suspected were smaller) was with a large, broad mean of historical *C. novemnotata* adult sizes.

The recently collected *C. novemnotata* adults were significantly smaller than adults from historical populations (see Fig. 1). This raises two possibilities. Either *C. novemnotata* sizes have evolved to be smaller or the potential for larger sizes remains but adult beetles are prevented from attaining larger adult sizes because of ecological conditions (resource limitation, infection, size-dependent predation). When reared in the laboratory the first generation descendants (F1) of the field-collected adults (P1) were physically larger than (although not quite significantly) historical populations. This implies that the potential for larger adult size is genetically maintained in *C. novemnotata* and that whatever factor that is preventing them from reaching this size in the field is not present in the laboratory.

Although several factors could contribute to smaller size in the field, we focus on one bottom-up factor: prey availability. We reason that although pathogens can directly lower adult size (Boots

and Begon, 1994), if infection was the cause of the smaller size of our beetles in the field it would have had the same impact in the captive population.

Following this line of reasoning we focus on prey availability as a plausible cause for the smaller size of recently collected *C. novemnotata* adults. Predatory insects show a range of responses to varying levels of prey availability during development. Lower prey levels during development can lead to decreases in survival to adulthood and smaller adult size (Pereyra and Archangelsky, 2007). This is the pattern that we found in our study (see Figs. 2 and 3). In fact, the mean size of field-collected adults fell within the range produced by our feeding trials, implying that prey limitation alone could produce the sizes we observed.

In our study, prey availability during development affected the probability of survival to adulthood as well as the size of the adults that emerged. The probability of survival through development increased with increasing prey availability in the range of 1–21 aphids/day but then declined going from 21 to 30. This implies that for this species it is possible to eat too many aphids and lose fitness. The mean size of *C. novemnotata* adults we collected in the field were nowhere near this “obesity” level. Their average size was almost 25 mm², which suggests they may have had access to approximately 5 aphids per day. This is a level of prey that exhibited only one third of the maximum survival we measured and could possibly have impacts on fecundity as well. The decreased fitness implied by the difference between current and historical adult *C. novemnotata* sizes appears severe enough to account for their observed decline in density and range.

If prey limitation is a contributing mechanism for decline, then the ultimate reason for the limited level of prey available to *C. novemnotata* and *C. septempunctata* deserves examination as a potential cause. This foreign species inhabits a very similar niche to *C. novemnotata* and usually in much higher densities. The Lost Ladybug Project lists 1758 records of *C. septempunctata* and only 45 for *C. novemnotata*. At the eastern Oregon collection site the ratio of *C. septempunctata* to *C. novemnotata* was greater than 20 to 1. In a recent submission from Idaho, students in Jerry Severe’s science class submitted over 730 identifiable lady beetle images to the Lost Ladybug Project. Over 75% of these were *C. septempunctata* while only 1 was a *C. novemnotata*. Beyond the disparity in density, *C. septempunctata* appears to be thriving at much closer to its potential than *C. novemnotata* based on the lack of difference between the size of field-collected, historical and laboratory-reared individuals (see Fig. 1). One insect predator may limit the prey available to another by interference with foraging (McPeck et al., 2001) or more commonly through scramble competition. There are documented cases of native predator species suffering lowered access to prey through scramble competition with the introduced species (Elliot et al., 1996; Evans, 1991).

If scramble competition is lowering prey densities available to *C. novemnotata* during development the immediate result could be smaller adult size. Day and Tatman (2006) suggest that north-eastern aphid populations are, in fact, steadily declining. Evans (2000) did not find evidence for a decline in body size for five native species as *C. septempunctata* densities increased in Utah in the 1990s. *C. novemnotata*, which was already very rare in Utah at that time, was not found in sufficient number to be included in the analysis (Evans, 2000). The evidence that (1) both species occur in the same habitats, (2) they are known to exploit the same prey, and (3) that lack of prey could cause the decline in mean *C. novemnotata* body size observed, inspired the hypothesis that *C. novemnotata* sizes have diminished due to lack of prey caused by scramble competition with *C. septempunctata* and that the smaller size of *C. novemnotata* is associated with lower fitness that led to their decline.

Some authors have suggested that *C. septempunctata* cannot be the sole reason for the decline of *C. novemnotata* because in some areas the decline seems to have preceded the invasion (Fothergill and Tindall, 2010). *C. septempunctata* may have moved in to occupy an incompletely filled niche after *C. novemnotata* had already begun its decline. In this case the main role of *C. septempunctata* may have been in preventing the reestablishment or rebounding of *C. novemnotata*. However, it also seems possible that upon arrival in new areas the earliest *C. septempunctata* invaders were misidentified or simply missed while still at low densities. Delayed detection has been documented for other invasive insects such as the cereal leaf beetle, *Oulema melanopus* (Haynes and Gage, 1981). While the introduced species is unlikely to have had a major impact at low initial densities its being overlooked could have caused the invasion clock to be initiated with an incorrect date. It also seems plausible a “wave of decline” could have hit a native species like *C. novemnotata* in a given area before the invading species arrived. If *C. novemnotata* dispersed equally in all directions but back dispersal was limited from one or more directions due to previous decline in another region, then decline could have preceded invasion. These types of sinks for populations, termed ecological traps, have been examined theoretically (Robertson and Hutto, 2006) and their potential to impact insect predator populations has been documented (Ries and Fagan, 2003).

Our study demonstrates that *C. novemnotata* adults are significantly smaller than they were historically and that prey availability alone is sufficient to account for this decrease in size. While our evidence is consistent with expectations if *C. septempunctata* were to negatively impact *C. novemnotata* through scramble competition, it is not exclusive of other potential mechanisms including competition for non-prey resources, intraguild predation, introduction of parasitoids or pathogens, and hybridization (Harmon et al., 2007). We suggest further study of current and historical distributions of *C. novemnotata* and *C. septempunctata* along with directed laboratory studies and modeling efforts to elucidate if *C. septempunctata* played a role in prey limitation leading to the decline of *C. novemnotata*.

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