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Agricultural practices for food safety threaten pest control services for fresh produce

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Running Head: Food safety practices threaten pest control

Summary

1. Over the past decade, several foodborne disease outbreaks provoked widespread reforms to the fresh produce industry. Subsequent concerns about wildlife vectors and contaminated manures created pressure on growers to discontinue use of manure-based composts and remove nearby semi-natural vegetation. Despite widespread adoption, impacts of these practices on ecosystem services such as pest control have not been assessed.
2. We used a landscape-scale field experiment to quantify associations between compost applications, semi-natural vegetation, pest control services, and lettuce yields on organic farms throughout California's Central Coast, a region experiencing food safety reforms.
3. We found that farms with surrounding semi-natural vegetation supported a diverse arthropod assemblage, whereas a herbivore-dominated assemblage occupied farms in simplified landscapes. Moreover, predatory arthropods consumed more herbivores at sites with more surrounding non-crop vegetation and reduced aphid pest infestations in lettuce.
4. Compost improved lettuce yields by increasing soil nutrients and organic matter, but affected neither pest control nor *E. coli* prevalence.
5. *Synthesis and applications.* Food safety concerns are prompting practices that simplify farms and landscapes. Our results demonstrate that two practices – elimination of manure-based composts and removal of non-crop vegetation – are likely having negative impacts on arthropod biodiversity, pest control, and soil quality. Critically, our findings and previous research suggest that compost can be applied safely and that habitat removal is likely ineffective at mitigating food

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safety risk. There is thus scope for co-managing fresh produce fields for food safety, ecosystem services, and biodiversity through applying appropriately treated composts and stopping habitat removal.

Key-words: agriculture, agro-ecology, biological control, co-management, compost, foodborne disease, food safety, produce, natural enemies, pest control

Introduction

Foodborne illness originating from fresh produce has emerged as a major public health concern (Painter et al. 2013), triggering sweeping reforms to the produce industry (LGMA 2013, FDA 2014). Reforms have been especially pervasive in California's Central Coast, the origin of a deadly, multi-state *E. coli* O157:H7 outbreak in 2006. Concerns that animals might carry foodborne illnesses created pressure on growers to reduce wildlife field intrusions by removing non-crop vegetation, perceived as wildlife habitat (Karp et al. 2015a). Following the outbreak, ~13% of the remaining riparian habitat along the Salinas River was degraded or destroyed (Gennet et al. 2013). Manures were targeted as another contamination source. Now, many organic growers apply liquid or heat-treated, pelleted fertilizers. The environmental consequences of these food-safety practices are largely unknown (Karp et al. 2015a), but may affect ecosystem services such as pest control (Letourneau et al. 2012) and building healthy soils (Mader et al. 2002).

In the Central Coast, a diverse suite of insect pests affects lettuce growers. Aphid and lepidopteran pests are usually the most damaging and often motivate insecticide applications (Bugg et al. 2008). Synthetic pesticides are unavailable to organic growers (~15% of growers in the region), who instead use a limited set of approved pesticides and/or rely on predators to consume aphids and other lettuce pests (Bugg et al. 2008, Smith et al. 2008, Nelson et al. 2012). Removal of predatory syrphid larvae from lettuce heads in one study caused aphid populations to explode, resulting in unmarketable

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plants (Smith et al. 2008). Food-safety induced vegetation removal may disrupt pest control because syrphids and other predators rely on non-crop vegetation to complete their lifecycles (Bugg et al. 2008). Generally, predator and parasitoid abundance, diversity, and consumption rates are higher on farms with more surrounding non-crop vegetation (Chaplin-Kramer et al. 2011). For example, semi-natural vegetation near broccoli farms increased syrphids as well as their predation on aphid pests, resulting in lower aphid population growth and less severe infestations at harvest (Chaplin-Kramer and Kremen 2013, Chaplin-Kramer et al. 2013).

Discontinuing manure-based compost applications could similarly disrupt predator-pest interactions. Mulches and composts may benefit predatory arthropods (Riechert and Bishop 1990, Settle et al. 1996, Mathews et al. 2004) if increases in organic matter enhance detritivore activity and thereby provide predators with a reliable food source (Settle et al. 1996). However, abundant detritivores could also satiate predators, and decrease pest consumption (Mathews et al. 2004).

We conducted a field experiment at 29 sites over two seasons in 2014 to evaluate how changing food-safety practices may influence pests, soils, and lettuce yields on organic farms in the Central Coast. We addressed four questions. First, how does surrounding semi-natural vegetation affect arthropods on produce farms? Second, does surrounding semi-natural vegetation provide a pest-control benefit to growers? Third, how does compost affect pests? Fourth, would retaining semi-natural vegetation and/or applying compost improve lettuce yields?

Materials and methods

Study Sites

We studied impacts of food-safety management in California's Central Coast. The region encompasses a mosaic of farms, estuaries, marshes, and riparian areas. Oak woodlands, upland scrub, and grasslands predominate in often-grazed, hillier areas. We utilized a natural gradient in this surrounding

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vegetation to study the effects of non-crop vegetation on pest control, crossed with an experimental manipulation of compost. We included sites in the San Juan (N= 8), Pajaro (N= 4), Northern Salinas (N= 12), and Southern Salinas (N= 5) Valleys (Fig. 1). We conducted our experiment over two seasons: spring (N=14 sites) and summer (N=15) of 2014. All farms were certified organic (<https://www.ams.usda.gov/rules-regulations/organic>).

Experimental Design

We established paired treatment plots (hereafter subplots)— receiving and not receiving compost— at farms along a gradient of surrounding semi-natural vegetation. Semi-natural vegetation occupied 2% to 56% (mean 29%) of the area within 1000m of plots (Table S1). To quantify pest control, we surveyed arthropod communities, measured pest consumption rates, and quantified lettuce yields. Specifically, we demarcated a 5m×10m plot at each site, encompassing either four 2m crop beds or eight 1m beds (depending on the grower’s planting design) near the field edge (Fig. S1). Planted lettuce plots were located within vegetable fields (lettuce: N=19, broccoli: N=7, celery: N=3). Management of plots aligned with the surrounding host farm. Beyond the normal fertilization regime, we applied organic compost on half (5m×5m) of each plot, 1-2 months prior to planting. Specifically, a cow, chicken, and green manure blend was applied at 25 tonnes/ha (Gabilan Ag Services, Salinas, CA). Prior to use, the compost was packed into windrows for 14-18 weeks, during which internal temperatures consistently reached levels necessary for inactivating pathogenic *E. coli* (55°C-75°C) (Jiang et al. 2003). Likely in part a result of recent food-safety concerns, compost is not regularly applied at most sites (N= 24), and no site applied compost in 2014. Instead, most growers used heat-treated, pelleted and/or liquid fertilizers (TRUE Organic Products, Inc., Spreckles, CA).

We planted sites from March 5-28 (spring) and May 30 to July 5 (summer). Each plot was planted with Romaine lettuce, either through direct seeding (N= 20; Rio Bravo variety; Syngenta Seeds Co. Minnetonka, MN) or transplanting (N= 9; Salvius variety; Johnny’s Selected Seeds Co., Fairfield,

ME) depending on grower preference. Plots that received transplants did not differ in any measured pest-control metric from direct-seeded plots (all $P > 0.05$). Seeds and seedlings were planted in 2 lines/1m bed or 4-5 lines/2m bed, and were subsequently thinned following industry standards.

Arthropod Sampling

We used a combination of survey methods to sample arthropod assemblages: pan traps (captured volant pests and enemies), pitfall traps (epigeal predators and detritivores), and collecting entire lettuce heads (pests and enemies that directly affect lettuce marketability). Sites were sampled twice: once at the seedling stage (~25 or ~7 days after direct seeding or transplanting, respectively) and once when the lettuce was mature (~60 or ~42 days).

For each sample round, we deployed 3 pitfall and 2 pan traps per subplot. Pitfall traps were constructed from 7.5cm diameter plastic cups, buried so that tops were flush with the soil, and filled with soapy water (~0.5tbsp Blue Dawn/L; Procter & Gamble Co., Cincinnati, OH). Traps were sheltered with Styrofoam bowls, suspended 5cm above the ground. Pan traps were 15cm plastic bowls, spray-painted fluorescent yellow or blue. Each subplot received 1 yellow and 1 blue pan filled with soapy water. Pitfall and pan traps were collected after 48hrs.

We also haphazardly collected three lettuce heads per subplot on the second site visit. Lettuce heads were cut at the base of the plant and transferred immediately into Ziploc bags. Bags were filled with water and drained through an arthropod funnel. Then, lettuce leaves were pulled off and rinsed above the funnel. RM identified 98% of all 38,418 arthropods captured to a family level. Known lettuce pests were identified to species, and RM, DSK, and SVJ assigned families to primary feeding guilds (see Appendix S1 in Supporting Information).

Predation Rates

We used a combination of sentinel pest (Meehan et al. 2012) and enclosure experiments (Chaplin-Kramer and Kremen 2013) to quantify predation rates. Our focus was on lepidopteran and aphid pests, important categories of lettuce pests (Bugg et al. 2008).

At seedling and mature lettuce phases, we placed live corn earworm eggs (*Helicoverpa zea*) and 2nd-3rd instar beet armyworm larvae (*Spodoptera exigua*) in our plots and monitored predation rates. Both pests were obtained from a commercial insectary (Benzon Research Inc., Carlisle, PA). We affixed 5 larvae and 25-70 (mean: 40) eggs to separate waterproof paper cards (~3.5cm x ~1.5cm; JL Darling LLC, Tacoma, WA). Larvae were adhered to cards at the posterior end of their bodies with hot glue to allow movement. Eggs were affixed by gluing small sections of their fabric substrates to cards. We used a microscope to count the initial number of eggs per card. During each visit, three each of egg and larvae cards were deployed in each subplot. After 48 hours, we collected all cards and counted the number of remaining eggs and larvae that had escaped predation.

We also conducted an enclosure experiment to quantify predation rates on potato aphid (*Macrosiphum euphorbiae*). Potato aphids were collected from lettuce plants at our study sites and reared in a greenhouse on Romaine lettuce. Once lettuce plants were mature, we first selected three lettuce plants at each sub-plot. Second, we removed all insects from each plant by visually inspecting leaves and removing encountered insects. Third, we washed the plant with water, and performed a second inspection. Fourth, we removed ~2cm of topsoil to exclude insects that fell off the plants and onto the soil during inspection. Fifth, we constructed bamboo frame, fine-meshed netted cages around each plant (0.4mmx6mm mesh, 7250NSW, Bioquip Products, Inc., Rancho Dominguez, CA). Each 40cmx40cmx40cm cage was buried ~10cm into the soil. Two of the three plants per sub-plot received control cages with no netting on two sides to allow natural enemies to forage and to ensure that all plants would experience similar micro-climates (Chaplin-Kramer and Kremen 2013). The third plant was completely enclosed.

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Finally, we inoculated the caged plant and one control plant with 50 aphids each. The other control plant did not receive aphids, and was used to monitor aphid colonization. Aphids were selected to reflect the colony's age structure, but alate (winged) aphids were excluded to prevent emigration (Chaplin-Kramer and Kremen 2013). After two weeks, plants were bagged and washed through a filter to collect all arthropods.

Yields and Soil Analyses

We calculated the average fresh weight (g) of harvested plants from a 1mx1m quadrat within each subplot (5-15 plants, mean= 7.5). In spring, we also collected soil samples to determine soil properties and pathogenic *E. coli* prevalence. We used a 1.25cm diameter soil probe to collect 1.4L of soil to a depth of 10cm per subplot. A commercial laboratory quantified concentrations of nitrate (mg/kg), phosphorous (mg/kg), and potassium (mg/kg), as well as pH, % organic matter, and % sand (Soiltest Farm Consultants Inc., Moses Lake, WA).

Samples were also tested for indicator and pathogenic (STEC) *E. coli*. From each sample, four 25g subsamples were inoculated into 225ml of buffered peptone water (BPW), shaken for 15min, and allowed to settle for 5min. One ml and 0.1ml aliquots of BPW from each subsample were spread-plated onto separate 150mm violet-red bile agar with VRB-MUG plates. The VRB-MUG plates and remaining BPW mixture were incubated overnight at 37°C. The red, fluorescent colonies on VRB-MUG were presumptively identified as *E. coli*, and three colonies per soil sample were then subjected to TSI, oxidase, and indole tests. All colonies yielding typical *E. coli* reactions to these tests were counted and banked. After counting, swabs were used to sweep the VRB-MUG plates and were stored in buffered glycerol at -20°C. All saved *E. coli* isolates were screened for shiga-toxins— the disease-causing agent present in some *E. coli* strains— using PCR. The PCR used multiplexed primers directed against markers stx1, stx2, eae, and fliCH7 (Olsvik et al. 1991, Gannon et al. 1997).

Habitat Classification

We contracted an Anderson Level II terrestrial land-use/land-cover map to calculate the extent of semi-natural vegetation in the Salinas, San Juan, and Pajaro Valleys (Total area: 1906km²; Aerial Information Services, Redlands, CA). Land cover was hand-classified into 16 categories from 1m² resolution National Agricultural Inventory Program imagery, taken in the summer of 2012. Due to the large spatial extent of our study area, only cropland and riparian features larger than 0.2ha were hand-classified (minimum mapping unit); the minimum mapping unit for other land cover classes was 0.4ha. We combined non-crop vegetation into one layer encompassing all semi-natural vegetation including: “Riparian Forest & Woodland,” “Southwestern North American Riparian Scrub,” “California Forest & Woodland”, “Upland Scrub,” “Mediterranean Grasslands & Forbs,” and “Meadow and Marsh.”

We developed an index to quantify the extent of surrounding semi-natural vegetation, recognizing that regions located closer to study sites were likely to have greater influence than those further away. First, we quantified the amount (m²) of semi-natural vegetation in 20 concentric rings, with inner radii at distances distributed uniformly on an exponential scale between 50m and 1500m. Next, we used a Gaussian function to assign weights to each ring, with farther rings given lower weightings than closer rings. Using a different decay function (*e.g.*, exponential decay) did not influence our results. Specifically, each ring was assigned weight *W*, given by:

$$W = \exp(-I^2 / (2 * d^2)) \quad \text{Equation 1}$$

where *I* is the inner edge distance of the ring and *d* is the decay rate that specifies how quickly weightings decrease with increasing distance. In order to determine the most predictive scale for subsequent analyses, we assessed several decay rates. We calculated the weighted sum of the area of semi-natural vegetation by summing across the 20 concentric rings the amount of semi-natural habitat within each ring, multiplied by that ring's weight. We standardized this index across study sites by subtracting the mean value and dividing by the standard deviation. Because the semi-natural vegetation index was highly negatively

correlated with a similar index of cropland extent (Fig. S2), we focus only on the effects of semi-natural vegetation on pest control.

We found the most predictive decay rate to be $\beta = 250$ (Fig. S3; Appendix S2). The amount of surrounding semi-natural vegetation less often predicted pest-control variables when it was calculated with larger decay rates ($\beta = 1000$ and 5000) that increased the importance of areas further from the study site (Appendix S2). We subsequently utilized $\beta = 250$ in our analyses.

Statistical Analyses of Arthropod Communities

We assessed the effects of semi-natural vegetation and compost on arthropod abundance, family richness, and biocontrol metrics with linear mixed-effects models (Zuur et al. 2009). Mixed effects models were used so that site and region could always be included as random effects. We first aggregated arthropod censuses to a sub-plot level, creating lists of all the arthropods captured in pitfall traps, pan traps, or lettuce heads during each visit to each sub plot. If traps were lost or broken (<2% of traps deployed), an alternate trap in the same sub-plot and visit was counted twice (1.3% of insects sampled).

We next evaluated whether to use generalized or linear mixed effect models. Simulations of null hypotheses (no effect of explanatory variables) suggested that evaluating significance using generalized linear mixed-effects models was anticonservative, with a type-I error rate of ~ 0.5 for a nominal P -value of 0.05 (Ives 2015). In contrast, linear mixed models yielded low type-I error rates (~ 0.05). Simulations were performed by randomly permuting values of semi-natural habitat and then comparing models with and without the predictor via likelihood ratio tests. The fraction of (1000) randomizations with P values < 0.05 was taken as the type I error rate.

Separate linear mixed-effects models were created for arthropods surveyed in lettuce heads, pitfall traps, or pan traps and for different response variables. Response variables of total arthropod, natural enemy, herbivore, aphid, non-aphid pest, syrphid fly, and detritivore captures were log-transformed to ensure normality and heteroscedasticity. Because residuals were normally distributed, it

was not necessary to transform family-level richness of all arthropods, natural enemies, and herbivores, or Pielou's evenness and Simpson diversity, except for pitfall traps (square-root transformation). To assess the relative abundance of enemies versus herbivores, we divided total enemy captures by the total captures of enemies and herbivores and square root transformed the result. The same approach was used for the relative abundance of enemies vs. aphids and enemies vs. non-aphid pests.

The effects of compost and semi-natural vegetation on each response variable were assessed in two steps. To investigate main effects, we created a first set of models with compost treatment, semi-natural vegetation (Gaussian-weighted index), season (spring or summer), and lettuce life stage (seedling or mature plant) as explanatory variables. Statistical significance of each predictor was evaluated via likelihood ratio tests that compared the full model to a reduced model without the variable (Zuur et al. 2009). Not all sites were included in each analysis: early harvests and mammalian herbivory caused some sites to be excluded (Table S2).

Next, we developed a second set of models to investigate whether effects of semi-natural habitat on response variables varied by season (summer vs. spring) and life stage (seedling vs. mature). To do so, we created four binary variables indicating whether the census took place in: (1) seedling stage/spring, (2) mature stage/spring, (3) seedling stage/summer, and (4) mature stage/summer. Each variable was allowed to interact with semi-natural habitat. With this parameterization, models could be compared via likelihood ratio tests with and without each interaction to determine whether semi-natural habitat significantly influenced response variables in each season-life stage combination. All other fixed effects (compost treatment, season main effect, and lettuce life stage effect) were included in these models. For analyses of detritivore abundance only, we additionally included interactions between compost treatment and season and life stage.

Analyses of Natural Enemy Activity

To assess landscape effects on enemy activity, we modeled associations between surrounding semi-natural habitat and predation rates on *H. zea* eggs, *S. exigua* larvae, and *M. euphorbiae* colonies. First, we aggregated data from egg and larvae cards to subplots, calculating the total number of eggs (or larvae) deployed and the fraction consumed. After calculating the proportion of eggs and larvae consumed at each plot, we modeled effects of surrounding semi-natural habitat as above. Egg and larvae proportion data were square-root transformed to ensure normality of residuals.

To quantify aphid predation from the enclosure experiment, we developed a biocontrol index (BCI) defined as:

$$\text{BCI} = \log\left(\frac{E+C}{O}\right) \quad \text{Equation 2}$$

where E is the final count of non-alate aphids in the exclusion treatment (closed to predation, received 50 initial aphids), C is the count in the colonization treatment (open to predation, no aphids), and O is the count in the control treatment (open to predation, 50 aphids). The colonization treatment was added to the exclusion treatment to account for aphids that colonized the open treatment during the experiment.

Ignoring the colonization treatment and re-calculating BCI as the ratio of aphids in predator enclosures versus controls did not change results. Higher values of BCI indicate more natural enemy suppression of aphids. Effects of semi-natural habitat on BCI were assessed using the same model structure as arthropod community analyses.

Yield Analyses

We assessed changes in lettuce yields relative to compost addition and landscape context, by calculating the average lettuce weight per sub-plot. To determine whether herbivores affect lettuce yields, we calculated an index of average herbivore abundance per sub-plot by: (1) quantifying total herbivores captures in lettuce heads, pan traps, and pitfall traps at each sub-plot, (2) scaling total captures across sub-plots for each capture method by subtracting the mean and dividing by the standard deviation, and (3)

calculating the average of the lettuce head, pan trap, and pitfall trap scaled values at each subplot. We then implemented linear mixed-effects models of average weight with season, compost, an interaction between semi-natural vegetation and season, an interaction between herbivore abundance and season, and the number of days between planting and collection as explanatory variables (three weeks were added to transplants to account for greenhouse growth). Lettuce weight was log-transformed to satisfy model assumptions.

Soil Analyses

We quantified effects of compost on soil properties and effects of soil properties on lettuce weights. We modeled effects of compost on nitrate, phosphorous, and potassium concentrations and on pH, percent organic matter, and percent sand. Next, we created models to predict changes in lettuce weights, including as fixed effects elapsed days past planting and soil properties (in separate models). Lettuce weights were log transformed and significance was assessed with likelihood ratio tests. All analyses were conducted in R version 3.2.2 (R Development Core Team 2010), using the “lme4” package (Bates *et al.* 2015).

Results

We captured a total of 38,418 arthropods across 21 orders and 128 families (Fig. S4; Appendix S1). Dominant guilds included herbivores (N= 24,670), detritivores (N= 8,183), and natural enemies (N= 3,015). 23% of herbivores were pests, mostly aphids (~90%). Plots surrounded by lettuce yielded more insect captures ($\chi^2 = 3.72$, $P = 0.05$), aphids ($\chi^2 = 6.36$, $P = 0.01$), and non-aphid pests ($\chi^2 = 4.80$, $P = 0.03$) than plots surrounded by other crops. Correspondingly, predation of *M. euphorbiae* and *S. exigua* larvae were lower in plots surrounded by lettuce ($\chi^2 = 6.03$, 3.84 ; $P = 0.01$, 0.05 respectively). Semi-natural vegetation did not differ between plots surrounded by lettuce vs. other crops ($F = 1.24$, $P = 0.28$).

Arthropod assemblages changed predictably across the vegetation gradient. First, fewer arthropods were captured on lettuce heads and in pitfall traps at sites with more surrounding semi-natural vegetation (Fig. 2; Table 1). Pan trap captures declined marginally ($P=0.09$). Second, while total captures declined, the number of arthropod families captured either remained constant (lettuce heads and pitfall traps) or increased (pan traps). As a result, sites with more surrounding semi-natural vegetation hosted more diverse assemblages (Table 1 and S2).

Higher arthropod diversity in landscapes with more semi-natural vegetation could be characterized by an increase in natural enemies relative to herbivores. Enemy richness in pan traps was higher at sites with more semi-natural vegetation (Table S3). Moreover, herbivore captures were lower in landscapes with more semi-natural vegetation under each survey method, while enemies were unaffected (Table S3). Therefore, the relative abundance of enemies versus herbivores in lettuce heads and pan traps was significantly higher at sites with more surrounding semi-natural vegetation, especially when lettuce plants were at the seedling stage (Fig. 2; Table 1). The one exception was pitfall traps, where both enemies and herbivores were less abundant at more vegetated sites.

Some pests (*i.e.*, aphids) were also less abundant at sites with more nearby semi-natural vegetation (Fig. 3; Table 2). Correspondingly, the relative abundance of natural enemies versus aphids was higher in areas with more surrounding semi-natural vegetation. Non-aphid pests, however, exhibited no clear response to surrounding landscape composition (Fig. 3; Table 2).

Exclusion experiments suggested that enemies consumed a significant fraction of pests. Excluding enemies for two weeks caused a 4.7 fold increase in aphid captures, from ~40 to ~170 aphids/plant when enemies were present versus excluded (Fig. S5; $\chi^2=54$, $P<0.001$). This estimate is likely conservative, as cages did not completely exclude predators. Still, at the conclusion of the experiment, significantly more enemies were found in the open versus caged treatment (~8 versus ~4 enemies/plant) ($\chi^2=33$, $P<0.001$).

Predation rates also varied along the landscape gradient (Fig. 4; Table 3). For example, we found that the biocontrol index (see methods) of aphid consumption was significantly higher at sites with more surrounding semi-natural vegetation. *H. zea* egg consumption was also higher in landscapes with more semi-natural vegetation, but only at the lettuce seedling stage. In contrast, *S. exigua* larval predation never changed across the land-use gradient. These non-significant results may have been influenced by the difficulty of separating consumed larvae from those dehydrated from sun exposure. Additionally, larvae may have occasionally escaped.

Unlike semi-natural vegetation, we found no consistent effects of compost on any measure of arthropod abundance or diversity (Table 1-3, S3), including detritivore captures (Table S4). Compost did, however, increase lettuce yields without compromising food safety (Fig. 5; Table S5). No samples tested positive for shiga-toxin producing *E. coli* (STEC), and only two samples tested positive for generic *E. coli* (one of which was in a control subplot; thus, compost was not an *E. coli* source). Compost increased soil phosphorous, potassium, and organic matter, all of which were positively correlated with lettuce weights (Fig. 5; Table S6). Compost, however, did not affect nitrates, which were also positively correlated with lettuce weights. Similarly, compost did not alter soil pH or the fraction of sand in the soil. While soil pH yielded no effects, lettuce weights were highest in soils with intermediate levels of sand (~40% sand; Fig. 5).

Herbivores and surrounding vegetation were negatively correlated with lettuce weights (Table S5). Weights declined with herbivore abundance in summer but not spring. Instead, sites with more surrounding semi-natural habitat had lower yields in spring.

Discussion

If fresh produce continues to be a source of disease outbreaks, then pressure on growers to enforce sterile environments and remove semi-natural vegetation may escalate (Karp et al. 2015a). Recent work from the Central Coast of California, however, has demonstrated that foodborne pathogens do not increase on farms with more surrounding semi-natural vegetation (Karp et al. 2015b). Similarly, food-

safety risk from composts can largely be eliminated by subjecting pathogens to the high temperatures experienced inside windrows (Jiang et al. 2003). Correspondingly, in this experiment, we did not observe elevated pathogen prevalence on composted plots or on plots surrounded by semi-natural vegetation. Our results do suggest, however, that removal of vegetation and cessation of compost could negatively affect growers through impacts on pests, soils, and yields.

Effects of semi-natural vegetation

We found that produce farms with more surrounding semi-natural vegetation had more diverse arthropod assemblages. In contrast, farms in more simplified landscapes hosted high abundances of fewer arthropod families, largely dominated by herbivores. Lettuce heads were also more infested with aphid pests on farms with less surrounding semi-natural vegetation. We likely observed these trends because we worked on organic farms— inorganic insecticides would likely lower arthropod abundances and mask trends.

Unlike herbivores, effects of semi-natural vegetation on enemies were unclear which was surprising given that enemies generally increase on farms with more surrounding semi-natural vegetation (Chaplin-Kramer et al. 2011). Indeed, other studies from the Central Coast have documented benefits of semi-natural vegetation for many of the enemies surveyed here (Letourneau et al. 2012, Chaplin-Kramer et al. 2013). One possibility is that enemies responded to fine-scale habitat elements (*e.g.*, hedgerows) that were too small to be classified in our land-use map. Another explanation is that enemies colonize farms from semi-natural habitats early on (Costamagna *et al.* 2015), but, over the season, enemies eventually equilibrate in population size across sites. Indeed, we observed that enemy captures and the relative abundance of enemies versus herbivores were more positively associated with semi-natural vegetation at the seedling than at the mature lettuce stage. Herbivore suppression at the seedling stage may be critical for some crops; tomatoes, for example, are vulnerable to insect pests at the seedling stage but can later withstand pests (Drinkwater et al. 1995).

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Early enemy colonization from surrounding habitats may be responsible for suppressing later pest population growth and lowering aphid abundances in landscapes with more semi-natural vegetation (Costamagna *et al.* 2015). Alternatively, because cropland and semi-natural habitat areas are negatively correlated, intensive farm management practices in more simplified landscapes may have reduced enemy activity and favored pests (*e.g.*, excessive pesticide applications, removal of floral resources etc.). Regardless, our finding that two weeks of enemy exclusion caused a nearly five-fold increase in aphid population size supports the idea that enemy activity could have driven spatial trends in pests. Indeed, at sites with more surrounding semi-natural habitat, we detected stronger effects of enemy exclusion on aphids. We also observed more egg predation at the seedling stage at sites with more surrounding semi-natural vegetation.

We found no evidence, however, that enhanced pest control increased lettuce weights. Yield effects are difficult to document in landscape analyses of pest control (Bianchi *et al.* 2006, Chaplin-Kramer *et al.* 2011). Because agriculture may expand more rapidly in productive regions, areas where natural habitat remains may be more marginal for agricultural production. Indeed, in our study, while we observed lower lettuce weights at more vegetated sites in the spring, this may have been due to lower soil quality on farms near remaining habitat. The two sites with lowest lettuce weights were surrounded by semi-natural vegetation, but also had the lowest concentrations of soil nitrate, organic matter, and phosphorous. Excluding either site caused semi-natural habitat to no longer significantly affect lettuce weights ($P > 0.05$).

While yields are a key incentive for pest management, other biocontrol metrics may be equally important. Even without yield impacts, increased predation on aphids may improve profits, as small numbers of aphids can render lettuce plants cosmetically unmarketable (Smith and Chaney 2007, Nelson *et al.* 2012). Because aphid infestations declined in landscapes with more semi-natural vegetation, surrounding habitat could thus enhance lettuce marketability. Moreover, on farms with more surrounding vegetation and lower aphid infestations, growers could conceivably apply less insecticide, reducing costs

and impacts on public health (Marks et al. 2010, Bouchard et al. 2011) and the environment (Anderson et al. 2003).

Effects of compost

Paralleling other work from the region (Jackson et al. 2004), applying compost did not initiate a beneficial trophic cascade that increased pest control. On the one hand, previous studies that have shown compost-driven increases in detritivore and/or predator abundances have generally applied significantly more mulch (Riechert and Bishop 1990), compost (Bell et al. 2008), or both (Mathews et al. 2004) than is typical for fresh produce growers in our region. On the other hand, some studies have found effects of compost on pest control at similar application rates to those used here (Settle et al. 1996). Consistent and multi-year compost applications may be necessary to sufficiently alter soil conditions before an effect on arthropods can be observed. Moreover, our plot size (5m×5m) may not have been large enough to attract detritivores and initiate a trophic cascade.

Despite little evidence that compost affects pest control, we did find that lettuce weights were higher on plots treated with compost and that *E. coli* prevalence did not change. The positive effect of compost on lettuce yields may have resulted from increases in organic matter and soil nutrients (*e.g.*, phosphorous and potassium) in compost-treated plots. Indeed, lettuce weights were significantly higher in plots with more phosphorous, potassium, and organic matter in their soils. Especially because no substitute fertilizer was used on non-composted plots, it is unclear whether increases in lettuce weights would also be observed on conventional farms that apply synthetic fertilizers.

Conclusion

Food-safety concerns have spurred pervasive reforms to produce management practices throughout the U.S. (LGMA 2013, FDA 2014). Our results demonstrate how two food safety practices—replacement of manure-based compost and removal of non-crop vegetation — may affect growers through impacting pests, soils, and yields. Specifically, we observed a negative association between crop

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pests and surrounding semi-natural habitat, and higher lettuce yields and healthier soils on plots with compost. All plots had low levels of *E. coli*, regardless of landscape context and compost treatment. Indeed, emerging evidence suggests that non-crop vegetation can be maintained (Karp et al. 2015b) and compost applied (Jiang et al. 2003) without compromising produce safety. Combined, our work suggests that growers may benefit from increased ecosystem services, without incurring food-safety costs, by applying appropriately treated compost and maintaining semi-natural habitat. Co-managing fresh produce for food safety and conservation goals is possible and likely beneficial for nature and for growers.

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Data accessibility

Arthropod surveys, enclosure experiments, sentinel pest experiments, lettuce weights, soil analysis, and surrounding semi-natural habitat indices are archived in Dryad Digital Repository <http://dx.doi.org/10.5061/dryad.37cc5>.

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Figure Legends:

Figure 1: Study site map. Our experiment encompassed 14 sites in spring (purple dots) and 15 in summer (red dots).

Figure 2: Effects of surrounding semi-natural vegetation on arthropod assemblages. Captures in lettuce heads (a) and pitfall traps (c) declined at sites with more surrounding semi-natural vegetation. Captures marginally declined in pan traps (b). Family richness in pans (e) but not lettuce (d) or pitfalls (f) increased at sites with more semi-natural vegetation. Family-level Simpson diversity also increased in pans (h) and pitfalls (i) but not lettuce heads (g). Sites with semi-natural vegetation had more enemies relative to herbivores in lettuce heads (j) and pans (k) but not pitfalls (l). In j, k, and l, enemy relative abundance is calculated as enemy captures divided by enemy plus herbivore captures. Model predictions are presented for significant effects— solid lines are slopes and dotted lines are 95% confidence intervals. Points are raw data at sub-plots. Light gray shading corresponds to samples obtained at the lettuce seedling stage; darker shading indicates mature lettuce. Panels a-c are on a log scale.

Figure 3: Sites with more surrounding semi-natural vegetation were less infested with aphids (a) but not non-aphid pests (b). Similarly, enemy abundance relative to aphids (c), but not non-aphid pests (d), increased with semi-natural vegetation. In c and d, enemy relative abundance is calculated as enemy

captures divided by enemy plus aphid (or non-aphid pest) captures. Points are subplots. For variables significantly affected by semi-natural vegetation, solid lines are model predictions and dotted lines are 95% confidence intervals. Panels a and b are on a log scale.

Figure 4: Effects of semi-natural vegetation on enemy activity. Aphid growth was more suppressed (Biocontrol Index- BCI, see methods) at sites with more surrounding semi-natural vegetation (a). Egg predation also increased with semi-natural vegetation, but only at the lettuce seedling stage (b). No landscape effects on larvae consumption were observed (c). Points are subplots. For significant models, solid lines are predicted slopes and dotted lines are 95% confidence intervals. Lighter shading indicates that samples were taken at the lettuce seedling stage; darker shading corresponds to mature lettuce.

Figure 5: Effects of compost and soil properties on yields. Adding compost caused an increase in lettuce weight (a). Black point in panel a is the mean effect of compost addition, lines are 95% confidence intervals, and gray points are sub-plots. Lettuce weight increased in subplots with more soil nitrate (b), phosphorous (c), potassium (d), and organic matter (f). Lettuce weight peaked at intermediate levels of sand (g), and was unaffected by pH (e). In regression plots of panels b-g, points are subplots, solid lines are model predictions, and dotted lines are 95% confidence intervals. Compost increased phosphorous (c), potassium (d), and organic matter (f), but not nitrate (b), pH (e), or sand (g). For compost effects in panels b-h, points are mean values per treatment and lines are confidence intervals. Filled circles correspond to compost plots; open circles are controls.

Table 1: Effect of semi-natural vegetation on arthropods. Significance was assessed with likelihood ratio tests (LRT), comparing models with and without predictors (see Methods). Because lettuce heads were not sampled at the seedling stage, no life stage interactions are included. Transformations to ensure normality and heteroscedasticity are noted in parentheses. Significant effects are bolded.

Response	Predictor	Lettuce Heads			Pan Traps			Pitfall Traps		
		β	χ^2	<i>P</i>	β	χ^2	<i>P</i>	β	χ^2	<i>P</i>
Total Arthropod Counts (log)	Intercept	4.78			4.65			4.14		
	Habitat	-0.35	4.66	0.03	-0.16	2.97	0.09	-0.32	14.36	<0.01
	Compost	-0.05	0.24	0.62	0.05	0.19	0.66	-0.06	0.38	0.54
	Summer	-0.32	1.06	0.30	0.11	0.39	0.53	0.08	0.27	0.61
	Mature Plant				0.31	6.38	0.01	0.44	17.97	<0.01
	Habitat: spring seedling				-0.27	2.47	0.12	-0.42	8.68	<0.01
	Habitat: spring mature	-0.40	2.72	0.10	0.13	0.63	0.43	-0.40	7.79	0.01
	Habitat: summer seedling				-0.23	2.63	0.10	-0.26	5.07	0.02
Habitat: summer mature	-0.31	2.25	0.13	-0.23	2.51	0.11	-0.25	4.66	0.03	
Arthropod Family Richness (log)	Intercept	13.92			11.60			8.44		
	Habitat	-0.09	0.02	0.90	1.71	7.73	0.01	0.31	0.58	0.45
	Compost	0.56	0.82	0.37	0.38	0.30	0.58	-0.43	0.65	0.42
	Summer	-0.11	0.01	0.93	0.68	0.36	0.55	2.06	5.81	0.02
	Mature Plant				7.43	72.02	<0.01	4.60	53.39	<0.01
	Habitat: spring seedling				1.90	3.19	0.07	1.00	1.76	0.19
	Habitat: spring mature	-1.09	1.02	0.31	1.21	1.32	0.25	0.16	0.04	0.83
	Habitat: summer seedling				1.46	2.82	0.09	-0.26	0.17	0.68
Habitat: summer mature	0.59	0.46	0.50	2.21	6.03	0.01	0.03	0.35	0.55	
Simpson Diversity (none but pitfalls which were squared)	Intercept	0.63			0.52			0.37		
	Habitat	0.04	2.17	0.14	0.04	5.11	0.02	0.09	8.36	<0.01
	Compost	0.00	0.00	0.94	0.01	0.26	0.61	0.00	0.00	0.97
	Summer	0.04	0.63	0.43	0.08	5.36	0.02	0.05	1.01	0.32
	Mature Plant				0.14	19.38	<0.01	0.09	6.54	0.01
	Habitat: spring seedling				0.10	6.62	0.01	0.12	6.05	0.01
	Habitat: spring mature	-0.01	0.02	0.90	-0.02	0.25	0.61	0.10	4.39	0.04
	Habitat: summer seedling				0.04	1.88	0.17	0.05	1.82	0.18
Habitat: summer mature	0.08	3.23	0.07	0.05	2.87	0.09	0.09	4.61	0.03	
Relative abundance of Enemies vs. Herbivores (square root)	Intercept	0.33			0.11			0.35		
	Habitat	0.05	6.09	0.01	0.04	11.20	<0.01	0.03	2.28	0.13
	Compost	0.02	0.50	0.48	0.02	0.95	0.33	0.00	0.02	0.90
	Summer	0.07	4.15	0.04	0.10	16.23	<0.01	0.25	29.32	<0.01
	Mature Plant				0.17	50.49	<0.01	0.17	33.28	<0.01
	Habitat: spring seedling				0.06	5.85	0.02	0.07	3.82	0.05
	Habitat: spring mature	0.03	1.12	0.29	0.00	0.00	1.00	0.02	0.30	0.58
	Habitat: summer seedling				0.04	4.80	0.03	0.01	0.11	0.74
Habitat: summer mature	0.06	5.83	0.02	0.06	7.56	0.01	0.02	0.66	0.42	

Table 2: Effect of semi-natural vegetation on pests on lettuce. Data transformations are indicated in parentheses. Significant effects are bolded.

Predictor	Aphid Count (log)			Non-Aphid Pest Count (log)			Relative Abundance of Enemies vs. Aphids (square root)			Relative Abundance of Enemies vs. Non-Aphid Pests (square root)		
	β	χ^2	<i>P</i>	β	χ^2	<i>P</i>	β	χ^2	<i>P</i>	β	χ^2	<i>P</i>
Intercept	3.63			-0.46	1.34	0.25	0.49			0.91		
Habitat (overall)	-0.52	3.90	0.05	-0.56	2.73	0.10	0.08	4.54	0.03	0.01	0.60	0.44
Compost	-0.01	0.00	0.97	-0.46	1.34	0.25	0.01	0.07	0.79	0.01	0.16	0.69
Summer	-0.16	0.10	0.76	-0.56	2.73	0.10	0.03	0.18	0.67	0.00	0.04	0.84
Habitat: spring	-0.46	1.34	0.25	-0.46	1.34	0.25	0.05	0.79	0.37	0.04	3.81	0.05
Habitat: summer	-0.56	2.73	0.10	-0.56	2.73	0.10	0.10	4.28	0.04	-0.01	0.22	0.64

Table 3: Effect of semi-natural vegetation on pest consumption. Data transformations are indicated in parentheses. Significant effects are bolded.

Predictor	Aphid Growth (log)			% Eggs Consumed (none)			% Larvae Consumed (none)		
	β	χ^2	<i>P</i>	β	χ^2	<i>P</i>	β	χ^2	<i>P</i>
Intercept	2.03			0.21			0.38		
Habitat (overall)	0.38	4.37	0.04	0.04	3.02	0.08	0.00	0.00	0.94
Predators in Cage	-0.54	4.46	0.03						
Compost	-0.29	0.68	0.41	0.01	0.06	0.81	-0.02	0.43	0.51
Summer	0.00	0.00	0.98	0.15	10.50	<0.01	0.03	0.23	0.63
Mature Plant				-0.09	8.06	<0.01	-0.02	0.32	0.57
Habitat: spring seedling				0.14	11.69	<0.01	-0.02	0.12	0.73
Habitat: spring mature	0.25	0.89	0.35	-0.02	0.23	0.63	0.05	0.59	0.44
Habitat: summer seedling				0.06	2.80	0.09	-0.03	0.31	0.58
Habitat: summer mature	0.47	3.98	0.05	-0.01	0.16	0.69	0.00	0.00	0.95









