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The potential of cleptoparasitic bees as indicator taxa for assessing bee communities

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Abstract – Many factors affect bee diversity and abundance, and knowledge of these is crucial for maintaining healthy bee communities. However, there are few means to fully evaluate the status of bee communities; most are based on monitoring species richness and abundance and do not consider the diverse life histories of bees. We propose that functional diversity of bee communities offers a more consistent means of evaluation and suggest that cleptoparasitic bees in particular show much promise as indicator taxa. Cleptoparasitic bees play a stabilising role within bee communities. They represent the apex of bee communities and are the first guild to respond to disturbances, are easily distinguished as such and are diverse enough to be representative of entire bee communities. The diversity and abundance of cleptoparasites in relation to all bees is indicative of the status of the total bee community, and monitoring them should form an integral part of assessing bee communities.

pollinator communities / guild structure / cleptoparasites / indicator taxa / ecosystem health

1. INTRODUCTION

There is no denying that human-assisted environmental changes are taking place on a global scale and that these changes are having strong impacts on vital ecological processes through their effects on biodiversity (Balvanera et al. 2006). One such ecological process evidencing these impacts is pollination (Biesmeijer et al. 2006; National Research Council 2007), the success of which for most plants is linked to pollinators, including bees (Michener 2007). Bees are keystone components of most terrestrial ecosystems, pollinating plants in natural and managed settings (Biesmeijer et al. 2006; Klein et al. 2007). In

the past decade, declines in bees and other pollinators have prompted much justified concern over the potential impacts to food production and ecosystem stability (e.g. Kremen et al. 2002, 2004; Biesmeijer et al. 2006; National Research Council 2007). As such, the need to understand which factors influence bee species richness (henceforth called “diversity”) are crucial to promote plant reproduction, bee conservation and stewardship, and prevent the continued decline of pollinators (Williams et al. 2010).

In general, the diversity of a community influences its stability, productivity and susceptibility to invasion (Hooper et al. 2005; Cardinale et al. 2006; Duffy et al. 2007). However, for most regions of the world, very little historic or “baseline” bee community data exist for comparison to present data. In response, much recent work has been done to develop and evaluate monitoring schemes,

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which accurately reflect bee diversity (e.g. Toler et al. 2005; Roulston et al. 2007; Westphal et al. 2008; Grundel et al. 2011; Nielsen et al. 2011), many concluding that sampling methods differ with respect to capture efficiency and representation of fauna and that they often complement each other. Clearly, habitats differ dramatically with respect to the bee fauna as a whole, as some habitats and/or areas are naturally more species rich than others (Michener 1979, 2007). A general consensus is that anthropogenic habitats (e.g. agro-ecosystems) typically have lower bee diversity than surrounding natural habitats (Kremen et al. 2002, 2004; Cane et al. 2006; Klein et al. 2007). However, interpreting diversity data can often be difficult; although comparisons of specific metrics among sites (and studies) are possible, the utility of such data for diagnosing the status of bee communities within the habitat(s), thus serving as an ecological indicator, is not always straight forward. For instance, in our recent study (Sheffield et al. 2013), we used traditional approaches for analysing bee diversity in response to ecological disturbances, finding that diversity differed significantly between intensely managed apple orchards and unmanaged habitats, though habitats of intermediate management intensity (i.e. differing proportions of unmanaged/natural habitat) were most similar to either extreme (i.e. the highly disturbed habitats, or to unmanaged habitats), depending on the metrics used to evaluate diversity data. As such, conclusions drawn on the impact of habitat enrichment for pollinators would be considered effective, or not, based on the estimators of diversity used. Moreover, these metrics do not consider the diverse life histories of the bees; in most studies, bees are usually considered on the whole a group with similar ecological behaviours.

A potential solution is to incorporate life history traits into such analyses (Tilman and Lehman 2001; Moretti et al. 2009; Williams et al. 2010; Scrosati et al. 2011) as not all species are functionally, evolutionarily and ecologically

equivalent (Chiarucci et al. 2011). Bees display a range of lifestyles, which can be assigned to non-taxonomic functional groups or “guilds” (*sensu* Blondel 2003) based on nesting biology, lifestyle, floral host specialisation, body size, etc. (Oertli et al. 2005; Moretti et al. 2009; Neame et al. 2012; Sheffield et al. 2013). This wide range of traits allows bees to exploit many habitats (Michener 2007; Williams et al. 2010), and monitoring the relative proportional abundance and diversity of members of these guilds provides additional information for describing and comparing communities (Sheffield et al. 2013). Bee guilds may show unique preferences and/or distinct requirements for habitat components (e.g. floral specialisation, nesting substrate preferences), and/or habitat type; different habitats will have specific guild profiles, and the guilds present in these habitats will show different responses to disturbance (Moretti et al. 2009; Neame et al. 2012; Sheffield et al. 2013). As such, guild profiles can be developed as baselines and, when considered together with other methods of assessing bee communities (e.g. standard diversity statistics, species abundance distributions, resulting fruit and seed set, seed yield, etc.), offer more information on the factors affecting bee communities within these habitats and their responses to disturbance, and better conclusions may be drawn with regards to the health of the ecosystem and/or how to improve it (Sheffield et al. 2013).

The main objective of this study is to demonstrate that assessment of bee communities should include the diverse life histories of bees in addition to traditional approaches of measuring species richness, diversity indices and/or abundance. We emphasise cleptoparasitic bees as indicator taxa for bee communities; a cleptoparasitic (or cuckoo) bee is one in which the adult female enters the nest of a non-conspecific host bee, oviposits in a natal cell and then departs from the nest; the cleptoparasite larva matures on the provisions stored for the host's larva (Rozen 2001). Benefits of a cleptoparasite-focussed approach are discussed, which is compatible with traditional approaches of bee community diversity.

2. MATERIALS AND METHODS

2.1. Study sites

The analysis presented here is of a dataset previously published by Sheffield et al. (2013). In 2001 and 2002, 19 sites within the Annapolis Valley, Nova Scotia were surveyed for bees, each selected to represent habitats within an ecological gradient. Habitats ranged from highly managed agro-ecosystems through levels of partial (i.e. adjacent) unmanaged and natural habitat to old fields. The four habitat categories were the following: (1) COMM-A (five sites), commercially managed apple orchards surrounded by adjacent orchard blocks or other agricultural crops, etc.; (2) COMM-B (five sites), commercially managed orchard blocks isolated from other agricultural areas and/or usually surrounded by adjacent woodland or non-agricultural land; (3) ABAND (four sites), abandoned and/or un-managed orchard sites that have not been sprayed or otherwise managed for production for at least 10 years; and (4) OLD FIELD (five sites), unmanaged meadow or open habitats with adjacent woodlands. Further details on the study sites can be found in Sheffield et al. (2008, 2013).

2.2. Site characterisation

The surrounding habitat landscape at each site was further categorised using satellite maps, overlaid with a 24×24 grid, each grid cell representing 25×25m, with the central sampling point at the centre of the grid (see Sheffield et al. 2013). A normalised measure of habitat dominance (h_D) (after O'Neill et al. 1988) was assessed; values of h_D range between 0 and 1, with higher values indicating a landscape dominated by only a few cover types; values closer to 0 indicating proportions of land cover that are nearly equal. The following landscape classifications were used: (A) tree fruit orchard under intense management, (B) non-tree fruit agricultural cropland under intense management, (C) woodland, (D) pasture (no spraying, seasonal mowing), (E) residence, (F) meadow and (G) abandoned orchard. Large bodies of water and paved areas occupying most of the surface of individual grid cells were subtracted from the total grid number. h_D was calculated at three

levels for each site: (A) the inner 8×8 grid; (B) a 16×16 grid, and (C) the total 24×24 grid.

2.3. Bee survey methods

At each site, nine 12-oz yellow pan traps (Solo® Cup Company; catalogue number PSB2Y 0099) were placed in a 3×3 grid, each pan trap separated by at least 10 m. Trap contents were collected weekly and pan traps replaced. Although it is known that the use of pan traps of several colours (e.g. yellow, blue and white) (Toler et al. 2005) and/or use of several sampling techniques (Roulston et al. 2007; Westphal et al. 2008; Grundel et al. 2011; Nielsen et al. 2011) typically offers a fuller representation of regional bee faunas in most circumstances, the sampling method used here is consistent across all sites, and captures were representative of the bee fauna of Nova Scotia (Sheffield et al. 2013). Voucher specimens of bees collected in this study are held at the Packer Collection, York University (Toronto, Ontario, Canada) and the Royal Saskatchewan Museum (Regina, Saskatchewan).

2.4. Data analysis

For each site, data from all nine pan traps were pooled prior to analysis. Total guild structure of the bee communities in each habitat types was previously compared (Sheffield et al. 2013); each species was assigned to one of eight guilds based on their known biology and included (A) solitary ground-nesters, (B) social ground-nesters, (C) honey bees, (D) non-parasitic bumble bees, (E) cavity-nesters, (F) cleptoparasites and (G) social parasites (*Bombus* subgenus *Psithyrus*). Sheffield et al. (2013) reported significant differences in the proportion of bees representing each guild for several guilds among habitats, including cavity-nesting bees, *Bombus*, and cleptoparasites; the present analysis focusses on the cleptoparasitic bee guild.

To demonstrate the diversity and abundance patterns of cleptoparasites within the four habitat types, data for this guild in each abundance octave were added to the truncated log-normal plots; species abundance data for each habitat were log transformed ($x=\log_2 n_i$) and fitted to a truncated log-normal

distribution following Magurran (2004). Although measuring deviations in the fit to the truncated log-normal distribution has been suggested as a diagnostic for the assessment of ecosystem health (Hubbell 2001), including for bee communities (Kevan et al. 1997), our purpose here is to show a summary of cleptoparasite species richness and abundance versus other bees. Sheffield et al. (2013) offer further discussion of this metric for assessing bee communities.

For individual sites, the number of cleptoparasitic species and the Berger–Parker dominance index were regressed against the number of cleptoparasitic individuals. To determine the impact of landscape on bee diversity, the reciprocal of the Simpson's diversity index ($1/D$) and the proportion of cleptoparasitic individuals were regressed against h_D .

3. RESULTS

3.1. Bee diversity

In both years, a total of 7,234 bee specimens, representing 146 species, were identified from yellow pans (see Sheffield et al. 2013, for complete list of species). Cleptoparasites were present in all habitat types, being proportionally most species rich and abundant in ABAND and OLD FIELD, but almost non-existent in COMM-A (Figure 1).

3.2. Landscape effects

Landscape structure at the smallest scale was the best predictor of both bee diversity and the percentage of cleptoparasitic bees; both variables increased in a similar fashion as h_D decreased (Figure 2A, D). As a larger proportion of the surrounding landscape was included, much smaller effects were observed (Figure 2B, E) on both bee diversity and cleptoparasites. At the largest scale, the reverse effect was observed, with bee diversity and the percentage of cleptoparasites decreasing as h_D increased (Figure 2C, F), though at this scale, unmanaged habitat types were the dominant land cover (Sheffield et al. 2013). In all cases, overall bee diversity and the cleptoparasites percentages followed the same pattern (Figure 2).

A strong relationship between the proportion of cleptoparasitic species and the proportion of cleptoparasitic individuals was observed across sites (Figure 3A); species richness and abundance increasing in ABAND and OLD FIELD habitats. This general trend was also supported in the truncated log-normal plots (Figure 1), indicating that abundance and diversity of cleptoparasites was much less in highly managed habitats (i.e. COMM-A) and increased in habitats with increasing levels of natural habitat. Dominant species were much more prominent in highly managed habitats, but decreased significantly as the proportion of cleptoparasites increased (Figure 3B).

4. DISCUSSION

Traditional approaches to bee diversity are not always fully informative with respect to the status of the habitats they represent, and in our previous analysis of these data, rarified estimates of species richness indicated that only in the most heavily disturbed habitats (i.e. COMM-A) was there significantly reduced diversity; no differences were found among all other habitat types (Sheffield et al. 2013). Other species richness estimators (i.e. “hidden” species in truncated log-normal plots, extrapolation-based methods) indicated that the natural habitats had significantly more species than all other habitats, which did not differ. In that study, Sheffield et al. (2013) indicate that these results suggest that agro-ecosystems with intermediate levels of natural habitat do or do not encourage bee species richness based on how the data are analysed. As such, diversity data alone may not always reflect habitat quality for bees, especially at intermediate levels of disturbance. Similarly, evaluating these bee communities based on diversity and abundance data, particularly measuring deviations in the fit to the truncated log-normal distribution, did not diagnose the “health” of these systems; all four habitats were not normally distributed (see Sheffield et al. 2013 for discussion). However, partitioning these plots to reveal cleptoparasite

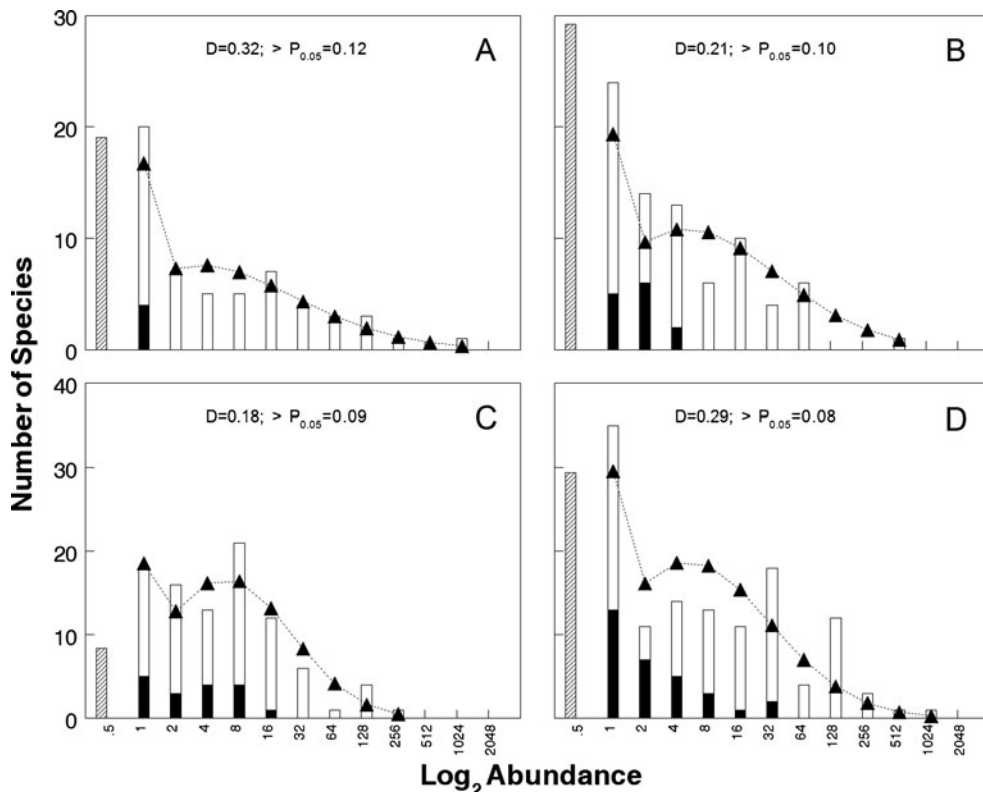


Figure 1. Truncated log-normal plot (\log_2 scale) of observed (*solid bars*) and expected (*triangles*) bee species abundances, and estimated “unseen” species (*diagonally striped bars*) in **A** COMM-A, **B** COMM-B, **C** ABAND and **D** WILD habitats for 2001–2002. Kolmogorov–Smirnov test for normality with Lilliefors adjustment do not support a log normal distribution for any habitat type. *Black sections of bars* represent the number of cleptoparasites in each abundance octave.

diversity and abundance (Figure 1) clearly demonstrated that this guild was virtually lacking in the COMM-A habitat, becoming more diverse and comprising a larger proportion of the bee community as the landscape became more diverse with unmanaged habitat (Figures 2 and 3A).

Additional analyses that incorporate life history traits have much to offer to further understand bee communities and their responses to disturbances (Williams et al. 2010; Neame et al. 2012; Sheffield et al. 2013). Bees constitute an incredibly diverse assemblage of life histories, social structure, nesting biology, lifestyles, etc. (Michener 2007; Williams et al. 2010), which make it unreasonable to generalise the group as a whole, especially if habitat compar-

isons and/or health evaluation are the subjects of investigation. Specific guilds respond differently due to specific requirements (Moretti et al. 2009; Neame et al. 2012), and incorporation of guilds into studies of bee communities may serve as an additional metric for measuring the status of the ecosystems (Sheffield et al. 2013). Natural habitats may have specific guild profiles; once determined (with enough regional sampling), these profiles may serve as benchmarks for pollinator community studies. As these profiles are guild-based and not exclusively taxonomy-based, comparisons across broader ecosystems may be possible, providing additional criteria to diagnose habitats and allowing comparisons among ecosystems that

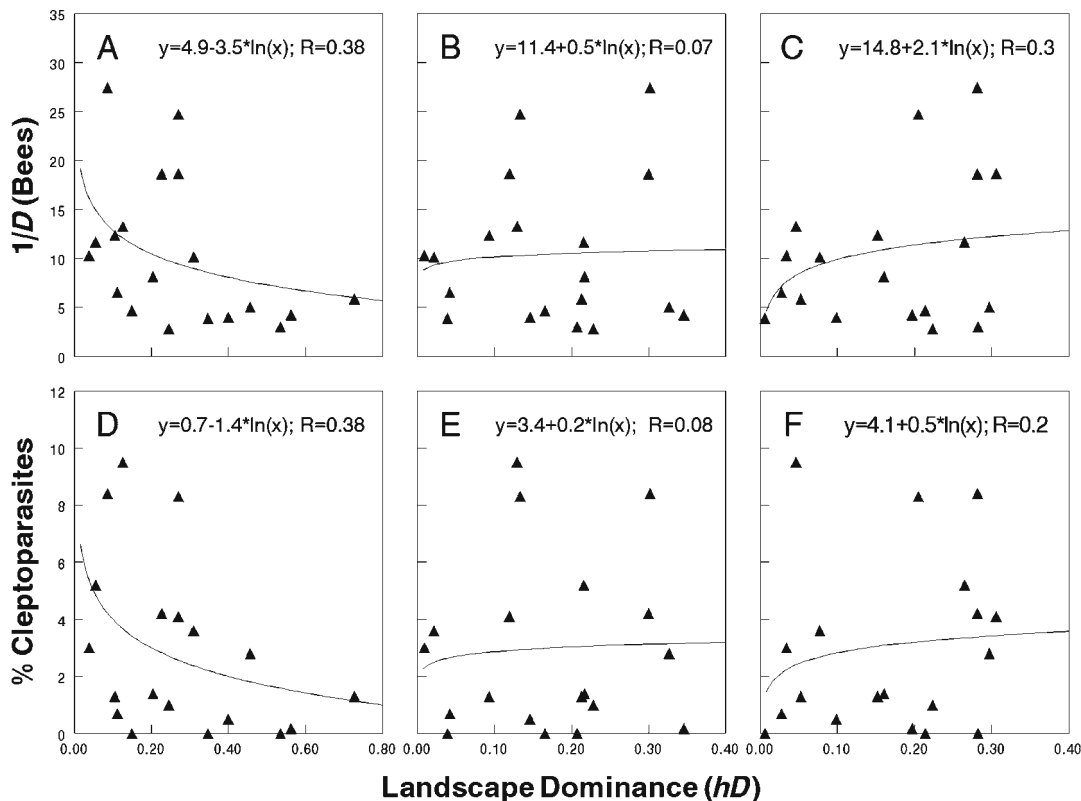


Figure 2. The diversity of bees ($1/D$) (A–C) and percent cleptoparasites (D–F) versus landscape dominance (h_D) for each site at three landscape levels surrounding the sampled area: $200 \times 200\text{m}$ (A, D), $400 \times 400\text{m}$ (B, E) and $600 \times 600\text{m}$ (C, F).

differ with respect to levels of disturbance or management (Tilman and Lehman 2001; Oertli et al. 2005). The rationale behind this approach is that loss of certain guilds, or functional diversity, within ecosystems can contribute to ecosystem collapse (O’Gorman et al. 2011), while the loss of species themselves may be buffered through overall species richness and redundancy (Walker 1992; Peterson et al. 1998; Elmqvist et al. 2003).

The quantity and quality of resources available to bees varies greatly among habitats, often attributable to the characteristics of the surrounding landscape (Kremen et al. 2002). Habitat type had a large impact on the proportional abundance of the different bee guilds (Sheffield et al. 2013), and strong negative responses to intense agriculture were observed

in cleptoparasites in general (Figure 1), but more specifically in cleptoparasite diversity (Figure 3A). Reductions in cleptoparasite abundance in itself will cause an increase in the calculated proportion of other guilds, but when species alone are considered, no differences in the proportion of the guilds nor their diversity were observed (Sheffield et al. 2013).

There is a growing body of research suggesting that parasites and parasitism (in the general sense), like predation, are important factors in maintaining ecosystem health and stability (e.g. Combes 1996; Morand and Gonzalez 1997; Horwitz and Wilcox 2005; Marcogliese 2004; Hudson et al. 2006; Wood et al. 2007). In essence, cleptoparasitic bees may perform many of the same roles as true parasites. Despite the fact they are free living and kill the offspring of

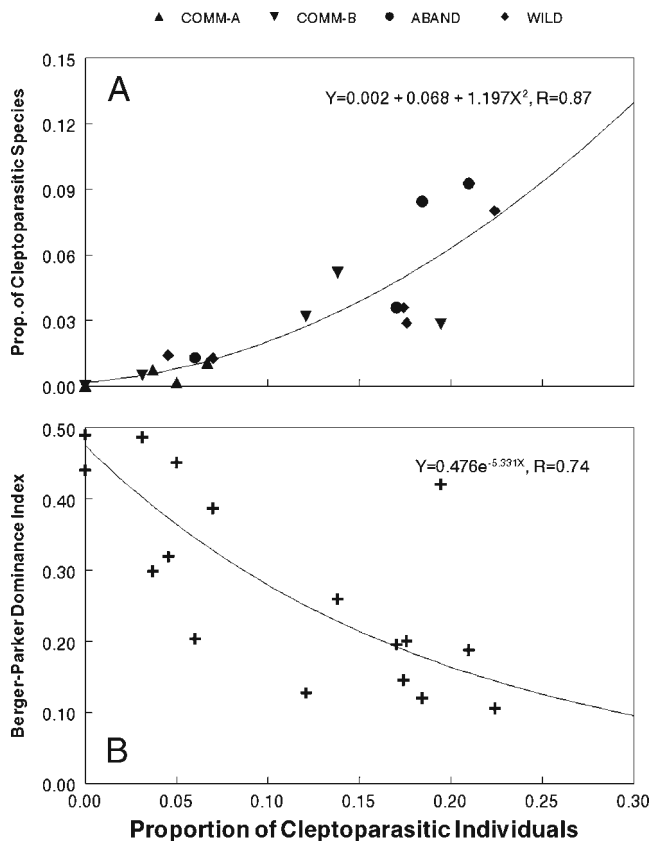


Figure 3. **A** The proportion of cleptoparasitic bee species versus individuals for sites from each habitat. **B** The Berger–Parker Dominance Index versus the proportion of cleptoparasitic individuals.

their host, like true parasites, they are reducing the fecundity of their host; depending on the taxon, either the cleptoparasitic larva kills the host egg (or developing larva) or the cleptoparasitic female destroys the host egg before she oviposits (Rozen 2001). Combes (1996) suggested that parasites perform a stabilising role, called “parasite arbitration” by effecting competition among hosts; cleptoparasitism itself is a form of competition (Iyengar 2008). In this study, sites with diverse and abundant cleptoparasite assemblages had lower dominance levels than in sites with few cleptoparasites (Figure 3B) supporting the idea that a stabilising role is being performed. Abundant host taxa are probably the most likely to be attacked in a given site and/or year, which may reduce competition among non-parasitic bee species;

cleptoparasite activity is positively correlated with host nest density, not number (Polidori et al. 2009), though Rosenheim (1990) suggested that pressure from parasites (including cleptoparasites) may favour either aggregated or widely dispersed nesting strategies of host taxa.

The proportion of cleptoparasitic species to host species is usually low (Wcislo 1981), indicating that many cleptoparasites are probably generalists with respect to their hosts, though individuals probably focus on one host species during their lifetime (Bogusch et al. 2006). Cleptoparasites can make up a significant proportion of bee species in a geographic area, and many of the “rare” species in surveys belong to this guild (Oertli et al. 2005). They are ubiquitous, and the structure of cleptoparasitic bee communities follows those of the

remaining bee community (Figure 2); there are generalist and specialist cleptoparasitic bees (though host information is known for very few), and their species richness and abundance is dictated by their hosts.

We believe that, among the guilds responding to different levels of disturbance, cleptoparasites have the greatest potential as indicator taxa for assessing bee communities. First, assigning bees to this guild is very easy; cleptoparasitic bees are, for the most part, easily recognised as such (Michener 2007). In contrast, the details of nesting biology of few bee species are actually known; megachilid bees, in particular, show such great variety of nesting habits, and some halictid bees (a large component of both solitary and social ground nesters) show even intraspecific variation in sociality (Michener 1974, 2007) that generalisations even at the level of genus are risky. Second, species richness often decreases upward from basal trophic levels in natural communities (Duffy 2003; Petchey et al. 2004). As such, the biology of cleptoparasites make them suitable as indicator taxa in a fashion similar to top predators and parasites (Marcogliese and Cone 1997) as they form the apex of bee communities; their presence within habitats is dependent on the presence of their host(s) and the resources available to and/or affecting these hosts (Finke and Denno 2004). The managed orchards had significantly fewer cleptoparasites (Figure 1A) despite these sites having the highest proportion of potential host species (i.e. solitary and social ground nesting bees) (Sheffield et al. 2013). As such, cleptoparasites were responding negatively to aspects of the sites in lieu of abundant host taxa. Disturbances within habitats that affect the species richness and/or fecundity of non-parasitic taxa through decline in availability of the resources needed by these bees may first be noticed in the relative abundance and diversity of cleptoparasites. Third, cleptoparasites are a diverse guild, and this lifestyle represents multiple independent origins in most bee families (Michener 2007) with a diverse assemblage of hosts. The effect of habitat disturbances on one or many host clades is likely to be observed and/or measurable within the cleptoparasitic guild as a whole. As such, and

unlike other guilds, the cleptoparasite guild responds in ways that are reflective of the entire bee community (Figure 2), and probably serve as sensitive indicator taxa for assessing the status of ecosystems.

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Le potentiel des abeilles cleptoparasites comme taxa indicateurs dans l'évaluation des communautés d'abeilles

Communauté des pollinisateurs / guildes / santé des écosystèmes / espèces indicatrices

Die Möglichkeit, kleptoparasitische Bienen als Indikatoren zur Beurteilung von Bienengemeinschaften zu verwenden

Bestäubergemeinschaften / Gildenstrukturen / Kleptoparasiten / Indikatortaxa / Ökosystemstatus

REFERENCES

- Balvanera, P., Pfisterer, A.B., Buchmann, N., He, J., Nakashizuka, T., Raffaelli, D., Schmid, B. (2006) Quantifying the evidence for biodiversity effects on ecosystem functioning and services. *Ecol. Lett.* **9**, 1146–1156
- Biesmeijer, J.C., Roberts, S.P.M., Reemer, M., Ohlemüller, R., Edwards, M., et al. (2006) Parallel declines in pollinators and insect-pollinated plants in Britain and The Netherlands. *Science* **313**, 351–354
- Blondel, J. (2003) Guilds or functional groups: does it matter? *Oikos* **100**, 223–231

- Bogusch, P., Kratochvíl, L., Straka, J. (2006) Generalist cuckoo bees (Hymenoptera: Apoidea: *Sphcodes*) are species-specialist at the individual level. *Behav. Ecol. Sociobiol.* **60**, 422–429
- Cane, J.H., Minckley, R., Roulston, T., Kervin, L.J., Williams, N.M. (2006) Complex responses within a desert bee guild (Hymenoptera: Apiformes) to urban habitat fragmentation. *Ecol. Appl.* **16**, 632–644
- Cardinale, B.J., Srivastava, D.S., Duffy, J.E., Wright, J.P., Downing, A.L., Sankaran, M., Jouseau, C. (2006) Effects of biodiversity on the functioning of trophic groups and ecosystems. *Nature* **443**, 989–992
- Chiarucci, A., Giovanni, B., Scheiner, S.M. (2011) Old and new challenges in using species diversity for assessing biodiversity. *Philos. Trans. R. Soc. London, Ser. B* **366**, 2426–2437
- Combes, C. (1996) Parasites, biodiversity and ecosystem stability. *Biodivers. Conserv.* **5**, 953–962
- Duffy, J.E. (2003) Biodiversity loss, trophic skew and ecosystem functioning. *Ecol. Lett.* **6**, 680–687
- Duffy, J.E., Cardinale, B.J., France, K.E., McIntyre, P.B., Thébault, E., Loreau, M. (2007) The functional role of biodiversity in ecosystems: incorporating trophic complexity. *Ecol. Lett.* **10**, 522–538
- Elmqvist, T., Folke, C., Nyström, M., Peterson, G., Bengtsson, J., Walker, B., Norberg, J. (2003) Response diversity, ecosystem change, and resilience. *Front. Ecol. Environ.* **1**, 488–494
- Finke, D.L., Denno, R.F. (2004) Predator diversity dampens trophic cascades. *Nature* **429**, 407–410
- Grundel, R., Frohnapple, K.J., Jean, R.P., Pavlovic, N.B. (2011) Effectiveness of bowl trapping and netting for inventory of a bee community. *Environ. Entomol.* **40**, 374–380
- Hooper, D.U., Chapin, F.S., Ewel, J.J., Hector, A., Inchausti, P., et al. (2005) Effects of biodiversity on ecosystem functioning: a consensus of current knowledge. *Ecol. Monogr.* **75**, 3–35
- Horwitz, P., Wilcox, B.A. (2005) Parasites, ecosystems and sustainability: an ecological and complex systems perspective. *Int. J. Parasitol.* **35**, 725–732
- Hubbell, S.P. (2001) *The Unified Neutral Theory of Biodiversity and Biogeography*. Princeton University Press, Princeton
- Hudson, P.J., Dobson, A.P., Lafferty, K.D. (2006) Is a healthy ecosystem one that is rich in parasites? *Trends Ecol. Evol.* **21**, 381–385
- Iyengar, E.V. (2008) Kleptoparasitic interactions throughout the animal kingdom and a re-evaluation, based on participant mobility, of the conditions promoting the evolution of kleptoparasitism. *Biol. J. Linn. Soc.* **93**, 745–762
- Kevan, P.G., Greco, C.F., Belaussoff, S. (1997) Log-normality of biodiversity and abundance in diagnosis and measuring of ecosystem health: pesticide stress on pollinators on blueberry heaths. *J. Appl. Ecol.* **34**, 1122–1136
- Klein, A.M., Vassiere, B.E., Cane, J.H., Steffan-Dewenter, I., Cunningham, S.A., Kremen, C., Tscharntke, T. (2007) Importance of pollinators in changing landscapes for world crops. *Proc. R. Soc. B* **274**, 303–313
- Kremen, C., Williams, N.M., Thorp, R.W. (2002) Crop pollination from native bees at risk from agricultural intensification. *Proc. Natl. Acad. Sci. USA* **99**, 16812–16816
- Kremen, C., Williams, N.M., Bugg, R.L., Fay, J.P., Thorp, R.W. (2004) The area requirements of an ecosystem service: crop pollination by native bee communities in California. *Ecol. Lett.* **7**, 1109–1119
- Magurran, A.E. (2004) *Measuring Biological Diversity*. Blackwell, Malden
- Marcogliese, D.J. (2004) Parasites: small players with crucial roles in the ecological theater. *EcoHealth* **1**, 151–164
- Marcogliese, D.J., Cone, D.K. (1997) Food webs: a plea for parasites. *Trends Ecol. Evol.* **12**, 320–325
- Michener, C.D. (1974) *The Social Behavior of the Bees*. Harvard University Press, Boston
- Michener, C.D. (1979) Biogeography of the bees. *Ann. Mo. Bot. Gard.* **66**, 277–347
- Michener, C.D. (2007) *The Bees of the World*, 2nd edn. Johns Hopkins University Press, Baltimore
- Morand, S., Gonzalez, E.A. (1997) Is parasitism a missing ingredient in model ecosystems? *Ecol. Model.* **95**, 61–74
- Moretti, M., de Bello, F., Roberts, S.P.M., Potts, S.G. (2009) Taxonomical vs. functional responses of bee communities to fire in two contrasting climatic regions. *J. Anim. Ecol.* **78**, 98–108
- National Research Council (2007) *Status of Pollinators in North America*. National Academies Press, Washington
- Neame, L.A., Griswold, T., Elle, E. (2012) Pollinator nesting guilds respond differently to urban habitat fragmentation in an oak-savannah ecosystem. *Insect Conserv. Divers.* **6**, 57–66. doi:10.1111/j.1752-4598.2012.00187.x
- Nielsen, A., Steffan-Dewenter, I., Westphal, C., Messinger, O., Potts, S.G., et al. (2011) Assessing bee species richness in two Mediterranean communities: importance of habitat type and sampling techniques. *Ecol. Res.* **26**, 969–983
- O’Neill, R.V., Krummel, J.R., Gardner, R.H., Sugihara, G., Jackson, B., et al. (1988) Indices of landscape pattern. *Landscape Ecol.* **1**, 153–162
- Oertli, S., Muller, A., Dorn, S. (2005) Ecological and seasonal patterns in the diversity of a species-rich bee assemblage (Hymenoptera: Apoidea: Apiformes). *Eur. J. Entomol.* **102**, 53–63
- O’Gorman, E.J., Yearsley, J.M., Crowe, T.P., Emmerson, M.C., Jacob, U., Petchey, O.L. (2011) Loss of functionally unique species may gradually undermine ecosystems. *Proc. R. Soc. Lond. B* **278**, 1886–1893

- Petchey, O.L., Hector, A., Gaston, K.J. (2004) How do different measures of functional diversity perform? *Ecology* **85**, 847–857
- Peterson, G., Allen, C.R., Holling, C.S. (1998) Ecological resilience, biodiversity, and scale. *Ecosystems* **1**, 6–18
- Polidori, C., Borruso, L., Boesi, R., Andrietti, F. (2009) Segregation of temporal and spatial distribution between kleptoparasites and parasitoids of the eusocial sweat bee, *Lasioglossum malachurum* (Hymenoptera: Halictidae, Mutillidae). *Entomol. Sci.* **12**, 116–129
- Rosenheim, J.A. (1990) Density-dependent parasitism and the evolution of nesting aggregations in the solitary Hymenoptera. *Ann. Entomol. Soc. Amer.* **83**, 277–286
- Roulston, T.H., Smith, S.A., Brewster, A.L. (2007) A comparison of pan trap and intensive net sampling techniques for documenting a bee (Hymenoptera: Apiformes) fauna. *J. Kansas Entomol. Soc.* **80**, 179–181
- Rozen Jr., J.G. (2001) A taxonomic key to mature larvae of cleptoparasitic bees (Hymenoptera: Apoidea). *Amer. Mus. Nov.* **3309**, 1–28
- Scrosati, R.A., van Genne, B., Heaven, C.S., Watt, C.A. (2011) Species richness and diversity in different functional groups across environmental stress gradients: a model for marine rocky shores. *Ecography* **34**, 151–161
- Sheffield, C.S., Kevan, P.G., Westby, S.M., Smith, R.F. (2008) Diversity of cavity-nesting bees (Hymenoptera: Apoidea) within apple orchards and wild habitats in the Annapolis Valley, Nova Scotia, Canada. *Can. Ent.* **140**, 235–249
- Sheffield, C.S., Kevan, P.G., Pindar, A., Packer, L. (2013) Bee (Hymenoptera: Apoidea) diversity within apple orchards and old fields habitats in the Annapolis Valley, Nova Scotia, Canada. *Can. Ent.* **145**, 94–114
- Tilman, D., Lehman, C. (2001) Biodiversity, composition, and ecosystem processes: theory and concepts. In: Kinzig, A.P., Pacala, S.W., Tilman, D. (eds.) *The Functional Consequences Of Biodiversity: Empirical Progress and Theoretical Extensions*, pp. 9–41. Princeton University Press, Princeton
- Toler, T.R., Evans, E.W., Tepedino, V.J. (2005) Pan-trapping for bees (Hymenoptera: Apiformes) in Utah's West Desert: the importance of color diversity. *Pan-Pac. Entomol.* **81**, 103–113
- Walker, B.H. (1992) Biological diversity and ecological redundancy. *Conserv. Biol.* **6**, 18–23
- Wcislo, W.T. (1981) The roles of seasonality, host synchrony, and behaviour in the evolutions and distributions of nest parasites in the Hymenoptera (Insecta), with special reference to bees (Apoidea). *Biol. Rev.* **62**, 515–543
- Westphal, C., Bommarco, R., Carré, G., Lamborn, E., Morison, N., et al. (2008) Measuring bee diversity in different European habitats and biogeographical regions. *Ecol. Mono.* **78**, 654–671
- Williams, N.M., Crone, E.E., Roulston, T.H., Minckley, R.L., Packer, L., Potts, S.G. (2010) Ecological and life-history traits predict bee species responses to environmental disturbances. *Biol. Conserv.* **143**, 2280–2291
- Wood, C.L., Byers, J.E., Cottingham, K.L., Altman, I., Donahue, M.J., Blakeslee, A.M.H. (2007) Parasites alter community structure. *Proc. Natl. Acad. Sci. USA* **104**, 9335–9339