1 Predicting changes in bee assemblages following state transitions in North American drylands

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16 Drylands worldwide are experiencing ecosystem state transitions: the expansion of some 17 ecosystem types at the expense of others. Bees in drylands are particularly abundant and diverse, with 18 potential for large compositional differences and seasonal turnover across ecotones. To better 19 understand how future ecosystem state transitions may influence bees, we compared bee assemblages 20 and their seasonality among three dryland ecosystem types of the southwestern U.S. (Plains grassland, 21 Chihuahuan Desert grassland, and Chihuahuan Desert shrubland). Using passive funnel traps, we caught 22 bees during two-week intervals from March through October during 2002 – 2014. The resulting dataset 23 included 302 bee species and >70,500 individuals. Bee abundance, composition, and diversity differed 24 among ecosystems, indicating the potential for future ecosystem state transitions to alter bee assemblage 25 composition in drylands. We also found strong seasonal turnover in bee species, suggesting that bee 26 phenological shifts may accompany ecosystem state transitions. Common rather than rare species drove 27 the observed trends, and both specialist and generalist bee species were indicators of each ecosystem 28 type or month; these species could be informative sentinels of community-wide responses to future shifts. 29 Our work suggests that predicting the consequences of global change for bee assemblages will require 30 accounting for both within-year and among-ecosystem variation.

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32 Introduction

33 Drylands worldwide are experiencing ecosystem state transitions: the expansion of some 34 ecosystem types at the expense of others^{1,2}. These transitions include encroachment of C₃ shrubland into 35 C_4 grassland³ and conversion of woodland to savanna⁴. It is through these transitions that the largest 36 changes in dryland ecosystem processes are occurring⁵⁻⁷. State transitions can produce dramatic 37 changes in carbon fluxes^{8,9}, nutrient dynamics^{10,11}, spatial heterogeneity in vegetation¹², and consumer 38 community composition^{13,14}. Because drylands cover ~45% of land area on Earth¹⁵ and support over 2 39 billion people¹⁶, understanding how much dryland ecosystems currently differ in community composition 40 can help to predict changes in future communities — and the ecosystem services they provide — under 41 state transitions.

Bees may serve as important bio-indicators of state transitions and sentinels of altered
 ecosystem services^{17,18}. In drylands, bees are important pollinators of both wild plants and agricultural

44 crops^{19,20}, and are particularly abundant and diverse. North America's highest bee diversity occurs in the 45 southwestern U.S. and northwest Mexico, and 75% of the continent's bee species are found in the 46 western U.S.^{21,22}. Relative to mesic ecosystems, drylands can also host higher proportions of specialist 47 bee species, which pollinate one or a few closely related plant species²³. For example, creosote bush 48 (Larrea tridentata (DC.) Coville), a widespread and abundant shrub in North American warm deserts²⁴, is 49 visited by 22 documented specialist bee species²⁵. Cacti also host many specialists²⁶. Communities 50 dominated by specialist bees may be less resilient to state changes or pollinator declines than 51 communities dominated by generalist bees, which can buffer plants against crashes in other bee 52 species^{27,28}. Future ecosystem state transitions could therefore substantially influence bees in drylands, 53 making it important to understand potential vulnerabilities of dryland bee assemblages to these shifts. 54 Understanding variation in bee composition among habitat types can shed light on how 55 ecosystem state transitions will influence bee assemblages. Prior studies have largely focused on bee 56 assemblage variation within agricultural environments, along urban-rural gradients, or with habitat 57 fragmentation²⁹⁻³¹, while fewer studies have compared natural ecosystems. For instance, in Spain, shrub 58 encroachment into grasslands corresponded with higher pollinator richness but fewer pollinator visits to 59 forbs³². In xeric environments, some studies have documented bee species turnover across relatively 60 small spatial scales^{25,33,34}. For instance, during a single growing season, one study found lower bee 61 abundance and richness in desert scrubland relative to riparian sites within a 4 km² area in the Sonoran 62 Desert³³. In contrast, abundances of insect pollinator functional groups did not differ between creosote 63 bush-dominated and adjacent annual forb-dominated microsites in the Mojave Desert³⁵, although this 64 study occurred on a smaller spatial scale with coarser taxonomic resolution. These contrasting results 65 highlight the need for additional data to better predict the potential consequences for bee assemblages of 66 specific state transitions in dryland ecosystems.

In addition, seasonal turnover in bee species composition suggests the potential for climate change to produce shifts in bee phenology³⁶. Some bees may cue on climate variables for their emergence as adults, with temperature or precipitation conditions triggering the emergence of bee species at different times of year^{37–39}. High temporal turnover in bee assemblage composition could thus indicate dominance of species with phenologies closely tied to climate, which may be particularly

72 susceptible to phenological shifts under climate change. Understanding bee assemblage seasonality in 73 ecosystem types predicted to expand or contract under climate change could thus be important for 74 predicting bee assemblage responses to state transitions. However, while bee composition is well 75 documented to vary seasonally within a community⁴⁰⁻⁴², few studies have compared seasonal patterns 76 among ecosystem types to discern how state transitions may shift bee phenology at the landscape scale. 77 Seasonal trends in bee abundance and richness were found to differ between natural and human-altered 78 landscape types during a single year in California, USA⁴³, and among agricultural land use classes during 79 3 years in New Hampshire, USA⁴⁴. However, we lack studies that use long-term data to elucidate how 80 general patterns of bee seasonality differ among natural ecosystem types that are expanding versus 81 contracting.

82 This study compared bee assemblages and their seasonality among sites representing three 83 dryland ecosystem types of the southwestern U.S.: Chihuahuan Desert shrubland, Chihuahuan Desert 84 grassland, and Plains grassland. Our sites occurred within a relatively small area (within 2-10 km of one 85 another) that encompassed ecotones between the types, and shared the same regional pool of bee 86 species. We used 13 years of monthly bee trap data to address two questions: (1) How much do bee 87 assemblage abundance, composition, and diversity differ among major southwestern U.S. ecosystem 88 types? (2) Do dryland ecosystem types differ in their degree of seasonal variation in bee abundance, 89 composition, or diversity? We examined patterns among ecosystem types and months of the year by 90 averaging across the time series, enabling us to identify general trends. Whereas this analysis focused on 91 intra-annual and among-habitat variation in bee composition, a companion study will report inter-annual 92 change over the time series, providing substantial additional complexity to the analysis.

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94 Methods

95 Ecosystem types

The Sevilleta National Wildlife Refuge (NWR; Socorro, NM) includes five ecosystem types that
together represent ~80 million ha of the southwestern U.S. We focused on three major ecosystem types:
Chihuahuan Desert shrubland, which is dominated by creosote bush (*Larrea tridentata*), Chihuahuan
Desert grassland, which is dominated by black grama grass (*Bouteloua eriopoda* (Torr.) Torr.), and Plains

grassland, which is dominated by blue grama grass (*Bouteloua gracilis* (Willd. Ex Kunth) Lag. Ex Griffiths)
(Table 1). Transitions among these ecosystem types are predicted to occur under climate change, with
Chihuahuan Desert shrubland encroaching upon Chihuahuan Desert grassland, which is predicted to
replace Plains grassland^{45–47}. In our study, the two Chihuahuan Desert sites were separated by ~2 km;
the Plains grassland site was ~10 km from the Chihuahuan Desert sites (Table 1).

105 Bee collection

106 Bees were sampled along five transects located within each of the three focal ecosystem types. 107 To sample bees, we installed one passive funnel trap at each end of five 200 m transects/site. Each trap 108 consisted of a 946 mL paint can filled with ~275 mL of propylene glycol and topped with a plastic 109 automotive funnel with the narrow part of the funnel sawed off (funnel height = 10 cm, top diameter = 14 110 cm, bottom diameter = 2.5 cm; Supplementary Fig. S1). The funnels' interiors were painted with either 111 blue or vellow fluorescent paint (Krylon, Cleveland, OH or Ace Hardware, Oak Brook, IL). On each 112 transect, we randomly assigned one trap to be blue and the other to be yellow (total across the three 113 sites: N = 30 traps, with 15 traps/color). Because different bee taxa are known to be attracted to blue 114 versus yellow⁴⁸, we summed the samples collected in the two traps on a given transect. Each trap was 115 placed on a 45 cm high platform that was surrounded by a 60 cm high chicken wire cage to prevent 116 wildlife and wind disturbance (Supplementary Fig. S1). Funnel traps provide a measure of bee activity, 117 not a measure of presence, and may be biased by bee taxon and sociality^{49,50}. From 2002 to 2014, bees 118 were sampled each month from March through October. Traps were opened each March as close as 119 possible to the first day of spring, and left open for 14 d, after which the bee specimens were collected. 120 The traps were then closed for 14 d. This two-week cycle was repeated through October. Bees were 121 rinsed and stored in 70% ethanol until processed.

122 Bee identification

Bees were identified to species by K.W.W. and T.L.G. Certain groups of bees could not be identified to species, either because there are no practicing experts in the bee group and species are unnamed for our study region, or because there are no revisions within the bee group to separate named from unnamed species. In these cases, we separated females into morphotypes as best as possible. The males of these groups could not be reliably linked to the females and were therefore excluded from the 128 dataset. The major groups treated in this manner were the genera Sphecodes, Pseudopanurgus, and 129 Nomada, the subgenera Dialictus and Evylaeus of the genus Lasioglossum, and the subgenus 130 Micrandrena of Andrena. We excluded Nomada from our analyses due to low abundance and lack of 131 ability to distinguish among species. New species of relatively well-known genera were recognized, and 132 the qualifiers nr. or aff. were used with uncertain identifications. Voucher specimens were deposited at 133 the University of New Mexico's Museum of Southwestern Biology and the USDA-ARS Pollinating Insects 134 Research Unit's U.S. National Pollinating Insects Collection. Information related to these specimens is 135 available via the Symbiota Collections of Arthropods Network (https://scan-bugs.org). 136 Analysis

137 Dataset. We created a species matrix in which cells contained the mean abundance of each bee 138 species for each month of collection, averaged over the years of collection (2002 - 2014). Each row was 139 a unique trapping transect, with five transects per ecosystem type per month (N = 120 observations). 140 Means were calculated using the <reshape2> package⁵¹ in R version 3.4.2⁵². To examine whether 141 assemblage-level patterns were driven by common or rare species, we ran all abundance, composition. 142 and diversity analyses (described below) on the full dataset, on a dataset with singleton bee species 143 (those caught only on a single transect, in a single month) removed, and finally on a subset of the dataset 144 containing only the bee species that were present in >5% of the samples.

145 Overview. Analyses addressed our two key guestions within one set of statistical models 146 (described below). First, (1) How much do bee assemblage abundance, composition, and diversity differ 147 among major southwestern U.S. ecosystem types? was determined by the statistical significance and 148 magnitude of the effect of ecosystem type in our models. We also compared the effect size of ecosystem 149 type against the effect size of month of sampling to estimate the relative importance of inter-ecosystem 150 versus seasonal variability. Then, to address (2) Do dryland ecosystem types differ in their degree of 151 seasonal variation in bee abundance, composition, or diversity? we evaluated whether the interaction 152 between ecosystem type and month of sampling was statistically significant, indicating that ecosystems 153 differed in the seasonality of bee abundance, composition, or diversity.

154Bee assemblage composition and turnover.For bee composition, we calculated Bray-Curtis155similarities in Primer version 6.1.1353. We then tested for the influence of ecosystem type, month of

156 sampling, and the random effect of transect, which was nested within ecosystem type to account for the 157 repeated measures design, using perMANOVA (version 1.0.3) with 9999 permutations of residuals under 158 a reduced model. We additionally examined whether ecosystem types or months differed in bee 159 assemblage dispersion using permDISP in Primer⁵³. We visualized assemblage composition with non-160 metric multidimensional scaling analysis (NMDS) implemented with 500 restarts in Primer. For each 161 ecosystem type, we assessed bee species turnover among months, as well as the rate of community 162 change, using the <codyn> package in \mathbb{R}^{54} . Finally, to identify which taxa contributed most to bee 163 assemblage (i) divergence among ecosystem types and (ii) divergence among months within each 164 ecosystem type, we calculated Dufrene-Legendre indicator species values using the indval function in the 165 <labdsv> R package⁵⁵. 166 Bee diversity and abundance. For bee diversity, we calculated the Shannon diversity index (H'), 167 species richness, and evenness (Pielou's J) using the $\langle vegan \rangle$ package in R⁵⁶. We then used linear 168 mixed effects models to examine the influences of ecosystem type, sampling month, and their interaction 169 (fixed effects), as well as transect identity (random effect nested within ecosystem type), on these three 170 responses, as well as on total bee abundance (function Imer, <Ime4> package in R)⁵⁷. When there was a 171 significant ecosystem type x sampling month interaction, we tested a priori contrasts for pairs of months

within each ecosystem type and for pairs of ecosystem types within each month using Tukey-Kramer
 multiple comparisons in the <emmeans> package in R⁵⁸.

174

175 Results

176 The dataset

We captured a total of 70,951 individuals representing 302 species during the 13 years of monthly trapping (see Supplementary Table S1 for a full species list). Species were distributed across 6 families and 56 genera (Supplementary Table S1 and Fig. S2). Our dataset was dominated by a small number of abundant species and contained a large number of rare species (Supplementary Fig. S3). The most commonly collected species were *Lasioglossum semicaeruleum* (36% of all collected specimens), *Agapostemon angelicus* (21%), *Diadasia rinconis* (7%), *Melissodes tristis* (5%), *Anthophora affabilis* (5%),

and *Eucera lycii* (3%). Amongst the collected species, 30% were singletons, and 58% were found in <5%
of all samples.

185 Bee assemblage composition: temporal variation surpassed differences among dryland ecosystem types 186 Variation among ecosystems. All ecosystems significantly diverged in bee assemblage 187 composition, and this pattern was present during all months (Table 2, Fig. 1). The greatest difference 188 among ecosystems occurred in October, when the Plains grassland bee assemblage diverged most 189 strongly from the Chihuahuan Desert shrubland (mean similarity: 41.4, P = 0.0089) and also diverged 190 from the Chihuahuan Desert grassland (mean similarity: 51.6, P = 0.0080). The three ecosystem types 191 did not differ in assemblage dispersion ($F_{2,117} = 0.52$, P = 0.71), indicating similar levels of temporal beta-192 diversity among ecosystem types (Fig. 1). 193 Indicators of variation among ecosystems. We identified 43 bee species as ecosystem indicators 194 according to their Dufrene-Legendre (DL) indicator species values (Table 3). Of these, 21 species were 195 indicators of Chihuahuan Desert shrubland, 14 species were indicators of Plains grassland, and 8 species 196 were indicators of Chihuahuan Desert grassland. All three ecosystems had indicator species within the 197 families Andrenidae, Apidae, Halictidae, and Megachilidae, and one Plains grassland indicator species 198 was in the family Colletidae (Table 3, Fig. 2). In all three ecosystems, Lasioglossum semicaeruleum (an 199 indicator of the Desert grassland), Agapostemon angelicus (an indicator of Plains grassland), Diadasia 200 rinconis, and Melissodes tristis were among the five most abundant bee species (Fig. 2). Anthophora 201 affabilis was also within the five most abundant species in the Plains and Chihuahuan Desert grasslands,

while *Perdita larreae* (a creosote bush specialist) was abundant in and an indicator of the Chihuahuan

203 Desert shrubland (Fig. 2).

204Temporal variation.The month of sample collection explained an order of magnitude more205variation in bee assemblage composition than did ecosystem type (Table 2, Fig. 3). Generally,206assemblages diverged between the early and late months of the year and converged during the middle of207the summer. Across ecosystems, the pair of months most divergent in bee composition was March versus208October (mean similarity = 12.7, P = 0.0001). In contrast, June and July were most similar in bee209composition (mean similarity = 64.0, P = 0.0001).

Months additionally differed from one another in the magnitude of assemblage dispersion, a metric that captures the degree of beta-diversity across both sites and transects (Table 2, Fig. 3). The strongest differences in beta-diversity were between March or June, which had the smallest multivariate dispersions (mean \pm s.e., March: 21.0 \pm 1.5, June: 20.4 \pm 0.8), against October, which had the largest average dispersion across ecosystems (29.8 \pm 1.8).

215 Bee abundance and diversity: temporal variation exceeded variation among dryland ecosystems

216Abundance. As with composition, across months, ecosystems diverged significantly from one217another in total bee abundance (Table 4). Bee abundance was on average 43% lower in the Chihuahuan218Desert shrubland relative to the two grassland sites from March through July (Fig. 4a). However,219abundances within the ecosystems converged in August, and abundance differences disappeared in220September and October (Fig. 4a), as indicated by a significant interaction between ecosystem type and221month of collection (Table 4: Ecosystem x Month, P < 0.0001).

222 Diversity. Ecosystems also diverged in bee diversity as measured by the Shannon index and 223 Pielou's evenness (Table 4). Differences in Shannon diversity (Fig. 4b) among ecosystems were more 224 strongly driven by evenness (Fig. 4d) than by richness (Fig. 4c). On average across all months of 225 sampling, the Chihuahuan Desert shrubland ecosystem had the highest bee Shannon diversity and 226 evenness, with these diversity metrics 5% (Shannon diversity) and 2% (evenness) higher than in the 227 Plains grassland. In turn, Plains grassland diversity metrics were 16% (Shannon diversity) and 12% 228 (evenness) higher than the Chihuahuan Desert grassland. In contrast, on average across months, the 229 ecosystems did not significantly differ in bee species richness (Table 4, Fig. 4c).

230 Importantly, differences among ecosystems in all diversity metrics varied by month of the year 231 (Fig. 4, Table 4: Ecosystem x Month – Shannon diversity: P < 0.0001, richness: P = 0.0137, evenness: P 232 < 0.0001), indicating that dryland ecosystem types differed in their degree of seasonal variation in bee 233 diversity (Question 2). Specifically, Shannon diversity was greater in the Chihuahuan Desert shrubland 234 than in the Desert grassland in all months except for March; differences in Shannon diversity were largest 235 in May and September, when Shannon diversity respectively was 38% and 33% higher in the 236 Chihuahuan Desert shrubland relative to grassland (Fig. 4b). Shannon diversity was also higher in the 237 Chihuahuan Desert shrubland relative to Plains grassland in April, July, August, and October (Fig. 4b).

The largest difference occurred in October, in which Shannon diversity was 31% higher in the Chihuahuan Desert shrubland than Plains grassland. However, this trend was reversed in both March and September, when Shannon diversity was 19% and 16% higher, respectively, in Plains grassland than in shrubland (Fig. 4b). The two grassland ecosystems differed in Shannon diversity in March, May, June, and September, with greater Shannon diversity in the Plains relative to Chihuahuan Desert grassland in all of these months (Fig. 4b).

244 Dryland ecosystems diverged in the magnitude of seasonal variation in bee assemblage composition,

245 abundance, and diversity

246 Assemblage composition. Bee assemblage composition varied strongly among months, with the 247 magnitude of seasonal change differing among ecosystems (Figs. 3,5; Table 2: Ecosystem X Month, P = 248 0.0001). The Chihuahuan Desert grassland had the greatest seasonal turnover in bee species 249 composition (Fig. 3b), and the highest rate of compositional change from month to month (Fig. 6). In 250 contrast, the Chihuahuan Desert shrubland had the lowest seasonal composition change (Figs. 3c,6), 251 with low turnover between July and August, and between August and September, compared to the other 252 ecosystems (Fig. 5). Among months, in all ecosystem types, bee species composition differed most 253 strongly between March and either September (Plains grassland: mean similarity = 16.4, P = 0.0077) or 254 October (Desert grassland: mean similarity = 9.2, P = 0.0091; shrubland: mean similarity = 11.7, P =255 0.0070) (Fig. 3). In contrast, in all ecosystems, June and July were most compositionally similar to one 256 another, with low turnover between them (Figs. 3.5; Plains grassland: mean similarity = 76.2, P = 0.0091; 257 Desert grassland: mean = 72.42. P = 0.0077; Desert shrubland mean = 70.4. P = 0.0156). Seasonal 258 patterns in bee assemblage composition were largely driven by common rather than rare species, as 259 indicated by very few qualitative differences in analysis outcomes when excluding singletons or 260 moderately rare species (see Supplementary Fig. S4). 261 Abundance. Like species composition, total bee abundance also varied seasonally across the

three ecosystem types (Table 4), and ecosystem types exhibited differing trends in total abundance over the course of the season (Fig. 4a). In the Chihuahuan Desert grassland, bee abundance increased from March to April, then generally declined through the rest of the season (Fig. 4a). In contrast, the Plains grassland had similar levels of bee abundance in March and April (df = 84, t = -1.13, P = 0.95), followed

by a ~50% decrease in abundance between April and May (df = 84, t = 11.12, P < 0.0001) and a 66% increase in abundance between May and June (df = 84, t = -6.99, P < 0.0001). Between July and August, while bee abundance decreased ~30% within both the Chihuahuan Desert grassland (df = 84, t = 4.62, P= 0.0004) and Plains grassland ecosystems (df = 84, t = 5.93, P < 0.0001), it increased by 40% within the Chihuahuan Desert shrubland (df = 84, t = -3.86, P = 0.0053) (Fig. 4a). Across ecosystem types, bee abundances were generally lower in September and October relative to all other months (Fig. 4a).

272 Diversity. Within each ecosystem, most months had similar levels of species richness, with some 273 exceptions (Fig. 4c). Notably, there was a sharp decline in richness between August and October across 274 all three ecosystems (Fig. 4c). During this period, richness declined by 70% within the Chihuahuan Desert 275 grassland (df = 84, t = 11.18, P < 0.0001) and by 60% within both the Plains grassland (df = 84, t = 10.92, 276 P < 0.0001) and Desert shrubland (df = 84, t = 11.18, P < 0.0001). However, month-to-month trends in 277 Shannon diversity and evenness diverged among ecosystems (Fig. 4b,d). Patterns in total abundance, 278 Shannon diversity, richness, and evenness were all largely driven by common rather than rare species 279 (see Supplementary Fig. S5).

Indicators of temporal variation within ecosystem types. Certain bee taxa were indicators of
specific months across all three ecosystems according to their DL indicator values (Supplementary
Tables S3-S6). These included Osmia species, Eucera lycii, Anthophora porterae, and Melecta pacifica
(March), Dioxys and Anthophora species (April), Diadasia australis and rinconis (June), Martinapis
lutericornis, Halictus ligatus, and Melissodes tristis (July), and Perdita semicaerulea and marcialis
(August) (Supplementary Tables S3-S6). September and October lacked indicator species shared by all
ecosystems.

In contrast, certain bee taxa were only characteristic of a given month within one or two
ecosystems (Supplementary Tables S3-S6). For instance, in June, the shrubland site had 5 indicator
species in the genus *Lasioglossum*; one of these was also characteristic of the desert grassland site
(Supplementary Tables S3,S5,S6). In July, *Perdita* species (Andrenidae) were indicators of the
Chihuahuan Desert sites but not the Plains site, the grassland sites had indicator species in the Halictidae
(especially *Lasioglossum*), and differing members of the Apidae were characteristic of different
ecosystems (Supplementary Tables S3-S6). In September, *Perdita* species (Andrenidae) were

characteristic of the Plains grassland, *Macrotera* (Andrenidae) were characteristic of the desert sites, and
differing members of the Apidae were characteristic of each site (Supplementary Tables S3-S6). Finally,
25 species were indicators of a particular month in one or two ecosystem types, and were then indicators
of a different month, often the following one, in the other ecosystem(s) (Supplementary Table S3).

298

299 Discussion

We found large variation in bee assemblages and their seasonality among three dryland ecosystem types of the southwestern U.S. These results indicate the potential for future ecosystem state transitions to alter bee assemblage composition in drylands. Overall, ecosystem types in our study had similar levels of bee species richness but differed from one another in species evenness and composition. These results imply that state transitions could alter the presence/absence and relative abundances of bee species in our system, bringing about substantial assemblage reordering.

306 Our data suggest that the most probable state transitions in our southwestern U.S. drylands – 307 shrub encroachment into Desert grassland and Desert grassland encroachment into Plains grassland ^{59,60} 308 - could substantially reshape bee communities. For the grass-to-shrub transition, our results suggest that, 309 on average over the season, total bee abundance will decrease while richness, Shannon diversity, and 310 Pielou's evenness will increase. In contrast, our findings predict that richness, Shannon diversity, and 311 evenness will decrease while total abundance will remain relatively unchanged if Desert grassland 312 replaces Plains grassland. The simultaneous occurrence of these state transitions could therefore 313 substantially alter the distribution of bees and their ecosystem services across the landscape.

314 A number of factors complicate accurately predicting the outcomes of ecosystem state transitions 315 for bees. First, bees may alter their foraging patterns based on floral resource availability⁶¹, and could 316 respond to shifting vegetation composition by foraging for greater distances if floral resources are scarce 317 in a particular location⁶². However, foraging range can differ greatly among bee species based on body 318 size⁶³, and the energetic costs of longer foraging distances may be high⁶⁴. These factors could mediate 319 the consequences of ecosystem state transitions for bee assemblage composition in ways that merit 320 further research, as they have been little-examined⁶¹. Our finding that particular bee species were 321 indicators of different months in different ecosystem types suggests that bees in our system may shift

322 their foraging locations based on floral availability (though they could alternatively be emerging at different 323 times of year in different ecosystems; see subsequent paragraphs) highlighting the importance of 324 examining foraging dynamics in the future.

325 Second, state transitions may create positive feedbacks that accelerate their pace and influence 326 bee responses¹. Creosote bush expansion, which is limited by minimum nighttime temperature, is aided 327 by a feedback in which a creosote individual creates a warmer microclimate around itself, which can 328 buffer it from low temperatures, in turn creating conditions favorable to further creosote establishment^{47,65}. 329 This accelerated temperature increase could influence the relative dominances of bee species, which 330 may differ in their temperature responses. For instance, in one Osmia species, increased temperature 331 during larval development caused decreased prepupal weight and increased adult mortality⁶⁶, and in 332 another species it increased the frequency of 1-year rather than 2-year lifecycles³⁹. If accelerated bee 333 assemblage shifts result in pollen limitation for plants already threatened by shrub encroachment, the 334 feedback could be enhanced, further increasing the pace of encroachment.

335 Our results contribute to a number of global studies suggesting how state transitions may 336 influence drylands arthropod communities. For instance, in the Chihuahuan Desert, ant species 337 composition varied with mesquite (Prosopis glandulosa) encroachment level, but richness and abundance 338 did not⁶⁷. At the Sevilleta NWR, grasshopper assemblage similarity decreased with elevational variation in 339 shrub and Bouteloua sp. grass cover⁶⁸. Similarly, at our study sites, the Desert grassland and shrubland 340 had distinct ground-dwelling arthropod assemblages, with higher abundance but similar richness in the 341 grassland relative to the shrubland, as we found¹⁴. These findings together suggest that Chihuahuan 342 Desert state transitions may similarly influence abundance and diversity patterns in several arthropod 343 groups. However, global evidence suggests that shrub encroachment can differentially affect arthropod 344 taxa³. For instance, shrub-encroached pastures in Spain had higher pollinator richness but fewer 345 pollinator visits to forbs relative to shrub-absent sites³², contrasting with our finding of little difference in 346 bee richness between shrub- and grass-dominated sites. In the Kalahari Desert, shrub encroachment 347 corresponded with greater abundances of some ground-dwelling arthropod groups but declines in 348 others⁶⁹. Our results thus add to understanding of how widely-occurring ecosystem transitions may affect 349 arthropods differentially across space.

350 Our work also bolsters evidence from human-altered landscapes about how bee assemblages 351 vary at a landscape scale. For instance, as in our study, bee abundance but not richness changed with 352 land use intensity in tropical agroecosystems for solitary bees, which comprised the majority of our 353 dataset²⁹. In contrast, a different study found shifts in bee species composition but not abundance or 354 diversity among forest fragments³⁰. Other studies have documented strong differences in both abundance 355 and richness of bees among habitat types^{33,70,71}. These findings highlight the importance of separately 356 examining trends within particular ecosystem and land-use types to comprehensively predict future 357 patterns.

358 Month-to-month differences in bee species composition were an order of magnitude stronger than 359 ecosystem differences in our study. This finding suggests potential susceptibility of bees in all ecosystem 360 types to climate change-induced phenological shifts³⁶. In particular, climate models for the southwestern 361 U.S. predict less precipitation in July and August, and more in September and October, resulting in an 362 extended period of aridity between spring rains and the start of the summer monsoon⁷². Evidence 363 suggests that desert bees, most of which nest underground, frequently cue on precipitation for their 364 emergence^{37,38}. Under altered monsoon precipitation timing, bees that currently emerge in July or August 365 might shift emergence to September and October, leading to higher levels of bee abundance and 366 richness at all sites during these months. These differences could be particularly pronounced in the 367 Chihuahuan Desert shrubland ecosystem, for which bee abundance, Shannon H', richness, and Pielou's 368 J (evenness) were all highest in July or August. The Chihuahuan Desert grassland could also be 369 particularly susceptible to altered dynamics, given that it had the strongest seasonal turnover. Substantial 370 assemblage reordering among months could occur if different bee species shift their phenological timing 371 to different degrees, which could have landscape-level consequences given that the Chihuahuan Desert 372 ecosystem types are expected to expand in the future⁴⁵⁻⁴⁷. In addition, for social bees that are active 373 throughout the growing season, such as those in the family Halictidae⁷³, loss of floral resources due to 374 midsummer aridity could cause abundance declines or colony death. Predicting the consequences of 375 ecosystem state transitions will thus require considering bee assemblage seasonality in ecosystem types 376 that are expanding versus contracting.

377 Regional climate predictions for the southwestern U.S. are dire – the probability of decadal 378 droughts is nearly 100% by the end of the century⁷⁴. Such droughts could differentially affect bees with 379 differing phenologies and life history strategies, and could lead to bee assemblage reordering. For 380 instance, many desert bees can remain in diapause for one year or more, emerging when conditions are 381 favorable^{37,38}. In one of the few studies on the topic, fewer bees emerged during a drought year compared 382 to the previous and following years in the northwestern Chihuahuan Desert³⁸. A greater proportion of 383 specialist than generalist bees remained in diapause, and the specialists that emerged were those whose 384 host plants bloom under low precipitation conditions. For Larrea tridentata, which requires precipitation to 385 bloom⁷⁵, few specialist bees emerged during the drought, suggesting that these specialists time their 386 emergence with their host plant³⁸. Differences among ecosystem types in their dominant flowering plant 387 species, their associated specialist versus generalist bee species, and their seasonality could therefore 388 lead to strong bee assemblage divergence among them as dominant bee species in each ecosystem 389 respond differentially to increased drought and shifted precipitation timing, with landscape-level bee 390 assemblage changes occurring as some ecosystems expand and others contract. Future analyses will 391 explore connections between bee abundance, diversity, and composition and individual aspects of 392 climate change over our time series.

Our findings of strong bee assemblage seasonality are consistent with work indicating high temporal turnover in plant-pollinator interactions in subalpine and alpine communities over the course of the growing season^{41,76}. Seasonal variation in plant-pollinator networks has also been documented in agricultural landscapes⁴⁴. While our study was not designed to examine plant-pollinator interactions, our results set the stage for considering how plant-pollinator networks could be altered by ecosystem state transitions and climate-induced phenological shifts of bee species.

In our dataset, common rather than rare bee species drove the trends in abundance, diversity, and composition that we observed over both space and time. Considering these species' ecologies may thus be particularly important for predicting the consequences of ecosystem state transitions, and some species may portend change in the bee assemblage as a whole⁷⁷. Among the most abundant bees in our dataset, three species (*Agapostemon angelicus, Lasioglossum semicaeruleum*, and *Melissodes tristis*) were broad generalists that collect pollen from plants of many families^{38,78}, suggesting that plants visited

by these and other bees could be buffered to a certain extent if there are future bee declines. However, specialist bees were also among the most abundant: *Diadasia rinconis* is a specialist on Cactaceae⁷⁹, *Anthophora affabilis* is a generalist with a strong preference for *Astragalus*, and *Perdita larreae* is a narrow specialist on *Larrea tridentata*^{23,80}. The consequences of ecosystem state transitions for these bee species may thus depend on shifts in their host plants. For instance, expansion of *Larrea tridentata* could benefit populations of *P. larreae* and other creosote bush specialist bees, and possibly lead to stronger competitive dynamics among creosote specialists and generalists under future conditions.

412 Our study identified bee species as indicators of each ecosystem type; monitoring these species 413 could help to illuminate the community-level consequences of ecosystem state transitions. The 414 Chihuahuan Desert shrubland had more indicator species than the other two ecosystem types, 415 suggesting that its future expansion could bring about distinctive assemblage shifts. The strongest 416 indicators of the shrubland included Perdita larreae, which specializes on Larrea tridentata, and P. 417 diversa, which specializes on *Tiquilia* spp., plants only found at the shrubland site. Abundance increases 418 of these bee species could thus signal effects of shrubland expansion on pollinator communities. 419 Similarly, in the Plains grassland, one indicator species (Colletes scopiventer) specializes on 420 Chamaesaracha spp., which are present in all three ecosystems but are most abundant in the Plains 421 grassland. Future decreases in the abundance of C. scopiventer could signal community-level shifts 422 accompanying the declining dominance of the Plains grassland. However, the remaining indicator species 423 of the two grassland ecosystems were broad generalists. Factors other than plant community 424 composition, such as nesting habitat preferences, may thus underlie their restriction to particular sites, 425 and they may be relatively less susceptible to climate-induced plant community shifts. This could also be 426 the case for Macrotera portalis, an indicator of the shrubland but a specialist on Sphaeralcea spp.⁸¹, 427 which is common in all three ecosystem types, and for generalist bee species that were indicators of the 428 shrubland. The spatial distribution of suitable nesting habitat also merits future consideration in that 429 specialist bees could be negatively affected if their plant hosts shift their ranges away from potential 430 nesting sites.

In addition, cleptoparasitic bees were among our identified indicator species. These included one
 indicator of the Chihuahuan Desert shrubland (*Neolarra vigilans*) and three of the Plains grassland

433 (Melecta alexanderi and two Sphecodes species). Not surprisingly, in the cases of Neolarra and Melecta. 434 their bee hosts, *Perdita* and *Anthophora*, were also among indicators of the same ecosystem types. 435 Cleptoparasitic bees may be particularly good indicators of environmental change, as they are relatively 436 diverse, and are known to be among the first bee functional groups to respond to disturbance⁸². 437 Monitoring the abundances of these species could indicate shifts in bee assemblage dynamics as 438 ecosystem state transitions occur. Cleptoparasitc species were also amongst indicators of particular 439 months, frequently in tandem with possible hosts, and may thus be useful for tracking phenological 440 responses to environmental change⁸². For example, both *Melecta alexanderi* and *M. bohartorum* were 441 indicators of the shrubland site in March, but were indicators of the grassland sites in April, suggesting the 442 possibility of altered emergence timing under the warmer microclimate conditions of the shrubland^{47,65}, 443 and thus susceptibility to phenological shifts in response to increasing temperature. 444 We also identified bee species that were characteristic of particular times of year across 445 ecosystem types. Monitoring these species could enable the detection of broad, cross-site phenological 446 shifts that may occur in the future. For instance, Osmia, Anthophora, and Diadasia species may be 447 monitored to consider shifting pre-monsoon bee phenology, and Perdita species may be used to study 448 shifts in emergence timed with monsoon rains. Future publications using these data will investigate inter-449 annual bee assemblage differences and relationships with climate variables. 450 Finally, we identified bee species that were characteristic of particular months only in specific 451 ecosystems. Among these, certain specialist bee species may be candidates for detecting important 452 phenological shifts within ecosystems, identifying phenological differences among ecosystems, and 453 tracking how specialists versus generalists respond to climate shifts. For instance, March in the 454 Chihuahuan Desert ecosystems had indicator species that likely specialize on Fabaceae (Ashmeadiella 455 erema and A. rubrella), and April in the grassland sites had a specialist on Brassicaceae (Dufourea 456 pulchricornis). In the Chihuahuan Desert ecosystem types, specialists of creosote bush were 457 characteristic of May, corresponding with creosote's spring bloom⁷⁵, June had Cactaceae specialists 458 (Diadasia sp.) timed with that family's bloom⁷⁹, and July had specialists on Asteraceae (Perdita ignota 459 ignota, P. callicerata, P. fallax, and P. albovittata) and Tiquilia sp. (P. diversa). In contrast, specialists on 460 Asteraceae (Melissodes coreopsis, P. ignota ignota, and P. callicerata) were indicators of August in the

461 Plains grassland, suggesting a shift in the importance of particular floral resources and/or differing 462 phenological patterns among ecosystem types. In August, the Chihuahuan Desert sites were 463 characterized by numerous creosote specialists, as documented in several studies^{23,38}, including 464 Hesperapis larreae, P. semicaerulea, and P. larreae. These species may be candidates for examining 465 how delayed monsoon influences bee phenology. The Sphaeralcea specialist Macrotera portalis⁸¹ was an 466 August indicator in the shrubland, but other Sphaeralcea specialists were characteristic of September in 467 both Chihuahuan Desert ecotypes; perhaps competitive dynamics were responsible for this difference. 468 These examples illustrate the suite of factors that could be important to consider to predict bee presence 469 and seasonality across the landscape. 470 Our analysis of 13 years of bee assemblage data spanning 302 species suggests that future 471 dryland ecosystem state transitions, by themselves, may alter bee species' relative abundances and 472 presence/absence. Strong bee assemblage seasonal turnover, particularly in ecosystems predicted to 473 expand, indicates the potential for bee phenological shifts to accompany state transitions, potentially 474 reordering communities. Our results indicate that predicting the consequences of global change for bee 475 assemblages will require accounting for both within-year and among-ecosystem variation. 476 477 Data availability 478 The data generated and analyzed during the current study will be available upon publication in 479 the Environmental Data Initiative (EDI) Data Portal (https://portal.edirepository.org). 480 481 References

- D'Odorico, P., Bhattachan, A., Davis, K. F., Ravi, S. & Runyan, C. W. Global desertification: drivers
 and feedbacks. *Adv. Water Resour.* **51**, 326–344 (2013).
- 484 2. Bestelmeyer, B. T. *et al.* Desertification, land use, and the transformation of global drylands. *Front.*485 *Ecol. Environ.* 13, 28–36 (2015).
- 486 3. Eldridge, D. J. *et al.* Impacts of shrub encroachment on ecosystem structure and functioning: towards
 487 a global synthesis. *Ecol. Lett.* 14, 709–722 (2011).

- 488 4. Allen, C. D., Breshears, D. D. & McDowell, N. G. On underestimation of global vulnerability to tree
- 489 mortality and forest die-off from hotter drought in the Anthropocene. *Ecosphere* **6**, art129 (2015).
- 490 5. Sala, O. E. & Maestre, F. T. Grass-woodland transitions: determinants and consequences for
- 491 ecosystem functioning and provisioning of services. J. Ecol. **102**, 1357–1362 (2014).
- 492 6. Biederman, J. A. *et al.* Terrestrial carbon balance in a drier world: the effects of water availability in
- 493 southwestern North America. *Glob. Change Biol.* 22, 1867–1879 (2016).
- 494 7. Bestelmeyer, B. T. *et al.* The grassland-shrubland regime shift in the southwestern United States:
 495 misconceptions and their implications for management. *Bioscience* 68, 678–690 (2018).
- 496 8. Anderson-Teixeira, K. J., Delong, J. P., Fox, A. M., Brese, D. A. & Litvak, M. E. Differential responses
- 497 of production and respiration to temperature and moisture drive the carbon balance across a climatic
- 498 gradient in New Mexico. *Glob. Change Biol.* **17**, 410–424 (2011).
- 9. Petrie, M. D., Collins, S. L., Swann, A. M., Ford, P. L. & Litvak, M. E. Grassland to shrubland state
 transitions enhance carbon sequestration in the northern Chihuahuan Desert. *Glob. Change Biol.* 21,
 1226–1235 (2015).
- 50210. Turnbull, L., Wainwright, J. & Brazier, R. E. Nitrogen and phosphorus dynamics during runoff events503over a transition from grassland to shrubland in the south-western United States. *Hydrol. Process.*
- 504 **25**, 1–17 (2011).
- 505 11. Wang, G. *et al.* Post-fire redistribution of soil carbon and nitrogen at a grassland–shrubland ecotone.
 506 *Ecosystems* 22, 174–188 (2019).
- 507 12. Ratajczak, Z. *et al.* Changes in spatial variance during a grassland to shrubland state transition. *J.*508 *Ecol.* **105**, 750–760 (2017).
- 509 13. Sanchez, B. C. & Parmenter, R. R. Patterns of shrub-dwelling arthropod diversity across a desert
 510 shrubland–grassland ecotone: a test of island biogeographic theory. *J. Arid Environ.* 50, 247–265
 511 (2002).
- 512 14. Lightfoot, D. C., Brantley, S. L. & Allen, C. D. Geographic patterns of ground-dwelling arthropods
 513 across an ecoregional transition in the North American Southwest. *West. North Am. Nat.* 68, 83–102
 514 (2008).

515 15. Pravalie, R. Drylands extent and environmental issues: a global approach. *Earth-Sci. Rev.* **161**, 259–

516 278 (2016).

- 517 16. UN EMG. Global drylands: a UN system-wide response. Environ. Manag. Group U. N. Geneva
- 518 http://www.unccd.int/Lists/SiteDocumentLibrary/Publications/Global_Drylands_Full_Report.pdf

519 (2011).

- 520 17. Kovács-Hostyánszki, A. et al. Earthworms, spiders and bees as indicators of habitat quality and
- 521 management in a low-input farming region—a whole farm approach. *Ecol. Indic.* **33**, 111–120 (2013).
- 522 18. Gonçalves, R. B., Sydney, N. V., Oliveira, P. S. & Artmann, N. O. Bee and wasp responses to a
- 523 fragmented landscape in southern Brazil. J. Insect Conserv. 18, 1193–1201 (2014).
- 524 19. Klein, A.-M. *et al.* Importance of pollinators in changing landscapes for world crops. *Proc. R. Soc. B*525 *Biol. Sci.* 274, 303–313 (2007).
- 526 20. Ollerton, J., Winfree, R. & Tarrant, S. How many flowering plants are pollinated by animals? *Oikos*527 **120**, 321–326 (2011).
- 528 21. Michener, C. D. *The Bees of the World.* (Johns Hopkins University Press, 2007).
- 529 22. Minckley, R. L. & Ascher, J. S. Preliminary survey of bee (Hymenoptera: Anthophila) richness in the

530 northwestern Chihuahuan Desert in *Merging Science and Management in a Rapidly Changing World:*

- 531 Biodiversity and Management of the Madrean Archipelago III and 7th Conference on Research and
- 532 Resource Management in the Southwestern Deserts, Proceedings RMRS-P-67 (eds. Gottfried, G. J.,
- 533 Ffolliott, P. F., Gebow, B. S., Eskew, L. G. & Collins, L. C.) 138–143 (U.S. Department of Agriculture,
- 534 Forest Service, Rocky Mountain Research Station, 2013).
- 535 23. Minckley, R. L., Cane, J. H. & Kervin, L. Origins and ecological consequences of pollen specialization
 536 among desert bees. *Proc. R. Soc. Lond. B Biol. Sci.* 267, 265–271 (2000).
- 537 24. Turner, R. M., Bowers, J. E. & Brugess, T. L. Sonoran Desert Plants: An Ecological Atlas. (The
 538 University of Arizona Press, 2005).
- 539 25. Minckley, R. L., Cane, J. H., Kervin, L. & Roulston, T. H. Spatial predictability and resource
- 540 specialization of bees (Hymenoptera: Apoidea) at a superabundant, widespread resource. *Biol. J.*

541 *Linn.* Soc. **67**, 119–147 (1999).

542 26. Simpson, B. & Neff, J. Pollination ecology in the Southwest. *Aliso* **11**, 417–440 (1987).

- 543 27. Memmott, J., Craze, P. G., Waser, N. M. & Price, M. V. Global warming and the disruption of plant-
- 544 pollinator interactions. *Ecol. Lett.* **10**, 710–717 (2007).
- 545 28. Kaiser-Bunbury, C. N., Muff, S., Memmott, J., Müller, C. B. & Caflisch, A. The robustness of
- 546 pollination networks to the loss of species and interactions: a quantitative approach incorporating
- 547 pollinator behaviour. *Ecol. Lett.* **13**, 442–452 (2010).
- 548 29. Klein, A.-M., Steffan-Dewenter, I., Buchori, D. & Tscharntke, T. Effects of land-use intensity in tropical
- agroforestry systems on coffee flower-visiting and trap-nesting bees and wasps. *Conserv. Biol.* 16,
 1003–1014 (2002).
- 30. Brosi, B. J., Daily, G. C., Shih, T. M., Oviedo, F. & Durán, G. The effects of forest fragmentation on
 bee communities in tropical countryside. *J. Appl. Ecol.* 45, 773–783 (2008).
- 553 31. Banaszak-Cibicka, W. & Żmihorski, M. Wild bees along an urban gradient: winners and losers. J.
- 554 Insect Conserv. **16**, 331–343 (2012).
- 32. Lara-Romero, C., García, C., Morente-López, J. & Iriondo, J. M. Direct and indirect effects of shrub
 encroachment on alpine grasslands mediated by plant-flower visitor interactions. *Funct. Ecol.* 30,
 1521–1530 (2016).
- 33. Minckley, R. Faunal composition and species richness differences of bees (Hymenoptera: Apiformes)
 from two north American regions. *Apidologie* **39**, 176–188 (2008).
- 560 34. Torné-Noguera, A. et al. Determinants of spatial distribution in a bee community: nesting resources,
- flower resources, and body size. *PLoS ONE* **9**, e97255 (2014).
- 35. Ruttan, A., Filazzola, A. & Lortie, C. J. Shrub-annual facilitation complexes mediate insect community
 structure in arid environments. *J. Arid Environ.* **134**, 1–9 (2016).
- 36. Renner, S. S. & Zohner, C. M. Climate change and phenological mismatch in trophic interactions
 among plants, insects, and vertebrates. *Annu. Rev. Ecol. Evol. Syst.* 49, 165–182 (2018).
- 566 37. Danforth, B. N. Emergence dynamics and bet hedging in a desert bee, *Perdita portalis. Proc. R. Soc.*
- 567 Lond. B Biol. Sci. 266, 1985–1994 (1999).
- 568 38. Minckley, R. L., Roulston, T. H. & Williams, N. M. Resource assurance predicts specialist and
- 569 generalist bee activity in drought. Proc. R. Soc. B Biol. Sci. 280, 20122703 (2013).

- 570 39. Forrest, J. R. K., Cross, R. & CaraDonna, P. J. Two-year bee, or not two-year bee? How voltinism Is
- affected by temperature and season length in a high-elevation solitary bee. *Am. Nat.* **193**, 560–574
 (2019).
- 573 40. Tylianakis, J. M., Klein, A.-M. & Tscharntke, T. Spatiotemporal variation in the diversity of
- 574 Hymenoptera across a tropical habitat gradient. *Ecology* **86**, 3296–3302 (2005).
- 575 41. Simanonok, M. P. & Burkle, L. A. Partitioning interaction turnover among alpine pollination networks:
- 576 spatial, temporal, and environmental patterns. *Ecosphere* **5**, art149 (2014).
- 42. Rollin, O., Bretagnolle, V., Fortel, L., Guilbaud, L. & Henry, M. Habitat, spatial and temporal drivers of
 diversity patterns in a wild bee assemblage. *Biodivers. Conserv.* 24, 1195–1214 (2015).
- 43. Leong, M., Ponisio, L. C., Kremen, C., Thorp, R. W. & Roderick, G. K. Temporal dynamics influenced
- 580 by global change: bee community phenology in urban, agricultural, and natural landscapes. *Glob*.
- 581 Change Biol. 22, 1046–1053 (2016).
- 44. Tucker, E. M. & Rehan, S. M. Farming for bees: annual variation in pollinator populations across
 agricultural landscapes. *Agric. For. Entomol.* 20, 541–548 (2018).
- 45. Peters, D. P. C. & Yao, J. Long-term experimental loss of foundation species: consequences for
 dynamics at ecotones across heterogeneous landscapes. *Ecosphere* 3, art27 (2012).
- 586 46. Collins, S. L. & Xia, Y. Long-term dynamics and hotspots of change in a desert grassland plant
- 587 community. Am. Nat. 185, E30–E43 (2015).
- 588 47. He, Y., D'Odorico, P. & De Wekker, S. F. J. The role of vegetation-microclimate feedback in
- promoting shrub encroachment in the northern Chihuahuan desert. *Glob. Change Biol.* 21, 2141–
 2154 (2015).
- 48. Wilson, J. S., Griswold, T. & Messinger, O. J. Sampling bee communities (Hymenoptera: Apiformes)
 in a desert landscape: are pan traps sufficient? *J. Kans. Entomol. Soc.* **81**, 288–300 (2008).
- 49. Cane, J. H., Minckley, R. L. & Kervin, L. J. Sampling bees (Hymenoptera: Apiformes) for pollinator
 community studies: pitfalls of pan-trapping. *J. Kans. Entomol. Soc.* **73**, 225–231 (2000).
- 595 50. Baum, K. A. & Wallen, K. E. Potential bias in pan trapping as a function of floral abundance. *J. Kans.*596 *Entomol. Soc.* 84, 155–159 (2011).

- 597 51. Wickham, H. Reshaping data with the reshape package. J. Stat. Softw. 21, 10.18637/jss.v021.i12
- 598 (2007).
- 599 52. R Core Team. R: A Language and Environment for Statistical Computing. (R Foundation for

600 Statistical Computing, 2017).

- 53. Clarke, K. R. & Gorley, R. N. Primer Version 6.1.10 User Manual and Tutorial. (Primer-E, 2009).
- 602 54. Hallett, L. *et al.* codyn: community dynamics metrics, R package version 2.0.2. 10.5063/F1N877Z6
- 603 (2019).
- 55. Roberts, D. W. labdsv: ordination and multivariate analysis for ecology. https://CRAN.R-
- 605 project.org/package=labdsv (2016).
- 606 56. Oksanen, J. et al. vegan: community ecology package, version 2.2-1. http://CRAN.R-
- 607 project.org/package=vegan (2015).
- 57. Bates, D., Mächler, M., Bolker, B. & Walker, S. Fitting linear mixed-effects models using lme4. *J. Stat.*Softw. 67, 1–48 (2015).
- 58. Lenth, R. V., Singmann, H., Love, J., Buerkner, P. & Herve, M. emmeans: estimated marginal means,
 aka least-squares means. https://cran.r-project.org/web/packages/emmeans/index.html (2018).
- 59. Collins, S. L. *et al.* A multiscale, hierarchical model of pulse dynamics in arid-land ecosystems. *Annu.*
- 613 Rev. Ecol. Evol. Syst. 45, 397–419 (2014).
- 614 60. Rudgers, J. A. *et al.* Climate sensitivity functions and net primary production: a framework for
- 615 incorporating climate mean and variability. *Ecology* **99**, 576–582 (2018).
- 616 61. Ogilvie, J. E. & Forrest, J. R. Interactions between bee foraging and floral resource phenology shape
- 617 bee populations and communities. *Curr. Opin. Insect Sci.* **21**, 75–82 (2017).
- 618 62. Redhead, J. W. *et al.* Effects of habitat composition and landscape structure on worker foraging
 619 distances of five bumble bee species. *Ecol. Appl.* 26, 726–739 (2016).
- 620 63. Greenleaf, S. S., Williams, N. M., Winfree, R. & Kremen, C. Bee foraging ranges and their
- 621 relationship to body size. *Oecologia* **153**, 589–596 (2007).
- 622 64. Horn, J., Becher, M. A., Kennedy, P. J., Osborne, J. L. & Grimm, V. Multiple stressors: using the
- 623 honeybee model BEEHAVE to explore how spatial and temporal forage stress affects colony
- 624 resilience. *Oikos* **125**, 1001–1016 (2016).

- 625 65. He, Y., D'Odorico, P., De Wekker, S. F. J., Fuentes, J. D. & Litvak, M. On the impact of shrub
- 626 encroachment on microclimate conditions in the northern Chihuahuan desert. J. Geophys. Res.–

627 Atmospheres **115**, D21120 (2010).

- 628 66. Radmacher, S. & Strohm, E. Effects of constant and fluctuating temperatures on the development of 629 the solitary bee *Osmia bicornis* (Hymenoptera: Megachilidae). *Apidologie* **42**, 711–720 (2011).
- 630 67. Bestelmeyer, B. T. Does desertification diminish biodiversity? Enhancement of ant diversity by shrub
- 631 invasion in south-western USA. *Divers. Distrib.* **11**, 45–55 (2005).
- 632 68. Rominger, A. J., Miller, T. E. X. & Collins, S. L. Relative contributions of neutral and niche-based
- processes to the structure of a desert grassland grasshopper community. *Oecologia* 161, 791–800
 (2009).
- 635 69. Blaum, N., Seymour, C., Rossmanith, E., Schwager, M. & Jeltsch, F. Changes in arthropod diversity
- along a land use driven gradient of shrub cover in savanna rangelands: identification of suitable
 indicators. *Biodivers. Conserv.* 18, 1187–1199 (2009).
- 638 70. Bates, A. J. *et al.* Changing bee and hoverfly pollinator assemblages along an urban-rural gradient.
 639 *PLoS ONE* 6, e23459 (2011).
- 640 71. Tonietto, R., Fant, J., Ascher, J., Ellis, K. & Larkin, D. A comparison of bee communities of Chicago
 641 green roofs, parks and prairies. *Landsc. Urban Plan.* **103**, 102–108 (2011).
- 642 72. Cook, B. I. & Seager, R. The response of the North American Monsoon to increased greenhouse gas
 643 forcing. *J. Geophys. Res.–Atmos.* **118**, 1690–1699 (2013).
- 644 73. Michener, C. D. *The Social Behavior of the Bees: A Comparative Study*. (Belknap Press of Harvard
 645 University Press, 1974).
- 646 74. Cook, B. I., Ault, T. R. & Smerdon, J. E. Unprecedented 21st century drought risk in the American
 647 Southwest and Central Plains. *Sci. Adv.* 1, e1400082 (2015).
- 648 75. Bowers, J. E. & Dimmitt, M. A. Flowering phenology of six woody plants in the Northern Sonoran
- 649 Desert. Bull. Torrey Bot. Club **121**, 215–229 (1994).
- 650 76. CaraDonna, P. J. *et al.* Interaction rewiring and the rapid turnover of plant–pollinator networks. *Ecol.*
- 651 *Lett.* **20**, 385–394 (2017).

- 652 77. Ellison, A. M. et al. Loss of foundation species: consequences for the structure and dynamics of
- 653 forested ecosystems. *Front. Ecol. Environ.* **3**, 479–486 (2005).
- 654 78. Schuh, R. T., Hewson-Smith, S. & Ascher, J. S. Specimen databases: a case study in entomology
 655 using web-based software. *Am. Entomol.* 56, 206–216 (2010).
- 79. Sipes, S. D. & Tepedino, V. J. Pollen-host specificity and evolutionary patterns of host switching in a
- 657 clade of specialist bees (Apoidea: *Diadasia*). *Biol. J. Linn. Soc.* **86**, 487–505 (2005).
- 658 80. Hurd, P. D. & Linsley, E. G. The principal Larrea bees of the southwestern United States

659 (Hymenoptera, Apoidea). Smithson. Contrib. Zool. 1–74 (1975).

- 660 81. Danforth, B. N., Ji, S. & Ballard, L. J. Gene flow and population structure in an oligolectic desert bee,
- Macrotera (Macroteropsis) portalis (Hymenoptera: Andrenidae). J. Kans. Entomol. Soc. 76, 221–235
 (2003).
- 82. Sheffield, C. S., Pindar, A., Packer, L. & Kevan, P. G. The potential of cleptoparasitic bees as
 indicator taxa for assessing bee communities. *Apidologie* 44, 501–510 (2013).
- 665

666 Acknowledgements

- Funding was provided by the NSF Long Term Ecological Research program (DEB-1655499) and
 the University of New Mexico Department of Biology. We thank David Lightfoot, Olivia Messinger Carril,
 and Jade McLaughlin for their contributions to this work.
- 670

671 Author contributions

672 M.R.K. assisted with bee specimen collection, analyzed the data, and wrote the manuscript.

673 K.W.W. designed the study and completed the majority of specimen collection and identification. J.B.

674 assisted with specimen collection and identification. T.L.G. identified specimens and provided taxonomic

675 expertise. J.A.R. and K.D.W. contributed to manuscript conceptual framing, data analysis, and writing. All

authors contributed to revising the manuscript.

677

678 **Competing interests**

679

The authors declare no competing interests.

680

681 Figure legends

- Figure 1. Non-metric multidimensional scaling (NMDS) plot depicting variation in bee species composition
- among three dryland ecosystem types: Plains grassland (blue points), Chihuahuan Desert grassland
- 684 (black points), and Chihuahuan Desert shrubland (green points). NMDS was run with 500 randomized re-
- starts and 2D stress = 0.13. On average, all ecosystem types significantly differed from one another
- 686 (Table 2): Plains grassland versus Chihuahuan Desert grassland (P = 0.0082), Plains grassland versus
- 687 Chihuahuan Desert shrubland (P = 0.0075), and Chihuahuan Desert grassland versus shrubland (P =

688 0.0084).

689

690 Figure 2. Mean yearly abundance + s.e. (darker, leftmost bar in each pair) and Dufrene-Legendre (DL)

691 indicator species value (lighter, rightmost bar in each pair) for important bee species within each

692 ecosystem type (Plains grassland, blue bars, B; Chihuahuan Desert grassland, black bars, G;

693 Chihuahuan Desert shrubland, green bars, C). Included bee species were within the 20 most abundant

694 species found across the study, and/or were top indicator species of particular ecosystem types

695 according to DL indicator value. Plots are arranged from left to right by mean yearly abundance across

696 ecosystem types.

697

698 Figure 3. Non-metric multidimensional scaling (NMDS) plots depicting variation in bee species

699 composition among months for three dryland ecosystems: (a) Plains grassland, (b) Chihuahuan Desert

grassland, and (c) Chihuahuan Desert shrubland. NMDS was run with all samples together, with 500

701 randomized re-starts and 2D stress = 0.13.

702

Figure 4. Variation across sampling months in per-transect bee abundance and diversity (± s.e.) as
measured by a) total bee abundance, b) Shannon diversity index (*H'*), c) richness, and d) evenness
(Pielou's *J*) for three dryland ecosystem types: Plains grassland (blue points), Chihuahuan Desert
grassland (black points), and Chihuahuan Desert shrubland (green points). Letters denote contrasts
between biomes within a given month; biomes labeled with different letters differed significantly from one

- another in the relevant abundance/diversity metric. Points lacking letters did not differ significantly from
 any other biome in the given month. For total abundance, s.e. values were <0.1.
- 710
- 711 Figure 5. Total bee species turnover between pairs of months (indicated on the x-axis) for three dryland
- ecosystem types: Plains grassland (blue points), Chihuahuan Desert grassland (black points), and
- 713 Chihuahuan Desert shrubland (green points).
- 714
- Figure 6. Average rate of bee assemblage change during March through October in three ecosystem
- 716 types: Plains grassland (slope = 9.34, s.e. = 4.99, t = 1.9, P = 0.0725), Chihuahuan Desert grassland
- 717 (slope = 24.32, s.e. = 3.99, *t* = 6.1, *P* < 0.0001), and Chihuahuan Desert shrubland (slope = 7.81, s.e. =
- 718 2.92, t = 2.7, P = 0.0128). Intervals (x-axis) represent time lags between all pairwise combinations of
- 719 months. Distances (y-axis) correspond with differences in bee assemblage composition between pairs of
- months, calculated as Euclidean distances. The slope of each line indicates the rate of bee assemblage
- 721 change in each ecosystem.
- 722

723 724 725 Table 1. Latitude, longitude, and elevation of study sites representing three ecosystem types of the southwestern U.S., along with current versus predicted future foundation species.

Ecosystem type	Latitude	Longitude	Elevation (m)	Current foundation species	Future foundation species
Desert shrubland	34.3431	-106.7417	1615	Creosote bush	Creosote bush
Desert grassland	34.3350	-106.7219	1616	Black grama	Creosote bush
Plains grassland	34.3325	-106.6328	1670	Blue grama	Black grama

727 728 729 730 731 Table 2. Results of 1) perMANOVA with 9999 permutations to test for the influence of ecosystem type

and month of sample collection on bee assemblage composition, using a Bray-Curtis similarity metric,

and 2) permDISP examining differences among ecosystem types and months in bee assemblage

dispersion.

			perMANOVA			permDISP			
	num. df	SS	MS	pseudo-F	Р	denom. df	F	Р	
Ecosystem	2	23252.00	11626.00	34.54	0.0001	117	0.52	0.7074	
Month	7	139860.00	19981.00	94.02	0.0001	112	5.92	0.0002	
Ecosystem x month	14	34139.00	2438.50	11.48	0.0001				
Transect (ecosystem)	12	4039.70	336.64	1.58	0.0003				
Residuals	84	17851.00	212.51						

Table 3. Indicator species for each ecosystem (Plains grassland, Chihuahuan Desert grassland, and

733 734 735 Chihuahuan Desert shrubland) according to Dufrene-Legendre indicator species value. Species are listed from highest to lowest indicator value within each ecosystem.

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Species	Family	Indicator value	P-value
Plains grassland			
Colletes scopiventer	Colletidae	0.64	0.0010
•	Halictidae	0.64	0.0010
Halictus ligatus			
Anthophora montana	Apidae	0.52 0.50	0.0010
Agapostemon angelicus	Halictidae Halictidae	0.50	0.0010
Halictus tripartitus		0.41	0.0010
Anthidium porterae Sphecodes sp. 1	Megachilidae Halictidae	0.27	0.0280
		0.27	0.0010 0.0190
Anthophora urbana	Apidae		0.0190
Melecta alexanderi	Apidae	0.20 0.18	
Sphecodes sp. 5	Halictidae		0.0250
Melissodes thelypodii thelypodii	Apidae	0.17	0.0090
Sphecodes sp. 6	Halictidae	0.15	0.0090
Megachile policaris	Megachilidae	0.14	0.0110
Pseudopanurgus sp. 2	Andrenidae	0.10	0.0210
Chibushuan Desart greesland			
Chihuahuan Desert grassland Lasioglossum (Dialictus) sp. 2	Halictidae	0.66	0.0010
		0.51	0.0010
Lasioglossum semicaeruleum	Halictidae		
Diadasia megamorpha	Apidae Andrenidae	0.20	0.0250
Perdita sphaeralceae alticola		0.18	0.0020
Megachile sublaurita	Megachilidae	0.16	0.0240
Perdita cara	Andrenidae	0.15	0.0160
Atoposmia aff. daleae	Megachilidae	0.10	0.0330
Atoposmia aff. daleae 2	Megachilidae	0.10	0.0420
Chihuahuan Desert shrubland			
Agapostemon melliventris	Halictidae	0.56	0.0010
Perdita larreae	Andrenidae	0.50	0.0010
Neolarra vigilans	Apidae	0.45	0.0010
Perdita marcialis	Andrenidae	0.43	0.0010
Perdita diversa	Andrenidae	0.42	0.0010
	Halictidae	0.40	0.0010
Lasioglossum aff. pervarum	Halictidae	0.40	0.0080
Lasioglossum (Dialictus) sp. 8 Ashmeadiella meliloti		0.39	
	Megachilidae		0.0440
Anthophorula completa	Apidae	0.33	0.0010
Lasioglossum morrilli	Halictidae	0.32	0.0240
Lasioglossum (Dialictus) sp. 7	Halictidae	0.31	0.0050
Ashmeadiella bigeloveae	Megachilidae	0.27	0.0160
Ashmeadiella cactorum	Megachilidae	0.25	0.0060
Macrotera portalis	Andrenidae	0.22	0.0090
Dianthidium implicatum	Megachilidae	0.20	0.0010
Anthophora cinerula	Apidae	0.19	0.0110
Anthidium cockerelli	Megachilidae	0.17	0.0070
Perdita austini	Andrenidae	0.16	0.0040
Apis mellifera	Apidae	0.16	0.0410
Megachile lobatifrons	Megachilidae	0.14	0.0140
Megachile spinotulata	Megachilidae	0.11	0.0370

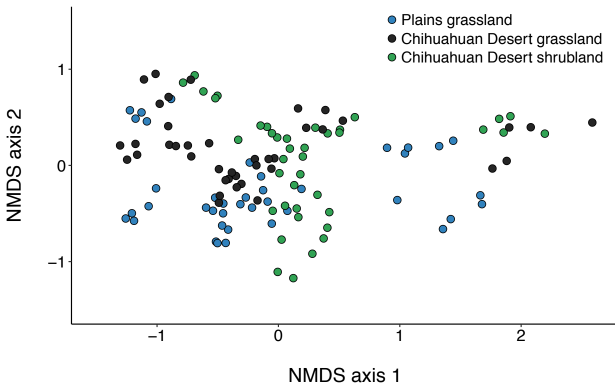
738 739 740 Table 4. Results of linear mixed effects models testing the influences of ecosystem type and month of

sample collection on total bee abundance, as well as bee assemblage Shannon diversity index (H'),

richness, and evenness (Pielou's J).

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	Total abundance		Shanno	Shannon H'		Richness		Evenness	
-	df	X ²	Р						
Ecosystem	2	45.19	< 0.0001	34.72	< 0.0001	1.38	0.50044	47.28	< 0.0001
Month	7	796.09	< 0.0001	105.64	< 0.0001	221.14	< 0.0001	85.33	< 0.0001
Ecosystem x month	14	359.31	< 0.0001	142.78	< 0.0001	28.13	0.01368	320.07	< 0.0001



743 744 Figure 1

