Running head: PLANT-POLLINATOR NETWORK ASSEMBLY

The temporal assembly of plant-pollinator networks following restoration

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1 Abstract

TO BE RE-WRITTEN The structure of networks is related to ability of commu-2 nities to maintain function in the face of species extinction. Understanding network structure and how it relates to network disassembly, therefore, is a priority for systemlevel conservation biology. We explore the assembly of plant-pollinator communities on native plant restorations in the Central Valley of California. The assembling communities are paired with unrestored field margins (controls) and mature (non-assembling) hedgerows. We determine whether there are change points in the assembly of the communities where the network undergoes significant reorganization. We are also ask how are the individual species changing their interaction patterns? What does 10 this mean for the topology/resilience of the network? We also attempted to adapt 11 a financial model to mutualistic networks. Our biggest difficulty with this approach 12 was to translate the price term to mutualistic systems. We explored a range of approaches, such as number of visits a species performs. However, it seems that financial systems 14 cannot be easily translated to mutualistic systems. In addition, we used a Changing 15 Point Detection Algorithm to assess weather the different communities went through 16 a critical reorganization on their interaction patterns. We were able to identify some 17 changing points in the communities, and also to explore some general patterns commonly 18 used to describe ecological networks. For example, on the network level, networks 19 become increasingly modular and less nested, whereas on the species level, species 20 become more specialized, as resources become more reliable. 21

Keywords: changing points, temporal networks, hedgerows, species interactions, prefer-

²³ ential attachment , mutualisms

Introduction

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- Global change has created a severe biodiversity crisis, and as species are lost, so are their
- interactions (Dunn et al., 2009; Barnosky et al., 2011). Because mutualistic interactions are
- essential for maintaining the diversity their component guilds of species , these systems
- are particularly at risk from coextinction cascades. The nature of these coextinction cascades 28
- depends on the interaction patterns within a community (Memmott et al., 2004; Rezende et al., 2007; Bas
- Recovering the lost biodiversity and interactions through ecological restoration has become 30
- increasingly imperative, and a 31
- The structure of networks is related to ability of communities to maintain function 32 in the face of species extinction. 33
- A key restoration aim is to facilitate assembly of robust interaction networks(Menz et al., 2010). We know little networks; thus it is critical to study how restoration influences the 35 assembly of plant-pollinator interactions.
- In general, however, about how to re-assemble interacting communities through 37 restoration, or the process of ecological network assembly more generally. 38
- few mechanisms of network assembly have been developed and analyzed. 39
- The mostly widely explored mechanism of network assembly, preferential attach-40 ment (Barabási & Albert, 1999), predicts that a new species (or node more generally) 41 is more likely to interact with species that are already well-connected ("the rich-get-42 richer" principle, Barabási & Albert, 1999). In pollination systems— a particularly 43 ubiquitous mutualism (Ollerton et al., 2011; Klein et al., 2007) — some studies have 44 found support for this mechanism of assembly. Investigating the 45
 - To date, only a handufullof field studies have examined how networks assemble

over time, often using space for time gradients.

- Olesen *et al.* (2008) was investigated day-to-day, temporal assembly of a plant-pollinator network within a season, taking advantage of the extreme seasonality of pollinator communities in Greenland. Olesen *et al.* (2008) found that new within a season, the network assembly was similar to preferential attachment. New species tended to interact with already well-connected species, likely because these species are either more abundant or more temporally persistent(Olesen *et al.*, 2008). In addition, using a space-for-time substitution to study.
 - Studying primary succession along a glacier foreland, Albrecht *et al.* (2010) found some indication assembly was occurring through preferential attachment. Network a similar pattern where nestedness, a pattern of interactions where a generalist core interacts with both specialist and generalist species, increased as the community aged(Albrecht *et al.*, 2010). Increasing nestedness could result from a process like preferential attachment where specialist species attach to the well-connected, generalist core.
 - Even non-successional temporal dynamics suggest a stable core of generalists persist
 despite high turnover of peripheral species (Fang & Huang, 2012; Díaz-Castelazo et al., 2010; Alar
- In contrast to the ordered network build-up described by preferential attachment,
 assembly can may be punctuated by significant reorganizations of interactions (Peel & Clauset, 201
 For example, as new species are added, resident species change their interaction
 partners to minimize competition, or become extinct. Such significant reorganizations of interactions, or changing points, have been observed in networks responding
 to abrupt shifts in the behavior of interacters (Peel & Clauset, 2014). No studies,
 however, have examined whether changing points occur during ecological network

- assembly. (Peel & Clauset, 2014).
- ⁷³ Understanding network assembly is particularly relevant to ecological restoration, which
- is essentially 'applied succession' (Parker, 1997, e.g.,). In pollination systems, the time
- since an area was restored has been shown to effect the structure of networks (Forup et al., 2008a,b; Devo
- suggesting interactions are evolving as the community develops. Understanding the
- mechanisms of network assembly will help to guide the restoration of particular communities.
- Facilitating effective restoration of networks is particularly imperative in areas where
- 80 species interactions provide essential ecosystem services, such as crop pollination. In
- 81 intensively managed agricultural landscapes, the demand for pollination services is the
- greatest (Kremen, 2008). However, honey bees, managed extensively around the world to
- provide crop pollination, are in global decline (Neumann & Carreck, 2010; van Engelsdorp et al., 2009).
- 84 In addition, native pollinators, which have the capacity to provide sufficient crop pollination
- ⁸⁵ (Kremen et al., 2002; Winfree et al., 2007; Kremen et al., 2004), are in short supply because
- these landscapes make poor habitats for pollinator populations (Kremen et al., 2002). To
- ensure provision the continued provision of ecosystem services and curb biodiversity
- 88 loss, effective restoration of pollinators and their interactions in agricultural landscapes
- 89 is critical.

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- ⁹⁰ To promote pollinator services in agriculture, farmers are increasingly turning to the
- 91 habitat restoration technique of planting strips of native plants along farm edges (hedgerows)
- 92 to help provide habitat for pollinators without removing arable land from production.
- 93 Hedgerows have been shown to augment the richness, abundance and trait diversity of
- pollinators in agricultural landscapes (Morandin & Kremen, 2013; M'Gonigle et al., 2015; Kremen & M
- 95 In addition, hedgerows promote the persistence and colonization of floral resource specialists
- (M'Gonigle et al., 2015). Little is known however, about the assembly of the network

97 following hedgerow restoration.

Using a long-term data-set of plant-pollinator communities assembling following hedgerow restoration in the highly simplified and intensively managed agricultural landscape of California's Central Valley, we explore the process of network development. We first 100 determine whether the mechanism underlying network assembly is a smooth build up of 101 interactions as would be predicted by preferential attachment, or punctuated by significant 102 reorganizations of interactions (i.e., changing points). Even with changing points in interaction 103 organization, networks could still be assembling via preferential attachment if the network 104 reorganizations were primarily driven the by peripheral, temporally variable species while 105 a stable, well-connected core of species still persists. We thus examine whether the species 106 are most variable in their network position — and thus important contributors network 107 reorganizations — are less persistent and connected species. Lastly, we examine whether 108 networks are assembling toward predictable interaction patterns, and the ramifications 109 for the robustness of the networks to species extinction and perturbation. 110

Materials & Methods

112 Study sites and collection methods

We surveyed plant-pollinator interaction networks of assembling hedgerows (N=5), as well as in two types of non-assembling communities to serve as controls: unrestored, weedy field margins (N=19XX) and established hedgerows (greater than 10 years since planting, N=29XX). The sites were located in the Central Valley of California in Yolo, Colusa and Solano Counties. This The area is comprised of intensively managed agriculture – primarily monocultures of conventional row crops, vineyards and orchards.

Hedgerows we were planted along field margins where they do not remove valuable land from production, and are ca. 3–6m wide and approximately 350m long and border large 120 (ca. 30-hectare) crop fields. Hedgerows consist of native, perennial, shrub and tree plant-121 ings including Rosa californica, Cercis occidentalis, Ceanothus spp., Heteromeles arbutifolia, 122 Sambucus mexicana, Eriogonum spp., Baccharis spp., Salvia spp. and others (Kremen & M'Gonigle, 2015; M'Gonig 123 The mean distance between monitoring sites was 15 km, and the minimum distance be-124 tween sites of the same type sampled in the same year was 2 km. The entire area surveyed 125 spanned almost 300 km². The and the crop fields adjacent to all sites were similarly man-126 aged as intensive, high-input monoculture. 127

Monitoring began in 2006 and continued through 2015. Sampling of the assembling hedgerows began the year before the area was restored. For logistical reasons, no sampling of assembling hedgerows was conducted in 2010. Sites were sampled between two and five times per year (Tables S1-S3Table XX). In each round of sampling, the order in which sites were sampled was randomized. Surveys were conducted under sunny conditions when the temperature was above 21°C and wind speed was below 2.5 meters/second.

Flower-visitors to plants in hedgerows and unrestored controls were netted for one hour of active search time (the timer was paused when handling specimens). Honeybees (*Apis mellifera*) were not collected because their abundance is determined largely by the placement of hives throughout the region by bee-keepers. All other insect flower visitors that touched the reproductive parts of the flower were collected; however, here we focus only on wild bees and syrphids (representing 49 and 19 XX and XX percent of records, respectively), the most abundant and effective pollinators in the system (C. Kremen, A. Klein and L. Morandin, unpublished data). Bee and syrphid specimens were identified to species (or morpho-species for some bee specimens in the genera *Nomada* and *Sphecodes*)

by expert taxonomists.

Quantitative networks were generative for each site through time. To account for the unequal number of surveys between years, the mean of the interactions between a pair of plants and pollinators across surveys within a year was used as a representation of the frequency of interactions.

149 Change point analysis

50 Identifying change points

We employed a change point detection method (Peel & Clauset, 2014) to identify funda-151 mental changes in large-scale pattern of interactions of plants and pollinators. A change point is caused by a merge, split, fragmentation or formation of communities (also called 153 modules or compartments). Change point detection methods have yet to be generalized 154 to quantitative networks, so for this analysis we focused on qualitative (binary) networks. 155 Following Peel & Clauset (2014), we first defined a probability distribution over the net-156 works using the generalized hierarchical random graph model (GHRG). The GHRG model 157 is able to capture both assortative and disassortative community structure patterns at all 158 scales in the network (Peel & Clauset, 2014). A network G is composed of vertices V 159 and edges $E \subseteq VV$. The GHRG model decomposes the N vertices into a series of nested 160 groups, the relationships among which are represented by the dendrogram T. The tips of 161 T are the vertices of G, and the probability that two vertices u and v connect is given by 162 the parameter p_r . The probability distribution of the network G thus defined as: 163

$$P(G|T, pr) = p_r^{E_r} (1 - p_r)^{N_r - E_r}$$
(1)

Where E_r is the observed number of edges between vertices with the common ancestor r, and N_r is the total possible edges.

Using Bayesian posterior inference and techniques from phylogenetic tree reconstruction, we fit the GHRG model to the networks (Peel & Clauset, 2014). This is accomplished by using a Markov Chain Monte Carlo (MCMC) procedure to first sample the posterior distribution of bipartitions, from which a consensus tree is derived (Peel & Clauset, 2014). ADD NEWMAN PAPER. We used β distributions with the hyperparameters $\alpha = \beta = 1$ to define priors for p_r .

Once the GHRG model has been fit to the networks, we determine whether a change 172 point occurred between two time slices. To detect a change point, we compare the fit of 173 two models – one where a change point had occurred between two networks, and one where no change occurred – using posterior Bayes factors. We chose a sliding window of 175 length, w, of four, within which to find change points. Larger windows allow for more 176 gradual changes, and four was the maximum possible with our maximum of nine years of 177 data. Lastly, to calculate a p-value for the Bayes factors, we use parametric bootstrapping 178 to numerically estimate the null distribution (Peel & Clauset, 2014). The change point 179 analysis was carried out using code published online by L. Peel. Analyses we conducted 180 in Python 3.4. .Peel. 181

We next test whether the change points occurring in maturing hedgerows were a component of the assembly process or a product of environmental shifts that lead to network
reorganizations in all types of communities. We model the number of change points as
successes and the total number of years each site was sampled as trails, and use a generalized linear model with Binomial error to test whether the probability of a change
point occurring varied by site type. For the non-assembling hedgerows and weedy field
margins, only sites with five or greater years of sampling was included in this analysis.

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190 Characteristics of species that contribute to change points" core" and "peripheral" communities

To further elucidate the nature of the change points, we examine the characteristics of the species that contributed the reorganization of interactions. Some species remain in relatively similar network positions through time, whereas others are more variable in their role and thus contribute more strongly to network reorganization. We test the that more persistent species with the highest degree are the most stable in their network position, as would be expected is the networks were assembling via preferential attachment.

We calculate species persistence as the proportion of the surveys a pollinator was observed. 198 Species observed consistently within and between years are thus maximally persistent. 199 Weighted species degree is calculated from interaction observations observed in more 200 extensive data-set from Yolo County (approx. 18000 interaction records) that included 201 both the data included in this study and additional data from sites where we collected 202 flower visitors using the same methods (M'Gonigle et al., 2015; Ponisio et al., 2015). To 203 represent the the variability of species within networks, we computed the coefficient 204 of variation of weighted closeness at each site through time. Closeness describes the 205 centrality of a species in the network by calculating path lengths to other vertices (species) 206 in the graph. We used linear mixed models to test whether the variability of species 207 closeness values was related to the persistence or degree of that species (Bates et al., 2014; Kuznetsova et 208 We included random effects for species, as well as site. We focused on the pollinator 209 species because the hedgerow flowers are planted and thus are not directly assembling. 210 Because degree and persistence were strongly correlated, ($\rho = 0.84$, p-value $< 2 * 10^{-16}$), 211 each explanatory variable was included in the model separately. Because a linear increase

in closeness, as might be expected with assembly by preferential attachment, would lead to a high variability in closeness scores, we To do so, we first generate dendrograms using 214 the GHRG model before and after each change point. We then determined which species 215 belonged to the "core" and "peripheral" network communities at each site. The "core" 216 network communities contain the majority of species and are more basal than the more 217 derived, less specious "peripheral" network communities. We next use a Permutational 218 Multivariate Analysis of Variance (PERMANOVA) (Anderson & Walsh, 2013) to determine 219 whether the species compositions of the species that belonged to the core and peripheral 220 communities differed. We also test whether closeness increases through timecore or peripheral 221 species had more variability in their species compositions (i.e., multivariate dispersion, Anderson et al.,

Temporal changes in interaction patterns

Network structure

The changing points in network structure may contribute to the reorganization of the assembling networks into predictable interaction patterns. Pollination networks exhibit two 226 main topologies structural patterns — modularity (e.g., Olesen et al., 2007) and nestedness (e.g., Bascompte et al., 2006, 2003). Modular In modular community interactions are in-228 sular, occurring within separate groups or "modules" more often than between modules. 229 Communities in the network may fragment as the network assembles, enhancing modu-230 larity. Conversely, nested networks are like pyramid of interactions, where there are some 231 species that interact with many species, other species that interact with a subset of those 232 species, and so on. If species entering the network tend to interact with the generalist 233 base of the network pyramid (i.e., preferential attachment), nestedness would increase 234 through time. Lastly Alternatively, if the network is accumulating specialist species or if

species or species are beginning to limit their interaction niche breath as the network assembles, this would lead to an increase in the network-level specialization (Blüthgen et al., 2006) and nestedness would decrease through time. To test whether network modularity, 238 nestedness or specialization changed linearly with assembly, we used linear mixed mod-239 els with the descriptive network metrics as the response variable, year of assembly as the 240 explanatory variable, and random effects of site and year. 241 We use NODF (weighted or unweighted) to evaluate network nestedness (Almeida-Neto et al., 2008). 242 NODF evaluates whether species with fewer partners interact with subsets of partners 243 with which more connected species interact (Almeida-Neto et al., 2008). To estimate modularity, 244 we use a hierarchical clustering algorithm (Newman & Girvan, 2004; Csardi & Nepusz, 2006). 245 We calculated standardized z-scores so that nestedness and modularity metrics could be 246 compared between communities. The z-scores were calculated by generating an ensemble 247 of 999 randomly assembled communities, subtracting the mean of the statistic calculated 248 across these communities from the observed value, and then dividing by the standard 249 deviation. To assemble random communities, we reshuffled the interactions between 250 species but fixed the total number of interactions, species and the distribution of the 251 interaction frequencies (Galeano et al., 2009). Lastly, Network specialization was measured 252 using H2, which estimate the deviation of the observed interaction frequency between 253 plants and pollinators from a null expectation where all partners interact in proportion 254 to their abundances (Blüthgen et al., 2006). It ranges from 0 for generalized networks to 1 255 for specialized networks.

Network robustness

Lastly, we test whether the changes in interaction patterns associated with network assembly affect the robustness. First, following of the network to species loss. Following Memmott *et al.* (2004), we simulate the extinction of plant species the subsequent extinction cascades of pollinator species. Because the reproduction of plant species if facilitated by active restoration efforts, it is unlikely the extinction of pollinators would effect affect plant populations in the hedgerows. However, plants ceasing to bloom, for example in response to drought, will likely effect affect the pollinators that depend on them. Plants species were eliminated based on their degree or abundance, and the number of pollinators that secondarily went extinct is calculated. The area below the extinction curve is used as a measure of network robustness(Memmott *et al.*, 2004).

Results

Over eight years and 747 samples, we collected and identified 19,547 wild bees and syrphids comprising 173 species from 50 genera. We observed 1,521 unique interactions between plants and pollinators.

72 Change point analysis

273 Identifying change points

The majority (76%) of the sites tests underwent at least one significant reorganization of interactions (Fig. 1). There were no consistent trends as to when change points occurred within assembling hedgerows or across all sites, except many site had changing points between year 2009 and 2011 (Fig. 1). In California, 2011 marked the beginning of a multi year drought. In the assembling hedgerows were not sampled in 2010, so disentangling whether the changing points are due to skipping a year of assembly or the drought is not impossible.

All five of the assembling hedgerows experienced changing points, whereas only 40% and 81% of non-assembling hedgerows and field margins, respectively, underwent significant interaction reorganizations. Assembling hedgerows experienced significantly more changing points than the non-assembling networks (difference in the odds ratios between assembling and non-assembling networks, 3.316, 95% CI [1.314, 8.572], p-value= 0.0117). Network assembly following restoration is thus punctuated by more interaction reorganizations than would be expected by environmental shifts that would effect assembling and non-assembling networks equally.

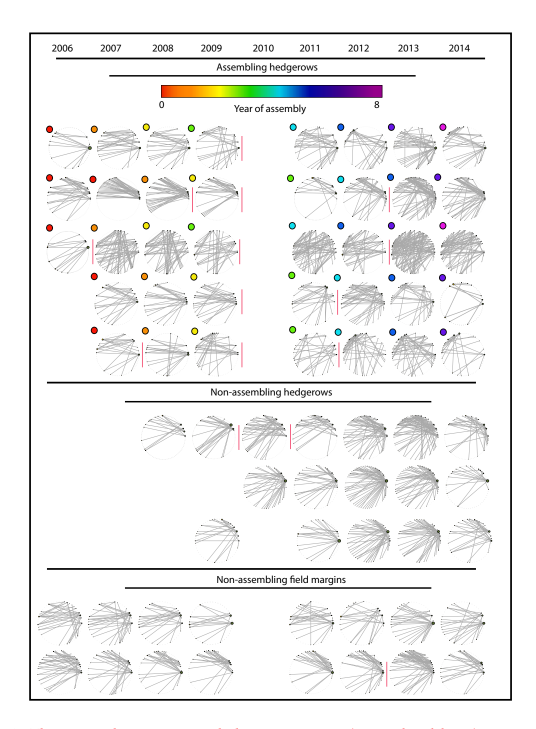


Figure 1: The network structure and changing points (vertical red lines) in assembling hedgerows and a representative sample of non-assembling hedgerows and weedy field margins. In each network, plants and pollinators are represented by green and yellow circles, respectively, weighted by their degree. Each species has has a consistent position in the network across years. In the assembling hedgerows, colored circles in the corner of each network represent the years post restoration. XXX

- 289 Characteristics of species that contribute to change points "core" and "peripheral" communities
- ²⁹⁰ In contradiction to the predictions of assembly by preferential attachment, both pollinator
- persistence and degree were positively related to network position variability. (ADD
- 292 STATS IF KEEPING RESULT).

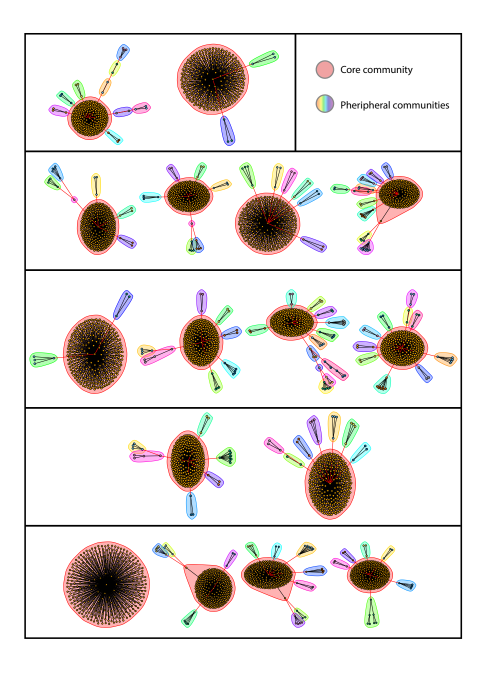


Figure 2: XXX

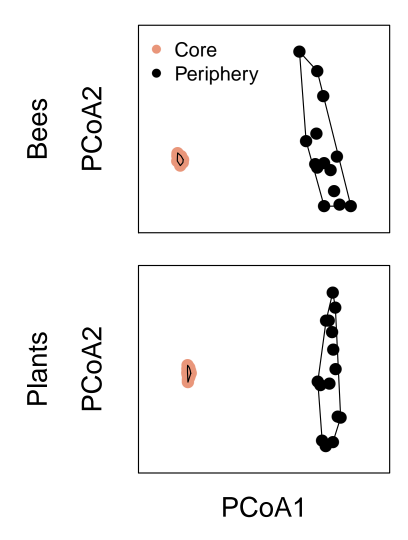


Figure 3: The coefficient of variation of network position, as represented by closeness, plotted against pollinator persistence and degree. Points represents means for each species across sites. The solid line indicates the mean slope estimate and the dashed lines are the 95% confidence intervals around the estimate. XXX

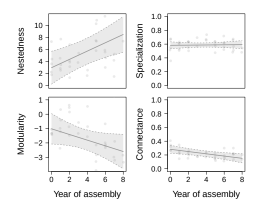


Figure 4: The change in nestedness, modularity, specialization and connectance as the networks assemble. The left panels represent z-scores. Scores greater than ~ 2 or less than ~ -2 are significantly more or less structured than randomly assembled networks. Points are the value for each site at each year of assembly. The solid line indicates the mean slope estimate and the dashed lines are the 95% confidence intervals around the estimate.XXX

33 Temporal changes in interaction patterns

Network structure

- 295 Network nestedness significantly increased with assembly (estimate of the slope of nestedness
- through time \pm standard error of the estimate, 1.834 ± 0.6142 , p-value=0.022, Fig. 4).
- 297 Modularity decreased (Fig. 4), though the slope was not significantly different from
- 298 zero (estimate of the slope of modularity through time ± standard error of the estimate,
- -0.524 ± 0.295 , p-value=0.124). Specialization remained relatively constant (estimate of
- the slope of specialization through time \pm standard error of the estimate, 0.003 ± 0.015 ,
- 301 *p*-value=0.827).

Network robustness

Assembly did not effect the robustness of the networks to species extinction when species where removed incrementally by degree (estimate of the slope of robustness through 304 time \pm standard error of the estimate, $6*10^{-5} \pm 4*10^{-3}$, p-value=0.987) or abundance 305 $(0.001 \pm 0.003, p$ -value=0.65, Fig. ??). 306 In contrast, the robustness of networks to perturbation, as measured by the algebraic connectivity of the network, increased as the network assembled (estimate of the slope of 308 robustness through time \pm standard error of the estimate, 0.6814 \pm 0.272, p-value=0.042, 309 Fig. ??). 310 The robustness of networks to species extinction and perturbation. The robustness to species extinction is measured by incrementally removing species by degree, through 312 removing species by abundance did not yield qualitatively different results. Points are 313 the value for each site at each year of assembly. The solid line indicates the mean slope 314 estimate and the dashed lines are the 95% confidence intervals around the estimate.

Discussion

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