

A needle in the haystack? Applying species co-occurrence frameworks with fish assemblage data to identify species associations and sharpen ecological hypotheses

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Abstract

Different species can associate or interact in many ways, and methods exist for inferring associations and underlying mechanisms from incidence data (e.g., co-occurrence frameworks). These methods have received criticism despite their recent resurgence in the literature. However, co-occurrence frameworks for identifying nonrandomly associated species pairs (e.g., aggregated or segregated pairs) have value as heuristic tools for sharpening hypotheses concerning fish ecology. This paper provides a case study examining species co-occurrence across 33 stream fish assemblages in southeastern Oklahoma, USA, which were sampled twice (1974 and 2014). This study sought to determine (a) which species were nonrandomly associated, (b) what processes might have driven these associations and (c) how consistent patterns were across time. Associations among most pairs of species (24 species, 276 unique pairs) were not significantly different from random (>80%). Among all significant, nonrandomly associated species pairs (54 unique pairs), 78% (42 pairs) were aggregated and 22% (12 pairs) segregated. Most of these (28 pairs, 52%) were hypothesized to be driven by nonbiotic mechanisms: habitat filtering (20 pairs, 37%), dispersal limitation (two pairs, 0.4%) or both (six pairs, 11%). The remaining 26 nonrandomly associated pairs (48%) had no detectable signal of spatial or environmental factors involved with the association, therefore the potential for biotic interaction was not refuted. Only five species pairs were consistently associated across both sampling periods: stonerollers *Camptostoma* spp. and orangebelly darter *Etheostoma radiosum*; red shiner *Cyprinella lutrensis* and bullhead minnow *Pimephales vigilax*; bluegill sunfish *Lepomis macrochirus* and redear sunfish *Lepomis microlophus*; redbfin shiner *Lythrurus umbratilis* and bluntnose minnow *Pimephales notatus*; and bigeye shiner *Notropis boops* and golden shiner *Notemigonus crysoleucas*. Frameworks for identifying nonrandomly associated species pairs can provide insight into broader mechanisms of species assembly and point to potentially interesting species interactions (out of many possible pairs). However, this approach is best applied as a tool for sharpening hypotheses to be investigated further. Rather than a weakness, the heuristic nature is the strength of such methods, and can help guide biologists toward better questions by

employing relatively cheap diversity survey data, which are often already in hand, to reduce complex interaction networks down to their nonstochastic parts which warrant further investigation.

KEYWORDS

biotic interactions, dispersal limitation, habitat filtering, nonrandom association, null model analysis, probabilistic species association

1 | INTRODUCTION

Interactions among species are an important piece of the ecological and evolutionary puzzle (Benton, 2009; Schemske *et al.*, 2009) and, because of interdependence among species, a key component of wholistic conservation (Gilman *et al.*, 2010; Wisz *et al.*, 2013). Within and among assemblages there are myriad ways individuals of different species can interact in space and time. Considerable attention has been given to competition (Bonin *et al.*, 2015; Schluter, 1994), which has waxed and waned considerably through time (Matthews, 1998; Wiens, 1977). Other examples include predator/prey (or parasite/host) interactions (Aragón & Sánchez-Fernández, 2013; Power *et al.*, 1985; Zurell *et al.*, 2018) and less commonly studied facilitative and mutualistic interactions (Peoples & Frimpong, 2016; Silknetter *et al.*, 2020; Stachowicz, 2001). Even in the absence of biotic interaction among species, other forces can lead to certain species co-occurring together more or less often than expected by chance alone (aggregation or segregation, respectively), including habitat filtering, niche partitioning and species packing (Geheber & Frenette, 2016; Heino & Grönroos, 2013; Werner, 1977), spatial filtering via dispersal limitation or historical contingency (Blois *et al.*, 2014; Gilpin & Diamond, 1982; Schamp *et al.*, 2015), and neutral or stochastic dynamics such as ecological drift (Bell, 2005; Hubbell, 2001; Ulrich, 2004). Further adding to the complexity of the interdependent web among interacting species is the element of time and space. For example, changes in density of one or both interactors through time due to predation or disturbance (McMeans *et al.*, 2020; Wiens, 1977) and/or seasonality (Brimacombe *et al.*, 2020), or through space due to environmental gradients (Case & Taper, 2000; Schwillk & Ackerly, 2005), can obscure—or at least alter—any pattern of association we might expect to observe.

Detecting species interactions has often relied on controlled experiments or direct natural history observation (Silknetter *et al.*, 2020). Even in small systems the number of interactions increases quickly with the number of species until it becomes intractable to assess each individually (Morales-Castilla *et al.*, 2015). Therefore, ecologists have developed methods to statistically test for nonrandom associations among species from incidence data as a means to circumvent such an arduous task (Forbes, 1907; Pielou & Pielou, 1968). These methods were largely born out of an interest in identifying competition among species (Diamond, 1975), but were later generalized using null modelling to provide more rigorous tests for identifying patterns of both aggregation and segregation (Connor & Simberloff,

1979; Gotelli, 2000; Peres-Neto, 2004). Because of the multitude of factors, both biotic and abiotic, which can drive patterns of association (Barner *et al.*, 2018; Blanchet *et al.*, 2020), more recent work has combined earlier methods for identifying nonrandom associations with additional multivariate approaches (*e.g.*, Blois *et al.*, 2014) which incorporate spatial and environmental data to disentangle the potential ecological mechanisms driving observed patterns (Cordero & Jackson, 2019; Devercelli *et al.*, 2016; Sfenthourakis *et al.*, 2006).

Both methods for identifying associations and those more recent extensions aimed at parsing them have faced considerable scrutiny (Blanchet *et al.*, 2020; Peterson *et al.*, 2020; Thurman *et al.*, 2019). Methods for identifying associations face several issues. For one, as with many ecological methods, there is a variety of approaches and they are not all equally robust for detecting associations (Barner *et al.*, 2018; Lavender *et al.*, 2019). Furthermore, ecological interactions may not lead to the clear spatial patterns that we expect (Freilich *et al.*, 2018) and thus these methods can fall short of detecting them (Cazelles *et al.*, 2016). In addition, and as with all things ecological, sampling biases caused by the difficult detection of rare species (Calatayud *et al.*, 2020; MacKenzie *et al.*, 2004) and effects of spatial scale of sampling grain (Perry *et al.*, 2020) can affect the ability of these methods to reliably identify real associations (Peterson *et al.*, 2020). As for interpreting ecological mechanisms from co-occurrence data, it can be difficult to determine whether the pattern is driven by habitat filtering or biotic processes because it is impossible to account for each environmental facet that may govern an association, therefore unmeasured factors confound such conclusions (Blanchet *et al.*, 2020). Also, interaction networks (more than two species) preclude pairwise analyses from detecting effects of additional species which may have direct or indirect effects on either member of a pair (Cazelles *et al.*, 2016; Holt & Bonsall, 2017). Other properties of ecosystems, such as habitat gradients, demographic dynamics and temporal changes (Brimacombe *et al.*, 2020), all play a role in shifting species associations and our ability to detect them, especially with incidence data that merely provide a ‘snapshot’ of diversity in space and time (Blanchet *et al.*, 2020).

Species incidence is among the easiest of ecological data to gather, and so it is reasonable to hope to extract as much information as possible (*e.g.*, species associations) from such data (Blanchet *et al.*, 2020). One of the biggest issues with these methods is what users expect to get out of them. Because of the issues described above, it is not reasonable to expect a definitive test of species associations and mechanisms from such methods. Rather, species co-occurrence

analyses are an exploratory tool (Silknetter *et al.*, 2020) for finding statistically nonrandom associations among large sets of potentially interacting pairs and sharpening hypotheses regarding the potential underlying mechanisms. These associations should be further evaluated through direct observation or experimental means to test these hypotheses. Given the large number of species interactions in even small communities (e.g., 10 species = 45 possible pairwise associations), these methods can help us find the ‘needles in the haystack’, so to speak, and ask more focused questions about how fish species interact.

Here, two frameworks are applied to investigate freshwater fish species associations and how they have changed between two sampling campaigns. These tests addressed several questions: (a) Which species pairs are nonrandomly associated and how many are there? (b) What do these frameworks tell us about mechanisms driving species associations? and (c) Are species associations consistent among two sampling campaigns executed 40 years apart?

2 | MATERIALS AND METHODS

2.1 | Study region and fish collection

The study region was in the south-central United States (southeastern Oklahoma) within a relatively rugged topographic region known as the U.S. Interior Highlands (Figure 1). Muddy Boggy Creek and Clear Boggy Creek merge to form the Muddy Boggy River, which is a tributary to the Red River (Mississippi Basin). The Muddy Boggy River

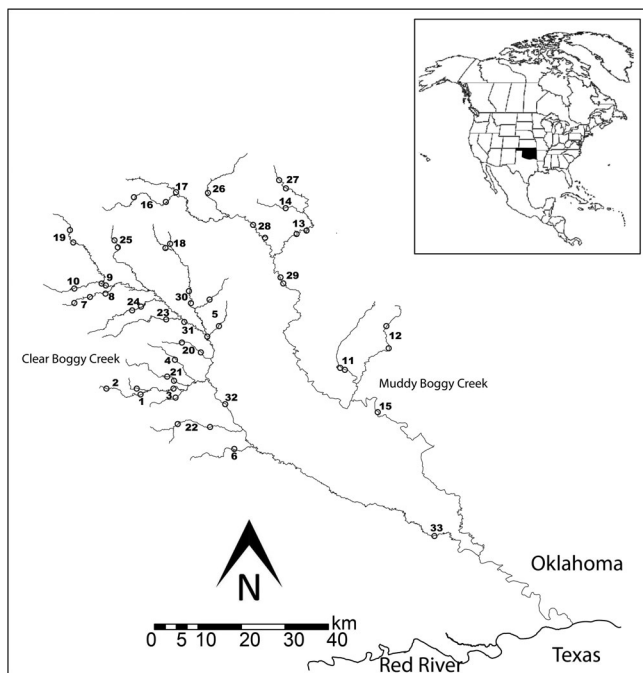


FIGURE 1 Map of the Muddy Boggy River in south-eastern, OK, USA. Sites where fish specimens were collected are marked with a hollow black circle and the sampling ID is noted next to each

flows southward and drains 6291 km² from an area approximately 113 km (north–south) by 48 km (east–west). See Pigg (1977) and Zbinden and Matthews (2017) for more detailed descriptions of the region.

Fish collections were made in the Muddy Boggy River drainage between 1974 and 1975 (Pigg, 1977) and between 2014 and 2015 (Zbinden & Matthews, 2017). From these collections, 33 sites sampled at both times were included based on a previous study that used these sites to quantify fish assemblage change over time (Zbinden, 2020). All collections were made between May and September during low flows. Seines of the appropriate size relative to the sampling reach (1.22×2.44 or 4.57 m) with mesh sizes approximately 4.88 mm were used to sample all available habitat within a stream reach transect 100 m in length. The fish collected at a site/locality constitute the ‘fish assemblage’, which is synonymous with the term ‘fish community’ used by others for the sake of consistency (e.g., Matthews & Marsh-Matthews, 2017). However, assemblage was chosen because fish alone are a phylogenetically restricted piece of the whole community, therefore an assemblage (*sensu* Fauth *et al.*, 1996). Prior to analyses, only species captured during both sampling periods were retained, and species occurring at fewer than six sites in both time periods were removed because of low statistical power for testing associations (Cordero & Jackson, 2019; Gotelli & McCabe, 2002; Veech, 2013). Two taxonomic changes were required. Central stoneroller *Campostoma anomalum* (Rafinesque 1820) and highland stoneroller *Campostoma spadiceum* (Girard 1856) collected in 2014 were collapsed into *Campostoma* spp. because recognition of *C. spadiceum* did not occur until after 1975 (Cashner *et al.*, 2010) and thus were not differentiated by J. Pigg. In the same manner, blackstripe topminnow *Fundulus notatus* (Rafinesque 1820) and blackspotted topminnow *Fundulus olivaceus* (Storer 1845) were collapsed into *Fundulus* spp. due to identification issues in this region (J.F. Schaefer and W.J. Matthews, pers. comm.)

2.2 | Spatial data

The pairwise fluvial network distance was calculated among all fish sampling locations using the Network Analyst Extension in ArcGIS v.10.8 (ESRI). This distance matrix was then decomposed into a rectangular matrix of spatial eigenvectors using distance-based Moran's eigenvector map analysis (dbMEM; Stéphane Dray *et al.*, 2006), also known as principal coordinates of neighbour matrices (Borcard & Legendre, 2002). The spatial eigenvectors were generated using the function `dbmem` in the R statistical software (v.3.6.1, R Core Team, 2019) package `adespatial` (Dray *et al.*, 2020), and only eigenvectors with positive spatial autocorrelation (six MEMs) were retained.

2.3 | Environmental data

Environmental factors for each collection site were characterized using data provided by HydroATLAS v.1.0 at 15 arc-second (~500 m)

resolution (Linke *et al.*, 2019). This global compendium of sub-basin and river reach characteristics contains 56 hydro-environmental factors partitioned into 281 attributes drawn from a multitude of global datasets (Linke *et al.*, 2019). The RiverATLAS v.10 shapefile for North America was trimmed in ArcMAP (ESRI v.10.8) to include only the subset of river network necessary for this study. This shapefile represented the river network and all associated environmental variables along the stream reaches, which are defined as stretches of stream between confluences or confluence and terminus. Geographic coordinate data of sampling sites were imported into the GIS and exported as a shapefile. Both shapefiles were projected using NAD 1983 (2011) Contiguous USA Albers projection. Environmental attributes of the river shapefile were joined to the appropriate sites using R packages *raster* (Hijmans & van Etten, 2015) and *sf* (Pebesma, 2019). From the total set (281 attributes), 22 variables were chosen because it was expected that they would be similar between sampling periods or because they represented long-term averages. To be clear, the environmental data was identical between each sampling period, but the characteristics were chosen specifically to match that assumption.

The environmental variables broadly characterize the hydrologic, physiographic and geologic nature of the sampling locations. These reach catchment variables (unless otherwise noted) included Strahler stream order, mean annual discharge (1971–2000), mean annual land surface runoff (1971–2000), maximum inundation extent (1993–2004), river area, river area in the upstream catchment, river volume, river volume in the upstream catchment, mean groundwater table depth (1927–2009), mean elevation, mean terrain slope, mean stream gradient, mean annual precipitation (1950–2000), extent of deciduous tree cover in reach catchment, extent of evergreen tree cover in reach catchment, extent of herbaceous cover in reach catchment, total forest cover in reach catchment, mean fraction of clay in soil of reach catchment, mean fraction of silt in soil of reach catchment, mean fraction of sand in soil of reach catchment, mean organic carbon fraction in soil of reach catchment and spatial extent of exposed karst in reach catchment. The descriptions above are based on the HydroATLAS catalogue (<https://hydrosheds.org/page/hydroatlas>).

All environmental factors were standardized by mean and unit standard deviation before being reduced using principal components analysis, using function `prcomp` of the R package *stats* (R Core Team, 2019) without scaling or centering because variables had already been standardized. Six of the principal components were retained to make the number of factors equal between spatial and environmental sets [six principal components (PCs) and six MEMs]. These six PCs accounted for 87.6% of variation in the environmental set. See the Supporting Information (Table S1) for full environmental PCA details and factor loadings on each component.

Two testing frameworks (Figure 2) aimed at detecting significant nonrandom species associations and subsequently hypothesizing mechanisms that govern associations by incorporating the spatial and environmental vectors were used (Blois *et al.*, 2014; D'Amen *et al.*, 2018; Gotelli & Ulrich, 2010; Veech, 2013). The Null Model Framework (Classic Framework *sensu* D'Amen *et al.*, 2018) and The Probabilistic Framework (Veech, 2013) differ only in how significantly

nonrandom species associations are tested, but the same decision tree approach for statistically deducing mechanisms *via* multivariate analysis of variance is employed (descriptions below).

2.4 | Null model framework

The null model framework (NMF) uses the C-score index, a measure of 'checkerboardedness' among species pairs (Stone & Roberts, 1990). The original index measured the average C-score for the entire species incidence matrix, but here an index for each species pair is desired. The following equation was used:

$$Cscore_{ab} = \frac{(R_a - S) \times (R_b - S)}{(R_a \times R_b)}$$

where the number of occurrences of species *a* is R_a and of species *b* is R_b , and *S* is the number of sites where both species were collected. A maximally aggregated species pair (all occurrences shared) will have a C-score of 0 and a maximally segregated species pair (no sites in common) will have a C-score of 1.

C-score significance for each species pair was tested by generating 10,000 null assemblages by randomly reshuffling the observed matrix. Column sums (species incidences) and row sums (per site species richness) of null assemblages were constrained to be equal to the observed matrix (fixed-fixed randomization *sensu* Connor & Simberloff, 1979; Gotelli, 2000). Observed C-scores for each pair were compared to the appropriate null distribution to test for significance ($\alpha = 0.05$). Where *n* is equal to the number of unique species in the presence-absence matrix, the total number of species pairs tested is equal to $n(n - 1)/2$. To reduce false detection error rate a mean-based Bayes approach (Bayes M criterion) from Gotelli and Ulrich (2010) and originally described by Efron (2005) was used. Significantly associated species pairs are grouped as either aggregated or segregated by comparing observed C-score with the mean randomized score for a given species pair. Z-scores were calculated based on 10,000 randomizations. Large positive scores are associated with segregating species and large negative scores are associated with aggregating species (D'Amen *et al.*, 2018; Gotelli & Ulrich, 2010). Results from null model analysis were then used with the deductive logic tree approach described below to hypothesize the processes underlying each significant association (Figure 2).

2.5 | Probabilistic framework

The probabilistic framework (PRF) is an entirely different approach for testing patterns of species co-occurrence based on probability theory. This approach uses combinatorics (*i.e.*, probability within finite sets) to estimate the probability of species co-occurring at greater than or less than observed sites by considering the possible combinations in which a given pattern of species distributions in *N* sites could possibly be observed (Equation 1 and Figure 2 from Veech, 2013). These

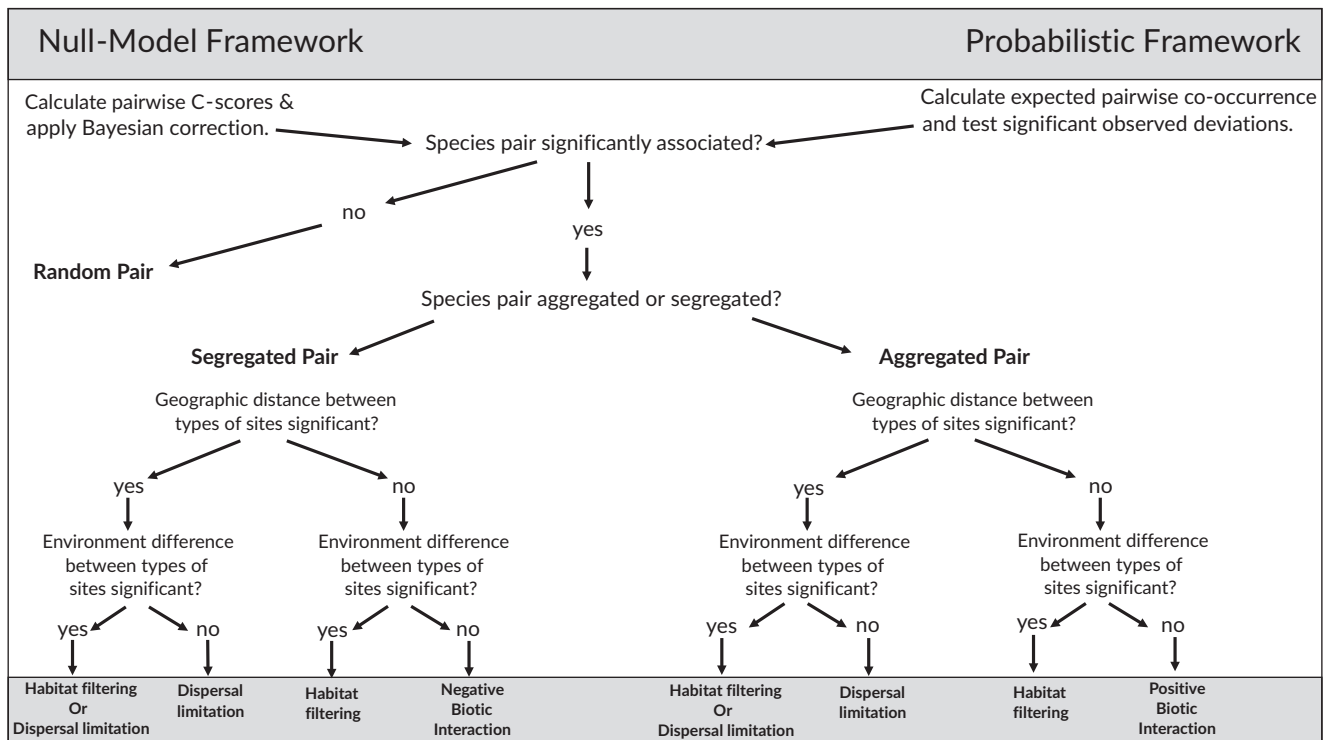


FIGURE 2 Logic tree for deducing inferences driving significant species associations from incidence data. Null model and probabilistic frameworks each use the same logic tree to make inferences (illustrated here). The frameworks differ in how significant associations are determined. The null model framework relies on randomly shuffling the data matrix to generate random expectations, and the probabilistic framework uses probabilistic combinatorics to quantify the expected probabilities of species aggregation and segregation based on the data set

probabilities may be obtained analytically and do not require a null distribution or data randomization, which can itself introduce Type I and II errors (Lavender *et al.*, 2019; Veech, 2014). The probabilistic approach determines whether species are positively (*i.e.*, aggregated), negatively (*i.e.*, segregated), or randomly associated. The probabilistic calculations are implemented in the R package *cooccur* (Griffith *et al.*, 2016). Although the NMF and the PRF differ in mathematical approach, both result in a set of significantly associated species pairs which can be further analysed as described below.

2.6 | Decision tree for statistically deducing mechanisms

The analyses of species pairwise co-occurrence presented above cannot determine the processes responsible for nonrandom species associations by themselves, although they are often inferred as indicative of biotic interaction (Peterson *et al.*, 2020). However, by including additional analyses of spatial and environmental data for each site, it is possible to determine whether dispersal limitation or habitat filtering are likely forces driving nonrandom associations, rather than biotic interaction. If neither of those forces are significant, it is not possible to refute a biotic interaction is possible. However, this hypothesis should be tested further through direct observation or experimentation. Both the NMF and PRF incorporate the same decision tree

approach for disentangling mechanisms (Figure 2). This approach is modified from Blois *et al.* (2014) and D'Amen *et al.* (2018).

Ultimately, multivariate analysis of variance (MANOVA) of different site categories is used to test for significant spatial and environmental effects on the nonrandom associations inferred from the analyses described above. For each significantly associated species pair, the sites in the incidence matrix belong to one of four mutually exclusive site-type categories: both species present (1,1), neither species present (0,0), or one species present without the other (1,0) and (0,1). Environmental and spatial data are partitioned between site-type categories to test for significant differences among them via MANOVA. These tests inform the decision tree and lead to logical, deductive inferences regarding potential mechanisms involved in species pair associations (Figure 2).

Allotopic sites, having one species without the other (1,0) and (0,1), are contrasted to discern mechanisms for *segregated* species pairs. If environmental factors and/or spatial arrangement differ among these two site classes, then the pattern of segregation is likely driven by those factors. However, if neither environment nor spatial factors differ, then negative biotic interaction (*e.g.*, competition or predation) is not rejected. However, this case cannot exclude other alternative hypotheses regarding segregation. For example, the pattern could be driven by differences in unmeasured environmental factors. Syntopic sites where both species were found (1,1) and sites with neither species (0,0) are compared to discern potential mechanisms for *aggregated* species pairs in the same manner. Again, in the absence of

significant environmental or spatial difference between site-type classes, the possibility of a biotic interaction remains. This could include several interactions, including facilitation, predation or indirect effects with other species (Blois *et al.*, 2014) or unmeasured environmental factors.

Within each of the four frameworks (NMF1974, NMF2014, PRF1974 and PRF2014), and following the identification of significant associations, each significant species pair was tested for differences in site-type categories (depending on aggregation or segregation as explained) using MANOVA. The test for environmental effects was based on six environmental PCs, while the test for spatial effects was based on six spatial eigenvectors, both retained from above. A significant ($\alpha = 0.05$) environmental effect is hypothesized to mean environmental filtering is the mechanism underlying the association. A significant ($\alpha = 0.05$) spatial effect is hypothesized to mean dispersal limitation is the driving force. In the case that both spatial and environmental effects are significant it is likely that spatially autocorrelated habitat filtering is responsible. If neither is significant, then it is not possible to reject the hypothesis of a biotic interaction underlying the association. Marginally significant ($P < 0.1$) effects are considered as enough evidence of spatial or environmental signal to reduce associations that were attributed to biotic interactions (see Supporting Information Tables S2-S7 for the full list of results).

R code is available in the Supporting Information as R-scripts: “null-model_framework.R” and “probabilistic_framework.R”. In addition, all data files, including species incidence plus environment factors, shapefiles, and fluvial distance matrix, are available in the Supporting Information.

2.7 | Ethical Statement

The collection of animals complied with the University of Oklahoma Institutional Animal Care and Use Committee guidelines and policies (OU-IACUC approval No. R14-002). All specimen collecting was permitted by the Oklahoma Department of Wildlife Conservation (No. 5995 and No. 6166).

3 | RESULTS

Twenty-four of the same native freshwater fish species were collected in both sampling periods and were present in at least six samples in at least one sampling period (= 276 unique species pairs

analysed). No matter the year or framework, most species associations were not different from random (Table 1). The PRF resulted in more nonrandom species associations in both years (Table 1). Results from NMF were skewed toward segregating associations, while those resulting from PRF were mostly aggregations (Table 1). For 1974, five of 10 associations from the NMF were also recovered with PRF; for 2014, it was only one of four associations. The associations found exclusively by NMF in either year were all segregations. Associations from each year and framework combination are available in Supporting Information (Tables S2-S5).

Fifty-four unique nonrandom species associations (19.6%) were found across years and frameworks (Supporting Information Tables S6 and S7). Of these associations, 22 (41%) were between species of the same family. Out of 24 species, only two were not involved in a single significant nonrandom association: slough darter *Etheostoma gracile* (Girard 1859) and orangespotted sunfish *Lepomis humilis* (Girard 1858). The number of associations per species ranged from 0 to 8, with a mean of 4.45 associations. The number of nonrandom species associations was not significantly associated with the number of occurrences per species ($R^2 = 0.09$, $P = 0.16$) or rank occurrence ($R^2 = 0.12$, $P = 0.10$) via linear regression (Figure 3).

Most nonrandom pairs were aggregated (42 pairs, 78%) compared to segregated (12 pairs, 22%). Most of these pairs were hypothesized as governed by nonbiotic processes (28 pairs, 52%): habitat filtering (20 pairs, 37%), dispersal limitation (two pairs, 0.4%) or both (six pairs, 11%). The remaining 26 nonrandomly associated pairs (48%) had no

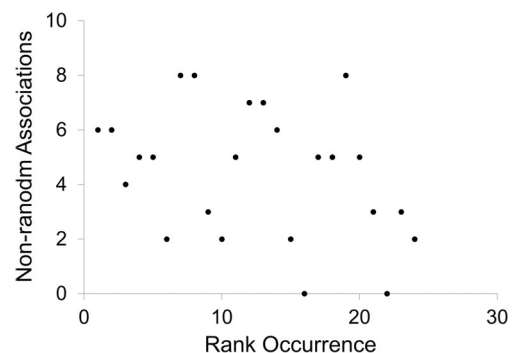


FIGURE 3 This scatterplot shows the relationship between the number of nonrandom associations and rank occurrence (based on number of occurrences across 66 samples) for each of the 24 species analysed here

Year	Framework	Total random	Total nonrandom	Total aggregated	Total segregated
1974	NMF	266 (94%)	10 (4%)	3	7
	PRF	241 (87%)	35 (13%)	31	4
2014	NMF	272 (99%)	4 (1%)	1	3
	PRF	260 (94%)	16 (6%)	15	1

TABLE 1 Summary of outcomes for species association modelling for two years (1974 and 2014) and two frameworks (null model framework and probabilistic framework)

Note: For both years and analyses, the majority of the 276 unique species combinations were not significantly different from random expectation (total random). Nonrandom pairs are broken down into total aggregated and total segregated.

TABLE 2 Significant species association that were recovered in at least two frameworks or sampling periods

No.	Species 1	Species 2	Hyp. Mech.	Null mod.	No. models	1974 NMF	1974 PRF	2014 NMF	2014 PRF
1	<i>Campostoma</i> spp.	<i>Etheostoma radiosum</i>	Biotic interaction	AGGR	3		X	X	X
2	<i>Cyprinella lutrensis</i>	<i>Cyprinella venusta</i>	Habitat filtering	SEGR	2	X	X		
3	<i>Cyprinella lutrensis</i>	<i>Pimephales vigilax</i>	Habitat filtering	AGGR	2		X		X
4	<i>Fundulus</i> spp.	<i>Micropterus punctulatus</i>	Spatial/habitat filtering	AGGR	2	X	X		
5	<i>Lepomis macrochirus</i>	<i>Lepomis microlophus</i>	Biotic interaction	AGGR	2		X		X
6	<i>Lythrurus umbratilis</i>	<i>Pimephales notatus</i>	Habitat filtering	AGGR	2		X		X
7	<i>Micropterus punctulatus</i>	<i>Notemigonus crysoleucas</i>	Biotic interaction	SEGR	2	X	X		
8	<i>Notropis boops</i>	<i>Notemigonus crysoleucas</i>	Spatial/habitat filtering	SEGR	2		X	X	
9	<i>Noturus nocturnus</i>	<i>Percina sciera</i>	Biotic interaction	AGGR	2	X	X		
10	<i>Percina sciera</i>	<i>Pimephales vigilax</i>	Habitat filtering	AGGR	2	X	X		

Note: Column headers indicate which two species are part of the interaction (species 1 and species 2), the hypothesized mechanism driving the association (hyp. mech.), the nature of the association, either aggregated or segregated (null mod.) and the number of models that returned this specific association as significant, four total models (no. models), and the remaining columns represent each model: null model 1974 (1974 NMF), probabilistic model 1974 (1974 PRF), null model 2014 (2014 NMF) and probabilistic model 2014 (2014 PRF).

detectable difference in the appropriate site-type categories via MANOVA, therefore the potential of biotic interaction was not refuted. For the 42 *aggregated* species pairs, 19 pairs (45%) were related to nonbiotic processes: habitat filtering (13 pairs, 31%), dispersal limitation (two pairs, 5%) or both (four pairs, 10%). The remaining 23 (55%) *aggregated* pairs were potentially driven by biotic interactions. For the 12 *segregated* species pairs, nine (75%) were driven by nonbiotic processes: habitat filtering (seven pairs; 58%), dispersal limitation (zero pairs, 0%) or both space and habitat (two pairs, 17%). Only three of 12 *segregated* pairs (25%) were potentially driven by biotic interaction.

Among the 54 significantly nonrandom species pairs recovered across analyses, only 10 were recovered in multiple scenarios, in either both frameworks and/or across sampling periods (Table 2). Only five of those pairs were found across both sampling periods. *Campostoma* spp. and orangebelly darter *Etheostoma radiosum* (Hubbs & Black 1941) were aggregated in both years with no discernable effect of dispersal limitation or habitat filtering. Red shiner *Cyprinella lutrensis* (Baird & Girard 1853) and bullhead minnow *Pimephales vigilax* (Baird & Girard 1853) were another aggregated pair found across years with an effect of habitat filtering found in both years. Bluegill sunfish *Lepomis macrochirus* Rafinesque 1819 and redear sunfish *Lepomis microlophus* (Günther 1859) were aggregated across times again without a signal of spatial or environmental effect. Redfin shiner *Lythrurus umbratilis* (Girard 1856) and bluntnose minnow *Pimephales notatus* (Rafinesque 1820) were yet another aggregated pair with an effect of habitat filtering found in 1974 but not in 2014. Finally, bigeye shiner *Notropis boops* Gilbert 1884 and golden shiner *Notemigonus crysoleucas* (Mitchill 1814) were the only segregated pair found consistently across time periods and this association is likely due to spatially autocorrelated habitat filtering.

4 | DISCUSSION

4.1 | Species co-occurrence

The results point to many associations among fish species in the Muddy Boggy River drainage. The frameworks used herein were able to identify 54 unique, significantly nonrandom associations from incidence data that contained 276 pairwise comparisons. However, the proportion of neutral associations (not different from random) was high across both time periods and frameworks. This is typical for studies of this kind (Lopes *et al.*, 2015; Lyons *et al.*, 2016), but see Cordero and Jackson (2019). Some authors have argued the large proportion of random associations points to the importance of stochasticity in structuring assemblages (Devercelli *et al.*, 2016). While stochasticity may play an important role, especially at smaller spatial scales (Zbinden, 2020), there is also evidence for deterministic processes, such as environmental filtering, structuring fish assemblages (Cordero & Jackson, 2019; Zbinden & Matthews, 2017). Some caution is warranted in rushing to conclusions regarding stochasticity (Rahel *et al.*, 1984) from analyses of this kind because of alternative explanations which are not explicitly tested. First, by considering only presence-absence data, information is lost and there is potential for associations to go undetected. If, for example, two species sometimes co-occur (weak signal), but when they do one of the species is invariably less abundant than when found without the other, that evidence might suggest a competitive interaction. However, this information is lost when considering only incidence. Second, pairwise comparisons have limited power to detect associations involving rare species due to low statistical power (Lavender *et al.*, 2019; Veech, 2013), which are often a large portion of diversity in a region (Lavoie *et al.*, 2009; Magurran & Henderson, 2003). Third, sample sizes typical of most biodiversity surveys are likely not large enough to reliably capture all

of the associations that may be present, with some estimates suggesting upwards of 500 or more samples may be necessary to do so (Blanchet *et al.*, 2020). So, while stochasticity is an important mechanism (Ellwood *et al.*, 2009; Vellend *et al.*, 2014), we must not neglect that methods that test co-occurrence are likely to result in many ‘random’ associations simply due to their limitations independent of stochasticity. Importantly, we should not neglect that an ‘assemblage’ (*i.e.*, fish alone) is only a part of the larger community (*e.g.*, macroinvertebrates, plants, etc.) in which interactions could be observed.

4.2 | What about mechanisms?

The decision tree framework for disentangling likely mechanisms that underly associations is constructed so that biotic interaction is essentially the null hypothesis. Unfortunately, that means biotic interaction can be refuted and alternatives supported, but in the absence of a signal from either spatial or environmental factors then the *de facto* explanation for a significant association is biotic interaction. Therefore, I emphasize that this inference should be viewed as hypothesis generation rather than hypothesis testing *per se*. Species associations that remain plausibly driven by biotic interaction should be judged by those conducting the analyses to determine whether the categorization makes ecological sense and warrants further investigation. It remains possible that unmeasured environmental factors could be involved in any of the associations identified. Additionally, the outcomes from the framework are not mutually exclusive (Blois *et al.*, 2014), and the expected patterns from processes such as competition and predation may not always be straightforward (Blanchet *et al.*, 2020; Cordero & Jackson, 2019).

Co-occurrence methods were largely developed with classic species interactions in mind (*e.g.*, competition and predation). However, the results here demonstrate that most significant associations, gleaned from incidence data gathered across this study drainage, could be explained by nonbiotic mechanisms, namely habitat filtering, but also dispersal limitation. It is intuitive that species which differ in resource requirements (*i.e.*, niche divergence) are less likely to be found in the same stream reaches, while those with similar requirements (*i.e.*, niche convergence) would more often co-occur. Indeed spatio-environmental filtering (Cordero & Jackson, 2019; Peres-Neto, 2004; Zbinden & Matthews, 2017) is a well-supported hypothesis of metacommunity theory (Erős, 2017; Leibold *et al.*, 2004). While habitat filtering is interesting in and of itself, if one is specifically seeking biotic interactions, then these frameworks can reveal which associations are likely due to alternative explanations instead.

No species were inferred to be segregated due to spatial factors alone (*i.e.*, dispersal limitation); perhaps the relatively small spatial scale of this study is not large enough to truly segregate species by distance (Heino *et al.*, 2015). Dispersal limitation was indicated, however, in the aggregation among *Camptostoma* spp. and *Pimephales notatus* and among *E. radiosum* and freckled madtom *Noturus*

nocturnus Jordan & Gilbert 1886. This suggests a pattern of nested, not entirely overlapping, distributions among species within the pair. Six interactions showed significant signal of both spatial and environmental factors in structuring the association and likely indicates the effect of spatially autocorrelated habitat filtering (Gotelli & Ulrich, 2012; Mykrä *et al.*, 2007). In other words, the environmental factors associated with the species interactions were themselves spatially structured.

Only a small subset of the associations found here were segregations that remained plausibly driven by negative biotic interactions which would fit the prediction for pairs of competitors or predator/prey, although predator/prey relationships could result in aggregations as well (Cordero & Jackson, 2019), especially at the spatial scale analysed here (Hoeinghaus *et al.*, 2007). These pairs were spotted bass *Micropterus punctulatus* (Rafinesque 1819) and *Notemigonus crysoleucas*, *E. radiosum* and white crappie *Pomoxis annularis* Rafinesque 1818, and *Lythrurus umbratilis* and *Notemigonus crysoleucas*. Both *M. punctulatus* and *P. annularis* are large centrarchid predators which could easily consume the other member of their respective pair. *N. crysoleucas* is a common forage fish consumed by many fish predators (Webber & Haines, 2003) and *E. radiosum* is a small benthic percid, although the only explicit test of predation on *E. radiosum* showed a lack of piscine predation (Scalet, 1974). *L. umbratilis* and *N. crysoleucas* are the only association that seems plausibly driven by competition. Another study in Oklahoma also found no co-occurrence among these two species at localities along a small creek (Stewart *et al.*, 1992). Both species occur in similar habitats and have similar diets and feeding modes (Robison & Buchanan, 2020), and so competition seems plausible. It is of note that of the 12 segregated associations recovered here, *N. crysoleucas* was a member of six of them. This species is highly tolerant of temperature and turbidity extremes, making it a good competitor (Robison & Buchanan, 2020). Overall, very little evidence of competition among the fish species analysed in this drainage was found, although it should be noted that only strongly competing species are likely to result in strong and significant segregated patterns. Furthermore, others have found competing species to sometimes aggregate rather than segregate, although this was at a much larger spatial scale (Cordero & Jackson, 2019).

A large portion of nonrandom aggregations (23 pairs) showed no discernable effect of spatial or environmental factors given the scale and environmental characteristics considered in this study. Therefore, the possibility of positive, or facilitative, interactions, but also associations due to predator prey interactions, driving these associations remains but should be scrutinized. Seven of these 23 pairs included two members of the sunfish and black bass family Centrarchidae, within which individuals display niche flexibility based on size that often leads to what is referred to as ‘niche packing’ or ‘species packing’ (Desselle *et al.*, 1978; Werner & Hall, 1976; Werner, 1977). In effect, these species forage on similar resources, but diverge on the specific size of the resources targeted. Target prey size is normally correlated with predator size and mouth morphology, and this allows for stable coexistence of predators. However, species packing is not mutually exclusive from facilitation, and there is evidence that

piscivorous predators using different strategies can benefit from one another (Eklöv & VanKooten, 2001; Schumann *et al.*, 2020).

An additional seven nonrandom pairs of those showing no discernable effect of spatial or environmental factors consisted of just one member of the Centrarchidae family plus a nonsunfish member (the other member being a minnow Leuciscidae in five of those cases). This may be explained, in part, by spawning nest associations (Johnson & Page, 1992; Peoples & Frimpong, 2016; Wallin, 1989). Members of the sunfish family make large nests by clearing depressions in stream-bottom substrate, which are often used by other species (Johnston, 1994a; Ross, 2013), and benefit from the protective parental care of the nest-builder and not necessarily the physical environment of the nest (Johnston, 1994b). Examples exist of minnow species parasitizing centrarchid nests, sometimes even predating largely on the hosts eggs (Fletcher, 1993) or otherwise lowering host fitness through fungal infection of its brood (Shao, 1997). All species in this group of seven pairs show strong overlap in spawning season (April–August in this region), and two of the species have been reported laying eggs in sunfish nests: *Cyprinella lutrensis* and *Lythrurus umbratilis* (Robison & Buchanan, 2020 and references therein). Any of these pairs not engaged in nest association could possibly represent a predator–prey interaction between the larger Centrarchid predator and smaller prey species (Hoeinghaus *et al.*, 2007). Additional data would be required to parse these alternative possibilities, along with the potential effect of unmeasured environmental factors.

Campostoma spp. and *Etheostoma radiosum* is an interesting association recovered here because these species were significantly aggregated in three of four models across both sampling periods (indicating a reliable association), and yet could never be associated with dispersal limitation or habitat filtering. *Campostoma* spp. occurred in 16/33 samples (1974) and 22/33 samples (2014). *E. radiosum* occurred in 19/33 samples (1974) and 21/33 samples (2014). These two co-occurred together in 13 and 18 samples in 1974 and 2014, respectively. This result suggests a facilitative relationship between *Campostoma* spp. and *E. radiosum* may exist. *Campostoma* spp. are benthic, algivorous minnows that forage in shoals of up to 1,000 individuals (Matthews *et al.*, 1987) and can have important effects on other trophic levels within their communities, such as producers and decomposers (Power *et al.*, 1985; Veach *et al.*, 2018). Grazing fish like *Campostoma* can also promote the downstream drift of macroinvertebrates by disturbing substrate and causing these organisms to enter the water column. Matthews *et al.* (1987) discussed the potential for *Campostoma* to have positive interactions on other fish by improving feeding opportunities for macroinvertebrate predators through substrate disturbance and provided anecdotal evidence of an association between dusky stripe shiner *Luxilus pilsbryi* (Fowler 1904) and *Campostoma* that often occurred in the same pools together, as observed during snorkelling surveys. A similar association has been reported between *Campostoma anomalum* and hornyhead chub *Nocomis biguttatus* (Kirtland 1840) (Gorman, 1988). Various suckers Catostomidae are known feeding facilitators that have a similar feeding mode that involves disturbing substrate (Pflieger, 1975; Reighard, 1920) as well as various other species of freshwater fish (Arnhold *et al.*, 2019; Ross, 2013). Therefore, it

is possible that large shoals of foraging *Campostoma* disturb substrate and force more macroinvertebrates into the water column which then drift downstream into riffles occupied by darters, like *E. radiosum*, that benefit from higher abundance of food. This is an intriguing hypothesis that deserves future consideration, as the importance of large populations of algivorous grazers is appreciated by fish ecologists (Matthews *et al.*, 1987), but perhaps not fully understood.

4.3 | Inconsistency through time

Similarity in the species associations recovered across sampling campaigns was low (only five consistent associations). Species interactions may be expected to change over time due to both species turnover and ‘rewiring’ of interactions (Brimacombe *et al.*, 2020). Species may associate only transiently (Bart, 1989; Grossman & Ratajczak, 1998), seasonally (Gido & Matthews, 2000; Matthews & Hill, 1980) or over long periods of time (Matthews & Marsh-Matthews, 2016; Matthews & Matthews, 2002; Ross *et al.*, 1985). Species associations may shift over time due to many nonmutually exclusive factors, including environmental change, population movement and demographic shifts, and local extinctions, all of which are all scale-dependent processes (Blanchet *et al.*, 2020; Chafin *et al.*, 2019; Harmon *et al.*, 2009).

A previous comparison of fish assemblage structure in the Muddy Boggy River drainage over the same period measured high temporal turnover at the local spatial scale (*i.e.*, sampling locality) (Zbinden, 2020). This and the similarity in season among the two sampling campaigns might suggest the importance of turnover as opposed to interaction rewiring (Brimacombe *et al.*, 2020) for explaining the low similarity in species association through time. As Lavender *et al.* (2019) showed, changes in species incidence can affect the ability of these tests to determine nonrandom interactions. These authors cautioned against comparing tests among sampling times or experimental treatments because of this reason. I take this point; however, I would argue not against comparison, but for more interest in similarities rather than differences. For the study presented here, many dissimilarities were found between sampling times and some of these may be due to changes in local incidences over time, among a host of other factors. However, nonrandom associations that persist through time (as discussed above) provide stronger and more reliable evidence for two species associating.

4.4 | Complimentary frameworks

The two co-occurrence frameworks used here appear to complement each other well because each provided unique associations not identified by the other. The PRF and NMF both aim to identify non-randomly associated species pairs, but they differ in statistical approach. Null model approaches with Bayes corrections had one of the lowest Type I error rates among co-occurrence methods, but consistently failed to detect known associations from simulated data in a study of co-occurrence methods (Lavender *et al.*, 2019). The PRF

model (Veech, 2013) used here actually performed best overall with regard to both error rates in that same comparison (Lavender *et al.*, 2019). Therefore, it was expected *a priori* that the NMF would result in fewer significant associations compared to the PRF, and indeed this was the case. Lavender *et al.* (2019) also found that negative (segregated) associations had lower Type I and higher Type II error rates. Generally, null methods seem to be more robust tests of negative associations, while probabilistic methods are more robust tests of positive associations. The findings here are also consistent with this result as there were many more aggregated than segregated pairs.

Despite low Type I error, the methods used here must be interpreted with caution. For example, this study analysed 276 unique species pairs and given an error rate of ~ 0.01 , it is possible that approximately three or fewer of the significantly nonrandom pairs were false positives. Future studies could also reduce error rate by only analysing species hypothesized *a priori* to have an association (e.g. *Campostoma* spp. and *Etheostoma* spp.), or by reducing the set to include only certain species of interest (e.g., species of conservation concern, species within families or known predator/prey species).

4.5 | Conclusion

Different species can associate or interact in many ways, and methods exist for inferring associations and underlying mechanisms from incidence data (e.g., co-occurrence frameworks). However, these methods have received much criticism despite their recent resurgence in the literature. Co-occurrence frameworks for identifying nonrandomly associated species pairs (e.g., aggregated or segregated pairs) should be used as heuristic tools for sharpening hypotheses concerning fish ecology (Silknetter *et al.*, 2020). The data presented here demonstrate that at least some species associations can be found in even a small number of samples. Roughly half of these associations were driven by nonbiotic processes (e.g., habitat filtering) and most associations did not show a consistent signal through time.

All the significant associations from this study can be explored further. Out of 276 possible associations, there were 54 that showed a nonrandom pattern. These can further be prioritized based on the number of models or times they were significant. These frameworks can provide insight into the broader mechanisms of species assembly and point to potentially interesting species interactions (out of many possible interactions). Rather than a weakness, the heuristic nature is the strength of such methods and can help guide biologists toward better questions by employing relatively cheap diversity survey data, which are often already in hand, to reduce complex interaction networks down to their nonstochastic parts. This can save fisheries biologists time and resources and help guide us toward species associations that warrant further investigation. In turn, this can lead to a fuller understanding of how fish assemblages are structured.

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DATA AVAILABILITY STATEMENT

All data and R code available in the supplementary material.

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