

Inferring social network structure in ecological systems from spatiotemporal data streams

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We propose a methodology for extracting social network structure from spatio-temporal datasets that describe timestamped occurrences of individuals. Our approach identifies temporal regions of dense agent activity and links are drawn between individuals based on their cooccurrences across these 'gathering events'. The statistical significance of these connections is then tested against an appropriate null model. Such a framework allows us to exploit the wealth of analytical and computational tools of network analysis in settings where the underlying connectivity pattern between interacting agents (commonly termed the *adjacency matrix*) is not given a *priori*. We perform experiments on two large-scale datasets (greater than 10^6 points) of great tit *Parus major* wild bird foraging records and illustrate the use of this approach by examining the temporal dynamics of pairing behaviour, a process that was previously very hard to observe. We show that established pair bonds are maintained continuously, whereas new pair bonds form at variable times before breeding, but are characterized by a rapid development of network proximity. The method proposed here is general, and can be applied to any system with information about the temporal co-occurrence of interacting agents.

> Keywords: network analysis; spatio-temporal data streams; animal social networks

1. INTRODUCTION

We use the terms graph or network to describe the simplified version of the pattern of interactions in a system, such as an animal population, where nodes are individual entities and edges represent some form of association, interaction, similarity or behavioural correlation between nodes. In the same way that a map is a simplified (though useful) version of a landscape, a network describes the *topology* of a real-world system by focusing on the connectivity patterns of its individual components [1].

The key motivation for employing network analysis tools is that the web of interconnections between individuals can provide insights into the underlying mechanisms that govern the system under study [2]. For example, in an ecological context, the position and role of animals in the network may have important fitness consequences [3] both for the individual and the population as a whole [4]. Additionally, the network paradigm gives us the flexibility to look at the system at various resolutions and model any type of interaction; sexual, cooperative, competitive, etc [4].

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Despite the advantages of the network paradigm and the wealth of analytical and computational tools for network analysis [5-8], the problem of capturing any given system as a graph is not always trivial. Not all systems possess an obvious 'web-like' structure (such as the Internet), where the interconnections between participating entities are apparent from direct observation (computers that are connected through physical cables). Additionally, collected data (from field studies, sensor observations, World Wide Web, etc.) may not capture the associations between the observed agents, thus no relational structure can be directly defined. For example, in systems, such as animal populations, the underlying network of social affiliations needs to be inferred through proxies such as the behaviour (mobility patterns, foraging habits, etc.) of individual animals.

This work focuses on the problem of finding the underlying social network structure of a population that can only be observed through the spatial trajectories of its individual members. We use as a case study a setting where individual wild birds are marked with transponder devices and through appropriate logging hardware we are able to identify their position at various sites in their natural habitat. The observation data collected in this manner consist of a long stream of timestamped records, where no obvious interaction or social affiliation is apparent. By assuming that *social structure* is a latent factor that affects the way birds visit locations (in the sense that socially affiliated individuals have similar mobility patterns), we propose a methodology that extracts a *social network* from such a spatio-temporal data stream. Although we demonstrate our method in an ecological context, our approach can be generalized to any setting where agents perform timestamped 'check-ins' at various locations.

The paper is organized as follows. In §2, we outline our experiment settings and discuss our data format. In §3, we present our contribution, which is a methodology for extracting network structure from timestamped observation data. In §4, we apply our method to the wild bird dataset and show that the extracted networks reflect actual processes that take place in the population, by focusing on mating pair formation. We conclude this paper in §5 by discussing the next steps of our research, both in terms of method development and data collection extensions. The Matlab code that implements the methods presented in the paper is made available¹ to the community.

2. DATA COLLECTION

This work lies within the context of a large ongoing study of the great tit *Parus major* population at Wytham Woods near Oxford, UK. Thousands of individual birds are marked with transponders and a grid of sensor-enabled locations generates hundreds of thousands of records each winter. At each one of the 67 locations in the forest, there is a feeder that acts as an attraction point for foraging individuals. By placing appropriate logging hardware at the feeder, we are able to record the presence of each individual bird. Owing to equipment constraints, there were only 16 loggers available at any time, and these were thus rotated around the 67 locations following a structured randomized design, so that each of eight approximately equally sized sections of the site always had two active loggers in it. More details on our experiment set-up is provided in the electronic supplementary material.

The data generated from this scheme consist of a long stream of timestamped observations as shown in table 1. Each row represents a single record that captures the ID of the bird along with the time and location where the foraging event took place. In this format, shown in table 1, our data stream is only a transactions table in a relational database context, which restricts our analysis to a handful of relatively simple counting operations such as finding the total appearances of a given bird, total birds that visited a specific feeder, etc.

What we are interested in is to find an appropriate mapping of this spatio-temporal stream to a relational space, where social affiliations between individuals are revealed by the similarity of their feeder visitation patterns. We seek to characterize the overall social network of the population of marked birds and explore the ability of this approach to recover relationships between mated pairs of individuals observed independently

¹See http://www.robots.ox.ac.uk/~parg/software.html.

Table 1. Sample format of our data.

bird ID	timestamp	location ID
N199642 TE80535 V260952 V260952 N199642	1/9/2007 10.02:15 (am) 1/9/2007 10.02:30 (am) 1/9/2007 10.02:30 (am) 1/9/2007 10.02:45 (am) 1/9/2007 10.12:15 (am)	1a 1a 2b 2b 1c
	(all)	

during breeding season data collection. We further wish to explore the temporal dynamics of the formation of mated pairs. In biological terms, the process by which pairs of individuals develop relationships that lead to mating is poorly understood in most natural populations, since the majority of work involves observations of pairs at the time of breeding, after pair formation has occurred. As a consequence, we have little knowledge of when such relationships form, and when they become distinguishable from other social relationships between individuals.

In $\S3$, we introduce a method, based on the above goals, that extracts network structure given such spatio-temporal data. In \$4, we present the application of this approach to the *P. major* dataset.

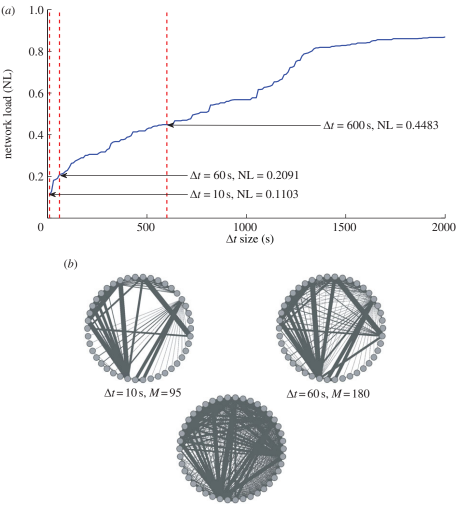
3. NETWORK INFERENCE FROM SPATIO-TEMPORAL DATA

3.1. The time-window problem

A typical approach for building a network from data such as those presented in §2 would involve discretizing the stream using a fixed aggregation or time window Δt and assuming that if two individuals are recorded within an interval Δt then there is a link between them in the network [9–13]. The most obvious problem with this approach is that of finding the appropriate size for the time window. An inappropriately small Δt may lead to a network that does not capture important connections, while a very large Δt would overload the graph with 'junk' links.

Using our wild-bird data as an example, we take a single day's worth of observations (in a format similar to the one shown in table 1) and split that stream into time intervals of size Δt . We then place links between the N individual birds (nodes) based on the number of times they were recorded within a temporal distance of Δt . We seek to examine the changes that take place in the network as we vary the time window size by monitoring the *network load* (NL), which is the fraction of M links in the network over all possible pair combinations $\frac{1}{2}(N^2-N)$ of N nodes.² We can see in figure 1a that NL increases along with the size of Δt , because more links are placed between nodes. An example of how network topology changes for various selections of time window size is shown in figure 1b, while Krings *et al.* [13] have performed similar experimentation considering

²In this example and throughout this paper we are considering networks that are *undirected* with nodes that have no self-edges.



 $\Delta t = 600 \, \text{s}, M = 978$

Figure 1. (a) We plot the network load for various time window sizes, spanning from 10 s to half an hour. We can see that especially for early increases of Δt , there is a large inclusion of links in the network. We also mark three cases of different time window sizes (dashed vertical line) and show in (b) how the graph topology changes based on the Δt value. (Online version in colour.)

more network metrics such as average degree, average weight, clustering coefficients, etc.

Between all these different network topologies that result from varying Δt , there is no direct way of showing which one is the most appropriate. Additionally, even if we had some prior knowledge on the appropriate time window size or even a specific quality function for finding its optimum value, we have still made the strong assumption that Δt is *fixed* throughout the data stream. This corresponds to the belief that the 'interaction radius' between individuals is *constant* across our observation period and is not affected by temporal changes in the overall system.

In the current work, we shall pursue a different approach for building networks from spatio-temporal records, which exploits the inhomogeneous *density profile* of our data stream thus avoiding schemes such as multiple runs [13] in order to select an appropriate Δt . This methodology, which we will call GMMEvents (Gaussian mixture model for event streams) is complemented with an appropriate null model that allows us to distinguish between links that denote social tie and the ones that result from coincidence.

3.2. Identification of gathering events

Let our spatio-temporal data \mathcal{D} , a sample of which we showed in table 1, be represented in the form $\mathcal{D} = \{b_z, t_z, \ell_z\}_{z=1}^Z$, where Z is the total number of records or *tuples* in our database (e.g. the number of rows of table 1). If we take a single tuple $\{b_z, t_z, \ell_z\}$, we read it as 'the bird b_z appeared at time t_z at the feeding location ℓ_z '. Note that $\{t_z\}_{z=1}^Z$ denotes event time; therefore, for every timestamp t_z , there is a corresponding bird appearance b_z . Additionally, given a specific bird *i* out of total N birds, there can be many records z for which $b_z = i$, as a single individual may appear many times in the data. Our goal is to find an appropriate mapping from the stream \mathcal{D} to an *adjacency matrix* $\mathbf{A} \in \mathbb{R}^{N \times N}$, where $a_{ij} \neq 0$ denotes a link between birds *i* and *j*. To keep the notation uncluttered, from now on we will focus on the case of a single location and show

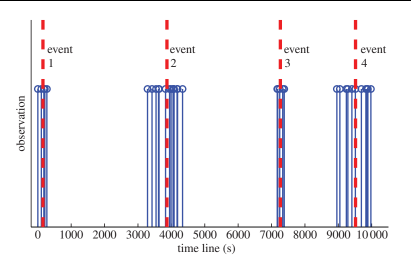


Figure 2. We plot bird arrivals as recorded at a specific location over the course of 3 h period. We can see that the visitation profile is temporally focused, consisting of bursts of bird activity. Our goal is to identify such regions of increased observation density and examine which individuals participate in these gathering events. (Online version in colour.)

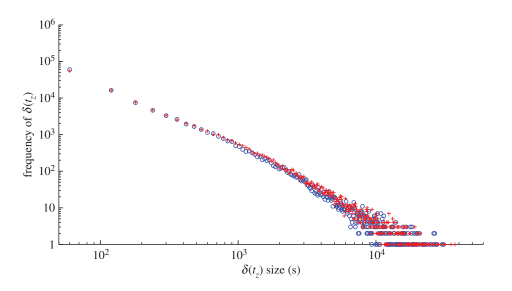


Figure 3. We calculate the time difference $\delta(t_z) = t_z - t_{z-1}$ between every pair of consecutive observations at each location in our two data streams (seasons 2007–2008 and 2008–2009) and plot the histogram of those values on a logarithmic scale. The $\delta(t_z)$ that refer to pairs where z - 1 is the last observation of day d - 1 and z the first observation of day d have been omitted, in order to avoid bias in the results (there is no bird feeding activity during night-time). Open circles, dataset 2007–2008; plus symbols, dataset 2008–2009. (Online version in colour.)

later that results can be easily generalized to the multi-site case.

Consider the plot of figure 2, which illustrates how bird arrivals at a particular feeding location are spread throughout a small sample of our observation timeline. Each stem represents an actual sensor capture of a specific bird b_z at time t_z . We can see that the records are not uniformly spread across time, but they are 'packed' in small observation-dense regions. Indeed, if we take the whole data stream and extract the histogram of the time differences $\delta(t_z) = t_z - t_{z-1}$ between every pair of consecutive observations, as seen in figure 3, we find a broad power-law tail with exponent $\simeq 2.5$ for $\delta(t_z) > 800$. This non-Poissonian decay of inter-record timestamps, along with the fact that most $\delta(t_z)$ take small values, implies that the observation profile comprises temporally focused bursts of recording activity, which can be seen as *flocks of foraging individuals*.

Our main hypothesis is that birds not only visit the feeder as part of such small flocks but also have a *pre-ference* to the members of the flock they choose to forage with. Such regions of increased observation density can be viewed as K gathering events of socially affiliated birds. We seek to cluster our Z observations in a way such that closely appearing individuals, based on their arrival timestamp t_z , are assigned to the same gathering event k.

We perform this clustering scheme using a Gaussian mixture model, with an appropriate configuration that allows us to automatically infer the effective number K of events/clusters (see the electronic supplementary material). The result is described by an observation-to-cluster responsibility matrix $\Gamma \in \mathbb{R}^{Z \times K}$, where Z is

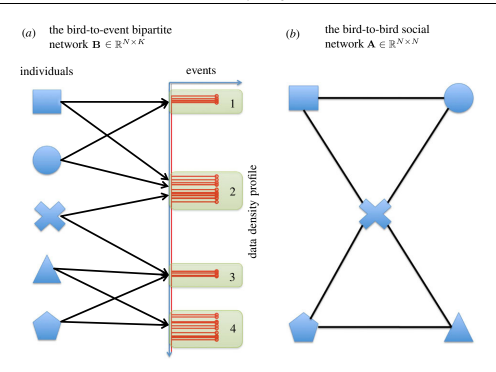


Figure 4. Our method identifies gathering events from the bursts in our observation stream as seen in (a). Then individuals are assigned to such events creating a bipartite network. In part (b), we recover the bird-to-bird social network, via an appropriate one-mode projection, based on the co-participation of individuals to these events. (Online version in colour.)

the total number of bird observations, K the number of clusters and the elements $\{\gamma_{z1}, \gamma_{z2}, \ldots, \gamma_{zK}\}$ of each row denote a membership score of a single observation z to an event k.

As there is a one-to-many correspondence between a given bird i and timestamps t_z , a single bird can be recorded many times in the observation stream or, in other words, there are many tuples $\{t_z, b_z\}$ for which $b_z = i$. Therefore, we seek to map the observation-to-cluster matrix $\Gamma \in \mathbb{R}^{Z \times K}$ to a bird-to-cluster matrix $\mathbf{B} \in \mathbb{R}^{N \times K}$. We start by taking each row $\boldsymbol{\gamma}_z = \{\boldsymbol{\gamma}_{zk}\}_{k=1}^K$ of $\boldsymbol{\Gamma}$ and set the largest element to 1 and all the others to zero. This allows us to describe participation scores γ_{zk} , and all the other measures we derive from them, as integer-valued occurrences. For each individual bird $i \in \{1, \ldots, N\}$, we identify the subset \mathcal{Z}_i of rows γ_z of $\boldsymbol{\Gamma}$ that correspond to observations regarding *i*. We thus set each row \mathbf{b}_i of \mathbf{B} as the sum $\mathbf{b}_i = \sum_{z \in \mathcal{Z}_i} \gamma_z$. The resulting matrix $\mathbf{B} \in \mathbb{R}^{N \times K}$ can be seen as a representation of a bipartite or two-mode network that is a graph with two types of nodes; N birds and K events, as shown in figure 4a. Each element b_{ik} denotes the number of times each bird was observed at a specific foraging group.

3.3. Building the social network

The bipartite network we extracted in §3*b* and shown in figure 4*a* describes the event participation structure of the bird population, which is the weighted allocation of *N* birds to *K* foraging events, encoded by $\mathbf{B} \in \mathbb{R}^{N \times K}$. Although this finding is important by itself, as it allows us to quantify the structure of such small foraging groups in terms of the number, individual characteristics, relatedness of their members etc, we seek to move one step further and extract the bird-to-bird *social network* based on the mutual participation of individuals to such events.

Therefore, we seek to define an appropriate one-mode projection $\mathbf{B} \in \mathbb{R}^{N \times K} \to \mathbf{A} \in \mathbb{R}^{N \times N}$, shown in figure 4, so that a link a_{ij} between a pair *i*, *j* in the resulting network will express how strongly the two birds forage together. We start by defining *co-occurrence* of individuals *i* and *j* as the number of times they were recorded in the same foraging group. Thus, given the event membership profiles \mathbf{b}_i and \mathbf{b}_j for *i* and *j*, respectively, we define the total co-occurrences a_{ij} as $a_{ij} = \sum_{k=1}^{K} \min(b_{ik}, b_{jk})$, where *K* is the number of foraging groups and a_{ij} is effectively the *link weight* between *i* and *j* in the resulting social network described by the adjacency matrix $\mathbf{A} \in \mathbb{R}^{N \times N}$. Note that other association indices, such as the ones presented in Ginsberg & Young [14] can be used depending on the problem context.

3.4. Co-occurrences: social tie versus coincidence

The next issue we seek to address is the statistical significance of the extracted link weights. Building the adjacency matrix $\mathbf{A} \in \mathbb{R}^{N \times N}$ in the manner described in §3*a* makes the very strong assumption that if two individuals participate in the same gathering event, they have some form of social affiliation. This assumption, known in the animal social network literature as the *Gambit of the Group* (GoG) [15], may lead us to adjacency matrices encumbered with 'junk' links, produced by co-occurrences that happened by chance. Such coincidences are also frequent in settings where there are natural peak-hours in the data collection period and also when the sensor hardware act as attraction points, as, for example, the bird feeders in our study. Hence, we seek to define an appropriate *null*

model that describes how 'statistically surprising' a given link weight would be, if there was no underlying social preference in the foraging habits of the bird population. From previous sections, we have discussed that observations occur in bursts (as shown in figure 2) that denote small foraging groups of birds that arrive together at the feeders. This is captured by the birdto-event matrix $\mathbf{B} \in \mathbb{R}^{N \times K}$, where each element b_{ik} in the row vector \mathbf{b}_i denotes the number of times bird *i* appeared at the gathering event k.

Consider each row vector \mathbf{b}_i as a draw from a multinomial distribution $\mathcal{M}(n_i, \mathbf{p}_i)$, with parameters $n_i = \sum_{k=1}^{K} b_{ik}$ and $p_{ik} = b_{ik}/n_i$. The values of the parameter vector $\{p_{ik}\}_{k=1}^{K}$ can be viewed as a *preference profile* of a bird i to each foraging event k. If our hypothesis that social affiliation between birds affects event membership holds, then closely interacting birds i, j will have similar preference profiles \mathbf{p}_i and \mathbf{p}_i .

Let us now propose an element shuffling σ of \mathbf{p}_i so that $\mathbf{p}_i \to \sigma(\mathbf{p}_i)$ and draw a new event occurrence vector $\mathbf{b}_{i}^{(0)}$ from the multinomial distribution $\mathcal{M}(n_{i}, \sigma(\mathbf{p}_{i}))$. Performing this permutation and sampling scheme independently for all birds $i \in \{1, ..., N\}$ leads to a new bipartite network described bird-to-event by $\mathbf{B}^{(0)} \in \mathbb{R}^{N \times K}$. This new matrix $\mathbf{B}^{(0)}$ preserves many key characteristics of the original data, among them the event membership structure, because bird appearances remain concentrated in K regions of increased observation density. Quantities such as the number of individuals N, and the total records n_i , of bird i in the data are also retained.

The key difference introduced in $\mathbf{B}^{(0)}$ is that, although a bird's uneven participation preference \mathbf{p}_i across foraging groups is preserved (as the permuted $\sigma(\mathbf{p}_i)$ has the same entropy as \mathbf{p}_i), the shuffling σ 'breaks' all correlations between \mathbf{b}_i and \mathbf{b}_j induced by latent social affiliation between individuals i and j. In other words, under our null model, birds still forage in small groups, but with no social preference to which other members of the group they will forage with. We repeat this process R-times and for each generated bird-to-event matrix $\mathbf{B}^{(0)}$ we extract the birdto-bird matrix $\mathbf{A}^{(0)}$ using the same one-mode projection presented in $\S 3c$. By generating multiple instances of $\mathbf{A}^{(0)}$ in this manner, we are effectively drawing samples from the *ensemble* or family of graphs $\mathcal{G}^{(0)}$ that contains all possible network configurations generated by the null model. Our goal is to examine if our observed network **A** is an unlikely case of $\mathcal{G}^{(0)}$.

The randomization process generates R values of the weight of each link between i and j. From the histogram, we get the empirical distribution $P(a_{ii}|H_0)$ that denotes the probability of having a link of weight a_{ij} given that the null hypothesis H_0 holds. We examine how statistically surprising is each observed link a_{ij} by performing a hypothesis test, given an appropriate significance level α , by examining the likelihood $p = P(x \ge a_{ij}|H_0)$ of co-occurrences as large as a_{ij} . Note that the key point of a null model is that co-occurrences happen between individuals, but not as a result of an underlying social structure. In other words, the links in $\mathbf{A}^{(0)}$ are independent under H_0 , hence $P(\mathbf{A}|H_0) = \prod_{ij} P(a_{ij}|H_0)$.

Thus, our significance test lies in examining how well this independence assumption can explain the observed co-occurrences encoded in each link of A.

3.5. Integrating information from multiple locations

We briefly expand on our graph inference scheme to the multi-location setting. For each record $\{t_z, b_z, \ell_z\}$ in our data stream, we now have an additional term $\ell_z \in \{1, \ldots, L\}$ that denotes the index of the location where observation z took place.

We start by segmenting our data $\mathcal{D} = \{t_z, b_z, \ell_z\}_{z=1}^Z$ into L streams, so that each $\mathcal{D}^{(\ell)}$ contains records referring only to location ℓ . For each $\mathcal{D}^{(\ell)}$, we perform the network extraction process as presented in §3b, c leading to L adjacency matrices $\mathbf{A}^{(\ell)} \in \mathbb{R}^{N_{\ell} \times N_{\ell}}$, where $N_{\ell} \leq N$, the subset of birds recorded at location ℓ . Significance tests, as described in $\S 3d$, are performed independently for each ℓ , in order to preserve the unique visitation and location load statistics of each site.

Each matrix $\mathbf{A}^{(\ell)} \in \mathbb{R}^{N_{\ell} \times N_{\ell}}$ generated in this scheme captures a subset of the overall connectivity profile in the population. As the interpretation of link weight is the number of co-occurrences between two individuals, the overall a_{ij} is simply the summation $a_{ij} = \sum_{\ell=1}^{L} a_{ij}^{(\ell)}$ over multiple sites. In §4, we will demonstrate how these methodologies

are applied to the wild-bird dataset described in $\S 2$.

4. RESULTS

4.1. Application on the wild-bird dataset

We apply GMMEvents on the dataset of wild-bird foraging records presented in §2. Our observations consist of two main streams: $\mathcal{D}_{7,8}$ that covers the activity of $N_{7,8} = 770$ birds from August 2007 to March 2008 and $\mathcal{D}_{8,9}$ that spans from August 2008 to March 2009 and contains $N_{8,9} = 753$ birds.

Instead of applying our method to the whole twoseason data stream directly, we start by breaking it down into 24 h segments. Our aim is to produce a collection of network snapshots that would allow us to study the day-by-day changes in the population's sociality. An example of the observation data is shown in figure 5a, where we can see the isolated observation-rich regions (blue stem lines) that refer to each particular day. Note that the night period (no-observation zones in between days) acts as a natural separator in our data stream, as no bird foraging activity takes place during that time.

We proceed by breaking down each daily segment of our data into sub-streams that correspond to Ldifferent feeding locations, shown in figure 5b for 9 September 2007. We then apply GMMEvents to each location ℓ separately, as co-occurrences need to be defined both in terms of temporal and spatial proximity. On each one of those feeder-specific streams for that day, our method identifies bursts in the observation density profile and builds a bipartite network $\mathbf{B}^{(\ell)}$ between birds and gathering events, as shown in figure 5*c*. The weight of each link $b_{ik}^{(\ell)}$ denotes the number of times bird i appeared in the gathering

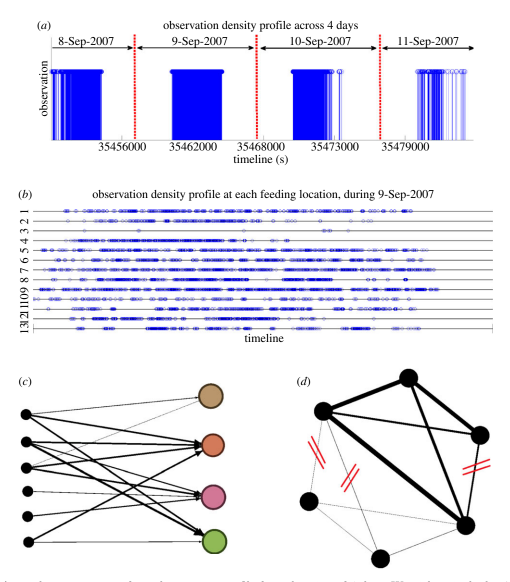


Figure 5. In (a), we show a segment of our data stream profile for a duration of 4 days. We pick a single day 'data-chunk' of observations and break it down into separate streams that refer to bird records at each particular location, as shown in (b). For each location-specific stream, we use our method to identify gathering events, as shown in coloured nodes on the right of the bipartite graph in (c). We assign birds (black nodes on the left of the graph) into such events based on their participation strength. We project the bird-to-event bipartite graph of (c) into an one-mode network based on co-occurrences in gathering events, as shown in (d). We remove any links (marked with double lines) that can be explained away by the null model. (Online version in colour.)

event k. Based on §3c, we then perform one-mode projection of this bipartite network into a bird-to-bird social network, shown in figure 5d, described by the adjacency matrix $\mathbf{A}^{(\ell)}$. The weight of each link $a_{ij}^{(\ell)} = \sum_{k}^{K} \min(b_{ik}^{(\ell)}, b_{jk}^{(\ell)})$ denotes the total number of cooccurrences between bird i and j across all K gathering events that took place at location ℓ . The statistical importance of each $a_{ij}^{(\ell)}$ is then tested against the null model we formulated in §3d, where all links below the significance threshold (marked with double lines in figure 5d) are removed. For our significance test, we used $R = 10^4$ samples of the null ensemble along with a standard $\alpha = 0.05$ importance threshold.

We repeat this process for all L locations and based on §3*e*, we combine all site-specific adjacency matrices $\mathbf{A}^{(\ell)}$ to a single one \mathbf{A}_t that captures the populationwide social structure on the given day T. An example is shown in figure 6, where we have summarized the subgraphs (such as the one shown in figure 5c) from all L = 13 locations shown in figure 5b into a single, global network that describes wild-bird social organization on 9 September 2007. We repeat the process for all T 24 h segments of our data stream, we get a stack of adjacency matrices $\{\mathbf{A}_t\}_{t=1}^T$ that represent daily snapshots of the wild-bird social network.

From an implementation perspective, GMMEvents runs L times for each day-segment of the data stream. For each location ℓ , R randomizations of the bird-toevent incidence matrix **B** are generated and for each one we perform one-mode projection in order to sample the weight distributions for each link pair i, j. Although it may appear computationally prohibitive for large datasets, our method is able to analyse 2 years' worth of data that correspond to about 1 million observations in approximately 6 h, run on a modern 8-core machine under a Matlab implementation. This is due to the fact that our method itself is executed on multiple small

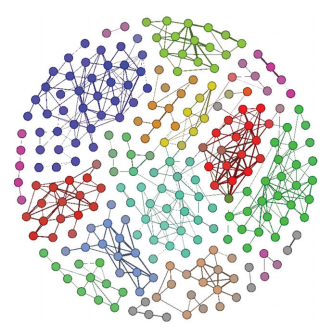


Figure 6. The Wytham woods *Parus major* wild bird social network at 9 September 2007, with N = 240 nodes, M = 491 edges, created by integrating all location-specific subgraphs shown in figure 5*d*. Note that not all 770 birds of the 2007–2008 season have been recorded during that day and also individuals no connections have been removed from the network. (Online version in colour.)

sub-streams (that refer to different locations per day) and can be directly parallelized. Our R randomization/ sampling schemes are also independent by definition, so they can run concurrently on different processing units. More details on computational issues are discussed in the electronic supplementary material and our source code documentation.

4.2. Using GMMEvents to study the dynamics of mating pair formation

In this section, we examine the validity of the graphs we extracted in $\S4a$ using GMMEvents. As the ground truth network is not available to us in such settings, we cannot directly compare our inferred topologies with some form of given solution. Although tests on simulated data streams have been performed (see the electronic supplementary material), our aim is to examine how well our dynamic network reflects meaningful quantities from our application-domain perspective.

We make use of an additional dataset, compiled from an *independent* field study at Wytham woods, which provides wild-bird mating records for each season. Such *pedigree* dataset logs the IDs of individuals that formed a breeding pair each year. Some bird pairs persist over several seasons while others last for only 1 year owing to either divorce or fatalities. We assume that if the extracted network structure is valid, then breeding individuals will be closely connected, either in terms of a direct link or being in the same *social circle*. Although looking for direct links between mated individuals is an obvious choice, it is a very strict case and thus very sensitive to missing data and noise. Therefore, our approach is to examine if breeding pairs belong more frequently and consistently than random into social circles that denote birds with similar foraging patterns.

Our first objective is to identify such social circles in our population. In figure 6, where we have visualized the network structure of the wild-bird population for a specific day, we can see certain regions in the graph (shown in different colour) where nodes are more *densely connected* with their immediate neighbours than the rest of the population. Such 'hot-spots' of increased link presence are called modules or *communities* in the network analysis jargon [7,16]. For each daily network described by \mathbf{A}_t , we extract such communities using a non-negative matrix factorization (NMF) approach [17].

We find that the majority of mated pairs in network communities are connected through a direct link in 77.26 per cent of cases for the 2007-2008 data and 71.57 per cent of cases for the 2008-2009 data. Reachability through a path of two links is reported for the 14.74 per cent of cases in 2007-2008 and 17.06 per cent of cases in 2008-2009. The average path length between two members, for the cases where both of them are observed in the data, is 1.33 (2007-2008) and 1.46 (2008-2009) with median value of 1 in both datasets. Finally, there are still cases (8% in 2007-2008 and 11.37% in 2008-2009) of pairs where their geodesic distance spans from three to six edges but still belong to the same community.

We monitor bird membership within these groups using a binary matrix \mathbf{C}_t , where each element $c_{ijt} = 1$ denotes that birds i, j appeared in the same community at day t. This leads us to a new collection of co-membership matrices $\{\mathbf{C}_t\}_{t=1}^T$ that encode temporal changes in the way birds participate with each other in communities. From a summation across t, we get a matrix $\mathbf{C}^{(s)} \in \mathbb{R}^{N \times N}$ where each element $c_{ij}^{(s)}$ denotes the total number of days in the season where the pair i, j participated in the same community. In figure 7, we plot a histogram of all co-membership values (y-axis on a logarithmic scale) based on two matrices $\mathbf{C}^{(s)}$ that refer to bird co-membership values in field seasons 2007-2008 and 2008-2009, respectively. We can see that for both seasons, the vast majority of pairs have never participated in the same group and the distribution is heavily skewed. This implies a strong preferential mechanism in the population, where random individuals rarely belong to the same social circle.

We now examine if the above distribution holds for certain sub-category of pairs in the network, which we know a priori are connected with actual social ties. This prior information is provided by the pedigree dataset we mentioned previously, which gives a list of node dyads i, j that denote breeding individuals. In this list, we also distinguish between mated pairs that were formed *during* our observation season, called *new* pairs, and others that already existed before, called old pairs. In figure 8, we plot the cumulative distributions $F(c_{ij})$, where c_{ij} are values co-membership matrix $\mathbf{C}^{(s)}$ and *i*, *j* can be (a) any node pair (blue circles, stem), (b) a new pair (green squares, stem) and (c) old pair (red triangles, stem). In figure 8a, we plot the distributions that refer to the 2007–2008 season, with N = 217 individuals, from which we have 49 new pairs and 20 old pairs. For season 2008–2009, shown in

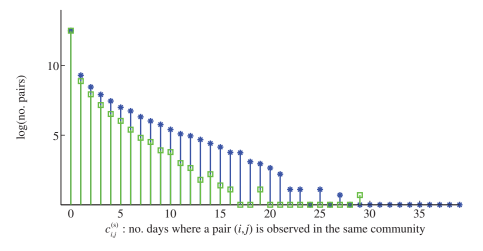


Figure 7. We plot the co-membership values of $\mathbf{C}^{(s)}$ on a base-2 logarithmic scale. Each value (*x*-axis) denotes the total number of days a random pair is observed in the same community. We can see that $\mathbf{C}^{(s)}$ is sparse and the vast majority co-membership values are zero. This shows that if we pick a random dyad in the population, it will most probably be never seen in the same social circle. Asterisks with continuous line, season 2007–2008; open square with continuous line, season 2008–2009. (Online version in colour).

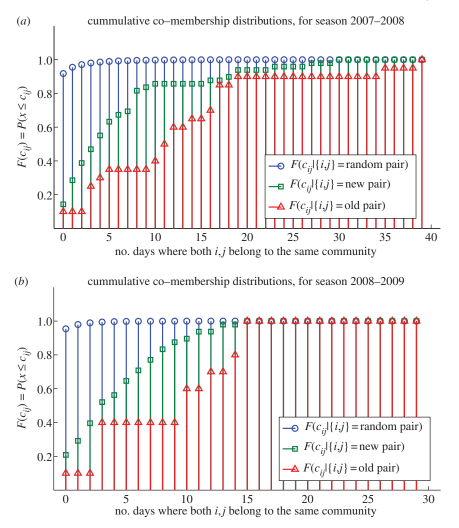


Figure 8. We plot the cumulative co-membership distributions for three different dyad types: random pairs, mating pairs formed in previous seasons and pairs that formed in the current season. Although for the majority of random bird pairs in the network comembership values are concentrated around zero, breeding individuals tend to participate much more frequently into the same flocks. (Online version in colour.)

figure 8b, we have N = 203 individuals that include 48 new pairs and 10 old pairs.

We can see that for both seasons presented in figure 8, the distributions that refer to mated pairs differ significantly from the one for random ones, with *p*-values less than 10^{-15} under a Kolmogorov–Smirnov test [18] with 5 per cent precision level for both seasons. In contrast to the random case, where values c_{ij} are mostly zero,

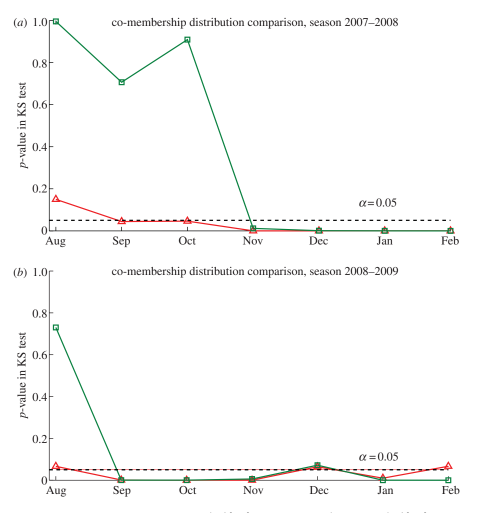


Figure 9. We compare the co-membership distributions $P(c_{ij}|\{i,j\} = \text{random pair})$ versus $P(c_{ij}|\{i,j\} = \text{old pair})$ (red triangles, line) and $P(c_{ij}|\{i,j\} = \text{random pair})$ versus $P(c_{ij}|\{i,j\} = \text{new pair})$ (green squares, line) in a month-by-month basis, using a Kol-mogorov–Smirnov test. Values above the proposed $\alpha = 0.05$ significance threshold imply that the two distributions under comparison are similar. We can see that from very early in the year old pairs differentiate themselves from random, by starting to participate frequently in the same communities. On the other hand, members of new pairs in the beginning of the year treat each other as random, while preferential mechanism that makes them flock together, starts to build-up during early winter. (a, b) Triangles with solid line, random versus old pairs; squares with solid line, random versus new pairs. (Online version in colour.)

co-membership for mated pairs achieves larger values, thus denoting stronger and consistent graph proximity. The differences between old and new pairs are also revealed between their respective cumulative distributions (green squares, stem and red triangles, stem), where old pairs achieve higher co-membership values owing to the fact that they existed before new pairs where formed, thus they had more opportunities during the season to participate in the same foraging flocks.

We have already seen that co-membership distributions differ between various pair types. We will now examine place that differentiation takeswhen during the observation season. We start by breaking down the observation period into eight months. For each month, we used the respective daily networks in order to find the three co-membership distributions of interest. We then compared $P(c_{ij}|\{i, j\} = \text{random pair})$ versus $P(c_{ij}|\{i, j\} = \text{old pair})$ and $P(c_{ij}|\{i, j\} = \text{random pair})$ versus $P(c_{ij}|\{i, j\} = \text{new pair})$, by calculating the *p*-value under a Kolmogorov–Smirnov test with a proposed significance level 0.05. In figure 9, we can see that at

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the beginning of the season, new pairs have similar co-membership patterns to random ones, as they have not been formed at such early point. But as we move through the year, this similarity drops and from the 'cloud' of random associations, breeding relationships emerge. On the other hand, old pairs that have been already formed from previous seasons have a consistent nonrandom co-membership pattern, even from very early points in the season.

5. DISCUSSION AND FUTURE WORK

The network paradigm is a powerful tool for studying real-world complex systems. As there is an extensive toolset of methods and algorithms for network analysis, in this work we have focused on the problem of constructing the network in the first place. In many applications, the collected data capture the behaviour of the system in some manner, like the spatial trajectories of participating agents, but not the underlying relations between them. We address this issue by assuming that mobility patterns of individuals may be correlated based on some form of underlying social connection. By identifying observation-dense regions in the data stream, which can be seen as *gathering events* of affiliated individuals, we propose a methodology of drawing links between agents based on their co-participation into those events.

Traditional approaches [9-12] to constructing social networks from spatio-temporal data involve discretizing the observation stream based on some fixed time window Δt and drawing links between individuals when they lie within such 'interaction-radius'. Our method overcomes the practical difficulties of such time-slicing approach in cases when we have no prior knowledge of how big or small the time window size should be, thus having to perform multiple runs across various Δt and select the appropriate one based on some ad hoc quality function. Additionally, we have proposed an appropriate null model, which allows us to examine if the co-occurrence of individuals into gathering events are a result of a latent social tie, or coincidence. Our null model retains the 'bursty' nature of the data stream but breaks all correlations between the individuals' appearance patterns through an appropriate randomization.

We applied GMMEvents into two large-scale datasets that provide wild-bird foraging records. We showed that the inferred network topologies reflect mating pair formation events that take place in the population, where breeding individuals tend to belong into the same foraging groups more often that random dyads. We also showed that the dynamics of community structure in the system reveal how newly formed pairs initially have a random-like behaviour, while as we approach the mating season they start to participate more often than random into the same communities.

The communities identified here are based on temporal occurrence at feeding stations, and while the data analysed here are extensive, they are incomplete, as observations are made for only a proportion of the time, and only for feeding-related activity. While more complete data would be expected to result in more completely connected communities (both in terms of link number and connection strength), it is not necessarily the case that all communities would ultimately be fully connected. For example, communities might be comprised of pairs of individuals that avoided each other (e.g. territorial males, competing females) relative to the other members of the community, even though they have links via other individuals. As expected for individuals linked via a network, there is a variety of direct and indirect ways that individuals within and between communities might influence each other. In the case of the present network, we might expect that an important source of direct effects lies in the flow of information between community members about the presence of food, but such information will also spread indirectly to other individuals via network links between communities [19]. Numerous other effects might also be considered. For example, like many animals, small passerine birds give alarm calls that alert other individuals about the presence of predators [20]. While the individuals in the same community may be

expected to be nearest to a focal individual, other linked communities may also be influenced directly by this sort of behaviour, and the overall inter-community network may serve as a hypothesis for the likelihood of such effects being transmitted between individuals. So far all feeding sites have been analysed in isolation until the last stage. Site-specific network adjacency matrices are extracted and tested for significance whence they are all combined to one single adjacency matrix. An alternative to this spatial aggregation over sites would be temporal aggregation. In this approach, temporal data could be aggregated and behaviour and feeding sites analysed directly. While such an analysis may account for popular feeders, it would not achieve the high temporal resolution of the existing approach. For instance, using a temporal aggregation strategy, a group of birds feeding in the morning and one in the evening would all be treated as one single group when the times of their feeding site visitation clearly suggest otherwise. A proper resolution of this conflict may require a full spatio-temporal clustering stage and another bespoke hypothesis test to detect both spatially and temporally insignificant events. Such a multivariate approach would alleviate the necessity to account for spatial correlations during hypothesis testing which otherwise would be extremely hard to extract from data. Thus, in our future work, we will focus on a full spatio-temporal analysis of bird behaviour and the development of clustering models that combine data of different characteristics, such as the bursty behavioural and the continuous spatial data.

The next stages of our research consist of two main modules. From the perspective of the model, we seek to extend the way we define the link a_{ijt} between two individuals at time t so that we take into account *prior knowledge* from previous observations. This has the advantage of capturing the uncertainty over the link weight, detect abrupt changes in the network topology and handling missing observations in a principled manner. From an ecological point of view, we currently run an improved scheme of our data collection, where we have sensors at each feeding location. This gives us the advantage of looking at the data at much greater resolutions thus having a more accurate view of the overall bird population's foraging patterns.

Although the methodology we presented is applied to animal observation records, it can be extended to any system where agents perform check-ins at certain locations and such observations are not uniformly spread in data stream, but *temporally focused*. We believe methodologies and theoretical results derived from the study of animal social networks will benefit the wider field of network analysis, as individuals can be monitored from the beginning to the end of their lifespan, there are no privacy issues associated with data collection and understanding the dynamics of animal interactions provides an insight into the behaviour and evolution of complex systems.

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