Why patterns of assortative mating are key to study sexual selection and how to measure them

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Abstract The study of sexual selection is being revolutionised by the realisation that most populations exhibit some degree of polyandry, i.e. females mating with multiple males. Polyandry can drastically change the operation of sexual selection on males as it reduces the reproductive success that males derive by mating with different females, by forcing their ejaculates to compete for fertilisation after copulation (sperm competition). Variation in polyandry within a population means that the impact of polyandry can differ drastically across males, depending on the polyandry of their own mating partners. Because the patterns through which males share mates within a population may have strong repercussions for variation in male reproductive success, measuring such patterns is critical to study the operation of sexual selection. Several methods have been proposed to measure the pattern of mate sharing at the population level. Here, we develop a new method (sperm competition intensity correlation, SCIC) and compare its performance against two established methods (Newman’s assortativity and nestedness), using both idealised model populations and random simulated populations, across a range of biologically relevant population parameters: (i) population size, (ii) sex ratio and (iii) the ‘mating density’ of the population. We conclude that SCIC may be the most promising approach, as it is both internally consistent and robust across the parameter range. We discuss some important caveats and provide advice regarding the choice of method for future studies of sexual selection.

Keywords Mating system · Sperm competition · Sexual selection · Sexual networks · Nestedness NODF · Newman’s assortativity · SCIC

Introduction

Sexual selection is the selective process arising from variation in reproductive success due to intrasexual competition over access to mates and their gametes (Darwin 1871; Andersson 1994). Traditionally, competition over mates was viewed as the main source of intrasexual variation in reproductive success, and the strength of sexual selection on mating success was measured by the Bateman gradient, i.e. the slope of the univariate regression of total reproductive success, \( T \), over mating success, \( M \):

\[
T(M) = (\beta_M \times M) + \varepsilon
\]  

(1)

The Darwin-Bateman paradigm predicts that the male Bateman gradient (i.e. \( \beta_{M\text{male}} \)) is typically steeper than...
that of females, due to stronger precopulatory sexual selec-
tion on male mating success (Bateman 1948; Andersson
1994; Parker and Pizzari 2015). Less appreciated, how-
ever, are the profound implications that female mating suc-
cess (M\text{females}) can have on the strength of sexual selec-
tion on males. When females are polyandrous (M\text{females} > 1)
such that the ejaculates of multiple males overlap at the
time of fertilisation (Pizzari and Wedell 2013), the scope
of male monopolisation is fundamentally changed. This is
because, for males, mating is no longer a zero sum game
as females become a shared resource, and males continue
to compete after mating for the fertilisation of sets of ova,
via sperm competition (Parker 1970) and under cryptic
female choice (Childress and Hartl 1972; Thornhill
1983). Therefore, in polyandrous populations (i.e. female
mating success can be greater than one), male reproductive
success is not only contingent on the number of females
mated and their fecundity but—crucially—also on the pro-
portion of ova produced by his partners that a male man-
ges to fertilise (Webster et al. 1995). Males that mate
with the same female share the paternity of her ova, and
the proportion of ova that a male fertilises is therefore a
function of the intensity of sperm competition (sensu
Parker 1998) faced by the male’s ejaculates across the
females that he inseminates: the more intense the sperm
competition, the lower the average share of paternity for
each male (Shuster and Wade 2003; Parker and Pizzari
2010). In polyandrous populations, sperm competition in-
tensity is therefore a key determinant of male reproductive
success (Parker and Pizzari 2010; Collet et al. 2012;
Kvarnemo and Simmons 2013; Parker and Birkhead
2013). Crucially, when individual females in a population
vary in their polyandry (variance in M\text{females} > 0), some
males have the potential to suffer higher sperm competi-
tion intensity than others. The way in which sperm com-
petition intensity is distributed across males dictates
population-level patterns in sexual selection, determining
both the total variation in male reproductive success and
the strength of precopulatory sexual selection on mating
success (i.e. \(\beta_{M\text{females}}\)). For example, consider a population
where males with relatively high mating success tend to
mate with the most polyandrous females and as a conse-
quency face intense sperm competition. Despite the rela-
tively high mating success of these males, their reproduc-
tive success is limited by the high polyandry of their
partners, which reduces their paternity share. At the pop-
ulation level, such non-random mating patterns may weak-
en sexual selection on male mating success (i.e. \(\beta_{M\text{females}}\)), as
further increases in mating success result in diminishing
 gains in fertilisation success (Sih et al. 2009; McDonald
et al. 2013; McDonald and Pizzari 2014). Similarly, a
negative relationship between male mating success and
the mating success of his mates at the population level
would strengthen \(\beta_{M\text{females}}\). Understanding the operation of
sexual selection therefore requires a measure of the rela-
tionship between the mating success of a male and the
sperm competitive environment faced by his ejaculates,
i.e. the mating success (i.e. polyandry) of his partners
within a population (Sih et al. 2009; McDonald and
Pizzari 2014).

Borrowing techniques developed in both social science
and ecological research focusing on food webs and mutu-
alistic interactions, two such measures have recently been
proposed: Newman’s assortativity (\(r\)) and nestedness metric
based on overlap and decreasing fill (NODF) (Newman 2002;
Almeida-Neto et al. 2008). Both methods utilise a network
perspective of sexual interactions, where mating populations
are described as a collection of nodes (males and females) that
are connected by edges representing copulations (Fig. 1).
Such sexual networks have been extensively explored in
studies investigating the spread of sexually transmitted
infections (Gupta et al. 1989; Liljeros et al. 2001), but their
application to sexual selection begun only recently
(McDonald et al. 2013; McDonald and Pizzari 2014;
Inghilesi et al. 2015; Muniz et al. 2015). Mating patterns show
a tremendous variety both between and within populations
(Emlen and Oring 1977; Thornhill and Alcock 1983;
Clutton-Brock 1989; Andersson 1994; Shuster and Wade
2003; Shuster 2009). The rapid surge in the availability of
fine-grained behavioural data sets in combination with
molecular parentage assignment (e.g. Preston et al. 2005;
Rodríguez-Muñoz et al. 2010; Collet et al. 2012; Péliissié
et al. 2014) is generating increasing scope for network
approaches, and metrics such as Newman’s assortativity (\(r\))
and nestedness (NODF) are gaining prominence as tools to
quantify mating patterns. Yet, despite this potential, we are
aware of no analytical or quantitative evaluations of these
methodological tools.

Here, we first review both, Newman’s assortativity (\(r\))
and nestedness (NODF), in the context of sexual networks
and introduce a third measure that we term ‘sperm com-
petition intensity correlation’ (SCIC). Secondly, we com-
pare the performance of these three metrics in quantifying
the assortativity of mating patterns based on mating suc-
cess. Thirdly, we use randomly organised simulated popu-
lations to explore the sensitivity of these three metrics to
each of three key axes of sexual selection variation: (i)
population size, (ii) population adult sex ratio and (iii)
the ‘mating density’ of the population. ‘Mating density’
measures the percentage of male–female pairs that copu-
late out of all possible heterosexual pairing combinations
in a given population and thus reflects the saturation of a
sexual network (i.e. the overall promiscuity of a popula-
tion), independently of population size or sex ratio (see
below). Furthermore, we examine how these three metrics
(i.e. Newman’s \(r\), NODF and SCIC) relate to each other

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across these three axes. Finally, in light of these results, we consider the application of these metrics to the study of sexual selection.

**Methods**

**Sexual networks and measures of the mating structure**

A sexual network is a collection of individuals (nodes) connected by links (edges) representing (inter- and/or intra-) sexual interactions, where the total number of links of a node is its degree (Croft et al. 2008; Wey et al. 2008; Sih et al. 2009; McDonald et al. 2013; Krause et al. 2014; McDonald and Pizzari 2014). In this study, we are interested in sperm competition so we focus on bipartite networks, that is networks with two sets of nodes, male and females, with edges between heterosexual mating partners and where an individual’s node degree represents its mating success. Conceptually, edges in these networks can either be directed or undirected. In an undirected network, when one male and one female are connected to each other, both are treated as having one link (male and female degree = 1). In a directed network, if one male and one female are connected, one individual (e.g. the male) is considered as having an outward link with that female (out-degree = 1, in-degree = 0) and the female is considered as having one inward link with that male (out-degree = 0, in-degree = 1). When females are polyandrous, their degree is greater than 1 and measures the intensity of sperm competition faced by their male partners (Parker 1970; Parker and Birkhead 2013). Below, we introduce three network metrics aimed at quantifying the relationship between a male’s mating success and that of his partners: Newman’s assortativity ($r_{\text{Newman}}$), nestedness (NODF) and SCIC.

**Newman’s assortativity ($r_{\text{Newman}}$)**

Newman’s assortativity ($r_{\text{Newman}}$) measures the assortativity of a network as the Pearson correlation coefficient between the trait values at either end of an edge (i.e. the link connecting a female to her mating partner), across all edges in the network (Newman 2002, 2003). The trait of interest in this context is node degree, which measures individual mating success (see above). Developed for unipartite networks (i.e. networks with one type of node, where edges can be potentially drawn between all individuals), $r_{\text{Newman}}$ can be defined as:

$$r_{\text{Newman}} = \frac{\sum j_i k_i E^{-1} \sum j_i \sum k_i}{\sqrt{\left[\sum j_i^2 E^{-1} \left(\sum j_i\right)^2\right] \left[\sum k_i^2 E^{-1} \left(\sum k_i\right)^2\right]}}$$  \hspace{1cm} (2)$$

where $E$ is the total number of edges in the network and $j_i$ and $k_i$ are node degrees of individuals $j$ and $k$ at either end of the $i$th edge. Therefore, $r_{\text{Newman}}$ ranges from 1 (positive correlation, positive assortativity) to −1 (negative correlation, disassortativity). In a network of one monogamous mating pair, $E$ equals 2 if we treat edges as undirected or 1 if we treat edges as directed. This is because in undirected networks, the edge is considered mutual and the correlation between traits at each end of the edge ranges from 1 to −1. In the case of directed networks, the edge is considered mutually exclusive and the correlation between traits at each end of the edge ranges from 1 to 0.
network with two males as rows and two females as columns, $[\begin{array}{cc} 1 & 1 \\ 1 & 0 \end{array}]$, where 1 means that the pair copulated. The directed Newman’s assortativity, $r_{Newman(D)}$, would treat this network as having three edges ($E = 3$), where the trait values (i.e. mating success) for the outgoing male links are $\binom{2}{1}$ and the trait values for the female inward links are $\binom{2}{1}$. Alternatively, Newman’s assortativity for undirected networks, $r_{Newman(U)}$, treats each link as a mutual connection and so would treat this network as having six edges ($E = 6$) where the trait values at either end of each link are $\binom{2}{1}$ and $\binom{2}{1}$.

This means that $r_{Newman(U)}$ can yield both positive and negative values of assortativity and behaves in a way similar to $r_{Newman(D)}$ when populations approach an even sex ratio, but the metrics can generate drastically divergent estimates of assortativity when sex ratios deviate from unity.

**Nestedness (NODF)**

Nestedness is a concept originally borne from ecological research, designed to quantify patterns of species co-occurrences in metacommunities, where sites with lower species richness contain reduced subsets of those sites with higher species richness (Ulrich and Almeida-Neto 2012). This concept was later applied to ecological networks such as mutualistic plant-pollinator networks (Ulrich et al. 2009). Mutualistic plant-pollinator networks are nested when specialist plants (those with few connections) tend to connect to the most generalist pollinators (those with many connections) and specialist pollinators tend to connect to the most generalist plants (Bascompte et al. 2003; Bascompte and Jordano 2007; Ulrich et al. 2009). In the context of sexual networks, a population is nested when males with few mating partners (i.e. low mating success; $M$) tend to mate with the most polyandrous females (i.e. females with high $M$), and males with the highest mating success tend to include amongst their partners the least polyandrous females (i.e. females with low $M$). Nestedness is therefore specific to patterns of disassortativity and bipartite networks, i.e. negative assortiment between male mating success and the mating success of their partners (McDonald and Pizzari 2014) and is perhaps best visualised by using the matrix representation of sexual networks (Fig. 1e). Accordingly, the most widely used methods to calculate nestedness is NODF (Almeida-Neto et al. 2008; Strona and Fattorini 2014), which measures nestedness based on the rules of percentage overlap (PO) and decreasing fill (DF) between pairs of rows and pairs of columns of such matrix and quantifies the extent to which the mating partners of individuals with low degree form subsets of the mating partners of those individuals with a higher degree (McDonald and Pizzari 2014). To calculate nestedness as outlined by Almeida-Neto et al. (2008) for sexual networks, we first order the mating population matrix of left to right (females) and top to bottom (males) in order of decreasing mating success. Then, for every pair of $i$ and $j$ rows (males), we calculate two values, DF (decreasing fill) and PO (percentage overlap). To calculate DF, we ask whether the upper row in the pair (row$i$) has a higher mating success ($M_i$) than the lower row (row$ji$). If $M_j < M_i$, then $DF_{ij} = 100$ and if $M_j \geq M_i$ then $DF_{ij} = 0$. The PO of pairs of rows is calculated as the percentage of row$i$’s mating partners shared with row$ji$. The same process is calculated for all pairs of columns (females), where the $j$th female is the leftmost and the $ji$th female is the female to its right. For all pairs of rows and columns, we then calculate their individual nestedness ($N_{ij}$) as:

If $DF_{ij} = 0$, then $N_{ij} = 0$

If $DF_{ij} = 100$, then $N_{ij} = PO_{ij}$

For example, consider the top male rows in Fig. 1e. The top male has six mates and the second male has five mates, therefore $DF_{ij} = 100$. The second male also shares all of his females with the top male, so the percentage overlap $PO_{ij} = 100$ and $N_{ij} = 100$. We can then measure the nestedness of the whole network as:

$$\text{NODF} = \frac{\sum N_{ij}}{\left(\frac{f(f-1)}{2}\right) + \left(\frac{m(m-1)}{2}\right)}$$

where $f$ and $m$ are the numbers of males and females, respectively, and NODF ranges between 100 (perfect nestedness) and 0 (no nestedness) (Almeida-Neto et al. 2008).

**SCIC**

The aim of quantifying population-level patterns in assortative mating by mating success is to understand its influence on the strength of sexual selection on male mating success (i.e. $\beta_M$). When females are polyandrous, males will be forced to sperm compete and, all else being equal, the reproductive success of a focal male is inversely proportional to the number of sperm competitors with which he competes (i.e. his ‘sperm competitive intensity’, SCI). The sperm competition intensity suffered by a male can be characterised by the harmonic mean of a male’s partners mating success (Shuster and Wade 2003; Wade and Shuster 2005). For example, consider a male that mates with two females; one of which does not remate, whilst the other copulates with two other males. The sperm of this male does not face sperm competition within the first female. For the second female however, the sperm of the focal male must compete with the sperm of two other males. Assuming
the simplest null model of sperm competition (i.e. fair raffle and similar ejaculate size and fertilising efficiency across competitors), the focal male’s share of paternity for this female is \( \frac{1}{99} \) (Shuster and Wade 2003). The average share of paternity of the focal male across the two partners is then \( \frac{1}{99} \times (\frac{1}{2} + \frac{1}{3}) = 0.667 \) where \( 0.667 \) is the harmonic mean number of sperm competitors faced by the focal male (i.e. \( \frac{1}{0.667} \approx 1.4 \) 99 competitor males). This can be calculated for the \( i \)th male simply as:

\[
SCI_i = \frac{1}{M_i \left( \sum_j M_j \frac{1}{k_j} \right)}
\]  

where \( k_j \) is the mating success (degree) of the \( j \)th female mated with the \( i \)th male, and \( M \) is the total number of mating partners of the \( i \)th male (i.e. his mating success). This parameter can then be added to our regression model in Eq. 1 as:

\[
T_i = \left( \beta_{MSCI} \times M_i \right) + \left( \beta_{SCI-M} \times SCI_i \right) + \varepsilon
\]  

where \( \beta_{M,SCI} \) represents the male Bateman gradient controlling for variation in SCI. Crucially, the way in which the univariate Bateman gradient, \( \beta_M \), is affected by variation in sperm competition intensity across males is determined by the slope of the partial regression of mating success on the mean sperm competition intensity faced by a male (i.e. \( \beta_{SCI-M} \)) and the correlation between \( \beta_{M,SCI} \) and \( \beta_{SCI-M} \) (i.e. SCI, Fig. 2) as:

\[
\beta_M = \beta_{M,SCI} + (SCI \times \beta_{SCI-M})
\]  

In general, theory predicts that \( \beta_{SCI-M} \) will be negative as increased sperm competition decreases a male’s reproductive success (Shuster and Wade 2003). Therefore, when SCI is positive, we predict a reduction in male Bateman gradients, whereas a negative SCI will steepen the slope of the male Bateman gradient (Wolf et al. 1999). Typically, Bateman gradients are standardised by dividing reproductive success and fertilising efficiency across competitors, the focal male (i.e. his mating success) and provide a standardised slope for SCI that facilitates comparisons across populations. If instead, \( M_{males} \) was standardised by subtracting the population mean and dividing by its standard deviation, as for selection gradients on phenotypic traits (Lande and Arnold 1983; Wolf et al. 1999), SCI could be calculated as the Pearson product correlation coefficient between SCI and \( M_{males} \). We provide R code to calculate SCI as supplementary material (R Core Team 2014).

**Model populations**

We develop model populations displaying a range of mating systems to test the performance of all the metrics (i.e. \( r^{Newman(U)} \), \( r^{Newman(D)} \), NODF and SCIC) in identifying mating success assortativity. Research has highlighted a tremendous variety both between and within populations (Emlen and Oring 1977; Thornhill and Alcock 1983; Clutton-Brock 1989; Andersson 1994; Shuster and Wade 2003; Shuster 2009). Variation in mating patterns (the mating topology) across populations can be decomposed into three main axes: (i) individual variation in male mating success (i.e. level of polygyny, \( var(M_{males}) \)), (ii) individual variation in female mating success (i.e. level of polyandry, \( var(M_{females}) \)) and (iii) the average level of polygyny and polyandry of the entire population (i.e. \( M_{males} \) and \( M_{females} \), respectively). In strictly monogamous populations with unitary sex ratio, \( M_{males} = M_{females} = 1 \), and there is no intrasexual variation in either male or female mating success (Fig. 1a). On the other hand, in populations with unitary sex ratio where the mating success of one sex (but not the other) exceeds 1, there is variation in mating success amongst members of that sex but no variation in mating success in the opposite sex (Fig. 1b). For sperm competition to occur, populations must exhibit some level of polyandry (i.e. \( M_{females} > 1 \); Pizzari and Wedell 2013). Furthermore, for males and females to mate assortatively by mating success, there must also be some variation in both male and female mating success (\( var(M_{males}) > 0 \), \( var(M_{females}) > 0 \), not including individuals that do not mate at all). Any metric measuring the assortativity of mating partners based on mating success should therefore only provide non-zero results when all three conditions are met, namely \( M_{females} > 1 \), \( var(M_{males}) > 0 \) and \( var(M_{females}) > 0 \).

None of the developed model populations in Fig. 3 display all three conditions and thus are not expected to generate assortative mating by \( M \) different from zero. On the other hand, the model populations in Fig. 4 meet all three conditions for assortative mating by \( M \). These populations represent different ‘mating densities’ (i.e. the proportion of possible mating pairs realised of all heterosexual pairing combinations possible in a given population) and display either positive or negative mating assortativity by \( M \).

**Simulated populations**

When measuring the relationship between male mating success and that of their partners, any useful metric should allow the estimation of such assortativity across a wide range of population parameters but should not be confounded by other naturally varying population parameters. Here, we test the performance of all metrics, \( r^{Newman(U)} \) and \( r^{Newman(D)} \), NODF and SCIC on randomly simulated mating populations along three fundamental gradients of variation in mating systems, namely (i) population size, (ii) sex ratio and (iii) ‘mating density’. We do so by varying each axis separately one at a time as outline below.
3. **Mating density**: We measured the ‘mating density’ of a population as the percentage of male–female pairs that mated out of the total possible combinations of heterosexual mating pairs (i.e. for a population of 10 males and 10 females, there are 100 possible mating pairs and if 50 of those pairs actually mated, the population would have a ‘mating density’ of 50 % or a density of 0.5 in network nomenclature). ‘Mating density’ therefore gives an indication of how well connected sexual networks are that is independent of population size or sex ratio. This parameter is important as many network analysis measures are strongly affected on the overall density of networks (Croft et al. 2008). We simulated populations that differed in the realised proportion of all possible mating combinations (‘mating density’) but not in population size or sex ratio. To do so, we created 1,000 populations, all with 20 males and 20 females (i.e. same population size and sex ratio), where either 25, 50 or 75 % of possible pairs mated. We simulated mating by randomly connecting males and females until 50 % of the total possible mating pairs was realised ensuring that every individual mated at least once. Of the 1,000 populations with 6 males and 6 females (6 × 6), ten populations had zero variation in female mating success and four populations had zero variation in male mating success, so no value could be calculated for \( r_{\text{Newman}(D)} \) in the former and SCIC in the latter case. These populations were removed from analyses that included these two metrics, respectively.

2. **Sex ratio**: We created 1,000 populations with 10 males and 50 females (1:5), 30 males and 30 females (1:1) or 50 males and 10 females (5:1) and simulated mating by randomly connecting males and females until 50 % of the total possible mating pairs was realised ensuring that every individual mated at least once. Each population therefore had the same population size and ‘mating density’.

To assess the independence of assortative mating measures across each level of population size, sex ratio and mating density, we report the Spearman rank correlation coefficient between all assortative mating measures: \( r_{\text{Newman}(U)}, r_{\text{Newman}(D)}, \text{NODF}, \text{and SCIC} \), with all population parameters. To explore whether each measure is likely to capture similar information, we also calculated the Pearson product moment correlation coefficient between each assortative mating measures at each level of population size, sex ratio and mating density. All analyses were conducted using R statistical software (R Core Team 2014). As this study is entirely based on in silico simulations rather than empirical data, no blinded protocol was required.

**Results**

**Model populations**

We first assessed the consistency of each measure of assortative mating (\( r_{\text{Newman}(U)}, r_{\text{Newman}(D)}, \text{NODF, and SCIC} \)) by applying all measures to the six model populations presented in Fig. 3. No population in Fig. 3 satisfies the criteria required for a correlation between male mating success and the mating success of his partners (i.e. polyandry and variation in both male and female mating success), and thus we should expect measures of assortative mating to return zero or no value for all six populations. As expected, all metrics returned zero or no value for all populations with the exception of \( r_{\text{Newman}(U)} \) that returned negative values for four populations (Fig. 3b, c, e and f and Table 1). This is because \( r_{\text{Newman}(U)} \) was originally
intended for unipartite networks (networks with one type of node, e.g. all males). Therefore, although there is no variation in mating success within one sex (which would preclude the calculation of a correlation coefficient), $r_{\text{Newman}(U)}$ instead correlates the mating success (degree) of individuals at either end of an edge (i.e. effectively counting each copulation twice from both the male and female end of an edge). Therefore, although there is no variation in degree within a sex, $r_{\text{Newman}(U)}$ utilises variation between sexes when calculating the correlation between male and female mating success.

We next assessed the sensitivity of all metrics to detect positive or negative assortativity by mating success across a range of mating structures in idealised populations. All metrics provided qualitatively logical values, consistent with expectations across all matrices, with the exception of NODF that returned zero values for two populations (Table 1 and Fig. 4a, e). This inconsistency arises for two reasons: firstly, because despite the apparent negative correlation between male mating success and the mating success of their partners in the population in Fig. 4a, this population does not show both decreasing fill and overlap in the mating partners between any males. Secondly, the population shown in Fig. 4e is a completely positively assorted, and NODF is a measure specific to negative assortment (i.e. disassortativity) and thus identifies this population as zero nestedness. Therefore, only two parameters $r_{\text{Newman}(D)}$ and SCIC provided consistent results across all model populations (Table 1).

### Simulated populations

1. **Population size**: All assortative mating measures were moderately correlated with population size, with the exception of NODF, which was decoupled from population size (Fig. 5a, d, g and j), confirming the result for NODF presented in Almeida-Neto et al. (2008). The

| Model population | SCIC | NODF | $r_{\text{Newman}(U)}$ | $r_{\text{Newman}(D)}$ | Overall assortative mating pattern |
|------------------|------|------|-------------------------|-------------------------|----------------------------------|
| Figure 3a        | –    | 0.000| –                       | –                       | None                             |
| Figure 3b        | 0.000| 0.000| –0.615                  | –                       | None                             |
| Figure 3c        | –    | 0.000| –1.000                  | –                       | None                             |
| Figure 3d        | –    | 0.000| –                       | –                       | None                             |
| Figure 3e        | 0.000| 0.000| –0.091                  | –                       | None                             |
| Figure 3f        | –    | 0.000| –0.615                  | –                       | None                             |
| Figure 4a        | –0.385| 0.000| –1.000                  | –1.000                  | Negative                         |
| Figure 4b        | –0.342| 33.333| –0.833                  | –0.833                  | Negative                         |
| Figure 4c        | –0.555| 100.000| –0.500                  | –0.500                  | Negative                         |
| Figure 4d        | –0.193| 33.333| –0.167                  | –0.167                  | Negative                         |
| Figure 4e        | 1.000| 0.000| 1.000                   | 1.000                   | Positive                         |
| Figure 4f        | 0.515| 20.000| 0.550                   | 0.550                   | Positive                         |

*– represents when no result is estimated by the method or is undefined (i.e. NA or NaN)*
4. Correlations between assortative mating measures. The correlations of \( r^{\text{Newman}(U)} \) and \( r^{\text{Newman}(D)} \) with population size tended towards zero as populations became larger. This observation is significant because it reveals that negative assortativity can emerge as a property of random mating in very small groups or populations, whereas deviations from 0 in larger populations more likely require a biological explanation (e.g., behavioral strategies).

2. Sex ratio: All assortative mating measures were very weakly or weakly correlated with population sex ratio (Fig. 5b, e, h and k). The measure \( r^{\text{Newman}(U)} \) however, showed a clear strong, non-monotonic relationship with sex ratio (Fig. 5b). When sex ratios were skewed in either direction, \( r^{\text{Newman}(U)} \) tended to be strongly negative. This is because uneven sex ratios resulted in one sex having a higher mean mating success than the other sex, and \( r^{\text{Newman}(U)} \) reflects the tendency for one sex with high mating success to mate with the other sex, which necessarily has a lower mean mating success.

3. Mating density: All assortative mating measures were very weakly correlated with the ‘mating density’ of populations (Fig. 5c, f and i), with the exception of NODF which was very strongly positively correlated with ‘mating density’ (Fig. 5l), confirming previous results for NODF (Almeida-Neto et al. 2008).

4. Correlations between assortative mating measures. Overall, for population size and ‘mating density’ simulations \( r^{\text{Newman}(U)} \) and \( r^{\text{Newman}(D)} \) showed a marked pattern across different sex ratios (Fig. 6). When populations diverged from an equal sex ratio, correlations between \( r^{\text{Newman}(U)} \) and all other measures reversed direction and weakened, reflecting the tendency for \( r^{\text{Newman}(U)} \) to identify increasingly negative assortative values when sex ratios are biased in either direction. Overall, \( r^{\text{Newman}(D)} \) and SCIC were consistently and strongly correlated with each other across all parameter ranges (Fig. 6).

**Discussion**

Multiple mating by females (polyandry) results in variance in male paternity share, creating opportunity for postcopulatory sexual selection via sperm competition and cryptic female choice (Parker 1970; Childress and Hartl 1972; Thornhill 1983). Consequently, in polyandrous populations, males can increase reproductive success by both mating with more females and/or increasing their paternity share (Alcock 1994; Alonzo and Warner 2000; Parker and Birkhead 2013), for example by maintaining exclusive access to mating partners by curtailing their polyandry (e.g., through mate guarding). Sexual selection on male mating success (\( \beta_{M,\alpha} \)) is stronger when males who mate with many females also enjoy a higher exclusivity of mating. Conversely, in populations with positive assortativity between male and female mating success, males with high mating success also face more intense sperm competition. This weakens selection on male mating success (\( \beta_{M,\alpha} \)) because increasing mating success results in decreasing returns in terms of reproductive success, due to reduced paternity share. Therefore, understanding inter-population variation in sexual selection hinges on characterising population-level patterns of the association between male mating success and the mating success of his partners (assortative mating by \( \beta \)).

In this work, we outlined multiple approaches to measures of the relationship between male mating success and the mating success of his partners, namely Newman’s assortativity for undirected and directed networks \( r^{\text{Newman}(U)} \) and \( r^{\text{Newman}(D)} \), and nestedness (NODF) and introduced a third quantitative measure, the SCIC. We then used (i) idealised model populations to assess the logical performance of these measures in capturing assortative mating by mating success and (ii) simulated random mating populations to test the dependency of these measures on three main population parameters, namely population size, sex ratio and the ‘mating density’ of the population.

Our model populations allowed us to test the performance of all measures of assortative mating in situations where there should be zero assortativity or where there should be qualitatively positive or negative assortativity. Our results highlighted limitations in two measures, NODF and \( r^{\text{Newman}(U)} \) as both measures either indicate zero assortative mating by mating success when there should be positive or negative assortativity, or alternatively indicated assortative mating when there should be none. Instead, \( r^{\text{Newman}(D)} \) and SCIC showed either no assortativity or were undefined when there should be zero assortative mating and qualitatively consistent results when there should be non-zero assortative mating.

Our simulations allowed us to test for the dependency of all assortative mating measures on population parameters expected to vary in nature, namely (i) population size, (ii) sex ratio and (iii) the ‘mating density’ of the population. Our results show that most measures were often not strongly associated with these population parameters. However, SCIC and \( r^{\text{Newman}(U)} \) showed a bias towards negative values at small populations. When populations are very small, those males that mate with many females will tend to on average compete with males who have lower mating success, purely as a result of small, restricted populations. Whether such patterns can be explained by random processes
alone may be investigated through the use of randomised null models (Ulrich and Gotelli 2007). However, in such small populations, even small deviations from random may have a large effect of the distribution of male reproductive success and a first step should be to explore the link between measures of assortative mating and the operation of sexual selection. In two cases, we identified strong relationships between population parameters and measures of assortativity. In our sex ratio simulation, \( r_{\text{Newman}(U)} \) showed a strong non-monotonic relationship across different sex ratios. This is because, when sex ratios are skewed, the mean mating success of one sex is higher than the other resulting in negative values of \( r_{\text{Newman}(U)} \). For ‘mating density’ simulations, we demonstrated that NODF is strongly positively correlated with ‘mating density’ (Almeida-Neto et al. 2008). However, the choice of null models used to simulate matrices may not be straightforward and would change the interpretation of empirical NODF values, to a deviation from randomness (Ulrich and Gotelli 2007; Strona and Fattorini 2014).

Overall, the two measures that were most consistent across all population parameters were \( r_{\text{Newman}(D)} \) and SCIC. Both measures correlated strongly and positively with each across all population parameters, suggesting they are likely to capture similar variation. However, arguably only SCIC has a clear relationship to sexual selection theory. This is because unlike \( r_{\text{Newman}(D)} \), SCIC can be directly included in the partitioning of the Bateman gradient into those components that reflect the

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**Fig. 5** The relationship between measures of assortative mating structure (\( r_{\text{Newman}(U)} \) (a–c), \( r_{\text{Newman}(D)} \) (d–f), SCIC (g–i) and NODF (j–l) and population level parameters (population size (a, d, g, j)), sex ratio (males/females) (b, e, h, k) and ‘mating density’ (c, f, i, l) for randomly simulated mating populations. Numbers on the bottom right are the Spearman rank correlation coefficients.
independent effect of mating success and the independent effect of sperm competition on male reproductive success. Furthermore, SCIC can be used directly to quantify the contribution of polyandry and assortativity in mating patterns to Bateman gradients, by multiplying SCIC and the partial regression coefficient of the sperm competition intensity faced by a male ($\beta_{\text{SCI-M}}$) (Fig. 2, Eq. 6).

An important caveat is that our approach calculates SCI assuming that each mating has the same competitive value (equality of mating). This assumption may generate simplistic expectations in terms of variation in paternity share. Variation in male traits such as ejaculate size, factors that load the sperm competition raffle (e.g., mating order) and mechanisms of cryptic female choice (Birkhead and Møller 1998; Simmons 2001) can all influence variation in paternity share and generate deviations from the predictions of our null model assuming of equality of mating. Such processes may themselves erode the predictive power of SCI in determining a male’s paternity share, as the number of males that inseminate a female becomes less important in determining male fertilisation success relative to other factors, e.g. the timing of insemination or ejaculate traits (Parker and Pizzari 2010). This may in turn reduce the predictive power of SCI. Such cases will themselves provide interesting empirical studies in understanding how the pattern of female multiple mating determines the operation of sexual selection. A particularly relevant example arises when individual partners copulate with each other multiple times (i.e. edges between males and females nodes are weighted by the number of copulations). Remating between male and female pairs may reflect postcopulatory selection on males and function as a trait that allows a male to defend his paternity by increasing the relative representation of his ejaculates versus competitors (Shuster and Wade 2003). In such situations, the calculation of a male’s SCI can be extended to include multiple copulations between individual males and females. Using this weighted approach, a male’s exclusivity with a female partner is determined not by the number of competitor males but instead by the relative representation of his ejaculates (Shuster and Wade 2003). This weighted version of male SCI is therefore shaped by both the remating rate of the focal male and the mating rates of competitor males, i.e. a low weighted SCI can be the result of both high focal male remating rate and low remating rates of other competitor males. Weighted and unweighted versions of SCIC can be compared to explore how patterns of remating accentuate or diminish the relationship between male mating success and the intensity of sperm competition.

The simulation approach used in this work has provided hypothetical conditions in which the researcher has detailed and complete knowledge of the mating patterns of a population. However, in many empirical studies, researchers may have less detailed knowledge, e.g. when mating success is inferred from genetic parentage assignment with little or no behavioural data (Collet et al. 2014; Taylor et al. 2014). Using genetic data alone may result in overestimates of variance in male mating success and Bateman gradients (Collet et al. 2014). With reference to the estimation of assortative mating structure, we expect that using genetic parentage as a means of determining male–female mating networks may also bias results. When behavioural data are not available, genetically determined mating networks may not accurately characterise the distribution of sperm competitive environments across males and instead may better represent the results of sperm competition rather than the competitive landscape, potentially underestimating the assortativity of a given mating network. This problem may be exacerbated when female clutch size is small (Collet et al. 2014). An extreme example comes from cases where females only lay one egg. In this case, using genetic parentage assignment removes completely the potential for female polyandry and any level of mating structure; however, the distribution of reproductive success across males may strongly depend on female polyandry and mating structure in such ‘winner fertilise all’ cases. A similar issue relates to the accuracy of behavioural data when available. Missing data may result in an underestimation of polyandry and bias estimates of mating network structure, which may be particularly strong when males of a given mating success are associated with a particular reproductive tactic that is consistently not recorded (e.g. alternative reproductive tactics; sneak mating). We therefore advocate that use of such methods requires both genetic and detailed behavioural data on copulations or genetic parentage assignment only in those cases where genetic data are strongly representative of behavioural mating patterns. Although these conditions may appear restrictive, there is great potential to explore these methods in experimental groups where behavioural data are more readily available (e.g. Collet et al. 2012; Pélissié et al. 2014) and in some

Fig. 6 Plots show the Pearson product correlation coefficient between measures of assortative mating structure ($r_{\text{Newman (U)}}$, $r_{\text{Newman(D)}}$, SCIC and NODF) across all levels of population level parameters (population size, sex ratio (males/females) and ‘mating density’ for randomly simulated mating populations.
natural populations of both vertebrates and invertebrates, where detailed behavioural data are becoming available (e.g. Preston et al. 2005; Rodríguez-Muñoz et al. 2010).

Finally, although this study has focused largely on the utility of assortment measures in studies on male Bateman gradients, assortment measures such as SCIC may also have useful applications for studies focusing on female reproduction. Consider, for example, populations where males are sperm limited and so those male that mate many times may fail to deliver sufficient sperm per copulation to fertilise all the ova of a female (Warner et al. 1995; Wedell et al. 2002). Patterns of positive assortative mating may then weaken female Bateman gradients, because increasing mating success is associated with fewer sperm delivered per copulation. Whereas in negatively assorted mating network, female Bateman gradients would be strengthened because females with few mates suffer reduced fertility as their partners tend to be the most promiscuous, and so potentially the most sperm depleted.

This work merges network theory from both social sciences and ecological literature to test the utility of a variety of measures in quantifying the relationship between male mating success and the mating success of their female mating partners, for use in the study of sexual selection and sperm competition. Overall, our results lead us to suggest that the measure of SCIC is the most promising approach as it is both logically consistent with the relationship between male mating success and the mating success of their partners, is not strongly confounded by variation in population parameters and has a clear relationship to sexual selection theory.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no competing interests.

Ethical approval This article does not contain any studies with human participants or animals performed by any of the authors.

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