Sampling scale can cause bias in positive assortative mating estimates: The first evidence in two intertidal snails

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Abstract

Assortative mating in the wild is commonly estimated by correlating between traits in mating pairs (e.g. size of males and females). Unfortunately such an approach may suffer from considerable sampling bias when the distribution of different expressions of a trait in the wild is non-random; for example, when segregation of different size classes of individuals occur in different microhabitats or areas. Consequently, any observed trait correlation in the wild can be an artifact of pooling heterogeneous samples of mating pairs from different microhabitats or areas rather than true non-random matings. This bias in estimating trait correlations due to sampling scale is termed the scale-of-choice effect (SCE). Here we use two intertidal littorinid species from Hong Kong to show how the SCE can bias size-assortative mating estimates from mating pairs captured in the wild, empirically demonstrating the influence of this effect on measures of positive assortative mating. This finding cautions that studies that have overlooked SCE may have misinterpreted the
magnitude and the cause of assortative mating, and we provide a new analytical approach to protect against this potential bias in future studies.

Keywords: SCE; mate choice; mating pair; size-assortative mating; mating preference; *Echinolittorina* spp.

**Introduction**

The decisions that dioecious organisms take in choosing their mates has key evolutionary importance, as these choices influence the probability of allele transmission and distribution of genotypes in the next generation (Lewontin et al., 1968; Coyne and Orr, 2004; Gavrilets, 2004). Mating preferences can be investigated by detecting deviations from random mating (i.e. assortative mating) in traits which are used to distinguish between individuals such as size and colour (Jiang et al., 2013). Perhaps the most simple and common approach to detect non-random mating in the wild is by studying any correlation between traits (assessed using Pearson’s $r$) of the observed mating pairs (Crespi, 1989; Arnqvist et al., 1996; Jiang et al., 2013). Trait correlations can be positive or negative, when there is a tendency of individuals mating preferentially with members of the opposite sex with similar or dissimilar trait values, respectively. Observed correlation of traits in mating pairs in the wild can, however, be the result of a variety of processes such as mating preferences and mating constraints (Crespi, 1989). Non-random distribution of different expressions of a trait may, however, also influence observed mating patterns (Arnqvist et al., 1996; Jiang et al., 2013), and a recent study has illustrated how such spatial patterns can lead to sampling bias and hence confound the assessment of mating patterns in the wild. This effect has been termed the scale-of-choice effect (SCE; Rolán-Alvarez et al., 2015),
and occurs when the sample of mating pairs is measured at a larger spatial scale than
the scale of mate choice in the organism and when there is spatial heterogeneity in
trait distribution at the true scale of mate choice (see Figure 1a). In the example (Fig.
1a), positive size-assortative mating may be ‘detected’ because different sized pairs
from different localities (e.g. sections of a seashore) have been measured and pooled
during estimation. This pooling of samples over an spatial scale inappropriate to the
movement range of the mating individuals will lead to a biased estimate of correlation,
or other similar statistics that may be used to estimate assortative mating
(Rolán-Alvarez et al., 2015), calculated at the wrong spatial scale for the hypothesis
being tested (Hassler and Thadewald, 2003). The practical consequence of this
phenomenon is that many correlation coefficients estimated from measurements in the
wild may be subject to this effect and, therefore, their support for the hypotheses
tested, and subsequent interpretation could be misleading, as the SCE can obscure the
true nature of any trait correlation.

Although the true scale of choice is usually unknown in most species, grouping
mating data according to the frequency of different expressions of a trait (e.g.
according to size classes or shell colour) of individuals surrounding mating pairs can
reduce the influence of spatial heterogeneity and, therefore, diminish or eliminate the
bias caused by SCE. As the SCE has two necessary conditions; firstly that there is a
mismatch between the scale over which individuals can make a choice and the scale at
which sampling occurs; and secondly that there is spatial heterogeneity at the scale of
sampling; diminishing or eliminating the effects of either condition will affect the
magnitude of SCE (Rolán-Alvarez et al., 2015), and therefore the bias affecting the
estimated correlation coefficient (r). Rolán-Alvarez et al., (2015), for example, have
shown that pooling samples of mating pairs of the intertidal snail, *Littorina fabalis,*
from three localities with similar trait (colour) frequency increased the estimated r when compared with non-pooled samples. The true strength of the negative mating preference for distinct shell colours was, therefore, accentuated due to the inappropriate analysis of the pairs. As such, a systematic bias in any meta-analysis on assortative mating may occur whenever the studied species has small dispersion ranges and the SCE is expected to be high. In reality, negative assortative mating is apparently a rare phenomenon in natural populations (Jiang et al., 2013), but generality may be an artifact of SCE which has not been accounted for in previous meta-analyses (Rolán-Alvarez et al., 2015). It is also desirable to investigate whether the SCE can also bias cases of positive assortative mating.

Positive size-assortative mating is common in gastropods and many other organisms (Erlandsson and Johannesson, 1994; Staub and Ribi, 1995; Erlandsson and Rolán-Alvarez, 1998; Rolán-Alvarez et al., 1999; Zahradnik et al., 2008; Avaca et al., 2012; Ng & Williams, 2012; see Jiang et al., 2013 for a review). In intertidal snails this mating pattern may be caused by males following female mucus trails in a size-dependent manner, which would result in positive assortative mating for size (Conde-Padín et al., 2008; Johannesson et al., 2008; Ng et al., 2013; Ng & Williams, 2014), but other mechanisms based on preferential mating can also be invoked to explain this pattern (Saur, 1990; Hull, 1998; Johannesson et al., 2008; Zahradnik et al., 2008; Saltin et al., 2013). Given their limited locomotion capacity relative to the habitat they occupy, it is reasonable to assume that snail species exhibit their choice at rather small scales, and therefore there is a strong potential for SCE to influence estimates of trait correlation coefficients using mating pairs captured in the wild. This theory was investigated using two tropical intertidal snails to determine whether SCE might influence assessment of their assortative mating patterns.
**Material and Methods**

**Study localities and sampling approach**

_Echinolittorina malaccana_ and _E. radiata_ are common grazers on the high shore and their distributions on the shore overlap, with _E. malaccana_ generally found above _E. radiata_ (Mak, 1996). Measurements were made in August 2015, during the snails reproductive season (Mak, 1996) at Cape D’ Aguilar Marine Reserve (22° 12’ 27” N, 114° 15’ 33” E) and Shek O (22° 13’44”N, 114° 15’ 22”E), Hong Kong (Fig. 1b). These two species were scored at seven localities (CD1 to CD7) in Cape D’ Aguilar and one in Shek O (SO) with relatively high densities of either one or both species (all localities for _E. malaccana_, and CD1, 2, 6 and 7 for _E. radiata_, Fig. 1b). These localities were all within 10-m stretches of the shoreline, and were separated by 10s to 1000s of metres in order to investigate the SCE (Fig. 1a).

Mating pairs were collected within the same tidal phase at each locality. In addition to the mating pairs, the four closest unmated conspecifics surrounding each mating pair were also scored to define a microarea unit (within a ~250 mm radius of each mating pair representing a small portion of area in each locality). Every mating pair plus the four unmated conspecifics, therefore, represents a different microarea replicate. All (4-80 microarea replicates among the localities) mating and nonmating snails were taken to the laboratory where sex (determined by the presence or absence of a penis as seen under dissection microscope) and shell length (± 0.1 mm, vernier calipers) were recorded. Previous studies have demonstrated very low parasite loading of littorinids in Hong Kong (maximum 0.6%, Tang, 1995), and hence this possible confounding effect on mating behaviour and mating pattern was considered to be insignificant. All snails collected were returned to their original shores after
investigation.

Statistical analyses

Size-assortative mating was estimated by Pearson’s correlation coefficient (r) among the observed mating pairs, and the significance evaluated by the non-parametric Kendall-tau-b test. The SCE is defined as the correlation coefficient of a trait in the pooled sample (r_{pooled}) minus the averaged correlation (r_{averaged}) of the trait among homogeneous sets of data (Rolán-Alvarez et al., 2015 see Figure 1a), and in this case, r_{pooled} refers to the correlation coefficient between sizes derived from pooled mating pairs of all localities in both Cape D’ Aguilar and Shek O, whereas r_{averaged} refers to the mean correlation coefficient derived from mating pairs in each homogeneous set of size classes (i.e. microareas that share similar sizes classes of individuals including the mated and unmated conspecifics). As such, SCE estimates the possibility of bias in estimating the correlation coefficient due to non-random distribution of different size classes among the localities by taking into account individual sizes in each microarea. Three to five homogeneous sets of size classes were used in the analyses, derived from the mean individual size in each microarea. The r_{averaged} was, therefore, calculated over these three to five homogeneous sets of size classes. The significance of the SCE was evaluated by comparing the r_{averaged} against the r_{pooled} value as null value by a t test. All analyses were performed with SPSS 20.0 (SPSS Inc., Chicago, IL, U.S.A).

Results and Discussion

When SCE was not considered by grouping data into homogenous sets of size classes, the calculated correlations supported the presence of positive size-assortative mating
in both species (Table 1). The averaged $r$ across all localities was positive and
significant ($p < 0.05$) in *E. radiata* and positive and marginally non significant ($p = 0.053$) in *E. malaccana*. In addition, there was at least one locality where significant, positive size-assortative mating was recorded in both species (Table 1) and the pooled samples among different homogeneous sets of size classes clearly demonstrated size-assortative mating. This pattern is typical for many littorinid species, and positive size-assortative mating is assumed to be the underlying mechanism (Erlandsson and Johannesson, 1994; Erlandsson and Rolán-Alvarez, 1998; Johnson, 1999; Rolán-Alvarez et al., 1999; Ito and Wada, 2006; Zahradnik et al. 2008; Ng & Williams, 2012). This assumption, however, may be incorrect if sampling biases such as the SCE are not taken into account (Rolán-Alvarez et al., 2015). Grouping the mating pairs in three to five homogeneous sets of size classes within the two study sites, for example, indicated close to random mating patterns in the two species (Table 1). The Pearson correlation coefficient averaged across the homogeneous sets of size classes was 0.21 (for 3 classes), 0.12 (for 4 classes) and 0.06 (for 5 classes) in *E. malaccana*, with 3 out of 6 estimates being significantly different from $r_{\text{pooled}}$, and the other 2 out 6 being marginally different, with the probability being close to 0.05. The SCE, and hence the bias in the estimation of correlation coefficients based on five homogeneous sets of size classes, was as large as 0.5 and 4.7 in Shek O and Cape’ D Aguilar respectively in *E. malaccana* (Table 1). The SCE was even more clear in *E. radiata*, as the correlation coefficients across averages was always significantly different from the pooled estimate, yielding a SCE of 0.54 (Table 1). In both species, therefore, the assumed trend towards positive assortative mating when mating pairs are pooled from multiple localities was likely an artifact due to the pooling of heterogeneous samples (i.e. mating pairs from different microareas consisting with individuals of different
size classes) rather than a true non-random mating pattern. To try and avoid this bias, we propose that investigators should analyse mating pairs in homogeneous subgroups, which can reveal whether the SCE is contributing to the estimates of correlation coefficients to detect relationships between traits. This approach requires large sample sizes as well as information about conspecifics close (i.e. within their movement range) to the mating pairs. Although the calculated correlation coefficients and hence SCEs are somewhat sensitive to the number of homogeneous subgroups used (3 to 5 in the present case; Table 1), this approach does provide an appropriate method to evaluate the potential bias due to the scale of sampling area.

As previous authors have pointed out, the study of assortative mating and its causes seems fraught with difficulties, especially as the problem has been defined and considered in different ways from theoretical and empirical frameworks (Gavrilets, 2004; Roff and Fairbairn, 2015). Theoretical mathematical functions to simulate assortative mating \textit{in silico}, for example, have recently suggested that assortative mating can be a consequence of either an increase of mating preference \textit{per se}, or by a drift in the distribution of the trait being used to describe the preference (Carvajal-Rodríguez et al., 2014). The difference between these two mechanisms in driving assortative mating is subtle but rather important in evolutionary terms, since only a genetic change in mating preference is evolutionary relevant for reinforcement (Kirkpatrick, 2000), speciation and/or sexual selection (Gavrilets, 2004), and we presently do not have the tools that distinguish between these phenomena (Carvajal-Rodríguez et al., 2014). Although the analytical approach provided here, which estimates assortative mating independently of the trait distribution, is not a perfect solution it does, however, provide an estimate of sampling bias and this alone is a considerable advantage over more traditional approaches (Jiang et al., 2013).
While this study demonstrates the need to consider sampling bias in estimating assortative mating, even greater efforts will be needed in the future to understand the true causes of this mating pattern.
Acknowledgements

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References


Figure 1. 1a) The scale-of-choice effect (SCE) results in this example when pooling a heterogeneous sets of subsamples (S₁ to S₅) with random mating within each (represented by the pairs within circles), causing an overall positive assortative mating in the pooled set of samples (S_pooled; represented by the ellipse). Note that the analytical approach proposed here consists of estimating correlation coefficients (r_averaged) from homogenous sets of size classes (Class₁=S₁, S₂ and S₅; Class₂=S₃ and S₄ in this example). The SCE would be r_pooled minus r_averaged. 1b) Sampled distribution of mating pairs captured in Shek O (SO) and Cape D' Aguilar (CD, inset), Hong Kong.
Table 1. Analysis of the scale-of-choice effect (SCE) for size assortative mating in *Echinolittorina malaccana* and *E. radiata*. The Pearson correlation coefficient ($r \pm SD$) for size of mates is provided within sample, pooled samples and different homogeneous sets (3-5) of samples within localities (SO or CD). The significance of the correlation coefficient, $r$, is given by the non-parametric Kendall-tau-b correlation test within sample and pooled ($r$ and $r_{pooled}$), and by a t-test for means ($r_{averaged}$; checking the null hypothesis = $r_{pooled}$). SCE is only calculated for the case when mating pairs are grouped according to five homogeneous sets of size classes.

<table>
<thead>
<tr>
<th>Species</th>
<th>Locality</th>
<th>All samples</th>
<th>( r_{averaged} ) across homogeneous sets of size classes</th>
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<tr>
<td></td>
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<td>N ( r ) ( N ) ( r_{pooled} ) 5 classes</td>
<td>4 classes</td>
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<tr>
<td>SO</td>
<td>40</td>
<td>0.52***</td>
<td>40 ( 0.52*** )</td>
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<tr>
<td>CD1</td>
<td>13</td>
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</tr>
<tr>
<td>CD2</td>
<td>40</td>
<td>-0.21</td>
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<tr>
<td>CD3</td>
<td>4</td>
<td>0.46</td>
<td></td>
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<td>E. malaccana</td>
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<tr>
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<tr>
<td>CD7</td>
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<td>mean</td>
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<td>0.25* ± 0.300</td>
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<th></th>
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<td>6 ( 0.90^* )</td>
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<td></td>
<td>CD2</td>
<td>31 ( 0.13 )</td>
<td>49 ( 0.67*** )</td>
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<tr>
<td></td>
<td>CD6</td>
<td>4 ( 0.016 )</td>
<td></td>
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<tr>
<td>E. radiata</td>
<td>CD7</td>
<td>8 ( 0.379 )</td>
<td></td>
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<tr>
<td>mean</td>
<td></td>
<td>0.36* ± 0.395</td>
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* \( p \leq 0.05; \)** \( p \leq 0.01; \)**\( p \leq 0.001; \)? \( p \leq 0.10 \)