

The Limits of Mean-Field Heterozygosity Estimates under Spatial Extension in Simulated Plant Populations

James L. Kitchen, Robin G. Allaby*

School of Life Sciences, University of Warwick, Coventry, United Kingdom

Abstract

Computational models of evolutionary processes are increasingly required to incorporate multiple and diverse sources of data. A popular feature to include in population genetics models is spatial extension, which reflects more accurately natural populations than does a mean field approach. However, such models necessarily violate the mean field assumptions of classical population genetics, as do natural populations in the real world. Recently, it has been questioned whether classical approaches are truly applicable to the real world. Individual based models (IBM) are a powerful and versatile approach to achieve integration in models. In this study an IBM was used to examine how populations of plants deviate from classical expectations under spatial extension. Populations of plants that used three different mating strategies were placed in a range of arena sizes giving crowded to sparse occupation densities. Using a measure of population density, the pollen communication distance (P_{cd}), the deviation exhibited by outbreeding populations differed from classical mean field expectations by less than 5% when P_{cd} was less than 1, and over this threshold value the deviation significantly increased. Populations with an intermediate mating strategy did not have such a threshold and deviated directly with increasing isolation between individuals. Populations with a selfing strategy were influenced more by the mating strategy than by increased isolation. In all cases pollen dispersal was more influential than seed dispersal. The IBM model showed that mean field calculations can be reasonably applied to natural outbreeding plant populations that occur at a density in which individuals are less than the average pollen dispersal distance from their neighbors.

Citation: Kitchen JL, Allaby RG (2012) The Limits of Mean-Field Heterozygosity Estimates under Spatial Extension in Simulated Plant Populations. PLoS ONE 7(8): e43254. doi:10.1371/journal.pone.0043254

Editor: Dirk Steinke, Biodiversity Institute of Ontario - University of Guelph, Canada

Received: April 27, 2012; **Accepted:** July 18, 2012; **Published:** August 27, 2012

Copyright: © 2012 Kitchen, Allaby. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work is supported by the Leverhulme Trust F/00 215/BC. The funder had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: Robin Allaby is an Editorial Board Member of PLoS ONE. This does not alter the authors' adherence to all the PLoS ONE policies on sharing data and materials.

* E-mail: r.g.allaby@warwick.ac.uk

Introduction

Understanding the evolutionary process is increasingly requiring in the integration of sources of data that are typically beyond classical population genetics models [1]. One such example is the inclusion of spatial extension. Classical population genetics approaches typically use a simplified mean-field approach in which individuals of a population or subpopulation conceptually occupy the same space, and are therefore subject to the same conditions and pressures, and purely random mating occurs. Along with other simplifying assumptions such as non-overlapping generations and constant population sizes, such populations behave in tractable ways that can be described through deterministic approaches leading to features such as Hardy-Weinberg equilibria [2] and the Wright-Fisher model [3]. The predictions of such models provide a useful starting point for evolutionary studies, for instance in establishing whether there has been significant deviation from neutrality indicative of selection. However, Mayr [4] observed that it was surprising how little classical population genetics has contributed to the understanding of one of the most important processes in evolution, speciation. This is because a mean-field based model is essentially based on an anagenic evolutionary system, rather than a cladogenic one [5]. The evolutionary differentiation of populations, which ultimately leads to speciation, requires different selective environments.

In reality populations are spread across environments that are often patchy with different sets of conditions. Environmental conditions may have a variety of effects, such as impeding dispersal [6] or providing different selective niches [7]. Plant mates are unlikely to be entirely chosen at random. For instance, plants have been known to have assortative mating due to pollinator foraging behaviour regarding petal color [8], inflorescence height [9] and overlapping flowering times [10]. Furthermore the spatial dimension is of particular importance even within a single constant environment because of their sedentary nature causing a bias for nearest-neighbor mating over panmixia [11]. The spatial density of a plant species affects the optimal mating strategy with higher out-crossing rates being more suited to higher densities [11] and selfing systems evolving to invade low-density environments [12]. The boundaries and margins between contrasting selection environments may be areas in which globally rare but locally abundant alleles occur through gene flow between areas in which they have high selective value to areas of low or neutral selective value, giving rise to processes such as the persistence of unfit alleles which would not be detected in mean field based model systems [1]. A problem with spatially extended plant populations in nature recently highlighted is that it is unclear to what extent classical population genetics tools can meaningfully be applied at all when individuals are continuously distributed across space [13].

It is therefore desirable to include the dimension of spatial extension into population genetics models in order to effectively incorporate these factors and more accurately reflect the evolutionary process. Individual-based modeling (IBM) is a versatile approach for the integration of non-parametric and complex factors into population genetics. Spatial extension can readily be incorporated into IBM and has been successfully applied to a variety of problems including plant domestication [14,15], predator-prey relationships [16–20] and the emergent field of landscape genetics [21–23]. As ecological boundaries, environmental conditions and varying dispersal distances can be readily incorporated into these models they have been used to study speciation events [24–26] and hybrid zone formation [27]. Furthermore as these models readily allow individual behaviors to be defined they have been useful for simulating gender specific mating preferences and the resultant effects of assortative mating on speciation and hybridization [28–29]. Cuddington and Yodzis [16] demonstrated that there is a reduced mobility in spatially explicit relative to mean-field models, resulting in reduced reproduction rates in the former. By reducing mobility parameters of the mean-field model, a close congruence between the two model types was achieved. The differentiating effects of spatial extension are likely to increase as individuals become more distantly spaced out. However, it is still unclear how the relationship between the two systems changes with increasing spatial extension.

We have developed an IBM to study the evolution of plant systems in order to integrate multiple diverse data sources and structures [1]. In brief, the IBM is implemented as follows: diploid individuals are arranged onto a two-dimensional grid made up of cells each of which can hold a single adult plant. The individuals contain genes that can accumulate neutral mutations and gametes are generated by Mendelian segregation of homologous chromosomes. The simulation updates in intervals each representing one month. Individuals begin as seeds and may germinate with a defined probability on condition of the absence of an adult in the cell. The individuals then move through the following life cycle stages: seed, vegetative growth, flowering and senescence, with the duration of each life cycle stage varying between individuals. Flowering individuals disperse pollen and seeds and may self or out-cross according to a user-input probability. Individuals senesce and are removed a number of updates after dispersing their seeds determined by randomly sampling.

In this study we aimed to determine under what conditions, if any, parameters estimated by classical population genetics approaches can be applied approximately to establish the presence of conditions of equilibria and neutrality in a spatially extended system. The IBM was used to generate plant populations of varying spatial densities, mating systems, and pollen and seed dispersal capabilities. We assessed the extent of deviation between classical population genetics and the spatially extended system by comparing values of the heterozygosity parameter. Heterozygosity was measured as expected from classical population genetics through the Hardy-Weinberg law (H_c), and compared to the real value obtained from individuals (H_o). As far as we are aware this is the first attempt to determine how perturbations of spatial extension affect the accuracy of mean-field based calculations of population parameters.

Results

All simulations in this study began with populations of 1000 individuals in a two dimensional matrix with M cells, in which a single cell could contain a single growing plant. Simulations were

carried out for 4000 updates with a burn-in period in which populations were allowed to equilibrate (see methods). For each set of conditions, simulations were repeated ten times. To prevent individuals from forming clusters with each other [30], we used a strategy in which cells were randomly blocked out (void cells) within all matrices from all simulations, such that they could not be occupied until just 1000 cells remained. On average this achieved an even spacing between individuals in our simulations with a greater degree of spacing between individuals within the larger matrices, Figure 1. In preliminary simulations where blocking was not applied we observed clustering behaviour even within larger matrices (data not shown).

Mating System

The first set of simulations investigated the effect of mating system in a spatially explicit population. In this case a matrix size of 35^2 cells was used in which 225 cells were randomly blocked. In this matrix size most individuals were immediately adjacent to other individuals in all eight surrounding cells. The model takes a user input of the probability of self-pollination, S_u , to allow control over the outcrossing exhibited by flowering individuals. The value S_u is the probability that a particular pollen grain will self-fertilize the parent plant rather than leave the flower and pollinate another plant. Simulations were carried out with S_u ranging in value from 0 to 1, Figure 2. At values of S_u at 0 and 0.001, the resulting values of H_o and H_c overlapped, and at a S_u of 0.1 were within 5% of each other suggesting H_o gave a good approximation of H_c . S_u equal to 0 is representative of a self-incompatible mating strategy, while a value of 0.001 represents a level of out-crossing equal to that found in panmictic systems. Increasing values of S_u in the spatially extended system resulted in a reduction in heterozygosity, as would also be expected of a mean field system [31].

Matrix Size

We investigated the effect of increased separation between individuals in a spatially extended population. Simulations were carried out with progressively increased matrix sizes ranging from 35^2 to 135^2 cells, with increments of 10 in both matrix dimensions between each set of simulation conditions, Figure 1. Each simulation was performed with $M - 1000$ void cells, allowing the population size to remain constant between different sets of simulation conditions. Three series of simulation experiments were carried out with different mating strategies defined by values of S_u of 0.001, 0.1 and 0.9, respectively, Figure 3. We used the parameter pollen communicating distance (P_{cd}) to express the degree of separation between individuals defined as:

$$P_{cd} = nm_d / P_d \quad (1)$$

Where nm_d is the average distance to the nearest neighbor in matrix cells, and p_d is the average dispersal distance of pollen in matrix cells. We also measured a second parameter, the seed communicating distance (S_{cd}) which is defined as:

$$S_{cd} = nm_d / S_d \quad (2)$$

Where S_d is the average dispersal distance of seeds in matrix cells. Constant values of maximum dispersal of S_d and P_d were used of 10 cells.

The population size per update in simulations with values of M over 13225 and S_u 0.001 was often low, leading to an increased variation of H_o which ranged in standard deviation 0.015–0.064. At smaller matrix sizes the standard deviation ranged

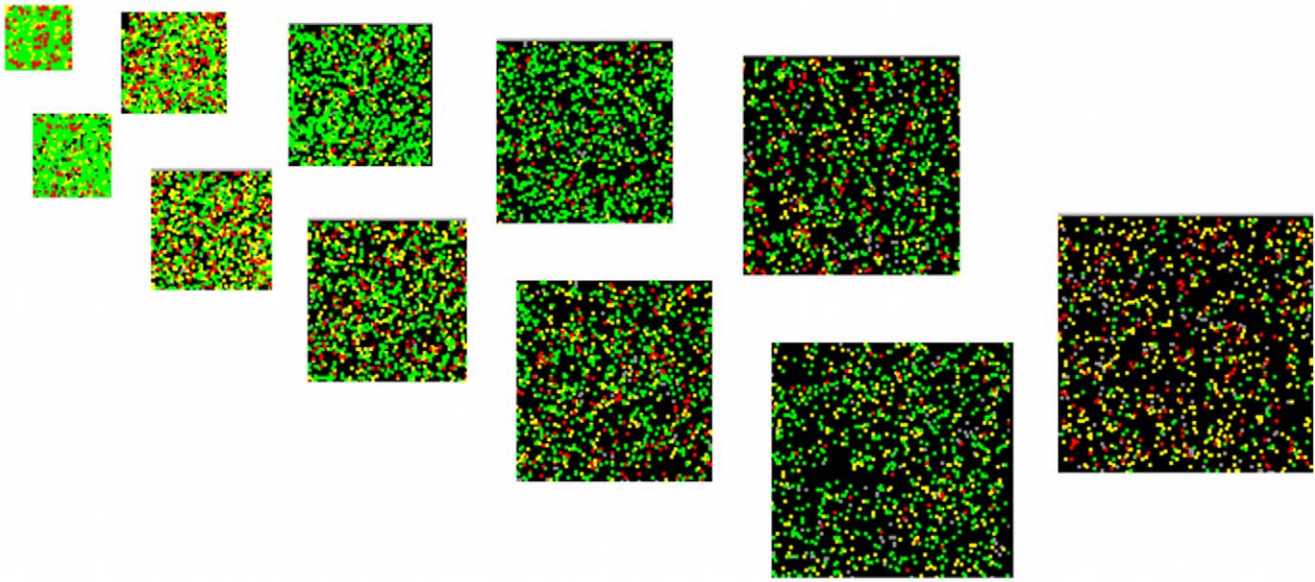


Figure 1. Model screen shots of the different matrix sizes used in the study. The void cells are in black with grey cells representing unoccupied non-void cells. The different colors of the individuals represent their different life-cycle stages, with red = seeds, green = vegetative and yellow = flowering/senescence.
doi:10.1371/journal.pone.0043254.g001

$4 \times 10^{-4} - 0.008$. This increase in variation was probably due to high mortality and low fertility rates under these conditions in the model resulting from low numbers of nearby vacant cells for seeds

to disperse to, and low numbers of nearby pollinating individuals (not shown). H_o decreased as the P_{cd} increased with matrix size. The relationship between P_{cd} and H_o in Figure 3 varied between

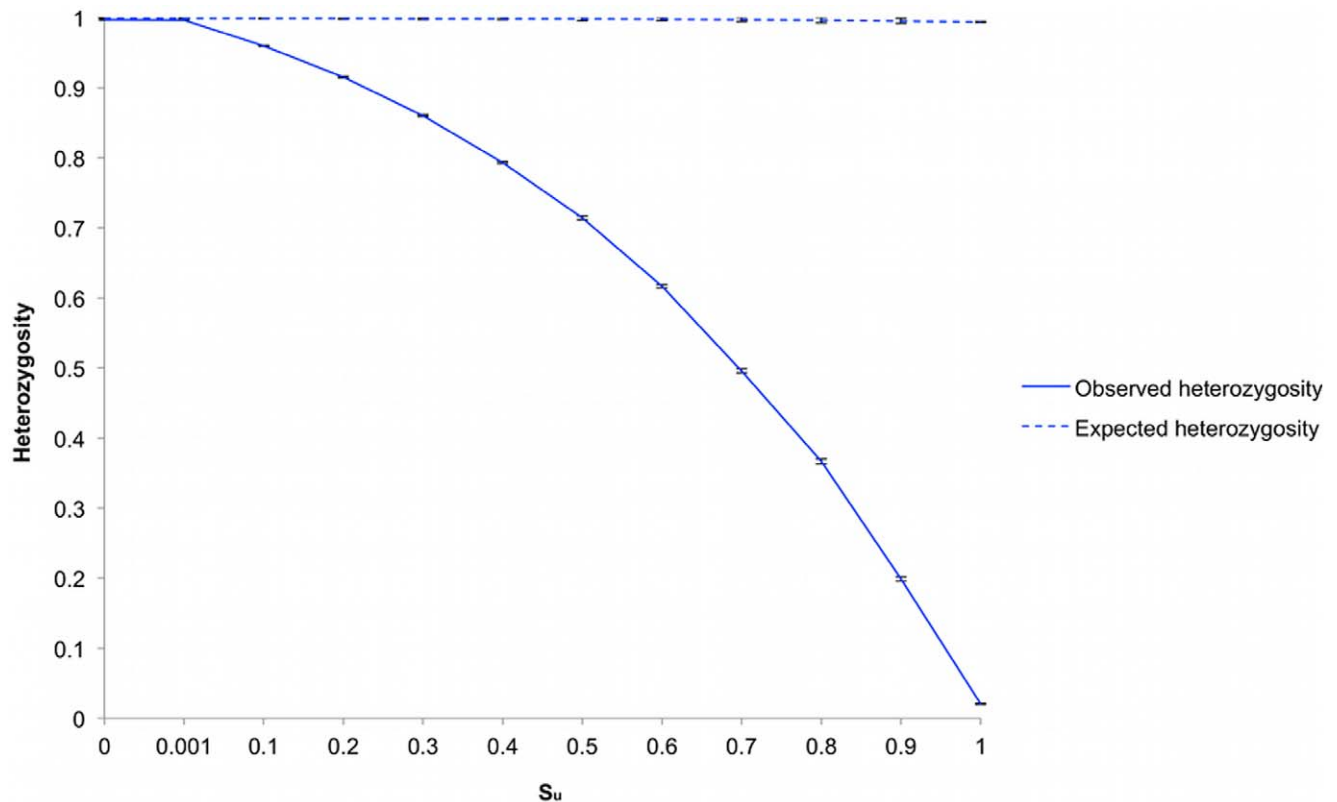


Figure 2. Heterozygosity as a function of selfing probability. H_o is plotted with the solid line and H_e using a dashed line. Simulations were repeated ten times. Error bars represent the standard error.
doi:10.1371/journal.pone.0043254.g002

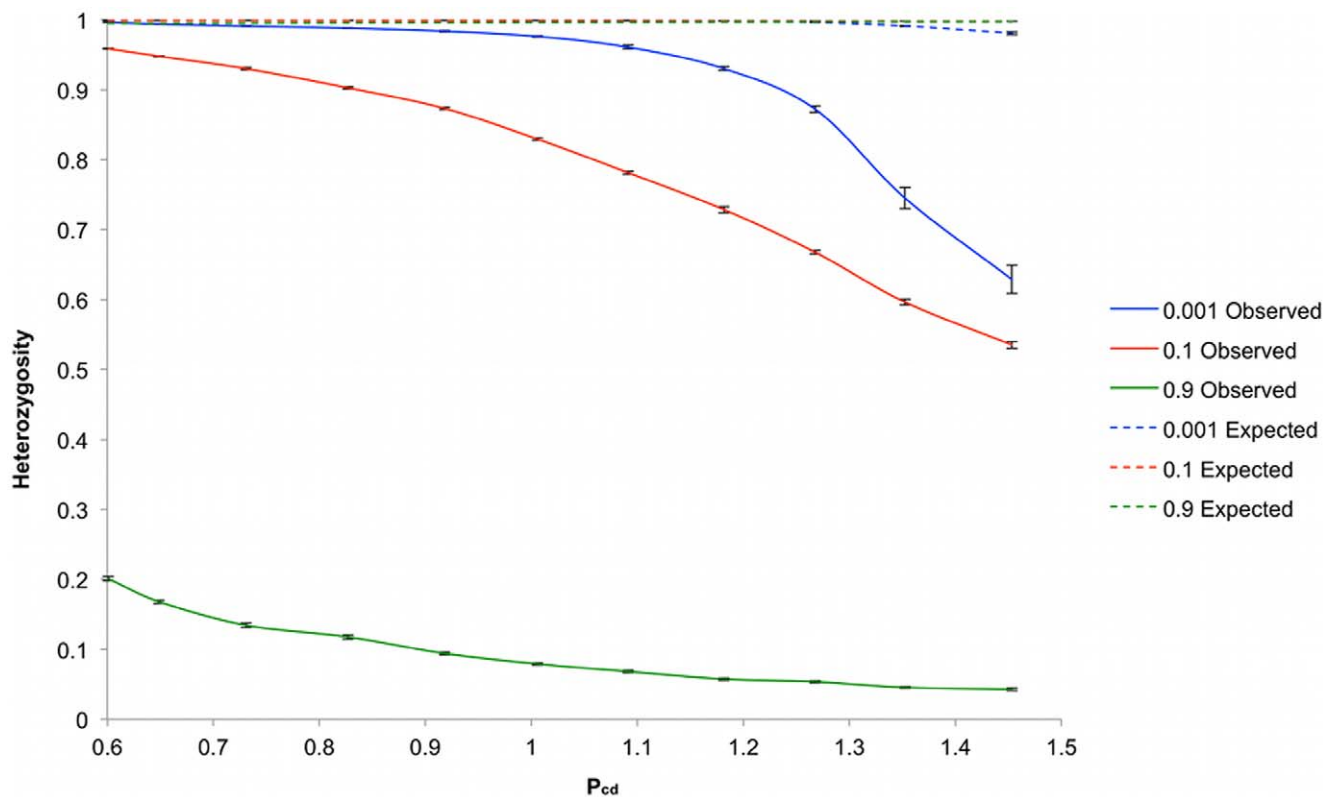


Figure 3. Heterozygosity as a function of matrix sizes. Heterozygosity was plotted at $S_u = 0.001$ (blue), $S_u = 0.1$ (red) and $S_u = 0.9$ (green). H_o is plotted with the solid line and H_e using a dashed line. Simulations were repeated ten times for each value of S_u , with error bars representing the standard error.

doi:10.1371/journal.pone.0043254.g003

the three mating systems tested. In the case of S_u equal to 0.001, the rate at which H_o decreased with increased P_{cd} , or r , as we shall use in the text, is given by the gradient of the line, which varies either side of a P_{cd} value of 1.1. At values of P_{cd} lower than 1.1 r is relatively shallow at $[-0.005]$, and steepens considerably to $[-0.103]$ at P_{cd} values higher than 1.1. In contrast, in simulations where S_u was equal to 0.1, an approximately linear relationship occurred between H_o and P_{cd} in which r was relatively constant. At the highest values of P_{cd} these two sets of simulations produced similar values of H_o . In the third set of simulations where S_u was equal to 0.9, values of H_o decreased to an asymptote.

The S_{cd} parameter describes the movement of seeds in relation to the spatial density of plants, and we expected it to have some influence on heterozygosity also. At the minimum matrix size a S_{cd} of 0.9 occurred, while at maximum matrix size this value was 2.2. Therefore, in the smallest matrix system the dispersal ranges of seeds encompassed neighboring individuals, but not in the case of the largest matrix suggesting that seed movement was contributing to gene flow in the former, but much less in the latter.

A threshold effect is apparent in these results, in which under a panmictic mating system deviation from the classical expectation is small, typically within 5%, while plants are within a range of their nearest neighbor, with an average nearest neighbor distance less than the average pollen dispersal distance. After this threshold the deviation from classical expectations increases rapidly with increasing separation. However, a more mixed mating strategy shows a steady deviation from classical expectation with increasing matrix size indicating that spatial separation has a more direct effect in these cases. Interestingly, these two sets of simulations reach similar levels of heterozygosity at the maximum extent of

separation investigated in this study. The isolation caused by the highly selfing system has a large effect on heterozygosity even in small matrices with low P_{cd} values, which then decreases to a minimum equilibrium which is probably kept from reaching zero by the influx of new mutations. In these cases, the selfing mating strategy is the principal influence on heterozygosity.

Pollen and Seed Dispersal

An alternative to increasing the space between individuals to increase spatial isolation is to reduce dispersal of pollen and seeds. If there are no other influencing factors introduced by increased spacing of individuals, then altering the dispersal behavior should allow similar levels of heterozygosity to be achieved under differentially extended systems. We tested this prediction by performing simulations at the largest matrix size ($M = 135^2$) in which we progressively increased pollen (Figure 4) or seed dispersal (Figure 5). Simulations were performed under the three different mating systems of S_u equal to 0.001, 0.1 and 0.9. We used the parameter P_{cd} to measure extent of separation of individuals expressed through pollen communication, and the equivalent parameter S_{cd} to express the extent of movement of seeds.

A threshold effect was again apparent in the pollen dispersal based perturbation simulations when S_u was set to 0.001, in which the gradient of the line varied either side of a P_{cd} value of 0.9, from -0.005 to -0.183 , Figure 4. Generally, lower values of H_o at each P_{cd} were obtained in this simulation than in the matrix size simulations for the same mating system. This is likely to be due in part at least to the effects of seed dispersal. The value of S_{cd} in the pollen dispersal based perturbations was consistently 2.3. However, S_{cd} varied in the matrix size based perturbation simulations

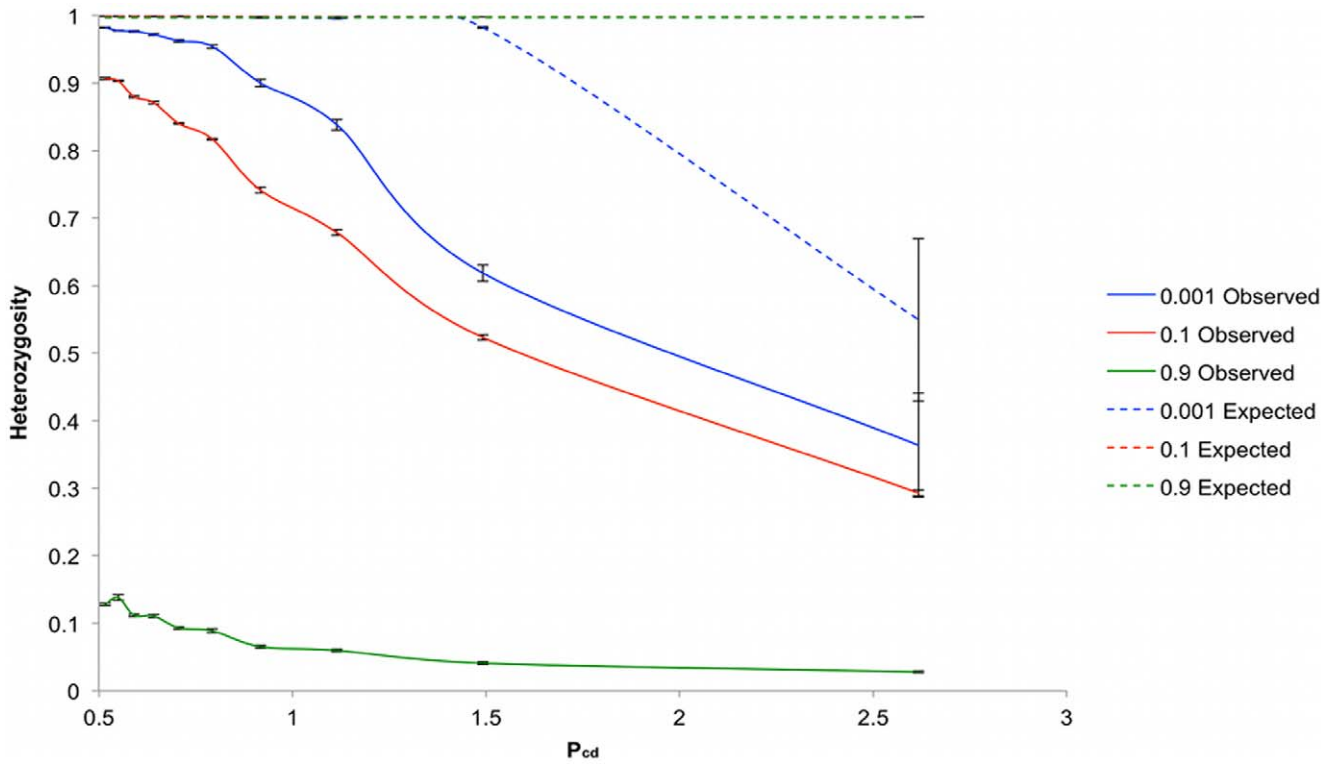


Figure 4. Heterozygosity as a function of P_{cd} . Heterozygosity was plotted at $S_u=0.001$ (blue), $S_u=0.1$ (red) and $S_u=0.9$ (green). H_o is plotted with the solid line and H_e using a dashed line. Simulations were repeated ten times for each value of S_u , with error bars representing the standard error.

doi:10.1371/journal.pone.0043254.g004

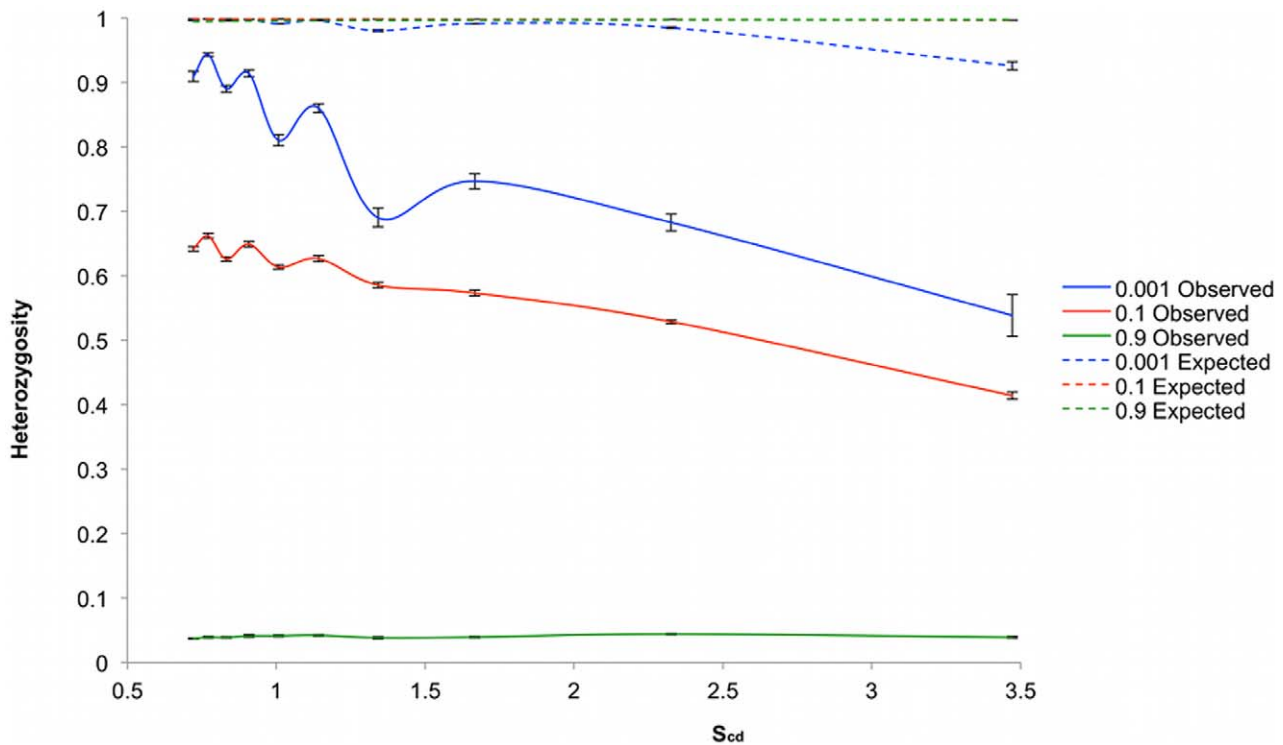


Figure 5. Heterozygosity as a function of S_{cd} . Heterozygosity was plotted at $S_u=0.001$ (blue), $S_u=0.1$ (red) and $S_u=0.9$ (green). H_o is plotted with the solid line and H_e using a dashed line. Simulations were repeated ten times for each value of S_u , with error bars representing the standard error.

doi:10.1371/journal.pone.0043254.g005

from 0.9–2.2. Consequently, at low values of P_{cd} there was a disparity in the effect of seed dispersal between the two systems, and the higher H_o of the matrix size perturbation based simulations is explicable due to the greater gene flow mediated by seed movement. The remaining two mating system simulations also produced results that echoed those of the matrix size based perturbation simulations (Figure 3), but with consistently lower values of H_o .

The effect of increasing seed dispersal had a more limited effect on increasing H_o than that observed with pollen dispersal, Figure 5. In these simulations, the P_{cd} value was consistently 1.5. In the case of simulations with S_u of 0.001, the results had a great deal of variance leading to a jagged line plotted of H_o against S_{cd} . In this case the jagged line correlates closely with the average population sizes generated in the simulations that varied considerably and included very low sizes (less than 100 individuals, data not shown). This is because low population densities (as defined by high values of P_{cd} and S_{cd}) are subject to a high risk of failure to reproduce under this mating system, which requires 99.9% outbreeding, because there may be an inadequate supply of neighbors sufficiently close for pollination. The other mating system simulations where S_u was 0.1 and 0.9 did not give rise to such an erratic set of results.

In all simulations the effect of increasing the seed dispersal on H_o was less than increasing the extent of pollen dispersal that demonstrates that although increasing seed dispersal has an influence on heterozygosity, it is weaker than the effect of pollen flow. Furthermore, we did not observe a clear threshold effect for seed dispersal.

Discussion

To our knowledge this is the only study where increasing spatial perturbations are made to examine the discrepancy between real observed values of population genetics variables, and those expected based on mean-field population genetics theory. The results we present here demonstrate that the deviation from classical population genetics introduced by the closer approximation to the real world through spatial extension is largely tractable, and different for different mating systems. We suggest that extent of spatial extension can be usefully viewed through a concept of effective population density which is described in the P_{cd} and S_{cd} variables, which can be measured directly for any plant species population in nature.

Our findings suggest that in outbreeding populations systems in which the P_{cd} and S_{cd} values are below 1, the deviation from classical expectations is generally low (less than 5%), and therefore classical population genetics approaches may make acceptable approximations of reality. The approximation breaks down rapidly once the threshold value of ~ 1 is exceeded for P_{cd} in particular. In the case of inbreeding systems, typical of sparsely populated systems, the effects of inbreeding are far greater than any introduced by spatial extension, consequently mean-field based adjustments for inbreeding may also be adequate in a spatially extended system. However, we found that populations with mixed mating strategies are more sensitive to increased spatial extension, and in these cases values generated from classical population genetics are likely to be poor approximations of the true values.

In order to study the effects of spatial extension we prevented clumping from occurring, which has been observed in previous studies [30]. It may be argued that in reality populations form a clumpy pattern of distribution. We found that increasing matrix size and nearest-neighbor distances had little effect on lowering H_o

when individuals were clumped together (not shown). Therefore clumping in this case may confound the effects of spatiality, even though clumped subpopulations maintain a spatial population structure. When we increased seed and pollen dispersal on clumped individuals, however, we saw similar increases in H_o as with spaced individuals, consistent with previous observations [30]. We suggest that clumped populations in the real world could usefully be considered within the effective population density framework presented here. Values of P_{cd} and S_{cd} could be considered within and between clumps. Often, these values of P_{cd} and S_{cd} within clumps would be low such that classical population genetics could be applied as an approximation. The P_{cd} and S_{cd} values between clumps may well be high enough to identify the expectation of genetic structure in the population. This in part answers Platt et al's [13] concern, that a lack of geographic barriers and continual changes in spatial genetic autocorrelation observed in *Arabidopsis thaliana* suggests that no single population structure could be identifiable, and therefore that application of classical population genetics could be problematic.

Our simulated results show that under certain conditions spatially extended populations closely approximate HWE, and this is verified by studies of plant populations in the real world [34–37]. These studies include plants with various pollen and dispersal systems, including both abiotic and biotic mechanisms. Plant populations of Sandalwood, for instance, with high dispersal and outcrossing rates conform to HWE [38] are consistent with our simulated results. In our simulations we observed that population sparsity leads to deviations from HWE. This deviation has also been observed in reality with sparse plant populations in species of juniper and poppy [39–40]. Conversely, it is expected that higher outcrossing frequencies should be associated with higher population densities, and so a tendency to conform to HWE, and this too has been observed in wind-pollinated conifers [41] and in *Mimulus ringens* [42]. Our simulations predict that some plant populations would be observed to conform to HWE in some cases and deviate from it in other cases due to the consequences of limited dispersal relative to stand density. This finding is concordant with that of Dering and Chybicki [43] who compared the genetic diversity of natural and artificial regenerations of *Quercus robur* (L.) and *Quercus petraea* (Matt.) Liebl. populations. In this case the natural regenerations were in HWE, but the artificially regenerated progeny plantations were not because of limitations in dispersal caused by the sowing regime. Similarly, limitations of dispersal caused deviations from HWE in 1 out of 14 sampled populations of *Striga hermonthica* [44]. Our simulations further showed that HWE is restored in sparse populations where dispersal rates are increased to rates in which values of P_{cd} are 1 or less. This effect is observed in *Desmodium nudiflorum*, which despite apparent population sparsity has long-range seed dispersal by animals [45].

This study demonstrates that our IBM system behaves as expected under neutral conditions, and has given some useful insight into how the relationship between stand density and dispersal as encompassed by the P_{cd} and S_{cd} values is a predictor of adherence to HWE that could be applied to the real world. The model therefore has utility in exploring hypotheses in which deviation from neutral expectations occurs due to other factors, such as selection. An emergent observation from studies of evolution at the systems and genomic level is that simple scenarios that affect single genes are rare, and that often many genes and regulatory networks are involved, as observed with humans [46–47]. Similarly, with the evolution of domesticated plants estimates of the number of genes underlying the domestication syndrome traits range from 27 to 70 in wheat [48–49]. Therefore there is a need to be able to consider evolutionary change in a way that

connects the interdependency of gene networks, genome architecture to spatially explicit populations. The IBM approach presented here is designed to achieve this end by having individuals that are capable of supporting gene networks in a virtual genome, such that a systems biology architecture can be connected to a spatially explicit population level of organization and selection. We believe that such approaches will enable the computational exploration of evolutionary genetics to move to a new level in which systems approaches and spatially explicit population genetics are integrated.

Methods

Model

Individuals are arranged onto a two-dimensional grid-like matrix. Cells may be occupied by a plant, or be empty. In each update the model processes each individual by iterating its age and allowing it to interact with other individuals. Each grid cell may contain only one adult and ten seeds. A ‘seed queue’ system was implemented to replace older seeds with newer seeds and discard the older seeds. Each simulation begins with 1000 diploid individuals each containing two genes per genome copy. When individuals germinate they can accumulate mutations, which are passed on to their progeny. Gametes are produced by Mendelian segregation of the individual’s genes at the floral stage of the life cycle. Individuals will then either pollinate themselves or other flowering individuals and then disperse their resultant seeds. Once individuals have dispersed their seeds they senesce and are removed from the model.

Burn-in. To ensure there had already been an accumulation of mutations as the simulations start, a burn-in algorithm was implemented. For each of the 1000 starting individuals 4000 updates with mutations occur to the genes at each copy of the genome, according to the input mutation rate of 10^{-4} . The resulting genotypes are then randomly assigned to each of the starting individuals.

Individuals. Each individual has the following life cycle stages: seed, vegetative, flowering and senescence. The number of updates for an individual to move to each life stage is drawn from a Poisson distribution. Once a seed has germinated it is considered to be an adult. Individuals may disperse pollen and reproduce within their flowering stage, with the number of ovules per individual set to 30. At the end of the flowering life stage all fertilized ovules are dispersed as seeds. At the end of the senescing life stage the individual dies and is removed from the model.

Matrix. Simulations were run with increasing matrix sizes of 35^2 cells to 135^2 cells in steps of 10 cells in each dimension. In each simulation, cells were made unoccupiable in the matrix thus preventing individuals from occupying these cells, termed ‘void cells’. The number of void cells per simulation was calculated according to the formula:

$$v = m^2 - 1000 \quad (3)$$

where v is the number of void cells and m^2 is the matrix size in cells. Simulations for each matrix size were repeated ten times. The matrix was constructed using a quadtree data structure [32]: this is a tree-like structure where objects are stored in two-dimensional space, with the whole matrix being the root node. As objects are added to the tree the tree nodes are recursively split into four more nodes, until the smallest node size has been reached (which was set to a 2×2 sub-matrix). When an individual is extracted for distance calculations the tree nodes are recursively searched for the specified (x,y) coordinates of the individual until

the smallest node containing the individual is obtained. Groups of individuals are extracted in a similar way using a radius. This method provided better performance than using an exhaustive search approach.

Dispersal. In this model pollen is dispersed by wind. A probability for self-fertilization per ovule, S_u is input by the user, if this probability value is overcome then the ovule may be pollinated by an outcrossing event. Seeds and pollen are dispersed using a cumulative form of the function, described in [33] for wind-pollinated plants such that the probability of a pollen grain travelling j cells is given by:

$$P(X = x_j) = \frac{ce^{ax_j}}{\sum_{i=1}^n ce^{ax_i}} \quad (4)$$

Where x_j is a distance in cells that a pollen grain is dispersed. The maximum number of cells pollen (P_{max}) can disperse is given by n . The variable c is the intercept and a is the gradient of the underlying function:

$$y = ce^{ax} \quad (5)$$

A value of 1.5 was used for c . For pollen x is considered in the range $1 \leq x \leq P_{max}$. The form of the equation used for seeds is similar, except that x is considered in the range $0 \leq x \leq S_{max}$. These variations ensure that pollen exiting the flower does outcross but that seeds can fall onto the same cell as their parent individual. P_{max} and S_{max} were user input. For pollen, a was calculated as:

$$a = \lambda / P_{max} \quad (6)$$

Where λ is a constant set to -2 for pollen dispersal. An equivalent equation is used for seed dispersal, with the denominator set to S_{max} and λ set to -4 . A maximum dispersal distance for both seed and pollen is input as a user defined parameter. The distance (D) a pollen or seed was dispersed was determined by generating a random number in the range of 0 to 1 that was used to sample under the probability distribution generated from equation (4).

Once distance of dispersal had been determined, the direction of dispersal was generated. Two different approaches were used for pollen and seeds respectively. The direction of dispersal was assumed to be due to a prevailing wind. The general direction of the wind was randomized for each update. The angle of the prevailing wind in degrees, θ , is drawn from a uniform distribution each update and is relative to the vertical axis. The specific direction of dispersal is generated from θ by resampling from a normal distribution with mean θ and a standard deviation of 180° to produce γ , the specific angle of dispersal. The destination cell is determined to which the seed will be dispersed using the values of γ and x .

For pollen dispersal, we began with the recipient of pollen and calculated the relative probabilities of pollination by all potential pollen donors. Potential pollen donors were those individuals that were within the range of the recipient defined by P_{max} . For each donor, the angle (ϕ) was calculated which is defined as the angle of the recipient to the donor relative to a vertical axis passing through the donor. A quantile value (q_i) was calculated for ϕ from the normal distribution of mean θ (angle of wind direction) and standard deviation of 15° (degrees), which gave a measure proportional to the probability of pollen emanating from the donor in the direction of the recipient. The probability of the

donor d_i being the successful pollen donor to the recipient was then calculated as:

$$P(d_i) = \frac{y_i q_i}{\sum_{j=1}^n y_j q_j} \quad (7)$$

Where y_j is calculated as in equation (4) for the distance x_j between potential donor and recipient, and n is the total number of potential pollen donors. The successful pollen donor was then selected through random sampling of all possible donors. The process was repeated independently for each ovule to be pollinated on the recipient plant.

Analysis

Heterozygosity calculations. The observed heterozygosity, H_o of remaining individuals after 4000 updates of each simulation was recorded and compared with the expected heterozygosity as calculated by an ideal population under Hardy-Weinberg equilibria, as in [23]. The Hardy-Weinberg formula for multiple alleles was used to calculate H_e .

Average nearest neighbor distances. To calculate the nearest neighbor distance an individual plant i , the distance (D) to all other individuals was calculated based on their Cartesian coordinates using:

$$D = \sqrt{(x_j - x_i)^2 + (y_j - y_i)^2} \quad (8)$$

Where x_i and y_i are the x and y coordinates of the individual i , and x_j and y_j are coordinates of the j th individual. The lowest value of D was recorded for each individual. All recorded D values were averaged at the end of the simulation to produce an average nearest neighbor distance. For most simulations, the pollen dispersal distance (P_d) or seed dispersal distance (S_d) was recorded for each dispersal event in number of cells. The effect of the

dispersal is context dependent on the density of individuals in the population, so we used an expression that captured this information defined in equations (1) and (2) in the main body of the text.

Selfing probabilities. In the experiment to explore the effect of mating strategy on heterozygosity, the model was run with input selfing probabilities (S_u) at 0, 0.001 and then from 0.1 to 1 in steps of 0.1. Each experiment at all selfing probabilities described was repeated ten times. A matrix size of 35^2 was used, with 225 void cells to limit the population to a maximum of 1000 individuals.

Pollen and seed dispersal. In the experiment to explore the effect of dispersal on heterozygosity, simulations were run with increasing maximum seed and pollen dispersal distances. To provide sufficient isolation of individuals simulations were run with a matrix size of 135^2 cells with 17225 void cells. Maximum seed and pollen dispersal were separately increased throughout simulations from 5 to 50 cells in steps of 5 cells. When maximum seed dispersal was increased, maximum pollen dispersal was set to 10 cells, and vice-versa for seed dispersal. With seed dispersal, seeds were given 1000 attempts to find a habitable cell, to avoid population crashes in the simulation. Distance values for pollen dispersal or seed dispersal were then calculated as described above.

Acknowledgments

We would like to thank Richie Smith at the University of Manchester useful discussions on data structures. We would like to thank Prof. Terry Brown at The University of Manchester and Dr Dorian Fuller at University College London for their help in designing this project. We are also thankful to Dr Jay Moore at the University of Warwick for his suggestions on design of some of the algorithms implemented.

Author Contributions

Conceived and designed the experiments: RA JK. Performed the experiments: JK. Analyzed the data: JK RA. Contributed reagents/materials/analysis tools: JK RA. Wrote the paper: RA JK.

References

- Allaby RG (2010) Integrating the processes in the evolutionary system of domestication. *Journal of Experimental Botany* 61: 935–944.
- Hardy GH (1908) Mendelian proportions in a mixed population. *Science* 28 (706): 49–50.
- Wright S (1943) Isolation by distance. *Genetics* 28: 114–138.
- Mayr E (1978) Review of Modes of Speciation by MJD White. *Syst. Zool.* 27: 478–482.
- Templeton A (1980) The theory of speciation *via* the founder principle. *Genetics* 94: 1011–1038.
- Garroway CJ, Bowman J, Wilson PJ (2011) Using a genetic network to parameterize a landscape resistance surface for fishers, *Martes pennanti*. *Mol Ecol* 20: 3978–3988.
- Rasic G, Keyghobadi N (2011) From broadscale patterns to fine-scale processes: habitat structure influences genetic differentiation in the pitcher plant midge across multiple spatial scales. *Molecular Ecology*: doi: 10.1111/j.1365-294X.2011.05280.x.
- Jones KN, Reithel JS (2001) Pollinator-mediated selection on a flower colour polymorphism in experimental populations of *Antirrhinum* (Scrophulariaceae). *Am. J. Bot.* 88: 447–454.
- Levin DA, Kerster HW (1972) Assortative pollination for stature in *Lythrum sativa*. *Evolution* 27: 144–152.
- Weis AE, Kossler TM (2004) Genetic variation in flowering time induces phenological assortative mating: quantitative genetic methods applied to *Brassica rapa*. *Am. J. Bot.* 91: 825–836.
- Bombliks K, Yant L, Laitinen RA, Kim ST, Hollister JD, et al. (2010) Local-scale patterns of genetic variability, outcrossing, and spatial structure in natural stands of *Arabidopsis thaliana*. *PLoS Genet* 6: e1000890.
- Baker HG (1955) Self-compatibility and establishment after “long-distance” dispersal. *Evolution* 9: 347–349.
- Platt A, Horton M, Huang YS, Li Y, Anastasio AE, et al. (2010) The scale of population structure in *Arabidopsis thaliana*. *PLoS Genet* 6: e1000843.
- Allaby RG, Fuller DQ, Brown TA (2008) The genetic expectations of a protracted model for the origins of domesticated crops. *Proceedings of the National Academy of Sciences USA* 105: 13982–13986.
- Allaby RG, Brown TA, Fuller DQ (2010) A simulation of the effect of inbreeding on crop domestication genetics with comments on the integration of archaeobotany and genetics: a reply to Home and Heun. *Vegetation History and Archaeobotany* 19: 151–158.
- Cuddington KM, Yodzis P (2000) Diffusion-limited predator-prey dynamics in euclidean environments: An allometric individual-based model. *Theoretical Population Biology* 58: 259–278.
- Wilson WG, McCauley E, De Roos AM (1995) Effect of dimensionality of Lotka-Volterra predator-prey dynamics: individual based simulation results. *Bull. Math. Bio.* 57: 507–526.
- Wilson WG, De Roos AM, McCauley E (1993) Spatial instabilities within the diffusive Lotka-Volterra system: Individual-based simulation results. *Theor. Popul. Biol.* 43: 91–127.
- McCauley E, Wilson WG, De Roos AM (1993) Dynamics of age-structured and spatially structured predator-prey interactions: Individual-based models and population-level formulations. *Am. Nat.* 142: 412–442.
- De Roos AM, McCauley E, Wilson WG (1991) Mobility versus density-limited predator-prey dynamics on different spatial scales. *Proc. R. Soc. Lond. B* 246: 117–122.
- Spear SF, Balkenhol N, Fortin MJ, McRae BH, Scribner K (2010) Use of resistance surfaces for landscape genetic studies: considerations for parameterization and analysis. *Mol Ecol* 19: 3576–3591.
- Kuparinen A, Schurr FM (2007) A flexible modelling framework linking the spatio-temporal dynamics of plant genotypes and populations: Application to gene flow from transgenic forests. *Ecological Modelling* 202: 476–486.
- Landguth EL, Cushman SA (2010) cdpop: A spatially explicit cost distance population genetics program. *Mol Ecol Resour* 10: 156–161.
- Gavrilets S, Li H, Vose MD (1998) Rapid parapatric speciation on holey adaptive landscapes. *Proc R. Soc. Lond* 265: 1483–1489.

25. Rice WR (1984) Disruptive selection on habitat preference and the evolution of reproductive isolation: a simulation study. *Evolution* 38 (6): 1251–1260.
26. Savill NJ, Hogeweg P (1998) Spatially induced speciation prevents extinction: the evolution of dispersal distance in oscillatory predator-prey models. *Proc R. Soc. Lond* 265: 25–32.
27. Durrett R, Buttel L, Harrison R (2000) Spatial models for hybrid zones. *Heredity* 84: 9–19.
28. Sadedin S, Hollander J, Panova M, Johannesson K, Gavrillets S (2009) Case studies and mathematical models of ecological speciation. 3: Ecotype formation in a Swedish snail. *Molecular Ecology* 18: 4006–4023.
29. Hartman PJ, Wetzel DP, Crowley PH, Westneat DF (2012) The impact of extra-pair mating behavior on hybridization and genetic introgression. *Theor Ecol* 5: 219–229.
30. Doligez A, Baril C, Joly HI (1998) Fine-scale spatial genetic structure with nonuniform distribution of individuals. *Genetics* 148: 905–919.
31. Hartl DL, Clark AG (1997) *Principles of Population Genetics*. Sinauer Associates, Sunderland, Massachusetts, 141–144.
32. Finkel RA, Bentley JL (1974) Quad Trees: A Data Structure for Retrieval on Composite Keys. *Acta Informatica* 4 (1): 1–9.
33. Beckie HJ, Hall LM (2008) Simple to complex: Modeling crop pollen-mediated gene flow. *Plant Science* 175: 615–628.
34. Barrat MD, Wallace MJ, Anthony JM (2012) Characterization and cross application of novel microsatellite markers for a rare sedge, *Lepidosperma Gibsonii* (Cyperaceae). *American Journal of Botany* 99(1): e14–e16.
35. King TL, Springmann JA, Young JA (2011) Tri- and tetra-nucleotide microsatellite DNA markers for assessing genetic diversity, population structure, and demographics in the Holmgren milk-vetch (*Astragalus holmgrenianus*). *Conservation Genet Resour* 4(1): 39–42.
36. Wöhrmann T, Guicking D, Khoshbakht K, Weising K (2011) Genetic variability in wild populations of *Prunus divaricata* Ledeb. in northern Iran evaluated by EST-SSR and genomic SSR marker analysis. *Genet Resour Crop Evol* 58: 115–1167.
37. Millar MA, Byrne M, Barbour E (2012) Characterisation of eleven polymorphic microsatellite DNA markers for Australian sandalwood (*Santalum spicatum*) (R.Br.) A.D.C. (Santalaceae). *Conservation Genet Resour* 4: 51–53.
38. Muir K, Byrne M, Barbour E, Cox MC, Fox JED (2007) High levels of outcrossing in a family trial of Western Australian Sandalwood (*Santalum spicatum*). *Silvae Genetica* 56: (5) 222–230.
39. Douaihy B, Vendramin GG, Boratynski A, Machon N, Bou Dagher-Kharrat M (2011) High genetic diversity with moderate differentiation in *Juniperus excelsa* from Lebanon and the eastern Mediterranean region. *Aob Plants*. plr003.
40. Allphin L, Windham MD, Harper KT (1998) Genetic diversity and gene flow in the endangered dwarf bear poppy, *Arctomecon Humilis*. *American Journal of Botany* 85(9): 1251–1261.
41. Restoux G, Silva DE, Sagnard F, Torre F, Klein E, et al. (2008) Life at the margin: the mating system of Mediterranean conifers. *Web Ecol* 8: 94–102.
42. Karron JD, Thumser NM, Tucker R, Hessenauer AJ (1995) The influence of population density on outcrossing rates in *Mimulus ringens*. *Heredity* 75: 175–180.
43. Dering M, Chybicki I (2012) Assessment of genetic diversity in two-species oak seed stands and their progeny populations. *Scandinavian Journal of Forest Research* 27(1): 2–9.
44. Olivier A, Glaszmann JC, Lanaud C, Leroux GD (1998) Population structure, genetic diversity and host specificity of the parasitic weed *Striga hermonthica* (Scrophulariaceae) in Sahel. *Plant Systematics and Evolution* 209: 33–45.
45. Smith WG, Schall BA (1979) Isozyme variation in *Desmodium nudiflorum*. *Biochemical Systematics and Ecology* 7: 121–123.
46. Pritchard JK (2010) How we are evolving. *Scientific American* 303: (4)41–47.
47. Sabeti PC, Schaffner SF, Fry B, Lohmueller J, Vardilly P, et al. (2006) Positive Natural Selection in the Human Lineage. *Science* 312: 1614–1620.
48. Peng J, Ronin Y, Fahima T, Röder MS, Li Y, et al. (2003) Domestication quantitative loci in *Triticum dicoccoides*, the progenitor of wheat. *Proc. Natl. Acad. Sci. USA* 100: 2489–2494.
49. Peleg Z, Fahima T, Korol AB, Abbo S, Saranga Y (2011) Genetic analysis of wheat domestication and evolution under domestication. *J. Exp. Bot.* 62: 5051–5061.