

The evolution of polyploidy and the extreme cost of pollen-swamping

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Abstract

Amazing summary of project here. . .

Keywords

polyploidy; evolution; angiosperm; mimulus; guttatus; pollen-swamping; mate-choice;

1 Introduction

Would some images of genome duplication be helpful?

Whole-genome duplication (polyploidisation) is an extreme mutation which occurs due to errors during cell division; such as non-disjunction where sister chromatids are not separated during anaphase. Failed disjunction leads to all the genetic material ending up in one of the two daughter cells; essentially doubling the genome within that cell. When this occurs during gametogenesis, it can lead to whole organisms with

more than two complete sets of chromosomes (polyploids). According to [Svartman et al. \(2005\)](#), the estimated rate of polyploidisation is one order of magnitude higher than that of standard mutation (10^{-5}). In humans, it coincides with around 5% of miscarriages ([Creasy et al. 1976](#)). Moreover, while there have been some live birth cases of triploid human infants, they usually die within hours. In irregular cases, they survive for several months, but none have survived more than six years ([Hashimoto et al. 2018](#)).

Newer data for miscarriage rate?

No polyploid mammal species has ever been found ([Svartman et al. 2005](#)), and the same is also usually true for birds (?). However, while in these taxa, genome-doubling is not well tolerated, for amphibians, fish, fungi, reptiles, and plants, the story is quite different. Amongst these, polyploidy is exceptionally well tolerated in plants. [Jiao et al. \(2011\)](#) found that genome-doubling is a ubiquitous feature in the lineages of almost all flowering plants (angiosperms), and [Wood et al. \(2009\)](#) also found that it has occurred even more recently (post genera formation) for 35% of all vascular plants. Plant surveys of natural populations over the last century further support this, reporting mean polyploid frequencies of 25% (?).

The Role of Genome-Doubling in Plant Diversification

Genome-doubling is thought to be fundamental in the diversification of plant species; having been found to coincide with 15% of angiosperm speciation events, and 31% in ferns ([Wood et al. 2009](#)). Furthermore, [Jiao et al. \(2011\)](#) showed that whole-genome duplication could even be linked with major biological innovations that have lead to diversification of the plant phyla. Their phylogenetic analysis brought to light two ancient groups of duplication events: the first grouping of duplications (around 319 millennia ago) influenced the diversification of regulatory genes that

were integral to seed development and the more recent grouping (around 192 millennia ago) affected genes that enabled flower development. The authors appropriately conclude that these events must have played a significant role in the rise of seed plants, as well as in the later diversification of angiosperms. Seeds and flowers were both crucial innovations that allowed plants to transition from an aquatic life-cycle to a terrestrial one by removing their dependence on water for reproduction — an event, broadly considered as one of the most significant in the history of our planet.

The full story, however, is not quite that clear-cut. According to [Mayrose et al. \(2011\)](#), genome-doubling significantly increases speciation rates of diploids, but new polyploid lines do not further speciate by that same mechanism. So, polyploid speciation rates are far smaller in comparison. Furthermore, they found polyploid extinction rates to be far higher than those of diploids because polyploids are clumped around the tips of phylogenies. This Likelihood-based analysis of vascular plants provided clear, and quantitative, support for the traditionally popular view that polyploidy most often leads to evolutionary dead ends.

First proposed by? Stebbins 71?

So, what is it then that allows this extreme mutation to succeed and become fixated? What scenarios and mechanisms combine to allow rare polyploid lines to succeed? Are the majority of the polyploids we see in natural populations doomed to extinction? Furthermore, can we predict which conditions will lead to their survival?

Despite having been studied for over a century, the factors that drive the success of polyploid establishment in the face of high extinction rates are still unclear. [Arrigo and Barker \(2012\)](#) conclude that polyploids tend to become extinct at the establishment phase due to reproductive

disadvantages such as triploid sterility, or limited mate-choice; the latter occurring via diploid pollen-swamping, or delayed flowering. On the other hand, polyploids are frequently linked with distinct traits such as “gigas effects”, which describes their increased size and plant organs. Additionally, benefits of polyploidy can include enhanced capabilities for buffering of deleterious mutation (due to increased heterozygosity), hybrid vigour (heterosis), and a reversal of selfing inhibition (Woodhouse et al. 2009; Ramsey and Ramsey 2014). These traits are thought to overcome the reproductive disadvantages of polyploidy and, in those rare cases, make this mutation key to the invasive and adaptive potential of plants.

The cost of being polyploid (limited mate-choice)

Diploid Pollen-Swamping

Delayed flowering

Delayed flowering (cost) is associated with gigas-effects of increased size (benefit). TRADE-OFF.

Mimulus study system

Polyploidisation events have been well documented in *Mimulus guttatus* (?).

? details 3-stage *M. guttatus* life-cycle and transition matrices.

Aims

Here we aimed to quantify the cost of pollen-swamping by testing how high the rate of whole-genome duplication would need to be in order to overcome the strength of negative selection. We asked:

- Will increasing the rate of whole-genome duplication lead to the fixation of polyploidy?

- And, if so, at what threshold point does this occur?

In addition, we aimed to discover how varying the starting population size affected these results:

- Does an established population react in the same way to increased duplication rates as a pioneering population entering an empty landscape?

2 Methods

An individual-based modelling approach was used to examine these aims. The model was created, from scratch, in *R* (R Core Team 2019) to simulate the life cycle of the yellow monkey flower (*Mimulus guttatus*) as laid out by ? and detailed in figure 1. The *M. guttatus* complex was a suitable choice due to whole-genome duplication being well documented throughout (?), and data being readily available.

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knitr::include_graphics("_images/m-guttatus-lifecycle-p")
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The model

The model was created as an open source R package {*sploidy*}, and is available along with the simulation scripts used for this work and the raw data, on Github (McKeon 2020).

Assumptions To test our hypotheses the model needed to make particular assumptions about the reproductive interactions between individuals, as well as how particular parameters (like whole genome duplication rate) were applied mechanistically. To see if the rate of duplication could overcome the effects of limited mate-choice, mate choice had to first be limited in some manner. This was achieved using a simple matching rule that defined whether attempts to fertilise were

C

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successful, based on parental ploidy level. Simulating, therefore, the cost of pollen-swamping.

The model assumptions were as follows:

- All individuals began as diploid.
- Polyploids arose at a set rate of whole-genome duplication during all sexually reproductive transitions.
- Diploids and polyploids could both out-cross, but never with each other. In addition, polyploids could not outcross with other polyploids of differing ploidy level, ie:- a tetraploid was not compatible with an octoploid.
- All individuals could self-fertilise.
- Offspring produced had the same ploidy level as their parents; two diploids produced a diploid, and so on.
- When whole-genome duplication occurred, the ploidy level of an individual was doubled; in this case two diploid parents would produce a tetraploid.
- All plants were hermaphrodite; mothers and fathers were chosen via random sampling from the same pool of individuals.
- Pollen range spanned the entire landscape.
- Seed dispersal spanned the entire landscape.

The *rbinom* function in base R was used throughout the model—where transitions reduced cohort size—to choose individuals that transitioned successfully. This allowed for a natural amount of stochasticity around the transition probabilities, similar to that which we would expect in nature. Where transition values increased cohort size, the function *sample_n* from the package *{Dplyr}* (Wickham et al. 2020) was used, instead, with replacement.

Table 1. Components of Low-elevation perennial *Mimulus guttatus** life-cycle transition. Seed survival (D) is a general value taken from ?. All other components taken from ? who recorded values specific to that *M. guttatus* population in the year 2013 (N = 77). G = germination rate, O = ovule number per flower, F = flower production, S = winter survival, R = rosette production, A = proportional recruitment success of ovules relative to rosettes.

D	G	O	F	S	R	A
0.534	0.652	494	5.34	0	0	0.00067

Simulation strategy

Rounds of simulations were run using the model function *sploidy* in order to home-in on the parameter ranges to include in the experiment design. These initial simulations were run for 1000 generations on a 100x100 landscape grid (to allow the default carrying capacity to maintain the adult population size at around one hundred thousand). The starting population contained sixty diploid individuals (twenty of each life stage), to simulate a pioneering species entering an empty space and the **ploidy_rate** parameter was varied between zero and 0.5. Seed dormancy was disabled by setting **seed_longevity** to zero. As was clonal reproduction, via the transition matrix provided to the **trans** parameter. Transition parameters—the transition matrix (**trans**), germination rate (**G**), and seed survival rate (**D**)—were taken from known demographic data (??).

Transition data Data for an *M. guttatus* population that had the most stable properties were used; minimal growth that could be kept in check with carrying capacity and low elasticity. ? described the formula for computing transition data as laid out in figure 2 and provided the values for a low-elevation perennial *M. guttatus* population as shown in table 1. These data produced the transition matrix (figure 2) provided to the model, along with the values of G (0.652) and D (0.534) shown in figure 1.

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knitr::include_graphics ("_images/m-guttatus-transformula
```


Seed_{t+1}

Seedling_{t+1}

Rosette_{t+1}

Seed_t

$\begin{pmatrix} D(1 - G) \\ DG \\ 0 \end{pmatrix}$

F

Figure 2. How the transition matrix (table 2) was calculated from the data provided in table 1, according to this formula taken from ?.

Table 2. Transition matrix used for the model, based on the formula shown in 2 and the data from figure 1.

	Seed (t)	Seedling (t)	Rosette (t)
Seed (t+1)	0.186	0.615	0.615
Seedling (t+1)	0.348	1.152	1.152
Rosette (t+1)	0.000	0.000	0.000

Varying the rate of whole-genome duplication In order to quantify how the rate of whole-genome duplication affected fixation of polyploid genotypes, a randomly sampled range of polyploidisation rates were chosen from an exponential distribution of 100000 random uniform numbers ranging between 0.001 and 0.010. Using an exponential distribution gave higher resolution around the lower parameter rates which we expected to be most interesting.

Diagnostic plotting of these data revealed that increased rates needed to be examined. Further simulations were then run using randomly sampled rates from 100000 random uniform numbers between 0.010 and 0.500 in

an attempt to determine how high the mutation rate needed to be in order for polyploids to break free from the inherent selection against them due to mate-choice limitation.

Germination rate benefit Further simulations which inferred a germination benefit to polyploids were run using the same randomly sampled rates of polyploidisation as above. A randomly sampled range of germination rate modifiers were chosen from 100000 random uniform numbers between 1 and 1.5, where the modifier was used to multiply the germination rate ($G = 0.652$). The max value of 1.5 would cause polyploid seeds to germinate at very close to complete success ($G = 0.978$), simulating the observation that polyploid seeds can have increased germination success due to germination over a wider temperature range (?).

3 Results

4 Discussion

Does limited mate-choice set the scene for the evolution of beneficial traits like selfing?

5 Conclusion

Does limited mate-choice set the scene for the evolution of beneficial traits like selfing?

A bit more manageable for a dissertation would be to try to flesh out one or two of these mechanisms and ask some questions about the conditions that it specifically is predicted to contribute substantially to diversification of polyploids – e.g., are there reasons to think that pollen swamping will lead to dead ends under some conditions, but diversification (or at least not dead ends) under others?

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