Through the Looking Glass: I. Why Cross-Fertilize?

Curtis M. Lively

27 November 2023

Contents

# Front Matter

## About

The following pages represent the first volume of a book. The main goal was to introduce the evolutionary problem of sexual reproduction, with a focus on competition between sexual and asexual females. But I also incorporated some ideas on genetic polymorphism and phenotypic plasticity with the goal of exploring “variation strategies” more generally. Finally, I tried to weave in some history of the field, along with some philosophy of science.

## Acknowledgements

I gratefully acknowledge helpful comments from Amrita Bhattacharya, Zoe Dinges, Kara Million, Deanna Soper, Mike Wade, Jukka Jokela, Dorota Paczesniak, Steve Howard, Lynda Delph, Clark Craddock, Robert Vrijenhoek, Jan McKenzie, Mike Winterbourn, Stuart West, Oren Harman, and especially Maurine Neiman and her lab group at the University of Iowa. Additional comments are welcome by [email](mailto:clively@indiana.edu).

Special thanks to Zoe Michelle Dinges (ZMD), who redrew the graphs and contributed original illustrations. Many thanks also to Adam Mazel of Indiana University Libraries for preparing the document for online publication.

This project was supported by the NSF OPUS program for the synthesis of biological research (DEB-1906465). I am also grateful to the Institute for Advanced Study in Berlin (Wissenschaftskolleg zu Berlin) for my stay as a “partner” during 2022-2023.

## Copyright and License

Copyright Curtis M. Lively 2023

Through the Looking Glass: I. Why Cross-Fertilize? by Curtis M. Lively is licensed under a Creative Commons Attribution 4.0 International License.

## Publisher Information

[Indiana University Libraries Publishing](https://libraries.indiana.edu/scholarly-communication)

## How to Cite

Lively, C.M. (2023). [*Through the Looking Glass: I. Why Cross-Fertilize?*](https://doi.org/10.5967/GBD3-KA07) Indiana University Libraries Publishing, Bloomington.

# 1. Why Sex?

|  |
| --- |
|  |

## 1.1 The Question

Most PhD programs require that students pass a preliminary examination. This was certainly true in my case. I was a PhD student at the University of Arizona studying rocky intertidal communities in the Northern Gulf of California. But the exams were not focused on our research. They were “depth-of-knowledge” exams. My question from Prof. Astrid Kodric-Brown instructed me to read the preface of G.C. Williams’ book, *Sex and Evolution*, which contains the following text (1975): “This book is written from a conviction that the prevalence of sexual reproduction in higher plants and animals is inconsistent with current evolutionary theory…. Many well informed readers may disagree with much of my reasoning, but I hope to at least convince them that here is a crisis at hand in evolutionary biology….”

The question was something like this: “Why does Williams think that sexual reproduction poses a crisis for evolutionary biology, and what is the solution?” A crisis? That was news to me. How could there be a crisis on evolutionary biology 40-plus years after the modern synthesis? My graduate course in theoretical population genetics did not mention any crises. I was not convinced. And a little freaked out.

The structure of our exams was very loose. I don’t remember having a deadline to produce a written answer, but I do remember that I spent several months on just this one question. During much of this time, I was doing field work in Sonora, Mexico, sometimes under very harsh conditions. But the more I studied the question, the more fascinated I became. I came to think that there was, indeed, a very real anomaly presented by sexual reproduction. Williams was right. Perhaps I was especially interested in this anomaly because I had read Thomas Kuhn’s *The Structure of Scientific Revolutions* as an undergraduate (1970). Kuhn made the case that dissecting anomalies can lead to interesting advances and that made sense to me. While I eventually produced an essay to address the question, the answer felt incomplete. I wanted to know more. There were many hypotheses, but there was no clear general explanation. Many years later, I am still working on my prelim question. This book is my revised answer.

## 1.2 The Problem

There are many problems with sexual reproduction, including the time spent finding mates and the risk of contracting sexually transmitted disease (review in Lehtonen *et al.* 2012). However, while important, these costs do not form the core of the paradox. Historically, the paradox of sex stems from two things: (1) the cost of meiosis, and (2) the cost of producing males.

### 1.2.1 The cost of meiosis: reduced relatedness

The “cost of meiosis” was proposed by George Williams (1975). His idea was simply that females are only half as related to their outcrossed offspring as they are to their self-fertilized or parthenogenetic offspring.[[1]](#footnote-33) (See [Box 1.1](#callout-1) for condensed definitions.) Williams’ idea also had theoretical support, as R.A. Fisher had already shown that an allele causing self-fertilization would rapidly spread to fixation, barring severe inbreeding depression (1941). So, why cross-fertilize? The persistence of cross-fertilization despite the cost of meiosis formed a paradox. This paradox created the crisis that Williams saw in evolutionary biology.

### 1.2.2 The cost of males

The other way to look at the problem was proposed by John Maynard Smith (1971, 1978). Here the issue is not relatedness. The problem stems rather from the difference between sexuals and asexuals in their per-capita birth rates ([Figure 1.1](#fig-1.1)). Imagine a population of sexual individuals at carrying capacity (). At the sexual females are, by definition, simply replacing themselves. This means that each sexual female is, on average, producing one son and one daughter. Both sons and daughters contribute genetically to the next generation, but only females give birth. Now, consider a mutation in a single female that causes her to reproduce asexually. She gives birth to two daughters instead of one daughter and one son. These two asexually produced daughters both give birth to two more daughters. Hence, after just two generations, the asexual female has four granddaughters, while the average sexual female has just one granddaughter ([Figure 1.1](#fig-1.1)). This asymmetry should lead to the rapid replacement of sexual females by asexual females ([Figure 1.2](#fig-1.2)). And by “rapid,” I mean within tens of generations, even for very large populations (Lively 1996). We thus seek a selective force that can give an advantage to sexual reproduction on a very short time scale.

|  |
| --- |
| Figure 1.1: The cost of males. Imagine a single clonal female in a sexual population at carrying capacity, . At , the sexual females are, on average, producing one daughter and one son. In contrast, the clonal female produces two daughters and four granddaughters. Hence, the clonal lineage should rapidly eliminate the sexual population ([Figure 1.2](#fig-1.2)). However, in nature, asexual reproduction is very rare in both plants (Whitton *et al.* 2008) and animals (Vrijenhoek 1998). Hence the paradox. Why is sexual reproduction so costly and yet so common? |

Several assumptions went into Maynard Smith’s model for the cost of males. In particular, he assumed that sexual females and asexual females make the same number of offspring, and that the survivorship of these offspring is also the same. Maynard Smith referred to this as the “all-else-equal assumption.” Unfortunately, some authors have taken the phrase “all-else-equal” to mean that everything else is exactly equal. But this is not the case. Maynard Smith did not assume, for example, that sexuals and asexuals have the same ploidy value.[[2]](#footnote-39) His model only assumes that sexual and asexual females have equal fecundities and survivorship probabilities (see [Box 1.2](#callout-2)). Under this assumption, a very rare clone would double in frequency in the next generation. Maynard Smith called this doubling-when-rare the two-fold cost of sex.

### 1.2.3 Contrasting the costs

The two alternative costs of sex raise an immediate question. Does the cost of sex result from reduced relatedness between mother and offspring, or from the cost of producing males? Or is the cost some combination of both? These questions are not easy to answer; but there is an algebraic solution, which suggests that the (1) two costs are mutually exclusive and (2) that they apply to different kinds of uniparental progeny (Lively & Lloyd 1990). Roughly speaking, I think we can adopt the following rules for the purpose of this book. When considering the spread of a rare allele that induces self-fertilization in hermaphrodites, the appropriate cost is Williams’ cost of meiosis. Here we have a single population in which the selfing allele is under positive selection because it has a transmission advantage. On the other hand, when we consider the spread of a clone into an obligately sexual population, we are dealing with competition between two different reproductively isolated groups. One group (the sexuals) produces males, which do not make offspring. The other group (asexuals) produces only females. Here the cost of sex stems from producing males. But the two costs do not combine. The cost of sex is not four-fold.

|  |
| --- |
| Figure 1.2: Clonal invasion dynamics. Results from a simulation study in which a single clonal individual was introduced into a sexual population (Lively 2009). **A** (top). Results for a 1:1 sex ratio in the sexual population. Here the frequency of daughters produced by sexual females was 1/2. The sexual population was initiated at carrying capacity: = 10,000. A single parthenogenetic female was introduced by the simulation at generation 1,000. Note that the asexual lineage replaces the sexual population in about 25 generations, and that it reaches a higher carrying capacity = 20,000. **B** (bottom). Results for a female-biased sexual population. Here the frequency of daughters produced by sexual females was 0.8. The sexual population was initiated at carrying capacity: = 17,500. As above, a single parthenogenetic female was introduced into the population at generation 1,000. Note that the asexual lineage replaces the sexual population, but it takes longer. The simulation assumes annual reproduction and non-overlapping generations. The R code for the simulation, including interactive graphical output, can be found [here](https://raw.githubusercontent.com/IULibScholComm/through-the-looking-glass/main/sim%20for%20fig%201.2(ZMD).R). The interactive graph can also be run [here](https://connect.posit.iu.edu/clonal-invasion-dynamics/) for users without R. |

### 1.2.4 The cost of recombination

There is another paradox of sexual reproduction known as the “cost of recombination.” Here the competition is not between sexual and asexual females, or between outcrossing and selfing alleles, but rather between alleles that modify the rate of recombination. So instead of asking “Why cross-fertilize?” we can assume cross-fertilization and ask, “Why is there excess crossing-over during meiosis?” Here is the paradox. If combinations of alleles at different loci are favored by natural selection (because together they create high-fitness offspring), then recombination would break these favorable allelic combinations apart. So, it makes no obvious sense to recombine more than needed for normal meiosis. Indeed, Lewontin (1971) formally showed that “the mean fitness of the population at equilibrium is a maximum in the absence of recombination.”[[3]](#footnote-48) Hence, there are two interrelated anomalies: cross-fertilization *per se* and meiotic recombination. Ideally, any theory that explains the persistence of biparental sex could also solve the paradox of recombination. But this need not be the case. They could have different solutions.

|  |
| --- |
| Box 1.1 |
| **Short definitions of terms as used in this book. These definitions do not include all possible nuances.**  **Carrying capacity:** The population density at which females have just enough food to replace themselves. Sexual females must make two offspring to replace themselves (assuming a 1:1 sex ratio), while asexual females must only produce one offspring. Hence, asexuals should have higher carrying capacities, as shown in [Figure 1.2](#fig-1.2).  **Cost of males:** The reduction in the per-capita growth rate of sexual populations due to the production of males. The cost of males is the appropriate cost for considering sexual subpopulations in competition with obligately asexual subpopulations.  **Cost of meiosis:** The reduction in relatedness between mother and offspring due to outcrossing. The cost of meiosis is the appropriate cost for considering the spread of alleles that induce self-fertilization.  **Clone:** A lineage of parthenogenetic females descended from the same asexual female. Members of the same clone may have small genetic differences, which accumulate by mutation over time.  **Cross-fertilization:** The exchange of gametes between different individuals, which may or may not be related.  **Outcrossing:** A form of cross-fertilization, which specifies crossing between unrelated individuals.  **Parthenogenesis:** Any form of asexual reproduction through ova.  **Recombination:** Genetic exchange between homologous chromosomes during meiosis, especially when the exchange leads to gametes with allele combinations not represented on the parental chromosomes.  **Self-fertilization:** The fusion of gametes from the same individual.  **Sex/rec:** Shorthand for sexual reproduction and recombination.  **Sexual reproduction:** I use the term here to mean cross-fertilization between unrelated individuals. However, the term is more general and can be used to mean the incorporation of novel genetic material by any mechanism. |

### 1.2.5 Darwin’s view

Even before the cost of males and meiosis were so dramatically revealed by Williams and Maynard Smith, biologists were reckoning with the anomaly of sex (Dagg 2016; reviews in Meirmans 2009). One of the earliest of these biologists was Charles Darwin. After he published the *Origin of Species*, Darwin was doing hand-pollination experiments at Down House on three species of a curious annual plant in the genus *Primula*. The plant is curious in that it has two morphs. One morph has a style that extends beyond the anthers (the long-style morph), and the other morph has anthers that extend beyond the style (the short-style morph). Botanists refer to this condition as distyly ([Figure 1.3](#fig-1.3)). Darwin found that crosses between the different morphs of the same species resulted in a very successful production of seeds, but crosses between unrelated individuals of the same morph were dramatically less successful (1862). In discussing these results, Darwin speculated that the two morphs may have evolved to insure cross-fertilization: “Whether or not the dimorphic condition of the *Primula* has any bearing on other points in natural history, it is valuable as showing how nature strives, if I may so express myself, to favour the sexual union of distinct individuals of the same species.”

|  |
| --- |
| Figure 1.3: Two flower morphs (distyly) in *Primula*. Darwin found that the short-styled morph (left) is incompatible with other short-style morphs and that the long styled morph (right) is incompatible with other long-style morphs. But the two different morphs can cross-fertilize. The arrows show movement of pollen from anthers to stigmas. The “X” indicates incompatibility. Redrawn from Darwin (1862) by ZMD. |

Darwin then asks a killer question. Why should the union of elements from distinct individuals be favored? Why, in fact, is there sex? “Nor do we know why nature should thus strive after the intercrossing of distinct individuals. We do not even in the least know the final cause of sexuality; why new beings should be produced by the union of the two sexual elements, instead of by a process of parthenogenesis. The whole subject is as yet hidden in darkness.” Darwin’s question shows that the cross-fertilization is curious, even without considering the costs of sex. It also shows how Darwin was drawn to anomalies on theory.[[4]](#footnote-57)

It is interesting to note that, in Darwin’s quote above, he switches from discussing mechanisms to prevent self-fertilization, such as distyly, to discussing parthenogenesis. Self-fertilization is a sexual process (involving the formation and fusion of gametes from the same parent), while parthenogenesis is an asexual process that does not generally involve meiosis and syngamy (review in Bell 1982). But parthenogenesis and self-fertilization are conceptually related, as they are both uniparental forms of reproduction. Hence, it makes sense that Darwin would switch back and forth between these two different forms of uniparental reproduction. Why cross-fertilize if either selfing or parthenogenesis is an option?

There may be another reason why Darwin pivots to parthenogenesis. Just prior to the publication of Darwin’s (1862) paper on *Primula*, Carl Theodor Ernst von Siebold (1856) published his observations on the successful development of adults from unfertilized eggs, which he called “parthenogenesis” (virgin birth). These were revolutionary observations, which caught Darwin’s attention. In a letter to his mentor, J.S. Henslow, Darwin mentioned von Siebold’s discovery as follows: “There is no greater mystery in the whole world, as it seems to me, than the existence of sexes, – more especially since the discovery of Parthenogenesis” (Darwin n.d.).

However, the discovery of parthenogenesis was met with some hostility.[[5]](#footnote-58) Consider, for example, the following statement by Rudolf Wagner in his 1857 review of von Siebold’s book on parthenogenesis (as translated from the original German by Churchill 1979): “I must unfortunately say that one of the most unpleasant of facts, [*Parthenogenesis*] has been introduced into physiology, which for the hope of so-called general laws of animal life-phenomena *is most distasteful*. It is impossible, considering the glorification of our highly vaunted progress in the theoretical understanding of the life processes, for it to be welcomed or particularly encouraged; and sincerely speaking, I can be as little pleased about it as a physicist would be if suddenly one or more exceptions to the law of gravitation were discovered” (Emphasis added).

Clearly, Wagner was not pleased with the discovery of asexual reproduction, calling it unpleasant, unwelcome, and distasteful. By contrast, Darwin did not find the idea to be distasteful in any way. He wondered instead why it was not more common. For example, Darwin (1868) wrote, “Parthenogenesis is no longer wonderful; in fact, the wonder is that it should not oftener occur.”[[6]](#footnote-59)

Over 100 years later, W. D. Hamilton (1975) was also pondering the evolution of outcrossing, and he wrote something conceptually similar: “[c]omplete inbreeding abandons the obviously important advantages of sexual reproduction, whatever these are.”

Whatever these are! The advantages of outcrossing were obviously important because cross-fertilization is so dominant. But the source of these advantages was not clear. At about the same time, Maynard Smith (1976) mused, “One gets the feeling that some essential feature of the situation has been overlooked.”

I now think that John Maynard Smith was correct. An essential feature had indeed been overlooked: parasites.

## 1.3 Summary

1. Obligate sexual reproduction is subject to invasion and replacement by all-female asexual lineages that do not pay the cost of males.
2. Obligate outcrossing in simultaneous hermaphrodites is subject to invasion and replacement by self-fertilization unless inbreeding depression is severe.
3. The exchange of DNA between different parental chromosomes (recombination) is similarly paradoxical.
4. Why then are recombination and cross-fertilization so common?

|  |
| --- |
| Box 1.2 |
| Maynard Smith’s (1978) model showing the cost of producing males.[[7]](#footnote-65) Let be the number of asexual females at time one, while gives the total number of sexual individuals (males plus females) at time one. Let give the number of offspring produced by asexual females, and gives the survival probability of asexual offspring to maturity. The number of surviving asexual offspring is then . Similarly, let be the number offspring produced by sexual females, and let give the survival probability of sexually produced offspring. Maynard Smith assumed that all individuals reproduce once and then die. Let be the frequency of females in the sexual population. The number of asexuals and sexuals at time two can then be calculated as in the table below. (Note, we do not assume that the population is at carrying capacity).  Table 1.1: Maynard Smith’s Model   |  | **Time One** | **Time Two** | | --- | --- | --- | | **N. of asexuals** |  |  | | **N. of sexuals** |  |  | | **Freq. of asexuals** |  |  |   The fold increase in frequency of asexuals, , is the ratio of the frequency of asexuals at time two divided by the frequency of asexuals at time one giving:  Under the all-else-equal assumption, and , giving:  Assuming that there is a single asexual female at time one, we get  If is very large, the solution reduces to . Hence, the fold increase in the frequency of asexuals, , is inversely related to the frequency of females in the sexual subpopulation. Assuming a 1:1 sex ratio, . Hence, for an equal sex ratio, the increase in asexuals is two-fold. This result gives the two-fold cost of males. Assuming “all-else-equal” a clone will double when rare when introduced into a large sexual population. |

# 2. The Ecological Hypotheses

|  |
| --- |
|  |

The sex/recombination (“sex/rec”) anomaly has attracted some of the best theoretical biologists over the last 50 years leading to at least two dozen hypotheses to explain selection for recombination and/or the persistence of obligate sexual reproduction in natural populations (Kondrashov 1993). In what follows, I first focus on the ecological hypotheses. The ideas underlying these hypotheses provide a handle for understanding some of the foundational concepts in evolutionary ecology.

## 2.1 The Lottery Model

As part of his book on the evolution of sex, Williams (1975) suggests that sex could be favored in fluctuating abiotic environments. The idea is intuitive: cross-fertilization generates variation among offspring. Hence, in fluctuating environments, sex could increase the probability that some offspring might survive. Williams likens the idea to a unique kind of lottery. For example, he writes, “Suppose you were offered this choice in a lottery: either you could have several different tickets, or you could have the same number of copies of the same ticket” (p. 15). If you choose *N* copies of the same ticket (asex) and your ticket wins, you get *N* times the reward. If you choose *N* different tickets (sex), you increase the probability of winning something, but the reward is smaller. Williams refers to the idea as the aphid-rotifer model, but the idea has since come to be known as the Lottery Model (following Bell 1982), which is a more descriptive phrase.[[8]](#footnote-72)

In my evolution course, I ask a form of Williams’ question but with a slight twist:

If you had a garden, upon which your descendants will depend for many generations to come, would you

1. plant a genetically variable crop, or
2. a monoculture with a two-fold higher yield?

Keep in mind that your descendants will follow your choice.

Often the students rightfully want some clarification. They ask, for example, “Can we use pesticides?” But every time, most students choose the variable crop. I remind them that selecting the variable crop will reduce their yield by one half. They don’t budge. I re-ask the question, doubling the relative yield for the monoculture from two-fold to four-fold. Then to 10-fold. Occasionally, one of the more risk-prone students will select the 10-fold higher yield, but most do not budge. They want genetic variation. I ask them why. Invariably, they say that the environment is going to change. They want to hedge their bets against an uncertain future.

Indeed, Williams’ Lottery Model is about bet hedging. The gist of bet hedging in evolutionary theory is that selection can act to reduce the variance in reproductive success over time, even if it also reduces the arithmetic mean across years (review in Philippi & Seger 1989). Suppose, for example, that we have the following data for both a monoculture and a genetically variable polyculture (in arbitrary units). Let’s assume that the variation in yield is driven by annual variation in abiotic factors such as temperature or precipitation. The effect of planting a polyculture (bet-hedging) can be estimated from the geometric mean, which incorporates the variation in yield over time.

Table 2.1: Bet Hedging

| Year | Monoculture | Polyculture |
| --- | --- | --- |
| 1 | 350 | 250 |
| 2 | 400 | 250 |
| 3 | 300 | 200 |
| 4 | 100 | 200 |
| 5 | 50 | 200 |
|  |  |  |
| **Arithmetic mean** | **240** | 220 |
| **Variance**[[9]](#footnote-73) | 19400 | 600 |
| **Geometric mean (GM)**[[10]](#footnote-74) | 184 | **219** |
| **Approximate GM** | 200 | **219** |

In this example, we find that the monoculture has an arithmetic mean of 240, while the polyculture has an arithmetic mean of only 220. So, I might be inclined to plant the monoculture. However, the among-year variance is very high for the monoculture (relative to the polyculture), driven in large part by the low yields in years 4 and 5. By contrast, the geometric mean for the monoculture is 184, while the geometric mean for the polyculture is 219. Based on this, I might be inclined to plant the polyculture, as it reduces the cost of very low yield in bad years. I think the students see this intuitively. Over the long term, it is better to be risk-averse and plant the genetically variable polyculture. What if, for example, the monoculture produced no food in the last year? The geometric mean would be zero. That would be catastrophic.

The effect of variance on the geometric mean (GM) can be seen by an approximation (Stearns 2000):

where is the mean, and var() is the variance in . Note that the approximation is equal to the arithmetic mean when the variance in is zero. Note too that the geometric mean increases as the variance in decreases. So, if selection operates to reduce the among-year variance in fitness, the outcome of selection will be reflected by an increase in the geometric mean. In general, evolutionary biologists use the geometric mean (rather than the arithmetic mean) to measure fitness over time.[[11]](#footnote-76)

Can sex be favored in variable environments as a bet-hedging strategy? It seems like a very sensible idea provided that the production of genetically variable progeny reduces the among-year variance in offspring survival. But remember, under a two-fold cost of sex, asexuals can replace large populations of sexuals in tens of generations (see [Chapter 1](#sec-why-sex)). So, if the Lottery Model is correct, significant environmental change must occur very rapidly. The many thousands of years between ice ages, for example, would be too long.

## 2.2 The Tangled Bank/Frozen Niche-Variation Model

Roughly speaking, the Lottery Model concerns the value of producing diverse offspring in a temporally variable abiotic environment. A different kind of model instead concerns the role of competition for different resource types that vary in space. Let us first consider the Frozen Niche-Variation Hypothesis of Robert Vrijenhoek. The key idea is that the clonal derivatives of sexual ancestors “freeze” some of the genetic variation in the sexual population. This frozen genotype then determines the resource niche of the clone. It seems reasonable to assume that the niche width of a single clone would be relatively narrow compared to the niche width of the genetically diverse sexual population. So, under this idea, a clone could invade a sexual population and perhaps displace it from one of its many niches. But a single clone could not completely replace the sexual population (Vrijenhoek 1979). This kind of process could explain those situations in which sexual and asexual females coexist, which was a major advance.[[12]](#footnote-78)

A conceptually similar model was independently developed by Graham Bell: the Tangled Bank Hypothesis (1982). Bell nabbed the name from the last paragraph of the *Origin of Species*, in which Darwin imagines life as an “entangled bank” of species interacting in a complex network. The core of the idea can be traced back to Howard Levene’s (1953) pioneering model, which showed that polymorphism could be maintained in a spatially heterogeneous environment provided that different genotypes specialize on different resources. Levene’s model was a major advance, as it showed that genetic diversity could be maintained without heterozygote advantage ([Box 2.1](#callout-2.1)). This was also one of the first models to fuse population genetics with ecology. But how does multiple niche polymorphism apply to sex? The idea is that if selection results in polymorphism, then a genetically diverse sexual population might be resistant to replacement by a clonal lineage that specializes on only one of the available resource types (as also in the Frozen Niche-Variation Model).

Here is how I pose the Tangled Bank idea to my undergraduate students. I start by giving them a choice between two hypothetical resources, which occur in different parts of the room. One is pizza, the other is broccoli. They all choose pizza. The problem, of course, is that the per-person value of the pizza resource declines as the pizza-eating population grows. At some point, there will be an advantage to specializing on broccoli. This could lead to a polymorphic population composed of obligate pizza eaters and obligate broccoli eaters, where (at equilibrium) the value of both resources is the same. Hence, selection for or against a particular strategy depends on the frequency of that strategy in the population. Perhaps this kind of “frequency-dependent selection” could favor sexual reproduction as a way to diversify offspring in environments where different resource types are patchily distributed in space. This reasoning forms the essence of the Tangled Bank Hypothesis.

One especially interesting aspect of the Tangled Bank Model is that the strength of frequency-dependent selection depends on population density. For example, there would be no selection to utilize the resource of lower value (broccoli) if there were no competition for pizza. This kind of selection, where the advantage to being rare depends on population density, is sometimes referred to as “soft” selection (Wallace 1975).[[13]](#footnote-79) In other words, soft selection is selection that is both frequency-dependent and density-dependent. This idea contrasts nicely with the Lottery Model, where selection is both frequency- and density-independent, which is called “hard” selection. For our purposes, we can use Wallace’s terminology to conceptually separate the Tangled Bank Hypothesis from the Lottery Model [Figure 2.1](#fig-2-2).

|  |
| --- |
| Figure 2.1: Partitioning the ecological hypotheses for the maintenance of obligate sexual reproduction. The figure is redrawn from Wallace (1975). The inserted blue text shows how the ecological hypotheses fit into Wallace’s matrix for density-dependent selection vs frequency-dependent selection. The Lottery Model relies on hard selection in temporally variable environments. The Tangled Bank relies on soft selection in spatially variable environments. The Red Queen relies on frequency-dependent selection generated by coevolving antagonistic species. |

We can think of the contrast like this. Under the Lottery Model, changes in the environment will select against certain genotypes independent of whether they are common or rare. Selection seems unconditional (hard). Under the Tangled Bank, selection is always conditional (soft); there is an advantage to having a rare genotype, but this advantage only accrues under strong competition (high density). Soft selection may not be exactly the best possible phrase, but it contrasts nicely with hard selection.[[14]](#footnote-84)

Two caveats are worth mentioning with respect to soft selection and the Tangled Bank Hypothesis. One is that polymorphism is only stable under a narrow range of patch-types frequencies. In addition, strong tradeoffs are required for the cost and benefits for morphs occupying different patches (Lively 1986a; Maynard Smith & Hoekstra 1980) (see also [Box 3.1](https://iulibscholcomm.github.io/through-the-looking-glass/eco-hyp-cont.html#callout-4)). The second caveat is that repeated mutation to asexual reproduction could lead to the accumulation of clonal diversity over time. Once all the niches are occupied by different specialized clones, there would be no advantage to sex. A diverse clonal population could then replace the sexual population (Bell 1982; Vrijenhoek & Parker 2009). This second caveat applies, in general, to any model of sex that relies on frequency-dependent selection. But the ideas could work if mutations to asex are rare. And, as I mentioned, sexuals and asexuals are known to coexist in some populations, which is consistent with the Tangled Bank and Frozen Niche-Variation Hypotheses (Vrijenhoek & Parker 2009). Coexistence, however, is also compatible with the Red Queen hypothesis, which we will now consider.

## 2.3 The Red Queen Hypothesis

The Red Queen Hypothesis is like the Lottery Model in that it focusses on environmental change over time. However, under the Red Queen idea, the change is mediated by changes in coevolving biological antagonists such as parasites, rather than changes in the abiotic environments.[[15]](#footnote-87) The distinction is important, as we will see.

It may be helpful to revisit [Figure 1.2](#fig-1.2), which shows the replacement of a sexual population by a clonal lineage within 25 generations. In this example, the clone is a single genotype, while the sexual population is composed of multiple recombining genotypes, only one of which is shared with the clone. Clearly, as the clone spreads, its genotype would become the most common in the host population. Now suppose that the host population is coevolving with a parasite population, which is composed of multiple strains. Assuming random contact between hosts and parasites, the parasite strain that could infect the most common host genotype would have a selective advantage over parasite strains that could only infect rare host genotypes. Let’s call this more successful parasite strain “strain **A**.” What would happen? It should be easy to see that strain **A** would increase in frequency. The parasite population would evolve.

Now, what if the parasite dramatically reduces the reproductive success of infected hosts? We might expect that, as the parasite evolves to infect the most-common host genotype, the reproductive advantage of the host clone is eroded. Moreover, if the parasite is *common and sufficiently virulent*, evolution by the parasite could prevent the clone from eliminating the sexual population. Under this scenario, there are at least two possible outcomes. One is that the sexuals and asexuals come to exist in stable frequencies, where the lost fecundity of the clone due to infection is equal to the cost of males, meaning that the mean fitnesses of sexuals and asexuals are equal. On the other hand, if the parasite is highly virulent, the frequencies of sexuals and asexuals can oscillate over time ([Figure 2.2](#fig-2-3) A). Under this second scenario, the new clone initially increases, but it is driven down sharply by infection ([Figure 2.2](#fig-2-3) B). Then, once the clone becomes very rare, it should become less infected than observed in the sexual population ([Figure 2.2](#fig-2-3) B). During this period, there is parasite-mediated selection against sex. Hence, the clone increases in frequency ([Figure 2.2](#fig-2-3) A), only to be driven down again by parasites after it becomes common ([Figure 2.2](#fig-2-3) B). Another cycle begins. **The key point is that parasites do not select against clonal reproduction per se; they only select against common genotypes.** But selection against common host genotypes might be sufficient to prevent fixation of a clone in the short term.

|  |
| --- |
| Figure 2.2: Simulation models showing that coevolving parasites can prevent the fixation of asexuals in the short term. **A.** The number of sexual and asexual individuals over time. **B.** The frequency of infection in sexual and asexual individuals over time. The simulation introduces a single clonal genotype into a sexual population at carrying capacity. After the clone becomes common (**A**) the parasites evolve to “target” it for infection (**B**). Note that after the parasites have driven the clone’s frequency down, the asexuals are less infected than the sexuals. Simulation model based on Lively (2009), which treats parasite virulence as a positive function of host density. |

This scenario of fluctuating selection for and against sex is just a special case of the more general idea that parasites will select against common genotypes within a diverse, sexual host population. As a rare host genotype becomes common, the parasites genotype that can infect it will be favored by natural selection. If the parasite is virulent (meaning that infection reduces host fitness), the targeted host genotype will decline in frequency, and a new host genotype will begin to increase in frequency. Under this logic, host-parasite coevolution will lead to the oscillation of genotypes in both the host and the parasite populations ([Figure 2.3](#fig-2-4)). These oscillations are now called Red Queen dynamics. Red Queen dynamics can lead to the maintenance of genetic polymorphism in sexual populations, and possibly protect sexual reproduction from replacement by asexual lineages. In addition, Red Queen dynamics could also favor recombination within a sexual population (Peters & Lively 1999, 2007; Salathe *et al.* 2008; Schmid-Hempel & Jokela 2002). These related ideas are now called the Red Queen Hypothesis (following Bell 1982).

|  |
| --- |
| Figure 2.3: Red Queen dynamics. The frequency of a single host genotype is shown along with the frequency of the only parasite genotype that can infect it. Note that the parasite tracks the host with a time lag. Results were extracted from a simulation of a sexual host population with nine possible genotypes coevolving with an asexual parasite with nine matching genotypes (Lively 2009). The dashed line shows the average genotype frequency for hosts and parasites. |

### 2.3.1 An intersection of science and literature

The name for the Red Queen Hypothesis comes from *Through the Looking Glass* (Carroll 1872). Here are the relevant bits of the story. After Alice goes through the looking glass (a mirror), she decides to follow a straight path to the top of a hill. But, in following the path, she ends up at her starting point. Talking to herself, she remarks, “But how curiously it twists! It’s more like a corkscrew than a path.” Repeated attempts were unsuccessful. In frustration, Alice addresses a tiger lily amongst a patch of flowers, “I wish you could talk!” The lily informs Alice that all the flowers can talk. The stunned Alice then begins a conversation with the flowers before finally asking, “Are there any more people in the garden besides me?” The rose answers “Yes, there is someone like you.” Alice sets out to follow this person (the Red Queen), but she quickly loses sight of her, and ends up back at her original starting point. Flustered, Alice decides to follow the advice of the rose: “*I* should advise you to walk the other way.” Alice then quickly finds the Red Queen.

Now it gets especially interesting. Alice mentions to the Red Queen that she would like to “find my way to the top of that hill.” The Red Queen replies, “*I* could show you hills, in comparison with which you’d call that a valley.” Alice protests: “a hill *ca’n’t* be a valley…. That would be nonsense.” This exchange between Alice and the Red Queen now seems prophetic, because, under frequency-dependent selection, locations on the adaptive landscape can rapidly change from fitness peaks to fitness valleys. Perhaps the Red Queen’s statement is correct: hills can become valleys, and valleys can become hills. More specifically, genotypes that were favored by natural selection when rare can become selected against after they become common, leading to a highly dynamic adaptive landscape.

In any case, Alice had clearly entered a crazy world. Straight paths become like corkscrews, progress is made by going the other way, and hills become valleys. Then, suddenly, Alice and the Red Queen began to run: “Alice never could quite make out, in thinking it over afterwards, how it was that they began: all she remembers is, that they were running hand in hand, and the Queen went so fast that it was all she could do to keep up with her.” During this furious run, Alice notices that they never pass anything. The trees remain in the same place as if they were moving along with them. Alice eventually asks: “Are we nearly there?” The Red Queen replies: “Why, we passed it ten minutes ago! Faster!”

When they finally stop, Alice is surprised to be where they started: “I do believe we’ve been under this tree the whole time! Everything’s just as it was!” The Red Queen replies that of course, and then asks: “What would you have it?” Alice replies that she would have expected to get somewhere else after running for a long time. The Red Queen then replies with this very famous quote: “Now, *here*, you see, it takes all the running *you* can do to keep in the same place.” It is a perfect metaphor for host-parasite coevolution.[[16]](#footnote-96) Host and parasite genotypes might oscillate as if they were running to stay in the same place.

It seems unlikely that Lewis Carroll had coevolution in mind when writing these passages. But he was a mathematician at Oxford University (his given name was Charles Dodgson), and at least one author has shown how his writings can be seen as metaphors for mathematical problems (Bayley 2009, 2010). Along these lines, mathematician Sanderson M. Smith has suggested that [Carroll simply inverted the equation for speed from “speed = distance/time” to “speed = time/distance”](http://www.herkimershideaway.org/writings/carroll.htm). Upon rearrangement, the latter gives “distance = time/speed.” Hence you must run very fast to stay in the same place. But how does the shifting landscape fit in? And why did Alice have to go the other way to meet the Red Queen? I would love to know.

It is perhaps worth pointing out that the phrase “Red Queen Hypothesis” can have two different meanings to evolutionary biologists. In the early 1970s, Leigh van Valen was grappling with data showing that the probability of extinction in very different organisms was independent of the age of the lineage. He reasoned that in coevolutionary interactions, the probability of one species driving the other species extinct could, in fact, be independent of lineage age (1973). It thus seems reasonable to suggest that both antagonists must run (coevolve) as fast as they can to prevent extinction. Graham Bell repurposed the phrase to mean within-population oscillations in host and parasite genotypes (1982). Hence, Van Valen’s idea is about macroevolution (speciation/extinction), while Bell’s idea is about microevolution. Even though van Valen’s use of the Red Queen metaphor was published first, I will use Bell’s microevolutionary meaning, as it perfectly captures the oscillating nature of genotype frequencies during host-parasite coevolution.[[17]](#footnote-98)

### 2.3.2 Conceptual roots of the Red Queen Hypothesis

Part of my goal is to show science as a process. As such it seems reasonable to discuss the origins of the Red Queen idea. One of the earliest statements alluding to the Red Queen Hypothesis came from W.D. Hamilton (1975). Hamilton was reviewing the books by Williams (1975) and Ghiselin (1974) for the *Quarterly Review of Biology*. Throughout the review, the reader can feel Hamilton’s frustration with their arguments. Towards the end, he makes these very abstract suggestions: “[I]t seems to me that we need environmental fluctuations around a trend line of change” and “For the source of these we may look to fluctuations and periodicities … generated by life itself.”

The quote does not specifically refer to parasites, but it does suggest that coevolutionary interactions, in general, could play a role in selecting for sex and recombination. In his memoirs, Hamilton (2001) clears this up, writing, “At that stage when I wrote the review, although I had not seen the particular relevance that parasitism might have, I had for many years seen sex looming ahead and had reached a stage of being excited by the possible primary role of biotic interaction. I had decided that it was in aspects of the interspecies struggle, and not survival in an inanimate environment, that I had to search for the main factor. Adaptation to new physical habitats might be made possible through sexuality but these adaptations could not be the main reason for its existence.”

Note that, here, Hamilton is specifically contrasting host-parasite coevolution with the Lottery Model, which relies on random changes in “physical habitats.”

At about the same time, a plant population biologist, Don Levin, was also writing on the paradox of sex/rec. In his paper, Levin specifically identified pathogens as a possible force selecting for recombination: “I propose that the persistent tracking of plant hosts by multiple pathogens and herbivores is a prime factor which prohibits the congealing of the genomes of species, especially those in closed communities” (Levin 1975).

Boom! By “prohibit the congealing of genomes,” Levin means, “selects for recombination.” The reference to “closed communities” means species that are tightly coevolving in the absence of homogenizing gene flow. This quote seems to be the first to specifically identify coevolving pathogens as a primary source of selection favoring the mixing of genomes. Levin’s idea was quickly followed by important conceptual contributions by Glesener & Tilman[[18]](#footnote-100) (1978), Jaenike[[19]](#footnote-101) (1978), and Lloyd (1980). In particular, Lloyd writes, “[B]iological interactions are more likely than unpredictable physical conditions to provide the kind of relentless, repetitive change that is necessary for sexual parents to be selected because of the genetic diversity that sex engenders.” Lloyd then turns this abstract idea into a specific prediction, which I would later test: “If this proves to be so, we will then be able to examine whether the occurrence of asexual reproduction is correlated with relaxation of the biological hostility of the environment.” Notice that, in the quotes presented above, both Hamilton and Lloyd were specifically predicting that coevolution is more important in selecting for sex than uncertain physical environments. But it is reasonable to ask, does it really matter? Aren’t both ideas fundamentally about bet hedging in uncertain environments? Yes: I think both ideas are about bet hedging. But the distinction still matters. The Lottery Model is about random shifts in the direction of selection; there is no selection against common genotypes unless the environment changes by chance in a way that disfavors them. By contrast, under the Red Queen, selection is frequency dependent. In fact, selection against common genotypes is the core of the model. Hence, the critical difference between the models is not so much about bet hedging but whether selection for sex/rec is directional (but randomly changing directions, a lottery) or frequency dependent (Red Queen). For example, parasites could be a source of directional selection for sex if they randomly changed which host genotypes they attacked. To my mind, that would be a Lottery Model. The Red Queen Hypothesis requires frequency-dependent selection generated by interactions between species.[[20]](#footnote-102) This is an important distinction.

Taking this view, the Red Queen Hypothesis may seem more closely related to the Tangled Bank model than to the Lottery Model, as both the Red Queen and the Tangled Bank rely on frequency-dependent selection. But the critical distinction here is that selection against common genotypes under the Tangled Bank relies on intraspecific competition in populations at carrying capacity (soft selection). The Red Queen relies on interspecific antagonistic coevolution, leading to parasite-mediated selection against common host genotypes.[[21]](#footnote-103)

In any case, looking back, it seems clear that the architects of the ecological hypotheses had two interrelated things in mind:

1. How can we explain sex/rec?
2. How do we understand the biogeographic and phylogenetic distributions of asexual reproduction?

As an evolutionary ecologist, I was drawn to the confluence of these questions. But other ideas were also interesting, such as the idea that sexual reproduction is favored because it reduces the interference between alleles at different loci (review in Otto 2021). I will cover some special cases of this latter idea in [Chapter 5](#sec-chap6).[[22]](#footnote-104)

## 2.4 Summary

1. Three ecological hypotheses have been proposed to explain the persistence of cross-fertilization in the face of competition with uniparental reproductive strategies, such as parthenogenesis or self-fertilization (following Bell 1982).
2. The Lottery Model is based on the possible advantages of diversifying offspring facing uncertain changes in the abiotic environment. Here selection is independent of both density and frequency.
3. The Tangled Bank and Frozen Niche-Variation Hypotheses are based on competition for resources when multiple resource types co-occur. Selection is frequency dependent, but the advantage to rare types only occurs when intraspecific competition is intense.
4. The Red Queen Hypothesis relies on parasite-mediated selection against common host genotypes. Such selection, when strong, can result in oscillatory changes in parasite and host alleles. These oscillations are sometimes called Red Queen dynamics.
5. A bet-hedging strategy reduces the variance in reproductive success over time, even if it reduces the arithmetic mean. Sexual reproduction under the Lottery Model is clearly a bet-hedging strategy. The Red Queen idea can perhaps also be seen as bet hedging.

|  |
| --- |
| Box 2.1. Levene’s model of multiple niche polymorphism. |
| It was widely thought that heterozygote advantage was required to maintain polymorphism at a single locus with two alleles. In the introduction to his paper, Levene wonders “whether it was in fact possible to have an equilibrium without the heterozygote being superior in any single niche” (1953). The paper is not easy to follow, eventhough the algebra is not difficult. Here I try to simplify the presentation.  Levene first assumes that the proportion of survivors coming from the niche is constant , independent of the genotypic composition of the niche (i.e., soft selection). He then assumes that the heterozygote has a relative fitness of one in all niches, giving . Let be the frequency of allele A, and let () be the frequency of allele B. The frequency of allele A in the next generation, , is then  The change in is simply . Under these assumptions, Levene showed that A allele will increase when rare (barring genetic drift) when |  where the left-hand side gives the harmonic mean fitness for genotype AA over all niches. The right-hand-side of the equation gives the harmonic mean fitness of the heterozygous genotype, AB, which is equal to one. Similarly, the B allele will increase when rare when  where the left-hand side gives the harmonic mean fitness for genotype BB over all niches. A genetic polymorphism is expected if both alleles can increase when rare; hence, **polymorphism is expected, in general, when the harmonic mean fitness for the heterozygote is greater than the harmonic mean fitness for either homozygote**.  But does this require that the AB genotype is the most fit in at least one niche? Levene gives a specific example to answer this question. He assumes two niches, where the proportion of survivors from both niches is equal (i.e., ). He then assumes genotypic fitness values, as given in the following table. It is important to note that the heterozygous genotype is not the most fit genotype in either niche.  Table 2.2: Levene’s Example   | **Genotype** | **Fitness Niche One** | **Fitness Niche Two** | **Arithmetic mean** | **Harmonic mean** | | --- | --- | --- | --- | --- | | **AA** | = 1.50 | = 0.67 | 1.09 | 0.93 | | **AB** | = 1.00 | = 1.00 | 1.00 | 1.00 | | **BB** | = 0.67 | = 1.50 | 1.09 | 0.93 |   For this example, the harmonic mean fitness for the heterozygote is greater than the harmonic mean fitness for either homozygote, thus meeting the conditions given by the equations above. Thus, the answer to Levene’s question is Yes. It is possible to have a genetic polymorphism without having heterozygote advantage in any single niche. And, interestingly, the polymorphism is expected even though the arithmetic mean fitness of the heterozygote is less than the arithmetic mean fitness of either homozygote. Finally, based on this example, it seems that a trade-off is required, such that the AA genotype does best in one niche, and the BB genotype does best in the other niche.  Nonetheless, Levene’s result suggests that overdominance for harmonic mean fitness is required for multiple niche polymorphism (Prout 1968). However, Timothy Prout showed that a polymorphism could be stable even if one allele is dominant, thus ruling out any kind of overdominance (1968). Let both the AA and AB genotypes have a fitness of one in both niches. Let the BB genotype have a fitness of 0.5 in niche one and a fitness of 1.67 in niche two. Assuming as above that both patches are equally common, we get the following table:  Table 2.3: Prout’s Example   | **Genotype** | **Fitness Niche One** | **Fitness Niche Two** | **Arithmetic mean** | **Harmonic mean** | | --- | --- | --- | --- | --- | | **AA** | = 1.00 | = 1.00 | 1.00 | 1.00 | | **AB** | = 1.00 | = 1.00 | 1.00 | 1.00 | | **BB** | = 0.50 | = 1.67 | 1.09 | 0.77 |   Prout showed that there would be a stable multiple niche polymorphism even under complete dominance, provided that the arithmetic mean fitness for BB is greater than one and the harmonic mean fitness for the BB genotype is less than one. So, clearly, overdominance for harmonic mean fitness is not required for a stable polymorphism.[[23]](#footnote-111)  The plot below shows as a function of for Prout’s model of dominance. Note that is positive when is near zero, and that is negative when is near one. There is an interior equilibrium near .   |  | | --- | |  | |

# 3. Contrasting the Ecological Hypotheses

|  |
| --- |
|  |

As I mentioned in [Chapter 1](#sec-why-sex), my dissertation focused on intertidal communities. I was especially interested in how two different barnacle morphs coexisted on rocky intertidal shores in the Northern Gulf of California. I had initially assumed that the two types were genetically determined and that they were likely to be different species ([Figure 3.1](#fig-3-2)). However, after years of false starts,[[24]](#footnote-120) I found that one of the two morphs was induced by chemical cues released by a predatory snail ([Figure 3.2](#fig-3-3)), and that the induced morph was more resistant to attack by this predator (1986c).[[25]](#footnote-121) Hence, the two morphs are not different species, but rather the result of phenotypic plasticity. In a blink of a field season, I went from being a community ecologist to an evolutionary biologist.

But why two morphs? Why didn’t selection favor unconditional development of the predator-resistant morph? Using predator-exclusion cages, I found that predation was concentrated near crevices in the reef, which the snails used during high tide as refuges (1986b). As the tide receded, the snails moved out from these crevices onto the exposed rock surfaces to forage on barnacles. When the tide returned, the snails motored back to the crevices, presumably to hide from snail-crushing rays that came in with the tide. This back-and-forth movement of snails created high-predation zones near crevices and low-predation zones far from crevices (about 20cm away). This finding explained why the predation-resistant morph was almost always found near crevices. Field experiments also showed that the predator-resistant morph grew more slowly and was less fecund than the typical volcano-shaped morph (Lively 1986b). Hence there is a trade-off. Taken together, the results suggested that plastic development was favored by natural selection to survive in the high-predation zones ([Box 3.1](#callout-4)). I would later come to think of adaptive plasticity as a type of variation strategy. Sexual reproduction can also be seen as a type of variation strategy (Lloyd 1984). And I was very fortunate to be able to study sexual reproduction after moving to New Zealand.

|  |
| --- |
| Figure 3.1: The two morphs of the intertidal barnacle, *Chthamalus anisopoma*. **Top**, the “bent” form is induced by exposure to chemical cues released by a specialized barnacle predator, the predatory gastropod *Acanthina angelica*. The bent or “hooded” form reduces the risk of successful attack by this predator. **Bottom**, the typical, conic form of the barnacle. The conic form is more fecund per unit size, and it grows more rapidly than the bent form, but it is also more susceptible to attack by the predator. Drawing by ZMD. |

|  |
| --- |
| Figure 3.2: Line drawing of the predatory snail *Acanthina angelica* attacking the bent form of the barnacle. Note that the predator has a spine on the outer margin of its shell. The spine is used to push through the opercular plates of barnacles, and it is very effective at penetrating and consuming the volcano-shaped form of the barnacle. The bent form of the barnacle is more resistant to attack of this kind because its aperture is less open to attack from above. Drawing by ZMD. |

I moved to New Zealand in 1984 just after defending my dissertation. My reason for moving to New Zealand was simple: my spouse (Lynda Delph) was there. Lynda had moved to New Zealand to study the evolution of plant breeding systems with Prof. David Lloyd. I did not have a job, but Lynda had a small stipend from the Fulbright Foundation. By the time I moved to New Zealand, we had only twelve dollars. But Lynda had found a flat in a dormitory at the University of Canterbury, where she worked as a “tutor.” Tutors at the time were usually graduate students who served as mentors for the resident students. We made many good friends during our time as tutors, and it was a fascinating total immersion into Kiwi culture. We did not have to pay rent, and we could eat for free in the cafeteria. We could then spend Lynda’s small Fulbright stipend on sampling trips.

Then I got very lucky. I was awarded a three-year postdoctoral fellowship from the NZ University Grants Committee. I had applied to work on the evolution of facultatively parthenogenetic nematodes, which represented a combination of my interests in developmental plasticity and sex.[[26]](#footnote-130) These topics were also very interesting to Wally Clark, a conceptual pioneer in the evolution of plasticity. He was also head of the Zoology Department at the University of Canterbury. I would not have received funding without the support of Prof. Clark. To my mind, the value of Clark’s work remains underestimated in general, but it had a big influence on me (e.g., Clark 1976).

I began looking for natural systems to study facultative parthenogenesis.[[27]](#footnote-131) To this end, I was reading Graham Bell’s incredible book on the evolution and genetics of sexual reproduction (1982). Searching the index, I found a reference to *Potamopyrgus antipodarum*, a New Zealand freshwater snail. Bell had cited Mike Winterbourn’s dissertation work on this snail (1970). Luckily for me, Prof. Winterbourn was just down the hall from me. I took the book to him, and I asked if the snails were, in fact, facultatively parthenogenetic. He said no; the snails were probably obligate asexuals, based on lab rearing experiments that he had done. He also said that most populations were all female, but some contained males. He then added that there was no obvious pattern to the distribution of males. Amazing! I immediately decided to work on these snails.[[28]](#footnote-132)

## 3.1 The Method of Multiple Working Hypotheses

As graduate students at the University of Arizona, we read some of the classics in the history and philosophy of science. Two of these papers concerned the method of contrasting multiple working hypotheses (Chamberlin 1890; Platt 1964).[[29]](#footnote-133) The idea is that multiple hypotheses should be simultaneously considered. Then, to the extent possible, the alternatives are forced to make different *a priori* predictions about the possible results. The hope is that all but one of the alternative hypotheses would be eliminated, leading to a “strong inference” that the remaining hypothesis is supported (Platt 1964). Thus, the focus is on falsifying one or more of the alternatives, rather than proving one of them (Popper 1959). Graham Bell used this same method to contrast the ecological models for sex by using data on the geographic distribution of asexual individuals across many plant and animal taxa (Bell 1982). The data led him to reject the Lottery Model ([Chapter 2](#sec-eco-hyp)). I decided to focus a similar test directly on the New Zealand snails ([Figure 3.3](#fig-3-4)).

|  |
| --- |
| Figure 3.3: The freshwater snail *Potamopyrgus antipodarum*. This small (3 – 6 mm) prosobranch snail evolved from marine ancestors (Phillips & Lambert 1990). Associated with the invasion of freshwater, the snail evolved an internal brood pouch, where the embryos hatch and develop before crawling out as juveniles. The snail also evolved parthenogenetic reproduction. Parthenogenesis and brooding are both rare traits in invertebrates, but they are often found together (Lively & Johnson 1994). Some *P. antipodarum* populations presently consist of a mixture of diploid sexual individuals and polyploid asexual females. The question under consideration here is, why have the sexual females persisted in these mixed population snails? What are the advantages of sexual reproduction? Photo credit: © [Bart Zijlstra](https://www.bartzijlstra.com). |

The snails (*Potamopyrgus antipodarum*) are often called mud snails, but I think the term is a misnomer. They live on rocks and vegetation in some of the most beautiful clear lakes, rivers, and streams in New Zealand (“Potamo” means river, not mud, in Greek). In any case, based on Winterbourn’s ecological work, streams seemed more unstable than lakes, as water flow can vary dramatically, especially during heavy rains in the mountains (Winterbourn *et al.* 1981). Hence, under the Lottery Model, streams should have more sexual females (and males) than lakes, because streams have more disturbance and less competition (see [Chapter 2](#sec-eco-hyp) for a comparison of models). By contrast, it seemed that competition for resources should be greater in lakes than in streams. Indeed, lake populations of the snail can be extremely dense. So, under the Tangled Bank, there should be more sexual females in lakes, where competition for resources is expected to be high. Finally, under the Red Queen Hypothesis, there should be more sexual females where the risk of infection by coevolving parasites is higher. As such, the different hypotheses could be forced to make different predictions, with the important caveat that infection might be correlated with habitat.

Some clarification regarding the prediction of the Red Queen Hypothesis might be useful here. Some people have asked me why the correlation between sex and infection is expected to be positive if, indeed, parasites are the selective force for sexual reproduction. For example, one could ask, if sex is so helpful in reducing infection risk, then shouldn’t the highly sexual populations have fewer, not more, parasites? That could, of course, be expected in an experiment where hosts across all populations were exposed to the same number of parasites. Then the more genetically diverse populations with higher frequencies of sexual females might be expected to have a lower prevalence of infection. But it is not the case that all natural populations have the same risk of infection. The idea under the Red Queen Hypothesis is that asexual females would replace sexual females where the risk of infection is low, and that sexual females would persist where the risk of infection is high, provided that the parasites are highly virulent. That is how the positive correlation could be generated. Nonetheless, the data could be expected to be very messy, especially if the frequency of sex oscillates over time in response to coevolutionary games with parasites.

The snails are infected by trematode worms, but I did not know anything about trematodes when I first began dissecting snails. I was just looking for males. Winterbourn told me that I would know a male snail when I saw one, as they have a penis just behind the right tentacle. But I had not observed any such structure on the many snails I collected from the streams around the university. I was beginning to think that I was missing something. Then one day, when I was dissecting a snail, hundreds of swimming things came out. Sperm, I thought. My first male! I took them to Wally Clark’s research technician, Jan McKenzie, to put under her fancy microscope. She informed me that sperm do not have eyes, that they do not have spines on their tails, and that they are, in fact, orders of magnitude smaller than these wiggling beasts under her lens. She was not impressed. I had perfectly fit the Kiwi stereotype of North American ecologists: good with statistics but no knowledge of real animals. She informed me that these swimming things were trematode larvae, **sterilizing parasites** of snails. Happily, we remained good friends, despite her disappointment in my training. And I had found my first infection, which meant that I might be able to test the Red Queen.

Perhaps embarrassingly, I had a scientific bias against the Red Queen going into the study. My bias was based on a study by May and Anderson (May & Anderson 1983). They showed that parasites had to kill infected individuals for sex to be favored over asex in hosts. Parasites are usually not that virulent; hence, it seemed to me that parasites could not provide sufficiently strong selection to *generally* favor sex. I will return to this important paper in another chapter and discuss how key assumptions of their model have been relaxed.

### 3.1.1 A side story on JMS

John Maynard Smith (JMS) was one of the most influential theoretical biologists in history of evolutionary thought. He was able to formulate and communicate novel ideas with apparent ease. Around the time that I was beginning to work on *Potamopyrgus*, JMS came to New Zealand, along with his wife, Sheila. He was invited by David Lloyd to spend time at University of Canterbury and to deliver three public lectures, which were all fantastic. During this time, JMS spent several weeks in New Zealand. Lynda, David, and I were lucky enough to hang out with him quite a bit. JMS was a remarkable individual. He could talk with anyone and show a sincere interest in their work. One morning, I was sitting next to JMS in the tearoom in the old Zoology Department. I was scared speechless. He kindly asked me what I was working on, so I told him about the snails. He knew of them! In fact, he had covered them in his book, *The Evolution of Sex*.[[30]](#footnote-139) He was very excited that I was working on these creatures, and he wanted to know my plan. I told him of my rough ideas for looking at the distribution of males as a way of contrasting the ecological hypotheses for sex. He looked directly at me, and said, “Interesting, but I hope the answer is not parasites” (or something like that). I asked him, why not parasites? He laughed out loud, and with a big smile he said: “Because Bill Hamilton thought of it first!” I could tell he was kidding. He then encouraged me to take the project on, and then he laughed again and added, “Whatever you do, don’t go and solve the problem of sex. Sex is too much bloody fun to have an answer!”

Toward the end of their time in New Zealand, JMS, Sheila, David, Lynda, and I did a trip together around the South Island. The whole time was incredible. Just listening to David and John talk about evolutionary theory was a scientific dream. Towards the end of our trip, we were all together in a restaurant at the Hermitage (near Mt. Cook) on the night that JMS turned 65 and formally retired. Our server, a young alpinist working to support his climbing in the Southern Alps, asked JMS, “I think that I saw a documentary about you. Are you famous?” JMS (smiling and intrigued) asked the alpinist what he remembered. Without hesitation, the alpinist recited a perfect overview of evolution by natural selection. JMS was clearly touched. Almost exactly half-way around the world from Sussex England, in a small township in New Zealand, JMS met someone whom he had influenced with his work. And this was on the very night of his retirement.

The next day, we drove to a small lake near Mt. Cook that David knew about: Lake Alexandrina. It was a glorious day, and we decided that we might as well collect some snails. JMS waded into the water and proceeded to collect a handful of *Potamopyrgus* from the shallow rocks. He handed the snails to me. He then laughed and said, “When you publish your study, I want to know the outcome for these exact snails.” As it turned out, Lake Alexandrina has a mixed population of sexual and asexual snails, and it has been the primary focus of our long-term studies on *Potamopyrgus*. The snail team still refers to this original site of collection as “JMS.” Interestingly, JMS is one of the most dynamic sites in the whole lake.

## 3.2 The Distribution of Male Snails

To contrast the alternative ecological hypotheses, I sampled snails from lakes and streams across the South Island of New Zealand. I could drive Lynda’s Volkswagen bug to most of the lakes, but I had to backpack into many. Unfortunately, my time working in the Sonoran Desert had not prepared me for the steep climbs, heavy rains, and chest-deep river crossings on the South Island. I did not take enough food or dry clothes on one trip, and my desert hiking boots disintegrated. I got my butt kicked. But it was wonderful to have an excuse to see remote parts of the South Island, especially after I got better gear and gained a better understanding of the New Zealand bush.

I collected and dissected hundreds of snails from each of 29 streams and 22 lakes, mostly on the South Island. I recorded sex (male or female) and infection by the trematodes that Winterbourn (1973) described. I reasoned that the frequency of males in a population must be strongly correlated with the frequency of sexual females simply because males are only produced by sexual females.[[31]](#footnote-142) The results showed that there were more males in lakes than in streams, which was inconsistent with the Lottery Model, but it was consistent with the Tangled Bank Model. However, male frequency was better predicted by the frequency of trematode infection than by habitat *per se* (Lively 1987). Hence, surprisingly, the results favored the Red Queen Hypothesis. I presented these findings to a small group at David Lloyd’s flat, and they convinced me to submit to *Nature* (1987).[[32]](#footnote-143)

A fascinating paper on the same topic was published in *Nature* at about the same time. This paper was also based on a strong-inference test comparing the Red Queen and the Tangled Bank. The authors, Austin Burt and Graham Bell, examined recombination in mammals (1987). They reasoned that under the Red Queen Hypothesis, longer-lived mammals would have higher rates of recombination because more genetic mixing would be favored as the asymmetry in host/parasite generation time increased. In contrast, the Tangled Bank Model predicted that shorter-lived mammals would have higher rates of recombination because they have larger litters, and recombination might lead to reduced competition among the more diverse offspring. Their results were stunning. Recombination was tightly and positively related to longevity in natural populations.[[33]](#footnote-144) The Red Queen was again supported.

Based on these studies in *Nature*, I was beginning to think that parasites might be a factor in selecting for cross-fertilization in hosts. But my study as well as the study of Burt and Bell (1987) were based on correlations. And every scientist knows that correlation is not causation. On the other hand, these correlations were predicted *a priori* by Lloyd (1980) and others (Bell 1982; e.g., Glesener & Tilman 1978). The Red Queen was supported by the data, but the data were not used to generate the hypothesis. Using the same data to both generate and substantiate hypotheses is where the problem arises with correlation, especially when multiple factors are considered in “fishing expeditions.” But forcing different hypotheses to make different *a priori* predictions about the direction of correlations is, to my mind, a powerful way to evaluate alternatives.

As a brief aside, I cannot help but mention the human toll taken by R.A. Fisher’s use of “correlation is not causation” as a way to plant doubt in the mind of smokers about the now-obvious risks of smoking (Gould 1991; Stolley 1991). Fisher was a consultant for the tobacco industry, and he did the industry a great service at the cost of human lives. I would also add that no test statistic is causation; F statistics derived from analysis of variance are not causation. Causation might be inferred from well-designed experiments, but no statistical test is causation. Analytical theory is not causation either, as is well demonstrated by the theoretical literature on sex/recombination. Causation instead may be inferred when multiple independent lines of evidence point to similar solutions. I think that Levins (1966) was correct when he wrote, “Hence our truth is the intersection of independent lies.”[[34]](#footnote-145) Although he was referring specifically to mathematical models, the same principle applies to biological systems. Ideally, the multiple lines of evidence would include long-term field observations of individual populations, broader biogeographic patterns across populations, and direct experiments on multiple independent systems, as well as multiple theoretical forays into the conditions under which the hypothesis is expected to hold.

In any case, my view by 1987 was that the Red Queen Hypothesis merited serious consideration.[[35]](#footnote-146) For my own data, I now asked whether the correlation between sex and infection was a “red herring.” In other words, could the correlation be generated because of something else? Yes, it could. Here is how it might work. First, infection could be higher in dense host populations as expected under theory (Anderson & May 1979; May & Anderson 1979). Second, there might be more sex in dense populations because asexual reproduction is favored in sparse populations as a way for individuals to ensure reproduction even in the absence of conspecific mates (Gerritsen 1980; Lloyd 1980; Tomlinson 1966). This latter idea is called the “Reproductive Assurance Hypothesis.” Hence, one could find a positive correlation between sex and infection as a simple consequence of epidemiology and selection for reproductive assurance. So, I decided to sample again, this time focusing on South Island lakes ([Figure 3.4](#fig-3-5) & [Figure 3.5](#fig-3-6)) while also collecting data on snail density. The results were consistent with the epidemiological expectations, as there was a marginally significant positive relationship between snail density and infection prevalence, but there was no support for the Reproductive Assurance Hypothesis (Lively 1992). Finally, the previously observed positive relationship between sex and infection held ([Figure 3.4](#fig-3-5)).[[36]](#footnote-147) The Red Queen was still in the running.

|  |
| --- |
| Figure 3.4: Results from surveys of New Zealand lakes and streams showing percent males against the prevalence of female infection by all species of trematodes. Note the upper left side of the graph. There are no highly sexual populations where parasites are rare or absent, which suggests that asexuals have replaced sexuals were parasite-mediated selection is weak. This result is consistent with Lloyd’s prediction given in [Chapter 2](#sec-eco-hyp). Circles represent stream populations (Lively 1987) plus two river samples. Gray triangles represent lake populations (Lively 1987). Black triangles represent lake and tarn populations (Lively 1992). The correlation is positive and statistically significant. |

These results suggested that parasite-mediated selection might contribute to the persistence of sex in mixed populations of sexual and asexual snails. It is of particular interest, perhaps, to note that there are no populations with a high proportion of males in samples where parasites were rare or absent ([Figure 3.4](#fig-3-5)). This finding is consistent with David Lloyd’s 1980 prediction that asexuals should dominate in populations “with a relaxation of biological hostility” (see above) (1980). But the results are messy.

There are several reasons for why the results might be expected to be messy. One is that prevalence of infection might not give a good estimate of the strength of parasite-mediated selection (Lively 2001). For example, infected snails might die at a faster rate than uninfected snails because of the energetic demands of infection. In addition, infected snails are more likely than uninfected snails to forage after sunrise, which exposes them to predation by their final hosts, ducks (Levri & Fisher 2000; Levri & Lively 1996). Prevalence of infection might also fluctuate over time as the genetic diversity in the host population changes and/or as the final hosts move among locations. We now know that the prevalence of infection varies greatly among years and among sites in the same lake (Gibson *et al.* 2016). Thus, detecting a significant correlation between sex and infection could be dicey, even if parasites were solely responsible for the short-term maintenance of sex in mixed populations.[[37]](#footnote-152)

Along these lines, many of the points in [Figure 3.4](#fig-3-5) represent a single sample taken at one site at one point in time. This limitation likely introduces “noise” into the data, especially for samples where parasites are only periodically common. For this reason, Jukka Jokela and I selected 20 of the best sampled lakes from the data set given in [Figure 3.4](#fig-3-5).[[38]](#footnote-153) We resampled all 20 lakes 10 – 15 years after my original samples. We found that prevalence of infection was highly correlated between sample periods as was male frequency (Lively & Jokela 2002). We then averaged the data for each lake under the assumption that the averages would better represent both the frequency of males and the prevalence of infection for each lake. With these data, the correlation between male frequency and infection prevalence was both positive and significant.[[39]](#footnote-154)

None of this is meant to imply proof of the Red Queen Hypothesis or that density dependence and random environmental change are not relevant for a full understanding of the problem.[[40]](#footnote-155) But the results do imply that the Red Queen Hypothesis was (and still is) worthy of further study.

## 3.3 Summary

1. The co-occurrence of discrete morphs is inherently interesting to evolutionary biologists. Genetic diversity is also inherently interesting.
2. Some populations of the New Zealand freshwater snail, *Potamopyrgus antipodarum*, contain both sexual and asexual females. Other populations are mostly or completely parthenogenetic. This makes the snail a very useful natural system for contrasting alternative hypotheses for the maintenance of sexual reproduction.
3. The prevalence of sterilizing trematode larvae is a better predictor sexual reproduction in the snail than habitat (lakes vs. streams), thus favoring the Red Queen Hypothesis over the alternative ecological hypotheses.
4. Comparing the *a priori* predictions of multiple working hypotheses can be helpful to evaluate competing ideas. Field studies of natural systems may be required to fully understand why cross-fertilization is so common.

|  |
| --- |
| Figure 3.5: The distribution of male and female *Potamopyrgus antipodarum* across New Zealand. The percentage of males is given in blue; the percentage of females is given in red. Pie charts enclosed in boxes are for lakes and tarns that are very close together. The large pie on the left-hand side shows the average frequencies of males and females across all samples. (Redrawn from Lively 1992.) |

|  |
| --- |
| Box 3.1. |
| As part of my dissertation research, I constructed a game-theoretic model of selection on three strategies:   1. canalized development into a high-fecundity morph, 2. canalized development to a low-fecundity, predation-resistant morph, 3. induced development into the low fecundity defended morph in the presence of predators.   The model examined evolutionary stability for a range of frequencies for two patches (high predation risk and low predation risk) across a range of values for the accuracy of the cue predicting future predation risk (Lively 1986a).  An example of the output is shown below. The results show that any of the three strategies can be an ESS in part of the parameter space.[[41]](#footnote-163) High reliability of the cue and intermediate patch frequencies favor the plastic strategy. Genetic polymorphism is expected under a relatively narrow set of conditions. Mixtures of constitutive and plastic strategies can also be stable. Note that changing the patch frequencies leads to evolutionary change. For example, reducing the frequency of the low-risk patch can lead to a selective sweep (arrow a). It can also lead to the evolution of plasticity (arrow b) and the evolution of canalized development (arrow c). Increasing the accuracy of the cue can also lead to the evolution of plasticity. It might be especially interesting to note that the trajectory of arrow b would give the appearance of saltatory change followed by stasis (i.e., punctuated equilibrium) (Levinton 1988). Also note that the conditions for a genetic polymorphism are relatively narrow. Redrawn from Lively (1999a) assuming a small cost to plasticity.   |  | | --- | |  | |

# 4. Self/Non-Self Recognition and Local Adaptation

|  |
| --- |
|  |

One way to falsify the Red Queen is to experimentally show that the expectations of the hypothesis are not met. One expectation is that parasites would quickly become adapted to infecting their local host populations. Here is the logic. If parasites are closely tracking common host genotypes in their local (sympatric) populations, then they should be better, on average, at infecting sympatric hosts than foreign (allopatric) hosts. If this is not the case, then parasites would seem unlikely (at least to me) to be a factor selecting for sexual reproduction.

I am often asked why we would expect the parasites to be better at infecting their local hosts instead of the opposite. Why shouldn’t hosts evolve to be more resistant to their local parasites than to allopatric parasites? It is a fair question. One common answer is that parasites are locally adapted to host populations because they have faster generation times. But that cannot be the whole answer. Theory has shown that parasites can be locally adapted even when there is no generation-time asymmetry (Gandon & Michalakis 2002; Lively 1999b). Instead, I think the answer has more to do with the underlying genetic basis for infection.

What, then, is the genetic basis for infection? This was unknown, but I was assuming that all animal hosts have a self/non-self recognition system, such that they can detect foreign tissues (e.g., parasites or tissue grafts) that do not match their own. Sponges, for example, accept tissue grafts from self, but reject grafts from unrelated individuals of the same species (review in Elda Gaino & Magnino 1999). This ability to reject foreign tissues seems widely conserved (Buss 1990). I was also assuming that the self-/non-self-recognition system is genetically variable and that different host genotypes would dominate in different populations. Parasite genotypes that match the most common local host genotypes would be favored by natural selection, and these parasite genotypes should increase in frequency. This should lead to local adaptation by the parasites. Fortunately, one can test for local adaptation using reciprocal cross-inoculation experiments.

## 4.1 Experimental Studies of Local Adaption

While I was still a post-doc in New Zealand, I set up two reciprocal cross-inoculation experiments to test for local adaptation by the parasites. I knew from my field surveys that one species of sterilizing trematode was especially common in lake populations of the snail. This species was not formally described, but Jan McKenzie sent it to a trematode expert in France, who thought it belonged in the genus *Microphallus*; hence I will refer to it as *Microphallus* sp.[[42]](#footnote-172) The life cycle of *Microphallus* turns out to be especially crucial to the story. The adult worms are tiny simultaneous hermaphrodites that live in the intestines of ducks. They cross-fertilize and produce eggs that are shed with the duck feces into the environment. In most trematodes, the eggs normally hatch in water, thereby releasing a swimming larval stage (miracidia), which actively swims to and penetrates the body of snails. This is the case for the trematodes that cause the human disease, Schistosomiasis. But, in this New Zealand species of *Microphallus*, the eggs hatch not in the environment but rather after being ingested by snails. The larvae then penetrate the snail from the inside. If the snail’s immune system does not recognize the larvae as foreign tissue, the larvae reproduce asexually, producing several hundred cysts (metacercaria) in the snail. These cysts completely replace the reproductive tissue in both males and females. Infected snails are sterilized ([Figure 4.1](#fig-4.1)).

|  |
| --- |
| Figure 4.1: Left panel: the life cycle of the trematode *Microphallus*. The adult worms live in the intestines of waterfowl and wading birds (black stilts). They produce cross-fertilized eggs, which are released into lakes and streams with the bird’s feces. The eggs hatch following ingestion by snails. Infection results in the asexual production of hundreds of cysts (called metacercaria). These cysts “hatch” and mature following ingestion by ducks, thus completing the life cycle. Drawing by Zoe M Dinges. |

# 5. The Ratchet and the Red Queen

In 1988, Indiana University advertised for an assistant professor in population biology, with emphasis on disease ecology. Lynda and I both applied. Happily, we were offered a split position in Biology in which we each got half salary. It may not sound like a good deal, but we were thrilled. It is not easy for a dual-career couple in the same field. We relocated to Bloomington in January of 1990, arriving during a cold snap (-20&degC). We moved into a university house; but we did not know enough to have the electricity turned on before arrival. Luckily, we still had our down sleeping bags, which we had purchased for field work in the Southern Alps. Aside from the chilly start, moving to Bloomington was the beginning of an academic dream come true. Most of this book aims to highlight the work of my incredible students and colleagues at IU.

## 5.1 The Problem

# References

Anderson, R.M. & May, R.M. (1979). Population biology of infectious diseases: Part 1. *Nature*, 280, 361–367.

Antonovics, J. & Ellstrand, N.C. (1984). Experimental studies of the evolutionary significance of sexual reproduction. I. A test of the frequency-dependent selection hypothesis. *Evolution*, 38, 103–115.

Bayley, M. (2009). Alice’s adventures in algebra: Wonderland solved. *New Scientist*.

Bayley, M. (2010). Algebra in wonderland. *New York Times*.

Bell, G. (1982). *The masterpiece of nature: The evolution and genetics of sexuality*. University of California Press, Berkeley.

Bell, T., Freckleton, R.P. & Lewis, O.T. (2006). [Plant pathogens drive density-dependent seedling mortality in a tropical tree](https://doi.org/10.1111/j.1461-0248.2006.00905.x). *Ecology Letters*, 9, 569–574.

Blasco-Costa, I., Seppälä, K., Feijen, F., Zajac, N., Klappert, K. & Jokela, J. (2019). [A new species of *atriophallophorus* deblock & rosé, 1964 (trematoda: Microphallidae) described from *in vitro*-grown adults and metacercariae from *Potamopyrgus antipodarum* (gray, 1843) (mollusca: tateidae)](https://doi.org/10.1017/S0022149X19000993). *Journal of Helminthology*, 94, 1–15.

Burt, A. & Bell, G. (1987). Mammalian chiasma frequencies as a test of two theories of recombination. *Nature*, 326, 803–805.

Buss, L. (1990). Competition within and between encrusting clonal invertebrates. *Trends in Ecology and Evolution*, 5, 352–356.

Chamberlin, T.C. (1890). The method of multiple working hypotheses. *Science*, 15, 92–96.

Churchill, F.B. (1979). Sex and the single organism: biological theories of sex in mid nineteenth century. *Studies in the History of Biology*, 3, 139–177.

Churchill, F.B. (1997). Life before model systems: General zoology at August Weismann’s institute. *American Zoologist*, 37, 260–268.

Clark, W.C. (1976). The environment and the genotype in polymorphism. *Zoological Journal of the Linnean Society*, 58, 255–262.

Dagg, J. (2016). [On recognising the paradox of sex](https://doi.org/10.3998/ptb.6959004.0008.003). *Philosophy, Theory, and Practice in Biology*, 8.

Darwin, C. (1862). [On the two forms, or dimorphic condition, in the species of *Primula*, and on their remarkable sexual relations.](https://doi.org/10.1111/j.1095-8312.1862.tb01218.x) *Journal of the Proceedings of the Linnean Society of London (Botany)*, 6, 77–96.

Darwin, C. (1868). *The variation of plants and animals under domestication*. 1st edn. John Murray, London.

Darwin, C. (n.d.). [Letter no. 2869](https://www.darwinproject.ac.uk/letter/?docId=letters/DCP-LETT-2869.xml). Darwin Correspondence Project.

Elda Gaino, G.B. & Magnino, G. (1999). [Self/non‐self recognition in sponges](https://doi.org/10.1080/11250009909356270). *Italian Journal of Zoology*, 66, 299–315.

Elliott, L.P. & Brook, B.W. (2007). Revisiting Chamberlin: Multiple working hypotheses for the 21st century. *Bioscience*, 57, 608–614.

Ellstrand, N.C. & Antonovics, J. (1985). Experimental studies of the evolutionary significance of sexual reproduction II. A test of the density-dependent selection hypothesis. *Evolution*, 39, 657–666.

Fisher, R.A. (1941). Average excess and average effect of a gene substitution. *Annals of Eugenics*, 11, 53–63.

Gandon, S. & Michalakis, Y. (2002). Local adaptation, evolutionary potential and host-parasite coevolution: Interactions between migration, mutation, population size and generation time. *Journal of Evolutionary Biology*, 15, 451–462.

Gerritsen, J. (1980). Sex and parthenogenesis in sparse populations. *American Naturalist*, 115, 718–742.

Ghiselin, M.T. (1974). *The economy of nature and the evolution of sex*. University of California Press, Berkeley.

Gibson, A.K., Jokela, J. & Lively, C.M. (2016). Fine-scale spatial covariation between infection prevalence and susceptibility in a natural population. *American Naturalist*, 188, 1–14.

Glesener, R.R. & Tilman, D. (1978). Sexuality and the components of environmental uncertainty: Clues from geographical parthenogenesis in terrestrial animals. *American Naturalist*, 112, 659–673.

Gould, S.J. (1991). The smoking gun of eugenics. *Natural History*, 100, 8–17.

Hamilton, W.D. (1975). Innate social aptitudes of man: An approach from evolutionary genetics. In: *Biosocial anthropology* (ed. Fox, R.). Malaby Press, London, pp. 133–153.

Hamilton, W.D. (2001). *Narrow roads of gene land: Volume 2: Evolution of sex*. Oxford University Press.

Hazel, W., Smock, R. & Lively, C.M. (2004). The ecological genetics of conditional strategies. *American Naturalist*, 163, 888–900.

Jaenike, J. (1978). A hypothesis to account for the maintenance of sex within populations. *Evolutionary Theory*, 3, 191–194.

Kelley, S.E. (1993). [Viruses and the advantage of sex in *Anthoxanthum odoratum*: A review](https://doi.org/10.1111/j.1442-1984.1993.tb00072.x). *Plant Species Biology*, 8, 217–223.

Kelley, S.E. (1994). [Viral pathogens and the advantage of sex in the perennial grass *Anthoxanthum odoratum*](https://doi.org/10.1098/rstb.1994.0146). *Philosophical Transactions of the Royal Society of London B, Biological Sciences*, 346, 295–302.

Kelley, S.E., Antonovics, J. & Schmitt, J. (1988). A test of the short-term advantage of sexual reproduction. *Nature*, 331, 714–716.

Kondrashov, A.S. (1993). Classification of hypotheses on the advantage of amphimixis. *Journal of Heredity*, 84, 372–387.

Kuhn, T.S. (1970). *The structure of scientific revolutions*. 2nd edn. University of Chicago Press.

Lehtonen, J., Jennions, M.D. & Kokko, H. (2012). The many costs of sex. *Trends in Ecology & Evolution*, 27, 172–178.

Levene, H. (1953). Genetic equilibrium when more than one ecological niche is available. *American Naturalist*, 87, 331–333.

Levin, D.A. (1975). Pest pressure and recombination systems in plants. *American Naturalist*, 109, 437–451.

Levins, R. (1966). The strategy of model building in population biology. *American Scientist*, 54, 421–431.

Levinton, J. (1988). *Genetics, paleontology, and macroevolution*. Cambridge University Press.

Levri, E.P. & Fisher, L.M. (2000). [The effect of a trematode parasite (*Microphallus sp.*) on the response of the freshwater snail *Potamopyrgus antipodarum* to light and gravity](https://doi.org/10.1163/156853900502565). *Behaviour*, 137, 1141–1151.

Levri, E.P. & Lively, C.M. (1996). The effects of size, reproductive condition, and parasitism on foraging behaviour in a freshwater snail, *Potamopyrgus antipodarum*. *Animal Behaviour*, 51, 891–901.

Lewontin, R.C. (1971). [The effect of genetic linkage on the mean fitness of a population](https://doi.org/10.1073/pnas.68.5.984). *Proceedings of the National Academy of Sciences of the United States of America*, 68, 984–986.

Lively, C.M. (1986a). Canalization versus developmental conversion in a spatially variable environment. *American Naturalist*, 128, 561–572.

Lively, C.M. (1986b). Competition, comparative life histories, and maintenance of shell dimorphism in a barnacle. *Ecology*, 67, 858–864.

Lively, C.M. (1986c). [Predator-induced shell dimorphism in the acorn barnacle *Chthamalus anisopoma*](https://doi.org/10.1111/j.1558-5646.1986.tb00466.x). *Evolution*, 40, 232–242.

Lively, C.M. (1987). Evidence from a New Zealand snail for the maintenance of sex by parasitism. *Nature*, 328, 519–521.

Lively, C.M. (1992). Parthenogenesis in a freshwater snail: Reproductive assurance versus parasitic release. *Evolution*, 46, 907–913.

Lively, C.M. (1996). Host-parasite coevolution and sex. *Bioscience*, 46, 107–109.

Lively, C.M. (1999a). Developmental strategies in spatially variable environments: Barnacle shell dimorphism and strategic models of selection. In: *The ecology and evolution of inducible defenses* (eds. Tollrian, R. & Harvell, C.D.). Princeton University Press, pp. 245–258.

Lively, C.M. (1999b). [Migration, virulence, and the geographic mosaic of adaptation by parasites.](https://doi.org/10.1086/303210) *The American Naturalist*, 153, S34–S47.

Lively, C.M. (2001). Trematode infection and the distribution and dynamics of parthenogenetic snail populations. *Parasitology*, 123, S19–S26.

Lively, C.M. (2006). The ecology of virulence. *Ecology Letters*, 9, 1089–1095.

Lively, C.M. (2009). The maintenance of sex: Host-parasite coevolution with density-dependent virulence. *Journal of Evolutionary Biology*, 22, 2086–2093.

Lively, C.M., Hazel, W.N., Schellenberger, M.J. & Michelson, K.S. (2000). Predator-induced defense: Variation for inducibility in an intertidal barnacle. *Ecology*, 81, 1240–1247.

Lively, C.M. & Johnson, S.G. (1994). [Brooding and the evolution of parthenogenesis: Strategy models and evidence from aquatic invertebrates](https://doi.org/10.1098/rspb.1994.0054). *Proceedings of the Royal Society of London. Series B: Biological Sciences*, 256, 89–95.

Lively, C.M., Johnson, S.G., Delph, L.F. & Clay, K. (1995). [Thinning reduces the effect of rust infection on jewelweed (*Impatiens capensis*)](https://doi.org/10.2307/1940718). *Ecology*, 76, 1859–1862.

Lively, C.M. & Jokela, J. (2002). Temporal and spatial distributions of parasites and sex in a freshwater snail. *Evolutionary Ecology Research*, 4, 219–226.

Lively, C.M. & Lloyd, D.G. (1990). The cost of biparental sex under individual selection. *American Naturalist*, 135, 489–500.

Lively, C.M., Xu, J. & Ben-Ami, F. (2021). [Causation without correlation: Parasite-mediated frequency-dependent selection and infection prevalence](https://doi.org/10.1098/rsbl.2021.0321). *Biology Letters*, 17, 20210321.

Lloyd, D.G. (1980). Benefits and handicaps of sexual reproduction. *Evolutionary Biology*, 13, 69–111.

Lloyd, D.G. (1984). Variation strategies of plants in heterogeneous environments. *Biological Journal of the Linnean Society*, 21, 357–385.

May, R.M. & Anderson, R.M. (1979). Population biology of infectious diseases: Part II. *Nature*, 280, 455–461.

May, R.M. & Anderson, R.M. (1983). [Epidemiology and genetics in the coevolution of parasites and hosts](https://doi.org/10.1098/rspb.1983.0075). *Proceedings of the Royal Society of London. Series B. Biological Sciences*, 219, 281–313.

Maynard Smith, J. (1971). What use is sex? *Journal of Theoretical Biology*, 30, 319–335.

Maynard Smith, J. (1976). A short-term advantage for sex and recombination through sib-competition. *Journal of Theoretical Biology*, 63, 245–258.

Maynard Smith, J. (1978). *The evolution of sex*. Cambridge University Press.

Maynard Smith, J. & Hoekstra, R. (1980). [Polymorphism in a varied environment: How robust are the models?](https://doi.org/10.1017/S0016672300013926) *Genetics Research*, 35, 45–57.

Meirmans, S. (2009). The evolution of the problem of sex. In: *Lost sex: The evolutionary biology of parthenogenesis* (eds. Schön, I., Martens, K. & Dijk, P. van). Springer, London, pp. 21–46.

Negovetic, S. & Jokela, J. (2001). Life history variation, phenotypic plasticity and maintenance of subpopulation structure in a freshwater snail. *Ecology*, 82, 2805–2815.

Otto, S.P. (2021). Selective interference and the evolution of sex. *Journal of Heredity*, 112, 9–18.

Peters, A.D. & Lively, C.M. (1999). The Red Queen and fluctuating epistasis: A population genetic analysis of antagonistic coevolution. *American Naturalist*, 154, 393–405.

Peters, A.D. & Lively, C.M. (2007). Short- and long-term benefits and detriments to recombination under antagonistic coevolution. *Journal of Evolutionary Biology*, 20, 1206–1217.

Philippi, T. & Seger, J. (1989). Hedging one’s evolutionary bets, revisited. *Trends in Ecology & Evolution*, 4, 41–44.

Phillips, N.R. & Lambert, D.M. (1990). [A cladistic analysis of species of the molluscan genus *Potamopyrgus* based on allozyme data](https://doi.org/10.1080/03014223.1990.10422600). *New Zealand Journal of Zoology*, 17, 257–263.

Platt, J.R. (1964). Strong inference. *Science*, 146, 347–353.

Popper, K. (1959). *The logic of scientific discovery*. Hutchinson & Company, London.

Prout, T. (1968). Sufficient conditions for multiple niche polymorphism. *The American Naturalist*, 102, 493–496.

Salathe, M., Kouyos, R.D., Regoes, R.R. & Bonhoeffer, S. (2008). Rapid parasite adaptation drives selection for high recombination rates. *Evolution*, 62, 295–300.

Schmid-Hempel, P. & Jokela, J. (2002). Socially structured populations and evolution of recombination under antagonistic coevolution. *American Naturalist*, 160, 403–408.

Siebold, C.T.E. von. (1856). *Wahre parthenogenesis bei schmetterlingen und bienen. Ein beitrag zur fortpflanzungsgeschichte der thiere*. William Engelmann, Leipzig.

Soper, D.M., Neiman, M., Savytskyy, O.P., Zolan, M.E. & Lively, C.M. (2013). Spermatozoa production by triploid males in the New Zealand freshwater snail *Potamopyrgus antipodarum*. *Biological Journal of the Linnean Society*, 110, 227–234.

Stearns, S.C. (2000). Daniel Bernoulli (1738): Evolution and economics under risk. *Journal of Biosciences*, 25, 221–228.

Stolley, P.D. (1991). When genius errs: R. A. Fisher and the lung cancer controversy. *American Journal of Epidemiology*, 133, 416–425.

Tomlinson, J. (1966). The advantages of hermaphroditism and parthenogenesis. *Journal of Theoretical Biology*, 11, 54–58.

Valen, L. van. (1973). A new evolutionary law. *Evolutionary Theory*, 1, 1–30.

Vrijenhoek, R.C. (1979). Factors affecting clonal diversity and coexistence. *American Zoologist*, 19, 787–797.

Vrijenhoek, R.C. (1998). Animal clones and diversity. *Bioscience*, 48, 617–628.

Vrijenhoek, R.C. & Parker, E.D. (2009). Geographical parthenogenesis: General purpose genotypes and frozen niche variation. In: *Lost sex: The evolutionary biology of parthenogenesis* (eds. Schön, I., Martens, K. & Dijk, P.). Springer, London, pp. 99–131.

Wallace, B. (1975). Hard and soft selection revisited. *Evolution*, 29, 465–473.

Whitton, J., Sears, C., Baack, E. & Otto, S. (2008). The dynamic nature of apomixis in the angiosperms. *International Journal of Plant Sciences*, 169, 169–182.

Williams, G.C. (1975). *Sex and evolution*. Princeton University Press.

Winterbourn, M.J. (1970). The New Zealand species of *Potamopyrgus* (Gastropoda: Hydrobiidae). *Malacologia*, 10, 283–321.

Winterbourn, M.J. (1973). Larval trematoda parasitising the New Zealand species of *Potamopyrgus* (Gastropoda: Hydrobiidae). *Mauri Ora*, 2, 17–30.

Winterbourn, M.J., Rounick, J.S. & Cowie, B. (1981). Are New Zealand stream ecosystems really different? *New Zealand Journal of Marine and Freshwater Research*, 15, 321–328.

1. The trematode worm was not formally described until 30 years later (Blasco-Costa *et al.* 2019). As it turns out, it belongs in the genus *Atriophallophorus*, rather than *Microphallus*, and it was very appropriately named after Mike Winterbourn: *A. winterbourni*. But I am going to call it *Microphallus* in this book, as that is what we called it in our early papers. [↑](#footnote-ref-33)
2. Plasticity is more mainstream now than it was in the 1980s. One reviewer of my paper was incredulous and recommended rejection from *Evolution* because the bent morph was not “genetically determined.” The Associate Editor (John Endler) rejected the review and accepted the paper. Clearly, it is the developmental strategy that is genetically determined, not the morph per se (Hazel *et al.* 2004; see also Lively *et al.* 2000). [↑](#footnote-ref-39)
3. Facultative parthenogenesis is used to mean environmentally cued production of parthenogenetic females. I was originally planning to work on a nematode population that produced a mixture of sexual males and females at high density but only parthenogenetic females at low density. [↑](#footnote-ref-48)
4. We were trained ask questions first and then seek suitable organisms to address the questions. This was the tradition before model-systems research took over (Churchill 1997). [↑](#footnote-ref-57)
5. Prof. Winterbourn was supportive of my work from this first day. He shared his knowledge of the snail system and of freshwater ecology, in general, with great enthusiasm. In addition, Mike met with my Ph.D. students and took them into the field. This book would not have been possible without Prof. Winterbourn. [↑](#footnote-ref-58)
6. See also Elliott and Brook (2007). They point out crucial differences between Chamberlin and Platt including that Chamberlin allowed for multiple ideas to be partially correct, which is important for [Chapter 5](#sec-chap6). [↑](#footnote-ref-59)
7. With respect to *Potamopyrgus* (along with a parthenogenetic beetle) Maynard Smith (1978) wrote, “Further Investigations of these cases could be most interesting.” When I met JMS, I did not know (or did not remember) that he had written this. But I think that he was correct. [↑](#footnote-ref-65)
8. The trematode worm was not formally described until 30 years later (Blasco-Costa *et al.* 2019). As it turns out, it belongs in the genus *Atriophallophorus*, rather than *Microphallus*, and it was very appropriately named after Mike Winterbourn: *A. winterbourni*. But I am going to call it *Microphallus* in this book, as that is what we called it in our early papers. [↑](#footnote-ref-72)
9. Plasticity is more mainstream now than it was in the 1980s. One reviewer of my paper was incredulous and recommended rejection from *Evolution* because the bent morph was not “genetically determined.” The Associate Editor (John Endler) rejected the review and accepted the paper. Clearly, it is the developmental strategy that is genetically determined, not the morph per se (Hazel *et al.* 2004; see also Lively *et al.* 2000). [↑](#footnote-ref-73)
10. Facultative parthenogenesis is used to mean environmentally cued production of parthenogenetic females. I was originally planning to work on a nematode population that produced a mixture of sexual males and females at high density but only parthenogenetic females at low density. [↑](#footnote-ref-74)
11. We were trained ask questions first and then seek suitable organisms to address the questions. This was the tradition before model-systems research took over (Churchill 1997). [↑](#footnote-ref-76)
12. Prof. Winterbourn was supportive of my work from this first day. He shared his knowledge of the snail system and of freshwater ecology, in general, with great enthusiasm. In addition, Mike met with my Ph.D. students and took them into the field. This book would not have been possible without Prof. Winterbourn. [↑](#footnote-ref-78)
13. See also Elliott and Brook (2007). They point out crucial differences between Chamberlin and Platt including that Chamberlin allowed for multiple ideas to be partially correct, which is important for [Chapter 5](#sec-chap6). [↑](#footnote-ref-79)
14. With respect to *Potamopyrgus* (along with a parthenogenetic beetle) Maynard Smith (1978) wrote, “Further Investigations of these cases could be most interesting.” When I met JMS, I did not know (or did not remember) that he had written this. But I think that he was correct. [↑](#footnote-ref-84)
15. This assumption turned out to be not strictly true. Polyploid females occasionally produce males, although they seem unlikely to be very fertile (Soper *et al.* 2013). [↑](#footnote-ref-87)
16. The group included Mark McKone. Mark was a post-doc with David, and his comments were especially influential. Fifteen years later, I would become Ph.D. advisor to one of Mark’s star mentees at Carleton College, Maurine Neiman. [↑](#footnote-ref-96)
17. There we also some very interesting outliers. Domesticated mammals had very strong positive residuals for the rate of recombination. This result suggests that recombination was selected by frequent changes in the targets of artificial selection by humans. [↑](#footnote-ref-98)
18. Therefore, we attempt to treat the same problem with several alternative models each with different simplifications but with a common biological assumption. Then, if these models, despite their different assumptions, lead to similar results we have what we can call a robust theorem which is relatively free of the details of the model. Hence our truth is the intersection of independent lies (Levins 1966). [↑](#footnote-ref-100)
19. I was also persuaded by elegant experimental studies on sweet vernal grass, which showed a density-independent advantage to having a rare genotype (Antonovics & Ellstrand 1984; Ellstrand & Antonovics 1985). Later studies showed that the rare advantage was likely due to escape from infection (Kelley *et al.* 1988; 1993, 1994). [↑](#footnote-ref-101)
20. The partial correlation between percent male and prevalence of infection, while controlling for habitat, is highly significant (, ). However, the partial correlation between percent male and habitat, while controlling for prevalence of infection, is marginally significant (, ). Similar results were gained after males were excluded from the calculation of infection prevalence, which controls for any sex-specific differences in susceptibility (as shown in [Figure 3.4](#fig-3-5)); specifically, prevalence of infection in females was significantly correlated with male frequency while controlling for habitat (, ), but the converse was not true (, ). [↑](#footnote-ref-102)
21. Using computer simulations, we recently found that detecting a significant positive correlation between clonal diversity and infection prevalence would only be expected in a fraction of parameter space, even when parasites were solely responsible for the maintenance of diversity (Lively *et al.* 2021). [↑](#footnote-ref-103)
22. In this smaller sample of 20 lakes, the correlation between male frequency and infection prevalence was positive but not statistically significant. [↑](#footnote-ref-104)
23. ; for log10 transformed data; . [↑](#footnote-ref-111)
24. The trematode worm was not formally described until 30 years later (Blasco-Costa *et al.* 2019). As it turns out, it belongs in the genus *Atriophallophorus*, rather than *Microphallus*, and it was very appropriately named after Mike Winterbourn: *A. winterbourni*. But I am going to call it *Microphallus* in this book, as that is what we called it in our early papers. [↑](#footnote-ref-120)
25. Plasticity is more mainstream now than it was in the 1980s. One reviewer of my paper was incredulous and recommended rejection from *Evolution* because the bent morph was not “genetically determined.” The Associate Editor (John Endler) rejected the review and accepted the paper. Clearly, it is the developmental strategy that is genetically determined, not the morph per se (Hazel *et al.* 2004; see also Lively *et al.* 2000). [↑](#footnote-ref-121)
26. Facultative parthenogenesis is used to mean environmentally cued production of parthenogenetic females. I was originally planning to work on a nematode population that produced a mixture of sexual males and females at high density but only parthenogenetic females at low density. [↑](#footnote-ref-130)
27. We were trained ask questions first and then seek suitable organisms to address the questions. This was the tradition before model-systems research took over (Churchill 1997). [↑](#footnote-ref-131)
28. Prof. Winterbourn was supportive of my work from this first day. He shared his knowledge of the snail system and of freshwater ecology, in general, with great enthusiasm. In addition, Mike met with my Ph.D. students and took them into the field. This book would not have been possible without Prof. Winterbourn. [↑](#footnote-ref-132)
29. See also Elliott and Brook (2007). They point out crucial differences between Chamberlin and Platt including that Chamberlin allowed for multiple ideas to be partially correct, which is important for [Chapter 5](#sec-chap6). [↑](#footnote-ref-133)
30. With respect to *Potamopyrgus* (along with a parthenogenetic beetle) Maynard Smith (1978) wrote, “Further Investigations of these cases could be most interesting.” When I met JMS, I did not know (or did not remember) that he had written this. But I think that he was correct. [↑](#footnote-ref-139)
31. This assumption turned out to be not strictly true. Polyploid females occasionally produce males, although they seem unlikely to be very fertile (Soper *et al.* 2013). [↑](#footnote-ref-142)
32. The group included Mark McKone. Mark was a post-doc with David, and his comments were especially influential. Fifteen years later, I would become Ph.D. advisor to one of Mark’s star mentees at Carleton College, Maurine Neiman. [↑](#footnote-ref-143)
33. There we also some very interesting outliers. Domesticated mammals had very strong positive residuals for the rate of recombination. This result suggests that recombination was selected by frequent changes in the targets of artificial selection by humans. [↑](#footnote-ref-144)
34. Therefore, we attempt to treat the same problem with several alternative models each with different simplifications but with a common biological assumption. Then, if these models, despite their different assumptions, lead to similar results we have what we can call a robust theorem which is relatively free of the details of the model. Hence our truth is the intersection of independent lies (Levins 1966). [↑](#footnote-ref-145)
35. I was also persuaded by elegant experimental studies on sweet vernal grass, which showed a density-independent advantage to having a rare genotype (Antonovics & Ellstrand 1984; Ellstrand & Antonovics 1985). Later studies showed that the rare advantage was likely due to escape from infection (Kelley *et al.* 1988; 1993, 1994). [↑](#footnote-ref-146)
36. The partial correlation between percent male and prevalence of infection, while controlling for habitat, is highly significant (, ). However, the partial correlation between percent male and habitat, while controlling for prevalence of infection, is marginally significant (, ). Similar results were gained after males were excluded from the calculation of infection prevalence, which controls for any sex-specific differences in susceptibility (as shown in [Figure 3.4](#fig-3-5)); specifically, prevalence of infection in females was significantly correlated with male frequency while controlling for habitat (, ), but the converse was not true (, ). [↑](#footnote-ref-147)
37. Using computer simulations, we recently found that detecting a significant positive correlation between clonal diversity and infection prevalence would only be expected in a fraction of parameter space, even when parasites were solely responsible for the maintenance of diversity (Lively *et al.* 2021). [↑](#footnote-ref-152)
38. In this smaller sample of 20 lakes, the correlation between male frequency and infection prevalence was positive but not statistically significant. [↑](#footnote-ref-153)
39. ; for log10 transformed data; . [↑](#footnote-ref-154)
40. I think density dependence is critically important. Disease transmission is certainly density dependent (Anderson & May 1979; May & Anderson 1979). Virulence may also be density dependent (Bell *et al.* 2006; Lively *et al.* 1995; Lively 2006). Habitat partitioning may also play a role in the distribution of sexual females among depth-stratified habitats (Negovetic & Jokela 2001). [↑](#footnote-ref-155)
41. “Parameter space” represents all possible combinations of variables as defined by the model. In [Box 3.1](#callout-4), different strategies are favored for different combinations of variables (i.e., different parts of the parameter space). [↑](#footnote-ref-163)
42. The trematode worm was not formally described until 30 years later (Blasco-Costa *et al.* 2019). As it turns out, it belongs in the genus *Atriophallophorus*, rather than *Microphallus*, and it was very appropriately named after Mike Winterbourn: *A. winterbourni*. But I am going to call it *Microphallus* in this book, as that is what we called it in our early papers. [↑](#footnote-ref-172)