

# Strong Selection is Necessary for Evolution of Blindness in Cave Dwellers

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## 1 **Abstract**

2 Blindness has evolved repeatedly in cave-dwelling organisms, and investigating loss of sight presents an  
3 opportunity to understand the operation of fundamental evolutionary processes, including drift, selection,  
4 mutation, and migration. The observation of blind organisms has prompted many theories for their  
5 blindness, including loss-by-disuse and selection against eye development when eyes are not used. Here  
6 we have developed a model that shows just how strong selection must be for blind populations of a cave-  
7 dwelling species to evolve. We used approximations to determine levels of selection that would result in  
8 caves containing only sighted individuals, only blind individuals, or a stable population of both. We then  
9 incorporated drift into the model using simulations. Based on our model, strong selection is necessary  
10 for the evolution of blindness unless immigration rates are extremely low. Drift decreased the fixation of  
11 blindness in populations, although for intermediate levels of migration the level of selection required to fix  
12 blindness decreased substantially. We hypothesize that this degree of selection may be due to phototaxis  
13 in sighted individuals, who move toward the light leaving only blind individuals in the cave.

## 14 Introduction

15 Blindness has evolved repeatedly across taxa in caves, creating nearly a thousand cave-dwelling species  
16 and many more populations (Culver et al., 2000; Dowling et al., 2002; Bradic et al., 2012; Coghill et al.,  
17 2014). However, many populations of blind individuals experience some level of immigration, which would  
18 be expected to prevent the fixation of blindness in a newly established population (Awise and Selander,  
19 1972; Bradic et al., 2012; Coghill et al., 2014). Thus, blind cave-dwelling populations of typically sighted  
20 species pose an interesting challenge to our understanding of evolutionary biology. Namely, how does a  
21 fixed phenotype evolve from low frequency despite immigration?

22 Darwin suggested that eyes would be lost by “disuse” (Darwin, 1859). We now consider this hypothesis the  
23 “neutral mutation hypothesis” — random mutations can accumulate in eye related genes or regulatory  
24 regions when, as in caves, there is no purifying selection to eliminate them. However, the accumulation of  
25 mutations (mutation pressure) causing blindness would take a long time to result in fixation of blindness  
26 in populations on its own (Barr, 1968). Thus, it is genetic drift combined with mutation pressure that would  
27 lead to blindness (Kimura and King, 1979; Borowsky, 2015). Genetic drift is the increases the frequency of  
28 blindness alleles created by mutations: eyes become increasingly less functional and finally disappear  
29 completely (Wilkins, 1988).

30 There are a variety of cave dwelling vertebrates and invertebrates (referred to as cavernicoles, troglaphiles,  
31 and troglobites). However, much of the work on the evolution of blindness has focused on the blind  
32 form of cavefishes, e.g. the Mexican tetra (*Astyanax mexicanus*) and Atlantic molly (*Poecilia mexicana*).  
33 For example, the hypothesis of relaxed selection is supported by the observation of a high number of  
34 mutations in cavefish putative eye genes (Hinaux et al., 2013; Protas et al., 2006; Gross et al., 2009). However,  
35 developmental evidence does not support this hypothesis: cavefish embryos begin eye development, but  
36 the eye disappears in larvae (Langecker et al., 1993; Jeffery et al., 2003). Random mutations should occur  
37 in genes controlling early eye development as well. Furthermore, although drift would often lead to loss  
38 of seeing individuals joining a population of blind individuals, this model depends on developing a high  
39 frequency of blindness in a cave population simply by drift in isolation.

40 Alternatively, the “adaptation hypothesis” suggests that there is a cost to an eye; thus, individuals without

41 eyes have greater fitness resulting in the eventual elimination of seeing individuals. This cost may either  
42 come from the energy required to develop a complex structure or due to the vulnerability of the eye (Barr,  
43 1968; Strickler et al., 2007; Jeffery, 2005; Protas et al., 2007; Niven, 2008; Niven and Laughlin, 2008; Moran  
44 et al., 2015). Another hypothesis states that sight is not lost not due to the “cost” of development but  
45 due to pleiotropic mutations selected for other traits. For example, in Mexican tetra increased expression  
46 of Hedgehog (Hh; Jeffery, 2005) likely affects feeding structures, allowing better foraging in low light  
47 conditions (Jeffery, 2005, 2001). However, increased Hh signaling inhibits pax6 expression, which results  
48 in eye loss during development (Jeffery, 2005; Yamamoto et al., 2004). Alternatively, cryptic variation may  
49 be maintained in normal conditions, and expressed as blindness only in case of stress, such as entry into  
50 the cave (Rohner et al., 2013). When the cryptic variation is “unmasked” it is then exposed to selection  
51 and could become fixed in the population.

52 These mechanisms of selection would result in the evolution of a blind population occurring quite slowly.  
53 Furthermore, given that there is often migration from surface to cave populations and that and these  
54 populations can interbreed, it seems that blind phenotypes should be lost (Avisé and Selander, 1972). One  
55 possibility is that the strength of selection for blindness is large enough to counter immigration (Avisé and  
56 Selander, 1972). Although blind fish maintained in the dark in the lab do not appear to have an advantage  
57 of this magnitude (Sadoglu, 1967), recent work suggests a very high cost to developing neural tissue,  
58 including eyes (Moran et al., 2015).

59 Due to the immigration of individuals from the surface and the expected level of selection for the eyeless  
60 phenotype, cave populations are an example of migration–selection balance (Wright, 1969; Hedrick, 2011;  
61 Nagylaki, 1992). However, much of the work in this area has explored the “invasion” of a novel allele or the  
62 maintenance of polymorphism, rather than fixation of different alleles in different populations (Yeaman  
63 and Otto, 2011; Yeaman and Whitlock, 2011).

64 Here we have developed a model that shows just how strong selection must be to generate blind popula-  
65 tions of a cave-dwelling species. Incorporating genetic drift into the model actually increases the level  
66 of selection required for fixation. This level of selection is not compatible with the hypothesis that eyes  
67 are lost due to drift or that eyes are lost due to selection for improved foraging and pleiotropy (assuming  
68 pleiotropy imposes only weak selection). However, if eye development imposes a high cost (Moran et al.,

69 2015) then the adaptation hypothesis is plausible. Alternatively, we suggest reconsidering the historic  
70 hypothesis that a high level of selection is due to migration of seeing individuals, who are strongly pho-  
71 totactic, out of the cave (Lankester, 1925; Romero, 1985). Thus, we suggest that a standing presence of  
72 blindness alleles, combined with extreme loss of sighted individuals in the cave, is a likely scenario leading  
73 to evolution of blind cave-dwelling populations.

## 74 **Model and Analysis**

### 75 **Assumptions**

76 Consider a species with two populations: surface-dwelling and cave-dwelling. We are interested in  
77 determining when the cave population will evolve blindness, i.e. become comprised of mostly blind  
78 individuals, as has occurred in numerous natural systems. We first assume that the surface and cave  
79 populations do not experience drift (i.e. populations are of infinite size). Additionally, immigration from  
80 the surface population into the cave affects the allele frequency in the cave, but emigration from the  
81 cave to the surface does not affect the surface population, as we assume that the surface population is  
82 significantly larger than the cave. Generations are discrete and non-overlapping, and mating is random.  
83 We track a single biallelic locus, where  $b^+$  is the dominant, seeing allele, and where  $b^-$  is the recessive,  
84 blindness allele. The frequencies of  $b^-$  are denoted by  $\tilde{q} \in [0, 1]$  on the surface and  $q \in [0, 1]$  in the cave.  
85 On the surface, we assume that blindness is selected against, and  $\tilde{q}$  is dictated by mutation-selection  
86 balance.

### 87 **Calculating the frequency of the blindness allele**

88 Within the cave, the life cycle is as follows. (1) Embryos become juveniles and experience constant selection  
89 with relative fitnesses of  $w_{b^- b^-} = 1 + s$  and  $w_{b^+ b^-} = w_{b^- b^+} = 1$ , where  $s \geq 0$ . (2) Juveniles migrate into  
90 and out of the cave such that a fraction  $m$  of adults come from the surface and  $1 - m$  from the cave, where  
91  $0 \leq m \leq 1$ . (3) Adults generate gametes with one-way mutation, where  $0 \leq u \leq 1$  is the probability that a  
92  $b^+$  becomes a non-functional  $b^-$ . (4) Gametes unite randomly to produce embryos. Given this life cycle,

93 we calculate the allele frequency of the daughter generation ( $q'$ ) via standard equations:

$$q_j = \frac{(1+s)q^2 + 1q(1-q)}{(1+s)q^2 + 1(1-q^2)} \quad \text{selection} \quad (1a)$$

$$q_a = q_j(1-m) + \tilde{q}m \quad \text{immigration} \quad (1b)$$

$$q' = q_a + (1-q_a)u \quad \text{mutation} \quad (1c)$$

94 The change in allele frequency in one generation is

$$\Delta q = q' - q = \frac{sq^2 [1 - q - m(1-u)(1-\tilde{q})] + [\tilde{q}m(1-u) + u - q(m(1-u) + u)]}{1 + sq^2} \quad (2)$$

95 Furthermore,  $b^-$  is maintained at a stable equilibrium on the surface:  $\tilde{q} = \sqrt{u/z}$ , where  $z$  is the selection  
 96 coefficient against  $b^-$  on the surface and  $u \leq z \leq 1$ .

### 97 **Identifying equilibrium frequencies of the blindness allele**

98 The model we have developed is an example of migration-selection balance (Figure 1; Wright, 1969; Hedrick,  
 99 2011; Nagylaki, 1992). An equilibrium exists for this model when  $\Delta q = 0$ . Assuming  $s > 0$  and setting  
 100  $\Delta q = 0$ , Equation 2 can be rearranged into the following cubic polynomial

$$-q^3 + Bq^2 + Cq + D = 0 \quad (3)$$

where

$$B = 1 - m(1-u)(1-\tilde{q})$$

$$C = -\frac{m(1-u) + u}{s}$$

$$D = \frac{\tilde{q}m(1-u) + u}{s}$$

101 There are three possible roots of this equation, corresponding to three possible equilibria. Depending  
102 on the parameter values, Equation 3 may have three real roots or one real root and two imaginary roots.  
103 While the values of the roots of this polynomial can be expressed analytically, these equations are too  
104 complex to be helpful for understanding the system. For simplicity, we will let  $\hat{q}$  represent any possible  
105 equilibrium, and  $\hat{q}_a \leq \hat{q}_b \leq \hat{q}_c$ , stand for the roots of Equation 3.

Rather than tackling the equilibria directly, we first demonstrate that the cave has a protected polymorphism. A protected polymorphism exists if the allele frequency moves away from both fixation and extinction, i.e.  $\Delta q < 0$  for  $q = 1$  and  $\Delta q > 0$  for  $q = 0$ . For  $q = 0$

$$\Delta q = \tilde{q}m(1 - u) + u = (1 - u)m\sqrt{u/z} + u$$

and  $q = 0$  will be an equilibrium if  $u = 0$ ; otherwise  $\Delta q > 0$  at  $q = 0$  due to immigration of individuals containing  $b^-$  (Figure 2). For  $q = 1$

$$\Delta q = -m(1 - \tilde{q})(1 - u) = -m(1 - \sqrt{u/z})(1 - u)$$

106 and  $q = 1$  will be an equilibrium if  $m = 0$  or  $u = 1$ ; otherwise  $\Delta q < 0$  at  $q = 1$  due to immigration of  
107 individuals containing  $b^+$  (Figure 2). Thus a protected polymorphism always exists except at the edge cases  
108  $m = 0$ ,  $u = 0$ , and  $u = 1$ . In biological terms, the cave population will be polymorphic despite directional  
109 selection for  $b^-$  if there is some immigration from the surface population and the surface population  
110 is polymorphic. For  $s = 0$ , there is only one equilibrium, and it is near 0. For large  $s$ , there is only one  
111 equilibrium, and it is near 1. Three equilibria will only exist for moderate levels of selection (Figure 2).

112 **Validity of equilibria.** An equilibrium is only valid in our model if it is real and between  $[0, 1]$ ; otherwise,  
113 it is not biologically interpretable in this system. The lower bound for any equilibrium is  $\frac{m\tilde{q}(1-u)+u}{m(1-u)+u}$  if  
114  $m > 0$  or  $u > 0$ ; otherwise it is 0 (Lemma 1). The upper bound for any equilibrium is  $1 - m(1 - u)(1 - \tilde{q})$   
115 (Lemma 2). Thus if any equilibrium is real it is valid. The only exception to this rule is the edge case when  
116  $s = m = u = 0$ . In this case, all evolutionary forces are eliminated, and  $q' = q$  for all  $q$ . Here every possible  
117 value of  $q$  is an equilibrium, although only  $q \in [0, 1]$  makes any sense. Furthermore, it is important to

118 note that if  $m > 0$ ,

$$\tilde{q} \leq \frac{m\tilde{q}(1-u) + u}{m(1-u) + u} \implies \tilde{q} \leq \hat{q} \quad (4)$$

119 indicating that the equilibrium frequencies in the cave are always greater than the allele frequency on the  
120 surface.

121 **Approximations.** In order to study equilibria, we will simplify our model by assuming that  $u \ll 1$  such  
122 that  $1 - u \approx 1$  and

$$\Delta q \propto sq^2 [1 - q - m(1 - \tilde{q})] + [\tilde{q}m + u - q(m + u)] \quad (5)$$

123 **Weak-selection approximation.** If selection is weak, then an equilibrium exists near  $\tilde{q}$  (Figure 2). We  
124 use a second-order Taylor series at  $q = 0$  to determine the upper bound on  $s$  for the presence of three  
125 equilibria (i.e. when selection is so strong that an equilibrium near  $\tilde{q}$  does not exist). The second-order  
126 series allows us to determine the lower two equilibrium points, although this approximation is inaccurate  
127 as  $q$  increases. This approximation gives us

$$\Delta q \approx s(1 - m)q^2 - (m + u)q + m\tilde{q} + u \quad (6)$$

after assuming that  $1 - \tilde{q} \approx 1$ . This equation has two roots, which are the lowest two of three total equilibria,

$$\hat{q}_{a,1} = \frac{m + u - \sqrt{(m + u)^2 - 4s(1 - m)(m\tilde{q} + u)}}{2s(1 - m)}$$

$$\hat{q}_{b,1} = \frac{m + u + \sqrt{(m + u)^2 - 4s(1 - m)(m\tilde{q} + u)}}{2s(1 - m)}$$

128 These two roots exist only if

$$0 < \sqrt{(m + u)^2 - 4s(1 - m)(m\tilde{q} + u)} \implies s < \frac{(m + u)^2}{4(1 - m)(m\tilde{q} + u)} \quad (7)$$

129 which provides us with an estimate of the upper bound on  $s$  for the presence of three equilibria.

130 The derivative of Equation 6 is  $\frac{d\Delta q}{dq}(q) = 2s(1 - m)q - (m + u)$ , and an equilibrium will be stable if  
 131  $-2 < \frac{d\Delta q}{dq}(\hat{q}) < 0$ . From this, it can be easily shown that  $\hat{q}_{a,1}$  is stable and  $\hat{q}_{b,1}$  is unstable.

132 **Strong-selection approximation.** In order to determine the lower bound on  $s$  for the presence of three  
 133 equilibria, we assume that selection is strong enough such that  $u/s \approx 0$  and  $\tilde{q}/s \approx 0$ . Therefore,

$$\Delta q \propto -q \left[ q^2 - [1 - m(1 - \tilde{q})]q + m/s \right] \quad (8)$$

and the equilibria can be described as

$$\begin{aligned} \hat{q}_{a,2} &= 0 \\ \hat{q}_{b,2} &= \frac{1}{2} \left( 1 - m(1 - \tilde{q}) - \sqrt{[1 - m(1 - \tilde{q})]^2 - \frac{4m}{s}} \right) \\ \hat{q}_{c,2} &= \frac{1}{2} \left( 1 - m(1 - \tilde{q}) + \sqrt{[1 - m(1 - \tilde{q})]^2 - \frac{4m}{s}} \right) \end{aligned}$$

134 The latter two equilibria will exist only if

$$s > \frac{4m}{[1 - m(1 - \tilde{q})]^2}$$

135 which provides us an estimate of the lower bound for the presence of three equilibria.

136 The derivative of Equation 8 is  $\frac{d\Delta q}{dq}(q) = -3q^2 + 2[1 - m(1 - \tilde{q})]q - m/s$ , and it can be easily shown  
 137 that  $\hat{q}_{b,2}$  is unstable and  $\hat{q}_{c,2}$  is stable.

138 **Validity of approximations.** By substituting  $\hat{q}_{a,1}$  and  $\hat{q}_{b,1}$  back into Equation 5, we obtain  $\Delta q = -sq^2(\hat{q} - \tilde{q}m)$ .

139 Thus,  $\Delta q \leq 0$ , which indicates that  $\hat{q}_{a,1}$  overestimates  $\hat{q}_a$  and that  $\hat{q}_{b,1}$  underestimates  $\hat{q}_b$ . By substituting  
 140  $\hat{q}_{b,2}$  and  $\hat{q}_{c,2}$  back into Equation 5, we find that  $\Delta q = \tilde{q}m + u(1 - q)$ . Thus  $\Delta q \geq 0$ , which indicates that  
 141  $\hat{q}_{b,2}$  overestimates  $\hat{q}_b$  and that  $\hat{q}_{c,2}$  underestimates  $\hat{q}_c$ . However, the error in our approximations is slight  
 142 (Figure 3).

143 **Dynamics.** The dynamics of the evolution of the cave population depend on the parameter values and  
 144 the starting allele frequency,  $q_0$  (Table 1). If there is one equilibrium value, then  $b^-$  will evolve to be the  
 145 major allele in the population if  $\hat{q} > 0.5$ . If there are three equilibria, and  $\hat{q}_c > 0.5$ , then  $b^-$  will become  
 146 the major allele only if its initial frequency is above the threshold  $\hat{q}_b$ .

147 Based on these approximations the dynamics of the system can be summarized as follows. First, there  
 148 are three possible equilibria:  $\hat{q}_a \approx \hat{q}_{a,1}$ ,  $\hat{q}_b \in [\hat{q}_{b,1}, \hat{q}_{b,2}]$ , and  $\hat{q}_c \approx \hat{q}_{c,2}$ . Second, there are four possible  
 149 equilibria configurations: 1, 2a, 2b, and 2c.

150 Case 1,  $\frac{(m+u)^2}{4(1-m)(m\hat{q}+u)} < \frac{4m}{[1-m(1-\hat{q})]^2}$ : only one equilibrium exists, and it is stable. The population will  
 151 always evolve towards it.

152 Case 2,  $\frac{4m}{[1-m(1-\hat{q})]^2} < \frac{(m+u)^2}{4(1-m)(m\hat{q}+u)}$ : depending on the strength of  $s$ , this case may have one of three  
 153 possible configurations:

154 Case 2a,  $0 \leq s < \frac{4m}{[1-m(1-\hat{q})]^2}$ : Only one equilibrium exists,  $\hat{q}_a$ , and it is stable. The population will always  
 155 evolve towards it.

156 Case 2b,  $\frac{4m}{[1-m(1-\hat{q})]^2} < s < \frac{(m+u)^2}{4(1-m)(m\hat{q}+u)}$ : All three equilibria exist;  $\hat{q}_a$  and  $\hat{q}_c$  are stable, while  $\hat{q}_b$  is  
 157 unstable. If the population starts below  $\hat{q}_b$ , it will evolve towards  $\hat{q}_a$ . If it starts above  $\hat{q}_b$ , it will evolve  
 158 towards  $\hat{q}_c$ .

159 Case 2c,  $\frac{(m+u)^2}{4(1-m)(m\hat{q}+u)} < s$ : only one equilibrium,  $\hat{q}_c$ , exists, and it is stable. The population will always  
 160 evolve towards it.

## 161 **The evolution of blindness**

162 When the cave population is founded, its initial allele frequency will likely match the equilibrium frequency  
 163 on the surface ( $q_0 = \hat{q}$ ). Since  $\tilde{q} < \hat{q}$ , the allele frequency in the cave will increase due to selection until it  
 164 reaches the lowest equilibrium. If this equilibrium is  $\geq \frac{1}{2}$ , we consider the population to have evolved  
 165 blindness. If there is only a single equilibrium, and  $s \geq \frac{4m-8\hat{q}m-4u}{1-2m(1-\hat{q})}$  (i.e.  $m \leq \frac{s+4u}{2s(1-\hat{q})+4-8\hat{q}}$ , Lemma 3),  $b^-$   
 166 will evolve to become the major allele. If there are three equilibria (Case 2b), then the population will not

167 evolve blindness: the maximum value of  $\hat{q}_a \approx \frac{1}{8}$ . Therefore, blindness will only evolve if

$$s \geq \max \left\{ \frac{4m - 8\tilde{q}m - 4u}{1 - 2m(1 - \tilde{q})}, \frac{(m + u)^2}{4(1 - m)(m\tilde{q} + u)} \right\} \approx s \geq \frac{m}{4(\tilde{q} + \frac{u}{m})} \quad (9)$$

168 This approximation is valid when  $\tilde{q} < \frac{1}{16}$  and  $u \ll m \ll 1$ . We analytically calculated ultimate allele  
169 frequencies for our model and compared it to the above approximation (Figure 4A), and we also explored  
170 the approximation when  $u$  and  $\tilde{q}$  are varied (Figure 5).

## 171 **Finite-population simulations**

### 172 **Constant migration**

173 To investigate the impact of drift on our model, we simulated diploid populations of size  $N = 1000$ , where  
174 the frequency of adults was determined by drawing  $2N$  alleles from a binomial distribution with mean  $q_a$   
175 (Equation 1b).  $q'$  was calculated based on the post-drift adult allele frequency and immigration. For each  
176 set of parameters, we recorded the average  $q'$  frequency across these 100 populations at specific time  
177 points.

178 For high migration rates, the average allele frequency is similar to the infinite model, except that drift allows  
179 some populations that have three equilibria to evolve blindness (Figure 4B). However, at low migration  
180 rates ( $m < u/\tilde{q} = 10^{-4}$ ), populations have low average frequency of  $b^-$  at 10 thousand generations,  
181 unless  $s > 1$ . As immigration decreased, these populations became dependent on *de novo* mutations  
182 to produce  $b^-$ , which is a slow process. At 5 million generations, which is close to the estimated age of  
183 cavefish populations Gross (2012), the average allele frequency is a better match to the results from the  
184 the infinite-population model (Figure 4C); although, selection is ineffective for  $s < 1/2N = 5 \times 10^{-4}$ .

### 185 **Episodic migration**

186 Because cave and surface populations are may be connected intermittently due to flooding, we simulated  
187 periods of immigration followed by periods of isolation following a first-order Markov process. The

188 probability of switching between isolation and immigration or vice versa was 10% every generation.  
189 Results for the intermittently connected simulations were nearly identical to previous simulations, with  
190 the exception that at high levels of migration and selection, drift was more effective in increasing allele  
191 frequencies (Figure 4D).

## 192 **Discussion**

193 Both our model and simulations show that strong selection (characterized as  $s > 0.05$ ; Rieseberg and  
194 Burke, 2001) is necessary for a cave population to evolve blindness. Our simulations demonstrate that  
195 genetic drift, which is likely to occur in small cave populations, markedly decreases the fixation of these  
196 rare alleles, resulting in the need for even greater selection.

## 197 **Model**

198 Our model demonstrates that without drift, blindness occurs in cave populations only when  $s$  is large or  
199  $m$  is very small. This result is logical: given low levels of immigration, selection increases the frequency  
200 of the blindness allele. In contrast, for high levels of immigration there will always be some sighted fish.  
201 This case essentially results in a single population, for which selection in the large surface population for  
202 sighted fish outweighs small-scale selection in the cave (Nagylaki and Lou, 2008). In other words, gene  
203 flow prevents local adaptation, as expected. For intermediate levels of immigration, there will be sighted  
204 fish unless selection removes them.

205 What is surprising about our result is the level of selection required to fix blindness in the population. This  
206 result contrasts with the level of the selection found in most cases of local adaptation. Sadoglu (1967)  
207 argued that given the observed number of populations of blind fish living in caves, drift would have fixed  
208 at least one for a useful eye, However, because all populations appear to be blind or with a significantly  
209 reduced eye, this indicates strong selection for “degenerative genes”.

## 210 **Drift**

211 The “neutral mutation hypothesis” is equivalent to no selection for the blindness allele in a small popula-  
212 tion. This hypothesis relies on drift to fix populations for blindness. Thus, we explored the interaction of  
213 immigration, selection and blindness with simulations. However, our simulations including drift produce  
214 similar results to our model, with two notable exceptions. First, as observed previously, drift removes the  
215 blindness allele when it occurs at low levels; thus, for low immigration rates, populations consisted primar-  
216 ily of sighted fish, except when very strong selection immediately increased the frequency of blindness.  
217 This is opposite of predicted by the “neutral mutation hypothesis”.

218 When migration is very high, drift has minimal effect, as the two populations essentially function as one,  
219 with the surface population swamping the smaller cave population. However, for intermediate levels of  
220 migration the level of selection required to fix blindness decreased substantially. In this case, immigration  
221 is increasing the frequency of the blindness allele in each generation, allowing more chances for selection  
222 to overcome drift. This result is consistent with the observation of Blanquart et al. (2012)

223 Allowing populations to evolve for longer periods of time does increase the likelihood that a population  
224 can drift to fixation for blindness. As for higher levels of immigration, more generations results in a greater  
225 chance for a blindness mutation to occur in the population, and for drift to increase the frequency of this  
226 allele. Similarly, increasing the mutation rate, either for the whole genome or as a way to allow multiple  
227 mutations to produce blindness, results in a greater likelihood of blind populations. As for higher levels of  
228 immigration, a greater chance of producing a mutation for blindness results in an increased number of  
229 chances for a population to evolve blindness.

## 230 **Effect of intermittent connections**

231 On the low end of migration, intermittent connections effectively result in a decrease in the immigration  
232 rate and “replenishment” of the blindness allele, which increases in the level of selection required to  
233 fix blindness in the population. In contrast, when immigration rates are high, disconnecting the two  
234 populations and effectively reducing the immigration rate allows populations to fix for blindness, at least  
235 until the next period of high immigration.

## 236 **Previous observations of strong selection**

237 The values we suggest here as “strong selection” are high, but not inconsistent with previous observations.  
238 Previous calculations of strong selection resulting in selective sweeps in wild populations range from  
239 0.02–0.7 (Sáez et al., 2003; Schlenke and Begun, 2004; Wootton et al., 2002; Nair et al., 2003). Estimated  
240 selection coefficients for drug resistance in *Plasmodium falciparum* were 0.1–0.7 (Wootton et al., 2002;  
241 Nair et al., 2003). These extremely high values for selection led to fixation in 20–80 generations. For a  
242 major advantageous allele, the average value of  $s$  has been estimated as 0.11 in plants and 0.13 in animals  
243 (Rieseberg and Burke, 2001; Morjan and Rieseberg, 2004). Thus, estimated selection coefficients for cave  
244 species are consistent with a selection mechanism that is stronger than previously proposed.

## 245 **Potential mechanisms of strong selection**

246 Previous mechanisms of selection proposed for cave species have primarily been weak (Darwin, 1859;  
247 Sadoglu, 1967). However, recent work has suggested that eye development imposes a high metabolic  
248 cost, particularly for juveniles (Moran et al., 2015). In a food-limited environment, like a cave, this cost  
249 could lead to the level of selection suggested by our model. Additionally, an alternative mechanism of  
250 selection exists: migration of seeing individuals, who are strongly phototactic, out of the cave (Lankester,  
251 1925; Romero, 1985). Emigration of sighted individuals functions like selection because it systematically  
252 removes  $b^+$  alleles from the cave population. Phototaxis has been observed in eyed cavefish (Espinasa  
253 and Borowsky, 2000). Migration thus imposes strong selection in the cave for blind individuals. This  
254 mechanism of strong selection would explain the observation that blindness in cave dwelling organisms  
255 evolved repeatedly. This hypothesis is consistent with all previous hypotheses of how blindness arises (i.e.  
256 random mutation or differential expression) but suggests an alternative mechanism of selection that is  
257 much stronger than previously proposed fitness advantages (i.e. reallocation of resources). Furthermore,  
258 it suggests a way of maintaining a mostly blind cave population despite interbreeding and gene flow from  
259 surface populations (Bradic et al., 2012; Avise and Selander, 1972). Interestingly, our work is consistent  
260 with work suggesting standing cryptic variation for eye size in cavefish (Rohner et al., 2013; Rohner, 2015).  
261 The primary effect of the variation being cryptic rather than *de novo* would be that the allele frequency in  
262 the surface population would be higher than expected. However, strong selection would still be required.

263 Alternatively, vibration attraction behavior, where individuals are attracted to moving objects, provides a  
264 strong advantage for some individuals by allowing them to find food (Yoshizawa et al., 2012). This behavior  
265 is observed in multiple cavefish populations (Yoshizawa et al., 2010). In contrast, this behavior would like  
266 result in predation in a sighted environment (Yoshizawa and Jeffery, 2011). Surface populations show a  
267 low frequency of this behavior (Yoshizawa and Jeffery, 2011). However, in this case the genetic basis of the  
268 behavior would have to be linked to blindness, and whether this is the case is unknown.

## 269 **Application to other local adaptation scenarios**

270 Although we have described our model in the context of cave and surface populations, with the allele  
271 under selection for blindness, this work also applies to other scenarios of local adaptation. Generally, our  
272 scenario can be considered a metapopulation with divergent selection (Blanquart et al., 2012; Yeaman  
273 and Otto, 2011). However, while previous work focused on stable polymorphism, here we have addressed  
274 how populations become fixed for a state.

## 275 **Acknowledgments**

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## 278 **Appendix**

279 **Lemma 1.** *If  $m > 0$  or  $u > 0$ , and  $s \geq 0$ , the minimum value of an equilibrium is  $\frac{m\bar{q}(1-u)+u}{m(1-u)+u}$ . If  $m = u = 0$   
280 and  $s > 0$ , the minimum value of an equilibrium is 0. If  $m = u = s = 0$ , all points are equilibria, and thus 0  
281 is the lowest possible valid equilibrium.*

282 *Proof.* Case 1. Let  $f(q)$  represent the change in allele frequency over one generation (Equation 2). Let  
283  $q^* = \frac{m\bar{q}(1-u)+u}{m(1-u)+u}$ . If  $s = 0$  and  $m > 0$  (or  $u > 0$ ),  $f(q^*) = 0$ , and therefore  $q^*$  is an equilibrium for these

284 parameters.

285 Now let  $s \geq 0$ , and  $q < q^*$ . The denominator of  $f(q)$  is positive, and the numerator is

$$sq^2 [1 - q - m(1 - u)(1 - \tilde{q})] + [\tilde{q}m(1 - u) + u - q(m(1 - u) + u)] \quad (10)$$

$$q < \frac{m\tilde{q}(1 - u) + u}{m(1 - u) + u} \implies q < 1 - m(1 - u)(1 - \tilde{q})$$

$$\iff 1 - q - m(1 - u)(1 - \tilde{q}) > 0$$

$\iff$  the first part of (10) is positive

$$q < \frac{m\tilde{q}(1 - u) + u}{m(1 - u) + u} \iff q(m(1 - u) + u) < \tilde{q}m(1 - u) + u$$

$$\iff \tilde{q}m(1 - u) + u - q(m(1 - u) + u) > 0$$

$\iff$  the second part of (10) is positive

$\therefore$  (10) is positive, and  $f(q) > 0 \forall q < q^*$

286 Since  $f(q) > 0 \forall q < q^*$ ,  $q^*$  is the lowest value that can be an equilibrium.

287 Case 2. Let  $m + u = 0$  and  $s > 0$ . Now (10) is  $sq^2(1 - q)$ , which has its lowest equilibrium at 0.

288 Case 3. If  $s = 0$ , all possible values of  $q$  are equilibria. □

289 **Lemma 2.** *The maximum value of an equilibrium is  $1 - m(1 - u)(1 - \tilde{q})$ .*

290 *Proof.* Let  $q^* = 1 - m(1 - u)(1 - \tilde{q})$ . Since  $\lim_{s \rightarrow \infty} f(q^*) = 0$ ,  $q^*$  is a potential equilibrium. Now let

291  $q > q^*$ .

$$\begin{aligned}
 q > 1 - m(1 - u)(1 - \tilde{q}) &\Leftrightarrow 1 - q - m(1 - u)(1 - \tilde{q}) < 0 \\
 &\Leftrightarrow \text{the first part of (10) is negative} \\
 q > 1 - m(1 - u)(1 - \tilde{q}) &\Leftrightarrow q > 1 - m(1 - u) + \tilde{q}m(1 - u) \\
 &\Leftrightarrow q - 1 + m(1 - u) + u > \tilde{q}m(1 - u) + u \\
 &\Leftrightarrow q(m(1 - u) + u) - (1 - q)(1 - m(1 - u) - u) > \tilde{q}m(1 - u) + u \\
 &\Rightarrow q(m(1 - u) + u) > \tilde{q}m(1 - u) + u \\
 &\Leftrightarrow \text{the second part of (10) is negative} \\
 &\therefore (10) \text{ is negative, and } f(q) < 0 \forall q > q^*
 \end{aligned}$$

292 Since  $f(q) < 0 \forall q > q^*$ ,  $q^*$  is the highest value that can be an equilibrium. □

293 **Lemma 3.** If  $m \leq \frac{s+4u}{(2s(1-\tilde{q})+4-8\tilde{q})(1-u)}$  and  $\tilde{q} < 1/2$ , there exists an equilibrium  $\geq 1/2$ .

294 *Proof.* By rearranging Equation 3, we find a formula for the migration rate that will generate an equilibrium  
 295 at  $q$ :

$$\hat{m}(q) = \frac{(1 - q)(q^2s + u)}{(q^2s(1 - \tilde{q}) + q - \tilde{q})(1 - u)} \tag{11}$$

296 First, we can show that  $\hat{m}(q)$  monotonically decreases for  $q \geq 1/2$ :

$$-\frac{d\hat{m}(q)}{dq} = sq \left( 2\tilde{q} + 2(1 - \tilde{q})u - (1 + 3\tilde{q} + (1 - \tilde{q})u)q + 2q^2 + s(1 - \tilde{q})q^3 \right) + (1 - \tilde{q})u$$

And

$$\begin{aligned}
 0 \leq \tilde{q} \leq 1/2 \leq q \leq 1 &\Rightarrow (2q - 1)(q - \tilde{q}) + \tilde{q}(1 - q) + (2 - q)(1 - \tilde{q})u \geq 0 \\
 &\Leftrightarrow 2\tilde{q} + 2(1 - \tilde{q})u - (1 + 3\tilde{q} + (1 - \tilde{q})u)q + 2q^2 \geq 0 \\
 &\Rightarrow sq \left( 2\tilde{q} + 2(1 - \tilde{q})u - (1 + 3\tilde{q} + (1 - \tilde{q})u)q + 2q^2 + s(1 - \tilde{q})q^3 \right) \\
 &\quad + (1 - \tilde{q})u \geq 0 \\
 &\Leftrightarrow \frac{d\hat{m}(q)}{dq} \leq 0
 \end{aligned}$$

297 Therefore,  $\hat{m}(q)$  monotonically decreases from  $\hat{m}(1/2)$  to  $\hat{m}(1) = 0$  as  $q$  increases from  $1/2$  to  $1$ . Stated  
298 another way, if  $0 \leq m \leq \hat{m}(1/2)$  there exists an equilibrium  $1/2 \leq q \leq 1$ .  $\square$

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392 **Figures**

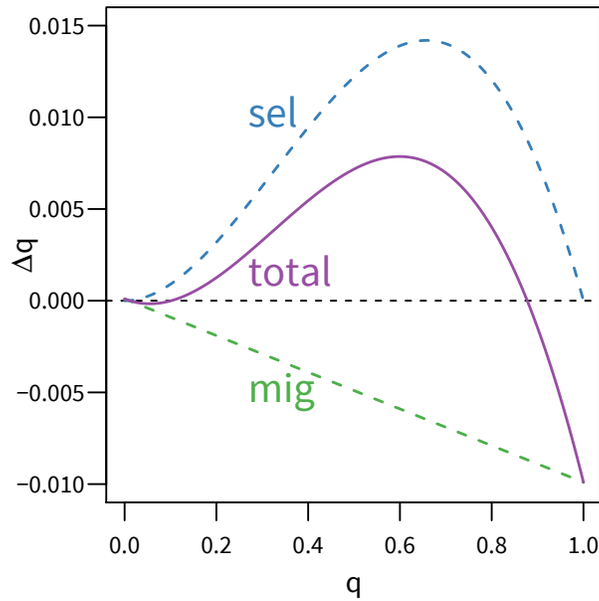


Figure 1: Dynamics of cave populations are governed by migration-selection balance. The top curve shows the change in allele frequency due to selection, while the bottom curve shows the change in allele frequency due to immigration. The middle curve shows how selection and migration are balanced to produced three equilibria points. Parameters:  $s = 0.1$ ,  $m = 0.01$ ,  $u = 10^{-6}$ ,  $\tilde{q} = 0.01$ .

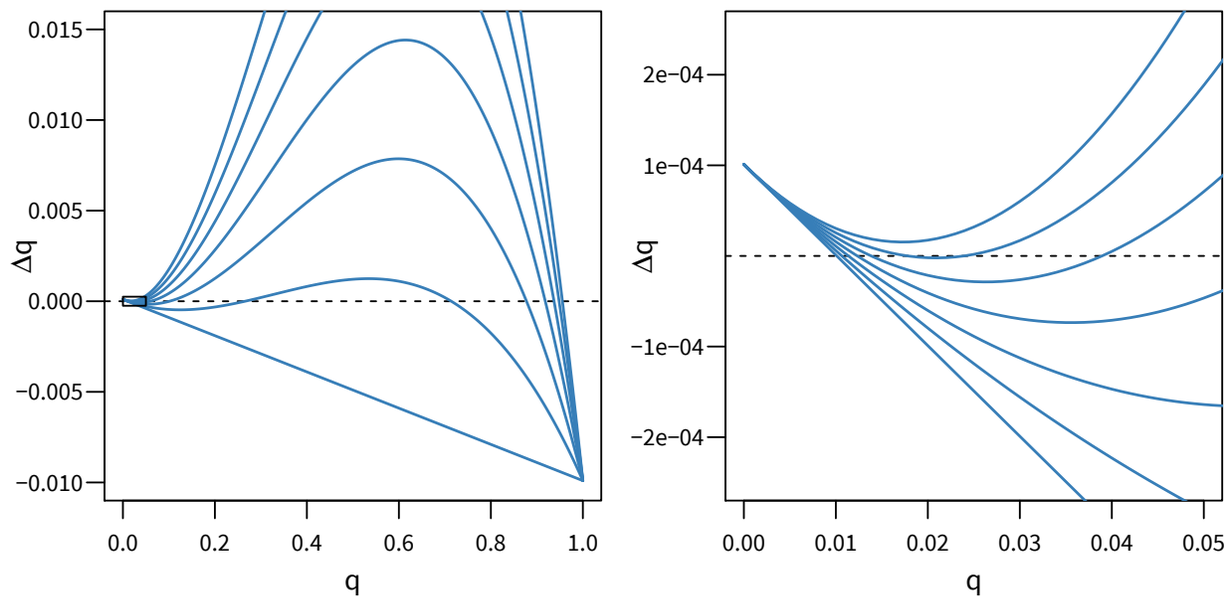


Figure 2: As selection increases, the evolutionary dynamics of the cave population changes. When  $s$  is low (bottom line;  $s = 0$ ), there is only one equilibrium: near 0. As  $s$  increases (middle five lines,  $s = 0.05, 0.1, 0.15, 0.2$ , and  $0.25$ ) the local maximum (upper hump) increases and crosses the x-axis, producing three equilibria. When  $s$  gets high enough (top line;  $s = 0.3$ ), the local minimum (lower valley) also crosses the x-axis, resulting in one equilibrium again. For all curves  $m = 0.01$ ,  $u = 10^{-6}$ , and  $\tilde{q} = 0.01$ . The figure on the right is an enlarged view of the box in the figure on the left.

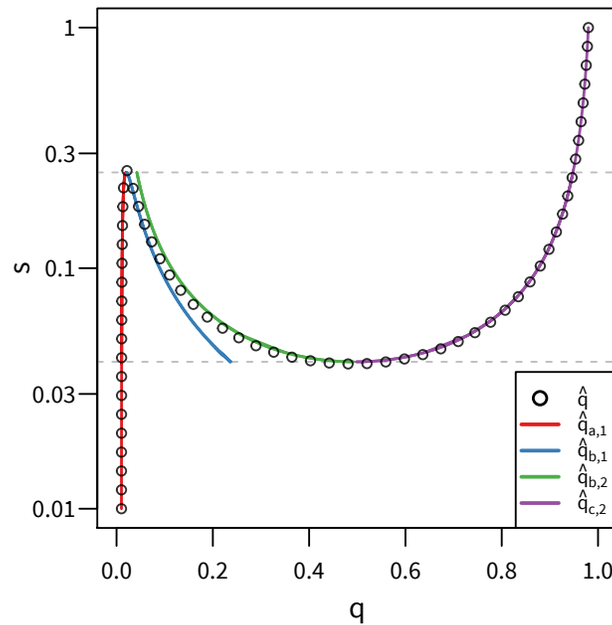


Figure 3: Our equilibria approximations are accurate. The approximations developed in this paper (solid lines) are a good fit for calculated values of  $s$  that result in equilibrium for a given  $q$  (circles) using Equation 3. Because these approximations are accurate we can use them to better understand this system. The dashed lines are our approximate bounds for the existence of three equilibria (i.e. for small and large values of  $s$ , there is one value of  $q$ ; for intermediate values of  $s$  there are three possible values of  $q$ ). Other parameters are  $m = 0.01$ ,  $u = 10^{-6}$ , and  $\tilde{q} = 0.01$ .

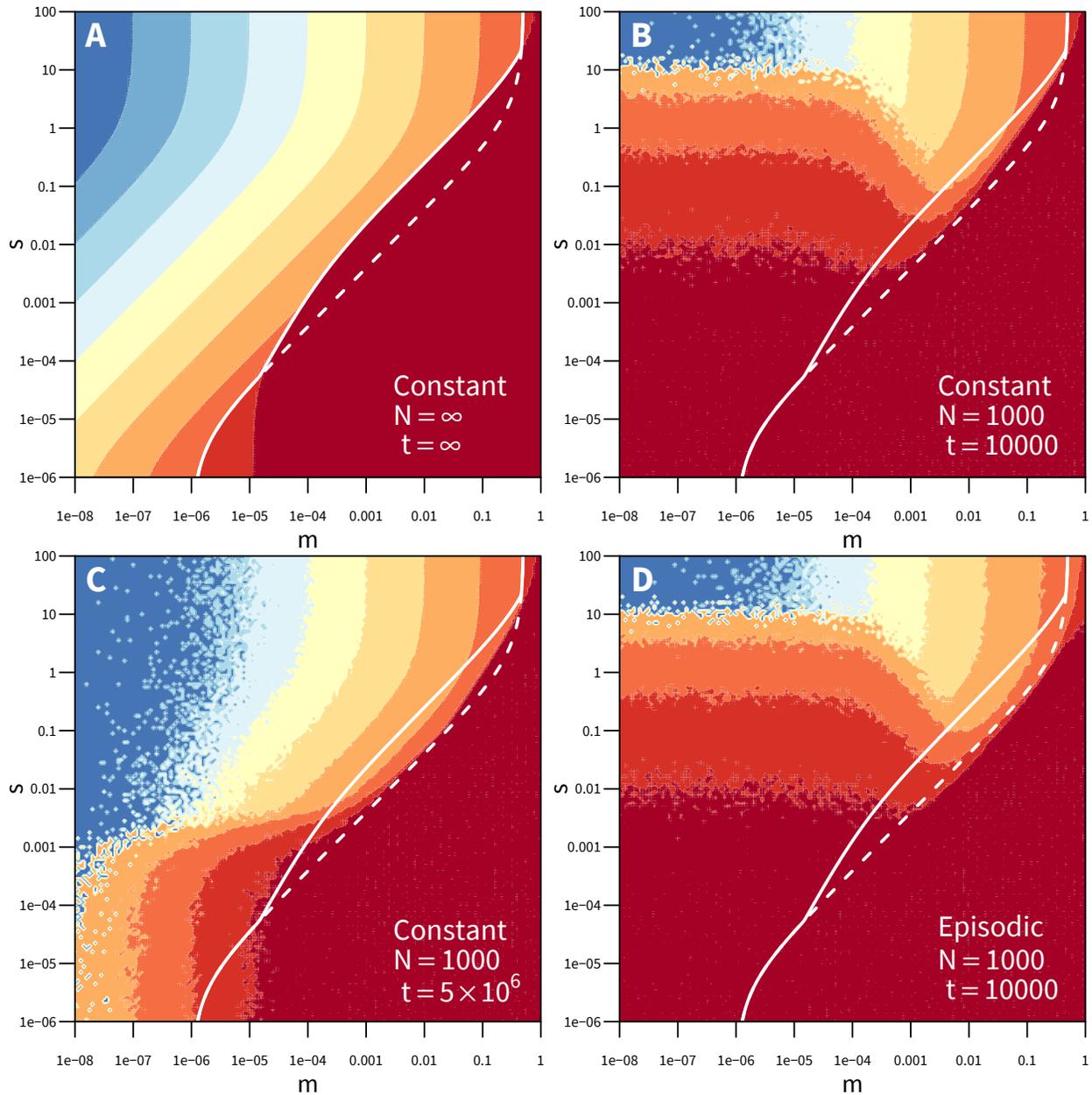


Figure 4: Populations evolve blindness with the help of strong selection. (A) The equilibrium  $b^-$  frequency for an infinite population and (B–D) average frequencies after  $t$  generations in finite populations (100 replicates). Colors correspond to the frequency of  $b^-$  for a given combination of  $s$  and  $m$ , where blue is high (blindness evolved) and red is low (blindness did not evolve). Colors levels are determined by a base-10 logit scale. The solid white line corresponds to the threshold calculated in Equation 9, and the dash white line corresponds to  $\hat{m}(1/2)$  (Equation 11). The area between the solid and dashed lines corresponds to the region where three equilibria exist. Other parameters are  $u = 10^{-6}$ ,  $\tilde{q} = 0.01$ , and  $q_0 = \tilde{q}$ .

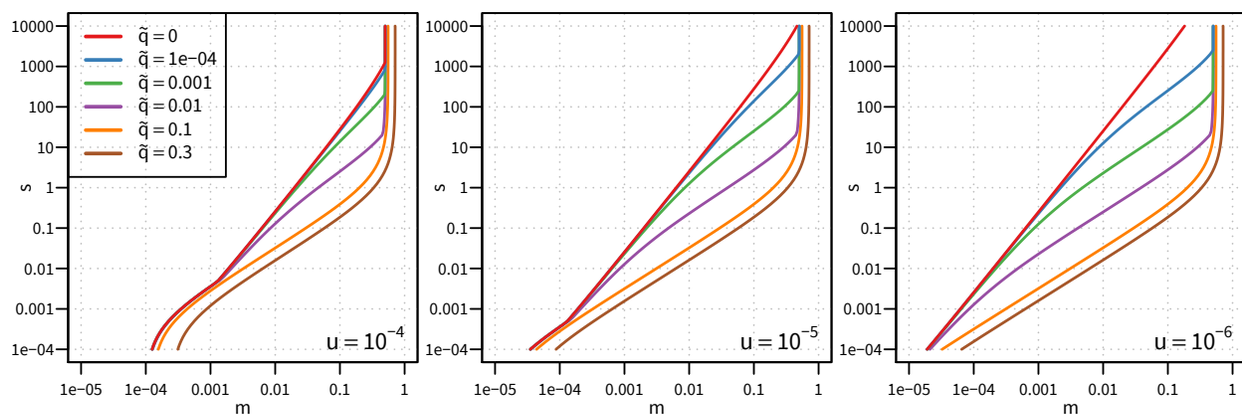


Figure 5: Strong selection is necessary for the evolution of blindness. Each panel represents a different mutation rate, and each line represents a threshold for the evolution of blindness based on  $\tilde{q}$  and  $q_0 = \tilde{q}$ . The rate of mutation of  $b^+$  to  $b^-$  alleles is a significant factor in the establishment of the thresholds, e.g. if  $\tilde{q} = 0.01$  and  $m = 0.01$ , then  $s > 0.129$  for  $u = 10^{-4}$  and  $s > 0.250$  for  $u = 10^{-6}$ ; if  $m = 0.001$ , then  $s > 0.004$  for  $u = 10^{-4}$  and  $s > 0.023$  for  $u = 10^{-6}$ .

393 **Tables**

Table 1: Evolutionary Dynamics of Blindness as  $t \rightarrow \infty$

**One Equilibrium**

$$0 \leq q_0 < \hat{q} \quad q_t \text{ increases to } \hat{q}$$

$$\hat{q} < q_0 \leq 1 \quad q_t \text{ decreases to } \hat{q}$$

**Three Equilibria**

$$0 \leq q_0 < \hat{q}_a \quad q_t \text{ increases to } \hat{q}_a$$

$$\hat{q}_a < q_0 < \hat{q}_b \quad q_t \text{ decreases to } \hat{q}_a$$

$$\hat{q}_b < q_0 < \hat{q}_c \quad q_t \text{ increases to } \hat{q}_c$$

$$\hat{q}_c < q_0 \leq 1 \quad q_t \text{ decreases to } \hat{q}_c$$