# **Analytics**

## Deriving the model dynamics

## Stationary population sizes of a wild-type only population

## In old - habitat patches

In old-habitat patches, we assume that the wild-type local populations are at carrying capacity  $K_{\text{old}}$ 

In[@]:= Nhatold = Kold;

#### In new - habitat patches

The stationary population size is found recursively, by considering the different events in the life cycle, and assuming here that carrying capacity is not reached, so that there is no need for density regulation.

Note: for simplicity, we write f for  $f_{old}$ 

$$\begin{aligned} & \text{In[*]:= Solve} \left[ x \ == \ \left( 1 - m + m \ \frac{\left( 1 - f \right)}{1 - f + \ e^{\pi w} \ f} \right) \omega w_{\text{new}} \ x + m \ \frac{\left( 1 - f \right)}{1 - f + \ e^{\pi w} \ f} \ \frac{f}{1 - f} \ \omega w_{\text{new}} \ K_{\text{old}}, \ x \right] \ / / \\ & \text{FullSimplify} \\ & \text{solNhatnew} \ = \ *[1]; \\ & \text{Out[*]=} \ \left\{ \left\{ x \rightarrow \frac{f \ K_{\text{old}} \ \omega w_{\text{new}}}{1 + \left( -1 + e^{\pi w} \right) \ f + \left( -1 + f + e^{\pi w} \ f \ \left( -1 + m \right) \ \right) \ \omega w_{\text{new}}} \right\} \right\} \end{aligned}$$

By definition, this population size cannot exceed  $K_{new}$ :

In[\*]:= Nhatnew = Min[K<sub>new</sub>, x /. solNhatnew];

## Population sizes after dispersal, wild - type only population

## In old - habitat patches

$$\label{eq:local_$$

#### In new - habitat patches

$$\begin{split} & \text{Ntildenew =} \\ & \left(1-\text{m} + \text{m} \frac{\left(1-f\right)}{1-\text{f} + \text{e}^{\pi \text{W}} \text{ f}}\right) \text{Nhatnew} + \text{m} \frac{\left(1-f\right)}{1-\text{f} + \text{e}^{\pi \text{W}} \text{ f}} \frac{\text{f}}{1-\text{f}} \text{Nhatold // FullSimplify} \\ & - \left(-1+\text{f} + \text{e}^{\pi \text{W}} \text{ f} \left(-1+\text{m}\right)\right) \text{Min} \left[\text{K}_{\text{new}}, \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{W}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f} + \left(-1+\text{m}\right) \, \omega \text{W}_{\text{new}}}\right] + \text{f} \text{m} \text{K}_{\text{old}} \\ & - \left(-1+\text{f} + \text{e}^{\pi \text{W}} \text{ f} \left(-1+\text{m}\right)\right) \, \omega \text{W}_{\text{new}}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{K}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{M}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{W}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{M}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{M}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{M}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{M}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{M}_{\text{old}} \, \omega \text{M}_{\text{old}} \, \omega \text{M}_{\text{new}}}{1+\left(-1+\text{e}^{\pi \text{M}}\right) \, \text{f}} + \frac{\text{f} \text{m} \text{M}_{\text{old}} \, \omega \text{M}_{\text{old}}$$

## Per capita growth rate of mutants

## In old - habitat patches

$$In[\bullet]:= a_{old} = \frac{\omega m_{old} K_{old}}{\omega w_{old} Ntildeold} - 1;$$

## In new - habitat patches

Since we assume that there is no density regulation in new - habitat patches because population size is below carrying capacity,

$$ln[\bullet] := a_{new} = \omega m_{new} - 1;$$

## Establishment probability

## System to be numerically solved

## Approximation of the establishment probability

## Definition of the Reproduction matrix

To avoid replacing  $a_{old}$  and  $a_{new}$  by their expressions, we write them differently, as  $s_{old}$  and  $s_{new}$ The matrix contain the expected number of successful offspring over the whole life cycle (including dispersal), i.e. the  $\lambda$  terms defined in the main text.

To avoid too lengthy expressions,

we use  $\mu_{\text{new}}$  and  $\mu_{\text{old}}$  for the probabilites that a disperser goes to a new or old patch

```
/// // reproMatrix =
                         \{\{(1-m\;\mu_{\text{new}})\;(1+s_{\text{old}})\;,\;m\;\mu_{\text{new}}\;(1+s_{\text{new}})\}\;,\;\{m\;\mu_{\text{old}}\;(1+s_{\text{old}})\;,\;(1-m\;\mu_{\text{old}})\;(1+s_{\text{new}})\}\}\;;
               reproMatrix // MatrixForm
                   \left( \begin{array}{ccc} (\mathbf{1} + \mathbf{S}_{\mathrm{old}}) \ (\mathbf{1} - \mathbf{m} \ \mu_{\mathrm{new}}) & & \mathbf{m} \ (\mathbf{1} + \mathbf{S}_{\mathrm{new}}) \ \mu_{\mathrm{new}} \\ & \mathbf{m} \ (\mathbf{1} + \mathbf{S}_{\mathrm{old}}) \ \mu_{\mathrm{old}} & & (\mathbf{1} + \mathbf{S}_{\mathrm{new}}) \ (\mathbf{1} - \mathbf{m} \ \mu_{\mathrm{old}}) \end{array} \right)
```

## Largest eigenvalue of the reproduction matrix and Taylor - expansion

We search for the largest eigenvalue of this reproduction matrix (the denominator is positive, so we pick the one with a + in front of the square root)

Info]:= Eigenvalues[reproMatrix]  $\rho = %[2] // FullSimplify;$  $Out[*] = \left\{ \frac{1}{2} \left( 2 + s_{\text{new}} + s_{\text{old}} - m \mu_{\text{new}} - m s_{\text{old}} \mu_{\text{new}} - m \mu_{\text{old}} - m \mu_{\text{old}} \right\} \right\}$  $\text{m s}_{\text{new}} \mu_{\text{old}} - \sqrt{\left( \left( -2 - \text{s}_{\text{new}} - \text{s}_{\text{old}} + \text{m } \mu_{\text{new}} + \text{m } \text{s}_{\text{old}} \mu_{\text{new}} + \text{m } \mu_{\text{old}} + \text{m } \text{s}_{\text{new}} \mu_{\text{old}} \right)^2 - \left( \left( -2 - \text{s}_{\text{new}} - \text{s}_{\text{old}} + \text{m } \mu_{\text{new}} + \text{m } \text{s}_{\text{old}} \mu_{\text{new}} + \text{m } \mu_{\text{old}} + \text{m } \text{s}_{\text{new}} \mu_{\text{old}} \right)^2 - \left( \left( -2 - \text{s}_{\text{new}} - \text{s}_{\text{old}} + \text{m } \mu_{\text{new}} + \text{m } \text{s}_{\text{old}} \mu_{\text{new}} + \text{m } \mu_{\text{old}} + \text{m } \text{s}_{\text{new}} \mu_{\text{old}} \right)^2 \right)}$ 4 (1 +  $s_{new}$  +  $s_{old}$  +  $s_{new}$   $s_{old}$  - m  $\mu_{new}$  - m  $s_{new}$   $\mu_{new}$  - m  $s_{old}$   $\mu_{new}$   $exttt{m } exttt{S}_{ exttt{new}} exttt{S}_{ exttt{old}} - exttt{m } exttt{S}_{ exttt{new}} - exttt{m } exttt{M}_{ exttt{old}} - exttt{m } exttt{S}_{ exttt{new}} + exttt{M}_{ exttt{old}} + exttt{M}_{ exttt{o$  $\frac{1}{2} \left( 2 + \mathsf{S}_{\mathsf{new}} + \mathsf{S}_{\mathsf{old}} - \mathsf{m} \ \mu_{\mathsf{new}} - \mathsf{m} \ \mathsf{S}_{\mathsf{old}} \ \mu_{\mathsf{new}} - \mathsf{m} \ \mu_{\mathsf{old}} - \mathsf{m} \ \mathsf{S}_{\mathsf{new}} \ \mu_{\mathsf{old}} + \right)$  $\sqrt{\left(-2-s_{\text{new}}-s_{\text{old}}+m\,\mu_{\text{new}}+m\,s_{\text{old}}\,\mu_{\text{new}}+m\,\mu_{\text{old}}+m\,s_{\text{new}}\,\mu_{\text{old}}\right)^2}$ 4 (1 + S<sub>new</sub> + S<sub>old</sub> + S<sub>new</sub> S<sub>old</sub> - m  $\mu$ <sub>new</sub> - m S<sub>new</sub>  $\mu$ <sub>new</sub> - m S<sub>old</sub>  $\mu$ <sub>new</sub> -

Define assumptions on the parameters to help Mathematical simplify solutions (note that both €>0 or €<0 work. But one needs to assume one for Mathematica to solve the square roots)

```
\ln[\bullet]:= assumptions = \{m > 0 \&\& m < 1 \&\& \pi_m \in \text{Reals \&\& } \}
                  \pi_{w} \in \text{Reals \&\& } \epsilon > 0 \&\& s_{\text{new}} > 0 \&\& \xi > 0 \&\& f > 0 \&\& f < 1 \&\& s_{\text{new}} > s_{\text{old}} \};
```

Define scaling to rescale parameters as function of  $\epsilon$ , assumed to be small

$$ln[\bullet]:=$$
 scaling = {m  $\rightarrow \epsilon$  mm,  $s_{old} \rightarrow \epsilon$  ss<sub>old</sub>,  $s_{new} \rightarrow \epsilon$  ss<sub>new</sub>};

Taylor - expand the eigenvalue, at the first order in  $\epsilon$  (after rescaling), and simplify

 $m_{\rm col} = 1$  Assuming[assumptions, Series[ $\rho$  /. scaling, { $\epsilon$ , 0, 1}] // Normal // FullSimplify]

$$\begin{aligned} \textit{Out[*]} &= 1 + \frac{1}{2} \in \left( \texttt{ss}_{\texttt{new}} + \texttt{ss}_{\texttt{old}} - \texttt{mm} \; \left( \mu_{\texttt{new}} + \mu_{\texttt{old}} \right) \; + \\ & \sqrt{ \left( \texttt{ss}_{\texttt{new}} - \texttt{ss}_{\texttt{old}} + \texttt{mm} \; \mu_{\texttt{new}} \right)^2 + 2 \; \texttt{mm} \; \left( - \; \texttt{ss}_{\texttt{new}} + \; \texttt{ss}_{\texttt{old}} + \; \texttt{mm} \; \mu_{\texttt{new}} \right) \; \mu_{\texttt{old}} + \; \texttt{mm}^2 \; \mu_{\texttt{old}}^2 } \right) \end{aligned}$$

#### Eigenvectors

Left eigenvector associated to the largest eigenvalue (second one in the order of vectors in the output)

```
In[⊕]:= u = Eigenvectors[Transpose[reproMatrix]] [2] // FullSimplify;
    Right eigenvector
In[*]:= v = Eigenvectors[reproMatrix][2] // FullSimplify;
```

Find the coefficients satisfying the following constraints (lev = left eigenvector, rev = right eigenvector, the numbers refer to the elements)

$$lole = Solve[{a (lev1 + lev2) == 1, a lev1 b rev1 + a lev2 b rev2 == 1}, {a, b}]$$

$$\textit{Out[*]=} \ \Big\{ \Big\{ a \rightarrow \frac{1}{\text{lev1} + \text{lev2}} \, , \, \, b \rightarrow \frac{\text{lev1} + \text{lev2}}{\text{lev1} \, \text{rev1} + \text{lev2} \, \text{rev2}} \Big\} \Big\}$$

a is the sum of the elements of the left eigenvector, b's denominator is the product of the vectors. The normalized vectors (n) are therefore

## Approximate establishment probability

Directly applying Theorem 5.6 from Haccou et al. (2005)

We define

$$\ln[\textbf{p}] = \mathbf{B} = \sum_{i=1}^{2} \left( \mathbf{un[i]} \left( \sum_{j=1}^{2} \mathbf{vn[j]} \ \mathbf{reproMatrix[i,j]} \right) \right) + \rho \ (\mathbf{1} - \rho) \left( \sum_{j=1}^{2} \mathbf{un[j]} \ \mathbf{vn[j]}^{2} \right);$$

and according to the theorem, the approximate establishment probabilities are then

$$ln[\circ] = exact \phi = \frac{2 (\rho - 1) vn}{B};$$

which we Taylor - Expand as (this takes a few seconds/minutes)

In[\*]:= solutions = Assuming[assumptions,

Assuming[assumptions, Normal[Series[exact $\phi$ /.scaling, { $\epsilon$ , 0, 1}]]]];

and we recover the original parameters (this also takes a few seconds/minutes)

$$log(s):=$$
 backscaling = {mm  $\rightarrow$  m /  $\epsilon$ ,  $ss_{old} \rightarrow s_{old} / \epsilon$ ,  $ss_{new} \rightarrow s_{new} / \epsilon$ };

backsolutions = Assuming[assumptions, FullSimplify[solutions /. backscaling]]

Extract the two establishment probabilities, and recover the model parameters

$$\lim_{\|x\| = 1} \text{recoverparams} = \left\{ \mu_{\text{new}} \to \frac{\left(1 - f\right)}{1 - f + e^{\pi m} f}, \mu_{\text{old}} \to \frac{e^{\pi m} f}{1 - f + e^{\pi m} f}, s_{\text{old}} \to a_{\text{old}}, s_{\text{new}} \to a_{\text{new}} \right\};$$

# **Figures**

## **Initializations**

#### **Parameters**

```
constantParam = {K_{old} \rightarrow 1000, K_{new} \rightarrow 500, \omega w_{old} \rightarrow 1.5, \omega w_{new} \rightarrow 0.75, \omega m_{new} \rightarrow 1.02};
       The other default parameters are
      f = 0.5,
       m = 0.06,
       \omegam<sub>old</sub> = 1.45 or 1.35
In[@]:= mdefault = 0.06;
       fdefault = 0.5;
       \omegamdefault1<sub>old</sub> = 1.35;
       \omegamdefault2<sub>old</sub> = 1.45;
       \pi\theta = \{\pi w \rightarrow \theta, \pi m \rightarrow \theta\};
       \pi 00 = \{\pi w \to 1/2, \pi m \to 1/2\};
       \pi ON = \{\pi w \to 1/2, \pi m \to -1/2\};
       \pi NO = \{\pi w \rightarrow -1/2, \pi m \rightarrow 1/2\};
       \pi NN = \{\pi w \rightarrow -1/2, \pi m \rightarrow -1/2\};
       Emigration probability m
ln[@]:= mmin = 10^{-3}; (* minimum value *)
       mmax = 1; (* max value *)
       nm = 101; (* number of values *)
       logmvals = Table [Log10[mmin] + \frac{\left(\text{Log10[mmax]} - \text{Log10[mmin]}\right)\left(i-1\right)}{\text{nm}-1}, {i, 1, nm}];
       mvals = Table[10<sup>logmvals[i]</sup>, {i, 1, nm}];
```

## Plotting parameters

```
In[*]:= col0 = Black;
    col00 = Blue;
    colON = Orange;
    colNO = Red;
    colNN = Green;
ln[•]:= mark0 = ●;
    mark00 = ▲;
    markON = ■;
    markNO = ◆;
    markNN = ▼;
```

## Population variables

Nhatold and Nhatnew, the population densities of the wild-type population at the beginning of a generation

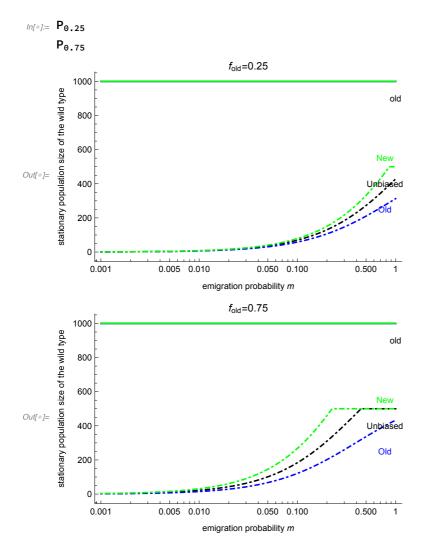
## Only for unbiaised dispersal

```
In[*]:= Show[LogLinearPlot[
        {Nhatold /. constantParam, Nhatnew /. constantParam /. {f \rightarrow fdefault} /. \pi0},
        {m, mmin, mmax}, PlotStyle → {{col0, Full}, {col0, DotDashed}}},
       AxesOrigin → {mmin, 0}, Frame → {True, True, False, False},
        FrameLabel \rightarrow {"emigration probability m",
          "stationary population size of the wild type"}],
      Graphics[Text["old", {0, 900}]], Graphics[Text["new", {-0.25, 400}]]]
     stationary population size of the wild type
                                                               old
        800
        600
        400
        200
                      0.005 0.010
                                       0.050 0.100
                                                         0.500
          0.001
                              emigration probability m
```

## For all WT dispersal biases

(The dispersal bias of the mutant does not affect the population size of the wild - type population)

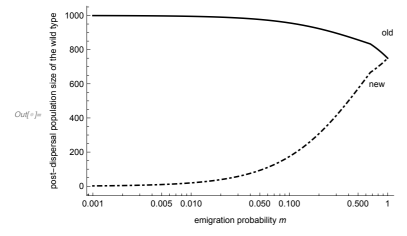
```
Infolia Show[LogLinearPlot[
       {Nhatold /. constantParam, Nhatnew /. constantParam /. {f \rightarrow fdefault} /. \pi 0},
       {m, mmin, mmax}, PlotStyle → {{col0, Full}, {col0, DotDashed}}},
       AxesOrigin → {mmin, 0}, Frame → {True, True, False, False},
       FrameLabel \rightarrow {"emigration probability m",
          "stationary population size of the wild type"}],
      LogLinearPlot[{Nhatold /. constantParam,
        Nhatnew /. constantParam /. {f \rightarrow fdefault} /. \pi00},
       {m, mmin, mmax}, PlotStyle → {{col00, Full}, {col00, DotDashed}}],
      LogLinearPlot[{Nhatold /. constantParam,
        Nhatnew /. constantParam /. {f \rightarrow fdefault} /. \pi NN},
       {m, mmin, mmax}, PlotStyle → {{colNN, Full}, {colNN, DotDashed}}],
      Graphics[Text["old", {0, 900}]],
      Graphics[Text[Style["Old", col00], {-0.25, 250}]],
      Graphics[Text[Style["New", colNN], {-0.25, 550}]],
     Graphics[Text[Style["Unbiased", Small], {-0.25, 400}]]]
       1000
    stationary population size of the wild type
                                                         old
       800
       600
       400
       200
         0
         0.001
                    0.005 0.010
                                    0.050 0.100
                                                   0.500
                           emigration probability m
    With other values of f_{old}
In[@]:= Do[Pthef = Show[LogLinearPlot[
          {Nhatold /. constantParam, Nhatnew /. constantParam /. {f \rightarrow thef} /. \pi 0},
         {m, mmin, mmax}, PlotStyle → {{col0, Full}, {col0, DotDashed}}},
         AxesOrigin → {mmin, 0}, Frame → {True, True, False, False},
         FrameLabel \rightarrow {"emigration probability m",
            "stationary population size of the wild type"}],
        LogLinearPlot[{Nhatold /. constantParam,
           Nhatnew /. constantParam /. {f \rightarrow thef} /. \pi00},
         {m, mmin, mmax}, PlotStyle → {{col00, Full}, {col00, DotDashed}}],
        LogLinearPlot[{Nhatold /. constantParam,
           Nhatnew /. constantParam /. \{f \rightarrow thef\} /. \pi NN\}, \{m, mmin, mmax\},
         PlotStyle → {{colNN, Full}, {colNN, DotDashed}}], Graphics[
         Text["old", {0, 900}]], Graphics[Text[Style["Old", col00], {-0.25, 250}]],
        Graphics[Text[Style["New", colNN], {-0.25, 550}]],
        Graphics[Text[Style["Unbiased", Small], {-0.25, 400}]],
        PlotLabel → "f<sub>old</sub>=" <> ToString[thef]]
      , {thef, {0.25, 0.75}}]
```



## Ntildeold and Ntildenew, the population densities of the wild - type population right after dispersal

## Only for unbiaised dispersal

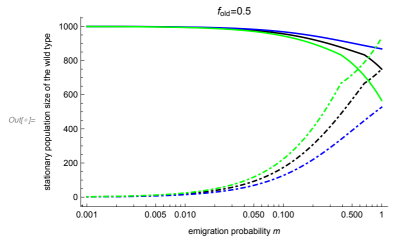
```
ln[\cdot]:= Show[LogLinearPlot[{Ntildeold /. constantParam /. {f \rightarrow fdefault} /. \pi0,
        Ntildenew /. constantParam /. \{f \rightarrow fdefault\} /. \pi 0\}, \{m, mmin, mmax\},
       PlotStyle → {{col0, Full}, {col0, DotDashed}}, AxesOrigin → {mmin, 0},
       Frame \rightarrow {True, True, False, False}, FrameLabel \rightarrow {"emigration probability m",
         "post-dispersal population size of the wild type"}],
     Graphics[Text["old", {0, 900}]], Graphics[Text["new", {-0.25, 600}]]]
```



## For all WT dispersal biases

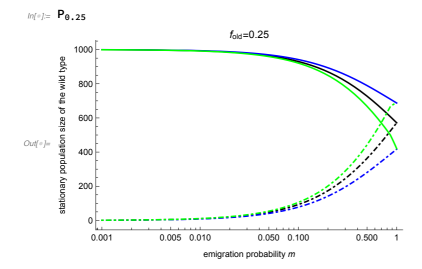
Again, dispersal bias of the mutant has no influence

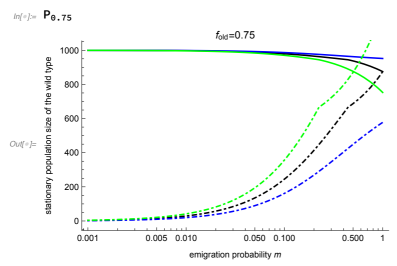
```
\ln |\pi| = \text{Show}[\text{LogLinearPlot}[\text{Ntildeold /. constantParam /. } \{f \rightarrow \text{fdefault}] /. \pi 0,
         Ntildenew /. constantParam /. {f \rightarrow fdefault} /. \pi0}, {m, mmin, mmax},
       PlotStyle → {{col0, Full}, {col0, DotDashed}}, AxesOrigin → {mmin, 0},
       Frame \rightarrow {True, True, False, False}, FrameLabel \rightarrow {"emigration probability m",
          "stationary population size of the wild type"}],
      LogLinearPlot[{Ntildeold /. constantParam /. {f \rightarrow fdefault} /. \pi 00,
         Ntildenew /. constantParam /. \{f \rightarrow fdefault\} /. \pi00\},
       {m, mmin, mmax}, PlotStyle → {{col00, Full}, {col00, DotDashed}}],
      LogLinearPlot[{Ntildeold /. constantParam /. {f \rightarrow fdefault} /. \pi NN,
         Ntildenew /. constantParam /. {f \rightarrow fdefault} /. \pi NN},
       {m, mmin, mmax}, PlotStyle → {{colNN, Full}, {colNN, DotDashed}}],
      PlotLabel \rightarrow "f_{old} = " \Leftrightarrow ToString[fdefault]]
```



For other values of  $f_{\text{old}}$ 

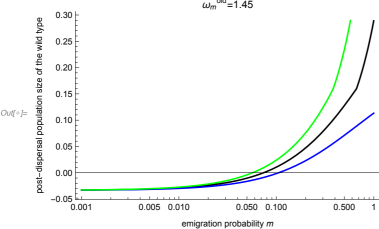
```
In[\circ]:= Do[P_{thef} =
       Show[LogLinearPlot[{Ntildeold /. constantParam /. {f \rightarrow thef} /. \pi0,
           Ntildenew /. constantParam /. \{f \rightarrow thef\} /. \pi0\},
          {m, mmin, mmax}, PlotStyle → {{col0, Full}, {col0, DotDashed}},
          AxesOrigin → {mmin, 0}, Frame → {True, True, False, False},
          FrameLabel \rightarrow {"emigration probability m",
             "stationary population size of the wild type"}],
        LogLinearPlot[{Ntildeold /. constantParam /. {f \rightarrow thef} /. \pi00,
           Ntildenew /. constantParam /. \{f \rightarrow thef\} /. \pi 00\},
          {m, mmin, mmax}, PlotStyle → {{col00, Full}, {col00, DotDashed}}],
        LogLinearPlot[{Ntildeold /. constantParam /. {f \rightarrow thef} /. \pi NN,
           Ntildenew /. constantParam /. \{f \rightarrow thef\} /. \pi NN\},
          {m, mmin, mmax}, PlotStyle → {{colNN, Full}, {colNN, DotDashed}}],
        PlotLabel → "f<sub>old</sub>=" <> ToString[thef]], {thef, {0.25, 0.75}}]
```



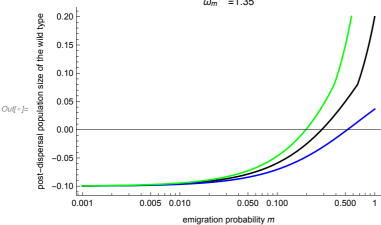


## $a_{\rm old}$ , the per capita local growth rate of the mutant

```
In[*]:= Show
      LogLinearPlot[a_{old} /. constantParam /. {f \rightarrow fdefault, \omega m_{old} \rightarrow \omega mdefault2_{old}} /. \pi 0,
        {m, mmin, mmax}, PlotStyle → {{col0, Full}, {col0, DotDashed}},
        AxesOrigin → {mmin, 0}, Frame → {True, True, False, False},
        FrameLabel \rightarrow {"emigration probability m",
           "post-dispersal population size of the wild type"}],
       LogLinearPlot[a_{old} /. constantParam /. {f \rightarrow fdefault, \omega m_{old} \rightarrow \omega mdefault2_{old} /. \pi 00,
        {m, mmin, mmax}, PlotStyle → {{col00, Full}, {col00, DotDashed}}],
       LogLinearPlot[a_{old} /. constantParam /. \{f \rightarrow fdefault, \omega m_{old} \rightarrow \omega mdefault2_{old}\} /. \pi NN,
        {m, mmin, mmax}, PlotStyle → {{colNN, Full}, {colNN, DotDashed}}],
       Graphics[Text["old", {0, 900}]], Graphics[Text["new", {-0.25, 600}]],
       PlotLabel \rightarrow "\omega_m^{\text{old}}=" <> ToString[\omegamdefault2<sub>old</sub>]]
                                    \omega_m^{\text{old}}=1.45
         0.30
```



```
In[*]:= Show[
       LogLinearPlot[a_{old} /. constantParam /. {f \rightarrow fdefault, \omega m_{old} \rightarrow \omega mdefault1_{old}} /. \pi 0,
        {m, mmin, mmax}, PlotStyle → {{col0, Full}, {col0, DotDashed}}},
        AxesOrigin → {mmin, 0}, Frame → {True, True, False, False},
        FrameLabel \rightarrow {"emigration probability m",
           "post-dispersal population size of the wild type"}],
       LogLinearPlot[a_{old} /. constantParam /. {f \rightarrow fdefault, \omega m_{old} \rightarrow \omega mdefault1_{old}} /. \pi 00,
        {m, mmin, mmax}, PlotStyle → {{col00, Full}, {col00, DotDashed}}],
       LogLinearPlot[a_{old} /. constantParam /. {f \rightarrow fdefault, \omega m_{old} \rightarrow \omega mdefault1_{old}} /. \pi NN,
        {m, mmin, mmax}, PlotStyle → {{colNN, Full}, {colNN, DotDashed}}],
       Graphics[Text["old", {0, 900}]], Graphics[Text["new", {-0.25, 600}]],
       PlotLabel \rightarrow "\omega_m^{old}=" <> ToString[\omegamdefault1<sub>old</sub>]]
                                    \omega_m^{\text{old}} = 1.35
```



# **Tests -- Sandbox**

Numerically solving the establishment probabilities

```
\textit{In[e]} := \texttt{FindRoot[Evaluate[\{Eq_{old},\ Eq_{new}\}\ /.\ constantParam\ /.\ \{f \to fdefault, fdefault, fdefault, fdefault, fdefault]}
                   \texttt{m} \rightarrow \texttt{mdefault}, \ \omega \texttt{m}_{\texttt{old}} \rightarrow \omega \texttt{mdefault1}_{\texttt{old}} \} \ \textit{/.} \ \pi \texttt{0}] \ , \ \{ \phi_{\texttt{old}}, \ \texttt{0.5} \} \ , \ \{ \phi_{\texttt{new}}, \ \texttt{0.5} \} \} ]
Out[\sigma]= {\phi_{old} \rightarrow 0.114666, \phi_{new} \rightarrow 0.0743409}
         Plotting simulation data
In[*]:= SetDirectory[
           "/Users/flo/Documents/Work/Projects/1_EnCours/2018_RescuePete/CodesPete"]
out|=]= /Users/flo/Documents/Work/Projects/1_EnCours/2018_RescuePete/CodesPete
```

```
In[*]:= aa = Import[
        "../evolutionary_rescue_and_dispersal/Fig2/a/vary_mu_phi1_pi1_0_pi2_0.csv"]
Out[v] = \{\{0.00229, 0.02\}, \{0.00135, 0.008\}, \{0.00156, 0.01\}, \{0, 0.06\}, \}
       \{0.00017, 0.001\}, \{0.00039, 0.003\}, \{0.00032, 0.0015\}, \{0, 0.15\},
       \{0.00044, 0.002\}, \{0.00067, 0.004\}, \{0, 0.08\}, \{0.00087, 0.005\},
       \{0.00098, 0.006\}, \{0, 0.1\}, \{0.00177, 0.015\}, \{0.00188, 0.03\},
       \{0.0013, 0.04\}, \{0.00019, 0.05\}, \{0.09898, 1\}, \{0.09258, 0.8\},
       \{0.00007, 0.2\}, \{0.0237, 0.3\}, \{0.04701, 0.4\}, \{0.08215, 0.6\}, \{0.06763, 0.5\}\}
log_{0} = ListLogLinearPlot[Table[{aa[[i, 2]], aa[[i, 1]]}, {i, 1, Length[aa]}]]
     0.10
     0.08
     0.06
Out[ • ]=
     0.04
     0.02
                    0.005 0.010
                                     0.050 0.100
                                                     0.500
```