LUKE HEATON - NICK JONES - MARK FRICKER

FUNGAL T2M -SOFTWARE MANUAL

PLANT SCIENCES, OXFORD FIRST EDITION

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The export_fig package was written by Oliver Woodford and Yair Altman and downloaded from: https://uk.mathworks.com/matlabcentral/fileexchange/23629-export-fig

The screen capture package was written by Yair Altman (2020) and downloaded from:.

https://www.mathworks.com/matlabcentral/fileexchange/ 24323-screencapture-get-a-screen-capture-of-a-figure-frame-or-component

The colorcet.m package was written by Peter Kovesi and downloaded from:

https://peterkovesi.com/projects/colourmaps/

Peter Kovesi. Good Colour Maps: How to Design Them. arXiv:1509.03700 [cs.GR] 2015 https://arxiv.org/abs/1509.03700

Installation

1.1 Overview

The code to run the simulation is provided as open source under a GNU General Public License v3.0 from:

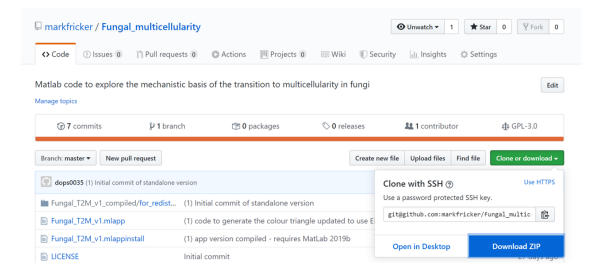
https://github.com/markfricker/Fungal_multicellularity

The code is provided in three alternative formats:

- A set of script files and functions that will run in MATLAB® on any platform;
- A MATLAB[®] app with a GUI interface that can be installed in MATLAB[®] on any platform (requires MATLAB[®] release 2019b or later, the Image Processing ToolboxTM and a screen resolution of 1600 x 900 or greater);
- A compiled standalone version that can be installed and run on Windows 10, 64 bit platforms and requires a screen resolution of 1600 x 900 or greater;

On the home page of the github repository, click on the green Clone or download button (Figure 1.1) and then the blue Download ZIP button.

Figure 1.1: Download the software in a zip file from the github repository



1.2 Download all files from the github repository

Once the file has downloaded, extract all the files. If you intend to run the script versions, the extracted folder needs to be on the MATLAB® path.

1.3 Installation of the MATLAB app

The MATLAB[®] app installer file, *Fungal_T2M_v1.mlappinstall*, contains everything necessary to install and run the App within the MATLAB[®] environment, including the source code, supporting data, information (such as product dependencies), and the app icon (Figure 1.2). The installation time is less than a minute.

Double-clicking on the *Fungal_T2M_v1.mlappinstall* file should launch MATLAB[®] if it is not already running, and install the app in the app toolbar (Figure 1.3). The program can be run, by clicking on the icon in the toolbar.



Figure 1.2: The program icon

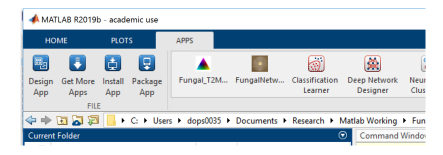


Figure 1.3: Location of the Fungal_T2M app in the APPS menu within MATLAB®

The app version of the program also requires the MATLAB® Image Processing Toolbox TM to be installed.

1.4 Installation of the stand-alone program

The software has been tested on Windows 10, and requires a minimum screen resolution of 1600 x 900. In addition, an appropriate version of the MATLAB® Compiler Runtime (MCR) is required to install the set of shared libraries that enable execution of the compiled MATLAB® application. The MCR should automatically download from the MathWorks website when the program is installed for the first time. Alternatively MCR can be downloaded from:

http://www.mathworks.com/products/compiler/mcr.

To install the MCR and standalone package, double-click the compiled MATLAB self-extracting <code>Fungal_T2M_v1.exe</code> file located in the subfolder:

\Fungal_T2M_v1_compiled\for_redistribution

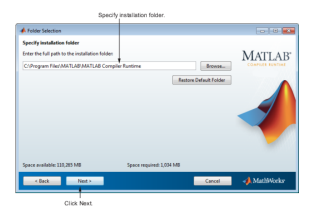
This extracts the MATLAB® Runtime Installer from the archive, along with all the files that make up the deployed MATLAB® en-

vironment. Once all the files have been extracted, the MATLAB® Runtime Installer starts automatically. When the MATLAB® Runtime Installer starts, it displays the following dialog box:



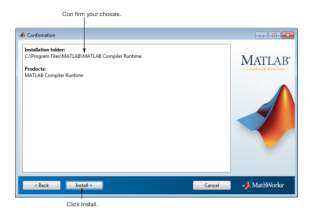
Read the information and then click Next to proceed with the installation.

Specify the folder in which you want to install the MATLAB® runtime in the Folder Selection dialog box and click Next.1 It is recommended to keep the default settings as this ensures the path to other program files is set automatically.



¹ On Windows systems, you can have multiple versions of the MATLAB® runtime on your computer, but only one installation for any particular version. If you already have an existing installation, the MATLAB® runtime Installer does not display the Folder Selection dialog box because you can only overwrite the existing installation in the same folder.

Confirm your choices and click Install. The MATLAB® Runtime Installer starts copying files into the installation folder. Installation takes about 10 minutes.



Click Finish to exit the installer.



MATLAB® Runtime Installer Readme File: A readme.txt file is included with the MATLAB Runtime Installer. This file, visible when the MATLAB Runtime Installer is expanded, provides more detailed information about the installer and the switches that can be used with it.

1.5 Installation of additional program files needed

A number of additional files needed to provide high quality output need to be installed at the same time as the main program. The latest version of Java needs to be installed, and is available from:

http://www.java.com/en/

Output of images at full resolution uses <code>export_fig.m</code> originally written by Oliver Woodford (2008-2014) and now maintained by Yair Altman (2015-). This is included in the installation. However, when exporting to vector format (PDF or EPS) this function requires that ghostscript is installed on your system. Ghostscript can be downloaded from:

http://www.ghostscript.com.

When exporting images to eps and pdf formats, <code>export_fig</code> additionally requires pdftops, from the Xpdf suite of functions. This is included in the xpdf tools package and can be downloaded from:

https://www.xpdfreader.com/download.html

Running the simulation using the GUI

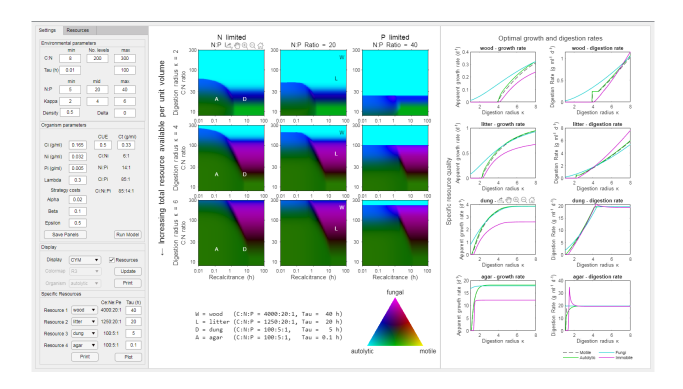


Figure 2.1: The GUI interface for the fungal multicellularity simulation

2.1 Introduction

The MATLAB app version and the standalone version both provide the same interface to run the simulation (Figure 2.1). The set of left-hand panels allow the user to change the parameters controlling the external resource environment and the internal organism parameters, including the cost terms for each of the different strategies. The definition of the parameters is given in Table 2.1.

Table 2.1: Definition and values of modelling parameters

Tunable Environmental Parameters	Symbol and Units	Interpretation
Supply of C, N and P	C_{E} , N_{E} , P_{E}	Grams of C, N and P per millilitre of substrate, set by the C:N and N:P ratio of the resource, and the resource density in g ml ⁻¹
Recalcitrance	τ hours	Time required for an exoenzyme to supply a mass of C, N or P equal to the total mass required to synthesise the exoenzyme in question
Relative digestion radius	κ	Cylindrical cells and hyphae digest resource over a distance times the cell radius
Resource accessibility	δ	Ratio of C that has to be digested to release each N, to reflect that N is embedded within C-rich polymers

Tuneable Organism Parameters	Symbol and Units	Interpretation
Core demand for C, N and P	$C_{\rm I} = 0.165 \ g \ ml^{-1}$ $N_{\rm I} = 0.032 \ g \ ml^{-1}$ $P_{\rm I} = 0.005 \ g \ ml^{-1}$	Core demand for C, N and P per unit volume of any organism.
Carbon Use Efficiency	CUE = 0.5	Ratio of C_1 for growth to total C required (C_T) , including respiration
Motility cost	$\alpha = 0.02$	Mass of C and N required to synthesise motile apparatus, relative to core demand for C and N
Vesicle transport cost	eta=0.1	Mass of C and N in vesicles, relative to the total C and N in exoenzymes and the fungal core
Recycling efficiency	$\epsilon = 0.5$	Fraction of C, N and P that is recouped by autolysis
Maximal rate of resource use	$\lambda = 0.3 \text{ g ml}^{-1} \text{ hour}^{-1}$	Maximal mass of resource any cell can use per unit time and volume

Model Variables	Symbol and Units	Interpretation
Relative density of exoen-	x	Mass of C and N used for exoenzymes for uni-
		cellular organisms relative to the core demand
zymes		for C and N in unicells
Polative density of	$x_{\rm C}, x_{\rm N}, x_{\rm P}$	Mass of C and N used for C, N or P digesting
Relative density of C, N or P exoenzymes		exoenzymes for fungal organisms relative to
C, N of F excenzymes		the core demand for C and N

Functions of x	Symbol and Units	Interpretation
Time to exhaust local resource	T hours	Time until one of C, N or P is locally exhausted
Time to exhaust C, N or P	$T_{\rm C}$, $T_{\rm N}$, $T_{\rm P}$ hours	Time until relevant element is locally exhausted
Specific growth rate of growing cell	$\eta~\mathrm{h}^{-1}$	Rate of synthesis of cell biomass, per unit of cell biomass, in the growing cells
Apparent growth rate	$\mu \ \mathrm{h}^{-1}$	Volumetric rate of colonisation, per unit volume colonised
Total limiting supply	Ω	Maximum number of daughter cells and their exoenzymes that can be synthesised, given the local supply of resource
Rate of resource use	$\Gamma \text{ g ml}^{-1} \text{ h}^{-1}$	Total mass of resource used per unit time and volume, in the growing cells

Tuneable environmental parameters

The C:N ratio of the resource ranges from the minimum value (min), typically around 5, to the maximum value (max), typically around 300, with the number of intermediate levels set by levels (Figure 2.2). The C:N ratio forms the y-axis for each of the resultant plots (on a log_{10} scale). The x-axis is set by the recalcitrance (τ), and ranges from min, typically around 0.01, to max, typically around 100, with the same number of intermediate levels. Results are plotted for three values of the N:P ratio, set by min, mid and max (default 5, 20 and 40, respectively) and three values for the overall resource availability, given by the radius of digestion (κ), again set by min, mid and max (default 2, 4 and 6, respectively).

The overall resource density in g ml⁻¹ is used to convert the resource C:N and C:P ratios into grams ml⁻¹ of C_E , N_E , and P_E .

The value of the additional accessibility parameter δ , reflects the fact that some fraction of the available C must be digested in order to access N, irrespective of the category of organism. For example, if the C:N ratio of the substrate is 200:1, imposing a value of $\delta = 0.1$ forces organisms to digest at least 20 C for every N they acquire.

Tuneable organism parameters 2.3

The internal C, N and P required by all organisms are set by C_i , N_i and P_i , respectively, in g ml⁻¹. These values are also displayed as the molar C:N, N:P, C:P and C:N:P ratio in the adjacent text boxes (Figure 2.3). The default values are typical of fungal and microbial cells, although there is considerable variation in both the absolute amounts and the relative ratios.

The total amount of C required (C_t) also includes that used in respiration. This is set by the carbon use efficiency (CUE), defined as the fraction of growth to total assimilation. The default value of CUE is set as 50%, but values can be much lower than this1.

The maximum rate that resource that can be used (by any organism) is set by Lambda, with a default value of 0.3 g ml⁻¹ h⁻¹. This would equate to a doubling time of just under 1h for the default resource density, and represents an upper limit that is unlikely to be achieved for organisms in the wild.

In addition, autolytic, motile and fungal cells have a single organism-specific cost parameter that helps to define their different strategies. Thus, Alpha is the additional cost of being motile (default 0.02), Beta is the cost associated with internal transport (default 0.1), whilst Epsilon is the fraction of an autolytic cell that can be recovered by recycling (default 0.5).

Running the simulation

The Run Model button will iterate through the external resource environmental parameters for each organism, and returns pseudo-

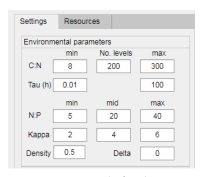


Figure 2.2: Controls for the environmental parameters that define the resource quantity and quality

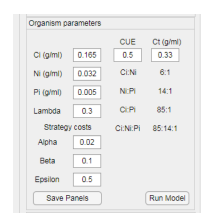


Figure 2.3: Parameter settings for the biological organisms

¹ Margarida Soares and Johannes Rousk. Microbial growth and carbon use efficiency in soil: Links to fungalbacterial dominance, SOC-quality and stoichiometry. Soil Biol. Biochem., 131: 195-205, 2019

colour coded maps showing which class of organism is predicted to grow most rapidly under each set of conditions, according to the inset colour triangle (Figure 2.4). The default settings are matched to the data reported in the main text. The run time is about 160-180 min on a Windows 10 machine with an Intel® Core™ i7-8750H CPU running at 2.20GHz with 32GB RAM. Reduced run times can be achieved by decreasing the number of levels (the minimum suggested is around 20) as run time increases with levels².

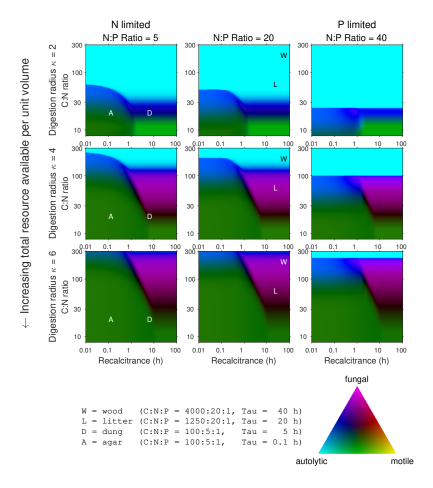


Figure 2.4: Colour-coded maps showing the relative performance of each type of organism across a wide range of resource quantity and quality.

Regions of parameter space where motile cells dominate are coloured cyan, regions where fungi dominate are magenta, and regions where autolytic cells dominate are yellow. In many regions, two or more organisms may fare equally well. Thus, there are typically extensive regions where motile cells and fungal organisms both thrive (blue), or motile and autolytic unicells are equivalent (green).

Displaying the colour-coded results

The cyan-magenta-yellow maps show the relative performance of each class of organism compared to the others. In addition, in the Display panel (Figure 2.5), the Display drop down menu can be used



Figure 2.5: Display options

to show how the individual growth rates compare to the fastest growing cell with no resource restriction, by selecting the *Growth* option (Figure 2.6). This provides comparative plots for each N:P Ratio for each digestion radius, whilst the *Organism* for comparison has to be selected from the corresponding drop down menu. The default lookup table (LUT) gives a heatmap scale. Alternative LUTs can be selected from the *Colormap* dropdown menu. the *Resources* checkbox superimposes letter codes on the plots that correspond to the set of four specific resources described in Section 2.6 below, and shown in the figure legend.

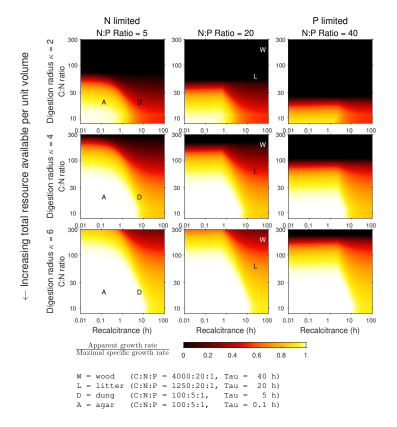


Figure 2.6: Colour-coded maps showing the relative growth of each type of organism compared to the maximum growth of any type of organism. The letter codes superimposed correspond to the region of parameter space associated with specific resources set out in the figure legend.

Alternatively, the relative growth rate of a fungal organism can be compared specifically with one of the other classes of organism to visualise directly the magnitude of the hyphal advantage (Figure 2.7). The default colormap is on a perceptually uniform rainbow scale², but other options can be selected from the *Colormap* dropdown menu, including a number of diverging LUTs (prefix 'D') that highlight the relative performance quite well. The **Print** button saves a copy of the display as a png or pdf file.

² P. Kovesi. Good colour maps: How to design them. *arXiv:1509.03700 [cs.GR]*, 2015

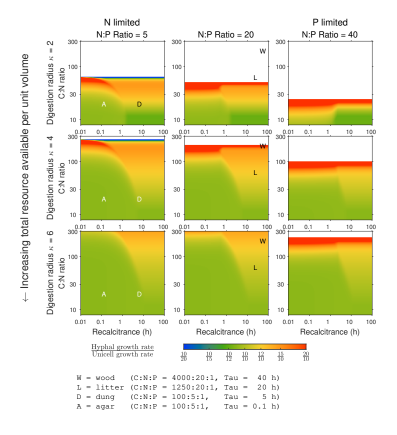
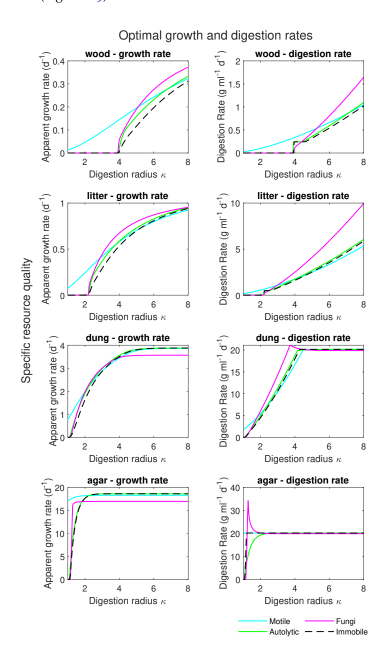


Figure 2.7: Colour-coded maps showing the relative growth of a hyphal organism compared to a specific class of unicellular organism.

Performance of different classes of organism on specific re-2.6

The colour-coded maps display results for a very extensive range of resource environments. The Specific resources panel (Figure 2.8) provides options to plot the performance for four specific resources, with the defaults set to wood (C:N:P 4000:20:1, τ = 40h), leaf litter (C:N:P 1250:20:1, τ = 20), dung (C:N:P 100:5:1, τ = 5h) and agar (C:N:P 100:5:1, τ = 0.1h).

The **Run** button runs the simulation for these specific resource configurations using the other parameters defined for the full model (Figure 2.9)



Whilst the values for C:N:P are widely reported in the literature, the values for τ are not defined experimentally for most substrates.

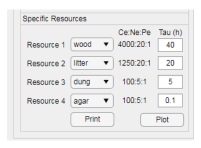


Figure 2.8: Selection of specific resources

Figure 2.9: Colour-coded maps showing the relative growth of a hyphal organism to a specific class of unicellular organism.

Thus, there is the option to vary τ for each resource using the adjacent edit box. For direct comparison, the same resource can be selected in all four dropdown menus and the simulation run with different values of τ . Increasing the value of tau significantly decreases the growth rate for all organisms, but in general, changing tau only has a small effect on the relative performance of different classes of organisms.

2.7 Inclusion of user defined resources

Given that there are many different resource possibilities other than the four defaults used, there is an option to define more substrates using the **Resources** tab at the top of the control panels (Figure 2.10)

The Table includes the four default resource types. Additional resources can be included using the **Add** button, which adds an additional row to the table. Likewise the **Remove** button will delete the currently selected entry. The modified table can be saved using the **Save** button, and re-loaded using the **Load** button. The **Update** button adds the new set of resources to the **Resource** drop down menus in the **Specific Resource** panel in the main **Settings** tab. The effect of changing resources can then be explored using the **Run** button.



Figure 2.10: Resource table to allow addition of specific resources with user-defined C:N:P ratios and recalcitrance (Tau)

Command Line Version

3.1 Overview

The set of MATLAB® scripts that can be used to run the different components of the simulation are given in Table 3.1. These have been tested on versions 2017a and should work in any subsequent release of MATLAB® on any platform. They are the same functions as called by the GUI, or embedded in the standalone versions of the program.

Script Name	Description
vary_CNP_and_tau.m	The main script to run the simulation and generate colour-coded maps of relative performance
vary_kappa.m	Script to generate the growth rate and rate of digestion for each class of organism in a specific resource environment

Function Name	Description
find_best_cell.m	Function to determine numerically the best growth rate (μ) for a given investment in digestive enzymes (x) using the parameter values for a cell growing at the maximum possible rate without substrate depletion which implements equations 1-2 of the Supplementary Information.
find_best_immobile.m	Function to determine numerically the best growth rate (μ) for a given investment in digestive enzymes (x) using the parameter values for immobile cells. Calls the function $find_mu_given_x_immobile.m$. Implements equations 1-6 of the Supplementary Information.
find_best_autolytic.m	Function to determine numerically the best growth rate (μ) for a given investment in digestive enzymes (x) using the parameter values for autolytic cells. Calls the function $find_mu_given_x_autolytic.m$. Implements equations 7-10 of the Supplementary Information.

find_best_motile.m	Function to determine numerically the best growth rate (μ) for a given investment in digestive enzymes (x) using the parameter values for motile cells. Calls the function $find_mu_given_x_motile.m$. Implements equations 11-13 of the Supplementary Information.
find_best_fungi.m	Function to determine numerically the best growth rate (μ) for a given investment in digestive enzymes (x) using the parameter values for fungal cells. Calls the function $find_mu_given_U_fungi.m$. Implements equations 14-17 of the Supplementary Information.

Display functions	Description
generate_CYM_merge.m	Function to generate the Cyan-Yellow-Magenta maps of relative growth rate. If the fastest growing colony has grown by a factor of n, and a slower growing colony has grown by a factor of m, the channels for the colonies are coloured accordingly with intensity 1 and m/n respectively. Higher n increases the apparent difference in colour, making a lighter CYM merge.
heatmap_comparison.m	Generates colour-coded ratio heatmap of the relative growth of fungal cells and autolytic cells in the standard set of resource environments. Calls <i>colorcet.m</i> to define a perceptually uniform ratio colourmap ('R3') and <i>applycolourmap.m</i> to apply it to the ratio growth image. See SI Figure 3.
heatmap_growth_inhibition.m	Generates a heatmap showing the relative growth of each class of organism compared to the maximum growth possible with unlimited resource. Calls <i>colorcet.m</i> to define a cold-hot colourmap ('L3') and <i>applycolourmap.m</i> to apply it to the relative growth inhibition image. See SI Figure 4.

Table 3.1: MATLAB scripts and functions

3.2 Running the simulation for varying resource quality and recalcitrance

Open the vary_CNP_and_tau.m script in the MATLAB® Editor. The first part of the code sets the resource and organism parameters (Listing 3.1), which can be modified manually to explore different environmental resources or impose different costs on the strategy used by each organism. For initial trials, the res1 parameter can be set to a much lower value (20) to reduce the computation time, whist still titrating the range of C:N and recalcitrance values (Figure 3.2).

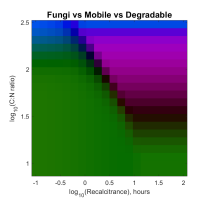


Figure 3.1: Coarse output for rapid prototyping

Listing 3.1: Setting parameter values at the start of the vary_CNP_and_tau code

```
1 %% set up parameters. These can be manually edited.
_{2} % resl is the number of different values for the C:N ratio and recalcitrance
3 % that are tried
_{4} res1 = 200;
6 % xres determines the number of different values for x that are tried in finding
7 % the optimal solution. Final value found is accurate to xres^2.
8 \text{ xres} = 200;
_{10} % sres is the number of iterations used in finding best solution for a given
% environment and a given rate of sythesis.
12 \text{ sres} = 60;
% N_to_P set the N:P ratio of the resource.
N_{to} = 20;
17 % kappa is the radius of digestion, normalised to the radius of the cells/hypha,
18 % and sets the total amount of resource available.
19 kappa = 6;
_{21} % C_to_N_min and C_to_N_max define the limits of the C:N ratio in the resource.
_{22} % The number of intermediate steps is controlled by resl. The C:N ratio is
23 % plotted on the y-axis of the resultant map.
24 C_to_N_min = 8;
C_{5} C_{5} = 300;
_{27} % tau_min and tau_max define the limits of the resource recalcitrance, where tau
  % is the time in hours for hydrolases to digest their own mass. The number of
_{29} % intermediate steps is controlled by res1. The recalcitrance is plotted on the
30 % x-axis of the resultant map.
_{31} tau_min = 0.1;
32 tau_max = 100;
_{34}\, % Ci is the mass of carbon needed per unit volume of growth, in g per ml
_{35} Ci = 0.165;
_{36} Ni = 0.032;
_{37} Pi = 0.005;
_{39} % the carbon use efficiency is the fraction of C for growth versus total C
40 % for respiration and growth
_{41} CUE = 0.5;
_{43} % Ct is the total C required, including respiratory C set by the carbon use
44 % efficiency
45 Ct = Ci/CUE;
47 % dry weight in grams per ml of the substrate.
_{48} density = 0.5;
_{50} % epsilon is the efficiency of recycling for senscent autolytic cells.
_{51} epsilon = 0.5;
_{53} % alpha is the additional mass of machinery needed for cell mobility, relative
_{\rm 54}\, % to the mass of essential metabolic machinery.
_{55} alpha = 0.02;
_{57} % beta is the mass of material in vesicles, relative to the mass of the rest of
58 % the fungus, including hydrolases.
59 beta = 0.1;
61 % lambda is the maximum rate of resource use per unit volume, in g per ml per
62 % hour.
63 lambda = 0.3;
_{65} % _{\Lambda} is the ratio of C that has to be digested to release each N, to reflect
66 % that N is embedded within C-rich polymers.
67 \quad \Delta = 0;
```

The output is a res1xres1 array of the relative growth (Mu) and the amount of digestive enzymes released (x) for each class of organism for the selected ranges of C:N and tau. The relative growth can be compared using the three different display scripts:

- generate_CYM_merge.m gives a three-way comparison using a cyan-yellow-magenta colour-code for motile, autolytic and fungal cells, respectively.
- *heatmap_growth_inhibition.m* compares the relative growth of each organism against a cell with no limitation on resource availability such that the colony continues to grow at the maximum rate under the conditions selected.
- heatmap_comparison.m compares the rate of fungal growth explicitly with autolytic cells to provide a more quantitative visual indication of the hyphal advantage in different resource environments.

3.3 Running the simulation for a specific resource quality in varying abundance

The second main script (Listing 3.2) requires the user to define the resource quality (C:N:P ratios and recalcitrance), but then calculates the relative growth rate for each class of organism and rate of digestion to give the optimal strategy. As with the vary_CNP_and_tau script, the first section has to be edited manually to set up different parameter configurations. The output is automatically generated as a plot of growth rate against resource abundance (varying kappa) and rate of digestion for the four classes of organism.

Listing 3.2: Setting parameter values at the start of the vary_kappa code

```
1 %% set up parameters. These can be manually edited.
2 % res1 is the number of different values for kappa that we try
_3 res1 = 200;
5 % xres determines the number of different values for x that are tried in finding
6 % the optimal solution. Final value found is accurate to xres^2.
_{7} xres = 200:
_{9} % sres is the number of iterations used in finding best solution for
10 % a given environment and a given rate of sythesis
11 sres = 60;
_{\rm 13} % k\_{\rm min} and k\_{\rm max} are the minimum and maximum values for the resource
_{14} % acquisition radius that we try
15 k_{min} = 1;
_{16} k_max = 8;
18 % Ci is the mass of carbon needed per unit volume of growth, in g per ml
_{19} Ci = 0.165;
_{20} Ni = 0.032;
Pi = 0.005;
_{23} % the carbon use efficiency is the fraction of C for growth versus total C
24 % for respiration and growth
```

```
<sub>25</sub> CUE = 0.5;
27 % Ct is the total C required, including respiratory C set by the carbon use
28 % efficiency
29 Ct = Ci/CUE;
_{31} % C_to_P and N_to_P set the C:P and N:P ratio of the resource, respectively.
C_{12} C_{13} = 2000;
_{33} N_to_P = 10;
_{35} % tau is the time in hours for hydrolases to digest their own mass and reflects
36 % the recalcitrance of the resource.
_{37} tau = 40;
39 % dry weight in grams per ml of the substrate
_{40} density = 0.5;
41
_{\rm 42} % _{\rm A} is the minimal fraction of C that must be digested in order to
_{43} % digest the available N
_{44} _{\Delta} = 0;
46 % epsilon is the efficiency of recycling for autolytic cells
_{47} epsilon = 0.5;
_{49} % alpha is the mass of machinery needed for cell mobility, relative
50 % to the mass of essential machinery
_{51} alpha = 0.02;
_{53} % beta is the mass of material in vesicles, relative to the mass of
_{\rm 54}\, % the rest of the fungus, including hydrolases
55 beta = 0.1;
57 % lambda is the maximum rate of resource use per unit volume,
58 % in g per ml per hour
```

 $_{59}$ lambda = 0.3;

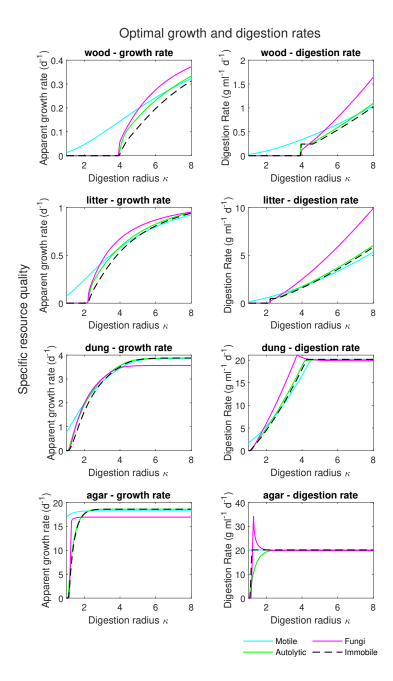


Figure 3.2: Growth rate and digestion rate for a specific resource quality and quantity