
Modelling Competition for Nutrients between Microbial Populations Growing on Solid Agar Surfaces

Author: Daniel Boocock; Supervisor: Dr Conor Lawless

August 9, 2016

ABSTRACT

1.1 Subsection

Motivation: Growth rate is a major component of the evolutionary fitness of microbial organisms. When nutrients are plentiful, fast-growing strains come to dominate populations whereas slower-growing strains are wiped out. This makes growth rate an excellent (a useful) surrogate for the health of cells. Measuring the health of cells grown in different genetic backgrounds or environments can inform about genetic interaction and drug sensitivity. In high-throughput procedures such as QFA and SGA, arrays of microbial cultures are grown on solid agar plates and quantitative fitness estimates are determined from growth measurements. Diffusion of nutrients along gradients in nutrient density arising between fast- and slow-growing neighbours is likely to be affecting growth rate and fitness estimates. However, current analyses assume that cultures grow independently. We study data from QFA experiments growing *Saccharomyces cerevisiae* to test a mass action kinetic model of nutrient dependent growth and diffusion. We try to correct for competition to provide more accurate and reliable fitness estimates.

Results: Don't know what to say yet.

Availability and Implementation: CANS, a Python package developed for the analysis in this paper, is freely available from [github](#).

Contact: daniel.boocock@protonmail.ch

1 INTRODUCTION

Dummy Lawless *et al.* (2010) citations (Heydari *et al.*, 2016) (Addinall *et al.*, 2008).

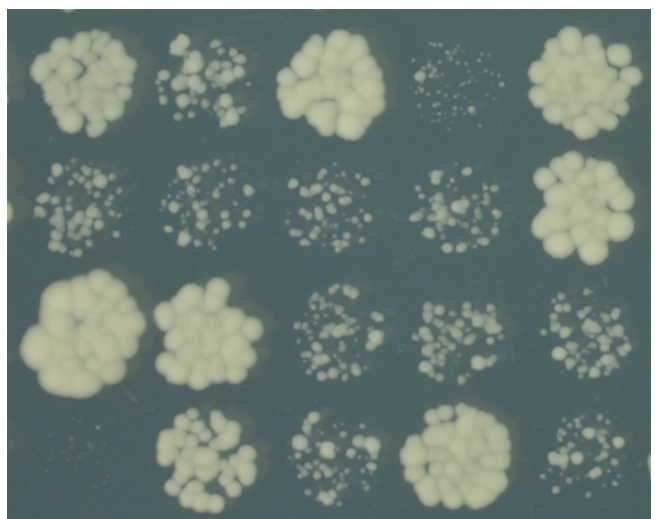


Figure 1: 4x5 section of a QFA plate. Taken from a 16x24 format solid agar plate inoculated with dilute *S. cerevisiae* cultures. Image captured at ~2.5d after inoculation and incubation at 27°C.

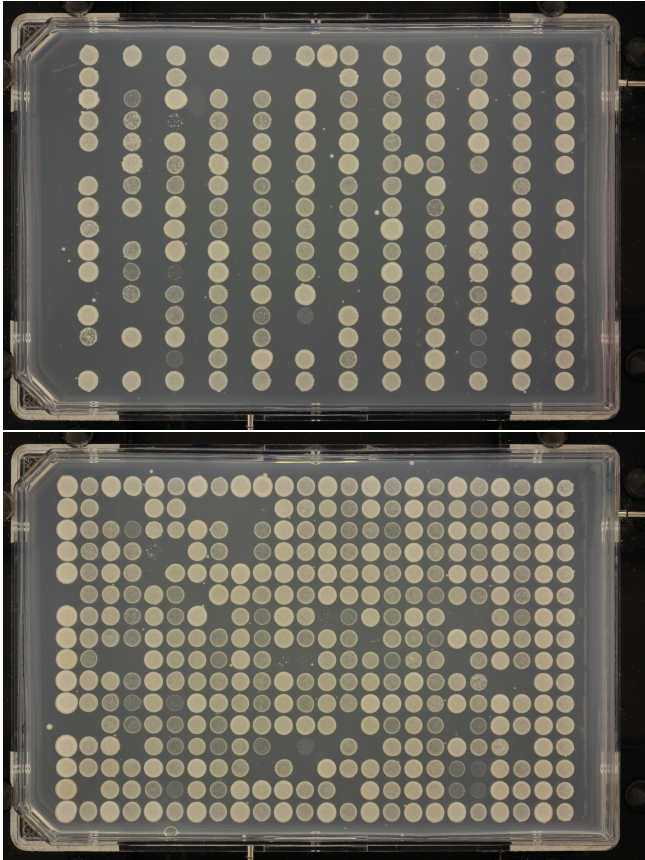


Figure 2: An experiment designed to examine competition.
A) QFA plate inoculated with a more concentrated *S. Cerevisiae* inoculum (no cells inoculated on alternative columns).
B) Same as in A, but with strains of similar growth rate inoculated in the positions missing in A.

2 METHODS

2.1 Subsection

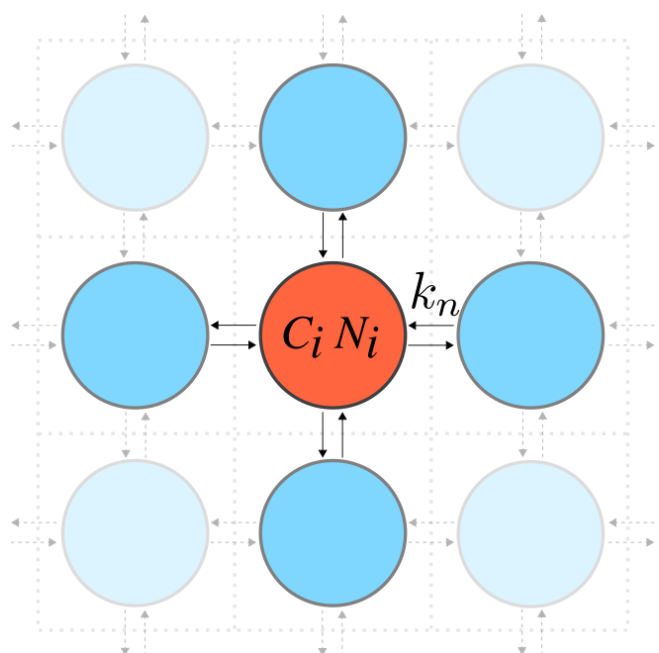


Figure 3: Schematic of the modelling approach. Each circle represents a culture, indexed i , growing in a grid on solid agar. Arrows represent the diffusion of nutrients in the network of cultures. C_i - amount of cells; N_i - amount of nutrients; k_n - plate level diffusion constant; darker blue circles - neighbourhood of culture i : δ_i .

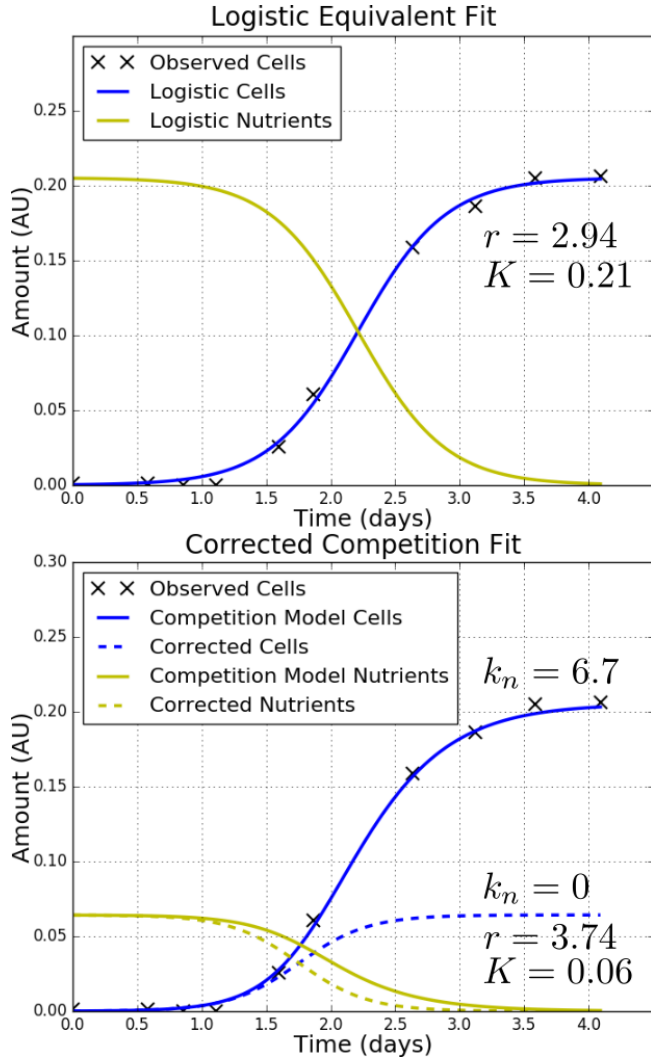


Figure 4: Using the competition model to correct for competition. Fits are to culture (R10, C3) of P15 which grew faster and reached a higher final cell density than its neighbours (not shown). According to the competition model, this is because this culture competed for more nutrients. To reach the same final cell density, the logistic equivalent model requires a higher amount of starting nutrients for this culture and a different amount for each neighbour. The correction to the competition model simulates how growth would have appeared without competition and allows us to return parameters r and K of the logistic model.

3 RESULTS

3.1 Guessing

N_0 estimated from average final cell amounts. See formula in code for two N_0 estimation.

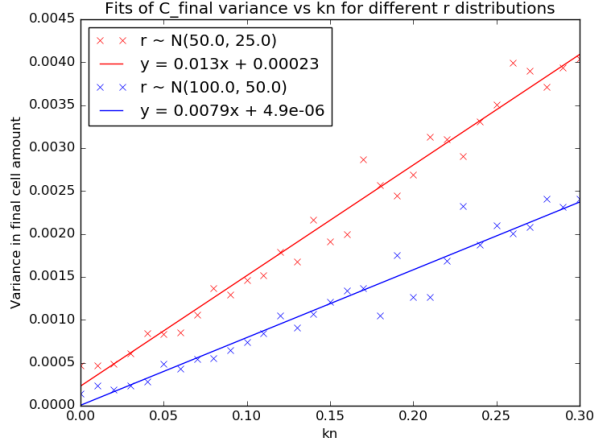


Figure 5: Guessing k_n from the variance in final cell amounts. The competition model is simulated for a 16x24 format plate using two random sets of culture-level b parameters drawn from different normal distributions. Each set of b parameters is simulated with a range of k_n parameter values. The variance in final cell density for all cultures is plotted against k_n for each simulation. Lines are shown for least squares fits to points from each set of b parameters.

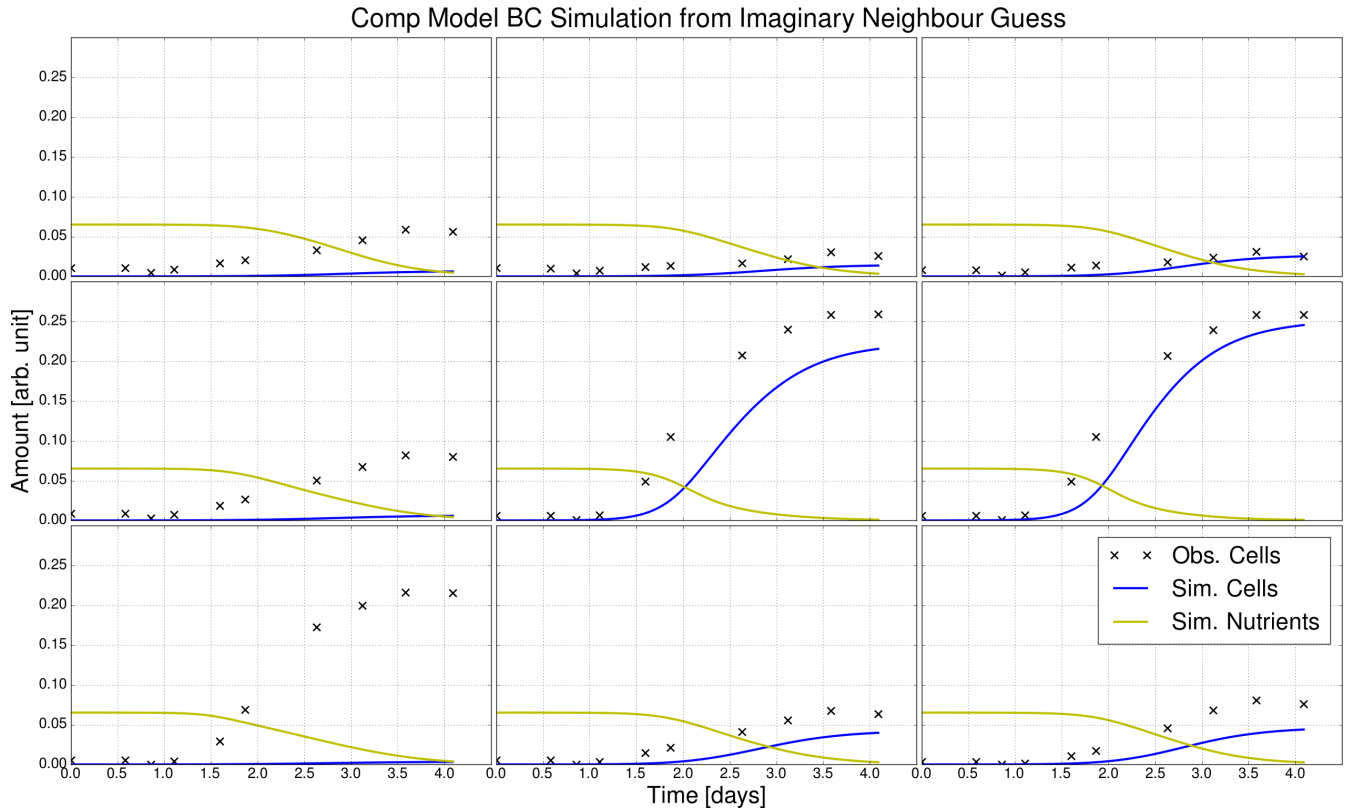


Figure 6: Competition model simulation using parameters from imaginary neighbour guessing. Shows a 3x3 zone with top-left coordinate (5, 18) from P15 with background *cdc13-1* at 27°C.

(NOT TO GO HERE This method of guessing requires a b-guess to be supplied to fix the faster growing neighbour. (I iterated through cell ratios. I iterated through a range of b guesses supplied at the plate level; running a different script with a C_0 guess, b_{guess} . It would probably have been better iterate through a list of b-guess values for each culture and choosing the estimated b value from the best fit of each culture. Guessing time is currently about four minutes which is fast compared to fitting which takes approximately three hours. However, this is unlikely to stop us from encountering local minima when we fit the Competition Model.)

Scripts were run with combinations of the following values. `cellratios = np.logspace(-3, -5, num=5)` `fittype = ["imagineigh", "logeq"]` `zerokn = [True, False]`

Each script looped through the following array of b values which were supplied to the initial guesser and used at the plate level.

```
for b-guess in [35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 95, 100, 150]:
```

Each b-guess value is used to guess a complete set of b parameters for every culture in the plate. Each of these parameter sets is then used as an initial guess to Competition Model fitting. For the 13 b-guesses we

must run 13 Competition Model fits. It would be better and more efficient to loop through the b-guesses at the culture level. Each culture still undergoes imaginary neighbour guessing with each of the 13 b-guess values, but now, for each culture, we choose just the b estimate from the best of the 13 fits. This will produce one set of b guesses which should be superior to any of the guesses attained when iterating through b-guess at the plate level. Then we only need to fit the Competition Model to 1 guess rather than 13. This will reduce the number of scripts that need to be run in parallel, or the use of a finer grid over C_0 , and should make convergence faster. However, if using a gradient method we are still likely to encounter local minima from these guesses. Instead, this improvement could be considered when developing a genetic algorithm (if initial guesses are required) or if fitting using a brute force method with a fine grid of fixed plate level parameters. We will see later that with true plate level parameters fixed we can recover good estimates for b using a gradient method. It may be possible to evolve candidates of plate level parameters, fix these, and minimise using the current gradient method.

3.2 Competition Model Fitting to P15

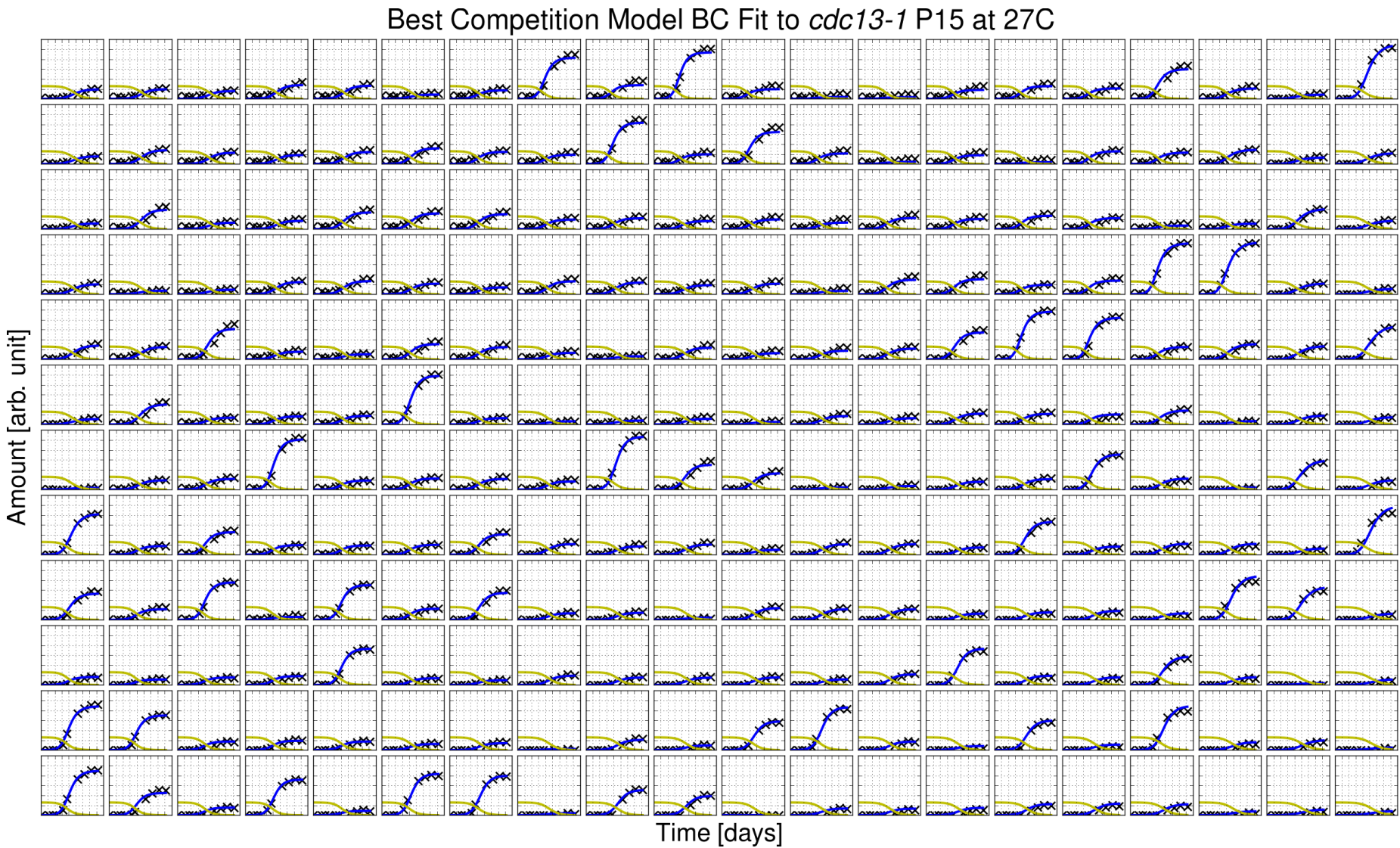


Figure 7: Fit of the competition model to a QFA plate. Data is for a 16x24 format plate (P15) with a background mutation *cdc13-1* incubated at 27°C. The plate contains 6 repeats of 50 genetic strains randomly arranged across the internal cultures. Repeats of a single strain are used for all edge cultures (removed in the plot). Model output for state variable, cell population size (blue curve), is fit to observed data (black crosses). Model predictions for unobserved variable (nutrient amount) are also plotted (yellow).

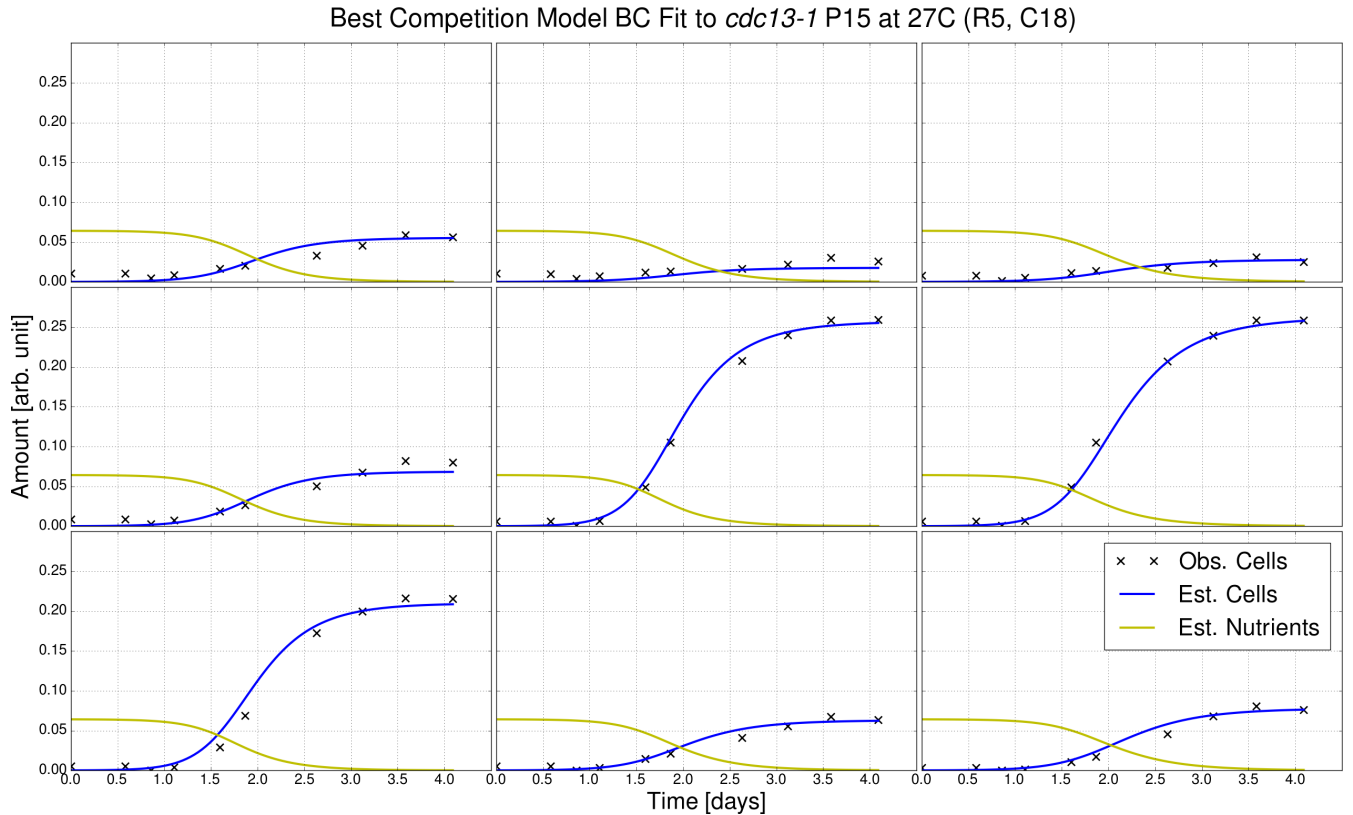


Figure 8: A 3×3 zone from Figure 7 with top-left coordinate $(5, 18)$.

3.3 Evaluating the treatment of boundaries

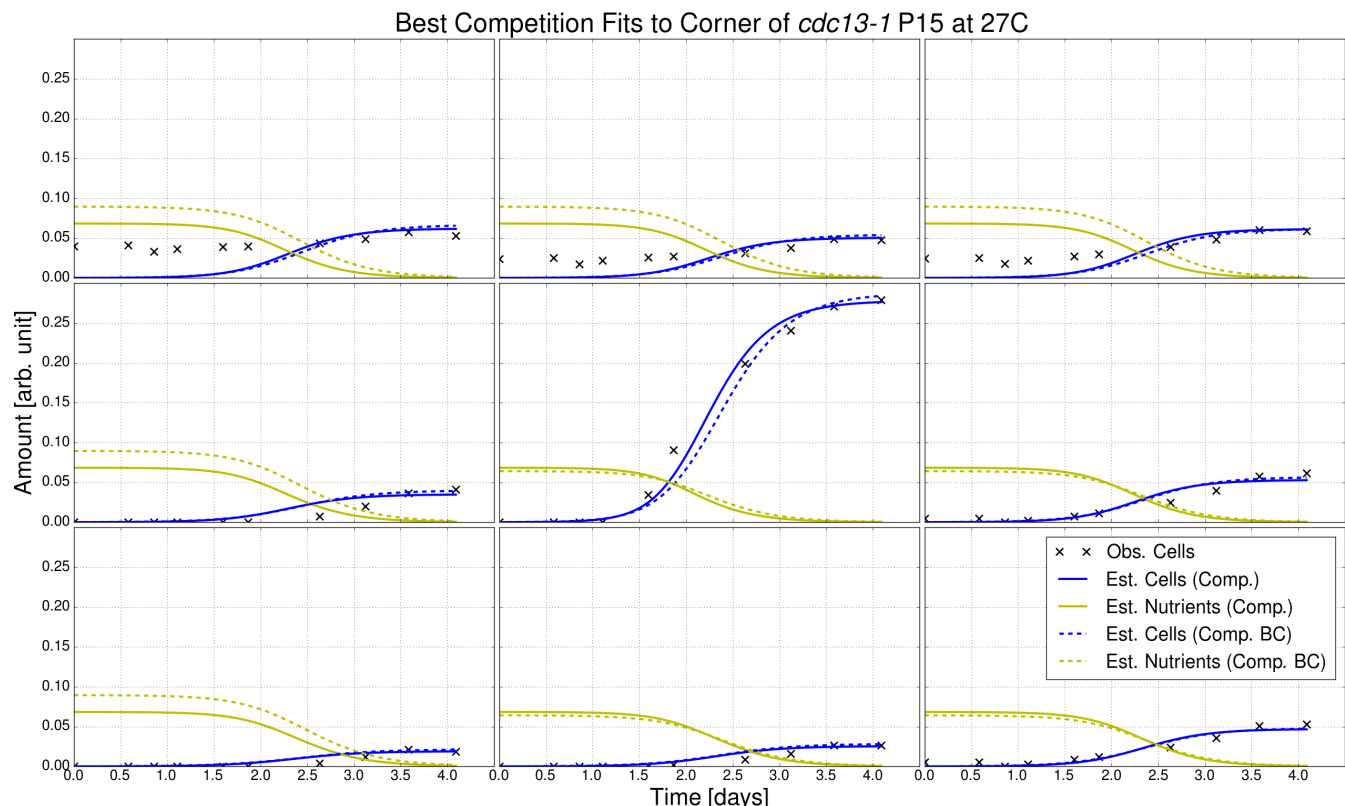


Figure 9: Treatment of boundary conditions in fits of the competition model. The top left corner of a 16x24 QFA plate fitted with two versions of the competition model: the first containing a single initial nutrient amount for all cultures; the second containing a separate initial nutrient amount for edge cultures.

3.4 Agreement of b rankings

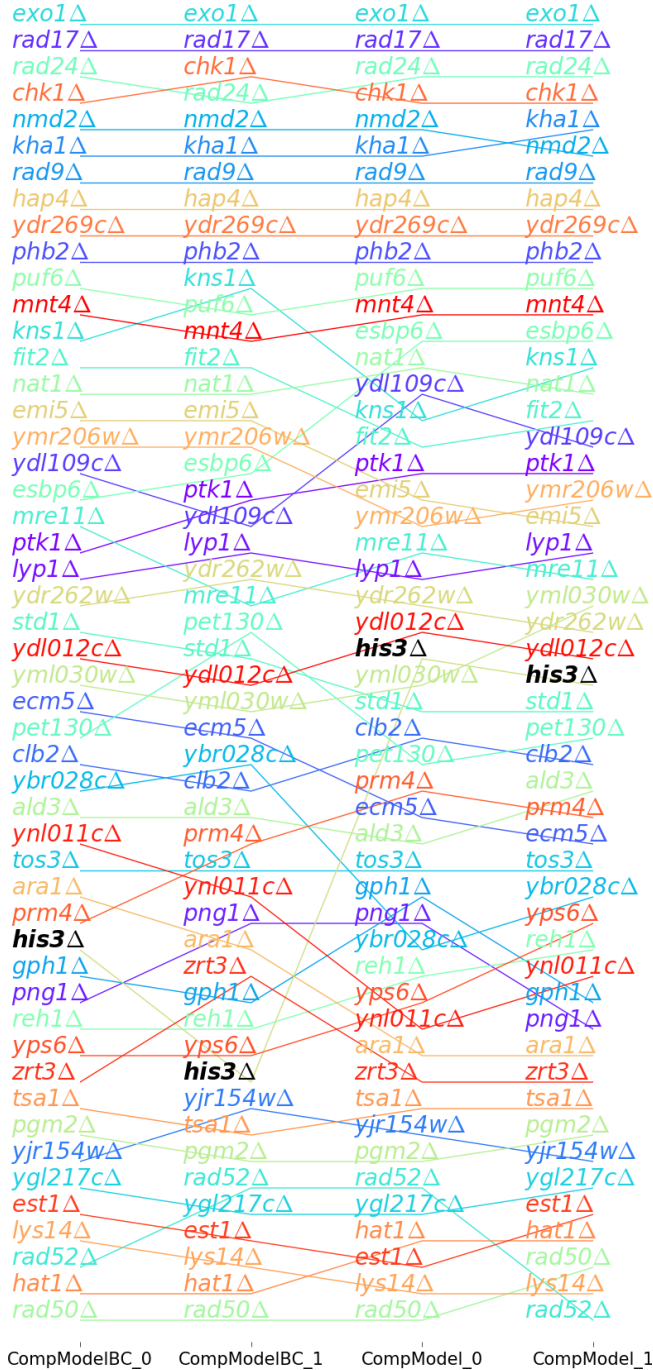


Figure 10: Comparison of b ranking for the best five competition model fits to P15. Ranking is calculated from the mean b estimate from the six repeats or each strain.

3.5 Comparison of fitness ranking



Figure 11: Comparison of r ranking for fits of the competition and logistic model to P15. Competition model r was converted from b, N_0 , and C_0 from the best competition model estimate. Logistic r was taken from fits using the QFA R package which makes heuristic checks for slow growing cultures.

3.6 Comparison of Variation in Fitness Estimates

Use repeats on plate 15 (6 per deletion) to calculate coefficient of variation (COV) of estimated r or MDR.

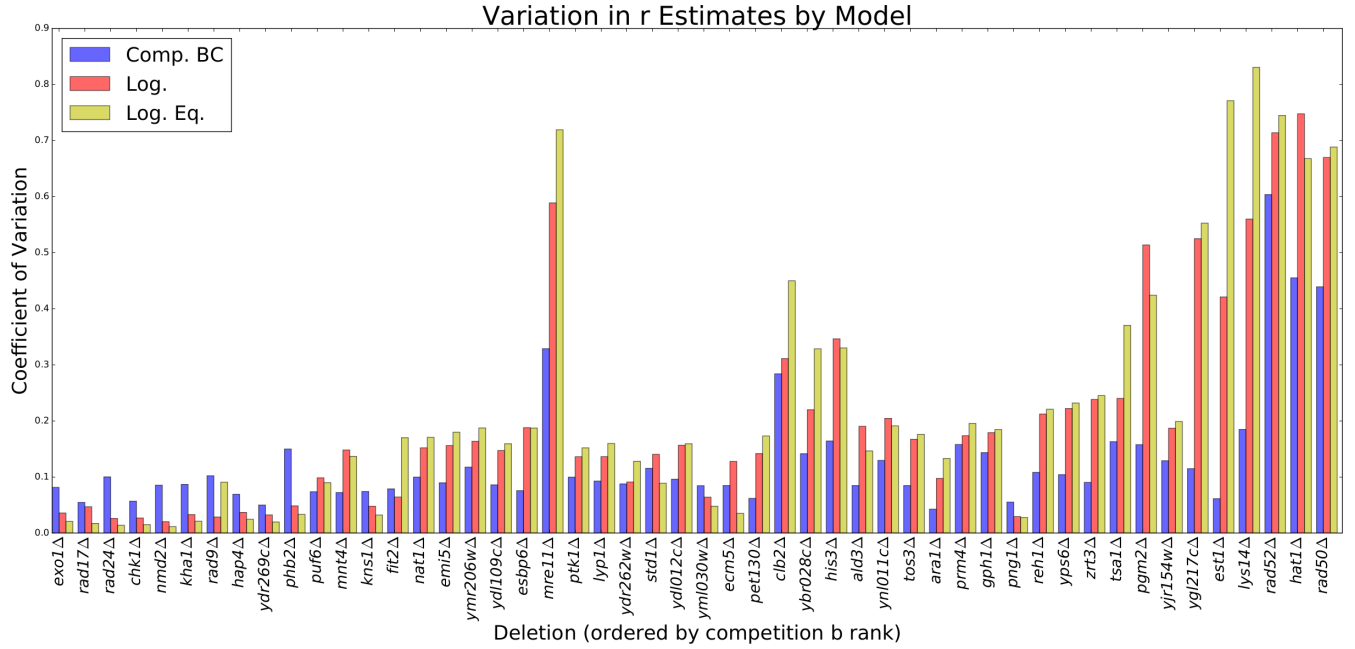


Figure 12: Coefficient of variation of r estimates. Strains are ordered left to right along the horizontal axis by highest to lowest competition model r ranking. Fits are for the competition model, the QFA R logistic model, and the logistic equivalent model.

3.7 Cross-plate validation

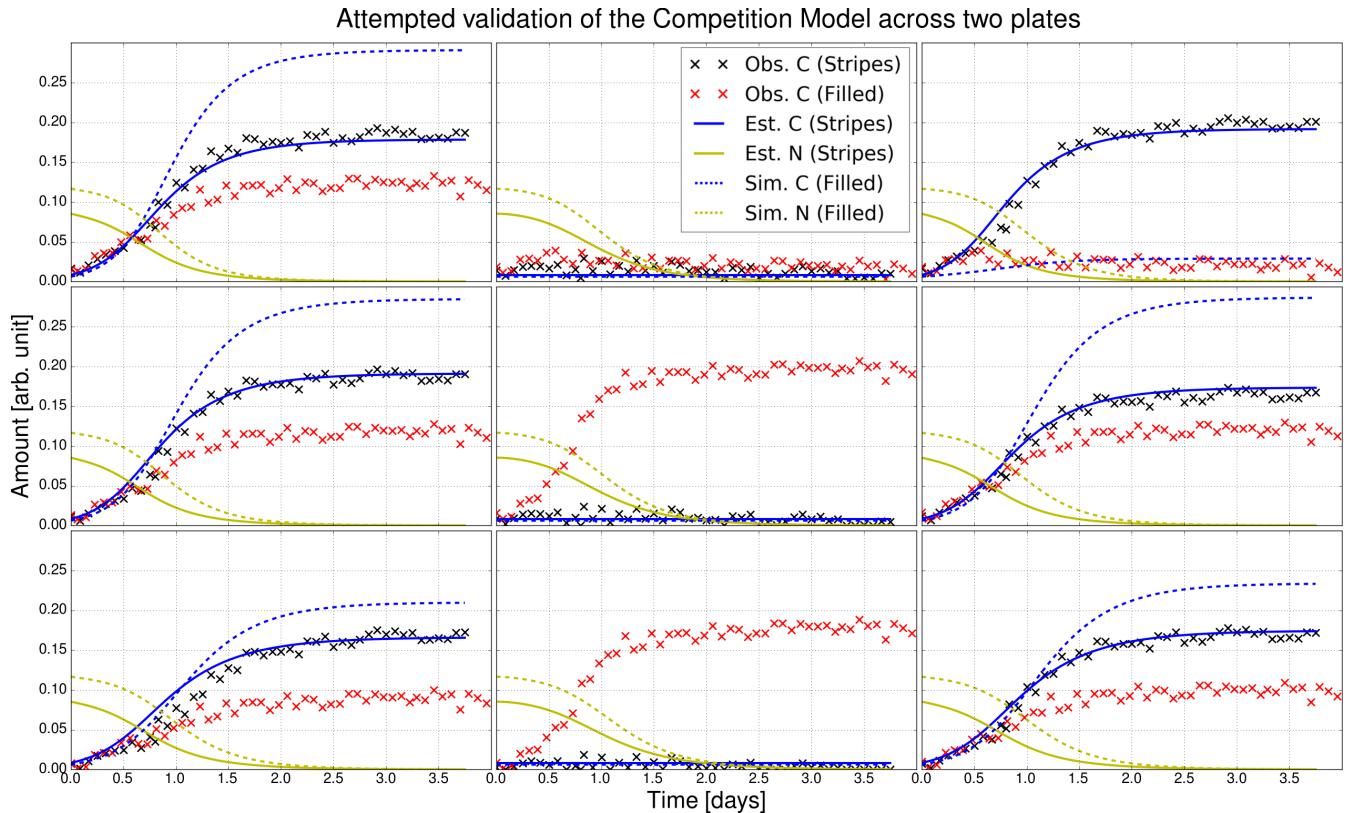


Figure 13: Calibration and validation of the competition model.

3.8 Towards a genetic algorithm

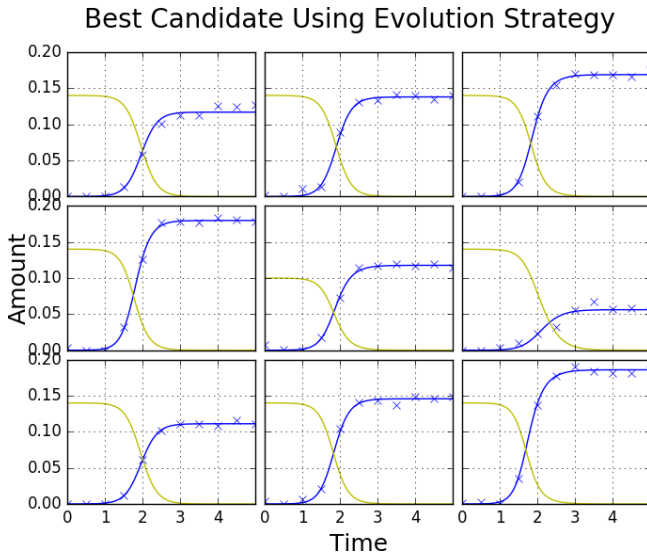


Figure 14: Genetic algorithm fit to a 3x3 simulation. MIGHT TAKE A LITTLE BIT OF WORK TO REPRODUCE AND COULD USE PARAMETERS FROM THE BEST P15 FIT RATHER THAN JUST PICKING/RANDOMIZING. NEED TO CHECK THAT PLATE LEVEL PARAMETERS WERE ALSO EVOLVED.

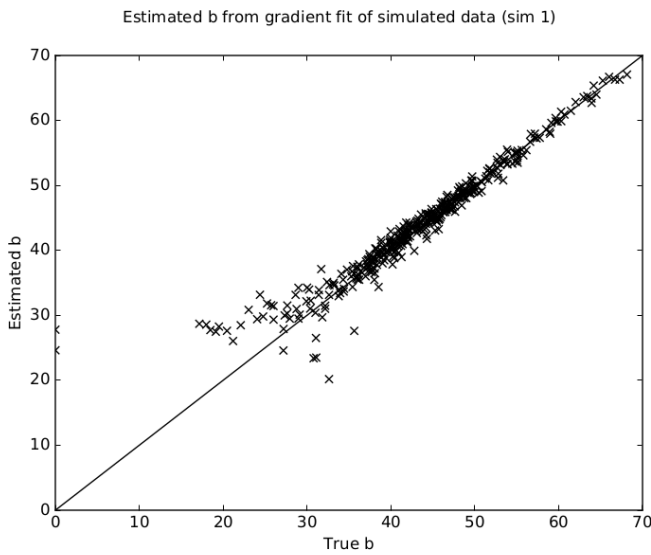


Figure 15: Recovery of true b values from a gradient method with fixed plate level parameters. I simulated timecourses from the best five (which model? all BC?) fits to p15, fixed the true plate level parameters, and used a gradient method to recover b . This plot shows the worst case from the five sets of values.

4 DISCUSSION

Fitness ranking from competition model fits may be better than from logistic model fits (Will comparing stripes rankings reveal anythin?). However, we cannot quantitatively compare fitness estimates between plates be-

cause we are not finding global minima. Work has begun to develop a genetic algorithm to do so. I am not convinced that this will succeed because growth is systematically overestimated when we move from the filled to striped plate for all of the current best parameter solutions. This suggests an issue with the modelling approach; below I suggest ways in which this could be improved. In any case, qualitative cross-plate validation using order of fitness ranking may still be better (for the competition model).

The first thing to notice about QFA data - from P15, the striped plate, and the filled plate - is the characteristic endpoint in growth on each plate (experiments could be designed to study variation in timescales over regions of a plate by inoculating cultures in columns left-to-right according to fitness). This suggests a plate-level or region-level growth-limiting effect. // Could this conceivably be an experimental limitation such as the drying out of an agar plate over time? // Comparison of the striped and filled data, shows that cultures grow larger when neighbours are removed and this suggests a direct interaction between cultures. The strongest candidates are competition for nutrients and growth limiting signalling such as ethanol poisoning. It is possible that other growth limiting effects may exist and could confound any attempt to fit a model which accounts for just one of these. It makes sense to investigate each likely effect in turn to determine its contribution and to start by validating the independent limit.

Spots can grow after a long time. Must be nutrients remaining or an encroachment? I have an image for the stripes plate showing cultures growing and believe after a very late stage. I need to check the data images but this may just be encroachment of another culture.

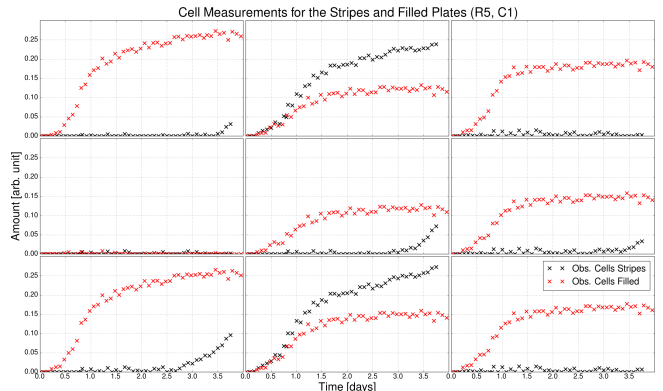


Figure 16: Observed cells for (R5, C1) 3x3 zone of Stripes and Filled plates showing (for Stripes data) slow growing cultures starting to grow after faster growing cultures have reached the stationary phase.

We have only studied data where cultures are grown

in an array on solid agar where we cannot validate the independent limit. In this limit, our model says that nutrients can only be converted to cells and all cultures starting with the same amount of nutrients will reach the same final cell density. This ignores metabolism which may differ between strains. Cell arrest could also limit growth (and this may occur in different strains at different rates). If present, differences in such effects could account entirely for differences in final cell density. However, they are unlikely to be the only effect, because this would not lead to the observed characteristic endpoint in growth. Using one-culture spot tests (in a petri-dish on agar?) or liquid cultures we can grow cultures independently and validate the independent limit. A current issue with methods for estimating fitness, is that identical strains grow differently on agar or in liquid culture leading to different fitness rankings (cite). This problem need not affect our validation as we can simply define a culture to have different parameters for growth in either medium. A greater difference may be caused by the dimensionality of the environment. Mass action kinetics is derived for reactions in a three-dimensional (gas or fluid?) (Guldberg and Waage C.M. Guldberg and P. Waage, Studies Concerning Affinity, C. M. Forhandlinger: Videnskabs-Selskabet i Christiana (1864), 35) and this approximation is more valid for liquid cultures than for cultures spotted onto a surface. I suggest to study first the more ideal case of liquid cultures and later see if the model holds for cultures grown on a surface. If it does not, it may be necessary to use a fractal kinetics model (I have references for this from the proposal) or, if the reaction is diffusion limited, consider a more detailed model of nutrient diffusion.

// Model equations for metabolism.//

Our model splits the agar into a grid with volume discretised per culture. In the stripes validation, we overestimate the effect of diffusion when neighbouring cultures are removed. I believe that we are not accurately capturing the point at which growth becomes diffusion limited and that nutrients are well approximated as being evenly distributed within the spatial scales that we model. A diffusion equation model could capture the local distribution of nutrients around a culture when the stationary phase is reached. Reo and Korolev (2014) use the diffusion equation (and Neumann and Dirichlet boundary conditions) to simulate nutrient dependent growth of a single bacterial culture on a petri dish in two-dimensions. They create a sink for nutrients from culture growth and equate the flux of nutrients through the culture area with the rate of increase in culture size. They model culture area as varying and keep culture

density constant. We may instead keep culture area constant, allow culture density to vary, and use our mass action kinetic model (eq no.) for the nutrient sink and culture growth. Simulating or fitting this model could help us learn more about diffusion in QFA experiments. It is probably computationally unfeasible to use such a detailed model to fit a whole plate. However, if necessary, it may be possible to use a finer grid to increase compartmentalisation of nutrients to capture spatial heterogeneity in the distribution of nutrients around a culture and the effect of diffusion limited growth. This could extend the validity of the competition model over a larger range of variability in culture growth rates (for instance when some cultures are left empty and others are very fast growing.)

It would also be useful to determine experimentally how nutrients are distributed throughout the agar at the stationary phase. Gaps could be left in an array of cultures and only inoculated once the stationary phase is reached. If they grow then nutrients remain. (Still require simulations to see distribution across depth). This could be extended by growing a single column of identical strains and, after the stationary stage has been reached, inoculating identical strains on the same plate at different distances from the row.

//Talk about an improvement to the imaginary neighbour model.//

Nutrients (sugars, nitrogen, etc.) in QFQ agars are of a standard composition, designed to reduce the excess of any single nutrient (check QFA paper and cite). ((background) What is the nutrient? Nitrogen is only used to build molecules for new cells, whereas sugars are also used for metabolism.) For modelling nutrient limited growth, especially across plates, it would be useful to know the identity of the limiting nutrient and ensure that it is always the same. We could achieve this using a different formula of agar.

The design of the stripes validation experiment could be improved. Rather than filling gaps with cultures not present on the stripes plate, and for which we have no b estimates, we could fill with repeats of the cultures already present on the stripes plate. (I'm not sure it makes any difference actually whether we validate from one direction to the other). It would also have been helpful to have repeats to study differences in COV between the competition and independent models. In order to make sure that competition effects were present in data, we made a drastic change between the stripes and filled plates. This provides a stern validation. The model assumes that competition effects are present whenever there is a difference in final cell amounts between cul-

tures. We could have first validated the model against a smaller change, by varying between slower and faster growing cultures rather than none and very strong growing cultures. If the model works well between such plates it may work well for the majority of QFA experiments which typically have smaller differences between cultures than the data we study. If we did want to test the in an extreme case we could have inoculated fast growing cultures next certain strains and not others to try to induce a change in ranking for which the competition model might compensate better than the logistic model.

//Signalling//

If we find that competition for nutrients is not a significant effect, for instance if growth becomes diffusion limited before nutrients from neighbours can be accessed, then we could instead model signalling by ethanol as the interaction effect. This may be modelled similarly to how we are already modelling nutrient diffusion.

//Signalling equation//

If there is any combination of competition, metabolism, signalling, or arrest contributing significantly to differences in the growth of cultures and the interaction between neighbours then it will be difficult to separate them when fitting a model to data. We may have to develop ways to calibrate effects in isolation (e.g. by adding/measuring ethanol?) and use this information when fitting to high-throughput data.

It is quicker to fit to small zones of a plate but as these have a larger proportion of edge cultures boundary conditions become important. In current data (e.g. P15), different cultures surround the edge and this makes accurate fitting difficult. As a result we must work with larger zones that take longer to analyse. We could surround small 3x3 and 4x4 zones with an empty ring and only need to consider net flux of nutrients across the boundary and not local variation due to different cultures surrounding the zone. We could also surround with the same, low-variance, strain to reduce net flux.

//Stochastic effects//

Unpublished work by Hermann and Lawless has investigated heterogeneity between cell lines within single QFA spots. They have found that a single or small number of extremely fast growing cell-lines come to dominate the population of a single culture. The implication is that cultures with a lower starting cell densities are likely to have greater variance between repeats. We could use higher starting cell concentrations to reduce this variance but then we study less of the growth phase. It may be possible to reduce heterogeneity by inoculating from the exponential growth phase rather than the stationary

phase and still study full growth curves. (Unless mean population growth constant is being studied...) Starting cell densities should ideally be as close to the lowest resolvable level as possible.

//Ways to measure C_0 // There is a confounding effect between initial cell density and b value with may justify using initial cell densities slightly above the minimum detectable level. Heterogeneity within cultures is an issue again here and cell density would effectively be lower than the measured value, e.g., if most inoculated cells are dead or slow growing.

To fit growth curves more accurately QFA has begun using the generalised logistic model (cite). Fitness estimates (MDR*MDP or MDR?) from this model have higher coefficient of variation than those from either the standard logistic or competition model. (accuracy and precision? Could for instance a step function be less variable than the standard logistic model?) Although the fits to data are qualitatively worse, it may be advisable to revert to the standard logistic model.

The logistic model requires different K parameters (N_0 for log. eq.) to be fit for each culture. The competition model shares information about N_0 between cultures and therefore has 383 fewer parameters for a full plate (Could this also explain the higher variance for the fastest growers?). For the slowest growing cultures, noise is more dominant and there is a confounding effect between r and K. To deal with this, the QFA R package uses heuristic checks. In the case of *est1Δ*, this has led to a dramatic disagreement in estimated fitness with the competition model. The estimate from (which model)? agrees better with existing biological knowledge (/Independent spot experiments?). (Is the competition model then useful?)

//QFA R is fixing C_0 rather than fitting (I used a grid)//

//Improvement to imaginary neighbour guessing//

REFERENCES

- Addinall, S.G. *et al.* (2008) A genomewide suppressor and enhancer analysis of *cdc13-1* reveals varied cellular processes influencing telomere capping in *Saccharomyces cerevisiae*. *Genetics*, **180**, 4, 2251–2266.
- Addinall, S.G. *et al.* (2011) Quantitative fitness analysis shows that nmd proteins and many other protein complexes suppress or enhance distinct telomere cap defects. *PLoS Genet*, **7**, 4, 1–16.
- Heydari, J. *et al.* (2016) Bayesian hierarchical modelling for inferring genetic interactions in yeast. *Journal of*

- the Royal Statistical Society: Series C (Applied Statistics), **65**, 3, 367–393.
- Lawless, C. et al. (2010) Colonyzer: automated quantification of micro-organism growth characteristics on solid agar. *BMC Bioinformatics*, **11**, 1, 1–12.
- Palková, Z. et al. (1997) Ammonia mediates communication between yeast colonies. *Nature*, **390**, 6659, 532–536.