



Recommendations for Fitting Microbial Growth Models

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March 2020

Computational Method in Ecology and Evolution MSc

3675 words

For Samraat

Abstract

The modelling of microbial growth is vital for developing food preservation measures to decrease food waste. By fitting a range of mechanistic and phenomenological models to a large dataset of 305 growth curves, I demonstrate that temperature has a major impact on microbial population dynamics. I also examine how methodological choices can influence the results of model fitting studies, including demonstrating that the choice of model can systematically bias parameter estimates to the extent that they become incomparable for meta-analysis. Improvements in the resolution of parameter estimates when models are fit in logarithmic space relative to linear space model fitting are also reported. These findings result in recommendations for microbial modelling methodology, particularly when data are aggregated across sources.

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1 Introduction

There is great interest in modelling the growth of microbial populations, such that the biology of microbial growth and its response to external variables can be understood [1]. Through this modelling, food shelf life can be predicted, and preservation methods can be optimised. In this project, I address four questions:

1. Which models best describe microbial population growth, and how does this vary with temperature?
2. How does temperature affect microbial growth?
3. How does choice of model affects parameter estimates?
4. Does the choice to fit models in linear or logarithmic space affect results?

I addressed these by fitting mechanistic and phenomenological models to a large dataset of 305 digitised growth curves. Each mechanistic models describes the growth curve with a mathematical relationship representing a hypothesis about the dynamics of microbial growth [2], so can be parameterised with meaningful biological quantities. In contrast, the phenomenological models are simply convenient mathematical functions used to describe patterns in the data, and can be used as a neutral comparison for the mechanistic models. By comparing the fit of

the mechanistic models to each other, and to the phenomenological models, we can determine which underlying biological hypothesis has the best support from the data. I also examine the effect of temperature, both on the relative support for the models, and on the growth rate and lag time of bacterial populations, as these play a central role in food spoilage.

By comparing parameter estimates between models, and comparing results from fits in logarithmic and linear space, I also investigate how methodological choices can impact results. In particular, I was interested in whether differences in methodology could invalidate comparison of parameter values from different sources in meta-analyses.

The Mechanistic Models

The mechanistic models can be split into three four-parameter models, and the three-parameter logistic model. The four-parameter models split the bacterial growth curve into a lag phase, exponential growth phase, and stationary phase, with the three-parameter logistic model discarding the lag phase.

Logistic The logistic model [3, 4] assumes that the microbial population immediately starts growing exponentially, with its growth rate falling towards zero as the population size approaches a carrying capacity. This is described by the logistic differential equation:

$$\frac{dN}{dt} = \mu_{max}N(1 - N)$$

With solutions:

$$N_t = \frac{N_0 N_{max} e^{\mu t}}{N_{max} + N_0(e^{\mu t} - 1)}$$

Four-parameter models

The four-parameter models parameterise the sigmoidal growth curve with minimum and maximum population sizes, maximum growth rate, and a time lag before the population starts growing - defined as the intercept between the tangent to the curve at the point of inflection, and the line $N = N_0$ (Figure 1). As such, the rate of growth and length of the lag phase can be independently varied (Figure 2)

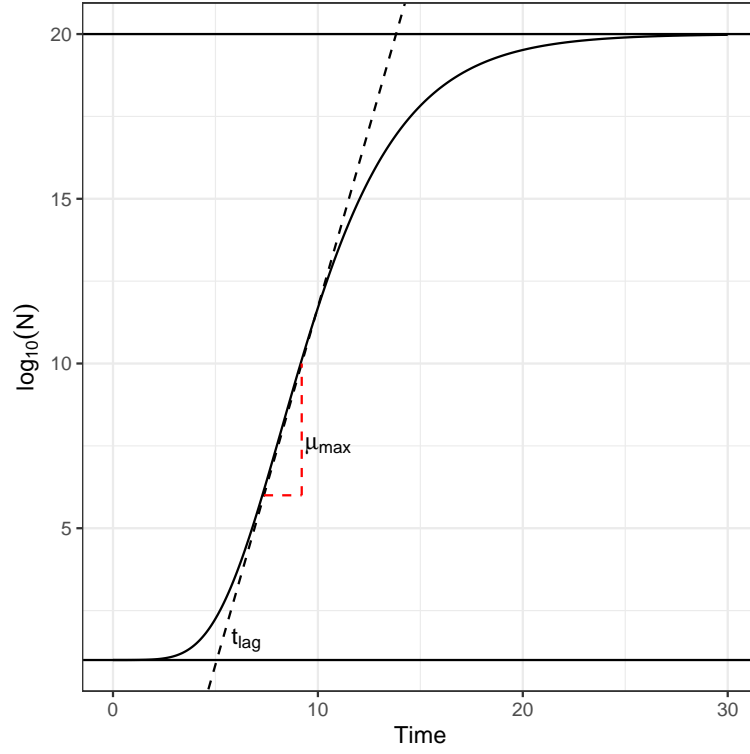


Figure 1: Illustration of μ_{max} and t_{lag} parameters. μ_{max} is the maximum gradient of the growth curve, and t_{lag} is the intercept between this tangent line and the line $N = N_0$

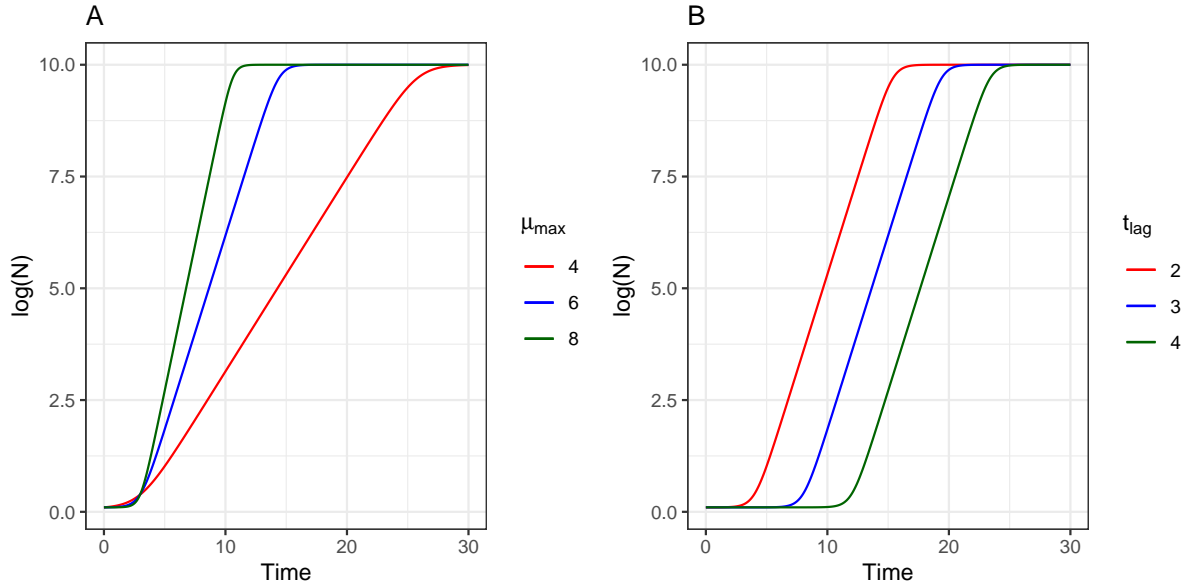


Figure 2: (A) μ_{max} and (B) t_{lag} parameters can be varied independently to control the rate of microbial growth, and the duration of the lag phase.

40 **Buchanan** The Buchanan model [2] is the simplest implementation of this, with a growth rate
 41 of zero during the lag and stationary phases, which surround a period of exponential growth:

$$\log(N_t) = \begin{cases} N_0 & \text{if } t \leq t_{lag} \\ N_0 + \mu_{max} \cdot (t - t_{lag}) & \text{if } t_{lag} \leq t \leq t_{max} \\ N_{max} & \text{if } t \geq t_{max} \end{cases}$$

Gompertz The Gompertz model [5, 1] similarly describes a sigmoidal growth curve, adding a gradual transition into and out of the exponential phase. The Gompertz model has long been popular for microbial population modelling, as it has empirically been found to fit well in a wide range of circumstances [6]. However, it should be noted that its derivation does not have a mechanistic basis in population growth modelling [7], but rather in describing human mortality rates [5].

$$\log_{10}(N_t) = N_0 + (N_{max} - N_0)e^{-e^{\frac{\mu_{max} \cdot e \cdot (t_{lag} - t)}{(N_{max} - N_0) \cdot \log(10)} + 1}}$$

Baranyi The Baranyi model is an attempt to formulate a truly mechanistic model for microbial population growth [8, 1], explicitly modelling the internal physiological state and external environmental conditions of the cell. This mechanistic formulation produces a much more flexible and widely applicable model, which can be applied to model population growth curves in time-varying environmental conditions. Here, we use a version of the model in which conditions are homogenous, with the curvature parameter into the exponential phase ν set to μ_{max} , and curvature parameter out of the exponential phase m set at an empirical estimate of 1:

$$\log_{10}(N_t) = N_{max} + \log_{10}\left(\frac{-1 + e^{\mu_{max} \cdot t} + e^{\mu_{max} \cdot t_{lag}}}{-1 + e^{\mu_{max} \cdot t} + e^{\mu_{max} \cdot t_{lag}} \cdot 10^{N_{max} - N_0}}\right)$$

Phenomenological models

As controls, I decided to fit quadratic and cubic polynomials to the data:

Quadratic $N_t = at^2 + bt + c$

Cubic $N_t = at^3 + bt^2 + ct + d$

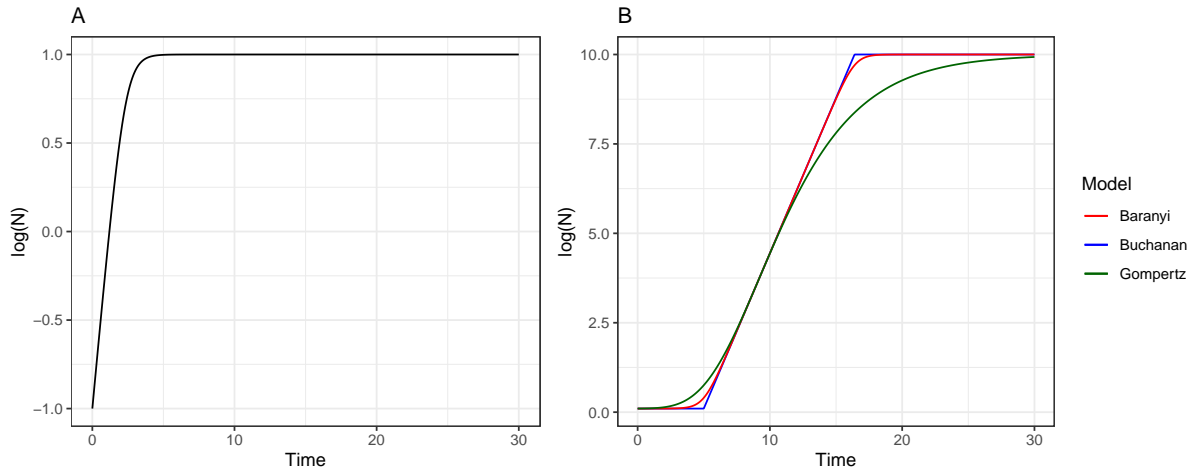


Figure 3: Demonstration of the shape of the 4 mechanistic models, plotted in log space. The logistic model (A) immediately starts growing at its maximum growth rate, whilst the three four-parameter models (B) have growth phases that transition into the exponential phase

Methods

Data

Data sources The provided microbial growth data contained 305 time series, digitised from graphs in 10 papers from the microbial growth literature. The data set contained time series from 47 different species, 18 media, and 17 temperatures, with population size measured in 4 different units. However, the data from one citation [9] clearly had incorrect time values, with times extending to 13,000 hours. I removed the data from this citation as there was not a clear correct transformation to produce the correct times, leaving 287 time series for analysis.

Log transformation As explained below, I decided to fit the model to $\log_{10}(N)$ as well as to linear data. In 5/287 data sets, there were negative population counts due to numerical error in growth curve digitization. One value of approximately -668 was removed, whilst other negative values were greater than -10^{-2} - in these cases, I transformed increased population sizes for that time series by the minimum population size in the time series, discarding the minimum value of 0 before taking the logarithm.

Time transformation I transformed each time series such that the first data point was at $t = 0$ when fitting the four-parameter models, as model fits were clearly suboptimal without this transformation. Since t_{lag} directly transforms the curve along the t-axis, t_{lag} could be adjusted back to obtain model parameters which fit the untransformed time series.

Model Fitting

When fitting the models, I used a least squares approach, both to \log_{10} transformed data, and to the original linear data (Figure S1). This required exponentiating the four-parameter models to predict linear population size, and similarly taking the logarithm of the logistic model to predict logarithmic population size. As such, the parameters of the four-parameter models remained in logarithmic space, and the parameters of the Logistic model in linear space.

Due to the advantages presented in Results, I decided to analyse parameter estimates from the log-fitted models (although linear fit results are presented in the Supplementary Figures), also choosing not to linearise residuals as this resulted in non-sensical fits (demonstrated in Figure S1). These fits can be explained by the fact that, in general:

$$\log(f(x)) \neq f(\log(x))$$

This means minimising errors in logarithmic space does not result in minimised errors in linear space, so evaluating fit in linear space would not capture the best possible fit for each model, (Figure S1).

Fitting linear models

The two polynomial phenomenological models are linear in their parameters, so were fitted using Ordinary Least Squares with the `lm()` function in R [10].

Fitting non-linear models

The mechanistic models are non-linear, so must be fitted using an optimisation approach. The `nlsLM` function in the `minpack.lm` [11] package uses the Levenberg-Marquardt algorithm to optimise initial values to determine best fitting parameter values for a specified model. If initial values are far from the optimal parameters, it is likely that the optimisation will not converge, or will converge to very suboptimal parameters. It is therefore important to estimate reasonable starting values.

Estimating parameter values N_{\min} and N_{\max} were set as the minimum and maximum observed population size respectively. μ_{\max} and t_{lag} estimates were determined using a rolling regression (Figure S2), which computed the regression line for every possible set of 5 adjacent

data points. The regression line with the greatest gradient was selected, with the gradient taken as an estimate of μ_{max} , and the t value at the intercept between this selected regression line and the line $N = \min(N)$ taken as the t_{lag} estimate. This was repeated in both logarithmic and linear space to provide parameter values for Logistic and four-parameter models.

Multiple fit attempts Despite well chosen initial values, models often fail to converge, or converge to poorly fitting parameter values. In order to increase the proportion of models which fit, and to ensure that I find parameter values nearer the global optimum, I used a grid search to explore a range of initial parameter values. For each parameter, I choose linearly spaced samples from a range surrounding the estimate of that parameter value, and fitted a model for every combination of these parameter values. For example, if I take 3 samples for each of N_0 and N_{max} , and 5 for t_{lag} and μ_{max} , I fit $3 \times 3 \times 5 \times 5 = 225$ models. From the model fits that converge, I select the model with the smallest RSS. If none of the models converge, I randomly sample parameter values with normally distributed noise around my parameter estimates until a model fits, or until there have been 1,000 failed attempts.

Using this method, I was able to fit all 287 data sets for the Logistic, Gompertz, and Baranyi models in both linear space and logarithmic space. In linear space, there was one failed fit for the Buchanan model, and in log space there were two failed fits for the Buchanan model.

Filtering

Some data sets show no pattern of population growth, or have too few points in the growth phase to constrain the μ_{max} and t_{lag} parameters. The 64 data sets with the lowest maximum R^2 value were visualised, of which 20 with no meaningful growth pattern were removed (Figure S3). For parameter estimate analysis, I further filtered out datasets with fewer than 2 points in the growth phase (Figure S4) by counting the number of points with a time value within the predicted exponential phase, which was determined using Akaike weighted parameter values. This removed a further 57 data sets, leaving 210 data sets.

Model Selection

Model fit was assessed using AIC [12] and BIC [13]. With the assumption of normally distributed errors, and simplification to remove constants, these are:

$$AIC = 2k + n \log(RSS)$$

$$BIC = n \log \frac{RSS}{n} + k \log n$$

Both AIC and BIC balance fit against model complexity, allowing direct comparison of fit for models with differing numbers of parameters. However, BIC has the disadvantage of making a number of unrealistic assumptions about the true model: that it exists, is in the model set, and has equal prior probability to other models.

As these are dependent on the number of parameters of the model, they can be used for direct comparison of fit for models with differing numbers of parameters. For each data set, the model with the lowest AIC/BIC value was selected as the best fit model. Additionally, the fact that AIC is based in Kullback–Leibler information theory means that standardised model likelihoods - Akaike weights - can be calculated to naturally compare model support and allow model averaging.

Parameter Estimation

Parameter estimation was performed using Akaike weights, which produce a single parameter estimate from a family of models [14], weighted by the model's relative likelihood. With AIC_{best} being the lowest observed AIC values for a dataset, Δ_{AIC} values are calculated with:

$$\Delta_i = AIC_i - AIC_{best}$$

The Akaike weight for the i^{th} model is then:

$$w_i = \frac{e^{-\frac{1}{2}\Delta_i}}{\sum_{r=1}^R e^{-\frac{1}{2}\Delta_r}}$$

The Akaike weighted parameter estimate for θ , with weights w_i and individual model parameter estimates θ_i is:

$$\theta_w = \sum_{i=1}^{i=I} \theta_i * w_i$$

Statistical Analysis

I ran linear regressions of $\Delta_{\text{Logistic - Gompertz}}$ against temperature, and $\Delta_{\text{Logistic - Gompertz}}$ against proportion of points in the lag phase, using `lm()` in R [10]. Their diagnostic plots were examined to confirm the suitability of a linear model.

I ran a mixed effects model of μ_{max} against temperature, and of t_{lag} against temperature, with species and medium used as random effects in each case. Again, diagnostic plots were used to confirm that residuals were homogeneous across the fitted values.

I ran a paired t-tests for μ_{lag} and t_{lag} between the each pair of four-parameter models to check for systematic differences in parameter estimates. I check for normality of differences by plotting histograms of these differences.

Computing Tools

R 3.6.1 [10] was used for initial data preparation script, for model fitting, for analysis, and for figure preparation. I chose R for these tasks because the native data structures are naturally suited to this type of data. The native support of vectorised operations also makes the code cleaner than it would be in Python. I chose to write my multi-start method to fit models over a range of initial parameter values in R because of the robustness and efficiency of non-linear least squares model fitting in the `minpack.lm` package. **minpack.lm** [11] was used for non-linear least squares model fitting using the more robust Levenberg-Marquardt algorithm. **ggplot2** [15] was used to create high quality graphics. **dplyr** [16] was used for some data wrangling tasks. **lme4** [17] was used to fit linear mixed models. **toOrdinal** [18] was used for creating ordinal numbers for annotation.

Python 3.7.4 [19] was used for calculation of initial values. **pandas** [20] was used for dataframes in Python. **matplotlib** [21] was used to create graphics in Python. **SciPy** [22] was used to fit linear regressions for the rolling regression. **NumPy** [22] was used for efficient arrays.

bash 3.2 was used to write scripts tying other elements of the project together.

L^AT_EX was used for compilation of write up.

Results

Model fitting is highly successful

The method of taking the best fitting model from a range of initial parameter values was very effective, with all models converging to every time series when those with no discernible growth pattern were removed, and only 3 unsuccessful fits otherwise (2293 / 2296 successful non-linear fits overall). The model fits were visually inspected, and found to describe the data well (Figure 4 for logarithmic space fits, Figure S5 for linear space fits). For the reasons expanded on in the Discussion, I present the results obtained from fitting models in logarithmic space first, before examining the results from linear space fits at the end of this section.

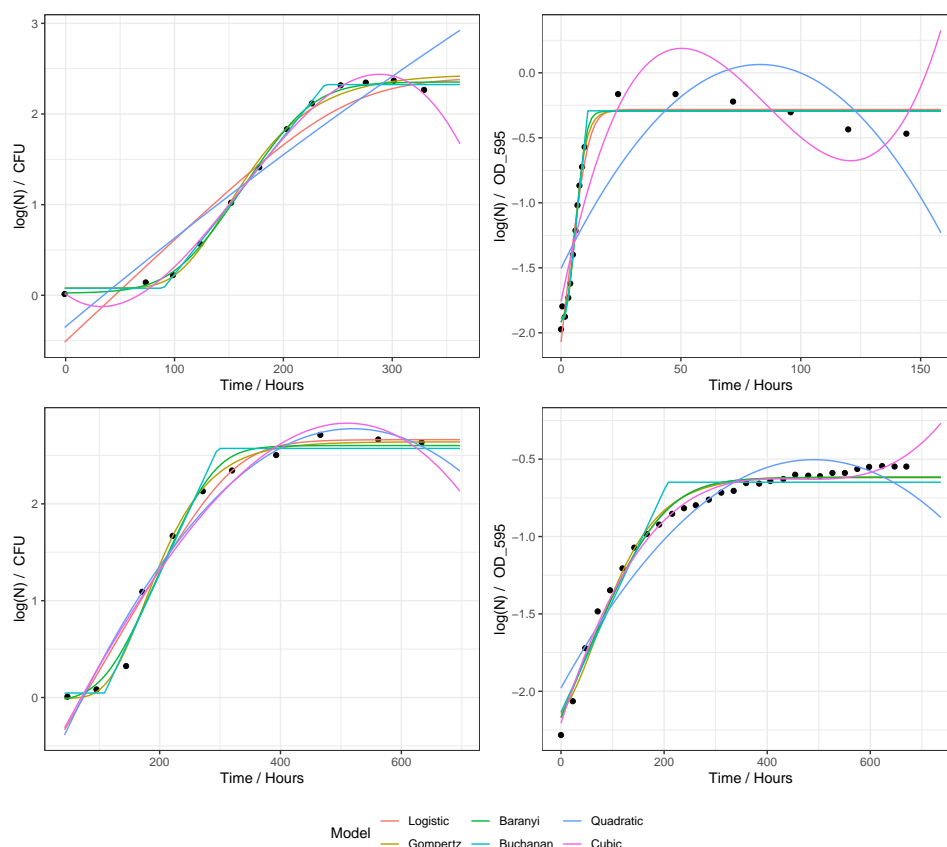


Figure 4: Demonstration of model fits in log space. Out of 287 data sets, all fits were successful apart from two fits of the Buchanan model, to datasets which were removed anyway.

Gompertz and Baranyi are best supported

The Gompertz model and Baranyi model are both well supported by the data. The Gompertz model most frequently has the lowest AIC (Figure 5 A), and has the highest mean Akaike weight (Figure ?? and Table 1). However, the Baranyi model has a lower mean ranking (Figure 5 C),

189 and lower mean AICs and BICs. (Table 1). It appears that the Baranyi model is a consistent
190 good fit to the data, achieving the lowest mean AIC despite relatively rarely being the best
191 fitting model - as evidenced by it placing second for nearly half of the datasets (Figure 5 B).

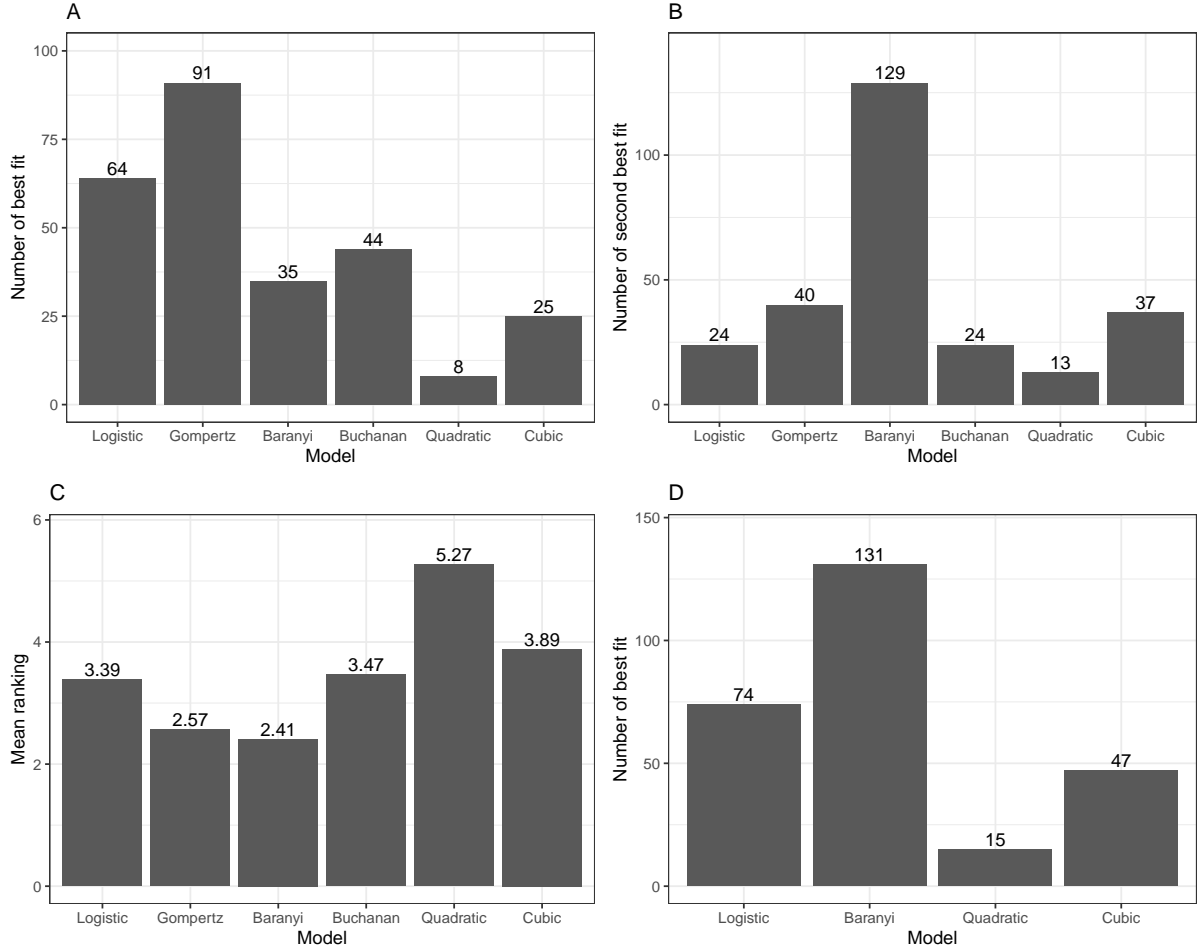


Figure 5: Plots showing the rankings of models in log space. (A) shows the number of times each model is the best fitting model for a dataset. (B) shows the number of times each model is the second best fitting model for a dataset. (C) shows the mean ranking for each model. (D) shows the number of times each model is the best fitting model for a dataset if we do not consider the Gompertz and Buchanan models.

	Logistic	Gompertz	Baranyi	Buchanan	Quadratic	Cubic
AIC	-58.5	-66.2	-66.3	-61.1	-43.7	-53.2
BIC	-57.0	-64.1	-64.2	-59.1	-42.2	-51.1
A_w	0.162	0.316	0.218	0.156	0.0319	0.116

Table 1: Mean AIC, BIC, and A_w values for each model, fitted in logarithmic space. Lower AIC represents greater relative support, whilst higher A_w represents greater relative likelihood.

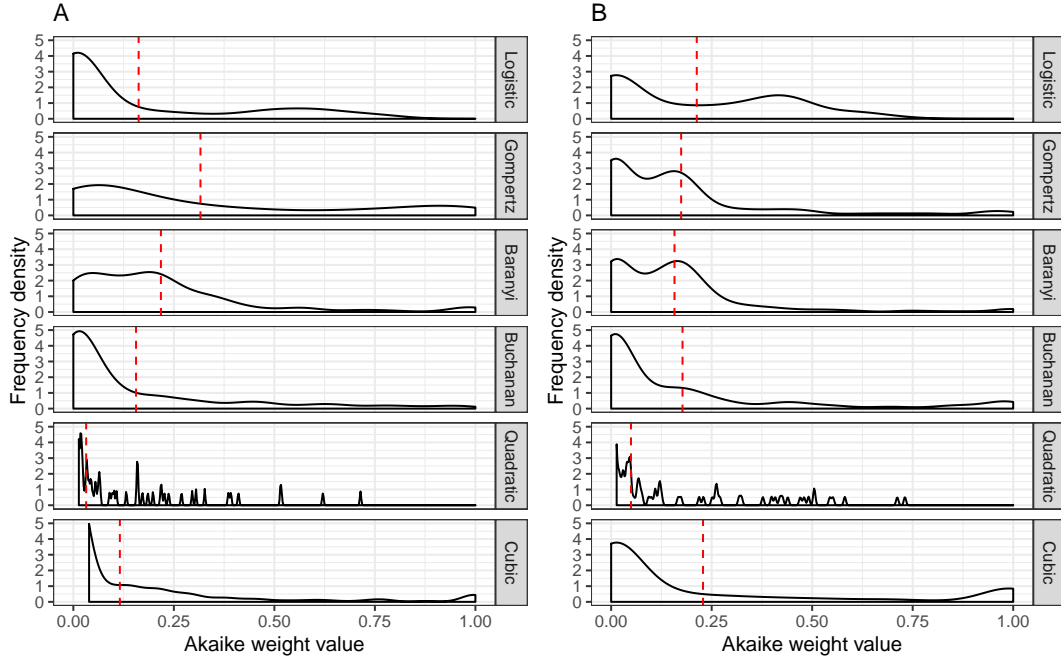


Figure 6: Gaussian Kernel Density Estimates for Akaike weights in logarithmic (A) and linear (B) space for each model.

Logistic fit is poor when t_{lag} is high

It is notable that the Logistic model is the best model second most frequently, but that its mean AIC and BIC scores are only fourth best overall - which implies great variability in its performance. In fact, we can see this variability in its Akaike weight distribution, with high probability density near zero but a relative high density in the 0.5 to 0.75 interval (Figure ??). This can be explained by the absence of a lag phase in the model: the Logistic model fits poorly when a lag phase is present, but fits well when it is absent without incurring the AIC/BIC penalty for a fourth parameter. This is evident in the increasing proportion of points in the lag phase as the ranking of the logistic model gets worse (Figure 7 A). More explicitly, we can see that the difference in AIC between the Logistic and Gompertz model increases with proportion of data points in the lag phase (Figure 7 B). This relationship is strongly significant, with a linear regression estimating an increase in $\Delta_{\text{Logistic} - \text{Gompertz}}$ of 6.9 for every increase in lag phase proportion of 0.1, with a standard error of 0.47.

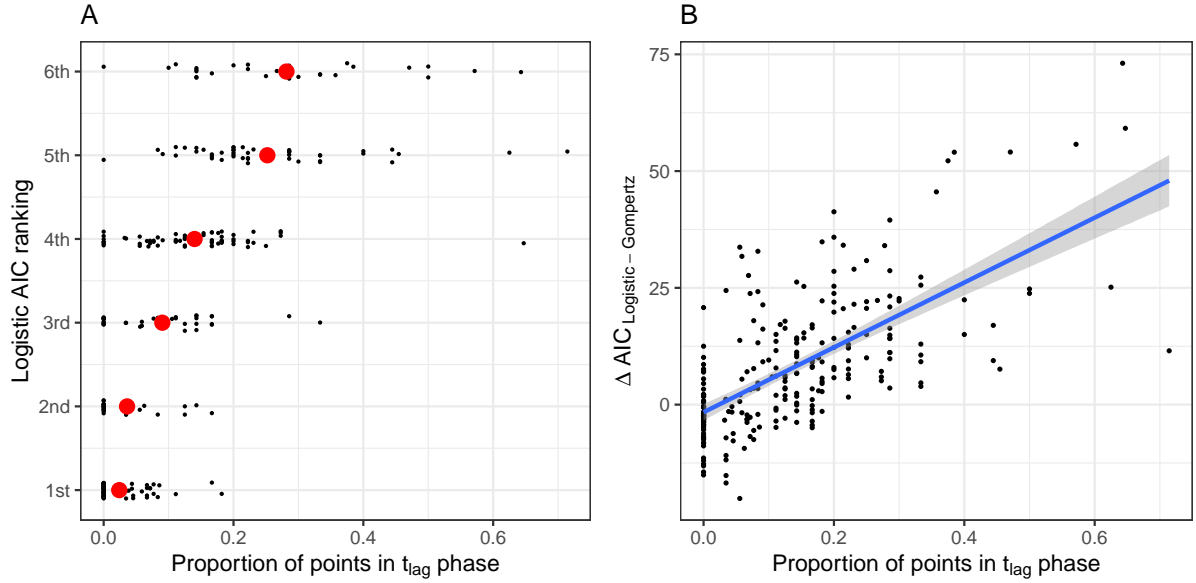


Figure 7: (A) is a plot of proportion of points in the lag phase, against the ranking of the Logistic model by AIC. We see that an increasing proportion of points in the lag phase is associated with a poorer ranking - with the mean proportion for each ranking shown by the red circle. (B) shows the difference in AIC score between the Gompertz and Logistic model against this proportion, with a statistically significant slope observed.

Temperature affects μ_{max} and t_{lag}

I ran a linear mixed model to determine the effect of temperature on μ_{max} and t_{lag} across the entire data set, including species and medium as random effects. Temperature was included as a categorical variable as it is unlikely that the response to temperature is linear across the entire temperature range. From this, estimates for $\log(\mu_{max})$ and t_{lag} values were made for each temperature, shown in Figure 8. It is clear that the log of the maximum rate of bacterial growth increases with temperature, approximately linearly across much of the temperature range, indicating an Arrhenius type functional response. t_{lag} decreases with temperature, quickly levelling off at around 10°C.

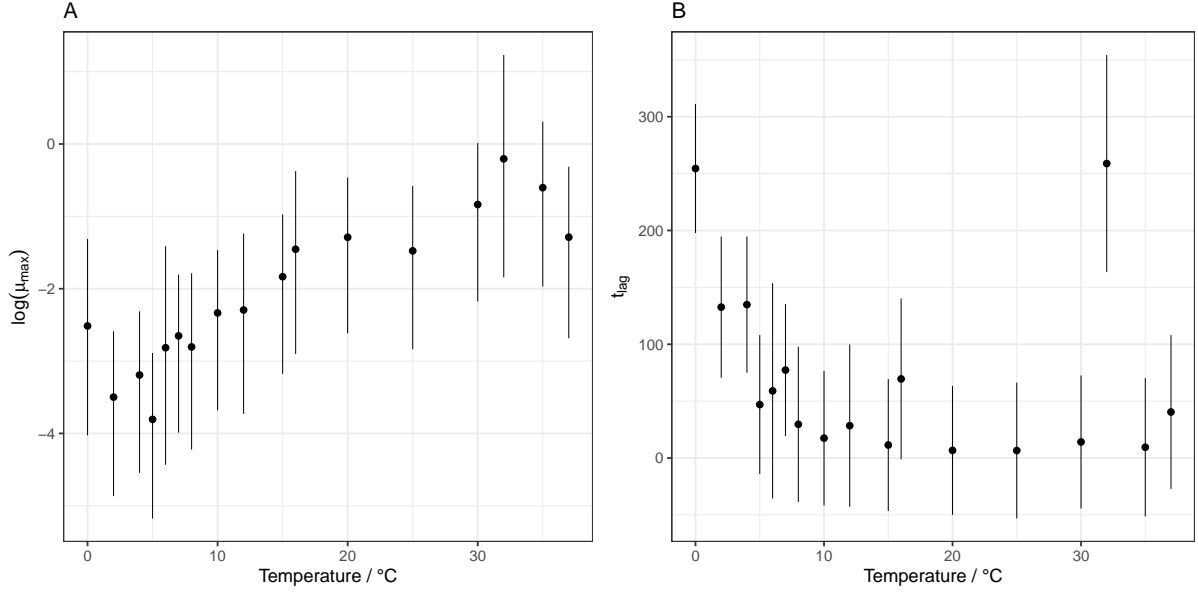


Figure 8: (A) shows the relationship between μ_{max} and temperature, as estimated by a linear mixed model with temperature as a categorical variable. Vertical lines indicate the 95% CI. μ_{max} increases steadily with temperature over the observed range. Similarly, (A) shows the relationship between t_{lag} and temperature. We see that below 10°C, t_{lag} increases rapidly as temperature decreases.

Logistic model fits better at higher temperatures

The decrease in t_{lag} with temperature implies that the Logistic model, which has better fit with smaller t_{lag} , should have better fit with increasing temperature. Indeed, Figure 9 A shows that $\Delta_{\text{Logistic} - \text{Gompertz}}$ decreases with increasing temperature. A linear regression suggests a decrease in $\Delta_{\text{Logistic} - \text{Gompertz}}$ of 3.30 for every 10°C, with standard error of 0.8.

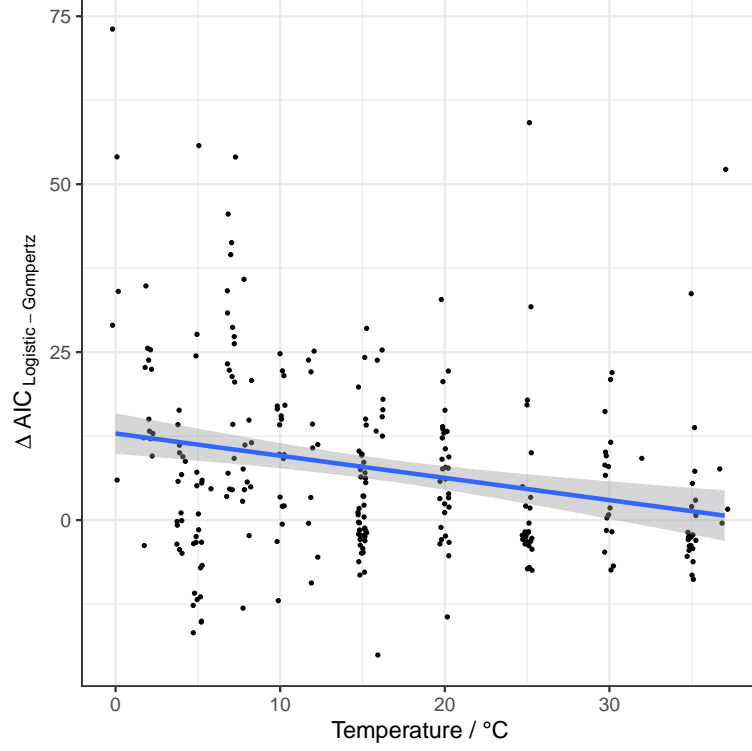


Figure 9: $\Delta_{\text{Logistic} - \text{Gompertz}}$ is seen to decrease by 3.30 for every 10°C , indicating an improvement in the fit of the Logistic model relative to the Gompertz model as temperature increases.

Model choice biases parameter estimates

Tables 2 and 3 show the mean pairwise differences in t_{lag} and μ_{max} for the 4-parameter models. The Buchanan model results in the lowest t_{lag} and lowest μ_{max} values. This is because the model assumes that population growth remains exactly at zero until $t = t_{lag}$, rather than smoothly transitioning to μ_{max} before this intercept. This means that Buchanan model fits must decrease t_{lag} to capture the first part of the growth phase, consequently lowering t_{lag} and μ_{max} values (Figure S6).

	Gompertz	Baranyi	Buchanan
Gompertz		-5.21 = - 9.8%	4.23 = 8.0%
Baranyi			9.44 = 17.8%
Buchanan			

Table 2: Mean pairwise differences in t_{lag} estimates for all models, and as proportions of the Akaike weighted mean t_{lag} value. The value in $(i, j) = t_{lag}^i - t_{lag}^j$. The Buchanan model produces lower t_{lag} estimates than either the Baranyi or the Gompertz models. All values are significant at the 5% level.

	Gompertz	Baranyi	Buchanan
Gompertz		-0.025 = -5.4%	0.11 = 23.4%
Baranyi			0.14 = 29.3%
Buchanan			

Table 3: Mean pairwise differences in μ_{max} estimates for all models, and as proportions of the Akaike weighted mean $t\mu_{max}$ value. The value in $(i, j) = \mu_{max}^i - \mu_{max}^j$. The Buchanan model produces lower μ_{max} estimates than either the Baranyi or the Gompertz models. All values are significant at the 5% level.

Model fitting in linear space fails to resolve the lag phase

As explained in Methods, fitting linearised models to minimise linear RSS reduces the ability of the models to resolve the lag phase. As such, the Logistic model is favoured when fitting in linear space despite its lack of a lag phase. When minimising linear residuals, the Logistic model has the highest mean Akaike weight (Figure ?? B), is most frequently the best model, and has the lowest mean ranking (Figure S7 and Table S1). The weakened penalisation of the lag phase also means that the relationship between Logistic model fit and proportion of points in the lag phase is much weaker (Figure S9), and the relationship with temperature becomes smaller and loses significance (Figure S9).

Discussion

Mechanistic modelling is limited in its resolution between biological hypotheses

The results are unequivocal about the importance of a lag phase for modelling microbial growth in log space, with the three four-parameter having the most support by Akaike weights, AICs, and BICs. Further, the fit of the Logistic model worsening with increasing lag phase duration. However, the difference in mean AIC between the Gompertz and Baranyi model is negligible, so no concrete mechanistic statements can be made - especially given that the Gompertz model is only mechanistic to the extent that it models the three canonical phases of the growth curve. Indeed, it has been argued that mechanistic modelling of this sort cannot adequately discern between biological hypotheses, which should be confirmed by direct observation or experimentation rather than argumentation about the cause of differences in model fit [6]. Indeed, direct observation of the process of microbial growth can also provide us with estimates of parameters such as t_{lag} and μ_{max} which currently have caveats in their interpretation, by directly counting

and timing cell divisions. However, for parameter estimation and prediction in its current form I propose that the Baranyi model is a stronger choice:

1. Consistency: The Baranyi model is a more consistently good fit, achieving a marginally lower AIC despite rarely being the best fitting model. This implies that the Baranyi model can be deployed more broadly and reliably than the Gompertz model, and that parameter estimates are likely to be more consistent.
2. Truly mechanistic: The Baranyi model has the advantage that it was formulated entirely with microbial growth in mind, giving it a more mechanistic basis in microbial growth modelling, and therefore more interpretable parameters. The full Baranyi model also has extra curvature parameters ν and m , which alter the transition into and out of the exponential phase. As more data becomes available, these parameters can be estimated to form a more detailed understanding of the mechanisms behind microbial growth. Similarly, the Baranyi model's capacity to make predictions with non homogenous environmental conditions in its differential equation form makes it a much more flexible tool for growth modelling.

Lowering temperature can significantly impede microbial growth

I find that $\log(\mu_{max})$ increases roughly linearly with temperature, corresponding to an exponential increase in μ_{max} - which in turn describes the rate of exponential growth. Similarly, I find that t_{lag} decreases rapidly with increasing temperature, such that the advantage of lower temperatures is almost entirely lost by 10 °C. Clearly, minimising the temperature of food storage can have a dramatic effect in decreasing microbial load - and conversely, increasing the temperature can result in growth orders of magnitude faster, with perhaps even brief exposure to higher temperatures able to spoil food. It will therefore be important to see how microbial growth patterns respond to temperatures that vary over time, which the Baranyi model will be very helpful in studying.

Parameter biases necessitate caution in meta-analyses

There is a clear hierarchy in the magnitude of parameter estimates from the three models. The Buchanan model produces the lowest t_{lag} and μ_{max} estimates, with the Baranyi producing the highest, and Gompertz producing intermediate values. The magnitude of the differences

between the Buchanan and Baranyi model reach 29% of the Akaike weighted estimate for μ_{max} , and 17% for t_{lag} . These are extremely significant differences, so great care, probably involving reevaluation of raw data sets, must be taken to ensure that parameter estimates from different sources are meaningfully compared, such that these biases are not interpreted as biological effects. A simulation study could be used to determine which model, or combination of models, produces the most reliable parameter estimate. Datasets produced by a variety of simulation mechanisms, with known 'true parameters' could be produced for a range of generative processes, and used to test how robust each of the models is to deviations from their mechanistic assumptions, and to noise in the data.

Fitting in logarithmic space is required to estimate relevant parameters

Fitting in linear space allows the logistic model to become the most supported model. This is a strong remit to recommend that model fitting should be carried out and evaluated in log space, as the improved ranking of the Logistic model demonstrates that fitting in linear space only weakly enforces accurate modelling of the lag phase. This is because linear space is concerned with modelling absolute rather than relative population size, such that mischaracterisation of the early growth phase is not strongly penalised. The expansion of variation at small population sizes in logarithmic space also allows higher confidence estimation of t_{lag} and μ_{max} , as shown by the narrower confidence intervals in Figure 8 relative to Figure S10. , Fitting in logarithmic space is therefore strongly recommended for food preservation work, in which early growth curve behaviour is centrally important as food is often spoiled before N_{max} is reached [6].

As such, this study has demonstrated that the choice of space and model in microbial growth modelling can have profound effects both on the parameters estimated, and may even even conceal findings. Further work examining the reliability of the various models to violations of their assumptions is needed to devise strategies that can provide the most accurate parameter estimates, which will allow increasingly quantitative approaches to food preservation.

Acknowledgements

I would like to thank Pablo Lechon for the stimulating and productive conversations we had on model fitting. And to Hovig Artinian, thank you for spending the night with me.

Supplementary Material

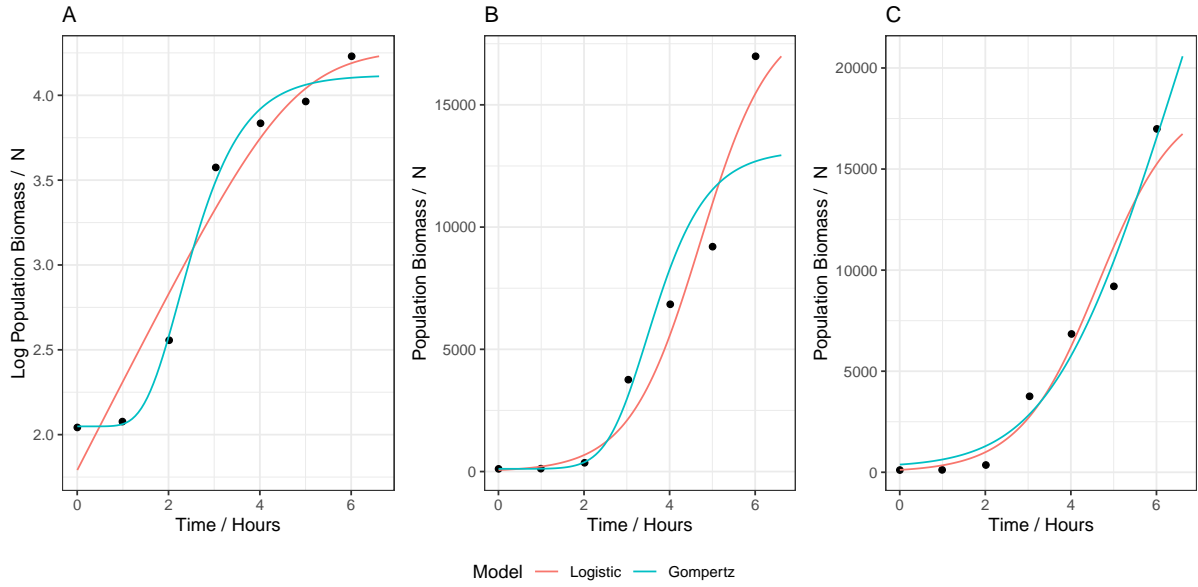


Figure S1: Comparison of: (A) fitting and evaluating modelling in log space. (B) fitting model in log space, evaluating fit in linear space. (C) fitting and evaluating model in linear space. When the models are fit and evaluated in log space, it is clear that the Gompertz model has a better fit. However, if evaluation is done in linear space of these fits, the logistic model appears to have a better fit. This is shown to be invalid, as fitting the models in linear space demonstrates that the Gompertz model can in fact achieve better fits in both spaces.

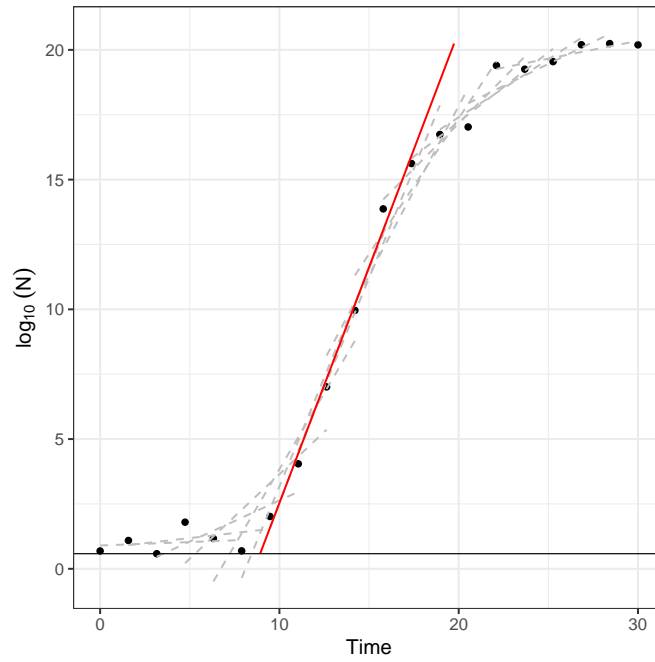
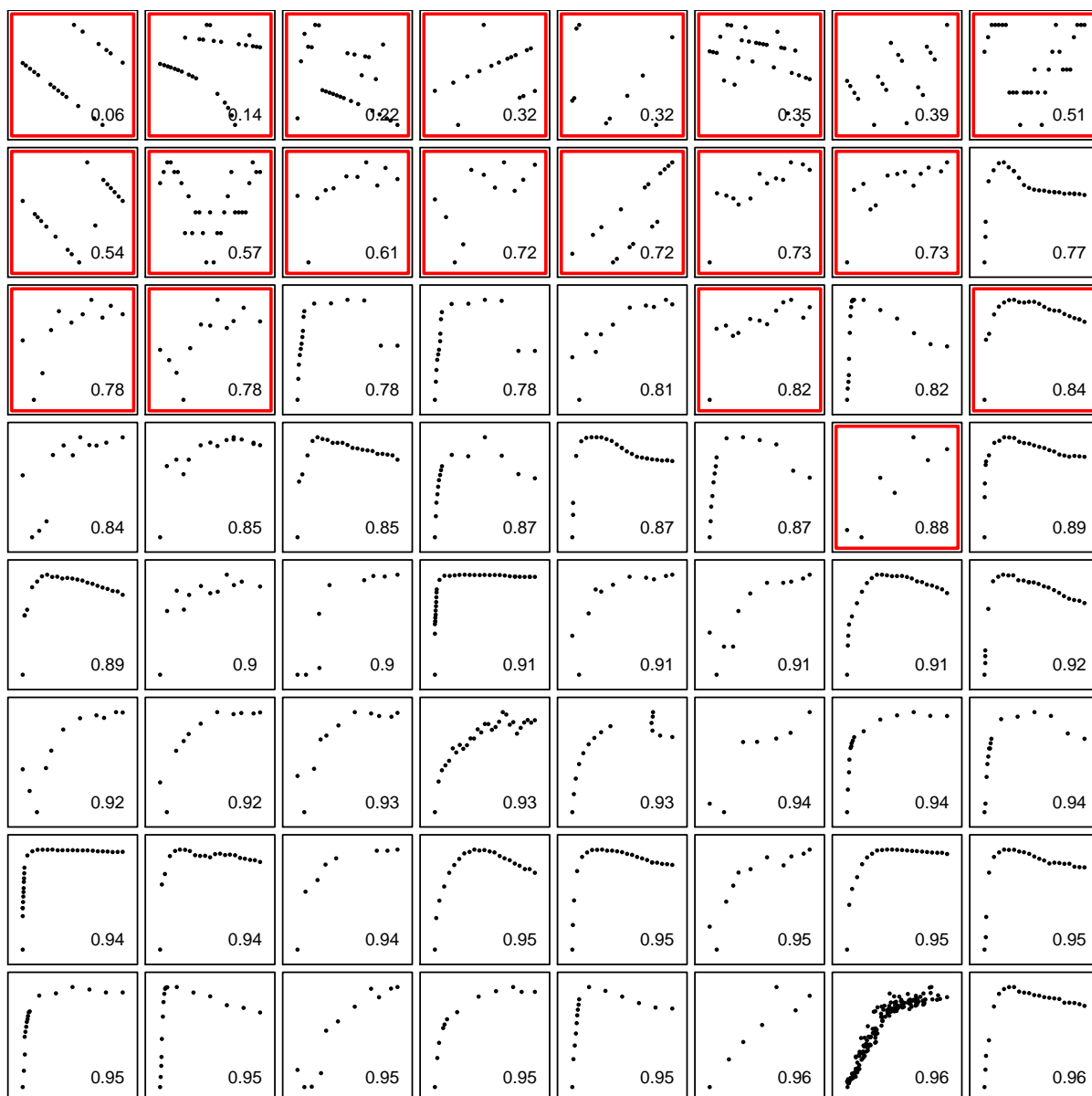


Figure S2: Rolling regression was used to produce starting parameters. The regression line with the greatest gradient was selected, with its gradient used as μ_{max} , and its intercept with $N = \min(N)$ used as t_{lag}



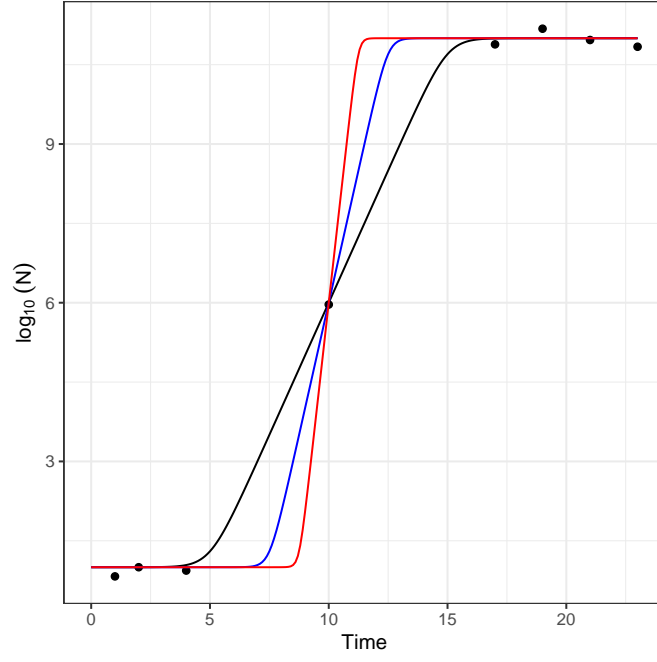


Figure S4: If there are too few points in the growth phase, μ_{max} and t_{lag} will not be well constrained. Here, we can see that there are infinitely many solutions that can fit equally well to the data as there is only one point in the growth phase.

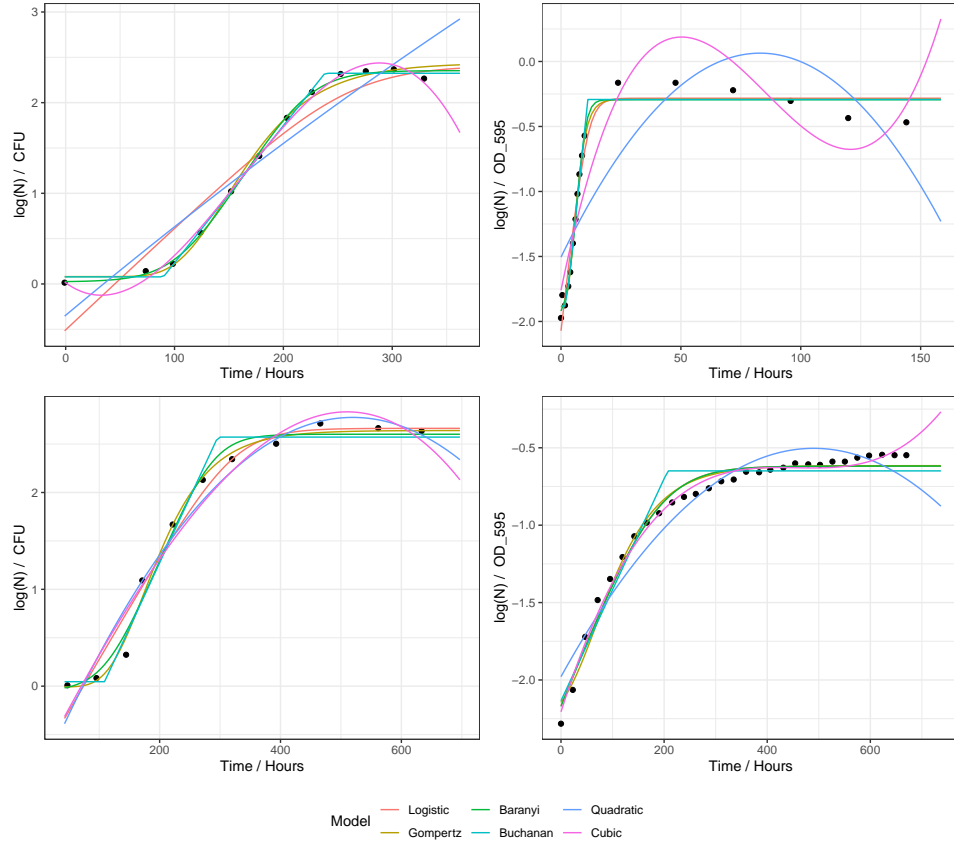


Figure S5: Demonstration of model fits in linear space. Out of 287 data sets, all fits were successful apart from a single fit of the Buchanan model, to a dataset which was removed anyway.

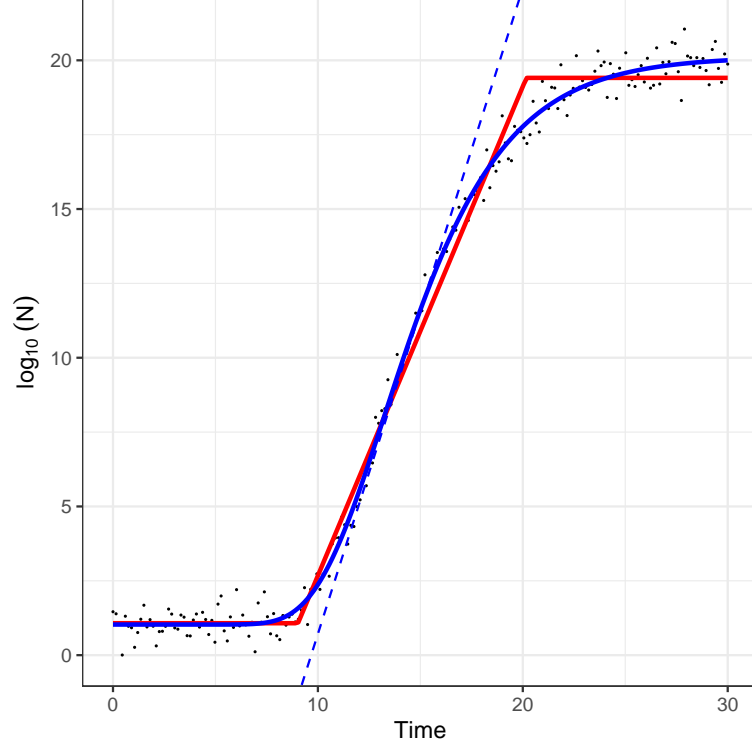


Figure S6: The Buchanan and Gompertz models are fit to the same hypothetical growth data, which results in smaller t_{lag} and μ_{max} estimates from the Buchanan model in order to capture the start of the growth phase. By contrast, there is some growth before $t = t_{lag}$ in the Gompertz model.

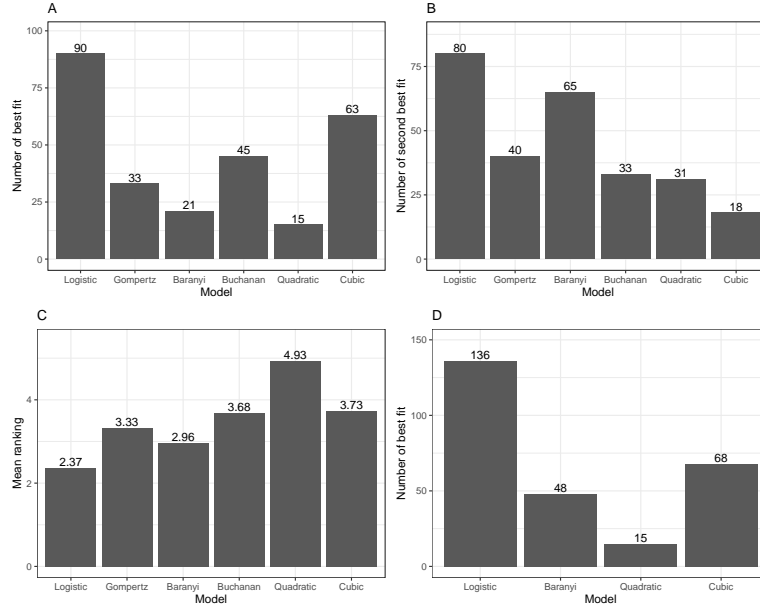


Figure S7: Plots showing rankings of models in linear space. (A) shows the number of times each model is the best fitting model for a dataset. (B) shows the number of times each model is the second best fitting model for a dataset. (C) shows the mean ranking for each model. (D) shows the number of times each model is the best fitting model for a dataset if we do not consider the Gompertz and Buchanan models. In linear space, the logistic model performs better than it does in log space, as less emphasis is put on the lag phase.

	Logistic	Gompertz	Baranyi	Buchanan	Quadratic	Cubic
AIC	62.3	61.8	61.3	65.5	76.3	67.7
BIC	63.8	63.8	63.3	67.5	77.8	69.7
A_w	0.213	0.174	0.158	0.178	0.0493	0.229

Table S1: Mean AIC, BIC, and Akaike weight values for each model fitted in linear space

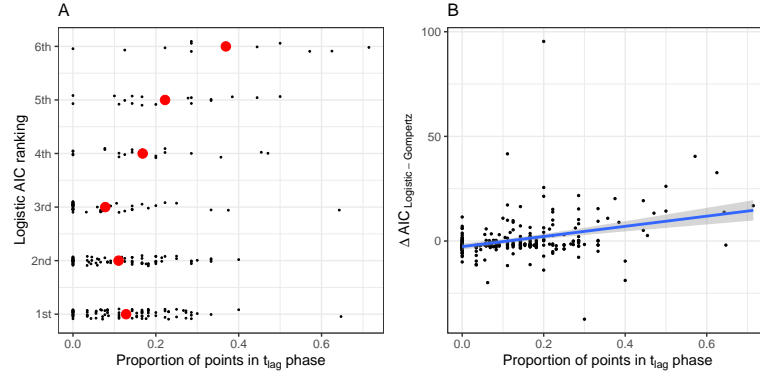


Figure S8: (A) is a plot of proportion of points in the lag phase, against the ranking of the Logistic model by AIC in linear space. (B) shows the difference in AIC score between the Gompertz and Logistic model against this proportion. In both cases, the relationship between Logistic model performance and t_{lag} is much weaker than in log space. The difference in AIC between Logistic and Gompertz decreases by 2.4 for every increase in lag phase proportion by 0.1 (in contrast to 6.9 for log space)

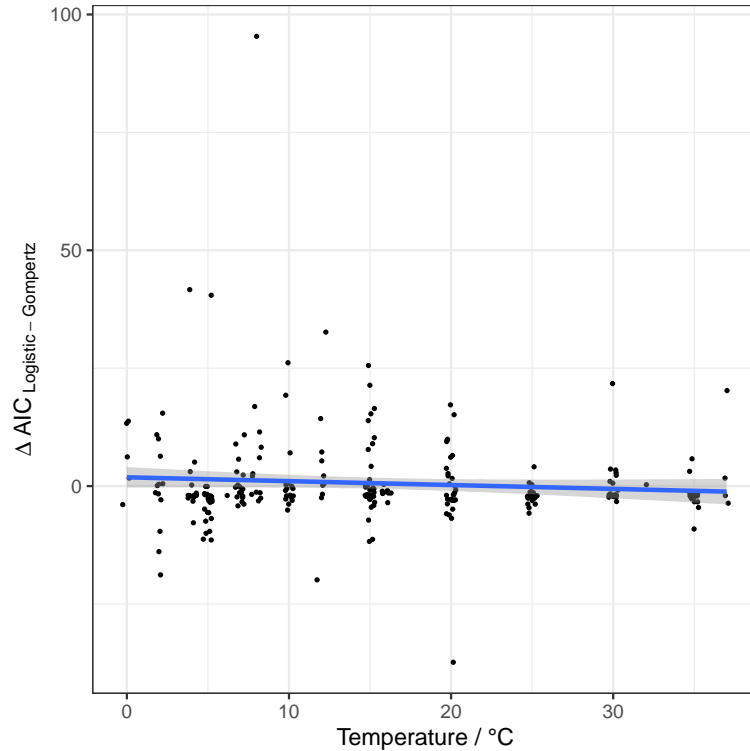


Figure S9: $\Delta_{Logistic - Gompertz}$ no longer has a statistically significant relationship with temperature when models are fit in linear space. This implies that the fit of the Logistic model does not become any weaker relative to the Gompertz model as temperature increases.

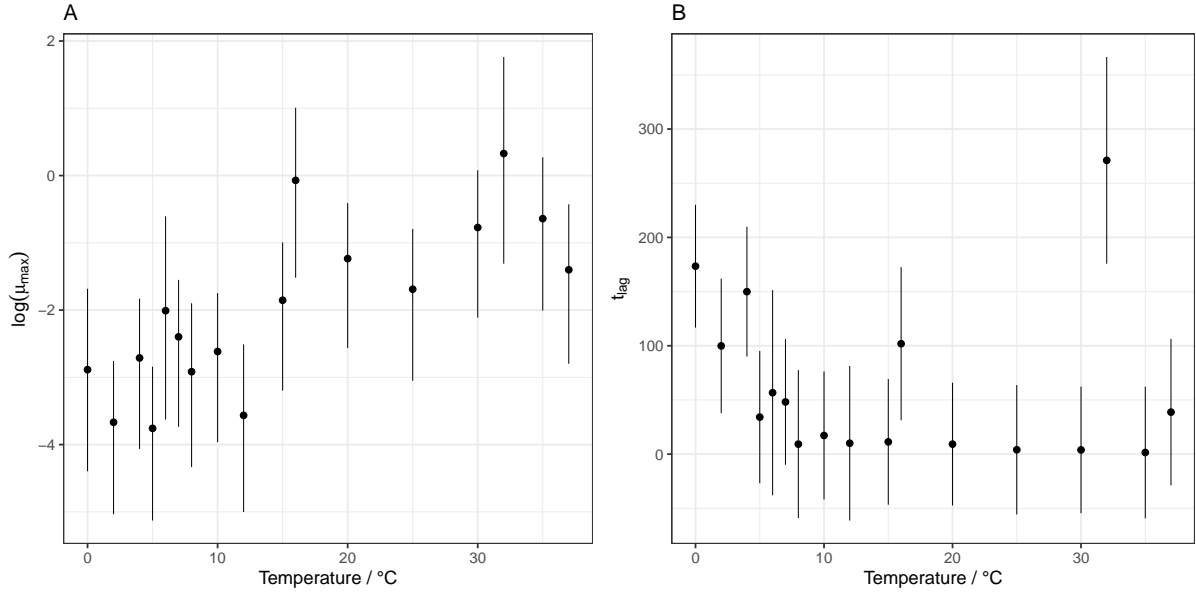


Figure S10: The broad patterns of μ_{max} and t_{lag} are the same when model fitting is in linear space, but the confidence intervals are much broader.

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