Mini Project

Comparing Mathematical Models for Bacterial Growth Curves Fitting

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Yuchen Yang

MRes Computational Method for Ecology and Evolution

Department of Life Sciences

Imperial College London

yuchen.yang19@imperial.ac.uk

1 Abstract

This study compares the Baranyi, Logistic, Gompertz, and the basic Polynomial(Cubic) model's behaviour in a bacterial growth data set using AIC, ΔAIC , and AICweight (and reasons to exclude other indicators). The result indicates that the **Gompertz** model is the best model for fitting this general data set(regarded as best fit for a total of **182** times among other models, taking up **62.5%** of the total data set groups), despite the fact that there are arguments saying the more mechanistic Baranyi model would be better. The study also used a subset of **Tetraselmis tetrahele** in the data set to examine temperature's effect on model fitting. The result from the Kruskal-Wallis Test indicates that models have shown a significant differences (Baranyi's p-value=0.001679, Logistic's p-value=0.01684, Gompertz's p-value=0.03801, Polynomial's p-value=0.01672) in their AICweight across different temperature groups. Finally, this study compared models' behaviour on the death phase of a growth curve and explore the reasons for those differences in performances.

13 1 Introduction

Mathematical models are used to understand and predict the growth of the bacterial population for quite a long time. Most bacterial growths in batch culture follow a distinct set of phases - lag 15 phase, exponential phase and stationary phase, which would perfectly fit a sigmoid shape. 16 Gompertz model (Gompertz 1825) and the Logistic model (Verhulst 1838) are the most frequently 17 used sigmoid models fitted to growth (Tjørve and Tjørve 2017). However, most of the time nowadays, taking measurement errors and the expected growth kinetics into consideration, food microbiology 19 researches would use the logarithm of cell concentrations as raw data. Some of the papers are using 20 those classical sigmoid growth models as above, but since the population is in the logarithm of the 21 cell concentrations, the results are not as good(Baranyi 2011). Baranyi also pointed out that the 22 reason those two models, while being very popular and useful, has their limitations, especially in 23 fitting bacterial data, is that the original model's curve does not remain sigmoid shape, but rather a monotone shape in log-space. Hence they could fail to capture the lag phases, and so that the use of

such models are limited (Baranyi 2011).

It has been argued that the empirical models such as modified Gompertz and Logistic models were not as preferred as the more mechanistic models such as the Baranyi model even with the modifications for 28 it to work better in log-space (Juneja et al. 2009). To compare and evaluate similarities and differences 29 between the models behaviour on a general bacterial database, this study has chosen the Baranyi 30 (Baranyi, Roberts, and McClure 1993), the Logistic, the Gompertz (both using the log-transformed 31 formulation of Zwietering) (Zwietering et al. 1990) and the basic Polynomial (Cubic) model, from the 32 most mechanistic to the least, to better understand the behavioural differences. 33 Aside from understanding how different models perform in a fitting practice, it is also interesting to understand what are the co-variates' role in this. Will a model's performance differ across different co-variate groups? Understanding this would be helpful for researchers to find appropriate models 36 to use in future analysis based on the condition of the data set. the Polynomial model is introduced 37 mainly to benchmark models' behaviour on a fourth stage that has not been very often mentioned death phase. Since A Cubic model have a declined phase at the end, it may be quite good at capturing 39 decline in population size after some maximum value (the carrying capacity) has been reached (the 40 "death phase" of population growth) (Buchanan 1918), it is also interesting to see how different models are behaving in that period. 42

- To recap on the objectives:
- This study would first compare the above-mentioned models' behaviour for a general bacterial data set, and determine which one fits the best.
- This study would help to understand different co-variates' (specifically focus on temperature)

 effect on model fitting performances.
- This study will touch base on the death phase performances.

Data 2

50 2.1 Basic Features

51 The data set consists of data from multiple pieces of research, which contains measurements of

52 change in biomass or number of cells of microbes over time and were collected through lab ex-

periments across the world. the original data can be found in the Data directory with the name

LogisticGrowthData.csv, detailed column metadata could be seen in LogisticGrowthMetaData.csv.

The two main fields of interest are called PopBio(abundance), and Time.

56 The data set consists of 4387 rows of data from 10 pieces of research, covering a total of 17 tempera-

ture groups from 0 to $37^{\circ}C$, 45 species, and 18 mediums. At the first glance, the min of the Time is

-25.2632, the min of the PopBio is -668.284, suggesting further data wrangling is required, but there

59 are no missing values in the data.

60 2.2 Pre-Processing

61 2.2.1 Re-Calibrate

62 The data set is grouped into 305 groups based on its Temp, Species, Medium, Citation, and Rep to

 $_{63}$ better facilitate the study. The group sample size is rather small, with a mean of 14.36066 and a

median of 12. Furthermore, each group's starting point $O(T_{0}, N_{0})$ is used as a reference to

move the whole group's data to the origin to get rid of any negative values and make better sense.

66 2.2.2 Logarithm

67 Since all the model we have used is either designed for the log-space or being log-transformed. To

better unify the results and deal with measurement errors, etc., a new column called logPopBio, which

is the natural log of the original PopBio, is added and used for all future analysis.

70 2.2.3 Death Phase

To better understand the fitting performance of all models during the "death phase", which is a decline

of the population after reaching the maximum population, a column named deathPhase is added. By

default the value is 0, and when $\frac{N_{max}-N_{tlast}}{N_{max}-N_{min}} > 20\%$, which would indicate a decline after hitting the maximum point in population density, the value is set to 1, which indicates there might be a death phase in the graph.

76 3 Methods

77 3.1 Models

For this particular study, I have used transformed/modified models to make sure they all have four parameters to fit at the end and should work well within a log-space. Take the **Baranyi** model as an example, the original model is defined as:

$$N_t = N_0 + r_{max} A_t - \ln\left(1 + \frac{e^{r_{max}} A_t - 1}{e^{N_{max} - N_0}}\right) \tag{1}$$

Where A_t is defined as:

$$A_t = t + \frac{1}{r_{max}} \cdot ln\left(\frac{e^{-r_{max}t} + h_0}{1 + h_0}\right) \tag{2}$$

And since t_{lag} can be obtained by using:

$$t_{lag} = \frac{ln(1 + \frac{1}{h_0})}{r_{max}} \tag{3}$$

Although the **Baranyi** model introduced a new parameter h_0 , according to 3, is possible to then re-write the whole model using only N_0 , N_{max} , r_{max} , and t_{lag} .

The final four models used for this study are listed here:

86 3.1.1 Baranyi

$$y = N_{max} + \ln\left(\frac{-1 + e^{r_{max} \cdot t_{lag}} + e^{r_{max} \cdot t}}{e^{r_{max} \cdot t} - 1 + e^{r_{max} \cdot t_{lag}} \cdot e^{N_{max} - N_0}}\right) \tag{4}$$

3.1.2 Logistic

$$y = N_0 + \frac{N_{max} - N_0}{1 + e^{\frac{4 \cdot r_{max} \cdot \left(t_{lag} - t\right)}{N_{max} - N_0} + 2}}$$
 (5)

88 3.1.3 Gompertz

$$y = (N_{max} - N_0) \cdot e^{-e^{\frac{r_{max} \cdot e \cdot \left(t_{lag} - t\right)}{N_{max} - N_0} + 1}} + N_0 \tag{6}$$

9 3.1.4 Polynomial(Cubic)

$$y = C_0 + C_1 \cdot t + C_2 \cdot t^2 + C_3 \cdot t^3 \tag{7}$$

In model 4, 5, and 6, the four parameters to fit are N_0 , N_{max} , r_{max} , and t_{lag} , where N_0 is initial cell culture (Population) density, N_{max} is maximum population density (aka "carrying capacity"), r_{max} is the maximum growth rate (the tangent to the inflection point), and t_{lag} is the x-axis intercept to this tangent. Parameters like A(in Gompertz) and h_0 (in Baranyi) are obtain and transformed using the four parameters. Meanwhile model 7 has four parameters with no biological meaning at all.

95 3.2 Starting Value

For this fitting practice, all starting values for model (4) has been set to 1, since the parameters are purely mathematical, and not biologically relevant, and it would fit regardless. For the rest of the models, methods for obtaining estimates for starting values are described in table 1:

Parameter	Method	Equation		
N_0	Get the min of N(PopBio)	$N_0 = min(PopBio)$		
N_{max}	Get the max of N(PopBio)	$N_{max} = max(PopBio)$		
r_{max}	Get the max of slopes for all adjacent	$r_{max} = max(\frac{\Delta N}{\Delta t})$		
	points(approximate tangent to the inflec-			
	tion point)			
t_{lag}	Get the $y = N_0$ intercept to the line with	$t_{lag} = t_{rmax} - \frac{N_{rmax}}{r_{max}} + N_0$		
	r_{max}			

Table 1: Overview on getting starting values

99 3.3 Evaluation

Coefficient of determination R^2 is used to determine converged but bad fit, and will be used to exclude
data set groups that do not make sense. Since the package used in python did not return R^2 , it is
calculated by using the already available SS_{res} and calculated SS_{tot} using values of the data set group

103 and model fit estimate.

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AIC and ΔAIC are used to populate the grading system for models. ΔAIC is calculated as 104 $\Delta AIC_i = AIC_i - AIC_{min}$. Most of the time, it is recommend not to use AIC without the bias 105 correction term, which is $AICc = AIC + \frac{2k^2 + 2k}{n - k - 1}$, unless $\frac{n}{k} < 40$, where k is the total number of pa-106 rameters of a model (degree of freedom for a model), and n is the sample size of the working data set 107 group (Burnham and Anderson 2002). While in this study, all of the models have the same parameter 108 count of 4, making AICc and AIC essentially the same. 109 BIC is calculated as $\ln(n)k - 2\ln(\hat{L})$ whereas $AIC = 2k - 2\ln(\hat{L})$, \hat{L} represents the maximized value 110 of the likelihood function of the model. Making the differences between AIC and BIC purely based 111 on the penalty part. For AIC and BIC to have a difference, $\ln n$ needs to be than 2, meaning the 112 sample size should at least be larger than 8. Since the average sample size of data set groups is 113 $n_{mean} = 14.36066$ as mentioned, there should be a difference between the two standards. Essentially 114 speaking, BIC is different because it prefers less complicated models when the sample size is larger. 115 However, all models used in this study have 4 parameters (n=4), which means they have the same 116 degrees of freedom, and that renders the AIC and BIC less different than each other. δBIC would 117 still be used in the grading process, but it is likely to give the same result. P are given based on each group's fitting results. In this particular study, 2 system would be used, 119 one using AIC, the other using BIC. Points will be given to a model using $\Delta AIC < 2$ and/or 120 $\Delta BIC < 2$, where 2 is set to be the threshold for a substantial difference in performance for both 121 indicators. (Burnham and Anderson 2004; Kass and Raftery 1995). 122 Since this study is going to compare model performance across different co-variate groups, and the 123 size of different co-variate groups S_{group} various. In order to be able to compare, a relative indicator is needed other than the P_{model} for the comparison, here we use Akaike weights $w_i(AIC)$ (Wagenmakers 125 and Farrell 2004), calculated as in equation 8 126

 $w_i(AIC) = \frac{e^{-0.5 \cdot \Delta_i AIC}}{\sum_{K=1}^K e^{-0.5 \cdot \Delta_k AIC}}$ (8)

128 3.4 Computing Tools

This project used *Python* for data preparation and fitting, *R* for plotting all graphs and doing analysis, *LaTeX* for report writing, and *Bash* to glue scripts together and make it automated and reproducible.

Packages used in *Python* are *pandas*, *math*, *numpy*, and *lmfit*. *ggplot*2 and *reshape*2 is used in *R*for plotting.

133 4 Result

4.1 Post-Processing

135 **4.1.1** Bad Fit

Using values obtained according to section 3.2, 2 of all fits returned a result.success == False, which 136 would indicate unsuccessful fits due to not being able to converge. Upon further investigation, this is most certainly caused by having inappropriate initial value (larger than it should) for r_{max} . In order 138 to get better initial values for the fit, a while loop is introduced for a fit that is failing to converge, 139 where it randomly samples a uniform distribution of $(\frac{r_{max}}{3}, r_{max})$ to be the new r_{max} . 140 There are other cases where there is an exception for valueError and the fit is aborted. Based on the 141 error message, it is caused by having negative values in the log() function for the Banranyi model or overflow for exp() for all models. This requires further investigation on either manipulate the raw 143 data more, or set boundaries to parameters, and will not be done in this particular study since the 144 number of cases are small enough to ignore.

146 **4.1.2** Bad Data

A first look at R^2 overview from figure 1 indicates some out-liners in the R^2 distribution for all models. These indicate that although there is a fit for the corresponding data using that model, the goodness of fit is relatively bad(a low value in R^2).

Here, a threshold is set to $R^2 = 0.8$, since this value, as shown in the graph would help to exclude a

maximum number of out-liners without losing too much information.

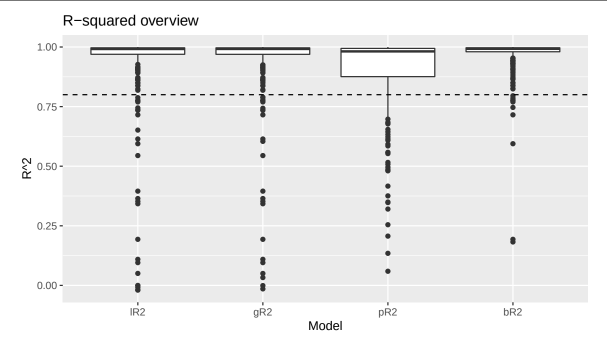


Figure 1: R^2 overview for each model, where lR2 = Logistic, gR2 = Gompertz, pR2 = Polynomial, bR2 = Baranyi

Figure 2 is an example of a data set where either all models' fit is $bad(R^2 < 0.8)$, or some of the model did not fit, and rest of the model indicated a bad fit. Before grading the model, data set groups that are similar to figure 2, are excluded from the final evaluation data. Upon examination, all of these cases happened because the data set group itself is showing a bad shape, as could be seen in figure 2. A total of 14 sets are excluded in this step, leaving the final count of data set groups to be 291. The majority of the data set groups are showing a good value in R^2 , Which also indicates a pretty decent starting value choice.

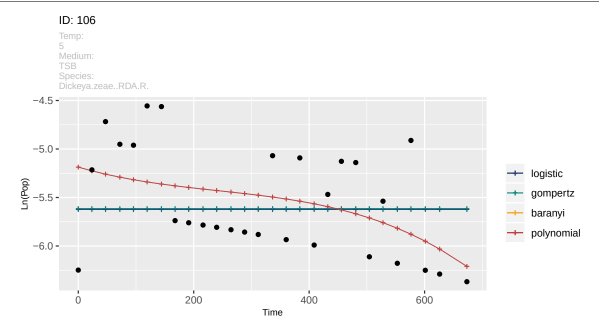


Figure 2: Example of deleted data set, where the Baranyi model didn't fit, and the rest three models showed bad fitting.

9 4.1.3 Grading

This part will take the data set group ID = 92 as an example to better illustrate the grading process.

As can be seen in the evaluation matrix shown in table 2 and graph shown in figure 3 for this specific data group, all four model fit well(all with a $R^2 > 0.91$). And according to the rule for the grading system illustrated in section 8, only the Baranyi model gets 1 point for having a $\Delta AIC < 2$, and all other models would not get a point for all of their ΔAIC is greater than 2. As can be seen, ΔBIC in this particular study is giving the same results(all models have the same parameter count and fitting same data set group at a time), therefore BIC would not be considered in the following analysis.

	AIC_{92}	$\Delta_{\it 92}AIC$	$w_{92}AIC$	BIC_{92}	$\Delta_{92}BIC$	R^2	Point
Baranyi	-50.92886	0	9.996976e-	-48.98923	0	0.9942026	1
			01				
Gompertz	-33.33531	17.59355	1.511741e-	-31.39568	17.59355	0.9748831	0
			04				
Logistic	-33.33458	17.59428	1.511190e-	-31.39495	17.59428	0.9748816	0
			04				
Polynomia	ul -18.89898	32.02988	1.108328e-	-16.95935	32.02988	0.9163562	0
			07				

Table 2: Evaluation matrix, ID=92

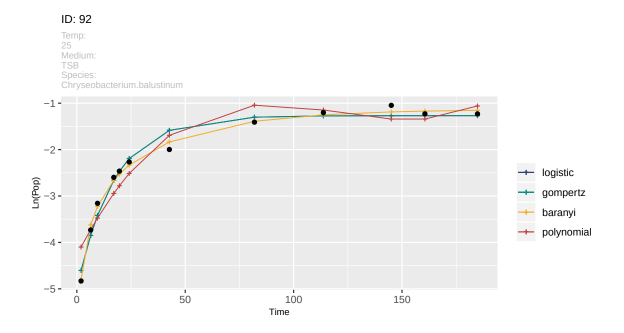


Figure 3: Example of a data set, where all models fit.

57 4.2 Fit Result

168 **4.2.1** Overall

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The final result for all models' fitting performances in the general data set can be seen in table 3. The

Gompertz model has the highest P amongst all. It is the the best model for fitting this general data

set, regarded as the best model to fit a data set group for a total of 182 times, taking up 62.5% of

the all groups there are. The Logistic and Baranyi performed quite similarly with a percentage of

54.6 and 52.6 respectively following up the Gompertz.

Baranyi	$\mathbf{Gompertz}$	Logistic	Polynomial	Total Groups
153(52.58%)	182 (62.5%)	159(54.6%)	77(24.6%)	291

Table 3: Overall Score, with percentage of Score over total size in bracket.

75 4.2.2 Co-variate Group Fit

Upon further examination of the data set, Tetraselmis tetrahele is used to conduct the analysis of 176 model's performance differences across temperature groups in the next section, as it is the only species in the data set that allows such analysis (5 repetitions under same medium in different temperature 178 groups). However, this part will only illustrate the overall fit results in different temperature groups, 179 a detailed analysis on **Tetraselmis tetrahele** will be done in the following section. The data set used 180 for this study contains 17 temperature groups from 0 to 37°C. The final fit result for each group using 181 each model can be seen in table 4. 10 out of 17 (58.82%) groups, the Gompertz model still kept 182 the lead amongst others. The Logistic and Baranyi performed quite similarly again, taking the lead 183 for 5 out of 17(29.41%) and 6 out of 17(35.29%) groups respectively. 184

Temperature(${}^{\circ}C$)	Baranyi	Gompertz	Logistic	Polynomial
0	0	5	1	0
2	5	8	7	5
4	8	6	5	6
5	13	11	12	7
6	4	3	5	4
7	4	14	6	2
8	5	9	5	3
10	10	18	11	4
12	10	7	7	5
15	24	29	24	14
16	6	4	7	0
20	19	21	20	15
25	19	15	17	5
30	7	9	9	4
32	3	3	3	0
35	14	16	17	3
37	2	4	3	0

 Table 4: Detailed point for temperature groups

Fit results in data set groups with and without a death phase are illustrated in table 5. The **Gompertz**model performed the best in data set groups without a death phase, and was out performed by **Baranyi**when there is a death phase.

Death Phase	Baranyi	Gompertz	Logistic	Polynomial
Yes	19(73.1%)	14(53.8%)	16(61.5%)	8(30.8%)
No	134(52.3%)	168(63.4%)	143(54.0%)	69(26.0%)

Table 5: Detailed point for death phase groups, with percentage of the total group size following in brackets.

4.3 Comparisons In Performances

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In order to have a more accurate and reasonable analysis on model's performance across different 189 temperature groups, **Tetraselmis tetrahele** is chosen for this part of the analysis. A total of 5 190 temperature groups are presented (T = 5, T = 8, T = 16, T = 25, T = 32). Differences in the models 191 performance which was indicated in wAIC can be seen in figure 4. Using the Kruskal-Wallis Test, All 192 models have shown a significant differences (Baranyi p-value = 0.001679, Logistic p-value = 0.01684, 193 Gompertz p - value = 0.03801, Polynomial p - value = 0.01672) in its wAIC across different tem-194 perature groups at .05 significance level, indicating temperature may have affected these models' 195 performance for this specific subset of data. The **Baranyi** have the most significant differences across 196 all groups. 197 Further comparing the pairwise difference shows that the **Baranyi** has the most significant differences 198 in pair $8^{\circ}C$ - $16^{\circ}C(p-value=0.007937)$, the **Gompertz** has the most significant differences in pair 199 $8^{\circ}C$ - $25^{\circ}C(p-value=0.01587)$, the **Logistic** has the most significant differences in pair $8^{\circ}C$ - $16^{\circ}C$ 200 and pair $5^{\circ}C$ - $16^{\circ}C(p-value=0.007937)$, while the **Polynomial** has significant differences in pair 201 $5^{\circ}C - 16^{\circ}C$, pair $8^{\circ}C - 16^{\circ}C$, and pair $16^{\circ}C - 25^{\circ}C(p - value = 0.007937)$. 202

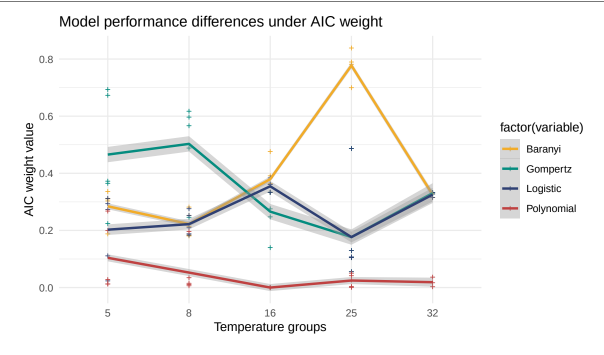


Figure 4: wAIC of different model across temperature groups, line with 2.5% CI.

Similarly for the death phase groups (seen in figure 5), **Baranyi** and **Gompertz** model have shown a significant differences (p - value = 0.04993, p - value = 0.02158) in date set groups with and without a death phase at .05 significance level, suggesting that having a death phase in the data set group would probably affect these models' performances. The **Polynomial** and **Logistic** model have not shown a significant differences (p - value = 0.6555, p - value = 0.8133) in their performance across the two groups.

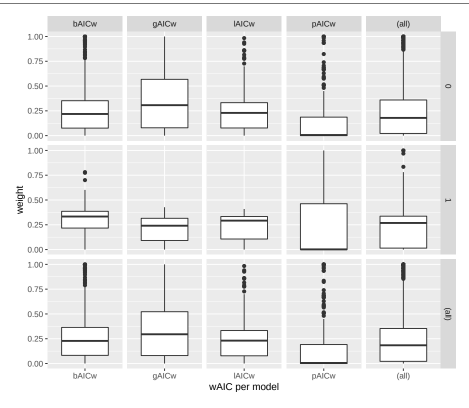


Figure 5: wAIC of different model for dataset with and without a death phase

²¹⁰ 5 Discussion

5.1 Reasons For Performance Difference Amongst Temperature Groups

- 212 Although the previous analysis has concluded there are performance differences for all four models,
- especially for the Baranyi, and some vague trends can be seen as in figure 4 amongst different tem-
- perature groups, the results are only based on a rather small data subset (size = 92) for **Tetraselmis**
- 215 **tetrahele**. It is hardly an indication for anything solid.
- Also, the difference could be caused by different sample sizes, measurement errors, and so on. How-
- ever, these errors are very hard to quantify.
- The fundamental question for model performance difference is probably much aligned with how tem-
- 219 perature will affect the shape of the population growth curve for bacteria since models(excluding the
- Polynomial) used in this study indicate a very similar shape regardless.
- A proper investigation in the relationship between temperature and models' performances or any other
- biological parameters like r_{max} should require much more experimental data, and the analysis should
- be conducted on a larger scale to limit the error brought by the data.

²⁴ 5.2 Death Phase And The Polynomial Model

The **Polynomial**(Cubic) model is introduced to mainly illustrate the disadvantages other models have regarding the fourth stage of bacterial growth curves - death phase - since it allows the curve to have a decreasing phase after hitting the maximum point.

In reality, as shown in table 5, the behaviour of the Polynomial model is not better than any of the other models(ranked the last), and there is no significant difference in perfromance between the two groups. A closer look at a data set group that has a death phase(figure 6) may help to understand why this is the case.

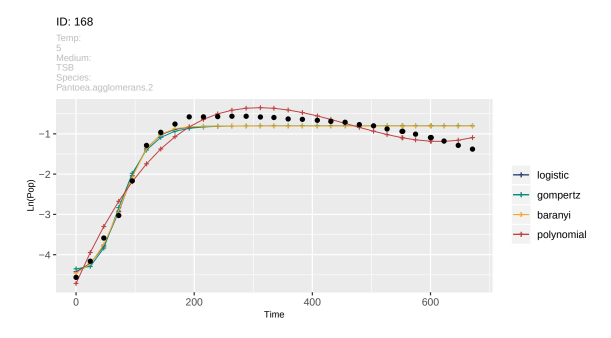


Figure 6: Example of a data set, where all models fit.

As can be seen, the data set group has shown a decline since Time = 200, and models other than
Polynomial are fitting the decline using a straight line. Although the Polynomial model is somehow
catching the decline tendency, the shape of the Polynomial(Cubic) model is relatively different from
the 4-phased growth curve - it needs two falls and one rise, or two rises and one fall. Because of this
feature of the Polynomial(Cubic) curve, even though it has catched a vague and overall trend of a
data set, it does not necessarily help with the overall fit result.

However, the comparison does show the disadvantages of the other three models used in this study at
catching the decline after the peak. It also shows that the Polynomial model, due to its nature, is not

240 the best at fitting population growth curves.

This could be a very rough first step of working on finding or building models that could work well

242 for the 4-phased growth curve.

5.3 Future Work

There are more biology related analysis that could be done after understanding that the Gompertz model is the best model to use for this data set. An initial thought would be trying to understand the effect temperature has over t_{lag} and r_{max} , since those are calculated each time the model fit a data set. A simple correlation and regression analysis could be done to understand the basic relationship, or using the new *time* and t_{lag} , *time* and r_{max} data set, more models in relation to those variables could be bench-marked. However, since the focus of this study is fitting and comparing models for bacterial growth curves, those topics are not discussed.

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