APPROXIMATE BAYESIAN COMPUTATION APPLIED TO MODEL SELECTION AND PARAMETER CALIBRATION IN CELL PROLIFERATION

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Abstract. Approximate Bayesian Computation is used in this work for the selection and calibration of cell proliferation models. Four competing models based on ordinary differential equations are analyzed, by using the measurements of the proliferation of DU-145 prostate cancer viable cells during seven days. The selection criterion of the ABC algorithm is based on the Euclidean distance between the model prediction and the experimental observations. The Richards Model and the Generalized Logistic Model were selected by the ABC algorithm used in this work, providing accurate estimates of the evolution of the number of viable cells. Bayes factor revealed that there was no evidence in favor of any of these two selected models.

1 INTRODUCTION

Cell proliferation is numerically given by the difference between the numbers of newly-divided and dying cells. In order to predict the number of viable cells, several mathematical
models have been proposed in the literature [1,2]. These models have been applied for tumors, since in cancer cells the proliferation process is increased due to the abnormal metabolic activity [3]. Costa et al. [4], for example, have used one of these proliferation models to represent the behavior of prostate cancer cells (DU-145) in vitro. In addition, they have analyzed a chemotherapy treatment using doxorubicin (DOX). Costa et al. [5] have applied Approximate Bayesian Computation via a Monte Carlo Sequential Method (ABC-SMC) [6-8] to select from competing models the one that best represented the proliferation of prostate cancer tumor cells during in vitro chemotherapy experiments. Distinct hypotheses are included in a specific model. Thus, the selection and calibration of these models are of great interest.

The goal of this work is to select among four continuous models the one that better represents in vitro experimental data of the proliferation of DU-145 human prostate cancer cells. In order to perform this analysis, the Approximate Bayesian Computation (ABC) algorithm of Toni et al. [6] is applied for model selection and calibration, since this algorithm is robust and indicated for cases that the likelihood is not exactly known [4], such as in this work.

2 MATHEMATICAL MODELS

Different approaches can be used to model cell proliferation, by applying continuous, discrete or hybrid models. The choice of the model type for the investigation depends on the type of experiment, goal of the study and mainly the biological characteristics of the cells under analysis. In this work, the experimental data was obtained from ATCC [9], as shown in Figure 1.

![Figure 1](https://www.scipedia.com)

Figure 1: In vitro experiments results provided by American Type Culture Collection (ATCC) [9].

Due to the characteristics of the experimental data, without cycling or repeating behavior, only continuous models are used here in the inverse analysis of model selection/calibration. As can be seen in Figure 1, the number of viable cells did not grow without bounds. For this reason, the Exponential and Mendelsohn models were not considered in the analysis. In order to consider a bound in the proliferation process, four models are investigated: Logistic, Gompertz, Richards and Generalized Logistic [1-3]. The predictions provided by these models assume
uniform cell distribution and proliferation in the cell culture. In addition, the experiments are considered isothermal, at a constant temperature of 37 °C.

In these models, \( N_i \) is the number of viable cells varying with time \( t \), with the initial number of cells given by \( N_{0,i} \). The rate of proliferation is given by parameter \( \alpha_i \) and the growth saturation by the dimensionless parameter \( \gamma_i \). The support capacity that takes into account the space condition, oxygen availability and nutrient source is considered by \( K_{\text{sup},i} \). The mathematical models are presented in Equations 1-8 where the subscripts \( i = 1, 2, 3, 4 \) designate the models.

### 2.1 Model 1: Logistic Model

\[
\frac{dN_1(t)}{dt} = \alpha_1 N_1(t) \left[ 1 - \frac{N_1(t)}{K_{\text{sup}1}} \right]; \quad t > 0
\]  
(1)

\[N_1(0) = N_{0,1}; \quad t = 0\]  
(2)

### 2.2 Model 2: Gompertz Model

\[
\frac{dN_2(t)}{dt} = \alpha_2 N_2(t) \ln \left( \frac{K_{\text{sup}2}}{N_2(t)} \right); \quad t > 0
\]  
(3)

\[N_2(0) = N_{0,2}; \quad t = 0\]  
(4)

### 2.3 Model 3: Richards Model

\[
\frac{dN_3(t)}{dt} = \alpha_3 N_3(t) \left[ 1 - \left( \frac{N_3(t)}{K_{\text{sup}3}} \right)^{\gamma_3} \right]; \quad t > 0
\]  
(5)

\[N_3(0) = N_{0,3}; \quad t = 0\]  
(6)

### 2.4 Model 4: Generalized Logistic Model

\[
\frac{dN_4(t)}{dt} = \alpha_4 N_4(t) \left[ 1 - \left( \frac{N_4(t)}{K_{\text{sup}4}} \right)^{\gamma_4} \right]; \quad t > 0
\]  
(7)

\[N_4(0) = N_{0,4}; \quad t = 0\]  
(8)
3 SENSITIVITY ANALYSIS

Before the solution of the inverse problem, it is important to analyze the sensitivity coefficients of the measured variables with respect to each parameter. An analysis of the reduced sensitivity coefficients ($X_r$) is performed here, which are obtained by multiplying the parameter by the first partial derivative of the response with respect to that parameter [10-21]. The reduced sensitivity coefficients of viable cells with respect to the parameters are presented in Figure 2 for all the four models.

![Figure 2: Reduced sensitivity coefficients: (a) Logistic Model, (b) Gompertz Model, (c) Richards Model, (d) Generalized Logistic Model](image)

The reduced sensitivity coefficients were calculated with the parameter values given in Table 1. The sensitivity coefficients of the parameter $N_0$ for all models suddenly increase and then decay until approximately a null value, as time increases. The sensitivity coefficients with respect to the parameter $K_{sup}$ increases until the steady state is reached. The sensitivity coefficients are not linearly dependent for the parameters of the Logistic model and of the Gompertz model. On the other hand, parameters $\alpha$ and $\gamma$ of Richards model and Generalized Logistic model are correlated, as shown by Figures 2c and 2d.
Table 1: Nominal values used for the sensitivity analysis

| Parameter | Value |
|-----------|-------|
| $N_0 \ [\text{cell}]$ | 10,000 |
| $\alpha \ [\text{day}^{-1}]$ | 0.9 |
| $K_{\text{sup}} \ [\text{cell}]$ | 220,000 |
| $\gamma$ | 1.7 |

4 APPROXIMATE BAYESIAN COMPUTATION ALGORITHM

The Approximate Bayesian Computation (ABC) algorithm of Toni et al [6] was used in this work for the simultaneous model selection and estimation of the model parameters. This algorithm is presented in Table 2.

Table 2: ABC Algorithm [6]

1. Define the tolerances $\varepsilon_1, \varepsilon_2, \ldots, \varepsilon_P$ for each of the iterations (populations) used for selecting the model and its parameters. Also, specify the distance function $d(Y, Y^*)$ that substitutes the likelihood function. Set the population indicator $p = 0$.
2. Set the particle indicator $i = 1$, where each particle represents, at each iteration, a model and its parameters.
3. Sample the model $M^*$ from the prior distribution for the models $\pi(M)$. If $p = 0$, sample the candidate parameters $P^{**}$ from the prior distribution for the parameters of model $M^*$, that is, $\pi(P(M^*))$. Else, sample $P^*$ from the parameters in the previous population $P(M^*)_{p-1}$, with weights $w(M^*)_{p-1}$, and perturb this particle to obtain $P^{**} = K_p(P^*, P^{**})$, where $K_p$ is a perturbation kernel.
4. If $\pi(P^{**}) = 0$, return to step 3. Else, simulate from the forward problem (operator $f$) a candidate set of observable variables with model $M^*$ and parameters $P^{**}$, that is, $Y^* = f(Y|P^{**}, M^*)$.
5. If $d(Y, Y^*) > \varepsilon_p$, return to step 3. Otherwise, set $M^*_p = M^*$, add $P^{**}$ to the population of particles $P(M^*)_p$ and calculate the particle weight

$$w(M^*)_p = \begin{cases} 1 & \text{if } p = 0 \\ \frac{\pi(P(M^*)_p)}{\sum_{j=1}^{N} w(M^*)_j P(M^*)_p, P(M^*)_p} & \text{if } p > 0 \end{cases}$$

6. If $i < N$, where $N$ is the number of particles, set $i = i + 1$ and go to step 3.
7. Normalize the weights.
8. If $p < P$, where $P$ is the number of iterations (populations), set $p = p + 1$ and go to step 2. Otherwise, terminate the iterations.

Instead of using the likelihood function, the ABC algorithm is based on a distance function calculated at each set of successive populations (formed by particles composed of the model selected and parameters estimated). A tolerance ($\varepsilon$) is prescribed at each population for the distance function given in this work by the Euclidean distance between the system dependent variable $Y^*$ and the experimental data $Y$. If the Euclidean distance is smaller than the tolerance,
the particle is accepted; otherwise, the particle is rejected and a new particle is generated.

5 RESULTS AND DISCUSSIONS

The ABC-SMC method with 4000 particles was applied to model selection and estimation of the model parameters. Uniform priors and uniform transition Kernels were adopted for the parameters, as presented in Table 3. Note that the upper and lower limits of the transition Kernel were assumed to be ±1% of the upper bound of the prior for each parameter.

Table 3: ABC Priors distribution and transition Kernels for the parameters

| Models | Priori  | Transition Kernel |
|--------|---------|-------------------|
| 1,2,3,4 | $N_0 \sim U(1000;19000)$ | $U(-190;190)$ |
| 1,2,3,4 | $\alpha \sim U(0.09;4.5)$ | $U(-0.045;0.045)$ |
| 1,2,3,4 | $K_{\text{sup}} \sim U(22000;418000)$ | $U(-4180;4180)$ |
| 3,4    | $\gamma \sim U(0.170;8.5)$ | $U(-0.085;0.085)$ |

The experimental data presented in Figure 1, obtained from ATCC™ (American Type Culture Collection) [9] for the number of viable cells of DU-145 prostate cancer cells during in-vitro experiments, were used in the inverse problem. In order to solve the inverse problem, the four mathematical models presented in Equations 1 to 8 were solved by the 4th order Runge-Kutta algorithm. The choice of the tolerances for the sequential populations of the ABC algorithm were set by starting at $5.42 \times 10^5$ and finishing at $5.42 \times 10^3$, along a total of fifty-seven populations. The last tolerance was imposed in accordance with Morozov’s discrepancy principle, assuming a standard deviation ($\sigma$) of 1% of the maximum value of viable cells, that is, $\varepsilon_{\text{last}} = \sigma \sqrt{N_{\text{measurements}}}$.

The ABC-SMC algorithm of Toni et al. [6] selected the Richards Model and the Generalized Logistic Model, as shown by Figure 3, to represent the experimental data presented in figure 1. In this figure it is possible to observe that after 27 populations only these two models have been selected. However, a total of 57 populations were needed to perform the correct calibration of the parameters of these two models.
The problem of selecting models can be associated to hypothesis tests, such as the Bayes factor proposed by Kass and Raftery [22]. The Bayes Factor for Models 3 and 4 is given by the posterior probability of each model in relation to the data, that is,

\[ B_{43} = \frac{\pi(M_4 | Y)}{\pi(M_3 | Y)} \]  

(9)

The criteria of Kass and Raftery [22] for interpreting the Bayes factor is presented in Table 4. At the final population shown in Figure 3, 2608 particles were selected for Model 4 and 1392 were selected for Model 3, which gives a Bayes factor of 1.87. In accordance with Table 4, there is no evidence in favor of any of the models 3 or 4.

| Evidence against Model 3       | 1 to 3 | 0 to 2 | Not worth more than a bare mention |
|-------------------------------|-------|-------|-----------------------------------|
| 3 to 20                       |       | 2 to 6| Positive                          |
| 20 to 150                     | 6 to 10|      | Strong                            |
| >150                          | > 10  |      | Very strong                       |

Table 4. Bayes factor [22]

The histograms of the model parameters at the final population are presented by Figures 4 and 5. These histograms exhibit approximate Gaussian behaviors, centered at mean values. The means, standard deviations and 95% credible intervals for the estimated parameters for both models 3 and 4 are presented in Table 5 and 6, respectively.
The numbers of viable cells computed with models 3 and 4, considering the mean of the accepted particles at the last population, are presented in figures 6 and 7, respectively. The light blue lines in these figures are the estimated curves calculated with each of the accepted particles at the final population. These figures show that both model estimations have an excellent agreement with the experimental data, thus confirming that either one of the competing models 3 or 4 (Richards or Generalized Logistic) could be used to represent in vitro experiments performed with DU-145 human prostate cancer cells.

We note that the results presented here were not influenced by the stochastic simulations performed. In fact, the results were qualitatively unchanged in four runs of the ABC-SMC algorithm used in this work.
Figure 5: Histograms for the parameters of Model 4: (a) $N_{04}$, (b) $\alpha_4$, (c) $K_{sup4}$, (d) $\gamma_4$.

Table 5: Parameters estimation - Model 3

| Parameter  | Mean   | Standard deviation | Lower limit 95% | Upper limit 95% |
|------------|--------|--------------------|-----------------|-----------------|
| $N_{03}$ [cell] | 5826.6 | 74.9072            | 5712.7          | 5998.6          |
| $\alpha_3$ [day$^{-1}$] | 0.7777 | 0.0049             | 0.7667          | 0.7858          |
| $K_{sup3}$ [cell] | 191,660 | 671.9028           | 190,460         | 193,040         |
| $\gamma_3$   | 2.8621 | 0.0662             | 2.7315          | 2.9938          |

Table 6: Parameters estimation - Model 4

| Parameter  | Mean   | Standard deviation | Lower limit 95% | Upper limit 95% |
|------------|--------|--------------------|-----------------|-----------------|
| $N_{04}$ [cell] | 8476.1 | 80.0335            | 8309.2          | 8615.0          |
| $\alpha_4$ [day$^{-1}$] | 2.2823 | 0.0244             | 2.2316          | 2.3273          |
| $K_{sup4}$ [cell] | 192,440 | 656.2398           | 191,200         | 193,700         |
| $\gamma_4$   | 3.4078 | 0.0457             | 3.3164          | 3.4912          |
6 CONCLUSIONS

The ABC-SMC algorithm of Toni et al. [6] was applied with 4000 particles for model selection and estimation of cell proliferation model parameters. The parameters were considered with uniform priors and uniform transition Kernels were used in the algorithm. In order to solve the inverse problem, the four mathematical models were solved with the Runge-Kutta’s 4\textsuperscript{th} order method. The tolerances for the sequential populations of ABC-SMC method
decreased from $5.42\times10^5$ to $5.42\times10^3$ along fifty-seven populations. The last tolerance was imposed in accordance with the assumed measurement uncertainty following Morozov’s discrepancy principle. The Richards Model and the Generalized Logistic Model were both selected by ABC-SMC algorithm, providing accurate estimations of the number of viable cells. An analysis of Bayes factor revealed that both models can be used to accurately represent in vitro measurements of the time evolution of the DU-145 human prostate cancer cells.

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