
Bayesian calibration, validation and uncertainty quantification for predictive modelling of tumour growth: a tutorial

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Received: date / Accepted: date

Abstract In this work we present a pedagogical tumour growth example, in which we apply calibration and validation techniques to an uncertain, Gompertzian model of tumour spheroid growth. The key contribution of this article is the discussion and application of these methods (that are not commonly employed in the field of cancer modelling) in the context of a simple model, whose deterministic analogue is widely known within the community. In the course of the example we calibrate the model against experimental data that is subject to measurement errors, and then validate the resulting uncertain model predictions. We then analyse the sensitivity of the model predictions to the underlying measurement model. Finally, we propose an elementary learning approach for tuning a threshold parameter in the validation procedure in order to maximize predictive accuracy of our validated model.

Keywords Bayesian Calibration, Tumour Growth, Model Validation

1 Introduction

The treatment of cancer represents a significant challenge in modern healthcare, and has given cause for the development of numerous mathematical

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and computational models of tumour growth and invasion over many decades. As our level of scientific understanding increases and we have access to ever greater computational power, we are able to create increasingly realistic mathematical models of biological phenomena and to compute numerical approximations of model solutions with greater accuracy. It is therefore natural to consider the transfer of mathematical and computational models from a purely theoretical, informative, or qualitative setting to the clinic, as a possible means of guiding patient therapy via prediction (Savage 2012; Gammon 2012; Yankeelev et al. 2013). However, if we wish to make clinically relevant, patient-specific predictions, it is of vital importance that these predictions are made in a safe and reliable manner.

Output from computational models differs from physical observations for a multitude of reasons. For instance, as parameter values are often inferred from experimental data, there is uncertainty associated with their values, and often solutions to systems of equations are subject to numerical errors associated with their discretization. Perhaps most fundamentally, however, mathematical models are abstractions of reality, necessarily simplifying or omitting phenomena and, as such, even exact solutions obtained from precise data may yield non-physical results. In order for computational model outputs to be viewed as sufficiently reliable for safety-critical applications, such as predictive treatment planning, any parametric or structural uncertainties must be quantified, as well as any inaccuracies resulting from numerical approximation.

There has been much recent work from the engineering and physical sciences communities surrounding the development of techniques for assessing the credibility of quantitative computational model predictions in safety critical applications. This field is often referred to as verification, validation and uncertainty quantification (VVUQ), and provides a formalism, techniques and best practices for assessing the reliability of complex model predictions (Oberkampf et al. 2004; Oden et al. 2010a,b; NRC 2012; Oberkampf and Roy 2010; Roache 2009). The growing importance of VVUQ in engineering and the physical sciences is highlighted by the extensive guidelines and standards for verification and validation in solid mechanics, fluid dynamics, and heat transfer produced by the American Society of Mechanical Engineers (ASME 2006, 2009, 2012). Furthermore, the US National Research Council (NRC) recently published an extensive report on VVUQ (NRC 2012) which, in addition to providing an extensive review of the literature with informative examples, highlights the importance of training young scientists in VVUQ as a field of importance in the 21st century. The primary purpose of this article is to serve as a pedagogical tool for members of the (continuum mechanics) cancer modelling community, introducing a range of concepts and techniques from the field of VVUQ for a simple and familiar biological example, in a manner similar to that adopted in Aguilar et al. (2015); Allmaras et al. (2013).

Following the terminology set out in NRC (2012), we refer here to *verification* as “the process of determining how accurately a computer program correctly solves the equations set out in the mathematical model”, *validation*

as “the process of determining the degree to which a model is an accurate representation of the real world from the perspective of the real world uses of the model”, and *uncertainty quantification* as “the process of quantifying uncertainties associated with model calculation of true physical quantities of interest”. Furthermore, we refer here to *calibration* as the process of inferring the values of model parameters from indirect measurements. In the context of predictions of tumour growth and invasion, these techniques provide us with a means of quantifying the robustness of our calibrated model predictions against empirical data, subject to measurement errors, and deficiencies in our biological understanding (manifested in an inherently incorrect description due to our mathematical model).

The application of validation and uncertainty quantification techniques in the mathematical modelling of solid tumour growth is currently limited. In Hawkins-Daarud et al. (2013) the authors set out a Bayesian framework for calibration and validation based on that described in Babuška et al. (2008) in a computational engineering context. However, the authors consider synthetic data as opposed to *in vitro* or *in vivo* experimental data. In Achilleos et al. (2013, 2014), a stochastic mixture model updated in a Bayesian manner is introduced and its tumour-specific predictions are validated against experimental data from a mouse model. In other areas of mathematical biology, VVUQ techniques are gaining recognition, e.g. in cardiac modelling (Pathmanathan and Gray 2013).

In order to make suitably accurate and reliable predictions, we are required to estimate parameter values, such as reaction rates and diffusion coefficients. Often, it is impossible to measure these parameters directly; they must be inferred. The classical, *deterministic* approach is to find the single set of parameter values that among all possible parameter choices best matches the observed data, in some appropriate sense. There are many available methods for determining this set, however, any approach that yields a single choice does not fully account for any uncertainty in the empirical data, nor any possible uncertainty regarding the mathematical model (Allmaras et al. 2013), and as such, does not account for uncertainty in the estimated parameters. Here, we consider a statistical (Bayesian) approach to parameter estimation to determine a probability density function (pdf) for the parameters, that updates any prior information we have regarding the parameters, by incorporating new information obtained from the observed data. In this setting, our model predictions are no longer the solution of a deterministic mathematical model, but rather a description of the random variable, or random field, that is a solution of the underlying *stochastic* model. We refer to Allmaras et al. (2013); Aguilar et al. (2015); Kaipio and Somersalo (2006); Tarantola (2005) for a more thorough discussion regarding Bayesian model calibration. We remark that it is also possible to infer information regarding parameter uncertainty in the classical inference setting, though we consider here the Bayesian approach only.

Once the model is suitably calibrated, we validate its performance by considering various behaviours and responses. Firstly, the model must

reproduce observed behaviour of the physical system for appropriate parameter values; we refer to Gelman et al. (2014a) for a thorough description of a range of checks for model fit in a Bayesian framework. Moreover, the output of the model must be robust to perturbations that are likely in the context of the intended use of the model. Such validation may simply involve direct comparison between model results and physical measurements. However, for complex models a sophisticated statistical approach may be required, combining hierarchical models¹ and multiple sources of physical data with subjective expert judgment.

In this article, we embark upon a process of model calibration and validation against *in vitro* experimental data, quantifying uncertainties in our model predictions and assessing the robustness of our modelling assumptions for a Gompertzian model of tumour spheroid growth (Gompertz 1825; Laird 1964). We adopt a similar procedure for prediction validation as set out in Hawkins-Daarud et al. (2013). However, we consider additional posterior predictive checks for our model, as described in Gelman et al. (2014a), and further assess the sensitivity of our predictions to assumptions in our statistical model. The primary contribution of this work is to demonstrate the application of existing VVUQ techniques to a mathematically simple model of tumour growth by means of an educational example. The simplicity of the Gompertzian model permits us to neglect any issues surrounding spatial and temporal discretizations, and focus solely on practical issues regarding the statistical approach adopted, in a similar manner to that in Allmaras et al. (2013); Aguilar et al. (2015). Whilst the computations presented here are not directly applicable to the clinic, over the longer term the statistical techniques discussed here could form the basis of a robust means of assessing quantitative model predictions for clinical applications based on *in vivo* data, potentially involving significantly more complex models.

The remainder of this article is organised as follows, in Section 2 we introduce the underlying model of tumour growth, describe the procedure used for collecting the experimental data, and specify the quantity of interest we wish to predict. In Section 3 we introduce the Bayesian framework and describe the computational techniques employed to determine the joint pdf for the parameters in the model. We assess how well our predictive model fits to the data employed in the calibration in Section 4, and in Section 5 we assess the extrapolative predictive capability of our calibrated model against a validation data set. In Section 6 we assess the robustness of our predictions against assumptions in our statistical model, and in Section 7 we consider the application of the techniques set out in the previous sections to multiple experiments. To conclude this article, we depart from the pedagogical example of the previous sections to discuss extensions to clinically relevant predictions in Section 8 and, finally, in Section 9 we draw conclusions about the work presented in this article, and highlight ongoing and future work.

¹ i.e. we model observable outcomes conditionally on parameters which themselves are given a probabilistic description in terms of further parameters known as hyperparameters.

2 Problem Description

In this section we formally describe the biological modelling problem under consideration, with minimal reference to the statistical framework employed in subsequent sections. To this end, we first set out the mathematical model for tumour spheroid growth considered in the remainder of this article. We then specify details of the experimental data available for the proceeding analysis and describe the calculation of an approximate error in the measurements. Finally, we define the quantity of interest (QoI) we wish to predict using our calibrated mathematical model.

2.1 Tumour Growth Model

In this work we consider a Gompertzian model of tumour spheroid growth (Gompertz 1825; Laird 1964), in which the tumour volume V at time t is given by

$$V(t) = K \exp \left(\log \left(\frac{V_0}{K} \right) \exp(-\alpha t) \right), \quad (1)$$

where V_0 denotes the initial tumour volume at $t = 0$, K denotes the carrying capacity (the maximal tumour size for nutrient-limited growth), and α denotes a growth rate related to the proliferative ability of the cells. In the deterministic setting, each of the parameters, V_0 , K , and α , takes a single constant value for a given data set, whereas here in the uncertain setting, we view each one as a random variable $V_0, K, \alpha : \Omega \rightarrow \mathbb{R}^+$, where Ω denotes a suitable sample space. Under this assumption, we note that the tumour volume $V(t)$ is also a random variable.

2.2 Experimental Data

Two-dimensional images of a tumour spheroid were captured at 14 time points over a period of 28 days. The times at which the measurements were taken are given in Table 6 in Appendix A. The resultant images were analysed using *SpheroidSizer* (Chen et al. 2014). In particular, the length of the major and minor axes of the spheroid, denoted ℓ_1 and ℓ_2 , respectively, were identified. Figure 1 shows a representative image of a tumour spheroid, and a processed image in which the boundary of the tumour is shown.

2.2.1 Partitioning into Subsets

In Hawkins-Daarud et al. (2013), the notion of calibration and validation data sets are discussed. The calibration set is employed in the calibration of the model and the validation set is utilized to validate the calibrated model. We adopt this same procedure here, and define two sets S_C and S_V corresponding to calibration and validation data respectively. For demonstrative purposes,

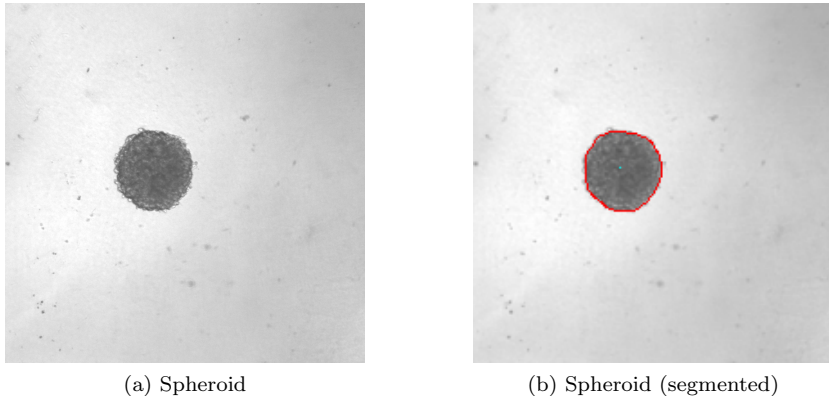


Fig. 1: Exemplar images of a tumour spheroid. (a) The raw image. (b) The boundary of the spheroid obtained using image segmentation software superimposed on the raw image.

however, we additionally reserve the data associated with the final time point for a final predictive check. Ordinarily, all data would be employed in calibration and validation. We depart from this approach here in order to demonstrate whether our model predictions coincide with the value measured in experiment, to make the tutorial more informative. The precise selection of the calibration and validation data employed in this study, along with the rationale behind their selection, is discussed in greater detail in Section 3.2.3.

2.2.2 Volume Calculation

In order to estimate the *true* length of the major and minor axes, we must calibrate the image scale, i.e. we must establish the physical dimensions of a single pixel in an image obtained from the microscope. To perform this calibration, we place a rule of length $100\ \mu\text{m}$ under the microscope and count how many pixels extend along its length. In our experimental configuration, $100\ \mu\text{m}$ corresponds to 40 pixels; thus the scale of each image is calculated as $s = 2.5\ \mu\text{m}$ per pixel. We estimate the volume of the spheroid by assuming that the length of the third axis, ℓ_3 , is given by the geometric mean of ℓ_1 and ℓ_2 , i.e. $\ell_3 = \sqrt{\ell_1\ell_2}$. The volume of the spheroid is then estimated by

$$V = \frac{\pi}{6} \ell_1 \ell_2 \ell_3 = \frac{\pi}{6} (\ell_1 \ell_2)^{3/2}. \quad (2)$$

2.2.3 Measurement Error Model

The volume measurements introduced in the previous section are subject to experimental noise. We assume that this noise is independently and normally distributed with a mean of zero and a standard deviation of σ_V . To estimate σ_V we approximate the error introduced at each of the image processing steps:

- 1. Image scale calibration:** The first point at which we may introduce an error into our calculation is in the estimation of the scale, s . Assuming we count n pixels (accurate to the nearest pixel), then the potential error in n is given by

$$\sigma_n = 0.5. \quad (3)$$

This results in a potential error in the calculation of s , denoted σ_s , given by

$$\sigma_s = \left| \frac{\partial s}{\partial n} \right| \sigma_n = \frac{100}{n^2} \sigma_n = 0.03125 \mu\text{m per pixel}. \quad (4)$$

- 2. Measurement of major and minor axes:** We assume that the automated segmentation procedure adopted by *SpheroidSizer* identifies the tumour boundary accurate to the nearest pixel. The length of the major and minor axes in units of pixels, d_1 and d_2 , respectively, are then subject to a potential error of $\sigma_{d_{1,2}} = \sqrt{2}$. The error in $\ell_{1,2} (= s \cdot d_{1,2})$ is then given by:

$$\sigma_{\ell_{1,2}} = \sqrt{d_{1,2}^2 \sigma_s^2 + s^2 \sigma_{d_{1,2}}^2}. \quad (5)$$

- 3. Inference of the length of the third axis:** We assume that the true value of ℓ_3 , ℓ_3^* , is subject to an error, ξ , i.e. $\ell_3^* = \sqrt{\ell_1 \ell_2} + \xi$. We assume that this error has zero mean and standard deviation $\sigma_{\ell_3} \approx \frac{(\ell_1 - \ell_2)}{2}$.

The above errors are combined using conventional error propagation to obtain the following estimate for σ_V :

$$\sigma_V = \sqrt{\left(\frac{\partial V}{\partial \ell_1} \right)^2 \sigma_{\ell_1}^2 + \left(\frac{\partial V}{\partial \ell_2} \right)^2 \sigma_{\ell_2}^2 + \left(\frac{\partial V}{\partial \ell_3} \right)^2 \sigma_{\ell_3}^2}. \quad (6)$$

Figure 2 shows the spheroid volume at $\{t_1, \dots, t_{14}\}$ as described above, with error bars corresponding to $\pm 2\sigma_V$.

2.3 Predictive Quantity of Interest

The process of validation and uncertainty quantification is applicable only for a specified QoI; acceptable predictive model performance for one particular QoI does not necessarily imply acceptable performance for all possible QoI. In particular, here we select a QoI that is of practical relevance to the real world applications of the model. In this study we consider the tumour volume at t_{14} as our predictive QoI because, in clinical applications of these techniques, it is likely that a QoI such as tumour volume at a given time may be employed as a proxy for patient prognosis. Recalling the discussion in Section 2.2.1, for illustrative purposes in the context of the tutorial nature of this article, we withhold the data obtained at t_{14} from all calibration processes so that we may compare our model prediction to experimental data.

We remark that typically, extrapolative predictions are more challenging than interpolative predictions. Intuitively, we see that the extrapolative case

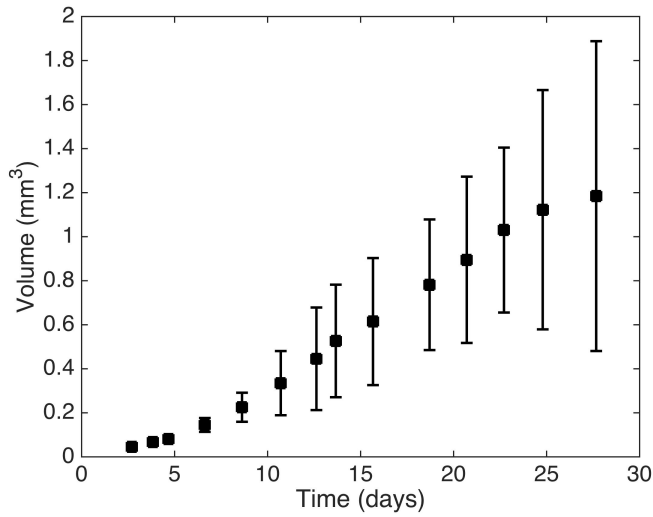


Fig. 2: A plot showing the experimental tumour volume data at times t_1, \dots, t_{14} , with error bars corresponding to $\pm 2\sigma_V$ where σ_V is defined by (6).

may introduce additional phenomena that are not well modelled, causing the prediction to have greater errors than indicated by a poorly designed validation study. As such, when we come to define our calibration and validation data sets, we adopt the best practice of validating our calibrated model against data that is as close to our predictive scenario as possible. We discuss the selection of calibration and validation data sets in greater detail in Section 3.2.3.

2.4 Overview of the Calibration and Validation Process

Now that the biological problem has been fully specified (i.e. the growth model, experimental data and QoI), we can outline the process employed in subsequent sections to predict the QoI and determine whether this prediction is not invalid. Algorithm 2.1 highlights the steps taken to perform model calibration and validation for our tumour growth example.

3 Model calibration

In this section we set out the Bayesian framework for model calibration. We then describe the numerical algorithms used to calibrate our model, briefly discussing the criteria we use to assess convergence of the algorithms. Finally, we apply these numerical algorithms to calibrate the Gompertzian model given in (1) against a subset of the experimental data described in Section 2.2.

Algorithm 2.1 Calibration and Validation Process

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- 1: Specify the calibration and validation data denoted S_C and S_V , respectively.
 - 2: Calibrate the model using the data S_C following the procedure set out in Section 3.
 - 3: Assess the ability of the model to reproduce the observed data S_C following the procedure set out in Section 4.
 - 4: Compute the PDF for the QoI using the model calibrated using S_C .
 - 5: Calibrate the model using the data S_V .
 - 6: Assess the ability of the model calibrated using S_V to reproduce the observed data S_V .
 - 7: Compute the PDF for the QoI using the model calibrated using S_V .
 - 8: Validate the prediction of the QoI made in 4. following the procedure set out in Section 5.
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3.1 Bayesian Calibration

We first recall that calibration refers to the process of inferring the values of model parameters from indirect measurements. The basis of the Bayesian approach is to enhance a subjective belief surrounding the probability of an event via the incorporation of experimental data. This process is inherently subjective and differs fundamentally from the *frequentist* approach, by which probabilities are assigned based on the frequency of their observations for large numbers of repeated experiments under identical conditions. The subjective nature of probability in the Bayesian framework provides a natural environment for assessing the chance of an event occurring when the concept of multiple repeated experiments under identical conditions is flawed. For instance, in a purely frequentist approach it is difficult to define an adequate notion of probability for patient mortality in a patient-specific model, as there can be no notion of assessing multiple patients with truly identical conditions. There are many examples in the literature of frequentist validation, e.g. Oberkampf and Barone (2006); in this work, however, we adopt a Bayesian framework and discuss the frequentist standpoint no further.

Before setting out the Bayesian method, we first introduce the relevant notation and terminology, where possible following the approach in Gelman et al. (2014a). We denote by $\boldsymbol{\theta}$ a vector of unobserved quantities, and we denote by $\mathbf{y} = (y_1, y_2, \dots, y_n)$ the observed data. Further, we denote conditional and marginal pdfs by $p(\cdot|\cdot)$ and $p(\cdot)$, respectively. In the Gompertzian model of Section 2.1, $\boldsymbol{\theta}$ corresponds to the model parameters in (1), i.e. $\boldsymbol{\theta} = (V_0, K, \alpha)$ and \mathbf{y} corresponds to the volume of the tumour spheroid obtained at the time points in the calibration data set S_C .

The model predictions of observable outputs are related to the input parameters by

$$\mathbf{y} = \mathcal{V}(t; \boldsymbol{\theta}, e), \quad (7)$$

where \mathcal{V} and e denote the measured volume of the tumour spheroid and measurement error, respectively. Given the parameter $\boldsymbol{\theta}$ and measurement error e , $\mathcal{V}(t; \boldsymbol{\theta}, e)$ invokes the solution of the forward problem and combination with the measurement error to yield \mathbf{y} , the observable variables. The relationship between the observable outputs, in our case the volume of the

spheroid, and model inputs at time t is then denoted by

$$\mathbf{y} = V(t; \boldsymbol{\theta}) + e, \quad (8)$$

where e denotes the error and the volume $V(\cdot; \cdot)$ is equivalent to that defined in (1), though now viewed as a function of t and $\boldsymbol{\theta} = (V_0, K, \alpha)$, via a simplifying abuse of notation. We note that there are also means of quantifying systematic discrepancies in the mathematical model, in which the data are modelled as

$$\mathbf{y} = V(t; \boldsymbol{\theta}) + \delta(t) + e,$$

where $\delta(t)$ denotes a discrepancy function. This approach may be suitable if we neglect significant biological effects in our underlying mathematical model or if we incur substantial discretization errors in the numerical approximation of PDE solutions. We proceed here employing the former model, and, as such, do not consider systematic model discrepancies explicitly. We refer to Kennedy and O'Hagan (2001); Higdon et al. (2005); Bayarri et al. (2007) for further details of these methodologies.

In order to make probabilistic statements regarding $\boldsymbol{\theta}$ and \mathbf{y} , we must introduce their joint probability density function, $p_{\text{JOINT}}(\boldsymbol{\theta}, \mathbf{y})$. This joint density can be written as the product of the prior distribution on $\boldsymbol{\theta}$, denoted $p_{\text{PRIOR}}(\boldsymbol{\theta})$, which corresponds to our *a priori* knowledge surrounding our model parameters, and the sampling distribution $p_{\text{SAMPLE}}(\mathbf{y}|\boldsymbol{\theta})$, thus yielding

$$p_{\text{JOINT}}(\boldsymbol{\theta}, \mathbf{y}) = p_{\text{PRIOR}}(\boldsymbol{\theta})p_{\text{SAMPLE}}(\mathbf{y}|\boldsymbol{\theta}). \quad (9)$$

We may view $p_{\text{PRIOR}}(\boldsymbol{\theta})$ as a summary of our subjective beliefs surrounding the distribution of the parameters at the outset of the calibration, which we further enhance via conditioning on observed experimental data. As such, we condition on \mathbf{y} and employ Bayes' theorem to obtain the conditional probability assigned to the parameter, referred to as the posterior density, that is given by

$$p_{\text{POST}}(\boldsymbol{\theta}|\mathbf{y}) = \frac{p_{\text{PRIOR}}(\boldsymbol{\theta})p_{\text{SAMPLE}}(\mathbf{y}|\boldsymbol{\theta})}{p_{\text{PRED}}^{\text{PRIOR}}(\mathbf{y})}, \quad (10)$$

where $p_{\text{PRED}}^{\text{PRIOR}}(\mathbf{y})$ denotes the marginal distribution

$$p_{\text{PRED}}^{\text{PRIOR}}(\mathbf{y}) = \int p_{\text{PRIOR}}(\boldsymbol{\theta})p_{\text{SAMPLE}}(\mathbf{y}|\boldsymbol{\theta}) d\boldsymbol{\theta}, \quad (11)$$

referred to as the prior predictive distribution. We consider the density $p_{\text{SAMPLE}}(\mathbf{y}|\boldsymbol{\theta})$ as a function of $\boldsymbol{\theta}$ rather than of \mathbf{y} , and refer to it as the likelihood function. The likelihood may then be interpreted as how 'likely' a parameter value is, given a particular outcome. The prior predictive distribution corresponds to the marginal distribution for the observable data obtained by averaging the likelihood over all possible parameter values with respect to the prior density. As such, the posterior distribution then corresponds to the enhanced degree of belief obtained via incorporation of the observed experimental data.

In order to make predictions regarding an unknown observable $\tilde{\mathbf{y}}$ from the same source as \mathbf{y} , we define the posterior predictive distribution, denoted $p_{\text{POST}}^{\text{PRED}}(\tilde{\mathbf{y}}|\mathbf{y})$, by

$$p_{\text{POST}}^{\text{PRED}}(\tilde{\mathbf{y}}|\mathbf{y}) = \int p_{\text{SAMPLE}}(\tilde{\mathbf{y}}|\boldsymbol{\theta})p_{\text{POST}}(\boldsymbol{\theta}|\mathbf{y}) d\boldsymbol{\theta}, \quad (12)$$

i.e. the posterior predictive distribution is the marginal distribution for new data $\tilde{\mathbf{y}}$ conditioned on the observed data \mathbf{y} that we obtain by averaging the likelihood over all possible parameter values with respect to the posterior density. We remark that all integrals are to be understood as being over the full range of the variable, which should be clear from context. Further discussion regarding these distributions may be found in Gelman et al. (2014a) and the references therein.

3.2 Model Identifiability

A key consideration in model calibration is identifiability. A model is deemed *identifiable* if it is possible to uniquely determine the values of the unobservable model parameters from the experimental data. Similarly, a model is non-identifiable if multiple parameterizations are observationally equivalent. Identifiability is of crucial importance to the field of clinical predictions because if a model's parameters are not well constrained, the resulting predictions of that model may be subject to an unacceptable degree of posterior uncertainty.

Two types of non-identifiability are distinguishable:

Structural: in which the model structure precludes the identification of parameters irrespective of the data (see, e.g., Cobelli and DiStefano (1980));

Practical: in which the data is insufficient (either in terms of quality or quantity) to identify the parameters.

While it is beyond the scope of this work either to discuss methods for determining structural identifiability, or to provide a wider exposition of practical identifiability, it is important to note that, given an amount of data of a certain quality, it is not necessarily guaranteed that model parameters may be determined unambiguously. Indeed it is often the case that experimental data is insufficient to calibrate even modestly complex mathematical models of biological systems. We refer the reader to Bellman and Åström (1970); Cobelli and DiStefano (1980); Raue et al. (2009) for further discussion.

3.2.1 Selection of Prior Distribution

We now specify the prior distribution of our parameters $\boldsymbol{\theta}$. The prior distribution indicates the degree of belief in the values of the parameters before any measurements are made. Where possible, the choice of prior should

incorporate any quantitative knowledge about the parameters, but may also incorporate subjective expert opinion.

We have no quantitative knowledge about the parameters, or additional expert opinion, other than biologically appropriate bounds on their ranges. It is clear from our biological understanding that V_0 , K and α are all greater than 0. Moreover, from the data presented in Figure 2 it is reasonable to suppose that $V_0 < 0.2\text{mm}^3$ and $K < 5.0\text{mm}^3$. In light of these observations, and the fact we have no further information regarding the parameter values, we take the marginal prior distribution of each parameter to be uniform over the interval given in Table 1, implying that prior distribution of $\boldsymbol{\theta}$ is given by

$$\boldsymbol{\theta} \sim U(0, 0.2) \ U(0.3, 5) \ U(0, 1), \quad (13)$$

where $U(a, b)$ denotes a uniform distribution over the interval (a, b) . As $p_{\text{PRIOR}}(\boldsymbol{\theta})$ is a pdf, it must integrate to 1 and since each parameter is uniformly distributed it must be constant. Therefore, $p_{\text{PRIOR}}(\boldsymbol{\theta})$ satisfies

$$1 = p_{\text{PRIOR}} \int_0^{0.2} \int_{0.3}^5 \int_0^1 1 d\boldsymbol{\theta}, \quad (14)$$

thus implying that $p_{\text{PRIOR}}(\boldsymbol{\theta})$ is given by

$$p_{\text{PRIOR}}(\boldsymbol{\theta}) = \frac{1}{0.94}. \quad (15)$$

The bounds we have chosen for our parameters and assumption of flat

Prior Knowledge			
Parameter	Units	Lower Bound	Upper Bound
V_0	mm^3	0.0	0.2
K	mm^3	0.3	5.0
α	s^{-1}	0.0	1.0

Table 1: Upper and lower bounds on the parameter values employed in specification of the prior distribution.

priors lead to a relatively uninformative prior distribution. As a consequence, the posterior distribution will be determined primarily by the data, via the likelihood function described in Section 3.2.2. If, however, we had some additional knowledge regarding the parameters, we could incorporate this into the prior distribution to (potentially) increase accuracy in our predictions.

We note that our choice of prior is not the only reasonable choice. If, for instance, we were less certain of the upper bound on the parameters we could impose a half-normal or half-Cauchy prior distribution. This would still impose the biologically motivated positivity constraint on the parameters, but would be weakly-informative in terms of determining the posterior distribution.

Further discussion on the choice of priors may be found in, e.g., Gelman et al. (2014a); Simpson et al. (2014).

Finally, we highlight that in the clinical setting, patient-specific data may be sparse due to the cost of imaging etc. In this context, the use of informative priors would provide a means of incorporating population data to potentially increase the accuracy and reliability of a patient-specific computation, a point to which we return to in Section 8.

3.2.2 Selection of the Likelihood

We now specify the likelihood function for the parameter $\boldsymbol{\theta}$, given data \mathbf{y} . In the Bayesian framework, it is the likelihood function that determines how the underlying biological model for the tumour volume given in Section 2.1 and the data, \mathbf{y} (described in Section 2.2), inform the posterior distribution.

We assume that the errors in the measurement of the tumour volume at each time point are independent and that the processes determining the *true* volume are deterministic. Furthermore, we assume that the experimental noise is normally distributed about 0, with variance $\sigma_V(t)$ (where $\sigma_V(t)$ denotes σ_V defined in (6) evaluated at time t). Under these assumptions, the likelihood is given by

$$p_{\text{SAMPLE}}(\mathbf{y}|\boldsymbol{\theta}) = \prod_{i \in S_C} \frac{1}{\sqrt{2\pi\sigma_V^2(t_i)}} \exp\left(-\frac{(y_i - V(t_i; \boldsymbol{\theta}))^2}{2\sigma_V^2(t_i)}\right). \quad (16)$$

In the framework described in Section 3.1, we require that the data are exchangeable. It is clear that the data \mathbf{y} are not themselves exchangeable. However, if we consider time as a covariate, then the set of pairs $\{(y_i, t_i)\}_{i=1}^{14}$ are exchangeable. We refer the reader to Schervish (1995) for a more thorough discussion on exchangeability.

We have assumed the particular form of the likelihood given in (16) based on the assumption of normally distributed measurement errors. However, this may prove to be incorrect. As such, it is important to investigate the robustness of any model predictions to this choice of likelihood, as this determines how the observed data impacts our calibrated model. We address this point further in Section 6.

3.2.3 Selection of the Calibration and Validation Sets

As discussed briefly in Section 2.2, we partition the data obtained at t_1, \dots, t_{13} into two sets, S_C and S_V , for calibration and validation, respectively. When specifying S_C and S_V several factors must be born in mind:

1. S_C should be sufficiently large and contain data of sufficient quality that the model is practically identifiable. In the context of this study, if only data from early time points is chosen (i.e. in the early nutrient-rich growth phase), it is possible that the parameter K may be unidentifiable as this parameter determines the long time behaviour of the system.

2. As with S_C , S_V should also be of sufficient size and quality to result in a practically identifiable model.
3. In line with the hierarchy of data described in NRC (2012); Oden et al. (2010a,b); Hawkins-Daarud et al. (2013), S_V should be of higher quality in the sense that the data is obtained for an experimental setup as close as possible to the predictive case. In our application, this corresponds to including volume data in S_V obtained at a time closer to t_{14} than is included in S_C .

In practical terms, we recommend a preliminary study employing synthetic data, in order to assess:

1. The identifiability of the model given various choices of S_C and S_V ;
2. Whether the validation procedure that results from a given choice of S_V is capable of discerning models, the predictions of which are not acceptable in the context of their intended real-world use.

In light of these considerations, we consider $S_C = \{t_1, \dots, t_{12}\}$ and $S_V = S_C \cup \{t_{13}\}$. We note that there are no strict rules regarding the selection of calibration and validation data. In the reporting of validation experiments the selection of data must be made explicit. The sparsity of data available in this tutorial presents particular challenges, as we would ideally have disjoint calibration and validation sets. However, if we impose this constraint, we typically arrive at the situation where either the calibration or validation posterior is dominated by the prior due to lack of data, thus leading to validated predictions of questionable practical use due to a large posterior uncertainty.

3.3 Sampling of the Posterior Distribution

While for certain combinations of prior distribution and likelihood it is possible to obtain analytical expressions for the posterior distribution, in general this is not the case. As such, we are often required to sample from the posterior distribution $p_{\text{POST}}(\boldsymbol{\theta}|\mathbf{y})$ via a discrete approximation. This sampling process represents a significant computational challenge for complex models with a large number of inferred parameters. For the problem at hand, it is possible for us to sample the posterior distribution employing a regular grid in the parameter space, cf. Hawkins-Daarud et al. (2013); Gelman et al. (2014a). However, we employ here a member of the popular family of methods for sampling the posterior distribution, known as Markov chain Monte Carlo (MCMC), so as to demonstrate how one may perform calibration for a more complex model.

It is beyond the scope of the current work to fully describe the theory associated with MCMC. As such, we refer the interested reader to Gilks et al. (1996); Chib and Greenberg (1995); Kaipio and Somersalo (2006); Gelman et al. (2014a) and the references contained therein, for a more complete discussion. We do, however, present a brief overview of the Metropolis-Hastings

MCMC algorithm, and an adaptive variant employed here. The key idea behind MCMC is to generate a Markov chain whose stationary distribution corresponds to the posterior distribution (10) in our Bayesian formulation. We refer the reader to Norris (1998) for an introduction to Markov chains. The Metropolis-Hastings algorithm (Hastings 1970) itself is a generalisation of a method first employed in Metropolis and Ulam (1949); Metropolis et al. (1953); the algorithm is shown in Algorithm 3.1. We forgo a discussion regarding selection of the proposal distribution J_i and initial distribution p_0 and instead refer the reader to the references provided above. In order to enhance the rate

Algorithm 3.1 Metropolis-Hastings

- 1: Draw $\boldsymbol{\theta}^0$ such that $p(\boldsymbol{\theta}^0) > 0$ from an initial distribution $p_0(\boldsymbol{\theta})$ based on sampling from a regular grid, or some other crude estimate.
 - 2: **for** $i = 1, i_{\max}$ **do**
 - 3: Sample a proposal $\boldsymbol{\theta}^*$ from a proposal distribution at time i , $J_i(\boldsymbol{\theta}^* | \boldsymbol{\theta}^{i-1})$.
 - 4: Calculate the ratio $r = \frac{p(\boldsymbol{\theta}^* | \mathbf{y}) / J_i(\boldsymbol{\theta}^* | \boldsymbol{\theta}^{i-1})}{p(\boldsymbol{\theta}^{i-1} | \mathbf{y}) / J_i(\boldsymbol{\theta}^{i-1} | \boldsymbol{\theta}^*)}$.
 - 5: Set $\boldsymbol{\theta}^i$ to $\boldsymbol{\theta}^*$ with probability $\min\{1, r\}$, or $\boldsymbol{\theta}^{i-1}$ otherwise.
 - 6: **end for**
-

at which the chains generated by the Metropolis-Hastings algorithm converge to the posterior distribution, various classes of adaptive algorithms have been proposed, see e.g. Andrieu and Thoms (2008) and the references therein. In this work, we employ Andrieu and Thoms (2008, Algorithm 4), which permits more rapid movement through regions in parameter space of low probability. A MATLAB implementation of this algorithm applied to the Gompertzian model of tumour growth is available in Connor (2016). In Algorithm 3.2, we provide pseudocode describing the generic form of the algorithm employed here, where $N(\cdot, \cdot)$ denotes a multivariate normal distribution. As Algorithm 3.2 proceeds, the proposal distribution is adapted to achieve more rapid convergence. Again, we forgo discussion regarding precise choices for β and the updates for λ_i , μ_i , and Σ_i and refer the reader to Andrieu and Thoms (2008).

Algorithm 3.2 Generic Adaptive MCMC

- 1: Draw $\boldsymbol{\theta}^0$ such that $p(\boldsymbol{\theta}^0) > 0$.
 - 2: **for** $i = 1, i_{\max}$ **do**
 - 3: Sample a proposal $\boldsymbol{\theta}^*$ from a proposal distribution $N(\boldsymbol{\theta}^{i-1}, \lambda_{i-1} \boldsymbol{\Sigma}_{i-1})$.
 - 4: Calculate the ratio β (defined in a similar manner to r in Alg. 3.1).
 - 5: Set $\boldsymbol{\theta}^i$ to $\boldsymbol{\theta}^*$ with probability $\min\{1, \beta\}$, or $\boldsymbol{\theta}^{i-1}$ otherwise.
 - 6: Compute λ_i , μ_i , and $\boldsymbol{\Sigma}_i$.
 - 7: **end for**
-

Intuitively, the MCMC algorithms presented correspond to generating sequences of points in parameter space by iteratively suggesting movement to new points in parameter space via a proposal distribution, whereby a

movement is accepted or rejected on the basis of the relative likelihood of the current and proposed points. It is in the computation of the acceptance ratios r in Algorithm 3.1 and β in Algorithm 3.2 that we may observe the dependence of the output from the algorithm on the experimental data, via the computation of the likelihood. Moreover, we may observe the potentially huge computational cost, in that each evaluation of the acceptance ratio necessitates a model evaluation. For the simple model considered in the current study this is not too demanding; however, if we were to consider a time dependent PDE model this would represent a significant cost.

As MCMC is an iterative algorithm, we must consider its convergence in the sense of whether the generated points are distributed according to the posterior distribution, as this clearly affects the reliability of any resultant analysis. In particular, if the iterative process has not proceeded for a sufficiently long period of time, then the simulations may not be representative of the target distribution. In order to diminish any dependence on the starting values we discard a number of early iterations as *warm-up*. Moreover, we compute multiple chains so that we may monitor whether ‘in’ chain variation is approximately equal to ‘between’ chain variation as an indicator of convergence.

We now describe, following Gelman et al. (2014a), how we assess the convergence of the algorithm. Let m denote the number of chains and n denote the length of each chain, and for each scalar estimand ψ we identify the simulations as ψ_{ij} , for $1 \leq i \leq n$ and $1 \leq j \leq m$. We now define the between- and within-chain variances, denoted B and W , respectively, as

$$B := \frac{n}{m-1} \sum_{j=1}^m (\bar{\psi}_{\cdot j} - \bar{\psi}_{..})^2, \quad (17)$$

and

$$W := \frac{1}{m} \sum_{j=1}^m s_j^2, \quad (18)$$

where

$$\bar{\psi}_{\cdot j} = \frac{1}{n} \sum_{i=1}^n \psi_{ij}, \quad \bar{\psi}_{..} = \frac{1}{m} \sum_{j=1}^m \bar{\psi}_{\cdot j}, \quad \text{and} \quad s_j^2 = \frac{1}{n-1} \sum_{i=1}^n (\psi_{ij} - \bar{\psi}_{\cdot j})^2.$$

The marginal posterior variance of the estimand $\text{var}(\psi|\mathbf{y})$ may then be estimated by a weighted average of W and B ,

$$\widehat{\text{var}}^+(\psi|\mathbf{y}) := \frac{n-1}{n} W + \frac{1}{n} B. \quad (19)$$

In order to monitor convergence of the algorithm, we estimate the factor by which the scale of the current distribution for ψ might be reduced if the simulations were continued for the limit $n \rightarrow \infty$ using the quantity

$$\hat{R} = \sqrt{\frac{\widehat{\text{var}}^+(\psi|\mathbf{y})}{W}}. \quad (20)$$

A large value of \hat{R} indicates that further simulations may improve the inference about the target distribution of ψ . As such, we specify a tolerance TOL , such that if $\hat{R} < \text{TOL}$ for some $i < i_{\max}$, we view the chains as having converged to the stationary distribution. Regarding the choice of the number of chains and i_{\max} , we refer the reader to the aforementioned references discussing MCMC.

In the proceeding examples we perform computations employing the following:

- We count the first 10,000 iterations as ‘warm up’ and do not include these points in the sample;
- We compute 3 chains ($m = 3$);
- We compute at least 25,000 iterations, but no more than 160,000 iterations ($25,000 < n < 160,000$); and
- We specify the TOL as 1.05.

Any further details required to reproduce our computations may be found in the complete code and data employed in the current work, available in the online resource Connor (2016).

3.4 Model Calibration for the Gompertzian Model

We proceed now by calibrating the Gompertzian model of tumour growth described in Section 2.1 with the experimental data described in Section 2.2. Figure 3 shows discrete approximations of the marginal posterior distributions for $\boldsymbol{\theta}$, obtained from draws of the posterior distribution generated by the adaptive MCMC algorithm, Algorithm 3.2. This corresponds to samples from the distribution defined in (10) obtained under the assumption of prior distribution (15) and likelihood (16). From this figure, we see that the marginal posterior distributions for V_0 , K , and α are unimodal, and are centred around values that are not close to the bounds we imposed in the definition of the prior distributions (thus indicating our assumption on the prior distribution is not inherently inconsistent with the data). Moreover, there are no issues surrounding identifiability of the parameters.

4 Model-Data Consistency

In Section 3 we calibrated the Gompertzian model introduced in Section 2.1, i.e. we obtained a posterior distribution $p_{\text{POST}}(\boldsymbol{\theta}|\mathbf{y})$ that combines our prior knowledge surrounding the unobservable parameters $p_{\text{PRIOR}}(\boldsymbol{\theta})$ and observed data \mathbf{y} in the calibration data set S_C . The first stage of the validation procedure we set out in this work is to verify whether outputs from the calibrated model are consistent with the observed calibration data. Or rather, does the calibration data look plausible under the posterior predictive distribution? In this section, we describe a selection of data misfit tests that seek to ascertain whether there are systematic differences between the

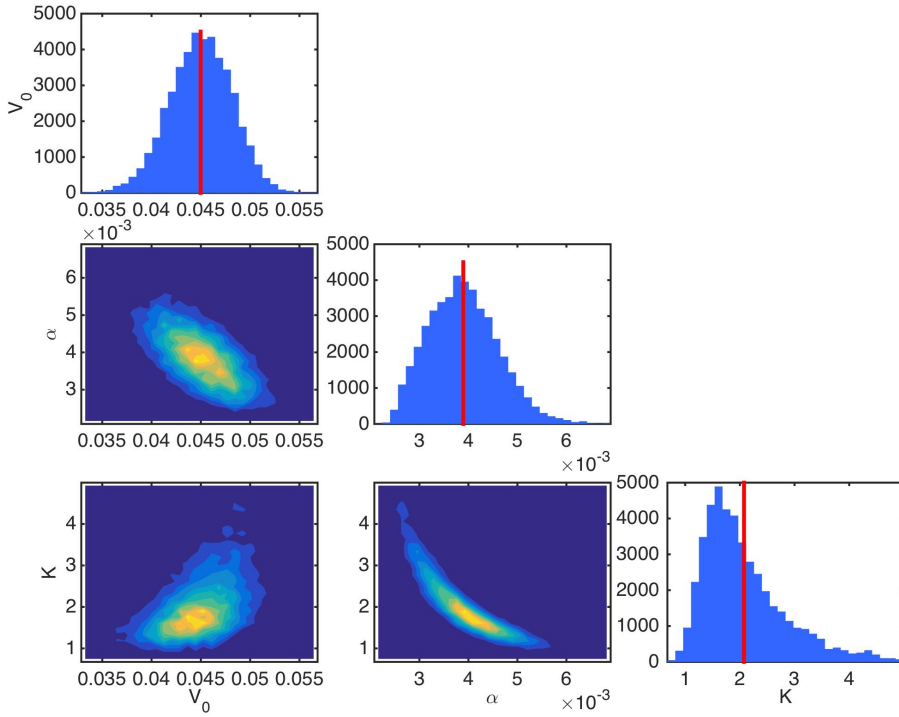


Fig. 3: Approximations of a range of marginal and joint posterior pdfs for V_0 (mm^3), K (mm^3), and α (s^{-1}) obtained via application of Algorithm 3.2 implemented in Connor (2016) to compute the joint posterior distribution (10). These approximations are computed employing the prior distribution (15) and likelihood (16), together with the calibration data S_C . The vertical line corresponds to the mean of the marginal posterior distribution for each parameter.

calibrated model outputs and the calibration data. We first describe graphical checks, and then describe more quantitative posterior predictive p -values test. We apply each methodology to our calibrated model from Section 3. We refer the reader to the bibliographic note in Gelman et al. (2014a, Chp. 6) for further references.

As noted in Hawkins-Daarud et al. (2013); NRC (2012), it is important to realise that we may never fully validate a model. The strongest statement we can make is that under certain tests, the model has not been invalidated. This observation is key when interpreting the implications of passing the model fit tests described in this section, or the model validation checks described in Section 5.

In each of the tests described below, we require the concept of replicated data, i.e. data that could have been observed, or data that could be obtained were we to perform the experiment again. Again, we follow the notation of Gelman et al. (2014a), and distinguish between the replicated data \mathbf{y}^{rep} and the notation for general predictive outcomes $\tilde{\mathbf{y}}$. We take the distribution for \mathbf{y}^{rep}

to be the current state of knowledge in the posterior predictive distribution, i.e.

$$p_{\text{POST}}^{\text{PRED}}(\mathbf{y}^{\text{rep}}|\mathbf{y}) = \int p_{\text{SAMPLE}}(\mathbf{y}^{\text{rep}}|\boldsymbol{\theta}) p_{\text{POST}}(\boldsymbol{\theta}|\mathbf{y}) d\boldsymbol{\theta}. \quad (21)$$

4.1 Graphical Checks

The main idea of graphical checks is to display the observed data with replicated data obtained from the calibrated model in order to assess whether there are any systematic discrepancies between the real and simulated data. In Gelman et al. (2014a), the authors describe three kinds of graphical display (direct, summary or parameter inference, and model-data discrepancy); however, given the relative simplicity of the model here we consider only direct display of the data against a collection of replications.

Figure 4 shows 5000 replications drawn from the posterior predictive distribution. From the figure, it appears as though there are no large systematic discrepancies between our observed data \mathbf{y} and the replicated data \mathbf{y}^{rep} .

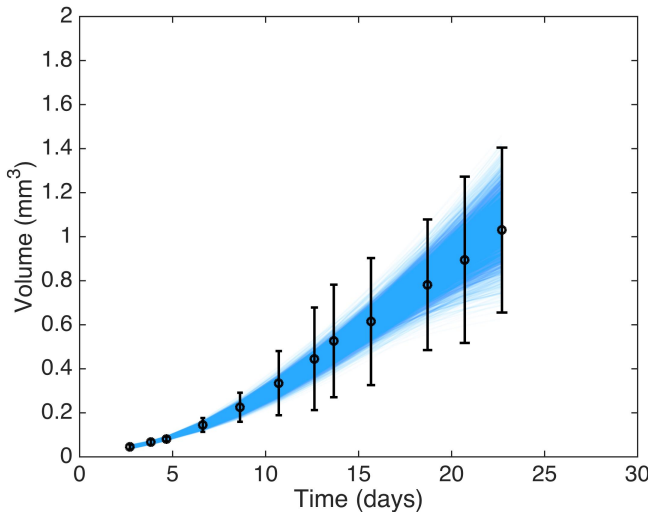


Fig. 4: Experimental data for times $\{t_1, \dots, t_{12}\}$ and errors bars showing $\pm 2\sigma_V$, together with 5000 replications obtained from the posterior predictive distribution for the calibration model. Replication data obtained by evaluating (1) at 5000 points drawn from the posterior distribution (10) obtained via application of Algorithm 3.2 with prior distribution (15) and likelihood (16), with the calibration data S_C .

4.2 Posterior Predictive p -Values

In measuring the discrepancy or degree of fit of the model to the data, it is necessary to define appropriate test quantities that we may check. A test quantity, or discrepancy measure, $T(\mathbf{y}, \boldsymbol{\theta})$, is a scalar summary of parameters and data that we may use as a standard for comparing data to predictive simulations. These quantities are analogous to test statistics in classical statistical testing.

Posterior predictive p -values in the Bayesian framework are defined as the probability that the test quantity for the replicated data \mathbf{y}^{rep} is more extreme than for the observed data, i.e.

$$\begin{aligned} p_B &= \Pr(T(\mathbf{y}^{\text{rep}}, \boldsymbol{\theta}) \geq T(\mathbf{y}, \boldsymbol{\theta}) | \mathbf{y}), \\ &= \int \int \mathbb{I}_{T(\mathbf{y}^{\text{rep}}, \boldsymbol{\theta}) \geq T(\mathbf{y}, \boldsymbol{\theta})} p_{\text{SAMPLE}}(\mathbf{y}^{\text{rep}} | \boldsymbol{\theta}) p_{\text{POST}}(\boldsymbol{\theta} | \mathbf{y}) d\mathbf{y}^{\text{rep}} d\boldsymbol{\theta}, \end{aligned} \quad (22)$$

where \mathbb{I}_A denotes the indicator function for event A .

We view a model as being questionable if p_B takes a value that is close to 0 or 1, indicating that it would be unlikely to observe \mathbf{y} in the replications of the data, if the model were true. Extreme values for p_B indicate significant discrepancies in the model that need to be addressed by expanding the model appropriately or altering assumptions in the model. However, finding an extreme value for p_B and deeming the model as suspect should not signal the end of the analysis. It will often be the case that the nature of the failure will suggest improvements to the model or identify certain data that are subject to additional error.

For our application, we consider the tumour volume at $\{t_1, t_2, \dots, t_{12}\}$ as the test quantities. In Table 2, we show the Bayesian p -values obtained for the tumour volume at each time point, for the posterior predictive distribution. We observe that there are no extreme values for p_B , and, as such, conclude that our model is not inconsistent with the observed data.

Time	p_B	$0.01 \leq p_B \leq 0.99$	Time	p_B	$0.01 \leq p_B \leq 0.99$
t_1	0.5162	True	t_7	0.2462	True
t_2	0.2440	True	t_8	0.1214	True
t_3	0.7544	True	t_9	0.4610	True
t_4	0.2830	True	t_{10}	0.6300	True
t_5	0.3108	True	t_{11}	0.6512	True
t_6	0.2360	True	t_{12}	0.5634	True

Table 2: Posterior predictive p -values at each time point computed employing the calibration model.

5 Model validation and prediction

Now that the calibrated model of Section 3 has passed the data consistency checks set out in Section 4, we investigate its predictive properties by attempting to validate it against a validation model that has been calibrated using the validation data, S_V . Recall that validation is the process of determining the degree to which a model represents the real world in the context of its real world uses. In this section, we explain how we validate our model by comparing the calibrated model against the validation model, based on the procedure set out in Hawkins-Daarud et al. (2013), and demonstrate its application to our example.

5.1 Model Validation Procedure

The validation data, S_V , contains additional information regarding the behaviour of the physical system that should lead to a more accurate model of tumour growth. We denote the data in the validation set by \mathbf{y}^{VAL} . The first stage of the validation process is to calibrate the Gompertzian model of tumour growth described in Section 2.1 against the validation data set to obtain a validation posterior density, denoted $p_{\text{POST}}^{\text{VAL}}(\boldsymbol{\theta}|\mathbf{y}^{\text{VAL}})$, and validation posterior predictive distribution, denoted $p_{\text{PRED}}^{\text{VAL}}(\hat{\mathbf{y}}|\mathbf{y}^{\text{VAL}})$, which are defined analogously to (10) and (12), respectively, but are instead calculated employing the data in S_V .

The next stage of the validation is to test whether the validation posterior predictive distribution passes the model-data consistency checks described in Section 4. If so, we proceed by testing whether the knowledge gained from the new data is very different from that obtained in the calibration experiments. To this end, we denote the pdf for our predictive quantity of interest, as described in Section 2.3, and obtained by calibrating the model with S_C and S_V , by p_C and p_V respectively. We say that the model is not invalidated if

$$M(p_C, p_V) \leq \text{THRESHOLD}, \quad (23)$$

where M is some appropriate metric, and **THRESHOLD** denotes a validation threshold we specify *a priori*. That is to say, our model is not invalidated if the posterior predictive distributions of our quantity of interest for the calibration and validation models are close in some appropriate sense. We recall our earlier remark that this is the strongest statement we might make, and further highlight that this statement is valid for our specified QoI only.

While there are many other appropriate choices of metric, in Hawkins-Daarud et al. (2013) the authors consider

$$M_1(p_C, p_V) = \sup_{y \in [\gamma_1, \gamma_2]} |F_C^{-1}(y) - F_V^{-1}(y)|, \quad (24)$$

where F_C and F_V denote the cumulative distribution functions (cdfs) of the calibration and validation models, respectively, and γ_1 and γ_2 are chosen to

exclude comparison of the tails of the distributions. Alternatively one could test the information gain from the additional data using such measures as the Kullback-Leibler divergence (Kullback and Leibler 1951). In addition to the metrics referenced in Hawkins-Daarud et al. (2013), we remark that it is possible to compare the empirical cdfs for the calibration and validation model via a two-sample Kolmogorov-Smirnov (KS) test (Massey 1951; Miller 1956). Furthermore, techniques for estimating out-of-sample predictive accuracy of the model may be applied as a possible means of comparing the calibration and validation models. In particular, formulae such as the Akaike information criterion (AIC) (Akaike 1973), deviance information criterion (DIC) (Spiegelhalter et al. 2002), and Watanabe-Akaike (or widely-available) information criterion (WAIC) (Watanabe 2010) provide means of estimating the predictive accuracy of the model in an approximately unbiased manner (see Gelman et al. (2014b) for a detailed discussion of these techniques). However, we seek to make extrapolative predictions, so the AIC, DIC, and WAIC are less applicable than if we were making interpolative predictions.

In this work, we compare the validation and calibration posterior predictive pdfs for the QoI via the test statistic of the two-sample KS test, in which the tails of the distributions are discounted (for details, see Connor (2016)), i.e. the quantity

$$M_2(p_C, p_V) = \sup_{x \in [\eta_1, \eta_2]} |F_C(x) - F_V(x)|, \quad (25)$$

for suitably chosen $\{\eta_1, \eta_2\}$. Furthermore, we set **Threshold** to 10%. In Section 7, we propose a possible method for selecting **Threshold** to maximize predictive accuracy of the validation procedure by means of comparison to multiple repeated experiments ².

Other possible validation procedures have been discussed in the literature (e.g. one-step forecast with re-estimation, multi-step forecast with/without re-estimation, 3-way cross-validation etc.). A complete discussion of these techniques is beyond the scope of this work; as such, we refer the reader to Arlot and Celisse (2010) and NRC (2012) and the references therein for a thorough review.

5.2 Validation of Gompertzian Model

The first stage in the validation of our calibrated model described in Section 3 is to obtain the validation model by calibrating the Gompertzian model of tumour growth, as described in Section 2.1, against the validation data S_V . When calibrating the validation model we employ the same prior distribution (13) and likelihood function (16) as for the calibrated model of Section 3. As we chose flat priors for the parameters and the posterior marginal pdfs in Figure 3 were far from the bounds on the parameters imposed through the prior, there is no reason to choose an alternative prior for the validation model.

² While repeated experiments are not available in the patient-specific clinical setting, we may view this as data from multiple individual patients in a similar population.

Furthermore, we chose the same likelihood function because here we wish to assess whether the knowledge gained from the validation data differs from that in the calibration experiments, rather than the sensitivity of the predictions to choice of likelihood function, a point we address in Section 6.

Figure 5 shows discrete approximations of the marginal posterior distributions for $\boldsymbol{\theta}$ obtained from draws of the posterior distribution generated by the adaptive MCMC algorithm, based on the validation dataset S_V . As was also the case for the calibrated model, we see that the distributions are unimodal and are not close to the bounds imposed in the definition of the prior.

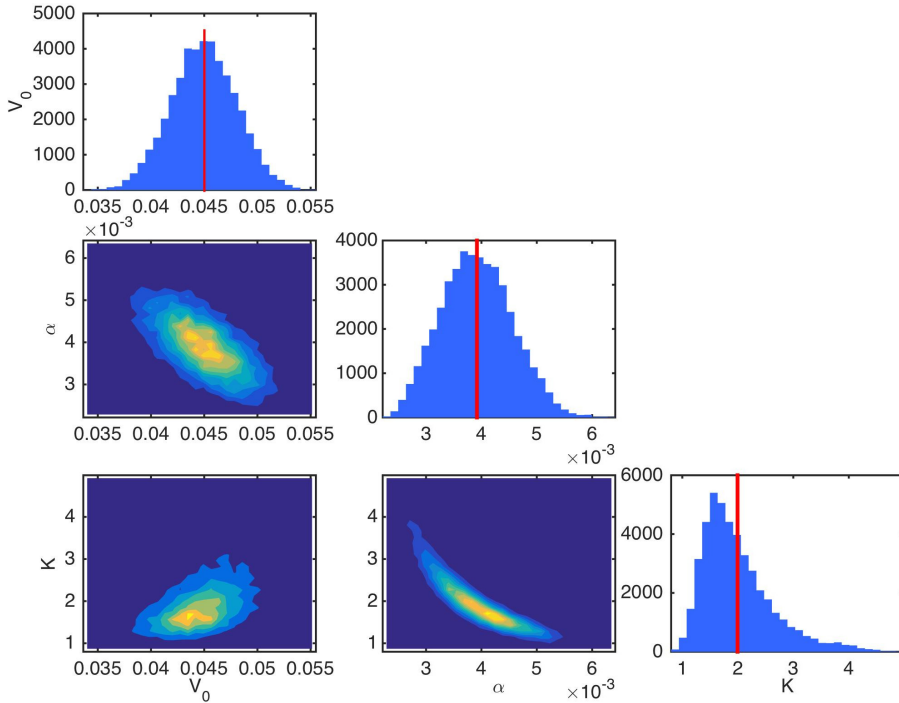


Fig. 5: Approximations of a range of marginal and joint posterior pdfs for V_0 (mm^3), K (mm^3), and α (s^{-1}) obtained with the validation data S_V . Results obtained via same methods as those in Figure 3. Vertical line indicates the mean of the marginal posterior distribution for each parameter.

Now we have obtained our validation model, we perform the same tests of model fit as for the calibration model outlined in Section 4, i.e. we assess whether the validation data is likely to occur as a replication obtained via the validation model posterior predictive distribution. Figure 6 presents 5000 replications of the data drawn from the posterior predictive distribution of the validation model, shown together with the experimental data and error bars

corresponding to $\pm 2\sigma_V$. Once more, we see no large structural discrepancies between the replications and the experimental data. In Table 3 we present the Bayesian p -values for the tumour volume at each of the 13 time points in the validation data set. As with the calibration data, we see no extreme p -values, and hence conclude that the posterior predictive distribution is not inconsistent with the observed experimental data.

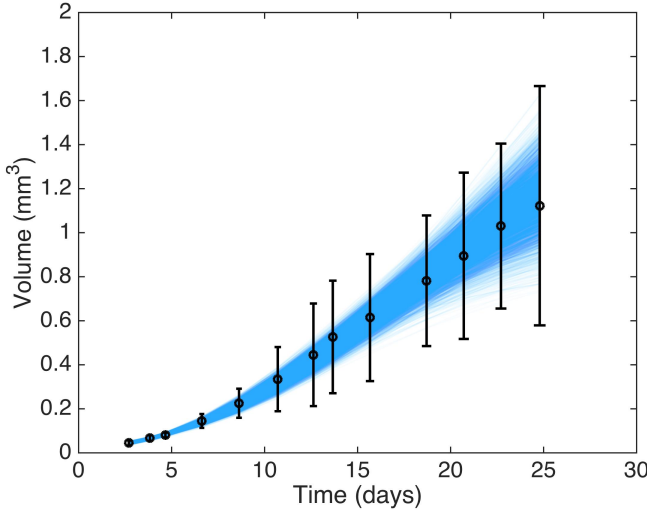


Fig. 6: Experimental data for times $\{t_1, \dots, t_{13}\}$ with error bars showing $\pm 2\sigma_V$, together with 5000 replications obtained from the posterior predictive distribution for the validation model. Replication data obtained by evaluating (1) at 5000 points drawn from the posterior distribution (10) obtained via application of Algorithm 3.2 with prior distribution (15) and likelihood (16), with the validation data S_V .

Time	p_B	$0.01 \leq p_B \leq 0.99$	Time	p_B	$0.01 \leq p_B \leq 0.99$
t_1	0.4976	True	t_8	0.1134	True
t_2	0.2316	True	t_9	0.4554	True
t_3	0.7720	True	t_{10}	0.6350	True
t_4	0.2838	True	t_{11}	0.6530	True
t_5	0.3138	True	t_{12}	0.5548	True
t_6	0.2334	True	t_{13}	0.5964	True
t_7	0.2400	True			

Table 3: Posterior predictive p -values at each time point computed employing the validation model.

Given that we have judged our validation model as suitably fitting the validation data based on the graphical checks and the posterior predictive p -

values, we proceed now to compare the posterior predictive distributions for the QoI set out in Section 2.3, i.e. we assess whether the predictive properties of our model are suitable for the proposed application. Figure 7 shows a comparison of the experimental data and errors, along with model predictions of the calibration and validation models (represented by the mean, with error bars corresponding to twice the standard deviation), and the experimental data for the tumour spheroid at t_{14} .

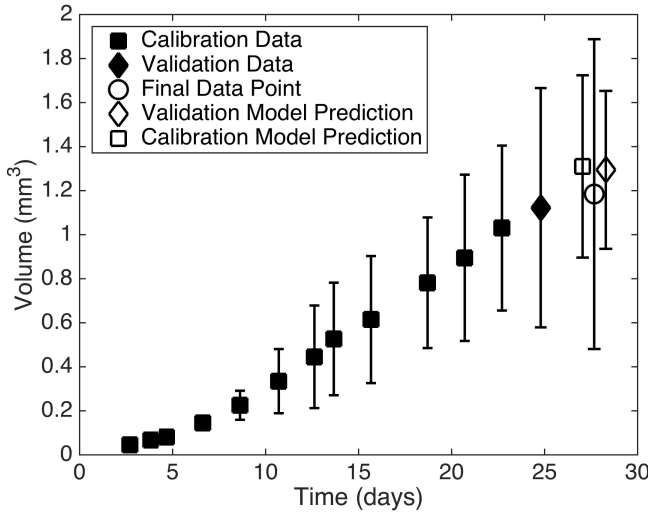


Fig. 7: Summary of all experimental data, errors, and model predictions. The data presented at the final time point has been artificially perturbed so that both model predictions and the experimental data are visible.

Figure 8 shows a discrete approximation of the posterior predictive pdf for our QoI obtained with both the validated calibration model and the validation model. In addition, as a comparison, we have also included the value of our QoI obtained from experiment. From Figure 8 we see that the observed tumour volume at t_{14} appears likely under the posterior predictive distribution, i.e. the model has made a good prediction for the data we have excluded in the calibration and validation procedure.

The value of the test statistic $M_2(p_C, p_V)$ computed for the calibration and validation models was approximately 6%, and as such, we deem the model not invalidated under the procedure set out above.

6 Sensitivity Analysis

While we have shown in Section 5 that our calibration model is not invalid under the validation procedure set out above, we have made several

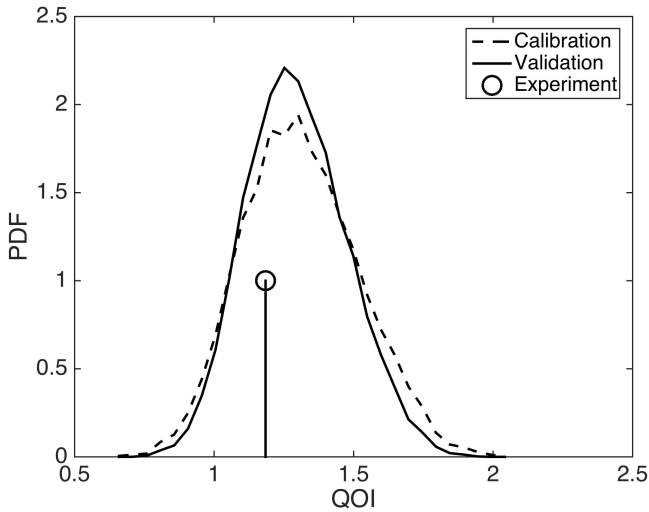


Fig. 8: Posterior predictive pdf for the QoI obtained employing the calibration and validation models, with spheroid volume obtained at t_{14} .

assumptions in the development of the statistical model which could be replaced with others that appear equally valid *a priori*. For instance, there are many appropriate choices for error measurement model, other than that detailed in Section 2.2.3, which may affect the predictive capability of the calibrated model. Similarly, the choice of normally distributed errors for the likelihood function, and the choice of prior, all hold influence over the model predictions.

In *robust Bayesian analysis*, a prediction is viewed as robust if it does not depend sensitively on the assumptions and inputs on which the model is based. Robust Bayesian methods address the difficulty associated with defining precise priors and likelihoods (Berger 1984; Pericchi and Pérez 1994; Insua and Ruggeri 2000; Lopes and Tobias 2011). It is beyond the scope of this work to perform a full robustness analysis. However, possible means of making the present analysis more robust include replacing the normally distributed errors in the likelihood with a Student's t -distribution, or a more flexible likelihood such as a mixture of normals.

As an example, we focus here on the inference of the length of third axis ℓ_3 , in the error calculation. In Section 2.2.3, we assume a given standard deviation in ℓ_3 . This will affect the resultant variation in our model predictions, possibly increasing the posterior uncertainty in the prediction of the QoI. In order to test the sensitivity of our prediction to this assumption, we now introduce an additional parameter, s_e , which we refer to as the *experimental scale*, that may be calibrated with experimental data. That is, we now consider $\theta = (V_0, K, \alpha, s_e)$. This parameter is introduced into the likelihood via an

alternative definition of the variance $\hat{\sigma}_V^2$, which we define as

$$\hat{\sigma}_V = \sqrt{\left(\frac{\partial V}{\partial \ell_1}\right)^2 \sigma_{\ell_1}^2 + \left(\frac{\partial V}{\partial \ell_2}\right)^2 \sigma_{\ell_2}^2 + \left(\frac{\partial V}{\partial \ell_3}\right)^2 \hat{\sigma}_{\ell_3}^2}, \quad (26)$$

where $\hat{\sigma}_{\ell_3} = s_e(\ell_1 - \ell_2)$. Assuming that the prior distribution for s_e is $U[1 \times 10^{-7}, 0.5]$, we may compute the posterior marginal distribution for s_e as shown in Figure 9. From this figure, we can observe a modal value that is of the order 10^{-3} , thus indicating we may have overestimated the variation in ℓ_3 (as previously, there was an implicit definition that $s_e = 1/2$, given the Gompertzian model for tumour growth and our experimental data. We proceed

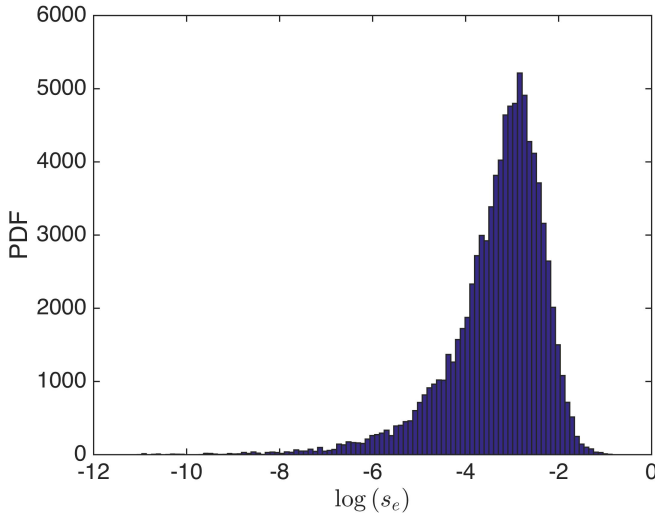


Fig. 9: Posterior marginal pdf (unscaled) for the experimental scale parameter, s_e .

now by assessing the fit of our enhanced model calibrated on S_C to the data and the validity of this model. This process is performed as described in Sections 4 and 5. Figure 10 shows 5000 replications with the experimental data and error bars corresponding to $\pm 2\hat{\sigma}_V$, in which we fix s_e at its modal value. Here, we see much tighter error bars on the experimental data, and greatly reduced variation in the replications when compared to Figure 4. However, here we observe that the data at t_8 lies outside of the replications. This is a concern; however, as we wish to predict late time behaviour, we place greater emphasis on the fact that the data and replications appear consistent for t_9 to t_{12} . Table 4 shows the Bayesian p -values for the data, based on the replications shown in Figure 10. The p -values confirm the inconsistency at t_8 and further highlight additional inconsistency at t_3 , which is not visible in the graphical data. However, we again proceed on the basis of the intended use of the model.

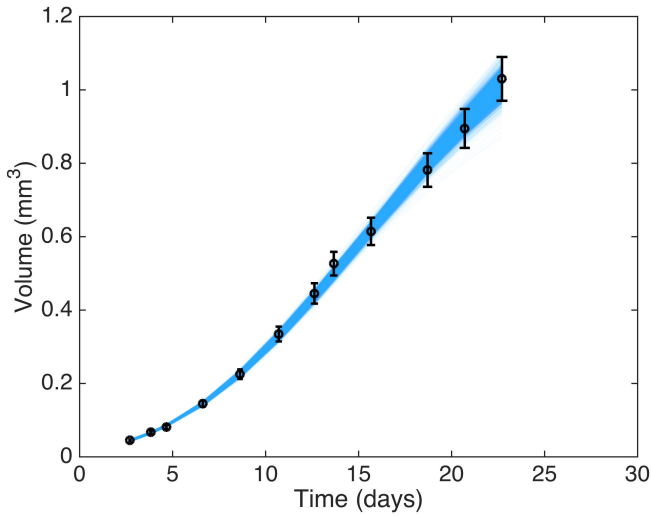


Fig. 10: Experimental data for times $\{t_1, \dots, t_{12}\}$ with error bars showing $\pm 2\hat{\sigma}_V$ employing the modal value for s_e , together with 5000 replications obtained from the posterior predictive distribution for the calibration model with experimental scale estimate for tumour volume.

Time	p_B	$0.01 \leq p_B \leq 0.99$	Time	p_B	$0.01 \leq p_B \leq 0.99$
t_1	0.2500	True	t_7	0.1312	True
t_2	0.0538	True	t_8	0.0006	False
t_3	0.9992	False	t_9	0.7858	True
t_4	0.5422	True	t_{10}	0.9396	True
t_5	0.6720	True	t_{11}	0.8500	True
t_6	0.2066	True	t_{12}	0.2216	True

Table 4: Posterior predictive p -values at each time point computed employing the calibration model.

We now perform the validation procedure set out in Section 5 for this enhanced model. Figure 11 shows a summary of all data, errors, and model predictions (cf. Figure 7) and Figure 12 shows the posterior predictive pdf for the QoI for the calibration and validation models (cf. Figure 8). The key difference we observe in this model, compared to the original, is the reduction in posterior variation of the QoI. However, when we evaluate the test statistic, we see $M_2(\hat{p}_C, \hat{p}_V) \approx 12\%$ and, as such, we deem this prediction invalid under the choice of **THRESHOLD** employed in Section 5.

As the focus of this work is to provide a pedagogical example, we proceed no further with this analysis; in practice, one might continue the analysis by investigating the sensitivity of all assumptions in the model (perhaps comparing models, via Bayes factors (Gelman et al. 2014a) for instance). However, we remark that this example serves to highlight potential difficulties

associated with obtaining accurate and reliable models in more complex settings, and the importance of reliable interpretation of any analysis.

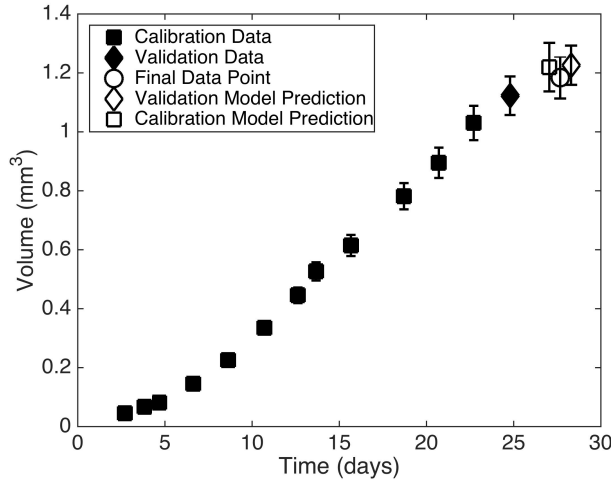


Fig. 11: Summary of all experimental data, errors, and model predictions for model with experimental scale estimate for tumour volume. The data presented at the final time point has been artificially perturbed so that both model predictions and the experimental data are visible.

7 Selection of Threshold

When the experimental data on which the previous sections were based was collected, 9 additional experiments were carried out on other spheroids from the same cell line. In this section, we use the data obtained from those additional experiments to assess the validity of our calibration model with experimental scale in a more informed manner. To this end, we adopt a simple learning approach based on assessing the accuracy of models obtained from calibration against additional sets of experimental data. Figure 13 shows data obtained from 9 additional experiments, which we now employ to tune **THRESHOLD**.

In this work, we judge a model to be valid (that is to say, not invalid) if the test statistic of the two-sample KS test applied to the calibration and validation posterior predictive PDFs of our QoI is not greater than some threshold, T_{KS} . Previously, we have employed the notation **THRESHOLD** for this quantity; we now depart from this notation to highlight that this is now a parameter that may be tuned to enhance the accuracy of the modelling/validation procedure set out here. Separately, a prediction obtained from a calibrated model is judged to be good if, after the QoI is measured

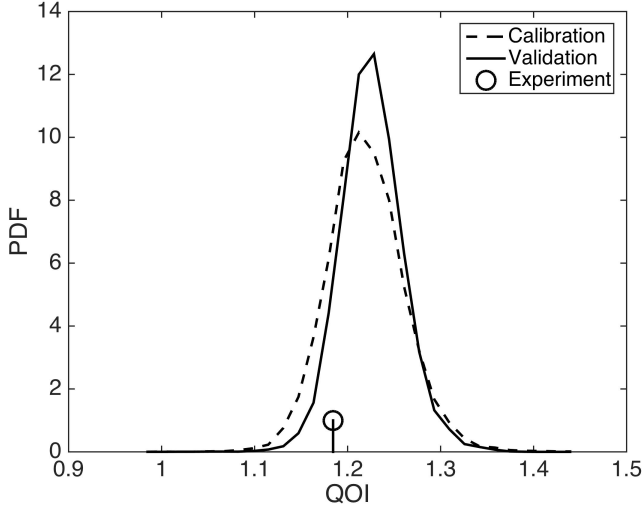


Fig. 12: Posterior predictive pdf for the QoI obtained employing the calibration and validation models with experimental scale estimate for tumour volume, together with spheroid volume obtained at t_{14} .

experimentally, that measurement value lies within the credible interval of the predicted QoI (details of how the credible interval is calculated can be found in Connor (2016)). In an ideal validation procedure, valid models would result in good predictions, whereas invalid models would not. As such, we seek here to jointly maximize

- i) The number of not invalid models for which the observed QoI falls within the credible interval of the predicted QoI, and
- ii) The number of invalid models for which the observed QoI falls outside the credible interval of the predicted QoI,

by tuning the threshold, T_{KS} . Table 5 describes the terms associated with all four possible combined outcomes of the prediction-experimental comparison.

	QoI Credible	QoI not Credible
Model not invalid	True Positive	False Positive
Model invalid	False Negative	True Negative

Table 5: Possible outcomes relating to validation process and experimental data.

We now proceed by performing the model calibration procedure using both the calibration and validation data sets, S_C and S_V , respectively, for each of the nine additional experimental data sets. For each experimental data set (discounting those for which the model is unable to adequately replicate the calibration data) we compare the calibration and validation posterior predictive PDFs of the QoI, calculating the two-sample KS test statistic. Further, for each model we check whether the measured QoI lies within the credible interval of the calibration posterior predictive PDF of the QoI. We then select $T_{KS} \in [0, 1]$, to maximize the accuracy of the prediction, defined by

$$\text{Accuracy} = \frac{(\# \text{ True Positive}) + (\# \text{ True Negative})}{(\text{Total } \# \text{ experiments})}. \quad (27)$$

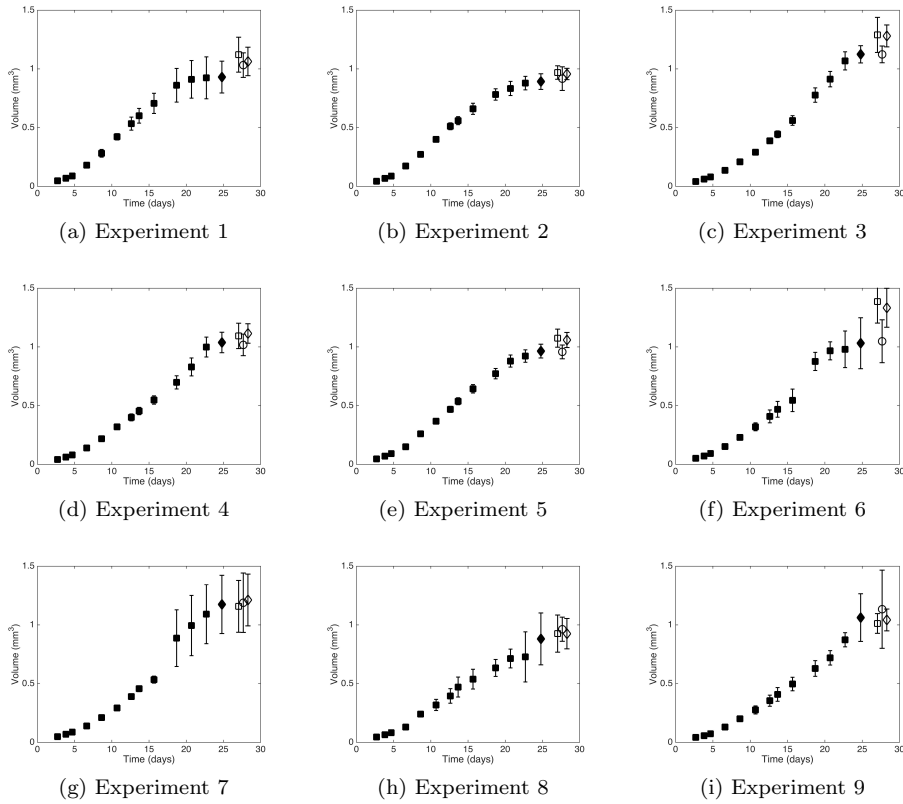


Fig. 13: Summaries of all experimental data, errors, and model predictions obtained from repeat experiments (cf. Figures 7 and 11). The legend for these plots is identical to that in Figures 7 and 11. The data presented at the final time point has been artificially perturbed so that both model predictions and the experimental data are visible.

We may then use the tuned threshold, T_{KS} , to assess the validity of the model obtained by calibrating the Gompertzian model against the initial experimental data set in a more informed manner. Through this process of evaluating the accuracy for multiple experiments for which data for the QoI is available, we are able to ameliorate some of the arbitrary nature associated with the choice of **THRESHOLD**.

Figure 14 demonstrates the variation of accuracy with T_{KS} , having neglected experiments 4, 6, and 7 on the basis of model-data consistency checks (as described in Section 4). We may observe that accuracy is optimized for values of T_{KS} between 7% and 18%. Moreover, if we choose **THRESHOLD** of 15%, then the calibration model obtained in Section 6 is now not invalid. This seems reasonable given the experimentally measured value for our predictive QoI and the PDF obtained from the calibration and validation models, as compared in Figure 12.

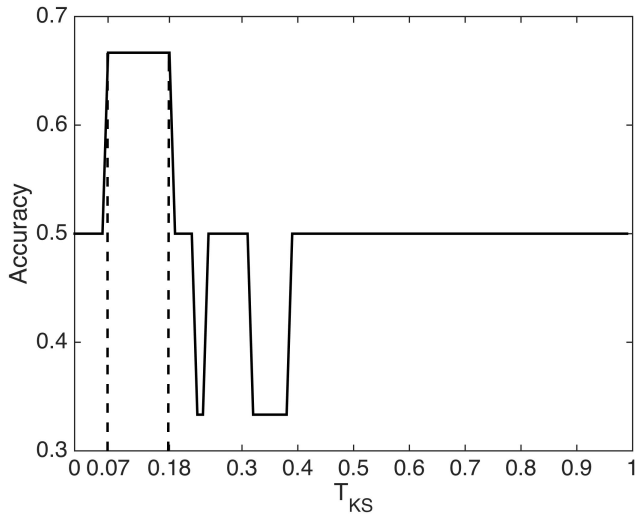


Fig. 14: Accuracy of the validated model for varying T_{KS} . Vertical dashed lines highlight the region for which the greatest accuracy is obtained.

7.1 Summary of the Calibration and Validation Process

In the preceding sections we have set out a process of calibration and validation, which, for an individual experiment, may be summarised as follows. Firstly, we calibrate a mathematical model against partitioned experimental data, employing the Bayesian approach, to obtain so-called calibration and validation posterior predictive distributions for a biological quantity of interest. We then compare these two distributions using an appropriate metric to assess the validity of our calibration model predictions. The **THRESHOLD** employed in the validation procedure is chosen based on an elementary optimization procedure that compares model predictions to experimental data obtained for the biological quantity of interest for similar, prior experiments. Figure 15 presents a flow chart highlighting the process described in this work.

The method outlined above is immediately transferable to more clinically relevant applications. In particular, the simple learning technique introduced in the previous section could be valuable in a clinical setting, in which model predictions of tumour growth (and response to therapy) could be compared with clinical outcomes to judge the accuracy and validity of a prediction for a range of **THRESHOLD** values. Such comparisons could then be used, as in our simple example, to improve the acceptance criterion for not invalid models for future patients. With further data, then, we could better establish which model predictions we should trust and which we should discard.

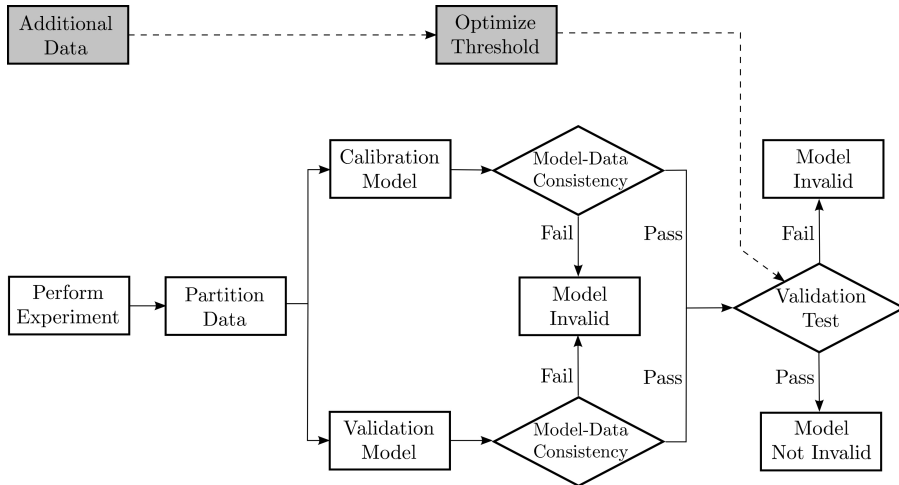


Fig. 15: Schematic diagram demonstrating the calibration and validation process.

8 Discussion

In Sections 2 - 7 we have presented a simple example of calibration, validation and uncertainty quantification for predictive modelling of tumour growth in a Bayesian framework. For this discussion, however, we depart from the example presented above and further discuss the wider application of these techniques, with a view to making patient-specific predictions in the clinic for the purpose of therapy planning.

As highlighted in Pathmanathan and Gray (2013), it is of vital importance to assess the reliability and accuracy of any model that might be used in a safety-critical application in the clinic. The Bayesian framework provides advantages over the frequentist framework in that it requires no notion of repeated experiments on an identical population, which is problematic for patient-specific predictions. Moreover, it provides advantages over methods which result in a single value for model parameters as opposed to distributions, even if confidence intervals for the parameters are supplied. A point we have not yet addressed in this article is the importance of presenting appropriate information regarding uncertainty to decision makers. While expectations and variances (covariances) provide full descriptions in the case of Gaussian random variables (fields), if the distribution is far from Gaussian this may be potentially insufficient. Furthermore, as the complexity of the underlying mathematical model increases, for instance to nonlinear partial differential equations, calibration to a single parameter value appears inadequate, as there may be significant skew or long tails in the distribution of the outputs, as a result of the nonlinearity, which could affect decisions. As such, we question whether in order to make well-informed clinical decisions, more information

regarding the uncertain outputs of the model is required than is attainable via classical means of calibration.

That is not to say these methods are without flaws. It may be the case that there is insufficient data available to properly identify the distributions of the parameters given complex models and expensive and/or noisy data acquisition (via medical imaging for instance). In fact, for even the simplest models (such as that considered here) it is likely that there would be insufficient data to parameterize a personalised patient model. For example, there may be only two data points available: the tumour volume at diagnosis and pre-treatment. In such case, the Bayesian approach provides a natural framework for incorporating population data via an informative prior, as described in Achilleos et al. (2013, 2014), or via expert opinion to constrain the priors.

In addition to the validation and uncertainty quantification procedures described here, in a clinical setting there is a vital need for verification of the computational model. We highlight two forms of verification here: software verification (i.e., is the computational model correctly implemented?), and solution verification (i.e., is the solution of the computational model sufficiently close to the solution of the mathematical model?). For the model considered in this work there are analytical solutions to the deterministic problem, and thus verification is not of vital importance. However, for more sophisticated models requiring the solution of PDEs verification is extremely important. It is beyond the scope of the current work to discuss verification fully, and we refer to NRC (2012) for a more complete introduction to these fields.

The computational cost of the methods and model implemented in the course of this work is low. However, as the complexity of the underlying mathematical model increases, so does the computational cost. As such, the construction of surrogate models via Gaussian process emulation (Kennedy and O'Hagan 2001), (generalized) polynomial chaos expansions (Ghanem and Spanos 1991; Ghanem and Red-Horse 1999; Ghanem 1999; Xiu and Karniadakis 2002, 2003; Najm 2009; Babuška et al. 2004), or stochastic collocation (Xiu and Hesthaven 2005; Tatang et al. 1997; Nobile et al. 2008a,b; Babuška et al. 2007) for instance, and model reduction techniques are extremely important. Again, it is beyond the scope of this work to review these fields (we refer to NRC (2012) and the references therein for a full discussion).

If computational models are to be integrated into clinical practice in order to make personalised predictions about tumour growth and treatment for models of greater complexity than that presented here, successful application of verification and emulation techniques will be of critical importance, in addition to the calibration, validation and uncertainty quantification techniques described here.

9 Conclusions

In this article we have presented an educational example in which we calibrate a simple mathematical model of tumour growth against experimental

data subject to measurement errors, and subsequently validate the model predictions. Moreover, we present an elementary learning approach for determining the validation threshold to maximize the predictive accuracy of the model. Despite the simplicity of the mathematical model, and the fact the experimental data was obtained *in vitro*; we feel that this example illustrates clearly how these methods might be applied to patient-specific models in the clinic.

There are many natural extensions to the work in this article. For example, the techniques we have presented could be applied to more complex models of tumour growth and spatially-resolved MRI and PET data from cancer patients undergoing treatment (Baldock et al. 2013). The use of more complex models will likely require the incorporation of surrogate models and/or model reduction. Finally, it is natural to consider the application of verification techniques in addition to the calibration, validation and uncertainty quantification described here.

Acknowledgements

J. Collis and M. E. Hubbard acknowledge the support of EPSRC grant number EP/K039342/1. This project has received funding from the European Unions Seventh Framework Programme for research, technological development and demonstration under grant agreement No 600841.

Appendix A Measurement Times

The times at which measurements were taken in the experiments described in Section 2.2 are given in Table 6.

Time Point	Time	Time Point	Time
t_1	65 h (2.7 d)	t_8	328 h (13.7 d)
t_2	92 h (3.8 d)	t_9	376 h (15.7 d)
t_3	112 h (4.7 d)	t_{10}	449 h (18.7 d)
t_4	159 h (6.6 d)	t_{11}	497 h (20.7 d)
t_5	207 h (8.6 d)	t_{12}	545 h (22.7 d)
t_6	257 h (10.7 d)	t_{13}	595 h (24.8 d)
t_7	303 h (12.6 d)	t_{14}	664 h (27.7 d)

Table 6: Times at which measurements of the spheroids were taken, measured after an initial seed of 2000 tumour cells per spheroid were implanted at $t = 0$.

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