The statistical analysis of non-linear longitudinal data in biomedical research using generalized additive models

Beyond repeated measures ANOVA and Linear Mixed Models

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33 1 Abstract

In biomedical research, the outcome of longitudinal studies has been traditionally analyzed using the repeated measures analysis of variance (rm-ANOVA) or more recently, a linear mixed model (LMEM). Although LMEMs are less restrictive that rm-ANOVA in terms of correlation and missing observations, both 36 methodologies share an assumption of linearity in the measured response, which results in biased estimates 37 and unreliable inference when they are used to analyze data where the trends are non-linear. In contrast, 38 generalized additive models (GAMs), are a class of models that relax the linearity assumption and allow 39 the data to determine the fit of the model while permitting missing observations and different correlation 40 structures, thereby being an excellent choice to analyze non-linear longitudinal data. This paper summa-41 rizes the limitations of LMEMs and rm-ANOVA, presents the basic theory of GAMs, and demonstrates their 42 implementation in R via the package mgcv using simulated data that follows longitudinal trends reported in 43 biomedical literature. To promote reproducibility in biomedical research, the code and data used to generate 44 this paper are available at:

46 2 Background

Longitudinal studies are designed to repeatedly measure a variable of interest in a group (or groups) of subjects, with the intention of observing the evolution of effect across time rather than analyzing a single 48 time point (e.g., a cross-sectional study). Biomedical research frequently uses longitudinal studies to analyze 49 the evolution of a "treatment" effect across multiple time points; and in such studies the subjects of analysis 50 range from animals (mice, rats, rabbits), to human patients, cells, or blood samples, among many others. 51 Tumor response [1-4], antibody expression [5,6], and cell metabolism [7,8] are examples of the different 52 situations where researchers have used longitudinal designs to study some physiological response. Because 53 the frequency of the measurements in a longitudinal study is dependent on the biological phenomena of 54 interest and the experimental design of the study, the frequency of such measurements can range from minute 55 intervals to study a short-term response such as anesthesia effects in animals[9], to weekly measurements 56 to analyze a mid-term response like the evolution of dermatitis symptoms in breast cancer patients [10], to monthly measurements to study a long-term response such as mouth opening following radiotherapy (RT) 58 in neck cancer patients [11]. 59

Traditionally, a "frequentist" or "classical" statistical paradigm is used in biomedical research to derive inferences from a longitudinal study. The frequentist paradigm regards probability as the limit of the expected outcome when an experiment is repeated a large number of times [12], and such view is applied to the analysis of longitudinal data by assuming a null hypothesis under a statistical model that is often an analysis of variance over repeated measures (repeated measures ANOVA or rm-ANOVA). The rm-ANOVA model makes three key assumptions regarding longitudinal data: 1) linearity of the response across time, 2) constant correlation across same-subject measurements, and 3) observations from each subject are obtained at all time points through the study (a condition also known as complete observations) [13,14].

The expected linear behavior of the response through time is a key requisite in rm-ANOVA [15]. This "linearity assumption" in rm-ANOVA implies that the model is misspecified when the data does not follow a linear trend, which results in unreliable inference. In biomedical research, non-linear trends are the norm

rather than the exception in longitudinal studies. A particular example of this non-linear behavior in longitudinal data arises in measurements of tumor response to chemo and/or radiotherapy in preclinical and clinical settings [1,8,16]. These studies have shown that the collected signal does not follow a linear trend over time, and presents extreme variability at different time points, making the fit of rm-ANOVA model inconsistent with the observed variation. Therefore, when rm-ANOVA is used to draw inference of such highly-variable data the estimates are inevitably biased, because the model is only able to accommodate linear trends that are far from adequately representing the biological phenomenon of interest.

A post hoc analysis is the statistical test used in conjunction with rm-ANOVA to perform repeated comparisons to estimate a p-value, which in turn is used as a measure of significance. Although it is possible that a post hoc analysis of rm-ANOVA is able to find "significant" p-values (p<0.05) from non-linear data, the validity of such metric is dependent on how adequate the model fits the data. In other words, p-values are valid only if the model and the data have good agreement; if that is not the case, a "Type III" error (known as "model misspecification") occurs[17]. For example, model misspecification will occur when a model that is only able to explain linear responses (such as rm-ANOVA) is fitted to data that follows a quadratic trend, thereby causing the resulting p-values and parameter estimates to be invalid [18].

Additionally, the *p-value* itself is highly variable, and multiple comparisons can inflate the false positivity rate (Type I error or α) [19,20], consequently biasing the conclusions of the study. Corrections exist to address the Type I error issue of multiple comparisons (such as Bonferroni [21]), but they in turn reduce statistical power $(1-\beta)[22]$, and lead to increased Type II error (failing to reject the null hypothesis when the null hypothesis is false) [23,24]. Therefore, the tradeoff of *post hoc* comparisons in rm-ANOVA between Type I, II and III errors might be difficult to resolve in a biomedical longitudinal study where a delicate balance exists between statistical power and sample size.

On the other hand, the assumption of constant correlation in rm-ANOVA (often known as the *compound* symmetry assumption) is typically unreasonable because correlation between the measured responses often diminishes as the time interval between the observation increases [25]. Corrections can be made in rm-ANOVA in the absence of compound symmetry [26,27], but the effectiveness of the correction is limited by the size of the sample, the number of measurements[28], and group sizes [29]. In the case of biomedical research, where living subjects are frequently used, sample sizes are often not "large" due to ethical and budgetary reasons [30] which might cause the corrections for lack of compound symmetry to be ineffective.

Due to a variety of causes, the number of observations during a study can vary between all subjects. For example, in a clinical trial patients may voluntarily withdraw, whereas attrition due to injury or weight loss in preclinical animal studies is possible. It is even plausible that unexpected complications with equipment or supplies arise that prevent the researcher from collecting measurements at certain time points. In each of these missing data scenarios, the *complete observations* assumption of classical rm-ANOVA is violated. When incomplete observations occur, a rm-ANOVA model is fit by excluding all subjects with missing observations from the analysis [13]. This elimination of partially missing data from the analysis can result in increased costs if the desired statistical power is not met with the remaining observations, because it would be necessary to enroll more subjects. At the same time, if the excluded observations contain insightful information that is not used, their elimination from the analysis may limit the demonstration of significant differences between groups.

During the last decade, the biomedical community has started to recognize the limitations of rm-ANOVA in the analysis of longitudinal information. The recognition on the shortcomings of rm-ANOVA is exemplified by the use of linear mixed effects models (LMEMs) by certain groups to analyze longitudinal tumor response data [8,16]. Briefly, LMEMs incorporate fixed effects, which correspond to the levels of experimental factors in the study (e.g., the different drug regimens in a clinical trial), and random effects, which account for random variation within the population (e.g., the individual-level differences not due to treatment such as weight or age). When compared to the traditional rm-ANOVA, LMEMs are more flexible as they can accommodate missing observations for multiple subjects and allow different modeling strategies for the variability within each measure in every subject [15]. However, LMEMs impose restrictions in the distribution of the errors of the random effects, which need to be normally distributed and independent [13,31]. And even more importantly, LMEMs also expect a linear relationship between the response and time [15], making them unsuitable to analyze non-linear data.

As the rm-ANOVA and the more flexible LMEM approaches make overly restrictive assumptions regarding the linearity of the response, there is a need for biomedical researchers to explore the use of additional statistical tools that allow the data (and not an assumption in trend) to determine the trend of the fitted model, to enable appropriate inference.

In this regard, generalized additive models (GAMs) present an alternative approach to analyze longitudinal 127 data. Although not frequently used by the biomedical community, these semi-parametric models are cus-128 tomarily used in other fields to analyze longitudinal data. Examples of the use of GAMs include the analysis of temporal variations in geochemical and palaeoecological data [32–34], health-environment interactions 130 [35] and the dynamics of government in political science [36]. There are several advantages of GAMs over 131 LMEMs and rm-ANOVA models: 1) GAMs can fit a more flexible class of smooth responses that enable 132 the data to dictate the trend in the fit of the model, 2) they can model non-constant correlation between 133 repeated measurements [37] and 3) can easily accommodate missing observations. Therefore, GAMs can 134 provide a more flexible statistical approach to analyze non-linear biomedical longitudinal data than LMEMs and rm-ANOVA. 136

The current advances in programming languages designed for statistical analysis (specifically R), have eased 137 the computational implementation of traditional models such as rm-ANOVA and more complex approaches such as LMEMs and GAMs. In particular, R[38] has an extensive collection of documentation and functions 139 to fit GAMs in the package mgcv [37,39] that not only speed up the initial stages of the analysis but also enable the use of advanced modeling structures (e.g. hierarchical models, confidence interval comparisons) 141 without requiring advanced programming skills from the user. At the same time, R has many tools that 142 simplify data simulation, an emerging strategy used to test statistical models [28]. Data simulation methods 143 allow the researcher to create and explore different alternatives for analysis without collecting information in the field, reducing the time window between experiment design and its implementation, and simulation 145 can be also used for power calculations and study design questions.

This work provides biomedical researchers with a clear understanding of the theory and the practice of using GAMs to analyze longitudinal data using by focusing on four areas. First, the limitations of LMEMs and rm-148 ANOVA regarding linearity of response, constant correlation structures and missing observations is explained in detail. Second, the key theoretical elements of GAMs are presented using clear and simple mathematical 150 notation while explaining the context and interpretation of the equations. Third, using simulated data that reproduces patterns in previously reported studies [16] we illustrate the type of non-linear longitudinal 152 data that often occurs in biomedical research. The simulated data experiments highlight the differences 153 in inference between rm-ANOVA, LMEMs and GAMs on data similar to what is commonly observed in 154 biomedical studies. Finally, reproducibility is emphasized by providing the code to generate the simulated 155 data and the implementation of different models in R, in conjunction with a step-by-step guide demonstrating 156 how to fit models of increasing complexity. 157

In summary, this work will allow biomedical researchers to identify when the use of GAMs instead of rmANOVA or LMEMs is appropriate to analyze longitudinal data, and provide guidance on the implementation
of these models by improving the standards for reproducibility in biomedical research.

3 Challenges presented by longitudinal studies

3.1 The repeated measures ANOVA

The repeated measures analysis of variance (rm-ANOVA) is the standard statistical analysis for longitudinal data in biomedical research. This statistical methodology requires certain assumptions for the model to be valid. From a practical view, the assumptions can be divided in three areas: 1) linear relationship between covariates and response, 2) a constant correlation between measurements, and, 3) complete observations for all subjects. Each one of these assumptions is discussed below.

3.2 Linear relationship

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3.2.1 The repeated measures ANOVA case

In a longitudinal biomedical study, two or more groups of subjects (e.g., human subject, mice, samples) are subject to different treatments (e.g., a "treatment" group receives a novel drug or intervention vs. a "control" group that receives a placebo), and measurements from each subject within each group are collected at specific time points. The collected response is modeled with *fixed* components. The *fixed* component can be understood as a constant value in the response which the researcher is interested in measuring, i.e., the average effect of the novel drug/intervention in the "treatment" group.

Mathematically speaking, a rm-ANOVA model with an interaction can be written as:

$$y_{ijt} = \beta_0 + \beta_1 \times time_t + \beta_2 \times treatment_j + \beta_3 \times time_t \times treatment_j + \varepsilon_{ijt}$$
 (1)

In this model y_{ijt} is the response for subject i, in treatment group j at time t, which can be decomposed in a mean value β_0 , fixed effects of time $(time_t)$, treatment $(treatment_j)$ and their interaction $time_t*treatment_j$ which have linear slopes given by β_1, β_2 and β_3 , respectively. Independent errors ε_{tij} represent random variation not explained by the fixed effects, and are assumed to be $\sim N(0, \sigma^2)$ (independently normally distributed with mean zero and variance σ_{μ}^2). In a biomedical research context, suppose two treatments groups are used in a study (e.g., "placebo" vs. "novel drug" or "saline" vs. "chemotherapy"). Then, the group terms in Equation (1) can be written as below with $treatment_j = 0$ representing the first treatment group (Group A) and $treatment_j = 1$ representing the second treatment group (Group B). The linear models then can be expressed as

$$y_{ijt} = \begin{cases} \beta_0 + \beta_1 \times time_t + \varepsilon_{ijt} & \text{if Group A} \\ \beta_0 + \beta_2 + \beta_1 \times time_t + \beta_3 \times time_t + \varepsilon_{ijt} & \text{if Group B} \end{cases}$$
 (2)

To further simplify the expression, substitute $\widetilde{\beta_0} = \beta_0 + \beta_2$ and $\widetilde{\beta_1} = \beta_1 + \beta_3$ in the equation for Group B.

This substitution allows for a different intercept and slope for Groups A and B. The model is then written as

$$y_{ijt} = \begin{cases} \beta_0 + \beta_1 \times time_t + \varepsilon_{ijt} & \text{if Group A} \\ \widetilde{\beta_0} + \widetilde{\beta_1} \times time_t + \varepsilon_{ijt} & \text{if Group B} \end{cases}$$
 (3)

Presenting the model in this manner makes clear that when treating different groups, an rm-ANOVA model is able to accommodate non-parallel lines in each case (different intercepts and slopes per group). In other words, the rm-ANOVA model "expects" a linear relationship between the covariates and the response, this means that either presented as Equation (1), Equation (2) or Equation (3), an rm-ANOVA model is only able to accommodate linear patterns in the data. If the data show non-linear behavior, the rm-ANOVA model will approximate this behavior with non-parallel lines.

3.2.2 The Linear Mixed Model Case

A linear mixed model (LMEM) is a class of statistical model that incorporates fixed effects to model the relationship between the covariates and the response, and random effects to model subject variability that is not the primary focus of the study but that might be important to distinguish [15,40]. A LMEM with interaction between time and treatment for a longitudinal study can be written as:

$$y_{ijt} = \beta_0 + \beta_1 \times time_t + \beta_2 \times treatment_j + \beta_3 \times time_t \times treatment_j + \mu_{ij} + \varepsilon_{ijt}$$
(4)

When Equation (1) and Equation (4) are compared, it is easily noticeable that LMEM and rm-ANOVA have the same construction regarding the fixed effects of time and treatment, but that the LMEM incorporates an 201 additional source of variation (the term μ_{ij}). This term μ_{ij} is the one that corresponds to the random effect, 202 accounting for variability in each subject within each group. The random component can also be understood as used to model some "noise" in the response, but that is intended to be analyzed and disentangled from 204 the "global noise" term ε_{ijt} from Equation (1). 205

For example, if the blood concentration of the drug is measured in certain subjects in the early hours of 206 the morning while other subjects are measured in the afternoon, it is possible that the difference in the 207 collection time introduces some "noise" in the data. As the name suggests, this "random" variability needs 208 to be modeled as a variable rather than as a constant value. The random effect μ_{ij} in Equation (4) is 209 assumed to be $\mu_{ij} \sim N(0, \sigma_{\mu}^2)$. In essence, the random effect in a LMEM enables to fit models with different slopes at the subject-level[15]. However, the expected linear relationship of the covariates and the response 211 in Equation (1) and in Equation (4) is essentially the same, representing a major limitation of LMEMs to 212 fit a non-linear response. 213

3.3 Covariance in rm-ANOVA and LMEMs

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In a longitudinal study there is an expected *covariance* between repeated measurements on the same subject, 215 and because repeated measures occur in the subjects within each group, there is a covariance between measurements at each time point within each group. The covariance matrix (also known as the variancecovariance matrix) is a matrix that captures the variation between and within subjects in a longitudinal study[41] (For an in-depth analysis of the covariance matrix see [40,42]).

In the case of an rm-ANOVA analysis, it is typically assumed that the covariance matrix has a specific 220 construction known as compound symmetry (also known as "sphericity" or "circularity"). Under this assumption, the between-subject variance and within-subject correlation are constant across time [26.42,43]. However, it has been shown that this condition is frequently not justified because the correlation between measurements tends to change over time [44]; and it is higher between consecutive measurements [13,25]. Although corrections can be made (such as Huyhn-Feldt or Greenhouse-Geisser)[26,27] the effectiveness of each correction is limited because it depends on the size of the sample, the number of repeated measurements[28], and they are not robust if the group sizes are unbalanced [29]. Because biomedical longitudinal studies are often limited in sample size and can have an imbalanced design, the corrections required to use an rm-ANOVA model may not be able to provide a reasonable adjustment that makes the model valid.

In the case of LMEMs, one key advantage over rm-ANOVA is that they allow different structures for the variance-covariance matrix including exponential, autoregressive of order 1, rational quadratic and others 231 [15]. Nevertheless, the analysis required to determine an appropriate variance-covariance structure for the data can be a challenging process by itself. Overall, the spherical assumption for rm-ANOVA may not 233 capture the natural variations of the correlation in the data, and can bias the inferences from the analysis.

3.4 Missing observations

Missing observations are an issue that arises frequently in longitudinal studies. In biomedical research, 236 this situation can be caused by reasons beyond the control of the investigator [45]. Dropout from patients 237 and attrition or injury in animals are among the reasons for missing observations. Statistically, missing 238 information can be classified as missing at random (MAR), missing completely at random (MCAR), and missing not at random (MNAR) [42]. In a MAR scenario, the pattern of the missing information is related 240 to some variable in the data, but it is not related to the variable of interest [46]. If the data are MCAR, this means that the missingness is completely unrelated to the collected information [47], and in the case of MNAR 242 the missing values are dependent on their value. An rm-ANOVA model assumes complete observations for all subjects, and therefore subjects with one or more missing observations are excluded from the analysis. This 244 is inconvenient because the remaining subjects might not accurately represent the population, and statistical power is affected by this reduction in sample size [48].

In the case of LMEMs, inferences from the model are valid when missing observations in the data exist that are MAR or MCAR [40]. For example, if attrition occurs in all mice that had lower weights at the beginning of a chemotherapy response study, the missing data can be considered MAR because the missigness is unrelated to other variables of interest.

This section has presented the assumptions for analyzing longitudinal data using rm-ANOVA and LMEMs and compared their differences regarding linearity, the covariance matrix and missing data. In particular, LMEMs offer a more robust and flexible approach than rm-ANOVA and if the data follows a linear trend, they provide an excellent choice to derive inferences from a repeated measures study. However, when the data presents high a non-linear behavior, LMEMs and rm-ANOVA fail to capture the trend of the data. To better convey the issues of linearity and correlation in linear models fitted to non-linear data, simulation is used in the next section.

3.5 What does an rm-ANOVA fit looks like? A visual representation using simulated data

To demonstrate the limitations of rm-ANOVA an LMEMs for non-linear longitudinal data, this section presents a simulation experiment of a normally distributed response of two groups of 10 subjects each. An rm-ANOVA model (Equation (1)) is fitted to each group, using R[38] and the package nlme[49]. Briefly, two cases for the mean responses for each group are considered: in the first case, the mean response in each group is a linear function with different intercepts and slopes; a negative slope is used for Group 1 and a positive slope is used for Group 2 (Figure 1, A). In the second case, a second-degree polynomial (quadratic) function is used for the mean response per group: the quadratic function is concave down for Group 1 and it is concave up for Group 2 (Figure 1, C). In both the linear and quadratic simulated data, the groups start with the same mean value at the first time point. This is intentional in order to simulate the expected temporal evolution of some physiological quantity.

Specifically, the rationale for the chosen linear and quadratic functions is the likelihood that a measured response in two treatment groups is similar in the initial phase of the study, but as treatment progresses a divergence in the trend of the response indicates an effect due to treatment. In other words, Group 1 can be thought as a "Control" group and Group 2 as a "Treatment" group. From the mean response per group (linear or quadratic), the variability or "error" of individual responses within each group is simulated using a covariance matrix with compound symmetry (constant variance across time). Thus, the response per subject in both the linear and quadratic simulation corresponds to the mean response per group plus the error (Figure 1 B,D). A more comprehensive exploration of the fit of rm-ANOVA for linear and non-linear longitudinal appears in Figure 5 and Figure 6 in the Appendix, where simulation with compound symmetry and independent errors (errors generated from a normal distribution that are not constant over time) and the plot of simulated errors, and fitted parameters in presented.

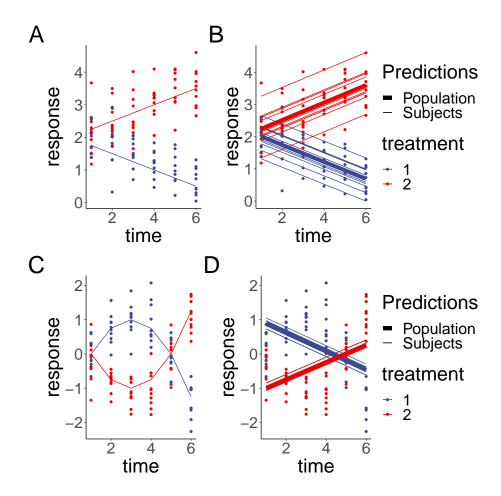


Figure 1: Simulated linear responses from two groups with correlated (top row) or independent (bottom row) errors using a rm-ANOVA model. A, C:Simulated data with known mean response (linear or quadratic, thin lines) and individual responses (points) showing the dispersion of the data. B,D: Estimates from the rm-ANOVA model for the mean group response (linear of quadratic). Thick lines are the predicted mean response per group, thin lines are the random effects for each subject and points represent the original raw data. The rm-ANOVA model not only fails to pick the trend of the quadratic data but it also incorrectly estimates the initial conditions.

The simulation shows that the fit produced by the rm-ANOVA model is good for linear data, as the predictions for the mean response are reasonably close to the "truth" of the simulated data (Figure 1,B). When the linearity and compound symmetry assumptions are met, the model approximates well the individual trends and the mean trends by group.

However, consider the case when the data follows a non-linear trend, such as the simulated data in Figure 1, C. Here, the mean response per group was simulated using a quadratic function but errors, individual responses and the rm-ANOVA model were produced in the same manner as in (Figure 1 A and B). The mean response in the simulated data with quadratic behavior is changing in each group through the timeline, and the mean value is the same as the initial value by the fifth time point for each group. Fitting an rm-ANOVA model (1) to this data produces the fit that appears in panel D in Figure 1.

A comparison of the fitted mean response of the rm-ANOVA model to the the simulated data in Figure ((1, D) indicates that the model is not capturing the changes within each group in a good way. Specifically, note that the fitted mean response of the rm-ANOVA model (panel D) shows that the change (increase for Treatment 1 or decrease for Treatment 2) in the response through time points 2 and 4 is not being captured by the model. Moreover, the rm-ANOVA model is not being able to capture the fact that the initial values are the same in each group, and instead fits non-parallel lines that have initial values that are

markedly different from the "true" initial values in each case (compare panels C and D). If such change has important physiological implications, the rm-ANOVA model omits it from the fitted mean response. Thus, even though the model correctly detects a difference in treatment groups, the exact nature of this difference is not correctly identified, limiting valuable inferences from the data.

This section has used simulation to better convey the limitations of linearity and correlation in the response in non-linear data. Although the model fitted to the simulated data was an rm-ANOVA model, the main issue of an expected linear trend in the response is the same in the case of a LMEM. In the following section, we present generalized additive models (GAMs) as a data-driven alternative method to analyze longitudinal non-linear data.

306 4 GAMs as a special case of Generalized Linear Models

4.1 GAMs and Basis Functions

Generalized linear models (GLMs) are a family of models that fit a linear response function to data that do not have normally distributed errors[50]. In contrast, GAMs are a family of regression-based methods for estimating smoothly varying trends and are a broader class of models that contain the GLM family as a special case[34,37,51]. A GAM model can be written as:

$$y_{ijt} = \beta_0 + f(x_t \mid \beta_j) + \varepsilon_{ijt} \tag{5}$$

Where y_{ijt} is the response at time t of subject i in group j, β_0 is the expected value at time 0, the change of y_{ijt} over time is represented by the function $f(x_t \mid \beta_j)$ with inputs as the covariates x_t and parameters β_j , and ε_{ijt} represents the residual error.

In contrast to the linear functions used to model the relationship between the covariates and the response in rm-ANOVA or LMEM, GAMs use more flexible *smooth functions*. This approach is advantageous as it does not restrict the model to a linear relationship, although a GAM will estimate a linear relationship if the data is consistent with a linear response. One possible set of functions for $f(x_t \mid \beta_j)$ that allow for non-linear responses are polynomials, but a major limitation is that polynomials create a "global" fit as they assume that the same relationship exists everywhere, which can cause problems with the fit [36]. In particular, polynomial fits are known to show boundary effects because as t goes to $\pm \infty$, $f(x_t \mid \beta_j)$ goes to $\pm \infty$ which is almost always unrealistic, and causes bias at the endpoints of the time period.

The smooth functional relationship between the covariates and the response in GAMs is specified using a 323 semi-parametric relationship that can be fit within the GLM framework, by using basis functions expansions of the covariates and by estimating random coefficients for these basis functions. A basis is a set of functions 325 that spans the space where the smooths that approximate $f(x_t \mid \beta_i)$ exist [34]. For the linear model in 326 Equation (1), the basis coefficients are β_1 , β_2 and β_3 and the basis vectors are $time_t$, $treatment_j$ and 327 $time_t \times treatment_i$. The basis function then, is the combination of basis coefficients and basis vectors that 328 map the possible relationship between the covariates and the response [52], which in the case of Equation 329 (1) is restricted to a linear family of functions. In the case of Equation (5), the basis function is $f(x_t | \beta_i)$, 330 which means that the model allows for non-linear relationships among the covariates. 331

A commonly used *basis function* is the cubic spline, which is a smooth curve constructed from cubic polynomials joined together in a manner that enforces smoothness [34,37]. Cubic splines have a long history in solving semi-parametric statistical problems and are often a default choice to fit GAMs as they are a simple, flexible and powerful option to obtain smoothness [53]. Therefore, this data-driven flexibility in GAMs overcomes the limitation that occurs in LMEMs and rm-ANOVA when the data is non linear.

To further clarify the concept of basis functions and smooth functions, consider the simulated response for Group 1 in Figure (1, C). The simplest GAM model that can be used to estimate such response is that of a single smooth term for the time effect; i.e., a model that fits a smooth to the trend of the group through time.

The timeline can be divided in equally spaced knots, each knot being a region where a different basis function will be used. Because there are six timepoints for this group, five knots can be used. The model with five knots to construct the smooth term means that it will have four basis functions (plus one that corresponds to the intercept). The choice of basis functions is already optimized in the package mgcv depending on the number of knots. In Panel A of Figure 2, the four basis functions (and the intercept) are shown. Each of the basis functions is composed of six different points (because there are six points on the timeline). To control the "wigliness" of the fit, each of the basis functions of Panel A is penalized by multiplying it by a coefficient according to the penalty matrix of Panel B. The penalty reduces the "wigliness" of the smooth fit to prevent overfitting: A weak penalty estimate will result in wiggly functions whereas a strong penalty 348 estimate provides evidence that a linear response is appropriate.

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In other words, the six points of each basis are multiplied by the corresponding coefficient in panel B, thereby increasing or decreasing the original basis functions of Panel A. In Figure 2, Panel C shows the resulting penalized basis functions. Note that the penalization for basis 1 has resulted in a decrease of its overall value (because the coefficient for that basis function is negative and less than 1); on the other hand, basis 3 has roughly doubled its value. Finally, the penalized basis functions are added at each timepoint to produce the smooth term. The resulting smooth term for the effect of time is shown in Panel D (orange line) along the simulated values per group, which appear as points.

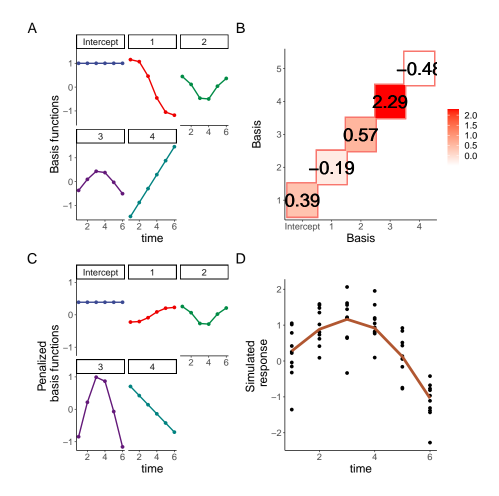


Figure 2: Basis functions for a single smoother for time with five knots. A: Basis functions for a single smoother for time for the simulated data of Group 1 from Figure 2, the intercept basis is not shown. B: Penalty matrix for the basis functions. Each basis function is penalized by a coefficient which can be positive or negative. The coefficient determines the overall effect of each basis in the final smoother. C: Penalized basis functions. Each of the four basis functions of panel A has been penalized by the corresponding coefficient shown in Panel B, note the corresponding increase (or decrease) of each basis. D: Smoother for time and original data points. The smoother (line) is the result of the sum of each penalized basis function at each time point, simulated values for the group appear as points.

5 The analysis of longitudinal biomedical data using GAMs

The previous sections provided the basic theoretical framework to understand the GAM framework and how these models are more advantageous to analyze non-linear longitudinal data when compared to LMEMs or rm-ANOVA. This section will use simulation to present the practical implementation of GAMs for longitudinal biomedical data using R and the package mgcv. The code for the simulated data and figures, and a brief guide for model selection and diagnostics appear in the Appendix.

53 5.1 Simulated data

The simulated data is based on the reported longitudinal changes in oxygen saturation (StO₂) in subcutaneous tumors that appear in Figure 3(C) in [16]. In the paper, diffuse reflectance spectroscopy was used to
quantify StO₂ changes in both groups at the same time points (days 0, 2, 5, 7 and 10). In the "Treatment"
group (chemotherapy) an increase in StO₂ is observed through time, while a decrease is seen in the "Control"
(saline) group. Following the reported trend, we simulated 10 normally distributed observations at each time
point with a standard deviation (SD) of 10% (matching the SD in the original paper). The simulated and
real data appear in Figure 3, A and the inlet, respectively.

5.2 An interaction GAM for longitudinal data

An interaction model is typically the main interest in longitudinal biomedical data, as it takes into account the effect of treatment, time, and their combination. In a practical sense, when a GAM is implemented for longitudinal data, a smooth can be added to the model for the *time* effect to account for the repeated measures over time. Although specific methods of how GAMs model correlation structures is a topic beyond the scope of this paper, it suffices to say that GAMs are flexible and can handle correlation structures beyond compound symmetry. A detailed description on basis functions and correlations can be found in [52].

For the data in Figure 3,A it is of interest to know the effect that each treatment has in StO_2 . The model then needs to incorporate independent smooths for *Group* and *Day*, respectively. The main thing to consider is that model syntax accounts for the fact that one of the variables is numeric (Day) and the other is a factor (Group). Because the smooths are centered at 0, the factor variable needs to be specified as a parametric term in order to identify any differences between the groups. Using R and the package mgcv the model syntax is:

m1<-gam(StO2 sim~Group+s(Day,by=Group,k=5,bs="gp"), method='REML',data=dat sim

This syntax specifies that m1 will store the model, and that the change in the simulated oxygen saturation (StO2_sim) is modeled using independent smooths for Group and Day (the parenthesis preceded by s) using 386 5 knots. The smooth is constructed using gaussian process smooths, which is indicated by bs="gp". These splines are used to model temporal trends and might be particularly suited for long-term studies where the 388 correlation between measurements changes as a function of the time intervals [34]. The parametric term 389 Group is added to quantify differences in the effect of treatment between groups, and the method chosen to 390 select the smoothing parameters is the restricted maximum likelihood (REML) [37]. When the smooths are 391 plotted over the raw data, it is clear that the model has been able to capture the trend of the change of StO₂ 392 for each group across time (Figure 3,B). Model diagnostics can be obtained using the gam.check function, 393 and the function appraise from the package gratia [54]. A guide for model selection and diagnostics is in the 394 Appendix, and an in-depth analysis can be found in [37] and [55]. 395

One question that might arise at this point is "what is the fit that an rm-ANOVA model produces for the simulated data?". The rm-ANOVA model, which corresponds to Equation (1) is presented in Figure 3,C.
This is a typical case of model misspecification: The slopes of each group are different, which would lead to a *p-value* indicating significance for the treatment and time effects, but the model is not capturing the changes that occur at days 2 and between days 5 and 7, whereas the GAM model is able to do so (Figure 3,B).

Because GAMs do not require equally-spaced or complete observations for all subjects, they are advantageous to analyze longitudinal data where missingness exists. The rationale behind this is that GAMs are able to pick the trend in the data even when some observations are missing. However, this usually causes the resulting smooths to have wider confidence intervals and less ability to pick certain trends. Consider the simulated StO₂ values from Figure (3, B). If 40% of the total observations are randomly deleted and the same interaction GAM fitted for the complete dataset is used, the resulting smooths are still able to show a different trend for each group, but the "Treatment" smooth takes an overall more linear profile (3, D). Although the confidence intervals have increased for both smooths, the model still shows different trends with as little as 4 observations per group at certain time points.

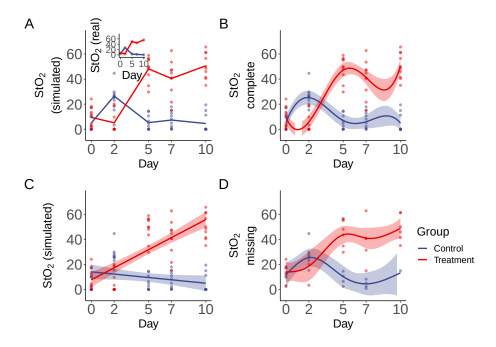


Figure 3: Simulated data and smooths for oxygen saturation in tumors. A: Simulated data that follows previously reported trends (inset) in tumors under chemotherapy (Treatment) or saline (Control) treatment. Simulated data is from a normal distribution with standard deviation of 10% with 10 observations per time point. Lines indicate mean oxygen saturation B: Smooths from the GAM model for the full simulated data with interaction of Group and Treatment. Lines represent trends for each group, shaded regions are 95% confidence intervals. C: rm-ANOVA model for the simulated data, the model does not capture the changes in each group over time. D: Smooths for the GAM model for the simulated data with 40% of its observations missing. Lines represent trends for each group, shaded regions are 95% confidence intervals.

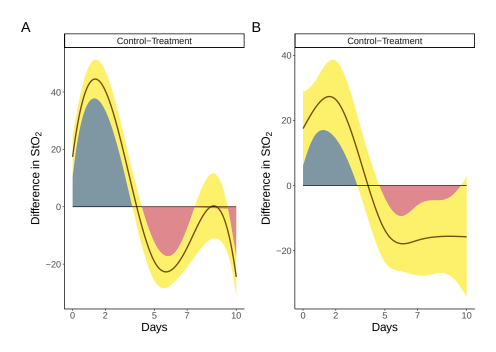


Figure 4: Pairwise comparisons for smooth terms. A: Pairwise comparisons for the full dataset. B: Pairwise comparisons for the dataset with missing observations. Significant differences exist where the interval does not cover 0. In both cases the effect of treatment is significant after day 5.

5.3 Determination of significance in GAMs for longitudinal data

At the core of a biomedical longitudinal study lies the question of a significant difference between the effect of two or more treatments in different groups. Whereas in rm-ANOVA a post-hoc analysis is required to answer such question by calculating some p-values after multiple comparisons, GAMs can use a different approach to estimate significance. In essence, the idea behind the estimation of significance in GAMs across different treatment groups is that if the difference between the confidence intervals of the fitted smooths for such groups is non-zero, then a significant difference exists at that time point(s). The absence of a p-value in this case might seem odd, but the confidence interval comparison can be conceptualized in the following manner: Different trends in each group are an indication of an effect by the treatment. This is what happens for the simulated data in Figure 3, A where the chemotherapy causes StO₂ to increase over time.

With this expectation of different trends in each group, computing the difference between the trends will identify if the observed change is significant. The difference between groups with similar trends is likely to yield zero, which would indicate that the treatment is not causing a change in the response in one of the groups (assuming the other group is a Control or Reference group).

Consider the calculation of pairwise differences for the smooths in 3 B and D. Figure 4, shows the comparison between each treatment group. Here, the effect of the "Control" group is compared to that of the "Treatment" group. The shaded region under the curve highlights the interval where each group has a higher effect than the other. Notice that the shaded region between days 0 and 4 indicates that through that time, the "Control" group has higher StO₂, but as therapy progresses the effect is reversed and by day 5 it is the "Treatment" group the one that has greater contribution. This would suggest that the effect of chemotherapy is significant after day 5 for the model used; which is noticeable in the data with missing observations albeit the missing data model has a wider confidence interval and is not able to pick the change between days 7 and 10 that the full dataset model is able to detect.

6 Conclusion

The traditional methods to analyze biomedical longitudinal data (rm-ANOVA, LMEMs) are restrictive when the data does not follow a linear pattern and their use in such situations leads to unreliable inference and biased estimates. GAMs present a suitable alternative to analyze non-linear longitudinal biomedical data, as they overcome the limitations of linearity, compound symmetry, and missing observations imposed by traditional methods. By presenting the implementation of GAMs using simulated data that follows previously reported trends in the literature, we aim at encouraging biomedical researchers to consider different statistical tools to analyze longitudinal data. Finally, by providing the data and code used in this paper we hope to address the need of creating and sharing reproducible work in biomedical research.

7 References

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A Code for Manuscript data

599

601

602

604

625

This section presents the code used to generate figures, models and simulated data from Sections 3 and 4 from the main manuscript.

A.1 Compound symmetry and independent errors in linear and quadratic responses

This section simulated linear and quadratic data in the same manner as in Section 3.5. The linear simulations using Figure 5 show in panels A and D the simulated mean responses and individual data points. Panels C and G show a visual interpretation of "correlation" in the responses: In panel C, subjects that have a value of the random error ε either above or below the mean group response are more likely to have other observations that follow the same trajectory, thereby demonstrating correlation in the response. In panel G, because the errors are independent, there is no expectation that responses are likely to follow a similar pattern. Panels D and H show the predictions from the rm-ANOVA model.

The following code produces a more comprehensive exploration of Figure 1 in the main manuscript.

```
606
     607
  #
608
     609
610
611
  ## Example with linear response
612
613
  #This function simulates data using a linear or quadratic mean response and
614
     each with correlated
615
  \#or\ uncorrelated\ errors. Each group has a different slope/concavity.
616
  example <- function(n_time = 6, #number of time points
617
                    fun_type = "linear", #type of response
                    error_type = "correlated") {
619
    if (!(fun_type %in% c("linear", "quadratic")))
621
      stop ('fun_type | must | be | either | "linear ", | or | "quadratic "')
    if \ (!(\texttt{error\_type \%in\% c("correlated", "independent"))})\\
623
      stop('fun_type_must_be_either_"correlated", _or_ "independent"')
```

```
x \leftarrow seq(1,6, length.out = n_time)
627
628
      #Create mean response matrix: linear or quadratic
629
      mu \leftarrow \mathbf{matrix}(0, \mathbf{length}(x), 2)
      # linear response
631
      if (fun type == "linear") {
632
        mu[, 1] \leftarrow -(0.25*x)+2
633
        mu[, 2] \leftarrow 0.25*x+2
      } else {
635
        \# quadratic response (non-linear)
636
637
        mu[, 1] \leftarrow -(0.25 * x^2) +1.5*x-1.25
638
        mu[, 2] \leftarrow (0.25 * x^2) -1.5*x+1.25
639
640
      #create an array where individual observations per each time point for each
642
          group are to be stored. Currently using 10 observations per timepoint
643
      y \leftarrow \operatorname{array}(0, \operatorname{dim} = \operatorname{c}(\operatorname{length}(x), 2, 10))
644
      #Create array to store the "errors" for each group at each timepoint. The "
646
          errors " are the
647
      #between-group variability in the response.
648
      errors \leftarrow array (0, dim = c(length(x), 2, 10))
      #create an array where 10 observations per each time point for each group
650
651
          are to be stored
652
      #The following cycles create independent or correlated responses. To each
653
          value of mu (mean response per group) a randomly generated error (
654
          correlated or uncorrelated) is added and thus the individual response is
655
           created.
656
      if (error_type == "independent") {
657
        ## independent errors
658
        for (i in 1:2) {
659
           for (j in 1:10) {
             errors [, i, j] \leftarrow \mathbf{rnorm}(6, 0, 0.25)
661
             y[, i, j] \leftarrow mu[, i] + errors[, i, j]
662
           }
663
      } else {
665
                                # number of treatments
        for (i in 1:2) {
           for (j in 1:10) { # number of subjects
667
             # compound symmetry errors: variance covariance matrix
             errors [, i, j] \leftarrow \text{rmvn}(1, \text{rep}(0, \text{length}(x)), 0.1 * \text{diag}(6) + 0.25 *
669
                 \mathbf{matrix}(1, 6, 6))
670
             y[, i, j] \leftarrow mu[, i] + errors[, i, j]
671
672
673
      }
674
675
676
      \#\#\ subject\ random\ effects
677
678
      ## visualizing the difference between independent errors and compound
679
          symmetry
680
```

```
## why do we need to account for this - overly confident inference
681
682
   #labelling y and errors
683
     dimnames(y) \leftarrow list(time = x,
                            treatment = 1:2,
685
                            subject = 1:10
687
     dimnames (errors) <- list (time = x,
688
                                  treatment = 1:2,
689
                                  subject = 1:10
690
691
     #labeling the mean response
692
     dimnames(mu) \leftarrow list(time = x)
693
                             treatment = 1:2)
694
     \#convert y, mu and errors to dataframes with time, treatment and subject
696
         columns
697
     dat <- as.data.frame.table(y,
698
                                    responseName = "y")
     dat_errors <- as.data.frame.table(errors,
700
                                            responseName = "errors")
701
     dat_mu <- as.data.frame.table(mu,
702
                                        responseName = "mu")
704
     #join the dataframes to show mean response and errors per subject
705
     dat <- left_join(dat, dat_errors,
706
                         \mathbf{by} = \mathbf{c}("time", "treatment", "subject"))
707
     dat <- left_join(dat, dat_mu,
708
                        \mathbf{by} = \mathbf{c}("time", "treatment"))
709
     \#add time
710
     dat$time <- as.numeric(as.character(dat$time))
711
     \#label subjects per group
712
     dat <- dat %>%
713
        mutate(subject = factor(paste(subject,
                                          treatment.
715
                                          sep = "-"))
716
717
718
     ## repeated measures ANOVA in R
719
   #time and treatment interaction model, compound symmetry required by the model
      fit_lme <- lme(y ~ treatment + time + treatment:time,
721
                      data = dat
722
                      random = \sim 1 \mid subject,
723
                       correlation = corCompSymm(form = ~ 1 | subject)
724
     )
725
726
     #create a prediction frame where the model can be used for plotting purposes
727
     pred_dat <- expand.grid(
728
        treatment = factor(1:2),
        time = unique(dat\$time)
730
     )
731
732
     #add model predictions to the dataframe that has the simulated data
733
     dat$y_pred <- predict(fit_lme)
734
```

```
735
     #return everything in a list
736
     return(list(
737
       dat = dat,
       pred_dat = pred_dat,
739
       fit lme = fit lme
741
     ))
743
   744
   745
   #This function will create the plots for either a "linear" or "quadratic"
746
      response
747
748
   plot_example <- function(sim_dat) {</pre>
     ## Plot the simulated data (scatterplot)
750
     p1 <- sim_dat$dat %%
751
       ggplot(aes(x = time,
752
753
                  y = y,
                  group = treatment,
754
                  color = treatment)
755
              ) +
756
       geom_point(show.legend=FALSE) +
       labs (y='response')+
758
       geom_line(aes(x = time,
759
                     y = mu,
760
                     color = treatment),
761
                 show.legend=FALSE) +
762
       theme_classic() +
763
       theme(plot.title = element_text(size = 30,
764
                                      face = "bold"),
765
           text = element\_text(size = 30)+
766
       scale_color_aaas()
767
     #plot the simulated data with trajectories per each subject
769
     p2 <- sim_dat$dat %%
770
       ggplot(aes(x = time,
771
                  y = y,
772
                  group = subject,
773
                  color = treatment)
774
              ) +
775
       geom_line(aes(size = "Subjects"),
                 show.legend = FALSE) +
777
       \# facet\_wrap(\sim treatment) +
778
       geom_line(aes(x = time,
779
                     y = mu,
780
                     color = treatment,
781
                     size = "Simulated_Truth"),
782
                 lty = 1, show. legend = FALSE) +
       labs (y='response')+
784
       scale\_size\_manual(name = "Type", values=c("Subjects" = 0.5, "Simulated_
785
          Truth'' = 3) +
786
       theme classic()+
787
        theme(plot.title = element text(size = 30,
788
```

```
face = "bold"),
789
         text=element text(size=30))+
790
        scale color aaas()
791
792
     #plot the errors
793
      p3 \leftarrow sim dat dat \%
794
        ggplot(aes(x = time,
795
                    y = errors
                    group = subject,
797
                    color = treatment)) +
798
        geom_line(show.legend=FALSE) +
799
         labs(y='errors')+
800
         theme_classic()+
801
         theme(plot.title = element_text(size = 30,
802
                                          face = "bold"),
            text = element\_text(size = 30)+
804
        scale_color_aaas()
805
806
      #plot the model predictions
807
     p4 <- ggplot (sim_dat$dat,
808
                    aes(x = time,
809
                         y = y,
810
                         color = treatment)) +
        geom_point()+
812
        labs (y='response')+
813
        geom_line(aes(y = predict(sim_dat\fit_lme),
814
                       group = subject , size = "Subjects")) +
815
        geom_line(data = sim_dat$pred_dat,
816
                   aes (y = predict (sim_dat $ fit_lme,
817
                                     level = 0,
818
                                     newdata = sim_dat$pred_dat),
819
                        size = "Population")) +
820
        scale_size_manual(name = "Predictions",
821
                            values=c("Subjects" = 0.5, "Population" = 3)) +
        theme classic() +
823
        theme(plot.title = element_text(size = 30,
824
                                          face = "bold"),
825
            text = element\_text(size = 30)+
        scale color aaas()
827
     return((p1+p3+p2+p4)+plot layout(nrow=1)+plot annotation(tag levels = 'A'))
829
830
831
   }
832
833
   t \times t < -18
834
835
   #Store each plot in a separate object
836
   Al = linear , error_type = "linear", error_type = "correlated"))
837
838
   Bl<-plot_example(example(fun_type = "linear", error_type = "independent"))
839
840
   Cl<-plot_example(example(fun_type = "quadratic", error_type = "correlated"))
841
842
```

B43 DK—plot_example(example(fun_type = "quadratic", error_type = "independent"))

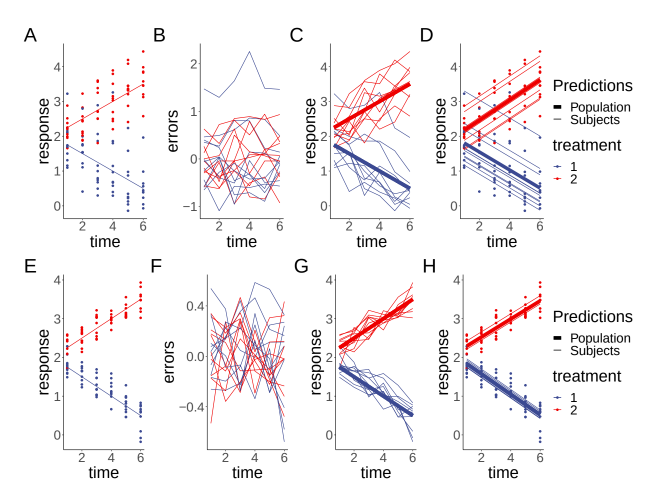


Figure 5: Simulated linear responses from two groups with correlated (top row) or independent (bottom row) errors using a rm-ANOVA model. A, C:Simulated data with known mean response (linear or quadratic, thin lines) and individual responses (points) showing the dispersion of the data. B,D: Estimations from the rm-ANOVA model for the mean group response (linear of quadratic). Thick lines are the predicted mean response per group, thin lines are the random effects for each subject and points represent the original raw data. The rm-ANOVA model does not pick the trend of the quadratic data.

For the quadratic response case, Figure 6 shows the simulated responses using compound symmetry and independent errors.

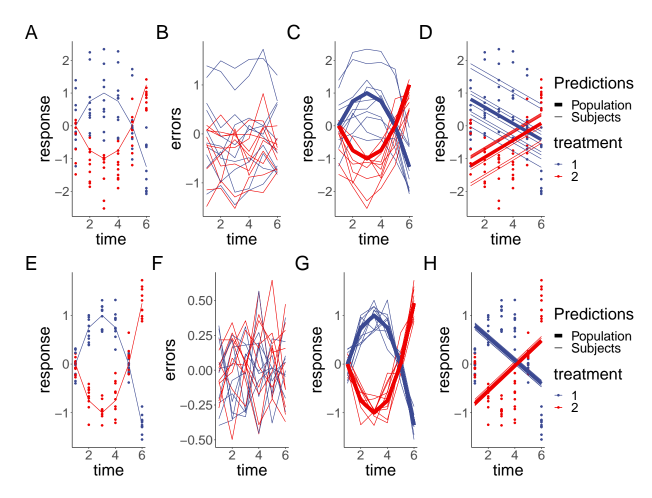


Figure 6: Simulated quadratic responses from two groups with a rm-ANOVA model fitted. A,E:Simulated data with known mean response (lines) and individual responses (points) showing the dispersion of the data. B,F: Generated errors showing the difference in the behavior of correlated and independent errors. C,G: Simulated known response per group (thick lines) with individual trajectories (thin lines), note that subjects with observations in the area above the mean response tend to stay in that region through the timeline. D,H: Estimations from the rm-ANOVA model for the mean group response. Thick lines are the predicted mean response per group, thin lines are the random effects for each subject and points represent the original raw data.

$_{^{16}}$ A.2 Basis functions and GAMs

847

848

849

850

This code produces Figure 2 from the main manuscript. Briefly, a non-linear (quadratic) response is simulated a gam model is fitted and the basis are extracted in order to explain how the smooth is constructed. The code for data simulation is used again here for the sake of keeping the same structure, although the data can be simulated in a more simple fashion.

```
\#generate the response: the same initial procedure from the previous section
851
         to simulate
852
    #the response
    n \text{ time} = 6
854
     x \leftarrow seq(1.6, length.out = n_time)
855
     mu \leftarrow \mathbf{matrix}(0, \mathbf{length}(x), 2)
856
     mu[, 1] \leftarrow -(0.25 * x^2) +1.5*x-1.25 \#mean response
857
             2] <- (0.25 * x^2) -1.5*x+1.25 \#mean response
858
     y \leftarrow \operatorname{array}(0, \operatorname{dim} = \operatorname{c}(\operatorname{length}(x), 2, 10))
859
```

```
errors \leftarrow array (0, \dim = \mathbf{c}(length(x), 2, 10))
860
                               # number of treatments
     for (i in 1:2) {
861
          for (j in 1:10) { # number of subjects
862
               # compound symmetry errors
               errors [, i, j] \leftarrow rmvn(1, rep(0, length(x)), 0.1 * diag(6) + 0.25 *
864
                   \mathbf{matrix}(1, 6, 6))
               y[, i, j] \leftarrow mu[, i] + errors[, i, j]
866
          }
     }
868
869
     #label each table
870
      dimnames(y) \leftarrow list(time = x, treatment = 1:2, subject = 1:10)
871
     dimnames(errors) \leftarrow list(time = x, treatment = 1:2, subject = 1:10)
872
873
     dimnames(mu) \leftarrow list(time = x, treatment = 1:2)
874
     #Convert to dataframes with subject, time and group columns
875
     dat <- as.data.frame.table(y, responseName = "y")
876
     dat errors <- as.data.frame.table(errors, responseName = "errors")
877
     dat_mu <- as.data.frame.table(mu, responseName = "mu")
878
     \mathrm{dat} \leftarrow \ \mathrm{left\_join} \left( \mathrm{dat} \,, \ \mathrm{dat\_errors} \,, \ \mathbf{by} = \mathbf{c} \left( \text{"time"} \,, \ \text{"treatment"} \,, \ \text{"subject"} \right) \right)
879
     dat \leftarrow left\_join(dat, dat\_mu, by = c("time", "treatment"))
880
     dat$time <- as.numeric(as.character(dat$time))
881
     #label subject per group
883
     dat <- dat %%
884
          mutate(subject = factor(paste(subject, treatment, sep = "-")))
885
     #extract "Group 1" to fit the GAM
887
      dat \leftarrow \mathbf{subset} (dat, treatment = = 1)
888
     #keep just the response and timepoint columns
889
       {\tt dat}{\leftarrow}{\tt dat}\left[\;, \mathbf{c}\left(\;'y\;'\;,\;'time\;'\right)\;\right]
890
891
       #GAM model of time, 5 knots
892
   gm \leftarrow gam(y \sim s(time, k=5), data = dat)
894
   #model_matrix (also known as) 'design matrix'
895
   #will contain the smooths used to create model 'qm'
896
   model_matrix<-as.data.frame(predict(gm, type='lpmatrix'))
898
   time < -c (1:6)
900
901
   basis — model_matrix[1:6,] #extracting basis (because the values are repeated
902
        after every 6 rows)
903
   \#basis \leftarrow model\_matrix[1:6,-1] \#extracting basis
904
   colnames (basis) [colnames (basis) == "(Intercept)"] <- "s (time).0"
905
   basis \square basis \%\% #pivoting to long format
906
      pivot_longer(
907
         cols=starts_with("s")
909
      arrange (name) #ordering
910
911
   #length of dataframe to be created: number of knots by number of timepoints (
912
        minus 1 for the intercept that we won't plot)
913
```

```
ln < -6*(length(coef(gm)))
914
915
   basis plot <- data.frame(Basis=integer(ln),
916
                              value_orig=double(ln),
917
                             time=integer(ln),
918
                              cof=double(ln)
   )
920
   basis_plot$time<-rep(time) #pasting timepoints
922
   basis_plot$Basis<-factor(rep(c(1:5),each=6)) #pasting basis number values
923
   basis_plot$value_orig<-basis$value #pasting basis values
924
   basis_plot$cof<-rep(coef(gm)[1:5],each=6) #pasting coefficients
925
   basis_plot<-basis_plot%>%
926
     mutate(mod_val=value_orig*cof) #the create the predicted values the bases
927
         need to be
   #multiplied by the coefficients
929
930
   #creating labeller to change the labels in the basis plots
931
932
   basis names<-c(
933
      '1'= "Intercept",
934
      2' = 1',
935
      "3 '= "2 °
      4' = 3"
937
      '5'="4"
938
939
   #calculating the final smooth by aggregating the basis functions
941
942
   smooth<-basis_plot%>%
943
     group_by(time)%>%
944
     summarize (smooth=sum(mod_val))
945
946
   #original basis
948
   sz < -1
949
   p11<-ggplot (basis_plot,
950
                 aes(x=time,
951
                     y=value_orig,
952
                     colour=as.factor(Basis)
953
954
                 )+
     geom_line(size=sz,
956
                 show.legend=FALSE)+
957
     geom\_point(size=sz+1,
958
                  show.legend = FALSE)+
     labs (y='Basis_functions')+
960
      facet_wrap(~Basis,
961
                  labeller = as_labeller(basis_names)
962
963
     theme_classic()+
964
     scale_color_aaas()
965
966
```

```
#penalized basis
968
    p12<-ggplot (basis_plot,
969
                  aes(x=time,
970
                       y=mod_val,
971
                       colour=as.factor(Basis)
972
973
                  )+
974
      geom\_line(show.legend = FALSE,
975
                  size=sz)+
976
      geom_point (show.legend = FALSE,
977
                    size=sz+1)+
978
      labs(y='Penalized_{\square} \setminus n_{\square} basis_{\square} functions')+
979
      scale_y_continuous (breaks=seq(-1,1,1))+
980
      facet_wrap(~Basis,
981
                    labeller=as_labeller(basis_names)
982
                    )+
983
      theme_classic()+
984
      scale_color_aaas()
985
    #heatmap of the penalization coefficient
987
    x_labels<-c("Intercept", "1", "2", "3", "4")
988
    p13<-ggplot (basis_plot,
989
                  aes (x=Basis,
                       v=Basis.
991
                        fill=cof))+
992
      geom_tile (aes (color='black'),
993
                  size=sz+1,
                  show.legend = FALSE)+
995
      geom\_tile(size=sz+1)+
996
      scale_fill_gradient(low = "white", high = "red")+
997
      labs (x='Basis',
998
            y='Basis')+
999
      scale_x_discrete(labels=x_labels)+
1000
      geom\_text(aes(label=round(cof,2)),
                  size = 10.
1002
                  show.legend = FALSE)+
1003
      theme classic ()+
1004
      theme(legend.title = element_blank())
1005
1006
    #plotting simulated datapoints and smooth term
    p14<-ggplot (data=dat,
1008
                  aes(x=time,y=y))+
1009
      geom\_point(size=sz+1)+
1010
      scale_color_aaas()+
1011
      labs (y='Simulated_\\n_response')+
1012
      geom_line (data=smooth,
1013
                  aes(x=time,
1014
                       y=smooth),
1015
                  color="#B15731",
1016
                   size=sz+1)+
1017
      theme_classic()
1018
1019
1020
    \#Combining \ all
1021
```

```
b_plot<-p11+p13+p12+p14+plot_annotation(tag_levels='A')& theme(
text=element_text(size=18)

)
```

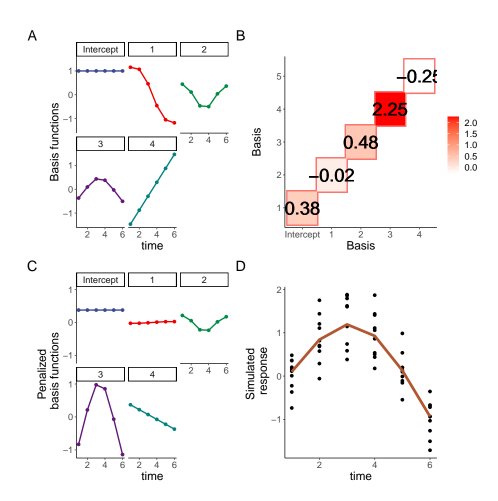


Figure 7: Basis functions for a single smoother for time with five knots. A: Basis functions for a single smoother for time for the simulated data of Group 1 from Figure 2, the intercept basis is not shown. B: Penalty matrix for the basis functions. Each basis function is penalized by a coefficient which can be positive or negative. The coefficient determines the overall effect of each basis in the final smoother. C: Penalized basis functions. Each of the four basis functions of panel A has been penalized by the corresponding coefficient shown in Panel B, note the corresponding increase (or decrease) of each basis. D: Smoother for time and original data points. The smoother (line) is the result of the sum of each penalized basis function at each time point, simulated values for the group appear as points.

B Longitudinal biomedical data simulation and GAMs

This section describes how to fit GAMs to longitudinal data using simulated data. First, data is simulated according to Section 5, where reported data of oxygen saturation (StO_2) in tumors under either chemotherapy or saline control is used as a starting point to generate individual responses in each group.

```
)
1034
1035
1036
    ## plot the mean response
1037
    f1 <-- ggplot (dat,
1038
                 aes(x = Day)
                      y = StO2,
1040
                      color = Group)) +
1041
         geom_line(size=1,
1042
                    show.legend = FALSE)+
1043
         geom\_point(show.legend = FALSE,
1044
                      size = 1.5,
1045
                      alpha = 0.5) +
1046
      labs (y=expression (paste (StO [2]),
1047
                                   '_(real)'))+
1048
      theme_classic()+
1049
      scale_color_aaas()+
1050
         scale x continuous (breaks=\mathbf{c}(0.5.10))+
1051
         scale_y_continuous (breaks=c(0,40))+
1052
      plot_layout (tag_level = 'new')+
1053
      theme (
1054
         plot.background = element_rect(fill = "transparent",
1055
                                              color = NA),
         axis.text=element text(size=14)
1057
1058
1059
1060
    #This function simulates data for the tumor data using default parameters of
1061
        10 observations per time point, and Standard deviation (sd) of 5%.
1062
    #Because physiologically StO2 cannot go below 0%, data is generated with a
1063
        cutoff value of 0.0001 (the "StO2_sim")
1064
1065
    simulate\_data \leftarrow function(dat, n = 10, sd = 5) {
1066
         dat_sim <- dat %>%
             slice (rep (1:n(), each = n)) %>%
1068
             group_by(Group, Day) %%
1069
             mutate (
1070
                      StO2\_sim = pmax(rnorm(n, StO2, sd), 0.0001),
1071
                      subject = rep(1:10),
1072
                      subject=factor(paste(subject, Group, sep = "-"))
1073
                      ) %>%
1074
             ungroup()
1075
1076
         return (dat_sim)
1077
    }
1078
1079
1080
    \#subject = factor(paste(subject, treatment, sep = "-")))
1081
1082
    n \leftarrow 10 \# number \ of \ observations
1083
    sd <- 10 #approximate sd from paper
1084
    set.seed(1) #set seed for reproducibility
1085
    df <- 6
1086
    dat_sim \leftarrow simulate_data(dat, n, sd)
```

```
1088
    #plotting simulated data
1089
    f2 <-- ggplot (dat sim,
1090
                  aes(x = Day,
                      y = StO2 \sin
1092
                       color = Group)) +
1093
         geom point (show.legend=FALSE,
1094
                       size = 1.5,
1095
                       alpha = 0.5) +
1096
         stat\_summary(aes(y = StO2\_sim,
1097
                              group=Group),
1098
                         fun=mean, geom="line",
1099
                         size=1.
1100
                         show.legend = FALSE)+
1101
      labs (y=expression (atop (StO [2]),
1102
                                   '(simulated)')))+
1103
      theme_classic()+
1104
      theme (
1105
         axis.text=element text(size=22)
1106
1107
      scale_color_aaas()+
1108
         scale_x continuous (breaks=c(0,2,5,7,10))
1109
```

1110 B.1 A basic Workflow for GAMs

This section shows a basic workflow to fit a series of increasingly complex GAMs to the simulated data from the previous section. Graphical and parameter diagnostics for goodness of fit are discussed, as well as model comparison via AIC (Aikake Information Criterion).

1114 B.1.1 First model

The first model fitted to the data is one that only accounts for different smooths by day. The model syntax specifies that gam_00 is the object that will contain all the model information, and that the model attempts to explain changes in StO2_sim (simulated StO₂) using a smooth per Day. The model will use 5 knots (k=5) for the smooth. And that the smooth is constructed using gaussian process basis (bs="gp"). The smoothing parameter estimation method used is the restricted maximum likelihood (REML).

```
gam_0 = gam(StO2\_sim \sim s(Day, k = 5, bs="gp"), method='REML', 
data = dat sim)
```

To obtain model diagnostics, two methodologies are to be used: 1) graphical diagnostics, and 2) a model check. In the first case, the functions appraise and draw from the package *gratia* can be used to obtain a single output with all the graphical diagnostics. For model check, the functions gam.check and summary from *mgcv* provide detailed information about the model fit and its parameters.

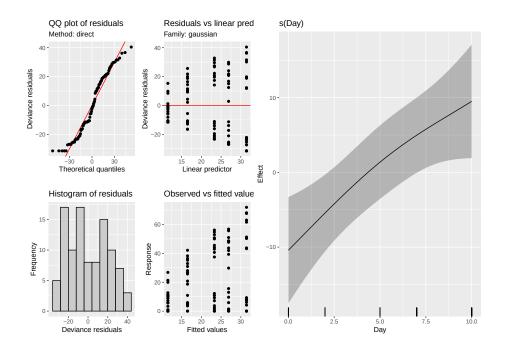


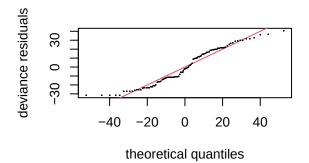
Figure 8: Graphical diagnostics for the first GAM model. Left: Graphical diagnostics provided by the function appraise from the package *gratia*. Right: Fitted smooth for the model, provided by the function draw.

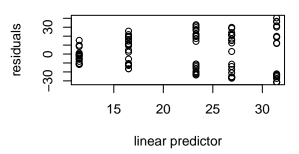
B.1.1.1 Graphical diagnostics From the output of the function appraise in Figure 8, the major indicators of concern about the model are the QQ plot of residuals and the histogram of residuals. The QQ plot shows that the errors are not reasonably located along the 45° line (which indicates normality), as there are multiple points that deviate from the trend, specially in the tails. The histogram also shows that the variation (residuals) is not following the assumption of a normal distribution.

The draw function permits to plot the smooths as ggplot2 objects, which eases subsequent manipulation, if desired. Because model gam_00 specifies only one smooth for the time covariate (Day), the plot only contains only one smooth. Note that the smooth shows an almost linear profile.

```
#need to add figure number and caption \operatorname{gam.\mathbf{check}}(\operatorname{gam}_{0}0)
```

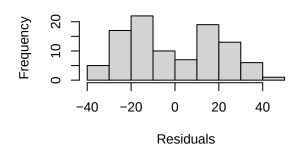
Resids vs. linear pred.





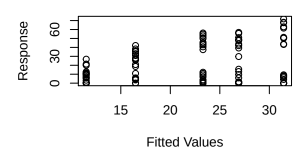
Histogram of residuals

Response vs. Fitted Values



1137

##



```
1138
   ## Method: REML
                       Optimizer: outer newton
1139
   ## full convergence after 6 iterations.
   ## Gradient range [-4.142968e-08,2.799316e-12]
1141
      (score 440.4108 & scale 414.2575).
   ## Hessian positive definite, eigenvalue range [0.04576008,49.0005].
1143
   \#\# \text{ Model rank} = 5 / 5
   ##
1145
   ## Basis dimension (k) checking results. Low p-value (k-index<1) may
   ## indicate that k is too low, especially if edf is close to k'.
1147
   ##
                     edf k-index p-value
   ##
1149
                             0.26 < 2e-16 ***
1150
   ## s(Day) 4.00 1.31
1151
                        0 '*** 0.001 '** 0.01 '* 0.05 '. 0.1 ' 1
   ## Signif. codes:
1152
   summary(gam_00)
1153
1154
   ## Family: gaussian
1155
   ## Link function: identity
1157
   ## Formula:
   ## StO2_sim \sim s(Day, k = 5, bs = "gp")
1159
   ##
       Parametric coefficients:
   ##
1161
                    Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept)
                       21.929
                                    2.035
                                             10.77
                                                      <2e-16 ***
1163
   ##
1164
                                                    '*' 0.05 '.' 0.1 ' '1
   ##
       Signif. codes:
                         0 '*** 0.001 '** 0.01
1165
   ##
   ##
       Approximate significance of smooth terms:
1167
                 edf Ref.df
                                  F p-value
   ##
1168
      s (Day) 1.314
                      1.536 9.151 0.00253 **
   ##
1169
   ##
1170
   ##
      Signif. codes:
                         0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' 1
1171
   ##
1172
   \# R-sq.(adj) = 0.105
                                Deviance explained = 11.7\%
1173
                         Scale est. = 414.26
   \#\# - REML = 440.41
                                                   n = 100
1174
```

Special attention must be paid to the 'k-index' from gam.check. This parameter indicates if the basis dimension of the smooth is adequate, i.e., it checks that the basis used to create the smooth are adequate to capture the trends in the data. If the model is not adequately capturing the trens in the data, this is indicated by a low k-index value (<1). From the output, it can be seen that the k-index is effectively <0.3, which indicates that the model is not capturing the variability in the data. The 'edf' (effective degrees of freedom) is an indicator of the complexity of the smooth. Here the complexity of the smooth is comparable to that of a 4th degree polynomial.

From the summary function, information about the assumed distribution of the errors (Gaussian in this case) and the link function can be obtained. The link function is 'identity' as the model does not make any transformation on the predictors. The 'significance of smooth terms' p-value indicates if each smooth is adding significance to the model. Here, the p-value is low but we have seen that there are issues with the model from the previous outputs. Finally, the 'deviance explained' indicates how much of the data the model is able to capture, which in this case corresponds to $\sim 12\%$.

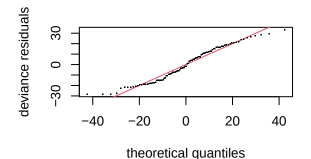
1189 B.1.2 Second model

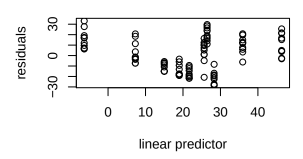
1188

The major flaw of gam_00 is that this model is not taking into account the fact that the data is nested in groups. The next iteration is a model where a different smooth of time (Day) is assigned for each group using by=Group in the model syntax.

```
1193 gam_01 \leftarrow gam(StO2\_sim \sim s(Day, by=Group, k = 5, bs="gp"),
1194 method= REML',
1195 data = dat\_sim)
1196
1197 gam.check(gam 01)
```

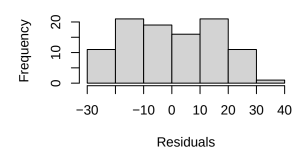
Resids vs. linear pred.

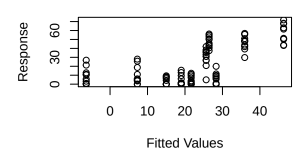




Histogram of residuals

Response vs. Fitted Values





```
1199
   ## Method: REML
                       Optimizer: outer newton
1200
   ## full convergence after 10 iterations.
   ## Gradient range [-0.0001703751, 9.561998e-05]
1202
       (score 418.612 & scale 270.7177).
   ## Hessian positive definite, eigenvalue range [0.0001702821,48.50255].
1204
   \#\# \text{ Model rank} = 9 / 9
   ##
1206
   ## Basis dimension (k) checking results. Low p-value (k-index<1) may
   ## indicate that k is too low, especially if edf is close to k'.
1208
   ##
1209
                                 k'
                                      edf k-index p-value
   ##
1210
   ## s(Day): GroupControl
                               4.00 \ 1.00
                                             0.32
                                                    <2e-16 ***
   ## s(Day):GroupTreatment 4.00 1.72
                                             0.32
                                                    <2e-16 ***
1212
1213
   ## Signif. codes:
                        0 '***' 0.001
                                        '**' 0.01
                                                  '*' 0.05 '.' 0.1 ' '1
1214
   summary(gam_01)
   ##
1216
   ## Family: gaussian
   ## Link function: identity
1218
   ## Formula:
1220
   ## StO2_sim \sim s(Day, by = Group, k = 5, bs = "gp")
1222
      Parametric coefficients:
```

```
Estimate Std. Error t value Pr(>|t|)
1224
                        21.929
                                       1.645
                                                13.33
   ##
       (Intercept)
                                                          < 2e - 16
1225
   ##
1226
                                    0.001
                                                 0.01
                                                        '*' 0.05 '.' 0.1 ' '1
       Signif. codes:
                          0
   ##
   ##
1228
       Approximate significance of smooth terms:
   ##
1229
                                    edf Ref. df
   ##
                                                       F p-value
1230
       s (Day): Group Control
                                          1.001
                                  1.001
                                                   4.099
                                                           0.0456
1231
       s (Day): Group Treatment 1.715
                                          1.979
                                                 35.551
                                                           <2e-16 ***
1232
1233
       Signif. codes:
                                                            0.05
1234
1235
   \# R - sq.(adj) =
                        0.415
                                  Deviance explained = 43.1\%
1236
                          Scale est. = 270.72
   \#\# - REML = 418.61
1237
```

Diagnostics for this model indicate that the k-index is still below 1 (0.32 from gam.check), and that the residuals are still not following a normal distribution (Figure 9). Moreover, the smooths (plotted via the draw() function) appear with a fairly linear profile, which indicates they are still not capturing the trends observed in the data. From summary(), the deviance explained by the model is ~43%.

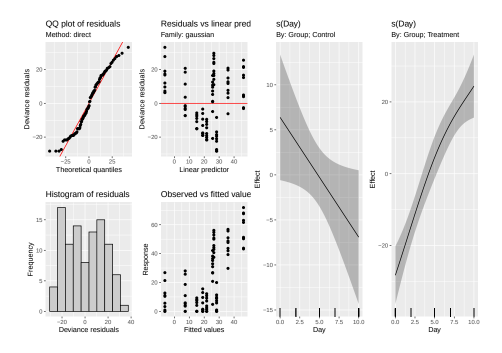


Figure 9: Graphical diagnostics for the second GAM model. Left: Graphical diagnostics provided by the function appraise from the package *gratia*. Right: Fitted smooth for the model, provided by the function draw.

B.1.3 Third model

1242

1244

1245

1247

Model gam_00 was built for didactic purposes to cover the simplest case, but it does not account for the nesting of the data by Group, which is apparent from the type of smooth fitted, the model diagnostics, and, the low variance explained by the model. On the other hand, gam_01 takes into account the nesting within each group and provides better variance explanation, but as indicated in Section 5, in order to differentiate between each group a parametric term needs to be added to the model for the interaction of *Day* and *Group*.

```
#GAM for StO2

1250

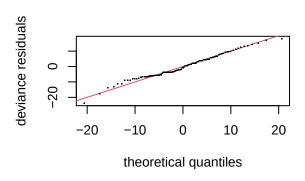
1251 gam1 <— gam(StO2_sim ~ Group+s(Day, by = Group, k = 5,bs="gp"),

1252 method='REML',

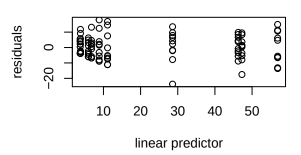
1253 data = dat_sim)

1254

1255 gam.check(gam1)
```

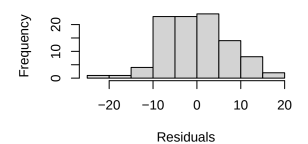


Resids vs. linear pred.



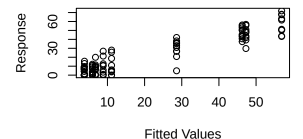
Histogram of residuals

Response vs. Fitted Values



1256

1272 ##



```
1257
   ## Method: REML
                       Optimizer: outer newton
      full convergence after 9 iterations.
1259
       Gradient range [-1.003557e-07,3.562136e-08]
       (score 362.7587 & scale 64.03804).
1261
   ## Hessian positive definite, eigenvalue range [0.9494021,48.08513].
       Model rank = 10 / 10
1263
   ##
1264
      Basis dimension (k) checking results. Low p-value (k-index<1) may
1265
      indicate that k is too low, especially if edf is close to k'.
1266
   ##
1267
   ##
                                 k'
                                      edf k-index p-value
1268
   ## s(Day): GroupControl
                               4.00 \ 3.83
                                              1.02
                                                      0.52
1269
   ## s(Day):GroupTreatment 4.00 3.84
                                             1.02
                                                      0.59
1270
   summary (gam1)
1271
```

```
## Family: gaussian
1273
      Link function: identity
1274
   ##
1275
   ## Formula:
       StO2\_sim \sim Group + s(Day, by = Group, k = 5, bs = "gp")
1277
       Parametric coefficients:
   ##
1279
                        Estimate Std. Error t value Pr(>|t|)
1280
       (Intercept)
                            9.781
                                         1.132
                                                  8.643
                                                         1.85e - 13 ***
1281
       GroupTreatment
                           24.296
                                         1.600
                                                 15.181
                                                          < 2e-16 ***
1282
1283
                            '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1
       Signif. codes:
1284
1285
       Approximate significance of smooth terms:
1286
                                   edf Ref.df
   ## s(Day): GroupControl
                                 3.825
                                         3.971
                                               16.81
                                                        <2e-16 ***
1288
   ## s(Day): GroupTreatment 3.835
                                         3.974 78.84
                                                        <2e-16 ***
1290
                            '*** ' 0.001 '** ' 0.01
                                                     '*' 0.05 '.' 0.1 ' '1
       Signif. codes:
1291
1292
   \#\# R-sq.(adj) =
                       0.862
                                 Deviance explained = 87.4\%
1293
   \#\# - REML = 362.76
                          Scale est. = 64.038
1294
```

The resulting model is model gam1, which is the model fitted in the main manuscript. By using appraise() and draw on this model (Figure 10) we see that the trend on the QQ plot has improved, the histogram of the residuals appears to be reasonably distributed, and the smooths are capturing the trend of the data within each group . From gam.check, the k-index is now at an acceptable value (\sim 1.02), and summary now indicates that the model is able to capture 87% of the variance data.

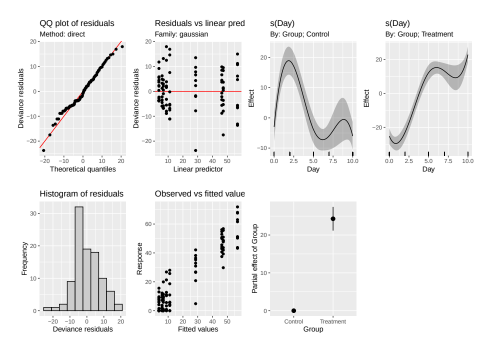


Figure 10: Graphical diagnostics for the final GAM model. Left: Graphical diagnostics provided by the function appraise from the package *gratia*. Right: Fitted smooths for the model, provided by the function draw.

1295

1297

B.1.4 Comparing models via AIC

One final comparison that can be made for model selection involves the use of the Aikake Information Criterion (AIC). This metric is used to estimate information loss, which we want to minimize with an appropriate model. Therefore, when 2 or more models are compared, the model with lower AIC is preferred.

In R, the comparison is done using the AIC function.

```
AIC (gam_00,gam_01,gam1)

1307 ## df AIC
1308 ## gam_00 3.536147 891.1671
1309 ## gam_01 4.980481 850.0698
1310 ## gam1 10.945191 711.4662
```

1315

1316

1321

The output in this case is expected: model gam1 has a lower AIC (711.46) whereas the initial two models have higher AICs (891 and 850). The AIC should not be considered as the only estimator of model goodness, instead to be used as complimentary information to the graphical diagnostics and model checks described above.

C GAM and Linear model plots and Missing data

This section covers the code used to generate Figure 3, where the simulated data, fit of the "final" GAM (gam1), linear model and GAM on data with missing observations are presented. Note that panel A in Figure 3 and the inlet are generated in the code chunk where the data is simulated in Section B, and are called later to build the figure.

C.1 GAM and Linear model plots

This code chunk creates panels B and D in Figure 3. Note that this code uses the final GAM from the previous section (gam1), so the simulated data and the model should be generated before running this section.

```
\#linear model
   lm1 \leftarrow lm(StO2\_sim \sim Day + Group + Day * Group, data = dat\_sim)
1327
1328
1329
   #creates a dataframe using the length of the covariates for the GAM
1330
   gam_predict <- expand_grid (Group = factor(c("Control", "Treatment")),
1331
                                Day = seq(0, 10, by = 0.1),
1332
                                subject = factor(rep(1:10))
1333
1334
    \#creates a dataframe using the length of the covariates for rm–ANOVA
    lm_predict<=expand_grid(Group = factor(c("Control", "Treatment")),</pre>
1336
                                Day = c(0:10),
1337
                               subject = factor(rep(1:10)),
1338
   lm_predict$subject<-factor(paste(lm_predict$subject, lm_predict$Group, sep =</pre>
1340
       -"))
1341
1342
    \#adds the predictions to the grid and creates a confidence interval for GAM
```

```
gam_predict<-gam_predict%>%
1344
         mutate(fit = predict(gam1,gam_predict,se.fit = TRUE,type='response')$fit
1345
                 se. fit = predict (gam1, gam predict, se. fit = TRUE, type='response')$
1346
                     se. fit)
1348
    \#using\ lm
1349
    lm predict<-lm predict%>%
1350
         mutate(fit = predict(lm1,lm_predict, se. fit = TRUE, type='response')$fit,
1351
                 se. fit = predict(lm1, lm_predict, se. fit = TRUE, type='response')$se.
1352
                     fit)
1353
1354
    #plot smooths and confidence interval for GAM
1355
    f3<-ggplot(data=dat_sim, aes(x=Day, y=StO2_sim, group=Group)) +
1356
         geom\_point(aes(color=Group), size=1.5, alpha=0.5, show.legend = FALSE)+
1357
      geom\_ribbon(aes(x=Day,ymin=(fit - 2*se.fit),
1358
                          ymax = (fit + 2*se.fit),
1359
                          fill=Group
                          ),
1361
                    alpha = 0.3,
1362
                    data=gam predict,
1363
                  show.legend=FALSE,
1364
                       inherit.aes=FALSE) +
1365
      geom_line(aes(y=fit,
1366
                       color=Group),
1367
                    size = 1, data = gam\_predict,
1368
                    show.legend = FALSE)+
1369
      \#facet\_wrap(\sim Group)+
1370
      labs (y=expression (atop (StO[2], 'complete')))+
1371
         scale_x_continuous(breaks=c(0,2,5,7,10))+
1372
           theme_classic()+
1373
      theme (
1374
         axis.text = element\_text (size = 22)
1375
1376
           scale_color_aaas()+
      scale_fill_aaas()
1378
1379
    #plot linear fit for rm-ANOVA
1380
    f4<-ggplot(data=dat_sim, aes(x=Day, y=StO2_sim, group=Group)) +
1381
         geom\_point(aes(color=Group), size=1.5, alpha=0.5, show.legend = FALSE)+
1382
      geom\_ribbon(aes(x=Day,ymin=(fit - 2*se.fit)),
1383
                          vmax = (fit + 2*se.fit), fill = Group),
1384
                    alpha = 0.3,
1385
                    data=lm predict,
1386
                    show.legend = FALSE,
1387
                       inherit.aes=FALSE) +
1388
      geom_line(aes(y=fit,
1389
                       color=Group),
1390
                    size = 1, data = lm\_predict,
1391
                    show.legend = FALSE)+
1392
      \#facet\_wrap(\sim Group) +
1393
      labs (y=expression (paste ('StO'[2], '\(\) (simulated)')))+
1394
         scale_x_continuous (breaks=c(0,2,5,7,10))+
1395
           theme classic()+
1396
      theme (
1397
```

```
axis.text=element_text(size=22)
1398
       )+
1399
             scale color aaas()+
1400
        scale__fill_aaas()
1401
1402
1403
1404
     #posthoc comparisons for the linear model
1405
     library (multcomp)
1406
1407
1408
    \#summary(glht(lm1, linfct = mcp(Group = 'Tukey')))
1409
     \#summary(\ g\ lh\ t\ (\ lm\ 1\ , \quad li\ nf\ c\ t=mcp\left(\ Group="Tukey", \quad in\ te\ ra\ c\ ti\ o\ n\_a\ ve\ ra\ g\ e=TRUE)\ )\ )
1410
1411
```

C.2Working with Missing data in GAMs

1412

##

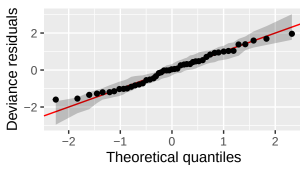
This code chunk randomly deletes 40% of the total observations in the original simulated data, and then an 1413 interaction GAM is fitted. Model diagnostics are presented, and an object that stores the fitted smooths is 1414 saved to be called in the final code chunk to build the figure. 1415

```
#missing data
1416
    \#create a sequence of 40 random numbers between 1 and 100, these numbers will
1417
   #correspond to the row numbers to be randomly erased from the original dataset
1418
    missing \leftarrow sample(1:100, 40)
1419
   #create a new dataframe from the simulated data with 40 rows randomly removed,
1421
         keep the missing values as NA
1422
1423
    ind <- which (dat_sim$StO2_sim %in% sample(dat_sim$StO2_sim, 40))
1424
1425
   #create a new dataframe, remove the StO2 column
1426
    dat_missing \leftarrow dat_sim[,-1]
1427
   #add NAs at the ind positions
1429
    dat_missing$StO2_sim [ind]<-NA
1431
    \#Count the number of remaining observations per day (original dataset had 10
1432
       per group per day)
1433
    dat_missing %%
1434
        group_by(Day, Group) %>%
1435
         filter (!is.na(StO2_sim))%>%
1436
      count (Day)
1437
   ## # A tibble: 10 x 3
1438
   ## # Groups:
                     Day, Group [10]
1439
             Day Group
   ##
                                 n
1440
   ##
          < dbl > < fct >
                             <int>
1441
               0 Control
                                  2
   ##
        1
1442
        2
               0 Treatment
                                  4
   ##
1443
               2 Control
                                 6
   ##
        3
        4
               2 Treatment
                                  5
   ##
1445
               5 Control
                                 6
        5
```

```
5 Treatment
1447
                  Control
                                  3
1448
                 Treatment
                                  5
1449
              10 Control
                                  3
        9
                                  8
       10
              10
                 Treatment
1451
   #the same model used for the full dataset
1452
   mod_m1 <- gam(StO2_sim ~ Group+s(Day, by=Group, k=5), data = dat_missing, family
1453
       =scat)
1454
   #appraise the model
1455
    appraise (mod_m1)
1456
```

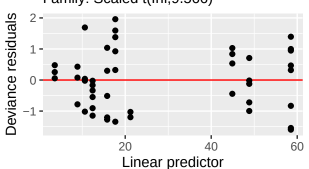
QQ plot of residuals

Method: simulate

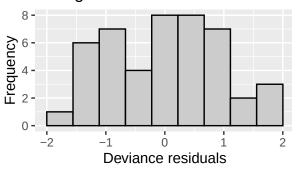


Residuals vs linear predictor

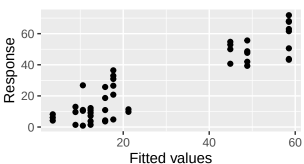
Family: Scaled t(Inf,9.566)



Histogram of residuals



Observed vs fitted values



 $\begin{array}{l} \text{f6} \leftarrow \text{ggplot}\left(\mathbf{data} = \text{dat_missing} \;,\;\; \operatorname{aes}\left(\mathbf{x} = \text{Day} \;,\;\; \mathbf{y} = \text{StO2_sim} \;,\;\; \operatorname{group} = \text{Group}\right) \;,\; \\ \text{geom_point}\left(\operatorname{aes}\left(\operatorname{color} = \text{Group}\right) \;,\; \operatorname{size} = 1.5 \;,\; \operatorname{alpha} = 0.5 \;, \\ \text{show} \;.\; \mathbf{legend} \;=\; \text{FALSE}\right) + \\ \text{geom_ribbon}\left(\operatorname{aes}\left(\;\; \mathbf{x} = \text{Day} \;,\; \mathbf{ymin} = \left(\operatorname{fit} \;-\; 2 * \mathbf{se} \;.\; \operatorname{fit}\right) \;,\;\; \\ \text{ymax} = \left(\operatorname{fit} \;+\; 2 * \mathbf{se} \;.\; \operatorname{fit}\right) \;, \\ \end{array}$

```
fill=Group
1472
                            ),
1473
                     alpha = 0.3,
1474
                     data=m_predict,
1475
                   show.legend=FALSE,
1476
                        inherit.aes=FALSE) +
1477
      geom_line(aes(y=fit,
1478
                        color=Group),
1479
                     size = 1, data = m_predict,
1480
                     show.legend = TRUE)+
1481
      \#facet\_wrap(\sim Group)+
1482
      labs(y=expression(atop(StO[2], 'missing')))+
1483
         scale_x_continuous (breaks=c(0,2,5,7,10))+
1484
           theme_classic()+
1485
      theme (
1486
         axis.text = element\_text(size = 22)
1487
1488
            scale_color_aaas()+
1489
      scale__fill_aaas()
1490
    mult_plot<-f2+inset_element (
1491
      f1, left = 0.01,
1492
      bottom = 0.5,
      right = 0.5,
1494
      top = 1.0) +
1495
       f3+f4+f6+
1496
        plot_annotation(tag_levels='A')&
1497
        y \lim (\mathbf{c}(-5,75)) \&
1498
      theme (
1499
          text=element_text (size=18)
1500
1501
       scale_color_aaas()
1502
1503
    mult\_plot
1504
```

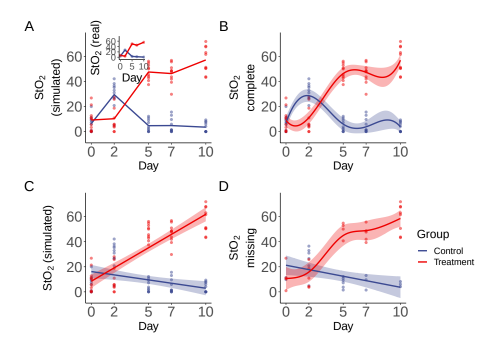


Figure 11: Simulated data and smooths for oxygen saturation in tumors. A: Simulated data that follows previously reported trends (inset) in tumors under chemotherapy (Treatment) or saline (Control) treatment. Simulated data is from a normal distribution with standard deviation of 10% with 10 observations per time point. Lines indicate mean oxygen saturation B: Smooths from the GAM model for the full simulated data with interaction of Group and Treatment. Lines represent trends for each group, shaded regions are 95% confidence intervals. C: rm-ANOVA model for the simulated data, the model does not capture the changes in each group over time. D: Smooths for the GAM model for the simulated data with 40% of its observations missing. Lines represent trends for each group, shaded regions are 95% confidence intervals.