SEMIPARAMETRIC MODELING OF AUTONOMOUS NONLINEAR DYNAMICAL SYSTEMS WITH APPLICATION TO PLANT GROWTH

BY DEBASHIS PAUL, JIE PENG, AND PRABIR BURMAN

University of California, Davis

We propose a semiparametric model for autonomous nonlinear dynamical systems and devise an estimation procedure for model fitting. This model incorporates subject-specific effects and can be viewed as a nonlinear semiparametric mixed effects model. We also propose a computationally efficient model selection procedure. We show by simulation studies that the proposed estimation as well as model selection procedures can efficiently handle sparse and noisy measurements. Finally, we apply the proposed method to a plant growth data used to study growth displacement rates within meristems of maize roots under two different experimental conditions.

1. Introduction. Continuous time dynamical systems arise, among other places, in modeling certain biological processes. This includes classical examples from population biology like the Lotka–Volterra equations for describing prey-predator dynamics [Perthame (2007)], or subject-specific processes like the progression of infectious diseases in individuals [Nowak and May (2000)]. Most of the existing approaches estimate the dynamical system by assuming known functional forms of the system. Moreover, many of them aim at estimating individual dynamics for one subject. However, in many scientific studies, there is a need to model the dynamical system nonparametrically due to insufficient knowledge of the problem at hand. In addition, there could be an interest to know the dynamics of a certain process at a population level in order to answer various scientific questions. Thus, in this paper, we propose a new method to bridge the gap and tackle these challenges.

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To motivate the model, we first briefly discuss a study on plant growth. There is a lot of research aiming to understand the effect of environmental conditions on the growth in a plant. For example, root growth in plants is highly sensitive to environmental factors such as temperature, water deficit or nutrients [Schurr, Walter and Rascher (2006); Walter et al. (2002)]. In Sacks, Silk and Burman (1997), an experiment is conducted to study the effect of water stress on cortical cell division rates through growth displacement rate within the meristem of the primary root of maize seedlings (Figure 1: left panel). In this study, for each plant, measurements are taken on the displacement, measured as the distance in millimeters from the root cap junction (root apex) of a number of markers on the root over a period of 12 hours (Figure 1: right panel). The growth displacement rate is defined as the rate of displacement of a particle placed along the root and, thus, it is a function of distance from the root apex. By its definition, growth displacement rate characterizes the relationship between the growth trajectory and its derivative (with respect to time). Therefore, it is the gradient function in the corresponding dynamical system. In this study there is a need to understand the dynamics at the population level, while accounting for subject-specific variations, in order to compare the growth displacement rates under two different water conditions.

Motivated by this study, in this paper, we focus on modeling and fitting the underlying dynamical system based on data measured over time, referred to as sample curves or sample paths, for a group of subjects. Moreover, for a given sample curve, instead of observing the whole sample path, measurements are taken only at a sparse set of time points together with possible measurement noise. In the plant data that we just mentioned, each plant is a subject, and the positions of the markers which are located at different
distances at time zero from the root cap junction correspond to different initial conditions. Each marker corresponds to one displacement trajectory (also referred to as growth trajectory/curve), and the number of measurements varies from two to seventeen, with measurement times varying across trajectories. (See Section 5 for a more detailed description.)

We first give a brief overview of the existing literature on fitting smooth deterministic dynamical systems in continuous time. A large number of physical, chemical or biological processes are modeled through systems of parametric differential equations [Ljung and Glad (1994); Perthame (2007); Strogatz (2001)]. For example, Ramsay et al. (2007) consider modeling a continuously stirred tank reactor and propose a method called parameter cascading for model fitting. Zhu and Wu (2007) adopt a state space approach for estimating the dynamics of cell-virus interactions in an AIDS clinical trial. Ramsay and Silverman (2002, 2005) consider fitting dynamical systems given by systems of linear differential equations where the coefficients of the differential operator may be time varying. They propose methods for estimating these (linear) differential operators based on principal differential analysis when the data are recorded at dense and regular time points. Poyton et al. (2006) also use the principal differential analysis approach to fit dynamical systems. Chen and Wu (2008b, 2008a) propose to estimate parametric differential equations with known functional forms and time-dependent parameters through a two-stage approach where the first stage involves estimation of the sample trajectories and their derivatives by nonparametric smoothing. Brunel (2008) gives a comprehensive theoretical analysis of such an approach. Cao, Fussmann and Ramsay (2008) propose a method for fitting nonlinear dynamical systems using splines with predetermined knots for describing the gradient function. This involves knowing the functional form of the differential equation and does not include any subject-specific effects. Wu and Ding (1999) and Wu, Ding and DeGruttola (1998) propose using the nonlinear least squares procedure for fitting parametric differential equations that take into account subject-specific effects.

For the problems that we address in this paper, measurements are taken on a sparse set of points for each sample curve so that estimation of individual sample trajectory or its derivative based on nonparametric smoothing is error-prone and results in a loss of information. Thus, numerical procedures for solving differential equations can become unstable if we treat each sample curve separately. Moreover, we are more interested in estimating the baseline dynamics at the population level than the individual dynamics of each subject. For example, in the plant study described above, we are interested in comparing the growth displacement rates under two different experimental conditions. On the other hand, we are not so interested in the individual displacement rate corresponding to each plant. Another important aspect in modeling data with multiple subjects is that adequate measures need to be
taken to model possible subject-specific effects, otherwise the estimates of
the model parameters can have inflated variability. In this paper we propose
a semiparametric approach for modeling dynamical systems which incor-
porates subject-specific effects while combining information across different
subjects. A nonparametric model is often essential because of insufficient
knowledge about the problem to suggest a reasonable parametric form of
the dynamical system. In addition, if realistic parametric models can be
proposed, then the nonparametric fit can be used for diagnostics of lack of
fit, for example, by employing a distance measure between the parametric
and nonparametric fits and studying its sampling variability. We propose
an estimation procedure that combines nonlinear optimization techniques
with a numerical ODE (ordinary differential equation) solver to estimate
the unknown parameters. In addition, we derive a computationally efficient
approximation of the leave-one-curve-out cross-validation score for model
selection. We show by simulation studies that the proposed approach can
efficiently estimate the baseline dynamics with noisy and sparsely measured
sample curves. Finally, we apply the proposed method to the plant data and
compare the estimated growth displacement rates under the two experimen-
tal conditions and discuss some scientific implications of the results.

To the best of our knowledge, modeling and fitting dynamical systems
nonparametrically while also allowing for subject-specific effects is new in the
literature. In particular, our model differs from traditional nonlinear mixed
effects models previously employed for fitting differential equations, which
are almost exclusively parametric [Wu, Ding and DeGruttola (1998); Wu
and Ding (1999); Guedj, Thiebaut and Commenges (2007); Li et al. (2002)].
For example, Guedj, Thiebaut and Commenges (2007) consider a nonlinear
state-space model where the state variable follows a parametric differen-
tial equation with subject-specific effects, and the parameters are estimated
through a maximum likelihood approach. In contrast, for the model pro-
posed in this paper, the form of the gradient function $g$ is not assumed to
be known and it is approximated in a sequence of bases with growing di-

dimension. Note that this gives rise to a sequence of parametric models with
increasing complexity, and one needs to adopt a model selection procedure to
select an appropriate model, as is typical in nonparametric function estima-
tion. The theoretical derivations in Paul, Peng and Burman (2009) also show
that the problem of estimating the gradient function $g$ nonparametrically is
intrinsically different from that under a parametric nonlinear mixed-effects
model.

The rest of the paper is organized as follows. In Section 2 we describe
the proposed model. In Section 3 we discuss the model fitting and model
selection procedures. In Section 4 we conduct simulation studies to illustrate
finite sample performance of the proposed method and compare the proposed
method with a two-stage procedure. In Section 5 we apply this method to
the plant data. Section 6 has a brief discussion. More details and additional simulation results are reported in the supplementary material [Paul, Peng and Burman (2011)].

2. Model. In this section we describe a class of autonomous dynamical systems that is suitable for modeling the problems discussed in Section 1. An autonomous dynamical system has the following general form:

\[ X'(t) = f(X(t)), \quad t \in [T_0, T_1]. \]

Without loss of generality, henceforth \( T_0 = 0 \) and \( T_1 = 1 \). Note that the above equation implies that \( X(t) = a + \int_0^t f(X(u)) \, du \), where \( a = X(0) \) is the initial condition. In an autonomous system, the dynamics, which is characterized by \( f \), depends on time \( t \) only through the “state” \( X(t) \). This type of system arises in various scientific studies such as modeling prey-predator dynamics, virus dynamics or epidemiology [Perthame (2007)].

In this paper we consider the following class of autonomous dynamical systems:

\[ X_{il}'(t) = g_i(X_{il}(t)), \quad l = 1, \ldots, N_i, \quad i = 1, \ldots, n, \tag{2.1} \]

where \( \{X_{il}(t) : t \in [0,1], l = 1, \ldots, N_i; i = 1, \ldots, n\} \) is a collection of smooth curves corresponding to \( n \) subjects, where \( N_i \geq 1 \) is the number of curves associated with the \( i \)th subject. For example, in the plant study, each plant is a subject and each marker corresponds to one growth curve and there are multiple markers for each plant. We assume that all the curves associated with the same subject follow the same dynamics and are only differentiated by different initial conditions. These are described by the functions \( \{g_i(\cdot)\}_{i=1}^n \). In this paper we model \( \{g_i(\cdot)\}_{i=1}^n \) as

\[ g_i(\cdot) = e^{\theta_i} g(\cdot), \quad i = 1, \ldots, n, \tag{2.2} \]

where:

1. the function \( g(\cdot) \) reflects the common underlying mechanism regulating all these dynamical systems. It is assumed to be a smooth function and is referred to as the gradient function.
2. \( \theta_i \)'s reflect subject-specific effects in these systems. The mean of \( \theta_i \)'s is assumed to be zero to impose identifiability.

Note that one may view the trajectories for each plant as multivariate functional data. However, here for each subject, the different trajectories correspond to different initial conditions of the same ODE describing the system. This means that given the initial condition and the subject-specific scaling parameter \( \theta_i \), the corresponding trajectory is completely determined by the underlying dynamical system and the only source of randomness is from measurement errors.
The simplicity and generality of this model make it appealing for modeling a wide class of dynamical systems. First, the gradient function $g(\cdot)$ can be an arbitrary smooth function. Second, the scale parameter $e^{\theta}$ provides a subject-specific tuning of the dynamics. This is motivated by the fact that, for a large class of problems, the variations of the dynamics in a population are in the scale of the rate of change rather than in the shape of the gradient function. For example, for the plant data, by examining the scatter plot of empirical derivatives versus empirical fits (Figure 2, for more details, see Section 5), we observe an excessive variability toward the end which reflects plant-specific scaling effects. Moreover, the above model is also flexible in incorporating time-independent covariates, say, $z_i$, for example, by expressing the scaling factor as $e^{\eta^T z_i}$ for some parameter $\eta$. In this paper our primary goal is to estimate the gradient function $g$ nonparametrically.

Assuming the gradient function $g$ to be smooth means that it can be well-approximated by a basis representation approach:

$$g(x) \approx \sum_{k=1}^{M} \beta_k \phi_{k,M}(x),$$

where $\phi_{1,M}(\cdot), \ldots, \phi_{M,M}(\cdot)$ are linearly independent basis functions, chosen so that their combined support covers the range of the observed trajectories. For example, we can use cubic splines with a suitable set of knots. Thus, for a given choice of the basis functions, the unknown parameters in
the model are the basis coefficients \( \beta := (\beta_1, \ldots, \beta_M)^T \), the scale parameters \( \theta := \{\theta_i\}_{i=1}^n \), and possibly the initial conditions \( a := \{a_{il} := X_{il}(0) : l = 1, \ldots, N_i\}_{i=1}^n \). Also, various model parameters, such as the number of basis functions \( M \) and the knot sequence, need to be selected based on the data. Therefore, in essence, this is a nonlinear, semiparametric, mixed effects model.

3. Model fitting.

3.1. Estimation procedure. In this section we propose an estimation procedure based on sparsely observed noisy data. Specifically, we assume that the observations are given by

\[
Y_{ilj} = X_{il}(t_{ilj}) + \varepsilon_{ilj}, \quad j = 1, \ldots, m_{il},
\]

where \( 0 \leq t_{il1} < \cdots < t_{ilm_{il}} \leq 1 \) are the measurement times for the \( l \)th curve of the \( i \)th subject, and \( \{\varepsilon_{ilj}\} \) are independently and identically distributed noise with mean zero and variance \( \sigma_\varepsilon^2 > 0 \). For model fitting with such data, we adopt an iterative updating procedure which imposes regularization on the estimates of \( \theta \) and \( a \). One way to achieve this is to treat them as unknown random parameters from some parametric distributions. Specifically, we use the following set of working assumptions: (i) \( a_{il} \)'s are independent and identically distributed as \( N(\alpha, \sigma_a^2) \) and \( \theta_i \)'s are independent and identically distributed as \( N(0, \sigma_\theta^2) \), for some \( \alpha \in \mathbb{R} \) and \( \sigma_a^2 > 0, \sigma_\theta^2 > 0 \); (ii) the noise \( \varepsilon_{ilj} \)'s are independent and identically distributed as \( N(0, \sigma_\varepsilon^2) \) for \( \sigma_\varepsilon^2 > 0 \); (iii) the three random vectors \( a, \theta, \varepsilon := \{\varepsilon_{ilj}\} \) are independent. Under these assumptions, the negative joint log-likelihood of the observed data \( Y := \{Y_{ilj}\} \), the scale parameters \( \theta \) and the initial conditions \( a \) is (up to an additive constant and a positive scale constant)

\[
(3.2) \quad \sum_{i=1}^n \sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} [Y_{ilj} - \bar{X}_{il}(t_{ilj}; a_{il}, \theta_i, \beta)]^2 + \lambda_1 \sum_{i=1}^n \sum_{l=1}^{N_i} (a_{il} - \alpha)^2 + \lambda_2 \sum_{i=1}^n \theta_i^2,
\]

where \( \lambda_1 = \sigma_a^2/\sigma_\varepsilon^2, \lambda_2 = \sigma_\theta^2/\sigma_\varepsilon^2 \), and \( \bar{X}_{il}(\cdot) \) is the trajectory determined by \( a_{il}, \theta_i \) and \( \beta \). This can be viewed as a hierarchical maximum likelihood approach [Lee, Nelder and Pawitan (2006)], which is considered to be a convenient alternative to the full (restricted) maximum likelihood approach. Define

\[
\ell_{ilj}(a_{il}, \theta_i, \beta) := [Y_{ilj} - \bar{X}_{il}(t_{ilj}; a_{il}, \theta_i, \beta)]^2 + \lambda_1 (a_{il} - \alpha)^2/m_{il} + \lambda_2 \theta_i^2/\sum_{l=1}^{N_i} m_{il}.
\]

Then the loss function in (3.2) equals \( \sum_{i=1}^n \sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} \ell_{ilj}(a_{il}, \theta_i, \beta) \). Note that the above distributional assumptions are simply working assumptions,
since the expression in (3.2) can also be viewed as a regularized $\ell_2$ loss with penalties on the variability of $\theta$ and $a$.

In many problems there are natural constraints on the gradient function $g$. Some of these constraints can be expressed in the form of quadratic constraints in certain derivatives of $g$. Thus, to add flexibility to our estimation procedure, we allow for incorporating penalties of the form: $\beta^T B \beta$ for an $M \times M$ positive semi-definite matrix $B$ in the loss function. Consequently, the modified objective function becomes

$$L(a, \theta, \beta) := \sum_{i=1}^n \sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} \ell_{ilj}(a_{il}, \theta_i, \beta) + \beta^T B \beta.$$  

(3.3)

The proposed estimator is then the minimizer of the objective function:

$$\hat{(a, \theta, \beta)} := \arg \min_{a, \theta, \beta} L(a, \theta, \beta).$$  

(3.4)

Note that here our main interest is the gradient function $g$. Thus, estimating the parameters of the dynamical system together with the sample trajectories and their derivatives simultaneously is the most efficient. In contrast, in a two-stage approach, the trajectories and their derivatives are first obtained via pre-smoothing [see, e.g., Chen and Wu (2008b, 2008a); Varah (1982)], and then they are used in a nonparametric regression framework to derive an estimate of $g$. This is inefficient since estimation errors introduced in the pre-smoothing step effectively cause a loss of information. Indeed, simulation studies carried out in Section 4 and the supplementary material [Paul, Peng and Burman (2011)] show that two-stage estimators suffer from significant biases in estimating the gradient function $g$. Alternative ways of estimating $g$ include using the reproducing kernel Hilbert space framework [Gu (2002)], and controlling the degree of smoothness of the fitted $g$ by tuning a roughness penalty.

In the following, we propose a numerical procedure for solving (3.4) that has two main ingredients:

- Given $(a, \theta, \beta)$, reconstruct the trajectories \(\{\tilde{X}_{il}(\cdot) : l = 1, \ldots, N_i\}_{i=1}^n\) and their derivatives. This can be implemented using a numerical ODE solver, such as the Runge–Kutta method [Tenenbaum and Pollard (1985)].
- Minimize (3.3) with respect to $(a, \theta, \beta)$. This amounts to a nonlinear regression problem. It can be carried out using either a specialized nonlinear least squares solver, like the Levenberg–Marquardt method, [Nocedal and Wright (2006)] or a general optimization procedure, such as the Newton–Raphson algorithm.

The above fitting procedure bears similarity to the local, or gradient-based, methods discussed by Li et al. (2002), Guedj, Thiébaut and Commenges.
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(2007) and Miao et al. (2009), even though their works focus on parametric ODEs. The main distinction of the proposed framework and those of Li et al. (2002) and Guedj, Thiébaut and Commenges (2007) lies in that, for the current setting, the complexity of the model is allowed to grow with increasing sample size and one eventually needs to adopt a model selection procedure to select an appropriate model (as is done in this paper). From purely a model-fitting point of view, nonlinear mixed-effects (NLME) model-based estimation procedures may be used in principle to fit each of these parametric submodels. The work of Ke and Wang (2001) on semiparametric mixed-effects model fitting also shares some common computational challenges with our model. However, unlike in Ke and Wang (2001), in our case, the likelihood for the nonparametric component (i.e., the gradient function) is not available in closed form.

We now briefly describe an optimization procedure based on the idea of the Levenberg–Marquardt method by linearization of \( \{ \tilde{X}_{il}(\cdot) \} \) with respect to \( a^*_il, \theta^*_i \) and \( \beta^* \). We break the updating step into three parts corresponding to the three different sets of parameters. For each set of parameters, we first derive a first order Taylor expansion of the curves \( \{ \tilde{X}_{il}(\cdot) \} \) around their current values and then update them by a least squares fitting, while keeping the other two sets of parameters fixed at the current values. This process is repeated until convergence.

For notational convenience, denote the current estimates by \( a^*: = \{ a^*_il \} \), \( \theta^*: = \{ \theta^*_i \} \) and \( \beta^* \), and define the current residuals as \( \tilde{\varepsilon}_{ilj} := Y_{ilj} - \tilde{X}_{il}(t_{ilj}; a^*_il, \theta^*_i, \beta^*) \). For each \( i = 1, \ldots, n \), and \( l = 1, \ldots, N_i \), define the \( m_{il} \times 1 \) column vectors

\[
J_{il,a^*_il} := \left( \frac{\partial}{\partial a^*_il} \tilde{X}_{il}(t_{ilj}; a^*_il, \theta^*_i, \beta^*) \right)_{j=1}^{m_{il}}, \quad \tilde{\varepsilon}_{il} = (\tilde{\varepsilon}_{ilj})_{j=1}^{m_{il}}.
\]

For each \( i = 1, \ldots, n \), define the \( m_i \times 1 \) column vectors

\[
J_{i,\theta^*_i} := \left( \frac{\partial}{\partial \theta^*_i} \tilde{X}_{il}(t_{ilj}; a^*_il, \theta^*_i, \beta^*) \right)_{j=1,l=1}^{m_{il,N_i}}, \quad \tilde{\varepsilon}_{i} = (\tilde{\varepsilon}_{ilj})_{j=1,l=1}^{m_{il,N_i}},
\]

where \( m_i := \sum_{l=1}^{N_i} m_{il} \) is the total number of measurements for the \( i \)th subject. For each \( k = 1, \ldots, M \), define the \( m. \times 1 \) column vectors

\[
J_{\beta^*_k} := \left( \frac{\partial}{\partial \beta^*_k} \tilde{X}_{il}(t_{ilj}; a^*_il, \theta^*_i, \beta^*) \right)_{j=1,l=1,i=1}^{m_{il,N_i,n}}, \quad \tilde{\varepsilon} = (\tilde{\varepsilon}_{ilj})_{j=1,l=1,i=1}^{m_{il,N_i,n}},
\]

where \( m. := \sum_{i=1}^{n} \sum_{l=1}^{N_i} m_{il} \) is the total number of measurements. Note that, given \( a^*, \theta^* \) and \( \beta^* \), the trajectories \( \{ \tilde{X}_{il}(\cdot) \} \)'s and their gradients (as well as Hessians) can be easily evaluated on a fine grid by using numerical ODE solvers such as the fourth order Runge–Kutta method [see Paul, Peng and
Burman (2011) for details]. Since, given the trajectories, their gradients satisfy linear differential equations, the solution may also be obtained explicitly (see the Appendix). The equation for updating $\beta$, while keeping $a^*$ and $\theta^*$ fixed, is
\[ [J_{\beta^*}^T J_{\beta^*} + \lambda_3 \text{diag}(J_{\beta^*}^T J_{\beta^*}) + B](\beta - \beta^*) = J_{\beta^*}^T \tilde{\varepsilon} - B\beta^*, \]
where $J_{\beta^*} := (J_{\beta^*_1} : \cdots : J_{\beta^*_M})$ is an $m_\cdot \times M$ matrix. Here $\lambda_3$ is a sequence of positive constants decreasing to zero as the number of iterations increases. They are used to avoid possible singularities in the system of equations. The normal equation for updating $\theta_i$ is
\[ (J_{i,\theta^*_i}^T J_{i,\theta^*_i} + \lambda_2)(\theta_i - \theta_i^*) = J_{i,\theta^*_i}^T \tilde{\varepsilon}_i - \lambda_2 \theta_i^*, \quad i = 1, \ldots, n. \]
After updating $\theta_i$'s, we re-center the current estimates such that their mean is set to be zero. This also helps in stabilizing the algorithm. The equation for updating $a_{il}$, while keeping $\theta_i$ and $\beta$ fixed at $\theta_i^*, \beta^*$ is
\[ (J_{l,a^*_il}^T J_{l,a^*_il} + \lambda_1)(a_{il} - a_{il}^*) = J_{l,a^*_il}^T \tilde{\varepsilon}_{il} + \lambda_1 \alpha_{il}^*, \quad l = 1, \ldots, N_i, \ i = 1, \ldots, n, \]
where $\alpha^* := \sum_{i=1}^n \sum_{l=1}^{N_i} a_{il}^*/N$, $\alpha_{il}^* = \alpha^* - a_{il}^*$ with $N := \sum_{i=1}^n N_i$ being the total number of sample curves. Note that on convergence, $\hat{\alpha} := \alpha^*$ provides an estimate of $\alpha$. The initial estimates can be conveniently chosen. For example, $a_{il}^{ini} = Y_{il1}$ and $\theta_{il}^{ini} \equiv 0$.

This procedure is quite stable and robust to the initial parameter estimates. However, it converges slowly in the neighborhood of the minima of the objective function as it is a first order procedure. On the contrary, the Newton–Raphson algorithm has a fast convergence when starting from estimates that are already near the minima. Thus, in practice, one could first use the above approach (referred to as the Levenberg–Marquardt step hereafter) to obtain a reasonable estimate and then use the Newton–Raphson algorithm to expedite the search of the minima. The derivation of the Newton–Raphson algorithm is rather standard and thus is omitted. If the true gradient function $g$ has high complexity, and/or if either the $\theta_i$’s or the noise are highly variable, the Newton–Raphson algorithm may be unstable, particularly when the initial conditions $a = \{X_{il}(0)\}$ are also estimated. Under such situations, we recommend using a (relatively) large number of Levenberg–Marquardt steps, followed by a one-step Newton–Raphson update.

Note that the tuning parameter $\lambda_3$ plays a different role than the penalty parameters $\lambda_1$ and $\lambda_2$. The parameter $\lambda_3$ is used to stabilize the updates of $\beta$ and thereby facilitate convergence. Thus, it needs to decrease to zero with increasing iterations in order to avoid introducing bias in the estimate. In this paper, we simply set $\lambda_{3j} = \lambda_3^0/j$ for the $j$th iteration, for some pre-specified $\lambda_3^0 > 0$. On the other hand, $\lambda_1$ and $\lambda_2$ are parts of the loss function (3.3). Their main role is to control the bias-variance trade-off of the estimators,
even though they also help in regularizing the optimization procedure. From the likelihood viewpoint, \( \lambda_1 \) and \( \lambda_2 \) are determined by the variances \( \sigma^2_\varepsilon, \sigma^2_\alpha \) and \( \sigma^2_\theta \) through \( \lambda_1 = \sigma^2_\varepsilon / \sigma^2_\theta \) and \( \lambda_2 = \sigma^2_\varepsilon / \sigma^2_\alpha \). We can estimate these variances from the current residuals and current values of \( \alpha \) and \( \theta \). By assuming that \( m_{il} > 2 \) for each pair \((i, l)\),

\[
\widehat{\sigma}^2_\varepsilon = \frac{1}{m_i - N_i - M} \sum_{i=1}^{n} \sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} (\bar{\varepsilon}_{ij})^2,
\]

\[
\widehat{\sigma}^2_\alpha = \frac{1}{N_i - 1} \sum_{i=1}^{n} \sum_{l=1}^{N_i} (a^*_i - a^*)^2, \quad \widehat{\sigma}^2_\theta = \frac{1}{n - 1} \sum_{i=1}^{n} (\theta^*_i)^2.
\]

We can then plug in the estimates \( \widehat{\sigma}^2_\varepsilon, \widehat{\sigma}^2_\alpha \) and \( \widehat{\sigma}^2_\theta \) to get new values of \( \lambda_1 \) and \( \lambda_2 \) for the next iteration. Instead, if we take the penalized los s function viewpoint, we can simply treat \( \lambda_1 \) and \( \lambda_2 \) as fixed regularization parameters which can be chosen by model selection criteria (see Section 3.3). Henceforth, we refer to the method as adaptive if \( \lambda_1 \) and \( \lambda_2 \) are updated after each iteration, and as nonadaptive if they are kept fixed throughout the optimization.

### 3.2. Standard error of the estimates.

It is important to obtain the standard error of the estimated gradient function. Since it is typically not possible to obtain an estimate of the bias for a nonparametric procedure, we ignore the bias term and use the best projection of true \( g \) in the model space as the surrogate center (this is the standard practice in nonparametric literature). Thus, equivalently, we provide an estimate of the asymptotic variance of \( \hat{\beta} \). Based on the asymptotic analysis presented in Paul, Peng and Burman (2009), we derive the following estimate:

(3.5) \[
V(\hat{\beta}) := \widehat{E}[\hat{\beta} - \beta](\hat{\beta} - \beta)^T = \widehat{\sigma}^2_\varepsilon W_n,
\]

with \( W_n = (A_n + B - C_n^T (D_n + \lambda_2 I_n)^{-1} C_n)^{-1} \), where, \( I_n \) is the \( n \times n \) identity matrix, \( A_n = \mathcal{G}_{\beta\beta}(\theta, \beta), \quad C_n = \mathcal{G}_{\theta\beta}(\hat{\theta}, \beta), \quad D_n = \mathcal{G}_{\theta\theta}(\hat{\theta}, \beta) \); where

\[
\mathcal{G}_{\beta\beta}(\theta, \beta) := \sum_{i=1}^{n} \sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} \left( \frac{\partial X_{il}}{\partial \beta}(t_{ij}; a_{il}, \theta_i, \beta) \right) \left( \frac{\partial X_{il}}{\partial \beta}(t_{ij}; a_{il}, \theta_i, \beta) \right)^T;
\]

\[
\mathcal{G}_{\theta\beta}(\theta, \beta) := \sum_{i=1}^{n} \sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} \left( \frac{\partial X_{il}}{\partial \theta_i}(t_{ij}; a_{il}, \theta_i, \beta) \right) \left( \frac{\partial X_{il}}{\partial \theta_i}(t_{ij}; a_{il}, \theta_i, \beta) \right)^T.
\]

\( \mathcal{G}_{\theta\beta}(\theta, \beta) \) is the \( n \times M \) matrix with the \( i \)th row being

\[
\sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} \frac{\partial X_{il}}{\partial \theta_i}(t_{ij}; a_{il}, \theta_i, \beta) \left( \frac{\partial X_{il}}{\partial \theta_i}(t_{ij}; a_{il}, \theta_i, \beta) \right)^T, \quad i = 1, \ldots, n;
\]
and \( \mathcal{G}_{\theta\theta}(\theta,\beta) \) is the \( n \times n \) diagonal matrix with the \( i \)th diagonal entry

\[
\sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} \left( \frac{\partial X_{il}(t_{ilj}; a_{il}, \theta_i, \beta)}{\partial \theta_i} \right)^2,
\]

\( i = 1, \ldots, n. \)

Note that the matrices \( A_n, C_n \) and \( D_n \) are obtained as byproducts of the estimation procedure. An estimate of the standard error of \( \hat{g}(x) \) for \( x \) in the domain of \( \{ \phi_{k,M} \}_{k=1}^{M} \) is therefore given by

\[
\hat{SE}(\hat{g}(x)) = (\phi_M(x)^T V(\hat{\beta}) \phi_M(x))^{1/2},
\]

where \( \phi_M(x) := (\phi_{1,M}(x), \ldots, \phi_{M,M}(x))^T \) and \( V(\hat{\beta}) \) is as in (3.5). Note that, in the given asymptotic framework, we treat \( \theta_i \)'s as random effects and the initial conditions \( \{ a_{il} \} \) are assumed to be known. In deriving (3.5), we have ignored the correlation structure between \( \theta_i \) and the gradient of the objective function with respect to \( \theta_i \), which yields a slightly conservative (i.e., upwardly biased) estimate of the standard error. Obtaining the asymptotic standard error estimates when the initial conditions \( \{ a_{il} \} \) are estimated from the data is beyond the scope of this paper.

As an alternative way of estimating the standard error, one may also use bootstrap where we resample the sample trajectories corresponding to each subject, in order to retain the overall structure of the model. The corresponding bootstrap estimates, though simple to obtain, are computationally expensive and we do not pursue this in this paper.

3.3. Model selection. After specifying a scheme for the basis functions \( \{ \phi_{k,M}(\cdot) \} \), we still need to determine various model parameters such as the number of basis functions \( M \), the knot sequence, penalty parameters, etc. In the literature, AIC/BIC/AICc criteria have been proposed for model selection of parametric dynamical systems; see, for example, Miao et al. (2009). Here we propose an approximate leave-one-curve-out cross-validation score for model selection. Under the current context, the leave-one-curve-out CV score can be defined as

\[
CV := \sum_{i=1}^{n} \sum_{l=1}^{N_i} \sum_{j=1}^{m_{il}} \ell_{ij}^{cv}(\hat{a}_{il}^{(-il)}, \hat{\theta}_i^{(-il)}, \hat{\beta}^{(-il)}),
\]

where \( \hat{\theta}_i^{(-il)} \) and \( \hat{\beta}^{(-il)} \) are estimates of \( \theta_i \) and \( \beta \), respectively, based on the data after dropping the \( l \)th curve of the \( i \)th subject; and \( \hat{a}_{il}^{(-il)} \) is the minimizer of \( \sum_{j=1}^{m_{il}} \ell_{ij}^{cv}(a_{il}, \hat{\theta}_i^{(-il)}, \hat{\beta}^{(-il)}) \) with respect to \( a_{il} \); and \( \ell_{ij}^{cv}(a_{il}, \theta_i, \beta) := (Y_{ilj} - \tilde{X}_{il}(t_{ilj}; a_{il}, \theta_i, \beta))^2 \) is the prediction error loss. When the initial conditions \( a_{il} = X_{il}(0) \) are i.i.d. random variables and are known (and thus we
set $\hat{a}_il = a_il$, the leave-one-curve-out CV score gives an asymptotically unbiased estimator of the prediction error. Calculating CV score (3.7) is computationally very demanding because one needs to obtain $\hat{\theta}_i{-il}$ and $\hat{\beta}(-il)$ for every pair of $(i, l)$. Therefore, we propose to approximate $\hat{\theta}_i{-il}$ and $\hat{\beta}(-il)$ through a first order Taylor expansion around the estimates $\hat{\theta}_i$, $\hat{\beta}$ based on the full data. We then obtain an approximation of $\hat{a}_il$ by minimizing the corresponding criterion with the approximations of $\hat{\theta}_i{-il}$ and $\hat{\beta}(-il)$ imputed. Consequently, we derive an approximate CV score $\tilde{CV}$ by plugging these approximations in (3.7), which is computationally inexpensive since all the quantities involved in computing $\tilde{CV}$ are byproducts of the Newton–Raphson step used in model fitting. This approximation scheme is similar to the one taken in Peng and Paul (2009) under the context of functional principal component analysis, which itself is motivated by the work of Burman (1990). Detailed derivations are given in the Appendix.

4. Simulation. In this section we conduct a simulation study to demonstrate the effectiveness of the proposed estimation and model selection procedures. Since we apply our method to study the plant growth dynamics in Section 5, we consider a simulation setting that partly mimics that data set. In the simulation, the true gradient function $g$ is represented by $M^* = 4$ cubic $B$-spline basis functions with knots at $\{0.35, 0.6, 0.85, 1.1\}$ and basis coefficients $\beta = (0.1, 1.2, 1.6, 0.4)^T$. It is depicted by the solid curve in Figure 4. We consider two different settings for the number of measurements per curve: moderate case—$m_il$’s are independently and identically distributed as Uniform[5, 20]; sparse case—$m_il$’s are independently and identically distributed as Uniform[3, 8]. Measurement times $\{t_{ilj}\}$ are independently and identically distributed as Uniform[0, 1]. The scale parameters $\theta_i$’s are randomly sampled from $N(0, \sigma_\theta^2)$ with $\sigma_\theta = 0.1$; and the initial conditions $a_il$’s are randomly sampled from a $c_\alpha \chi^2_{k_\alpha}$ distribution (to ensure positivity as well as to study model robustness), with $c_\alpha, k_\alpha > 0$ chosen such that $\alpha = 0.25, \sigma_\alpha = 0.05$. Finally, the residuals $\varepsilon_{ilj}$’s are randomly sampled from $N(0, \sigma_\varepsilon^2)$ with $\sigma_\varepsilon = 0.01$. Throughout the simulation, we set the number of subjects $n = 10$ and the number of curves per subject $N_i = N = 20$. Observations $\{Y_{ilj}\}$ are generated using the model specified by equations (2.1), (2.2) and (3.1). For all settings, 50 independent data sets are used to evaluate the performance of the proposed procedure. The sample trajectories are evaluated using the 4th order Runge–Kutta method [as described in Paul, Peng and Burman (2011)] on an equally spaced grid with grid spacings $h = 0.0005$.

In the estimation procedure, we consider cubic $B$-spline basis functions with knots at $0.1 + j/M, j = 1, \ldots, M$, to model $g$, where $M$ varies from 2
Table 1
Convergence and model selection based on 50 independent replicates

| Model | a known | a estimated |
|-------|---------|-------------|
|       | 2 3 4 5 6 | 2 3 4 5 6 |
| moderate | Number converged | 50 50 50 50 50 | 50 7 50 50 46 |
|         | Number selected | 0 0 46 1 3 | 0 0 49 1 0 |
| sparse  | Number converged | 50 50 50 50 50 | 50 5 49 44 38 |
|         | Number selected | 0 0 45 0 5 | 1 0 47 1 1 |

to 6. Note that here \( M = 4 \) corresponds to the true gradient function. The Levenberg–Marquardt step is chosen to be nonadaptive, and the Newton–Raphson step is chosen to be adaptive (see Section 3.1 for the definition of adaptive and nonadaptive). We examine three different sets of initial values for \( \lambda_1 \) and \( \lambda_2 \): (i) \( \lambda_1 = \sigma^2 / \sigma^2_a = 0.04, \lambda_2 = \sigma^2 / \sigma^2_b = 0.01 \) (“true” values); (ii) \( \lambda_1 = 0.01, \lambda_2 = 0.0025 \) (“deflated” values); (iii) \( \lambda_1 = 0.16, \lambda_2 = 0.04 \) (“inflated” values). It turns out that the estimation and model selection procedures are quite robust to the initial choice of \( (\lambda_1, \lambda_2) \), thereby demonstrating the effectiveness of the adaptive method used in the Newton–Raphson step. Thus, in the following, we only report the results when the “true” values are used.

We also compare results when (i) the initial conditions \( a \) are known, and hence not estimated; and (ii) when \( a \) are estimated. As can be seen from Table 1, the estimation procedure converges well and the true model \( (M = 4) \) is selected most of the times for all the cases. Mean integrated squared error (MISE) and Mean squared prediction error (MSPE) and the corresponding standard deviations, SD(ISE) and SD(SPE), based on 50 independent data sets, are used for measuring the estimation accuracy of \( \hat{g} \) and \( \hat{\theta} \), respectively. Since the true model is selected most of the times, we only report results under the true model in Table 2. As can be seen from this table, when the initial conditions \( a \) are known, there is not much difference in

Table 2
Estimation accuracy under the true model

|       | MISE(\(\hat{g}\)) | SD(ISE) | MSPE(\(\hat{\theta}\)) | SD(SPE) |
|-------|------------------|--------|------------------------|--------|
| a known | moderate | 0.069  | 0.072  | 0.085  | 0.095  |
|         | sparse   | 0.072  | 0.073  | 0.085  | 0.095  |
| a estimated | moderate | 0.088  | 0.079  | 0.086  | 0.095  |
|         | sparse   | 0.146  | 0.129  | 0.087  | 0.094  |

*All the numbers are multiplied by 100.*
the performance between the moderate case and the sparse case. On the other hand, when $\mathbf{a}$ are estimated, the advantages of having more measurements become more prominent. We also conduct further simulation studies (results not reported in details here) to check the effect of increasing the noise level, as well as the dispersion of the initial conditions $\mathbf{a}$. When $\mathbf{a}$ are known, even with $\sigma_x = 0.05$, the convergence is almost unaffected, and in about 75% of the cases the true model ($M = 4$) is selected. Increasing $\sigma_a$ to 0.1 does affect convergence, especially for larger $M$. But under this setting, even with $\sigma_x = 0.05$, the true model converges in 90% of the cases and is selected to be the best in more than 75% of the cases. When $\mathbf{a}$ are estimated, the convergence deteriorates more obliviousy under increased noise levels.

In Figure 4 we have a graphical comparison of the fits when the initial conditions $\mathbf{a}$ are known versus when they are estimated in the sparse case. In the moderate case, there is very little visual difference under these two settings. We plot the true $g$ (solid curve), the pointwise mean of $\hat{g}$ (broken curve), and 2.5% and 97.5% pointwise quantiles (dotted curves) under the true model. These plots show that both fits are almost unbiased. Also, when $\mathbf{a}$ are estimated, there is greater variability in the estimated $g$ at smaller values of $x$, mainly due to a scarcity of data in that region. Indeed, the larger MISE of the estimator of $g$ when initial conditions are estimated mainly results from the larger MISE on the domain of $g$ where there is essentially no observed data. Due to the extrapolation effect, no method without using true initial conditions is expected to work well on such a domain, especially under a nonparametric setting. This point is illustrated in more detail later in this section (cf. Table 3), as well as in the supplementary material [Paul, Peng and Burman (2011), Section S3]. Overall, as can be seen from these tables and figures, the proposed estimation and model selection procedures perform effectively.

To evaluate the accuracy of the pointwise standard error estimator given in (3.6), in Figure 6 we plotted the average of the estimate (blue curve) over 50 independent data sets and the ±2 standard error bands of the estimates (broken red curves) based on the same 50 independent data sets under the true model ($M = 4$) when $\mathbf{a}$ is known. The pointwise standard errors are also computed empirically from the converged replicates (black curve) among the 50 simulation runs. We observe that, although being somewhat conservative, (3.6) gives a quite satisfactory estimate of the pointwise standard error of $\hat{g}$.

We also compare the performance of the proposed procedure with a two-stage approach. Following Chen and Wu (2008b), in the first stage, each individual trajectory $X_{il}(\cdot)$ and its derivative $X'_{il}(\cdot)$ are estimated by local linear and local quadratic smoothing, respectively. The bandwidths are chosen by cross-validation. In the second stage, two different methods for
Table 3
Comparison of estimation accuracy of two-stage estimators (either local quadratic smoothing or parametric regression using true model in the second stage) with hierarchical likelihood estimators (for the selected model, among models with $M = 2, \ldots, 6$ B-spline basis functions) under the sparse case.

Two-stage estimator

| Method in stage 2 | Bandwidths in stage I | Summary statistics | $x \in [-0.5, 0.2]$ | $x \in (0.2, 1]$ | $x \in (1, 1.5]$ |
|-------------------|-----------------------|---------------------|-------------------|----------------|----------------|
| Local quadratic smoothing | Optimal bandwidths | Mean(ISE($\hat{g}$)) | $3.8 \times 10^7$ | 20.177 | $7.3 \times 10^6$ |
| | | Median(ISE($\hat{g}$)) | $4.1 \times 10^5$ | $2.398$ | $1.8 \times 10^4$ |
| | | (SD(ISE($\hat{g}$))) | $(2.3 \times 10^8)$ | (330.146) | $(5.1 \times 10^7)$ |
| Regression (true model) | Optimal bandwidths | Mean(ISE($\hat{g}$)) | 27.592 | 28.492 | 0.063 |
| | | Median(ISE($\hat{g}$)) | $3.812$ | $2.094$ | $0.004$ |
| | | (SD(ISE($\hat{g}$))) | $(423.283)$ | $(565.281)$ | $(1.249)$ |

Hierarchical likelihood estimator

| Method | Summary statistics | $x \in [-0.5, 0.2]$ | $x \in (0.2, 1]$ | $x \in (1, 1.5]$ |
|--------|---------------------|-------------------|----------------|----------------|
| a known | Mean(ISE($\hat{g}$)) | 0.006 | 0.083 | 0.001 |
| | Median(ISE($\hat{g}$)) | $0.003$ | $0.041$ | $0.000$ |
| | (SD(ISE($\hat{g}$))) | $(0.009)$ | $(0.106)$ | $(0.002)$ |
| a estimated | Mean(ISE($\hat{g}$)) | 0.710 | 0.195 | 0.007 |
| | Median(ISE($\hat{g}$)) | $0.025$ | $0.054$ | $0.000$ |
| | (SD(ISE($\hat{g}$))) | $(4.751)$ | $(0.789)$ | $(0.048)$ |

estimating $g$ are considered with $\{\hat{X}_i(t)\}$ as response and $\{\hat{X}_i(t)\}$ as predictor: (i) a least squares regression fit of the basis coefficients using the true model; (ii) a local quadratic smoothing. A more detailed description of the two-stage approach and more simulation studies are given in the supplementary material [Paul, Peng and Burman (2011), Section S2].

In Table 3 we report the integrated squared errors of the two-stage estimators as well as those of the hierarchical likelihood estimators (under the model selected by $CV$) for the sparse case. While reporting the risk of the estimators, we divide the domain of $x$ into three regions: $[-0.5, 0.2]$, $(0.2, 1]$ and $(1, 1.5]$. In this simulation, even though the true gradient function $g$ has support effectively on $[-0.5, 1.5]$, the observed measurements $Y_{ilj}$’s are almost entirely confined in the region $(0.2, 1]$. Due to the extrapolation effect, methods without using the true initial conditions are expected to perform (relatively) poorly in the domains where there is no data. Thus, we divide the domain into different regions for more informative comparisons across methods. We also plot the pointwise mean and median and pointwise 95% bands around the mean for the two-stage estimators of $g$ in Figure 5. These
results show that the two two-stage estimators are highly biased and variable. The one using the true model in the second stage has better behavior in the regions where there is no data, compared to the fully nonparametric estimator. However, the level of bias and variability is much higher than the proposed estimator on all three regions. Another important observation is that, for the hierarchical likelihood estimator, the median of integrated squared errors over the data domain \((0, 1]\) is comparable for the cases when the initial condition \(a\) is known and when \(a\) is estimated.

To further compare these two approaches, we conduct another simulation study where all \(\theta_i\)'s are taken to be zero (equivalently, \(\sigma_\theta = 0\)), so that there is no subject-specific variability. For this simulation, we also consider a sampling design, referred to as “very dense,” in which the number of measurements per curve is Uniform\([60, 100]\) so that the first stage estimates of the two-stage methods are more accurate. The number of subjects is chosen to be \(n = 10\) and there is only one curve per subject (i.e., \(N_i \equiv 1\)). The results [reported in Table S5-5 in Paul, Peng and Burman (2011)] show that the proposed method again gives better estimates and it is much less biased (even when the initial conditions are estimated). The mean integrated squared error over the data domain \((0, 1]\) of the hierarchical likelihood estimator, when \(a\) is estimated, is much smaller than that of the two-stage method, even when the true model is used in the second stage. For a more detailed comparison of the two approaches, see Section S2 of Paul, Peng and Burman (2011). Moreover, we also do simulations when the true gradient function \(g\) is more complex and does not belong to the model space. The overall picture for the performance of the proposed estimation and model selection procedures, as well as the comparison with the two-stage methods, is consistent with the results presented here. See Section S3 of Paul, Peng and Burman (2011) for details.

Finally, we comment on the computational time and the rate of convergence of the proposed procedure. These depend on several factors, especially the model complexity and bias, noise level and criteria for convergence. Typically, the convergence is faster when \(a\) is treated as known, as opposed to when it is estimated from the data. For the simulation study presented here, under the true model \((M = 4)\), with \(a\) known, convergence is generally achieved in about 30 to 40 Levenberg–Marquardt steps and often in only 2 to 3 Newton–Raphson steps. The number of Levenberg–Marquardt steps required for convergence almost doubles when \(a\) is estimated. For biased models [including those presented in Section S3 of Paul, Peng and Burman (2011)], the convergence often takes more steps (up to 150 Levenberg–Marquardt steps and several Newton–Raphson steps). The computational times for the simulation study presented in this section are summarized in Table 4. These computations were carried out on a 64-bit Linux machine with Intel Core 2 Quad processors running at 3.2 GHz and with 8 GB RAM.
Table 4
Computational cost for the simulation study in Section 4. Reported quantities are the average time in seconds and standard deviations (within brackets) over 50 replicates (including the ones without convergence)

| Model (M) | 2    | 3    | 4    | 5    | 6    |
|-----------|------|------|------|------|------|
| moderate  | a known | 11.40 | 20.34 | 28.14 | 41.51 | 42.29 |
|           | (0.24) | (0.68) | (0.73) | (1.53) | (2.39) |
| a estimated | 21.22 | 89.20 | 44.23 | 56.34 | 69.89 |
|           | (1.18) | (18.38) | (4.54) | (9.33) | (23.05) |
| sparse    | a known | 11.50 | 20.35 | 28.25 | 41.53 | 42.57 |
|           | (0.33) | (0.63) | (0.80) | (1.58) | (3.05) |
| a estimated | 24.01 | 93.58 | 47.06 | 68.57 | 89.57 |
|           | (1.59) | (17.02) | (11.55) | (25.08) | (38.75) |

5. Application: Plant growth data. In this section we apply the proposed method to the plant growth data from Sacks, Silk and Burman (1997) described in the earlier sections. One goal of this study is to investigate the effect of water stress on growth displacement rate within the meristem of the primary root of maize seedlings. Note that, meristem is the tissue in plants consisting of undifferentiated cells and found in zones of the plant where growth can take place. The growth displacement rate is defined as the rate of displacement of a particle placed along the root and it should not be confused with “growth rate” which usually refers to the derivative of the growth trajectory with respect to time. For more details, see Sacks, Silk and Burman (1997). Growth displacement rate is important to infer the cell division rate—the local rate of formation of cells—that is not directly observable in a changing population of dividing cells. The growth displacement rate is also needed for understanding some important physiological processes such as biosynthesis [Silk and Erickson (1979); Schurr, Walter and Rascher (2006)]. Moreover, a useful growth descriptor called the “relative elemental growth rate” (REGR) can be calculated as the gradient of the growth displacement rate (with respect to distance), which shows quantitatively the magnitude of growth at each location within the organ.

The data consist of measurements on ten plants from a control group and nine plants from a treatment group where the plants are under water stress. The meristem region of the root, where the measurements are taken, is shown in Figure 1 (left panel). The primary roots had grown for approximately 18 hours in the normal and stressed conditions before the measurements were taken. The roots were marked at different places using a water-soluble marker and high-resolution photographs were used to measure the displacements of the marked places. The measurements were in terms of distances from the root cap junction (in millimeters) and were taken for each of these
marked places, hereafter markers, over an approximate 12-hour period while the plants were growing. The measurement process is shown schematically in the right panel of Figure 1. In Figure 3 the growth (displacement) trajectories of one plant with 28 markers in the control group and another plant with 26 markers in the treatment group are depicted. Note that measurement times are different for these two plants. Also, measurements were only taken in the meristem. Thus, whenever a marker grew outside of the meristem, its displacement would not be recorded at later times anymore. This, together with possible technical failures (in taking measurements), is the reason that in Figure 3 some growth trajectories were cut short. More sophisticated data acquisition techniques are described in Walter et al. (2002) and Basu et al. (1998), where the proposed method is also potentially applicable.

Many studies in plant science such as Silk (1994), Sacks, Silk and Burman (1997) and Fraser, Silk and Rost (1990) all suggest reasonably steady growth velocity across the meristem under both normal and water-stress conditions at an early developmental stage. Moreover, exploratory regression analysis based on empirical derivatives and empirical fits of the growth trajectories indicates that time is not a significant predictor and, thus, an autonomous model is reasonable. This also means that time zero does not play a role in terms of estimating the dynamical system and there is also no additional variation associated with individual markers. In addition, the form of the gradient function $g$ is not known to the plant scientists, only its behavior at root cap junction and at some later stage of growth are known [Silk (1994)]. Figure 2, the scatter plot of empirical derivatives versus empirical fits in the treatment group, indicates that there is an increase in the growth displacement rate starting from a zero rate at the root cap junction, followed by a nearly constant rate beyond a certain location. This means that growth stops beyond this point and the observed displacements are due to
growth in the part of the meristem closer to the root cap junction. Where
and how growth stops is of considerable scientific interest. These boundary
behaviors also imply that a linear ODE model is obviously not appropri-
ate. In addition, popular parametric models such as the Michaelis–Menten
type either do not satisfy the boundary constraints and/or have parameters without clear interpretations in the current context. Moreover, there is some controversy among plant scientists about the possible existence of a “growth bump” in the middle of the meristem. Taking all these features into consideration, the semiparametric model proposed in this paper is ap-

FIG. 5. True and fitted gradient function $g$ by two-stage approach for the sparse case. Top panel: the second stage uses local quadratic smoothing. Bottom panel: the second stage uses regression under the true model.
Fig. 6. Standard error estimates for the simulation study in Section 4. Pointwise standard error of $\hat{g}$ for the sparse case with initial conditions $a$ known. The true model (with $M = 4$ B-spline basis functions) is used in fitting. Solid black curve: pointwise standard error computed from 50 replicates. Solid blue curve: averaged pointwise standard error estimates from (3.6) (based on 50 replicates). Broken red curve: 2 standard deviations bands for the estimated pointwise standard error (based on 50 replicates).

appropriate for investigating the scientific questions associated with this study, in particular, comparing the baseline growth displacement rates between the treatment and control groups. Notice that, in order for the proposed estimation method to give an accurate estimate of the gradient function, we need only that the measurement on the state variable $x$ is dense in its domain, and that the measurement errors are independent across time. These are satisfied for the plant data since, even though each trajectory is recorded at a relatively small number of time points, there is a fairly large number of trajectories for each plant, corresponding to the different initial conditions. Note that, for each plant, the number of measurements is indeed the sum total of all the measurements for its different trajectories. Moreover, the proposed method combines information across different plants (subjects), which allows one to fit the model reasonably well even with relatively few measurements per subject.

Now consider the model described in Section 2. For the control group, we have the number of curves per subject $N_i$ varying in between 10 and 29; and for the water stress group, we have $12 \leq N_i \leq 31$. The observed growth displacement measurements $\{Y_{ij}: j = 1, \ldots, m_{il}, l = 1, \ldots, N_i\}_{i=1}^n$ are assumed to follow model (3.1), where $m_{il}$ is the number of measurements taken
for the \(i\)th plant at its \(l\)th marker, which varies between 2 and 17; and \(\{t_{ilj} : j = 1, \ldots, m_d\}\) are the times of measurements, which are in between [0, 12] hours. Altogether, for the control group there are 228 curves with a total of 1,486 measurements and for the treatment group there are 217 curves with 1,712 measurements in total. Note that the constraint at the root cap junction corresponds to \(g(0) = 0 = g'(0)\), which is imposed by simply omitting the constant and linear terms in the spline basis. The flatness of \(g\) at a (unknown) distance away from the root cap junction means that \(g'(x) = 0\) for \(x \geq A\) for some constant \(A > 0\). In order to impose this, as part of the objective function (3.3), we use

\[
\beta^T B \beta := \lambda_R \int_A^{2A} (g'(x))^2 \, dx = \lambda_R \beta^T \left[ \int_A^{2A} \phi'(x)(\phi'(x))^T \, dx \right] \beta,
\]

where \(\phi = (\phi_{1,M}, \ldots, \phi_{M,M})^T\) and \(\lambda_R\) is a large positive number quantifying the severity of this constraint; and \(A > 0\) determines where the growth displacement rate becomes a constant. \(A\) and \(\lambda_R\) are both adaptively determined by the model selection scheme discussed in Section 3.3. Moreover, since the initial conditions (marker positions) \(\{a_{il}\}\) are chosen according to some fixed experimental design (though measured with errors), it is not appropriate to shrink their estimates toward a fixed number. Hence, we set \(\lambda_1 = 0\) in the loss function (3.3).

Before fitting the proposed model, we first describe a simple regression-based method for getting a crude initial estimate of the function \(g(\cdot)\), as well as selecting a candidate set of knots. This involves (i) computing the re-scaled empirical derivatives \(e^{-\hat{\theta}^{(0)}_{ij}} \tilde{X}_{ilj}^r\) of the sample curves from the data, where the empirical derivatives are defined by taking divided differences: \(\tilde{X}_{ilj}^r := (Y_{il(j+1)} - Y_{ilj})/(t_{il(j+1)} - t_{ilj})\), and \(\hat{\theta}^{(0)}_{ij}\) is a preliminary estimate of \(\hat{\theta}_i\); and (ii) regressing the re-scaled empirical derivatives onto a set of basis functions evaluated at the corresponding sample averages: \(\tilde{X}_{ilj} := (Y_{il(j+1)} + Y_{ilj})/2\). In this paper we use the basis \(\{x^2, x^3, (x - x_k)^3\}_{k=1}^K\) with a pre-specified, dense set of knots \(\{x_k\}_{k=1}^K\). Then, a model selection procedure, like the stepwise regression, with either AIC or BIC criterion, can be used to select a set of candidate knots. In the following, we shall refer to this method as stepwise-regression. A similar method is employed by Sacks, Silk and Burman (1997). The resulting estimate of \(g\) and the selected knots can then act as a starting point for the proposed procedure. We expect this simple method to work reasonably well only when the number of measurements per curve is moderately large. Comparisons given later (Figure 10) demonstrate a clear superiority of the proposed method over this simple approach.

We fit the proposed model to the control group and the treatment group separately. For the control group, we first use the procedure described in
Section 3.1 with $g$ represented in cubic $B$-splines with $M$ (varying from 2 to 12) equally spaced knots. At this stage, we set $\beta_{ini} = 1_M$, $\theta_{ini} = 0_n$, $a_{ini} = (X_{il}(t_{il}): l = 1, \ldots, N_i)_{i=1}^n$. The criterion based on the approximate CV score [equation (A.7) in the Appendix] selects the model with $M = 9$ basis functions. This is not surprising since, when equally spaced knots are used, usually a large number of basis functions are needed to fit the data adequately. In order to get a more parsimonious model, we consider the stepwise-regression method to obtain a candidate set of knots. We use 28 equally spaced candidate knots on the interval $[0, 14]$ and use the fitted values $\{\hat{\theta}_{i}^{(0)}\}_{i=1}^{10}$ from the previous $B$-spline fit. The AIC criterion selects a model with 10 knots among these 28 candidate knots, plus the quadratic term. We then consider various submodels with knots chosen from this set of selected knots and fit them again using the proposed estimation procedure. The approximate CV scores for a number of different submodels are reported in Table 5. The parameters $A$ and $\lambda_R$ are also varied and selected by the approximate CV score. Based on the approximate CV score, the model with knot sequence $(3.0, 4.0, 6.0, 9.0, 9.5)$ and $(A, \lambda_R) = (9, 10^5)$ is selected. Also note that the model selected by stepwise-regression has a larger CV score than those of the models reported in Table 5. A similar procedure is applied to the treatment group. It turns out that the model with knot sequence $(3.0, 3.5, 7.5)$, which is also selected by stepwise-regression, has

Table 5

Model selection for real data. Control group: approximate CV scores for four submodels of the model selected by the AIC criterion in the stepwise-regression step. $M_1$: knots = $(3.0, 4.0, 5.0, 6.0, 9.0, 9.5)$; $M_2$: knots = $(3.0, 4.0, 5.5, 6.0, 9.0, 9.5)$; $M_3$: knots = $(3.0, 4.0, 6.0, 9.0, 9.5)$; $M_4$: knots = $(3.0, 4.5, 6.0, 9.0, 9.5)$. Treatment group: approximate CV scores for the model $M$: knots = $(3.0, 3.5, 7.5)$

| Model | $\lambda_R = 10^3$ | $\lambda_R = 10^5$ |
|-------|------------------|------------------|
|       | $A = 8.5$ | $A = 9$ | $A = 9.5$ | $A = 8.5$ | $A = 9$ | $A = 9.5$ |
| $M_1$ | 53.0924 | 53.0877 | 53.1299 | 54.6422 | 53.0803 | 53.1307 |
| $M_2$ | 53.0942 | 53.0898 | 53.1374 | 54.5190 | 53.0835 | 53.1375 |
| $M_3$ | 53.0300 | 53.0355 | 53.0729 | 53.8769 | $\text{53.0063}$ | 53.0729 |
| $M_4$ | 53.0420 | 53.0409 | 53.0723 | 54.0538 | 53.0198 | 53.0722 |

| $\lambda_R = 10^5$ | $\lambda_R = 10^5$ |
|------------------|------------------|
| $M$ | 64.9707 | 64.9817 |
| $A = 7$ | 64.9835 | 64.9817 |
| $A = 7.5$ | 64.9843 | 64.9817 |
| $A = 8$ | 65.5798 | 64.9817 |

*No convergence.
considerably smaller CV score compared to all other candidate models, and, hence, we only report the CV scores under this model in Table 5 with various choices of \((A, \lambda_R)\). It shows that \((A, \lambda_R) = (7, 10^3)\) has the smallest approximate CV score.

Figure 7 shows the estimated gradient functions \(\hat{g}\) under the selected models for the control and treatment groups, respectively. Apart from \(\hat{g}\), we also plot the estimated pointwise two-standard error bands using (3.6). The fact that the bands are generally nonoverlapping except for a small region clearly indicates that the baseline growth displacement rates for the control and treatment groups are different. The plot also shows that there is no growth bump for either group. In the part of the meristem closer to the root cap junction (distance within \(\sim 5.5\) mm), the growth displacement rate for the treatment group is higher than that for the control group. This is probably due to the greater cell elongation rate under water stress condition in this part of the meristem so that the root can reach deeper in the soil to get enough water. This is a known phenomenon in plant science. The growth displacement rate for the treatment group flattens out beyond a distance of about 6 mm from the root cap junction. The same phenomenon happens for the control group, however, at a further distance of about 8 mm from the root cap junction. Also, the final constant growth displacement rate of the control group is higher than that of the treatment group. This is due to the stunting effect of water stress on these plants, which results in an
Fig. 8. Fitted relative elemental growth rate (REGR) under the selected models for control and treatment groups, respectively. The REGR is computed by differentiating the estimated gradient function $g$.

earlier stop of growth and a slower cell division rate. Figure 8 shows the estimated relative elemental growth rates (i.e., $\hat{g}'$) for these two groups. Relative elemental growth rate (REGR) relates the magnitude of growth directly to the location along the meristem. For both groups, the growth is fastest in the middle part of the meristem ($\sim$3.8 mm for control group and $\sim$3.1 for treatment group), and then growth dies down pretty sharply and eventually stops. We observe a faster growth in the part of the meristem closer to the root cap junction for the water stress group and the growth dies down more quickly compared to the control group. The shape of the estimated $g$ may suggest that it might be modeled by a logistic function with suitably chosen location and scale parameters, even though the scientific meaning of these parameters is unclear and the boundary constraints are not satisfied exactly. As discussed earlier, there is insufficient knowledge from plant science to suggest a functional form beforehand. This signifies the major purpose and advantage of nonparametric modeling, which is to provide insights and to suggest candidate parametric models for further studies.

In order to check how our method performs in terms of estimating individual sample trajectories, we solved the differential equation model for each plant $i$ with fitted values of $X_{it}(0)$, $\theta_i$ and $g$. Figure 9 shows the fitted (under the selected model) and observed trajectories for three plants each from the control and the treatment groups. As can be seen from this figure, although there are subject-specific variabilities in the fits, the over-
all shapes of the trajectories are captured fairly well. Figure 10 shows the residual versus time plot for the treatment group. The plot for the control group is similar and thus is omitted. This plot shows that the proposed procedure based on minimizing the objective function (3.3) has much smaller and more evenly spread residuals (SSE = 64.50) than the fit by stepwise-regression (SSE = 147.57), indicating a clear benefit of the
more sophisticated approach. Overall, the estimation and model selection procedures give reasonable fits under both experimental conditions. Note that, for the first six hours, the residuals (right panel of Figure 10) show some time-dependent pattern, which is not present for later times. Since throughout the whole 12 hour period the residuals remain small compared to the scale of the measurements, the autonomous system approximation seems to be adequate at least for practical purposes. Nevertheless, modeling growth dynamics through nonautonomous systems may enable scientists to determine the stages of growth that are not steady across a region of the root. This aspect is discussed briefly in Section 6.

6. Discussion. The model and the fitting procedures presented in this paper are quite flexible and effective in terms of modeling autonomous dynamical systems nonparametrically when the data are from a number of subjects and when the underlying population level dynamics is of interest. When applying the proposed method to the plant growth data, we obtain results that are scientifically sensible. For the plant data, \( g \) is nonnegative and, thus, a modeling scheme imposing this constraint may be more advantageous. However, the markers are all placed at a certain distance from the root cap junction, where the growth displacement rate is already positive, and the total number of measurements per plant is moderately large. These mean that explicitly imposing nonnegativity is not crucial for the plant data, a fact also supported by the estimates which turn out to be nonnegative and the simulation results where the resulting estimators of \( g \) are always nonnegative for the moderate and/or “a known” cases. In general, if \( g \) is strictly positive (strictly negative) over the domain of interest, then we can model the logarithm of \( g \) (resp., \( -g \)) by basis representation.

The proposed approach is flexible in terms of incorporating various constraints on the dynamics and is able to capture features of the dynamical system which are not known to us a priori. It can also be extended to incorporate covariate effects, as well as to model nonautonomous systems which are currently under investigation. Even though in this paper we use the plant growth data as an illustration, the proposed framework is potentially useful to many other studies with similar types of data, where estimating the underlying dynamical system is of interest. For example, the data set collected as part of the Multicenter AIDS Cohort Study [Kaslow et al. (1987); Diggle et al. (2002)] can be used to study the dynamics of the CD4+ counts. Investigating the dynamics of CD4+ counts at a population level, while also taking into account individual effects, is of great importance to understand the progression of AIDS. This data set consists of 2,376 measurements of CD4+ cell counts against time since seroconversion (time when HIV becomes detectable) for 369 infected men enrolled in the study. In this data set, each patient is a subject and there is one sample curve associated with
each subject which reflects CD4+ counts over time. Moreover, each curve is only observed at a few time points and the set of measurement times is different across patients. The estimation procedure proposed in this paper can be adjusted appropriately to deal with such scenarios more effectively. Specifically, in order to deal with a large number of random effects, instead of the hierarchical likelihood approach, we can adopt a marginal maximum likelihood approach. These are topics of our ongoing research.

APPENDIX

Gradient of the sample trajectories. Note that $X_{il}(\cdot)$ satisfies

$$(A.1) \quad X_{il}(t) = a_{il} + \int_0^t e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}(X_{il}(s)) \, ds, \quad t \in [0,1].$$

Differentiating $(A.1)$ with respect to the parameters, we have

$$X_{a_{il}} (t):= \frac{\partial X_{il}(t)}{\partial a_{il}} = 1 + \int_0^t \frac{\partial X_{il}(s)}{\partial a_{il}} e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}(X_{il}(s)) \, ds,$$

$$X_{\theta_i} (t):= \frac{\partial X_{il}(t)}{\partial \theta_i} = \int_0^t \left[ \frac{\partial X_{il}(s)}{\partial \theta_i} e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}'(X_{il}(s)) + e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}(X_{il}(s)) \right] \, ds,$$

$$X_{\beta_k} (t):= \frac{\partial X_{il}(t)}{\partial \beta_k} = \int_0^t \left[ \frac{\partial X_{il}(s)}{\partial \beta_k} e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}'(X_{il}(s)) + e^{\theta_i} \phi_{k,M}(X_{il}(s)) \right] \, ds$$

for $i = 1, \ldots, n; \ l = 1, \ldots, N_i; \ r = 1, \ldots, M$. In other words, these functions satisfy the linear differential equations:

$$\frac{d}{dt} X_{a_{il}} (t) = X_{a_{il}} (t) e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}'(X_{i}(t)), \quad X_{a_{il}} (0) = 1,$$

$$\frac{d}{dt} X_{\theta_i} (t) = X_{\theta_i} (t) e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}'(X_{il}(t)) + e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}(X_{il}(t)), \quad X_{\theta_i} (0) = 0,$$

$$\frac{d}{dt} X_{\beta_k} (t) = X_{\beta_k} (t) e^{\theta_i} \sum_{k=1}^M \beta_k \phi_{k,M}'(X_{il}(t)) + e^{\theta_i} \phi_{k,M}(X_{il}(t)), \quad X_{\beta_k} (0) = 0.$$

If the $a_{il}$'s are positive and the function $g_\beta := \sum_{k=1}^M \beta_k \phi_{k,M}$ is positive on the domain of $a_{il}$'s, then the trajectories $X_{il}(t)$ are nondecreasing in $t$. In this
case, and more generally, whenever the solutions exist on the time interval [0, 1] and $g_\beta$ is continuously differentiable the gradients of the trajectories can be solved explicitly:

(A.2) \[ X^a_{il}(t) = \frac{g_\beta(X_{il}(t))}{g_\beta(X_{il}(0))}, \]

(A.3) \[ X^{\theta}_{il}(t) = e^{\theta t}g_\beta(X_{il}(t)), \]

(A.4) \[ X^{\beta}_{il}(t) = g_\beta(X_{il}(t)) \int_{X_{il}(0)}^{X_{il}(t)} \frac{\phi_{r,M}(x)}{(g_\beta(x))^2} dx. \]

We verify (A.4). Proofs (A.2) and (A.3) are similar. We can express

\[ X^{\beta_r}_{il}(t) = e^{\theta_i t} \int_0^t \phi_{r,M}(X_{il}(s)) \exp \left( e^{\theta_i} \int_s^t g_\beta(X_{il}(u)) du \right) ds \]

[using $X_{il}'(u) = e^{\theta_i g_\beta(X_{il}(u))}$]

\[ = e^{\theta_i t} \int_0^t \phi_{r,M}(X_{il}(s)) \exp \left( \log g_\beta(X_{il}(t)) - \log g_\beta(X_{il}(s)) \right) ds \]

\[ = g_\beta(X_{il}(t)) \int_{X_{il}(0)}^{X_{il}(t)} \frac{\phi_{r,M}(x)}{(g_\beta(x))^2} dx. \]

**Derivation of $\tilde{CV}$**. Observe that, when evaluated at the estimate $\hat{a}$, $\hat{\theta}$ and $\hat{\beta}$ based on the full data,

\[ \frac{\partial}{\partial \theta_i} \left( \sum_{l,j} \ell_{il}^{cv} \right) + 2\lambda_2 \theta_i = 0, \quad i = 1, \ldots, n; \]

(A.5)

\[ \frac{\partial}{\partial \beta} \left( \sum_{i,l,j} \ell_{ilj}^{cv} \right) + 2B \beta = 0. \]

Whereas, when evaluated at the drop $(i, l)$-estimates, $\hat{a}_{il}^{(-il)}, \hat{\theta}_i^{(-il)}, \hat{\beta}^{(-il)}$,

\[ \frac{\partial}{\partial \theta_i} \left( \sum_{l^* : l^* \neq l} \ell_{il^*}^{cv} \right) + 2\lambda_2 \theta_i = 0; \]

(A.6)

\[ \frac{\partial}{\partial \beta} \left( \sum_{i^*, l^*: (i^*, l^*) \neq (i, l)} \ell_{i^*l^*}^{cv} \right) + 2B \beta = 0. \]
Expanding the left-hand side of (A.6) around $\widehat{\beta}$ and $\widehat{\theta}$, and using (A.5), we obtain the following first order approximations:

$$
\widehat{\theta}^{(-il)}(t) \approx \tilde{\theta}^{(-il)}(t) := \hat{\theta}^{(-il)} + \left[ \sum_{i' = 1}^{N_i} \sum_{l' = 1}^{m_i} \frac{\partial^2 \ell^{cv}_{il}^{i'}}{\partial \theta_{i}^2} \cdot \partial^2 \ell^{cv}_{i'l'}^{i'} + 2\lambda_2 \right]^{-1} \left[ \sum_{j = 1}^{m_{il}} \left( \frac{\partial \ell^{cv}_{ilj}}{\partial \theta_{i}} \right) \right].
$$

$$
\widehat{\beta}^{(-il)}(t) \approx \tilde{\beta}^{(-il)}(t) := \hat{\beta}^{(-il)} + \left[ \sum_{i' = 1}^{N_i} \sum_{l' = 1}^{m_i} \sum_{j' = 1}^{m_{i'l'}} \frac{\partial^2 \ell^{cv}_{i'l'}^{i'}}{\partial \beta \partial \beta^T} + 2B \right]^{-1} \left[ \sum_{j = 1}^{m_{il}} \left( \frac{\partial \ell^{cv}_{ilj}}{\partial \beta} \right) \right].
$$

In the above, the gradients and Hessians of $\ell^{cv}_{ilj}$ are all evaluated at $(\tilde{\alpha}, \tilde{\theta}, \tilde{\beta})$, and, thus, they have already been computed on a fine grid in the course of obtaining these estimates. Hence, there is almost no additional computational cost to obtain these approximations. Now for $i = 1, \ldots, n; l = 1, \ldots, N_i$, define

$$
\tilde{a}^{(-il)} = \arg\min_a \sum_{j = 1}^{m_{il}} [Y_{ilj} - \tilde{X}_{ilj}(t_{ilj}; a, \tilde{\theta}^{(-il)}(t_{ilj}), \tilde{\beta}^{(-il)}(t_{ilj}))]^2 + \lambda_1 (a - \hat{\alpha})^2,
$$

where $\hat{\alpha}$ is the estimator of $\alpha$ obtained from the full data. Finally, the approximate leave-one-curve-out cross-validation score is

$$
(CV)_{i = 1}^{n} \sum_{l = 1}^{N_i} \sum_{j = 1}^{m_{il}} \ell^{cv}_{ilj}(\tilde{a}^{(-il)}(t_{ilj}), \tilde{\theta}^{(-il)}(t_{ilj}), \tilde{\beta}^{(-il)}(t_{ilj})).
$$

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SUPPLEMENTARY MATERIAL

Supplement to “Semiparametric modeling of autonomous nonlinear dynamical systems with application to plant growth” (DOI: 10.1214/11-AOAS459SUPP; .pdf). The supplementary materials provide additional details on the computational schemes. It also contains further simulation studies elucidating the performance of the proposed estimators under scenarios not covered in the main article.

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Department of Statistics
University of California
Davis, California 95616
USA
E-mail: debashis@wald.ucdavis.edu
jie@wald.ucdavis.edu
burman@wald.ucdavis.edu