PARAMETER ESTIMATION FOR DYNAMICAL SYSTEMS

by

Shelby R. Stanhope

B.S., Mathematics, Colorado State University, 2007
M.S., Mathematics, Colorado State University, 2010

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This dissertation was presented

by

Shelby R. Stanhope

It was defended on

June 20th 2016

and approved by

Dr. Jonathan Rubin, Professor, Department of Mathematics
Dr. David Swigon, Associate Professor, Department of Mathematics
Dr. Catalin Trenchea, Associate Professor, Department of Mathematics
Dr. Gilles Clermont, MD, Department of Critical Care Medicine UPMC

Dissertation Advisors: Dr. Jonathan Rubin, Professor, Department of Mathematics,
Dr. David Swigon, Associate Professor, Department of Mathematics
Parameter estimation is a vital component of model development. Making use of data, one aims to determine the parameters for which the model behaves in the same way as the system observations. In the setting of differential equation models, the available data often consists of time course measurements of the system. We begin by examining the parameter estimation problem in an idealized setting with complete knowledge of an entire single trajectory of data which is free from error. This addresses the question of uniqueness of the parameters, i.e. identifiability. We derive novel, interrelated conditions that are necessary and sufficient for identifiability for linear and linear-in-parameters dynamical systems. One result provides information about identifiability based solely on the geometric structure of an observed trajectory. Then, we look at identifiability from a discrete collection of data points along a trajectory. By considering data that are observed at equally spaced time intervals, we define a matrix whose Jordan structure determines the identifiability. We further extend the investigation to consider the case of uncertainty in the data. Our results establish regions in data space that give inverse problem solutions with particular properties, such as uniqueness or stability, and give bounds on the maximal allowable uncertainty in the data set that can be tolerated while maintaining these characteristics. Finally, the practical problem of parameter estimation from a collection of data for the system is addressed. In the setting of Bayesian parameter inference, we aim to improve the accuracy of the Metropolis-Hastings algorithm by introducing a new informative prior density called the Jacobian prior, which exploits knowledge of the fixed model structure. Two approaches are developed to systematically analyze the accuracy of the posterior density obtained using this prior.
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PREFACE

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I dedicate my dissertation in loving memory of my father Hurban Bolls. Thank you for always telling me that you were proud of me.
The motivation for this investigation is rooted in the modeling of biological systems with differential equations. In such an endeavor, one aims to model a physical phenomenon to better understand it, explain it, reproduce it, and possibly to make predictions about it. Insight can be gained through both quantitative and qualitative analysis of the model equations. The advantages of employing differential equation models for biological systems are discussed in [19]. In particular, the dynamics of how a system behaves over time, under varying conditions can be captured using such a modeling approach.

Development of a model is a process consisting of several stages. First, the model structure for the system of interest is designed based on physical laws and \textit{a priori} knowledge of how certain components in the system interact. Model complexity is a fundamental issue that must be confronted. The reductionist approach, in which small scale interactions are studied, or the systemic approach, of representing higher levels of organization, may be appropriate, depending on the modeling application [53, 60, 86]. Next, the identifiability of model parameters is analyzed and the model is appropriately adjusted. Data about the physical system is then collected using experimental studies, or already existing data is utilized, and values of model parameters are estimated from the data. Further analysis of the model is conducted and modifications to the model structure may be necessary. It is the estimation of the parameters on which this work is focused.

Differential equation models of physical systems often involve parameters whose values are not known and which may represent physical properties that cannot be directly measured. Other model parameters might arise from approximations or model reductions and they may have no direct biological interpretation [57]. Parameter estimation relies on the comparison of experimental observations of the modeled system with corresponding model output to
obtain values for these parameters. Due to this, some assert that the usefulness of the model is ultimately linked to the quality and abundance of the data [19]. Unfortunately, biologically realistic models are likely to be parameter rich and data poor [7]. More recently, experimental techniques such as microarray analysis, genome sequencing, high throughput flow cytometry and ELISA biochemical assays, have made a wealth of biological data available for certain applications [77].

Our particular interests stem from applications of mathematical modeling in medicine. In this setting, experimental data often consists of time course measurements of test subjects exposed to the same stimulus. In this work, we will refer to data collected from a single subject at multiple points in time as single trajectory data. This data is often sparsely available, for example, because an experiment is not repeatable due to the disease’s alteration of the subject’s immune system [36, 62]. It is also common that not all model variables can be measured. In this work, we will look at the problem of parameter estimation from single trajectory data and collections of single trajectory data. We do not focus on a particular application of mathematical modeling in biology, but instead try to address more broad aspects of parameter estimation for dynamical systems.

Our investigation of parameter estimation from single trajectory data follows a natural progression. We begin by examining an idealized setting with complete knowledge of an entire trajectory of data which is free from error, and then we consider discrete points along this trajectory. Next, we allow these discrete points to have uncertainty, and finally, we look at the case of collections of discrete single trajectory data.

In the setting of error free data, we begin in Chapter 2, by addressing the question of identifiability. A model is said to be identifiable if the parameter estimation problem has a unique solution given some amount of error free data. Identifiability analysis is an a priori study that should be conducted once a model structure has been fixed. It is important to establish identifiability prior to implementing computational techniques for parameter estimation to avoid misinterpretation of the numerical results.

There is a rich body of literature addressing identifiability for both linear and nonlinear systems of differential equations, [8, 9, 44, 58, 63, 89]. For linear systems of differential equations, the transfer function method utilizes Laplace transforms of the model equations
to derive a response function whose coefficients determine identifiability [8]. This method ultimately relies on the ability to solve simultaneous nonlinear equations. In another approach for linear systems, power series expansions of the trajectory as a function of the unknown parameters are analyzed to determine identifiability [89]. Alternatively, instead of trying to identify the parameter matrix of the linear system directly, the modal matrix approach aims to identify the eigenvalues and eigenvectors of the matrix [44]. These methods are only feasible for low dimensional systems as the analysis becomes difficult or impossible as dimensions increase. For nonlinear systems of differential equations, a differential algebra approach employs differential polynomials to define an input-output relation with rational coefficients which depend on the model parameters. These coefficients are used to define a map whose injectivity determines the identifiability. Computer algorithms have been introduced to perform the symbolic computations required for the differential algebra approach [9]. Another approach for nonlinear systems is the implicit function method. Here, derivatives of the system outputs with respect to time are taken and any unobservable system states are eliminated. Then the rank of a matrix whose entries consist of partial derivatives with respect to the parameters determines the identifiability of the system [90]. In this approach, higher order derivatives can become complicated and the singularity of the matrix can be difficult to determine [63]. Each of these treatments are highly reliant on computations that are only feasible in low dimensions or with the help of computer programs. Also, in the methods discussed above, the models typically include an input which serves as a control for the system. Identifiability is then addressed with full access to the set of all admissible controls. Our treatment is instead geared toward the case of single trajectory data. Without access to a set of controls, resulting in a variety of observations for the system, we have only a single initial condition, and thus one trajectory, from which to determine the parameters. This work complements this existing literature and approaches the identifiability problem in a uniquely different way, see Figure 1.

In Chapter 2, we investigate identifiability by asking whether the parameters of linear and linear-in-parameters dynamical systems can be uniquely determined from a single trajectory. We provide precise definitions of several forms of identifiability, and derive some novel, interrelated conditions that are necessary and sufficient for these forms of identifiability to
arise. We also show that the results have a direct extension to a class of nonlinear systems that are linear in parameters. One of our results provides information about identifiability based solely on the geometric structure of an observed trajectory, while other results relate to whether or not there exists an initial condition that yields identifiability of a fixed but unknown coefficient matrix and depend on its Jordan structure or other properties. Several examples are presented to illustrate identifiability for various systems and trajectories. In the later part of the chapter, we transition to looking at identifiability from a discrete collection of data points along a trajectory. By considering data that are observed at equally spaced time intervals, we form a special matrix whose Jordan structure determines the identifiability. Lastly, we show that the sensitivity of parameter estimation with discrete data depends on a condition number related to the data’s spatial confinement. The content of this chapter appeared as a publication in [76].

In Chapter 3, we continue our study of parameter estimation with discrete data from a single trajectory and extend the investigation to consider the case of uncertainty in the data. The problem of estimation of parameters of a dynamical system from discrete data can be formulated as the problem of inverting the map from the parameters of the system to points along a corresponding trajectory. Solving the inverse problem becomes even more
challenging in the presence of uncertainty in experimental measurements, as may arise due to measurement errors and fluctuations in system components. In Chapter 3, we focus on linear systems of differential equations and derive necessary and sufficient conditions for single trajectory data to yield a matrix of parameters with specific dynamical properties. To address the key issue of robustness, we go on to establish conditions that ensure that the desired properties of the solution to the inverse problem are maintained on an open neighborhood of the given data. We then build from these results to find bounds on the maximal uncertainty in the data that can be tolerated without a change in the nature of the inverse problem. In particular, both analytical and numerical estimates are derived for the largest allowable uncertainty in the data for which the qualitative features of the inverse problem solution, such as uniqueness or attractor properties, persist. We also derive the conditions and bounds for the non-existence of a real parameter matrix corresponding to the given data, which can be utilized in modeling practice to prescribe a level of uncertainty under which the linear model can be rejected as a representation of the data. One result establishes regions in data space on which uniqueness of the inverse problem solution is guaranteed. The invertibility of the solution map on these sets will play a vital role in the analysis of Chapter 4. Regions in data space on which the inverse problem maintains a unique solution can also be mapped back to define regions in parameter space on which the model is identifiable. The contents of this chapter have been submitted for publication and are currently undergoing review.

Our final study relates to the practical problem of parameter estimation given a collection of discrete single trajectory data. As we have seen, such a collection of data may arise from time series data which was measured experimentally from several individuals in a population. This situation provides the motivation for our study and presents several challenges in the parameter estimation problem. If the parameters of the model have biological significance, for example virus clearance rate or initial concentration of virus particles, it is natural for their values to vary from individual to individual within a population. With this perspective, the solution to the inverse problem is not a single parameter value, but instead, is a distribution of parameters representing variability over a population [77]. The objective is then to find the best approximation to the parameter density that describes this distribution. In a
Bayesian inference approach to parameter estimation, the parameter density is approximated by a posterior density, which makes use of the data that is available for the system. An introduction to Bayesian inference for parameter estimation can be found in [33, 74]. The posterior density is formed using a likelihood function, which quantifies the deviation of the model from the available data, and a prior density which incorporates any previous knowledge about the parameters. A sample of the posterior can be constructed using Markov chain Monte Carlo techniques, and in particular the Metropolis-Hastings algorithm. Several recent papers have used the Metropolis-Hastings algorithm for Bayesian parameter estimation in models of infectious disease and intrahost response to bacterial and viral infections, [59, 64, 65, 69].

Prescription of a likelihood function and prior density are necessary to implement Bayesian parameter estimation. The prior density reflects any information that is known about the parameters before data is considered, which may include bounds on the values of the parameters which are obtained from literature (e.g. biological experiments) or from qualitative analysis of the system (e.g. analysis of existence and stability of equilibria). A noninformative prior (also commonly referred to as an objective or flat prior) is used when no information about the parameter values is known. Alternatively, an informative prior can be used when characteristics of the parameters are known a priori. Unlike noninformative priors, informative prior is not dominated by the likelihood. The choice of a prior is a topic of intense debate among practitioners of Bayesian inference [10, 32, 68, 74]. In Chapter 4, we contribute to the conversation on prior density selection by introducing a new informative prior density called the Jacobian prior, which exploits knowledge of the fixed model structure, Figure 2. After presenting a theoretical derivation of the prior, we develop two approaches to systematically analyze the accuracy of the posterior density obtained by implementing the Metropolis-Hastings algorithm with the Jacobian prior. In first approach, we employ a linear system of differential equations and use key results of Chapter 3 to define the parameter density solution to the inverse problem as means for comparison with the posteriors obtained from the Metropolis-Hastings algorithm. The posterior obtained with the Jacobian prior is compared to the parameter density and to other posteriors that are computed using two other commonly employed priors. The second approach can be used with any system
of differential equations; we apply it again with the linear system and then with a nonlinear model. For the linear system of differential equations, there is a clear evidence that the Jacobian prior makes the posterior a better approximation of the parameter density. In the nonlinear case, the posterior is approximated more accurately with the Jacobian prior in some cases and equally well using all three of the priors in other cases.

Figure 2: Our contribution to the problem of prior density selection.
2.0 IDENTIFIABILITY OF LINEAR AND LINEAR-IN-PARAPRAMETERS DYNAMICAL SYSTEMS FROM A SINGLE TRAJECTORY

2.1 INTRODUCTION

Mathematical models of physical systems often include parameters for which numerical values are not known \textit{a priori} and cannot be directly measured. Parameter estimation relies on the comparison of experimental observations of the modeled system with corresponding model output to obtain values for these parameters. There are many computational techniques for parameter estimation that can be employed, most of which rely on minimization of the difference between model output and observed data. However, before numerical estimates of parameter values are pursued, it is important to address the question of whether the parameters of the model are identifiable; that is, does the parameter estimation problem have a unique solution, given access to some amount of error free data? If the model’s parameters are not identifiable from such idealized data, then numerical estimates of parameters from data may be misleading. Clearly, the answer to this question depends on the structure of the underlying model, the amount and type of the data given, and the precise definition of uniqueness.

Identifiability analysis of dynamical systems has been an area of intense study \cite{8, 15, 18, 63, 67, 89} and it is usually employed to aid model development, which ideally proceeds in the following order: first a model of a system of interest is designed; second, the identifiability of model parameters is analyzed and the model is appropriately adjusted; third, sufficient amount of data about the physical system is collected using experimental studies; and finally, values of model parameters are estimated from the data. In many biological studies, however, the order is often reversed, and modelers use data that were collected before any thought was
given to modeling of the system. The modeler then faces the challenging task of designing a model that represents and explains the existing data for a system that is no longer available or for which measurements cannot be repeated under the original conditions. In some cases, repeated collection of data from a single subject is impossible because it has been destroyed during the process of data collection. This happens frequently in studies of disease models in clinical settings, in which either the disease itself or the manipulations performed to assay the subject’s state may be fatal to the subject (laboratory animal). In immunological studies on mice, for example, each data point is an aggregate of data obtained from several different animals which are sacrificed during the process [82, 46]. In disease studies on larger mammals or humans, longitudinal data may be obtained for a single subject, but the experiment is not reproducible because the subject’s immune system has been altered by the disease [36, 62]. In this chapter we take this problem to the extreme and address the question of *identifiability of parameters of dynamical systems from a single observed trajectory*.

Linear models are a natural starting point for our study because they have a simple structure, but the identifiability question in this setting is nonetheless nontrivial because the solution to such a system depends nonlinearly on its parameters. This setting is also convenient because there is existing theory to build on and one can exploit invertible operators for numerical techniques for handling linear systems. Parameter identifiability for linear dynamical systems has been studied extensively in control theory and related areas [8, 30, 34, 48, 67, 70, 88, 89]. A time-invariant linear control model typically has the form

\[
\begin{align*}
\dot{x}(t) &= Ax(t) + Bu(t) \\
y(t) &= Cx(t)
\end{align*}
\]  

(2.1)

where \( A, B, C \) are parameter matrices, \( x(t) \) is a vector function that defines the state of the system at time \( t \), \( u(t) \) is the vector of input (control), and \( y(t) \) is the vector output. Zero initial conditions are typically chosen, but delta-function controls can be used to represent any desired initial condition. The impulse response function \( Y(s) = C(sI - A)^{-1}B \) fully describes the (Laplace transformed) solutions of the system. In this context the matrix parameters \( A, B, C \) are said to be identifiable if they can be determined from the set of output
observations $y$ obtained by varying the control $u$. It is known that even with all control functions available, the matrices $A, B, C$ are not all simultaneously identifiable, since the impulse response function is invariant under the transformation $(A, B, C) \rightarrow (T^{-1}AT, T^{-1}B, CT)$, where $T$ is an invertible matrix [8]. Kalman [48] devised a nomenclature in which the system (2.1) is transformed into a new set of variables that can be divided into four classes: (a) controllable but unobservable, (b) controllable and observable, (c) uncontrollable and unobservable, and (d) uncontrollable but observable. Bellman and Åström showed that if $C$ is square and has full rank (i.e., all states $x$ are observable) and the matrix $[B | AB | \cdots | A^{n-1}B]$ has full rank (i.e., the system is controllable), then both $A$ and $B$ are identifiable from the impulse response of the system when a full set of controls is available [8]. In special situations in which the matrices $A, B, C$ depend on a parameter, that parameter may be identifiable even when $C$ does not have full rank. It is not clear whether enough information can be obtained to identify the parameters from a smaller set of controls, perhaps even a single trajectory (i.e., single control), however, which is all that may be available in our motivating applications. Sontag has shown that when the model is identifiable, the parameters can be estimated even from information about a single variable extracted from a single trajectory, provided that enough data points have been measured: no more than $2r + 1$ data points are required to identify $r$ parameters in the case in which the dynamical system is real analytic [75]. Since the classical results discussed above imply that $C$ must be full rank for identifiability to be possible, we predominantly focus here on the case in which $C$ is full rank in (2.1), in which case, without loss of generality, we can in fact assume that $C = I$. However, we also briefly consider the issue of partial identifiability when $C$ is of lower rank.

As the next step, we examine nonlinear dynamical systems that depend linearly on parameters. Such systems can be written in the form

$$\dot{x}(t) = Af(x(t)) + u(t) - \mu(t)x(t) \quad (2.2)$$

where $A$ is a parameter matrix, $f$ is a known, locally Lipschitz continuous map (to insure existence and uniqueness of solutions), $u(t)$ is a time-dependent input and $\mu(t)$ is a decay control. Such systems commonly arise in differential equation models of chemical reaction networks that are derived from mass-action kinetics; in those cases, $A$ is the stoichiometric
matrix, \( f \) is the vector of reaction rate functions (each of which is a product of a reaction rate constant and a monomial in the components of \( x \)), \( u \) is the inflow into the reaction chamber, and \( \mu \) is the outflow \([41, 24, 25]\). Among models of type (2.2) one can also include the generalized Lotka-Volterra models that are commonly used in ecology or population dynamics \([21, 66, 43]\), in which cases \( A \) is usually a sparse matrix. The problem of identifying the model (2.2) consists of two separate tasks: (i) identification of the parameter matrix \( A \) and (ii) identification of the reaction rate functions \( f(x) \). Each of these problems has been studied extensively \([38, 84, 50]\). Several conditions for identifiability have also been derived. For example, Chen and Bastin \([13]\) found necessary and sufficient conditions for identifiability of \( A \) in the case when full response of the system to the controls \( u \) and \( \mu \) is available. Farina et al. \([23]\) addressed the problem of identifying the reaction rate constants in \( f(x) \) by expanding the system and linearizing about an equilibrium state and found that the Jacobian must be full rank and that knowledge of dynamical data for any time-dependent input to the system is essential. Craciun and Pantaea \([16]\) related the identifiability of chemical reaction network systems to the topology of the reaction network. Here we assume that the functions \( f(x) \) are known and focus on task (i).

We begin in Section 2.2 by presenting several definitions of identifiability and reviewing established theorems on identifiability for linear dynamical systems. In Section 2.3, we expand on these results to provide a complete rigorous characterization of identifiability from a single trajectory for linear dynamical systems. More specifically, we discuss several equivalent characterizations of the identifiability criterion, which ultimately yield an identifiability condition solely based on geometric properties of the known trajectory, namely whether or not the trajectory is confined to a proper subspace of the relevant phase space. This criterion provides practical utility, since it can be applied using what is known about the trajectory, without knowledge of the structure of the parameter matrix. Subsequently, we investigate the existence of a trajectory for which the parameter matrix is identifiable, obtaining a necessary and sufficient requirement for existence based on the properties of the coefficient matrix associated with the model. Several examples illustrate identifiability for various systems and initial conditions. In Section 2.4, we briefly discuss some implications of our confinement result for partial identifiability and for identifiability when not all model
variables are observable. In Section 2.5, we extend our results to a broader class of dynamics by deriving necessary and sufficient conditions for identifiability of a nonlinear system that is linear in parameters. The sufficient condition again has geometrical character and refers to the confinement of an image of the trajectory in the space of reaction rates. In Section 2.6, we discuss the problem of finding parameter values explicitly using discrete data and show that some linear models are identifiable from a complete trajectory but not from a finite set of data, and that the accuracy of parameter estimation is related to how significantly the available data deviates from confinement. The chapter concludes with a brief summary of results and possible future directions in Section 2.7.

### 2.2 DEFINITIONS AND PRELIMINARIES

We consider a model defined as a linear dynamical system in which data for all of the state variables is available (i.e. $C = I$) and the set of inputs (controls) consists of the set of initial conditions:

$$
\dot{x}(t) = Ax(t) \quad x(0) = b.
$$

In equation (2.3), $x(t) \in \mathbb{R}^n$ is the state of the system at time $t$, the system parameters are the entries of the coefficient matrix $A \in \mathbb{R}^{n \times n}$, and $b \in \mathbb{R}^n$ is the initial condition. For clarity of exposition we will refer to the entire matrix $A$ as the (matrix) parameter $A$. Analysis of linear systems of the form (2.3) is greatly simplified since we have an explicit formula for their solutions, which we generally refer to as trajectories:

$$
x(t; A, b) = e^{At}b = \sum_{j=0}^{\infty} \frac{A^j t^j}{j!} b.
$$

In the context of this work the term model identifiability is meant to represent the identifiability of the parameters of the model. In other words, we fix the model structure and ask whether the matrix parameter $A$ can be uniquely determined from error-free data consisting of a trajectory or some subset thereof. This question is strongly related to whether, independent of the data, two distinct parameters can lead to identical solutions. A term
parameter distinguishability has been used in this context in the literature [15, 34]. It is clear that these characteristics depend not only on the properties of the system, but also on the set of possible parameters being considered for comparison. The identifiability problem should therefore be reformulated as determining whether, on a particular subset of $\mathbb{R}^{n \times n}$, the map from parameter space to solution space is injective. The extreme cases are to allow parameters from the full parameter space $\mathbb{R}^{n \times n}$ or to restrict to a single point in the parameter space. In the latter case, identifiability is trivial since there are no other competing parameters to consider.

The initial condition $b$ determines the trajectory (2.4) of the system (2.3) uniquely for any given $A$. It needs to be taken into consideration, since we may or may not know or be able to select the initial values of the states represented in a model before running an experiment, and hence we may wish to consider identifiability from a given initial condition or from a larger set of initial conditions. We shall discuss here three progressively more constrained definitions of identifiability. In the first definition, we allow for an initial condition to be chosen to aid identifiability:

**Definition 1.** Model (2.3) is identifiable in $\Omega \subseteq \mathbb{R}^{n \times n}$ if and only if for all $A, B \in \Omega$, $A \neq B$, there exists $b \in \mathbb{R}^n$ such that $x(t; A, b) \neq x(t; B, b)$.

For identifiability in the sense of Definition 1, we immediately have the following result:

**Theorem 2.** Model (2.3) is identifiable in $\mathbb{R}^{n \times n}$.

**Proof.** Consider the negation of the statement: model (2.3) is not identifiable in $\Omega$ if there exist $A, B \in \Omega$, $A \neq B$, such that for all $b \in \mathbb{R}^n$, for all $t \in \mathbb{R}$, $x(t; A, b) = x(t; B, b)$. This negation cannot hold on a set $\Omega$ containing distinct elements $A$ and $B$, since if $x(t; A, b) = x(t; B, b)$ for all $b \in \mathbb{R}^n$, for all $t \in \mathbb{R}$, then differentiation of equation (2.4) and evaluation at $t = 0$ gives $Ab = Bb$ for all $b$. Applying this result to $n$ linearly independent choices of $b$ gives $AW = BW$ for the invertible matrix $W$ with the selected $b$ vectors as its columns.

---

1 The definitions above employ the term *identifiability* in the same sense in which *global identifiability* has been used in some literature (e.g., [80]) to distinguish this concept from that of *local identifiability*, which focuses on a small neighborhood of a given parameter. However, since $\Omega$ may be just a proper subset of the full parameter space, the use of the word global in this context could be misleading and thus we omit it.

2 The notation $x(t; A, b) \neq x(t; B, b)$ indicates that there exists at least one $t > 0$ such that $x(t; A, b) \neq x(t; B, b)$. Analogously, $x(t; A, b) = x(t; B, b)$ indicates that $x(t; A, b) = x(t; B, b)$ are identical for all $t$. 

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hence \( A = B \). As a consequence, we see that for any two distinct linear systems (2.3), there is an initial condition \( b \in \mathbb{R}^n \) that will distinguish the solutions.

Theorem 2 implies that if we are free to choose the initial condition, then any two linear models with distinct parameter matrices in \( \mathbb{R}^{n \times n} \) can be distinguished, because we can choose the initial condition \( b \) such that the corresponding trajectories will be distinct. In practice, however, control over the initial condition may not be available, and more restrictive definitions of identifiability are needed. Nonetheless, Definition 1 is important in a setting in which the matrix \( A \) itself is parametrized by an auxiliary parameter. For example, for \( \dot{x} = (p_1 + p_2)x \), the parameter sets \( (p_1, p_2) = (c_1, c_2) \) and \( (p_1, p_2) = (c_2, c_1) \) will result in the same trajectory, with no initial condition \( b \) that will distinguish them.

Identifiability from a single trajectory is addressed by the following definition:

**Definition 3.** Model (2.3) is identifiable in \( \Omega \) from \( b \in \mathbb{R}^n \) if and only if for all \( A, B \in \Omega \) with \( A \neq B \), it holds that \( x(\cdot; A, b) \neq x(\cdot; B, b) \).

Since in most practical applications with biological data, the initial condition is fixed and cannot be chosen at will, most of our results in Section 2.3 relate to identifiability in the sense of Definition 3. It is also of interest, however, to consider the problem of identifiability from any initial condition:

**Definition 4.** Model (2.3) is unconditionally identifiable in \( \Omega \) if and only if for all \( A, B \in \Omega \), \( A \neq B \) implies that for each nonzero \( b \in \mathbb{R}^n \), \( x(\cdot; A, b) \neq x(\cdot; B, b) \).

Sufficient conditions for unconditional identifiability will be revealed in the end of Section 2.3.

We will now show that necessary and sufficient conditions for identifiability of model (2.3) in the sense of Definition 3 follow from published results of Thowsen [80] and Bellman and Åström [8]. First, suppose that \( \Omega \) is an open set, in which case we have:

**Theorem 5.** For \( \Omega \subset \mathbb{R}^{n \times n} \) open and a fixed \( b \in \mathbb{R}^n \), model (2.3) is identifiable in \( \Omega \) from \( b \) if and only if \( \{b, Ab, ..., A^{n-1}b\} \) are linearly independent for all \( A \in \Omega \).

**Remark.** An alternative but equivalent formulation would replace the condition of linear independence with the condition that the matrix \([b|Ab|...|A^{n-1}b]\) has full rank, which would
bring the wording closer to that of Bellman and Åström.

We will obtain Theorem 5 from the following theorem, presented by Thowsen [80] (and simplified later by Gargash [28]), which we state without proof:

**Theorem 6.** Let \( \Omega \subset \mathbb{R}^{n \times n} \) be a set of matrices such that \( \Omega = \{ \sum_{j=1}^{p} A_j \alpha_j | \alpha \in \Theta \} \), where \( A_j \in \mathbb{R}^{n \times n} \) for all \( j \) and \( \Theta \) is an open subset of \( \mathbb{R}^p \). Fix \( b \in \mathbb{R}^n \) and let

\[
G(A) = \begin{bmatrix}
A_1b & \ldots & A_pb \\
A_1Ab & \ldots & A_pAb \\
\vdots & \ddots & \vdots \\
A_1A^{n-1}b & \ldots & A_pA^{n-1}b
\end{bmatrix}_{n^2 \times p}
\]

Model (2.3) is identifiable in \( \Omega \) from \( b \) if and only if \( \text{rank} G(A) = p \) for all \( A \in \Omega \).

**Proof.** (of Theorem 5) Consider the decomposition of \( A \) into elementary matrices, \( A = \sum_{i,j=1}^{n} E_{ij}a_{ij} \) where \( \{E_{ij}\} \) is the standard basis for \( n \times n \) matrices, and the set \( \Theta \) is equal to \( \mathbb{R}^{n^2} \). Under this decomposition, the matrix \( G(A) \) in Theorem 6 takes the special form

\[
\begin{bmatrix}
b^T & 0 & \ldots & 0 \\
0 & b^T & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & b^T
\end{bmatrix}
\]

\[
\begin{bmatrix}
(Ab)^T & 0 & \ldots & 0 \\
0 & (Ab)^T & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & (Ab)^T
\end{bmatrix}
\]

\[
\begin{bmatrix}
(A^{n-1}b)^T & 0 & \ldots & 0 \\
0 & (A^{n-1}b)^T & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & (A^{n-1}b)^T
\end{bmatrix}
\]
Let
\[
\tilde{G}(A) = \begin{bmatrix}
    b^T \\
    (Ab)^T \\
    \vdots \\
    (A^{n-1}b)^T
\end{bmatrix}_{n \times n}.
\]

With this definition, \( \text{rank } G(A) = n \cdot \text{rank } \tilde{G}(A) \). Hence, \( \text{rank } G(A) = n^2 \) if and only if \( \text{rank } \tilde{G}(A) = n \), which holds if and only if \( \{b, Ab, ..., A^{n-1}b\} \) are linearly independent.

Next, we note that the sufficiency of the condition of linear independence of \( \{b, Ab, ..., A^{n-1}b\} \) is not tied to the requirement that \( \Omega \) is an open set. This observation becomes obvious from a reformulation of a result of Bellman and Åström obtained for the control system (2.1) [8]:

**Theorem 7.** For \( \Omega \subset \mathbb{R}^{n \times n} \) and a fixed \( b \in \mathbb{R}^n \), if \( \{b, Ab, ..., A^{n-1}b\} \) are linearly independent for all \( A \in \Omega \), then model (2.3) is identifiable in \( \Omega \) from \( b \).

Let us define, for a fixed \( b \in \mathbb{R}^n \), the set \( \Omega_b = \{ A \in \mathbb{R}^{n \times n} : \{b, Ab, ..., A^{n-1}b\} \text{ are linearly independent} \} \). This set has a special significance for the identifiability of the model from \( b \) in view of the following corollary of Theorem 5:

**Corollary 8.** Let \( b \in \mathbb{R}^n \) be fixed. The set \( \Omega_b \) is the largest open set in which model (2.3) is identifiable from \( b \).

**Proof.** Clearly, by Theorem 5, any open set \( \Omega \subseteq \mathbb{R}^{n \times n} \) in which the model (2.3) is identifiable from \( b \) satisfies \( \Omega \subseteq \Omega_b \). That \( \Omega_b \) is an open set follows from Corollary 18, which is stated at the end of the next Section.

In view of Theorem 5, the sets \( \Omega_b \) can be employed to characterize sets \( \Omega \) in which the model (2.3) is unconditionally identifiable:

**Corollary 9.** Let \( \Omega \in \mathbb{R}^{n \times n} \) be open. Model (2.3) is unconditionally identifiable in \( \Omega \) if and only if \( \Omega \subseteq \bigcap_{b \in \mathbb{R}^n \setminus \{0\}} \Omega_b \).

Although the conditions stated in Theorems 5, 7, and Corollary 9 reveal the properties of sets in which the model is identifiable, their practical applicability is limited, because we cannot test whether the system obeys the condition \( A \in \Omega_b \) unless we know the matrix parameter \( A \). In the next section, we derive a more practical condition based on the properties
of a model trajectory. That is, let \( \gamma(A, b) = \{x(t; A, b) : t \in \mathbb{R} \} \subset \mathbb{R}^n \) denote the orbit of the model (2.3) corresponding to the trajectory with initial condition \( b \). The condition that we obtain specifies, for any given trajectory, whether or not we can identify the model from that trajectory. Specifically, we show that the model (2.3) is identifiable from a trajectory if and only if the orbit corresponding to that trajectory is not confined to a proper subspace of \( \mathbb{R}^n \).

2.3 IDENTIFIABILITY CONDITIONS BASED ON TRAJECTORY BEHAVIOR OR COEFFICIENT MATRIX PROPERTIES

2.3.1 Analysis

Now that we have established that the identifiability of the model from a fixed initial condition \( b \) is characterized by the linear independence of the set \( \{b, Ab, ..., A^{n-1}b\} \), we will discuss certain equivalent characterizations of that property that reveal its implications for the geometrical behavior of the corresponding orbit.

We will denote the space formed by linear combinations of the vectors \( \{b, Ab, ..., A^{n-1}b\} \) as \( K_n(A, b) = \text{span}\{b, Ab, ..., A^{n-1}b\} \). This space is the Krylov subspace generated by \( A \) and \( b \) [92] which is also called the \( A \)-cyclic subspace generated by \( b \) [40] and is the range of the controllability matrix \([b | Ab | ... | A^{n-1}b]\).

**Lemma 10.** \( \{b, Ab, ..., A^{n-1}b\} \) are linearly dependent, or equivalently \( \dim(K_n(A, b)) < n \), if and only if \( b \) is contained in an \( A \)-invariant proper subspace of \( \mathbb{R}^n \).

**Proof.** Recall that a space \( V \) is called \( A \)-invariant if for all \( v \in V, Av \in V \). For the forward direction, assume that \( \dim(K_n) < n \). Certainly, \( b \in K_n \). The result thus follows from showing that \( K_n \) is invariant under \( A \). To establish this invariance, let \( x \in K_n \). Then, \( x = c_0b + c_1Ab + ... + c_{n-1}A^{n-1}b \) and \( Ax = c_0Ab + c_1A^2b + ... + c_{n-1}A^n b \). By the Cayley-Hamilton Theorem, \( A^n \) can be written as a linear combination of \( I, A, A^2, ..., A^{n-1} \). Hence, \( Ax = d_0b + d_1Ab + ... + d_{n-1}A^{n-1}b \in K_n \), as desired.
The reverse direction follows immediately, as \( b \in V \) for an \( A \)-invariant set \( V \) with \( \dim(V) < n \) implies that, since \( K_n \subseteq V \), \( \dim(K_n) \leq \dim(V) < n \) as well. □

**Lemma 11.** Orbit \( \gamma(A,b) \) is confined to a proper subspace of \( \mathbb{R}^n \) if and only if \( b \) is contained in an \( A \)-invariant proper subspace of \( \mathbb{R}^n \).

**Proof.** For the backward direction, let \( V \subseteq \mathbb{R}^n \) be an \( A \)-invariant proper subspace of \( \mathbb{R}^n \). If \( b \in V \), then \( A^j b \in V \) for all \( j = 0,1,\ldots \). Hence, \( x(t;A,b) = e^{At}b = \sum_{j=1}^{\infty} \frac{t^j A^j b}{j!} \in V \) for all \( t \in \mathbb{R} \). So, \( \gamma(A,b) \subseteq V \).

For the forward direction, assume that \( \gamma(A,b) \) is confined to a proper subspace of \( \mathbb{R}^n \). Then there exists \( v \neq 0 \in \mathbb{R}^n \) such that \( v^T x(t;A,b) = 0 \) for all \( t \in \mathbb{R} \). Since \( x(t;A,b) = e^{At}b \), we have \( v^T e^{At}b = 0 \) for all \( t \in \mathbb{R} \) as well. Furthermore, differentiation gives \( v^T Ae^{At}b = 0, \ldots, v^T A^j e^{At}b = 0 \) for all \( t \in \mathbb{R} \), for \( j = 0,1,\ldots,n-1 \). In particular, for \( t = 0 \), \( v^T A^j b = 0 \) for \( j = 0,1,\ldots,n-1 \). Hence, \( v^T [b Ab\ldots A^{n-1}b] = 0 \). Since \( v \neq 0 \), \([b Ab\ldots A^{n-1}b]\) cannot have full rank, and thus Lemma 10 gives the desired result. □

In light of Theorem 5, Lemmas 10 and 11 yield the following corollary:

**Corollary 12.** For \( \Omega \subseteq \mathbb{R}^{n\times n} \) open, model (2.3) is identifiable in \( \Omega \) from \( b \) if and only if for all \( A \in \Omega \), \( \gamma(A,b) \) is not confined to a proper subspace of \( \mathbb{R}^n \).

Combining the above results we obtain one of the main results of this chapter, a concise relation between the uniqueness of the parameters of the model (2.3) and the geometric structure of its orbits:

**Theorem 13.** For the model (2.3) there exists no \( B \in \mathbb{R}^{n\times n} \) with \( A \neq B \) such that \( x(\cdot;A,b) = x(\cdot;B,b) \) if and only if the orbit \( \gamma(A,b) \) is not confined to a proper subspace of \( \mathbb{R}^n \).

**Proof.** Suppose that the orbit \( \gamma(A,b) \) of model (2.3) is not confined to a proper subspace of \( \mathbb{R}^n \), i.e., the parameter matrix \( A \) that supplied the orbit obeys \( A \in \Omega_b \). If \( B \in \Omega_b \) with \( A \neq B \), then the trajectories \( x(t;A,b) \) and \( x(t;B,b) \) are distinct as a result of the identifiability of the model (2.3) in \( \Omega_b \) from \( b \). If, on the other hand, \( B \notin \Omega_b \), then the set of vectors \( \{b,Bb,\ldots,B^{n-1}b\} \) is linearly dependent and hence Lemmas 10 and 11 imply that
the orbit \( \gamma(B, b) \) is confined to a proper subspace of \( \mathbb{R}^n \). Therefore \( x(t; B, b) \) is not equal to \( x(t; A, b) \).

Theorem 13 implies that the parameter matrix of model (2.3) is uniquely defined by any trajectory that has an orbit that is not confined to a proper subspace. It is a practical result that provides immediate information about the possibility of model identification from a single trajectory while relying solely on the geometrical description of the trajectory. Note that the full trajectory is needed to identify the matrix \( A \) since the orbit provides no information about the transit time.

With this relation of identifiability to trajectory behavior, we can also observe that identifiability is invariant under similarity transformation.

**Corollary 14.** Model (2.3) is identifiable in \( \Omega \in \mathbb{R}^{n \times n} \) from \( b \in \mathbb{R}^n \) if and only if model (2.3) is identifiable in \( \tilde{\Omega}_S = \{ C = SAS^{-1} : A \in \Omega \} \) from \( Sb \), for all \( S \in \mathbb{R}^{n \times n} \) invertible.

**Proof.** For the forward direction, assume that model (2.3) is identifiable in \( \Omega \) from \( b \) and let \( A \in \Omega \). Let \( S \) be an invertible \( n \times n \) matrix and define \( C = SAS^{-1} \). Assume \( C \) and \( D \) yield the same trajectory for the initial condition \( Sb \). Since \( C \) and \( D \) yield the same trajectory, \( e^{Ct}Sb = e^{Dt}Sb \) and hence \( e^{S^{-1}CSt}b = e^{S^{-1}DSt}b \) and \( e^{At}b = e^{S^{-1}DSt}b \), for all \( t \in \mathbb{R} \). Since \( A \) is identifiable from \( b \), \( A = S^{-1}DS \). So, \( SAS^{-1} = D \), which yields \( C = D \). Hence, model (2.3) is identifiable in \( \Omega \) from \( Sb \), for all \( S \in \mathbb{R}^{n \times n} \) invertible. The reverse direction follows similarly.

Having related identifiability on \( \Omega \) to the linear independence of \( \{ b, Ab, \ldots, A^{n-1}b \} \) for \( A \in \Omega \), it is natural to ask whether for every parameter matrix \( A \) the set \( \Omega_b \) is non-empty, i.e., whether for every \( A \) there is an initial condition \( b \) such that such that \( \{ b, Ab, \ldots, A^{n-1}b \} \) are linearly independent. We will now show that such a \( b \) need not necessarily exist and hence there are models (2.3) that cannot be identified from any single trajectory.

The following result determines under what conditions on the structure of \( A \) the set \( \Omega_b \) is non-empty.

**Theorem 15.** There exists \( b \) such that \( \{ b, Ab, \ldots, A^{n-1}b \} \) are linearly independent if and only if \( A \) has only one Jordan block for each of its eigenvalues.
Proof. The result follows from Cyclic Decomposition Theorem for square matrices. In particular, a vector \( b \) such that \( \{b, Ab, \ldots, A^{n-1}b\} \) are linearly independent is called a cyclic vector for \( A \). A corollary of the Cyclic Decomposition Theorem ([40] p. 237) states that \( A \) has a cyclic vector if and only if the characteristic and minimal polynomials for \( A \) are identical, which holds if and only if the matrix \( A \) has only one Jordan block for each of its eigenvalues.

This condition results from the fact that a nilpotent matrix \((J - \lambda I)\), where \( J \) is an \( k \times k \) elementary Jordan matrix with eigenvalue \( \lambda \), has minimal polynomial of the form \( x^k \), and furthermore the minimal polynomial of \( A \) is of order \( m = \sum_{i=0}^{s} k_i \) where \( k_i \) is the size of the largest Jordan block corresponding to the eigenvalue \( \lambda_i \) of \( A \), with the sum being taken over all distinct eigenvalues of \( A \). Now, the property that \( A \) has only one Jordan block for each of its eigenvalues holds if and only if \( m = \sum_{i=0}^{s} k_i = n \), which is equivalent to the characteristic and minimal polynomials for \( A \) being identical. \( \square \)

An additional Proposition elucidates a certain structure for \( b \) that is useful for attaining identifiability.

**Proposition 16.** For a \( k \times k \) elementary Jordan matrix \( J \) with eigenvalue \( \lambda \) and vector \( b \in \mathbb{R}^k \) with \( b_k \neq 0 \), \((J - \lambda I)^m b = 0\) if and only if \( m \geq k \).

**Proof.** The backward direction is immediate since for \( m \geq k \), \((J - \lambda I)^k = 0\). The forward direction follows because for \( p < k \), \((J - \lambda I)^p b = [b_{p+1}, \ldots, b_k, 0, \ldots, 0]^T \neq 0 \) whenever \( b_k \neq 0 \). \( \square \)

Theorem 15 establishes the existence of \( b \) from which model (2.3) is identifiable under the linear independence condition and Proposition 16 specifies a structure for such \( b \). In particular, for such \( b \), the component corresponding to each \( k_i \times k_i \) Jordan block \( J_i \) of \( A \) must not be annihilated by \((J_i - \lambda_i I)^{k_i-1}\). For diagonalizable \( A \), this result reduces to the requirement that such \( b \) has a nonzero component in each eigenspace of \( A \) (see Lee [56]). This property, sometimes referred to as persistent excitability for systems in control theory, has been discussed by [5, 58]. A result closely related to Theorem 15 can be found in [52]. More interesting examples arise for matrices that do not have distinct real eigenvalues and are discussed in examples below.
Another helpful characterization of the linear independence of \{b, Ab, ..., A^{n-1}b\} can be stated in terms of left eigenvectors of \(A\), using the well known PBH controllability test for linear systems [47]:

**Theorem 17.** The vectors \{b, Ab, ..., A^{n-1}b\} are linearly independent if and only if there is no left eigenvector of \(A\) orthogonal to \(b\).

In our case we employ this theorem for the following result:

**Corollary 18.** For \(\Omega \subset \mathbb{R}^{n \times n}\) open, model (2.3) is identifiable in \(\Omega\) from \(b\) if and only if there is no left eigenvector of \(A\) orthogonal to \(b\) for any \(A \in \Omega\).

The Corollary 18 provides a new criterion for identifiability of the model (2.3) and can be used to characterize the set \(\Omega_b\) from Section 2: \(\Omega_b = \{A \in \mathbb{R}^{n \times n} : w^T A = \lambda w \Rightarrow w^T b \neq 0\}\) defines the largest open set in which the model is identifiable from \(b\). Likewise,

\[
\bigcap_{b \in \mathbb{R}^n \setminus \{0\}} \Omega_b = \{A \in \mathbb{R}^{n \times n} : w^T A = \lambda w \Rightarrow w^T b \neq 0 \text{ for all } b \in \mathbb{R}^n \setminus \{0\}\}
\]

defines the largest set \(\Omega\) in which the model is unconditionally identifiable.

For matrices of even dimensions we can use this representation to define sets \(\Omega\) in which model (2.3) is unconditionally identifiable. For example, let \(\Omega_c = \{A \in \mathbb{R}^{2 \times 2} : A\) has a complex pair of eigenvalues\}. Any left eigenvector \(w\) of a matrix \(A\) in this set has nonzero imaginary part and hence \(w\) is not orthogonal to any vector \(b \in \mathbb{R}^2\). By Corollary 18, \(\{b, Ab\}\) are linearly independent (over \(\mathbb{R}\)), hence the model is unconditionally identifiable in \(\Omega_c\) by Corollary 9.

Interestingly, matrices with odd dimensions always have a real-valued left eigenvector \(w\) and our results imply that there is no set \(\Omega \subseteq \mathbb{R}^{(2n+1) \times (2n+1)}\) in which model (2.3) is unconditionally identifiable, since in that case one can find \(b \in \mathbb{R}^{2n+1}\) such that \(w^T b = 0\).

### 2.3.2 Examples

We will now discuss several examples of identifiability in \(\Omega\) from \(b\) for various sets \(\Omega\) in parameter space and initial conditions \(b\). We start by discussing certain special choices
of $\Omega$ selected based on the properties we have established as important for identifiability. Subsequently, we consider how the behaviors of trajectories of (2.3) for some particular matrices relate to our results.

As we have discussed, $\Omega_b = \{ A \in \mathbb{R}^{n \times n} : \{ b, Ab, \ldots, A^{n-1}b \} \text{ are linearly independent} \} = \{ A \in \mathbb{R}^{n \times n} : w^T A = \lambda w \Rightarrow w^T b \neq 0 \}$ is the largest open set in which the model is identifiable from $b$. Thus, identifiability from $b$ holds in any subset of $\Omega_b$, regardless of whether that set is open. Also, we can define a larger (non-open) set $\Omega$ in which we have identifiability from $b$ by extending $\Omega_b$ in trivial ways, such as by combining $\Omega_b$ with a single matrix $A_0 \in \Omega_b^c$. For $\Omega = \Omega_b \cup \{ A_0 \}$, the model will be identifiable in $\Omega$ from $b$ because the orbit $\gamma(A_0, b)$ is confined to a proper subspace of $\mathbb{R}^n$ and will not coincide with any orbit for a matrix in $\Omega_b$ (which cannot be so confined). This type of trivial extension could be continued with a sequence of matrices that have confined trajectories for the initial condition $b$, but with no two solutions that are the same. On the other hand, define $\Omega = \mathbb{R}^{n \times n}\setminus\Omega_b$, which is not open, such that Theorem 5 and Corollary 12 do not apply. It is clear that any two distinct matrices $A, B$ with the same eigenvalues that share $b$ as an eigenvector for the same eigenvalue will yield the same solution, $x(t; A, b) = x(t; B, b)$ for all $t \in \mathbb{R}$, and such matrices can be found in $\Omega$. Hence, by Definition 3, model (2.3) is not identifiable in $\Omega$ from $b$. This argument shows that we do not have identifiability from $b$ on the complement of $\Omega_b$.

Next, define $\Omega_J = \{ A \in \mathbb{R}^{n \times n} : A \text{ has more than one Jordan block for some eigenvalue} \}$. From Theorem 15 we know that for any $A \in \Omega_J$, $\{ b, Ab, \ldots, A^{n-1}b \}$ are linearly dependent for any $b \in \mathbb{R}^n$. $\Omega_J$ is a set of measure zero and is not open in $\mathbb{R}^{n \times n}$, however, so Theorem 5 does not apply to the identifiability of $\Omega_J$. We can again appeal directly to Definition 3 to show that the model is not identifiable in $\Omega_J$ from $b$ for any choice of $b \in \mathbb{R}^n$. For example, in $\mathbb{R}^{3 \times 3}$, fix $b = [1, 1, 1]^T$. Let

$$A = \begin{bmatrix} -2 & 0 & 0 \\ 0 & -2 & 0 \\ 2 & -1 & -3 \end{bmatrix} \quad \text{and} \quad B = \begin{bmatrix} -2 & 0 & 0 \\ 1 & -2.5 & -0.5 \\ 1 & -0.5 & -2.5 \end{bmatrix}.$$
$A$ and $B$ both have the Jordan matrix

$$D = \begin{bmatrix} -2 & 0 & 0 \\ 0 & -2 & 0 \\ 0 & 0 & -3 \end{bmatrix}$$

which has two Jordan blocks for $\lambda = -2$, so $A, B \in \Omega_J$. $A$ and $B$ also share $b$ as an eigenvector for $\lambda = -2$. So, $x(t; A, b) = x(t; B, b)$ for all $t \in \mathbb{R}$, but $A \neq B$, hence by Definition 3, model (2.3) is not identifiable in $\Omega_J$ from $b$. An analogous pair of matrices that violate identifiability can be obtained for any $b \in \mathbb{R}^n$ by simply choosing two matrices with the same Jordan form, having more than one Jordan block for some eigenvalue and sharing $b$ as an eigenvector for that eigenvalue. In fact, $\Omega_J \cap \Omega_b = \emptyset$. So it is straightforward to construct sets $\Omega \subseteq \mathbb{R}^{n \times n}$ on which the model is not identifiable from $b$, using elements of $\Omega_J$.

To illustrate the application of Theorem 13, we will now discuss several examples of $2 \times 2$ and $3 \times 3$ matrices. For each, we consider under what conditions they have confined trajectories and correspondingly, for what initial conditions $b$ we have $A \in \Omega_b$.

Figure 3 shows the phase plane for each of four matrices with trajectories plotted for a few different initial conditions. Matrix

$$A_a = \begin{bmatrix} -2 & 1 \\ 1 & -2 \end{bmatrix}$$

has distinct eigenvalues. For any initial condition $b$ lying on an eigenvector, the corresponding orbit will be confined to a proper subspace of $\mathbb{R}^2$. These are the only initial conditions that lead to confined trajectories, so $A_a \in \Omega_b$ for all $b$ not on an eigenvector. This observation is consistent with the requirements on the structure of $b$ for identifiability, as given in Proposition 16 and the associated discussion. That is, if we consider $A_a$ and $b$ written in the basis in which $A_a$ is in Jordan form (diagonalized), then $b$ would have a zero component if and only if it were an eigenvector.

Matrix

$$A_b = \begin{bmatrix} -5/2 & 1/2 \\ -1/2 & -3/2 \end{bmatrix}$$
Figure 3: Phase portraits for system (2.3) generated by four matrices with different eigenvalue structures. In this and subsequent figures, red curves are trajectories from which the corresponding model is not identifiable while blue curves are trajectories that yield identifiability. Matrices $A_a, A_b, A_c, A_d$, as given in the text, were used to generate panels (a), (b), (c), and (d) respectively.

has a repeated eigenvalue of $-2$ and is not diagonalizable. Since it only has one Jordan block, the requirement on $b$ for the orbit to not be confined is that in the basis in which $A_b$ is given in Jordan form, $b$ must have a nonzero component in the last entry. Equivalently, this requirement means that $b$ cannot lie on the genuine eigenvector $[1, 1]^T$. From the phase plane
it is easy to see that the orbit arising from any initial condition lying on this eigenvector will be confined to a proper subspace of $\mathbb{R}^2$. So, $A_b \in \Omega_b$ for all $b$ not on the genuine eigenvector.

Matrix

$$A_c = \begin{bmatrix} -2 & -3 \\ 3 & -2 \end{bmatrix}$$

has a complex conjugate pair of eigenvalues. In this circumstance, $\mathbb{R}^2$ has no nontrivial $A_c$-invariant proper subspaces. Therefore, the orbit from any nonzero initial condition is not confined to a proper subspace of $\mathbb{R}^2$. This conclusion is clear from the phase plane. Hence, $A_c \in \Omega_b$ for all $b \in \mathbb{R}^2 \setminus \{0\}$.

Finally, matrix

$$A_d = \begin{bmatrix} -2 & 0 \\ 0 & -2 \end{bmatrix}$$

represents the case of a repeated Jordan block for $\lambda = -2$. This is a star shaped system in which every initial condition lies on an eigenvector and hence every orbit is confined to a proper subspace. Thus, in this case, there exists no $b$ such that $A_d \in \Omega_b$.

![Figure 4: Phase space structures for matrix $A_e$.](image)
Figure 4 corresponds to the matrix

\[
A_e = \begin{bmatrix}
-1 & -3 & 2 \\
0 & -4 & 2 \\
0 & 0 & -2 \\
\end{bmatrix},
\]

which has three distinct real eigenvalues. The corresponding eigenvectors are plotted in black and the planes represent the two-dimensional invariant subspaces spanned by pairs of eigenvectors. Any initial condition lying in one of these planes will have a zero component (in the basis in which \(A\) is diagonalized) and the corresponding orbit will be confined to that plane. Any initial condition outside of these planes will have a orbit that is not confined to a proper subspace. Hence, \(A_e \in \Omega_b\) for all \(b\) not lying in one of the planes. This example is the three-dimensional analogue of \(A_a\) but is more interesting because for \(A_e\) to be in \(\Omega_b\), not only must \(b\) not lie on an eigenvector of \(A_e\), but it also may not land on any two-dimensional plane spanned by two eigenvectors of \(A_e\).

![Figure 5: Phase space structures for matrix \(A_f\).](image)

Figure 5 was generated from

\[
A_f = \begin{bmatrix}
-0.2 & -1 & 0 \\
1 & -0.2 & 0 \\
0 & 0 & -0.3 \\
\end{bmatrix},
\]
which has a complex conjugate pair of eigenvalues and one real eigenvalue: \( \lambda_{1,2} = -0.2 \pm i \) and \( \lambda_3 = -0.3 \). Any orbit with an initial condition on the \( xy \) plane will stay confined to that plane. This is the only proper \( A_f \)-invariant subspace of \( \mathbb{R}^3 \). It is clear from the phase plane that any orbit from an initial condition outside of this plane will not be confined to a proper subspace. Hence, \( A_f \in \Omega_b \) for all \( b \) not lying in the \( xy \) plane.

Figure 6 corresponds to

\[
A_g = \begin{bmatrix}
-1 & 1 & 0 \\
0 & -1 & 0 \\
0 & 0 & -1
\end{bmatrix},
\]

which has two Jordan blocks for the eigenvalue \( \lambda = -1 \). Because of the repeated block, the orbit from any initial condition is confined to a proper subspace of \( \mathbb{R}^3 \). Hence, there is no \( b \in \mathbb{R}^3 \) such that \( A_g \in \Omega_b \). This conclusion can also be verified from Theorem 17. The left eigenvectors of \( A_g \) are: \( x_1 = [0, 1, 0] \), \( x_2 = [0, 0, 1] \). For an arbitrary \( b = [b_1, b_2, b_3]^T \), there exists a left eigenvector, namely, \( x = [0, -b_3, b_2] \), such that \( xb = 0 \). Thus, Theorem 17 implies that \( \{b, A_g b, A_g^2 b\} \) are linearly dependent for all \( b \in \mathbb{R}^3 \).
2.4 PARTIAL IDENTIFIABILITY FROM A CONFINED TRAJECTORY

2.4.1 Analysis

The examples presented in the last section bring up the following question: If the model (2.3) is not identifiable from a single trajectory, how much information about $A$ can be obtained from such a trajectory? The answer to this question is provided by a natural generalization of the results leading to Theorem 13.

**Theorem 19.** Suppose $V$ is a proper linear subspace of $\mathbb{R}^n$ invariant under $A$ with $k = \dim V$. Let $A|_V$ denote the linear operator on $V$ that is obtained as a restriction of $A$ to the subspace $V$. The following statements are equivalent:

(i) $V$ is the minimal $A$-invariant subspace such that $b \in V$.
(ii) $\{b, Ab, \ldots, A^{k-1}b\}$ are linearly independent in $V$.
(iii) The orbit $\gamma(A,b)$ is in $V$ and is not confined to a proper subspace of $V$.
(iv) There exists no $B \in \mathbb{R}^{n \times n}$ such that $B|_V \neq A|_V$ and $x(\cdot; B, b) = x(\cdot; A, b)$.

**Proof.** The proof of the theorem can be constructed by generalization of the proofs discussed above. In particular, proof of the equivalence of (i) and (ii) is analogous to the proof of Lemma 10, proof of the equivalence of (i) and (iii) is analogous to the proof of Lemma 11, and proof of the equivalence of (iii) and (iv) is analogous to the proof of Theorem 13.

It may be of concern that the subspace $V$ depends on the matrix to be identified and hence is not known in advance. However, in view of the statement (iii) of Theorem 19, the subspace $V$ is clearly defined by the orbit $\gamma(A,b)$ as the smallest linear subspace of $\mathbb{R}^n$ containing $\gamma(A,b)$. Therefore, given the trajectory $x(t; A, b)$ one can identify both the invariant subspace $V$ and the restriction $A|_V$ but no more information about the model (2.3).

The restriction operator $A|_V$ is identical to a submatrix of $A$ if the subspace $V$ is a span of vectors from the standard basis $\{e_1, e_2, \ldots, e_n\}$ of $\mathbb{R}^n$. In general, $A|_V$ can be decomposed using a basis $\{v_1, v_2, \ldots, v_k\}$ of $V$ as follows:

$$A|_V = \sum_{i=1}^{k} \sum_{j=1}^{k} \alpha_{ij} A_{ij}$$

(2.5)
where $A_{ij}$ are rank-1 matrices such that $A_{ij}v_j = v_i$, and $\alpha_{ij}$ with $i, j = 1, ..., k$ are determined by the trajectory $x(t; A, b)$. By completing $\{v_1, v_2, \ldots, v_k\}$ into a basis $\{v_1, v_2, \ldots, v_n\}$ of $\mathbb{R}^n$ one can write the unidentified part of the parameter matrix, $A - A|_V$, as

$$A - A|_V = \sum_{i=1}^{n} \sum_{j=k+1}^{n} \alpha_{ij} A_{ij} + \sum_{i=k+1}^{n} \sum_{j=1}^{k} \alpha_{ij} A_{ij}$$

(2.6)

where the remaining coefficients $\alpha_{ij}$ are free parameters.\(^3\)

The full parameter matrix $A$ can be reconstructed from several confined trajectories of the system. The minimum number of such trajectories needed to fully identify $A$ depends on the number and relative positions of the subspaces to which those trajectories are confined.

As a final comment in this section let us note that the results above can also be used to characterize the case in which the model variables are not fully observable, i.e., the case

$$\dot{x}(t) = A x(t), \quad x(0) = b, \quad y(t) = C x(t)$$

(2.7)

in which the known matrix $C$ is not of rank $n$. Suppose that the orbit $\gamma(A, b)$ of the trajectory $x(t; A, b)$ of the system is confined to a subspace $V$. Suppose, in addition, that $\dim V \leq \text{rank } C$ and that $V \cap \text{null } C = 0$. Then one can construct an invertible map $\tilde{C}$ that takes $V$ into range $C$ and determine $x(t; A, b)$ from the observed image of the trajectory $y(t; A, b, C)$ as $x(t; A, b) = \tilde{C}^{-1} y(t; A, b, C)$. Using the procedure above, one can then identify $A|_V$ from the trajectory $x(t; A, b)$. Unfortunately, the information on whether $x(t; A, b)$ is confined and to which subspace cannot be directly ascertained by observing $y(t; A, b, C).

\(^3\)The matrix $[A]_B = [\alpha_{ij}]$ is comprised of the coefficients of $A$ in the basis $B = \{v_1, v_2, \ldots, v_n\}$. 

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2.4.2 Example

The matrix $A_e$ defined in the last section has a two-dimensional invariant subspace $V = \text{span}\{v_1, v_2\}$ where $v_1 = [1, 1, 0]^T$, $v_2 = [0, 0, 1]^T$. Trajectory $x(t; A_e, b)$ starting at $b = [-1, -1, 1]^T$ is confined to $V$ but not to any lower-dimensional subspace of $V$. In accord with Theorem 19, we can use the trajectory $x(t; A_e, b)$ to identify $A_e|_V$. Specifically, we transform $A_e$ to the coordinate system with the basis $B = \{v_1, v_2, v_3\}$, where $v_3 = [0, 1, 0]^T$, as

$$
[A_e]_B = \begin{bmatrix}
-4 & 2 & -3 \\
0 & -2 & 0 \\
0 & 0 & -1
\end{bmatrix}.
$$

The expansion in the basis $\{v_1, v_2\}$ of the restriction $A_e|_V$ is given by the upper left 2x2 submatrix of $[A_e]_B$.

Now, for any matrix $B$ with invariant subspace $V$ and such that $B|_V = A_e|_V$, the trajectory $x(t; B, b)$ is identical to $x(t; A_e, b)$. Each such matrix, in the coordinates of $V$, must agree with the upper left 2x2 block of $[A_e]_B$ and take the form

$$
[B]_B = \begin{bmatrix}
-4 & 2 & \beta_{13} \\
0 & -2 & \beta_{23} \\
0 & 0 & \beta_{33}
\end{bmatrix}
$$

for some $\beta_{13}, \beta_{23}, \beta_{33} \in \mathbb{R}$. Transforming $[B]_B$ to the standard coordinate system yields

$$
B = \begin{bmatrix}
-\beta_{13} - 4 & \beta_{13} & 2 \\
\beta_{13} - \beta_{33} - 4 & \beta_{13} + \beta_{33} & 2 \\
-\beta_{23} & \beta_{23} & -2
\end{bmatrix}.
$$

Note that in this particular case, the third column of matrix $A$ is identifiable from the confined trajectory $x(t; A_e, b)$. 

30
2.5 SYSTEMS THAT ARE LINEAR IN PARAMETERS

2.5.1 Analysis

Consider now a nonlinear dynamical system that depends linearly on parameters and has only its initial condition as a control:

\[ \dot{x}(t) = Af(x(t)) \]
\[ x(0) = b. \]  

(2.8)

In equation (2.8), \( x(t) \in \mathbb{R}^n \) is the state of the system at time \( t \), the system parameters are the entries of the coefficient matrix \( A \in \mathbb{R}^{n \times m} \), \( b \in \mathbb{R}^n \) is the initial condition, and \( f = (f_1, ... , f_m) : \mathbb{R}^n \rightarrow \mathbb{R}^m \), where \( f_j \) are functions that are locally Lipschitz continuous in \( x \).

The initial condition \( b \) determines the trajectory \( x(t; A, b) \) of the system (2.8) for any given \( A \). A definition of identifiability, analogous to Definition 1, can be stated as follows:

**Definition 20.** Model (2.8) is identifiable in \( \Omega \subseteq \mathbb{R}^{n \times m} \) if and only if for all \( A, B \in \Omega \), \( A \neq B \), there exists \( b \in \mathbb{R}^n \) such that \( x(\cdot; A, b) \neq x(\cdot; B, b) \).

Recall that for the linear model (2.3), Theorem 2 implies identifiability in \( \mathbb{R}^{n \times n} \). Such general identifiability, however, does not hold for systems that are linear in parameters. Instead, we can obtain a necessary condition for identifiability of the model (2.8) using the properties of the map \( f \). We first introduce \( \mathcal{F} \), the largest linear subspace of \( \mathbb{R}^m \) that contains the range of \( f \), as \( \mathcal{F} = \text{span}\{f(x) | x \in \mathbb{R}^n\} \subseteq \mathbb{R}^m \). The dimension of \( \mathcal{F} \) determines the identifiability of the model (2.8) as follows:

**Theorem 21.** If \( \dim \mathcal{F} < m \), then model (2.8) is not identifiable in \( \mathbb{R}^{n \times m} \).

**Proof.** Assume \( \dim \mathcal{F} = r < m \). Let \( \{v_1, ..., v_r\} \) be a basis for \( \mathcal{F} \). Complete \( \{v_1, ..., v_r\} \) to a basis \( \{v_1, ..., v_r, v_{r+1}, ..., v_m\} \) of \( \mathbb{R}^m \) and let \( V = [v_1 | ... | v_m] \). For fixed \( A \in \mathbb{R}^{n \times m} \), choose \( C \in \mathbb{R}^{n \times (m-r)} \) such that \( C \neq [Av_{r+1} | ... | Av_m] \) and let \( D = [Av_1 | ... | Av_r | C] \). Finally, let \( B = DV^{-1} \). This procedure yields \( B \in \mathbb{R}^{n \times m} \) such that (i) \( Av_j = Bv_j \), \( j = 1, ..., r \) and (ii) \( Av_j \neq Bv_j \) for some \( j \in \{r + 1, ..., m\} \). Since (i) implies that \( Af(x) = Bf(x) \) \( \forall x \in \mathbb{R}^n \), the trajectories \( x(t; A, b) \) and \( x(t; B, b) \) of the model (2.8) are identical for any initial condition \( b \in \mathbb{R}^n \), yet (ii) implies that \( B \neq A \). Hence, the model (2.8) is not identifiable in \( \mathbb{R}^{n \times m} \).
One of the main results for linear systems described in Section 2.3 was Theorem 13, which provides a connection between the uniqueness of the parameters of a model (2.3) and the confinement of the orbit of that model. Interestingly, a similar result can be shown for model (2.8), but here, parameter uniqueness is instead linked to the geometric structure of the image of the orbit of the model in the flux space. Let \( \phi_{A,b}(t) = \{ f(x(t; A, b)) \mid t \in \mathbb{R} \} \) be a curve in \( \mathbb{R}^m \) that represents the image under the map \( f \) of the orbit \( \gamma(A, b) \) of the model (2.8).

**Theorem 22.** There exists no \( B \in \mathbb{R}^{n \times m} \) with \( A \neq B \) such that \( x(\cdot; A, b) = x(\cdot; B, b) \) if and only if \( \phi(A, b) \) is not confined to a proper subspace of \( \mathbb{R}^m \).

**Proof.** To prove the reverse implication, suppose that \( \phi(A, b) \) is not confined to a proper subspace of \( \mathbb{R}^m \). Then one can find the matrix \( A \) uniquely as follows: Since \( \phi(A, b) \) is not confined, one can choose \( m \) distinct time points \( t_1, \ldots, t_m \), such that the matrix \( F = [f(x(t_1)) \mid \ldots \mid f(x(t_m))] \) is invertible. From knowledge of the trajectory, one knows \( \dot{x}(t_1), \ldots, \dot{x}(t_m) \). Appending this information into a matrix yields the linear equation \( \begin{bmatrix} \dot{x}(t_1) \mid \ldots \mid \dot{x}(t_m) \end{bmatrix} = AF \), which has a unique solution \( A = [\dot{x}(t_1) \mid \ldots \mid \dot{x}(t_m)]F^{-1} \). Therefore there is no other matrix \( B \) that would produce a trajectory identical to \( A \).

The forward direction is proven by contrapositive utilizing an argument similar to the one used in the proof of Theorem 21. Assume that \( \phi(A, b) \) is confined to a proper subspace of \( \mathbb{R}^m \) and define \( F_\gamma = \text{span}\{f(x(t, A, b)), t \in [0, \infty)\} \). By the confinement of \( \phi(A, b) \), \( \dim F_\gamma = r < m \). Let \( V = \{v_1, \ldots, v_r\} \) be a basis for \( F_\gamma \). Complete \( V \) to a basis of \( \mathbb{R}^m \), \( \hat{V} = \{v_1, \ldots, v_r, v_{r+1}, \ldots, v_m\} \). Construct \( B \in \mathbb{R}^{n \times m} \) such that \( Av_j = Bv_j, j = 1, \ldots, r \), but \( Av_j \neq Bv_j \) for some \( j \in \{r+1, \ldots, m\} \) (as in the previous proof). With this construction, \( A \neq B \), but \( Af(x) = Bf(x) \forall x \in \mathbb{R}^n \), and hence \( x(t; A, b) = x(t; B, b) \).

As in the linear case, this theorem is a statement about identifiability of the model from a single trajectory. In fact, since the linear model (2.3) is a special case of the nonlinear model (2.8) with \( f \) being the identity map, Theorem 13 is a special case of Theorem 22. In that case, the confinement of the orbit image \( \phi(A, b) \) in the flux space is equivalent to the confinement of the orbit \( \gamma(A, b) \) of the trajectory.
2.5.2 Examples

As an illustrative example, consider the system,

\[
\begin{align*}
\dot{x} &= a_{11}x + a_{12}xy + a_{13}y \\
\dot{y} &= a_{21}x + a_{22}xy + a_{23}y
\end{align*}
\] (2.9)

System (2.9) is linear in parameters and can be represented in the form of model (2.8) with

\[
A = \begin{bmatrix}
a_{11} & a_{12} & a_{13} \\
a_{21} & a_{22} & a_{23}
\end{bmatrix}
\]

and \(f(x, y) = [x, xy, y]^T\). For the matrix

\[
A = \begin{bmatrix}
-2 & 1 & -2 \\
-1 & 1 & -3
\end{bmatrix}
\]

and initial condition \(b = [1, 1]^T\), \(\phi(A, b)\) is confined to a proper subspace of \(\mathbb{R}^3\), as shown in figure (7). Theorem 22 implies that there exists a matrix \(B \in \mathbb{R}^{2 \times 3}\) with \(A \neq B\) such that \(x(t; A, b) = x(t; B, b)\). One can construct \(B\) using the procedure described in the proof of the theorem. Since \(x(t) = y(t)\) for the solution of the IVP, we have \(F_{\gamma} = \)
span\{(x, x^2, x)^T | x \in \mathbb{R}\}. Thus \{(1, 0, 1)^T, (0, 1, 0)^T\} is a basis for \(F_\gamma\). Complete this basis to \(\{v_1, v_2, v_3\} = \{(1, 0, 1)^T, (0, 1, 0)^T, (1, 0, -1)^T\}\), a basis of \(\mathbb{R}^3\). Let
\[
D = [Av_1 | Av_2 | w] = \begin{bmatrix}
-4 & 1 & w \\
-4 & 1 \\
\end{bmatrix}
\]
such that \(w \neq Av_3 = [0, 2]^T\). For example, let
\[
D = \begin{bmatrix}
-4 & 1 & -2 \\
-4 & 1 & -6
\end{bmatrix}.
\]
Then
\[
B = D[v_1 | v_2 | v_3]^{-1} = \begin{bmatrix}
-3 & 1 & -1 \\
-5 & 1 & 1
\end{bmatrix}
\]
is one such example.

Another example is provided by the Lotka-Volterra model of competing species [21, 66],
\[
\begin{align*}
\dot{x} &= x(1 - ax - cy) \\
\dot{y} &= y(1 - ay - cx).
\end{align*}
\]  \tag{2.10}
Model (2.10) is linear in parameters with
\[
A = \begin{bmatrix}
1 & -a & -c & 0 & 0 \\
0 & 0 & -c & 1 & -a
\end{bmatrix}
\]
and \(f(x, y) = [x, x^2, xy, y, y^2]^T\).

For any initial condition of the form \(b = [x_0, x_0]^T\), the trajectory of the system obeys \(x(t) = y(t)\) and depends on the sum \(a + c\) but not the individual values of the parameters \(a, c\). This is consistent with Theorem 22, which concludes that since \(f(x, x) = [x, x^2, x^2, x, x^2]^T\) and \(\phi(A, b)\) is confined to a proper subspace of \(\mathbb{R}^5\), the parameter matrix which yields this trajectory is not unique, and therefore model (2.10) is not identifiable in \(\mathbb{R}^{2 \times 5}\).
2.6 DISCRETE DATA

In practical applications, one does not typically have knowledge of a full trajectory of the system, but rather a sample of discrete data points that lie on a trajectory, possibly perturbed by measurement noise. Suppose for now that we have \(m\) accurate data points \(x_0, x_1, \ldots, x_{m-1}\) in \(\mathbb{R}^n\) that lie on the trajectory of the model (2.3), defined as 
\[x_k = x(t_k; A, b), k = 0, 1, \ldots, m - 1,\]
where \(t_0, t_1, \ldots, t_{m-1}\) are distinct time points. Non-confinement of the orbit \(\gamma(A, b) = \{x(t; A, b) : t \in \mathbb{R}\}\) is obviously determined by the dimension of the span of the data vectors.:

**Lemma 23.** Orbit \(\gamma(A, b)\) is not confined to a proper subspace of \(\mathbb{R}^n\) if and only if there exist \(t_0, t_2, \ldots, t_{n-1}\) such that \(x_0, x_2, \ldots, x_{n-1}\) are linearly independent.

Note that the confinement of an orbit cannot be established from a fixed, finite data set. For example, in a two-dimensional system, the available data may lie in a straight line, but the underlying solution may be a spiral, which is not a confined orbit. Model (2.3) is identifiable from a full trajectory in this example, since the corresponding orbit is not confined to a proper subspace.

It now remains to show how the parameter matrix \(A\) is determined from the data. The parameter matrix \(A\) can be computed explicitly without the need for optimization from \(n + 1\) data points which are uniformly spaced in time. Let \(x_0, \ldots, x_n\) be such that \(x_k = x(t_k; A, b)\) for each \(k = 0, \ldots, n\), with \(t_{k+1} - t_k = \Delta t\) for all \(k = 0, \ldots, n - 1\). Assume that every collection of \(n\) of the data points are linearly independent. Let \(\Phi(\Delta t)\) denote the principal matrix solution of model (2.3) and let \(X_0\) and \(X_1\) denote the matrices \([x_0 \mid x_1 \mid \ldots \mid x_{n-1}]\) and \([x_1 \mid x_2 \ldots \mid x_n]\), respectively. The principal matrix solution provides a relation between the data given by \(\Phi(\Delta t)x_k = x_{k+1}\), thus \(\Phi(\Delta t)X_0 = X_1\). Without loss of generality, we can let \(\Delta t = 1\) and define \(\Phi = \Phi(1)\). Then, by the invertibility of \(X_0\) we find that,

\[\Phi = X_1(X_0)^{-1}.\]  

(2.11)

In theory, the matrix \(A\) can be computed by taking the matrix logarithm of \(\Phi\), since \(\Phi = e^A\). It is important to note, however, that the logarithm of a matrix does not always exist, and if
it does, it is not necessarily unique. Requirements for the existence of a real matrix logarithm are given in the following theorem [17].

**Theorem 24.** Let \( \Phi \) be a real square matrix. Then there exists a real solution \( A \) to the equation \( \Phi = e^A \) if and only if \( \Phi \) is nonsingular and each Jordan block of \( \Phi \) belonging to a negative eigenvalue occurs an even number of times.

In our case, \( \Phi = X_1(X_0)^{-1} \) is nonsingular due to the linear independence of the data. The second condition of Theorem 24 is satisfied trivially if \( x_0, x_1, \ldots, x_n \) are indeed discrete points on a trajectory of the linear model (2.3). More importantly, uniqueness of the matrix logarithm, which is directly related to the identifiability of the model (2.3), is addressed in the following theorem [17].

**Theorem 25.** Let \( \Phi \) be a real square matrix. Then there exists a unique real solution \( A \) to the equation \( \Phi = e^A \) if and only if all the eigenvalues of \( \Phi \) are positive real and no Jordan block of \( \Phi \) belonging to any eigenvalue appears more than once.

Given the data \( x_0, x_2, \ldots, x_n \), the matrix logarithm yields a unique corresponding parameter matrix \( A \) if and only if \( \Phi \), defined using (2.11), satisfies the hypotheses of Theorem 25. We have established earlier (in Theorem 15) that model (2.3) can be identified from some initial condition \( b \) if and only if \( A \) has only one Jordan block for each of its eigenvalues. If \( A \) satisfies this requirement and has real eigenvalues, then \( \Phi \) has positive eigenvalues and has one Jordan block for each of them and hence satisfies the hypotheses of Theorem 25. We can then conclude that model (2.3) is identifiable from the data \( x_0, x_1, \ldots, x_n \). If, however, \( A \) has one Jordan block for each of its eigenvalues and has complex eigenvalues, then \( \Phi \) has a negative eigenvalue and a pair of Jordan blocks associated to each pair of complex eigenvalues of \( A \). For example,

\[
A = \begin{bmatrix} 0 & \pi \\ -\pi & 0 \end{bmatrix}
\]

has eigenvalues \( \pm \pi i \) and has no repeated Jordan blocks, yet

\[
e^A = \begin{bmatrix} -1 & 0 \\ 0 & -1 \end{bmatrix}
\]
with a repeated Jordan block and negative eigenvalues. In such a case, Theorem 25 implies that one can find a matrix $B \neq A$ such that $e^B = \Phi = e^A$. Although the trajectories $x(t; A, x_0)$ and $x(t; B, x_0)$ must necessarily differ (since the model with matrix $A$ is identifiable from the full trajectory $x(t; A, x_0)$), the data obtained from these trajectories for the same set of time points $\{t_0, t_2, \ldots, t_n\}$ are identical. Thus, we have the following observation.

**Corollary 26.** The model (2.3) with matrix $A$ that has a pair of complex eigenvalues is not identifiable from any set of data $x_0, x_2, \ldots, x_n$ that are uniformly spaced in $t$.

Figure 8 illustrates the non-identifiability that arises with complex eigenvalues in the case of discrete data for 2x2 linear systems. In this example, the solution to the system with parameter matrix

$$A_r = \begin{bmatrix} -3/2 & -4 \\ 4 & -3/2 \end{bmatrix} + 2\pi r \begin{bmatrix} 0 & -1 \\ 1 & 0 \end{bmatrix}$$

will satisfy the data for any integer value of $r$. The solutions shown in Figure 8 are for the cases $r = 0, 1, \text{ and } -1$.

The extension of the above computation to the model (2.8) is simple, provided one can measure not only the values $x_1, x_2, \ldots, x_m$ of the variables at $t_1, t_2, \ldots, t_m$, but also their
rates of change $y_1, y_2, \ldots, y_m$, where $y_k = dx(t; A, b)/dt|_{t=t_k}$. Just as in the proof of Theorem 22, if $f(x_1), f(x_2), \ldots, f(x_n)$ are linearly independent (and hence $\phi(A, b)$ is not confined to a proper subspace), then the unique parameter matrix that yields this data is given by

$$A = YF^{-1} \tag{2.12}$$

where $Y = [y_1 | \ldots | y_m]$ and $F = [f(x_1) | \ldots | f(x_m)]$.

The computations outlined above are valuable not only because they offer direct methods for computing the parameter matrix $A$ that do not rely on minimization of an error function, but also because they can be used to provide insight into the sensitivity of $A$ to the data. Since both computations are based on linear algebraic operations, one can use the tools of numerical analysis to determine the conditioning of the problem (see, e.g., [83]). In the case of the linear model (2.3), equation (2.11) implies that the problem of computing $\Phi$ has condition number $\kappa(X_0) = \|X_0\|\|X_0^{-1}\|$, and indeed one can compute that any perturbations of the data ($\delta X_0$ of $X_0$ and $\delta X_1$ of $X_1$) induce a perturbation $\delta \Phi$ of $\Phi$ that obeys

$$\frac{\|\delta \Phi\|}{\|\Phi\|} \leq \kappa(X_0) \left( \frac{\|\delta X_0\|}{\|X_0\|} + \frac{\|\delta X_1\|}{\|X_1\|} \right) \tag{2.13}$$

where $\|\cdot\|$ is any norm of choice. Note that if $\|\cdot\|$ is the Euclidean norm $\|\cdot\|_2$, then $\kappa(X_0)$ equals the ratio of the largest to the smallest singular values of $X_0$. It follows that the closer are the data vectors $x_0, x_1, \ldots, x_{n-1}$ (i.e., the columns of $X_0$) to being linearly dependent, the more ill-conditioned is the inverse problem.

Likewise, in the case of the model (2.8), equation (2.12) implies that the problem of computing $A$ has the condition number $\kappa(F) = \|F\|\|F^{-1}\|$, and hence any perturbations $\delta F$ and $\delta Y$ induce a perturbation $\delta A$ that obeys

$$\frac{\|\delta A\|}{\|A\|} \leq \kappa(F) \left( \frac{\|\delta F\|}{\|F\|} + \frac{\|\delta Y\|}{\|Y\|} \right). \tag{2.14}$$

Again, the closer are the vectors $f(x_1), f(x_2), \ldots, f(x_m)$ (i.e., the columns of $F$) to being linearly dependent, the more ill-conditioned is the inverse problem.

The observations made at the end of this section indicate that the highest accuracy in the inverse problem (and hence the lowest sensitivity to measurement errors) can be achieved by selecting data so as to minimize the condition number of the data matrix. This result
has an important practical implication: although the results in Sections 2.3 and 2.5 indicate that for an identifiable model any infinitesimally small portion of a trajectory is sufficient to identify the parameter matrix, in any practical situation a small segment of trajectory will have a nearly linear orbit and hence any selection of data from that segment will yield a data matrix with high condition number. Thus, in order to minimize the condition number, one must have a sufficiently large portion of the trajectory that explores all dimensions of the underlying space.

2.7 CONCLUSIONS

In this chapter, for both linear and nonlinear dynamical systems, we have derived necessary and sufficient conditions for identifiability of parameters from a single trajectory based solely on the geometry of the trajectory or the geometry of an image of the trajectory. Furthermore, we have shown that an improved accuracy of parameter estimation can result when the trajectory deviates farther from being confined. These results have a practical utility since the criterion can be applied using only what is known about the trajectory, without any knowledge of the model parameters. Additional results for linear systems include a link between identifiability from single trajectory with initial condition \( b \) and the linear independence of \( \{b, Ab, ..., A^{n-1}b\} \), several characterizations of the linear independence of \( \{b, Ab, ..., A^{n-1}b\} \) including a condition on the Jordan form of \( A \), and the result that unconditional identifiability cannot occur outside of \( \mathbb{R}^{2n \times 2n} \). Finally, we addressed the question of explicitly computing model parameter values from a discrete collection of data points.

There are several directions for possible extension of the results in this chapter. First, our results imply that discrete data contained within a lower-dimensional subspace of the full state space will not yield identifiability of an underlying system. Such data may arise, however, from particular samplings of a trajectory that is not confined in this way. The derivation of more general identifiability conditions from discrete data remains for future exploration. Second, the condition \( C = I \) is highly restrictive, because in real scenarios not all variables of the system may be observable. We have shown that when \( C \) is not of full
rank, then confinement of an orbit to an invariant subspace of dimension not greater than the rank of $C$ can yield partial identifiability of the parameter matrix. A natural extension of the present study would be to investigate if there are ways to enhance the practical applicability of this result or to obtain more general identifiability results in this case. A third direction for future study would be the consideration of general nonlinear systems. We have shown that identifiability conditions based on confinement can be derived for systems that feature linearity in parameters, regardless of whether the dynamics is linear or nonlinear. For nonlinear dynamical systems lacking this form of parameter dependence, the assessment of identifiability will likely require new ideas. The prospects for addressing this problem using linearization about trajectories appear to be limited, based on our observations about sensitivity associated with parameter estimation from small segments of trajectories. A final direction to consider is the identifiability of systems that are linear or linear in parameters with time-dependent parameter matrices. Our methods would likely be useful for systems with rather trivial time-dependence, such as piecewise constant parameter matrices, where switching times between different constant values are known and full solution trajectories are available, but handling more general time-dependence appears to be a difficult problem. These and other related topics represent important directions for follow-up studies.

In the next chapter, we will continue to look at the case of discrete single trajectory data. Moving beyond the setting of error free data, we will investigate properties of the inverse solution in the presence of uncertainty in the data.
3.0 ROBUSTNESS OF SOLUTIONS OF THE INVERSE PROBLEM FOR LINEAR DYNAMICAL SYSTEMS WITH UNCERTAIN DATA

3.1 INTRODUCTION

A fundamental problem in modeling temporally evolving systems is the determination of model parameter values from experimental observations collected at specific time points. Since models can be viewed as forward mappings from sets of parameter values to time-dependent states of model variables, the problem of parameter estimation is often referred to as an inverse problem. Although parameter estimation has received significant attention in the literature, certain fundamental questions about the inverse problem still remain open. Solving the inverse problem becomes even more challenging in the presence of uncertainty in experimental measurements, as may arise due to measurement errors and fluctuations in system components. The overall goal of this chapter is to derive estimates of the degree of uncertainty in data to which properties of the inverse problem, such as existence and uniqueness of solutions, are robust.

We start by addressing fundamental issues of existence and uniqueness of solutions to the inverse problem based on a discrete collection of linearly independent data points assumed to be known without uncertainty, before turning to the uncertain case. We focus our analysis on linear models, which are prevalent in the study of many important applications including pharmacokinetics, linear response theory for mechanical and electronic systems, continuous time Markov chain probabilistic models, and near-equilibrium responses of nonlinear systems [11, 31, 3, 73]. In addition to their applicability, linear systems are convenient because in the linear case, there is an explicit structure of the associated forward solution map that can be exploited. Furthermore, we mostly consider data points that are equally spaced in time, as
may be obtained from experiments with regimented data collection schedules and for which it may be possible to explicitly solve for the linear system parameter matrix. Despite these advantages, the inverse problem is nontrivial because the solution to a linear dynamical system depends nonlinearly on its parameters.

Specifically, in Section 3.2, we begin by considering data with no uncertainty, where classical results on matrix logarithms yield necessary and sufficient conditions for the data to specify a unique parameter matrix $A$ that solves the inverse problem. Contrary to our expectations, we find that the existence of model parameters corresponding to given data is guaranteed only for a restricted subset of potential data sets and that there is only a limited region in data space that yields a unique set of parameters. Subsequently, we explore how uncertainty in the data impacts the existence and uniqueness of $A$. In Section 3.3, we provide conditions that ensure that existence or uniqueness of the inverse problem solution is guaranteed to hold in an open neighborhood of data, and we make use of a convenient transformation to better characterize regions in which $A$ is unique. These steps prepare us for Section 3.4, where we present the main results of the chapter, consisting of analytical and numerical estimates of the maximum uncertainty in the data under which the properties of the inverse problem are certain to be preserved. Examples for 2-dimensional systems are shown in Section 3.5, where we first define regions in data space for which the solution to the inverse problem has various properties and then illustrate bounds on the maximal permissible uncertainty for those properties. Finally, in Section 3.6, we briefly remark on the case of non-equally-spaced data points, and we conclude with a discussion in Section 3.7, which includes some comments on open directions and related work in the past literature.

3.2 DEFINITIONS AND PRELIMINARIES

As in Chapter 2, we consider the model defined as a finite-dimensional linear dynamical system; recall equation (2.3):

$$
\begin{align*}
\dot{x}(t) &= A x(t) \\
x(0) &= b.
\end{align*}
$$
In equation (2.3), $x(t) \in \mathbb{R}^n$ is the state of the system at time $t$, the system parameters are the entries of the coefficient matrix $A \in \mathbb{R}^{n \times n}$ and $b \in \mathbb{R}^n$ is the initial condition. For clarity of exposition we will refer to the entire matrix $A$ as the parameter $A$. We shall define the parameter space $\mathcal{P}$ as the set of all parameters $A$ and initial conditions $b$.

For a fixed $A$, system (2.3) has a well defined solution, or trajectory, given by $x(t; A, b) = \Phi(t)b$, where $\Phi(t) = e^{At}$ is the principal matrix solution. The data representing the system is a set of observations of the trajectory values. We assume that data for all of the state variables is available and denote by $\mathcal{D}$ the data space consisting of a set of $(n + 1)$-tuples $d = (x_0, x_1, x_2, \ldots, x_n)$ of points $x_j \in \mathbb{R}^n$. Each such $d$ will be referred to as a data set and each $x_j$ as a data point. Sampling the trajectory $x(t; A, b)$ at equally spaced times (without loss of generality $\Delta t = 1$) yields an element of $\mathcal{D}$, namely a data set composed of the specific data points $x_j = x(j; A, b) \in \mathbb{R}^n$, $j = 0, 1, \ldots, n$; non-uniformly spaced data are discussed in Section 3.6. We use solution map to refer to the map $F : \mathcal{P} \to \mathcal{D}$ from parameter space to data space defined as $F(A, b) = (x_0, x_1, x_2, \ldots, x_n)$ for this choice of $\{x_j\}$ sampled from $x(t; A, b)$. From the definition of $x_j$, it is immediately clear that $b = x_0$ and thus, we focus on the problem of determining the parameter $A$. The inverse problem is then the problem of inverting the map $F$ to find $F^{-1}(d)$ (i.e., $A$ such that $F(A, b = x_0) = d$) for a given data set $d$. If the data set $d$ is obtained from experimental measurements or some other outside source, then this problem may or may not have a solution.

We set out to derive necessary and sufficient conditions that a data set $d \in \mathcal{D}$ must satisfy so that there exists a unique real matrix $A$ for which the dynamical system (2.3) produces data $d$ (i.e., so that $d = F(A, b)$). These conditions define a subset of the data space on which the inverse map $F^{-1}$ is well defined. Given a uniformly spaced data set $d \in \mathcal{D}$, one can attempt to solve the inverse problem as presented in Section 2.6. We review the constructs here, as this notation is referenced several times throughout this chapter. Denote by $X_0$ and $X_1$ the $n \times n$ matrices $[x_0 \mid \ldots \mid x_{n-1}]$ and $[x_1 \mid \ldots \mid x_n]$, respectively. The principal matrix solution provides a relation between the data points; letting $\Phi := \Phi(1) = e^A$, we have $x_{j+1} = \Phi x_j$, which implies that $X_1 = \Phi X_0$ and hence $\Phi = X_1 X_0^{-1}$. All that remains is to find $A$ as the matrix logarithm of $\Phi$. Thus, from an operational standpoint, the map $F^{-1}$ is a composition of two nonlinear maps: (i) The map $G : \mathcal{D} \to \mathbb{R}^{n \times n}$ defined by $G(d) = \Phi$, which
is defined (and continuous) at all points $d$ such that $x_0, \ldots, x_{n-1}$ are linearly independent (i.e. wherever $X_0^{-1}$ exists), and (ii) the matrix logarithm map, denoted here as $L : \mathbb{R}^{n \times n} \rightarrow \mathcal{P}$ and defined as $A = L(e^A)$. Hence, $F^{-1} = L \circ G$ is well defined if (i) $X_0$ is invertible, (ii) the matrix logarithm of $\Phi$ exists, and (iii) the matrix logarithm of $\Phi$ is unique. The case when condition (i) fails was studied extensively in Chapter 2; in that case the system (2.3) generating the data $d$ either does not exist or is not identifiable. Conditions (ii) and (iii) can be addressed with the help of theorems 24 and 25 by Culver [17], presented in Section 2.6, which characterize the existence and uniqueness of a real matrix logarithm.

We note that even when a matrix logarithm exists, there are still issues with how to compute it. Numerical methods for computing the logarithm of a matrix are discussed in [37, 14, 1].

### 3.3 EXISTENCE AND UNIQUENESS OF THE INVERSE IN $N$ DIMENSIONS

As will become clear shortly, Theorems 24 and 25 identify matrices $\Phi$ that are not robust in the sense that they form a set of zero measure. Since we aim to discuss the properties of inverse problem solutions for uncertain data, such as data that is perturbed due to noise, it makes sense to determine conditions that ensure that a given matrix $\Phi$ is inside an open set of matrices with particular existence or uniqueness properties.

#### 3.3.1 Inverse problems on open sets

We now state and prove three corollaries of Theorems 24 and 25 that are useful for considering uncertain data and, as it turns out, avoid the conditions on Jordan blocks that can become overly cumbersome for practical use in $n$ dimensions. The first corollary characterizes open sets of matrices that have real logarithms, the second corollary characterizes open sets of matrices that have unique real logarithms, and the third corollary characterizes open sets of matrices that do not have real logarithms.
Corollary 27. (to Theorem 24) Let $\Phi^*$ be an $n \times n$ real matrix. The following statements are equivalent:

(a) There exists an open set $U \subset \mathbb{R}^{n \times n}$ containing $\Phi^*$ such that for any $\Phi \in U$ the equation $\Phi = e^A$ has an $n \times n$ real solution $A$.
(b) $\Phi^*$ has only positive real or complex eigenvalues.

Proof. Suppose that $\Phi^* \in \mathbb{R}^{n \times n}$ has only positive real or complex eigenvalues. Then the result follows immediately by Theorem 24 and the continuous dependence of eigenvalues on matrix entries.

For the converse, suppose that $\Phi^* \in \mathbb{R}^{n \times n}$ has a real matrix logarithm, i.e., $\Phi^* = e^A$ where $A$ is $n \times n$ real matrix. By Theorem 24, either $\Phi^*$ satisfies (b), or $\Phi^*$ has at least one negative eigenvalue and the corresponding Jordan block occurs an even number of times. We now show that the second alternative contradicts the existence of the open set $U$. To this end, let $\Phi^* = QJQ^{-1}$, where

$$J = \begin{bmatrix} J_1 & 0 \\ 0 & J_2 \\ \vdots & \ddots \end{bmatrix}$$

is a Jordan canonical form of $\Phi^*$ with $J_1$ a Jordan block corresponding to a negative eigenvalue. Let $B = QKQ^{-1}$ where,

$$K = \begin{bmatrix} aI & 0 \\ 0 & 0 \end{bmatrix},$$

with $a \in \mathbb{R}$ and $I$ the identity matrix of the same size as $J_1$. Then for every sufficiently small nonzero $a$, $\Phi + B$ has a negative eigenvalue for which the corresponding Jordan block occurs exactly once and hence there is no real $A$ such that $\Phi + B = e^A$.

Corollary 28. (to Theorem 25) Let $\Phi^*$ be an $n \times n$ real matrix. The following statements are equivalent:

(a) There exists an open set $U \subset \mathbb{R}^{n \times n}$ containing $\Phi^*$ such that for any $\Phi$ in $U$ the equation $\Phi = e^A$ has a unique $n \times n$ real solution $A$.
(b) $\Phi^*$ has $n$ distinct positive real eigenvalues.
Proof. The proof is similar to the previous one. Condition (b) implies condition (a) based on Theorem 25 and continuity. For the converse, suppose that $\Phi^* \in \mathbb{R}^{n \times n}$ has a unique real matrix logarithm, i.e., there is a unique $A \in \mathbb{R}^{n \times n}$ such that $\Phi^* = e^A$, and $\Phi^*$ is in the interior of an open set of such matrices. By Theorem 25, $\Phi^*$ has positive real eigenvalues and no Jordan block appears more than once. There can be more than one Jordan block for the same eigenvalue but those Jordan blocks must be of different sizes. We can write $\Phi^* = QJQ^{-1}$ where $J$ is the Jordan canonical form of $\Phi^*$ as defined in (3.1) and $J_i$ are ordered in size from largest to smallest, such that no two $J_i$ are the same.

Suppose now that the largest Jordan block, $J_1$, is not a $1 \times 1$ matrix. Let $p_J(t)$ be the characteristic polynomial of $J$, i.e., $p_J(t) = \prod_{i=1}^{k} (\lambda_i - t)^{d_i}$ where $\lambda_i$ are all real positive and $d_1 \geq 2$. Let $B = QKQ^{-1}$ where $K$ is an $n \times n$ matrix with all zero entries except $L_{12} = a$, $L_{21} = -a$. Then, for every $a > 0$, $\tilde{J} = J + K$ has entries $\tilde{J}_{12} = 1 + a$, $\tilde{J}_{21} = -a$, which implies that $\tilde{J}$ has characteristic polynomial

$$p_{\tilde{J}}(t) = [(\lambda_1 - t)^2 + a(1 + a)] (\lambda_1 - t)^{d_1-2} \prod_{i=2}^{k} (\lambda_i - t)^{d_i}$$

with two complex roots. It follows that $\Phi^* + B = e^A$ does not have a unique real solution $A$ for $a > 0$, which contradicts the assumption that $A$ is unique for each $\Phi$ in an open set containing $\Phi^*$. Thus, all Jordan blocks $J_i$ are of size 1 and furthermore, since no two $J_i$ can be the same, $\Phi^*$ has $n$ distinct eigenvalues. \qed

Statements similar to Corollaries 27, 28 can be made to establish the existence of open sets of matrices with other properties. An example follows.

**Corollary 29.** (to Theorem 24) Let $\Phi^*$ be $n \times n$ real matrix. The following statements are equivalent:

(a) There exists an open set $U \subset \mathbb{R}^{n \times n}$ containing $\Phi^*$ such that for each $\Phi \in U$ the equation $\Phi = e^A$ does not have an $n \times n$ real solution $A$.

(b) $\Phi^*$ has at least one negative eigenvalue of odd multiplicity.

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Proof. Suppose that $\Phi^* \in \mathbb{R}^{n \times n}$ has at least one negative eigenvalue of odd multiplicity. Then, there is at least one Jordan block associated to it that occurs an odd number of times and hence, by Theorem 24, there is no $A \in \mathbb{R}^{n \times n}$ such that $\Phi^* = e^A$. Moreover, there exists an open neighborhood of $\Phi^*$ for which there remains at least one negative eigenvalue of odd multiplicity. $^1$

For the converse, suppose that there exists an open set $U \subset \mathbb{R}^{n \times n}$ containing $\Phi^*$ such that for each $\Phi \in U$ the equation $\Phi = e^A$ does not have an $n \times n$ real solution $A$. Since $\Phi^* = e^A$ has no real solution, by Theorem 24, either $\Phi^*$ is singular, or there is a negative eigenvalue of $\Phi^*$ which belongs to a Jordan block that appears an odd number of times, or both.

If $\Phi^*$ is singular with no Jordan block associated to a negative eigenvalue occurring an odd number of times, then in every neighborhood of $\Phi^*$ there exists a $\Phi$ which is nonsingular with the same condition on Jordan blocks, and $\Phi = e^A$ will have a real solution $A$. This contradicts the existence of $U$.

Thus, there is at least one negative eigenvalue of $\Phi^*$ for which a Jordan block repeats an odd number of times. Denote these negative eigenvalues by $\lambda_1, \ldots, \lambda_r$, $r \geq 1$. Let $m_{\lambda_i}$ be the multiplicity of $\lambda_i$.

Suppose that all $m_{\lambda_i}$ are even. Let $\Phi^* = QJQ^{-1}$ where $J$ is the Jordan canonical form of $\Phi^*$ as defined in (3.1), let $J_{\lambda_1}^1, J_{\lambda_1}^2, \ldots, J_{\lambda_1}^k$ denote all of the Jordan blocks associated to $\lambda_1$ (the blocks may have the same size), and denote by $J_1$ the block-diagonal matrix comprised of $J_{\lambda_1}^1, J_{\lambda_1}^2, \ldots, J_{\lambda_1}^k$.

For $i = 1, \ldots, k$ define $W_{i}^{\lambda_1}$ to be a block-diagonal matrix of the same dimension as $J_{i}^{\lambda_1}$ comprised of $K$, where

$$K = \begin{bmatrix} 0 & a \\ -a & 0 \end{bmatrix}$$

for some $a > 0$. If the dimension of $J_{i}^{\lambda_1}$ is even and equal to $2s$, then $W_{i}^{\lambda_1}$ is comprised of exactly $s$ blocks $K$, the characteristic polynomial of $J_{i}^{\lambda_1} + W_{i}^{\lambda_1}$ is $p(t) = [(\lambda_1 - t)^2 + a(1 + a)]^s$ and, since $a > 0$, $J_{i}^{\lambda_1} + W_{i}^{\lambda_1}$ has only complex eigenvalues.

$^1$A perturbation may split it into a collection of distinct negative eigenvalues of various multiplicities and/or complex pairs, but at least one negative eigenvalue with odd multiplicity will remain.
If is the dimension of $J_{i}^{\lambda_{1}}$ is odd and equal to $2s + 1$, then $W_{i}^{\lambda_{1}}$ contains $s$ blocks $K$ and a zero block of size one, the characteristic polynomial of $J_{i}^{\lambda_{1}} + W_{i}^{\lambda_{1}}$ is $p(t) = [(\lambda_{1} - t)^{2} + a(1 + a)]^{s} (\lambda_{1} - t)$, and since $a > 0$, $J_{i}^{\lambda_{1}} + W_{i}^{\lambda_{1}}$ has $s$ complex conjugate pairs of eigenvalues and one real negative eigenvalue $\lambda_{1}$.

Define the $m_{\lambda_{1}} \times m_{\lambda_{1}}$ matrix,

$$C_{1} = \begin{bmatrix} W_{1}^{\lambda_{1}} & \cdots & \cr & \ddots & \cr \end{bmatrix}.$$ 

Since $m_{\lambda_{1}}$ is even, the number of odd sized blocks is even, therefore for every $a > 0$, the matrix $J_{1} + C_{1}$ has an even number of size one Jordan blocks corresponding to eigenvalue $\lambda_{1}$, and all other eigenvalues of $J_{1} + C_{1}$ are complex.

By repeat this process for $\lambda_{2}, \ldots, \lambda_{r}$, we can construct the matrix

$$F = \begin{bmatrix} C_{1} & \cdots & \cr & \ddots & \cr C_{r} & 0 & \cr \end{bmatrix}.$$ 

Let $B = QFQ^{-1}$. Then it follows that for every $a > 0$, all negative eigenvalues of $\Phi^{*} + B$ will have Jordan blocks that repeat an even number of times. If $\Phi^{*}$ is nonsingular, then $\Phi^{*} + B = e^{A}$ has real solution $A$ for every $a > 0$. If $\Phi^{*}$ is singular, then $\Phi^{*} + B + aI = e^{A}$ has real solution $A$ for every $a > 0$ sufficiently small. In either case, this implication contradicts the existence of the open set $U$. Thus it must be that some $m_{\lambda_{i}}$ is odd and therefore, $\Phi^{*}$ has at least one negative eigenvalue of odd multiplicity.

\[ \square \]

For a given $d$ such that the associated matrix $\Phi$ has $n$ distinct positive eigenvalues, Corollary 27 (28, or 29, respectively) gives an open set $U \subset \mathbb{R}^{n \times n}$ on which the matrix logarithm exists (exists and is unique, or does not exist, respectively). By the continuity of $G$ at $d$, $G^{-1}(U)$ (the preimage of $U$) is an open set in $\mathcal{D}$ containing $d$ such that every data set in $G^{-1}(U)$ is generated by a real $A$ (a unique real $A$, or no real $A$, respectively). This
concept will be important in Section 3.4, as we seek sets in \( \mathcal{D} \) that define the largest allowable uncertainty about \( d \) such that the inverse problem has a solution (a unique solution, or no solution, respectively).

3.3.2 Companion matrix formulation

Suppose we wish to identify a region of data on which the inverse map \( F^{-1} \) is well defined, i.e., a region \( R \subset \mathcal{D} \) such that for every \( d \in R \) there is a unique parameter \( A \) that defines a linear system (2.3) with a trajectory that generates the data set \( d \). Corollary 28 tells us that, for a data set \( d \) and corresponding fundamental matrix \( \Phi = X_1X_0^{-1} \), the uniqueness of \( A \) on a neighborhood of \( \Phi \) is determined by the eigenvalue structure of \( \Phi \). Since eigenvalue properties are invariant under similarity transformation, we will proceed by first simplifying the form of the matrix.

Let \( \hat{\Phi} \) be the matrix similar to \( \Phi \) that has the form of a companion matrix, i.e.,

\[
\hat{\Phi} = P\Phi P^{-1} = \begin{bmatrix}
0 & 0 & \cdots & 0 & y_1 \\
1 & 0 & \cdots & 0 & y_2 \\
0 & 1 & \cdots & 0 & y_3 \\
\vdots & \ddots & \ddots & \vdots & \vdots \\
0 & 0 & \cdots & 1 & y_n
\end{bmatrix}.
\]

One can think of \( \hat{\Phi} \) as the fundamental matrix for a trajectory for which the data set \( \hat{d} \) is comprised of the vectors of the standard basis of \( \mathbb{R}^n \), together with the data vector \( y = (y_1, y_2, \ldots, y_n)^T \), i.e., \( \hat{\Phi} = G(\hat{d}) \) for \( \hat{d} = (e_1, e_2, e_3, \ldots, e_n, y) \). The matrix \( P \) defines the affine transformation that takes the data \( d \) into the standard (normalized) data \( \hat{d} \), i.e., \( P : \mathbb{R}^n \rightarrow \mathbb{R}^n \) where \( Px_j = e_{j+1}, Px_n = y \). This implies that \( P = X_0^{-1} \) and \( \hat{\Phi} = X_0^{-1}X_1 \).

Companion matrices have several nice properties, one of which is that their characteristic polynomials have a very simple form, given by \( p(t) = y_1 + y_2t + \cdots + y_n t^{n-1} - t^n \). Matrices \( \hat{\Phi} \) and \( \Phi \) have the same eigenvalues; hence, \( \Phi \) has real, positive, distinct eigenvalues if and only if \( p(t) \) has real, positive, distinct roots. A method for determining the number of real distinct roots of a polynomial was presented in [91] using foundational theory from [27]. Given that the roots are real, Descartes’ rule of signs with the criterion that the maximum
number of negative roots is zero gives necessary and sufficient conditions for positivity of the roots, which translate into conditions on $y$. A discussion of this approach is included in Appendix A.

### 3.3.3 Examples

As a first illustrative example, consider the case $n = 2$. The conditions on $y = (y_1, y_2)^T$ so that $\hat{\Phi}$ has 2 distinct positive eigenvalues, obtained from the Yang theorem [91] and Descartes’ rule of signs, are: $y_2^2 + 4y_1 > 0$, $y_2 > 0$, and $y_1 < 0$ (see Appendix A.0.1). They define a region $\hat{R} \subset \mathbb{R}^2$ pictured in Figure 13(a). To gain insight into how these conditions translate into conditions on the data set $(x_0, x_1, \ldots, x_n)$, one can look at the transformation of the region $\hat{R}$ back to a region $R \subset D$ using the inverse transformation $P^{-1}$. One can obtain several different cross-sections of $R$ by fixing two data points from $x_0, x_1, x_2$ and allowing the third to vary. The shaded regions in Figure 13(b)-(d) represent all the possible coordinates the third data point could have such that the data set yields a unique inverse $A$ when the two other data points are fixed as shown.

In case of $n = 3$, the conditions on $y = (y_1, y_2, y_3)^T$ are given in Appendix A.0.2 and the region $\hat{R}$ in $y$-space where all inequalities are satisfied is depicted in Figure 10. Just as in the two-dimensional case, the region is simply connected and unbounded. If we substitute $y_2 = -q_1 y_3^2$ and $y_1 = q_2 y_3^3$, where $q_1$ and $q_2$ are new parameters, we can describe the region using inequalities $y_3 > 0, 0 < q_1 < 1/3, -27q_2^3 + (18q_1 - 4)q_2 - 4q_1^3 + q_1^2 > 0$. The benefit of the new parametrization (which can be introduced for arbitrary dimension) is that it demonstrates that the region $\hat{R}$ consists of curved rays that start at the origin and extend to infinity.

There is an important remark we wish to make while discussing the companion matrix associated with given data, concerning the problem of incomplete data. Since $\hat{\Phi} = X_0^{-1}X_1$ and $\hat{\Phi}$ is determined by the eigenvalues of $\Phi$, for any choice of such eigenvalues, i.e., for any choice of $\hat{\Phi}$, there exists $x_n$ such that the data set $d = (x_0, x_1, x_2, \ldots, x_n)$ is compatible with $\hat{\Phi}$, namely $x_n = X_0y$. In other words, the knowledge of $X_0$, i.e., the knowledge of the first $n$ data points $x_0, x_1, \ldots, x_{n-1}$ for an $n$-dimensional linear dynamical system, does not provide
Figure 9: (a) Region $\hat{R}$ in $y$-space for which $\hat{\Phi} \in \mathbb{R}^{2 \times 2}$ has two distinct positive eigenvalues and hence $\Phi$ has a unique matrix logarithm, which solves the inverse problem. (b)-(d) Regions in data space $D$ for which $\Phi \in \mathbb{R}^{2 \times 2}$ has a unique matrix logarithm, obtained as cross-sections of $R$ formed by fixing two data points as shown. For example, in (b), $x_1, x_2$ are fixed and the shaded region indicates where the coordinates of $x_0$ can lie; note that (d) is strongly related to Figure 13.

any information about the eigenvalues of that system. Thus, we cannot deduce from just $n$ data points whether the observed $n$-dimensional linear system is stable or unstable, whether it is a node or spiral or saddle, or even whether the data are generated by a system with a real parameter matrix. Figure 11 illustrates this point by showing trajectories of linear dynamical systems (2.3) that all share an identical matrix $X_0$ but differ widely in dynamical properties. The same observation can be made about similar cases in which any one of the
Figure 10: Region $\hat{R}$ in $y$-space for which $\hat{\Phi} \in \mathbb{R}^{3 \times 3}$ has a unique matrix logarithm.

$n + 1$ data points $x_0, x_1, x_2, \ldots, x_n$ is missing.

3.4 ANALYSIS OF UNCERTAINTY IN THE DETERMINATION AND CHARACTERIZATION OF INVERSE

Realistic data are never exact, but are subject to uncertainty caused by measurement error, fluctuation in experimental conditions, or variability in experimental subjects. A natural question arises as to how large an uncertainty in the data can be tolerated without altering the properties of the solution to the inverse problem. Several scenarios are of interest; for example,

- The data imply that the inverse system has a stable node. What is the largest uncertainty in the data that ensures the maintained stability of the equilibrium for the inverse system? What is the largest uncertainty that maintains the node property?
- The data imply that the inverse system has oscillations (damped or undamped). What
Figure 11: Trajectories of system (2.3) that share 5 data points in 5 dimensions need not have the same dynamical behaviors. Each panel shows time courses of all 5 components of system (2.3) with \( n = 5 \) for a particular choice of parameter \( A \). Within each panel, each color represents a different component of the system, while the same components share the same color across panels. Five points that are equally spaced in time, through which the trajectories pass in all panels where trajectories exist, are marked with circles. The next equally spaced point is labeled with a star; these points differ across panels and lead to different properties of the inverse problem solution, including non-existence of real parameter matrix for the data in the lower right panel.

is the largest uncertainty that maintains the oscillatory property of the system?

• The inverse does not exist for given data. What is the largest uncertainty for which we
can still rule out the linear model?

The theory developed in Section 3.3 implies that all of the above criteria can be formulated as conditions on the eigenvalues of the perturbed fundamental matrix. For example, a system with a stable node still has a stable node under perturbation if and only if the eigenvalues of the perturbed fundamental matrix remain real, positive, distinct, and smaller than 1. Thus, in principle, we could construct algebraic constraints on the data similar to those given in Section 3.3.2 to define regions of the data space in which data correspond to systems with specific dynamical properties. Generalizing the examples of Section 3.3.3, however, reveals that even in the case of a unique inverse in 4 dimensions, the algebraic constraints are very complex. We will therefore focus instead on the scenario where a specific data set \( d \) is given and derive bounds on the maximal perturbation of \( d \) for which a particular property of the system is conserved.

Note that any affine transformation of the data preserves the eigenvalue structure and hence the existence, uniqueness, and stability of the system with respect to the inverse problem. Thus, the data can be varied in a coordinated fashion to an arbitrary extent without affecting qualitative properties of the inverse. Here, however, we focus on finding limits on uncorrelated perturbations of the data. Let \( C(d, \epsilon) = \times_{i=0}^{n} c(x_i, \epsilon) \) where \( c(z, \epsilon) = \{ \tilde{z} \in \mathbb{R}^n | \max_{1 \leq j \leq n} |\tilde{z}_j - z_j| < \epsilon \} \) for \( z_i \) denoting the components of the vector \( z \). The definition of the neighborhood \( C(d, \epsilon) \) is chosen so that the parameter \( \epsilon \) controls the maximum perturbation \( \Delta x_0, \Delta x_1, \ldots, \Delta x_n \) in any component of the data \( x_0, x_1, \ldots, x_n \), i.e., \( \tilde{d} \in C(d, \epsilon) \) if and only if \( \max_{i,j} |(\Delta x_i)_j| < \epsilon \) where \( |(\Delta x_i)_j| = |\tilde{x}_i)_j - (x_i)_j| \). Neighborhood \( C(d, \epsilon) \) of the data set \( d \in \mathcal{D} \) will be called permissible for some qualitative property of the inverse of \( d \) (such as existence, uniqueness, stability, and so on) if and only if that qualitative property is shared by inverses of all data sets \( \tilde{d} \in C(d, \epsilon) \). The value \( \epsilon > 0 \) is called the maximal permissible uncertainty for some qualitative property of the inverse of \( d \) if and only if \( C(d, \epsilon) \) is a permissible neighborhood of \( d \) for that property and \( C(d, \tilde{\epsilon}) \) is not a permissible neighborhood for that property for all \( \tilde{\epsilon} > \epsilon \).

We begin with an analytical and numerical description of the maximal permissible uncertainty for existence and uniqueness of the inverse \( F^{-1}(d) \) of \( d \).
properties will be described in Section 3.4.4. We derive both lower and upper analytical bounds on maximal permissible uncertainty, describe a numerical procedure for computing the bounds, and then compare the estimates with direct numerical results for several examples.

### 3.4.1 Analytical lower bound

Consider a fixed data set \( d = (x_0, x_1, x_2, \ldots, x_n) \in \mathcal{D} \) such that the associated matrix \( \Phi = X_1X_0^{-1} = [x_1 | \ldots | x_n][x_0 | \ldots | x_{n-1}]^{-1} \) has \( n \) distinct positive eigenvalues. Thus, \( A = F^{-1}(d) \) is unique and there is a neighborhood of \( d \) for which uniqueness persists. For any perturbed data set \( \tilde{d} = (\tilde{x}_0, \tilde{x}_1, \tilde{x}_2, \ldots, \tilde{x}_n) \in \mathcal{D} \) with \( \tilde{x}_i = x_i + \Delta x_i \) define the perturbed data matrices \( \tilde{X}_0 = X_0 + \Delta X_0 = [x_0 | \ldots | x_{n-1}] + [\Delta x_0 | \ldots | \Delta x_{n-1}] \) and \( \tilde{X}_1 = X_1 + \Delta X_1 \) (analogously). Let \( \tilde{\Phi} = \tilde{X}_1\tilde{X}_0^{-1} \) be the fundamental matrix of the perturbed data.

Let \( \epsilon_u \) be the maximal permissible uncertainty in the data \( d \) to ensure the existence of a unique inverse. By definition, for any perturbation of the data with \( \max_{i,j} |(\Delta x_i)_j| < \epsilon_u \) the matrix \( \tilde{\Phi} \) has a unique logarithm \( \tilde{A} \), and for any \( \hat{\epsilon} > \epsilon_u \) there exists a perturbation of the data with \( \epsilon_u < \max_{i,j} |(\Delta x_i)_j| < \hat{\epsilon} \) such that \( \tilde{\Phi} \) does not have a unique logarithm.

A lower bound \( \epsilon_u \) on \( \epsilon_u \) can be obtained by the following result, where \( \|\cdot\| \) denotes a matrix norm that is either the maximum row sum norm \( \|\cdot\|_\infty \) or the maximum column sum norm \( \|\cdot\|_1 \), defined as

\[
\|A\|_\infty = \max_{1 \leq i \leq n} \sum_{j=1}^n |a_{ij}|, \quad \|A\|_1 = \max_{1 \leq j \leq n} \sum_{i=1}^n |a_{ij}|.
\]

**Theorem 30.** Let \( d \in \mathcal{D} \) be such that \( \Phi \) has \( n \) distinct positive eigenvalues \( \lambda_1, \ldots, \lambda_n \). Let

\[ m_1 = \frac{1}{2} \min_{i<j} |\lambda_i - \lambda_j|, \quad m_2 = \min_{1 \leq i \leq n} \{\lambda_i\} > 0, \quad \text{and} \quad \delta_u = \min\{m_1, m_2\}. \]

If \( \epsilon > 0 \) is such that \( \epsilon \leq \epsilon_u := f(\delta_u, d) \), where

\[
f(\delta, d) = \frac{\delta}{n (\delta + 1 + \|A\|) \|S^{-1}\| \|X_0^{-1}S\|^2}, \quad (3.2)
\]

\( \Phi = SAS^{-1}, \) and \( \Lambda = \text{diag}(\lambda_1, \ldots, \lambda_n) \), then for any \( \tilde{d} \in C(d, \epsilon) \), \( \tilde{\Phi} \) has \( n \) distinct positive eigenvalues and hence the equation \( e^{\tilde{\hat{A}}} = \tilde{\Phi} \) has a unique solution \( \tilde{\hat{A}}. \)
The proof of Theorem 30 utilizes several preliminary results that we now present. The first result and its proof make use of Theorems 6.1.1 (Gershgorin Disc Theorem) and 6.3.2 in [42] and the proofs presented therein.

**Theorem 31.** Let \( \Phi \in \mathbb{R}^{n \times n} \) be diagonalizable with \( \Phi = SAS^{-1} \) and \( \Lambda = \text{diag}(\lambda_1, \ldots, \lambda_n) \). Let \( E \in \mathbb{R}^{n \times n} \). If \( \tilde{\lambda} \) is an eigenvalue of \( \Phi + E \), then \( \tilde{\lambda} \in D \), where

\[
D = \bigcup_{i=1}^{n} D_i, \quad D_i = \{ z \in \mathbb{C} : |z - \lambda_i| \leq \|S^{-1}ES\| \}.
\]

Furthermore, if \( \lambda_i \) are all distinct and the discs \( D_i \) are pairwise disjoint, then each \( D_i \) contains exactly one eigenvalue of \( \Phi + E \).

**Proof.** By similarity, \( \Phi + E \) has the same eigenvalues as \( \Lambda + S^{-1}ES \). Denote by \( e_{ij} \) the elements of \( S^{-1}ES \). Then, by the Gershgorin Disc Theorem, the eigenvalues of \( \Lambda + S^{-1}ES \) are contained in the union of the discs

\[
Q_i = \{ z \in \mathbb{C} : |z - (\lambda_i + e_{ii})| \leq \sum_{j=1}^{n} |e_{ij}| \}.
\]

Clearly, each disc \( Q_i \) is contained in the disc

\[
P_i = \{ z \in \mathbb{C} : |z - \lambda_i| \leq \sum_{j=1}^{n} |e_{ij}| \}.
\]

Furthermore, in view of

\[
\sum_{j=1}^{n} |e_{ij}| \leq \max_{1 \leq i \leq n} \sum_{j=1}^{n} |e_{ij}| = \|S^{-1}ES\|_\infty
\]

each disc \( P_i \) is contained in the disc

\[
D_i = \{ z \in \mathbb{C} : |z - \lambda_i| \leq \|S^{-1}ES\|_\infty \}.
\]

Thus, if \( \tilde{\lambda} \) is an eigenvalue of \( \Phi + E \), then \( \tilde{\lambda} \in Q_i \subseteq P_i \subseteq D_i \) for some \( i \) and therefore, \( \tilde{\lambda} \in D \).

The argument for the norm \( \|\cdot\|_1 \) is constructed in a similar fashion by replacing row sums with column sums in the relations above.

If \( \lambda_i \) are all distinct and the sets \( D_i \) are pairwise disjoint, then the discs \( Q_i \) are pairwise disjoint, and Gershgorin Disc Theorem implies that there is exactly one eigenvalue of \( \Phi + E \) in each \( Q_i \) and hence each \( D_i \).
Lemma 32. Suppose the eigenvalues $\lambda_1, \ldots, \lambda_n$ of $\Phi$ are real, positive and distinct. Let $\Phi = SAS^{-1}$ with $\Lambda = \text{diag}(\lambda_1, \ldots, \lambda_n)$ and let $m_1 = \frac{1}{2} \min_{i<j} |\lambda_i - \lambda_j|$, $m_2 = \min_{1 \leq i \leq n}\{\lambda_i\}$, and $\delta = \min\{m_1, m_2\}$. If $\|S^{-1}ES\| < \delta$, then the eigenvalues of $\Phi + E$ are real, positive, and distinct.

Proof. Suppose the eigenvalues $\lambda_1, \ldots, \lambda_n$ of $\Phi$ are real, positive and distinct. Then $\Phi$ is diagonalizable as $\Phi = SAS^{-1}$ with $\Lambda = \text{diag}(\lambda_1, \ldots, \lambda_n)$. Let $R_i$ be the disc

$$R_i = \{z \in \mathbb{C} : |z - \lambda_i| \leq m_1\}$$

and $D_i$ be as in the statement of Theorem 31. Since $\|S^{-1}ES\| < \delta \leq m_1$, it follows that $D_i \subseteq R_i$. The sets $R_i$ are pairwise disjoint, by the definition of $m_1$, so the sets $D_i$ are pairwise disjoint and, by Theorem 31, each $D_i$ contains exactly one eigenvalue of $\Phi + E$. The center of $D_i$ is $\lambda_i \in \mathbb{R}$, so if $D_i$ were to contain a complex eigenvalue of $\Phi + E$, it would also contain its conjugate, which is a contradiction. Thus, the eigenvalues of $\Phi + E$ are real and distinct. Furthermore, the inequality $\|S^{-1}ES\| < \delta \leq m_2$ implies that $D_i \subseteq \{z \in \mathbb{C} : |z - \lambda_i| \leq m_2\} \subseteq \{z \in \mathbb{C} : \text{Re}(z) > 0\}$, and hence the eigenvalues of $\Phi + E$ are all positive. \hfill \Box

Thus, to guarantee that the eigenvalues $\tilde{\lambda}_1, \ldots, \tilde{\lambda}_n$ of $\tilde{\Phi} = \Phi + E$ are real, positive, and distinct, it suffices to choose the perturbation matrix $E$ such that $\|S^{-1}ES\| < \delta$. Theorem 30 provides a lower bound on the largest allowable perturbation of the data points such that this condition holds.

Proof. (Theorem 30) Let $\tilde{d} \in C(d, \epsilon)$ and let $\tilde{\Phi}$ be the associated fundamental matrix as previously defined. Let $E = \tilde{X}_1\tilde{X}_0^{-1} - X_1X_0^{-1}$. Applying the definitions of $\tilde{X}_0, \tilde{X}_1$ yields $E(X_0 + \Delta X_0) = (X_1 + \Delta X_1) - \Phi(X_0 + \Delta X_0)$, which implies that $E = \Delta X_1X_0^{-1} + \Phi\Delta X_0X_0^{-1} - E\Delta X_0X_0^{-1}$ and $S^{-1}ES = S^{-1}\Delta X_1X_0^{-1}S + \Lambda S^{-1}\Delta X_0X_0^{-1}S - S^{-1}ESS^{-1}\Delta X_0X_0^{-1}S$. Therefore,

$$\|S^{-1}ES\| \leq \frac{\|S^{-1}\Delta X_1X_0^{-1}S\| + \|\Lambda S^{-1}\Delta X_0X_0^{-1}S\|}{1 - \|S^{-1}\Delta X_0X_0^{-1}S\|}.$$

After introducing $\|S^{-1}\Delta X\| := \max\{\|S^{-1}\Delta X_0\|, \|S^{-1}\Delta X_1\|\}$, we obtain the following bound on $\|S^{-1}ES\|$ in terms of $\|S^{-1}\Delta X\|$ (provided that $\|S^{-1}\Delta X\|\|X_0^{-1}S\| < 1$):

$$\|S^{-1}ES\| \leq \frac{\|S^{-1}\Delta X\|(1 + \|\Lambda\|)\|X_0^{-1}S\|}{1 - \|S^{-1}\Delta X\|\|X_0^{-1}S\|}. \quad (3.3)$$

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Now, let
\[ q = \frac{\delta}{1 + \|A\|} + 1, \quad \hat{\epsilon} = n\|S^{-1}\|\epsilon \]
and let \( \epsilon \) be as in the statement of the theorem. It follows from \( \epsilon \leq \epsilon U \) that \( \hat{\epsilon} \leq \frac{q-1}{q\|X_0^{-1}S\|} \), or equivalently \( \frac{1}{q} \leq 1 - \hat{\epsilon}\|X_0^{-1}S\| \). The condition \( \hat{d} \in C(d, \epsilon) \) implies that \( |(\Delta x_i)_j| < \epsilon \) for all \( i \in \{1, \ldots, n\}, j \in \{0, \ldots, n\} \), and so \( \|S^{-1}\Delta X\| \leq \|S^{-1}\|\|\Delta X\| \leq \hat{\epsilon} \) (which holds for both \( \|\cdot\|_1 \) and \( \|\cdot\|_x \)). Thus,
\[ \frac{1}{q} < 1 - \|S^{-1}\Delta X\|\|X_0^{-1}S\|. \tag{3.4} \]
Substitution of (3.4) into inequality (3.3) can be used to conclude that \( \|S^{-1}ES\| < \delta_U \). By Lemma 32, \( \hat{\Phi} \) has \( n \) distinct positive eigenvalues.

In the special case where the first \( n \) data points \( x_0, \ldots, x_{n-1} \) are fixed, we can obtain a tighter bound on the size of \( \Delta x_n \). We look for the largest uncertainty \( \epsilon \) of the final data point \( x_n \) so that for any \( \hat{x}_n \in c(x_n, \epsilon) \), \( \hat{d} = (x_0, \ldots, x_{n-1}, \hat{x}_n) \) gives an associated \( \hat{\Phi} \) with \( n \) distinct positive eigenvalues. Fixing the first \( n \) data points implies that \( \Delta X_0 = 0 \) and hence \( E = \Delta X_1 X_0^{-1} \).

**Theorem 33.** Let \( d \in D \) be such that \( \Phi \) has \( n \) distinct positive eigenvalues, and let \( \epsilon > 0 \) satisfy
\[ \epsilon < \max \left\{ \frac{\delta}{\|S^{-1}\|_\infty\|X_0^{-1}S\|_\infty}, \frac{\delta}{n\|S^{-1}\|_1\|X_0^{-1}S\|_1} \right\}. \]
Then for any \( \hat{d} = (x_0, \ldots, x_{n-1}, \hat{x}_n) \) where \( \hat{x}_n \in c(x_n, \epsilon) \), the associated matrix \( \hat{\Phi} \) has \( n \) distinct positive eigenvalues and hence the equation \( \epsilon \hat{A} = \hat{\Phi} \) has a unique solution \( \hat{A} \).

**Proof.** Given fixed \( x_0, \ldots, x_{n-1} \) and \( \hat{x}_n \in c(x_n, \epsilon) \), we have \( \|\Delta X_1\|_\infty = \|[0 \cdots 0 \Delta x_n]\|_\infty = \max_{1 \leq j \leq n} |(\Delta x_n)_j| < \epsilon \), and \( \|\Delta X_1\|_1 < n\epsilon \). If \( \|X_0^{-1}S\|_\infty < n\|X_0^{-1}S\|_1 \), then
\[ \|S^{-1}ES\|_\infty \leq \|S^{-1}\Delta X_1\|_\infty\|X_0^{-1}S\|_\infty < \epsilon\|S^{-1}\|_\infty\|X_0^{-1}S\|_\infty < \delta. \]

In the opposite case,
\[ \|S^{-1}ES\|_1 \leq \|S^{-1}\Delta X_1\|_1\|X_0^{-1}S\|_1 < n\epsilon\|S^{-1}\|_1\|X_0^{-1}S\|_1 < \delta. \]

In both cases, by Lemma 32, \( \hat{\Phi} = \Phi + E \) has \( n \) distinct positive eigenvalues. \( \Box \)
3.4.2 Analytical upper bound

To construct an upper bound on $\epsilon_U$, we only need to provide a technique for constructing a perturbation $\Delta x_0, \Delta x_1, \ldots, \Delta x_n$ for which the corresponding matrix $\tilde{\Phi}$ does not have $n$ distinct real positive eigenvalues. Naively, for any specified diagonal form $\tilde{\Lambda}$ of $\tilde{\Phi}$, we can choose an arbitrary set of eigenvectors $\tilde{S}$, compute $\tilde{\Phi} = \tilde{S}\tilde{\Lambda}\tilde{S}^{-1}$, choose the first data point $\tilde{x}_0$, compute the remaining data points as $\tilde{x}_k = \tilde{\Phi}^k\tilde{x}_0$, and compute the error $\epsilon = \max_{i,j}|(\Delta x_j)_i|$. This procedure can be used to provide a valid upper bound on $\epsilon_U$, but the resulting bound will be too large to be useful. Instead, we suggest the following approach.

Let $\hat{\Phi} = X_0^{-1}\Phi X_0$ be the companion matrix defined in Section 3.3. The vector $y$ (the last column of $\hat{\Phi}$) is uniquely determined by the eigenvalues of $\Phi$. Likewise, we can specify $\tilde{y}$ of the companion matrix $\hat{\Phi} = (\tilde{X}_0)^{-1}\tilde{\Phi}\tilde{X}_0$ by prescribing the eigenvalues of $\tilde{\Phi}$. The vector $\tilde{y}$ satisfies the relation

$$\tilde{y} = (\tilde{X}_0)^{-1}\tilde{x}_n = (X_0 + \Delta X_0)^{-1}(x_n + \Delta x_n),$$

which, using $x_n = X_0y$, implies a formula for the perturbation of $x_n$ in terms of the perturbation of all other data points:

$$\Delta x_n = \Delta X_0\tilde{y} + X_0(\tilde{y} - y).$$

This formula provides a linear constraint on the perturbation of the data in terms of the imposed eigenvalue properties (as represented by vector $\tilde{y}$) that does not require the knowledge of the eigenvector matrix $S$. Let $C(d, \tilde{\epsilon})$ be the smallest neighborhood of $d$ that contains a data set $\tilde{d}$ corresponding to a companion matrix defined by $\tilde{y}$. In view of (3.5), the problem of finding $\tilde{\epsilon}$ can be reformulated as a linear programming problem of minimizing $\epsilon$ while satisfying the constraints

$$w_i = \sum_{j=0}^{n-1} (\Delta x_j)_i\tilde{y}_{j+1} - (\Delta x_n)_i, \quad 1 \leq i \leq n$$

$$-\epsilon \leq (\Delta x_j)_i \leq \epsilon, \quad 1 \leq i \leq n, 0 \leq j \leq n$$
where \( w = X_0(y - \tilde{y}) \). The solution of this problem is given by
\[
\bar{\epsilon} = \max_{\tilde{y}} \frac{|w|}{\|\tilde{y}\|_1 + 1} = \frac{\|X_0(y - \tilde{y})\|_\infty}{\|\tilde{y}\|_1 + 1}. \tag{3.6}
\]
Equation (3.6) can provide an upper bound \( \bar{\epsilon}_U \) on \( \epsilon_u \) for any appropriate choice of \( \tilde{y} \); this approach does not provide an explicit formula for the minimizer of the linear programming problem, however.

If one desires an analytical upper bound together with an explicit formula for the perturbations \( \Delta x_i \) that realize this bound, one can use the inequality
\[
\epsilon_u \leq \min_{\Delta X_0} \max_{0 \leq j \leq n} \{ \|\Delta x_j\|_\infty \} \leq \min_{\Delta X_0} \{ \|\Delta X_0\|, \|\Delta X_0\tilde{y} + X_0(\tilde{y} - y)\| \} \tag{3.7}
\]
where the matrix norm \( \|\cdot\| \) can be either \( \|\cdot\|_\infty \) or \( \|\cdot\|_1 \). Two crude estimates of (3.7) can be obtained by putting \( \Delta X_0 = 0 \), which implies \( \epsilon_u \leq \|X_0(\tilde{y} - y)\| \) or by choosing \( \Delta X_0 \) such that \( \Delta X_0\tilde{y} + X_0(\tilde{y} - y) = 0 \) (for example as \( \Delta X_0 = X_0(y - \tilde{y})\tilde{y}^T/\|\tilde{y}\|_2^2 \)) which yields \( \epsilon_u \leq \|X_0(y - \tilde{y})\tilde{y}^T\|/\|\tilde{y}\|_2^2 \). A more refined approximation is then provided by the following convex interpolation of the two crude estimates: \( \Delta X_0 = \alpha w\tilde{y}^T \) with \( 0 \leq \alpha \leq \|\tilde{y}\|_2^{-2} \) and \( w = X_0(y - \tilde{y}) \). An optimum in (3.7) is reached when
\[
\|\Delta X_0\| = \|\Delta X_0\tilde{y} + X_0(\tilde{y} - y)\|,
\]
which implies
\[
\alpha \|w\tilde{y}^T\| = \|w\|(1 + \alpha \|\tilde{y}\|_2^2)
\]
and hence
\[
\alpha = \frac{\|w\|}{\|w\|\|\tilde{y}\|_2^2 + \|w\|\|\tilde{y}\|_2^2}.
\]
An upper bound on \( \epsilon_u \) is therefore provided by the quantity
\[
\bar{\epsilon}_U = \frac{\|X_0(y - \tilde{y})\|\|X_0(y - \tilde{y})\tilde{y}^T\|}{\|X_0(y - \tilde{y})\tilde{y}^T\| + \|X_0(y - \tilde{y})\|\|\tilde{y}\|_2^2}. \tag{3.8}
\]

The upper bound estimates given above, whether they are obtained as a solution of the linear programming problem (3.6) or using (3.8), depend on the choice of \( \tilde{y} \), i.e., the choice of eigenvalues of the perturbed matrix \( \tilde{\Phi} \). One can refine these bounds by further optimization over all appropriate values of those eigenvalues.
3.4.3 Numerical bound

In addition to finding analytical upper and lower bounds on the uncertainty of the data using the techniques described above, one can also take a numerical approach to estimating $\epsilon_U$. For simplicity in representing the set $C(d, \epsilon)$ graphically, we will focus our discussion on the case of a 2-dimensional system; however, the approach can be extended to $n$-dimensions.

Fix $d = (x_0, x_1, x_2) \in (R^2)^3$. To estimate $\epsilon_U$, we will discretize the surface of $C(d, \epsilon)$ and examine whether each grid point yields a unique inverse. By gradually increasing $\epsilon$ we can find the bound as the largest value of $\epsilon$ for which a grid point fails to give unique inverse. In practice, we surround each data point $x_j$ with a collection $M_j$ of points of equally spaced along the edge of a square with center point $x_j$ and side length $2\epsilon$. Depending on the desired precision, we choose $M_j$ to consist of either 8, 16, or 32 grid points. Then, we pair any point

\[ p_0 \in M_0 \text{ with any points } p_1 \in M_1 \text{ and } p_2 \in M_2 \text{ to define the matrix } \Phi = [p_1 \mid p_2][p_0 \mid p_1]^{-1}. \]

In accordance with Theorem 25, the eigenvalues of $\Phi$ will determine whether the solution to $\Phi = e^A$ is unique.

3.4.4 Analytical bounds for additional properties

Using the results we have obtained so far, we can derive upper and lower bounds on the uncertainty in data that preserves additional qualitative properties of the solution to the inverse problem, as long as these properties can be defined as conditions on the eigenvalues of the matrix $\Phi$. For example, let $d \in D$ be such that $\Phi$ has $n$ distinct real eigenvalues $\lambda_1, \ldots, \lambda_n$ satisfying $0 < \lambda_j < 1$, $j = 1, \ldots, n$. Then, the associated matrix $A$ (with $\Phi = e^A$) has $n$
distinct negative eigenvalues and hence the equilibrium is a stable node. Let $\epsilon_{SN}$ define the maximal permissible uncertainty in the data $d$ under which the equilibrium remains a stable node. A lower bound on $\epsilon_{SN}$ can be obtained using the same argument as in Theorem 30, except with the bound on maximum perturbation of eigenvalues, $\delta_u$, replaced by the quantity $\delta_{SN}$ that guarantees that the perturbed eigenvalues remain real, distinct, and between 0 and 1. Below we present, without proofs, analytical lower bound statements analogous to Theorem 30 for the cases of a stable node and a stable system, as well as for the case in which we require no solution to exist. We leave it to the reader to derive upper bounds using the line of reasoning presented in Section 3.4.2.

**Theorem 34.** Let $d \in D$ be such that $\Phi$ has $n$ distinct positive eigenvalues $\lambda_1, \ldots, \lambda_n$ satisfying $0 < \lambda_j < 1$, $j = 1, \ldots, n$. Let $m_1 = \frac{1}{2} \min_{i < j} |\lambda_i - \lambda_j|$, $m_2 = \min_{1 \leq j \leq n} \{\lambda_j\}$, $m_3 = \min_{1 \leq j \leq n} \{1 - \lambda_j\}$ and $\delta_{SN} = \min\{m_1, m_2, m_3\}$. If $\epsilon > 0$ is such that $\epsilon \leq \epsilon_{SN} = f(\delta_{SN}, d)$ with $f$ as defined in (30), then for any $\tilde{d} \in C(d, \epsilon)$, $\tilde{\Phi}$ has $n$ distinct positive eigenvalues $\tilde{\lambda}_j$ with $0 < \tilde{\lambda}_j < 1$, $j = 1, \ldots, n$, and hence the equation $e^{\tilde{A}} = \tilde{\Phi}$ has a unique matrix solution $\tilde{A}$ for which the origin of (2.3) is a stable node.

To guarantee stability of the equilibrium with respect to all inverse problem solutions without demanding uniqueness of a solution, we require that $d \in D$ be such that the eigenvalues of $\Phi$ are not real negative and satisfy $0 < |\lambda_j| < 1$, $j = 1, \ldots, n$. Then, the associated matrix $A$ (with $\Phi = e^A$) has (possibly complex) eigenvalues with negative real part and thus the equilibrium at the origin is stable. Let $\epsilon_s$ define the maximal permissible uncertainty in the data $d$ such that the equilibrium remains stable. A lower bound on $\epsilon_s$ is obtained in the following result.

**Theorem 35.** Let $d \in D$ be such that the eigenvalues of $\Phi$ are not real negative and satisfy $0 < |\lambda_j| < 1$, $j = 1, \ldots, n$. Let $m_1 = \min_{1 \leq j \leq n} \{1 - |\lambda_j|\}$, $m_2 = \min_j |\lambda_j|$ for all $j$ such that $\text{Re}(\lambda_j) > 0$, $m_3 = \min_j |\text{Im}(\lambda_j)|$ for all $j$ such that $\text{Re}(\lambda_j) < 0$ and $\delta_s = \min\{m_1, m_2, m_3\}$. If $\epsilon > 0$ is such that $\epsilon \leq \epsilon_s = f(\delta_s, d)$ with $f$ as defined in 30, then for any $\tilde{d} \in C(d, \epsilon)$, the eigenvalues of $\tilde{\Phi}$ satisfy $0 < |\tilde{\lambda}_j| < 1$, $j = 1, \ldots, n$ and are not real negative, and hence the equation $e^{\tilde{A}} = \tilde{\Phi}$ has a solution $\tilde{A}$ and every such solution has all eigenvalues with negative real part.
Finally, it is interesting to consider the case of nonexistence of an inverse. In particular, given data $d$ for which a real inverse $A$ does not exist, i.e., data that do not represent the trajectory of any real linear system, what is the greatest amount of uncertainty for which nonexistence of a real solution persists, and hence a linear model should not be considered as a possible mechanism underlying the observed uncertain data? Let $d \in \mathcal{D}$ be such that $\Phi$ has at least one negative real eigenvalue of odd multiplicity, which implies that there is no real matrix $A$ such that $\Phi = e^A$. Let $\epsilon_{\text{DNE}}$ define the maximal permissible uncertainty in the data $d$ under which the inverse problem will be guaranteed to remain without a real solution. A lower bound on $\epsilon_{\text{DNE}}$ is obtained in the following result.

**Theorem 36.** Let $d \in \mathcal{D}$ be such that $\Phi$ has at least one negative real eigenvalue of odd multiplicity. Let the collection of such eigenvalues be denoted by $\lambda_1, \ldots, \lambda_k$, where $\lambda_k$ is the closest to zero. Let $m_1 = \max_{1 \leq i \leq k} (\min_{j (j \neq i)} \frac{1}{2} |\lambda_j - \lambda_i|)$, $m_2 = |\lambda_k|$ and $\delta_{\text{DNE}} = \min \{m_1, m_2\}$. If $\epsilon > 0$ is such that $\epsilon \leq \epsilon_{\text{DNE}} = f(\delta_{\text{DNE}}, d)$ with $f$ as defined in (30), then for any $\tilde{d} \in C(d, \epsilon)$, $\tilde{\Phi}$ has at least one negative eigenvalue of odd multiplicity, and hence the equation $e^{\tilde{A}} = \tilde{\Phi}$ has no real solution.

### 3.5 EXAMPLES FOR TWO-DIMENSIONAL SYSTEMS

In the case of two-dimensional linear systems, one can represent several of the previous results in a more explicit fashion and extend upon stability results to encompass various classifications of the equilibrium. We present these extensions here, along with several numerical examples that can be conveniently depicted in the phase plane.

#### 3.5.1 Regions of existence and uniqueness of the inverse

The criteria in Theorems 24 and 25 are based on the Jordan structure of $\Phi$, which for real $2 \times 2$ matrices can only take a few different forms, and hence can be analyzed completely. Utilizing the relationship between the eigenvalues of a matrix and its trace and determinant, the criteria for existence and uniqueness of the matrix logarithm of $\Phi$ can be fully charac-
characterized by conditions on the trace and determinant of $\Phi$ which, in turn, can be expressed as conditions on the data from which $\Phi$ is constructed. For notational simplicity, let $D = \det \Phi$ and $T = \tr \Phi$.

The analysis is based on the following straightforward corollaries of Culver’s theorems:

**Corollary 37.** Let $\Phi \in \mathbb{R}^{2 \times 2}$. There exists $A \in \mathbb{R}^{2 \times 2}$ such that $\Phi = e^A$ if and only if $D \neq 0$ and any of the following hold:

1. $D > 0$, $T > 0$, and $T^2 \geq 4D$,
2. $T^2 < 4D$,
3. $\Phi = \lambda I$, with $\lambda < 0$.

**Corollary 38.** Let $\Phi \in \mathbb{R}^{2 \times 2}$. There exists a unique $A \in \mathbb{R}^{2 \times 2}$ such that $\Phi = e^A$ if and only if $T > 0$, $T^2 \geq 4D > 0$, and $\Phi \neq \lambda I$ for all $\lambda \in \mathbb{R}$.

The diagram in Figure 13(a) summarizes outcomes with respect to the existence and uniqueness of $A$, based on the trace and determinant of $\Phi$. The label DNE indicates that

![Diagram](image1.png)

Figure 13: (a) Existence and uniqueness of matrix logarithm of $\Phi$ classified in terms of $D$ and $T$. (b) An example of how the existence and uniqueness of $A$ depend on the coordinates of $x_2$, given fixed $x_0, x_1$.

in this region, the matrix logarithm results in complex matrices $A$, whereas we are only interested in data resulting from systems with real parameters ($A \in \mathbb{R}^{2 \times 2}$). $\Phi$ is singular on
the red dashed line that cuts through the DNE region, and therefore no inverse \( A \) exists there. The dotted green boundary represents the case where \( \Phi \) has two real, negative eigenvalues that are equal. Here, \( \Phi \) is either a \( 2 \times 2 \) Jordan block, in which case, \( A \) is a complex matrix, or \( \Phi \) is a negative scalar multiple of the identity matrix, in which case, \( A \) is non-unique and a continuum of real matrices \( A \) exists [17]. The dashed blue boundary demarcates the case where \( \Phi \) has two real, positive eigenvalues that are equal. If \( \Phi \) is a \( 2 \times 2 \) Jordan block, then \( A \) is unique, but if \( \Phi \) is a positive scalar multiple of the identity, then \( \Phi \) has multiple Jordan blocks for a given eigenvalue and hence is non-unique.

Let us now focus on data \( d = (x_0, x_1, x_2) \in \mathcal{D} \), where \( x_0, x_1, x_2 \in \mathbb{R}^2 \) are three data points spaced uniformly in time along the trajectory of the system. Let \( X_0 = [x_0 \mid x_1], X_1 = [x_1 \mid x_2] \), and \( \hat{X} = [x_0 \mid x_2] \). The following two theorems summarize the results obtained above as well as those covering the case in which \( X_0 \) is not invertible.

**Theorem 39.** There exists a 2-dimensional system (2.3) for which the solution map gives the data \( x_0, x_1, x_2 \) spaced uniformly in time if and only if any of the following conditions hold:

1. \( \det X_0 \neq 0, \det X_1/\det X_0 > 0, \det \hat{X}/\det X_0 > 0, \text{ and } (\det \hat{X})^2 > 4 \det X_1 \det X_0, \)
2. \( \det X_0 \neq 0 \text{ and } (\det \hat{X})^2 < 4 \det X_1 \det X_0, \)
3. \( \det X_0 \neq 0 \text{ and } X_1 = \lambda X_0 \text{ with } \lambda < 0, \)
4. \( x_2 = \lambda x_1 = \lambda^2 x_0, \text{ with } \lambda \neq 0. \)

**Theorem 40.** There exists a unique 2-dimensional system (2.3) for which the solution map gives the data \( x_0, x_1, x_2 \) spaced uniformly in time if and only if

\[
\det X_0 \neq 0, \quad \det \hat{X}/\det X_0 > 0, \\
(\det \hat{X})^2 > 4 \det X_1 \det X_0 > 0, \text{ and } X_1 \neq \lambda X_0 \text{ for all } \lambda \neq 0.
\]

The conditions defined in Theorems 39 and 40 define regions in 6-dimensional data space in which the inverse \( A \) exists or exists and is unique. By fixing two of the data points, one can visualize two-dimensional cross-sections of these regions defined by conditions on the third data point. Figure 13(b) shows the outcomes associated with different regions where \( x_2 \) can be located, given example locations of \( x_0 \) and \( x_1 \). The label DNE indicates that for
In this region, the matrix logarithm results in complex matrices $A$, which means there is no real 2-dimensional system (2.3) that gives the data $x_0, x_1, x_2$.

### 3.5.2 Classifying the equilibrium point associated with the inverse

Theorems 39 and 40 give results on the existence and uniqueness of $A$ based on the data. Once we know that a real matrix $A$ exists, additional conditions on the data may be found that define certain important properties of the system. The classification of the equilibrium point at the origin is easily determined by eigenvalues of $A$ and hence can be transformed into conditions on the data. The conditions on the classification of $A$ in terms of properties of $\Phi = e^A$ are given by the following statement, where again, $D = \det \Phi$ and $T = \text{tr} \Phi$:

**Theorem 41.** The equilibrium point $x = 0$ of system (2.3) with matrix $A \in \mathbb{R}^{2\times2}$ is

1. a stable node, if $T > 0$, $T^2 > 4D > 0$, and $1 > D > T - 1$.
2. an unstable node, if $T > 0$ and $T^2 > 4D > 4(T - 1)$.
3. a saddle, if $T - 1 > D > 0$.
4. a stable spiral, if $T^2 < 4D < 4$.
5. an unstable spiral, if $T^2 < 4D$ and $D > 1$.
6. a center, if $D = 1$ and $T^2 < 4$.

The region in Figure 13(a) where $A$ is unique contains all of the systems where the equilibrium point is a saddle or node and the region corresponding to a non-unique $A$ contains all of the systems where the equilibrium point is a spiral or a center (with stars on the boundary between uniqueness and non-uniqueness). The regions described in Theorem 41 are depicted in Figure 14(a). Note that although the theorem as stated translates conditions on $A$ into conditions on $\Phi$, the partitioning it provides can also be used to determine the classification of $x = 0$ for any $A$ derived as the logarithm of a given $\Phi$.

As in the case of Theorems 39 and 40, the conditions of Theorem 41 can be converted to conditions on the data $x_0, x_1, x_2 \in \mathbb{R}^2$. Such conditions were used to construct Figure 14 (b), which describes regions in which the placement of $x_2$, given fixed locations of $x_0$ and $x_1$, yields a system with a particular type of equilibrium and corresponding asymptotic behavior. Note that, consistent with Figure 14, when data lies in the region where it uniquely
specifies $A$, the equilibrium of the resulting system is a saddle or a node, while data that gives non-unique real $A$ yields spirals and centers.

Figure 14: (a) Classification of the origin for (2.3) with $A$ such that $\Phi = e^A$, depicted in terms of conditions on $D = \det \Phi$ and $T = \tr \Phi$. Solid lines form boundaries between regions; dashed lines do not. Note that the origin is a center for (2.3) on the solid line separating stable and unstable spirals. (b) Conditions on the position of $x_2$ to give dynamical systems with specified equilibrium point types when $x_0$ and $x_1$ are fixed. Solid lines form boundaries between regions; dashed lines do not.

### 3.5.3 Bounds on maximal permissible uncertainty

In this section we will illustrate the dependence of the maximal permissible uncertainty on the data and the property being maintained.

**Example 1. Differences between analytical bounds and numerical estimates of maximal permissible uncertainty.** Let $d = (x_0, x_1, x_2) = ((10, 2)^T, (6.065, -4.44)^T, (7, -10)^T)$, which are points that are equally spaced in time on a trajectory of the system (2.3) with

$$A = \begin{bmatrix} -0.6724 & -0.7201 \\ -0.8610 & -0.0244 \end{bmatrix}.$$ 

Since $\Phi = e^A$ has two distinct positive eigenvalues, $A$ is the unique matrix that yields the data $d$. For this data set, the direct numerical estimate of the maximum permissible
uncertainty for uniqueness is $\tilde{\epsilon}_u = 1.075$. The analytical lower bounds on $\epsilon_u$ are substantially smaller than the numerical bound, namely $\epsilon_u^\infty = 0.083$ and $\epsilon_u^1 = 0.149$ (where $\epsilon_u^\infty$ is found by applying the norm $\|\cdot\|_{\infty}$ and $\epsilon_u^1$ by using the norm $\|\cdot\|_1$ in Theorem 30).

The analytical upper bound depends on the choice of eigenvalues for the perturbed matrix $\tilde{\Phi}$. Using (3.8) and choosing a perturbed matrix with one zero eigenvalue and the second eigenvalue equal to the average of eigenvalues of $\Phi$ yields the upper bound $\tau_u = 1.245$. Optimization over the value of the second eigenvalue yields a better estimate $\tau_u = 1.078$, which is essentially identical to the numerical upper bound. Choosing a perturbed matrix with identical eigenvalues equal to the average of eigenvalues of $\Phi$ leads to $\tau_u = 1.841$ and optimization over the position of the double eigenvalue gives the upper bound $\tau_u = 1.570$. Using the linear programming estimate of (3.6) gives the same bounds as the zero eigenvalue choice of perturbation. Choosing a perturbed matrix with identical eigenvalues equal to the average of eigenvalues of $\Phi$ and solving (3.8) yields $\tilde{\epsilon}_u = 1.224$ and optimization over the position of the double eigenvalue gives $\tilde{\epsilon}_u = 1.140$. A summary of the optimal bounds is given in the second column of Table 1.

**Example 2. Dependence of maximal permissible uncertainty on $x_2$.** As in Example 1, let $x_0 = (10, 2)^T$, $x_1 = (6.065, -4.44)^T$. Theorem 41 defines regions in $\mathbb{R}^2$ that specify the nature of the equilibrium based on the location of the last data point $x_2$. The boundaries of the regions are shown in Figure 15(a). We select a sample point $x_2$ from each labeled region, and in each case, we compute various estimates of the maximal permissible uncertainty $\epsilon$ to preserve the corresponding property. We depict each uncertainty by outlining in 15(b) the square-shaped sets $c(x_0, \epsilon), c(x_1, \epsilon)$, and $c(x_2, \epsilon)$ in the phase plane $\mathbb{R}^2$, which can be interpreted as follows: given any $\tilde{x}_0 \in c(x_0, \epsilon)$, $\tilde{x}_1 \in c(x_1, \epsilon)$, and $\tilde{x}_2 \in c(x_2, \epsilon)$, the matrix $A$ that yields the data $d = (\tilde{x}_0, \tilde{x}_1, \tilde{x}_2)$ has the appropriate property. Table 1 summarizes the location of $x_2$, the property being preserved, and the best estimate of $\epsilon$ in each case. As can be seen in Figure 15, it appears that the proximity of $x_2$ to the boundary of the region in which the desired property holds impacts the size of the resulting $\epsilon$.

**Example 3. Dependence of maximal permissible uncertainty on the choice of solution property.** Consider the data set $d$ with $x_0, x_1$ as in Examples 1 and 2 and with fixed $x_2 = (3.6, -4.3)^T$, corresponding to a dynamical system with a stable node at the origin. The
maximal permissible uncertainty of the data depends on what property we require to be preserved. In this case we can choose between uniqueness, the stable node property, or stability. If we want to guarantee a unique solution to the inverse problem, we find that \(\tilde{\epsilon}_U = 0.072\). This is quite small due to the proximity of \(x_2\) to the border of the stable spiral region where \(A\) is non-unique, as seen in Figure 16(a). Note that \(c(x_2, \tilde{\epsilon}_U)\) does not extend all the way out to the boundary of the stable spiral region, because these boundary lines are derived with \(x_0\) and \(x_1\) fixed, but we allow uncertainty in all three data points. For preservation of the stable node property, we observe that in this case \(\tilde{\epsilon}_{SN} = \tilde{\epsilon}_U\); however, this relation does not hold universally. It would not be true, for example, if \(x_2\) were located within
the stable node region but closer to the saddle region where $A$ is unique (e.g., see Example 4 below). If we want to ensure that the data lie on a trajectory that converges to the origin (i.e., preserve the stability of the system), we find that the maximal permissible uncertainty is $\tilde{\epsilon}_s = 0.647$, which is significantly larger than $\tilde{\epsilon}_u$. Thus we can guarantee stability for larger uncertainty in the data than what is needed to preserve the uniqueness of solutions. Figure 16(a) depicts the data in the phase plane and the numerically computed maximal permissible uncertainties for the uniqueness, stable node, and stability properties. Within the set $C(d, \tilde{\epsilon}_s)$, any choice of data will lie on a stable trajectory, however, the uniqueness of the inverse problem may not be preserved. In 16(b) we illustrate two different data sets contained in $C(d, \tilde{\epsilon}_s)$; one data set on a stable node trajectory (red), corresponding to a unique $A$, and a second data set belonging to a stable spiral trajectory (blue), where $A$ is not unique.

**Example 4. Dependence of maximal permissible uncertainty on the choice of solution property.** Consider the data set $d$ with $x_0, x_1$ as in Examples 1-3 and with $x_2 = (5, -5.6)^T$, which corresponds to a stable node equilibrium. Again, the maximal permissible uncertainty of the data depends on what property we require to be preserved. Here we find that $\tilde{\epsilon}_{ss} = 0.251$. 

### Table 1: Best Estimates of $\epsilon_X$, $X \in \{ SN, U, S, DNE \}$, for Example 2

<table>
<thead>
<tr>
<th>Property (X)</th>
<th>Stable Node (SN)</th>
<th>Unique (U)</th>
<th>Stable (S)</th>
<th>Nonexistence (DNE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_2$</td>
<td>(3.68, -3.46)</td>
<td>(7, -10)</td>
<td>(3.1, -5.5)</td>
<td>(3.6)</td>
</tr>
<tr>
<td>$\tilde{\epsilon}_x$ Numerical estimate</td>
<td>0.216</td>
<td>1.075</td>
<td>0.519</td>
<td>3.055</td>
</tr>
<tr>
<td>$\epsilon_x$ by Theorem 30</td>
<td>0.059</td>
<td>0.149</td>
<td>0.207</td>
<td>0.609</td>
</tr>
<tr>
<td>$\tilde{\epsilon}_x$ by (3.6)</td>
<td>0.217*</td>
<td>1.078†</td>
<td>0.519‡</td>
<td>3.059#</td>
</tr>
<tr>
<td>$\tilde{\epsilon}_x$ by (3.8)</td>
<td>0.251*</td>
<td>1.078†</td>
<td>0.665‡</td>
<td>3.683†</td>
</tr>
</tbody>
</table>

* $\lambda_1 = \lambda_2$, † $\lambda_1 = 0$, ‡ $\lambda_2 = 1$, # $\lambda_2 \to \infty$
\(\bar{\varepsilon}_s = 0.091\). If we relax the constraint that \(A\) be a stable node, and just require that it is unique, then unlike the previous example, we find a larger maximal permissible uncertainty, \(\bar{\varepsilon}_U = 0.487\). The data \(d\) and the numerically computed \(\bar{\varepsilon}_{SN} = \bar{\varepsilon}_s\) and \(\bar{\varepsilon}_U\) are pictured in the phase plane in Figure 17(a). Any data set \(\tilde{d} \in C(d, \bar{\varepsilon}_s)\) will be generated by a unique \(A\); however, we find that the trajectories on which the data lie may have vastly different behavior. For example, \(d\) as previously defined and \(\tilde{d} = (x_0, x_1, (3.1, -4.3)^T)\) both belong to \(C(d, \bar{\varepsilon}_U)\); however, \(d\) lies on a trajectory converging to the origin and \(\tilde{d}\) belongs to an unstable trajectory as shown in Figure 17(b).

**Example 5. Analytical lower bound for uncertainty in the last data point.** Now consider
Figure 17: (a) Numerical estimates $\tilde{\epsilon}_S = \tilde{\epsilon}_{SN}$ (red) and $\tilde{\epsilon}_U$ (blue) for the data depicted. (b) Stable trajectory (red) through the data $d$ and unstable trajectory (blue) through perturbed data $\tilde{d} \in C(d, \epsilon_U)$.

In the case where the data points $x_0$ and $x_1$ are fixed at the same values as in Examples 1-4 and we seek to find the maximum uncertainty $\epsilon_U$ allowed in the last data point $x_2$ such that for any $\tilde{x}_2 \in c(x_2, \epsilon_U)$, there is a unique $A$ that gives the data $\tilde{d} = (x_0, x_1, \tilde{x}_2)$ at equally spaced time points. Let $x_2 = (3.679, -3.459)^T$. We find the numerical estimate $\tilde{\epsilon}_U = 0.423$ and the analytical estimates $\epsilon_U = 0.07$ and $\epsilon_U^\infty = 0.187$ (using Theorem 33). In this case, the difference between the numerical and analytical lower bounds is much improved from Example 1, and the lower bound, which is the only easily computable bound in higher dimensions, becomes quite useful. Pictured in Figure 18 are the boundaries of the sets $c(x_2, \epsilon_U)$ (red) and $c(x_2, \epsilon_U^\infty)$ (green) along with the functions dividing the plane into regions where $x_2$ may lie such that the parameter $A$ that produces that data is unique or non-unique. As expected, we observe that $c(x_2, \epsilon_U)$ and $c(x_2, \epsilon_U^\infty)$ are contained in the region corresponding to a unique $A$. 
Figure 18: The square regions surrounding the data point $x_2$ represent numerical (red) and analytical (green) estimates of the maximum uncertainty in that data point that allows for a unique inverse.

3.6 REMARK ON NONUNIFORM SPACING OF DATA

In the results presented so far we have assumed that the data available about the system (2.3) were spaced uniformly in time, i.e., that $d$ contains the data points $x_0, x_1, x_2, \ldots, x_n \in \mathbb{R}^n$ where $x_j = x(j; A, b) = e^{Aj}b$. In this section we shall make few remarks on how the results can be extended to the case in which data are spaced non-uniformly, with the restriction that the sampling times are still integer multiples of some $\Delta t$, assumed without loss of generality to be equal to 1. Such situations occur frequently for medical data, which are usually collected more frequently during the early course of a disease and less frequently during recovery.

The following lemma relates the eigenvalues of $\Phi = e^A$ to the data $d = (x_{j_0}, x_{j_1}, x_{j_2}, \ldots, x_{j_n})$:

Lemma 42. Assume that for $i = 0, \ldots, n$, $x_{j_i} = \Phi^{j_i}b$ where $j_i$ are integers such that $0 = j_0 < j_1 < \cdots < j_n$. Let $X_0 = [x_{j_0} \mid \ldots \mid x_{j_{n-1}}]$ and assume that $X_0$ is invertible and $\Phi$ is
nonsingular. Let \( y = X_0^{-1} x_{j_n} \) be a vector with entries \( y_1, y_2, \ldots, y_n \). Then \( \lambda \) is an eigenvalue of \( \Phi \) only if \( \lambda \) is a root of the polynomial

\[
\lambda^{j_n} - y_n \lambda^{j_{n-1}} - \cdots - y_2 \lambda^{j_1} - y_1 = 0.
\] (3.9)

Proof. Since \( y = X_0^{-1} x_{j_n} \), it follows that \( x_{j_n} = X_0 y \), i.e.,

\[
\Phi^{j_n} b = \sum_{i=0}^{n-1} y_{i+1} \Phi^{j_i} b. \tag{3.10}
\]

The assumption that \( X_0 \) is invertible is equivalent to the statement that the vectors \( \{x_{j_i}\}_{i=0}^{n-1} \) are not confined to a proper subspace of \( \mathbb{R}^n \) and hence, if \( x_{j_i} = \Phi^{j_i} b = e^{A_{j_i}} b \), \( b \) is not confined to a proper \( \Phi \)-invariant subspace of \( \mathbb{R}^n \). Thus \( b \), when decomposed in the basis of ordinary and generalized eigenvectors of \( \Phi \), has a nonzero component along the ordinary or generalized eigenvector for every eigenvalue of \( \Phi \). Let \( v \) be the eigenvector corresponding to the eigenvalue \( \lambda \) of \( \Phi \). If \( b \) has a nonzero component along \( v \), then (3.10) implies that

\[
\lambda^{j_n} v = \sum_{i=0}^{n-1} y_{i+1} \lambda^{j_i} v,
\]

which proves the statement. If the component of \( b \) along \( v \) is zero but \( b \) has a nonzero component along \( u \) such that \((A - \lambda I)^{k-1} u \neq 0 \) and \((A - \lambda I)^{k} u = 0\), then (3.10) implies that

\[
\lambda^{j_n+k} u = \sum_{i=0}^{n-1} y_{i+1} \lambda^{j_i+k} u,
\]

which for nonzero \( \lambda \) reduces to the previous case. \( \square \)
Lemma 42 is employed in solving the inverse problem as follows: First, the vector \( y \) is computed from the data \( d = (x_{j_0}, x_{j_1}, x_{j_2}, \ldots, x_{j_n}) \). Second, all roots of the polynomial (3.9) are found using numerical techniques. Third, a combination of \( n \) distinct roots is chosen from the collection of roots. Fourth, the companion matrix \( \Phi \) of \( \Phi \) is formed from the chosen roots and a set of vectors \( z_{j_i}, i = 0, 1, \ldots, n-1 \) is computed by taking \( z_0 = e_1 \) and \( z_{j_i+1} = \Phi^{j_{i+1}-j_i} z_{j_i} \). Finally, the matrix \( \Phi \) is found as \( \Phi = P^{-1} \hat{\Phi} P \) where \( P \) is the (unique) \( n \times n \) matrix such that \( P x_{j_i} = z_{j_i} \) for \( i = 0, 1, \ldots, n-1 \).

The procedure outlined above can find any matrix \( \Phi \) that is robust, in the sense of Corollaries (27)-(29). The condition that \( X_0 \) be invertible (i.e., \( b \) not be confined to a proper \( \Phi \)-invariant subspace) is essential for identifiability of \( \Phi \), as we have already observed in Chapter 2 for the special case of \( j_i = i \). If the indices \( j_i \) differ by more than 1 then the polynomial (3.9) has more roots than the matrix \( \Phi \) has eigenvalues. Arbitrary combinations of such roots will lead to different alternative matrices \( \Phi \) and hence to non-uniqueness of solutions of the inverse problem. By an appropriate choice of the roots that make up the eigenvalues of \( \Phi \), one may be able to control the properties of the matrix \( \Phi \) and, in turn, the existence, uniqueness, and stability properties of the parameter matrix \( A \) of the system (2.3).

3.7 DISCUSSION

We have analyzed the inverse problem for linear dynamical systems, i.e., the problem of finding the value of the parameter matrix for which a linear system generates a given discrete data set consisting of points equally spaced in time on a single trajectory. Our results establish regions in data space that give solutions with particular properties, such as uniqueness or stability, and give bounds on the maximal allowable uncertainty in the data set that can be tolerated while maintaining these characteristics.

Three types of bounds on uncertainties are presented: analytical lower bounds, below which properties are guaranteed to hold for all perturbations of data; analytical upper bounds, which provide proven perturbations of data for which properties are guaranteed
to be lost; and numerical bounds, derived from direct sampling of data points. Our results indicate that the upper bounds, when optimized over all potential eigenvalues, provide excellent agreement with the numerical estimates. The numerical methods are hypothetically applicable to systems of arbitrary size; however, the combinatorial problem of pairing together all possible data points along grid points can pose a challenge as the dimension of the system increases. Similarly, the computation of the analytical upper bound (via (3.6) or (3.8)) requires optimization that becomes computationally expensive for larger systems.

Although the analytical lower bound significantly underestimates the maximal permissible uncertainty, it provides a bound that is immediately accessible for systems of higher dimension, without increased computation. Since we focused on the derivation of these bounds, the question of how these bounds scale with system size remains open for future investigation. Furthermore, in this work, we have considered only random perturbations of the data matrix $\Phi$. Due to the special construction of the matrix $\Phi$, it may be possible to improve the analytical lower bound by considering structured matrix perturbations. Many results have been established concerning the bounds on eigenvalues for structured perturbations of matrices [42, 39] that may prove useful in this effort.

A variety of earlier works considered identification of linear systems or parameter matrices from discrete data. Allen and Pruess proposed a method for approximating $A$ in system (2.3) from a discrete collection of data points [2]. Their approach begins by defining an approximating function for the data (e.g., a cubic spline approximation), and they use equally spaced points along this curve to compute a matrix $\hat{A}$ that approximates the true parameter matrix $A$. A key distinction between their work and the initial analysis presented here is that they use points on an approximation of the trajectory, while we assume that the data represent exact points on the actual trajectory; their results also do not treat uncertainty in data.

In other past work, Singer and Spilerman investigated the problem of identifying the matrix $Q$ in the Markov model $P' = Q P$ where $P$ and $Q$ are $n \times n$ matrices [73]. They derive conditions for $P = e^Q$ to have a unique solution. Their results are consistent with the findings of Culver, but with additional constraints to account for the requirement that the model is a continuous time Markov structure. They additionally comment on the case of
identifying $Q$ from noisy observations and suggest exploring in a neighborhood of $P$ in order to detect non-uniqueness of the matrix logarithm through observations on the eigenvalues of the matrices in this neighborhood.

It is of interest to note that much work has been done in determining the maximal allowable uncertainty in the parameter matrix $A$ such that the solution to (2.3) remains stable [39]. This well known bound is called the stability radius. In our investigation of the inverse problem, $\epsilon_s$ has an analogous meaning, but we quantify the uncertainty in the data space rather than in parameter space.

Our results also include bounds on regions of data space where the inverse problem cannot be solved. The utility of such results is that they can provide an approach for model rejection. That is, suppose we have a data set $d$ acquired from measurements of some physical phenomenon that we believe can be modeled with a linear system of differential equations. Perhaps it is known that the measurement error for any data point $x_i$ is approximately given by $\epsilon$. If we find that there is no real matrix $A$ that yields the collected data $d$ and further find that $\epsilon_{d_{\text{DNE}}} > \epsilon$, then we can conclude with certainty that the data cannot come from a system that can be modeled with a linear system of differential equations, and thus we can reject the linear model.

Our work is also related to the important problem of determining identifiability in parameter estimation. The connection to identifiability is apparent if we consider the set in parameter space defined by $F^{-1}(C(d, \epsilon_u))$. On this set we have that $F^{-1}(d_1) \neq F^{-1}(d_2)$ implies $d_1 \neq d_2 \in C(d, \epsilon_u)$, which is to say that two distinct parameter sets must yield distinct data. So, $F^{-1}(C(d, \epsilon_u))$ defines a set in parameter space on which the model is identifiable.

In the next chapter, important aspects of parameter estimation from a collection of single trajectory data will be investigated. In this practical setting, we will study improvements to Monte Carlo techniques used in Bayesian parameter estimation.
4.0 THE JACOBIAN PRIOR IMPROVES PARAMETER ESTIMATION
WITH THE METROPOLIS-HASTINGS ALGORITHM

4.1 INTRODUCTION

Once a model structure is fixed and experimental data is collected, the problem of determining the unknown model parameters must be confronted. In a Bayesian inference approach to parameter estimation, a posterior distribution of parameters which describes the probability of a parameter governing the system given the available data, is sought. Markov chain Monte Carlo methods provide a means to sample this distribution. One widely used computational method which employs this approach is the Metropolis-Hastings algorithm. Implementation of this algorithm for Bayesian parameter estimation requires the prescription of a prior density of parameters, which reflects any previously known information about the parameters. The choice of a prior may greatly impact the posterior obtained from the algorithm and has been a topic of debate among practitioners [10, 32, 68, 74]. In this chapter, we introduce a new informative prior which does not rely on previous information about the parameters alone, but exploits knowledge of the fixed model structure.

The presentation of the chapter is as follows: In the first section, the notation for the models of interest and other important constructs that will be used in this chapter are defined. In Section 4.2.2, background on Bayesian inference for parameter estimation is discussed, followed by an introduction to the Metropolis-Hastings algorithm. In Section 4.3, we present the theoretical derivation of the Jacobian prior. We then work to systematically analyze the accuracy of the posteriors obtained using this newly proposed prior in the Metropolis-Hastings algorithm. In Section 4.4.1, the known solution to systems of linear differential equations is exploited and the analysis of Chapter 3 is utilized in order to define the
parameter density explicitly. This provides the exact solution to the parameter estimation problem as a means for comparison to the posteriors obtained from the Metropolis-Hastings algorithm. Two different systematic approaches for analyzing the accuracy of the posteriors are presented. Through a series of several examples, we find that using the Jacobian prior results in posteriors that closely match the parameter density and frequently yields more accurate results than with other commonly used priors. In Section 4.4.2, we conduct a similar analysis of comparison between the parameter density and the computed posteriors, instead for a nonlinear system of differential equations. The chapter concludes with a discussion in Section 4.5.

4.2 PRELIMINARIES

4.2.1 Model and notation

The model of interest is formulated as an initial-value problem for a system of ordinary differential equations,

\[
\begin{align*}
\dot{x}(t) &= g(x(t), \Lambda) \\
x(0) &= b.
\end{align*}
\]

In equation (4.1), \(x(t) \in \mathbb{R}^n\) is the state of the system at time \(t\), \(\Lambda\) is a vector of parameters, and \(b \in \mathbb{R}^n\) is the initial condition. The initial condition or a subset of its components may be unknown and thus considered as parameters. Then, the model parameters, denoted by \(a \in \mathbb{R}^p\), is a vector comprised of \(\Lambda\) and any parameter elements of \(b\). For notational simplicity, the parameter space will be called \(\mathcal{A}\).

We shall assume that model (4.1) is well posed and the solution (or trajectory), denoted by \(x(t; \Lambda, b)\), is unique for all \(\Lambda\) and \(b\). We assume that data for all of the state variables is available and denote by \(\mathcal{Y}\), the data space consisting of a set of \(m\)-tuples \(y = (x_1, x_2, \ldots, x_m)\) of points \(x_j \in \mathbb{R}^n\), representing observations of the system at times \(t_1, t_2, \ldots, t_m\). Observations of the trajectory \(x(t; \Lambda, b)\) at times \(t_1, t_2, \ldots, t_m\) yields an element of \(\mathcal{Y}\), namely an \(m\)-tuple of points \(x_j = x(t_j; \Lambda, b) \in \mathbb{R}^n, \ j = 1, \ldots, m\). In this work, we will require \(p = nm\). The
term *solution map* will refer to the map $F : A \rightarrow \mathcal{Y}$ from parameter space to data space, defined as $F(a) = (x_1, x_2, \ldots, x_m) = y$ for a choice of $\{x_j\}$ sampled from $x(t; A, b)$.

$Y$ will denote the available data for the system which will be used in the inverse problem of determining the model parameters. In the case of a collection of single trajectory data, $Y = \{y^1, y^2, \ldots y^N\}$ with $y^j \in \mathcal{Y}$ may represent $N$ repeated observations of the system or $N$ subjects from which experimental measurements are drawn. Alternatively, $Y$ may be provided as a set of mean values $\{\bar{x}_1, \ldots, \bar{x}_m\}$ and standard deviations $\{\sigma_1, \ldots \sigma_m\}$, or in an idealized setting, $Y$ may be a density on the data space, denoted by $Y = \nu(y)$.

A number of approaches have been developed for parameter estimation of differential equation models. In traditional parameter fitting techniques, a cost function, which quantifies the agreement between the measured data and the model predicted values of the output, is constructed. For example, if the data $Y$ are assumed to be normally distributed random variables with mean values $\{\bar{x}_1, \ldots, \bar{x}_m\}$ and standard deviations $\{\sigma_1, \ldots \sigma_m\}$, then the cost function may be represented as a sum of squared residuals between the measured data and the model trajectory:

$$E(a, Y) = \sum_i \sum_j \frac{(x_j(t_i, a) - \bar{x}_{i,j})^2}{2\sigma_{i,j}^2}.$$  

The solution to the parameter estimation problem is the parameter $a^*$ which minimizes the cost function. The vector $a^*$ is called the maximum likelihood estimate of the parameters. One commonly employed algorithm for minimizing the cost function is the Levenberg-Marquardt scheme [51].

Another parameter estimation approach is the multiple shooting algorithm. In this method, the initial value problem is converted to multiple boundary value problems that are solved for segments of the full trajectory [12, 85]. This method is particularly useful for stiff or chaotic systems. Another popular approach is the Kalman filter technique. Here, a recursive algorithm updates the parameters by minimizing the discrepancy between the model predicted output and the data as successive data points are incorporated [49]. There are several variations and extensions of the Kalman filter for nonlinear systems [87].

Each of the methods discussed above produces a single parameter vector as the solution to the inverse problem. Alternatively, probabilistic approaches, such as Bayesian inference, output a distribution of parameters rather than a unique set of parameters. Such an approach
may be especially relevant in biological modeling. In this setting, the model parameters may represent biological quantities that naturally vary from individual to individual. In this case, the parameter distribution represents parameter variability between individuals in a population. Together, the parameter distribution and the system of differential equations form a collection of models often referred to as an ensemble. A discussion of ensemble modeling for biological systems may be found in [77]. The objective is then to find the best approximation to the density $\rho(a)$ that describes the parameter distribution.

### 4.2.2 Bayesian inference for parameter estimation

In a Bayesian inference approach to parameter estimation, one aims to approximate the parameter density $\rho(a)$ with a posterior density $\rho(a|Y)$, which makes use of the available data. Given a fixed model structure and the data $Y$, the posterior density of the parameters $\rho(a|Y)$ quantifies the probability that the system is governed by the parameters $a$ given the data $Y$. Fundamental theory of Bayesian inference and its applications to parameter estimation are discussed in [33, 74]. Bayes’ Theorem provides a crucial relation between the unknown posterior density and a known likelihood function $L(Y|a)$ and prior density $\pi(a)$.

**Bayes’ Theorem:**

$$
\rho(a|Y) = \frac{L(Y|a)\pi(a)}{\int L(Y|a)\pi(a)da}.
$$

(4.2)

The likelihood function $L(Y|a)$ quantifies the deviation of the model with parameters $a$ from the available data, $Y$. The function $L(Y|a)$ (with $a$ as the dependent variable and $Y$ fixed) is defined by the statistical model of the data (where the error comes from and what is the distribution of the error). For example, suppose the data observations $x_i$ are assumed to be normally distributed random variables, and $Y$ is provided as a set of mean observation values $\bar{x}_1, \ldots, \bar{x}_m$ and standard deviations $\{\sigma_1, \ldots, \sigma_m\}$. If the errors in the data measurements are independent and identically distributed, then the likelihood function can be defined as

$$
L(Y|a) = \prod_i L_i(\bar{x}_i|a)
$$

where,

$$
L_i(\bar{x}_i|a) = \prod_j \frac{1}{\sigma_{i,j} \sqrt{2\pi}} e^{-\frac{(x_j(t_i,a)-\bar{x}_i)^2}{2\sigma_{i,j}^2}}.
$$
Alternatively, in an idealized setting, the data $Y$ could be given as a density function $\eta(y)$. The solution to model (4.1) is unique and due to the absence of error in the forward model solution, it holds that the likelihood can be defined as $L(\eta|a) = \eta(F(a))$ ([79], page 35). Several other likelihood functions may be defined based upon properties of the data available, for example by taking into account covariances in the data. Thoughtful selection of the likelihood function is important because the data informs the posterior only through the likelihood.

The prior density $\pi(a)$ reflects any information that is known about the parameters before data is considered. This information may include bounds on the values of the parameters which are obtained from literature (e.g. biological experiments) or from qualitative analysis of the system (e.g. analysis of existence and stability of equilibria).

A noninformative prior (also commonly referred to as an objective or flat prior) may be used when no information about the parameter values is known. This prior is flat relative to the likelihood, thus minimizing its impact on the posterior distribution. A uniform density posed on the support of the parameters is a commonly used noninformative prior [74]; for example $\pi(a) = \chi_{[0,\infty)}(a)$ is a uniform prior density for a positive parameter (this is an improper prior because its integral is infinite). Jeffreys prior is an objective prior based on the Fisher information matrix [45]. This prior does not vary much over the region in which the likelihood is significant and does not take on large values outside that range, thus satisfying the local uniformity property [29]. It also has the property of scale invariance, meaning that it is invariant under reparameterization by injective transformations. An informative prior can be used when characteristics of the parameters are known a priori. Unlike noninformative priors, this prior is not dominated by the likelihood.

The choice of a prior is a topic of debate among practitioners of Bayesian inference [10, 32, 68, 74]. The possible dangers of employing an informative prior are illustrated by Smith in [74]. He shows that a poorly chosen informative prior can degrade the accuracy of the posterior far more than an objective prior and recommends that unless good previous information is known, an objective prior should be used. This chapter represents a contribution to the investigation of prior density selection by introducing a new informative prior which is specifically relevant for Bayesian inference in the setting of parameter estimation.
4.2.3 Metropolis-Hastings algorithm

In practice, a sample of the posterior density \( \rho(a|Y) \) can be obtained using Markov chain Monte Carlo (MCMC) techniques. In the setting of Bayesian inference, the Metropolis-Hastings algorithm constructs a Markov chain \( a^1, a^2, \ldots, a^M \) with the posterior density as its limiting distribution [35, 61]. This approach to parameter estimation is widely used in practice and the Metropolis-Hastings algorithm has been referred to as one of the top ten algorithms of the twentieth century [74]. Accessible introductions to the algorithm can be found, for example, in [74, 81]. In each iteration of the algorithm, \(\rho(\hat{a}|Y)\) will be computed using Theorem 4.2 with the normalization constant omitted, as will be discussed below.

Given the model (4.1) and the available data \(Y\), the algorithm is formulated as follows:

1. Choose an initial parameter value \(a^1\) satisfying \(\rho(a^1|Y) > 0\).
2. For \(k = 1, \ldots, M\),
   (i) Propose a new parameter value \(\hat{a}\) from a proposal (jumping) distribution \(q(\hat{a}|a^k)\).
   (ii) Using \(\hat{a}\), solve the ODE system (4.1) and compute \(\rho(\hat{a}|Y) = L(Y|\hat{a})\pi(\hat{a})\).
   (iii) Set
       \[
       a^{k+1} = \begin{cases} 
       \hat{a}, & \text{with probability } \min \left\{ 1, \frac{\rho(\hat{a}|Y) q(a^k|\hat{a})}{\rho(a^k|Y) q(\hat{a}|a^k)} \right\} \\
       a^k, & \text{otherwise}
       \end{cases} 
       \]
       (\(\hat{a}\) accepted)
       \[
       a^{k+1} = a^k, \quad \text{otherwise}
       \]
       (\(\hat{a}\) rejected).

The proposal distribution \(q\) for selecting \(\hat{a}\) is based only on the previous parameter \(a^k\), thus creating a Markov process. In the case that \(q\) is symmetric, it follows that \(q(a^k|\hat{a}) = q(\hat{a}|a^k)\) and the probability of accepting \(\hat{a}\) as \(a^{k+1}\) in step 2(iii) simplifies to

\[
\min \left\{ 1, \frac{\rho(\hat{a}|Y)}{\rho(a^k|Y)} \right\}.
\]

The variance of the proposal distribution greatly influences the exploration in parameter space. Large variance may cause most of the proposals to be rejected due to small likelihoods, thus the resulting chain remains stationary for many iterations. If the variance is chosen to be too small, the acceptance ratio will be high, but the exploration of the parameter space will be slow. Additionally, estimating an anisotropic posterior using an isotropic proposal distribution will decrease the efficiency of the exploration of the space [74]. It is recommended...
that the variance of the proposal distribution should be chosen so that on average, 25% of the proposals are accepted [77].

Step 2(ii) of the algorithm provides the connection to Bayes’ Theorem. The likelihood function $L(Y|a)$ and the prior $\pi(a)$ must be prescribed, and the resulting posterior may be highly influenced by this choice, as will be revealed in later examples. Note that in the computation of $\rho(\hat{a}|Y) = L(Y|\hat{a})\pi(\hat{a})$, the normalization constant is omitted. It is unnecessary to include this constant because in the ratio of posteriors used for the acceptance criterion, it is eliminated.

The acceptance criteria in step 2(iii) provides a means through which the chain can escape a local maximum in $\rho(a|Y)$. In the case of a symmetric proposal density, if $\rho(\hat{a}|Y) \geq \rho(a^k|Y)$ then $\hat{a}$ is accepted with probability 1, otherwise when $\rho(\hat{a}|Y) < \rho(a^k|Y)$ the probability of accepting $\hat{a}$ decreases as $\rho(\hat{a}|Y)$ decreases.

The sequence $a^1, a^2, \ldots, a^M$ produced by the algorithm is a Markov chain that, in the limit of $M \to \infty$, has the posterior density as its stationary distribution. How long the chains must be run to converge and sufficiently sample from the posterior is a challenging question, and analytic convergence and stopping criteria are lacking [74]. A burn in period, where the first $j$ elements of the chain are thrown out, is often employed because in the initial part of the algorithm, the proposals are not sampled from the stationary distribution. Several techniques have been introduced to improve exploration of the space and speed up convergence. For example, parallel tempering makes use of several Markov chains run simultaneously with different variances in the proposal distribution [20]. Swapping between the chains allows for refined sampling in regions near extrema and improved mixing, so that chains do not become stuck in local extrema. This approach can be computationally expensive. A less sophisticated, but useful means of improving mixing is to introduce a random proposal $\hat{a}$ selected from a uniform distribution on the parameter support every $X$ iterations of the algorithm (e.g. $X = 10$). This approach allows the chain to escape local extrema and increases the convergence rate of the chain.
In this section, we assume that the data is given as the distribution $\eta(y)$ and derive a prior density that will facilitate exact solution of the inverse problem when Bayesian inference is employed. We will propose the use of this prior in the Metropolis-Hastings algorithm and examples will be presented in following sections.

As defined in section (4.2.1), $A$ is the parameter space, $Y$ is the data space of observations, and $F : A \rightarrow Y$ represents the forward solution map from model parameters to data. Define $\rho(a)$ to be the parameter density, representing the distribution of parameters for the model. Let $\eta(y)$ be a density on the data space, such that for any measurable $\Omega \subseteq A$, we have that $F(\Omega) \subseteq Y$ and

$$\int_{F(\Omega)} \eta(y) dy = \int_{\Omega} \rho(a) da.$$ \hspace{1cm} \text{(4.3)}

If $\Omega$ is any subset of $A$ (of positive measure) on which $F$ is injective, then by the well known change of variables formula (Theorem 2.47 \[26\]),

$$\int_{F(\Omega)} \eta(y) dy = \int_{\Omega} \eta(F(a)) | \det D_a F(a) | da = \int_{\Omega} \eta(F(a)) J(a) da \text{ where } J(a) = | \det D_a F(a) | .$$

Thus, from equation (4.3), for any $\Omega \subseteq A$ on which $F$ is injective,

$$\int_{\Omega} \rho(a) da = \int_{\Omega} \eta(F(a)) J(a) da,$$

and therefore,

$$\rho(a) = \eta(F(a)) J(a).$$ \hspace{1cm} \text{(4.4)}

In the Bayesian inference approach, the posterior density is determined by making use of the available data. The density $\eta(y)$ on the data provides an ideal setting, in which full information about the data is known. The Bayes relation (4.2) with $Y = \eta(y)$, gives that the posterior density, $\rho(a|\eta)$, is proportional to the likelihood times the prior. That is,

$$\rho(a|\eta) = k L(\eta|a) \pi(a),$$

where $\pi(a)$ is the prior density on the parameters, $L(\eta|a)$ is the likelihood of the data given an element $a$ of the parameter space, and $k$ is a normalization constant. In the absence of
error in the forward model solution, it holds that \( L(\eta|a) = \eta(F(a)) \), ([79], page 35) and thus,

\[
\rho(a|\eta) = k\eta(F(a))\pi(a).
\]  

(4.5)

In order for the posterior \( \rho(a|\eta) \) to be equal to the parameter density \( \rho(a) \), it follows from equations (4.4) and (4.5), that the prior should be chosen as

\[
\pi(a) = \frac{J(a)}{k}.
\]  

(4.6)

We will refer to \( J(a) \) as the Jacobian prior, following the conventional terminology in which \( D\eta F(a) \) is called the Jacobian. We note that the term Jacobian prior has appeared previously [78], in the computation of a marginal with a change of variables, however the context is different and it does not appear to be a commonly used term. This informative prior is not simply based on previous knowledge of the parameters, but rather, on prior knowledge of the model structure and the connection between model parameters and the data (i.e. the solution map \( F \)). This derivation relies heavily on the injectivity of \( F \), which is guaranteed when the model (4.1) is identifiable. In the following section, we will use the analysis of Chapter 3, where explicit sets on which \( F^{-1} \) exists were established, in order to systematically study the accuracy of the posterior densities obtained by using the Jacobian prior in the Metropolis-Hastings algorithm.

4.4 SYSTEMATIC ANALYSIS OF THE INFLUENCE OF THE PRIOR

In this section, a systematic approach will be used to demonstrate the superiority of the performance of the Jacobian prior in the Metropolis-Hastings algorithm with that of other commonly used priors. In particular, the parameter density \( \rho(a) \) will be constructed for comparison with the posterior densities \( \rho(a|\eta) \) obtained from implementing the Metropolis-Hastings algorithm with various priors.

Model (4.1) is selected to be locally identifiable, thus guaranteeing the injectivity of \( F \). We will assume there is full knowledge of the data, given by the density \( \eta(y) \). The injectivity of \( F \) directly relates \( \eta(y) \) to a parameter density \( \rho(a) \) by equation (4.3). With
In the Bayes relation (4.2), the data directly informs the posterior only through the likelihood. Using \( \eta(y) \) as the data we have seen that \( L(\eta|a) = \eta(F(a)) \) (due to the absence of error in the forward model solution). This eliminates uncertainty in the estimation of \( \rho(a|\eta) \) due to the data and isolates the prior as the primary source of error, thus truly revealing the effect of the prior.

In the following section, we will use the approach introduced above to study the effectiveness of the Jacobian prior in obtaining accurate estimates of the posterior density with the Metropolis-Hastings algorithm. In the case of linear systems of ordinary differential equations, we will use the analysis of Chapter 3 in order to explicitly define the density \( \eta(y) \). In other circumstances, \( \eta(y) \) cannot be defined exactly and must be estimated. Two different approaches will be presented in order to address both of these situations.
4.4.1 Linear systems

Consider the linear system of differential equations defined as

\[
\begin{align*}
\dot{x}(t) &= \Lambda x(t) \\
x(0) &= b
\end{align*}
\]  

(4.7)

where \( x(t) \in \mathbb{R}^n \) is the state of the system at time \( t \), \( \Lambda \in \mathbb{R}^{n \times n} \) is a matrix of parameters, and \( b \in \mathbb{R}^n \) is the initial condition, which is now taken as a fixed known value rather than a free parameter. Since the initial condition is independent of the parameters, the model parameters are the entries of the matrix \( \Lambda \) written as a vector \( a \in \mathbb{R}^{n^2} \) and the parameter space is \( \mathcal{A} = \mathbb{R}^{n^2} \). As in Chapter 3, the data will be observed at \( n \) equally spaced time intervals (without loss of generality \( \Delta t = 1 \)) \( t_j = j, \ j = 1, \ldots, n \), yielding \( y = (x_1, \ldots, x_n) \in \mathcal{Y} \) with \( x_j \in \mathbb{R}^n \). Using \( n \) time points ensures that \( D_a F(a) \) is a square matrix.

Defining model (4.1) to be a linear system of differential equations provides several advantages in using our systematic approach to study the Jacobian prior. In Chapter 3, the existence of sets in the data space \( \mathcal{Y} \) on which \( F^{-1} \) exists for the linear dynamical system (4.7) was established. Due to the injectivity of \( F \), the Jacobian, \( J(a) \), can be used as the prior in the Metropolis-Hastings algorithm. The invertibility of \( F \) will be exploited to construct a direct sample of the parameter density \( \rho(a) \), which will be used to compare with the posteriors obtained by employing various priors in the Metropolis-Hastings algorithm.

4.4.1.1 Comparison of priors: Approach 1

Implementation of the systematic approach described by Figure 19, in practice, requires sampling of the densities and additional modifications from the abstract setting. The following description details our first method for applying this approach and Figure 20 provides a corresponding schematic diagram.

1. Fix \( y \in \mathcal{Y} \) such that \( F^{-1}(y) = a \) is unique (i.e. \( y \) such that the associated \( \Phi = [x_1 | \ldots | x_n][b | x_1 | \ldots | x_{n-1}]^{-1} \) has \( n \) positive distinct eigenvalues (Corollary 28)).
2. Numerically estimate \( C(y, \epsilon_\nu) \) using the technique introduced in Chapter 3, and define \( \eta(y) \) to be a uniform density on \( C(y, \epsilon_\nu) \).
3. Sample \( \eta(y) \) to obtain \( Y = \{y^1, y^2, \ldots, y^N\}, \ y^j \in \mathcal{Y} \) and compute the associated parameter values (via \( F^{-1} \), employing the techniques of Chapter 3). Denote this sample by \( A = \{a^1, a^2, \ldots, a^N\}, \ a^j \in \mathcal{A} \). Since \( F : A \to Y \) is a bijection, \( A \) is, by construction, a sample of the parameter density \( \rho(a) \).

4. Implement the Metropolis-Hastings algorithm using various priors \( \pi \) to obtain samples, \( M_\pi \), of the posterior density \( \rho(a|\eta) \).

5. Compare \( M_\pi \), a sample of the posterior \( \rho(a|\eta) \) obtained through from the Metropolis-Hastings algorithm, to \( A \), a sample of the parameter density \( \rho(a) \).

![Diagram of parameter space and data space](image.png)

Figure 20: Visual summary of the method used to construct a sample of the parameter density \( \rho(a) \) to order to compare with posteriors \( \rho(a|\eta) \) obtained using various priors \( \pi(a) \) in the Metropolis-Hastings algorithm. The red coloring indicates the first density that is defined to begin the approach.

### 4.4.1.2 Notes on the implementation of Approach 1 and the Jacobian prior in the Metropolis-Hastings algorithm

Implementation of the Metropolis-Hastings algorithm requires the prescription of the likelihood function, proposal distribution, and prior density. In the case of Approach 1, the data are supplied with full knowledge of the
density $\eta(y)$, which is defined as a uniform density over the set $C(y, \epsilon_v)$. For a given $a \in \mathcal{A}$, the solution $F(a)$ is uniquely defined and free from uncertainty, thus $L(\eta|a) = \eta(F(a))$. In step 2(ii) of the Metropolis-Hastings algorithm, the ODE is solved with parameter $\hat{a}$ and the likelihood is given by,

$$L(Y|\hat{a}) = \begin{cases} 1, & F(\hat{a}) \in C(y, \epsilon_v) \\ 0 & \text{otherwise} \end{cases}.$$ 

At the $k^{th}$ step of the algorithm, the proposal distribution is defined to be a multivariate Gaussian with mean $a^k$, specifically $\hat{a} \sim \mathcal{N}(a^k, \gamma I)$, where $I$ is the $p \times p$ the identity matrix. The variance $\gamma$ is tailored to achieve a proposal acceptance of approximately 25%.

In this study three priors will be employed, the uniform prior, Jeffreys prior, and the Jacobian prior. Jeffreys prior is defined to be $\pi(a) = (1/a_1, 1/a_2, \ldots, 1/a_p)$. It is important to note that Jeffreys prior is only valid when the signs of the parameters are fixed, hence it will only be used in such cases. The mixing procedure described in section 4.2.3 is used in order to speed up convergence; proposals are selected randomly one of every ten iterations of the algorithm. When bounds on the parameter values are known a priori, this information can be included in addition to the Jacobian prior. In practice, bounds on the parameter values are imposed during the parameter proposal step of the algorithm. Including this prior knowledge is also helps to improve convergence rates. Here, parameter bounds were imposed by broadening the observed bounds on $A$.

Since the use of the Jacobian prior in the Metropolis-Hastings algorithm is being proposed for the first time, the details of its inclusion in the algorithm will be discussed here. For a map $F : \mathbb{R}^p \rightarrow \mathbb{R}^p$, the Jacobian determinant is classically defined as $\det(D_aF(a))$, where $D_aF(a)$ is the $p \times p$ matrix of partial derivatives with components,

$$D_aF(a)_{ij} = \frac{\partial F_i}{\partial a_j}(a).$$

The inclusion of the Jacobian prior in the Metropolis-Hastings algorithm occurs in the application of Bayes’ Theorem in step 2(ii), when $\rho(\hat{a}|Y) = L(Y|\hat{a})\pi(\hat{a})$ is computed. Although equation (4.6) suggests that the prior should be chosen as $J(a)/k$, in practice, the inclusion of the normalization constant $k$ is not necessary because it cancels in the ratio $\rho(\hat{a}|Y)/\rho(a^k|Y)$ within the acceptance criterion. Thus, we take $\pi(a) = J(a)$ with the division by $k$ omitted.
Here, \( J(\hat{a}) = |\text{det}(D_a F(\hat{a}))| \), and \( D_a F(\hat{a}) \) can be numerically estimated can be numerically estimated using, for example,

\[
\frac{\partial F}{\partial a_j}(\hat{a}) = \frac{F(\hat{a} + \epsilon e_j) - F(\hat{a})}{\epsilon},
\]

where \( e_j \) a standard basis vector in \( \mathbb{R}^p \) and \( \epsilon \) is positive number very close to 0 (note that this equation estimates the \( j^{th} \) column of \( D_a F(\hat{a}) \)). This approximation will be used in the computation of \( J(a) \) in the Metropolis-Hastings algorithm for the examples presented in the subsequent sections.

As a final note, the phrasing “sample the density \( \eta(y) \)” means to sample the distribution represented by the density function \( \eta(y) \).

### 4.4.1.3 Approach 1 examples

In this section, several examples of Approach 1 will be presented to study the accuracy of the posterior densities obtained when the Jacobian prior is used. For ease of visualization, model (4.7) will be restricted to the case of two-dimensional linear systems. All numerical codes were written and implemented using MATLAB.

**Example 1.** Select the initial condition \( b \) and the data \( y \) to be

\[
b = \begin{bmatrix} 10 \\ 2 \end{bmatrix} \quad \text{and} \quad y = (x_1, x_2) = \left( \begin{bmatrix} 5 \\ -2.3 \end{bmatrix}, \begin{bmatrix} -4 \\ -2.2 \end{bmatrix} \right)
\]

representing observations of the system at times \( t = 1, 2 \). The corresponding matrix \( \Phi \) has two distinct positive eigenvalues, thus \( F^{-1}(y) = a \) is unique. For this choice of \( b \) and \( y \), the parameter vector \( a = (\lambda_{11}, \lambda_{12}, \lambda_{21}, \lambda_{22}) \) gives a linear system (4.7) with a stable node fixed point at the origin. Numerical computation of the maximal permissible uncertainty in the data such that uniqueness of the inverse is maintained yields \( \epsilon_u = 0.12 \). Note that because the initial condition \( b \) is a fixed value, \( C(y, \epsilon_u) = \{b\} \times c(x_1, \epsilon_u) \times c(x_2, \epsilon_u) \) as defined in Chapter 3.

Define the data density \( \eta \) to be a uniform density on the set \( C(y, \epsilon_u) \). \( N = 800,000 \) random elements from \( C(y, \epsilon_u) \) are selected to obtain a sample of \( \eta \) denoted by \( Y = \{y^1, y^2, \ldots y^N\} \). Figure 21 (a) depicts squares bounding the sets \( c(x_1, \epsilon_u) \) and \( c(x_2, \epsilon_u) \) in the phase plane, with the sample \( Y \) shown in green (the boxes appear to be colored solid simply due to the large number of samples, \( N \).) Figure 21 (b) shows the marginalized density function \( \eta(y) \)
in black and marginals of the sample $Y$ are displayed with histograms. The notation $x_{i,j}$ is used to denote the $j^{th}$ coordinate of the $i^{th}$ observation $x_i$.

Figure 21: (a) In the phase plane, boundaries of the sets $c(x_1, \epsilon_u)$ and $c(x_2, \epsilon_u)$ are shown in black, and the sample $Y$ is depicted in green. (b) The marginalized density function $\eta(y)$ is displayed black, and the blue histograms depict marginals of the sample $Y$.

The inverses $F^{-1}(y^j) = a^j$ are computed to form $A = \{a^1, a^2, \ldots a^N\}$, a sample of the parameter density $\rho(a)$. $N$ was chosen to be large to obtain an accurate representation of $\rho(a)$. Next, the Metropolis-Hastings algorithm is implemented and Jeffreys prior, the uniform prior, and the Jacobian prior are employed to obtain samples of the posterior $\rho(a|\eta)$, denoted by $M_{Jeff}$, $M_{Unif}$, and $M_{Jac}$, respectively. The algorithm was run for 2.5 million parameter proposals and convergence was checked through multiple runs. Figure 22 represents marginalized histograms of $A$, $M_{Jeff}$, $M_{Unif}$, and $M_{Jac}$, where the curves are defined by the bin centers and heights (i.e. a frequency polygon). As is common in practice, only every 100th element of the posterior sample $M$ is included in the histogram to reduce the effects of correlation inherent in MCMC sampling. In this and all subsequent examples, the following colors will be used, $A$ is red, $M_{Jeff}$ is blue, $M_{Unif}$ is black, and $M_{Jac}$ is green. Thus, for the sample of the posterior obtained from the Metropolis-Hastings algorithm to accurately represent the parameter density, we seek to match the red curve. In this figure, $A$, $M_{Jeff}$, and $M_{Jac}$ overlap nicely, with a slightly better matching between $M_{Jac}$ and $A$ in
the last two parameters. Thus, employing both the Jacobian and Jeffreys priors provides good representation of the parameter density. However, in this example, the uniform prior does not yield an accurate estimate of $\rho(a)$, as seen in the vastly different curves for $A$ (red) and $M_{Unif}$ (black).

Visual comparison of the marginals of $A$ and $M_\pi$ provides an effective way to view the shapes of the posteriors and analyze their similarity. Histograms depend greatly on the bin width, so in addition to this comparison, a two-sample Kolmogorov-Smirnov test was performed on the marginals to determine if the samples $A$ and $M_\pi$ are drawn from the same continuous distribution. For a significance level $\alpha$, the Bonferroni correction for testing 4 hypotheses is $\alpha/4$. Using the kstest2 function in MATLAB, the computed $p$-values for $M_{Jeff}$ and $M_{Unif}$ were all less than $1 \times 10^{-30}$ and the $p$-values for $M_{Jac}$ were computed as \{0.05, 0.08, 0.31, 0.68\} for the parameters $\{\lambda_{11}, \lambda_{12}, \lambda_{21}, \lambda_{22}\}$ respectively. As in the histogram representations, the only every 100th element of the posterior sample $M_\pi$ was included in the computation of the $p$-values in order to de-emphasize the false correlations in the chain due to the random walk procedure. The $p$-values indicate that $M_{Jac}$ is the most likely to have been drawn from the same distribution as $A$.

![Figure 22: Example 1: Curves representing marginalized histograms for $A$ (red), $M_{Jeff}$ (blue), $M_{Unif}$ (black), and $M_{Jac}$ (green).](image)

**Example 2.** Select the initial condition $b$ and the data $y$ to be

$$b = \begin{bmatrix} 10 \\ 2 \end{bmatrix} \quad \text{and} \quad y = (x_1, x_2) = \left( \begin{bmatrix} 6.1 \\ -4.4 \end{bmatrix}, \begin{bmatrix} 7 \\ -10 \end{bmatrix} \right).$$
In this example, the unique matrix $a = F^{-1}(y)$ corresponds to a linear system with a saddle fixed point at the origin. Numerical computation of the maximal permissible uncertainty in the data such that uniqueness of the inverse is maintained yields $\epsilon_u = 1.08$. Using the same procedures as in Example 1, $A$, $M_{Unif}$, and $M_{Jac}$ are computed and are pictured in Figure 23. In this case, the Jeffreys prior should not be used because the parameters do not have a fixed sign. Here, we find that the Jacobian prior produces a posterior which nicely matches the parameter density and again, the uniform prior yields an inaccurate estimate. The computed $p$-values for $M_{Jac}$ are $0.05, 0.02, 0.004, 0.002$ and essentially 0 for $M_{Unif}$ (less than $1 \times 10^{-200}$).

![Figure 23: Example 2: Curves representing marginalized histograms for $A$ (red), $M_{Unif}$ (black), and $M_{Jac}$ (green).](image)

**Example 3.** Select the initial condition $b$ and the data $y$ to be

$$b = \begin{bmatrix} 5 \\ 15 \end{bmatrix} \quad \text{and} \quad y = (x_1, x_2) = \begin{bmatrix} 10 \\ 10 \\ 30 \\ -15 \end{bmatrix},$$

so that the unique matrix $a = F^{-1}(y)$ corresponds to a linear system with an unstable node fixed point at the origin. Numerical estimation gives $\epsilon_u = 1.03$. Using the same procedures as in Example 1, $A$, $M_{Jeff}$, $M_{Unif}$, and $M_{Jac}$ are computed and are pictured in Figure 24. Here, the the posteriors using the Jacobian prior approximate the parameter density well in all four parameters and the posterior using the uniform prior gives good approximations in the first three parameters. In this example, the Jeffreys prior yields poor estimates.
The computed $p$-values for $M_{Jeff}$ and $M_{Unif}$ are all less than $1 \times 10^{-16}$, and for $M_{Jac}$ are \{0.26, 0.15, 0.22, 0.006\}.

![Figure 24: Example 3: Curves representing marginalized histograms for $A$ (red), $M_{Jeff}$ (blue), $M_{Unif}$ (black), and $M_{Jac}$ (green).](image)

In this series of examples, employing the Jacobian prior consistently produced good approximations of parameter density, while accuracy resulting from using the Jeffreys and uniform priors varied from case to case. Visual inspection of the marginals showed instances where the uniform and Jeffreys priors seemed to give results comparable to the Jacobian prior, however, the computed $p$-values indicate that the posterior sample $M$ is most likely to be drawn from the parameter distribution when the Jacobian prior is used.

The ability to explicitly define the data density $\eta(y)$ was ideal because it eliminated error in the posterior estimation due to uncertainty about the available data. In applications, experimental measurements are sparse in the data space and it is not possible to determine the data density exactly. In the following section, we will introduce a second approach and use it to study the accuracy of the posteriors obtained with the Jacobian prior in this practical setting.

### 4.4.1.4 Comparison of priors: Approach 2

Typically, the data that will be used to estimate the parameters of a system of ordinary differential equations are a finite collection of experimental observations taken at incremented time intervals. The available data $Y$ may then be supplied as a collection of $N$ repeated observations, $\{y^1, y^2, \ldots, y^N\}$. In this case,
decisions must be made concerning how to represent the data in order to define the likelihood function \( L(Y|a) \). The more accurately the data can be represented, the more reliable will be the estimated posterior density. To study the influence of the prior in this setting, the following approach will be used. This technique is again an adaptation of the systematic approach described in Figure 19, while accounting for practical limitations. The following description details our method for applying the second approach and Figure 25 provides a corresponding schematic diagram.

1. Define a density \( \rho(a) \) on the parameters.
2. Sample \( \rho(a) \) to obtain \( A = \{a^1, a^2, \ldots a^N\} \), \( a^j \in \mathcal{A} \), restricting the sampling so that \( F \) is injective on \( A \).
3. Fix the time points for observation as \( t_1, \ldots, t_n \) and compute the forward solution \( F(a^j) = y^j \) for all \( j \), yielding the collection of simulated data \( Y = \{y^1, y^2, \ldots y^N\} \), \( y^j \in \mathcal{Y} \). Since \( F : A \rightarrow Y \) is a bijection, \( A \) is, by construction, a sample of the parameter density that we wish to approximate with the Metropolis-Hastings algorithm.
4. Use a density estimation method (such as multivariate kernel density estimation) to represent the sample \( Y \) with a density function \( \tilde{\eta}(y) \).
5. Implement the Metropolis-Hastings algorithm using various priors \( \pi \) to obtain samples, \( M_\pi \), of the posterior density \( \rho(a|\tilde{\eta}) \).
6. Compare \( M_\pi \), a sample of the posterior \( \rho(a|\tilde{\eta}) \) obtained through from the Metropolis-Hastings algorithm, to \( A \), a sample of the parameter density.

4.4.1.5 Notes on the implementation of Approach 2 A few comments will be made here to explain how the steps outlined in Approach 2 shall be implemented. The initial condition \( b \) is chosen as a fixed, known value and thus the parameters of the model are the entries of the matrix \( \Lambda \). Then, \( a \in \mathbb{R}^{n^2} \) is the parameter vector consisting of the matrix entries of \( \Lambda \). \( \rho(a) \) is defined as a multivariate Gaussian distribution \( a \sim \mathcal{N}(\bar{a}, \Sigma) \) with mean vector \( \bar{a} \) and covariance matrix \( \Sigma = \sigma^2 I \) where \( I \) is the \( n^2 \times n^2 \) the identity matrix. Then, the density function is defined to be
\[
\rho(a) = \frac{1}{\sqrt{(2\pi)^{n^2}|\Sigma|}} \exp \left( -\frac{1}{2} (a - \bar{a})^\top \Sigma^{-1} (a - \bar{a}) \right).
\]
Figure 25: Visual summary of approach 2. The red coloring indicates the first density that is defined to begin the approach. KDE refers to the kernel density estimation of $Y$ with a density $\tilde{\eta}(y)$.

When, the sample the $A = \{a^1, a^2, \ldots a^N\}$ is randomly drawn from the density $\rho(a)$, we restrict to parameters for which the data is uniquely produced by that parameter (for $a \in A$, $F(a) = y$ and there is no other $b \in A$ such that $F(b) = y$). This can easily be achieved in practice by selecting a mean parameter $\bar{a}$ with real, distinct eigenvalues and taking $\sigma$ to be small. Even if $\sigma$ is small, $a^j$ drawn from a Gaussian distribution might fail to have real, distinct eigenvalues, in theory. Only $a^j$ with real, distinct eigenvalues are included in the sample. The solution map $F$ is then used to find the corresponding data $F(a^j) = y^j$, which yields the sample $Y = \{y^1, y^2, \ldots y^N\}$. Due to the restriction on the sampling, it follows that $F : A \rightarrow Y$ is a bijection as desired. The value $N$ is chosen to be large, to gain as much information about the structure of the data density as possible.

We aim to diminish the error in the posterior sample, $M$, resulting from the choice of
how the data are represented and which likelihood function is used. In order to achieve this objective, we seek a density function which is as close as possible to the underlying distribution that generated the sample $Y$. For the finite sample of observed data, density estimation constructs an estimate of the underlying probability density function everywhere, including where no data are observed. Histograms are of the most simple density estimators; however, they are not smooth and the choice of bin width leads to a constant struggle between bias and variance [72]. Kernel density estimation is a more sophisticated alternative. In this approach, a kernel function is centered at each data point, so that a discrete data point is smoothed over the region surrounding it. The kernels are then summed, resulting in single function that is the estimated density function. Background on kernel density estimation is provided in Appendix B.

In MATLAB, the built in function ‘ksdensity’ performs a kernel density estimation for finite sets of univariate and bivariate data. In the setting of this work, the data have dimension $nm$, which will be larger than 2 in all of the following examples. MATLAB code for multivariate kernel density estimation based on the papers [54, 55] has been made freely available by the authors. One desirable feature of this toolbox is that the bandwidth does not need to be prescribed and is optimally determined within the program. It can also handle large data sets and performs a pre-clustering in order to do so. This code, with minor modifications, was used for all density estimates presented in this work. Figure 26 shows a 4-dimensional kernel density estimate obtained using this code. In this figure, marginals of the data are represented by blue histograms and marginals of the density estimate are given by the red curves. The small nest of black curves depict marginals of the individual kernels that were summed for the density estimate.

Using the same data set as Figure 26, two-dimensional projections of the data and density estimate are given in Figure 27. In part (a), projections of the discrete data set are depicted and in part (b), two-dimensional projections of the density estimate are shown, with the third dimension represented by a color shading. Both of these figures indicate that the density estimate produces an accurate representation of the data.

Returning to the implementation of Approach 2, an approximate density of sample $Y$ is constructed with multivariate kernel density estimation, using the freely available MATLAB
Figure 26: Example kernel density estimate for 4-dimensional data. Marginal histograms of the data are shown in blue, marginals of the density estimate are given by the red curves, and marginals of the kernels surrounding the data points are given by the black curves.

Figure 27: Example kernel density estimate for 4-dimensional data. Two dimensional projections of the discrete data set are pictured in (a) and projections of the kernel density estimate are given in (b).

code from [54] and is denoted by $\tilde{\eta}(y)$. Next, the Metropolis-Hastings algorithm is performed and Jeffreys prior, the uniform prior, and the Jacobian prior are employed in order to obtain $M_{\text{Jeff}}$, $M_{\text{Unif}}$, and $M_{\text{Jac}}$, respectively.

In the implementation of the Metropolis-Hastings algorithm, the density estimate $\tilde{\eta}(y)$
will be used in the computation of the likelihood in the same manner that the exact data density \( \eta(y) \) was used in Approach 1, namely \( L(\bar{\eta}|a) = \bar{\eta}(F(a)) \). The proposal density and the numerical estimation of the Jacobian will be the same as presented in section 4.4.1.2.

The accuracy of the kernel density estimate reduces the error in the posterior estimate due to incomplete knowledge of the data and the choice of likelihood, and therefore the effect of the prior density will be revealed.

4.4.1.6 Approach 2 examples In this section, several examples of Approach 2 will be presented in order study the accuracy of the posterior densities obtained when the Jacobian prior is used. For ease of visualization, model (4.7) will again be restricted to the case of two-dimensional linear systems.

Example 4. This example follows the steps outlined in Approach 2 and discussed in section 4.4.1.5. The initial condition is fixed as \( b = (10, 2)^T \). The parameters are the entries of the matrix \( \Lambda \) written as a vector, \( a = (\lambda_{11}, \lambda_{12}, \lambda_{21}, \lambda_{22}) \). The mean parameter vector used to define \( \rho(a) \) is \( \bar{a} = (-1, -1.5, -1, -2) \), and the observation times are set as \( t_1 = 1 \) and \( t_2 = 2 \). Here, the mean parameter value corresponds to a linear system with a stable node equilibrium. A series of three examples is presented with increasing values for \( \sigma \) across all three. The following values are used in each example: (a) \( \sigma = 0.15 \), (b) \( \sigma = 0.2 \), and (c) \( \sigma = 0.25 \). \( Y \) was sampled from \( \rho(a) \) using \( N = 5000 \). In Figure 28, the marginals of \( Y \) are pictured as histograms and overlaying each is the marginalization of \( \bar{\eta}(y) \) drawn as a red curve. The individual kernels, which were summed over in the density estimation, are depicted by the (nest of) black curves. As \( \sigma \) increases, the spread of the data increases and in each case, the kernel density estimate \( \bar{\eta}(y) \) appears to accurately represent the data.

The Metropolis-Hastings algorithm is implemented to obtain \( M_{Jeff} \), \( M_{Unif} \), and \( M_{Jac} \). In the same manner as the examples for Approach 1, Figure 29 displays the marginalized histograms of \( A \), \( M_{Jeff} \), \( M_{Unif} \), and \( M_{Jac} \). As before, we seek to match the red curve which depicts \( A \), the sample of the parameter distribution.

In part (a), \( M_{Jac} \) and \( A \) match very closely, while \( M_{Jeff} \) and \( M_{Unif} \) are quite similar to one another, but do approximate \( A \) as well. As \( \sigma \) increases in parts (b) and (c), we observe that the \( M_{Jac} \) continues to approximate \( A \) well, but the accuracy of \( M_{Jeff} \) and \( M_{Unif} \) degrades
Figure 28: Marginal histograms of $Y$ are pictured, along with marginalizations of the density $\tilde{\eta}(y)$ obtained from kernel density approximation, which are displayed as red curves. The black curves represent the individual kernels which are summed in the density estimation. In part (a) $\sigma = 0.15$, (b) $\sigma = 0.2$, and (c) $\sigma = 0.25$, all with the same mean parameter $\tilde{a} = (-1, -1.5, -1, -2)$. severely. In part (c) we finally see a difference between $M_{Jeff}$ and $M_{Unif}$ with Jeffreys performing slightly better in the last parameter. The marginals for $M_{Jeff}$ in part (c) for the parameters $\lambda_{12}$ and $\lambda_{21}$ increase near zero due to properties of Jeffreys prior. Jeffreys prior is
Figure 29: Curves representing marginalized histograms for $A$ (red), $M_{Unif}$ (black), and $M_{Jac}$ (green). In part (a) $\sigma = 0.15$, (b) $\sigma = 0.2$, and (c) $\sigma = 0.25$, all with the same mean parameter $\bar{a} = (-1, -1.5, -1, -2)$.

uniform in the log scale and must be bounded away from zero, as an infinite likelihood occurs there. The $p$-values were computed using a two-sample Kolmogorov-Smirnov test comparing $A$ with the various samples $M_\pi$ to determine the likelihood that the samples were drawn from the same distribution. The $p$-values are reported in Table 2. They further indicate that
employment of the Jacobian prior produces the most accurate estimates of the parameter density.

<table>
<thead>
<tr>
<th></th>
<th>(a)</th>
<th>(b)</th>
<th>(c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$M_{Jac}$</td>
<td>${0.14, 0.63, 0.44, 0.02}$</td>
<td>${0.09, 0.51, 0.28, 0.16}$</td>
<td>${0.99, 0.05, 0.52, 0.1}$</td>
</tr>
<tr>
<td>$M_{Jeff}$</td>
<td>${0, 0, 0, 0}$</td>
<td>${0, 0, 0, 0}$</td>
<td>${0, 0, 0, 0}$</td>
</tr>
<tr>
<td>$M_{Unif}$</td>
<td>${0, 0.0005, 0, 0}$</td>
<td>${0, 0, 0, 0}$</td>
<td>${0, 0, 0, 0}$</td>
</tr>
</tbody>
</table>

Values less than $1 \times 10^{-5}$ were rounded to 0.

Table 2: Two sample Kolmogorov-Smirnov test computed $p$-values comparing the marginals of $A$ with the marginals of $M_\pi$ for the parameters $\{\lambda_{11}, \lambda_{12}, \lambda_{21}, \lambda_{22}\}$.

**Example 5.** Choosing how to represent the available data and selecting a likelihood function are important issues that must be confronted in order to implement the Metropolis-Hastings algorithm. Given a collection of $N$ observations $Y = \{y^1, \ldots, y^N\}$, there are several alternatives for how to represent these data. One option is to compute the mean and standard deviations at each time $t_1, t_2, \ldots, t_m$, denoting them by $\{\bar{x}_1, \ldots, \bar{x}_m\}$ and $\{\sigma_1, \ldots, \sigma_m\}$, respectively. Then the point averaged likelihood function can be defined as

$$L(Y|a) = \prod_i L_i(\bar{x}_i|a)$$

where,

$$L_i(\bar{x}_i|a) = \prod_j \frac{1}{\sigma_{i,j}\sqrt{2\pi}} e^{-\frac{(x_j(t_i|a) - \bar{x}_{i,j})^2}{2\sigma_{i,j}^2}}.$$ 

Vital characteristics of the data may be lost in selecting this representation. Additionally, summarizing the data in this way, inherently assumes that the data at each time point are normally distributed, which may not be accurate. This representation of the data is frequently used by practitioners of the Metropolis-Hastings algorithm for parameter estimation.

An alternative to summarizing the data $Y = \{y^1, \ldots, y^N\}$ with these statistical quantities, is to represent it using a multidimensional histogram. This approach is most relevant when the system size, $n$, and the number of time points, $m$, in which the data are collected are
small, and the number of repeated observations, \( N \), is large. In this case, an \( nm \)-dimensional histogram could be used to summarize the data. If the histogram is normalized to represent a probability density, then the likelihood function can be easily defined using the bin heights. As is the case with one dimensional histograms, the bin widths must be prescribed and may greatly affect the structure of the histogram. After the bin widths are selected and the normalized histogram is constructed, it can be used to define the likelihood for use in the Metropolis-Hastings algorithm in the following way: for a proposed parameter \( \hat{a} \), the data \( F(\hat{a}) \) is computed and the histogram bin containing that data point is identified, then the likelihood \( L(Y|\hat{a}) \) is the height of that bin.

As we have seen, kernel density estimation provides an accurate representation of the data, and the likelihood is the evaluation of the estimated probability density function, \( L(\tilde{\eta}|\hat{a}) = \tilde{\eta}(F(\hat{a})) \).

In Approach 2, we aimed to study the effect of the prior on the accuracy of the posterior obtained from the Metropolis-Hastings algorithm. In order to minimize the error in posterior due to the representation of the data, we employed kernel density estimation to define the density \( \tilde{\eta}(y) \). In this example we would like to explore the effect of the choice of data representation and likelihood function on the posteriors obtained from the Metropolis-Hastings algorithm.

As in Example 4(a), we fix \( \mathbf{b} = (10, 2)^\top \), \( \mathbf{a} = (-1, -1.5, -1, -2) \), \( \{t_1, t_2\} = \{1, 2\} \), and \( \sigma = 0.15 \). The data \( Y = \{y^1, \ldots, y^N\} \) are constructed using \( N = 10,000 \) observations at times \( \{1, 2\} \). \( Y \) is then represented using the three different alternatives described above as shown in Figure 30. Here, the green curves depict the marginalized normal density used to describe the data in the point averaged approach, with computed means \( \{\bar{x}_{1,1}, \bar{x}_{1,2}, \bar{x}_{2,1}, \bar{x}_{2,2}\} = \{5.2, -2.4, 4.3, -2, 4\} \) and standard deviations \( \{0.85, 0.56, 1.41, 0.85\} \). Marginalizations of the 4-dimensional histogram, constructed with 30 bins in each dimension, are shown in yellow. Lastly, the blue curves represent the marginalized probability density function obtained from the kernel density estimation. There is a clear discrepancy between the point averaged and kernel density estimation representations. The Metropolis-Hastings algorithm is then implemented with the Jacobian prior, and the appropriate likelihood function is employed in each case. Figure 31 compares the marginals of \( M_{Jac} \) computed using
Figure 30: Marginalizations of the three different representations of the data are shown. The point averaged representation is depicted by the green curve, the probability density function from the kernel density estimation is depicted by the blue curve, and the histograms are shown in yellow.

three different representations of the data. The marginals for $A$ are depicted in red. The marginals of $M_{Jac}$ are green for the point averaged approach, yellow for the 4-dimensional histogram, and blue for the kernel density estimation approach. In this example, the point

Figure 31: Marginal histograms for $A$ (red) and three estimates $M_{Jac}$ obtained employing kernel density estimation (blue), point averaged (green), and multidimensional histogram (yellow).

averaged approach resulted in the poorest estimate of the parameter density. The histogram
4.4.2 Jacobian prior with nonlinear systems of differential equations

In this section, the use of the Jacobian prior for parameter estimation in a nonlinear system of differential equations will be studied. The system that will be used to conduct this study is a model for influenza A virus infection from [6]:

\[
\begin{align*}
\dot{V} &= rI - cV \\
\dot{H} &= -\beta HV \\
\dot{I} &= \beta HV - \delta I
\end{align*}
\] (4.8)

where \( V \) is the concentration of infectious viral particles with units TCID\(_{50}\)/ml, \( H \) is the number of uninfected target cells, and \( I \) is the number of productively infected cells. By shedding viral titers, infected cells increase the concentration of viral particles at a rate of \( r \) per cell, and free viral particles are cleared at a rate of \( c \) per day. Uninfected cells interact with virus particles and become infected at a rate \( \beta HV \). The infected cells die at a rate of \( \delta \) per cell (where \( 1/\delta \) is the average life span of an infected cell). The initial number of infected cells is taken to be zero, and the initial viral concentrations, \( V(0) = V_0 \), and initial number of target cells, \( H(0) = H_0 \), are considered as parameters. Then, the vector of model parameters is \( a = (V_0, H_0, \beta, r, c, \delta) \in \mathbb{R}^6 = \mathcal{A} \), with units (TCID\(_{50}\)/ml, cells, (TCID\(_{50}\)/ml\()^{-1}\)day\(^{-1}\), (TCID\(_{50}\)/ml)day\(^{-1}\), day\(^{-1}\)).

Approach 2 again provides a systematic way to study which prior makes the posterior a better approximation of the parameter density. Figure 25 gives a roadmap for the approach and the details for this specific setting are discussed here. The parameter density \( \rho(a) \) is defined to be a normal multivariate density with mean parameter vector \( \bar{a} \) and covariance matrix \( \Sigma = \text{diag} \left( (\bar{a}_1\sigma_1)^2, (\bar{a}_2\sigma_2)^2, \ldots, (\bar{a}_6\sigma_6)^2 \right) \). In the examples below, \( \rho(a) \) is constructed by prescribing \( \bar{a} \) and the vector \( \sigma = (\sigma_1, \sigma_2, \ldots, \sigma_6) \). As before, \( \rho(a) \) is sampled to obtain \( A = \{a^1, a^2, \ldots, a^N\} \), \( a^j \in \mathcal{A} \). The number of observation times for the data will be taken
to be $m = 2$ (so that $p = mn = 6$ and $D_a F(a)$ is a square matrix). With this setup, for a fixed $a$, the solution map gives, $F(a) = ((V(t_1), H(t_1), I(t_1))^T, (V(t_2), H(t_2), I(t_2))^T) = y$. The forward solution, $F(a^j) = y^j$, is computed for all $j$, yielding the collection of data $Y = \{y^1, y^2, \ldots, y^N\}$, $y^j \in \mathcal{Y}$. As in the previous treatment, multivariate kernel density estimation is used to represent the sample $Y$ with a density function $\tilde{\eta}(y)$, and the Metropolis-Hastings algorithm is implemented using various priors, to obtain $M_\pi$. The posteriors, $M_\pi$, will be compared to the $A$, the sample of the parameter density, and to each other. A series of four examples using this approach is now presented.

Example 6. In this example, $\rho(a)$ is constructed by selecting $\bar{a} = (0.093, 4 \times 10^8, 2.7 \times 10^{-5}, 0.012, 3.0, 4.0)$ and $\sigma = (0.01, 0.05, 0.01, 0.05, 0.02, 0.01)$. Here, the mean parameter values are chosen to be the average of the best-fit parameter values from [6] and $\sigma$ was prescribed to contain small values. The observation times are taken to be $\{t_1, t_2\} = \{1, 2\}$, and the number of samples for $A$ and $Y$ is $N = 1000$. In Figure 32 (a), the solution to the system for the mean parameter $\bar{a}$ is given by the blue curves, and the box plots represent the sample of data $Y$, graphed at the associated time points for the corresponding variables. The figure shows that there is a large spread in the data for this selection of $\sigma$. Part (b) of Figure 32 shows marginal histograms of $Y$, and marginals of the kernel density estimate are given by the red curves. Finally, in part (c) of the figure, $A$ is compared to the posterior samples $M_{\text{eff}}$, $M_{\text{Unif}}$, and $M_{\text{Jac}}$ obtained from the Metropolis-Hastings algorithm. We can see that $M_{\text{Jac}}$ most accurately approximates $A$ for the parameters $H_0, \beta, r, \text{ and } \delta$. None of the posteriors approximate the marginals for $V_0$ or $c$ well. The parameter $c$ is related to the virus clearance; we can see from Figure 32, that for this choice of $t_1$ and $t_2$, the data do not capture any information about the decay of the virus. As before, a two sample Kolmogorov-Smirnov test was conducted, however the $p$-values were too small in each of the cases to provide any useful insight in comparing $M_{\text{eff}}$, $M_{\text{Unif}}$, and $M_{\text{Jac}}$, thus we will rely on the marginal depictions for comparison. This will also be the case in the subsequent examples.

Example 7. In this example, everything is the same as Example 6, but the value of $\sigma$ is increased to $\sigma = (0.05, 0.05, 0.05, 0.1, 0.04, 0.02)$ for the construction of $\rho(a)$. Figure 33 (a) again shows the solution for the mean parameter and the box plot representations of $Y$. As expected, the broadness in the spread of the data is increased. In Figure 33 (b), we compare
Figure 32: Example 6: (a) Solutions curves from the mean parameter value and box plot representations of $Y$. The box plots at $t = 1$ for $V$ and $I$ are difficult to see because they are tight relative to the figure scale. (b) marginal histograms of $Y$ and the kernel density estimate, (c) curves representing marginalized histograms for $A$ (red), $M_{Jeff}$ (blue), $M_{Unif}$ (black), and $M_{Jac}$ (green).

$A$, $M_{Jeff}$, $M_{Unif}$, and $M_{Jac}$ and see that again, $M_{Jac}$ most accurately approximates $A$ for the parameters $H_0$, $\beta$, $r$, and $\delta$. In this example, there is a more pronounced discrepancy between $M_{Unif}$ and $M_{Jeff}$ and $A$ than in Example 6. The observation times remained the same; still no information is gathered about the decay rate of the virus, and thus the marginal for $c$ is not approximated well by any of the posteriors.

Example 8. In this example, different time points are chosen in an effort to obtain more information about $c$. Let $\bar{a}$ and $N$ be the same as the previous two examples, and choose
Figure 33: Example 7: (a) Solutions curves from the mean parameter value and box plot representations of $Y$. The box plots at $t = 1$ for $V$ and $I$ are difficult to see because they are tight relative to the figure scale. (b) curves representing marginalized histograms for $A$ (red), $M_{eff}$ (blue), $M_{Unif}$ (black), and $M_{Jac}$ (green).

$$\sigma = (0.01, 0.02, 0.01, 0.01, 0.02, 0.01)$$ and $\{t_1, t_2\} = \{2, 3\}$. Figure 34 (a) shows the solution for the mean parameter and the box plot representations of $Y$. The new observation times now capture data in both the increasing and decreasing portions of the $V$ solution. Figure 34 (b) depicts marginals of $A$, $M_{eff}$, $M_{Unif}$, and $M_{Jac}$. There is a vast improvement in the estimation of $c$. We find that all of the priors produce very similar posteriors, and all of the marginals approximate $A$ well, except for $V_0$. In this example, the accuracy of the approximation of the parameter density between the different priors is indistinguishable.

Example 9. For the final example, the mean parameter used to define $\rho(a)$ is changed to $\bar{a} = (0.25, 4 \times 10^8, 1.4 \times 10^{-2}, 2.7 \times 10^{-5}, 3.2, 3.2)$, from [77]. We choose $\sigma = (0.01, 0.0001, 0.01, 0.05, 0.02, 0.01)$, $N = 1000$, and $\{t_1, t_2\} = \{1, 2\}$. As before, Figure 35 (a) shows the solution for the mean parameter, $\bar{a}$, and the box plot representations of $Y$, and (b) depicts marginals of $A$, $M_{eff}$, $M_{Unif}$, and $M_{Jac}$. In a similar manner as Example 8, the observation times capture data in both the increasing and decreasing portions of the $V$ solution, so the posteriors for
Figure 34: Example 8: (a) Solutions curves from the mean parameter value and box plot representations of $Y$. The box plot at $t = 3$ for $H$ is difficult to see because it is tight relative to the figure scale. (b) curves representing marginalized histograms for $A$ (red), $M_{Jeff}$ (blue), $M_{Unif}$ (black), and $M_{Jac}$ (green).

c are meaningful. In the marginals of the posteriors, we see that both initial condition parameters are not approximated well with any of the three priors. For the remaining parameters, the marginals of $M_{Jeff}$, $M_{Unif}$, and $M_{Jac}$ are all quite similar to each other. $A$ is matched well for the parameters $\beta, c, \delta$, but less accurately for $r$. Again, we cannot distinguish between the performance of the three priors.

4.5 CONCLUSION AND DISCUSSION

In this chapter, a new informative prior was introduced as an alternative prior density to employ in the Metropolis-Hastings algorithm. The Jacobian prior does not solely rely on previous information about the parameters, but instead, exploits the known structure of the
Figure 35: Example 9: (a) Solutions curves from the mean parameter value and box plot representations of $Y$. The box plot at $t = 2$ for $H$ is difficult to see because it is tight relative to the figure scale. (b) curves representing marginalized histograms for $A$ (red), $M_{Jeff}$ (blue), $M_{Unif}$ (black), and $M_{Jac}$ (green).

differential equation model. This prior may not have been previously considered in the setting of Bayesian inference, because it is specific to the problem of parameter estimation. The model solution provides the map $F$, which does not appear in Bayes’ Theorem explicitly. $F$ is introduced in the Bayesian inference approach to parameter estimation through the likelihood function, which makes the connection between available data and the model solution for a given parameter.

Two systematic approaches were used to study which prior makes the posterior a better approximation of the parameter density. In each approach, a sample of the parameter density was constructed in order to compare with the posteriors obtained by implementing the Metropolis-Hastings algorithm with various priors. In the case of linear systems of ordinary differential equations, we found that the Jacobian prior consistently produced better approximations of the parameter density than both the uniform and Jeffreys priors. In our study using the nonlinear influenza model, the results were less conclusive. We obtained better ap-
proximations of the parameter density using the Jacobian prior in two examples, and found that the performance of the Jeffreys, Jacobian, and uniform priors was indistinguishable in the other two examples. Analysis with additional nonlinear models is needed.

In the case of linear systems of differential equations, the theory of Chapter 3 guaranteed the invertibility of $F$. In the influenza model, we were unable to verify the invertibility of $F$; some inaccuracy in the approximation may be due to this. The parameters $\beta, r, c, \delta$ of model (4.8) were shown to be globally identifiable from 3 measurements of $V$, 2 of $H$, and 2 of $T$ in [63]; however, they did not consider any initial conditions as parameters. Further investigations are needed to determine the importance of the invertibility of $F$ in practice. It may be the case that local identifiability (local invertibility) is sufficient.

In this chapter, we required that $p = nm$ (number of parameters is equivalent to the number of model equations multiplied by the number of observation times). This restriction was imposed to ensure that the Jacobian matrix $D_aF(a)$ was square. A natural next step would be to relax this constraint and study the use of the Jacobian prior with more observation times. In this case, a restriction of $F$ may need to be imposed.

Finally, we note that the numerical approximation of the Jacobian determinant increases computation time in the algorithm. For example, one iteration of the Metropolis-Hastings algorithm with the Jacobian prior averaged 0.0225 seconds, while with the uniform and Jeffreys priors, it was 0.01 sec.
5.0 CONCLUSIONS

In this work, several aspects concerning the estimation of parameters for dynamical systems using single trajectory data were addressed. The first investigation of this dissertation confronted the question of identifiability. Specifically, we studied whether the parameters of a linear, or linear-in-parameters, dynamical system can be uniquely determined from a single error free trajectory of data. Several equivalent characterizations of the identifiability criterion, were presented. For linear systems, this yielded an identifiability condition solely based on geometric properties of the known trajectory, namely whether or not the trajectory is confined to a proper subspace of the phase space. This criterion provides practical utility, since it can be applied using what is known about the trajectory, without knowledge of the structure of the parameter matrix. Several examples were presented to illustrate identifiability for various systems and initial conditions. Additional results for linear systems include a link between identifiability from single trajectory with initial condition \( b \) and the linear independence of \( \{b, Ab, ..., A^{n-1}b\} \), several characterizations of the linear independence of \( \{b, Ab, ..., A^{n-1}b\} \) including a condition on the Jordan form of \( A \). A straightforward extension was made to the case of linear-in-parameters systems, where now the geometry of the image of the orbit in the flux space determined identifiability. Although a further extension to the general class of nonlinear systems of differential equations would be desirable, the lack of a unifying parameter structure within the class of these models would cause the need for a different analytical approach than ours. We then shifted our focus to investigate identifiability from a discrete collection of points along the trajectory and showed that some linear models are identifiable from a complete trajectory but not from a finite set of data. Other works which investigate identifiability of linear dynamical systems, typically include an input which serves as a control for the system. Identifiability is then addressed with full access to
the set of all admissible controls. Our treatment is more geared toward the case of single trajectory data. Without access to a set of controls, resulting in a variety of observations for the system, we have only a single initial condition, and thus one trajectory, from which to determine the parameters. This work complements this existing literature and approaches the identifiability problem in a uniquely different way.

We continued with the theme of discrete data in Chapter 3, where we investigated the robustness of the solution to the inverse problem. By considering a collection of data observed at equally spaced time intervals along a trajectory, we sought to define the largest allowable uncertainty in the data that could be tolerated such that properties of the inverse solution, such as uniqueness and equilibrium stability, were maintained. We first proved that such properties could be maintained in a neighborhood of the data, and subsequently worked to define bounds on the maximal permissible uncertainty. Three types of bounds on uncertainties were presented: analytical lower bounds, below which properties are guaranteed to hold for all perturbations of data; analytical upper bounds, which provide proven perturbations of data for which properties are guaranteed to be lost; and numerical bounds, derived from direct sampling of data points. Our results indicate that the upper bounds, when optimized over all potential eigenvalues, provide excellent agreement with the numerical estimates. All three of the bound types are hypothetically applicable to systems of arbitrary size; however, the numerical and analytical upper bounds become computationally expensive for larger systems. Although the analytical lower bound significantly underestimates the maximal permissible uncertainty, it provides a bound that is immediately accessible for systems of higher dimension, without increased computation. Further analysis is needed to determine whether a more accurate analytical lower bound can be obtained using structured perturbations. Several illustrative examples were presented and additional extensions were developed for the case of two-dimensional linear systems. Additionally, the ability to define the set, $C(d, \epsilon_U)$, on which a unique solution to the inverse problem is guaranteed at each point, became a vital tool in the analysis conducted in Chapter 4. The results of this chapter provide a useful contribution to identifiability in parameter estimation. Often, a model can be determined to be identifiable either globally in all of parameter space or locally, in a neighborhood of a point in parameter space. Defining the set $C(d, \epsilon_U)$ and applying the
inverse map, provides an explicitly defined set in parameter space on which the model is identifiable, that is neither an arbitrarily small neighborhood or the entire parameter space. Additionally, our results also include bounds on regions of data space where the inverse problem cannot be solved. The utility of such results is that they can provide an approach for model rejection.

In Chapter 4, our focus shifts to practical aspects encountered in estimating the parameters of a dynamical system from a collection of discrete single trajectory data. In this case, the parameter estimation problem is addressed by seeking an approximation to the parameter density, given in the Bayesian inference approach by a posterior density. A new informative prior was introduced as an alternative prior density to employ in the computation of the posterior with the Metropolis-Hastings algorithm. The Jacobian prior does not solely rely on previous information about the parameters, but instead, exploits the known structure of the differential equation model. Two systematic approaches were used to study which prior makes the posterior a better approximation of the parameter density. In each approach, a sample of the parameter density was constructed in order to compare with the posteriors obtained by implementing the Metropolis-Hastings algorithm with various priors. In the case of linear systems of ordinary differential equations, we found that the Jacobian prior consistently produced better approximations of the parameter density than both the uniform and Jeffreys priors. In our study using the nonlinear influenza model, the results were less conclusive, however in our examples, that the Jacobian prior performed at least as well as the uniform and Jeffreys priors and in a few cases, better. These findings will be submitted as a paper to a peer-reviewed journal. This chapter provides an important first step for introducing the use of the Jacobian prior in the Metropolis-Hastings algorithm, however further studies are needed. An analysis with additional nonlinear models where $F$ is known to be invertible is our next task. A deeper look into the requirement of the injectivity of $F$ and the importance of model identifiability in practice is needed. An extension to the case of a non-square Jacobian needs to be addressed. In this case, a restriction of $F$ to gain injectivity may be necessary, and defining the Jacobian as the product of nonzero eigenvalues, may be the correct formulation in this case.

Although the models employed in this dissertation were all systems of differential equa-
tions, when we consider the setting of discrete data sampled from a trajectory at a sequence of incremented time points, the systems are in fact discrete dynamical systems. There may be more existing theory for discrete dynamical systems which can be drawn from to extend some of our results to additional systems or classes of systems.

Throughout this work, linear systems of differential equations provided a convenient setting where properties of the principal matrix solution could be exploited to develop insightful theoretical results. The structure of these systems also allowed for an extension to nonlinear systems which are linear in parameters. Although a system of linear equations has a simple formulation as an ODE, the solution to the system is highly nonlinear in the parameters and thus provides a nontrivial investigation in the setting of parameter estimation. It is also the case that many biological systems are modeled using linear systems of differential equations. Although the analysis of Chapter 3 is exclusive to linear systems of differential equations, these results provided an important means to study the accuracy of the posteriors obtained in the Metropolis-Hastings algorithm. This algorithm can be used for parameter estimation for both linear and nonlinear differential equations. In biological modeling in particular, Bayesian inference provides a means to represent parameter variability in a population. Standard parameter fitting techniques result in a single optimal parameter value as a solution to the inverse problem; however, parameters with biological relevance are likely to differ between individuals. Ensemble models arising from estimation of parameter densities with Bayesian techniques are highly useful for biological modeling. Due to the wide use and popularity of the Metropolis-Hastings algorithm for Bayesian parameter inference, an advancement in improving the accuracy of the algorithm, I believe, would be the most impactful contribution of this dissertation to the modeling community.
DETERMINING THE NUMBER OF REAL DISTINCT ROOTS OF A POLYNOMIAL

A method for determining the number of real distinct roots of a polynomial was presented by Yang [91], based on results of Gantmacher [27]. For a polynomial

\[ p(t) = y_1 + y_2 t + \ldots + y_n t^{n-1} - t^n, \quad (A.1) \]

one defines the \((2n+1) \times (2n+1)\) discrimination matrix

\[
\Delta = \begin{bmatrix}
-1 & y_n & y_{n-1} & y_{n-2} & \ldots & y_1 & 0 & 0 & 0 & \ldots & 0 \\
0 & -n & (n-1)y_n & (n-2)y_{n-1} & \ldots & y_2 & 0 & 0 & 0 & \ldots & 0 \\
0 & -1 & y_n & y_{n-1} & y_{n-2} & \ldots & y_1 & 0 & 0 & \ldots & 0 \\
0 & 0 & -n & (n-1)y_n & (n-2)y_{n-1} & \ldots & y_2 & 0 & 0 & \ldots & 0 \\
\vdots & & & & & & \ddots & & & & \\
0 & 0 & \ldots & 0 & 0 & -1 & y_n & y_{n-1} & y_{n-2} & \ldots & y_1 \\
\end{bmatrix} \quad (A.2)
\]

comprised of alternating rows of two types.

Let \(D_k = \det(\Delta_{1:2k,1:2k})\) for \(k = 1, \ldots, n\) where \(\Delta_{1:2k,1:2k}\) is the upper left \(2k \times 2k\) sub-matrix of \(\Delta\). Define \(s_k = \text{sign}(D_k)\) and call \(S = [s_1, s_2, \ldots, s_n]\) the sign list. If, within \(S\), there is some subsequence \([s_{i+1}, s_{i+2}, \ldots, s_{i+j-1}]\) of zero entries, with \(j \geq 3\) and with \(s_i, s_{i+j} \neq 0\), replace \([s_{i+1}, s_{i+2}, \ldots, s_{i+j-1}]\) with \([-s_i, -s_i, s_i, -s_i, \ldots]\). After all such replacements are made, the resulting list is termed the revised sign list.
Using the revised sign list, we can determine the number of real distinct roots of $p(t)$ from the following theorem.

**Theorem 43** (Yang). The number $\nu$ of sign changes in the revised sign list equals the number of pairs of complex conjugate roots of $p(t)$. Furthermore, if the number of non-zero entries in the revised sign list is $l$, then the number of distinct real roots is $l - 2\nu$.

Hence, if the revised sign list has no sign changes and no zero entries, then the roots of $p(t)$ are all real and distinct.

In addition to Yang’s theorem, Descartes’ rule of signs can be used to determine whether the roots are positive. That is, given that the roots are all real, Descartes’ rule of signs with the criterion that the maximum number of negative roots is zero gives necessary and sufficient conditions for positivity of the roots. Namely, if $y_k > 0$ for $k = n, n - 2, n - 4, \ldots$ and $y_k < 0$ for $k = n - 1, n - 3, n - 5, \ldots$ for $y_k$ as defined in equation (A.1), then the roots of $p(t)$ are real, positive, and distinct.

**A.0.1 2 × 2 Case**

The polynomial $p(t) = y_1 + y_2 t - t^2$ has discrimination matrix

$$
\Delta = \begin{bmatrix}
-1 & y_2 & y_1 & 0 & 0 \\
0 & -2 & y_2 & 0 & 0 \\
0 & -1 & y_2 & y_1 & 0 \\
0 & 0 & -2 & y_2 & 0 \\
0 & 0 & -1 & y_2 & y_1
\end{bmatrix}
$$

Here $D_1 = 2 > 0$, $D_2 = y_2^2 + 4y_1$, and $S = [1, \text{sign}(D_2)]$. The revised sign list will have no sign changes and no zero entries as long as $y_2^2 + 4y_1 > 0$. In order to ensure that the roots are positive, Descartes’ rule of signs requires $y_2 > 0$ and $y_1 < 0$.

In summary, $p(t)$ has distinct, real, positive roots iff $y_2 > 0$ and $-y_2^2 < 4y_1 < 0$. 

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A.0.2 3 × 3 Case

The polynomial \( p(t) = y_1 + y_2t + y_3t^2 - t^3 \) has discrimination matrix

\[
\Delta = \begin{bmatrix}
-1 & y_3 & y_2 & y_1 & 0 & 0 & 0 \\
0 & -3 & 2y_3 & y_2 & 0 & 0 & 0 \\
0 & -1 & y_3 & y_2 & y_1 & 0 & 0 \\
0 & 0 & -3 & 2y_3 & y_2 & 0 & 0 \\
0 & 0 & -1 & y_3 & y_2 & y_1 & 0 \\
0 & 0 & 0 & -3 & 2y_3 & y_2 & 0 \\
0 & 0 & 0 & -1 & y_3 & y_2 & y_1
\end{bmatrix}
\]

Here \( D_1 = 3 > 0 \), \( D_2 = 2y_3^2 + 6y_2 \), \( D_3 = -27y_1^2 - (4y_3^3 + 18y_2y_3)y_1 + 4y_2^3 + y_2^2y_3^2 \), and \( S = [1, \text{sign}(D_2), \text{sign}(D_3)] \). The revised sign list will have no sign changes and no zero entries as long as \( D_2 > 0 \) and \( D_3 > 0 \). In order to ensure that the roots are positive, Descartes’ rule of signs requires \( y_3 > 0 \), \( y_2 < 0 \), and \( y_1 > 0 \).

In summary, \( p(t) \) has distinct, real, positive roots iff \( y_3 > 0 \), \(-y_3^2 < 3y_2 < 0\), and \( \max(0, -b - 2\sqrt{C}) < 27y_1 < -b + 2\sqrt{C} \), where \( b = 2y_3^3 + 9y_2y_3 \) and \( C = y_3^6 + 9y_2y_3^4 + 27y_2^2y_3^2 + 27y_2^3 \).
APPENDIX B

KERNEL DENSITY ESTIMATION

For a finite sample of observed data, density estimation constructs an estimate of the underlying probability density function everywhere, including where no data are observed. Histograms are one of the most simple density estimators; however, they are not smooth and the choice of bin width leads to a constant struggle between bias and variance [72]. Kernel density estimation is a more sophisticated alternative. In this approach, a kernel function is centered at each data point, so that a discrete data point is smoothed over the region surrounding it. The kernels are then summed, resulting in a single function that is the estimated density function.

A kernel density estimation in one dimension is depicted in Figure 36 (a). The discrete collection of data points are plotted as stars along the x-axis and the individual kernels are pictured in blue. The density estimate resulting from summing these kernels is represented by the black curve. The case of kernel density estimation in two dimensions is pictured in Figures 36 (b) and (c) as presented in [72]. The contours represent the individual kernels surrounding the data points in part (b), and the resulting density estimate in part (c).

The mathematical formulation for the multivariate kernel density estimation is as follows: Given n data points \( w_1, w_2, \ldots, w_n \) in \( \mathbb{R}^d \), the estimated density function is given by

\[
\hat{f}_H(w) = \frac{1}{n} \sum_{i=1}^{n} K_H(w - w_i).
\]
The bandwidth matrix, $H \in \mathbb{R}^{d \times d}$, is symmetric positive definite.

$$K_H(w) = |H|^{-1/2} K(H^{-1/2}w),$$

where $K(w)$ is the kernel function. The standard multivariate normal kernel $K(w) = (2\pi)^{-d/2} \exp(-\frac{1}{2}w^Tw)$ is often employed, however other kernel functions are also used. Selection of the bandwidth matrix is the most important task as it controls the orientation and the amount of smoothing and thus the accuracy of the density estimate [71].
BIBLIOGRAPHY


