# THE EFFECT OF TERM ORDER ON IDENTIFYING MODELS OF GENE REGULATORY NETWORKS

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ABSTRACT. Gene regulatory networks (GRNs) control many life processes. An important problem in systems biology is to identify a model of the network for a given set of laboratory data. A recently developed method builds a space of polynomial models from a given data set; however, it requires that an order on the variables (genes) and an order on the monomial terms (gene interactions) to be established. No rigorous analyses have been performed to determine the impact of these orders on the construction of the model space.

We chose a well-studied GRN in *E. coli* to study the effect of these orders on the identification of an existing model of the network. We also determined how much data, as well as which data explicitly are required for model identification.

#### 1. Introduction

Gene regulatory networks (GRNs) are fundamental in orchestrating life processes. Recent technology has allowed for amassing of large amounts of data collected from probing GRNs. A growing problem is to infer the network from the data. A recently developed method constructs the space of polynomial models that fit the data. An advantage of the model space is that it permits further analyses; however, currently no extensive analyses have been performed. As the modeling approach imposes an order on the variables (genes) and an order on the monomial terms (gene interactions), rigorous analyses will help to determine the impact of these orders on the construction of the model space.

We highlight some results from exploring the relationship between the data and the model space. In particular we computed the size of data sets necessary to guarantee a unique model in the space. Additionally we identified which data sets specifically guarantee a unique model. As a consequence, we have established the hypothesis that the choice of variable order has a greater impact on modeling than the term order. Solving this open problem will make significant contributions towards refined experimental design and more predictive models of GRNs.

In the following sections, we use the well-studied model of the *lac* operon to determine the minimal size of data, and which data, are necessary to reversely engineer the correct model. We also look at the sufficiency of the data size, or more explicitly, the number of data which always guarantees the return of the correct model.

# 2. Case Study: The lac Operon

The lac operon is a network of genes that controls the metabolism, or breakdown, of lactose in  $E.\ coli.$ 

FIGURE 1. The lac operon.

FIGURE 2. Wiring diagram of the model F.

FIGURE 3. The state space of the model F.

We used a simplified version of a published Boolean model of the *lac* operon [?], in which glucose concentrations are fixed.

## 3. Preliminaries

3.1. **Models of GRNs.** A polynomial model for a GRN is a collection of polynomials, in which the behavior of each gene is described by a polynomial function.

The model for the *lac* operon is given by  $F = (f_M, f_L, f_G)$  where

$$\begin{array}{rcl} f_M & = & L \vee L_e \\ f_L & = & M \\ f_G & = & 1. \end{array}$$

Using the translation rules between Boolean functions and polynomial functions

$$x \lor y = xy + x + y$$
$$x \land y = xy$$
$$\neg x = x + 1$$

as well as the representation of  $M, L, L_e, G$  with the variables  $x_1, x_2, x_3$  respectively, then the coordinate functions of F can be rewritten as

(3.1) 
$$f_1 = x_2x_3 + x_2 + x_3$$

$$f_2 = x_1$$

$$f_3 = 1$$

where each  $f_i \in \mathbb{Z}_2[x_1, x_2, x_3]$ .

We simulate the GRN by evaluating F on all possible inputs.

Probing a GRN typically produces incomplete data sets. It is a problem to identify the model from these limited data.

3.2. Model Building and Selection Using Algebraic Geometry. The algebraic method in [?] computes all models that fit given data.

We define the *model space* to be the collection of functions of the form

$$F_p + F_h$$

where  $F_p$  is a particular function that fits the data, and  $F_h$  is any function that is zero on the data. If  $I = \{\text{all possible } F_h\}$ , then

$$F_p \mod I := \text{remainder of } F_p \text{ upon division by all } F_h$$

is a "minimal" model that fits the data. However, polynomial division is well defined only if I is written as a Gröbner basis.

All definitions are taken from [?].

FIGURE 4. Two sample data sets which return the model F.

FIGURE 5. Two sample data sets which return the model F.

**Definition 3.1.** A *Gröbner basis* is a multivariate nonlinear version of Gauss-Jordan Elimination and requires a canonical way to write polynomials.

**Definition 3.2.** A variable order (VO) is a sorting of variables, and a monomial order (MO) is a rule for comparing monomials. A monomial order requires a variable order.

**Example 3.3.** If x > y, then  $xy > y^3$  in lexicographic MO, i.e. dictionary order, while  $y^3 > xy$  in graded lexicographic MO, in which total degree is counted first. If y > x, then  $y^3 > xy$  in both monomial orders.

#### 4. Methods

We considered all possible VOs on 3 variables (3! = 6). For each of these VOs, we used the MOs lex, glex, and grevlex. As I is computed from data, we created all subsets of the complete set of  $2^3 = 8$  data points (see Section 3.1). We computed minimal models using each of the  $2^8 = 256$  subsets and reported which data sets, MOs, and VOs result in the original model F. The algorithms were executed in Macaulay2 [?].

# 5. Results

- (1) Any subset of 7 data points is sufficient to return F.
- (2) A minimum of 5 data points is necessary to return F.
  - (a) Only 32 of the 56 possible subsets of 5 data points return F (two such sets shown below).
  - (b) In these 32 subsets, each data point appears with the same frequency (20).
- (3) Each MO produced identical Gröbner bases, resulting in identical minimal models, though not always F.
- (4) Only 2 VOs returned F.

## 6. Analysis

6.1. **Identifying Minimal Data Size.** To understand why the model requires 5 points as the minimal data size, we will refer to the following result by Robbiano.

**Theorem 6.1** (Robbiano[?]).  $V \subseteq k^n$  identifies a polynomial f if and only if the evaluation matrix X(V, f) has full rank.

We will also need to define the support and closure of a polynomial.

**Definition 6.2.** Let f be a polynomial. The *support* s(f) of  $\underline{f}$  is the collection of monomials  $\{x^{\alpha_1}, \ldots, x^{\alpha_m}\}$  that appear in f. The *closure*  $\underline{s(f)}$  of s(f) is the set  $\{x^{\beta}: x^{\beta} \mid x^{\alpha}, x^{\alpha} \in s(f)\}$  of monomials that divide elements of s(f). These monomials are also referred to as *standard monomials*.

Before we can prove the minimal data size for an F collection of polynomials, we need to look at the minimal data size for one polynomial f.

**Lemma 6.3.** Let f be a polynomial, s(f) its support, and  $m = |\overline{s(f)}|$ . Then there exists  $V \subseteq k^n$  that identifies f and m is the smallest integer such that |V| = m.

Proof. Let f be a polynomial and s(f) its support. Let  $m = |\overline{s(f)}|$ . Then there exists  $V \subseteq k^n$  (could be all of  $k^n$ ) such that V identifies f. By Theorem 6.1, X(V, f) has rank  $m \leq |V|$ . Since X(V, f) is potentially rectangular, we know we can find a square submatrix of the same rank. So there exists  $V' \subseteq V$  such that |V'| = m. Hence there is a subset V' such that X(V', f) has rank m and thus identifies f by Theorem 6.1.

To show that m is the smallest such integer, suppose that there is  $W \subseteq k^n$  that identifies f with |W| < m. Then X(W, f) has rank m', which is strictly less than m. Again, we can find a square submatrix of rank m'. However this means that one of the monomials in s(f) is not identified. Hence, by Theorem 6.1, W does not identify f, a contradiction to the initial assumption. Therefore, m is the smallest such integer.

Now that we know the minimal data size for a single polynomial f is equal to the number of standard monomials, let's consider a system of polynomials  $F = f_1, \ldots, f_n$  like the one we used for the *lac* operon. We then have the following definitions.

**Definition 6.4.** Let  $F = (f_1, \ldots, f_n)$  be a PDS and  $V = \{p_1, \ldots, p_m\}$  a subset of point in  $k^n$ . V identifies  $f_i$  if the coefficients of  $f_i$  can be found uniquely by evaluation of the monomials of f on the points in V. Furthermore, V identifies F if V identifies each  $f_i$ .

**Definition 6.5.** Let  $F = (f_1, \ldots, f_n)$  be a polynomial dynamical system and  $V \subseteq k^n$ . Let  $X(V, f_1, \ldots, f_n)$  be the evaluation matrix of the monomials in the support of each  $f_i$  on the points in V. In short, we write X(V, F).

We now try to extend Robbiano's result for a single polynomial to a system of polynomials.

**Theorem 6.6.**  $V \subseteq k^n$  identifies a polynomial dynamical system  $F = (f_1, \ldots, f_n)$  if and only if the evaluation matrix X(V, F) has full rank.

*Proof.* Let  $V \subseteq k^n$ ,  $F = (f_1, \ldots, f_n)$  a PDS with  $s(f_i)$  being the support of each  $f_i$ , and  $m_i = |\overline{s(f_i)}|$ .

Suppose X(V, F) has full rank. Then X(V, F) has full column rank. Since  $m_i = |\overline{s(f_i)}|$ , then  $X(V, f_i)$  has rank  $m_i$ . So by Theorem 6.1, V identifies  $f_i$ . By definition, V identifies F.

Suppose V identifies F. So V identifies each  $f_i$ . So by Theorem 6.1,  $X(V, f_i)$  has rank  $m_i$ . Since  $\bigcup s(f_i)$  contains m linearly independent monomials (as they are standard monomials), then the rank of X(V, F) = m.

Using the above theorem, we can now prove that the smallest size of V is equal to the number of standard monomials in F.

**Theorem 6.7.** Let  $F = (f_1, \ldots, f_n)$  be a PDS,  $s(f_i)$  the support of each coordinate function  $f_i$ , and  $m = |\overline{\bigcup_{i=1}^n s(f_i)}|$ . Then there exists  $V \subseteq k^n$  that identifies F and m is the smallest integer such that |V| = m.

Proof. Suppose  $F = (f_1, \ldots, f_n)$ ,  $s(f_i)$  the support of each  $f_i$ ,  $m = |\overline{\bigcup_{i=1}^n s(f_i)}|$ , and  $m_i = |\overline{s(f_i)}|$ . So  $s(f_i) \subseteq \overline{\bigcup_{i=1}^n s(f_i)}|$ , implying that  $m_i \le m$ . By Lemma 6.3, there exists  $V_i$  such that  $m_i$  is the smallest integer where  $V_i$  identifies  $f_i$  and  $|V_i| = m_i$ . Let  $V = \bigcup_{i=1}^n V_i$ . By construction, V identifies F since a subset of V identifies  $f_i$ . If |V| = m, then we are done. Suppose |V| < m. Then X(V, F) has rank m', which is strictly less than m. But by Theorem ??, if V identifies F, then X(V, F) must have rank m, a contradiction. Now suppose that |V| > m. So we can find  $V' \subset V$  such that X(V', F) has rank m. By Theorem ??, V' identifies F.

To show that m is the smallest such integer, suppose that there is  $W \subseteq k^n$  that identifies F with |W| < m. Then X(W, F) has rank m', which is strictly less than m. Again, we can find a square submatrix of rank m'. However this means that one of the monomials in s(f) is not identified. Hence, by Theorem 6.6, W does not identify F, clearly a contradiction to the initial assumption. Therefore, m is the smallest such integer.

Returning to our case study of the *lac* operon model, since F has *closure*  $\overline{s(f)} = \{1, x_1, x_2, x_3, x_2x_3\}$  and  $|\overline{s(f)}| = 5$ , then 5 data points are necessary to identify the model.

#### 7. Discussion

The results suggest that the choice of variable order has a greater impact on identifying models than the monomial order. Solving this open problem will make significant contributions towards refined experimental design and more predictive models of GRNs. To test the validity of the new hypothesis, further studies of monomial orders and models of higher degree will be conducted.

## 8. Acknowledgements