Plan for my Research at INRA in Montpellier during July 2012- July 2013

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May 27, 2011

My research over the last ten years has focussed largely on the application of computer science to the life sciences. Most of that work has had to do with systematic experimental design for biologists, visualization software, and machine learning tools. The current thrust of my work has to do with large scale inference and analysis of networks. My plan in Montpellier is to continue that work. This will lead to a tool that may be useful to the analysis of the data generated by the Integration of Nutritional Functions team (and possibly others) at INRA.

1 Brief Review of the Most Relevant of My Previous Work

My work in scientific computing is driven by the philosophy that the problems and questions should come from the lab scientists themselves. I attempt to solve those problems in a way that the scientist can use and to generalize that solution as much as possible. The following branches of my work all follow from that philosophy.

1. Adaptive Combinatorial Design for the Design of Experiments Lab scientists would like to answer a scientific question as quickly (in person time) and as economically (in lab equipment and materials) as possible. A typical "search space" in a lab setting will include many possible perturbations (e.g. light, carbon, nitrogen, knock-outs,...) and the goal is to find the values of those purturbations that optimize a particular output (e.g. biomass, seed size). The expensive approach is to explore the entire search space. An alternative is to design a small number of experiments and then to use the results of those experiments to design the next group with a view towards finding the optimal conditions very rapidly. We have used and improved a technique from statistics called combinatorial design to this end[24]. Our basic strategy is to use combinatorial design (i) to design a well-spaced and very small set of initial experiments and (ii) to use the results of that first set of experiments to design a second set of experiments that focusses on the features that seem most influential. Thus, combinatorial design is used "recursively" to find the optimal values of the influential features and the other features. We have used this successfully in our plant biology group at NYU, but the method has been used by collaborators looking at bacteria and bioenergy.

2. Visualization of Multiple Experiments

A frequent genomics question is "Which genes are most affected by all of these experiments or a subset of those experiments?" A common way to appreciate this visually is to use a Venn diagram. Because Venn diagrams generalize poorly beyond three experiments, we have developed a visual representation known as a Sungear (http://virtualplant.bio.nyu.edu/cgibin/sungear/index.cgi) that does generalize. Sungear is an interactive search interface that supports statistical conclusions and that is particularly strong in performing metaanalyses of many different experiments[22], [21]. For example, a sister lab is using Sungear for cancer studies. It is a general tool and we have seen applications ranging from science to marketing to the evaluation of sports teams.

3. Data Analysis to infer gene or module function

Most of my work with Gloria Coruzzi, Ken Birnbaum, and Phil Benfey over the years has had to do with data analysis to discover gene function. Sometimes, this has meant the inference of the individual or combinatorial genetic causes of traits[26], sometimes in a cell-specific manner[25]. Most frequently though, the idea has been to take a holistic "systems biology" approach to try to understand the role of modules of genes, often using machine learning [23], [19], [18], [16], [13]. Many of the tools we have developed are now incorporated into our system Virtual Plant[15].

4. Network inference in genomic networks

The goal in this work is to determine which genes influence which other genes and the strength of those relationships. In the case of genomics, the experimental strategy consists of measuring the effect of perturbations (such as the introduction of stress, genetic change, or the insertion of nutrients) to organisms over time. The end result is a network that predicts causal relationships among genes. We have just begun that work [6, 4] and will outline our plan to continue it below.

5. Subgraph queries

Related to the question of network inference is what to do when one has a large network or several large networks and one wants to find common motifs. The paradigmatic question is "where is a certain labeled query graph q in a large database D of graphs?" Our fundamental strategies have been to use filters to prune away graphs from D that cannot match q and then to use a location data structure to find good starting points for searches in the graphs of D that remain. We have explored several variants of this problem [7, 9, 20] Another question has to do with clustering graphs that are similar.[17]

6. Time series analysis to find correlations and bursts among tens of thousands of time series over sliding windows

A paradigmatic problem in this area is to find highly correlated instruments in financial markets, where correlations can come and go over time. The two central techniques are to use dimensionality reduction techniques such as wavelets and sketches (random vectors) to avoid comparing all pairs of instruments and to update previous correlations efficiently. This work does not directly relate to our plant biology work, but could be used for other scientific applications having far longer time series [3, 11, 12].

2 The Basic Plan: Scalable Network Inference on a Workflow Platform

In our review of genomic network inference algorithms [29, 6, 32, 35, 38, 39], we have observed that there are several stages of analysis depending on the kind of data that is available. We can divide those data types along two

dimensions: (i) whether that data is generated based on genetic perturbations or not; and (ii) whether that data consists of steady state data or time series data.

Example of genetic perturbations include the suppression of gene function or its enhancement, whether cell-specific of not. In a non-genomic setting, the equivalent of a genetic perturbation is any direct modification of a node in a network. Non-genetic perturbations, by contrast, are analogous to manipulations of the inputs of networks.

What constitutes time series data depends on the system under examination. For example, formally, measurements taken every 4 hours constitute a time series. For our purposes, however, time series data means a series of experiments having the property that the state at experimental time point k+1 depends on the state at time point k but not on the interaction among data components at k+1. For us, then, for a series of experiments to be considered a time series, causality edges flow from the state of elements (e.g., genes) at k to elements at k+1. Whether this is true for measurements taken every four hours or not will depend on the rate of reaction of elements in the organism under study. We might call the kind of time series we are interested in *causal time series*.

From steady state experiments, algorithms can derive correlation, clusters, and biclusters[31, 33, 37] Clustering reduces the number of nodes to a small group of "super-nodes" that rise and fall together. From genetic perturbations, one can determine the direct or indirect influence of a perturbed element on others.[29, 30, 39] From causal time series data, one can determine causal links – if there is enough data relative to the number of super-nodes.[34] The overall workflow can be represented in following Vistrails figure [36]

Whereas this offers a systematic way to infer networks, the resulting networks are not always so good. In our own work for example[6], we were able to predict the direction of gene expression (whether expression rose or fell) on out-of-sample data quite accurately, but not its magnitude. The main open problems in my view then are to (i) improve the quality of the algorithms given the data available and (ii) to determine which next experiment to do to improve the prediction accuracy.



Figure 1: Steps of the network inference workflow: each step contains several optional algorithms and many parameters for each algorithm

2.1 Algorithms

There are two issues regarding algorithms: quality and speed. They are related, because faster algorithms make it possible to search more parameter combinations and thus achieve better quality.

The quality of a network inference workflow depends on the algorithms chosen and the parameters fed to those algorithms. For network inference the main algorithms are Inferelator 2.0[29, 34], ARACNE[40, 32], TSNI[35], BANJO[38], and NIR[39]. Different authors claim that each algorithm is best overall, but we suspect that each has a "sweet spot" which we must find. Finding the best parameters on the other hand will require an exploration of the parameter space. For this, we will use genetic algorithms[41, 42] in combination with combinatorial design. Here is where quality and speed interact: fast algorithms permit more exploration of the parameter space.

The speed issue comes up when the inference problem concerns large networks. The core problem concerns the identification of edges that could cause changes in the value of a target element and assigning values to them. This is essentially a regression problem. Luckily, there has recently been a flurry of excellent work on machine learning algorithms which are both sufficiently fast and parallelizable to be used on data sets with milions of elements. Below, we list a few of the algorithms we think might useful in training regression models on large-scale data.

• Random Forests [2]

Random forests are ensembles of decision trees which are constructed from random subsets of the data. They're fast to train, easy to parallelize, and perform extremely well.

• Large-Scale SVM Regression [1]

Bottou demonstrated that a stochastic gradient descent solver for a variety of learning problems (including support vector machine optimization) is able to scale with extremely large datasets while converging to the predictive performance of traditional optimization algorithms.

• Large-Scale ℓ_1 Regularized Learning [10]

Stochastic coordinate descent can be used to learn sparse regression models, with small training times even for data sets where both the dimensionality and the number of training points is large.

2.2 Experiments to Do Next

Regardless of the quality of algorithms, insufficient or excessively noisy data can prevent an algorithm from inferring good networks. Because experiments take time and expense, we want to guide the experimenter to do the "right" experiment. One way to do this is to determine which existing experiment has been most valuable and doing another one like that. To determine the value of an existing experiment, one can remove that experiment, rerun the inference algorithm and then re-compute prediction accuracy.

For example, in the case of our Arabidopsis time-course study[6], removing two replicate experiments from two different timepoints prior to 15 minutes was less harmful to the accuracy of out-of-sample prediction of the network state at 20 min than removing both replicates from a single time-point prior to 15 min. This suggests that measurements at different time-points may be more valuable than replicates.

Whereas this rather naive approach may work well in some cases, it will not lead to radically different experimental designs. One way to discover better designs will be to simulate the data under different noise and variance assumptions. Given such simulation results, one should be able to approach a given application, characterize its noise and variance properties, and design a series of experiments.

2.3 The Importance of Workflow

Workflow systems help solve two important problems in scientific computation:

- Just as it is important for experimental procedures to be repeatable so it is important for computational procedures to be repeatable. I have been active in urging computer scientists in the large database community to create repeatable experiments[47, 46] This year we featured the use of Vistrails to help make this possible (http://www.sigmod2011.org/calls_papers_sigmod_r By storing workflows along with associated software and data, experimenters ensure that a whole computational flow can be reproduced.
- 2. Workflow systems support a disciplined appoach to parameter exploration. For example, Vistrails will soon have a genetic algorithms module so that parameter values can be varied and optimized.

For these reasons, our scalable network software will be wrapped in a workflow system, probably Vistrails to start.

3 Expected Results

Over the course of my sabbatical year (July, 2012 to July, 2013), I expect to design and build useful network inference software and apply it to important problems, I also intend to keep my eyes and ears open as new problems come to my attention. In my previous sabbaticals at INRIA Rocquencourt, I consistently found that interactions with colleagues led to new research directions and excellent publications[43, 44, 45]. I have every expectation that serendipity will play an equally positive role this time.

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