Introduction

To explain how a complex trait works, systems biology begins with organizing macromolecules into a genetic network [1]. The biological clock is an example of how a complex trait with numerous pleiotropic phenotypes can emerge from the interaction of only a few regulatory macromolecules. For two reasons much of what we know about the clock at the molecular level comes from the study of the filamentous fungus, Neurospora crassa [2]. First, this complex trait is easy to observe and to manipulate in N. crassa (Fig. 1). Second, as a well-studied microbial system, it has been possible to identify three molecular building blocks of the clock, the genes white-collar-1 (wc-1) and white-collar-2 (wc-2), and frequency (frq). The genes wc-1 and wc-2 encode PAS-domain containing transcription factors [3] that turn on the clock oscillator. The WC-1 protein also acts as a blue-light receptor [4]. The gene frq encodes the clock oscillator FRQ [5] and is activated by the WHITE-COLLAR transcription factor protein complex WCC = WC-1/WC-2. The FRQ protein in turn appears to function as a cyclin to recruit an as yet to be identified kinase/phosphatase pair for the phosphorylation-dependent inactivation of WCC [6].

This information enabled formulation of the detailed genetic network shown in Fig. 2 that explains how the clock functions [7]. In this network model, the WCC protein activates the oscillator gene frq. The active frq+ gene is then transcribed into its cognate mRNA frq1, which in turn is translated into its cognate protein FRQ. The FRQ protein, in turn, deactivates the WCC in the P reaction. FRQ thereby closes a loop of dynamical frustration wherein WCC turns on the oscillator gene whose product shuts down the activator WCC. This dynamical frustration (i.e., negative feedback loop between WCC and FRQ) explains in part how clock oscillations arise [7]. In addition, WCC activates a number of clock-controlled genes (ccgs) that serve as outputs of this clock mechanism. The number of these ccgs in the genome, and hence the extent of clock control over metabolism, is largely unknown but see [8].

Our goal is to refine systematically the genetic network model of the clock mechanism [7] and to explore the metabolic context of the clock. To achieve this goal Locke et al. [9] proposed using an iterative process of modeling and experimentation to identify and validate genetic networks. Along these lines, we introduce a model-driven discovery process called Computing Life in Fig. 3 [1,10]. In this paradigm, a cycle of modeling and genomics experiments are used to identify and, with each cycle, tighten our estimates on model parameters and on model predictions for the biological clock. The biological system is first perturbed. Measurements on all relevant species are made by RNA and protein profiling [1]. An ensemble of genetic network model parameters is generated for the
process of interest [11–12,7]. Predictions are made from the model ensemble and compared with available data. Revision of the model then poses the difficult choice of what perturbation experiment is to be done next to improve maximally our knowledge of the genetic network?

One approach to the problem of “informative” experiment design has been to assume that genetic networks are in steady state and/or are linear, and under these conditions predictions are made about the next round of perturbations [13–16]. This cannot be done here because the biological clock is usually not in steady state but rather approaching a stable limit cycle [17]. Also, the steady-state approach discards most information contained in observations on network dynamics, i.e., its time-dependent behavior. Another approach is to generate an entire compendium of profiling experiments for varied genetic and environmental perturbations [18]. Such profiling experiments are costly, however, and it is desirable that every experiment, at each stage, be maximally informative about the underlying genetic network. Here, we describe how this process of choosing the Maximally Informative Next Experiment (or MINEing) can be guided by the continuously refined network model in an intelligent and cost-

Figure 1. The clock of *N. crassa* is remarkably adaptive in its entrainment to varied artificial days. Replicate race tubes are inoculated at one end and subject to a 6 hr, 18 hr, and 48 hr artificial day over 7 ordinary days. The clock is manifested by the appearance of orange bands (i.e., asexual production of spores) as the culture grows to the other end of the tube. In each artificial day the race tubes experienced (A) 3 hrs light and 3 hrs dark, (B) 9 hrs light and 9 hrs dark, or (C) 24 hrs light and 24 hrs dark. It can be seen that the number of conidial bands tracked the number of artificial days experienced. Race tubes were prepared as described in [30] and in Materials and Methods and were inoculated using the *bd* mutation (FGSC 1858).

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Figure 2. A genetic network for the biological clock from [7]. Molecular species (i.e., reactants or products) in the network are represented by boxes. The white-collar-1 (wc-1), white-collar-2 (wc-2), frequency (frq), and clock controlled gene (ccg) gene symbols can be superscripted 0, 1, r0, r1, indicating, respectively, a transcriptionally inactive (0) or active (1) gene or a translationally inactive (r0) or active (r1) mRNA. Associated protein species are denoted by capitals. A photon (in yellow) denotes a photon species. Reactions in the network are represented by circles. Arrows entering circles identify reactants; arrows leaving circles identify products; and bi-directional arrows identify catalysts. The labels on each reaction, such as *S*0, also serve to denote the rate coefficients for each reaction. Reactions labeled with an *S, L, or D* denote transcription, translation, or degradation reactions, respectively. Reactions without products, such as *D*, are decay reactions. Reactions, such as *A* and *P*, have cooperative kinetics: (A) *n* WC-C-frq<sup>r</sup>-frq<sup>r</sup> and (P) WC-C-m FRQ→WC-2-m FRQ. The *n* and *m* are Hill coefficients or cooperativities. Only one reaction, the “A” reaction, has a back reaction, (7), frq<sup>r</sup>→*n* WC-C-frq<sup>r</sup>, included, with non-zero rate. The rate constants specify the right hand side of the kinetics model in equation (1) through the Law of Mass Action in Materials and Methods.

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Figure 3. Computing Life Paradigm. The “perturb” and “observe” steps represent the experimentation phase; the “fit”, “predict” and “evaluate” steps are the main components of the genetic network ensemble simulation phase; and the “select” step is the MINE design phase which closes the Computing Life workflow cycle.

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effective way while fully exploiting the information contained in the observed network dynamics. Our tracing of three cycles through the Computing Life paradigm in the context of refining our network model for the biological clock’s mechanism in *Neurospora crassa* illustrates this approach.

**Materials and Methods**

**Describing the genetic network**

All stages of the Computing Life paradigm in Fig. 3 involve the use of the genetic network. The methods of describing, fitting, predicting with, and evaluating the genetic network are first described, and then we continue to trace the methodology used to complete the cycle in Fig. 3, providing a methodological walk through the Computing Life paradigm in Materials and Methods.

**Kinetics model and the model ensemble.** The starting point for our MINE design approach is a kinetic rate equation model for the time-dependence of the molecular species concentrations in the network, based, e.g., on mass-action kinetics. The model in Fig. 2, for example, specifies a system of 16 ordinary differential equations (ODEs) that describe the temporal profiles of genes and their products. In general, these ODEs have the form:

\[
\frac{dX}{dt} = G(X,t; \theta, u)
\]

where \(X = [X_1, \ldots, X_N]^T\) is a \(N \times 1\) vector of species concentrations, with \(N\) denoting the number of molecular species evolving according to the kinetics rate equations; \(G = [G_1, \ldots, G_N]^T\) specifies the kinetics, i.e., \(G_n(X,t; \theta, u)\) is the net rate of production of species \(n\) at time \(t\), given the species concentrations \(X\). The model parameter vector or, for the short, “the model” \(\theta = [\theta_1, \ldots, \theta_M]^T\) in \(G\) is the \(M \times 1\) vector of all unknown model parameter variables, including, for example, unknown reaction rate constants, species’ initial concentrations and unit conversion factors. The rate functions \(G\) also depend on an array of “control variables” which are known and can be varied by the experimenter. These control variables specify, for example, the nature of the perturbations and external conditions to be applied to the biological system or, more generally, the experiment to be done. This array of control variables is denoted by a vector \(u\) (of unspecified dimension) and, for short, is referred to as “the experiment” in the following.

**Fitting, Predicting with, and evaluating ensembles of genetic networks**

Profiling experiments tend to generate data at only a few time points, the reaction networks are large, and their kinetics models are rich in unknown \(\theta\)-parameters. An ensemble method of genetic network identification [11,12,19,20,7] is therefore used to constrain the model parameters \(\theta\), using the model likelihood or some other criterion to select members of the model ensemble. The model ensemble \(Q(\theta)\) is a probability distribution on the parameter space of rate coefficients and initial species concentrations [21]. When viewed as a function of \(\theta\), the ensemble \(Q(\theta)\) can be the likelihood function. This model ensemble summarizes what we know and, equally importantly, what we do not know about the biological network, given the prior or “old” experimental data. We refer to Ref. [7] for a detailed description of the construction of \(Q(\theta)\) from prior experimental data and its numerical implementation by way of a Metropolis Monte Carlo ensemble simulation algorithm *ens.fld*.

With the ensemble in hand it is possible to make predictions from the ensemble means of the species concentrations, as shown in Results. It is also possible to take the expected species trajectories of the ensemble member \(\theta\) and compare them to the observed species trajectories using a figure of merit, such as the likelihood \(Q(\theta)\) or \(\chi^2 = -2 \ln Q(\theta) + \text{const}\), to evaluate goodness of fit of the ensemble as described in [11,7]. A direct graphical evaluation of the ensemble’s goodness of fit by can be assessed also by overlaying the observed trajectories of species concentrations onto the ensemble mean trajectories +/− the ensemble standard errors in these mean trajectories, as again shown in Results.

**Selecting an optimal perturbation**

The next stage in the Computing Life paradigm is selecting a perturbation in Fig. 3. We describe for the first time a novel method for selecting an optimal perturbation involving evaluating the Maximally Informative Next Experiment.

**MINE design as “microscopy” in model space.** For a given choice of model \(\theta\), let \(f(\theta, u)\) denote a kinetics model prediction for a single species log-concentration, \(\log(y_i)\), to be measured for a single time point by the next profiling experiment, i.e., \(y_i\) is one of the elements of \(X_i\) to be measured at some specific observation time \(t\). The vector \(u\), as explained above, comprises all control variables which are known and describe the externally imposed conditions of the experiment. However, \(u\) should now be understood also to comprise all control variables defining the specific data point \(y\) to be measured, including, for example, the choice of molecular species to be observed and the time of observation. We will need to generalize this notation when the planned next experiment measures multiple variables \(y_1, \ldots, y_d\). Let \(F(\theta, U)\) denote \(d \times 1\) vector of the corresponding predicted log-outcomes and \(U = [u_1, \ldots, u_d]\) the (super-)vector of corresponding control parameter vectors \(u_i\) where \(u_i\) specifies the control variables for the measurement of the data point \(y_i\) for \(i = 1, \ldots, d\). The log-variables to be observed, \(\log(y_i)\), will also be referred to, for short, as the “observables” in the following and \(U\), for short, as “the next experiment”. We are using log-concentrations instead of the concentrations themselves here, in order to obtain scale-free (i.e. concentration-unit-independent) MINE criteria, as explained below.

Clearly, the question of which next experiment \(U\) is “maximally informative” is not a mathematically well-defined problem. We have to make an *ad hoc* choice for a design criterion and then try it out in real-life applications. The basic conceptual ideas underlying this *ad hoc* construction of a MINE criterion are borrowed from microscopy: we want to use whatever experimental technique is available to us to “look into” or “image” the inner workings of the cell. A microscope generates images of the cell’s material components in three-dimensional *physical space* or in some lower-dimensional projection thereof. Profiling experiments, by analogy, generate images of the cell’s (very!) high-dimensional *kinetics parameter space*.

Ideally, we would like to be able to obtain highly resolved images, allowing us to determine accurately a genetic network’s “location” in kinetics parameter space, specified by a unique choice of parameter vector \(\theta\). Unfortunately, and again in analogy to microscopy, the images we do get from present-day profiling experiments do not allow us to completely re-construct \(\theta\): our “vision” in \(\theta\)-space is seriously blurred. The model ensemble \(Q(\theta)\) captures the constraints imposed on \(\theta\), i.e., what we know; but the spread of \(Q(\theta)\) within those constraints in \(\theta\)-space, also captures the blurring, i.e., what we do not know, given the prior experimental data. Our goal is therefore to reduce this blurring as much as possible when performing the next experiment: we want to tune our “model parameter microscope” to get a different view of \(\theta\)-space with the maximum possible resolution.

An important aspect to keep in mind here is that every imaging procedure, be it optical microscopy or RNA profiling, requires a
mathematical model which relates the observed image \( F \) to the underlying object \( \theta \). The mapping function \( F(\theta,U) \) captures that imaging model. Without such an imaging model, we cannot, for example, re-construct the shape, size and location \( \theta \) of a cellular organelle from the light intensity pattern \( F \) of the cell’s magnified image produced by an optical microscope. In the case of optical and electron microscopes, the imaging model is, by now, well-established, highly reliable, and commonly known as physical optics. In the case of profiling experiments, an appropriate imaging model framework may well be mass balance kinetics, but the details, as illustrated by Fig. 2 or 4 are still very much subject to debate.

Since the profiling experiment is sparse and noisy, we do not have a sufficient amount of sufficiently diverse experimental data to “look in all directions” of the kinetics parameter space. Each experiment only yields a (in general non-linear) projection of object points \( \theta \) in the M-dimensional kinetics parameter space onto the image points \( F(\theta,U) \) in d-dimensional image space. Sparsity and noise imply that typically only a lower dimensional image sub-space, of dimension \( d_{eff} \leq M \), can actually be resolved by the experiment. The MINE design procedure cannot eliminate the blurring of our vision; but it can help to minimize the blur.

**Criterion 1: MINE by maximal distance in image space.** To develop these notions into a quantitative MINE criterion, let us first consider the simplest case: the design of a Maximally Informative Next Experiment to measure only a single data point \( y \). Suppose we randomly draw two possible choices of models from the model ensemble \( Q \), denoted by \( \theta \) and \( \theta' \), which both give predictions consistent with the “old” experimental data (within the experimental uncertainties). To distinguish between these two choices, we want to perform the next experiment with control vector \( u \). The predicted outcomes for this next experiment would be, respectively, \( f(\theta,u) \) and \( f(\theta',u) \). The crucial point to notice here is this: the more these two predicted outcomes \( f(\theta,u) \) or \( f(\theta',u) \) differ from each other, the “better” the next experiment will allow us to discriminate between the two model choices. As a “metric” of the difference between the two members of the ensemble, we could choose, for example,

\[
V_{\theta,\theta'}(u) = \frac{|f(\theta,u) - f(\theta',u)|^2}{2}.
\]

The Maximally Informative Next Experiment \( u \) is then the one that maximizes this difference metric. Letting the joint distribution

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**Figure 4. Alternate genetic network for the biological clock from [33].** Molecular species (i.e., reactants or products) in the network are represented by boxes. The terms are the same as described in the legend of Fig. 2. The main difference is that the WCC has a light and dark form denoted WCC\(^L\) and WCC\(^D\). When these two forms bind upstream of \( frq \) and \( ccg \) genes, this leads to two different transcriptionally active forms of the gene, such as \( frq^{1D} \) and \( frq^{1L} \). In addition, photons (in yellow) can enter the system to interact with WC-1 in four ways, depending on the bound state of the WCC in the reactions E1, E2, E3, and E4. All four of these reactions have been given non-zero back reaction rates. The final difference is that the two forms of WCC lead to two deactivation reactions of WCC by FRQ, labeled P and Q. Reactions, such as A and P, have cooperative kinetics: (A) \( n \) WCC\(^D\)+\( frq^{0} \rightarrow frq^{1} \) and (P) WCC\(^D\)+m FRQ \rightarrow WC-2+m FRQ. The n and m are Hill coefficients or cooperativities. Only for 6 reactions, such as the “A” reaction, a back reaction, such as (7) frq\(^{1D} \rightarrow n \) WCC+\( frq^{0} \), included, with non-zero rate. The rate constants specify the right hand side of the kinetics model in equation (1) through the Law of Mass Action in Materials and Methods.

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of the randomly drawn pair \( Q(0,0') = Q(0|Q(0')) \), the foregoing criterion can be applied to an ensemble of models by choosing \( u \) such that it maximizes the average of \( V_{0,0'}(u) \):

\[
V(u) = \int_{0}^{\theta} V_{0,0'}(u) Q(\theta) Q(\theta') = E \left[ f(.,u) \right] - E \left[ f(.,u) \right] ^2. 
\]

(3)

Where \( | . \) denotes integration or summation over all \( \theta \)-components and \( E[ . ] \) denotes the mean over the ensemble probability distribution \( Q(\theta) \). Our MINE criterion \( V(u) \) is then the variance in our prediction within the ensemble and can be evaluated by Monte Carlo (MC) methods \([11]\). To achieve maximum “resolution” in model \((\theta)\) space, we thus “tune” the next experiment \( u \) so that it will take us into those regions of prediction \((\theta)\) space where there is the most uncertainty: the experiment \( u \) so that it will take us into those regions of \( n \)-dimensional image difference space. That is, \( n \) components of \( V(\theta, 0', U) : = F(0, U) - F(0', U) \).

(4)

That is, we replace \( V_{0,0'}(u) \) in \([2]\) by

\[
V_{0,0'}(U) = \left| \Delta F(0,0', U) \right|^2 / 2
\]

(5)

where \([ . ] \) denotes the Euclidean norm, i.e., \( | \Phi | = | \Phi |^{1/2} / | \Phi |^{1/2} \). Inserting this into \([3]\) we get the MINE design criterion

\[
V(U) = E \left[ | f(.,u) |^2 - | E[f(.,u)] |^2 \right] - \sum_{i=1..d} \left( E \left[ f(.,u_i) \right]^2 - | E[f(.,u_i)] |^2 \right)
\]

(6)

It is easy to see from \([6]\) that maximizing this \( V(U) \) will collapse \( u \) at the same \( u \)-point where the individual variance in the \( 1 \)-dimensional (single-variable) prediction space, \( E \left[ | f(.,u) |^2 - | E[f(.,u)] |^2 \right] \), is largest. So, this MINE criterion would demand that the next experiment simply observe the same \( y \)-variable \( d \) times, instead of observing \( d \) independent \( y \)-variables. Clearly, this criterion lacks the ability to enforce independence of multiple observables.

**Criterion 2: MINE by maximal volume in image space.** To construct a likely more useful MINE criterion, which does enforce some measure of independence of the observables, we are again guided by the microscopy analogue. Suppose we are “viewing” a certain “object space volume” \( V_o \) through our microscope. By way of the mapping function, this produces from \( V_o \) an “image difference volume” \( V_A \) in \( d \)-dimensional image difference space. That is, \( V_A \) is the volume swept out by the image difference vector \( \Delta F(0,0', U) \) for all pairs of object points \((0,0')\) in \( V_o \times V_o \), or, formally, \( V_A(V_o,U) : = \Delta F(V_o,V_o,U) \).

Our notation makes it explicit here that \( V_A \) depends on the choice of the control vector \( U \), as well as on \( V_o \).

To formulate an improved MINE criterion, we propose to invoke the volume \( V_A \) swept out by \( \Delta F \), instead of the Euclidean norm of \( \Delta F \) used in \([6]\). The basic microscopy-inspired idea here is this: the greater the volume amount contained in \( V_A(V_o,U) \), the more detail we should be able to discern in \( V_o \). In other words, we should be able to gain more information about the contents of \( V_o \) if we tune our microscope’s control vector \( U \) so as to increase the \( d \)-dimensional image difference volume amount, denoted by \( | V_A(V_o,U) | \).

However, unlike the Euclidean distance criterion \([6]\), the requirement of sweeping out a higher-dimensional volume \( V_A \) will naturally enforce a certain degree of independence of the observables. Notice here that the Euclidean norm measures just the length of the \( \Delta F \)-vector and this can be maximized even if \( \Delta F \) sweeps out only a \( 1 \)-dimensional sub-manifold. By contrast, \( V_A \) is by construction a higher-dimensional manifold with a dimensionality of up to \( d \) or \( M \), whichever is less.

The next question is then how to choose an appropriate \( V_o \) or the corresponding \( V_A \), in terms of the ensemble pair distribution \( Q(0,0') = Q(0|Q(0')) \). Again, this requires an ad hoc decision and we are guided in making it by computational expediency. In fact, the foregoing considerations of constructing a \( V_A \) from an underlying \( V_o \) in object \((\theta)\) space should only be regarded as a heuristic motivation for introducing such a \( V_A \). As a practical matter, our \( V_o \)-based MINE approach is greatly simplified if we do not try to construct an exact \( V_A \) from a given \( V_o \). Rather, we will define a “representative” \( V_A \) swept out by \( \Delta F(0,0', U) \) when \( \theta \) and \( \theta' \) are drawn from “typical” values prescribed by the ensemble pair distribution \( Q(0,0') = Q(0|Q(0')) \).

This \( V_A \) will be constructed from the characteristic variance/co-variance ellipsoid of \( \Delta F \) and it will be again dependent on the control vector \( U \).

To that end, it is conceptually (but not computationally) useful to first define the ensemble distribution of \( \Delta F \):

\[
Q_A(\Phi, U) : = \int_{0}^{\theta} \delta(\Phi - \Delta F(0,0', U)) Q(\theta) Q(\theta').
\]

(7)

where \( \Phi = [\Phi_1, \Phi_2, \Phi_3] \) is any point in \( \Delta F \)-space and \( \delta_{\theta \theta'} \) is the Dirac delta-function in \( d \) dimensions. \( Q_A(\Phi, U) \) is the probability density for \( \Delta F(0,0', U) \) to take on the value \( \Phi \), given that \( \theta \) and \( \theta' \) are independently distributed according to \( Q(\theta) \) and \( Q(\theta') \), respectively. \( Q_A(\Phi, U) \) defines an effective \( V_A(U) \) in the image difference \( \Delta F \)-space by way of the characteristic elliptoid of \( \Delta F \)'s \( d \times d \) variance/co-variance matrix \( D(U) \), given by

\[
D_{\Delta}(U) : = \int_{\Phi} \Phi D_{\Delta} \Phi_{\Delta}(\Phi, U)/2.
\]

(8)

Note that \( \Delta F \)'s characteristic variance/co-variance ellipsoid is centered at the origin, \( \Phi = 0 \), since \( Q_A(0, U) = Q(\Phi, U) \) due to \( \Delta F(0,0', U) \). The squared half-axis lengths of the characteristic ellipsoid are the \( D \)-matrix eigenvalues, with corresponding eigenvectors defining the respective half-axis orientations. The ellipsoid’s half-axes are orthogonal to each other and they define a rectangular prism in \( \Delta F \)-space whose volume amount, by a universal constant prefactor, is proportional to that of the ellipsoid. Instead of the ellipsoid itself, we therefore choose this “variance/co-variance prism” of \( \Delta F \) as our image difference volume \( V_A(U) \). The square of its volume amount, \( | V_A(U) |^2 \), is the determinant of \( D(U) \) and this is what we can use as a possible MINE criterion to be maximized:

\[
V(U) = \det(D(U)) | V_A(U) |^2.
\]

(9)

This is sometimes referred to as the generalized variance, and its distribution is known exactly if the predictions are Gaussian over the ensemble \([22, \text{Th. 3.2.1.5}]\).

We should strongly emphasize here that our invocation of the (co-)variance ellipsoid of \( \Delta F \) does not imply or require the \( D \)-distribution \( Q_A(\Phi, U) \) to be Gaussian. Via the matrix \( D(U) \) in \([8]\),
such an ellipsoid can be constructed for any $Q^h\Phi(U)$. Given the ad
hoc character of the entire MINE approach, the resulting $v_k$ is “as
good as any” for purposes of representing a “characteristic
volume” swept out by $\Delta F(0,0',U)$. The main advantage of this
construction is its computational feasibility: combining [4,7] and
[0], we can re-write $D_k(U)$ in terms of ensemble means over $Q^h$

$$D_{ik}(U) = E[f_i(.,u_i)f_j(.,u_k)] - E[f_i(.,u_i)]E[f_j(.,u_k)]$$  \hspace{1cm} (10)

which can be calculated by ensemble Monte Carlo evaluation
[7,11] of the required ensemble means $E[...]$. We also note here
that the D-matrix, as well as the E-matrix defined below, is scale-
free, that is, independent of the choice of model concentration
units, since the definition of $D_k(U)$ invokes only $\Delta F(0,0',U)$ [4]
which involves only the differences of log-concentrations or,
equivalently, logs of only concentration ratios. Hence, the $D_k(U)$
matrix elements, the corresponding $E_{ik}(U)$ matrix elements defined
below, and all MINE criteria developed here based on these two
matrices are scale-free.

**Hilbert Space picture of MINE formalism.** The foregoing
results can also be re-stated in terms of a Hilbert Space (HS)
formalism, using a HS of functions defined on the kinetics model
parameter ($\theta$)-space where the (co-)variance serves as the HS
inner product. That is, for any pair of such functions, $g(\theta)$ and $h(\theta)$,
we define the HS inner product by:

$$ (g,h) = E[g(.)h(.)] - E[g(.)]E[h(.)].$$  \hspace{1cm} (11)

The log variables $\log(y_i)$ are now represented by HS vectors $f_i$:

$$ f_i(\theta) = f(\theta,u_i) \hspace{1cm} \text{for } i = 1, \ldots, d;$$  \hspace{1cm} (12)

and the (co-)variance matrix element $D_k$ is simply the inner
product of HS vectors $f_i$ and $f_k$.

$$ D_{ik} = \langle f_i|f_k \rangle. \hspace{1cm} (13)$$

The ensemble standard deviation of the observable predicted by $f_i$
is the HS vector norm or “length”, denoted by $\|f_i\|$, where $\|\| \|_2$
is defined by $\|g\| = (E[g(.)^2])^{1/2}$. For notational convenience, we are
classically suppressing the dependence on $U$ or $u_i$ for quantities
like $f_i$ or $D_k$.

Independence of observables $\log(y_1), \ldots, \log(y_d)$ is very naturally
represented in this formality in terms of linear independence of
the corresponding HS vector set $f_1, \ldots, f_d$. These d HS vectors span
a finite-dimensional subspace in HS which has dimension d, if
$f_1, \ldots, f_d$ are linearly independent; else it is less than d. A linearly
independent HS vector set $f_1, \ldots, f_d$ also spans a d-dimensional
prism in HS, and the MINE criterion [9] has a simple interpretation
in terms of this HS prism: from [13], it is easy to show that $\det(D)$ is the square of the volume of this HS prism.

Hence, the characteristic (co-variance) prism $v_k$ in d-dimensional
$\Delta \theta$-space has an alternative (“dual”) representation in terms of
the HS subspace prism volume. In contrast $v_k$ in $\Delta \theta$-space, the HS
prism spanned by $f_1, \ldots, f_d$ is, in general, not rectangular since
$f_1, \ldots, f_d$ are not guaranteed to be mutually orthogonal with respect
to their HS inner product [11].

If the $f_1, \ldots, f_d$ become linearly dependent, their HS prism
collapses to a lower-dimensional one, and $\det(D)$ vanishes. In the
other extreme, if the observables are uncorrelated, the corre-
sponding $f_1, \ldots, f_d$ are mutually orthogonal in terms of the HS inner
product, i.e., they are maximally independent: their HS prism
volume is simply given by the product of their vector lengths $\|f_i\|$
and hence $\det(D) = (\|f_1\| \cdots \|f_d\|)^2$. In general, for correlated
observables, the $f_1, \ldots, f_d$ are non-orthogonal and we have
$\det(D) = (\|f_1\| \cdots \|f_d\|)^2$. The ratio $\det(D)/(\|f_1\| \cdots \|f_d\|)^2$
can be regarded as a composite measure of the degree of independence
of the observables and this ratio, as discussed further below,
depends on the HS “angles” between pairs of $f_i$-vectors, but not
on their individual lengths $\|f_i\|$. Maximizing $V(U) = \det(D(U))$
therefore requires a compromise between maximal mutual
independence of all observables and maximal variance of each
individual observable.

We note in passing that the Euclidean distance criterion [6] can
also be expressed in terms of the (co-)variance matrix $D$: the right-
hand side of [6] is the trace of $D(U)$. However, in contrast to the
volume criterion [9], for the Euclidean distance criterion [6], $V(U)$
is simply the sum of the squared HS vector lengths, i.e.,
$V(U) = \text{trace}(D(U)) = \|f_1\|^2 + \cdots + \|f_d\|^2$, and that is maximized
when each individual HS vector length is maximal, that is, when
each observable has maximal variance, regardless of any co-
variance correlations between observables, as already discussed
under [6].

**Criterion 3: MINE by maximal observational independence.** Based on these considerations, we propose one further MINE criterion which more strongly than [9]
emphasizes independence of the observables. Instead of the
original HS vectors $f_i$, we use normalized HS vectors

$$ g_i(\theta) : = f_i(\theta)/\|f_i\| \hspace{1cm} \text{for } i = 1, \ldots, d;$$  \hspace{1cm} (14)

to define a new “normalized” (co-)variance matrix or correlation
matrix, denoted by $E$, analogous to [13],

$$ E_{ik}(U) = \langle g_i|g_k \rangle = D_{ik}(U)/(\|f_i\| \|f_k\|)$$

$$ = E[g_i(.,u_i)g_k(.,u_k)] - E[g_i(.,u_i)]E[g_k(.,u_k)]$$  \hspace{1cm} (15)

This is the well known correlation matrix between the predictions
[22]. Our proposed third MINE criterion is then to maximize

$$ V(U) : = \det(E(U)) = \det(D(U))/(\|f_1\| \cdots \|f_d\|)^2.$$  \hspace{1cm} (16)

In contrast to [9], the variances of the observables do not affect
$\det(E(U))$, only their degree of linear independence does. In HS
geometrical language, $\det(E(U))$ is the squared volume of a prism
spanned by the HS unit vectors $g_1, \ldots, g_d$ and that volume is
determined entirely by the pairwise “angular” relations between
the $g_i$ not by their individual lengths which are all fixed at
$\|g_i\| = 1$.

Such a MINE criterion is likely advantageous in applications
where the HS vectors of the observables, $f_1, \ldots, f_d$, are “almost”
linearly dependent. The greatest gain in information from the next
experiment is then likely achieved by improving the independence
of the observables, rather than by maximizing their individual
variances. This scenario of “almost” linearly dependent (i.e., highly
correlated) observables is what we have in fact encountered,
consistently, in our MINE calculations for the three Computing
Life cycles reported in this work. Maximizing $\det(E(U))$ is therefore
the MINE criterion we have employed to guide the design of our
life cycle 1, cycle 2 and cycle 3 experiments, subject to additional
numerical modifications now to be discussed.

There are several additional reasons why the $\det(E(U))$ is the
preferred MINE criterion. The correlation matrix is a well known
measure of linear dependence between variables (i.e., the
predictions \( F \) as well as a well known measure of stochastic dependence when the predictions \( F \) are Gaussian over the ensemble. The \( \det(E(U)) \) MINE measure is bounded between 0 and 1. The value of 1 denotes linear independence, and in the case of Gaussian predictions, complete stochastic independence of the predictions. The value 0 means perfect linear dependence of the predictions, and in the case of Gaussian predictions, perfect stochastic dependence. The measure is familiar and easy to interpret, and finally, the \( \det(E(U)) \) has well known distributional properties, particularly when the predictions are Gaussian [22].

**Volume collapse pathology.** The lack of sufficient linear independence of the \( g_1,\ldots,g_d \) HS vector set (or, equivalently, of the \( f_1,\ldots,f_d \) set) is most easily diagnosed numerically by calculating the \( d \) eigenvalues of the D-matrix, denoted by \( \lambda_n = \lambda_n(U) \) and enumerated by \( n = 1,\ldots,d \) in descending order, with corresponding complete, orthonormal \( d \times 1 \) eigenvectors \( e^{(n)} \). Since \( D \) is non-negative, so are the exact \( \lambda_n \). However, in our actual MINE calculations, we encounter the numerical difficulty which the numerical inaccessible exact \( \det(D) \) is then replaced by the numerically stable determinant of a modified \( D \)-matrix, \( \det(E(\text{cut})(U)) \). This is what we have actually done in the MINE calculations reported here. We also note in closing that the Euclidean distance criterion [6], \( V(U) = \text{trace}(D(U)) \), is numerically not affected by the ellipsoid volume collapse: \( \text{trace}(D) \) is the sum, not the product, of the D-eigenvalues and it is therefore dominated by the numerically well-controlled largest eigenvalues only.

**Cycle 1–3 kinetics ensemble simulations and MINE calculations.** In all MINE calculations reported here, the observables \( \log(y_1),\ldots,\log(y_d) \) were chosen to be the log-concentrations of the 3 clock RNA species which are represented in our network model shown in Fig. 4: \( \log([f_0]),\log([w_0-f]),\log([w_0-f-1]) \) and \( \log([w_0-2]) \) where \( [w_0-1] \) is the combined total of the \( "w0" \) and \( "c1" \) versions of the \( [w_0-f] \) RNA, i.e., \( [w_0-1] = [w_0-1] + [w_0-2] \). Each of these 3 RNA concentrations was to be measured at 13 observation time points, \( t_j \) with \( j = 1,\ldots,13 \); hence there were \( d = 3 \times 13 = 39 \) data points \( y_{1j},\ldots,y_{dj} \) to be observed in the next experiment and the index \( "i" \) in the above sections therefore represents both the time index \( j \) and the species index \( n \), i.e., \( i=(j,n) \) with \( n = 1,2,3 \) for the 3 clock RNA species. The observation times \( t_j \) are chosen to have equidistant spacing \( t_b \), after an initial time lag of \( t_0 \), measured from the starting time of the experiment, \( t=0 \), where \( \tau = 0 \) is defined by the initial Light-to-Dark (L/D) transition. Hence,

\[
t_j = t_L + t_S (j - 1) \quad \text{for} \quad j = 1,\ldots,13.
\]

In each MINE cycle, an ensemble simulation was performed for the kinetics model for Fig. 4, using the same ensemble simulation procedure as described in [7], to generate a “representative” MC sample of 40,000 random \( \theta \)-vectors, drawn from the respective ensemble distribution \( Q(\theta) \) for that cycle. A subset of 200 random \( \theta \)-vectors from this sample was then used to calculate MC estimates for the ensemble expectation values \( E[\ldots] \) for evaluation of the E-matrix via [15].

In the cycle 1 ensemble simulation, \( Q(\theta) \) was constructed, as described in [7], from the same prior (“old”) experimental input data as shown in Figs. 2 and 4 of Ref. [7], taken from the literature, Refs. [3,23,24] and [25]. For the cycle 2 (cycle 3) ensemble simulation, \( Q(\theta) \) was revised by adding the new experimental data, from the new cycle 1 (cycle 2) experiment to the cycle 1 (cycle 2) prior experimental data set. In addition, the cycle 2 and cycle 3 ensemble simulations included conidiation density data from the 48 hr artificial day, interpreted as a measure of the CCG protein concentration in Fig. 1, as described in [7]. These conidiation data were generated in a race tube experiment with a periodic light/dark (L/D) exposure with a 48 h period and are the data shown in the right-most panel of Fig. 1.

In all new experiments reported here, where light exposure was applied, the light intensity at the sample location was about 70 \( \mu \text{mol}(\text{photons})/(\text{s}\cdot \text{m}^2) \) in Einsteinian units, or about 15 \( \text{W} / \text{m}^2 \) in radiometric units, or about 5,300lux–490 ft-candles in photometric units, assuming a “cool white” spectral distribution, generated by our fluorescent light source. (The approximate conversion factors are 1 \( \text{W} / \text{m}^2 \leftrightarrow 4.622 \, \mu \text{mol}(\text{photons})/\text{(s} \cdot \text{m}^2) \leftrightarrow 350.7\text{lux} = 32.58 \text{ft-candles} \) for the “cool white” spectrum, as given in [26].) By contrast, the light intensity was only 20 \( \mu \text{mol}(\text{photons})/(\text{s}\cdot \text{m}^2) \), with unspecified spectral distribution, for the light exposure experimental data we have taken from the literature [23]. Since our kinetics ensemble simulations for cycle 2 and 3 incorporate both our new experimental data and the literature data [25] into the respective distributions \( Q(\theta) \), we have assumed that both, the literature experiments and ours, were performed with the same photon spectral distribution. We have
therefore modeled all light exposure experiments in terms of photon “concentrations” [entering into the reaction rate function $G$ in [1]] which are chosen proportional to the respective light intensities. For all light-exposed experimental data used [25] or reported here, the light exposure was periodic, starting at the initial $(t = 0)$ L/D transition, with a 50% duty cycle (i.e., the same duration of D and L) and a phasing of either D/L (i.e., dark first, then light) or L/D, as indicated in the discussion of the respective results. This time-dependent light exposure was modeled, as in [7], by a corresponding time-dependent periodic photon concentration of rectangular pulse shape, entering into the rate function $G$.

In cycle 1, $V(U) = \det(E(U))$ was maximized with respect to $t_S$ and $t_L$ for a “next” experiment designed to measure the three clock RNA species in the dark. The resulting optimal MINE values of $t_S = 0$ and $t_L = 5$ h (see Fig. 5) were slightly modified, to $t_S = 0$ and $t_L = 4$ h, so as to keep the total duration of the cycle 1 experiment below the ~50 h limit imposed by experimental clock stability constraints in liquid cultures. The latter values of $t_S$ and $t_L$ were then used, without further adjustments, in all subsequent cycle 1, cycle 2, and cycle 3 RNA profiling experiments and in the corresponding cycle 2 and cycle 3 MINE calculations.

In cycle 2, $V(U) = \det(E(U))$ was maximized with respect to the period $t_P$ of the alternating light exposure, for a “next” experiment designed to measure the three clock RNA species subject to a 50% duty cycle and D/L phasing. From the resulting optimal range of $t_P \sim 20$ h–24 h (see Results), $t_P = 24$ h was chosen for the actual cycle 2 experiment.

In cycle 3, $V(U) = \det(E(U))$ was maximized with respect to the “gene knock-down” transcription ratio (TR), and with respect to the choice of the gene species to be knocked down, for a “next” experiment designed to measure the three clock RNA species in the dark, but with one of the three clock genes subjected to an experimentally controlled reduction in its transcription rate coefficient. The resulting “most informative” gene was found to be $we-1$, with a transcription rate coefficient reduced to TR = 10% of the wild-type value (see Results). In the actual cycle 3 experiment, this MINE-recommended knock-down value for $we-1$ was approximated, within the limitations of experimental control, by a TR of about 30% of the wild-type value. The clock RNA data from this $we-1$ knock-down experiment were then incorporated with all other prior literature, cycle 1 and cycle 2) experimental data into the Q(0) for one “terminal” ensemble simulation for the network in Fig. 2 and 4. The clock RNA results from this terminal ensemble simulation, along with the respective new RNA data generated in the 3 MINE cycles, are shown in Results for the network in Fig. 4.

**Perturbing the genetic network**

Once the MINE perturbation experiment is designed, the perturbation is implemented at the next stage in the cycle in Fig. 3., as now described.

**Strains.** All but one strain used (namely OR-74A below) carry a $bd$ ($bd$) mutation, permitting the observation of the clock in race tubes. The $bd$ mutation (Fungal Genetics Stock Center 1858) was used for the first series of microarray experiments in the dark as well as the second series of microarray experiments examining a light-response. Strain 87-84-6-8 [7] carrying a mutation in $we-1$ and an inducible copy $qa:we-1$ at the his-3 locus was used in the dial down of $we-1$ expression [7]. Dial-down was achieved by shifting liquid cultures from 0.3% quinic acid to 2% galactose. Strain 74A-OR23-1A was used in shift experiments from sucrose (1.5%) to quinic acid (0.3%) as a control to identify QA inducible genes [27]. Strain 93-4 ($frq^{+} qa:frq$) transformed with pDE3dBiH$qa:frq^+$ was kindly provided by Deborah Bell-Pedersen (Biology Department, Texas A & M University) to test for auto-feedback loops in $we-1$ and $we-2$.

**Liquid Growth Conditions for harvesting RNAs.** Establishment of liquid cultures followed Nakashima [28] and [5], and the cultures were grown for 48 hrs in petri plates. Half-cm mycelial disks were cut from the mat and dropped into 300 ml flasks with 100 ml Fries+2.0% glucose+5.0% arginine+supplements [29].

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Figure 5. MINE calculation to determine when to start sampling ($t_L$) and how often ($t_S$). The MINE surface is plotted as function of the lag $t_L$ in hrs and spacing $t_S$ in hrs; higher values on the MINE surface suggest the preferred design points ($t_L$, $t_S$). Color contours of the log of the MINE criterion $\det(E)$ are overlayed as a function of the lag ($t_L$) and spacing ($t_S$) to show points on the surface of similar MINE values. The MINE surface suggests to start sampling immediately (small $t_L$) and to make the spacing ($t_S$) between observations as large as possible. The maximum permissible spacing ($t_S$) between observations is 5 hrs, as determined by two constraints. One, there is the cost constraint of 13 microarray chips per cycle, and two, beyond a 50 hr experiment in cycle 1 stable oscillations in liquid culture are not guaranteed.

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Cycle 1. In the first series of microarray experiments to identify circadian genes (cycle 1) all flasks were placed in a shaker (New Brunswick Scientific, Edison, NJ. Series 25) at 150 rpm and at 25 °C for the same period of time, 50 hrs, and also were given at least 2 hrs of 70 micromoles per second per meter squared (µM/s/m²) before L/D (light to dark) transition. A total of 13 flasks were harvested by vacuum filtration, one flask every 4 hrs starting at time 0, the L/D transition time, in such a way that the total growth time of each liquid culture was kept constant [42]. After the L/D transition initiating the experiment, the flasks were shaken at 150 rpm and left in the dark. Cells were placed at −70 °C to await RNA harvesting using a High Pure RNA kit (Roche, Inc.). The harvested RNA was subjected to RT-PCR and then microarray analysis as described below.

Cycle 2. In the second series of microarray experiments to study the light-response (cycle 2) the total growth time was not controlled, and 24 hrs after the L/D transition the light (70 µM/s/m²) was turned back on for 24 hrs. 17 flasks were harvested at time points 0 hr, 4 hrs, 8 hrs, 12 hrs, 16 hrs, 20 hrs, 24 hrs, 24 hrs+20 m, 24 hrs+40 m, 25 hrs, 26 hrs, 28 hrs, 32 hrs, 36 hrs, 40 hrs, 44 hrs, and 48 hrs.

Cycle 3. In the third series of microarray experiments to study the WCC-response (cycle 3), prior to the L/D transition, mycelial disks were transferred into 500 ml flasks with 100 ml liquid medium and placed in the dark. Flasks were harvested at 0, 10 m, 20 m, 30 m, 40 m, 50 m, 1 hr, 2 hrs, 4 hrs, and 8 hrs. Four additional replicate 0 time points were harvested as well.

Observing the outcome of the perturbation

Observing the outcome of the perturbation completes the cycle through Fig. 3. Measurements are made on the system by a combination of race tube assays, real-time polymerase chain reaction (RT-PCR), and profiling with oligonucleotide arrays, as now described.

Race tube assay for biological clock. Starter cultures were made on 0.1% glucose+0.17% arginine+Vogel's medium [29] and subject to a 25 µM/s/m² light source. Conidia were filtered with glass wool as described in [30] and used to inoculate race tubes layered with 20 ml of 0.1% Glucose+0.17% arginine+Vogel's medium or 0.001 M quinic acid+0.17% arginine+Vogel's medium [29]. A total of 135 replicates tubes were inoculated and either subject to low (25 µM/s/m²) or high intensity pulses (70 µM/s/m²) once per hour for 90 sec over a twelve hour period or subjected to no light pulses to measure period and a phase response as described in [30] and [31].

RNA isolation. RNAs were isolated using the High Pure RNA isolation kit (Roche, Inc.). Their quality and quantify was assessed using an RNA Nano LabChip (Agilent Technologies, Inc.) was subjected to one RNA isolation kit (Roche, Inc.). Generally only samples with a ratio of at least 1.00 for 28S/18S RNA were detected by RT-PCR (ABI-Prism 7500, Applied Biosystems, Inc.) according to manufacturer directions using Taq Man probes against an rRNA standard. Triplicate reactions (50 µl) were analyzed using the ΔAC method (Applied Biosystems, Inc.).

Design of 12K Oligonucleotide Arrays (Combimatrix, Inc.). These arrays were constructed with Version 3 of the Neurospora crassa genome sequence [32] from a file labeled neurospora_3_gene_dna_3203.txt downloaded from the Broad Institute Web site. From this sequence Combimatrix, Inc. designed 12,000 oligonucleotides of ~35 nt to synthesize electrochemically on their chips. Several genes were represented multiple times with 53 oligonucleotides derived from the following genes: wc-1 (5 duplicates); uex-2 (5); fqr (5); rDNA (>7), qa-x (5); qa-2 (5); qa-4 (5); qa-3 (5); qa-y (5); qa-z (5); qa-1F (5); qa-1S (5). In addition, 633 negative control oligonucleotides were added from bacterial, plant, and λ-phage sequences as well as features with no oligonucleotides and quality control oligonucleotides. The rDNA oligonucleotide sequences were treated as negative controls in the microarray experiments as well because none of the at least eight rDNA-derived oligonucleotides on each chip showed up in an expanded Fig. 6 with 4721 circadian genes [33] with or without a known LRE upstream. In addition 22 duplicate oligonucleotides from each of 4 distinct λ-phage sequences were added to be used as positive controls for all but 4 arrays (by spiking them into the amplified RNA (aRNA) probe). These positive and negative controls were scattered at random on each array. The arrays with their design are at http://www.yale.edu/townsend/Links/ffdatabase/introduction.htm [34]. Their accession numbers are 13, 34, and 36 for cycle 1, cycle 2, and cycle 3, respectively. The same design was used on all Combimatrix chips except for samples 48 hrs, 44 hrs, 40 hrs, and 36 hrs on cycle 1.

RNA amplification and oligonucleotide array hybridization. 750 ng of total RNA (as determined by a Nano LabChip (Agilent Technologies, Inc.)) was subjected to one round amplification using the MessageAmp aRNA Amplification Kit (Ambion Inc), which uses an “Eberwine type” amplification. Biotin-11-UTP and CTP (Enzo Life Sciences, Inc.) were incorporated during the in vitro transcription reaction. A total of 5 µg of amplified RNA (aRNA) was fragmented, A total of 10 pM each bioinylated spike-in oligonucleotide (phage) was added with hybridization solution, and hybridized according to manufacturer’s protocol rev 2.03 (http://www.combimatrix.com). Hybridization was performed at 45 °C for 24 hrs using a 25% formamide based solution. Washing was done according to manufacturers protocol rev 2.03. Streptavidin Alexa Fluoro® 647 conjugate (Invitrogen) was used at a final concentration of 1.0 µg/ml to visualize hybridized targets. Laser confocal scanning was performed on a GSI Lumonics ScanArray 5000 (now manufactured by Perkin-Elmer, Inc.) using a single laser power and a photomultiplier (PMT) gain setting adjusted less than 10% between arrays. Versions of image software MI_Version_5_4_3, MI_Version_5_5_0, MI_Version_5_6_0, and MI_Version_5_7_0 (Combimatrix, Inc.) were used to obtain spot intensities (such as median foreground count) on each array feature for microarray analysis.

Quality Control on RNAs. RNA samples was confirmed to have a ratio of at least 1.00 for 28S/18S RNA on the LabChips (Agilent Technologies, Inc.). RNAs used for aRNA synthesis and hybridization to chips were visually scanned for trends in the foreground median count in control sequences in the (x,y) coordinates. For the λ-oligonucleotides spiked into each aRNA, the coefficient of variation (CV) in median foreground count was computed, and if the chip had a CV greater than 0.65 (n = 88), the sample was usually not used and redone. All chips were verified to have 51% of its features above median background (with the exception of samples 48 hrs, 44 hrs, 40 hrs, and 36 hrs on cycle 1, which were at 44–45%). This percentage (51%) of identified genes with expression above...
background is higher than that reported (38%) in [35] and close to the 60% reported in [36]. RT-PCR was done in parallel on all RNA samples, and if the relative quantification by RT-PCR with DDCt method did not agree quantitatively with the median foreground counts obtained on wc-1, wc-2, wc-2, wc-2, ccg-4, wc-1, wc-2, al-1, CAD11415.1, con-6, bli-3 rel, ccg-1, ckb-1, vvd, bli-3 rel, ccg-1 rel, ccg-15, ccg-2, ccg-6, ccg-7, al-3, FWD-1, ccg-13, ccg-14, lyz, camk, ccg-15 rel, wc-1, ccg-4 rel, al-2, bli-3 rel 2, frq, frq.

Figure 6. Transcriptional profile of approximately 2436 putative genes with LREs upstream at 0, 4, 8, ..., 48 hrs (values staggered on x-axis) after shift from light to dark (L/D) after background subtraction, normalization within arrays relative to the grand median of each chip, logging, and clustering with average linkage using Euclidean distance between mRNA profiles of different genes [38]. The bright green is —3, and the bright red is +3 is expression level on a decadic log scale. Data arose from 13 chips probed with a biotin labeled aRNA. Over 43 known clock-associated genes are overlayed in the right margin of this microarray experiment including varied known ccg genes. Genes in Fig. 2 are represented at least 5 times on each chip (explaining why frq appears 5 times in the margin). The 2436 putative genes with LREs upstream were selected by fitting $A_0 + A \sin (\omega t + \phi)$ by nonlinear least squares [44] to the profile of each of the 11,000 genes, and those with a significant regression sum of squares contribution with respect to the amplitude $A$ in $F_{1,9} > 5.12 (\alpha = 0.05)$ [94] and with a period between 16 hrs and 30 hrs are displayed (see Fig. 1 in [5] for frq mRNA peak separation of 16 hrs and 30 hrs). This work [5] establishes a standard for what is considered acceptable variation in the estimated period of the oscillator. See text for a reexamination of this standard. The smallest significant $F_{1,9} (= 5.12)$ observed among circadian genes with upstream LREs had an estimated amplitude of 497 and 3.98 fold variation in mRNA levels over time in contrast to the amplitude of frq mRNA, 974+/−79. The smallest significant amplitude (63 with an $F_{1,9} = 8.29$) estimated among circadian genes with LREs upstream had 1.44 fold variation in mRNA levels over time. A gray bar at the bottom indicates lights off. doi:10.1371/journal.pone.0003105.g006

Microarray Analysis. The median foreground (FG) counts were used on all 12K features. From each median foreground count on an oligonucleotide array a background subtraction was performed using the 5th percentile of the following negative control oligonucleotide features: (1) plant; (2) bacterial; (3) phage not spiked into aRNA; (4) quality control oligonucleotides (QC); and (5) empty. Then the median foreground counts were normalized within arrays by multiplying each feature’s median FG count on a particular chip $x$ (median FG across all chips in the cycle) / (median FG count on the particular chip). A MIPS functional classification was assigned each feature on a chip [37]. Hierarchical clustering of genes was implemented using the methods proposed in [38] and implemented in Cluster 3.0 [39].
available at http://bonsai.ims.u-tokyo.ac.jp/~mdehoon/software/cluster. Options selected for analysis were log transformation, mean-centering and normalization followed by average linkage (i.e., UPGMA) using Euclidean distance. Trees were displayed with Java TreeView 1.0.12 [40] available at http://treeview.sourceforge.net.

**Searching for WCC and QA-1F binding sites in silico.** Putative WCC GATX-binding sites or Light-Response Elements (LREs) were identified with the program pattern (Accelrys, Inc.) operating on the 1000 nt upstream of each identified gene [32] from file neurospora_crasa_#10BD2C.fasta from the Broad Institute Web site. The offset used was 1 and overhang 0. A mismatch of 2 was allowed. Patterns searched for were:

- \text{GATG\{5,20\}}
- \text{GATC\{5,20\}}
- \text{GATA\{5,20\}}
- \text{GATT\{5,20\}}

Putative QA-1F-binding sites were identified with the program pattern (Accelrys, Inc.) operating on the 1000 nt upstream of each identified gene [32] from file neurospora_crasa_#10BD2C.fasta from the Broad Institute Web site. The offset used was 1 and overhang 0. Patterns [41] searched for were:

- \text{GAT\{5,20\}GATT\{5,20\}}
- \text{GATT\{5,20\}GATT\{5,20\}}
- \text{GATT\{5,20\}GATT\{5,20\}}
- \text{GATT\{5,20\}GATT\{5,20\}}
- \text{GATT\{5,20\}GATT\{5,20\}}
- \text{GATT\{5,20\}GATT\{5,20\}}

**Results**

**Genetic networks for the biological clock guide the MINEing for clock-controlled genes**

The genetic network models in Figs. 2 and 4 [7,43] make three predictions about each gene in the genome under clock-control: a clock-controlled gene should: (1) maintain an endogenous circadian rhythm when the organism is grown in the dark; (2) be light-responsive when the organism is moved from Dark to Light (D/L); (3) change its expression when the level of the transcription factor WCC is dialed down (see Materials and Methods). The Computing Life paradigm is used below to discover these clock-controlled genes.

**Maximally Informative Next Experiment (MINE)**

The objective of the MINE approach is to develop a quantitative criterion (or criteria) for the amount of additional information that can be gained about the genetic network from the “next” experiment to be performed; and then to maximize this “measure of additional information”, denoted by \(V(U)\), with respect to the choice of the design or “control” parameters of the next experiment, denoted by the control vector \(U\). Control vector \(U\) comprises all those parameters which are known to, and are to some extent controllable by, the experimenter and which completely characterize measurements to be performed and the external conditions and perturbations applied to the biological system during the experiment. Two critical inputs for the MINE calculation are the underlying network kinetic rate equation model of the genetic network and any available “prior” or “old” experimental data. In a recently developed ensemble simulation approach [7,11], these two inputs are combined both to constrain the unknown kinetics model parameters and to predict the likely information content \(V(U)\) for the next experiment, given \(U\). Technical details and underlying conceptual ideas of the MINE approach are described in the Materials and Methods. Here we have used one of the MINE criteria in the Materials and Methods, \(V(U) = \text{det}(E(U))\), Eq. (16), to guide the design of new experiments on the biology of the clock. This criterion is the determinant of the ensemble correlation matrix \(E(U)\) between predictions. The predictions here are of the log concentrations of \(\text{wc-1}, \text{wc-2},\) and \(\text{fg}\) mRNAs over time in the next experiment.

When the predictions of two models in the fitted ensemble (the collection of all models consistent with available data) are highly correlated, the models will be difficult to distinguish by the next experiment \(U\); when the predictions of two models in the ensemble are less correlated, they will be more easily distinguished in the next experiment \(U\). A higher value of the MINE criterion \(V(U)\) recommends the experiment \(U\) for which predictions between any two randomly selected models in the fitted ensemble are more uncorrelated and hence more distinguishable. Each MINE calculation is done within the constraint of a fixed budget (i.e., 13 microarray chips per experimental cycle or equivalently, 13 time points to be sampled). The budget and hence the number of time points determine the dimension of the correlation matrix \(E(U)\).

Two possible hypotheses have been developed for the clock mechanism in Figs. 2 and 4, from [7] and [33]. An older and slightly more realistic genetic network [33,43] in Fig. 4 was used to guide the MINEing because the simpler genetic network in Fig. 2 was developed while the MINE experiments were underway. The older network [33] allows a light and dark form of WCC [4]. At the conclusion of the Computing Life enactment, these two different networks are tested against each other.

**Cycle 1 - Which genes are circadian?**

The first series of microarray experiments were designed to determine how many genes are under clock-control. If such genes were outputs of the clock mechanism in Fig. 4, then they should be able to maintain an endogenous rhythm of \(\sim 22\) hrs in the dark. The first experiment involves growing the organism in the dark for 48 hrs to observe the endogenous rhythm. The initial MINE design question concerns how often should we sample and when should we start sampling. The spacing between observations is denoted by \(t_s\), and the delay till the first observation by \(t_L\). The maximum in spacing (\(t_S\)) is limited by the time over which circadian rhythms are maintained in liquid culture and the cost constraint of 13 microarray chips (see Materials and Methods). A MINE calculation using published data [3,23–25] results in Fig. 5, based on the genetic network in Fig. 4.

The “best” experiment, with maximum \(\text{det}(E(U))\) – in the upper back corner of Fig. 5 – is to start sampling immediately and to use the maximum spacing of 4 hrs between observations. This was the microarray experiment, performed (see Materials and Methods) with the results shown in Fig. 6. These experimental results would suggest that as many as 43% of the genes could be clock-controlled. A more detailed statistical analysis below reduces this percentage to 25%. There are 2436 (22%) circadian genes, which is still considerably higher than 2–10% of circadian genes reported for \textit{Drosophila} [44,45] and \textit{Arabidopsis} [46–48], and 10% higher than that reported in [8] for \textit{Neurospora}. Our percentage, however, is not out of line with estimates of 36% based on in vivo enhancer traps in \textit{Arabidopsis} [49].

In addition to the 11,000 \textit{N. crassa} genes on each chip (including 43 genes used as positive controls), the chips carried 633 negative control oligonucleotide sequences including those derived from plant, bacterial, phage, and \textit{N. crassa} sDNA sequences. The
empirical false positive and false negative rates are reported in Table 1 for each microarray experiment. For the first microarray experiment (cycle 1), the empirical false positive rate is 18% with the nominal significance level of the periodicity test used. Of these 4721 circadian genes, 2436 of them have a LRE upstream [4]. With a multiple test correction suggested for microarray analysis by Storey [50] and implemented as in Benjamini and Hochberg [51], the number of circadian genes with upstream LREs drops to 1460. In this multiple test correction the ranked list of genes sorted by their P-values is simply trimmed from the high end near the nominal significance level (of P-values) to control the False Discovery Rate (FDR) as described in [52]. The target FDR is set to the nominal significance levels in Table 1. If we subtract the 18% of false positives, then 43–18 = 25% of the genome would appear to be under clock-control. The estimated percentage of circadian genes (25% corrected for false positives) is close to the uncorrected percentage of circadian genes with LREs upstream (22%). Requiring the presence of an upstream LRE appears to be a good filter for circadian genes.

Circadian oscillations are seen in Fig. 6 along rows by the alternating pattern of red (high expression) and green (low expression) for different genes, albeit with different phases. There also appear to be two clusters of known clock-associated genes with similar transcriptional profiles at the top and half way up Fig. 6 with different phases. The distribution of periods of oscillation is found in Fig. 7A with a mean of 24.92+/-0.09 hrs, implying the oscillations are circadian as predicted. This compares well with the average period between conidiation bands of 21.64+/-0.05 hrs obtained from 135 race tube assays (see Materials and Methods).

The phase of the clock-controlled genes is also interesting in Fig. 7B. Dusk (L/D transition) is taken as the zero time. The phase φ of 100% corresponds to being 360 degrees out of phase with genes at midnight. There are morning and evening genes in Fig. 7B. The frq mRNA has a phase of 48%+/-16% as a typical morning gene, and the wc-1 mRNA has a phase of 69%+/-12% as a typical dawn gene, being a blue-light receptor [4]. RNA metabolism genes tend to be dawn genes as well with a mean of 62%+/-5% (Fig. 7B), while regulators tend to be dusk genes with a mean of 28%+/-9%. Cell cycle genes among clock-controlled genes (in red in Fig. 7B) tend to be morning genes with a mean of 47%+/-8%, as is the cell cycle checkpoint kinase prd-4 [53]. This is consistent with light triggering conidiation in Fig. 1. The phase of genes may provide some clues as to how the clock allows the organism to adapt to its environment (see Discussion).

A naı¨ve expectation might be that while the phase would vary between different cognate genes as, in Fig. 7B, the gene periods in Fig. 7A would be expected to be the same. Several possible causes for this variation in period of clock-controlled genes in Fig. 7A present themselves. There is noise in mRNA profiling measurements on which the period estimates are based; there is also intrinsic noise in mRNA levels from cell to cell [54]; and we are observing the system over a short time interval covering only 2 periods of oscillation. As illustrated in [7], it can take longer than 2 periods before the limit cycle is established and, during the transient prior to that, neither period nor phase of oscillation are well-defined. The finite observation time also limits the accuracy of the measured period by way of an uncertainty principle [55]: the shorter the observation time, the greater the uncertainty in frequency and period.

Cycle 1 microarray results were validated by RT-PCR (see next section) on twelve genes including wc-1, wc-2, and frq relative to an rRNA standard with excellent quantitative agreement (Fig. 8). Oscillations in wc-1 mRNA with a period of 22.17 hrs +/- 1.66 hrs, which have not been reported before (see validation in the next section). The presence of a LRE upstream of the wc-2 gene would suggest additional feedback loops to wc-1 and wc-2 in Figs. 2 and 4 to make them autoregulatory. Evidence for an autoregulatory loop for wc-1 has recently been provided [56]. Oscillations in wc-1 mRNA are weak if plotted on the same scale as frq mRNA levels, as expected [24,7]. The periods for frq, wc-1, and wc-2 mRNA oscillations of 21.40hrs +/- 1.69 hrs, 23.5hrs +/- 2.47 hrs, and 22.17hrs +/- 1.66 hrs, respectively, in Fig. 8 agree well with the period of banding in race tubes above, namely 21.64hrs +/- 0.05 hrs [31,33].

Table 1. Observed fraction of false positives and false negatives among 633 negative controls on each microarray chip (see Materials and Methods) and among 43 distinct genes as positive controls using reported clock-associated genes [93].

| Microarray Experiment | Nominal significance level (α) | Observed fraction of false positives | Observed fraction of false negatives | estimated power |
|-----------------------|--------------------------------|-------------------------------------|-------------------------------------|-----------------|
| Circadian cycle (cycle 1) (in the dark) | 0.05 | 0.18 | 0.37 | 0.63 |
| Light-response (cycle 2) (D/L) | 0.20* | 0.17 | 0.47 | 0.53 |
| WCC response (cycle 3) (turn WCC off) | 0.20* | 0.22 | 0.44 | 0.56 |

The estimated power is 1 – fraction of false negatives observed. The fraction of false positives observed can be compared with the nominal significance level used to identify genes that are: (1) circadian; (2) light-responsive; (3) WCC-responsive.

The nominal significance level was adjusted using the positive and negative controls to insure that the estimated power was high.

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Regulators. An entire replicate of the cycle 1 experiment was conducted (results not shown and same replicate used for validating wc-2 microarray results), and the levels of kal-1 and rpn-4 mRNAs were measured every 4 hrs over a 48 hr window in the dark by RT-PCR (see Materials and Methods) in two replicates of cycle 1 (including the original cycle 1 experiment in the dark). A combined estimate of the amplitudes based on 26 time points was tested with an F-test (as described in the legend of Fig. 6) and found to be significant (F_{1,24} = 4.95, P = 0.04 with a period of 23 hrs for kal-1 and F_{1,24} = 6.65, P = 0.02 with a period of 30 hrs for rpn-4).

Signaling. The gene rrg-1 has a role in the osmotic response signaling pathway [57–58]. An entire replicate of the cycle 1 experiment was conducted (results not shown and same replicate used for validating wc-2 microarray results), and the levels of rrg-1 mRNA were measured every 4 hrs over a 48 h window by RT-PCR (see Materials and Methods) in two replicates of cycle 1 (including the original cycle 1 experiment in the dark). A combined estimate of the amplitudes based on 26 time points was tested with an F-test (as described in the legend of Fig. 6) and found to be significant (F_{1,24} = 6.82, P = 0.02) with a period of 17 hrs.

Cycle 2 - Which genes are light-responsive?
Each of these 2436 circadian genes from cycle 1 could be under the control of: (1) WCC; (2) a different oscillator [8]; or (3) multiple oscillators [59]; or be false positives. The chance of the latter is only 18% (see Table 1). As shown in Fig. 1, an important element to the clock is light-entrainment. As the organism is grown under different “artificial days”, that is, different periods of alternating light exposure, the organism speeds up or slows down its biological clock.

If these genes were under WCC-control, then they should also be light-responsive according to the genetic network hypotheses in Figs. 2 and 4. This poses the question of what artificial day period should be used for the experiment. A MINE calculation results in Fig. 9 using published results [3,23–25] plus the data in Fig. 6 (cycle 1), i.e., the MINE calculations are cumulative with respect to data already obtained.

The MINE results, shown in Fig. 9, suggest a long artificial day with a half-period of daylight of between 19 and 24 hrs. A second cycle of microarray experiments was therefore performed in which the light was turned back on after 24 hrs in a 48 hour observation period (see Materials and Methods). Results are shown in Fig. 10. Among these 3374 light-responsive genes (or 31% of N. crassa
genes), 1725 (or 16% of the genes) responded to light in Fig. 10 and possessed LREs upstream. With the Benjamini and Hochberg [51] multiple-test correction, 1026 out of these 1725 genes with upstream LREs remain significant. In a similar experiment Lewis et al. [60] report detecting 22 light-responsive genes induced out of 1343 distinct genes arrayed as cDNAs, or 3%, and Ma et al. [61] report 34% of the unique genes in Arabidopsis thaliana induced or repressed. Among the 31% of light-responsive genes detected here, up to 17% could be false positives, leaving 14% = 31%−17% as light-inducible. Among the 31% of light-responsive genes, 56% were induced (as opposed to repressed). The percentage of 0.56×14% = 8% is still higher than the 3% of Lewis et al. [60] and less than the 31% of A. thaliana. The estimated percentage of light-responsive genes (14% corrected for false positives) is close to the uncorrected percentage of light-responsive genes with LREs upstream (16%). Requiring an upstream LRE also appears to be a good filter for light-responsive genes.

A total of 768 genes were both circadian and light-responsive (Fig. 11), and the chance that any one of these 768 genes is a false positive would be (0.18)×(0.17) = 0.03 (Table 1), since the experiments were done independently. These 768 genes then remain candidates for cgs in Fig. 2 or 4. The response by genes to light falls into two clusters, one cluster being turned off (in the top part of Fig. 10) and one cluster being turned on (in the bottom part of Fig. 10). The positive response of some genes to light appears largely transient with a burst of expression after the Dark-to-Light (D/L) transition while other genes appear to have a sustained response after the D/L transition. Most of the known clock-associated genes fall within the bottom cluster of light-responsive genes with LREs upstream, as expected.

As a control, these results from cycle 2 were compared with a near replicate of this experiment reported in [60], using a different microarray technology and only a sample of the genes in N. crassa. With a power of 53% (Table 1) in our experiments, we would expect to see around 53% concordance with the experiments reported in [60]. In fact, we saw 64%+/−20% of the genes reported as light-responding in [60] in our cycle 2 experiments with good agreement to the 53% expectation.

**Cycle 3 - Which genes are under WCC-control?**

Another prediction of the genetic networks in Fig. 2 or 4 is that if WCC were dialed down (i.e., the mRNA level of wc-1 is reduced by use of a QA-inducible promoter as described in Materials and Methods), then a gene under its direct or indirect control should experience a sudden change in its mRNA level. To test this with a gene knock-down experiment, it is necessary to ascertain first what gene should be perturbed to yield maximum information about the genetic network in Fig. 4. A MINE calculation was done using published results [3,23–25] plus the data in Fig. 6 (cycle 1) and the data in Fig. 10 (cycle 2) as described in Materials and Methods.

The MINE calculation in Fig. 12, suggests that the most informative knock-down is to reduce wc-1 to 10% of its original...
transcriptional activity. As detailed in Methods and Materials, the knock-down was engineered with a mutation in the native \textit{wc-1} and with a quinic acid inducible copy of \textit{wc-1} introduced at another locus, producing a knock-down to 30\% of its original activity; the results are shown in Fig. 13. A total of 4655 WCC-responsive genes were found to respond, but only 2323 of these genes had a LRE upstream as reported in Fig. 13. With the Benjini and Hochberg [51] multiple-test correction, 1445 out of these 2323 genes with upstream LREs remain significant. The estimated percentage of WCC-responsive genes (20\% corrected for false positives) is close to the uncorrected percentage of WCC-responsive genes with LRE(s) upstream (21\%). Requiring an upstream LRE remains a good filter for WCC-responsive genes.

Most of the \textit{frq} gene mRNAs were dialed down, as expected. The \textit{frq} gene belongs to a cluster of genes being turned off in the top part of Fig. 13. Both \textit{wc-1} and \textit{wc-2} responded differently than \textit{frq} and belong to the second larger cluster of genes being turned on (at least transiently) in the bottom part of Fig. 13. They have a fast transient response about 20–40 m after the QA/GAL (L/D) transition and then a drop off (see Fig. 8).

Also of interest are the 440 genes that are both circadian and light-responsive, but not under WCC control in Fig. 11. These 440 light-responsive and circadian genes which are not apparently under WCC-control, could be responding indirectly to genes under the control of WCC. They could also be false-negatives under the WCC-responsive assay (Table 1); they could be responding through a yet to be identified oscillator [8]; or they could be responding through multiple oscillators [59]. The chance that any one of them is a false positive is 0.03 = (0.18) \times (0.17). The genes \textit{cpe-1}, NCU05429, and \textit{cgq-16} (WO6H02), have been previously identified as candidates for being under the control of another oscillator [8]. All three were found to be circadian here (confirming the results in [8]), and \textit{cpe-1} and \textit{cgq-16} were found to be light-responsive as well. All of them were found not to be WCC-responsive here, although in the case of \textit{cgq-16} this appears to be a false negative [59].

Lewis \textit{et al.} [60] conducted a related experiment that over-expressed \textit{wc-1}. Only 18\% of the induced genes corresponded to the ones that we detected. A similar result was seen with overexpression of \textit{CLOCK} in Drosophila [56]. There could be a variety of explanations, but it is not unexpected in a signaling system that over-expression leads to a different outcome than a knock-down, particularly when there are other coupled interacting pathways to those in Fig. 2 or 4. See, for example, the platelet-derived growth factor \(\beta\) receptor (PDGFR\(\beta\)) signaling system [63] or the sonic Hedgehog (\textit{shh}) signaling system in neurogenesis [64], where high and low levels of \textit{shh} have different neurogenetic outcomes. The response of the clock to \textit{wc-1} over-expression is apparently not the same as lowering \textit{wc-1} expression. An additional MINE calculation analogous to Fig. 12 (results not shown) also suggested that that an over-expression experiment would not be as informative as a knock-down.

The Computing Life paradigm has led us to the discovery of 328 \textit{clock-controlled genes} supported by all three series of microarray experiments and having an upstream LRE. Among these 328 genes, the chance of a false positive is \((0.18) \times (0.17) \times (0.22) = 0.0067\) (Table 1), the three microarray experiments having been done independently. A total of 104 of these 328 genes survive the multiple test correction in all three cycles [51]. These genes satisfy the three predictions of the genetic network and constitute \textit{clock-controlled genes} (Fig. 11). Of these 328 \textit{clock-controlled genes}, 314 of them are distinct on the arrays (some genes are represented multiple times; see Materials and Methods).

**Direct test of the auto-feedback loops activating \textit{wc-1} and \textit{wc-2}**

All three cycles of microarray experiments support the presence of auto-feedback loops for WCC activating \textit{wc-1} and \textit{wc-2}. In cycle 1 there was evidence that \textit{wc-1} and \textit{wc-2} mRNAs were circadian in Fig. 8. In cycle 2 there was a fast light-response by \textit{wc-1} and \textit{wc-2} (of less than an hour) in Fig. 8. In cycle 3 both \textit{wc-1} and \textit{wc-2} were WCC-responsive in Fig. 8 and have upstream LRE elements. An experiment with a short 6 hr artificial day is predicted by a MINE calculation in Fig. 9 to be highly informative about the genetic network in Fig. 4. A prediction of the genetic network in Fig. 4 is that the auto-feedback loops added should permit entrainment to a short artificial day of 6 hrs duration (as in Fig. 1) independent of FRQ. To test this hypothesis, a strain (93-4) with a \textit{frq} mutation was subjected to a short artificial day as seen in Fig. 14A. As can be seen, the rapid condensation pattern with \textit{frq} in Fig. 14A is indistinguishable from a \textit{bd} mutant in Fig. 14B. To rule out that the condensation response to a short artificial day is under the control of an independent light-response pathway, a mutant in \textit{bd}, \textit{his-3}, \textit{wc-1} (87-84-6) was generated by a cross, \textit{bd} \textit{his-3} (87-84)\times\textit{wc-1} (FGSC 3914). As can be seen (Fig. 14B), the \textit{wc-1} mutation almost entirely removed banding under the artificial day of 6 hrs. To confirm this finding, the \textit{bd}, \textit{his-3}, \textit{wc-1} (87-84-6) strain was transformed with a plasmid containing a QA-inducible \textit{wc-1} as described ([7] and Materials and Methods) and was found to band weakly when \textit{wc-1} was induced and not to band when \textit{wc-1} was not induced (results not shown). This establishes that banding under a short artificial day is under the direct clock control of \textit{wc-1}. In a similar entrainment experiment a double mutant \textit{wc-2}dm, \textit{bd} from the cross \textit{wc-2}dm\times\textit{bd} (FGSC 11124, 65)\times\textit{bd} (FGSC 1858) was subjected to a 6 hr artificial day. As can be seen in Fig. 14C, \textit{wc-2} also nearly removed all banding. As a final confirmation of these experiments, the \textit{frq} gene was over-expressed on 0.01 M QA as well as turned
off on glucose in a strain with a QA-inducible copy of the frq gene (see Materials and Methods), and these two conditions had no effect on the rapid banding (results not shown). These results serve to confirm that the rapid light response to a 6 hr day is due to the auto-feedback loops to wc-1 (and wc-2) and not due to the FRQ oscillator.

The clock mechanism of three genes wc-1, wc-2, and frq is pleiotropic in its effects on metabolism

These 314 clock-controlled genes identified are involved in a broad range of biological functions: DNA metabolism, replication, repair, cell cycle, RNA metabolism, transport, carbon and energy metabolism, isoprenoid biosynthesis (including carotenoids), development, and signaling (Fig. 15A). Periods and phases of all 295 (314–19; see below about 19) ccgs are similar in distribution to all circadian genes (Fig. 7) with upstream LREs.

The connection of the clock to development has been reported (ccg-2, ccg-4, ccg-6, and con-6). A recent connection to RNA metabolism has been through the frequency RNA helicase (fjh) gene, whose product FRH co-immunoprecipitates with FRQ [66]. The microarray experiments here identified fhj and 16 additional genes in RNA metabolism under clock control (Fig. 15B). In addition to fhj, 4 additional genes with products homologous to ATP-dependent RNA helicases in S. cerevisiae, namely ROK1, HAS1, PRP16, and RRP3, are among the 295 clock-controlled genes. At least three of these RNA helicases are involved in ribosomal RNA processing. While ribosome transcription is not under clock control (the ribosomal RNAs are not circadian in Cycle 1), almost all of the ccgs in RNA metabolism are involved in ribosome processing and assembly, i.e. ribosome biogenesis. These include SEN2, SOF1, LHP1, RRP3, POP4, UTP5, RCL1, ABD1, and PAB1 homologs in S. cerevisiae. The yeast PAB1 is a poly-A binding protein, and LHP1 is another distinct RNA-binding protein involved in the maturation of tRNAs and snRNAs. LHP1 has been implicated not only in the biogenesis of non-coding RNAs, but a recent ChiP/chip experiment in S. cerevisiae has demonstrated that

Figure 10. Transcriptional profile of approximately 1725 genes with LREs upstream at 0, 4, 8, ..., 24, 24.3333, 24.6667, 25, 26, 28, ..., 48 hrs (values staggered on x-axis) after shift from light to dark (L/D) followed by D/L transition 24 hrs later, after background subtraction, normalization within arrays relative to grand median of each chip, logging, and clustering with average linkage using Euclidean distance between mRNA profiles of different genes [38]. The bright green is −3, and the bright red is +3 is expression level on a decadic log scale. Data arose from 16 chips probed with a biotin labeled aRNA. Over 43 known clock-associated genes are overlayed in the right margin of this microarray experiment including varied known ccg genes. Genes in Fig. 2 are represented at least 5 times on each chip (explaining duplicate entries of frq, for example). The 1725 genes with upstream LREs were selected by a t-test comparing the mean of the first seven time points in the dark with the mean of time points 24,333, 24,6667, 25, 26, 28, and 32 hrs in the light with those having |t11|>1.363 (α = 0.20) displayed. The minimum observed t-value corresponded to a fold variation of 1.23 in the means before and after the light was turned back on. The mean observed t11 of the frq mRNA was 1.92. Grey bar denotes lights off; white bar denotes lights on.

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it targets a number of mRNAs [67]. The RNAse P (P0P4) binds to the RP21 RNA, which is also a target component of Lhp1p, and the PAB1 mRNA is also apparently a target of Lhp1p in yeast as well [67]. Inada and Guthrie [67] also report enrichment of the gene encoding the snoRNA U3 among targets of Lhp1p. The product of UTP5 is part of the preosome containing the U3 snoRNA and involved in ribosome biogenesis. The clock’s regulation of the ribosome appears to occur through its biogenesis rather than its transcription. This is a novel mechanism by which the clock can regulate clock-controlled genes post-transcriptionally.

Connections of the clock to DNA metabolism are recently reported in humans [68]. A human clock CLK2 protein physically associates with S-phase checkpoint components ATR, ATRIP, clasinp, and the checkpoint kinase, Chk1. Also human CLOCK-depleted cells accumulate DNA damage, engage in radiosensitive DNA synthesis, and fail to recruit proteins, such as RAD51, functioning in human recombination pathways. Here several putative checkpoint-associated proteins (e.g., NCU00560 and NCU04326) as well as 8 genes involved directly in purine/pyrimidine metabolism (NCU0 7590, 8359, 4323, 6262, 3194, 5542, and 4195) and repair (ncw-6/NCU00901) appear to be clock-controlled genes. The ncu-6 gene is a homolog of RAD50 in S. cerevisiae involved in double-stranded break repair. As predicted from the results on humans [68], the RAD51 homolog (mer-3/NCU02741) was circadian and light-responsive in N. crassa in cycles 1 and 2, but not WCC-responsive (cycle 3).

The clock connection to the cell cycle has been only recently reported in Neurospora through pdt-4, a homolog of Chk2 in yeast, a second checkpoint kinase [53]. In addition to pdt-4, we have identified 2 other putative cell cycle checkpoint genes as clock-controlled genes (NCU00560 and NCU04326), homologs of CDC4 and CDC28 in S. cerevisiae. Up to 16% of rhythmic genes (cycle 1) may be involved in the cell cycle in some mouse tissues in contrast to the 3% in Fig. 1A identified by more stringent criteria (i.e., positive in cycles 1–3) in N. crassa [69].

In that carbon metabolism showed up as significant and may have arisen due to the use of the QA-inducible switch in the last series of microarray experiments, one additional control was performed with wild type (OR74A – see Materials and Methods) in which many QA-inducible genes were identified with micro-arrays by a shift from 1.5% sucrose to 0.3% QA over a 0 to 9 hr window [11,27]. Of the 314 distinct clock-controlled genes identified, only 19 of them were QA-inducible (with most of them being unclassified in function). Only 2 of the QA-inducible clock-controlled genes were involved in carbon metabolism. Subtracting the 19 QA-inducible ccs from the 314 distinct ccs, 295 clock-controlled genes remained.

Approximately one-half of these 295 clock-controlled genes are of unknown function. The most prevalent known function among these genes are phosphatases and kinases. They make up almost half [11] of the 23 genes with products involved in protein synthesis (Fig. 16B), processing, and degradation, and at least three of the genes under DNA metabolism are known kinases/phosphatases as well (CK1, HHPI homolog, and PPI). This plethora of phosphatases and kinases may reflect the role they play in modifying/linking the functions of wc-1, wc-2, and frq as regulators of (to) other pathways, as well as in the coupling of the clock to varied signaling pathways and the cell cycle. For example, the phosphatases PPI and PPIa, dephosphorylate FRQ in vitro, thereby altering oscillator behavior [70], and the kinases CK1α and CKII mediate the phosphorylation of WCC [71]. After that, DNA metabolism, RNA metabolism, and carbon/energy metabolism represent equally important outputs of the clock. The clock outputs are representative of the frequency of these functions in the proteome [37] with two exceptions: a deficit of transport and unclassified genes that are ccs.

While only one ccc in Fig. 15 has a product classified as a transcriptional activator (PRO1 homolog) involved in fruiting body development (kal-1, NCU07392, [72]), four other ccs genes were classified as regulators. Their individual cyclical transcriptional profiles are given in Fig. 16A. One of these putative regulators is an inferred ornithine decarboxylase antizyme involved in sulfur and nitrogen regulation (NCU07155) [37]. This connection has also been reported in Arabidopsis [47,73]. The remaining putative regulators identified under clock control in Fig. 2 or 4 were NCU00045, NCU01640 (rpm-4), and NCU06108.

Earlier work has suggested a link between signal transduction pathways for conidiation and the clock [74]. From microarray analysis here the clock is tied into a number of other signal transduction pathways as well, including stress (cg-9/NCU09559), oxidative stress (NCU05169), light (lid/NCU03967), mating (mat/ NCU02500, NCU03378, NCU07335), and osmo-sensing (os-1, rrg-1, hpt-1). The last output to the clock has only been reported recently (Fig. 16C). For example, cat-1 involved in osmo-sensing has been reported to be under WCC control [75]. Jones et al. [57] have reported a role of rrg-1 in osmo-sensing reminiscent of the os mutants. The genes rrg-1 and hpt-1 have an upstream LRE and were found to be circadian in cycle 1, but not light-responsive or WCC-responsive [58].

Impact of standard that the period of a ccc is between 16 and 30 hrs

To provide insights on the impact of the standard [5] of requiring genes to have a period between 16 and 30 hrs to be declared clock-controlled genes, we tightened the standard to 17–29 hrs because of the up turn at the extremes of the distribution of periods in Fig. 7A. The result was declaring 2172 genes with QA-inducible switch identified in cycle 1 and 2, with 23 of the 314 clock-controlled genes identified.
three cycles of microarray experiments to 290. If the 14 duplicates of *frq*, *wc-1*, and *wc-2* as well as 18 remaining QA-inducible genes are removed, we are left with 258 clock-controlled genes with a range of periods between 17 hrs and 29 hrs. Genes affected in the text can be determined from their reported periods. Further trimming of the range of acceptable periods in Fig. 7A will gradually shrink the number of clock-controlled genes identified.

**Identifying an ensemble of genetic networks for the biological clock of N. crassa**

The culmination of Computing Life is the identification of an ensemble of genetic networks describing how the clock functions from 3 cycles of microarray experiments initiated from published data [3,23–25]. Results are summarized in Table 2. For 69% = (100\times18/26) of the rate constants in common with Table 1 [7], standard errors were reduced by the addition of data from cycles 1–3. Measured lifetimes of the *wc-1* mRNA and FRQ protein remain concordant with estimated values in Table 2 with an order of magnitude increase in the amount of data (see Table 3). The long lifetime of the *wc-1* mRNA provided a critical test of the genetic networks [7], and the long lifetime of the *wc-1* mRNA of 7.4 hrs = $D_7^{-1}$ continues to be supported by microarray data here.

Transcription rates of *frq*, $A$ and $A'$, as well as the deactivation rate of WCC, $P$, were previously identified as critical parameters for maintaining oscillations through the negative feedback loop in Fig. 2 [7]. These constants are now more sharply defined in Table 2. Eleven of the 26 parameters identified in Table 1 of [7] are not significantly different from those in Table 2, although a majority of the rate constants are estimated more precisely. Precision of cycles 1–3 are assessed further in the next section and Table 3.

The behavior of the ensemble is displayed in Fig. 17. In cycle 1 the predicted oscillations of *frq* mRNAs are displayed with microarray measurements. The predicted oscillations in *wc-1* and *wc-2* mRNAs are much reduced relative to the *frq* mRNAs. In cycle 2 in the first 24 hrs the measurements and predictions track those in cycle 1; the correlation over all 87808 = 12544 $\times$ 7 microarray features between cycles 1 and 2 for the first 7 time points is 0.82. When the light is turned back on at 24 hrs into the cycle 2 experiment, the coordinated response of the ensemble and microarray data (particularly the *frq* mRNA) to light can be seen as the clock resets [62]. In cycle 3, the slow decline in the *wc-1* mRNAs is seen corresponding to a lifetime of 7.4 hrs = $D_7^{-1}$. An alternative ensemble in Fig. 2 from Yu *et al.* [7] was tested and

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**Figure 12. A 90% knock-down of the *wc-1* gene is the MINE experiment.** The decadic log of the MINE criterion $\det(E)$ is displayed as a function of percent remaining activity of the three clock genes *wc-1*, *wc-2*, and *frq*. The matrix $E$ is the correlation matrix of the predictions, emphasizing independence of predicted data points $f_i(u_i)$. The predictions are for the mRNA levels of *wc-1*, *wc-2*, and *frq* over time. doi:10.1371/journal.pone.0003105.g012
performed about the same as the genetic networks in Fig. 4; the distribution of chi-squared statistics [7] for the ensemble fitted to Fig. 2 is largely overlapping with chi-squared statistics for the networks in Fig. 4 (results not shown). By Occam’s Razor the simpler network with fewer parameters in Fig. 2 is then preferred. The series of model-guided experiments now has identified and selected an ensemble of genetic networks describing the clock mechanism in Fig. 2.

**Comparison of the precision and power of microarray experiments in cycles 1–3 with other microarray experiments**

A standardized way of assessing progress in the Computing Life paradigm as well for comparing the power of different microarray experiments here with others in the literature and in the future would be useful. Progress here is measured by the error per observation $\sigma^2$ or error variance.

In linear and nonlinear models a standard approach to estimating the precision of an experiment is to estimate the error variance $\sigma^2$ [76], as it appears in the likelihood for the genetic network [7]. Townsend [77] illustrates by simulation and data analysis that such a common variance component can be extracted from each of a variety of microarray experiments and used to compare different experiments. Under the multivariate Gaussian assumption leading to the likelihood in [7], a simple estimator for the error per observation $\sigma^2$ can be constructed for successive cycles of the Computing Life paradigm:

$$\hat{\sigma}^2 = \frac{1}{n} \chi^2_{\text{min}} \sigma_0^2$$

where $n$ is the number of observations, $\chi^2_{\text{min}}$ is the minimum chi-squared statistic over the ensemble [7], and $\sigma_0^2$ is a preliminary estimate of the error per observation in the multivariate Gaussian.
Figure 14. Light entrainment response under short artificial day by frq and wc-1 mutations provides evidence for autofeedback loops on wc-1 and wc-2 in Fig. 2. (A) Light entrainment response of a frq, bd mutation (FGSC 93-4) during artificial days with 3 hrs of light followed by 3 hrs of dark in race tubes. (B) Light entrainment response of a wc-1, bd mutation (87-84-6) during artificial days with 3 hrs of light followed by 3 hrs of dark in race tubes. (C) Light entrainment response of a wc-2, bd mutation [65] during artificial days with 3 hrs of light followed by 3 hrs of dark in race tubes. A bd mutation was cultured in race tubes as a control. See Fig. 1 as well for “wild type”, namely the bd mutation.

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Discussion

What we know and do not know about clock-controlled genes

Model-guided microarray experiments through MINEing have revealed much of what we know and do not know about the biological clock in Fig. 11. At the center of the Venn Diagram there is a highly cross-validated set of 295 distinct clock-controlled genes behaving as the clock mechanisms in Fig. 2 or 4 would predict. To date, only sixteen clock-controlled genes have been discovered in N. crassa in over 40 years of clock biology [59]. This set of 295 genes is circadian, light-responsive, and under WCC control and spans a broad array of functions (Fig. 15). It is quite remarkable that only three genes, wc-1, wc-2, and frq, could have such diverse and pleiotropic effects on the organism’s transcriptome, and the full extent of the clock’s role in the metabolic web has not been evidenced till now.

The series of model-guided array experiments also point to an intriguing set of 440 genes or 57% ($= 100 \times 440 / (328+440)$ in Fig. 11) of the genes that are circadian and light-responsive and not under direct WCC control. No more than half of these 440 genes can be explained by false negatives in the cycle 3 experiment assaying for WCC control according to Table 1. These genes could be under the indirect control of the clock mechanism. Some candidate regulators implicated by all three microarray experiments include five genes, NCU00045, NCU01640 (gpa-4), NCU06108, NCU07155, and NCU07392 (kal-1). One of these may be a regulator of nitrogen and sulfur metabolism (NCU07155), and another is a homolog to PRO1 (called kal-1), a homeo-box containing transcription factor with a typical GALL4-like DNA binding domain involved in fruiting body development [72]. Another possibility is that some of these mRNAs are modified post-transcriptionally by RNA-binding proteins which are cegs, such as the LHP1 and PAB1 homologs, to control expression. In that RNA-binding proteins appear to have specific mRNA target populations [78], they may provide another mechanism for the combinatorial clock control of gene expression post-transcriptionally.
Figure 15. The clock of *N. crassa* has 295 distinct clock-controlled genes of diverse function as outputs. A. 328 Genes (with putative WCC binding sites) which are circadian, light-responsive, and WCC-responsive are classified by function (MIPS, [37]) in the outer wheel. 19 QA-inducible genes are not included in the outer wheel. With the exception of the unclassified category, the standard errors on the percentages of the outer wheel are 0.01–0.02. Gene products of the *N. crassa* proteome are classified by function as well (MIPS, [37]) in the inner wheel. The numbers below are a MIPS coding of functional categories. The definition of the categories [37] is DNA, 1.03, 10.01; cell cycle, 10.03, 10; transcriptional control & regulation,
There are WCC-responsive genes that are circadian only (612), but not light-responsive as well as those that are light-responsive only (396), but not circadian in Fig. 11. The genes, which are circadian only and not light-responsive, could be explained by other regulators involved with the clock mechanism that suppress a light-response (or be under the control of an oscillator not connected to the light-response through WCC as suggested by Correa et al., [8]). The possibility of an oscillator not connected to WCC is discussed in the next section below. Similarly, genes that are light-responsive only, but not circadian, might also be explained by regulators of the light-response that interact with WCC to repress a circadian response (or be part of a light-response pathway not connected to the known clock mechanism, but see below). One potential list of candidates as modulators of the circadian- and light-response are the 11 phosphatases and kinases that are clock-controlled genes in Fig 15. These two functions of being circadian and being light-responsive can be separated in the action of WCC through the outputs of the clock. Other kinds of posttranslational modifications of histones could be involved in chromatin-remodeling of upstream sequences to clock-controlled genes to determine gene activation through WCC [79].

Data on wc-1 cis-regulation by Kaldli et al. [56] and on frq cis-regulation by Belden et al. [79] suggest a simpler hypothesis that invokes only one new regulator class. The wc-1 gene has at least two kinds of promoters upstream responding to WCC. The CCAAT box promoter in front of wc-1 is not light-responsive, while the TAATTA promoter in front of wc-1 for WCC is light-responsive. The LREs determine light responsiveness [4], while other genes, such as clockswitch (caw-1), regulate accessibility to promoters through chromatin-remodeling around the Clock Box or C box upstream of frq to determine the endogenous circadian rhythm. Froehlich et al. [80] have shown that the C box is necessary and sufficient for endogenous circadian expression of frq. An additional LRE more proximal to frq is light-responsive [4]. Genes (295 out of 328) that have both types of elements, C boxes and LREs, might be circadian and light-responsive. Genes (612) with a C box and no LRE might be circadian but not light-responsive. Light-responsive genes (396) could be explained by only having the LRE upstream. This alternative would also not require any new regulators beyond the enzyme GSV-1 enabling chromatin-remodeling around the C box. In Table 4 there is a significant association of genes being light-responsive and not circadian vs. being circadian and not light-responsive and the number of upstream LREs. Unfortunately, there was no difference in the incidence of the known Clock box among the circadian/ non-light-responsive (612) versus the 396 light-responsive/non-circadian genes. Further characterization of Clock boxes in front of other genes besides frq appears warranted.

Fig. 11 would suggest considerable limits on what we know about the clock mechanism. There are 1056 genes that are circadian only and not WCC-responsive and 361 genes that are light-responsive only and not WCC-responsive, and no more than half of these can be explained as false negatives in cycle 3 (WCC response array experiment). These two sets of genes are not explainable directly by WCC. A similar observation was made on CLOCK in Drosophila [36]. This percentage of the unexplained has increased from the earlier array experiments carried out by Lewis et al. [60]. This would suggest that there are other genes involved in the biological clock mechanism and other genes involved in a light response.

From the model-guided array experiments there is a concluding suggestion on a new direction for MINEing for clock-controlled genes. There are 967 genes that apparently are not circadian or light-responsive, but under WCC control. The clock is well-known to respond to a variety of environmental cues or zeitgebers, such as temperature [81]. The fact that these 967 genes do not respond to light would suggest future cycles of discovery to examine how these 967 genes respond to temperature [82].

The circadian response of genes with upstream LREs is stochastically independent of their light response conditional on the WCC response

The microarray experiments in cycles 1–3 allow us to infer a broader relationship of circadian, light, and WCC-responses by different genes in the genome from Fig. 11. We have established in Fig. 11 that the circadian and light-responses can be separated by the response of clock-controlled genes. It is natural then to ask how are these responses related in different genes? For example, we can ask if the circadian and light responses of different genes with LREs upstream are stochastically independent across all genes with an LRE element. A simple contingency table model in which the circadian and light- response of each of the 5702 genes with LRE elements are conditionally independent given the WCC response can be fitted as shown in Table 5 fairly well. The resulting goodness of fit chi-squared statistic $X^2 = 7.20$, which is barely a significant departure from the model at the 0.05 level. What this implies is that the circadian and light responses of genes with LREs upstream are almost entirely explained by whether or not they have a response to WCC. When the WCC response is off, then the circadian response of a gene with an upstream LRE is more probable ($p_C > p_L$ in Table 5). When the WCC response is on, then the light response of a gene with an upstream LRE is more probable ($q_L > q_C$ in Table 5). This would suggest that any gene, such as caw-1 [79], acting on the light or circadian response of a gene with an upstream LRE would need to act through WCC. The only explanation for light and circadian responses of egs in Fig. 2 or 4 is WCC. If there were another oscillator acting on genes with upstream LREs, then its outputs would be constrained to behave in the same way as the known clock mechanism. That is, an additional oscillator would be coupled to the known clock mechanism in Fig. 2 through WCC and would have an independent light and circadian gene outputs conditional on a gene’s response to WCC. If there were additional information, such as the number of upstream LREs (as in Table 4), then more could be said. This conditional independence result does not hold across all 11,000 genes, if the same model in Table 5 is fitted to all genes ($X^2 = 771.70$, P<0.001).

Clock as adaptation

The clock-controlled genes and their time of action in Fig. 7B provide a possible narrative on adaptation. This complex trait controls levels of asexual reproduction as shown in Fig. 1. The complex trait has a clear genetic basis in Fig. 2. Variation in the clock trait in natural populations has been demonstrated in D.
Figure 16. Transcriptional profiles of individual genes with upstream LRE elements in the dark (cycle 1) in the functional categories of: (A) regulation (MIPS functional classification categories 1.02.04, 11.02.03, and 16, [37]); (B) putative phosphatases/kinases (functional classification category 14.07.03); (C) signal transduction (categories 30.01, 32.01, 32.05, 34.11); (D) Development and growth (categories 40.01, 40.02, 41.01, 43.01). The mean mRNA level of each gene was subtracted from the 13 individual mRNA levels measured on each gene in Fig. 6. Data are from Fig. 6. doi:10.1371/journal.pone.0003105.g016

Table 2. Rate coefficients in the genetic network model (Fig. 4) of the biological clock (n = m = 4) based on data from cycles 1–3 predicting the clock’s observed oscillations, light response, and wc-1 perturbation (Fig. 17).

| X     | k       | <X>     | s(X) |
|-------|---------|---------|------|
| A     | 5       | 0.0313  | 0.00974 |
| B     | 5       | 4.010E-4 | 1.020E-4 |
| T     | 1       | 0.1108  | 0.00498 |
| S1    | 1       | 0.000420| 0.00000048 |
| S2    | 1       | 0.0220  | 0.00838 |
| S3    | 1       | 5.47E-5 | 1.597E-4 |
| S4    | 1       | 1.252   | 0.286 |
| D1    | 1       | 6.607   | 1.399 |
| D2    | 1       | 0.153   | 0.0247 |
| D3    | 1       | 0.798   | 0.134 |
| C1    | 2       | 1.047   | 0.220 |
| L1    | 1       | 94.39   | 4.346 |
| L2    | 1       | 0.3698  | 0.2207 |
| L3    | 1       | 63.93   | 21.5 |
| D4    | 1       | 0.00451 | 0.0118 |
| D5    | 1       | 0.00890 | 0.00242 |
| D6    | 1       | 0.205   | 0.00899 |
| D7    | 1       | 0.135   | 0.0148 |
| D8    | 1       | 0.0122  | 0.00304 |
| C2    | 2       | 3.322   | 0.912 |
| P     | 5       | 0.2233  | 0.2701 |
| A_1   | 5       | 0.1293  | 0.0826 |
| B_1   | 1       | 0.6091  | 0.1718 |
| S_1   | 1       | 2.572   | 2.757 |
| L_1   | 1       | 3.664   | 8.993 |
| D_1   | 1       | 0.579   | 0.137 |
| D_2   | 1       | 0.5536  | 0.1173 |
| E1    | 2       | 0.003125| 9.863E-4 |
| E2    | 2       | 0.0965  | 0.0104 |
| E3    | 2       | 2.614   | 2.607 |
| E4    | 2       | 0.0128  | 0.0298 |
| S_5   | 1       | 8.924   | 0.696 |
| D_9   | 1       | 1.234E-4| 3.259E-4 |
| A_c   | 5       | 0.0524  | 0.0156 |
| Q     | 5       | 4.812E-4| 6.111E-4 |
| D_o   | 1       | 2.865E-4| 9.257E-4 |
| E_1   | 2       | 0.00974 | 0.00288 |
| E_2   | 1       | 0.000542| 0.00188 |
| E_3   | 2       | 1.335E-5| 3.456E-5 |
| E_4   | 2       | 0.0121  | 0.00682 |

Ensemble mean <X> and ensemble standard deviation σ(X)=|<X>-<X>^2|1/2 for rate coefficients (X) in the n=m=4 biological clock model of Fig. 4. For a kth order reaction (with k = 1, 2, or 5), the rate coefficient is given in units of 1/(hour^k cu^k1) where “cu” represents the arbitrary, but common model unit of concentration for all species, except for the photon species where 1 cu photons = 0.20 μmole photons/s m^2. See also Materials and Methods. The estimated value of 1/(D_0)=5 hrs is consistent with the FRQ protein life-time of ~4–7 hrs, estimated from the FRQ-decay data of [96]. doi:10.1371/journal.pone.0003105.t002

Systems Biology

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Setting the phase may set the level of WCC experienced by the ccg of different clock-controlled genes as shown in (Fig. 7B). This adaptationist interpretation of the role genetic network for ccg has been reported in other model systems [90].

Experiments. There appears to be a strong connection between the developmental program is timed through clock-controlled genes immediate genes (IEGs)'' in the evening. Implementation of the structures to produce these spores requires carbon and energy as synthesis must be reinitiated and completed before entry into cell-cycle initiation in the morning to produce the spores, DNA is positioned in the Central Dogma played out in ribosome assembly, protein synthesis, modification, and degradation taking place in the morning. One of the features of spores that make them so resistant to environmental insults is the synthesis and incorporation of the isoprenoids as pigments. Apparently their biosynthesis is necessary to obtain more information about the genetic network in each successive cycle through the Computing Life paradigm. This still leaves the choice of a set of possible experiments, from which U* is drawn, in the hands of the experimenter. In the present case, the biological goal of identifying clock-controlled genes sets the stage on which MINE plays. Identifying clock-controlled genes leads us to ask the genetic network in Fig. 4 for their predicted behavior, thus establishing the set of experiments in cycles 1–3 to be considered. In this context MINE is a tool to achieve a particular biological goal. One could naively make the biological goal coincident with the unconstrained objective of learning the most about the genetic network, the criterion of MINE, but then the set of possible experiments U becomes very large and the optimization of V(U) computationally intractable. Adopting a particular biological goal, such as finding ccg, puts structure on the design question and thereby enables the researcher to parameterize the optimization of the next experiment U.

There have been several consequences to the use of model-guided discovery by MINE through the choice of experiments. In detecting circadian rhythms in cycle 1 MINE involved a design with even spacing between observations of 4 hrs and with sampling starting immediately without delay (Fig. 5). This design in cycle 1 conforms to practice in previous experiments [5]. However, in detecting the light-response in cycle 2 and the WCC-response in cycle 3 there is a departure from conventional wisdom. In cycle 2 prior experiments on light-entrainment, researchers have avoided a very short 6 hr artificial day [25], and as a consequence have missed an opportunity to examine the autofeedback in the networks as in Fig. 14. In detecting a response to perturbations in the clock mechanism in cycle 3, conventional wisdom would have us focus on perturbing the oscillator gene frq [8]. In contrast the MINE calculations in Fig. 12 suggested that a knockdown would be more informative than overexpression. The result has been a paucity of responding genes [36,60]. The MINE calculations in Fig. 12 suggested that a knockdown would be more informative about the genetic network in Fig. 2 or 4, and this was the set of possible experiments that are ultimately specified by the goal(s) of the experimenter. Within the constraint of a particular biological goal, such as finding clock-controlled genes, finding the MINE U* provides an avenue to obtain the most information about the genetic network in each successive cycle through the Computing Life paradigm.
experiment adopted here. As a consequence, by following conventional wisdom without the input of MINE, it is likely our screen would have missed the clear and unbiased identification of 295 clock-controlled genes in Fig. 15 or as precise an identification of the genetic network (see Table 3). The connection of the clock to ribosome biogenesis in Fig. 15B in particular is then an outcome of MINE suggesting a new direction of exploration in microarray experiments to refine our understanding of the clock mechanism.

It may at first seem surprising, or accidental, that MINE design actually helps increase the experimental sensitivity for ccg detection: the MINE approach, as described in Materials and Methods, optimizes the experimental sensitivity for discerning between unknown model parameter vectors $\Theta$; it does not per se optimize the next experiment for detection of new molecular or gene species (such as ccgs) that are not explicitly included in the network model in Fig. 2. However, these two seemingly unrelated features are actually closely linked, as we will now explain.

The MINE optimization does tend to select experimental conditions $U^*$ which enhance the predicted response for the molecular species to be observed in the next experiment. (If the to-be-observed species were responding weakly their observation would hardly improve discrimination between different $\Theta$). One of those to-be-observed molecular species in each of our 3 MINE cycles was the $frq$ RNA. The $frq$ gene, however, is co-regulated (by WCC) with all clock-controlled genes. For purposes of external perturbation response, the $frq$ gene itself is in fact like a typical ccg. Thus, by choosing the MINE $U^*$ to enhance the $frq$ response we are in effect also enhancing the response of all other ccgs, since they are subject to similar regulatory control. Hence, the MINE-directed discovery of our 295 new clock-controlled genes is not accidental; it is an expected and highly desirable by-product of the MINE optimization. The broader conclusion to be drawn from these considerations is that, for gene-regulatory systems, MINE optimization, in general, will improve the conditions for

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**Figure 17.** An ensemble of genetic networks predicts the mRNA levels of $wc-1$, $wc-2$, and $frq$ for cycles 1–3. The decadic log (lg) of each gene’s mRNA level is measured at least 5 times on an array for each time point. Some data points are from the literature [3,23–25]. The curves represent the mean prediction of the ensemble of genetic networks in Fig. 4 +/- 2 ensemble standard errors about the ensemble mean. In Fig. 8 the averages of the 5–6 replicates of each mRNA level at each time points are displayed. Grey bars denote lights off; white bars denote lights on. doi:10.1371/journal.pone.0003105.g017
experimental detection of new, unknown gene species which are co-regulated with the already known species targeted by the next experiment.

There are three advances here over earlier analyses of the genome-wide logical circuitry [1,91,92] of fundamental processes, such as development, carbon metabolism, and the clock. One, a working kinetics model of the clock [7] gives a complete quantitative description of genetic network dynamics in Fig. 17. Two, an ensemble method was developed for identifying a genetic network with many parameters from limited data [7] with the results in Table 2. Three, a new methodology (i.e., MINE design) was developed for a model-driven discovery workflow cycle (Computing Life) in profiling experiments. This new methodology resulted in the identification of most downstream clock-controlled genes in Fig. 15A and an unexpected connection between the clock and ribosome biogenesis in Fig. 15B.

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Author Contributions

Conceived and designed the experiments: RN JA HBS. Performed the experiments: WD YY RN RK JG JA. Analyzed the data: WD XT RN JA HBS. Contributed reagents/materials/analysis tools: XT RN JA HBS. Wrote the paper: RN JA HBS.

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Table 4. There is an association between whether or not a WCC-responsive gene is light-responsive and the number of its LREs upstream.

| Gene Response | Number of LREs upstream |
|---------------|-------------------------|
|               | 1       | 2       | 3       | >3      | N       |
| circadian/no light-response | O 405 | 137 | 47 | 23 | 612 |
| light response/not circadian (cycle 2) | O 228 | 91 | 46* | 30* | 396 |

WCC-responsive genes are classified as circadian and not light responsive or not circadian and light-responsive and by their number of putative upstream WCC-binding sequences (GATX sequences) or LREs. Observed (O) and Expected (E) counts of these genes under the hypothesis of no association are reported along with the chi-squared test of no association. See Fig. 11 for categories of LRE counts conditioned on N (N = 612 or 396).

The chi-squared statistic is $X^2 = 12.8445$ with df = 3 and critical value $C = 11.34$ for $\alpha = 0.01$. The residuals (O–E)/sqrt(E) of the starred counts make a large contribution to the $X^2$. doi:10.1371/journal.pone.0003105.t004

Table 5. The circadian response of each gene with an upstream LRE is conditionally independent of its light response.

| Gene Response | Not WCC-responsive | WCC-responsive |
|---------------|--------------------|----------------|
|               | NL L               | NL L           |
| not circadian | O 1337 561 | 986 396 | E 1338 560 | 957 425 |
| circadian     | O 1056 440 | 612 314 | E 1055 441 | 641 285 |

The independence is achieved by conditioning on the WCC-response. Genes are classified as “not circadian” or “circadian,” “not light-responsive” (NL) or “light-responsive” (L), or not WCC-responsive or WCC-responsive (WCC) and Expected (E) counts of these genes under the hypothesis of no association between the circadian and light responses conditional on the WCC-response are reported along with the chi-squared test of no association. See Fig. 11 for categories of genes by category of response. The no response category was computed as $5702–4365$ (sum of counts in the remaining 7 categories). All duplicates of wc-1, wc-2, frq, and other genes were removed from the observed counts of genes (O) with upstream LREs in Fig. 11 to be classified into 8 categories below. The probability estimate of a gene being WCC-responsive is $p_C = (2308/5702)$. The probability estimates of a gene being circadian or light-responsive when it is known not be WCC-responsive are $p_{L} = (1496/3394)$ or $p_{L} = (1001/3394)$, respectively. The probability estimates of a gene being circadian and light-responsive when it is known to be WCC-responsive are $p_{L} = (926/2308)$ or $q_{L} = (710/2308)$. These probability estimates can be used to compute the expected (E) counts below.

As an example $441 = p_C \times p_{L} = (1496/3394) \times (2308/5702) = 1176/5702$. The chi-squared statistic is $X^2 = 7.20$ with df = 2 and critical value $C = 5.99$ for $\alpha = 0.05$. doi:10.1371/journal.pone.0003105.t005
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