Identifying sources of variation and the flow of information in biochemical networks

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AUTHOR SUMMARY

Cells respond to change stochastically because biochemical mechanisms are stochastic: Intermolecular collisions trigger biochemical events at seemingly random times. Yet cells can both control and exploit such stochasticity. To understand how, we must identify the sources of stochasticity and distinguish variation due to information flow, say from the extracellular environment, from that due to confounding “noise”. Here, we present a general scheme for dynamic, stochastic systems to decompose variation into as many components as there are potential sources of stochasticity. Further, we identify a particular component that measures information flow and describe conditions that experimental probes, or “reporters,” should satisfy to determine this component and all others.

Intuitively, the effect of fluctuations in one stochastic variable on the variation in a variable of interest can be determined by comparing the expected variation in this variable when the first, “explanatory” variable freely fluctuates to that when it is somehow “fixed”. Our general decomposition of variance incorporates this idea by mathematically “fixing” the explanatory variable by conditioning on its history (i.e., its current value and its values at all previous times) (1). If we would like to determine how fluctuations of n sources affect the variation of an “output” variable, Z, then the decomposition has n + 1 terms: one term corresponding to each of the n variables and another term to describe any additional variation in Z that is generated through other sources.

For biochemical systems, we describe conditions that reporters should satisfy to measure each of the components. A reporter, such as a fluorescently tagged protein, is needed to measure the output of the system. To assess the effect of fluctuations in a variable, Y, on the output, Z, we must construct a second “conjugate” reporter so that its level only correlates with the level of Z because of fluctuations in the levels of Y (the two reporters are conditionally independent, given the history of Y). Then, provided that the reporters have the same conditional means and the same conditional variances, both the covariance of the two reporters and their mean squared difference describe terms in the decomposition. Therefore, for n terms in the decomposition, n reporters are needed.

To illustrate our theory, we define transcriptional and translational variation and show how such variation can be measured experimentally using fluorescent proteins translated from a bicistronic mRNA (Fig. P1). For a standard model of gene expression, we also provide mathematical expressions for these components by...
including conjugate reporters in the mathematical model we analyze. We expect this analysis technique to be widely applicable for evaluating variance components using stochastic models of biochemical networks.

Much gene expression is initiated by signaling networks, and we go on to explore how fluctuations in the environment and upstream signaling can affect downstream gene expression. Using a three-component decomposition, we reanalyze previous measurements of the response of budding yeast to extracellular pheromones (2) and conclude that, in this example, variation from signal transduction is less substantial than variation arising from other processes extrinsic to gene expression.

Variation in gene expression is also a consequence of information flowing through signaling networks from the extracellular environment. We identify a component in our decomposition whose size relative to the variance of the output, $Z$, typically indicates how readily a network can “decide” the state of the extracellular environment from the level of $Z$. For an environmental input, $X$ (for example, the concentrations of a collection of ligands), we express this informational fraction of the output variance as $V\{E[Z|X]\}/V[Z]$, where $E$ denotes expectation and $V$ denotes variance. As the informational fraction tends to its maximum value of one, the distributions of outputs corresponding to each state of the input become more distinct from one another. Consequently, it is easier to unambiguously identify the state of the input: More values of output can be uniquely identified with an input state because the overlap between the (conditional) output distributions decreases.

We apply our results to measurements of sensing of osmotic stress by budding yeast (3) and show that informational variation can be the dominant source of variation. As much as 80% can be informational if we include variation generated in response to changes in the environment’s osmolarity. By finding probability distributions of osmotic stress that maximize the informational fraction we can, with some caveats, make predictions about the types of environment most faithfully detected by budding yeast’s osmosensing network. These environments have frequent low levels of osmotic stress and infrequent high levels.

A challenge when investigating variation in any system is the influence of the wider stochastic environment in which the system is embedded. Although previous work decomposed variation into intrinsic and extrinsic components (4, 5), there was no means to identify the processes generating variation or to quantify their effects. Our general decomposition and conditions for conjugate reporters provide just such a means. They are essential knowledge for experimental design: Although constructing reporters may sometimes be difficult, these difficulties may not even be apparent, let alone solved, without precisely knowing the properties required of the reporters. Our results hold for any stochastic dynamic system at any point in time and, taken together, thus provide a general mathematical foundation for studies of cellular variation.