Explicit Tracking of Uncertainty Increases the Power of Quantitative Rule-of-Thumb Reasoning in Cell Biology

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ABSTRACT Back-of-the-envelope or rule-of-thumb calculations involving rough estimates of quantities play a central scientific role in developing intuition about the structure and behavior of physical systems, for example in so-called Fermi problems in the physical sciences. Such calculations can be used to powerfully and quantitatively reason about biological systems, particularly at the interface between physics and biology. However, substantial uncertainties are often associated with values in cell biology, and performing calculations without taking this uncertainty into account may limit the extent to which results can be interpreted for a given problem. We present a means to facilitate such calculations where uncertainties are explicitly tracked through the line of reasoning, and introduce a probabilistic calculator called CALADIS, a free web tool, designed to perform this tracking. This approach allows users to perform more statistically robust calculations in cell biology despite having uncertain values, and to identify which quantities need to be measured more precisely to make confident statements, facilitating efficient experimental design. We illustrate the use of our tool for tracking uncertainty in several example biological calculations, showing that the results yield powerful and interpretable statistics on the quantities of interest. We also demonstrate that the outcomes of calculations may differ from point estimates when uncertainty is accurately tracked. An integral link between CALADIS and the BioNumbers repository of biological quantities further facilitates the straightforward location, selection, and use of a wealth of experimental data in cell biological calculations.

INTRODUCTION

Rule-of-thumb, or back-of-the-envelope, calculations are of great utility across the sciences, allowing estimates of quantities to be obtained while gleaning intuition about the important numerical features of a system. In physics, the paradigm of the Fermi problem has been used for decades to develop intuition about the structure and behavior of systems by employing reasonable approximations, order-of-magnitude estimates, dimensional analysis, and clearly stated assumptions. The use of the napkin (often more readily available than an envelope in modern cafés and conferences) as a medium to perform rough calculations and gain understanding of a system given limited experimental information is well known in the physical sciences and has recently gained popular attention (1). Recent mathematical approaches to complex problems in wider scientific fields have employed these back-of-the-envelope approaches, including bioestimates in physical biology (2) and cell biology (3) and the popular “street-fighting mathematics” for use throughout the sciences (4).

However, these calculations currently do not have as central a role in cell biology as they do in the physical sciences, despite receiving substantial recent attention as powerful tools for reasoning in quantitative biology (5,6), and being facilitated by quantitative resources like the excellent BioNumbers database (7). One reason for this absence is that many of the quantities involved in cell biology are either intrinsically highly variable or have large measurement errors. Calculations that do not take these uncertainties into account (yielding a mean value estimate without associated uncertainties), although powerful in their own right, may represent only part of the story (Fig. 1A).

In some back-of-the-envelope circumstances, accuracy may be maintained without the explicit tracking of uncertainties. An example of this is in calculations involving the multiplication of several terms, each of which may be reasonably assumed to be normally distributed with similar coefficients of variation. In such a calculation, the logarithm of the error in an estimate scales with the square-root of the number of terms in the calculation. However, quantitative cell biology often involves distributions that cannot be assumed to be normally distributed, as well as calculations more general than simple multiplications of terms. In these circumstances, where individual uncertainties can differ between terms and may be over many orders of magnitude, the risk of inaccuracy associated with calculations without uncertainty is increased. If uncertainties are included in such calculations, it is often through standard propagation-of-uncertainty approaches (8), which typically track a limited
number of distribution moments and can thereby fail to accurately represent the distribution of the final result for nonnormal distributions. Of course, the process of performing rough calculations and obtaining estimated answers is immensely valuable in its own right, for the reasons discussed above. To complement this powerful process of Fermi reasoning in biology, we here suggest a complementary form of envelope reasoning, allowing for calculations including uncertain quantities.

FIGURE 1 An example back-of-the-envelope calculation. Current technology is unable to measure the number of protons in a cell, so we estimate this number from measured quantities. (A) An estimate without uncertainty, combining rough estimates of pH and cell volume to obtain a guess for the number of protons. In this example, mean values are chosen to match the means of known measurements, but no associated uncertainty is analyzed. (B) An estimate using CALADIS to explicitly account for uncertainties in the measured quantities and reporting explanatory statistics about the final quantity, using uniform distributions to represent the uncertainty in the variables involved. Other representative distributions are possible and can be analyzed using our approach (see Results). (C) CALADIS also finds that in this example calculation, more of the final uncertainty arises from uncertainty in cell volume than pH; refining volume estimates is slightly preferred as the optimal experimental strategy to lower overall uncertainty. To see this figure in color, go online.

METHODS
Explicitly tracking uncertainty in cell biological calculations

We propose an approach to biological rule-of-thumb calculations involving uncertain quantities that does not solely rely on point estimates of quantities of interest. Instead, our approach involves treating every uncertain quantity in a rule-of-thumb calculation as a probability distribution describing this uncertainty. The following iterative process is then performed: in each iteration, a sampled value is taken from each distribution of interest in the calculation. The value of the complete calculation is computed given this set of samples. This process is iterated many times to build up a distribution of values describing the output of the calculation. This output distribution then provides an interpretable and statistically rigorous answer to the rule-of-thumb question. We present this approach as a complement to, and not a substitute for, the valuable process of Fermi estimation, and stress again the value of “just having a go with the numbers”.

We emphasize that our approach, calculation of quantities using samples from distributions rather than point estimates, can be used to obtain interpretable results in cases where we do not have access to the full set of original measurements. This situation is likely to apply, for example, when using summarized results from previous independent experiments. In this case, our method can be viewed as a generalization of the resampling approaches that could be used if we had full access to the original data, such as bootstrapping or jackknifing (9). In addition to adding statistical power to rule-of-thumb questions in cell biology, this approach can also be used to facilitate efficient design of experiments to reduce uncertainty in a given quantity. In the picture of calculations performed using probability distributions, this goal can be accomplished using a simple variant of a sensitivity analysis approach. Consider artificially decreasing the variance of each distribution in a calculation one-by-one. Decreasing the variance of each individual distribution will lead to a decrease in the overall variance of the output distribution, and the magnitudes of these induced overall decreases can be recorded. The quantity with the most power to decrease overall variance in the calculation output can then be identified, and its value further refined through experiment. Conceptually, this approach resembles performing a sensitivity analysis on the variance of the solution distribution with respect to the variances of individual input distributions.

An important point to consider when attempting to quantify uncertainty in scientific calculations is the source and meaning of the word “uncertainty”. A degree of measurement error may be associated with an experimental protocol, causing uncertainty in the resulting value due to imprecision. Alternatively, a given physical or biological quantity may exhibit genuine variability independent of the measurement process, in that its value fluctuates or changes with time and/or other controlling factors. The degree to which calculations involving uncertain quantities are interpretable is contingent on the types of uncertainty involved (see Discussion).

CALADIS: a probabilistic calculator for biology

We introduce a web-based calculator called CALADIS (from “calculate a distribution”), available for free use (and free source code download) at www.caladis.org. CALADIS, in addition to computing with constant quantities and standard mathematical operators and functions, naturally incorporates probability distributions as fundamental calculation elements, yielding as its output a probability distribution over the final answer. As described above, this probabilistic calculation approach allows uncertainties to be tracked throughout a calculation, providing a wealth of output data and allowing a complete view of the statistical details of the output of a probabilistic calculation (Fig. 1B) and further information about the sources of uncertainty (Fig. 1 C; see later). We underline that our web tool requires no knowledge of computer programming and no access to mathematical software tools, and, in addition to
functioning on desktop and laptop browsers, is compatible with a range of hand-held devices. Our aim in designing this tool is to facilitate fast and easy calculations involving uncertain biological quantities for users including those who lack the background or software to produce their own machinery for performing such calculations. The ability of our site to function on mobile devices makes it a plausible substitute for the well-known napkin over coffee or a conference dinner, facilitating informal but rigorous rough estimates of quantities as new ideas emerge.

CALADIS presents the user with a field (Fig. 2 A) to input calculation expressions, which may involve probability distributions identified with a prepended # symbol, where # functions as a sigil denoting a distribution, e.g.,

\[ \frac{4}{3} \pi \# \text{cellRadiusDist}^3. \]

For every probability distribution found in the input expression, CALADIS prompts the user to choose a distribution type, and appropriate parameters to describe that distribution (for example, perhaps specifying that \#cellRadiusDist is a uniform distribution between 1 and 1.5 \(\mu\)m), or, in the case of BioNumbers (see below), automatically populates the distribution details with the appropriate parameters (Fig. 2 B). Users may also use a built-in browser to input distributions corresponding to recorded quantities (Fig. 2 C; see BioNumbers below). The user may then click “Calculate”, whereupon CALADIS computes a probability distribution describing the final answer using the above approach, sampling many times from each distribution the user entered to build up a set of samples from the resultant distribution, which is then displayed graphically (Fig. 2 D). This interface includes a tool to estimate the probability mass between two given values, user-controlled display of the probability of lying in each bin, summary statistics of the distribution (Fig. 2 E), results from the optional standard deviation (SD) analysis (Fig. 2 F), and a URL that serves as a permanent link to that calculation. This collection of output statistics and graphics allows a complete overview of the probabilistic result of the user’s calculation.

CALADIS also facilitates the aforementioned efficient design of experimental strategies, through consideration of the contributions of different quantities to the overall uncertainty in a calculation. The user has the option of performing a standard deviation analysis for common types of input distribution in the web interface. In this analysis, the SD of each input distribution of this type is artificially reduced by 10%, and the resulting effect on the SD and interquartile range of the resultant distribution is recorded (Fig. 2 F). Intuition about the input variable with the most power to refine the overall output estimate can then be gained straightforwardly.

BioNumbers

We have embedded the data provided by the BioNumbers repository (7) within CALADIS. BioNumbers contains a huge range of biological measurements, spanning scales from microscopic chemical reaction rates and cellular concentrations to ecosystem- and planetwide statistics of

![FIGURE 2 Elements of CALADIS interface. (A) The expression input box: a user enters a calculation here, providing any required information about distributions (for example, perhaps specifying that a certain distribution is uniform between 0 and 1, or normal with mean 1 and SD 0.1). (B) Each probability distribution in the input expression must then be characterized, either through the user’s entry of appropriate parameters, or (as depicted) through the automatic recognition of a BioNumber. (C) The BioNumbers Browser allows the identification, selection, and inclusion of values from the BioNumbers database. (D) The resultant distribution for the calculation is then displayed, along with summary statistics of the distribution (E) and (optionally) SD analysis (F) assessing the sensitivity of overall variance with respect to the variance of individual elements. This illustration involves, as an example calculation, the proton number calculation discussed in the Results. To see this figure in color, go online.](https://doi.org/10.1016/j.bpj.2014.10.041)
biological populations. Our link to the database allows us to perform powerful rule-of-thumb biophysical and cell-biological calculations with BioNumbers (5) while tracking uncertainties to estimate the ranges of the final answer.

Within our web tool, the BioNumbers database is parsed to obtain, for each BioNumber, a corresponding probability distribution, units, and a URL to the source data. Probability distributions are assigned based on the format of the source data and according to a user-defined protocol (see the Supporting Material). The units of each value are automatically obtained from the database. Users may then use a variety of approaches to identify and select BioNumbers for use in a probabilistic calculation, and the corresponding probability distributions are automatically included as calculation elements (see the Supporting Material).

RESULTS

Problems with reasoning with mean values in nonlinear contexts

We first illustrate how reasoning using only mean estimates may lead to incorrect results in calculations. Consider two measured quantities \( X \) and \( Y \), perhaps corresponding to the abundance of two different types of entity in a population. We are interested in the proportion of \( X \) in the population

\[
P = P(X + Y).
\]

Say we have the information that the measured quantities follow log-normal distributions, with \( X \) having mean \( m_X = 0.1 \) and SD (of the log-normal distribution itself, as opposed to the underlying normal distribution) \( s_X = 0.1 \), and \( Y \) having mean \( m_Y = 0.9 \) and SD \( s_Y = 0.9 \). In this artificial example, estimating the expected proportion of \( X \) in the population from the means alone would give

\[
\hat{P} = \frac{m_X}{m_X + m_Y} = 0.1.
\]

However, accurately tracking uncertainty in this calculation produces the counterintuitive result that \( E(P) \approx 0.144 \), rather more than the population proportion estimated from mean values (Fig. 3 A).

This illustration contrasts with the cases where a calculation is straightforwardly additively or multiplicatively separable. In such cases, the fact that functions \( f(X) \) and \( g(X) \) of independent random variables \( X \) and \( Y \) are themselves independent leads to the results \( E(f(X)g(Y)) = E(f(X))E(g(Y)) \) and \( E(f(X) + g(Y)) = E(f(X)) + E(g(Y)) \), implying that calculations based on the individual means of \( X \) and \( Y \) will accurately estimate the overall mean. The error in the mean-based estimate \( \hat{P} \) in our example arises from the structure of the expression used to calculate the population proportion: the fraction cannot be separated into independent functions of \( X \) and \( Y \). Generally in such inseparable cases, calculations based solely on mean values may not provide correct estimators. In such cases, explicitly tracking uncertainty not only provides a powerful characterization of the uncertainty in the final answer but also guarantees that such errors in the mean outcome are not made.

Next, we give two example calculations from the BioNumber of the Month website (3) to illustrate the process of explicitly tracking uncertainties in cell biological calculations with BioNumbers. The details of the BioNumber distributions used are shown in the Supporting Material.

The number of hydrogen ions in a cell

Given measurement of the pH and volume \( V \) of a system, the number of hydrogen ions in the system can be deduced as

\[
n = 10^{-pH}N_AV, \text{ where } N_A \approx 6 \times 10^{-23} \text{ is Avogadro’s number}.
\]

In the December 2011 entry of Milo (3), measurements of pH and cell volume are used to estimate that an \textit{Escherichia coli} cell contains \(~60\) hydrogen ions. Using CALADIS’ BioNumbers browser to search for “cell volume” and “cytoplasm pH” identifies BioNumbers 100003 (\textit{E. coli} cell volume) and 106518 (\textit{E. coli} pH). These values appear in BioNumbers as (100003) 0.1–3.5 \( \mu \text{m}^3 \), interpreted as \( U(0.1, 3.5) \mu \text{m}^3 \); and (106518) 7.2–7.8, interpreted as \( U(7.2, 7.8) \). It is possible to interpret these results in terms of different probability distributions—a facility supported by CALADIS (see the Supporting Material). For example, the quantity 0.1–3.5 \( \mu \text{m}^3 \) could be interpreted as a log-normal distribution with 0.1 \( \mu \text{m}^3 \) and 3.5 \( \mu \text{m}^3 \) as \( \pm 1\sigma \) points of the distribution. However, in this specific example, we use a uniform distribution, because the corresponding log-normal distribution exhibits extremely high variance with a range over more than an order of magnitude, which does not intuitively match the expected distribution of cell sizes in a population. Additionally, analytic results for the distribution of exponentially growing, dividing cells suggest a quadratic distribution that bears a stronger resemblance to the uniform than the log-normal picture (10). The ability to explore these different interpretations, and quantitatively debate the properties of each, are valuable scientific processes which our approach facilitates.

We can automatically access these BioNumbers and their associated uncertainties in CALADIS, then calculate the above equation while tracking uncertainties (this calculation forms the example used illustratively in Fig. 1 B). We find that the resultant distribution (see Fig. 3 B) easily spans an order of magnitude, with 14% of the density \(< 10\) protons and 3% \( > 100 \) protons (statistics straightforwardly found using CALADIS’ interface). Use of SD analysis suggests that more of this uncertainty originates from the spread of cell volumes. We now have a mean estimate at \(~37\) protons and a full characterization of the uncertainty associated with this answer, allowing a quantified degree of confidence to be associated with our reasoning.

Diffusion times in cells

In the March 2010 entry of Milo (3), the characteristic time-scales for diffusion through cells of various sizes are
explored, using the expression $t = x^2/6D$, where $x$ is the
length scale of diffusion and $D$ is the diffusion constant of
the species of interest. Milo (3) uses a rough estimate
of the diffusion constant for GFP in *E. coli* and order-of-
magnitude reasoning to obtain an estimate of 10 ms to tra-
verse a root mean square distance of 1 μm.

Using CALADIS’ BioNumbers browser to search for
“diffusion rate” identifies BioNumber 100193 (diffusion
rate in *E. coli*), recorded as $7.7 \pm 2.5 \, \mu \text{m}^2 \, \text{s}^{-1}$ and inter-
preted as $N(7.7, 2.5) \, \text{m}^2 \, \text{s}^{-1}$. We follow the calculation in
Milo (3) by including this BioNumber in the above equation,
using $x = 1 \, \mu \text{m}$, and performing the probabilistic calculation
of $t$ in CALADIS, tracking uncertainties. We observe that the
resultant distribution (see Fig. 3 C) is highly skewed, with an
apparent coefficient of variation (the ratio of the SD to the
mean, illustrating the spread of the distribution) of $\approx 2.4$.
This example, where a probability distribution appears in
the denominator of an expression for a quantity of interest,
illustrates how the resultant uncertainty can behave unintui-
tively when variables are combined even in relatively simple
ways. Calculation of a resultant distribution provides a more
robust method in these circumstances than traditional prop-
agation-of-uncertainty approaches, and construction of a
full probability distribution for the output of a calculation
allows interpretation of details like skewness that are missed
by a simple estimate of the SD alone.

**DISCUSSION**

We have described an approach for performing rule-of-
thumb calculations in biophysics and cell biology while
incorporating the considerable uncertainty often involved
in such biological contexts. This approach, which does not
rely solely on point estimates of relevant quantities, allows
the treatment and interpretation of the uncertainty involved
in such calculations, increasing their trustworthiness and
their power to assist intuitive reasoning. In addition, it
may be used to optimize experimental design, by helping
to identify measurements with the greatest power to refine
knowledge of the overall quantity of interest.

To facilitate the straightforward use of this approach, both
at a computer and on mobile devices, we have introduced
CALADIS, an online tool for performing calculations
involving probability distributions, available for free use
and with its source code open and available to download.
CALADIS has a particular link to rule-of-thumb calcula-
tions with BioNumbers in cell biology, and we have illus-
trated its use in deriving distributions of quantities of
biophysical and cell biological interest. In employing these
calculations in a scientific context, it is important to note
that tracking the uncertainties in calculations is only useful
if the underlying model is appropriately trusted: hygienic
treatment of errors is a separate consideration from picking
the right model for the world. It is unlikely that the model
probability distributions employed in our approach (and
many other analyses) represent the perfect description of a
quantity arising in the real world; however, we hope that
our approach, with the broad range of distributions sup-
ported by CALADIS, provides a means of reasonably esti-
mating a wide range of real quantities. As we highlight
above, the discussion of appropriate models for uncertainty,
and their quick quantitative comparison, is a scientifically
beneficial feature facilitated by our approach.

Back-of-the-envelope calculations (though used
throughout history) have become increasingly popular
recently as tools for developing quantitative reasoning and

**FIGURE 3** Point estimates and biological distri-
butions. (A) The distribution resulting from the
illustrative $X/(X + Y)$ calculation in the text. The
value obtained by considering mean estimates
alone differs from the mean of the true distribution,
which is heavily skewed, highlighting the impor-
tance of explicitly tracking uncertainty. (B and C)
Estimates, using data from biological experiments
via the BioNumbers database, and tracking uncer-
tainties for (B) number of protons in an *E. coli*
cell and (C) time for GFP to diffuse 1 μm in
*E. coli*. All distributions are direct outputs from
CALADIS. To see this figure in color, go online.
intuition (1,4,5). Despite this increase in popularity, their use is not yet as prevalent in biology as in the physical sciences. We hope that this tool provides support for, and may increase trust in, the use of back-of-the-envelope calculations in quantitative cell biology (and across the biosciences) by exposing the role of uncertainties. We have shown that in some cases (for example, in calculating proportions), failing to track uncertainties can lead to rough guesses that do not represent the full truth of the calculation.

In our work with biological calculations, we have found that CALADIS plays a useful role in quality control for rule-of-thumb reasoning: after having made an approximate estimate on a real napkin, it is helpful to check whether the biological question at hand remains adequately answered if variability/imprecision is appropriately accommodated. We further note the reverse possibility: rather than serving as a sanity check for our envelope calculations, CALADIS can help create optimism about our estimates. For example, in settings where the uncertainty of some calculation elements is known to be very substantial, it might be the case that the final distribution of the estimated quantity is, in fact, sufficiently constrained for scientific advance. As discussed, an uncertainty appended to an estimated quantity needs to be treated with care (since it can depend on distribution choice) but it can still serve as a partial certificate for the relevance of the estimate. We suggest that researchers may present links to their calculations within CALADIS, so that readers are then free to use their prior beliefs to modify the component distributions (if, for example, a reader is less confident about a variable than the author) to see whether the conclusions are still robust.

As mentioned previously, the interpretation of calculations tracking uncertainty is contingent on the source of the uncertainty in the elements of the calculation, which may arise from imprecision (for example, measurement errors associated with an experimental protocol) or variability (the natural fluctuations intrinsic to a system of interest). Care must be taken in the interpretation of the resultant distribution depending on the sources of uncertainty in the calculation. For example, consider a quantity $X$ which is subject to natural variability, stationary but fluctuating with time, and which has been characterized by a distribution involving a finite number $N$ of measurements of $X$ at different times. If we are interested in the behavior of $X$ over an infinitesimally small time window, it makes sense to draw from this distribution of $X$, since this distribution represents plausible states of the system. If we are interested in the time-averaged behavior of $X$, we may instead consider the distribution of $\bar{E}(X)$, an estimate of the mean of $X$. $E(X)$ is a single number about which we are uncertain: the distribution of $\bar{E}(X)$ derived from our measurements will have a finite width (the standard error on the mean, dependent on $N$), corresponding to imprecision rather than natural variability. Mixing uncertainties due to imprecision with those due to variability may lead to results that are not trivial to interpret. We underline the importance of transparency in the meaning of a probabilistic calculation to avoid misinterpretation—in the above example, it should be explicitly stated whether a calculation involves (variable) single instances of a measurement ($X$) or (imprecise) time-averaged behavior ($\bar{E}(X)$).

The process of sampling from distributions describing individual quantities, performing a calculation using these samples, and building a final distribution is akin to several methodologies of use in Bayesian statistics (11). The difference between our approach and Bayesian sampling approaches is that after establishing our distributions we condition on no further data, instead assuming that the individual distributions (which could be pictured as priors) already contain all information on the likelihood of individual values. In this sense, the Bayesian interpretation of our approach is not as a method for extracting posteriors from priors given data, but is instead a method for performing calculations with priors without new data, thus constructing new prior distributions over more complicated quantities.

SUPPORTING MATERIAL

Additional supplemental information are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(14)01124-2.

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