The volumes and transcript counts of single cells reveal concentration homeostasis and capture biological noise

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ABSTRACT Transcriptional stochasticity can be measured by counting the number of mRNA molecules per cell. Cell-to-cell variability is best captured in terms of concentration rather than molecule counts, because reaction rates depend on concentrations. We combined single-molecule mRNA counting with single-cell volume measurements to quantify the statistics of both transcript numbers and concentrations in human cells. We compared three cell clones that differ only in the genomic integration site of an identical constitutively expressed reporter gene. The transcript number per cell varied proportionally with cell volume in all three clones, indicating concentration homeostasis. We found that the cell-to-cell variability in the mRNA concentration is almost exclusively due to cell-to-cell variation in gene expression activity, whereas the cell-to-cell variation in mRNA number is larger, due to a significant contribution of cell volume variability. We concluded that the precise relationship between transcript number and cell volume sets the biological stochasticity of living cells. This study highlights the importance of the quantitative measurement of transcript concentrations in studies of cell-to-cell variability in biology.

INTRODUCTION

Spontaneous fluctuations in the activities of molecular processes cause heterogeneity in the molecular composition of isogenic cells (Ozbudak et al., 2002; Elowitz et al., 2002; Sigal et al., 2006). Cell-to-cell variability, often referred to as “noise,” has been observed in mRNA (Golding et al., 2005; Raj et al., 2006; Zenklusen et al., 2008) and protein levels (Ozbudak et al., 2002; Elowitz et al., 2002; Sigal et al., 2006; Yu et al., 2006), in the timing of molecular processes (Amir et al., 2007; Di Talia et al., 2007), and in cellular growth rates (Boulineau et al., 2013). The causes of molecular noise involve molecules occurring at low numbers per cell, such as transcription factors or mRNAs, that tend to show large, spontaneous deviations relative to their mean number within the cell population (Paulsson, 2004). These deviations (fluctuations) can be caused by cell division (Huh and Paulsson, 2011), transcription bursting (Suter et al., 2011), or transient imbalances between molecular synthesis and degradation rates that occur spontaneously through thermal noise (“intrinsic” noise) or due to fluctuations in the number of regulators (“extrinsic” noise) (Thattai and van Oudenaarden, 2001; Elowitz et al., 2002; Swain et al., 2002; Paulsson, 2004). Extrinsic noise indicates that fluctuations can propagate through the entire molecular network of a cell (Pedraza and Paulsson, 2008). As a result, the molecular composition of cells can be highly variable and cause heterogeneity in differentiation decisions (Wernet et al., 2006), stress response magnitude (Veening et al., 2005), and the survival prospects of cells after drug exposure (Spencer et al., 2009). Two highly informative reviews covering gene expression noise and its consequences were written by Kaern et al. (2005) and Raj and van Oudenaarden (2008).

Early studies on stochastic gene expression relied on fluorescent proteins to assess protein noise by either taking snapshots (Elowitz et al., 2002; Ozbudak et al., 2002) or by using real-time fluorescence imaging (Rosenfeld et al., 2005; Sigal et al., 2006). More recently, single-molecule mRNA counting has been introduced as a method for absolute quantification of mRNA number (Golding et al., 2005;
using confocal microscopy. We studied three human cell line variants (HEK293; described in Gierman et al., 2007). Each clone has a single random insertion of the same green fluorescent protein (GFP).

**RESULTS**

Single-cell transcript data indicate gene location–dependent mRNA expression

We analyzed three clones derived from the same human cell line (HEK293; described in Gierman et al., 2007). Each clone has a single random insertion of the same green fluorescent protein (GFP).
reporter gene controlled by a constitutive phosphoglucomutase (PGK) promoter (Gierman et al., 2007). We determined the statistics of the GFP mRNA levels in single cells with smFISH (Figure 1A). The probe set contained 35 probes of 17–18 nucleotides coupled with a fluorescent label (Supplemental Information 7). Images of single cells were obtained with confocal microscopy on smFISH-treated cells counterstained with 4′,6-diamidino-2-phenylindole (DAPI). Individual cells were recorded as 52 z-stack images (300 nm/slice; Figure 1D and Supplemental Figure S2). Laminar staining confirmed that the DAPI staining correctly identifies the nuclear envelope (Supplemental Figure S1). On the basis of the colocalization with the DAPI signal, we assigned each mRNA molecule to be either nuclear or cytoplasmic. An overview of the transcript counting statistics is given in Figure 1B and Supplemental Figure S14, and the mRNA distributions are shown in Figure 1, E–G, and Supplemental Figure S13. The measured volume distributions for the whole cell (Supplemental Figure S5), as well as the cytoplasmic and nuclear volume distributions (Figure 2C), are positively skewed, which means there are, relative to the mean, more small than large cells. Similarly shaped distributions have previously been reported for stationary, growing cell populations (Tzur et al., 2009). These positively skewed distributions are due to the formation of two (smaller) daughter cells from each (large) mother cell. The cell volume distributions can be well approximated by the functional cell volume distributions derived from, balanced, exponential growth of the cells (Supplemental Information 3.3 and Supplemental Figure S7).

The obtained volume measurements are summarized in Figure 2B. The mean cell volume (V) for clone I is 1800 µm³, with a larger nucleus (979 µm³), on average, than cytoplasm (822 µm³). For the volume distribution of clone I, 70% of the cells have a volume deviation maximally one SD of the mean. The correlation coefficients between the nuclear and the cytoplasmic volumes indicate a weak but significant positive correlation, indicating growth of both the nuclear and cytoplasmic volume during cell maturation.

Because the clones are isogenic, except for the integration site of the construct, the measured volume statistics would be expected to be similar. Figure 2B and Supplemental Figure S5 confirm this expectation for the three volume distributions measured.

**mRNA concentration statistics of single cells indicate mRNA concentration homeostasis**

Next we combined the mRNA number and volume data of each cell to determine the statistics of cellular, cytoplasmic, and nuclear mRNA concentrations (Figure 3A). Figure 3B shows that the mean mRNA concentration differs among the three clones, indicating the dependence of expression levels on the gene location (Supplemental Information 2). Scaling of the mean and SD are independent. We observe higher mRNA concentrations in the cytoplasm than in the nucleus. Similar conclusions can be drawn from the mRNA numbers. The comparison of the mRNA concentration and mRNA number in terms of the coefficient of variation indicates that the concentration displays smaller cell-to-cell variability across all clones. Thus mRNA number noise overestimates the biological mRNA noise of the cells. The concentration variability is smaller because of the positive correlation of the cell volume with mRNA number per cell. This is indicated in Figure 1C, showing a linear dependency of the mean mRNA number (at a specific volume) with the mRNA number per cell. The dependency of the mean mRNA number on volume is proportional, such that a doubling in cell volume is accompanied by a doubling in the mRNA number. The cellular mRNA concentration conditional on the cell volume is therefore constant, indicating homeostasis of the mRNA concentration (Figure 3C). The same proportionality of the mRNA numbers and homeostasis of the mRNA concentrations is observed in the nucleus (Supplemental Figure S11) and cytoplasm (Supplemental Figure S12) of the cells.

**Volume scaling of the mRNA concentration statistics explains the concentration variability**

To address the origins of the differences between the mRNA concentration (c) and the mRNA number (m) noise, we apply the law of

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The small deviations are volume-dependent noise, with \( \alpha \) decreasing with volume (Figure 4B and Supplemental Information 3.3), which provides a lower bound for the volume-induced noise contribution to the mRNA number noise. This indicates that mRNA number noise is entirely determined by gene expression noise.

The experimental data indicate that the gene expression contributions to mRNA concentration and mRNA number noise are similar in absolute values (Figure 4B). Because volume-derived noise in mRNA concentration is close to zero, the mRNA concentration noise is approximately equal to the gene expression–derived noise in mRNA numbers. It turns out that this is a direct consequence of the observed mRNA concentration homeostasis and, in addition, the scaling of the conditional mRNA number variance, \( \text{var}(x|V) \), with cell volume. At mRNA concentration homeostasis, the exact relationship between the gene expression–induced mRNA number and concentration noise is given by (Supplemental Information 3.5)

\[
\text{var}(c|V) = \frac{(\text{var}(m|V))}{(1/2)} \text{var}(m|V) + \alpha^2 \left[ \frac{\text{var}(m|V)}{\text{var}(V)} \right]
\]

with \( \alpha^2 \) denoting the covariance between \( x \) and \( y \). Because we observe that \( \text{var}(m|V) \) increases with volume (Figure 4B and Supplemental Figure S9), the covariance will be negative. Therefore \( \gamma_{xV}^2 \) is an upper bound for the relative deviation between the conditional noise in mRNA concentration and mRNA number in cases of concentration homeostasis. This upper bound is reached when the covariance equals zero. From the volume probability distribution,
volume-dependent noise term, which is zero for concentrations but equals values between 0.04 and 0.06 for mRNA numbers. Under conditions of mRNA concentration homeostasis, this term is expected to be independent of the average expression level. The relative difference between mRNA number and concentration noise then depends on the magnitude of the volume-independent noise contribution. For the three clones we investigated, this amounted to differences of 36, 33, and 45% between mRNA concentration and mRNA number noise for I, II, and III, respectively. Thus biologically relevant mRNA concentration noise differs greatly from mRNA number noise, indicating the importance of the combined measurement of mRNA numbers and volumes of single cells.

DISCUSSION

Single-molecule RNA FISH is a powerful method for assessing cell-to-cell variation in gene expression. It does not require genetic engineering, and it gives the exact number of mRNA molecules per cell. In this study, we combined smFISH with cell volume measurements to obtain insight into the cell-to-cell variation of the mRNA number variance conditional on volume, that is, from $\beta = \gamma$ varm $VV (c | V)$. The experimental data (Supplemental Figure S9) indicate that this scaling is maximally quadratic with volume. The volume growth model predicts that, in cases of linear dependence, $\text{var}(c | V)$ is 4% higher than $\text{var}(c | V)$; in cases of quadratic dependence, it is 4% lower; and only when $\text{var}(m | V) = \beta V^2$ is the maximal deviation of 12% achieved (Supplemental Information 4.6). As can be seen from Figure 4B, the relative difference between the two conditional noise terms indeed is close to the ± 4% region, as predicted by theory in combination with the experimentally observed volume scaling.

As a result of these relations, we can conclude that, for our data, the difference between mRNA concentration noise $\text{var}(c)$ and mRNA number noise $\text{var}(m)$ is dominated by the contribution of the volume-dependent noise term, which is zero for concentrations but equals values between 0.04 and 0.06 for mRNA numbers. Under conditions of mRNA concentration homeostasis, this term is expected to be independent of the average expression level. The relative difference between mRNA number and concentration noise then depends on the magnitude of the volume-independent noise contribution. For the three clones we investigated, this amounted to differences of 36, 33, and 45% between mRNA concentration and mRNA number noise for I, II, and III, respectively. Thus biologically relevant mRNA concentration noise differs greatly from mRNA number noise, indicating the importance of the combined measurement of mRNA numbers and volumes of single cells.

FIGURE 3: Statistics of single-cell mRNA concentrations. (A) The previously obtained mRNA number (Figure 1) and volume data (Figure 2) were used to determine the concentration of mRNA in single cells ($c$), in their nuclei ($c_N$), and in their cytoplasm ($c_C$). (B) Statistics of the different mRNA concentrations for the three different clones (color coded). Notation: $\mu = \text{mean}; \sigma = \text{SD}; \text{cv} = \text{coefficient of variation}; \Delta^v = \text{the fraction of samples between } \mu - \sigma \text{ and } \mu + \sigma; \rho = \text{correlation between } m_t \text{ and } m_v; \text{ and } * p < 0.001 (H_0: \rho = 0).$ The 95% confidence intervals of the statistics are given in Supplemental Figure S15. (C) For a specific cell volume ($V$), the mean mRNA concentration is calculated. A least-squares linear fit is shown for all three clones, indicating mRNA concentration homeostasis. The conditional variances of the data set are shown in Supplemental Figure S10. (D–F) Scatter plots of ($c_n$) and ($c_c$) for the three different clones. Marginal histograms show the distribution of ($c_c$) (top) and ($c_n$) (right). The given concentration (number per cubic micrometer) can be converted to picomoles (pM) by multiplying with a conversion factor of 1660. The sample size is given by $n$. The bin size for the marginal histograms is 0.001 pm$^3$. As V$^2$ (1/7) is estimated to be 1.12 (Supplemental Information 3.6). The deviation between $\frac{\text{var}(c)}{\text{var}(m)}$ and $\frac{\text{var}(m)}{\text{var}(c)}$ requires the calculation of the covariance from the volume-scaling relation of the mRNA number variance conditional on volume, that is, from $\text{var}(m | V) = \beta V^2$. The experimental data (Supplemental Figure S9) indicate that this scaling is maximally quadratic with volume. The volume growth model predicts that, in cases of linear dependence, $\text{var}(c | V)$ is 4% higher than $\text{var}(c | V)$; in cases of quadratic dependence, it is 4% lower; and only when $\text{var}(m | V) = \beta V^2$ is the maximal deviation of 12% achieved (Supplemental Information 4.6). As can be seen from Figure 4B, the relative difference between the two conditional noise terms indeed is close to the ± 4% region, as predicted by theory in combination with the experimentally observed volume scaling.

As a result of these relations, we can conclude that, for our data, the difference between mRNA concentration noise $\frac{\text{var}(c)}{\text{var}(m)}$ and mRNA number noise $\frac{\text{var}(m)}{\text{var}(c)}$ is dominated by the contribution of the...
mRNA concentration per cell. We studied three clones that only differed in the location of an identical reporter construct controlled by a constitutive PGK promoter. The differences in the mRNA statistics of these clones indicate gene location dependency, which presumably results from the different genomic contexts at the integration site. We found that mRNA number noise overestimates cell-to-cell variation. Noise in the mRNA concentration circumvents this problem by taking into account the correlation of mRNA number with cell volume. One candidate source of this correlation is cell growth.

We found that the mean mRNA number conditional on volume, $\langle m(V) \rangle$, scaled linearly with volume, that is, $\langle m(V) \rangle = cV$, which indicates a constant mRNA concentration ($c$) as a function of cell volume (homeostasis). This we interpret as a constant mRNA concentration while the cell volume grows. In addition, we found that the mRNA number variance conditional on the volume, $\text{var}(m(V))$, displayed a stronger than linear scaling with volume, that is, $\text{var}(m(V)) = \beta V^\gamma$ with $1 \leq \gamma \leq 2$. The latter scaling explained the difference between the volume-independent mRNA concentration $\text{var}(m(V))$ with volume hints at another not yet understood detail of constitutive gene expression. We observed the volume scaling of $\text{var}(m(V))$ and $\text{var}(m(V))$ with the same construct expressed from different genomic locations, and therefore the volume scaling is not likely to be a genome location–dependent phenomenon. The observed volume scaling of mRNA numbers is not likely to be a property of our reporter construct, as several other studies reported similar effects Rosenfeld et al., 2005; Sigal et al., 2006; Cohen et al., 2009; Cookson et al., 2010). Cookson et al. (2010) found in yeast a peak dependency of a GFP expressed from a constitutive promoter as function of cell volume, which is indicative of constant synthesis and an accelerating growth of cell volume as a function of the cell cycle. Similar data were reported for several human proteins (Sigal et al., 2006).

The three clones investigated show 36, 33, and 45% differences between mRNA concentration and mRNA number noise for I, II, and III, respectively. We conclude that functional mRNA noise differs greatly from mRNA number noise, indicating the importance of...
measuring mRNA noise in terms of concentrations. For inducible or cell cycle–dependent promoters, the assessment of mRNA concentration noise is even more relevant. For such systems, a non-linear relation between volume growth and mRNA synthesis is expected. This would introduce much larger differences between the mRNA number and mRNA concentration noise than reported in this study, which is limited to a constitutive promoter.

Real-time monitoring of mRNA number (e.g., by using MS2 labeling; Fusco et al., 2003) and volume growth of single cells for a set of (classical) constitutive or inducible promoters could provide more information about the origins and effects of the volume-scaling relations of the mean and variance of the transcript numbers. A downside is that such studies would require the tracking of several hundred cell divisions to obtain robust statistics on the volume dependencies of mRNA number statistics. To attain robust statistics in our experiments, we used a confocal microscopy setup and had to study roughly 1000 cells per cell clone.

Single-molecule methods are a great addition to single-cell biology. They are quantitative and exact, as they give insight into the actual molecular composition of individual cells. In addition, they directly relate to stochastic theory and model predictions, giving deep insight into how cells exploit the inherent stochasticity of molecular processes to diversify isogenic populations. In this study, we have shown that, in order to profit fully from the exactness of single-molecule methods in cell biology, these methods should ideally be combined with single-cell volume measurements. This finding is highly relevant, because concentration noise captures biological noise.

**MATERIALS AND METHODS**

**Cell clones and cell culture**

Experiments were performed on human embryonic kidney cells (HEK293) with a single integration of a phosphoglycerate kinase (PGK)-driven GFP gene construct obtained from Gierman et al. (2007). We analyzed three different clones with the integration at different genomic locations: clone I (HG19:chr1:225684028, within the ENAH gene), clone II (HG19:chr1:150379508, within the RPRD2 gene), and clone III (HG19:chr1:150664232, within the GOLPH3L gene). The cells were cultured in DMEM (31965023; Life Technologies, Carlsbad, CA) supplemented with 10% (vol/vol) fetal calf serum (16140; Life Technologies) and 100 U/ml penicillin–streptomycin (15140; Life Technologies). Incubation was at 37°C in a humidified 5% CO₂ atmosphere. Before any experiments, the cells were grown for at least 2 wk after thawing to achieve steady-state cell growth and steady-state expression statistics for the integrated construct.

**Single-molecule RNA FISH**

Samples were treated according to the Protocol for Adherent Mammalian Cell Lines for the Custom Stellaris FISH probes. Cells were cultured for 3 h in Lab-Tek chambered coverglasses (155380; Lab-Tek, Waltham, MA) before fixation. ETOH permeabilization was done overnight at 4°C. For hybridization, we used a 125-nM probe in the hybridization buffer and overnight incubation at 37°C. Imaging was done without using anti-fade. The cells were counterstained with 5 ng/ml DAPI. The sequence of the probe targeting the eGFP insertion can be found in Supplemental Information 7. The DNA probes were coupled to CAL Fluor Red 590 fluorophores by the manufacturer (Biosearch Technologies, Petaluma, CA).

**Image acquisition**

Samples were imaged using a Nikon Ti-E scanning-laser confocal inverted microscope (A1) with a 60x oil objective in combination with Nikon NIS-Elements imaging software. Excitation was by 561.5-nm diode-pumped solid-state and 402.1-nm diode lasers. Detection was via 595-50 nm and 450-50 nm band-pass filters. Optical sections were captured at 0.300-μm intervals and a resolution of 256 by 256 pixels and a zoom factor of 6.8, resulting in a voxel size of 0.0047 μm³ (0.1243 μm × 0.1243 μm × 0.3 μm). Averaging was used four times to reduce photon and camera noise.

**Image analysis**

Image analysis software was adapted from Raj et al. (2008). Images are filtered with a semi–three-dimensional Laplacian of Gaussian filter that removes noise and enhances the signal-to-noise ratio (filter width = 1.5). The number of mRNA spots was found by applying a threshold for which the number of mRNA was least sensitive to changes in this threshold. The threshold was determined by using a window function calculating the average spot numbers over 7 constitutive thresholds divided by the sum of the SD of these spot counts and a constant α (= 10; Itzkovitz et al., 2012). To measure the cell volume, we performed the following operations on all individual z-slices: 1) median filtering (20 × 20 pixels), 2) image thresholding (graythresh, Otsu’s method; Otsu, 1979), 3) fill image regions and holes, and 4) morphological closing with a disk (radius = 4 pixels). The nuclear volume was obtained from the processed images of the DAPI channel (Supplemental Information 1). The cell volume is defined by the presence of signal from either the nucleus (DAPI), the cytoplasm (red channel), or both. The cytoplasmic size is given by the difference between nuclear and cell volume. The obtained pixel size was multiplied by the voxel size of 0.0047 μm³ to provide the cell size in cubic micrometers.

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