Extensive post-transcriptional regulation across human tissues

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Abstract

Transcriptional and post-transcriptional regulation shape tissue-type-specific proteomes, but their relative contributions remain contested. Estimates of the factors determining protein levels in human tissues do not distinguish between (i) the factors determining the variability between the abundances of different proteins and (ii) the factors determining the physiological variability of the same protein across different tissue types. We estimated the factors determining these two orthogonal sources of variability and found that mRNA levels can account for most of the variability between the levels of different proteins but not for the protein variability across tissue-types. After accounting for measurement noise, most of the across-tissue protein variability remains unexplained by mRNA levels, suggesting extensive post-transcriptional regulation. These results reconcile existing estimates in the literature, caution against estimating protein fold-changes from mRNA fold-changes between different cell-types, and highlight the contribution of post-transcriptional regulation in shaping tissue-type-specific proteomes.
Introduction

The relative ease of measuring mRNA levels has facilitated numerous investigations of how cells regulate their gene expression across different pathological and physiological conditions (Sørlie et al, 2001; Slavov and Dawson, 2009; Spellman et al, 1998; Slavov et al, 2011, 2012; Djebali et al, 2012). However, often the relevant biological processes depend on protein levels, and mRNA levels are merely proxies for protein levels (Alberts et al, 2014). If a gene is regulated mostly transcriptionally, its mRNA level is a good proxy for its protein level. Conversely, post-transcriptional regulation can set protein levels independently from mRNA levels, as in the cases of classical regulators of development (Kuersten and Goodwin, 2003), cell division (Hengst and Reed, 1996; Polymenis and Schmidt, 1997) and metabolism (Daran-Lapujade et al, 2007; Slavov et al, 2014a). Thus understanding the relative contributions of transcriptional and post-transcriptional regulation is essential for understanding their trade-offs and the principles of biological regulation, as well as for assessing the feasibility of using mRNA levels as proxies for protein levels.

Previous studies have considered single cell-types and conditions in studying variation in absolute mRNA and protein levels genome-wide, often employing unicellular model organisms or cell cultures (Gygi et al, 1999; Smits et al, 2014; Schwanhäusser et al, 2011; Li et al, 2014; Jovanovic et al, 2015; Csárdi et al, 2015). However, analyzing per-gene variation in relative mRNA and protein expression across different tissue types in a multicellular organism presents a potentially different and critical problem which cannot be properly addressed by examining only genome-scale correlations between mRNA and protein levels (Csárdi et al, 2015). Wilhelm et al (2014) have measured protein levels across human tissues, thus providing a valuable dataset for analyzing the regulatory layers shaping tissue-type-specific proteome. The absolute levels of proteins and mRNAs in this dataset correlate well, highlighting that highly abundant proteins have highly abundant mRNAs. Such correlations between absolute levels of mRNA and protein mix/conflate many sources of variation, including variability between the levels of different proteins, variability within the same protein across different conditions and cell-types, and the variability due to measurement error and technological bias.

However, these different sources of variability have very different biological interpretations and implications. A major source of variability in protein and mRNA data arises from differences
between the levels of mRNAs and proteins corresponding to different genes. That is, the mean levels (averaged across tissue-types) of different proteins and mRNAs vary widely. We refer to this source of variability as *mean-level variability*. This mean-level variability reflects the fact that some proteins, such as ribosomal proteins, are highly abundant across all profiled conditions while other proteins are orders of magnitude less abundant across all profiled conditions. Another principal source of variability in protein levels, intuitively orthogonal to the mean-level variability, is the variability within a protein across different physiological conditions or cell-types. This variability reflects normal physiological regulation across tissues, and thus we refer to it as *across-tissue variability*. The across-tissue variability is usually much smaller in magnitude, but it is frequently the most relevant source of variability for understanding different phenotypes across cells-types and physiological conditions.

Here, we separately quantify the contributions of transcriptional and post-transcriptional regulation to the mean-level variability and to the across-tissue variability across human tissues. Our results suggest that the across-tissue protein variability is poorly explained by transcriptional regulation, while the mean-level variability can be explained well by transcriptional regulation. These results add to previous results in the literature (*Gygi et al.*, 1999; *Schwanhausser et al.*, 2011; *Li et al.*, 2014; *Wilhelm et al.*, 2014; *Jovanovic et al.*, 2015; *Csárdi et al.*, 2015; *Smits et al.*, 2014), and suggest that the post-transcriptional regulation is a major contributor in determining the variability in the levels of a protein across human-tissues. We then suggest a simple and general approach for deconvolving the contributions of transcriptional and post-transcriptional regulation to measured protein levels.

**Results**

**The correlation between absolute mRNA and protein levels conflates distinct sources of variability**

We start by outlining the statistical concepts underpinning the common correlational analysis and depiction (*Gygi et al.*, 1999; *Schwanhausser et al.*, 2011; *Wilhelm et al.*, 2014; *Csárdi et al.*, 2015) of estimated absolute protein and mRNA levels as displayed in Figure 1a. The magnitude of the
correlation between the absolute mRNA and protein levels of different genes and across different tissue-types is used to estimate the level at which the protein levels are regulated (Wilhelm et al., 2014). If the across-tissue variability of a gene is dominated by transcriptional regulation, its protein-to-mRNA ratio in different tissue-types will be a gene-specific constant. Based on this idea, Wilhelm et al. (2014) estimated these protein-to-mRNA ratios. They suggested that the median ratio for each gene can be used to scale its tissue-specific mRNA levels and that this “scaled mRNA” predicts accurately tissue-specific protein levels.

Indeed, scaled mRNA levels explain large fraction of the total variance ($R^2_T = 0.77$, across 6104 measured proteins, Figure 1a) as previously observed (Schwanhäusser et al., 2011; Wilhelm et al., 2014). However, $R^2_T$ quantifies the fraction of the total protein variance explained by mRNA levels between genes and across tissue-types; thus, it conflates the mean-level variability with the across-tissue variability. This conflation is shown graphically in Figure 1b for a subset of 100 genes measured across 12 tissues. The across-tissue variability is captured by the variability within the regression fits and the mean-level variability is captured by the variability between the regression fits.

The aggregation of distinct sources of variation, where different subgroups of the data show different trends, may lead to the effect known as Simpson’s or amalgamation paradox in the statistical literature, which can lead to counter-intuitive results and incorrect conclusions (Blyth, 1972). To illustrate the Simpson’s paradox in this context, we chose a subset of genes for which the scaled mRNA and measured protein levels are negatively correlated across tissues, and the mean-level variability spans the full dynamic range of the data. For this subset of genes, the overall (conflated/amalgamated) correlation is large and positive, despite the fact that all within-gene trends are negative. This counter-intuitive result is possible because the conflated correlation is dominated by the variability with larger dynamical range, in this case the mean-level variability. This conceptual example taken from the Wilhelm et al. (2014) data demonstrates that, $R^2_T$ is not necessarily informative about the across-tissue variability, i.e., the protein variance explained by scaled mRNA within a gene ($R^2_P$). Thus the conflated correlation is not generally informative about the level — transcriptional or post-transcriptional — at which across-tissue variability is regulated. See Supporting Figure S1 and Methods for a simulation that further demonstrates the insensitivity of $R^2_T$ for quantifying across-tissue variability.
However, it is exactly the across-tissue variability that contributes to the biological identity of each tissue type. This across-tissue variability has a dynamic range of about $2 - 10$ fold and is thus dwarfed by the $10^4$ fold dynamic range of abundances across different proteins. To further demonstrate the implications of this vast difference in the dynamic ranges, we generate data from a simple model using the observed between-tissue variability in the protein-to-mRNA ratio. This protein/mRNA ratio has been referred to as a gene’s “translational efficiency” because it reflects, in large part, its translational rate. Since this ratio also reflects other layers of regulation, such as protein degradation (Jovanovic et al., 2015), and noise we will refer to it descriptively as protein-to-RNA (PTR) ratio.

### Across-tissue mRNA variability is a poor predictor of the across-tissue protein variability

Figure 1 illustrates the statistical problems with using the fraction of the total protein variance explained by scaled mRNA levels ($R^2_T$) as an indication about the extent to which mRNA changes contribute to protein changes across tissues (i.e., $R^2_P$). To investigate the significance of this conflation further, we next evaluated the differences between scaling mRNA with the median PTR ratio (as in Figure 1a) and scaling mRNA with the PTR ratio of a specific tissue. That is, instead of using the median PTR ratio, we can use the PTR ratio estimated from one tissue to scale the mRNA level from another tissue. For instance, we correlate the protein levels measured in uterine to the uterine mRNA levels scaled by the prostate PTR ratio, Figure 2a. This correlation is lower compared to the correlation when mRNA is scaled by the median PTR ratio shown in Figure 1a. This reduction underscores that the PTR ratio for a gene varies enough between tissue types to affect even the conflated variability $R^2_T$. Extending this analysis to more pairs of tissues (Figure 2b, c) indicates very similar results; In all cases, the correlation is around 0.5, substantially smaller than the 0.9 correlation observed when mRNA is scaled by the median PTR ratio, Figure 1a.

Despite the very similar correlations between measured protein and scaled mRNA levels for all 3 comparisons in Figure 2a-c, the corresponding correlations between the protein and mRNA fold-changes differ substantially, Figure 2d-f. Unlike the buffering of mRNA variability observed across species and individuals (Khan et al., 2013; Battle et al., 2014), the protein levels generally vary much
more across tissues than the mRNA levels (Supporting Figure S2); thus the protein and the mRNA fold changes in Figure 2d-f are plotted on different scales. The fold change comparisons in Figure 2d-f demonstrate that in fact, the fraction of variance explained in protein fold-changes by mRNA fold-changes is usually small and depends strongly on the compared pair of tissues. For instance, the mRNA fold-changes between the uterus and prostate have essentially no predictive power for protein fold-changes across these tissues. At the same time, other tissues show a moderate fold-change correlation (e.g., prostate vs. kidney and uterus vs. kidney). The fraction of the variance in protein fold-changes that can be explained by mRNA fold-changes varies significantly across the three examples. However, $R^2_T$ remains high and constant in all three examples (Figure 2a-c), because it is dominated by the mean-level variability. This result underscores the general problem of variance conflation in the analysis of measured mRNA and protein levels.

Next, we sought to evaluate whether across-tissue variability of mRNAs can serve as a proxy for the across-tissue variability of proteins. For this analysis, we extend our results on fold-change correlations from Figure 2 to all pairwise combinations of tissue-types. The range of correlations in Figure 3a indicates that for some pairs of tissues, across-tissue variability in mRNA explains a significant fraction of the across-tissue protein variance but for other tissue-type pairs it does not. This result indicates that for most genes, the observed variability across at least some tissues is either due to measurement noise or to post-transcriptional regulation. Before focusing on distinguishing between these two possibilities, we investigated whether the across-tissue variability of some genes is regulated primarily transcriptionally. If so, the protein fold-changes of such genes may be predicted reliably from mRNA fold-changes. To this end, we quantify the error in predicting protein fold-changes from mRNA fold-changes. The cumulative distribution of errors (Figure 3b) indicates that the protein fold-changes for less than 1000 genes can be estimated from mRNA fold-changes with less than 100% error. For over 30% of proteins, estimating protein levels using a single gene-specific PTR ratio results in over 1000% error; see methods.

**Coordinated post-transcriptional regulation of functional gene sets**

The lack of correlation between protein and mRNA fold-changes can reflect large measurement noise rather than post-transcriptional regulation (Li et al, 2014; Franks et al, 2015). The noise contribution to the variability of the PTR ratios of a gene is likely independent from the
of the gene; see Methods. Conversely, genes with similar functions are likely to be regulated similarly and thus have similar tissue-type-specific PTR ratios. Thus, we explored whether the across-tissues variability of the PTR ratios of functionally related genes reflects such tissue-type-specific and biological-function-specific post-transcriptional regulation.

For this analysis, we define the “relative protein to RNA ratio” (rPTR) of a gene in a given tissue to be the PTR ratio in that tissue divided by the median PTR ratio of the gene across the other 11 tissues. We evaluated the significance of rPTR variability for a gene-set in each tissue-type by comparing the corresponding gene-set rPTR distribution to the rPTR distribution for those same genes pooled across the other tissues (Figure 4); we use the KS-test to quantify the statistical significance of differences in the rPTR distributions; see Methods. Our results indicate that the genes from many GO terms (Consortium et al., 2004) have much higher rPTR in some tissues than in others. For example the ribosomal proteins of the small subunit (40S) have high rPTR in kidney but low rPTR in stomach (Figure 4a-b).

Some of these trends can account for fundamental physiological differences between tissue types. For example, the kidney is by far the most metabolically active (energy consuming) tissue among the 12 profiled tissues (Hall, 2010) and it has very high rPTR for many gene sets involved in energy production (Figure 4a). In this case, post-transcriptional regulation very likely plays a functional role in meeting the high energy demands of kidneys. Moreover, the fact that we observe a highly significant (posterior error probability < $10^{-10}$) mode of rPTR (such as increased rPTR for mitochondrial genes and decreased rPTR for focal adhesion in kidney) indicates that at least some of the variability in post-transcriptional regulation across tissue-types reflects regulatory activity rather than measurement noise.

**Quantifying post-transcriptional regulation across human tissues**

The results in Figure 4 demonstrate the some of the across-tissue variability of protein levels is due to post-transcriptional regulation, not noise. To further quantify the fractions of across-tissue protein variability due to transcriptional regulation, post-transcriptional regulation, and noise, we need to take noise into account. Both RNA-seq (Marioni et al., 2008; Consortium et al., 2014) and mass-spectrometry (Schwanhäusser et al., 2011; Peng et al., 2012) have relatively large and systematic error in estimating *absolute* levels of mRNAs and proteins, i.e., the ratios between
different proteins/mRNAs. These errors originate from DNA sequencing GC-biases, and variations in protein digestion and peptide ionization. However, relative quantification of the same gene across tissue-types by both methods is much more accurate since systematic biases are minimized when taking ratios between the intensities/counts of the same peptide/DNA-sequence measured in different tissue types (Ong et al., 2002; Blagoev et al., 2004; Consortium et al., 2014; Jovanovic et al., 2015). It is this relative quantification that is used in estimating across-tissue variability, and thus noise levels are much smaller compared to the noise of absolute quantification.

To quantify the transcriptional and post-transcriptional contributions to across-tissue protein variability, we start by estimating the reliability of the measurements, Figure 5a-d. Reliability is simply defined as the fraction of the observed/empirical variance due to signal. Thus reliability is proportional to the signal strength and decreases with the noise levels. The analysis of across-tissue variability depends on the relative quantification with a gene and across tissues, and thus we estimate the reliability of relative quantification, i.e., fold-changes across tissues.

For both protein and mRNA, we estimate the reliability as the $R^2$ between independent estimates of their fold-changes. In the case of mRNA, the data contain multiple independent measurements for each tissue, and we split these measurements into two sets. The levels of each mRNA were estimated for each set and the estimates correlated, averaging across tissues (Figure 5a). The squares of these correlations provide estimates for the reliability of each mRNA and their median provides a global /representative estimates for the reliability of relative RNA quantification. An alternative approach to estimating this average reliability is from the $R^2$ between independent estimates of the fold-changes across two tissues (Figure 5b), and it provides very similar estimate; see Methods.

The reliability of the relative protein measurements was estimated as the $R^2$ between separate estimates of the relative protein levels. For each protein, Estimate 1 was derived from 50% of the quantified peptides and Estimate 2 from the other 50%. Since most analytical noise (related to protein digestion, chromatographic mobility and peptide ionization) is peptide-specific, such non-overlapping sets of peptides provide mostly independent estimates for the relative protein levels. The correlations between the estimates for each protein (averaging across 6 tissues) are displayed as a distribution in Figure 5c. To make this reliability estimate relevant to the iBAQ values used throughout the paper, we did not limit this estimate only to ratios between the intensities
of corresponding ions; rather, in each tissue, each protein was quantified by the mean intensities of its peptides; see Methods. As an additional reliability estimate, we compared the protein fold-changes between the salivary and the adrenal glands that were estimated from non-overlapping sets of peptides. For this comparison, we used only peptides detected in both tissues and derived the two separate estimates only from corresponding-ion-ratio. Since corresponding-ion-ratios control better for peptide-specific analytical noise, the reliability estimate is higher, Figure 5d; see Methods.

Taking into account the reliability of the measurements, we depict the upper-bound for the fractions of across-tissue protein variability that can be explained by mRNA levels (i.e., transcriptional regulation) in Figure 5e. To account for any uncertainty in the reliability estimates, we depict the fraction of explained variance for a wide range of reliability estimates. At the reliability estimated for this dataset (Figure 5a-d), at most 30 % of the across-tissue variability – the variability of protein levels across-tissue types that is left after accounting for the measurement noise – can be explained by the mRNA levels. The remaining 70 % is most likely due to post-transcriptional regulation. This result underscores the different modes of regulation for the mean-level variability (mostly transcriptional) and for the across-tissue variability (mostly post-transcriptional). Since the exact estimate depends on the noise/reliability levels, we show estimates for higher and lower levels of reliability. Even if measurement error is larger and most measured variance in both mRNA and protein levels is due to noise, not signal, i.e., reliability $< 50 \%$, transcriptional regulation still can explain at most about 50 % of the across-tissue protein variability. Thus even in this extreme case, post-transcriptional regulation is likely a major determinant of across-tissue protein variability, and thus tissue-type specific proteomes.

**Discussion**

Highly abundant proteins have highly abundant mRNAs. This dependence is consistently observed (Jovanovic *et al*, 2015; Csárdi *et al*, 2015; Gygi *et al*, 1999; Smits *et al*, 2014; Schwanhäusser *et al*, 2011) and dominates the explained variance in the estimates of absolute protein levels (Figure 2a, Figure 2a-c, Figure 5c). This underscores the role of transcription for setting the full dynamic range of protein levels. In stark contrast, differences in the proteomes of distinct human tissues are poorly
explained by transcriptional regulation (Figure 2d-f, Figure 5a-b). Rather, the mechanisms shaping the tissue-specific proteomes involve post-transcriptional regulation. This result underscores the role of translational regulation and of protein degradation for mediating physiological functions within the range of protein levels consistent with life.

The estimates of absolute protein levels are affected by technological biases and measurement error (Peng et al., 2012; Franks et al., 2015) which can contribute to overestimating post-transcriptional regulation. These biases can difficult to estimate and influential (Csárdi et al., 2015), potentially leading to underestimates of the variance in protein levels explained by transcription. However, such systematic biases do not affect the relative changes of protein levels and the estimates of across-tissue variability. Indeed, the strong enrichment of rPTR within gene sets (Figure 4) demonstrates a concerted regulation at the post-transcriptional level. It is thus unlikely that bias and measurement error alone explain the weak correlations between tissue-specific differences in mRNA and protein levels (Figure 2).

As with all analysis of empirical data, the results depend on the quality of the data and the estimates of their reliability. This dependence on data quality is particularly strong given that some conclusions rest on the failure of across-tissue mRNA variability to predict across-tissue protein variability. Such inference based on unaccounted for variability is substantially weaker than measuring directly and accounting for all sources of variability. If the reliability of the data are significantly below 50 %, the data would be consistent with mRNA levels accounting for most of the across-tissue variability, as an upper limit estimate for the transcriptional contribution. In that case, however, the signal is dominated by noise, and thus the data cannot accurately quantify the contributions of different regulatory mechanisms. The strong functional enrichment for rPTR (Figure 4) and the error estimates by Wilhelm et al (2014) suggest that the across-tissue variance that is not explained by mRNA is likely due to post-transcriptional regulation, not to signal dwarfed by noise.

For both mRNA and protein, the reliability estimates in Figure 5a-d quantify and account for measurement noise but not for systematic biases in sample collection and handling. The existing data do not allow estimating the contribution of such sample-related biases. Such biases may differ between the protein and the mRNAs data since the two datasets were collected separately (Wilhelm et al., 2014; Fagerberg et al., 2014). Still, some of these dataset-dependent biases are likely to
average out in the estimates of relative protein and mRNA levels that are used for quantifying across-tissue variability. Another limitation of the data is that isoforms of mRNAs and proteins are merged together, i.e., using razor proteins. This latter limitation is common to all approaches quantifying proteins and mRNAs from peptides/short-sequence reads. It stems from the limitation of existing approaches to their to infer isoform and quantify them separately.

The correlations between the fold-changes of mRNAs and proteins in Figure 3 indicate that the relative contributions of transcriptional and post-transcriptional regulation can vary substantially depending on the tissues compared. Thus, the level of gene regulation depends at least to come extent on context. For example transcriptional regulation is contributing significantly to the dynamical responses of dendritic cells (Jovanovic et al., 2015) and to the differences between spleen and kidney (Figure 3a) but much less to the differences between spleen and thyroid gland (Figure 3a). All data, across all profiled tissues, suggest that post-transcriptional regulation contributes substantially to the across-tissue variability of protein levels. The degree of this contribution depends on the context.

Indeed, if we only increase the levels for a set of mRNAs without any other changes, the corresponding protein levels must increase proportionally as demonstrated by gene inductions (McIsaac et al., 2011). However, the differences across cell-types are not confined only to different mRNA levels. Rather, these differences include different RNA-binding proteins, alternative untranslated regions (UTRs) with known regulatory roles in protein synthesis, specialized ribosomes, and different protein degradation rates (Mauro and Edelman, 2002; Gebauer and Hentze, 2004; Rojas-Duran and Gilbert, 2012; Castello et al, 2012; Arribere and Gilbert, 2013; Slavov et al, 2014b; Katz et al, 2014). The more substantial these differences, the bigger the potential for post-transcriptional regulation. Thus cell-type differentiation and commitment may result in much more post-transcriptional regulation than observed during perturbations preserving the cellular identity. Consistent with this possibility, mRNA fold-changes can account for less than 50% of the measured across-tissue variability; the remaining variability is likely due to substantial tissue-specific post-transcriptional regulation; in contrast, stimulating dendritic cells elicits a strong transcriptional response but not change in the cell-type and thus less cell-type-specific post-transcriptional regulation (Jovanovic et al, 2015).
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Methods

Data and scaled mRNA levels

We used data from Wilhelm et al. (2014); Fagerberg et al. (2014) containing estimates for the mRNA levels (based on RNA-seq) and for the protein levels (based on mass-spectrometry) of \( N = 6104 \) genes measured in each of twelve different human tissues: adrenal gland, esophagus, kidney, ovary, pancreas, prostate, salivary gland, spleen, stomach, testis, thyroid gland, and uterus. For these genes, about 8% of the mRNA measurements and about 40% of the protein measurements are missing.

First, denote \( m_{ij} \) the log mRNA levels for gene \( i \) in condition \( j \). Similarly, let \( p_{ij} \) denote the corresponding log protein levels. First, we normalize the columns of the data, for both protein and mRNA, to different amounts of total protein per sample. Any multiplicative factors on the raw scale correspond to additive constants on the log scale. Consequently, we normalize data from each tissue-type by minimizing the sum of squared differences between data from that tissue and the first tissue (chosen to serve as a baseline). Specifically, for all proteins and conditions \( j > 1 \), we normalize each measurement by setting

\[
p_{ij}^n \leftarrow p_{ij}^u - \frac{1}{N} \sum_i (p_{i1}^u - p_{ij}^u)
\]

Where \( p_{ij}^n \) and \( p_{ij}^u \) represent the normalized and non-normalized protein measurements respectively. We conduct the same normalization for mRNA. This normalization corrects for any multiplicative differences in the raw mRNA or protein.

After normalization, we define \( r_{ij} = p_{ij} - m_{ij} \) as the log PTR ratio of gene \( i \) in condition \( j \).
If the post-transcriptional regulation the $i^{th}$ gene were not tissue-specific, then the $i^{th}$ PTR ratio would be independent of tissue-type and can be estimated as

$$\hat{T}_i = \text{median}_j (p_{ij} - m_{ij})$$

Then the log “scaled mRNA” (or mean protein level) can be defined as

$$\bar{p}_{ij} = m_{ij} + \hat{T}_i$$

On the raw scale this amounts to scaling each mRNA by its median PTR ratio and represents an estimate of the mean protein level. The residual difference between the log mean protein level and the measured log protein level

$$r_{ij} = p_{ij} - \bar{p}_{ij}$$

consists of both tissue-specific post-transcriptional regulation and measurement noise.

The fraction of total protein variance explained by scaled mRNA levels is not informative about the across-tissue variance explained by scaled mRNA levels.

We use the measured mRNA levels and the estimated median PTR ratios from Wilhelm et al (2014) to simulate protein levels from a log-normal distribution (see Methods). Since the dynamic range of meal-level variability (between different proteins) is large, increasing the variability in the PTR ratios across tissue-types has negligible impact on $R^2_T$. However, the dynamic range of across-tissue variability (within a protein and across tissues) is significantly smaller, and thus increasing the variability in the PTR ratios has a dramatic impact on the within-gene explained variance, $R^2_P$ (Figure S1). As we demonstrate below, this simple model is consistent with the data for all tissue-types. Tissue-specific mRNA levels and estimated median PTR ratios together explain most of the total (conflated) variance simply because this variance is dominated by the mean-level protein variability. Within a gene, however, the variability in the PTR ratio may contributed substantially to across-tissue variability, and we next explore this possibility.
Across-Tissue Correlations

For each gene, $i$, we can compute the correlation between mRNA and protein across tissues. Unlike the between gene correlations which are consistently large after scaling for each tissue (Figure S3a), Figure S3b shows that the across-tissue correlations are highly variable between genes. Thus, to find a representative population estimate of this within gene correlation, we pool information between genes.

First, for a gene $i$, we compute the fisher transformed within gene correlation as $z_i = \text{arctanh}(\hat{r}_i)$ which is approximately normal:

$$z_i \sim N \left( \frac{1}{2} \log\left(\frac{1 + \rho}{1 - \rho}\right), \frac{1}{\sqrt{N_i - 3}} \right)$$

where $N_i$ are the number of observed mRNA-protein pairs for gene $i$ (at most 11) and $\rho$ corresponds to the population correlation. We can then easily find the maximum likelihood estimate of the Fisher transformed population correlation by weighting each observation by its variance:

$$\omega_i = \frac{1}{n_i - 3}$$
$$W_i = \frac{\omega_i}{\sum_j \omega_j}$$
$$\hat{z}_{pop} = \sum W_i z_i$$

Transforming this estimate back to the correlation scale, we find the population correlation to be:

$$\hat{\rho} = \frac{e^{2\hat{z}_{pop}} - 1}{e^{2\hat{z}_{pop}} + 1} \approx 0.35$$

For each gene we compute the z-score, $z^*_i = \frac{z_i - \frac{1}{2} \log\left(\frac{1 + \hat{\rho}}{1 - \hat{\rho}}\right)}{\sqrt{N_i - 3}}$, and compute the corresponding q-value. Only three genes had FDR less than 1% and only 9 genes had an FDR of less than 10%.
% The empirical variability of the across-tissue correlations is thus consistent with the sampling variability we would expect if the true population correlation was 0.35 and suggests that the lack of correlation between mRNA and protein across tissues may simply be due to noise. Figure S3b-c demonstrate the similarity between the observed empirical correlations and simulated data. Below we address this by directly estimating and correcting for noise.

**Noise Correction**

*measured* mRNA and protein across tissues. Measurement noise attenuates estimates of correlations between mRNA and protein level (Franks et al., 2015). A simple way to quantify this attenuation of correlation due to measurement error is via Spearman’s correction. Spearman’s correction is based on the fact that the variance of the measured data can be decomposed into the sum of variance of the noise and the signal. If the noise and the signal are independent, this decomposition and the Spearman’s correction are exact (Csárdi et al., 2015).

Below is a proof that the observed empirical variance is the sum of the variance of the signal and the variance of the noise:

- $e_i$ - Expectation at the $i^{th}$ data point; $\bar{e}_i = e_i - \langle e \rangle$

- $\zeta_i$ - Noise at the $i^{th}$ data point; $\langle \zeta \rangle = 0$

- $x_i$ - Observation at the $i^{th}$ data point; $\bar{x}_i = x_i - \langle x \rangle$, $x_i = e_i + \zeta_i$;

\[
\sigma_x^2 = \frac{1}{n} \sum_i \bar{x}_i^2 = \frac{1}{n} \sum_i (\bar{e}_i + \zeta_i)^2 = \\
= \frac{1}{n} \sum_i \bar{e}_i^2 + \frac{1}{n} \sum_i \zeta_i^2 + \frac{2}{n} \sum_i \bar{e}_i \zeta_i \\
\approx \frac{1}{n} \sum_i \bar{e}_i^2 + \sigma_e^2 + \sigma_\zeta^2 + 2 \approx 0
\]

Spearman’s correction is based on estimates of the “reliability” of the measurements, which is defined as the fraction of total measured variance due to signal rather than to noise:
Reliability  \[= \frac{\sigma^2_{signal}}{\sigma^2_{total}}\]

\[= 1 - \frac{\sigma^2_{err}}{\sigma^2_{err} + \sigma^2_{signal}}\]  

Where the noise corrected correlation is then simply

\[\frac{\text{Cor}(X,Y)}{\sqrt{\text{Rel}(X)\text{Rel}(Y)}}\]  

The most direct estimate for across-tissue reliabilities are computed per gene from independent measurements in which all steps, from sample collection to level estimation, are repeated independently. The RNA data contain multiple estimates of mRNA levels per tissue type (Fagerberg et al, 2014). For each tissue type, we split the estimates into two non-overlapping subsets and computed the average mRNA levels for each gene in each subsets. For each gene, we compute the correlation between these two independent estimates (across the tissues) and summarize the distribution in Figure 5a.

The protein data published by Wilhelm et al (2014) do not contain multiple biological replicates. To estimate the replicate reliability of the protein measurements, we downloaded the raw mass-spectrometry data for six tissues (stomach, salivary gland, esophagous, ovary, testis and adrenal gland) and searched them against the human uniprot database with MaxQuant (Cox and Mann, 2008). In the evidence files (supplied as supporting information) we summed the integrated-precursor-areas for each peptide to estimate the abundance of the peptide.

We generate “pseudo-replicates” for each razor protein, by randomly spliting all peptides mapped to that protein (across all tissues) into two non-overlapping subsets. For each tissue, we take the average of the observed intensities in each subset to get an independent estimate of the protein abundance. We then correlate these two distinct estimates of across-tissue protein abundance to estimate reliability of the relative quantification of each protein (Figure 5c).

We also introduce an alternative approach for assessing reliabilities using pairs of tissues. For
a tissue pair, we found all peptides quantified in both tissues and computed the ratios between their corresponding integrated-precursor-areas. For each razor protein, we again split all peptides mapped to it into two non-overlapping subsets and estimated the median fold-change for each subset as an estimate for the fold-change of the protein across the two tissues. These estimates of relative protein quantification are plotted in Figure 5d and may in some cases represent a reasonable approximation to the reliability of relative-protein quantification.

Below, we show why this approach of estimating the reliability of relative-protein quantification provides a reasonable approximation to the reliability needed for estimating the across-tissue mRNA-protein correlations. Here, we decompose the observed log protein level for protein $i$ in tissue $j$ measured in replicate $k$ in terms of the mean protein level across tissues, $\mu_i$, plus tissue specific biological variation, $r_{ij}$, and replicate specific noise, $\epsilon_{ijk}$:

$$p_{ij} = \mu_i + r_{ij} + \epsilon_{ijk}$$

The reliability is the squared correlation between two independent replicates Csárdi et al (2015). Under the above decomposition, the correlation can be expressed as

$$Cor(p_{ijk1}, p_{ijk2}| i = i_0) = \frac{Cov(\mu_{i0} + r_{ioj} + \epsilon_{iojk1}, \mu_{i0} + r_{ioj} + \epsilon_{iojk2})}{\sqrt{Var(p_{iojk1})Var(p_{iojk1})}}$$

$$= \frac{\sqrt{Var(r_{ioj})}}{\sqrt{Var(r_{ioj}) + Var(\epsilon_{iojk})}}$$

Where the last line holds as long as the noise terms, $\epsilon$, are iid. Now we consider an approximation to this correlation, by instead computing the correlation between the replicates of log fold-changes between two tissues. As long as the noise terms, $\epsilon$ and across-tissue protein variations, $r_{ij}$ are iid then:
\[
\text{Cor}(p_{ij_1k_1} - p_{ij_2k_1}, p_{ij_1k_2} - p_{ij_2k_2} | j_1, j_2) = \\
\frac{\text{Cov}(r_{ij_1} - r_{ij_2}) + (\epsilon_{ij_1k_1} - \epsilon_{ij_2k_1}), (r_{ij_1} - r_{ij_2}) + (\epsilon_{ij_1k_2} - \epsilon_{ij_2k_2})}{\sqrt{\text{Var}(p_{ij_1k_1} - p_{ij_2k_1}) \text{Var}(p_{ij_1k_2} - p_{ij_2k_2})}} \\
\frac{\sqrt{\text{Var}(r_{ij_1} - r_{ij_2}) + (\epsilon_{ij_1k_1} - \epsilon_{ij_2k_1})} \text{Var}(r_{ij_1} - r_{ij_2}) + (\epsilon_{ij_1k_2} - \epsilon_{ij_2k_2})}{2 \text{Var}(r_{ij})} \\
\frac{\sqrt{\text{Var}(r_{ij}) + \text{Var}(\epsilon_{ijk})}}{2 \text{Var}(r_{ij}) + \text{Var}(\epsilon_{ijk})} \\
\]

where the last two lines only follow if \(r_{ij_1}\) and \(r_{ij_2}\) have equal variances.

We estimated representative across-tissue reliabilities of the mRNA and the protein measurements from independent estimates for the mRNA and the protein levels, Figure 5a-d. Given these estimates of mRNA and protein reliabilities, we computed the de-noised fraction of across-tissue protein variability explained by transcript levels using Equation 3 a the representative observed data correlation, 0.35. Figure 5e depicts the \(R^2\) regions as a function of the reliabilities of relative mRNA and protein estimates across tissues.

**Functional gene set analysis**

To identify tissue-specific PTR for functional sets of genes, we analyzed the distributions of PTR ratios within functional gene-sets using the same methodology as Slavov and Botstein (2011). We restrict our attention to functional groups in the GO ontology (Consortium et al., 2004) for which at least 10 genes were quantified by Wilhelm et al. (2014). Let \(k\) index one of these approximately 1600 functional gene sets. First, for every gene in every tissue we estimate the relative PTR (rPTR) or equivalently, the difference between log mean protein level and measured protein level:

\[
\hat{r}_{ij} = p_{ij} - \text{median}_{j' \neq j}(p_{ij'} - m_{ij'})
\]

To exclude the possibility that \(\hat{r}_{ij} = 0\) exactly, we require that \(j' \neq j\). When the estimated
rPTR is larger than zero, the measured protein level in tissue $j$ is larger than the estimated mean protein level. Likewise, when this quantity is smaller than zero, the measured protein is smaller than expected. Measured deviations from the mean protein level are due to both measurement noise and tissue specific PTR. To eliminate the possibility that all of the variability in the rPTR ratios is due to measurement error we conduct a full gene set analysis.

For each of the gene sets we compute a vector of these estimated log ratios so that a gene set is comprised of

$$G_{kj} = \{\hat{r}_{i_{1}j}, ..., \hat{r}_{i_{n_{k}}j}\}$$

where $i_{1}$ to $i_{n_{k}}$ index the genes in set $k$ and $j$ indexes the tissue type.

Let $KS(G_1, G_2)$ be the function that returns the p-value of the Kolmogorov-Smirnov test on the distribution in sets $G_1$ and $G_2$. The KS-test is a test for a difference in distribution between two samples. Using this test, we identify gene sets that show systematic differences in PTR ratio in a particular tissue ($j$) relative to all other tissues.

Specifically, the p-value associated with gene set $k$ in condition $j$ is

$$\rho_{kj} = KS(G_{kj}, \bigcup_{j' \neq j} G_{kj'})$$

To correct for multiple hypotheses testing, we computed the false discovery rate (FDR) for all gene sets in tissue $j$ (Storey, 2003). In Figure 4a-c, we present only the functional groups with FDR less than 2% and report their associated p-values. The significance of many of these groups, controlling for false discoveries suggests that not all of the variability in rPTR is due to measurement noise.

**References**


Li JJ, Bickel PJ, Biggin MD (2014) System wide analyses have underestimated protein abundances and the importance of transcription in mammals. *PeerJ* 2: e270


Figure 1. The fraction of total protein variance explained by scaled mRNA levels is not informative about the across-tissue variance explained by scaled mRNA levels. (a) mRNA levels scaled by the median protein-to-mRNA (PTR) ratio correlate strongly with measured protein levels ($R^2_T = 0.77$ over 6104 measured mRNAs and proteins in each of 12 different tissues). (b) A subset of 100 genes are used to illustrate an example Simpson’s paradox: regression lines reflect within-gene and across-tissue variability. Despite the fact that the overall correlation between scaled mRNA and measured protein levels is large and positive $R_T = 0.89$, for any single gene in this set, mRNA levels scaled by the median PTR ratio are negatively correlated to the corresponding measured protein levels ($R_P < 0$).
Figure 2. The total protein variance explained by scaled mRNA levels is not indicative of the correlations between mRNA and protein fold-changes across the corresponding tissue pairs. While scaled mRNA is predictive of the absolute protein levels (a-c, top row), the accuracy of these predictions does not generally reflect the accuracy of protein fold-changes across tissues that are predicted from the corresponding mRNA fold-changes (d-f, bottom row).
Figure 3. Measured protein fold-changes can be predicted from the corresponding mRNA fold-changes only for very few genes (a) As in Figure 2d-f, we computed all pairwise correlations between the mRNA and protein fold-changes for 11 tissue-types. (b) The cumulative distribution of the maximum fold error that results from estimating protein fold-changes by mRNA fold-changes. The error in quantifying protein fold change from mRNA exceeds 10-fold for 1441 proteins (24%, red) and exceeds two-fold for 4952 proteins (84%, blue); the fold-changes of less than 1000 proteins can be estimated from mRNA fold-changes with less than twofold (100%) error (blue).
Figure 4. Concerted variability in the relative protein-to-RNA (rPTR) ratio of functional gene-sets across tissue-types (a) mRNAs coding for the small ribosomal subunit, NADH dehydrogenase and respiratory proteins have much higher protein-to-mRNA ratios in kidney as compared to the median across the other 11 tissues (FDR < 2%). In contrast mRNAs coding for focal adhesion have lower protein-to-mRNA ratios (FDR < 2%). (b) The stomach also shows very significant rPTR variation, with low rPTR for the small ribosomal subunit and high rPTR for tRNA-aminoacylation (FDR < 2%). (c) Summary of rPTR variability, as depicted in panel (a-b), across all tissues and many gene ontology (GO) terms. Metabolic pathways and functional gene-sets that show statistically significant (FDR < 2%) variability in the relative protein-to-mRNA ratios across the 12 tissue types. All data are displayed on a log10 scale, and functionally related gene-sets are marked with the same color.
Figure 5. Contributions of transcriptional and post-transcriptional regulation to the observed across-tissue variability in protein levels (a) The reliability – defined as the fraction of the measured variance due to the signal – of relative mRNA levels is estimated as the $R^2$ between the mRNA levels measured in eleven different tissues. Replica estimates for the levels of each transcript were correlated (averaging across the 11 tissue-types) and the results for all analyzed transcripts displayed as a distribution. (b) The reliability of relative mRNA levels is estimates as the $R^2$ between independent estimates of the fold-changes between the salivary and the adrenal glands. (a) The reliability of relative protein levels is estimated as the $R^2$ between the protein levels measured in six different tissues. Separate estimates for each protein were derived from non-overlapping sets of peptides and were correlated (averaging across the six tissue-types) and the results for all analyzed proteins displayed as a distribution; see Methods. (d) The reliability of relative protein levels is estimates as the $R^2$ between two separate estimates of the fold-changes between the salivary and the adrenal glands. Estimate 1 and Estimate 2 are derived from non-overlapping sets of peptides. (e) The fraction of across-tissue protein variance that can be explained by across-tissue mRNA variance is plotted for reliability levels of the measurements ranging from 35 % to 100 %. 
Figure S1. Simulated data demonstrating change in $R_p^2$ and $R_T^2$ (total correlation) as across-tissue variation increases. As the across-tissue variability increases, the fraction of the total protein variance explained by scaled mRNA decreases ($R_T^2$) only slightly (red), while the fraction of the across-tissue variability that can be explained by mRNA levels plummets sharply (blue, dashed). Here, the red line corresponds exactly to the function $1 - R_T^2$. The larger the total variance is relative to the across-tissue variability, the more pronounced this effect. The grey region represents the approximate range for the Wilhelm et al (2014) data.
Figure S2. The dynamic range of physiological variability is larger for proteins than for mRNAs (a) Distributions of dynamic ranges for mRNAs and proteins quantified by the standard deviations computed on a log scale. Note that the log-scale makes the standard deviation independent of scalar scaling on the linear scale. (b) Distribution of differences between the standard deviations of proteins and their corresponding mRNAs. The larger than zero median indicates that the physiological variability of most genes is larger at the protein level than at the mRNA level.
Figure S3. (a) Histogram for each of correlations for each tissue. The between-gene correlations ($R_T$) of measured protein with unscaled protein are generally low (red), but after scaling mRNA by the median PTR ratio correlations are quite high (blue). (b) Within-gene correlations ($R_P$) of measured protein and scaled mRNA are highly variable due to noise and between gene variability in post-transcriptional regulation. (c) Simulated correlations from a bivariate normal with population correlation 0.35. Empirical across-tissue correlations are largely consistent with sampling variability due to noise. Less than 10 genes show significant difference from true correlation 0.35 at 10% FDR (see Methods)