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A Proteomics Approach to Profiling the Temporal Translational Response to Stress and Growth

MITNCAT combines BONCAT, pSILAC, and TMT to quantify protein synthesis rates

MITNCAT quantified up-regulation of protein folding chaperones during the UPR

MITNCAT revealed EGF-driven protein synthesis in four distinct temporal waves

MITNCAT identified delayed synthesis proteins with enriched rare codons

DATA AND SOFTWARE AVAILABILITY
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A Proteomics Approach to Profiling the Temporal Translational Response to Stress and Growth

Daniel A. Rothenberg,1,2 J. Matthew Taliaferro,3,4,5 Sabrina M. Huber,1 Thomas J. Begley,6 Peter C. Dedon,1,7 and Forest M. White1,2,8,9,*

SUMMARY
To quantify dynamic protein synthesis rates, we developed MITNCAT, a method combining multiplexed isobaric mass tagging with pulsed SILAC (pSILAC) and bio-orthogonal non-canonical amino acid tagging (BONCAT) to label newly synthesized proteins with azidohomoalanine (Aha), thus enabling high temporal resolution across multiple conditions in a single analysis. MITNCAT quantification of protein synthesis rates following induction of the unfolded protein response revealed global down-regulation of protein synthesis, with stronger down-regulation of glycolytic and protein synthesis machinery proteins, but up-regulation of several key chaperones. Waves of temporally distinct protein synthesis were observed in response to epidermal growth factor, with altered synthesis detectable in the first 15 min. Comparison of protein synthesis with mRNA sequencing and ribosome footprinting distinguished protein synthesis driven by increased transcription versus increased translational efficiency. Temporal delays between ribosome occupancy and protein synthesis were observed and found to correlate with altered codon usage in significantly delayed proteins.

INTRODUCTION
Cellular response to perturbation often leads to a change in cell state, accompanied by dynamic alterations in protein synthesis and degradation that ultimately result in changes in protein expression levels (Golan-Lavi et al., 2017). Measuring changes in mRNA abundance is commonly used to estimate changes in protein expression; however, relative mRNA abundance has been shown to be an incomplete predictor of protein synthesis and abundance (Schwanha¨usser et al., 2011; Jovanovic et al., 2015) because translation is a highly regulated process that can be modulated by signaling pathways (Rowlands et al., 1988; Feng et al., 1992; Chen and London, 1995; Berlanga et al., 1999; Gingras et al., 2001; Novoa et al., 2003), RNA structural elements (Filbin and Kieft, 2009), and tRNA isoacceptor availability (Chan et al., 2010, 2012; Chionh et al., 2016).

Ribosome footprint (RFP) analysis, the identification of mRNA transcript fragments that are shielded by ribosomes, presumes that ribosome-bound transcripts are being translated into proteins and is considered the gold-standard RNA-based approach to estimate translation rates. RFP analysis involves the isolation and sequencing of ~30 nucleotide mRNA fragments shielded by the ribosome from nuclease degradation (Ingolia et al., 2009; Ingolia, 2016). Since increased RFP abundance could result from increased ribosome density or increased transcript expression with constant ribosome density, normalizing RFP with transcript expression (typically measured by mRNA sequencing [mRNA-seq]) provides a metric known as translational efficiency (TE), effectively reading out the ribosome occupancy per transcript. Although RFP and TE provide a fairly accurate estimate of potential protein synthesis rates, these measurements do not account for stalled ribosomes and have been shown to be less representative of protein synthesis rates during cell stress response (Iwasaki and Ingolia, 2017; Liu et al., 2017).

Proteomics approaches quantify the protein product rather than the RNA precursors of protein synthesis. Two techniques, pulsed SILAC (pSILAC) (Schwanha¨usser et al., 2009) and bio-orthogonal non-canonical amino acid tagging (BONCAT) (Dieterich et al., 2006), enable direct measurement of newly translated proteins. In pSILAC, heavy-isotope labeled amino acid analogs are added to cells in culture and are incorporated into newly synthesized proteins over a defined time period before mass spectrometry (MS)-based analysis. This approach allows for an estimation of protein turnover by comparing the abundance of the...
heavy, newly made peptides with the light, preexisting peptides (Doherty et al., 2005) and can be performed on up to two conditions simultaneously (Schwanhäusser et al., 2009). Owing to dynamic range and sensitivity limitations, it can be difficult to detect pSILAC-labeled, newly translated proteins against the large background of pre-existing proteins in the cell (Eichelbaum and Krijgsveld, 2014). These challenges effectively limit minimum incorporation time and make it difficult to monitor low-abundance proteins. However, targeted MS approaches such as multiple reaction monitoring have been used with pSILAC to quantify synthesis rates for selected proteins (Liu et al., 2017). In BONCAT, azidohomoalanine (Aha) (Dieterich et al., 2006), an azide-modified methionine analog used naturally by the native methionyl tRNA synthetase (MetRS) (Kiick et al., 2002), is added to cells and incorporated into newly synthesized proteins. The azide functional group on Aha enables selective enrichment through click chemistry-mediated solid-phase capture of Aha-labeled proteins. Combining BONCAT with pSILAC (e.g., BONLAC [Bowling et al., 2016]/QuanCAT [Howden et al., 2013] and HILAQ [Ma et al., 2017]) improves the sensitivity and coverage of pSILAC and provides a quantitative comparison of protein synthesis rates across two conditions (Eichelbaum et al., 2012). Using multiple MS analysis, this combined approach has been used for temporal analysis of newly synthesized proteins following macrophage activation (Eichelbaum and Krijgsveld, 2014). However, prolonged overlapping metabolic labeling periods prevented analysis of rapid changes in protein synthesis, and the use of pSILAC limited the number of time points assayed (Eichelbaum and Krijgsveld, 2014).

With the goal of developing a method that would allow for high sensitivity analysis of newly translated proteins at multiple time points with high temporal resolution, we developed MITNCAT (multiplex isobaric tagging/non-canonical amino acid tagging), combining BONCAT with pSILAC and using multiplexed isobaric tandem mass tagging (TMT) (Thompson et al., 2003) to quantitatively compare translation rates for thousands of proteins across ten different conditions in a single MS experiment. Here the combination of BONCAT and pSILAC enables enrichment for newly translated proteins and post hoc removal of non-specifically retained proteins from BONCAT enrichment, since newly translated proteins should all have pSILAC labels. Multiplex isobaric tagging generates quantification of newly synthesized proteins for discrete time bins within a single experiment. Previous studies have combined pSILAC and TMT to monitor protein turnover (Welle et al., 2016), whereas here we combine BONCAT, TMT, and pSILAC to describe the temporal dynamics of protein synthesis rates at discrete time points following stimulation. Application of MITNCAT to the unfolded protein response enabled the temporal analysis of thousands of protein synthesis rates and highlighted the differential translation regulation of a large number of metabolic and translational regulatory proteins.

We also applied MITNCAT to quantify protein synthesis rates following epidermal growth factor (EGF) stimulation of HeLa cells, a system whose dynamic response has been well characterized across miRNA (Avraham et al., 2010), transcript (Amit et al., 2007), protein expression (Waters et al., 2012; Shi et al., 2016; Golan-Lavi et al., 2017), and protein post-translational modification (Zhang et al., 2005; Reddy et al., 2016). Although the temporal dynamics at each of these levels has been shown to affect the cellular response to EGF, the effect on dynamic protein synthesis rates has yet to be characterized. Here we applied MITNCAT with discretely timed pulses of Aha and pSILAC to quantify the temporal dynamics of protein synthesis rates for thousands of proteins at multiple time points following EGF stimulation. These data document the temporal control of protein synthesis, including increased synthesis rates of dozens of proteins within the first 15 min, previously unprecedented temporal resolution. Comparison of proteomic synthesis rate data to mRNA-seq and RFP data at each time point established the transcriptional versus translational efficiency-based control of protein synthesis. Furthermore, we compared the temporal dynamics of protein synthesis and RFP, uncovering a potential role of codon bias in regulating temporal delays in protein synthesis.

**RESULTS**

**Isobaric Mass Tags Allow for Robust Multiplexed Quantitative Analysis of Newly Synthesized Proteins**

Several publications over the past decade have documented the dynamic regulation of miRNA, mRNA, and protein expression following cell stimulation. Realizing that protein synthesis rates are likely similarly dynamic, we developed MITNCAT, a method to accurately quantify the temporal dynamics of protein synthesis rates at a global scale. In MITNCAT, multiplex isobaric tags enable the simultaneous analysis of newly synthesized proteins from multiple discrete time points following cellular stimulation. Briefly, Aha-based BONCAT is used to label newly synthesized proteins with a bio-orthogonal chemical handle and
heavy-isotope labeled arginine and lysine pSILAC amino acids are added to the media concurrently with Aha. Proteins synthesized during the labeling period therefore incorporate Aha (for enrichment) and pSILAC (as a marker for specificity). Aha-labeled proteins are solid-phase captured onto a dibenzocyclooctyne (DBCO)-functionalized resin through a copper-free click chemistry reaction. After multiple rounds of highly stringent washes, the bound proteins are digested, on bead, to liberate pSILAC-labeled peptides. In the same tube, multiplex isobaric tandem mass tags are added to each sample to allow for quantitative MS analysis. The samples are then combined, subjected to high pH reverse phase fractionation, and analyzed via liquid chromatography-tandem mass spectrometry (LC-MS/MS) for discovery-mode, quantitative measurement of temporal dynamics of protein synthesis rates.

To validate the accuracy of this quantitative approach to measure newly translated proteins, we chose a model system, MCF10a cells, and incubated the cells with 3 mM Aha for increasing periods of time: 15, 30, 60, and 120 min. Assuming that proteins were being synthesized at a continuous rate over this experiment, Aha-labeled newly synthesized proteins should increase linearly with time. Two negative controls were included in this experiment: methionine (Met) was substituted for Aha to assess the level of background binding, and 300 μg/mL cycloheximide (CHX) was added to the media 30 min before labeling with Aha for 120 min (Figure 1A, left) to determine if blocking translation could prevent Aha incorporation into proteins. Peptides were labeled with one of six different tandem mass tags (Figure 1A, right) that allow for relative quantitation of newly translated proteins among the different samples. The samples were then combined, subjected to high pH reverse phase fractionation, and analyzed via liquid chromatography-tandem mass spectrometry (LC)-MS/MS for discovery-mode, quantitative measurement of temporal dynamics of protein synthesis rates.

Figure 1. Analysis of Aha Labeling Time Course Demonstrates Reliable Quantitation of Newly Synthesized Proteins across Large Dynamic Range

Newly synthesized proteins were labeled with Aha and pSILAC for variable amounts of time, enriched onto a DBCO-functionalized resin, digested with trypsin, and eluted peptides were labeled with isobaric mass tags (A); TMT reporter ions used for quantitation from exemplar spectra (B); a fitted line to the medians of all Aha-labeled samples demonstrates accuracy of quantification over a broad dynamic range (C); western blot for biotinylated Aha residues of newly synthesized proteins in inputs before click enrichment (left) and in the supernatant following click enrichment (right) (D). Data are median ± SEM.
Since TMT-based quantification is relative and not absolute, the 15-min Aha-stimulated channel was used for normalization. Taking the median of all proteins observed in each channel, a more than 2-fold decrease was observed in the Met and CHX samples compared with the 15-min Aha-labeled sample, whereas the 30-, 60-, and 120-min labeled Aha channels each featured an increase in intensity that corresponded with the increase in duration of Aha labeling (Figure 1C). A line fitted to the four Aha-labeled time points has a slope of 0.87, slightly below the expected value of 1. This slight compression in the MS-based quantification was repeated in a subsequent replicate (Figure S1A). To determine the source of this deviation, we used an orthogonal, fluorescence-based method to quantify Aha-labeled proteins from each time point. An aliquot of the input and supernatant from each pull-down was analyzed by reacting each protein sample with DBCO-biotin to click on a biotin tag. Quantitative western blotting was then performed using a fluorophore-conjugated streptavidin as a probe. This analysis revealed time-dependent increases in Aha-labeling with a slope of 1 (Figure S1B) and robust depletion of Aha-labeled proteins from the supernatant (Figure 1D), suggesting that the decreased slope of the MS-based quantification is mostly likely not due to translation suppression associated with the decreased incorporation of Aha in place of Met (Kiick et al., 2002) but may instead be due to suppression of dynamic range in MS-based quantification (Savitski et al., 2013). Individual proteins, although showing increased variation, still demonstrate a robust linear increase in protein synthesis in response to longer labeling periods (Figure S1C), further suggesting compression due to isolation interference. Increasing the degree of sample fractionation could potentially improve the dynamic range, but given that we were able to fairly accurately quantify an 8-fold change in translation rate, we determined that the current approach should be suitable for further applications. Stratifying proteins based on degradation kinetics (McShane et al., 2016) yielded no differences (Figure S1E), suggesting that protein degradation kinetics do not play a major role in quantification on the timescales tested.

Analysis of Protein Synthesis during the Unfolded Protein Response Shows an Up-regulation of Stress Proteins in the Background of Global Translational Repression

To validate the robustness of this method in a biological context, we selected the unfolded protein response (UPR), a stress response known to drive inhibition of protein synthesis to prevent further protein misfolding while also up-regulating protein folding chaperones to aid in the repair of misfolded proteins.

To induce the UPR, 10 μg/mL tunicamycin was added to cells and newly synthesized proteins were labeled with Aha, K8, and R10 for 30 min starting at either 0, 1, 2, 3 or 4 hr after treatment (Figure S2A). DMSO-treated controls were collected at matched time points to account for differences in starvation length as well as any artifacts that may arise owing to the presence of Aha (Figure S2A).

MITNCAT led to the quantification of a total of 3,178 unique proteins, with 2,007 unique proteins appearing in at least two of the three replicates and therefore considered for subsequent analyses. When comparing the median of log2-transformed time points, we observed a global decrease in protein synthesis starting between 30 and 90 min following tunicamycin treatment compared with DMSO controls (Figures 2A and S2B). This observation is consistent with previous reports showing that global translation decreases owing to phosphorylation of eIF2α at approximately 30 min following endoplasmic reticulum (ER) stress (Novoa et al., 2003).

All proteins were subjected to clustering using a self-organizing map. To identify the most robust clusters, this process was repeated 1,000 times using random initial seeds to generate a co-clustering frequency map (Figure 2B). This approach led to the identification of two distinct clusters: one characterized by protein synthesis down-regulated below the median (cluster 1, Figure S2C) and one characterized by protein synthesis up-regulated above the median (cluster 2, Figure S2C).

Gene ontology (GO) term analysis revealed that the down-regulated cluster 1 is statistically significantly enriched for the terms protein targeting to the ER ($p = 2.87 \times 10^{-19}$), translational initiation ($p = 2.80 \times 10^{-19}$), and ribosome biogenesis ($p = 7.15 \times 10^{-10}$). The down-regulation of proteins involved in
translation, such as ribosomal proteins, translation initiation factors, and translation elongation factors, (Figure 2C, left and center) may enhance the suppression of protein synthesis in cells undergoing the UPR.

The down-regulated cluster also contained proteins associated with glycolysis. The down-regulation of proteins involved in glycolysis, including GAPDH (Figure 2C, right), ENO1, TP11, PKM, and ALDOA (Figure S2D), suggests that metabolic flux through the glycolytic pathway may be reduced during the UPR. It has been previously reported that tunicamycin treatment reduces glucose uptake, lactate production, and ATP levels (Wang et al., 2011), consistent with the observed down-regulation of proteins associated
with these pathways. Interestingly, proteins involved in the tricarboxylic acid cycle, such as citrate synthase, SDHA, and IDH3, were not included in this down-regulated cluster (data not shown).

Cluster 2 contained proteins that were down-regulated to a lesser extent and also included proteins that were up-regulated in the background of global translation repression. One of the most highly up-regulated proteins in this cluster was BIP, which is the canonical ER stress response chaperone. Other proteins that exhibited a large increase in translation rate following tunicamycin treatment were HERPUD1, which is involved in targeting proteins for degradation via the ERAD pathway, and HYOU1, another member of the heat shock family of proteins involved in protein folding and cell survival in response to stress (Figure 2D). Other stress-related proteins were also observed to be up-regulated (Figure S2E).

Since the inhibition of translation during the UPR is mediated by the phosphorylation of eIF2α by PERK, we hypothesized that the observed overall repression of translation would not be seen by measuring RNA transcript abundance. Indeed, mRNA-Seq did not recapitulate this global down-regulation of protein synthesis measured by MITNCAT (Figures S2F–S2H), as only selected transcripts were affected by tunicamycin treatment. Altogether, these data demonstrate the reliability of MITNCAT for quantifying the time course of protein synthesis changes at a global level in a complex biological system.

EGF Stimulation Results in Temporally Distinct Waves of Protein Synthesis

It has been shown that epidermal growth factor receptor (EGFR) activation leads to temporally distinct waves of transcription, in which immediate-early genes (IEGs) are followed by delayed early genes (DEGs) and finally late response genes (LRGs), which are up-regulated 2 to 4 hr after stimulation (Amit et al., 2007; Avraham and Yarden, 2011; Feldman and Yarden, 2014). To determine whether temporally distinct waves also occurred for protein synthesis, we applied MITNCAT to quantify proteome-wide protein synthesis rates temporally distinct time windows following EGF stimulation. HeLa cells were serum starved for 24 hr and stimulated with 20 nM EGF, and Aha, K8, and R10 were concurrently applied in consecutive 30-min windows following EGF addition, resulting in time points collected at 30, 60, 90, 120, and 150 min (Figure S3A). To account for the effects of the KRM-free media starvation and Aha labeling, matched PBS controls were also collected at the same time points.

Following processing and analysis, 1,749 unique proteins were identified across four replicates, with 1,007 unique proteins observed in at least two replicates and retained for subsequent analysis. To visualize the temporal dynamics of protein synthesis following EGF stimulation, proteins whose synthesis was statistically significantly altered in at least one time point were subjected to hierarchical clustering. The resulting heatmap highlights the temporal dynamics of synthesis of selected proteins, with some proteins peaking at 60–90 min post-treatment, whereas others feature increased synthesis as early as within the first 30 min of EGF stimulation (Figure 3A). Although there was a trend toward increased global protein synthesis following EGF stimulation, this trend did not reach statistical significance (Figure S3B).

To group proteins into temporally distinct clusters, k-means clustering analysis was performed using six clusters and Pearson correlation as the distance metric. To identify the most robust clusters, this process was repeated 10,000 times using a random initial seed; the results of this analysis were then plotted on a co-clustering frequency map (Figure 3B). This analysis revealed four distinct clusters, each with a unique temporal profile (Figure S3C).

Cluster 1 was characterized by increased protein synthesis as early as 30 min following EGF stimulation, with maximum synthesis at 60 min before returning to baseline levels by 150 min. This cluster contained many of the canonical IEGs, including EGR1 (Figure 3C), JUN, CYR61, and IER2 (Figure S3D), consistent with a model of rapid up-regulation of these genes within the first hour following stimulus. Intriguingly, this cluster also contained many of the canonical DEGs, such as DUSP1, ATF3, and ZFP36L1 (Figure S3D). This rapid up-regulation of DEGs was in contrast to the established literature that shows most DEG transcripts being up-regulated between 1 and 2 hr following EGF stimulation (Avraham and Yarden, 2011; Feldman and Yarden, 2014). One notable IEG absent from the dataset was MYC, which has previously been shown to be up-regulated on the transcript level in response to EGF stimulation (Amit et al., 2007). MYC was modestly up-regulated between 30 and 90 min, but not to an extent that was statistically significant in this analysis. This observation suggests that previously observed increases in MYC protein levels could
be due to a combination of increased stability (Sears et al., 2000) and a modest increase in protein synthesis.

In response to EGF stimulation, synthesis of proteins in cluster 2 increased at 60 min, was maximal at 90 min, and subsequently decreased back toward basal levels. Similar to cluster 1, cluster 2 contained IEGs such as NR4A1 (Figure 3D) and NR4A3 and the DEG KLF10 (Figure S3E). Within clusters 1 and 2, synthesis of both IEGs and DEGs were observed between 30 and 90 min following EGF stimulation. Rather than showing a separate wave of IEGs followed by a wave of DEGs, as has been implicated by transcriptional analysis, IEGs and DEGs appeared to be collectively expressed within the same time frame with maximal synthesis occurring between 60 and 90 min following EGFR activation.

Cluster 3 was characterized by a delayed response to EGFR activation, with protein synthesis beginning at around 90 min and increasing through the final time point, 150 min after stimulation. This cluster contained several LRGs involved in cytoskeletal dynamics and cell motility, such as CTEN (Figure 3E), VASP, EZR, and EPPK1 (Figure S3F).

Finally, cluster 4 was characterized by an increase in synthesis beginning around 30 min and remaining elevated across all time points. This cluster almost exclusively contained proteins associated with the translational machinery, including ribosomal proteins, translation initiation factors such as eIF4B (Figure 3F), and translation elongation factors (Figure S3G). These data suggest that, in response to pro-growth cues such as EGF stimulation, cells increase their translational capacity by synthesizing more ribosomal proteins and associated translation factors to further increase synthesis of new proteins. This observed increase in ribosomal protein synthesis is consistent with the observation that ribosomal RNAs (rRNAs) also show an increase in transcription within 30 min following EGFR activation (Stefanovsky et al., 2001).

Figure 3. Changes in Protein Synthesis following EGF Stimulation Were Clustered into Four Groups Based on Temporal Behavior

Temporal changes in protein synthesis over the first 150 min following EGF stimulation (A) were clustered by k-means clustering, resulting in four groups that have distinct temporal profiles (B). Exemplar proteins demonstrate the unique protein synthesis profiles of each group (C–F). n = 4 biological replicates. Data are mean ± SEM, * = p < 0.05.
Increasing Sampling Frequency Yields New Insights on Protein Synthesis Dynamics

To better characterize the temporal response to EGF, especially for proteins that displayed dynamic protein synthesis such as those in clusters 1 and 2, we tested whether we could increase the temporal resolution of MITNCAT. Aha/K8/R10 labeling times were reduced from 30 min to 15 min, with samples collected at 15, 30, 45, 60, and 75 min following EGF treatment along with matched negative controls as described previously. In this analysis, 1,857 proteins were identified in total, with 1,135 proteins identified in two or more replicates. Somewhat to our surprise, decreasing labeling time to 15 min did not lead to a significant decrease in the number of identified proteins relative to the 30-min labeling experiments, suggesting that the dynamic range of the experiment (the difference between the most abundant and least abundant detectable proteins), rather than the overall sensitivity, might be the limiting factor in number of identified proteins. To visualize the dynamic profiles of protein synthesis, proteins with significantly altered synthesis following EGF stimulation were clustered to generate a heatmap (Figure 4A). This heatmap demonstrates that some proteins are rapidly synthesized within 15 min of EGF stimulation, whereas many additional proteins feature a strong increase between 30 and 60 min post-treatment. Analysis of this 15-min temporal resolution data by k-means clustering yielded similar clusters as compared with the 30-min resolution data. However, because the time course ended at 75 min, before cluster 2 proteins and LRGs in cluster 3 reach their maximum, LRGs and late IEG/DEGs previously found in clusters 2 and 3 were clustered together. Additionally, although ribosomal proteins and translation factors from cluster 4 remained clustered, the improved temporal resolution of the analysis enabled a bifurcation (clusters 4 and 4*) between those...
proteins that demonstrated an increased synthesis by 15 min compared with those that increased synthesis starting at 30 min (Figure S4A).

Despite the relatively small changes in overall temporal clusters, increasing temporal resolution provided interesting insights at specific time points. For instance, 24 proteins, including DUSP1 (Figure 4B, arrowhead), JUN, and several ribosomal proteins, demonstrated significantly increased protein synthesis as early as 15 min following EGF stimulation. Of these proteins, DUSP1 was one of the fastest responders, with expression increasing nearly 60% within the first 15 min. This rapid up-regulation of a DEG, even before most IEGs, has not been previously observed and stands in contrast to the classical model of DEGs being up-regulated following the expression of IEGs. The ATF3 transcription factor was also notable, as improved temporal resolution highlighted an immediate-early decrease below basal levels in ATF3 synthesis rates at 15 min following EGF stimulation, followed by an increase in synthesis at 30 min that peaked between 45 and 60 min (Figure 4C, arrowhead). This immediate down-regulation of ATF3 is corroborated by RFP analysis (Figure S5B, Table S5) and has not been previously observed.

Comparison of Protein Synthesis to RNA Abundance and Ribosome Footprints Reveal Transcriptionally and Translationally Controlled Groups of Proteins

Increased synthesis of a given protein could be due to many different factors, including an increase in the abundance of the protein-coding transcript or an increase in translation efficiency due to increased number of ribosomes binding to each transcript or the rate at which the ribosomes move along the transcript. To determine the relationship between dynamic transcript expression, ribosome binding, and protein synthesis rates, we performed mRNA-seq and RFP on identical samples corresponding to the 30-, 60-, and 90-min time points following EGF stimulation. TE rates were calculated based on the mRNA-seq and RFP data and represent ribosome occupancy normalized to transcript abundance (see Methods). These data were then compared with the quantitative protein synthesis data at these same time points.

On comparison of protein synthesis with RNA abundance and TE measurements, two distinct groups of genes emerged (Figures 5A and S5A). In the first group, changes in protein synthesis correlated strongly with changes in RNA abundance (Figure 5A blue bar and Figure S5B), suggesting regulation predominantly at the level of transcription. This group contains most of the IEGs and DEGs that are transiently expressed at high levels, such as EGR1 (Figure S5C). This rapid up-regulation of EGR1 was also orthogonally confirmed by western blot, with new EGR1 protein increasing 1.9-fold from 30 to 60 min after EGF treatment (Figure S5C), compared with a 2.2-fold increase as measured by MITNCAT over that time (Figure 5B). Furthermore, co-treatment with proteasome inhibitor MG132 yields no change in newly synthesized EGR1 (Figure S5C), demonstrating that the difference in magnitude between mRNA/RFP levels and protein synthesis is not due to EGR1 protein degradation. In the second group, changes in protein synthesis correlated with variations in TE, with minimal change in RNA abundance (Figure 5A, green bar and Figure S5D), suggesting regulation on the level of translation. This cluster was highly enriched for ribosomal proteins and several translation initiation and elongation factors such as eIF4B (Figure S5C). These proteins were rapidly up-regulated and sustained throughout the duration of the analysis, albeit to a lesser magnitude than the IEGs, DEGs, and LRGs.

One notable difference between the two clusters was the magnitude of the change in protein synthesis. In the transcriptionally driven cluster, protein synthesis increased as high as 7-fold, whereas the most highly up-regulated protein in the translationally driven cluster experienced a 2-fold change in synthesis. Up-regulation solely through increased translation may be limited by the maximum rate at which ribosomes can bind and translate a transcript. Larger changes in protein expression may require increasing the number of mRNA transcripts available to be translated. This observation highlights a potential trade-off between fast but limited up-regulation through translational control and slower but more potent up-regulation through transcriptional control.

Codon Bias Correlates with Temporal Delays between Ribosome Binding and Translation of New Proteins

Generating time course data for ribosome binding and protein synthesis offered the unique opportunity to characterize the temporal relationships between these processes. Because protein synthesis and ribosome binding occur on scales of different magnitudes, the values were standardized to allow for a direct comparison of the temporal profiles for each. To prevent the analysis of random fluctuations in proteins with
unchanging RFP values or protein synthesis rates, only proteins with at least one statistically significant time point (p < 0.05) in both the RFP and MITNCAT datasets were considered, and thus only proteins present in at least two of the four MITNCAT replicates were included in the analysis of temporal delay. These restrictions limited the protein dataset to 90 and the RFP dataset to 400; the overlap between these datasets was 27 proteins. The TE data was compared with MITNCAT data for these 27 proteins. From this analysis, a subset of 17 proteins exhibited a clear delay between ribosome binding and protein synthesis (Figures 6A and S6A and Table S6). One potential cause for this delay is a difference between codon frequency usage in these genes and the corresponding tRNA isoacceptor availability. To determine if these genes exhibit codon usage frequencies that deviate from the genome averages, the average frequency for each codon in each gene (Table S6) was averaged across all the genes in the group. Statistical significance was
Figure 6. Proteins Exhibiting a Delay between Ribosome Binding and Protein Synthesis Have a Significant Bias in Codon Usage

A comparison of RFP analysis and protein synthesis yielded a group of 17 proteins exhibiting a delay between ribosome binding (n = 2) and protein synthesis (n = 3) (A). An analysis of codon usage reveals a statistically significant codon bias present in this set of proteins (B) not present in a randomly generated set of proteins (C). After an analysis of 500 randomly generated protein sets, the extent of this codon bias appears to be unique to these proteins exhibiting a delay between ribosome binding and protein synthesis (D). When considering 500 sets of 100 randomly selected genes, the codons enriched in the set of 17 delayed proteins (Figure 6B, upper right quadrant) occupy the A-site of ribosomes with a higher frequency than other codons (E). Data are mean ± SEM.

determined by generating randomized groups of the same size and comparing the codon usage frequency with the genome average. This process was repeated 10^6 times, and a p value was generated by counting how many randomly generated groups showed a frequency deviation from the genome greater than that of the queried group for each codon. Reported p values were statistically significant if they were less than the
Bonferroni corrected $\alpha = 7.81 \times 10^{-4}$ (accounting for 64 different codons). The cohort of 17 genes exhibiting delayed translation showed a statistically significant codon bias in 14 different codons for 9 different amino acids (Figure 6B). The other 10 genes that did not show delayed translation did not have codon bias compared with the genome average (Figure S6B). To determine if this bias was unique to this group of proteins, 17 random proteins were selected from the whole dataset and queried for codon bias, and this process was repeated 500 times. Of the 500 sets of randomly selected proteins, 469 sets exhibited no statistically significant deviations from the genome average and no sets had a codon bias in more than three amino acids (Figures 6C and 6D). It has been previously reported that longer proteins tend to exhibit a codon bias (Duret and Mouchiroud, 1999), but the distribution of protein length in the set of proteins with delayed translation is not statistically different from that of the entire dataset (Figure S6C).

A majority of the proteins in the delayed translation set share the GO annotation “response to growth factor.” To test whether this codon bias is common among proteins with this GO annotation, we changed the background set from proteins in the MITNCAT dataset to all proteins that have the GO annotation “response to growth factor.” Even after making this change, the proteins exhibiting a delay in translation have a codon bias in nine different amino acids (Figures S6D and S6G) compared with six in the most extreme outlier set drawn from proteins annotated as “response to growth factor” (Figures S6F–S6G). This analysis suggests codon bias is unique to these proteins exhibiting delayed translation and may provide a potential mechanism to explain the temporal discrepancy between ribosome binding and protein synthesis.

If the delay in protein synthesis is indeed due to increased codon bias in these particular transcripts, then ribosomes would be expected to wait longer for a charged tRNA at these codons, and therefore these codons would occupy the A-site of the ribosome with a higher frequency than other codons. To investigate this possibility, we analyzed the frequency of ribosome A-site occupancy for each codon across 500 subsets of 100 randomly selected transcripts from our dataset. When codons were stratified based on whether they were enriched in the 17 proteins with delayed synthesis (Figure 6B, upper right quadrant), we found that these enriched codons occupied the A-site with a greater frequency than other codons across all genes (Figure 6E). This suggests that these particular codons may take longer to get translated, and therefore transcripts enriched for these codons may undergo slower translation, resulting in a delay between ribosome binding and protein synthesis.

**DISCUSSION**

Here, we combined BONCAT, pSILAC labeling, and isobaric mass tagging in a novel method, MITNCAT, that enables highly multiplexed quantitative measurements of protein synthesis across multiple time points. Previous studies directly analyzing new protein synthesis have either relied on targeted approaches with pSILAC (Liu et al., 2017) or BONCAT with a limited number of overlapping time points across multiple MS analyses, resulting in poor temporal resolution (Eichelbaum and Knijgs-veld, 2014).

One of the critical features of MITNCAT is the improved sensitivity associated with multiplex analysis of pSILAC-labeled peptides from Aha tagged and enriched proteins. On-bead proteolytic digestion of captured proteins and subsequent detection of released peptides eliminates the need to release and detect the Aha-tagged peptide and provides multiple peptides per protein to improve quantification accuracy. Critically, incorporation of pSILAC differentiates newly synthesized proteins from potential non-specifically retained background proteins, improving the stringency of the analysis. Finally, labeling peptides from each condition with isobaric tags enables the multiplexed analysis of many conditions while simultaneously increasing the MS and MS/MS signal intensity by the summation of the isobarically tagged peptides. Overall, combining these techniques into a coherent strategy improved the sensitivity of the method, allowing for decreased labeling times and thus improved temporal resolution, as demonstrated by the 15-min temporal resolution following EGF stimulation. Further improvements to the sensitivity, combined with more stringent washing and increased sample loading, should allow for temporal resolution in the minute time frame.

Application to the UPR demonstrated the reliability of this method to measure translationally controlled changes in protein synthesis. Global down-regulation of translation was observed, a result that was not captured via analysis of RNA abundance. Furthermore, one cluster of proteins was down-regulated to a
greater extent than the global average. This cluster contained proteins associated with translational machinery and the glycolysis pathway. The mechanism behind the specific down-regulation of these proteins is not known, but many of these proteins are classified as housekeeping genes that are expressed at high levels across tissue types (Eisenberg and Levanon, 2013). Perhaps because these proteins were expressed at such high levels, global inhibition of translation resulted in a greater degree of down-regulation of these proteins compared with other proteins with lower basal translational rates. We also observed the up-regulation of protein-folding chaperones and other proteins associated with protein degradation and survival consistent with an acute stress response.

Application of MITNCAT to quantify protein synthesis following EGF stimulation enabled us to group proteins based on temporal changes in synthesis. We quantified dynamic synthesis rates for over a thousand proteins and observed IEGs and DEGs to be synthesized simultaneously, peaking between 60 and 90 min post-treatment. LRGs began to be synthesized starting around 90 min and continued to increase in synthesis through 150 min post-treatment. These increases in protein synthesis were matched by observed changes in RNA abundance, suggesting control at the level of transcription. Interestingly, ribosomal proteins and associated translation factors demonstrated increased synthesis as early as 15 min following EGF stimulation, and many of these proteins maintained increased synthesis throughout the time course, yet these changes were not matched by corresponding changes in RNA abundance. The disparity between protein synthesis rates and transcript levels suggested regulation at the level of TE, which was further confirmed by RFP. It is worth noting that the altered synthesis rates of this very large group of proteins could not have been detected by mRNA-seq (transcript expression) alone. RFP would have suggested this increase, but MITNCAT was easily able to detect these changes, including directly measuring increased protein synthesis in the first 15 min following EGF stimulation. Therefore, MITNCAT could also serve as an orthogonal method to validate RFP datasets.

Finally, the comparison of temporal changes in RFP analysis and protein synthesis revealed a class of proteins that demonstrated a temporal delay between ribosome binding and protein synthesis. Analysis of the sequences coding for these proteins revealed a statistically significant bias in their codon usage frequencies that was unique to this group. This correlation opens the possibility for regulation of protein expression based on the availability of specific tRNA isoacceptors and presence of modified nucleosides in tRNAs.

Overall, MITNCAT is broadly applicable to a range of biological systems, provides synthesis rate information for thousands of proteins in a high-throughput, discovery-based approach, and yet can also be coupled to targeted MS-based approaches to quantify temporal dynamics of protein synthesis for a priori selected proteins. The sensitivity of MITNCAT provides high temporal resolution, and application of this approach led to the identification of many proteins whose synthesis was significantly altered as rapidly as 15 min following stimulation. Future application of this approach to other biological systems will provide novel insights into the regulation between transcription and translation.

Limitations of the Study
To efficiently label newly synthesized proteins with SILAC and Aha labels, cells must be subjected to a 30-min starvation of arginine, lysine, and methionine. This amino acid starvation may trigger a stress response in the cells. In the absence of proper controls, this may result in artificially high expression of stress proteins. In this study, all data points are normalized to untreated controls to normalize out these effects. However, when examining protein synthesis during the UPR using MITNCAT, increased stress response proteins in the negative controls may normalize out a portion of the response that would have occurred in response to the UPR. This limitation should be considered when using MITNCAT for studying stress responses.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

DATA AND SOFTWARE AVAILABILITY
All methods can be found in the accompanying Transparent Methods supplemental file. The accession number for the ribosome footprint and transcript sequencing data reported in this paper is NCBI BioProject: PRJNA478455. The accession number for the proteomics data reported in this paper is ProteomeXchange Consortium (PRIDE): PXD009592.
SUPPLEMENTAL INFORMATION

Supplemental Information includes Transparent Methods, six figures, and six tables and can be found with this article online at https://doi.org/10.1016/j.isci.2018.11.004.

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AUTHOR CONTRIBUTIONS

D.A.R., S.M.H., P.C.D., and F.M.W. designed research; D.A.R., J.M.T., and S.M.H. performed research; T.J.B. contributed analytical tools for codon usage bias; D.A.R., J.M.T., S.M.H., P.C.D., and F.M.W. analyzed data; D.A.R., J.M.T., S.M.H., T.J.B., P.C.D., and F.M.W. wrote the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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Supplemental Information

A Proteomics Approach to Profiling the Temporal Translational Response to Stress and Growth

Daniel A. Rothenberg, J. Matthew Taliaferro, Sabrina M. Huber, Thomas J. Begley, Peter C. Dedon, and Forest M. White
Figure S1. Analysis of Aha labeling time course shows reliable quantitation of newly-synthesized proteins across replicates, related to Figure 1. A replicate of the time course in HeLa cells demonstrates excellent quantitation independent of cell line (A). Quantitation of the western blot from Figure 1D demonstrates a linear 1-to-1 relationship between labeling time and labeling intensity (B). A selection of individual protein trajectories reveals a robust labeling time-dependent increase in enrichment (C). Proteins with increased reporter ion abundance have a stronger 1:1 correlation with Aha-labeling time, but even the bottom 10% of proteins by ion intensity have a correlation of over 75% (D). The relationship between labeling time and protein abundance is not impacted by varying protein degradation kinetics characterized in [36] (E). Data are medians +/- SEM.
Figure S2. Temporal profiles of protein synthesis during the unfolded protein response following tunicamycin treatment, related to Figure 2. Newly synthesized proteins were labeled with Aha and SILAC amino acids for 30 minutes every hour following tunicamycin treatment, with matched negative controls to account for changes due to starvation (A). Tunicamycin treatment results in a decrease in global protein synthesis (B), with one cluster of proteins showing a decrease below the median and one cluster showing an increase above the median (C). Proteins associated with glycolysis are in cluster 1, demonstrating especially strong down-regulation (D), whereas other stress response proteins are up-regulated following tunicamycin treatment (E). Measurement of RNA abundance by mRNA-Seq (F) does not capture the global decrease in protein synthesis as measured by BONCAT (G,H). n=3 biological replicates for MS data, n=1 for mRNA-Seq. Data are mean +/- SEM. * = p < 0.05
Figure S3. Changes in protein synthesis following EGF stimulation were clustered into four groups based on temporal behavior, related to Figure 3. Newly synthesized proteins were labeled with Aha and SILAC amino acids in consecutive 30 minute windows following EGF treatment, with matched negative controls to account for changes due to starvation (A). EGF treatment does not result in a statistically significant change in global translation (B). Each protein cluster demonstrates a unique temporal profile in response to EGF treatment (C). Clusters 1 and 2 are transiently expressed proteins and contain IEGs (green) and DEGs (red) (D,E), whereas cluster 3 is up-regulated at a later time and consists of LRGs (blue) (F). Cluster 4 contains primarily proteins associated with translational machinery, and is up-regulated across all time points sampled (G). n=4 biological replicates. Data are mean +/- SEM. * = p < 0.05
Figure S4. Decreasing the duration of Aha labeling increased temporal resolution and yielded new insights into protein synthesis dynamics, related to Figure 4. Temporal changes in protein synthesis were sampled every 15 minutes over the first 75 minutes, and clustered by k-means clustering. n=3 biological replicates.
Figure S5. Temporal response to EGF stimulation compared between transcript expression, ribosome binding, and protein synthesis, related to Figure 5. Statistical significance of the network-wide temporal response to EGF stimulation was assessed at the transcript expression level by mRNA-Seq (n=2), while translation rates were estimated by RFP (n=2), and protein synthesis rates measured by MITNCAT (n=3) (A). Comparing these datasets revealed a set of proteins whose altered synthesis correlated with changes in RNA abundance (blue bar) (B), suggesting regulation at transcription. Another group demonstrated changes in synthesis correlating with changes in translational efficiency (green bar) (C), suggesting regulation at translation. Measuring protein synthesis of Aha-labeled EGR1 by immunoprecipitation followed by click-labeling yielded relative protein synthesis changes in agreement with MITNCAT data. Blocking protein degradation by co-treating with proteasome inhibitor MG132 did not alter the synthesis profile of EGR1 (D).
Figure S6. Proteins exhibiting a delay between ribosome binding and protein synthesis have a significant bias in codon usage, related to Figure 6. A comparison of RFP analysis (n=2) and protein synthesis (n=3) yielded a group of 17 proteins exhibiting a delay between ribosome binding and protein synthesis (A). The other 10 proteins that in this dataset that did not exhibit a delay also did not demonstrate a codon bias (B). Analysis of protein length reveals that the proteins exhibiting delayed translation (red lines) do not differ in length from the rest of the proteins in the dataset (C). Changing the background from all proteins in our dataset to all proteins with the GO annotation “response to growth factor” (RTGF) did not affect the number of amino acids with codon bias in the delayed protein set (D) compared to randomly selected groups of 17 proteins from that background (E-G). Data are mean +/- SEM.
Transparent Methods

Cell Culture

For Aha time course and UPR experiments, MCF10a cells (courtesy of Joan Brugge, Harvard Medical School) were cultured in DMEM:F12 media supplemented with 5% horse serum, 20 ng/mL EGF, 500 µg/mL hydrocortisone, 100 ng/mL cholera toxin, 10 µg/mL insulin, 1% penicillin/streptomycin, and 2 mM glutamine. Cells were passaged every third day at a 1:4 ratio, and all experiments were performed on the third day following passaging. For EGF stimulation experiments, HeLa cells (from ATCC) were cultured in DMEM supplemented with 10% fetal bovine serum (FBS), 1% penicillin/streptomycin, and 2 mM glutamine. Cells were passaged every 3 days and split between 1:4 and 1:6 ratio, and serum starvation was initiated on the second day following passaging.

Western Blot Quantitation

Protein concentration of lysates were measured via bicinchoninic acid (BCA) assays and normalized to 1 mg/mL. For measurements of Aha incorporation, DBCO-biotin was added to the lysates for 1 hour at a concentration of 25 µM. The reaction was quenched by adding sodium azide to roughly 100 mM. LDS buffer was added to 1x, and beta-mercaptoethanol was added to 1%. Samples were loaded onto a NuPAGE Novex 4-12% Bis-Tris Midi Protein Gels (Invitrogen) and run at 160V for 1 hour. Samples were transferred to a nitrocellulose membrane at 100V for 1 hour. Membranes were blocked with LICOR PBS blocking buffer for 30 minutes at room temperature. Biotinylated proteins were probed with IRDye 680RD-conjugated streptavidin (LICOR #32230). The following primary antibodies were used: GAPDH 1:10,000 (Cell Signaling #5174). The following secondary antibodies were used: IRDye 800CW-conjugated goat-anti-rabbit 1:10,000 (LICOR # 926-32211) and IRDye 680LT-conjugated goat-anti-mouse 1:10,000 (LICOR #926-68031). Western blots were imaged on a LICOR Odyssey instrument.
Quantitation of the Western blot was performed using ImageJ software (https://imagej.nih.gov/ij). The average signal intensity of each lane was measured starting at a molecular weight below endogenously biotinylated proteins [1], normalized to the methionine control lane, and log2 transformed.

Metabolic Labeling and Cell Lysis

For all EGF experiments, cells were changed to serum-free media 24 hours before stimulation. In all experiments, growth media was removed and replaced with lysine/arginine/methionine (KRM)-free DMEM:F12 media 30 minutes prior to time course initiation. For EGF experiments, EGF was added to the media to achieve a final concentration of 20 ng/mL or an equivalent volume of PBS was added as a control. For UPR experiments, tunicamycin was added to the media to achieve a final concentration of 10 µg/mL or an equal volume of DMSO was added as a control. At the appropriate time points following treatment, Aha was added to the media to achieve a final concentration of 3 mM, and $^{15}$N$_4^{13}$C$_6$ arginine (R10) and $^{15}$N$_2^{15}$C$_6$ lysine (K8) were added to 0.5 mM. After 30 minutes of Aha/K8/R10 labeling, the media was aspirated, and the cells were washed in ice cold PBS supplemented with 300 µg/mL cycloheximide (CHX). Cells were lysed in 1% SDS in PBS supplemented with 50 mM N-ethylemaleimide (NEM) and 300 µg/mL CHX. 1.5 mL of -20°C acetone was added to each tube immediately following lysis to precipitate proteins. Proteins were precipitated at -20°C for at least an hour and up to overnight.

Sample Processing and Fractionation

Following precipitation, the samples were centrifuged at maximum speed (~21,000 x g) for 15 minutes at room temperature to pellet the precipitated proteins. The supernatant was aspirated off the pellet, and the pellet was allowed to air dry for 15 minutes to evaporate residual acetone. 250 µL of 1% SDS in PBS supplemented with 50 mM NEM was added to the pellet,
along with ~1/10th of a vial of 0.7 mm garnet homogenizing beads to aid in pellet disruption. Samples were alternately vortexed and centrifuged at maximum speed until the pellet was completely solubilized.

Following resuspension, protein concentration was measured via BCA assay, and total protein was normalized to 100-300 µg in 500 µL of 1% SDS in PBS. Each sample was diluted 2-fold in 8 M urea + 850 mM NaCl. 12.5 µL of the lysate was aliquoted for biotin labeling of Aha-labeled proteins and western blot analysis. The remaining volume for each sample was applied to 30 µL of DBCO-agarose beads that had pre-equilibrated by washing 3x in 1 mL 0.8% SDS in PBS. Click enrichment took place overnight at room temperature on a rotor.

After overnight click incubation, the bead supernatant mixture was transferred to an empty spin column and allowed to drain into Eppendorf tubes to collect the supernatant. 12.5 µL of supernatant was aliquoted for biotin labeling of Aha-labeled proteins and western blot analysis. The tubes were rinsed out with 1 mL MilliQ water and added to the spin column, allowing to drain into a waste container. To reduce disulfide bridges, 1 mL 10 mM DTT in 0.8% SDS in PBS was added to the columns and the columns were capped and placed on a rotor at 50°C. Following reduction, the column was drained, and 1 mL 50 mM NEM in 0.8% SDS in PBS was added to the column to alkylate the newly reduced cysteines. The columns were placed on a rotor at room temperature for 30 minutes. The column was once again drained, and the beads were washed 8x with 1 mL 0.8% SDS in PBS, 8x with 1 mL 8 M urea, and 8x with 1 mL 20% acetonitrile (MeCN). After the second wash in each step, the column was capped and allowed to stand for 10 minutes.

Following the final wash step, the beads were completely dried by spinning the excess wash buffer into an empty Eppendorf tube. The beads were resuspended in 300 µL of digest buffer (200 mM triethyl ammonium bicarbonate (TEAB) + 10% MeCN) and transferred to a fresh tube. The column was twice rinsed with 300 µL digest buffer, with the rinses being combined with the sample. The beads were pelleted by centrifuging at 5000 x g for 5 minutes at room
temperature. The supernatant was carefully removed and replaced with 100 µL 1 ng/µL trypsin in digest buffer. The on-bead digest proceeded overnight at room temperature on a rotor.

Following the overnight digest, an aliquot of TMT dissolved in 30 µL anhydrous MeCN was added directly to the tube (beads included). TMT labeling proceeded for 1 hour at room temperature, after which the reaction was quenched with the addition of 15 µL 1 M Tris pH 7.4. The volumes were reduced to about 50 µL in a vacuum centrifuge. All samples (including beads) were combined into a single Eppendorf tube. The individual tubes were rinsed 3x with 40 µL 50% MeCN + 0.1% AcOH, with the rinses being combined with the pooled sample. The sample was completely dried in a vacuum centrifuge.

Following drying, the sample (beads included) was resuspended in 500 µL 10 mM TEAB pH 8. The fritted end of a 200 µm i.d. capillary was placed in the bead pack, and the sample was loaded onto a ZORBAX Extend 300 C18 column (Agilent #770995-902) at 750 psi. The C18 column was placed in line with an HPLC, and the following gradient was run at a flow rate of 1mL/min. (A = 10mM TEAB pH 8, B = 99% MeCN, 10mM TEAB): 0-5 min, 0-5% B; 5-50 min, 5-40% B; 50-59 min, 40-70% B; 59-64 min, 70% B; 64-65 min, 70 to 1% B. Fractions were collected every minute between 5 min and 65 min. Every 12th fraction was concatenated together to give 12 total samples (5 fractions per sample). Samples were placed into a vacuum centrifuge overnight or until the sample reached dryness.

**Mass Spectrometry Analysis**

Dried samples were resuspended in 50 µL 0.1% formic acid (FA). The samples were placed in a ThermoFisher Easy nLC 1000 autosampler and analyzed on a ThermoFisher QExactive Plus mass spectrometer using 25 µL of the resuspended sample. The sample was analyzed using the following gradient over a C18 column (A = 0.1% FA, B = 80% MeCN in 0.1% FA): 0-4 min, 0-14% B; 4-50 min, 13-42% B; 50-57 min, 42-60% B; 57-60 min, 60-100% B; 60-68 min, 100% B; 68-69 min, 100-0% B; 69-75 min, 0% B. The instrument was operated in data
dependent acquisition mode, with the top 15 most abundant precursors with charge of +2 or greater selected for fragmentation and dynamic exclusion set to 15 s. Precursors were isolated with a window of 0.4 \( m/z \) and fragmented via HCD at 33 NCE. Precursor scan settings were set to AGC = 3e6, maximum IT = 50 ms, and resolution of 70,000. MS2 scan settings were set to AGC = 1e5, maximum IT = 300 ms, and resolution of 35,000. The total acquisition time was 75 minutes per sample.

MS data files were searched on MASCOT version 2.4 with fixed modifications for NEM alkylation on cysteines (+125.047 Da), addition of TMT 6-plex to N-termini and lysine residues (+229.163 Da). Variable modifications were SILAC R10 on arginine residues (+10.008 Da), SILAC K8 on lysine residues (+8.014 Da), addition of TMT 6-plex to SILAC K8 lysine residues (+237.177 Da), Aha substitution for methionine residues (-4.986 Da), diaminobutyrate (reduced Aha) substitution for methionine residues (-30.976 Da), oxidation of methionine residues (+15.995 Da) and phosphorylation on tyrosine, threonine, and serine residues (+79.966 Da). Precursor tolerance was 10 ppm, fragment tolerance was 15 mmu, two missed cleavages were allowed, and the enzyme was set to trypsin. Peptides were considered to be positively identified if they had a score of at least 25 and newly translated if they contained a SILAC or Aha residue. Peptides with TMT reporter ion intensities less than 1000 in any one channel were discarded. To control for technical variation between channels, in the absence of a statistically significant change in global protein synthesis rate across replicates, values were normalized to the median of each channel within each replicate. Therefore, the EGF dataset was median normalized, but the tunicamycin dataset was not. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE [2] partner repository with the dataset identifier PXD009592.

**RNA Sequencing and Ribosome Footprint Analysis**

Cells were subjected to identical KRM-free media pretreatment, Aha/K/R labeling, and
PBS+CHX washing conditions as described previously to account for any effects that may be caused by these treatments. Cells were lysed and processed using the Illumina TruSeq Ribo Profile kit (Illumina #RPHMR12126) according to the manufacturer’s protocol. RFP samples were sequenced on an Illumina NextSeq instrument with 50 nt single-end reads and 6 nt barcodes with 6 samples per lane. Adapter sequences were removed from the 3’ end using Cutadapt. Reads were then mapped to hg38 rRNA sequences using STAR. Reads that aligned to rRNA sequences were removed. The remaining reads were then mapped to an hg38 annotation (Gencode release 26). Reads that uniquely map to this annotation were then quantified using Salmon and a FASTA file containing all hg38 cDNA sequences to generate transcripts per million (TPM) values and counts. A library for total RNA was prepared using the Illumina NeoPrep System and sequenced on an Illumina NextSeq instrument with 40 nt paired-end reads and 6 nt barcodes with 12 samples per lane. For analyses requiring alignment, these reads were mapped to the same hg38 (Gencode release 26) annotation. Quantification was done using Salmon and a FASTA file containing all hg38 cDNA sequences to generate TPM and count values. Only ribosome footprints that mapped to coding regions were considered for quantification and subsequent analyses. Ribosome profiling TE changes and associated p-values were calculated using the Salmon-derived count data and the Xtail package [3]. All ribosome footprint and transcript sequencing data have been deposited to NCBI’s BioProject database under accession number PRJNA478455 (https://www.ncbi.nlm.nih.gov/bioproject/478455).

**Self-Organizing Map (SOM) Clustering Analysis**

A self-organizing map (SOM) was used to cluster proteins from UPR experiments that exhibited similar protein synthesis dynamics following tunicamycin treatment. Clustering analysis was performed using the Self Organizing Map Toolbox MATLAB package (http://www.cis.hut.fi/projects/somtoolbox). A 5-by-5 neural network was initiated with hexagonal lattice structure. The input was the log-2 fold-change in protein synthesis following tunicamycin
treatment relative to DMSO controls for each time point. The network was randomly initiated and used Euclidean distance as the metric for classifying proteins to specific neurons. The SOM algorithm was repeated 1,000 times, and a co-clustering map was generated indicating the frequency with which any two proteins clustered in the same neuron. This co-clustering map was then subjected to hierarchical clustering using Euclidean distance as the metric for clustering proteins.

**K-Means Clustering Analysis**

K-means clustering was used to group proteins from EGF experiments into clusters with distinct temporal responses. Data was filtered by removing proteins that appeared in less than two out of four replicates, as well as removing proteins that did not show a statistically significant change (according to Student’s t-test) in synthesis between EGF and PBS controls in at least one time point. After plotting the within-cluster distance against number of clusters, six clusters were selected for analysis, as increasing the number of clusters above six only marginally decreased the within-cluster distance. The input was log-2 fold-change in protein synthesis for all time points normalized to the 30 minute PBS control. Cluster centroids were initialized randomly, and Pearson correlation was used as the distance metric. K-means clustering was repeated 10,000 times, and a co-clustering map was generated indicating the frequency with which any two proteins shared the same cluster. This co-clustering map was then subjected to hierarchical clustering using Euclidean distance as the metric for clustering proteins.

**Analysis of Temporal Delay Between RFP and MITNCAT Datasets**

To prevent the analysis of random fluctuations in proteins with unchanging RFP values or protein synthesis rates, only proteins with at least one statistically significant time point ($p<0.05$) in both the RFP and MITNCAT datasets were considered, and thus only proteins present in at least two of the four MITNCAT replicates were included in the analysis of temporal delay. These
restrictions limited the protein data set to 90 and the RFP data set to 400; the overlap between these data sets was 27. Log2 PBS normalized values were centered around the mean and normalized to the standard deviation across time points. Because RFP analysis considers only 30, 60, and 90 minute time points whereas MITNCAT examines 30, 60, 90, 120, and 150 minute time points, MITNCAT values were standardized using the mean and standard deviation of the first three time points. Proteins were manually classified as “delayed” based on the relationship between the RFP and protein synthesis rate curves.

**Codon Analytics**

Human full-length open reading frames (ORFs) were obtained from the Mammalian Gene Collection (https://genecollections.nci.nih.gov/MGC/) [1]. Gene specific codon usage data was obtained for 32,751 human coding sequences using a described algorithm [5], which was previously used on yeast, rat and mouse genes and transcripts [6,7]. Briefly, human ORFS were computationally validated to ensure they contained start and stop codons and designated as gene sequences. Next all gene sequences were individually read from start to stop codon. The number of times each of the 64 possible in frame codons was used in each gene was recorded and used to determine gene specific codon frequencies, with the frequencies for all codons for a specific amino acid in a gene adding up to 1.00. Genome values for each codon frequency and standard deviation values were then obtained using data from the 32,751 analyzed genes. Codon usage frequencies from the “delayed” translation set were centered around the genome average and then averaged to calculate an average deviation for the set. The same calculation was performed for randomly generated sets when determining statistical significance (see Statistical Methods).

To determine A-site enrichments for each codon, the codon in the A-site for each ribosome protected fragment (RPF) read was determined. After considering all read lengths from 24 to 40 nt, we found that the majority of our RPF reads were 30-35 nt long. Further, only reads between 30 and 35 nt long showed a strong enrichment for the triplet periodicity common in ribosome
profiling experiments. For these reasons, only reads of these lengths were considered for all RPF analyses. For these reads, the P site codon is at nucleotides 14, 15 and 16. The A-site is therefore at nucleotides 17, 18 and 19. We only considered reads where the A-site codon was in the reading frame of the coding sequence (approximately 60-65% of all reads). For each codon, the frequency with which it was in the A-site was recorded. The A-site frequency was then compared to the null expected frequency. This expected frequency was the frequency of each codon in the longest open reading frame of each gene weighted by the abundance of the gene in the ribosome profiling dataset. Enrichments were calculated as the observed frequency of a codon in A-sites compared to this expected background frequency. These enrichments were then calculated for every codon on 500 random subsets of 100 genes each. Based on their abundance in the 17 genes that displayed delayed increases in protein abundance (Figure 6B, upper right quadrant), codons were separated into “enriched” and “nonenriched” classes. The median A-site enrichment across codons in each class was calculated for each random subset.

Statistical Methods

For MITNCAT experiments, Student’s t-test was used to calculate p-values comparing treated and control samples at all time points in GraphPad Prism. For RNA-Seq and RFP experiments, p-values were generated via the Wald Test using DESeq2. Translational Efficiency (TE) p-values were calculated using the Xtail algorithm [3]. Statistical significance was assigned for \( p < 0.05 \). For codon bias analysis, empirical p-values were calculated using the random permutation test. Briefly, 1e6 random sets of genes with the same size as the query set were generated, and the codon frequency usage was calculated for that set. The p-value was calculated as the fraction of random sets with a more extreme codon frequency than the query set. Significance was assigned if \( p \) was less than the Bonferroni corrected \( \alpha = 7.81 \times 10^{-4} \) (corresponding to an expected FWER of 0.05 across the 64 unique codons). GO term enrichment was performed using the PANTHER classification system version 12.0 [8]. Cluster members were
queried against the background of all proteins included in the clustering analysis. P-values were generated from PANTHER.

Supplemental Methods References

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