An RNA structure-mediated, posttranscriptional model of human α-1-antitrypsin expression

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Chronic obstructive pulmonary disease (COPD) affects over 65 million individuals worldwide, where α-1-antitrypsin deficiency is a major genetic cause of the disease. The α-1-antitrypsin gene, SERPINA1, expresses an exceptional number of mRNA isoforms generated entirely by alternative splicing in the 5′-untranslated region (5′-UTR). Although all SERPINA1 mRNA isoforms encode exactly the same protein, expression levels of the individual mRNA isoforms vary substantially in different human tissues. We hypothesize that these transcripts behave unequally due to a posttranscriptional regulatory program governed by their distinct 5′-UTRs and that this regulation ultimately determines α-1-antitrypsin expression. Using whole-transcript selective 2′-hydroxyl acylation by primer extension (SHAPE) chemical probing, we show that splicing yields distinct local 5′-UTR secondary structures in SERPINA1 transcripts. Splicing in the 5′-UTR also changes the inclusion of long upstream ORFs (uORFs). We demonstrate that disrupting the uORF results in markedly increased translation efficiencies in luciferase reporter assays. These uORF-dependent changes suggest that α-1-antitrypsin mRNA expression levels are controlled at the posttranscriptional level. A leaky-scanning model of translation based on Kozak translation initation sequences alone does not adequately explain our quantitative expression data. However, when we incorporate the experimentally derived RNA structure data, the model accurately predicts translation efficiencies in reporter assays and improves α-1-antitrypsin expression prediction in primary human tissues. Our results reveal that RNA structure governs a complex posttranscriptional regulatory program of α-1-antitrypsin expression. Crucially, these findings describe a mechanism by which genetic alterations in noncoding gene regions may result in α-1-antitrypsin deficiency.

Significance

Protein and mRNA expression are in most cases poorly correlated, which suggests that the posttranscriptional regulatory program of a cell is an important component of gene expression. This regulatory network is still poorly understood, including how RNA structure quantitatively contributes to translational control. We present here a series of structural and functional experiments that together allow us to derive a quantitative, structure-dependent model of translation that accurately predicts translation efficiency in reporter assays and primary human tissue for a complex and medically important protein, α-1-antitrypsin. Our model demonstrates the importance of accurate, experimentally derived RNA structural models partnered with Kozak sequence information to explain protein expression and suggests a strategy by which α-1-antitrypsin expression may be increased in diseased individuals.

Human α-1-antitrypsin is of particular clinical interest because deficiencies in this protein are associated with chronic obstructive pulmonary disease (COPD), liver disease, and asthma (1–4). Smoking is the major environmental factor that contributes to COPD risk, although the inconsistency of COPD rates among smokers points to additional genetic factors that modulate risk (5–7). Multiple genetic variants in the gene encoding α-1-antitrypsin, SERPINA1, cause the disease α-1-antitrypsin deficiency (8–10), which can result in COPD, liver failure, and inflammatory conditions like panniculitis, vasculitis, and glomerulonephritis (9, 11, 12). α-1-Antitrypsin is a protease inhibitor that specifically targets neutrophil elastase, which is present at chronic low levels in the lungs (1). Deficiency of α-1-antitrypsin thus results in higher levels of neutrophil elastase, which in turn degrades elastin (especially in the lungs), resulting in COPD (13). Thus, the role of SERPINA1 in COPD etiology is well described at the protein level; however, little is known about SERPINA1 at the transcript level and whether alteration of potential posttranscriptional controls can contribute to α-1-antitrypsin deficiency and ultimately COPD. Genome-wide association studies identified COPD-associated variants that map to the SERPINA1 untranslated regions (UTRs), introns, and promoter region (5, 14). Furthermore, genetic variants shown to alter SERPINA1 splicing patterns were identified in the SERPINA1 introns of patients with COPD (15, 16). The presence of disease-associated variants in noncoding regions suggests that posttranscriptional regulation of SERPINA1 mRNA is an important component of disease risk. Nevertheless, variants in noncoding regions of SERPINA1 comprise only a small fraction of its disease-associated variants discovered to date, which may reflect the tendency of variant discovery studies to focus exclusively on coding exons (10, 17).

Several features of SERPINA1 emphasize the importance of its transcripts and their regulation. The SERPINA1 gene is exceptionally complex; 11 different splicing isoforms occur in human tissues (18). While alternative splicing occurs in 95% of human multixen genes (19, 20), the 11 SERPINA1 transcripts are extreme, placing SERPINA1 in the top 0.5% of human genes in terms of transcriptional complexity (18). A particularly salient feature of SERPINA1 alternative splicing is that all variants differ only within their 5′-UTRs (21). Therefore, all SERPINA1 mRNA isoforms code for the same α-1-antitrypsin protein; however, their translation efficiency | RNA secondary structure | uORFs | SERPINA1 | α-1-antitrypsin deficiency

Data deposition: The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih.gov/geo (accession no. GSE61520). SHAPE data are also available in SRA format at https://trace.ncbi.nlm.nih.gov/TRACE/traceศาสพเษดส์/9R_Rp8Toro1-UEmK-ocdfpGMMDYyntt1ALuaATX7YqB2Aedit?hap=drive_web.

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differing 5′-UTRs likely determine transcript-specific differences in posttranscriptional processes such as mRNA translation efficiency, subcellular localization, and stability (22, 23). Importantly, the SERPINA1 transcript isoforms are differentially expressed across tissue types (24), suggesting that posttranscriptional regulatory mechanisms adjust α-1-antitrypsin production based on the transcripts expressed in each tissue. The presence of up to three upstream ORFs (uORFs) in the SERPINA1 5′-UTRs (25, 26) suggests a potentially important yet unstudied mechanism for the translation efficiency regulation of these transcripts. In addition to the sequence-based differences between SERPINA1 transcripts, RNA secondary structure differences in the 5′-UTR could also determine their regulation (27–29).

We propose here that noncoding features of SERPINA1 transcripts make up a posttranscriptional regulatory program that ultimately determines α-1-antitrypsin expression. We describe a complex interplay between alternative splicing and translation efficiency mediated by uORFs and RNA structure, which together control tissue-specific expression of α-1-antitrypsin in humans. Our quantitative and predictive model reveals an important and overlooked aspect of α-1-antitrypsin deficiency and suggests RNA-based targets for therapeutic consideration.

Results

Transcript Complexity in SERPINA1. As a clinically important gene harboring numerous COPD and α-1-antitrypsin deficiency-associated variants (30) (Fig. 1A), SERPINA1 is of additional interest for the exceptional number of transcript isoforms it produces. Two transcription start sites (TSSs), six splicing donor (SD), and three acceptor (SA) sites yield a total of 11 transcript isoforms (21) (Fig. 1A and and Fig. S1), which places SERPINA1 in the top 0.5% of transcriptionally complex human genes (18). Remarkably, all of the alternative splicing occurs in the 5′-UTR of SERPINA1 mRNA (Fig. 1A). Thus, in healthy adults, α-1-antitrypsin exists as a single protein isoform that is produced from 11 different mRNAs. We sought to determine whether the mRNAs are functionally different and how any differences relate to α-1-antitrypsin production or deficiency.

We therefore began this investigation by quantifying the expression of the various SERPINA1 transcripts in human tissues. Using data from the Illumina BodyMap 2.0 transcriptome-wide RNA-seq project, we quantified the relative amount of total SERPINA1 transcripts in 16 human tissues (Fig. 1B) and show the relative amount of each SERPINA1 transcript in the form of a heat map (Fig. 1C). There are clear differences in the total amount of SERPINA1 present in each tissue. Liver noticeably yields the highest total SERPINA1 read counts (Fig. 1B), reflecting that α-1-antitrypsin is primarily expressed by hepatocytes and secreted into the bloodstream (1, 31). While the lungs are thought to acquire α-1-antitrypsin from the bloodstream (1, 32), we found that lung tissue transcribes non-trivial amounts of SERPINA1 (Fig. 1B), thus potentially producing its own α-1-antitrypsin. Although some SERPINA1 transcript isoforms are more prevalent than others, we detected all of the transcripts, with some tissues like liver expressing every transcript (Fig. 1C). To verify these findings with greater specificity, we designed SERPINA1 5′-UTR-specific primers and amplified RNA extracted from liver and lung epithelial cells (HepG2 and A549 cell lines, respectively). All 11 transcripts were expressed in HepG2 cells, and all save 1 in A549 cells (Fig. 1C, Bottom). The varied expression of the SERPINA1 transcript isoforms across the tissues suggests that each SERPINA1 transcript has a distinct posttranscriptional function. Given that these transcripts vary only in their 5′-UTR, we hypothesize that the splicing complexity in the 5′-UTR of SERPINA1 plays an important role in its posttranscriptional regulation, especially, as detailed next, in SERPINA1 mRNA translation.

Translation Efficiency Analysis. The 5′-UTR in an mRNA regulates translation of the coding sequence and ultimately controls the expression of protein products (22). To test the effect of different SERPINA1 5′-UTRs on mRNA translation, we measured the translation efficiencies of six representative SERPINA1 5′-UTRs with luciferase assays. Strikingly, we found significant differences in translation efficiency for the six SERPINA1 5′-UTRs (Fig. 2A). Alternative splicing determines the inclusion (or exclusion) of up to three uORFs in the final SERPINA1 transcript isoform (26) (Figs. 1A and 2B, and Fig. S1). Because uORFs can affect translation efficiency (33, 34), the uORFs in SERPINA1 may modulate translation of the different transcripts [an idea acknowledged decades ago (25) but untested until now]. To evaluate the effect of uORFs on SERPINA1 translation, we mutated the start codon of a single uORF in each luciferase construct from “AUG” to “AAG” (Fig. 2B). In this group of mutants, we mutated every possible SERPINA1 uORF in at least one construct. Although it is possible that translation initiation at the mutated start codons could still occur (35), the initiation efficiency of an “AAG” start codon is very low—between 0 and 3% (36).

Mutating the uORF start codon(s) resulted in large increases in the translation efficiency of three of the six transcripts (Fig. 2C), suggesting that these uORFs typically inhibit translation. The three transcripts with inhibitory uORFs are NM_001002235, NM_001002236, and NM_001127705, and their mutated uORFs were uORFγ, uORFδ, and uORFγ′, respectively (Fig. 2B). Interestingly, uORFγ is too close to the mRNA 5′-terminus to be translated based on canonical understanding of translation initiation (37). However, our luciferase assays clearly suggest that it is functional, as it both significantly represses translation of...
To control for potential inaccuracies in the Kozak sequence prediction efficiency (39, 40). Indeed, uORF and uORF_E10246 relative to the luciferase assay control. The number of uORFs in each transcript measured by luciferase reporter assays. Replicate TE values are shown as gray squares in Table S1. The inhibitory uORF scripts (Fig. 2A) predicted with a leaky-scanning model of translation (Eq. 1). The leaky-scanning model moderately predicted the translation efficiencies given by our luciferase assay data ($r^2 = 0.40$, Table S1). Another factor that could affect translation efficiency is reinitiation after uORF translation (42). We therefore fit a “reinitiation leaky-scanning” model (Eq. 2) to the experimental translation efficiencies, but observed no improvement ($r^2 = 0.33$, Table S1). The rules that govern uORF reinitiation are admittedly poorly understood (43). It is possible that uORFα, which has a strong Kozak sequence (Fig. 2B and Dataset S1), nevertheless fails to inhibit coding sequence (CDS) translation (Fig. 2) due to efficient reinitiation after uORFα translation. Adjusting uORFα Kozak strength in the leaky-scanning model to reflect this idea, we observe a moderate improvement in fit ($r^2 = 0.60$, Table S1). An additional factor that can modulate translation efficiency is mRNA secondary structure. Evidence for the effect of secondary structure on translation has been conflicting (27–29, 34, 40), but such studies have typically relied on theoretical structure prediction, which falls far short of the accuracy achieved with direct chemical probing experiments (44, 45). We next sought to predict translation efficiency for the SERPINA1 transcripts using secondary-structure features derived from chemical structure probing.

**Secondary Structure of SERPINA1 Transcripts.** Recent advances in RNA structural mapping techniques, in particular selective 2'-hydroxyl acylation by primer extension and mutational profiling (SHAPE-MaP) (46), have enabled accurate, high-throughput, whole-transcript structural interrogation of RNA (47–49). SHAPE-MaP interrogates the reactivity of each 2'-hydroxyl in an RNA toward the reagent 1-methyl-7-nitroisatoic anhydride where the relative reactivity estimates the tendency of each nucleotide to be structured (i.e., base paired) or unstructured (i.e., unpaired). To measure structure differences between the SERPINA1 transcripts, we performed SHAPE-MaP separately on the six SERPINA1 transcript isoforms whose 5'-UTRs were examined in luciferase assays. The resulting data are highly correlated between replicates (Fig. S4), with average correlation coefficients of 0.89 or more. Our experimental SHAPE-MaP data provide SHAPE reactivity profiles at nucleotide resolution for each of the six SERPINA1 transcripts. Regions with lower median SHAPE values (low SHAPE reactivities) consist of largely unreactive nucleotides (Fig. 3A), whereas regions with higher median SHAPE values indicate the reverse (Fig. 3B). The median-centered SHAPE reactivities of each transcript illustrate the relative reactivity of regions in the transcripts and indicate structured regions (Fig. 3C). The high-reproducibility of SHAPE-MaP is immediately apparent in the median-centered SHAPE profiles: the reactivity patterns in the coding sequences (CDS) are nearly identical across the six transcripts, corresponding to the transcripts’ identical CDS sequences (Fig. 3C and Fig. S3). In addition, shared exons in the 5'-UTRs also exhibit comparable SHAPE reactivities despite existing in unique contexts in the different transcripts (Fig. S5).

We next used our SHAPE-MaP data to derive minimum free-energy structure models for the six SERPINA1 transcripts (44, 50). SHAPE reactivities were incorporated as pseudo-free-energy terms to guide RNA structure modeling with RNAfold (51). Importantly, this approach has been extensively validated and generally yields structure models with accuracies above 90% (44, 46, 51, 52). Even in the case where there is SHAPE data for the entirety of an RNA (as is common at the ends of transcripts), incorporating available SHAPE data still greatly improves the accuracy of structure predictions (53, 54). As an internal control, we initially compared the structure...
models derived for the CDS regions of different transcripts. Consistent with the high correlation observed between SHAPE-MaP profiles, the secondary-structure models are highly similar in the CDS regions, supporting the robustness of the models (Fig. S6). However, we were most interested in the structures around the uORFs and the beginning of the CDS, and how this information could be used to model translation efficiency.

**Modeling Translation Efficiency with Structure.** We next sought to gain a quantitative understanding of the contribution of RNA structure to translation. The interplay of transcript structural elements with the translational machinery is not well understood, although studies in bacterial and mammalian systems suggest that secondary structures near start codons are most likely to affect mRNA translation (27, 28, 55, 56). We established above that the uORFs in SERPINA1 affect translation efficiency and found that a model that incorporates only Kozak sequence strength did not quantitatively explain a large portion of the translation efficiency differences (Fig. 2D). We hypothesized that, in addition to Kozak sequence strength, the model requires structural data encompassing the Kozak sequence to accurately capture the probability of the ribosome initiating at a given ORF. SHAPE-MaP data provided us with a high-confidence structure of each transcript (Fig. 3D and Fig. S6), including the structures surrounding each Kozak sequence (Fig. 4A and B and Fig. S5). Studies in prokaryotes suggest that translation initiation occurs in proportion to the exponent of the free energy ($\Delta G$) of unfolding of the local structure (57), which is the energy required to “unfold” a region of RNA (and is thus a positive value). We modified the leaky-scanning model from Eq. 1 to include the $\Delta G$ of unfolding around the Kozak sequence (Methods and Eq. 3). The SHAPE data-driven “structure leaky-scanning” model dramatically improves the predictive power of the model to 94% (Fig. 4C). The structural terms in the model weigh each Kozak sequence by its accessibility in addition to its strength. From their location in uORF secondary structures, it is immediately clear that not all of the uORF Kozak sequences are equally accessible (Fig. 4A and B). For example, the Kozak sequence for uORF6 resides in a single-stranded loop, while the Kozak sequence for uORF4 is engaged in a base-paired stem structure. It appears that uORFs $\delta$, $\delta'$, and $\gamma$ are the only uORFs that have a Kozak sequence that is both strong and structurally accessible (Figs. 2C and 4A and B; and Fig. S7), potentially explaining why only these uORFs inhibit SERPINA1 translation in our assays. The specific $\Delta G$ of unfolding associated with each uORF Kozak sequence is important: permitting the $\Delta G$ values’ assignments and refitting the structure leaky-scanning model never produces $r^2$ values reaching 0.94 (value of $P < 0.001$). Furthermore, refitting the structure leaky-scanning model using $\Delta G$ values predicted without SHAPE data yields a lower correlation ($r^2 = 0.79; \text{Dataset S1}$), supporting the importance of using accurate SHAPE-based structure models. Finally, we also varied the size of the unfolding region around the Kozak sequence used for calculating $\Delta G$ of unfolding values. Supporting the physical relevance of our structure leaky-scanning model, the optimal predictive power was obtained for an unfolding window size of 30 nt, consistent with the known size of the eukaryotic ribosomal footprint (58, 59). Either smaller or larger unfolding regions exhibited significantly worse agreement with the translation efficiency data (Table S2).

It is important to note that our structure models for the SERPINA1 transcripts focus on local structures (Methods). Although long-range interactions in large RNAs can occur, local structure is thought to dominate the folding of mRNAs (60, 61). To explore the possibility of longer-range secondary structures, we recalculated the $\Delta G$ of unfolding values, allowing for greater pairing distances in RNA structure predictions and refit our structure leaky-scanning model in each case (Table S3). Predictive
performance of the model generally decreases as the max pairing distance increases (Table S3), suggesting that local structure is most important in determining translation of these transcripts (Table S3). However, we cannot exclude the possibility of long-range interactions. While a few recent structure probing methods can directly detect long-range interactions (62, 63), SHAPE-directed modeling accuracy decreases for long-range interactions, which could also contribute to the decreased performance as max pairing distance increases.

Based on this analysis, we propose that the Kozak sequence determines the likelihood of initiating translation, but the secondary structure determines whether the Kozak sequence can in fact be accessed. Thus, the translation efficiency of each SERPINA1 transcript is a combination of the initiation strength and structure of its CDS Kozak sequence, attenuated by the translation efficiency of any uORFs as governed by the same parameters.

**Mutating Secondary Structure to Change Translation Efficiency.** While our luciferase assays suggest that little to no translation occurs at uORFα (Fig. 2), available ribosomal profiling data (64) show minimal yet detectable signal at uORFα (Fig. S8). This indicates that uORFα is capable of translation and is thus capable of being translationally regulated, including by structural manipulation. To further assess the role of secondary structure in controlling uORF function, we designed structure mutants for uORFα (in transcript NM_001002235.2). We designed three mutants with low free energies of unfolding to be predominantly single stranded within 30 nt of the uORF Kozak sequence. Mutants contained altered sequences upstream and downstream of the Kozak sequence, without altering the Kozak sequence itself. The wild-type structure of uORFα has an unfolding energy of 22.4 kcal/mol, while the three mutants have unfolding energies below 4 kcal/mol and are expected to enhance the activity of uORFα, thus diminishing translation of the CDS. The translation efficiency of each structure mutant was measured by luciferase assays relative to wild type. As expected, the mutants show reduced translation efficiencies (of the CDS) relative to wild type (Fig. S4) that are consistent with predictions from the structure leaky-scanning model (Fig. 5B).

In transcript NM_001002236.2, which contains uORFβ, we designed four mutants with increased energies of unfolding within 30 bases of the uORF Kozak sequence and an additional mutant with greatly decreased ΔG of unfolding. As with uORFα, a reduction in the ΔG of unfolding causes a reduction in overall translation efficiency relative to wild type (Fig. 5C and Fig. S9). Conversely, increasing the ΔG of unfolding around uORFβ increases translation efficiency in one structure mutant, but as the structure-mutant energies of unfolding increase above ~25 kcal/mol, overall translation efficiency begins to decrease (Fig. S9). These results could be interpreted to indicate that the ΔG of unfolding is not a significant factor controlling uORFβ translation initiation. Alternatively, the overall decrease in translation efficiency as hairpin sizes increase exactly replicates multiple experiments in which hairpins of increasing size added to the 5' UTR progressively reduce translation efficiency (39, 65-67). Thus, increasing the ΔG of unfolding around a uORF may increase overall translation efficiency up to a point (56), beyond which strong secondary structures begin to impede ribosomal scanning altogether (65).

Overall, our data from SERPINA1 wild type, uORF mutant, and structure mutant luciferase assays strongly support that the ΔG of unfolding around the Kozak sequence is an important determinant of translation efficiency. Including the additional structure mutants, the leaky-scanning model moderately predicts translation efficiencies (r² = 0.55), but most of the variation in translation efficiency is explained by the structure leaky-scanning model (r² = 0.83). Changing the ΔG of unfolding around a single Kozak sequence in a given 5'-UTR leads to changes in translation efficiency that are well predicted by the structure leaky-scanning model, but unanticipated by the leaky-scanning model (Fig. 5 C and D).

**Modeling α-1-Antitrypsin Expression in Tissue.** A goal of transcriptomics is to develop models that accurately describe transcript dynamics and expression in living tissue. As we have seen from tissue-specific transcriptome data, SERPINA1 transcription is not limited to the liver, and different tissues express different combinations of the SERPINA1 transcript isoforms. Assuming the overall translation efficiency in each tissue as the ratio of α-1-antitrypsin protein produced in each tissue. Based on available protein quantification data (68), we calculated the overall SERPINA1 translation efficiency in each tissue as the ratio of α-1-antitrypsin protein to SERPINA1 transcript totals. If the translation efficiency of SERPINA1 mRNA were equal in every tissue, then we expect to observe that α-1-antitrypsin amounts and total SERPINA1 transcript amounts are correlated. However, we observed no such correlation (Fig. 6A), indicating that different tissues have different net α-1-antitrypsin translation rates, potentially due to their unique combinations of SERPINA1 transcript isoforms. Assuming that the overall translation efficiency in a tissue is the average of the translational efficiencies of all its SERPINA1 transcripts weighted by abundance, we can use the two scanning models described above to predict SERPINA1 translation efficiency in tissues (Eq. 4). While our luciferase assays show uORFα to repress translation in transcript NM_000295.4 (Fig. 2C), it is likely that this uORF is not functional in vivo given its close proximity to the transcript 5' termini and lack of a canonical translation initiator of short 5'-UTR (TISU) sequence (37). Indeed, when we assessed our ability to model translation efficiency treating the uORFα as functional (Fig. S10) or nonfunctional (Fig. 6B and C), the nonfunctional assumption yielded better prediction of
tissue-specific translation efficiencies. The leaky-scanning model of translation (Fig. 2D and Eq. 1) explains 59% of the variation in translation efficiency between tissues (Fig. 6B), whereas the structure leaky-scanning model (Fig. 4C and Eq. 3) explains 66% (Fig. 6C). The addition of RNA structural data to the model of translation thus improves predictions of translation efficiency in human tissues.

Discussion

The amount of protein produced from a gene is not a simple function of the abundance of the transcript (69, 70). The complex path between transcript expression and protein expression is often a missing link in our understanding of cellular phenotype, indicating a need for integrative models that bridge this divide. **SERPINA1** is exemplary of the effects of posttranscriptional regulation on protein output. While each of the **SERPINA1** transcripts produces the same protein isoform, they do so with different translation efficiencies. Differences in uORF content and 5′-UTR secondary structure combine to differentiate the translational efficiencies of **SERPINA1** transcripts. Secondary structure plays a surprisingly important role in accounting for these differences, and in determining the repressive effect of individual **SERPINA1** uORFs. When considering the role of secondary structure in a system, correctly defining an RNA secondary structure demands more than a cursory computational prediction. Structural data accurate enough for successful biological models require comprehensive chemical or enzymatic probing of the RNA molecules of interest (46-48). Previously, no correlation was found between secondary structure and translation rate in experiments that measured the protein expression of constructs with varied uORF or CDS Kozak sequences (34, 40). In these studies, it is likely that the purely computational RNA structure models were inadequate for predicting structures around Kozak sequences. In this study, we used SHAPE-MaP chemical probing to successfully improve mRNA translation efficiency predictions (44) (Figs. 2D, 4C, and 5). While this model aptly describes the translation of **SERPINA1** transcripts, additional experiments measuring the translation efficiencies of simultaneous uORF and structure mutants are necessary to determine the contribution of secondary structure in more detail. Additionally, a more generalizable model of translation efficiency will require modifications to capture additional factors that regulate translation, including overlapping uORFs, reinitiation after uORF translation (43), non-AUG uORF translation (71), and 5′ cap secondary structure (65).

Transcript-specific translation efficiencies may play an important role in tissue-specific protein expression, especially in the case of α-1-antitrypsin, which shows a complex and varied expression pattern across human tissues. However, overall α-1-antitrypsin output in a tissue is not solely a consequence of translation efficiency. Transcripts travel through a coordinated posttranscriptional program, or “regulon” (72), and may diverge from their fellow isoforms at each step. Tissues could also have different overall rates of translation (for example, in a fast- versus slow-growing tissue) or have different rates of protein export.

These additional layers of regulation likely explain why our model of translation efficiency performs better in tissue culture cells than in tissues. However, our model still provides insights into the regulation of α-1-antitrypsin expression in tissues. First, liver tissue is a considerable outlier in both models of **SERPINA1** translation efficiency (Fig. 6B and C). Interestingly, predicting much higher translational efficiencies in liver tissue than observed based on measured levels of α-1-antitrypsin is consistent with the understanding that liver exports most of its α-1-antitrypsin into the bloodstream (1, 31, 73). This artifact indicates a need for tissue-specific cellular import/export dynamics to inform models of protein expression. Conversely, the models predict translation efficiency in lung tissue fairly accurately, suggesting that translation of **SERPINA1** mRNA is a major source of its α-1-antitrypsin, which contrasts the paradigm that lung tissue derives its α-1-antitrypsin from the bloodstream (1, 32). Our detection of **SERPINA1** transcripts in cultured lung cells (A549 cells; Fig. 1C) and recent quantification of **SERPINA1** transcripts in lung tissue (24) further support the conclusion that cells in the lung itself express α-1-antitrypsin. This surprising conclusion contradicts current models of the role of α-1-antitrypsin in disease. The most common genetic variant in **SERPINA1** associated with COPD and α-1-antitrypsin deficiency, the Pi*Z allele, is thought to cause α-1-antitrypsin to be poorly exported from the liver, leading to deficient α-1-antitrypsin levels in the lungs and eventual neutrophilic overload (1, 8, 32). If lung tissue produces its own α-1-antitrypsin, however, then this disease model is likely incomplete. Instead, disease-associated variants must also impact α-1-antitrypsin levels in lung tissue, either by producing unviable α-1-antitrypsin or reducing its translation. For example, genetic variants could reduce α-1-antitrypsin production if they shift **SERPINA1** transcription to isoforms with the lowest translation efficiencies. A recent study quantified α-1-antitrypsin and different **SERPINA1** transcripts in the serum of α-1-antitrypsin deficiency patients and healthy controls to determine whether patients have different combinations of the transcripts (24). Unfortunately, the primer design in that study did not differentiate between the transcripts with the lowest and highest translational efficiencies, but the data did show a change in transcript proportions for at least one patient population (24).

Ultimately, COPD in α-1-antitrypsin deficiency is caused by the diminished levels of α-1-antitrypsin. Current therapies attempt to deliver donor serum-derived α-1-antitrypsin i.v. to affected individuals, but this treatment is costly and of unknown efficacy (74).
Our work suggests a therapeutic strategy: α-1-antitrypsin levels could be increased in situ, perhaps with antisense oligonucleotides (ASOs) that target the Kozak sequences around the uORFs in SERPINA1 transcripts, as shown recently for other uORF-containing mRNAs (56). ASOs would likely act as double-stranded regions that increase the ΔG of unfolding around uORF Kozak sequences, blocking the uORFs in SERPINA1 transcripts and increasing in situ α-1-antitrypsin expression. Our findings illustrate the importance of the numerous SERPINA1 transcript isoforms and their translation in disease and the impact of posttranscriptional regulation and secondary structure on phenotype in general.

Materials and Methods

SERPINA1 Annotation. The known SERPINA1 transcript annotations were taken from RefSeq, version hg38. In each transcript, uORFs are defined by a start and stop codon in the same frame within the 5′-UTR. Distinct uORFs are named here with the Greek letters α, β, γ, δ, ε, and ζ.

Heat Map of Tissue-Specific Isoform Expression. Pair-ended RNA-seq reads from 16 different tissues were downloaded from the Illumina BodyMap 2.0 project [Gene Expression Omnibus (GEO) accession number GSE30611]. Abundance estimates of the 11 known SERPINA1 transcripts were quantified with Sailfish, version Beta 0.7.6 (75), using the full human transcriptome (RefSeq, version hg38) as the reference. Estimates of total SERPINA1 expression in each tissue were calculated as the sum of transcripts per million (TPM) of each transcript. For better visualization in Fig. 1 and 2, the total expression in liver was adjusted by constant to 106, and all other tissues’ TPM estimates of each transcript. For better visualization in Fig. 1 transcripts: NM_001002235.2, NM_000295.4, NM_001127700.1, and SERPINA1 we built six luciferase constructs containing 5′-UTRs of each transcript. For luciferase assays, Luciferase Assays.

Luciferase Assays. To assess the translation efficiency of SERPINA1 transcripts, we built six luciferase constructs containing 5′-UTRs from six selected SERPINA1 transcripts: NM_001002235.2, NM_000295.4, NM_001127700.1, NM_001127700.4, NM_001127705.1, and NM_00010236.2. The 5′-UTRs were cloned into a modified pGL3 that minimizes the amount of plasmid 5′-UTR in the product. For each of the SERPINA1 and control constructs, 0.5 μg of plasmid was transfected into HEK cells. Cells were harvested with Cell Culture Lysis Reagent (Promega; E153A) 24 h posttransfection. Luciferase activity of the samples was measured by Luciferase Assay Substrate (Promega; E151C) and Luciferase Assay Buffer (Promega; E152B) with a luminometer (Molecular Devices). Luciferase activity measurements were taken in duplicate and averaged for each sample. The luciferase activity measurement for each sample was normalized to total sample protein concentration, as determined by Bradford assay (n = 4), and reported in Dataset S1. Luciferase measurements were further normalized to the abundance of luciferase RNA in each sample to obtain (luciferase activity)/luciferase RNA, as described previously (76). To quantify luciferase RNA abundance, after measuring luciferase activity, total RNA was extracted with TRIzol. Samples were depleted of DNA with Ambion Turbo DNA-free (AM1907) and reverse transcribed with High Capacity cDNA Reverse Transcription Kit (Applied Biosystems; 4368814). Luciferase and GAPDH cDNAs were quantified by real-time PCR (qRT-PCR) on a Bio-Rad CFX96 Real-Time System. Luciferase and GAPDH primers used were 5′-ACAAAGGCTATACGGTGGCT-3′ (forward), 5′-CGTGTCCCTACCAAAACAAA-CAGC-3′ (reverse), and 5′-CTGTTGCTGTACCAAAATTTGCT-3′ (forward), respectively. Luciferase RNA abundance was determined by the ΔΔCT method (n = 4). All (luciferase activity)/luciferase RNA measurements are reported relative to an empty vector control to correct for systematic variations between experiments.

uORF Mutants. To disrupt uORFs in the original six SERPINA1 plasmid constructs, we designed primers to start the codon of selected uORFs from AUG to AAG using the NEB QS site-directed mutagenesis kit. uORFs and uORFδ were mutated in NM_001002236.2 and NM_001127705.1 luciferase plasmids using primers uORFT4A1F: CCGAGTACAAAGAAGCCTTCUCCG4SAR: CCTA-GGACCCCTGTCGTCA. uORFδ was mutated in NM_001127705.1 luciferase plasmids using primers uORFT285AF: ACTACGATAAGGATAGTCTCTGTCATGJTUB85AR: CACCCAAAATGCTCTAGT. uORFs were mutated in NM_001002235.2 and NM_001127700.1 luciferase plasmids using primers uORFT32AF: CCGGACCG-GCAAGCTTGGCTCGT32AR: ACAGTCGACCGTCTCGGC. uORFε was mutated in NM_000295.4 luciferase plasmid using primers uORF44A: CCGGGCAAG-ACTCTCTTGCTTG44AR: CCTGCCGCTGCTCAAAAT. Mutant constructs were sequenced (Dataset S2). Luciferase assays were performed in the mutant constructs as above, and results are reported in Fig. 2, Fig. 5, and Dataset S1.

uORF Structure Mutants. The structure mutants of uORFs in transcript NM_001002235.2 and of uORFs in transcript NM_001002236.2 were computationally designed by altering sequences adjacent to the uORF Kozak sequences and predicting the resulting change in secondary structure. To design structure mutants, codons in the given uORF were either permuted or mutated with CodonShuffle (77) to preserve dinucleotide frequency and codon usage. To design mutants with increased uORF ΔG of unfolding, sequence upstream of the uORF was changed to complement the new uORF sequence. To design mutants with decreased uORF ΔG of unfolding, sequence upstream of the Kozak sequence was substituted with CAA repeats, using codon adjacency (1 single-stranded structure (SSS) and 2 nt downstream of the AUG) were left unchanged. The ΔG of unfolding was then predicted for all ORFs in the transcript, selecting mutants that exhibited the desired change in uORF ΔG of unfolding without affecting the predicted ΔG of other open read frames. Three mutants were selected for NM_001002235.2 and five for NM_001002236.2. The structure mutant 5′-UTRs were cloned into double digestion into modified pGL3 plasmids and verified by sequencing (Dataset S2) as described above. Luciferase assays were performed as before, except without normalization to luciferase RNA levels. Previous luciferase assay data (wild type and uORF mutant constructs) show a very strong linear correlation (r = 0.95) between luciferase activities alone and luciferase activities normalized by luciferase RNA, indicating that luciferase activity alone is sufficient to estimate translation efficiency for these constructs. Luciferase assays on wild-type NM_001002236.2 were performed in parallel for comparison. Structure mutant luciferase activities adjusted to the scale of RNA-normalized luciferase values according to the following: adjusted luciferase activity = (luciferase activity)*0.20650 + 0.20141. Structure mutant luciferase assay results are reported in Fig. 5, Fig. 59, and Dataset S1.

S-RACE. To characterize the 5′ ends of luciferase construct transcripts, HEK cells were transfected with NM_001002235.2-luciferase were treated with the RLM-RACE kit (Ambion). Briefly, total RNA was extracted from calf intestinal phosphatase, and intact 5′-methylguanosine caps were removed by treatment with tobacco acid phosphatase. The 5′ ends of transcripts were ligated to a linker sequence and primed with random hexamers.
in reverse transcription. cDNA was then amplified via nested PCR with forward primer CTGATACAGCATTTGTGATT and reverse primer CCTGAGTGTCATCCGAGTCA, complements of the 5' linked pool reagent used for each luciferase coding sequence, respectively. PCR products were shot gun cloned into pCRII/Blunt (Invitrogen) and Sanger sequenced using forward primer M13F to determine S-end sequence.

Ribosome Profiling Data. Ribosomal profiling datasets with sufficient coverage over the entire SERPINA1 locus were identified by RPdDB (Eichhorn et al. (64); U2OS cells (78)). Single-end ribosomal profiling sequencing reads were downloaded from SRA (identifiers SRX860698 and SRX860702), trimmed on the 3' end to 26 nt as described in ref. 78, and mapped to the human genome build hg38 by Bowtie2, allowing for multimapped reads. Read coverage mapping to the SERPINA1 transcripts is visualized in Fig. S8.

SHAPE-MaP Sequencing and Analysis. 5'-UTRs and coding sequences of six selected SERPINA1 transcripts were cloned into pBLUNTII using overlap extension PCR and verified by sequencing (Dataset S2). The selected transcripts are the same set analyzed by luciferase assays. Plasmids were named as follows: NM_000295: pAL0108; NM_001002235: pAL0096; NM_001127700: pAL0110 and NM_001127705: pAL0103 and pAL0105. Templates for transcription were amplified from the same set analyzed by luciferase assays. Plasmids were named as follows: NM_000295: pAL0108; NM_001002235: pAL0096; NM_001127700: pAL0110 and NM_001127705: pAL0103 and pAL0105. Templates for transcription were amplified using an Ambion MEGAClear Transcription Clean-up kit (Thermo Fisher) or an RNeasy mini kit (Qiagen). Transcripts were verified using denaturing agarose gel electrophoresis with 2% SEAkem gold agarose and the Amresco Formamide RNA Gel Loading buffer. cDNA was reverse transcribed into SHAPE-MaP library preparation, as described previously (46) with some modifications. Briefly, RNA was diluted in water, denatured at 95 °C for 1 min, and snap cooled on ice. After the addition of freezing buffer (100 mM KCl, –10 mM MgCl2, 100 mM Hepes, pH 8.0, final concentration), the RNA was folded at 37 °C for 10–15 min. Then 45 μL of folded RNA was either mixed with 5 μL of DMSO (untreated control) or 5 μL of 100 mM 1-methyl-1-tris(methylamino)pyridinium iodide (M7) in DMSO (treated sample). After 5 min, reactions were desalted using G25 or G50 columns. A denatured control was performed in parallel in which the RNA was diluted into 50 mM Hepes, pH 8.0, 4 mM EDTA, and 50% formamide, then heated to 95 °C and treated with 5 μL of 100 mM 1M7 in DMSO. After 1 min, reactions were desalted using G25 or G50 columns. The RNA was reverse transcribed using SuperScript II (Life Technologies) and random nonamers followed by cleanup with a G25 or G50 column. The second strand was synthesized using the Northern mRNA Second Strand Synthesis Module (NEB). The double-stranded DNA was then prepared using a Nextera or Nextera XT DNA Library Prep Kit (Illumina). Following DNA library concentration determination via Qubit and analysis by Bioanalyzer, libraries were run on a mSeq (Illumina) and resulting data were analyzed using the ShapeMapper pipeline (46), version 1.2, which calculates the SHAPE reactivity of each nucleotide i as follows:

\[
R_i = \frac{mutr_i - mutr_o}{mutr_o}
\]

where mutr is the mutation rate in the sample treated with the SHAPE reagent, mutr is the mutation rate in the untreated control, and mutr is the mutation rate in the denatured control.

 SHAPE-MaP sequencing data and processed SHAPE reactivity profiles are available in the National Center for Biotechnology Information (NCBI) GEO database accession number GSE81525. SHAPE data are also available in SRNNSM format at https://docs.google.com/spreadsheets/d/1_RpB9t01-U6ekm-o99pGMOYRe-t1IALuA7K7qf98AAD/edit?usp=drive_web. SHAPE-MaP experiments were performed twice for each transcript, and the average of the two replicate publications was used for subsequent analyses requiring SHAPE data. For visualization, the median SHAPE profiles in Fig. 3C were generated for each transcript by calculating the median SHAPE value in windows of 20 bases (step size = 1) and subtracting the global median.

Secondary-Structure Analysis. Each transcript with SHAPE-MaP data was folded with RNAfold, version 2.2.4, incorporating their respective SHAPE data with the –shape option and a max distance of 50 (--maxGapSpan = 50) to model on local structures. SHAPE reactivities were incorporated into structure modeling as pseudo free energies according to ref. 44 using a slope of 1.8 and an intercept of −0.6. The 3'-UTRs were excluded from structure modeling since these regions were not covered by our SHAPE-MaP experiments. Structure models were also generated for the six luciferase constructs, which consist of a specific SERPINA1 5'-UTR followed by 700 bases of luciferase coding sequence. For these luciferase construct models, SHAPE data from the endogenous transcripts were used to restrain the 5'-UTRs; SHAPE data for the luciferase coding sequence are unavailable. ΔG of unfolding measurements were calculated around Kozak sequences in the structure models by removing base pairs that occur within ±15 bases around the “A” in the start codon. The free energy of the “relaxed” structure was subtracted by the free energy of the original structure to arrive at the ΔG of unfolding around the Kozak sequence. (The ΔG of unfolding = −ΔG of folding.) The ΔG of unfolding was calculated around the coding sequence and uORF Kozak sequences in the luciferase constructs when fitting to experimental translation efficiencies, and in the wild-type transcripts when fitting to tissue translation efficiencies (Dataset S1). SHAPE-MaP was not performed on uORF mutant SERPINA1 transcripts, but, because point mutations rarely cause perceptible changes in secondary structure (79, 80), it is assumed that the structures of the wild-type transcripts closely approximate the structures of the uORF mutants. ΔG of unfolding values around SERPINA1 structure mutant uORFs were calculated in the same manner as above, with the exception that the underlying structure models were generated using naive prediction due to the absence of SHAPE-MaP data for structure mutants.

Models. The performance of a number of different translation efficiency models is described in Dataset S1. Predictor(s) were fit with simple linear regression to the (luciferase activity)/(luciferase RNA) measurements of the six SERPINA1 constructs and six uORF mutant constructs. Adjusted r2 values and predictor P values were determined by the lm function in R, version 3.2.3. The models we feature in the results are the leaky-scanning and the structure leaky-scanning model (equations 1 and 3). Eq. 1, the naive translation model, is an expansion of a previously published model (34) to allow multiple nonoverlapping uORFs. Eq. 2 is a rederivation of Eq. 1 that allows for translation reinitiation after uORF translation, dependent on a logistic model that assumes the ribosome has a 50% probability of reinitiating 35 bases downstream from the end of an uORF (42). Eq. 3, the structure leaky-scanning model, is our variation of Eq. 1 that incorporates the ΔG of unfolding around Kozak sequences, as assumed to be exponentially related to ribosomal initiation (57). TE is “translation efficiency”; k, k', and i are constants; and P is the strength of the given Kozak sequence as determined previously (40). Kozak strengths are converted to probabilities by dividing by the maximum Kozak strength, 150. Because uORF order matters in Eq. 2, Pji refers to the Kozak sequence strength of the uORF that is jth closest to the CDS. di refers to the distance between the end of the nth uORF and the beginning of the next ORF. ΔGji in Eq. 3 corresponds to the ΔG of unfolding ±15 bases around the given Kozak sequence (calculation described above). The subscripts of Pji and ΔGji indicate either the coding sequence or the nth uORF, numbered 5'-3' in each transcript. Pji and ΔGji values for a transcript without an nth uORF are simply zero. P and ΔGji values are provided for every transcript’s CDS and uORFs in Dataset S1. Constants k and i were optimized in the structure leaky-scanning model fit to SERPINA1 wild type and uORF mutant (luciferase activity/luciferase RNA) values, and the constant k is the original published value (34): k = 0.86 k' = 0.39 i = 0.037

\[
\begin{align*}
TE & = kP_{cds} \left(1 - kP_i \frac{1}{1 + e^{(1 - kP_j)(1 - kP_k)}}\right) \\
TE & = kP_{cds} \left(1 - kP_i \frac{1}{1 + e^{(1 - kP_j)(1 - kP_k)}}\right) \\
TE & = kP_{cds} e^{\alpha G_{ij}} \left(1 - kP_i e^{\alpha G_{ij}}\right) \left(1 - kP_j e^{\alpha G_{ij}}\right) \left(1 - kP_k e^{\alpha G_{ij}}\right)
\end{align*}
\]
[4]

The predicted model values for tissue translation efficiencies were then fit to the measured tissue translational efficiencies with simple linear regression. Models fit best to the log of the measured tissue translational efficiencies. R-squared values and model P values are reported in Dataset S1. The structure of the SERPINA1 transcript contains alternative 5′ and 3′ UTRs, as well as alternative splicing in intron 1. The α1-antitrypsin protein is encoded by the SERPINA1 gene on chromosome 14q32.3. The SERPINA1 translation efficiency in each tissue was measured by dividing the α1-antitrypsin protein concentration by the total SERPINA1 concentration. To predict tissue translation efficiencies with a given model, the model’s translation efficiency estimates of all 11 SERPINA1 transcripts were used to predict the average translation efficiency of each tissue as in Eq. 4. $TE$ is the model-predicted translation efficiency of tissue $\Delta G$, and $m_l$ is the function for the translation efficiency of transcript $i$ with parameters from fitting the model to the luciferase data:

$$TE_j = \sum_i (m_l(i) \times TP_{III}) \div \sum_i TP_{III}.$$
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