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Correspondence and requests for materials should be addressed to J.D. (jduan@uchicago. edu)

Genome-wide survey of interindividual differences of RNA stability in human lymphoblastoid cell lines

Jubao Duan^{1,2}, Jianxin Shi³, Xijin Ge⁴, Lars Dölken⁵, Winton Moy¹, Deli He¹, Sandra Shi¹, Alan R. Sanders^{1,2}, Jeff Ross⁶ & Pablo V. Gejman^{1,2}

¹Center for Psychiatric Genetics, NorthShore University HealthSystem Research Institute. Evanston, Illinois 60201, USA, ²Department of Psychiatry and Behavioral Sciences, The University of Chicago, Chicago, Illinois 60637, USA, ³Biostatistics Branch, Division of Cancer Epidemiology and Genetics, National Cancer Institute, Bethesda, Maryland 20892, USA, ⁴Department of Mathematics and Statistics, South Dakota State University, Brookings, South Dakota 57007, USA, ⁵Department of Medicine, Addenbrooke's Hospital, University of Cambridge, Cambridge CB2 0QQ, United Kingdom, ⁶McArdle Laboratory for Cancer Research, University of Wisconsin, Madison, Wisconsin 53706, USA.

The extent to which RNA stability differs between individuals and its contribution to the interindividual expression variation remain unknown. We conducted a genome-wide analysis of RNA stability in seven human HapMap lymphoblastoid cell lines (LCLs) and analyzed the effect of DNA sequence variation on RNA half-life differences. Twenty-six percent of the expressed genes exhibited RNA half-life differences between LCLs at a false discovery rate (FDR) < 0.05, which accounted for \sim 37% of the gene expression differences between individuals. Nonsense polymorphisms were associated with reduced RNA half-lives. In genes presenting interindividual RNA half-life differences, higher coding GC3 contents (G and C percentages at the third-codon positions) were correlated with increased RNA half-life. Consistently, G and C alleles of single nucleotide polymorphisms (SNPs) in protein coding sequences were associated with enhanced RNA stability. These results suggest widespread interindividual differences in RNA stability related to DNA sequence and composition variation.

Population variation of RNA abundances is widespread^{1,2}, influenced by DNA sequence variation³⁻¹¹, and has a heritability of $> 30\%^{4.7,9,12-14}$. RNA abundance is determined by the RNA transcription and RNA degradation (referred to here as RNA stability). Of the two, RNA transcription has been shown to have a larger effect than RNA stability on the cellular variance of total RNA abundances between genes^{15,16}. However, at a population level, the extent of RNA stability differences between individuals and how much the interindividual RNA stability differences contribute to the overall RNA abundances remain unknown.

RNA stability is regulated through interactions between cis-regulatory sequences and various RNA-binding proteins and microRNAs (miRNAs)¹⁷. DNA sequence variants (or polymorphisms) can modify the regulation of RNA stability and thereby affect interindividual RNA stability differences. The effects of nonsense mutations and 3'-UTR sequence variants have received the most attention¹⁸; transcripts containing premature stop codons (i.e., nonsense mutations) undergo accelerated RNA decay, known as nonsense-mediated RNA decay (NMD)¹⁸. Within the 3'-UTR, AU-rich elements (AREs) and miRNA target sites mediate RNA decay and block RNA translation¹⁹. *Cis*-regulatory sequences can also be present in the 5'-UTRs, protein coding regions^{18,20,21}, and intron/exon junctions^{22,23}. Synonymous sequence variants are often thought to be functionally silent, but we and others have shown that they can affect RNA stability²⁴⁻²⁹, where, as predicted, G and C alleles were associated with more stable transcripts. In humans, RNA stability differences have been implicated in inflammation³⁰⁻³⁴, cardio-vascular disease^{35,36}, cancer³⁷⁻³⁹, glaucoma²⁷, age-related macular degeneration⁴⁰, and neuropsychiatric disorders such as mental retardation^{41,42}, Alzheimer's disease⁴³, and some forms of muscular atrophy^{44,45}.

Using LCLs as a cellular model, we explored the extent of interindividual RNA stability differences, the contribution of RAN stability differences to variation of total RNA abundances between genes and between individuals, and the effects of gene sequence features and sequence variants on RNA stability differences in a genome-wide scale (outlined in Table 1). We employed a non-invasive approach to measure genome-wide RNA half-lives through assaying the ratio of 4sU-labeled nascent (newly transcribed) RNAs and total⁴⁶⁻⁴⁸. Seven HapMap LCLs derived from genetically unrelated HapMap individuals (hapmap.ncbi.nlm.nih.gov)^{3-7,10,11} were

Table 1 Summary of analyses			-
Motivation	Analysis	Result	Conclusion
Interindividual differences of total RNA abundances are prevalent, but extent of interindividual RNA stability differences is unknown.	Use ANOVA to test for RNA stability variation among 3 LCLs (with 3 biological replicates).	$\sim 25.7\%$ (n = 2884) genes with FDR < 0.05, of which 452 with > 2-fold differences.	Interindividual RNA stability differences are common.
Relative contribution of de novo transcription and of RNA decay to the total RNA abundance variation between genes in LCLs is unknown.	Examine the Pearson correlation of total RNA abundance with RNA half-life and with nascent RNA abundance across all the genes.	Nascent RNA differences explained ~ 80%, and RNA half-life differences explained ~ 6.4% of the variance in total RNA abundances between genes.	Transcriptional differences are a major determinant of the variance in transcript abundances between genes.
At population level, how much the interindividual RNA stability differences contribute to the interindividual total RNA abundance differences is unknown.		interindividual total RNA abundance	Transcriptional differences are a major determinant of the variance in transcript abundances between individuals.
RNA stability differences can be influenced by cis-regulatory sequences and genetic variants, trans-factors (e.g., miRNA or RNA-binding proteins) and environmental factors (e.g., cell line growth condition). We investigate whether the effects of sequence features and genetic variants are detectable in LCLs.	correlation between RNA half-life and different gene features across	RNA half-lives were weakly correlated with a number of gene features (transcript length, 3'-UTR length, miRNA target sites, GC and GC3 contents, etc.).	Multiple gene features contribute to RNA half-life differences between genes but with weak effect.
	Use linear regression to analyze the association of inter-individual RNA half-life differences with specific type of DNA sequence variants.	RNA half-lives, and for genes showing	individuals but with

analyzed. Three of the analyzed LCLs were part of the 1,000 Genomes Project dataset, thus allowing the examination on a genome-wide scale of the effect of specific types of genetic variants on RNA halflife. We found that $\sim 25.7\%$ of the expressed genes exhibited interindividual RNA stability differences, and $\sim 37\%$ of genes presenting interindividual expression variation showed detectable interindividual RNA stability differences that could explain the direction of gene expression differences. At a genome-wide scale, we found that nonsense mutations correlated with reduced RNA half-life (as expected) while G or C alleles of synonymous or non-synonymous SNPs weakly correlated with increased RNA half-life.

Results

RNA stability in LCLs and interindividual RNA half-life differences. We examined RNA stability by measuring the ratio of nascent RNAs and total RNAs employing two hour (h) 4sU labeling⁴⁶⁻⁴⁹ in seven LCLs. Three of them (GM07029, GM10835, and GM12813) had biological replicates (i.e., independent cell cultures for the same LCL) and each had technical duplicates (i.e., separate RNA aliquots from the same cell culture) which allowed us to test the reproducibility of the RNA measurements.

The purified 4sU-labeled nascent RNA accounted for ~ 3% of total RNA. 11,301 RefSeq genes (hg18) passed array quality control metrics and showed detectable expression in both nascent and total RNA samples (see Methods). RNA abundances showed high correlation (Pearson R = 0.91 ~ 0.93; See Supplementary Fig. S1 online) with the measurements on the same specimens that were also used in a previous study⁵⁰. RNA half-lives were determined for 11,132 genes, and correlated well (Pearson R > 0.5) with those previously reported for a human B cell line⁴⁸ (See Supplementary Fig. S2 online). While 2 h of 4sU-labeling increases the precision of RNA half-lives for medium and long-lived genes (compared to 1 h of 4sU-labeling), measurements of short RNA half-lives (< 1 h) becomes less precise due to the relatively large nascent/total RNA ratios (> 75%)⁵¹. As expected, genes with very short RNA half-life (< 1 h) thus showed rather poor reproducibility of RNA half-life measurements

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(Figure 1A). However, these genes only accounted for ~ 0.5% of all the expressed genes, and 2 h 4sU-labeling generated reproducible data for the more stable transcripts (Figure 1A). The correlation of RNA half-life (Figure 1B) followed the expected descending order from technical replicates (Pearson R = 0.92 - 0.95) to biological replicates (Pearson R = 0.85 - 0.96), and to unrelated LCLs (Pearson R = 0.72 - 0.82).

Gene ontology (GO) analysis of genes with different RNA stability in LCLs using the DAVID tool⁵² showed consistency with previous reports in other cell types^{51,53}. Rapidly decaying RNAs (n = 396, halflife < 2 h) were found enriched for GO-terms related to transcription (Benjamini family-wide FDR < 0.05; See Supplementary Table S1 online), while slowly decaying RNAs (n = 199, half-life > 15 h) were enriched for GO-terms related to protein synthesis, modification, and trafficking (FDR < 0.05; See Supplementary Table S2 online). For rapidly decaying genes, there was a trend towards enrichment of the GO-term of cell cycle (FDR = 0.058)⁵³.

We then analyzed RNA stability variation in the three LCLs with biological replicates of RNA half-life measurements. Out of the 11,132 genes with measured RNA half-life, analysis of variance (ANOVA) detected 2,884 genes (~ 25.7%) with interindividual RNA half-life differences at a Benjamini & Hochberg⁵⁴ FDR < 0.05 (corresponding nominal p = 0.013). 452 genes showed \geq 2-fold interindividual RNA half-life differences (Figure 2A and See Supplementary Table S3). To investigate whether the ANOVA pvalue for each transcript is robust to the normality assumption, we performed 1,000 random permutations to approximate the p-values. We found that the p-values based on standard ANOVA and permutations agreed very well (see Supplementary Fig. S3 online), and 21.7% (vs. \sim 25.7% in initial ANOVA) of expressed genes show interindividual RNA half-life differences at FDR < 0.05 (corresponding nominal p = 0.011) in permutation test. Adding the four additional LCLs (GM07019, GM12812, GM12814, and GM12815) for which there were no biological/technical replicates, we found more genes (total n = 3,589) showing interindividual half-life differences at FDR < 0.05 (See Supplementary Table S3), and the statistical significances (ANOVA p-values) of the result from all 7 LCLs

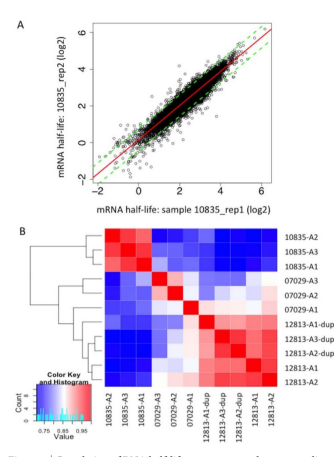


Figure 1 | Correlation of RNA half-life measurements between replicates and between cell lines. (A) A representative scatter plot of RNA half-life (\log_2 scale) from two biological replicates of GM10835. RNA half-life for each gene was calculated from the ratio of total RNA abundances and nascent RNA abundances after 2 h of 4sU-labeling. Green dashed lines indicate the 1.5-fold deviation from the trend line (red) at either side. Pearson R between biological replicates for the three LCLs ranged from 0.85 to 0.96. (B) Heatmap of RNA half-life correlation for all 3 cell lines (GM07029, GM10835, and GM12813) with biological replicates (denoted as A1, A2, and A3) and technical duplicates (denoted as "dup"). As shown in the color key, blue color represents lower correlation (Pearson R) and red color indicates higher correlation.

correlated well (Pearson R = 0.81; Figure 2B) with that of the three LCLs with biological replicates.

For an independent validation of interindividual differences of RNA half-life, we re-grew the three LCLs (GM07029, GM10835 and GM12813) and performed quantitative PCR (qPCR) on nascent and total RNAs for 15 selected genes. These 15 genes were randomly selected from those exhibiting interindividual RNA half-life differences > 2-fold at FDR < 0.05. We considered a gene as validated if the gene showed significant RNA stability differences (by ANOVA) across the three tested LCLs in qPCR with the same directional changes as in array experiment (based on the positive correlation of RNA stability between qPCR and array experiments). For 12 out of the 15 tested genes, qPCR yielded results consistent with those observed in our array experiment (Table 2). Among the 12 validated genes, 10 showed strong correlations of R > 0.70 (except for LANCL22 with R = 0.49 and C20orf43 with R = 0.59) between qPCR and microarray results, which we considered as a typical validation between qPCR and microarray^{55,56}. One possible explanation for the weaker correlations of those two genes (LANCL22 and C20orf43) could be that multiple probes for each gene were used in exon array experiment while only single probe was used in the qPCR.

De novo transcription, RNA half-life, and RNA abundance. Total RNA abundances are a result of de novo transcription and RNA decay. We estimated the relative contribution of de novo transcription (represented by nascent RNA abundance) and of RNA decay (indexed as RNA half-life) on RNA abundance by examining the coefficient of determination (R^2) of the correlations between total RNA, RNA half-life, and nascent RNA abundance. We initially analyzed the 3 LCLs with biological replicates. Total RNA abundances showed a Pearson correlation coefficient (R) of 0.89 ($p < 2.2 \times 10^{-16}$) with nascent RNA abundances, and an R of $0.25 (p < 2.2 \times 10^{-16})$ with RNA half-lives (Figures 3A and 3B). Hence, based on the R², nascent RNA abundance differences explained \sim 80%, and RNA half-life differences independently explained $\sim 6.4\%$ of the variance in total RNA abundances between genes. For the subset of genes showing interindividual RNA half-life differences (FDR < 5%), the proportion of the variance of total RNA abundances between genes explained by RNA half-life differences between genes increased to \sim 16% as estimated from the correlation between RNA half-life and total RNA abundance (R = 0.39, $p < 2.2 \times 10^{-16}$; Table 3 and Figure 3C and 3D). Analyzing the other 4 LCLs without replicates of measurements gave similar results (See Supplementary Fig. S4 online), with the proportion of variance in RNA abundances explained by RNA half-life differences were at $\sim 14.4\%$ (Pearson R = 0.38) for all the expressed genes and $\sim 21.2\%$ for the subset of genes showing interindividual RNA half-life differences (FDR < 5%). Including nascent RNA abundance and RNA halflife in a single linear model, we found that de-novo transcription and decay together explained 96.8% of the variance of variance of total RNA.

We next estimated the contributions of RNA decay and de novo transcription to the interindividual variation of total RNA abundance. At FDR < 0.05, between the same three LCLs for which we detected 2,884 genes with interindividual RNA half-life differences, 6,536 genes showed differences of total RNA abundances and 6,453 genes showed differences of nascent RNA abundances between LCLs. About 83% of the genes showing interindividual differences of total RNA abundances also showed differences of nascent RNA abundances and/or RNA half-lives (Figure 4). We further determined whether the interindividual differences of total RNA abundances were in the same directions as the interindividual differences of nascent RNA abundances and RNA half-lives by examining the directions of the correlations of total RNA abundances with nascent RNA abundances and with RNA half-lives across the three tested LCLs (See Supplementary Table S4). Out of the 6,536 genes with interindividual total RNA abundance differences, 2,997 (46%) showed differences only at the transcription level and 584 (9%) showed differences only at the RNA decay level, with > 99% of the differences in the same direction as gene expression changes; 1,808 (28%) showed differences at both transcriptional and decay levels, with > 74% of the differences in the same direction as gene expression changes (See Supplementary Table S4 online). These results indicate that transcriptional differences between individuals play a larger role than RNA stability differences in explaining the overall interindividual variation of gene expression. Nevertheless, \sim 37% of all genes presenting significant gene expression variation between individuals also showed RNA half-life differences (28% showing both transcription and RNA half-life differences while 9% showing only RNA_half-life differences), where most ($\sim 88\%$) showed the same direction of the differences in RNA half-lives and in total RNA abundances (See Supplementary Table S4).

Gene features correlated with genome-wide RNA stability. RNA decay is a process that involves both cis-sequence features and transprotein factors (e.g., RNA-binding proteins and miRNAs). We investigated here whether the effects of cis-sequence features on



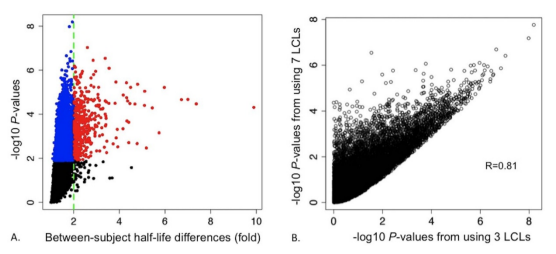


Figure 2 | **Interindividual differences of RNA half-life.** (A) ANOVA result from using three LCLs (GM07029, GM10835 and GM12813) with 3 biological replicates. The ANOVA *p*-values (in $-\log_{10}$ scale) are plotted against the between-subject differences of RNA half-life (expressed as the maximum fold of changes between the three LCLs). Red and blue dots represent genes (n = 2,884) showing RNA half-life differences at FDR < 0.05 (*p* = 0.013), of which red dots indicate genes (n = 452) with 2-fold or higher between-subject RNA half-life differences. (B) ANOVA *p*-values from using 3 LCLs (GM07029, GM10835 and GM12813) and from using all 7 LCLs (GM07019, GM07029, GM10835, GM12812, GM12813, GM12814, and GM12815) correlate very well (Pearson R = 0.81).

mRNA stability variation are detectable. As shown in Table 3, using the data from the 7 LCLs, we found a genome-wide negative correlation of RNA half-life with both transcript length (Spearman rank correlation R = -0.222; $p < 2.2 \times 10^{-16}$) and 3'-UTR length (R = -0.142; $p < 2.2 \times 10^{-16}$), which is consistent with previous reports^{22,51,53,57-60}. There was also a positive correlation (R = 0.191; $p < 2.2 \times 10^{-16}$) of RNA half-life with the number of exon/intron junctions per kb transcript (i.e., adjusted for transcript length) as previously reported^{22,60}, which has been suggested to be a result of a stabilizing effect of the exon/intron junction complex (EJC) on

Gene	Method	t _{1/2} -07029 (h)	SD	t _{1/2} -10835 (h)	SD	t _{1/2} -12813 (h)	SD	Fold-diff	ANOVA p	Pearson R (qPCR vs. array)
ARPC5L	array	2.96	0.50	4.16	0.31	2.03	0.36	2.05	6.9E-04	0.996
	qPCŔ	2.29	0.32	3.16	0.15	1.37	0.17	2.31	2.3E-04	
MESDC2	array	2.61	0.25	3.87	0.45	1.81	0.44	2.14	1.4E-03	0.964
	qPCŔ	1.36	0.23	1.78	0.20	0.67	0.11	2.67	9.7E-04	
NFE2L3	array	5.64	0.99	3.07	1.07	7.81	1.81	2.54	4.4E-03	-0.998
	qPCR	1.76	0.22	3.50	0.33	0.59	0.19	5.96	2.5E-05	
AMICA1	array	4.35	0.91	9.73	2.20	5.18	0.45	2.24	8.8E-04	0.999
	qPCŔ	1.70	0.11	3.67	0.34	2.10	0.41	2.16	6.0E-04	
HBXIP	array	5.54	1.13	7.95	1.08	3.84	0.78	2.07	1.7E-03	0.695
	qPCŔ	5.74	0.98	5.34	0.53	3.62	0.29	1.58	1.7E-02	
LANCL2 c	array	2.32	0.42	2.83	0.29	1.34	0.38	2.10	2.5E-03	0.493
	qPCŔ	2.44	0.10	1.70	0.18	1.33	0.08	1.83	1.3E-04	
GPC4	array	1.45	0.30	5.70	0.56	1.19	0.21	4.80	4.3E-06	0.837
	qPCŔ	3.11	0.21	4.10	0.26	1.68	0.75	2.43	2.3E-03	
C20orf43	array	3.64	0.48	6.93	0.25	2.43	0.43	2.86	2.5E-05	0.590
	qPCŔ	5.38	0.41	4.97	0.40	2.93	0.33	1.83	4.9E-04	
RGS1	array	8.41	1.60	4.18	0.57	5.45	0.46	2.01	6.6E-04	-0.156
	qPCŔ	1.84	0.07	2.59	0.23	0.60	0.12	4.28	1.3E-05	
PDCD4	array	4.43	0.76	5.16	0.37	2.01	0.25	2.57	9.4E-06	0.719
	qPCŔ	3.27	0.22	6.39	0.20	3.06	0.11	2.09	9.7E-07	
	array	4.44	0.34	6.98	0.35	2.58	0.55	2.70	1.3E-04	0.945
	qPCŔ	2.98	0.67	3.62	0.27	1.36	0.19	2.67	1.7E-03	
DNAJA2	array	2.60	0.60	2.80	0.24	1.21	0.09	2.32	3.6E-05	0.984
	qPCŔ	1.50	0.12	1.90	0.11	0.56	0.15	3.39	2.2E-04	
RPL14	array	7.72	1.43	5.18	0.55	3.07	0.62	2.51	6.4E-04	0.844
	qPCŔ	26.60	1.31	26.50	0.84	17.49	0.38	1.52	2.8E-05	
SFXN2	array	5.57	0.55	7.05	0.55	11.32	0.54	2.04	2.1E-06	-0.259
	qPCŔ	3.19	0.31	7.55	0.68	3.24	0.47	2.37	6.3E-05	
GKC	array	1.98	0.19	4.44	0.41	3.72	0.25	2.24	3.7E-06	0.985
	qPCŔ	0.77	0.17	13.76	0.65	12.18	0.39	17.77	6.3E-08	

Note: RNA half-life (1_{1/2}) measurements from array and qPCR methods are listed. The RNA samples used in array and qPCR experiments were from independent cell cultures for each of the 3 LCLs (GM07029, GM10835, and GM12813). There are 3 biological replicates (i.e., from independent cell cultures) for each measurement and the average RNA half-life and standard deviation (SD) are listed. Fold-diff=the ratio between the largest RNA half-life value and the smallest RNA half-life value among three LCLs. ANOVA is used to test the interindividual differences of RNA half-life. Pearson correlation (R) of RNA half-life measurements between array and qPCR experiments is calculated. Only genes showing both significant ANOVA p value (<0.05) in qPCR validation and positive Pearson R (qPCR vs. array) are are considered as independently confirmed (Pearson R in bold).



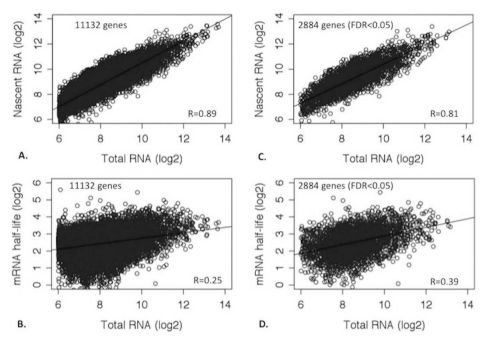


Figure 3 | Correlations of total RNA abundances with nascent RNA abundances and RNA half-lives in LCLs. Total RNA abundance, nascent RNA abundance and RNA half-life for each of the 11,132 genes (with RNA half-life calculated) were averaged from the 3 LCLs with biological and technical replicates. (A) shows a strong Pearson correlation (R = 0.89) between total RNA and nascent RNA abundances (log_2 scale) for all 11,132 genes, indicating that variance of total RNA abundances between genes was predominately determined by transcription. (B) Pearson correlation (R = 0.25) of total RNA abundances with RNA half-life for all 11,132 genes is shown. (C) and (D) show the Pearson correlations of total RNA abundances with nascent RNA abundances and RNA half-life (R = 0.81 and R = 0.39, respectively), for a subset of genes showing interindividual RNA half-life differences at FDR < 0.05 (n = 2,884).

spliced transcripts^{22,61}. In addition, we observed a negative correlation of RNA half-life and the number of 3'-UTR miRNA targets (R = -0.146; $p < 2.2 \times 10^{-16}$), an observation previously reported in plants⁵⁹ and consistent with the role of miRNAs in destabilizing its target mRNAs in mammalian cells⁶². For the subset of genes showing interindividual differences of RNA stability, we found a position correlation of RNA half-life with coding GC content (R = 0.141, $p = 9.0 \times 10^{-6}$) and with coding GC3 content (R = 0.197, $p = 8.4 \times 10^{-10}$) (Table 3). The correlation of RNA stability with coding GC3 content (R = 0.224, after correction for GC

Table 3 | Spearman rank correlation of RNA stability with total RNA abundance and gene features

content, $p = 2.8 \times 10^{-12}$). As GC3 content represents the nucleotide sequence composition at the third codon positions (which are mostly synonymous sites), our result suggests a role of synonymous SNPs on affecting RNA stability but with weak effect. The magnitudes of the correlations of RNA half-life with various genomic features in LCLs were low in general (R = 0.1 ~ 0.3), which were consistent to previous reports in other cell types or species^{22,51,53,57-60}.

As some of these tested genomic features are not fully independent (e.g., 3'-UTR length and the number of miRNA targets), we further examined the overall contribution of these genomic features to the

	All genes (n = 11,132)		Genes with FDR $<$ 0.05 (n = 2,884)		Genes with FDR < 0.01 (n = 1,077)	
	R	<i>p</i> -value	R	<i>p</i> -value	R	<i>p</i> -value
Total RNA level	0.253	< 2.2E-16	0.389	< 2.2E-16	0.389	2.2E-16
Transcript length	-0.222	< 2.2E-16	-0.274	< 2.2E-16	-0.242	8.5E-16
3'-UTR length	-0.142	< 2.2E-16	-0.171	< 2.2E-16	-0.167	5.2E-08
# of miRNĂ targets	-0.146	< 2.2E-16	-0.174	< 2.2E-16	-0.143	4.7E-07
# of miRNA targets/kb 3'-UTR	-0.133	< 2.2E-16	-0.151	1.7E-15	-0.120	1.0E-04
3'-UTR folding energy (G)	0.129	< 2.2E-16	0.142	1.1E-13	0.115	2.4E-04
5'-UTR length	-0.087	< 2.2E-16	-0.094	8.6E-10	-0.143	3.6E-06
5′-UTR folding energy (G)	0.094	< 2.2E-16	0.111	6.9E-09	0.131	3.0E-05
Coding region length	-0.189	< 2.2E-16	-0.209	< 2.2E-16	-0.217	1.9E-12
Coding GC content	0.004	7.3E-01	0.070	3.5E-04	0.141	1.3E-05
Coding GC3 content	0.029	3.7E-03	0.113	7.9E-09	0.197	1.0E-09
Coding GC3/GC content	0.044	1.2E-05	0.136	4.2E-12	0.224	2.9E-12
# of Intron	-0.014	1.4E-01	-0.038	4.6E-02	-0.064	3.8E-02
Intron length	-0.035	2.9E-04	-0.087	3.8E-06	-0.139	6.0E-06
Exon junctions/kb transcript	0.191	< 2.2E-16	0.217	< 2.2E-16	0.176	9.2E-09

Note: Spearman rank correlation (R) between RNA half-life and each gene feature and the corresponding *p*-value are listed for all genes (n = 11, 132) and genes showing interindividual differences of RNA half-life at FDR < 0.05 (n = 2,884) or FDR < 0.01 (n = 1,077). RNA half-life data are the averages from the 7 LCLs. Gene features for each gene are downloaded from UCSC hg 18 (RefSeq genes). RNA folding energies (pre-computed in UCSC hg 18) are predicted by RNAfold of Vienna RNA Package⁸¹. Bolded are correlations specific to or stronger in genes with interindividual differences of RNA half-life.

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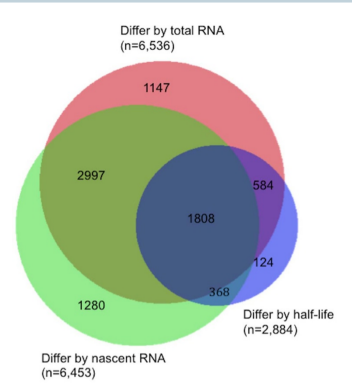


Figure 4 | Venn diagrams showing the number of genes with interindividual differences of total RNA abundances, nascent RNA abundances, RNA half-lives and both. A total of 11,132 genes were tested by ANOVA for interindividual differences (FDR < 0.05) in total RNA abundances, nascent RNA abundances and RNA half-lives between 3 LCLs. 6,536 genes showed interindividual total RNA abundance differences, of which 2,997 (46%) showed differences only at the transcription level and 584 (9%) showed differences only at the RNA decay level, and 1,808 (28%) showed differences at both transcriptional and RNA decay levels, indicating that transcriptional differences between individuals play a larger role than RNA stability differences in explaining the overall interindividual variation of gene expression. However, genes showing interindividual RNA half-life differences contributed to ~ 37% of all genes showing significant gene expression variation.

variance of RNA half-life by multiple linear regression analysis. For all the expressed genes, we found that genomic features listed in Table 3 explained ~ 9.5% of the RNA half-life variance ($R^2 = 0.095$). For the subset of genes showing interindividual RNA half-life differences, the fraction of the RNA half-life variance explained by these genomic features increased to ~ 13.4% ($R^2 = 0.134$).

Effect of DNA sequence variant type on RNA stability. Out of the seven LCLs for which we have RNA half-life data, three (GM12812, GM12814 and GM12815) have been sequenced by the 1000 Genomes Project (July 2012 release). Using the resultant genotype data for these three subjects, we examined whether interindividual RNA half-life differences were correlated with specific sequence changes, namely, nonsense SNPs (45 in 41 genes), SNPs disrupting miRNA-target sites (115 in 111 genes), and coding synonymous or non-synonymous SNPs with allele changes between "stable alleles" G or C and "unstable alleles" A or T (14,137 in 5,671 genes) (See Supplementary Table S5 online).

Nonsense mutations represent the most extreme protein coding sequence changes, some of which are known to cause RNA to undergo NMD¹⁸. Consistently, nonsense SNPs showed the strongest correlation with RNA half-life differences (See Supplementary Table S6), with premature stop-codons associated with reduced RNA half-lives (regression coefficient $\beta = 0.138$, P = 0.009; Figure 5A and See

Supplementary Table S5), supporting the validity of our RNA halflife measurements. For SNPs in miRNA target sequences, alleles disrupting the sequence pairing between miRNA and its target sequence are expected to block the miRNA-mediated RNA degradation, thereby increasing RNA half-lives. However, we did not observe a significant correlation between RNA half-life differences and different SNP alleles in miRNA target sites ($\beta = 0.022$, p = 0.312; Figure 5B), although the SNP effect appeared to be in the expected direction, which may be due to our underpowered sample size.

For coding synonymous and non-synonymous SNPs, our hypothesis was that G or C allele were "stable allele" and thus associated with increased RNA half-life. Our initial analysis of all the available 14,137 synonymous and non-synonymous SNPs in 5,671 genes did not reveal the expected correlation between G or C alleles and increased RNA half-life (See Supplementary Table S5). Given our underpowered sample size, we restricted the analysis to a total of 1,080 SNPs in 505 genes that had interindividual RNA half-life differences at the FDR < 0.01 level to increase the power to detect possible weak effect from G or C alleles. Such analysis showed a trend towards positive correlation between G or C alleles and RNA half-life $(\beta = 0.023, p = 0.075$ for untransformed RNA half-life; Figure 5C). Because the correlation analysis for genes with multiple coding SNPs can be confounded by non-independent tests (i.e., multiple SNPs linked to the same RNA half-life measurement) and complicated by possible balancing effects from multiple functional SNPs (e.g., the "stable" allele G or C allele of one SNP may cancel out the effect from an A or T allele of another SNP), we further restricted the analysis to only 261 genes with a single coding SNP (See Supplementary Table S7). We found that G or C allele was nominally associated with increased RNA half-life ($\beta = 0.071$, p = 0.021; Figure 5D and see Supplementary Table S5 online), but with very weak effect.

Discussion

About 80% of genes show interindividual transcript abundance differences^{1,2}, resulting from a balance between transcription and RNA decay. This pilot study represents the first attempt to characterize genome-wide interindividual RNA stability differences and the effects from specific types of genetic variants. We found that \sim 25.7% of all the expressed genes presented interindividual RNA stability differences (FDR < 0.05). Factoring in our qPCR validation rate of 80%, \sim 21% of all expressed genes in LCLs showed RNA halflife differences between individuals. Although transcriptional differences remain the largest known factor influencing the variance of total RNA abundances^{15,16}, RNA stability differences explained a larger fraction of the total RNA abundance variance in the subset of genes showing interindividual RNA half-life differences than in all the expressed genes (16% vs. 6%). For genes with interindividual total RNA abundance differences, 37% also showed interindividual RNA half-life differences with most ($\sim 88\%$) in the same direction as the gene expression differences. On a genome-wide scale, we have replicated the effect of nonsense SNPs in accelerating RNA decay, and detected weak effect of allele substitutions from A or T to G or C of synonymous and nonsynonmous SNPs in stablizing RNA transcripts.

While mRNA stability analysis in LCLs might be confounded by differences of sex or age of the donor, EBV transformation, and other technical factors (as would be total RNA abundance analysis), these confounding factors are unlikely to explain the observed extensive RNA stability differences between our LCLs. For the three LCLs used to compare RNA half-life differences, the donors' ages at the time of phlebotomy were similar (~ 40 s). Although the donors were not of the same sex, over-representation of sex-chromosome genes among those showing interindividual differences of RNA half-life was not observed (85/301 X chromosome genes, Fisher's exact test p = 0.46). In addition, there was no enrichment of EBV-affected genes⁶³ among

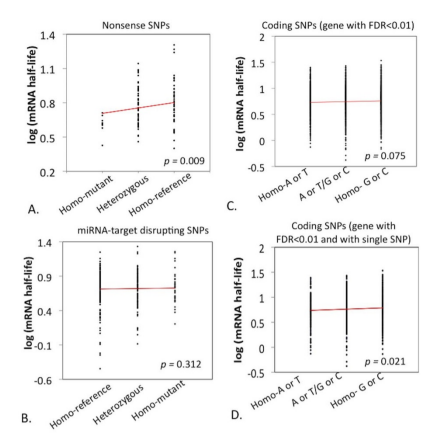


Figure 5 | Correlation of RNA half-life with sequence variants with 1,000 Genomes genotype data. For genes with SNPs polymorphic between the three individuals (GM12812, GM12814, and GM12815), RNA half-life (in log10 scale) and genotype for each individual were fitted in a regression analysis to estimate the regression coefficient β and p value for the correlation. (A) For 45 nonsense SNPs in 41 genes, mutant (stop-codon) allele of a nonsense SNP significantly correlates with decreased RNA half-life (homozygous mutant allele < heterozygous < homozygous reference allele) ($\beta = 0.138$, p = 0.009). This is consistent with expected NMD effect. One nonsense SNP, rs16982743, in *SIGLEC12* was not included in the analysis because it is in exon 1 of the gene and is expected to escape from NMD^{83,84}. (B) For 115 SNPs in miRNA sites of 111 genes, miRNA target-disrupting allele dose not show significant correlation with RNA half-life ($\beta = 0.022$, p = 0.312). (C) For 1,080 SNPs in 505 genes showing interindividual RNA half-life differences at FDR < 0.01, G or C allele does not show a correlation with RNA half-life ($\beta = 0.023$, p = 0.075). (D) For a subset of genes (n = 261) containing single coding SNPs, the G or C allele shows significant correlation with increased mRNA half-life (homozygous G or C alleles > heterozygous > heterozygous A or T alleles) ($\beta = 0.071$, p = 0.021). Single-side *p*-values are presented, because either a positive or negative correlation is expected between different genotypes and RNA half-life for each type of tested sequence variants.

those showing RNA half-life differences (32/103 EBV-affected genes, Fisher's exact test p = 0.39), suggesting that the EBV-transformation condition of different LCLs had no major effect on our results. Furthermore, the observed interindividual RNA half-life differences were unlikely to merely be the result of measurement problems and technical artifacts since: (1) The approach of 4sU-labeling of newly synthesized RNAs is less disruptive to cellular function compared to the traditional method of blocking transcription with actinomycin D, thus minimizing the confounding effects on global gene expression inherent to actinomycin D usage48,51. (2) The measurements of RNA half-life were highly reproducible (Pearson correlation R = 0.92 -0.95 between technical replicates), and the degree of correlation of the RNA half-life measurements was in the expected order of technical replicates > biological replicates > different LCLs, indicating the reliability of the RNA half-life measurements. (3) The arraymeasured interindividual RNA half-life differences for a random set of genes were validated by qPCR using independent cell cultures, suggesting that systematic bias is unlikely. Because trans-acting factors stabilize/destabilize RNA transcripts in a tissue/cell-specific manner^{64,65}, other cell/tissue types may show different sets of genes with interindividual RNA half-life differences. Nonetheless, our results indicated that interindividual RNA half-life differences are measurable and widespread on a genome-wide scale.

differences between genes and between individuals are detectable, such effects were very weak as indicated by the weak correlations between gene/sequence features and RNA stability ($R = 0.1 \sim 0.3$) (Table 3). On the other hand, the observed weak correlations are consistent with previous reports^{22,51,53,57-60}, and have similar magnitudes to that of reported correlation between expression level and gene features, where the maximum R is -0.26 with 3'-UTR length out of ~ 200 gene/sequence features examined⁶⁶. Some explanations for the observed weak correlations may include: (1) Other unexamined gene/DNA sequence features may contribute to the RNA stability variance, each with a small effect similar to what have been demonstrated for total RNA abundances⁶⁶. (2) Gene/sequence features in some subset of genes may show stronger effect on RNA halflife. (3) More importantly, as RNA decay is a complicated process that involves not only cis-sequence features but also various transfactors (e.g., RNA-binding proteins or miRNAs)17, expression variation of such trans-factors may also contribute to the RNA stability differences between cell line. In our dataset, although we did not find over-representation of genes related to RNA decay or RNA binding among (by DAVID tool; data not shown) those showing interindividual differences of total RNA abundances, we did find significant interindividual differences of total RNA abundance for a number of

Although the effects of gene sequence features on RNA stability

critical RNA decay-related genes¹⁷ (17/26 expressed genes with interindividual differences at FDR < 5%) (Supplementary Table S8).

For genes showing RNA stability differences, we have found a weak correlation between RNA half-life and coding GC and GC3 contents (Table 3). We also found that G or C alleles of coding SNPs were also weakly correlated with more stable RNAs for a subset of genes showing interindividual RNA half-life differences and with a single SNP (Figure 4D). Our results are thus consistent with that GC base pairings are more stable than an AU pairings (3 vs. 2 hydrogen bonds), suggesting a role of thermodynamic stability of RNA molecules in determining RNA stability. On a genome-wide scale, our results are in line with previous observations in individual genes where coding SNPs (synonymous or nonsynonymous SNPs) affecting RNA stability all had allele changes between G or C and A or $T^{24-26,67,68}$.

It is important to acknowledge a number of limitations of this pilot study. Since both transcription and RNA decay are dynamic processes subject to extensive temporal and spatial regulation^{15,69,70}, our analytical model is a simplified representation of a very complex system (e.g, alternative RNA splicing was not considered here). Furthermore, while our results suggest a relatively weaker contribution of RNA decay (compared to de novo transcription) to the overall gene expression variation, RNA stability may have a larger contribution to the plasticity of RNA levels, including the kinetics and the overall response to cellular perturbations. Lastly, due to the small sample size, we have restricted our analyses to examining the collective effects of different types of DNA sequence variants on mRNA stability, rather than identifying specific DNA sequence variants that influence mRNA stability by a genome-wide association study. Nonetheless, our study suggested that interindividual RNA stability differences are pervasive and influenced by specific gene features and sequence variants. These results thus warrant further genome-wide mapping of individual sequence variants associated with interindividual RNA stability differences in larger cohorts of samples. Given the substantial role of gene expression regulation in common disease pathogenesis^{29,71,72}, delineating the regulatory effect of genetic variants on RNA transcription and RNA decay will help elucidate novel causal mechanisms underlying genetic associations with common diseases.

Methods

Cell culture. We obtained LCLs from Coriell Cell Repositories (ccr.coriell.org). These LCLs were derived from 7 unrelated HapMap CEU individuals, GM07019 (female), GM07029 (male), GM10835 (male), GM12812 (male), GM12813 (female), GM12814 (male), and GM12815 (female). All donors were genetically unrelated Caucasians, of which only GM07019, GM07029, and GM10835 had age information (45, 47, and 39 years old, respectively). To minimize the effect of non-genetic factors on gene expression, we kept cell culture conditions consistent across different cell lines, including (1) using the same batch of media and drug (4sU; Sigma), (2) seeding cells at the same density before treatment of 4sU, (3) harvesting the 4sU-treated cells at the same time under similar conditions across cell flasks, and (4) assuring that different LCLs and biological replicates have the same growth rates (i.e., cell number doubling time). In brief, we grew the cells in 15 ml RPMI 1640 media (from ATCC) supplemented with 15% fetal bovine serum (FBS; from ATCC) in T-25 cell culture flasks. We passaged the cells at an initial cell density of 5×10^5 cells/ml and change media every other day. Upon cell growth reaching log phase of growth (usually ~ 5 days after the first passaging), we seeded the cells at a density of 2.5×10^5 cells/ml and counted the viable cell number the next day to make sure the cell growth was in log phase for all the cell lines. To start the 4sU-labeling experiment, we seeded the logphase growing cells at a density of 2.5×10^5 cells/ml and 24 h later, we treated cells with 200 μ M 4sU for 2 h and harvested cells (cell density reached to 5–6 \times 10⁵ cells/ml). The concentration of 200 µM of 4sU has been shown to allow efficient labeling of nascent RNAs without affecting global gene expression^{47,48}. The experiments were approved by Northshore Healthsystem Research Institute Institutional Review Board.

Total RNA and nascent RNA isolation. We used TRIzol reagent (Invitrogen) to isolate total RNAs from ~ 7.5 × 10° LCL cells and further purified the total RNAs by the RNeasy Mini Kit (Qiagen). To separate the 4sU-labeled nascent RNAs from total RNAs, we followed the protocol developed by Dolken et al.⁴⁸. Briefly, ~ 60 µg of total RNAs were subjected to biotinylation in the presence of 1 mg/ml of Biotin-HPDP (Pierce) for 3 h at room temperature, where only 4sU-labeled RNAs an be

biotinylated, and then precipitated by 2-propanol. The precipitated RNAs were resuspended and incubated with 100 µL of µMACS streptavidin-coated beads (Miltenyi Biotec) to allow the biotinylated nascent RNAs to bind to µMACS beads. We then applied the mixture onto the top of a column placed in the magnetic field of a µMACs separator (Miltenyi Biotec), where only the nascent RNAs bind with streptavidin-coated beads in high affinity and were retained in the column by the strong magnetic field. After three washes (100 mM Tris-HCl, pH 7.4, 10 mM EDTA, 1 M NaCl, 0.1% Tween20) at 65°C and then another three washes at room temperature, the nascent RNAs were eluted from the column by the addition of 100 mM dithiothreitol (DTT). We found that the binding of nascent RNAs to the streptavidin-coated microbeads was specific and time/dosage-dependent (See Supplementary Fig. S5 online). We were able to recover 2 µg of nascent RNAs, which was sufficient for subsequent exon array profiling, from a total of \sim 60 µg total RNAs derived from $\sim 7.5 \times 10^6$ LCLs treated with 200 μ M of 4sU for 2 h. The recovered nascent RNAs were further purified using RNeasy MinElute Spin columns (Qiagen). Before sending the RNA samples to the array facility, we assessed RNA quality for both total and nascent RNAs by measuring the A_{260}/A_{280} ratio (all > 1.9) by Biophotometer (Eppendorf). We also examined RNA integrity on agarose gel by visually checking bands of 28S and 18S rRNAs.

Affymetrix Human Exon 1.0 ST array processing. The abundance of total and nascent RNAs was assayed in parallel with Affymetrix Human Exon arrays at the University of Chicago functional genomics core. The array facility rechecked the total and nascent RNAs for concentration, A_{260}/A_{280} ratio, and RNA integrity by Nanodrop (Thermo Scientific) and an Agilent 2100 Bioanalyzer. As a routine procedure for the exon array assay at the array facility, ribosomal RNA (rRNA) reduction was performed with the RiboMinus Transcriptome Isolation Kit (Invitrogen) before cDNA synthesis to minimize the background hybridization signal. 100 ng of total RNAs was used in cDNA synthesis with the GeneChip WT cDNA Synthesis Kit (Affymetrix), and the synthesized cDNAs were fragmented and end-labeled with the GeneChip WT terminal Labeling Kit (Affymetrix). 5.5 µg fragmented/labeled cDNAs were used to hybridize with probes on the exon array at 45°C tor 16–18 h. The hybridized chips were washed and stained on a GeneChip Fluidics Station 450, and scanned on a GCS3000 Scanner (Affymetrix).

Exon array data analyses. We used the Affymetrix Power Tools (APT) software package to normalize the probe intensities and summarize the expression values at the probe set level (representing exon expression) and at the meta-probe set level (representing gene expression). See Supplementary Table S9 summarizes key quality control (QC) metrics. ~ 597 K SNP-containing probes (based on dbSNP build 129)7 were removed before probe intensity normalization, since such a SNP may affect the probe's binding affinity in the array hybridization and thus lead to biased estimates of gene expression level^{1,74,75}. Hybridization intensities of the remaining probes on an array were log₂-transformed and then quantile normalized^{76,77}. We carried out the normalization process for the "C" series of arrays with total RNAs and the "N" series of arrays with nascent RNAs separately, because N and C arrays represent different biology and therefore may show different gene expression patterns. To determine the reliability of the signal intensity, we generated a detection P-value of each probe set (core exon) or each meta-probe set (transcript cluster or gene) using the "Detection Above Background" (DABG) algorithm implemented in APT. A detection $P \le 0.01$ (recommended by Affymetrix) was used as a cut-off for determining whether a probe was expressed (i.e., detectable). Furthermore, only genes expressed in > 95% samples and with log-scale expression values > 6 in the "C" series of arrays were considered for further expression analyses. The number of the expressed probe sets (< average plus two standard deviations) on each array was used as one of the key QC metrics to detect array outliers. We identified a "C" array, 12813_rep3 (A12813A3 in Supplementary Fig. S6 online), as an outlier and excluded it from further analysis for the following reasons: (1) it had \sim 120 K expressed probe sets, which is > 2 standard deviations lower than the average number of expressed probe sets; (2) the correlation coefficient of gene expression values in 12813_rep3 with other biological or technical

replicates (R = $0.94 \sim 0.98$) appeared lower than that between other pairs of replicates of sample 12813 (R = $0.98 \sim 0.99$); and (3) a multidimensional scaling (MDS) analysis also indicated that A12813A3 was not clustered together with other replicates.

Estimation of RNA half-life from microarray data. The steady-state level of RNA transcripts measured from "C" arrays reflects the kinetic balance between transcription and RNA degradation. At steady state level, the RNA decay process in the period of 4sU-labeling can be expressed as^{48,49,78}.

$$1 - \frac{N}{C} = e^{-k_d t}$$

where k_d is the decay constant, N is the amount of nascent RNAs measured from "N" arrays and C is the amount of the total RNAs measured from "C" arrays. A commonly used term for describing RNA stability, RNA half-life (T_{1/2}), is dependent on the k_d and can be calculated by⁴⁹:

$$T_{1/2} = \frac{\ln 2}{k_d}$$

Therefore, to determine the RNA decay constant k_d and subsequently to estimate the RNA half-life in each cell line, we needed to determine the ratio N/C from the microarray data with $t=2\,$ h (i.e., the 4sU-labeling time). Because the N and C arrays

for each cell line were processed independently, we normalized the scale of N/C arrays using the global median RNA half-life method⁴⁸. Briefly, based on the median global RNA half-life in human B-cells (\sim 5.3 h⁴⁸, we determined a correction factor for the expression values between a pair of N and C arrays for each LCL, and then calculated the ratio of N/C from which the decay constant k_d and $T_{1/2}$ were calculated⁴⁹. Because RNA half-life in o be calculated only when N/C <1 (see the formula above), we were able to estimate RNA half-life for 11,132 out of 11,301 expressed genes. Applying such a correction factor based on predefined B-cell median half-life to calculate RNA half-life between individuals, because the median N/C ratios before applying the correction factor were very similar across different samples (1.66 to 1.69), implying that the median degradation rates and RNA half-lives were very similar between samples.

qPCR. Total and 4sU-labeled nascent RNAs were extracted from LCLs cultured independently from those used in the exon array experiment. We performed reverse transcription from RNAs to cDNAs with random hexamer primers using the Applied Biosystem (ABI) Reverse-transcription kit. With cDNAs as templates, we used one ABI TaqMan gene expression assay per gene in real-time PCR to quantify the expression of a target gene in total and nascent RNAs, respectively. Real-time PCR was performed in 384-well plates with 3 replicates on an ABI 7900 with standard cycling parameters. For quantification of gene expression, we used the comparative delta-Ct method with *glyceraldehyde 3-phosphate dehydrogenase* (*GAPDH*), a constitutively expressed gene, as an endogenous control to normalize the quantity of target transcripts (i.e., the relative expression value of a target gene = $2^{(C_{Larget} - C_{L}GAPDH)}$). RNA half-life was then estimated from the ratio of nascent/total RNAs as described above in the microarray experiment.

Bioinformatics. We downloaded RNA length, exon and intron lengths, miRNA target sites, 5' - and 3'-UTR sequences and folding energies from UCSC hg18 (RefSeq genes). For genes with multiple transcripts, the longest transcript was chosen in the analyses. We calculated coding GC content and GC3 content using locally installed CodonW version 1.4.2 (codonw.sourceforge.net//culong.html). Whole genome sequencing data for three individuals (GM12812, GM12814, and GM12815) were from the 1,000 Genomes Project (June 2011 release). The downloaded genotype file from the 1,000 Genomes Project (in VCF 4.0 format) was further processed in VCFtools (vcftools.sourceforge.net) and PLINK79 to keep only genotypes for coding SNPs (synonymous, non-synonymous, and nonsense) and 3'-UTR SNPs disrupting miRNA target sites. We used ANOVAR⁸⁰ to systematically perform functional SNP annotation and to extract RNA sequences carrying different alleles of a SNP. The minimum free energy of RNA folding structures was predicted by RNAfold of Vienna RNA Package (www.tbi.univie.ac.at/~ivo/RNA/)⁸¹ using the default setting. DAVID tools were used for enrichment analyses of GO terms and pathways⁵². For clustering genes with co-regulated RNA stability, we used the R package for WGCNA, which identifies modules of densely interconnected genes based on a correlation matrix containing all pair-wise Pearson correlations between all genes across all subjects⁸². We carried out here hierarchical clustering across all 7 LCLs (to boost the power) for RNA half-life of the 4,533 genes showing interindividual RNA half-life differences with a p < 0.05.

Statistical analyses. Statistical analyses were performed using R (www.r-project.org). For each gene, we performed ANOVA to test the null hypothesis that the measurements (total RNA abundances, nascent RNA abundances, and RNA half-life) had no interindividual differences. A P-value was calculated for each gene and each measurement. Genes showing significant interindividual differences in each measurement were identified by controlling the Benjamini & Hochberg FDR $< 0.05^{54}$, corresponding to the ANOVA nominal *P* < 0.013. To investigate whether the ANOVA p-value for each transcript is robust to the normality assumption, we first calculated the F statistic using ANOVA, which measured how well the replicates were separated among different cell lines. Then, we performed 1,000 permutations and approximated the p-value as the proportion of permutation with F statistic greater than the observed one. We then examined whether the p-values of standard ANOVA analysis based on normality assumption and the permutations agree well. For the Spearman rank correlation of RNA half-life with gene features, we averaged RNA half-life for each gene from the 3 LCLs with biological replicates to reduce the noise. To evaluate the overall contribution of various genomic features to the RNA half-life variance between genes, we performed multivariate linear regression using log10-transformed RNA half-life as the dependent variable and log₁₀-transformed genomic features (except 5' - and 3'-UTR folding energies were not log₁₀-transformed as they are negative values) as predictors (Table 3).

Furthermore, we investigated whether the RNA half-life was associated with sequence variants from different functional categories, e.g. nonsense SNPs, miRNA target-disrupting SNPs in 3'-UTRs, and coding synonymous and non-synonymous SNPs. Ideally, a linear regression analysis could be done for each SNP with a sufficient sample size. In our study, however, we had no power to detect any association based on only three LCLs with whole genome sequence data available. Therefore, we aggregated the SNPs for each category and tested the overall association for each category. Our test involves the following major steps: (1) We extracted the SNPs polymorphic in the three samples sequenced by the 1000 Genomes Project and classified the extracted SNPs into functional categories. (2) We coded homozygotes of alleles putatively decreasing RNA half-life (i.e., nonsense allele of a nonsense mutation, reference allele of a miRNA target-disrupting SNP, and "unstable" allele A or T of a coding synonymous or non-synonymous SNP) as 0, heterozygotes as 1, and

homozygotes of the other allele (i.e., the reference allele of a nonsense SNP, miRNA target-disrupting allele, and "stable" allele G or C of a coding SNP) as 2. (3) A linear regression analysis was performed to test the overall correlation between the RNA half-life and the genotypes across subjects and qualified SNPs from each functional category. We computed single-sided *P*-values because only positive correlations were expected based on our SNP coding scheme.

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Author contributions

J.D. and P.V.G. conceived and designed the experiments. D.H., S.S. and J.D. performed the experiments. J.S., X.G., L.D., W.M. and J.D. analyzed the data. A.R.S. and J.R. helped to interpret the results. J.D. wrote the paper. All authors reviewed the manuscript.

Additional information

Accession codes: The microarray data have been submitted to GEO under accession no. GSE34204.

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