A Computational Approach for the Identification of Novel L1 Transcriptional Regulators

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\textbf{KEYWORDS:} LINE1, transposons, regulators, eQTLs, screen, aging, GWAS
ABSTRACT

Long interspersed element 1 (L1) are a family of autonomous, actively mobile transposons that occupy ~17% of the human genome. The pleiotropic effects L1 induces in host cells—promoting genome instability, inflammation, or cellular senescence—are established, and L1’s associations with aging and aging diseases are widely recognized. However, because of the cell type-specific nature of transposon control, the catalogue of L1 regulators remains incomplete.

Here, we employ an eQTL approach leveraging transcriptomic and genomic data from the GEUVADIS and 1000Genomes projects to computationally identify new candidate regulators of L1 expression in lymphoblastoid cell lines. To cement the role of candidate genes in L1 regulation, we experimentally modulate the levels of top candidates in vitro, including IL16, STARD5, HSDB17B12, and RNF5, and assess changes in TE family expression by Gene Set Enrichment Analysis (GSEA). Remarkably, we observe subtle but widespread upregulation of TE family expression following IL16 and STARD5 overexpression. Moreover, a short-term 24 hour exposure to recombinant human IL16 was sufficient to transiently induce subtle but widespread upregulation of L1 subfamilies. Finally, we find that many L1 expression-associated genetic variants are co-associated with aging traits across genome-wide association study databases.

Our results expand the catalogue of genes implicated in L1 transcriptional control and further suggest that L1 contributes to aging processes. Given the ever-increasing availability of paired genomic and transcriptomic data, we anticipate this new approach to be a starting point for more comprehensive computational scans for transposon transcriptional regulators.
BACKGROUND

Transposable elements (TEs) constitute ~45% of the human genome [1]. Among these, the long interspersed element-1 (LINE-1 or L1) family of transposons is the most abundant, accounting for ~16-17% [1, 2], and remains autonomously mobile, with humans harboring an estimated 80-100 retrotransposition-competent L1 copies [3]. These retrotransposition competent L1s belong to evolutionarily younger L1PA and L1Hs subfamilies, are ~6 kilobases long, carry an internal promoter in their 5'-untranslated region (UTR), and encode two proteins — L1ORF1p and L1ORF2p — that are necessary for transposition [4]. The remaining ~500,000 copies are non-autonomous or immobile because of the presence of inactivating mutations or truncations [1] and include L1 subfamilies of all evolutionary ages, including the evolutionarily older L1P and L1M subfamilies. Though not all copies are transposition competent, L1s can nevertheless contribute to aspects of aging [5, 6] and aging-associated diseases [7-10].

Though mechanistic studies characterizing the role of L1 in aging and aging-conditions are limited, it is clear that its effects are pleiotropic. L1 can contribute to genome instability via insertional mutagenesis and an expansion of copy number with age [11] and during senescence [12]. L1 can also play a contributing role in shaping inflammatory and cellular senescence phenotypes. The secretion of a panoply of pro-inflammatory factors is a marker of senescent cells, called the senescence associated secretory phenotype (SASP) [13]. Importantly, the SASP is believed to stimulate the innate immune system and contribute to chronic, low-grade, sterile inflammation with age, a phenomenon referred to as “inflamm-aging” [13, 14]. During deep senescence,
L1 are transcriptionally de-repressed and consequently generate cytosolic DNA that initiates an immune response consisting of the production and secretion of pro-inflammatory interferons [15]. Finally, L1 is causally implicated in aging-associated diseases like cancer. L1 may contribute to cancer by (i) serving as a source for chromosomal rearrangements that can delete tumor-suppressor genes [16] or (ii) introducing its promoter into normally-silenced oncogenes [17]. Thus, because of the pathological effects L1 can have on hosts, it is critical that hosts maintain precise control over L1 activity.

Eukaryotic hosts have evolved several pre- and post-transcriptional mechanisms for regulating TEs [18, 19]. Nevertheless, our knowledge of regulatory genes remains incomplete because of cell type-specific regulation and the complexity of methods required to identify regulators. Indeed, one clustered regularly interspaced short palindromic repeats (CRISPR) screen in two cancer cell lines for regulators of L1 transposition identified >150 genes involved in diverse biological functions, such as chromatin regulation, DNA replication, and DNA repair [20]. However, only about ~36% of the genes identified in the primary screen exerted the same effects in both cell lines [20], highlighting the cell type-specific nature of L1 control. Moreover, given the complexities of in vitro screens, especially in non-standard cell lines or primary cells, in silico screens for L1 regulators may facilitate the task of identifying and cataloguing candidate regulators across cell and tissue types. One such attempt was made by generating gene-TE co-expression networks from RNA sequencing (RNA-seq) data generated from multiple tissue types of cancerous origin [21]. Although co-expression modules with known TE regulatory functions, such as interferon signaling, were
correlated with TE modules, it is unclear whether other modules may harbor as of now uncharacterized TE-regulating properties, since no validation experiments were carried out. Additionally, this co-expression approach is limited, as no mechanistic directionality can be assigned between associated gene and TE clusters, complicating the prioritization of candidate regulatory genes for validation. Thus, there is a need for the incorporation of novel “omic” approaches to tackle this problem. Deciphering the machinery that controls TE activity in healthy somatic cells will be crucial, in order to identify checkpoints lost in diseased cells.

The 1000Genomes Project and GEUVADIS Consortium provide a rich set of genomic resources to explore the mechanisms of human TE regulation in silico. Indeed, the 1000Genomes project generated a huge collection of genomic data from thousands of human subjects across the world, including single nucleotide variant (SNV) and structural variant (SV) data [22, 23]. To accomplish this, the project relied on lymphoblastoid cell lines (LCLs), which are generated by infecting resting B-cells in peripheral blood with Epstein-Barr virus (EBV). Several properties make them advantageous for use in large-scale projects, e.g. they can be generated relatively uninvasively, they provide a means of obtaining an unlimited amount of a subject’s DNA and other biomolecules, and they can serve as an in vitro model for studying the effects of genetic variation with any phenotype of interest [24, 25]. Indeed, the GEUVADIS Consortium generated transcriptomic data for a subset of subjects sampled by the 1000Genomes Project, and used their genomic data to define the effects genetic variation on gene expression [26]. Together, these resources provide a useful toolkit for investigating the genetic regulation of TEs, generally, and L1, specifically.
In this study, we (i) develop a pipeline to identify novel candidate regulators of L1 expression in lymphoblastoid cell lines, (ii) provide experimental evidence for the involvement of top candidates in L1 expression control, and (iii) expand and reinforce the catalogue of diseases linked to L1.
RESULTS

In silico scanning for L1 subfamily candidate regulators by eQTL analysis

To identify new candidate regulators of L1 transcription, we decided to leverage publicly available human “omic” datasets with both genetic and transcriptomic information. For this analysis, we focused on samples for which the following data was available: (i) mRNA-seq data from the GEUVADIS project, (ii) SNVs called from whole-genome sequencing data overlayed on the hg38 human reference genome made available by the 1000Genomes project, and (iii) repeat structural variation data made available by the 1000Genomes project. This yielded samples from 358 European and 86 Yoruban individuals, all of whom declared themselves to be healthy at the time of sample collection (Figure 1A). Using the GEUVADIS data, we obtained gene and TE subfamily expression counts using TEtranscripts [27]. As a quality control step, we checked whether mapping rates segregated with ancestry groups, which may bias results. However, the samples appeared to cluster by laboratory rather than by ancestry (Figure S1A). As additional quality control metrics, we also checked whether the SNV and SV data segregated by ancestry following principal component analysis (PCA). These analyses demonstrated that the top two and the top three principal components from the SNV and SV data, respectively, segregated ancestry groups (Figure S1B, Figure S1C).

We then chose to do a three-part integration of the available “omic” data (Figure 1B). Since TEtranscripts quantifies TE expression aggregated at the TE subfamily level
and discards TE position information, we chose to carry out a *trans*-eQTL analysis against global expression of each L1 subfamily. We reasoned that there would have to be factors (i.e., miRNAs, proteins, non-coding RNAs) mediating the effects of SNVs on L1 subfamily expression. Thus, to identify candidate genic mediators, we searched for genes with *cis*-eQTLs that overlapped with L1 *trans*-eQTLs. As a final filter, we reasoned that for a subset of regulators, L1 subfamily expression would respond to changes in the expression of those regulators. Consequently, we chose to quantify the association between L1 subfamily expression and candidate gene expression by linear regression. We hypothesized that this three-part integration would result in combinations of significantly correlated SNVs, genes, and L1 subfamilies (*Figure 1B*).

The *trans*-eQTL analysis against every expressed L1 subfamily led to the identification of 499 *trans*-eQTLs distributed across chromosomes 6, 11, 12, 14, and 15 that passed genome-wide significance (*Figure 1C, Supplementary Table S1A*). The *cis*-eQTL analysis led to the identification of 845,260 *cis*-eQTLs that passed genome-wide significance (*Supplementary Figure S2, Supplementary Table S1B*). After integrating the identified *cis* - and *trans*-eQTLs and running linear regression, we identified 1,272 SNV-Gene-L1 trios that fulfilled our three-part integration approach (*Supplementary Table S1C*). Among this pool of trios, we identified 7 unique protein-coding genes including *IL16, STAR*D5, HLA-DRB5, HLA-DQA2, HSD17B12, RNF5, and FKBPL* (*Figure 1C*). We note that although *EHMT2* did not pass out screening approach, it does overlap *EHMT2-AS1*, which did pass our screening thresholds (*Figure 1C*). We also note that several other unique non-coding genes, often
overlapping the protein-coding genes listed, were also identified (Figure 1C). For simplicity of interpretation, we focused on protein-coding genes during downstream experimental validation.

Next, to define first and second tier candidate regulators, we clumped SNVs in linkage disequilibrium (LD) by L1 trans-eQTL p-value to identify the most strongly associated genetic variant in each genomic region (Figure 2A, Supplementary Figure S3A). LD-clumping identified the following index SNVs (i.e. the most strongly associated SNVs in a given region): rs11635336 on chromosome 15, rs9271894 on chromosome 6, rs1061810 on chromosome 11, rs112581165 on chromosome 12, and rs72691418 on chromosome 14 (Supplementary Table S1D). Genes linked to these SNVs were considered first tier candidate regulators and included IL16, STARD5, HLA-DRB5, HLA-DQA2, and HSD17B12 (Figure 2B, Supplementary Table S1E). The remaining genes were linked to clumped, non-index SNVs and were consequently considered second tier candidates and included RNF5, EHMT2-AS1, and FKBPL (Supplementary Figure S3B). Additionally, for simplicity of interpretation, we considered only non-HLA genes during downstream experimental validation, since validation could be complicated by the highly polymorphic nature of HLA loci [28] and their involvement in multi-protein complexes.

Finally, to computationally determine whether candidate genes may causally influence L1 subfamily expression, we carried out mediation analysis on all SNV-gene-L1 trios (Supplementary Figure S4A). Interestingly, 868 out of the 1,272 (68.2%) trios
exhibited significant (FDR < 0.05) mediation effects (Supplementary Table S1F).

Among the 1st tier candidate regulators, significant, partial, and consistent mediation effects could be attributed to STARD5, IL16, HSD17B12, and HLA-DRB5 (Supplementary Figure S4B, Supplementary Table S1F). To note, while significant mediation could be attributed to the index SNV for STARD5, significant mediation could only be attributed to clumped SNVs for IL16 and HSD17B12. Given that STARD5 and IL16 share cis-eQTL SNVs, this suggests that STARD5 may be the more potent mediator. Among the 2nd tier candidate regulators, significant, partial, and consistent mediation effects could be attributed to RNF5, EHMT2-AS1, and FKBPL (Supplementary Figure S4C, Supplementary Table S1F). These results suggest that candidate genes may mediate the effects between linked SNVs and L1 subfamilies.

In silico scanning for L1 subfamily candidate regulators in an African population

We next sought to assess the cross-ancestry regulatory properties of candidate genes by repeating our scan using the Yoruban samples as a smaller but independent replication cohort. Here, rather than conduct a genome-wide scan for cis- and trans-associated factors, we opted for a targeted approach focusing only on gene cis-eQTLs and L1 subfamily trans-eQTLs that were significant in the analysis with European samples (Supplementary Figure S5A). The targeted trans-eQTL analysis led to the identification of 227 significant (FDR < 0.05) trans-eQTLs distributed across chromosomes 6 and 11 (Supplementary Table S2A). The targeted cis-eQTL analysis led to the identification of 1,248 significant (FDR < 0.05) cis-eQTLs (Supplementary
Table S2B). After integrating the identified cis- and trans-eQTLs and running linear regression, we identified 393 SNV-Gene-L1 trios that fulfilled our three-part integration approach (Supplementary Table S2C). Among this pool of trios, we identified 2 unique protein-coding genes—HSD17B12 and HLA-DRB6—as well as several unique non-coding genes (Supplementary Table S2C). Again, we clumped SNVs in linkage disequilibrium (LD) by L1 trans-eQTL p-value. LD-clumping identified the following index SNVs: rs2176598 on chromosome 11 and rs9271379 on chromosome 6 (Supplementary Table S2D). Genes linked to these SNVs were considered first tier candidate regulators and included both HSD17B12 and HLA-DRB6 (Supplementary Figure S5B, Supplementary Table S2E). Finally, we carried out mediation analysis on all SNV-gene-L1 trios; however, no significant (FDR < 0.05) mediation was observed (Supplementary Table S2F). These results implicate HSD17B12 and the HLA loci as candidate, cross-ancestry L1 expression regulators.

To assess why some candidate genes did not replicate in the Yoruba cohort, we manually inspected cis- and trans-eQTL results for trios with those genes (Supplementary Figure S6A). Interestingly, we identified rs9270493 and rs9272222 as significant (FDR < 0.05) trans-eQTLs for L1MEb expression. However, those SNVs were not significant cis-eQTLs for RN5 and FKBPL expression, respectively. For trios involving STARD5, IL16, and EHMT2-AS1, neither the cis-eQTL nor the trans-eQTL were significant. We note that for most of these comparisons, although the two genotypes with the largest sample sizes were sufficient to establish a trending change in cis or trans expression, this trend was often broken by the third genotype with
spurious sample sizes. This suggests that replication in the Yoruba cohort may be limited by the small cohort sample size in the GEUVADIS project.

TE families and known TE-associated pathways are differentially regulated across L1 trans-eQTL variants

Though our eQTL analysis identified genetic variants associated with the expression of specific, evolutionarily older L1 subfamilies, we reasoned that there may be more global but subtle differences in TE expression profiles among genotype groups, given that TE expression is highly correlated [21]. Thus, for each gene-associated index SNV identified in the European eQTL analysis, we carried out differential expression analysis for all expressed genes and TEs (Supplementary Table S3A-S3C; Figure 3A). At the individual gene level, we detected few significant (FDR < 0.05) changes: 4 genes/TEs varied with rs11635336 genotype (IL16/STARD5), 4 genes/TEs varied with rs9271894 genotype (HLA), and 5 gene/TEs varied with rs1061810 genotype (HSD17B12) (Supplementary Table S3A-S3C). Importantly, however, these genes/TEs overlapped the genes/TEs identified in the cis- and trans-eQTL analyses, providing an algorithmically independent link among candidate SNV-gene-TE trios.

In contrast to gene-level analyses, Gene Set Enrichment Analysis (GSEA) provides increased sensitivity to subtle but consistent/widespread transcriptomic changes at the level of gene sets (e.g. TE families, biological pathways, etc.). Thus, we leveraged our differential expression analysis in combination with GSEA to identify repeat family and biological pathway gene sets impacted by SNV genotype in the
GEUVADIS dataset (Supplementary Table S3D-S3O; Figure 3A). Interestingly, changes in the genotype of rs11635336 (IL16/STARD5), rs9271894 (HLA), and rs1061810 (HSD17B12) were associated with an upregulation, upregulation, and downregulation, respectively, of multiple TE family gene sets (Figure 3B, Supplementary Table S3P). Differentially regulated TE family gene sets included DNA transposons, such as the hAT-Charlie family, and long terminal repeat (LTR) transposons, such as the endogenous retrovirus-1 (ERV1) family (Figure 3B, Supplementary Table S3P). Noteworthy, the L1 family gene set was the only TE gene set whose expression level was significantly altered across all three SNV analyses (Figure 3B, Supplementary Table S3P). Consistent with their relative significance in the L1 trans-eQTL analysis, the L1 family gene set was most strongly upregulated by alternating the IL16/STARD5 SNV (NES = 3.74, FDR = 6.43E-41), intermediately upregulated by alternating the HLA SNV (NES = 1.90, FDR = 7.19E-5), and least strongly changed by alternating the HSD17B12 SNV (NES = -1.57, FDR = 2.11E-2) (Figure 3C). We briefly note here that rs9270493, a clumped SNV linked to RNF5, was also linked to upregulation of the L1 family gene set (Supplementary Table S3Q-S3R). These results suggest that TE subfamily trans-eQTLs are associated with subtle but global differences in TE expression beyond a lone TE subfamily.

Next, we asked if other biological pathways were regulated concomitantly with TE gene sets in response to gene-linked index SNVs, reasoning that such pathways would act either upstream (as regulatory pathways) or downstream (as response pathways) of TE alterations. GSEA with the MSigDB Hallmark pathway gene sets [29,
identified 5 gene sets fitting this criterion, including “oxidative phosphorylation”, “mTORC1 signaling”, “fatty acid metabolism”, “adipogenesis”, and “cholesterol homeostasis” (Figure 3D, Supplementary Table S3S). Interestingly, several of these pathways or genes in these pathways have been implicated in TE regulation before. Rapamycin, which acts through mTORC1, has been shown to alter the expression of L1 and other repeats [31, 32]. Estrogens, which are involved in cholesterol and lipid metabolism, have been found to drive changes in repeat expression, and the receptors for both estrogens and androgens are believed to bind repeat DNA [32, 33]. Pharmacological inhibition of the mitochondrial respiratory chain and pharmacological reduction of endogenous cholesterol synthesis have also been shown to induce changes in L1 protein levels or repeat expression more broadly [34, 35]. GSEA with the GO Biological Process gene sets (Figure 3E, Supplementary Table S3T) and the Reactome gene sets (Figure 3F, Supplementary Table S3U) also identified several metabolism-related pathways including “ATP metabolic process”, “Generation of precursor metabolites and energy”, and “metabolism of amino acids and derivatives”. These results add to the catalogue of pathways associated with differences in L1 expression.

In our eQTL analysis, we also identified two orphan index SNVs, rs112581165 and rs72691418, to which we could not attribute a protein-coding gene mediator. To determine whether these SNVs also regulate any transposon families or biological pathways, we repeated the differential expression analysis (with all expressed genes and TEs) (Supplementary Table S4A-S4B) and the GSEA (Supplementary Table
S4C-S4J) with these SNVs (Supplementary Figure S7A). At the individual gene level, we detected 3193 genes/TEs that varied significantly (FDR < 0.05) with rs112581165 genotype and 1229 genes/TEs that varied significantly with rs72691418 genotype (Supplementary Table S4A-S4B). Similar to above, we next carried out GSEA to identify changes in functionally relevant gene sets. Like the gene-linked index SNVs, changes in the genotype of rs112581165 and rs72691418 were both associated with a downregulation and upregulation, respectively, of 10 TE families (Supplementary Figure S7B, Supplementary Table S4K). Noteworthy, the L1 family gene set was among the most strongly dysregulated TE family gene sets for both rs112581165 (NES = -4.32, FDR = 5.18E-89) and rs72691418 (NES = 4.01, FDR = 5.38E-79) (Supplementary Figure S7C). These results suggest that TE subfamily trans-eQTLs are associated with subtle differences in TE expression beyond the lone TE subfamily, even in the absence of a protein-coding gene cis-eQTL.

Like before, we asked if other biological pathways were regulated concomitantly with TE gene sets in response to orphan index SNVs. The top 10 Hallmark pathway gene sets identified by GSEA included gene sets that were previously identified (“oxidative phosphorylation”, “fatty acid metabolism”, and “mTORC1 signaling”), as well as several new pathways (Supplementary Figure S7D, Supplementary Table S4L). Among the new pathways, “DNA repair” [20] and the “P53 pathway” [36, 37] have also been linked to L1 control, and proteins in the “Myc targets v1” gene set interact with L1 ORF1p [38]. GSEA with the GO Biological Process gene sets (Supplementary Figure S7E, Supplementary Table S4M) and the Reactome gene sets (Supplementary
Figure S7F, Supplementary Table S4N) identified several metabolism-related pathways and several translation-related pathways, such as “cytoplasmic translation”, “eukaryotic translation initiation”, and “eukaryotic translation elongation”. Importantly, proteins involved in various aspects of proteostasis have been shown to be enriched among L1 ORF1p-interacting proteins [38]. Again, these results add to the catalogue of pathways associated with differences in TE expression, even in the absence of a candidate cis mediator.

Modulation of top candidate gene activity in a lymphoblastoid cell line induces small but widespread TE expression changes

We decided to validate the L1 regulatory properties of top candidate genes associated with L1 trans-eQTLs. For experimental purposes, we selected the GM12878 lymphoblastoid cell line, because (i) it is of the same cell type as the transcriptomic data used here for our eQTL analysis, and (ii) its epigenomic landscape and culture conditions have been well well-characterized as part of the ENCODE project [39, 40]. For validation purposes, we selected IL16, STARD5, HSD17B12, and RNF5 out of the 7 protein-coding gene candidates. We chose these genes for validation because the first 3 are associated with top trans-eQTL SNVs and the fourth one had very strong predicted mediation effects. To note, although GM12878 was part of the 1000Genomes Project, it was not included in the GEUVADIS dataset. However, based on its genotype, we can predict the relative expression of candidate regulators (Supplementary Figure S8A), which suggest that GM12878 may be most sensitive to modulations in IL16 and
STARD5 expression, given their relatively low endogenous expression. Interestingly, examination of the ENCODE epigenomic data in GM12878 cells [39] demonstrated that the region near the IL16/STARD5-linked index SNV (rs11635336) was marked with H3K4Me1 and H3K27Ac, regulatory signatures of enhancers (Supplementary Figure S8C). Similarly, the region near the HLA-linked index SNV (rs9271894) was marked with H3K4Me1, marked with H3K27Ac, and accessible by DNase, suggesting regulatory properties of the region as an active enhancer (Supplementary Figure S8C). These results further highlight the regulatory potential of the IL16-, STARD5-, and HLA-linked SNVs.

First, we decided to test the transcriptomic impact of overexpressing our top candidates in GM12878 LCLs. Cells were electroporated with overexpression plasmids (or corresponding empty vector), and RNA was isolated after 48h (Figure 4A, Supplementary Figure S9A). Differential expression analysis comparing control and overexpression samples confirmed the overexpression of candidate genes (Supplementary Figure S9B, Supplementary Table S5A-S5D). Intriguingly, we observed that IL16 was significantly upregulated following STARD5 overexpression (Supplementary Figure S9C, Supplementary Table S5B), although the inverse was not observed (Supplementary Figure S9C, Supplementary Table S5A), suggesting that IL16 may act downstream of STARD5. We note here that, consistent with the use of a high expression vector, the IL16 upregulation elicited by STARD5 overexpression (log₂ fold change = 0.45) was weaker than the upregulation from the IL16 overexpression (log₂ fold change = 1.89) (Supplementary Table S5A-S5B).
To further assess the biological relevance of each overexpression, we carried out GSEA using the GO Biological Process, Reactome pathway, and Hallmark pathway gene sets (Supplementary Table S5E-S5P). Importantly, GSEA using GO Biological Process and Reactome pathway gene sets highlighted differences that were consistent with the known biology of our candidate genes. Firstly, \textit{IL16} is involved in regulating T-cell activation, B-cell differentiation, and functions as a chemoattractant [41-46]. Moreover, it modulates macrophage polarization by regulating \textit{IL-10} expression [47]. \textit{IL16} overexpressing cells showed upregulation for “phagocytosis recognition” and “positive chemotaxis”, downregulation for “negative regulation of cell differentiation”, and downregulation for “Interleukin 10 signaling” (Figure 4B-4C). Secondly, \textit{STARD5} encodes a cholesterol transporter and is upregulated in response to endoplasmic reticulum (ER) stress [48-50]. \textit{STARD5} overexpressing cells showed downregulation of various cholesterol-related gene sets such as “sterol biosynthetic process”, “sterol metabolic process”, and “regulation of cholesterol biosynthesis by SREBP (SREBF)” (Figure 4D-4E). Thirdly, \textit{HSD17B12} encodes a steroid dehydrogenase involved in converting estrone into estradiol and is essential for proper lipid homeostasis [51-53]. \textit{HSD17B12} overexpressing cells showed downregulation of cholesterol-related gene sets, including “sterol biosynthetic process” and “regulation of cholesterol biosynthesis by SREBF (SREBP)” (Supplementary Figure S9D-S9E). Finally, \textit{RNF5} encodes an ER and mitochondrial-bound E3 ubiquitin-protein ligase that ubiquitin-tags proteins for degradation [54-57]. \textit{RNF5} overexpressing cells demonstrated alterations in gene sets involved in proteostasis and ER biology, including upregulation of “ERAD pathway”, \textit{CC-BY-NC-ND 4.0 International license available under a which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.
“response to endoplasmic reticulum stress”, and “intra-Golgi and retrograde Golgi-to-ER traffic” (Supplementary Figure S9F-S9G). These results suggest that our approach leads to biological changes consistent with the known biological impact of the genes being overexpressed.

Next, we sought to determine whether modulation of candidate genes had any impact on TE expression in general, and L1 in particular. Although there were no significant changes for individual TE subfamilies following IL16 and STARD5 overexpression (Supplementary Table S5A-S5B), we identified subtle but widespread upregulation of various TE families across both conditions by GSEA (Figure 4F, Supplementary Table S5Q-S5R). Interestingly, 7 families, including L1, ERV1, ERVL-MaLR, Alu, ERVL, TcMar-Tigger, and hAT-Charlie families, were commonly upregulated under both conditions (Figure 4F). In contrast, cells overexpressing HSD17B12 or RNF5 did not drive widespread changes in L1 family expression, as assessed by GSEA (Supplementary Table S5S-S5T). Noteworthy, the L1 family gene set was more strongly upregulated following STARD5 overexpression (NES = 2.25, FDR = 6.14E-7) compared to IL16 overexpression (NES = 2.24, FDR = 2.40E-5) (Figure 4G, Supplementary Table S5Q-S5R). Since IL16 is upregulated in response to STARD5 overexpression, this suggests that STARD5 may synergize with IL16 for the regulation of L1 transcription.

Then, we decided to further characterize the impact of IL16 activity on TEs, since (i) its overexpression led to a global upregulation of TE transcription, and (ii) it was itself...
upregulated in response to \textit{STARD5} overexpression, which also led to increased TE expression. Thus, since IL16 is a soluble cytokine, we independently assessed its regulatory properties by exposing GM12878 cells to recombinant human IL16 peptide [rhIL16] for 24 hours (\textbf{Figure 5A, Supplementary Figure S10A}). Differential gene expression analysis (\textbf{Supplementary Table S6A}) and comparison with the \textit{IL16} overexpression results demonstrated that differentially expressed genes were weakly but significantly correlated (\textbf{Supplementary Figure S10B}). Additionally, we carried out GSEA using the GO Biological Process, Reactome pathway, and Hallmark pathway gene sets (\textbf{Supplementary Table S6B-S6E}) and compared those results with the GSEA from the \textit{IL16} overexpression (\textbf{Supplementary Table S6F-S6H}). Consistent with the known biology of \textit{IL16}, GSEA highlighted a downregulation of many immune cell-related gene sets, including “leukocyte differentiation”, “mononuclear cell differentiation”, and “Interleukin-10 signaling” (\textbf{Figure 5B-5C, Supplementary Table S6F-S6H}). Like the overexpression results, exposure of GM12878 to rhIL16 for 24 hours led to an upregulation of an L1 family gene set by GSEA, although the effect was less pronounced than with the overexpression (\textbf{Figure 5D}). Even though treatment of GM12878 with rhIL16 for 48 hours exhibited known features of IL16 biology (\textbf{Supplementary Figure S10B-S10D, Supplementary Table S6J-S6Q}), the L1 upregulation was no longer detectable, though other TEs remained upregulated (\textbf{Supplementary Figure S10E, Supplementary Table S6Q}). These results further support the notion that \textit{IL16} acts as a modulator of L1 expression.
Finally, we sought to define the biological pathways regulated concomitantly with the L1 family gene set under all experimental conditions where it was upregulated (i.e., IL16 overexpression, STARD5 overexpression, and 24 hours of rhIL16 exposure) (Figure 6A, Figure 6B, Supplementary Table S7A). Again, we reasoned that such pathways would act either upstream (as regulatory pathways) or downstream (as response pathways) of TE alterations. GSEA with the Hallmark pathway gene sets identified 7 gene sets fitting this criterion, including “TNFα signaling via NF-κB”, “IL2 STAT5 signaling”, “inflammatory response”, “mTORC1 signaling”, “estrogen response early”, “apoptosis”, and “UV response up” (Figure 6C, Supplementary Table S7B). GSEA with the GO Biological Process gene sets (Figure 6D, Supplementary Table S7C) and the Reactome pathway gene sets (Figure 6E, Supplementary Table S7D) also identified MAPK signaling, virus-related pathways like “HCMV early events”, pathways involved in cell differentiation, and pathways involved in cholesterol and steroid metabolism like “signaling by nuclear receptors”. These results further cement the catalogue of pathways associated with differences in TE expression.

*L1 trans-eQTLs are co-associated with aging traits in GWAS databases.*

Although TE de-repression has been observed broadly with aging and age-related disease [5, 58], whether this de-repression acts as a causal driver, or a downstream consequence, of aging phenotypes remains unknown. We reasoned that if increased TE expression at least partially drives aging phenotypes, L1 trans-eQTLs
should be enriched for associations to aging traits in genome-wide association studies [GWAS] or phenome-wide association studies [PheWAS].

To test our hypothesis, we queried the Open Targets Genetics platform with our 499 trans-eQTL SNVs, mapped traits to standardized MeSH IDs, and then manually curated MeSH IDs related to aging-related traits (Figure 7A). Consistent with our hypothesis, a large proportion of L1 trans-eQTL SNVs (222/499 or 44.5%) were either (i) associated with an aging MeSH trait by PheWAS or (ii) LD-linked to a lead variant associated with an aging MeSH trait (Figure 7B). Moreover, among the 222 SNVs with significant aging-trait associations, we observed frequent mapping to more than a single age-related trait by PheWAS, with many SNVs associated with 10-25 traits (Figure 7C, Supplementary Table S8A). Additionally, many of the 222 SNVs mapped to 1-5 aging traits through a proxy lead variant (Figure 7D, Supplementary Table S8A). Among the most frequently associated or linked traits, we identified type 2 diabetes mellitus, hyperparathyroidism, thyroid diseases, coronary artery disease, hypothyroidism, and psoriasis, among many others (Figure 7E, Supplementary Table S8B).

As a parallel approach, we queried the Open Targets Genetics platform with our L1 trans-eQTL SNVs, as well as 500 combinations of random SNVs sampled from all SNVs used in the eQTL analyses. We then leveraged broader phenotype categories annotated by the platform, including 14 disease categories that we considered aging-related, to determine whether L1 eQTL associations were enriched for any disease categories (Supplementary Figure S11A). L1 eQTL associations were significantly
enriched (FDR < 0.05 and ES > 1) for 13 out of 14 disease categories, including cell proliferation disorders, immune system diseases, and musculoskeletal diseases (Supplementary Figure S11B-N). The cardiovascular diseases category was the only disease category for which we did not observe a significant enrichment (Supplementary Figure S11O). The enrichment for cell proliferation disorders is consistent with the associations of L1 activity with cellular senescence [12, 15] and cancer [59, 60]. The enrichment for immune system diseases is consistent with the role of L1 as a stimulator of the interferon pathway, inflammation, and senescence [15], as well as the more general notion that transposons can mimic viruses and stimulate immune responses from their hosts [61]. The enrichment for musculoskeletal diseases is consistent with an increase in L1 expression and copy number with age in muscle tissue from aging mice [11]. These results reinforce the notion that L1 activity is strongly and non-randomly associated with an assortment of age-related diseases.

Intriguingly, a large fraction of co-associated SNVs were on chromosome 6 near the HLA locus, which has previously been shown to be a hotspot of age-related disease traits [62]. Despite its association to our strongest L1 trans-eQTL SNV, little is known about the regulation and impact of IL16 during aging. One study, however, found that IL16 expression increases with age in ovarian tissue, and the frequency of IL16 expressing cells is significantly higher in ovarian tissue from women at early and late menopause, compared to premenopausal women [63]. Given these findings, and since L1 expression levels and copy number have been found to increase with age [5], we asked whether circulating IL16 levels may also change with age, using C57BL/6NJia
mice as a model (Figure 7F, Supplementary Table S8C). Consistent with the notion that increased IL16 levels may, at least partially, drive age-related TE de-repression, we observed a significant increase in circulating IL16 levels in female mice with age, and a trending increase with age in male mice (although the levels showed more animal-to-animal variability). By meta-analysis, circulating IL16 levels changed significantly with age across sexes (Figure 7F). These results further support the hypothesis that IL16 is involved in L1 biology and may modulate L1 age-related changes. In sum, our results provide one of the first pieces of evidence of a causal link between L1 expression levels and age-related decline.
In this work, we developed a pipeline to computationally identify candidate L1 transcriptional regulators by eQTL analysis. We provide experimental evidence for the involvement of top candidates in regulating L1 expression, demonstrating as a proof-of-principle that this approach can be broadly used on other large “omic”-characterized cohorts with human (i.e. GTEx [64, 65] or HipSci [66]) or mouse (i.e. DO mice [67]) subjects to identify other regulators of L1 activity. These datasets, combined with our approach, could be utilized to rigorously characterize conserved or group-specific TE regulatory mechanisms on multiple layers, such as across TE families (like Alu or ERVs), across cell or tissue types, across ancestry groups, and across species. This approach, which leverages existing datasets to perform in silico screening, could be a powerful method to expand our knowledge of TE regulation in non-diseased cells and tissues.

While we believe this approach can readily be applied to other datasets, we would like to note potential limitations with the approach implemented here, some of which were simply beyond the scope of this paper. Firstly, though it is common to use probabilistic estimation of expression residuals (PEER) [68] to enhance detection of cis-eQTLs, PEER was not implemented in our analysis as a precautionary measure, in order to avoid potentially blurring global TE signals, which likely led to a more conservative list of candidate cis gene mediators. Second, given the technical complexity in generating the vast amount of mRNA-seq data used for the eQTL analysis, it is possible that technical covariates introduced non-linear effects that would
not be easily removed by approaches like PEER or SVA [69]. For that reason, we opted to supplement our computational predictions with experimental data. Third, the L1 \textit{trans}-eQTLs identified were specific to older L1 subfamilies (L1P and L1M) and were not shared across subfamilies. One factor that may partially explain this is the heightened difficulty of quantifying the expression of evolutionarily younger L1 subfamilies using short-read sequencing [70]. More generally, significant single gene differences are often difficult to reproduce across studies, and it is for this reason that methods like GSEA were developed, to robustly identify broader changes in sets of genes [29]. Consistently, GSEA suggests that many TE families, beyond the single L1 subfamilies identified in the eQTL analysis, are differentially regulated among samples with different genotypes for \textit{trans}-eQTL SNVs and among samples where \textit{IL16/IL16} and \textit{STARD5} were manipulated. We note that although \textit{HLA} and \textit{HSD17B12} loci were significant in both the European and African cohorts, we were not able to independently identify all of the same candidate regulators. This is likely due to a combination of small sample size for the African cohort and the existence of population-specific L1 regulation. Future studies with larger sample sizes may be useful for expanding the catalogue of loci that are biologically meaningful for L1 expression across more than one population. Importantly, our computational scan is limited to loci exhibiting genomic variation among tested individuals. This will vary with factors like the ancestry groups of the populations being studied. Moreover, variants that confer extreme fitness defects may not exist at a sufficiently high level in a population so as to allow for an assessment of their involvement as eQTLs. Also, a potential “blindspot” of our current approach is that it does not distinguish between TE reads of intronic or intergenic origin. Intronic TE RNA
is often considered less interesting from a biological perspective since their presence may be attributed to readthrough transcription [71]. However, all analyses carried out in this study relied on polyA-selected libraries, which should be enriched for mature gene and transposon transcripts, minimizing the presence of readthrough transcribed L1 RNA. Additionally, even if L1-containing transcripts were quantified, these have been implicated in tightly controlled functions like T-cell quiescence maintenance [31], suggesting that intronic L1 RNA possesses biologically meaningful properties. Thus, this pipeline should help expand our understanding of L1 biology in either case. Finally, although we focused on protein-coding candidate regulators, it is possible that the non-coding genes identified in our scan may also causally drive differences in L1 expression. Though not explored here, other regulatory factors like small RNAs may also act as partial mediators. Since the GEUVADIS Consortium generated small RNA data in parallel to the mRNA data used in this study [26], in the future, our pipeline could be adapted to scan for cis small RNA mediators relatively easily. These unexplored factors may explain the associations between orphan SNV genotypes and TE family gene set changes.

Despite potential limitations, our approach identified IL16, STARD5, HLA-DRB5, HLA-DQA2, HSD17B12, RNF5, FKBPL, and EHMT2-AS1 as candidate L1 regulators in the European cohort. Moreover, the upregulation of several TE gene sets following IL16/IL16 and STARD5 manipulation highlights a causal role for these genes in TE control. Between these two genes, multiple lines of evidence suggest that STARD5 is the more potent mediator. First, the three-part integration statistics are more significant
for \textit{STARD5} than for \textit{IL16}. Second, the index SNV on chromosome 15 exhibited significant mediation effects through \textit{STARD5} but not \textit{IL16}; the most significant mediation effect for \textit{IL16} was linked to a clumped SNV. Third, the upregulation for the L1 family gene set was stronger under \textit{STARD5} overexpression than under \textit{IL16} overexpression. Importantly, we observed that \textit{IL16} was upregulated following \textit{STARD5} overexpression, but this upregulation was less than the upregulation from the \textit{IL16} overexpression. This suggests that \textit{IL16} can participate in the L1-regulating properties exerted by \textit{STARD5}. Moving forward, it will be informative to assess the effects of \textit{IL16} and \textit{STARD5} on L1 expression in other cell types. Interestingly, other genes like \textit{EHMT2} have previously been linked to retrotransposons [72, 73]. For the remaining genes that we experimentally tested, we note that GM12878 is predicted to have relatively high endogenous \textit{HSD17B12} expression and intermediate \textit{RNF5} expression, based on the GM12878 genotypes at \textit{cis}-eQTLs for these genes. Given these expression patterns, GM12878 may not be sensitive to overexpression of \textit{RNF5}, and especially insensitive to \textit{HSD17B12} overexpression. For these two candidates, cells may be more sensitive to knockdown- or knockout-based approaches. Indeed, \textit{HSD17B12} is essential for mouse development, \textit{HSD17B12} knockout in adult mice results in reduced body weight and liver toxicity, and knockdown of the \textit{Caenorhabditis elegans} ortholog for \textit{HSD17B12} reduces lipid stores and blocks induction of the unfolded protein response of the endoplasmic reticulum [52, 74, 75]. Given that TEs are often derepressed when homeostasis is challenged [61], such as following \textit{HSD17B12} knockout/knockdown, it remains possible that \textit{HSD17B12} possesses L1 regulatory properties that were not detectable by our approach. Future work could avert such
technical limitations by testing the impact of candidate genes by both up- and down-
regulation, or by selecting LCL samples with endogenous target gene expression levels
that would be most sensitive to our overexpression approach.

As another, theoretical line of evidence for the potential involvement of candidate
genes in L1 regulation, we highlight known interactions between tested candidate genes
and viral infections, which may be relevant under conditions where transposons are
recognized as viral mimics [61]. Indeed, *IL16* has been extensively studied for its ability
to inhibit human immunodeficiency virus (HIV) replication, partly by suppressing mRNA
expression [76-78]. Additionally, but in contrast to its HIV-suppressive properties, *IL16*
can enhance the replication of influenza A virus (IAV) and facilitate its infection of hosts,
potentially through its repression of type I interferon beta and interferon-stimulated
genes [79]. *IL16* can also contribute to the establishment of lifelong gamma herpesvirus
infection [80]. *STARD5* is another candidate implicated in the influenza virus replication
cycle [81]. *HSD17B12* promotes the replication of hepatitis C virus via the very-long-
chain fatty acid (VLCFA) synthesis pathway and the production of lipid droplets
important for virus assembly [82, 83]. Additionally, HSD17B12 has been found
interacting with the coronavirus disease 2019 (COVID-19) protein nonstructural protein
13 (NSP13), which is thought to antagonize interferon signaling [84]. Finally, *RNF5* has
been implicated in both promoting and antagonizing severe acute respiratory syndrome
coronavirus 2 (SARS-CoV-2) by either stabilizing the interactions of membrane protein
(M) [85] or inducing degradation of structural protein envelope (E) [86], respectively.
Fundamentally, *RNF5* regulates virus-triggered interferon signaling by targeting the
stimulator of interferon genes (STING) or mitochondrial antiviral signaling protein (MAVS) for ubiquitin-mediated protein degradation [56, 57]. These studies reinforce the roles of tested candidate regulators in virus-associated processes, including interferon-mediated signaling.

Consistent with the notion that L1 is associated with aging and aging phenotypes [5, 58], we observed that L1 trans-eQTL SNVs were associated with aging phenotypes in GWAS/PheWAS databases. This is very surprising, but interesting, given that all 1000Genomes Project participants declared themselves to be healthy at the time of sample collection. Assuming this to be true, our results suggest that L1 expression differences exist in natural, healthy human populations, and these expression differences precede onset of aging diseases. Though it is often unclear whether L1 misregulation is a consequence or driver of aging phenotypes, our results suggest that L1 levels may drive aging phenotypes. As we continue to expand the catalogue of L1 regulators, especially in healthy cells and tissues, the L1 regulatory processes that are disrupted over the course of aging will become increasingly clear. To that end, this work may serve as a guide for conducting more comprehensive scans for candidate TE regulators.
CONCLUSIONS

We developed an eQTL-based pipeline that leverages genomic and transcriptomic data to scan the human genome for novel candidate regulators of L1 subfamily expression. Though the initial scan identified genetic variants associated with expression differences in specific L1 subfamilies, secondary analyses by GSEA suggest that genetic variants are associated with subtle but global differences in the expression of many TE families. Our pipeline identified candidate genes, including HSD17B12 and HLA genes, that likely play a conserved role in L1 regulation across human populations of different ancestries. Though some top candidates from the European cohort scan, such as IL16, STARD5, and RNF5, were not significant in the African cohort analysis, it is likely that some of these genes would appear in cross-ancestry scans with larger samples sizes. We detected subtle but global differences in L1 family expression following IL16 overexpression, STARD5 overexpression, and rhIL16 treatment for 24 hours, further suggesting that some candidate genes have regulatory potential. We generate a list of pathways, such as mTORC1 signaling and cholesterol metabolism, that may act upstream of L1 expression. Finally, the co-association of some genetic variants with both L1 expression differences and various age-related diseases suggests that L1 differences may precede and contribute to the onset of disease. Our results expand the potential mechanisms by which L1 expression is regulated and by which L1 may influence aging-related phenotypes.
METHODS

Publicly available data acquisition

The eQTL analysis was carried out on 358 European (EUR) individuals and 86 Yoruban (YRI) individuals for which paired single nucleotide variant, structural variant, and transcriptomic data were available from Phase 3 of the 1000 Genomes Project [22, 23] and from the GEUVADIS consortium [26]. Specifically, Phase 3 autosomal SNVs called on the GRCh38 reference genome were obtained from The International Genome Sample Resource (IGSR) FTP site (http://ftp.1000genomes.ebi.ac.uk/vol1/ftp/data_collections/1000_genomes_project/release/20190312_biallelic_SNV_and_INDEL/). Structural variants were also obtained from the IGSR FTP site (http://ftp.1000genomes.ebi.ac.uk/vol1/ftp/phase3/integrated_sv_map/). mRNA-sequencing fastq files generated by the GEUVADIS consortium were obtained from ArrayExpress under accession E-GEUV-1.

Aggregating and pre-processing genotype data for eQTL analyses

To prepare SNVs for association analyses, all SNVs were first annotated with rsIDs from dbSNP build 155 using BCFtools v1.10.2 [87]. VCFtools v0.1.17 [88] was then used to remove indels and keep variants with the following properties in each of the two populations: possessed a minimum and maximum of two alleles, possessed a minor allele frequency (MAF) of at least 1%, passed Hardy-Weinberg equilibrium thresholding at $p < 1 \times 10^{-6}$, with no missing samples, and located on an autosome. We
note here that sex chromosomes were not included in the analysis since (i) Y chromosome SNVs were not available and (ii) analyses with X chromosome SNVs require unique algorithms and cannot simply be incorporated into traditional association pipelines [89, 90]. VCF files containing these filtered SNVs were then converted to PLINK BED format using PLINK v1.90b6.17 [91], keeping the allele order. PLINK BED files were subsequently used to generate preliminary 0/1/2 genotype matrices using the ‘--recodeA’ flag in PLINK. These preliminary matrices were manipulated in terminal, using the gcut v9.0 function to remove unnecessary columns and datamash v1.7 to transpose the data, to generate the final 0/1/2 matrices used for the eQTL analyses. Finally, PLINK was used to prune the list of filtered SNVs, using the “--indep-pairwise 50 10 0.1” flag, and to generate principal components (PCs) from the pruned genotypes.

To control for inter-individual differences in genomic transposon copy number load, we applied 1 of 2 approaches, depending on the analysis. For approach 1, the net number of L1 and Alu insertions was quantified across the 444 samples. We chose to aggregate the L1 and Alu copy numbers, since Alu relies on L1 machinery for mobilization [92], and so the aggregate number may provide a finer view of L1-associated copy number load. Briefly, VCFTools was used to extract autosomal structural variants from the 1000Genomes structural variant calls. L1 and Alu insertions and deletions were then extracted with BCFtools by keeping entries with the following expressions: ‘SVTYPE=”LINE1”’, ‘SVTYPE=”ALU”’, ‘SVTYPE=”DEL_LINE1”’, and ‘SVTYPE=”DEL_ALU”’. The resulting VCF files were then transformed to 0/1/2 matrices in the same manner as the SNVs. A net copy number score was obtained for each
sample by adding the values for the L1 and Alu insertions and subtracting the values for the L1 and Alu deletions. For approach 2, the complete structural variant matrix was filtered with VCFtools using the same parameters as with the SNV matrices. The filtered structural variant matrix was then pruned with PLINK, and these pruned structural variant genotypes were used to generate principal components, in the same fashion as with the SNV matrix. The net copy number score or the structural variant principal components, depending on the analysis, were included as covariates.

**mRNA-seq read trimming, mapping, and quantification**

Fastq files were first trimmed using fastp v0.20.1 [93] with the following parameters: detect_adapter_for_pe, disable_quality_filtering, trim_front1 17, trim_front2 17, cut_front, cut_front_window_size 1, cut_front_mean_quality 20, cut_tail, cut_tail_window_size 1, cut_tail_mean_quality 20, cut_right, cut_right_window_size 5, cut_right_mean_quality 20, and length_required 36. Read quality was then inspected using fastqc v0.11.9.

Next, the GRCh38 primary human genome assembly and comprehensive gene annotation were obtained from GENCODE release 33 [94]. Since LCLs are generated by infecting B-cells with Epstein-Barr virus, the EBV genome (GenBank ID V01555.2) was included as an additional contig in the human reference genome. The trimmed reads were aligned to this modified reference genome using STAR v2.7.3a [95] with the following parameters: outFilterMultimapNmax 100, winAnchorMultimapNmax 100, and
outFilterMismatchNoverLmax 0.04. Finally, the TEcount function in the TEtranscripts v2.1.4 [27] package was employed to obtain gene and TE counts, using the GENCODE annotations to define gene boundaries and a repeat GTF file provided on the Hammell lab website (downloaded on February 19 2020 from https://labshare.cshl.edu/shares/mhammelllab/www-data/TEtranscripts/TE_GTF/GRCh38_GENCODE_rmsk_TE.gtf.gz) to define repeat boundaries.

**Gene cis-eQTL and L1 trans-eQTL analyses**

Gene and TE count files were loaded into R v4.2.1. Lowly expressed genes were first filtered out if 323/358 European samples and 78/86 Yoruban samples did not have over 0.44 counts per million (cpm) or 0.43 cpm, respectively. These fractions were selected because they corresponded to expression in ~90% of samples and thus helped maintain maximal statistical power by focusing on genes ubiquitously expressed across each entire population. The cpm thresholds were selected because they corresponded to 10 reads in the median-length library within each set of samples.

Then, counts underwent a variance stabilizing transformation (vst) using DESeq2 v1.36.0 [96]. The following covariates were regressed out from vst normalized expression data using the ‘removeBatchEffect’ function in Limma v3.52.2 [97]: lab, population category, principal components 1-2 of the pruned SNVs, biological sex, net L1/Alu copy number, and EBV expression levels. Since the Yoruban samples were all
from the same population, the population variable was omitted in their batch correction.

Here, we note several things. First, EBV expression was included as a covariate because heightened TE expression is often a feature of viral infections [98]. Secondly, although PEER [68] is often used to remove technical variation for cis-eQTL analysis, this can come at the expense of correcting out genome-wide biological effects. This can be problematic in some settings, such as trans-eQTL analysis. Thus, PEER factors were not included. The batch-corrected data underwent a final inverse normal transformation (INT), using the RankNorm function in the R package RNOmni v1.0.1, to obtain normally distributed gene expression values.

The INT expression matrices were split into genes and L1 subfamilies, which were used to identify gene cis-eQTLs and L1 subfamily trans-eQTLs in the European superpopulation using MatrixEQTL v2.3 [99]. For gene cis-eQTLs, SNVs were tested for association with expressed genes within 1 million base pairs. We opted to use a trans-eQTL approach using aggregate subfamily-level TE expression since the trans approach should allow us to identify regulators of many elements rather than one. The Benjamini-Hochberg false discovery rate (FDR) was calculated in each analysis, and we used the p-value corresponding to an FDR of < 5% as the threshold for eQTL significance. In addition, the cis-eQTL and trans-eQTL analyses were also repeated using 20 permuted expression datasets in which the sample names were scrambled, and the p-value corresponding to an average empirical FDR of < 5% was used as a secondary threshold. To note, we calculated the average empirical FDR at a given p-value $p_i$ by (i) counting the total number of null points with $p \leq p_i$, (ii) dividing by the
number of permutations, to obtain an average number of null points with $p \leq p_i$, and (iii) dividing the average number of null points with $p \leq p_i$ by the number of real points with $p \leq p_i$. eQTLs were called as significant if they passed the stricter of the two thresholds.

SNV-gene and SNV-L1 associations that were significant in the European superpopulation were then targeted and tested in the Yoruban population using R's built-in linear modelling functions. In this case, only the Benjamini-Hochberg FDR was calculated, and significant eQTLs were called if they possessed an FDR < 5%.

Defining SNV-gene-L1 trios and mediation analysis

For each population, the *cis*- and *trans*-eQTL results were integrated to identify SNVs associated with both gene and L1 subfamily expression. We reasoned that L1 expression would respond to differences in expression of *bona fide* regulators. Consequently, gene expression and L1 subfamily expression associations were assessed by linear regression, and the p-values from this analysis were Benjamini-Hochberg FDR-corrected. Candidate SNV-gene-L1 trios were defined as those with *cis*-eQTL, *trans*-eQTL, and expression regression FDRs < 5%. To identify top, index SNVs in regions of linkage disequilibrium (LD), SNVs within 500 kilobases of each other with an $R^2 > 0.10$ were clumped together by *trans*-eQTL p-value using PLINK v1.90b6.17. Mediation analysis was carried out using the ‘gmap.gpd’ function in eQTLMAPT v0.1.0 [100] on all candidate SNV-gene-L1 trios. Empirical p-values were calculated using 30,000 permutations, and Benjamini-Hochberg FDR values were calculated from
empirical p-values. Mediation effects were considered significant for trios with FDR < 5%.

Differential expression analysis across trans-eQTL SNV genotypes

Transcriptomic changes associated with alternating the allele of each SNV of interest were evaluated using DESeq2 v1.36.0. Using the same filtered counts prepared for the eQTL analysis, a linear model was constructed with the following covariates for each SNV: SNV genotype in 0/1/2 format, biological sex, lab, population category, principal components 1-2 of the pruned SNVs, and principal components 1-3 of the pruned SVs (to account for structural variant population structure). As before, the population label was omitted from the Yoruban population analysis. Significant genes and TEs were those with an FDR < 5%.

Functional enrichment analyses

We used the Gene Set Enrichment Analysis (GSEA) paradigm as implemented in the R package clusterProfiler v4.4.4 [101]. Gene Ontology, Reactome, and Hallmark pathway gene sets were obtained from the R package msigdb v7.5.1, an Ensembl ID-mapped collection of gene sets from the Molecular Signature Database [29, 30]. Additionally, TE subfamilies were aggregated into TE family gene sets using the TE family designations specified in the TE GTF file (downloaded on February 19 2020 from https://labshare.cshl.edu/shares/mhammellab/www-
data/TEtranscripts/TE_GTF/GRCh38_GENCODE_rmsk_TE.gtf.gz) used during the RNA-seq quantification step. The DESeq2 v1.36.0 Wald-statistic was used to generate a combined ranked list of genes and TEs for functional enrichment analysis. All gene sets with an FDR < 5% were considered significant. For plots with a single analysis, the top 5 downregulated and top 5 upregulated gene sets were plotted, at most. For plots with multiple analyses, shared gene sets with the desired expression patterns in each individual analysis were first identified. Then, the p-values for shared gene sets were combined using Fisher’s method, and this meta-analysis p-value was used to rank shared gene sets. Finally, the top 5 gene sets with one expression pattern and the top 5 gene sets with the opposite expression pattern were plotted. If there were less than 5 gene sets in either group, those were replaced with gene sets exhibiting the opposite regulation, in order to plot 10 shared gene sets whenever possible.

**Cell lines and cell culture conditions.**

GM12878 (RRID: CVCL_7526) lymphoblastoid cells were purchased from the Coriell Institute. We opted to use GM12878 as a well-characterized representative cell line for candidate validation, given that (i) it is of the same cell type as the transcriptomic data used here for our eQTL analysis, and (ii) its epigenomic landscape and culture conditions are well-characterized as part of the ENCODE project [39, 40].

GM12878 cells were maintained in RPMI (Corning cat. 15-040-CV) containing 15% FBS and 1X Penicillin-Streptomycin-Glutamine (Corning cat. 30-009-CI).
were cultured in a humidified incubator at 37°C and 5% CO₂, subculturing cells 1:5 once cells reached a density of ~10⁶ mL⁻¹. All cells used were maintained below passage 30 and routinely tested for mycoplasma contamination using the PlasmoTest Mycoplasma Detection Kit (InvivoGen).

**Plasmids**

The empty pcDNA3.1(+) backbone (Invitrogen cat. V79020) was a kind gift from the lab of Dr. Changhan David Lee at the University of Southern California Leonard Davis School of Gerontology. Overexpression vectors for **IL16** (CloneID OHu48263C), **STARD5-FLAG** (CloneID OHu07617D), **HSD17B12-FLAG** (CloneID OHu29918D), and **RNF5-FLAG** (CloneID OHu14875D) on a pcDNA3.1 backbone were purchased from GenScript. Plasmid sequences were verified for accuracy using Plasmidsaurus’s whole plasmid sequencing service.

**Transfections**

*Escherichia coli* were cultured in LB Broth (ThermoFischer Scientific) supplemented with 50 μg/mL carbenicillin to an optical density 600 (OD₆₀₀) of 2 – 4. Plasmid extractions were carried out using the Nucleobond Xtra Midi Plus EF kit (Macherey-Nagel) following manufacturer recommendations. Plasmids were aliquoted and stored at -20°C until the time of transfection. On the day of transfection, GM12878 cells were collected in conical tubes, spun down (100xG, 5 minutes, room temperature),
resuspended in fresh media, and counted by trypan blue staining using a Countess II FL automated cell counter (Thermo Fisher). The number of cells necessary for the experiment were then aliquoted, spun down, and washed with Dulbecco’s phosphate-buffered saline (DPBS) (Corning, cat. #21-031-CV).

GM12878 cells were transfected by electroporation using the Neon Transfection System (Invitrogen) with the following parameters: 1200 V, 20 ms, and 3 pulses for GM12878 cells in Buffer R. Per reaction, we maintained a plasmid mass:cell number ratio of 10 μg : 2*10^{6} cells. For mRNA-sequencing, 8*10^{6} GM12878 cells were independently transfected for each biological replicate, with 4 replicates per overexpression condition, and cultured in a T25 flask. Immediately after transfection, cells were cultured in Penicillin-Streptomycin-free media for ~24 hours.

Afterwards, to promote selection of viable and healthy transfected GM12878 cells, we enriched for viable cells using the EasySep Dead Cell Removal (Annexin V) Kit (STEMCELL Technologies) before seeding 2*10^{6} live cells in the same media used for cell maintenance. After another 24 hours, cell viability was measured by trypan blue staining on a Countess automated cell counter and cells were spun down (100xG, 5 min, room temperature) and lysed in TRIzol Reagent (Invitrogen) for downstream total RNA isolation (see below).

Recombinant human IL16 (rhIL16) peptide treatment
Human rIL16 was obtained from PeproTech (cat. #200-16) and resuspended in 0.1% bovine serum albumin (BSA) solution (Akron, cat. #AK8917-0100). GM12878 cells were seeded at a concentration of 500,000 live cells per mL of media on 6-well suspension plates with 3 independent replicates per condition. Cells were exposed to 0, 24, or 48 hours of 100 ng mL\(^{-1}\) of rhIL16. To replace or exchange media 24 hours after seeding, cells were transferred to conical tubes, spun down (100xG, 5 min, room temperature), resuspended in 5 mL of the appropriate media, and transferred back to 6-well suspension plates. After 48 hours, cell viability was measured by trypan blue staining and cells were spun down (100xG, 5 min, room temperature) and lysed in TRIzol Reagent (Invitrogen).

**RNA extractions and mRNA sequencing**

RNA was extracted using the Direct-zol RNA Miniprep kit (Zymo Research) following manufacturer recommendations. The integrity of RNA samples was evaluated using an Agilent High Sensitivity RNA ScreenTape assay (Agilent Technologies), ensuring that all samples had a minimum eRIN score of 8 before downstream processing. We then submitted total RNA samples to Novogene (Sacramento, California) for mRNA library preparation and sequencing on the NovaSeq 6000 platform as paired-end 150 bp reads.

**Analysis of overexpression and rhIL16 exposure mRNA-seq**
mRNA-seq reads were trimmed, mapped, and quantified like for the eQTL analysis, except for the overexpression sample data. For this data, one modification was made: the EBV-inclusive reference genome was further modified to include the pcDNA3.1 sequence as an additional contig. Lowly expressed genes were filtered using a cpm threshold as in the eQTL processing, but that cpm threshold had to be satisfied by as many samples as the size of the smallest biological group. For the overexpression data, surrogate variables were estimated with the ‘svaseq’ function [69] in the R package ‘sva’ v3.44.9, and they were regressed out from the raw read counts using the ‘removeBatchEffect’ function in the R package Limma v3.52.2. DESeq2 was used to identify significantly (FDR < 5%) differentially expressed genes and TEs between groups. Functional enrichment analysis was carried out as previously described.

PheWAS analysis

To gather the known associated traits for the 499 TE-related SNVs, we used Open Targets Genetics (https://genetics.opentargets.org/), a database of GWAS summary statistics [102]. First, we queried the database using the 499 TE-related SNVs and collected traits that were directly associated (with P < 5x10^{-8}) with the SNVs, as well as traits associated with lead variants that were in linkage disequilibrium (LD) with the queried SNPs (with R^2 > 0.6). For age-related traits (ARTs), we used the comprehensive list of 365 Medical Subject Headings (MeSH) terms reported by [103] (downloaded from https://github.com/kisudsoe/Age-related-traits). To identify known age-related traits, the known associated traits were translated into the equivalent MeSH
terms using the method described by [103]. Then, the MeSH-translated known
associated traits for the 499 TE-related SNVs were filtered by the MeSH terms for age-
related traits.

As a parallel approach, we mapped the RsIDs for all SNVs used during the eQTL analyses to their corresponding bi-allelic Open Targets variant IDs, when available. The variant IDs corresponding to L1 trans-eQTL SNVs were extracted, and 500 different equal-length combinations of random SNVs were generated. Next, we queried the Open Targets database using the lists of L1-associated and random SNVs and collected the associated traits (with \( P < 5 \times 10^{-5} \)). Importantly, the database assigns traits to broader categories, including 14 disease categories that we considered age-related. We counted the number of L1-associated or random SNVs mapping to each category, and we used the random SNV counts to generate an empirical cumulative distribution function (ecdf) for each category. We calculated enrichment p-values using the formula \( p = 1 - \text{ecdf}(\text{mapped eQTLs}) \) and then Benjamini-Hochberg FDR-corrected all p-values. An enrichment score (ES) was also calculated for each category using the formula \( \text{ES} = \frac{\text{number of mapped L1 eQTLs}}{\text{median number of randomly mapping SNVs}} \). Categories with an ES > 1 and FDR < 0.05 were considered significantly enriched.

**Quantification of mouse serum IL16 by ELISA**

Serum was collected from male and female C57BL/6JNia mice (4-6 and 20-24 months old) obtained from the National Institute on Aging (NIA) colony at Charles
Rivers. All animals were euthanized between 8-11 am in a “snaking order” across all groups to minimize batch-processing confounds due to circadian processes. All animals were euthanized by CO₂ asphyxiation followed by cervical dislocation. Circulating IL16 levels were quantitatively evaluated from mouse serum by enzyme-linked immunosorbent assay (ELISA). Serum was diluted 1/10 before quantifying IL16 concentrations using Abcam’s Mouse IL-16 ELISA Kit (ab201282) in accordance with manufacturer instructions. Technical replicates from the same sample were averaged to one value before statistical analysis and plotting. P-values across age within each sex were calculated using a non-parametric 2-sided Wilcoxon test, and p-values from each sex-specific analysis were combined using Fisher’s method.
DECLARATIONS

Ethics approval and consent to participate

All animals were treated and housed in accordance with the Guide for Care and Use of Laboratory Animals. All experimental procedures were approved by the University of Southern California’s Institutional Animal Care and Use Committee (IACUC) and are in accordance with institutional and national guidelines. Samples were derived from animals on approved IACUC protocol #20770.

Availability of data and materials

Sequencing data generated in this study is accessible through the Sequence Read Archive (SRA) under BioProject PRJNA937306. All code is available on the Benayoun lab GitHub (https://github.com/BenayounLaboratory/TE-eQTL_LCLs). Analyses were conducted using R version 4.2.1. Code was re-run independently on R version 4.3.0 to check for reproducibility.

Competing interests

The authors declare that they have no competing interests.

Funding

This material is based upon work supported by the National Science Foundation Graduate Research Fellowship Program (NSF GRFP) under Grant No. DGE-1842487 (to J.I.B.), the National Institute on Aging (NIA) under Grant No. T32 AG052374 (to
J.I.B.), the University of Southern California with a Provost Fellowship (to J.I.B.), the
USC Gerontology Enriching MSTEM (GEMSTEM) to Enhance Diversity in Aging
Program under Grant No. R25 AG076400 (to C.R.M.), and the National Institute of
General Medical Sciences (NIGMS) under Grant No. R35 GM142395 (to B.A.B).

Authors’ contributions

J.I.B. and B.A.B designed the study. J.I.B., L.Z., and S.K. performed data
analyses, with guidance from Y.S. and B.A.B. J.I.B. and C.R.M. carried out
experiments. J.I.B., B.A.B., S.K., and Y.S. wrote the manuscript. All authors contributed
to the editing of the manuscript.

Acknowledgements

We would like to thank Prof. Rachel Brem for her feedback and insights on the
eQTL analyses. We would also like to thank Dr. Minhoo Kim for her feedback on the
manuscript. Some panels were created with BioRender.com.
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FIGURE LEGENDS

Figure 1. Overview of the pipeline developed to scan for L1 transcriptional regulators in silico.

(A) An illustration of the samples and “omic” data used in this study. Of the 358 European individuals, 187 were female and 171 were male. Of the 86 African individuals, 49 were female and 37 were male. (Note that Utah subjects are of Northern European ancestry, and thus part of the European cohort for analytical purposes). (B) A schematic illustrating how genetic variants, gene expression, and TE expression can be integrated to identify highly correlated SNV-Gene-TE trios. (C) A Manhattan plot for the L1 subfamily trans-eQTL analysis in the European cohort. The genes that passed our three-part integration approach are listed next to the most significant trans-eQTL SNV they were associated with in cis. The dashed line at p = 3.44E-8 corresponds to an average empirical FDR < 0.05, based on 20 random permutations. One such permutation is illustrated in the bottom panel. The solid line at p = 2.31E-8 corresponds to a Benjamini-Hochberg FDR < 0.05. The stricter of the two thresholds, p = 2.31E-8, was used to define significant trans-eQTLs. FDR: False Discovery Rate. Some panels were created with BioRender.com.

Figure 2. Identification of 1st tier candidate L1 expression regulators in the European cohort.

(A) A schematic for how 1st tier candidate genes were defined. In short, these were genes in trios with index SNVs that were at the top of their respective peak. (B) The
three-part integration results for three protein-coding genes—STARD5, IL16, HSD17B12—that we considered first tier candidates for functional, in vitro testing. In the left column are the trans-eQTLs, in the middle column are the cis-eQTLs, and in the right column are the linear regressions for gene expression against L1 subfamily expression. Expression values following an inverse normal transform (INT) are shown. The FDR for each analysis is listed at the top of each plot. FDR: False Discovery Rate.

**Figure 3.** L1 trans-eQTLs are associated with subtle, widespread differences in TE families and known TE-associated pathways.

(A) Scheme for functionally annotating gene-linked index SNVs by GSEA. (B) GSEA analysis for shared, significantly regulated TE family gene sets across genotypes for rs11635336 (IL16/STARD5), rs9271894 (HLA), and rs1061810 (HSD17B12). (C) GSEA plots for the L1 family gene set results summarized in (B). For these plots, the FDR value is listed. GSEA analysis for top, shared, concomitantly regulated (D) MSigDB Hallmark pathway, (E) GO Biological Process, and (F) Reactome pathway gene sets across genotypes for rs11635336 (IL16/STARD5), rs9271894 (HLA), and rs1061810 (HSD17B12). Shared gene sets were ranked by combining p-values from each individual SNV analysis using Fisher’s method. In each bubble plot, the size of the dot represents the -log10(FDR) and the color reflects the normalized enrichment score. FDR: False Discovery Rate.

**Figure 4.** Impact of IL16 and STARD5 overexpression on LCL gene and TE expression landscapes.
**IL16** and **STARD5** overexpression induce changes consistent with their known biology, as well as subtle but widespread upregulation of TE families. (A) Scheme for experimentally validating the roles of **IL16** and **STARD5** in L1 regulation. GSEA analysis for top, differentially regulated (B) GO Biological Process and (C) Reactome pathway gene sets following **IL16** overexpression. GSEA analysis for top, differentially regulated (D) GO Biological Process and (E) Reactome pathway gene sets following **STARD5** overexpression. (F) GSEA analysis for shared, significantly regulated TE family gene sets following **IL16** and **STARD5** overexpression. (G) GSEA plots for the L1 family gene set results summarized in (F). For these plots, the FDR value is listed. In each bubble plot, the size of the dot represents the $-\log_{10}(FDR)$ and the color reflects the normalized enrichment score. FDR: False Discovery Rate. Some panels were created with BioRender.com.

**Figure 5.** rhIL16 treatment is sufficient to transiently upregulate an L1 family gene set.

(A) Scheme for experimentally validating the role of rhIL16 in L1 regulation. GSEA analysis for top, shared, concomitantly regulated (B) GO Biological Process and (C) Reactome pathway gene sets following **IL16** overexpression and rhIL16 exposure for 24 hours. Shared gene sets were ranked by combining p-values from each individual treatment analysis using Fisher’s method. (D) GSEA analysis for top, differentially regulated TE family gene sets following rhIL16 exposure for 24 hours. The GSEA plot for the L1 family gene set result summarized in the bubble plot is also shown. For this plot, the FDR value is listed. In each bubble plot, the size of the dot represents the -
\[
\log_{10}(\text{FDR}) \quad \text{and the color reflects the normalized enrichment score. FDR: False Discovery Rate. Some panels were created with BioRender.com.}
\]
PheWAS. (D) Histogram depicting the distribution of number of aging MeSH traits linked with the 222/499 SNVs through a proxy lead SNP in LD with the SNVs. (E) A diagram highlighting the organ targets of the top 10 most frequently associated aging traits. (F) The concentrations of circulating IL16 in aging mice of both sexes was assessed by ELISA. Significance across age in each sex was assessed using a Wilcoxon test. The p-values from each sex (females in pink and males in blue) were combined by meta-analysis using Fisher’s method. Any p-value < 0.05 was considered significant. Some panels were created with BioRender.com.
Figure 1

A  Samples and data used in this study

B  Scheme for in silico identification of candidate transposon-subfamily regulators

C  Manhattan plots for European L1 trans-eQTL analysis
**Figure 2**

**A** Scheme for defining 1st tier candidate genes

*Scan for significant L1 trans-eQTLs*

*Clump trans-eQTL SNVs by p-value and define index SNVs*

*Identify genes linked to index SNVs*

**B** In silico screen results for 1st tier candidate regulators

*rs11635336*

*rs1061810*

*Gene Scan for significant L1 trans-eQTLs*

*Clump trans-eQTL SNVs by p-value*

*Identify genes linked to index SNVs*
Figure 3

A Scheme for functionally annotating SNVs

B Index SNV associations with TE family gene sets

C Index SNV associations with an L1 family gene set

D Index SNV associations with MSigDB Hallmark gene sets

E Index SNV associations with GO Biological Process gene sets

F Index SNV associations with Reactome gene sets
Figure 4

A) Scheme for assessing the effects of candidate genes on L1 expression

- Extract plasmid DNA
- Electroporation with GM12878 LCLs
- Remove dead cells
- Pellet cells and extract RNA
- Sequence mRNA

B) Top enriched GO Biological Processes following IL16 overexpression

<table>
<thead>
<tr>
<th>GO Biological Processes</th>
<th>NES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regulation of response to extracellular stimulus</td>
<td>5.0</td>
</tr>
<tr>
<td>Phagocytosis recognition</td>
<td>-2.5</td>
</tr>
<tr>
<td>Positive chemotaxis</td>
<td>0.0</td>
</tr>
<tr>
<td>Complement activation</td>
<td>-2.5</td>
</tr>
<tr>
<td>Interleukin-12 production</td>
<td>-5.0</td>
</tr>
<tr>
<td>MAPK cascade</td>
<td>-log(FDR)</td>
</tr>
<tr>
<td>ERK1 and ERK2 cascade</td>
<td>1.5</td>
</tr>
<tr>
<td>Negative regulation of cell differentiation</td>
<td>2.0</td>
</tr>
<tr>
<td>Sterol biosynthetic process</td>
<td>2.5</td>
</tr>
<tr>
<td>Gliogenesis</td>
<td>3.0</td>
</tr>
</tbody>
</table>

IL16 OE

C) Top enriched Reactome gene sets following IL16 overexpression

<table>
<thead>
<tr>
<th>Reactome Pathways</th>
<th>NES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Creation of C4 and C2 activators</td>
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</tr>
<tr>
<td>SRP-dependent cotranslational protein targeting to membrane</td>
<td>2.5</td>
</tr>
<tr>
<td>Initial triggering of complement</td>
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<tr>
<td>Eukaryotic translation elongation</td>
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<tr>
<td>Eukaryotic translation initiation</td>
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<tr>
<td>Scavenging of heme from plasma</td>
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<tr>
<td>Interleukin-10 signaling</td>
<td>2.4</td>
</tr>
<tr>
<td>Class A1/2 rhodopsin-like receptors</td>
<td>2.7</td>
</tr>
<tr>
<td>HCMV early events</td>
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<tr>
<td>Signaling by GPCR</td>
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</tr>
<tr>
<td>GPCR ligand binding</td>
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</tr>
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</table>

IL16 OE

D) Top enriched GO Biological Processes following STARD5 overexpression

<table>
<thead>
<tr>
<th>GO Biological Processes</th>
<th>NES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cytoplasmic translation</td>
<td>5.0</td>
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<tr>
<td>Protein folding</td>
<td>2.5</td>
</tr>
<tr>
<td>Positive regulation of mRNA splicing</td>
<td>0.0</td>
</tr>
<tr>
<td>Post-translational protein targeting to ER membrane</td>
<td>-2.5</td>
</tr>
<tr>
<td>Oxidative phosphorylation</td>
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</tr>
<tr>
<td>Sterol biosynthetic process</td>
<td>-log(FDR)</td>
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<td>Steroid biosynthetic process</td>
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<tr>
<td>Sterol metabolic process</td>
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<td>Negative regulation of cell differentiation</td>
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<td>Alcohol biosynthetic process</td>
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STARD5 OE

E) Top enriched Reactome gene sets following STARD5 overexpression

<table>
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<th>Reactome Pathways</th>
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</thead>
<tbody>
<tr>
<td>Translation</td>
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</tr>
<tr>
<td>SRP-dependent cotranslational protein targeting to membrane</td>
<td>2.5</td>
</tr>
<tr>
<td>Regulation of expression of SLITs and ROBOs</td>
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<tr>
<td>Eukaryotic translation elongation</td>
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</tr>
<tr>
<td>Eukaryotic translation initiation</td>
<td>-5.0</td>
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<tr>
<td>Signaling by GPCR</td>
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<td>Cholesterol biosynthesis</td>
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<td>Regulation of cholesterol biosynthesis by SREBF (SREBP)</td>
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<td>GPCR ligand binding</td>
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STARD5 OE

F) IL16 and STARD5 associations with TE family gene sets

<table>
<thead>
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<tbody>
<tr>
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<tr>
<td>Alu subfamilies</td>
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<tr>
<td>ERV1 subfamilies</td>
<td>0.0</td>
</tr>
<tr>
<td>ERVL–MaLR subfamilies</td>
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</tr>
<tr>
<td>hAT–Charlie subfamilies</td>
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</tr>
<tr>
<td>ERVL subfamilies</td>
<td>-log(FDR)</td>
</tr>
<tr>
<td>TcMar–Tigger subfamilies</td>
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</tr>
<tr>
<td>hAT–Tip100 subfamilies</td>
<td>3.0</td>
</tr>
</tbody>
</table>

IL16 OE

<table>
<thead>
<tr>
<th>TE Family</th>
<th>NES</th>
</tr>
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<tbody>
<tr>
<td>L1 subfamilies</td>
<td>4.0</td>
</tr>
<tr>
<td>Alu subfamilies</td>
<td>2.0</td>
</tr>
<tr>
<td>ERV1 subfamilies</td>
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<tr>
<td>ERVL–MaLR subfamilies</td>
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</tr>
<tr>
<td>hAT–Charlie subfamilies</td>
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<tr>
<td>ERVL subfamilies</td>
<td>-log(FDR)</td>
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<td>TcMar–Tigger subfamilies</td>
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<td>hAT–Tip100 subfamilies</td>
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STARD5 OE

G) IL16 and STARD5 associations with an L1 family gene set

<table>
<thead>
<tr>
<th>L1 gene set enrichment</th>
<th>NES</th>
</tr>
</thead>
<tbody>
<tr>
<td>hAT–Charlie subfamilies</td>
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</tr>
<tr>
<td>ERVL subfamilies</td>
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Rank in Ordered Dataset

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Figure 5

**A** Scheme for assessing the effects of IL16 protein on L1 expression

Seed GM12878 LCLs

Treat cells with recombinant human IL16

Pellet cells and extract RNA

Sequence mRNA

Differential gene expression and GSEA

**B** Top shared GO Biological Processes

<table>
<thead>
<tr>
<th>Differentially regulated GO BPs</th>
<th>IL16 OE</th>
<th>rhIL16 T24</th>
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<tr>
<td>MAPK cascade</td>
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<tr>
<td>Leukocyte differentiation</td>
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<td>-4</td>
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<tr>
<td>Mononuclear cell differentiation</td>
<td>-4</td>
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<tr>
<td>Giogenesis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Negative regulation of cell differentiation</td>
<td></td>
<td></td>
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<tr>
<td>Regulation of hemopoesis</td>
<td>-log₁₀(FDR) 1.5</td>
<td>2.0</td>
</tr>
<tr>
<td>Positive regulation of cell death</td>
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<td>3.0</td>
</tr>
<tr>
<td>Regulation of leukocyte differentiation</td>
<td>3.0</td>
<td>3.5</td>
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<tr>
<td>Positive regulation of MAPK cascade</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive regulation of cell adhesion</td>
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</table>

**C** Top shared Reactome gene sets

<table>
<thead>
<tr>
<th>Differentially regulated Reactome pathways</th>
<th>IL16 OE</th>
<th>rhIL16 T24</th>
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</thead>
<tbody>
<tr>
<td>Regulation of cholesterol biosynthesis by SREBP (SREBF)</td>
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</tr>
<tr>
<td>Interleukin-10 signaling</td>
<td></td>
<td></td>
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<td>HCMV infection</td>
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<tr>
<td>HCMV early events</td>
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<tr>
<td>Activation of gene expression by SREBF (SREBP)</td>
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<td>Formation of tubulin folding intermediates by CCT/TriC</td>
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<td>L1cam interactions</td>
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<td>Interleukin-3, Interleukin-5, and GM-CSF signaling</td>
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<tr>
<td>Neuronal system</td>
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</table>

**D** rhIL16 associations with TE family gene sets

<table>
<thead>
<tr>
<th>Differentially regulated TE families</th>
<th>NES</th>
<th>rhIL16 T24</th>
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<tbody>
<tr>
<td>L1 subfamilies</td>
<td>5.0</td>
<td></td>
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<td>Gypsy subfamilies</td>
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<td></td>
</tr>
<tr>
<td>ERVL subfamilies</td>
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</tbody>
</table>

**Additional Information:**

- The figure includes a scheme for assessing the effects of IL16 protein on L1 expression, with steps including treatment with recombinant human IL16, cell pelletation, RNA extraction, and differential gene expression analysis.
- **B** highlights the top shared Gene Ontology (GO) Biological Processes, with differentially regulated processes such as MAPK cascade, leukocyte differentiation, and mononuclear cell differentiation.
- **C** shows the top shared Reactome gene sets, including regulated pathways like cholesterol biosynthesis, interleukin-10 signaling, and HCMV infection.
- **D** details differentially regulated TE families, with L1 subfamilies showing a NES of 5.0.

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Figure 6

A Scheme for assessing shared pathway changes in conditions with L1 regulation

B Shared upregulated TE family gene sets

C Top shared Hallmark pathways

D Top shared GO Biological Processes

E Top shared Reactome pathways
Figure 7

A Scheme for identifying TE eQTL co-associated aging traits

1. Generate a list of TE trans-eQTLs
2. Identify co-associated phenotypes catalogued on Open Targets Genetics
3. Group phenotypes by age-related MeSH trait IDs

B Number of SNVs with an age-related MeSH trait

- No 55.5%
- Yes 44.5%

C Number of associated age-related traits per SNV

D Number of linked age-related traits per SNV

E Top traits with the most number of associated SNVs

- Hyperparathyroidism
- Thyroid Diseases
- Hypothyroidism
- Hemoglobinopathies
- Coronary Artery Disease
- Diabetes Mellitus, Type 2
- Diabetes Mellitus
- Body mass index
- Arthritis, Rheumatoid
- Psoriasis

F Mouse serum [IL16] with age

- Young (4 - 6 months)
- Old (20 - 24 months)

Meta-Analysis p = 0.0479
Meta-Analysis p = 0.0402