SHORT COMMUNICATION

Comprehensive translational control of tyrosine kinase expression by upstream open reading frames

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Post-transcriptional control has emerged as a major regulatory event in gene expression and often occurs at the level of translation initiation. Although overexpression or constitutive activation of tyrosine kinases (TKs) through gene amplification, translocation or mutation are well-characterized oncogenic events, current knowledge about translational mechanisms of TK activation is scarce. Here, we report the presence of translational cis-regulatory upstream open reading frames (uORFs) in the majority of transcript leader sequences of human TK mRNAs. Genetic ablation of uORF initiation codons in TK transcripts resulted in enhanced translation of the associated downstream main protein-coding sequences (CDSs) in all cases studied. Similarly, experimental removal of uORF start codons in additional non-TK proto-oncogenes, and naturally occurring loss-of-uORF alleles of the c-met proto-oncogene (MET) and the kinase insert domain receptor (KDR), was associated with increased CDS translation. Based on genome-wide sequence analyses we identified polymorphisms in 15.9% of all human genes affecting uORF initiation codons, associated Kozak consensus sequences or uORF-related termination codons. Together, these data suggest a comprehensive role of uORF-mediated translational control and delineate how aberrant induction of proto-oncogenes through loss-of-function mutations at uORF initiation codons may be involved in the etiology of cancer. We provide a detailed map of uORFs across the human genome to stimulate further research on the pathogenic role of uORFs.

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INTRODUCTION

Systems biology data have demonstrated the importance of many layers of protein expression control beyond transcriptional gene regulation, including regulation at the translational and post-translational levels. Specifically, ribosome profiling has uncovered abundant non-canonical translational initiation at upstream open reading frames (uORFs) located in the transcript leader sequence (TLS) of eukaryotic mRNA. A potential uORF is defined by a translational initiation site that precedes the initiation codon of the main protein-coding sequence (CDS) and by a subsequent in-frame termination codon within a given transcript. Translation of a uORF is considered to affect the translation rates of a subsequent CDS by interfering with unrestricted progression of the scanning ribosome or by consuming a functional preinitiation complex. In uORF-containing transcripts, leaky ribosomal scanning across the uORF initiation codon or reinitiation of ribosomes after termination at the uORF stop codon, permit translation of the CDS. Multiple modes of translational control through uORFs have been reported for ~100 eukaryotic genes, yet systematic analyses on the relevance of uORF-regulation in specific groups of functionally important genes are lacking. Although there is evidence that uORF-activating or -creating point mutations in the tumor-suppressor genes CDKN1B and CDKN2A are involved in the development of multiple endocrine neoplasia syndrome type 4 and hereditary melanoma, respectively, few data exist on a potential role of loss-of-uORF mutations in the development of human disease.

Tyrosine kinases (TKs) represent a prototype family of oncogenic proteins that are key enzymatic regulators of cellular signaling cascades involved in proliferation and differentiation control. TKs are frequently overexpressed or constitutively activated in malignant cells and oncogenic functions of TKs have been attributed to (I) genetic translocations (for example, BCR/ABL in chronic myeloid leukemia), (II) gene amplifications (for example, ERBB2 in breast cancer) or (III) mutations within their kinase domains causing enhanced or constitutive signaling activity (for example, c-Kit in acute myeloid leukemia). Here, we investigated the prevalence and regulatory impact of uORF-mediated translational control in TKs and other proto-oncogenes by sequence analysis and by targeted mutational ablation of uORF initiation and termination codons. Our data suggest frequent uORF-mediated translational repression of human TKs and imply a potential driver function for loss-of-uORF mutations in tumorigenesis.

RESULTS AND DISCUSSION

Characterization of prevalence and properties of TK uORFs The set of TKs analyzed in this study includes all human proteins with assigned ‘tyrosine kinase activity’ according to the amigo.geneontology.org website (GO:0004713) accessed in April 2012.

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A total of 140 individual proteins were identified that were encoded by 296 transcript variants annotated in the RefSeq (hg19) database. Within this set, a total of 409 distinct AUG-initiated uORFs were detected. For simplicity, and because translational initiation at uAUG codons appeared to be most relevant in downstream translational control, the analysis of near cognate non-AUG upstream initiation codons was excluded from this study, despite a potential role in serving as alternative uORF start sites.

Of the 140 human TK genes, 89 (63.6%) encoded at least one transcript variant that contained one or more uORF(s) (Figure 1a and Supplementary Table 1). The observed proportion of uORF-bearing genes in the TK subset was higher as compared with the average uORF prevalence of 55.5% in the whole genome (Supplementary Table 2). Most likely, longer TLSs of TK transcripts accounted for the higher proportion of uORF-bearing genes in the TK subset, as the ratio of uORF initiation codons per base of TLS was not increased above genome-wide frequencies (data not shown).

Within the 296 individual TK transcripts, one, two or more uORFs were found in 20.6%, 18.9% and 27.0%, respectively, showing that the occurrence of multiple uORFs within the same transcript is a rather frequent event (Figure 1a). In all, 18.6% of the uORFs overlapped with the CDS initiation codon (Supplementary Table 1), meaning that translation of these uORFs would prevent ribosomal reinitiation at the downstream CDS. By mapping uORF genomic positions to the '100 vertebrates conserved elements' provided by the UCSC, we found that uAUGs were significantly higher conserved as compared with the remaining uORF or TLS sequences in both, the TK subset and genome wide (Figure 1b).

Looking at all human genes, also uORF sequences excluding the uAUGs were significantly higher conserved as compared with

![Figure 1](image-url). Presence, conservation and position of uORFs in human tyrosine kinases and all human genes. (a) Pie charts indicating the prevalence of uORFs in TK genes and the number of uORFs per transcript for alternative TK mRNAs. (b) Bar graph displaying the frequency of conserved bases within TLSs, uORFs and uAUGs of TKs and all human genes. Note that columns labeled 'TLS' do not include uORF bases and columns labeled 'uORF' do not include uAUGs, respectively. For genome-wide transcript analyses, Hg19 annotations were downloaded from the UCSC genome browser database. Of 44 113 entries, we excluded 7794 non-coding RNAs, 1861 sequences not aligning to autosomes or sexual chromosomes, and 1237 transcripts lacking a TLS. Finally, we positively selected the 32 332 mRNAs with unique genome alignments corresponding to 17 893 distinct genes (ftp://ftp.ncbi.nlm.nih.gov/refseq/H_sapiens/mRNA_Prot/). Conservation was defined by mapping the respective genomic positions to the '100 vertebrates conserved elements' provided by UCSC (http://genome.ucsc.edu/cgi-bin/hgTrackUi?g=cons100way). Statistical significance was analyzed using the $\chi^2$ test and indicated as **P < 0.01 and NS (not significant). (c) Graph indicating the prevalence of uORFs in respect to the length of the TLS for TKs and all human genes (solid green and black lines, respectively). Dashed lines indicate expected uORF frequencies for TKs and all human genes following a 10 000-times randomization of TLS nucleotides (green and black, respectively). (d) Box plots summarizing the variable length of TLSs and uORFs in TK transcripts. The graph indicates the range of nucleotide distances from the transcriptional start site (TSS) to the uAUG and from the uAUG to the main AUG (mAUG).
non-uORF TLS sequences; yet this observation did not hold significance for the smaller gene set of TKs. Extending a previous report that identified AUG as the triplet most frequently conserved among human and mouse,16 our data based on conserved elements among 100 vertebrate species suggest functional importance of uAUGs and occasionally also of the subsequent uORF sequences.

In all human transcripts and in the TK subset, uORFs were less prevalent than expected, as shown by quantification of uAUG frequencies in natural and randomized TLSs (Figure 1c). These observations are in line with previous findings implying negative selection of uAUGs.17 Moreover, we observed a high variability of uORF length and position within all transcripts (data not shown) and individual TK-TLSs (Figure 1d and Supplementary Table 1), suggesting that the regulatory potential of a uORF may be highly dependent on the structural vicinity within individual transcripts.

Translational repression of TKs by uORFs
Having characterized the high prevalence of uORFs in TK transcripts, we set out to monitor a potential cis-regulatory function on downstream translation. A genetically modified luciferase reporter plasmid was generated to facilitate the insertion of complete TLSs, including the gene-specific CDS initiation codon and the core-Kozak nucleotide at position +4 of the main AUG, to retain the endogenous initiation context of individual transcripts (Figure 2a and Supplementary Figure 1). The inserted TLSs either contained the wild-type uORF initiation codon (wt uORF) or a point mutation at the uORF start site (UUU), known to abolish ribosomal initiation (ΔuORF).18 The functionality of the TLS luciferase reporter system was tested by inserting wt or ΔuORF versions of the TLSs of CECA, CEBPB and ERBB2 (Figures 2b–d, positive controls), representing paradigm transcripts with conserved uORFs and previously documented regulatory potential.15,20,22 As expected, elevated luciferase activity was detected for the ΔuORF-containing reporter constructs, as compared with the wt uORF controls, whereas mRNA abundance was not significantly affected. Hence, the TLS reporter system proved to be suited to determine the regulatory impact of uORFs on downstream translation.

Following the genome-wide identification of potential uORFs in human TK transcripts, 10 TLSs harboring a single uORF were selected to exclude competing impacts of subsequent uORFs within the same TLS during functional analyses. Selection was further based on uORF properties that implied functional importance,3 including the conservation among vertebrate species and the quality of the Kozak initiation context. Transcripts of HCK, MAP2K3, ERBB3 (short and long transcript variants), LCK, MAP2K2, TYRO3 and RET displayed considerably high conservation of the uORF initiation codon, whereas in the transcripts of YES1 and ZAP70 interspecies conservation was lower (Figure 2b).

Introduction of wt and ΔuORF variants of the selected TK TLSs into the luciferase reporter construct revealed that deletion of the uORF initiation codon enhanced downstream translation in all cases (Figure 2c). Luciferase activity increased >3-fold after removal of conserved uORFs in HCK, MAP2K3, ERBB3-short and LCK transcripts. Here, the ΔuORF-induced de-repression of translation was similar as observed with the positive controls CECA, CEBPB and ERBB2. Of note, deletion of the less conserved uORFs (ZAP70 and YES1) resulted in lower, yet still significant de-repression of reporter activity, implying translational activity also at less conserved uORFs. Quantification of mRNA excluded the possibility that an increase in mRNA abundance predominately accounted for the higher luciferase activity observed (Figure 2d), although mildly elevated transcript levels of ΔuORF TLS versions were seen on some occasions. Similar results of consistently enhanced luciferase translation in response to functional ablation of the TK uORFs were obtained by using HEK293 instead of HeLa cells (Supplementary Figure 2), suggesting cell type independence.

Next, the analysis was extended to additional non-TK proto-oncogenic transcripts, known to be overexpressed or amplified in human cancer (Figures 2b–d).21 Two independent ribosomal profiling studies in human and mouse had demonstrated translational activity at specific uORFs of SKP2, MDM2, CDK4 and SHC1.2,14 Similar to the observations for TKs, deletion of uORF initiation codons of aforementioned proto-oncogenes resulted in increased downstream translation of their CDS, whereas occasional stabilizing effects on the related mRNAs were much smaller. Taken together, the results imply that translation of many TKs and other proto-oncogenic proteins is constitutively repressed by uORFs. Beyond previous observations in individual transcripts,20,22 the universal regulatory effect of uORF deletions in all cases analyzed here support the existence of a widespread translational control mechanism, where proto-oncogenes may gain enhanced expression and tumorigenic potential through loss-of-uORF mutations within their TLSs.

Mutation-induced elongation of uORFs sustains translational repression
Within the sample set of TKs and proto-oncogenes described above, no obvious difference in the regulatory impact of overlapping vs non-overlapping uORFs could be observed. This is in line with a previous report, where naturally occurring overlapping- and non-overlapping uORFs, on average, showed similar inhibitory effects on downstream translation.3

To investigate the consequences of de-novo mutations that may convert a non-overlapping into an overlapping uORF, stop codons of the uORFs (uStop) in CECA, CEBPB, ERBB2, HCK, MAP2K3 and RET were replaced by non-stop CUU codons (ΔuStop) that created overlapping uORFs in luciferase constructs and prevented the possibility of ribosomal reinitiation (Figure 3a). In all transcripts analyzed, luciferase translation efficacy was markedly reduced in response to the loss-of-uStop mutations (Figure 3b), suggesting that in the wt TLSs reinitiating ribosomes contribute substantially to the translation of the CDS. Consistently, re-introduction of alternative uStop codons downstream of the natural uStop and upstream of the CDS in HCK, MAP2K3 and RET reverted this inhibitory effect to various extents (Figures 3a and b). Of note, for most of the ΔuStop versions of the TLS, the luciferase mRNA levels mildly increased as compared with the wt TLS (Figure 3c). As previously reported, reduced nonsense-mediated mRNA decay in the absence of uORF termination codons may account for the observed elevation of respective mRNAs levels.23

The observation of reduced CDS translation after deletion of uStop codons in all four TK transcripts suggested that the translation of these and other uORF-bearing TKs may substantially depend on reinitiating ribosomes that previously translated a uORF and terminated at a uStop codon. As translational reinitiation is dependent on the reconstitution of a functional reinitiation complex,24 TK translation may be highly sensitive to environmental signals and global translational conditions of a cell.

Widespread prevalence of natural polymorphic uORFs
To investigate whether naturally occurring sequence variations in TK uORF initiation codons or the surrounding Kozak consensus sequences may alter translational control, we screened TK TLSs for single-nucleotide polymorphisms (SNPs) using the UCSC genome browser15 and dbSNP.25 In the TK gene set, we found eight SNPs and one deletion depleting a uAAUG (ΔpuORF). Accordingly, ~5.8% of TKs may be subject to translational variability due to polymorphic uORF initiation codons (puORF; Figure 4a and Supplementary Table 5). In addition, 26 SNPs and 1 deletion in
14.3% of TK genes were found to affect uORF-related Kozak consensus sequences (pKozak).

For experimental exploration of these SNPs, we focused on KDR, where the uORF-deleting SNP (rs7667298) affected one single uORF, and on MET, where the second of two uORF initiation codons was altered by two independent SNPs (rs13235174 and rs13222425, Figure 4b). Allele frequencies annotated in the dbSNP were 44.5% vs 55.5% for the puORF vs ΔpuORF allele of KDR and unknown for the MET SNPs. The ΔpuORF allele of KDR was associated with mildly enhanced translation of the luciferase reporter gene (Figure 4c). In the MET transcript, the two alternative polymorphic ablations of the second MET uORF (UUG and AAG), or a combination of both (UAG), resulted in mild de-repression of downstream translation. Additional deletion of the first MET uORF, located upstream and in-frame to the polymorphic uAUG, resulted in further enhancement of reporter expression, but did not alter the regulatory potential of the ΔpuORF variants observed at the second MET uORF (Supplementary Figure 3), suggesting more complex functions of translation initiation control. As we observed slightly higher mRNA levels for some of the polymorphic ΔpuORF
versions of the KDR and MET TLSs, we cannot exclude some contribution of mRNA stabilization to the increased reporter function (Figure 4d and Supplementary Figure 3). Nevertheless, active translation of the KDR and MET uORFs could be confirmed independently by the detection of C-terminally HA-tagged uORF peptides in immunoblot analyses (Figure 4e).

Irrespective of the mode of translational induction, our data suggest that naturally occurring uORF polymorphisms may alter the expression of the respective downstream proteins. In support of the hypothesis of loss-of-uORF-mediated proto-oncogene activation, the KDRΔuORF allele was independently found to be associated with increased KDR protein levels in lung cancer samples26 and, together with two additional KDR SNPs, with increased risk of glioma development.27 Furthermore, the ApuORF allele alone was associated with an acute course of sarcoidosis28 and a trend toward shorter overall survival of patients suffering from pancreatic carcinoma.29 For the MET SNP, no clinical association data are available to date.

Given the highly reproducible translational activity of uORFs in TKs and other proto-oncogenes described above, we determined the genome-wide prevalence of SNPs within uORF start codons and the surrounding Kozak consensus sequences. First, computational sequence analyses generated a current map of all human uORFs and provided information on the transcripts affected, the genomic position and the length of each individual uORF (Supplementary Table 2). Second, we analyzed how many of the 56,248,699 human SNPs listed in dbSNP 137 mapped to specific uORF sequences. We identified 1375 SNPs affecting uAUGs and 2724 SNPs affecting uAUG-related Kozak sequences in 2610 individual genes (Supplementary Table 6). These observations reveal that the translation rates of up to 14.6% of annotated human genes may be dependent on SNPs affecting the uORF initiation context (Figure 4a). In addition, we detected 697 SNPs at uStop codons in 3.4% of the genes (Supplementary Table 6), further increasing the number of proteins whose expression may be subject to inter-individual variability in response to uORF-related SNPs. For eight of the uORF-related SNPs described here, clinical association data have been documented according to the current dbSNP annotations in the UCSC database (Supplementary Table 7). The low number may in part reflect the fact that...
genome-wide sequence data from clinical samples largely focused on coding exons and excluded uORFs in the TLSs. In addition, dbSNP annotations may be incomplete, as recent publications reported higher numbers of genotype–phenotype associations and potential clinical associations of altered uORF-mediated translational control in various types of diseases.3,10

The data described above demonstrate a wide range of regulatory potential of uORF-mutations at initiation and termination codons. Our results demand for systematic searches for loss-of-uORF mutations in proto-oncogenes, and gain-of-uORF or loss-of-uStop mutations in tumor-suppressor genes, as both types of translational deregulation may cause misexpression of the respective proteins and may result in dominant phenotypes.3,10 Ultimately, ongoing re-sequencing of whole genomes, together with resources implemented to survey functional data from ribosome profiling51 and individual studies,4 will help to precisely characterize the contribution of uORF-related genetic variants to the etiology of disease and, more general, to phenotypic divergence. Our data suggest that the translational activation of proto-oncogenic proteins through loss-of-uORF mutations may contribute to increased tumor susceptibility.

Figure 4. Polymorphic uORFs affect TK translation. (a) Pie charts indicating the fraction of TK genes and all human genes with SNPs in uORF start codons (puORF) and/or corresponding Kozak sequence contexts (pKozak). SNPs were analyzed for all RefSeq-annotated transcript variants of TKs and the whole human genome using dbSNP 137.25 (b) Schematic representation of the position and length of a uORF and puORF within the TLSs of KDR and MET. SNPs that disrupt the uORF initiation codon are underlined and the resulting alternative codons are displayed in gray. Conservation of the uAUG among human and mouse (check mark) and among a total of nine vertebrate species is depicted. The weak quality of the Kozak context (−, no core Kozak base matches) is indicated. (c) Bar graph representing the relative luciferase activity in the presence of the wt puORF and the ΔpuORF containing TLSs of indicated TKs. For the two MET SNPs, all three possible SNP combinations were analyzed. (d) Bar graph indicating the relative luciferase mRNA levels of wt puORF and ΔpuORF reporter constructs for indicated TLSs. Error bars represent the s.e.m. of at least three independent experiments. Asterisks indicate statistical significance (**P < 0.01 and *P < 0.05). (e) Immunoblot shows HA-tagged peptides translated from KDR and MET uORFs. The TLSs of KDR and MET including the cap-to-uORF sequence, the uORF-coding sequences (with a disrupted uStop codon) and a C-terminally added triple HA-tag (Supplementary Table 4) were cloned into the pcDNA3 vector (Invitrogen). HEK293 cells were transfected following standard protocols and protein lysates were separated 42 h later on 18% SDS–polyacrylamide gels. After transfer, polyvinylidene fluoride membranes (Roth) were probed with specific antibodies: HA (HA.11, Covance) and β–Actin (Clone AC-15, Sigma-Aldrich) and horseradish peroxidase-linked secondary antibody (GE Healthcare Life Sciences).
CONFLICT OF INTEREST
The authors declare no conflict of interest.

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