ORFLine: a bioinformatic pipeline to prioritise small open reading frames identifies candidate secreted small proteins from lymphocytes

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Abstract
The annotation of small open reading frames (smORFs) of less than 100 codons (<300 nucleotides) is challenging due to the large number of such sequences in the genome. The recent development of next generation sequence and ribosome profiling enables identification of actively translated smORFs. In this study, we developed a computational pipeline, which we have named ORFLine, that stringently identifies smORFs and classifies them according to their position within transcripts. We identified a total of 5744 unique smORFs in datasets from mouse B and T lymphocytes and systematically characterized them using ORFLine. We further searched smORFs for the presence of a signal peptide, which predicted known secreted chemokines as well as novel micropeptides. Five novel micropeptides show evidence of secretion and are therefore candidate mediators of immunoregulatory functions.

Keywords: Lymphocytes, micropeptides, small open reading frames, Riboseq

Introduction
Open reading frames (ORFs) are the regions of the genome which contain the triplet nucleotide codons that direct the sequence of amino acids (AA) in a protein. ORFs of less than 100 codons, referred to here as small ORFs (smORFs), are particularly numerous and have been challenging to annotate and to functionally characterise (reviewed in (1-3)). smORFs have been classified according to their location relative to the main ORF within the host transcript (4). The translation products of smORFs, termed micropeptides, have been shown to be involved in many aspects of life (5-12). These new discoveries complement already characterised peptides and small proteins known to be important biological regulators. Within the immune system the best characterised of these include host defence anti-microbial peptides, chemokines and cytokines that are known to play essential roles in normal and pathological immune reactions.

The advent of next-generation sequencing technologies and proteomic approaches has led to a more comprehensive annotation of genes, transcripts and their translated protein products (3). Several large-scale genomic studies have revealed that a much larger fraction of the genome is transcribed and translated than was anticipated (13,14). A large collection of putative translatable smORFs have been identified by computational methods based on the level of DNA and protein sequence conservation across species, coding potential and context of the initiation codon. Ribosome profiling (Ribo-Seq), an approach based on deep sequencing of isolated ribosome-protected mRNA fragments, has provided extensive evidence for the translation of smORFs (14-21). A variety of metrics and algorithms can use Ribo-Seq data to annotate translated regions of the genome. Amongst them ORFscore is a metric to quantify the bias of the trinucleotide periodicity pattern of ribosome protected footprints (RPFs) towards the first reading frame in an ORF (18). The periodicity pattern has been used by several algorithms and pipelines including ORF-RATER (19), RibORF (20), Ribotaper (22), RP-BP (23), and Ribocode (24). A recently described integrated platform called RiboToolkit provides a one-step server for comprehensive analysis of Ribo-seq data and utilises Ribocode as part of its packages (25). In addition to ORFscore, other metrics can be used in conjunction to improve actively translated ORF identification. For example, the Ribosome Release Score (RRS) detects the termination of translation at the stop codon and can robustly distinguish protein-coding transcripts from ncRNAs (26).
Here we describe a new analytical pipeline that we call “ORFLine” that performs a comprehensive and systematic analysis of RNA-Seq and Ribosome profiling to identify actively translated smORFs. In comparison to previously published pipelines, ORFLine reduces computational demands by focusing on smORFs. Also, ORFLine applies a series of logical filters to improve the stringency of prediction. Predicted smORFs are classified according to their host transcript type and the position of the smORF relative to other ORFs within the same transcript. We have applied ORFLine to datasets of mouse lymphocytes and discovered 5744 actively translated smORFs during B and T cell activation. We also analysed the genetic conservation, translation efficiency, and related biological processes of the predicted smORFs. We further identified smORFs containing signal peptides which have a potential to be secreted and could act as immune regulators.

Results

Overview of ORFLine

ORFLine takes Ribo-Seq, RNA-Seq, reference genome, transcriptome and gene annotation as input data and produces an output list of predicted smORFs with genomic coordinates and classification (Figure 1A). The three main pipeline components to process the raw Illumina sequences and perform smORF prediction are: 1) Prediction of putative smORFs; 2) Sequencing data QC and processing; and 3) Identification of translated smORFs. Prediction of putative smORFs and sequencing data processing are independent components and can be executed in parallel. The identification of translated smORFs utilises the output of the previous two components as input (Table 1). ORFLine is applicable to data from any species.

The output of ORFLine is a list of smORFs that have passed the filters in the identification of translated smORFs. They are identified as smORFs with ribosome protected RNA fragments (RPF) signal. The output file (Table 2) contains the genomic location and splicing information (including number of exons and exon lengths) of a smORF are clearly annotated and can be loaded and visualized in a genome browser. The quantitative information about a smORF is also calculated including translation efficiency, RNA expression and Ribosome-protected RNA expression (FPKM value). The nucleotide sequences are retrieved and translated into amino acid sequences and presented in column 25 of Table 2.
Comparison between ORFLine and RiboCode

We characterised smORFs in mouse lymphocytes using datasets obtained from ex vivo non-activated B cells; a published dataset from our lab of Lipopolysaccharide (LPS)-activated B cells (28); a new dataset of B cells activated with LPS plus interleukins IL-4 and IL-5 for 48 hours, by which time most cells had divided once, and some started a second division; naïve CD4+ T cells stimulated with antibodies to CD3 and CD28 which mimics activation by antigen; and a published time-course of Th1 T cells re-stimulated with anti-CD3+anti-CD28 (29) see Table S1. ORFLine predicted a total of 5744 unique smORFs in all samples analysed (union of 2607 smORFs predicted in B cells and 4935 smORFs predicted in T cells) (Table S3). A lower number (568) of smORFs were predicted for the resting B cells than for LPS-activated B cells (2444), most likely reflecting an overall increase in RNA abundance associated with elevated rates of transcription in activated B cells.

We also analysed the same datasets with the recently published ORF-detection pipeline RiboCode (24) using its default settings. RiboCode predicted a total of 15,920 unique smORFs, of which 3,667 are smORFs nested in longer smORFs in the same reading frame and 48 smORFs are from non-expressed transcripts. We removed those 3,715. In the remaining 12,205 smORFs, 3,337 were predicted as internal or frameshift smORFs. These are found nested in the CDS, but in a different reading frame. Considering that frameshift translation is a rare event (30), they are not included in our results. We removed all 3,337 frameshift smORFs predicted by RiboCode and compared the remaining 8,868 non-internal smORFs predicted by RiboCode with the 5,744 predicted from ORFLine (Figure S2). Of these, 1,957 (22.1% in RiboCode and 34.1% in ORFLine) are found as exact genomic coordinate matches by both pipelines (Table 3). For un-annotated smORFs, we are not certain they are translated, and we lack a reference set of true-positives, therefore we sampled the smORFs which are differentially identified by the two pipelines and noticed that smORFs predicted by RiboCode typically have low RPF coverage or are assigned a low or negative ORFScore, or low RRS, and are filtered out by ORFLine (Figure S3). Our criteria for metrics have shown to be robust in smORF prediction in previous studies (18,26). ORFLine also predicted 356 smORFs encoded by low abundance transcripts (25% percentile) that are not predicted by RiboCode. Taken together, these comparisons demonstrate that ORFLine allows a reliable identification of smORFs residing in transcripts of low abundance.
smORF classification

ORFLine classified smORFs according to their relative position with the annotation, if any, of the host transcript (Figure 1B). It predicted canonical smORFs and extended variants of annotated coding DNA sequences (CDS) of 100 codons or less in protein-coding mRNAs. ORFLine also found upstream ORFs (uORFs) which we subdivided into either uORFs starting in the 5' untranslated region (5' UTR) of annotated protein-coding mRNAs and overlapping with the coding region (ouORFs) or non-overlapping uORFs (nuORFs) which terminated before the start of the annotated CDS. uORFs are known to be prevalent in the genome and they represent ~80% of all smORFs predicted by ORFLine (Figure 1C). In addition, ORFLine identified downstream ORFs (dORFs) as the rarest class of smORFs. These can be subdivided as overlapping dORFs (odORFs) that overlap with the CDS and extend into the 3' UTR of known protein-coding mRNAs or non-overlapping dORFs (ndORFs) located exclusively in annotated 3'UTRs. Lastly, 501 smORFs in putative non-coding RNAs (long non-coding RNAs and pseudogenes) were predicted, which are termed ncORF. Direct biochemical and functional evidence is available for only ~40% (338) of canonical smORFs in protein databases such as UniProt (The UniProt Consortium, 2019) for their protein products, it includes diverse entities such as chemokines and subunits of mitochondrial complexes. The remainder (~5340) have either not been functionally characterised or have not been annotated at all.

smORF conservation

To examine the conservation of smORF-encoded micropeptides between species, we employed PhyloCSF to analyse signatures of evolutionary conservation. 11.4% of smORFs had a PhyloCSF score > 50, thus showing strong evidence of conservation (Figure 2A), with canonical smORFs being enriched among these (Figure 2B). A small subset (~6.5%) of uORFs, ncORFs and dORFs showed high PhyloCSF scores, indicating conservation of smORFs CDS. There are over 60% of smORFs lacking signs of selective pressure to maintain their amino acid sequences (no cross-species sequence alignment and not conserved, Figure 2A), in which uORFs, ncORFs and dORFs are enriched (Figure 2B). The median length of canonical smORFs is 79 codons, however, the median length of uORF, dORF and ncORF are 24, 34 and 33 codons respectively. By comparison with other classes, canonical smORFs are, on average, longer and more highly conserved (Figure 2C). Having distinct transcript organization, size, conservation and peptide structure, the cellular and molecular functions of canonical smORFs, uORFs, dORFs and ncORFs are likely to differ from each other, with less conserved classes primarily independent of peptide sequences. However, we observed that the PhyloCSF score positively correlates with the length of coding sequence (data not shown). Therefore, it is likely that the conservation of shorter smORFs is underestimated.

Canonical smORFs

A total of 338 canonical smORFs were predicted in B and T cells. 88% of these are conserved or weakly conserved between species (Figure 3A). We divided canonical smORFs into “short CDS” and “short isoforms”, the latter are the products of alternative splicing of transcripts from genes annotated as encoding proteins longer than 100 amino acids (4). Among the predicted canonical smORFs, 54.4% are short CDSs and 45.6% are short isoforms. There are hundreds of putative short CDSs in mouse and human, these are typically located on monocistronic transcripts and their host transcripts are structurally shorter and simpler compared with canonical mRNAs (4). We have predicted 184 short CDSs and they have a median size of 79 codons. We find short isoforms have a median size of 80 codons and resemble short CDSs in size and conservation (Figure 3B). As short isoforms share conserved protein sequences with their longer canonical protein isoforms, they may have functions that are directly related to their longer protein isoforms (4). To increase confidence that predicted canonical smORFs were indeed translated we calculated the translation efficiency of the short CDSs and short isoforms. When compared to long CDSs of expressed protein-coding transcripts, we found their median translation efficiency to be greater (Figure 3C). We also conducted GO term enrichment analysis comparing the 184 short CDS and the 154 short isoforms against the remaining 3536 smORF-encoding genes of B and T cells. The top hits of short CDSs are related to chemokine activity and mitochondrial biology (Figure 3D, Table S4). Seven chemokines are predicted (Ccl1, Ccl22, Cc13, Cc14, Cc15, Cc11). We also observed enrichment of gene products involved in mitochondrial complexes, for example, Uqcr10 is a subunit of Coenzyme Q:cytochrome c reductase (Complex III); this complex has a critical role in the oxidative phosphorylation pathway for the generation of ATP. Another mitochondrial protein is Romo1, which is located in the mitochondrial membrane and is responsible for increasing the level of reactive oxygen species (ROS) in cells (31). Romo1 also has antimicrobial activity against a variety of bacteria by penetrating the bacterial membrane (32). Short isoform encoding genes are associated with a broad range of GO terms, with no single term for GO biological processes reported as enriched.
We then investigated the influence of uORFs on translation of the downstream CDS. Several studies have shown a repressive effect of uORFs on mRNA translation of CDS during the first four hours of T cell activation (Figure 4A). About 4% of the uORFs have a high PhyloCSF score and TE (above the median TE of long CDS) and potentially encode conserved functional micropeptides (Figure 4B, for B cells activated with LPS and IL-4+IL-15). However, the sequences of the majority of uORFs are not conserved, suggesting that any potential function is largely independent of the encoded peptide. The proportion of expressed uORF-containing transcripts in B cells and T cells is between 6.2% and 12.4%, except resting B cells (2.7%). It has been demonstrated that uORFs may regulate the translation of the downstream CDS. Several studies have shown a repressive effect of uORFs on the translation of CDS (21,36,37). We analysed the effect of uORFs on mRNA translation of their associated genes by comparing the translation efficiency of the CDS in all uORF-containing transcripts versus those lacking uORFs. As expected, the presence of uORFs and overlapping uORFs was associated with a translation repression (Figure 4C). We performed GO enrichment analysis for all uORF-containing genes to discover their associated biological processes (2881 target genes against 3536 background genes) and these genes are mostly enriched in processes linked to protein modification, regulation of gene expression and cellular response to stimulus (Figure 4D, Table S4). This indicates that uORF-containing genes are broadly involved in complex biological pathways such as protein or RNA production and cell signalling. Regulatory uORFs may be suited to allow the rapid changes in gene expression in response to stress and environmental stimuli.

We then investigated the influence of uORFs on translation of the downstream CDS during the first four hours of T cell activation (Figure 4E). During the first two hours of activation the median RNA and RPF abundance for both non-uORF-containing transcripts and for transcripts containing a single uORF remains mostly unchanged as the median Log2 fold change is close to 0. There were few transcripts with two or more uORFs and these had a median 0.35 Log2 fold change of RNA and RPF abundance that is not statistically significantly different from the other two groups. However, we cannot rule out that we would find a difference if the numbers of transcripts in this class was greater. By four hours all mRNAs, irrespective of the presence of uORFs, show a median Log2 fold change of RNA abundance of around 0.5 compared to 0 hours. At 4h RPFs did not increase substantially, indicating a possible delay of translation of induced transcripts. As transcripts with two or more uORFs show a negative and the lowest median Log2 fold change of RPF abundance at 4h compared to 0h (P=0.04256 when compared to non-uORF-containing transcripts) the presence of uORFs appears to have a negative impact on translation of the downstream CDS. For translation efficiency, there is little change during the first two hours. The negative median Log2 fold change in TE for all groups in 4h vs 0h reflects the increase of RNA abundance at 4h and the lack of change of transcripts in RPFs. These results indicate that the rate of translation lags behind the increase in transcript abundance and the presence of uORFs can affect translation in activated T cells.

### smORFs in non-coding RNAs

Non-coding ORFs (ncORFs) are smORFs that are found in annotated long non-coding RNAs (lncRNAs) and pseudogenes. They are typically short with a median length of 33 codons. By definition, non-coding RNAs are not translated into protein. However, annotated lncRNAs have been predicted from their sequences to contain six smORFs on average (4). We have predicted 501 translated ncORFs and 14.4% of these are considered conserved or weakly conserved. We noticed very different distributions of size and PhyloCSF score between ncORFs and canonical smORFs (Figure 5A). The distribution of translation efficiency for ncORFs is also different from that for long CDS, the median TE of ncORFs is greater than long CDS (Figure 5B). Three ncORFs identified in LPS-activated B cells (Cct6a, Gm16675 and EDL19200.1) were found to have a high PhyloCSF score and TE, so we infer them to encode functional micropeptides (Figure 5C). We searched the micropeptides they encode in NCBI BLASTp database (38), but did not find any match for Gm16675. The 6330418K02Rik gene is annotated as an antisense lncRNA gene in GENCODE and only one match was reported for its predicted micropeptide (sequence ID: EDL19200.1). The micropeptide was partially aligned to three uncharacterized proteins with 35.59% to 78.18% identity in Habropoda.
containing transcripts may be translatonally enhanced. Cumulative distribution of translation efficiency in expressed dORF-containing transcripts versus transcripts lacking dORFs as control. Significance was computed using a two-sample Kolmogorov–Smirnov test, P = 0.437. *Laboriosa* and *Gulo gulo*. The examples likely reflect that these genes are misclassified as non-coding, although it is possible that they could also have functions as a noncoding RNA in addition to their peptide coding capacity.

**dORFs**

243 ndORFs and 17 odORFs were predicted with a median length of 34 AA. Only 20 (~7.7%) are conserved or weakly conserved (Table S5). The translation efficiency of dORFs is lower than the long CDSs in general (data not shown). In transcripts that contain multiple ORFs, a translation reinitiation mechanism is able to prevent recycling of some or all ribosome subunits upon termination of the first translated ORF and thereby enable the translation of the dORF (39). The low TE indicates a very low level of translational reinitiation after the stop codon of the upstream CDS. In B cells activated with LPS and IL-4+IL-12, we noticed that dORF-containing transcripts show no significant difference in TE compared to those without. (Figure 6).

**Signal sequence containing micropeptides**

An N-terminal signal peptide sequence of 16-30 amino acids is characteristic of proteins destined to be secreted, resident within cellular membranes or within compartments of the secretory pathway. We predicted the presence of signal peptides in amino acid sequences of micropeptides using SignalP server (40). This predicted 80 candidates including known chemokines (CCL-1, -2, -4, -5 and -22) and the cell surface protein CD52, as well as the recently identified lncRNA encoded Awi112010 (12) and 1810058I24Rik micropeptides (41). Amongst these, 28 are canonical micropeptides and they typically have high levels of conservation. Of the remaining 52 non-canonical micropeptides, 12 show conservation (Figure 7A).

We selected eight putative smORFs for further characterisation (Figure 7B). First of all, except for the uORF from Zdhhc5, all of uORF-encoded micropeptides are in different reading frames from the main CDS. The coding sequence of Zdhhc5 uORF does not overlap with the ZDHHC5 ORF, suggesting it is not an N-terminal extension of the main CDS. To examine the expression of signal sequence-containing micropeptide host transcripts, we compared mouse RNA-Seq datasets for lymphocytes spanning B cell terminal differentiation (42), Th1 cell activation (29), and regulatory T cells (43) as well as epidermal cells (44). These data revealed dynamic expression patterns for several of the host transcripts. For example, BC031181 was downregulated during B cell differentiation but upregulated during Th1 cell activation, it was also highly expressed in epidermal cells (Figure 7B). Host transcript expression patterns provide a lead to where and what stage of cell differentiation micropeptides may be produced and can help with experimental validation of micropeptide prediction. To determine if the selected putative micropeptides are likely functional, we compared the conservation of amino acid sequence between different mammalian species (Figure S4). All of the micropeptides including those encoded in uORFs show evidence of conservation. This indicates positive selection pressure of the coding sequence of these micropeptides.

**In vitro characterisation of predicted micropeptides with signal sequence**

We sought to validate the potential for secretion of micropeptides with signal sequences. To this end, we selected and cloned eight putative smORFs with predicted signal sequences into a dicistronic mammalian expression vector which allowed synthesis of the micropeptide with a FLAG epitope tag at its C-terminus and of GFP driven by an IRES from the same transcript. HEK293T cells transfected with micropeptide-encoding plasmids displayed anti-FLAG signals in both total cell lysates (C) and supernatant (S) fractions (Figure 7C). GFP was detected in all total cell lysates, but the smORF encoded in the Slc39a9 transcript showed no evidence of expression. The smORFs encoded by Ph21a (uORF), 1500011B03Rik and BC031181 (both canonical smORFs) showed the most abundant expression and secretion. The small ORFs encoded by Zdhhc5 and Tbp1 expressed less strongly but showed evidence of secretion. By contrast, the smORFs encoded by Opa1 and 1190007I07Rik were weakly or not at all secreted (Figure 7C). A recent report demonstrates that the human ortholog of 1190007I07Rik, named C12orf73, encodes a functional micropeptide named BRAWNIN found at the inner mitochondrial membrane and required for respiratory complex III assembly (45). These results demonstrate that putative smORFs can be expressed and secreted, but further investigations are required to validate their subcellular localisation and to demonstrate their biological functions.

**Discussion**

In this study we have developed an improved pipeline for discovery of novel smORFs expressed with low abundance. We used this pipeline to predict 5744 unique smORFs that show evidence of being translated in B and T lymphocytes in different conditions. Apart from 368 annotated as short CDSs or isoforms, the others are novel and located in long non-coding RNAs, pseudogenes or in the 5'UTR and 3'UTRs of canonical protein coding transcripts. By assessing the conservation of the amino acid sequences and their translation efficiency relative to long proteins we inferred whether the translation products of these smORFs were likely to be functional.

We compared our pipeline with Ribocode which assesses the triplet periodicity of RPFs in an ORF with modified Wilcoxon signed-rank test and is claimed to outperform other existing pipelines including RiboTaper, Rp-Bp and ORF-RATER (19,22,23). Ribocode has been incorporated into a recently published integrated tool (RiboToolkit) to analyse ribosome profiling data (25). When compared with ORFLine,
Among the predicted smORFs, 80% were located within 5’UTRs. Many of these were not conserved at the amino acid level between species and may be regulatory. A recent study has shown that non-canonical Hoogsteen-paired G-quadruplex (rG4) structures present upstream of some uORFs promote 80S ribosome formation on start codons, causing inhibition of translation of the downstream CDS (46). Identification of rG4 motifs in an uORF context thus may help to distinguish regulatory uORFs and this feature could be incorporated into new iterations of ORF calling pipelines.

About four percent of our predicted smORFs are dORFs, which is consistent with previously reported data (20). dORFs of less than 100 aa are poorly characterised. Our pipeline predicted three dORFs encoded by Pgs1, Sszr1 and Dpm2 genes which are conserved by amino acid sequences between human and mouse (data not shown). These dORFs likely encode functional proteins and deserve further investigation. Recently, Wu et. al. also reported that dORFs are poorly conserved in human and zebrafish genomes, but are readily translated and can enhance translation of the main CDS (47). Our results also demonstrate no evidence of a statistically significant effect of dORFs on translation of the upstream CDS. However, it is important to point out that we considered only 32 dORFs in LPS-activated B cells, compared with 1406 for human and 1153 for zebrafish by Wu et. al and therefore our analysis of TE may be underpowered to detect an effect.

We predicted 8 micropeptides with signal peptide sequences and five were found to be secreted in a model system. Little is known regarding the abundance or stability of these micropeptides in physiological settings. In our attempts to validate the predictions of novel smORFs, we observed that overexpression of smORFs in mammalian cells yielded varied levels of expression. Specifically, the uORF of Skc39a9 did not produce any detectable proteins despite codon optimisation, indicating possible short half-life of this smORF encoded protein. It has been suggested that many peptide products are selectively and rapidly degraded within cells, and hence are difficult to detect biochemically (48,49). These factors impede their identification by regular mass spectrometry as they are often lost during sample preparation thus not available for detection. Recently it was proposed that immunopeptidomics based on the repertoire of peptides presented by MHC class I molecules may be suitable for detection of smORFs translation products (50). The immunopeptidomics differs from the proteome in that it skews away from abundant gene products, enriching peptides from non-canonical translation initiation and micropeptides with short half-lives (51).

As the ability to predict smORFs far outstrips the ability to validate them experimentally only a small number of predicted smORFs have so far been validated. For further investigation of the biological functions of the potentially secreted micropeptides, investment into the generation of antibodies and model organisms will be required. For micropeptides with signal sequences, they have the potential to be novel cytokines. If so, it will be exciting to validate the existence of receptors and to shed light onto the biology of these micropeptides.

**Materials and Methods**

**Tissue culture**

B lymphocytes from the spleen or lymph nodes (LN) of C57BL/6 mice were isolated using the B Cell Isolation Kit (Miltenyi Biotech). For activation, B cells were cultured for 48 hours in RPMI 1640 Medium (Dutch Modification) supplemented with 10% FCS, 100 IU/ml penicillin, 100 μg/ml streptomycin, 2 mM L-GlutamaMAX (Gibco), 1 mM Sodium Pyruvate and 50 μM 6-mercaptopoethanol in the presence of 10 μg/ml of LPS (Sigma, E. Coll 0127: B8), 10ng/ml of IL4 and 10ng/ml of IL5. Naïve CD4+ T lymphocytes from spleen and LNs were isolated with CD4+CD62L+ T Cell Isolation Kit (Miltenyi Biotech) and stimulated in the same medium as for B cells using plate bound anti-CD3 (2C11) and 1 μg/ml of anti-CD28 (37.51) for 24 hours. HEK293T cells were maintained in DMEM (Gibco) with 10% FBS (Gibco) and 1× GlutaMAX-I (Gibco).

**cDNA Library preparation**

RNA-Seq libraries were generated using the TruSeq Stranded mRNA Sample Prep Kit (Illumina Inc). Ribo-Seq libraries were prepared using the ARTseq Ribosome Profiling Kit (Epicentre, Illumina) from cells treated with cycloheximide (CHX, 100 μg/ml) prior to cell lysis. cDNA libraries were sequenced using Illumina HiSeq1000, Illumina HiSeq2000 or Illumina HiSeq2500 system in a 100-bp single-end (RNA-Seq) or 50-bp single-end (Ribo-Seq) mode. Summary metrics of libraries used are described in Table S1.

**Reference genome, transcriptome and annotation**

GENCODE (52) reference genome sequences (mouse GRCh38/mm10) are downloaded from the GENCODE website (Table S2). Transcriptome sequences (cDNA sequences) and gene annotation downloaded from the same GENCODE source are used to search for putative ORFs. RNA sequences are downloaded from UCSC Table Browser (53), RNA sequences are downloaded from GENCODE (version M20, we have also tested M13 and M15) as well as published studies (18,19). The transcriptome is defined as the collection of all transcripts on the reference chromosomes. GENCODE Transcript biotypes are defined here - https://www.gencodegenes.org/pages/biotypes.html. In our pipeline, we remove the following biotypes:

- Ig_* and TR_* (Immunoglobulin variable chain and T-cell receptor genes)
- miRNA
- misc_RNA
- Mt_rRNA and Mt_tRNA
- rRNA and ribozyme
- scaRNA, scRNA, snRNA, snRNA and sRNA
- nonsense_mediated_decay
- Non_stop_decay

Fig. 7. Predicted signal sequence containing micropeptides and their host transcripts expression under different conditions. (A) Scatter plots show the distributions of length (codon) and PhyloCSF score for each predicted signal peptide containing micropeptides. (B) Heatmap analysis of host transcript expression during B cell terminal differentiation. Thy1 cell activation, B cells were cultured for 48 hours in RPMI 1640 Medium (Dutch Modification) supplemented with 10% FCS, 100 IU/ml penicillin, 100 μg/ml streptomycin, 2 mM L-GlutamaMAX (Gibco), 1 mM Sodium Pyruvate and 50 μM 6-mercaptopoethanol in the presence of 10 μg/ml of LPS (Sigma, E. Coll 0127: B8), 10ng/ml of IL4 and 10ng/ml of IL5. Naïve CD4+ T lymphocytes from spleen and LNs were isolated with CD4+CD62L+ T Cell Isolation Kit (Miltenyi Biotech) and stimulated in the same medium as for B cells using plate bound anti-CD3 (2C11) and 1 μg/ml of anti-CD28 (37.51) for 24 hours. HEK293T cells were maintained in DMEM (Gibco) with 10% FBS (Gibco) and 1× GlutaMAX-I (Gibco).
Overview of strategy to identify candidate smORFs

Using the nucleotide sequences of all transcripts downloaded from GENCODE (release M13) (54) as a reference, ORFLine searches for putative ORFs beginning with “AUG”, “TUG”, “CUG”, “GUG” and ending with “UAG”, “UAA”, “UGA” in each of the three reading frames. It then removes ORFs that are not n*3 (n > 1) nucleotides long and designates those that are between 10 to 100 codons in length as putative smORFs. The ORF coordinates are initially transcript coordinates and are converted to genomic coordinates given exon location information in the gene annotation (in GTF/GFF format), the output of this step are genomic coordinates and strands for putative smORFs in BED format. Each ORF will be assigned two different identifiers, one is called RegionId, the second is called ORFId. RegionId is created based on genomic coordinates, ORFId is created based on the transcript coordinates. An ORF has a unique genomic location, thus RegionId is unique, but it may arise from multiple overlapping transcripts, so it may have multiple ORFIds (Figure S1). This step is carried out once only and needs to be updated when transcriptome annotation is changed.

Sequencing data processing

Raw Illumina sequencing data in FASTQ format are trimmed of adapter sequences and the resultant reads are aligned against specific sequences assembled from a collection of rRNA, tRNA, Mt_rRNA and Mt_rRNA snRNA, snoRNA, misc_RNA and miRNA sequences using Bowtie v1.2.2 (55) to remove these sequences. The remaining reads are aligned to the reference genome (GRCm38). Adaptor trimming and quality trimming (including poor quality “N” base at the 5’ end of some of the reads) were performed with Trim Galore v0.4.5 (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore), quality checked with FastQC v 0.11.8 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc). Authentic RPFs will be ~28-30nt in length, therefore, we have kept trimmed reads that have a length between 25 and 35 nt, as they account for ~75% of the total reads on average (refer to information in Table S1).

Alignment to reference genome

The reads were mapped to the GRCm38/mm10 reference genome using the STAR aligner v2.5.2a (56). The aligner reports only uniquely mapped reads (mapping quality MAPQ ≥ 255). The following shows an example command, and parameters are in bold:

```
STAR --runThreadN $THREAD --genomeDir SRFGENOMESTAR --readFilesIn SOUTPATH/bowtie-contaminant-removal/$NAME_trimmed_unfiltered.tqz --readFilesCommand zcat --outReadsUnmapped Fastx --outFileNamePrefix $OUTPATH/star-genome/$NAME/
```

--alignIntronMin $ALIGNINTRON_MIN
--alignIntronMax $ALIGNINTRON_MAX
--alignEndsType EndToEnd
--outFilterMismatchNmax $MISMATCH_MAX
--outFilterMismatchNoverLmax $MISMATCH_NOVERL_MAX
--outFilterType $FILTER_TYPE
--outFilterIntronMotifs RemoveNoncanonicalUnannotated
--outSAMattributes $SAM_ATTR
--outSAMtype BAM SortedByCoordinate
--outBAMsortingThreadN $THREAD

Transcript expression quantification

In each experiment, sequence alignments (in BAM format) of all biological replicates were combined for RNA-Seq and Ribo-Seq respectively. Transcript expression was quantified using StringTie v1.3.6 (57) in FPKM (Fragments Per Kilobase per Million). From a given dataset, a minimal expression level was set to FPKM > 0.5 (27) to exclude non-conserved transcripts.

P-site offset determination

A majority of RPFs has a length between 28-31 nucleotides (nt). The 5' P-site offset is the distance from the 5' end of the read to the ribosomal P-site (15). To determine P-site offset, we separated footprints into groups based on their length, P-site offset was calculated for each read length, using plastid python library v0.4.8 (58). We observed P-site offsets are 12 nt long for RPF in 28-31 nt in our experiments.

Identification of translated smORFs

ORFLine takes the gene annotation, putative smORFs, Ribo-Seq and RNA-Seq alignment as input to predict actively translated smORFs. ORFLine combines alignment files of all biological replicates (pooled analysis) to increase the signal intensity in case the smORFs are lowly expressed. This component consists of several metrics and filters, putative smORFs that have exceeded a confidence threshold for each metric (as indicated in Table 1) were kept.

RPF coverage

To filter ORFs which are insufficiently covered by reads, we calculated the proportion of codons being covered by RPFs. We consider a codon covered if there is a mapped RPF with the P-site aligned to nucleotide 1 of that codon. An ORF is discarded if the ratio of covered codons to the total number of codons in the ORF < 0.1 (18).

ORFScore

ORFScore was proposed by Bazzini and colleagues (18) and we re-implemented the ORFScore algorithm in R. The ORFScore was then calculated as:

where \( F_n \) is the number of reads in reading frame \( n \), \( \overline{F} \) is the total number of reads across all three frames divided by 3. RPFs were counted at each position within an ORF, excluding the first and last coding codons. To filter out putative artificial peaks, the most abundant read position was masked if reads aligning to that position comprised more than 70% of the total reads in the ORF. The ORFScore is a log-scaled chi-squared goodness of fit test statistic, the p-values associated with the test were adjusted using Benjamini-Hochberg FDR-controlling method and smORFs with ORFScore > 0 and adjusted p-value < 0.01 were retained.

Ribosome Release Score (RRS)

Firstly, the 3’UTRs of smORFs is defined. For canonical smORFs, we used annotated 3’UTRs. For other classes of smORFs, their 3’UTRs were defined as the region between the stop codon and the next possible stop codon in any frame. The RRS score is defined as the ratio of the two normalized ratios and calculated with the following equation: RRS = (FPKM_RF ORF/FPKM_RF 3’-UTR)/ (FPKM_RNA ORF/FPKM_RNA 3’-UTR). Based on the original study, smORF with RRS > 5 is considered to be translated (26).

Inside/outside read ratio

The ribosome footprints typically show precise positioning between the start and the stop codon of translated ORFs. Low density of footprints before start codons and after stop codons and high inside/outside ratio is expected. By considering the read distribution of the nearest 3 upstream codons outside and the first 3 codons inside an ORF, we used a feature called inside/outside read ratio (total RPF of inside 3 codons/total RPF of outside 3 codons) to assess whether genuine translation takes place. ORFs will be discarded if the ratio ≤ 1 (more reads mapping outside than inside).

Analysis of predicted smORFs

Translation efficiency (TE)

A measure of the rate of translation for a given feature (e.g. the CDS of a mRNA or a smORF), obtained in ribosome profiling experiments. It was calculated as the base 2 logarithmic ratio of RPF expression (FPKM) over mRNA expression (FPKM).

Conservation of the amino acid sequences

To examine the conservation of smORF-encoded micropeptide sequences between species, we performed PhyloCSF (59), a likelihood-based method to analyze signatures of evolutionary conservation in multiple species sequence alignments. PhyloCSF assigns a score to each smORF based on conservation within selected vertebrate species (https://github.com/mlin/PhyloCSF/wiki#available-phylogenies). For each smORF, we selected alignments of mouse, human, chimpanzee, gorilla, cow, dog and zebrafish from a publicly available whole genome multiple alignment using Galaxy “stitch gene blocks” tool (60). smORFs were considered conserved if their PhyloCSF score was > 50 (26), and weakly
conserved if they had a PhyloCSF score > 0. PhyloCSF score = 0 indicates that there is no DNA sequence alignment cross species and PhyloCSF score < 0 is considered not conserved.

**Gene ontology (GO) enrichment analysis**

We used the g:Profiler server (61) to perform GO analysis in two unranked lists of genes mode. The background list comprised the combined expressed transcripts (FPKM > 0.5) of B and T cells. The target list contains the host transcripts of the smORFs. For the significance threshold, we chose the default option g:SCS threshold and combined expressed transcripts (FPKM > 0.5) of B and T cells. The unranked lists of genes mode. The background list comprised the unranked lists of genes mode. The background list comprised the unranked lists of genes mode. The background list comprised the unranked lists of genes mode. The background list comprised the unranked lists of genes mode. The background list comprised the unranked lists of genes mode. The background list comprised the unranked lists of genes mode.

**Cloning and expression of candidate secreted micropeptides**

The coding sequences of predicted smORFs were codon optimised for mammalian expression (GeneArt from ThermoFisher) and cloned into a mammalian expression vector. A C-terminal FLAG epitope tag was placed downstream of the micropeptide cDNA sequences flanked by a spacer sequence of Glycine-Alanine-Alanine. This is followed by an EMCV-IRES upstream of GFP cDNA. The di-cistronic mRNA is controlled by a CAG promoter. For transfection, HEK293T were seeded at 60-90% confluency the day before, and transfected at 60-90% confluency. Transfected cells were replaced with fresh media to remove possible plasmid contamination in downstream assays. 21-24 hours post transfection, cells were collected by centrifugation at 300 × g for 5 minutes at 4 °C. For Western blot analysis, total cell lysates and total supernatant were acquired and analysed using the Odyssey CLx (LI-COR).

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**Conflict of Interest**

We declare no conflicts of interest.

**End Matter**

**Data Availability**

**Cell type** | **RNA-Seq** | **Ribo-Seq**
--- | --- | ---
B cell setup 1 | GSE62129 | GSE62134
B cell setup 2 | GSE146073 (GSM4364117-121) | GSE154491 (all samples)
CD4+ T cell | GSE155087 (GSM4694962-64) | GSE155087 (GSM4694968-972)
Th1 reactivation | GSE83351 | GSE83351

**Pipeline code availability**

Pipeline code is publicly available on the source code hosting platform GitHub. The URL is https://github.com/boboppie/ORFLine. We also created a Singularity image (https://singularity.lbl.gov/) which enables the users to execute and test the pipeline easily in a virtual environment. All dependencies including bioinformatics tools are pre-installed in the image, the URL is https://github.com/boboppie/ORFLine-singularity.

**Supplementary Data**

Supplementary Data are available at BioRxiv online.
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## Tables

| Component | Step | Cut-offs and rationales | Input | Output |
|-----------|------|-------------------------|-------|--------|
| **Prediction of all putative smORF** | Computational prediction of all theoretically possible smORFs | Scan all annotated transcripts | Transcriptome sequences; user defined start codons (default: “AUG”, “TUG”, “CUG”, “GUG”) | List of genomic regions with predicted smORFs |
| **Ribo-Seq and RNA-Seq QC and processing** | Trimming of sequencing adapters and low-quality bases | Trim adapters and low-quality bases from 3’ end of reads | Ribo-Seq and RNA-Seq FASTQ files | Trimmed FASTQ files |
| | Removal of reads mapped to rRNA and tRNA loci | Align all reads to rRNA/tRNA sequences | Trained Ribo-Seq FASTQ files; rRNA/tRNA sequences | rRNA/tRNA depleted FASTQ files; rRNA/tRNA alignment BAM files |
| | Alignment to the reference genome | Unique mapped reads are kept; read length < 36 nt | rRNA/tRNA depleted FASTQ files | Genome alignment BAM files; unmapped FASTQ files |
| | Calculation of P-site offset for Ribosome protected fragments (RPFs) of each length | Reads are split into groups on the basis of length and the 5’ end mapping rule is applied to each group | Genome alignment BAM files | List of P-site offset values for reads of each length |
| | Evaluation of triplet periodicity for all RPFs | The first reading frame should show a greater share of reads (> 50%) | Genome alignment BAM files | Summary of reads share for each reading frame |
| | Estimate of transcript expression (by FPKM values) | Transcripts with FPKM > 0.5 are considered expressed (27) | Genome alignment BAM files | List of FPKM values for each transcript |
| **Identification of translated smORFs** | Filtering out reads of lengths for which triplet periodicity pattern is not observed | Reads are split into groups on the basis of length; only groups which show triplet periodicity are kept | Merged genome alignment BAM file; read phasing summary | Read length filtered BAM files |
| | Filtering out regions with putative smORFs which have no aligned reads | Read count > 0 | Read length filtered BAM files; full list of predicted smORFs | Read count filtered smORF regions |
| | Filtering out regions with putative smORFs which have no aligned RPFs | RPF count > 0 (some regions retained in the previous step may contain only reads not derived from RPFs) | Filtered BAM files; read count filtered smORFs | RPF count filtered smORF regions |
| | Filtering smORFs based on expression of host transcripts | Only keep smORFs located on transcript with FPKM > 0.5 | Host transcript FPKM values; RPF count filtered smORFs | Transcript expression filtered smORFs |
| | Assigning classes to smORFs on the basis of their position on the host transcripts | Class is added as annotation; the number of smORFs is unchanged | Transcript expression filtered smORFs | Class annotated smORFs |
| | Filtering smORFs on the RPF coverage > 0.1, ORFScore > | | Class annotated smORFs | ORFScore filtered |
| basis of ORFScore | 0, adjusted p-value < 0.01 | smORFs |
|-------------------|--------------------------|--------|
| Filtering out smORF overlapping with a CDS | The estimated ratio (RPF countCDS/RPF countsmORF) > 1 | Read length filtered BAM files; ORFScore filtered smORFs |
| Filtering smORFs on the basis of Ribosome release score (RRS) | RRS > 5 | Read length filtered BAM files; RNA-Seq alignment BAMs; Region filtered smORFs |
| Filtering smORFs with same stop but different start codons (nested smORFs) | The smORF with the maximum ORFScore is retained, otherwise a smORF with AUG start codon is retained | RRS filtered smORFs |
| Filtering smORFs on the basis of Inside/outside (I/O) ratio | smORFs are retained if inside/outside read ratio > 1 (more reads mapping inside than outside) | Read length filtered BAM files; nested filtered smORFs |

Table 1. Summary of steps in the ORFLine.
| Column | Description |
|--------|-------------|
| 1 - 12 | The first 12 columns are in BED12 format, the fields are described here - https://genome.ucsc.edu/FAQ/FAQformat.html#format1. The 4th column is ORFId (transcript-based). |
| 13     | smORF class, including canonical, five_prime... |
| 14     | Peptide length |
| 15     | RegionId (genomic-based) |
| 16     | Ensembl transcript Id |
| 17     | Gene symbol |
| 18     | Gene description |
| 19     | ORF score |
| 20     | Ribosome release score |
| 21     | Ribo FPKM |
| 22     | RNA FPKM |
| 23     | Translation efficiency (TE) |
| 24     | CDS TE (NA if host transcript is noncoding) |
| 25     | AA sequence |

Table 2. Pipeline final output format.
| Class                                                | ORFLine          | RiboCode | Commonality | Unique to ORFLine | Unique to RiboCode |
|------------------------------------------------------|------------------|----------|-------------|-------------------|--------------------|
| Annotated (including canonical/canonical_extended)   | 368 (338 canonical + 30 canonical_extended) | 290      | 183         | 185               | 107                |
| Novel (noncoding)                                    | 501              | 990      | 69          | 432               | 921                |
| nuORF                                                | 4,174            | 5,133    | 1,401       | 2,773             | 3,732              |
| ouORF                                                | 441              | 1,718    | 185         | 256               | 1,533              |
| ndORF                                                | 243              | 506      | 121         | 122               | 385                |
| odORF                                                | 17               | 231      | 3           | 14                | 228                |
| Total                                                | 5,744            | 8,868    | 1,957       | 3,787             | 6,911              |

Table 3. ORFLine and RiboCode prediction commonality/difference by class.