Widespread translational control regulates retinal development in mouse

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ABSTRACT

Retinal development is tightly regulated to ensure the generation of appropriate cell types and the assembly of functional neuronal circuitry. Despite remarkable advances have been made in understanding regulation of gene expression during retinal development, how translational regulation guides retinogenesis is less understood. Here, we conduct a comprehensive translatome and transcriptome survey to the mouse retinogenesis from the embryonic to the adult stages. We discover thousands of genes that have dynamic changes at the translational level and pervasive translational regulation in a developmental stage-specific manner with specific biological functions. We further identify genes whose translational efficiencies are frequently controlled by changing usage in upstream open reading frame during retinal development. These genes are enriched for biological functions highly important to neurons, such as neuron projection organization and microtubule-based protein transport. Surprisingly, we discover hundreds of previously uncharacterized micropeptides, translated from putative long non-coding RNAs and circular RNAs. We validate their protein products in vitro and in vivo and demonstrate their potentials in regulating retinal development. Together, our study presents a rich and complex landscape of translational regulation and provides novel insights into their roles during retinogenesis.

INTRODUCTION

The vertebrate retina is a specialized part of the central nervous system with diverse cell types, high-level organization and an evolutionarily conserved structure (1). It can serve as an ideal model to study neural development, such as deciphering the developmental gene regulatory patterns and understanding mechanisms of morphogenesis formation and specificity (2). To date, genome-wide molecular characterization of retinogenesis has been understood using transcriptomic (3), epigenomic (4) and proteomic (5) approaches and has identified many molecules that play important roles in regulating the development of retina.

The central dogma of molecular biology states two major steps during the detailed residue-by-residue transfer process of genetic information: transcription and translation, by which information encoded in DNA flows into RNA via transcriptional regulation and ultimately to proteins via translational regulation (6). Similar to transcription, translation involves a series of highly temporally orchestrated events directed by cis-elements and trans-factors (7). Increasing evidence emphasizes the importance of translation of gene expression (8,9). Dynamic, tight, and coordinated translational regulation can contribute to the growth of multicellular organisms, particularly during a rapid morphological transition, such as development of red blood cell (10), embryonic stem cell differentiation (11), cortical neurogenesis (12), myogenic differentiation (13) and spermioteleosis (14). In addition, it is suggested that translational regulation may influence plasticity of visual pathway development and function (15). Nevertheless, a systematic, genome-wide analysis of gene translation for retinogenesis is lacking at present.

Recently, accumulating evidence has shown that a fraction of small open reading frames (ORFs) within putative long non-coding RNAs (lncRNAs) are translated to encode functional micropeptides (16). For example, translational product of a lncRNA, Dworf, is of critical importance in regulating contraction-relaxation cycles in muscle (17). More recently, some circular RNAs (circRNAs) have been shown to encode bioactive micropeptides, with specific cellular and physiological functions (18,19). These unexpected findings have further emphasized the complex translational regulation in modulating gene expression. However, it is en-
tirely unclear whether the translation of non-coding RNAs and non-canonical ORF-mediated translational control exist during retinal development.

Herein, we conducted the first survey of the translational landscape of retinal development in mouse. We applied ribosome profiling (Ribo-seq), mRNA sequencing (mRNA-seq), and circRNA sequencing (circRNA-seq) to the developing mouse retina from the embryonic to the adult stages. We found that translation was dynamically uncoupled with transcription, particularly with larger expression divergence before eye-opening. We revealed diverse regulatory changes fulfilling the requirements of gene expression outputs at different developmental stages. We further detected dynamic changes in translational efficiency (TE) and discovered thousands of upstream ORFs (uORFs) fine-tuning gene translation. Surprisingly, we identified hundreds of actively translated IncRNAs and ribosome-associated circRNAs, which were highly developmental stage specific. We validated their translation in vivo and in vitro and annotated their potential functions in retinal development. Overall, our study provides a snapshot of complex dynamic patterns of gene translation in mouse retinal development and gains new insights into the regulatory principles of gene translation, particularly non-canonical ORF-mediated translational control, during retinogenesis and neural development.

MATERIALS AND METHODS

Animals and tissue collection

Wild-type mice of C57BL/6J genetic background were purchased from Guangdong Medical Experimental Animal Center. Mice from seven different developmental stages including E13 (Embryonic day), E15, P0 (Postnatal day), P6, P13, P21 and M9 (Month) were euthanized, and the eyes were enucleated immediately after sacrifice, and further retinal tissues from enucleated eyes of each mouse were harvested. In detail, two litters of mice per stage were used to represent two biological replicates. The retinal tissues of mice from the same litter were pooled, snap-frozen in liquid nitrogen, and then stored at −80°C prior to library preparation for next-generation sequencing. Notably, retinal tissues of E13, E15, P0, P6, P13 and P21 were separately used for paired mRNA-seq and Ribo-seq; retinal tissues of E15, P0, P6, P21 and M9 were separately used for circRNA-seq; and retinal tissues of M9 were also used for Ribo-seq. All experimental procedures were approved by the Animal Ethics Committee of Zhongshan Ophthalmic Center, Sun Yat-sen University (Guangzhou, China; License No.: SYXX (YUE) 2018-0189).

Library preparation and sequencing

Frozen retinal samples per stage were lysed using 1 ml of mammalian lysis buffer (200 μl of 5× Mammalian Polysome Buffer, 100 μl of 10% Triton X-100, 10 μl of DTT (100 mM), 10 μl of DNase I (1 U/μl), 2 μl of cycloheximide (50 mg/ml), 10 μl of 10% NP-40 and 668 μl of nuclease-free water). After incubation for 20 min on ice, the lysates were cleared by centrifugation at 10 000 × g and 4°C for 3 min. The lysate was divided into 300-μl aliquots for Ribo-seq and in parallel mRNA-seq, respectively. In detail, for the 300-μl aliquots of clarified lysates, 5 units of ARTseq Nuclease were added to each A260 lysate, and the mixtures were incubated for 45 min at room temperature. Nuclease digestion was stopped by additional 15 μl of SUPERase In RNase Inhibitor (Ambion). Subsequently, the lysates were applied to Sephaacyl S400 HR spin columns (GE Healthcare Life Sciences), and ribosome-protected fragments were purified using the Zymo RNA Clean & Concentrator-25 kit (Zymo Research). Ribosomal RNA was depleted using the Ribo-Zero magnetic kit (Epicentre). Sequencing libraries of ribosome-protected fragments were generated using the ART-seq™ Ribosome Profiling Kit (Epicentre, RPHMR12126) according to the manufacturer’s instructions. From the 300-μl aliquots of clarified lysates, poly(A)+ RNAs were extracted and purified, and sequencing libraries of poly(A)+ RNAs were then generated using the VAHTSTM mRNA-seq v2 Library Prep Kit from Illumina (Vazyme Biotech, NR601-01) according to the manufacturer’s instructions. The resulting 26 barcoded libraries were pooled and sequenced using an Illumina HiSeq 2500 instrument in the single-end mode and in a randomized order across lanes.

For circRNA sequencing, total RNA was extracted using Trizol reagent (Invitrogen, Carlsbad, CA, USA), the RNA integrity was assessed by Agilent 2100 with RIN number > 7.0. Approximately 5 μg of total RNA was used for library preparation, the Ribo-Zero™ rRNA Removal Kit (Illumina, San Diego, USA) was used to deplete ribosomal RNA according to the manufacturer’s instructions, the left RNAs were treated with RNase R (Epicentre Inc., Madison, WI, USA) to remove linear RNAs and to enrich circRNAs. The circRNA sequencing libraries were then generated using TruSeq Stranded Total RNA HT Sample Prep Kit (Illumina RS-122–2203) according to the manufacturer’s instructions with minor modifications. Briefly, cDNAs were reverse transcribed using ProtoScript II Reverse Transcriptase. The ligated products were amplified with PCR by the following conditions: initial denaturation at 95°C for 3 min; 8 cycles of denaturation at 98°C for 15 s, annealing at 60°C for 15 s, and extension at 72°C for 30 s; and then final extension at 72°C for 5 min. At last, the resulting 10 barcoded libraries were sequenced using an Illumina Hiseq X Ten instrument in the paired-end mode.

mRNA-seq and Ribo-seq alignment

The raw Ribo-seq and mRNA-seq sequencing reads were demultiplexed using CASAVA (version 1.8.2), followed by adapter trimming with Cutadapt (v1.9.1, -e 0.1 -o 6 -m 20) (20) and removal of poor-quality reads with Sickle (v1.33, -q 20 -l 20 -x -n; available at https://github.com/najoshi/sickle). The reads aligned to mouse rRNA and rRNA sequences using Bowtie (v1.0.1, -l 20) (21) were further removed. All remaining reads were mapped to the mouse reference genome (GENCODE, Release M18: GRCh38.p6) using STAR (v2.5.2) with default parameters. Only uniquely mapped
reads selected by samtools (v1.6, Phred score ≥ 20) were used for subsequent analysis.

**Gene quantification and expressed genes definition**

Raw counts for different genomic features were obtained using featureCounts from the subread package in Bioconductor (v1.5.0, -t CDS for Ribo-seq data and -t exon for mRNA-seq data) (22). The raw counts from all 24 samples were then combined and normalized together by a pooled-based size factor yielded from the DESeq2 R package (23) to minimize the batch effects among samples. After that, the expression level of each gene was estimated as reads per kilobase of transcript per million reads mapped (RPKM) using an in-house R script. At the transcriptome layer, only those genes with RPKM > 1 across two replicate samples of each developmental stage were kept, defined as well-transcribed genes. At the translome layer, only those genes that are well-transcribed and undergo active translation (see below) were defined as well-translated genes.

**Actively translated ORF detection**

Actively translated ORF detection in Ribo-seq data was performed using RiboTISH (v0.2.4) with ‘-framebest’ strategy to select the best candidate ORF (24). To increase statistical power of ORF finder, we merged the aligned BAM files from two replicated samples of each developmental stage using ‘samtools merge’ (v1.6). Upstream ORFs (uORFs) were defined as ORFs originating from the 5′-UTR of annotated protein-coding genes (i.e. with TisType: 5′-UTR); downstream ORFs (dORFs) were defined as ORFs originating from the 3′-UTR of annotated protein-coding genes (i.e. with TisType: 3′-UTR); and IncORFs were defined as ORFs originating from annotated IncRNA genes.

**Genes with dynamical changes detection**

Genes with dynamic expression patterns over time were identified by the maSigPro R package, which is specifically designed for the analysis of time-course gene expression data (25). The method is a two-regression step approach where the experimental groups are identified by dummy variables. It first adjusts a global regression model with all the defined variables to identify differentially expressed genes, and in second a variable selection strategy is applied to study differences between groups and to find statistically significant different profiles. Genes with significant temporal changes in retinal development were selected at a FDR <0.05 and R-squared threshold equal to 0.6.

**Translational efficiency estimation**

The TE for each protein-coding gene was estimated as the ratio of the normalized RPKM values of Ribo-seq to mRNA-seq reads in annotated CDS regions (26). Given a high degree of TE correlation between two replicated samples of each developmental stage, particularly when only focusing on those well-transcribed genes (mean Pearson’s correlation coefficient = 0.95), TE values of the two replicates for each protein-coding gene were then averaged to form a final TE for subsequent comparative analysis.

**Differential expression analysis, gene classification, and gene ontology (GO) enrichment analysis**

Differential expression analysis was performed using deltaTE (27), which incorporates Ribo-seq and mRNA-seq data. Briefly, deltaTE considers sample-to-sample variance and introduces an interaction term into the statistical model of DESeq2, and the interaction term is used to model condition (i.e. developmental stages) and sequencing methodology (i.e. Ribo-seq or mRNA-seq). In order to identify significant differences between conditions that are discordant between sequencing methodologies, its generalized linear model incorporates three components: the condition, the sequencing type, and an interaction term containing both. For each gene, a false discovery rate (FDR) threshold of 5% is used to determine statistical significance of the resulting change in RPFs (ΔRPF), mRNA counts (ΔRNA), and TE (ΔTE), respectively, and the ΔRPF, ΔRNA, and ΔTE are combined to determine its regulatory group of belonging. In detail, genes with change in mRNA and RPF levels at the same rate were defined as differentially transcribed genes (DTG), and genes with change in RPF level independent of change in mRNA level, which lead to a change in TE, were defined as differential translational efficiency gene (DTEG). DTGs and DTEGs between adjacent stages could further be categorized into four classes: buffered, intensified, forwarded and exclusive. Specifically, translationally buffered genes have a significant change in TE that offsets the change in RNA; translationally intensified genes have a significant change in TE that bases on the effect of transcription; translationally forwarded genes are DTGs that have a significant change in mRNA and RPF at the same rate and with no significant change in TE. Conversely, translationally exclusive genes are DTEGs that have a significant change in RPF but with no change in mRNA leading to a significant change in TE.

GO enrichment analysis was used to reveal biological functions of differentially expressed genes (DEGs), which was achieved by the ClusterProfiler R package (available at http://bioconductor.org/packages/release/bioc/html/clusterProfiler.html) using all genes as the background. Only those GO terms with false discovery rate (FDR) < 0.01 were regarded as statistically significant.

**Principal component analysis**

To define the main contributing layer of gene expression regulation (buffered, exclusive, forwarded, and intensified) for each coregulatory biological function, principal component analysis (PCA) was performed as described in a previous study (19). For each GO term, the relative fractions of four defined classes of differential genes were used as an input for the PCA. The prcomp and fviz_pca_biplot functions from the factoextra R package (available at http://www.sthda.com/english/rpkgs/factoextra) were used for the PCA and visualizing the output of the PCA, respectively. Thus, the placement of each GO term in the PCA plot was
based on the directionality of four layers of gene expression regulation.

Sequence conservation analysis
To calculate evolutionary conservation of non-canonical ORFs in sequence, we downloaded phastCons tracks from UCSC based on 60 vertebrate species (http://hgdownload.soe.ucsc.edu/goldenPath/mm10/phastCons60way, August 2019), and then retrieved the base-by-base conservation scores from ‘mm10.60way.phastCons60wayPlacental.bw’. For each ORF, the mean PhastCons score at each base was used to evaluate the degree of its cross-species conservation. It varies on a scale from 0 to 1, with 0 indicating to be poor conserved and 1 indicating to be strong conserved.

CircRNA identification and quantification
Raw circRNA-seq reads were pre-processed with Perl scripts, including the removal of adaptor-polluted reads, low-quality reads (Phred score \( \geq 20 \)) and reads with number of \( N \) bases accounting for \( \geq 5\% \). The clean reads were then mapped to the mouse reference genome (GENCODE, GRCh38.p6) using BWA-MEM (-T 19, v0.7.17; available at http://bio-bwa.sourceforge.net/). Next, two different detection tools were used to identify transcribed circRNAs, namely, CIRI2 (v2.0.6) (28) and CIRCexplorer2 (v2.3.6) (29). To further reduce false positives of circRNA identification, only those circRNAs meeting all of the following three criteria were kept, including (i) having at least two unique backsplice junction (BSJ) reads, (ii) being simultaneously identified in both tools and (iii) being simultaneously detected in two replicated samples of each developmental stage. Finally, the BSJ reads of CIRI2 were used to quantify the transcriptional level of circRNA using CPM (counts per million mapped reads) (30).

Identification of ribosome associated-circRNAs
To determine ribosome-associated circRNAs, we first extracted the 40-base pair (bp) sequences on either side of the backsplice junction site of each transcribed circRNA, and then the sequence was ligated in tandem to generate a pseudo circRNA reference. Next, all Ribo-seq reads that failed to map to the linear reference genome were realigned to the pseudo circRNA reference sequences using Tophat2 (v 2.1.1) with default parameters except \( N \), which was set to 0 (the default is 2) (31). Finally, ribo-associated-circRNAs (ribo-circRNAs) were defined as having (i) at least one unique backsplice junction-spanning Ribo-seq reads and (ii) a minimum read-junction overlap of three nucleotides (nt) on either side of the backsplice junction site. Notably, the backsplice junction-spanning Ribo-seq reads might arise from linear, transspliced products (trans-spliced RNAs; tsRNAs) rather than circRNA molecules. To further reduce false positive results, we took advantage of our poly(A)+ RNA-seq (mRNA-seq) data to detect BSJ reads and identify potential tsRNAs expressed in each stage, considering that circRNAs are generally non-polyadenylated but tsRNAs are not. The ribo-circRNAs sharing the same splicing junction site with these tsRNAs were then excluded from subsequent analyses.

CircRNA ORF prediction
To predict putative circRNA-encoded ORF (cORF), the cORF_prediction_pipeline was used with some modifications (32). Briefly, the full-length sequence of each ribo-circRNA was retrieved and multiplied four times to allow for rolling circle translation. All possible cORFs with an AUG-start codon followed by an in-frame stop codon in the exonic sequence were identified and then filtered based on the requirements of a minimum length of 20 amino acids (aa) and of spanning the backsplice junction site. Notably, only the longest cORF was retained for each of the three frames per ribo-circRNA. If a cORF does not contain stop codon, it was defined as an INF (infinite)-cORF, representing that the circRNA could be translated via a rolling circle amplification mechanism.

Two-dimensional liquid chromatography–tandem mass spectrometry
Retina tissues at E13, E15, P0, P6, P13 and P21 were mixed, fully ground in liquid nitrogen and lysed with lysis buffer L3 (Fitgene Biotech, #FP1801), 0.2% SDS and 1x PMSF (Sangon Biotech, #P0754). Lysates were sonicated on ice and cleared by centrifugation at 12 000 rpm for 10 min. The protein extracts were purified by overnight acetone precipitation and re-solubilized using L3, concentration of the samples was determined by Bradford assay. Proteins were reduced using 50 mM TCEP (1 h at 60°C), alkylated using 55 mM MMTS (45 min at RT in the dark), then the protein sample was loaded on a 10 kDa ultrafiltration tube, washed twice with 8 M urea and three times with 0.25 M TEAB. For protein digestion, 50 \( \mu \)l of 0.5 M TEAB and trypsin (Promega, enzyme: protein ratio of 1:50) were added to the membrane, reaction was incubated overnight at 37°C. The resulting peptides were first fractionated on a Gemini-NX 3u C18 110Å 150’2.00 mm columns (Phenomenex, #00F-4453-B0) using high-pH reversed-phase chromatography (Dionex Ultimate 3000 RSLCnano) with increasing concentration of acetonitrile, 20 fractions were collected according to the 214 nm absorbance and running time. After vacuum drying, 3 \( \mu \)g of fractionated peptide samples were separated on an Acclaim PepMap RSLC C18 2 \( \mu \)m 100 Å 75 \( \mu \)m i.d. \( \times \) 150 mm column (Dionex, #160321) and analysed using Thermo Scientific™ Q Exactive™. Full MS spectra from \( m/z 375–1800 \) were acquired at a resolution of 70 000 with an automatic gain control (AGC) target value of 3e6 and maximum injection time (IT) of 40 ms. MS/MS spectra were obtained at a 17 500 resolution with an AGC target of 1e5 and maximum injection time (IT) of 60 ms, TopN was set to 20 and NCE/stepped NCE was set to 27. In addition, two unfractionated peptide samples were also analysed by LC–MS/MS using the same parameters as fractionated peptide samples.

Analysis of mass spectrometry-based proteomic data
Three publicly available proteomic datasets obtained from the PRIDE database (accession numbers: PXD003441,
PXD002247 and PXD009909) and an in-house MS dataset were used to detect protein products from translatable lncRNAs and circRNAs. The raw data files were analysed using MaxQuant software (v1.6.15.0) (33) against a custom-made database, which combined all mouse sequences from UniProt/Swiss-Prot (MOUSE.2020-08) with sequences derived from translatable lncRNAs and circRNAs, based on the target decoy strategy (Reverse) with the standard search parameters with the following exceptions: (i) the peptide-level FDR was set to 5%, and the protein-level FDR was excluded; (ii) the minimal peptide length was set to seven amino acids and (iii) a maximum of two missed cleavages was allowed. Each search included carbamidomethylation of cysteine as fixed modification, methionine oxidation, N-terminal acetylation as variable modifications, but for PXD002247 and our own MS data, a variable modification of deamidation of asparagine and glutamine was also included.

Functional annotation of IncORF-encoded peptides

Conserved domain and protein homology detection. Each of putative micropeptides encoded by IncORFs was annotated against the InterPro database using InterProScan (v5.44) with the default parameters (34). In total, 212 of 603 micropeptides were mapped to known homologous records in the InterPro database, of which 68 were annotated with specific functional domains involving in vital important pathways (Supplementary Table S1).

Guilt-by-association approach. A total of 198 translatable lncRNAs with dynamical changes in the retinal development were grouped into three clusters by using k-means clustering algorithm. To infer biological functions of each lncRNA cluster, the guilt-by-association approach was used, where (i) Pearson’s correlation coefficient between each lncRNA-mRNA pair was computed in all samples in our dataset; (ii) candidate lncRNA-mRNA pairs were selected as those with correlation coefficients > 0.70 and significance level of 0.05 for Pearson’s correlation (FDR < 0.05) and (iii) protein-coding gene co-expressed with any one lncRNA of each cluster were merged into a union set of genes for GO enrichment analysis.

Validation of uORF-mediated translation repression

To validate translation repression mediated by uORF, Neuro-rod1 was selected to construct reporter vector. The 5’-UTR and CDS sequence of 1242 bp was cloned into pCAG-eGFP, and the main ORF of Neuro-rod1 was Flag-tagged at its C-terminus. The EGFP with an IRES was used as an internal control for transfection and RNA expression. To measure the levels of NEUROD1 protein with a C-terminal FLAG-tag, Neuro-2a cells were transfected with wild-type (WT) or mutant (MUT) reporter plasmids by using Lipofectamine 3000 (Invitrogen). At 24 h post transfection, cells were passaged to 24-well plate and cultured with complete media (10% FBS) or differentiate media (1% FBS) for 2 days. And then cells were harvested, washed in phosphate-buffered saline (PBS) and lysed in RIPA lysis buffer (Invitrogen) added with Benzonase and protease inhibitor. Proteins were separated by sodium dodecylsulphate-polyacrylamide gel electrophoresis (SDS-PAGE), and then transferred to 0.2 μm polyvinylidene fluoride (PVDF) membranes. The membrane was blocked in 3% nonfat dry milk, and then probed with anti-GFP, anti-FLAG and anti-Histone H3.1 primary antibody. The protein bands were developed with HRP-Substrate ECL (Millipore) detected with the Alliance Q9 system.

Translation validation of uORFs and IncORFs

Plasmid construction. We constructed a series of expression vectors for the detection of non-canonical ORF translation, including 4 uORFs (u-Rnf10, u-Rnft1, u-Usp8 and u-Zkscan17) and 10 IncORFs (Brip1os, Cct6a, Gas5, Malat1, Miat, Peg13, RP23-410L16.1, RP24-11214.1, Six3os1 and Zfas1), together with two known translatable lncRNAs (Mrln and Dworf) and one protein-coding gene (DHFR) as positive controls. The uORFs and IncORFs (including the predicted 5’- and 3’-UTR) were cloned from mouse retinal cDNAs; Mrln were cloned from mouse muscle cDNAs; Dworf were cloned from mouse heart cDNAs; and DHFR were cloned from Hela cells cDNAs. After obtaining the sequence of these candidates and controls, a HiBiT tag was inserted upstream the stop codon of each predicted ORF by inverse PCR to enable luminescence-based detection of translation products. In order to confirm that peptides were indeed translated from the corresponding ORFs, mutations were introduced in the start codons of predicted ORFs using ClonExpress II One Step Cloning Kit (Vazyme, #C112-02). Additionally, 10 ORFs (Cct6a, Gas5, Malat1, Mrln, RP23-410L16.1, RP24-11214.1, Six3os1, Zfas1, u-Rnf11 and u-Usp8) and their mutants were added with 3xFlag tags downstream the stop codon of predicted ORFs. The primers and oligos used for cloning, inverse PCR, mutation generation, and epitope tagging were listed in Supplementary Table S2.

Luminescence-based detection of translation products. The translation of ORFs was detected using both in vitro translation assays (IVT assays) and cultured cells. For IVT assays, TiT® Quick Coupled Transcription/Translation System was used for in vitro translation of all HiBiT-tagged ORFs. According to the manufacturer’s instructions, 1 μg of plasmid DNA was used as template for each reaction and the reaction was incubated for 90 min at 30°C. The detection of translated products was performed using Nano-Glo® HiBiT Lytic Detection System (Promega, #N3030) according to the manufacturer’s instructions with minor modifications. In brief, 10 μl IVT products were diluted to 50 μl using nuclease free water, added with 50 μl lytic buffer and mixed by pipet, after 10 minutes incubation at room temperature, the luminescence was measured on a Promega GloMax®-Multi Detection System. To further explore the translational potential of the candidates in cellular context, 500 ng of plasmid DNA was transfected into N2A, Hela and ARPE19 cells using Lipofectamine 3000. Cells were
harvested at 48 h post-transfection for subsequent analysis. The plate was equilibrated to room temperature, then 300 μl lytic buffer was added to each well and incubated for 10 min on orbital plate shaker. The lysates were divided into three tubes and the luminescence of each tube was measured on a Promega GloMax®-Multi Detection System.

**Western blots.** The translated peptides were further validated by western blot. Transfected N2A Cells were harvested with 100 μl of RIPA buffer (25 mM Tris-HCl pH 7.6, 150 mM NaCl, 1% NP-40, 0.1% sodium deoxycholate, 0.1% SDS) added with 1 x PIC (Merck, #539131) and Benzonase (NovoProtein, #M046-01B) and incubated 10 min on ice. For detection of peptides from Brip1, Miap, u-Rnf10 and DHFR, lysates were added with 5 x SDS-PAGE Sample Buffer (GenStar, #E153) and denatured at 95°C for 5 min, and samples were loaded on 4-20% Mini-PROTEAN® TGX™ Precast Protein Gel and transferred to a 0.2 μm NC membrane. For detection of the remaining peptides, lysates were added with 2 x Novex Tricine SDS Sample Buffer (Invitrogen, #LC1676) and denatured at 85°C for 2 min, and samples were loaded on 16.5% GLASS Gel® Tricine gel (WSHT, #TCH2001-16.5T) and transferred to a 0.1 μm NC membrane. The blot was carried out using Nano-Glo® HiBIT Blotting System (Promega, #N4210) according to the manufacturer’s instructions.

**Validation of circRNA expression**

**Divergent PCR validation.** Divergent primers were designed to amplify the circRNA backsplice junction sequence and retinase cDNA were used as template for divergent PCR (Supplementary Table S2). Divergent PCR was performed using green Taq mix (Vazyme), and the reaction was carried out for 3 min at 95°C and 30 cycles of 15 s at 95°C, 15 s at 60°C and 30 s at 72°C. The PCR products were then analysed on 1.5% agarose gels in 1 x TAE buffer. All of the PCR products were sanger sequenced with forward and reverse primers to find the backsplice junction sequence.

**RNase R treatment assay.** RNase R treatment and circRNA quantification was performed according to a published protocol (Panda and Gorospe, 2018) with minor modification. In brief, RNA was treated with 20 μl RNase R digestion reaction (2 μg RNA, 1 μl RNase R (Lucigen, #RNR07250), 2 μl 10× RNase R reaction buffer) and control reaction without RNase R. The reactions were incubated for 30 min at 37°C and immediately purified using ZYMO RNA Clean & Concentrator (ZYMO RESEARCH, D7011). The purified RNA samples were eluted in 20 μl of nuclease free water, and 12 μl of RNase R treated RNA and control RNA were used for reverse transcription. Quantification of circRNAs were performed using iTaq Universal SYBR Green Supermix (BioRad, #1725124) according to the manufacturer’s instructions. The qPCR reactions were prepared as follows: 0.1 μl cDNA, 10 μl of SYBR Green Supermix, 1.2 μl primer mix (5 μM each) and 6.8 μl nuclease-free water. Reactions were carried on CFX Connect Real-Time PCR Detection System for 2 min at 95°C and 40 cycles of 5 s at 95°C and 20 s at 60°C followed by melting curve analysis. The enrichment of RNA after RNase R treatment was calculated using the delta CT method, mouse gene Gapdh and Rps14 were used as linear control.

**RESULTS**

**Ribo-seq, mRNA-seq and circRNA-seq on the mouse retinal tissue**

To understand the translational regulation of gene expression during mouse retinogenesis, we performed Ribo-seq and mRNA-seq to generate translatome and transcriptome profiles of mouse retina at six developmental stages, including E13, E15, P0, P6, P13 and P21, temporally spanning two major developmental events in retina, birth and eye opening (day11-12) (Figure 1A). We also performed circRNA-seq specifically for transcriptomic profiling of circRNAs at E15, P0, P6, P21 and M9. All the sequencing experiments were done with two biological replicates. In total, the Ribo-seq, mRNA-seq, and circRNA-seq yielded 1.17, 0.37 and 0.87 billion raw reads, with an average of around 83.25, 30.98 and 87.38 million reads per library, respectively (Supplementary Table S3).

Ribosome-protected fragments (RPFs) generated from the Ribo-seq had a typical length range of 25- to 35-nucleotide (nt), tightly distributed around a peak of 29- to 30-nt (Figure 1B and Supplementary Figure S1A), a preference mapped to annotated coding sequences (CDS) and 5’ untranslated region (Supplementary Figure S1B), a strong bias toward the translated frame (Figure 1C and Supplementary Figure S1C), and a characteristic three-nucleotide (3-nt) periodic subcodon pattern (Figure 1D and Supplementary Figure S1D). As expected, these characteristics were absent in the mRNA-seq datasets. There was a high degree of agreement between biological replicates, with an average Pearson’s correlation coefficient of 0.981, 0.993 and 0.990 for Ribo-seq, mRNA-seq and circRNA-seq, respectively. Principal component analysis (PCA) showed that the samples had a clear segregation of development stages (Figure 1E) and the samples from the same stage were more similar to each other than the ones from the different stages (Supplementary Figure S1G). In addition, expression of some known marker genes in retinal development on our datasets, such as Otx2, Pax6 and Neurod4, well agreed with previous studies (Supplementary Figure S2) (35–37). Collectively, these results demonstrated that our experimental data were of high quality.

**Coordination of translational and transcriptional regulation during retinal development**

Based on the Ribo-seq and mRNA-seq datasets, we identified an average of 11,150 well-translated protein-coding genes and 12,458 well-transcribed protein-coding genes per stage (Figure 2A, see Materials and Methods). Most of the well-translated and well-transcribed protein-coding genes were shared across all the stages (Figure 2B-C). The shared genes underwent dynamic expression changes during retinal development, with 91.4% (9294 of 10 172 at the transcriptional level; Supplementary Figure S3A) and 76.3% (6961 of 9119 at the translational level; Supplementary Figure S3B) genes showing significant changes in temporal differential
expression (FDR < 0.05; see Materials and Methods), including many well-known transcription factors directing retinogenesis such as Rax, Crx, Otx2, Vsx2 and Neurod1 (Supplementary Figure S3C). Correlation analysis showed that transcriptome and translatome were more different at the early stage but became more similar after eye-opening, suggesting uncoupled changes between transcriptome and translatome during retinal development (Figure 2D).

To capture the overall transcriptional and translational changes in gene expression, we integrated mRNA-seq and Ribo-seq datasets to perform differential expression analysis between adjacent stages using deltaTE (Figure 2E, FDR < 0.05; see Materials and Methods). We observed that the total number of differentially expressed genes (DEGs) gradually increased with development until a peak between P6 and P13, up to a total of 5753, then followed by a dramatic decline. This pattern was consistent with the progression of retinal differentiation and maturation (Supplementary Figure S3D). To capture the specific translational changes independent of changes in transcription, we further identified differential translational efficiency genes (DTEGs) and combined the differential direction of these DEGs and DTEGs to categorize them into four distinct regulatory classes: forwarded, exclusive, buffered, and intensified (Figure 2F and Supplementary Table S4; see Materials and Methods). The forwarded genes have RPF changes that are explained by the mRNA changes. The exclusive genes have changes in TE without mRNA changes. The buffered genes have changes in TE that offsets the mRNA changes and the intensified genes also have changes in TE that amplifies the mRNA changes.

We observed that differences in transcription were not always forwarded to the translational level, and on average, > 38% of differentially transcribed genes were translationally buffered or intensified (Figure 2G). Of these, translational buffering was more prominent than translational intensification, which further emphasized the existence of extensive translational regulation that shaped gene expression changes during retinal development. Moreover, translational regulation could also influence gene expression independently, with an average of 517 differential genes found in each comparison of adjacent stages whose changes occurred exclusively at the translational level without underlying mRNA changes. For instance, between E15 and P6, ribosomal occupancy significantly changed for many well-studied genes related to the functionality of neurons, such as...
as Dhx36 that helps specific microRNA localize to neuronal dendrite (38), Eeal that restores homeostatic synaptic plasticity (39), and Tk2 that ensures neuronal function in vivo (40) (Figure 2H).

Notably, genes in the same regulatory class exhibited developmental stage-specific functional enrichment (Figure 2I), while genes in the different regulatory classes exhibited class-specific functional enrichment (Figure 2J). For instance, 'neuron projection organization' which is critical for the establishment of retinal visual function was particularly enriched for the exclusive genes during P6 to P13 and intensified genes during P13 to P21. Overall, these results revealed that both translational and transcriptional regulation had important but different roles in the development of retina.

Functional importance of translational regulation

We next attempted to examine the relative contribution of translational and transcriptional regulation for the biological functions important to retinogenesis. We first performed GO enrichment analysis for the above-detected differentially transcribed and translated genes (FDR < 0.01, Supplementary Table S5; see Materials and Methods) and iden-
identified specific functions enriched throughout the development of the mouse retina (Figure 3A). Particularly, many of these functions were associated with synapse formation and synaptic transmission at neuromuscular junctions, such as ‘visual system development’, ‘postsynapse assembly’, ‘postsynapse organization’, and ‘signal release from synapse’. We also observed that some biological functions showed stage-specific enrichment. For instance, ‘positive regulation of chromosome separation’ and ‘microtubule-based protein transport’ were exclusively enriched between P0 and P6 as well as between P6 and P13, respectively.

We then calculated the relative percentages of regulatory classes (i.e. forwarded, exclusive, intensified, and buffered) of differential genes for each functional term and performed a PCA to dissect their contributions to the overall expression change of each term. The manifestations showed that these important biological functions were under different degrees of translational and transcriptional regulation (Figure 3B). For instance, the ‘visual system development’ was under combinatorial regulation of transcriptional and translational control mechanisms. At the early stage, this function was subjected to the major transcriptional forwarded regulation, but as development progressed, transcriptional regulation gradually weakened and translational regulation, particularly translationally exclusive and intensified regulation, gradually strengthened (Figure 3C). Signal transmission-related function ‘axon transport’, enriched frequently after birth, was mainly subjected to translational regulation. Specifically, about 62.7% of total contributions in ‘axon transport’ could be attributed to translational regulation (exclusive + intensified + buffered), of which 72.9% came from exclusive and intensified regulation. Besides these, the stage-specific enriched functions that played important roles in the homeostatic maintenance of mature synapses, such as ‘microtubule-based transport’ and ‘postsynaptic modulation of chemical synaptic transmission’, were mainly under translational intensified and exclusive regulation, respectively. Our results showed a rich and complex regulation of gene expression during retinal development.

**Dynamics of translational efficiency and contribution of regulatory uORFs**

Due to long half-lives (>2 h) for the majority of eukaryotic mRNAs, regulation of their encoded proteins is achieved by controlling mRNA TEs and protein degradation rates (41). We detected a total of 5945 differential translational efficiency (DTE) genes between adjacent stages (see Materials and Methods). Unsupervised clustering by k-means revealed the temporal dynamics of TE that were categorized into seven clusters (Figure 4A). Each cluster was composed of distinct genes with specific biological functions (Supplementary Table S6). Particularly, the 1049 DTE genes in the cluster C were significantly enriched in ‘synapse organization’, ‘postsynapse organization’, and ‘neuron projection organization’. TEs of these genes in the cluster C were specifically enhanced around E15 (embryonic wave) and P13 (postnatal wave), which would help promote neuronal differentiation, thereby facilitating production of functionally active neurons. The 925 DTE genes in the cluster E were significantly enriched in ‘mRNA/tRNA 5′-end processing’, ‘midbody abscission’ and ‘microtubule-based transport’. Notably, these genes showed peak TE at P6 and might play a critical role in maintaining retina homeostasis and later neurogenic divisions, given typical characteristics of extensive alternative splicing (42) and active cell division events (43) during this period of retinal development. In addition, the 816 DTE genes in the cluster B showed peak TE at the embryonic period and were mainly enriched in ‘mitotic nuclear division’, ‘negative regulation of chromatin organization’, and ‘positive regulation of chromatin organization’, fulfilling the requirements for early neurogenic divisions.

Previous studies have shown that regulatory elements located on mRNA transcripts might affect gene translation, particularly uORFs in the 5′-UTRs (44–47). In total, we detected 2971 actively translated uORFs in 2123 protein-coding genes (Figure 4B and Supplementary Table S7; see Materials and Methods). We randomly selected four uORFs and successfully validated their translated products by IVT assays (Figure 4C). In further support of uORF translation, we used mass spectrometry (MS)-based proteomics data to provide direct in vivo evidence for translation of 181 uORFs (Supplementary Table S7; see Materials and Methods). Notably, only a minority of uORFs (327) showed strong evidence of conservation (PhastCons score > 0.95; see Materials and Methods). This result indicated the majority of uORFs lacking signs of selective pressure to maintain their amino acid sequences, consistent with the findings of previous study (44), suggesting that uORF function might be largely independent of their translated products.

We next sought to understand the effect of uORFs on CDS translation. By comparing the CDS TEs of genes with and without uORFs, we consistently observed a significant uORF-mediated translational repression (Figure 4D, Wilcoxon rank test). One representative uORF residing in Neurod1 was experimentally validated to further confirm the repressive effect of uORFs on downstream CDS translation (Supplementary Figure S4A). Comparative analysis of the CDS TEs of genes with single versus multiple uORFs further showed multiple uORFs tended to have an additive repressive effect (Supplementary Figure S4B), with similar trends observed in five out of six developmental stages displaying significant non-randomness (Fisher’s exact test, P-value = 0.0313). Given that the majority (57.9%, 1719/2971) of uORFs were found exclusively in a single stage, suggesting that uORF-mediated translational regulation could occur in a stage-specific manner (Figure 4E). The GO enrichment analysis revealed that uORF-containing genes participated in many important biological processes required for retinal development, such as ‘axon guidance’, ‘dopaminergic synapse’, and ‘signaling pathways regulating pluripotency of stem cells’ (Figure 4E and Supplementary Table S8). Moreover, we detected actively translated dORFs in the 3′-UTRs. In addition to thousands of translated uORFs, however, a total of only 266 dORFs were detected in 217 protein-coding genes, with an average of 64 dORFs per stage. Given that translation of dORFs significantly enhance translation of their corre-
Figure 3. Dissecting Translational regulation in the developing mouse retina. (A) Frequency distribution of occurrences of enriched GO terms in all comparisons of adjacent stages. Colours represent overlap degrees of term among the five comparisons of adjacent stages (that is, E15 vs E13, P0 vs E15, P6 vs P0, P13 vs P6, and P21 vs P13). (B) Scatter plot of PCA of each developmental stage separating the manifestations of individual GO terms within the global regulatory programs (see Materials and Methods). Each numbered point represents a functional GO term, and its position along each axis indicates the relative contribution of transcriptional and translational regulation to the overall differential patterns. For each arrangement, the assigned GO term can be found in Supplementary Table S5. (C) Examples of stage-specific coregulatory functional arrangements. The barplots show the relative fractions of expressed genes in retina with distinct regulatory modes.
Figure 4. Analysis of translational efficiencies across retinal development. (A) Hierarchically-clustered heatmap of TE of 5945 protein-coding genes (FDR < 0.05; as described in (27)). Different colours indicate different clusters; numbers indicate significant DTE genes in the cluster; and temporal patterns of TE changes are portrayed through lines. Representative genes and enriched GO terms in the two largest clusters are shown on the right. (B) Number of actively translated uORFs detected in each stage by using Ribo-TISH. The uORF distribution is shown below. (C) Translation validation of uORFs. Left: luminescence of uORFs in IVT assays, where all candidates were compared with the negative control and the dashed line indicates the luminescence of the negative control (negative control: pcHiBiT, positive controls: Dworf and Mrln, protein-coding control DHFR; ns, \( P > 0.05; ^* P < 0.05; ** P < 0.01; *** P < 0.001 \)); Right: western blots of uORF coded peptides (HiBiT tagged) and corresponding ATG mutants. (D) Left: Comparison of TEs between protein-coding genes without, and with and with validated uORFs in each stage; Right: summary of TE between genes without/with with validated uORF (Wilcoxon rank-sum test). (E) Frequency distribution of occurrences of uORFs during retinal development, showing developmental stage-specific usage. Colours represent numbers of uORFs with different usage frequency during the mouse retinal development.
sponding canonical ORFs (48), we compared TEs of genes with and without dORFs, but did not observe significant enhance effect of dORFs on their CDS translation possibly due to their much smaller number per stage (Supplementary Figure S4C).

Translation of long noncoding RNA genes

Micropeptides encoded by presumed long noncoding RNAs (lncRNAs) have frequently been overlooked, and their prevalence and potential function in retinal development, even in neural development, remain unknown. To discover translated lncRNAs in the developing mouse retina, we searched for actively translated ORFs in lncRNAs (lncORFs) (see Methods). In total, we identified 603 unique lncORFs in 290 lncRNAs, with a median length of 48 amino acids (aa) per lncORF (Figure 5A and Supplementary Table S9). Majority of these lncORFs (57.5%, 347/603) could be found in another translatome dataset of mouse retina (GSE94982), including previously well-characterized micropeptides B230154K17Rik (49) and Crude (50). Comparing with MS-based proteomics data, we provided direct in vivo evidence for translation of 75 out of the 290 lncRNAs (Supplementary Figure S5A and Supplementary Table S9; see Materials and Methods).

To further experimentally validate the translation products, we first performed IVT assays for 10 randomly chosen lncORFs. Considering the relatively low molecular weight of potential micropeptides, we specifically fused a 11-amino-acid HiBiT epitope tag to the C-terminal of each lncORF that could produce bright and quantitative luminescence through high affinity complementation with LgBiT (51). The quantifiable results demonstrated that all of them successfully produced micropeptides in IVT assays (Figure 5B). Furthermore, we separately transfected the expression vectors of these lncORFs into N2a (Neuro-2a), Hela, and ARPE19 cells and detected micropeptide products of all of them in at least two cell types (Figure 5C). Subsequent start codon mutation of these lncORFs prevented their translation or caused truncated translation, as evidenced by significant decreases in the luminescence intensity (Figure 5D and Supplementary Figure S5B). Loss of signal in the predicted size range of micropeptide products was additionally confirmed by western blot analysis (Figure 5E and Supplementary Figure S5C).

Translation of lncRNAs showed a strong stage-specificity, with an average of 112 lncRNAs per stage detected to undergo active translation but only 19 shared across all developmental stages (Supplementary Figure S5D). Among the 290 translated lncRNAs, 198 exhibited significant temporal expression changes, of which 12 were found exclusively in translatome, suggesting that lncRNA translation might independently contribute to changes in developmental programs (Supplementary Figure S5E-F). We also sought to understand the tissue-specificity of these 290 translated lncRNAs. We conducted Ribo-seq and mRNA-seq for six tissues in mouse and found 172 (59.3%), 102 (35.2%), 133 (45.9%), 79 (27.2%) and 128 (44.1%) of them that also had translation evidence in the brain, heart, kidney, liver and lung, respectively, with 18 translated in all six tissue types (GSE94982). Importantly, the majority of these lncRNAs (86.1–91.5%) shared at least one identical lncORF with those detected in the retina. These results suggested that some of the translated lncRNAs detected in the developing retina had potential multi-tissue activities. Functional annotation of these micropeptides translated from lncRNAs using InterProScan 5 revealed 68 (~11.3%) of 603 lncORFs with identifiable features found in known proteins (Supplementary Table S1), such as conserved domains, protein families, and functional sites, some of which even could be assigned to defined molecular functions. For instance, micropeptide encoded by Pispmt1 participated in ‘protein tyrosine/serine/threonine phosphatase activity’ and micropeptide encoded by RP23-95L9.6 participated in ‘nucleic acid binding’. Our analysis suggested that these lncORFs had functional relevance which could be tested in the future study.

Translation of circRNAs

An increasing number of studies have indicated that circRNAs can also be translated into detectable peptides with physiological functions (52). We used circRNA-seq and Ribo-seq data to explore translation potentials of circRNAs. Using stringent identification and filtering strategies (Supplementary Figure S6A; see Materials and Methods), 255 of 28 910 transcribed circRNAs with at least two supporting back-spliced footprints were defined as ribosome-associated circRNAs (ribo-circRNAs) (Supplementary Table S10). Notably, the majority (82.0%) of them were found in the stages after eye opening, suggesting their potential functional importance for the later stage of retinal development (Figure 6A).

Ribo-circRNAs had distinguishing properties from untranslated circRNAs (ut-circRNAs). Ribo-circRNAs had significantly shorter exonic length and fewer exons than the ut-circRNAs, with at least 29% and 40% decreases in the median length and number of exons, respectively (Figure 6B; Wilcoxon rank-sum test, P-value < 2.2e–16), in agreement with previous observations in human cell lines (53). Ribo-circRNAs had significantly longer flanking introns (Figure 6C; Wilcoxon rank-sum test, P-value = 4.87e–12) that harboured more repetitive elements than ut-circRNAs (Figure 6D; Wilcoxon rank-sum test, P-value = 2.45e–12). In addition, evolutionary conservation of micropeptides translated from ribo-circRNAs was significantly higher than that of ut-circRNAs (Figure 6E; Wilcoxon rank-sum test, P-value < 1.61e–03), suggesting that they were likely to have functional roles, given that strong conservation in sequence is a general indicator of important biological function (54).

Next, we randomly selected 20 candidates including 19 ribo-circRNAs and 1 ut-circRNA and performed Sanger sequencing of RT-PCR products using divergent primers to confirm the back-splice junction sites of these circRNAs (Figure 6F and Supplementary Figure S6B). After RNase R treatment, all these circRNAs were resistant to digestion with RNase R exonuclease, validating the existence of these ribo-circRNAs (Supplementary Figure S6C). Using MS-based proteomics data, we further provided direct in vivo evidence for translation of 13 ORFs spanning the splice junctions. After minimizing the effect of trans-spliced

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Table 3. Developmental stage-dependent translation of lncORFs

| ORF type | Summary | E13 | E15 | P0 | P6 | P13 | P21 |
|----------|---------|-----|-----|----|----|-----|-----|
| IncRNA gene | Number of IncORFs (translated lnRNAs) | 131 | 173 | 197 | 260 | 356 | 114 |
| IncORFs | Median length (codon) | 70 | 65 | 51 | 52.5 | 54 | 77 |

**Figure 5.** Translation of IncRNAs. (A) Number of actively translated IncORFs detected in each stage. (B) Luminescence of 10 candidate IncORFs and controls in IVT assays. All candidates are compared with the negative control and the dashed line indicates the luminescence of the negative control. (C) Luminescence of 10 candidate IncORFs in N2A, Hela and ARPE19 cells. All candidates are compared with the negative control. (D) Luminescence of all IncORFs and corresponding ATG mutant ORFs in N2A cells. (E) Western blots of IncORF coded peptides (HiBiT tagged) and corresponding ATG mutants. Note: negative control: pcHiBiT, positive controls: Dworf and Mrln, protein-coding control DHFR; ns, $P > 0.05$; * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$.

RNAs (see Materials and Methods), these micropeptides were more likely to derive from ribo-circRNA themselves. Of the 13 ribo-circRNAs producing putative micropeptides, five were found to be differentially transcribed (Figure 6G), including circKmt2a, circArhgap10, circMacf1, circAp3b2 (up-regulated in P21 compared to P6) and circZfp532 (down-regulated in P21 compared to P6), emphasizing that their translation might be important for retinal development.

**DISCUSSION**

Our integrative analysis of transcriptome and translatome during retinal development revealed specific changes in translation and regulatory roles of translational regulation in retinogenesis. We found that retinogenesis is accompanied by dynamic, rapid and coordinated changes in gene translation and translational regulation. Thousands of important regulatory protein-coding genes were subjected to significant changes in the translational level in a stage-
specific manner, potentially redefining functional architecture and diversity of the retinal cells.

Specific translational regulation could be achieved by controlling TEs, thereby determining quantitative differences in protein abundance during retinogenesis. TE dynamics during developmental transition could further be triggered by some regulatory elements, such as the most significant at certain stages. Moreover, many uORF-encoded regulatory mechanisms may contribute to an uORF's role in translational regulation, including the length of the 5′UTR, the secondary structure and GC content, as well as the strength of the surrounding Kozak context, and the uORF length, which have been substantially discussed in detail in previous studies (46,56,57). A uORF with a suboptimal context may benefit leaky scanning and hence allow its main CDS translation (58). Due to these complexities of uORF-mediated translational regulation, this might partially explain why the change of translation efficiency by uORF is not significant at certain stages. Moreover, many uORF-encoded
peptides were here detected. Recent studies have suggested that they may modulate translation or have other cellular functions. For instance, uORF-encoded micropeptides act in cis to cause ribosome stalling or limit ribosomal access to the main CDS or act in trans to suppress translation (59). uORF-encoded peptides can serve as MHC class I ligands, thereby contributing to the antigen repertoire and possible immunogenicity (60). Detailed physiological functions and underlying mechanisms of uORFs in shaping retinogenesis via producing micropeptides remain to be elucidated.

Surprisingly, we additionally discovered pervasive translation of IncORFs from IncRNAs. We experimentally confirmed their translation products and showed that these micropeptides had regulatory potential and biological relevance. For instance, we found that the micropeptide encoded by RP23-15O6.3 contained C2H2-type zinc fingers. The micropeptide encoded by AC166710.3 shared a common structure with ribosomal L18e/L15P superfamily. Functional enrichment analysis revealed potential roles of these translated IncRNAs in retinal development (Supplementary Figure S5F). One representative IncRNA was Miat (also known as Rner2 or Gomafu) that contributed to mitosis of retina progenitor cells, illustrating its essential role for retinogenesis (61). Besides the direct involvement of retinal development, some of the encoded micropeptides may serve as regulatory elements to play important roles in modulating neighbouring gene activity. This is partially supported by the observation that neighbouring genes of many translated IncRNAs such as Six3os1 (IVT assays and in vivo translation), Vax2os, Otx2os1, Pax6os1, and Zeb2os1 (MS detection) were important determinants of retinal cell fate (62). In our study, micropeptides were only evident for a relatively small subset of uORFs, dORFs, and IncORFs. The possible reasons behind this were: (i) the majority of their encoded peptides might lack signs of selective pressure to maintain their amino acid sequence; (ii) their encoded peptides may be relatively short lived and rapidly degraded; (iii) their encoded peptides might exist at only very low levels within the cell and (iv) our high-pH reversed-phase fractionated retinal proteomics data did not cover the full peptide isoelectric point range so that it was unable to explore the entire tryptic peptidome. To obtain a comprehensive map of hidden micropeptides during retinal development, innovative technologies are needed.

Notably, 13 micropeptides encoded by non-canonical ORFs spanning the splice junctions were detected using MS-based proteomics data. These micropeptides might be partially associated with the functions of their host proteins due to the existence of overlapping sequences between them, likely competing with (63) or protecting (64) their host proteins. It should be noted here that circRNAs possess coding potential and can indeed be translated, but translation may originate from their linear trans-spliced RNA by-products (tsRNAs) (65). Although we take advantage of our poly(A)+ mRNA-seq data to minimize the impact of potential tsRNAs, additional experiments to assess whether these micropeptides are translation products of circRNA themselves remain necessary particularly when concerning with the mechanistic aspect of their synthesis.

In addition, some limitations might exist in our current study. In particular, a high-resolution view of cellular translational heterogeneity and cell-type-specific translational regulation was obscured by our bulk-seq data. Single-cell sequencing can be used to analyse the differences in gene expression between individual cells, but regrettably, technologies enable genome-wide investigation of in vivo gene translation with subcodon resolution at the single-cell level remain unavailable to date. Nevertheless, our bulk-seq data measure the average behaviour of gene expression from heterogeneous tissue samples, this tends to reduce the sparsity of values within the expression matrix, making the data’s parameters more rich and less susceptible to dropouts. Our dataset can serve as a valuable resource for future studies of the translational machinery during retinogenesis, providing crucial and complementary information to single-cell omics data. In general, our present study provides an overall snapshot of gene translation and translational regulation during retinal development.

DATA AVAILABILITY

All Ribo-Seq, RNA-Seq, and circRNA-seq sequencing data have been deposited in the SRA database at NCBI (https://www.ncbi.nlm.nih.gov/sra/) under accession number PRJNA589677. The mass spectrometry proteomics data have been deposited in the PRIDE database under accession number PXD023439.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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Author contributions: Z.X. and H.W.W. supervised the project; J.Q.Y. performed RNA-seq and Ribo-seq library construction; C.Y.C. designed and performed experiments; M.Q.X. provided help and suggestions for experimental design; K.N.C. and H.H.L. analysed and interpreted the data; K.N.C., C.Y.C., H.W.W. and Z.X. wrote the manuscript; all authors approved the manuscript.

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