



Express: A database of transcriptome profiles encompassing known and novel transcripts across multiple development stages in eye tissues

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ABSTRACT

Advances in sequencing have facilitated nucleotide-resolution genome-wide transcriptomic profiles across multiple mouse eye tissues. However, these RNA sequencing (RNA-seq) based eye developmental transcriptomes are not organized for easy public access, making any further analysis challenging. Here, we present a new database “Express” (<http://www.iupui.edu/~sysbio/express/>) that unifies various mouse lens and retina RNA-seq data and provides user-friendly visualization of the transcriptome to facilitate gene discovery in the eye. We obtained RNA-seq data encompassing 7 developmental stages of lens in addition to that on isolated lens epithelial and fibers, as well as on 11 developmental stages of retina/isolated retinal rod photoreceptor cells from publicly available wild-type mouse datasets. These datasets were pre-processed, aligned, quantified and normalized for expression levels of known and novel transcripts using a unified expression quantification framework. Express provides heatmap and browser view allowing easy navigation of the genomic organization of transcripts or gene loci. Further, it allows users to search candidate genes and export both the visualizations and the embedded data to facilitate downstream analysis. We identified total of > 81,000 transcripts in the lens and > 178,000 transcripts in the retina across all the included developmental stages. This analysis revealed that a significant number of the retina-expressed transcripts are novel. Expression of several transcripts in the lens and retina across multiple developmental stages was independently validated by RT-qPCR for established genes such as *Pax6* and *Lhx2* as well as for new candidates such as *Elavl4*, *Rbm5*, *Pabpc1*, *Tia1* and *Tubb2b*. Thus, Express serves as an effective portal for analyzing pruned RNA-seq expression datasets presently collected for the lens and retina. It will allow a wild-type context for the detailed analysis of targeted gene-knockout mouse ocular defect models and facilitate the prioritization of candidate genes from Exome-seq data of eye disease patients.

1. Introduction

The eye is a complex sensory organ that consists of an anterior segment that comprises of the cornea, iris, lens, ciliary body and anterior sclera, and a posterior segment that comprises of the retina, choroid and the optic nerve. Eye development is coordinated by a complex regulatory program that involves a myriad of signaling, transcriptional and post-transcriptional events (Lachke and Maas, 2010; Cvekl and Ashery-Padan, 2014; Zagozewski et al., 2014; Dash et al., 2016). With the advancement of sequencing technologies (such as Next Generation Sequencing (NGS)) and its broad application on a genome

wide scale (ENCODE Project Consortium, 2004; Chin et al., 2011; GTEx Consortium, 2015), it is possible to explore the mechanisms governing the developmental “oculome” (Lachke and Maas, 2010). Indeed, over the past decade, several studies have reported on the transcriptome of specific eye tissues at various development stages (Lachke et al., 2012; Khan et al., 2015; Tian et al., 2015; Chaitankar et al., 2016; Anand and Lachke, 2017; Kakrana et al., 2017).

RNA sequencing (RNA-seq) provides a high-resolution comprehensive platform to define cell or tissue-specific transcriptomes and monitor changes therein (Kang et al., 2015). Further, it has advanced our knowledge and understanding of the structure and composition of

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protein-coding and non-coding transcript isoforms in higher eukaryotes, in turn facilitating the downstream functional and comparative analysis (Chen et al., 2014; Calahorra et al., 2015; Fernandez-Valverde et al., 2015; Zimmermann et al., 2015). Transcriptome studies reported for various developmental stages on the lens (Hoang et al., 2014; Khan et al., 2015, 2016) and retina (Busskamp et al., 2014; Roger et al., 2014; Sundermeier et al., 2014; Uren et al., 2014; Andzelm et al., 2015; Ruzyski et al., 2015; Zhang et al., 2015) were mostly limited to comparative gene expression analysis, by restricting to known or annotated genes. However, the complete transcriptome and various isoforms in the context of developmental stages in tissues of eye are not fully characterized. In this study, we present a comprehensive and user-friendly platform termed “Express”, which enables the investigation of the transcriptomic profiles in mouse lens and retina tissues across various developmental stages. Express provides a one-stop portal for investigating gene expression at the resolution of individual transcripts encoded by not just the annotated coding and non-coding genes, but importantly also many novel gene loci in the mouse genome. Express facilitates this by allowing users to view the transcript level expression profiles of a gene across multiple developmental stages as heatmaps and simultaneously enables the visualization of the genomic location of the transcripts in an embedded genome browser. Users can view and download the various visualizations as well as the underlying data to facilitate rational design of experiments to study transcript structure, expression and splicing alterations across different developmental stages.

2. Materials and methods

To obtain a comprehensive understanding of the transcriptome during development in lens and retinal tissues in mouse eye, we collected multiple publicly available RNA-seq datasets corresponding to the raw RNA sequence reads of mouse eye subcomponents from different developmental stages (Table 1 and Table 2). Briefly, these datasets were aligned to the mouse reference genome, quantified for expression levels of known and novel transcripts followed by the normalization of the expression levels across samples. Resulting raw and normalized expression levels were then organized into an open-source Relational DataBase Management System (RDBMS) MySQL database as illustrated in the workflow (Fig. 1). PHP: Hypertext Preprocessor (PHP) backend Application Program Interface (API) helps query the database and a user-friendly

frontend enables the visualization of the query results as heatmap and browser views across development stages. Each of the major steps employed in processing and analysis were described in further detail in the following sections.

2.1. Data collection and processing

We collected the raw RNA-seq reads of multiple development stages (each with its biological replicate) of mouse eye from Gene Expression Omnibus (GEO) (Barrett et al., 2013) and European Nucleotide Archive (ENA) (Gibson et al., 2016). Tables 1 and 2 show the relevant source of the RNA-seq datasets along with several metrics for lens and retina respectively, resulting from the alignment of the reads to the mouse reference genome (mm10). We downloaded the single end datasets in FASTQ format (A text-based format for storing both the nucleotide sequence and its corresponding quality scores) using the Sequence Read Archive (SRA) Toolkit (fastq-dump command), and the paired end datasets were directly downloaded from ENA (European Nucleotide Archive). We ensured the quality of the aligned sequence reads was a minimum of Phred quality score 20 for each sample using FASTX-Toolkit (http://hannonlab.cshl.edu/fastx_toolkit/).

We employed Hierarchical Indexing for Spliced Alignment of Transcripts (HISAT, version 0.1.6) for aligning short reads from RNA-seq experiments onto reference genome (Kim et al., 2015). HISAT (with default parameters) can rapidly align the quality filtered reads collected from different sources (Tables 1 and 2) against the mouse reference genome mm10. SAM (Sequence Alignment/Map) files obtained as outputs from HISAT were post-processed using SAMtools (version 0.1.19) (Wheeler et al., 2008; Li et al., 2009) for converting SAM to BAM (Binary Alignment/Map) followed by sorting the output BAM files, and finally these BAM files were indexed using SAMtools. The sorted BAM files obtained after post-processing were used to quantify the expression levels of known and novel transcripts across development stages.

Transcript quantification and discovery from the aligned RNA-seq datasets was accomplished using StringTie (version 1.2.1) (Pertea et al., 2015). StringTie is a novel network flow algorithm based on a fast and highly efficient assembler to quantify the transcripts of each genomic locus considering all possible multiple splice events. In addition to annotated transcripts, it can also provide the information of possible novel transcripts in each sample. The transcript level expression data for each sample quantified using StringTie were stored as GTFs (Gene

Table 1

RNA-seq samples for mouse lens. The table shows SRA ID (Sequence Read Archive) for the sample, PMID (PubMed) for the study, developmental stage, read type, read length, read count, base count, and overall alignment rate using HISAT (Hierarchical Indexing for Spliced Alignment of Transcripts).

#	SRA ID	PMID	Development Stage	Read Type	Read Length	Read Count	Base Count	Overall Alignment Rate (%)
1	SRR2039769	26225632	E15	PE	100	13772390	2754478000	94
2	SRR2039770	26225632	E15	PE	100	13542500	2708500000	95
3	SRR953395	24161570	E15.5	SE	52	48552190	2524713880	94
4	SRR953394	24161570	E15.5	SE	52	47574424	2473870048	94
5	SRR953393	24161570	E15.5	SE	52	42525381	2211319812	94
6	SRR2039771	26225632	E18	PE	100	17810970	3562194000	93
7	SRR2039772	26225632	E18	PE	100	18019388	3603877600	93
8	SRR1222595	25489224	P0	SE	51	33174286	1691888586	88
9	SRR1222596	25489224	P0	SE	51	29919226	1525880526	87
10	SRR1222672	25489224	P0	SE	51	29965660	1528248660	89
11	SRR1222673	25489224	P0	SE	51	28652759	1461290709	86
12	SRR1222674	25489224	P0	SE	51	30661663	1563744813	89
13	SRR1222675	25489224	P0	SE	51	24833352	1266500952	89
14	SRR2039773	26225632	P0	PE	100	17766309	3553261800	93
15	SRR2039774	26225632	P0	PE	100	14533000	2906600000	93
16	SRR2039775	26225632	P3	PE	100	15495833	3099166600	93
17	SRR2039776	26225632	P3	PE	100	13072393	2614478600	93
18	SRR2039777	26225632	P6	PE	100	16965754	3393150800	93
19	SRR2039778	26225632	P6	PE	100	17658286	3531657200	93
20	SRR2039779	26225632	P9	PE	100	18874309	3774861800	93
21	SRR2039780	26225632	P9	PE	100	13563853	2712770600	93

Table 2

RNA-seq samples for mouse retina. The table shows SRA ID (Sequence Read Archive) for the sample, PMID (PubMed) for the study, developmental stage, read type, read length, read count, base count, and overall alignment rate using HISAT (Hierarchical Indexing for Spliced Alignment of Transcripts).

No	SRA ID	PMID	Development Stage	Read Type	Read Length	Read Count	Base Count	Overall Alignment Rate (%)
1	SRR1023063	24382353	P2	SE	76	29939891	2275431716	93
2	SRR1023064	24382353	P2	SE	76	36280541	2757321116	94
3	SRR1784052	26324254	P10	SE	50	27028041	1351402050	94
4	SRR1784053	26324254	P10	SE	50	24267068	1213353400	94
5	SRR1784054	26324254	P10	SE	50	28777255	1438862750	94
6	SRR1574329	25801704	P11	PE	90	108113500	19460430000	96
7	SRR1574330	25801704	P11	PE	90	106003809	19080685620	97
8	SRR1574333	25801704	P11	SE	90	176162120	8631943880	87
9	SRR1574334	25801704	P11	SE	90	176206610	8634123890	87
10	SRR1023073	24382353	P21	SE	76	39574659	3007674084	94
11	SRR1023074	24382353	P21	SE	76	38855951	2953052276	94
12	SRR1784070	26324254	P21	SE	50	34792825	1739641250	95
13	SRR1784071	26324254	P21	SE	50	31461693	1573084650	95
14	SRR1784072	26324254	P21	SE	50	37264811	1863240550	95
15	SRR1176996	24812086	P28	SE	50	45056397	2252819850	92
16	SRR1176997	24812086	P28	SE	50	51508183	2575409150	91
17	SRR1176998	24812086	P28	SE	50	52339450	2616972500	92
18	SRR1687694	25712131	P30	PE	100	16507045	3334423090	84
19	SRR1687695	25712131	P30	PE	100	16384187	3309605774	84
20	SRR1687696	25712131	P30	PE	100	14348324	2898361448	81
21	SRR1687697	25712131	P30	PE	100	14251738	2878851076	81
22	SRR1687698	25712131	P30	PE	100	25674252	5186198904	83
23	SRR1427139	25002228	P30	SE	51	44636149	2276443599	68
24	SRR1427140	25002228	P30	SE	51	40396217	2060207067	66
25	SRR1427141	25002228	P40	SE	51	52430453	2673953103	74
26	SRR1427142	25002228	P40	SE	51	46344921	2363590971	75
27	SRR1213798	25489233	P48	PE	90	6779886	1220379480	96
28	SRR1213799	25489233	P48	PE	90	6803332	1224599760	97
29	SRR1213800	25489233	P48	PE	90	6826280	1228730400	96
30	SRR1427143	25002228	P50	SE	51	42904725	2188140975	70
31	SRR1427144	25002228	P50	SE	51	41243569	2103422019	68
32	SRR1427145	25002228	P60	SE	51	56910039	2902411989	48
33	SRR1427146	25002228	P60	SE	51	46737505	2383612755	56
34	SRR1427147	25002228	P90	SE	51	40366735	2058703485	76
35	SRR1427148	25002228	P90	SE	51	45096977	2299945827	71

Transfer Format files) providing expression levels for both known and novel transcripts. All the GTFs obtained for each sample were grouped and provided as an input for StringTie “merge” mode along with mouse reference genome (mm10) to obtain a reference annotation file (in GTF) including novel transcripts. Next, the reference merged GTF was used in re-running StringTie with the sorted BAM files for the corresponding samples, to obtain GTFs per sample having the same transcript identifier for a given novel transcript across all the samples. The known transcripts were defined as the transcripts that were annotated as reference mouse transcripts in the Ensembl database (Yates et al., 2016). In contrast, novel transcripts were defined as the transcripts that were exclusively predicted by StringTie with little or no overlap with existing mouse transcript annotations in mm10. We examined the length of the discovered transcript onto annotated reference transcript coordinates and calculated a novelty score for each novel transcript by using the below formula,

$$\text{Novelty Score} = \left(1 - \frac{\text{length of overlapping region}}{\text{full length of novel transcript}} \right) \times 100$$

The novel transcripts having a novelty score (NS) ≥ 70 were considered as completely novel and the novel transcripts having novelty score < 70 were considered as unannotated transcripts. Since sequencing or processing artifacts at various steps of the transcript quantification analysis could potentially contribute to high number of transcript isoforms, we have classified transcripts into three categories namely a) known transcripts which are annotated in Ensembl database (https://www.ensembl.org/Mus_musculus/Info/Index) b) completely novel transcripts i.e., transcripts which exhibit a novelty score of at least 70 and c) the remaining transcripts were classified as unannotated

transcripts and excluded from all the downstream analysis. A quantification matrix was generated for both the lens and retinal transcriptomes with respect to different developmental stages by extracting the TPM (Transcripts Per Million reads sequenced) values from StringTie outputs. These matrices were utilized for downstream analysis such as normalization of expression levels and the corresponding datasets (raw and normalized expression levels) were employed to show the transcript levels across stages in lens and retina in *Express*.

2.2. Normalization of transcript expression levels across samples in a tissue

Although RNA-seq samples originating from the same laboratory are unlikely to have significant technical variation among the replicates and developmental stages, there could still be variations arising due to factors like tissue preparation, RNA extraction and sequencing depth differences. In our analyzed datasets for both lens and retina, RNA sequencing datasets originating from multiple labs and protocols were analyzed. Hence, in addition to providing the default option of raw expression levels of a transcript across developmental stages, we have performed a widely adopted quantile normalization of the samples using the preprocessCore package (Bolstad, 2001) in R and used the resulting normalized expression data for showing the expression heatmaps in *Express*. Quantile normalization is a global adjustment method that assumes the statistical distribution of each sample under study is the same (Bolstad et al., 2003). Normalization is achieved by forcing the observed distributions to be the same and the average distribution, obtained by taking the average of each quantile across samples, is used as the reference. Its application on both microarray and RNA sequencing data has consistently shown its superior performance compared to

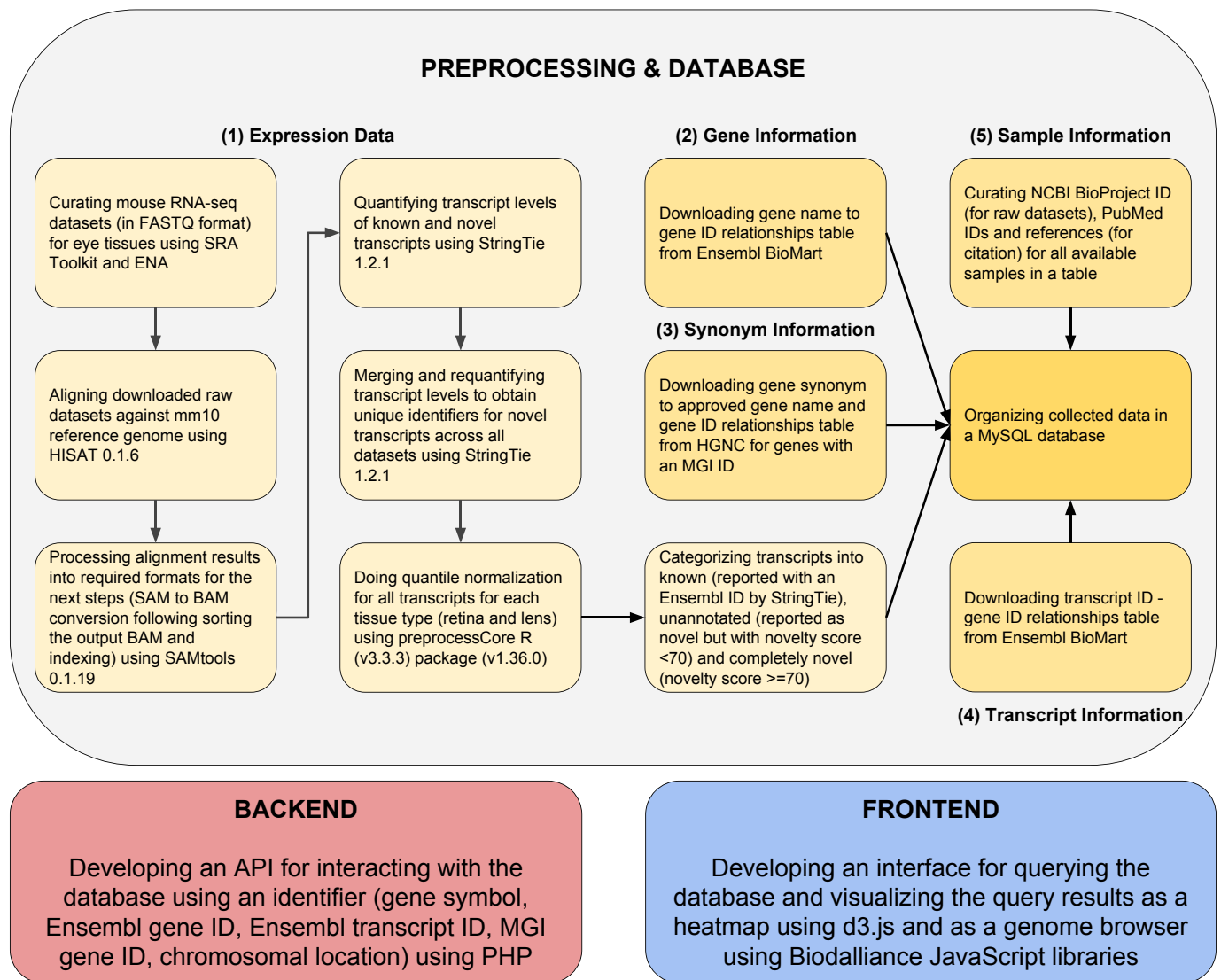


Fig. 1. Overview of the transcriptome profiling and database construction for *Express*. Transcriptomes of mouse lens and retina spanning several developmental stages (with biological replicates) were collected from published sources listed in Tables 1 and 2. Curated RNA sequence data was quality filtered using FASTX Toolkit. High quality raw sequence reads were processed and aligned to mouse reference genome mm10 using HISAT and outputs were collected as Sequence Alignment Map (SAM) files. Post-processing (i.e. conversion of SAM to sorted Binary Alignment Map (BAM)) of aligned reads was accomplished using SAMTools. Aligned and post-processed RNA-seq BAM files associated with each developmental stage were utilized for identifying and quantifying the expression levels of known and novel transcripts across respective developmental stages of tissue subtypes using StringTie. Quantile normalization was performed for samples per tissue type using preprocessCore R package. The novel transcripts reported by StringTie were categorized into unannotated (novelty score < 70) and completely novel transcripts (novelty score ≥ 70). These normalized expression levels of known, unannotated and completely novel transcripts were organized into a table. Gene information mapping gene names to gene IDs was downloaded from Ensembl BioMart. Synonym information mapping gene synonyms to approved gene names and gene IDs was downloaded from Hugo Gene Nomenclature Committee (HGNC) for the genes with an MGI ID. Sample information was manually curated for samples and NCBI BioProject ID, PubMed ID and study reference were obtained per sample. These collected data were then organized into a My Structured Query Language (MySQL) database. Following abbreviations and web resources have been employed in this study: SRA - Sequence Read Archive (<https://www.ncbi.nlm.nih.gov/sra>), ENA - European Nucleotide Archive (<https://www.ebi.ac.uk/ena>), HISAT - <https://ccb.jhu.edu/software/hisat/>, SAM - Sequence Alignment Map, BAM - Binary Alignment Map, StringTie - <https://ccb.jhu.edu/software/stringtie/>, R - <https://www.r-project.org/about.html>, Ensembl Biomart - <https://www.ensembl.org/biomart>, HGNC - HUGO Gene Nomenclature Committee, MGI - Mouse Genome Informatics (www.informatics.jax.org), MySQL - My Structured Query Language, API - Application Programming Interface, PHP - Hypertext Preprocessor.

other competing methods (Bolstad et al., 2003; Dillies et al., 2013). Hence, we used this normalization on our RNA-seq expression profile matrices across developmental stages in lens and retina respectively. Raw or normalized expression levels of replicates of a developmental stage were averaged for display purposes on *Express*. In addition to the quantile normalization, to exhibit only high quality relevant transcripts, the end user has the option to select only highly expressed transcripts for visualization. This is facilitated by including a selection filter which allows the visualization of the expression levels for only those transcripts of a gene which have at least a certain level of expression observed in at least one of the developmental stages shown.

2.3. Database construction and implementation

In order to build the *Express* database of transcriptome profiles encompassing known and novel transcripts across multiple developmental stages in eye tissues in mouse, we employed several steps. These steps are illustrated in Fig. 1. Briefly, as described in the above sections, the aligned, quantified and then normalized datasets organized as final matrices for each tissue type were stored into an SQL table. *Express* stores both the raw as well as the quantile normalized expression levels of transcripts in Transcripts Per Million Reads (TPM) units. Moreover, the sample metadata information is manually curated with NCBI

Table 3
Primers for qPCR validation of transcripts.

Gene	Forward primer sequence (5' -> 3')	Reverse primer sequence (5' -> 3')
Pax6	AGTTCTTCGCAACCTGGCTA	ACTTGGACGGGAAGTACAC
Elavl4	GGCAGAAGAAGCCATCAAAG	GCAAATGTGTCAGCCTGAAT
Rbm5	GCACCACAGTGACTACCACT	GCACGTAGGTCTCCTTCTCG
Lhx2	CGCGCTTAGCTGTAACGAGAA	CGCTTTGTCTTTTGGCTGCT
Pabpc1	GAGACCAGCTTCTCACAGG	ACCTTGACATGAACAGCAG
Tia1	GGCTTGGTGGAAGACAAATC	TCACCCCTCCACAGTACACA
Tubb2b	ATCGGTGCCAAGATCGGT	CCTGAAGATCTGCCCAAATG

BioProject ID, PubMed ID and a reference for citing the corresponding dataset. Also, the table containing Ensembl gene ID, MGI (Mouse Genome Informatics) gene ID and chromosomal location for all genes in mouse genome was downloaded from Ensembl BioMart and the table containing gene synonym, approved gene name and Ensembl gene ID was downloaded from HGNC (HUGO Gene Nomenclature Committee) for genes that are linked to an MGI gene ID. Similarly, we obtained transcript ID - gene ID relationships table from Ensembl BioMart for linking gene information to the expression data. We then organized these tables into a MySQL database whose schema is shown in Fig. S1.

2.4. User interface and access

2.4.1. Backend

We developed a PHP: Hypertext Preprocessor (PHP) Application Programming Interface (API) for interacting with the database using a query (e.g. gene symbol, Ensembl gene ID, MGI gene ID, Ensembl transcript ID or chromosomal location) for the user given TPM cutoff and tissue type. Upon sending the query, TPM cutoff and tissue type to the API, the query type is identified, and the corresponding quantile normalized transcript level expression data is retrieved from the database. Next, the expression values are normalized between 0 and 1 per transcript and the final data is returned in JSON (JavaScript Object Notation) format to be visualized by the frontend. The backend PHP API can also be used programmatically to obtain expression data, which is documented on documentation page of *Express* (<http://www.iupui.edu/~sysbio/express/docs.html>).

2.4.2. Frontend

The frontend interacts with the user to accept input (a tissue type, a TPM cutoff, value type and a query) and a visualization of the retrieved data from the MySQL database is provided. We show the structure of queried transcripts in a genome browser developed using Biodalliance JavaScript library (<http://www.biodalliance.org>). We obtained mouse transcript structures from GENCODE version M7 (GRCm38.p4) in BigBed format (A binary file format that is created by conversion from a Browser Extensible Data format file) available on http://www.gencodegenes.org/mouse_biodalliance.html. To this BigBed file, we added the structures of novel transcripts discovered by StringTie from our analysis. We renamed the default identifiers obtained from StringTie to include corresponding tissue type in the identifier for easy understanding in the genome browser. To modify the GENCODE transcript annotation, we first converted the BigBed file into BED file, added the structures of novel transcripts and then converted back to BigBed format using UCSC utilities (Kent et al., 2010). Also, the expression data per transcript across multiple developmental stages is shown as a heatmap developed by using d3.js JavaScript library (<https://d3js.org>). The heatmap is sorted by transcript groups (as introduced in the section “Data collection and preprocessing”) as known transcripts, completely novel transcripts and unannotated transcripts. The transcripts in each group are also sorted by the averaged expression value for all developmental stages for keeping highly expressed transcripts at the top. The front end provides two select boxes for choosing an available TPM

cutoff (0, 1, 2, and 5), and the tissue type and a textbox for entering the query. The frontend interface allows the user to choose a minimum expression cutoff for a transcript, which enables the display of only those transcripts resulting from search exhibiting this minimum expression level cutoff in at least one developmental stage. The default cutoff is set to 5 TPM. The value type select box can also be used to query for raw expression values or quantile normalized expression values. After search is performed, the results are shown as a heatmap along with a genome browser to view the transcript structure. The heatmap and browser view can be toggled using the button on the right-hand side of the navigation bar. Also, using the Export dropdown menu, it is possible to export heatmap view and browser view in SVG (Scalable Vector Graphics) format and heatmap data in TSV (Tab Separated Values).

2.5. Experimental validation of the RNA-seq identified transcripts for lens and retinal expressed genes

The University of Delaware animal facility hosted all the mice used in these experiments, which were performed following the guidelines defined in the Association for Research in Vision and Ophthalmology (ARVO) statement for the use of animals in ophthalmic and vision research. C57Bl/6 mouse lenses were microdissected at three stages, namely, embryonic day (E) 15.5, post-natal day (P) 0 and P10. Retina was dissected from four stages, namely P10, P20, P30 and P48. The day of detection of vaginal plug was defined as E0.5. Each of three biological replicates at E15.5 comprised of six lenses, and at P0 and P10 comprised of two lenses. Each of the biological replicates for retinal expression comprised of 2 retinas from P10, P20, P30 and P48. Total RNA was extracted from lenses using RNeasy Mini kit (Qiagen Inc, Valencia, CA) and cDNA was synthesized using Bio-Rad iScript™ cDNA Synthesis Kit (Bio-Rad Laboratories, Hercules, CA), for use as a template in quantitative PCR (RT-qPCR) analysis. Forward and reverse primers were designed on the longest isoform of the transcript on exonic sequence flanking an intronic region such that the product sizes were < 300 bp (Table 3). RT-qPCR was performed using Power SYBR Green PCR Master Mix (Invitrogen life technology, Grand Island, NY). Several house-keeping genes namely, *Actb*, *B2m* (Beta 2-microglobulin), and *Hprt* were used for normalization (Wigle et al., 1999; Shaham et al., 2013; Cavalheiro et al., 2014; Mamuya et al., 2014; He et al., 2016). Fold-change differences between target gene expression compared to specific housekeeping gene expression was estimated using the $\Delta\Delta Ct$ method. The first comparison of gene expression in the $\Delta\Delta Ct$ method was performed independently with several housekeeping genes. The second comparison was calculated based on expression at E15.5 (lens samples), and at P10 (retina samples). Statistical significance was calculated using two-way ANOVA as described (Bookout and Mangelsdorf, 2003).

3. Results and discussion

3.1. Overview of *Express* database

Express is a database of transcriptome profiles encompassing known and novel transcripts across multiple development stages in mouse eye tissues. Several steps involved in preprocessing, post-processing, quantification and normalization of collected data followed by its organization in *Express* are illustrated in Fig. 1 (see Materials and Methods). *Express* contains 81,779 distinct transcripts for mouse lens and 178,367 distinct transcripts for mouse retinal samples. Novel transcripts are defined as those that are not annotated in the reference genome annotation (see Materials and Methods). The proportions of the known and completely novel transcripts for each developmental stage at 5 TPM threshold in lens and retina are shown in Fig. 2A and B, respectively (Fig. S2 shows the distributions for different TPM thresholds). In the following sections, we illustrate the composition of the

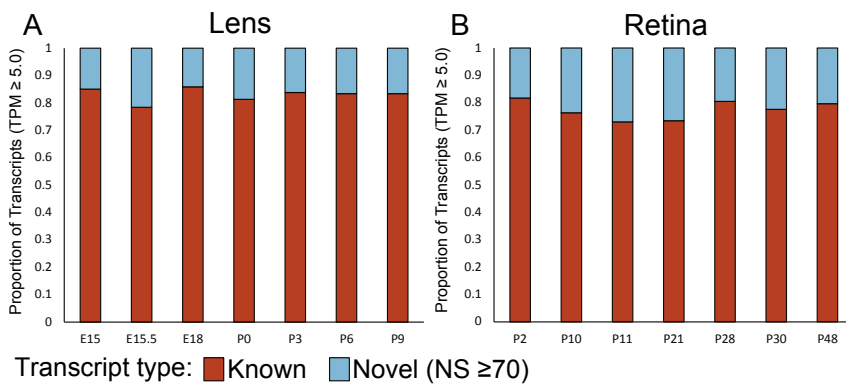


Fig. 2. Histograms showing the proportion of known and completely novel transcripts documented in Express across developmental stages at 5 Transcripts Per Million (TPM) mapped reads threshold. (A) Proportion of transcripts for each stage available for lens samples from E15 to P9. (B) Proportion of transcripts for each stage available for retinal samples from P2 to P90. Multiple datasets associated with a given developmental stage are merged to facilitate the ease of comparison across stages.

datasets and functionality of the database as well as present validations of several genes in lens and retinal tissues, to demonstrate the utility of *Express* for studying eye development.

3.2. Analysis of lens and retinal RNA-seq data for building *Express*

Express contains gene and transcript level expression data obtained from 21 lens and 35 retinal RNA-seq mouse samples. As shown in Table 1 for lens samples and in Table 2 for retinal samples, the information about the datasets used in this study is given per dataset as Sequence Read Archive (SRA) ID, PubMed (PM) ID, developmental stage, read type (single-end or paired-end), read length (in bp), read count, base count and overall alignment rate. The lens samples include developmental stages from E15 to P9 whose read types and read lengths vary with alignment rates ranging from 86% to 94%. The retinal samples include developmental stages from P2 to P90 whose read types and read lengths also vary. Majority of the retinal samples exhibited a high overall alignment rate varying from 80% to 97% (See Materials and Methods, Tables 1 and 2). We downloaded the raw datasets for lens and retinal samples from Gene Expression Omnibus (GEO) (Barrett et al., 2013) and European Nucleotide Archive (ENA) (Gibson et al., 2016). We performed a quality control using FASTX-Toolkit. Reads with a Phred quality score lower than 20 were filtered out. Next, we aligned them to the mouse reference genome (mm10) using Hierarchical Indexing for Spliced Alignment of Transcripts (HISAT) and quantified the aligned datasets for transcript expression levels for known and novel transcripts identified by StringTie, which is an established method for identifying new transcripts (See Materials and Methods, Fig. 1). The gene and transcript information from mouse reference genome were also downloaded from Ensembl (Yates et al., 2016) and organized along with the expression data in a MySQL database (See Materials and Methods, Fig. 1). To control for technical variation in expression levels between samples, we performed a quantile normalization of all the samples in a given tissue type (See Materials and Methods). Both raw as well as normalized expression levels in Transcripts Per Million (TPM) reads sequenced units, are stored in the database and are available to download from the *Express* website. We also developed a PHP: Hypertext Preprocessor (PHP) backend to interact with the database and a frontend to interact with the user and to visualize the query results (See Materials and Methods, Fig. 1).

3.3. User guide for exploring *Express* database

To retrieve gene expression data from *Express*, we have added the following features to the web interface. Step-by-step instructions for using *Express* are also available as a User Manual (see Fig. 3A and the web interface of *Express* following the webpage <http://www.iupui.edu/~sysbio/express/user-guide.html>).

- (1) The parameters to investigate the expression of a gene are (a) tissue

type, namely lens, retina and lens cell subtype; (b) expression level, namely gene or transcript (splice isoform) level; (c) TPM (transcripts per million) cutoff of 0, 1, 2 and 5 and tpm values which could be raw or values after quantile normalization.

- (2) The query to investigate gene expression could be gene name, ENSEMBL ID or chromosome location.
- (3) The output can be viewed in (a) heatmap or (b) browser view using toggle buttons on the top right side of the web interface.
 - a Heatmap view shows gene expression at different developmental stages. A heatmap is provided on the panel with gradations of blue color intensity. Higher intensity of blue indicates high gene expression compared to other developmental stages investigated in this study. The output is shown as gene name, ENSEMBL ID and chromosome location and the relative expression of query gene.
 - b Browser view shows all the genes and transcripts expressed in lens and retina in the query chromosomal location. Unannotated genes are displayed as *MSTRG.XXXX.XXXXX.X*.
- (4) The chromosomal window on the browser view can be increased or decreased using the magnification slider provided on the top right of the browser view panel.
- (5) The heatmap view and the browser view can be downloaded as high-resolution images using a dropdown export menu provided on the top right-hand side of the web interface. Along with the views, the raw and quantile normalized values for each isoform of a gene can be downloaded using the heatmap data option on the export menu.
- (6) Further sources for the RNA seq-data used for the analysis in this study are provided at the bottom of the web interface.

As shown in Fig. 3A, for instance – on investigating the gene expression profile of *Bfsp1* in the lens at the gene level, at a threshold of 5 TPM for raw expression level, an output is generated with both heatmap and browser view. In the browser view, all genes expressed in the lens and retina at the chromosomal location as *Bfsp1* can be visualized. In the heatmap view, relative expression of *Bfsp1* at various developmental stages is shown. We can compare the expression of splice forms of *Bfsp1* at different developmental stages using the transcript level option i.e. while the expression of isoform ENSMUST00000099296 increases with development, the expression of ENSMUST00000028907 is highest at P0.

Express re-captures the general expression trends in developing lens tissue for various transcription factors as shown in Fig. 3B. For example, Pax6, Sox2, Mab2111, Foxe3, Pitx3 and Mafg exhibit high expression in early (embryonic) lens development stages compared to late (postnatal) stages. In contrast, Maf, Prox1, and Sox1 expression is lower in early lens developmental stages and higher in later stages. Similarly, *Express*-based analysis of gene expression for Crystallin genes in lens development revealed an increased expression of Crystallin genes in postnatal stages compared to embryonic stages in developing lens tissue as shown

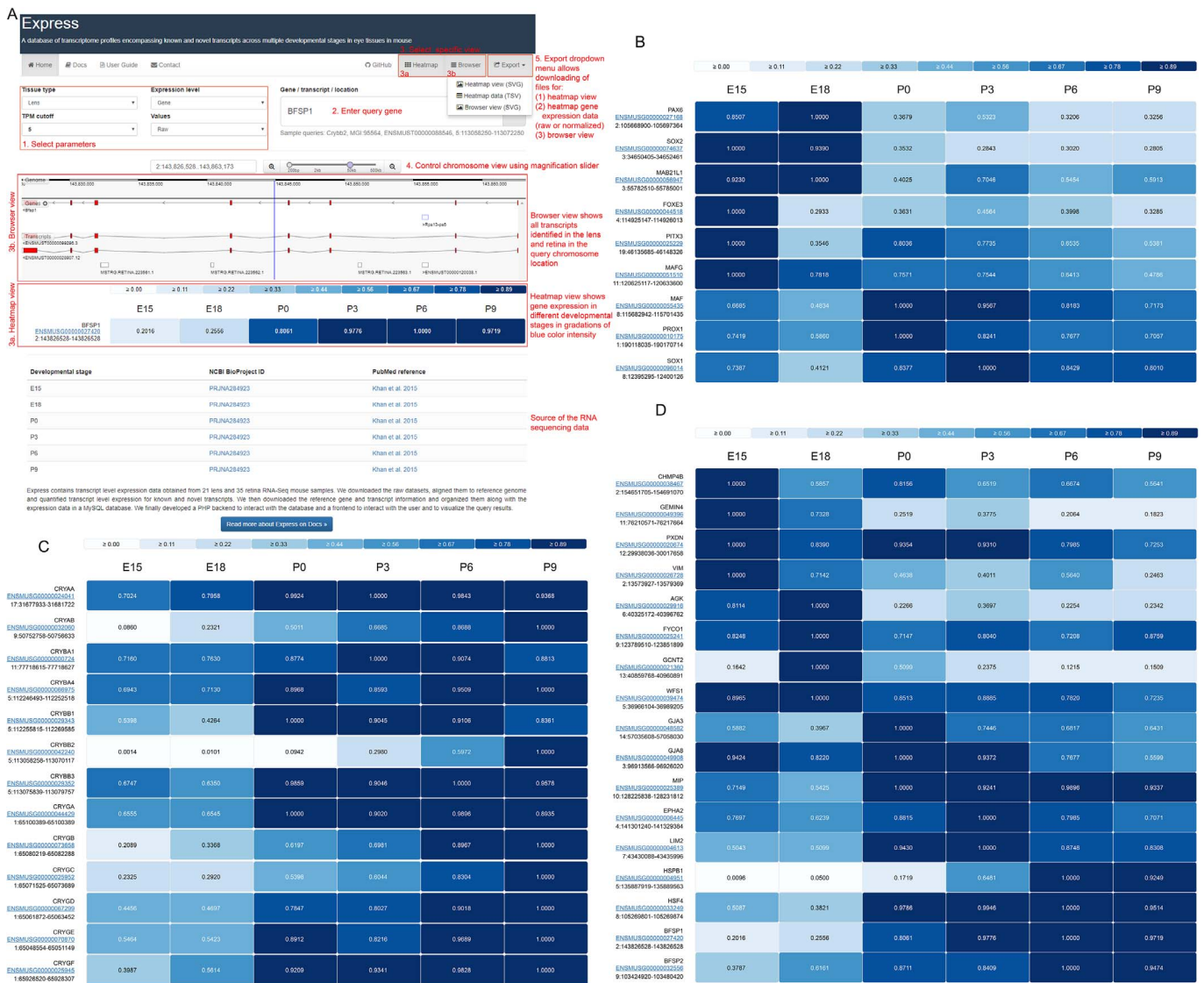


Fig. 3. User guide for employing Express to investigate eye gene expression is highlighted in panel 3A. 1) User selects parameters, 2) Enters query gene or chromosomal region, 3) Selects view options, 3a. with heatmap view can visualize gene expression in various developmental stages, 3b. with Browser view can visualizes different transcripts, 4) Uses magnification slider to controls chromosomal range, 5) Can use the Export dropdown menu to download heatmap view, raw or normalized gene expression data or browser view. Heatmaps showing the expression profiles of selected (B) transcription factors (C) crystallin genes and (D) non-crystallin genes in multiple developmental stages of mouse lens. Expression data is normalized by the maximum expression level of a given transcript across stages and visualized as heatmap in Express.

in Fig. 3C. Express also shows varying expression profile of various non-crystallin genes that are linked to human congenital/pediatric cataract (Fig. 3D). For example, Chmp4b, Gemin4, Pxdn, Vim, Agk, Fyco1 and Wfs1 are expressed highly in early embryonic stages, while Epha2, Gja3, Gja8, Lim2, Mip, Bfsp1 and Bfsp2 exhibit high expression in late embryonic and postnatal stages.

3.4. Development of Express as a user-friendly tool

Express provides transcript level expression data for mouse lens and retina across different developmental stages for known and novel transcripts as identified by StringTie. The mouse developmental stages are expressed as embryonic (E) or post-natal (P) followed by a number that indicates the number of days after fertilization or birth, respectively (e.g. E18 corresponds to an embryo dissected 18 days after the vaginal plug was observed, while P0 corresponds to the day of birth). RNA-seq data is available for 7 developmental stages of the lens (E15, E15.5, E18, P0, P3, P6, and P9) and 11 developmental stages of the retina (P2, P10, P11, P21, P28, P30 P40, P48, P50, P60 and P90). In express,

users can also search for cell-type specific expression profiles where available. For instance, a representation for lens dataset such as P0:E and P0:F stands for the epithelial and fiber compartments in lens. The fraction of transcripts for each developmental stage for lens and retinal samples is shown in Fig. 2. Although majority of the lens developmental stages exhibit ~17% of completely novel transcripts, the proportion of completely novel transcripts in retina were found to be significantly higher and varying with expression threshold. Observed fraction of completely novel transcripts was found to be < 25% across majority of the retina stages when transcripts were filtered to include only those expressed greater than 5 TPM in retinal samples (see Fig. 2 and Supplementary Fig. S2). The number of retina-expressed transcripts that are identified to be novel in this study is comparable to that previously reported in the human retina (Farkas et al., 2013), and therefore supports the finding that retinal cells potentially express a large number of uncharacterized transcripts. In Express, users can filter to view only those transcripts resulting from a search that satisfy one of the four levels of confidence in expression levels – 1) transcripts exhibiting a non-zero expression level in TPM in at least one developmental stage, 2)

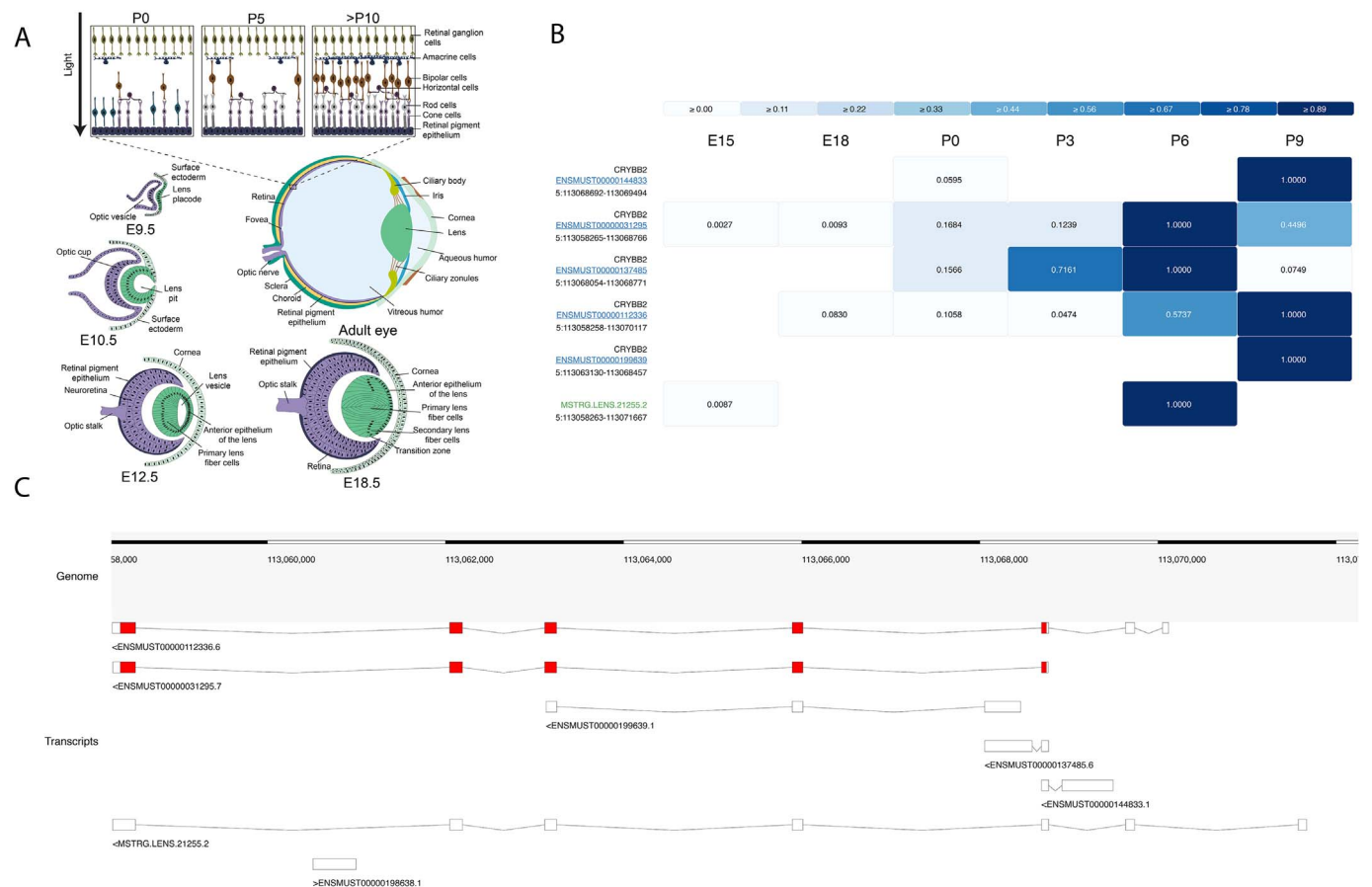


Fig. 4. Overview of the mouse eye development and user interface. (A) In the initial stages of eye development, the optic vesicle interacts with the overlying non-neural surface ectoderm at embryonic day (E) 9.5 in mouse and induces its thickening to form the lens placode. Subsequently at E10.5 the optic vesicle and the lens placode interact to develop into the optic cup and the lens pit, respectively. The lens pit closes to form the lens vesicle and the overlying ectoderm contributes to the corneal epithelium. The posterior cells of the lens vesicle differentiate to form the primary lens fiber cells while cells of the anterior epithelium of the lens divide to form new epithelial cells that migrate towards the transition zone. Cells at the transition zone exit the cell cycle and terminally differentiate to form the secondary fiber cells. Further, the fiber cells migrate towards the center of the lens, and as they terminally differentiate, undergo organelle degradation, resulting in an organelle free zone in the center of the lens by E18.5. Further development and differentiation events lead to the formation of the adult eye where the anterior region consists of the cornea, iris, ciliary body and ciliary zonules. The posterior of the lens consists of the retina, retinal pigment epithelium, choroid and sclera. A more detailed diagram of the retina shows that it is composed of several distinct cell types, including the retinal ganglion cells, amacrine cells, bipolar cells, horizontal cells and the rod and cone photoreceptors. Retinal ganglion cells and cone cells are differentiated and functional by E18.5 and by postnatal day (P) 5, amacrine cells, bipolar cells, horizontal cells and rod cells are formed. By P10, all the neuronal cells in the retina have completely connected synaptic junctions. Rod and cone cells synapse with horizontal cells for communicating with other photoreceptors and with bipolar cells, which further synapse with amacrine cells. The amacrine cells in turn synapse with the retinal ganglion cells. (B) Heatmap view from Express resulting from a query for the chromosomal location “5:113058250-113072250” when lens is selected as tissue with TPM cutoff = 5 using d3.js JavaScript package. The transcripts are sorted by novelty category (known transcripts colored in blue and linked to its Ensembl transcript page, completely novel transcripts colored in green and unannotated transcripts colored in black). The transcripts within each novelty category are also sorted by averaged normalized expression values for the row. In lens datasets some development stages are marked as “E” for epithelium and “F” for fiber cells and unmarked development stages represent whole tissue. Also in retina some development stages are marked as “C” for cone and “R” for rod cells and unmarked development stages represent whole tissue. The marked datasets are derived from the given cell type. (C) Browser view from Express from a query for the chromosomal location “5:113058250-113072250” using BioDalliance JavaScript package.

transcripts with at least 1 TPM in at least one developmental stage 3) transcripts with at least 2 TPM in at least one developmental stage and 4) transcripts expressed with at least 5 TPM in at least one developmental stage (default threshold). A summary of eye developmental stages for ready comparison of ocular morphological changes with *Express* data stages is shown in Fig. 4A. At 5 TPM cut-off, the lens samples were found to exhibit ~16% completely novel transcripts across stages. In contrast, the retinal samples were found to comprise of ~22% completely novel transcripts. When lower expression thresholds were used the fraction of completely novel transcripts significantly increased in retinal samples. It is likely that the high number of novel transcripts in retinal samples is likely due to the several distinct types of cells in the retina (Fig. 4A and Supplementary Fig. S2). Indeed, the total number of transcripts identified in mouse retina in this study are very similar to the numbers reported in human retinal samples (Farkas et al., 2013). *Express* allows a heatmap view for any specific query, as shown in the example query of

the chromosomal location “5:113058250-113072250” for the lens (Fig. 4B). A legend is provided at the top to indicate the intensity of the color for normalized expression values across the heatmap (dark color corresponds to high expression), while the heatmap itself shows transcripts in rows and different developmental stages in columns that represent progressive stages. The row labels show the gene symbol, Ensembl transcript ID linked to its official page or novel transcript ID and the chromosomal location of the transcript. The heatmap view can be exported using the Export dropdown menu in the navigation bar in SVG format, which can provide high-quality vector image for use in publications. Fig. 4C shows the browser view for the same query (for the chromosomal location “5:113058250-113072250”). The mouse genome is provided in the “Genome” track and the “Transcripts” track provides the view showing the various transcript structures for known and novel transcripts encoded by this genomic location. The browser is interactive and one can drag and navigate through neighboring transcript structures in the browser. The browser view can also be exported

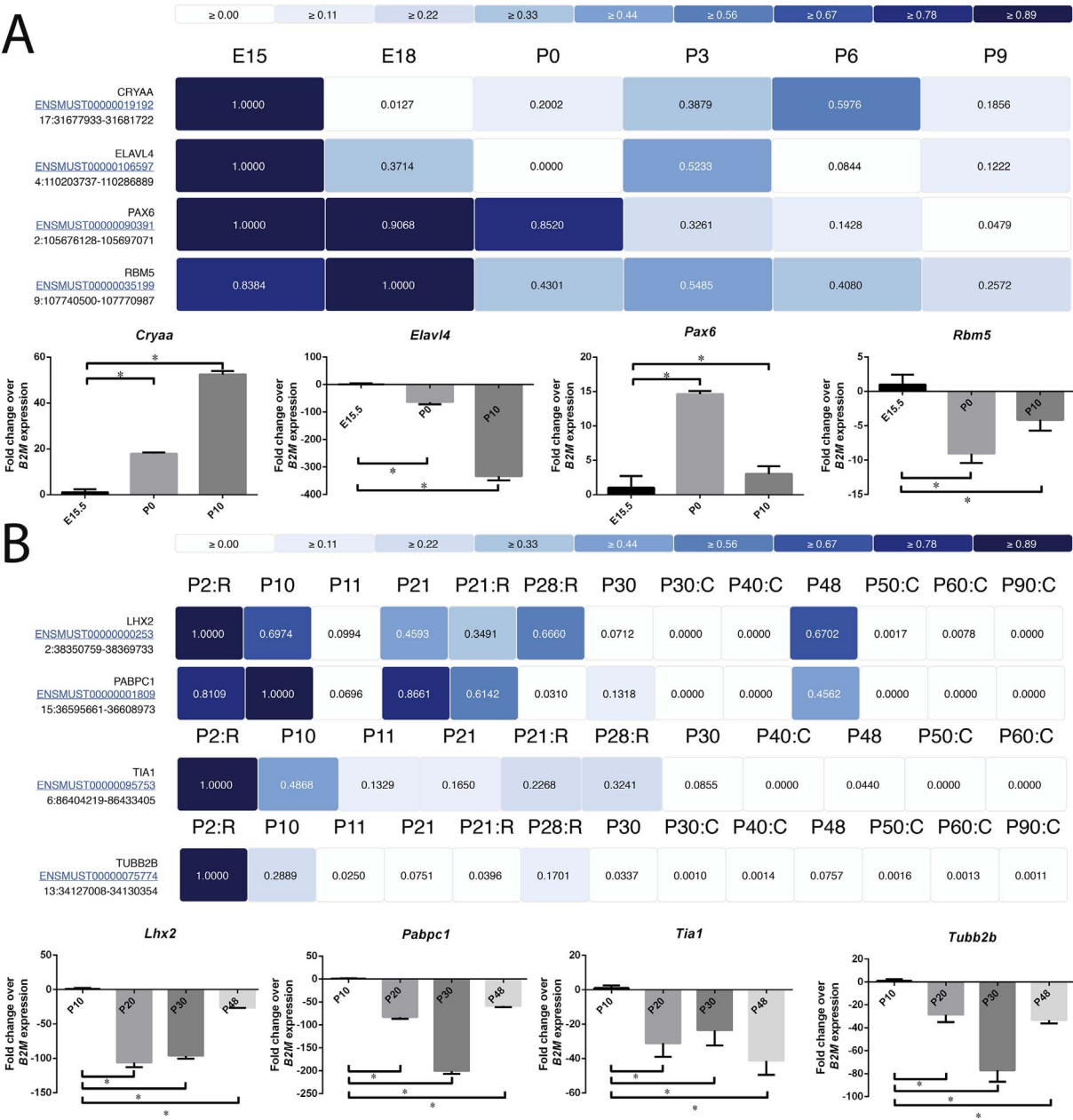


Fig. 5. Heatmaps showing the expression profiles of selected transcripts in multiple development stages of mouse eye tissues. Expression data were normalized by the maximum expression level of a given transcript across stages and visualized as heatmap in Express. Expression profile of selected transcripts for (A) lens and (B) retina were downloaded from Express and shown in form of heatmap. In retina datasets some development stages are marked as “C” for cone and “R” for rod cells and unmarked development stages represent whole tissue. The marked datasets are derived from the given cell type. Their expression profile was also verified for multiple development stages using qPCR with B2M as housekeeping control and shown as additional panels.

using the Export dropdown menu in the navigation bar in SVG format, which can serve to provide high-quality vector image for use in publications. In addition, users can also download both the raw and normalized expression levels resulting from the search for a gene of interest as tab-delimited text files.

3.5. Validation of transcript-expression in lens and retina

We identified several genes and their corresponding transcripts that were found to be significantly altered across the developmental stages in lens and retina. We validated the expression pattern of these genes as well as other established genes as a representative set of very significantly altering transcripts across stages to evaluate expression levels reported in Express. In particular, we downloaded the expression profile

of the selected transcripts (in the form of a heatmap) from Express for each tissue subtype and experimentally validated their levels for multiple development stages using RT-qPCR (see Materials and Methods). In the lens, we validated the expression of Pax6, Elavl4 and Rbm5 (Fig. 5A, Fig. S3A). Pax6 (Paired box 6) is a transcription factor essential for eye development in mice and humans. Mutations in Pax6 have been linked to congenital cataract, aniridia and anophthalmia in humans (Glaser et al., 1994) and haplo-insufficiency of Pax6 in mice results in small eyes (Sey) in mice (Hogan et al., 1986; Hill et al., 1991). RT-qPCR shows that Pax6 expression is elevated in early postnatal stages in agreement with Express (Fig. 5A). Elavl4 (ELAV (Embryonic Lethal, Abnormal Vision, Drosophila)-like 4 (Hu antigen D) belongs to ELAV protein family and is expressed in the mouse lens and frog retina (Amato et al., 2005; Bitel et al., 2010). Elevated expression of Elavl4 in

the mouse lens increases the expression level of its targets (*GAP43* and *CamKII α*), which is a similar outcome to its overexpression in brain tissue (Bitel et al., 2010). We find that the expression of *Elavl4* in mouse lens is high during embryonic stages and reduces gradually in postnatal stages (Fig. 5A), as predicted by the transcriptome datasets in *Express*. *Rbm5* (RNA binding motif protein 5) belongs to the Rbm protein family and is associated with lung cancer (Shao et al., 2012; Li et al., 2014; Su et al., 2014; Yang et al., 2016). We find that *Rbm5* is expressed highly at embryonic stages and its expression reduces during early postnatal stages (Fig. 5A). While this is the first report of *Rbm5* expression in the lens, another member of the Rbm family, *Rbm24* is expressed in the vertebrate eye and its deficiency in zebrafish causes microphthalmia (Lachke et al., 2012; Maragh et al., 2014).

Further, we also validated the expression of *Express*-predicted genes *Lhx2*, *Pabpc1*, *Tia1* and *Tubb2b* in the retina (Fig. 5B, Fig. S3B). *Lhx2* (LIM homeobox 2) encodes an eye field transcription factor that is expressed from the earliest stages of optic development. *LHX2* mutations in human as well as its knockout in mice causes anophthalmia (Porter et al., 1997; Desmaison et al., 2010). As indicated by *Express*, RT-qPCR shows that *Lhx2* expression reduces in the retina in late postnatal stages (Fig. 5B). *Pabpc1* (Poly A-binding protein, cytoplasmic 1) binds to the poly A tail of mRNA and modulates its susceptibility to cap-mediated mRNA decay (Walters et al., 2010). RT-qPCR shows that *Pabpc1* is expressed highly at early postnatal stages and its expression reduces significantly at later developmental stages (Fig. 5B) until P30 when its expression increases again. *Tia1* (T-Cell-Restricted Intracellular Antigen-1) promotes the recruitment of U1 snRNP to splice sites and is implicated in lymphoma and leukemia (Forch et al., 2002; Milne et al., 2009; Koreishi et al., 2010), which is also expressed in the mouse lens (Lachke et al., 2011). We find that *Tia1* expression gradually decreases in the retina with age (Fig. 5B). *Tubb2b* (Tubulin, Beta 2B Class IIb) is a component of microtubules. *Tubb2b* mutations result in congenital fibrosis of extraocular muscles (CFOEM), which leads to ptosis (drooping eyelids) and restricted eye movements in humans (Cederquist et al., 2012). RT-qPCR confirms that *Tubb2b* expression is high at P10 and reduces sharply at P30 before increasing again at P48 (Fig. 5B) as predicted by *Express*. We also verified the levels of the control genes compared across time points for reference as shown in Fig. S3C.

We also investigated how *Express* compares with the established expression pattern for the gamma-Crystallin family of genes. A previous study describes the expression of different *Cryg* family transcripts at the mouse stages E16.5, P1, P10, P20, P30, P40, P80, P120, and P180 (Goring et al., 1992). We compared *Cryg* gene expression for the stages in *Express* that are closest in developmental time to the stages in the Goring et al. study. Specifically, we compared *Cryg* expression in *Express* for E15, P0 and P9 that are close to the stages E16.5, P1 and P10 in the Goring et al. study. Using raw expression and TPM cut-off of 5, we

find that there is good agreement between the *Express* and previous findings for the general trends of the *Cryg* genes, namely *Cryge*, *Crygf*, *Crygb*, *Crygc* and *Crygd* (Fig. 6). *Cryga* showed a slight deviation from the Goring et al. study in that it did not exhibit a slight reduction at P0 prior to being high at P10 (although it exhibits general agreement with the previous study in that the expression of *Cryga* was higher at P9 compared to E15). Therefore, these findings offer further support that gene expression data in *Express* reflects the experimentally validated and established gene expression patterns in the lens.

4. Conclusion

A number of studies in the past have focused on studying the expression landscape of genes using microarrays across developmental stages (Farjo et al., 2002; Blackshaw et al., 2004; Zhang et al., 2006) in mouse eye development and specialized databases (Lachke et al., 2012; King et al., 2015) have been built. However, our understanding of the transcript structure, expression and their splicing alterations is just beginning to emerge during lens development across model systems (Srivastava et al., 2017). Here, we present *Express* which to our knowledge is the first large-scale carefully pruned transcriptomic resource based on eye tissue RNA-seq data to provide a user-friendly portal for studying and visualizing the expression levels of both the known and novel transcript isoforms across developmental stages in mouse eye tissues. Further, we validate several transcripts using RT-qPCR across multiple developmental stages in mouse lens and retinal tissues to confirm that the *Express*-quantified levels of transcripts are in agreement with the detected expression levels from RNA-seq quantification pipeline employed in this study. We found several transcripts encoding RNA-binding proteins to be highly expressed in embryonic development that are significant down-regulated in post-natal stages suggesting that post-transcriptional control of gene expression may function in early eye development.

Our analysis suggests that retinal samples exhibit a significant number of novel transcripts comparable to a recent analysis of human retinal transcriptomes (Farkas et al., 2013). It can be speculated that these novel RNA transcripts may reflect cell type specific functions. Hence resources like *Express* can not only further our understanding of the tissue-specific developmental transcriptome but can also serve to improve gene annotations in mouse. Although several of these novel transcripts identified across developmental stages in our analysis could be non-coding, with some isoforms resulting from errors in transcript assembly process, we include them in *Express* since some of these transcripts could have regulatory roles that are yet to be discovered. However, since *Express* enables users to filter the gene and transcripts resulting from a search based on their expression levels, a user can choose to analyze and explore only the most promising candidates.

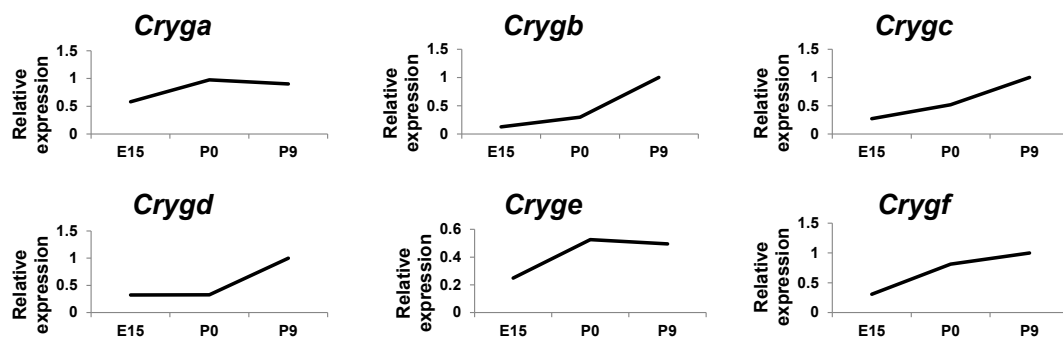


Fig. 6. Gene expression analysis of *Cryg* family of genes. Raw Expression profiles of *Cryg* genes in mouse lens were downloaded from the *Express* database to investigate if they are in agreement with previously described patterns (reference: Goring et al., 1992). Expression values for specific *Cryg* genes in the stages closest to the developmental time points in the previous study are plotted (E15 in *Express* in lieu of E16.5 in Goring et al., 1992; P0 in lieu of P1, P9 in lieu of P10). The general expression patterns for the *Cryg* genes correlate well between *Express* and the previous findings. Specifically, *Crygb*, *Crygc*, *Crygd* and *Crygf* expression increases as development progresses and is highest at P9, while *Cryge* expression elevates at P0 and beyond. The only minor deviation is exhibited by *Cryga* whose expression increases from E15.5 through P9 in *Express*, instead of the slight decrease at P1 prior to increasing again at P10 as previously described.

We anticipate that future versions of *Express*, which will continue to be manually curated and pruned to comprise of only high quality RNA-seq expression data across eye tissues, can be a useful resource for prioritization of candidate genes from exome sequencing analysis for patients with ocular defects as well as for providing a functional and developmental context to investigate the significance of differentially expressed genes in mouse mutants with eye defects.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at <http://dx.doi.org/10.1016/j.exer.2018.01.009>.

Authors' contribution

GB, RS and SCJ designed the study, implemented the computational approaches and performed the data analysis. GB, RS, and SD interpreted the data. SD and SAL designed and validated the predicted targets experimentally. RS, GB, SD wrote the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare no competing financial interests.

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