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unified and comprehensive single-cell atlas of the mouse retina, integrating both unpublished and

Experimental enrichment for large data generation of

integration analysis and cell type annotation

dissemination through various interactive web

Li et al., iScience 27, 109916 June 21, 2024 © 2024 The Authors. Published by Elsevier Inc.

[https://doi.org/10.1016/](https://doi.org/10.1016/j.isci.2024.109916) [j.isci.2024.109916](https://doi.org/10.1016/j.isci.2024.109916)

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Comprehensive single-cell atlas of the mouse retina

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SUMMARY

Single-cell RNA sequencing (scRNA-seq) has advanced our understanding of cellular heterogeneity by characterizing cell types across tissues and species. While several mouse retinal scRNA-seq datasets exist, each dataset is either limited in cell numbers or focused on specific cell classes, thereby hindering comprehensive gene expression analysis across all retina types. To fill the gap, we generated the largest retinal scRNA-seq dataset to date, comprising approximately 190,000 single cells from C57BL/6J mouse retinas, enriched for rare population cells via antibody-based magnetic cell sorting. Integrating this dataset with public datasets, we constructed the Mouse Retina Cell Atlas (MRCA) for wild-type mice, encompassing over 330,000 cells, characterizing 12 major classes and 138 cell types. The MRCA consolidates existing knowledge, identifies new cell types, and is publicly accessible via CELLxGENE, UCSC Cell Browser, and the Broad Single Cell Portal, providing a user-friendly resource for the mouse retina research community.

INTRODUCTION

The retina is a highly heterogeneous part of the eye that captures and processes the light signal.^{[1–3](#page-10-0)} The processing is enabled through five classes of retinal neurons: photoreceptors (PR), horizonal cells (HC), bipolar cells (BC), amacrine cells (AC), and retinal ganglion cells (RGC), which form an intricate circuitry necessary for processing and relaying the light signal to the visual cortex. Non-neuronal cells such as Müller glia cells (MG), microglia, astrocytes, and retinal pigment epithelial cells (RPE) provide structural integrity of the tissue and carry out various supporting roles such as metabolism and neuronal homeostasis in the retinal microenvironment.^{[4](#page-10-1),[5](#page-10-2)} Characterization of distinct retinal cell types is, therefore, critical in advancing our understanding of the fine intricacies of cell interactions involved in retinal biology and visual disorders.

Single cell technologies have opened a window into knowledge of cellular heterogeneity and intricate cell-to-cell interactions that cannot currently be resolved at the tissue level and have allowed exploration of individual cellular expression signatures, which can be mapped to unique molecular cell types.^{6,[7](#page-10-4)} The resulting cell atlas can serve as a foundation for numerous applications, including the annotation of cell types in other scRNA-seq experiments, 8 the identification of differentially expressed targets for purification or manipulation, 9 and the gen-eration of marker panels useful for single-molecule imaging, including spatial profiling.^{[10](#page-10-7)} While studies have demonstrated cell type heterogeneities in various tissues, several perplexing issues remain to be addressed in establishing a comprehensive cell atlas such as the agreement on cell type definitions across different experiments or whether enough cells have been profiled to exhaust all existing cell types. Integrated analyses of various scRNA-seq datasets from different studies, therefore, can provide an important insight that comprehensively addresses such issues.

The mouse retina provides an important model for the study of neurobiology, with more than 130 distinct cell types characterized through previous scRNA-seq studies.^{[7](#page-10-4)[,9](#page-10-6)[,11–15](#page-10-8)} However, the scRNA-seq datasets have been generated separately for BC^{[11](#page-10-8)} AC,¹² and RGC,^{[9](#page-10-6)[,13](#page-10-10)[,15](#page-10-11)} with the largest dataset containing just under 36,000 cells, making it difficult to use in aggregate. Though most of these datasets are independently browsable on the Broad Single Cell Portal^{[16](#page-10-12)} and accessible through separate databases such as the Gene Expression Omnibus (GEO) repository, it can be challenging to assess gene expression patterns across all retinal cell types. Ensuring these atlases define a complete set of retinal cell types remains a major challenge that can only be addressed by powering studies to sufficiently profile the rarest retinal cell types. Here, we generated scRNA-seq data of over 189,000 cells in the mouse retina to complement 141,000 cells from six publicly available

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Figure 1. Overview of single cell atlas of the mouse retina

(A) The workflow for generating unpublished scRNA-seq datasets. The data generation process involved using mice aged from P14 to 12 months. Following retina dissection and cell dissociation, single cells were enriched using autoMACS with Anti-CE73-PE antibodies or Anti-CD90.1 beads for specific amacrine, bipolar, and retinal ganglion cells. Subsequently, 10X single-cell RNA sequencing was performed on both the unenriched and enriched single cells. The retained single cells were then utilized in downstream atlas construction.

(B) The integrated analysis workflow for constructing the MRCA. To construct a comprehensive unified single-cell reference of the mouse retina, we generated 16 unpublished scRNA-seq samples of the mouse retina and incorporated four curated public datasets to enhance specific amacrine, bipolar and retinal ganglion cells. The collected data were processed using the Cell Ranger and CellQC pipeline to produce feature count matrices. Feature counts were then processed to remove estimated empty droplets, ambient RNA, and doublets. The retained cells were integrated using scVI to eliminate batch effects across samples. The trained low-dimensional embeddings were used to calculate cell dissimilarities and identify clustering through a two-level clustering approach. Major class and subclass cell types were annotated using canonical marker genes and public labeling. To facilitate user-friendly access and exploration, the MRCA was deployed on accessible interactive web browsers, including CELLxGENE, UCSC Cell Browser, and Single Cell Portal.

Figure 1. Continued

(C) Pie chart displaying the percentage of cells contributed by each dataset used in the MRCA. (D) UMAP visualization of the MRCA colored by major classes. (E) Dot plot illustrating the expression of canonical markers for major classes. See also [Figure S1](#page-10-13) and [Table S1](#page-10-13).

scRNA-seq datasets,^{[9](#page-10-6)[,11–15](#page-10-8)} creating a unified cell atlas of the wild-type mouse retina containing over 330,000 cells. Our integrated analysis presents a comprehensive characterization of all major cell classes in the retina, including non-neuronal types, as well as a consensus cell type annotation of BCs, ACs, and RGCs. Accessible, interactive web browsers have facilitated easy visualization of atlas characterizations and exploration of gene expression in the MRCA. The comprehensive unified MRCA will serve as a valuable resource for the community.

RESULTS

Generation of scRNA-seq dataset for wild-type mouse retina

To establish a comprehensive atlas of the mouse retina, we performed scRNA-seq profiling with C57BL/6J mouse retina tissue samples, aged from P14 to 12 months, for over 189,000 cells ([Figure 1A](#page-2-0) and [STAR Methods\)](#page-12-0). As summarized in [Table S1](#page-10-13), six retinal samples of varying ages were dissociated without enrichment, and ten samples of eight weeks old were enriched using surface markers CD73 and CD90.1 to enrich for rare cell populations. Depletion of rod PR was achieved by removing cells positive for CD73 using anti-CD73-PE antibody and anti-PE mag-netic beads, which primarily label PR precursors and mature rod PR in mice.^{12,[17](#page-10-14)} To enrich ACs and RGCs, CD90.1 positive cells are selected.^{[18](#page-10-15),1}

Integration of scRNA-seq datasets for the mouse retina

To compile the most comprehensive scRNA-seq data for the MRCA, we curated and obtained six publicly available scRNA-seq datasets, each enriched for a specific cell type using transgenic labels or immunolabeling combined with FACS. Together, they consisted of over 141,000 cells. To consolidate the transcript annotation between different datasets, we used the Cell Ranger (version 7.0.1) pipeline to align raw FASTQ files from four datasets obtained from GEO and Sequence Read Archive (SRA) repositories. Count matrices of these datasets were generated using the mm10 reference genome obtained from 10x Genomics ([https://cf.10xgenomics.com/supp/cell-exp/refdata](https://cf.10xgenomics.com/supp/cell-exp/refdata-gex-mm10-2020-A.tar.gz)[gex-mm10-2020-A.tar.gz](https://cf.10xgenomics.com/supp/cell-exp/refdata-gex-mm10-2020-A.tar.gz)). Five of the published studies were sequenced on the 10x Genomics 3' platform, and one (Shekhar et al.) was generated using the Drop-seq protocol.⁷ The Drop-seq data were aligned against mm10 and processed into count matrices using the Drop-seq pipeline (<https://github.com/broadinstitute/Drop-seq>). The cell type labels of previous annotations were obtained from the Broad Single Cell Portal website.¹⁶ To remove technical variations introduced across different experiments or studies, scVI^{[20](#page-11-0)} was applied to integrate all newly generated and public datasets, generating a low-dimensional representation ([Figure 1](#page-2-0)B and [STAR Methods\)](#page-12-0). Putative cell doublets were further removed using the deep learning doublet identification method Solo^{21} Solo^{21} Solo^{21} ([Figure S1](#page-10-13)A).

In the integrated data, the public dataset accounts for 43%, while the newly generated data accounts for the remaining 57% [\(Figure 1](#page-2-0)C). Within the integrated UMAP, 97 clusters were identified ([Figure S1](#page-10-13)B). These clusters were annotated as one of 12 major classes, including PR, BC, AC, RGC, HC, MG, RPE, astrocyte, microglial, endothelial, and pericyte, using known marker gene expression^{22,[23](#page-11-3)} ([Figure S1](#page-10-13)C). Cells from non-enriched retina samples showed a distribution across major classes at an expected proportion, with rod PR as the biggest proportion.^{[2](#page-10-17)} In contrast, enriched samples from both newly generated data and previous studies showed the expected skewed distribution of cell types in BCs, ACs, and RGCs [\(Figure S1](#page-10-13)D). The two newly generated samples with enrichment methods, CD73⁻ and CD90.1⁺ samples, were primarily composed of BCs and ACs, respectively, contributing to 83% (122.6K out of 147.7K) and 25% (11.2K out of 44K) of all BCs and ACs in the integrated data, respectively.

Previous studies have identified 15 distinct types of BCs, 63 ACs, and 45 RGCs.^{9,[11,](#page-10-8)[12](#page-10-9)} To determine the consensus annotation of neuronal types for these subclasses, we performed clustering analysis at higher resolution within individual BC, AC, and RGC classes ([Figures 1D](#page-2-0) and 1E).

15 types of bipolar cells

A total of 147,700 BCs were identified in the integrated datasets, with 122,600 cells from our newly generated CD73- sample and 19,800 cells from the Shekhar et al. study.¹¹ The integrated analysis identified 15 BC clusters, corresponding to previously annotated BC types ([Figures 2](#page-4-0) and 2B and [STAR Methods](#page-12-0)). The 15 clusters of integrated BCs showed a generally even distribution of cells from various samples, with the exception of two types, BC1A and BC1B, where more than 90% of populations came from the study by Shekhar et al. possibly due to differences in enrichment methods ([Figures 2](#page-4-0)A, 2D, [S2](#page-10-13)D, and S2E). The final annotation of BCs revealed consistent expression profiles of previously identified BC type marker genes^{[11](#page-10-8)[,24](#page-11-4)} ([Figures 2B](#page-4-0) and 2C). With a significant addition of BCs in the MRCA, clear separation of BC8 and BC9 is observed, which were merged but demonstrated substructure in the Shekhar et al. dataset [\(Figures 2](#page-4-0)A and 2B). The separate clusters showed proper expression patterns of known markers like Cpen9 in BC9.^{11,[25](#page-11-5)} In addition, additional BC type markers were identified via differential gene expression analysis, which showed more specific expressions than previous marker genes, such as Tafa4 in BC4, Ptprt in BC5A, and Gm13986 in BC8 ([Figure 2E](#page-4-0)). Interestingly, despite an almost 10-fold increase in the number of BCs in our analysis, we did not observe any sign of a novel cell type, which suggests that the mature mouse retina likely only contains 15 BC types.

Figure 2. Bipolar cells

(A) UMAP visualization of BCs colored by public cell type labels from Shekhar et al. 2016. The newly discovered cells without public labeling are colored in gray. (B) BCs colored by the 15 annotated annotated cell types.

(C) Dot plot of BC type marker gene expression in the 15 types.

(D) Pie chart showing the percentage each data source making up BC1A and BC1B population.

(E) Dot plot of new markers for three BC types: BC4, BC5A, and BC8. The three new markers exhibit more exclusive expression patterns. See also [Figures S2](#page-10-13) and [S3](#page-10-13).

Amacrine cells

Through CD90.1 positive enrichment, the newly generated samples contributed 11,200 ACs, in addition to the 27,600 ACs from Yan et al.^{[12](#page-10-9)} in the integrated dataset ([Figures S4A](#page-10-13) and S4B). Utilizing the collected data, the integrated analysis annotated 63 AC types, revealing consistent expression profiles of known marker genes [\(Figures 3A](#page-5-0) and 3B and [STAR Methods](#page-12-0)). While a minimal batch effect in each cluster was observed across different sample sources, CD90.1⁺ and Ma et al. RGC samples showed biased enrichment toward GABAergic types except for AC4, AC10, and AC28 ([Figure S6](#page-10-13)A). The bias in cell type population appears to be directly tied to the preferential expression of Thy1 (CD90) in sub-populations of ACs [\(Figure S5](#page-10-13)D). In particular, Thy1 is characterized as being expressed primarily in GABAergic AC types. 26 26 26

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Figure 3. Amacrine cells

(A) UMAP visualization of AC cells colored by the annotated types.

(B) Dot plot of canonical marker gene expression in AC types.

(C) Four previously under-clustered AC types, i.e., AC18, AC20, AC36, and AC45, are split into two distinct clusters at a high resolution of clustering. (D) Feature plot showing the marker gene expression of the four previously under-clustered AC types.

(E) Dot plot showing the top 10 differentially expressed genes between split clusters for the previously under-clustered AC types. See also [Figures S4–S7.](#page-10-13)

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Figure 4. AC types by a high-resolution clustering

(A) Dot plot of DEGs expressed in two split clusters for AC_36, stratifying $Gbx2^+$ AC types in AC_36. (B) Visualization of AC cells colored by AC types at a high clustering resolution. See also [Figures S4–S7.](#page-10-13)

The integrated analysis of ACs demonstrated that four types, AC18, AC20, AC36, and AC45, have been previously under-clustered, each splitting into two clusters in the integrated UMAP [\(Figure 3](#page-5-0)C). AC18, which expresses Cck neuropeptide, 27 27 27 is split into C24 and C27 in our clustering and has been labeled as AC18A and AC18B in the final annotation [\(Figure 4B](#page-6-0)). Interestingly, the cell type marker Cck is highly expressed in AC18A, but not in AC18B [\(Figure 3D](#page-5-0)). AC20, which does not contain any known marker, is divided into C42 and C60 (AC20A and AC20B), with its marker Sema3a also expressed highly in AC20A, but not in AC20B ([Figure 3D](#page-5-0)). A non-GABAergic non-glycinergic (nGnG) type 4, AC36, is split into C58 and C61 (AC36A and AC36B), consistent with previous finding of two morphologically distinct AC36 types in the INL and displaced in the GCL, stratifying to S3 and S5 sublaminae of the IPL.^{[10](#page-10-7)[,28](#page-11-8)} By examining the list of differentially expressed genes (DEG) between the two broadly isolated types,²⁸ we annotated AC36A as the S3 type by the increased markers such as Gbx2, Tac1, and Pcdh8 and AC36B as the S5 type by Gad1, Gad2, and Id4. ([Figure 4A](#page-6-0)). Lastly, a catecholaminergic type 1 cell type,²⁹ AC45, is split into C64 and C66 (AC45A and AC45B). The expression of Chl1, which distinguishes catecholaminergic type 1 from type 2, was increased specifically in AC45A. The DEG analysis between the clusters of the previously under-clustered cell type revealed many genes enriched specifically in each cluster, with Cck, Sema3a, Chl1 being one of the top-ranked genes in AC18A, AC20A, and AC45A, respectively ([Figure 3E](#page-5-0)). Out of the four under-clustered cell types, only one, AC20, showed a biased sample source from Yan et al. data. Furthermore, while cells from Yan et al. were distributed across both AC45A and AC45B, AC45B contains an increased number of cells from the newly generated CD90.1 sample ([Figure S6](#page-10-13)A).

As a result, we have identified 67 AC types that can be grouped into four AC subclasses: 49 GABAergic, 10 Glycinergic, 3 Both, and 5 nGnG ACs. Within the final dataset, GABAergic ACs make up 67.7% of the total AC population, followed by Glycinergic ACs at 22.5%, GABA/ Glycinergic ACs at 1%, and nGnG ACs at 8.7%. However, these distributions are likely biased toward GABAergic ACs due to the inclusion of cells from CD90.1⁺ and CD90.2⁺ enriched collections.

Retinal ganglion cells

The integrated data contains 77,[9](#page-10-6)00 RGCs, primarily from the three publicly available datasets: Tran et al., $^{\circ}$ Jacobi et al., 13 13 13 and Ma et al.. 15 15 15 The integration of the collected data identified all 45 previously identified RGC types ([Figure 5A](#page-7-0) and [STAR Methods\)](#page-12-0). Examination of known cell type markers in the integrated data with the final annotation showed proper expression profiles in corresponding types^{[9](#page-10-6),[18](#page-10-15),[30](#page-11-10)} ([Figure 5](#page-7-0)B). Our integrated analysis of RGCs further divide two cell types, 16_ooDS_DV (ON-OFF direction-selective dorsal and ventral) and 18_Novel, into four clusters [\(Figures 5C](#page-7-0), 5D, and [S9](#page-10-13)A–S9C). 16_ooDS_DV, which contains both types with dorsal and ventral orienta-tion selective functional roles,^{31,[32](#page-11-12)} was split into 16_ooDS_D (C31) and 16_ooDS_V (C39), similar to the supervised clustering analysis done in the Tran et al. $^{\circ}$ and the clustering in Jacobi et al., 13 13 13 and Ma et al. 15 studies. Examination of the marker genes Calb1 and Calb2 demonstrated that C39 is the ventral selective type with high expression of Calb2, and C31 is the dorsal selective type with Calb1 expression.⁹ In addition, the 18_Novel type could also be split into 18_Novel_a (C36) and 18_Novel_b (C40), in agreement with the Ma et al. study.¹⁵ Examination of 18_Novel markers Pcdh20 and 4833424E24Rik revealed increased expression of both markers in 18_Novel_b, yet Pcdh20 expression was absent in 18_Novel_a [\(Figure S9](#page-10-13)E). The DEG analysis further demonstrated many genes selectively expressed in these two clusters, suggesting they are transcriptionally similar but independent types ([Figure S9D](#page-10-13)). In total, we have identified 47 RGC types in the MRCA ([Figure 5](#page-7-0)D).

Figure 5. Retinal ganglion cells

(A) UMAP visualization of RGC cells colored by the annotated types.

(B) Dot plot of canonical marker gene expression in RGC types.

(C) Two previously under-clustered RGC types, i.e., 16_ooDS_DV and 18_Novel, are split into two distinct clusters at a high resolution of clustering. Dot plot of Calb1 and Calb2 in the two split clusters of 16_ooDS_DV.

(D) Visualization of RGC cells colored by RGC types at a high clustering resolution. See also [Figures S8–S10](#page-10-13).

Non-neuronal retinal cells

To include the comprehensive set of cell types in the retina in the MRCA, 18,500 non-neuronal cells were integrated for six non-neuronal cell types, including astrocyte, endothelial, MG, microglia, pericyte, and RPE [\(Figures 1E](#page-2-0) and [S11](#page-10-13)A). These cells are evenly distributed in the collected datasets, except for astrocytes solely from the Benhar et al. dataset^{[14](#page-10-18)} [\(Figure S11](#page-10-13)B). After being combined with neuronal retinal cells, the MRCA consisted of 12 major classes and 138 cell types.

Data dissemination at accessible interactive web browsers

The MRCA has been made available for public access using the CELLxGENE platform ([https://cellxgene.cziscience.com/collections/](https://cellxgene.cziscience.com/collections/a0c84e3f-a5ca-4481-b3a5-ccfda0a81ecc) [a0c84e3f-a5ca-4481-b3a5-ccfda0a81ecc](https://cellxgene.cziscience.com/collections/a0c84e3f-a5ca-4481-b3a5-ccfda0a81ecc) and [https://mouseatlas.research.bcm.edu/\)](https://mouseatlas.research.bcm.edu/) [\(Figures 6A](#page-8-0)–6C). The MRCA is also accessible on

Figure 6. Visualization of MRCA in accessible interactive browsers

(A) Visualization of the MRCA in the CELLxGENE browser. The homepage depicts three panels to explore the MRCA. The left panel contains the pre-computed features facilitating the selection of cells by interested categories. The middle panel is the UMAP of the MRCA, colored by the annotated major classes. The right panel allows input of quick gene symbols and gene sets.

(B) Visualization of the subclass RGC atlas in the CELLxGENE browser. The middle panel depicts RGCs colored by the reclassified names selected in the left panel. (C) Visualization of gene expression for a BC9 marker, Cpne9, in the BC atlas. The left subfigure shows the BC types, and the right subfigure highlights the normalized gene expression values of Cpne9 for BC9 type in the middle panel. See also [Figure S11.](#page-10-13)

UCSC Cell Browser (<https://retina.cells.ucsc.edu>) and the Broad Single Cell Portal [\(https://singlecell.broadinstitute.org/single_cell/study/](https://singlecell.broadinstitute.org/single_cell/study/SCP2560) [SCP2560\)](https://singlecell.broadinstitute.org/single_cell/study/SCP2560). Pre-computed gene expression profiles of all cells included in the integrated analysis can be examined and visualized. Users also have access to the metadata information, including major class and cell type labels in the database. The accessible interactive web browsers of the MRCA can aid in easy access to the transcriptome profiles of any given mouse retinal cells without the bioinformatic burden and provides a valuable tool for the vision community.

DISCUSSION

As part of the central nervous system, the retina contains numerous neuronal types with distinct morphologies and functional roles.^{[1,](#page-10-0)[33](#page-11-13)} The heterogeneous cell type composition and the stereotypically patterned structure of the tissue makes the retina an ideal model for single-cell sequencing studies in establishing the single-cell atlas.^{7,[22](#page-11-2),[34](#page-11-14)} Although several scRNA-seq studies focusing on the retina tissue have been done previously,^{[7,](#page-10-4)[9](#page-10-6),[11–15](#page-10-8)} each available dataset contains single-cell profiles primarily of one or a few retinal cell classes with a limited number of cells. Furthermore, no systematic evaluation or comparison of the datasets has been done yet to cross-validate the cell type transcriptomes and address annotation consensus.

In this study, we generated scRNA-seq profiles of 189,000 retinal cells from 16 scRNA-seq experiments to perform an integrated analysis with 141,000 retinal cells from six previously reported datasets. Six out of the newly generated collections were done using endogenous retina

tissues with simple dissociation and without enrichment. PR constitute over 70% of the cell proportion in the retina,^{[2](#page-10-17)[,35](#page-11-15)} and there are only two subclasses of PR, which are well studied. Therefore, we utilized two methods for rare population cell type enrichment. The first way was depleting the rod PR. To achieve this goal, the rod PR cell surface marker, CD73, was used in seven of the 16 experiments. Though this marker is generally considered as a specific marker for rod PR, it is also expressed on the surface of a subset of ACs, HCs, and MGs. Depletion increased the enrichment of BCs from 12% to 90%. Furthermore, CD90.1 was used to enrich certain retinal neurons such as ACs and RGCs in three experiments. Enrichment of retinal cells with CD90.1 also showed an increased number of ACs with some RGCs.

One of the challenges in integrating and comparing publicly available data is that they are generated using different single-cell experi-mental platforms and analysis pipelines.^{36,[37](#page-11-17)} One public data enriched with BCs from Shekhar et al.^{[11](#page-10-8)} was generated using the Drop-seq^{[7](#page-10-4)} technology and was processed separately using the Cell Ranger transcript annotation. The four other sources of publicly available data were done using the 10x Genomics platform. A minimal batch effect across data sources was observed in the integrated analysis, with the expected distribution and clustering of major classes from corresponding sources. While the newly generated data without enrichment were primarily composed of rod PR, cells from the newly generated data with enrichment and publicly available data showed a proper distribution across BCs, AC, and RGCs.

Integrated analysis of various scRNA-seq datasets allowed us to examine AC, BC, and RGC types, which together comprise over 100 distinct cell types, including novel clusters. Through the integrated analysis, we addressed two key questions on the neuronal cell types in the retina: to confirm the consensus cell type signatures and to examine whether the total number of cell types of retinal neurons is exhausted. Following the initial integrated analysis to identify major classes, subsets of each major class were subjected to further integration and twolevel clustering to annotate all previously identified cell types, which showed an even distribution of data sources in general. The cell type annotation was achieved through examining known marker gene expressions and previous annotation labels when available. Although our newly generated data resulted in a significantly increased number of cells in the integrated analysis of BCs, ACs, and RGCs, we did not observe significant increases of novel cluster. As such, the previously reported set of BC types in the adult mouse retina is likely complete, supported by the more than 7-fold increase in BCs in the integrated data. On the other hand, our integrated analysis updates annotations of AC and RGC types. In particular, we observed several instances of previously under-clustered AC and RGC types splitting into distinct clusters in our analysis. For example, we confirmed the separation of 16_ooDS_DV types into two distinct clusters in the integrated data of RGCs, which was separated into dorsal and ventral selective types only through supervised clustering in the Tran et al. study⁹ and later confirmed in Jacobi et al.^{[13](#page-10-10)} study. Furthermore, we identified the separation of AC36 and assigned its clusters to S3 and S5, stratifying Gbx2⁺ AC types,^{[28](#page-11-8)} which strengthens our analysis by connecting to biologically distinct cell types. The separation of previously merged cell types into distinct clusters can be attributed to the increased number of cells in our integrated analysis. An integrated study may introduce new clusters caused by differences in expression or cell qualities among datasets. However, the QC metrics show that the identified novel clusters for ACs and RGCs have high-quality cells and are represented in dataset sources ([Figures S3](#page-10-13), [S6,](#page-10-13) and [S10\)](#page-10-13). Altogether, this suggests that, while our AC and RGC type annotations are comprehensive, they will likely continue to be refined by future studies.

Finally, we have deposited the MRCA into interactive web browsers that are user-friendly and publicly accessible. This allows for the examination of raw and normalized gene expression profiles of all retinal cells, along with their metadata such as major class and cell type annotation. The MRCA not only provides the consensus signature of mouse retinal cell types by comparing multiple scRNA-seq data but also alleviates the bioinformatics burden for many vision researchers who wish to examine transcriptome signatures in any cell type of their interest.

Limitations of the study

This study presents the first unified single-cell atlas of the wild-type mouse retina that integrates both unpublished and several public datasets. However, our study is subject to several limitations. There is a lack of experimental validation for the discovered novel types among amacrine and retinal ganglion cells. Potential validations through morphology or histology would bolster the credibility of these findings within the mouse retina. The dissociation technology utilized in this study focuses solely on single cells, excluding nuclei, due to limited access to singlenuclei RNA-seq (snRNA-seq) data from both unpublished and public resources. Incorporating snRNA-seq data would greatly enhance the transcriptomic reference of the mouse retina at the single-cell resolution.

STAR+METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at [https://doi.org/10.1016/j.isci.2024.109916.](https://doi.org/10.1016/j.isci.2024.109916)

ACKNOWLEDGMENTS

We thank Alice Tian for her meticulous proofreading of the manuscript. This project was funded by NIH/NEI R01EY022356, R01EY018571, S10OD032189, Chan Zuckerberg Initiative (CZI) award CZF2019-002425, RRF to R.C., and CZI 2023-330377 to J.L.

AUTHOR CONTRIBUTIONS

J.L., J.C., X.C., and R.C. conceptualized and designed the study. R.C. supervised the study. X.C. and Y.L. generated scRNA-seq data in this study. J.L., J.M., and S.P. compiled dataset collection. J.L., J.C., and S.P. developed the integrated analysis pipeline and performed the integration and annotation analysis. J.R.S, G.M., and B.J.F. provided public datasets before publishing. N.M.T. provided input for various annotation. All authors wrote, reviewed, and contributed to the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: February 19, 2024 Revised: March 18, 2024 Accepted: May 3, 2024 Published: May 8, 2024

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STAR**★METHODS**

KEY RESOURCES TABLE

(Continued on next page)

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources should be directed to and will be fulfilled by the lead contact, Rui Chen [\(ruichen@bcm.edu\)](mailto:ruichen@bcm.edu).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- Data: The raw sequencing reads from sixteen newly generated samples have been deposited at NCBI GEO and are publicly accessible under the accession GSE243413. The landing page for the MRCA data resources is accessible at [https://rchenlab.github.io/resources/](https://rchenlab.github.io/resources/mouse-atlas.html) [mouse-atlas.html.](https://rchenlab.github.io/resources/mouse-atlas.html) Processed cell-by-gene count matrices, along with cell type annotations, are available on Zenodo [\(https://doi.org/](https://doi.org/10.5281/zenodo.10815031) [10.5281/zenodo.10815031\)](https://doi.org/10.5281/zenodo.10815031). Furthermore, both raw and normalized count matrices and cell type annotations are publicly accessible on the CELLxGENE data collection at [https://cellxgene.cziscience.com/collections/a0c84e3f-a5ca-4481-b3a5-ccfda0a81ecc.](https://cellxgene.cziscience.com/collections/a0c84e3f-a5ca-4481-b3a5-ccfda0a81ecc) The MRCA is also hosted on the Baylor College of Medicine data portal at [https://mouseatlas.research.bcm.edu.](https://mouseatlas.research.bcm.edu) Additionally, access to the MRCA is provided on the UCSC Cell Browser at <https://retina.cells.ucsc.edu> and the Broad Single Cell Portal (SCP) at [https://](https://singlecell.broadinstitute.org/single_cell/study/SCP2560) singlecell.broadinstitute.org/single_cell/study/SCP2560. The SCP accession numbers for the three subclass atlases are SCP2557 for amacrine cells, SCP2558 for retinal ganglion cells, and SCP2559 for bipolar cells.
- Code Availability: All code used for the MRCA project can be found in the MRCA reproducibility GitHub repository ([https://github.](https://github.com/RCHENLAB/MouseRetinaAtlas_manuscript) [com/RCHENLAB/MouseRetinaAtlas_manuscript\)](https://github.com/RCHENLAB/MouseRetinaAtlas_manuscript). The pipeline to process the unpublished and collected public datasets is accessible at <https://github.com/lijinbio/cellqc>.
- Additional information: Any additional information required to reanalyse the data reported in this paper is available from the [lead con](#page-13-3)[tact](#page-13-3) upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

This study did not collect new samples.

METHOD DETAILS

Generation of scRNA-seq datasets of the mouse retina

We have generated 16 scRNA-seq samples of the mouse C57BL/6J retina [\(Table S1\)](#page-10-13). All mice were male. All procedures were approved by the Institutional Animal Care and Use Committee (IACUC) and followed the Association for Research in Vision and Ophthalmology (ARVO)

Statements for the Use of Animals in Ophthalmic and Vision Research, in addition to the guidelines for laboratory animal experiments (Institute of Laboratory Animal Resources, Public Health Service Policy on Humane Care and Use of Laboratory Animals). After dissection, retinas were dissociated into single cells using papain-based enzyme following the published protocol.^{[48](#page-11-28)} With activated 45U of papain (Worthington, Cat. #LS003126) solution (1mg L-Cystine, Sigma; 8 KU of DNase I, Affymetrix; in 5 mL DPBS), retina was incubated at 37C for ~20min, followed by the replacement of buffer with 2mL ovomucoid solution (15 mg ovomucoid, Worthington Biochemical; 15 mg BSA Thermo Fisher Scientific; in 10 mL DPBS) and 500ul deactivated FBS. Following the enzymatic digestion step, the retina tissues were carefully triturated and filtered using 20 µm plastic meshes. Trituration steps were repeated with additional 1mL ovomucoid solution until no tissue was visible. Singlecell suspension was spun down at 300g, 4C for 10 min and used in the next step.

To deplete the PR, cells were resuspended in 0.5% BSA and stained with CD73-PE antibody (MACS, Catalog: 130-102-616) for 10 min at 4C (for each million cells, add 98ul 0.5% BSA with 2ul CD73-PE antibody) and washed with 35 mL 0.5% BSA at 4C for 10min. After being stained with Anti-PE microbeads (MACS, Catalog: 130-105-639) (80µl 0.5% BSA and 20ul microbeads per each million cells) for 15 min at 4C, cells were washed and resuspended in 0.5% BSA. CD73 negative neuronal cells were enriched by autoMACS Pro Separator (Miltenyi Biotec) DEPLETES mode. Similarly, CD90.1 positive neuronal cells were enriched with CD90.1 microbeads (MACS, LOT: 130-094-523; 90ul 0.5% BSA and 10ul CD90.1 microbeads per each million cells) and autoMACS POSSEL-S mode. Cells viability was 87%–94% when checked using DAPI staining under microscope.

Guided by 10X manufacturer's protocols [\(https://www.10xgenomics.com\)](https://www.10xgenomics.com), single-cell cDNA library was prepared and sequenced. Briefly, single-cell suspension was loaded on a Chromium controller to obtain single cell GEMS (Gel Beads-In-Emulsions) for the reaction. The library was prepared with Chromium Next GEM single cell 3' kit V2 (10X Genomics) and sequenced on Illumina Novaseq 6000 [\(https://www.illumina.](https://www.illumina.com) [com](https://www.illumina.com)). Our newly generated single cell data were sequenced at the Single Cell Genomics Core at Baylor College of Medicine.

QUANTIFICATION AND STATISTICAL ANALYSIS

Data collection and preprocessing of the mouse retinal scRNA-seq

To recover high-quality cells, data samples were processed through a quality control pipeline [\(https://github.com/lijinbio/cellqc\)](https://github.com/lijinbio/cellqc). In brief, raw sequencing reads of 10x Genomics were first analyzed by the 10x Genomics Cell Ranger pipeline (version 7.0.1)^{[40](#page-11-20)} using the mm10 genome reference obtained from 10x Genomics [\(https://cf.10xgenomics.com/supp/cell-exp/refdata-gex-mm10-2020-A.tar.gz](https://cf.10xgenomics.com/supp/cell-exp/refdata-gex-mm10-2020-A.tar.gz)). Potential empty droplets in the filtered feature count matrices were further detected by dropkick.⁴⁷ Background transcripts contamination in the retained true cells were eliminated using SoupX.⁴⁵ DoubletFinder then was utilized to estimate and exclude potential doublets with high proportions of simulated artificial doublets.⁴³ In the resulting singlets, we extracted high-feature cells that contain ≥ 300 features, ≥ 500 transcript counts, and \leq 10% of reads mapped to mitochondrial genes.

In addition to our own data, we have incorporated well-characterized public datasets. Specifically, we have integrated cell-type-enhanced profiling data for ACs (accession: GSE14[9](#page-10-6)715),¹² bipolar cells (accession: GSE81904),¹¹ retinal ganglion cells (accession: GSE133382⁹ and GSE254587^{[15](#page-10-11)}), and non-neuronal retinal cells (accession: GSE199317).¹⁴ Furthermore, we have included four samples from wild-type mice were also collected from GSE201254 to account for retinal ganglion cells.¹³ To account for non-neuronal retinal cells, nine control samples were collected from GSE199317.^{[14](#page-10-18)} These cell-type specific single-cell datasets form the basis for subclass clustering in our mouse retina reference. To generate the updated transcriptome measurement of the GSE81904 from Shekhar et al., which was derived from the Drop-seq protocol, we applied the Drop-seq pipeline using the source code available at <https://github.com/broadinstitute/Drop-seq>. To ensure consistent gene feature annotation with the Cell Ranger pipeline, we used the gene annotation GTF file from the 10x Genomics mm10 genome reference package during the alignment of Drop-seq reads. In addition, GSE149715, GSE133382, GSE201254, and GSE199317 were also processed from scratch using raw sequencing reads using the 10x Genomics Cell Ranger pipeline (version 7.0.1).^{[40](#page-11-20)} To incorporate the high-quality cell type annotation of six public datasets, released count matrices and cell labeling were downloaded for meta-analysis. To further eliminate potential multiples in the integrated analysis, Solo doublet detection algorithm was used to identify potential multiples.

Data integration of scRNA-seq datasets

To eliminate technical variations in samples derived from different studies and experiments, 52 samples were integrated to remove the batch effect by scVI.²⁰ scVI explicitly formulates the batch effect as a latent variable in the deep generative model of observed expressions. Normalized expression was applied to detect highly variable genes (HVGs) using the Seurat algorithm (flavor: seurat). The ''sampleID'' was used as the batch key for calculating HVGs and the batch variable in the scVI modeling. The scVI model utilized 2 hidden layers (n_layers: 2) and a 30-dimensional latent space (n_latent: 30). The trained low-dimensional representation was used for cluster detection with the Leiden algo-rithm.⁴⁹ UMAP of low-dimensional visualization was generated by the Scanpy package.^{[41](#page-11-21)}

Cell clustering and cell type annotation

To annotate major classes of cell clusters, we incorporated well-annotated cell labels released from public datasets, i.e., Yan et al. for ACs, Shekhar et al. for BCs, and Tran et al. and Jacobi et al. for RGCs. Cells from Yan et al. were annotated into 63 AC types. Cells from Shekhar et al. were 15 BC types showing in 14 clusters with small numbers of cells annotated as ACs, rod, and cone. Tran et al. cells were identified as 45 RGC types. The cell type labels of these well-annotated cells are used to annotate integrated cell clusters. To annotate isolated cell clusters that

were isolated from existing cell labels of the public datasets, cluster-specific markers were examined from the top ranked genes generated by the Wilcoxon rank-sum test using the rank genes groups() function in the Scanpy package.⁴¹

To annotate subclass BC, AC, and RGC, subclass-specific cells were isolated and integrated using scVI. The generated low-dimensional embeddings were used to detect clusters using the Leiden algorithm. The parameter ''resolution'' of the Leiden algorithm affects the number of resulting clusters, and a typical resolution of 1.0 tends to cause over-clustering, where multiple clusters are identified within a compact cluster due to a high resolution. Therefore, it is necessary to test various resolutions when using the Leiden algorithm on specific atlases. Additionally, a single resolution fails to capture the hierarchical nature of similarity among cell clusters in complex retina cell classes, such as BC, AC, and RGC. Hence, to determine the optimal number of clusters for subclasses, a two-level clustering approach was applied. In the first level of clustering, various resolutions (e.g., [0.1, 1.0]) were tested for the Leiden algorithm. A resolution was selected where over-clustering was not observed in UMAP visualization, while some under-clustered clusters were deemed acceptable. Those under-clustered clusters from the firstlevel clustering would be refined into smaller clusters by testing various resolutions in the second-level clustering. In the second-level clustering, cells and low-dimensional embeddings from each cluster identified in the first-level clustering were extracted, and the extracted low-dimensional embeddings were used in the Leiden algorithm. Again, various resolutions (e.g., [0.01, 0.09]) were tested, and a resolution was selected to achieve clustering without over-clustering on UMAP. Consequently, the two-level clustering approach achieved an optimal number of clusters for BCs, ACs, and RGCs.

To construct the BC atlas, data samples for BCs were integrated using scVI. Initially, 33 clusters were identified, of which 30 could be matched and merged to individual BC types by examining previously generated cell labels and their known marker gene expression,^{[11](#page-10-8),[24](#page-11-4)} while the remaining 3 clusters (C30, C31, and C32) were excluded from the analysis as they contained non-BCs from previous annotation labels or had high UMI counts [\(Figures 2A](#page-4-0) and [S2](#page-10-13)A–S2C). Consequently, 15 BC types were identified and annotated.

To construct the AC atlas, the data integration analysis for ACs using scVI identified a total of 71 clusters, of which 62 clusters could be matched and merged to 49 individual AC types via previous annotation labels and known marker expression. However, 8 clusters were over-clustered that contained two or more previous AC type labels, and one cluster (C70) was excluded from the AC reference due to non-AC cells [\(Figures S4](#page-10-13)C and S4D). To further address the 8 remaining over-clustered clusters ([Figure S5](#page-10-13)A), we utilized a two-level annotation approach. This involved isolating cells from each cluster and refining the clustering. The two-level annotation allowed the separation of the remaining 14 types: AC11, AC16, AC29, AC42, AC47, AC50, AC53, AC54, AC55, AC56, AC60, AC61, AC62, and AC63 [\(Figures S5](#page-10-13)A–S5C). This revealed clusters that primarily consisted of RGCs, which have been removed in the integrated AC map [\(Figure S5C](#page-10-13)). As a result, 63 AC types were identified and annotated.

Three AC types, AC16, AC53, and AC62, were identified as dual types expressing both canonical GABAergic and glycinergic receptors in the study by Yan et al. AC16, however, was shown as a suspected doublet in their study, alongside AC60. To examine the transcriptomic similarities between the two clusters, AC16 and AC60, a hierarchical clustering tree was generated using the phylogram R package $\rm{^{50}}$ based on the average normalized expression of clusters. AC16 and AC60 showed proximity in the hierarchical clustering ([Figure S7](#page-10-13)) with relatively high UMI counts [\(Figure S6](#page-10-13)A). Although the AC53 cluster was resolved in the second-level annotation, the loose clustering of AC53 cells is quite apparent. The third dual type, AC62, was also under-clustered and merged with AC42 and AC55. While AC62 was resolved in the secondlevel annotation, it also appears near its neighboring cluster, AC42, in the hierarchical clustering ([Figure S7\)](#page-10-13). With very few cells being annotated as dual types in CD90.1 and Ma et al. samples, which express high levels of Thy1 (data not shown), further validations of the dual types are required.

To construct the RGC atlas in the MRCA, the integrated analysis identified 54 clusters with an even distribution of cells from different data sources in most clusters [\(Figures S8](#page-10-13)A–S8D). Out of these clusters, 48 can be mapped and merged into 39 individual RGC types previously identified using marker gene expression and previous annotation labels [\(Figures S8A](#page-10-13) and S8B), while five clusters were over-clustered that contained multiple previous RGC types, and one cluster (C8) contained a mixture of several RGC type labels with high UMIs and was excluded from the downstream analysis as multiplets. To annotate the remaining seven types found in the five clusters with multiple labels, the secondlevel annotation was performed, which resulted in a clear separation of all 46 previously identified RGC types [\(Figures 5](#page-7-0)A and [S9](#page-10-13)A–S9C).

Differentially expressed gene analysis

To identify genes that are differentially expressed between cell types, we generated pseudo-bulk transcriptome of each annotated cell type in individual sample id. We used pyDESEQ2^{[51](#page-11-31)} to compare two clusters or types using the Wald test and identified genes specifically expressed in each cluster or type. Differentially expressed genes are identified under q-value <0.05. The Wald statistics (log2FoldChange divided by lfcSE) was used to rank and select the top 10 genes expressed in each type.