Analysis of gene network bifurcation during optic cup morphogenesis in zebrafish

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Abstract:

Sight depends on the tight cooperation between photoreceptors and pigmented cells, an ancestral tandem in metazoans. In vertebrates, both cell types derive from common progenitors in which a single gene regulatory network (GRN) bifurcates into the neural retina (NR) and retinal-pigmented epithelium (RPE) specification programs. Classical genetic studies have identified the main upstream nodes controlling these networks; however, their detailed architecture and cis-regulatory logic remain poorly investigated. Here, we have characterized transcriptome dynamics (RNA-seq) and chromatin accessibility (ATAC-seq) in segregating NR/RPE populations in zebrafish. Analysis of differentially active cis-regulatory modules and enriched transcription factor (TF) motives suggest extensive network redundancy and contextdependent TF activity. Downstream targets identification highlights an early recruitment of desmosomal genes in the flattening RPE cells, revealing Tead factors as their upstream regulators. Dynamic investigation of GRNs uncovers an unexpected sequence of TF recruitment during RPE specification, which is conserved in humans. This systematic interrogation of the NR/RPE bifurcating networks should improve both counselling of congenital eye disorders and hiPSCs-to-RPE differentiation protocols for cell-replacement therapies applicable to retinal degenerative diseases.

Key-words: Eye development, Gene regulatory network (GRN), cis-regulatory modules, Retinal degenerative disease, neural retina (NR), retinal pigmented epithelium (RPE).

Running title: Retinal network bifurcation in zebrafish

Introduction:

Darwin in *The Origin of Species* already hinted to the possibility of a single photoreceptor escorted by a pigmented cell, as the ancestral structure to all animal eyes. Several authors have further elaborated this notion during the last decades (Arendt & Wittbrodt, 2001, Gehring, 2014, Vopalensky & Kozmik, 2009). An intimate association between photoreceptors and pigmented ancillary cells is found in the eyes of all metazoan groups (Land & Nilsson, 2002), even in cnidarians (Kozmik, Ruzickova et al., 2008). Thus, the origin of this cellular tandem can be traced back to a common ancestor dating to at least 650 mya. Such remarkable evolutionary conservation can be explained by the fact that vision depends on the close physical and physiological interaction between the two cell types (Strauss, 2005). Photoreceptors, with a very active metabolism and a constant exposure to light, are particularly prone to degeneration due to DNA damage and oxidative stress. Pigmented cells protect photoreceptors by maintaining their homeostasis through growth factors secretion, visual pigments recycling and outer-segments phagocytosis (Letelier, Bovolenta et al., 2017).

Fate map experiments in vertebrates have shown that the precursors of the neural retina (NR) and retinal pigmented epithelium (RPE) derive from the undifferentiated eye field (Li, Joseph et al., 2000). In the permanently growing retinas of teleost fish, stem cells located at the ciliary marginal zone are capable of producing both cell types (Tang, Gao et al., 2017). Furthermore, in tetrapods retinal precursors retain certain potential to trans-differentiate from the pigmented to the neural retina identity (Coulombre & Coulombre, 1965, Pittack, Jones et al., 1991), and *vice versa* (Rowan, Chen et al., 2004).

In zebrafish, the RPE and NR presumptive domains start differentiating at the optic vesicle stage from the medial (ML) and lateral (LL) epithelial layers, respectively (Li et al 2000; Kwan et al 2012). The specification of both retinal domains occurs simultaneously to the folding of the vesicle into a bi-layered cup, and entails profound cell shape changes in each domain. Precursors at the LL elongate along their apico-basal axis, differentiate as NR, and constrict basally to direct the folding of the retinal neuroepithelium (Ivanovitch, Cavodeassi et al., 2013, Nicolas-Perez, Kuchling et al., 2016). In contrast, precursors at the ML either acquire a squamous epithelial shape and differentiate as RPE, or flow into the LL to contribute to the NR domain (Heermann, Schutz et al., 2015, Moreno-Marmol, Cavodeassi et al., 2018, Picker, Cavodeassi et al., 2009, Sidhaye & Norden, 2017). In other vertebrates, the specification of the NR/RPE domains and the morphogenesis of the optic cup follow a pattern similar to that of the zebrafish, though the relative weight of the different morphogenetic mechanisms may vary among species (Martinez-Morales, Cavodeassi et al., 2017).

During the last decades numerous studies have investigated how the RPE and NR domains get genetically specified (Fuhrmann, 2010). Eye identity is established in the anterior neural plate by the early activation of a gene regulatory network (GRN) pivoting on a few key regulators collectively known as eye field transcription factors (EFTF): including Lhx2, Otx2, Pax6, Rx, or Six3 (Zuber, Gestri et al., 2003). Upon the influence of inductive signals derived from the lens epithelium (FGFs) or the extraocular mesenchyme (Wnts and BMPs), the eye field network branches into the mutually exclusive developmental programs of the NR and RPE (Cardozo, Almuedo-Castillo et al., 2020, Fuhrmann, 2010). Classical genetic experiments performed mainly in mice have identified key nodes of the NR and RPE specification networks. The TF-encoding gene *Vsx2* (previously known as *Chx10*) was identified as the earliest determination gene differentially expressed in NR but not in RPE precursors (Liu, Chen et al., 1994). *Vxs2*

plays an essential role in specifying the NR domain by restraining the RPE identity through direct repression of the TF Mitf (Bharti, Liu et al., 2008, Horsford, Nguyen et al., 2005, Rowan et al., 2004, Zou & Levine, 2012). Other homeobox regulators, many inherited from the eye field specification network, contribute to the network of NR specifiers. This list includes *Lhx2*, *Sox2*, *Rx*, *Six3* and *Six6* genes, which are required either for NR maintenance or for suppressing RPE identity (Roy, de Melo et al., 2013, Wang, Yasugi et al., 2016). The establishment of the RPE network depends instead on the cooperative activity between Mitf and Otx factors (Lane & Lister, 2012, Martinez-Morales, Dolez et al., 2003). Classical experiments in mice showed that inactivation of either *Mitf* or *Otx* genes results in an RPE differentiation failure, with the alternative acquisition of NR molecular and morphological features (Martinez-Morales, Signore et al., 2001, Mochii, Ono et al., 1998, Nguyen & Arnheiter, 2000). More recently, mutation of the main effectors of the Hippo pathway, *Yap* and *Taz*, showed the essential role of these coregulators in the differentiation of the RPE lineage (Kim, Park et al., 2016, Miesfeld, Gestri et al., 2015).

Despite the identification of these key regulators, the architecture of the NR and RPE specification subnetworks is far from being well understood (Martinez-Morales, 2016). Systematic attempts to reconstruct retinal GRNs using next-generation sequencing (NGS) methods have focused mainly in the differentiation of neuronal types at later stages of development or in epigenetic changes linked to retinal degeneration (Aldiri, Xu et al., 2017, Buono & Martinez-Morales, 2020, Wang, Zibetti et al., 2018). Very recently, scRNA-seq has been proved a useful tool to characterize cell heterogeneity and infer differentiation trajectories in human tissues and retinal organoids (Collin, Queen et al., 2019, Hu, Wang et al., 2019). However, a comprehensive analysis of the NR and RPE bifurcating networks would require a deeper sequencing coverage approach, while dealing with the limited cell population size at optic cup stages.

In this study, we dealt with previous limitations by applying a combination of RNA-seq and ATAC-seq on sorted NR and RPE populations. Using the zebrafish retina as a model system, we followed transcriptome dynamics and chromatin accessibility changes in both cell populations, as they depart from a common pool of progenitors. The cross-correlation of RNA-seq and ATAC-seq data allowed us to identify activating and repressing cis-regulatory elements (CREs), discover enriched transcription factor (TF) binding motifs, and unveil relevant downstream targets for the main specifiers. Our analyses confirmed previously known TFs as central nodes of the eye GRNs, and more importantly, provide information on their recruitment sequence both in zebrafish and human cells.

Results:

1.- Analysis of specification networks in isolated NR and RPE precursors.

To examine the bifurcation of the regulatory networks specifying the NR and RPE domains in zebrafish, we focused on the developmental window comprised between stages 16 hpf and 23 hpf (Fig 1 A). Within this period, the optic vesicle transits from a flattened disk in which the lateral and medial layers are still similar in terms of cell shape and volume, to an optic cup stage in which the NR and RPE cells are morphologically differentiated: bottle-shaped for retinal neuroblasts and flat for RPE precursors (Li et al., 2000). It is important to state that at the end of this window (23 hpf), neurogenesis have not started yet in the retina, neither the RPE display pigmentation (Masai, Stemple et al., 2000). We used the transgenic lines E1 bHLHe40:GFP and vsx2.2:GFP to isolate the NR and RPE populations at 18 and 23 hpf by FACS (see methods; Fig 1B). Both vsx2 and bhlhe40 are among the earliest markers reported in zebrafish for the neural retina and RPE domains respectively (Barabino, Spada et al., 1997, Cechmanek & McFarlane, 2017). To isolate early precursors at stage 16 hpf we took advantage of the fact that the vsx2.2:GFP transgene is transiently expressed in all progenitors and gets restricted to the NR as progenitors of the medial layer start expressing bHLHe40 (Nicolas-Perez et al., 2016); Moreno-Marmol et al., 2020). Populations isolated by flow cytometry were analysed by RNA-seq, at all stages, and by ATAC-seq at 23 hpf (Fig 1A).

The transcriptomic analysis of the different cell populations highlighted thousands of differentially expressed genes (DEGs) throughout development and between domains (Appendix Fig S1A,B; Dataset EV1). A multidimensional analysis of these changes revealed that an extensive divergence between the NR and RPE transcriptomes already occurs within the first two hours (16 to 18 hpf) of optic vesicle folding (Fig 1C). Later on, between 18 hpf and 23 hpf, the transcriptome divergence between the NR and RPE progresses, although is more modest within each domain (Fig 1C, Appendix Fig S1B).

2.-Chromatin accessibility in NR vs RPE precursors identifies cis-regulatory modules.

To gain further insight into the architecture of the optic cup GRNs, we sought to identify differentially open chromatin regions in the morphologically divergent NR and RPE domains. To this end, we performed ATAC-seq experiments from isolated NR and RPE precursors at 23 hpf. A total of 238369 open chromatin regions (OCRs) were detected using this approach. After statistical analysis, a fraction of the peaks was identified as differentially open chromatin regions (DOCRs), corresponding to putative cis-regulatory elements (CRE) that are more active in the NR than in the RPE or vice versa. Approximately 12.6% of all peaks (30172 peaks) were differentially open with an adjusted p-value < 0.05. This proportion was reduced to 4.8% when the adjusted p-value was lowered to <0.001 (Fig EV1A,B; Dataset EV2). Regardless the adjusted p-value, we observed a larger number of DOCRs in the RPE than in the NR (18909 vs 11263; adjusted p-value <0.05), and this was also accompanied by a higher average fold change in the RPE than in the NR associated DOCRs (Fig EV1B,C). Analysis of the distribution of open chromatin regions in the genome showed no evident differences between NR and RPE for the whole set of OCRs. However, there was a difference in average localization between the entire OCRs and the subsets of DOCRs in the genome. When compared to the distribution of all the OCRs a noticeable depletion of DOCRs located near the promoter regions was evident in both the NR and RPE domains. The proportion of NR peaks near the promoter was 11.91% for OCRs vs 4.6% for DOCRs; whereas for the RPE peaks the proportion was 10.03% for OCRs vs 3.09% for DOCRs (Fig EV1D), showing that domain specific cis-regulatory modules tend to occupy more distal positions in the genome. Gene ontology analysis of terms from the category Biological Process enriched in the list of genes associated with DOCRs yield results consistent with the analysed tissue. Genes associated with NR DOCRs were enriched in terms related to nervous system development, neuron differentiation, and eye morphogenesis, whereas genes associated with RPE DOCRs were enriched in terms such as melanocyte differentiation and epithelial differentiation (Appendix Fig S2).

Previous reports have shown the importance of correct DNA methylome patterning during human iPSC reprograming towards RPE (Araki, Miura et al., 2019), and have proved a requirement for active demethylation during eye formation in vertebrates (Xu, Xu et al., 2012). Thus, to further characterize the NR and RPE specific DOCRs, we next interrogated their DNA methylome profiles. We utilised base-resolution DNA methylome (WGBS) and hydroxymethylome (TAB-seq) data of zebrafish embryogenesis and adult tissues (Bogdanovic, Smits et al., 2016), to obtain insight into the temporal dynamics of DNA methylation (5mC) over these open chromatin regions. Supervised clustering (k=2) of 5mC patterns separates both datasets into two well-defined clusters (Fig EV2A,B). The first cluster consists of distal regulatory regions that initiate active demethylation at ~24hpf, as demonstrated by strong 5hydroxymethylcytosine (5hmC) enrichment and the progressive loss of 5mC starting at 24hpf, and then are most visible in the adult brain. Importantly, this cluster is highly enriched in H3K27ac, an active enhancer mark, and depleted of H3K4me3, which indicates their identity as bona fide enhancers (Bogdanovic, Fernandez-Minan et al., 2012). The second cluster of both NR and RPE datasets is hypomethylated, depleted of hmC, and enriched in the promoter mark H3K4me3, in line with a CpG island promoters character. To test whether active, tet-protein dependent DNA demethylation is required for the establishment of open chromatin structure at NR and RPE ATAC-seq peaks, we interrogated chromatin accessibility in wild type and triple tet (tet1/2/3) morphants in whole 24hpf embryos (Bogdanovic et al., 2016). Data from two independent MO knockdown experiments revealed that their chromatin accessibility signature is significantly (Mann-Whitney-Wilcoxon Test, P < 2.2e-16) reduced for NR ATAC-seq peaks in tet morphants when compared to wild type embryos, even when whole embryos were assessed (Fig EV2C). This decrease was not visible in RPE peaks, likely because RPE enhancers are active in a much smaller cell population and their chromatin mark is masked by the en masse approach (Fig EV2D).

3.- Cross-correlation of RNA-seq and ATAC-seq data reveals activating and repressing CREs.

Integration of ATAC-seq and RNA-seq data has been used as a powerful tool to explore the architecture of developmental GRNs (Buono & Martinez-Morales, 2020, Lowe, Cuomo et al., 2019). By intersecting RNA-seq and ATAC-seq datasets, we observed a substantial overlap between DEGs and genes associated with DOCRs at 23 hpf. Thus, 47% of all DEGs are associated with at least one DOCR (Fig 1D). Specific examples of ATAC-seq and RNA-seq tracks are shown for the *vsx1* and *mitfa* loci, NR and RPE respective markers (Fig 1E,F). The cross-analysis of ATAC-seq and RNA-seq data allowed us to classify DOCRs (putative CREs) in two groups: those correlating with up regulated genes, here termed "activating CREs" and those correlating and repressing peaks are provided for the *six3a* and *otx2* loci (Fig EV3A,B). When

their average distance to the closest Transcription Start Site (TSS) was examined, we observed a significant trend: both for NR and RPE peaks, activating CREs tend to occupy positions closer to the TSS than the repressing regions (Fig EV3C). A possible explanation for this difference is the positive bias introduced by proximal promoters, which are typically enriched in binding sites for activators (Maston, Evans et al., 2006). To gain insight into the regulatory logic of the NR and RPE domains, we calculated the number of associated activating and repressing CREs for both the whole DEGs and the subset of differentially expressed TFs. Notably, as expected from their regulatory complexity, the average number of CREs per gene (either activating or repressing) was consistently higher for TFs than for the whole collection of DEGs (Fig 2C). More importantly, in the neural retina TFs associated with activating CREs outnumbered TFs associated with repressing regions (182 vs 69 respectively), whereas the opposite was observed in the RPE. Indeed, in this tissue we detected a robust repressive cis-regulatory logic, with 173 TFs associated with at least one repressing CREs and only 110 associated with activating ones (Fig 2B,C). This opposite trend, better appreciated in histogram graphs, suggests that the NR specification program is sustained mainly by the activation of transcriptional regulators, whereas RPE determination seems to require primarily the repression of the NR-specific-TFs (Fig 2D,E).

4.- Distinctive sets of TFs and cytoskeletal components are progressively recruited during NR and RPE specification.

To efficiently analyse expression dynamics, as the NR and RPE networks bifurcate, we used a gene clustering approach. We applied both hierarchical and partitioning soft clustering for the classification of genes encoding for TFs or cytoskeletal components. We focused on these two gene categories to examine not only transcriptional lineage specifiers, but also terminal effectors operating in the divergent cell shape changes observed between the NR and RPE domains. Using partitioning soft clustering we established 25 groups of gene expression variation from progenitors towards NR or RPE, for both TFs and cytoskeleton components (Appendix Fig S3A,B; Datasets EV3 and EV4). This approach identified several expression clusters, which are distinctive for each domain and developmental stage: e.g. clusters 1, 13, and 23 for 16hpf progenitors in the TFs category (Appendix Fig S3A); or 6, 11, and 21 for 23hpf RPE precursors in the cytoskeletal components' analysis (Appendix Fig S3B). Using a hierarchical clustering approach, we could aggregate the small sub-groups into 6 main clusters for TFs and 8 for cytoskeletal components, all of them linked mainly to a specific domain and/or developmental stage (Fig 3; Fig EV4; Datasets EV5 and EV6). To define precise time windows of expression and to infer regulators order of action, we focused our attention on the identity of TFs belonging to the different large clusters. Significant TFs in cluster #5, corresponding to 16hpf progenitors, include rx3, an early eye specifier with a known role in optic vesicle evagination (Loosli, Winkler et al., 2001, Mathers, Grinberg et al., 1997, Rembold, Loosli et al., 2006); and her factors required to maintain the progenitor neural state (Chapouton, Webb et al., 2011) (Fig 3B). As expected, Cluster #1, corresponding to NR 23 hpf precursors, contains many of the acknowledged retinal specifiers, such as rx1, rx2, sox2, six3a, six3b, six6b, vsx1, vsx2, hmx1, hmx4 and lhx2b (Fuhrmann, 2010), which already increased their expression at 18 hpf (Fig 3B). Surprisingly, known RPE specification genes such as otx2 and mitfa (Martinez-Morales, Rodrigo et al., 2004), included in cluster #4, do not increase their levels significantly in RPE precursors at 18hpf, peaking only later at 23 hpf (Fig 3B). In contrast, TFs included in cluster #3, such as *tead3b*, *tfap2a* and *tfap2c* started to rise, when not peaking, in the RPE at 18 hpf. Notably, some of these TFs, including *tcf12*, *smad6b*, and especially *vgll2b* not only peak at 18 hpf, but also rapidly decrease at 23 hpf (Fig 3B). This observation suggests the existence of two waves of TFs regulating the identity of the RPE domain.

Next, we focused on the content of the hierarchical clusters associated to cytoskeletal components (Fig EV4A). While the analysis of the cytoskeletal genes upregulated in PG returned no significant enrichment for terms annotated in the Cellular Components, similar analysis for cytoskeletal genes in the NR clusters yielded significantly enriched terms associated to microtubules and centrosomes (Fig EV4B). This observation is in line with the elongation of the apico-basal axis and the polarization of the microtubules reported for NR precursors in zebrafish (Ivanovitch et al., 2013). In contrast, the only enriched term in the RPE cytoskeletal clusters was "intermediate filaments" (Fig EV4B), consistent with the RPE acquiring a squamous epithelial character.

5.- Motif enrichment analysis suggests redundancy and cooperativity in the NR and RPE networks.

To further explore the regulatory logic of the NR and RPE gene networks, we investigated overrepresented motifs within the DOCRs associated with each domain. Motif enrichment analysis of the differentially open regions in the NR identified a highly significant overrepresentation of the homeobox and sox TF binding motifs (Fig 4A; Dataset EV7). The core homeobox binding motif (5'- TAATT-3') is shared by TFs from the homeodomain K50 PRD-class; such as vsx1, vsx2, rx1, rx2, and rx3; LIM-class, such as lhx2b; and NKL class; such as hmx1 and hmx4 (Fig 4A). All of them are well-known retinal specifiers contained in group #1 of our hierarchical clustering analysis (NR 23 hpf). As they are co-expressed in the retina, it is likely that they cooperate to target a partially overlapping set of cis-regulatory modules. To explore this possibility, we retrieved the individual position weight matrixes (PMW) associated to both homeobox and sox TFs from available databases and used this information to explore their functional synergy in the retinal DOCRs (see Method sections). Two different approaches were used: (i) calculating the co-occurrence rate of binding sites for two different TFs in the same CRE (co-occupancy); and (ii) estimating the percentage of binding sites for TFs located in different CREs but still associated with the same gene (co-regulation) (Fig 4B,C). These analyses predicted an extremely high degree of inter-connectivity within the NR network. Thus, the average co-occurrence rate of two different theoretical binding sites in the same peak (cooccupancy) was 25.6%. This combinatorial activity becomes more pronounced when examining the binding sites in different CREs associated to the same gene (co-regulation), which was in average 46.6%. Indeed, when individual TF binding sites were investigated, the number of co-occupied peaks or co-regulated genes was in general much larger than those instances in which the binding site was found isolated (Fig 4D,E). Even when these are predicted binding sites, our analyses suggest a strong cooperative activity of homeobox and sox factors in driving the NR developmental program.

In the RPE, a motif enrichment analysis of the specific DOCRs also revealed a number of binding motifs that correspond to the TFs identified as up-regulated in the RNA-seq analysis (Fig 5 A; Dataset EV8). We found a very significant enrichment for tfap2a and tfap2c binding motifs, in agreement with the early expression of these factors in the RPE. This considerable overrepresentation points to a prominent position of these factors within the regulatory

hierarchy, which is in line with their role in the specification of the pigmented tissue (Bassett, Williams et al., 2010). In addition, a significant enrichment for bHLH, tead and otx2 binding motifs was also observed (Fig 5A). Importantly, the core bHLH motif (5'-CACGTG-3') is shared by members of the family expressed in RPE, such as mitfa, bHLHE40, bHLHE41, and tfec (Fig 5A). As already investigated for the NR TFs, we examined the potential cooperation among these RPE specifiers (Fig 5B-E). Interestingly, whereas the average rate of co-regulated genes in the RPE did not differ much from that found in NR (48% vs 46.6%; Fig 5E), the rate of cooccupied peaks dropped from 25.6% in the NR to 11.7% in the RPE (Fig 5D). This observation suggests that the RPE network is less dependent on TFs cooperativity within the same CRE than the NR network. In such regulatory scenario, each TF may trigger transcriptional subprograms within a broader developmental network. To explore this possibility, we assessed GO enrichment for the genes associated with RPE DOCRs containing the different TFBS and then we grouped the results using hierarchical clustering (Appendix Fig S4). The outcome of this clustering approach supported a branched regulatory scenario. Thus, whereas some TFs (i.e. tcf12, tfap2c, otx2) appeared associated to many GO terms, others were more specific: such is the case of *mitfa* that was associated only to pigment cell differentiation (Appendix Fig S4).

Finally, to investigate the regulatory logic of activating and repressing CREs in each domain, we scanned these regions for enriched motifs separately. Strikingly, those enriched motifs ranking higher for the NR activating CREs, such as the homeobox and sox TF binding sites, were also ranking higher for the repressing regions in this tissue (Fig 6A; Dataset EV9). A very similar scenario was observed for the RPE peaks, which regardless being activating or repressing regions showed tfap2c, tfap2a, bHLH, and otx2 consensus binding sites as top enriched motifs (Fig 6B; Dataset EV9). When GO terms associated with genes linked to these regions were examined, we observed terms linked to eye morphogenesis enriched for both NR activating and RPE repressing regions (Fig 6). This finding reveals context-dependent TFs activity, and points to a common set of genes activated in the NR and repressed in the RPE by antagonistic GRNs. In fact, a detailed analysis of the gene lists associated to NR activating and RPE repressing regions showed that many of the retinal specifiers themselves (including hmx4, *lhx9, mab21l1, nr2f2, pax6a, pax6b, rx2, six3a,* or *sox2*) are under the antagonistic regulation of the NR and RPE GRNs (Dataset EV10). In addition, the analysis of GO terms enrichment also suggested domain-specific functions for the NR and RPE GRNs. Thus, NR repressing CREs are associated to genes involved in mesoderm formation, whereas RPE activating regions are link with epidermal differentiation genes.

6.- Desmosomal components are activated during RPE specification.

The integration of our ATAC-seq and RNA-seq data allows formulating and testing different hypotheses related to genetic programs controlling the specification of retinal tissues. As an example, we followed up the observation that genes encoding for intermediate filaments are enriched in the RPE domain (Fig EV4B). Keratin looping into desmosomal plaques plays a fundamental role in maintaining tissue architecture under mechanical load (Hatzfeld, Keil et al., 2017) (Fig 7A). We thus investigated the transcriptional profile of other desmosomal components as the RPE and NR networks diverge. A detailed analysis revealed that not only keratin genes, but also many desmosomal genes such as *dspa*, *evpla*, *pleca*, or *ppl*, are among the most upregulated genes (i.e. highest fold change) in committed RPE cells (Fig 7B). Many of these genes increased their expression already at 18 hpf, when RPE cells start to differentiate

morphologically, but hours before the tissue acquires pigmentation (Fig 7B). To gain insight into the regulation of these cytoskeletal components, we performed a motif enrichment analysis of the subset of DOCRs associated with keratins and other desmosomal genes upregulated in the RPE (Dataset EV11). This analysis revealed Tead motifs as top ranked overrepresented binding sites in the CREs regulating desmosomal genes (Fig 7C). When compared to that of the entire set of DOCRs, the average number of motifs per peak was 4.5 fold higher for those CREs specifically associated with components of the desmosome machinery (1.35 vs 0.3). This observation suggests a key role for Tead family proteins in the regulation of the intermediate filament cytoskeleton. To test this hypothesis, we took advantage of available zebrafish double mutants for the main Tead coactivators yap and taz, which have been reported to display RPE differentiation defects (Miesfeld et al., 2015). In agreement with our observations, the expression of keratins (krt4 and krt8) as well as pigmentation genes (*tyr* and *tyrp1b*) were significantly reduced in *yap -/- taz -/-* double mutant head tissue at 18 hpf as determine by RT-qPCR (Fig 7D). As a control, the expression of the NR markers vsx2 and six3q was unaffected in the mutant tissue. These results further support a role for Tead factors in the transcriptional regulation of desmosomal genes at the RPE.

The high efficiency of the CRISPR/Cas9 technology allows performing F0 mutagenesis screens in zebrafish (Shankaran, Dahlem et al., 2017). We took advantage of this technology to test functionally some of the genes identified in our analyses. A total of 21 genes, selected on the bases of their expression profile, CRE composition and/or dynamics, associated GO terms, and number of paralogues, were investigated in an F0 pilot screen (Fig EV5). A substantial proportion of the tested genes (i.e. 12 out of 21 sgRNA combinations tested) displayed eye malformations, such as microphthalmia, eye fissure closure defects and/or hypopigmentation, thus confirming that many of the candidate genes play an essential role as components of the eye GRNs (Fig EV5, Dataset EV12). Interestingly, among the sgRNA combinations tested, the injection of those directed against the structural desmosomal components *dspa/b*, the Tead regulators *vgll2a/b*, and particularly the cocktail *tead1a/b* + *tead3a/b* resulted in a significant proportion (30.4%, 40.6%, and 80% respectively) of the F0 embryos displaying reduced eye size and severe hypopigmentation (Fig 7E; Fig EV5). These results further indicate that both desmosomal assembly and Tead activity are required during RPE differentiation.

7.- Gene expression analysis during hiPSCs-to-RPE differentiation

A second important finding derived from our gene expression clustering analysis was the identification of two waves of transcriptional regulators during the specification of the RPE in zebrafish. Understanding this TF recruitment sequence in humans may have important basic and translational applications. We thus asked whether the same *consecutio temporum* identified in zebrafish was conserved in human iPSCs differentiating to RPE. During the first four weeks of differentiation in culture from hiPSCs (see methods), pluripotent stem cells progressively change their morphology to a cobblestone appearance, acquiring a light pigmentation at the end of the fourth week (Fig 8A). Using RT-qPCR, we tested mRNA levels for a total of 26 RPE genes including genes encoding for stemness markers (*NANOG* and *OCT4*); mature RPE markers (i.e. *CRALBP, RPE65* and *TYR*); known RPE specifiers activated in the second wave of gene expression (i.e. *BHLHE40, MITF, OTX2,* and *TFEC*); desmosomal components (i.e. *DSP, EVPL, KRT4, KRT5, KRT8*); and TFs and signalling molecules activated in the first wave of gene expression (NOTCH1, NOTCH2, NOTCH3, SMAD6, TCF12, TEAD1, TEAD2,

TEAD3, TEAD4, TFAP2A, TFAP2C, and VGLL2) (Fig 8B; Dataset EV13). This analysis showed that most of the human genes orthologous to the zebrafish early specifiers cluster together according to their expression profiles, and reach maximal expression within the first weeks of culture. This was the case for *SMAD6*, *TEAD1*, *TEAD3*, *TFAP2A*, *TFAP2C*, and particularly for *VGLL2 and TEAD2*, the expression of which peaked very transiently. Similarly, the human orthologs of most of the genes that in zebrafish were identified as activated in a second transcriptional wave (including *MITF*, *BHLHE40*, *TFEC*, *CRALBP*, *RPE65* and *TYR*) also clustered together reaching a maximum expression level in the fourth week of culture. These findings indicated that despite the very different time scales of the developmental programs in these far-related vertebrate species, hours in zebrafish and days in humans, their RPE regulatory networks share a common logic of two-waves in the TFs recruitment sequence.

Discussion:

A system level approach to explore GRNs architecture. Cell differentiation, as classically illustrated by Waddington's epigenetic landscapes (Waddington, 1966), often entails sequential binary decisions that progressively restrict cell competence. At a genetic level this translates in the branching of "mother" gene regulatory networks into child sub-networks. Here we have used a double RNA-seg/ATAC-seg approach to zoom into the bifurcation of the NR/RPE developmental programs, which will give rise to specialized neurons and pigmented squamous cells respectively. The transient nature of this differentiation process, as well as the limiting size of the cell populations involved (Kwan, Otsuna et al., 2012), have hindered any systemic approach to investigate the architecture of the specification networks as they branch. The high sensitivity and deep sequencing coverage of the RNA-seq and ATAC-seq methods (Buenrostro, Giresi et al., 2013), combined with the isolation of NR and RPE precursors by FACS allowed us to overcome previous constrains. Our approach permitted not only the detection of transcriptomic variations and active cis-regulatory modules, but also helped to define hierarchical relationships among the core components of the network. In our analyses we have used a proximity method to associate genes and open chromatin regions. Previous studies have confirmed that this link to the adjacent gene is correct in 90% of the cases (Yoshida, Lareau et al., 2019). Therefore, although this estimate should be taken cautiously when individual cis-regulatory elements are examined, it is a valid approach when conclusions are driven from a genome-wide analysis. In this study we identified approximately 30000 chromatin regions differentially open (DOCRs) between the NR and RPE domains. The examination of their methylation status, as well as their overlap with H3K27ac and H3K4me3 marks indicates that most of these regions (≈95%) correspond to initially inactive enhancers that get progressively demethylated and activated at the phylotypic period, a behaviour reported for developmental genes in general (Bogdanovic et al., 2016). A much smaller cluster (\approx 5%) matches to hypomethylated constitutively active promoter regions (Figs EV1, EV2). It is therefore likely that most of the regions here identified as DOCRs correspond to cis-regulatory elements (CREs) active during the segregation of the NR/RPE networks.

The NR as a default program. Our data, together with previous observations by others, are consistent with the hypothesis that the neural retina is the default state of the optic vesicle precursors. Morphologically, optic vesicle precursors either at the medial or the lateral layers share a similar neuroepithelial character: i.e. elongated cells polarized along the apico-basal axis arranged in a pseudostratified epithelium (Ivanovitch et al., 2013, Kwan et al., 2012). During optic cup morphogenesis, NR precursors retain this neuroepithelial morphology, whereas differentiating RPE cells undergo profound cell shape changes as they progressively flatten into a squamous epithelium (Moreno-Marmol et al., 2018). These observations correlate in our analyses with a larger number of differentially upregulated genes and differentially opened chromatin regions at the RPE (Fig EV1). We have shown that repressing CREs associated with TFs dominate in the RPE network, whereas activating elements are more abundant in the NR program (Fig 2). Furthermore, the motifs analysis of activating elements in the NR and repressing elements in the RPE indicates that they are antagonistically regulating a similar set of genes (Fig 6). It is therefore very likely that the acquisition of the RPE fate requires the global repression of the NR program. However, although the role of *Vsx2* in the

suppression of RPE identity via direct *Mitf* repression has been well documented (Horsford et al., 2005, Rowan et al., 2004, Zou & Levine, 2012), the mechanisms underlining the repressive activity of the RPE network are much less understood (Fuhrmann, Levine et al., 2000). Interestingly, *Pax6* seems to cooperate both with NR and RPE specifiers acting as a balancing factor between the two networks depending on the cellular context (Bharti, Gasper et al., 2012, Raviv, Bharti et al., 2014). The complexity of the transcriptional regulatory logic is often exemplified by its context dependence, which may happen at very different levels. As illustrated by a recent report in *Drosophila*, cis-regulatory modules can play a dual role as enhancers or silencers in different cellular environments (Gisselbrecht, Palagi et al., 2020). Our motif analysis also points to the same set of TFs acting as repressors or activators within the same tissue, suggesting their activity is conditioned by their chromatin environment. This is in line with previous work showing that TFs have diverse regulatory function depending on the enhancer context (Stampfel, Kazmar et al., 2015).

GRNs robustness. One of our main observations is that NR and RPE networks are remarkably robust. This is the case particularly in teleosts, in which whole genome duplication resulted in additional copies of eye specifiers. In zebrafish, several homeobox TFs expressed in the neural retina converge on the 5'- TAATT-3' motif, including vsx1, vsx2, rx1, rx2, rx3, lhx2b, lhx9, hmx1 and hmx4. Our motif discovery predictions suggest these proteins, together with sox factors, may coregulate the same genes and cooperate within the same CREs, although these are theoretical predictions that may require validation by direct ChIP-seq studies. This approach though is challenging in our experimental setting due to the limited starting material and the reduced availability of ChIP grade antibodies in zebrafish. Yet, we believe that considered in bulk these are informative predictions, as they are restricted to differentially open chromatin regions and differentially expressed TFs. Supporting a co-regulation, the mutation of many retinal homeobox TFs, such as rx2 in medaka (Reinhardt, Centanin et al., 2015) or lhx2 in zebrafish (Seth, Culverwell et al., 2006) does not compromise severely the identity of the tissue. In the RPE, redundancy pertains mainly the core bHLH motif (5'-CACGTG-3'), which is shared by mitfa, bHLHE40, bHLHE41, and tfec. In agreement with this, the simultaneous elimination of *mitfa* and *mitfb* has no apparent consequences on RPE specification or pigmentation in zebrafish (Lane & Lister, 2012). All these observations further point to a broad redundancy and cooperativity among the main nodes of the eye specification networks.

Coupling of transcriptional and cell shape changes in the RPE. As expected, the cell shape changes associated to the acquisition of the NR and RPE identities have a clear reflection at the transcriptomic level. The differentiation of the NR entails the activation of cytoskeletal components associated to microtubules polarization, in agreement with the neuroepithelial character of this tissue (Ivanovitch et al., 2013). In contrast, here we show that a significant number of keratins and other desmosomal genes get recruited in the presumptive RPE hours before this tissue acquires its distinctive pigmentation. Although keratins and plaque proteins have been reported as RPE markers in several vertebrate species (Owaribe, Kartenbeck et al., 1988), their contribution to the morphogenetic program of this tissue remains unexplored. Given the role of desmosomal plaques in conferring resistance to mechanical load (Hatzfeld et al., 2017), it is tempting to hypothesize that the activation of these cytoskeletal genes may be linked to RPE precursors adapting their cytoskeleton to increased tissue tension as they

flatten. Motif enrichment analysis of CREs linked to desmosomal genes, together with the genetic evidence here provided by *yap -/-; taz -/-* double mutants and Tead crispants, strongly support a role for Tead TFs in the activation of keratin genes at the RPE. Indeed, several keratin (e.g. *krt5, krt8,* and *krt97*), and desmosomal genes (e.g. *evpla, pleca, plecb,* and *ppl*) have been identified as targets for Yap/Tead complexes by ChIP-seq studies in mammalian cells (Estaras, Hsu et al., 2017, Lian, Kim et al., 2010, Zanconato, Forcato et al., 2015), as well as DamID-seq studies in zebrafish embryos (Vazquez-Marin, Gutierrez-Triana et al., 2019). Thus, the direct transcriptional regulation of keratins and plaque genes by Yap/Tead complexes seems a conserved theme across different tissues and vertebrate species.

Two-waves of TFs recruitment during RPE specification. Arguably one of the most notable observations in this study is the relatively late peak of expression of genes such as mitfa or otx2, which were previously considered among the earliest RPE specifiers (Martinez-Morales et al., 2004). Their delayed peak of expression occurs after the specification and flattening of RPE precursors have commenced, and rather coincides with the onset of pigmentation in the tissue at 23hpf. This agrees with previous studies indicating a cooperative role for Otx and Mitf in the direct activation of the melanogenic gene battery (Martinez-Morales et al., 2003). A detailed expression study in zebrafish also reported two separated phases for RPE specification, the first of which is *mitf* independent (Cechmanek & McFarlane, 2017). Here we show that the early phase of specification entails the recruitment of a different set of transcriptional regulators, including *smad6b*, *tead1b*, *tead3b*, *tfap2a*, *tfap2c*, *tcf12*, and *vgll2b*. Some of these TFs, or their cofactors, act as upstream regulators of Otx and Mitf, such is the case for Tcf (Westenskow, Piccolo et al., 2009) and Tead-cofactors Yap and Taz (Kim et al., 2016, Miesfeld et al., 2015). Others, such as *tfap2a*, play an important role in RPE specification, although their hierarchical role within the RPE GRN remains unclear (Bassett et al., 2010). Finally, here we also describe a similar sequence of TF recruitment in human differentiating RPE cells. Thus, despite the different morphology of the cells (i.e. cuboidal in humans and squamous in zebrafish) and the large evolutionary distance (\approx 450 mya), the regulatory logic that specifies the RPE seems largely conserved across vertebrates.

Conclusions. Our data shed light on the bifurcation of the NR and RPE programs not only in zebrafish but in other vertebrate species. The obtained results may be relevant to identify novel causative genes for eye hereditary diseases, as mutations in many nodes of the eye GRNs result in congenital eye malformations (Fitzpatrick & van Heyningen, 2005). Importantly, our findings uncover an unanticipated regulatory logic within the RPE specification network. This provides critical information to improve hiPSCs-to-RPE differentiation protocols, a key step in cell replacement strategies for retinal degenerative diseases.

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Author contribution.

LB conducted most experiments, performed bioinformatic analyses, and had a main contribution in figures and manuscript edition. SN, TMM, and RP contributed to protocols refining, FACS experiments, and fish maintenance. BdC and FJDC carried out hiPSCs experiments together with LB. OB performed 5mC analysis and contributed to manuscript edition. JRMM conceived the project and assisted LB in data analysis. The manuscript was edited and written by JRMM and PB, who co-supervised all the work.

Conflict of Interest.

The authors declare that they have no conflict of interest

Data Availability

Datasets supporting the conclusions of this article are available in the Gene Expression Omnibus (GEO) repository (<u>https://www.ncbi.nlm.nih.gov/geo</u>) under the following accession numbers: RNA-seq (GSE150346) and ATAC-seq (GSE150189).

Materials and Methods

Fish maintenance

The zebrafish (Danio rerio) AB/Tübingen (AB/TU) wild-type strains, the transgenic lines tg(vsx2.2:GFP-caax) (Nicolas-Perez et al., 2016) and tg(E1_bHLHe40:GFP) (Moreno-Marmol et al., 2020) and the mutant strain yap +/- taz +/- (Miesfeld et al., 2015) were maintained and breed under previously described experimental conditions (Kimmel et al., 1995, Westerfield, 2000). All animal experiments were carried out according to the guidelines of our Institutional Animal Ethics Committee.

Cell cytometry

Zebrafish cells were dissociated and prepared for FACS as previously described (Manoli & Driever, 2012). PG at 16 hpf, NR at 18 and 23 hpf were isolated from dissected heads of the tg(*vsx2.2:caax-GFP*). RPE at 18 and 23 hpf were isolated from whole tg(E1_*bHLHE40*:GFP) embryos. A FACSAriaTM Fusion flow cytometer was used to recollect only the GFP+ cells. GFP+ cells were isolated directly in Trizol for RNA extraction, or in ATAC-seq tagmentation buffer for open chromatin detection.

RNA extraction

Total RNA was extracted using 750 ul TRIzol LS (Invitrogen) following manufacturer's protocol. Possible DNA contamination was eliminated treating the RNA samples with with TURBO DNAse-free (Ambion). Concentration of the RNA samples was evaluated by Qubit (Thermo Fisher), and then the samples were used for subsequent applications.

qPCR

cDNA retrotranscription and qPCR were performed as described (Vazquez-Marin et al., 2019). For primer sequences, see Appendix tables S1 and S2. *HPRT1* and *GAPDH* were used as housekeeping genes for human samples, whereas *eef1a111* was used for zebrafish samples.

RNA-seq analyses.

RNA was extracted from sorted cells and then treated with DNAse as described above. rRNAs were eliminated from the samples with Ribo-Zero[®] rRNA Removal Kit (Illumina) prior library preparation. Samples were sequenced in SEx125bps or PEx125bps reads with an Illumina Hiseq 2500. We obtained at least 35 M reads from the sequencing of each library. Three biological replicates were used for each analyzed condition. Reads were aligned to the danRer10 zebrafish genome assembly using Tophat v2.1.0. Transcript abundance was estimated with Cufflinks v2.2.1. Differential gene expression analysis was performed using Cuffdiff v2.2.1, setting a corrected p value < 0.05 as the cutoff for statistical significance of the differential expression. Multidimensional Scaling Analysis (MDS) was performed using the function MDSplot of the CummeRbund package in R 3.6.1. Soft clustering of time-series gene expression data was done for all the transcripts with a variance among the five conditions \geq 3 using the R package Mfuzz with a m = 1.5 (Kumar & M, 2007). The TF transcript subset was extracted from the total list of genes using the tool "Classification System" of PANTHER (Thomas, Campbell et al., 2003) filtering for the protein class "transcription factors" (PC00218). Some TFs not present in the database for an annotation issues (i.e. mitfa, vsx1, vsx2, rx1, rx2,

rx3, lhx2b, hmx1, hmx4, sox21a) were added manually. The cytoskeleton component subset was obtained retrieving all the genes belonging to GO term "cytoskeleton" (GO:0005856), including all the child and further descendant GO terms, with biomaRt. All the heatmaps representing transcriptomic variations were plotted with the R package *pheatmap* using exclusively the transcripts that resulted to be differentially expressed from the comparison between at least two of our experimental conditions TFs and cytoskeleton components were filtered using the same methodology used for Mfuzz clustering. Gene ontology analysis was performed with the online tool GOrilla (Eden, Navon et al., 2009) or Panther (Thomas et al., 2003) using two unranked lists of genes (target and background lists).

ATAC-seq

ATAC-seq was performed starting from 5000 sorted cells using a FAST-ATAC protocol previously described (Corces, Buenrostro et al., 2016). All the libraries were sequenced 2x50 bp with an Illumina Hiseq 2500 platform. We obtained at least 100 M reads from the sequencing of each library. For data comparison, we used two biological replicates for each condition. Reads were aligned to the danRer10 zebrafish genome and differential chromatin accessibility was calculated as reported (Magri, Jimenez-Gancedo et al., 2019). All chromatin regions reporting a differential accessibility with an adjusted p-value < 0.05 were considered as differentially open chromatin regions (DOCRs). All the DOCRS have been associated with genes using the online tool GREAT (McLean, Bristor et al., 2010) with the option "basal plus extension". Gene ontology analysis of all the genes associated with DOCRs was also performed with GREAT. De novo motif enrichment of TF binding sequences in the sets of DOCRs was performed using HOMER (Sven et al., 2010). Top enriched TF PWMs from the HOMER results and PWMs from JASPAR database (Fornes, Castro-Mondragon et al., 2020) were used as input for the online tool FIMO (Grant, Bailey et al., 2011) to assess the exact TFBS genome position in the DOCRs. Before estimating the rate of TF co-occupancy in same peak among the binding motifs for the different TFs, all the binding motif sequences overlapping for more than 3 bps were eliminated, keeping only the TF binding sequences with the lowest p-value.

Activating/repressing cis-regulatory element configuration

For this analysis only the DEGs and the DORCs that could be associated to each other were taken into account. The log2FoldChange values of transcript expression and chromatin accessibility of NR and RPE at 23 hpf were used to discriminate four clusters of activating and repressing CREs. Gene Ontology enrichment analysis of the genes belonging to the different clusters was performed with FishEnrichr (Kuleshov, Jones et al., 2016).

CRE 5mC analysis

Whole genome bisulfite sequencing and TET-assisted bisulfite sequencing (Bogdanovic et al., 2016) data was trimmed with Trimmomatic software (Bolger, Lohse et al., 2014) and mapped onto the danRer10 genome assembly using WALT (Chen, Smith et al., 2016) keeping only the reads mapping to a unique genomic location. Duplicates were removed with sambamba (Tarasov, Vilella et al., 2015) and DNA methylation levels were calculated using MethylDackel (https://github.com/dpryan79/MethylDackel). All heatmaps were made using Deeptools (Ramirez, Dundar et al., 2014) and Gene ontology enrichments were calculated with GREAT

(McLean et al., 2010). ATAC-seq reads were counted using BEDtools (Quinlan & Hall, 2010) and statistics were performed in R.

CRISPR/Cas9 F0 screening

All the sgRNAs were designed using the online tool CRISPscan (https://www.crisprscan.org/) (Moreno-Mateos, Vejnar et al., 2015) and synthetized following described protocols (Vejnar, Moreno-Mateos et al., 2016). All the sgRNAs were selected to target the first half of the CDS in exons resulting actually expressed in the eye tissues from our RNA-seq data (trying to avoid the first exon to prevent the usage of an alternative start codon that would produce a possibly functional protein), with an efficiency score > 58 and no predicted off-targets (Appendix Table S3). Two different sgRNAs were used together to target the same gene. The sgRNAs were injected in the zebrafish yolk at 1-cell stage at final concentration of 80 ng/µl together with the Cas9 endonuclease at a concentration of 300 ng/µl. 1 nl of the mixture was injected in each embryo. In the case a target gene had a close paralogue, the sgRNAs targeting both of the paralogues were injected at the same time, adjusting the final concentration of the sgRNA-Cas9 mixture. Lethality, phenotypic features and penetrance were assessed at 24 and 48 hpf.

Cell culture

Human hiPSCs were obtained, after informed consent, from peripheral blood monocytes by cell reprogramming using a non-integrative Sendai virus vector as described (Garcia Delgado, de la Cerda et al., 2019). Cells were maintained in feeder-free adherent conditions onto Matrigel-covered plates in standard incubation at 37° C, 5% CO₂, 20% O₂. hiPSCs were fed every two days with mTser1 serum-free culture medium and passaged every 5-7 days depending on confluency. Dispase was used for gentle dissociation for passage. A 6-well plate with undifferentiated, well-grown hiPSCs was the starting point of the experiment (Day 0). To induce RPE differentiation culture medium was changed to the following: KO DMEM, KSR 15%, Glutamax 2 mM, non-essential aminoacids 0.1 mM, β -mercaptoethanol 0.23 mM, Peniciline/streptomycin. Differentiating cells were harvested directly in Trizol LS (Invitrogen) at day 0, and weeks 1, 2, 3 and 4 for gene expression studies.

Figure Legends: (max 300 words)

Figure 1: Experimental setup and raw data (A) Schematic representation of optic vesicle morphogenesis from undifferentiated retinal progenitors cells (PG, blue), at 16 hpf, to segregation and differentiation of the neural retina (NR, purple) and pigmented epithelium (RPE, orange) domains at 18 and 23 hpf. NR and RPE populations isolated by flow cytometry were analysed by RNA-seq and ATAC-seq at the stages indicated. (B) Stage 23 hpf confocal sections from the zebrafish transgenic lines used to mark and isolate the RPE (*E1_bHLHe40:GFP*) and NR (*vsx2.2:GFP*) populations. (C) Multidimensional scaling analysis of the RNA-seq data showing a progressive transcriptomic divergence between the NR and RPE domains. (D) Percentage of overlapping between differentially expressed genes (DEGs) and genes associated with differentially opened chromatin regions (DOCRs). (E-F) Overview of ATAC-seq and RNA-seq tracks (UCSC browser) for representative NR (*vsx1*; E) and RPE (*mitfa*; F) specific genes. Solid bars on the top indicate DOCRs. If purple, the DOCR is more accessible in the NR. On the contrary, if orange, the DOCR is more accessible in the RPE. In the depicted case all the DOCRs are accompanied by an increased transcription of the associated gene in the corresponding tissue.

Figure 2: **CRE configuration in the NR and RPE domains**. (A) Schematic representation of the functional relationship between DOCRs, either activating (green) or repressing (red), and their associated DEGs. (B) The graph illustrates the correlation between the levels of differentially expressed TFs (log2FC) and the accessibility of their associated DOCRs (log2). (C) The table summarizes the number of activating or repressing CREs associated with either to all the DEGs or only with the differentially expressed TFs. (D-E) Histograms correlating the number of TFs associated with activating or repressing CREs to the number of peaks per gene, for both the NR (D) and RPE (E) domains.

Figure 3: **Hierarchical clustering of TFs gene expression variations during optic cup development.** (A) Hierarchical clustering output shows TFs expression trends in the distinct domains and stages. Gene expression values, normalized by row, are indicated with a red to blue graded colour. Note that most clusters comprise a particular domain and developmental stage. (B) Relative transcript level changes for significant TFs (indicated by dotted lines) within each cluster.

Figure 4: **NR motif enrichment analysis**. (A) Representative TF binding motifs enriched in NR DOCRs as identified by HOMER. The binding motif similarity among neural retina TFs of the homeobox family is indicated (<u>http://jaspar.genereg.net</u>). (B) Circoplot illustrating the co-occupancy rate of TFBS in the same DOCRs for the main TFs identified in the motif enrichment analysis (C) Circoplot illustrating the degree of co-regulation between TFs regulating the same gene through different DOCRs. (D) Number of CREs containing the main TFs identified in the motif enrichment analysis, classified according to their co-occupancy (E) Number of genes associated to CREs containing the main TFs identified in the motif enrichment analysis, classified according to their co-regulation.

Figure 5: **RPE motif enrichment analysis**. (A) Representative TF binding motifs enriched in RPE DOCRs as identified by HOMER. Analysis of the binding motif similarity among TFs of the bHLH family (<u>http://jaspar.genereg.net</u>). (B) Circoplot illustrating the co-occupancy rate of TFBS in the same DOCRs for the main TFs identified in the motif enrichment analysis (C) Circoplot illustrating the degree of co-regulation between TFs regulating the same gene through different DOCRs. (D) Average percentage of co-occupancy in the same DOCR for two different TF in the RPE and NR. (E) Average percentage of two different TF regulating the same gene through different DOCRs in the RPE and NR.

Figure 6: *Motif enrichment analysis of activating and repressing CREs*. Representative TF motifs enriched in activating (left column) and repressing (right column) CREs in both the NR (A) and RPE (B) domains. Analysis of GO terms enrichment for genes associated with each set of elements is indicated.

Figure 7: **Regulation of desmosomal components during RPE specification.** (A) Schematic representation of a desmosome junction. (B) Intermediate filament and desmosomal genes expression variations during optic cup development. Expression values are reported as log(FPKM) (C) Motif enrichment analysis of the DOCRs associated with genes encoding intermediate filament or desmosome components. (D) mRNA levels of keratin genes as well as RPE and NR markers as determined by RT-qPCR in wild type and *yap -/- taz -/-* double mutant zebrafish samples (dissected heads) at 18 hpf. Significant differences are indicated (n=3; T-test; ***= p<0.001). (E) Representative stereo microscope images of zebrafish embryo heads at 48 hpf. Magnification Bar = 0.5 mm): wild type and embryos injected with Cas9 (300 ng/µl) together with the following sgRNAs (80 ng/µl) combinations: *vgll2a* and *vgll2b* (vgll2); *tead1a* and *tead1b* (tead1) and *dspa and dspb* (dsp). Note the reduced eye size and RPE hypopigmentation in the crispants.

Figure 8: Gene expression during hiPSCs-to-RPE differentiation. (A) Bright field microscopy images (10-x and 20x) of hiPSCs before (Day 0) and during their differentiation towards RPE. Note the progressive acquisition of epithelial morphology and pigmentation. (B) Hierarchical clustered heatmap showing gene expression level variations, as determined by RT-qPCR during the differentiation towards RPE from hiPSCs. Note the conservation of the two clusters (early and late transcriptional waves) identified in zebrafish.

Expanded View. Figure Legends:

Figure EV1: Analysis of differentially opened chromatin regions (DOCRs) during optic cup morphogenesis. (A) Volcano plot illustrating the chromatin accessibility changes during eye morphogenesis. Each dot corresponds to a peak (i.e. open chromatin region). Black dots indicate not significant variations; colour shades chromatin accessibility changes with different ranges of adjusted p-value (darker: p<0.05; medium= p<0.01, lighter= p<0.001). (B) Table including the number of peaks significantly more or less accessible between the two conditions. (C) Frequency histogram showing the distribution of the DOCRs in relation to their accessibility fold change. (D) Pie charts displaying the percentage of peaks falling in distinct regions of zebrafish genome. Top: genome distribution of the whole set of open chromatin regions (OCRs) identified by ATAC-seq in each condition. Bottom: genome distribution of the DOCRs.

Figure EV2: DNA methylation profiles of NR and RPE peaks. Clustering of NR (A) and RPE (B) ATAC-seq peaks according to 5mC pattern across developmental stages and in adult brain (left columns), as well as 5hmC, K4me3, and K27ac profiles at 24 hpf (right columns). Note that the signature of cluster #1 corresponds to active enhancers, whereas that of cluster #2 corresponds to hypomethylated promoter regions. (C) Chromatin accessibility quantification for NR and RPE peaks in wild type and triple tet (tet1/2/3) morphants. Analysis was performed in whole 24hpf embryos. Statistical significance was evaluated using a Mann-Whitney-Wilcoxon Test, P < 2.2e-16. (D) Chromatin accessibility levels for NR and RPE ATAC-seq peaks in wild type and triple tet (tet1/2/3) zebrafish morphants.

Figure EV3: Activating and repressing CREs localization. (A) Schematic representation of the functional relations among DOCRs, either activating (green) or repressing (red), and their associated DEGs. (B) Illustrative examples of ATAC-seq and RNA-seq tracks (UCSC browser) for representative NR (*six3*; E) and RPE (*otx2*; F) genes. Bars on the top indicate DOCRs. If purple, the DOCR is more accessible in NR; if orange, the DOCR is more accessible in RPE. Some of the more accessible DOCRs are accompanied by a decreased transcription of the associated gene in the corresponding tissue. (C) Average distance to the TSS of all activating and repressing DOCRs for the NR and RPE domains (see Figure 2C). Note that activating CREs are significantly (T-test) closer to the TSS than repressing regions in both domains.

Figure EV4: Hierarchical clustering of gene expression variations during optic cup development for cytoskeletal components. (A) Hierarchical clustering output showing the expression trends in distinct domains and stages. Gene expression values, normalized by row, are indicated with a red to blue graded colour. Note that all stages and domains are represented by at least one gene cluster (B) Cellular component GO enrichment for cytoskeletal genes belonging to NR or RPE clusters.

Figure EV5: CRISPR CRISPR/Cas9 F0 screening. (A) Eye phenotypes resulting from the injection of Cas9 protein and sgRNAs for the candidate genes. (B) Percentage of embryos showing an impaired eye phenotype upon the injection of Cas9/sgRNAs complex (n= total number of

injected embryos). The % of lethality upon injection is also shown. See also Supplementary data set 12.

Appendix . Figure Legends:

Appendix Figure S1: Eye domain transcriptome variations during optic cup morphogenesis. (A) Volcano plots illustrating the transcriptome variations during eye morphogenesis. Each dot corresponds to a gene. Black dots indicate not significant variations, whereas coloured dots point out significant expression variations among domains and developmental stages. (B) Table summarizing the number of DEGs (upregulated or downregulated) in each one of the conditions.

Appendix Figure S2: Gene ontology enrichment of the genes associated with DOCRs. Bar chart showing GO terms for biological processes enriched in the genes associated to DOCRs. Bar length is proportional to enrichment significance.

Appendix Figure S3: Partitioning clustering of gene expression variations during optic cup development. Partitioning clustering output (k= 25) showing the expression trends in the distinct domains and stages examined for two classes of genes: TFs (A) and cytoskeleton components (B). Stages and domains examined are linearly represented with a colour code and numbered from 1 (NR 23hpf) to 5 (RPE 23hpf), following the lineage trajectory.

Appendix Figure S4: GO terms associated with RPE TFs. Hierarchical clustering of the main RPE TFs according to the enriched GO terms (Biological process) associated to their putative downstream genes (genes associated with RPE DOCRs that contain each TFBS).

References

Aldiri I, Xu B, Wang L, Chen X, Hiler D, Griffiths L, Valentine M, Shirinifard A, Thiagarajan S, Sablauer A, Barabas ME, Zhang J, Johnson D, Frase S, Zhou X, Easton J, Zhang J, Mardis ER, Wilson RK, Downing JR et al. (2017) The Dynamic Epigenetic Landscape of the Retina During Development, Reprogramming, and Tumorigenesis. Neuron 94: 550-568 e10

Araki H, Miura F, Watanabe A, Morinaga C, Kitaoka F, Kitano Y, Sakai N, Shibata Y, Terada M, Goto S, Yamanaka S, Takahashi M, Ito T (2019) Base-Resolution Methylome of Retinal Pigment Epithelial Cells Used in the First Trial of Human Induced Pluripotent Stem Cell-Based Autologous Transplantation. Stem Cell Reports 13: 761-774

Arendt D, Wittbrodt J (2001) Reconstructing the eyes of Urbilateria. Philos Trans R Soc Lond B Biol Sci 356: 1545-63

Barabino SM, Spada F, Cotelli F, Boncinelli E (1997) Inactivation of the zebrafish homologue of Chx10 by antisense oligonucleotides causes eye malformations similar to the ocular retardation phenotype. Mech Dev 63: 133-43

Bassett EA, Williams T, Zacharias AL, Gage PJ, Fuhrmann S, West-Mays JA (2010) AP-2alpha knockout mice exhibit optic cup patterning defects and failure of optic stalk morphogenesis. Hum Mol Genet 19: 1791-804

Bharti K, Gasper M, Ou J, Brucato M, Clore-Gronenborn K, Pickel J, Arnheiter H (2012) A regulatory loop involving PAX6, MITF, and WNT signaling controls retinal pigment epithelium development. PLoS Genet 8: e1002757

Bharti K, Liu W, Csermely T, Bertuzzi S, Arnheiter H (2008) Alternative promoter use in eye development: the complex role and regulation of the transcription factor MITF. Development 135: 1169-78

Bogdanovic O, Fernandez-Minan A, Tena JJ, de la Calle-Mustienes E, Hidalgo C, van Kruysbergen I, van Heeringen SJ, Veenstra GJ, Gomez-Skarmeta JL (2012) Dynamics of enhancer chromatin signatures mark the transition from pluripotency to cell specification during embryogenesis. Genome Res 22: 2043-53

Bogdanovic O, Smits AH, de la Calle Mustienes E, Tena JJ, Ford E, Williams R, Senanayake U, Schultz MD, Hontelez S, van Kruijsbergen I, Rayon T, Gnerlich F, Carell T, Veenstra GJ, Manzanares M, Sauka-Spengler T, Ecker JR, Vermeulen M, Gomez-Skarmeta JL, Lister R (2016) Active DNA demethylation at enhancers during the vertebrate phylotypic period. Nat Genet 48: 417-26

Bolger AM, Lohse M, Usadel B (2014) Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 30: 2114-20

Buenrostro JD, Giresi PG, Zaba LC, Chang HY, Greenleaf WJ (2013) Transposition of native chromatin for fast and sensitive epigenomic profiling of open chromatin, DNA-binding proteins and nucleosome position. Nat Methods 10: 1213-8

Buono L, Martinez-Morales JR (2020) Retina Development in Vertebrates: Systems Biology Approaches to Understanding Genetic Programs: On the Contribution of Next-Generation Sequencing Methods to the Characterization of the Regulatory Networks Controlling Vertebrate Eye Development. Bioessays 42: e1900187

Cardozo MJ, Almuedo-Castillo M, Bovolenta P (2020) Patterning the Vertebrate Retina with Morphogenetic Signaling Pathways. Neuroscientist 26: 185-196

Cechmanek PB, McFarlane S (2017) Retinal pigment epithelium expansion around the neural retina occurs in two separate phases with distinct mechanisms. Dev Dyn 246: 598-609

Chapouton P, Webb KJ, Stigloher C, Alunni A, Adolf B, Hesl B, Topp S, Kremmer E, Bally-Cuif L (2011) Expression of hairy/enhancer of split genes in neural progenitors and neurogenesis domains of the adult zebrafish brain. J Comp Neurol 519: 1748-69

Chen H, Smith AD, Chen T (2016) WALT: fast and accurate read mapping for bisulfite sequencing. Bioinformatics 32: 3507-3509

Collin J, Queen R, Zerti D, Dorgau B, Hussain R, Coxhead J, Cockell S, Lako M (2019) Deconstructing Retinal Organoids: Single Cell RNA-Seq Reveals the Cellular Components of Human Pluripotent Stem Cell-Derived Retina. Stem Cells 37: 593-598

Corces MR, Buenrostro JD, Wu B, Greenside PG, Chan SM, Koenig JL, Snyder MP, Pritchard JK, Kundaje A, Greenleaf WJ, Majeti R, Chang HY (2016) Lineage-specific and single-cell chromatin accessibility charts human hematopoiesis and leukemia evolution. Nat Genet 48: 1193-203

Coulombre JL, Coulombre AJ (1965) Regeneration of neural retina from the pigmented epithelium in the chick embryo. Dev Biol 12: 79-92

Eden E, Navon R, Steinfeld I, Lipson D, Yakhini Z (2009) GOrilla: a tool for discovery and visualization of enriched GO terms in ranked gene lists. BMC Bioinformatics 10: 48

Estaras C, Hsu HT, Huang L, Jones KA (2017) YAP repression of the WNT3 gene controls hESC differentiation along the cardiac mesoderm lineage. Genes Dev 31: 2250-2263

Fitzpatrick DR, van Heyningen V (2005) Developmental eye disorders. Curr Opin Genet Dev 15: 348-53

Fornes O, Castro-Mondragon JA, Khan A, van der Lee R, Zhang X, Richmond PA, Modi BP, Correard S, Gheorghe M, Baranasic D, Santana-Garcia W, Tan G, Cheneby J, Ballester B, Parcy F, Sandelin A, Lenhard B, Wasserman WW, Mathelier A (2020) JASPAR 2020: update of the open-access database of transcription factor binding profiles. Nucleic Acids Res 48: D87-D92

Fuhrmann S (2010) Eye morphogenesis and patterning of the optic vesicle. Curr Top Dev Biol 93: 61-84

Fuhrmann S, Levine EM, Reh TA (2000) Extraocular mesenchyme patterns the optic vesicle during early eye development in the embryonic chick. Development 127: 4599-609

Garcia Delgado AB, de la Cerda B, Alba Amador J, Valdes Sanchez ML, Fernandez-Munoz B, Relimpio Lopez I, Rodriguez de la Rua E, Diez Lloret A, Calado SM, Sanchez Pernaute R, Bhattacharya SS, Diaz Corrales FJ (2019) Subretinal Transplant of Induced Pluripotent Stem Cell-Derived Retinal Pigment Epithelium on Nanostructured Fibrin-Agarose. Tissue Eng Part A 25: 799-808

Gehring WJ (2014) The evolution of vision. Wiley Interdiscip Rev Dev Biol 3: 1-40

Gisselbrecht SS, Palagi A, Kurland JV, Rogers JM, Ozadam H, Zhan Y, Dekker J, Bulyk ML (2020) Transcriptional Silencers in Drosophila Serve a Dual Role as Transcriptional Enhancers in Alternate Cellular Contexts. Mol Cell 77: 324-337 e8

Grant CE, Bailey TL, Noble WS (2011) FIMO: scanning for occurrences of a given motif. Bioinformatics 27: 1017-8

Hatzfeld M, Keil R, Magin TM (2017) Desmosomes and Intermediate Filaments: Their Consequences for Tissue Mechanics. Cold Spring Harb Perspect Biol 9

Heermann S, Schutz L, Lemke S, Krieglstein K, Wittbrodt J (2015) Eye morphogenesis driven by epithelial flow into the optic cup facilitated by modulation of bone morphogenetic protein. Elife 4

Horsford DJ, Nguyen MT, Sellar GC, Kothary R, Arnheiter H, McInnes RR (2005) Chx10 repression of Mitf is required for the maintenance of mammalian neuroretinal identity. Development 132: 177-87

Hu Y, Wang X, Hu B, Mao Y, Chen Y, Yan L, Yong J, Dong J, Wei Y, Wang W, Wen L, Qiao J, Tang F (2019) Dissecting the transcriptome landscape of the human fetal neural retina and retinal pigment epithelium by single-cell RNA-seq analysis. PLoS Biol 17: e3000365

Ivanovitch K, Cavodeassi F, Wilson SW (2013) Precocious acquisition of neuroepithelial character in the eye field underlies the onset of eye morphogenesis. Dev Cell 27: 293-305

Kim JY, Park R, Lee JH, Shin J, Nickas J, Kim S, Cho SH (2016) Yap is essential for retinal progenitor cell cycle progression and RPE cell fate acquisition in the developing mouse eye. Dev Biol 419: 336-347

Kozmik Z, Ruzickova J, Jonasova K, Matsumoto Y, Vopalensky P, Kozmikova I, Strnad H, Kawamura S, Piatigorsky J, Paces V, Vlcek C (2008) Assembly of the cnidarian camera-type eye from vertebrate-like components. Proc Natl Acad Sci U S A 105: 8989-93

Kuleshov MV, Jones MR, Rouillard AD, Fernandez NF, Duan Q, Wang Z, Koplev S, Jenkins SL, Jagodnik KM, Lachmann A, McDermott MG, Monteiro CD, Gundersen GW, Ma'ayan A (2016) Enrichr: a comprehensive gene set enrichment analysis web server 2016 update. Nucleic Acids Res 44: W90-7

Kumar L, M EF (2007) Mfuzz: a software package for soft clustering of microarray data. Bioinformation 2: 5-7

Kwan KM, Otsuna H, Kidokoro H, Carney KR, Saijoh Y, Chien CB (2012) A complex choreography of cell movements shapes the vertebrate eye. Development 139: 359-72

Land MF, Nilsson D-E (2002) Animal eyes. Oxford University Press, Oxford ; New York

Lane BM, Lister JA (2012) Otx but not Mitf transcription factors are required for zebrafish retinal pigment epithelium development. PLoS One 7: e49357

Letelier J, Bovolenta P, Martinez-Morales JR (2017) The pigmented epithelium, a bright partner against photoreceptor degeneration. J Neurogenet 31: 203-215

Li Z, Joseph NM, Easter SSJ (2000) The morphogenesis of the zebrafish eye, including a fate map of the optic vesicle. Developmental Dynamics 218: 175-188

Lian I, Kim J, Okazawa H, Zhao J, Zhao B, Yu J, Chinnaiyan A, Israel MA, Goldstein LS, Abujarour R, Ding S, Guan KL (2010) The role of YAP transcription coactivator in regulating stem cell self-renewal and differentiation. Genes Dev 24: 1106-18

Liu ISC, Chen JD, Ploder L, Vidgen D, Vanderkooy D, Kalnins VI, Mcinnes RR (1994) Developmental expression of a novel murine homeobox gene (Chx10): Evidence for roles in determination of the neuroretina and inner nuclear layer. Neuron 13: 377-393

Loosli F, Winkler S, Burgtorf C, Wurmbach E, Ansorge W, Henrich T, Grabher C, Arendt D, Carl M, Krone A, Grzebisz E, Wittbrodt J (2001) Medaka *eyeless* is the key factor linking retinal dertermination and eye growth. Development 128: 4035-4044

Lowe EK, Cuomo C, Voronov D, Arnone MI (2019) Using ATAC-seq and RNA-seq to increase resolution in GRN connectivity. Methods Cell Biol 151: 115-126

Magri MS, Jimenez-Gancedo S, Bertrand S, Madgwick A, Escriva H, Lemaire P, Gomez-Skarmeta JL (2019) Assaying Chromatin Accessibility Using ATAC-Seq in Invertebrate Chordate Embryos. Front Cell Dev Biol 7: 372

Manoli M, Driever W (2012) Fluorescence-activated cell sorting (FACS) of fluorescently tagged cells from zebrafish larvae for RNA isolation. Cold Spring Harb Protoc 2012

Martinez-Morales JR (2016) Vertebrate Eye Gene Regulatory Networks. In Organogenetic Gene Networks, J. C-GH, P. B (eds) Switzerland: Springer, Cham

Martinez-Morales JR, Cavodeassi F, Bovolenta P (2017) Coordinated Morphogenetic Mechanisms Shape the Vertebrate Eye. Front Neurosci 11: 721

Martinez-Morales JR, Dolez V, Rodrigo I, Zaccarini R, Leconte L, Bovolenta P, Saule S (2003) OTX2 activates the molecular network underlying retina pigment epithelium differentiation. J Biol Chem 278: 21721-31

Martinez-Morales JR, Rodrigo I, Bovolenta P (2004) Eye development: a view from the retina pigmented epithelium. Bioessays 26: 766-77

Martinez-Morales JR, Signore M, Acampora D, Simeone A, Bovolenta P (2001) Otx genes are required for tissue specification in the developing eye. Development 128: 2019-2030

Masai I, Stemple DL, Okamoto H, Wilson SW (2000) Midline signals regulate retinal neurogenesis in zebrafish. Neuron 27: 251-263

Maston GA, Evans SK, Green MR (2006) Transcriptional regulatory elements in the human genome. Annu Rev Genomics Hum Genet 7: 29-59

Mathers PH, Grinberg A, Mahon KA, Jamrich M (1997) The Rx homeobox gene is essential for vertebrate eye development. Nature 387: 603-607

McLean CY, Bristor D, Hiller M, Clarke SL, Schaar BT, Lowe CB, Wenger AM, Bejerano G (2010) GREAT improves functional interpretation of cis-regulatory regions. Nat Biotechnol 28: 495-501

Miesfeld JB, Gestri G, Clark BS, Flinn MA, Poole RJ, Bader JR, Besharse JC, Wilson SW, Link BA (2015) Yap and Taz regulate retinal pigment epithelial cell fate. Development 142: 3021-32 Mochii M, Ono T, Matsubara Y, Eguchi G (1998) Spontaneous transdifferentiation of quail pigmented epithelial cell is accompanied by a mutation in the Mitf gene. Dev Biol 196: 145-59 Moreno-Marmol T, Cavodeassi F, Bovolenta P (2018) Setting Eyes on the Retinal Pigment Epithelium. Front Cell Dev Biol 6: 145

Moreno-Mateos MA, Vejnar CE, Beaudoin JD, Fernandez JP, Mis EK, Khokha MK, Giraldez AJ (2015) CRISPRscan: designing highly efficient sgRNAs for CRISPR-Cas9 targeting in vivo. Nat Methods 12: 982-8

Nguyen MT, Arnheiter H (2000) Signaling and transcriptional regulation in early mammalian eye development: a link between FGF and MITF. Development 127: 3581-3591

Nicolas-Perez M, Kuchling F, Letelier J, Polvillo R, Wittbrodt J, Martinez-Morales JR (2016) Analysis of cellular behavior and cytoskeletal dynamics reveal a constriction mechanism driving optic cup morphogenesis. Elife 5

Owaribe K, Kartenbeck J, Rungger-Brandle E, Franke WW (1988) Cytoskeletons of retinal pigment epithelial cells: interspecies differences of expression patterns indicate independence of cell function from the specific complement of cytoskeletal proteins. Cell Tissue Res 254: 301-15

Picker A, Cavodeassi F, Machate A, Bernauer S, Hans S, Abe G, Kawakami K, Wilson SW, Brand M (2009) Dynamic coupling of pattern formation and morphogenesis in the developing vertebrate retina. PLoS Biol 7: e1000214

Pittack C, Jones M, Reh TA (1991) Basic fibroblast growth factor induces retinal pigment epithelium to generate neural retina in vitro. Development 113: 577-88

Quinlan AR, Hall IM (2010) BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26: 841-2

Ramirez F, Dundar F, Diehl S, Gruning BA, Manke T (2014) deepTools: a flexible platform for exploring deep-sequencing data. Nucleic Acids Res 42: W187-91

Raviv S, Bharti K, Rencus-Lazar S, Cohen-Tayar Y, Schyr R, Evantal N, Meshorer E, Zilberberg A, Idelson M, Reubinoff B, Grebe R, Rosin-Arbesfeld R, Lauderdale J, Lutty G, Arnheiter H, Ashery-Padan R (2014) PAX6 regulates melanogenesis in the retinal pigmented epithelium through feed-forward regulatory interactions with MITF. PLoS Genet 10: e1004360

Reinhardt R, Centanin L, Tavhelidse T, Inoue D, Wittbrodt B, Concordet JP, Martinez-Morales JR, Wittbrodt J (2015) Sox2, Tlx, Gli3, and Her9 converge on Rx2 to define retinal stem cells in vivo. EMBO J 34: 1572-88

Rembold M, Loosli F, Adams RJ, Wittbrodt J (2006) Individual cell migration serves as the driving force for optic vesicle evagination. Science 313: 1130-4

Rowan S, Chen CM, Young TL, Fisher DE, Cepko CL (2004) Transdifferentiation of the retina into pigmented cells in ocular retardation mice defines a new function of the homeodomain gene Chx10. Development 131: 5139-52

Roy A, de Melo J, Chaturvedi D, Thein T, Cabrera-Socorro A, Houart C, Meyer G, Blackshaw S, Tole S (2013) LHX2 is necessary for the maintenance of optic identity and for the progression of optic morphogenesis. J Neurosci 33: 6877-84

Seth A, Culverwell J, Walkowicz M, Toro S, Rick JM, Neuhauss SC, Varga ZM, Karlstrom RO (2006) belladonna/(Ihx2) is required for neural patterning and midline axon guidance in the zebrafish forebrain. Development 133: 725-35

Shankaran SS, Dahlem TJ, Bisgrove BW, Yost HJ, Tristani-Firouzi M (2017) CRISPR/Cas9-Directed Gene Editing for the Generation of Loss-of-Function Mutants in High-Throughput Zebrafish FO Screens. Curr Protoc Mol Biol 119: 31 9 1-31 9 22

Sidhaye J, Norden C (2017) Concerted action of neuroepithelial basal shrinkage and active epithelial migration ensures efficient optic cup morphogenesis. Elife 6

Stampfel G, Kazmar T, Frank O, Wienerroither S, Reiter F, Stark A (2015) Transcriptional regulators form diverse groups with context-dependent regulatory functions. Nature 528: 147-51

Strauss O (2005) The retinal pigment epithelium in visual function. Physiol Rev 85: 845-81

Tang X, Gao J, Jia X, Zhao W, Zhang Y, Pan W, He J (2017) Bipotent progenitors as embryonic origin of retinal stem cells. J Cell Biol 216: 1833-1847

Tarasov A, Vilella AJ, Cuppen E, Nijman IJ, Prins P (2015) Sambamba: fast processing of NGS alignment formats. Bioinformatics 31: 2032-4

Thomas PD, Campbell MJ, Kejariwal A, Mi H, Karlak B, Daverman R, Diemer K, Muruganujan A, Narechania A (2003) PANTHER: a library of protein families and subfamilies indexed by function. Genome Res 13: 2129-41

Vazquez-Marin J, Gutierrez-Triana JA, Almuedo-Castillo M, Buono L, Gomez-Skarmeta JL, Mateo JL, Wittbrodt J, Martinez-Morales JR (2019) yap1b, a divergent Yap/Taz family member, cooperates with yap1 in survival and morphogenesis via common transcriptional targets. Development 146

Vejnar CE, Moreno-Mateos MA, Cifuentes D, Bazzini AA, Giraldez AJ (2016) Optimization Strategies for the CRISPR-Cas9 Genome-Editing System. Cold Spring Harb Protoc 2016

Vopalensky P, Kozmik Z (2009) Eye evolution: common use and independent recruitment of genetic components. Philos Trans R Soc Lond B Biol Sci 364: 2819-32

Waddington CH (1966) Principles of development and differentiation. New York, Macmillan New York

Wang J, Zibetti C, Shang P, Sripathi SR, Zhang P, Cano M, Hoang T, Xia S, Ji H, Merbs SL, Zack DJ, Handa JT, Sinha D, Blackshaw S, Qian J (2018) ATAC-Seq analysis reveals a widespread decrease of chromatin accessibility in age-related macular degeneration. Nat Commun 9: 1364 Wang Z, Yasugi S, Ishii Y (2016) Chx10 functions as a regulator of molecular pathways controlling the regional identity in the primordial retina. Dev Biol 413: 104-11

Westenskow P, Piccolo S, Fuhrmann S (2009) Beta-catenin controls differentiation of the retinal pigment epithelium in the mouse optic cup by regulating Mitf and Otx2 expression. Development 136: 2505-10

Xu Y, Xu C, Kato A, Tempel W, Abreu JG, Bian C, Hu Y, Hu D, Zhao B, Cerovina T, Diao J, Wu F, He HH, Cui Q, Clark E, Ma C, Barbara A, Veenstra GJ, Xu G, Kaiser UB et al. (2012) Tet3 CXXC domain and dioxygenase activity cooperatively regulate key genes for Xenopus eye and neural development. Cell 151: 1200-13

Yoshida H, Lareau CA, Ramirez RN, Rose SA, Maier B, Wroblewska A, Desland F, Chudnovskiy A, Mortha A, Dominguez C, Tellier J, Kim E, Dwyer D, Shinton S, Nabekura T, Qi Y, Yu B, Robinette M, Kim KW, Wagers A et al. (2019) The cis-Regulatory Atlas of the Mouse Immune System. Cell 176: 897-912 e20

Zanconato F, Forcato M, Battilana G, Azzolin L, Quaranta E, Bodega B, Rosato A, Bicciato S, Cordenonsi M, Piccolo S (2015) Genome-wide association between YAP/TAZ/TEAD and AP-1 at enhancers drives oncogenic growth. Nat Cell Biol 17: 1218-27

Zou C, Levine EM (2012) Vsx2 controls eye organogenesis and retinal progenitor identity via homeodomain and non-homeodomain residues required for high affinity DNA binding. PLoS Genet 8: e1002924

Zuber ME, Gestri G, Viczian AS, Barsacchi G, Harris WA (2003) Specification of the vertebrate eye by a network of eye field transcription factors. Development 130: 5155-67

Figure 1



Figure 2



С		all genes			TFs			
		peaks	genes	peaks/gene		peaks	genes	peaks/gene
	NR activating CREs	3434	754	4.55		1174	182	6.45
	NR repressing CREs	1999	552	3.62		388	69	5.62
	RPE activating CREs	5984	1119	5.35		869	110	7.90
	RPE repressing CREs	3358	883	3.80		706	173	4.08



Figure 4









4000
 not co-occupied peaks
 same TFBS co-occupied peak
 other TFBS co-occupied peaks
 2000
 0
 1/hx
 rx
 vsx
 hmx
 sox



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Figure 5



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Figure 6



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Figure 7



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Figure 8



NR 23 hpf vs RPE 23 hpf

Α



В

adj p-value	DOCRs	NR	RPE
<0.05	30172 (12.6%)	11263	18909
<0.01	19012 (7.9%)	7062	11950
<0.001	11387 (4.8%)	4237	7150



RPE

- Promoter (<=1kb) (10.03%)</p>
- Promoter (1–2kb) (2.21%)







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Figure EV3





NR 23 hpf

microtubule organizing center

microtubule organizing center part

microtubule associated complex

intermediate filament

centrosome

microtubule

nuclear part

В

NR 18 hpf

0

0

2

4

1

6

2

-LOG(p-value)

8

10

3

12

4

PG 16 hpf

-LOG(p-value)

RPE 18 hpf

RPE 23 hpf

NR

RPE

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Figure EV5



В

% of sgRNA/Cas9 injected embryos displaying eye phenotype





D
F

DOWN	PG 16 hpf	NR 18 hpf	RPE 18 hpf	NR 23 hpf	RPE 23 hpf
PG 16 hpf	-	650	1154	977	1518
NR 18 hpf	1474	-	1974	498	2105
RPE 18 hpf	1538	1808	-	2017	826
NR 23 hpf	1690	470	2105	-	2383
RPE 23 hpf	1849	1867	610	2158	-

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RPE



melanocyte differentiation neuroepitelial cell differentiation epithelial cell migration epithelial cell differentiation response to yeast endothelial cell migration integrin mediated signaling pathway response to fungus retinoic acid receptor signaling pathway

B

Appendix Figure S3



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Appendix Figure S4



organic acid transmembrane transport drug transmembrane transport carboxylic acid transmembrane transport cardiocyte differentiation animal organ development regulation of cellular process regulation of nitrogen compound metabolic process tissue regeneration regeneration polysaccharide metabolic process polysaccharide biosynthetic process pottern specification process growth embryonic neurocranium morphogenesis developmental growth
cellular polysaccharide biosynthetic process cell junction organization cell–cell adhesion via plasma–membrane adhesion molecules cell junction assembly
cell migration cell motility signal transduction
regulation of cell shape regulation of cell migration regulation of cell adhesion pigmentation mornhogenesis of an enithelium
establishment or maintenance of cell polarity epithelium development
cell migration involved in gastrulation biological adhesion cell adhesion regulation of locomotion
regulation of cellular component movement regulation of cell motility regulation of cell morphogenesis
movement of cell or subcellular component negative regulation of multicellular organismal process tissue development niament cell differentiation
iridophore differentiation vascular endothelial growth factor receptor signaling pathway transmembrane receptor protein tyrosine kinase signaling pathway
neuron projection guidance negative regulation of response to external stimulus negative regulation of axon guidance negative regulation of axon extension involved in axon guidance
morphogenesis of a branching structure melanocyte differentiation dorsal aorta development
cell-cell adhesion cell fate specification blood vessel development aorta development
axon guidance Wnt signaling pathway vasculature development
sensory organ development regulation of response to stimulus regulation of plasma membrane bounded cell projection organization regulation of growth
regulation of developmental growth regulation of cell projection organization positive regulation of exocytosis
fin morphogenesis embryonic appendage morphogenesis anatomical structure formation involved in morphogenesis cell surface receptor signaling pathway involved in cell–cell signaling



associated GO-term
not associated GO-term



Appendix Supplementary Tables

Table SI. Primers used for Real Time q-PCR of human samples.

gene	gene ID	Fw_primer (5'->3')	Rv_primer (5'->3')
OCT4	ENSG00000204531	CTTCAGGAGATATGCAAAGCAGA	TGATCTGCTGCAGTGTGGG
NANOG	ENSG00000111704	GGATCCAGCTTGTCCCCAAA	AGGAAGGAAGAGAGAGAGACAGT
RPE65	ENSG00000116745	ACCACCTGTTTGATGGGCAA	AGTGCGGATGAACCTTCTGT
CRALBP	ENSG00000116745	GTCACAACTTGGCCCTGACT	GGTCCATGGTCCTTGGTTGT
TYR	ENSG0000077498	GATTCAGACCCAGACTCTTTTCA	ACGACACAGCAAGCTCACAA
HPRT1	ENSG00000165704	CCCTGGCGTCGTGATTAGTG	TCGAGCAAGACGTTCAGTCC
GAPDH	ENSG00000111640	AGGTCGGAGTCAACGGATTT	TGGAATTTGCCATGGGTGGA
MITF	ENSG00000187098	CCGGGCTCTGTTCTCACTTT	GGAACTGCTGCTCTTCAGCG
OTX2	ENSG00000165588	CCTCACTCGCCACATCTACT	AGTGGAACTTACAGCCTCATGG
BHLHE40	ENSG00000134107	ATTAACGAGTGCATCGCCCA	AGCTCACCAGCTTGTAAACCA
TFEC	ENSG00000105967	GATAAAATCCACTCATTGCTGGTCC	GGGCTTTCTGTAGCTGAGGC
TFAP2A	ENSG00000137203	GAGAGTAGCTCCACTTGGGTG	CCGTCGTGACGGTCCTCG
TFAP2C	ENSG0000087510	GAAGAGGACTGCGAGGATCG	GCTGATATTCGGCGACTCCA
TEAD1	ENSG00000187079	CCATTCCAGGGTTTGAGCCT	GCTTGGTTGTGCCAATGGAG
TEAD2	ENSG0000074219	TCGGAATGAACTGATCGCCC	CCTGGTCCTTCAACTTGGACT
TEAD3	ENSG0000007866	GACCGTACCATTGCCTCCTC	TTGCTGTACGTGTCAGGGTC
TEAD4	ENSG00000197905	GGCACCATTACCTCCAACGA	CAGCTCGTTCCGACCATACA
TCF12	ENSG00000140262	CCATGAAGGCTTGTCCCCAA	GGAGACTAGATTGACAGCCTGG
VGLL2	ENSG00000170162	GCTTTGCTCCGCCTGATGAC	ATAGGCTAGTTTCTGGTGGTAGG
SMAD6	ENSG00000137834	GGGCCCGAATCTCCGC	GGTCGTACACCGCATAGAGG
KRT5	ENSG00000186081	CGAGGAATGCAGACTCAGTG	GCTGCTGGAGTAGTAGCTTCC
KRT4	ENSG00000170477	TCCTTCATCGACAAGGTGCAG	GGGCTCAAGGTTTTTGCTGG
KRT8	ENSG00000170421	CAGCAAATGTTTGCGGAATGAA	AACCAGGCGGAGATCCCTTC
DSP	ENSG0000096696	AGGCTGGAGTACGATGACCT	TAGATGCCTCTAAAGCCTGC
EVPL	ENSG00000167880	CGACTTCCGACTGCTCCATCT	CCAAGTCCTCCAAGGGTGTG
NOTCH1	ENSG00000148400	CTGCCTCTTCGACGGCTTT	AAGTGGAAGGAGCTGTTGCG
NOTCH2	ENSG00000134250	CGAGTGTGTCCCAGGCTATC	CTTCACAGAGTAGGCCCCGA
NOTCH3	ENSG0000074181	GTCTTCCTGGGTTTGAGGGTC	GGGCACTGGCAGTTATAGGT

Table SII. Primers used for Real Time q-PCR of zebrafish samples.

gene	gene ID	Fw_primer (5'->3')	Rv_primer (5'->3')
tyr	ENSDARG00000039077	ACGGATACTTCATGGTGCCC	CGCTGACCTGGATCCTGTAAAT
tyrp1b	ENSDARG00000056151	GCCCGTCCAATGGTTCAAAG	GGAGCGCTGTAACCCTCAAT
krt4	ENSDARG00000017624	CTTCGTTGCGGCTCCTATCA	TCCAGGAAGCGCACTTTGTC
krt8	ENSDARG00000058358	TCCGCGCTCAGTATGAAGAC	AAGTTGGCTCGCTGTCCTTT
six3a	ENSDARG00000058008	AAAAACAGGCTCCAGCATCAA	AAGAATTGACGTGCCCGTGT
vsx2	ENSDARG0000005574	GGGATTAATTGGGCCTGGAGG	GCTGGCAGACTGGTTATGTTCC
eef1a1l1	ENSDARG00000020850	TCCACCGGTCACCTGATCTAC	CAACACCCAGGCGTACTTGA

Table SIII. sgRNAs used for CRISPR/Cas9 screen

gene	gene ID	sgRNA seg (5'->3')
mphosh10	ENSDARG0000053912	GGTGGCTTTCGTGGACGAGGCGG
mphosh10	ENSDARG0000053912	GGATTTCGAGGAGGCAGGGGTGG
heatr1	ENSDARG0000099742	GAGGTGCTGGCTCTCCGTCATGG
heatr1	ENSDARG0000099742	GAGGGAGGGCCAATCAGCAAAGG
hells	ENSDARG0000057738	TGGGGCTGCTGTGCTGGCACAGG
hells	ENSDARG0000057738	AGACAGGTTATTCTGGAGGGGGG
nop58	ENSDARG00000104353	AGAGATCTCGATGGGCACAGAGG
nop58	ENSDARG00000104353	GGGCATCAGAAACCAGATGGAGG
mcm5	ENSDARG0000019507	GCGTAACCCTGCAGCCCCGGTGG
mcm5	ENSDARG0000019507	GTGGCGCAGACCAAAGCCAAAGG
dkc1	ENSDARG0000016484	GAGCTGCGACGAGTCCGTTCCGG
dkc1	ENSDARG0000016484	GGGCTGCCTGATCGTGTGTGTGG
cirh1a	ENSDARG0000017675	GGGCCAATCTGGGCCATAACAGG
cirh1a	ENSDARG0000017675	AGAGGGTACGGGACGTCCCGCGG
wdr12	ENSDARG0000003287	GGGGAAGGCTGTGATGACTGTGG
wdr12	ENSDARG0000003287	GAGATCTGCAACCTCGGAGGAGG
ttc27	ENSDARG0000007918	GGAAGTTGCTCTGTTGGCGGTGG
ttc27	ENSDARG0000007918	GTGGCTCCTCTGCTCTTCGGTGG
dhx33	ENSDARG00000051785	GAGGCGGGCATCGGCCGGCAGGG
dhx33	ENSDARG00000051785	GTGTTTGGAGATGTCCCGGCAGG
mcm2	ENSDARG00000102798	GGGCCACACGGTGCGCGAGTGGG
mcm2	ENSDARG00000102798	GAGCGACTGACACTCAGGACAGG
tsr2	ENSDARG0000005772	GTGTGAGCAGGGCAGATTGGCGG
tsr2	ENSDARG0000005772	TGGAGCGTTCAGTCAGCAGAAGG
mif	ENSDARG00000071336	GTGACAGTACATCGCCGTACAGG
mif	ENSDARG00000071336	GTGAGCGAGCAGAGCGCACACGG
fbl	ENSDARG00000053912	GGTGGCTTTCGTGGACGAGGCGG
fbl	ENSDARG00000053912	GGATTTCGAGGAGGCAGGGGTGG
tcf12	ENSDARG0000004714	AGCTTCCAGTGGCGTTCCCTCGG
tcf12	ENSDARG0000004714	GTGGGCGACACCGAGTGTGGCGG
smad6b	ENSDARG0000031763	TGTGCTGCAGGTCAGACCACCGG
smad6b	ENSDARG0000031763	GGGGAAAGTCTTGAGTATGGAGG
vgli2a	ENSDARG00000041706	AGGGGACATCAGTTCGGTGGTGG
vgli2a	ENSDARG00000041706	GTGTATGCGGCTGCAAAATACGG
vgli2b	ENSDARG0000053773	
vgli2b	ENSDARG0000053773	GAGGICGGCICGGGGGAGAGGGG
neurod4	ENSDARG0000003469	
neuroa4	ENSDARG0000003469	
nr2f10		
<u>1112/10</u>		
nr2f1b		
1112/10 toad1a		
tead1a	ENSDARG0000028159	
tead1h	ENSDARG0000028133	GACACITARAATTICCCOCOACO
tead1b	ENSDARG0000059483	
tead?a	ENSDARG0000074321	GATGATCTTTCTGCGGCCACAGG
tead3a	ENSDARG0000074321	TGAAGGGTAGGGTACGGGCTCGG
tead3h	ENSDARG0000063649	TGAAGGTATGCGCTTTCCTGCGG
tead3b	ENSDARG0000063649	GATGGGGGTCGGCCAGAACTGGG
dspa	ENSDARG0000022309	TGGCACGTGACTGGACCTGGAGG
dspa	ENSDARG0000022309	AGGCTGATGCTCAGAGGGTTTGG
wu:fi04e1	ENSDARG0000076673	AGGATCTGAATATTCAGCGGCGG
wu:fi04e1	ENSDARG00000076673	GACGACGGCCTGGAGGAGTGCGG