

A New Strategy to Identify and Annotate Human RPE-Specific Gene Expression

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Abstract

Background: To identify and functionally annotate cell type-specific gene expression in the human retinal pigment epithelium (RPE), a key tissue involved in age-related macular degeneration and retinitis pigmentosa.

Methodology: RPE, photoreceptor and choroidal cells were isolated from selected freshly frozen healthy human donor eyes using laser microdissection. RNA isolation, amplification and hybridization to 44 k microarrays was carried out according to Agilent specifications. Bioinformatics was carried out using Rosetta Resolver, David and Ingenuity software.

Principal Findings: Our previous 22 k analysis of the RPE transcriptome showed that the RPE has high levels of protein synthesis, strong energy demands, is exposed to high levels of oxidative stress and a variable degree of inflammation. We currently use a complementary new strategy aimed at the identification and functional annotation of RPE-specific expressed transcripts. This strategy takes advantage of the multilayered cellular structure of the retina and overcomes a number of limitations of previous studies. In triplicate, we compared the transcriptomes of RPE, photoreceptor and choroidal cells and we deduced RPE specific expression. We identified at least 114 entries with RPE-specific gene expression. Thirty-nine of these 114 genes also show high expression in the RPE, comparison with the literature showed that 85% of these 39 were previously identified to be expressed in the RPE. In the group of 114 RPE specific genes there was an overrepresentation of genes involved in (membrane) transport, vision and ophthalmic disease. More fundamentally, we found RPE-specific involvement in the RAR-activation, retinol metabolism and GABA receptor signaling pathways.

Conclusions: In this study we provide a further specification and understanding of the RPE transcriptome by identifying and analyzing genes that are specifically expressed in the RPE.

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Introduction

The retinal pigment epithelium (RPE) is a monocellular retinal layer that plays a particularly important role in visual function. This is illustrated by its involvement in a large number of severe retinal disorders like age-related macular degeneration and retinitis pigmentosa.

The RPE has multiple functions including supplying the photoreceptors with nutrients, recycling retinal from the photoreceptors and regulating the ion balance in the subretinal space. The RPE secretes a number of growth factors. Thereby it is involved in the maintenance of the structure and cellular differentiation of adjacent (cell) layers, the photoreceptors on the apical side, and Bruch's membrane and the choroid on the basolateral side (See Figure 1) [1].

Embryologically, both the RPE and the photoreceptors develop from the neuro-epithelium. A fold in the neuro-epithelium causes two layers of this structure to face each other [1]. One of these layers develops into the RPE, the other into the neural retina. The neural retina further differentiates and photoreceptors develop in close interaction with the microvilli of the RPE [1]. The development of the choroid from neural crest cells also depends on cues from the mature RPE [2]. The choroidal vasculature is created through angiogenesis from existing blood vessels from the paraocular mesenchyme [2]. Finally, Bruch's membrane (BM) is created from the basement membrane of the RPE and the basement membrane of the endothelium. In this way, the retina develops into a neatly arranged multi-layered structure (Figure 1).

We recently analyzed the RPE transcriptome [3]. We reported on the expressed genes and correlated molecular pathways in the RPE from cells that were specifically isolated from healthy human donor eyes [3]. Functional annotation of the RPE transcriptome showed that the RPE has high levels of protein synthesis, strong

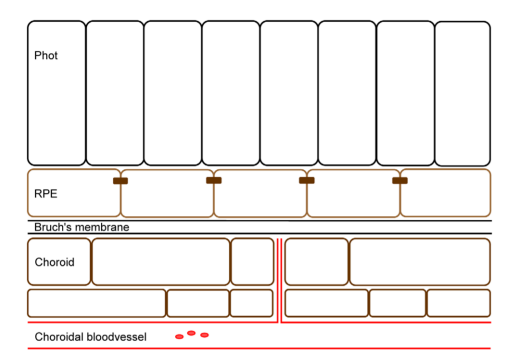


Figure 1. Overview of the (cell) layers surrounding the RPE. The dark brown rectangles connecting the RPE cells indicate tight junctions present between the cells. Phot: photoreceptors. doi:10.1371/journal.pone.0009341.q001

energy demands, is exposed to high levels of oxidative stress and has a variable degree of inflammation [3]. These data confirmed and expanded our knowledge on functional properties of the RPE, previously identified in a number of studies [4–7]. Nonetheless, due to the study design, our previous study was limited by unavoidable contamination from adjacent cell types [3,5,8].

In the current study we have overcome this limitation by using a new 44 k microarray strategy that includes both the RPE cell layer and the adjacent cell layers in the experiment. We compared the transcriptomes of the photoreceptor, RPE and choroidal cells using a single platform. We deduced that at least 114 genes are specifically expressed in the RPE and we describe their corresponding pathways.

Results

We performed microarray analyses on RNA specifically isolated from RPE, photoreceptors and choroid cells from healthy human donor eyes. Recently, we defined the RPE transcriptome by measuring gene expression levels and identifying functional properties of the RPE [3]. The current study provides a further specification of the RPE transcriptome by identifying genes expressed at much higher levels in the RPE than in either adjacent cell layer, the photoreceptors and the choroid. In addition, we expand our dataset from a 22 k microarray platform to a 44 k microarray platform, resulting in a more extensive coverage of the human genome [9], and we used more advanced bioinformatics to analyze the data.

RPE Gene Expression Compared to either Photoreceptors or Choroid

Of the 33,712 features present on the microarray (GSE20191 [10], see Text S1), 1,904 (5.6%) genes had at least 2.5-fold higher expression in the RPE than in the photoreceptors (RPE>phot) on average over three arrays (see Table 1 and Text S2). There was a significant overrepresentation of genes that encode signal proteins,

glycoproteins, secreted proteins, membrane proteins, cell adhesion proteins, extracellular matrix proteins, proteins involved in the immune response, Ca²⁺-binding and actin binding (David [11]). Functional analysis of these genes revealed an overrepresentation of genes in the following pathways: cell adhesion molecules, melanogenesis and type I diabetes mellitus.

Furthermore, 1,126 (3.3%) of 33,712 genes on the array had at least 2.5-fold higher expression in the RPE than in the choroid (RPE>chor) on average over three arrays (see Table 2 and Text S3). In this group of genes there was a significant overrepresentation of genes coding for proteins involved in vision, retinitis pigmentosa and (membrane) transport, as well as genes with a symport function and genes in the olfactory transduction pathway.

A Novel Strategy to Detect RPE-Specific Gene Expression

Until now, all RPE transcriptome studies, including our own [3], were aimed at excluding the cell layers adjacent to the RPE from the experiment in order to prevent cellular contamination and to achieve the highest RPE tissue specificity possible [3,4]. However even isolation of RPE cells by meticulous laser dissection microscopy resulted in unavoidable contamination with adjacent cell types [5,8] (this study).

In the current study however, we did not discard the adjacent cell layers as possible contaminants, but we included them as valuable resources for comparison of gene expression. Our primary objective was to further specify the expression level of genes in the RPE relative to their expression in the photoreceptors and the choroid. In this way, we deduced RPE-specific gene expression.

RPE-Specific Gene Expression

As a first analysis we identified all genes with average expression levels at least 2.5-fold higher in the RPE than in both photoreceptors and choroid. This resulted in a list of 458 entries with RPE-specific expression (see Text S4). Functional annotation showed an overrepresentation of genes involved in inositol

Table 1. Top 30 genes with highest expression in RPE compared to photoreceptors [27,30].

Gene symbol	Genbank ID	FC RPE/chor&phot	
ITGB8	BC042028		
C1orf168	AK093468	41	
COL8A1	AL359062	37	
SLC26A4	NM_000441	37	
SLC26A7	NM_052832	37	
DKFZp761G0122	AL713743	37	
CLIC6	NM_053277	34	
VASN	NM_138440	33	
SMOC2	NM_022138	31	
SLC16A12	AK124901	29	
C6orf105	NM_032744	28	
SLC6A13	BC020867	28	
PRDM16	NM_022114	28	
LGI1	NM_005097	27	
PLD5	NM_152666	26	
A_32_P114831	A_32_P114831	26	
KCNS3	NM_002252	25	
BEST1	NM_004183	23	
IL8	NM_000584	23	
TMEM27	NM_020665	23	
NTN4	NM_021229	23	
RWDD3	AK126344	23	
LRP8	NM_004631	23	
SLCO1C1	NM_017435	23	
COL8A2	NM_005202	22	
MYRIP	NM_015460	22	
GPNMB	NM_002510	20	
PLA2G7	NM_005084	20	
KCNJ13	NM_002242	19	
FAM40B	AB032996	19	

FC: fold change, average expression level in RPE compared to choroid and photoreceptors in six arrays.

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metabolism, retinol metabolism, genetic disorders and ophthalmic diseases (data not shown).

In order to illustrate the usefulness of our approach to identify RPE specific transcripts, we next employed even stricter criteria (i.e. in *all* six arrays *at least* 2.5-fold higher expression levels in the RPE than in both photoreceptors and choroid (RPE>phot&chor FC>2.5)). This yielded 114 genes (see Table 3). We deduced from our recently published data set [3] that 39 out of these 114 genes are very highly expressed in the RPE (see Table 4).

We next hypothesized that these 39 genes, given their very high expression, could easily have been detected by other means in previous studies. We compared our 39 highly expressed RPE-specific genes from the current study to a database containing 13,037 genes described to be expressed specifically in either the retina or the RPE, described by Schulz et al [4] and were able to identify 29 of our genes in their database (75%) (see Table 4). A more thorough manual search of individual studies on the expression of genes in the RPE showed that 85% of the 39 genes were previously identified to be expressed in the RPE (Table 4).

Table 2. Top 30 genes with highest expression in RPE compared to choroid [27,30].

Gene symbol	Genbank ID	FC RPE/chor&phot
RBP3	NM_002900	37
RPE65	NM_000329	24
MPP4	NM_033066	24
ELOVL4	NM_022726	23
GUCA1C	NM_005459	23
PDE6G	NM_002602	22
NEUROD1	NM_002500	20
BEST1	NM_004183	20
SLC6A13	BC020867	19
ITGB8	BC042028	19
RP1	NM_006269	19
GUCA1B	BX537393	19
PDC	NM_002597	19
CNGB3	NM_019098	18
PROM1	NM_006017	18
PRPH2	NM_000322	18
HCN1	AK094523	18
DKFZp761G0122	AL713743	17
HOOK1	AK027250	17
GNAT2	NM_005272	16
RLBP1	NM_000326	16
C6orf105	NM_032744	16
CNGA1	NM_000087	16
ABCA4	NM_000350	16
TMEM27	NM_020665	16
RWDD3	AK126344	16
OPCML	BX537377	15
NRL	NM_006177	15
C1orf168	AK093468	15
TMEM16B	NM_020373	15

Note that cellular contamination of the RPE cells may be present, identified by the *ABCA4* transcript, which is truly a photoreceptor-specific transcript [31]. FC: fold change, average expression level in RPE compared to choroid and photoreceptors in six arrays.

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For the remaining 15% (6 genes) no data was available in the literature. Using semi quantitative QPCR (s-QPCR), we confirmed RPE expression of these genes (Table 4 and Figure 2). Both ERMN (AB033015) and SLC6A20 (NM_020208) appear to be more highly expressed in RPE compared to both choroid or photoreceptors, thereby fully confirming the microarray data. TMEM27 (NM_020665) and LMO1 (NM_002315) appear also to more highly expressed in the RPE than in the choroid, but show approximate equal expression in the photoreceptors. SGK1 (NM_005627) is ubiquitously expressed in all three cells examined. The expression of SPOCK1 (NM_004598) in the RPE is rather low compared to the photoreceptors and does apparantly not confirm the microarray data.

Functional Annotation of RPE-Specific Gene Expression

We used the list of 114 RPE-specifically expressed genes for further functional annotation of the major pathways specific for the RPE. The online database DAVID did not identify any Kegg pathways in

Table 3. 114 genes with at least 2.5 fold higher RPE expression in the RPE than in both the photoreceptors and the choroid in all six microarrays, defined as RPE-specific expression.

Gene symbol	Genbank ID	FC	gene symbol	Genbank ID	FC	gene symbol	Genbank ID	FC
ITGB8	BC042028	31	SLC16A14	NM_152527	10	ACOT11	NM_147161	7
C1orf168	AK093468	28	WFDC1	NM_021197	10	CDH3	NM_001793	6
DKFZp761G0122	AL713743	27	SLC6A13	NM_016615	10	FRZB	NM_001463	6
SLC6A13	BC020867	24	CACNB2	BG428517	10	LOC439949	AY007155	6
C6orf105	NM_032744	22	SLC2A12	NM_145176	10	SERPINF1	NM_002615	6
CLIC6	NM_053277	22	SLC6A12	NM_003044	10	GPAM	NM_020918	6
BEST1	NM_004183	21	KIAA0953	AF131834	10	SPOCK1	NM_004598	6
RPE65	NM_000329	20	ADORA2B	NM_000676	10	FLJ30594	NM_153011	6
PLD5	NM_152666	19	CA14	NM_012113	10	MUPCDH	NM_031264	6
TMEM27	NM_020665	19	PNPLA3	NM_025225	10	C1orf168	AK125198	6
RWDD3	AK126344	19	RGR	NM_002921	10	CLDN19	NM_148960	6
LRP8	NM_004631	19	STRA6	NM_022369	9	LMO1	NM_002315	6
LGI1	NM_005097	18	C7orf46	NM_199136	9	GLDC	NM_000170	6
SLC16A12	AK124901	17	KIRREL2	NM_199180	9	A_24_P186746	A_24_P186746	5
FAM40B	AB032996	17	RDH5	NM_002905	9	RDH11	NM_016026	5
PRDM16	NM_022114	16	BMP4	NM_001202	9	SFRP5	NM_003015	5
MYRIP	NM_015460	16	TMEM56	NM_152487	9	SGK1	NM_005627	5
BMP7	NM_001719	15	THC1892753	THC1892753	9	KRT18	NM_000224	5
ERMN	AB033015	14	CNKSR3	NM_173515	9	OPHN1	NM_002547	5
SLC13A3	NM_022829	14	CCNO	NM_021147	8	TDRD9	NM_153046	5
SLCO1C1	NM_017435	14	RDH10	NM_172037	8	EZR	NM_003379	5
LRAT	NM_004744	13	PBX4	NM_025245	8	FAM40B	BC019064	5
OPCML	BX537377	13	SKIP	NM_130766	8	C7orf46	BC042034	5
RLBP1	NM_000326	13	SLC7A10	NM_019849	8	CTSD	AK022293	5
TRPM3	NM_206948	13	CXCL14	NM_004887	8	DHCR7	NM_001360	4
KCTD4	NM_198404	13	A_24_P234871	A_24_P234871	8	ITGAV	NM_002210	4
THC1934449	THC1934449	13	COL20A1	NM_020882	8	GALNT11	NM_022087	4
LRP8	NM_017522	12	LHX2	NM_004789	8	THC1967593	THC1967593	4
SLC39A12	NM_152725	12	C1QTNF5	NM_015645	7	LOC650392	BC036550	4
DUSP6	NM_001946	12	SLC22A8	NM_004254	7	HPD	NM_002150	4
ERMN	BC026345	11	ROBO2	AK074780	7	BCAT1	NM_005504	4
SLCO1A2	NM_005075	11	THC1970019	THC1970019	7	A_23_P122650	A_23_P122650	4
RBP1	NM_002899	11	ADAD2	NM_139174	7	A_32_P226525	A_32_P226525	4
SLC16A3	NM_004207	11	OR51E2	NM_030774	7	PCP4	NM_006198	4
CNDP1	NM_032649	11	SLC6A20	NM_020208	7	A_32_P112546	A_32_P112546	4
SLCO1A2	NM_134431	11	SLC16A8	NM_013356	7	KIAA1576	NM_020927	4
THC1839330	THC1839330	11	THC2004763	THC2004763	7	A_23_P73096	A_23_P73096	4
SULF1	NM_015170	11	A 24 P109661	A_24_P109661	7	BASP1	NM_006317	3

FC: fold change, average expression level in RPE compared to choroid and photoreceptors in six arrays. doi:10.1371/journal.pone.0009341.t003

this group of genes. However, among the 95 genes recognized by the Ingenuity software, there was a significant overrepresentation of genes in the following functional categories: symport, membrane transport, vision, glycoprotein, transport (p<0.0001). Moreover, using the Ingenuity database we identified three canonical pathways (see Figure 3): RAR-activation (Figure 4), retinol metabolism (Figure 5) and GABA receptor signaling (Figure 6). We found four biological functions with a significant overrepresentation of genes: ophthalmic disease, visual system development and function, genetic disorder and nervous system development and function (see Figure 7).

Finally, we identified four networks correlated with our RPE-specific gene list (see Figures 8, 9, 10, and 11).

Similar functional annotations were found for the group of 39 highly expressed RPE-specific genes (data not shown). Upon closer inspection of this group we also identified eight (almost one in five!) known retinal disease genes (BEST1 [NM_004183], C1QTNF5 [NM_015645], CDH3 [NM_001793], LRAT [NM_004744], RDH5 [NM_002905], RDH11 [NM_016026], RGR [NM_002921], RLBP1 [NM_000326] and STRA6 [NM_022369]). Fifteen entries represented membrane bound or transmembrane genes.

Table 4. 39 RPE-specific genes with high expression levels as determined in our previous study [3].

gene symbol	Genbank ID	FC RPE/ chor&phot	RPE expression in literature	ref	review Schulz
C6orf105	NM_032744	22	microarray	[32]	
BEST1	NM_004183	21	immunohistochemical staining	[33]	
TMEM27	NM_020665	19	this study, Figure 11		
LRP8	NM_004631	19	microarray	[32]	1
LGI1	NM_005097	18	review, exclusively in RPE studies	[4]	1
FAM40B	AB032996	17	cDNA clones	[34]	
ERMN	AB033015	14	this study, Figure 11		1
LRAT	NM_004744	13	western blot, northern blot	[35]	
RLBP1	NM_000326	13	fluorescence immunocytochemistry	[36]	1
DUSP6	NM_001946	12	est database	[37]	1
RBP1	NM_002899	11	review, genes expressed in retina/RPE	[4]	1
SLC16A3	NM_004207	11	immunofluorescence	[38]	1
WFDC1	NM_021197	10	immunocytochemistry, microarray	[8,39]	1
KIAA0953	AF131834	10	review, genes expressed in retina/RPE	[4]	
CA14	NM_012113	10	immunocytochemistry	[40]	1
RGR	NM_002921	10	microarray	[32]	1
STRA6	NM_022369	9	immunohistochemistry	[41]	
RDH5	NM_002905	9	northern blot	[20]	1
BMP4	NM_001202	9	RNAse protection assay	[42]	1
CXCL14	NM_004887	8	microarray, RT-PCR	[39]	1
LHX2	NM_004789	8	in situ hybridization	[43]	1
C1QTNF5	NM_015645	7	cDNA library, RT-PCR	[44,45]	1
SLC6A20	NM_020208	7	this study, Figure 11		
SLC16A8	NM_013356	7	PCR	[46]	1
CDH3	NM_001793	6	western blot	[47]	1
FRZB	NM_001463	6	review, genes expressed in retina/RPE	[4]	1
SERPINF1	NM_002615	6	amino acid sequencing	[48]	1
SPOCK1	NM_004598	6	this study, Figure 11		
LMO1	NM_002315	6	this study, Figure 11		
RDH11	NM_016026	5	bovine and monkey, immunohistochemistry and in situ hybridization	[21]	1
SFRP5	NM_003015	5	northern blot	[49]	1
SGK1	NM_005627	5	this study, Figure 11		1
KRT18	NM_000224	5	CNV RPE RT-PCR	[50]	1
EZR	NM_003379	5	rat immunofluorescence and immunoelectron microscopy	[51]	1
DHCR7	NM_001360	4	microarray	[32]	1
ITGAV	NM_002210	4	microarray	[32]	1
GALNT11	NM_022087	4	review, genes expressed in retina/RPE	[4]	1
PCP4	NM_006198	4	review, genes expressed in retina/RPE	[4]	1
BASP1	NM_006317	3	microarray	[32]	1

Column four and five show 33 of the 39 genes (85%) were previously described (RNA or protein level) in individual studies in the literature using several different techniques to have RPE expression. Column six shows that 29 of the 39 genes (74%) were also present in a study by Schulz et al. [4] reporting on 13,037 genes expressed in the retina/RPE (their supplementary table 2) ref: literature reference, FC: fold change, average expression level in RPE compared to choroid and photoreceptors in six arrays, 1 in column six: present. doi:10.1371/journal.pone.0009341.t004

Discussion

Study Design

In this study we provide a further specification of the RPE transcriptome, by analyzing the genes expressed in the RPE in reference to their expression in photoreceptors and choroid. We performed functional analyses on our microarray data

using the online database DAVID and Ingenuity software in order to categorize the data and identify important functional properties in 114 genes specifically expressed in the RPE. Moreover, we identified 39 genes with RPE-specific expression and high expression levels in the RPE and in 85% of genes we were able to confirm RPE expression using the literature.

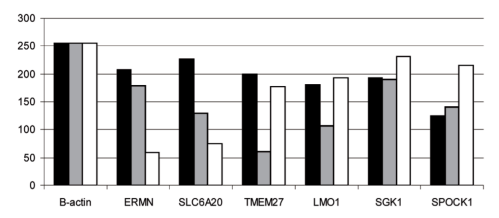


Figure 2. Confirmation of microarray results by s-QPCR. Beta actin, a household gene, was used to normalize gene expression in between all cells of the retina. The black bars indicate RPE expression levels, the grey bars indicate choroid expression and the empty bars indicate photoreceptor expression. For all genes RPE expression was shown in the microarray. See also text and Table 4. doi:10.1371/journal.pone.0009341.g002

For the remaing 15%, sQPCRs were carried out which in most cases confirmed RPE enriched expression. Our current study design was focused on comparative gene expression, genes with high expression levels in two or multiple tissues were not included.

As discussed extensively elsewhere [5,8], some degree of cellular contamination by applying LDM procedures to the retina is unavoidable. An important issue in this study was if we could overcome the cellular contamination problem by comparing the expression profiles of three adjacent cellular monolayers, in other words, did we succeed in identifying RPE specific transcripts? A number of considerations are important: 1) Obviously, if we would choose our comparative criteria even more strict (for example

expression RPE >20 x than in photoreceptor or choroid, versus current criteria RPE >2.5 x than in photoreceptor or choroid), we would obtain an even more specific RPE expressed data set. 2) In our current dataset of 114 RPE genes, we did not find any obvious contamination of highly expressed (known) photoreceptor transcripts, like opsins. 3) The majority of the 114 genes interact with each other via a limited number of molecular pathways and networks, with functional annotations which more or less can can be attributed to the RPE. This further confirms the RPE specific origin of these transcripts. For the majority of completely unknown genes, our sQPCR data confirmed the microarray data. Only one entry (SPOCK1) showed apparent higher photoreceptor expression than the RPE equivalent, and did not confirm the microarray

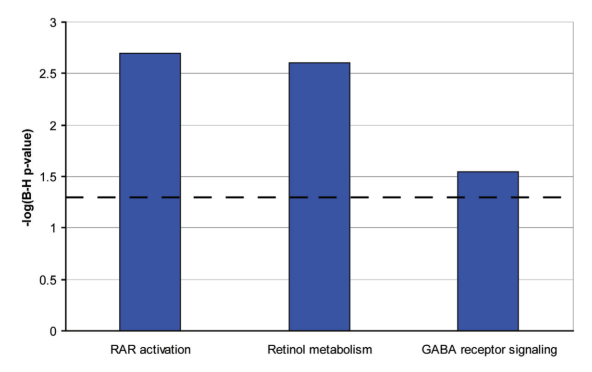


Figure 3. Three canonical pathways identified by Ingenuity software [25] in the group of RPE-specific genes. The three bars represent the canonical pathway identified, the x-axis identifies the pathways. The y-axis shows the —log of the Benjamini-Hochberg (B—H) p-value. The dotted line represents the threshold above which there are statistically significantly more genes in a pathway than expected by chance. doi:10.1371/journal.pone.0009341.q003

RAR Activation

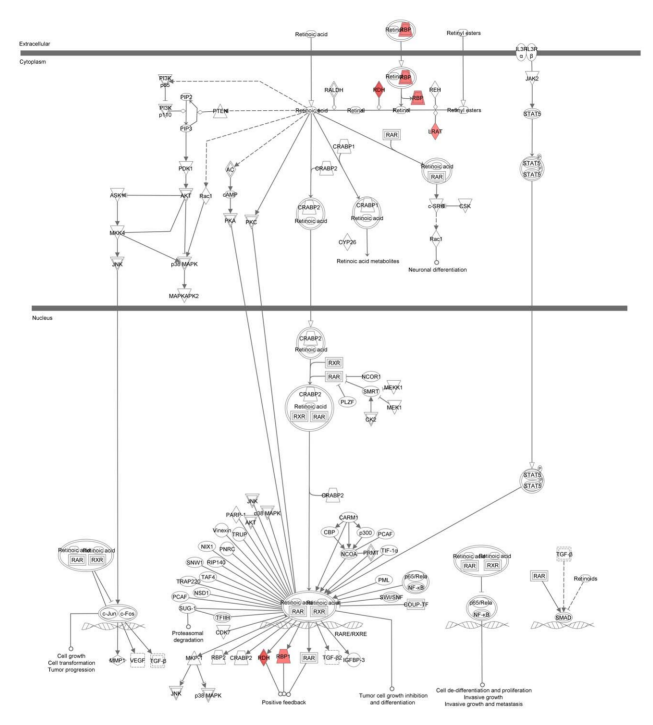


Figure 4. The RAR-activation pathway identified by the Ingenuity software. This is one of the canonical pathways that contains statistically significantly more genes than expected by chance in the group of 114 genes with RPE-specific expression. This figure shows the proteins corresponding to the overrepresented genes. Colored fields indicate their presence among the 114 genes with RPE-specific expression, uncolored genes are added by the software to form pathways. Solid lines between molecules indicate direct physical relationships between molecules (such as regulating and interacting protein domains); dotted lines indicate indirect functional relationships (such as co-regulation of expression of both genes in cell lines). doi:10.1371/journal.pone.0009341.g004

data. The reason for this discrepancy remains to be elucidated. All data taken together, RPE specificity for the majority of 114 transcripts identified is highly likely; better than previously, although some minor degree of contamination can still not be excluded.

Functional Properties of the RPE

RPE versus photoreceptors only. Compared to the photoreceptors, the RPE cells express genes from three different functional categories at significantly higher levels. The first is cell adhesion molecules. The RPE represents the outer blood-retina

Retinol Metabolism

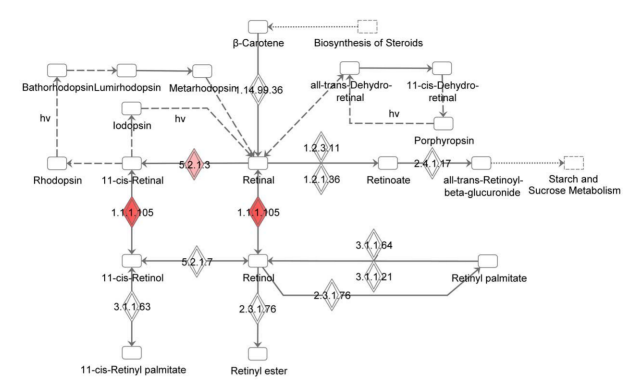


Figure 5. The retinol metabolism pathway identified by the Ingenuity software. This is one of the canonical pathways that contains statistically significantly more genes than expected by chance in the group of 114 genes with RPE-specific expression. This figure shows the proteins corresponding to the overrepresented genes. Colored symbols indicate their presence among the 114 genes with RPE-specific expression, uncolored genes are added by the software to form pathways. Solid lines between molecules indicate direct physical relationships between molecules (such as regulating and interacting protein domains); dotted lines indicate indirect functional relationships (such as co-regulation of expression of both genes in cell lines). doi:10.1371/journal.pone.0009341.g005

barrier (BRB) and this group of molecules most likely illustrated the importance for the RPE to adhere firmly to BM in order to maintain the integrity of the barrier. The second pathway is the melanogenesis pathway uniquely present in the RPE. Indeed, in the heavily pigmented RPE, the pigment granules protect against oxidative stress [12]. The third RPE pathway is the type I diabetes mellitus pathway. Twelve out of 13 genes are members of the major histocompatibility complex (MHC). This pathway was also present among the genes with highly variable expression levels in the RPE in our previous study [3]. The MHC genes are responsible for antigen presentation and are implicated in the RPE specific immune response in both health and disease [13].

In addition to the Kegg pathways described above, there were genes overrepresented in a number of functional categories, related to major functions of the RPE. There were secreted proteins, proteins involved in signaling and Ca²⁺-binding proteins. It is well known that many RPE-specific Ca²⁺ channels are involved in intracellular signaling, cellular signal transduction and the regulation of secretion of various factors [14].

Additional functional categories included membrane proteins, cell adhesion proteins and extracellular matrix (ECM) proteins, that correlate with the structural role of the RPE and its interaction with BM [1,15].

Finally, there was an overrepresentation of glycoproteins. Glycoproteins are crucial for the phagocytosis of photoreceptor outer segments [16].

RPE versus choroid only. In the group of genes with expression levels higher in the RPE than in the choroid, there was an

overrepresentation of genes in the olfactory transduction pathway. This pathway contains mainly guanylate cyclases and calcium/calmodulin-dependent protein kinases, naturally present in such highly active cells as the RPE. In addition, there was a high abundance of genes involved in vision and transport. Obviously, the RPE plays a dominant role in the transport of many signaling molecules and in the transport of waste material from the photoreceptor cells.

Genes with RPE-specific expression. We functionally annotated the 114 genes with RPE-specific expression using both David and Ingenuity software. As expected, both programs yielded vision, visual and nervous system development and function, ophthalmic disease and genetic disorder as significantly overrepresented groups of genes. Indeed, most of the genes in which mutations lead to retinal disorders are genes with high and specific expression in either RPE or photoreceptors [3].

Using David we also found an overrepresentation of genes involved in sym- or transport. This most likely represents one of the major functions of the RPE, which is the transport of biomolecules from the choroid toward the photoreceptors and vice versa. The photoreceptors heavily depend on nutrients, oxygen, hormones, etc. from the bloodstream. Meanwhile, waste products like oxidized cholesterol, visual cycle intermediates and excess water leave the retina through the RPE [1].

The Ingenuity database also revealed three canonical pathways, RAR-activation, retinol metabolism and GABA receptor signaling. Binding of retinoic acid to the retinoic acid receptor (RAR), leads to tissue-specific activation or suppression of downstream transcription [17,18]. In mature RPE, this process is invaluable for

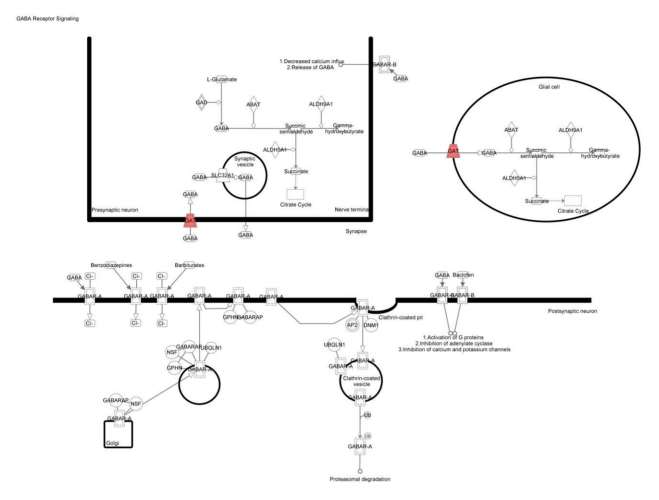


Figure 6. The GABA receptor signaling pathway identified by the Ingenuity software. This is one of the canonical pathways that contains statistically significantly more genes than expected by chance in the group of 114 genes with RPE-specific expression. This figure shows the proteins corresponding to the overrepresented genes. Colored symbols indicate their presence among the 114 genes with RPE-specific expression, uncolored genes are added by the software to form pathways. Solid lines between molecules indicate direct physical relationships between molecules (such as regulating and interacting protein domains); dotted lines indicate indirect functional relationships (such as co-regulation of expression of both genes in cell lines). doi:10.1371/journal.pone.0009341.g006

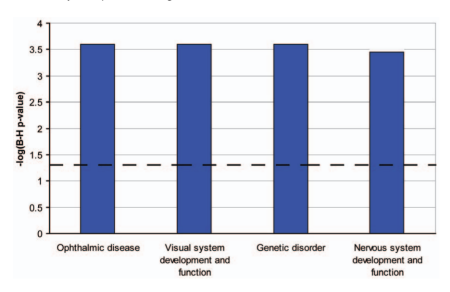


Figure 7. Four most significant biological functions identified by Ingenuity software [25] in the group of RPE-specific genes. The four bars represent the canonical pathway identified, the x-axis identifies the pathways. The y-axis shows the —log of the Benjamini-Hochberg (B—H) p-value. The dotted line represents the threshold above which there are statistically significantly more genes in a pathway than expected by chance. doi:10.1371/journal.pone.0009341.g007

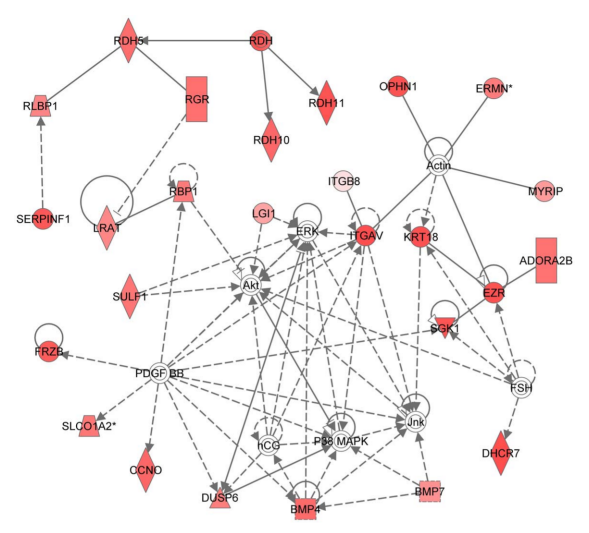


Figure 8. Most significant molecular network generated by the Ingenuity software. Network is generated from our dataset with RPE-specific expression (114 entries; see text). Note that the colored symbols represent gene entries that occur in our data set, while the transparent entries are molecules from the knowledge database, inserted to connect all relevant molecules in a single network. Solid lines between molecules indicate direct physical relationships between molecules (such as regulating and interacting protein domains); dotted lines indicate indirect functional relationships (such as co-regulation of expression of both genes in cell lines). Abbreviation of gene names are according to standard abbreviations used in Genbank [29]. The main functionalities given by Ingenuity for this molecular network are Nervous system development and function, Visual system development and function, organismal development. This network overlaps with the network in Figure 2). Highlights in this network include: (a) the regulating role of the MAPK/ERK pathway, a very complex signal transduction pathway that couples intracellular responses to the binding of growth factors to cell surface receptors; (b) platelet-derived growth factor (PDGF BB), known to induce RPE cell proliferation and migration and the development of proliferative vitreoretinopathy (PVR), acts indirectly on multiple molecules in this network, and c) the RPE retinol metabolism is present in the periphery of this molecular network.

maintenance of differentiation and homeostasis [17]. The significant presence of this pathway may thus be explained by the need of the RPE to counteract local insults (oxidative stress, lipid digestion, lipofuscin accumulation) and maintain homeostasis

The retinol metabolism pathway contained the *RDH5* [NM_002905], *RDH10* [NM_172037] and *RDH11* [NM_016026] genes, all three genes are involved in the RPE part of the visual cycle [19]. It is well known that the RPE converts all-*trans* retinoids to 11-cis isomers. More specifically, *RDH5* [NM_002905] and *RDH11* [NM_016026] convert 11-cis retinol to 11-cis retinal, while *RDH10* [NM_172037] converts 11-cis retina to all-*trans* retinal [19–22].

Finally, GABA is an important inhibitory neurotransmitter from the GABA receptor signaling pathway (Grsp) present in both brain and retina [23]. This pathway is involved in the retinal reuptake of GABA from the subretinal space. Interestingly, at least one protein from this pathway (SLC6A12 [NM_003044]) was previously observed in the rat and bullfrog RPE [23].

Comparison of RPE-Expressed Genes to the Literature

Previous studies, combined into one review, claimed to reveal 246 genes to be expressed exclusively in the RPE [4]. Strikingly, in the current study, we found that 23 out of these 246 genes (9%) had higher expression levels in the photoreceptors than in the RPE. In addition, 72 of the 246 genes (29%) had higher expression levels in the choroid than the RPE. This indicates that at least a certain level of contamination is present in the RPE signal of previously performed studies. Consequently, care should be taken when interpreting these results.

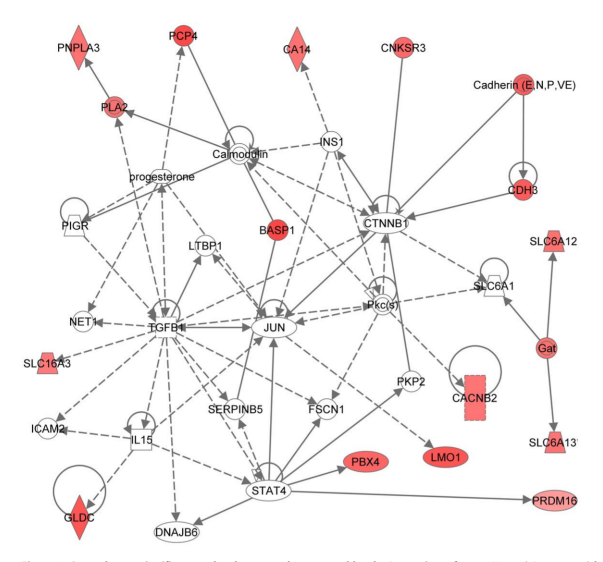


Figure 9. Second most significant molecular network generated by the Ingenuity software. Network is generated from our microarray dataset with RPE-specific expression (114 entries; see text). For explanation of symbols on the diagrams see legend Figure 7. The main functionalities given by Ingenuity for this molecular network are cellular development, hematological system development and function and connective tissue development and function. Highlights in this network include: (a) The dual presence of GABA receptors *SLC6A12* [NM_003044] and *SLC6A13* [NM_016615]; the presence and interactions of (b) the insulin-1 (INS1) protein and (c) the hormone progesterone. doi:10.1371/journal.pone.0009341.g009

Conclusions

Our study provides a detailed description of RPE-specific gene expression, as compared to both adjacent cell layers, photoreceptors and the choroid. In addition we provide a detailed functional analysis of the functional properties of the RPE-specific genes. We show the involvement of the RPE in RAR-activation, retinol metabolism and GABA receptor signaling. Moreover, for 85% of the genes we call RPE-specific with high expression levels, we could more or less verify our results using the literature. Finally, we added a substantial number of new genes significantly expressed in the RPE.

Methods

Human Donor Eyes

This study was performed in agreement with the declaration of Helsinki on the use of human material for research. Material used in this study was provided to us by the Corneabank Amsterdam. In accordance with Dutch law, the Corneabank ensured none of the donors objected to the use of their eyes for scientific purposes. Approval of the medical ethics committee was not required as data were analyzed anonymously. A detailed description of our methods can be found elsewhere [3]. In brief, we selected five eyes from five human postmortem donors. Globes were enucleated between 16 and 22 hours post mortem and frozen several hours later according to a standard protocol. Donors were aged 63 to 78 years at time of death. We chose older donors in order to minimize the likelihood of the presence of yet undiagnosed monogenic eye diseases. The donors died of cardiovascular or cerebrovascular causes or of chronic obstructive pulmonary disease. Donors did not have a known ophthalmic disorder. Visual examination and histological examination, including periodic acid Schiff (PAS) staining, indicated no retinal pathology. Three eyes were selected for the analysis of RPE vs. choroid, due to limited tissue availability only one of these eyes was also used for the analysis of RPE vs. photoreceptors. For the second and third comparison of RPE vs. photoreceptors, two additional eyes were selected.

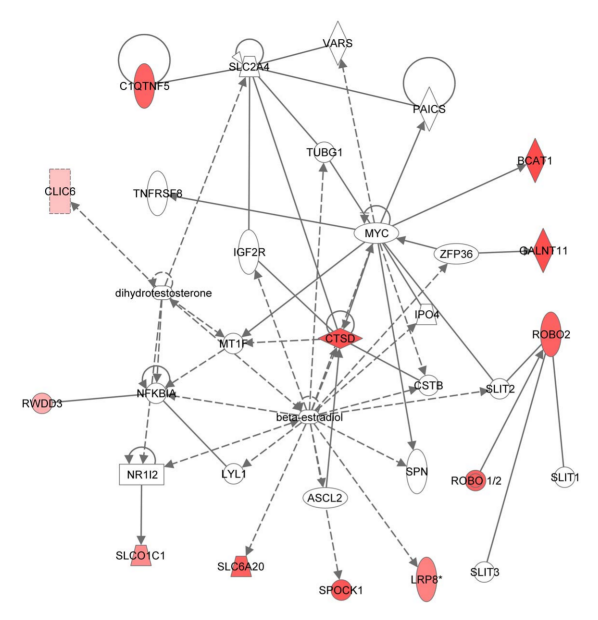


Figure 10. Third significant molecular network generated by Ingenuity. Network is generated from our microarray dataset with RPE-specific expression (114 entries; see text). For explanation of symbols on the diagram see legend Figure 7. The main functionalities given by Ingenuity for this molecular network are cellular movement, nervous system development and function and gene expression. Please note the central signaling role of beta-estradiol; known to affect retinal function and disease. doi:10.1371/journal.pone.0009341.g010

Cell Sampling

Globes were snap-frozen and stored at -80° C until use. A macular fragment of 16 mm² with the fovea in its centre was cut from each of the retinas, as described previously [8]. For each eye, multiple cryosections were stained with periodic-acid Schiff and microscopically examined for abnormalities. Twenty μ m sections from the macular areas were used for the isolation of choroid, RPE cells and photoreceptor cells. A Cresyl Violet staining (LCM Staining Kit, Ambion) was applied to the sections intended for the isolation of photoreceptor cells, according to the manufacturer's protocol. No staining was applied to sections to be used for the isolation of RPE cells or choroid. All sections were dehydrated with ethanol and air-dried before microdissection with a Laser Microdissection System (PALM, Bernried, Germany) (Figure 12). Cells were stored at -80° Celsius.

RNA Isolation and Amplification

Total RNA was isolated and the mRNA component was amplified [8]. Amplified RNA (aRNA) was quantified with a nanodrop (Isogen Life Science B.V., The Netherlands) and the quality was checked on a BioAnalyzer (Agilent Technologies, Amstelveen, The Netherlands). Subsequently, aRNA samples were labeled with either a Cy3 or a Cy5 fluorescent probe.

Microarray Design and Handling

For all hybridizations a 44 k microarray was used (Agilent Technologies, Amstelveen, The Netherlands). For three of the donors, photoreceptor RNA was hybridized against RPE RNA. In addition, for three donors, choroidal RNA was hybridized against RPE RNA. Hybridization, washing and scanning were performed as described previously [8].

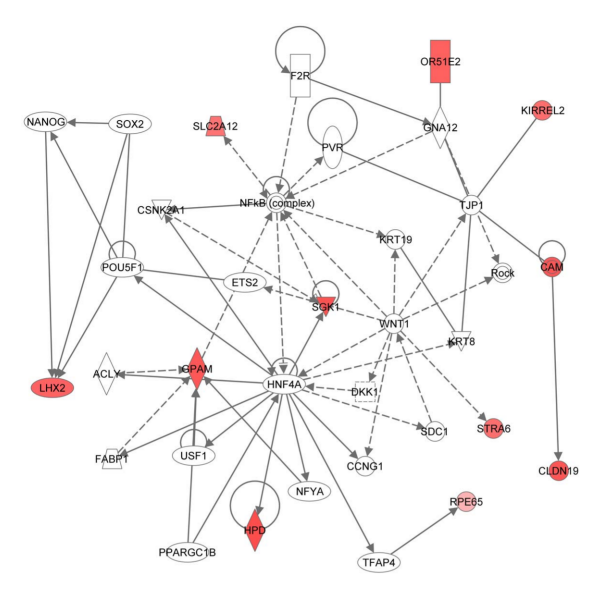


Figure 11. Fourth significant molecular network generated by Ingenuity. Network is generated from our microarray dataset with RPE-specific expression (114 entries; see text). For explanation of symbols on the diagrams see legend Figure 7. This network overlaps with the network in Figure 7. The main functionalities given by Ingenuity for this molecular network are lipid metabolism, molecular transport and nucleic acid metabolism. The highlights of this network include: The central roles for (a) the *HNF4A* transcription factor and (b) the NFkappa Beta and (c) Wnt signaling pathways. doi:10.1371/journal.pone.0009341.g011

Data Analysis

Scanned images were processed and analyzed with Feature Extraction software (v 8.5 Agilent) and Rosetta Resolver software (Rosetta Inpharmatics). All data is MIAME compliant and the raw data has been deposited in the Gene Expression Omnibus(GEO) database. For each gene we calculated either the ratio between the RPE and the photoreceptor signal, or the ratio between the RPE and the choroid signal, depending on the array. This resulted in three ratio's for the RPE versus photoreceptors and three ratio's for the RPE versus choroid for each gene. Only when all three ratio's for a single gene were greater than 2.5, we considered a gene to have a meaningfully higher gene expression (GE) in one tissue compared to the other. We chose differences in GE of at least two-fold (fold change (FC)>2.5) as cut off criterion for RPE compared to photoreceptor GE (RPE>phot), RPE compared to choroid GE (RPE>chor) and for RPE compared to photoreceptor as well as choroid GE (RPE>phot&chor). The same criteria were

applied to photoreceptor compared to RPE GE (phot>RPE) and choroid compared to RPE GE (chor>RPE). A functional analysis of Kegg pathways (Kyoto Encyclopedia of Genes and Genomes) and functional categories was performed on all groups of genes with an average FC>2.5 using the David online software [11]. Cut off criteria used were a p-value of less than 0.001 using either a Benjamini-Hochberg correction or an Ease score, a modified Fisher's exact test [11,24]. More advanced analyses of our RPEspecific genes was performed with the Ingenuity knowledge database [25] (version IPA 7.4, april 2009) yielding biological functions, canonical pathways and gene networks. We searched the literature for proof of RPE expression of our RPE-specific genes with high expression levels using the Genecards website [26] the online Mendelian inheritance in man (OMIM) website [27] and the Pubmed website [28] using the gene name combined with 'RPE' or 'retina' or 'retinal pigment epithelium' or 'expression' as search criteria.

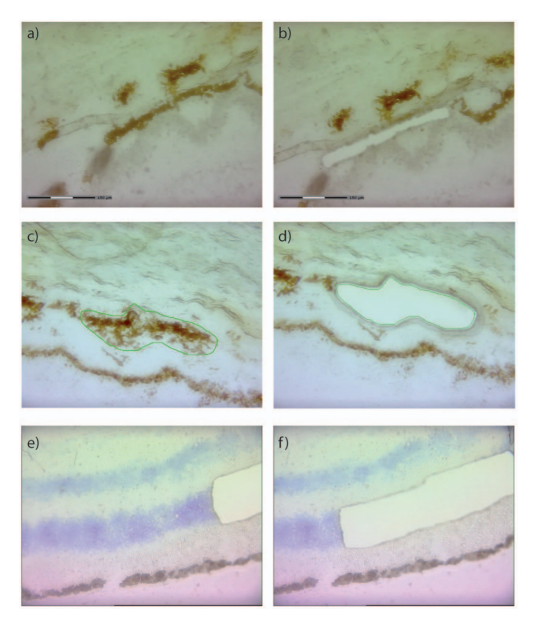


Figure 12. Frozen sections of the different cell types used in the study before and after laser dissection microscopy. Sections were used to isolate a) and b) RPE cells, c) and d) choroid, and e) and f) photoreceptor cells. Note the relatively poor morphology due to the use of frozen sections. The sections used for the isolation of photoreceptor cells were stained with cresyl violet (see methods section), the other sections were unstained. The scale can be found in figures a and b. doi:10.1371/journal.pone.0009341.g012

Confirmation of Microarray Results

For confirmation of our microarray data sQRT-PCR was used (and not QPCR) since sQRT-PCR is less sensitive for a) relatively poor RNA quality which is unavoidably obtained from human donor eyes, and b) for adjacent cell contamination of the LDM samples. Moreover, our aim was to find an approximation of expression in the RPE, choroid and photoreceptors.

sQRT-PCR was carried out, in triplicate, using exon spanning primers on RNA from LDM derived cell samples of the RPE, choroid and photoreceptors. Primer sequences are available on request. Six genes for which no further literature data on RPE gene expression was available, were tested (Table 4). B-actine, a household gene, was used to normalize gene expression in between all cells of the retina.

Supporting Information

Text S1 All features on the array. All 33,712 features present on the array.

Found at: doi:10.1371/journal.pone.0009341.s001 (2.32 MB XLS)

Text S2 All genes with higher expression in the RPE than the photoreceptors. List of all genes with expression levels 2.5 fold higher in RPE than in photoreceptors (average).

Found at: doi:10.1371/journal.pone.0009341.s002 (0.18 MB XLS)

Text S3 All genes with higher expression in the RPE than the choroid. List of all genes with expression levels 2.5 fold higher in RPE than in choroid (average).

Found at: doi:10.1371/journal.pone.0009341.s003 (0.11 MB XLS)

Text S4 All genes with RPE specific expression. List of all genes with expression levels 2.5 fold higher in RPE than in both choroid and the photoreceptors (average).

Found at: doi:10.1371/journal.pone.0009341.s004 (0.05 MB XLS)

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Author Contributions

Conceived and designed the experiments: JCB JBtB AABB. Performed the experiments: JCB JBtB AHE. Analyzed the data: JCB SS AJV AHE PvdS AABB. Contributed reagents/materials/analysis tools: SS AJV PvdS. Wrote the paper: JCB AABB. Supervised: AABB.

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