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The Genetics of Central Serous Chorioretinopathy



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Rosa Louise Schellevis

The Genetics of Central Serous Chorioretinopathy

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Voor mijn opa's en oma's

“The potential people who could have been here in my place but who will in fact never see the light of day outnumber the sand grains of Arabia. Certainly those unborn ghosts include greater poets than Keats, scientists greater than Newton. We know this because the set of possible people allowed by our DNA so massively exceeds the set of actual people. In the teeth of these stupefying odds it is you and I, in our ordinariness, that are here. We privileged few, who won the lottery of birth against all odds.”

Richard Dawkins, Unweaving the Rainbow: Science, Delusion and the Appetite for Wonder

“Life before death,
strength before weakness,
journey before destination.”

First Ideal of the Knights Radiant - The stormlight archive by Brandon Sanderson

Table of Contents

LIST OF ABBREVIATIONS.....	8
1. INTRODUCTION.....	12
2. TARGETED APPROACHES.....	30
2.1 Chronic central serous chorioretinopathy is associated with genetic variants implicated in age-related macular degeneration.....	32
2.2 Genomic copy number variations of the complement component <i>C4B</i> gene are associated with chronic Central Serous Chorioretinopathy.....	52
2.3 Association of a haplotype in the <i>NR3C2</i> gene, encoding the mineralocorticoid receptor, with chronic Central Serous Chorioretinopathy.....	66
2.4 Genetic risk factors in acute Central Serous Chorioretinopathy.....	80
3. UNBIASED APPROACHES.....	98
3.1 Role of the complement system in chronic Central Serous Chorioretinopathy: a genome-wide association study.....	100
3.2 Exome sequencing identifies <i>PIGZ</i> , <i>DUOX1</i> , <i>LAMB3</i> and <i>RSAD1</i> as susceptibility genes for chronic Central Serous Chorioretinopathy in females.....	136
4. FAMILIAL CENTRAL SEROUS CHORIORETINOPATHY.....	170
4.1 Familial Central Serous Chorioretinopathy.....	172
4.2 Exome sequencing in familial chronic Central Serous Chorioretinopathy.....	196
5. STEROID MEASUREMENTS IN CENTRAL SEROUS CHORIORETINOPATHY.....	224
5.0 Elevated steroid hormone levels in active chronic Central Serous Chorioretinopathy.....	226
6. GENERAL DISCUSSION.....	244
7. SUMMARY/SAMENVATTING - DANKWOORD - LIST OF PUBLICATIONS - CV.....	270
English Summary.....	272
Nederlandse samenvatting.....	276
Dankwoord.....	281
List of Publications.....	285
Curriculum Vitae.....	286
Phd Portfolio.....	287
Data Management page.....	289

List of abbreviations

Genes

<i>ADAMTS9</i>	ADAM metallopeptidase with thrombospondin type 1 motif 9
<i>ALX1</i>	ALX homeobox protein 1
<i>APOE</i>	Apolipoprotein E
<i>ARFGEF1</i>	Adenosine diphosphate ribosylation factor guanine nucleotide exchange factor 1
<i>ARMS2</i>	Age-related macular susceptibility 2
<i>B3GALT1</i>	Beta 3-glucosyltransferase
<i>C1QL3</i>	Complement c1q like 3
<i>C3</i>	Complement component 3
<i>C4 (A/B)</i>	Complement component 4 (A/B)
<i>C5orf63</i>	Chromosome 5 open reading frame 63
<i>CD34</i>	Cluster of differentiation 34
<i>CD46</i>	Cluster of differentiation 46
<i>CDH5</i>	Cadherin 5
<i>CETP</i>	Cholesteryl ester transfer protein
<i>CFH</i>	Complement factor H
<i>CFHR1/4</i>	CFH related 1/4
<i>CFI</i>	Complement factor I
<i>COL8A1</i>	Collagen type 8 alpha 1 chain
<i>COL10A1</i>	Collagen type 10 alpha 1 chain
<i>DUOX1</i>	Dual oxidase 1
<i>G3BP1</i>	Guanosine triphosphatase activating protein (SH3 Domain) binding protein 1
<i>GAPVD1</i>	Guanosine triphosphatase-activating protein and VPS9 domains 1
<i>GATA5</i>	GATA binding protein 5
<i>GSTM1</i>	Glutathione-S-transferase M1
<i>KCNT2</i>	Potassium sodium-activated channel subfamily T member 2
<i>IER3-DDR1</i>	Immediate early response 3-Discoidin Domain receptor tyrosine kinase 1
<i>LAMB3</i>	Laminin subunit beta 3
<i>LIPC</i>	Lipase C, hepatic type
<i>LIPH</i>	Lipase member H
<i>MEGF6</i>	Multiple epidermal growth factor like domains 6

<i>NOB1</i>	Nin one 1 binding protein 1 homolog
<i>NOP14</i>	Nucleolar protein 14
<i>NR3C1</i>	Nuclear receptor subfamily 3 group C 1
<i>NR3C2</i>	Nuclear receptor subfamily 3 group C 2
<i>OR5H14</i>	Olfactory receptor family 5 subfamily H member 14
<i>PAI-1</i>	Plasminogen activator inhibitor 1
<i>PIGZ</i>	Phosphatidylinositol glycan anchor biosynthesis class Z
<i>PITPNC1</i>	Phosphatidylinositol transfer protein cytoplasmic 1
<i>PTPRB</i>	Protein tyrosine phosphatase, receptor type B
<i>RAD51B</i>	RAD51 paralog B
<i>RCCX</i>	RP-C4-CYP21-TNX
<i>RNF144A</i>	Ring finger protein 144A
<i>RORA</i>	Retinoic acid receptor-related orphan receptor A
<i>RSL1D1</i>	Ribosomal L1 domain containing 1
<i>RSAD1</i>	Radical S-adenosyl methionine domain containing 1
<i>RSPO2</i>	R-spondin 2
<i>SGK-1</i>	Serum glucocorticoid kinase 1
<i>SKIV2L</i>	Ski2 Like RNA Helicase
<i>SLC16A8</i>	Solute carrier family 16 member 8
<i>SPATA7</i>	Spermatogenesis associated 7
<i>SYN3</i>	Synapsin 3
<i>TDGF1</i>	Teratocarcinoma-derived growth factor 1
<i>TGFBR1</i>	Transforming growth factor beta receptor 1
<i>TNFRSF10A</i>	Tumor necrosis factor receptor superfamily member 10a
<i>VEGFA</i>	Vascular endothelial growth factor A
<i>VIPR2</i>	Vasoactive intestinal peptide receptor 2
<i>ZNF713</i>	Zinc finger protein 713
<i>ZNF300P1</i>	Zinc finger protein 300 pseudogene 1

Other abbreviations

<i>17-OHP</i>	17 α -Hydroxyprogesterone
<i>ACTH</i>	Adrenocorticotrophic hormone
<i>aCSC</i>	Acute central serous chorioretinopathy
<i>aHUS</i>	Atypical hemolytic syndrome
<i>AM</i>	Adrenomedullin
<i>AMD</i>	Age-related macular degeneration
<i>BCVA</i>	Best corrected visual acuity
<i>BM</i>	Bruch's membrane
<i>Bp</i>	Base pair
<i>C3G</i>	C3 glomerulopathy
<i>CCP</i>	Complement control protein
<i>Chr</i>	Chromosome
<i>CI</i>	Confidence interval
<i>(c)CSC</i>	(chronic) central serous chorioretinopathy
<i>DHEA(S)</i>	Dehydroepiandrosterone (sulfate)
<i>DHT</i>	Dihydrotestosterone
<i>DOC</i>	Deoxycorticosterone
<i>E1</i>	Estrone
<i>E2</i>	Estradiol
<i>EDI</i>	Enhanced depth imaging
<i>EPACTS</i>	Efficient and parallelizable association container toolbox
<i>eQTL</i>	Expression quantitative trait locus
<i>ETDRS</i>	Early Treatment Diabetic Retinopathy Study
<i>EUGENDA</i>	European Genetic Database
<i>FA</i>	Fluorescein angiography
<i>FAF</i>	Fundus autofluorescence
<i>FH</i>	Factor H
<i>GC</i>	Glucocorticoids
<i>GR</i>	Glucocorticoid receptor
<i>GTE_x</i>	Genotype-tissue expression
<i>GWAS</i>	Genome-wide association study
<i>HPA</i>	Hypothalamic-pituitary-adrenal
<i>ICGA</i>	Indocyanine green angiography
<i>INDEL</i>	Insertion deletion

<i>IQR</i>	Interquartile range
<i>LE</i>	Left eye
<i>LD</i>	Linkage disequilibrium
<i>MAF</i>	Minor allele frequency
<i>MAGMA</i>	Multi-marker analysis of genomic annotation
<i>MHC</i>	Major histocompatibility complex
<i>MR</i>	Mineralocorticoid receptor
<i>MSigDB</i>	Molecular signatures database
<i>NA</i>	Not annotated
<i>NBS</i>	Nijmegen biomedical study
<i>No.</i>	Number
<i>Nr.</i>	Number
<i>OCT</i>	Optical coherence tomography
<i>OR</i>	Odds ratio
<i>PCA</i>	Principal component analysis
<i>PCV</i>	Polypoidal choroidal vasculopathy
<i>PDT</i>	Photodynamic therapy
<i>QC</i>	Quality control
<i>RE</i>	Right eye
<i>RPE</i>	Retinal pigment epithelium
<i>SCR</i>	Short consensus repeat
<i>SD</i>	Standard deviation
<i>SE</i>	Standard error
<i>SKAT(-O)</i>	Sequence Kernel Association Test (-Optimal unified test)
<i>SLE</i>	Systemic lupus erythematosus
<i>SNP</i>	Single nucleotide polymorphism
<i>SRF</i>	Subretinal fluid
<i>UTR</i>	Untranslated region
<i>VEGAS2</i>	Versatile gene-based association 2

Adapted from:

Central serous chorioretinopathy, Section I: Basics, Genetics

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Eiko K. de Jong

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1. Introduction



The human eye

Our vision is mediated by a complex process involving our eyes and the visual cortex of the brain. When light initially enters our eyes through the cornea, pupil and lens it is focused on the retina located in the back of the eye. The human retina is comprised of various specialized neuronal layers containing different types of cells. Light is converted into a biochemical signal by the photoreceptor cells through an elaborate process called the phototransduction cascade. The synapses of multiple photoreceptor cells are connected to one bipolar cell, and also have a connection with the horizontal cells, which provide lateral interaction. The signal of multiple bipolar cells in turn is transferred to the ganglion cells, which are laterally interconnected with amacrine cells. The axons on the ganglion cells make up the optic nerve, which sends the signal to the visual cortex in the brain, where it is interpreted and the image is perceived (Figure 1).^{1,2}

There are two different types of photoreceptors cells in the retina: the rods and the cones. The rods are mainly active during dim light conditions and mediate our black, white, and contrast vision, while the cones are important for color and high resolution vision. Rods are mainly present in the peripheral retina, whereas cones are more abundant in the macula, the central part of the retina which is responsible for central and sharp vision.^{1,2} To maintain proper functioning of the visual process the retinal pigment epithelial (RPE) cells located beneath the neuronal layers and the choroidal blood flow beneath the RPE are of great importance.^{4,5}

The retinal pigment epithelium (RPE)

The RPE cells encapsulate the outer segments of the photoreceptors and are connected to each other through tight-junctions (Figure 4A). Functions of the RPE cells, among others, include providing nutrients to the photoreceptor cells, preventing photo-

15

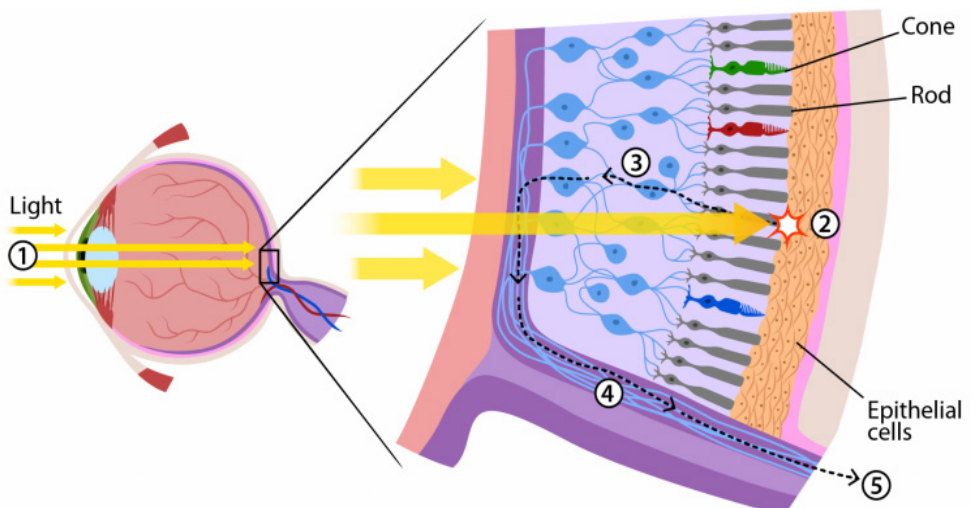


Figure 1: Schematic representation of the visual process. Light enters the eye through the lens and passes through the eye to fall on the retina (1). In the retina the light signal is received by two types of photoreceptors, the rods and cones (2) and is converted to a biochemical signal. The signals of multiple photoreceptors is passed on to a single bipolar cell, multiple bipolar cells in turn transfer their signal to a ganglion cell. (3) The axons of the ganglion cells make up the optic nerve (4) and pass on the signal to the visual cortex of the brain (5).³

oxidation, recycling waste products of the visual cycle and performing phagocytosis of the photoreceptor outer segment discs.⁴ Together with the Bruch's membrane (BM) the RPE cells form the outer blood-retina barrier. This barrier effectively controls the transport of water, ions and nutrients from the neuroretina to the underlying choroidal structure and vice versa.^{4,6,7} Waste products from the photoreceptors are transported through the subretinal space, to the RPE cells. Here the products move from the apical- to the basolateral side, pass through the Bruch's membrane (BM) and are taken up by the choriocapillaris, a fine network of permeable blood vessels that allow rapid product exchange. The choriocapillaris transports the waste products towards more distal parts of the choroid, where blood vessels are larger and no longer permeable, eventually releasing the waste products in the blood stream.⁵ Nutrients and oxygen on the other hand follow the reversed route, and are transported from the choroid through the BM and RPE to the photoreceptor cells.

Due to the large metabolic turnover of the photoreceptor cells, a large volume of water is produced and released in the subretinal space. Elimination of the water from the retina is mediated by the RPE cells through a tightly controlled ion-balance, and generates an adhesion force between the RPE and retina.⁴ The transport of chloride (Cl⁻) ions mainly drives the apical to basolateral water transport in the RPE, which is estimated to be 1.4-11 $\mu\text{L}/\text{cm}^2/\text{h}$.⁴ Multiple ion channels, ion exchangers and cotransporters are involved in this process (Figure 2).^{4,6} Additionally, aquaporin 1 molecules, which are known to transport water, are found on both the apical and basolateral side of the RPE, providing additional water efflux.⁴

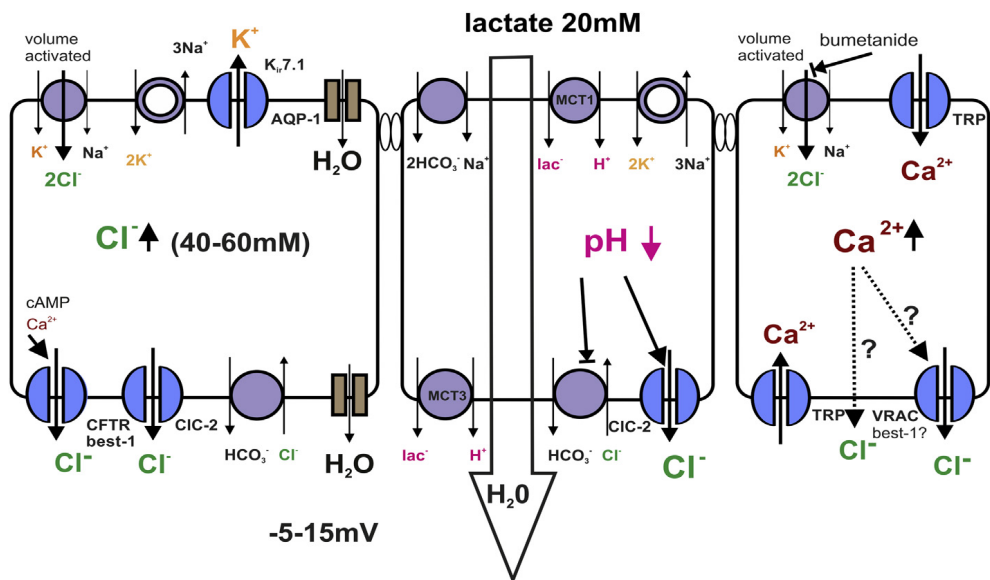


Figure 2: Summary of the mechanisms for transepithelial transport adapted from Reichart and Strauss 2014.⁶ Transepithelial transport of water and Cl⁻: A gradient for Na⁺ into the cell is provided by the activity of the apical Na⁺/K⁺-ATPase and inward rectifier K⁺ channel Kir7.1. The gradient is used to transport Cl⁻ into the cell via the activity of the Na⁺/K⁺/2Cl⁻ cotransporter, leading to an intracellular Cl⁻ concentration of 40-60 mM. Cl⁻ leaves the cell across the basolateral membrane via different types of Cl⁻ channels, osmotically driving water through aquaporins across the epithelium. AQP1: Aquaporin 1; best-1: Bestrofin 1; CIC-2: Cl channel-2 of the CIC family; CFTR: cystic fibrosis transmembrane conductance regulator; Kir7.1: inward rectifier K⁺ channel.

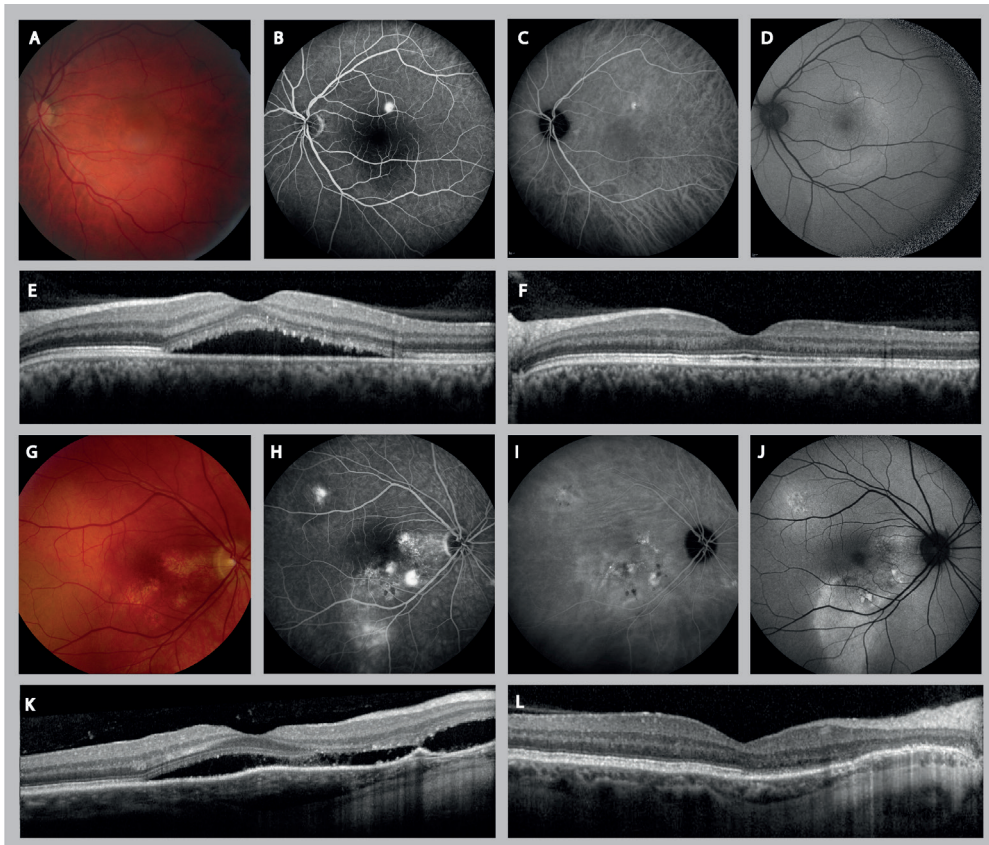


Figure 3: Clinical features of acute and chronic CSC. Multimodal imaging of the left eye of a 41-year-old male patient (A-F) with acute central serous chorioretinopathy (aCSC) and the right eye of a 40 year-old male patient (G-L) with chronic CSC (cCSC). B) Fluorescein angiography (FA) revealed a single “hot spot” of leakage and no atrophic retinal pigment epithelium (RPE) changes in the aCSC patient. C) On mid-phase indocyanine green angiography (ICGA) a small hyperfluorescent lesion was observed at the location of the “hot spot” on FA. D) Fundus autofluorescence (FAF) imaging showed granular hyper-autofluorescent changes at the site of the serous neuroretinal detachment. E) Optical coherence tomography (OCT) scan at first presentation revealed a subretinal serous fluid (SRF) accumulation, F) which resolved after four weeks. H) FA imaging in the cCSC patient revealed a large area of atrophic RPE changes and multiple leakage spots. I) ICGA imaging in this patient revealed diffuse choroidal hyperpermeability which was slightly larger than the area of leakage visible on FA. J) FAF imaging showed a mixture of intense areas of hyper-autofluorescence together with granular hypo-autofluorescent changes. K) At diagnosis, foveal SRF and a small RPE detachment were observed on the OCT scan of the cCSC patient, L) which both resolved within three weeks after treatment with half-dose photodynamic therapy. Image from chapter 2.4 of this thesis.

Central serous chorioretinopathy

One of the main defining features of central serous chorioretinopathy (CSC) is the accumulation of serous fluid underneath the retina (Figure 3E/K and Figure 4B). This subretinal fluid disrupts the interaction between the photoreceptor cells and the retinal pigment epithelium (RPE) leading to visual complaints such as blurred vision, metamorphopsia and loss of visual acuity when the subretinal fluid is present in the central part of the retina.⁸⁻¹⁰ The diagnosis of CSC is determined using combined information from different imaging modalities such as fundus photography, optical coherence tomography (OCT), fluorescein angiography (FA) and indocyanine green angiography (ICGA) (Figure 3).

Two forms of CSC can be distinguished: acute and chronic CSC. One of the discriminating factors between the two subtypes is the duration of complaints; where acute CSC (aCSC) usually spontaneously resolves within 3-6 months with minimal visual acuity loss, chronic CSC (cCSC) is characterized by prolonged fluid accumulation.^{8,11} The subtypes can be further distinguished by the number of retinal abnormalities; patients with acute CSC usually present with only a single focal leakage spot on FA (Figure 3B), whereas chronic CSC is characterized by more widespread RPE alterations or atrophy, more numerous RPE detachments and leakage points (Figure 3H) and in general a poorer visual prognosis.^{8,11} Besides RPE alterations, increased permeability of the choroid and an overall thickening of the choroid can also be observed in CSC.^{8,9,12} It is still unclear if acute and chronic CSC represent two distinct entities or whether they belong to a spectrum within the same disease.

The mechanism behind the occurrence of CSC is currently unknown. It has been hypothesized that increased pressure from the (thickened) choroid in combination with a dysfunctional pump-function of the RPE underlie the subretinal fluid accumulation.^{8,9} However, there is no consensus how both the thickened choroid and RPE dysfunction itself occur. Several clinical risk factors for CSC have been described, the most predominantly being (perceived) stress and the use of corticosteroids.¹⁰

Other risk factors include pregnancy, *Helicobacter pylori* infection, type-A personality, hypertension and Cushing's disease.^{8,13} A large meta-analysis of 17 studies on CSC risk factors was performed by Liu et al, providing an overview of the odds ratios associated with the different risk factors for CSC (Table 1).¹³

Patients with CSC usually present with their first symptoms between 40-55 years of age, when they are still professionally active and visual complaints cause major consequences for their day-to-day life.⁸ A skewed disease incidence in men compared to women is observed (9.9 cases per 100,000 men, 1.7 per 100,000 women).³⁰ Therefore, it has been suggested that the stress axis, or other hormonal imbalances may underlie the disease.^{8,31} So far, one animal study demonstrated that features of CSC, namely a thickened choroid and increased leakage, could be modeled in rats after injection of a high dose of the mineralocorticoid aldosterone.³² This animal work was translated into humans by testing mineralocorticoid receptor (MR) antagonists as a potential treatment for CSC.

Table 1. Overview of reported risk factors for CSC from Liu et al. 2016¹³

Risk factor	Meta-analysis	Meta-analysis	Studies included in
	P-value	OR (95% CI)	meta-analysis
Coronary heart disease	NS	1.31 (0.94-1.82)	14-16
Hypertension	0.0002	1.70 (1.28-2.25)	14-21
<i>Helicobacter pylori</i> infection	<0.0001	3.12 (1.81-5.40)	22-24
Steroid usage	0.0002	4.29 (2.01-9.15)	14-17,19,21,25,26
Sleeping disturbance	0.002	1.90 (1.28-1.83)	18-20
Autoimmune disease	0.0001	3.44 (1.90-6.26)	17,21,25
Psychopharmacologic medication use	0.0001	2.69 (1.63-4.45)	15-17
Type-A behavior	0.03	2.53 (1.08-5.96)	21,27,28
Gastroesophageal reflux disease	<0.05	3.29 (1.04-10.34)	17
Peptic Ulcer	<0.05	1.56 (1.30-1.88)	14,22
Organ transplantation	NS	6.30 (0.85-46.94)	17,19
Urbanization level	<0.05	0.86 (0.76-0.98)	14,19
Antihistamines	<0.05	2.93 (1.62-5.31)	17,29
Smoking	NS	1.16 (0.81-1.67)	17,29

CI: Confidence Interval; NS: Not significant; OR: Odds Ratio

Although some cases of CSC spontaneously resolve, treatment of the disease is warranted in most cases of cCSC. Currently, treatment options for CSC are limited, but most commonly used treatments are either micropulse laser or photodynamic therapy (PDT). For PDT, verteporfin is administered intravenously and a laser beam is used to activate the substance when it enters the choroid.³³ In micropulse laser treatment the RPE cells are targeted, and subjected to ultrashort laser pulses.³⁴ Although it has been postulated that PDT reduces choroidal congestion and micropulse laser stimulates RPE pumping function, the exact biological mechanism of action of both techniques is not known.^{8,33,35} Recent studies indicate a better disease outcome while using half-dose PDT compared to high-density subthreshold micropulse laser.^{36,37} Additionally, several reports have discussed the effectiveness of MR antagonists in CSC,³⁸⁻⁴⁰ but to date no randomized controlled clinical trial has been performed to properly assess superiority over other treatments in CSC.

Familial CSC - The basis of a genetic component for CSC

For CSC, reports about familial CSC occurrence go back many decades with the first report published in 1960 and the latest, to date, published in 2017.^{41,42} These studies have led to the first hints on a genetic component for CSC and form the basis for the work presented in this thesis.

In 1960, an interesting case was presented in literature about a monozygotic twin pair that almost simultaneously presented with CSC.⁴¹ The two twin brothers, 36 years of age, visited an ophthalmologist within 9 months from each other, presenting with unilateral CSC. They were treated with rest, vasodilator drugs and anti-allergics

and vision improved in both instances to normal in one brother (10/10) and almost normal in the other (8/10). Even though the brothers had experienced hardship during World War II, they were described as 'happy and cheerful men'. The author was not able to pin-point a specific current stressor in their lives, nor any commonality in their exposure to environmental circumstances that might trigger the disease. The author suggested that this particular case supports a theory in which constitutional hereditary factors may play a role in CSC.⁴¹

Eight years after this initial report of two twin brothers, the first parent-child relationship was presented. In 1968, Haik et al described the case of a 30-year old housewife from Costa Rica, who had moved to the United States with her husband, presenting with CSC.⁴³ According to the report she had experienced "unaccustomed stresses from loneliness, household duties, and long hours of study to learn English" before waking up one day with a dark spot in the vision of her left eye. Treatment with a vasodilator, tranquilizer and corticosteroids led to disappearance of most of the fluid after four weeks, while vision returned to normal after six weeks. Interestingly, nine days after initial onset of CSC in the patient, her mother also presented in the clinic with complaints of a dark spot in her left eye. The mother was described as being "unduly concerned about illnesses within her family" and she had also suffered a recent loss of uninsured property in Costa Rica around the same time.⁴³ The course of disease followed a similar pattern as that of her daughter, and after eight weeks both the fundus and visual acuity returned to normal.

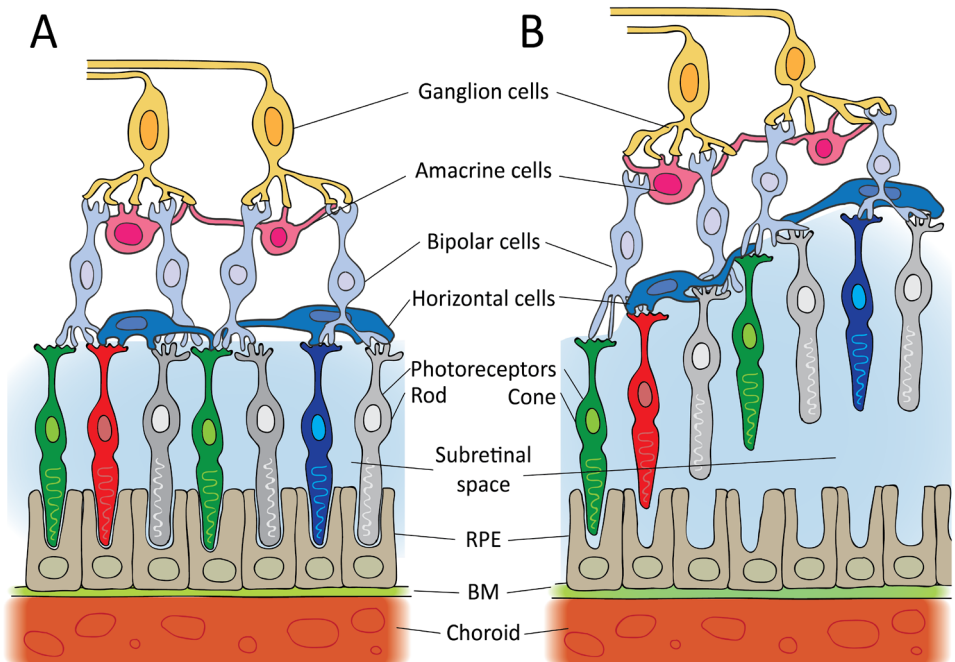


Figure 4: Graphical representation of the retinal anatomy in a normal and an eye affected with CSC. A) Normal retina, indicated are the various different cell layers involved in the visual process. B) Retina affected with CSC, fluid accumulation is present between the photoreceptor layer and RPE cells, leading to disruption of their interaction. BM: Bruch's membrane. RPE: Retinal Pigment Epithelium. Original figure courtesy of Maartje Geerlings, PhD.

During the following years scattered reports of familial occurrence emerged in literature, but in 1996, the first series of eleven families presenting with CSC were described. In each family, two to four family members were affected with CSC.⁴⁴ Most of these families contained affected siblings, but one family showed CSC in a mother and her son. Of note, in most of these patients both eyes were affected in contrast to the earlier reports where only one eye was affected. Moreover, the mean age of the patients was 44.9 years, which is in the range of the demography of cCSC patients, whereas the patients in the initial reports were quite young, suggesting that they may represent acute cases.⁴⁴

After this collection of eleven families was described, a few additional families were reported in literature,⁴⁵⁻⁴⁸ but the next major collection of families was described more than 20 years later in 2017.⁴² This study expanded on the families described in the 1996 publication of Oosterhuis et al, and now included a total of 23 families with 103 subjects.⁴² Of particular interest in this work was the long-term follow up of subjects that had been enrolled in the initial study, demonstrating that 24% of subjects that were previously unaffected showed (sub)clinical signs of CSC after 20 years.⁴² These results suggest that familial occurrence is a strong risk factor of developing the disease, underscoring the possible involvement of genetic factors.

Although these observations of familial CSC occurrence do not completely rule out other attributing factors, such as simultaneous exposure to environment triggers or circumstances that are a consequence of *nurture* rather than *nature*, they do suggest that a genetic predisposition or susceptibility to such environmental factors can be passed on through the generations, leading to a higher chance of CSC development in family members.

21

A brief history of human genetics

In 1866, Mendel discovered that “traits” are transmitted to offspring in certain reproducible patterns,⁴⁹ a discovery that can be seen as the start of the modern genetics era. However, only after the observation in 1902 that chromosomes were the carriers of these “traits” or also called genes,^{50,51} and the discovery in 1911 that certain genes are located on specific chromosomes,⁵² advancements in the genetics field really took flight. In rapid succession, the DNA double helix was discovered to be the carrier of genetic information, RNA was identified to be transcribed from DNA and proteins from RNA, establishing the central dogma of molecular biology.⁵³

Inheritance patterns

Genetic diseases can be divided in several categories based on the contribution of the genetic component to the disease. Mendelian diseases are single gene disorders that can be inherited through different inheritance patterns. In a dominant disease, a single (rare) variant in a gene (Box 1) causes the disease, and subsequently the disease is transmitted from generation to generation.^{49,54} Recessive disorders are caused by the inheritance of two disease-causing variants in a single gene from each parent, only leading to disease when both gene-copies are affected. In addition more complex inheritance patterns may occur, including sex-linked inheritance patterns, which are caused by disease-causing variants on the X- or Y-chromosomes, incomplete penetrance in which individuals that carry a disease-causing variant remain

unaffected, and digenic disorders, in which disease-causing variants in two genes underlie the disease.⁵⁴ However, the common consensus is that disorders following these inheritance patterns are caused by (rare) variants that often have a large effect on protein function, and are absent or very rare in non-affected individuals.

On the contrary, in complex diseases, such as age-related macular degeneration (AMD), the combined effect of multiple genetic variants (that are generally common in the population) and environmental risk factors determines the disease susceptibility.^{55,56} In complex diseases, the effect size of common variants on disease is represented by the odds ratios (ORs). These allelic ORs are calculated by comparing the allele (Box 1) frequency of a variant between patients and an unaffected control population. A higher allele frequency of the variant in the patient population compared to healthy individuals indicates that the variant increases risk for disease ($OR > 1$), whereas the opposite applies when a variant is more common in healthy individuals ($OR < 1$) and is thus protective for the disease.

For complex diseases, a concurrence of circumstances might cause disease such as variations in one or multiple genes or pathways, influenced by one or more environment triggers.^{55,56} These multiple layers of complexity make it challenging to study the genetic origin of complex diseases. By comparing genetic data of large cohorts of affected and unaffected individuals, genetic association studies aim to find

Box 1: Commonly used genetic terminology

Allele: Variation of a gene which can span one or multiple consecutive nucleotides. The most common occurrence of the variation is referred to as the major allele, whereas the less occurring change is indicated as the minor allele.

expression Quantitative Trait Loci (eQTLs): A locus or SNP that directly influences the expression of a close by (cis-eQTL) or distal (trans-eQTL) gene. Due to different activation of transcription factors and regulatory elements eQTLs are often tissue-specific.

Gene: Region of the genome that is translated into a functional protein. Genes consist of coding regions (exons), non-coding regions (introns), and untranslated regions (UTRs). After transcription, intronic regions are spliced out and the exonic regions are translated into amino-acids forming the functional protein.

Haplotype (block): A stretch of DNA containing multiple alleles inherited together due to the process of recombination and presence of linkage disequilibrium in the genome.

Imputation: The prediction of alleles on the same haplotype block based on the information of alleles that have been genotyped. This method can be used to greatly increase the amount of variants in a study.

Linkage disequilibrium (LD): The non-random distribution of alleles at a locus, indicating that in a population certain alleles are not independently inherited, but co-occur more often than expected by chance. This gives rise to the occurrence of haplotype blocks. The amount of LD between variants is expressed as a R^2 value from 0=no linkage to 1=perfect linkage.

Locus: Location of a gene on the genome.

Recombination: Merging and shuffling of chromosomal parts during cell division when the chromosomes are paired. During meiosis maternal and paternal chromosomes can be mixed leading to the formation of new haplotypes. The amount of recombination varies between regions of the genome.

Single-Nucleotide Polymorphism (SNP): A variation in the DNA that spans only one nucleotide and has a frequency of $> 1\%$ in the population.

variants that occur at different frequencies in patients compared to controls, and may contribute to the disease.⁵⁷

Modern genotyping methods

Initially, the only way to observe genetic abnormalities was by karyotyping, which uses microscopy and chromosomal staining to identify relatively large chromosomal abnormalities.^{58,59} Later, linkage analysis was used in families to pinpoint the approximate location of disease specific mutations. This technique uses SNP information to identify haplotype blocks segregating with the disease due to linkage disequilibrium (Box 1).⁶⁰ Later, after the development of Sanger sequencing,⁶¹ these locations could be sequenced to identify disease-causing genetic variants at the level of single nucleotides. A rapid growth of research into the genetics of diseases followed, identifying an ever growing list of causal genes in for example Huntington's disease (1983), color blindness (1986), cystic fibrosis (1989), breast cancer (1994) and many others.⁶²⁻⁶⁶

Currently, sequencing and genotyping techniques are evolving to cheaper and high-throughput approaches, allowing us to delve deeper into the genetic origins of disease. Depending on the study design, various methods can be used to determine genetic variation. In a classic candidate variant/gene approach, a specific single nucleotide variant (SNP) or gene is selected and genotyped, and its association to a disease is tested. SNPs can be genotyped with various methods including uniplex genotyping assays targeting one particular variant, or multiplex genotyping chips or arrays targeting hundreds of thousands of genetic variants simultaneously.^{67,68} The entire coding regions of genes can be sequenced with Sanger sequencing, but the field is shifting to faster and scalable new techniques, for example molecular inversion probes (MIPs) combined with next-generation sequencing.^{69,70} The candidate gene/variant-approach can be very effective when variants are known or suspected to be involved in disease. However, problems with scalability occur when a large number of variants or genes needs to be tested, or when there is no prior knowledge about the genetic loci involved in the disease.⁷¹

23

Therefore, genome-wide unbiased approaches can be used in order to test many variants and genes at the same time and to find new loci in the genome that are associated to disease.⁵⁷ Hundreds of thousands of genetic variants spread throughout the genome can be genotyped using commercially available chips or arrays.⁶⁷ Data of these (mostly) common variants can be imputed with reference panels, in which non-genotyped variants are predicted based on known inheritance patterns present in the genome (linkage disequilibrium, Box 1), increasing the data sets to millions of variants.⁷² In a genome-wide association study (GWAS) the frequency of these (mostly) common variants are compared between affected and unaffected individuals, and regions of the genome can be identified that increase susceptibility to disease.⁵⁷ As an example, the GWAS method has been very successful in identifying over 52 independent variants at 34 genomic loci associated with AMD.⁷³

Alternatively, all the coding regions of the genome (the exome) or the entire genome of an individual can be sequenced and the frequencies of both rare and common variants can be compared to healthy individuals. However, as on average every individual carries 4.1-5 million variations compared to the reference genome, of

which ~149-182 sites are protein-truncating and 10,000-12,000 variants alter an amino-acid,⁷⁴ the large datasets generated by such techniques provide a challenge to determine causality. Another challenge when using exome or genome sequencing is that due to the low frequencies of most rare variants, large cohorts are necessary to obtain sufficient statistical power to prove that a rare variant is associated with a disease. To overcome this hurdle, the cumulative effect of multiple variants in one gene can be determined using gene-based tests, or the combined effect of variants in multiple genes can be analyzed using pathway enrichment analyses.^{75,76}

Aims and outline of this thesis

Currently, the etiology of CSC is unknown and treatment options are limited. Studying the genetic architecture of a disease has the potential to uncover novel insights on the disease pathogenesis and to work towards therapies targeting the biological mechanisms underlying the disease. This approach has been followed for several other multifactorial eye diseases such as AMD, glaucoma and myopia.⁷⁷ In particular for AMD the genetic architecture of the disease has been thoroughly investigated over the past 15 years, yielding significant insights into the molecular mechanisms of AMD development.

The aim of this doctoral thesis is to investigate the genetic components contributing to cCSC development in order to increase our knowledge of the disease etiology and on the long-term to improve treatment options for patients affected with cCSC.

In **Chapter 1**, an introduction is given about the eye and the disease characteristics of aCSC and cCSC. The first familial studies on CSC are described, which provided the initial hypothesis that a genetic component may underlie CSC. A brief history of genetics is supplied along with the most commonly used techniques in genetic studies.

Section 2 of this thesis describes various candidate studies performed for CSC. In **Chapter 2.1**, previously associated AMD SNPs as well as variants in the complement factor H (*CFH*) gene are investigated in a cohort of cCSC patients. Frequencies of these genetic variants are compared between cCSC patients, healthy controls and AMD patients, and haplotypes of the *CFH* gene are constructed.

Chapter 2.2 investigates the involvement of copy number variations of the *C4* gene in cCSC. The protein encoded by this gene (*C4*) plays an important role in the complement system, and has also been shown to increase cortisol response, making it an interesting candidate gene for cCSC.

Functional variants in the glucocorticoid receptor gene (*NR3C1*) and mineralocorticoid receptor (*NR3C2*) are investigated in cCSC patients in **Chapter 2.3**. Additionally, haplotypes of both genes are constructed.

In **Chapter 2.4** all variants that were previously associated with cCSC are investigated in patients with aCSC. Additionally, minor allele frequencies of the selected variants were compared between cCSC and aCSC patients.

In order to identify new loci in the genome implicated in cCSC **Section 3** focuses on unbiased case-control studies for cCSC.

Chapter 3.1 describes the first GWAS study conducted on a large cohort of cCSC patients and population controls. Genetic variants throughout the genome are compared between these groups and information from eQTL databases is used to predict gene expression levels in cCSC patients and controls to identify genes that might be differentially expressed in cCSC patients.

An exome sequencing study on cCSC patients is conducted in **Chapter 3.2**. In order to find new genes implicated in the disease, multiple variants in the same gene are grouped to perform gene-burden analyses. In addition, males and females are

analyzed separately to identify genetic associations that are sex-specific.

Section 4 focuses on familial cases of CSC.

In **Chapter 4.1**, the disease characteristics of familial CSC patients and their family members are described. A number of families described in a previous study are investigated again to provide long-term follow-up, and more families are included compared to the previous study.

Multiple families included in Chapter 4.1 are investigated on a genetic level in **Chapter 4.2**. Rare variant segregation analysis is performed for the families, and gene-based analyses of rare variants are also provided.

Chapter 5 investigates levels of 17 steroid hormones in cCSC patients with an active episode of fluid accumulation compared to health controls.

Chapter 6 summarizes the primary findings of this thesis, and provides an overview of all genetic studies performed on cCSC to date. The current knowledge on the genetics of CSC is placed into current literature and future perspectives and follow-up studies are discussed.

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2.1 Chronic central serous chorioretinopathy is associated with genetic variants implicated in age-related macular degeneration

2.2 Genomic copy number variations of the complement component *C4B* gene are associated with chronic central serous chorioretinopathy

2.3 Association of a haplotype in the *NR3C2* gene, encoding the mineralocorticoid receptor, with chronic central serous chorioretinopathy

2.4 Genetic risk factors in acute central serous chorioretinopathy

2. Targeted Approaches





2.1 Chronic central serous chorioretinopathy is associated with genetic variants implicated in age-related macular degeneration

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Abstract

Purpose: In this study, single nucleotide polymorphisms (SNPs) at 19 loci, previously associated with age-related macular degeneration (AMD), were systematically tested for association in patients with chronic central serous chorioretinopathy (cCSC). In addition, we evaluated the effect of detailed phenotyping on these genetic associations.

Design: Case-control study.

Participants: We included 292 cCSC patients, 1147 AMD patients, and 1311 control individuals.

Methods: We genotyped SNPs at 19 AMD-associated loci and 6 additional SNPs at the complement factor H (*CFH*) locus. Phenotyping of all patients was based on fundoscopy, spectral-domain optical coherence tomography, fluorescein angiography (FA), and indocyanine green angiography.

Main Outcome Measures: We measured the allele frequencies of 25 AMD-associated SNPs and *CFH* haplotype frequencies in patients with cCSC and the effect of phenotypic subdivision of cCSC on genetic associations.

Results: One SNP in *ARMS2* (rs10490924) was significant after Bonferroni correction ($P_{\text{unadjusted}} = 0.002$; odds ratio [OR] = 0.64). The SNPs at 3 other AMD loci (*CFH*, *TNFRSF10A*, *ADAMTS9*) showed a trend toward association with typical cCSC. Further analysis of the *CFH* locus identified 2 SNPs that significantly conferred increased risk for cCSC and 1 that was protective. The *CFH*-H3 haplotype was also found to be protective ($P=0.01$; OR=0.54). Using multimodal imaging, 197 patients were classified as having typical cCSC, 52 patients had unilateral abnormalities on FA that were otherwise typical of cCSC, and 43 patients had a clinical picture that could be compatible with cCSC, but with features that could also indicate other macular diseases. Significant differences of the minor allele frequencies of the tested SNPs were observed between these 3 phenotypic subgroups.

Conclusions: Chronic CSC is associated with genetic variants in *ARMS2* and *CFH*, indicating a genetic and pathophysiologic overlap between cCSC and AMD. Intriguingly, alleles in *ARMS2* and *CFH* that confer risk of AMD may be protective for cCSC, and alleles in *CFH* that are protective for AMD confer risk for cCSC. Significant differences in allele frequencies were found among the phenotypic subgroups for several SNPs, illustrating the importance of correct clinical classification.

Introduction

Central serous chorioretinopathy (CSC) is among the most common forms of macular disease in the Western world.^{1,2} It is characterized by a serous detachment of the neuroretina from the underlying retinal pigment epithelium (RPE) in the macula due to fluid leakage through a dysfunctional RPE. Clinical evidence from multimodal imaging, such as choroidal congestion and thickening and hyperpermeability of the choroid, implies an important role for choroidal abnormalities as an underlying cause for RPE dysfunction and subretinal fluid leakage in CSC.¹⁻³

Two main subtypes of CSC can be distinguished. Patients with acute CSC present with sudden and marked central vision loss. Acute CSC is characterized by a focal leakage spot on fluorescein angiography (FA), beneath a macular neurosensory retinal detachment.¹⁻⁹ This so-called hot spot indicates leakage at the level of the RPE.^{1,4,10} The acute form of CSC generally has a favorable prognosis because the accumulated subretinal fluid often subsides spontaneously within 2 to 3 months, with (near-) normal recovery of vision (Figure 1A-C).¹ In contrast, chronic CSC (cCSC) is typically not self-limiting and subretinal fluid remains present for >3 months.^{2,4,10}

Patients have more diffuse multifocal leakage on FA and indocyanine green angiography (ICGA), as well as often irregularly distributed widespread RPE changes associated with various degrees of low-grade leakage (Figure 1D-F).¹⁻¹⁰ Because of persistent serous neuroretinal detachments with progressive and irreversible photoreceptor damage, cCSC has a poorer visual prognosis than acute CSC.^{2,6,11,12}

Although the etiology of CSC is largely obscure, clinical observations point toward an association with the use of corticosteroids, hypercortisolism, stress, and type A personality.^{1,10,13} The incidence of CSC is approximately 6 times higher in men than in women,¹ although this male-to-female proportion seems to be less pronounced in cCSC and steroid-related CSC.

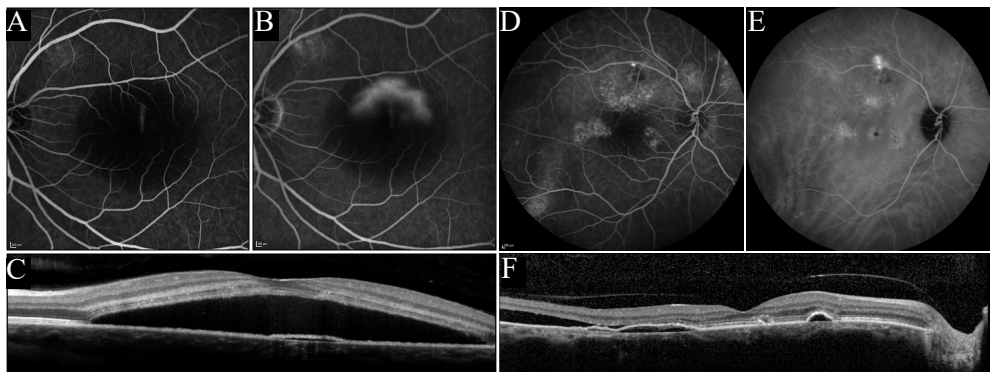


Figure 1: Examples of fluorescein angiography (FA), indocyanine green (ICG) angiography, and spectral-domain optical coherence tomography (SD OCT) in acute and chronic central serous chorioretinopathy (CSC). A+C, The left eye of a patient with classical acute CSC, showing a small focal leak in the early phase of FA and a typical smokestack fluorescein leak on late-phase FA (A, B); the corresponding SD OCT image showed a large central subretinal fluid (SRF) accumulation associated with a focal, shallow retinal pigment epithelial (RPE) detachment (C). D-E, The right eye of a patient with chronic CSC with more widespread leakage on FA (D) and diffuse hyperfluorescent areas on ICG angiography (E), as well as SRF and multiple RPE detachments on SD OCT (F).

Interracial differences in the prevalence of cCSC and the familial occurrence of cCSC suggest strong genetic involvement.^{2,14-16} However, only a limited number of possible genetic associations have been reported so far.^{17,18} Certain phenotypic features of cCSC, such as serous RPE detachments, neurosensory retinal detachment, and patchy atrophy of the RPE, are also observed in other macular diseases, such as age-related macular degeneration (AMD) and polypoidal choroidal vasculopathy.¹⁹ This phenotypic overlap suggests that a genetic overlap may also exist; AMD is known to be a multifactorial and genetically complex disorder, and 19 genetic loci have been associated with the disease.²⁰⁻²²

To assess the degree of phenotypic and genotypic overlap between cCSC and AMD, this study used a combination of detailed phenotyping based on multimodal imaging and systematic analyses of single nucleotide polymorphisms (SNPs) at the 19 known AMD-associated loci in cCSC patients. Furthermore, a detailed analysis of *complement factor H (CFH)* haplotypes in cCSC patients was performed. Finally, the effect of phenotypic differentiation on these genetic associations was evaluated.

Methods

In this study, we included 292 patients with cCSC who visited the outpatient clinic of the Department of Ophthalmology at the Radboud University Medical Center, Nijmegen, the Netherlands (Table 1). The diagnosis of cCSC and phenotyping with multimodal imaging was based on an extensive ophthalmologic examination, including funduscopy, spectral-domain optical coherence tomography (SD-OCT), FA, and ICG angiography. The definition of cCSC used in this study was based on the currently available literature, taking the following characteristics into account (all had to be present): serous subretinal fluid on OCT, 2:1 areas of multifocal diffuse leakage on FA, and corresponding hyperfluorescence on ICG angiography in 2:1 eye.¹⁻¹⁰ Patients suffering from acute CSC as recognized by a focal leakage spot or a smokestack pattern on FA, and patients with a disease period of <3 months, were excluded.¹⁻¹⁰ A total of 1311 control subjects were recruited from the blood bank of the Radboud University Medical Center, Nijmegen, the Netherlands (n = 177), and the European Genetic Database (EUGENDA, www.eugenda.org; n = 1134). For control subjects recruited from EUGENDA, ophthalmologic grading was performed excluding signs of AMD. In addition, 1147 AMD patients from the EUGENDA (www.eugenda.org) were used to calculate the minor allele frequencies of SNPs at AMD-associated loci (Table 1). The diagnosis of AMD was defined as described previously.²³ Informed consent for the use of DNA for genetic studies was obtained from all subjects. The study adhered to the tenets of the Declaration of Helsinki. Institutional review board/ethics committee approval was obtained.

Single Nucleotide Polymorphism Genotyping

Genomic DNA was extracted from peripheral blood samples using standard procedures. Nineteen SNPs at loci previously associated with AMD [(rs10490924 (ARMS2), rs12144939 (CFH), rs429608 (C2-CFB), rs2230199 (C3), rs9621532 (TIMP3), rs4420638 (APOE), rs3764261 (CETP), rs943080 (VEGFA), rs13278062 (TNFRSF10A), rs493258 (LIPC), rs10033900 (CFI), rs3812111 (COL10A1), rs13081855 (COL8A1-FILIP1L), rs3130783 (IER3-DDR1), rs8135665 (SLC16A8), rs334353 (TGFBFR1),

Table 1. Demographics of the Study Population

Characteristic	Subgroup 1	Subgroup 2	Subgroup 3	Controls	AMD
No. of subjects	197	52	43	1311	1147
Sex (male/female)	154/43	38/14	26/17	629/682	448/699
Mean age \pm SD (yrs)	53 \pm 10	55 \pm 12	57 \pm 13	66 \pm 12	76 \pm 9
Age range (yrs)	29-74	32-78	30-78	19-97	55-101

AMD=age-related macular degeneration; SD=standard deviation.

rs8017304 (*RAD51B*), rs6795735 (*ADAMTS9*), and rs9542236 (*B3GALT1*)] were genotyped in 292 cCSC patients and 1311 controls by outsourcing to the genotyping service of LGC Genomics (LGC Limited, Fordham, UK).²⁰⁻²² Genotyping of 6 *CFH* SNPs (rs1061170, rs3753394, rs800292, rs2284664, rs1329428, rs1065489) was performed with KASP genotyping assays (LGC Genomics) in 292 cCSC patients and 881 controls. The SNP-specific KASP primer mix and KASP master mix were added to 10 ng of DNA according to the manufacturer's instructions. Polymerase chain reaction amplification was performed (Veriti 384 thermal cycler; Applied Biosystems, Foster City, CA). Fluorescent FAM and HEX signals were read out with the 7900HT Fast Real-Time Polymerase Chain Reaction System (Applied Biosystems) and converted to genotype information with the SDS program (version 2.3; Applied Biosystems).

Statistical Analysis

Differences in allele frequencies were calculated using the Fisher exact test (2 sided for the 19 AMD loci and 1-sided for *CFH* SNPs, based on previous reports), and a Bonferroni correction was performed. The individual tests for the 19 AMD loci were considered significant at $P < 0.0026$, corrected for 19 tests. The significance threshold for the 6 *CFH* variants was set at $P < 0.0083$, correcting for 6 tests. Haplotypes were generated with R (version 3.0.2) and R Studio (version 0.98.501), using the haplo.stats package (version 1.6.8). For each SNP, information for both alleles was inserted into R and the haplo.cc command was used to determine associations between haplotype frequencies and cCSC. The most frequent haplotypes (frequency in controls $>5\%$) were named H1 to H5 according to Hageman et al.²⁴ Haplotypes were considered to be significantly associated to CSC if $P < 0.01$, correcting for 5 tests. Minor allele frequencies of significantly associated SNPs were compared between subgroups of CSC using a Fisher's exact test and were considered significant if $P < 0.0125$, correcting for 4 tests.

Results

Phenotyping and Classification of Chronic Central Serous Chorioretinopathy Patients

All cCSC patients were classified into 3 subgroups based on their phenotypic characteristics on multimodal imaging. Patients in phenotypic subgroup 1 (n=197) showed the most typical clinical picture of cCSC. This typical picture was defined as the presence of chronic serous subretinal fluid in ≥ 1 eye on SD OCT, bilateral irregular RPE window defects on FA with ≥ 1 "hot spot" of leakage in the affected

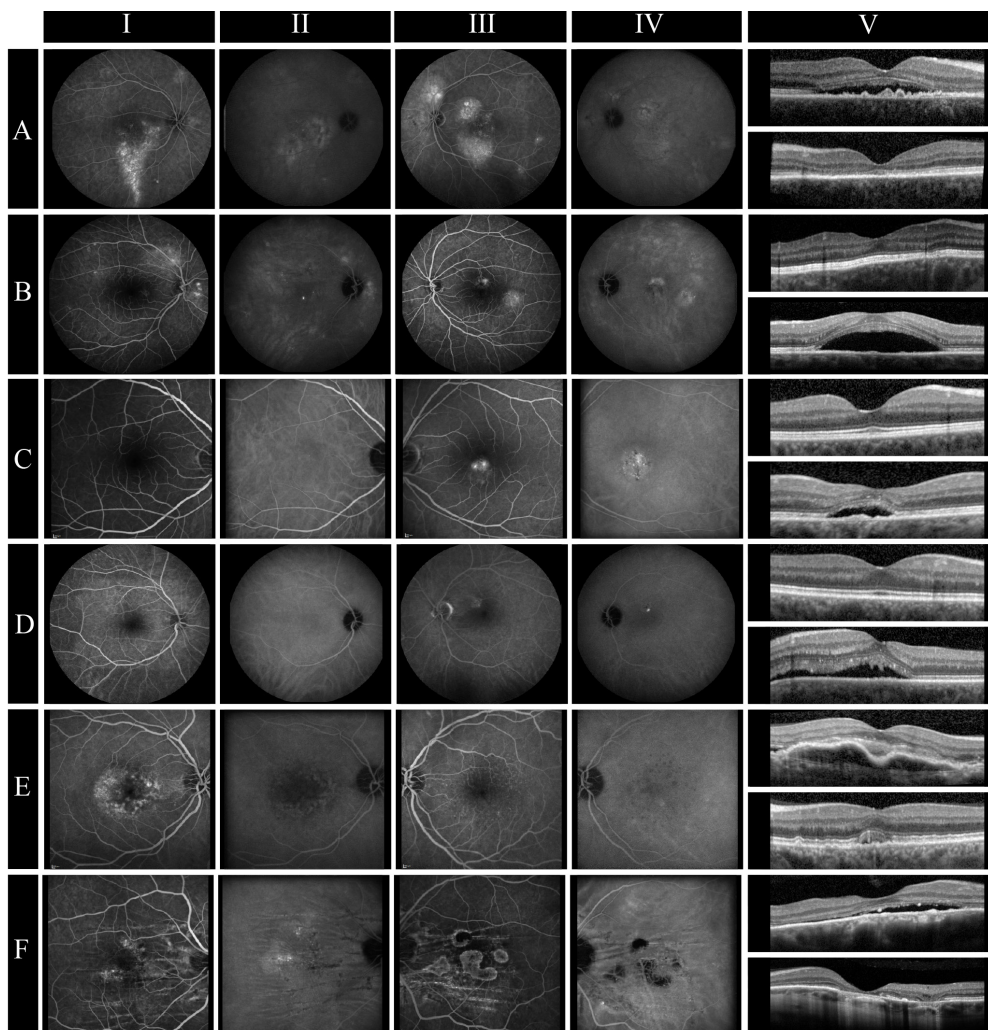


Figure 2: Examples of fluorescein angiography (FA; of right eyes [REs] in column I and left eyes [LEs] in column III), indocyanine green (ICG) angiography (REs in column II and LEs in column IV), and spectral-domain optical coherence tomography (SD OCT; of REs in upper images of column V and LEs in lower images of column V) imaging of each phenotypic subgroup. A, B, The REs and LEs of 2 patients representing subgroup 1, demonstrating diffuse areas of leakage on FA and ICG angiography (A/I-II, B/III-IV) and subretinal fluid (SRF) beneath the fovea (A/V [upper image], B/V [lower image]), illustrating typical active chronic central serous chorioretinopathy (cCSC). A/I, Classic example of a gravitational tract on FA in cCSC. The fellow eyes of these patients did not have SRF on SD OCT (A/V [lower image], B/V [upper image]) but had clear hyperfluorescent areas on FA and ICG angiography, indicating cCSC without active leakage (A/III-IV, B/I-II). C, D, Imaging of 2 patients belonging to subgroup 2 with unilateral, more localized leakage on FA and ICG angiography (C/III-IV, D/III-IV) and central SRF on SD OCT (C/V [lower image], D/V [lower image]). E, F, The REs and LEs of 2 patients as an example of subgroup 3. In the first patient, an irregular hyperfluorescent pattern was seen on FA and ICG angiography of the right eye (E/I-II). On SD OCT, there was a large retinal pigment epithelium detachment in association with serous SRF (E/V [upper image]). In the fellow eye, central mild hyperfluorescence was seen on FA, with hypofluorescence on ICG angiography (E/III-IV). Also, minimal central SRF was present on SD OCT (E/V [lower image]). The second patient had diffuse leakage on FA and ICG angiography (F/I-II) accompanied by SRF on SD OCT (F/V [upper image]) in the RE. Imaging of the LE showed multifocal areas of atrophy on FA and ICG angiography (F/III-IV) and retinal atrophy on SD OCT (F/V). Both eyes had choroidal folds on FA and ICG angiography (F/I-IV).

eye(s), and corresponding hyperfluorescent zones on ICG angiography (Figure 2A, B). No evidence of choroidal neovascularization, polypoidal choroidal vasculopathy, or other atypical findings were seen in this subgroup.

Phenotypic subgroup 2 (n=52) included patients with unilateral (instead of bilateral) abnormalities on FA that were otherwise typical for cCSC. In addition, subgroup 2 included patients with no clear “hot spot” and/or more focal leakage on FA, with an absence of subretinal fluid on SD OCT but otherwise typical FA features of cCSC. Also, patients in whom ICG angiography imaging did not show clear hyperfluorescence corresponding with FA abnormalities were included in subgroup 2 (Figure 2C, D). Furthermore, these patients showed no evidence of choroidal neovascularization, polypoidal choroidal vasculopathy, or other atypical findings.

For patients in phenotypic subgroup 3 (n=43), the clinical picture was primarily compatible with cCSC, but with more atypical features suggestive of other macular diseases; alternatively, it possibly constituted a combination of cCSC and another diagnosis. For instance, atypical clinical features in patients in subgroup 3 included evidence of choroidal neovascularization, the presence of drusen, or (highly) myopic fundus changes (Figure 2E, F).

Association of Single Nucleotide Polymorphisms at Age-related Macular Degeneration Loci with Chronic Central Serous Chorioretinopathy

To investigate the extent to which cCSC and AMD overlap genetically, the association of 19 SNPs at previously described AMD loci was tested in 197 typical cCSC patients (subgroup 1) versus 1311 controls (Table 2). Of these 19 loci, only rs10490924 in *ARMS2* remained significant ($P=0.002$; odds ratio [OR]=0.64) after correcting for multiple testing (Table 2).

Association of CFH Single Nucleotide Polymorphisms and Haplotypes with Chronic Central Serous Chorioretinopathy

To investigate the *CFH* locus in more detail, we tested 6 additional SNPs for association in 197 typical cCSC patients (subgroup 1) and 881 controls (Table 3). After correction for multiple testing, 2 *CFH* SNPs conferred an increased risk for cCSC (rs800292 [$P=7.5 \times 10^{-4}$; OR=1.50] and rs1329428 [$P=4.6 \times 10^{-4}$; OR=1.47]) and 1 SNP was protective (rs1065489; $P=0.003$; OR=0.63; Table 3).

Of the 5 observed *CFH* haplotypes, H3 (TGTCCT) was significantly associated ($P=0.01$) with cCSC. The H3 haplotype was protective against the development of cCSC (OR=0.54), being present in 15.7% of the control population and in 9% of cCSC patients. The H2 haplotype (CATTG) showed a trend toward association ($P=0.072$), conferring risk for cCSC (OR=1.33) and being present in 20.8% of control individuals and in 25.8% of cCSC patients (Table 4).

Differences in Minor Allele Frequencies of Tested Single Nucleotide Polymorphisms and Haplotypes among Phenotypic Subgroups of Chronic Central Serous Chorioretinopathy

To investigate whether classifying cCSC into 3 distinct phenotypic subgroups had an effect on the underlying genetic associations, the minor allele frequencies of the significantly associated SNPs [(rs10490924 (*ARMS2*), rs800292 (*CFH*), rs1329428

(*CFH*), and rs1065489 (*CFH*)] were compared between subgroups 1, 2, and 3 (Table 5). When comparing cCSC subgroups 1 and 2, none of the SNPs showed a difference in frequency, suggesting that these subgroups are genetically similar. When the atypical patients (subgroup 3) were compared with subgroup 1, rs10490924 in *ARMS2* ($P=0.002$) and rs800292 in *CFH* ($P=0.002$) showed a significant difference in allele frequencies between these clinical subgroups, suggesting that subgroups 1 and 3 are genetically different (Table 5).

Table 3. Complement Factor H Locus in Chronic Central Serous Chorioretinopathy

SNP	Location	Alleles (Major /Minor)	MAF in Subgroup 1 (n=197)	MAF among Controls (n=881)	Un-adjusted Allelic P Value	Allelic Odds Ratio (95% CI)
rs3753394	Promotor	C/T	0.245	0.295	0.027	0.78 (0.60-1.00)
rs800292 (I62V)	Exon 2	G/A	0.315	0.235	7.5×10^{-4}	1.50 (1.18-1.90)
rs1061170 (Y402H)	Exon 9	T/C	0.310	0.350	0.065	0.83 (0.66-1.05)
rs2284664	Intron 15	C/T	0.276	0.218	0.009	1.37 (1.07-1.76)
rs1329428	Intron 15	C/T	0.526	0.431	4.6×10^{-4}	1.47 (1.17-1.83)
rs1065489 (D936E)	Exon 18	G/T	0.118	0.176	0.003	0.63 (0.45-0.87)

CI=confidence interval; MAF=minor allele frequency; SNP=single nucleotide polymorphism. For this analysis, 1-sided P values < 0.0083 were considered significant.

Table 4. Complement Factor H Haplotypes in Chronic Central Serous Chorioretinopathy

Haplo-types	SNP						HF in Sub-group 1 (n=197)	HF among Controls (n=881)	Un-adjusted P-value	Odds Ratio (95% CI)
	rs3753394	rs800292	rs1061170	rs2284664	rs1329428	rs1065489				
H1	C	G	C	C	C	G	0.291	0.328	0.164	0.83 (0.59-1.17)
H2	C	A	T	T	T	G	0.258	0.208	0.072	1.33 (0.93-1.90)
H3	T	G	T	C	C	T	0.090	0.157	0.010	0.54 (0.32-0.91)
H4	C	G	T	C	T	G	0.125	0.132	0.480	0.96 (0.60-1.52)
H5	T	G	T	C	T	G	0.102	0.076	0.150	1.37 (0.81-2.32)

CI=confidence interval; HF=haplotype frequency; SNP=single nucleotide polymorphism. For this analysis, 1-sided P values < 0.01 were considered significant.

Table 2. Analysis of 19 Age-Related Macular Degeneration Loci in Chronic Central Serous Chorioretinopathy

Single Nucleotide Polymorphism (Locus)	Alleles (Major /Minor)	MAF in Subgroup 1 (n=197)	MAF among Controls(n= 1311)	Unadjusted AllelicP value	Allelic Odds Ratio (95% CI)
rs10490924 (ARMS2)	G/T	0.17	0.24	0.002	0.64 (0.49-0.85)
rs12144939 (CFH)	G/T	0.25	0.20	0.031	1.33 (1.03-1.70)
rs429608 (C2-CFB)	G/A	0.16	0.13	0.133	1.25 (0.94-1.68)
rs2230199(C3)	G/C	0.19	0.17	0.393	1.16 (0.83-1.61)
rs9621532 (TIMP3)	A/C	0.05	0.05	0.807	1.05 (0.65-1.68)
rs4420638 (APOE)	A/G	0.16	0.17	0.470	0.89 (0.66-1.19)
rs3764261(CETP)	G/T	0.30	0.32	0.484	0.92 (0.73-1.16)
rs943080 (VEGFA)	T/C	0.44	0.48	0.211	0.87 (0.70-1.08)
rs13278062 (TNFRSF10A)	T/G	0.40	0.48	0.004	0.73 (0.59-0.90)
rs493258 (LIPC)	C/T	0.49	0.46	0.064	1.23 (0.99-1.52)
rs10033900 (CFI)	C/T	0.50	0.48	0.514	1.08 (0.87-1.34)
rs3812111 (COL10A1)	A/T	0.34	0.36	0.498	0.92 (0.74-1.15)
rs13081855 (COL8A1-FILIP1L)	G/T	0.12	0.09	0.074	1.38 (0.98-1.93)
rs3130783 (IER3-DDR1)	A/G	0.20	0.19	0.836	1.03 (0.78-1.34)
rs8135665 (SLC16A8)	C/T	0.21	0.22	0.600	0.92 (0.71-1.20)
rs334353 (TGFBRI1)	T/G	0.26	0.25	0.662	1.06 (0.83-1.35)
rs8017304 (RAD51B)	A/G	0.36	0.38	0.467	0.92 (0.73-1.14)
rs6795735 (ADAMTS9)	C/T	0.46	0.41	0.047	1.25 (1.01-1.54)
rs9542236 (B3GALT1)	T/C	0.44	0.44	1.000	1.01 (0.81-1.25)

CI=confidence interval; MAF=minor allele frequency. For this analysis, 2-sided P values < 0.0026 were considered significant.

Discussion

Both risk-conferring and protective associations of polymorphisms in the *ARMS2* and *CFH* genes with cCSC were identified in this study. Moreover, by introducing a subclassification of cCSC, our data demonstrate clear differences as well as a certain degree of phenotypic and genetic overlap between cCSC and AMD.

Chronic CSC is a heterogeneous disease that sometimes shows a clinical overlap with other macular diseases, such as polypoidal choroidal vasculopathy and AMD. With detailed phenotyping based on SD OCT, FA, and ICG angiography, we defined 3 clinically distinctive subgroups, including typical cCSC (subgroup 1), slightly atypical cCSC (subgroup 2), and a third more atypical phenotypic subgroup of cCSC patients who had evidence of an overlap with other macular diseases (subgroup 3).

From an analysis of 19 SNPs at known AMD loci,²⁰⁻²² 1 SNP in *ARMS2* remained significant after correction for multiple testing, and 3 SNPs at other AMD loci (*CFH*, *TNFRSF10A*, *ADAMTS9*) showed a trend toward association with typical cCSC (subgroup 1). The association of these SNPs with cCSC was not exactly in concordance with the strength of the previously reported association with AMD.²⁰ Indeed, the effect sizes for *ARMS2* (rs10490924) and *CFH* (rs1061170, Y402H) are greater in AMD (ORs=2.76 and 2.25, respectively),^{20,24} whereas an inverse and weaker effect was observed in cCSC (ORs=0.64 and 0.83, respectively). A protective effect was observed for *TNFRSF10A* (rs13278062) in cCSC (OR=0.73), whereas an opposite and weaker effect was previously detected in AMD (OR=1.15).²⁰ *ADAMTS9* (rs6795735) conferred increased risk for both diseases, but with a less strong effect in AMD (OR=1.10) compared with cCSC (OR=1.37).²⁰ For some of these associations the effect was mainly carried by the males (Table 6).

43

Table 5. Comparison of Allele Frequencies among the Subgroups

SNP	Location	MAF in Sub-group 1 (n=197)	MAF in Sub-group 2 (n=52)	MAF in Sub-group 3 (n=43)	MAF among Controls (n=1311 /881)	MAF among Those With AMD (n=1147)	P-value	
							Sub-group 1 vs. Sub-group 2	Sub-group 1 vs. Sub-group 3
rs10490924 (A69S)	<i>ARMS2</i> exon 1	0.17	0.26	0.31	0.24	0.41	0.033	0.002
rs800292 (I62V)	<i>CFH</i> exon 2	0.31	0.26	0.15	0.24	0.18	0.337	0.002
rs1329428	<i>CFH</i> intron 15	0.53	0.41	0.40	0.43	N/A	0.047	0.032
rs1065489 (D936E)	<i>CFH</i> exon 18	0.12	0.16	0.12	0.18	N/A	0.246	1

AMD=age-related macular degeneration; MAF=minor allele frequency. SNP=single nucleotide polymorphism For this analysis, 2-sided P values < 0.0125 were considered significant.

However, because cCSC predominantly occurs in male patients, the small number of female patients present in our cohort made the interpretation of sex-specific associations unreliable. Genotyping of more female cCSC patients can shed light on the occurrence of a sex-specific effect and on potential differences in disease etiology.

It is well known that AMD is a multifactorial disease in which a number of subcellular systems and pathways are involved, such as the complement system, angiogenesis, lipid metabolism, and the extracellular matrix.²⁰ Our current findings suggest a partial pathophysiologic overlap of cCSC with AMD that could be connected with 2:1 of these associated pathways.

In both AMD and cCSC, the reported association with *ARMS2* (rs10490924) has the strongest effect compared with other disease-associated SNPs (OR=2.76 for AMD and OR=0.64 for cCSC).²⁰ Interestingly, the association of *ARMS2* (rs10490924) with cCSC is protective, whereas the previously reported association with AMD confers risk.^{20,25} The potential protective effect of *ARMS2* (rs10490924) in cCSC was not reported before. However, an association of *ARMS2* (rs10490924) with a lesser incidence of serous retinal detachments has been reported in polypoidal choroidal vasculopathy, a disease that shows clinical overlap with cCSC.²⁶ Intriguingly, a history of CSC is more prevalent in patients with polypoidal choroidal vasculopathy and choroidal hyperpermeability,²⁷ and CSC in combination with a type 1 neovascularization or polypoidal choroidal vasculopathy has also been described.^{28,29} The protective effect of *ARMS2* (rs10490924) in both conditions may suggest a functional involvement of *ARMS2* at the level of the RPE and/or choroid. Although the exact role of the *ARMS2* protein product is unknown, a recent study demonstrated that *ARMS2* interacts with components of the extracellular matrix.³⁰ Disturbances in the extracellular matrix at the level of the RPE and/or choroid could result in an increased susceptibility to cellular detachments, which may indicate a possible link between the supposed molecular function of *ARMS2* and the clinical phenotype of cCSC.

Our data also suggest involvement of other extracellular matrix molecules. Although these results did not remain significant after correction for multiple testing and require replication in other cohorts for confirmation, it is worthwhile to consider their association. *ADAMTS9* (rs6795735; $P=0.045$) encodes a metalloproteinase that cleaves large aggregated proteoglycans, has thrombospondin domains, and is an inhibitor of angiogenesis.³¹ *COL8A1* (rs13081855; $P=0.074$) encodes collagen type VIII that is expressed in the endothelium of blood vessels and is involved in endothelial cell and vascular smooth muscle proliferation and migration.^{32,33} *TNFRSF10A* (rs13278062; $P=0.004$) is part of the tumor necrosis factor-receptor superfamily and acts as a death receptor after binding its cytokine ligand TNFSF10/TRAIL, known to play an important role in the induction of apoptosis,³⁴ vascular smooth muscle proliferation, and regulation of angiogenic responses in the brain after stroke.^{35,36} These potential associations suggest that the extracellular environment as well as angiogenic processes may be involved in cCSC.

We also found a significant association with several SNPs in the *CFH* gene (rs800292, rs1329428, and rs1065489), but the effect sizes were slightly smaller than those

recently reported in a Japanese CSC cohort (e.g., OR= 1.47 for rs1329428 reported in the current cohort versus 1.79 reported by Miki et al¹⁷). This may partially be explained by genetic differences in the frequency of the minor alleles for the tested SNPs at the *CFH* locus that exist between the Japanese and Western European populations (hapmap.org; Table 7), as well as differences in phenotyping. Similar to our observation for *ARMS2*, *CFH* SNPs that confer risk for cCSC were previously found to be protective for AMD, whereas SNPs that were found to be protective for cCSC were previously found to increase risk for AMD.²⁴ The H2 haplotype, previously found to be protective for AMD (OR=0.54),²⁴ was found to increase risk for cCSC (OR=1.33). Conversely, the H3 haplotype that was previously not found to be associated with AMD was found to be protective for cCSC (OR=0.54).²⁴ The observation that *CFH* SNPs have opposite effects in CSC versus AMD was also recently reported by Miki et al,¹⁷ but the underlying mechanisms that could explain these observations are unclear.

The functional implications of the associated *CFH* SNPs in the present cCSC cohort and their possible significance for the pathogenesis of cCSC are unknown, but there seems to be an important role for the *CFH* protein in the RPE and choroid. *CFH* is an inhibitor of the alternative complement pathway (AP), and choroidal cells play a central role in complement AP activity in the eye. C3, an acute phase protein and the central player in complement activation, is most highly expressed in the choroid.^{37,38} The *CFH* protein is the predominant cell surface-associated complement inhibitor in the RPE-choroid complex, where it down-regulates the complement AP activation via inhibition of C3 activity.³⁹

45

The choroid seems to play a pivotal role in the pathogenesis of CSC because choroidal thickening, congestion, and hyperpermeability seem to be the most pronounced and extensive abnormalities in cCSC patients.⁴⁰ In addition, patients with CSC show increased choroidal blood flow during exercise compared with controls.⁴¹ Homozygosity for the *CFH* rs1061170 (Y402H) risk allele was associated with increased choroidal blood flow and ocular perfusion pressure during exercise using Doppler flowmetry,⁴² a process that may be linked to the *CFH*-binding partner adrenomedullin via (steroid-) hormone-dependent nitric oxide bioactivity in vascular endothelial cells.^{39,43,44} Altered complement AP activity, extravasation of complement related proteins, and deposition of terminal complement membrane attack complexes may lead to RPE damage and dysfunction. Damage to the RPE cell layer, further destabilized by changes in the extracellular matrix and under stress from increasing pressure from the thickened choroid, could eventually result in subretinal fluid leakage.^{43,44}

Three distinct phenotypic subgroups within our analyzed cCSC patient population were identified based on multimodal imaging, ranging from typical to less typical phenotypes. The minor allele frequencies of the SNPs (rs10490924, rs800292, rs1329428, and rs1065489) that were significantly associated with typical cCSC (subgroup 1) differed from the minor allele frequencies observed in the other subgroups. Subgroup 2 differs only slightly from the most typical subgroup (subgroup 1), suggesting that considerable genetic overlap exists. In fact, the major distinction in the reported phenotypic criteria are bilateral versus unilateral occurrence. The atypical subgroup (subgroup 3) differs significantly from subgroup 1 for *ARMS2*

(rs10490924; $P=0.002$) and *CFH* (rs800292; $P=0.002$) minor allele frequencies, which suggests that subgroup 3 is genetically different from the typical cCSC group. Based on the phenotypic criteria, subgroup 3 shows overlap with other maculopathies such as AMD, highly myopic macular changes, and polypoidal choroidal vasculopathy. The minor allele frequencies of these 2 SNPs in subgroup 3 were even similar to those normally observed in AMD, with the same direction of effect seen in AMD opposed to typical cCSC. Our results demonstrate that detailed phenotyping in cCSC patients is important to obtain reliable genetic results.

In conclusion, we show that genetic variants in *ARMS2* and *CFH* are associated with cCSC. Detailed phenotyping and classification of cCSC are important to establish such genetic associations. Our findings indicate that complement dysregulation in the RPE/choroid, and potentially dysregulation of genes involved in extracellular matrix and angiogenesis factors, are involved in the pathogenesis of cCSC. Further genetic and phenotypic analysis of cCSC, as well as genotype-phenotype correlation analyses, can provide important clues about the genetic background and pathogenesis of cCSC and may lead to the identification of possible new preventive and therapeutic targets.

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The authors have no proprietary or commercial interest in any materials discussed in this article.

Acronyms and Abbreviations:

AMD=age-related macular degeneration; AP= alternative complement pathway; CFH=complement factor H; cCSC=chronic central serous chorioretinopathy; CSC=central serous chorioretinopathy; EUGENDA=European Genetic Database; FA=fluorescein angiography; ICG=indocyanine green; OR=odds ratio; RPE=retinal pigment epithelium; SD OCT=spectral-domain optical coherence tomography; SNP=single nucleotide polymorphism.

Supplementary Files

Table 7. Minor Allele Frequencies of CFH SNPs in European (CEU) and Japanese (JPT) Population.

SNP	MA (HapMap-CEU)	MA frequency HapMap-CEU	MA frequency HapMap-JPT
<i>CFH</i> rs3753394	T	0.279	0.509
<i>CFH</i> rs800292	A	0.217	0.416
<i>CFH</i> rs2284664	T	0.195	0.406
<i>CFH</i> rs1329428	T	0.424	0.473
<i>CFH</i> rs1065489	T	0.161	0.469

CFH = complement factor H; SNP = single nucleotide polymorphism; MA = minor allele

Table 6. Analysis of 19 Age-related Macular Degeneration Loci in Chronic Central Serous Chorioretinopathy split based on sex

SNP (locus)	Alleles (Major/Minor)	Gender		MAF	MAF controls (M=629/ F=682)	Unadjusted Allelic P-value	Allelic Odds Ratio (95% CI)
		subgroup 1 (M=154/ F=43)					
rs10490924 (ARMS2)	G/T	M	0.15	0.22	0.006	0.63 (0.45-0.88)	
		F	0.21	0.25	0.601	0.83 (0.49-1.42)	
rs12144939 (CFH)	G/T	M	0.27	0.20	0.006	1.51 (1.13-2.01)	
		F	0.18	0.21	0.675	0.86 (0.49-1.54)	
rs429608 (C2-CFB)	G/A	M	0.17	0.14	0.277	1.22 (0.87-1.71)	
		F	0.14	0.13	0.615	1.15 (0.61-2.17)	
rs2230199 (C3)	G/C	M	0.18	0.16	0.355	1.21 (0.81-1.82)	
		F	0.21	0.19	0.630	1.16 (0.63-2.17)	
rs9621532 (TIMP3)	A/C	M	0.05	0.05	0.883	0.93 (0.51-1.69)	
		F	0.08	0.05	0.224	1.60 (0.71-3.60)	
rs4420638 (APOE)	A/G	M	0.15	0.19	0.181	0.78 (0.55-1.10)	
		F	0.18	0.16	0.649	1.13 (0.63-2.01)	
rs3764261 (CETP)	G/T	M	0.31	0.32	0.681	0.94 (0.72-1.23)	
		F	0.27	0.32	0.468	0.81 (0.50-1.33)	
rs943080 (VEGFA)	T/C	M	0.44	0.48	0.337	0.88 (0.68-1.13)	
		F	0.44	0.48	0.501	0.85 (0.55-1.33)	

rs13278062										
(TNFRSF10A)	T/G	M	0.41	0.50	0.005	0.69 [0.54-0.89]				
rs493258		F	0.38	0.46	0.139	0.71 [0.45-1.12]				
(LIPC)	C/T	M	0.49	0.45	0.140	1.22 [0.95-1.56]				
rs10033900		F	0.48	0.47	1.000	1.02 [0.65-1.58]				
(CFI)	C/T	M	0.50	0.48	0.443	1.11 [0.86-1.42]				
rs3812111		F	0.49	0.48	0.911	1.03 [0.66-1.60]				
(COL10A1)	A/T	M	0.35	0.35	1.000	1.00 [0.77-1.30]				
rs13081855		F	0.31	0.37	0.295	0.76 [0.48-1.23]				
(COL8A1-FILIP1L)	G/T	M	0.12	0.09	0.049	1.49 [1.01-2.21]				
rs3130783		F	0.10	0.09	0.844	1.07 [0.50-2.26]				
(IER3-DDR1)	A/G	M	0.22	0.20	0.579	1.10 [0.81-1.49]				
rs8135665		F	0.12	0.18	0.184	0.62 [0.31-1.21]				
(SLC16A8)	C/T	M	0.21	0.24	0.408	0.84 [0.65-1.18]				
rs334353		F	0.18	0.20	0.675	0.85 [0.48-1.51]				
(TGFBRI)	T/G	M	0.27	0.25	0.513	1.10 [0.83-1.46]				
rs8017304		F	0.21	0.25	0.601	0.84 [0.49-1.43]				
(RAD51B)	A/G	M	0.36	0.37	0.742	0.95 [0.73-1.23]				
rs6795735		F	0.35	0.38	0.563	0.85 [0.54-1.35]				
(ADAMTS9)	C/T	M	0.47	0.41	0.052	1.29 [1.00-1.66]				
rs9542236		F	0.44	0.41	0.570	1.14 [0.73-1.77]				
(B3GALT1)	T/C	M	0.43	0.44	0.748	0.96 [0.74-1.23]				
		F	0.48	0.44	0.498	1.18 [0.76-1.83]				

SNP= single nucleotide polymorphism; M= male; F= female; MAF= minor allele frequency; CI= confidence interval. For the analysis displayed in this table two sided P-values < 0.0026 were considered to be significant

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2.2 Genomic copy number variations of the complement component *C4B* gene are associated with chronic central serous chorioretinopathy

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Abstract

Purpose: Chronic central serous chorioretinopathy (cCSC) has recently been associated to variants in the *complement factor H* gene. To further investigate the role of the complement system in cCSC, the genomic copy number variations in the *complement component 4 (C4)* gene were studied.

Methods: *C4A* and *C4B* copy numbers were analyzed in 197 cCSC patients and 303 healthy controls using a Taqman copy number determination assay. Copy numbers of *C4A*, *C4B* and the total *C4* load were compared between cases and controls, using a Fisher Exact test. For this analysis Bonferroni correction was performed for three tests, and P values <0.017 were considered to be significant. A logistic regression model was constructed to calculate the odds ratios (ORs) of each of the *C4B* copy numbers, using two copies as a reference. For this model P values <0.05 were considered to be significant.

Results: *C4B* genomic copy numbers differed significantly between cCSC patients and healthy controls (P=0.0018). Absence of *C4B* significantly conferred risk of cCSC (P=0.039, OR=2.61 [95% confidence interval (CI)=1.05–6.52]), whereas three copies of *C4B* significantly decreased the risk of cCSC (P=0.014, OR=0.45 [95%CI=0.23–0.85]). The *C4A* genomic copy numbers and total *C4* load did not significantly differ between cases and controls.

Conclusions: This study showed that copy numbers of *C4B* are significantly associated with cCSC. Carrying no copies of *C4B* significantly increases the risk of cCSC, whereas carrying three *C4B* copies is protective. These findings reinforce the hypothesis of a possible involvement of the complement system in the pathogenesis of cCSC.

Introduction

Chronic central serous chorioretinopathy (cCSC) is characterized by fluid accumulation under the neuroretina. It has been postulated that this serous fluid derives from the choroid, and that it leaks through a dysfunctional retinal pigment epithelium (RPE) causing a detachment of the neuroretina.¹⁻⁴ Classically, cCSC patients are relatively young (middle-aged) men who are still professionally active.⁶ Besides male sex, also the use of corticosteroids, type A personality and stress have been associated with cCSC.^{4,6,7} Although the exact pathophysiological mechanism of the disease remains unknown, we and others have previously suggested involvement of the complement system, and in particular the *complement factor H (CFH)* gene in cCSC.^{8,9} Interestingly, *CFH* variants that confer increased risk in cCSC have previously been described to be protective in age-related macular degeneration, and vice versa.⁹⁻¹¹

The complement system consists of three major pathways: the classical, the lectin, and the alternative pathway.¹² *CFH* is involved in the alternative pathway and can influence C3b production by blocking one of the two C3-convertases (C3bBb). The classical and lectin pathways also play an important role in C3b production mediated by the other C3-convertase (C4b2a), of which the complement component 4 (C4) protein is a key factor.¹³

Copy number variations of the *C4* gene have been associated with several eye diseases and autoimmune disorders (e.g. Vogt- Koyanagi-Harada and Beçhet's disease).¹⁴⁻¹⁶ In addition, Banlaki et al.¹⁷ have found that the genomic copy number of *C4B* is associated with cortisol release after adrenocorticotrophic hormone (ACTH) stimulation. This is of particular interest considering that stress, which appears to be associated with cCSC,⁴ has a strong influence on the hypothalamic-pituitary-adrenal (HPA) axis and increases ACTH release.¹⁸

Because of its role in the complement system, and its association with the HPA axis, we hypothesized that copy number variation in the *C4* gene may be associated with cCSC. In this study we assessed the copy number variations of the *C4* gene in a cCSC cohort.

Materials and Methods

Subjects

In this study, 197 patients diagnosed with cCSC who visited the outpatient clinic of the department of Ophthalmology at the Radboud University Medical Center, Nijmegen, the Netherlands were included (Table 1). The diagnosis cCSC was based on an extensive ophthalmological examination including funduscopy, spectral-domain optical coherence tomography, fluorescein angiography and indocyanine green angiography. The definition of typical cCSC used in this study was based on the previously published subgroups by de Jong et al.,⁹ and patients in this study were phenotypes by an experienced retina specialist (CJFB) (Figure 1A-F). Additionally, a total of 303 control subjects were recruited from the blood bank of the Radboud University Medical Center, Nijmegen, the Netherlands (n = 154), and the European Genetic Database (EUGENDA, www.eugenda.org; provided in the public domain by the University Hospital of Cologne, Cologne, Germany and Radboud university medical center) (n = 149) (Table 1). For this last group fundus photographs were graded

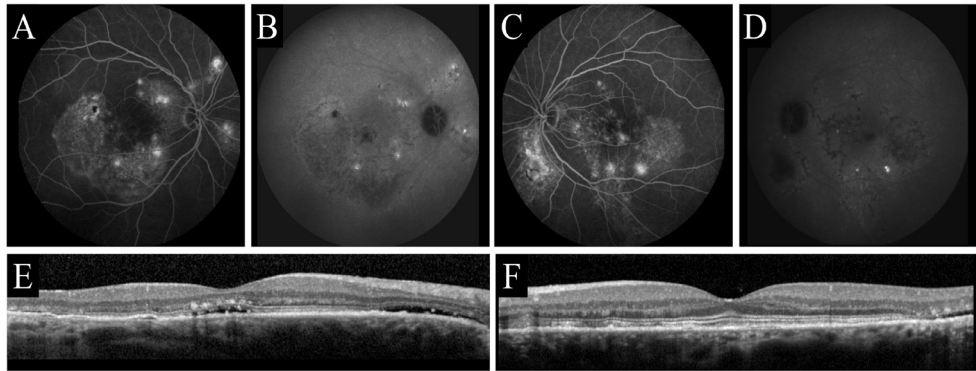


Figure 1: Example of FA of an RE (A), and an LE (C), ICG angiography of an RE (B), and an LE (D), and spectral-domain optical coherence tomography of an RE (E), and an LE (F) imaging of the phenotypic characteristics of the cCSC cohort used for this study. (A-F) The RE and LE of a patient demonstrate diffuse hyperfluorescent areas of leakage on FA and ICG angiography (A-D) and subretinal fluid beneath the fovea (E), illustrative for typical cCSC. FA, fluorescein angiography; ICG, indocyanine green; LE, left eye; RE, right eye.

to rule out any ophthalmological abnormalities at the moment of inclusion. Informed consent for the use of DNA for genetic studies was obtained from all subjects. This study followed the guidelines of the Declaration of Helsinki and was approved by the local ethics committee.

Copy number determination

DNA was isolated from peripheral blood using standard procedures. *C4A* and *C4B* copy numbers were determined by real-time PCR using Taqman® genotyping assays (Applied Biosystems, Thermo Fisher Scientific, Waltham, USA). The FAM-labeled *C4A* (Hs07226349_cn) or *C4B* (Hs07226350_cn) Taqman copy number assay was combined together with the VIC-labeled Ribonuclease P (*RNaseP*) reference assay (Catalog nr. 4403326), and Taqman genotyping mastermix (Catalog nr. 4381656). All samples were tested in duplicate for *C4A* and *C4B* on 384-wells plates using 10 ng of DNA in a total reaction volume of 10 μ l.

Samples with known copy numbers for either *C4A* (0-4) or *C4B* (0-3) were kindly provided by C. Yung Yu.¹⁹ These samples were included as a reference on each plate to facilitate accurate copy number determination, using the method described previously.¹⁹ In each run the amplification efficiencies of the *C4A/C4B* and *RNaseP* probes were calculated using a serial dilution (50-1.56 ng) of a sample with two *C4A* and *C4B* copies. The primer efficiencies of the probes were compared, and deemed

	cCSC patients	Controls	P value
No. of subjects	197	303	NA
Sex (male/female)	154/43	226/77	0.392
Mean age \pm SD (years)	53 (\pm 10)	53 (\pm 11)	0.755
Age range (years)	29-74	29-77	NA

cCSC; chronic central serous chorioretinopathy, NA; not annotated SD; standard deviation

similar if they differed <2%, this was the case in all runs. Therefore, the efficiencies were not incorporated into the calculations of the copy numbers. Polymerase chain reaction was performed with a 7900HT thermocycler (Applied Biosystems, Thermo Fisher Scientific) using the following program: 2 minutes at 50°C, 10 minutes at 95°C, and 40 cycles of 15 seconds at 95°C and 60 seconds at 60°C. Data were analyzed with the Copycaller software (V2.0; Applied Biosystems, Thermo Fisher Scientific). Copy numbers determined by the Copycaller software were corrected by using the trend line based on the reference samples, as described before.¹⁹ If the results were inconsistent between the Copycaller output and the values corrected with the trend line, samples were retested on a new plate in triplicate.

Statistics

The comparison of *C4A*, *C4B* and total *C4* copy number distribution between cCSC patients and controls was performed by means of a Fisher Exact test using SPSS Statistics (V20; IBM Corp., Armonk, USA). Bonferroni correction for multiple testing was performed for three tests and P values < 0.017 were considered to be statistically significant. A logistic regression model was constructed to determine the odds ratios (ORs) for the various copy numbers of *C4B*. According to previously published studies, two genomic copy numbers of *C4B* are considered to be most common in the healthy population.^{16,20} We were able to confirm this in our cohort, and therefore this copy number was set as reference. In this model, P values <0.05 were considered to be significant. Graphs were generated using Graphpad Prism (V5; Graphpad Software, San Diego, CA, USA).

Results

The copy numbers of *C4A* and *C4B* were successfully determined in 197 cCSC cases and 303 controls. No significant difference was observed between cases and controls for the *C4A* genomic copy number (range: 0-6, P=0.649; Figure 2A). The *C4B* distribution was significantly different between cCSC patients and controls (range: 0-4, P=0.0018; Figure 2B). Overall, cases carried lower copy numbers of *C4B* than the control population. The total *C4* genomic copy number was not different in cases compared to controls (P=0.148; Figure 2C). Age and sex were not associated with either *C4A*, *C4B*, or total *C4* genomic copy number (Table 2; data for *C4A* and total *C4* not shown).

To assess the effect size of the different copy numbers of *C4B* on development of cCSC, a logistic regression was performed (Table 2). The logistic regression model based on the distribution of *C4B* between cases and controls was significant (P=0.0035, Table 2). Carrying no copies of *C4B* conferred increased risk of cCSC (P=0.039, OR=2.61, 95% confidence interval [CI]=1.05-6.52). A similar trend was observed for carriers of one copy of *C4B*, but the results were not significant (P=0.080, OR=1.47, 95% CI=0.96-2.26). Carrying three *C4B* copies was associated with a significantly decreased risk of cCSC (P=0.014, OR=0.45, 95% CI=0.24-0.85), whereas no significant association with cCSC was observed in individuals carrying four copies of *C4B* (P=0.81).

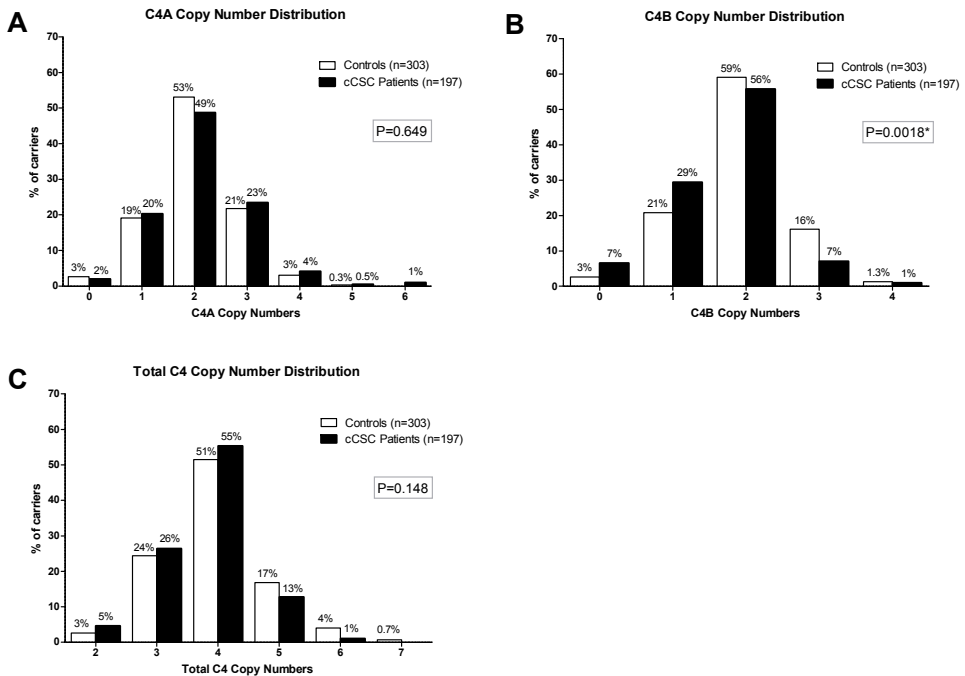


Figure 2: Distribution of the genomic copy numbers of *C4A* (A), *C4B* (B), and total *C4* (C) in patients with cCSC and controls. Displayed P values were generated with a Fisher Exact test. To correct for multiple testing, P values <0.017 were considered to be significant.

Table 2. Logistic regression model for *C4B* load

Overall significance model, $P=0.0035$

<i>C4B</i> copy nr.	cCSC patients (n)	Controls (n)	P value	OR	95% CI
Age	197	303	0.605	NA	NA
Gender	197	303	0.346	NA	NA
0	13	8	0.039	2.613	1.048-6.518
1	58	63	0.080	1.469	0.956-2.259
2	110	179	Ref.	1	NA
3	14	49	0.014	0.445	0.234-0.849
4	2	4	0.808	0.809	0.145-4.503

OR, Odds Ratio; CI, Confidence Interval, Ref, Reference

Discussion

Our study results demonstrate that cCSC patients have a significantly different *C4B* load as compared to healthy controls ($P=0.0018$). Carrying no copies of *C4B* was associated with an increased risk of cCSC (OR=2.61, 95% CI=1.05-6.52), whereas carrying three *C4B* copies was associated with a decreased risk of cCSC (OR=0.45, 95% CI=0.23-0.85). No association with cCSC was observed in individuals carrying four *C4B* copies, which is likely due to the limited sample size of this group (cases, $n=4$; controls, $n=2$). No significant differences were observed between cases and controls for *C4A* and total *C4* load.

The *C4* gene lies within the *RP-C4-CYP21-TNX* (*RCCX*) locus located in the major histocompatibility complex (MHC) region III on chromosome 6 of the human genome.²⁰ The MHC region contains an elevated level of genomic copy number variations that are presumably present to increase immunological diversity.²¹ Duplications and deletions in the region have led to the formation of haplotypes containing variable copies of the *RCCX* locus in the human population (Figure 3B).^{5,21,22} Haplotypes containing two or more duplications of the *RCCX* locus show extensive variability in their gene content, generally with complete duplications of the *C4* gene (Figure 3B).^{22,23} The *C4* gene encodes for the C4 protein, of which two variants have been described (*C4A/C4B*), differing in only four amino acids encoded by exon 26 (Figure 3A).²⁴

Copy number variations of either *C4A* or *C4B* have been associated with several systemic diseases with ocular involvement, such as Vogt-Koyanagi-Harada disease,

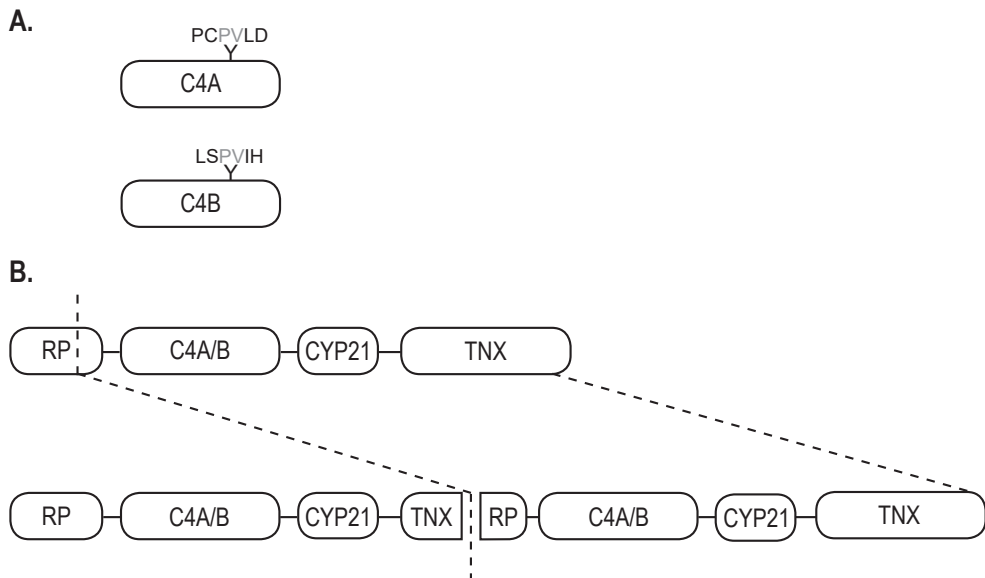


Figure 3: A schematic illustration of the *RCCX* locus. (A) A display of the two variants of the *C4* gene, differing in four amino acids in exon 26 (p. 1101-1106). (B) An example of the structure of the *RCCX* locus with a single copy (upper panel) and two copies (lower panel) of the *C4* gene. In case of duplication, a complete *C4A* or *C4B* and the *CYP21* gene are duplicated, while the *RP* and *TNX* gene are only partially duplicated. The dotted lines indicate the approximate site where duplication has occurred in the past. (Figure based on Banlaki et al. 2013).⁵

Behçet's disease, and systemic lupus erythematosus (SLE).¹⁴⁻¹⁶ Hou et al.¹⁴ have shown that a lower copy number of *C4A* and *C4B* increases the risk of Vogt-Koyanagi-Harada, an autoimmune disorder characterized by bilateral granulomatous panuveitis. The same group¹⁵ has also demonstrated that higher copy numbers of *C4A* confer risk of Behçet's disease, an autoinflammatory disease, which presents with acute anterior uveitis. Several studies^{16,25,26} have shown an association between low copy numbers of *C4* and an increased risk for SLE, an autoimmune disease that is typically mediated by immune complexes. In the past, CSC has been described in SLE patients,^{27,28} but it remains unclear whether this is a primary manifestation of SLE, or whether it is a consequence of corticosteroid treatment for SLE.^{29,30} Several studies have reported a positive linear correlation between serum *C4* and *C4* genomic copy number,³¹⁻³³ suggesting that the lower number of *C4B* copies in cCSC patients leads to lower systemic *C4B* levels. This may indicate that an overall lower activity of the complement system might be present in cCSC patients.

Recently, low copy numbers of *C4B* have been associated with hyperreactivity of the HPA axis.¹⁷ Banlaki et al.¹⁷ have shown that in patients with adrenal incidentaloma and low (<2 copies) *C4B* genomic copy number, baseline ACTH was significantly reduced compared to high (≥ 2 copies) genomic copy number of *C4B*. Moreover, a significantly higher cortisol response is observed after ACTH stimulation in the patients with low *C4B* genomic copy number.¹⁷

These results are of interest in the context of cCSC because of the described clinical associations with stress and the use of corticosteroids that both exert physiological effects at the level of the HPA axis.¹⁸ Various relatively small studies have studied cortisol levels in cCSC patients. Although 24h urine samples showed elevated cortisol levels in cCSC patients in certain studies,^{34,35} these results are not observed in single serum measurements during set times in other studies.^{36,37} These discrepancies could be explained by variable cortisol fluctuations between individuals during the day, and therefore changes in endogenous cortisol levels cannot be ruled out as a hallmark of cCSC. It is possible that patients with cCSC generally have normal cortisol levels, but respond differently to stimulation of the HPA axis. Stress, which also appears to be associated with cCSC,^{2,38} stimulates the HPA axis and could lead to temporarily elevated cortisol levels in patients as compared to healthy individuals. How high levels of cortisol can lead to subretinal fluid accumulation is currently unknown. A study in rats suggested that the disease mechanism could be mediated by binding of corticosteroids to the mineralocorticoid receptor.³⁹ In this study, activation of the mineralocorticoid receptor causes vascular effects similar to those observed in cCSC,³⁹ but the underlying pathways still remain to be elucidated.

The mechanism through which low copy numbers of *C4B* may lead to hyperresponsiveness of the HPA axis is unclear. It has been hypothesized that it is not the *C4B* gene, but rather the neighboring *CYP21A2* gene, that mediates this effect.¹⁷ The *CYP21A2* gene encodes the enzyme 21-hydroxylase, which plays an important role in the steroid metabolism pathway by converting progesterone and 17- α -hydroxyprogesterone to 11-deoxycorticosterone and 11-deoxycortisol, respectively. Because of the genomic structure of the *RCCX* locus, variation in the *CYP21A2* gene is in high linkage disequilibrium with variation in the neighboring *C4* gene (*C4A* or *C4B*).¹⁷ Therefore, further exploration of the precise structure

and specific variations present in the *RCCX* locus may reveal new insights into the pathogenesis of cCSC.

The current study and previous studies identified an association between complement genes and cCSC,^{8,9} suggesting that the complement system may be dysregulated in cCSC. Taken together, these findings may indicate that the immune system, influenced by environmental factors, such as stress, could play a pivotal role in the pathophysiology of cCSC. Further studies are necessary to determine the physiological effects of genetic variation at the *C4* gene and the *RCCX* locus in cCSC.

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The authors alone are responsible for the content and writing of the paper.

No conflicting relationships exist for any author.

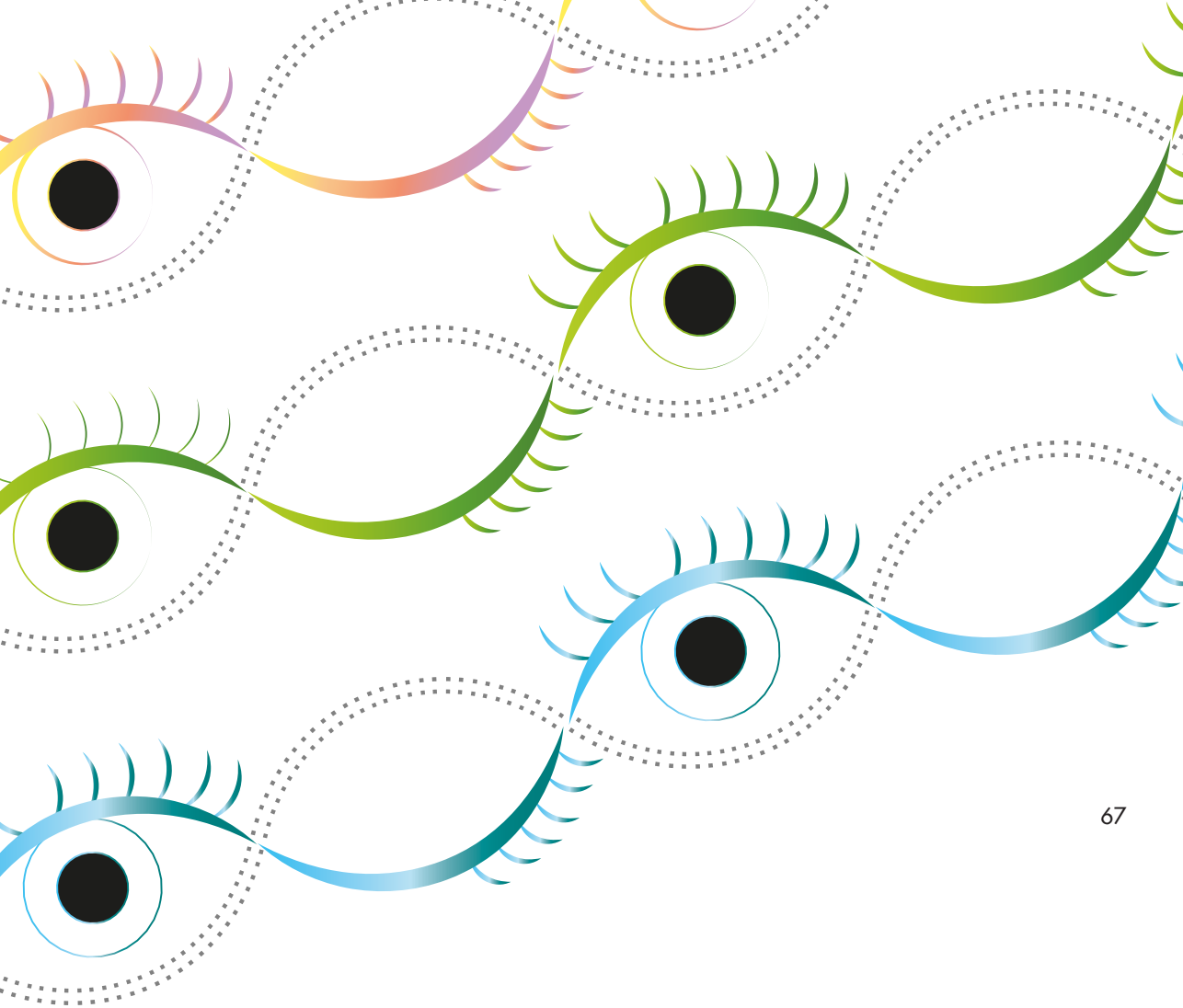
Running head: C4B in chronic central serous chorioretinopathy

Keywords: chronic central serous chorioretinopathy, cCSC, complement component 4, C4, C4A, C4B

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2.3 Association of a haplotype in the *NR3C2* gene, encoding the mineralocorticoid receptor, with chronic central serous chorioretinopathy

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Abstract

Importance: Chronic central serous chorioretinopathy (cCSC) is a chorioretinal disease with unknown disease etiology. The glucocorticoid receptor and the mineralocorticoid receptor, 2 glucocorticoid-binding receptors, might be involved in the pathogenesis of cCSC.

Objective: To assess the association of functional variants and haplotypes in the glucocorticoid receptor (*NR3C1*) and mineralocorticoid receptor genes (*NR3C2*) with cCSC.

Design, setting, and participants: In this case-control genetic association study, 336 patients with cCSC and 1314 unaffected controls, collected at 3 university medical centers from September 1, 2009, to May 2016, underwent KASP genotyping for selected variants in *NR3C1* (rs56149945, rs41423247 and rs6198) and *NR3C2* (rs2070951 and rs5522).

Main outcome measure: Genetic associations of 3 *NR3C1* variants and 2 *NR3C2* variants with cCSC.

Results: Among the 336 patients (274 men and 62 women; mean [SD] age, 52[10] years), after correction for multiple testing, rs2070951 in the *NR3C2* gene was significantly associated with cCSC (odds ratio, 1.29; 95% CI, 1.08-1.53; P=.004). Moreover, the GA haplotype of single nucleotide polymorphisms rs2070951 and rs5522 in *NR3C2* conferred risk for cCSC (odds ratio, 1.39; 95% CI, 1.15-1.68; P=.004), whereas the CA haplotype decreased risk for cCSC (odds ratio, 0.72; 95% CI, 0.60-0.87; P<.001). Three known variants in *NR3C1* that alter the activity of the glucocorticoid receptor (rs56149945, rs41423247 and rs6198) were not associated with cCSC.

Conclusions and Relevance: In this study, the variant rs2070951 and the GA haplotype in *NR3C2* were associated with an increased risk for cCSC. Results of this genetic study support a possible role for the mineralocorticoid receptor in the pathogenesis of cCSC. Since these haplotypes have previously been associated with perceived stress, this study provides a clue to bridging clinical risk factors for cCSC to underlying genetic associations.

Introduction

In central serous chorioretinopathy (CSC), it has been suggested that dysfunction of the retinal pigment epithelium (RPE) due to congestion, thickening, and hyperpermeability of the underlying choroid leads to subretinal fluid accumulation with an associated detachment of the neuroretina.¹⁻⁵ The exact etiology of the disease is currently unknown, but clinical associations point towards an involvement of steroid signaling. Endogenous hypercortisolism (Cushing's syndrome), exogenous glucocorticoid exposure, and possibly stress and type A personality are associated with CSC.⁶⁻¹¹ It has been hypothesized that the glucocorticoid receptor (GR) and the mineralocorticoid receptor (MR), 2 glucocorticoid-binding receptors, may also be involved in the pathogenesis of CSC.²

The involvement of the MR in the pathogenesis of CSC has been suggested based on the results of studies in rats, in which choroidal findings similar to those seen in CSC occurred after intravitreal injection of either corticosterone or aldosterone.¹² Involvement of the MR was further supported by ophthalmological findings in patients with primary hyperaldosteronism (Conn syndrome).¹³ Moreover, studies evaluating the administration of MR antagonists for patients with cCSC have shown possible beneficial effects.^{12,14-17} However, clinical results were variable and not permanent, and no prospective randomized placebo-controlled clinical trials have been published to this date to study the role of MR antagonists in the treatment of CSC.^{2,16}

The GR is the most widely expressed cortisol receptor in the body; it regulates metabolism and the cardiovascular system, and it plays a role in immune suppression and stress response.¹⁸ As stress and both exogenous and endogenous hypercortisolism may be involved in the etiology of CSC,^{4,7,10,11} the GR may also be an interesting player in the pathogenesis of CSC.

There are several genetic variants in the genes encoding the MR and GR that are known to alter the MR and GR protein activity.¹⁹⁻²⁶ The MR is encoded by the *NR3C2* gene (OMIM 600983), consisting of 10 exons, with 2 alternative 5'-UTR exons 1 α and 1 β allowing tissue-specific promoter activation.²⁷ The GR is encoded by the *NR3C1* gene (OMIM 138040), consisting of 10 exons of which 1 to 9 α are translated into the functional GR α receptor.²² In this study, we assessed whether genetic variants in *NR3C2* (rs2070951 and rs5522) and *NR3C1* (rs56149945, rs41423247, and rs6198) are associated with cCSC.

Key Points

Question: Are functional variants and haplotypes in the glucocorticoid receptor (*NR3C1*) and mineralocorticoid receptor (*NR3C2*) genes associated with chronic central serous chorioretinopathy (cCSC)?

Findings: In this case-control, genetic association study, rs2070951 in the *NR3C2* gene was significantly associated with cCSC. The GA haplotype of single-nucleotide polymorphisms rs2070951 and rs5522 in *NR3C2* conferred risk for cCSC, whereas the CA haplotype decreased the risk for cCSC.

Meaning: Results of this genetic study support a possible role for the mineralocorticoid receptor in the pathogenesis of cCSC.

Materials and methods

We included 336 patients with cCSC in this study. Phenotyping was performed from September 1, 2009, to May 1, 2016, by an experienced retina specialist (C.J.F.B.) and was based on results of a complete ophthalmological examination, including fundoscopy, optical coherence tomography, fluorescein angiography, and indocyanine green angiography. The patients showed the most typical clinical cCSC characteristics (serous subretinal fluid affecting the fovea on optical coherence tomography, a disease period of >3 months, ≥ 1 area of “hot spot” leakage [the point where fluid is presumed to leak into the serous fluid pocket] or diffuse leakage in combination with irregular retinal pigment epithelial window defects on fluorescein angiography, and a corresponding hyperfluorescence detected on indocyanine green angiography), described as phenotypic subgroup 1 in a previous article on genetic associations in cCSC.²⁸ Patients with high myopia, evidence of choroidal neovascularization, polypoidal choroidal vasculopathy, and other atypical findings were excluded. For this study, neither previous nor current steroid use was considered an exclusion criterion. The patient cohort consisted of 234 patients from the Radboud university medical center (Nijmegen, the Netherlands), 72 patients from the Leiden University Medical Center (Leiden, the Netherlands), and 30 patients from the University Hospital of Cologne (Cologne, Germany). A total of 1314 unaffected individuals (ie, controls) enrolled in the European Genetic Database (www.eugenda.org) were included in the study; they has no signs of either cCSC or age-related macular degeneration when evaluated by multimodal imaging. The study adhered to the tenets of the Declaration of Helsinki,²⁹ and was approved by the institutional review boards and the ethics committees of the Radboud university medical center, the Leiden University Medical Center, and the University Hospital of Cologne. Written informed consent was received from all participants.

71

Genotyping of selected variants was performed using KASP assays (LGC Genomics) according to manufacturer’s instructions. Specific primers with 5-carboxyfluorescein and 5'-hexachloro-fluorescein labels were designed per variant (*NR3C1*: *rs6198*, *rs5614994*, and *rs41423247*; *NR3C2*: *rs5522* and *rs2070951*) (eTable 1 in the Supplement) and the polymerase chain reaction (PCR) conditions per primer pair were provided by LGC Genomics. Data were read with the 7900HT Fast Real-Time PCR system (Applied Biosystems by Life Technologies) and analyzed with SDS version 2.4 (Applied Biosystems).

Table 1. Demographic characteristics of the cCSC patients and controls

	n	Mean age \pm stdev (years)	Males (%)
cCSC cases			
Nijmegen	234	52 \pm 9	188 (80%)
Cologne	30	50 \pm 9	24 (80%)
Leiden	72	52 \pm 10	62 (86%)
Total	336	52 \pm 10	274 (81,5%)
Controls	1314	70 \pm 7	549 (42%)

cCSC, chronic central serous chorioretinopathy; stdev, standard deviation

Statistical Analysis

In IBM SPSS Statistics version 22 (SPSS Inc.), the 2-sided Pearson's χ^2 test was used to compare both the genotype and allele frequencies between cases and controls. Bonferroni correction for multiple testing was performed for 5 variants and $P < .01$ was considered statistically significant. Logistic regression was performed for the associated rs2070951 variant with Firth bias-corrected likelihood ratio test implemented in EFACTS version 3.2.6 (Efficient and Parallelizable Association Container Toolbox, <http://genome.sph.umich.edu/wiki/EFACTS>), correcting for sex.³⁰

Using a haplotype analysis, we assessed the combined effect of the 2 variants in NR3C2. Haplotype analysis was performed using R version 3.0.2 (R Core Team; <https://www.R-project.org>), using the haplo.stats package (version 1.6.8). The 2 most frequent haplotypes were separately used as a reference in the haplo.cc command, to determine odds ratios (ORs) for both haplotypes. A logistic regression analysis (haplo.glm) including sex and haplotypes was performed using the most common haplotype as a reference. Only haplotypes with a frequency higher than 5% are shown. $P < .05$ was considered statistically significant.

Power analysis was performed with CaTS version 0.0.2 (University of Michigan), using a multiplicative model in a joint analysis.³¹ The power per variant was calculated based on the minor allele frequency in controls, a disease prevalence of 0.0001, and a variable genotype relative risk (1-2.6), and the graph was created with Graphpad Prism version 5.03 (Graphpad Software).

Results

Among the 336 patients, 274 were men and 62 were women; the mean (SD) age was 52 (10) years. The demographic characteristics of the patients and controls enrolled in this study are summarized in Table 1. All described variants were in Hardy-Weinberg equilibrium, both for controls and patients with cCSC. No statistically significant associations between cCSC and variants in the NR3C1 gene (rs56149945,

rs41423247 and rs6198) were found (Table 2). After correction for multiple testing, a significant association between cCSC and the rs2070951 variant in the NR3C2 gene was observed (OR, 1.29; 95% CI, 1.08-1.53; $P = .004$). No association between the variant rs5522 in NR3C2 and cCSC was found (Table 2).

Haplotype analysis of the NR3C2 single-nucleotide polymorphisms rs2070951 and rs5522 showed a significant decreased risk for cCSC for the CA haplotype (OR, 0.72; 95% CI, 0.60-0.87, $P < .001$) and an increased risk for the GA haplotype (OR, 1.39; 95% CI 1.15-1.68, $P = .004$) (Table 3). To account for potential confounding effects of sex between cases and controls, we corrected for this factor in a logistic regression model. When including this variable in the model, the association of rs2070951 was independent of sex (OR, 1.28; 95% CI, 1.16-1.41; $P = .009$). Similarly, when correcting for sex in the haplotype analysis, setting the most common haplotype (GA) as reference, we found that the association of the

A multiplicative model was used to calculate the power of the study, because this model produces a genetic relative risk score that is an estimation of the OR of an allelic model.³² For each variant, the power of detecting an association was calculated

Table 2. Association analysis of variants in *NR3C1* and *NR3C2* in cCSC patients

Variant	Gene	Location	Major/ Minor allele	Minor Allele Frequency		P value	Geno- type	Allelic	OR (95% CI)
				Controls	Patients				
rs56149945	<i>NR3C1</i>	Exon 2	A/G	0.0415	0.0357	.37	.50	0.86 (0.55-1.34)	
rs41423247	<i>NR3C1</i>	Intron 2	C/G	0.360	0.390	.16	.15	1.14 (0.95-1.35)	
rs6198	<i>NR3C1</i>	Exon 9 UTR	A/G	0.169	0.158	.334	.48	0.92 (0.73-1.16)	
rs2070951	<i>NR3C2</i>	c.-2	C/G	0.464	0.527	.008	.0040	1.29 (1.08-1.53)	
rs5522	<i>NR3C2</i>	Exon 2	A/G	0.129	0.137	.84	.57	1.08 (0.83-1.38)	

cCSC, chronic central serous chorioretinopathy; CI, confidence interval; MAF, Minor allele frequency; OR, odds ratio; UTR, untranslated region. Bonferroni correction for multiple testing was performed for 5 variants and p-values < 0.01 were deemed significant

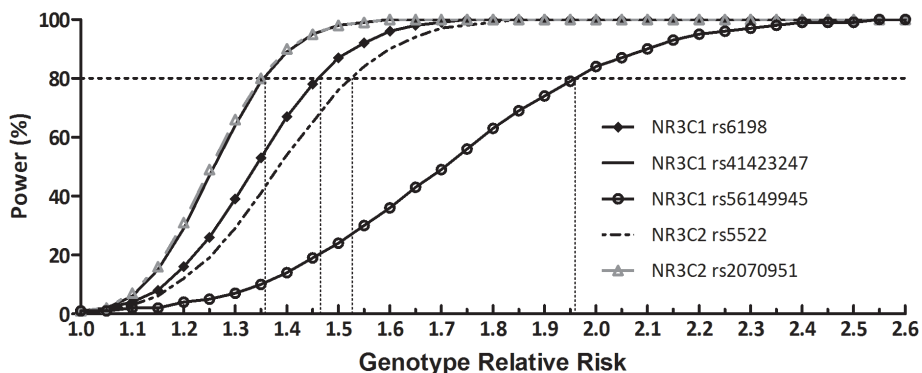


Figure 1: Power calculation for each variant in a cohort of 336 patients with chronic central serous chorioretinopathy patients and 1314 unaffected controls. For each variant, the power to detect a certain genotype relative risk was assessed using a multiplicative model, with minor allele frequency in controls and disease frequency of 0.0001 as input. The 80% power detection limits per variant were 1.35 (rs2070951), 1.36 (rs41423247), 1.46 (rs6198), 1.52 (rs5522), and 1.95 (rs56149945).

for the current cohort size. For all variants, genotype relative risks less than 2 could be detected with 80% power using this cohort. The detection limits of the genetic relative risk at 80% power were 1.35, 1.36, 1.46, 1.52, and 1.95 for rs2070951, rs41423247, rs6198, rs5522, and rs56149945, respectively (Figure 1).

Discussion

In this study, we analyzed a possible association with cCSC for 3 known functional variants in *NR3C1* and 2 known functional variants in *NR3C2*. The rs2070951 variant in *NR3C2* was significantly associated with cCSC, whereas the rs5522 variant was not associated with cCSC. The *NR3C2* CA and GA haplotypes were both significantly associated with cCSC, with a protective and a risk-conferring effect, respectively. Odds ratios of the associated *NR3C2* variant and the haplotypes were similar to previously described associations for the *CFH* (OMIM 134370) and *ARMS2* (OMIM 611313) genes, and lower compared to the previously described associations in the *C4A* (OMIM 120810), *C4B* (OMIM 120820), and *CDH5* (OMIM 601120) genes.^{28,33-35} The 3 variants in the *NR3C1* gene (encoding the GR) were not associated with cCSC, which may suggest that MR functionality is more relevant than GR functionality in the etiology of cCSC. However, a larger cohort size is required to exclude the involvement of the 4 variants that were not associated with cCSC in this study.

An abnormal response to the administration of corticosteroids in a subset of patients with cCSC is the strongest risk factor for the disease with described ORs of up to 37.⁹ However, the precise mechanism of action of steroids in the pathogenesis of cCSC is unknown. One study showed that both mineralocorticoids and glucocorticoids can activate the MR on choroidal endothelial cells in a rat model.¹² In this animal model, MR activation resulted in vessel dilation via up-regulation of the endothelial vasodilatory calcium-dependent potassium channel KCa2.3,^{2,12} producing choroidal thickening that is also commonly observed in patients with cCSC.² The MR is also present on retinal pigment epithelial cells, and clearance of subretinal fluid through the retinal pigment epithelium towards the choriocapillaris may be influenced by

Table 3. Haplotype analysis of *NR3C2* in cCSC patients

Haplo- type	rs2070951	rs5522	P value	Frequency		Direction of effect	OR (95% CI)	Direction of effect	OR (95% CI)
				Controls	Patients				
H1	C	A	<.001	0.408	0.336	Base	NA	Protective	0.72 (0.60- 0.87)
H2	C	G	.54	0.127	0.137	NS	1.32 (1.00- 1.75)	NS	0.95 (0.73- 1.24)
H3	G	A	.004	0.463	0.527	Risk	1.39 (1.15- 1.68)	Base	NA

cCSC, chronic central serous chorioretinopathy; CI, confidence interval; Freq, frequency; NA, not annotated; NS, not significant; OR, odds ratio

differences in MR haplotypes.³⁶ In addition, on Müller glial cells, the MR regulates water homeostasis in the eye; dysregulation of this mechanism may contribute to the intraretinal edema observed in a subset of cCSC patients.^{2,37} However, direct GR overactivation without MR involvement seems to be sufficient to induce cCSC in some patients because synthetic glucocorticoids with strong selectivity for GR over MR have also been described as a risk factor for the disease.^{7,9}

Both variants in *NR3C2* that were tested in this study influence the transactivational capacity of the MR after exposure to both cortisol and dexamethasone,²¹ and they have been shown to affect salivary cortisol levels, especially during the morning cortisol awakening peak.^{9,21,38} The rs2070951 G variant, which is associated with cCSC in this study, leads to lower expression of MR and reduced transactivation. One study showed that male carriers of the rs2070951 GG genotype in *NR3C2* had a higher systolic blood pressure.²⁰ This finding is particularly relevant in the context of cCSC because hypertension is a known risk factor for the disease.^{39,40} The effect of this genetic variant on systolic blood pressure was observed only in male patients, which is interesting because cCSC is more common in men than in women.⁴¹ In our dataset, the association for rs2070951 was also observed only in male cCSC patients when the data was stratified for sex (males: OR, 1.21; 95% CI, 1.04-1.58; P=.02 vs females: OR, 1.21; 95% CI, 0.84-1.75; P=.31, eTable 2 in the Supplement). However, this is likely owing to the limited number of female patients with cCSC included in the analysis, and based on the current data we therefore cannot definitively conclude that sex differences exist in this genetic association.

Haplotypes in *NR3C2* have previously been associated with differences in perceived chronic stress,²⁶ another postulated risk factor for cCSC.^{11,42-44} We found that the haplotype of the single-nucleotide polymorphism rs2070951 and rs5522, GA, which has been previously associated with increased susceptibility to stress,²⁶ conferred risk for developing cCSC in our cohort. The haplotype that was associated with an optimistic attitude (and a tendency to recover from or adjust easily to misfortune or change), CA,⁴⁵ was protective against the development of cCSC. This finding could indicate that, for patients with cCSC who have the GA haplotype, both the MR-mediated pathways and chronic stress are of significant importance, whereas other not-yet-identified factors could play a bigger role in patients with cCSC who have the CA haplotype. In addition, there is a likelihood that patients with the different haplotypes carry additional unknown genetic variants that might also contribute to an increased or decreased cCSC risk.

Clinical studies that tested the potential of MR antagonists in the treatment of cCSC have yielded mixed results.¹⁴⁻¹⁷ Our findings may partly explain this variable response to MR antagonists because carriers of different MR haplotypes may respond differently to MR antagonists. The results of our study may lead to the stratification of patients with cCSC into subgroups, based on MR haplotype. Treatment of these stratified patient subgroups with MR antagonists could result in group-specific effects. For patients with the CA haplotype, other (thus far unknown) factors could contribute to the development of CSC to a greater extent. The results of this study may therefore indicate that a more personalized treatment approach in cCSC may be useful. Further studies on the response to treatment of patients with who have different MR genotypes are needed to test this hypothesis.

Limitations

Our study contains some limitations. The sample size of our cohort is insufficient to detect genetic variants with low ORs. A larger cohort is needed to completely rule out the association of the 4 remaining variants with cCSC.

Conclusions

In this study rs2070951 in the *NR3C2* gene, encoding the MR receptor, is significantly associated with cCSC. Additionally, haplotypes of *NR3C2* that have previously been associated with perceived stress also associate with cCSC in this study, which may be a first clue bridging clinical risk factors for cCSC to underlying genetic associations. Functional effects of this variant and the associated haplotype in the MR gene may contribute to the disease mechanisms of cCSC.

Footnotes and Financial Disclosures

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Author Contributions:

Dr van Dijk and Ms Schellevis had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Dr van Dijk and Ms Schellevis contributed equally to the article.

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Study supervision: Hoyng, den Hollander, Boon, de Jong.

Conflict of Interest Disclosures

All authors have completed and submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest and none were reported.

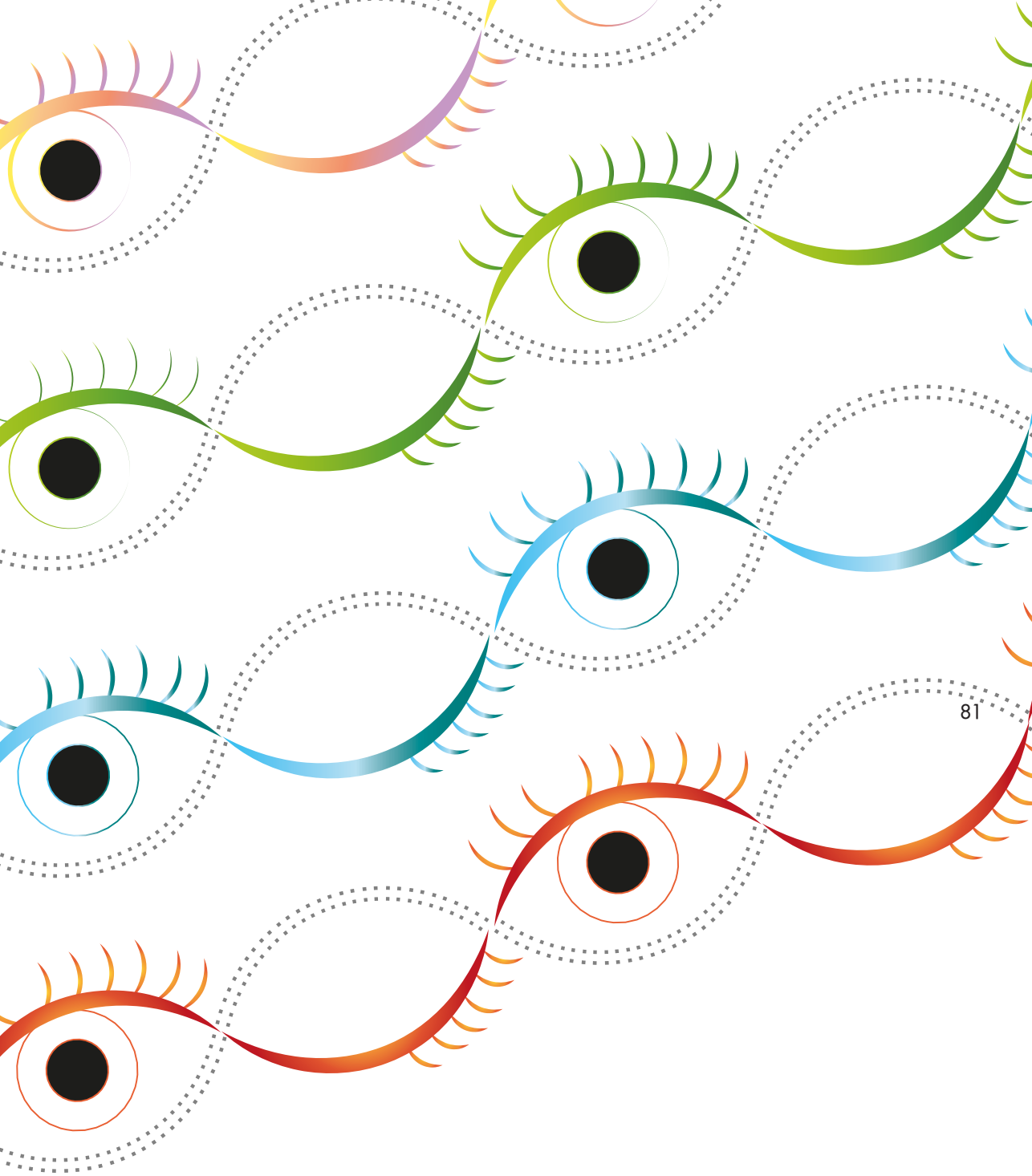
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2.4 Genetic risk factors in acute central serous chorioretinopathy

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Abstract

Purpose: To investigate genetic associations in Caucasian patients with acute central serous chorioretinopathy (aCSC), and to assess genetic differences between aCSC and chronic CSC (cCSC).

Methods: A total of 135 aCSC patients, 272 cCSC patients, and 1385 control individuals were included. Eight single nucleotide polymorphisms (SNPs) were genotyped for *ARMS2* (rs10490924), *CFH* (rs800292, rs1061170, rs1065489, rs1329428, rs2284664, rs3753394), and *NR3C2* (rs2070951). Also, *C4B* gene copy numbers were analyzed.

Results: Three SNPs in the *CFH* gene were significantly associated with aCSC: rs800292 ($P = 0.003$, OR = 1.53 [95% CI = 1.15-2.03]), rs1061170 ($P = 0.002$, OR = 0.64 [95% CI = 0.48-0.86]), and rs1329428 ($P = 5.87 \times 10^{-6}$, OR = 1.83 [95% CI = 1.40-2.38]). A significant difference was found in the distribution of *C4B* gene copy numbers in aCSC patients compared to controls ($P = 0.0042$). No differences could be found among the selected variants between aCSC and cCSC patients.

Conclusions: Three variants in the *CFH* gene and copy number variations in *C4B* were found to be significantly associated with the risk of aCSC development. Despite the differences in clinical presentation, acute and chronic CSC may share a similar genetic predisposition based on our present analysis. Other genetic and/or non-genetic risk factors may be more influential in the differentiation toward an acute or a chronic phenotype of CSC.

Introduction

Acute central serous chorioretinopathy (aCSC) is a sudden-onset and relatively common macular disease.¹ It is characterized by a neuroretinal detachment with serous subretinal fluid (SRF) accumulation as seen on optical coherence tomography (OCT).² Patients with aCSC characteristically show a single focal “hot spot” of leakage on fluorescein angiography (FA).³ This leakage occurs because of a small defect in a focally detached retinal pigment epithelium (RPE), which normally constitutes the outer blood-retina barrier.^{3,4} Acute CSC has been described to be a self-limiting condition and visual acuity recovers completely in most cases.⁵

In contrast to aCSC, the phenotype of chronic CSC (cCSC) is characterized by prolonged and usually persistent SRF accumulation, larger and/or multiple RPE detachments, often more diffuse RPE leakage, and more extensive multifocal atrophic RPE changes.¹ A timely diagnosis and treatment is required in order to accelerate SRF resolution, and to prevent irreversible photoreceptor damage, vision loss and decreased vision-related quality of life.⁶ It is hypothesized that a congested and hyperpermeable choroid lies at the pathophysiological basis of CSC, as part of the pachychoroid spectrum.^{1,7} Dysfunction of the RPE, secondary to these choroidal abnormalities, would then result in SRF leakage and neuroretinal detachment, but the exact etiology of the disease is still unknown.^{7,8} There is ongoing debate about whether aCSC and cCSC form two distinct entities, or whether they belong to a continuum of the same disease.^{9,10}

Recently, single nucleotide polymorphisms (SNPs) in the *ARMS2* gene and the *CFH* gene (involved in the complement system) were found to be significantly associated with cCSC.¹¹⁻¹³ An association of these SNPs was previously identified in age-related macular degeneration (AMD), pointing to a genetic and pathophysiologic overlap between CSC and AMD. An important role for the choroid has been postulated in both diseases, which both manifest at the choriocapillaris-Bruch's membrane-RPE-neuroretina interface. Interestingly, some risk-conferring alleles in *ARMS2* and *CFH* in AMD were found to be protective in cCSC and vice versa.¹¹ We also identified an association of a SNP in the *NR3C2* gene that encodes the mineralocorticoid receptor in cCSC.¹⁴ Furthermore, genomic copy number variations in the *complement component 4 (C4B)* gene were shown to be associated with cCSC.¹⁵ To the best of our knowledge, no genetic studies have been performed to date in patients with an acute phenotype of CSC characterized by only a single focal leak on FA and without any other signs of chronicity. Additionally, clinically distinct aCSC and cCSC phenotypes have not been compared genetically thus far.

In the present study, we therefore assessed whether SNPs in *ARMS2* (rs10490924), *CFH* (rs800292, rs1061170, rs1065489, rs1329428, rs2284664, rs3753394), and *NR3C2* (rs2070951), and the copy numbers of *C4B* gene are associated with aCSC in a Caucasian patient cohort. Furthermore, these genetic variants were compared between Caucasian aCSC and cCSC patients, to assess whether there are significant differences in these genetic risk factors that could indicate that these disease subtypes are (patho)genetically distinct.

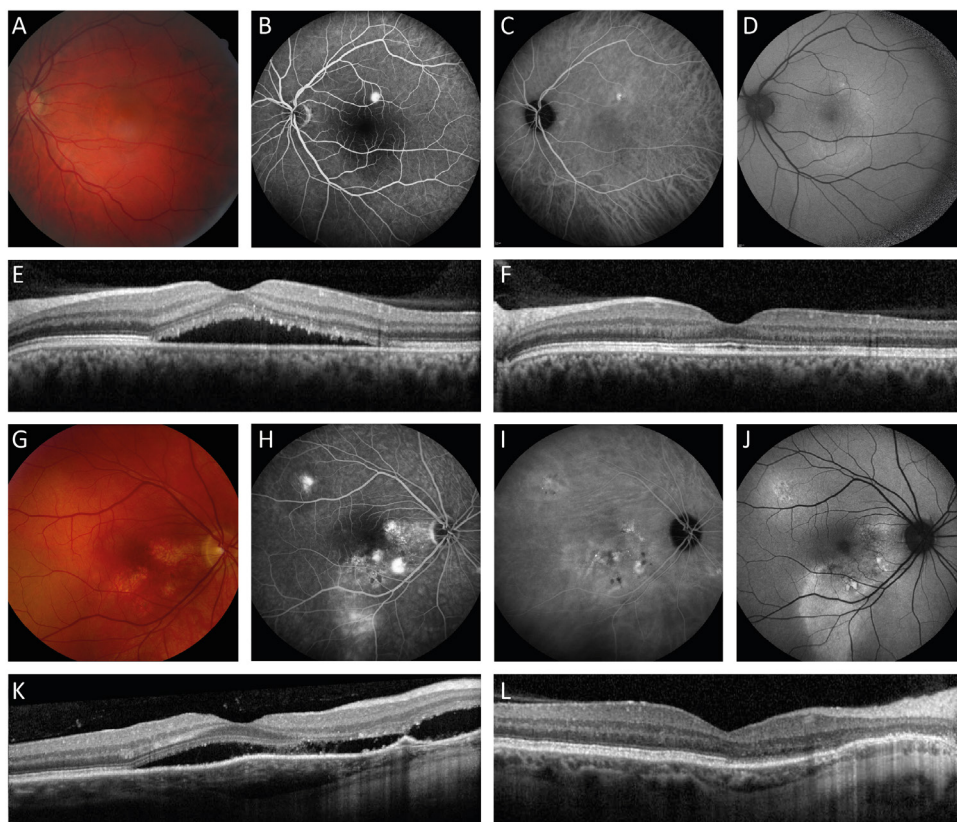


Figure 1: Clinical features visible on multimodal imaging of the left eye of a 41-year-old male patient (A-F) with acute central serous chorioretinopathy (aCSC) and the right eye of a 40 year-old male patient (G-L) with chronic CSC (cCSC). (B) Fluorescein angiography (FA) revealed a single “hot spot” of leakage and no atrophic retinal pigment epithelium (RPE) changes in the aCSC patient. (C) On mid-phase indocyanine green angiography (ICGA) a small hyperfluorescent lesion was observed at the location of the “hot spot” on FA. (D) Fundus autofluorescence (FAF) imaging showed granular hyper-autofluorescent changes at the site of the serous neuroretinal detachment. (E and F) Optical coherence tomography (OCT) scan at first presentation revealed a subretinal serous fluid (SRF) accumulation (E), which resolved after four weeks (F). (H) FA imaging in the cCSC patient revealed a large area of atrophic RPE changes and multiple leakage spots. (I) ICGA imaging in this patient revealed diffuse choroidal hyperpermeability which was slightly larger than the area of leakage visible on FA, and FAF imaging showed a mixture of intense areas of hyper-autofluorescence together with granular hypo-autofluorescent changes. At diagnosis, foveal SRF and a small RPE detachment were observed on the OCT scan of the cCSC patient (K), which both resolved within three weeks after treatment with half-dose photodynamic therapy (L).

Methods

A total of 135 Caucasian subjects with aCSC was included in the study. Subjects were selected from a large cohort of CSC patients from three referral centers: 47 patients from the Department of Ophthalmology at Leiden University Medical Center (Leiden, the Netherlands), 72 patients from the Rotterdam Eye Hospital (Rotterdam, the Netherlands), and 16 patients from University Hospital of Cologne (Cologne, Germany).

Phenotyping of aCSC patients was performed by two experienced retina specialists (SY, CJFB), who had to agree on the aCSC phenotype being typical, which was based on findings on fundoscopy, OCT, FA, and indocyanine green angiography (ICGA) when available. For purposes of comparison to chronic CSC strict criteria for the diagnosis of aCSC were used. Also, only patients who met the definition of aCSC were included when there was at least one follow-up visit and complete resolution of SRF during the first CSC episode (Figure 1). For this study, aCSC was identified on multimodal imaging as a combination of: 1. Documented serous SRF accumulation on OCT; 2. A single focal leakage point ("hot spot") on FA; 3. Atrophic RPE alterations (including RPE detachments) limited to less than one optic disc diameter in size in the affected eye. Also, the contralateral eyes were not allowed to show any signs of chronicity, such as presence of atrophic RPE changes or chronic SRF leakage. Patients with other possible causes of SRF accumulation such as choroidal neovascularization, or polypoidal choroidal vasculopathy were excluded. In this study, previous steroid use was not an exclusion criterion.

The control group consisted of Caucasian individuals enrolled in the European Genetic Database (EUGENDA; www.eugenda.org), in whom no signs of maculopathy were found when evaluated by multimodal imaging, and 176 subjects from the blood bank of the Radboud University Medical Center (Nijmegen, the Netherlands). The control group for the analysis of *ARMS2* and *CFH* included 826 controls, whereas the analysis of *NR3C2* and *C4B* included 1385 and 250 controls, respectively. Additionally, to assess the genetic difference between aCSC and cCSC we included a cohort of 272 Caucasian patients with typical cCSC (Figure 1), as described in a previous genetic analysis on cCSC by our group.¹¹ Both controls and a subgroup of the cCSC patients were genotyped in previous studies.^{11,14,15} Approval for this study was obtained at the local institutional review boards in all participating centers and the study was in accordance with the tenets of the Declaration of Helsinki. Written informed consent was obtained from all subjects prior to blood collection for genetic analysis.

SNP and copy number genotyping

DNA was isolated from peripheral blood by using standard procedures. The choice of the most relevant genetic variants to be analyzed was based on findings in earlier studies.¹²⁻¹⁵ Genotyping of the selected SNPs was performed using KASP assays (LGC Genomics; Berlin, Germany) as described previously according to the manufacturer's instructions. Data were read out with the 7900HT Fast Real-Time PCR system (Applied Biosystems by Life Technologies, Austin, TX, USA) and were analyzed with SDS (version 2.4, Applied Biosystems). *C4B* copy numbers were measured as previously described using a TaqMan genotyping assay (Hs07226350_cn, Applied Biosystems, Thermo Fisher Scientific, Waltham, MA, USA) with RNaseP as a reference assay.¹⁵

Table 1. Demographic characteristics of the study population

	aCSC patients	cCSC patients	Controls <i>ARMS2</i> & <i>CFH</i>	Controls <i>C4B</i>	Controls <i>NR3C2</i>
Number of subjects	135	272	826	250	1385
Number of males	92 (68%)	216 (79%)	424 (51%)	198 (79%)	635 (46%)
Mean age \pm SD (years)	47 \pm 10	51 \pm 10	64 \pm 12	51 \pm 10	51 \pm 10

aCSC = acute central serous chorioretinopathy; cCSC = chronic central serous chorioretinopathy; SD = standard deviation

Statistical analysis

The allele frequency of the SNPs was compared between aCSC and unaffected controls or cCSC patients using a 2-sided Pearson's chi-square test (IBM SPSS Statistics, version 22, SPSS Inc., Chicago, IL, USA). The *C4B* copy numbers distribution was compared with a 2-sided Fisher's exact test and a logistic model correcting for gender was performed setting two copies of *C4B* as a reference, as previously described.¹⁵ Bonferroni correction for multiple testing was performed for nine variants and *P*-values <0.0056 were considered statistically significant. The combined effect of the selected six variants in *CFH* was assessed using a haplotype analysis correcting for gender. Haplotype analysis was performed using R (v3.0.2) using the haplo.stats package (v1.6.8). The two most frequent haplotypes were separately used as a reference in the haplo.glm command to determine odds ratios (ORs) for the haplotypes with a frequency >5% and the aggregate of the haplotypes with a frequency <5%.

Results

Of the 135 aCSC patients included, 92 patients (68%) were men, with a mean age of 47 \pm 10 years (Table 1). Fifty-six aCSC patients (41%) underwent an ICGA imaging and none had a sign of choroidal neovascularisation. Recent steroid use (< 3 months prior to diagnosis) was reported in 29 aCSC patient (21%). The demographic characteristics of aCSC patients, cCSC patients and controls are summarized in Table 1.

Association of SNPs in *ARMS2*, *NR3C2*, and *CFH* genes with aCSC

No association could be found with the rs10490924 variant in *ARMS2* in aCSCs (Table 2). An initial significant association in the rs2070951 variant in *NR3C2* was lost after correction for multiple testing (Table 2). Among the six tested variants in *CFH* gene, five variants showed an association with aCSC. Among these, two variants, rs1065489 (*P* = 0.019, odds ratio (OR) = 0.63 [95% confidence interval (CI) = 0.43-0.93]) and rs2284664 (*P* = 0.013, OR = 1.44 [95% CI = 1.08-1.93]) showed an association, which was lost after correction for multiple testing (Table 2). Three variants were significantly associated with aCSC after correction for multiple testing: rs800292 (*P* = 0.003, OR = 1.53 [95% CI = 1.15-2.03]), rs1061170 (*P* = 0.002, OR = 0.64 [95% CI = 0.48-0.86]), and rs1329428 (*P* = 5.87 \times 10⁻⁶, OR = 1.83 [95% CI = 1.40-2.38]).

Association of *CFH* haplotypes with aCSC

Haplotype analysis corrected for gender identified five haplotypes in the *CFH* gene

with a frequency above 5% and an aggregate of the haplotypes with a frequency lower than 5%. When using the most common haplotype (H1) as a reference, an association with aCSC was found for the risk-conferring H2 ($P = 0.003$, OR = 1.75 [95% CI = 1.21-2.53]), H4 ($P = 0.0180$, OR = 1.69 [95% CI = 1.09-2.6]) and H5 ($P = 0.001$, OR = 2.3 [95% CI = 1.39-3.83]), of which H2 and H5 were significant after correction for multiple testing (Table 3). Using the H2 haplotype as a reference, a protective effect for the H1 ($P = 0.003$ OR = 0.57 [95% CI = 0.39-0.83]) and H3 ($P = 0.010$, OR = 0.54 [95% CI 0.33-0.86]) haplotypes was identified, but only the association with H1 remained after correction for multiple testing.

C4B copy number determination in aCSC

Carriers of two copies of the *C4B* gene were more frequent in the aCSC group (68%) compared to the control group (57%), whereas carrying three *C4B* gene copies was observed less frequently in the aCSC group (5.3% versus 18% in controls) (see Figure 2, Supplemental Digital Content 1, which demonstrates the *C4B* gene copy distribution). The distribution of *C4B* in aCSC patients compared to controls was significantly different after correction for multiple testing ($P = 0.0042$). The effect size of different *C4B* copy numbers on aCSC was assessed by a logistic regression model corrected for gender. The overall model was not significant ($P = 0.051$) (Table 4), but carriers of three *C4B* copies appeared to have a reduced risk of aCSC ($P = 0.002$, OR = 0.27 [95% CI = 0.12-0.63]) (Table 4).

Table 2. Analysis of 8 single nucleotide polymorphisms (SNPs) in acute central serous chorioretinopathy

SNP (gene)	Alleles (Major /Minor)	aCSC (n)	MAF aCSC	Controls (n)	MAF controls	Un-adjusted allelic P-value	Allelic odds ratio (95% CI)
rs10490924 (<i>ARMS2</i>)	G/T	132	0.174	812	0.217	0.111	0.76 (0.54-1.07)
rs2070951 (<i>NR3C2</i>)	C/G	132	0.538	1385	0.468	0.0287	1.33 (1.03-1.71)
rs800292 (<i>CFH</i>)	G/A	133	0.320	798	0.235	3.06×10^{-3}	1.53 (1.15-2.03)
rs1061170 (<i>CFH</i>)	T/C	133	0.259	803	0.353	2.82×10^{-3}	0.64 (0.48-0.86)
rs1065489 (<i>CFH</i>)	G/T	134	0.119	794	0.177	0.0199	0.63 (0.43-0.93)
rs1329428 (<i>CFH</i>)	C/T	133	0.579	787	0.429	5.87×10^{-6}	1.83 (1.40-2.38)
rs2284664 (<i>CFH</i>)	C/T	134	0.287	805	0.219	0.0132	1.44 (1.08-1.93)
rs3753394 (<i>CFH</i>)	C/T	131	0.263	800	0.293	0.324	0.86 (0.64-1.16)

aCSC = acute central serous chorioretinopathy; CI = Confidence interval; MAF = minor allele frequency; SNP = single nucleotide polymorphism; P-values < 0.0055 were considered significant.

Table 3. Complement factor H (CFH) gene haplotypes in aCSC

Haplo- types	Variants											
	rs3753394	rs800292	rs1061170	rs2284664	rs1329428	rs1065489	HF aCSC	HF controls	Un- adjusted allelic P-value	Allelic odds ratio (95% CI)	Un- adjusted allelic P-value	Allelic odds ratio (95% CI)
H1	C	G	C	C	C	G	0.249	0.329	Base	Base	0.003	0.57 (0.39-0.83)
H2	C	A	T	T	T	G	0.272	0.209	0.003	1.75 (1.21-2.53)	Base	Base
H3	T	G	T	C	C	T	0.102	0.158	0.799	0.94 (0.58-1.52)	0.010	0.54 (0.33-0.86)
H4	C	G	T	C	T	G	0.164	0.133	0.018	1.69 (1.09-2.6)	0.859	0.96 (0.64-1.46)
H5	T	G	T	C	T	G	0.114	0.072	0.001	2.3 (1.39-3.83)	0.280	1.32 (0.8-2.17)
Rare	*	*	*	*	*	*	0.100	0.098	0.107	1.52 (0.91-2.52)	0.578	0.87 (0.53-1.43)

aCSC = acute central serous chorioretinopathy; CI = Confidence interval; HF = haplotype frequency; MAF = minor allele frequency; P-values < 0.0083 were considered significant.

Differences between aCSC and cCSC

The minor allele frequencies of the tested *ARMS2*, *NR3C2*, and *CFH* variants were not significantly different between aCSC and cCSC patients (See Table 5, Supplemental Digital Content 2, which demonstrates minor allele frequencies in aCSC versus cCSC). Haplotype H4 in *CFH* showed a higher frequency in aCSC compared to cCSC (0.164 in aCSCs versus 0.111 in cCSCs, $P = 0.0250$, OR = 1.78 [95% CI = 1.08-2.95]), but this was not significant after correction for multiple testing (See Table 6, Supplemental Digital Content 3, which demonstrates *CFH* haplotypes in aCSC versus cCSC). The distribution of *C4B* copy numbers was not significantly different between aCSC and cCSC patients ($P = 0.345$, see Figure 2, Supplemental Digital Content 1), and the logistic regression model was also not significant ($P = 0.472$) (See Table 7, Supplemental Digital Content 4, which demonstrates logistic regression model for *C4B* load in aCSC versus cCSC).

Discussion

To the best of our knowledge, this is the first study to analyze potential genetic associations specifically in aCSC patients, and to compare them with known genetic associations that were previously identified in cCSC. We have found a significant association between three variants in the *CFH* gene and copy numbers of the *C4B* gene in patients with aCSC compared to healthy individuals. Among these *CFH* variants, two SNPs were risk-conferring and one was protective. Additionally, the H1 haplotype in the *CFH* gene was protective, whereas H2 and H5 were risk-conferring for aCSC. Three copy number of *C4B* conferred a protective effect for aCSC. No association was found between polymorphisms in *ARMS2* or *NR3C2* and the risk of aCSC. Finally, no significant differences were identified in these variants between aCSC and cCSC patients.

Genetic variation in different components of the complement system, which is an essential part of innate immunity,¹⁶ such as factor H (FH) and complement component 4B (*C4B*) proteins, have previously been associated with cCSC. Of the six tested variants in the *CFH* gene, five variants were associated with aCSC of which three were significant after correction for multiple testing. When comparing our findings

Table 4. Logistic regression model for *C4B* load

	Controls (n = 250)	aCSC patients (n = 133)	P-value	Odds ratio (95% CI)
Overall significance model P-value = 0.051				
Male sex	198 (79%)	91 (68%)	0.066	1.58 (0.97-2.57)
<i>C4B</i> copy number				
0	6 (2.4%)	4 (3.0%)	0.788	1.20 (0.33-4.39)
1	55 (22%)	32 (24%)	0.704	0.91 (0.54-1.51)
2	142 (57%)	90 (68%)	Base	Base
3	44 (18%)	7 (5.3%)	0.002	0.27 (0.12-0.63)
4	3 (1.2%)	0	0.999	NA

aCSC = acute central serous chorioretinopathy; CI = Confidence interval; NA = not annotated; P-values .0055 were considered significant.

in aCSC patients with available literature in cCSC patients, our data confirmed the protective effect of rs1061170, and the risk-conferring effects of rs1329428 and rs800292. However, for these variants the observed effect size in aCSC was larger than previously described in Caucasian cCSC patients, respectively, rs1061170 (OR = 0.64 versus 0.83), rs1329428 (OR = 1.83 versus 1.47) and rs800292 (OR = 1.53 versus 1.50).¹¹

Additionally, as observed in cCSC patients, the H1 haplotype in *CFH* was found to be protective for aCSC. Similar to the single variants, the protective effect of the H1 haplotype was stronger for aCSC compared to cCSC (OR = 0.57 versus 0.83).¹¹ The H2 and H5 haplotypes which were previously reported to increase the risk of cCSC, showed the same association in aCSC patients and their effect size was again larger in aCSC patients compared to cCSC (OR = 1.75 versus 1.33 and OR = 2.30 versus 1.37, respectively).¹¹ It has been suggested that factor H, which is encoded by the *CFH* gene, can influence the choroidal hemodynamic properties.¹⁷ Also, it has been suggested that an altered activity of factor H protein could cause RPE damage and dysfunction,^{11,17} but the exact mechanism of factor H in the etiology of CSC is still unknown. Both an absence and low copy numbers of *C4B* are known to increase the risk of cCSC, whereas carrying three copies is protective against cCSC.¹⁵ In our study, this protective effect was confirmed in aCSC patients, with an even larger effect size (OR = 0.27) compared to previous reports of cCSC.¹⁵

Exogenous administration of glucocorticoids, or an endogenous excess (Cushing syndrome) was previously found as an important risk factor in development of CSC.¹⁸ The glucocorticoid receptor and mineralocorticoid receptor are the most important targets for glucocorticoids, and therefore their involvement in the pathogenesis of CSC is conceivable.¹ We have previously found a significant association of a genetic variant in the *NR3C2* gene, encoding the mineralocorticoid receptor, with an increased risk of cCSC.¹⁴ In the present study, we did not find a significant association between the rs2070951 SNP in *NR3C2* with aCSC patients after correction for multiple testing, which could have occurred due to a lack of statistical power. Future larger studies can shed a light on whether this finding is indeed due to a lack of power or if it reflects a true difference in *NR3C2* rs2070951 risk SNP load between aCSC and healthy individuals.

In a previous study, we have found an association between genetic variations in the *ARMS2* gene and cCSC.¹¹ A possible mechanistic explanation for this association was speculated to be the potential interaction of the *ARMS2* protein with the extracellular matrix at the level of the choroid and RPE, which are also primarily affected in CSC.¹¹ Although the mechanism of action is not fully understood, presence of the rs10490924 variant in the *ARMS2* gene was shown to be protective against cCSC development.¹¹ This association with the rs10490924 SNP in *ARMS2* was not found in the current aCSC cohort. Again, this may be due to a lack of power, but could also indicate a difference in genetic predisposition between aCSC and healthy controls.

Acute CSC and cCSC generally show contrasting clinical presentations in terms of extent of retinal abnormalities and final visual outcome.^{1,19} There is currently no consensus on the classification of CSC, the definition of chronicity, and the exact period of time after which CSC should be considered chronic differs between studies,

ranging from two to six months.²⁰⁻²³ Besides a time based definition, cCSC is usually distinguished from aCSC by its more extensive retinal abnormalities on multimodal imaging, which includes multiple focal or diffuse leakage spots and widespread bilateral RPE alterations.^{2,9} A typical aCSC, on the other hand, presents with a single leakage spot, with only very few RPE changes. Although some patients with cCSC have a history of aCSC, many patients present with a chronic phenotype at the first presentation.^{9,24} Therefore, it is still unclear whether these two are part of a continuum with the same pathophysiological background, or if they are essentially different entities. A combination of genetic and non-genetic risk factors such as steroid use, hypertension, and pregnancy,²⁵⁻²⁷ may play a role in the aspect and severity of CSC, and the risk of progression of aCSC towards a chronic disease course.

Our data suggest a genetic overlap between aCSC and cCSC. No genetic difference could be found when comparing the selected variants in the cCSC and aCSC cohort. However, the effect size of the genetic variants associated with both aCSC and cCSC appears to be systematically larger in aCSC compared to cCSC. The lack of a significant difference between aCSC and cCSC with regard to the associated genetic variants may be partially caused by the small sample size of the current aCSC cohort, and thus a limited power. The larger effect size observed for aCSC suggests that genetic risk factors may play a larger role in the development of aCSC. It has been previously suggested that in multifactorial retinal diseases with genetic involvement such as age-related macular degeneration, patients who develop the disease at a younger age have a stronger genetic predisposition.^{28,29} A similar mechanism could explain the larger genetic predisposition among aCSC patients, who are generally younger than cCSC patients.⁹ Other limitations in the present study concern the control groups with different sample sizes, which might have influenced the observed associations. Also, there might be ethnical differences between German and Dutch patients, whom we considered as one Caucasian population.

In conclusion, variants rs800292 and rs1329428 in *CFH* gene were found to be significantly associated with a higher risk of aCSC, whereas variant rs1061170 in this gene was protective against aCSC. Three copy number of the *C4B* gene was protective against aCSC, and copy number of the gene differed between aCSC patients and controls. These specific *CFH* SNPs and the *C4B* copy numbers showed an even stronger association with aCSC than previously reported for cCSC. Our findings indicate that despite the differences in clinical presentation, acute and chronic CSC might share genetic risk and protective factors, at least among the currently known variants. Presumably, other non-genetic risk factors, or other currently unknown genetic variants are more influential in the differentiation toward an acute or a chronic disease course in CSC. Future genotype-phenotype correlation analyses in larger cohorts may provide important clues about interaction between these different risk factors.

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Footnotes and Financial Disclosures

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Key words: Acute central serous chorioretinopathy, *ARMS2* gene, chronic central serous chorioretinopathy, complement factor H, complement component 4, *CFH* gene, *C4B* copy numbers, genetic association, *NR3C2* gene

Summary Statement: Three variants in the *CFH* gene and copy numbers of *C4B* were found to be significantly associated with the risk of acute CSC in Caucasians. Acute and chronic CSC may share a similar genetic predisposition based on the tested genetic variants in *ARMS2*, *CFH*, and *NR3C2* genes, and copy numbers of the *C4B* gene.

Running head: Genetics in acute CSC

The authors report no conflicts of interest.

Supplementary Files

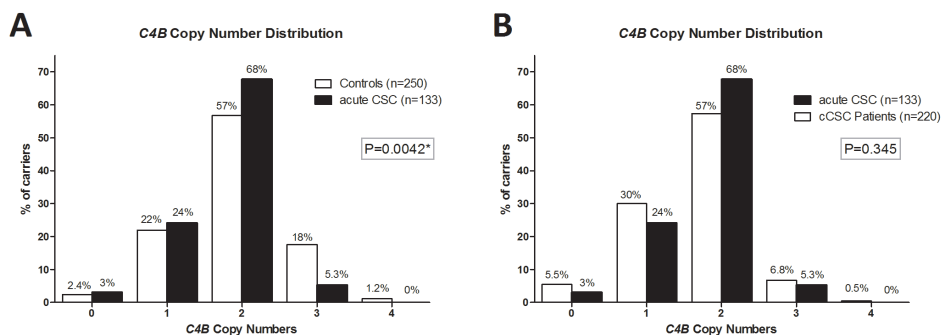


Figure 2: (supplementary) Distribution of *C4B* copy numbers. (A) Comparison between the *C4B* distribution among acute central serous chorioretinopathy (CSC) and controls, (B) and acute CSC and chronic CSC.

Table 7 (supplementary). Logistic regression model for *C4B* load aCSC versus cCSC

	cCSC patients (n = 220)	aCSC patients (n = 133)	P-value	Odds ratio (95% CI)
Male sex	216 (79%)	91 (68%)	0.026	1.75 (1.07-2.88)
<i>C4B</i> copy number				
0	12 (5.4%)	4 (3.0%)	0.247	0.50 (0.16-1.61)
1	66 (30%)	32 (24%)	0.142	0.69 (0.41-1.14)
2	126 (57%)	90 (68%)	Base	Base
3	15 (6.8%)	7 (5.2%)	0.361	0.64 (0.25-1.66)
4	1 (0.4%)	0 (0%)	1.000	NA

aCSC = acute central serous chorioretinopathy; cCSC = chronic central serous chorioretinopathy; CI = confidence interval; NA = not annotated; P-values <0.0055 were considered significant.

Table 5 (supplementary). Comparison of allele frequencies in aCSC versus cCSC

SNP (gene)	aCSC (n)	MAF aCSC	cCSC (n)	MAF cCSC	Unadjusted allelic P-value	Allelic odds ratio (95% CI)
rs10490924 (ARMS2)	132	0.174	243	0.193	0.520	0.88 (0.60-1.30)
rs2070951 (NR3C2)	132	0.538	269	0.520	0.642	1.07 (0.80-1.44)
rs800292 (CFH)	133	0.320	245	0.296	0.500	1.12 (0.81-1.54)
rs1061170 (CFH)	133	0.259	245	0.320	0.0801	0.74 (0.53-1.04)
rs1065489 (CFH)	134	0.119	244	0.133	0.587	0.88 (0.56-1.39)
rs1329428 (CFH)	133	0.579	244	0.510	0.0707	1.32 (0.98-1.78)
rs2284664 (CFH)	134	0.287	244	0.275	0.709	1.07 (0.76-1.48)
rs3753394 (CFH)	131	0.263	242	0.273	0.783	0.95 (0.68-1.34)

aCSC = acute central serous chorioretinopathy; cCSC = chronic central serous chorioretinopathy; CI = confidence interval; MAF = minor allele frequency; SNP = single nucleotide polymorphism; P-values < 0.0055 were considered significant.

Table 6 (supplementary). Complement factor H (CFH) gene haplotypes in aCSC versus cCSC

Variants												
Haplo- types	rs3753394	rs800292	rs1061170	rs2284664	rs1329428	rs1065489	HF aCSC	HF cCSC	Un- adjusted allelic P-value	Allelic odds ratio (95% CI)	Un- adjusted allelic P-value	Allelic odds ratio (95% CI)
H1	C	G	C	C	C	G	0.249	0.301	Base	Base	0.219	0.77 (0.51-1.16)
H2	C	A	T	T	T	G	0.272	0.262	0.219	1.29 (0.86-1.94)	Base	Base
H3	T	G	T	C	C	T	0.102	0.104	0.408	1.28 (0.71-2.30)	0.976	0.99 (0.57-1.73)
H4	C	G	T	C	T	G	0.164	0.111	0.025	1.78 (1.08-2.95)	0.201	1.38 (0.84-2.26)
H5	T	G	T	C	T	G	0.114	0.115	0.373	1.28 (0.75-2.18)	0.963	0.99 (0.59-1.66)
Rare	*	*	*	*	*	*	0.100	0.108	0.515	1.21 (0.68-2.16)	0.822	0.94 (0.54-1.64)

aCSC = acute central serous chorioretinopathy; cCSC = chronic central serous chorioretinopathy; CI = confidence interval; HF = haplotype frequency; MAF = minor allele frequency, P-values < 0.0083 were considered significant.

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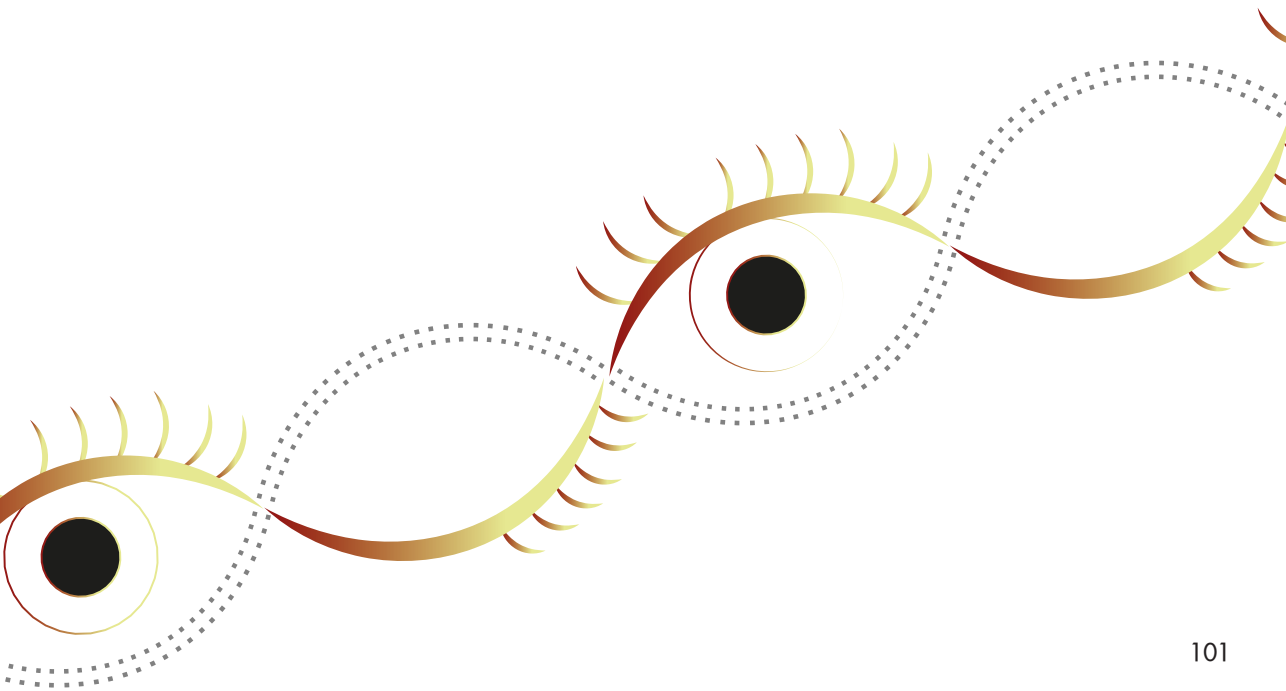
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3.1 Role of the complement system in chronic central serous chorioretinopathy: a genome-wide association study

3.2 Exome sequencing identifies *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* as susceptibility genes for chronic central serous chorioretinopathy in females

3. Unbiased Approaches





3.1 Role of the complement system in chronic central serous chorioretinopathy: A Genome-wide association study

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Abstract

Importance: To date, several targeted genetic studies on chronic central serous chorioretinopathy (cCSC) have been performed; however, unbiased genome-wide studies into the genetics of cCSC have not been reported. To discover new genetic loci associated with cCSC and to better understand the causative mechanism of this disease, we performed a genome-wide association study (GWAS) on patients with cCSC.

Objective: To discover new genetic loci and pathways associated with cCSC and to predict the association of genetic variants with gene expression in patients with cCSC.

Design, Setting and Participants: This case-control GWAS was completed in the general community, 3 referral university medical centers, and outpatient care on European individuals with cCSC and population-based control participants. Genotype data was collected from May 2013 to August 2017, and data analysis occurred from August 2017 to November 2017.

Main outcomes and measurements: Associations of single-nucleotide polymorphisms, haplotypes, genetic pathways, and predicted gene expression with cCSC.

Results: A total of 521 patients with cCSC (median age 51 years; interquartile range [IQR], 44-59 years; 420 [80.6%] male) and 3577 European population-based control participants (median age, 52 years; IQR, 37-71 years; 1630 [45.6%] male) were included. One locus on chromosome 1 at the complement factor H (*CFH*) gene reached genome-wide significance and was associated with an increased risk of cCSC (rs1329428; odds ratio [OR], 1.57 [95% CI, 1.38-1.80]; $P = 3.12 \times 10^{-11}$). The *CFH* haplotypes H1 and H3 were protective for cCSC (H1: OR, 0.64 [95% CI, 0.53-0.77]; $P = 2.18 \times 10^{-6}$; H3: OR, 0.54 [95% CI, 0.42-0.70]; $P = 2.49 \times 10^{-6}$), whereas haplotypes H2, H4, H5, and the aggregate of rare *CFH* haplotypes conferred increased risk (H2: OR, 1.57 [95% CI, 1.30-1.89]; $P = 2.18 \times 10^{-6}$; H4: OR, 1.43 [95% CI, 1.13-1.80]; $P = 2.49 \times 10^{-3}$; H5: OR, 1.80 [95% CI, 1.36-2.39]; $P = 4.61 \times 10^{-5}$; rare haplotypes: OR, 1.99 [95% CI, 1.43-2.77]; $P = 4.59 \times 10^{-5}$). Pathway analyses showed involvement of the complement cascade and alternative open reading frame (*ARF*) pathway in cCSC. Using PrediXcan, we identified changes in predicted expression of complement genes *CFH*, complement factor H related 1 (*CFHR1*), complement factor related 4 (*CFHR4*), and membrane cofactor protein (*MCP/CD46*). Additionally, the genes potassium sodium-activated channel subfamily T member 2 (*KCNT2*) and tumor necrosis factor receptor superfamily member 10a (*TNFRSF10A*) were differentially expressed in cCSC patients.

Conclusions and relevance: In this GWAS on cCSC, we identified a locus on chromosome 1 at the *CFH* gene that was significantly associated with cCSC, and we report protective and risk-conferring haplotypes in this gene. Pathway analyses were enriched for complement genes, and gene expression analysis suggests a role for *CFH*, *CFHR1*, *CFHR4*, *CD46*, *KCNT2*, and *TNFRSF10A* in the disease. Taken together, these results underscore the potential importance of the complement pathway in the causative mechanism of cCSC.

Introduction

Central serous chorioretinopathy (CSC) is characterized by subretinal fluid accumulation between the neuroretina and the retinal pigment epithelium (RPE).¹⁻³ Patients usually present with clinical symptoms of metamorphopsia and central vision impairment.^{4,5} Although acute CSC can show spontaneous resolution within 3 months, patients with chronic CSC (cCSC) have prolonged presence of fluid with progressive loss of vision, permanent RPE alterations, and a reduced quality of life.⁴ Generally, CSC occurs more frequently in males (9.9 per 100 000) than females (1.7 per 100 000) and often presents at an age at which patients are still professionally active.⁶

The exact causative mechanism of cCSC is unknown, but clinical and experimental studies have implicated dysfunction of the RPE and hyperpermeability of the choroid in CSC.⁴ The disease has been associated with the use of corticosteroids, stress, and type A personality. Reports of familial occurrences of cCSC support a genetic component for the disease.^{1,3,4,7-11} Genetic studies on cCSC have been limited to candidate gene approaches; several associations have been reported with single-nucleotide polymorphisms (SNPs) in the genes age-related macular degeneration susceptibility 2 (*ARMS2*), cadherin 5 (*CDH5*), complement factor H (*CFH*) and nuclear receptor subfamily 3 group C member 2 (*NR3C2*), as well as copy number variations in the complement factor 4B (*C4B*) gene.¹²⁻¹⁶

Unbiased genome-wide association studies (GWAS) have identified hundreds of genomic loci implicated in complex diseases, such as age-related macular degeneration (AMD), myopia, and glaucoma.¹⁷ Such approaches have shed light on the genetic architecture of these diseases; however, establishing functional links between identified loci and disease remains challenging. Besides having direct effects on protein function or structure, SNPs can also influence the expression of nearby (cis) or distal (trans) genes. Such regulatory genetic variants are called expression quantitative trait loci (eQTL).¹⁸ Several eQTL databases, linking genotype information to tissue expression of genes, have been established. One of the largest of these projects is the GTEx project, in which expression profiles of 44 tissues of 449 donors have been collected in version 6.¹⁸ These eQTL databases offer functional information on SNPs identified in GWAS, contribute to a better understanding of the studied disease trait, and can be used to predict expression in genotyped samples.¹⁹

In this study, we performed a GWAS involving European patients with cCSC and population control participants to identify new cCSC disease loci and to increase our knowledge on the causative mechanism of this disease. Additionally, we performed pathway analyses to discover new pathways implicated in cCSC. Using publicly available eQTL data, we aimed to identify new candidate genes predicted to be differentially expressed in patients with cCSC compared with control participants.

Key points

Question: Which genetic loci associate with chronic central serous chorioretinopathy (cCSC) when using an unbiased genome-wide approach?

Findings: In this case-control genome-wide association study, a locus containing multiple variants in the *CFH* gene was associated with cCSC. Additionally, pathway

analysis implicated the complement system and based on genetic data, components of the complement system were predicted to have an association with altered expression in patients with cCSC.

Meaning: This study of participants with cCSC demonstrates the potential role of the *CFH* gene and the complement system in the causative mechanism of cCSC.

Methods

Study participants

Genomic DNA was extracted from blood using standard procedures. In total, 546 European patients with cCSC recruited from the outpatient clinics of the Radboud university medical center (Radboudumc), Nijmegen, the Netherlands, University Hospital of Cologne (UHC), Cologne, Germany, and Leiden University Medical Center (LUMC), Leiden, the Netherlands, were included. Grading of all patients was performed by an experienced retinal specialist (C.J.F.B.), and was based on extensive ophthalmological examination including fundoscopy, spectral-domain optical coherence tomography, fluorescein angiography and indocyanine green angiography. Diagnosis of cCSC was based on subgroups 1 and 2, which have been previously described.¹³ Briefly, patients that were included in this study showed the presence of serous fluid on optical coherence tomography in 1 or both eyes, either bilateral (subgroup 1) or unilateral (subgroup 2) RPE irregularities with 1 or more hot spots of leakage on fluorescein angiography, and corresponding hyperfluorescence on indocyanine green angiography.¹³ Patients diagnosed with acute CSC as recognized by a focal leakage spot (ink blot) or a smokestack pattern on fluorescein angiography, with less than 1 disc diameter of adjacent atrophic RPE alterations, and/or a duration of disease of less than 3 months were excluded from the study. Patients in whom evidence of another explanatory diagnosis or complication was present (subgroup 3), such as polypoidal choroidal vasculopathy, choroidal neovascularization, drusen or other signs of AMD, were excluded from this study.

This study was carried out in accordance with the tenets of the Declaration of Helsinki and was approved by the local ethics committees of the Radboudumc, LUMC and UHC. Written informed consent was obtained for all subjects involved in the study.

Genotyping data of controls was obtained from the Nijmegen Biomedical Study (NBS), a population-based survey conducted by the Department for Health Evidence and the Department of Laboratory Medicine of the Radboudumc. (eMethods in supplement, <http://www.nijmegenbiomedischestudie.nl/>).²⁰ In this population-based study, no ophthalmologic grading was performed. Only controls for which genotyping was available on the OmniExpress platform (n=3654) were included in this study. Genotype data was collected from May 2013 to August 2017, and data analysis occurred from August 2017 to November 2017.

The mean age of patients with cCSC and control participants was compared with a Mann-Whitney U test. The sex distribution was compared with a X^2 test using SPSS version 22 (IBM). All values of $P < .05$ were deemed significant.

Genome-wide association and haplotype analyses

Genotyping was performed with the OmniExpress-12 or OmniExpress-24 chips, and data were imputed with the Haplotype Reference Consortium release 1.1.2016. After stringent quality control (eMethods in the Supplement), 521 patients with cCSC and 3577 NBS control participants were included in the analysis, carrying 11 261 291 autosomal SNPs and 265 428 X-chromosomal SNPs. Single-variant association analysis was performed using the Firth bias-corrected likelihood ratio test implemented in EPACTS (version 3.2.6, <http://genome.sph.umich.edu/wiki/EPACTS>; Univeristy of Michigan),²¹ correcting for sex and the first 2 components of the ancestry analysis, to correct for potential population stratification (eMethods in the Supplement). To assess the potential role of confounding by AMD, the analysis was repeated in patients and controls below younger than 51 years.

Haplotype analysis, combining multiple variants in the *CFH* gene, was performed with the haplo.stats²² (version 1.7.7) package in R (R Foundation for Statistical Computing).²³ The SNPs described by Hageman et al.²⁴ and the *CFH* SNPs that were previously associated with cCSC were used as input.^{12,13} Haplotypes with a frequency greater than 1% were analyzed individually, while rare haplotypes (with a frequency less than 1%) were aggregated. Haplotype association with cCSC was performed using the haplo.glm in R, correcting for sex and the first 2 principal components of the ancestry analysis. The analysis was performed using the most common protective haplotype (H1) and cCSC risk-carrying haplotype (H2) as references. The frequency of the haplotypes in patients with cCSC and control participants was obtained using the haplo.cc command in R.

Pathway and PrediXcan analysis

We used GWAS summary statistics to perform competitive gene-set analysis to identify pathways associated with cCSC using 2 different programs: MAGMA (version 1.06, <https://ctg.cncr.nl/software/magma>) and VEGASv2 (version 2, <https://vegas2.qimrberghofer.edu.au/vegas2v2>).^{25,26} Additionally, PrediXcan (Hakyimlab; <https://github.com/hakyim/PrediXcan>) was used to predict gene expression levels based on publicly available eQTL data of GTEx.²⁷ PrediXcan was performed on all 44 provided GTEx tissues using genetic variants in 17 742 genes. We selected the genes that were associated in at least 1 tissue after Bonferroni correction for multiple testing ($P < .05/17742$ genes = 2.21×10^{-6}) and were nominally associated ($P < .05$) in at least 50% of the tissues they were expressed in. (The eMethods in the Supplement provide further clarification). This study had sufficient power to detect common variants with OR greater than 1.5 and had more than 80% power to detect the lead variant (eMethods and eFigure 1 in the Supplement).

Results

Genome-wide association analysis

A total of 546 patients with cCSC were recruited from Radboudumc (n = 319), UHC (n = 74), and LUMC (n = 153). In addition, 3654 European population-based control participants were included. After quality control (eFigures 2 and 3 in the Supplement), a total of 521 European patients with cCSC (median age, 51 years; interquartile range [IQR], 44-59 years; 420 [80.6%] male) and 3577 controls (median age, 52 years; IQR,

37-71 years; 1630 [45.6%] male; eTable 1 in the Supplement) were analyzed for 11 261 291 autosomal and 265 428 X-chromosomal SNPs. Because of the significant difference in sex between patients with cCSC and control participants (eTable 1 in the Supplement, $P = 1.78 \times 10^{-50}$), sex was included as a covariate in the analysis along with 2 principal components of the ancestry analysis.

The GWAS identified 20 SNPs that reached genome-wide significance (using a cutoff value for significance set at $P < 5.0 \times 10^{-8}$) that all resided at 1 locus on chromosome 1 in the *CFH* gene (lead variant: rs1329428; odds ratio [OR], 1.57 [95% CI, 1.38-1.80]; $P = 3.12 \times 10^{-11}$; Figure 1; Table 1; eFigure 5, and eTable 2 in the Supplement). Additionally, six suggestive signals ($P < 1 \times 10^{-6}$) were found on chromosome 1 (OR, 0.64 [95% CI, 0.54-0.77]; $P = 6.69 \times 10^{-7}$), chromosome 8 (OR, 42.25 [95% CI, 9.55-186.89]; $P = 1.33 \times 10^{-7}$), chromosome 9 (OR, 8.36 [95% CI, 3.67-19.06]; $P = 6.28 \times 10^{-7}$), chromosome 15 (OR, 206.11 [95% CI, 4.50-9436.68]; $P = 7.84 \times 10^{-7}$), chromosome 17 (OR, 5.25 [95% CI, 2.94-9.48]; $P = 1.51 \times 10^{-7}$), and chromosome 20 (OR, 0.67 [95% CI, 0.57-0.79]; $P = 7.14 \times 10^{-7}$; Figure 1; Table 1). Conditioned analysis of the lead SNP on chromosome 1 did not reveal any other independent signal in the *CFH* gene (eFigure 4 in the Supplement), but did retain the suggestive signal at the *CD46* gene on chromosome 1 indicating that the *CD46* signal is independent of the *CFH* association (unconditioned and conditioned data are shown in eFigure 4 the Supplement). Stratified analysis on patients and controls younger than 51 years identified associations with the same 20 SNPs in the *CFH* gene, but ORs were higher for the lead variant than in the complete cohort (OR, 1.89 [95% CI, 1.56-2.32]; $P = 1.23 \times 10^{-10}$; eTable 2 in the Supplement).

107

Haplotype analysis

To further characterize the association at the *CFH* gene, haplotype analysis was performed. A logistic model corrected for sex and 2 principal components was performed using either the most common cCSC protective (H1) haplotype or the most common cCSC risk-conferring (H2) haplotype as reference (Table 2). The H2, H4, and H5 haplotypes, all containing the minor allele of the lead variant, were associated with an increased risk of cCSC (H2: OR, 1.57 [95% CI, 1.30-1.89]; $P = 2.18 \times 10^{-6}$; H4: OR, 1.43 [95% CI, 1.13-1.80]; $P = 2.49 \times 10^{-3}$; H5: OR, 1.80 [95% CI, 1.36-2.39]; $P = 4.61 \times 10^{-5}$). The H1 and H3 haplotypes, containing the major allele of the lead variant, were protective (H1: OR, 0.64 [95% CI, 0.53-0.77]; $P = 2.18 \times 10^{-6}$; H3: OR, 0.54 [95% CI, 0.42-0.70]; $P = 2.49 \times 10^{-6}$). Additionally, the aggregate of rare haplotypes also increased the risk of cCSC (OR, 1.99 [95% CI, 1.43-2.77]; $P = 4.59 \times 10^{-5}$). All associations were significant after Bonferroni correction for multiple testing for 10 tests (using a cutoff value of $P < .005$).

Gene-set/Pathway analysis

Using the GWAS summary statistics, we performed competitive gene-set analysis using 2 programs. MAGMA identified 3 pathways that were associated with cCSC after correction for multiple testing (Table 3, genes in eTable 3 in the Supplement). Two of the pathways are implicated in the complement cascade, and 1 is the *ARF* pathway. We identified the same reactome pathways of the complement cascade using VEGAS2. However, these findings were not significant after correction for multiple testing (using a cutoff value of $P < 1 \times 10^{-5}$; Table 3, genes in eTable 4 in the

Supplement).

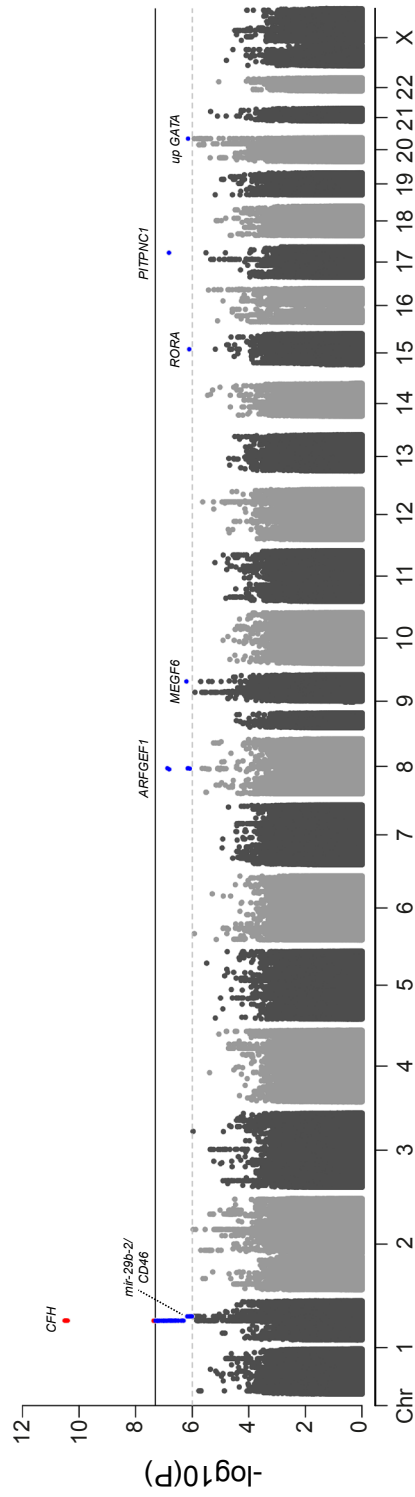


Figure 1 : Manhattan Plot of Genome-Wide Association Study on 521 Patients With Chronic Central Serous Chorioretinopathy and 3577 Population Control Participants. Genome-wide association analysis was performed correcting for sex and 2 principal components. The genome-wide significant signals are depicted in red (significant with cutoff value set at $P < 5 \times 10^{-8}$), whereas the suggestive variants are depicted in blue (significant with cutoff value set at $P < 1 \times 10^{-9}$).

Table 1. Top hits in chronic central serous chorioretinopathy GWAS study

Top Single-Nucleotide Polymorphism	Chr	Base Pair (hg19)	Major Allele/ Minor allele	Gene	Minor Allele Frequency in Patients with cCSC (n=521)	Minor Allele Frequency in control participants (n=3577)	OR (95% CI)	P-value
rs1329428 ^a	1	196702810	C/T	CFH	0.543	0.434	1.57 (1.38-1.80)	3.12x10 ⁻¹¹
rs561753392	8	68131981	C/T	ARFGEF1	0.0070	0.0005	42.25 (9.55-186.89)	1.33x10 ⁻⁷
rs76766498	17	65418162	A/G	PITPNC1	0.021	0.005	5.25 (2.94-9.48)	1.51x10 ⁻⁷
rs118083394	9	123367552	C/T	MEGF6	0.014	0.002	8.36 (3.67-19.06)	6.28x10 ⁻⁷
rs4844392	1	207991209	C/G	mir-29b-2/ CD46	0.159	0.230	0.64 (0.54-0.77)	6.69x10 ⁻⁷
rs2379120	20	61030580	T/A	upstream GATA5	0.218	0.300	0.67 (0.57-0.79)	7.14x10 ⁻⁷
rs541395042	15	61175324	C/T	RORA	0.006	0.00	206.11 (4.50-9436.68)	7.84x10 ⁻⁷

ARFGEF1: adenosine diphosphate ribosylation factor guanine nucleotide exchange factor 1; bp: base pair; cCSC: chronic central serous chorioretinopathy; CD46: cluster of differentiation 46; CFH: complement factor H; chr: chromosome; CI: confidence interval; GATA5: GATA-5 binding protein 5; GWAS: genome-wide association study; hg: human genome; MAF: minor allele frequency; MEGF6: multiple epidermal growth factor like domains 6; mir-29b-2: micro ribonucleic acid 29b-2; OR: odds ratio; PITPNC1: phosphatidylinositol transfer protein cytoplasmic 1; RORA: retinoic acid receptor-related orphan receptor alpha; ^a This is a genome-wide significant hit

Table 2. Haplotype analysis of the Complement factor H gene

Haplo type	Frequency in Controls		Frequency in patients with cCSC		Per CFH-H1 ^b		Per CFH-H2 ^c		P-value Corrected ^{d,e}
					Odds Ratio (95% CI)	P-value Corrected ^{d,e}	Odds Ratio (95% CI)	P-value Corrected ^{d,e}	
rs3753394	C	G	0.339	0.273	1 [REF]	1 [REF]	0.64 (0.53-0.77)	2.18x10 ^{-6f}	1 [REF]
rs529825	A	A	0.221	0.274	1.57 (1.30-1.89)	2.18x10 ^{-6f}	0.54 (0.42-0.70)	2.49x10 ^{-6f}	0.442
rs800292	G	G	0.135	0.094	0.85 (0.66-1.09)	0.204	0.91 (0.72-1.15)	0.343	
rs3766404	T	G	0.115	0.128	1.43 (1.13-1.80)	4.61x10 ^{-5f}	1.15 (0.86-1.53)	0.191	
rs1061170	C	T	0.052	0.078	1.80 (1.36-2.39)	0.492	0.74 (0.48-1.16)	0.021	
rs3753396	A	A	0.030	0.025	1.17 (0.75-1.81)	0.485	0.53 (0.30-0.91)	0.972	
rs203674	G	T	0.027	0.018	0.82 (0.48-1.42)	0.048	1.01 (0.64-1.59)	0.421	
rs2284664	C	C	0.022	0.027	1.58 (1.00-2.48)	0.418	0.80 (0.46-1.38)	0.163	
rs1329428 ^g	C	T	0.017	0.017	1.25 (0.73-2.17)	4.59x10 ^{-5f}	1.27 (0.91-1.77)		
rs1065489	G	G	0.042	0.066	1.99 (1.43-2.77)				

cCSC: chronic central serous chorioretinopathy; CI: confidence interval; NA: not applicable; REF: reference; ^a Results shown in the rs1329428 column are top hits for the genome-wide association study findings. ^b Analysis performed with the most common protective haplotype CFH-H1 set as a reference. ^c Analysis performed with the most common risk-conferring haplotype CFH-H2 set as a reference. ^d Bonferroni correction for multiple testing for 10 haplotypes with significance set at $P < .05/10 = .005$. ^e Analysis was performed correcting for sex and the first 2 principal components of the ancestry analysis. ^f Values are significant based on a cutoff level of $P < .005$. ^g The data on rare haplotypes includes an accumulation of alleles that could be either minor or major.

Table 3. Competitive Gene-Set Analysis of Multi-Marker Analysis of Genomic Annotation (MAGMA) and Versatile Gene-Based Association Study 2 (VEGAS2)

Pathway name	Genes, No	Beta (se)	P value	
			Nominal	Empirical corrected
MAGMA				
Reactome Regulation of Complement Cascade	13	1.29 (0.265)	5.49×10^{-7}	8.90×10^{-6a}
Reactome Complement Cascade	29	0.786 (0.179)	5.56×10^{-6}	7.37×10^{-3a}
Biocarta ARF pathway	17	0.848 (0.194)	6.49×10^{-6}	8.65×10^{-3a}
VEGAS2				
Reactome: R-HSA-977606 Regulation of Complement Cascade	13	NA	1.03×10^{-7}	1.60×10^{-5}
Reactome: R-HSA-166658 Complement Cascade	22	NA	6.09×10^{-7}	4.20×10^{-5}

arf: alternative open reading frame; NA: not annotated; se: standard error; ^a significant after correction for multiple testing; the MAGMA nominal significance threshold is $P < 4.12 \times 10^{-5}$ and the VEGAS2 empirical significance threshold is $P < 1 \times 10^{-5}$

Table 4. PrediXcan results, genes differentially expressed between cCSC patients and controls

Gene	Tissues with expression, No	Tissues with differential expression, No. (%) ^a	Tissues ^b	Odds Ratio (95% CI)	P value
CFHR4	1	1 (100)	Liver	0.91 (0.85-0.98)	1.29x10 ⁻⁶
CD46	25	21 (84)	Esophagus Muscularis	1.05 (1.03-1.07)	1.79x10 ⁻⁶
			Skin Not Sun Exposed Suprapubic	1.32 (1.18-1.48)	2.26x10 ⁻⁶
			Heart Atrial Appendage	1.16 (1.09-1.24)	2.39x10 ⁻⁶
			Colon Sigmoid	1.07 (1.04-1.09)	2.57x10 ⁻⁶
TNFRSF10A	18	15 (83)	Cells Transformed fibroblasts	0.96 (0.94-0.98)	2.41x10 ⁻⁶
			Adrenal Gland	0.95 (0.93-0.97)	2.60x10 ⁻⁶
			Hypothalamus	0.78 (0.72-0.85)	1.73x10 ⁻⁸
			Adipose Subcutaneous	1.15 (1.1-1.21)	1.85x10 ⁻⁸
CFHR1	12	9 (75)	Liver	0.95 (0.93-0.97)	1.87x10 ⁻⁷
			Cells Transformed fibroblasts	0.79 (0.71-0.87)	1.41x10 ⁻⁶
KCNT2	5	3 (60)	Esophagus Gastroesophageal Junction	3.70 (2.27-6.04)	1.62x10 ⁻⁷
			Nerve Tibial	1.13 (1.08-1.18)	4.99x10 ⁻⁷
CFH	5	3 (60)	Adipose Subcutaneous	1.09 (1.05-1.13)	1.17x10 ⁻⁶
			Nerve Tibial	1.10 (1.06-1.14)	1.63x10 ⁻⁶

Abbreviations: **CD46**: cluster of differentiation 46; **CFH**: complement factor H; **CFHR1/4**: complement factor H related 1/4; **GTEx**: genotype-tissue expression; **KCNT2**: Potassium Sodium-Activated Channel Subfamily T Member 2; **OR**: odds ratio; **TNFRSF10A**: tumor necrosis factor receptor superfamily member 10a; ^a The cutoff value for significance in differential expression was P < .05; only those genes that showed expression differences in more than 50% of their expressed tissues are depicted; ^b This section used a cutoff value for significance at P < 2.81 x 10⁻⁶.

PrediXcan analysis

For each individual, we predicted the association of genotypes with expression levels of 17 742 genes in 44 different GTEx (version 6) tissues using PrediXcan. The predicted gene expression of genes at the *CFH* locus (*CFH*, *CFHR1*, *CFHR4* and *KCNT2*) was different between cCSC patients and controls after correction for multiple testing, using a cutoff value for significance of $P < 2.82 \times 10^{-6}$ (for example, *CFH* in subcutaneous adipose tissue: OR, 1.09 [95% CI, 1.05-1.13]; $P = 1.17 \times 10^{-6}$; *CFHR1* in subcutaneous adipose tissue: OR, 1.15 [95% CI, 1.10-1.21]; $P = 1.85 \times 10^{-8}$; *CFHR4* in liver: OR, 0.91 [95% CI, 0.85-0.98]; $P = 1.29 \times 10^{-6}$; *KCNT2* in gastroesophageal junction: OR, 3.70 [95% CI, 2.27-6.04]; $P = 1.62 \times 10^{-7}$; full results in Table 4). Additionally, we observed altered predicted expression of the *CD46* and *TNFRSF10A* genes (Table 4, results of all tissues in eTable 5 in the Supplement).

Discussion

In this unbiased genome-wide association study of cCSC, we identified a locus for cCSC on chromosome 1, at the *CFH* gene, that had previously been described in 2 targeted candidate gene studies.^{12,13} We discovered protective and risk haplotypes in *CFH* and found evidence for involvement of rare *CFH* haplotypes in the disease. Moreover, using gene-set analysis and publicly available expression databases, we uncovered additional evidence for altered regulation of the complement system in cCSC and identified novel candidate genes and pathways implicated in this disease.

So far, to our knowledge, the association of cCSC with the *CFH* gene is the only genetic association for cCSC that has been replicated in multiple independent studies.^{12,13,28} The effect size of the lead-variant (rs1329428) in *CFH* in this unbiased study was higher compared to the previous targeted study,¹³ but lower compared to the study by Miki et al.¹² (with ORs in the 3 studies 1.57, 1.47, and OR=1.79, respectively). Stratified analysis of patients younger than 51 years to excluded potential confounding with AMD showed the same genome-wide significant 20 SNPs with similar direction of effect, confirming the cCSC specific association. Additionally, haplotype analysis of the *CFH* locus confirmed a previously reported protective effect of *CFH*-H3,¹³ and identified associations with the protective *CFH*-H1 and risk carrying *CFH*-H2, *CFH*-H4, and *CFH*-H5 haplotypes. The risk associated with the *CFH*-H5 haplotype was higher than the risk caused by the single lead variant alone and higher to our knowledge than any previously reported SNP in *CFH*, with an OR of 1.80 for *CFH*-H5 compared to OR of 1.57 for rs1329428. Interestingly, an aggregate of all haplotypes with a frequency of less than 1% also showed a higher risk for cCSC (OR, 1.99). Higher ORs of *CFH*-H5 and the rare haplotype aggregate suggest that other (rare) variants in these haplotypes might play a role in cCSC.

The factor H protein, encoded by the *CFH* gene, is able to block the formation of C3-convertases and therefore is an important regulator of the complement system.²⁹ The *CFH* gene has been widely studied for its role in AMD, and, interestingly, variants that confer risk for AMD are protective for cCSC and vice versa.¹³ These genetic associations imply that these 2 diseases might have an opposite disease mechanism, with an overactivation of the complement system in AMD²⁹ and a reduced activity of the complement system in cCSC.

In the current study, we further substantiated the involvement of the complement system in cCSC. Pathway analysis with MAGMA and VEGAS2 showed associations of 2 gene sets of the complement system. We used PrediXcan to predict the expression of genes based on the genotype information of patients with cCSC and control participants. Unfortunately, no eye-specific or retina-specific eQTL database is publicly available, and these tissues are not yet implemented in GTEx nor in any other large eQTL data set. Therefore, as an indicator for general differences, we used all GTEx tissues and observed that the expression of 4 complement genes (*CFH*, *CFHR1*, *CFHR4* and *CD46*) was predicted to be different in patients with cCSC compared to control participants. All GTEx tissues that express *CFH* showed an upregulation of *CFH*, suggesting that complement system activity may be reduced in cCSC (Table 4). Depending on the tissue, *CFHR1* showed upregulation or down-regulation, indicating that regulation of this gene might be tissue specific; meanwhile *CFHR4* was consistently down-regulated (Table 4). The factor H-associated (FHR) proteins, encoded by the *CFHR* genes, show sequence similarities to factor H and can compete with factor H for C3b binding and in this manner can influence complement activation.³⁰ Deletions of *CFHR1* have been found to be protective for AMD and *CFHR4* deletions have been implicated in atypical hemolytic uremic syndrome,³⁰ but the genes have not previously been associated with cCSC. to our knowledge.

Interestingly, we also observed an association with another complement gene independent of the *CFH* locus: *CD46* was identified as one of the sub-threshold hits in the GWAS (Table 1), and PrediXcan predicted up-regulation of *CD46* expression (Table 4). The *CD46* gene encodes the membrane cofactor protein (MCP/*CD46*), a complement inhibitor that blocks all pathways of the complement cascade through inactivation of C3b and C4b.^{31,32} In light of the clinical characteristics of cCSC, in which patients present with hyperpermeability of the choroid and with fluid leakage through the retinal pigment epithelium, the involvement of *CD46* is particularly interesting. It is an important regulator in the maintenance of epithelial cell barrier integrity through interaction with, for example, cadherins and integrins.^{32,33} Previous studies have shown that the activation of *CD46* leads to a decrease in transepithelial resistance and loss of tight junctions in intestinal cells and a decreased membrane adhesion in RPE cells.^{32,33} The predicted increased *CD46* expression in patients with cCSC therefore suggests a downregulation of complement activity, similarly as observed for factor H, and in addition destabilization of the integrity of the RPE, one of the main hallmarks of CSC. Together, both the complement and epithelial regulatory function of *CD46* make it an interesting protein for further study in cCSC.

Taken together, the associations and predicted altered expression of *CD46*, *CFH*, *CFHR1*, and *CFHR4* with cCSC imply a reduced activity of the complement system in cCSC. To date and to our knowledge, limited information is available on the actual activity of the complement system in patients with cCSC. One study did not find obvious systemic alterations in blood complement components of patients with cCSC, but this study had limited power, and *CD46*, *FHR1*, and *FHR4* levels were not measured.³⁴ Combined with previously described associations of *CFH* and *C4B*¹²⁻¹⁴ and the associations described here, a central role for the complement system in the disease mechanism of cCSC emerges. Larger studies measuring systemic complement regulators and activation products in patients with cCSC (specifically *FH*, *FHR1*,

FHR4, C4B, and CD46) are warranted.

Besides consolidating the involvement of the complement system, we also identified a new cCSC candidate pathway and 2 new cCSC candidate genes. MAGMA identified an association between cCSC and the *ARF* pathway. This pathway is involved in ribosomal biogenesis, and activation of the pathway leads to termination of ribosomal RNA production and cell cycle arrest.³⁵ In addition, PrediXcan showed decreased expression of *TNFRSF10A* and up-regulation of *KCNT2*. Involvement of *TNFRSF10A* in cCSC was suggested previously in our candidate gene study of the main AMD loci. We showed a protective effect for a *TNFRSF10A* variant, which was also observed in this study (OR, 0.74 [95% CI, 0.64-0.85]; $P = 1.47 \times 10^{-5}$; data not shown) and the largest AMD GWAS (OR, 0.90; $P = 4.5 \times 10^{-11}$).^{13,36} The *KCNT2* gene encodes a potassium sodium-activated outward rectifier channel (*KCNT2*) with an unknown function. The *ARF* pathway and *TNFRSF10A* and *KCNT2* genes are interesting candidates for cCSC, but their exact role in the eye remains to be elucidated.

Limitations

To our knowledge, this study includes the largest reported cohort of European patients with cCSC; however, even larger sample sizes are necessary to find associations with rare variants or variants with a low effect size. Additionally, in this study, only European individuals were included, and therefore genetic associations in other races/ethnicities still remain to be discovered. Replication of this study by centers in- or outside of Europe will be necessary to increase sample size, determine population specific associations, and replicate the suggestive signals observed in this study.

The use of PrediXcan on the GTEx data allows for prediction of expression levels in 44 tissues, excluding the eye. Although we only regarded those genes that showed differential expression in at least 50% of tissues they were expressed in, this does not necessarily mean that this also applies to expression in the retina. Likewise, retina-specific genes might have been missed due to tissue-specific expression. A retina-specific eQTL database would be necessary to determine these associations, but to date, such a database has not been available.

Conclusions

In this study, we describe a GWAS for cCSC and confirmed the association of genetic variants in *CFH* with the disease. Additionally, we identified *CFHR1*, *CFHR4*, *CD46*, *KCNT2*, and *TNFRSF10A* as cCSC candidate genes because of their genetic associations and predicted altered expression. With *CFH*, *CFHR1*, *CFHR4*, and *CD46* being important regulators of the complement cascade, this study strengthens the involvement of the complement system in cCSC. Further work on the expression of the proteins encoded by these genes is warranted. Additionally, the use of next-generation sequencing techniques, such as exome sequencing, will enable the identification of (rare) coding variants influencing protein function of these genes and might provide more insight in the causative mechanisms of cCSC.

Footnotes and Financial Disclosures

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Author Contributions:

Ms Schellevis had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. Drs van Dijk, Breukink, Altay, Boon, and de Jong contributed equally to the article. Drs van Dijk, Breukink, and Altay shared the second authorship, while Boon and de Jong share the last authorship position.

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Drafting of the manuscript: Schellevis, de Jong, van Dijk, Keunen, den Hollander, Boon.

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Supervision: Kiemeney, Keunen, Hoyng, den Hollander, Boon, de Jong.

Conflict of Interest Disclosures

All authors have completed and submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Dr den Hollander reports grants from Macula Vision Research Foundation during the conduct of the study, and personal fees from Ionis Pharmaceuticals outside the submitted work. Dr Fauser reports receiving support from Roche Pharmaceuticals outside the submitted work. No other disclosures were reported.

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Additional Information

The Nijmegen Biomedical Study is a population-based survey conducted at the Department for Health Evidence, and the Department of Laboratory Medicine of the Radboud university medical center. Principal investigators of the Nijmegen Biomedical Study are L.A.L.M. Kiemeney, A.L.M. Verbeek, D.W. Swinkels and B. Franke.

Supplementary Files

eMethods

DNA extraction and study participants

Genomic DNA isolation was performed from blood using a Hamilton Microlab STAR autoloader system with an integrated Chemagen MSM I separation module (Hamilton Robotics GmbH, Martinsried, Germany). DNA isolation was performed using the Chemagic DNA blood kit special (PerkinElmer) according to the manufacturer's instructions. An integrated barcode system was used throughout the process to track all samples and isolated DNA fractions. Following DNA isolation, concentrations of DNA fractions were determined (Hamilton Microlab Starlet robot with integrated Tecan Infinite 200 Pro reader).

In the Nijmegen Biomedical Study 21,756 age- and gender-stratified randomly selected inhabitants of the municipality of Nijmegen received an invitation to complete a postal questionnaire on, e.g., lifestyle and medical history, and to donate an 8.5 ml blood sample in a serum separator tube and a 10 ml EDTA blood sample. The response to the questionnaire was 43% (n=9,350), and 69% (n=6,468) of the responders donated blood samples

Quality Control and Filtering

The 3,654 NBS controls were genotyped with the HumanOmniExpress-12 (n=1034) or Infinium OmniExpress-24 (n=2620) bead chips. cCSC patients (n=547) were genotyped in three separate batches. Using the HumanOmniExpress-12 genotyping of 114 cCSC samples was performed by the Illumina facility of the Department of Genetics, University Medical Center Utrecht, The Netherlands, under supervision of B.P.C. Koeleman in May 2013 (cCSC1). To expand the study, two additional batches of cCSC patients were genotyped at the Genome Analysis Center, Helmholtz Zentrum München, Germany with the Infinium OmniExpress-24 bead chip, 96 samples in December 2014 (cCSC2) and 336 samples in August 2017 (cCSC3).

Quality control steps were applied to the separate batches using PLINK (v1.90b4.1, <https://www.cog-genomics.org/plink/1.9/>).^{1,2} Samples with a call rate of <97% or discordant reported gender were removed (71 controls and 7 cCSC patients). In each batch, SNPs that had genotype call rates <98% and/or showed deviations from Hardy-Weinberg equilibrium ($P < 10^{-6}$) were excluded and only variants with a minor allele frequency (MAF) >1% were retained. The cCSC3 data was lifted from GRCh38 to genome build GRCh37, to match the other batches (<http://genome.ucsc.edu/cgi-bin/hgLiftOver>).³ Only variants common to all datasets were kept and the data of the various batches was merged into one dataset. Strand inconsistencies were identified between batches with a flipscan implemented in PLINK and variants were flipped until concordance was reached in all datasets. Only variants with a call rate >98% in the full dataset were preserved, leaving 589,487 autosomal and 13,282 X-chromosomal variants that could be used for down-stream analysis.

Population stratification and cryptic relatedness

In order to assess population stratification, the dataset was merged with the Hapmap3 data containing individuals of known genetic ancestry.⁴ Data was pruned with a window size of 50kb, step size of 5 and variance inflation factor of 2 and

a principal component analysis (PCA) was performed with PLINK. Only individuals of European ancestry were retained for further analysis (Excluded 2 controls and 13 cCSC patients). Cryptic relatedness within the dataset was analyzed with KING (v2.0).⁵ A kinship coefficient threshold of <0.0884 was used to remove duplicates and individuals with a first or second degree relationship from the dataset (4 controls and 5 cCSC patients). Plots of the PCA and KING analyses were made with R (v3.2.0, <https://cran.r-project.org/>). In total, 521 cCSC patients and 3,577 controls remained for downstream analysis.

Imputation

After quality control a total of 589,487 autosomal and 13,282 X-chromosomal variants were used to impute the dataset. Autosomal genotype data was phased using Eagle (v2.3),⁶ while the X chromosome was phased with ShapeIT (v2. r790).⁷ After phasing, the data was imputed with the Haplotype Reference Consortium (HRC) release 1.1.2016 using the Michigan Imputation server (<https://imputationserver.sph.umich.edu>).^{8,9} SNPs were filtered on an imputation quality score of $R^2 > 0.3$ for common variants (MAF $>5\%$) and a $R^2 > 0.8$ for low frequency variants (MAF $<5\%$).

Pathway and PrediXcan analysis

In both MAGMA and VEGAS, first gene-based P-values were calculated which were used as input for the competitive gene-set analysis on predefined gene-sets. MAGMA gene-sets were downloaded from the molecular signatures database v6.1 (MSigDB, <http://software.broadinstitute.org/gsea/msigdb>, C2: curated gene-sets, canonical pathways) and 100,000 permutations were performed to apply empirical multiple testing correction.¹⁰ In VEGASv2 permutations ranging from 1,000 to 1,000,000 were performed depending on the p-value, and the provided biosystems gene-pathway annotation file was used, which includes genes from gene ontology, curated gene-sets from MSigDB, PANTHER, and pathway commons.¹¹

119

Transcriptome prediction models of the GTEx (V6) data containing 44 tissues, based on the 1000 genomes files were downloaded from PredictDB (<http://predictdb.hakymilab.org/>).¹² As PrediXcan does not allow for covariate implementation, the covariates gender, PC1 and PC2 were regressed out on the phenotype using the `lm` and `resid` function in R (`fit=lm(Disease~Sex+PC1+PC2)`, `resid(fit)`), and the resulting residuals were used as input for the analysis as recommended by PrediXcan in its google forum discussions.¹³ Because the resulting residuals were continuous, the linear regression option was used to perform the association analysis.

Power calculation

Power analysis was performed implementing the effective sample size calculation for alleles [$n_{\text{eff}} = 4 / (1/n_{\text{cases}} + 1/n_{\text{controls}})$]¹⁴ in an altered version of the `pwr` (v.1.2.1) R package provided by Dr. L.G. Fritsche (<https://github.com/ilarsf/gwasTools>). Power calculation input parameters for the current study were 521 cases and 3,577 controls, $\alpha = 5 \times 10^{-8}$ and $\alpha = 1 \times 10^{-6}$. Top variants were plotted according to their odds ratio (OR) and their MAFs in controls.

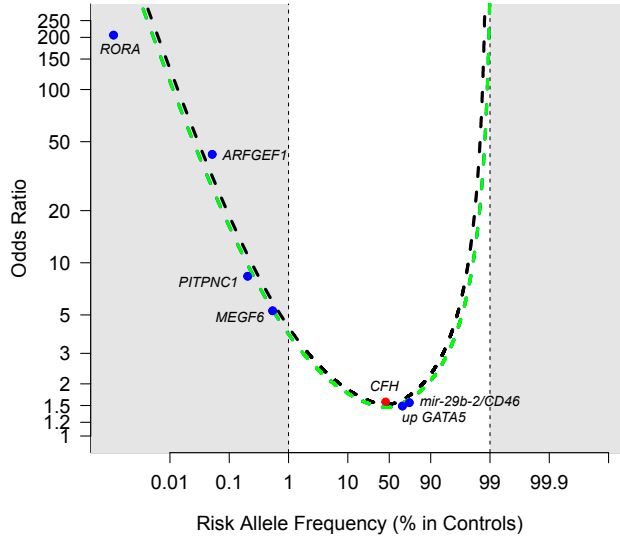


Figure 1. Power of the cCSC GWAS. The black line indicates the power of the current study at genome-wide significance ($P=5 \times 10^{-8}$), whereas the green line indicates power to detect suggestive signals ($P=1 \times 10^{-6}$). The lead variant in CFH is depicted in red, and suggestive signals are depicted in blue.

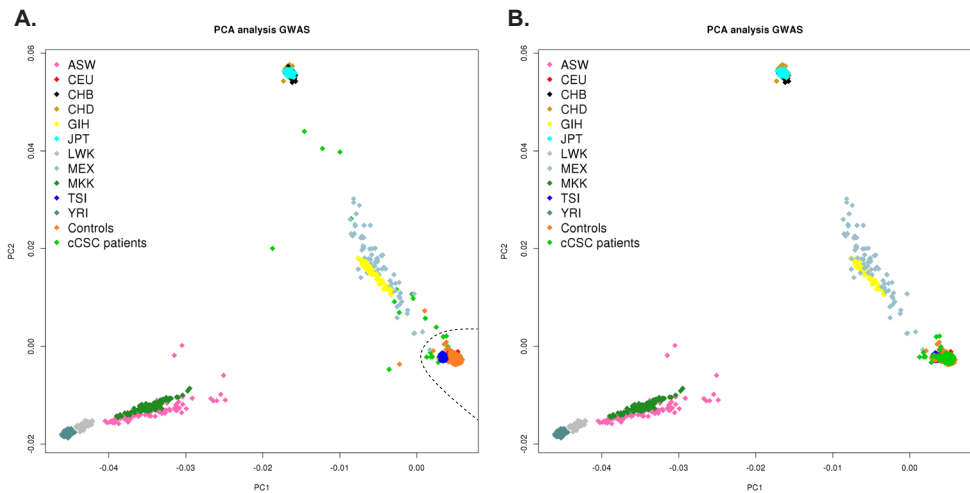


Figure 2. Principal component analysis (PCA) of the cCSC patients and controls. Graphical representation of the first 2 principal components of the PCA analysis, before (A) and after (B) filtering based on European ancestry. Various colours indicate the Hapmap3 samples according to the legend, orange depicts the controls and light green depicts the cCSC patients. Dotted line in A depicts cut-off used to exclude samples.

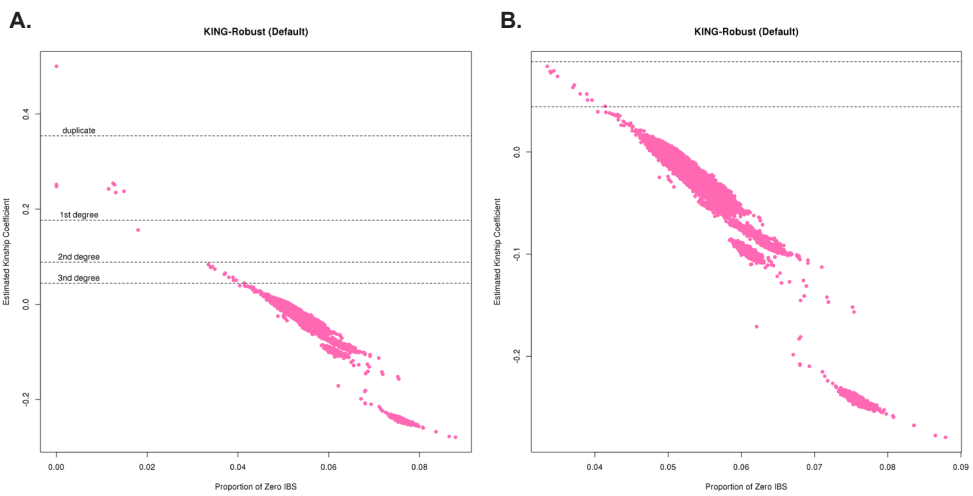


Figure 3. Kinship analysis of cCSC patients and controls. Kinship analysis was performed with KING, and estimated kinship coefficients of individuals in the study were calculated (A). Individuals with a kinship > 0.0884 were removed from the analysis (B).

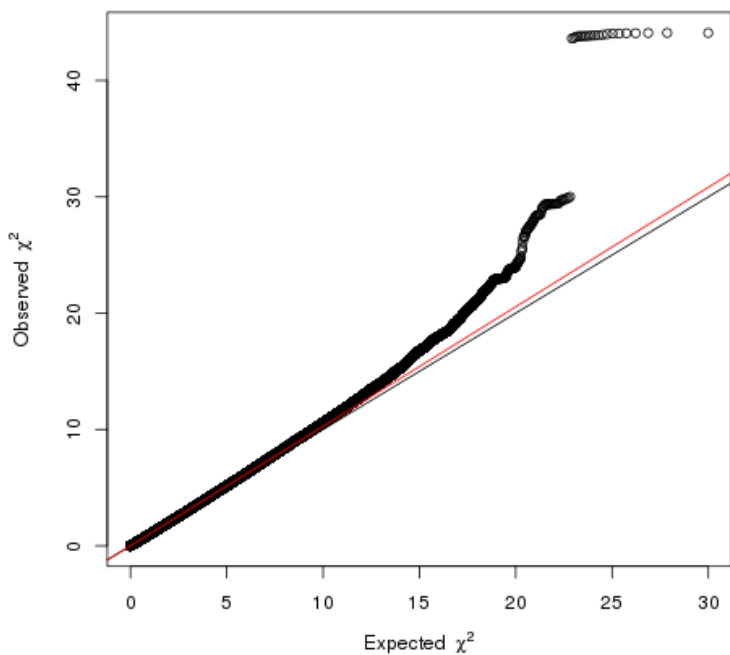


Figure 5. Quantile-quantile plot of the cCSC GWAS. Each round represents a genetic variant with its expected and observed chi-square statistics. The black line indicates the predicted ratio between observed and expected chi-square statistics. The red line indicates deviations due to inflation $\lambda=1.03$.

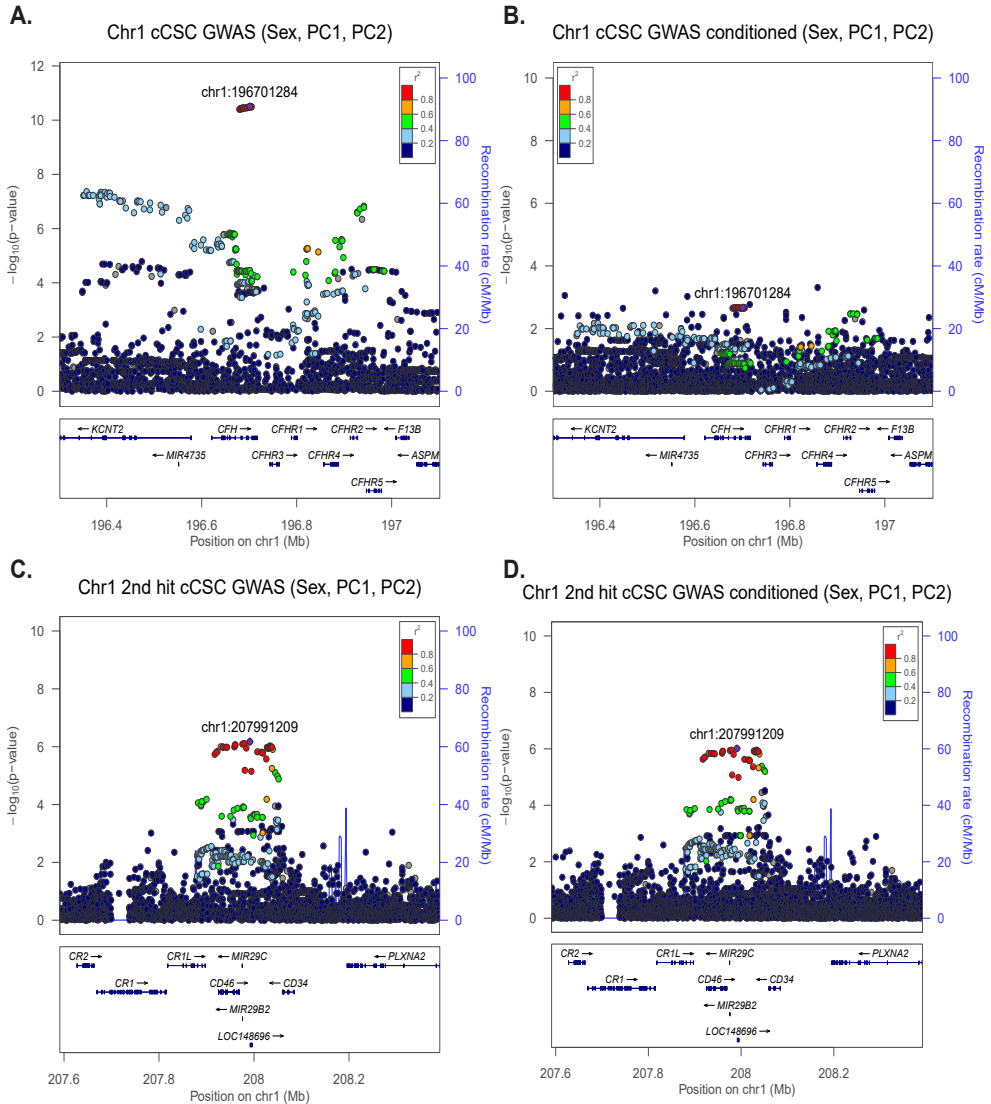


Figure 4. Conditioned analysis. Conditioned analysis of the lead variant on chromosome 1 was performed to identify independent signals on chromosome 1. No independent signal was found in the *CFH* gene (B), the additional signal in the *CD46* gene was independent of the lead variant (D).

eMethods References

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eTable 1. Demographic characteristics of the GWAS cohort

	Discovery cohort		
	n	Median age in years (IQR)	Males (%)
cCSC patients			
Nijmegen	307	51 (44-59)	242 (79%)
Cologne	71	51 (43-59)	55 (78%)
Leiden	143	51 (44-59)	123 (86%)
Total	521	51 (44-59)	420 (81%)
Controls			
NBS	3577	52 (37-71)	1630 (46%)
P-value cCSC patients vs Controls		0.192	1.78E-50

cCSC, chronic central serous chorioretinopathy; GWAS, genome-wide association study; NBS, Nijmegen Biomedical Study; Age difference between cCSC patients and controls was compared with a Mann-Whitney U test, and gender differences were compared with a chi-square test.

Table 2. Genome-wide significant hits in the *CFH* gene

SNP	Chr	bp (hg19)	Major/Minor allele	Complete Cohort				Cohort 50 years and younger			
				MAF cCSC (n=521)	MAF controls (n=3577)	P-value	OR (95% CI)	MAF cCSC (n=254)	MAF controls (n=1721)	P-value	OR (95% CI)
rs1329428	1	196702810	C/T	0.543	0.434	3.12E-11	1.57 (1.38-1.80)	0.583	0.433	1.23E-10	1.89 (1.55-2.31)
rs7540032	1	196701284	C/T	0.543	0.434	3.12E-11	1.57 (1.38-1.80)	0.583	0.433	1.23E-10	1.89 (1.55-2.31)
rs10922108	1	196701473	A/T	0.543	0.434	3.13E-11	1.57 (1.38-1.80)	0.583	0.433	1.23E-10	1.89 (1.55-2.31)
rs7514261	1	196700914	G/A	0.543	0.434	3.16E-11	1.57 (1.38-1.80)	0.583	0.434	1.24E-10	1.89 (1.55-2.31)
rs10801559	1	196704204	G/A	0.543	0.434	3.17E-11	1.57 (1.38-1.80)	0.583	0.433	1.24E-10	1.89 (1.55-2.31)
rs1329427	1	196704559	C/T	0.543	0.434	3.22E-11	1.57 (1.38-1.80)	0.583	0.433	1.26E-10	1.89 (1.55-2.31)
rs10922109	1	196704632	C/A	0.543	0.434	3.23E-11	1.57 (1.38-1.80)	0.583	0.433	1.27E-10	1.89 (1.55-2.31)
rs10801558	1	196699044	T/G	0.543	0.434	3.31E-11	1.57 (1.37-1.80)	0.583	0.434	1.32E-10	1.89 (1.55-2.31)
rs1410996	1	196696933	G/A	0.543	0.434	3.39E-11	1.57 (1.37-1.80)	0.583	0.434	1.37E-10	1.89 (1.55-2.31)
rs3766405	1	196695161	C/T	0.543	0.434	3.47E-11	1.57 (1.37-1.80)	0.583	0.434	1.37E-10	1.89 (1.55-2.31)
rs10922106	1	196691464	A/G	0.543	0.434	3.51E-11	1.57 (1.37-1.80)	0.583	0.433	1.11E-10	1.9 (1.56-2.32)
rs10922105	1	196690250	A/C	0.543	0.434	3.53E-11	1.57 (1.37-1.80)	0.583	0.433	1.12E-10	1.9 (1.56-2.31)
rs10922104	1	196687730	A/G	0.543	0.434	3.58E-11	1.57 (1.37-1.80)	0.583	0.434	1.13E-10	1.9 (1.56-2.31)
rs10465586	1	196687329	A/T	0.543	0.434	3.59E-11	1.57 (1.37-1.80)	0.583	0.434	1.13E-10	1.9 (1.56-2.31)
rs6688272	1	196684392	G/T	0.543	0.434	3.61E-11	1.57 (1.37-1.80)	0.583	0.434	1.14E-10	1.9 (1.56-2.31)
rs3753395	1	196686652	A/T	0.543	0.434	3.61E-11	1.57 (1.37-1.80)	0.583	0.434	1.14E-10	1.9 (1.56-2.31)
rs6677089	1	196684313	A/C	0.543	0.434	3.63E-11	1.57 (1.37-1.80)	0.583	0.434	1.15E-10	1.9 (1.56-2.31)
rs7535263	1	196682346	G/A	0.543	0.434	3.73E-11	1.57 (1.37-1.80)	0.583	0.434	1.29E-10	1.89 (1.55-2.31)
rs2274700	1	196682947	G/A	0.543	0.434	3.91E-11	1.57 (1.37-1.80)	0.583	0.434	1.28E-10	1.9 (1.55-2.31)
rs10737680	1	196679455	A/C	0.543	0.434	3.99E-11	1.57 (1.37-1.80)	0.582	0.434	1.31E-10	1.89 (1.55-2.31)

Table 3. Gene contents of the significant MAGMA gene-sets.

ALPHA = 0.05

TOT_SETS = 1329

SET1_SET = BIOCARTE_ARF_PATHWAY

SET1_NGENES = 17

SET1_P = 6.49E-06

SET1_P_CORR = 0.00865

SET1	GENE	CHR	START	STOP	NSNPS	NPARAM	N	ZSTAT	P
SET1	POLR1A	2	86253451	86333278	312	40	4098	1.3454	0.089243
SET1	POLR1B	2	113298676	113334727	168	28	4098	1.582	0.05682
SET1	PIK3CA	3	178866311	178952500	302	45	4098	2.6021	0.0046327
SET1	PIK3R1	5	67511584	67597649	387	91	4098	2.5764	0.0049915
SET1	POLR1C	6	43484777	43497114	36	13	4098	1.5684	0.058395
SET1	RAC1	7	6414126	6443598	172	32	4098	0.45552	0.32437
SET1	TWIST1	7	19039315	19157295	421	58	4098	3.0537	0.0011301
SET1	PIK3CG	7	106505723	106547592	116	35	4098	1.7093	0.0437
SET1	MYC	8	128748315	128753680	22	10	4098	0.72783	0.23336
SET1	CDKN2A	9	21967751	21994490	61	19	4098	0.43356	0.3323
SET1	ABL1	9	133588266	133763062	673	87	4098	1.747	0.040323
SET1	MDM2	12	69201952	69239324	123	28	4098	-0.94153	0.82678
SET1	POLR1D	13	28194880	28241559	190	42	4098	0.73589	0.2309
SET1	RBI	13	48877883	49056026	534	45	4098	0.85083	0.19743
SET1	TP53	17	7571720	7590868	81	25	4098	-0.54942	0.70864
SET1	TBX2	17	59477257	59486827	31	7	4098	0.29473	0.3841
SET1	E2F1	20	32263292	32274210	25	15	4098	1.4725	0.070444

SET2_SET = REACTOME_REGULATION_OF_COMPLEMENT_CASCADE

SET2_NGENES = 13

SET2_P = 5.49E-07

SET2_P_CORR = 0.00089

SET2	GENE	CHR	START	STOP	NSNPS	NPARAM	N	ZSTAT	P
SET2	CFH	1	196621008	196716634	383	42	4098	6.0264	8.38E-10
SET2	CFHR3	1	196743925	196763203	28	7	4098	2.2713	0.011565
SET2	C4BPB	1	207262210	207273338	38	15	4098	1.8251	0.03399
SET2	C4BPA	1	207277583	207318317	154	33	4098	0.97315	0.16524
SET2	CD55	1	207494817	207534311	131	22	4098	1.6713	0.047326
SET2	CR1	1	207669473	207815110	416	59	4098	1.0065	0.15709
SET2	CD46	1	207925383	207968861	138	23	4098	4.4686	3.94E-06
SET2	PROS1	3	93591881	93692934	168	55	4098	-0.85506	0.80374
SET2	CFI	4	110661848	110723381	245	33	4098	1.5119	0.065281
SET2	C2	6	31865562	31913449	211	60	4098	1.5594	0.059454
SET2	CFB	6	31913721	31919861	34	18	4098	2.6371	0.0041807
SET2	C4A	6	31949834	31970457	2	1	4098	-2.2974	0.9892
SET2	C3	19	6677846	6720662	202	32	4098	1.7533	0.039777

SET3_SET = REACTOME_COMPLEMENT_CASCADE

SET3_NGENES = 29

SET3_P = 5.56E-06

SET3_P_CORR = 0.00737

SET3	GENE	CHR	START	STOP	NSNPS	NPARAM	N	ZSTAT	P
SET3	MASP2	1	11086580	11107296	66	10	4098	1.0515	0.14651
SET3	C1QA	1	22963118	22966175	8	4	4098	-1.1141	0.86738

SET3	C1QC	1	22969969	22974603	25	6	4098	-0.26136	0.60309
SET3	C1QB	1	22979682	22988130	42	14	4098	-0.90879	0.81827
SET3	C8A	1	57320443	57383894	315	49	4098	0.31115	0.37784
SET3	C8B	1	57394883	57433213	182	45	4098	0.59985	0.2743
SET3	CRP	1	159682079	159684396	9	5	4098	-0.8472	0.80156
SET3	CFH	1	196621008	196716634	383	42	4098	6.0264	8.38E-10
SET3	CFHR3	1	196743925	196763203	28	7	4098	2.2713	0.011565
SET3	C4BPB	1	207262210	207273338	38	15	4098	1.8251	0.03399
SET3	C4BPA	1	207277583	207318317	154	33	4098	0.97315	0.16524
SET3	CD55	1	207494817	207534311	131	22	4098	1.6713	0.047326
SET3	CR1	1	207669473	207815110	416	59	4098	1.0065	0.15709
SET3	CD46	1	207925383	207968861	138	23	4098	4.4686	3.94E-06
SET3	PROS1	3	93591881	93692934	168	55	4098	-0.85506	0.80374
SET3	MASP1	3	186933873	187009810	363	87	4098	1.4588	0.07231
SET3	CFI	4	110661848	110723381	245	33	4098	1.5119	0.065281
SET3	C9	5	39284377	39364655	461	50	4098	1.5884	0.056098
SET3	C7	5	40909599	40983041	427	59	4098	0.18546	0.42644
SET3	C6	5	41142248	41261588	573	82	4098	0.18287	0.42745
SET3	C2	6	31865562	31913449	211	60	4098	1.5594	0.059454
SET3	CFB	6	31913721	31919861	34	18	4098	2.6371	0.0041807
SET3	C4A	6	31949834	31970457	2	1	4098	-2.2974	0.9892
SET3	C5	9	123714613	123812554	314	43	4098	1.5377	0.062059
SET3	C8G	9	139839333	139841426	15	4	4098	0.18851	0.42524
SET3	MBL2	10	54525140	54532578	52	16	4098	0.42175	0.3366
SET3	C15	12	7167819	7178336	29	9	4098	-1.1184	0.86831
SET3	CFD	19	859665	863624	9	4	4098	1.4475	0.073874
SET3	C3	19	6677846	6720662	202	32	4098	1.7533	0.039777

eTable 4. Gene contents of the VEGAS2 most significant gene-sets

Pathway	nGenes		nSamples	ObservedP	empiricalP	Genes
	Mapped	Used				
1269250 R-HSA-977606 Regulation of Complement cascade	22	13	1000000	1.03E-07	1.60E-05	C8B, CFHR3, CD46, PROS1, CFI, C9, C6, C4B, 2, C5, C8G, CD59, VTN, C3
1269241 R-HSA-166658 Complement cascade	36	22	500000	6.09E-07	4.20E-05	MASP2, C1QB, FCN3, C8B, CRR, CFHR3, CD46, PROS1, MASP1, CFI, C9, C6, C4B, 2, C5, FCN1, C8G, MBL2, CD59, C1R, VTN, CFD, C3

eTable 5. PrediXcan results of genes significant after Bonferroni correction for multiple testing.

Tissue	Gene	Gene	T	P-value	OR (95% CI)
Artery_Aorta	ENSG00000144031	ANKRD53	-4.725	2.38E-06	0.84 (0.79-0.91)
Stomach	ENSG00000144031	ANKRD53	-2.698	7.00E-03	0.91 (0.85-0.98)
Thyroid	ENSG00000144031	ANKRD53	-2.593	9.56E-03	0.89 (0.81-0.97)
Artery_Tibial	ENSG00000144031	ANKRD53	1.909	5.64E-02	1.07 (1-1.14)
Brain_Nucleus_accumbens_basal_ganglia	ENSG00000144031	ANKRD53	1.193	2.33E-01	1.01 (0.99-1.03)
Brain_Cortex	ENSG00000144031	ANKRD53	-0.471	6.38E-01	0.99 (0.96-1.03)
Skin_Not_Sun_Exposed_Suprapubic	ENSG00000144031	ANKRD53	-0.449	6.53E-01	0.99 (0.94-1.04)
Anterior_cingulate_cortex	ENSG00000144031	ANKRD53	-0.437	6.62E-01	0.98 (0.91-1.07)
Colon_Transverse	ENSG00000144031	ANKRD53	-0.042	9.66E-01	0.98 (0.32-2.96)
Esophagus_Muscularis	ENSG00000117335	CD46	4.783	1.79E-06	1.05 (1.03-1.07)
Skin_Not_Sun_Exposed_Suprapubic	ENSG00000117335	CD46	4.736	2.26E-06	1.32 (1.18-1.48)
Heart_Atrial_Appendage	ENSG00000117335	CD46	4.724	2.39E-06	1.16 (1.09-1.24)
Colon_Sigmoid	ENSG00000117335	CD46	4.709	2.57E-06	1.07 (1.04-1.09)
Artery_Tibial	ENSG00000117335	CD46	4.573	4.96E-06	1.05 (1.03-1.07)

Pituitary	ENSG00000117335	CD46	4.567	5.10E-06	1.1 (1.05-1.14)
Brain_Cortex	ENSG00000117335	CD46	4.565	5.15E-06	1.05 (1.03-1.07)
Artery_Coronary	ENSG00000117335	CD46	4.530	6.05E-06	1.11 (1.06-1.16)
Brain_Caudate_basal_ganglia	ENSG00000117335	CD46	4.478	7.73E-06	1.07 (1.04-1.1)
Artery_Aorta	ENSG00000117335	CD46	4.438	9.34E-06	1.06 (1.03-1.09)
Nerve_Tibial	ENSG00000117335	CD46	4.212	2.58E-05	1.05 (1.03-1.07)
Heart_Left_Ventricle	ENSG00000117335	CD46	4.201	2.71E-05	1.12 (1.06-1.18)
Brain_Cerebellar_Hemisphere	ENSG00000117335	CD46	3.999	6.47E-05	1.05 (1.02-1.07)
Cells_EBV-transformed_lymphocytes	ENSG00000117335	CD46	3.920	9.00E-05	1.05 (1.03-1.08)
Adipose_Subcutaneous	ENSG00000117335	CD46	3.902	9.68E-05	1.06 (1.03-1.1)
Esophagus_Gastroesophageal_Junction	ENSG00000117335	CD46	3.577	3.52E-04	1.04 (1.02-1.06)
Brain_Putamen_basal_ganglia	ENSG00000117335	CD46	3.327	8.87E-04	1.05 (1.02-1.08)
Brain_Cerebellum	ENSG00000117335	CD46	3.290	1.01E-03	1.03 (1.01-1.05)
Ovary	ENSG00000117335	CD46	3.246	1.18E-03	1.08 (1.03-1.13)
Brain_Hippocampus	ENSG00000117335	CD46	3.206	1.36E-03	1.05 (1.02-1.08)
Small_Intestine_Terminal_Ileum	ENSG00000117335	CD46	2.751	5.97E-03	1.05 (1.01-1.09)
Adrenal_Gland	ENSG00000117335	CD46	1.582	1.14E-01	1.03 (0.99-1.06)
Anterior_cingulate_cortex	ENSG00000117335	CD46	0.713	4.76E-01	1.01 (0.99-1.02)
Testis	ENSG00000117335	CD46	0.423	6.73E-01	1.01 (0.98-1.03)
Cells_Transformed_fibroblasts	ENSG00000117335	CFH	0.133	8.94E-01	1 (0.94-1.07)
Adipose_Subcutaneous	ENSG00000000971	CFH	4.869	1.17E-06	1.09 (1.05-1.13)
Nerve_Tibial	ENSG00000000971	CFH	4.801	1.63E-06	1.1 (1.06-1.14)
Ovary	ENSG00000000971	CFH	3.032	2.44E-03	1.05 (1.02-1.09)
Artery_Tibial	ENSG00000000971	CFH	1.045	2.96E-01	1.03 (0.97-1.09)
Pituitary	ENSG00000000971	CFH	-0.376	7.07E-01	0.99 (0.96-1.03)
Hypothalamus	ENSG00000244414	CFHR1	-5.648	1.73E-08	0.78 (0.72-0.85)

Tissue	Gene	Gene	T	P-value	OR (95% CI)
Adipose_Subcutaneous	ENSG00000244414	CFHR1	5.637	1.85E-08	1.15 (1.1-1.21)
Liver	ENSG00000244414	CFHR1	-5.221	1.87E-07	0.95 (0.93-0.97)
Cells_Transformed_fibroblasts	ENSG00000244414	CFHR1	-4.831	1.41E-06	0.79 (0.71-0.87)
Ovary	ENSG00000244414	CFHR1	4.564	5.15E-06	1.11 (1.06-1.16)
Artery_Tibial	ENSG00000244414	CFHR1	4.355	1.37E-05	1.24 (1.13-1.37)
Esophagus_Gastroesophageal_Junction	ENSG00000244414	CFHR1	4.346	1.42E-05	1.1 (1.05-1.15)
Adipose_Visceral_Omentum	ENSG00000244414	CFHR1	-3.302	9.68E-04	0.86 (0.79-0.94)
Spleen	ENSG00000244414	CFHR1	-2.457	1.40E-02	0.96 (0.93-0.99)
Pancreas	ENSG00000244414	CFHR1	-1.014	3.11E-01	0.6 (0.23-1.6)
Nerve_Tibial	ENSG00000244414	CFHR1	0.533	5.94E-01	1.01 (0.98-1.04)
Lung	ENSG00000244414	CFHR1	0.058	9.54E-01	1 (0.92-1.09)
Liver	ENSG00000134365	CFHR4	-4.849	1.29E-06	0.88 (0.83-0.93)
Esophagus_Gastroesophageal_Junction	ENSG00000162687	KCNT2	5.247	1.62E-07	3.7 (2.27-6.04)
Nerve_Tibial	ENSG00000162687	KCNT2	5.035	4.99E-07	1.13 (1.08-1.18)
Artery_Tibial	ENSG00000162687	KCNT2	3.540	4.05E-04	1.11 (1.05-1.17)
Esophagus_Muscularis	ENSG00000162687	KCNT2	1.578	1.15E-01	1.02 (1-1.05)
Adipose_Subcutaneous	ENSG00000162687	KCNT2	0.892	3.72E-01	1.06 (0.93-1.2)
Ovary	ENSG00000120539	MASTL	-5.027	5.19E-07	0.93 (0.9-0.96)
Adipose_Visceral_Omentum	ENSG00000120539	MASTL	-1.927	5.40E-02	0.94 (0.89-1)
Vagina	ENSG00000120539	MASTL	-1.525	1.27E-01	0.98 (0.95-1.01)
Adipose_Subcutaneous	ENSG00000120539	MASTL	-1.228	2.20E-01	0.99 (0.97-1.01)
Heart_Atrial_Appendage	ENSG00000120539	MASTL	-1.141	2.54E-01	0.99 (0.96-1.01)
Brain_Cerebellar_Hemispheres	ENSG00000120539	MASTL	-1.106	2.69E-01	0.99 (0.97-1.01)
Brain_Putamen_basal_ganglia	ENSG00000120539	MASTL	-1.072	2.84E-01	0.97 (0.92-1.03)
Liver	ENSG00000120539	MASTL	-1.061	2.89E-01	0.98 (0.94-1.02)

Cells_Transformed_fibroblasts	ENSG00000120539	MASTL	-1.030	3.03E-01	0.99 (0.97-1.01)
Stomach	ENSG00000120539	MASTL	-1.001	3.17E-01	0.99 (0.96-1.01)
Adrenal_Gland	ENSG00000120539	MASTL	-0.961	3.37E-01	0.99 (0.96-1.01)
Artery_Coronary	ENSG00000120539	MASTL	-0.961	3.37E-01	0.98 (0.95-1.02)
Esophagus_Gastroesophageal_Junction	ENSG00000120539	MASTL	-0.950	3.42E-01	0.99 (0.97-1.01)
Heart_Left_Ventricle	ENSG00000120539	MASTL	-0.948	3.43E-01	0.98 (0.95-1.02)
Skin_Sun_Exposed_Lower_leg	ENSG00000120539	MASTL	-0.946	3.44E-01	0.99 (0.97-1.01)
Muscle_Skeletal	ENSG00000120539	MASTL	-0.940	3.47E-01	0.99 (0.97-1.01)
Colon_Transverse	ENSG00000120539	MASTL	-0.911	3.63E-01	0.99 (0.96-1.01)
Artery_Tibial	ENSG00000120539	MASTL	-0.909	3.63E-01	0.99 (0.97-1.01)
Lung	ENSG00000120539	MASTL	-0.881	3.78E-01	0.99 (0.97-1.01)
Esophagus_Mucosa	ENSG00000120539	MASTL	-0.877	3.80E-01	0.99 (0.97-1.01)
Artery_Aorta	ENSG00000120539	MASTL	-0.875	3.82E-01	0.99 (0.96-1.01)
Esophagus_Muscularis	ENSG00000120539	MASTL	-0.821	4.12E-01	0.99 (0.97-1.01)
Breast_Mammary_Tissue	ENSG00000120539	MASTL	-0.782	4.34E-01	0.99 (0.97-1.01)
Colon_Sigmoid	ENSG00000120539	MASTL	-0.759	4.48E-01	0.99 (0.96-1.02)
Thyroid	ENSG00000120539	MASTL	-0.737	4.61E-01	0.99 (0.97-1.01)
Spleen	ENSG00000120539	MASTL	-0.722	4.70E-01	0.99 (0.97-1.02)
Skin_Not_Sun_Exposed_Suprapubic	ENSG00000120539	MASTL	-0.672	5.02E-01	0.99 (0.97-1.01)
Brain_Caudate_basal_ganglia	ENSG00000120539	MASTL	-0.616	5.38E-01	0.99 (0.96-1.02)
Anterior_cingulate_cortex	ENSG00000120539	MASTL	-0.613	5.40E-01	1 (0.98-1.01)
Brain_Cerebellum	ENSG00000120539	MASTL	-0.574	5.66E-01	0.99 (0.97-1.02)
Nerve_Tibial	ENSG00000120539	MASTL	-0.557	5.77E-01	0.99 (0.98-1.01)
Uterus	ENSG00000120539	MASTL	-0.536	5.92E-01	0.99 (0.97-1.02)
Whole_blood_assoc	ENSG00000120539	MASTL	-0.512	6.09E-01	0.99 (0.96-1.02)
Pituitary	ENSG00000120539	MASTL	-0.498	6.18E-01	0.99 (0.97-1.02)

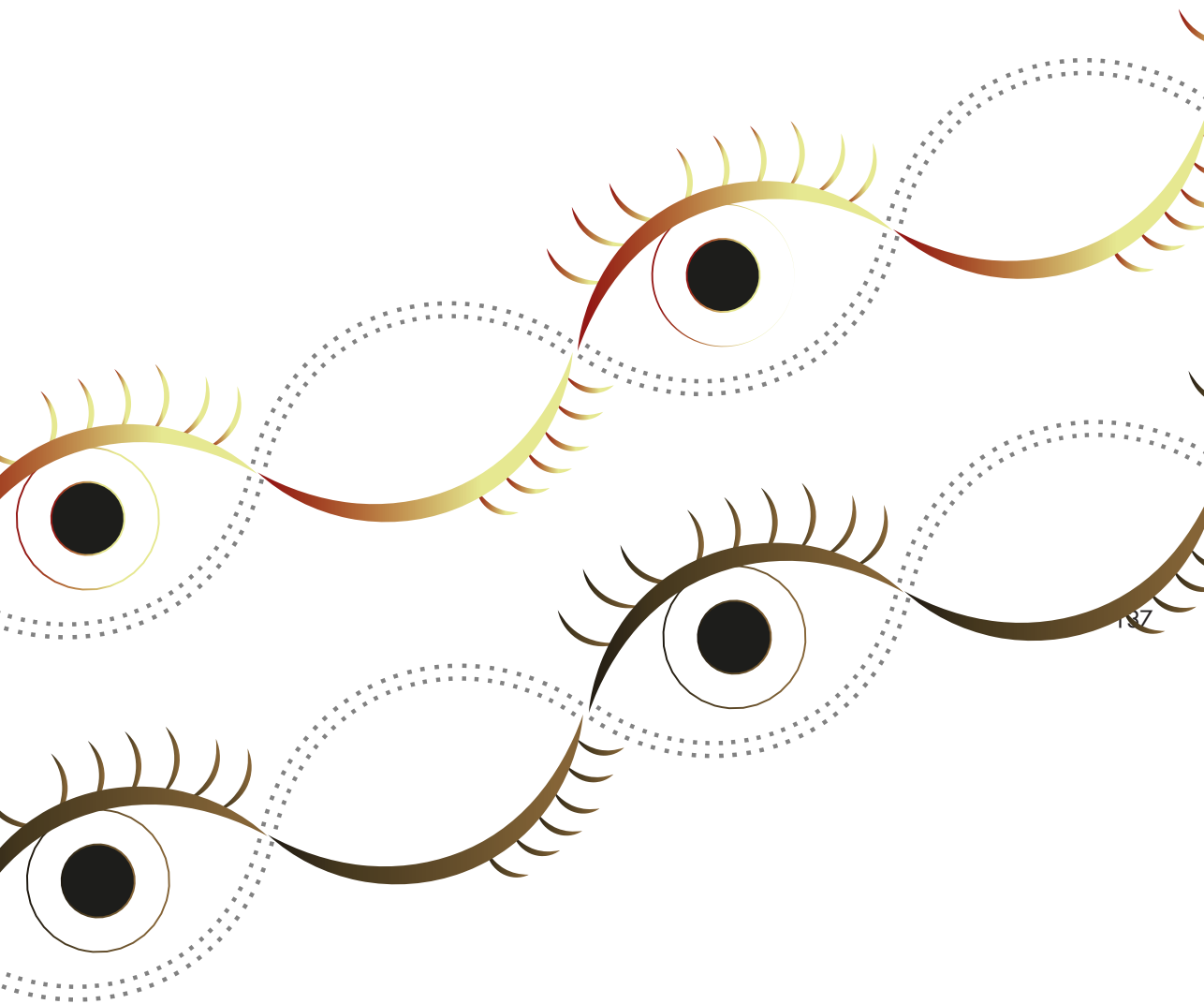
Tissue	Gene	Gene	T	P-value	OR (95% CI)
Pancreas	ENSG00000120539	MASTL	-0.490	6.24E-01	0.99 (0.97-1.02)
Small_Intestine_Terminal_Ileum	ENSG00000120539	MASTL	-0.474	6.36E-01	1 (0.98-1.01)
Cells_EBV-transformed_lymphocytes	ENSG00000120539	MASTL	-0.459	6.46E-01	0.99 (0.96-1.03)
Prostate	ENSG00000120539	MASTL	-0.281	7.79E-01	1 (0.98-1.02)
Hypothalamus	ENSG00000120539	MASTL	0.105	9.17E-01	1 (0.98-1.02)
Cells_Transformed_fibroblasts	ENSG00000104689	TNFRSF10A	-4.722	2.41E-06	0.96 (0.94-0.98)
Adrenal_Gland	ENSG00000104689	TNFRSF10A	-4.707	2.60E-06	0.95 (0.93-0.97)
Artery_Tibial	ENSG00000104689	TNFRSF10A	-4.282	1.90E-05	0.88 (0.83-0.93)
Esophagus_Gastroesophageal_Junction	ENSG00000104689	TNFRSF10A	-4.245	2.23E-05	0.62 (0.5-0.78)
Heart_Atrial_Appendage	ENSG00000104689	TNFRSF10A	-4.220	2.49E-05	0.87 (0.81-0.93)
Nerve_Tibial	ENSG00000104689	TNFRSF10A	-4.164	3.19E-05	0.74 (0.64-0.85)
Adipose_Subcutaneous	ENSG00000104689	TNFRSF10A	-4.051	5.19E-05	0.94 (0.92-0.97)
Artery_Aorta	ENSG00000104689	TNFRSF10A	-3.713	2.07E-04	0.97 (0.95-0.98)
Adipose_Visceral_Omentum	ENSG00000104689	TNFRSF10A	-3.466	5.33E-04	0.91 (0.87-0.96)
Esophagus_Muscularis	ENSG00000104689	TNFRSF10A	-3.446	5.76E-04	0.95 (0.92-0.98)
Lung	ENSG00000104689	TNFRSF10A	-3.214	1.32E-03	0.93 (0.9-0.97)
Colon_Sigmoid	ENSG00000104689	TNFRSF10A	-3.080	2.09E-03	0.96 (0.93-0.98)
Whole_blood_assoc	ENSG00000104689	TNFRSF10A	-2.770	5.63E-03	0.95 (0.91-0.98)
Testis	ENSG00000104689	TNFRSF10A	-2.045	4.09E-02	0.97 (0.94-1)
Liver	ENSG00000104689	TNFRSF10A	-2.037	4.18E-02	0.97 (0.94-1)
Brain_Cerebellar_Hemisphere	ENSG00000104689	TNFRSF10A	-1.358	1.75E-01	0.99 (0.97-1.01)
Anterior_cingulate_cortex	ENSG00000104689	TNFRSF10A	1.128	2.59E-01	1.02 (0.98-1.07)
Prostate	ENSG00000104689	TNFRSF10A	-0.951	3.41E-01	0.98 (0.95-1.02)

Red color indicates p-values that reached genome-wide significance $P < 0.05/17742 = 2.82 \times 10^{-6}$. Blue color indicates p-values < 0.05

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3.2 Exome sequencing identifies *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* as susceptibility genes for chronic central serous chorioretinopathy in females

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Abstract

Background: Chronic central serous chorioretinopathy (cCSC) is a multifactorial eye disease characterized by subretinal fluid accumulation that leads to vision loss. Clinically, cCSC is associated with stress, hypercortisolism and corticosteroid use, and is more frequent in males (80%) than in females (20%). Current genetic studies on cCSC have thus far focused on common variants, but familial occurrence of cCSC also suggests a role for rare variants in the disease susceptibility. Therefore, in this study, we performed exome sequencing of cCSC patients to elucidate the role of rare (protein-altering) variants in the disease.

Methods: Exome sequencing was performed on 269 cCSC patients and 1,586 controls. Data were processed according to the Genome-Analysis-Toolkit (GATK) best practices. Principal component analysis was performed to check for genetic ancestry and only unrelated subjects of European descent were retained. Burden, SKAT and SKAT-O tests were performed using 2 different grouping criteria. One group included protein-altering variants only, while the other contained synonymous and splice site variants as well. The gene-based analyses were performed using the SKAT R-package correcting for two principal components using two approaches; (1) on the entire cohort correcting for sex and (2) on males and females separately. Additionally, the gene-based associations of genes at previously reported cCSC loci were investigated.

Results: After filtering, the dataset contained 263 cCSC patients (208 males [79%]) and 1352 controls (671 males [50%]) carrying 197,915 protein-altering variants in 16,370 genes and 330,689 exonic variants in 18,173 genes. Analysis stratified by sex identified significant associations with the *PIGZ* ($P_{SKAT}=9.19 \times 10^{-7}$ & $P_{SKAT-O}=2.48 \times 10^{-6}$), *DUOX1* ($P_{SKAT}=1.03 \times 10^{-6}$), *RSAD1* ($P_{SKAT}=1.92 \times 10^{-7}$ & $P_{SKAT-O}=8.57 \times 10^{-8}$) and *LAMB3* ($P_{Burden}=1.40 \times 10^{-6}$ & $P_{SKAT-O}=1.14 \times 10^{-6}$) genes in female cCSC patients, after correction for multiple testing. The number of rare variant carriers in these genes was significantly higher in the female cCSC cohort compared to female controls (45,5% vs. 18.5%, $P=1.92 \times 10^{-6}$, $OR=3.67$ [95%CI=2.09-6.46]). No significant associations were identified in the entire cohort nor in the male patients.

Conclusions: In this exome study on cCSC patients, we have identified *PIGZ*, *DUOX1*, *RSAD1* and *LAMB3* as new susceptibility genes for cCSC in females. The sex-specific associations suggest a possible interaction between genetic factors and sex for cCSC, and indicate that rare (protein-altering) variants play a role in the etiology of cCSC in females.

Keywords (3-10): exome sequencing, central serous chorioretinopathy, CSC, Burden, SKAT, SKAT-O

Background

Central serous chorioretinopathy (CSC) is a vision-threatening eye disease characterized by serous fluid accumulation between the neuroretina and retinal pigment epithelium (RPE), supposedly secondary to a dysfunctional choroid and outer blood-retina barrier of the RPE.¹ The incidence of cCSC is estimated to be 1.7 in 100,000 in women and 9.9 in 100,000 in men with most cohorts confirming this ~80% male predominance.^{1,2} Two distinct forms of CSC have been described, distinguished by the duration of symptoms and extent of the RPE alterations. Acute CSC usually resolves spontaneously within ~3 months, whereas in chronic CSC (cCSC) fluid accumulation persists and more widespread RPE atrophy occurs.¹ The etiology of CSC is unknown, but associations with stress, hypertension and the use of corticosteroids have been described.¹ A genetic component for the disease was suggested because of familial occurrence of CSC,^{3,4} and so far a limited number of genetic studies have been performed for cCSC.

Candidate gene studies have described associations between cCSC and common single nucleotide polymorphisms (SNPs) in the *CFH*, *ARMS2*, *CDH5* and *NR3C2* genes,⁵⁻⁸ as well as copy number variations of the *C4B* gene.⁹ Additionally, the first genome-wide association study on cCSC confirmed the role of *CFH* in the disease.¹⁰ Other genes suggested to be involved in the genetics of cCSC include *CD46*, *CFHR1*, *CFHR4*, *KCNT2*, *TNFRS10A* and *VIPR2*.^{10,11} So far, only common variants have been associated with cCSC, however, familial clustering of cCSC has also hinted towards a possible role for rare variants in disease susceptibility.^{3,4}

Therefore, to elucidate the role of rare (protein-altering) variants in cCSC we performed exome sequencing on 269 cCSC patients. We performed single-variant association and gene-based tests, on the entire cohort and stratifying the analyses for sex. Additionally, we investigated the gene-based associations of genes at loci that have previously been associated with cCSC. Here, we report new genetic susceptibility loci and suggest a sex-specific genetic component for cCSC and further broaden our knowledge on the disease etiology.

Materials & Methods

Library preparation and sequencing

cCSC patients

In this study, 269 cCSC patients who visited the outpatient clinic of the Department of Ophthalmology of the Radboud university medical center, Nijmegen, the Netherlands were included. Patients were diagnosed with cCSC based on multimodal retinal imaging as described previously.^{6,10} DNA of cCSC patients, isolated from blood, was purified with the QIAamp DNA purification kit (Qiagen), according to manufacturer's instructions. 3µg of purified DNA was used as input for the SureSelect^{XT} target enrichment system for Illumina paired-end multiplexed sequencing (Version B4, August 2015, Agilent Technologies).

Library preparation was performed according to the manufacturer's instructions. Samples were hybridized with the SureSelect All Exon V4 (Agilent Technologies) capture library and post-hybridization addition of the index tags was performed on the

Dynabeads MyOne Streptavidin T1 beads (ThermoFisher Scientific). Midway and final quality checks were performed with TapeStation high sensitivity D1000 screentape (Agilent Technologies) and a Qubit dsDNA high sensitivity assay (ThermoFisher Scientific). A total of 8 samples was pooled for sequencing in one lane. Sequencing was performed at the Department of Genetics of Maastricht University Medical Center+, Maastricht, The Netherlands, using an Illumina HiSeq2000 with 2*100bp chemistry.

Controls

As population controls, 1586 parents that were part of a large cohort of intellectual disability trios were used;¹² no ophthalmologic imaging was available for these individuals. Control samples were sequenced with the Agilent SureSelect V4 enrichment kit in a diagnostic setting at the Beijing Genomics Institute (BGI) in Copenhagen between August 2013 and March 2015. Sequencing was performed on an Illumina HiSeq instrument with 101-bp paired-end reads.¹² The median coverage of the exomes of both cCSC patients and controls was 75X.

This study was performed according to the guidelines of the Declaration of Helsinki and was approved by the local ethics committee. Written informed consent was obtained for all subjects involved in the study.

Variant calling

Bam to FastQC extraction was performed for both controls and cCSC patient data with Picard-tools (v 1.90). Files of patients and controls were processed together to minimize batch effects.¹³ Duplicate reads were marked with Picard-tools and reads were aligned to the reference genome (GRCh37.p5 with alternate haplotypes excluded) using BWA-MEM (version v.0.7.12). Sequencing data was processed according to the Genome-Analysis-Toolkit (GATK) best practices (v3.8). Briefly, base recalibration was performed using the following known sites: dbsnp version 144, Mills and 1000G gold standard and 1000G phase1 high confidence insertion/deletions (INDELs). Variant calling was performed with haplotypcaller in g.vcf mode specified on the SureSelect V4 regions flanked with 200bp. Individual g.vcfs were merged into batches of 200 samples and joint genotyping on the whole cohort was performed per chromosome using genotypeGVCFs with the newQual option. Sites with a minimal average coverage of 15X in both the cCSC patients and control datasets were retained for downstream analyses.

Variant quality score recalibration (VQSR)

Variant recalibration was performed with GATK (v3.8) using the recommended settings.¹³ For SNPs, Hapmap3.3 and OmniExpress 2.5 chip sites were used as truth and training sets, while the 1000G phase1 data was used for training only. For INDELs, the Mills and 1000G gold dataset was used as truth and training sets. For both SNPs and INDELs “dbsnp v138 excluding sites after dbsnp v129 sites” were used to indicate known variants. The following arguments were used during the generation of the variant recalibration model for SNPs: Quality by Depth (QD), Mapping Quality (MQ), MQRankSum, ReadPosRankSum, Fisher strand (FS), Strand odds ratio (SOR), InbreedingCoeff, and MQCap = 60. For INDELs the MQ option was removed, as recommended. The variant recalibration model was applied to the dataset for SNPs

and INDELs separately and the data was filtered based on the 99.5 and 99 sensitivity tranche, respectively.

Population stratification, Cryptic relatedness & Sex-check

All variants overlapping with the Hapmap3 data were extracted and combined with the original Hapmap3 data containing individuals of various genetic ancestries. The merged dataset was pruned using PLINK (v1.90b3y), with a window size of 50kb, stepsize of 5kb and variant inflation factor of 2. Principal component analysis (PCA) was performed on the data to account for population admixture. The first 2 PCA components were plotted with R (v 3.2.0), and ancestry labels were added to the Hapmap data. After filtering only individuals from European descent were retained (Figure S1 in the Supplement).

To assess cryptic relatedness between individuals in the dataset, kinship coefficients were calculated with KING (v2.0).¹⁴ Proportion of zero identity by state (IBS) was plotted against the estimated kinship coefficient with R (v 3.2.0) (Figure S2 in the Supplement). Samples were removed that had a kinship coefficient indicative of being either first degree (0.177-0.354), second degree (0.177-0.0884) or third degree (0.0884-0.0442) relatives. Finally, reported sex was compared with the data available for the X chromosome using the sex-check option in PLINK, with a threshold of $F < 0.35$ for women and $F > 0.7$ for men.

Variant filtering

Variants located in low complexity regions of the genome were removed.¹⁵ Multiallelic variants were extracted with VCFtools (v0.1.13) and split using the splitMultiallelic and LeftAlignandTrimVariants option in GATK (v3.8). After splitting, biallelic and multiallelic SNPs and INDELs were combined and tested for Hardy-Weinberg equilibrium $P > 1 \times 10^{-8}$ and a minor allele count of 1. Variants from the adult-onset cancer genes captured in the American College of Medical Genetics and Genomics recommendations for incidental findings (*BRCA1*, *BRCA2*, *MLH1*, *MSH2*, *MSH6*, *PMS2*, *MUTYH*) were removed to reduce the risk of secondary findings.¹⁶ Remaining variants were annotated with TabAnno (<https://github.com/zhanxw/anno>). In order to perform the downstream gene analysis, two different groupfiles were made using a Python script (available on request). The protein-altering variant groupfile contained nonsynonymous, stop gain/loss, start gain/loss, codon gain/loss, frameshift and essential splice site variants, while the exonic groupfile included both protein-altering variants and synonymous, exon and variants in the complete splice site.

Statistical analysis

Data was converted to binary PLINK files and imported into the SKAT R-package (v. 1.2.1).^{17,18} As the SKAT package does not offer a function to analyze a large set of single variants, one large “dummy” geneset was generated containing all the variants in the input file. The association of each variant was calculated with a loop through the “dummy” geneset (Script File 1 in the Supplement). The SKAT R-package uses the efficient resampling method to also include extremely rare variants in the analysis.¹⁷ The SKAT null model adjusted for small sample size was performed for a binary phenotype correcting for two principal components and sex (for the entire cohort) with the resampling method “bootstrap.fast”. The generated file was also used to

calculate the variant weights based on minor allele frequency using the “Get_Logistic_Weights” function of the SKAT package (v. 1.2.1).^{17,18}

Next, a minor allele frequency (MAF) weighted Burden, SKAT, and SKAT-O analysis was performed with the “SKATBinary.SSD.All” option. These analyses were performed on either the entire cohort or the cohort stratified by sex using the protein-altering and exonic groupfiles. The default settings were used for kernel (linear.weighted), imputation methods (bestguess), number of resampling (2×10^6) and p-value calculation (hybrid). Only genes with at least two variants were retained and multiple testing correction was performed using Bonferroni correction. Plots were generated with the R-package CMplot. An example of the gene-based test pipeline is available in Script File 1 in the Supplement. Sanger sequencing was performed to confirm the presence of the variants in the top- and suggestive genes of the different gene-based tests in the cCSC patients ($P < 5 \times 10^{-5}$). Additionally, Burden, SKAT and SKAT-O p-values of previously associated cCSC genes and genes in their 100kb up- and downstream region were extracted from the dataset.

The number of rare variant carriers (MAF < 0.05) in the four identified cCSC susceptibility genes was counted per gene in female and male cCSC patients and controls. The carrier and non-carrier groups were compared between controls and cCSC patients using a Chi-square test or Fisher’s exact test when at least 25% of the cells had an expected count <5. Odds ratios for single variants in the susceptibility genes were calculated using the Firth corrected likelihood method in SAS (v9.4),¹⁹ when variants were significantly associated with cCSC, because the SKAT single variant analysis only provides P-values and no risk estimates. Haplotypes in the *LAMB3*

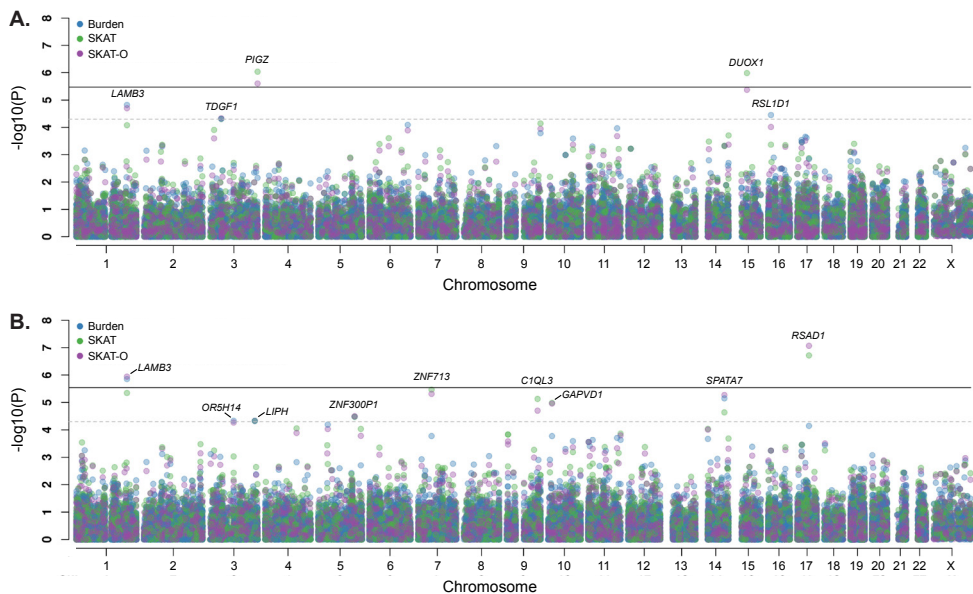


Figure 1: Manhattan plot of gene-based analysis in female cCSC patients. Burden (blue), SKAT (green) and SKAT-O (purple) association results are depicted for the female cohort including 55 cCSC patients vs. 681 controls, using the protein-altering (A) and exonic (B) group files. The dark horizontal line indicates the significance threshold after correction for multiple testing: $0.05/17,370 \text{ genes} = 2.88 \times 10^{-6}$ (A), $0.05/14,935 \text{ genes} = 3.35 \times 10^{-6}$ (B), while the dotted line indicates the suggestive threshold of $P = 5 \times 10^{-5}$.

gene were constructed with the haplo.stats R-package as described before.^{10,20}

Results

Quality control

In this study, 269 cCSC patients and 1586 controls were analyzed by exome sequencing. Among these samples 232 controls and six cCSC patients were not of European descent, and were therefore excluded from downstream analysis. Additionally, three control samples were removed due to cryptic relatedness (n=2) or uncertain sex based on the X-chromosome data (n=1). After individual quality control the dataset contained 263 cCSC patients (208 males [79%]) and 1352 controls (671 males [50%]). For these individuals, joint genotyping with GATK resulted in a dataset containing 746,264 SNPs and 45,048 INDELS. Of these 3,793 SNPs and 1,661 INDELS were multi-allelic, and were split. Additional quality control included HWE filtering ($P = 1 \times 10^{-8}$) and removal of the low complexity regions of the genome.¹⁵ Finally, a total of 579,602 variants remained for downstream analysis, of which 547,595 were SNPs and 32,007 were INDELS.

Unbiased single variant association and gene-based analyses

Single variant association analysis corrected for sex and two principal components did not identify any genome-wide significant hits in the entire datasets, nor in the datasets stratified by sex (Figure S3 in the Supplement). To increase power to detect associations, gene-based analyses using the Burden, SKAT and SKAT-O tests were performed. These analyses were carried out on the entire cohort and on male and female cohorts stratified by sex, testing either all protein-altering variants or using all exonic variants. The number of genes and variants included in the gene-based analyses is described in Table 1.

In the protein-altering variant analysis, genome-wide significant associations with the *PIGZ* ($P_{SKAT} = 9.19 \times 10^{-7}$ & $P_{SKAT-O} = 2.48 \times 10^{-6}$) and *DUOX1* ($P_{SKAT} = 1.03 \times 10^{-6}$) genes were identified in females (Table 2, Figure 1A). Additionally, suggestive associations in the protein-altering analysis ($P < 5 \times 10^{-5}$) were identified in the entire cohort, as well as the male and female cohort for *NOP14*, *RSPO2*, *LAMB3*, *TDGF1* and *RSL1D1* (Figure 1A, Table S2 and Figure S4 the Supplement). Gene-based analyses of the exonic variants identified genome-wide significant associations in the *RSAD1* ($P_{SKAT} = 1.92 \times 10^{-7}$ & $P_{SKAT-O} = 8.57 \times 10^{-8}$) and *LAMB3* ($P_{Burden} = 1.40 \times 10^{-6}$ & $P_{SKAT-O} = 1.14 \times 10^{-6}$) genes in the female cohort (Table 2, Figure 1B). Gene-based analysis of the exonic variant on the entire

Table 1. Number of variants and genes included in the gene-based analyses

	Protein-altering variants [#]			Exonic variants [#]		
	Variants (n)	Genes (n)	Bonferroni correction p-value threshold	Variants (n)	Genes (n)	Bonferroni correction p-value threshold
All individuals	197,915	16,370	3.05×10^{-6}	330,689	18,173	2.75×10^{-6}
Males	135,468	15,083	3.31×10^{-6}	232,667	17,348	2.88×10^{-6}
Females	123,898	14,935	3.35×10^{-6}	215,244	17,370	2.88×10^{-6}

[#]Only genes with more than 2 variants were retained and used for Bonferroni correction

cohort and the male cohort did not identify any genome-wide associations. Suggestive hits for the entire cohort and the stratified male and female cohorts were identified in *ALX1*, *NOP14*, *G3BP1*, *RNF144A*, *C5orf63*, *NOB1*, *LOC283332*, *ZNF713*, *SPATA7*, *C1QL3*, *GAPVD1*, *ZNF300P1*, *LIPH* and *OR5H14* (Figure 1B, Table S2 and Figure S4 in the Supplement).

Rare variant carrier frequency in PIGZ, DUOX1, LAMB3 and RSAD1 in cCSC patients and controls

Gene-based analyses identified four cCSC-associated susceptibility genes in the female cohort. An overview of all variants found in *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* in females is presented in Table S2 in the Supplement. The *PIGZ* gene-based association was explained by 14 rare (MAF <0.05) and 4 common (MAF >0.05) nonsynonymous variants. The number of rare variant carriers was higher in the female cCSC patient group compared to the female controls (Table 3, P=0.014, Odds ratio (OR)=3.67 [95% confidence interval (CI)=1.42-9.47]). The p.A124V variant in *PIGZ* was individually associated with cCSC in females (P=0.0003, OR=16.40 [95% CI= 3.64-73.94]) (Table S2 in the Supplement).

The *DUOX1* gene-based association was composed of 12 rare and 2 common nonsynonymous variants and 1 rare stop-gain variant. Of the rare variants, 3 were found in the cCSC group, whereas 11 were found in the control group. The number of carriers was significantly higher in the cCSC group compared to controls (Table 3, P= 0.039, OR=3.74 [95%CI=1.19-11.77]) The p.R925W variant was individually associated with cCSC (P=0.03, OR=62.69 [95%CI=1.52->999.99]), and was only found in the female cCSC group (Table S2 in the Supplement).

145

In the *LAMB3* gene, a total of 49 variants was observed in females. These variants included 31 nonsynonymous, 13 synonymous, 3 intronic variants near splice sites and 2 stop-gain variants. Three of the synonymous variants were common, while the remaining 46 variants were rare. In the control group 94 individuals (13.8%) carried at least one *LAMB3* rare variant, while 17 (30.9%) female cCSC patients carried one or more rare variants in *LAMB3* (Table 3, P=0.00056, OR=2.83 [95%CI=1.53-5.22]). Haplotype analysis identified a significant association with cCSC for a haplotype carrying 7 variants in *LAMB3* (p.S97S, p.N181D, p.V527M, p.Y588Y, p.S708S, p.N690S, p.A926D) which had a frequency of 7.3% in the cCSC patients and 1.6% in the control cohort (Table S3 in the Supplement, P=7.22x10⁻⁵, OR=6.60 [95%CI=2.61-16.67]).

Table 2. Significant gene-based association results for the female cCSC cohort using the Burden, SKAT and SKAT-O tests

Gene	Variants	Burden p-value	SKAT p-value	SKAT-O p-value
<i>PIGZ</i>	Protein-altering	6.99x10 ⁻³	9.19x10 ^{-7#}	2.48x10 ^{-6#}
<i>DUOX1</i>	Protein-altering	7.18x10 ⁻³	1.03x10 ^{-6#}	4.23x10 ⁻⁶
<i>RSAD1</i>	Exonic	7.14x10 ⁻⁵	1.92x10 ^{-7*}	8.57x10 ^{-8*}
<i>LAMB3</i>	Exonic	1.40x10 ^{-6*}	4.51x10 ⁻⁶	1.14x10 ^{-6*}

Significant after Bonferroni correction for 14,935 genes; P≤3.35x10⁻⁶

* Significant after Bonferroni correction for 17,370 genes; P≤2.88x10⁻⁶

Table 3. Female carriers of rare variants (MAF <0.05) in *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1*

Gene	Nr. rare variants in gene	Nr. carriers in controls (n _{total} =681)	Nr. carriers in cCSC patients (n _{total} =55)	P-value	OR (95% CI)
<i>PIGZ</i>	14	22 (3.2%)	6 (10.9%)	0.014	3.67 (1.42-9.47)
<i>DUOX1</i>	13	14 (2.1%)	4 (7.3%)	0.039	3.74 (1.19-11.77)
<i>LAMB3</i>	46	93 (13.8%)	17 (30.9%)	5.57x10 ⁻⁴	2.83 (1.53-5.22)
<i>RSAD1</i>	8	5 (0.7%)	3 (5.5%)	0.017	7.80 (1.81-33.55)
Variant in 1 or more genes		126 (18.5%)	25 (45.5%)	1.92x10 ⁻⁶	3.67 (2.09-6.46)

Nr. number; P-values are calculated with Chi-square test or Fisher's exact when at least 25% of the cells had an expected count <5.

A total of 11 variants was present in *RSAD1* in females, of these 9 were nonsynonymous (7 rare, 2 common) variants, 1 was a common synonymous variant, and 1 was a rare intronic variant. The number of rare variant carriers was higher among female cCSC patients, compared to the control group (Table 3, P= 0.017, OR=7.80 [95%CI=1.81-33.55]). Three variants were only observed in the patient group (p.P250L, p.V299I, p.R317Q).

In total, 18.5% of the female control individuals carried at least one rare variant in *PIGZ*, *DUOX1*, *LAMB3* or *RSAD1*, whereas this number was 45.5% in the female cCSC patients. The number of rare variant carriers in these genes was significantly higher in the female cCSC patients group compared the control group (Table 3, P=1.92x10⁻⁶, OR=3.67 [95%CI=2.09-6.46]). In males, no differences in the number of carriers of *PIGZ*, *DUOX1*, *LAMB3* or *RSAD1* rare variants were observed between controls and cCSC patients (Table S4 in the Supplement). Also, the total number of carriers of at least one variant in these 4 genes was not different in males (Table S4 in the Supplement).

Gene-based association of genes located in known cCSC loci

Genes at previously associated cCSC loci and genes within ±100kb of the top-associated variant were extracted from the dataset including the *ARMS2-HTRA1*, *C4-RCCX*, *CD46*, *CDH5*, *KCNT2-CFH-CFHR1/4*, *NR3C2*, *TNFRS10A* and *VIPR2* loci. In total, 56 genes were located in these regions, of which 39 genes contained at least 2 protein-altering variants and 44 genes had at least 2 variants in the exonic regions. The gene-based association results of these genes were extracted from the genome-wide analysis. However, no associations were identified between these genes and cCSC after correction for multiple testing (Table S5 in the Supplement, Threshold P<0.00114 [corrected for 44 genes in the exonic variant analysis] and P<0.00128 [corrected for 39 genes in the protein-altering variant analysis]).

Discussion

In this study, in order to increase our knowledge and understanding of the genetics of cCSC, we performed exome sequencing on a cohort of cCSC patients. Using a gene-based approach, we identified 4 new candidate genes for the disease in female cCSC patients specifically.

Due to a relative small sample size this study had limited power to identify single-variant associations, especially for rare variants, therefore, three different gene-based analyses with varying assumptions were performed. Previous studies have shown that the *CFH* gene carried both protective and risk-conferring variants,^{5,6} therefore, we used the SKAT test, which assumes variants within a gene can have both directions of effect. The Burden test on the other hand relies on the assumption that all variants in a gene have the same direction of effect, and the SKAT-O test combines both assumptions into one test.¹⁸ The gene-based analyses were performed using two different inclusion criteria. One analysis contained protein-altering variants only, since they are most likely to affect protein function. The other analysis contained all exonic variants, including synonymous (silent) variants and splice site variants, of which the effect on protein function is more difficult to predict. Nevertheless, such variants may alter gene expression levels and were therefore included in the analysis.²¹ In the female cohort we identified a significant burden of protein-altering variants in two genes (*PIGZ* and *DUOX1*) and a significant burden of exonic variants in two additional genes (*LAMB3* and *RSAD1*).

Nearly half of the females in the cCSC cohort carried at least one variant in the 4 identified cCSC susceptibility genes (45.5%), while this number was significantly lower in the female control cohort (18.5%, $P=1.92 \times 10^{-6}$). Carrying a rare variant in one of these genes increases the risk for developing cCSC 3.7-fold in females, while carrying a variant in *RSAD1* even increases risk 7.8-fold. Notably, a number of high-risk variants were identified in the gene-based tests, including the p.A124V variant in *PIGZ* ($P=0.0003$, OR=16.40 [95% CI= 3.64-73.94]) and the p.R925W variant in the *DUOX1* gene ($P=0.03$, OR=62.69 [95%CI=1.52->999.99]), and three variants (p.P250L, p.V299I, p.R317Q) in *RSAD1* were observed only in the patient group but not in the controls. The number of carriers of rare variants in the *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* genes was not different between male cCSC patients and controls (Table S4 in the Supplement, 15.9% vs. 20.4%, respectively, $P=0.146$), indicating that the observed associations with cCSC are sex-specific. Replication in an independent female cCSC cohort is recommended to substantiate these findings.

Interestingly, genome-wide significant associations were only observed in female cCSC patients but not in the males. The enrichment of rare variants in these genes in females is likely not due to population stratification, sequencing artefacts or the presence of related individuals, because stringent quality control was performed to avoid these confounders. It remains unclear why no associations in the male cCSC cohort were observed, despite accounting for the largest proportion of the cohort. One could hypothesize that cCSC is more genetically heterogeneous in males, and that a larger sample size is therefore necessary to identify associations in this group. The biological mechanism behind these sex-specific associations remains to be elucidated, but sex-biased gene expression or regulation could be one of the

contributing factors.^{22,23} Interactions between sex and genetics have been described for other complex traits such as hypertension, schizophrenia and juvenile idiopathic arthritis,²³⁻²⁵ and for cCSC the previously described *CDH5* association was observed in males only.⁷ In light of future clinical trials and genetic studies on cCSC, sex-stratification should be taken into account to further elucidate a possible interaction between sex, genetics and possibly treatment outcome.

For cCSC specifically, an interaction between genetic variants and sex hormone regulation and expression seems likely, since the association of cCSC with the stress axis, hypercortisolism and the use of corticosteroids has been described extensively, and pregnancy is known to increase the risk of CSC in females.^{1,26} An interaction with the steroid hormone system has already been described for *LAMB3* and *DUOX1*,^{27,28} and all four identified susceptibility genes are expressed in different tissues of the eye including the retina and RPE (Figure S5 in the Supplement).²⁹

The *LAMB3* gene encodes the beta chain of laminin-5, which increases the adhesion properties of RPE cells and promotes the epithelial appearance of these cells *in vitro*.³⁰ Laminin-5 shows upregulation in endometrial biopsies after treatment with progesterone and 17 β -estradiol,²⁷ suggesting a hormonal regulation of the gene. The dual oxidase 1 protein encoded by *DUOX1*, is a member of the NADPH oxidase family and generates hydrogen peroxide, an important reactive oxygen species (ROS). *DUOX1* is mainly expressed in thyroid and lung epithelial cells, however after serum starvation both rods and cones expressed *DUOX1* indicating that it plays a role in ROS response in these cells as well.³¹ Furthermore, the expression of *DUOX1* is stimulated by testosterone in skin keratinocytes, indicating a steroid hormone-dependent activation of this gene at least in these cells.²⁸ No steroid response has been described yet for the *PIGZ* and *RSAD1* genes and their general function is still obscure.^{32,33} In light of the cCSC phenotype, for all four genes the effect of the identified variants on the disease mechanisms needs to be determined. Additionally, future experiments may focus on determining if a steroid hormone response is also present in the retina, RPE and/or choroid for these genes, and if the identified rare variants might alter this response.

Conclusions

In this exome sequencing study on a cohort of cCSC patients and population controls, we identified *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* as new susceptibility genes for cCSC in females specifically. Our results suggest that the contribution of rare genetic risk variants is different between the sexes and that the cCSC disease mechanism might differ between males and females. Further genetic studies and clinical trials on cCSC may take a sex-interaction component for cCSC into account.

Footnotes and Financial Disclosures

Declarations:

This study was performed according to the guidelines of the Declaration of Helsinki and was approved by the local ethics committee. Written informed consent was obtained for all subjects involved in the study.

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

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The authors declare that they have no competing interests

Authors' Contributions

RLS performed library preparation of the cCSC samples, performed all bioinformatics analyses including variant calling, filtering, and statistical analysis and drafted the manuscript. MBB was instrumental in collecting DNA samples from cCSC patients. CBH collected DNA samples and supervised MBB. CG provided the control samples used in this study and offered bioinformatics support. CJFB collected DNA samples, supervised MBB, and performed clinical grading of the cCSC patients. EKdJ and AldH obtained funding for the research and were responsible for design of the study, critical revisions of the manuscript and supervision of RLS. All authors provided critical feedback on the manuscript and approved the final manuscript.

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Supplementary Files

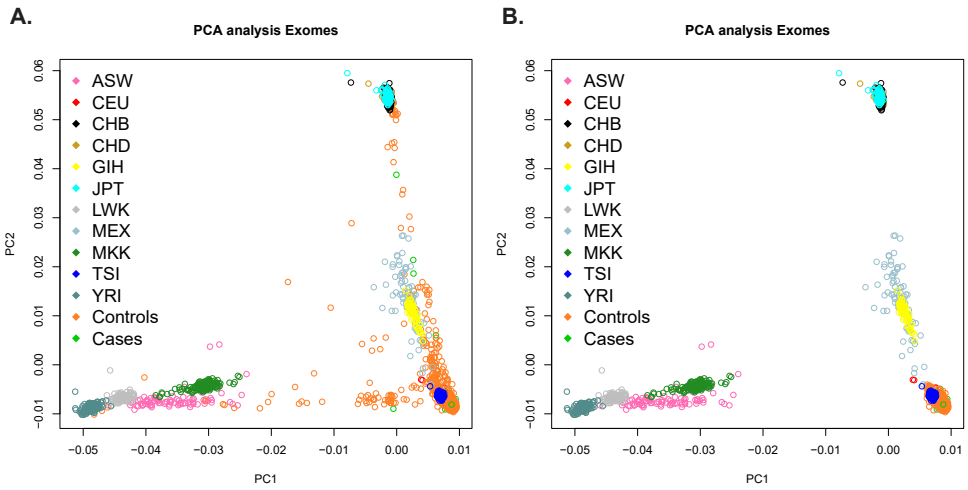


Figure S1. Principal component analysis (PCA) of the cCSC patients and controls. Graphical representation of the first 2 principal components of the PCA analysis, before (A) and after (B) filtering based on European ancestry. Various colours indicate the Hapmap3 samples according to the legend, orange depicts the controls and light green depicts the cCSC patients.

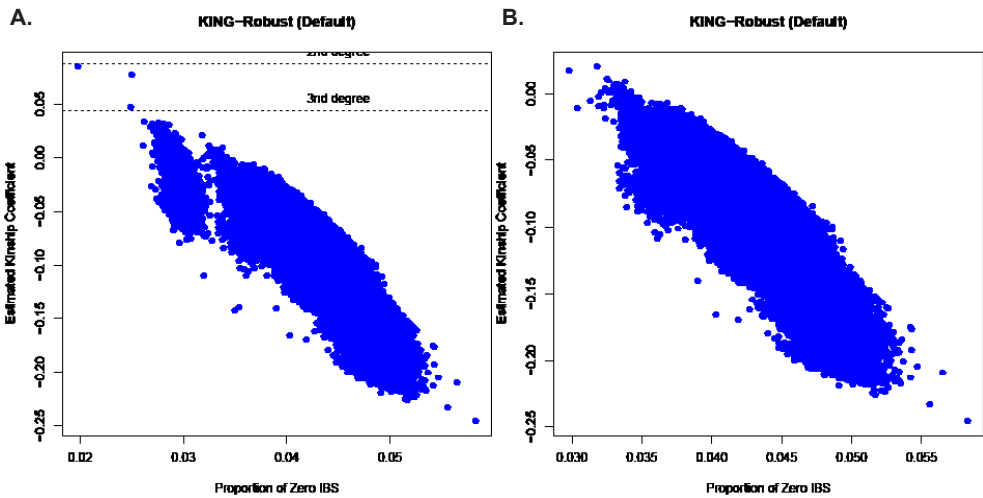


Figure S2. Kinship analysis of cCSC patients and controls. Kinship analysis was performed with KING, and estimated kinship coefficients of individuals in the study were calculated (A). Individuals with a kinship > 0.0442 were removed from the analysis (B).

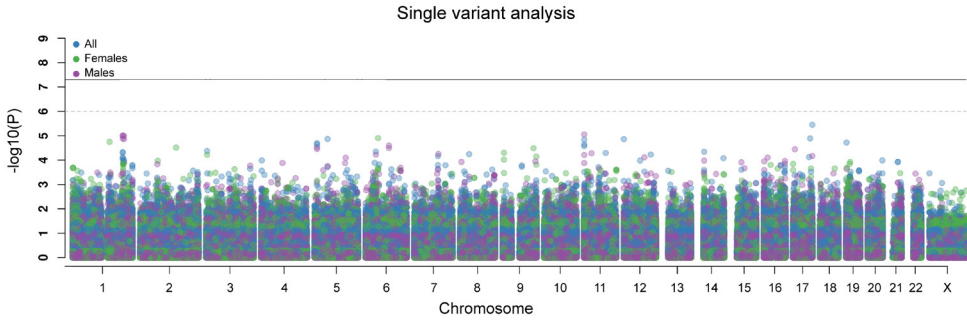


Figure S3. Single variant analysis of the exome sequencing results in cCSC patients and controls. Single variant analysis was performed with correction for 2 principal components and gender in the complete cohort (263 cCSC patients vs. 1352 controls [blue]), while for the females (55 cCSC patients vs. 681 controls [green]) and males (208 cCSC patients vs. 671 controls [purple]) only correction with the 2 principal components was applied. The dark horizontal line indicates the significance threshold after correction for multiple testing of 5×10^{-8} , while the dotted line indicates the suggestive threshold of $P = 1 \times 10^{-6}$.

151

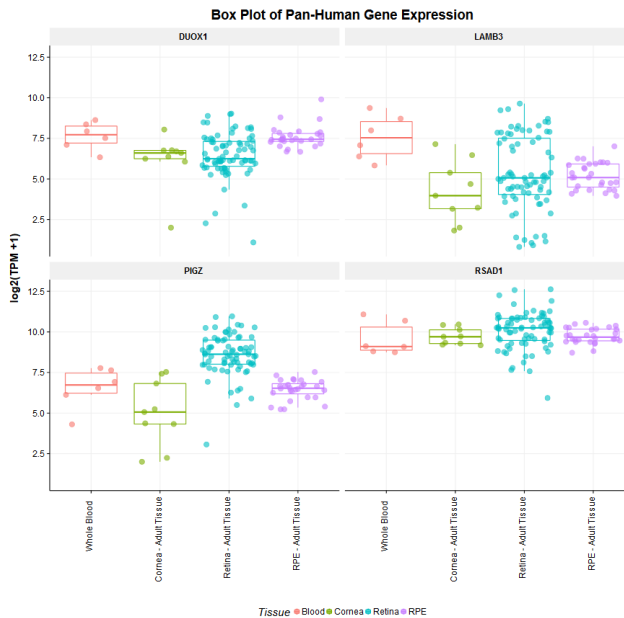


Figure S5. Gene expression of *DUOX1*, *LAMB3*, *PIGZ* and *RSAD1* in different tissues. Pan-Human Gene Expression levels of the four susceptibility genes in females according to the eyeintegration database v0.63 (27). Whole blood (red), Cornea (green), retina (blue) and retinal pigment epithelium (RPE, purple) expression levels are depicted.

Table S1. Suggestive gene-based associations using the Burden, SKAT and SKAT-O tests

Gene	Variants	Burden p-value	SKAT p-value	SKAT-O p-value
All individuals				
ALX1	Exonic	1.57x10 ⁻⁵	3.71x10 ⁻⁵	2.17x10 ⁻⁵
NOP14	Protein-altering/ Exonic	3.02x10 ⁻² / 2.49x10 ⁻²	3.76x10 ⁻⁵ / 1.52x10 ⁻⁵	8.88x10 ⁻⁵ / 3.88x10 ⁻⁵
	Exonic	4.72x10 ⁻⁵	1.71x10 ⁻³	8.45 x10 ⁻⁵
G3BP1	Exonic			
Males				
RSP02	Protein-altering	3.81x10 ⁻⁵	1.95x10 ⁻²	4.37x10 ⁻⁵
ALX1	Exonic	1.43x10 ⁻⁵	2.80x10 ⁻⁵	1.74x10 ⁻⁵
RNF144A	Exonic	2.25x10 ⁻⁵	2.25x10 ⁻⁵	2.12x10 ⁻⁵
C5orf63	Exonic	2.48x10 ⁻⁵	4.92x10 ⁻⁵	3.49x10 ⁻⁵
NOB1	Exonic	4.24x10 ⁻⁵	4.24x10 ⁻⁵	4.24x10 ⁻⁵
LOC283332	Exonic	2.77x10 ⁻⁴	4.81x10 ⁻⁵	4.64x10 ⁻⁵
G3BP1	Exonic	2.63x10 ⁻⁵	1.36x10 ⁻³	4.81x10 ⁻⁵
Females				
ZNF713	Exonic	1.68x10 ⁻⁴	3.40x10 ⁻⁶	4.86x10 ⁻⁶
SPATA7	Exonic	7.11x10 ⁻⁶	2.30x10 ⁻⁵	5.35x10 ⁻⁶
C1QL3	Exonic	1.68x10 ⁻⁴	1.03x10 ⁻⁵	1.10x10 ⁻⁵
GAPVD1	Exonic	1.45x10 ⁻²	7.43x10 ⁻⁶	1.99x10 ⁻⁵
ZNF300P1	Exonic	3.33x10 ⁻⁵	3.33x10 ⁻⁵	3.16x10 ⁻⁵
LIPH	Exonic	4.73x10 ⁻⁵	4.73x10 ⁻⁵	4.64x10 ⁻⁵
TDGF1	Protein-altering	4.89x10 ⁻⁵	4.89x10 ⁻⁵	4.64x10 ⁻⁵
OR5H14	Exonic	4.67 x10 ⁻⁵	9.33x10 ⁻⁴	5.42x10 ⁻⁵
RSL1D1	Protein-altering	3.52x10 ⁻⁵	4.28x10 ⁻⁴	9.66x10 ⁻⁵

	Counts female controls (n=681)	MAF cCSC	MAF controls	Exac_NFE	PhyloP	Gramham	Mutation Taster (score)	CADD	Sift (score)	Polyphen (score)	Trap(score)
1	0	7,34E-04	1,00E-04	0,25	45	N(1)	15,8	T (0.361)	B (0.036)	0,147	
2	0	1,47E-03	7,00E-04	1,61	58	N(0.953)	13,6	T (0.159)	B (0.124)	0,001	
1	0	7,34E-04	6,00E-04	1,29	155	D(0.558)	20,9	T (0.45)	B (0.138)	0,012	
0	9,09E-03	0	7,50E-05	1,53	125	N(0.643)	24,4	T (0.068)	D (0.999)	0,234	
1	0	7,34E-04	1,70E-03	.	.	A(1)	44,0	.	.	0,773	
2	0	1,47E-03	2,00E-04	0,06	15	N(1)	8,1	T (0.266)	B (0.003)	0,004	
1	0	7,34E-04	3,00E-05	3,56	61	D(1)	17,0	T (0.831)	D (0.957)	0,021	
0	0,018	0	2,40E-03	0,99	101	D(1)	34,0	D (0.004)	D (1)	0,048	
2	0	1,47E-03	5,00E-04	3,23	180	D(1)	26,6	D (0.014)	D (1)	0,038	
146	0,136	0,107	0,127	0,98	180	P(0.899)	15,4	T (0.364)	B (0)	0,026	
1	0,01	7,34E-04	1,52E-05	1,46	60	D(1)	10,9	T (0.38)	Prob. D (0.661)	0,014	
467	0,336	0,343	0,674	0,85	22	P(0.999)	20,5	T (0.439)	B (0.002)	0,09	
1	0	7,34E-04	1,60E-03	3,38	29	D(1)	33,0	D (0.002)	D (1)	0,059	
1	0	7,34E-04	1,30E-03	3,38	43	D(1)	23,8	T (0.277)	Prob. D (0.706)	0,332	
1	0	7,34E-04	4,50E-05	2,20	180	D(1)	33,0	D (0)	D (1)	0,116	
1	0	7,34E-04	2,00E-04	0,26	98	N(1)	16,23	D (0.015)	B (0.002)	0,021	
1	0	7,34E-04	9,98E-05	2,40	89	D(1)	15,80	D (0.018)	B (0.036)	0,214	
1	0	7,34E-04	2,00E-04	0,29	26	D(1)	9,08	T (1)	B (0)	0,475	
3	0	2,20E-03	4,00E-04	-0,93	98	N(1)	0,00	T (0.407)	B (0)	0,123	
3	0,036	2,20E-03	5,70E-03	-1,40	64	N(1)	0,00	T (1)	B (0.002)	0,144	
1	0	7,34E-04	1,30E-03	0,66	29	D(1)	23,20	T (0.129)	D (1)	0,128	
4	0	2,94E-03	2,00E-04	0,78	109	D(0.821)	16,08	T (0.164)	Prob. D (0.934)	0,327	
334	0,218	0,245	0,744	0,20	58	P(1)	0,27	T (0.323)	B (0.001)	0,063	
2	0	1,47E-03	1,50E-03	1,61	64	N(1)	0,00	T (1)	B (0.136)	0,064	
1	0	7,34E-04	1,57E-04	0,52	94	.(.)	0,03	T (0.49)	B (0.002)	0,011	
1	0	7,34E-04	4,00E-04	0,76	64	N(1)	7,99	T (0.164)	Prob. D (0.919)	0,078	
557	0,391	0,409	0,596	0,46	43	P(1)	8,52	T (0.505)	B (0)	0,166	
1	0	7,34E-04	4,00E-04	-2,76	101	N(1)	0,00	T (0.188)	B (0.001)	0,009	
189	0,164	0,139	0,141	0,98	64	P(0.947)	7,25	T (0.252)	B (0.007)	0,081	
2	9,09E-03	1,47E-03	1,90E-03	3,82	10	D(0.937)	25,20	D (0.036)	D (1)	0,043	
117	0,045	0,086	0,084	2,20	10	N(0.712)	22,10	T (0.143)	Prob. D (0.524)	0,431	
1	0	7,34E-04	3,00E-05	3,82	98	D(0.931)	24,90	D (0.005)	D (0.998)	0,297	
0	9,09E-03	0	5,00E-04	-0,60	21	N(1)	2,73	T (0.18)	Prob. D (0.539)	0,01	

LAMB3 (NM_001017402)

Gene	Variant	Exon	Effect	Nucleotide change	Protein change	SKAT P-value	SAS P-value	OR (95% CI)	Weight	Counts female C S C (n=55)
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RSADI (NM_018346)

LAMB3 (NM_001017402)

Gene	Variant	Exon	Effect	Nucleotide change	Protein change	SKAT P-value	SAS P-value	OR (95% CI)	Weight	Counts female C C C (n=55)
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Table S3. Haplotypes of LAMB3 in the female cohort

Haplotype	Pooled HF		Control HF		CSC patients HF		OR (95% CI)	
	0	1	0	1	0	1	Base	F-value
rs745444340	1	0	0	0	0	0	0	0
rs147931502	0	0	0	0	0	0	0	0
rs1475272725	0	0	0	0	0	0	0	0
1:209791343	0	0	0	0	0	0	0	0
rs140769823	0	0	0	0	0	0	0	0
rs143164025	0	0	0	0	0	0	0	0
rs77389254	0	0	0	0	0	0	0	0
rs775023132	0	0	0	0	0	0	0	0
rs2076222	0	0	0	0	0	0	0	0
rs199946321	0	0	0	0	0	0	0	0
rs539752226	0	0	0	0	0	0	0	0
rs151006337	0	0	0	0	0	0	0	0
1:209796424	0	0	0	0	0	0	0	0
rs901375494	0	0	0	0	0	0	0	0
rs113745536	0	0	0	0	0	0	0	0
rs78247685	0	0	0	0	0	0	0	0
rs2072937	0	0	0	0	0	0	0	0
rs2229466	0	0	0	0	0	0	0	0
1:209799011	0	0	0	0	0	0	0	0
1:209792027	0	0	0	0	0	0	0	0
rs80356682	0	0	0	0	0	0	0	0
rs760667887	0	0	0	0	0	0	0	0
rs2229465	0	0	0	0	0	0	0	0
rs771720883	0	0	0	0	0	0	0	0
rs370280436	0	0	0	0	0	0	0	0
rs2179402	0	0	0	0	0	0	0	0
rs767004520	0	0	0	0	0	0	0	0
rs2076349	0	0	0	0	0	0	0	0
1:209800233	0	0	0	0	0	0	0	0
rs771727419	0	0	0	0	0	0	0	0
1:209800257	0	0	0	0	0	0	0	0
rs7624400142	0	0	0	0	0	0	0	0
rs2229468	0	0	0	0	0	0	0	0
1:209801457	0	0	0	0	0	0	0	0
rs779219301	0	0	0	0	0	0	0	0
rs2076351	0	0	0	0	0	0	0	0
rs548096567	0	0	0	0	0	0	0	0
rs114875539	0	0	0	0	0	0	0	0
rs2814161	0	0	0	0	0	0	0	0
rs886045867	0	0	0	0	0	0	0	0
rs141059189	0	0	0	0	0	0	0	0
rs375846575	0	0	0	0	0	0	0	0
rs121912482	0	0	0	0	0	0	0	0
rs150895872	0	0	0	0	0	0	0	0
rs2235542	0	0	0	0	0	0	0	0
rs2076356	0	0	0	0	0	0	0	0
rs757259426	0	0	0	0	0	0	0	0
rs2228339	0	0	0	0	0	0	0	0
rs140470350	0	0	0	0	0	0	0	0

Base: Haplotype used as reference in the association model; HF: Haplotype frequency; NA: Not annotated; PC: Principal component of the ancestry analysis

Corrected with PC1 and PC2

0.020	0.077	0.014	0.116	0.062	0.143	0.012	0.508
0.016	0.074	0.012	0.114	0.060	0.145	0.013	0.518
0.073	0.112	NA	0.116	0.053	0.129	NA	0.412
7.22E-05	0.045	0.108	0.521	0.447	0.929	0	0
6.60	2.03	2.72	1.24	1.40	1.03	0 (0.0)	Base
(12.61-	(1.02-	(0.80-	(0.64-	(0.59-	(0.53-	NA	OR (95% CI)
16.67)	4.04)	9.18)	2.41)	3.30)	2.00)		

Table S4. Male rare variant (MAF <0.05) carriers in *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1*

Gene	No. rare variants in gene	Controls (n=671)	cCSC patients (n=208)	P-value	OR (95% CI)
<i>LAMB3</i>	32	81 (12.1%)	20 (9.6%)	0.38	0.78 (0.46-1.30)
<i>PIGZ</i>	19	35 (5.2%)	7 (3.4%)	0.35	0.63 (0.28-1.45)
<i>DUOX1</i>	23	24 (3.6%)	8 (3.8%)	1.00	1.08 (0.48-2.44)
<i>RSAD1</i>	4	7 (1.0%)	1 (0.5%)	0.69	0.46 (0.06-3.73)
Variant in 1 or more genes		137 (20.4%)	33 (15.9%)	0.15	0.74 (0.49-1.12)

Table S5. Gene-burden results of genes in the known cCSC loci

Gene	Locus	All individuals								Males		
		exonic				protein altering				exonic		
		Variants	Burden	SKAT	SKAT-O	Variants	Burden	SKAT	SKAT-O	Variants	Burden	SKAT
ARMS2	ARMS2-HT-RA1	6	0,358	0,681	0,498	5	0,593	0,865	0,725	5	0,377	0,602
HTRA1	ARMS2-HT-RA1	10	0,763	0,910	0,918	6	0,844	0,632	0,866	9	0,716	0,859
PLEKHA1	ARMS2-HT-RA1	18	0,572	0,267	0,432	12	0,118	0,248	0,201	15	0,209	0,149
CR1L	CD46	8	0,542	0,844	0,645	7	0,546	0,877	0,663	7	0,673	1,000
CD46	CD46	10	0,405	0,689	0,504	7	0,347	0,666	0,431	7	0,163	0,151
CD34	CD46	16	0,219	0,033	0,047	9	0,113	0,033	0,047	12	1,000	0,575
BEAN1	CDH5	24	0,758	1,000	0,892	16	0,583	0,668	0,694	16	0,589	0,633
CDH5	CDH5	39	0,787	0,892	1,000	23	0,867	0,577	0,798	25	0,549	0,596
CFHR2	CFH	6	0,090	0,389	0,136	4	0,193	0,494	0,270	6	0,249	0,685
CFHR4	CFH	7	0,294	0,146	0,242	4	0,262	0,709	0,400	6	1,000	0,410
KCNT2	CFH	26	0,733	0,188	0,271	11	0,140	0,057	0,119	14	0,552	0,278
CFHR5	CFH	24	0,374	0,358	0,524	19	0,350	0,338	0,497	17	0,943	0,641
CFH	CFH	33	0,717	0,333	0,515	19	0,276	0,464	0,423	24	0,184	0,073
NR3C2	NR3C2	15	0,227	0,737	0,348	8	0,109	0,782	0,185	12	0,332	0,881
ARHGAP10	NR3C2	31	0,570	0,630	0,742	16	0,202	0,932	0,276	24	0,866	0,363
PPT2	RCCX	4	0,170	0,109	0,151	3	0,789	0,793	1,000	4	0,128	0,143
STK19	RCCX	10	0,590	0,600	0,754	4	0,187	0,519	0,380	8	0,846	0,965
ZBTB12	RCCX	8	0,482	0,617	0,566	5	0,372	0,776	0,481	5	0,516	0,599
FKBPL	RCCX	7	0,699	0,729	0,852	6	0,889	0,746	0,885	7	0,740	0,264
RDBP	RCCX	10	0,040	0,018	0,040	8	0,046	0,021	0,028	7	0,035	0,011
DOM3Z	RCCX	19	1,000	0,777	1,000	9	1,000	0,783	1,000	15	1,000	0,502
EGFL8	RCCX	14	0,655	0,223	0,332	10	0,064	0,162	0,107	10	1,000	0,374
CFB	RCCX	17	0,350	0,703	0,505	11	0,204	0,646	0,310	15	0,836	1,000
ATF6B	RCCX	21	0,173	0,682	0,268	11	0,248	1,000	0,373	16	0,522	1,000
SLC44A4	RCCX	25	0,483	0,641	0,656	15	0,404	0,706	0,597	14	0,257	1,000
EHMT2	RCCX	36	0,208	0,183	0,284	16	0,153	0,073	0,113	22	0,656	0,538
C2	RCCX	24	0,811	0,645	0,786	20	0,660	0,591	0,728	18	0,682	0,676
AGER	RCCX	31	0,209	0,745	0,324	20	0,275	0,368	0,385	25	0,320	0,818
SKIV2L	RCCX	45	0,922	0,283	0,429	24	0,798	0,215	0,327	33	0,596	0,050
NOTCH4	RCCX	72	0,685	1,000	0,856	47	0,544	0,939	0,740	55	0,591	1,000
TNXB	RCCX	116	0,519	0,404	0,576	84	0,221	0,346	0,351	87	0,228	0,048
GPSM3	RCCX	2	0,221	0,221	0,221	NA	NA	NA	NA	2	0,094	0,094
MIR1236	RCCX	3	0,104	0,379	0,196	NA	NA	NA	NA	2	0,076	0,279

Gene	Locus	All individuals							Males			
		exonic				protein altering			exonic			
		Va-ri-ants	Burden	SKAT	SKAT-O	Va-ri-ants	Burden	SKAT	SKAT-O	Vari-ants	Burden	SKAT

0,716	NA	NA	NA	NA	2	0,037	0,037	0,037	NA	NA	NA	NA
0,372	NA	NA	NA	NA	3	0,729	0,729	0,729	NA	NA	NA	NA
0,541	NA	NA	NA	NA	3	0,115	0,115	0,115	NA	NA	NA	NA
0,926	NA	NA	NA	NA	7	0,176	0,335	0,338	3	0,079	0,079	0,079
0,787	4	0,527	0,753	0,787	2	0,585	0,585	0,585	2	0,585	0,585	0,585
0,813	6	0,632	1,000	0,736	13	0,785	1,000	1,000	6	0,934	1,000	1,000
0,181	8	0,317	0,535	0,439	18	0,337	0,179	0,275	8	0,823	0,782	1,000
0,203	12	0,890	0,057	0,100	12	0,250	0,193	0,317	10	0,097	0,118	0,143
0,290	11	1,000	0,603	0,806	23	0,131	0,569	0,218	10	0,079	0,102	0,148
0,798	4	0,764	0,683	0,781	15	0,867	0,590	0,800	5	0,954	0,307	0,307
0,512	20	0,396	0,817	0,569	22	0,979	0,596	0,754	13	1,000	0,495	0,674

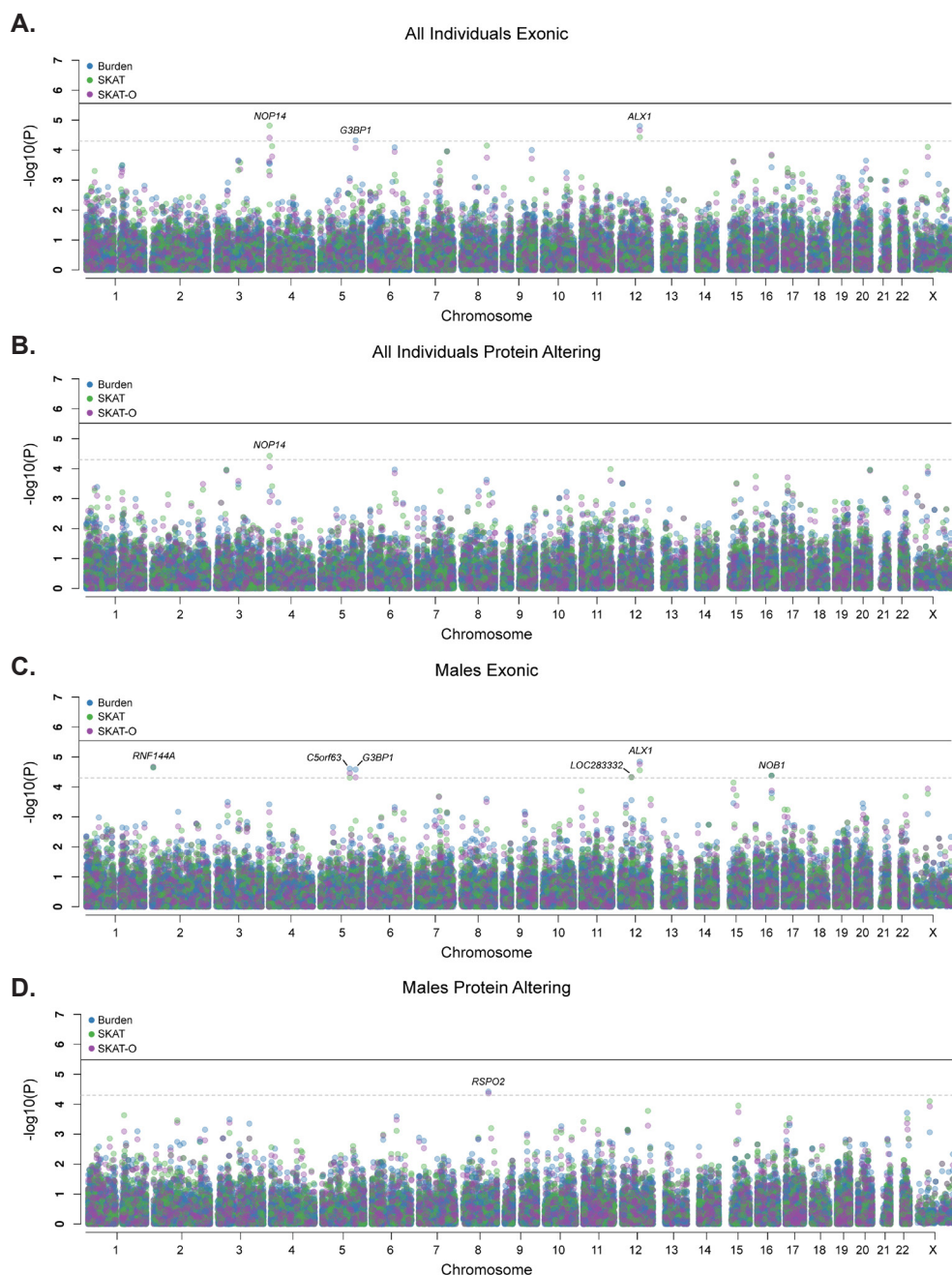


Figure S4. Gene-burden analysis all cCSC individuals and the male cCSC cohort. Burden (blue), SKAT (green) and SKAT-O (purple) association results are depicted for the complete cohort of 263 cCSC patients and 1352 controls corrected for gender and 2 principal components, using the Exonic (A) and Protein Altering (B) group files. The results the gene-burden test in the 208 male cCSC patients and 671 male controls, for both group files are also depicted (C+D). The dark horizontal line indicates the significance threshold after correction for multiple testing: $0.05/18,173 \text{ genes} = 2.75 \times 10^{-6}$ (A), $0.05/16,370 \text{ genes} = 3.05 \times 10^{-6}$ (B), $0.05/17,348 \text{ genes} = 2.88 \times 10^{-6}$ (C) and $0.05/15,083 \text{ genes} = 3.31 \times 10^{-6}$ (D), while the dotted line indicates the suggestive threshold of $P = 5 \times 10^{-5}$.

Script File S1

```
#Convert data to plink files and make a dummy gene list based on the variants
in the bim files
#In the following format:
#dummygene    SNP1
#dummygene    SNP2
awk '{print("dummygene""\t"$2)}' input.bim | sort -k2 -u > SNPsets_dummy.txt

#Generate weight file based on minor allele frequency
R
library(SKAT)
#Generate the File.SSD and File.Info
Generate_SSD_SetID("input.bed", "input.bim", "input.fam", "SNPsets_dummy.
txt", "File.SSDdummy", "File.Infodummy")
#read the covariate file and fam file if applicable
FAM_Cov<-Read_Plink_FAM_Cov("input.fam", "input_cov.txt", Is.binary=TRUE,
flag1=0, cov_header=TRUE)
SSD.INFOdummy<-Open_SSD("File.SSDdummy", "File.Infodummy")
#Get genotype matrix of the dummy geneset
dummy_geneset<-Get_Genotypes_SSD(SSD.INFOdummy, 1, is_ID = TRUE)
weight<-Get_Logistic_Weights(dummy_geneset, par=0.07, par2=150)
weights<-data.frame(weight)
weights[,2] <- colnames(dummy_geneset)
weights = weights[, c(2,1)]
write.table(weights, file="snp_weights.txt", col.names=FALSE, row.names=FALSE,
sep="\t", quote = FALSE)

#perform single variant association with dummy geneset dataset
#Define the covariates if applicable
Phenotype = FAM_Cov$Phenotype
PC1 = FAM_Cov$PC1
PC2 = FAM_Cov$PC2
Sex = FAM_Cov$Sex
#Run the null model
obj<-SKAT_Null_Model_MomentAdjust(Phenotype ~ PC1 + PC2 + Sex, type.
Resampling="bootstrap.fast")
#Run the loop through the genotype matrix and output the p-values
single_pvalue <- NULL
for(snp in colnames(dummy_geneset)){out <- SKATBinary_Single(dummy_
geneset[,snp], obj); single_pvalue <- rbind(single_pvalue, data.frame(snp,
out$p.value, out$method.bin, out$m))}
#print the output file
write.table(single_pvalue, file="Single_pvalue_all.txt", col.names=TRUE, row.
names=FALSE, sep="\t", quote = FALSE)

#Run the burden, SKAT and SKAT-O test using the snp_weights_file.txt
Generate_SSD_SetID("input.bed", "input.bim", "input.fam", "groupfile.txt",
"File.SSD_groupfile", "File.Info_groupfile")
#read the covariate file and fam file if applicable
FAM_Cov<-Read_Plink_FAM_Cov("input.fam", "input_cov.txt", Is.binary=TRUE,
flag1=0, cov_header=TRUE)
SSD.INFO_groupfile<-Open_SSD("File.SSD_groupfile", "File.Info_groupfile")
Phenotype = FAM_Cov$Phenotype
PC1 = FAM_Cov$PC1
PC2 = FAM_Cov$PC2
Sex = FAM_Cov$Sex.x
obj<-SKAT_Null_Model_MomentAdjust(Phenotype ~ Sex + PC1 + PC2, type.
Resampling="bootstrap.fast")
```

```

custom_weights<-Read_SNP_WeightFile("snp_weights.txt")
#Burden test
out.burden<-SKATBinary.SSD.All(loaded_SSD, obj, method="Burden", obj.
SNPWeight=custom_weights, kernel="linear.weighted")
#SKAT test
out.skate<-SKATBinary.SSD.All(loaded_SSD, obj, method="SKAT", obj.
SNPWeight=custom_weights, kernel="linear.weighted")
#SKAT-O test
out.skato<-SKATBinary.SSD.All(loaded_SSD, obj, method="SKATO", obj.
SNPWeight=custom_weights, kernel="linear.weighted")

#keep only genes with more than 2 variants and perform correction for multiple
testing
out.burden_select <- out.burden[out.burden$N.Marker.Test>1,]
out.burden_select$FDR <- p.adjust(out.burden_select$P.value, method="fdr")
out.burden_select$Bonf <- p.adjust(out.burden_select$P.value,
method="bonferroni")
write.table(out.burden_select, file="burden_output_groupfile.txt", col.
names=TRUE, row.names=FALSE, sep="\t", quote = FALSE)

out.skate_select <- out.skate[out.skate$N.Marker.Test>1,]
out.skate_select$FDR <- p.adjust(out.skate_select$P.value, method="fdr")
out.skate_select$Bonf <- p.adjust(out.skate_select$P.value, method="bonferroni")
write.table(out.skate_select, file="skate_output_groupfile.txt", col.names=TRUE,
row.names=FALSE, sep="\t", quote = FALSE)

out.skato_select <- out.skato[out.skato$N.Marker.Test>1,]
out.skato_select$FDR <- p.adjust(out.skato_select$P.value, method="fdr")
out.skato_select$Bonf <- p.adjust(out.skato_select$P.value,
method="bonferroni")
write.table(out.skato_select, file="skato_output_groupfile.txt", col.
names=TRUE, row.names=FALSE, sep="\t", quote = FALSE)

#plot the data in CMplot
library(CMplot)
ref_genes=read.table("genes_location.txt", header=T)
data_combined1 <- merge(out.burden_select[,c(1,2)], out.skate_select[,c(1,2)],
by="SetID", all=TRUE)
data_combined2 <- merge(data_combined1, out.skato_select[,c(1,2)], by="SetID",
all=TRUE)
data_combined3 <- merge(ref_genes, data_combined2 , by=1, no.dups=TRUE)
colnames(data_combined3) = c("GENE", "CHR", "BP", "burden", "skate", "skato")
nrow(data_combined3)
#genome-wide significant line --> 0.05/nrow=pvalue_cutoff
data_combined3[is.na(data_combined3)] <- 1

CMplot(data_combined3, plot.type="m", multitracks=TRUE, threshold=c(pvalue_
cutoff,suggestive_line), threshold.lty=c(1,2),
threshold.lwd=c(1,1), threshold.col=c("black","grey"), amplify=TRUE,bin.
size=1e6,
chr.den.col=NULL, signal.col=c("red","green"),signal.cex=c(1,1),
file="pdf",memo="results")

```

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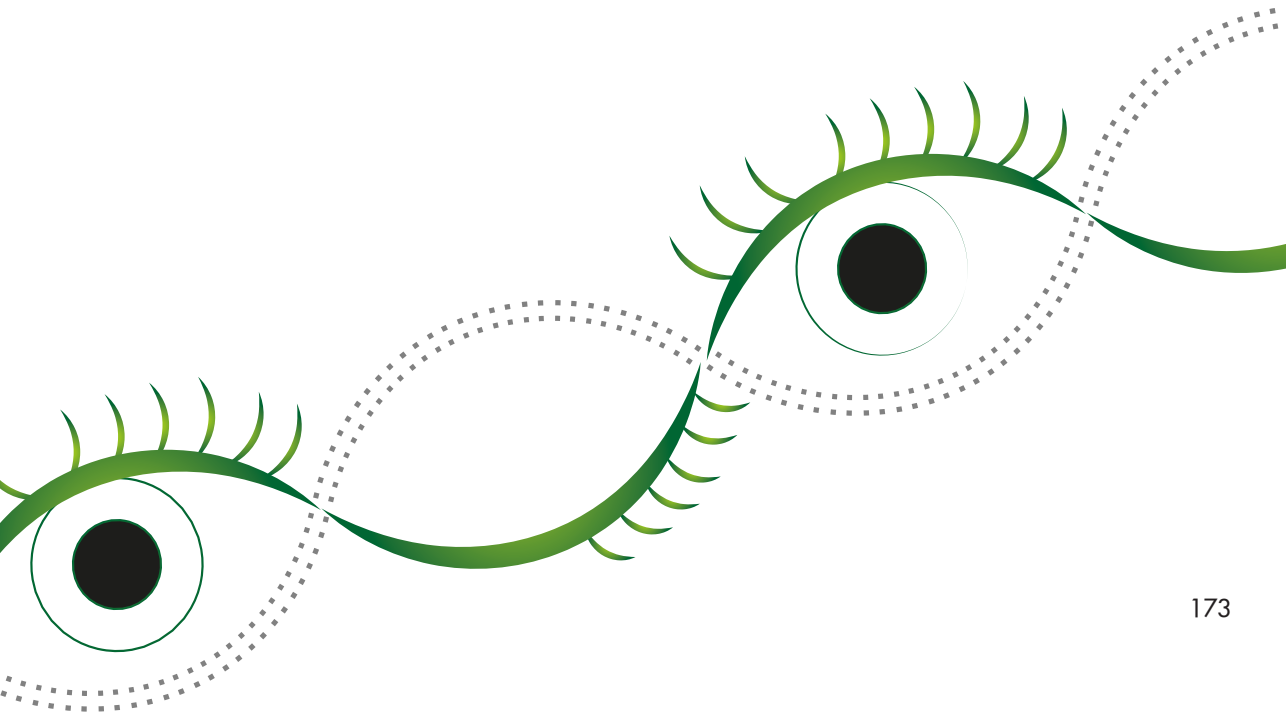
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4.1 Familial central serous chorioretinopathy

4.2 Exome sequencing in familial chronic central serous chorioretinopathy

4. Familial Central Serous Chorioretinopathy





4.1 Familial central serous chorioretinopathy

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* These authors contributed equally to this study

Abstract

Purpose: To assess ophthalmological characteristics in patients and unaffected individuals in families with multiple members affected by central serous chorioretinopathy (CSC), both at presentation and long-term follow-up.

Methods: In 103 subjects from 23 families, with at least 2 affected patients with CSC per family, prospective extensive ophthalmological examination was performed, including best-corrected visual acuity, indirect ophthalmoscopy, digital color fundus photography, optical coherence tomography, fundus autofluorescence, and fluorescein angiography imaging. From these, 24 individuals from 6 families had undergone extensive ophthalmological examination in either 1994 or 1995 and were followed up in this study.

Results: Subretinal fluid accumulation on optical coherence tomography and/or “hot spots” of leakage on fluorescein angiography indicative of CSC were detected in 45 of 103 phenotyped subjects (44%). Findings suggestive of CSC, but without the presence of subretinal fluid on optical coherence tomography and/or “hot spots” of leakage on fluorescein angiography, were observed in an additional 27 family members (26%). In 4 out of 17 previously non-affected subjects (24%) from the 24 individuals that were followed up after more than 20 years, we found more severe abnormalities.

Conclusion: Extensive ophthalmological phenotyping resulted in the detection of (suggestive) CSC in 52% of family members of patients with CSC. Genetic factors may play an important role in these specific CSC cases. Moreover, during follow-up progressive disease can occur in a noteworthy number of patients.

Introduction

Central serous chorioretinopathy (CSC) is a chorioretinal disease characterized by a detachment of the neurosensory retina because of serous subretinal fluid (SRF). The disease is hypothesized to occur as a result of hyperpermeability and thickening of the choroid (pachychoroid), and subsequent damage to the retinal pigment epithelium (RPE).¹⁻⁵ Exposure to exogenous corticoids is the most important risk factor for CSC, with odds ratios of up to 37.⁶⁻⁸ Genetic risk and protective factors have also been identified in chronic CSC, although the exact pathogenetic mechanism of the disease is still unclear.⁹⁻¹³ Single nucleotide polymorphisms (SNPs) in the *age-related maculopathy susceptibility 2* (ARMS2), *complement factor H* (CFH), *cadherin 5* (CDH5), and *nuclear receptor subfamily 3 group C member 2* (NR3C2, mineralocorticoid receptor) genes, and the absence of the *complement component 4B* (C4B) gene have been associated with an increased risk for CSC in whites.¹⁰⁻¹³ The risk-conferring CFH association was also observed in an Asian cohort.⁹ In addition, the presence of three copies of the C4B gene and several other single nucleotide polymorphisms in the CFH gene were associated with a protective effect for CSC.^{11,13}

Familial occurrence of CSC has been described previously, but its occurrence appears to be rare.¹⁴⁻¹⁷ A recent study described a thickened choroid (>395 μm) in 50% of eyes from family members of patients with CSC, suggestive of an autosomal-dominant inheritance pattern for pachychoroid.¹⁸ A previous study in which ophthalmological imaging was performed in at least 2 members per family described multiple affected family members in 52% of 27 families and fundus lesions suggestive of CSC in 44% out of 80 screened unaffected relatives.¹⁵ That work, performed over 20 years ago, did not address whether any progression was observed in the unaffected family members that had suspected lesions. In our current study, through extensive ophthalmological imaging in newly recruited families with multiple members affected by CSC, we aimed to corroborate earlier observations on the occurrence of suspected lesions in unaffected family members. Moreover, we followed up a number of individuals with reported suspected lesions over 20 years ago to ascertain whether these individuals demonstrated any sign of progression of the disease.

Methods

Subject selection

In this multicenter prospective cohort study, 103 subjects from 23 families including probands were phenotyped either at the Department of Ophthalmology of the Radboud university medical center ([Radboudumc] Nijmegen, the Netherlands), the Leiden University Medical Center ([LUMC] Leiden, the Netherlands), or the Rotterdam Eye Hospital (Rotterdam, the Netherlands). All patients were recruited at the outpatient clinic of the participating hospitals, after the proband had reported a history of having additional family members affected by CSC. Thirty subjects (from six families) out of 103, were previously phenotyped at the Department of Ophthalmology of the LUMC, before the era of optical coherence tomography (OCT) and multimodal imaging, using only indirect ophthalmoscopy and fluorescein angiography (FA), and were invited for a long-term follow-up visit. At the previous visit, findings characteristic for CSC were seen in 9 subjects (30%) and a suspicion of CSC in 16 subjects (53%).¹⁵ To update and extend the available ophthalmological information for these patients, we were able

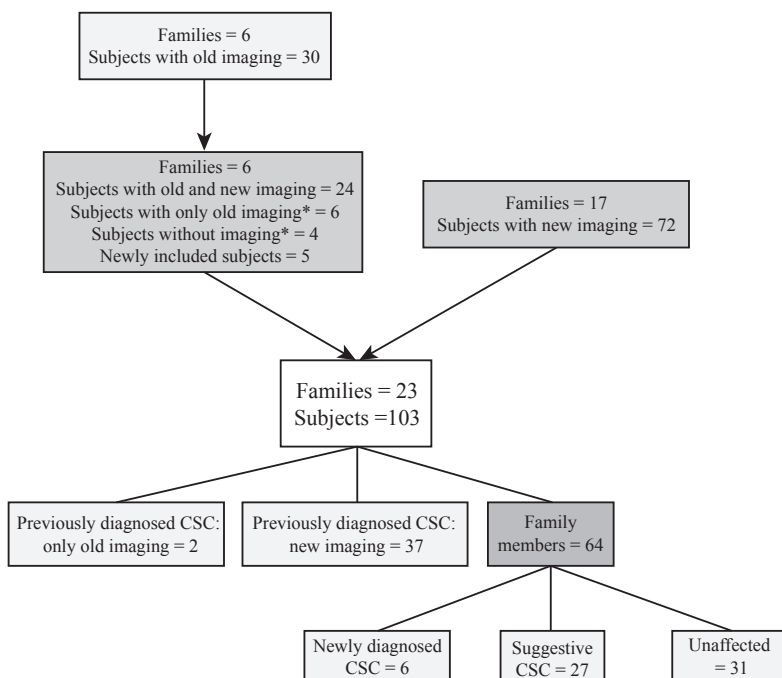


Figure 1: Outline of selection of patients with CSC who underwent ophthalmologic phenotyping. *Family members who were excluded for ophthalmologic analysis, except for the two family members in whom CSC had been previously diagnosed.

177

to include 24 of these 30 subjects for new imaging. Five additional new members of these families were also recruited and studied. Another 72 members from 17 other families, in which the proband reported either a possible or a confirmed CSC diagnosis in one or more additional family members were invited for phenotyping (Figure 1).

In total, before ophthalmologic phenotyping for this study, in 39 subjects (30 male subjects and 9 female subjects) of the 23 families, the diagnosis of CSC had already been confirmed. Of these 39 patients, 37 patients could be phenotyped again for this study. Two patients with confirmed diagnosis of familial CSC could not be invited for new phenotyping, but were included in the analysis. Subjects were included from April 2011 to August 2016. Written informed consent for the enrollment was obtained from all subjects. The study adhered to the tenets of the Declaration of Helsinki. Approval of the institutional review board and the ethics committee was obtained for all centers involved.

Ophthalmological imaging

Subjects included in this study received an extensive ophthalmological examination. First, an Early Treatment of Diabetic Retinopathy Study (ETDRS) best-corrected visual acuity (BCVA) measurement was performed. A previously published method was used to convert Snellen BCVA to ETDRS BCVA, when this was not available.¹⁹ Next, pupils were dilated with 1% tropicamide and 5% phenylephrine. Subsequently, indirect ophthalmoscopy and digital color fundus photography (Topcon Corp., Tokyo, Japan)

were performed. Using the spectral-domain OCT device (SPECTRALIS HRA+OCT; Heidelberg Engineering, Dublin, CA), regular OCT, enhanced depth imaging (EDI) OCT, fundus autofluorescence (FAF), and FA images were obtained. Fluorescein angiography imaging was performed at 10, 15, 20, 25, and 30 minutes after oral administration of 10 ml of 20% fluorescein, after a subject had fasted for at least 3 hours. When treatment or follow-up was necessary on the basis of the obtained images, subjects were invited for a regular visit to the outpatient clinic. An experienced retina specialist (C.J.F.B.) assessed the obtained images in a masked manner.

Subjects were categorized as having CSC when serous SRF could be detected on OCT and when ≥ 1 "hot spot" of leakage or diffuse leakage in combination with irregular RPE window defects was present on FA. No signs of either polypoidal choroidal vasculopathy or age-related macular degeneration, or other atypical findings could be present. In subjects who could not be reinvited for phenotyping and for whom OCT images were not available, a description of a serous neurosensory detachment on ophthalmoscopy in combination with FA abnormalities was considered confirmative for the diagnosis of CSC. Based on multimodal imaging, a diagnosis of a clinical picture suggestive of a CSC background (suggestive CSC) could be established in subjects who did show RPE abnormalities either typical for CSC without SRF on OCT or typical for pachychoroid pigment epitheliopathy.⁵

Choroidal thickness (CT) was defined as the distance from the outer part of the hyperreflective RPE layer to the hyperreflective line of the inner surface of the sclera, and was assessed manually using the caliper tool provided by the SPECTRALIS HRA+OCT machine in subjects for whom EDI-OCT was available. Subjects with a CT of $>395 \mu\text{m}$ were considered to have pachychoroid.¹⁸

Statistical analysis

Statistical analyses were performed in IBM SPSS Statistics, version 23.0 (IBM Corp., Armonk, NY) and GraphPad Prism, version 5.03 (GraphPad Software Inc., La Jolla, CA). The distribution of the ETDRS BCVA and CT values of affected and unaffected eyes, and eyes in which suggestive CSC was found at moment of diagnosis were analyzed with the D'Agostino-Pearson omnibus normality test. Non-parametric tests were used for further analysis, when at least 1 of the compared groups was

Table 1. Clinical characteristics of different subgroups

	Known CSC	New CSC	Suspected CSC	Un-affected	Total
No. of patients	39	6	27	31	103
% Males	77	50	44	42	56
Mean age \pm SD (years)	49 \pm 12	48 \pm 15	53 \pm 9	52 \pm 11	51 \pm 11
History of cardiovascular diseases, %	41	50	52	36	43
History of depression, %	8	0	0	0	3
Hypertension, %	36	33	44	23	34
Steroid use <1 year diagnosis (%)	4/39 (10)	0	5/27 (18)	7/31 (23)	16/103 (16)

not normally distributed. The CT values of affected and unaffected eyes, and eyes with suggestive CSC at moment of diagnosis were compared using a Kruskal-Wallis test with Dunn's post-hoc comparison adjustment. ETDRS BCVA of affected eyes at moment of diagnosis and at follow-up was compared using Wilcoxon's matched-pairs signed-rank test. *P* values <0.05 were considered to be statistically significant.

Results

In total, 23 families with multiple individuals diagnosed with CSC were included in this study, resulting in the inclusion of 103 subjects (Figure 1). This included 39 previously diagnosed CSC patients and 64 undiagnosed family members, who were extensively phenotyped. Families were of variable size, with a range of 2 to 10 family members participating in the study (Figure 2 and Supplementary Figure 1). In 6 families (26%), CSC could be diagnosed in multiple generations, with a father-to-son transmission in three families. Reduced penetrance seemed to be present in 2 families (Supplementary Figure 1 P and W), and possible maternal transmission was found in one family (Supplementary Figure 1O).

Characteristics of previously diagnosed CSC patients

Thirty-nine patients (30 men and 9 women), who were previously diagnosed with CSC, were phenotyped in this study. The age at diagnosis of these patients was 49 ± 12 years (Table 1). The median ETDRS BCVA of the 63 affected eyes at diagnosis was 80 letters (Snellen equivalent: 20/25; Q1:70-Q3:85) and 85 (Snellen equivalent: 20/20; Q1:81-Q3:89) in the 15 unaffected eyes. The median ETDRS BCVA of the 56 affected eyes, for which new ophthalmological phenotyping could be performed after

179

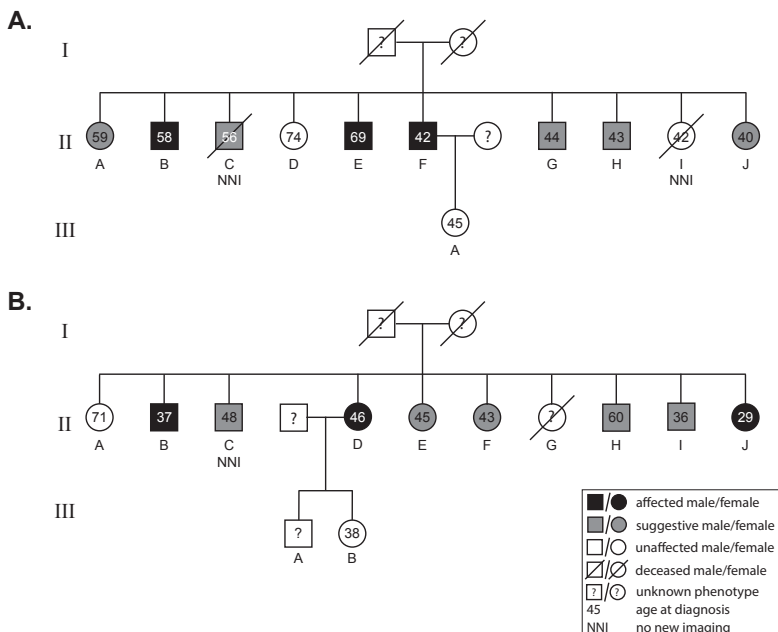


Figure 2: Pedigrees of two CSC families (A and B) with multiple affected family members. All family members were phenotyped again for this study with the exception of the individuals indicated with NNI (no new images).

a median follow-up of 38 months (Q1:11-Q3:160), was 80 letters (Snellen equivalent: 20/25; Q1:68-Q3:85) and 84 (Snellen equivalent: 20/21; Q1:78-Q3:89) in the 12 unaffected eyes, which did not differ from the ETDRS BCVA at the moment of CSC diagnosis.

In 14 of the 39 patients (36%), a history of hypertension was present and in a total of 16 patients (41%), cardiovascular diseases had been previously diagnosed. Three patients were clinically diagnosed with depression. One patient was diagnosed with Cushing disease after she had presented with CSC and three patients were previously known with amblyopia. A total of four patients (10%) reported steroid use within a year before the first diagnosis of CSC, out of whom three patients had received a renal transplant, for which low-dose steroids were prescribed at the moment of ophthalmological phenotyping.

Of the 39 patients, 23 had previously received treatment in a total of 27 eyes: 18 eyes received photodynamic therapy, 11 eyes underwent high-density subthreshold micropulse laser, and 6 eyes were treated with focal thermal laser. In eight other eyes of seven patients, intravitreal injections with either bevacizumab (7 eyes) or ranibizumab (1 eye) were performed because of either the suspicion or the presence of a choroidal neovascularization. Of these eight eyes, two eyes of two different patients exclusively received intravitreal anti-vascular endothelial growth factor injections, after which SRF disappeared in one eye.

The remaining six eyes also received other types of treatment, after which SRF leakage disappeared in five eyes (83%). Thirteen eyes had received treatment with a single treatment modality, which lead to complete resolution of SRF in 8 eyes (62%), whereas 14 eyes were treated with multiple treatment modalities, leading to an

Table 2. Performed treatments for central serous chorioretinopathy and outcome of these treatments, in 23 treated probands

Type of treatment(s)	No. of patients	No. of eyes	No SRF (leakage) at final follow-up
Treatment with one treatment modality	13	13	8/13 (62%)
Focal thermal laser	2	2	2/2 (100%)
HSML	4	4	2/4 (50%)
Intravitreal bevacizumab injections	2	2	1/2 (50%)
PDT	5	5	3/5 (60%)
Treatment with multiple treatment modalities	14	14	10/14 (71%)
Focal thermal laser + intravitreal bevacizumab injections + PDT	1	1	1/1 (100%)
Focal thermal laser + intravitreal bevacizumab injections	1	1	0/1 (0%)
Focal thermal laser + PDT	2	2	2/2 (100%)
HSML + PDT	6	6	4/6 (67%)
HSML + intravitreal ranibizumab injections + PDT	1	1	0/1 (0%)
Intravitreal bevacizumab injections + PDT	2	3	3/3 (100%)

HSML, high-density subthreshold micropulse laser; PDT: photodynamic therapy

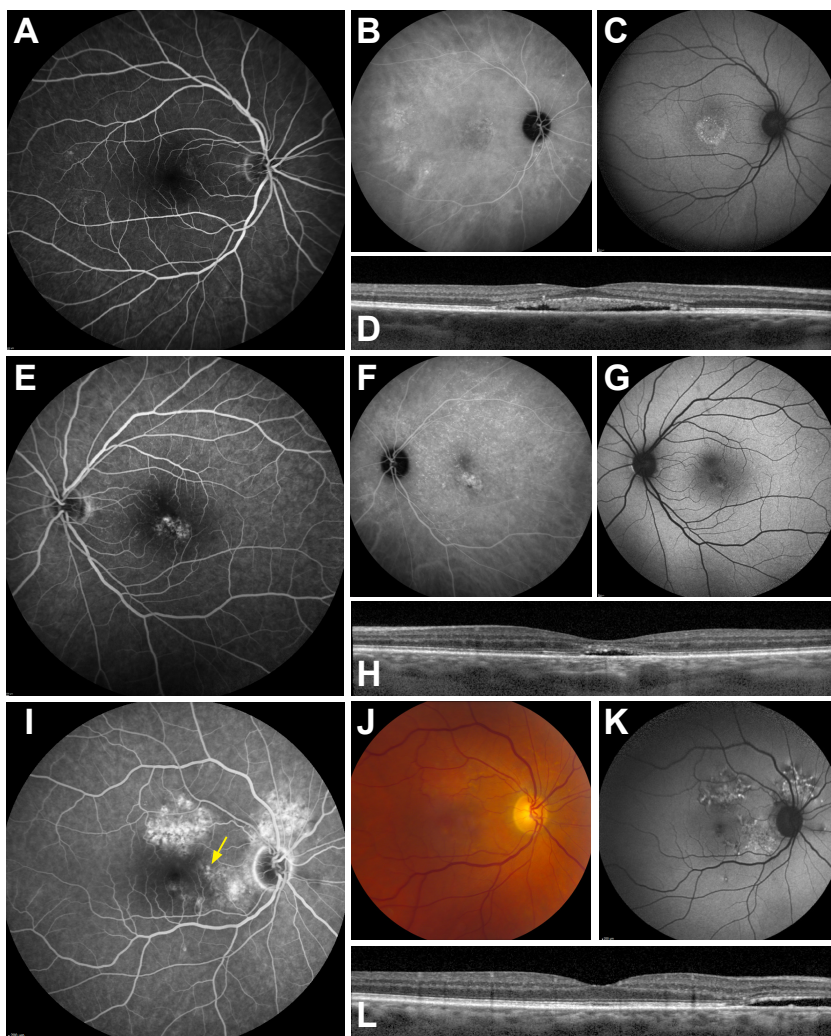


Figure 3: Spectrum of observed phenotypes in previously diagnosed patients with central serous chorioretinopathy. A-D. Multimodal imaging of the right eye of a 40-year-old female patient with unilateral chronic CSC. On intravenous FA (A), some hyperfluorescent changes, but no clear “hot spot” of leakage could be detected. Indocyanine green angiography (B) showed areas of hyperfluorescent changes, which were larger compared to changes on FA. Fundus autofluorescence (C) imaging revealed hyperautofluorescent changes in the fovea. The EDI-OCT (D) scan showed a foveal accumulation of SRF, with the presence of debris within the SRF. Moreover, dilated Haller vessels (pachyvessels) were present, and the subfoveal CT was $513\ \mu\text{m}$. E-H. Multimodal imaging of the left eye of a 55-year old male patient with unilateral chronic CSC. Intravenous FA (E) revealed foveal hyperfluorescent changes with a “hot spot” of leakage. Indocyanine green angiography imaging (F) showed diffuse zones of hyperfluorescent changes, without signs of either polypoidal choroidal vasculopathy or a choroidal neovascularisation. On FAF (G), mostly hypoautofluorescent changes were present foveally. On EDI-OCT (H), a foveal SRF accumulation and pachyvessels could be detected. Subfoveal CT was $420\ \mu\text{m}$. I-L. Multimodal imaging of the right eye of a 76-year-old male patient with bilateral chronic CSC. Oral FA (I) revealed multiple hyperfluorescent areas, with a possible “hot spot” of leakage nasally of the fovea (arrow). Fundus photography (J) showed multiple areas of alterations of the RPE. On FAF (K), both hyperautofluorescent and hypoautofluorescent changes were present. On EDI-OCT (L), SRF could be observed nasally of the fovea. A large pachyvessel could be found below the area of leakage on FA. Subfoveal CT was $245\ \mu\text{m}$, at that moment.

absence of SRF in 10 eyes (71%). The performed treatments and outcome of these treatments have been summarized in Table 2.

A wide range of phenotypical characteristics typical for CSC could be detected on multimodal imaging of the 39 previously diagnosed CSC patients (Figure 3). Either bilateral SRF on OCT or bilateral “hot spots” of leakage on FA was detected in 24 patients (62%). Until the moment of phenotyping for this study, a recurrence of SRF leakage had appeared in 15 patients (38%). On multimodal imaging, a unilateral epiretinal membrane could be observed in two patients, choroidal folds were found in both eyes of one patient, and cuticular drusen were observed bilaterally in another patient. Median CT in all 48 eyes of 25 patients, for whom EDI-OCT was available, was 323 μm (Q1: 275.5-Q3:365.5). Median CT in the 38 affected eyes was 325 μm (Q1:279-Q3:361), whereas median CT in the 10 unaffected eyes was 287 μm (Q1:247-Q3:372). Pachychoroid could be detected in 6 eyes of 5 patients.

Phenotyping of family members

Sixty-four family members (28 men and 36 women), in whom no previous CSC was diagnosed, were phenotyped for this study. Medical history of these subjects revealed hypertension in 22 subjects (34%) and other cardiovascular diseases in 9 other subjects (14%; total 48%). One subject had received two panretinal laser treatments because of a central retinal artery occlusion with secondary ischemia. Twelve of these screened family members (19%) reported steroid use within a year before screening. No link of the outcome of phenotyping with either steroid use or medical history could be detected.

Based on serous SRF on OCT and corresponding abnormalities on multimodal imaging, six screened family members (9%) were diagnosed with CSC. Mean age at diagnosis of these 3 men and 3 women was 48 ± 15 years (Table 1) and median ETDRS BCVA was 84 letters (Snellen equivalent: 20/21; Q1:80-Q3:85) in the 7 affected eyes and 85 (Snellen equivalent: 20/20; Q1:71-Q3:89) in the 5 unaffected eyes (Table 3). Moreover, multimodal imaging showed RPE changes suggestive of CSC in 27 screened subjects (42%). The mean age at diagnosis was 53 ± 9 years for these 27 suggestive CSC subjects, and 44% of these subjects were male (Table 1).

The extent of RPE changes on fundus photography and corresponding either hyperfluorescent or hypofluorescent changes on FAF imaging in suggestive CSC patients were variable. Abnormalities on FA ranged from subtle hyperfluorescent RPE changes to diffuse areas of hyperfluorescence, but without the presence of SRF leakage (Figure 4). On OCT, either subtle or obvious RPE alterations were present, but without SRF (Figure 4). Choroidal thickness was variable among the suggestive patients with CSC, but in only one of these patients bilateral pachychoroid was present.

Within the group of screened family members without a previous history of CSC, multimodal imaging revealed bilateral cuticular drusen in two subjects, bilateral choroidal folds in two subjects, a unilateral macular epiretinal membrane in three subjects, a macular cystoid lesion in one eye, a lamellar macular hole in one eye, and findings characteristic of a previous vascular occlusion in one eye.

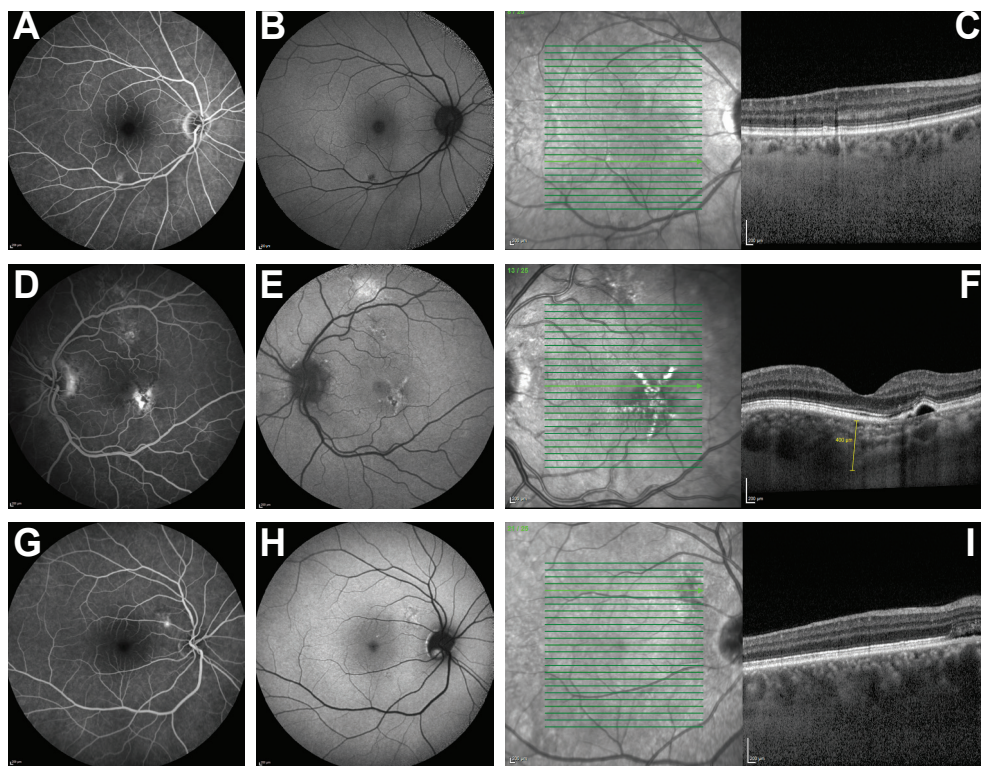


Figure 4: Spectrum of observed phenotypes in family members of patients with CSC. Various phenotypes resembling CSC were observed in family members of patients with CSC. A-C. Multimodal imaging of the right eye of a 61-year-old female patient. On oral FA (A) and FAF (B) imaging, subtle hyper (auto) fluorescent changes suggestive of CSC could be detected inferior of the fovea. Moreover, OCT (C) showed subtle RPE abnormalities, and a subfoveal CT of $247 \mu\text{m}$. Similar hyperfluorescent lesions on FA and FAF could be detected in the left eye of this patient. D-F. Multimodal imaging of the left eye of a 60-year-old female patient. On oral FA (D) diffuse areas of hyperfluorescence were found without an obvious “hot spot” of leakage, which was considered to be suggestive of CSC. Both hyperautofluorescent and hypoautofluorescent changes were present on FAF (E), with an area of likely prior SRF superiorly. On OCT (F), a detachment of the RPE could be detected temporally of the fovea, without the presence of serous SRF. Subfoveal CT was $400 \mu\text{m}$ in this eye, and dilated Haller layer (pachyvessels) and bilateral choroidal folds were also present in this patient. G-I. Multimodal imaging of the right eye of a 60-year-old female patient. On oral FA (G), a characteristic “hot spot” of leakage superonasally of the fovea could be detected, with a surrounding area of hyper (auto) fluorescence on FA and FAF (H). On OCT (I), unilateral SRF could be detected superonasally of the fovea. A pachyvessel was present below the SRF, and the subfoveal CT in this eye was $528 \mu\text{m}$.

Choroidal thickness in affected and suspected patients and unaffected individuals

Choroidal thickness data of CSC affected, CSC suggestive, and unaffected eyes in the three subgroups were combined (Table 3). Choroidal thickness measurements on EDI-OCT images of 44 affected eyes of patients in whom CSC had been diagnosed showed a median subfoveal CT of $325 \mu\text{m}$ (Q1:282-Q3:360), whereas 41 eyes of suggestive CSC patients with both detected abnormalities and available EDI-OCTs, had a median CT of $272 \mu\text{m}$ (Q1:237-Q3:337), and the 84 unaffected eyes showed a CT of $280 \mu\text{m}$ (Q1:240-Q3:335). There was a significant difference in CT between affected and

both the unaffected and suggestive CSC eyes ($P < 0.05$). No significant difference was found between unaffected and suggestive CSC eyes (Table 3). Pachychoroid could be detected in 18 eyes in 12 family members in both the affected (6 patients, eight eyes), suggestive CSC (2 patients, four eyes), and unaffected (4 family members, six eyes) subgroups.

Follow-up of individuals who received ophthalmological phenotyping in 1994-1995

Out of the 30 patients who were previously phenotyped at the Department of Ophthalmology of the LUMC,¹⁵ 24 patients could be included for new imaging, which was performed at a median follow-up of 20.8 years (Q1: 20.7-Q3:21.2) after the original phenotyping. In 7 out of these 24 patients, the CSC diagnosis had already been established at baseline. In 4 of the remaining 17 patients (24%), the outcome of new phenotyping led to a diagnosis of more severe abnormalities. Namely, 2 patients out of 13 patients (15%) in whom RPE alterations were previously observed proved to have active CSC at the moment of new imaging. Moreover, in one out of four previously non-affected subjects (25%), findings suggestive of CSC were found during our phenotyping, and in another nonaffected subject (25%), active CSC was found.

Discussion

Based on the results of this extensive phenotyping, occurrence of CSC and RPE abnormalities suggestive of CSC appears to be relatively common in families with CSC. Family members of patients with CSC could therefore be at risk to develop CSC. Within this study, SRF on OCT and/or “hot spots” of leakage on FA could be detected in 45 of the 103 included subjects (44%) and a CSC suspicion in 27 others (26%). In 31 subjects (30%), no signs of CSC were present. The 45 individuals affected by CSC included 6 previously undiagnosed family members of diagnosed CSC patients. Moreover, in 24% of patients who were invited for a long-term follow-up visit after phenotyping more than 2 decades ago, either progression or new occurrence of CSC could be detected.

Active CSC was found only in a minority of patients in whom findings characteristic for CSC could be detected during ophthalmological screening. Based on the outcome of current ophthalmological screening, none of the screened patients in whom CSC was newly diagnosed required treatment for active CSC. In this study, the range of ophthalmological abnormalities in subjects who were diagnosed as having suggestive CSC was wide (Figure 4) and the clinical relevance of RPE abnormalities suggestive of CSC remains to be elucidated because similar findings have been described not to be present in healthy controls.²⁰ Therefore, the exact relevance of the suggestive CSC diagnosis is unclear, as is the exact overlap between suggestive CSC and CSC.⁵ The diagnosis of suggestive CSC has features similar to the previously described pachychoroid pigment epitheliopathy, for which either unilateral or bilateral reduced fundus tessellation with overlying RPE changes on fundoscopy and abnormalities on FAF have been described.⁵ However, in contrast with pachychoroid pigment epitheliopathy, an increased subfoveal CT was not seen in all patients with suggestive CSC.⁵ In this study, either unilateral or bilateral pachychoroid could be detected in only 6 of the 61 screened family members (10%), for whom EDI-OCT imaging was available. Pachychoroid, using a CT cut-off point of 395 μm , occurred

Table 3. Ophthalmologic characteristics of different subgroups

	Known CSC	New CSC	Suggestive CSC	Unaffected	Total
Affected subjects	39	6	27	31	103
Amblyopia	3	-	1	1	5
Bilateral choroidal folds	1	1	1	-	3
Bilateral cuticular drusen	1	-	2	-	3
Central retinal artery occlusion	-	-	1	-	1
Glaucoma	1	-	-	-	1
Ocular hypertension	1	1	-	-	2
Unilateral epiretinal membrane	2	1	2	-	5
Unaffected eyes					
No. of eyes with available BCVA/CT	15/10	5/4	10/10 ^x	62/60	92/84
Median BCVA (ETDRS) (Q1-Q3)	85 (81-89)	85 (71-89)	87 (73-90)	89 (82-91)	88 (81-90)
Median BCVA (Snellen equivalent)	20/20	20/20	20/18	20/17	20/17
Median CT in μm (Q1-Q3)	287 (247-372)	339 (268-386)	270 (252-332)	277 (227-334)	280 (240-335)
Suspected eyes					
No. of eyes with available BCVA/CT	-	-	43/41	-	43/41
Median BCVA (ETDRS) (Q1-Q3)	-	-	89 (85-91)	-	89 (85-91) ^{ns}
Median BCVA (Snellen equivalent)	-	-	20/17	-	20/17
Median CT in μm (Q1-Q3)	-	-	272 (237-337)	-	272 (237-337) ^{ns}
Affected eyes					
No. of eyes with available BCVA/CT	63/38	7/6	-	-	70/44
Median BCVA (ETDRS) (Q1-Q3)	80 (70-85)	84 (80-85)	-	-	80 (70-85)**
Median BCVA (Snellen equivalent)	20/25	20/21	-	-	20/25
Median CT in μm (Q1-Q3)	325 (279-361)	320 (296-376)	-	-	325 (282-360)*

ns: non-significant compared to unaffected eyes

^x: one eye was excluded due to poor visual acuity after a central retinal artery occlusion

**P value compared with unaffected eyes: < 0.001

* P value compared with unaffected eyes: < 0.05

both in CSC affected, CSC suggestive, and unaffected screened family members.¹⁸ Nevertheless, the CT of affected eyes of patients with CSC was significantly higher compared to all eyes of unaffected individuals (Supplementary Figure 2). More recently, both the maximal CT and pathologically dilated outer choroidal vessels (in the Haller layer) have been described to be even more typical than CT for various diseases within the pachychoroid spectrum including pachychoroid pigment epitheliopathy, CSC, and pachychoroid neovasculopathy.^{21,22} These abnormalities in the choroid, together with a loss in choriocapillary volume, could also be found in our patients (Figures 3 and 4), addressing the importance of the previously reported pachychoroid spectrum.^{5,21,22}

The underlying pathogenesis of familial clustering of CSC and CSC-like RPE abnormalities remains obscure. Age at CSC diagnosis, sex, steroid use, response to different CSC treatments, medical history, and presence of choroidal neovascularisation do not differ when comparing the current familial CSC cohort with the occurrence of these factors in previously described nonfamilial CSC patient cohorts.^{1-3,23-26} This leads to a challenge in recognizing family members who may be at risk for developing active CSC because we did not find phenotypic characteristics in the probands that could be indicative of a familial occurrence of CSC. However, both the percentage of recurrent and/or bilateral CSC among the probands in the CSC families in our study was relatively high, compared with sporadic CSC patients in available literature.²⁷ The presence of these signs of a relatively severe CSC could lead to a higher risk for family members to develop CSC.

Central serous chorioretinopathy is believed to be a complex, multifactorial disease, in which both environmental and genetic factors play a role, and its inheritance pattern is currently unknown. In the current study, most of the affected family members involved sibships, but in some families, CSC could be detected in several generations. In most families, the mode of inheritance appears to be autosomal dominant (Supplementary Figure 1). The possible autosomal-dominant inheritance pattern is in line with a recent study on the familial occurrence of pachychoroid, in which this pattern has been suggested.¹⁸ However, the lack of large families with data from multiple generations could lead to an unreliable determination of the inheritance pattern. Ophthalmologic phenotyping in multiple generations was only performed when members of the younger generation reported a possible history of CSC or were within the expected age range to develop CSC. Our findings could, therefore, be an underestimation of the actual occurrence of clinical findings characteristic for CSC in multiple generations.

Based on the familial occurrence of CSC, it has already been suggested that genetic factors might play an important role in CSC. This hypothesis is supported by studies on common genetic variants (single nucleotide polymorphisms) that have previously been associated with sporadic CSC.⁹⁻¹¹ Single nucleotide polymorphisms in the *ARMS2*, *CDH5*, and *CFH* genes and copy number variations in the *C4B* gene involved in extracellular matrix, cell adhesion, and complement system, respectively, have already been associated with CSC.⁹⁻¹¹ The detection of (suggestive) CSC in 52% of family members of CSC patients reported in this study may indicate that there is also a role for more severe rare genetic variants in CSC. This has already been described in families with age-related macular degeneration, a disease that shows overlapping features with CSC.²⁸

In conclusion, CSC and lesions that could be predisposing to CSC can cluster in families, which may indicate that family members of CSC patients are at risk to develop the disease. However, many of these individuals may remain asymptomatic. During a follow-up at which multimodal imaging is performed, progression of disease can occur in a noteworthy number of patients. Comparing next generation genetic sequencing data of both affected and unaffected family members, ideally from multiple generations, may shed a light on which genetic factors are involved in familial CSC. In future studies, a combination of extensive phenotyping and in-depth genotyping can provide new clues on the etiology of CSC.

Footnotes and Financial Disclosures

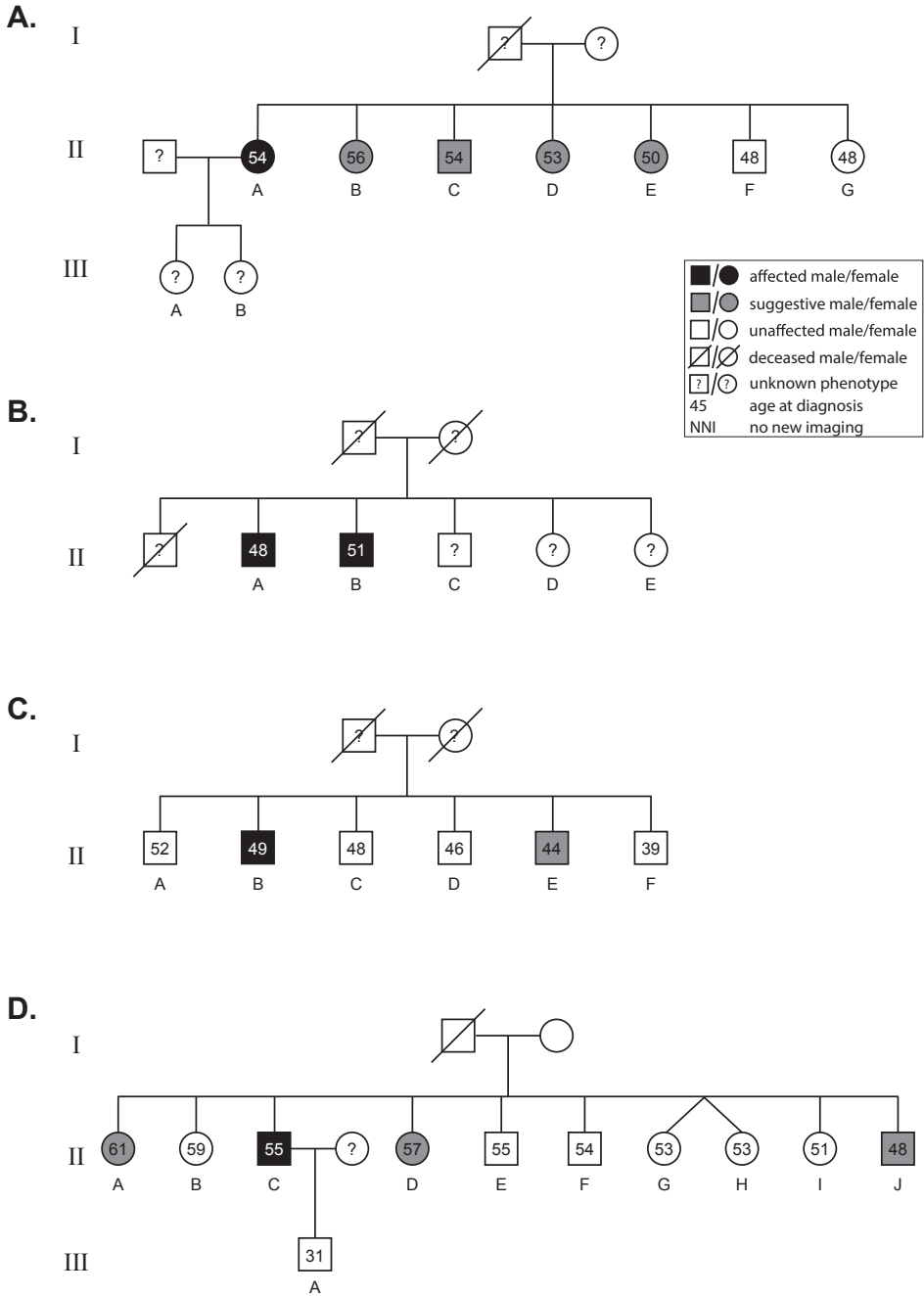
This research was supported by the following foundations: MaculaFonds, Retina Netherlands, BlindenPenning, and Landelijke Stichting voor Blinden en Slechtzienden, that contributed through UitZicht, as well as Rotterdamse Stichting Blindenbelangen, Haagse Stichting Blindenhulp, ZonMw VENI Grant, and Gisela Thier Fellowship of Leiden University (CJFB), and Macula Vision Research Foundation, Stichting Nederlands Oogheelkundig Onderzoek, Stichting Blindenhulp, Stichting A.F. Deutman Oogheelkunde Researchfonds, Radboud Institute of Molecular Life Sciences, Gelderse Blindenstichting, and Nijmeegse Oogonderzoek Stichting (AidH&EKdJ). The funding organizations provided unrestricted grants.

No conflicting relationship exists for any author.

Key words: central serous chorioretinopathy; disease progression; familial; genetics

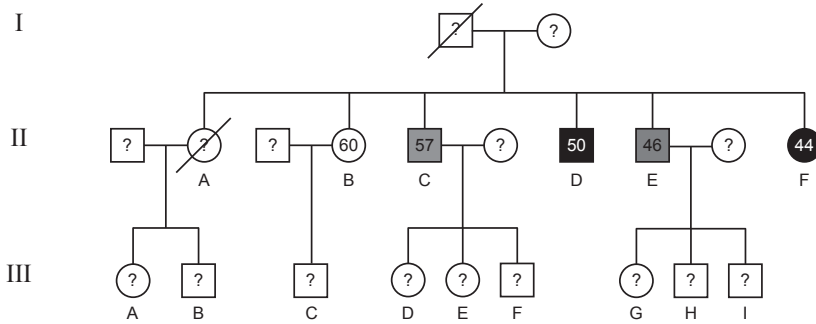
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Supplementary Files

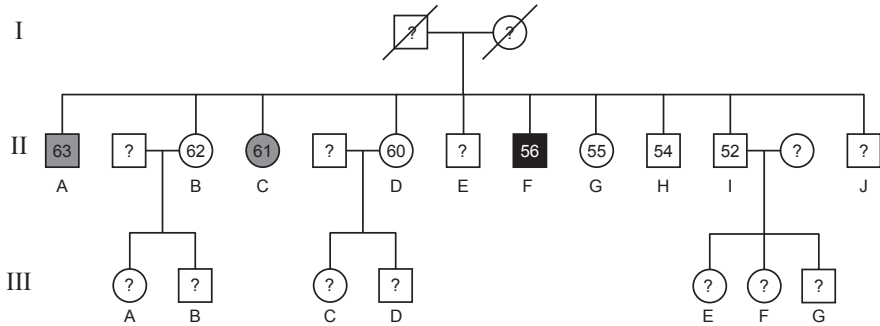


Supplementary Figure 1. Pedigrees of the families who have been invited for ophthalmological phenotyping

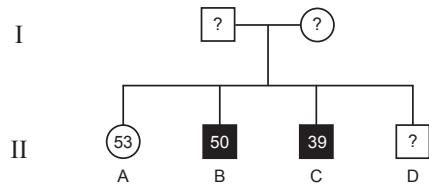
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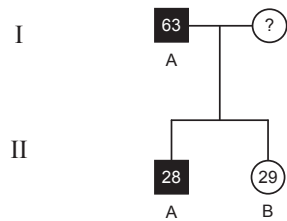
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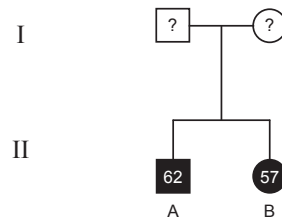
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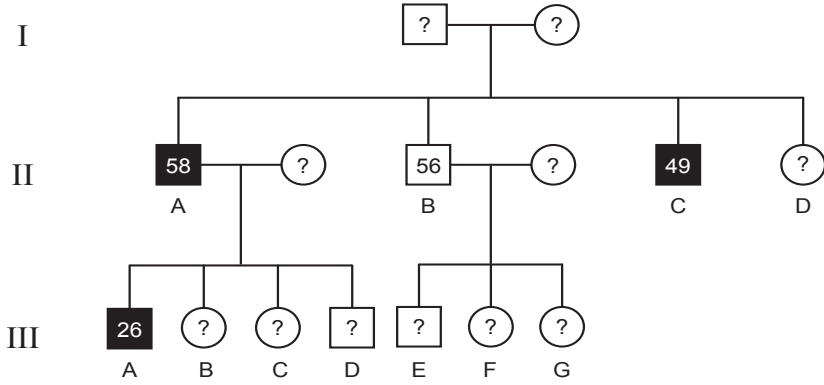
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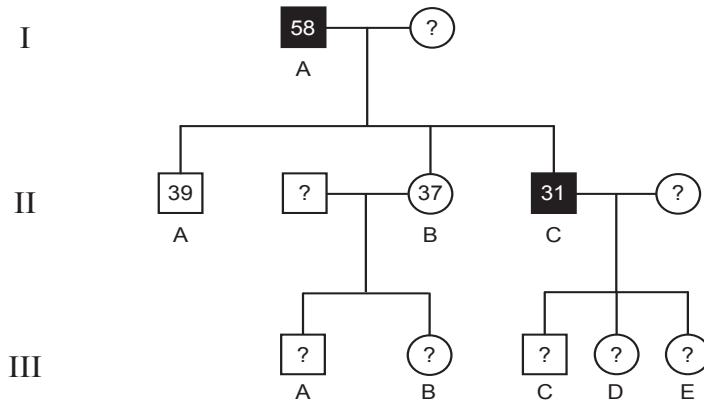
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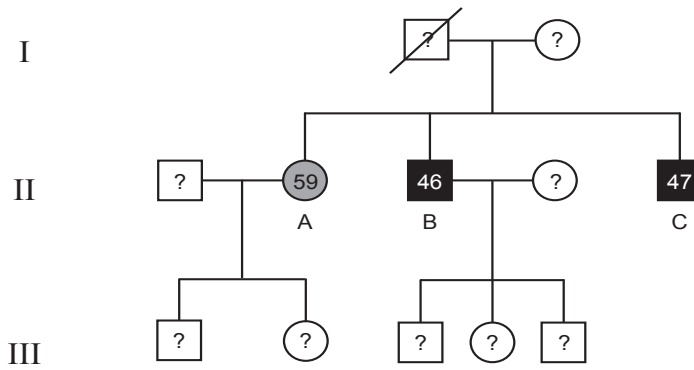


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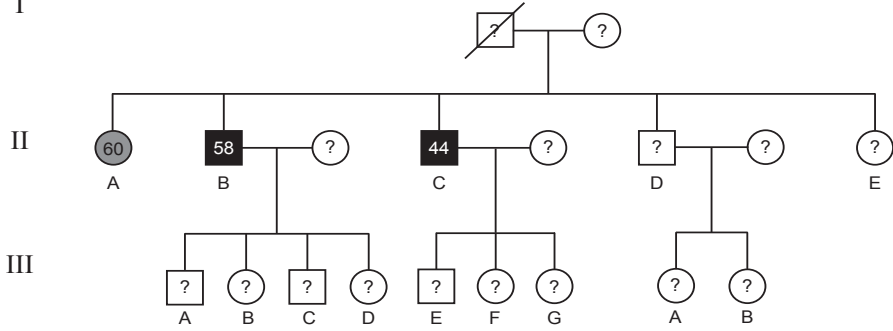


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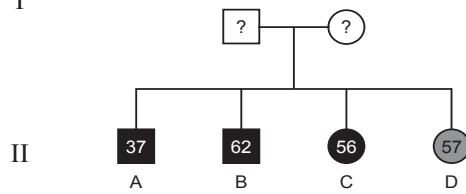
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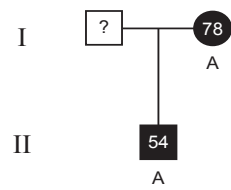
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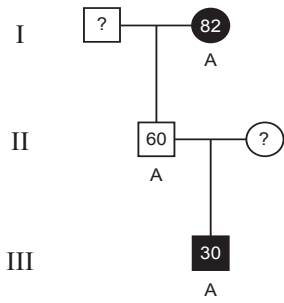
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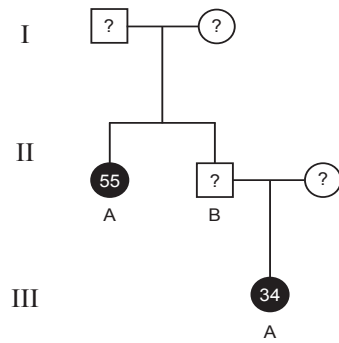
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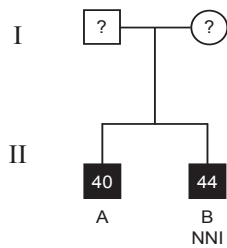
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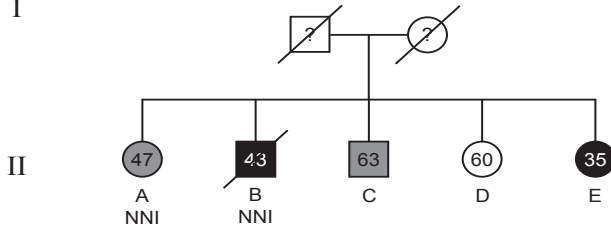
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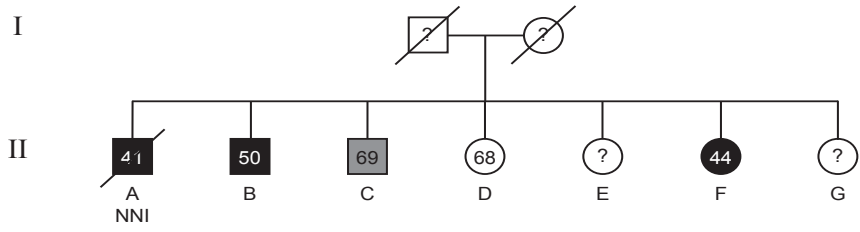
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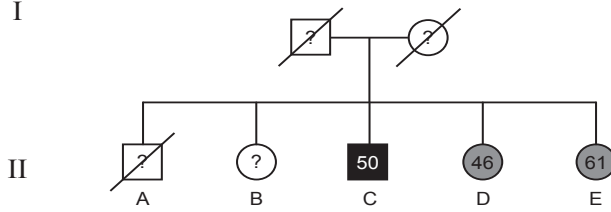
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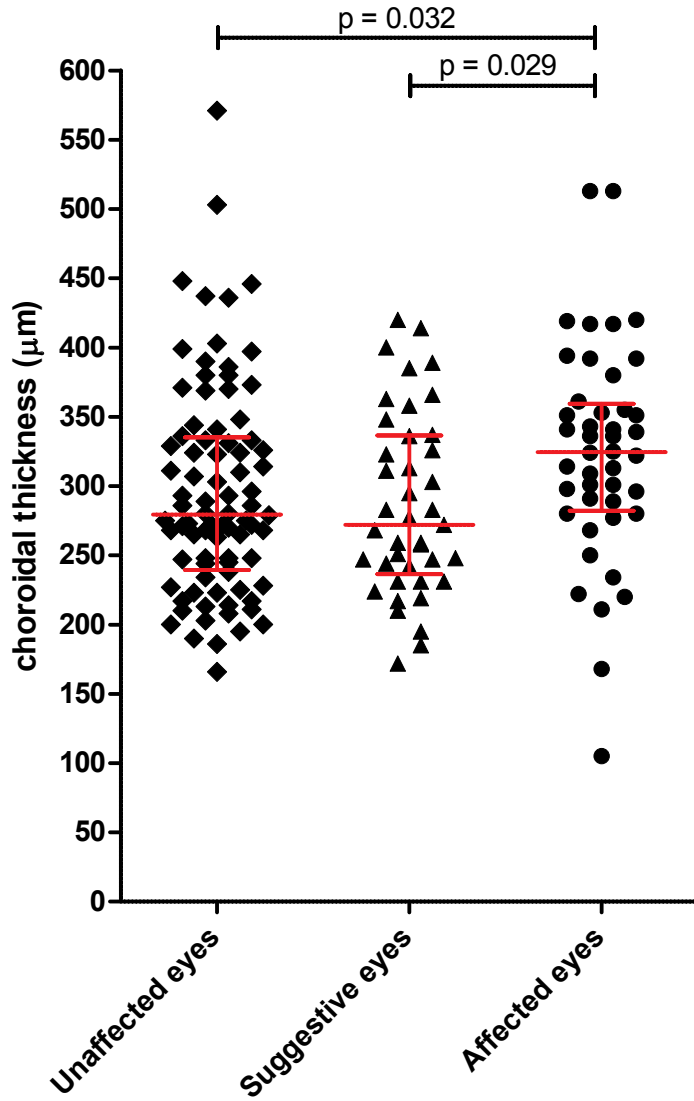


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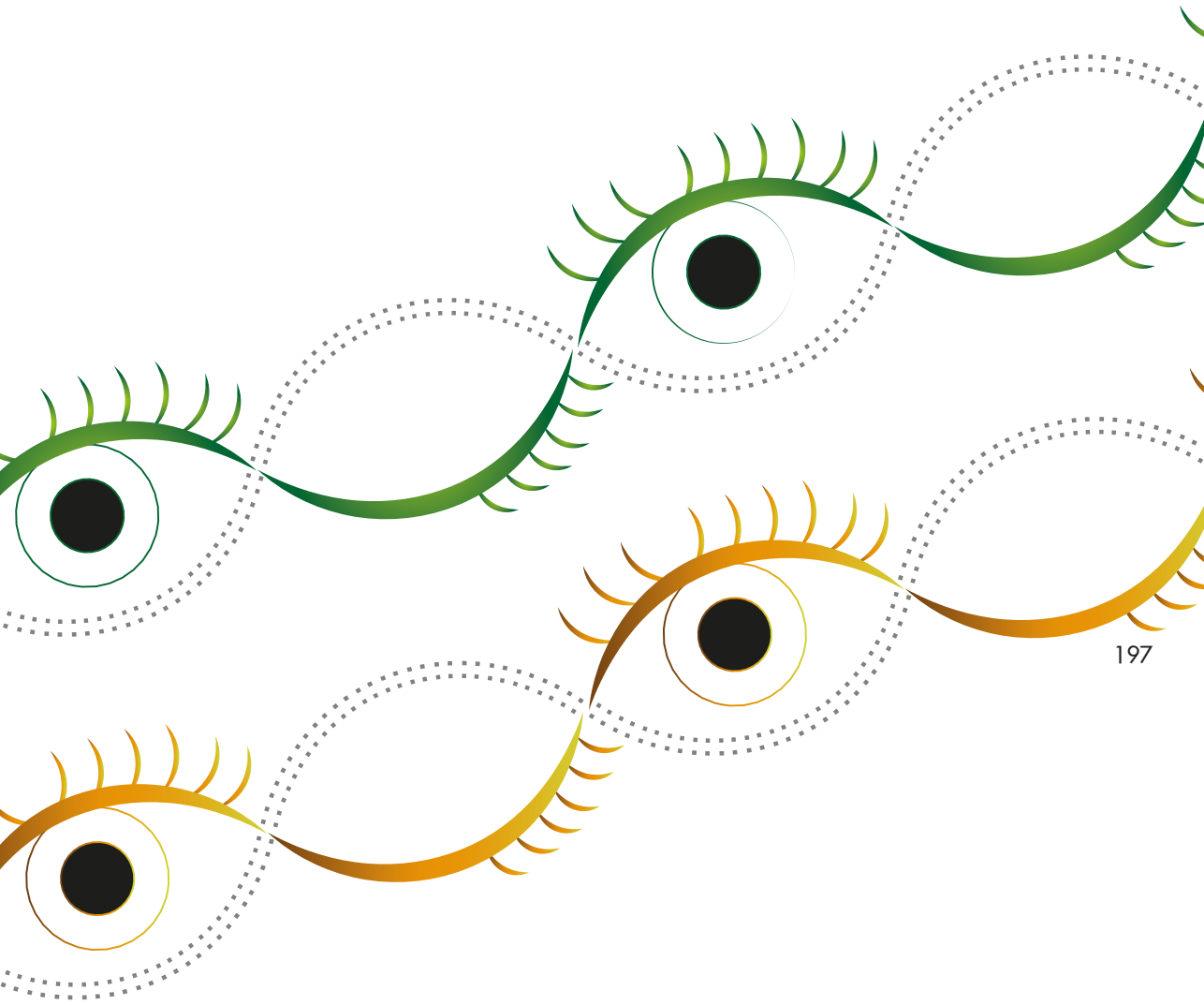


Supplementary Figure 2: Measurement of choroidal thickness in unaffected, central serous chorioretinopathy suggestive, and affected eyes. Medians with interquartile range are depicted in red.

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4.2 Exome sequencing in familial chronic central serous chorioretinopathy

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Abstract

Central serous chorioretinopathy (CSC) is a chorioretinal disease characterized by fluid accumulation between the neuroretina and retinal pigment epithelium with unknown etiology. Family studies have suggested that CSC might be an inherited disease with an autosomal dominant inheritance pattern. We therefore performed exome sequencing on 72 individuals of 18 families with CSC, and determined whether rare genetic variants (minor allele frequency <1%) segregated in two or more families with CSC. In addition, familial gene-burden analysis was performed and compared to gene-based results of an exome sequencing study on a large sporadic CSC case-control cohort. In total, 11 variants were found to segregate in two out of 18 families. One of these variants, c.4145C>T; p.T1382I (rs61758735) in the *PTPRB* gene, was also associated with CSC in the case-control cohort ($P=0.008965$). Additionally, in 28 genes two or more different heterozygous variants were found to segregate in two or more families, but none of these genes showed consistent associations in both the family gene-burden results and gene-burden analysis in the case-control cohort. In this study, we identified possible candidate genes for familial CSC. However, Mendelian inheritance of variants in one or a limited number of genes can be excluded based on this study. Instead, familial CSC may be a heterogeneous Mendelian disease caused by variants in many different genes, or alternatively CSC may represent a complex disease to which both environmental factors and genetics contribute.

Introduction

In central serous chorioretinopathy (CSC), choroidal congestion, thickening, and hyperpermeability have been suggested to cause leakage through the retinal pigment epithelium (RPE). Subsequently, a neuroretinal detachment occurs due to the accumulation of serous subretinal fluid.¹⁻⁵ The exact etiology of CSC is still unclear, but male gender, administration of exogenous corticoids have been described to be the most pronounced risk factors for CSC.⁶⁻⁸ Other risk factors include endogenous hypercortisolism, stress and pregnancy.^{2,9,10} Moreover, genetic variants that confer risk or are protective for CSC have been identified by genetic association studies in case-control cohorts.¹¹⁻¹⁶

Although familial occurrence of CSC appears to be rare, several reports on familial CSC and the occurrence of CSC in multiple generations within a single family have been published, pointing to a potential role for genetic factors in familial CSC.¹⁷⁻²⁰ A Mendelian inheritance of CSC has been proposed previously based on observations that at least two family members proved to have findings characteristic for CSC in 52% of 27 families.¹⁸ In some families affected individuals were present in multiple generations, suggesting an autosomal dominant mode of inheritance of familial CSC.²⁰ Additionally, in 50% of eyes from screened family members of CSC patients a thickened choroid (pachychoroid) of more than 395 μ m was detected, which has been described to be the underlying choroidal abnormality in various diseases that are part of the pachychoroid spectrum.²¹ However, thus far no genetic studies on familial CSC have been conducted.

Whole exome sequencing has proven to be a powerful tool to identify novel disease-associated genes and gene variants in many disorders.^{22,23} Exons are presumed to harbor about 85% of disease-causing mutations, making them a primary target to search for disease-associated variants in CSC families.²⁴ Therefore in this study, we performed exome sequencing on 72 individuals of 18 families in which multiple members were found to have CSC, in order to determine whether Mendelian inheritance of rare genetic variants causes familial CSC.

Materials and methods

Editorial Policies and Ethical Considerations

Written informed consent for the enrollment was obtained from all subjects. The study adhered to the tenets of the Declaration of Helsinki. Approval of the institutional review board and the ethics committee was obtained for all centers involved.

Subject selection

In this multicenter study, 72 subjects from 18 families, including patients with CSC and unaffected family members, visited either the Department of Ophthalmology of the Radboud university medical center ([Radboudumc] Nijmegen, the Netherlands) or the Leiden University Medical Center ([LUMC] Leiden, the Netherlands). The participants were recruited at the outpatient clinic of the participating hospitals, after the proband had reported a family history of CSC. The majority of the individuals were included in our previously published study on the phenotypic characteristics of familial CSC and were divided in the following groups: "Affected with CSC", "Suggestive of CSC" or "Unaffected", using the criteria described before.²⁰

Briefly, subjects were categorized as having CSC when serous fluid was detected on OCT and when one or more 'hot spots' of leakage or diffuse leakage was present in combination with irregular RPE window defects on fluorescein angiography. Patients were excluded if signs of either polypoidal choroidal vasculopathy or age-related macular degeneration (AMD), or other atypical findings were present. Suggestive CSC was characterized by RPE alterations typical for CSC, but without the presence of either subretinal fluid or 'hot spots' of leakage on fluorescein angiography.²⁰ Unaffected individuals showed no abnormalities on any of the modalities using multimodal imaging.

Exome sequencing

Library preparation of the 52 family members of 12 families collected at the Radboudumc, Nijmegen, was performed with the SureSelect^{XT} target enrichment system for Illumina paired-end multiplex sequencing according to manufacturer's instructions (Version B4, August 2015, Agilent Technologies). Finished libraries were sent to the Department of Genetics of Maastricht University Medical Center+, Maastricht, the Netherlands, where sequencing was performed with 8 samples per lane using an Illumina HiSeq2000 with 2*100bp chemistry, together with a large cohort of 269 sporadic CSC patients (Schellevis et al. 2018 submitted).

The 20 samples of the 6 families collected in Leiden University Medical Center, Leiden, the Netherlands were sent to GenomeScan BV, Leiden, for sequencing. For these samples the Agilent SureSelect V5 enrichment kit was used and sequencing was performed with 2*125bp chemistry on the HiSeq2500.

Variant calling and recalibration

Data of all individuals were processed according to the Genome-Analysis-Toolkit (GATK) best practices (v3.8) together with the case-control cohort consisting of 269 sporadic cCSC patients and 1586 population controls (Schellevis et al. 2018 submitted) to improve variant calling. Briefly, BAM to FastQC extraction was performed with Picard-tools (v 1.90), duplicate reads were marked with Picard-tools, and reads were aligned to the reference genome (GRCh37.p5 with alternate haplotypes excluded) using BWA-MEM (version v.0.7.12), as described before (Schellevis et al. 2018 submitted). Base recalibration was performed and subsequent variant calling was performed with the HaplotypeCaller algorithm. All GVCFs were merged together, and joint genotyping was performed on the dataset together.

Variant recalibration was performed on the entire dataset with GATK using the recommended settings,²⁵ as described before. Genetic variants located in low complexity regions of the genome were removed.²⁶ Multiallelic variants were extracted with VCFtools (v0.1.13) and split using the splitMultiallelic and LeftAlignandTrimVariants option in GATK (v3.8). Remaining variants were tested for Hardy-Weinberg equilibrium ($P=1 \times 10^{-8}$). Variants from the adult-onset genes captured in the American College of Medical Genetics and Genomics recommendations for incidental findings (*BRCA1*, *BRCA2*, *MLH1*, *MSH2*, *MSH6*, *PMS2*, *MUTYH*) were removed to reduce the risk of secondary findings.^{27,28} Finally, family members were extracted from the dataset and only variants with a minor allele count ≥ 1 in the combined family file were retained. Data was annotated with Tabanno (<https://github.com/zhanxw/anno>) and Annovar.²⁹

Variant filtering

Variants were filtered based on a minor allele frequency (MAF) of $\leq 1\%$ or $\geq 99\%$ in the following populations: 1000Genomes_all, 1000Genomes_European, Exac_All, Exac_NFE, Exac_FIN, and esp6500siv2_all. Additionally, variants with MAF $\geq 1\%$ in the 1586 controls of the sporadic CSC case-control cohort were removed. Variants that were annotated by Annovar to be present in the exonic or splice site regions were retained and synonymous variants were excluded. Remaining variants included: frameshift insertions and deletions (INDELs), nonframeshift INDELs, nonsynonymous variants, stop gain or loss variants, and variants with unknown effects. All variants present in one or more unaffected individuals in any of the families were removed, with the exception of the unaffected individual of *Family 14*, because based on the pedigree structure (Supplementary Figure 1) reduced penetrance appeared to be present in this family. Only variants with a CADD score above 20, corresponding to the 1% most deleterious variants of the human genome, or with an unknown CADD score in case of INDELs, were retained.

Next, for each family segregation analysis of variants was performed, where variants were retained if they were present in all affected individuals of the family and not present in unaffected individuals. No filtering was performed for individuals with suggestive CSC. Variants that segregated in two or more families, or genes that contained multiple variants that segregated in two or more families were retained for further evaluation. Familial gene-burden associations were calculated with RareIBD

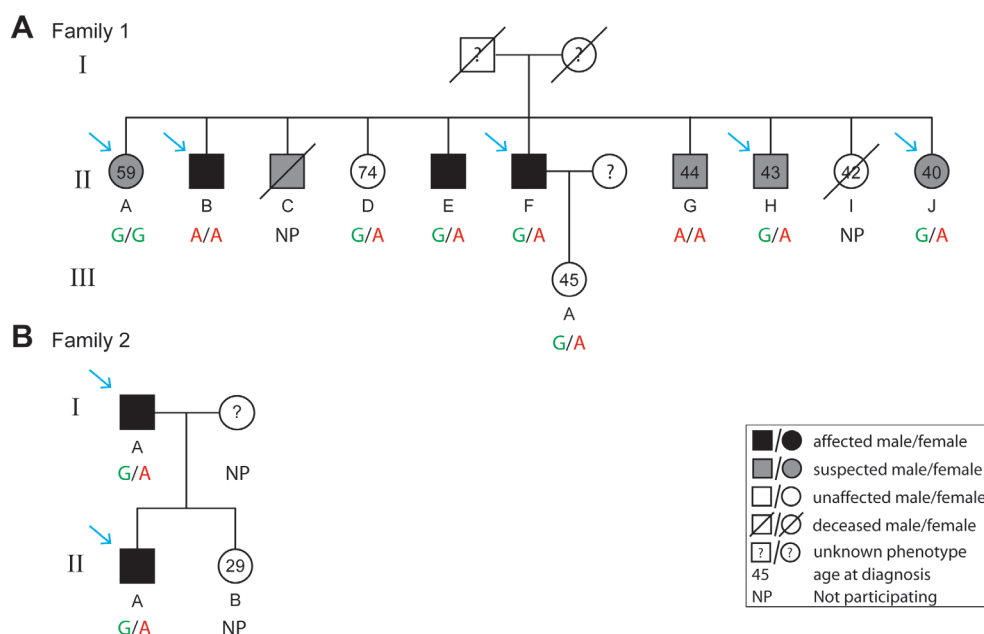


Figure 1: Segregation analysis of rs61758735G>A in *PTPRB* for Family 1 and Family 2. Genotypes for rs61758735 are depicted below each individual in the pedigree. Individuals whom were sent in for exome sequencing are indicated with a blue arrow, while other individuals for which DNA was available were sent in for Sanger sequencing. Individual B11B did not want to participate in the study, and therefore could not be sequenced.

for those genes that carried multiple segregating variants in two or more families.³⁰ The region encompassing the c.4145C>T; p.T1382I (rs61758735) variant in the *PTPRB* gene was amplified in additional family members of *Family 1* using AmpliTaq DNA polymerase with the following PCR program: 5 min. at 95°C, 10 cycles of touchdown starting at 62°C for 45 sec. and lowering the annealing temperature 0.5°C each cycle, followed by 25 cycles of an annealing temperature of 57°C, all these cycles started with 30 sec. at 95°C and ended 45 sec. at 72°C. The PCR was concluded with 5 min. at 72°C and the following primers were used forward primer: AGCCTTTGAGCAGCTTTTTC and reverse primer: TGATGCTAGTGCCCCATAAG. The PCR product was analyzed by Sanger sequencing by the core sequencing facility at the Department of Human Genetics of the Radboudumc.

Results

In this exome sequencing study on familial CSC we included 72 individuals from 18 different families. Out of these 72 individuals, 33 subjects were affected with CSC, 18 had characteristics suggestive of CSC, and 21 were unaffected (Table 1, Figure 1 and Supplementary Figure 1 for pedigrees). After variant filtering based on MAF ($\leq 1\%$), CADD score (≥ 20), absence in unaffected individuals, and protein-altering function (frameshift INDELs, nonframeshift INDELs, nonsynonymous, stop gain/loss variants and variants with unknown effects), the dataset contained 2806 variants present in 2368 genes.

Segregation analysis was performed for all 18 families, retaining only variants that were present in the affected individuals and absent in unaffected individuals. The average number of segregating variants in each family was 44 and ranged from three to 124 (Table 1). In four families two segregating heterozygous rare variants in the same gene were observed. *Family 12* carried two variants in *KCNMA1*, *Family 7* in *RBPJL*, *Family 8* in *SLC26A10*, and *Family 1* in *SP9* (Supplementary Table 1). The entire list of segregating variants for each family is available in Supplementary Table 2.

Variants that segregated in two or more families were retained for further evaluation. In total, 11 rare variants were found to segregate in two families, of which one variant in the *PTPRB* gene was found homozygous in one individual, while the remaining variants in the *SETD2*, *PWP1*, *ABCA9*, *AT2B2*, *ZFAND4*, *MROH5*, *ZAN*, *SHISA6*, *DCP1A*, and *PPM1E* genes were heterozygous (Table 2). Expression of these genes in the retina and RPE was evaluated using the Eye Integration Database (Supplementary Figure 1).³¹ The single variant association results of these variants were extracted from the sporadic CSC case-control dataset (Schellevis et al. 2018 submitted), and the variant in the *PTPRB* gene (c.4145C>T; p.T1382I, rs61758735) was significantly associated with CSC in the case-control cohort ($P=0.008965$, Odds Ratio=2.83, 95% Confidence Interval=1.34-5.97). Extended segregating analysis of the *PTPRB* variant in additional available family members of *Family 1* revealed that two individuals carried the variant homozygously (one individual is an affected subject and one is a subject with findings suggestive of CSC), six individuals carried the variant heterozygously (two affected individuals, two individuals with findings suggestive of CSC, and two unaffected individuals) and one individual did not carry the variant (this individual had characteristics suggestive of CSC) (Figure 1A). Since two unaffected individuals (of which one individual was 74 years at examination) carried the variant heterozygously,

complete segregation of this variant with the disease in this family was not observed. For *Family 2* no additional individuals were available for extended segregation analysis (Figure 1B).

As a next step, genes that contained multiple variants segregating in two or more families were evaluated. In 28 genes, we observed two or more different heterozygous variants that segregated in two or more families, including in two genes known to cause a retinal phenotype (*ABCA4* and *VCAN*, full list of variants in Table 3). Expression of these genes in the retina and RPE was evaluated using the Eye Integration Database, and all genes with the exception of *CDMD2* and *TCF25* showed moderate to high expression in the RPE or retina (Supplementary Figure 2).³¹ For all 28 genes the results of the gene-based analysis (Burden, SKAT, and SKAT-O) were extracted from the case-control analysis (Schellevis et al. 2018 submitted). Also, a family gene-burden analysis was performed with RareIBD including all rare variants found in all families in the 28 genes. Multiple genes were nominally associated with CSC, but no significant associations were observed in either of the tests after correction for multiple testing for 28 genes (Table 4). Additionally, no genes showed consistent associations in both the case-control cohort and the family cohort.

Table 1. Results of per family segregation

Family	No. Affected individuals	No. Suggestive individuals	No. Unaffected individuals	No. Variants left after segregation
Family 1	2	3	-	75
Family 2	2	-	-	69
Family 3	1	3	2	36
Family 4	1	1	4	14
Family 5	1	2	4	3
Family 6	2	2	1	37
Family 7	1	2	3	17
Family 8	3	-	1	17
Family 9	2	-	2	12
Family 10	2	1	-	58
Family 11	2	1	-	79
Family 12	3	-	-	29
Family 13	2	-	-	71
Family 14*	2	-	1	37
Family 15	2	-	-	60
Family 16	2	1	1	21
Family 17	1	1	-	124
Family 18	2	1	2	29

* Possibly reduced penetrance in this family

Table 2. Rare segregating variants observed in two families

Family	Chr	Position	Variant	Ref	Alt	Gene	Accession number	Codon change	Protein Change	P-value (Schellevis et al. 2018 submitted)	CADD score
Family 7 + 14	3	47162897	rs114719990	T	C	SETD2	NM_014159	c.A3229G	p.T1077A	0.856	23.6
Family 13 + 17	12	108105893	rs11547909	G	A	PWP1	NM_007062	c.G1402A	p.E468K	0.801	24
Family 2 + 16	17	67013913	rs143651746	A	G	ABCA9	NM_080283	c.T2785C	p.F929L	0.167	22.4
Family 1 + 17	3	10413597	rs144118750	C	T	ATP2B2	NM_001683	c.G1420A	p.V474I	0.092	21.8
Family 9 + 10	10	46122195	rs144142701	A	T	ZFAND4	NM_174890	c.T1076A	p.L359H	0.592	26.1
Family 2 + 15	8	142487895	rs147691391	T	A	MROH5	NM_207414	c.A1346T	p.E449V	0.273	24
Family 6 + 17	7	100365542	rs183014219	C	A	ZAN	NM_173059	c.C4949A	p.T1650K	0.407	24.7
Family 3 + 13	17	11459147	rs185956842	C	T	SHISA6	NM_001173461	c.C890T	p.P297L	0.241	33
Family 7 + 14	3	53326592	rs35988197	G	C	DCP1A	NM_018403	c.C890G	p.A297G	0.278	24
Family 1 + 2	12	70949014	rs61758735	G	A	PTPRB	NM_001206971	c.C4145T	p.T1382I	0.00896	27.8
Family 1 + 10	17	56833502	rs770124556	G	G	PPM1E	NM_014906	c.144_145insCCCGAA	p.E48delinsEPE	0.375	-0.01

Alt, alternative allele; chr, chromosome; Ref, Reference allele

Table 3. Genes with multiple different segregating variants in two or more families

Family	Chr	Position	Variant	Ref	Alt	Gene	Accession number	Codon change	Protein Change	P-value (Schellevis et al. 2018 submitted)	CADD score
Family 18	1	94502780	1:94502780	C	T	ABCA4	NM_000350	c.G3734A	p.S1245N	NA	21.2
Family 14	1	94508969	rs61751374	G	A	ABCA4	NM_000350	c.C3113T	p.A1038V	0.800	20.5
Family 7	3	183905991	rs566108440	G	A	ABCF3	NM_001351298	c.G614A	p.R205Q	0.797	26.1
Family 2	3	183908940	rs779795407	C	T	ABCF3	NM_001351298	c.C1448T	p.P483L	NA	35
Family 17	2	241815411	rs140992177	T	C	AGXT	NM_000030	c.T836C	p.I279T	0.201	23.3
Family 1	2	241810796	rs121908524	T	A	AGXT	NM_000030	c.T454A	p.F152I	0.588	28.3
Family 7	16	1394822	rs148966323	C	T	BAIAP3	NM_001199096	c.C1673T	p.T558I	0.496	25.2
Family 2	16	1394491	rs114280977	G	A	BAIAP3	NM_001199096	c.G1516A	p.D506N	0.052	22
Family 2	8	139833569	rs145361557	C	T	COL22A1	NM_152888	c.G1055A	p.R352Q	0.823	23.1
Family 12	8	139838971	rs72731614	C	T	COL22A1	NM_152888	c.G899A	p.R300Q	NA	23.8
Family 18	1	34015872	rs149704396	C	T	CSMD2	NM_052896	c.G8390A	p.R2797Q	0.079	22.8
Family 14	1	34066488	rs755952714	C	T	CSMD2	NM_001281956	c.G6833A	p.G2278E	NA	24.9
Family 1	17	1944871	rs200625064	C	T	DPH1	NM_001346576	c.C778T	p.Q260X	NA	35
Family 2	17	1943099	rs80150196	C	G	DPH1	NM_001346576	c.C326G	p.P109R	0.429	28.4
Family 4	16	15733081	rs150196755	C	T	MARF1/ KIAA0430	NM_001184998	c.G10A	p.G4R	0.718	28.6
Family 6	16	15729650	rs192438053	C	T	MARF1	NM_001184998	c.G694A	p.G232R	0.495	25.8
Family 1	6	138655606	rs777828045	AAGG	A	ARFGEF3	NM_020340	c.5624_5626del	p.L1875 1876del	0.795	-0.01
Family 2	6	138615130	rs755891726	G	T	ARFGEF3	NM_020340	c.G3369T	p.R1123S	0.799	25.3
Family 18	1	109740175	rs940837035	TCA	T	KIAA1324	NM_001284353	c.1188_1189del	p.I396fs	0.582	-0.01
Family 4	1	109734349	rs41279690	G	A	KIAA1324	NM_001284353	c.G533A	p.G178D	0.374	23.3
Family 15	8	28989925	rs145324154	C	T	KIF13B	NM_015254	c.G2842A	p.A948T	0.573	32

Family	Chr	Position	Variant	Ref	Alt	Gene	Accession number	Codon change	Protein Change	P-value (Schellevis et al. 2018 submitted)	CADD score
Family 11	8	29102864	8:29102864	G	C	KIF13B	NM_015254	c.C148G	p.R50G	NA	27.8
Family 13	7	91871373	rs34358665	C	T	KRIT1	NM_001350672	c.G77A	p.R26Q	0.542	23.6
Family 12	7	91851344	7:91851344	T	C	KRIT1	NM_001013406	c.A1291G	p.K431E	NA	24.2
Family 17	18	44089726	rs571539488	G	A	LOXHD1	NM_001145473	c.C169T	p.P57S	NA	25.6
Family 7	18	44140215	rs759237437	C	C	LOXHD1	NM_144612	c.2891_2892insCTCAT-CAGAGGAGTCTC	p.S964del-insSSSESS	0.386	-0.01
Family 1	14	74971538	rs760036288	C	T	LTP2	NM_000428	c.G4396A	p.G1466R	NA	32
Family 15	14	74967643	14:74967643	A	C	LTP2	NM_000428	c.T5410G	p.C1804G	NA	27.2
Family 6	17	10426647	rs769778269	C	T	MYH2	NM_001100112	c.G5555A	p.R1852Q	NA	33
Family 11	17	10424643	rs34161789	C	T	MYH2	NM_001100112	c.G5780A	p.R1927Q	0.854	35
Family 17	10	95137125	rs367618675	G	A	MYOF	NM_133337	c.C2024T	p.A675V	NA	22
Family 11	10	95079636	rs146626145	G	A	MYOF	NM_133337	c.C552T	p.A1851V	0.248	23.8
Family 3	6	32180684	rs150079294	A	C	NOTCH4	NM_004557	c.T2443G	p.C815G	0.555	24.5
Family 12	6	32163648	rs764118051	G	T	NOTCH4	NM_004557	c.C5578A	p.R1860S	NA	34
Family 9	20	47364384	20:47364384	C	T	PREX1	NM_020820	c.G253A	p.D85N	NA	23.2
Family 11	20	47282854	rs149524742	C	T	PREX1	NM_020820	c.G1705A	p.V569M	0.044	34
Family 6	20	43945573	rs199904334	A	C	RBPIL	NM_001281449	c.A1525C	p.N509H	NA	25.1
Family 7	20	43942164	20:43942164	G	C	RBPIL	NM_001281448	c.G676C	p.V226L	NA	27.1
Family 7	20	43942170	20:43942170	A	C	RBPIL	NM_001281448	c.A682C	p.T228P	NA	26
Family 3	8	145736819	rs41555416	G	A	RECQL4	NM_004260	c.C3622T	p.R1208C	0.500	23.5
Family 13	8	145737142	rs61755067	C	G	RECQL4	NM_004260	c.G3424C	p.D1142H	0.585	32
Family 18	3	53126560	rs201230044	C	T	RFT1	NM_052859	c.G1283A	p.S428N	NA	22
Family 12	3	53126512	rs147740901	G	A	RFT1	NM_052859	c.C1331T	p.T444M	0.588	22.7

Family 17	9	135173569	9:135173569	C	T	SETX	NM_001351527	c.G5679A	p.M1893I	NA	25.1
Family 12	9	135210039	rs527394446	T	C	SETX	NM_001351527	c.A794G	p.D265G	0.659	28.5
Family 17	16	89965023	rs144328773	C	A	TCF25	NM_014972	c.C1081A	p.R361S	0.673	34
Family 18	16	89972604	rs137901241	G	A	TCF25	NM_014972	c.G1631A	p.R544Q	0.811	28.4
Family 17	2	179554624	rs202234172	C	T	TTN	NM_133378	c.28031-1G>A	-	0.360	26.1
Family 15	2	179628969	rs139504522	G	A	TTN	NM_003319	c.C9911T	p.P3304L	NA	23.6
Family 13	2	179396568	rs201218828	T	G	TTN	NM_003319	c.A7759C	p.E25860A	0.241	22.1
Family 11	2	179411137	2:179411137	C	T	TTN	NM_003319	c.G67726A	p.G22576S	NA	23.3
Family 14	9	132636952	rs142714756	G	A	USP20	NM_001008563	c.G1838A	p.R613H	NA	32
Family 12	9	132630423	rs148425010	G	A	USP20	NM_001008563	c.G830A	p.S277N	0.379	23.2
Family 10	5	82835589	rs146630369	T	C	VCAN	NM_001164097	c.T3806C	p.L1269P	0.796	24.2
Family 11	5	82850808	rs768896921	A	G	VCAN	NM_001126336	c.A1463G	p.N488S	NA	28.9
Family 13	14	75245347	14:75245347	TGAG	T	YLP1	NM_019589	c.1072_1074del	p.358_358del	NA	-0.01
Family 11	14	75265490	14:75265490	C	G	YLP1	NM_019589	c.C3490G	p.R1164G	NA	25.5
Family 17	3	102183076	rs375032047	C	T	ZPLD1	NM_175056	c.C790T	p.R264X	0.603	48
Family 15	3	102175036	3:102175036	G	A	ZPLD1	NM_175056	c.376-1G>A	-	NA	25.7

Alt, alternative allele; chr, chromosome; Ref, Reference allele

Burden, SKAT and SKAT-O results were obtained from our study and was performed with 263 sporadic CSC patients and 1352 population controls (Schellevis et al. 2018 submitted).

Table 4. Gene-based analysis results of the genes with multiple segregating variants in two or more families.

Gene	Case-control analysis			Family analysis
	Burden	SKAT	SKAT-O	RareIBD
ABCA4	0.074	0.295	0.129	0.051
ABCF3	0.386	1.000	0.523	0.047
AGXT	0.374	0.367	0.517	0.301
ARFGEF3	0.126	0.166	0.198	0.085
BAIAP3	0.076	0.663	0.132	0.143
COL22A1	0.564	0.506	0.684	0.020
CSMD2	0.460	0.266	0.471	0.328
DPH1	0.039	0.220	0.061	0.086
KIAA1324	0.074	0.035	0.051	0.097
KIF13B	0.053	0.026	0.044	0.288
KRIT1	0.345	0.821	0.446	0.021
LOXHD1	0.567	0.960	0.761	0.049
LTBP2	0.378	0.827	0.553	0.177
MARF1	0.710	1.000	0.878	0.094
MYH2	0.858	0.452	0.655	0.037
MYOF	0.016	0.128	0.025	0.144
NOTCH4	0.544	0.939	0.740	0.018
PREX1	0.044	0.165	0.073	0.061
RBPJL	0.131	0.262	0.195	0.014
RECQL4	0.234	0.538	0.391	0.144
RFT1	0.073	0.341	0.130	0.019
SETX	0.587	0.494	0.667	0.169
TCF25	0.782	0.594	0.829	0.191
TTN	0.577	0.435	0.596	0.072
USP20	0.226	0.948	0.354	0.215
VCAN	0.271	1.000	0.414	0.144
YLPM1	0.524	0.871	0.715	0.047

Burden, SKAT and SKAT-O results were obtained from our study and was performed with 263 sporadic CSC patients and 1352 population controls (Schellevis et al. 2018 submitted).

Discussion

In this exome sequencing study on familial CSC we included 72 individuals from 18 different families and focused on rare genetic variants that segregated with the disease in these families. We observed 11 variants that segregated in two families, of which one was also associated with CSC in a recent case-control study (Schellevis et al. 2018 submitted). In addition in 28 genes two different variants were found to segregate in two families, and 26 of these genes showed expression in the retina or RPE according to the Eye Integration Database.³¹

For AMD, a well-studied multifactorial eye disease with phenotypic overlap with CSC, exome sequencing studies in families have been successful in identifying rare variants that fully or partially segregate with the disease.³²⁻³⁷ Most variants in these AMD families were identified in genes of the complement system, in which common and rare variants were previously identified to be associated with AMD in case-control cohorts.³⁸ So far the only well-established genetic association for CSC has been found in common variants in the *complement factor H (CFH)* gene.^{11,13-15} However, in this study we did not observe any segregating rare variants in *CFH* in our families.

In this first unbiased exome sequencing study in familial CSC, we did not identify either a single variant nor multiple variants in a single gene that segregated with the CSC phenotype. This excludes that familial CSC is a Mendelian disease caused by mutations in a single gene. Analysis of exome sequencing data identified numerous variants that segregate with the disease in each individual family. However with the current study setting it is impossible to identify which of these variants might have an effect on the phenotype. Therefore, we focused on variants that segregated with CSC in at least two families, and on genes that carried multiple variants that segregated with CSC in at least two families.

In total, 11 segregating variants were observed in two families, of which the c.4145C>T; p.T1382I (rs61758735) variant in the *PTPRB* gene showed an association in the sporadic CSC case-control cohort (Schellevis et al. 2018 submitted). However, extended segregation analysis in additional family members excluded complete segregation of the *PTPRB* variant with the disease in one of two families. This is in line with results obtained in AMD families in which rare variants did not always fully segregate with the disease.^{32,34,39} Nevertheless, these variants are likely to contribute to the disease in these families, since several rare variants significantly associated with AMD in case-control studies often also show partial segregation in families.^{32,37}

The *PTPRB* gene encodes the vascular endothelial protein tyrosine phosphatase (VEPTP) protein. VEPTP is an important modulator of vascular endothelium morphogenesis and is involved promoting angiogenesis and regulates endothelial barrier functions by interacting with cadherin.^{40,41} Additionally, intra-ocular injection of anti-VEPTP suppress neovascularization in mice.⁴² As it is hypothesized that choroidal hyperpermeability and dysfunction is one of the underlying problems in CSC,² the *PTPRB* gene is an interesting candidate gene for CSC and should be investigated in future studies. The remaining 11 variants were extremely rare in the case-control cohort, and we can neither rule out nor confirm their possible role in the CSC disease mechanism at this time.

In 28 genes, multiple different segregating variants were observed in at least two families. For these genes gene-based associations in the case-control cohort (using SKAT, SKAT-O, and Burden test) and gene-based associations in the CSC families (using RareIBD) were evaluated. Five genes were nominally associated in the gene-burden analyses of the case-control cohort and 9 genes were nominally associated in the family dataset. However, none of the genes showed consistent associations in both the gene-burden analysis in the case-control cohort and in the family gene-burden analysis. Of the 5 genes associated in the case-control cohort, the *DPH1*, *KIAA1324*, and *PREX1* genes were close to significance in our families and might be interesting genes for replication in a larger family dataset.

In summary, we aimed to identify rare variants associated with familial CSC. In each family many variants segregated with the disease, but only few were found to segregate in at least two families. One variant was also associated with CSC in a recent case-control cohort (Schellevis et al. 2018 submitted), and this gene, *PTPRB*, has a function that could be of importance in the etiology of CSC. Therefore, *PTPRB* might be an interesting candidate gene for future studies on CSC. Future analyses should include more families with more individuals to increase the chance of finding segregating variants in multiple families, and to increase the power of the RareIBD analysis. Additionally, future studies may focus on the possible role for genetic variants in non-coding genetic regions, such as introns and promoter regions or large structural alterations that cannot be detected with exome sequencing, such as copy number variations.

In general, in familial CSC a Mendelian inheritance pattern of variants in one or a limited number of genes can be excluded based on this study. Instead, familial CSC may be a heterogeneous Mendelian disease caused by variants in many different genes, or alternatively CSC may represent a complex disease to which both genetic and environmental factors contribute.

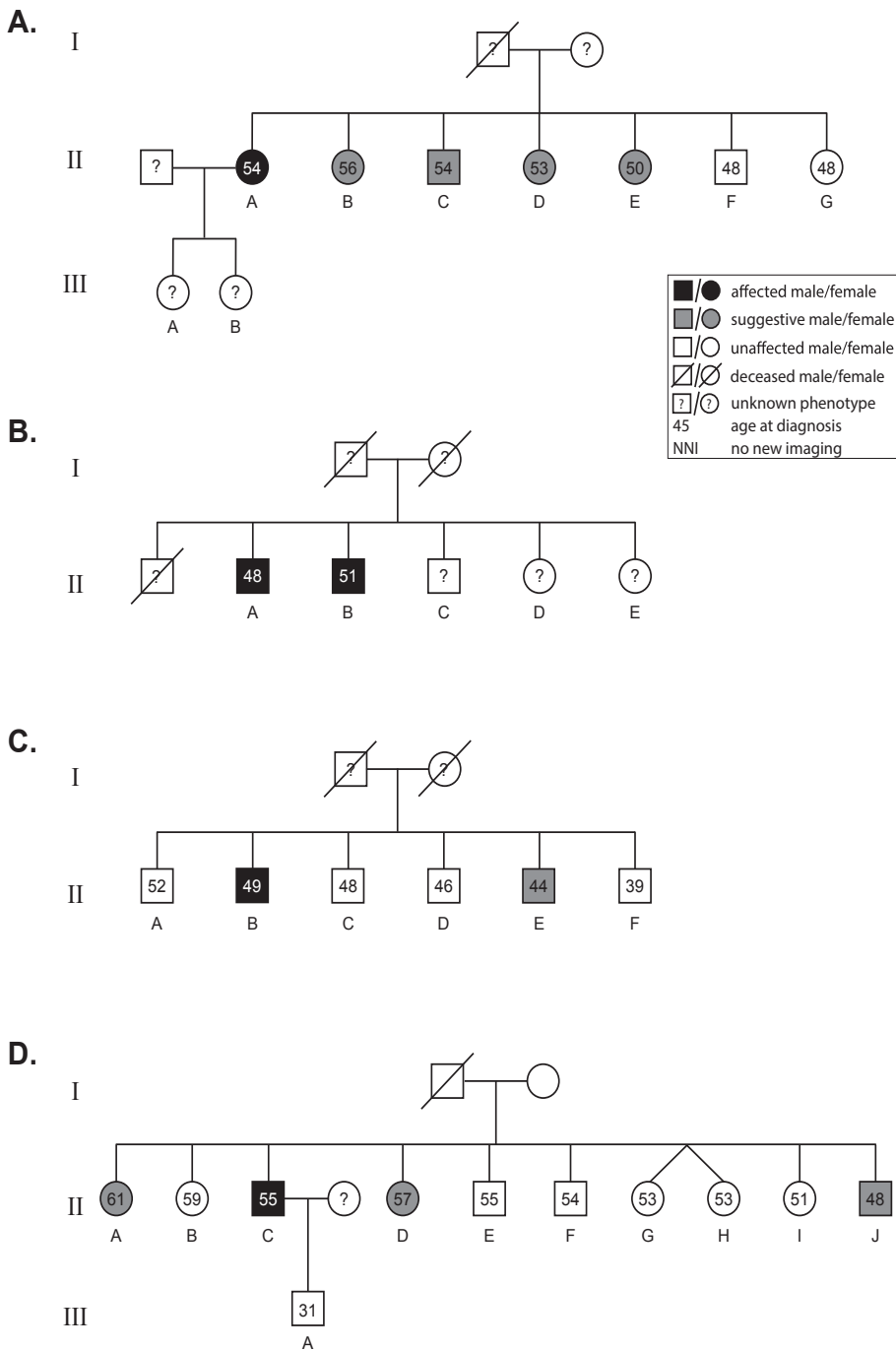
Footnotes and Financial Disclosures

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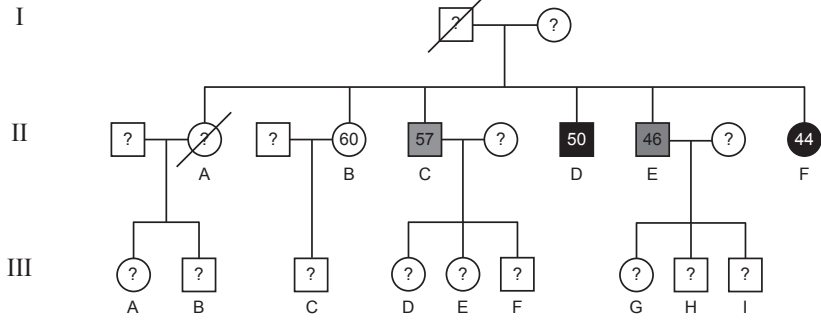
Conflict of interest: no conflicting relationship exists for any author.

Supplementary Files

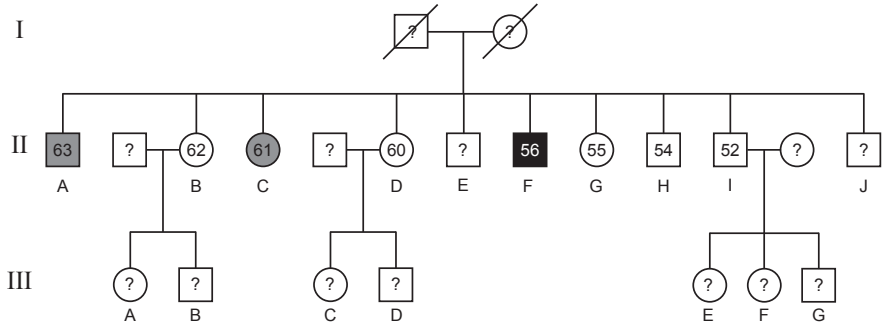


Supplementary Figure 1: Pedigrees of families included in the current study. Individuals that were sent in for exome sequencing are depicted with a blue arrow.

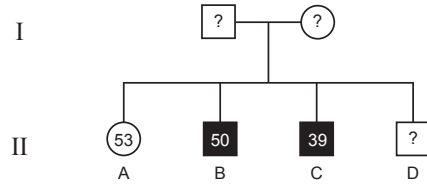
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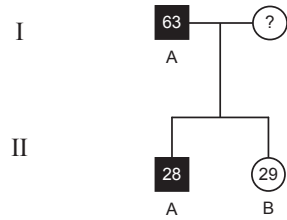
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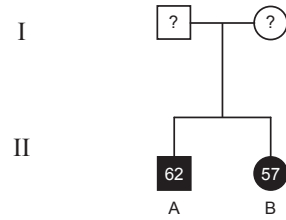
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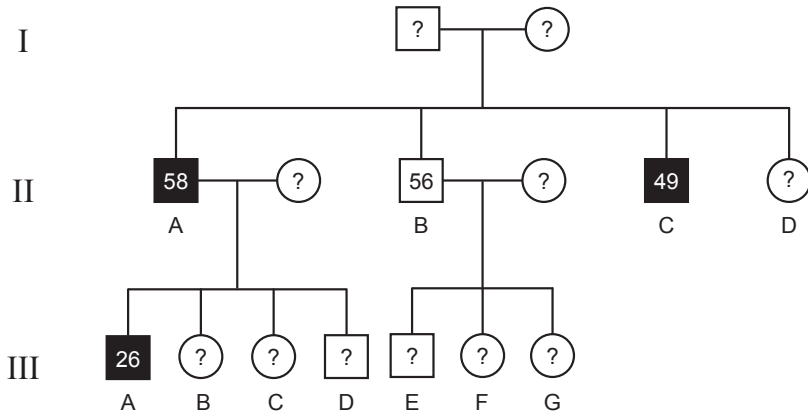
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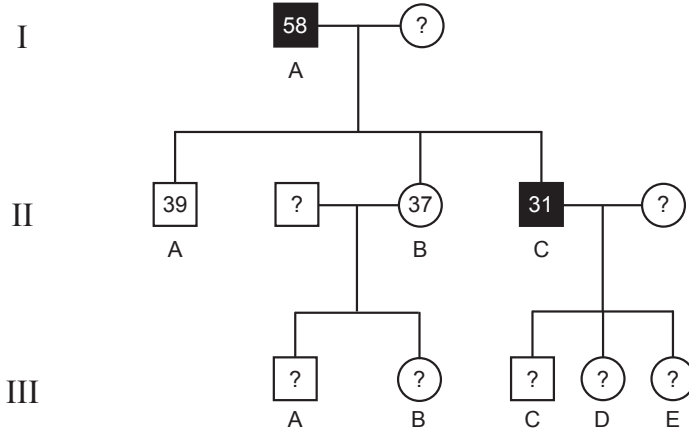
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J.

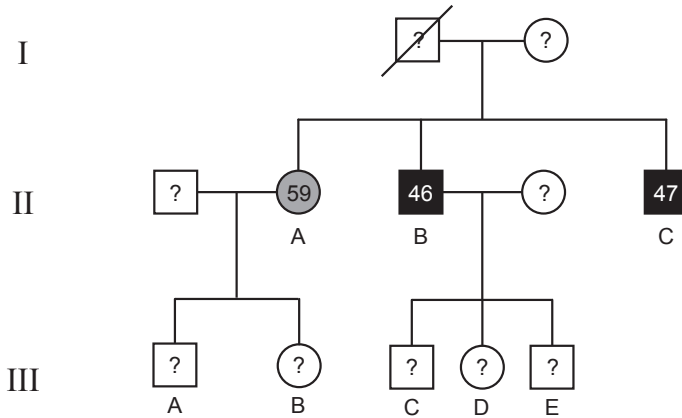


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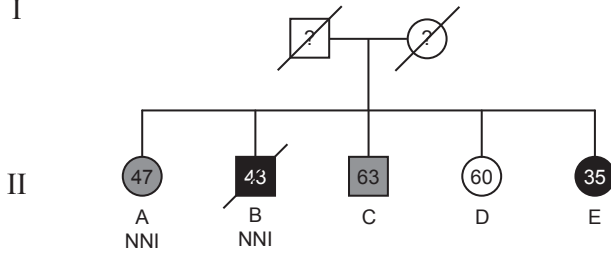


215

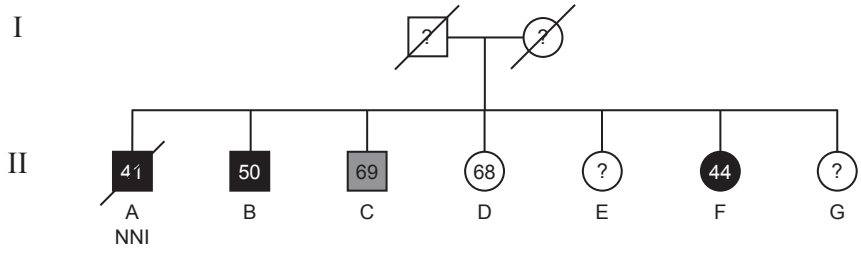
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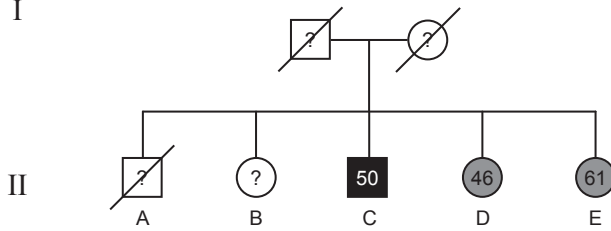
S. I



T. I



U. I



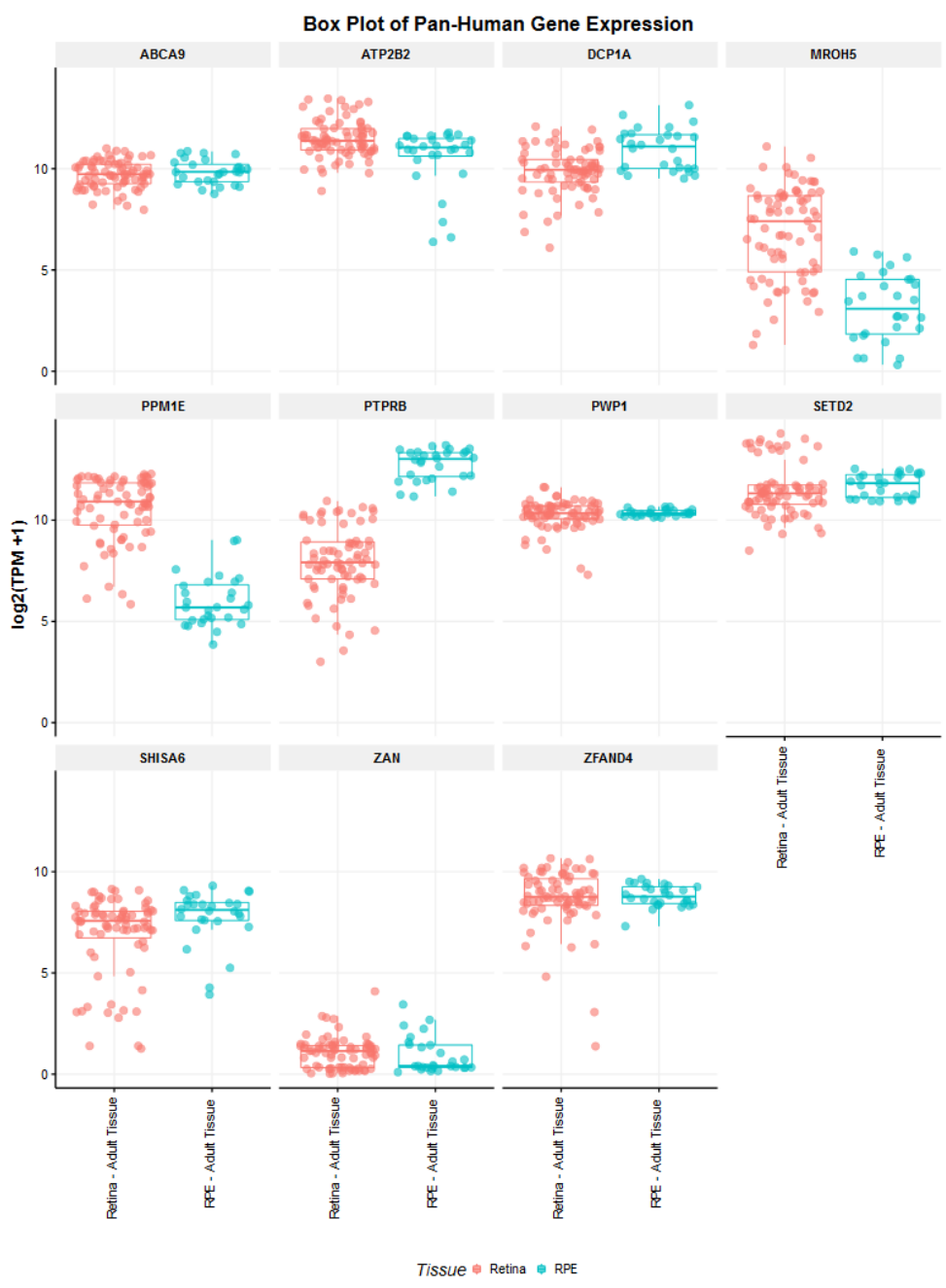
Supplementary Table 1. Genes with multiple variants within one family

Family	Chr	Position	Variant	REF	ALT	Gene	Accession number	Codon change	Protein change	P-value (Schelis et al. 2018 submitted)	CADD score
Family 12	10	79397312	rs200474297	T	C	KCNMA1	NM_001014797	c.A89G	p.H30R	0,37	24,4
Family 12	10	79397236	rs778736171	AGAG	*	KCNMA1	NM_001014797	-	-	NA	NA
Family 7	20	43942164	20:43942164	G	C	RBPJL	NM_001281448	c.G676C	p.V226L	NA	27,1
Family 7	20	43942170	20:43942170	A	C	RBPJL	NM_001281448	c.A682C	p.T228P	NA	26
Family 8	12	58017860	rs113207856	G	A	SLC26A10	NM_133489	c.G1206A	p.W402X	0,14	39
Family 8	12	58018668	rs111924104	T	G	SLC26A10	NM_133489	c.T1247G	p.L416R	0,19	27,5
Family 1	2	175202184	2:175202184	AGGGCGG- GCGGCGG- CAGGGCGG- GCGGCG	*	SP9	NM_001145250	-	-	NA	NA
Family 1	2	175202187	rs889377194	GGCGGGCGG- CGGCA	*	SP9	NM_001145250	-	-	NA	NA

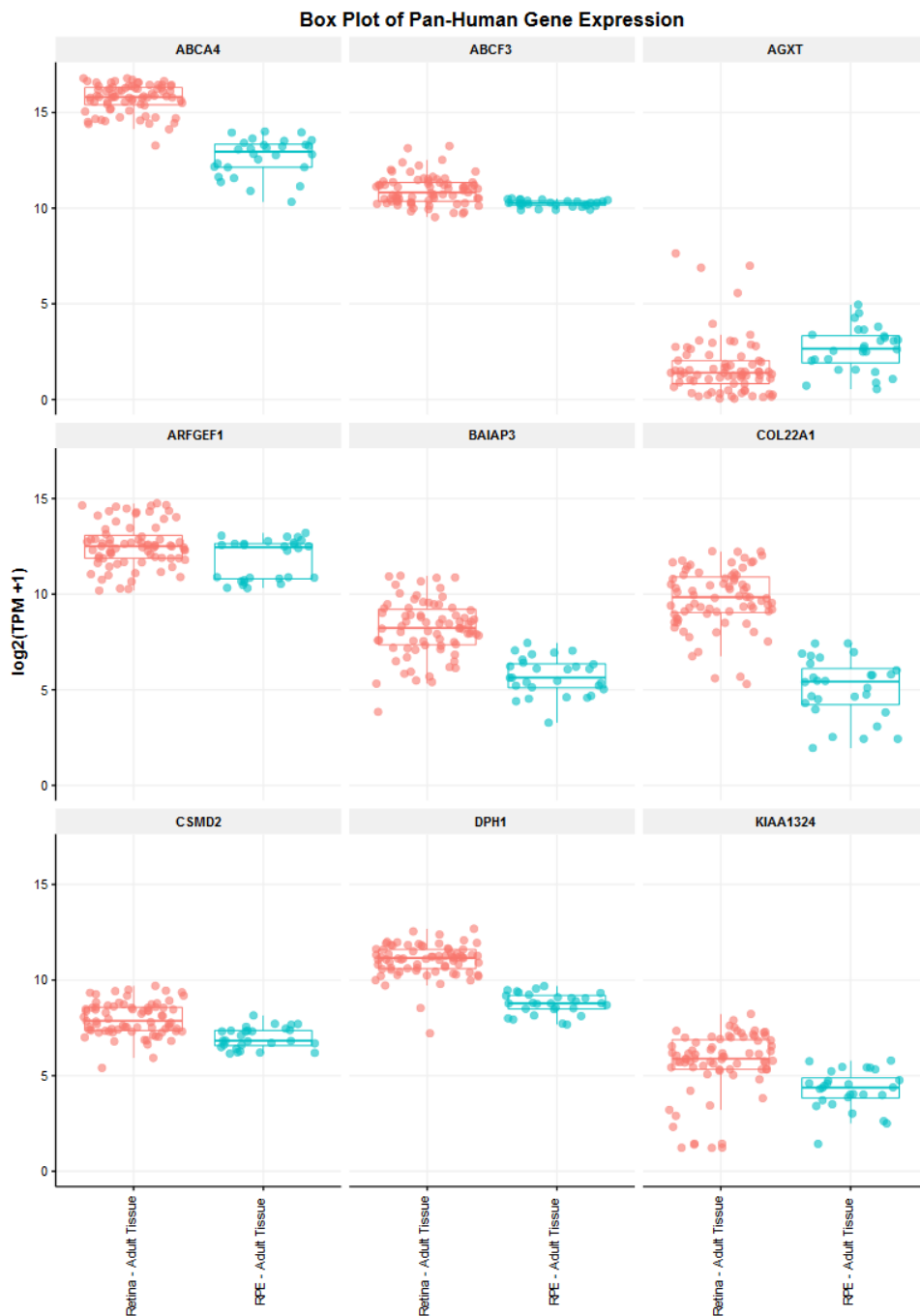
Alt, alternative allele; chr, chromosome; Ref, Reference allele

Supplementary Table 2



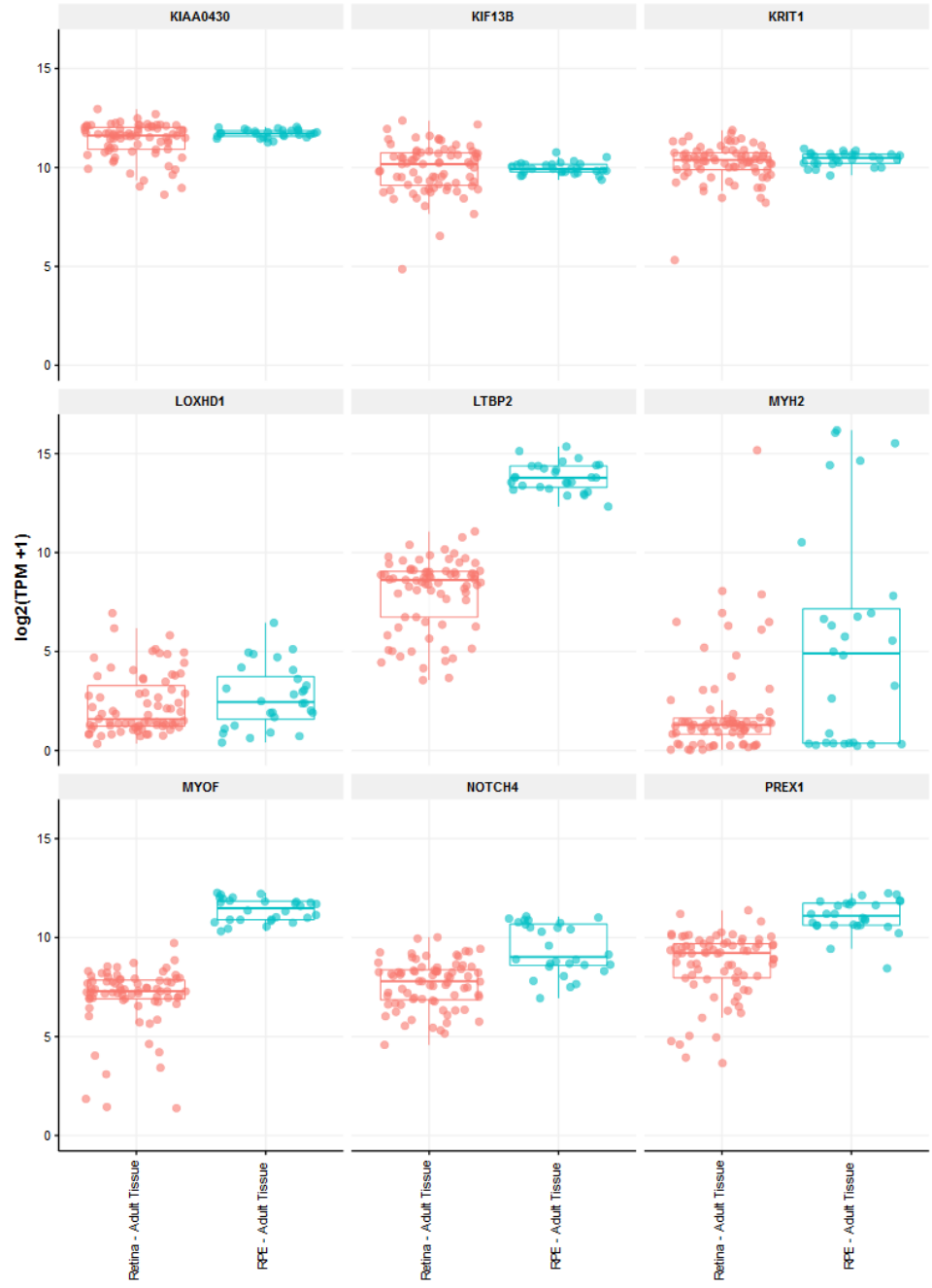


Supplementary Figure 2. Expression according to the Eye integration database³¹ for the 11 genes with the same variant in two families. Adult retina is depicted in red, whereas RPE is blue.

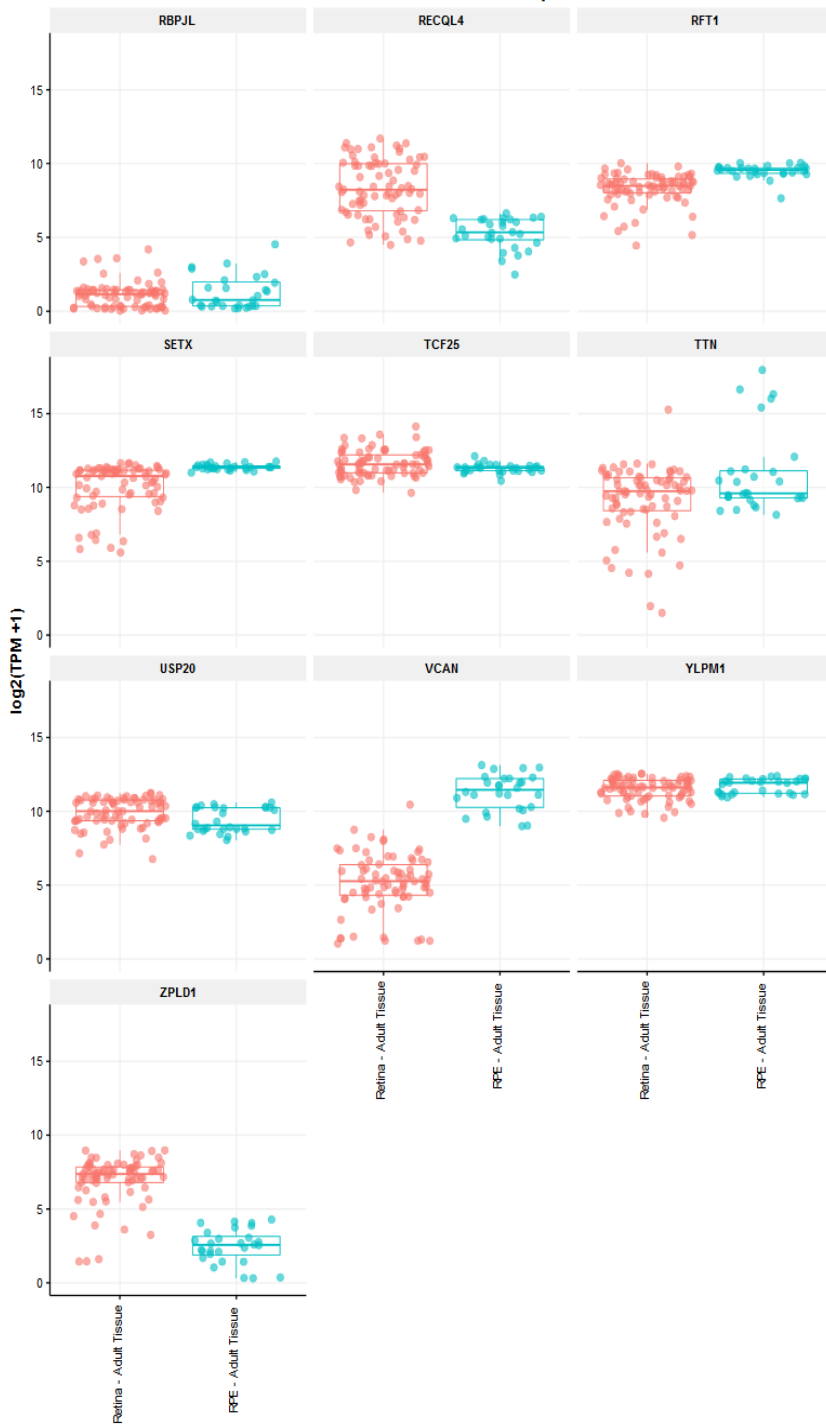


Supplementary Figure 3. Expression according to the Eye integration database³¹ for the 28 genes with multiple different variant in two families. Adult retina is depicted in red, whereas RPE is blue.

Box Plot of Pan-Human Gene Expression



Box Plot of Pan-Human Gene Expression



References

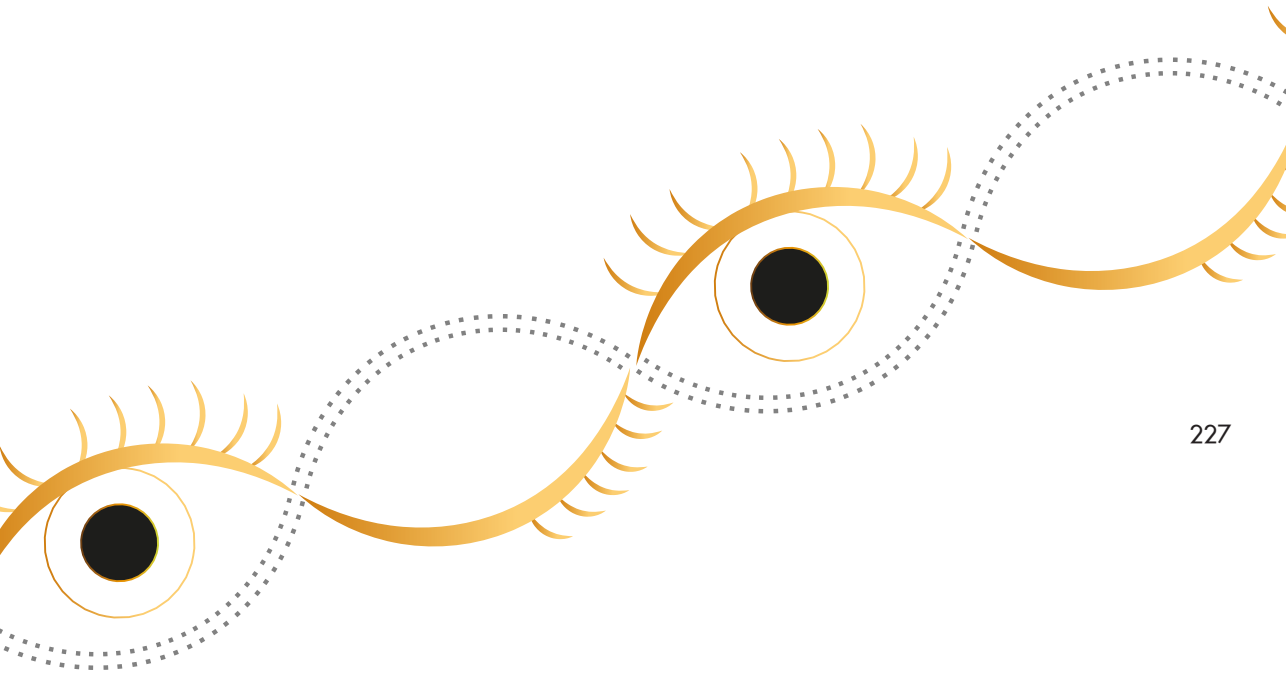
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5. Elevated steroid hormone levels in active chronic central serous chorioretinopathy

5. Steroid Measurements In Central Serous Chorioretinopathy





5 Elevated steroid hormone levels in active chronic central serous chorioretinopathy

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Manuscript in preparation

Abstract

Importance: Chronic central serous chorioretinopathy (cCSC) is a vision-threatening eye disease characterized by fluid accumulation between the photoreceptors and the retinal pigment epithelium. The etiology of cCSC is currently unknown, but reported associations with the use of steroids, stress, pregnancy and the male gender suggest an important role for the steroid hormone system in the disease.

Objective: To perform a comprehensive analysis of the steroid hormone system in cCSC by measuring levels of 17 different steroid hormones simultaneously.

Design: Case-control study

Setting: Two referral university medical centers for outpatient care

Participants: 46 male patients with active cCSC and 46 male age-matched controls.

Main outcome measurements: Serum hormone levels of 17 steroid hormones and Spearman's correlations between steroid hormones. The AbsoluteIDQ stero17 kit was used, measuring cortisol, cortisone, 11-deoxycortisol, aldosterone, corticosterone, deoxycorticosterone [DOC], 17 α -hydroxyprogesterone [17-OHP], progesterone, estrone [E1], estradiol [E2], androstenedione, androsterone, dehydroepiandrosterone [DHEA], dehydroepiandrosterone sulfate [DHEAS], dihydrotestosterone [DHT], etiocholanolone and testosterone.

Results: Elevated levels of androsterone, estrone, etiocholanolone and androstenedione were observed in 46 male cCSC patients (mean age: 50 [range: 28-72] years) compared to 46 male controls (mean age: 50 [range: 22-72]). Median hormone levels in cCSC patients vs. controls, respectively, were: androsterone: 0.84 ng/ml (Inter quartile range [IQR]=0.61-1.06) vs. 0.69 ng/ml (IQR=0.48-0.96, $P=0.031$), estrone: 0.12 ng/ml (IQR=0.10-0.15) vs. 0.10 ng/ml (IQR=0.08-0.11, $P=0.0048$) and etiocholanolone: 0.19 ng/ml (IQR=0.15-0.29) vs. 0.13 ng/ml (IQR=0.099-0.20, $P=0.0061$). Mean levels of androstenedione, which were normally distributed, were 3.10 ng/ml ($SD=1.03$) vs. 2.55 ng/ml ($SD=0.95$), in cCSC patients vs. controls, respectively. Additionally, Spearman's correlations between aldosterone and 11-deoxycortisol, androsterone, DHEA, DHEAS and E1 differed between cCSC patients and controls, as well as between androsterone and E1, and between DHT and 17OHP.

Conclusions and relevance: In this study, by simultaneously measuring levels of 17 steroid hormones, a comprehensive overview of the status of the steroid hormone system in active cCSC was obtained. Levels of four hormones were significantly elevated in cCSC patients compared to controls. Also, the relationships between steroid hormones differed between cases and controls, suggesting that the balance in the steroid hormone system is altered in cCSC patients. This study suggests that steroid hormones might be useful for predicting active phases of cCSC, and that restoring steroid hormone homeostasis might be a target for treatment of cCSC.

Introduction

Central serous chorioretinopathy (CSC) is a multifactorial retinal disease characterized by fluid accumulation between the photoreceptors and the retinal pigment epithelial (RPE) leading to vision impairment.¹⁻³ The etiology of CSC is currently unknown and treatment options are limited. Two forms of CSC are described: acute CSC and chronic CSC (cCSC). Acute CSC usually spontaneously resolves, while patients with chronic CSC (cCSC) have persisting complaints or multiple separate episodes of fluid accumulation accompanied with permanent loss of visual acuity and RPE alterations.^{1,4} CSC is more frequent in males (~80% of patients) and has been associated with stress, the use of steroids, type A personality and pregnancy,¹⁻³ suggesting an important role for the steroid hormone system in the disease.

Steroid hormones regulate a large number of physiological processes, such as sex differentiation, metabolism and immune responses.⁵ To date, only a limited number of studies have investigated hormone levels in CSC patients. The most widely studied hormone is cortisol with over 11 studies,⁶⁻¹⁶ aldosterone was measured in 2 studies,^{8,17} testosterone levels were measured in 4 studies,^{7,8,10,12} dehydroepiandrosterone (DHEA) and dehydroepiandrosterone sulfate (DHEAS) were both described in only one study.^{8,18} However, results between studies were not consistent, showing associations with CSC and cortisol or testosterone levels in some studies, but not in others.⁶⁻¹⁶ Notably, these studies focused only on a limited number of hormones, while it is known that many complex interactions exist between steroid hormones and that they are highly correlated.⁵ A comprehensive analysis of the steroid hormone system would not only provide information on differences in levels on individual steroid hormones, but would also allow for the assessments of interactions between hormones.

Therefore, in order to better understand the role of the hormone system and to observe possible imbalances in this system in cCSC, we simultaneously measured 17 different steroid hormones in serum samples of 46 patients with active cCSC and 46 age- and sex-matched controls. With this study we aimed to increase our knowledge on the status of the steroid hormone system in cCSC patients, in order to gain more insight into the mechanisms underlying the disease.

Key Points

Question: Which steroid hormones are associated with active chronic central serous chorioretinopathy (cCSC)?

Findings: By measuring 17 steroid hormones simultaneously in patients with cCSC compared to age- and sex-matched controls, we observed elevated levels of androsterone, estrone, etiocholanolone and androstenedione in cCSC patients. Additionally, correlations between steroid hormones, in particular of the aldosterone hormone, were altered in cCSC patients.

Meaning: This study indicates involvement of sex-hormones in cCSC and suggests an altered balance of the hormone system in patients with active cCSC.

Methods

Study participants

In this multicenter study, we included 46 patients with cCSC from 2 different university hospitals. As controls 46 age- and sex-matched individuals from the EUGENDA database were used (www.eugenda.org). In total, 16 cCSC patients and 36 controls were collected at the Radboud university medical center (Nijmegen, The Netherlands) and 30 cCSC patients and 10 controls were collected at the University Hospital of Cologne (Cologne, Germany). The diagnosis of cCSC was based on multimodal grading of images obtained via color fundus photographs, optical coherence tomography, fluorescein angiography (FA) and indocyanine green angiography (ICGA) by two independent graders (LA, VS). Included were patients with active chronic cCSC, defined as focal or diffuse leakage in FA/ICGA with retinal pigment epithelium (RPE) alterations and visible subretinal fluid on spectral domain optical coherence tomography (SD-OCT). Patients with extensive RPE alterations (≥ 5 disc areas) were excluded. None of the controls and cCSC patients showed signs of other retinal diseases (drusen, age-related macular degeneration, former or present choroidal neovascularization, myopic degeneration, dome-shaped maculopathy, hemangioma, vitreomacular traction, diabetic macular edema etc.) on OCT, fundus photography or on FA/ICGA. Individuals who reported steroid or eplerenon/spirinolacton use within last 6 months at the moment of blood withdrawal were excluded from the study.

Informed consents were obtained for all individuals participating in this study. This study was performed according to the guidelines of the Declaration of Helsinki and the local ethic committees of the participating hospitals approved the study.

231

Steroid measurements

Serum samples were collected using a standardized coagulation and centrifugation protocol, and samples were stored at -80°C within 1 hour after collection. Time of blood withdrawal was recorded for the cCSC patients as they were collected specifically for this study, but was not known for the EUGENDA control samples which were collected previously for the EUGENDA database. For each participant an aliquot of frozen serum ($500\mu\text{l}$) was shipped to Biocrates Life Sciences (Innsbruck, Austria) and 17 different steroid hormones were measured using the AbsolutelDQ® stero17 assay. The 17 steroids include glucocorticoids (cortisol, cortisone and 11-deoxycortisol), mineralocorticoids (aldosterone, corticosterone and deoxycorticosterone [DOC]), progestogens (17 α -hydroxyprogesterone [17-OHP] and progesterone), estrogens (estrone [E1] and estradiol [E2]), and androgens (androstenedione, androsterone, dehydroepiandrosterone [DHEA], dehydroepiandrosterone sulfate [DHEAS], dihydrotestosterone [DHT], etiocholanolone and testosterone). Measurements of the 17 hormones were obtained from Biocrates in an Excel sheet along with a corresponding data report.

Statistical Analysis.

All statistical analyses were conducted using SPSS (IBM, version 25). Hormone levels in cCSC patients and controls were tested for normality with the Shapiro-Wilk test. Levels of normally distributed hormones were compared between cCSC patients and controls with an independent sample t-test, whereas non-normally distributed

hormones were tested with a Mann-Whitney *U* test. The levels of associated steroids (P-values <0.05) were compared between collection sites (Cologne and Nijmegen) with the same tests, depending on the normality of the distribution of the individual hormones. Additionally, levels of the associated steroids were compared between time of blood withdrawal with a Kruskal-Wallis test with the following hourly bins: 8.00-9.00, 9.00-10.00, 10.00-12.00, 12.00-13.00, and 14.00-15.00.

Because all steroid hormones are derived from cholesterol, and are thus highly correlated, the Spearman's rho rank correlation between the different steroid hormones was calculated. The correlation coefficients of cCSC patients and the control individuals were compared using the "r.test" function of the "psych" package. The r.tests first transforms the rho correlations into Z-scores using the Fisher Z transformation, and compares the differences divided by the standard error of the differences to obtain a p-value as proposed by Cohen and Cohen.¹⁹

Results

This study included 46 male patients with an active episode of cCSC and 46 unaffected male controls collected at two university hospitals. The mean age of the included patients was 50 (range: 28-72) years, and the mean age of the age-matched-controls was 50 (range: 22-72) years. Fourteen of the 17 steroid hormones were successfully measured, whereas 3 steroids had measurements outside of the limit of detection (LOD) in some samples [aldosterone (10 samples; 10.8%), estradiol (E2) (1 sample; 1.1%), etiocholanolone (11 samples; 12.0%)]. The distribution of samples that ranged outside of the LOD was similar between cCSC patients and controls. All hormone levels, except for androstenedione, cortisone and DHEAS, showed deviation from normality in either cCSC patients or controls, therefore nonparametric tests were used for these hormones and median levels with inter quartile ranges (IQR) are described.

Higher levels of androsterone, androstenedione, E1 and etiocholanolone were observed in cCSC patients compared to controls (Figure 1, Table 1). Median levels for androsterone were 0.84 ng/ml (IQR=0.61-1.06) in cCSC patients and 0.69 ng/ml (IQR=0.48-0.96) in controls (P=0.031). The median levels of E1 in cCSC patients were 0.12 ng/ml (IQR=0.10-0.15) and 0.10 ng/ml (IQR=0.08-0.11) in controls (P=0.0048). Median concentration of etiocholanolone was 0.19 ng/ml (IQR=0.15-0.29) in cCSC patients and 0.13 ng/ml (IQR=0.099-0.20) in controls (P=0.0061). Finally, mean androstenedione levels were 3.10 ng/ml (Standard deviation [SD]=1.03) in cCSC patients and 2.55 ng/ml (SD=0.95) in controls (P=0.0094). No difference was observed between samples collected from Nijmegen or Cologne for the associated steroids (Supplementary Table 1) and no association was observed between levels of these four steroids and the time of blood withdrawal (Supplementary Table 2).

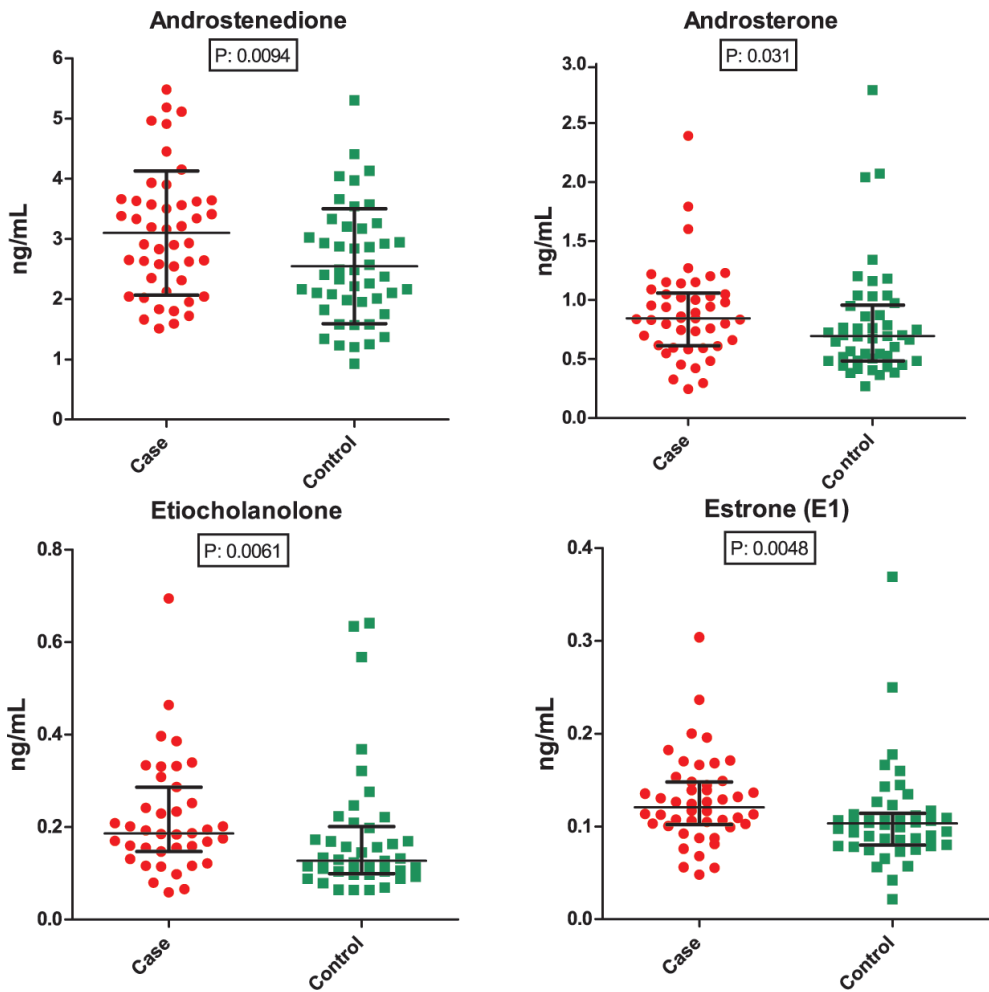


Figure 1: Hormone levels associated with cCSC. Different panels indicate the absolute hormone levels measured in cCSC patients (case) in red and controls in green. For Androstenedione error bars indicate mean and standard deviation, while for the other hormones the median and inter quartile ranges are depicted. Values for mean and medians can be found in Table 1.

Next, the relationship between steroids was investigated in cCSC patients and controls by calculating Spearman's rho correlations between all measured steroids. Numerous steroid hormones were highly correlated in both the cCSC patients and controls (Figure 2). Comparison of the Spearman's correlation coefficients between cCSC patients and controls identified a difference in correlations for aldosterone with androsterone (0.316 vs. -0.379, P-value=0.00076), DHEA (0.029 vs. -0.445, P-value=0.019), DHEAS (0.117 vs. -0.333, P-value=0.031), E1 (0.381 vs. -0.085, P-value=0.024) and 11-deoxycortisol (0.208 vs. -0.255, P-value=0.029) in controls and cCSC patients, respectively. Additionally, a difference in the correlation of androsterone with E1 (0.613 vs. -0.182, P-value=0.014) and DHT with 17-OHP (-0.100 vs. 0.538, P-value=0.0011) was observed between controls and cCSC patients,

Table 1. Association results of steroids in 46 cCSC patients compared to 46 controls

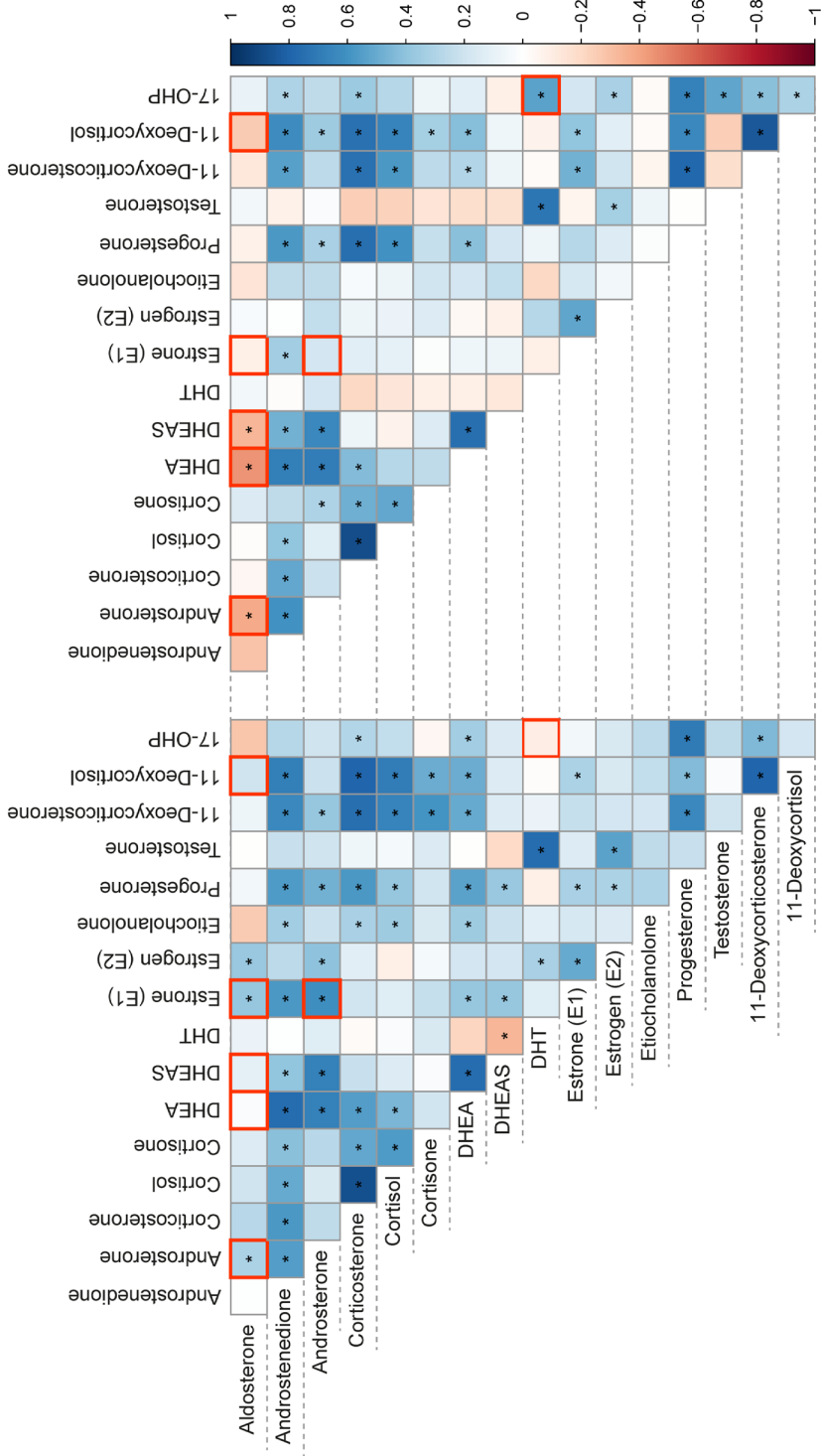
	Concentration in ng/ml				Mann Whitney U 2-sided P-value
	Controls		cCSC patients		
Non-normally distributed hormones	Median ng/ml	IQR	Median ng/ml	IQR	
11-Deoxy-corticosterone	0.070	(0.053-0.11)	0.088	(0.052-0.13)	0.323
11-Deoxycortisol	0.55	(0.35-0.9)	0.59	(0.41-1.26)	0.492
17 α -Hydroxyprogesterone (17-OHP)	2.11	(1.6-2.64)	2.14	(1.63-2.63)	0.854
Aldosterone	0.14	(0.085-0.21)	0.16	(0.13-0.23)	0.303
Androsterone	0.69	(0.48-0.96)	0.84	(0.61-1.06)	0.031
Corticosterone	5.12	(3.048-8.49)	4.96	(2.65-9.82)	0.975
Cortisol	259.35	(209.65-360.13)	275.60	(232.05-357.35)	0.331
Dehydroepiandrosterone (DHEA)	10.05	(5.15-15)	11.25	(7.63-16.35)	0.414
Dihydrotestosterone (DHT)	1.28	(1.04-1.73)	1.24	(0.90-1.81)	0.468
Estrone (E1)	0.10	(0.080-0.11)	0.12	(0.10-0.15)	0.0048
Estradiol (E2)	0.067	(0.052-0.081)	0.060	(0.052-0.083)	0.654
Etiocholanolone	0.13	(0.099-0.20)	0.19	(0.15-0.29)	0.0061
Progesterone	0.17	(0.13-0.26)	0.16	(0.12-0.25)	0.797
Testosterone	14.75	(10.40-18.40)	14.50	(10.8-17.45)	0.587
Normally distributed hormones	Mean ng/ml	SD	Mean ng/ml	SD	Independent T-Test P-value
Androstenedione	2.55	0.95	3.10	1.029	0.0094
Cortisone	61.78	16.56	68.01	13.62	0.052
Dehydroepiandrosterone sulphate (DHEAS)	5064.82	3030.96	5759.03	2638.17	0.244

cCSC: chronic central serous chorioretinopathy, IQR: Inter quartile range depicting the 25 and 75 percentile, SD: Standard deviation

respectively (Table 2, Figure 2, red bordered squares). The four steroids associated with cCSC were positively correlated in all groups (Supplementary Figure 1).

Discussion

In order to obtain an overview of the steroid hormone system in cCSC we studied 17 steroids in 46 patients with active cCSC and 46 age-matched controls. We measured elevated levels of androsterone, androstenedione, estrone and etiocholanolone in patients with active cCSC compared to controls. Additionally, alterations in the balance of the steroid hormone system were observed in cCSC patients, mainly involving relationships of aldosterone with other steroids.



cCSC Patients

Controls

Figure 2: Spearman's correlations between steroid hormones in cCSC patients and controls. On the left side correlations are depicted as observed in the controls, whereas correlations in cCSC patients are found on the right. Colors indicate positive (blue) and negative (red) correlations between the different hormones, and significant correlations are indicated with an asterisk (*). Red bordered squares are those correlations that were significantly different between cCSC patients and controls (Table 2).

Table 2. Difference in correlations of steroid hormones between cCSC patients and controls

Correlation	Controls		cCSC Patients		Delta Rho	Delta Z	P-value
	Spearman Rho	Correlation P-value	Spearman Rho	Correlation P-value			
Aldosterone - 11-Deoxycortisol	0.208	0.198	-0.255	0.103	0.463	2.188	0.029
Aldosterone - Androsterone	0.316	0.047	-0.379	0.013	0.695	3.367	0.00076
Aldosterone - DHEA	0.029	0.859	-0.445	0.0031	0.474	2.354	0.019
Aldosterone - DHEAS	0.117	0.473	-0.333	0.031	0.450	2.152	0.031
Aldosterone - Estrone (E1)	0.381	0.0153	-0.085	0.592	0.466	2.255	0.024
Estrone (E1) - Androsterone	0.613	6.00E-06	0.182	0.227	0.431	2.456	0.014
DHT - 17OHP	-0.100	0.510	0.538	1.13E-04	-0.638	3.255	0.0011

17-OHP: 17 α -Hydroxyprogesterone, DHEA: Dehydroepiandrosterone, DHEAS: Dehydroepiandrosterone sulphate, DHT: Dihydrotestosterone. cCSC: chronic central serous chorioretinopathy

Several steps were performed in order to prevent confounding by factors that are known to influence steroid levels. First, to homogenize the dataset we include only males in this study, because the disease is most prevalent in this group (9.9:100,000 vs. 1.7:100,000 in women).²⁰ Secondly, the controls used in this study were age-matched with the cCSC patients. Also, all individuals that reported steroid use at blood withdrawal were excluded. Finally, for the cCSC-associated hormones we confirmed that levels were not dependent on time at blood withdrawal and did not differ between recruitment sites.

A limited number of studies have been performed on steroid hormone imbalances in cCSC patients till date, and these studies focused on either single measurements or at most on four steroids simultaneously. Association results were variable between studies and reported hormone levels also varied greatly between studies. Some associations with testosterone or morning, evening or 24h cortisol levels have been reported,^{6,8-10,13,15} but other studies did not confirm these results.^{10,11,14} Aldosterone, DHEA and DHEAS were measured, but no association with cCSC were observed in the studies investigating these hormones.^{8,18}

To our knowledge, no studies have been published that have reported higher levels of androsterone, androstenedione, E1 and etiocholanolone in cCSC patients as observed in the current study. Androsterone, androstenedione, etiocholanolone and estrone are important products of the sex-hormone branch of the steroid hormone system. Androstenedione, androsterone and etiocholanolone are weak androgens and mainly function as precursors or metabolites for testosterone (Figure 3).⁵ However, androsterone and etiocholanolone have additional neurosteroid properties acting on the GABA_A receptor and are able to cross to the blood-brain barrier to prevent seizures.^{21,22} Androstenedione is the main precursor of testosterone and estrone.⁵

Estrone is a weak estrogen hormone that can readily be converted to estradiol, has binding affinity to the estrogen receptors ER α and ER β and its levels are known to be increased in the blood of postmenopausal women.²³ Currently, the role of these hormones in the etiology of cCSC is still unknown. Future studies might focus on determining the influence of elevated levels of these hormones on the RPE and choroid using for example cell models.

We assessed the steroid hormone system as a whole by investigating relationships between the different hormones because steroid hormones are highly correlated, and multiple testing correction with Bonferroni would be too conservative. Interestingly, we observed several altered correlations between the measured hormones when comparing controls and cCSC patients. In controls, a triangular positive correlation was observed between aldosterone, androsterone and estrone. In the cCSC patients, a negative correlation for aldosterone with androsterone was observed and the correlation with estrone with both aldosterone and androsterone was lost. Additionally, in cCSC patients DHEA, DHEAS and 11-deoxycortisol were negatively correlated with aldosterone, and DHT was negatively correlated with 17-OHP, while these correlations were not observed in controls. Although aldosterone itself was not associated with cCSC in this study, its altered relationship with multiple other hormones suggests an altered balance in the steroid hormone system in cCSC patients. In light of the current clinical trials targeting the mineralocorticoid receptor (the binding site of aldosterone), this observation might be clinically relevant. In future studies, follow-up measurements of patients with active cCSC when they either have spontaneously resolved, have responded to treatment (micropulse laser, photodynamic therapy or eplerenone) or have persisting complaints might be informative to determine whether alterations in the hormone level balance might be used as predictor of disease status or treatment response, or might in itself be a target for treatment.

237

Here, by simultaneously measuring 17 components of the steroid hormone system in a group of active cCSC patients the status of the hormone system could be investigated. This approach allowed us to describe elevated single hormone levels in addition to altered relationships between various hormones in cCSC patients compared to controls. Even though this is one of the larger studies on steroid measurements in cCSC patients to date, the current sample size is still limited and future studies should focus on replication of the current study in an independent cCSC cohort. Additionally, as only male cCSC patients were included in this study, measurements of the 17 steroids and the hormone system balance in female cCSC patients could provide new insights on sex-specific differences in the disease.

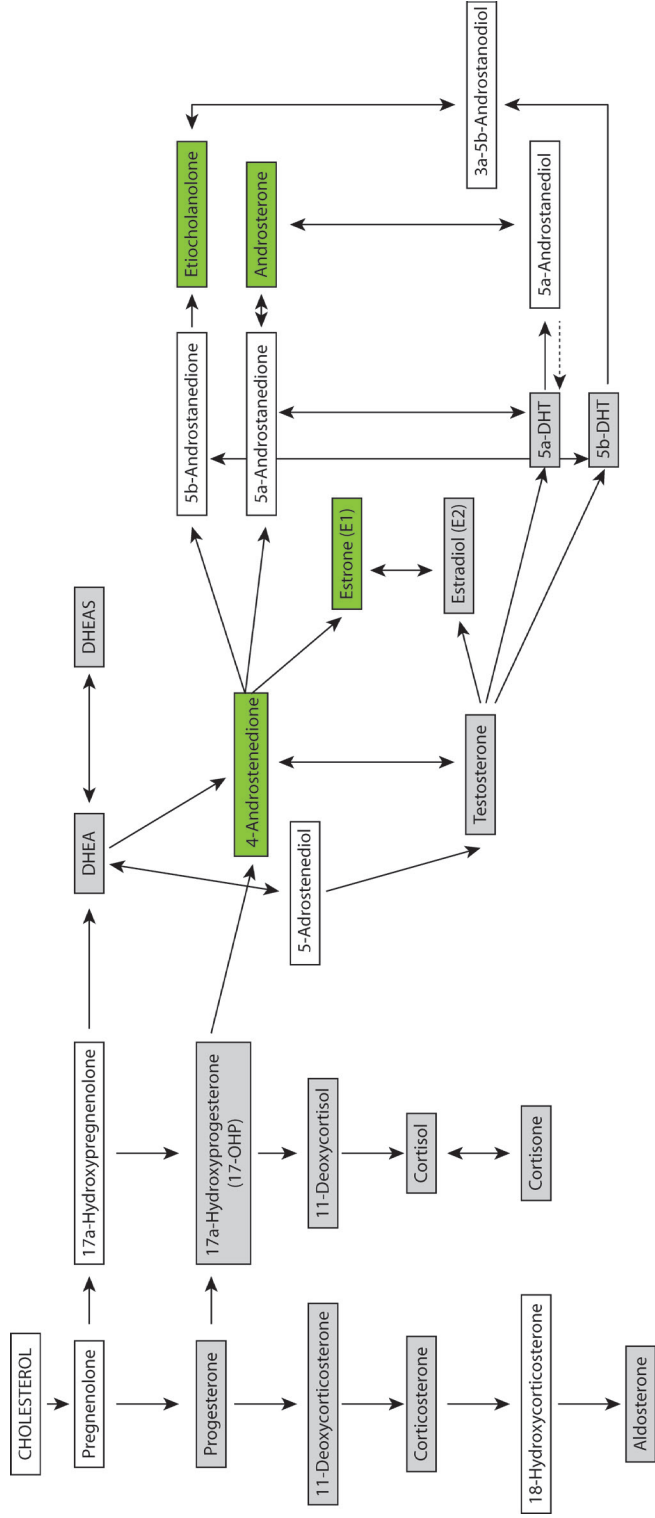


Figure 3: Overview of the steroid hormone pathway and alterations in cCS. Conversion steps between hormones are indicated with arrows. The hormones measured in this study are indicated as grey boxes, while green boxes indicate significantly elevated hormone levels in cCS patients. Figure based on .5

Conclusions

In this study, we measured elevated levels of androsterone, androstenedione, E1 and etiocholanolone in patients with an active episode of cCSC compared to controls. Additionally, we observed altered correlations between hormones in cCSC patients, mainly with aldosterone, indicative of an altered hormone system balance in individuals with cCSC. This study provides new insights into the status of the steroid hormone system in active cCSC and suggests new leads for future studies into the role of steroid hormones in cCSC. These studies might focus on the potential predictive value of the hormone system balance for active phase cCSC and on whether restoring steroid hormone balance might be an avenue for treatment of cCSC.

Footnotes and Financial Disclosures

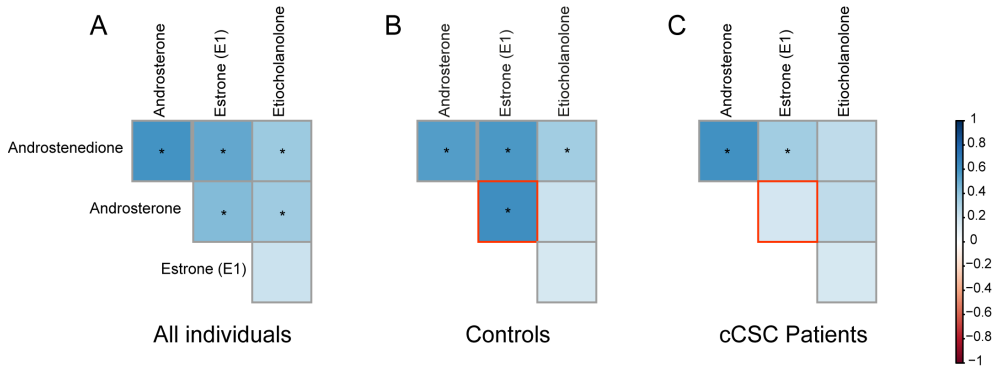
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Conflict of interest: No conflicting relationship exists for any author

Rosa L. Schellevis had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Supplementary Files



Supplementary Figure 1: Correlations between the four associated steroids. Colors indicate positive (blue) and negative (red) correlations between the different hormones, and significant correlations are indicated with an asterisk (*). Red bordered squares are those correlations that were significantly different between cCSC patients and controls (Table 2).

Supplementary Table 1. Differences between Cologne and Nijmegen cohort for the 4 significant steroids

		Concentration in ng/mL				Controls				
		cCSC patients		Nijmegen (n=16)		Cologne (n=10)		Nijmegen (n=36)		
Non-normally distributed hormones		Median	IQR	Median	IQR	Median	IQR	Median	IQR	Mann Whitney U 2-sided P-value
Androsterone		0.948	(0.89-1.15)	0.838	(0.65-0.93)	0.501	(0.43-0.79)	0.713	(0.53-0.97)	0.132
Estrone (E1)		0.128	(0.11-0.15)	0.115	(0.090-0.15)	0.088	(0.075-0.11)	0.104	(0.086-0.12)	0.116
Etiocholanolone		0.202	(0.16-0.33)	0.164	(0.13-0.22)	0.156	(0.11-0.16)	0.119	(0.098-0.22)	0.829
Normally distributed hormones		Mean	Stdev.	Mean	Stdev.	Mean	Stdev.	Mean	Stdev.	Independent T-Test P-value
Androstenedione		3.088	1.00	3.113	1.11	2.353	1.41	2.603	0.80	0.603

cCSC: chronic central serous chorioretinopathy, IQR: Inter quartile range depicting the 25 and 75 percentile, SD: Standard deviation

Supplementary Table 2. Difference between time of blood withdrawal for the 4 associated steroids

Hormone	Grouping value				12-13h (n=7)				14-15h (n=6)				
	8-9h (n=12)	9-10h (n=14)	10-12h (n=8)		Median	IQR	Median	IQR	Median	IQR	Median	IQR	2-sided P-value
Androstenedione	3.38	2.03-4.95	3.19	2.05-3.63	3.05	2.47-3.84	2.62	2.04-3.50	2.88	2.04-3.50	2.88	2.62-3.34	0.738
Androsterone	0.92	0.56-1.08	0.83	0.59-0.96	1.07	0.81-1.24	0.76	0.75-1.05	1.05	0.75-1.05	1.05	0.84-1.19	0.344
Estrone (E1)	0.14	0.08-0.17	0.12	0.10-0.15	0.11	0.11-0.14	0.11	0.09-0.15	0.12	0.09-0.15	0.12	0.09-0.14	0.897
Etiocholanolone	0.19	0.16-0.33	0.15	0.12-0.18	0.20	0.12-0.25	0.18	0.10-0.23	0.20	0.10-0.23	0.20	0.14-0.34	0.393

IQR: Inter quartile range depicting the 25 and 75 percentile, SD: Standard deviation

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Central serous chorioretinopathy, Section I: Basics, Genetics

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6. General Discussion



Although the familial occurrence of CSC already hinted towards a genetic component for CSC in 1960, the first publications linking genetic variants to the occurrence of cCSC only appeared in 2014 just before the start of this thesis. To date, various candidate gene studies and one unbiased genome-wide association study focusing on CSC have been performed. Additionally, exome sequencing in both sporadic and familial CSC patients was performed as part of this thesis. In this general discussion, we present the published studies on the genetics of CSC specified per gene in order of publication date, discuss the robust association between the *CFH* gene and cCSC, and suggest future studies to be conducted.

Candidate gene studies

CFH

Involvement of the *complement factor H (CFH)* gene in CSC was initially investigated because the factor H protein is able to bind adrenomedullin. This molecule is a vasodilator in the choroid important for regulation of choroidal flow, which was previously suggested to be the cause of CSC.¹

The first publication on genetic associations of *CFH* with CSC was reported in a Japanese CSC cohort (n=140) indicating that four variants in the *CFH* gene conferred increased risk for cCSC (rs3753394, rs800292 [I62V], rs2284664 and rs1329428), whereas one (rs106548) was protective for CSC (Table 1).² These associations were confirmed with the same direction of effect in a Dutch cCSC cohort (n=197), although the minor allele frequencies greatly differed and not all variants reached significance after correction for multiple testing (Table 1 and **Chapter 2.1**).³ In addition, two different *CFH* SNPs (rs12144939 and rs1061170 [Y402H]) were also tested in this cohort, but both were not associated with CSC after correction for multiple testing.³ Associations of the same *CFH* variants in a small Greek CSC cohort (n=41) showed an opposite effect for the rs3753394 and rs1065489 variants,⁴ as the respective C and G alleles were protective instead of risk-conferring compared to the Japanese and Dutch populations.^{2,3} In this study, the rs800292 and rs2284664 variants were not associated, while the rs1329428 variant was risk-conferring as observed in both other studies (Table 1).⁴

In the Dutch study, different phenotypes of CSC were also compared, ranging from a very typical group with bilateral CSC (n=197), towards a unilateral CSC-affected group (n=52), and finally a group with a more mixed phenotype (n=43), including signs of neovascularisation and/or drusen on top of the subretinal fluid accumulation.³ Comparison of the *CFH* variants between these groups, indicated that the typical CSC group was genetically distinct from the more admixed group, stressing the importance for extensive and thorough phenotyping of CSC.³

Haplotype analysis of the *CFH* gene was performed in both the Japanese and the Dutch study (Figure 1). In the Japanese study, two different haploblocks were constructed (J1: rs3753394|rs800929 and J2: rs2284664|rs1329428|rs1065489), while in the Dutch study the two blocks were combined and rs1061170 was also included (D: rs3753394|rs800929|rs1061170|rs2284664|rs1329428|rs1065489). The protective J1-H1: TG (OR=0.69) and J2-H1: CCT (OR=0.55) and the risk-carrying J1-H2: CA (OR=1.63), and J2-H2: TTG (OR=1.66) haplotypes were associated with

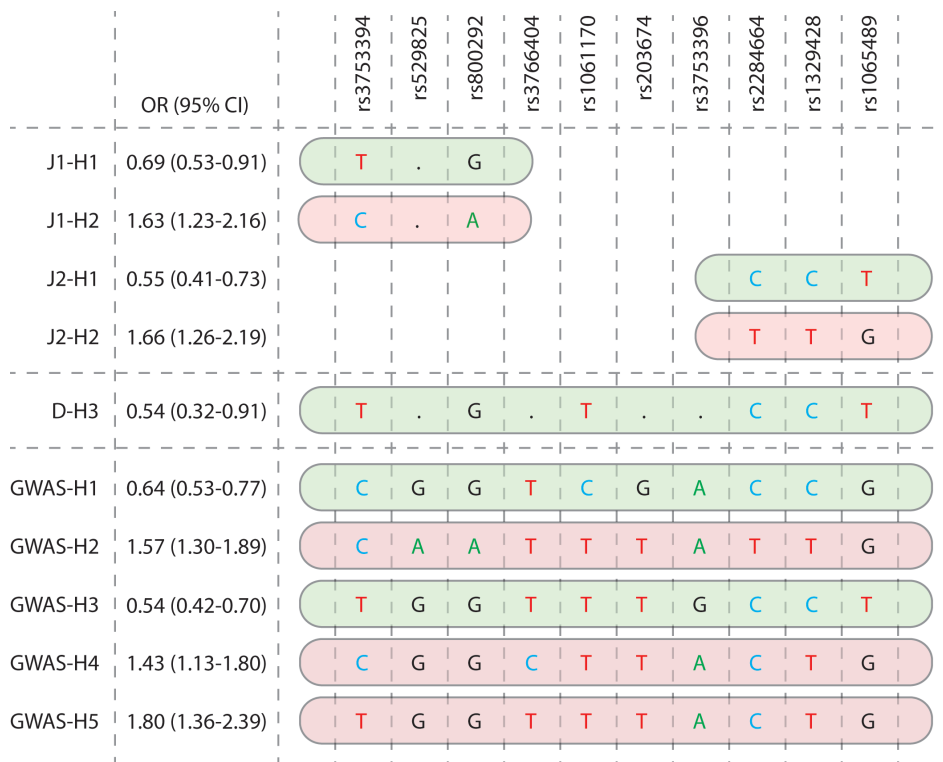


Figure 1: Haplotypes of the *CFH* gene in the Japanese (J1-J2) and Dutch (D) candidate gene studies and the genome-wide association study (GWAS).^{2,3,5} Protective haplotypes are shaded in green, whereas red shaded haplotypes indicate a risk-carrying effect. Nucleotides A/C/T/G are colored for better distinction, and missing genotype information is indicated with a point. OR: Odds ratio, CI: confidence interval.

CSC after correction for multiple testing.² In the Dutch cohort, only the protective D-H3:TGTCCT (OR=0.54) haplotype was associated after correction for multiple testing, whereas the D-H2:CATTG (OR=1.33) haplotype showed a trend towards association.³ Thus, although minor allele frequency differences exist between the Japanese and European population, similar haplotypes were risk-conferring or protective for CSC.

CDH5

In a candidate gene study containing patients from the USA (n=145) and Denmark (n=254), the relationship between CSC and 82 SNPs in 44 genes was tested based on their role in three pathways: stress response, steroid metabolism, and choroidal/epithelial permeability.⁶ Variants with suggestive associations in the American cohort (the p-value cutoff was not defined in this study) in the *cadherin 5* (*CDH5*), *heat shock protein 90* (*HSP90*), *Mitogen-Activated Protein Kinase 1* (*MAPK1*) and *CFH* genes were replicated in the Danish cohort. Of these selected variants, three SNPs in *CDH5* were independently associated to cCSC in both the initial USA and the Danish replication cohorts, with two variants carrying a protective effect (Table 2, rs1130844 [OR=0.78] and rs1073584 [OR=0.77]), and one risk-conferring variant (rs7499886 [OR=1.30]). These associations were reported to be mainly driven by the male subgroup of the

cohort.⁶ It must be noted that the number of SNPs selected for replication was not reported, therefore it is uncertain if these associations remain significant after correction for multiple testing.

Haplotype analysis of the *CDH5* gene was performed with two haploblocks (Figure 2, C1: rs1130844|rs2304527|rs1073584 and C2:rs10852432|rs7499886|rs1073584). The protective C1-H1: CTT (OR=0.61) and C2-H4: TGT (OR=0.66) haplotypes and the risk carrying C2-H3: CAC (OR=1.57) haplotypes were significantly associated to cCSC after correction for multiple testing (5 haplotypes: P<0.01).

The authors suggest that the *CDH5* protein might be involved in the CSC disease mechanism due to its function in endothelial cell-cell junctions, and its localization in the choriocapillaris. However, the genetic associations presented in this study have not been reported in other CSC cohorts, and therefore the involvement of the *CDH5* gene in the disease remains uncertain.

PAI-1

One small study using plasma of 17 CSC patients, showed that levels of the plasminogen activator inhibitor 1 (PAI-1) were elevated in CSC patients.⁷ PAI-1 is an important factor in inhibiting fibrin degradation, but also has been indicated in other processes such as inflammation.⁸

Previous studies have indicated that a G-insertion/deletion polymorphism in the promoter of *PAI-1* (4G/5G) directly influences the plasma PAI-1 levels. Therefore, in order to potentially explain the increased levels of PAI-1 in CSC, this polymorphism was investigated in a Turkish CSC cohort (n=60).⁹ Although the CSC patients group again showed significantly higher serum PAI-1 levels, no significant differences were found in the frequency of the 4G/5G polymorphism between CSC patients and controls (Table 2).⁹ To date, it remains unclear what might cause this PAI-1 elevation, and larger studies investigating a potential role for the *PAI-1* gene in CSC are warranted.

249

ARMS2 and other AMD-associated genes

After the association of *CFH* with CSC was identified, the question arose whether other age-related macular degeneration (AMD)-associated variants are associated with CSC. Therefore, a total of 18 variants previously described to be associated with AMD¹⁰ were investigated in the same Dutch cCSC cohort that replicated the *CFH* association (n=197, Table 2 and **Chapter 2.1**).³ Variants in *age related macular susceptibility 2 (ARMS2)*, *TNF Receptor Superfamily Member 10A (TNFRSF10A)* and *A disintegrin and metalloproteinase with thrombospondin motifs 9 (ADAMTS9)* were associated with CSC. However only the association with rs10490924 (A69S) in *ARMS2* remained significant after correction for multiple testing, conferring a protective effect for the minor T-allele (OR=0.64).³ The association of CSC with *ARMS2* has not been reported in other cohorts, however the rs10490924 variant has been associated with polypoidal choroidal vasculopathy, one of the endophenotypes of CSC (see section: The genetics of CSC endophenotypes).

SGK-1

The protein encoded by the *serum glucocorticoid kinase gene-1 (SGK-1)* is able to regulate several channels and transporters in epithelial cells, including *EnaC*, *SCN5A*,

Table 1. Published associations between the CFH gene and CSC in candidate gene studies

SNP	Location	Miki et al. 2014 ²				de Jong, Breukink et al. 2015, ³				Moschos et al. 2016 ^{4*}						
		Ma/Mi	MAF CSC (n=140)	MAF control (n=934)	P-value	OR (95% CI)	Ma/Mi	MAF CSC (n=197)	MAF control (n=881)	P-value	OR (95% CI)	Ma/Mi	MAF CSC (n=41)	MAF control (n=78)	P-value	OR (95% CI)
rs3753394	Promotor	T/C	0.59	0.49	1.70×10^{-3}	1.50 (1.16-1.94)	C/T	0.25	0.30	2.70×10^{-2}	0.78 (0.60-1.00)	T/C	0.29	0.47	6.80×10^{-3}	0.46 (0.26-0.81)
rs800292	Exon 2 (62V)	G/A	0.53	0.40	6.75×10^{-5}	1.66 (1.29-2.14)	G/A	0.32	0.24	7.50×10^{-4}	1.50 (1.18-1.90)	G/A	0.39	0.45	3.86×10^{-1}	0.79 (0.46-1.36)
rs2284664	Intron 15	C/T	0.50	0.39	7.84×10^{-4}	1.54 (1.19-1.98)	C/T	0.28	0.22	9.00×10^{-3}	1.37 (1.07-1.76)	C/T	0.27	0.36	1.57×10^{-1}	0.66 (0.36-1.18)
rs1329428	Intron 15	C/T	0.59	0.45	6.44×10^{-6}	1.79 (1.39-2.31)	C/T	0.53	0.43	4.60×10^{-4}	1.47 (1.17-1.83)	T/C	0.60	0.38	1.24×10^{-3}	2.44 (1.41-4.22)
rs1065489	Exon 18 (D936E)	G/T	0.35	0.48	6.56×10^{-5}	0.59 (0.45-0.77)	G/T	0.12	0.18	3.00×10^{-3}	0.63 (0.45-0.87)	G/T	0.57	0.42	2.15×10^{-2}	1.88 (1.10-3.23)
rs12144939	Intron 15						G/T	0.25	0.20	3.10×10^{-2}	1.33 (1.03-1.70)					
rs1061170	Exon 9 (Y402H)						T/C	0.31	0.35	6.50×10^{-2}	0.83 (0.66-1.05)					NA

CI: confidence interval; CSC: central serous chorioretinopathy; MAF: minor allele frequency; Ma/Mi: Major/Minor allele; n: number; OR: odds ratio; SNP: single nucleotide polymorphism; Bold p-values indicate significance after correction for multiple testing in the respective publication; *Allelic associations were calculated based on genotype data presented in the publication

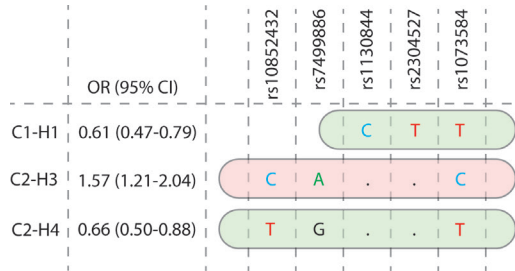


Figure 2: Haplotypes of the *CDH5* gene in the candidate gene study.⁶ Protective haplotypes are shaded in green, whereas red shaded haplotypes indicate a risk-carrying effect. Nucleotides A/C/T/G are colored for better distinction, and missing genotype information is indicated with a point. OR: Odds ratio, CI: confidence interval.

ROMK1, KCNE1/KCNQ1, Kv1.3, NHE3 and the Na⁺/K⁺-ATPase.¹¹ Because of this ability to activate channels and because of its glucocorticoid responsive activation, the involvement of *SGK-1* was investigated in a small Turkish cCSC cohort (n=32). The coding region of the *SGK-1* gene was sequenced, and three variants were identified (Table 2, rs1057293, rs1743966 and rs147657480). Although the rs147657480 variant was only observed in CSC patients and not in controls, none of the identified variants was significantly associated with CSC, potentially due to the small sample size and consequent lack of power.

C4B

Complement factor 4 (C4) is encoded by two nearly identical genes, *C4A* and *C4B*, and plays a role in the classical and lectin pathway of the complement system.¹² Copy numbers of the *C4A* and *C4B* genes have occurred in the genome due to the process of recombination (Box 1 in Chapter 1), leading to a large variation in the amount of copies of these genes in individuals. The amount of *C4A* and *C4B* copies an individual carries has a direct influence on the expression of C4.¹³ Also, the regulation of stress through cortisol release after adrenocorticotrophic hormone (ACTH) stimulation, has been indicated to vary based on the amount of C4 copies.¹³ Therefore, because of the involvement of C4 in both the complement system and the stress pathway, the influence of copy numbers of the C4 gene were studied in relation to cCSC (Chapter 2.2).³

Copy numbers of both the *C4A* and *C4B* gene and the combination of those two were studied. A significant difference in copy numbers of the *C4B* gene was observed in cCSC patients (n=197) compared to controls. A complete absence of *C4B* (*C4B**Q0) conferred risk for CSC, with the largest OR reported for a genetic variant in CSC so far (OR=2.61), while carrying three copies of the gene was protective for cCSC (OR=0.45).³ The association of the *C4B* gene with cCSC has so far not been reported in other cohorts.

GSTM1

Because of its role in oxidative stress, and potential role in choroidal vasoconstriction, variants in the *Glutathione-S-Transferase M1* (*GSTM1*) gene were studied in a Greek CSC cohort (n=41).⁴ However, in this small cohort the frequencies of the normal and null alleles of the *GSTM1* polymorphism were not significantly different between CSC patients and controls (Table 2).⁴

NR3C1 and NR3C2

The *NR3C1* and *NR3C2* genes encode the glucocorticoid and mineralocorticoid receptors, respectively. Because of the known association of stress and use of corticosteroids with CSC, genetic variants that influence the function of these receptors were tested in a Dutch cCSC cohort (Table 2, n=336, **Chapter 2.3**).¹⁴ Three variants in *NR3C1* (rs56149945, rs41423247, and rs6198) and one variant in *NR3C2* (rs5522) were not associated with cCSC. However, the rs2070951 variant in *NR3C2* was associated with an increased risk for cCSC (OR=1.29). Haplotype analysis of the two studied variants (Figure 3, N: rs2070951|rs5522) identified a protective effect for the N-H1: CA haplotype (OR=0.72), while N-H3: GA conferred risk for CSC (OR=1.39).

The association of the rs2070951 variant and *NR3C2* haplotypes have not been confirmed in other CSC cohorts, however, a GWAS on choroidal thickness indicated a nominal significant association for another variant in *NR3C2* (rs10519952, P = 0.03) with this CSC endophenotype.¹⁵

Unbiased studies

Genome-wide association analysis

Most studies published so far have used a candidate gene approach, however recently the first genome-wide association study for cCSC has been performed (**Chapter 3.1**).⁵ In this study, the allele frequencies of ~11 million SNPs were compared between 521 European cCSC patients and 3577 population controls. A genome-wide significant association with variants at the *CFH* gene was identified, and 6 variants with suggestive association ($P < 1 \times 10^{-6}$) were observed.⁵

Haplotypes of *CFH* were extended compared to the previous publication of the same group (Figure 1, GWAS:rs3753394|rs529825|rs800929|rs3766404|rs1061170|rs203674|rs3753396|rs2284664|rs1329428|rs1065489).³ The GWAS-H2:CAATTTATTG, GWAS-H4:CGGCTTACTG, GWAS-H5:TGGTTTACTG, all containing the minor allele of the top-GWAS variant (bold) were risk carrying for cCSC (OR=1.57, 1.43 and 1.80, respectively), whereas the GWAS-H1: CGGTTCGACCG and GWAS-H3: TGGTTTGCCT containing the wild-type allele were protective (OR=0.64 and 0.54, respectively).⁵

In the same publication, gene expression in the cCSC patients and controls was predicted using expression quantitative trait loci (eQTLs) data of the Genotype-Tissue Expression (GTEx) database. Significant differences of predicted expression were found for genes in the *CFH* locus, namely, *CFH* itself, but also *CFH related 1 and 4* (*CFHR1/4*), and the nearby *Potassium Sodium-Activated Channel Subfamily T Member 2* (*KCNT2*) gene.⁵ Additionally, the *cluster of differentiation 46* (*CD46*) and *tumor necrosis factor receptor superfamily member 10a* (*TNFRSF10A*) genes located elsewhere in the genome were predicted to be differentially expressed. The predicted altered expression makes *CFHR1*, *CFHR4*, *CD46* and *TNFRSF10A* interesting candidate genes for future studies on cCSC. Finally, pathway analyses of the data indicated the involvement of the complement cascade in cCSC.⁵

Table 2. Published associations of genetic variants with CSC

Gene	Accession number	SNP	Location	Chr:BP (hg19)	Ma/ Mi allele	N. CSC/ controls	MAF CSC	MAF control	Nominal Allelic P-value	OR (95% CI)
Schubert et al. 2014, Human Mutation ⁶										
CDH5	NM_001795.4	rs10852432	Intron 1	16:66402515	C/T*		0.50	0.53	0.080	0.87
CDH5	NM_001795.4	rs23444564	Intron 1	16:66413150	C/T*		0.29	0.25	0.060	1.22
CDH5	NM_001795.4	rs7499886	Intron 1	16:66413195	A/G*	399/1421	0.48	0.41	0.001	1.3
CDH5	NM_001795.4	rs1130844	Exon 3 (Gly-128Gly)	16:66420885	C/T*		0.42	0.49	0.002	0.78
CDH5	NM_001795.4	rs1073584	Intron 8	16:66430467	T/C*		0.26	0.31	0.004	0.77
Sogutlu Sari et al. 2014, Cutan Ocul Toxicol ⁹										
PAI-1		rs1799768/ rs1799889		7:100769706/ 7:100769711	4G/5G	60/50	0.45	0.45	1	1.00 (0.59-1.71)
de Jong & Breukink et al. 2015, Ophthalmology ³										
ARM52	NM_001099667.2	rs10490924	Exon 1 (Ala-69Ser)	10:124214448	G/T		0.17	0.24	0.002	0.64 (0.49-0.85)
SKIV2L	NM_006929.4	rs429608	Intron 11	6:31930462	G/A		0.16	0.13	0.133	1.25 (0.94-1.68)
C3	NM_000064.2	rs2230199	Exon 3 (Arg-102Gly)	19:6718387	G/C		0.19	0.17	0.393	1.16 (0.83-1.61)
SYN3	NM_003490.3	rs9621532	Intron 6	22:33084511	A/C	292/1311	0.05	0.05	0.807	1.05 (0.65-1.68)
APOE	-	rs4420638	Downstream	19:45422946	A/G		0.16	0.17	0.470	0.89 (0.66-1.19)
CETP	-	rs3764261	Upstream	16:56993324	G/T		0.30	0.32	0.484	0.92 (0.73-1.16)
VEGFA	-	rs943080	Downstream	6:43826627	T/C		0.44	0.48	0.211	0.87 (0.70-1.08)
TNFRSF10A	-	rs13278062	Upstream	8:23082971	T/G		0.40	0.48	0.004	0.73 (0.59-0.90)
LIPC	-	rs493258	Upstream	15:58687880	C/T		0.49	0.46	0.064	1.23 (0.99-1.52)

CFI	-	rs10033900	Downstream	4:110659067	C/T	0.50	0.48	0.514	1.08 (0.87-1.34)	
COL10A1	NM_000493.3	rs3812111	Intron 2	6:116443735	A/T	0.34	0.36	0.498	0.92 (0.74-1.15)	
COL8A1	NM_001850.4	rs13081855	Intron 3	3:99481539	G/T	0.12	0.09	0.074	1.38 (0.98-1.93)	
IER3-DDR1	-	rs3130783	Upstream	6:30774357	A/G	0.20	0.19	0.836	1.03 (0.78-1.34)	
SLC16A8	NM_013356.2	rs8135665	Intron 4	22:38476276	C/T	292/1311	0.21	0.22	0.600	0.92 (0.71-1.20)
TGFBR1	NM_004612.2	rs334353	Intron 6	9:101908365	T/G	0.26	0.25	0.662	1.06 (0.83-1.35)	
RAD51B	NM_002877.5	rs8017304	Intron 8	14:68785077	A/G	0.36	0.38	0.467	0.92 (0.73-1.14)	
ADAMTS9	-	rs6795735	Upstream	3:64705365	C/T	0.46	0.41	0.047	1.25 (1.01-1.54)	
B3GALT1	NM_194318.3	rs9542236	Intron 4	13:31819325	T/C	0.44	0.44	1	1.01 (0.81-1.25)	
Akyol et al. 2015, Int J Ophthalmology ¹¹										
SGK-1	NM_001143676.1	rs1057293	Exon 10 (Asp-p335Asp)	6:134493397	C/T	0.06	0.05	1	1.36 (0.29-6.32)	
SGK-1	NM_001143676.1	rs1743966	Intron 8	6:134493947	A/G	32/32	0.22	0.12	0.160	1.96 (0.76-5.06)
SGK-1	NM_001143676.1	rs147657480	Exon2 (Met-32Val)	6:134583262	A/G	0.03	0.00	0.496	NA	
Moschos et al. 2016, Retina ⁴										
GSTM1	NM_000561.3	null allele (tag: rs366631)	Complete gene	1:110230418-110236367	Normal/Null	41/78	0.50	0.722	0.87 (0.41-1.87)	
van Dijk, Schellevis et al. 2017 ¹⁴										
NR3C1	NM_000176.2	rs56149945	Exon 2 (Asn363Ser)	5:142779317	A/G	0.04	0.04	0.500	0.86 (0.55-1.34)	
NR3C1	NM_000176.2	rs41423247	Intron 2	5:142778575	C/G	336/1314	0.39	0.36	0.150	1.14 (0.95-1.35)
NR3C1	NM_000176.2	rs6198	Exon 9 UTR	5:142657621	A/G	0.16	0.17	0.480	0.92 (0.73-1.16)	
NR3C2	NM_000901.4	rs2070951	c. -2	4:149358014	C/G	0.53	0.46	0.004	1.29 (1.08-1.53)	
NR3C2	NM_000901.4	rs5522	Exon 2 (Val-180Ile)	4:149357475	A/G	0.14	0.13	0.570	1.08 (0.83-1.38)	

*Minor allele not specified in study; BP, base pair; CSC: central serous chorioretinopathy; Chr: chromosome; CI: confidence interval; Ma: major; Mi: minor; N: number; OR: odds ratio

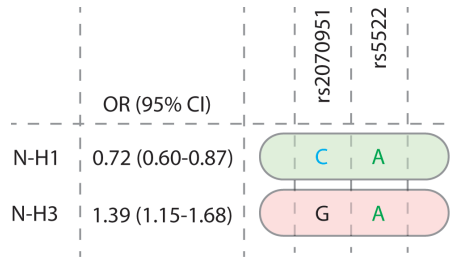


Figure 3: Haplotypes of the *NR3C2* gene in the candidate gene study.¹⁴ Protective haplotypes are shaded in green, whereas red shaded haplotypes indicate a risk-carrying effect. Nucleotides A/C/T/G are colored for better distinction. OR: Odds ratio, CI: confidence interval.

Exome sequencing analyses

For this thesis the first exome sequencing studies on cCSC were performed on both a case-control cohort and in a family-based study (**Section 4**). So far, no other studies have been performed using exome sequencing on cCSC patients.

The analysis of exome sequencing data of the cCSC case-control cohort was performed on 263 European cCSC patients and 1352 population controls and mainly consisted of gene-based analyses of the data (**Chapter 4.1**). A significant association was observed with cCSC in females for the *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* genes. Almost half of the female cCSC patients (45.5%) carried at least 1 rare variant in one of these genes compared to a carrier rate of 18.5% in female controls. In this study, we did not observe any significant associations with cCSC in the entire cohort nor in the males only after correction for multiple testing. Additionally, no gene-burden associations were observed in the previously associated cCSC genes and loci described in the **Candidate gene studies** section of this discussion. The sex-specific association of rare variants in the *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* genes suggest that a different disease mechanism might underlie CSC in females compared to males, but replication of these results in other cohorts with cCSC is warranted.

255

For the family-based data analysis, 72 individuals of 18 families in which CSC was present were analyzed by exome sequencing (**Chapter 4.2**). In total, 12 different segregating variants were present in 2 families. Of these, a variant (rs61758735) in the *PTPRB* gene was also significantly associated with CSC in the case-control exome sequencing study. Segregation of rs61758735 in additional family members of one of the families showed partial segregation with the phenotype, as also previously observed for rare variants associated with familial AMD.^{16,17} The function of the VEPTP protein encoded by *PTPRB* in angiogenesis and endothelial barrier functioning makes it an interesting candidate gene for CSC. Gene burden analysis of the family data did not identify the genes as in the case-control cohort. In this study, our analysis of familial CSC did not confirm the proposed Mendelian inheritance for CSC, but suggests an interaction between genetics and the environment, which would rather be in line with a complex disease. Extension of genetic studies on familial CSC might confirm the complex inheritance, or might increase the chance of finding multiple families carrying segregating variants in the same gene that could support a Mendelian inheritance instead.

The genetics of CSC endophenotypes

A disease generally is a composite of several features, or traits, that can be evaluated independently from the disease itself. These traits, or endophenotypes of a disease, can be tested for genetic associations, revealing potential susceptibility loci for the disease as a whole. This approach has been successful in other eye diseases such as myopia, where large-scale genetic association studies on several endophenotypes of myopia including axial length,¹⁸ refractive error,¹⁹ and corneal curvature,²⁰ discovered several loci of interest for myopia.

One of the main endophenotypes for cCSC is choroidal thickness. Although it is not apparent in all CSC cases, thickening of the choroid in particular has been reported to be a hallmark of the disease. Because of this connection, CSC has been regarded as part of the pachychoroid spectrum of diseases.²¹ Pachychoroid is a feature that can be measured in OCT images, with enhanced depth imaging and several studies have been published that have measured choroidal thickness in normal individuals.²²

So far only two groups have investigated whether pachychoroid itself, independent of diagnosed CSC, is a heritable trait. In one prospective observational study relatives (n=16) of CSC patients (n=5) were given an extensive eye exam, including detailed retinal imaging.²² In 50% of the eyes of relatives of CSC patients, pachychoroid could be observed and this feature seemed to be dominantly inherited, independent of a clinical diagnosis of CSC.²² A second study sought to quantify the level of heritability of the pachychoroid feature and did so using extended Amish families with early/intermediate AMD. Choroidal thickness was measured in almost 700 individuals of these families and was shown to be transmitted across generations independent of AMD status. The authors concluded that choroidal thickness was moderately heritable with a heritability score of 0.40, but did not evaluate genetic associations with thick choroid.²³

Candidate gene studies

Because of the tight links between choroidal thickness in AMD, CSC and polypoidal choroidal vasculopathy (PCV), and the respective genetic associations to *CFH* and *ARMS2* in those diseases, most genetic studies exploring associations to choroidal features used variants in *CFH* or *ARMS2* as the starting point. The first of these studies investigated the link between choroidal thickness and vascular hyperpermeability and polymorphisms in *CFH* and *ARMS2* in patients with PCV (n=63) and AMD (n=58).²⁴ This study reported an association between an increased choroidal thickness and the minor-A allele of the variant rs800292 (I62V) in *CFH* in PCV patients. The rs10490924 (A69S) variant in *ARMS2* and rs1061170 (Y402H) in *CFH* were not associated to choroidal thickness in this study.²⁴

A similar study also investigated associations with choroidal vascular hyperpermeability and choroidal thickness, testing *ARMS2* and *CFH* variants in 149 patients with PCV.²⁵ Increased choroidal thickness and vascular permeability in these PCV patients was associated with the major-G allele of the rs10490924 (A69S) variant in *ARMS2* and the minor-T allele of rs1329428 in *CFH*. The rs800929 (I62V) in *CFH* was not associated to any of the traits.²⁵

One study investigated choroidal blood flow regulation in young healthy subjects

(n=96).²⁶ This study showed that homozygous carriers of the AMD-risk genotype (CC) of the rs1061170 (Y402H) variant in *CFH* demonstrated impaired choroidal blood flow after squatting exercises. The authors argue that dysregulation of the choroidal blood stream, in part related to a person's genotype, could predispose them, from an early age, to precipitating events leading to ocular disease.²⁶ However, due to absence of overlap between associations in the above mentioned studies, no solid conclusions can be drawn about the involvement of the *CFH* and *ARMS2* variants in choroidal thickness based on these candidate studies.

Genome-wide association study

Only one large study investigated the genetic underpinnings of choroidal thickness in an unbiased, hypothesis-free manner by performing a GWAS on choroidal thickness in 3418 individuals.¹⁵ This work made use of the Japanese Nagahama Study in which, among others, detailed ophthalmic examinations including measures of choroidal thickness were coupled to available genome-wide genotyping information. One locus in *CFH* (top-associated variant: rs3753394) was discovered to be associated to choroidal thickness, while another novel locus in the *VIPR2* (top-associated variant: rs7782658) gene showed a suggestive association. SNPs that showed moderate linkage disequilibrium with the top-associated variants (*CFH* [rs800292] & *VIPR2* [rs3793217]) were replicated in 2692 independent individuals from the same Nagahama study. In the meta-analysis both loci reached genome-wide significance.

Additionally, these SNPs were tested for association in a Japanese (n=539/701) and Korean (n=425) CSC group including both acute and chronic patients. Both rs800292 and rs3793217 were associated with CSC, although these associations did not reach genome-wide significance.¹⁵ Since the *CFH* association was recently identified in a GWAS for CSC and has been shown in multiple candidate studies,²⁻⁵ this finding can be considered robust. However, the *VIPR2* association to CSC warrants replication in other independent cohorts, especially since the odds ratio of this variant in the Japanese study is relatively small (OR=1.24) and the allele frequency varies greatly among different populations (range minor allele frequency: 0.04-0.18).²⁷

257

Discussion

Although the field of CSC-genetics is still emerging, we have observed a growth in studies on the genetics of CSC during the last 5 years. The studies described in this thesis confirm the genetic component contributing to the disease, and pave the way for future studies to understand the etiology of CSC.

Although various candidate genes for CSC have been identified, the associations with variants in *CDH5*, *ARMS2*, *C4B*, *NR3C2*, *VIPR2*, *CD46*, *KCNT2*, *TNRSF10A*, *CFHR1/4*,^{3,5,6,14,15} *LAMB3*, *DUOX1*, *RSAD1*, *PIGZ* and *PTPRB* need to be tested in independent cohorts in order to be confirmed. Therefore, the implications of these associations are not discussed in detail here.

In this section of this general discussion we focus on the robust genetic association of *CFH* with CSC in light of current literature. We elaborate on questions that still need to be answered regarding the etiology of CSC and propose studies that can be conducted to shed more light on the disease mechanisms underlying CSC.

The role of CFH in CSC

The genetic association of the *CFH* gene with CSC has been proven in multiple candidate gene studies and one genome-wide association study,²⁻⁵ indicating that *CFH* is involved in the development of CSC, albeit via a still unknown mechanism. The *CFH* gene encodes the factor H (FH) protein, which is an inhibitor of the complement system, but FH also has been implicated in adrenomedullin (AM) binding. Here, we briefly discuss the potential role of FH in the CSC disease mechanism.

The complement system

The FH protein is most widely known for its function in preventing over activation of the complement system by blocking the formation of C3-converstases.¹² Together with the cofactor factor I, another complement inhibitor, it regulates the feedback loop of the alternative pathway of the complement system.¹² FH consists of a stretch 20 complement control protein (CCP) domains, which are also referred to as short consensus repeats (SCRs). The N-terminal part of FH (CCP1-5) is thought be involved in complement regulation, whereas the C-terminal domains mainly regulate cell surface binding (CCP19-20).²⁸

In the GWAS study on CSC, the genome-wide signal in *CFH* was driven by 20 SNPs that are in high linkage disequilibrium.⁵ Most of these SNPs are located in introns 9-15, and one synonymous SNP is located in exon 10 (rs2274700), suggesting that they may have a regulatory effect on gene expression, or they may be co-inherited with other (yet unidentified) protein-altering variants on the same haplotype. The specific function of the protein domains (CCP10-15) around which the 20 variants cluster is still obscure, but because eQTL data in the same study predicts that *CFH* and surrounding genes are differentially expressed in multiple tissues, a regulatory role of these variants seems likely.

So far, no rare protein-altering variants in the *CFH* gene have been associated with CSC. However, the common variants and haplotypes associated with CSC have been studied before in relation to AMD, which is the most common cause of vision loss in the elderly population characterized by lipid accumulations between the Bruch's membrane and RPE.²⁸ Interestingly, their direction of effect appears to be opposite between both diseases; variants protective for AMD are risk-conferring for CSC and vice versa, suggesting an opposite disease mechanism for both conditions.^{2,3,5} As over-activation of the complement system is thought to underlie AMD development, this suggests that the opposite mechanism in CSC would be reduced activity of the complement system. Associations of other complement genes besides *CFH* in the *CFHR1*, *CFHR4*, *C4* and *CD46* genes identified in this thesis certainly suggest an important role for the complement system in the disease. Currently, studies on complement measurements in CSC are limited and have not confirmed this hypothesis yet,¹⁴ therefore measurements in larger cohorts and of additional complement components need to be conducted to test the role of the complement system in CSC.

Adrenomedullin

Besides its function in the complement system, the first hypothesis that FH might have a function in the CSC disease mechanism was suggested based on its ability to bind to adrenomedullin (AM).¹ This regulatory peptide is expressed by many cells,

including the RPE, is sensitive to corticosteroid stimulation, and is an important factor in blood pressure homeostasis and vasodilation, making it a very interesting molecule in light of the pathogenesis of CSC.^{1,29}

The CCP 15-20 domains of the FH protein are thought to be involved in binding of AM, which stabilizes AM and prevents it from degradation by metalloprotease 2, while also increasing the C3b-conversion ability of FH itself.²⁹ Studies in rats indicate that administration of AM leads to reduced pro-inflammatory signals, accelerated wound healing and improved gut barrier function after ischemia.²⁹ Although the role of AM in the CSC disease mechanisms was speculated already in 2003,¹ no research, besides the *CFH* associations, has been published so far linking the *ADM* gene or AM protein to CSC, therefore it remains to be determined if this molecule via binding of FH has an influence on the development of CSC.

Rare variants in *CFH*

Rare protein-altering variants in *CFH* have been associated to a range of diseases namely C3 glomerulopathy (C3G), characterized by kidney failure due to accumulation of C3 in the glomeruli; atypical hemolytic syndrome (aHUS), characterized by acute renal failure, microangiopathic hemolytic anemia and thrombocytopenia, and AMD.²⁸ Rare protein-altering variants identified in AMD tend to cluster at the N-terminal part of the protein in CCP3/5/7, indicating an effect on complement interactions, whereas aHUS/C3G mutations were more frequent in the SCR19/20 domain at the C-terminus, suggesting loss of binding interactions with cells.²⁸

In our exome sequencing study (**Chapter 3.2**), we did not observe a genome-wide significant association with a rare variant in *CFH* for cCSC (data not shown). The gene-burden analysis of the *CFH* gene also did not show significant associations with the gene, but showed nominally significant associations with the neighboring *KCNT2* and *CFRH5* genes. The absence of an association for *CFH* might be explained by the limited sample size of the exome sequencing study and the accompanied limited power to find associations, especially with rare variants. Besides the power issues, we can hypothesize that the association of *CFH* with cCSC might be driven by the common variants and their effect on gene-expression levels of *CFH* and neighboring genes. Another option is that the rare variants associated with cCSC are not located in *CFH* itself, but that the associated *CFH* haplotypes actually tag alterations of the neighboring *CFHR* genes, which are poorly covered in the exome sequencing study due to their repetitive nature. Deep sequencing of the region with for example molecular inversion probes and next-generation sequencing might shed light on potential rare variants in these genes, but first steps should focus on increasing sample size.

Data acquisition and analysis of whole exome sequencing data comes with a set of challenges. When adding additional samples to the exome sequencing study various bioinformatics steps need to be taken into account. The GATK pipeline used to process the data in our studies includes an n+1 option for data, meaning that new samples can easily be joined to the dataset. However, one must take note to try to minimize the differences between different batches of sequencing data. Read depth, enrichment kit version, sequencing site and machinery all have to be as similar as possible, otherwise the possibility of false positives greatly increases. After merging the datasets it is still necessary to perform extended quality control on for example

population substructure or kinship to achieve homogeneity between the different batches, before association analyses can be performed. Even then, confirmation of variants with Sanger sequencing is recommended, especially for insertions and deletions, which are in general more difficult to call.

Steroid hormones in cCSC

The most described risk-factors for CSC are the male sex, the use of steroids and the presence of stress (Table 1 in **Chapter 1**). Both the stress- and sex-hormones are members of the steroid hormone system. Conversion of cholesterol by various enzymes produces hormones of three main pathways, namely, aldosterone involved in water homeostasis, the stress-hormones cortisol and cortisone, and finally the various sex-hormones with their androgenic, estrogenic and progestinic effects.³⁰

In **Chapter 5**, the only chapter of this thesis focusing on non-genetic risk factors, we investigated the steroid hormone status in cCSC patients. To date, a limited number of studies have looked at steroid hormone levels in individuals affected with cCSC, and the results are not very consistent between studies (Table 3).³¹⁻⁴³ All previous studies focused on one or at maximum four steroids simultaneously, whereas we chose to measure 17 steroid hormones at the same time. Measuring multiple steroid in the same sample simultaneously allows for the assessment of interactions between steroids, is more cost efficient than measuring hormones separately and gives an overview of the complete hormone system balance with only one test. We observed elevated levels of androsterone, androstenedione, estrone and etiocholanolone, in patients with an active episode of cCSC. These hormones are all members of the sex-hormone section of the steroid hormone cascade, and have not been associated with cCSC before. Although it still remains to be elucidated what the role of these hormones is in the occurrence of CSC, these results shed the first light on the involvement of sex-hormones in the etiology of cCSC.

It is perhaps surprising that no association with cortisol or aldosterone was observed in our steroid study, as these hormones are thought to be involved in the occurrence of CSC.^{44,45} Absence of association with cortisol might be explained by the current study set-up. For this study we used a single time point measurement, and as cortisol is known to fluctuate during the day, multiple measurements of the same individual at different time points might shed more light on the levels of this hormone in active cCSC. For aldosterone, although the levels itself were not altered in cCSC patients, we did observe an altered correlation of aldosterone with several other hormones in the system. Therefore, we hypothesize that there might be a disruption in the balance of the entire system, which might have an effect that leads to the cCSC phenotype, via a still unknown mechanism. It is however necessary to replicate these results in a larger sample of cCSC patients.

Unanswered questions and future studies

Because the number of studies focused on the genetics of CSC is still limited, many questions still remain unanswered. In this section, we briefly address some of the remaining questions and suggest possible studies in which they might be answered.

Table 3. Studies on hormone levels in CSC patients performed till September 2018

Hormone	Study	Controls (n)		CSC patients (n)		Reported levels		Acute CSC	Chronic CSC	Case-control P-value
		Controls (n)	Men/Women	Chronic	Acute	Controls	Chronic CSC			
Aldosterone	38	-	50/11	-	-	-	143.0 ± 53.6 ng/L	-	-	-
Aldosterone	35	32	Men only	15	15	9.73 ± 1.29 ng/L	6.36 ± 0.67 ng/L	6.83 ± 1.28 ng/L	0.108	0.108
Cortisol (7AM)	36	54	44/11 both	54	54	11.73 ± 4.62 µg/dl	14.03 ± 6.6 µg/dl	-	-	0.039
Cortisol (8-9AM)	40	-	16/1 & 12/1	17	13	-	12.79 ± 5.38	12.76 ± 3.77	-	-
Cortisol (8-9AM)	35	32	Men only	15	15	9.50 ± 0.59 µg/dl	10.48 ± 0.98 µg/dl	9.54 ± 0.50 µg/dl	0.494	0.494
Cortisol (8AM)	34	30	Men only	30	30	18.76 ± 8.26 µg/dl	29.97 ± 12.44 µg/dl	-	-	<0.001
Cortisol (8AM)	33	12	11/1 & 22/1	23	23	362.25±51.54 nmol/L	495.02±169.47 nmol/L	-	-	0.002
Cortisol (8AM)	37	37	??	37	37	8.53 µg/dl	8.76 µg/dl	8.76 µg/dl	0.975	0.975
Cortisol (8AM)	32	-	Men only	-	6	-	-	11.6 (4.74-18.3) µg/dl	-	-
Cortisol (4PM)	37	37	??	37	37	6.07 µg/dl	6.13 µg/dl	-	-	0.948
Cortisol evening (11PM)	36	54	44/11	54	54	10.96 ± 4.29 µg/dl	13.1 ± 6.49 µg/dl	-	-	0.047
Cortisol (11PM)	34	30	Men only	30	30	13.06 ± 6.9 µg/dl	22.03 ± 11.54 µg/dl	-	-	<0.001
Cortisol	43	41	?? & ??	44	44	??	296.53 +/- 77.03 ng/ml	-	-	<0.05
Cortisol (24h urine)	34	30	Men only	30	30	7.39 ± 2.47 mg/day	11.01 ± 4.14 mg/day	-	-	<0.001
Cortisol (24h urine)	39	16	14/2 both	16	16	115.3 +/- 63.4 nmol/l	188.20 +/- 34.1 nmol/l	-	-	<0.05
Cortisol (24h urine)	31	??	?? & 19/5	24	24	28.4 ± 3.6 µg/ml	36.7 ± 25.1 µg/ml	-	-	0.17
Cortisol (24h urine)	42	24	19/5 & 7/7/9	-	86	51.55 (28.49) nmol/24h	-	83.99 (49.04) nmol/24h	<0.000	<0.000
DHEA	35	32	Men only	15	15	293.96 ± 45.01 µg/dl	342.82 ± 60.91 µg/dl	369.10 ± 68.24 µg/dl	0.297	0.297
DHEAS	41	40	27/13 & 36/10	46	-	249.36 ± 122.93 µg/dl	203.79 ± 84.75 µg/dl	-	-	0.046
Testosterone	35	32	Men only	15	15	255.94 ± 7.43 ng/ml	351.67 ± 15.56 ng/ml	362.33 ± 14.41 ng/ml	<0.001	<0.001
Testosterone	40	-	16/1 & 12/1	17	13	-	5.22 ± 1.54 ng/dl	5.79 ± 1.13 ng/dl	-	-
Testosterone	33	12	11/1 & 22/1	23	23	4.23±1.89 ngm/ml	3.85±1.81 ngm/ml	-	-	0.58
Testosterone	32	-	Men only	-	6	-	-	549.5 (246-794) ng/ml	-	-

Immune system-hormone system interactions

Within this thesis we have found multiple lines of evidence that the complement system, an important part of our innate immune system, is associated with cCSC, and that the system might be downregulated in cCSC patients. Additional lines of evidence of involvement of the immune system in CSC come from the association of CSC with stress and the use of steroids.⁴⁴ Prolonged or repeated stress stimuli, and use of exogenous glucocorticoids are known to suppress the immune system and inflammatory responses.⁴⁶ Thus, multiple associations suggest that a downregulation of the immune system might underlie the biological mechanism of CSC.

A significant lowering of the immune system activity would however result in a decreased resistance to pathogens as well. Although CSC patients are generally healthy besides their visual complaints, the increased frequency of *Helicobacter pylori* infection in CSC patients might suggest that this lowered immune system activation makes patients more susceptible to infections.⁴⁷⁻⁵¹ However, instead of a significant downregulation giving rise to increased infection rates, it is more likely that the changes in the immune system are more subtle or maybe even only locally affecting the eye. Because the retina has a high metabolic turnover and energy demand, subtle changes in the biological mechanisms underlying these processes can lead to detrimental changes, as might also be the case for CSC. As studies on the immune system activity in CSC patients have not been performed on large scale yet, the exact status of the immune system in CSC patients still remains to be elucidated. Studies of aqueous humor or subretinal fluid, rather than systemic measurements in the blood, might be instrumental in discovering local effects present in the eye specifically.

It is hypothesized that the interaction between stress and the immune system is partly mediated by the levels of glucocorticoids in the blood stream. Low levels of glucocorticoids (acute stress) will primarily occupy the MR, which has the highest binding affinity for GCs, whereas higher levels of GCs (chronic stress) will also promote binding to the GR when the MR is saturated, initiating downstream immunosuppressive processes mediated by the GR.⁴⁶ However, the exact mechanism of these interactions and if they might be tissue-specific is not yet known.⁴⁶ Interestingly, overactivation of the MR by glucocorticoids has been indicated as one of the underlying causes of CSC and currently clinical trials are performed to investigate the efficacy of eplerenone, a MR antagonist, as treatment for CSC.^{52,53} In our analysis of steroids in Chapter 5, however, we did not observe significant elevation of glucocorticoids or mineralocorticoids in patients with active cCSC, but did observe alterations in correlations with aldosterone and other steroid hormones in the system.

Phenotypic differences in CSC

Currently, there is not a clear consensus of what defines the different CSC phenotypes, where some studies use an arbitrary threshold of 3-6 months to distinguish acute from chronic CSC, others use the amount of RPE alterations as differentiating factor.⁴⁴ In **Chapter 2.1**, we proposed new phenotypic definitions for groups of patients with chronic CSC. We observed that genetic differences could be identified between the most typical form of cCSC and a group of patients that showed signs of other retinal diseases together with CSC. This study underscores the necessity of accurate phenotyping for cCSC. Therefore, for future studies, it is important that a

consensus is reached about the different phenotypes observed in the CSC spectrum, and that these separate phenotypes are taken into account when conducting genetic association analyses.

It is still unclear why some people are affected with acute CSC and others are immediately affected with the more severe chronic CSC phenotype or why some people progress from acute to chronic CSC. In **Chapter 2.4** we performed, to our knowledge, the first genetic analysis on acute CSC patients specifically. In this study we did not observe a genetic difference between acute and chronic CSC for the currently identified genetic risk variants. Although risk effects for acute CSC seemed higher than compared to chronic CSC patients, these results were not significantly different between both groups. This suggests that, based on the currently identified genetic variants, the biological mechanism of both forms of the disease might be similar.

However, we cannot exclude that not yet identified genetic factors specific for acute or for chronic CSC might exist. When identified, these genetic variants might be used to distinguish the two forms of the disease, which might have clinical implications for treatment options for patients. One can hypothesize that multiple genetic variants combined could make a person more likely to develop chronic CSC instead of acute CSC and that for acute CSC, an environmental trigger might be more important for disease onset. Therefore, it might be interesting to investigate how the genetic burden of an individual interacts with potential environmental triggers, and might make some individuals more susceptible for one form of CSC compared to the other.

The male-female difference

To date, genetic studies have not shed light on the difference in incidence of CSC in males (9.9:100,000) compared to females (1.7:100,000).⁵⁴ For the studied *CDH5* variants, it was reported that the association was mainly driven by the male CSC patients. However, due to the low number of females included in the study (n=109),⁶ it is likely that there was not enough power to conclusively determine absence of association in the female cohort and the results need to be validated. In the single GWAS performed on cCSC in **Chapter 3.1**, the X-chromosome was included in the analysis, however no associations were found on this chromosome and no sex-specific associations were reported.⁵ In our exome sequencing study in **Chapter 3.2**, we observed a genome-wide significant association in female cCSC patients specifically. This suggests that cCSC might be more genetically heterogeneous in male cCSC patients and larger samples sizes are required to find significant associations in this group.

It has been reported that sex differences occur in many organs, including the eye, but there is still much uncertainty about the role of gonadal hormones on these tissues. One recent review has summarized current literature about sex-differences in the eye and indicates a neuro-protective role for estrogens.⁵⁵ As certain physiological processes in the eye appear to be different in males and females, we need to take into account that the observed sex-differences in CSC could occur due to an interaction between intrinsic sex-differences due to hormones, environmental exposures, and genetic susceptibility.

In our analysis of steroid hormones in Chapter 5 we observed a significant association of four hormones of the sex-hormone system in males. Confirmation of these associations in female CSC patients would be very interesting to ascertain whether the hormone system is similarly altered in this group. Additionally, it would be very interesting to compare measurements of the same individual during an active phase of disease and an inactive phase, either after spontaneous resolution of the symptoms or after treatment. In this way the balance of the hormone system might be used as a predictive marker for disease status or treatment response. Also new treatment modalities might be developed in order to resolve the CSC episode by directly targeting the hormone system imbalance.

The genetic component of CSC

After the completion of this thesis, the inheritance pattern of CSC still remains partly obscure. On one hand family pedigree data suggests a Mendelian inheritance pattern, while on the other hand the predominantly sporadic occurrence of CSC along with the connection with stress or steroid use hints at a complex disease. In our exome sequencing study on familial CSC in **Chapter 4.2** we could not confirm nor exclude the proposed Mendelian inheritance pattern for CSC. We did not observe one variant or gene that clearly segregated with the disease, rather in each family multiple variants were found to segregate with the disease. This suggests that a complex inheritance pattern might indeed be more likely for CSC. However, it must be noted that included families were small and data of multiple generations were often not available. Therefore, including more and extended pedigrees in the analysis of familial CSC might be beneficial to obtain a more clear picture of variants and genes that might segregate in multiple families, and thus could contribute to familial CSC.

Additionally, at the moment, all genetic studies have been conducted on individuals of Asian and European descent, and studies on other ethnicities are still lacking. It still remains unclear if incidences for CSC differ between ethnicities, and if there might be a genetic component contributing to these differences. Some studies have reported a higher incidence of CSC in Hispanic, Caucasian and Asian populations, compared to African-Americans, but others could not confirm these differences.⁴⁴ Also, it has been suggested that severity of CSC differs between populations, with a more severe phenotype in African-American and Asian populations. Therefore, in addition to replication of current findings in other ethnical groups, to determine population specific genetic associations and disease mechanisms, studies on African and Hispanic CSC patients will be a valuable addition to the field.

Future studies for CSC as a complex disease

Conducting genetic association studies to identify rare variants associated with complex diseases has its challenges. First of all, the sample sizes required for finding single variant associations with rare genetic variants need to be very large due to the very low frequency of the variants involved, and significant costs are associated with the generation of such large datasets. Additionally, once conducted, stringent multiple-testing correction needs to be applied to the results. The latter issue can be partially solved with the use of gene-based analyses in which single variants are clustered into genes, thereby reducing the number of performed tests. However, for the studies included in this thesis, we mainly ran into the first issue, namely the

problem of sample size. Collaborations with the Leiden University Medical Center and the University Hospital of Cologne increased the number of cCSC patients included in the studies, but we did not have sufficient power to identify associations with single rare variants in our exome sequencing study in **Chapter 4.1**. In the future, collaborations with additional research institutes and possibly the initiation of a CSC consortium would be beneficial for all studies into the genetics of CSC.

Besides including more samples in the analysis for both sporadic and familial CSC, studies on the interaction between genetic and environmental factors are necessary. In other complex diseases, such as AMD, it has already been observed that rare variants in families do not completely segregate with the phenotype.¹⁶ This is likely due to the presence of additional genetic variants and environmental triggers in affected individuals, and an absence of these variants and triggers in unaffected family members. Detailed questionnaires on lifestyle events, psychological stress, diet and other habits might be used to obtain information on potential environmental triggers that together with the genetic background of an individual increases risk for cCSC. Combining both genetic and environmental factors might help to generate a prediction model for the occurrence of CSC, which would allow for taking appropriate preventive steps in patients at risk for developing CSC and would aid in designing personalized treatment.

One could even take this a step further by combining steroid or complement measurements in serum or plasma of individuals with CSC with genetic information, or environmental interactions. However, even more interesting would be measuring these factors in the target tissue, the eye. So far, one study has measured subretinal fluid composition during an episode of CSC.⁵⁶ One male patient, with consistent complaints of CSC was treated with eplerenone, however due to a fibrous clot the subretinal fluid did not completely disappear and the patient was operated to remove the clot. During the operation subretinal fluid was collected, subsequent protein and metabolome analysis was performed on this sample and the measurements were compared to fluid from two patients with retinal detachment.

Interestingly, multiple components of the complement system were altered in the CSC-subretinal fluid compared to fluid of the retinal detachments. This included downregulation of complement component 4B levels, whereas FH and FHR1 levels were upregulated, which is consistent with our genetic associations of the corresponding genes. These results give the first hints of which processes are biologically altered in the eyes of CSC patients and might be causative for the accumulation of subretinal fluid. Although these results seem very promising and confirm our genetic associations, we do have to take into account that the patient included in this study was not treatment naïve, and that the administration of eplerenone might have already had impact on proteins and metabolites residing in the subretinal fluid pocket.

Occasions for obtaining serous fluid from patients that at the same time have an episode of CSC and have complications that warrant eye surgery will be very rare. An intermediate solution could be to perform anterior chamber taps on CSC patients, although withdrawal of fluid from the anterior chamber has some associated risks and there are ethical considerations since this procedure is not routinely indicated

for CSC.

Finally, besides replication of current associations and investigating environmental interactions, genetic studies might investigate the relationships between genetic variants and treatment response to MR antagonists, micropulse laser or photodynamic therapy. Also, in light of the described copy number alterations for the *C4B* gene, genome-wide copy number alterations might be explored in CSC.

Naturally, the identified genetic associations need to be translated into underlying biological mechanisms present in the RPE cells or choriocapillaris. Functional studies should be conducted in which the associated genetic variants are introduced into cultured RPE cells and choroidal endothelial cells, or such cells could be generated from induced pluripotent stem cells of CSC patients. The effect of different steroid hormones or complement factors on these cells can then be studied, in light of the CSC phenotype. For these models, possible read-outs might be RPE pump function, transmembrane resistance, choroidal neovascularization and morphology. Models combining multiple cell layers in transwell cultures, or organ-on-a-chip systems have the potential to uncover molecular interactions between RPE cells and the choroidal endothelium that are relevant for cCSC.

In this thesis, we aimed to shed light on the genetic background of CSC. To do this, we performed candidate gene studies, unbiased genome-wide studies and measured hormone levels in patients with cCSC. We have identified new associations with CSC and have proposed studies in this general discussion that might be conducted in the future. Additional research on the disease will help us to better understand the processes behind the occurrence of CSC, and might help to improve treatment for patients affected by this curious disease.

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English summary

Nederlandse Samenvatting

Dankwoord

List of Publications

Curriculum Vitae

PhD Portfolio

Data Management Page

7. Summary/Samenvatting

Dankwoord

List Of Publications

CV



English summary

Central serous chorioretinopathy (CSC) is a chorioretinal disease characterized by fluid accumulation between the photoreceptor cells and the retinal pigment epithelium (RPE). Two different forms of CSC can be distinguished: acute CSC and chronic CSC. Acute CSC usually spontaneously resolves within three months with minimum RPE alterations and visual acuity loss. In chronic CSC (cCSC) the subretinal fluid persists and more wide-spread RPE atrophy and permanent vision loss occurs. The biological mechanisms behind the occurrence of CSC are unknown. The current hypothesis about the etiology suggests that an increased pressure from the underlying vasculature, the choroid, alone or in combination with a dysfunctional pump-function of the RPE leads to the subretinal fluid accumulation.

CSC is more frequent in males and primarily affects men of 45-55 years of age, whom are still in their active working years. Complaints of CSC include metamorphopsia and blurred vision, which has a significant influence on daily life. Besides the male gender, risk factors for CSC include the use of steroids, stress, hypertension, *Helicobacter pylori* infection and pregnancy. Currently, treatment options for CSC are limited, have variable success rates and their biological mechanisms are not clear.

In this thesis we aimed to investigate the genetic components contributing to CSC. We used both candidate gene approaches and unbiased genome-wide approaches and focused mainly on the chronic form of the disease because of its more prominent effect on visual acuity loss. Studying the genetic architecture of CSC might provide us with new insights on the biological mechanisms underlying the disease and could uncover potential targets for new treatment options in the future.

273

Clinical overlap exists between age-related macular degeneration (AMD) and CSC, and a previous study of a Japanese group suggested a genetic overlap as well. Therefore, in **Chapter 2.1** genetic variants that were already known to be associated with AMD were investigated in a large cohort of cCSC patients. The cCSC patients were extensively phenotyped for this study and divided into three phenotypic groups. Group 1 was characterized by bilateral fluid accumulation and RPE alterations, group 2 showed only unilateral abnormalities, whereas group 3 included individuals with retinal abnormalities indicative of other macular diseases. DNA was extracted for 292 cCSC patients, 1311 healthy controls and 1147 AMD patients and was genotyped for 19 single nucleotide polymorphisms (SNPs) in AMD-associated genes and 6 SNPs in *complement factor H (CFH)*. A significant association with cCSC was observed for 3 SNPs (rs800292, rs1329428 and rs1065489) in the *CFH* gene, 1 SNP (rs10490924) at the *Age-Related Macular Susceptibility 2 (ARMS2)* locus and the *CFH*-H3 haplotype. Interestingly, the variants that were risk-conferring for AMD were protective for cCSC, and vice versa. Additionally, a significant difference between phenotype group 1 and group 3 was observed for two of the *CFH* SNPs and the *ARMS2* variant, underscoring the importance of extensive phenotyping.

In **Chapter 2.2** the role of copy numbers of the *complement component 4 (C4)* gene was investigated in cCSC. The *C4* gene, which has two forms (*C4A* and *C4B*), is located on a region of the genome which is prone to recombination, which has led to the occurrence of a variable amount of copies of *C4* in different individuals. *C4* encodes the *C4* molecule which plays an important role in the classical and lectin pathway of

the complement system. We identified a significant association between cCSC and the *C4B* gene in 197 cCSC patients of phenotype group 1 and 303 healthy controls. The absence of *C4B* increased the risk for cCSC, whereas carrying 4 copies of the gene was protective. These findings further support that the complement system might play a role in cCSC.

Chapter 2.3 focuses on genetic variants in the genes for the glucocorticoid (*NR3C1*, GR) and mineralocorticoid (*NR3C2*, MR) receptors. The associations of cCSC with stress, the use of steroids and other hormonal influences such as pregnancy have been extensively described in literature. Also, animal models have suggested a role for the mineralocorticoid aldosterone in the disease mechanisms. We selected 3 functional variants in the GR and 2 in the MR to investigate their potential role in the cCSC disease mechanism in 336 cCSC patients of phenotype group 1 and 1314 unaffected controls. We observed an association with cCSC for the rs2070951 SNP in the *NR3C2* gene, and haplotypes in the same gene. The associated haplotypes have previously been associated to perceived stress, bridging the clinical risk factors for cCSC to an underlying genetic association.

In **Chapter 2.4** we tested the variants in *CFH*, *ARMS2*, *NR3C2* and the copy numbers of *C4B* which were previously associated to cCSC in a cohort of 135 patients with acute CSC (aCSC). We also compared the minor allele frequencies of these variants with 272 cCSC patients and 1385 controls whom were genotyped previously. Three SNPs (rs800292, rs1329428 and rs1061170) in the *CFH* gene and copy number variations of *C4B* were observed to be associated with aCSC. Although odds ratios appeared to be higher in aCSC compared to cCSC, no significant differences were observed for these variants between aCSC and cCSC. This suggests that, despite the difference in clinical presentation, the diseases appear to have a similar genetic predisposition.

In **Chapter 3.1** the first genome-wide association (GWAS) on cCSC was performed using a large cohort of 521 cCSC patients collected in three different university hospitals and 3577 European population controls. Single variant association analysis corrected for sex and ancestry background identified a genome-wide significant association with 20 SNPs located at the *CFH* locus on chromosome 1. Additionally, six suggestive association signals were observed that require replication. Haplotype analysis with 10 variants in the *CFH* gene identified an association with the risk-conferring *CFH*-H2, *CFH*-H4, *CFH*-H5 haplotypes, which all carried the minor allele of the GWAS lead-variant, and the protective *CFH*-H1 and *CFH*-H3 haplotypes containing the major allele of the GWAS lead-variant. Pathway analysis with VEGAS2 and MAGMA identified a significant association with the complement cascade. Finally, the GWAS genotyping data was combined with information of the GTEx database, which contains expression quantitative trait loci (eQTL). These eQTLs allowed us to predict the expression levels of genes in cCSC patients and controls using the PrediXcan program. A significantly different predicted expression in cCSC patients was observed for genes at the *CFH* locus including *CFH*, *CFHR1*, *CFHR4* and *KCNT2*. Additionally, the complement inhibitor gene *CD46* and the *TNFRSF10A* gene were also predicted to be differentially expressed in cCSC patients. This study underlines the importance of the complement system in the molecular disease mechanisms of cCSC and provides further leads for functional and genetic studies on cCSC.

Chapter 3.2 focuses on exome sequencing on a cohort of 269 cCSC patients and 1586 population controls. Data was jointly processed and stringent quality control was performed on the dataset. This retained 263 patients and 1352 controls on which gene-burden analyses were performed using the Burden, SKAT and SKAT-O tests. The gene-based tests were performed including protein-altering variants or all exonic variants combined. Stratification of the data by sex identified a genome-wide significant association in female cCSC patients for the *PIGZ* and *DUOX1* genes when protein-altering variants were used. Analysis including all exonic variants resulted in the associations of two additional genes, *RSAD1* and *LAMB3*, in females specifically. Various suggestive associations were found in the male cohort and in the entire cohort corrected for sex, however none of these associations reached genome-wide significance. The number of carriers of rare variants in the individual *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* genes was significantly higher in female cCSC patients compared to female controls. In total for the 4 genes combined, 45.5% of the female cCSC patients carried at least one rare variant in either of the 4 genes, compared to 18.5% carriers in the female controls. This sex-specific associations observed in this study suggest that the contribution of rare genetic variants to cCSC might be different between the sexes and that the disease mechanisms might differ between males and females.

In **Chapter 4.1** we investigated the phenotype of 103 individuals of 23 different families in which cCSC was present in at least two individuals. Extensive multimodal imaging was performed on cCSC patients and their family members to assess the phenotypic profile of familial cCSC. In 45 individuals (44%) subretinal fluid accumulation or leakage was observed indicative of cCSC. In the family members of these patients, 27 individuals showed findings suggestive of cCSC characterized by a wide spectrum of RPE alterations resembling cCSC but without subretinal fluid accumulation. The remaining individuals were unaffected. Additionally, this study included 24 individuals that had been phenotyped in 1995-1996, and for whom long-term follow-up was investigated in this study. In total, 24% of these individuals showed more severe phenotypes than observed in the previous study, such as subretinal fluid accumulation or RPE alterations not present before. With this study we have shown that CSC clusters in families, and that although many individuals remain asymptomatic they might be at risk for developing lesions predisposing to CSC.

Chapter 4.2 investigates the genetics of familial CSC using most of the families that were included in **Chapter 4.1**. Whole exome sequencing was performed on 72 individuals of 18 different families, including 33 cCSC patients, 18 individuals with suggestive cCSC and 18 unaffected family members. Segregation analysis was performed for each family for rare genetic variants with a frequency <1% in the population, selecting those variants only present in affected individuals and not in unaffected family members. Combining the segregating variants in the families identified 12 different variants present in 2 families. One of these variants in the *PTPRB* gene was also associated with CSC in our case-control cohort (**Chapter 3.2**) and has potential as a cCSC candidate gene because of its biological function. In total, in 28 genes multiples different segregating variants were observed in the families. For these genes RareIBD familial burden analysis was performed and the results were compared with our gene-burden analyses on the case-control cohort. However, no

overlap was observed between the analyses. Although a Mendelian inheritance was suggested for CSC based on pedigree data, in this study we did not observe a clear inheritance with a single rare variant nor with different rare variants within a single gene for cCSC. This underlines the notion that CSC is a complex disease for which both genetic and environmental factors should be taken into account.

In **Chapter 5** we investigated the steroid hormone status of male cCSC patients by simultaneously measuring 17 different steroid hormones in a cohort of 46 patients with active cCSC and 46 age- and sex-matched unaffected controls. Levels of androsterone, androstenedione, estrone and etiocholanolone were elevated in patients compared to controls. Additionally, correlations between the different hormone measurements were assessed and compared between cCSC patients and controls. Correlations between aldosterone and androsterone, estrone, DHEA, DHEAS and 11-deoxycortisol were different compared to controls as well as androsterone with estrone and DHT with 17-OHP. These differences indicate an altered hormone system balance in patients with active cCSC. This study is the first to show associations of cCSC with increased levels of androsterone, androstenedione, estrone and etiocholanolone and the first to suggest that evidence for an altered hormone system balance in these patients by screening multiple components of the steroid hormone system simultaneously.

In the last chapter of this thesis, **Chapter 6**, all studies on the genetics of CSC to date have been summarized and are discussed in the context of literature. Genetic studies on pachychoroid, one of the CSC endophenotypes, are also discussed and the role of the complement system in the disease is further elaborated. Finally, future studies are proposed that might help to unravel the disease mechanisms underlying cCSC.

Nederlandse Samenvatting

Centrale sereuze chorioretinopathie (CSC oftewel serosa) is een chorioretinale aandoening die wordt gekenmerkt door een vochtophoping, ook wel subretinaal vocht genoemd, tussen de het netvlies en het retinaal pigmentepitheel (RPE) in het oog. We onderscheiden twee verschillende vormen van serosa namelijk acute en chronische serosa. Acute serosa verdwijnt gewoonlijk spontaan binnen drie maanden na het ontstaan van de eerste klachten met minimale aantasting van het RPE en weinig verlies van het gezichtsvermogen. Bij chronische serosa blijft de subretinale vloeistof aanwezig, treden er meer en grotere RPE veranderingen op en is er permanent verlies van het gezichtsvermogen. De biologische mechanismen die ten grondslag liggen aan het optreden van serosa zijn nog onbekend. De huidige hypothese over het ontstaan van de aandoening is dat een verhoogde druk van het onderliggende vaatstelsel, het vaatvlies of choroidea genoemd, alleen of in combinatie met een verminderde pompfunctie van het RPE leidt tot de subretinale vochtophoping.

Serosa komt het meest voor bij mannen tussen de 45-55 jaar oud, die nog professioneel actief zijn. De klachten die gepaard gaan met het hebben van serosa zijn onder andere metamorfopsie (vervorming van rechte lijnen) en wazig zicht, wat een aanzienlijke invloed heeft op het dagelijks leven van de patiënt. Naast het mannelijke geslacht zijn er verschillende andere risicofactoren voor serosa beschreven zoals het gebruik van steroïden, het hebben van stress, hypertensie, *Helicobacter pylori* infectie en zwangerschap. Op dit moment zijn de behandelingsopties voor serosa beperkt en hebben ze variabele slagingspercentages. Daarnaast zijn de biologische mechanismen die ten grondslag liggen aan de behandelingen nog niet geheel duidelijk. Om de behandelingsopties voor de patiënt te verbeteren is het belangrijk meer te weten te komen de oorzaken van serosa.

In dit proefschrift hebben we de genetische componenten die bijdragen aan het ontstaan van serosa onderzocht. Hiervoor hebben we kandidaatgen studies en genomwijde studies uitgevoerd en concentreerden we ons voornamelijk op de chronische vorm van serosa, omdat deze vorm van de aandoening een grotere invloed heeft het gezichtsvermogen en de kwaliteit van leven van de patiënt. Het bestuderen van de genetische achtergrond van serosa kan ons nieuwe inzichten geven over welke biologische mechanismen ten grondslag liggen aan de ziekte en welke mechanismen in de toekomst potentiële aangrijpingspunten zouden kunnen zijn voor nieuwe behandelingsmogelijkheden.

Er bestaat klinische overlap tussen leeftijdsgebonden maculadegeneratie (LMD) een andere netvlies aandoening die vooral bij mensen boven de 60 jaar voorkomt, en serosa. Het onderzoek van een Japanse groep suggereerde ook een genetische overlap tussen de twee aandoeningen. Daarom hebben we in **Hoofdstuk 2.1**, in een groot cohort van chronische serosa patiënten, genetische varianten onderzocht waarvan al bekend was dat ze geassocieerd waren met LMD. Voor deze studie hebben we de klinische karakteristieken van de chronische serosa patiënten uitgebreid bekeken en hebben we de patiënten in drie groepen opgedeeld (fenotypering). Groep 1 werd gekenmerkt door vochtophoping en RPE veranderingen in beide ogen, groep 2 vertoonde slechts afwijkingen in één oog, terwijl groep 3 bestond uit personen met retinale afwijkingen die aanwijzingen gaven voor andere aandoeningen van de macula in combinatie met

serosa. Voor deze studie hebben we DNA verzameld van in totaal 292 chronische serosa patiënten, 1311 gezonde controles en 1147 patiënten met LMD. Daarna zijn er 19 nucleotide veranderingen, oftewel single nucleotide polymorfismen (SNPs), getest in deze groepen. De varianten lagen in LMD geassocieerde genen, en daarnaast werden zes varianten in het *complement factor H (CFH)* gen getest in deze individuen. Een significante associatie met chronische serosa werd waargenomen voor drie varianten in het *CFH* gen (rs800292, rs1329428 en rs1065489) en één variant in het *Age-Related Macular Susceptibility 2* (rs10490924, *ARMS2*) locus. Toen we naar de combinatie van de verschillende varianten (haplotype) in het *CFH* gen keken, vonden we ook een associatie met het *CFH*-H3 haplotype. Verassend genoeg zagen we dat de varianten die risicodragend waren voor LMD juist beschermend waren voor chronische serosa en vice versa. Bovendien werd een significant verschil tussen fenotypegroep 1 en groep 3 waargenomen voor twee van de *CFH* varianten en de *ARMS2* variant, hetgeen het belang van uitgebreide fenotypering aangeeft.

In **Hoofdstuk 2.2** werd de rol van het aantal kopieën van het *complement component 4 (C4)* gen onderzocht in chronische serosa. Het *C4* gen, dat twee vormen heeft (*C4A* en *C4B*), ligt op een gebied van het genoom dat gevoelig is voor recombinatie. Dit heeft er toe geleid dat er een variabel aantal kopieën van *C4* in verschillende individuen aanwezig kan zijn. Het *C4* gen codeert voor het *C4* eiwit dat een belangrijke rol speelt in de klassieke en lectine routes van het complementsysteem, een onderdeel van ons afweersysteem. In deze studie vonden we een significante associatie tussen chronische serosa en het *C4B* gen bij 197 chronische serosa-patiënten van fenotypegroep 1 en 303 gezonde controles. De afwezigheid van *C4B* verhoogde het risico op chronische serosa, terwijl het dragen van vier kopieën van het gen beschermend was. Aangezien *C4* een rol speelt in het complement systeem ondersteunt deze studie dat het complementsysteem een rol zou kunnen spelen in chronische serosa.

De associaties van chronische serosa met stress, het gebruik van steroïden en andere hormonale veranderingen zoals zwangerschap zijn uitgebreid beschreven in de literatuur. Daarnaast hebben diermodellen ook een rol gesuggereerd voor het mineralocorticoïde hormoon aldosteron in het ziektemechanisme van serosa. De verschillende hormonen in het lichaam binden aan receptoren om hun functie uit te voeren en veranderingen in deze receptoren kunnen invloed hebben op het effect van het hormoon. Daarom richt **Hoofdstuk 2.3** zich op genetische varianten in de genen voor de glucocorticoïd (*NR3C1*, GR) en mineralocorticoïd (*NR3C2*, MR) receptoren. We selecteerden 3 functionele varianten in de GR en 2 in de MR om hun mogelijke rol in het ziektemechanisme van chronische serosa te onderzoeken. De aanwezigheid van deze varianten werd bepaald in 336 patiënten met chronische serosa van fenotypegroep 1 en 1314 gezonde controles. We zagen dat er een variant in het *NR3C2* (rs2070951) gen en haplotypen in hetzelfde gen vaker voorkwam in de patiënten met chronische serosa. Deze haplotypen waarvoor we een associatie zagen, werden eerder ook gevonden in mensen die een groter gevoel van stress hadden, waardoor er een brug geslagen wordt tussen de klinische risicofactoren voor chronische serosa en de onderliggende genetische associatie.

In **Hoofdstuk 2.4** hebben we de varianten die eerder beschreven zijn voor chronische serosa in een groep van 135 patiënten met acute serosa getest. Deze varianten waren aanwezig in de *CFH*, *ARMS2*, *NR3C2* genen en het aantal kopieën van *C4B* werd

ook onderzocht. De aanwezigheid van de varianten in de acute serosa groep werd vergeleken met 272 chronische serosa-patiënten en 1385 controles die eerder waren getest. Drie genetische varianten in het *CFH* gen (rs800292, rs1329428 en rs1061170) en het aantal kopieën van het *C4B* gen bleken ook geassocieerd te zijn met acute serosa. Hoewel de effecten van deze varianten hoger leken te zijn voor acute serosa, werden er geen significante verschillen waargenomen voor de frequentie van deze varianten tussen acute serosa en chronische serosa. Dit suggereert dat, ondanks het verschil in klinische presentatie, de ziekten een vergelijkbare genetische achtergrond lijken te hebben.

In **Hoofdstuk 3.1** werd de eerste genomwijde associatie analyse (GWAS) voor chronische serosa uitgevoerd met behulp van een groot cohort van 521 chronische serosa patiënten uit drie verschillende universitaire ziekenhuizen en 3577 Europese populatiecontroles. De analyse gecorrigeerd voor geslacht- en genetische etniciteit identificeerde een genomwijde associatie met 20 varianten in het *CFH* locus op chromosoom 1. Daarnaast werden zes suggestieve associaties waargenomen die nog moeten worden gerepliceerd. Haplotype analyse met 10 varianten in het *CFH* gen identificeerde een associatie met de risico-verhogende *CFH*-H2, *CFH*-H4, *CFH*-H5-haplotypen, welke allemaal het secundaire allel van de GWAS-topvariant droegen. De beschermende *CFH*-H1 en *CFH*-H3-haplotypen droegen daarentegen het wildtype allel van de GWAS-topvariant. De VEGAS2 en MAGMA programma's werden gebruikt om de informatie van meerdere genen te combineren om een gemeenschappelijk biologisch mechanisme te identificeren. Deze analyse toonde een significante associatie met de complementcascade aan. Tenslotte werden de GWAS gegevens gecombineerd met informatie van de GTEx-database. Deze database bevat informatie over de invloed van een genetische variant op de expressie van een gen in de vorm van eQTLs. Met deze eQTLs konden we de expressieniveaus van genen in chronische serosa patiënten en controles met behulp van het PrediXcan programma voorspellen, zonder deze te hoeven meten. Het programma voorspelde een verschillende expressie voor de genen in het *CFH* locus (*CFH*, *CFHR1*, *CFHR4* en *KCNT2*) in chronische serosa patiënten. Bovendien werd er ook voorspeld dat het *CD46* gen, dat codeert voor een complement remmer, en het *TNFRSF10A* gen een afwijkende expressie hebben in chronische serosa patiënten. Deze studie onderstreept het belang van het complementsysteem in het moleculaire ziektemechanismen van chronische serosa en biedt verdere aanknopingspunten voor functionele en genetische studies naar de aandoening.

279

Hoofdstuk 3.2 richt zich op het exoom sequenzen van een groep van 269 chronische serosa patiënten en 1586 populatiecontroles. Bij exoom sequencing wordt er gekeken naar alle regio's in het genoom die coderen voor een eiwit, namelijk de exonen van een gen. De genetische informatie van alle individuen werd samen verwerkt en er werd een strenge kwaliteitscontrole op de dataset uitgevoerd. Daardoor bleven er totaal 263 patiënten en 1352 controles over waarop de analyses konden worden uitgevoerd. Voor deze studie hebben we gen-testen gebruikt (Burden, SKAT en SKAT-O) deze testen gebruiken meerdere varianten in het zelfde gen als input en combineren de statistiek van al deze varianten tot één associatie signaal voor elk gen. We hebben de gen-testen eerst uitgevoerd met alleen varianten die het eiwit van een gen direct veranderden en daarna ook met alle varianten die in de exonen

aanwezig waren. Door de dataset op te delen in mannen en vrouwen vonden we een genoom-wijde associatie in de vrouwelijke chronische serosa patiënten voor de *PIGZ* en *DUOX1* genen wanneer we alleen de eiwit-veranderende varianten includeerden in de analyse. De analyse waarbij alle varianten in de exonen werden gebruikt resulteerde in de associaties van twee andere genen, *RSAD1* en *LAMB3*, in de vrouwen. Verschillende suggestieve associaties werden gevonden in het mannelijke cohort en in het volledige cohort gecorrigeerd voor geslacht, maar geen van deze associaties bereikte de genoom-wijde significantie drempel. Het aantal dragers van zeldzame varianten in de *PIGZ*, *DUOX1*, *LAMB3* en *RSAD1* genen was significant hoger in de vrouwen met chronische serosa vergeleken met gezonde vrouwen. In totaal droeg 45,5% van de vrouwen met chronische serosa ten minste één zeldzame variant in één van de 4 genen, vergeleken met 18,5% van de controles. De geslachtsgebonden associaties die we in deze studie hebben gevonden, suggereren dat de bijdrage van zeldzame genetische varianten in chronische serosa mogelijk verschillend is en dat het ziektemechanisme kan verschillen tussen mannen en vrouwen.

In **Hoofdstuk 4.1** onderzochten we de fenotypes van 103 individuen van 23 verschillende families waarin chronische serosa aanwezig was in tenminste twee familieleden. Uitgebreide beeldvorming werd uitgevoerd op de patiënten en hun familieleden om het fenotypische profiel van familiale chronische serosa te bepalen. Bij 45 personen (44%) werd een subretinale vochtophoping of lekkage waargenomen die kan wijzen op chronische serosa. In de familieleden van deze patiënten vertoonden 27 personen bevindingen die wijzen op chronische serosa, gekenmerkt door een breed spectrum van RPE-veranderingen die op chronische serosa lijken maar zonder subretinale vochtophoping. De overige personen vertoonden geen oogafwijkingen die konden duiden op serosa. Daarnaast bevatte deze studie ook 24 individuen die in 1995-1996 al waren onderzocht, en waarvoor we nu lange termijn follow-up gegevens konden verzamelen. In totaal vertoonde 24% van deze personen een ernstiger fenotype dan werd waargenomen in de vorige studie, zoals een subretinale vochtophoping of RPE-verandering die eerder niet aanwezig was. Met deze studie hebben we aangetoond dat serosa voorkomt in families, en dat hoewel veel individuen asymptomatisch blijven, ze mogelijk een risico lopen op het ontwikkelen van laesies die lijken op serosa.

In **Hoofdstuk 4.2** hebben we de genetica van familiale serosa onderzocht in de meeste families die zijn opgenomen in **Hoofdstuk 4.1**. Exoom sequencing werd uitgevoerd op 72 individuen van 18 verschillende families, waaronder 33 chronische serosa patiënten, 18 individuen met suggestieve chronische serosa en 18 gezonde familieleden. Segregatie-analyse werd in elke familie uitgevoerd voor zeldzame genetische varianten met een frequentie <1% in de populatie, waarbij die varianten werden geselecteerd die alleen aanwezig waren in individuen met serosa en niet in gezonde familieleden. Toen we alle segregerende varianten die aanwezig waren in de families combineerden, vonden we 12 verschillende varianten die aanwezig waren in 2 families. Een van deze varianten in het *PTPRB* gen was ook geassocieerd in ons case-control cohort (**Hoofdstuk 3.2**) en is een mogelijk kandidaatgen voor serosa vanwege zijn biologische functie. In totaal werden er in 28 genen verschillende segregerende varianten waargenomen in de families. Voor deze genen werd een familiäre gen-analyse uitgevoerd met behulp van het programma RareID. Deze

resultaten werden vergeleken met onze analyse van de gen-testen in het case-control cohort. Er werd echter geen overlap waargenomen tussen de analyses. Hoewel een dominante overerving werd voorgesteld voor serosa op basis van stamboomgegevens, observeerden we in deze studie geen duidelijke overerving met een enkele zeldzame variant noch met meerdere zeldzame varianten in hetzelfde gen voor chronische serosa. Dit onderstreept het idee dat serosa een complexe ziekte is waarbij rekening moet worden gehouden met zowel genetische factoren als omgevingsfactoren.

In **Hoofdstuk 5** hebben we de steroïde hormoonstatus van chronische serosa patiënten onderzocht door gelijktijdig 17 verschillende steroïde hormonen te meten in een groep van 46 mannelijke patiënten met actieve chronische serosa en 46 gezonde mannelijke controles van dezelfde leeftijd. De niveaus van androsteron, androstenedione, oestron en etiocholanolone in het bloed waren verhoogd bij patiënten in vergelijking met controles. We vergeleken ook de correlaties tussen de verschillende hormoonmetingen in de patiënten en controles. De correlaties tussen aldosteron en androsteron, oestron, DHEA, DHEAS en 11-deoxycortisol waren verschillend in de patiënten met serosa in vergelijking met de controles, evenals de correlaties van androsteron met oestron en DHT met 17-OHP. Deze verschillen duiden op een verandering in de hormoonhuishouding bij patiënten met actieve chronische serosa. Dit is de eerste studie die verhoogde spiegels van androsterone, androstenedione, oestron en etiocholanolone laat zien in patiënten met actieve chronische serosa. Bovendien is het de eerste keer dat een verandering in de balans van het hormoonsysteem is aangetoond bij deze patiënten door meerdere componenten van het steroïde hormoon systeem tegelijkertijd te meten.

281

In het laatste hoofdstuk van dit proefschrift, **Hoofdstuk 6**, zijn alle studies over de genetica van serosa tot op heden samengevat en worden ze besproken in de context van de literatuur. Genetische studies over pachychoroid, een van de serosa endofenotypen, worden ook besproken en de rol van het complementsysteem bij de ziekte wordt verder uitgewerkt. Tenslotte worden toekomstige studies voorgesteld die kunnen helpen om het ziektemechanisme dat ten grondslag ligt aan chronische serosa verder te ontrafelen.

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283

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Wat zullen we vieren? Bata-vieren!! Alle kano-maatjes heel erg bedankt voor alle gezellige trainingen, toernooien, spelletjesavonden, borrels, zaal- en zwembadtrainingen. **Ineke** ons weekendje Lissabon was super! Snel nog maar een keer overdoen, dit keer met de mannen? **Eva, Elke, Hein, Lein, Abel, Maartje** en **Bram** bedankt voor alle gezellige spelletjesavonden!

Fietsend de PhD door, letterlijk en figuurlijk. Bedankt **Jules** voor alle ongein op de weg en het eindeloos kletsen over capibara's en lievelingskleuren. **Fokko** en **Kenneth** ook bedankt voor het wind-tegenhouden, ik ga jullie nog missen in Nieuw Zeeland ;).

Het is bijna onmogelijk om de intriges in de tijden van King Arthur of de tweede wereldoorlog, de avonturen op Skye, de ervaringen met het runnen van restaurants, energiemaatschappijen, boerderijen en onze vaardigheden als koeienhandelaars, wereldveroveraars en treinmachinisten samen te vatten in zo'n klein stukje tekst. Lieve **Rob, Annemarie, Esther, Maarten, Linsey, Joost, Laurent** en **Rosalie** heel erg bedankt voor alle leuke middagen, avonden en weekenden vol met avontuur, veel bier en lekker eten! Langzaamaan is de spelletjesgroep uitgegroeid tot wat hij nu is, en ik hoop dat ondanks dat we zo allemaal onze aparte wegen gaan, we nog steeds zo door kunnen blijven spelen!

Annemarie en **Esther**, samen zijn we ooit begonnen met studeren en binnenkort

breekt echt de tijd aan dat we na 9 jaar niet meer bij elkaar in de buurt wonen. Esther, laten we vooral doorgaan met onze carnavalstraditie, dan wordt het misschien nog wat met mijn Limburgs, roetewisser!! Annie, ik ben heel benieuwd wanneer ik je voor het eerst op tv zie bij de Nationale wetenschapskwis, waar je ook terecht komt, volg je hart!

Dear **Maartje** en **Laura**, dear paranimfas, I am so happy that we did our PhD together and to have you both next to me today. I cannot imagine what it would have been without you! Maartje bedankt voor alle gezellige koffie-wijn-escaperoom-spelletjes-pokémon-kano-dates, laten we daar vooral mee doorgaan! Guapa, thank you so much for everything, all our moments together including Dollars and meeting all your great friends. Maybe one day we should still open our beachbar! I cannot express how much I love you both! <3

Marjo en **Heleen** ineens heb ik er twee moeders bij, bedankt dat ik me vanaf dag 1 welkom bij jullie voel, het voelt als thuiskomen in een warm nest.

Lieve **Abel**, wat is het fijn dat jij er bent. Bij jou kan ik mezelf zijn, lekker nerdy en prettig-gestoord, slechts enkele woorden nodig en je begrijpt me, (toch die kloon-eigenschappen hé) en ik mag nog eens een keer magic van je winnen ook ;). Ik ben benieuwd wat de toekomst ons gaat brengen, laten we het gezellig maken. Suki desu!

Lieve **familie**, bedankt voor de continue steun en interesse in mijn onderzoek de afgelopen jaren. En fijn dat jullie soms genoeg namen met mijn redelijk korte antwoorden als ik er even niet over wou praten. ;)

Janine wat kan ik een zeurpiet zijn als het om opmaak gaat, sorry daarvoor! Bedankt dat je het met me hebt uitgehouden, het resultaat is geweldig! Je kan er heel erg trots op zijn! Super bedankt voor het maken van mijn opmaak en al de moeite die je erin hebt gestoken!

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Lieve **Maarten**, kleine broertjes worden groot en wat is het speciaal dat je vandaag naast me staat. Waar we vroeger vaak ruzie hadden, zijn we nu steeds dichterbij elkaar toe gegroeid. Blijf vooral jezelf, kijk af en toe "met het mes op tafel" zonder geluid, en er ligt een gouden toekomst voor je. Ik ben trots op je!

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List of Publications

RL Schellevis, LAltay, A Kalisingh, V Sitniska, CB. Hoyng, CJF Boon, JMM Groenewoud, AI den Hollander, EK de Jong. "Elevated steroid hormone levels in active chronic central serous chorioretinopathy" Submitted to *JAMA Ophthalmology*

A Castells-Nobau; I Eidhof; M Fenckova; DB Brenman-Suttner; JM Scheffer-de Gooyert; S Christine; **RL Schellevis**; K van der Laan; C Quentin; L van Nijnhuijs; F Hofmann; R Ejsmont; SE Fisher; JM Kramer; SJ Sigrist; AF Simon; A Schenck. "Conserved regulation of neurodevelopmental processes and behavior by FoxP in *Drosophila*". Accepted for publication in *PLOS ONE*

RL Schellevis*, EHC van Dijk*, MB Breukink, JE Keunen, GWE Santen, CB Hoyng, AI den Hollander, CJF Boon, EK de Jong. "Exome sequencing in familial chronic central serous chorioretinopathy" Accepted for publication in *Molecular Genetics & Genomics Medicine*

RL Schellevis, MB Breukink, C Gilissen, CJF Boon, CB Hoyng, AI den Hollander* & EK de Jong*. "Exome sequencing identifies *PIGZ*, *DUOX1*, *LAMB3* and *RSAD1* as susceptibility genes for chronic central serous chorioretinopathy in females" Submitted to *Scientific Reports*

D Mohabati*, **RL Schellevis***, EHC van Dijk, LAltay, S Fauser, CB. Hoyng, EK. de Jong, CJF Boon#, S Yzer# "Genetic risk factors in acute central serous chorioretinopathy" *Retina*, In Press

Schellevis RL, van Dijk EHC, Breukink MB, Altay L, Bakker B, Koeleman BPC, Kiemeny LA, Swinkels DW, Keunen JEE, Fauser S, Hoyng CB, den Hollander AI, Boon CJF, de Jong EK. "Role of the Complement System in Chronic Central Serous Chorioretinopathy: A Genome-Wide Association Study." *JAMA Ophthalmol.* 2018 Aug 2. doi: 10.1001/jamaophthalmol.2018.3190.

Kersten E, Paun CC, **Schellevis RL**, Hoyng CB, Delcourt C, Lengyel I, Peto T, Ueffing M, Klaver CCW, Dammeier S, den Hollander AI, de Jong EK. "Systemic and ocular fluid compounds as potential biomarkers in age-related macular degeneration." *Surv Ophthalmol.* 2018 Jan - Feb;63(1):9-39. doi: 10.1016/j.survophthal.2017.05.003.

van Dijk EHC*, **Schellevis RL***, Breukink MB, Mohabati D, Dijkman G, Keunen JEE, Yzer S, den Hollander AI, Hoyng CB, de Jong EK, Boon CJF. "Familial Central Serous Chorioretinopathy." *Retina.* 2017 Nov 28. doi: 10.1097/IAE.0000000000001966.

van Dijk EHC*, **Schellevis RL***, van Bergen MGJM, Breukink MB, Altay L, Scholz P, Fauser S, Meijer OC, Hoyng CB, den Hollander AI, Boon CJF, de Jong EK. "Association of a Haplotype in the *NR3C2* Gene, Encoding the Mineralocorticoid Receptor, With Chronic Central Serous Chorioretinopathy." *JAMA Ophthalmol.* 2017 May 1;135(5):446-451. doi: 10.1001/jamaophthalmol.2017.0245.

Breukink MB*, **Schellevis RL***, Boon CJ, Fauser S, Hoyng CB, den Hollander AI, de Jong EK. "Genomic Copy Number Variations of the Complement Component C4B Gene Are Associated With Chronic Central Serous Chorioretinopathy." *Invest Ophthalmol Vis Sci.* 2015 Aug;56(9):5608-13. doi: 10.1167/iovs.15-17343.

de Jong EK, Breukink MB, **Schellevis RL**, Bakker B, Mohr JK, Fauser S, Keunen JE, Hoyng CB, den Hollander AI, Boon CJ. "Chronic central serous chorioretinopathy is associated with genetic variants implicated in age-related macular degeneration." *Ophthalmology*. 2015 Mar;122(3):562-70. doi: 10.1016/j.ophtha.2014.09.026.

Curriculum Vitæ

Rosa (Roos) Louise Schellevis was born on the 15th of October, 1991 in Doetinchem, the Netherlands. In 2009, she graduated from the Rietveld Lyceum in Doetinchem with a secondary school diploma focused on Nature and Technical sciences and started her bachelor on Molecular Life Sciences at the Radboud University Nijmegen, The Netherlands. During her Bachelor degree she spent a semester abroad at the Washington & Jefferson College in Washington, Pennsylvania in the United States of America. Her Bachelor was completed in 2012 with an internship at the department of Pathology of the Radboud university medical center (Radboudumc) under supervision of Patricia Groenen. For this internship she studied the integration of the Epstein Barr virus in the genome using fluorescent in situ hybridization.



During her Master on Molecular Life Sciences she followed courses in Medical Biology and Neurobiology. She conducted her first internship at the department of Human Genetics of the Radboudumc under supervision of Annette Schenck and Ana Castells Nobau, studying the role of the FOXP gene in communication using *Drosophila* as a model organism. Her second Master internship was performed at the department of Ophthalmology of the Radboudumc where she worked on the genetics of chronic central serous chorioretinopathy (cCSC) under supervision of Eiko de Jong and Anneke den Hollander.

In 2014, after finishing her Masters with the honor “cum laude”, she continued her work on the genetics of cCSC to obtain her PhD degree. During her PhD she generated and analyzed large datasets of genetic information of cCSC patients and controls in order to unravel the mechanisms of the disease. Her interests lie in using computational science and the latest bio-informatics tools to answer biologically relevant questions regarding disease mechanisms. She was co-organizer of the Dutch Ophthalmology PhD Students (DOPS) congress in 2016, was awarded the Bayer Ophthalmology Research Award (BORA) in 2018 and obtained a travel grant from the Association for Research in Vision and Ophthalmology (ARVO) in the same year. Her four years of doctoral research are presented in this thesis entitled “The Genetics of Central Serous Chorioretinopathy”. She plans to defend the work of her doctoral thesis on the genetics of cCSC in the beginning of 2019.

PhD Portfolio

Rosa L. Schellevis Phd Period: 01-10-2014 to 01-10-2018
 Department of Promotor: Prof. A.I. den Hollander
 Ophthalmology Co-promotor: Dr. Eiko K. de Jong
 Donders Institute for
 Brain, Cognition and
 Behaviour
 Radboud university
 medical center

	Training activities	Year (s)	ECTS
I. Courses & Workshops			
	Graduate school introduction RIMLS	2015	2
	Introduction day Radboudumc	2014	0.5
	Statistics introduction	2014	0.2
	Genetic Epidemiology Course (MED-5E005)	2014	5.7
	Scientific Integrity for PhD Students	2015	0.6
	Complex Trait Analysis of Next Generation Sequencing Data	2015	1.75
	Intermediate & Advanced workshop UCSC	2015	0.4
	eBROK Academy	2016	1.3
	Linux Course	2016	0.8
	BBMRI-omics: Introduction	2017	0.8
	MBSR: Mindfulness Based Stress Reduction for PhD Students	2017	2
	GTEx Resources: eQTLs and Gene Expression workshop	2017	0.1
II. Seminars & Lectures			
	Radboud Research Rounds (9x)	2014-2018	0.9
	RIMLS Technical Forum (4x)	2014-2016	0.4
	RIMLS PhD Retreat* (2x)	2015-2016	1.5
	Sensory Disease Meeting (weekly; 12 presentations [#])	2014-2018	4
	Theme Discussion Human Genetics (weekly, 3 presentations [#])	2014-2018	3
	Research Meeting Oogheelkunde (weekly; 12 presentations [#])	2014-2018	4
	Radboud Science Day: Sensory Disorders*	2015	0.5

Lecture Prof. Evan E. Eicler	2015	0.1
Seminar Jorge Cham	2016	0.1
Donders Sessions (1x)	2016	0.1
Introductory seminar about long-read sequencing and the Sequel	2017	0.1
Transitioning from academia to industry seminar	2017	0.1
III. Symposia & congresses		
RIMLS New Frontiers Symposium (2x)	2014-2015	2
Refereeravonden Oogheekunde (2x)	2014-2015	2
Science Day: Maastricht-Nijmegen	2015	0.25
EUGENDA Retreat & AMD-day ^{##} (4x)	2014-2017	1.5
National PhD Day	2016	0.25
Perception Day	2016	0.5
Donders Discussions*	2016	0.75
ASHG 2017 conference *	2017	1.25
Nederlands Oogheekunde Gezelschap (NOG) 2017 [#]	2017	0.5
Dutch Ophthalmology PhD Students conference (DOPS) ^{##} (n=4)	2015-2018	2.75
Annual ARVO conference ^{*#} (2x)	2016+ 2018	3
IV. Teaching activities and other		
Treasurer DOPS 2016	2016	2
Supervision Master Student Maaïke van Bergen	2014-2015	3
Supervision Master Student Lisa Schiffelers	2016-2017	2
Total		52.7
* Poster presentation; # Oral presentation		

Data Management Page

Type of Data	Subject to privacy (yes/no)	Way of anonymization	Storage
DNA of patients used in this thesis	Yes	A DNA number is assigned to each individual by the cell culture facility of the Department of Human Genetics.	The key is stored in the MCCD database and is only accessible by clinicians and members of the cell culture facility. DNA samples are stored at the Department of Human Genetics. Contact person for the DNA samples is Saskia van der Velde-Visser, saskia.vandervelde-visser@radboudumc.nl
Genotype data of individual SNPs	No	Samples in the dataset are already anonymized	Genotype information is stored in an SPSS file in the following folder on the Human Genetics H-drive: I:\GR Theme groups\06 PI Group Anneke den Hollander\01 Personal folders\Schellevis, Roos\02_KASPAR\01_Statistiek Masterfile\Serosa_genetica_FINAL.sav Results per SNP can be found on the Ophthalmology R-drive: R:\KASPAR\
GWAS genotyping data	No	Samples in the dataset are already anonymized	All files regarding the GWAS study can be found in the following folder on the Ophthalmology R-drive: R:\GWASData
Exome sequencing data	No	Samples in the dataset are already anonymized	All files regarding the exome sequencing study can be found in the following folder on the Ophthalmology R-drive: R:\ExomeData
Clinical characteristics of familial CSC patients and family members	Yes	Samples are coded according to their position in the pedigree and the key is locked with a password	An excel file for conversion can be found on the Ophthalmology H-drive: H:\Onderzoek\Roos Schellevis Additional clinical information can be found here: H:\Onderzoek\Roos Schellevis\Beeldvorming families extra.7z Both files are encrypted with a password which is stewarded by Prof. Anneke den Hollander, Anneke.denhollander@radboudumc.nl

Type of Data	Subject to privacy (yes/no)	Way of anonymization	Storage
Contact details of patients and family members participating in the family studies	Yes	Files are zipped and protected with a password	The zipped files is stored on the Ophthalmology H-drive: H:\Onderzoek\Roos Schellevis\Serosadag\Bellen info.7z The file is encrypted with a password which is stewarded by Prof. Anneke den Hollander, Anneke.denhollander@radboudumc.nl
Contact information of all patients included in the study until 2015	Yes	Files are zipped and protected with a password	The zipped file is stored on the Ophthalmology H-drive: H:\Onderzoek\Roos Schellevis\ Adressen.7z The file is encrypted with a password which is stewarded by Prof. Anneke den Hollander, Anneke.denhollander@radboudumc.nl
Export of Castor database at the end of this thesis containing all CSC for which DNA was collected and whom have been included in the studies in this thesis	Yes	Patients are anonymized and are coded with a database number	The Castor database is set offline and prof. Anneke den Hollander is still able to login at: https://data.castoredc.com/login The final export can be found on the H-drive of Ophthalmology: H:\Onderzoek\Roos Schellevis\SEROSA_DATABASE\ The key for conversion to DNA and Radboud number is stored in the following file: H:\Onderzoek\Roos Schellevis\SEROSA_DATABASE\ CASTOR_sleutel_serosa_database.xlsx The file is encrypted with a password which is stewarded by Prof. Anneke den Hollander, Anneke.denhollander@radboudumc.nl
Informed Consents of all patients included in the studies	Yes	Pdf files of all patients are zipped in a password protected file	The zipped file can be found on the H-drive of Ophthalmology: H:\Onderzoek\Roos Schellevis\IC\ICs.7z The file is encrypted with a password which is stewarded by Prof. Anneke den Hollander, Anneke.denhollander@radboudumc.nl
Clinical files for grading of all patients	Yes	PowerPoint files of all patients are zipped in a password protected file	The zipped file can be found on the H-drive of Ophthalmology: H:\Onderzoek\Roos Schellevis\graden.7z The file is encrypted with a password which is stewarded by Prof. Anneke den Hollander, Anneke.denhollander@radboudumc.nl
Information of steroid use for patients included in the study	Yes	Patient data is stored in an excel file protected with a password	The excel file van be found on the H-drive of Ophthalmology: H:\Onderzoek\Roos Schellevis\ steroid check.xlsx The file is encrypted with a password which is stewarded by Prof. Anneke den Hollander, Anneke.denhollander@radboudumc.nl

Type of Data	Subject to privacy (yes/no)	Way of anonymization	Storage
Serum and plasma samples of patients	Yes	Anonymized with a number and clinical information is stored in a password protected excel file	<p>Stored in the -80C freezers at the Department of Ophthalmology and in the Radboudumc Biobank.</p> <p>Contact person where to find the samples is Bjorn Bakker; bjorn.bakker@radboudumc.nl</p> <p>Clinical information of the samples can be found on the H-drive of Ophthalmology: H:\Trialcentrum\Studies\promovendi\Serosa\Serum samples serosa patienten.xlsx</p> <p>The file is encrypted with a password which is stewarded by Prof. Anneke den Hollander, Anneke.denhollander@radboudumc.nl</p>
All files for the publications presented in this thesis	No	NA	<p>All files can be found on the H-drive of Human Genetics:</p> <p>I:\GR Theme groups\06 PI Group Anneke den Hollander\01 Personal folders\Schellevis, Roos\09_Publications</p>

Donders Graduate School for Cognitive Neuroscience

For a successful research Institute, it is vital to train the next generation of young scientists. To achieve this goal, the Donders Institute for Brain, Cognition and Behaviour established the Donders Graduate School for Cognitive Neuroscience (DGCN), which was officially recognised as a national graduate school in 2009. The Graduate School covers training at both Master's and PhD level and provides an excellent educational context fully aligned with the research programme of the Donders Institute.

The school successfully attracts highly talented national and international students in biology, physics, psycholinguistics, psychology, behavioral science, medicine and related disciplines. Selective admission and assessment centers guarantee the enrolment of the best and most motivated students.

The DGCN tracks the career of PhD graduates carefully. More than 50% of PhD alumni show a continuation in academia with postdoc positions at top institutes worldwide, e.g. Stanford University, University of Oxford, University of Cambridge, UCL London, MPI Leipzig, Hanyang University in South Korea, NTNU Norway, University of Illinois, North Western University, Northeastern University in Boston, ETH Zürich, University of Vienna etc.

Positions outside academia spread among the following sectors:

- specialists in a medical environment, mainly in genetics, geriatrics, psychiatry and neurology,
- specialists in a psychological environment, e.g. as specialist in neuropsychology, psychological diagnostics or therapy,
- higher education as coordinators or lecturers.

A smaller percentage enters business as research consultants, analysts or head of research and development. Fewer graduates stay in a research environment as lab coordinators, technical support or policy advisors. Upcoming possibilities are positions in the IT sector and management position in pharmaceutical industry. In general, the PhDs graduates almost invariably continue with high-quality positions that play an important role in our knowledge economy.

For more information on the DGCN as well as past and upcoming defenses please visit:

<http://www.ru.nl/donders/graduate-school/phd/>

