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GENETIC RISK FACTORS IN SEVERE, NONSEVERE AND ACUTE PHENOTYPES OF CENTRAL SEROUS CHORIORETINOPATHY

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Purpose: To study genetic predispositions and differences between severe chronic central serous chorioretinopathy (cCSC), nonsevere cCSC, and acute central serous chorioretinopathy (aCSC).

Methods: One hundred seventy-three severe cCSC patients, 272 nonsevere cCSC patients, 135 aCSC patients, and 1,385 control individuals were included. Eight single-nucleotide polymorphisms were genotyped in the *ARMS2* (rs10490924), *CFH* (rs800292, rs1061170, rs1065489, rs1329428, rs2284664, rs3753394), and *NR3C2* (rs2070951). Additionally, *C4B* gene copy numbers were analyzed.

Results: A significant association in 5 single-nucleotide polymorphisms in the *CFH* gene could be reproduced among severe cCSC patients, including rs800292 (P = 0.0014; odds ratio [OR] = 1.93; 95% confidence interval [CI] = 1.51–2.47), rs1065489 ($P = 2.22 \times 10^{-4}$; OR = 0.49; 95% CI = 0.34–0.72), rs1329428 (P = 0.001; OR = 1.89; 95% CI = 1.49–2.40), rs2284664 ($P = 1.21 \times 10^{-4}$; OR = 1.65; 95% CI = 1.28–2.13), and rs3753394 ($P = 6.10 \times 10^{-4}$; OR = 0.61; 95% CI = 0.46–0.81). Carrying three *C4B* copies was protective for severe cCSC (P = 0.001; OR = 0.29; 95% CI = 0.14–0.61). No significant differences in allele frequencies could be found among the CSC phenotypes.

Conclusion: Acute CSC, nonsevere cCSC, and severe cCSC all showed a similar association with the *CFH* and *C4B* genes, and the three phenotypes could not be distinguished based on the genetics. This shows that despite the differences in clinical presentation and severity, there is an overlap in the genetic predisposition of different CSC phenotypes. Nongenetic factors may play a more important role in determining the clinical course of CSC.

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Central serous chorioretinopathy (CSC) is a chorioretinal disease, characterized by serous fluid accumulation in the subretinal space, often affecting the macula with subsequent visual impairment.¹ The underlying pathophysiology of CSC is not fully understood. However, a congested, hyperpermeable, and leaking choroid, together with a damaged and dysfunctional retinal pigment epithelium (RPE) are thought to underlie the subretinal fluid (SRF) accumulation in CSC.²

At least two different CSC phenotypes can be distinguished: acute and chronic CSC. Acute CSC (aCSC) is generally considered self-limiting with a near-complete visual recovery, thus not requiring

treatment in most cases. In contrast, chronic CSC (cCSC) often has persistent SRF with more extensive atrophic RPE changes, in which treatment can be beneficial. There is no consensus on the duration threshold that distinguishes acute and chronic CSC, but an arbitrary period of 4 to 6 months of duration of active disease (SRF leakage) is often considered for the definition of chronicity. Apart from chronic SRF leakage, patients with cCSC may present with a wide spectrum of retinal abnormalities. In mild cCSC cases, there are limited areas of RPE atrophy, few RPE detachments, and a circumscribed area of leakage. More severe cCSC cases show widespread or multifocal (or both) areas of RPE atrophy, more numerous RPE

detachments, diffuse areas of leakage, and intraretinal cystoid degeneration.^{4–6} Moreover, this spectrum of severe cCSC was previously shown to have the worst visual prognosis among all cCSC cases, even after treatment and complete resolution of SRF.⁷ Therefore, severe cCSC may be considered a distinct clinical subgroup within the spectrum of CSC.

Recently, specific single-nucleotide polymorphisms (SNPs) in the *age-related maculopathy susceptibility 2* (*ARMS2*), the *complement factor H (CFH)*, and the *nuclear receptor subfamily three Group C member 2* (*NR3C2*) genes were found to be associated with the risk of cCSC.^{8–10} Genomic copy number variations in the *complement component 4 (C4B)* gene were also shown to be associated with cCSC.¹¹ As aCSC, "non-severe" cCSC, and "severe" cCSC appear substantially distinct CSC subgroups with regard to clinical manifestation and prognosis (Figure 1), these different CSC forms may also have different genetic risk profiles.

In the present study, we analyzed the association of SNPs in the *ARMS2*, *CFH*, and *NR3C2* genes, and copy numbers of *C4B* gene, in a cohort of cCSC patients who showed a severe disease presentation based on previously published disease characteristics.⁷ In addition, we analyzed and compared the association

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of the aforementioned risk SNPs between three Caucasian CSC subgroups, including aCSC, cCSC without characteristics of severity, and cCSC patients with severity characteristics.

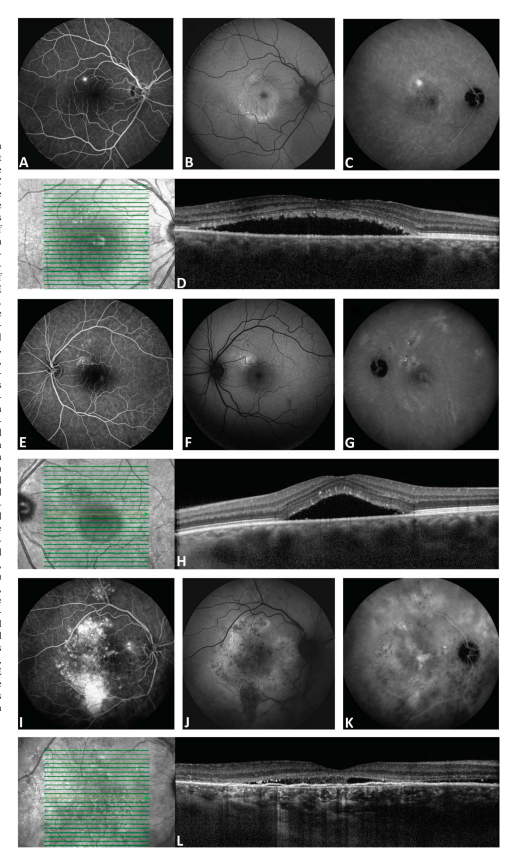
Materials and Methods

In total, 173 white subjects with a severe cCSC phenotype were included, originating from four tertiary referral centers: 65 patients from the Department of Ophthalmology of Leiden University Medical Center (Leiden, the Netherlands), 67 patients from the Radboud University Medical Center (Nijmegen, the Netherlands), 24 patients from the Rotterdam Eye Hospital (Rotterdam, the Netherlands), and 17 patients from the University Eye Hospital of Cologne (Cologne, Germany).

Patients were phenotyped by two experienced retina specialists (S.Y. and C.J.F.B.). For phenotyping, a complete ophthalmological examination was used, including fundoscopy, optical coherence tomography, fluorescein angiography (FA), and, when available, indocyanine green angiography. White patients were included in the severe group of cCSC when they had a history of active disease for more than 6 months, in combination with at least one of the following abnormalities: 1) cumulative areas of larger than five optic disk diameters of diffuse atrophic RPE alterations visible on midphase FA, 2) at least 2 "hot spots" of leakage on midphase FA, 3) an area of diffuse fluorescein leakage larger than one disk diameter on midphase FA, without an evident leaking focus, or 4) presence of posterior cystoid retinal degeneration assessed on optical coherence tomography.^{7,12} Subjects were excluded when there was a suspicion of a (secondary) choroidal neovascularization, aneurysmal choroidal vasculopathy, age-related macular degeneration, multifocal choroiditis, retinal vascular occlusions, or high myopia. The presumably steroid-induced CSC cases (steroid use within 3 months before CSC diagnosis) were not excluded from analysis.

The cohort of severe cCSC was genetically compared with a cohort of 272 white patients with nonsevere cCSC, who had a history of persistent disease but did not have any of the 4 previously mentioned characteristics of severity. Additionally, severe cCSC was compared with 135 white patients with aCSC, defined as a combination of 1) documented serous SRF accumulation on optical coherence tomography, 2) a single focal leakage point on FA, and 3) atrophic RPE alterations limited to less than one disk diameter in size. The control group included white individuals enrolled in the European Genetic

Fig. 1. Clinical features on multimodal imaging in different CSC phenotypes. The right eye of a 34-year-old man with aCSC is shown in A-D. In E-H, the left eye of a 43-year-old male patient with nonsevere cCSC is shown. In I-L, the right eye of a 61-year-old male patient with severe cCSC is shown. Fluorescein angiography imaging revealed a single "hot spot" of leakage and no atrophic RPE changes in the aCSC patient (A). Fluorescein angiography in the nonsevere cCSC showed a leakage spot and multifocal small areas of RPE changes (E), whereas in the severe cCSC case, large and widespread RPE atrophy and diffuse leaking areas were seen (I). On midphase indocyanine green angiography, in the aCSC case, a small hyperfluorescent lesion was observed at the site of the "hot spot" on fluorescein angiography (C). In contrast, indocyanine green angiography in the severe and nonsevere cCSC patients showed more extensive multifocal hyperfluorescent changes (G and K). Fundus autofluorescence imaging showed a mix of granular hyperautofluorescent and hypoautofluorescent changes, which were most prominent in the severe cCSC patient (B, F, and J). Optical coherence tomography scan at first presentation revealed a subretinal serous fluid accumulation and subretinal debris in all patients (D, H, and L). Furthermore, a typical irregular shallow RPE detachment was present in the severe cCSC case (L), which is often observed in combination with chronic SRF leakage.



Database (EUGENDA; www.eugenda.org), in whom no signs of macular disease were found on multimodal imaging, and 176 subjects included in the blood bank of the Radboud University Medical Center. Approval for this study was obtained at the local institutional review boards in all participating centers, and the study adhered to the tenets of the Declaration of Helsinki. Written informed consent was obtained from all subjects before blood collection for genetic analysis.

Single-Nucleotide Polymorphism Genotyping

DNA was isolated from peripheral blood using standard procedures. The most relevant genetic variants to be analyzed were chosen based on findings in earlier genetic studies in CSC and included the following variants: ARMS2 (rs10490924), CFH (rs800292, rs1061170, rs1065489, rs1329428, rs2284664, rs3753394), and NR3C2 (rs2070951), and copy number variations in the C4B gene.8-11 KASP assays (LGC Genomics, Berlin, Germany) were used for SNP genotyping, as described previously and according to manufacturer's instructions. A 7900HT Fast Real-Time PCR system (Applied Biosystems by Life Technologies, Austin, TX) was used to read out the genotyping data. Data analysis was performed with SDS (version 2.4, Applied Biosystems). A TagMan genotyping assay (Hs07226350 cn; Applied Biosystems, Thermo Fisher Scientific, Waltham, MA) with RNaseP as a reference assay was used to measure C4B gene copy numbers, as described previously.

Statistical Analysis

The allele frequency of the SNPs in severe cCSC patients was compared with unaffected controls, non-severe cCSC, or aCSC subjects using a two-sided Pearson's chi-square test (IBM SPSS Statistics, version 22; SPSS, Inc, Chicago, IL). The C4B copy numbers distribution was compared with a two-sided Fisher's exact test. Additionally, a logistic model correcting for gender was designed, and two copies of *C4B* were set as a reference. ¹¹ *P* values of <0.0056 were considered statistically significant after a Bonferroni correction for multiple testing for 9 variants. Hap-

lotype analysis correcting for gender was performed to assess the combined effect of the selected six variants in *CFH* using R (R Core Team, v3.0.2) with the haplo.stats package (v1.7.7). As a reference, the two most frequent haplotypes were used 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 of <5%.

Results

In the present study, we included 173 patients with severe cCSC (mean age: 54 ± 10 years; 151 [87%] males), 272 patients with nonsevere cCSC (mean age: 51 ± 10 years; 216 [79%] males), and 135 patients with aCSC (mean age: 47 ± 10 years; 92 [68%] males). The demographic characteristics are summarized in Table 1.

Association With Single-Nucleotide Polymorphisms in the ARMS2, NR3C2, and CFH Genes

No significant association was found with the rs10490924 variant in ARMS2 gene or with the rs2070951 variant in the NR3C2 gene in the severe cCSC group after correction for multiple testing (Table 2). Also, no difference was observed in allele frequencies of these tested variants when comparing severe cCSC with nonsevere cCSC or aCSC (Table 3). An association could be found in six tested variants in the CFH gene in the severe cCSC group (Table 2). Associations of five CFH variants remained significant after correction for multiple testing: rs800292 (P = 0.0014; OR = 1.93; 95% confidence interval [CI] = 1.51 to 2.47), rs1065489 ($P = 2.22 \times 10^{-4}$; OR = 0.49; 95% CI = 0.34-0.72), rs1329428 (P = 0.001; OR = 1.89; 95% CI = 1.49–2.40), rs2284664 ($P = 1.21 \times 10^{-4}$; OR = 1.65; 95% CI = 1.28–2.13), and rs3753394 (P = 6.10×10^{-4} ; OR = 0.61; 95% CI = 0.46–0.81). No difference was observed when comparing allele frequencies of the six tested variants in the CFH gene between severe cCSC and either nonsevere cCSC or aCSC (Table 3).

Table 1. Demographic Characteristics of the Study Population and Control Subjects per Tested Gene

	Severe cCSC	Non-severe cCSC	aCSC	Controls ARMS2 and CFH	Controls C4B	Controls NR3C2
No. of subjects	173	272	135	826	250	1,385
No. of male subjects	151 (87%)	216 (79%)	92 (68%)	424 (51%)	198 (79%)	635 (46%)
Mean age ± SD (years)	54 ± 10	51 ± 10	47 ± 10	64 ± 12	51 ± 10	51 ± 10

ARMS2, age-related maculopathy susceptibility 2; CFH, complement factor H; C4B, complement component 4; NR3C2, nuclear receptor subfamily 3 Group C member 2.

Table 2.	Analysis	of	Eight	SNPs	in	Severe	cCSC

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SNP (Locus)	Alleles in Controls (Major/Minor)	Severe cCSC (n)	MAF in Severe cCSC	Controls (n)	MAF Among Controls	Unadjusted Allelic <i>P</i>	Allelic OR (95% CI)
rs10490924 (ARMS2)	G/T	171	0.187	812	0.217	0.214	0.83 (0.62–1.11)
rs2070951 (NR3C2)	C/G	172	0.494	1,385	0.468	0.350	1.11 (0.89–1.39)
rs800292 (CFH)	G/A	172	0.372	798	0.235	0.0014*	1.93 (1.51–2.47)
rs1061170 (CFH)	T/C	172	0.282	803	0.353	0.012	0.72 (0.56–0.93)
rs1065489 (CFH)	G/T	172	0.096	794	0.177	$\textbf{2.22}\times\textbf{10}^{-\textbf{4}\star}$	0.49 (0.34–0.72)
rs1329428 (CFH)	C/T	171	0.588	787	0.429	0.0010*	1.89 (1.49–2.40)
rs2284664 (CFH)	C/T	171	0.316	805	0.219	$\textbf{1.21}\times\textbf{10}^{-4\star}$	1.65 (1.28–2.13)
rs3753394 (CFH)	C/T	171	0.202	800	0.293	$\textbf{6.10}\times\textbf{10}^{-\textbf{4*}}$	0.61 (0.46–0.81)

^{*}Two-sided P values < 0.00556 were considered significant after correction for multiple testing.

Association With CFH Haplotypes

Five haplotypes in the *CFH* gene with a frequency higher than 5% and an aggregate of the haplotypes with a frequency lower than 5% were identified. When using the most common haplotype (H1) as a reference and correcting for gender, severe cCSC showed an association with H2, H3, H4, H5, and the low frequency aggregated haplotypes (Table 4). However, only H2 remained significant after correction for mul-

tiple testing, which was risk carrying for severe cCSC (P = 0.001; OR = 1.73; 95% CI = 1.24–2.41; Table 4). Using the H2 haplotype as a reference, H1 and H3 were found to be associated with severe cCSC after correction for multiple testing, carrying a protective effect (P = 0.0013; OR = 0.58; 95% CI = 0.41–0.81 and $P = 4.14 \times 10^{-6}$; OR = 0.30; 95% CI = 0.18–0.50, respectively; Table 4). When comparing the haplotype frequencies of severe cCSC to this frequencies in

Table 3. Comparison of Allele Frequencies in Severe cCSC Versus Nonsevere cCSC and aCSC

	Group	1*	Group 2	†	Grou	p 3‡	Group	2 Versus 1	Group	3 Versus 1
SNPs (Locus)	Severe cCSC (n)	MAF	Non-severe cCSC (n)	MAF	aCSC (n)	MAF	Unadjusted Allelic P	Allelic OR (95% CI)	Unadjusted Allelic P	Allelic OR (95% CI)
rs10490924 (ARMS2)	171	0.187	243	0.193	132	0.174	0.821	0.96 (0.67–1.37)	0.683	1.09 (0.72–1.66)
rs2070951 (NR3C2)	172	0.494	269	0.520	132	0.538	0.447	0.90 (0.69–1.18)	0.216	0.82 (0.59–1.13)
rs800292 ((CFH)	172	0.372	245	0.296	133	0.320	0.021	1.41 (1.05–1.89)	0.177	1.26 (0.90–1.77)
rs1061170 (CFH)	172	0.282	245	0.320	133	0.259	0.235	0.83 (0.62–1.13)	0.534	1.12 (0.78–1.61)
rs1065489 (CFH)	172	0.096	244	0.133	134	0.119	0.101	0.69 (0.44–1.08)	0.350	0.78 (0.47–1.31)
rs1329428 (CFH)	171	0.588	244	0.510	133	0.579	0.0275	1.37 (1.04–1.81)	0.828	1.04 (0.75–1.43)
rs2284664 (CFH)	171	0.316	244	0.275	134	0.287	0.199	1.22 (0.90–1.65)	0.448	1.14 (0.81–1.62)
rs3753394 (CFH)	171	0.202	242	0.273	131	0.263	0.0192	0.67 (0.48–0.94)	0.073	0.71 (0.48–1.0)

Two-sided P values < 0.00556 were considered significant after correction for multiple testing.

ARMS2, age-related maculopathy susceptibility 2; CFH, complement factor H; MAF, minor allele frequency; NR3C2, nuclear receptor subfamily 3 Group C member 2.

ARMS2, age-related maculopathy susceptibility 2; CFH, complement factor H; MAF, minor allele frequency; NR3C2, nuclear receptor subfamily 3 Group C member 2.

^{*}Group 1: severe cCSC.

[†]Group 2: nonsevere cCSC.

[‡]Group 3: aCSC.

Table 4. Complement Factor H Haplotypes in Severe cCSC

			Var	Variants			HF Amond	HF Among	Unadiusted	Allelic OB	Unadiusted	Allelic OB
Haplotypes	Haplotypes rs3753394	rs800292	rs1061170	rs2284664	.664 rs1329428 rs1065489	rs1065489	Controls	Severe cCSC	Allelic, P	(12 % S6)	Allelic, P	(95% CI)
Ξ	O	σ	O	O	O	g	0.329	0.255	Base	Base	$1.25 \times 10^{-3*}$	$1.25 \times 10^{-3*}$ 0.58 (0.41–0.81)
H2	O	∢	_	⊢	-	ഗ	0.209	0.300	0.0013*		Base	Base
¥	-	ڻ ت	-	O	O	-	0.158	0.065	0.012		$4.14 \times 10^{-6*}$	0.30 (0.18-0.50)
H	O	ڻ ت	_	O	-	ഗ	0.133	0.164	0:030	1.57 (1.05–2.35)		0.90 (0.61–1.34)
H5	-	ڻ ت	_	O	-	ഗ	0.072	0.094	0.011	1.91 (1.16–3.15)	0.697	1.10 (0.67–1.82)
Rare	*	*	*	*	*	*	0.098	0.122	0.028	1.65 (1.06–2.57)		0.95 (0.61–1.49)
* O. I.o.	10 +000 13 10 10 10 10 10 10 10 10 10 10 10 10 10	Nobiodo o	tacoitiania bo	+ 002200 2040	201 + 001 + 01 + 01 + 01 + 01 + 01 + 01	Saitoct of						

*P-values < 0.0083 were considered significant after correction for multiple testing. HF, haplotype frequency; MAF, minor allele frequency. nonsevere cCSC and aCSC, no significant differences were found after correction for multiple testing (see **Tables 1 and 2, Supplemental Digital Content 1**, http://links.lww.com/IAE/B132 and http://links.lww.com/IAE/B133, respectively).

C4B Copy Number Determination

The distribution of C4B copy numbers was significantly different in severe cCSC compared with controls after correction for multiple testing (P = 0.0020) (see Figure 1, Supplemental Digital Content 3, http:// links.lww.com/IAE/B134). A logistic regression model showed that carrying three C4B copies was protective for severe cCSC (P = 0.001; OR = 0.29; 95% CI = 0. 14–0.61) (Table 5). The distribution of C4B copy numbers was not significantly different between severe cCSC, nonsevere cCSC, and aCSC groups (see Figure 1, Supplemental Digital Content 3, http://links.lww. com/IAE/B134). In addition, the overall logistic regression model for effect size was not significant when comparing severe cCSC with nonsevere cCSC (P = 0. 665) or when comparing severe cCSC with aCSC (P =0.551) (see Tables 4 and 5, Supplemental Digital Content 4, http://links.lww.com/IAE/B135 and http:// links.lww.com/IAE/B136, respectively).

Discussion

There is a wide variety in the clinical presentation of CSC, ranging from aCSC to severe chronic CSC,^{1,7,12–14} and it is unclear whether these subgroups are different with regard to pathogenesis and genetic background. In the present study, we analyzed specific genetic risk factors in severe cCSC patients and compared them with nonsevere cCSC and aCSC patients. Our data showed that in patients with severe cCSC, three variants (rs800292, rs1329428, and rs2284664) in the CFH gene were significantly associated with an increased risk of the disease, whereas two variants (rs1065489 and rs3753394) were protective. Also, having three copies of the C4B gene was protective against severe cCSC. However, no differences were identified between severe CSC, nonsevere CSC, and aCSC phenotypes.

A comparison of the genetic associations in the three phenotypic subgroups indicated similar risk and protective profiles in the *CFH* gene variants, *CFH* haplotypes, and *C4B* gene copy numbers. Interestingly, although the groups were not significantly different, the genetic effect size, in terms of protective or risk-conferring ORs, was systematically larger in the severe cCSC subgroup compared with nonsevere cCSC and aCSC subgroups. This was also true when

	Overall S	ignificance Model P = 0.007		
	Controls (n = 250)	Severe cCSC (n = 164)	Р	OR (95% CI)
Male sex C4B copy number	198 (79%)	143 (87%)	0.010	0.48 (0.27–0.84)
0	6 (2.4%)	4 (2%)	0.781	0.83 (0.23-3.04)
1	55 (22%)	51 (31%)	0.225	1.33 (0.84–2.12)
2	142 (57%)	99 (60%)	Base	Base
3	44 (18%)	10 (6%)	0.001*	0.29 (0.14-0.61)
4	3 (1.2%)	Ö	0.999	`NA

Table 5. Logistic Regression Model for C4B Load in Severe cCSC Patients

*P-values < 0.0055 were considered significant after correction for multiple testing. C4B, complement component 4; NA, not annotated.

comparing the genetic effect size of *CFH* variants rs800292, rs1329428, and rs1065489 in severe cCSC with cCSC patients in the literature. Severe cCSC may therefore have a stronger genetic predisposition than milder CSC subtypes. Our findings indicate that there is a significant overlap in the known genetic risk factors and therefore likely also pathophysiological overlap between CSC subtypes, despite clinical differences.

A role for the complement system, and the CFH gene in particular, in the pathogenesis of CSC was suggested previously based on genetic association studies.^{8,9,11} Our present study confirms this association in all three CSC phenotypic subtypes. The choroid and choriocapillaris play a central role in the pathogenesis of CSC: although complement activity is abundant in choroidal tissue, 15 complement system dysregulation may be a key factor in CSC disease mechanism. A range of variants in genes involved in the complement system have also been identified in age-related macular degeneration.^{16,17} In contrast to age-related macular degeneration, no systemic complement abnormalities were found in a relatively small group of cCSC patients. 18,19 Local complement system effects may be more important in CSC, rather than systemic complement system abnormalities. However, larger studies on systemic complement differences in cCSC patients are necessary.

Patients with CSC share certain clinical characteristics with age-related macular degeneration, such as macular fluid leakage and RPE abnormalities, as well as possible complication of choroidal neovascularization, 20 but there are also clear differences such as an earlier age at onset, an absence of drusen, the presence of pachychoroid, and association with steroid use. The *CFH* variants reported in this study seem to have opposite effects in CSC compared with age-related macular degeneration, which may point to a different role of the complement system in the pathophysiology of these diseases as suggested before. 8 In our current

cohorts, we could not replicate the associations with the *ARMS2* gene and *NR3C2* gene variants as demonstrated previously.^{8,10} This lack of a significant association may be explained by the smaller sample size of the subgroups.

In the present study, a possible role of other, currently unknown, genetic variants cannot be excluded. Other factors may have a more prominent role than genetic factors in determining the course and severity of the disease. Daruich et al²¹ suggested that older age (>40 years), presence of high (>50 μ m) RPE detachments, and a thickened (>500 μm) choroid are significantly correlated with a prolonged episode of aCSC. Long-term steroid use has been suggested not only to increase the risk of CSC but also to cause a more severe bilateral chronic disease with multiple RPE leaking sites, more extended areas of RPE atrophy, and even bullous retinal detachments.^{22–24} Piccolino et al¹² have shown that presence of posterior cystoid retinal degeneration, which was considered a sign of severity in our study, is specifically associated with steroid use, longer duration of symptoms, and subretinal fibrin accumulation. Furthermore, severe cCSC presentations were previously described in pregnant women²⁵ and among certain ethnic groups.²⁶ Our findings suggest that the profile of known genetic risk SNPs between phenotypically different CSC patients is similar, and therefore, it is likely that other factors, such as described above, determine disease course and outcome.

In conclusion, associations between *CFH* genetic variants and *C4B* copy numbers and severe CSC were demonstrated, but no marked genetic differences were found between acute, nonsevere, and severe chronic phenotypes of CSC in the tested variants. This study indicates that different phenotypes of CSC may not develop as a result of genetic predisposition, at least among the currently known CSC-associated *CFH* variants. Presumably, other nongenetic risk factors such as environmental factors or currently unknown genetic

variants may play a role in the clinical course of CSC. Future genetic and clinical studies in larger cohorts may provide important clues about the different risk factors associated with CSC disease severity.

Key words: acute central serous chorioretinopathy, *ARMS2*, chronic CSC, *complement factor H, complement component 4*, genetic association, *NR3C2*, severe CSC.

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