Genome-wide association analyses identify multiple loci associated with central corneal thickness and keratoconus


Central corneal thickness (CCT) is associated with eye conditions including keratoconus and glaucoma. We performed a meta-analysis on >20,000 individuals in European and Asian populations that identified 16 new loci associated with CCT at genome-wide significance (P < 5 × 10−8). We further showed that 2 CCT-associated loci, FOXO1 and FNDC3B, conferred relatively large risks for keratoconus in 2 cohorts with 874 cases and 6,085 controls (rs2721051 near FOXO1 had odds ratio (OR) = 1.62, 95% confidence interval (CI) = 1.4–1.88, P = 2.7 × 10−10, and rs4894535 in FNDC3B had OR = 1.47, 95% CI = 1.29–1.68, P = 4.9 × 10−7). FNDC3B was also associated with primary open-angle glaucoma (P = 5.6 × 10−4; tested in 3 cohorts with 2,979 cases and 7,399 controls). Further analyses implicate the collagen and extracellular matrix pathways in the regulation of CCT.

Human ocular biometric parameters comprise a set of highly heritable and often correlated quantitative traits. One notable example is CCT, which has an estimated heritability of up to 95% (ref. 1). Whereas extreme corneal thinning is a dramatic clinical feature for rare congenital connective tissue disorders, including brittle cornea syndrome (BCS) and several types of osteogenesis imperfecta2–3, mildly reduced CCT is involved in more common and late-onset eye diseases. It is a hallmark of keratoconus and a risk factor for primary open-angle glaucoma (POAG) in individuals with ocular hypertension4–5. Previous genome-wide association studies (GWAS) conducted on both European and Asian populations have identified 11 CCT-associated loci6–9. Among these loci, mutations in ZNF469 (refs. 10–12), COL5A1 (ref. 13) and COL5A2 (refs. 14,15) are known to cause rare disorders of BCS, Ehlers-Danlos syndrome (EDS) and corneal dystrophy, respectively. However, none was found to be associated with common eye diseases.

Keratoconus is a common corneal ectasia, affecting 1 in 2,000 in the general population16. It is a progressive eye disease characterized by thinning and asymmetrical conical protrusion of the cornea, which causes variable and severe visual impairment. Owing to the limited availability of medical treatments, keratoconus is one of the leading causes of corneal transplantation worldwide17. Two GWAS have been conducted on susceptibility for keratoconus, and these studies suggested some new genetic associations, but neither study reported genome-wide significant loci18,19. POAG is the most common form of glaucoma, which is the second leading cause of blindness worldwide20. Several risk loci for POAG have been identified through early linkage and candidate gene studies21,22 and recent

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GWAS\textsuperscript{23–27}. In both diseases, affected individuals have reduced CCT relative to the general population.

Dissecting the genetics of key risk factors may provide insights into associated disease etiology. Therefore, we conducted a large meta-analysis of GWAS on CCT from over 20,000 individuals, including individuals of European and Asian descent who were affected or unaffected with glaucoma to identify new CCT-associated loci. To evaluate potential clinical relevance, we tested the identified CCT-associated loci in 2 keratoconus susceptibility studies (totaling 874 keratoconus cases and 6,085 controls) and 3 studies on POAG risk (totaling 2,979 POAG cases and 7,399 controls). The overall study design is shown in Supplementary Figure 1.

RESULTS

Meta-analysis of CCT from >20,000 samples

We collected 13 GWAS on CCT, totaling over 20,000 individuals (Supplementary Table 1). Because of differences in the sample attributes, we performed meta-analyses of CCT within each subset, including 13,057 individuals with European ancestry who were unaffected with eye disease in set 1, 6,963 individuals with Asian ancestry who were unaffected with eye disease in set 2, 1,936 POAG cases with European ancestry in set 3 and 198 normal tension glaucoma (NTG), a common form of open-angle glaucoma) cases with Asian ancestry in set 4. All samples were genotyped on commercially available genotyping arrays. The majority of studies in the first two sets were imputed according to the phased haplotypes of HapMap reference samples, whereas few studies in the other two sets had imputation data available by the time of this study.

We separated the twin studies (Australian BATS and TEST twin studies and UK twin studies) from the first set of samples with European ancestry. Because family studies are less affected by potential population stratification, these twin studies were analyzed as a direct replication of association results from the discovery set consisting of all the remaining set 1 samples. All loci from the discovery set were found in the twin studies with similar effect sizes and with at least nominal replication (Table 1). This result, together with a low genomic control parameter ($\lambda = 1.05$; Supplementary Fig. 2), suggested that population stratification had little effect on the meta-analysis. We then performed meta-analysis on the results from the discovery set and the twin studies, reported as combined set 1. This combined set identified 13 loci that were associated with CCT at genome-wide significance in the populations of European descent (Table 1, Supplementary Fig. 3a–m and Supplementary Table 2).

Those included five known loci, COL5A1, AVGR8 (reported here as FGFR9-SGCC), FOXO1, AKAP13 and ZNF469, in European populations\textsuperscript{28,29}, one locus, LRRK1-CHSY1, previously identified in Asian populations\textsuperscript{6} but unknown in European populations and seven new loci (Fig. 1 and Table 1).

Using a recent approach of approximate conditional and joint multiple-SNP analysis\textsuperscript{30}, we found that two loci harbored multiple independent variants that were associated with CCT (Table 1). One example was the COL5A1 region. Previous studies based on European and Asian samples both suggested the presence of two independent signals in this region\textsuperscript{6,9}. In the current meta-analysis, we observed that the two variants rs3118520 and rs7044529, which were 126 kb apart with linkage disequilibrium (LD) $r^2 < 0.01$ in our reference samples from the Queensland Institute of Medical Research (QIMR) cohort\textsuperscript{28,29}, represented two independent LD blocks. The first LD block was located upstream of COL5A1 (RXRA-COL5A1), and the second was in an intron of COL5A1. Similarly, the locus at LRRK1-CHSY1 harbored three independent associations that could be captured by rs2034809, rs930847 and rs752092. Because the trait-increasing alleles of rs2034809 and rs930847 were negatively correlated ($r = -0.17$), the association at rs2034809 was not found to reach genome-wide significance in the single-SNP analysis ($P = 7.5 \times 10^{-5}$), and the marginal effects of both SNPs were underestimated (Table 1).

The five known loci (including two independent signals at COL5A1) explained 4% of additive variance in CCT in European populations. The eight loci not known in European populations (including multiple signals at the LRRK1-CHSY1 locus) explained an additional 3.5% of additive variance, adding up to 7.5% of the total variance explained in European populations.

The two published GWAS of CCT in three Asian ancestry groups—Indian, Malay and Chinese—successfully replicated associations at COL5A1, AKAP13 and ZNF469 (refs. 6,9). To investigate whether CCT-associated loci are important across populations on a larger scale, we tested the loci identified from European samples in set 1 in these three Asian populations. All loci had effect directions consistent with the ones found in European samples except for rs10189064 at the USP37 locus, which was very rare in the Asian samples. Furthermore, 11 of 15 lead SNPs (or their proxies) were at least nominally significant at $P < 0.05$ (Supplementary Table 3). These observations suggest that most of the CCT-associated loci identified from populations of European descent are shared in Asian populations. Therefore, we performed another meta-analysis using Fisher’s method to combine association $P$ values from GWAS of European samples and Asian samples. Fisher’s method makes no assumptions regarding trait distribution or allele frequencies across studies. This meta-analysis showed that the known CCT-associated loci in the Asian populations, including COL5A2, FAM46A-IBTK, C7orf42 and 9p23 (reported here as MPDZ-NF1B), replicated in European samples and further identified ten new loci associated with CCT, including COL4A3, which had been suggested in a set of Croatian samples but had not reached genome-wide significance (Fig. 2a and Table 2).

Many studies observed that individuals with glaucoma, especially those with the advanced form, have thinner corneas than the general population\textsuperscript{31}. Owing to phenotypic differences and potential confounding factors, for example, the fact that individuals with glaucoma take intraocular pressure-lowering medication, which has been shown to be associated with corneal thinning\textsuperscript{31}, we conducted separate meta-analyses of CCT in the clinical cohorts. We collected 5 cohorts comprising a total of 1,936 POAG cases with European ancestry (set 3) and an additional cohort from Hong Kong comprising 198 NTG cases with Han Chinese ancestry (set 4), all of whom had CCT measured. Neither of these two sample sets had adequate statistical power to identify genetic loci significantly associated with CCT (in the lower range observed in individuals with glaucoma). We tested the CCT-associated loci identified from the general population in these two sets. The results from meta-analysis in set 3 showed that all genome-wide significant loci except one had the same direction of effect in open-angle glaucoma (OAG) samples compared to those in the general population, and 12 loci were at least nominally significant (Supplementary Table 4). In set 4, most of the loci were found to have effect directions consistent with those from the other sets, but, owing to the very limited power in this study, only one locus showed nominal association (Supplementary Table 4). We applied polygenic modeling in a subset of POAG cases from set 3 and found that polygenic overlap existed beyond the loci detected in healthy individuals and in POAG cases (Supplementary Fig. 4a). Despite the slight phenotypic differences, we have shown that there is substantial overlap between the genetic variants that account for CCT variation in the general population and the variants found in...
individuals with glaucoma. This suggests that similar pathway(s) regulate CCT regardless of eye disease status.

Pathway analysis of CCT-associated loci
To investigate whether the CCT-associated loci were enriched in functional units in terms of genes and pathways, we performed a gene-based analysis using the Versatile Gene–based Association Study (VEGAS) and a further pathway analysis using a new tool extended from VEGAS (VEGAS-Pathway). These analyses were first run on the meta-analysis results from set 1; therefore, HapMap phase 2 Utah residents of Northern and Western European ancestry (CEU) samples were used as the reference to estimate patterns of LD. We defined the gene regulatory region as 50 kb both upstream and downstream of a gene. This definition includes most but not all regulatory effects. For the lead SNPs at the ZNF469 locus, we identified several independent associations, as suggested from the results of the conditional and joint analysis. The top-ranking pathways were collagen (GO 0005581, empirical $P = 1.1 \times 10^{-8}$), followed by ECM and fibrillogenesis (GO 0003179, empirical $P = 4.8 \times 10^{-10}$). The significance of the ECM pathway, as expected, but the pathway analysis reduced the number of ECM-containing genes and repeating the pathway analysis reduced the results of the conditional and joint analysis. The leading pathways defined by the Gene Ontology (GO) database. We found that the top-ranking pathways were collagen (GO 0005581, empirical $P = 1 \times 10^{-8}$), followed by extracellular matrix (ECM; GO 0031012), ECM part, fibrillar collagen and collagen fibril organization, as well as multicellular organismal metabolic process and myosin binding (Supplementary Table 6). A number of collagen genes were common in these pathways, for example, COL2A1, COL5A1, COL5A2, COL5A3, COL11A1 and COL11A2. Removing all of the genome-wide significant genes and repeating the pathway analysis reduced the significance of the collagen pathway, as expected, but the pathway remaining was significant (empirical $P = 0.005$), suggesting that more of the remaining collagen pathway genes also contribute to variation in CCT. The three pathways involving ECM contain various components, for example, connective tissue components, such as HAPLN1, and molecules regulating cell migration and adhesion, for example, EDIL3 and TGFBR2. We ran another gene set enrichment analysis using the MAGENTA tool (Supplementary Table 5).
among the significant pathways (empirical $P$ values = 5.2 $\times$ 10$^{-5}$ and 0.01, respectively) (Supplementary Table 7).

The same framework of VEGAS-Pathway analysis was applied to the meta-analysis of the three Asian populations, with the HapMap 2 Japanese in Tokyo, Japan (JPT) and Han Chinese in Beijing, China (CHB) populations set as the reference for LD patterns. None of the GO pathways was significant after Bonferroni correction for multiple testing. But the top-ranking pathways overlapped well with the ones from the European populations, which included collagen fibril organization, ECM organization and ECM part (Supplementary Table 8). Notably, two pathways that were among the top—face morphogenesis and head morphogenesis—although not significant after correction for multiple testing, contained the gene PDGFRα, which has been reported to associate with human corneal curvature in Asian populations$^{34}$. In a meta-analysis of pathways obtained from both European and Asian samples using Fisher’s method, four pathways were significant after correction for multiple testing: collagen fibril organization (Fisher’s $P = 3.2 \times 10^{-7}$, 1.5 $\times$ 10$^{-3}$ after correction), collagen (Fisher’s $P = 3.4 \times 10^{-6}$, 0.016 after correction), fibrillar collagen (Fisher’s $P = 4.8 \times 10^{-6}$, 0.022 after correction) and ECM part (Fisher’s $P = 5.2 \times 10^{-6}$, 0.024 after correction), and ECM was borderline significant (Fisher’s $P = 1.3 \times 10^{-5}$, 0.06 after correction) (Supplementary Table 9).

Finally, we used the GRAIL tool$^{35}$ to explore the functional connectivity between the CCT-associated loci using the GO database as the knowledge base. The top two pathways suggested from this gene set enrichment approach were ECM structural constituents conferring tensile strength (GO 0030020) and the collagen pathway (Supplementary Table 10). The results were thus consistent using different pathway approaches. Using PubMed Text as a knowledge base, the top key words showing functional connection between the 27 loci included collagen, syndrome, cornea, mutation and mitochondrial. We also considered each locus singly, but none was significantly connected with the remaining loci (GRAIL $P_{ext}$ > 0.05 for all genes).

![Manhattan plot](image_url)  
**Figure 1** Manhattan plot of set 1 meta-analysis results. Association results (–log$_{10}$P values) are plotted for each chromosome. A blue dot and the corresponding gene name indicate that the locus is known in European populations. The red dot and the corresponding gene name indicate that the locus has not previously been reported as associated with CCT in European populations.

![Association plot](image_url)  
**Figure 2** Association with CCT in European and Asian populations, and with keratoconus risk in European populations. (a) Association with CCT in European populations (set 1). (b) Association with CCT in Asian populations (set 2). (c) Association with keratoconus risk in European populations. This plot is a standard forest plot, but, instead of showing study-specific effects, it shows the association effects in the three panels for each locus. The allele is chosen as the CCT-reducing allele (boxes that represent the point estimates in a and b are on the left side of the dashed line, effect on CCT < 0). Effect on CCT is in standardized units. Given the association between reduced CCT and elevated keratoconus risk, for the effect directions to be consistent, the CCT-reducing allele would be the keratoconus risk allele (boxes that represent the OR of keratoconus in c on the right side of the dashed line, OR > 1). Five loci in blue, FOXO1, two SNPs at COL5A1, FND3C3B and MPDZ-NF1B, are the CCT-associated loci that are significantly associated with keratoconus risk after correction for multiple testing and also show consistent effect directions. The ZNF469 locus is marked in red because the CCT-reducing allele is significantly associated with a lower risk of keratoconus. rs1564892 and rs7620503 were not genotyped in the keratoconus case-control studies and are therefore presented as missing values (NA).
These results together highlight the importance of the collagen and ECM pathways in the regulation of CCT.

Clinical relevance to rare and common eye diseases
Corneal thinning is reported as one of the clinical features in a number of rare disorders that affect a variety of connective tissues. Rare mutations in ZNF469 (refs. 10–12) and a range of collagen genes2,3,13–15 have been mapped to these disorders. We and others have identified associations of the common variants in these genes, for example, ZNF469, COL5A1 and COL8A2, with normal corneal thickness7–9. In a more recent study, mutations in PRDM5 (4q27) were identified in two affected families with BCS who were known to be negative for mutations in the ZNF469 gene46. Although none of the CCT-associated loci were mapped to PRDM5, the common SNP rs10518367 70 kb upstream of the gene was associated with CCT in the European samples at the significance level of \( P = 8.9 \times 10^{-5} \).

Keratoconus and POAG are common conditions with milder corneal thinning but more complex genetic etiologies. To investigate whether the CCT-associated loci influence genetic susceptibility to these conditions, we tested association of these loci in case-control studies with a prespecified \( P \)-value threshold for testing 26 independent SNPs (Table 2) in 2 diseases of 0.001 (\( P < 0.05/52 \)). As reduced CCT is associated with POAG5 and progressive corneal thinning is observed in keratoconus17, we hypothesized that, for the effect directions to be consistent in the epidemiological sense, the CCT-reducing allele would also be the keratoconus or POAG risk allele.

We genotyped the CCT-associated loci from Table 2 in 652 keratoconus cases and 2,761 controls and also evaluated these loci in a published GWAS of keratoconus susceptibility with 222 cases and 3,324 controls19 (Supplementary Table 11). The meta-analysis of these 2 sets (874 keratoconus cases and 6,085 controls in total) identified 6 loci that were strongly associated with keratoconus risk: FOXO1 (OR = 1.62, 95% CI = 1.4–1.88, \( P = 2.7 \times 10^{-10} \) for rs2721051), FNDC3B (OR = 1.47, 95% CI = 1.29–1.68, \( P = 4.9 \times 10^{-9} \) for rs4894535), RXRA-COL5A1 (OR = 1.32, 95% CI = 1.19–1.47, \( P = 2.6 \times 10^{-7} \) for rs1536482), MPDZ-NF1B (OR = 1.33, 95% CI = 1.18–1.51, \( P = 5.2 \times 10^{-6} \) for rs1324183), COL5A1 (OR = 1.37, 95% CI = 1.19–1.57, \( P = 8 \times 10^{-6} \) for rs7620503) and ZNF469 (OR = 1.25, 95% CI = 1.11–1.40, \( P = 1.9 \times 10^{-4} \) for rs9938149) (Fig. 2c and Table 3). However, among the six significant loci, the rs9938149 SNP in ZNF469 showed an unexpected effect direction, with the CCT-increasing allele leading to increased risk for keratoconus (Fig. 2c). Applying polygenic modeling to the 222 keratoconus cases and 3,324 controls, we found that the aggregate effects of loci with more modest effect on CCT did not clearly enhance the prediction of keratoconus risk over and above the risk conferred by the six significantly associated loci (Supplementary Fig. 4b). We note, however, that this lack of increased prediction might have been caused by the polygenic modeling only being applicable to a subset of our available keratoconus cases (only 222 of the total 874 cases had GWAS genotyping data available, meaning that most keratoconus samples did not contribute to the polygenic modeling result).
Similarly, we examined the CCT-associated loci for a role in POAG risk in 3 clinical cohorts (2,979 cases and 7,399 controls in total): advanced POAG cases and controls from the Australian and New Zealand Registry of Advanced Glaucoma (ANZRAG)\textsuperscript{24,37} and POAG cases and controls from GLAUGEN and NEIGHBOR\textsuperscript{25,38,39} (Supplementary Table 12). Performing meta-analysis of these results, we found that \textit{FNDC3B} was also significantly associated with POAG risk ($P = 5.6 \times 10^{-4}$) (Table 3). However, unlike its association with keratoconus, the CCT-reducing allele of rs4894535 was associated with reduced risk of POAG ($OR = 0.83$, $95\% CI = 0.74–0.92$). Furthermore, polygenic modeling on these three sets showed limited evidence for CCT aggregate effects predicting POAG risk (Supplementary Fig. 4c).

Taken together, variants identified through GWAS of CCT were not only relevant to rare disorders but to more common and complex eye diseases. Other CCT-associated loci might also be valuable candidate genes when studying the genetics of keratoconus.

### Gene expression in human and inbred mouse cornea

To further establish genes from the identified loci as candidates for influencing CCT, we examined publicly available microarray gene expression data from the corneas of two human donor samples and three strains of inbred mice\textsuperscript{40,41} (Supplementary Table 13). For lead SNPs with a single associated nearby gene, the vast majority were expressed in the corneas of both humans and mice. For lead SNPs with two nearby genes, presence-absence of corneal expression suggested prioritization among the two candidates in two loci (expressed in \textit{CWC27}, not expressed in \textit{ADAMTS6}; expressed in \textit{PTGDS}, not expressed in \textit{LCN12}). To our knowledge, except for collagen loci, none of the lead genes are known to influence corneal thickness in mice, although some strains with mutations influencing collagen or ECM have previously been described\textsuperscript{42-45}.

We also queried the publicly available expression quantitative trait locus (eQTL) database GHS Express\textsuperscript{46} to determine potential SNP function in terms of transcript regulation. We found that rs930847 and rs752092 were cis eQTLs for \textit{LRRK1} ($P = 1.04 \times 10^{-11}$, $R^2 = 3.4\%$) and \textit{CHSY1} ($P = 1.2 \times 10^{-6}$, $R^2 = 1.8\%$), respectively, both in monocytes. There is limited evidence of functional roles for the other CCT-associated loci identified in the meta-analysis.

### DISCUSSION

Previous GWAS of CCT have identified 11 loci, 5 of which were found in studies of individuals with European ancestry\textsuperscript{2-6} and 6 of which came from studies of 3 Asian populations\textsuperscript{9-11}. We performed CCT meta-analysis on an overall sample size of >20,000 individuals and brought the number of CCT-associated loci to 27. Using a recent approach of approximate conditional and joint analysis\textsuperscript{19}, we found that two loci harbored multiple independent variants in association with CCT. In the subset of normal samples with European ancestry, we estimated that the five known loci explained 4% of additive variance in CCT and that eight new loci in European populations explained another 3.5%. The loci associated with CCT in European and Asian populations (Table 2) together explained 8.3% of additive variance in Europeans and 7% in Asians.

The differences in the sample attributes enabled various comparisons between the subsets of meta-analysis results. We observed
similar CCT distributions in European and Asian populations and found that the underlying genetic effects were largely shared between the two ancestry groups. Despite a slightly lower CCT range in the group of individuals with glaucoma, there was significant overlap in CCT loci from samples affected and unaffected with glaucoma, suggesting similar pathways regulating CCT regardless of disease status. Our results also showed that genes harboring rare variants causing Mendelian disorders with clinical feature of extreme corneal thinning (for example, BCS and osteogenesis imperfecta) also harbor common variants that influence CCT in the general population.

We evaluated the relevance of CCT-associated loci for two major eye diseases, keratoconus and glaucoma, using independent case-control sets. We found that, despite the modest effect on CCT, 11 SNPs showed nominal association with keratoconus, with 6 significant after correction for multiple testing. The effect of the CCT-associated loci on glaucoma was less pronounced, although the FNDC3B locus was significantly associated with POAG. Our results showed that part of the genetic predisposition to these diseases was mediated through the genes underlying CCT, with the remaining predisposition attributable to alternative mechanisms.

We expected CCT-reducing alleles to be associated with increased risk of keratoconus and POAG, given the known association between thin corneas and these two diseases. In general, this was true for keratoconus, with most CCT-reducing alleles associated with increased risk of keratoconus. However, there were exceptions. The CCT-reducing allele near ZNF469 (rs9938149) led to decreased keratoconus risk. At FNDC3B, the CCT-reducing allele resulted in elevated keratoconus risk (OR = 1.47, 95% CI = 1.29–1.68), but it lowered POAG risk (OR = 0.83, 95% CI = 0.74–0.92). The pattern of complex associations seen at ZNF469 for keratoconus and at FNDC3B for POAG, where the effect directions did not always agree with the epidemiological prediction, might either reflect a false positive association or a genuine pleiotropic action of these genes in disease. An example was shown in which a body fat percentage-decreasing allele near IRS1 also associated with adverse metabolic profile, and the authors showed that it could be elucidated by a complex mechanism of pleiotropic actions.

The six SNPs that were significantly associated with keratoconus each had a relatively large OR (range of 1.25–1.62). Unlike the inflated ORs obtained from testing thousands of variants for association between a trait and marker in GWAS, our targeted typing of SNPs on the basis of the CCT findings, led to unbiased estimates for the ORs. Because all six SNPs had moderately high risk allele frequencies, a risk profile based on even just these SNPs yielded reasonable risk prediction. For example, ~1% of the population carry 1 or 2 risk-associated loci at each of the 6 loci, and these individuals are at 7.2-fold increased risk (assuming a multiplicative model) of keratoconus relative to the ~1% of the population who are homozygous for the protective allele at each SNP. Although these six loci combined only explained ~2% of the variation in CCT, the effect of these SNPs on keratoconus risk is large, and further evaluation of the clinical relevance of these SNPs is merited.

Our data clearly show that the endophenotype approach yields disease-relevant loci. Notably, the new CCT-associated locus from our meta-analysis, FNDC3B, was also significantly associated with both keratoconus and POAG risk. This locus did not reach genome-wide significant association with CCT in either set 1 (European ancestry) or set 2 (Asian ancestry), and, owing to the small effect on CCT, meta-analysis across groups was required to detect genome-wide significance. This raises the possibility that, among the hundreds of SNPs approaching genome-wide significance, there remain additional loci that may prove clinically relevant in determining keratoconus and glaucoma risk.

The steady increase in the number of significant loci as sample size has increased for the CCT GWAS agrees with the findings that, for most quantitative traits, in a rough approximation, doubling the sample size leads to doubling of the number of significant loci. Therefore, performing genome-wide meta-analysis of CCT and of other ocular biometric parameters on an ever-larger sample set would likely lead to further insights into associated eye diseases, such as keratoconus and glaucoma.

Finally, using three different methods of pathway analysis, we showed that CCT-associated loci converge on collagen and ECM pathways (and their related pathways). We also showed that similar pathways regulate cornea thickness in both European and Asian populations. The identified collagen pathway allows rationalization of the existing findings on the associated collagen genes and suggests more collagen genes may be important for CCT. We showed that the causal genes for BCS, ZNF469 (ref. 10) and PRDM5 (ref. 36), had a role in CCT. Although neither of the genes was included in any of the GO pathway definitions, recent microarray expression analysis showed that both genes participate in a pathway regulating ECM development and maintenance. Therefore, the ECM pathway probably has a significant role in the regulation of CCT.

Taken together, the results from this meta-analysis further illuminate the genetic architecture of CCT and reveal that CCT-associated loci convey eye disease risk.

Note: Supplementary information is available in the online version of the paper.
The role of dermatopontin in the stromal organization of the cornea. A versatile gene-based test for genome-wide association studies.

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Meta-analysis of set 1 (13,057 individuals with European ancestry unaffected by eye disease). Details of the specific studies are provided in the Supplementary Note. The majority of the sites controlled for demographic variables, such as age and sex. Additional covariates that were controlled in the analysis, for example, principal components, study site or CCT measurement, were variable across individual studies. Most of the study sites have imputed data available (with the HapMap phase 2 CEU population set as the reference). Before the meta-analysis, additional checks were imposed on the SNPs, including for MAF (MAF > 1%) and imputation quality ($R^2$ from IMPUTE/SNPTST > 0.3), and we required that SNPs be autosomal and not strand ambiguous if strand information was not available. We also compared the sample allele frequencies with HapMap allele frequencies to ensure that the allele frequencies were concordant to the ones from the reference. The quantile–quantile plot and the genomic control parameters from individual studies were also examined to assure no obvious inflation in association $P$ values owing to residual population stratification or other confounding factors at each site.

We applied inverse variance–based meta-analysis, using METAL, to combine individual association results in each of the four sample sets. Even though the genomic control parameter from each study was close to 1, we applied genomic control correction within each study before inclusion in the meta-analysis. A second genomic control correction on the meta-analysis statistics was suggested to be overly conservative; therefore, we did not apply it in this study. The overall $λ$ value was 1.048 (Supplementary Fig. 2). The heterogeneity test based on Cochran's $Q$ test was also used to examine whether observed effect sizes were homogeneous across samples. We also searched for the proxies of the most associated variants, that is, the genotyped SNPs that were in high LD with the lead SNPs (Supplementary Table 2). This process was carried out to determine whether there was any spurious association due to genotyping or imputation artifacts.

We separated the Australian and UK twin studies from set 1, so that the remaining set 1 samples were analyzed as a discovery set, and Australian and UK twin studies were analyzed as an independent target set. Because all associated variants from the discovery set had the same effect direction and nominal significance in the target set, we conducted a meta-analysis on the whole set referred to as combined set 1 (Table 1). Regional association plots are shown in Supplementary Figure 3a–m.

We also applied the recent approach of approximate conditional and joint multiple-SNP analysis to the set 1 meta-analysis results. This approach uses the genotypes from a large reference set to approximate LD, instead of requiring the actual genotypes from the study samples as in conventional conditional analysis. Here, we used the QIMR cohort as the reference, which included 3,924 unrelated individuals with European ancestry and with genotypes available for 2,410,957 SNPs.

Testing CCT-associated loci in set 2 (6,963 Asian samples unaffected by eye disease). To investigate whether the associated variants identified in European populations had similar effects on corneal thickness in Asian individuals, we collected samples from three Asian populations (Indian, Malay and Chinese) and tested the CCT-associated loci identified from set 1 in these samples. These Asian samples were imputed using HapMap reference panels: the phase 2 JPT and CHB populations as an approximate reference to identify proxies for the variants that were not genotyped or imputed in any of the Asian populations. To increase power, we conducted a meta-analysis on the three Asian data sets using the same scheme as described for the meta-analysis of the set 1 samples (Supplementary Table 3).

Meta-analysis of set 1 and set 2. We performed a meta-analysis on the normal samples with European and Asian ancestry using Fisher's method (Table 2). This method combines the $P$ values from the two sets of results; therefore, it evaluates the strength of association instead of weighting on the basis of effect size. We also performed this meta-analysis using the inverse–variance weighting method—this method is potentially more powerful than Fisher's method—but is inappropriate if the trait distribution or allele frequencies vary across studies. We considered loci to be independent if LD was very low ($r^2 < 0.01$) in both European and Asian samples.

Testing CCT-associated loci in the clinical cohorts (set 3: 1,936 POAG cases with European ancestry and set 4: 198 normal-tension glaucoma cases with Asian ancestry). We analyzed the set of individuals with glaucoma for separate association testing to determine whether the CCT-associated loci identified in the general population also influenced CCT values in glaucoma cases. The diagnostic criteria for POAG are provided in the Supplementary Note. We performed meta-analysis on the set 3 samples, which included five sets of POAG cases with European ancestry, using a similar protocol to that described for the meta-analysis of set 1 (Supplementary Table 4). An additional 198 NTG cases, all of Chinese ancestry, formed set 4 and were tested for CCT-associated loci separately (Supplementary Table 4).

Testing CCT-associated loci in glaucoma case-control studies and keratoconus case-control studies. To investigate whether the CCT-associated loci were relevant to disease, we directly tested the lead SNPs for association with risk of two common ophthalmological disorders, glaucoma and keratoconus. The sample included 652 keratoconus cases from Australia ($n = 517$) and Northern Ireland ($n = 135$), who were genotyped in 2 Sequenom multiplexes at the Australian Genotyping Research Facility, 2,761 samples typed on Illumina HumanHap610 arrays from the Blue Mountains Eye Study were used as controls. US keratoconus cases ($n = 222$) and controls ($n = 3,324$) were genotyped on Illumina HumanHap370 arrays. The combined test of association was determined by a weighted fixed-effects meta-analysis of the two clinical cohorts (Supplementary Table 11). Similarly, we obtained three cohorts to evaluate POAG risk, 590 advanced POAG cases and 3,956 controls from South Australia, and GLAUGEN and NEIGHBOR case-control collections, including 1,669 and 720 high-tension and normal-tension glaucoma cases and 3,443 controls (Supplementary Table 12). The diagnostic criteria for both diseases are provided in the Supplementary Note.

Pathway analysis. We used three pathway analysis approaches, an extension of the VEGAS gene-based test to pathway analysis (VEGAS-Pathway), MAGENTA and GRAIL.

For VEGAS-Pathway analysis, we first performed gene-based tests using VEGAS software on the set 1 and set 2 meta-analysis results separately. VEGAS incorporates information from the full set of markers within a gene and accounts for LD between markers by using simulations from the multivariate normal distribution. Because the set 1 samples were all of European descent, we used the HapMap 2 CEU population as the reference to estimate patterns of LD. The set 2 samples were from three Asian ancestry groups; however, we used the combined HapMap 2 JPT and CHB populations as the reference to approximate LD patterns. The gene regulatory region was defined as 50 kb both upstream and downstream of a gene. The gene-based results for meta-analysis of European samples were presented for the genes of interest, including known CCT-associated loci, newly identified loci and their neighboring genes (Supplementary Table 5).

Pathway analysis was carried out using prespecified pathways from the GO database. Pathways with 10 to 1,000 genes were selected, yielding 4,628 pathways. Pathway analysis was based on combining gene-based test results from VEGAS. Pathway $P$ values were computed by summing $χ^2$ test statistics derived from VEGAS $P$ values. Empirical VEGAS-Pathway $P$ values for each pathway were computed by comparing the summed $χ^2$ test statistics from real data with those generated in 500,000 simulations where the relevant number (according to the size of the pathway) of randomly drawn $χ^2$ test statistics was summed. To ensure that clusters of genes did not adversely affect results, within each pathway, gene sets were pruned such that each gene was $>500$ kb.
Previously published microarray data were used to define the connectivity between the CCT-associated loci available in MAGENTA were used to query the meta-analysis results from the gene regulatory region used for VEGAS was maintained here, that is, 50 kb both upstream and downstream of a gene. The first difference in the methods used in MAGENTA and in VEGAS-Pathway is that MAGENTA assigns the best-SNP P value within the gene regulatory region as the gene score and corrects this gene score for confounding factors, such as gene size, whereas VEGAS takes the full set of markers and accounts for LD between markers by simulation. The second difference is in how the gene-based results are used to form a pathway test statistic. VEGAS-Pathway uses the full set of VEGAS P values within a given pathway to form the test statistic. In contrast, MAGENTA uses a non-parametric approach to conduct the pathway enrichment test, only using genes that reach a prespecified enrichment cutoff to construct a test statistic. We used the 95th percentile of all gene scores as the enrichment cutoff. The GO and PANTHER (2010) databases available in MAGENTA were used to query the meta-analysis results from the set 1 samples of European ancestry (Supplementary Table 7).

GRAIL is an algorithm examining the functional connection between associated loci. We queried the GO database (available in GRAIL) to identify the key pathways that show the connectivity between the CCT-associated loci listed in Table 2 (Supplementary Table 10).

**Polygenic modeling.** We applied the previously described polygenic approach, using CCT meta-analysis results as the discovery set, to the following target sets: ANZRAG subjects (to test whether aggregation of CCT effects predicts CCT values in individuals with POAG), the US keratoconus case-control set (to test whether aggregation of CCT effects predicts keratoconus risk) and ANZRAG, GLAUGEN and NEIGBOR case-control sets (similarly, to test whether CCT effects predict POAG risk on a polygenic basis). We used genotyped data in the target sets, retaining only SNPs with clear, non-ambiguous strand coding. The polygenic profile score for each individual in the target sets was calculated as the summation of the SNP genotypes in different P-value bins defined by the CCT meta-analysis, weighted by the CCT effects. Finally, polygenic scores were used to predict trait values in the target sets. When the target set included keratoconus or POAG cases and controls, logistic regression was used to assess association between the disease trait and the polygenic profile score. Analogously, when testing CCT in POAG cases, linear regression was used.

**Gene expression analyses.** Previously published microarray data were used to initially characterize the expression of genes associated with lead SNPs. To confirm the mouse microarray data, quantitative RT-PCR (qRT-PCR) analysis was performed using independent tissue samples. Corneas were dissected at the limbus from enucleated eyes in PBS, and the corneas from each mouse were pooled to form one sample; three samples (mice) were analyzed per strain. Corneal samples were homogenized, and RNA was extracted, treated with DNase I, purified (Aurum Total RNA Fatty and Fibrous Tissue Pack, Bio-Rad) and converted to cDNA (iScript cDNA Synthesis kit, Bio-Rad). Quantitative PCR was performed with a SYBR green master mix (iQ SYBR Green Supermix, Bio-Rad) in an RT-PCR detection system (CFX Connect, Bio-Rad). Each reaction contained 5.9 µl of water, 7.5 µl of 2× iQ SYBR green master mix, 0.3 µl of 5′-primer (5 µM), 0.3 µl of 3′ primer (5 µM) and 1 µl of cDNA (2 ng/µl). Sequences for the primer pairs used in the PCR reactions are available upon request. PCR conditions involved incubation at 95 °C for 3 min and 40 cycles of 95 °C for 10 s, 55 °C for 10 s and 72 °C for 30 s. PCR products were subjected to melting curve analysis to ensure that only a single product was amplified. Each experiment included three technical replicates of each RNA sample. Expression data were quantified on the basis of threshold cycle (Ct) values. For each transcript, Ct values for each sample were averaged and normalized to the values for β-actin (ACTB). Change analysis was based on ΔΔCt values and the amplification efficiency of the transcripts.