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Authors

Kamitaki, Nolan Sekar, Aswin Handsaker, Robert E <u>et al.</u>

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Complement genes contribute sex-biased vulnerability in diverse illnesses

Nolan Kamitaki^{1,2}, Aswin Sekar^{1,2}, Robert E. Handsaker^{1,2}, Heather de Rivera^{1,2}, Katherine Tooley^{1,2}, David L. Morris³, Kimberly E. Taylor⁴, Christopher W. Whelan^{1,2}, Philip Tombleson³, Loes M. Olde Loohuis^{5,6}, Schizophrenia Working Group of the Psychiatric Genomics Consortium⁷, Michael Boehnke⁸, Robert P. Kimberly⁹, Kenneth M. Kaufman¹⁰, John B. Harley¹⁰, Carl D. Langefeld¹¹, Christine E. Seidman^{1,12,13}, Michele T. Pato¹⁴, Carlos N. Pato¹⁴, Roel A. Ophoff^{5,6}, Robert R. Graham¹⁵, Lindsey A. Criswell⁴, Timothy J. Vyse³, Steven A. McCarroll^{1,2}

¹Department of Genetics, Harvard Medical School, Boston, Massachusetts 02115, USA

²Stanley Center for Psychiatric Research, Broad Institute of MIT and Harvard, Cambridge, Massachusetts 02142, USA

³Department of Medical and Molecular Genetics, King's College London, London WC2R 2LS, UK

⁴Rosalind Russell / Ephraim P Engleman Rheumatology Research Center, Division of Rheumatology, UCSF School of Medicine, San Francisco, California 94143, USA

⁵Department of Human Genetics, David Geffen School of Medicine, University of California, Los Angeles, California 90095, USA

⁶Center for Neurobehavioral Genetics, Semel Institute for Neuroscience and Human Behavior, University of California, Los Angeles, California 90095, USA

⁷A full list of collaborators is in Schizophrenia Working Group of the Psychiatric Genomics Consortium.

Competing interests

Supplementary Information is available for this paper.

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Correspondence and requests for materials should be addressed to Nolan Kamitaki (nolan_kamitaki@hms.harvard.edu), Timothy J. Vyse (timothy.vyse@kcl.ac.uk) and Steven A. McCarroll (mccarroll@hms.harvard.edu). Author Contributions

N.K., A.S., T.J.V., and S.A.M. conceived the genetic studies. M.T.P., C.N.P., and M.B. collected and contributed whole-genome sequence data for the Genomic Psychiatry Cohort. R.E.H. and C.W.W. genotyped *C4* structural variation in the Genomic Psychiatry Cohort and optimized variant selection for use as a reference panel in the imputation of *C4* variation into lupus and schizophrenia cohorts (Extended Data Fig. 1). T.J.V., R.R.G., L.A.C., C.D.L., R.P.K., J.B.H., K.M.K., D.L.M., and P.T. contributed genotype data and imputation of non-*C4* variation for analysis of SLE cohorts. K.E.T. and L.A.C. contributed genotype and phenotype data along with imputation of non-*C4* variation for analysis of the SJS cohort. Investigators in the Schizophrenia Working Group of the Psychiatric Genomics Consortium collected and phenotyped cohorts and contributed genotype data for analysis of schizophrenia cohorts. N.K did the imputation and association analysis (Figs. 1, 2, 3a, b and Extended Data Figs. 2, 3, 4, 5, and 6). T.J.V., R.R.G., and D.L.M. provided valuable advice on the analysis and interpretation of SLE association results. R.A.O. and L.M.O.L collected and provided CSF samples composing the group from Utrecht, Netherlands. C.E.S. collected and provided CSF samples composing the Brigham & Women's Hospital group. H.d.R and K.T. performed the C4 and C3 immunoassay experiments on CSF samples (Fig. 3c, d and Extended Data Fig. 7a). N.K. did the analysis of plasma C4 and C3 concentrations (Extended Data Fig. 7b–f). S.A.M and N.K. wrote the manuscript with contributions from all authors.

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⁸Department of Biostatistics and Center for Statistical Genetics, University of Michigan, Ann Arbor, Michigan 48109, USA

⁹Division of Clinical Immunology and Rheumatology, University of Alabama at Birmingham, Birmingham, Alabama 35294, USA

¹⁰Center for Autoimmune Genomics and Etiology (CAGE), Department of Pediatrics, Cincinnati Children's Medical Center & University of Cincinnati and the US Department of Veterans Affairs Medical Center, Cincinnati, Ohio, USA

¹¹Department of Biostatistical Sciences, Wake Forest School of Medicine, Winston-Salem, North Carolina 27101, USA

¹²Howard Hughes Medical Institute, Chevy Chase, Maryland 20815, USA

¹³Cardiovascular Division, Brigham and Women's Hospital, Boston, Massachusetts 02115, USA

¹⁴SUNY Downstate Medical Center, Brooklyn, New York 11203, USA

¹⁵Human Genetics, Genentech Inc, South San Francisco, California 94080, USA

Abstract

Many common illnesses differentially affect men and women for unknown reasons. The autoimmune diseases lupus and Sjögren's syndrome affect nine times more women than men¹, whereas schizophrenia affects men more frequently and severely². All three illnesses have their strongest common genetic associations in the Major Histocompatibility Complex (MHC) locus, an association that in lupus and Sjögren's syndrome has long been thought to arise from alleles of the human leukocyte antigen (HLA) genes at that locus^{3–6}. Here we show that the complement component 4 (C4) genes, which are also in the MHC locus and were recently found to increase risk for schizophrenia⁷, generate 7-fold variation in risk for lupus (95% CI: 5.88–8.61; $p < 10^{-117}$ in total) and 16-fold variation in risk for Sjögren's syndrome (95% CI: 8.59–30.89; $p < 10^{-23}$ in total) among individuals with common C4 genotypes, with C4A protecting more strongly than C4B in both illnesses. The same alleles that increase risk for schizophrenia greatly reduced risk for lupus and Sjögren's syndrome. In all three illnesses, C4 alleles acted more strongly in men than in women: common combinations of C4A and C4B generated 14-fold variation in risk for lupus, 31fold variation in risk for Sjögren's syndrome, and 1.7-fold variation in schizophrenia risk among men (vs. 6-fold, 15-fold, and 1.26-fold among women respectively). At a protein level, both C4 and its effector C3 were present at greater levels in men than women in cerebrospinal fluid (p <10⁻⁵ for both C4 and C3) and plasma^{8,9} among adults ages 20–50, corresponding to the ages of differential disease vulnerability. Sex differences in complement protein levels may help explain the larger effects of C4 alleles in men, women's greater risk of SLE and Sjögren's, and men's greater vulnerability in schizophrenia. These results implicate the complement system as a source of sexual dimorphism in vulnerability to diverse illnesses.

Systemic lupus erythematosus (SLE, or "lupus") is a systemic autoimmune disease of unknown cause. Risk of SLE is heritable ($66\%^{10}$), although SLE may have environmental triggers, as its onset often follows events that damage cells, such as infections and severe

sunburns¹¹. Most SLE patients produce autoantibodies against nucleic acid complexes, including ribonucleoproteins and DNA¹².

In genetic studies, SLE associates most strongly with variation across the major histocompatibility complex (MHC) locus, which contains the human leukocyte antigen (*HLA*) genes³. However, conclusive attribution of this association to specific genes and alleles has been difficult; the identities of the most likely genetic sources have been frequently revised as genetic studies have grown in size^{4,5}. In several other autoimmune diseases, including type 1 diabetes, celiac disease, and rheumatoid arthritis, strong effects of the MHC locus arise from *HLA* alleles that cause the peptide binding groove of HLA proteins to present a disease-critical autoantigen^{13,14}. In SLE, by contrast, genetic variants in the MHC locus (including SNPs and *HLA* alleles) associate broadly with the presence of diverse autoantibodies¹⁵.

The complement component 4 (*C4A* and *C4B*) genes are also present in the MHC genomic region, between the class I and class II *HLA* genes. Classical complement proteins help eliminate debris from dead and damaged cells, attenuating the visibility of diverse intracellular proteins to the adaptive immune system. *C4A* and *C4B* commonly vary in genomic copy number¹⁶ and encode complement proteins with distinct affinities for molecular targets^{17,18}. SLE frequently presents with hypocomplementemia that worsens during flares, possibly reflecting increased active consumption of complement¹⁹. Rare cases of severe, early-onset SLE can involve complete deficiency of a complement component (C4, C2, or C1Q)^{20,21}, and one of the strongest common-variant associations in SLE maps to *ITGAAM*, which encodes a receptor for C3, the effector of C4²². Although total *C4* gene copy number associates with SLE risk^{23,24}, this association is thought to arise from linkage disequilibrium (LD) with alleles of nearby *HLA* genes²⁵, which have been the focus of finemapping analyses^{3,4}.

The complex genetic variation at C4 – arising from many alleles with different numbers of C4A and C4B genes – has been challenging to analyze in large cohorts. A recently feasible approach to this problem is based on imputation: people share long haplotypes with the same combinations of SNP and C4 alleles, such that C4A and C4B gene copy numbers can be imputed from SNP data⁷. To analyze C4 in large cohorts, we developed a way to identify C4 alleles from whole-genome sequence (WGS) data (Extended Data Fig. 1a, b), then analyzed WGS data from 1,265 individuals (from the Genomic Psychiatry Cohort^{26,27}) to create a large multi-ancestry panel of 2,530 reference haplotypes of MHC-region SNPs and C4 alleles (Extended Data Fig. 1c) – ten times more than in earlier work⁷. We then analyzed SNP data from the largest SLE genetic association study³ (ImmunoChip 6,748 SLE cases and 11,516 controls of European ancestry) (Extended Data Fig. 2a, b), imputing C4 alleles to estimate the SLE risk associated with common combinations of C4A and C4B gene copy numbers (Fig. 1a).

Groups of research participants with the eleven most common combinations of *C4A* and *C4B* gene copy number exhibited 7-fold variation in their relative risk of SLE (Fig. 1a, Extended Data Fig. 2c). The relationship between SLE risk and *C4* gene copy number exhibited consistent, logical patterns across the 11 genotype groups. For each *C4B* copy

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number, greater *C4A* copy number associated with reduced SLE risk (Fig. 1a, Extended Data Fig. 2c). For each *C4A* copy number, greater *C4B* copy number associated with more modestly reduced risk (Fig. 1a). Logistic-regression analysis estimated that the protection afforded by each copy of *C4A* (OR: 0.54; 95% CI: [0.51, 0.57]) was equivalent to that of 2.3 copies of *C4B* (OR: 0.77; 95% CI: [0.71,0.82]). We calculated an initial *C4*-derived risk score as 2.3 times the number of *C4A* genes, plus the number of *C4B* genes, in an individual's genome. Despite clear limitations of this risk score – it is imperfectly imputed from flanking SNP haplotypes ($t^2 = 0.77$, Extended Data Table 1) and only approximates *C4*-derived risk by using a simple, linear model (to avoid over-fitting the genetic data) – SNPs across the MHC genomic region tended to associate with SLE in proportion to their level of LD with this risk score (Extended Data Fig. 3a).

Combinations of many different C4 alleles generate the observed variation in C4A and C4B gene copy number; particular C4A and C4B gene copy numbers have also arisen recurrently on multiple SNP haplotypes⁷ (Extended Data Fig. 1c). Analysis of SLE risk in relation to each of these C4 alleles and SNP haplotypes reinforced the conclusion that C4A contributes strong protection, and C4B more modest protection, from SLE, and that C4 genes (rather than nearby variants) are the principal drivers of this variation in risk levels (Fig. 1b).

These results prompted us to consider whether other autoimmune disorders with similar patterns of genetic association at the MHC genomic region might also be driven in part by C4 variation. Primary Sjögren's syndrome (SjS) is a heritable $(54\%^{28})$ systemic autoimmune disorder of exocrine glands, characterized primarily by dry eyes and mouth with other systemic effects. At a protein level, SjS is (like SLE) characterized by diverse autoantibodies, including antinuclear antibodies targeting ribonucleoproteins²⁹, and hypocomplementemia³⁰. The largest source of common genetic risk for SiS lies in the MHC genomic locus³¹, with associations to the same haplotype(s) as in SLE⁶ and with heterogeneous HLA associations in different ancestries³². We imputed C4 alleles into existing SNP data from a European-ancestry SjS case-control cohort (673 cases and 1153 controls). As in SLE, logistic-regression analyses found both C4A copy number (OR: 0.41; 95% CI: [0.34, 0.49]) and C4B copy number (OR: 0.67; 95% CI: [0.53, 0.86]) to be protective against SjS. The risk-equivalent ratio of C4B to C4A gene copies was similar in SjS and SLE (about 2.3 to 1); also, as with SLE, nearby SNPs associated with SjS in proportion to their LD with a C4-derived risk score ((2.3)C4A+C4B) (Extended Data Fig. 3b). The distribution of SjS risk across the individual C4 alleles and haplotypes revealed a pattern that, as in SLE, supported greater protective effect from C4A than C4B, and little effect of flanking SNP haplotypes (Fig. 1b).

The association of SLE and SjS with *C4* gene copy number has long been attributed to the *HLA-DRB1**03:01 allele. In European populations, *DRB1**03:01 is in strong LD ($r^2 = 0.71$) with the common *C4*-B(S) allele, which lacks any *C4A* gene and is the highest-risk *C4* allele in our analysis (Fig. 1b); many MHC-region SNPs associated with SLE and SjS in proportion to their LD correlations with both *C4* and *DRB1**03:01 (Extended Data Fig. 4a, b). Cohorts with other ancestries can have recombinant haplotypes that disambiguate the contributions of alleles that are in LD in Europeans. Among African Americans, we found that common *C4* alleles exhibited far less LD with *HLA* alleles; in particular, the LD

between C4-B(S) and DRB1*03:01 was low ($r^2 = 0.10$) (Extended Data Table 2). Thus, genetic data from an African American SLE cohort (1,494 cases, 5,908 controls) made it possible to distinguish between these potential genetic effects. Joint association analysis of C4A, C4B, and DRB1*0301 implicated C4A ($p < 10^{-14}$) and C4B ($p < 10^{-5}$) but not DRB1*0301 (p = 0.29) (Extended Data Table 3). Each C4 allele associated with effect sizes of similar magnitude on SLE risk in Europeans and African Americans (Fig. 2a). An analysis specifically of combinations of C4-B(S) and DRB1*03:01 allele dosages in African Americans showed that C4-B(S) alleles consistently increased SLE risk regardless of DRB1*03:01 status, whereas DRB1*03:01 had no consistent effect when controlling for C4-B(S) (Fig. 2b). Although C4 alleles had less LD with nearby variants on African American than on European haplotypes, SNPs across the genomic region associated with SLE in proportion to LD correlations with C4 in African Americans as well (Extended Data Fig. 4c).

Accounting for *C4* alleles in jointly analyzing the SLE association data from African American and European ancestry cohorts also enabled the mapping of an additional, moremodest genetic effect independent of *C4*; this effect (tagged by rs2105898 and rs9271513) appeared to involve noncoding variation in the *HLA* class II XL9 region that associates most strongly with expression levels (rather than the coding sequence) of many *HLA* class II genes (Extended Data Figs. 3c, d, 4d–l, 5, and Supplementary Note 1).

Alleles at *C4* that increase dosage of *C4A* (and to a more modest extent *C4B*) appear to protect strongly against SLE and SjS (Fig. 1a, b); by contrast, alleles that increase expression of *C4A* in the brain are more common among research participants with schizophrenia⁶. These same illnesses exhibit striking, and opposite, sex differences: SLE and SjS are nine times more common among women of childbearing age than among men of a similar age¹, whereas in schizophrenia, women exhibit less severe symptoms, more frequent remission of symptoms, lower relapse rates, and lower overall incidence². Though the vast majority of genetic associations in complex diseases are shared between men and women³³, the SNPs that most strongly associate with SLE risk within the MHC region associate to larger potential effect sizes in men³⁴. Hence, we sought to evaluate the possibility that the effects of *C4* alleles on risk in SLE, SjS, and schizophrenia might differ between men and women.

Analysis indicated that the effects of *C4* alleles were stronger in men. When a sex-by-*C4* interaction term was included in association analyses, this term was significant for both SLE (p = 0.002) and schizophrenia (p = 0.0024), with larger *C4* effects in men for both disorders. (Analysis of SjS had limited power due to the small number of men affected by SjS.) For both SLE and schizophrenia, the individual *C4* alleles consistently associated with stronger effects in men than women (Fig. 3a, b). SNPs across the MHC genomic region exhibited sex-biased association to SLE, SjS, and schizophrenia to the extent of their LD with *C4* (Extended Data Fig. 6a–c).

The stronger effects of C4 alleles on male relative to female risk could arise from sex differences in C4 RNA expression, C4 protein levels, or downstream responses to C4. Analysis of RNA expression in human tissues, using data from GTEx³⁵, identified no sex

differences in *C4* RNA expression in brain, blood, liver, or lymphoblastoid cells (a moredetailed description of this analysis can be found in Supplementary Note 2). We then analyzed C4 protein in cerebrospinal fluid (CSF) from two panels of adult research participants (n = 589 total) in whom we had also measured *C4* gene copy number (by direct genotyping or imputation). CSF C4 protein levels correlated strongly with *C4* gene copy number ($p < 10^{-10}$, Extended Data Fig. 7a), so we normalized C4 protein measurements to the number of *C4* gene copies. CSF from adult men contained on average 27% more C4 protein per *C4* gene copy than CSF from women (meta-analysis $p = 9.9 \times 10^{-6}$, Fig. 3c). C4 acts by activating the complement component 3 (C3) protein, promoting C3 deposition onto targets in tissues. CSF levels of C3 protein were also on average 42% higher among men than women (meta-analysis $p = 7.5 \times 10^{-7}$, Fig. 3d).

The elevated concentrations of C3 and C4 proteins in CSF of men parallel earlier findings that, in plasma, C3 and C4 are also present at higher levels in men than women^{8,9}. The large sample size (n > 50,000) of the plasma studies allows sex differences to be further analyzed as a function of age. Both men and women undergo age-dependent elevation of C4 and C3 levels in plasma, but this occurs early in adulthood (age 20-30) in men and closer to menopause (age 40–50) in women, with the result that male-female differences in complement protein levels are observed primarily during the reproductive years (ages 20-50)^{8,9}. We replicated these findings using measurements of C3 and gene copy numbercorrected C4 protein in plasma from adults, finding (as in the earlier plasma studies^{8,9} and in CSF, Fig. 3c, d) that these differences are most pronounced during the reproductively active years of adulthood (ages 20-50) (Extended Data Fig. 7b-d). We also observed that SjS patients have lower C4 serum levels than controls ($p < 1 \times 10^{-20}$, Extended Data Fig. 7e) even after correcting for C4 gene copy number ($p \le 1 \times 10^{-8}$, Extended Data Fig. 7f), suggesting that hypocomplementemia in SjS is not simply due to C4 genetics but also reflects disease effects on ambient complement levels, for example due to complement consumption. The ages of pronounced sex difference in complement levels corresponded to the ages at which men and women differ in disease incidence: in schizophrenia, men outnumber women among cases incident in early adulthood, but not among cases incident after age 40^2 ; in SLE, women greatly outnumber men among cases incident during the child-bearing years, but not among cases incident after age 50 or during childhood³⁶; in SjS, the large relative vulnerability of women declines in magnitude after age 50^{37} .

Our results indicate that the MHC genomic region shapes vulnerability in lupus and SjS – two of the three most common rheumatic autoimmune diseases – in a very different way than in type I diabetes, rheumatoid arthritis, and celiac disease. In the latter diseases, precise interactions between specific *HLA* protein variants and specific autoantigens determine risk 13,14 . In SLE and SjS, however, the genetic variation implicated here points instead to the continuous, chronic interaction of the immune system with very many potential autoantigens. Because complement facilitates the rapid clearance of debris from dead and injured cells, elevated levels of C4 protein likely attenuate interactions between the adaptive immune system and ribonuclear self-antigens at sites of cell injury, pre-empting the development of autoimmunity. The additional *C4*-independent genetic risk effect described here (associated with rs2105898) may also affect autoimmunity broadly, rather than antigenspecifically, by regulating expression of many HLA class II genes (including *DRB1*, *DQA1*,

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and *DQB1*). Mouse models of SLE indicate that once tolerance is broken for one selfantigen, autoreactive germinal centers generate B cells targeting other self-antigens³⁸; such "epitope spreading" could lead to autoreactivity against many related autoantigens, regardless of which antigen(s) are involved in the earliest interactions with immune cells. Further supporting such a model, higher copy number of *C4* associates with lower risk of AQP4-IgG-seropositive neuromyelitis optica (NMO-IgG+)³⁹, in which seropositive patients have increased incidence of other non-organ-specific autoantibodies such as those seen in SLE and SjS⁴⁰. B-cells also express the complement receptors CR1 and CR2⁴¹, providing an additional candidate mechanism for regulation by C4 and C3.

We note that the role of complement proteins in preventing the emergence of autoimmunity may be very different than their (potentially disease-exacerbating) role once autoimmunity has been established. Also, our genetic findings address the development of SLE and SjS rather than complications that arise in any specific organ. A few percent of SLE patients develop neurological complications that can include psychosis⁴²; though psychosis is also a symptom of schizophrenia, neurological complications of SLE do not resemble schizophrenia more broadly, and likely have a different etiology.

The same C4 alleles that increase vulnerability to schizophrenia appeared to protect strongly against SLE and SjS. This pleiotropy will be important to consider in efforts to engage the complement system therapeutically. The complement system contributed to these pleiotropic effects more strongly in men than in women. Moreover, though the natural allelic series at C4 allowed human-genetic analysis to establish dose-risk relationships for C4 in men and women, sexual dimorphism in the complement-protein levels also included complement component 3 (C3). Why and how biology has come to create this sexual dimorphism in the complement system in humans presents interesting questions for immune and evolutionary biology.

Methods

Creation of a C4 reference panel from whole-genome sequence data

We constructed a reference panel for imputation of C4 structural haplotypes using wholegenome sequencing data for 1265 individuals from the Genomic Psychiatry Cohort²⁶. The reference panel included individuals of diverse ancestry, including 765 Europeans, 250 African Americans, and 250 people of reported Latino ancestry.

We estimated the diploid *C4* copy number, and separately the diploid copy number of the contained HERV segment, using Genome STRiP⁴⁴. Briefly, Genome STRiP carefully calibrates measurements of read depth across specific genomic segments of interest by estimating and normalizing away sample-specific technical effects such as the effect of GC content on read depth (estimated from the genome-wide data). To estimate *C4* copy number, we genotyped the segments 6:31948358–31981050 and 6:31981096–32013904 (hg19) for total copy number, but masked the intronic HERV segments that distinguish short (S) from long (L) *C4* gene isotypes. For the HERV region, we genotyped segments 6:31952461–31958829 and 6:31985199–31991567 (hg19) for total copy number. Across the 1,265 individuals, the resultant locus-specific copy-number estimates exhibited a strongly multi-

modal distribution (Extended Data Fig. 1a) from which individuals' total C4 copy numbers could be readily inferred.

We then estimated the ratio of *C4A* to *C4B* genes in each individual genome. To do this, we extracted reads mapping to the paralogous sequence variants that distinguish *C4A* from *C4B* (hg19 coordinates 6:31963859–31963876 and 6:31996597–31996614) in each individual, combining reads across the two sites. We included only reads that aligned to one of these segments in its entirety. We then counted the number of reads matching the canonical active site sequences for *C4A* (CCC TGT CCA GTG TTA GAC) and *C4B* (CTC TCT CCA GTG ATA CAT). We combined these counts with the likelihood estimates of diploid *C4* copy number (from Genome STRiP) to determine the maximum likelihood combination of *C4A* and *C4B* estimate from the likelihood ratio between the most likely and second most likely combinations.

To phase the *C4* haplotypes, we first used the GenerateHaploidCNVGenotypes utility in Genome STRiP to estimate haplotype-specific copy-number likelihoods for *C4* (total *C4* gene copy number), *C4A*, *C4B*, and HERV using the diploid likelihoods from the prior step as input. Default parameters for GenerateHaploidCNVGenotypes were used, plus - genotypeLikelihoodThreshold 0.0001. The output was then processed by the GenerateCNVHaplotypes utility in Genome STRiP to combine the multiple estimates into likelihood estimates for a set of unified structural alleles. GenerateCNVHaplotypes was run with default parameters, plus -defaultLogLikelihood –50, -unknownHaplotypeLikelihood –50, and -sampleHaplotypePriorLikelihood 2.0. The resultant VCF was phased using Beagle 4.1 (beagle_4.1_27Jul16.86a) in two steps: first, performing genotype refinement from the genotype likelihoods using the Beagle gtgl= and maxlr=1000000 parameters, and then running Beagle again on the output file using gt= to complete the phasing.

Our previous work suggested that several C4 structures segregate on different haplotypes, and probably arose by recurrent mutation on different haplotype backgrounds⁷. The GenerateCNVHaplotypes utility requires as input an enumerated set of structural alleles to assign to the samples in the reference cohort, including any structurally equivalent alleles, with distinct labels to mark them as independent, plus a list of samples to assign (with high likelihood) to specific labeled input alleles to disambiguate among these recurrent alleles. The selection of the set of structural alleles to be modeled, along with the labeling strategy, is important to our methodology and the performance of the reference panel. In the reference panel, each input allele represents a specific copy number structure and optionally includes a label that differentiates the allele from other independent alleles with equivalent structure. We use the notation $\langle H_n_n_L \rangle$ to identify each allele, where the four integers following the H are, respectively, the (redundant) haploid count of the total number of C4 copies, C4A copies, C4B copies and HERV copies on the haplotype. For example, <H 2 1 1 1> was used to represent the "AL-BS" haplotype. The optional final label L is used to distinguish potentially recurrent haplotypes with otherwise equivalent structures (under the model) that should be treated as independent alleles for phasing and imputation.

To build the reference panel, we experimentally evaluated a large number of potential sets of structural alleles and methods for assigning labels to potentially recurrent alleles. For each evaluation, we built a reference panel using the 1265 reference samples, and then evaluated the performance of the panel via cross-validation, leaving out 10 different samples in each trial (5 samples in the last trial) and imputing the missing samples from the remaining samples in the panel. The imputed results for all 1265 samples were then compared to the original diploid copy number estimates to evaluate the performance of each candidate reference panel (Extended Data Table 1).

Using this procedure, we selected a final panel for downstream analysis that used a set of 29 structural alleles representing 16 distinct allelic structures (as listed in the reference panel VCF file). Each allele contained from one to three copies of *C4*. Three allelic structures (AL-BS, AL-BL, and AL-AL) were represented as a set of independently labeled alleles with 9, 3, and 4 labels, respectively.

To identify the number of labels to use on the different alleles and the samples to "seed" the alleles, we generated "spider plots" of the *C4* locus based on initial phasing experiments run without labeled alleles, and then clustered the resulting haplotypes in two dimensions based on the Y-coordinate distance between the haplotypes on the left and right sides of the spider plot. Clustering was based on visualizing the clusters (Extended Data Fig. 1c) and then manually choosing both the number of clusters (labels) to assign and a set of confidently assigned haplotypes to use to "seed" the clusters in GenerateCNVHaplotypes. This procedure was iterated multiple times using cross-validation, as described above, to evaluate the imputation performance of each candidate labeling strategy.

Within the data set used to build the reference panel, there is evidence for individuals carrying seven or more diploid copies of C4, which implies the existence of (rare) alleles with four or more copies of C4. In our experiments, attempting to add additional haplotypes to model these rare four-copy alleles reduced overall imputation performance. Consequently, we conducted all downstream analyses using a reference panel that models only alleles with up to three copies of C4. In the future, larger reference panels might benefit from modeling these rare four-copy alleles.

The reference panel will be available in dbGaP (accession # pending) with broad permission for research use.

Genetic data for SLE

For analysis of systemic lupus erythematosus (SLE), collection and genotyping of the European-ancestry cohort (6,748 cases, 11,516 controls, genotyped by ImmunoChip) as previously described³. Collection and genotyping of the African-American cohort (1,494 cases, 5,908 controls, genotyped by OmniExpress) as previously described⁵.

Genetic data for SjS

For analysis of Sjögren's syndrome (SjS), collection and genotyping of the Europeanancestry cohort (673 cases, 1,153 controls, genotyped by Omni2.5) as previously described³² and available in dbGaP under study accession number phs000672.v1.p1.

Genetic data for schizophrenia

The schizophrenia analysis made use of genotype data from 40 cohorts of European ancestry (28,799 cases, 35,986 controls) made available by the Psychiatric Genetics Consortium (PGC) as previously described⁴³. Genotyping chips used for each cohort are listed in Supplementary Table 3 of that study.

Imputation of C4 alleles

The reference haplotypes described above were used to extend the SLE, SjS, or schizophrenia cohort SNP genotypes by imputation. SNP data in VCF format were used as input for Beagle v4.1^{45,46} for imputation of *C4* as a multi-allelic variant. Within the Beagle pipeline, the reference panel was first converted to bref format. From the cohort SNP genotypes, we used only those SNPs from the MHC region (chr6:24–34 Mb on hg19) that were also in the haplotype reference panel. We used the conform-gt tool to perform strand-flipping and filtering of specific SNPs for which strand remained ambiguous. Beagle was run using default parameters with two key exceptions: we used the GRCh37 PLINK recombination map, and we set the output to include genotype probability (i.e., GP field in VCF) for correct downstream probabilistic estimation of *C4A* and *C4B* joint dosages.

Imputation of HLA alleles

For *HLA* allele imputation, sample genotypes were used as input for the R package HIBAG⁴⁷. For both European ancestry and African American cohorts, publicly available multi-ethnic reference panels generated for the most appropriate genotyping chip (i.e. Immunochip for European ancestry SLE cohort, Omni 2.5 for European ancestry SjS cohort, and OmniExpress for African American SLE cohort) were used⁴⁸. Default parameters were used for all settings. All class I and class II *HLA* genes were imputed. Output haplotype posterior probabilities were summed per allele to yield diploid dosages for each individual.

Associating single and joint C4 structural allele dosages to SLE and SjS in European ancestry individuals

The analysis described above yields dosage estimates for each of the common *C4* structural haplotypes (e.g., AL-BS, AL-AL, etc.) for each genome in each cohort. In addition to performing association analysis on these structures (Fig 1b), we also performed association analysis on the dosages of each underlying *C4* gene isotype (i.e. *C4A*, *C4B*, *C4L*, and *C4S*). These dosages were computed from the allelic dosage (DS) field of the imputation output VCF simply by multiplying the dosage of a *C4* structural haplotype by the number of copies of each *C4* isotype that haplotype contains (e.g., AL-BL contains one *C4A* gene and one *C4B* gene).

C4 isotype dosages were then tested for disease association by logistic regression, with the inclusion of four available ancestry covariates derived from genome-wide principal component analysis (PCA) as additional independent variables, PC_c.

$$logit(\theta) \sim \beta_0 + \beta_1 C 4 + \Sigma_c \beta_c P C_c + \varepsilon$$
⁽¹⁾

where θ =E[SLE]X]. For SjS, the model instead included two available multiethnic ancestry covariates from dbGaP that correlated strongly with European-specific ancestry covariates (specifically, PC5 and PC7) and smoking status as independent variables. Coefficients for relative weighting of *C4A* and *C4B* dosages were obtained from a joint logistic regression,

$$logit(\theta) \sim \beta_0 + \beta_1 C 4A + \beta_2 C 4B + \Sigma_c \beta_c P C_c + \varepsilon$$
⁽²⁾

The values per individual of $\beta_1 C4A + \beta_2 C4B$ were used as a combined C4 risk term for estimating both association strength (Extended Data Fig. 3a, b) as well as evaluating the relationship between the strength of nearby variants' association with SLE or SjS and linkage with C4 variation (Extended Data Fig. 4a–c).

Joint dosages of *C4A* and *C4B* for each individual in the same cohort were estimated by summing across their genotype probabilities of paired structural alleles that encode for the same diploid copy numbers of both *C4A* and *C4B* (Extended Data Fig. 2a, b). For each individual/genome, this yields a joint dosage distribution of *C4A* and *C4B* gene copy number, reflecting any possible imputed haplotype-level dosages with nonzero probability. Joint dosages for *C4A* and *C4B* diploid copy numbers were tested for association with SLE in a joint model with the same ancestry covariates (Fig. 1a),

$$logit(\theta) \sim \beta_0 + \Sigma_{i,j}\beta_{i,j}P(C4A = i, C4B = j) + \Sigma_c\beta_cPC_c + \varepsilon$$
(3)

Calculation of composite C4 risk for SLE

Because SLE risk strongly associated with C4A and C4B copy numbers (Fig. 1a) in a manner that can be approximated as – but is not necessarily linear or independent – a composite C4 risk score was derived by taking the weighted sum of joint C4A and C4B dosages multiplied by the corresponding effect sizes from the aforementioned model of the joint C4A and C4B diploid copy numbers. The weights for calculating this composite C4 risk term were computed from the data from the European ancestry cohort, and then applied unchanged to analysis of the African American cohort.

Associations of variants across the MHC region to SLE and SjS

Genotypes for non-array SNPs were imputed with IMPUTE2 using the 1000 Genomes reference panel; separate analyses were performed for the European-ancestry and African American cohorts. Unless otherwise stated, all subsequent SLE analyses were performed identically for both European ancestry and African American cohorts. Dosage of each variant, v_i , was tested for association with SLE or SjS in a logistic regression including available ancestry covariates (and smoking status for SjS) first alone (Extended Data Fig. 3a, b),

$$logit(\theta) \sim \beta_0 + \beta_1 v_i + \Sigma_c \beta_c P C_c + \varepsilon$$
⁽⁴⁾

then with C4 composite risk (Extended Data Fig. 3c),

$$logit(\theta) \sim \beta_0 + \beta_1 v_i + \beta_2 C 4 + \Sigma_c \beta_c P C_c + \varepsilon$$
⁽⁵⁾

where θ =E[SLE|X]. For SjS, the simpler weighted (2.3)*C*4*A*+*C*4*B* model was used instead of composite risk term, as the cohort's size gave poor precision to estimates of risk for many joint (*C*4*A*, *C*4*B*) copy numbers (Extended Data Fig. 3d). The Pearson correlation between the *C*4 composite risk term and each other variant was computed and squared (r²) to yield a measure of linkage disequilibrium between *C*4 composite risk and that variant in that cohort.

Association analyses for specific C4 structural alleles

The *C4* structural haplotypes were tested for association with disease (Fig. 1b, 2a) in a joint logistic regression that included (i) terms for dosages of the five most common *C4* structural haplotypes (AL-BS, AL-BL, AL-AL, BS, and AL), (ii) (for SLE and SjS) rs2105898 genotype, and (iii) ancestry covariates and (for SjS) smoking status,

$$logit(\theta) \sim \beta_0 + \beta_1 BS + \beta_2 AL + \beta_3 ALBS + \beta_4 ALBL + \beta_5 ALAL + \beta_6 rs2105898 + \Sigma_c \beta_c PC_c + \varepsilon$$
(6)

where θ =E[SLE|X]. Several of these common *C4* structural alleles arose multiple times on distinct haplotypes; we term the set of haplotypes in which such a common allele appeared as "haplogroups". The haplogroups can be further tested in a logistic regression model in which the structural allele appearing in all member haplotypes is instead encoded as dosages for each of the SNP haplotypes in which it appears. These association analyses (Fig. 1b, 2a) were performed as in (6), with structural allele dosages for ALBS, ALBL, and ALAL replaced by multiple terms for each distinct haplotype.

To delineate the relationship between *C4*-BS and *DRB1**03:01 alleles – which are highly linked in European ancestry haplotypes – allelic dosages per individual in the African American SLE cohort were rounded to yield the most likely integer dosage for each. Although genotype dosages for each are reported by BEAGLE and HIBAG respectively, probabilities per haplotype are not linked and multiplying possible diploid dosages could yield incorrect non-zero joint dosages. Joint genotypes were tested as individual terms in a logistic regression model (Fig. 2b),

$$logit(\theta) \sim \beta_0 + \Sigma_{i, j} \beta_{i, j} P(C4 - BS = i, DRB1 * 03:01 = j) + \Sigma_c \beta_c PC_c + \varepsilon$$
(7)

Sex-stratified associations of C4 structural alleles and other variants with SLE, SjS, and schizophrenia

Determination of an effect from sex on the contribution of overall C4 variation to risk for each disorder was done by including an interaction term between sex and C4; ie. (2.3)C4A + C4B for SLE and SjS and estimated C4A expression for schizophrenia:

$$logit(\theta) \sim \beta_0 + \beta_2 C 4 + \beta_3 I_{Sex} + \beta_4 I_{Sex} C 4 + \Sigma_c \beta_c P C_c + \varepsilon$$
(8)

Each variant in the MHC region was tested for association with among European ancestry cases and cohorts in a logistic regression as in models (4)–(6) using only male cases and controls, and then separately using only female cases and controls (Extended Data Fig. 6a–c). Likewise, allelic series analyses were performed as in (7), but in separate models for men and women (Fig. 3a, b).

To assess the relationship between sex bias in the risk associated with a variant and linkage to C4 composite risk (as non-negative r^2), male and female log-odds were multiplied by the sign of the Pearson correlation between that variant and C4 composite risk before taking the difference.

Analyses of cerebrospinal fluid

Cerebrospinal fluid (CSF) from healthy individuals was obtained from two research panels. The first panel, consisting of 533 donors (327 male, 126 female) from hospitals around Utrecht, Netherlands, was described previously^{49,50}. The donors were generally healthy research participants undergoing spinal anesthesia for minor elective surgery. The same donors were previously genotyped using the Illumina Omni SNP array. To estimate *C4* copy numbers, we used SNPs from the MHC region (chr6:24–34 Mb on hg19) as input for *C4* allele imputation with Beagle, as described above in Imputation of *C4* alleles.

The second CSF panel sampled specimens from 56 donors (14 male, 42 female) from Brigham and Women's Hospital (BWH; Boston, MA, USA) under a protocol approved by the institutional review board at BWH (IRB protocol ID no. 1999P010911) with informed consent. These samples were originally obtained to exclude the possibility of infection, and clinical analyses had revealed no evidence of infection. Donors ranged in age from 18 to 64 years old. Blood samples from the same individuals were used for extraction of genomic DNA, and *C4* gene copy number was measured by droplet digital PCR (ddPCR) as previously described⁷. Samples were excluded from measurements if they lacked *C4* genotypes, sex information, or contained visible blood contamination.

C4 measurements were performed by sandwich ELISA of 1:400 dilutions of the original CSF sample using goat anti-sera against human C4 as the capture antibody (Quidel, A305, used at 1:1000 dilution), FITC-conjugated polyclonal rabbit anti-human C4c as the detection antibody (Dako, F016902–2, used at 1:3000 dilution), and alkaline phosphatase–conjugated polyclonal goat anti-rabbit IgG as the secondary antibody (Abcam, ab97048, used at 1:5000 dilution). C3 measurements were performed using the human complement C3 ELISA kit (Abcam, ab108823).

Because *C4* gene copy number had a large and proportional effect on C4 protein concentration in these CSF samples (Extended Data Fig. 7a), we corrected for *C4* gene copy number in our analysis of relationship between sex and C4 protein concentration, by normalizing the ratio of C4 protein (in CSF) to *C4* gene copies (in genome). Therefore, these analyses included only samples for which DNA was available or *C4* was successfully imputed. In total, 495 (332 male, 163 female) C4 and 304 (179 male, 125 female) C3 concentrations were obtained across both cohorts. Log-concentrations of C3 (ng/mL) and C4

(ng/[mL, per *C4* gene copy number]) protein were then used separately in linear regression models to estimate a sex-unbiased cohort-specific offset for each protein,

$$\log_{10}(C3 \text{ or } C4 \text{ concentration}) \sim \beta_0 + \beta_1 I_{male} + \beta_2 I_{cohort} + \epsilon$$
 (9)

to be applied to all concentrations for that protein. Estimation of average measurements by age for each sex was done by local polynomial regression smoothing (LOESS) (Fig. 3c, d). To evaluate the significance of sex effects, we used these cohort-corrected concentrations estimates and analyzed them with the non-parametric unsigned Mann-Whitney rank–sum test comparing concentration distributions for males and females.

Analyses of blood plasma

Blood plasma was collected and immunoturbidimetric measurements of C3 and C4 protein in 1,844 individuals (182 men, 1662 women) by Sjögren's International Collaborative Clinical Alliance (SICCA) from individuals with and without SjS as previously described⁵¹. *C4* copy numbers for these individuals were previously imputed for use in logistic regression of SjS risk. As *C4* copy number has an effect on measured C4 protein similar to CSF (Extended Data Fig. 7b), we normalized C4 levels to them in all following analyses. Estimation of average measurements by age for each sex was done by local polynomial regression smoothing (LOESS) on log-concentrations of C3 (mg/dL) and C4 (mg/[dL, per *C4* gene copy number]) protein (Extended Data Fig. 7c, d). To evaluate the significance of sex bias within age ranges displaying the greatest difference (informed by LOESS), we analyzed individuals in these bins with the non-parametric unsigned Mann-Whitney rank– sum test comparing concentration distributions for males and females.

Difference in C4 protein levels between individual with and without SjS was done by performing a non-parametric unsigned Mann-Whitney rank–sum test on C4 protein levels with and without normalization to *C4* genomic copy number (Extended Data Fig. 7e, f).

Data Availability Statement

Individual genotype data for Sjögren's syndrome cases and controls and individual plasma concentrations for C4 and C3 are available in dbGaP under accession number phs000672.v1.p1. Individual genotype data for schizophrenia cases and controls are available by application to the Psychiatric Genomics Consortium (PGC). Questions regarding individual genotype data for SLE cases and controls of European and/or African American ancestry can be directed to Timothy J. Vyse (timothy.vyse@kcl.ac.uk). Data resources (reference haplotypes), software scripts and instructions for imputing C4 alleles into SNP data sets are available on the McCarroll lab web site at http://mccarrolllab.org/resources/resources-for-c4/. Genotype and protein concentration data for CSF samples are available upon request.

Extended Data





(a) Distributions (across 1,265 individuals) of total C4 gene copy number (C4A + C4B), as measured from read depth of coverage across the C4 locus, in whole-genome sequencing data.

(b) The relative numbers of reads that overlap sequences specific to C4A or C4B (together with the total C4 gene copy number as in **a**) are used to infer the underlying copy numbers of the C4A and C4B genes. For example, in an individual with four C4 genes, the presence

of equal numbers of reads specific to *C4A* or *C4B* suggests the presence of two copies each of *C4A* and *C4B*. Precise statistical approaches (including inference of probabilistic dosages), and further approaches for phasing *C4* allelic states with nearby SNPs to create reference haplotypes, are described in Methods.

(c) The SNP haplotypes flanking each C4 allele are shown as rows (SNPs as columns), with white and black representing the major and minor allele of each SNP. Gray lines at the bottom indicate the physical location of each SNP along chromosome 6. The differences among the haplotypes are most pronounced closest to C4 (toward the center of the plot), as historical recombination events in the flanking megabases will have caused the haplotypes to be less consistently distinct at greater genomic distances from C4. The patterns indicate that many combinations of C4A and C4B gene copy numbers have arisen recurrently on more than one SNP haplotype, a relationship that can be used in association analyses (Fig. 1b).

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Extended Data Figure 2. Aggregation of joint *C4A* and *C4B* genotype probabilities per individual across imputed *C4* structural alleles for estimation of SLE risk for each combination.
(a) An individual's joint *C4A* and *C4B* gene copy number can be calculated by summing the *C4A* and *C4B* gene contents for each possible pair of two inherited alleles. Many pairings of possible inherited alleles result in the same joint *C4A* and *C4B* gene copy number.
(b) Each individual's *C4A* and *C4B* gene copy number was imputed from their SNP data, using the reference haplotypes summarized in Extended Data Fig. 1c. For >95% of individuals (exemplified by samples 1–6 in the figure), this inference can be made with >90% certainty/confidence (the areas of the circles represent the posterior probability distribution over possible *C4A/C4B* gene copy numbers). For the remaining individuals (exemplified by samples 7–9 in the figure), greater statistical uncertainty persists about *C4* genotype. To account for this uncertainty, in downstream association analysis, all *C4*

genotype assignments are handled as probabilistic gene dosages – analogous to the genotype dosages that are routinely used in large-scale genetic association studies that use imputation. (c) Odds ratios and 95% confidence intervals underlying each of the *C4*-genotype risk estimates in Fig. 1a presented as a series of panels for each observed copy number of *C4B*, with increasing copy number of *C4A* for that *C4B* dosage (x-axis). Data are from analysis of 6,748 SLE cases and 11,516 controls of European ancestry.

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Extended Data Figure 3. Conditional association analyses for genetic markers across the extended MHC genomic region within the European-ancestry SLE and Sjögren's syndrome (SjS) cohort.

(a) Association of SLE with genetic markers (SNPs and imputed *HLA* alleles) across the extended MHC locus within the European-ancestry SLE cohort (6,748 cases and 11,516 controls). Orange diamond: an initial estimate of *C4*-related genetic risk, calculated as a weighted sum of the number of *C4A* and *C4B* gene copies: (2.3)C4A+C4B, with the weights derived from the relative coefficients estimated from logistic regression of SLE risk vs. *C4A* and *C4B* gene dosages. This risk score is imputed with an accuracy (r^2) of 0.77. Points representing all other genetic variants in the MHC locus are shaded orange according to their level of linkage disequilibrium–based correlation to this *C4*-derived risk score. (b) As in **a**, but for a European-ancestry Sjögren's syndrome (SjS) cohort (673 cases and 1,153 controls). The orange diamond here also represents (2.3)*C4A+C4B*, with this

weighting derived from the relative coefficients estimated from logistic regression of SjS risk vs. *C4A* and *C4B* gene dosages

(c) Association of SLE with genetic markers (SNPs and imputed *HLA* alleles) across the extended MHC locus within the European-ancestry SLE cohort controlling for *C4* composite risk (weighted sum of risk associated with various combinations of *C4A* and *C4B*). Variants are shaded in purple by their LD with rs2105898, an independent association identified from trans-ancestral analyses.

(d) As in **c**, but in association with a European-ancestry SjS cohort. Here a simpler linear model of risk contributed by *C4A* and *C4B* was used instead of a weighted sum across all possible combinations.



Extended Data Figure 4. Using *C4* **gene variation to understand the appearance of transancestral disparity in MHC association signals, and to fine-map an additional genetic effect** All panels show association signals (for SLE and SjS) for variants in a multi-megabase region of human chromosome 6 containing the MHC region including the *HLA* and *C4* genes.

(a) Relationship between SLE association [$-\log_{10}(p)$, y-axis] and LD to the weighted *C4* risk score (x-axis) for genetic markers and imputed *HLA* alleles across the extended MHC locus. In this European-ancestry cohort, it is unclear (from this analysis alone) whether the association with the markers in the predominant ray of points (at a ~45° angle from the x-

axis) is driven by variation at C4 or by the long haplotype containing DRB1*03:01 (green), DQA1*05:01 (blue), and B*08:01 (red). In addition, at least one independent association signal (a ray of points at a higher angle in the plot, with strong association signals and only weak LD-based correlation to C4 and DRB1*0301) with some LD to DRB1*15:01 (maroon) is also present.

(b) Analysis as in **a**, but for associations to SjS in a cohort of European ancestry. As in SLE, it is initially unclear whether the genetic association signal is driven by variation at *C4* or by linked *HLA* alleles, *DRB1**03:01 (green), *DQA1**05:01 (blue), and *B**08:01 (red). There is also an independent association signal with LD to *DRB1**15:01 (maroon). (c) Analysis as in **a**, but of an African American SLE case–control cohort (in which LD in the MHC region is more limited). Many MHC-region SNPs associate with SLE in proportion to their LD with the weighted *C4* risk score inferred from the earlier analysis of the European-ancestry cohort; this *C4*-derived risk score itself associates with SLE at $p = 4.3 \times 10^{-19}$ in a logistic regression on 1,494 SLE cases and 5,908 controls. No similarly strong association is observed for *DRB1**03:01, *DQA1**05:01, or *B**08:01, *HLA* alleles which are in strong LD with *C4* risk on European-ancestry (but not African American) haplotypes. An independent association signal is also present in this cohort, more clearly in LD with the *DRB1**15:03 allele (maroon).

(d) LD in the European-ancestry SLE cohort between the composite *C4* risk term (weighted sum of risk associated with various combinations of *C4A* and *C4B* from Fig. 2a) and variants in the MHC region as r^2 (y-axis).

(e) As in **d**, but for the African American SLE cohort.

(f) LD (to *C4* composite risk) for the same variants in European-ancestry individuals (x-axis) and African Americans (y-axis). Note the abundance of variants that have greater LD with *C4* risk among European-ancestry individuals than among African Americans. Also, several groups of variants have equivalent LD (to *C4* risk) in European ancestry individuals but exhibit a range of LD to *C4* risk among African Americans.

(g) Associations with SLE ($-\log_{10} p$ -values) for the same variants in European ancestry (x-axis) and African American (y-axis) case-control cohorts. Orange shading represents the extent of LD with *C4* risk in European ancestry individuals. Variants with strong European-specific association to SLE are generally in strong LD with *C4* risk among Europeans-ancestry individuals.

(h) Comparison of the inferred effect size from association of genetic markers with SLE (unconditioned log-odds ratios) among European-ancestry (x-axis) and African American (y-axis) research participants. As also seen in **g**, variants with discordant associations to SLE (across populations) tend also to be in strong LD to *C4* risk among European-ancestry individuals.

(i) As in **g**, but now controlling for the effect of C4 variation in analysis of the Europeanancestry cohort (x-axis). Note that controlling for C4 risk in European-ancestry individuals alone greatly aligns (relative to **g**) the patterns of association between European ancestry and African American cohorts.

(j) As in **i**, but now also controlling for the effect of C4 in associations of the African American cohort. Note that due to the lack of strong LD relationships between C4 and variants in the MHC region in African Americans (**e**), this further adjustment does not change results strongly (relative to **i**). The independent signal, rs2105898, and *HLA* alleles,

DRB1*15:01 and DRB1*15:03, are also highlighted. LD with rs2105898 in European-ancestry individuals is indicated by purple shading.
(k) Comparison of the inferred effect sizes from association of genetic markers with SLE (log-odds ratios) controlling for C4-derived risk among European-ancestry (x-axis) and African American (y-axis) research participants. Two SNPs (rs2105898 and rs9271513) that form a short haplotype common to both ancestry groups are among the strongest associations in both cohorts. (Their association to SLE in the European-ancestry cohort was initially much less remarkable than that of other SNPs that are in strong LD with C4.) LD with rs2105898 in European-ancestry individuals is indicated by purple shading.
(l) As in i, but with variants shaded by whether they exhibit greater LD to rs2105898 in

Europeans (blue) or African Americans (red).



b

ZNF143 consensus motif

rs2105898 reference allele

T G A A T A C A T T A A C C A G G G G G C

rs2105898 alternate allele

T G A C T A C A T T A A C C A G G G G G C

Extended Data Figure 5. Relationship of rs2105898 alleles to a known ZNF143 binding motif in the XL9 region of the MHC class II locus

(a) Location of rs2105898 (yellow line at center) within the XL9 region, with relevant tracks showing overlapping histone marks and transcription factor binding peaks (from ENCODE⁵²), visualized with the UCSC genome browser⁵³.

(b) ZNF143 consensus binding motif as a sequence logo, with the letters colored if the base is present in >5% of observed instances. The alleles of rs2105898 are indicated by outlined box surrounding the base.

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Extended Data Figure 6. Relationships between sex bias of disease associations and LD to C4 risk for variants in the MHC region.

(e) Relationship between male bias in SLE risk (difference between male and female log– odds ratios) and LD with *C4* risk for common (minor allele frequency [MAF] > 0.1) genetic markers across the extended MHC region. For each SNP, the allele for which sex risk bias is plotted is the allele that is positively correlated (via LD) with *C4*-derived risk score. (f) Relationship between male bias in SjS risk (log-odds ratios) and LD with *C4* risk for common (minor allele frequency [MAF] > 0.1) genetic markers across the extended MHC region. For each SNP, the allele for which sex risk bias is plotted is the allele that is positively correlated (via LD) with *C4*-derived risk score.

(g) Relationship of male bias in schizophrenia risk (log–odds ratios) and LD to *C4A* expression for common (MAF > 0.1) genetic markers across the extended MHC region. For each SNP, the allele for which sex risk bias is plotted is the allele that is positively correlated (via LD) with imputed *C4A* expression, as previously described⁷.



Extended Data Figure 7. Correlation of C4 protein measurements in cerebrospinal fluid and blood plasma with imputed C4 gene copy number and relationship of plasma complement to sex and SjS status

(a) Measurements of C4 protein in CSF obtained by ELISA are presented as $\log_{10}(ng/mL)$ (y-axis) for each observed or imputed copy number of total *C4* (x-axis, here showing most likely copy number from imputation). Because *C4* gene copy number affects C4 protein levels so strongly, we normalized C4 protein measurements to each donor's *C4* gene copy number in subsequent analyses (Fig. 3c). Bars indicate median values for each *C4* copy number.

(b) Measurements of C4 protein in blood plasma obtained by immunoturbidimetric assays are presented as $\log_{10}(mg/dL)$ (y-axis) for each imputed most-likely copy number of C4 genes (x-axis). Because C4 gene copy number affects C4 protein levels so strongly, we normalized C4 protein measurements by C4 gene copy number in subsequent analyses as in c. Due to the number of observations (n = 1,844 total), the plot is downsampled to 500 points; the median bars shown are for all individuals (before downsampling). (c) Levels of C4 protein in blood plasma from 182 adult men and 1662 adult women as a function of age. Concentrations are normalized to the number of C4 gene copies in an individual's genome (a strong independent source of variance) and shown on a \log_{10} scale as a LOESS curve. Shaded regions represent 95% confidence intervals derived during LOESS. (d) Levels of C3 protein in blood plasma as a function of age from the same individuals in panel c. Concentrations are shown on a log_{10} scale as a LOESS curve. Shaded regions represent 95% confidence intervals derived during LOESS. (e) C4 protein in blood plasma was measured in 670 individuals with SiS (red) and 1,151 individuals without SjS (black) and is shown on a log₁₀ scale (x-axis). Vertical stripes represent median levels for cases and controls separately. Comparison of the two sets was done with a non-parametric two-sided Mann-Whitney rank–sum test ($p = 4.8 \times 10^{-21}$). (f) As in \mathbf{e} , but concentrations are normalized to the number of C4 gene copies in an individual's genome and this per-copy amount is shown on a log_{10} scale (x-axis). Comparison of the two sets was done with a non-parametric two-sided Mann-Whitney rank-

sum test ($p = 7.6 \times 10^{-9}$).

Extended Data Table 1. Imputation accuracy for *C4* copy numbers in European ancestry and African American haplotypes.

Imputation accuracy was evaluated by correlation of imputation results to C4 gene copy numbers directly inferred from WGS data. Aggregated copy numbers imputed from each round of leaving 10 individuals out were correlated with the directly-typed measurements and are reported as r^2 for each feature of C4 structural variation for European ancestry and African American members of the reference panel separately.

	Imputation accuracy (r ²)									
Gene copy number	European ancestry	African Americans								
<i>C4</i>	0.80	0.58								
C4A	0.78	0.65								
C4B	0.74	0.61								
C4-HERV	0.91	0.76								
2.3(<i>C</i> 4 <i>A</i>)+ <i>C</i> 4 <i>B</i>	0.77	0.64								

Extended Data Table 2. Frequency of common *C4* alleles and their LD-based correlation with *HLA* alleles in European ancestry and African American cohorts.

For each common *C4* allele and *HLA* gene, the allele with strongest LD (r^2) is listed if present on more than half of the haplotypes with that *C4* allele (exact fraction in %). r^2 values greater than 0.4 are highlighted to point out particularly strong *C4-HLA* allele correlations, such as for several *HLA* alleles with the *C4*-B(S) allele in European ancestry individuals. Some common *C4* alleles are further subdivided into distinct haplotypes used in imputation (and in Fig. 1b), as defined by shared alleles from variants flanking *C4*. Note that some alleles such as *C4*-A(L)-A(L)-3 are present at a low frequency in African Americans that might reflect their presence on admixed European-origin haplotypes spanning this region, whereas others such as *C4*-B(S) are likely to also exist on African haplotypes – these differences between *C4* alleles are also reflected in the similarity of LD with *HLA* alleles to the corresponding row of the European ancestry section.

Europe	ean ai	ncestry																
	A		В		СС		C4 Allele	1	DRB1		1	DQA1		1	DQB1			
allele	%	r^2	allele	%	r ²	allele	%	r^2	allele	Frequency	allele	%	r^2	allele	%	r ²	allele	%
01:01	69	0.27	08:01	93	0.75	07:01	93	0.57	B(S)	13.7%	03:01	94	0.71	05:01	94	0.7	02:01	94
									A(L)	4.8%								
						06:02	69	0.31	A(L)- B(S)-1	6.1%	07:01	74	0.25	02:01	74	0.25		
			44:03	54	0.28	16:01	53	0.39	A(L)- B(S)-2	4.5%	07:01	57	0.1	02:01	57	0.1	02:02	55
									A(L)- B(S)-3	3.8%								
									A(L)- B(S)-4	4.5%								
			07:02	64	0.42	07:02	63	0.35	A(L)- B(L)-1	15.5%	15:01	73	0.49	01:02	74	0.32	06:02	70
									A(L)- B(L)-2	23.1%								
			35:01	55	0.2	04:01	57	0.09	A(L)- A(L)-1	3.2%	01:01	65	0.14	01:01	65	0.11	05:01	64
									A(L)- A(L)-2	2.1%	13:01	67	0.16	01:03	65	0.13	06:03	67
02:01	65	0.03	44:02	74	0.24	05:01	72	0.23	A(L)- A(L)-3	4.5%	04:01	80	0.29	03:03	79	0.37	03:01	82
Africa	n Am	erican																
	A			В			С		C4	مامالا		DRB1			DQA1			DQB1
allele	%	r ²	allele	%	r ²	allele	%	r ²	allele	Frequency	allele	%	r ²	allele	%	r ²	allele	%
									B(S)	5.0%				01:02	51	0.01		
									A(L)	7.5%								

									A(L)- B(S)-1	14.1%								
									A(L)- B(S)-2	18.1%								
									A(L)- B(S)-3	17.7%								
									A(L)- B(S)-4	6.5%								
									A(L)- B(L)-1	4.4%	15:01	67	0.2	01:02	72	0.04	06:02	59
									A(L)- B(L)-2	4.5%								
									A(L)- A(L)-1	0.7%	01:01	57	0.07	01:01	53	0.01		
									A(L)- A(L)-2	0.8%								
02:01	72	0.03	44:02	86	0.31	05:01	78	0.17	A(L)- A(L)-3	0.8%	04:01	93	0.27	03:03	86	0.14	03:01	87

Extended Data Table 3. Results of association analyses of SLE risk against *C4* variation, *HLA* alleles, and/or rs2105898 in European ancestry and African American cohorts.

Coefficients (beta, standard error) and p-values (as $-\log_{10}(p)$) for individual terms composing several relevant logistic regression models for predicting SLE risk in a European ancestry cohort of 6,748 SLE cases and 11,516 controls and an African American cohort of 1,494 SLE cases and 5,908 controls. Each analysis also included ancestry-specific covariates. For each model, the Akaike information criterion (AIC) and overall p-value (as determined by Chi-squared likelihood-ratio test) are given at the right to indicate the relative strengths of similar models for each ancestry cohort.

European and	European ancestry													
		C4			C4A			C4B			DRB1*03:01			B
Model	beta	se	-log10(p)	beta	se	-log10(p)	beta	se	-log10(p)	beta	se	-log10(p)	beta	
<i>C4</i>	-0.55	0.027	92.7											
C4A				-0.53	0.024	105.3								
C4A+C4B				-0.62	0.028	112	-0.27	0.037	12.3					
DRB1*03:01										0.7	0.03	117.1		
<i>B</i> *08:01													0.69	0.0
rs2105898														
<i>C4A</i> + <i>C4B</i> + <i>DRB1</i> *03:01				-0.35	0.041	17.2	-0.11	0.041	2.3	0.4	0.046	17.5		
<i>C4A</i> + <i>C4B</i> + <i>B</i> *08:01				-0.41	0.039	24.6	-0.17	0.039	4.7				0.35	0.0

C4A + C4B	-0.67	0.028	122.8	-0.32	0.038	16.4
+ rs2105898						

African Ame	African American													
	C4			C4A			C4B			DRB1*03:01				B
Model	beta	se	-log10(p)	beta	se	-log10(p)	beta	se	-log10(p)	beta	se	-log10(p)	beta	
<i>C4</i>	-0.51	0.059	17.3											
C4A				-0.43	0.062	11.2								
C4A+C4B				-0.62	0.068	18.7	-0.41	0.068	8.6					
DRB1*03:01										0.41	0.091	5.2		
<i>B</i> *08:01													0.78	0.1
rs2105898														
<i>C4A</i> + <i>C4B</i> + <i>DRB1</i> *03:01				-0.59	0.073	15	-0.38	0.071	7.1	0.1	0.099	0.5		
<i>C4A</i> + <i>C4B</i> + <i>B</i> *08:01				-0.51	0.073	11.7	-0.37	0.069	7.2				0.49	0.12
<i>C4A</i> + <i>C4B</i> + rs2105898				-0.52	0.07	13.2	-0.43	0.069	9.4					

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Schizophrenia Working Group of the Psychiatric Genomics Consortium

Stephan Ripke^{16,17}, Benjamin M. Neale^{16,17,18,19}, Aiden Corvin²⁰, James T. R. Walters²¹, Kai-How Farh¹⁶, Peter A. Holmans^{21,22}, Phil Lee^{16,17,19}, Brendan Bulik-Sullivan^{16,17}, David A. Collier^{23,24}, Hailiang Huang^{16,18}, Tune H. Pers^{18,25,26}, Ingrid Agartz^{27,28,29}, Esben Agerbo^{30,31,32}, Margot Albus³³, Madeline Alexander³⁴, Farooq Amin^{35,36}, Silviu A. Bacanu³⁷, Martin Begemann³⁸, Richard A Belliveau Jr¹⁷, Judit Bene^{39,40}, Sarah E. Bergen ^{17,41}, Elizabeth Bevilacqua¹⁷, Tim B Bigdeli ³⁷, Donald W. Black⁴², Richard Bruggeman⁴³, Nancy G. Buccola⁴⁴, Randy L. Buckner^{45,46,47}, William Byerley⁴⁸, Wiepke Cahn⁴⁹, Guiqing Cai^{50,51}, Murray J. Cairns^{54,135,185}, Dominique Campion⁵², Rita M. Cantor⁵³, Vaughan J. Carr^{54,55}, Noa Carrera²¹, Stanley V. Catts^{54,56}, Kimberly D. Chambert¹⁷, Raymond C. K. Chan⁵⁷, Ronald Y. L. Chen⁵⁸, Eric Y. H. Chen^{58,59}, Wei Cheng⁶⁰, Eric F. C. Cheung⁶¹, Siow Ann Chong⁶², C. Robert Cloninger⁶³, David Cohen⁶⁴, Nadine Cohen⁶⁵, Paul Cormican²⁰, Nick Craddock^{21,22}, Benedicto Crespo-Facorro²²⁵, James J. Crowley⁶⁶,

David Curtis^{67,68}, Michael Davidson⁶⁹, Kenneth L. Davis⁵¹, Franziska Degenhardt^{70,71}, Jurgen Del Favero⁷², Lynn E. DeLisi^{143,144}, Ditte Demontis^{32,73,74}, Dimitris Dikeos⁷⁵, Timothy Dinan⁷⁶, Srdjan Djurovic^{29,77}, Gary Donohoe^{20,78}, Elodie Drapeau⁵¹, Jubao Duan^{79,80}, Frank Dudbridge⁸¹, Naser Durmishi⁸², Peter Eichhammer⁸³, Johan Eriksson^{84,85,86}, Valentina Escott-Price²¹, Laurent Essioux⁸⁷, Ayman H. Fanous^{88,89,90,91}, Martilias S. Farrell⁶⁶, Josef Frank⁹², Lude Franke⁹³, Robert Freedman⁹⁴, Nelson B. Freimer⁹⁵, Marion Friedl⁹⁶, Joseph I. Friedman⁵¹, Menachem Fromer^{16,17,19,97}, Giulio Genovese¹⁷, Lyudmila Georgieva²¹, Elliot S. Gershon²²⁴, Ina Giegling^{96,98}, Paola Giusti-Rodríguez⁶⁶, Stephanie Godard⁹⁹, Jacqueline I. Goldstein^{16,18}, Vera Golimbet¹⁰⁰, Srihari Gopal¹⁰¹, Jacob Gratten¹⁰², Lieuwe de Haan¹⁰³, Marina Mitjans³⁸, Marian L. Hamshere²¹, Mark Hansen¹⁰⁴, Thomas Hansen^{32,105}, Vahram Haroutunian^{51,106,107}, Annette M. Hartmann⁹⁶, Frans A. Henskens^{54,108,109}, Stefan Herms^{70,71,110}, Joel N. Hirschhorn^{18,26,111}, Per Hoffmann^{70,71,110}, Andrea Hofman^{70,71}, Mads V. Hollegaard¹¹², David M. Hougaard¹¹², Masashi Ikeda¹¹³, Inge Joa¹¹⁴, Antonio Julià¹¹⁵, René S. Kahn⁴⁹, Luba Kalaydjieva^{116,117}, Sena Karachanak-Yankova¹¹⁸, Juha Karjalainen⁹³, David Kavanagh²¹, Matthew C. Keller¹¹⁹, Brian J. Kelly¹³⁵, James L. Kennedy^{120,121,122}, Andrey Khrunin¹²³, Yunjung Kim⁶⁶, Janis Klovins¹²⁴, James A. Knowles¹²⁵, Bettina Konte⁹⁶, Vaidutis Kucinskas¹²⁶, Zita Ausrele Kucinskiene¹²⁶, Hana Kuzelova- Ptackova¹²⁷, Anna K. Kähler⁴¹, Claudine Laurent^{34,128}, Jimmy Lee Chee Keong^{62,129}, S. Hong Lee¹⁰², Sophie E. Legge²¹, Bernard Lerer¹³⁰, Miaoxin Li^{58,59,131} Tao Li¹³², Kung-Yee Liang¹³³, Jeffrey Lieberman¹³⁴, Svetlana Limborska¹²³, Carmel M. Loughland^{54,135}, Jan Lubinski¹³⁶, Jouko Lönnqvist¹³⁷, Milan Macek Jr¹²⁷, Patrik K. E. Magnusson⁴¹, Brion S. Maher¹³⁸, Wolfgang Maier¹³⁹, Jacques Mallet¹⁴⁰, Sara Marsal¹¹⁵, Manuel Mattheisen^{32,73,74,141}, Morten Mattingsdal^{29,142}, Robert W. McCarley^{143,144}, Colm McDonald¹⁴⁵, Andrew M. McIntosh^{146,147}, Sandra Meier⁹², Carin J. Meijer¹⁰³, Bela Melegh^{39,40}, Ingrid Melle^{29,148}, Raquelle I. Mesholam-Gately^{143,149}, Andres Metspalu¹⁵⁰, Patricia T. Michie^{54,151}, Lili Milani¹⁵⁰, Vihra Milanova¹⁵², Younes Mokrab²³, Derek W. Morris^{20,78}, Ole Mors^{32,73,153}, Kieran C. Murphy¹⁵⁴, Robin M. Murray¹⁵⁵, Inez Myin-Germeys¹⁵⁶, Bertram Müller-Myhsok^{157,158,159}, Mari Nelis¹⁵⁰, Igor Nenadic¹⁶⁰, Deborah A. Nertney¹⁶¹, Gerald Nestadt¹⁶², Kristin K. Nicodemus¹⁶³, Liene Nikitina-Zake¹²⁴, Laura Nisenbaum¹⁶⁴, Annelie Nordin¹⁶⁵, Eadbhard O'Callaghan¹⁶⁶, Colm O'Dushlaine¹⁷, F. Anthony O'Neill¹⁶⁷, Sang-Yun Oh¹⁶⁸, Ann Olincy⁹⁴, Line Olsen^{32,105}, Jim Van Os^{156,169}, Psychosis Endophenotypes International Consortium¹⁷⁰, Christos Pantelis^{54,171}, George N. Papadimitriou⁷⁵, Agnes A. Steixner³⁸, Elena Parkhomenko⁵¹, Michele T. Pato¹²⁵, Tiina Paunio^{172,173}, Milica Pejovic-Milovancevic¹⁷⁴, Diana O. Perkins¹⁷⁵, Olli Pietiläinen^{173,176}, Jonathan Pimm⁶⁸, Andrew J. Pocklington²¹, John Powell¹⁵⁵, Alkes Price^{18,177}, Ann E. Pulver¹⁶², Shaun M. Purcell⁹⁷, Digby Quested¹⁷⁸, Henrik B. Rasmussen^{32,105}, Abraham Reichenberg⁵¹, Mark A. Reimers¹⁷⁹, Alexander L. Richards²¹, Joshua L. Roffman^{45,47}, Panos Roussos^{97,180}, Douglas M. Ruderfer^{21,97}, Veikko Salomaa⁸⁶, Alan R. Sanders^{79,80}, Ulrich Schall^{54,135}, Christian R. Schubert¹⁸¹, Thomas G. Schulze^{92,182}, Sibylle G. Schwab¹⁸³, Edward M. Scolnick¹⁷, Rodney J. Scott^{54,184,185}, Larry J. Seidman^{143,149}, Jianxin Shi¹⁸⁶, Engilbert Sigurdsson¹⁸⁷, Teimuraz Silagadze¹⁸⁸, Jeremy M. Silverman^{51,189}, Kang Sim⁶², Petr Slominsky¹²³, Jordan W. Smoller^{17,19}, Hon-Cheong So⁵⁸, Chris C. A. Spencer¹⁹⁰, Eli A. Stahl^{18,97}, Hreinn Stefansson¹⁹¹, Stacy Steinberg¹⁹¹, Elisabeth Stogmann¹⁹², Richard E. Straub¹⁹³, Eric Strengman^{194,49}, Jana Strohmaier⁹², T. Scott Stroup¹³⁴, Mythily

Subramaniam⁶², Jaana Suvisaari¹³⁷, Dragan M. Svrakic⁶³, Jin P. Szatkiewicz⁶⁶, Erik Söderman²⁷, Srinivas Thirumalai¹⁹⁵, Draga Toncheva¹¹⁸, Paul A. Tooney^{54,135,185}, Sarah Tosato¹⁹⁶, Juha Veijola^{197,198}, John Waddington¹⁹⁹, Dermot Walsh²⁰⁰, Dai Wang¹⁰¹, Qiang Wang¹³², Bradley T. Webb³⁷, Mark Weiser⁶⁹, Dieter B. Wildenauer²⁰¹, Nigel M. Williams²¹, Stephanie Williams⁶⁶, Stephanie H. Witt⁹², Aaron R. Wolen¹⁷⁹, Emily H. M. Wong⁵⁸, Brandon K. Wormley³⁷, Jing Qin Wu^{54,185}, Hualin Simon Xi²⁰², Clement C. Zai^{120,121}, Xuebin Zheng²⁰³, Fritz Zimprich¹⁹², Naomi R. Wray¹⁰², Kari Stefansson¹⁹¹, Peter M. Visscher¹⁰², Wellcome Trust Case-Control Consortium ²²⁰⁴, Rolf Adolfsson¹⁶⁵, Ole A. Andreassen^{29,148}, Douglas H. R. Blackwood¹⁴⁷, Elvira Bramon²⁰⁵, Joseph D. Buxbaum^{50,51,106,206}, Anders D. Børglum^{32,73,74,153}, Sven Cichon^{70,71,110,207}, Ariel Darvasi²⁰⁸, Enrico Domenici²⁰⁹, Hannelore Ehrenreich³⁸, Tõnu Esko^{18,26,111,150}, Pablo V. Gejman^{79,80}, Michael Gill²⁰, Hugh Gurling⁶⁸, Christina M. Hultman⁴¹, Nakao Iwata¹¹³, Assen V. Jablensky^{54,117,201,210}, Erik G. Jönsson^{27,29}, Kenneth S. Kendler²¹¹, George Kirov²¹, Jo Knight^{120,121,122}, Todd Lencz^{212,213,214}, Douglas F. Levinson³⁴, Oingqin S. Li¹⁰¹, Jianjun Liu^{203,215}, Anil K. Malhotra^{212,213,214}, Steven A. McCarroll^{17,111}, Andrew McQuillin⁶⁸, Jennifer L. Moran¹⁷, Preben B. Mortensen^{30,31,32}, Bryan J. Mowry^{102,216}, Markus M. Nöthen^{70,71}, Roel A. Ophoff^{53,95,49}, Michael J. Owen^{21,22}, Aarno Palotie^{17,19,176}, Carlos N. Pato¹²⁵, Tracey L. Petryshen^{17,143,217}, Danielle Posthuma^{218,219,220}, Marcella Rietschel⁹², Brien P. Riley²¹¹, Dan Rujescu^{96,98}, Pak C. Sham^{58,59,131} Pamela Sklar^{97,106,180}, David St Clair²²¹, Daniel R. Weinberger^{193,222}, Jens R. Wendland¹⁸¹, Thomas Werge^{32,105,223}, Mark J. Daly^{16,17,18}, Patrick F. Sullivan^{41,66,175} & Michael C. O'Donovan^{21,22}

¹⁶Analytic and Translational Genetics Unit, Massachusetts General Hospital, Boston, Massachusetts 02114, USA.

¹⁷Stanley Center for Psychiatric Research, Broad Institute of MIT and Harvard, Cambridge, Massachusetts 02142, USA.

¹⁸Medical and Population Genetics Program, Broad Institute of MIT and Harvard, Cambridge, Massachusetts 02142, USA.

¹⁹Psychiatric and Neurodevelopmental Genetics Unit, Massachusetts General Hospital, Boston, Massachusetts 02114, USA.

²⁰Neuropsychiatric Genetics Research Group, Department of Psychiatry, Trinity College Dublin, Dublin 8, Ireland.

²¹MRC Centre for Neuropsychiatric Genetics and Genomics, Institute of Psychological
 Medicine and Clinical Neurosciences, School of Medicine, Cardiff University, Cardiff, CF24
 4HQ, UK.

²²National Centre for Mental Health, Cardiff University, Cardiff, CF24 4HQ, UK.

²³Eli Lilly and Company Limited, Erl Wood Manor, Sunninghill Road, Windlesham, Surrey, GU20 6PH, UK.

²⁴Social, Genetic and Developmental Psychiatry Centre, Institute of Psychiatry, King's College London, London, SE5 8AF, UK.

²⁵Center for Biological Sequence Analysis, Department of Systems Biology, Technical University of Denmark, DK-2800, Denmark.

²⁶Division of Endocrinology and Center for Basic and Translational Obesity Research, Boston Children's Hospital, Boston, Massachusetts, 02115 USA.

²⁷Department of Clinical Neuroscience, Psychiatry Section, Karolinska Institutet, SE-17176 Stockholm, Sweden.

²⁸Department of Psychiatry, Diakonhjemmet Hospital, 0319 Oslo, Norway.

²⁹NORMENT, KG Jebsen Centre for Psychosis Research, Institute of Clinical Medicine, University of Oslo, 0424 Oslo, Norway.

³⁰Centre for Integrative Register-based Research, CIRRAU, Aarhus University, DK-8210 Aarhus, Denmark.

³¹National Centre for Register-based Research, Aarhus University, DK-8210 Aarhus, Denmark.

³²The Lundbeck Foundation Initiative for Integrative Psychiatric Research, iPSYCH, Denmark.

³³State Mental Hospital, 85540 Haar, Germany.

³⁴Department of Psychiatry and Behavioral Sciences, Stanford University, Stanford, California 94305, USA.

³⁵Department of Psychiatry and Behavioral Sciences, Atlanta Veterans Affairs Medical Center, Atlanta, Georgia 30033, USA.

³⁶Department of Psychiatry and Behavioral Sciences, Emory University, Atlanta Georgia 30322, USA.

³⁷Virginia Institute for Psychiatric and Behavioral Genetics, Department of Psychiatry, Virginia Commonwealth University, Richmond, Virginia 23298, USA.

³⁸Clinical Neuroscience, Max Planck Institute of Experimental Medicine, Göttingen 37075, Germany.

³⁹Department of Medical Genetics, University of Pécs, Pécs H-7624, Hungary.

⁴⁰Szentagothai Research Center, University of Pécs, Pécs H-7624, Hungary.

⁴¹Department of Medical Epidemiology and Biostatistics, Karolinska Institutet, Stockholm SE-17177, Sweden.

⁴²Department of Psychiatry, University of Iowa Carver College of Medicine, Iowa City, Iowa 52242, USA.

⁴³University Medical Center Groningen, Department of Psychiatry, University of Groningen NL-9700 RB, The Netherlands.

⁴⁴School of Nursing, Louisiana State University Health Sciences Center, New Orleans, Louisiana 70112, USA.

⁴⁵Athinoula A. Martinos Center, Massachusetts General Hospital, Boston, Massachusetts 02129, USA.

⁴⁶Center for Brain Science, Harvard University, Cambridge, Massachusetts, 02138 USA.

⁴⁷Department of Psychiatry, Massachusetts General Hospital, Boston, Massachusetts, 02114 USA.

⁴⁸Department of Psychiatry, University of California at San Francisco, San Francisco, California, 94143 USA.

⁴⁹University Medical Center Utrecht, Department of Psychiatry, Rudolf Magnus Institute of Neuroscience, 3584 Utrecht, The Netherlands.

⁵⁰Department of Human Genetics, Icahn School of Medicine at Mount Sinai, New York, New York 10029 USA.

⁵¹Department of Psychiatry, Icahn School of Medicine at Mount Sinai, New York, New York 10029 USA.

⁵²Centre Hospitalier du Rouvray and INSERM U1079 Faculty of Medicine, 76301 Rouen, France.

⁵³Department of Human Genetics, David Geffen School of Medicine, University of California, Los Angeles, California 90095, USA.

⁵⁴Schizophrenia Research Institute, Sydney NSW 2010, Australia.

⁵⁵School of Psychiatry, University of New South Wales, Sydney NSW 2031, Australia.

⁵⁶Royal Brisbane and Women's Hospital, University of Queensland, Brisbane, St Lucia QLD 4072, Australia.

⁵⁷Institute of Psychology, Chinese Academy of Science, Beijing 100101, China.

⁵⁸Department of Psychiatry, Li Ka Shing Faculty of Medicine, The University of Hong Kong, Hong Kong, China.

⁵⁹State Key Laboratory for Brain and Cognitive Sciences, Li Ka Shing Faculty of Medicine, The University of Hong Kong, Hong Kong, China.

⁶⁰Department of Computer Science, University of North Carolina, Chapel Hill, North Carolina 27514, USA.

⁶¹Castle Peak Hospital, Hong Kong, China.

⁶²Institute of Mental Health, Singapore 539747, Singapore.

⁶³Department of Psychiatry, Washington University, St. Louis, Missouri 63110, USA.

⁶⁴Department of Child and Adolescent Psychiatry, Assistance Publique Hopitaux de Paris, Pierre and Marie Curie Faculty of Medicine and Institute for Intelligent Systems and Robotics, Paris, 75013, France.

⁶⁵ Blue Note Biosciences, Princeton, New Jersey 08540, USA

⁶⁶Department of Genetics, University of North Carolina, Chapel Hill, North Carolina 27599–7264, USA.

⁶⁷Department of Psychological Medicine, Queen Mary University of London, London E11BB, UK.

⁶⁸Molecular Psychiatry Laboratory, Division of Psychiatry, University College London, London WC1E 6JJ, UK.

⁶⁹Sheba Medical Center, Tel Hashomer 52621, Israel.

⁷⁰Department of Genomics, Life and Brain Center, D-53127 Bonn, Germany.

⁷¹Institute of Human Genetics, University of Bonn, D-53127 Bonn, Germany.

⁷²Applied Molecular Genomics Unit, VIB Department of Molecular Genetics, University of Antwerp, B-2610 Antwerp, Belgium.

⁷³Centre for Integrative Sequencing, iSEQ, Aarhus University, DK-8000 Aarhus C, Denmark.

⁷⁴Department of Biomedicine, Aarhus University, DK-8000 Aarhus C, Denmark.

⁷⁵First Department of Psychiatry, University of Athens Medical School, Athens 11528, Greece.

⁷⁶Department of Psychiatry, University College Cork, Co. Cork, Ireland.

⁷⁷Department of Medical Genetics, Oslo University Hospital, 0424 Oslo, Norway.

⁷⁸Cognitive Genetics and Therapy Group, School of Psychology and Discipline of Biochemistry, National University of Ireland Galway, Co. Galway, Ireland.

⁷⁹Department of Psychiatry and Behavioral Neuroscience, University of Chicago, Chicago, Illinois 60637, USA.

⁸⁰Department of Psychiatry and Behavioral Sciences, NorthShore University HealthSystem, Evanston, Illinois 60201, USA.

⁸¹Department of Non-Communicable Disease Epidemiology, London School of Hygiene and Tropical Medicine, London WC1E 7HT, UK.

⁸²Department of Child and Adolescent Psychiatry, University Clinic of Psychiatry, Skopje 1000, Republic of Macedonia.

⁸³Department of Psychiatry, University of Regensburg, 93053 Regensburg, Germany.

⁸⁴Department of General Practice, Helsinki University Central Hospital, University of Helsinki P.O. Box 20, Tukholmankatu 8 B, FI-00014, Helsinki, Finland

⁸⁵Folkhälsan Research Center, Helsinki, Finland, Biomedicum Helsinki 1, Haartmaninkatu 8, FI-00290, Helsinki, Finland.

⁸⁶National Institute for Health and Welfare, P.O. BOX 30, FI-00271 Helsinki, Finland.

⁸⁷Translational Technologies and Bioinformatics, Pharma Research and Early Development,F. Hoffman-La Roche, CH-4070 Basel, Switzerland.

⁸⁸Department of Psychiatry, Georgetown University School of Medicine, Washington DC 20057, USA.

⁸⁹Department of Psychiatry, Keck School of Medicine of the University of Southern California, Los Angeles, California 90033, USA.

⁹⁰Department of Psychiatry, Virginia Commonwealth University School of Medicine, Richmond, Virginia 23298, USA.

⁹¹Mental Health Service Line, Washington VA Medical Center, Washington DC 20422, USA.

⁹²Department of Genetic Epidemiology in Psychiatry, Central Institute of Mental Health, Medical Faculty Mannheim, University of Heidelberg, Heidelberg, D-68159 Mannheim, Germany.

⁹³Department of Genetics, University of Groningen, University Medical Centre Groningen,9700 RB Groningen, The Netherlands.

⁹⁴Department of Psychiatry, University of Colorado Denver, Aurora, Colorado 80045, USA.

⁹⁵Center for Neurobehavioral Genetics, Semel Institute for Neuroscience and Human Behavior, University of California, Los Angeles, California 90095, USA.

⁹⁶Department of Psychiatry, University of Halle, 06112 Halle, Germany.

⁹⁷Division of Psychiatric Genomics, Department of Psychiatry, Icahn School of Medicine at Mount Sinai, New York, New York 10029, USA.

⁹⁸Department of Psychiatry, University of Munich, 80336, Munich, Germany.

⁹⁹Departments of Psychiatry and Human and Molecular Genetics, INSERM, Institut de Myologie, Hôpital de la Pitiè-Salpêtrière, Paris, 75013, France.

¹⁰⁰Mental Health Research Centre, Russian Academy of Medical Sciences, 115522 Moscow, Russia.

¹⁰¹Neuroscience Therapeutic Area, Janssen Research and Development, Raritan, New Jersey 08869, USA.

¹⁰²Queensland Brain Institute, The University of Queensland, Brisbane, Queensland, QLD 4072, Australia.

¹⁰³Academic Medical Centre University of Amsterdam, Department of Psychiatry, 1105 AZ Amsterdam, The Netherlands.

¹⁰⁴Illumina, La Jolla, California, California 92122, USA.

¹⁰⁵Institute of Biological Psychiatry, Mental Health Centre Sct. Hans, Mental Health Services Copenhagen, DK-4000, Denmark.

¹⁰⁶Friedman Brain Institute, Icahn School of Medicine at Mount Sinai, New York, New York 10029, USA.

¹⁰⁷J. J. Peters VA Medical Center, Bronx, New York, New York 10468, USA.

¹⁰⁸Priority Research Centre for Health Behaviour, University of Newcastle, Newcastle NSW 2308, Australia.

¹⁰⁹School of Electrical Engineering and Computer Science, University of Newcastle, Newcastle NSW 2308, Australia.

¹¹⁰Division of Medical Genetics, Department of Biomedicine, University of Basel, Basel, CH-4058, Switzerland.

¹¹¹Department of Genetics, Harvard Medical School, Boston, Massachusetts 02115, USA.

¹¹²Section of Neonatal Screening and Hormones, Department of Clinical Biochemistry, Immunology and Genetics, Statens Serum Institut, Copenhagen, DK-2300, Denmark.

¹¹³Department of Psychiatry, Fujita Health University School of Medicine, Toyoake, Aichi, 470–1192, Japan.

¹¹⁴Regional Centre for Clinical Research in Psychosis, Department of Psychiatry, Stavanger University Hospital, 4011 Stavanger, Norway.

¹¹⁵Rheumatology Research Group, Vall d'Hebron Research Institute, Barcelona, 08035, Spain.

¹¹⁶Centre for Medical Research, The University of Western Australia, Perth, WA 6009, Australia.

¹¹⁷The Perkins Institute for Medical Research, The University of Western Australia, Perth, WA 6009, Australia.

¹¹⁸Department of Medical Genetics, Medical University, Sofia1431, Bulgaria.

¹¹⁹Department of Psychology, University of Colorado Boulder, Boulder, Colorado 80309, USA.

¹²⁰Campbell Family Mental Health Research Institute, Centre for Addiction and Mental Health, Toronto, Ontario, M5T 1R8, Canada.

¹²¹Department of Psychiatry, University of Toronto, Toronto, Ontario, M5T 1R8, Canada.

¹²²Institute of Medical Science, University of Toronto, Toronto, Ontario, M5S 1A8, Canada.

¹²³Institute of Molecular Genetics, Russian Academy of Sciences, Moscow123182, Russia.

¹²⁴Latvian Biomedical Research and Study Centre, Riga, LV-1067, Latvia.

¹²⁵Department of Psychiatry and Zilkha Neurogenetics Institute, Keck School of Medicine at University of Southern California, Los Angeles, California 90089, USA.

¹²⁶Faculty of Medicine, Vilnius University, LT-01513 Vilnius, Lithuania.

¹²⁷ Department of Biology and Medical Genetics, 2nd Faculty of Medicine and University Hospital Motol, 150 06 Prague, Czech Republic.

¹²⁸ Department of Child and Adolescent Psychiatry, Pierre and Marie Curie Faculty of Medicine, Paris 75013, France.

¹²⁹Duke-NUS Graduate Medical School, Singapore 169857, Singapore.

¹³⁰Department of Psychiatry, Hadassah-Hebrew University Medical Center, Jerusalem 91120, Israel.

¹³¹Centre for Genomic Sciences, The University of Hong Kong, Hong Kong, China.

¹³²Mental Health Centre and Psychiatric Laboratory, West China Hospital, Sichuan University, Chengdu, 610041, Sichuan, China.

¹³³Department of Biostatistics, Johns Hopkins University Bloomberg School of Public Health, Baltimore, Maryland 21205, USA.

¹³⁴Department of Psychiatry, Columbia University, New York, New York 10032, USA.

¹³⁵Priority Centre for Translational Neuroscience and Mental Health, University of Newcastle, Newcastle NSW 2300, Australia.

¹³⁶Department of Genetics and Pathology, International Hereditary Cancer Center, Pomeranian Medical University in Szczecin, 70–453 Szczecin, Poland.

¹³⁷Department of Mental Health and Substance Abuse Services; National Institute for Health and Welfare, P.O. BOX 30, FI-00271 Helsinki, Finland

¹³⁸Department of Mental Health, Bloomberg School of Public Health, Johns Hopkins University, Baltimore, Maryland 21205, USA.

¹³⁹Department of Psychiatry, University of Bonn, D-53127 Bonn, Germany.

¹⁴⁰Centre National de la Recherche Scientifique, Laboratoire de Génétique Moléculaire de la Neurotransmission et des Processus Neurodégénératifs, Hôpital de la Pitié Salpêtrière, 75013, Paris, France.

¹⁴¹Department of Genomics Mathematics, University of Bonn, D-53127 Bonn, Germany.

¹⁴²Research Unit, Sørlandet Hospital, 4604 Kristiansand, Norway.

¹⁴³Department of Psychiatry, Harvard Medical School, Boston, Massachusetts 02115, USA.

¹⁴⁴VA Boston Health Care System, Brockton, Massachusetts 02301, USA.

¹⁴⁵Department of Psychiatry, National University of Ireland Galway, Co. Galway, Ireland.

¹⁴⁶Centre for Cognitive Ageing and Cognitive Epidemiology, University of Edinburgh, Edinburgh EH16 4SB, UK.

¹⁴⁷Division of Psychiatry, University of Edinburgh, Edinburgh EH16 4SB, UK.

¹⁴⁸Division of Mental Health and Addiction, Oslo University Hospital, 0424 Oslo, Norway.

¹⁴⁹Massachusetts Mental Health Center Public Psychiatry Division of the Beth Israel Deaconess Medical Center, Boston, Massachusetts 02114, USA.

¹⁵⁰Estonian Genome Center, University of Tartu, Tartu 50090, Estonia.

¹⁵¹School of Psychology, University of Newcastle, Newcastle NSW 2308, Australia.

¹⁵²First Psychiatric Clinic, Medical University, Sofia 1431, Bulgaria.

¹⁵³Department P, Aarhus University Hospital, DK-8240 Risskov, Denmark.

¹⁵⁴Department of Psychiatry, Royal College of Surgeons in Ireland, Dublin 2, Ireland.

¹⁵⁵King's College London, London SE5 8AF, UK.

¹⁵⁶Maastricht University Medical Centre, South Limburg Mental Health Research and Teaching Network, EURON, 6229 HX Maastricht, The Netherlands.

¹⁵⁷Institute of Translational Medicine, University of Liverpool, Liverpool L69 3BX, UK.

¹⁵⁸Max Planck Institute of Psychiatry, 80336 Munich, Germany.

¹⁵⁹Munich Cluster for Systems Neurology (SyNergy), 80336 Munich, Germany.

¹⁶⁰Department of Psychiatry and Psychotherapy, Jena University Hospital, 07743 Jena, Germany.

¹⁶¹Department of Psychiatry, Queensland Brain Institute and Queensland Centre for Mental Health Research, University of Queensland, Brisbane, Queensland, St Lucia QLD 4072, Australia.

¹⁶²Department of Psychiatry and Behavioral Sciences, Johns Hopkins University School of Medicine, Baltimore, Maryland 21205, USA.

¹⁶³Department of Psychiatry, Trinity College Dublin, Dublin 2, Ireland.

¹⁶⁴Eli Lilly and Company, Lilly Corporate Center, Indianapolis, 46285 Indiana, USA.

¹⁶⁵Department of Clinical Sciences, Psychiatry, Umeå University, SE-901 87 Umeå, Sweden.

¹⁶⁶DETECT Early Intervention Service for Psychosis, Blackrock, Co. Dublin, Ireland.

¹⁶⁷Centre for Public Health, Institute of Clinical Sciences, Queen's University Belfast, Belfast BT12 6AB, UK.

¹⁶⁸Lawrence Berkeley National Laboratory, University of California at Berkeley, Berkeley, California 94720, USA.

¹⁶⁹Institute of Psychiatry, King's College London, London SE5 8AF, UK.

¹⁷⁰A list of authors and affiliations appear in the Supplementary Information.

¹⁷¹Melbourne Neuropsychiatry Centre, University of Melbourne & Melbourne Health, Melbourne, Vic 3053, Australia.

¹⁷²Department of Psychiatry, University of Helsinki, P.O. Box 590, FI-00029 HUS, Helsinki, Finland.

¹⁷³Public Health Genomics Unit, National Institute for Health and Welfare, P.O. BOX 30, FI-00271 Helsinki, Finland.

¹⁷⁴Medical Faculty, University of Belgrade, 11000 Belgrade, Serbia.

¹⁷⁵Department of Psychiatry, University of North Carolina, Chapel Hill, North Carolina 27599–7160, USA.

¹⁷⁶Institute for Molecular Medicine Finland, FIMM, University of Helsinki, P.O. Box 20 FI-00014, Helsinki, Finland.

¹⁷⁷Department of Epidemiology, Harvard School of Public Health, Boston, Massachusetts 02115, USA.

¹⁷⁸Department of Psychiatry, University of Oxford, OXford, OX3 7JX, UK.

¹⁷⁹Virginia Institute for Psychiatric and Behavioral Genetics, Virginia Commonwealth University, Richmond, Virginia 23298, USA.

¹⁸⁰Institute for Multiscale Biology, Icahn School of Medicine at Mount Sinai, New York, New York 10029, USA.

¹⁸¹PharmaTherapeutics Clinical Research, Pfizer Worldwide Research and Development, Cambridge, Massachusetts 02139, USA.

¹⁸²Department of Psychiatry and Psychotherapy, University of Gottingen, 37073 Göttingen, Germany.

¹⁸³Psychiatry and Psychotherapy Clinic, University of Erlangen, 91054 Erlangen, Germany.

¹⁸⁴Hunter New England Health Service, Newcastle NSW 2308, Australia.

¹⁸⁵School of Biomedical Sciences and Pharmacy, University of Newcastle, Callaghan NSW 2308, Australia.

¹⁸⁶Division of Cancer Epidemiology and Genetics, National Cancer Institute, Bethesda, Maryland 20892, USA.

¹⁸⁷University of Iceland, Landspitali, National University Hospital, 101 Reykjavik, Iceland.

¹⁸⁸Department of Psychiatry and Drug Addiction, Tbilisi State Medical University (TSMU), N33, 0177 Tbilisi, Georgia.

¹⁸⁹Research and Development, Bronx Veterans Affairs Medical Center, New York, New York 10468, USA.

¹⁹⁰Wellcome Trust Centre for Human Genetics, Oxford, OX3 7BN, UK.

¹⁹¹deCODE Genetics, 101 Reykjavik, Iceland.

¹⁹²Department of Clinical Neurology, Medical University of Vienna, 1090 Wien, Austria.

¹⁹³Lieber Institute for Brain Development, Baltimore, Maryland 21205, USA.

¹⁹⁴Department of Medical Genetics, University Medical Centre Utrecht, Universiteitsweg 100, 3584 CG, Utrecht, The Netherlands.

¹⁹⁵Berkshire Healthcare NHS Foundation Trust, Bracknell RG12 1BQ, UK.

¹⁹⁶Section of Psychiatry, University of Verona, 37134 Verona, Italy.

¹⁹⁷Department of Psychiatry, University of Oulu, P.O. BOX 5000, 90014, Finland

¹⁹⁸University Hospital of Oulu, P.O.BOX 20, 90029 OYS, Finland.

¹⁹⁹Molecular and Cellular Therapeutics, Royal College of Surgeons in Ireland, Dublin 2, Ireland.

²⁰⁰Health Research Board, Dublin 2, Ireland.

²⁰¹School of Psychiatry and Clinical Neurosciences, The University of Western Australia, Perth WA6009, Australia.

²⁰²Computational Sciences CoE, Pfizer Worldwide Research and Development, Cambridge, Massachusetts 02139, USA.

²⁰³Human Genetics, Genome Institute of Singapore, A*STAR, Singapore 138672, Singapore.

²⁰⁵University College London, London WC1E 6BT, UK.

²⁰⁶Department of Neuroscience, Icahn School of Medicine at Mount Sinai, New York, New York 10029, USA.

²⁰⁷Institute of Neuroscience and Medicine (INM-1), Research Center Juelich, 52428 Juelich, Germany.

²⁰⁸Department of Genetics, The Hebrew University of Jerusalem, 91905 Jerusalem, Israel.

²⁰⁹Neuroscience Discovery and Translational Area, Pharma Research and Early Development, F. Hoffman-La Roche, CH-4070 Basel, Switzerland.

²¹⁰Centre for Clinical Research in Neuropsychiatry, School of Psychiatry and Clinical Neurosciences, The University of Western Australia, Medical Research Foundation Building, Perth WA 6000, Australia.

²¹¹Virginia Institute for Psychiatric and Behavioral Genetics, Departments of Psychiatry and Human and Molecular Genetics, Virginia Commonwealth University, Richmond, Virginia 23298, USA.

²¹²The Feinstein Institute for Medical Research, Manhasset, New York, 11030 USA.

²¹³The Hofstra NS-LIJ School of Medicine, Hempstead, New York, 11549 USA.

²¹⁴The Zucker Hillside Hospital, Glen Oaks, New York,11004 USA.

²¹⁵Saw Swee Hock School of Public Health, National University of Singapore, Singapore 117597, Singapore.

²¹⁶Queensland Centre for Mental Health Research, University of Queensland, Brisbane 4076, Queensland, Australia.

²¹⁷Center for Human Genetic Research and Department of Psychiatry, Massachusetts General Hospital, Boston, Massachusetts 02114, USA.

²¹⁸Department of Child and Adolescent Psychiatry, Erasmus University Medical Centre, Rotterdam 3000, The Netherlands.

²¹⁹Department of Complex Trait Genetics, Neuroscience Campus Amsterdam, VU University Medical Center Amsterdam, Amsterdam 1081, The Netherlands.

²²⁰Department of Functional Genomics, Center for Neurogenomics and Cognitive Research, Neuroscience Campus Amsterdam, VU University, Amsterdam 1081, The Netherlands.

²²¹University of Aberdeen, Institute of Medical Sciences, Aberdeen, AB25 2ZD, UK.

²²²Departments of Psychiatry, Neurology, Neuroscience and Institute of Genetic Medicine, Johns Hopkins School of Medicine, Baltimore, Maryland 21205, USA.

²²³Department of Clinical Medicine, University of Copenhagen, Copenhagen 2200, Denmark.

²²⁴Departments of Psychiatry and Human Genetics, University of Chicago, Chicago, Illinois 60637, USA.

²²⁵University Hospital Marqués de Valdecilla, Instituto de Formación e Investigación Marqués de Valdecilla, University of Cantabria, ED39008 Santander, Spain.

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Figure 1. Association of SLE and Sjögren's syndrome (SjS) with C4 alleles

(a) Levels of SLE risk associated with 11 common combinations of *C4A* and *C4B* gene copy number. The color of each circle reflects the level of SLE risk (odds ratio) associated with a specific combination of *C4A* and *C4B* gene copy numbers relative to the most common combination (two copies of *C4A* and two copies of *C4B*) in gray. The area of each circle is proportional to the number of individuals with that number of *C4A* and *C4B* genes. Paths from left to right on the plot reflect the effect of increasing *C4A* gene copy number (greatly reduced risk); paths from bottom to top reflect the effect of increasing *C4B* gene copy number (modestly reduced risk); and diagonal paths from upper left to lower right reflect the effect of exchanging *C4B* for *C4A* copies (modestly reduced risk). Data are from analysis of 6,748 SLE cases and 11,516 controls of European ancestry. The odds ratios are reported with confidence intervals in Extended Data Fig. 2c.

(b) SLE and SjS risk associated with common combinations of *C4* structural allele and MHC SNP haplotype. For each *C4* locus structure, separate odds ratios are reported for each "haplogroup," i.e., the MHC SNP haplotype background on which the *C4* structure segregates. Data are from analyses of 6,748 SLE cases and 11,516 controls for the left plot and 673 SjS cases and 1,153 controls for the right plot. Error bars represent 95% confidence intervals around the effect size estimate for each allele.

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а

Odds ratio in African Americans 2.4 2.0 B(S) 1.5 1.2 A(L)1.0 A(L)-B(S)A(L) - B(L) $\mathbf{A}(L) - \mathbf{A}(L)$ 1.0 3.0 1.2 1.5 2.0 2.4 Odds ratio in Europeans b C4-B(S) alleles 2 0 0 2 0 2 1 1 16 8 4 Odds ratio 2 1 0.5 0.25 0.125 0.0625 ò 2 1 DRB1*03:01 alleles

> Figure 2. *C4* and trans-ancestral analysis of the MHC association signal in SLE (a) Common *C4* alleles exhibit similar strengths of association (odds ratios) in Europeanancestry and African American (1,494 SLE cases; 5,908 controls) cohorts. Error bars represent 95% confidence intervals around the effect size estimate for each sex. (b) Analysis of SLE risk across combinations of *C4*-B(S) and *DRB1**03:01 genotypes in an African American SLE case–control cohort, in which the two alleles exhibit very little LD ($r^2 = 0.10$). On each *DRB1**03:01 genotype background, additional *C4*-B(S) alleles increase risk (i.e. within each grouping). Whereas on each *C4*-B(S) background, *DRB1**03:01 alleles

have no appreciable relationship with risk (this can be seen by comparing, for example, the first of the three points from each group). Error bars represent 95% confidence intervals around the effect size estimate for each combination of C4-B(S) and DRBI*03:01.

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Figure 3. Sex differences in the magnitude of *C4* genetic effects and complement protein concentrations.

(a) SLE risk (odds ratios) associated with the four most common *C4* alleles in men (x-axis) and women (y-axis) among 6,748 affected and 11,516 unaffected individuals of European ancestry. For each sex, the lowest-risk allele (*C4*-A(L)-A(L)) is used as a reference (odds ratio of 1.0). Shading of each point reflects the relative level of SLE risk (darker = greater risk) conferred by *C4A* and *C4B* copy numbers as in Fig. 2b. Error bars represent 95% confidence intervals around the effect size estimate for each sex.

(b) Schizophrenia risk (odds ratios) associated with the four most common C4 alleles in men (x-axis) and women (y-axis) among 28,799 affected and 35,986 unaffected individuals of European ancestry, aggregated by the Psychiatric Genomics Consortium⁴³. For each sex, the lowest-risk allele (C4-B(S)) is used as a reference (odds ratio of 1.0). For visual comparison with **a**, shading of each allele reflects the relative level of SLE risk. Error bars represent 95% confidence intervals around the effect size estimate for each sex.

(c) Concentrations of C4 protein in cerebrospinal fluid sampled from 340 adult men (blue) and 167 adult women (pink) as a function of age with local polynomial regression (LOESS) smoothing. Concentrations are normalized to the number of *C4* gene copies in an individual's genome (a strong independent source of variance, Extended Data Fig. 7a) and shown on a \log_{10} scale as a LOESS curve. Shaded regions represent 95% confidence intervals derived during LOESS.

(d) Levels of C3 protein in cerebrospinal fluid from 179 adult men and 125 adult women as a function of age. Concentrations are shown on a \log_{10} scale as a LOESS curve. Shaded regions represent 95% confidence intervals derived during LOESS.