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Genome-wide association study of CNVs in 16,000 cases of eight common diseases and 3,000 shared controls

The Wellcome Trust Case Control Consortium*

Copy number variants (CNVs) account for a major proportion of human genetic polymorphism and have been predicted to have an important role in genetic susceptibility to common disease. To address this we undertook a large, direct genome-wide study of association between CNVs and eight common human diseases. Using a purpose-designed array we typed ~19,000 individuals into distinct copy-number classes at 3,432 polymorphic CNVs, including an estimated ~50% of all common CNVs larger than 500 base pairs. We identified several biological artefacts that lead to false-positive associations, including systematic CNV differences between DNAs derived from blood and cell lines. Association testing and follow-up replication analyses confirmed three loci where CNVs were associated with disease—*IRGM* for Crohn's disease, HLA for Crohn's disease, rheumatoid arthritis and type 1 diabetes, and *TSPAN8* for type 2 diabetes—although in each case the locus had previously been identified in single nucleotide polymorphism (SNP)-based studies, reflecting our observation that most common CNVs that are well-typed on our array are well tagged by SNPs and so have been indirectly explored through SNP studies. We conclude that common CNVs that can be typed on existing platforms are unlikely to contribute greatly to the genetic basis of common human diseases.

Genome-wide association studies (GWAS) have been extremely successful in associating SNPs with susceptibility to common diseases, but published SNP associations account for only a fraction of the genetic component of most common diseases, and there has been considerable speculation about where the 'missing heritability' might lie. Chromosomal rearrangements can cause particular rare diseases and syndromes², and recent reports have suggested a role for rare CNVs, either individually or in aggregate, in susceptibility for a range of common diseases, notably neurodevelopmental diseases³⁻⁶. So far, there have been relatively few reported associations between common diseases and common CNVs (see for example refs 7–11), which might simply reflect incomplete catalogues of common CNVs or the lack of reliable assays for their large-scale typing. Here we report the results of our direct association study, identify the population properties of the set of CNVs studied, describe novel analytical methods to facilitate robust analyses of CNV data, and document artefacts that can afflict CNV studies.

We designed an array to measure copy number for the majority of a recently compiled inventory of CNVs from an extensive discovery experiment¹², and several other sources. We then used the array to type 3,000 common controls and 2,000 cases of each of the diseases: bipolar disorder, breast cancer, coronary artery disease, Crohn's disease, hypertension, rheumatoid arthritis, type 1 diabetes and type 2 diabetes. These eight diseases make a major impact on public ill health¹³, cover a range of aetiologies and genetic predispositions, and have been extensively studied via SNP-based GWAS, including our earlier Wellcome Trust Case Control Consortium (WTCCC) study¹⁴.

Pilot experiment, array content, assay and samples

Pilot experiment. We undertook a pilot experiment to compare three different platforms for assaying CNVs and to assess the merits

of different experimental design parameters (see Supplementary Information for full details). On the basis of the pilot data, we chose the Agilent Comparative Genomic Hybridization (CGH) platform, and aimed to target each CNV with ten distinct probes, although in the analyses below we include any CNV targeted by at least one probe (Supplementary Fig. 9). Our analysis of the pilot CGH data indicated that the quality of the copy number signal for genotyping (rather than for discovery) at a CNV is reduced when the reference sample is homozygous deleted, in effect because the reference channel then just measures noise. To minimize this effect we used a fixed pool of DNAs as the reference sample throughout our main experiment.

Array content. Informed by our pilot experiment, we designed the CNV-typing array in a collaboration with the Genome Structural Variation Consortium (GSV) in which a preliminary set of candidate CNVs was shared at an early stage with the WTCCC. Table 1 summarizes the design content of the array, and Fig. 1 illustrates the various categories of designed loci unsuitable for association analysis. (See Methods for further details.)

Assay. In brief (see Supplementary Information for further details), the Agilent assay differentially labels parallel aliquots of the test sample and reference DNA (a pool of genomic UK lymphoblastoid cell-line DNAs from nine males and one female prepared in a single batch for all experiments) and then combines them, hybridizes to the array, washes and scans. Intensity measurements for the two different labels are made at each probe separately for the test and reference DNA. These act as surrogates for the amount of DNA present, with analyses typically relying on the ratio of test to reference intensity measurements at each probe.

Samples. A total of 19,050 case-control samples were sent for assaying: \sim 2,000 for each of the eight diseases and \sim 3,000 common controls (these were equally split between the 1958 British Birth Cohort

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Table 1 Discovery source for regions targeted on the genotyping array

		<u> </u>	<u> </u>			
Source of loci	Number of loci targeted	Number of loci analysed	Number of loci polymorphic with good calls			
CNVs						
GSV discovery project	10,835	10,217	3,096			
Affymetrix 500k	18	14	12			
Affymetrix 6.0	83	81	47			
Illumina 1M	82	81	18			
WTCCC CNV loci	231	209	108			
Novel sequence						
Novel insert regions	292	292	151			
Total	11,541	10,894	3,432			

GSV CNVs were prioritized according to extent of polymorphism in European discovery samples. See Methods for full details of other sources.

(58C) and the UK Blood Services (UKBS) controls). These were augmented by 270 HapMap1 samples (see ref. 12 for additional analyses of the HapMap data) and 610 duplicate samples for quality control purposes. About 80% of samples from the WTCCC SNP GWAS were used here. (See Supplementary Information for further details of sample collections, inclusion criteria, and so on.)

Data pre-processing, CNV calling and quality control

Data pre-processing. For each sample, raw data from the CNV experiment consist of intensity measurements for the test and reference sample for each probe. There are numerous choices at the data pre-processing stage, including how to normalize data to reduce inter-individual variation, and how to combine the information across the set of probes within a CNV. Several novel analytical tools substantially improved data quality, but no single approach works well for every CNV, so we carried through 16 pre-processing pipelines to maximize the number of CNVs that can be tested for association. (See Supplementary Information Section 4 for illustrations and a sense of the challenges.)

CNV calling. The objective in CNV calling at each CNV is to assign each assayed sample to a diploid copy-number class, which represents the sum of copy numbers on each allele. This step is analogous

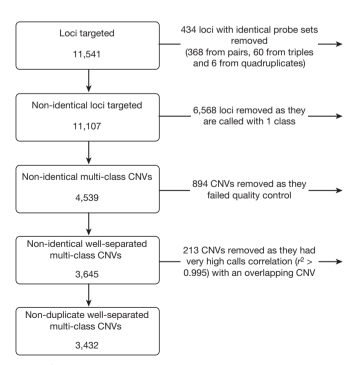


Figure 1 | Flowchart showing which CNVs are included on the array. The chart shows the reasons for CNVs being removed from consideration (the column of arrows and text to the right of the figure) from those originally targeted on the array, and the number of CNVs remaining at each stage of filtering.

to, but typically considerably more challenging than, calling genotypes from SNP-chip data. Available assays for SNPs are more robust and have better signal-to-noise properties than do available assays for CNVs¹⁵. We used two different statistical methods ('CNVtools', which is available as a Bioconductor package, and 'CNVCALL') in parallel to estimate the number of copy-number classes at each CNV and assign individuals to these classes. (See Supplementary Information for further details.) Figure 2 illustrates three multi-allelic CNVs that have attracted attention in the literature in part due to the difficulties in obtaining reliable data.

Quality control. After the application of quality control metrics to each sample and each CNV (see Methods), 17,304 case-control samples (of 19,050 initially) were available for association testing. There were 3,432 CNVs with more than one copy-number class which passed quality control and were included in subsequent analyses. At these CNVs, concordance of calls between pairs of duplicate samples was 99.7%.

Properties of CNVs

Single-class CNVs. Of the 10,894 distinct putative CNVs typed on the array after removal of detectable redundancies, 60% are called with a single copy-number class, and so cannot be tested for association. After detailed analyses (see Methods) we estimate that just under half of these are probably not polymorphic. For the remainder, the combination of the experimental assay and analytical methods we have used do not allow separate copy-number classes to be distinguished. Multi-class CNVs. A total of 4,326 CNVs were called with multiple classes. Of these, 3,432 passed quality control filters, which in practice means that the classes were well separated and thus that it was possible to assign individuals to copy-number classes with high confidence. Most of these CNVs (88%) have two or three copy-number classes, consistent with their having only two variants, or alleles, present in the population (we refer to these as bi-allelic CNVs). Note that some loci involving both duplications and deletions could be called with only three classes if both homozygote classes are very rare.

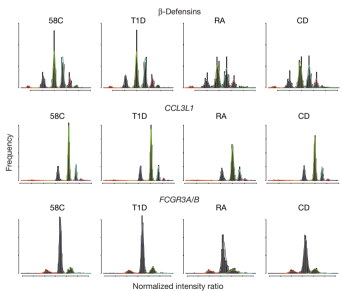


Figure 2 | Illustrative CNVs. Histograms of three multi-allelic CNVs (one per row) previously reported to be associated with autoimmune diseases: β-defensin (CNVR3771.10), *CCL3L1* (CNVR7077.12) and *FCGR3A/B* (CNVR383.1), showing 6, 5 and 4 fitted copy number classes, respectively. The histogram of normalized intensity ratios is shown for one control and the three autoimmune collections. Histograms are overlaid by the fitted distribution used to model each class (variously the red, blue, light-green, cyan, magenta and dark green curves). In all such figures, the area under the fitted curve of a particular colour is the same for all collections at the same CNV. 58C, 1958 British Birth Cohort; CD, Crohn's disease; RA, rheumatoid arthritis; T1D, type 1 diabetes.

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Allele frequencies. Supplementary Fig. 21 shows the distribution of minor allele frequency (MAF) for bi-allelic CNVs passing quality control. For example, 44% of autosomal CNVs passing quality control had MAF <5%. This is shifted towards lower MAFs compared to commonly used SNP chips. One consequence is that for given sample sizes association studies will tend to have lower power than for SNP studies. (See Supplementary Fig. 22 for power estimates.) Extrapolating from analyses described in ref. 12 gives an estimate that the 3,432 CNVs we directly tested represent 42–50% of common (MAF >5%) CNVs greater than 0.5 kilobases (kb) in length which are polymorphic in a population with European ancestry.

Tagging by SNPs. In the literature discussing the possible role of common CNVs in human disease there has been controversy over the extent to which CNVs will be in linkage disequilibrium with SNPs. If linkage disequilibrium between CNVs and SNPs were similar to that between SNPs, SNPs typed in GWAS would act as tags not only for untyped SNPs but also for untyped CNVs, and in turn SNP-based GWAS would have indirectly explored CNVs for association with disease. (See refs 16 and 17 for opposite views.) Our large-scale genotyping of an extensive CNV catalogue allows us to settle this question. In fact, CNVs that are typed well in our experiment are in general welltagged by SNPs—almost to the same extent that SNPs are well-tagged by SNPs (Supplementary Fig. 20). Among variable 2- and 3-class CNVs passing quality control with MAF>10%, 79% have $r^2 > 0.8$ with at least one SNP; for those with MAF <5%, 22% have $r^2 > 0.8$ with at least one SNP. This is consistent with the vast majority having arisen from unique mutational events at some time in the past. It follows that genetic variation in the form of common CNVs which can be typed on our array, has already been explored indirectly for association with common human disease through the SNP-based GWAS. In passing, we note that the high correlations between our CNV calls and SNP genotypes provide strong indirect evidence that our CNV calls are capturing real variation. It is possible that the CNVs that we cannot type well are systematically different from those that we can type, for example in having many more copy-number classes, and hence perhaps that they arise from repeated mutational events in the same region, in which case their linkage disequilibrium properties with SNPs could also be systematically different from the CNVs that we can type. We have no data that bear on this question, and it seems likely that such CNVs will be difficult to type genome wide on any currently available platforms.

Association testing

We performed association testing at each of the CNVs that passed quality control, in two parallel approaches. First, we applied a frequentist likelihood ratio association test that combines calling (using CNVtools) and testing into a single procedure, using an extension of an approach previously described¹⁸. Second, we undertook Bayesian association analyses in which the posterior probabilities from CNVCALL were used to calculate a Bayes factor to measure strength of association with the disease phenotypes. Important features of both sets of analyses are that they correctly handle uncertainty in assignment of individuals to copy-number classes, and by allowing for some systematic differences in intensities between cases and controls, that they provide robustness against certain artefacts which could arise from differences in data properties between cases and controls. There were no substantial differences between the broad conclusions from the frequentist and Bayesian approaches.

Our association analyses were based on a model in which a single parameter quantifies the increase in disease risk between successive copy-number classes, analogous to that underlying the trend test for SNP data. Various analyses of the robustness of our procedure, adequacy of the model, and lack of population structure were encouraging (see Methods and Supplementary Information). For example, Supplementary Fig. 23 shows quantile—quantile plots for the primary comparison of each case collection against the combined controls, and for the analogous comparisons between the two control

groups. These show generally good agreement with the expectation under the null hypothesis.

Careful analysis of our association testing revealed several sophisticated biological artefacts that can lead to false-positive associations. These include dispersed duplications, whereby the variation at a CNV is not in the chromosomal location in the reference sequence to which the probes in the CNV uniquely match, and a DNA source effect whereby particular CNVs, and genome-wide intensity data, can look systematically different according to whether the assayed DNA was derived from blood or cell lines. (See Box 1 for illustrations and further details.)

Independent replication of putative association signals is a routine and essential aspect of SNP-based association studies. Particularly in view of the differences in data quality between SNP assays and CNV assays, and the wide range of possible artefacts in CNV studies, replication is even more important in the CNV context. Several possible approaches to replication are available. When a CNV is well tagged by a SNP (or SNPs), replication can be undertaken by assessment of the signal at the tag SNP(s) in an independent sample, either by typing the SNP or by reference to published data. Where no SNP tag is available, direct typing of the CNV in independent samples is necessary, either using a qualitative breakpoint assay or a quantitative DNA dosage assay. In most cases there will be a choice of assays. Notably, replication via SNPs was possible for 15 out of 18 of the CNVs for which we undertook replication based on analysis of our penultimate data freeze

Figure 3 plots *P* values for the primary frequentist analysis for each CNV in each collection. Table 2 provides details of the top, replicated, association signals in our experiment after visual inspection of cluster plots to detect artefacts not removed by earlier quality control. Cluster plots for each CNV in Table 2 are shown in Supplementary Figs 18 and 19, and Supplementary Files 2 and 3.

There is one positive control for the diseases we studied, namely the known CNV association at the *IRGM* locus in Crohn's disease⁷. Reassuringly, our study found this association ($P = 1 \times 10^{-7}$, odds ratio (OR) = 0.68; throughout, all ORs are with respect to increasing copy number).

We identified three loci—HLA for Crohn's disease, rheumatoid arthritis and type 1 diabetes; *IRGM* for Crohn's disease; and *TSPAN8* for type 2 diabetes—at which CNVs seemed to be associated with disease, all of which we convincingly replicated through previously typed SNPs that tag the CNV, and a fourth locus (CNV7113.6) at which there is suggestive evidence for association and replication in both Crohn's disease and type 1 diabetes.

We observed CNVs in the HLA region associated variously with Crohn's disease (CNVR2841.20, $P=1.2\times10^{-5}$, OR = 0.80), rheumatoid arthritis (CNVR2845.14, $P=1.4\times10^{-39}$, OR = 1.77) and type 1 diabetes (CNVR2845.46, $P=8\times10^{-153}$, OR = 0.2). Copy number variation has previously been documented on various HLA haplotypes¹9 and owing to the extensive linkage disequilibrium in the region it is perhaps not unexpected to have found CNV associations in our direct study. Linkage disequilibrium across the HLA region has hampered attempts to fine-map causal variation across this locus, and we have no evidence that suggests that the HLA CNVs associated with autoimmune diseases in this study represent signals independent of the known associated haplotypes.

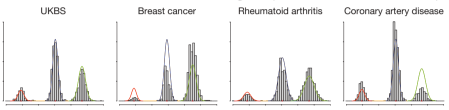
We identified two distinct CNVs 22 kb apart upstream of the *IRGM* gene, both of which are associated with Crohn's disease. The longer CNV (CNVR2647.1, $P=1.0\times10^{-7}$, OR = 0.68) has previously been identified⁷ as a possible causal variant on an associated haplotype first identified through SNP GWAS¹⁴, and acted as our positive control; however, the association of the smaller CNV (CNVR2646.1, $P=1.1\times10^{-7}$, OR = 0.68, located <2 kb downstream from a different gene, *C5orf62*) is a novel observation. Although direct experimental evidence links the associated haplotypes with variation in expression of the *IRGM* gene, it does not bear on the question of which of the two CNVs or the associated SNPs

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Box 1 | Some artefacts in CNV association testing

Some types of artefacts, such as population structure and calling artefacts, are very similar to those seen in SNP studies. Others, related to differences in data properties between cases and controls, can be potentially more serious for CNVs^{26,27}. In this box we draw attention to some specific artefacts of biological interest that we observed and which researchers should consider as explanations of putative disease-relevant associations. We note that, for the unwary, some of these artefacts could easily survive 'replication' of an association.

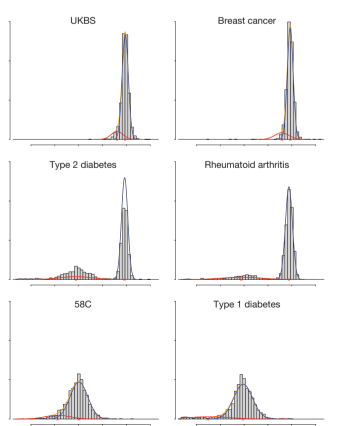
First, we consider dispersed CNVs. Box 1 Fig. 1 shows cluster plots for a particular CNV (CNVR2664.1) that shows a strong case-control association signal for breast cancer cases ($P = 5 \times 10^{-143}$, higher copy number for disease) with a related signal for rheumatoid arthritis ($P = 3 \times 10^{-27}$), and a signal in the opposite direction for coronary artery disease ($P = 4 \times 10^{-30}$). The right-hand class (green curve) has a higher frequency in breast cancer (and rheumatoid arthritis), and a lower frequency in coronary heart disease. (The area under the green curve is the same for each collection.) This turned out to be an artefact caused by differences in sex ratio in the various case and control samples (breast cancer, 100% female; rheumatoid arthritis, 74% female; coronary artery disease, 22% female; controls, 50% female). Comparing breast cancer cases against female controls abolished the signal. The CNV is annotated as being on chromosome 5 and all 10 probes in the CNV map uniquely to chromosome 5 in the human reference sequence. However, we found that SNPs which tagged the variation at this CNV all mapped to the X chromosome and that the region containing the probes for this CNV is present on the X chromosome in the Venter genome. We conclude that the CNV is a dispersed duplication, with the variation actually occurring on the X chromosome, and not on chromosome 5. We found one similar example, of a CNV (CNVR1065.1, featured in Table 2 as a replicated association) annotated as mapping uniquely to chromosome 2 that shows a strong signal in type 1 diabetes and rheumatoid arthritis. Careful examination shows it to be another dispersed duplication where the polymorphism is located in the HLA region, and is well tagged by HLA SNPs known to be associated with both diseases. Supplementary Fig. 27 shows the clear evidence from interchromosomal linkage disequilibrium that these two loci are dispersed duplications.



Box 1 Fig. 1 | Dispersed duplications leading to false-positive associations.

Second, we consider variation in DNA source. Box 1 Fig. 2 shows cluster plots for a different CNV (CNVR866.8) with marked differences in type 2 diabetes as compared with the UKBS controls (or against just the 58C controls). The plots show histograms of normalized intensity ratios for six collections. Examination of the pattern across collections is interesting. The collections in the top row show a single tight peak towards the right of the plot. Those in the bottom row show a single, more dispersed peak to the left. The collections in the middle row show evidence of both peaks. It turns out that for collections with the tight peak all DNA samples were derived from blood whereas all samples in the two collections with the single dispersed peak had DNA derived from cell lines. The remaining collections contain some DNAs derived from both sources. This CNV (and many others) thus exhibit systematically different behaviour depending on the DNA source. Box 1 Fig. 3 shows a plot of the second (PC2) and third (PC3) principal components of the array-wide intensity data (plot created using all samples after quality control from all ten collections using data from all CNVs, with each point representing one sample, with the points coloured according to whether that sample was derived from blood (red) or cell lines (blue)). It is clear that these two components can almost perfectly classify samples according to the source of the DNA.

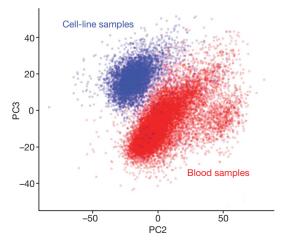
Lymphoblas to id cell lines are typically grown from transformed B cells, whereas DNA extracted from blood comes largely from a mixture of white blood cells. One specific feature of B cells is that each B cell has been subject to its own pattern of rearrangements around the immunoglobulin genes via the process of the



Box 1 Fig. 2 | DNA source effect leading to false-positive associations.

of V(D)J recombination. This suggests a natural candidate for our observed DNA source effect, and indeed the CNV illustrated in Box1Fig. 2 is located close to one of the immunoglobulin genes, as are the other instances we have found of similar gross DNA source effects. But it is not the whole story. Principal components analysis of genome-wide intensity data with any probe mapping to within 1 megabase of an immunoglobulin gene excluded from analysis (Supplementary Fig. 29) shows reasonably clear discrimination by DNA source (although less clear than when all probes are included), with many probes, genome-wide, contributing to the discrimination.

Dispersed duplications and DNA source effects represent interesting biological artefacts. We also observed more prosaic effects. As one example, Supplementary Fig. 30 shows that there are systematic effects on probe intensity of the row of the plate in which a sample was run.



Box 1 Fig. 3 | Principal component analysis showing DNA source effect.

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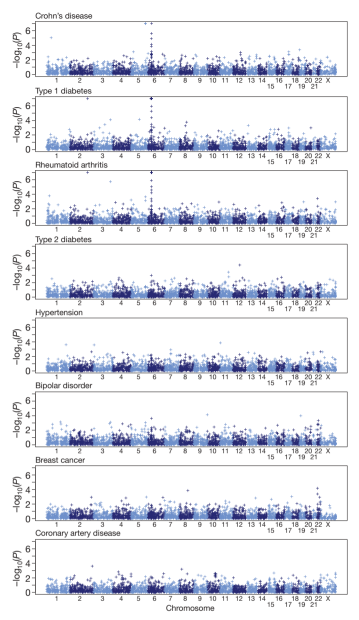


Figure 3 | **Genome-wide association results.** Distribution of $-\log_{10}(P)$ along the 23 chromosomes where P is the P-value for the one degree-of-freedom test of association for each disease. The x-axis shows the chromosomes numbered from 1 (on the left) to X (on the right). CNVs included in these plots were filtered on the basis of a clustering quality score (see Supplementary Information for details) and manual inspection of the most significant associations. The two apparent associations on chromosome 2 for rheumatoid arthritis and type 1 diabetes result from a dispersed duplication in which the variation is actually located within the HLA locus (see Box 1).

might be driving this variation⁷. Our conditional regression analyses on the two CNVs and SNPs on this haplotype do not point significantly to any one of these as being more strongly associated.

SNP variation in the *TSPAN8* locus was recently shown to be reproducibly associated with type 2 diabetes²⁰, but the potential role of a CNV is a novel observation. This CNV (CNVR5583.1, $P=3.9\times10^{-5}$, OR = 0.85) potentially encompasses part or all of an exon of *TSPAN8* and so is a plausible causal variant. The most significantly associated SNP identified in the recent meta-analysis is only weakly correlated with the CNV as originally tested ($r^2=0.17$), and so the CNV may simply be weakly correlated with the true causal variant. Closer examination of probe-level data at this CNV indicates a series of different events (including an inverted duplication and a deletion) resulting in more complex haplotypes than those tested for

association by our automated approach. With this more refined definition of haplotypes the signal is stronger. (See Supplementary Information for details.)

CNVR7113.6 lies within a cluster of segmentally duplicated sequences that demarcate one end of a common 900-kb inversion polymorphism on chromosome 17 that has previously been shown to be associated with number of children and higher meiotic recombination in females²¹. The CNV shows weak evidence for association with Crohn's disease ($P = 1.8 \times 10^{-3}$, OR = 1.15) and type 1 diabetes ($P = 1.1 \times 10^{-3}$, OR = 1.13), but is in extremely high linkage disequilibrium ($r^2 = 1$) with SNPs known to tag the inversion, and so is in tight linkage disequilibrium with a long haplotype spanning many possible causal variants. This CNV encompasses at least one spliced transcript, but no high-confidence gene annotations. Finemapping the causal variant within such a long, tightly linked haplotype is likely to prove challenging.

In addition to the loci in Table 2, we undertook replication on 13 other loci, detailed in Supplementary Table 13, for which there was some evidence of association ($P < 1 \times 10^{-4}$ or $\log_{10}(\text{Bayes factor (BF)}) > 2.1)$ in our analysis of the penultimate data freeze. Replication results were negative for all these loci. Several other loci for which there is weak evidence ($P < 1 \times 10^{-4}$ or $\log_{10}(\text{BF}) > 2.6$) for association in our final data analysis are listed in Supplementary Table 14.

To investigate further the potential role of CNVs as pathogenically relevant variants underlying published SNP associations, we took 94 association intervals in type 1 diabetes, Crohn's disease and type 2 diabetes (excluding the HLA), and for the index SNP in each association interval assessed its correlation with our calls at 3,432 CNVs. We identified two index SNPs as being correlated with an r^2 of greater than 0.5 with a called CNV. The SNPs were: rs11747270 with both CNVR2647.1 and CNVR2646.1 (*IRGM*), and rs2301436 with CNVR3164.1 (*CCR6*), both for Crohn's disease. Both of these association intervals were also identified in an independent analysis using CNV calls on HapMap samples by ref. 12.

As a further test of our approach, we examined three multi-allelic CNVs that have attracted attention in the literature, both for the challenges of obtaining reliable data and for putative associations with a range of autoimmune diseases: CCL3L1 (our CNVR7077.12); β -defensins (CNVR3771.10); and FCGR3A/B (CNVR383.1)^{10,22–24}. Encouragingly, all three CNVs pass quality control and give good quality data. Figure 2 shows cluster plots for these CNVs in our experiment. The best calls for the three CNVs required the use of two analysis pipelines (sets of choices about normalization and probe summaries) different from our standard pipeline. None of the CNVs shows significant association with the three autoimmune diseases in our study after allowance for multiple testing. In particular, we do not see formally significant evidence to replicate the reported association for CCL3L1 and rheumatoid arthritis²⁴ (nominal P=0.058).

We also assessed whether CNVs that delete all or part of exons might be enriched among disease susceptibility loci, even if our study were not well-powered enough to see statistically significant evidence of association for individual CNVs. To do so, we compared the 53 exonic deletion CNVs¹² that passed quality control with collections of CNVs of the same size, matched for MAF and numbers of classes. We used a (two-sided) Wilcoxon signed-rank test²⁵ to ask whether the strength of signal for association (measured by Bayes Factors) was systematically different for the exon-deletion CNVs as compared to the matched CNVs. We found no evidence that deletion of an exon systematically changed evidence for association (see Supplementary Information). In a related analysis, we compared CNVs passing quality control that were well tagged by SNPs ($r^2 > 0.8$) to those passing quality control that were not, again matching for MAF and number of classes (excluding low MAF CNVs and those failing Hardy–Weinberg equilibrium tests to avoid calling artefacts). There was no evidence that CNVs passing quality control that are not well tagged by SNPs are enriched for stronger signals of association compared to those which were well tagged (see Supplementary Information).

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Table 2 | Replicated CNV associations and those at replicated loci

Disease	Chr.	Start (bp) (CNV)	Length (kb)		Fitted no. classes*	Combined controls (P)†	Extended reference (P)	Combined controls (log ₁₀ (BF))‡	reference		Extended reference (OR)	MAF		Replication size		Replication size (P)
													Cases#	Ctrls	Cases	3126 (1)
T2D	12	69,818,942 (CNVR5583.1)	1.0	TSPAN8	3	3.9×10 ⁻⁵	2.5×10 ⁻⁶	2.8	4.3	0.85	0.85	0.40	0.36	5,579	4,549☆	3.9×10 ⁻⁵
CD	5	150,157,836 (CNVR2646.1)	3.9	IRGM	3	1.1×10^{-7}	5.5×10 ⁻⁵	5.8	4.1	0.68	0.75	0.07	0.10	7,977	6,894☆	7.5×10 ⁻¹³
CD	5	150,183,562 (CNVR2647.1)	20.1	IRGM	3	1.0×10^{-7}	4.3×10 ⁻⁵	6.1	3.8	0.68	0.76	0.07	0.10	7,977	6,894☆	3.9×10 ⁻¹⁰
CD	6	31,416,574 (CNVR2841.20)	5.1	HLA	3	1.7×10^{-5}	1.1×10^{-5}	3.6	3.9	0.80	0.82	0.19	0.23	NA	NA	NA
T1D	6	32,582,950 (CNVR2845.46)	6.7	HLA	2	8.0×10 ⁻¹⁵³	2.1×10 ⁻¹⁹⁶	125.5	154.4	0.20	0.26	0.14	0.01	NA	NA	NA
RA	6	32,609,209 (CNVR2845.14)	4.0	HLA	4	1.4×10^{-39}	8.1×10^{-60}	51.5	73.5	1.77	1.83	NA	NA	NA	NA	NA
RA	2→6		0.8	HLA	3	6.8×10 ⁻⁴⁹	1.6×10 ⁻⁶⁹	51.0	73.7	1.85	1.94	0.36	0.49	NA	NA	NA
T1D	2→6		0.8	HLA	3	1.3×10 ⁻²⁹	1.1×10^{-39}	28.0	38.4	1.62	1.61	0.36	0.47	NA	NA	NA
RA	NA	NA (AC 000138.1 44)	5.6	HLA	3	8.3×10 ⁻⁴	1.1×10^{-5}	1.3	2.7	0.87	0.86	0.25	0.28	2,743	3,398	1.1×10 ⁻³
T1D	NA	NA (AC 000138.1 44)	5.6	HLA	3	2.0×10 ⁻³¹	2.7×10 ⁻⁴⁵	31.0	45.1	0.59	0.57	0.25	0.36	2,649	3,883	7.3×10 ⁻⁵⁰
CD	17	40,930,407 (CNVR7113.6)		Chr17inv	3	1.2×10^{-3}	5.8×10 ⁻⁴	1.4	1.6	1.15	1.14	0.24	0.21	6,069	4,978☆	8.6×10 ⁻⁵
T1D	17	40,930,407 (CNVR7113.6)	33.9	Chr17inv	3	1.6×10 ⁻³	7.5×10 ⁻⁴	1.0	1.2	1.13	1.12	0.24	0.21	9,395	7,911☆	4.6×10 ⁻⁶

Only one of the several associated CNVs mapping to the HLA in the reference sequence is shown for each of rheumatoid arthritis, type 1 diabetes and Crohn's disease. Further details of replication assays and methods are given in Supplementary Information. AC_000138.1_44 is a novel sequence insertion present in the Venter genome sequence but not in the reference sequence and hence no chromosomal location is presented. Minor allele frequency is only estimated for CNVs with three or fewer copy number classes. CD, Crohn's disease; RA, rheumatoid arthritis; T1D, type 1 diabetes; T2D, type 2 diabetes.

Discussion

We have undertaken a genome-wide association study of common copy-number variation in eight diseases by developing a novel array targeting most of a recently discovered set of CNVs. Our findings inform understanding of the genetic contributions to common disease, offer methodological insights into CNV analysis, and provide a resource for human genetics research.

One major conclusion is that considerable care is needed in analysing copy-number data from array CGH experiments. Choices of normalization, probe summary and probe weighting can make major differences to data quality and utility in association testing. Notably, the optimal choices vary greatly across the CNVs we studied.

A second major conclusion is that CNV association analyses are susceptible to a range of artefacts that can lead to false-positive associations. Some are a consequence of the less-robust nature of the data compared to SNP chips. But others, such as systematic differences depending on DNA source (for example, blood versus cell lines) and dispersed duplications, are more subtle. Several artefacts could survive replication studies. Simultaneously studying eight diseases helped greatly in identifying these artefacts, and stringent quality control was invaluable in eliminating false-positive associations. At least for currently available CNV-typing platforms, we recommend considerable care in interpreting putative CNV associations combined with independent replication on a different experimental platform.

Despite the important technical challenges and potential artefacts discussed above, we have demonstrated that high-confidence CNV calls can be assigned in large, real-world case-control samples for a substantial proportion of the common CNVs estimated to be present in the human genome. We have identified directly several CNV loci that are associated with common disease. Such loci could contribute to disease pathogenesis. However, the loci identified are well tagged by SNPs and, hence, the associations can be, and were, detected indirectly via SNP association studies.

There is a marked difference between the number of confirmed, replicated associations from our CNV study (3 loci) and that from the comparably sized WTCCC1 SNP GWAS of seven diseases and its immediate follow-up ($\sim\!24\,\mathrm{loci}$). (In assessing the importance of CNVs in disease, it is the absolute number of associations, rather than the proportion among loci tested, that is important.) Following ref. 12 we estimated that our study directly tests approximately half of all autosomal CNVs $>\!500\,\mathrm{bp}$ long, with MAF $>\!5\%$. For such CNVs, our power averages over 80% for effects with odds ratios $>\!1.4$, and $\sim\!50\%$ for odds ratio =1.25 (Supplementary Fig. 22). We conclude that at least for the eight diseases studied, and probably more generally, there are unlikely to be many associated CNVs with effects of this magnitude.

Might there be many more common disease-associated CNVs each of small effect, in the way that we now know to be the case with SNP associations for many diseases? The total number of CNVs over 500 bp with MAF >5% is limited (estimated to be under 4,000 (ref. 12)), so unless many of these simultaneously affect many different diseases (something for which we saw no evidence outside of the HLA region) there would seem to be insufficient such CNVs for hundreds to be associated with each of many common diseases. In addition, most common CNVs (MAF >5%) are well tagged by SNPs, and thus amenable to indirect study by SNP GWAS. Examining the large meta-analyses of SNP GWAS for Crohn's disease, type 1 diabetes and type 2 diabetes, there were 95 published associated loci of which only 3, including HLA, had the property that CNVs correlated with the associated SNPs; two of these were detected in our direct study.

We conclude that common CNVs typable on current platforms are unlikely to have a major role in the genetic basis of common diseases, either through particular CNVs having moderate or large effects (odds ratios >1.3, say) or through many such CNVs having small effects. In particular, such common CNVs seem unlikely to account for a substantial proportion of the 'missing heritability' for these diseases. Among the CNVs that we could type well, those not well

^{*} The number of diploid copy-number classes.

[†] P value from the frequentist association test combining UKBS and 58C as controls.

 $[\]dot{z}$ The \log_{10} of the Bayes factor from the Bayesian association analysis combining UKBS and 58C as controls.

[§] The odds ratio estimated for each additional copy of the CNV based on both UKBS and 58C as controls.

^{||} Extended reference refers to the analogous quantities calculated in comparing cases of the disease in question with UKBS, 58C and aetiologically unrelated cases.

The minor allele frequency in controls (UKBS plus 58C).

[#]The minor allele frequency in cases.

[☆]Replication sample includes WTCCC samples.

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tagged by SNPs have the same overall association properties as those which are well tagged. We saw no enrichment of association signals among CNVs involving exonic deletions.

We have argued elsewhere¹⁴ that the concept of 'genome-wide significance' is misguided, and that under frequentist and Bayesian approaches it is not the number of tests performed but rather the prior probability of association at each locus that should determine appropriate *P* value thresholds. Here, to reduce the possibility of missing genuine associations, we deliberately set relaxed thresholds for taking CNVs into replication studies. Having completed these analyses the hypothesis that, a priori, an arbitrary common CNV is much more likely than an arbitrary common SNP to affect disease susceptibility is not supported by our data.

Limitations. Our findings should be interpreted within the context of several limitations. First, despite our successes in robustly testing some of the previously noted challenging CNVs in the genome, for some CNVs we could not reliably assign copy-number classes from our assay. We estimate that just under half of these were not polymorphic in our data, being either false positives in the discovery experiment, or very rare in the UK population. For the remainder, we were also unable to perform reliable association analyses based directly on intensity measurements (that is, without first assigning individuals to copy number classes; data not shown). Such CNVs might plausibly be systematically different from those that we do type successfully, in which case it is not possible to extrapolate from our results to their potential role in human disease. Second, we note that we have not studied CNVs of sequences not present in the reference assembly, high-copy-number repeats such as LINE elements, or most polymorphic tandem repeat arrays, and our findings may not generalize to such variation. Finally, our experiment was powered to detect associations with common copy number variation and our observations and conclusions do not necessarily generalize to the study of rare copy number variants. Different approaches will be necessary to investigate the contribution of such variation to common disease.

METHODS SUMMARY

Pilot study. A total of 384 samples spanning a range of DNA quality were assayed for 156 previously identified CNVs on each of three different platforms: Agilent CGH, NimbleGen CGH and Illumina iSelect. The pilot experiment contained many more probes per CNV than we anticipated using in the main study, and replicates of these probes, to allow an assessment of data quality as a function of the number of probes per CNV and of the merits of replicating probes predicted in advance to perform well, compared to using distinct probes.

Sample selection. Case samples came from previously established UK collections. Control samples came from two sources: half from the 1958 Birth Cohort and half from a UK Blood Service sample. Approximately 80% of samples had been included within the WTCCC SNP GWAS study. The 610 duplicate samples were drawn from all collections.

Array design. The main study used an Agilent CGH array comprising 105,072 long oligonucleotide probes. Probes were selected to target CNVs identified mainly through the GSV discovery experiment¹², with some coming from other sources. Ten non-polymorphic regions of the X chromosome were assayed for control purposes.

Array processing. Arrays were run at Oxford Gene Technology (OGT). The samples were processed in batches of 47 samples drawn from two different collections, with each batch containing one control sample for quality control purposes. These batches were randomized to protect against systematic biases in data characteristics between collections.

Data analysis. Primary data and low-level summary statistics were produced at OGT. All substantive data analyses were undertaken within the consortium. Plates failing quality control metrics were rerun, as were 1,709 of the least well-performing samples. Details of the common CNVs assayed in this study, including any tag SNP, are given at http://www.wtccc.org.uk/wtcccplus_cnv/supplemental.shtml.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

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Author Contributions are listed in Supplementary Information.

Author Information Summary information for the CNVs studied, including genomic locations, numbers of classes and SNP tags on different platforms is available at http://www.wtccc.org.uk/wtccplus_cnv/supplemental.shtml. Full data are available, under a data access mechanism, from the European Genome-phenome Archive (http://www.ebi.ac.uk/ega/page.php). Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Correspondence and requests for materials should be addressed to P.D. (peter.donnelly@well.ox.ac.uk).

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METHODS

Pilot experiment. Full details of Methods are given in the Supplementary Information, but in brief a total of 384 samples from four different collections spanning the range of DNA quality encountered in our previous WTCCC SNP-based association study¹⁴ were assayed for 156 previously identified CNVs on each of three different platforms: Agilent Comparative Genomic Hybridization (CGH), and NimbleGen CGH (run in service laboratories) and Illumina iSelect (run at the Sanger Institute). The pilot experiment contained many more probes per CNV (40–90 depending on platform) than we anticipated using in the main study, and replicates of these probes, to allow an assessment of data quality as a function of the number of probes per CNV and of the merits of replicating probes predicted in advance to perform well, compared to using distinct probes.

The Agilent CGH platform performed best in our pilot and we settled on an array that comprised 105,072 long oligonucleotide probes. On the basis of the pilot data we aimed to target each CNV with 10 distinct probes. Actual numbers of probes per CNV on the array varied from this for several reasons (see Supplementary Information and Supplementary Fig. 9), and we included in our analyses any CNV with at least one probe on the array.

Array content, assay and samples for the main experiment. Array content: the GSV discovery experiment¹² involved 20 HapMap Utah residents with European ancestry (CEU) and 20 HapMap Yoruban (YRI) individuals, and 1 Polymorphism Discovery Resource individual, assayed via 20 NimbleGen arrays containing a total of 42,000,000 probes tiled across the assayable portion of the human reference genome. We prioritized CNVs for our experiment based on their frequency in the discovery sample, with those identified in CEU individuals given precedence. A total of 10,835 out of 11,700 CNVs were included from the list provided by the GSV, with those not included on the array being present in discovery in only 1 YRI individual and not overlapping genes or highly conserved elements. This list was augmented by any common CNVs not present among the GSV list found from analyses of Affymetrix SNP 6.0 data in HapMap 2 samples (83 CNVs), Illumina 1M data in HapMap 3 samples (82 CNVs), analyses of Affymetrix 500K samples (18 CNVs)^{7,29,30}, and from our own analyses of WTCCC1 SNP data (231 CNVs). In addition, we sought to identify CNVs not present in the human reference sequence through analyses of published 31,32 novel sequence insertions (292 CNVs in total). Thus in total, our array targeted 11,541 putative CNVs. Ten non-polymorphic regions of the X chromosome were also assayed for control purposes.

Most loci targeted on the CNV-typing array derive from microarray-based CNV discovery, which is inherently imprecise. In contrast to SNP discovery by sequencing, arrays do not provide nucleotide-level resolution, nor do they locate additional copies of a sequence in the genome. As a result, when CNVs called in different individuals overlap, but are not identical, these could be called as one or two different CNVs, and where discovered CNVs involve probes which map to multiple places in the reference genome, they might be called as CNVs in each of these locations. Interpretation of counts of CNVs from discovery experiments is thus not straightforward. Data on CNVs across thousands of individuals provide added power to refine CNV definitions and derive a non-redundant set of CNVs. In addition, our CNV-typing array draws together CNVs from different sources, and additional redundancy between these, although minimized during array design, can be identified and removed. Analyses of the final array design revealed 434 of the 11,541 CNVs to be redundant because they were targeted by exactly the same probes as other CNVs on the array, and analysis of our array data revealed a further 213 of 562 CNVs to be redundant from instances where overlapping CNVs passing quality control were called as distinct in discovery yet had effectively identical copy-number calls. See Supplementary Information Section 3.1 for further details on array content.

Assay: arrays were run at Oxford Gene Technology (OGT), with each plate containing one control sample for quality control purposes. Primary data and low-level summary statistics were produced at OGT. All substantive data analyses were undertaken within the consortium. Plates that failed pre-specified quality control metrics were rerun on the array, and in addition we repeated 1,709 of the least well-performing samples, chosen according to our own quality control analyses. (See Supplementary Information for further details.)

Samples: the WTCCC CNV study analysed cases from eight common diseases (breast cancer, bipolar disorder, coronary artery disease, Crohn's disease, hypertension, rheumatoid arthritis, type I diabetes, and type 2 diabetes) and two control cohorts (1958 Birth Cohort (58C) and the UK Blood Service collection (UKBS)). The number of subjects from each cohort that were analysed and the numbers that passed each phase of the quality control procedures within this study are shown in Supplementary Table 7. For bipolar disorder, coronary heart disease, Crohn's disease, hypertension, rheumatoid arthritis, type 1 diabetes, type 2 diabetes and the two control cohorts, a large proportion of the subjects studied in this experiment were the same as those in the WTCCC1 SNP GWAS (Supplementary

Table 2). Where sufficient DNA was not available for the original WTCCC1 individuals, additional new samples from the same cohorts were used, selected using the same approaches used for the WTCCC1 samples. Any samples that failed any of the relevant quality control metrics in WTCCC1 were excluded from consideration for this experiment. The breast cancer cohort was not included in the WTCCC1 SNP GWAS.

Data pre-processing, CNV calling and quality control. Data pre-processing: for each of the targeted loci, the subset of probes that target the locus of interest (at least 1-bp overlap) while also targeting the least number of additional CNVs was selected for assaying (see Supplementary Information Section 4.2 for more details). A total of 16 different analysis 'pipelines' were used to create onedimensional intensity summaries for each CNV. First, a range of different methods were used to create single intensity measurements for each probe from the red channel (test DNA) and green channel (reference DNA) intensity data. This included different methods for normalization of the signals (see Supplementary Information Section 4.3 for details). Second, some pipelines incorporated a new method called probe variance scaling (PVS) that weights probes based on information derived from duplicate samples (see Supplementary Information Section 4.5 for details). Third, some pipelines used the first principal component of the normalized probe intensities to summarize the probe-level data to CNV-level data, whereas other pipelines used the mean of the probe intensities. Finally, some pipelines additionally used a linear discriminant function (LDF) to refine further the summaries based on information from an initial round of genotype calling (see Supplementary Information Section 4.4 for details).

CNV calling: algorithmic details of the two calling methods used (CNVtools and CNVCALL) are provided in Supplementary Information Section 6. Each method was applied separately to the intensity summaries created from each of the 16 pre-processing pipelines for each CNV locus.

Quality control: samples were excluded on the basis of sample handling errors, evidence of non-European ancestry, evidence of sample mixing, manufacturer's recommendations on data quality, outlying values of various pre-calling and post-calling quality metrics, and identity or close relatedness to other samples (see Supplementary Information Section 5.1 for further details). To choose which pipeline to use for a given CNV we used the pipeline that gave the highest number of classes and the highest average posterior probability in cases where more than one pipeline gave the same maximum number of classes. CNVs were excluded that had identical probe sets to other CNVs, that were called with one class in all pre-processing pipelines, that had low average posterior calls in all pre-processing pipelines, or that had a high calls correlation with an overlapping CNV (see Supplementary Information Section 5.2 for further details).

Properties of CNVs. Single class CNVs: Supplementary Table 15 shows the proportion of the single-class CNVs from the GSV discovery project broken down according to the number of individuals and population(s) in which they were discovered. Of the GSV CNVs discovered in CEU, 52% are single class in our data, whereas a higher proportion (74%) of GSV CNVs discovered exclusively in YRI are single class, as would be expected. CNVs at which distinct copy number classes cannot be distinguished might result because: (1) although polymorphic, the signal to noise ratio at that CNV does not allow reliable identification of distinct copy-number classes; (2) the copy-number variant has an extremely low population frequency; or (3) the CNV was a false positive in a discovery experiment and is in fact monomorphic. In a genuinely polymorphic CNV, the intensity measurements within a pair of duplicates should be more similar than between a random pair of distinct individuals. Intensity comparisons between duplicates and random pairs of individuals thus allow estimates of the proportion of single-class CNVs which are not copy-number variable in our data (see Supplementary Information). These estimates range from ~23% for single-class CNVs discovered in two or more CEU individuals to ~50% of single-class CNVs discovered exclusively in YRI (see Supplementary Information for details). We estimate that 18% of GSV CNVs discovered in CEU do not exhibit polymorphism in our UK sample. This figure is similar to the GSV estimate for false positives in discovery of 15%12. Overall, considering CNVs on the array from all sources, we estimate that 26% do not exhibit polymorphism, so that just under half of the CNVs that seem in our data to have a single class are likely not to be polymorphic. Not all of these will be false positives in discovery; some represent CNVs that are either unique to the individual in which they were discovered or are extremely rare in the UK population.

Multi-class CNVs: a companion study¹² estimated that 83% of the bi-allelic CNVs it genotyped represent deletions, with the remainder being duplications. Supplementary Table 7 compares the number of copy-number classes estimated by the two calling algorithms used in the analyses for each of the CNVs passing quality control. Most differences in numbers of called classes between the algorithms arise from CNVs where one class is very rare and is handled differently by the algorithms (for example, called as a separate class in one algorithm but classed as outlier samples or merged with a larger class by the other).

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These 3,432 CNVs include 80% of the CNVs genotyped on the Affymetrix 6.0 array that are common (MAF >5%) in a population with European ancestry³³; conversely only 15% of the common CNVs we called could be called using the Affymetrix 6.0 array.

Allele frequencies: we calculated minor allele frequencies (MAFs) for 2- and 3-class CNVs by assuming that these CNVs were biallelic and using the expected posterior genotype counts (see Supplementary Information Section 7.3 for further details).

Tagging by SNPs: to determine how well tagged the CNVs analysed in our experiment were by SNPs, we carried out correlation analyses using control samples that were common to the current studies and other WTCCC studies. We analysed three different collections of SNPs. We used imputed HapMap2 SNP calls in the WTCCC1 study that used the Affymetrix 500k array, and actual calls from the WTCCC2 study using both the Affymetrix 6.0 array and a custom Illumina 1.2M array. In all cases we used samples from the UKBS collection (see Supplementary Information Section 7.1 for further details).

Geographical variation: geographical information, at the level of 13 pre-defined regions of the UK, was available for 82% of the samples in our study and we undertook analyses for differences in copy-number class frequencies between regions. The results, shown in Supplementary Fig. 24, confirm that there is no major genome-wide population structure, but that, unsurprisingly, there is differentiation at CNVs within HLA. It does not seem easy to determine whether other regions with low P values in this test represent genuine departures from the null hypothesis of no differentiation, rather than chance effects, although we note that the third most regionally differentiated CNV outside the HLA (CNVR7722.1, $P=3\times10^{-5}$, 12-d.f.) is a deletion located within the gene LILRA3, which may act as soluble receptor for class I MHC antigens, and so would be consistent with the observed HLA stratification. This deletion is also the subject of a reported disease association 34 in multiple sclerosis, a finding that may require some caution given the level of geographical stratification at this CNV in our data. (See Supplementary Information Section 9.1 for further details.)

Association testing. Diagnostic plots such as quantile–quantile and cluster plots were created using R. Cluster plots were visually inspected for all CNVs with putative associations.

Principal component analysis (PCA) was applied to the summarized intensity levels for all CNVs, and for all samples that passed quality control. Plots of the first ten principal components were coloured by various sample parameters and these revealed some of the artefacts described in Box 1.

Where possible, replication was carried out by using data from other studies for SNPs that tag the CNVs of interest. Where there was no SNP tag available, breakpoint or direct quantitative CNV assays were designed (see Supplementary Information Section 9 for further details).

We used a two-sided Wilcoxon signed-rank test to test for differences between distributions of Bayes factors between different subsets of CNVs (those that delete all or part of an exon versus those that do not, and CNVs that are well-tagged by SNPs versus those that are not well tagged). (See Supplementary Information Section 9.5 for further details.)

Testing for population stratification: all our samples are from within the UK, and we have excluded any for which the genetic data suggest evidence of non-European ancestry. All collections in this study, apart from breast cancer, were involved in the WTCCC SNP GWAS, and across these collections 80% of samples coincided between the two studies. Analysis of the WTCCC SNP data¹⁴ established that population structure was not a major factor confounding association testing. Similar analyses using SNP data available for the breast cancer samples yielded similar results (data not shown). These SNP results reinforce the evidence from the quantile—quantile plots in Supplementary Fig. 23 and our geographical analyses of the CNV data.

Expanded reference group analysis: in addition to our primary case-control analyses, following ref. 14 we also undertook expanded reference group analyses, in which copy-number class frequencies in cases for a particular disease are compared with those for controls and the other diseases with no aetiological or known genetic connection (see Supplementary Table 10 for details).

Other analyses. We used information on variability between duplicate samples to determine whether CNVs called with one class show signals of polymorphism (details are given in Supplementary Information Section 9.2).

We used estimates of the number of common autosomal CNVs segregating in a population of European ancestry from ref. 12 to estimate the coverage of common autosomal CNVs in our study (see Supplementary Information Section 9.3 for further details).

We designed a series of PCR primers to analyse further the complex signals associated with CNVR5583.1 found in the *TSPAN8* region. (See Supplementary Information Section 9.4 for further details.)

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