Integrating *de novo* and inherited variants in over 42,607 autism cases identifies mutations in new moderate risk genes

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Abstract

Despite the known heritable nature of autism spectrum disorder (ASD), studies have primarily identified risk genes with *de novo* variants (DNVs). To capture the full spectrum of ASD genetic risk, we performed a two-stage analysis of rare *de novo* and inherited coding variants in 42,607 ASD cases, including 35,130 new cases recruited online by SPARK. In the first stage, we analyzed 19,843 cases with one or both biological parents and found that known ASD or neurodevelopmental disorder (NDD) risk genes explain nearly 70% of the genetic burden conferred by DNVs. In contrast, less than 20% of genetic risk conferred by rare inherited loss-of-function (LoF) variants are explained by known ASD/NDD genes. We selected 404 genes based on the first stage of analysis and performed a meta-analysis with an additional 22,764 cases and 236,000 population controls. We identified 60 genes with exome-wide significance (p < 2.5e-6), including five new risk genes (*NAV3, ITSN1, MARK2, SCAF1, and HNRNPUL2*). The association of *NAV3* with ASD risk is entirely driven by rare inherited LoF variants, with an average relative risk of 4, consistent with moderate effect. ASD individuals with LoF variants in the four moderate risk genes (*NAV3, ITSN1, SCAF1, and HNRNPUL2*, n = 95) have less cognitive impairment compared to 129 ASD individuals with LoF variants in well-established, highly penetrant ASD risk genes (*CHD8, SCN2A, ADNP, FOXP1, SHANK3*) (59% vs. 88%, p = 1.9e-06). These findings will guide future gene discovery efforts and suggest that much larger numbers of ASD cases and controls are needed to identify additional genes that confer moderate risk of ASD through rare, inherited variants.

NOTE: This preprint reports new research that has not been certified by peer review and should not be used to guide clinical practice.
Introduction

Autism spectrum disorder (ASD) is a neurodevelopmental condition characterized by impaired social communication and repetitive behaviors\textsuperscript{1}. Previous studies in ASD utilized family-based designs to focus on \textit{de novo} variants (DNVs) identified from parent-offspring trios\textsuperscript{2-8}. Over one-hundred high confidence ASD genes enriched with likely deleterious DNVs have been identified\textsuperscript{8}, most of which are also enriched for DNVs in other neurodevelopmental disorders (NDDs)\textsuperscript{9-11}. Statistical modeling suggests there are \textasciitilde1000 genes with DNV variants in ASD\textsuperscript{12,13}. However, despite the large effect size of individual pathogenic DNVs, all DNVs together only explain \textasciitilde2\% of variance in liability for ASD\textsuperscript{8,14}.

On the other hand, ASD is highly heritable (estimated heritability over 0.5)\textsuperscript{14-16}. Previous studies estimated that common variants explain up to half of the heritability\textsuperscript{14}, although only five genome-wide significant loci have been identified\textsuperscript{17}. The role of inherited coding variants has been evaluated using familial segregation of loss-of-function (LoF) variants (stop-gain, splice site and frameshift variants) carried by parents without ASD diagnoses or intellectual disability. Rare LoF variants only in genes intolerant of variation\textsuperscript{9,18} are over-transmitted to probands compared with siblings without ASD\textsuperscript{7,8,19-22}. However, identification of the individual risk genes enriched by such inherited variants has remained elusive.

We have created a large longitudinal research cohort, SPARK (SPARKForAutism.org\textsuperscript{23}) to advance research on the genetic, behavioral, and clinical features associated with ASD. SPARK represents the largest ASD cohort in the world, with over 100,000 individuals with ASD enrolled.

Rare, LoF variants are enriched in developmental disorders including ASD\textsuperscript{22,24}, but LoF variants in the general population are also enriched for sequencing and annotation artefacts\textsuperscript{25}, which present technical challenges in large sequencing studies. Methods to distinguish between high and low confidence LoF variants\textsuperscript{18,26,27} have been used to quantify gene level LoF intolerance\textsuperscript{18,26,28,29} and to refine the role of \textit{de novo} LoF variants in NDDs\textsuperscript{20}.

Here we present an integrated analysis of \textit{de novo} and inherited coding variants in over 42,607 ASD cases, including cases from previously published ASD cohorts and 35,130 new cases from SPARK. To our knowledge, this analysis is the largest sequencing study of ASD to date. In our two-stage design, we first characterized the contribution of DNVs and rare inherited LoF variants to ASD risk. Results from the first stage informed the second stage, in which we conducted a meta-analysis of 404 genes. By combining evidence from DNVs, over-transmission, and case-control comparison, we identified 60 ASD risk genes with exome-wide significance, including five new genes not previously implicated in neurodevelopmental conditions. Finally, we estimated the effect sizes of known and newly significant genes and used them for power calculations to inform the design of future studies.
Results
Overview of data and workflow
We aggregated exome or whole genome sequencing (WGS) data of 35,130 new cases from the SPARK study and 7,665 cases from published ASD studies (ASC\textsuperscript{3,8}, MSSNG\textsuperscript{6}, and SSC\textsuperscript{2,30}) (Supplementary Table S1) and performed a two-stage analysis (Figure 1). In the first stage, we analyzed de novo coding variants (DNVs) in 16,877 ASD trios and assessed transmission of rare LoF variants in 20,491 parents without ASD diagnoses or intellectual disability to offspring with ASD (including 9,504 trios and 2,966 single-parent-proband duos). For DNVs, we characterized the enrichment pattern in known and candidate risk genes, mutation intolerance (ExAC pLI\textsuperscript{18} and gnomAD metrics\textsuperscript{26}) and performed gene-based burden tests of de novo LoF and missense variants by DeNovoWest\textsuperscript{11}. For rare inherited LoFs, we estimated the over-transmission from parents without an ASD diagnosis to ASD cases in all genes and gene sets predefined by functional genomic data or results from DNV analysis. Based on DNV enrichment and over-transmission patterns in gene sets, we selected 404 genes for meta-analysis in stage 2 utilizing 22,764 new cases with exome or WGS data. In stage 2, we applied DeNovoWEST on DNVs, conducted transmission-disequilibrium tests on inherited LoFs in trios or duos, performed burden tests on rare LoFs in cases compared with population controls (104,068 subjects from gnomAD exome, non-neuro subset v2.1.1 and 132,345 TOPMed subjects), and combined the p-values to estimate a final p-value for each of the 404 genes. Finally, we performed a mega-analysis of rare LoFs in all cases and controls to estimate the effect sizes of known or new candidate ASD genes to inform future studies.

Known ASD or NDD risk genes explain two-thirds of population attributable risk of de novo coding variants in ASD
In the first stage, we combined data from four large-scale ASD cohorts, resulting in 16,877 unique ASD trios and 5,764 unaffected trios (Supplementary Table S1). The four cohorts show similar exome-wide burden of DNVs in simplex families. The burden of de novo LoF variants in cases with a family history of ASD is significantly lower than those without a reported family history (p=1.1e-4 by Poisson test), whereas the burden of predicted de novo damaging missense (D-mis, defined by REVEL score\textsuperscript{31}>=0.5) and synonymous variants are similar (Supplementary Figure S1).

Compared to unaffected offspring, the excess of damaging DNVs (de novo LoF and D-mis variants) in individuals with ASD is concentrated in LoF-intolerant genes, defined as genes with a probability of being LoF intolerant (pLI)\textsuperscript{18} >=0.5 in the Exome Aggregation Consortium (ExAC). Using LoF observed/expected upper-bound fraction (LOEUF), a recently developed gene constraint metric\textsuperscript{26}, the burden of damaging DNVs is highest among genes ranked in the top 20% of LOEUF scores (Figure 2A). Overall, the population attributable risk (PAR) from damaging DNVs is about 10%. We assembled 618 previously established dominant (“known”) ASD or NDD risk genes (Supplementary Table S2). These genes explained about 2/3 of the PAR from damaging DNVs. Excluding these genes, the fold enrichment of damaging DNVs was greatly attenuated (Figure 2A).
To assess the evidence of DNVs in individual genes, we applied DeNovoWEST, which integrates DNV enrichment with clustering of missense variants in each gene. The initial DeNovoWEST scan of DNVs in 16,877 ASD trios identified 159 genes with p<0.001 (Supplementary Table S3).

**Rare inherited LoF variants contribute to ASD risk mostly through unknown risk genes**

To analyze the contribution of rare inherited LoF variants to ASD risk, we evaluated transmission disequilibrium in ultra-rare (allele frequency < 1e-5) high-confidence (by LOFTEE and pext; see Methods and Supplementary Note) LoF variants from parents without ASD diagnoses or intellectual disability to affected offspring with ASD in 9,504 trios and 2,966 duos from the first stage (Supplementary Table S4). For a given set of genes, we quantified transmission disequilibrium using the number of over-transmitted (excess in transmission over non-transmission) LoF variants per trio; parent-offspring duos were considered half-trios.

Among autosomal genes, the overall transmission disequilibrium signal of ultra-rare LoF variants is enriched in LoF intolerant genes (ExAC pLI>=0.5) and in genes within the top 20% of LOEUF scores (Figure 2B), similar to the burden of damaging DNVs. We observed both over-transmission to affected and under-transmission to unaffected offspring, especially in genes within the top 10% of LOEUF scores. However, known ASD/NDD genes only explain ~20% of over-transmission of LoF variants to affected offspring (Figure 2B). On the X chromosome, we only considered transmission from mothers without ASD diagnoses to 9,883 affected sons and 2,571 affected daughters (Supplementary Table S4). Rare LoF variants in mothers without ASD diagnoses only show significant over-transmission to affected sons but not affected daughters and remain significant after removing known ASD/NDD genes (Supplementary Figure S2).

Together, these data suggest that most genes conferring inherited ASD risk are yet to be identified. Autosomal rare D-mis variants also show evidence of transmission disequilibrium to affected offspring, although the signal is much weaker and dependent on gene set, D-mis prediction method, pExt and allele frequency filters (Supplementary Figure S3).

To characterize the properties of genes contributing to ASD risk through rare inherited variants, we defined 25 gene sets from five categories representing both functional and genetic evidence relevant to ASD (Supplementary Table S5 and Supplementary Figure S4). We limited the genes to 5,754 autosomal constrained genes (ExAC pLI>=0.5 or top 20% of LOEUF scores) and performed TDT (Supplementary Table S6). For each gene set, we tested if high-confidence rare LoF variants show a higher frequency of transmission to ASD offspring than the remaining genes in the overall constrained gene set. As a comparison with DNVs, we also tested if the same set of genes are more frequently disrupted by damaging DNVs than the rest of the genes in ASD trios using the framework of dnEnrich.

We first considered functional gene sets derived from the neuronal transcriptome, proteome, or regulome. We confirmed significant enrichment in damaging DNVs (p<0.005 by simulation) in the gene sets that were previously suggested to be enriched for ASD risk genes including expression module M2/3, RBFOX1/3 targets, FMRP targets, and CHD8 targets. However, this enrichment can be largely explained by known ASD/NDD genes (Supplementary Figure S5). For ultra-rare inherited LoF variants, we found the proportion of transmission to ASD individuals in most functional gene sets is close to all genes in the background; only RBFOX
targets show a weak enrichment but can be largely explained by known genes (Figure 3). We also applied two recently developed machine learning methods to prioritize ASD risk genes: forecASD\textsuperscript{37} that integrates brain expression, gene network, and other gene level metrics, and A-risk\textsuperscript{38} that uses cell-type specific expression signatures in developing brain. Although enrichment of DNVs in genes predicted by these methods are mainly explained by known genes, genes prioritized by A-risk are significantly enriched with inherited LoFs that cannot be explained by known genes. Using A-risk\textgtr=0.4 (recommended threshold), 30\% of constrained genes (n=1,464) were prioritized and explain 64\% of the over-transmission of LoF variants to ASD offspring (p=2.6e-5 by chi-squared test). The enrichment is even higher than genes prioritized by the LOEUF score: 33\% of genes (N=1,777) in the top decile of LOEUF account for 55\% over-transmission (p=3.5e-4 by chi-squared test) (Figure 3).

We also considered gene sets that have evidence of genetic association with DNVs. Genes nominally enriched by DNVs (p<0.01 by DeNovoWEST; N=300) in ASD from the current study have a significantly higher over-transmission rate than other constrained genes (Odds ratio=1.39, p=3.0e-5 by chi-squared test) (Figure 3), although these genes only account for 21\% of the over-transmission. Genes nominally enriched by DNVs in other NDDs\textsuperscript{11} are also significantly enriched by DNVs in ASD and weakly enriched by inherited LoFs in ASD; however, both can be largely explained by known genes (Figure 3). This suggests that a subset of ASD genes increase risk by both de novo and inherited variants, and new genes can be identified by integrating evidence from DNV enrichment and TDT.

**DNVs and a subset of rare inherited LoFs are associated with cognitive impairment**

To evaluate the association of genotypes with phenotype in ASD, we used self-reported cognitive impairment in SPARK, a Vineland score of <70 in the SSC or the presence of intellectual disability in ASC. Damaging DNVs in genes ranked within the top 10\% of LOEUF scores show a higher burden (p=1.1e-24, by chi-squared test) in ASD cases with evidence of cognitive impairment than other cases, consistent with previous results\textsuperscript{2,8} (Figure 4A). Once known ASD/NDD genes were excluded, the residual burden of damaging DNVs in genes at the top 10\% LOEUF is greatly reduced and not significantly associated with cognitive phenotype in ASD (Figure 4A). Over-transmission of rare LOFs in genes within the top 10\% of LOEUF genes to ASD cases with cognitive impairment is about 2.7 times higher than to the cases without cognitive impairment (p=4.6e-3 by chi-squared test) and is still 2x higher (p=0.04 by chi-squared test) once known ASD/NDD genes were excluded (Figure 4B). However, rare LoFs in genes prioritized by A-risk, in which there is significant over-transmission to all cases overall, are not associated with cognitive impairment (Supplementary Figure S6). Taken together, these results suggest that rare variants in the top 10\% of LOEUF genes—most of which are already known to be ASD/NDD risk genes—are associated with cognitive impairment. However, a subset of rare, inherited variants, particularly those prioritized by A-risk, are not associated with cognitive impairment.
Meta-analysis of de novo and rare inherited LoF variants identifies 5 new risk genes with exome-wide significance

Based on results from the first stage of analysis, 404 genes showed plausible evidence of contributing to ASD risk, including: 1) 260 genes with evidence of TDT (TDT statistic \(39>1\)) and in gene sets enriched with rare inherited LoFs (top 10% LOEUF or within top 20% LOEUF and A-risk \(>0.4\)) (Supplementary Table S6) and 2) 159 genes with \(p<0.001\) from the DeNovoWEST analysis of DNVs (with 15 genes by both) (Supplementary Table S3). We performed a meta-analysis on the 367 autosomal genes with all data from Stage 1 and Stage 2, which includes 6,174 new ASD trios, 1,942 new duos, 15,780 unrelated cases (see Methods), and 236,000 population controls.

In the meta-analysis, we used Fisher’s method\(^{40}\) to combine 3 p-values that estimate independent evidence of DNVs, TDT, and case-control comparison: (1) DeNovoWEST with DNVs from both Stage 1 and 2 (\(n=23,039\) trios, Supplementary Table S1) using the parameters estimated in Stage 1, (2) TDT with rare LoF variants in parents without ASD diagnoses or intellectual disability with affected offspring in 15,586 trios and 4,907 duos (Supplementary Table S4), and (3) unrelated cases (Supplementary Table S7) compared to population controls using a binomial test. We used two sets of controls: gnomAD exome v2.1.1 non-neuro subset (\(n=104,068\)) and TOPMed WGS (freeze 8, \(n=132,345\)). We performed a case-control burden test using the two sets separately and input the larger p-value for the Fisher’s method. This approach avoids any sample overlap and provides sensitivity analysis to ensure that significant genes are not dependent on the choice of population reference. Although population reference data were processed by different bioinformatics pipelines, the cumulative allele frequencies (CAF\( s\)) of high-confidence (HC, see Methods) LoF variants are similar between internal pseudo-controls (see Methods) and the two population references after applying the same LoF filters (Supplementary Figure S7). Previous population genetic simulations predict that for genes under moderate to strong selection (selection coefficient>0.001), deleterious variants are expected to arise within 1,000 generations and population demographic histories do not confound the CAF\( s\) of deleterious alleles in these genes\(^{41}\). For 367 selected autosomal genes, the point estimates of selection coefficient under mutation-selection balance model\(^{42}\) are all greater than 0.01 (Supplementary Figure S8). Consistent with the theoretical predictions, most HC LoF variants in these genes are ultra-rare (Supplementary Figure S9) and the CAF\( s\) of HC LoF variants in European and non-European population samples are highly correlated (Supplementary Figure S10). Thus, we included population samples across all ancestries as controls. To make use of all genetic data collected, we also included rare variants of unknown inheritance from autism cases that were analyzed in the first stage. These variants come from cases that are part of parent-autism duos; such variants were either inherited from the parent not participating in the study or occurred de novo. Therefore, these data represent data independent of the transmission disequilibrium testing, even though the same cases were included in TDT.

We identified 60 genes with exome-wide significance (\(p<2.5e^{-6}\)). Figure 5 summarizes the distribution of LoF variants (with different modes of inheritance) in genes that reached experimental-wide significance by DNV enrichment (Figures 5A) and other significant genes by meta-analysis (Figure 5B, Supplementary Figure S11). Genes that are significant only in meta-
analysis tend to harbor more inherited LoF variants than de novo variants, consistent with their lower penetrance for ASD or NDD.

Although most significant genes were previously known, we identified five new genes that are exome-wide significant regardless of the choice of population reference: NAV3, MARK2, ITSN1, SCAF1, and HNRNPU2 (Table 1). As expected, most supporting variants are ultra-rare, and results are robust to the allele frequency filter. These five new genes together explain 0.27% population attributable risk ratio (PAR) (Supplementary Table S8). NAV3 has a similar PAR as CHD8 and SCN2A (~0.095%). ITSN1 is similar to PTEN (~0.065%).

The association of NAV3 with ASD risk is entirely driven by rare inherited variants (Table 1). NAV3 harbors a single HC de novo LoF variant in an unaffected sibling in the SSC and was previously included in the negative training set by A-risk. Despite this, NAV3 still has a high A-risk score, suggesting NAV3’s expression pattern is highly similar to known ASD genes (Supplementary Data 1). NAV3 has high expression in inner cortical plate of developing cortex, and in pyramidal neurons (hippocampus CA1 and somatosensory cortex) and cortical interneurons, consistent with the signatures of known ASD genes (Supplementary Figure S12).

The association of MARK2 with ASD risk is primarily driven by DNVs. MARK2 is also associated with other NDDs (P=2.7e-5 by DeNovoWEST) including Tourette syndrome and epilepsy. We find that 3/8 of autistic offspring with variants in MARK2 report epilepsy, 2/8 report Tourette syndrome and 7/8 have evidence of cognitive impairment (Supplementary Table S9).

The remaining three novel genes have support from both DNVs and rare LoFs. Two genes have suggestive evidence from other NDD studies. ITSN1 and SCAF1 shows nominal significance of DNV enrichment in 31,058 NDD trios (P<0.05 by DeNovoWEST). SCAF1 was among the top 50 genes from gene-based burden test in a recent schizophrenia case-control study (P=0.0027 by burden test). Both ITSN1 and NAV3 have moderate effect sizes (point estimate of relative risk 3~6, Supplementary Table S8). ITSN1 has been highlighted in our previous study with evidence of enriched inherited LoFs. ITSN1 and NAV3 also show increased CAF of LoF variants in a recent study by ASC although the association was not significant. We also assessed deletions in these new genes. For both ITSN1 and NAV3, we identified four partial or whole gene deletions in 33,083 parents without ASD diagnoses or intellectual disability that also show transmission disequilibrium to affected offspring (Supplementary Figure S13).

While both de novo and rare inherited LoFs in the most constrained genes are strongly associated with intellectual disability (ID) in ASD (Figure 4), the association of such variants in individual genes is heterogenous, as suggested by the lack of association of rare inherited variants in genes with high A-risk (Supplementary Figure S5). We calculated the burden of cognitive impairment (see Methods) in 87 ASD individuals with HC LoF variants in the four novel moderate risk genes and compared it to 129 individuals with HC LoF in the well-established ASD risk genes CHD8, SCN2A, SHANK3, ADNP and FOXP1 as well as 8,731 individuals with ASD in SPARK (Supplementary Figure S14). Although most individuals with variants in well-established ASD risk genes have some evidence of cognitive impairment (88%,) individuals with LoF variants in the moderate risk genes had significantly lower burden (56%, p=4.5e-7 by chi-squared test). Individuals with HC LOFs in the moderate risk genes did not have a significantly different
burden of cognitive impairment than 8,731 individuals with ASD in SPARK (56% vs. 50%, p = n.s.). Individuals with LoF variants in the moderate risk genes also had a similar male: female (4:1) ratio compared to the larger cohort whereas individuals with variants in the well-established ASD risk genes showed significantly less male bias (1.6: 1, p = 0.009 by chi-squared test) (Supplementary Figure S14), as previously reported. We also predicted full-scale IQ on all participants based on parent-reported data using a machine learning method. Carriers of rare LoFs in three (NAV3, SCAF1, and HNRNPUL2) of the four new genes with substantial contribution from rare inherited variants have similar IQ distribution as the overall SPARK cohort (Figure 6A), which is substantially higher than heterozygotes with rare LoFs in well-established, highly-penetrant genes that contribute to ASD primarily through de novo variants (“DN genes”), such as CHD8, SHANK3, and SCN2A. In fact, both novel and established genes with significant contribution from rare inherited LoFs are less associated with ID than DN genes (Figure 6B). Across these genes, there is a significant negative correlation (r=0.78, p=0.001) of estimated relative risk of rare LoFs with average predicted IQ of the individuals with these variants (Figure 6C). These genes could be associated with other neurobehavioral phenotypes.

Most known ASD/NDD genes that are enriched by de novo LoF variant harbor more de novo than inherited HC LoF variants in ~16,000 unrelated ASD trios (Figure 5A and Supplementary Figure S15), consistent with their high penetrance for ASD/NDD phenotypes and strong negative selection. Using population exome or WGS data, we calculated a point estimate of selection coefficient ($\hat{s}$) of LoFs in each gene (Supplementary Table S8) and found that the fraction of de novo LoFs in ASD genes is higher in genes with large $\hat{s}$, and smaller in genes with small $\hat{s}$ (Supplementary Figure S7B), consistent with population genetic theory. We also estimated average effect size of rare LoFs in ASD genes by comparing cumulative allele frequency (CAF) in 31,976 unrelated cases and population exome or WGS data. As expected, known and newly significant ASD genes with higher risk to ASD are under stronger selection (larger $\hat{s}$) (Supplementary Figure S16).

Functional similarity of new genes with known ASD genes

To better appreciate the probable functional implications of the new exome-wide significant genes that confer inherited risk for ASD, we integrated mechanistic (STRING) and phenotypic (HPO) data into a single embedding space (six dimensions, one for each archetype coefficient) using a combination of canonical correlation analysis and archetypal analysis. This embedding space serves as an interpretive framework for putative ASD risk genes (N=1,776). Six functional/phenotypic archetypes were identified (Figure 7) that represent pathways that are well-understood to play a role in ASD: neurotransmission (archetype 1 or A1), chromatin modification (archetype 2 or A2), RNA processing (archetype 3 or A3), membrane trafficking and protein transport (archetype 4 or A4), extracellular matrix, motility, and response to signal (archetype 5 or A5), and KRAB domain and leucine-rich region proteins (archetype 6 or A6), also enriched for intermediate filaments. These archetypes organize risk genes in a way that jointly maximizes their association with mechanisms (STRING clusters) and phenotypes (HPO terms). For instance, A1 genes (neurotransmission) are enriched for the STRING cluster CNR1 (ion channel and neuronal system) and are also associated with seizure and epileptic phenotypes. A2 genes (chromatin modifiers) are enriched for nuclear factors and genes linked to growth and
morphological phenotypes (Supplementary Table S10). We call genes that strongly map to an archetype (i.e., > 2x the next highest-ranking archetype) “archetypal” and “mixed” if this criterion is not met (see methods). Archetypal genes are generally less functionally ambiguous than “mixed” genes. Of the five novel inherited risk genes, two are archetypal (suggesting function within known risk mechanisms): NAV3 (A6: KRAB domain & LRR) and ITSN1 (A4: membrane trafficking and protein transport). SCAF1, MARK2, and HNRNPUL2 are mixtures of the identified archetypes, largely A4 and A5. That these new genes did not resolve clearly into archetypes (that were defined by known and suspected autism risk genes) suggests that they may operate in potentially novel or under-appreciated mechanisms. To elucidate these possibilities, we constructed an ad hoc “archetype,” defined by the centroid between SCAF1, MARK2, and HNRNPUL2 (see Figure 7C). Cell-cell junction (CL:6549) was the STRING cluster most associated with this centroid (p = 4.12 x 10^{-14} by the K-S test, Fig. 7D), which fits with its location between A4 (membrane trafficking) and A5 (ECM).

Power analysis

The power of identifying risk genes with rare or de novo variants monotonically increases with increasing effect size or expected CAF under the null. New ASD genes to be discovered are likely to have smaller effect size than known ASD genes, as suggested by our results. Additionally, known ASD genes are biased toward longer genes with higher background mutation rate of damaging variants (“long genes”) (Supplementary Figure S17). Even though longer genes are more likely to be expressed in brain and relevant to ASD/NDD, among most constrained genes, long genes (LoF mutation rate above 80% quantile) and short genes (below 80%) have similar enrichment of damaging de novo variants and rare inherited LoFs (Supplementary Figure S18). Notably, for small genes, known genes have virtually no contribution to over-transmitted HC LoFs to affected offspring (Supplementary Figure S18B). It suggests that many smaller genes contributing to ASD risk remain to be identified. We focus on the power of detecting new ASD genes with a moderate effect size and the full range of background mutation rate.

We use a published framework to analyze power based on case-control association of rare variants. For rare variants in genes under strong selection, CAF is largely determined by mutation rate and selection coefficient. We therefore modeled power of discovering risk genes as a function of relative risk and selection coefficient. With about 5,500 constrained genes, the power of the current study was calculated for 31,976 unrelated cases and experiment-wise error rate of 9e-6 (Supplementary Figure S19).

We inversed the power calculation to determine required sample size to achieve 90% power under the same assumptions (Supplementary Figure S20). For genes at median LoF mutation rate across all genes, we estimated that it requires about 96,000 cases (three times the current sample size) to identify genes with similar effect size as NAV3 (RR=4.5) and ITSN1 (RR=5), about 64,000 (twice the current sample size) to find genes with similar effect sizes as SCAF1 (RR=8) and HNRNPUL2 (RR=9). We note that it requires 10 and 5 times the current sample size to detect these types of genes by de novo variants alone.
Discussion

In this study, we assembled the largest sequencing data set of individuals with ASD to date, including 35,130 ASD cases and their family members collected by SPARK. We characterized the contribution of rare inherited variants to ASD risk and identified five new ASD risk genes by both de novo and rare inherited coding variants. We identified rare LoF variants in new ASD risk genes with modest effect size that are not strongly associated with ID. This finding represents a difference in phenotypic association with ID compared with other well-established, highly penetrant ASD genes. To find new risk genes with relative risks of 2-5 (comparable to the low relative risk genes from this study: NAV3 and ITSN1) in the 50-percentile for gene-wide LoF mutation rate (2e-6) and the 50-percentile for selection among known risk genes (0.2), our power analysis suggests that 52,000, 73,000, 116,000 or 227,000 total ASD cases are necessary, respectively (cf. eq 1 from power calculation in Supplementary material). Larger ASD cohorts with phenotypic data will be necessary to identify new ASD risk genes and may help to understand the biology of core symptoms of ASD in individuals without ID.

Our results suggest that identification of new risk genes with rare inherited variants can substantially improve genetic diagnostic yield. We found that rare inherited LoF variants account for 6% of PAR, similar to de novo LoF variants. Over two thirds of the PAR from de novo coding variants are explained by known ASD or NDD genes. In contrast, less than 20% of PAR from rare inherited LoFs variants is explained by known genes, suggesting most genes contributing to ASD risk through rare inherited variants are yet to be discovered. These unknown risk genes are still largely constrained to LoFs in the general population and/or have similar expression profiles in developing brains to known ASD risk genes. Combining evidence from both de novo and rare inherited variants, we identified 60 genes associated with ASD with exome-wide significance, including five novel genes. Rare LoFs in these five new genes account for a PAR of 0.27%, about half of the PAR of the 5 most common highly penetrant ASD genes (KDM5B, GIGYF1, CHD8, SCN2A, SHANK3).

NAV3, to our knowledge, is the first autosomal ASD risk gene discovered by association of solely rare inherited variants. Carriers of rare LoFs in NAV3 have an average predicted IQ of 81, slightly above the SPARK cohort average (79). The prevalence of ID among NAV3 heterozygotes is similar to the SPARK cohort average. This is distinctly different from established ASD risk genes (e.g., CHD8, SHANK3, SCN2A), nearly all identified by highly penetrant de novo variants, associated with ID in ASD cohorts\(^2\). The absence of ID is also observed in other genes (e.g., SCAF1, HNRNPUL2, GIGYF1, KDM5B, KMT2C) with substantial contribution from rare inherited variants and modest effect size. Nevertheless, the data show that variants in these new ASD genes have effects on core symptoms of ASD, cognition, and other behaviors including schizophrenia, Tourette syndrome, ADHD and other behavioral conditions. Detailed phenotyping of individuals carrying these rare inherited variants is needed to understand the phenotypic effects of each gene. Such strategies should include a genetic and phenotypic assessment of family members who also carry the rare variant but may not have an ASD diagnosis. Since all individuals consented in SPARK are re-contactable, such studies will enable a more complete picture of the broad phenotypic effects of these variants without the bias of clinical ascertainment. Overall, these risk genes with modest effect size may represent a
different class of ASD genes that are more directly associated with core symptoms of ASD
and/or neuropsychiatric conditions rather than global brain developmental and ID.

The approaches employed in this study made full use of rare variation, and this analytical
method is generalizable to many conditions. In particular, the multiple methods used to reduce
noise in LoF alleles present in control samples were particularly effective in assessing the signal
within the novel genes of moderate effect. We also leveraged gene expression profiles
informed by machine learning methods to help prioritize genes for the meta-analysis stage of
our analysis. Future studies that leverage additional multi-omic data such as dGTEx may
further improve signal to noise.

Our archetypal analysis provides some clues as to the potential risk mechanisms of the five
newly identified risk genes. *ITSN1* was unambiguously mapped to A4: membrane trafficking and
protein transport and has a role in coordinating endocytic membrane traffic with the actin
cytoskeleton. *NAV3* (A6: KRAB domain and LRR), is associated with both axon guidance and
malignant growth and invasion and is thought to regulate cytoskeletal dynamics. Indeed, A6 is
enriched for processes related to intermediate filaments (Supplementary Table S10) a known
determinant of cell motility and polarity. Although *MARK2, SCAF1*, and *HNRNPUL2* were not
identified as archetypal (potentially suggesting divergence from well-known autism risk
mechanisms) a search for functional enrichment of this interstitial region between A4 and A5
found that their roles in developmental risk may be most relevant at the cell-cell junction,
particularly as it relates to migration (see Figure 7D).

Taken together, our results suggest that a continued focus on *de novo* variants for ASD gene-
discovery may yield diminishing returns. By contrast, studies designed to identify genomic risk
from rare and common inherited variants will not only yield new mechanistic insight but help
explain the high heritability of ASD. SPARK is designed to recruit individuals across the autism
spectrum, without relying on ascertainment at medical centers. As a result, SPARK may be
better suited to identify genes with transmitted variants that have lower penetrance and to
identify the genetic contributions to the full spectrum of autism. The strategies employed by
SPARK — to recruit and assess large numbers of individuals with autism across the spectrum
and their available family members without costly, in-depth clinical phenotyping — is necessary
to achieve the required sample size to fully elucidate genetic contributions to ASD. SPARK’s
ability to recontact and follow all participants will also be critical to deeply assess the
phenotypes associated with the newly discovered genes and to develop and test novel
treatments.
| Gene   | Prioritization | Enrichment of de novo damaging variants | Transmission disequilibrium of HC LoFs | case-control comparison of HC LoF rate |
|--------|----------------|-----------------------------------------|--------------------------------------|---------------------------------------|
|        |                | dnLoF | μ\text{LoF} | dnDmis | μ\text{Dmis} | P\text{DNV} | Trans: Non-Trans to affected | P\text{TDT} | Number (rate) of LoFs in cases | Rate of LoFs in controls: gnomAD exome, TOPMed | P\text{CC} | P\text{meta} |
| NAV3   | TDT            | 1     | 1.1e-5    | 1       | 1.1e-5     | 0.23      | 17        | 17:2       | 3.6e-4  | 22 (1.4e-3)       | 3e-4, 2.6e-4 | 4.4e-7, 2.1e-8 | 1.2e-8 |
| MARK2  | De novo        | 5     | 4.4e-6    | 3       | 4.8e-6     | 8.9e-9    | 3         | 3:1        | 0.31    | 4 (2.5e-4)       | 2e-5, 6e-5  | 4.5e-3, 0.03 | 2.3e-8 |
| SCAF1  | TDT            | 2     | 4.8e-6    | 0       | 1.7e-7     | 1.3e-3    | 4         | 3:1        | 0.31    | 13 (8.2e-4)      | 3e-5, 7e-5  | 2.1e-6, 1.4e-6 | 2.1e-7 |
| ITSN1  | TDT            | 3     | 1.2e-5    | 2       | 1.3e-5     | 2.6e-3    | 18        | 17:2       | 3.6e-4  | 10 (6.3e-4)      | 1.6e-4, 2e-4 | 2e-3, 4e-3  | 4.3e-7 |
| HNRNPUL2 | De novo     | 3     | 5.8e-6    | 0       | 3.8e-6     | 1.8e-3    | 2         | 2:0        | 0.25    | 10 (6.3e-4)      | 4e-5, 5e-5  | 2.6e-6, 8.2e-7 | 2.7e-7 |

Table 1: Statistical evidence for the five novel exome-wide significant ASD risk genes identified in this study.

Control HC LoF rates are estimated from two population-based reference panels: gnomAD exome (v2.1.1, non-neuro subset, 104,068 individuals), and TopMed (freeze 8, 132,345 individuals). Meta-analysis is done by combining p-values from de novo, TDT and pseudo case-control analysis using Fisher’s method. For pseudo case-control, we conservatively took the largest p-value for meta-analysis. P\text{DNV}: One-sided p-value for enrichment of all DNVs in 23,053 ASD trios, P\text{TDT}: One-sided p-value of over-transmission of HC LoFs to affected offspring in 28,556 trios and 4,526 duos, P\text{CC}: One-side p-value for increased HC LoF rate in 15,811 unrelated cases compared with population controls (showing two p-values from comparison with gnomAD exome and TOPMed data respectively).
**Figure 1. Analysis workflow.** In the discovery stage, we identified *de novo* variants in 16,877 ASD trios and rare LoF variants in 20,491 parents without ASD diagnoses and intellectual disability. We compared properties of *de novo* and rare variants to identify rare LoFs that contribute to genetic risk in individuals with ASD. We also evaluated their associations with cognitive impairment and enriched gene sets. We performed an initial exome-wide scan of genes enriched by *de novo* variants or showing transmission disequilibrium (TD) of rare LoFs to affected offspring and selected a total of 404 genes for further replication, including 159 *de novo* enriched genes and 260 prioritized TD genes from enriched gene sets (15 genes were in both). In the meta-analysis stage, we first evaluated evidence from *de novo* enrichment and TD of rare, inherited LoFs in an expanded set of family-based samples including over 6,000 additional ASD trios and around 2000 additional duos. The *de novo* variants in ASD were combined with those from additional 31,565 NDD trios to refine the filters of high confidence (HC) LoFs in *de novo* enriched genes. We also constructed an independent dataset of LoF variants of unknown inheritance from 15,780 cases that were not used in *de novo* or transmission analysis. We compared LoF rates in cases with two population-based sets of controls (n ~104,000 and ~132,000, respectively). For 367 LoF intolerant genes on autosomes, the final gene level evidence was obtained by meta-analyzing p-values of *de novo* enrichment, TD of HC rare, inherited LoFs, and comparison of HC LoFs from cases and controls not used in the *de novo* or transmission analysis. We also performed a mega-analysis that analyzed HC LoFs identified in all 31,976 unrelated ASD cases and compared their rates with population-based controls.
Figure 2. Comparison of burden between de novo damaging variants and rare, inherited LoFs in ASD. (A) The burden of de novo variants was evaluated by the rate ratio and rate difference between 16,877 ASD and 5,764 unaffected trios. The exome-wide burden of de novo LoF and Dmis (REVEL>=0.5) variants are concentrated in constrained genes (ExAC pLI>=0.5) and in genes with the highest levels of LoF-intolerance in the population—defined by the top two deciles of gnomAD LOEUF scores. Burden analysis was repeated after removing known ASD/NDD genes. The number of genes before and after removing known genes in each constraint bin is shown below the axis label. Among constrained genes (ExAC pLI>=0.5 or the top 20% of gnomAD LOEUF scores), close to two thirds of case-control rate differences of de novo LoF and Dmis variants can be explained by known genes. (B) The burden of inherited LoFs was evaluated by looking at the proportion of rare LoFs in 20,491 parents without ASD/NDD genes. The number of genes before and after removing known genes in each constraint bin is shown below the axis label. Among constrained genes (ExAC pLI>=0.5 or the top 20% of gnomAD LOEUF scores), close to two thirds of case-control rate differences of de novo LoF and Dmis variants can be explained by known genes. (B) The burden of inherited LoFs was evaluated by looking at the proportion of rare LoFs in 20,491 parents without ASD/NDD genes. The number of genes before and after removing known genes in each constraint bin is shown below the axis label. Among constrained genes (ExAC pLI>=0.5 or the top 20% of gnomAD LOEUF scores), close to two thirds of case-control rate differences of de novo LoF and Dmis variants can be explained by known genes.
Figure 3. Association of rare, inherited LoFs with cognitive impairment in ASD cases. Ultra-rare inherited LoFs with pExt>=0.1 in genes with the top 10% gnomAD LOEUF scores also show a higher proportion of transmission and a higher over-transmission rate to ASD offspring with cognitive impairment than those without. Rare LoFs in other constrained genes are not significantly associated with phenotypic severity. The increased burden of inherited LoFs in cases with cognitive impairment remains significant after removing known ASD/NDD genes.
Figure 4. Enrichment of rare LoF variants in ASD cases across gene sets. Gene sets were defined and grouped by transcriptome proteome, neuronal regulome, ASD gene prediction scores, genetic evidence from neuropsychiatric diseases, and gene level constraint. Analyses were repeated after removing known ASD/NDD genes. (Number of genes in each set before and after removing known genes are shown in bracket below gene set.) Dots represent fold enrichment of DNVs or odds ratios for over-transmission of LoFs in each set. Horizontal bars indicate the 95% confidence interval. For each gene set, we show the percentage of over-transmission of rare LoFs to cases.

Enrichment of rare, inherited LoFs was evaluated by comparing the transmission and non-transmission of ultra-rare LoFs with pExt>=0.1 in the gene set versus those in all other constrained genes using a 2-by-2 table. P-values were given using the chi-squared test.
Figure 5. Distribution of *de novo* and inherited LoF variants in known and novel ASD genes in cases and population controls. From left to right: pyramid plots summarizing the number of *de novo* LoF variants in 15,857 ASD trios, inherited HC LoFs in 18,720 unrelated offspring included in transmission analysis, and HC LoFs in 15,780 unrelated cases; bar plot of transmission vs. non-transmission for rare HC LoFs identified in parents without ASD diagnoses or intellectual disability; three plots comparing the HC LoF rate in 31,976 unrelated ASD cases with gnomAD exomes (non-neuro subset, 104,068 individuals). Horizontal bars indicate standard errors. (A) The upper panel shows 28 known ASD/NDD genes in which LOEUF scores are in the top 30% of gnomAD, have a p-value for enrichment among all DNVs (p <9e-6) in 23,039 ASD trios, and have more than 10 LoFs. (B) The lower panel shows 9 additional ASD risk genes that achieved a p-value of <9e-6 in Stage 2 of this analysis. The majority of genes in the lower panels harbor more inherited LoFs than *de novo* variants. All five novel genes (*Error! Reference source not found.*) are shown in the lower panel. Note that the x-axes of LoF rates are in the squared root scale.
Figure 6. Predicted full-scale IQ (FSIQ) in individuals with pathogenic variants in inherited or de novo genes in SPARK. We examined the distribution of predicted IQ by a machine learning method for individuals with ASD with a LoF mutation in one of the five novel exome-wide significant genes (MARK2, NAV3, ITSN1, SCAF1, HNRNPUL2) and nine known ASD genes (CHD8, SHANK3, SCN2A, ADNP, ARID1B, FOXP1, KDM5B, GIGYF1, KMT2C), compared with 2,545 SPARK participants with ASD and known IQ scores. We denote the genes contributing to ASD primarily through de novo LoF variants in our analysis as “De novo” (in red), and the genes primarily through inherited LoF variants as “Inherited” (in blue). (A) Distribution of predicted IQ between individuals with ASD with LoF mutations in the five novel genes, 9 known genes and all participants with ASD and known IQ scores in SPARK (n=2,545). We compared the mean predicted IQ between participants with LoF mutations in ASD genes and all participants by two-sample t-test. Significance level is denoted by the star sign above each violin plot (*: 0.01 ≤ p<0.05, **: 0.001≤p<0.01, ***: p<0.001). Individuals with pathogenic variants in de novo risk genes have significantly lower predicted IQ than overall SPARK participants with ASD and known IQ scores, while individuals with LoF variants in moderate risk, inherited genes with show similar predicted IQ as the overall SPARK participants, with the exception of ITSN1. (B) Distribution of predicted IQ between individuals with ASD gene grouped by both inheritance status (“De novo” or “Inherited”) and whether the ASD genes are novel (“Novel” or “Known”). We compared the mean predicted IQ between individuals with pathogenic variants in “De novo” genes and “Inherited” genes among our five novel genes and nine known genes. Overall, people with LoF mutations in “De novo” genes have an average of 13-16 points lower predicted IQ than individuals with LoF mutations in “Inherited” genes, regardless of whether the ASD genes are novel or known. (C) Average relative risk of ASD and average predicted IQ among different groups. Each dot shows the average of individuals with rare LoFs of a gene selected in panel A. The relative risk is estimated from mega analysis and capped at 60. Pearson correlation between average IQ and log relative risk is -0.78 (p=0.001). The horizontal line represents the average IQ (IQ=79) of all SPARK individuals with predicted IQs. ITSN1 is an outlier at the bottom left corner.
Figure 7. Functional/phenotypic embedding of ASD risk genes. Using a combination of archetypal analysis and canonical correlation analysis, putative autism risk genes were organized into $k=6$ archetypes that represent distinct mechanistic (STRING) and phenotypic (HPO) categorizations (A; neurotransmission, chromatin modification, RNA processing, transport, extracellular matrix, motility and response to signal, and leucine-rich repeat/KRAB domain containing genes). Genes implicated by our meta-analysis are indicated by their label, with novel genes indicated in red. For each of the five novel genes, we identified the five nearest neighbors in the embedding space among the 62 meta-analysis genes (B). SCAF1, MARK2, and HNRNPUL2 were identified as “mixed” rather than “archetypal” in their probable risk mechanisms. To gain further insight into possible risk mechanisms, we calculated the embedding distance to the centroid of these three genes (C), which was then used as an index variable to perform gene set enrichment analysis. A STRING cluster (CL:6549) containing genes related to cell-cell junctions and the gap junction was identified as being highly localized in this region of the embedding space ($p = 4.12 \times 10^{-14}$ by the KS test) (D). This may suggest that these genes confer autism risk through dysregulation of processes related to cell adhesion and migration.
**Methods**

We performed an integrated analysis of coding variants in over 35,130 new ASD cases in SPARK and additional cases from previously published autism cohorts (ASC\textsuperscript{3,8}, MSSNG\textsuperscript{6}, and SSC\textsuperscript{2,30}), using a two-stage analysis workflow (0 1). In the first stage, we analyzed over 10,000 ASD cases from family-based samples and systematically compared damaging DNVs and rare, inherited LoF variants. Then we performed an exome-wide scan of genes enriched by DNVs in ASD cases and prioritized genes with suggestive evidence of DNV enrichment. We filtered for high-confidence (HC) LoF variants and searched for genes enriched by inherited HC LoFs using a transmission disequilibrium test (TDT)\textsuperscript{54}. In the second stage, we added 22,764 ASD cases and used meta-analysis to further assess the prioritized genes for enrichment of DNVs and TDT of HC LoFs. For LoF intolerant genes, we compared frequency of HC LoF variants in unrelated cases, population controls, and pseudo-controls in ASD families. Finally, we performed a case-control analysis of ASD cases vs population controls to estimate effect sizes for known and newly significant genes and used them for power calculations to estimate sample sizes needed for future studies.

**ASD Cohorts**

**SPARK**

We established SPARK (Simons Foundation Powering Autism Research for Knowledge) cohort to facilitate genotype driven research of ASD at scale\textsuperscript{23}. Eligibility criteria for SPARK study is residence in the United States and a professional diagnosis of ASD or a family member of a proband in SPARK. SPARK has recruited over 50,000 re-contactable families with ASD cases at 31 different clinical centers across the United States as well as through social and digital media. Individuals with known genetic diagnoses and individuals with and without a family history of autism are included. Whenever possible, parents and family members with or without autism were enrolled and included in the genetic analysis.

Saliva was collected using the OGD-500 kit (DNA Genotek) and DNA was extracted at PreventionGenetics (Marshfield, WI). The samples were processed with custom NEB/Kapa reagents, captured with the IDT xGen capture platform, and sequenced on the Illumina NovaSeq 6000 system using S2/S4 flow cells. Samples were sequenced to a minimum standard of >85% of targets covered at 20X. 97% of samples have at least 20x coverage in >95% of region (99% of samples — in 89% of regions). Pending sample availability, any sample with 20X coverage below 88% was re-processed and the sequencing events were merged to achieve sufficient coverage. The Illumina Infinium Global Screening Array v1.0 (654,027 SNPs) was used for genotyping. The average call rate is 98.5%. Less than 1% of samples have a call rate below 90%.

In the first stage of analysis, we included 28,649 SPARK individuals including 10,242 ASD cases from over 9,000 families with exome sequencing data that passed QC (Error! Reference source not found.). A subset of 1,379 individuals was part of the previously published pilot study\textsuperscript{7}. To replicate prioritized genes from the discovery stage, we performed a second stage analysis that included an additional 39,926 individuals with 16,970 ASD cases from over 20,000 families with...
exome or whole genome sequencing (WGS) data available after of the analysis in discovery cohort was completed. For new samples in this study, exome sequences were captured by IDT xGEN research panel and sequenced on the Illumina NovaSeq system. DNA samples were also genotyped for over 600K SNPs by Infinium Global Screening Array.

We used KING[^55] to calculate statistics for pairwise sample relatedness from genotypes of known biallelic SNPs, and validated participant-reported familial relationships (Supplementary Figure S21A-B). The relatedness analysis also identified cryptically related families that are connected by unreported parent-offspring or full sibling pairs. Pedigrees were reconstructed manually from inferred pairwise relationships and validated by PRIMUS[^56] and we used inferred pedigree for all analyses. Sample sex was validated by normalized sequencing depths or array signal intensities of X and Y chromosomes which also identified X and Y chromosome aneuploidies (Supplementary Figure S21C-D). To infer genetic ancestry, we first performed principal component (PC) analysis on SNP genotypes of non-admixed reference population samples from 1000 Genomes Projects[^57] (Africans, Europeans, East Asians and South Asians) and Human Genome Diversity Project[^58,59] (Native Americans), then projected SPARK samples onto PC axes defined by the five reference populations using EIGENSOFT[^60] (Supplementary Figure S22). The projected coordinates on first four PC axes were transformed into probabilities of five population ancestries using the method of SNPweights[^61]. The inferred ancestral probabilities show general concordance with self-reported ethnicities (Supplementary Figure S22B).

Samples were predicted from a reference population if the predicted probability was >=0.85.

The phenotypes of participants are based on self- or parent-report provided at enrollment and in a series of questionnaires from the Simons Foundation Autism Research Initiative database, SFARI Base. We used SFARI Base Version 4 for the discovery cohort and Version 5 for the replication cohort. In the discovery cohort, information about self-reported cognitive impairment (or intellectual disability/developmental delay) was available for 99.2% of ASD cases and 83.5% of other family members at recruitment or from the Basic Medical Screening Questionnaire available on SFARIbase. For phenotype-genotype analyses in individuals with variants in specific ASD risk genes, we defined an individual as having cognitive impairment if 1) there was self- or parent-report of cognitive impairment at registration or in the Basic Medical Screening Questionnaire, 2) the participant was at or over the age of 6 at registration and was reported to speak with less than full sentences or the participant was at or above age 4 at registration and reported as non-verbal at that time, 3) the parent reported that cognitive abilities were significantly below age level, 4) the reported IQ or the estimated cognitive age ratio (ratio IQ[^62,63] was <80 or 5) the parent reported unresolved regression in early childhood without language returning and the participant does not speak in full sentences. The continuous full-scale IQ was imputed based on a subset of 521 samples with full scale IQ and phenotypic features by the elastic net machine learning model[^48]. In a subset of cases for which full-scale IQ data or standardized Vineland adaptive behavior scores (version 3) was available, we found self-reported cognitive impairment shows higher correlation with Vineland score than full-scale IQ (Supplementary Figure S23). ASD cases with self-reported cognitive impairment were defined as Cognitively Impaired cases, and other cases as Not Cognitively Impaired cases. Other non-ASD family members were considered as unaffected if they were also not indicated to have cognitive impairment. In total of 18.5% families, proband has at least one first-degree
relative with ASD who was recruited in the study and/or reported by a family member. Those families were referred to as multiplex, and other families with only a single ASD individual as simplex. The majority (>85%) of affected relative pairs in multiplex families were siblings. Multiplex families have slightly lower male-to-female ratio and lower proportion of cognitive impairment among affected offspring (Supplementary Figure S24A-B). In comparison, only 1% of parents in the discovery cohort are affected of which two thirds are females and less than 3% have cognitive impairment (Supplementary Figure S24A-B). In addition, non-ASD family members in multiplex families show significantly higher frequency of self-reported cognitive impairment, learning/language disorders, other neuropsychiatric conditions, and other types of structural congenital anomalies (Supplementary Figure S24C). Non-ASD parents in multiplex families also have lower educational attainment (Supplementary Figure S24D).

SSC

SSC (Simon Simplex Collection) collected over 2,500 families with only one clinically confirmed ASD cases who have no other affected first or second degree relatives as an effort to identity de novo genetic risk variants for ASD. SSC data have been published before. Here we included 10,032 individuals including 2,633 cases with exome or WGS data available and passed QC. The data were reprocessed using the same pipeline as SPARK. For 91 trios that are not available or incomplete, we collected coding DNVs from published studies. In analysis to associate genetic variants with phenotype severity, we used standardized Vineland adaptive behavior score to group affected cases because it shows higher correlation than full-scale IQ with self-reported cognitive impairment in SPARK (Supplementary Figure S23). Cases with cognitive impairment in SSC were defined by Vineland score<=70, and cases with no cognitive impairment by score>70.

ASC

ASC (Autism Sequencing Consortium) is an international genomics consortium to integrate heterogenous ASD cohorts and sequencing data from over 30 different studies. Individual level genetic data are not available. So we included 4,433 published trios (4,082 affected and 351 unaffected) merged from two previous studies for DNV analysis. To define low and high functioning cases, we used binary indicator of intellectual disability which was available for 66% of cases. Families with multiple affected trios are considered multiplex, others are simplex.

MSSNG

The MSSNG initiative aims to generate WGS data and detailed phenotypic information of individuals with ASD and their families. It comprehensively samples families with different genetic characteristics in order to delineate the full spectrum of risk factors. We included 3,689 trios in DB6 release with whole genome DNV calls are available and passed QC in DNV analysis, of which 1,754 trios were published in the previous study. A total of 3,404 offspring with a confirmed clinical diagnosis of ASD were included as cases. Among individuals without a confirmed ASD diagnosis, 222 who did not show broader or atypical autistic phenotype or other developmental disorders were used as part of controls. Multiplex families were defined as families having multiple affected siblings in sequenced trios or in phenotype database. Information about cognitive impairment was not available at the time of analysis.
Variant calling and quality control

Supplementary Table S11 describes software version and parameter settings for each analysis below.

Data processing

Sequencing reads were mapped to human genome reference (hg38) using bwa-mem\(^7\) and stored in CRAM format\(^8\). Duplicated read pairs in the same sequencing library of each individual were marked up by MarkupDuplicates of Picard Tools\(^9\). Additional QC metrics for GC bias, insert size distribution, hybridization selection were also calculated from mapped reads by Picard Tools\(^9\). Mosdepth\(^10\) was used to calculate sequencing depth on exome targets (or 500 bp sliding windows for WGS) and determine callable regions at 10X or 15X coverage. Cross-sample contamination was tested by VerifyBamID\(^11\) using sequencing only mode. Samples were excluded if it has insufficient coverage (less than 80% targeted region with >=20X), shows evidence of cross-sample contamination (FREEMIX>5%), or discordant sex between normalized X and Y chromosome depth and self/parent reports that cannot be explained by aneuploidy.

Variants for each individual were discovered from mapped reads using GATK HaplotypeCaller\(^12\), weCall\(^13\), and DeepVariant\(^14\). Individual variant calls from GATK and weCall were stored in gVCF format and jointly genotyped across all samples in each sequencing batch using GLnexus\(^15\).

Variants were also jointly discovered and genotyped for individuals of the same family using GATK HaplotypeCaller\(^12\) and freebayes\(^16\), and then read-backed phased using WhatsHap\(^17\). To verify sample relatedness, identify overlapping samples with other cohorts, and verify sample identity with SNP genotyping data, genotypes of over 110,000 known biallelic SNPs from 1000 Genomes or HapMap projects that have call rate >98% and minor allele frequency (MAF) >1% in the cohort were extracted from joint genotyping VCFs. SNP array genotypes were called by Illumina GenomeStudio. We kept samples with >90% non-missing genotype calls and used genotypes of over 400,000 known SNPs that have call rate >98% and MAF>0.1 for relatedness check and ancestry inference.

De novo variants

We identified candidate de novo SNVs/indels from SPARK and SSC cohorts from per-family VCFs generated by GATK and freebayes and cohort-wide population VCF by weCall using a set of heuristic filters that aim to maximize the sensitivity while minimizing false negatives in parents\(^1\).

We then reevaluated the evidence of all de novo candidates from all input sources. Candidate was removed if there was contradictory evidence against from any input source (“contradiction filters”, see Supplementary Table S11). Further, we only kept candidates if they can be called by DeepVariant in offspring but have no evidence of variant in parents. For candidates that were identified in multiple offspring (recurrent), we only kept the ones that passed DeepVariant filter in all trios. For candidates that were shared by siblings in the same family, we only kept the ones with de novo quality estimated by triodenovo higher than 8 (or 7 for SNVs in CpG context). Before creating the final cleaned call set, we selected subsets of variants (see Supplementary Table S11) for manual evaluation by IGV to filter out candidates with failed review. Finally, we merged nearby clustered de novo coding variants (within 2bp for SNVs or 50bp for indels) on the same haplotype to form multi-nucleotide variants (MNVs) or complex
indels. We removed variants located in regions known to be difficult for variant calling (HLA, mucin, and olfactory receptors). DNVs in the final call set follow a Poisson distribution with an average 1.4 coding DNVs per affected and 1.3 per unaffected offspring (Supplementary Figure S25). The proportion of different types of DNVs, the mutation spectrum of SNVs, and indel length distributions were similar between SPARK and SSC (Supplementary Figure S2). A small fraction of variants in the final call set are likely post-zygotic mosaic mutations (Supplementary Figure S2).

Rare variants

Rare variant genotypes were filtered from cohort-wide population VCFs with QC metrics collected from individual and family VCFs (Supplementary Figure S27A). Briefly, we initially extracted high quality genotypes for each individual for variants that appear in less than 1% of families in the cohort. Evidence for the variant genotypes were re-evaluated by DeepVariant from aligned reads and collapsed over individuals to create site level summary statistics including fraction of individual genotypes that passed DeepVariant filter and mean genotype quality over all individuals. For variant genotypes extracted from GLnexus VCFs, we re-examined variant genotype from per-family VCFs by GATK to collect GATK site level metrics (including QD, MQ, SOR, etc.) then took read-depth weighted average over families to create cohort-wide site metrics. For variant genotypes extracted GATK joint genotyping VCFs, these site metrics were directly available directly from INFO fields.

Variant site level QC filters were calibrated using familial transmission information, assuming that false positive calls are more likely to show Mendelian inheritance error (Supplementary Figure S27B). Briefly, we first applied a baseline site level filter that favors high sensitivity, then optimized thresholds for filters with additional QC metrics. The selected QC metrics were reviewed first to determine a small number of optional thresholds. Then the final set of QC parameters were optimized from a grid search over the combinations of available thresholds such that: 1. presumed neutral variants identified from parents (silent variants or variants in non-constrained genes) shows equal transmission and non-transmission to offspring; 2. rates of neutral variants are similar in different sample groups from the same population ancestry; 3. vast majority variants identified in trio offspring are inherited from parents. In case when multiple sets of QC thresholds give similar results, priority will be given to the set that also recovers maximum number of DNV calls in trio offspring. The optimized filtering parameters were used in final QC filters to generate analysis-ready variants.

For a rare coding variant initially annotated as LoF (including stop gained, frameshift, or splice site), we searched for nearby variants on the same haplotype (within 2bp for SNVs or 50bp for indels). If nearby variants can be found, they were merged to form MNVs or complex indel and re-annotated to get the joint functional effect. If the joint effect was not LoF, then the original variant was removed from LoF analysis.

Variant annotations

The genomic coordinates of QC passed variants were lifted over to hg19 and normalized to the leftmost positions. Functional effects of coding variants were annotated to protein coding transcripts in GENCODE V19 Basic set using variant effect predictor. The gene level effect
was taken from the most severe consequences among all transcripts (based on the following
priority: LoF>missense>silent>intronic). pExt for each variant can be operationally defined as
the proportion of expression levels of transcripts whose variant effects are the same as gene
effect over all transcripts included in the annotation\textsuperscript{27}. We used transcript level expressions in
prenatal brain development from Human Developmental Biology Resource\textsuperscript{81} to calculate pExt.
Missense variants were annotated by pathogenicity scores of REVEL\textsuperscript{31}, CADD\textsuperscript{82}, MPC\textsuperscript{83} and
PrimateAI\textsuperscript{84}. Population allele frequencies were queried from gnomAD\textsuperscript{26} and ExAC\textsuperscript{18} using all
population samples. All rare variants were defined by cohort allele frequency <0.001 (or <0.005
for X chromosome variants). To filter for ultra-rare variants, we keep variants with cohort allele
frequency <1.5e-4 (or allele count=1) and population allele frequency <5e-5 in both gnomAD\textsuperscript{26}
and ExAC\textsuperscript{18}.

LoF variants on each coding transcript were further annotated by LOFTEE\textsuperscript{26} (v1.0, default
parameters). We also annotated splice site variants by SpliceAI\textsuperscript{85}, and removed low confidence
splice site variants with delta score <0.2 from LoF variants. pExt for LoF variants was calculated
by the proportion of expression level of transcripts that harbor HC LoFs evaluated by LOFTEE
over all transcripts included in the analysis. Thus, the pExt filter for LoFs already incorporated
LOFTEE annotations. The baseline filter to analyze rare, inherited LoFs and LoFs of unknown
inheritance is pExt\textgreater=0.1. To refine gene-specific pExt threshold in the second stage, we selected
95 known ASD/NDD genes plus a newly significant DNV enriched gene MARK2 which harbor at
least four de novo LoF variants in combined ASD and other NDD trios, and for each gene choose
the pExt threshold from \{0.1,0.5,0.9\} that can retain all de novo LoF variant with pExt\geq 0.1
(Supplementary Table S1).

Copy number variants
Copy number variants (CNVs) were called from exome read depth using CLAMMS\textsuperscript{86}. CNV calling
windows used by CLAMMS were created from exome targets after splitting large exons into
equally sized windows of roughly 500bp. Calling windows were annotated by average
mappability score\textsuperscript{87} (100mer) and GC content assuming average insert size of 200. Depths of
coverage for each individual on the windows were calculated using Mosdepth\textsuperscript{70} and then
normalized to control for GC-bias and sample’s overall average depth. Only windows with GC
content between 0.3 and 0.75 and mappability \textgeq=0.75 were included in further analyses. For
each given sample, we used two approaches to reduce the dimension of sample’s coverage
profile and automatically selected 100 nearest neighbors of the sample under analysis as
reference samples. The first approach used seven QC metrics calculated by Picard Tools from
aligned reads as recommended by the CLAMMS developer\textsuperscript{86}, we further normalized those
metrics in the cohort by its median absolute deviation in the cohort. The second approach used
singular value decomposition of the sample by read-depth matrix to compute the coordinates
of the first 10 principal components for each sample.

Model fitting and CNV calling for each individual using custom reference samples were
performed using default parameters. From raw CNV calls, neighboring over-segmented CNVs of
the same type were joined if joined CNVs include over 80% of the calling windows of original
calls. For each sample, we kept CNV calls made from one set of reference samples that have
smaller number of raw CNV calls. Outliers with excessive raw CNV calls (>400) were removed.
For each CNV, we counted the number of CNVs of the same type in parents that overlap >50%
of the calling windows. High-quality rare CNVs were defined as <1% carrier frequency among parents and have Phred-scaled quality of CNV in the interval >90. We queried high-quality rare copy number deletions to look for additional evidence to support new genes.

Genetic analysis

**De novo** variants analysis

In the discovery stage analysis, the DNV call sets of SPARK and SSC were merged with published DNVs from ASC\(^3^\)\(^8^\) and MSSNG\(^6^\) and additional SSC trios of which we did not have sequencing data. To infer likely samples overlaps with published trios of which we do not have individual level data, we tallied the proportion of shared DNVs between all pairs of trios. For a pair of trios, let \(N_1\) and \(N_2\) be the number of coding DNVs and \(O\) the number of shared DNVs between pair. To account for mutation hotspots, if a DNV is a SNV within CpG context or a known recurrent DNVs identified in SPARK and SSC, it contributes 0.5 to the count. Likely overlapping samples were identified if \(\frac{O}{N_1} \geq 0.5\) or \(\frac{O}{N_2} \geq 0.5\) and they have identical sex.

To determine the expected number of DNVs in the cohort, we used a 7-mer mutation rate model\(^52^\) in which the expected haploid mutation rate of each base pair (bp) depends on the 3bp sequence context on both sides. The per-base mutation rates were adjusted by the fraction of callable trios at each base pair which was the fraction of trios with \(\geq 10X\) coverage in parents and \(\geq 15X\) coverage in offspring. For published trios, we used an inhouse WGS data of 300 trios with average 36X coverage to approximate the callable regions. Gene level haploid mutation rates for different classes of DNVs were calculated by summing up the depth-adjusted per-base mutation rate of all possible SNVs of the same class. The rate for frameshift variants was presumed to be 1.3 times the rate of stop gained SNVs\(^53^\). Mutation rates in haploid X chromosome regions were adjusted for the observed male-female ratio (4.2) assuming mutation rates in spermatogenesis is 3.4 times higher than oogenesis\(^9^\). The exome-wide rate of synonymous DNVs closely matches the observed number of DNVs (Supplementary Figure S12). We also observed similar fold enrichment of damaging DNVs (vs. expected rate) in ASD cases across four cohorts after accounting for samples with family history (Supplementary Figure S12).

To perform gene-based test of DNVs, we applied DeNovoWEST\(^11^\) a simulation-based approach to test the enrichment of weighted sum of different classes of DNVs compared to the expected sum based on per-base mutation rates in each gene. We used empirical burden of DNVs to derive weights for different variant classes in constrained genes (ExAC pLI\(\geq 0.5\)) and non-constrained genes separately based on positive predictive values (PPV) (Supplementary Table S13). For ASD, we defined **de novo** D-mis variants by REVEL score \(\geq 0.5\), and the rest of **de novo** missense variants are taken as benign missense (B-mis). For other NDDs, we defined two classes of **de novo** D-mis variants by MPC score \(\geq 2\) or MPC\(\leq 2\) and CADD score \(\geq 25\), and the remaining **de novo** missense variants are B-mis. We first ran DeNovoWEST to test the enrichment of all nonsynonymous DNVs (pEnrichAll). To account for risk genes that harbor only missense variants, we ran DenovoWEST to test the enrichment of **de novo** missense variants only and applied a second test for spatial clustering of missense variants using DenovoNear\(^9^\), then combined evidence of missense enrichment and clustering (pCombMis). The minimal of pEnrichAll and pCombMis was used as the final p-value for DeNovoWEST. The exome-wide
significance threshold was set to 1.3e-6 (=0.05/(18,000 genes*2 tests)) to account for the two
tests. The analysis on replication cohort used the same weights as derived from discovery
cohort. Compared with the original publication\textsuperscript{11}, our implementation of DeNovoWEST used
different ways to stratify genes, determine variant weights, and calculate per-base mutation
rates. We applied our DeNovoWEST implementation on 31,058 NDD trios and compared with
published results on the same data set. The p-values from re-analysis show high overall
concordance with published results (Supplementary Figure S28). We used p-values from our re-
analysis on other NDD trios in comparative analysis with ASD.

Gene set enrichment analysis of DNVs was performed by DnEnrich framework\textsuperscript{32}. We included
all \textit{de novo} LoF and D-mis variants in 5,754 constrained genes from 16,877 ASD and 5,764
control trios. For each gene set, we calculated the fraction of weighted sums of damaging DNVs
in the set using PPV weights of constrained genes (Supplementary Table S13) for cases and
controls respectively. The test statistics for each gene set is the ratio of such fractions in cases
over controls. To determine the distribution of test statistic under the null hypothesis, we
randomly placed mutations onto the exome of all constrained genes, while held the number of
mutations, their tri-nucleotide context and functional impact to be the same as observed in
cases and controls separately. Note that by conditioning on the observed number of damaging
DNVs in cases and controls, we tested enriched gene sets in cases that are not due to an
increased overall burden. At each round of simulation, the permuted test statistic in each gene
set was calculated. Finally, the p-value was calculated as number of times the permuted
statistic is greater than or equal to observed statistic. Fold enrichment (FE) was calculated as
the ratio of between observed and average of test statistics over all permutations. We also
approximated 95% confidence interval for FE by assuming log(FE) follows normal distribution
with mean 0 and standard deviation determined by the p-value.

In all DNV analyses above, DNVs shared by full or twin siblings represent single mutational
events and were counted only once. When an individual carry multiple DNVs within 100bp in
the same gene, only one variant with most severe effects was included in the analysis.

Transmission disequilibrium analysis
The effect of inherited LoF variants was analyzed using TDT in each individual genes or in gene
sets. Rare LoF variants were first identified in parents without ASD diagnoses or intellectual
disability who have at least one offspring, then for each parent-offspring pair, the number of
times the LoF variant was transmitted from parents to offspring was tallied. For variants in
(non-PAR part of) X chromosome, we only used rare LoF variants carried by mothers without
ASD diagnoses or intellectual disability and analyzed transmission in different types of mother-
offspring pairs. For TDT analysis of rare, inherited missense variants in selected gene sets,
different D-mis definitions and allele frequency cutoffs were used (Supplementary Figure S3).

The over-transmission of LoFs to affected offspring was evaluated by a binomial test assuming
transmission equilibrium under the null hypothesis of 50% chance of transmission. In the
discovery stage, ultra-rare LoFs with pExt>0.1 were used in exome-wide transmission
disequilibrium and gene set enrichment analysis. For gene-based test, all rare LoFs with
pExt>0.1 were also used, and TDT statistic\textsuperscript{39} for each gene was calculated by \( z = \frac{T - NT'}{\sqrt{T + NT'}}, \) where
\( T'(NT') \) is the number of times LoF variants were transmitted (not transmitted) to affected
When offspring include monozygotic twin pairs, only one was kept in the transmission analysis. We prioritized 244 autosomal genes with z>1 in top 10% LOEUF or in top 20% LOEUF and A-risk>=0.4. In the second stage gene-based test, if a gene-specific pExt threshold is available, we used HC LoF variants passed the gene-specific pExt filter.

In gene set enrichment analysis of inherited LoFs, the rate of transmission to affected offspring in each gene set was compared with the transmission rate in rest of the genes in the background using chi-squared test.

Case control analysis

Pseudo-controls are constructed from parents without ASD diagnoses or intellectual disability in simplex families, using alleles that were not transmitted to affected offspring. Each parent without ASD diagnoses or intellectual disability contributes sample size of 0.5 to pseudo-controls. Rare LoFs in ASD cases whose parent data are not available and from other cases that were not utilized in DNV enrichment or TDT analysis were analyzed in this stage. Specifically, for each ASD case, we found out all his/her most recent unaffected ancestors without ASD diagnoses or intellectual disability in the pedigree and calculated the contributing sample size as 1 minus the summation of kinship coefficients with these ancestors. If the contributing sample size is greater than 0, then the sample was included in pseudo-cases after removing alleles that were observed in any unaffected ancestors without ASD diagnoses or intellectual disability used in TDT and alleles included in DNV analysis if any. Examples of such rare LoFs in cases and their contributing sample sizes are given in Supplementary Figure S29.

Rare LoFs in cases and controls for X chromosome were categorized separately for males and females. For male controls, because fathers do not transmit X chromosomes to sons, male controls include all fathers. In contrast, male cases only include those whose mothers do not have ASD diagnoses or intellectual disability (thus not included in TDT analysis). For females, because we only include mothers without ASD diagnoses or intellectual disability and affected sons in TDT, female pseudo-cases include all affected females. Female pseudo-controls were established from unaffected mothers in simplex families using alleles that do not transmit to affected sons. Each unaffected mother contributes a sample size of 0.5 to pseudo-controls. In both sexes, DNVs were removed from pseudo-cases.

For gene-based tests in Stage 2, case-control comparisons are not independent of TDT. So we used population references as controls, including gnomAD exomes26 (v2.1.1 non-neuro subset), gnomAD genomes26 (v3.1 non-neuro subset), and TopMed genomes88 (Freeze 8). Variants in the population references were filtered to keep those passed default QC filter in released data. For variants in gnomAD data set, we further removed variants located in low complexity region, because such regions are enriched with false positive calls89 but the default filter does not effectively remove variants in those regions. QC filters in the inhouse ASD cohort and in TopMed had already removed most of variants located in such regions. Variants from population references were re-annotated in the same way as rare variants identified in ASD cohort. In gene level case-control comparison of LoF burden, we used baseline pExt>=0.1 filter or gene-specific pExt threshold if available to define HC LoF variants. For LoF variants in selected genes, we also extracted curation results by gnomAD to remove curated non-LoF variants and manually reviewed IGV snapshots from gnomAD browser if available to remove
likely variant calling artifacts (Supplementary Data 1). Number of HC LoF variants were obtained from the summation of allele count in site level VCF files. Gene level burden of HC LoF variants between cases and population controls are tested by comparing the HC LoF variant rates between cases and controls using Poisson test. To account for different in depth of coverage, sample sizes are multiplied by the fraction of callable coding regions of each gene (>=15X for autosomes or female X chromosome, >=10X for male X chromosome) in ASD cases and in population controls respectively.

To account for sample relatedness in case-control analysis, we created a relationship graph in which each node represents an individual and each edge represents a known first or second-degree relationship between two individuals. We also add edges to pairs of individuals without known familial relationship but have estimated kinship coefficient >=0.1. From the graph, we select one individual from each connected component to create unrelated case-control samples. For chromosome X, father and sons were treated as unrelated. For population controls, only gnomAD data included sex specific allele counts and were used in the sex-specific analysis.

Meta-analysis was performed for prioritized autosomal genes among top 30% LOEUF. We integrated evidence from the enrichment of all DNVs, transmission disequilibrium, and increased burden in case compared with population controls by combining p-values using Fisher’s method. Experiment-wide error rate was set at 9e-6 (=0.05 divided by 5340 autosomal genes at LOEUF 30%). In mega-analysis, we combined all unrelated ASD cases together and compared CAFs of HC LoF variants with three population references.

Power calculation
To calculate statistical power of the current study and to estimate sample size for future gene discovery efforts, we adopted the statistical framework by Zuk et al. 2014 comparing CAF of LoF variants in \(N\) unrelated cases \(f_{\text{case}}\) with CAF \(f\) in natural population. The effect of LoFs in the same gene are assumed to be the same and increase ASD risk by \(\gamma\) fold. The population CAF \(f\) is assumed to be known with high precision from large cohorts. Since we only focus on LoF-intolerant genes in the population, \(f\) is assumed to be at selection-mutation equilibrium \(f = \frac{\mu_{\text{LoF}}}{s}\) where \(\mu_{\text{LoF}}\) is LoF mutation rate and \(s\) is selection coefficient. The test statistic asymptotically follows a non-central chi-squared distribution with 1-df and non-centrality parameter (NCP):

\[
\lambda = 4N \left[ \gamma f \ln \gamma + (1-\gamma) \ln \frac{1-\gamma f}{1-f} \right]
\]

Given the significance threshold \(\alpha\), power can be calculated analytically by

\[
1 - \beta = 1 - F(F^{-1}(1 - \alpha, 0), \lambda)
\]

where \(F(x, \lambda)\) is the cumulative distribution of \(\chi^2\) with NCP \(\lambda\).

To calculate sample size to achieve desired power \(1 - \beta\) at significance level \(\alpha\), we first solve NCP \(\lambda_{\alpha, \beta}\) from the above equation. Then sample size can be approximated by:

\[
n_{\alpha, \beta} \approx \frac{\lambda_{\alpha, \beta}}{4f[\gamma \ln \gamma - (\gamma - 1)]}
\]
For current study in ASD, sample size is $N = 31,976$ unrelated cases, experimental wide error rate is $\alpha = 9e-6$. Given continuing expansion of population reference, treating $f$ as known without error is a reasonable assumption for future studies. To calculate power for new genes identified in this study, we used point estimates of $\gamma$ and $f$ from mega-analysis using gnomAD exomes as population controls, and used $\mu_{\text{LoF}}$ computed from the 7mer context dependent mutation rate model$^{52}$ to convert $f$ to $s = \frac{\mu_{\text{LoF}}}{f}$. The required sample sizes were calculated to achieve 90% of power.

Power and sample size are both calculated as a function of relative risk for ASD ($\gamma$) and selection coefficient ($s$) across different haploid LoF mutation rates ($\mu_{\text{LoF}}$). We only considered $s$ between 0.01 and 0.5, because most prioritized genes have point estimates of $s>0.01$ (Error! Reference source not found.) and genes with $s>0.5$ are expected to harbor to de novo than inherited LoF variants and can to be identified from the enrichment of DNVs. Relative risk to ASD ($\gamma$) was constrained between 1 and 20 since we are mainly interested in discovering genes with moderate to small effects. The reduction in fitness $s$ is correlated with the increases in ASD risk $\gamma$ by $s = \gamma \pi s_D$ under the assumption of no pleiotropic effect, where $\pi$ is ASD prevalence and $s_D$ is decreased reproductive fitness of ASD cases. Based on epidemiological studies, current estimated prevalence of ASD is $\hat{\pi} = 1/54$,$^{30}$ estimated $s_D$ is for 0.75 male and for 0.52 female$^{91}$ so sex averaged $\hat{s}_D = 0.71$ (assuming male-to-female ratio of 4.2). In reality, most known ASD genes also show pleiotropic effects with other NDDs or associated with prenatal death and therefore $s \geq \gamma \pi s_D \approx \gamma \hat{\pi} \hat{s}_D = 0.013 \gamma$. So we only considered combinations of $(s, \gamma)$ that satisfy the condition: $s \geq 0.013 \gamma$.

Gene sets

To evaluate the contribution of known ASD risk genes to the burdens of DNVs and inherited LoF variants identified in this study, we collected 618 known dominant ASD/NDD genes from the following sources:

1. Known developmental disorder genes from DDG2P$^{52}$ (2020-02) that are dominant or X-linked and have organ specificity list includes brain or cause multi-system syndrome.
2. High confidence ASD genes collected by SFARI$^{93}$ (2019-08) with score of 1 or 2 excluding known recessive genes.
3. Newly emerging dominant ASD genes reported in recent literatures and included in SPARK genes list$^{94}$ (2020-07).

To evaluate the gene sets enriched by damaging DNVs or inherited HC LoFs, we used all constrained genes by ExAC pLI>0.5 or in top 20% of LOEUF as the background. Gene sets of the following five categories were collected for gene sets enrichment analysis.

Transcriptome and proteome

- For genes with brain-specific expression, we used processed RNA-seq data from Fagerberg et al. 2014$^{95}$ and selected genes with average reads per kilobase of transcript per million mapped reads (RPKM)>1 in brain and over four times of median RPKM of 27 tissues.
- Genes in co-expression modules M2 and M3 derived from weighted gene correlation network analysis (WGCNA) analysis of BrainSpan developmental RNAseq data were
previously reported to enrich for known ASD genes and collected from Table S1 from that reference.

• To find genes expressed in excitatory or inhibitory neurons, we selected genes from Mo et al. 2015 that have average transcripts per million (TPM) greater than 100 in excitatory and inhibitory neurons respectively.

• Synaptic genes including those encode presynaptic proteins, presynaptic active zone, synaptic vesicles, and postsynaptic density were collected from SynaptomeDB.

Neuronal regulome

• Putative CELF4 target genes are defined as genes whose iCLIP occupancy>0.2 in Wagnon et al. 2012.

• CHD8 target genes are defined as genes whose promoter or enhancer region overlap with CHD8 binding peaks in human neural stem cells or mid-fetal brain in Cotney et al. 2015.

• FMRP target genes in mouse were first collected from Table S2C of Darnell et al. 2011 with FDR<0.1. They were then mapped to orthologous human genes using homology mapping provided by MGI (2018-07).

• Genes targeted by RBFOX2 were selected from Weyn-Vanhentenryck et al. 2014 to have Rbfox2 tag counts greater 8. Due to high correlations between RBFOX1 and RBFOX3, targeted genes by the two RNA binding proteins were merged in one gene set and selected to have total tag counts of Rbfox1 and Rbfox3 greater than 24. Selected mouse genes symbols were then mapped to orthologous human genes using homology mapping provided by MGI.

Autism gene predictions

• ForecASD is an ensemble classifier that integrates brain gene expression, heterogeneous network data, and previous gene-level predictors of autism association to yield a single prediction score. We created two sets of genes with forecASD prediction score greater than 0.4 or 0.5.

• A-risk is a classifier that uses a used gradient boosting tree to predict autism candidate genes using cell-type specific expression signatures in fetal brain. We created three sets of genes with prediction score greater 0.4, 0.5 or 0.6.

Genetic evidence

• For genes enriched by DNVs in ASD, we selected genes showing nominal statistical evidence (P<0.01 or P<0.05 by DeNovoWEST) in discovery cohort of 16,877 trios.

• For genes implicated by in other NDD, we selected genes nominally enriched by DNVs in 31,058 NDDs (P<0.01 or P<0.01 by DeNovoWEST using our implementation).

• For genes in implicated in schizophrenia, we selected genes nominally significant (P<0.05) by gene-based test in latest schizophrenia case-control study of 24,248 cases and 97,322 controls.

Archetypal analysis: STRING v11 clusters and Human Phenotype Ontology (HPO) terms were formatted as gene-by-term binary matrices. The working gene list was taken as the union of forecASD top decile genes and the 62 autism-associated gene from this study (total 1,776 genes). A total of 583 genes from this set had annotations in both STRING and
HPO, and using these genes, a canonical correlation analysis (CCA) was carried out using the RGCCA package for R (https://cran.r-project.org/web/packages/RGCCA/index.html) using five components and sparsity parameter c1 set to 0.8 for both the HPO and STRING matrices. Component scores for all 1,776 genes were calculated using the STRING cluster annotations and the corresponding coefficients from the CCA. This 1,776 gene by 5 CC component matrix was used as input for archetypal analysis, and the optimal \( k \) (number of archetypes) was selected using the elbow plot heuristic, with the residual sums of squares (RSS) plotted as a function of \( k \). We displayed the archetypal embedding using the simplexplot() function of the archetypes R package. Genes were identified as “archetypal” if their top archetype coefficient was > 2x the next highest archetypal coefficient. Those genes that did not fulfill this criterion were classified as “mixed”, while those that did were assigned to their maximally-scoring archetype. Each of the six identified archetypes were given a human-readable summary description based on review of the top associated STRING clusters (Figure 7). Further cluster/term association results are available in Supplementary Table S10. Representative genes for each archetype were chosen from among the list of 62 risk genes identified in this study, using the top 6 genes for each archetype (note that these genes do not necessarily fulfill the “archetypal” criterion described above, but are simply the top six of the 62 for each archetype).
**Author Contributions**

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Competing interests

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References

1. Lord, C. *et al.* Autism spectrum disorder. *Nat Rev Dis Primers* **6**, 5 (2020).

2. Iossifov, I. *et al.* The contribution of de novo coding mutations to autism spectrum disorder. *Nature* **515**, 216-21 (2014).

3. De Rubeis, S. *et al.* Synaptic, transcriptional and chromatin genes disrupted in autism. *Nature* **515**, 209-15 (2014).

4. O’Roak, B.J. *et al.* Recurrent de novo mutations implicate novel genes underlying simplex autism risk. *Nat Commun* **5**, 5595 (2014).

5. Yuen, R.K. *et al.* Whole-genome sequencing of quartet families with autism spectrum disorder. *Nat Med* **21**, 185-91 (2015).

6. Yuen, R.K. *et al.* Whole-genome sequencing resource identifies 18 new candidate genes for autism spectrum disorder. *Nature* **515**, 209-15 (2014).

7. Feliciano, P. *et al.* Exome sequencing of 457 autism families recruited online provides evidence for autism risk genes. *NJP Genom Med* **4**, 19 (2019).

8. Satterstrom, F.K. *et al.* Large-Scale Exome Sequencing Study Implicates Both Developmental and Functional Changes in the Neurobiology of Autism. *Cell* **180**, 568-584 e23 (2020).

9. Study, D.D.D. Large-scale discovery of novel genetic causes of developmental disorders. *Nature* **519**, 223-8 (2015).

10. Study, D.D.D. Prevalence and architecture of de novo mutations in developmental disorders. *Nature* **542**, 433-438 (2017).

11. Kaplanis, J. *et al.* Evidence for 28 genetic disorders discovered by combining healthcare and research data. *Nature* **586**, 757-762 (2020).

12. He, X. *et al.* Integrated model of de novo and inherited genetic variants yields greater power to identify risk genes. *PLoS Genet* **9**, e1003671 (2013).

13. Nguyen, H.T. *et al.* Integrated Bayesian analysis of rare exonic variants to identify risk genes for schizophrenia and neurodevelopmental disorders. *Genome Med* **9**, 114 (2017).

14. Gaugler, T. *et al.* Most genetic risk for autism resides with common variation. *Nat Genet* **46**, 881-5 (2014).

15. Sandin, S. *et al.* The familial risk of autism. *JAMA* **311**, 1770-7 (2014).

16. Sandin, S. *et al.* The Heritability of Autism Spectrum Disorder. *JAMA* **318**, 1182-1184 (2017).

17. Grove, J. *et al.* Identification of common genetic risk variants for autism spectrum disorder. *Nat Genet* **51**, 431-444 (2019).

18. Lek, M. *et al.* Analysis of protein-coding genetic variation in 60,706 humans. *Nature* **536**, 285-91 (2016).

19. Krumm, N. *et al.* Excess of rare, inherited truncating mutations in autism. *Nat Genet* **47**, 582-8 (2015).

20. Kosmicki, J.A. *et al.* Refining the role of de novo protein-truncating variants in neurodevelopmental disorders by using population reference samples. *Nat Genet* **49**, 504-510 (2017).

21. Ruzzo, E.K. *et al.* Inherited and De Novo Genetic Risk for Autism Impacts Shared Networks. *Cell* **178**, 850-866 e26 (2019).
Wilfert, A.B. et al. Recent ultra-rare inherited variants implicate new autism candidate risk genes. *Nat Genet* **53**, 1125-1134 (2021).

pfelicano@simonsfoundation.org, S.C.E.a. & Consortium, S. SPARK: A US Cohort of 50,000 Families to Accelerate Autism Research. *Neuron* **97**, 488-493 (2018).

MacArthur, D.G. & Tyler-Smith, C. Loss-of-function variants in the genomes of healthy humans. *Hum Mol Genet* **19**, R125-30 (2010).

MacArthur, D.G. et al. A systematic survey of loss-of-function variants in human protein-coding genes. *Science* **335**, 823-828 (2012).

Karczewski, K.J. et al. The mutational constraint spectrum quantified from variation in 141,456 humans. *Nature* **581**, 434-443 (2020).

Cummings, B.B. et al. Transcript expression-aware annotation improves rare variant interpretation. *Nature* **581**, 452-458 (2020).

Cassa, C.A. et al. Estimating the selective effects of heterozygous protein-truncating variants from human exome data. *Nat Genet* **49**, 806-810 (2017).

Fuller, Z.L., Berg, J.J., Mostafavi, H., Sella, G. & Przeworski, M. Measuring intolerance to mutation in human genetics. *Nat Genet* **51**, 772-776 (2019).

An, J.Y. et al. Genome-wide de novo risk score implicates promoter variation in autism spectrum disorder. *Science* **362**(2018).

Ioannidis, N.M. et al. REVEL: An Ensemble Method for Predicting the Pathogenicity of Rare Missense Variants. *Am J Hum Genet* **99**, 877-885 (2016).

Fromer, M. et al. De novo mutations in schizophrenia implicate synaptic networks. *Nature* **506**, 179-84 (2014).

Parikshak, N.N. et al. Integrative functional genomic analyses implicate specific molecular pathways and circuits in autism. *Cell* **155**, 1008-21 (2013).

Weyn-Vanhentenryck, S.M. et al. HITS-CLIP and integrative modeling define the Rbfox splicing-regulatory network linked to brain development and autism. *Cell Rep* **6**, 1139-1152 (2014).

Darnell, J.C. et al. FMRP stalls ribosomal translocation on mRNAs linked to synaptic function and autism. *Cell* **146**, 247-61 (2011).

Cotney, J. et al. The autism-associated chromatin modifier CHD8 regulates other autism risk genes during human neurodevelopment. *Nat Commun* **6**, 6404 (2015).

Brueggeman, L., Koomar, T. & Michaelson, J.J. Forecasting risk gene discovery in autism with machine learning and genome-scale data. *Sci Rep* **10**, 4569 (2020).

Chen, S. et al. Dissecting Autism Genetic Risk Using Single-cell RNA-seq Data. *bioRxiv*, 2020.06.15.153031 (2020).

Ewens, W.J. & Spielman, R.S. The transmission/disequilibrium test: history, subdivision, and admixture. *Am J Hum Genet* **57**, 455-64 (1995).

Fisher, R.A. *Statistical methods for research workers, 11th ed. rev*, (Edinburgh, Oliver and Boyd, 1925).

Zuk, O. et al. Searching for missing heritability: designing rare variant association studies. *Proc Natl Acad Sci U S A* **111**, E455-64 (2014).

Wright, S. Evolution in Mendelian Populations. *Genetics* **16**, 97-159 (1931).
43. Skene, N.G. & Grant, S.G. Identification of Vulnerable Cell Types in Major Brain Disorders Using Single Cell Transcriptomes and Expression Weighted Cell Type Enrichment. *Front Neurosci* **10**, 16 (2016).
44. Willsey, A.J. *et al.* Coexpression networks implicate human midfetal deep cortical projection neurons in the pathogenesis of autism. *Cell* **155**, 997-1007 (2013).
45. Wang, S. *et al.* De Novo Sequence and Copy Number Variants Are Strongly Associated with Tourette Disorder and Implicate Cell Polarity in Pathogenesis. *Cell Rep* **24**, 3441-3454 e12 (2018).
46. Heyne, H.O. *et al.* De novo variants in neurodevelopmental disorders with epilepsy. *Nat Genet* **50**, 1048-1053 (2018).
47. Singh, T., Neale, B.M., Daly, M.J. & Consortium, o.b.o.t.S.E.M.-A. Exome sequencing identifies rare coding variants in 10 genes which confer substantial risk for schizophrenia. *medRxiv*, 2020.09.18.20192815 (2020).
48. Shu, C., Snyder, L.G., Shen, Y., Chung, W.K. & Consortium, o.b.o.t.S. Imputing cognitive impairment in SPARK, a large autism cohort. *medRxiv*, 2021.08.25.21262613 (2021).
49. Arnheim, N. & Calabrese, P. Understanding what determines the frequency and pattern of human germline mutations. *Nat Rev Genet* **10**, 478-88 (2009).
50. Rees, E., Moskvina, V., Owen, M.J., O'Donovan, M.C. & Kirov, G. De novo rates and selection of schizophrenia-associated copy number variants. *Biol Psychiatry* **70**, 1109-14 (2011).
51. Raychaudhuri, S. *et al.* Accurately assessing the risk of schizophrenia conferred by rare copy-number variation affecting genes with brain function. *PLoS Genet* **6**, e1001097 (2010).
52. Carlson, J. *et al.* Extremely rare variants reveal patterns of germline mutation rate heterogeneity in humans. *Nat Commun* **9**, 3753 (2018).
53. Samocha, K.E. *et al.* A framework for the interpretation of de novo mutation in human disease. *Nat Genet* **46**, 944-50 (2014).
54. Spielman, R.S. & EWens, W.J. The TDT and other family-based tests for linkage disequilibrium and association. *Am J Hum Genet* **59**, 983-9 (1996).
55. Chen, W.M., Manichaikul, A. & Rich, S.S. A generalized family-based association test for dichotomous traits. *Am J Hum Genet* **85**, 364-76 (2009).
56. Staples, J. *et al.* PRIMUS: rapid reconstruction of pedigrees from genome-wide estimates of identity by descent. *Am J Hum Genet* **95**, 553-64 (2014).
57. Genomes Project, C. *et al.* A global reference for human genetic variation. *Nature* **526**, 68-74 (2015).
58. Li, J.Z. *et al.* Worldwide human relationships inferred from genome-wide patterns of variation. *Science* **319**, 1100-4 (2008).
59. Bergstrom, A. *et al.* Insights into human genetic variation and population history from 929 diverse genomes. *Science* **367**(2020).
60. Patterson, N., Price, A.L. & Reich, D. Population structure and eigenanalysis. *PLoS Genet* **2**, e190 (2006).
61. Chen, C.Y. *et al.* Improved ancestry inference using weights from external reference panels. *Bioinformatics* **29**, 1399-406 (2013).
62. Bishop, S.L., Farmer, C. & Thurm, A. Measurement of nonverbal IQ in autism spectrum disorder: scores in young adulthood compared to early childhood. *Journal of autism and developmental disorders* **45**, 966-974 (2015).

63. Munson, J. *et al.* Evidence for latent classes of IQ in young children with autism spectrum disorder. *American journal of mental retardation : AJMR* **113**, 439-452 (2008).

64. Fischbach, G.D. & Lord, C. The Simons Simplex Collection: a resource for identification of autism genetic risk factors. *Neuron* **68**, 192-5 (2010).

65. Werling, D.M. *et al.* An analytical framework for whole-genome sequence association studies and its implications for autism spectrum disorder. *Nat Genet* **50**, 727-736 (2018).

66. Buxbaum, J.D. *et al.* The autism sequencing consortium: large-scale, high-throughput sequencing in autism spectrum disorders. *Neuron* **76**, 1052-6 (2012).

67. Li, H. Aligning sequence reads, clone sequences and assembly contigs with BWA-MEM. *arXiv* (2013).

68. Fritz, M.H.Y., Leinonen, R., Cochrane, G. & Birney, E. Efficient storage of high throughput DNA sequencing data using reference-based compression. * Genome Res* **21**, 734-740 (2011).

69. Institute, P.T.-B.B. https://broadinstitute.github.io/picard/.

70. Pedersen, B.S. & Quinlan, A.R. Mosdepth: quick coverage calculation for genomes and exomes. *Bioinformatics* **34**, 867-868 (2018).

71. Jun, G. *et al.* Detecting and estimating contamination of human DNA samples in sequencing and array-based genotype data. *Am J Hum Genet* **91**, 839-48 (2012).

72. Poplin, R. *et al.* Scaling accurate genetic variant discovery to tens of thousands of samples. *bioRxiv*, 201178 (2018).

73. GitHub - Genomicsplc/wecall: Fast, accurate and simple to use command line tool for variant detection in NGS data. https://github.com/Genomicsplc/wecall.

74. Poplin, R. *et al.* A universal SNP and small-indel variant caller using deep neural networks. *Nat Biotechnol* **36**, 983-987 (2018).

75. Lin, M.F. *et al.* GLnexus: joint variant calling for large cohort sequencing. *bioRxiv*, 343970 (2018).

76. Garrison, E. & Marth, G. Haplotype-based variant detection from short-read sequencing. *arXiv* (2012).

77. Martin, M. *et al.* WhatsHap: fast and accurate read-based phasing. *bioRxiv*, 085050 (2016).

78. Tan, A., Abecasis, G.R. & Kang, H.M. Unified representation of genetic variants. *Bioinformatics* **31**, 2202-4 (2015).

79. Frankish, A. *et al.* GENCODE reference annotation for the human and mouse genomes. *Nucleic Acids Res* **47**, D766-d773 (2019).

80. Autism Spectrum Disorders Working Group of The Psychiatric Genomics, C. Meta-analysis of GWAS of over 16,000 individuals with autism spectrum disorder highlights a novel locus at 10q24.32 and a significant overlap with schizophrenia. *Molecular autism* **8**, 21-21 (2017).

81. Lindsay, S.J. *et al.* HDBR Expression: A Unique Resource for Global and Individual Gene Expression Studies during Early Human Brain Development. *Front Neuroanat* **10**, 86 (2016).
Kircher, M. et al. A general framework for estimating the relative pathogenicity of human genetic variants. Nat Genet 46, 310-5 (2014).

Samocha, K.E. et al. Regional missense constraint improves variant deleteriousness prediction. bioRxiv, 148353 (2017).

Sundaram, L. et al. Predicting the clinical impact of human mutation with deep neural networks. Nat Genet 50, 1161-1170 (2018).

Jaganathan, K. et al. Predicting Splicing from Primary Sequence with Deep Learning. Cell 176, 535-548 e24 (2019).

Packer, J.S. et al. CLAMMS: a scalable algorithm for calling common and rare copy number variants from exome sequencing data. Bioinformatics 32, 133-5 (2016).

Koehler, R., Issac, H., Cloonan, N. & Grimmond, S.M. The uniqueome: a mappability resource for short-tag sequencing. Bioinformatics 27, 272-4 (2011).

Taliun, D. et al. Sequencing of 53,831 diverse genomes from the NHLBI TOPMed Program. Nature 590, 290-299 (2021).

Li, H. Toward better understanding of artifacts in variant calling from high-coverage samples. Bioinformatics 30, 2843-51 (2014).

Maenner, M.J. et al. Prevalence of Autism Spectrum Disorder Among Children Aged 8 Years — Autism and Developmental Disabilities Monitoring Network, 11 Sites, United States, 2016. MMWR Surveill Summ 69, 1-12 (2020).

Power, R.A. et al. Fecundity of patients with schizophrenia, autism, bipolar disorder, depression, anorexia nervosa, or substance abuse vs their unaffected siblings. JAMA Psychiatry 70, 22-30 (2013).

Wright, C.F. et al. Genetic diagnosis of developmental disorders in the DDD study: a scalable analysis of genome-wide research data. Lancet 385, 1305-14 (2015).

Abrahams, B.S. et al. SFARI Gene 2.0: a community-driven knowledgebase for the autism spectrum disorders (ASDs). Mol Autism 4, 36 (2013).

Singer, E. What Makes an Autism Gene? . (SPARK for Autism).

Fagerberg, L. et al. Analysis of the human tissue-specific expression by genome-wide integration of transcriptomics and antibody-based proteomics. Mol Cell Proteomics 13, 397-406 (2014).

Mo, A. et al. Epigenomic Signatures of Neuronal Diversity in the Mammalian Brain. Neuron 86, 1369-84 (2015).

Pirooznia, M. et al. SynaptomeDB: an ontology-based knowledgebase for synaptic genes. Bioinformatics 28, 897-9 (2012).

Wagnon, J.L. et al. CELF4 regulates translation and local abundance of a vast set of mRNAs, including genes associated with regulation of synaptic function. PLoS Genet 8, e1003067 (2012).

Eppig, J.T. Mouse Genome Informatics (MGI) Resource: Genetic, Genomic, and Biological Knowledgebase for the Laboratory Mouse. Ilar j 58, 17-41 (2017).

Szklarczyk, D. et al. STRING v11: protein-protein association networks with increased coverage, supporting functional discovery in genome-wide experimental datasets. Nucleic Acids Res 47, D607-d613 (2019).

Köhler, S. et al. The Human Phenotype Ontology in 2021. Nucleic Acids Res 49, D1207-d1217 (2021).
102. Cutler, A. & Breiman, L. Archetypal Analysis. Technometrics 36, 338-347 (1994).
103. Thorndike, R.L. Who belongs in the family? Psychometrika 18, 267-276 (1953).