Autism risk in offspring can be assessed through quantification of male sperm mosaicism

Martin W. Breuss1,2, Danny Antaki3,4,5, Renee D. George12, Morgan Kleiber3,4,5, Kiely N. James1,2, Laurel L. Ball1,2, Oanh Hong3,4,5, ileena Mitra7,8, Xiaoxu Yang1,2, Sara A. Wirth1,2, Jing Gu1,2, Camila A. B. Garcia1,2, Madhusudan Gujral3,4,5, William M. Brandler3,4,5,6, Damir Musaev1,2, An Nguyen1,2, Jennifer McEvoy-Venneri1,2, Renatta Knox1,2,9, Evan Sticca1,2, Martha Cristina Cancino Botello10, Javiera Uribe Fenner10, Maria CárceI PérezI, Maria Arranz11, Andrea B. Moffitt12, Zihua Wang12, Amaia Hervás13, Orrin Devinsky1,2, Jonathan Sebat3,4,5,* and Joseph G. Gleeson1,2*

De novo mutations arising on the paternal chromosome make the largest known contribution to autism risk, and correlate with paternal age at the time of conception. The recurrence risk for autism spectrum disorders is substantial, leading many families to decline future pregnancies, but the potential impact of assessing parental gonadal mosaicism has not been considered. We measured sperm mosaicism using deep-whole-genome sequencing, for variants both present in an offspring and evident only in father’s sperm, and identified single-nucleotide, structural and short tandem-repeat variants. We found that mosaicism quantification can stratify autism spectrum disorders recurrence risk due to de novo mutations into a vast majority with near 0% recurrence and a small fraction with a substantially higher and quantifiable risk, and we identify novel mosaic variants at risk for transmission to a future offspring. This suggests, therefore, that genetic counseling would benefit from the addition of sperm mosaicism assessment.

Clinicians are facing an ever-increasing incidence of autism spectrum disorders (ASD) in the population, without effective strategies available to prevent disease or counsel families. Recent studies have identified gene-damaging de novo mutations (DNMs) in at least 10–30% of simplex ASD cases1–4, along with the realization that the number of DNMs increases as a function of paternal age at the time of conception, doubling in DNM number in an offspring every 16.5 years of the father’s age at the time of conception5,6. A DNM, defined as a genetic variant present in an offspring but not detectable in either parent, can have any of several different origins7,8. While classically considered as occurring in the fertilized egg at the one-cell stage, most probably occur either postzygotically in the offspring or in a parent, either in the gonads or broadly in a mosaic pattern9. DNMs that occur during embryogenesis of a parent cause mosaicism in the soma, the gonads or both, and remain throughout life yet may be undetectable or barely detectable in blood10. However, the balance of gonadal-specific compared to broadly distributed DNMs in the father has not been carefully assessed, and thus the role of gonadal mosaicism in DNM recurrence risk remains uncertain.

Knowledge of the rates and mechanisms by which gonadal mutations arise has been advanced through assessment of multiple transmissions of DNMs within families, where approximately 1.3% of DNMs are shared by siblings11. Although only 3.8% of offspring DNMs are detectably mosaic in parental blood, this increases to 57.2% if shared by two or more offspring9,11. Countertuitively, DNM recurrence risk decreases by 1.8–2.3% per year of parent age, due to an increase in aging-associated DNMs9,11, thereby decreasing the relative contribution of parental mosaic variants to mutation burden.

Results
Sperm sequencing allows stratification of variants into low and high recurrence risk. We recruited eight families from our ASD cohort12,13, where each father agreed to submit a sperm sample for sequencing (Supplementary Dataset 1). Employing 30× whole-genome sequencing (WGS) from blood12,13, we defined 912 de novo single-nucleotide variants (dSNVs) in the 14 offspring (Fig. 1a and Methods). We then isolated sperm from the ejaculates and performed 200× WGS on paternal blood and sperm cells to determine which dSNVs were detectable in sperm based on three or more mutant reads (Extended Data Fig. 1 and Methods). We found 23 (2.5%) dSNVs that were also detected in paternal blood or sperm, leaving 889 (97.5%) dSNVs undetectable (Fig. 1b

1Department of Neurosciences, Howard Hughes Medical Institute, University of California, San Diego, La Jolla, CA, USA. 2Rady Children’s Institute for Genomic Medicine, San Diego, CA, USA. 3Beyster Center for Genomics of Psychiatric Diseases, University of California, San Diego, La Jolla, CA, USA. 4Department of Psychiatry, University of California, San Diego, La Jolla, CA, USA. 5Department of Cellular and Molecular Medicine, University of California, San Diego, La Jolla, CA, USA. 6Department of Pediatrics, University of California, San Diego, La Jolla, CA, USA. 7Department of Medicine, University of California, San Diego, La Jolla, CA, USA. 8Department of Computer Science and Engineering, University of California, San Diego, La Jolla, CA, USA. 9Department of Child Neurology, Weill Cornell Medical College, New York, NY, USA. 10Child and Adolescent Mental Health Unit, Hospital Universitari Mutua de Terrassa, Barcelona, Spain. 11Fundació Docència i Recerca Mutua Terrassa, Barcelona, Spain. 12Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, NY, USA. 13Research Laboratory Unit, Fundació Docència i Recerca Mutua Terrassa, Barcelona, Spain. 14Department of Neurology, Epilepsy Division, New York University School of Medicine, New York, NY, USA. *e-mail: jsebat@ucsd.edu; jogleeson@ucsd.edu
and Supplementary Dataset 2). Orthogonal validation of a subset with ultra-deep target amplicon sequencing (TAS) showed a validation rate of ~83% (15/18; Extended Data Fig. 2, Supplementary Dataset 3 and 4 and Methods). All three nonvalidated variants were at allelic fractions (AFs) <3% or located within repetitive elements (SINE or LINE).
Using the ratio of mutant to reference reads in blood and sperm, we defined four dSNV classes: sperm-detectable only (SDO); sperm-detectable enriched (SDE)—for which the AF was >3-fold higher in sperm than in blood (i.e., Δ > 3); sperm–blood equal (SBE, enrichment <3-fold); and blood-detectable only (BDO). Of the 23 variants, 34.8, 30.4, 26.1 and 8.7% were SDO, SDE, SBE and BDO, respectively. Nanopore long-read sequencing of the children allowed phasing of 501 of the 912 dSNVs to the paternal haplotype. Of the 23 mosaic variants, 20 resided on the paternal chromosome (40% SDO, 35.0% SDE, 25.0% SBE and 0% BDO; Fig. 1c). Thus, assessment of blood or using population risk underestimates paternal gonadal mosaicism (PGM) for most mosaic dSNVs (Fig. 1d). Furthermore, most dSNVs are not present in paternal sperm at this sensitivity level and thus have little measurable likelihood of recurrence.

The PGM burden was roughly equally distributed among the eight families (0–5 PGM variants/male), with AFs varying from 17% to the lower detection limit of 1.3% (Fig. 1e). Neither the number of mosaic variants nor their AF correlated with paternal age (Extended Data Fig. 3). We observed a mutational signature for PGM variants consistent with a developmental origin, not observed for nonPGM DNMs (for example, relative decrease in T>C variants14–16) (Extended Data Fig. 4). While PGM variants >7% AF were often also detectable in blood (10/11 SBE or BDO), variants below this level were typically restricted to sperm (7/12 SDO). Together, these data are consistent with an origin of PGM during embryonic development of the father, with those occurring earlier showing broader tissue distribution and higher AFs1. We next assessed the potential of sperm/blood sequencing to measure PGM for de novo structural variants (dSVs) and de novo short tandem-repeat variants (dSTRs; see Methods). Among the eight families, F01 had two de novo deletions (dDels) and F06 had one de novo duplication (dDup; Fig. 2a). One of these variants was detectably mosaic in paternal sperm, with an AF of 2–6% (Fig. 2b–d and Extended Data Fig. 5a–d). Among the eight families we identified 126 different dSTRΔs, of which 15 (11.9%) were mosaic (Fig. 2e–j, Extended Data Fig. 5e–h and Supplementary Dataset 5). Because 12 of 15 variants were SDO or SDE, recurrence risk assessment from blood alone would be erroneous for 80%.

**PGM extends to ASD pathogenic variants.** We next assessed whether clinically pathogenic DNMs could be detected in parental sperm, which could impact clinical decision making. We assessed a cohort of 14 families in which an offspring had ASD attributed to a dSNV or a 1–base pair (bp) dDel based on American College of Medical Genetics guidelines (Fig. 3a, Supplementary Text, Supplementary Table 1 and Supplementary Dataset 1). Using Droplet Digital PCR (ddPCR), three of 14 (21.4%) DNMs were detected as mosaic in sperm, with AFs of 14.47% (F09), 0.56% (F10) and 8.09% (F13) (Fig. 3b, Extended Data Fig. 6a–d and Supplementary Dataset 3). We were successful in phasing the 14.47% and 0.56% AF variants to the paternal haplotype (Supplementary Dataset 2 and 6). Three variants phased to the maternal haplotype posed no risk of PGM, but seven could not be phased, including the 8.09% AF variant. The F13 variant was absent in paternal blood (SDO), and the F09 variant was substantially reduced (SDE) (Extended Data Fig. 7a–c). These results, while representing a small number of DNMs, suggest that a substantial fraction of both paternally phased and unphased disease-related DNMs are detectable as PGM, and thus recurrence risk can be estimated directly.

Two variants showed sperm AF that predicted substantially elevated recurrence above the basal 1% recurrence risk in families (F09 at 14.47% AF and F13 at 8.09% AF). While F13 had a single child, F09, with a c.1007+1G>A known pathogenic variant in GRIN2A15,16 (Fig. 3c), had two older siblings lacking criteria for ASD, but deeper questioning revealed that both siblings showed neurodevelopmental abnormalities with no known cause (Fig. 3d and Supplementary Table 2). The middle child showed ADHD and speech impairment and the oldest child had ADHD and seizures, all consistent with GRIN2A haploinsufficiency. We collected DNA samples from the whole family and found that the GRIN2A c.1007+1G>A variant was heterozygous in all three children (Extended Data Fig. 6e). Thus, the mosaic variant in the father's sperm at 14.47% was transmitted to all three offspring, an unlikely but confirmed event, resulting in pleiotropic clinical features.

In our larger ASD cohort, five families had dSVs detected with standard WGS that were considered risk alleles (Supplementary Text). These included the de novo 22q12.3 dDel in F01 and the 1p36.32 dDup in F06 (Fig. 2a), as well as F18 with a 7q11.23 dDup and a 16p13.11 dDel, F19 with a 15q13.1-q13.3 dDel and F20 with a 1q21.3-q22.1 dDup (Fig. 3e). Several of the CNVs (for example, 15q13.1-q13.3) were flanked by directly oriented segmental duplications, suggesting that they may have arisen during meiosis through nonallelic homologous recombination17,18. A meiotic origin of these variants would preclude any possibility of PGM; however, as nonallelic homologous recombination may also occur during mitosis, these were still included in this analysis19.

We phased all of these variants and found that all bar one, the 7q11.23 dDup, phased to the paternal haplotype. Probe sets were designed to interrogate these variants from sperm using PCR and ddPCR copy number assessment (Fig. 3f and Supplementary Dataset 7). These assays confirmed the presence of the dSV in all tested probands, but did not reveal sperm mosaicism in any additional cases beyond F01 (Fig. 3g–i and Extended Data Fig. 7e–h). The 22q12.3 variant in F01 was mosaic in the father's sperm sample, based on the presence of a junction fragment matching the band in the proband and assessment of the deletion by nested PCR (Fig. 3g–i). ddPCR quantification showed 0.9382 mutant allele abundance in the proband (that is, heterozygous), whereas the father's sperm showed a 0.1358 abundance and his blood 0.0023 abundance (Fig. 3i), suggesting that ~7–8% of sperm carries the deletion. Thus, one of five dSVs was detectably mosaic in paternal sperm at an AF that could be considered clinically significant, since this would increase recurrence risk by ~7–8-fold (Extended Data Fig. 7e–h). The specificity of these assays precluded the confidencial exclusion of mosaicism in paternal sperm except for one additional variant (F20 with a 1q21.3-q22.1 dDup), and thus a negative predictive value is more difficult to calculate for most dSVs.

For three of the four pathogenic variants that were mosaic in sperm, a second semen sample, collected 1–4 months after the first, was subjected to mosaicism analysis by ddPCR (Extended Data Fig. 7d). While all three tested variants were detected at similar AF, the NR2F1 mutation exhibited a slight, but significant, difference between the two samples (P < 0.001). This suggests that mosaicism at these higher AFs is relatively stable over time.

**Unbiased analysis of sperm mosaicism detects 9–23 mosaic variants in sperm.** We next assessed the value in identifying PGM for variants not yet observed in children. Using the 200x sperm WGS on the eight fathers, we identified mosaic variants using both the intersection of variants of MutTect2 and Strelka2 (ref. 20,21), both optimized for mosaic variant detection in one tissue compared to another, as well as MosaicHunter22, optimized for mosaic variant detection shared between two tissues (Fig. 4a, Extended Data Fig. 8 and Supplementary Dataset 8). Combined, these methods identified 6/23 DNMs (from Fig. 1b) as PGM, since many of these occurred in repetitive sequences that were masked by these callers. This low recall rate was partially due to optimization of the pipeline for specificity (TAS, ~90% validation rate; Extended Data Fig. 9). To increase power for subsequent analyses on variants detected in blood and sperm, we defined three major groups of mosaic mutations—SDO, BSS (blood/sperm shared; includes SDE, SBE and blood detectable mosaic); and BDO, only detectable in blood. Combined, these methods identified 9–23 mosaic variants (Fig. 4b) in sperm.
enriched/BDE) and BDO (Fig. 4b). We identified 62 SDO, 61 BSS and 568 BDO, the last of these probably reflecting clonal hematopoiesis primarily arising from the father of F02 (Fig. 4c). There were 9–23 variants in the sperm of each father, all with the potential to transmit to an offspring.

The AF of PGM variants ranged from a maximum of ~35% to the lower limit of detection, ~1.5% (Fig. 4d). Compared with sperm AF, blood AF showed two trends: at higher sperm AFs, blood AFs were similar to sperm AF while at lower sperm AFs, blood AFs were very low or undetectable (Extended Data Fig. 9d–f).
This suggested two separate origins of PGM during paternal embryogenesis—the former occurring before and the latter after germ cell specification. The AF distribution of SDO, BSS and BDO was consistent with this model, where most SDO variants occurred at AFs <10%, whereas BSSs showed an AF range up to 35% (Fig. 4e,f). BDO AFs tended to mimic those of SDO, but there was a distribution tail with higher AFs probably reflecting clonal hematopoiesis. These BDO variants, while numerous, had little chance of being transmitted to an offspring because they were absent in sperm. Therefore, sequencing of blood only to identify potentially transmissible variants would not distinguish BDO from BSS and would miss SDO variants completely.

Mutational signatures suggest an embryonic origin of PGM. We then combined all mosaic SNVs detected in both approaches (Figs. 1 and 4) to observe common patterns for these variants. While there was no clustering along, or enrichment across, chromosomes (Extended Data Fig. 10a,b and Supplementary Dataset 9), we observed distinct mutational signatures differentiating variant classes. Assessing the relative contribution of each of the six possible...
base substitutions, mosaic variants differed from the background of Genome Aggregation Database (gnomAD) variants in several categories (Fig. 5a,b and Extended Data Fig. 10c). The early shared BSS mosaics differed from SDO and BDO variants, which were similar to each other. Supporting an embryonic origin of these variants, they were all depleted in T>C variants, a class that was correlated with environmental damage and aging gonads and depleted in variants that were shared among siblings. The differential signals for SSS variants enriched in C>A and T>G mutations relative to gnomAD and SDO and BDO mosaics are consistent with distinct mutational mechanisms in early embryonic development compared to those at later stages.

Discussion

Our results represent a significant improvement over previous strategies, where assessment of parental blood mosaicism only was used in combination with population statistics. The role of sperm mosaicism has been increasing recognized in single-gene disorders, and our work complements these efforts by providing a more general assessment of sperm mosaicism. Our data suggest a model of three major types of PGM (Fig. 4c): type I arises during the terminal differentiation of sperm and never recovers. Type II arises in proliferating spermatogonial stem cells (SSCs) and includes those that are extant clonally (IIa) or those under positive selection (IIb), akin to the ‘selfish sperm’ hypothesis. Type IIA probably represents mutations accumulating in individual SSCs and proposed to underlie the increased mutational load with age, although its importance in this process is controversial. Multiple inheritance is rare for IIA whereas IIB is similar to IIA because they have the same origin, but their selective advantage results in overproliferation of the SSC clone and the potential for population-wide recurrence.

Type III arises during paternal embryonic development, before primordial germ cell (PGC) specification or within the PGC population, and may be detectably mosaic in sperm, resulting in the potential for recurrence. The timing of a mutation probably determines its abundance and patterns of mosaicism between spermatogenic tissue, and our data suggest distinct mutational mechanisms between BSS and SDO variants.

Employing our methods, a distinction between the contribution of type I and type II mosaicism to the male-specific mutational burden is not possible, as both are below the detection limits; similarly, type III PGM occurring after PGC specification is probably not possible unless it is positively selected. In contrast, our work focuses on the detection of type III mosaicism, which can stratify the risk of recurrence. Considering the fraction of mosaic variants detected for each father, we estimate that on average 2.9% (95% CI: 1.6–7.1%) if a variant can be phased to the paternal haplotype. Of variants fall within this category, and this increases to 4.3% (95% CI: 1.6–7.1%) if a variant can be phased to the paternal haplotype. However, based on our data and those of previous, gene-centric studies on sperm mosaicism, even within this group, risk can vary by an order of magnitude.

Thus, the patterns of sperm mosaicism and the resulting framework for its detection that we present have the potential to impact clinical testing in two ways. First, direct assessment of previously
transmitted pathogenic variants in paternal sperm allows for the stratification of fathers with low and high recurrence risk through TAS or ddPCR analysis. Second, even without any previous risk or family history, prospective fathers who may want to know their risk of transmitting a high-impact variant to their child could undergo deep sequencing of their sperm, followed by mosaic analysis of these data. This potential is highlighted by our finding that one of the SDO variants (F06: chr9:131380333 G>A; NP_001123910.1:p.Arg1849Gln; 3.7% AF) was located in SPTAN1, a gene known to cause infantile epileptic encephalopathy (MIM: 613477). While this specific variant has not previously been reported, it was predicted to be ‘potentially disease-causing’ by MutationTaster, had a MutPred2 score of 0.687 (ref. 32) and a different nonsynonymous change in this same amino acid residue has been reported in...
affected children in ClinVar (SCV000243194.10, SCV000553140.2; p.Arg1849Trp). Based on our results, we would predict that this variant, which has the potential to be pathogenic, has a 3.7% inheritance risk for any subsequent child of the father in F06.

There are still several limitations and impediments regarding the application of sperm mosaicism testing. First, both approaches require the assessment of suspected high-penetrate variants and currently ignore modifiers and polygenic risk scores. This limitation is exemplified by the GRIN2A variant in family F09, where it is unclear whether the variability in expressivity is due to environment, genetic modifiers or stochasticity. Second, the absence of detectable mosaicism in paternal sperm can stratify the family into low risk only if the mutation of interest has been phased to the paternal haplotype. While phasing can be achieved through several experimental approaches, including the nanopore sequencing we present in this manuscript, its implementation in clinical practice is still uncommon. Third, while we show examples of resampling for three of the pathogenic variants and the relative stability of mosaicism between samples, it is unclear whether this is true across all mosaic variants and should be studied systematically in future. However, it is a problem that would be less relevant when testing sperm samples that are directly used for in vitro fertilization. Finally, our framework for the unbiased detection of mosaicism is tuned for specificity and may therefore miss clinically relevant variants. Similarly to mosaic analysis of cancer, implementation of sperm analysis for mosaic risk mutation has to be tuned for clinical application and may require large-scale secondary validation by methods such as TAS.

Online content
Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41591-019-0711-0.

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**Methods**

**Binomial modeling of detection threshold.** Depicted curves were based on a classic binomial model assuming that the AF of a mutation represents the probability of encountering a mutant read. The cumulative probability was calculated using the integrate-quad function of the scipy module from Python.

**Simulation and analysis.** To determine our sensitivity to detect mosaic variants, we created simulated datasets that contained known mosaic variants at low frequencies. We first randomly generated 10,000 variants from chromosome 22 as our set of mosaic variants. We then used Pysim to simulate Illumina paired-end sequencing reads from reference chromosome 22 and a version of chromosome 22 that contained the alternate alleles from our 10,000 mosaic variants. These two sets of reads were then combined to create a dataset of sequences with mosaic variants at 1, 2, 3, 4, 5, 10, 15, 20, 25 and 50% AF. The coverage of these datasets was 200×. We processed these reads through our standard mapping and somatic variant-calling pipeline before calculating sensitivity of detection of mosaic variants at each AF as the fraction of simulated variants called by our dsNV pipeline, or by both MuTect 2/Strelka 2 and MosaicHunter.

**Patient recruitment.** Patients were enrolled, according to approved human subjects protocols at the University of California, for blood, saliva and semen sampling. Semen was collected for all fathers of families F01–20. For F09–12, saliva from the fathers and their family members was obtained; for F01–08 and F13–20, DNA from blood was extracted. WES trio analysis for F09–12 was performed on DNA extracted from lymphocyte cell lines (generated by the NIMH Repository) and results of saliva samples. WES trio analysis for F01–08 and F13–20 was performed on DNA derived from blood. Each father provided a single sperm sample, with the exception of F01, F09 and F13, where a second sample was obtained, 3.5 and 4 months, respectively, after the first. Patients were part of two independent cohorts, assembled to identify dSNVs and dSVs through trio sequencing: ① the REACH cohort, consisting of 265 families with a proband with general features of ASD and recruited at Rady Children’s Hospital San Diego (J.S.) and at Mutua Terrassa Hospital Barcelona (M.A., A.H. and J.S.), and one focusing on 98 probands with ASD and an additional diagnosis of epilepsy, recruited at NYU Medical School (O.D. and J.G.G.; unpublished). The REACH cohort has been described previously. ② The cohort assembled by J.G.G. and O.D. represents a new recruitment effort that focused on patients with a diagnosis of ASD with associated epilepsy. Patients were evaluated by a child neurologist and a clinical geneticist for general and neurological assessment after referral from their primary care physician for concern about developmental delay and autism. Intellectual function was assessed by IQ score. Speech was assessed by a speech therapist fluent in the child’s native language. Brief videos of each affected member were created during the examination as part of the clinical assessment. Autism was assessed by a clinical psychologist using the Autism Diagnostic Interview-Revised, the Autism Diagnostic Observation Schedule and the Childhood Autism Rating Scale, and developmental milestones were assessed with the Vineland Scale and hyperactivity with the Conners Parent/Teacher Scale, all administered in the child’s native language by a trained psychologist. Epilepsy was assessed by a trained specialist and included history of daytime and night-time seizures, seizure types, length, onset and resolutions, and treatment history. Electroencephalography (EEG) was assessed awake and asleep using a minimum of 21 electrodes and ‘10 to ‘20 system placements as recommended by the International Federation of Clinical Neurophysiology. For all subjects, a 24-h EEG was recorded to assess the possibility of night-time seizures and to identify seizure foci. All subjects were recruited between the ages of 3–8 years. Patients were followed longitudinally to assess response to anticonvulsant therapy and behavioral therapy. All patients were seen at the NYU School of Medicine and were recruited through the ethical framework at the University of California, San Diego.

**Blood and saliva extraction.** DNA was extracted in an Autopure LS instrument (Qiagen).

**WES and WGS trio analysis.** WGS sequencing and analysis for F01–08 and F13–20 was performed as described previously. Exome capture and sequencing of F09–12 were performed at the New York Genome Center (Agilent Human All Exon 50 Mb kit, Illumina HiSeq 2000, paired-end, 2×100 bp) and the Broad Institute (Agilent Sure-Select Human All Exon v2.0, 44-Mb baited target, Illumina HiSeq 2000, paired-end, 2×76 bp). Sequencing reads were aligned to the hg19 reference genome using BWA (v.0.7.8). Duplicates were marked using Picard’s MarkDuplicates (v.1.83). Duplicate reads were realigned around indels (inDels) with GATK’s IndelRealigner. Variant calling for SNVs and InDels was performed according to GATK’s best practices by first calling variants in each sample with HaplotypeCaller and then jointly genotyping them across the entire cohort using CombinedGVCFs and GenotypeGVCFs. Variants were annotated with SnpEff (v.4.2) and SnpSift (v.4.2), and allele frequencies from the 1000 Genomes Project and the Exome Aggregation Consortium (ExAC) were also used. New variants were called for probands using Triodeno (v.0.06) with a minimum de novo quality score of 2.0 and subjected to manual inspection. Variants from F01–F08 were further interrogated for postzygotic mosaic variants (PMVs) that might be present in the children. Among all 912 variants, only four showed significant deviation from an expected 0.5 AF by using a binomial model, this effect was seen before multiple testing and disappeared following Bonferroni correction. This lack of PMVs in our data is most probably a reflection of limited sequencing depth (~40×) and cannot conclusively exclude the existence of PMVs in our data. Nevertheless, conservatively, we assumed that all 912 dSNVs were true DNsMs. We further interrogated F01–08 for possible paternally mosaic variants that might have been erroneously reported as inherited heterozygous variants; such artifacts might have resulted in an underestimation of mosaicism and overestimation of SDO and SDE variants. However, multiple filtering approaches did not result in the identification of any such variants. While we cannot exclude their existence, we believe that their contribution to mosaicism—if any—is minor in our dataset.

**Sperm extraction.** Extraction of sperm cell DNA from fresh ejaculates was performed as previously described. In short, sperm cells were isolated by centrifugation of the fresh (up to 2-d) ejaculate over an isotonic solution (90%) (Sage/Origio, no. ART-2100; Sage/Origio, no. ART-1006) using up to 2 ml of the sample. Following a washing step, quantity and quality were assessed using a cell-counting chamber (Sigma-Aldrich, no. BR717805-1EA). Cells were pelleted and lysis was performed by the addition of RLT lysis buffer (Qiagen, no. 79216). Bond-Breaker TCEP solution (Pierce, no. 77770) and 0.2-mm stainless steel beads (Next Advance, no. SS802) on a Disruptor Genie (Scientific Industries, no. SI-2381). The lysate was processed using reagents and columns from an AmpPrep DNA/RNA Mini Kit (Qiagen, no. 80204). Concentration of the final elution was assessed employing standard methods. Concentrations ranged from ~0.5–300 ng/µl.

**WGS of matched sperm and blood samples.** WGS was performed using an Illumina TrueSeq PCR-free kit (350-bp insert) or a TrueSeq Nano Kit (350-bp insert) on an Illumina HiSeq X Ten (paired-end FASTQ) for blood samples, and an Illumina HiSeq 3000 (paired-end FASTQ) sequenced blood and sperm samples from fathers were aligned to the hg19 reference genome (1000 Genomes v37) with BWA mem (v.0.7.15-r1410), specifying the −M option that tags chimeric reads as secondary and that are required for certain downstream applications that implement this legacy option. The resulting average mean coverage was 227× for blood samples and 222× for sperm samples, with an average read length of 150 bp for both sets. Duplicates were removed with the markdup command from sambamba (v.0.6.6), and base quality scores were recalibrated with the Genome Analysis Toolkit (GATK v.3.5-0-g36282e4). SNPs and InDels were called with Haplo tunerCaller jointly genotyping within pedigrees, consisting of the deep-coverage (~200×) genomes from the father’s blood and sperm and ~40×coverage genomes derived from the blood of both parents and children.

**Oxford Nanopore (ONP) sequencing and analysis.** Whole-genome sequencing libraries were generated with ONP 1D-long reads for all children (except for F03-II-2, due to lack of sufficient DNA) in deep-whole genome families (F01–F08) and deep-pedigree families with PMVs (F13–F20). reads were aligned to the GRCh38/hg19 reference genome with BW A mem with the -x ont2d option for ONP reads. Coverage of proband samples ranged from 3× to 15× (average, 8.6x), with an average read length of 5,349bp.

**Haplotype phasing.** To phase dSNVs, a set of phase-informative single-nucleotide polymorphisms (SNPs) from the WGS germline variant calls, or from an assembly of the local area using Nextera sequencing (see below) of a 20-kb region around the dSNV, was determined. Phase-informative SNPs were those where the child was heterozygous and either (1) one parent was heterozygous or homozygous for the alternate allele while the other was homozygous for the reference allele, or (2) one parent was heterozygous while the other was homozygous for the alternate allele. Second, where applicable, long reads (ONP reads, average length 5,349bp) were identified that contained both a dSNV and one or more phase-informative SNPs. The number of dSNV and phase-informative SNP combinations that were present in reads and consistent with the dSNV occurring on a maternal or paternal haplotype were counted. Reads containing an InDel flanking either the dSNV or the phase-informative SNP were excluded from the analysis. Finally, dSNVs were assigned to maternal and paternal haplotypes if there were: (1) a minimum of two counts and (2) the haplotype with the majority of counts had at least two-thirds of total counts. For F09–F12, F16 and F17, we attempted phasing using a Drop-Phase approach. In short, a complementary assay to the maternal allele at the dSNV position was designed for both the wild-type and the variant allele (see ddPCR design, validation, and setup of experiments for SNV analysis). Co-occurrence of the mutant dSNV was then assessed for both genotypes and quantified as described previously (Supplementary Dataset 8).

**Sanger sequencing of SNVs.** PCR and Sanger sequencing were performed according to standard methods. Primer sequences can be found in Supplementary Dataset 10. Validated mutations and surrounding SNPs were also used as basis for the design of ddPCR assays, where applicable.
ddPCR design, validation and setup of experiments for SNP analysis. Using the Primer3Plus web interface\textsuperscript{42–44}, the amplicon and probes for wild-type and mutant were designed to distinguish reference and alternate allele (settings are given in Supplementary Information under Additional information). Primers were required to be located within 15 bp up- and 15 bp downstream of the mutation and adjusted, so melting temperatures were matched between reference and alternate probe. In addition, if possible, amplicons were kept at 100 bp or shorter and probes at 20 bp or shorter. Specificity of the primers was assessed using Primer-BLAST. Custom primer and probe mixes (primer/probe ratio of 3:6) were ordered from IDT with FAM-labeled probes for the alternate, and HEX-labeled probes for the reference, allele (Supplementary Dataset 10). Optimal annealing temperature, specificity and efficiency were tested using custom gblocks (IDT) or patient DNA at a range of dilutions. ddPCR was performed on a Bio-Rad platform using a QX200 droplet generator, a C1000 touch cycler, a PX1 PCR Plate Sealer and a QX200 droplet reader. The following reagents were used: ddPCR Supermix (Bio-Rad, no. 1863024), droplet generation oil (Bio-Rad, no. 1863005), cartridge (Bio-Rad, no. 1864008) and PCR plates (Eppendorf, no. 95102036). Aiming for 30–60 ng per reaction, up to 8µl of DNA solution was used in a single reaction. Data analysis was performed using the software packages QuantaSoft and QuantaSoft Analysis Pro (Bio-Rad). Each run included technical duplicates or triplicates (as indicated in figure legends). For direct comparison of sperm samples we used seven technical replicates, except for F09 where the total amount of sperm DNA was limiting. Across all ddPCR reactions that were designed for SNP detection, we determined that the minimum AF that could reliably be detected was 0.1%. Therefore, we set this as the threshold of detection. Raw data for ddPCR experiments can be found in Supplementary Dataset 3.

TAS. PCR products for sequencing were designed with a target length of 160–190 bp, with primers being at least 60 bp distant from the base of interest. Primers were designed using the command-line tool of Primer3 with a Python wrapper script (Supplementary Dataset 10). PCR products were amplified according to the following procedures using GoTaq Colorless Master Mix (Promega, no. M7832) on sperm, blood and an unrelated control. Amplicons were either enzymatically cleaned with Exon (NEB, no. M0293S) and SAP (NEB, no. M0371S) treatment or gel extraction (Zymo Research, no. D4007) where necessary. Following normalization with the Qubit HS Kit (Thermo Fisher Scientific, no. Q33231), amplification products were processed according to the manufacturer’s protocol with SureSelect SPRI Beads (Beckman Coulter, no. A63881) at 1.2x. Library preparation was performed according to the manufacturer’s protocol using a Kapa Hyper Prep Kit (Kapa Biosystems, no. KK8501) and barcoded independently with unique dual indices (IDT for Illumina, no. 200223). After sequencing on an Illumina HiSeq 4000 with 100-bp paired-end reads, reads were mapped to the hg19 reference genome (IDT for Illumina, no. 200223). To assess the distribution of mosaic variants along the chromosomes, an equal number of variants (for mosaic dSNVs and unbiased calls that were sperm-specific, sperm mosaic, blood mosaic or blood-specific) was randomly generated with BEDTools from the called region from Strelka 2 with or without subtraction of the repeatMasker (rmsk.txt) file from UCSC as appropriate. This process was repeated 10,000 times to generate a distribution of the mean and s.d. of the distance between neighboring variants according to a broken-stick model\textsuperscript{18}.

Assessment of location of genome-wide distribution of mosaic variants. To assess the distribution of mosaic variants along the chromosomes, an equal number of variants (for mosaic dSNVs and unbiased calls that were sperm-specific, sperm mosaic, blood mosaic or blood-specific) was randomly generated with BEDTools from the called region from Strelka 2 with or without subtraction of the repeatMasker (rmsk.txt) file from UCSC as appropriate. This process was repeated 10,000 times to generate a distribution of the mean and s.d. of the distance between neighboring variants according to a broken-stick model\textsuperscript{18}.

Mosaicism analysis. Using the read depth information generated by HaplotypCaller, the AF for previously called dSNVs was determined. Additionally, dSNVs that fell within repetitive regions of the human genome were annotated using the repeatMasker (rmsk.txt) file from the University of California, Santa Cruz (UCSC). Variants that were homozygous alternate in the father and heterozygous in the proband, as well as those that were present in both blood and sperm, were processed according to the manufacturer’s protocol with SureSelect SPRI Beads (Beckman Coulter, no. A63881) at 1.2x. Library preparation was performed according to the manufacturer’s protocol using a Kapa Hyper Prep Kit (Kapa Biosystems, no. KK8501) and barcoded independently with unique dual indices (IDT for Illumina, no. 200223). After sequencing on an Illumina HiSeq 4000 with 100-bp paired-end reads, reads were mapped to the hg19 reference genome (IDT for Illumina, no. 200223). To assess the distribution of mosaic variants along the chromosomes, an equal number of variants (for mosaic dSNVs and unbiased calls that were sperm-specific, sperm mosaic, blood mosaic or blood-specific) was randomly generated with BEDTools from the called region from Strelka 2 with or without subtraction of the repeatMasker (rmsk.txt) file from UCSC as appropriate. This process was repeated 10,000 times to generate a distribution of the mean and s.d. of the distance between neighboring variants according to a broken-stick model\textsuperscript{18}.

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Normalized sequencing depth calculations generated by CNVView were derived from binned coverages in 45-kb nonoverlapping windows.

dSTrA calling and mosaicism detection. Analysis of STR expansions and contractions were performed using HipSTR \(^{50}\) (v.0.6) jointly on all BAM files (40x trios and >200x blood and sperm of fathers). The reference STR set provided by HipSTR for GRCh37 (GRCh37.hipstr_reference.bed) and default options were used, except for –del-stutter-model and –output-gfx. Furthermore, a modified version of HipSTR’s denofinder tool was run on each of the 40x trios.

The posterior probability of a de novo mutation was calculated using HipSTR' genotype likelihood and STR loci mutation rates as priors. Strict quality filters to detect de novo STRs were applied within trios. STR loci were excluded from analysis if they were in segmentally duplicated (UCSC hg19 genome SuperDups table)\(^{51}\) regions. Genotype STR Callings in all family members were required to have a minimum genotyping quality of 0.8, a maximum of 15% of reads with stutter or InDel, at least 10 spanning reads and at least 20% of reads to support each allele. STR loci were excluded if homozygous in the child or if they contained homopolymers and dinucleotide repeat motifs. de Novo STR mutations were further required to have a posterior probability of de novo mutation ≥0.8. Mutations were excluded if they were not a multiple of the repeat motif unit, or if the de novo allele was found in one of the parents at ≥0.1 allele frequency. STR mutations were further considered only if the repeat unit was ≥2, because homopolymers and dinucleotide repeats were enriched for false-positive calls. The remaining loci were annotated with their phase where possible, and de novo allele frequencies in the >200x sperm and blood samples. dSTrA were qualified as inconclusive if mosaicism was detected in both mother and father, as true de novo if no mosaicism was detected in the parents, as maternal if mosaicism was detected in the mother only, and as paternal if mosaicism was detected in blood, sperm or both. In regard to dSNVs, sperm-enriched variants were annotated as such if the AF was ≥3-fold higher in sperm than blood. Phase of STR was inferred from genotype: if a unique allele was inherited from one of the parents, the STRA was assumed to be derived from the other.

Nextera sequencing to identify informative SNPs. PCR products for sequencing were designed to encompass 1 kbp for the assembly of the local region around the mutation for phasing of F09–12 (Supplementary Dataset 10). Parallelized primer design was achieved using the web interface of PCRTiler\(^{53}\). PCR was performed consisting of 2 min initiation at 95 °C, 35 cycles of 95 °C for 30 s, 55 °C anneal for 30 s and 72 °C for 1 min, followed by a final extension at 72 °C for 2 min. Products were resolved on 2% agarose gels. For dPCR analysis, primer and probe sets for the SVs (copy number and break point analysis) were designed using Primer3Plus (Supplemental Text and Supplementary Dataset 10). Primers were designed to span the deletion breakpoints within the region or to lie within an intron of a coding gene. Preprint at bioRxiv https://doi.org/10.1101/049536 (2016).

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

Aligned BAM files generated for this study through deep WGS or TAS are available on SRA (accession no. PRINAS88332). WGS data used for de novo calling are available through the NIH Data Archive (NDA; collection ID: 2019).

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Author contributions
M.W.B., J.S. and J.G.G. conceived the project and designed the experiments. M.W.B., M.K., L.L.B., X.Y., S.A.W., C.A.B.G. and A.N. performed the experiments. D.A., R.D.G., I.M., X.Y., J.G., M.Gymrek, W.M.B., M.Gujral and M.W.B. performed the bioinformatic and data analyses. D.M., R.K. and E.S. performed de novo analysis of the cohort collected and provided by O.D. K.N.J., O.H., J.M.-V., M.C.C.B., J.U.F., M.C.P., M.A., A.H. and M.W.B. requested, organized and handled patient samples. A.B.M. and Z.W. performed the orthogonal sensitive detection of mosaic variants. M.W.B., J.G.G. and J.S. wrote the manuscript with input from R.D.G. and K.N.J. All authors saw and commented on the manuscript before submission.

Competing interests
M.W.B., D.A., M.K., K.N.J., W.M.B., J.S. and J.G.G. are inventors on a provisional patent (PCT ref. no. SD2017-181-2PCT) filed by UC, San Diego, titled ‘Assessing risk of de novo mutations in males’.

Additional information
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Correspondence and requests for materials should be addressed to J.S. or J.G.G.
Peer review information Kate Gao was the primary editor on this article and managed its editorial process and peer review in collaboration with the rest of the editorial team.
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Extended Data Fig. 1 | 200×WGS allows detection of mosaic variants down to 1% sensitivity. **a**, Plot showing the fraction of the genome that is covered at a given depth for blood and sperm following WGS with a target coverage of 200×. **b**, Plot showing the insert size of the reads for blood and sperm. **c**, Nanopore long-read technology (average read length 5,349 bp) was able to assign parental haplotype to 601/832 dSNVs in 13 children. Out of these, 501 were paternal, resulting in $\alpha \sim 4$ as reported previously. **d-e**, Binomial models for the detection limit of mosaic variants. Plots show the probability of detecting a given variant at a specific allelic fraction (AF) when requiring at least 3 alternate reads at different read-depths (**d**) or including a magnified inset for AF between 0.05 and 0 at 200× (**e**). **f**, Analysis of the power of detection assuming a minimum requirement of 3 reads at 200× sequencing. Plot shows the integrated probability of detection for the indicated tiers based on the curve seen in **e**. **g-h**, Plot of the fraction of detected variants (**g**) and the integrated detected fraction for the indicated AF ranges (**h**) of simulated data using Pysim. Results are from 10,000 variants simulated at 0.25, 0.20, 0.15, 0.10, 0.05, 0.02, and 0.01 AF. HaplotypeCaller was employed to detect variants as for data in Fig. 1.
Extended Data Fig. 2 | Orthogonal validation of a subset of mosaic dSNVs. a, 18 variants that could be assessed by ultra-deep target amplicon sequencing (TAS): shown are the reported 200× WGS results (square with horizontal line) and the results from TAS (closed circle) (shown are estimated fraction ± binomial 95% CI). Sperm (left, green) and blood (right, orange). Dashed line and grey box: upper 95% CI of an unrelated control and the area beneath to visualize likely false positive variants. y-axis: allelic fraction (%) for a log2 transformation of the data. Red text: variants that were considered to have failed orthogonal validation: 15/18 variants were successfully confirmed. Underlined variants were confirmed, but likely annotated as the wrong class (all 5 are probably SDO rather than SDE). For all data points, the estimated fraction and CI are based on the fraction of mutant reads, see Supplementary Data 2 and 4. b, Allelic fraction (determined by ddPCR or WGS read counts) of the mutant allele with the highest allelic fraction in sperm (F05: Chr22:23082101A>G). Sperm and Blood indicate samples from the father, other samples (Blood/ddPCR) were derived from the mother, the child harboring the dSNV (II-2), or control (Ctrl) blood. Graph shows individual data points (experimental triplicates) and mean ± SEM for the ddPCR data.
Extended Data Fig. 3 | Age correlation of all and mosaic dSNVs. **a**, Plot showing the increase in dSNV number with paternal age at birth, as described previously. Dashed line shows a regression curve demonstrating this dependence (n = 14 trios, adjusted R² = 0.526, P = 0.0020). **b**, Plot showing the increase in dSNV number with paternal age at birth for paternal variants only. As expected, this correlation was stronger than for non-phased variants (n = 13 trios, adjusted R² = 0.736, P = 0.000107). **c–d**, Plots showing correlation for paternal age and the number of mosaic variants or the mean AF in sperm. Paternal age/number of mosaic variants (c; n = 14 trios, adjusted R² = -0.048, P = 0.536) and paternal age/mean AF in sperm (d; n = 14 trios, adjusted R² = -0.047, P = 0.463) did not show any significant correlation. Adjusted R², coefficient of determination, and F-statistic nominal P-values are derived from a linear regression model through ordinary least squares. All graphs show individual data points, a regression line, and the 95% CI.
Extended Data Fig. 4 | Mutational signature for non-mosaic and mosaic dSNVs. 

**a.** Mutational signatures (6 categories) for non-mosaic and mosaic dSNVs, compared to the overall gnomAD signature and a permuted subset (n = 1,000 permutations for n = 889 (non-mosaic) and n = 23 (mosaic) dSNVs; shown is the 95% band). Asterisks indicate observed signatures that lie outside the 95% band of the permuted variants. Non-mosaic variants are largely reminiscent of the gnomAD signature (with the exception of a significant depletion of T > G). Mosaic variants exhibit some differences, but none reach significance due to the low number of available mutations. 

**b.** Mutational signatures (96 categories; trinucleotide environment for non-mosaic and mosaic dSNVs. 

**c.** Detailed view of the 96 mutational categories for non-mosaic and mosaic dSNVs, compared to the overall gnomAD signature and a permuted subset (n = 1,000 permutations for n = 889 (non-mosaic) and n = 23 (mosaic) dSNVs; shown is the 95% band). Dots indicate the observed mutational signature (black: within 95% band; red: outside the 95% band).
Extended Data Fig. 5 | See next page for caption.
Extended Data Fig. 5 Sperm mosaicism stratifies recurrence risk for dSV and dSTRΔ variants. a-c, Calculated copy number (a, c) and fraction of supporting reads (b) for the 6q16.1 deletion in F01 and the 1p36.32 duplication as indicated. Orange band in a and c ±1 SD of the CN using similarly sized regions across the genome (n = 1,000 random regions, see Methods). Plot in b shows the estimated fraction of supporting reads (estimated fraction ± binomial 95% CI; based on the fraction of mutant reads, see Supplementary Data 7). Together, these approaches suggest that these dSVs are not mosaic in paternal sperm. Note that the fraction of supporting reads could not be used for the duplication due to the repetitive elements flanking this SV. d, Copy number variant plot for the duplication in F06 for the Proband (40 × ), Father (200 × both), and the mother (40 × ). Visualization was performed with the CNView tool (see Methods). e, Correlation of the number of dSTRΔs with paternal age at birth. Dashed line shows a regression curve (n = 14 trios, adjusted R² = -0.058, P = 0.598). Adjusted R², coefficient of determination, and F-statistic nominal P-value are derived from a linear regression model through ordinary least squares. Graph shows individual data points, a regression line, and the 95% CI. f, Number of STR repeat units for non-mosaic dSTRΔs or those that are mosaic. No significant difference can be observed between the two groups (n = 111 non-mosaic variants and n = 15 mosaic variants; two-tailed Mann Whitney test; nominal P = 0.5490). Boxplots show median and quartiles with outliers as well as individual values. g, Detailed analysis of the TCTA repeat numbers in paternal, maternal, and child’s blood at low sequencing depth. Results show a de novo 13 × repeat in the child that is neither present in the father nor the mother. h, Sample reads showing the presence of a 10 × and 13 × allele in the child, a homozygous 10 × allele in the mother, a 10 × and a 12 × allele in the father, and the presence of a mosaic 13 × allele exclusively in paternal sperm.
Extended Data Fig. 6 | Sperm mosaicism stratifies risk for pathogenic ASD mutations. a–c AF (determined by ddPCR) of the mutant allele in paternal sperm (sperm) and maternal blood (mother) for the relevant dSNV in the 14 families. Part of this panel is also presented in Fig. 3. Ctrl—an unrelated sperm or blood sample, as indicated, acting as control. Graphs show individual data points (experimental triplicates) and mean ± SEM. d, Sanger sequencing results of paternal sperm for the locus harboring the dSNV for each family. Confirming the ddPCR results, F09, F10, and F13 showed mosaicism at their respective positions. e, Sanger sequencing results showing the C>T conversion locus in GRIN2A in F09 for all family members. The mutation was absent in the saliva of both parents, but present as a heterozygous allele in all 3 children.
Extended Data Fig. 7 | See next page for caption.
Extended Data Fig. 7 | ddPCR assessment of pathogenic structural variants and recurrent sampling of pathogenic DNMs in F01, F09, and F13. a–c, AF (determined by ddPCR) of the mutant alleles in F09 (a), F10 (b), and F13 (c). DNA tested was derived from paternal sperm and the saliva (a and b) or blood (c, bl.) of the father, mother, or affected child. In addition, controls for sperm (sp) and blood (bl) are provided. d, AF (determined by ddPCR) comparing two biological replicates of paternal sperm for F01, F09, and F13. The samples showed comparable levels of AF over time for all three samples, however, F13 exhibited a minor, but statistically significant difference. ***P < 0.001 (unpaired t-test, two-tailed, degrees of freedom = 12). e–g, Relative copy number (determined by ddPCR) for the three indicated dSVs for blood- and sperm-derived samples. Note that there is no detectable abnormality in the paternal sperm copy number above noise level, suggesting absence of sperm mosaicism in these samples. h, Direct copy number quantification of the duplication by ddPCR. All graphs show individual data points (experimental triplicates except for Affected in g [experimental duplicate], and F01 and F13 in d [7 experimental replicates]) and mean ± SEM.
Extended Data Fig. 8 | Limit of detection analysis for the unbiased analysis of gonadal mosaic SNVs. a-d, Plots of the fraction of detected variants (a, c) and the integrated detected fraction for the indicated AF ranges (b, d) of simulated data using Pysim for the intersection of MuTect 2/Strelka 2 (a, b) and MosaicHunter (c, d). Results were from 10,000 variants simulated at 0.25, 0.20, 0.15, 0.10, 0.05, 0.02, and 0.01 AF. This was the same data set as used in Extended Data Fig. 1. The MuTect 2/Strelka 2 and MosaicHunter pipelines were employed with the same filters as for the data in Fig. 4.
Extended Data Fig. 9 | Mosaic SNVs identified by unbiased analysis have a high validation rate and their AF differs depending on their origin. a-c, 74 variants that could be assessed by ultra-deep target amplicon sequencing (TAS); shown are the reported 200x WGS results (square with horizontal line) and the results from TAS (closed circle) (shown are estimated fraction ± binomial 95% CI). Sperm (left, green) and blood (right, orange). Dashed line and grey box: upper 95% CI of an unrelated control and the area beneath to visualize likely false positive variants. y-axis: allelic fraction (%) for a log2 transformation of the data. Plots are split by the three categories: SDO (a), BSS (b), and BDO (c). Red text denotes variants that were considered to have failed orthogonal validation: 13/19 (a), 21/21 (b), and 33/34 (c) were successfully confirmed. Underlined variants were confirmed, but likely annotated as the wrong class (that is, they are actually BSS for SDO and BDO variants in a and c, or are SDO (green text) or BDO (orange text) for BSS variants in c). For all data points, the estimated fraction and CI are based on the fraction of mutant reads, see Supplementary Data 2 and 8. d-f, Ranked plot of the estimated sperm and blood AF with 95% confidence intervals (estimated fraction ± binomial CI; based on the fraction of mutant reads, see Supplementary Data 8) for all variants detected in the three categories. SDO (d) and BDO (f) variants both show curves that are reminiscent of exponential decay, consistent with an increase of the number of mutations with expansion of the progenitor pool at a constant mutational rate. However, BSS (e) mosaicism for the first 40 variants appears to be more linear, suggesting that mutation rates for early division might be higher than those for later. This is consistent with previous models that estimated an elevated mutation rate in early embryonic development14.
Extended Data Fig. 10 | See next page for caption.
Extended Data Fig. 10 | Mosaic variants do not exhibit clustering but differ in their mutational signatures depending on their origin. a, Plot of the chromosomal location for each of the mosaic variants and their allelic fraction found in sperm from F01-08. Circles, triangles, and squares denote variants found to be mosaic by the dSNV approach, by the unbiased approach, or by both, respectively. b, Permutation simulations (n = 10,000 simulations of n = 23 mosaic dSNVs, n = 62 SDO mosaics, n = 123 SDO + BSS mosaics, n = 568 BDO mosaics, and n = 629 BDO + BSS mosaics) of variant locations to obtain mean and SD of broken stick fragment lengths. Vertical lines mark the observed value from mosaic dSNVs and mosaic variants from the indicated classes. These simulations illustrate that the observed distributions of variants along the chromosomes (as visualized in a for those that were mosaic in sperm) were within expectation. c, Detailed view of the 96 mutational categories for SDO, shared, and BDO mosaic variants, compared to the overall gnomAD signature and a permuted subset (n = 1,000 permutations for n = 68 (SDO), 72 (BSS), and 568 (BDO) gnomAD SNVs; shown is the 95% band). Dots indicate the observed mutational signature (black: within 95% band; red: outside the 95% band).
Reporting Summary

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Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

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- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

No software was used to collect the data in this study.

Data analysis

Commercial and available software used directly to analyze and visualize data: QuantaSoft (1.7.4.0917) and QuantaSoft Analysis Pro (1.0.596), GraphPad Prism 5, R (3.4.1 and ggbio package), Python (3.6.5 and matplotlib, seaborn, pysam, pandas, scipy modules), CNView, Strelka 2 (v2.9.2), MuTect 2 (v2.1), MosaicHunter (v1.0), HipSTR (v0.6), Pysam, Picard’s MarkDuplicates (v1.83), BWA (v0.7.8), Genome Analysis ToolKit (GATK version 3.5-0-g36282e4), Snpeff (v4.2), SnpSift (v4.2), Triodenovo (v0.06), BWA mem (version 0.7.15-r1140), sambamba (version 0.6.6), samtools (v1.9), and bedtools (v2.25.0). Variant calling was done using standard pipelines and programs as described in the methods section of this paper.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The datasets generated during the current study will be made available in public data bases and are available upon request from J.G.G. on reasonable request. Additionally, summary tables of all data are included as supplementary information files.
Field-specific reporting

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | Sample size for deep sequencing was based on previous mosaic studies on blood and the expected number of de novo mutations per trio. Based on this, we estimated that we require at least 10 trios to obtain a minimum of 500 DNMs for interrogation (current n=14 trios with n=912 dSNVs), to obtain >10 mosaic variants for analysis. Sequencing depth was determined by theoretical considerations (binomial model) and simulated data. For the analysis of pathogenic DNMs, based on previous sperm mosaicism studies on epileptic disorders, we estimated that around 10% of pathogenic variants may be mosaic. Thus, we expected that we need at least 15 DNM families to observe a mosaic variant with ~80% chance (n=20 pathogenic DNMs). |
| Data exclusions | No full data sets were excluded in this study. Individual variants were filtered based on previously established best practices for variant calling and mosaicism detection. |
| Replication | This is a descriptive study of fixed, unique cohort. Replication was not attempted. |
| Randomization | This is a descriptive study. No randomization was performed. |
| Blinding | No groups were allocated by the scientists. |

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

| Materials & experimental systems | Methods |
|----------------------------------|---------|
| n/a | n/a |
| [x] Antibodies | [x] ChIP-seq |
| [x] Eukaryotic cell lines | [x] Flow cytometry |
| [x] Palaeontology | [x] MRI-based neuroimaging |
| [x] Animals and other organisms | |
| [x] Human research participants | |
| [ ] Clinical data | |

Human research participants

Policy information about studies involving human research participants

Population characteristics

Human participants were derived from two autism cohorts (O.D. and J.S.). The study included parents, affected children and unaffected children of all sexes and ages.

Recruitment

The sole inclusion criterion was a medical diagnosis of autism spectrum disorder with or without epilepsy and a molecular diagnosis by next generation sequencing. The patient cohort was solely used for genetic analysis and the ascertainment of appropriate samples was performed as stated in our IRB protocol. In short, fathers of individuals with autism were previously enrolled through independent studies and subjected to whole genome or exome sequencing. All fathers with a pathogenic DNM who agreed to be recontacted for additional studies were informed of this study. As the existence of sperm mosaicism is independent of any possible selection bias (e.g. religious conflicts, history of vasectomy), willingness to participate and provide a semen sample is expected to be independent from measured outcomes.

Ethics oversight

IRB at UC, San Diego

Note that full information on the approval of the study protocol must also be provided in the manuscript.