Submegabase copy number variations arise during cerebral cortical neurogenesis as revealed by single-cell whole-genome sequencing

Suzanne Rohrback,a,b,1 Craig Apri,f, Fiona Kaper,f, Richard R. Rivera,f Christine S. Liu,a,b, Benjamin Siddoway,b and Jerold Chun a,2

aSanford Burnham Prebys Medical Discovery Institute, La Jolla, CA 92037; bBiomedical Sciences Graduate Program, University of California, San Diego, La Jolla, CA 92039; and Applied Genomics Tools, Research and Technology Development, Illumina Inc., San Diego, CA 92122

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Somatic copy number variations (CNVs) exist in the brain, but their generation, prevalence, forms, and biological impact remain unclear, even within experimentally tractable animal models. We combined a transposase-based amplification (TbA) methodology for single-cell whole-genome sequencing with a bioinformatic approach for filtering unreliable CNVs (FUnc), developed from machine learning trained on lymphocyte V(D)J recombination. TbA–FUnc offered superior genomic coverage and removed >90% of false-positive CNV calls, allowing extensive examination of submegabase CNVs from over 500 cells throughout the neurogenic period of cerebral cortical development in Mus musculus. Thousands of previously undocumented CNVs were identified. Half were less than 1 Mb in size, with deletions 4× more common than amplification events, and were randomly distributed throughout the genome. However, CNV prevalence during embryonic cortical development was nonrandom, peaking at mid-neurogenesis with levels triple those found at younger ages before falling to intermediate quantities. These data identify pervasive small and large CNVs as early contributors to neural genomic mosaicism, producing genomically diverse cellular building blocks that form the highly organized, mature brain.

Significance

Reports of copy number variations (CNVs) within single human brain cells have been limited to megabase-scale alterations in relatively few cells, leaving unclear when CNVs first arise and whether their generation is regulated. Answering these questions has been limited by an absence of experimentation with model organisms that allow developmental assessments infeasible with human samples. Here, we identify the existence and developmental dynamics of cerebral cortical CNVs in mouse, showing that their prevalence increases through midneurogenesis. Our improved sequencing approach also allowed characterization of previously undocumented neural CNVs below 1 Mb in size, comprising half of all alterations. These data demonstrate the existence of myriad CNVs, which genomically diversify neural cells before incorporation into the mature organization of the brain.

Conflict of interest statement: S.R., C.A., and F.K. declare competing financial interests in patents and patent applications filed by Illumina may encompass the methods, reagents, and data disclosed in this article. Some work in this study is related to technology described in patent applications 2010/0120098, 2011/0287435, 2013/0196860, and US2015/0376608.

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Data deposition: Sequencing data have been deposited in the National Center for Biotechnology Information Sequence Read Archive database, accession no. SRP121705.

1Present address: Applied Genomics Tools, Research and Technology Development, Illumina Inc., San Diego, CA 92122.
2To whom correspondence should be addressed. Email: jchun@sbpdiscovery.org.

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cerebral cortical neurogenesis, along with samples of cortical neurons and splenocytes from adults for control comparison (Fig. 1A). The DNA from a total of 688 single cells was amplified, from which 488 cells passed dataset quality controls (QC) for further interrogations (Table 1). To obtain sufficient DNA from a single cell for sequencing, we utilized TbA, which integrates whole-genome amplification and sequencing library preparation, analogous to optimizing the Nextera library preparation approach for scWGS applications (Fig. 1B). To assess the appropriateness of this method for accurate identification of CNVs, we compared samples collected from the same biological preparation but amplified by TbA vs. GenomePlex (Sigma), as the latter kit has been the most widely applied amplification methodology for scWGS applications (12–15). TbA generated datasets with consistently lower noise (Fig. 2A), which appears to be due to increased library complexity as indicated by higher genomic coverage per read (Fig. 2B). This allowed more and smaller CNVs to be identified when using TbA (Fig. 2C and D).

While developing our data analysis pipeline, we discovered that cells with highly altered genomes were erroneously subject to being discarded by several widely applied QC noise metrics (12–15). Specifically, male samples exhibited consistently higher values for two noise measurements, median absolute difference (MAD) and variability score (VS) (SI Appendix, Fig. S1A). This was caused by a systematically higher variability in normalized read depth across the monosomic X chromosome (SI Appendix, Fig. S1B). This increased noise appeared to be a technical artifact of hypoploidy: when data from two separately amplified, monosomic X chromosomes were combined, there was a reduction in read depth fluctuations to that of a disomic, female X chromosome (SI Appendix, Fig. S1C). This phenomenon was confirmed by analyzing independently generated aneuploid datasets (SRP041670; NCBI SRA) (15); cells containing at least one hypoploidy had significantly higher MAD and VS scores (SI Appendix, Fig. S1D). These results suggest that independent amplification of homologous chromosomes creates an averaging effect that produces a reduced error profile for disomic chromosomes. Since the median absolute pairwise difference (MAPD) noise statistic was the least impacted by hypoploidy across all assessments, it was selected for noise quantification in subsequent analyses (18).

Use of appropriate controls is critical in scWGS because the methodology destroys the original template, precluding direct replication of findings. Prior scWGS studies have used cells with constitutively present CNVs or aneuploidies to assess technical sensitivity (12–15, 19–21); however, constitutive CNVs are not somatically produced and may create analytical bias for the detection of a singular form of identical size and genomic location. We therefore established a positive control for stochastically generated CNVs by using splenic immune cells, which include B and T lymphocytes that have undergone somatic DNA recombination by means of V(DJ) recombination and (for B cells) heavy chain class switching. Blinded assessments of these cells identified deletions (≤2.5 Mb) mapping to the known V(DJ) recombination loci for B and T cells (Fig. 3), including the expected size range and chromosomal location. A total of 68 distinct recombination events were identified in 51 TbA-amplified splenocytes, forming the largest stochastically generated positive control dataset involving normal (not genetically diseased) cells to calibrate identification of somatic, neural CNVs. These controls were used both to determine appropriate

**Table 1.** Samples prepared and analyzed for somatic CNVs

| Group    | Animals*Method | Samples amplified/analyzed | No. of CNVs |
|----------|----------------|---------------------------|-------------|
| E11.5    | TbA            | 44/38                     | 232         |
| E12.5    | GP             | 39/28                     | 227         |
| E13.5    | TbA            | 38/28                     | 314         |
| E14.5    | GP             | 35/29                     | 120         |
|          | TbA            | 74/56                     | 1093        |
| E14.5    | GP             | 37/26                     | 134         |
| E16.5    | TbA            | 74/56                     | 1093        |
| E19.5    | TbA            | 46/39                     | 540         |
| Adult neurons | TbA        | 47/32                     | 302         |
| Splenic cells | GP        | 59/55                     | 524         |
|          | TbA            | 129/188                   | 837         |
| Totals   | GP             | 139/92                    | 506         |
|          | TbA            | 519/396                   | 4888        |
| All      | 658/488        | 5394                      |             |

*Multiple brains from one litter were pooled for embryonic samples.

Samples analyzed/amplified indicates both to the total number of datasets produced and the number passing QC requirements (Reads > 600,000; MAPD < 0.40; confidence score > 0.80). GP, GenomePlex (Sigma).

Fig. 1. Overview of study design and methods. (A) An extensive range of murine tissue samples was collected for analysis, including NPCs at six embryonic ages, adult cortical neurons, and adult splenocytes. (B) The TbA method to amplify genomic DNA was performed on single nuclei isolated by FANS, which involves tagmentation—enzymatic DNA fragmentation via insertion of universal sequencing adapters—followed by PCR with unique sample indexes. (C) Bioinformatic processing of data begins with calculating sequencing depth in ~0.1-Mb genomic regions, followed by CNV calling with the CBS algorithm, and finished with application of FUnC, which removes CNV calls that do not conform sufficiently to an integer copy number state.
TBa improves CNV detection capabilities. Comparisons were made and TbA (< 23) or TbA (n = 155). (A) Noise, measured by MAPD, is lower when using TbA. (B) This corresponds with increased genomic coverage. The red line shows the maximum coverage possible, to which TbA samples conform closely. The use of degenerate oligonucleotides for PCR in GenomePlex is less able to amplify unique genomic regions. (C and D) TbA detects more CNVs per cell (C) because many more small CNVs can be identified with the reduced noise (D). *p < 10^-4.

QC thresholds (SI Appendix, Fig. S2) and to characterize the appearances of genuine CNV events.

Another major challenge in scWGS is identifying and excluding false positives; therefore, FUNc was developed to filter unreliable CNV calls (Fig. 1C). Size has typically been the primary factor in removing uncertain CNV calls (12–14, 20) (Fig. 4A). However, V(D)J recombination events are comparatively small (Fig. 4B), and essentially all would be excluded by applying the previously published size cutoffs. To overcome this problem, we introduced the metric integer distance (intD), which reflects the difference between sequencing depth and integer copy number (Fig. 4A). Smaller values indicate higher quality. intD is typically close to 0 for positive controls of V(D)J recombination and euploid regions, but is higher for novel CNV calls, which are a mixture of genuine alterations and false positives (Fig. 4C).

We then employed a machine-learning approach to combine size and intD in an unbiased, nonlinear fashion. Separate models were built from euploid and V(D)J segments (from distinct chromosomal loci for Ig or T cell receptor recombination) with bagging (bootstrap aggregation) of one-class support vector machines (SVMs) to define the variable space within which reliable CNV calls appeared (Dataset S1). FUNc uses the 95% confidence interval around these models as cutoffs to assess the validity of CNV calls made by circular binary segmentation (CBS) (Fig. 4D and Dataset S2). Differences in intD were minimized after FUNc, but size distributions were nominally impacted (SI Appendix, Fig. S3A and B). Critically, this strategy enabled assessment of CNVs as small as 0.25 Mb, an order of magnitude smaller than many previous reports (average ~3.5 Mb) (12–14). TbA combined with FUNc identified myriad CNVs during cortical neurogenesis, only 12% of which were ≥3.5 Mb (Fig. 4E and Dataset S3).

We obtained multiple lines of evidence to validate use of FUNc. First, FUNc had little impact on our ability to classify splenocytes (SI Appendix, Fig. S3C). Second, splenocytes prepared by a different researcher after FUNc development showed a high rate of positive control CNV inclusion (SI Appendix, Fig. S3D). Third, when analyzing an independent dataset (SRP041670; NCBI SRA) (13), FUNc retained nearly all constitutive, validated CNVs from characterized cell lines (SI Appendix, Fig. S3E) and excluded a majority of likely false-positive CNVs (SI Appendix, Fig. S5F).

Fourth, simulated datasets were created to generate large numbers of false- and true-positive CNVs (Datasets S4 and S5). FUNc eliminated 90.8% of false positives, reducing the false discovery rate from 56.2 to 4.9% (Fig. 4F). In contrast, only 16.9% of true positives were excluded by FUNc, causing the false-negative rate to increase from 16.0 to 30.2% (Fig. 4G).

Our large sample size and moderate sequencing depth provided an opportunity to assess a topic of controversy, aneuploidy in nonmitotic NPCs. The gender of nonmitotic (interphase and postmitotic) cells and nuclei—which contain uncondensed DNA and an intact nuclear envelope sufficient to allow fluorescence-activated nuclear sorting (FANS)—could be clearly identified by sex chromosome aneuploidies (male) (Fig. 5A). These samples also produced identifiable but rare cells that were fully aneuploid (two cells) (Fig. 5B), consistent with the pattern predicted by metaphase spread imaging. However, cells much more often displayed one or more of what could be called “fragmented aneuploidies”: numerous CNVs along large, distinct chromosome regions, reminiscent of chromothryptic chromosomes in RPE-1 cell lines (5) (Fig. 5 C and D). If aneuploidy arbitrarily included cells with ≥50% of a chromosome altered (14), aneuploidy rates increased threefold, to ~2% of all cells (Fig. 5 E and F), with higher frequencies at younger ages consistent with prior studies (2, 6, 7).

Cortical CNVs followed a genome-wide, apparently random distribution, in clear contrast with V(D)J recombination (Fig. 6A). The absence of obvious genomic hot spots affected by CNVs was confirmed by hierarchical clustering of genome-wide copy number profiles: only B and T cell loci formed compelling clusters (SI Appendix, Fig. S4). Likewise, dimensionality reduction of the genomic element composition of CNV loci could not distinguish between sample types any better when using true data than randomly selected genomic regions (SI Appendix, SI A and B). Lymphocyte deletions showed elevated vertebrate evolutionary conservation because of the highly conserved recombination signal sequences in V(D)J loci (SI Appendix, Fig. S6).
Aneuploidy follows a fragmented pattern.

Previous Studies

Appendix, Fig. S5D) (22, 23). At younger ages, amplification events were weakly enriched for S-phase genomic regions associated with early DNA replication, consistent with the increased proportion of proliferating NPCs at such ages (SI Appendix, Fig. S5C) (24). Otherwise, only centromeres and telomeres were consistently more affected by CNVs than expected by chance.

The most remarkable aspect of CNV prevalence was variation during cerebral cortical neurogenesis. The number of CNVs per cell increased until midneurogenesis (E14.5), peaking at double

Fig. 4. Machine learning on true positive copy number assignments allows reliable analysis of sub-Mb somatic CNVs. (A) Illustration of metrics used to describe CNV calls. Size is measured in “bins” – 0.1-Mb intervals of equal mappability (blue diamonds). The pale blue lines show the average read-depth in segments (contiguous regions predicted to have the same copy number state by the circular binary segmentation algorithm), and the red line shows the predicted copy number states. intD is the difference between these lines. (B and C) Segment size (B) and intD (C) follow distinct distributions for euploid genomic regions (n = 3,174), immune recombination events (n = 68), and putative CNVs (n = 1,711). (D) Results after 10,000 iterations of building one-class support vector machines by bootstrapping the immune recombination events or euploid regions. Darker background colors indicate more frequent model inclusion; the creation of separate models accounts for the artificial gap between small and large sizes. The yellow line is a combined 95% confidence interval, used by FUnC as the maximum allowable intD for a given CNV size. (E) CNV size distributions for all biological groups assessed. The red line indicates the interpreted average minimum size of previously reported neural CNVs (3.5 Mb), which excludes >85% of CNVs identified here. Sample sizes are listed in Table 1. (F and G) Computationally simulated false positives (F; n = 87) rarely pass FUnC while simulated true positive CNVs (G; n = 2,910) fit well within the reliable-CN bound. *P < 0.0001 vs. euploid; †P < 10^−10 vs. immune.

Fig. 5. Aneuploidy follows a fragmented pattern. (A) A euploid male cell, where monosomy X is identified as a single copy number state. (B) An E16.5 NPC with classical, whole-chromosome aneuploidy. (C and D) Numerous CNVs along distinct chromosomes—fragmented aneuploidies—creates either apparent hyperploidies (C) or hypoploidies (D). (E) The amount of a chromosome altered by CNVs forms a continuous distribution (n = 7,920; chromosomes from all high-quality TbA-amplified samples). (F) Aneuploidy rates when considering chromosomes with ≥50% amplified or deleted to be aneuploid. Error bars show SEM; sample sizes are listed in Table 1.
the average adult neuron rate, and triple that observed at E11.5 (Fig. 6B). Over 93% of samples contained one or more CNVs, and elevated rates corresponded with an increase in the proportion of cells containing high, rather than moderate, numbers of CNVs (Fig. 6C). While both amplifications and deletions increased with CNV frequency, deletions predominated (Fig. 6D) and produced a cumulative DNA loss from E12.5 to E14.5 (Fig. 6E). Multiple biological preparations of E13.5, E14.5, and lymphocyte samples confirmed the differences among age groups (SI Appendix, Fig. S6).

**Discussion**

The use of both the TbA whole-genome amplification technique and unbiased modeling of true positive CNVs (to create FUnC) expanded the range of detectable neural GM, particularly allowing the identification of submegabase (sub-Mb) CNVs that were invisible to previous approaches and revealing many more CNV-positive neural cells than previously recognized. It is virtually certain that even more CNVs exist below 0.25 Mb, especially considering the nonidentical alterations (e.g., SNVs) observed by high depth sequencing after massive amplification or clonal expansion (4, 25).

Within the field of single-cell genomics, there has been a trend toward collecting ultra-low-coverage data to screen very large numbers of cells for CNVs (20, 26). This approach is advantageous when CNVs are prevalent, large, and clonal. That does not describe the characteristics that we have observed in cortical samples, where nearly half of all CNVs were small; medium-resolution, low-noise data were essential to identify such alterations. CNVs have been considered by some as unimportant for neuroscience (13, 15, 20) based upon studies limited to multimegabase-sized CNVs. However, our findings support a marked prevalence of sub-Mb alterations, suggesting their potential to alter neural phenotypes. Indeed, proof-of-concept for physiological roles of CNVs. However, our findings support a marked prevalence of sub-Mb alterations, suggesting their potential to alter neural phenotypes. Indeed, proof-of-concept for physiological roles of CNVs.

The observed developmental differences in CNV prevalence have parallels to DNA double-strand break generation, programmed cell death, and aneuploidy. Nonhomologous end joining (NHEJ) proteins are essential for viable cortical neurogenesis at similar developmental stages as the peak in CNV prevalence and can play a role in CNV generation (5, 8, 27, 28). Further research to elucidate the potential involvement of NHEJ pathways in cortical CNV presence could provide insights into the importance of these somatic alterations.

Programmed cell death is elevated around E14 in the cortex (17, 29), and while the CNVs reported here are much larger than DNA fragments associated with apoptosis (29), some cells with a high CNV burden probably die (17, 30). Cell death might in part explain the developmental reduction in CNVs reported here. In further support of this relationship, very large CNVs that are aneuploidies (2) are altered in form and number by inhibiting apoptosis via caspase genetic deletion or pharmacological inhibition (30). Notably, the peaks of programmed cell death (17, 29) and CNV prevalence coincide, implicating a relationship between these two phenomena in the embryonic cerebral cortex. The genome-wide location of CNVs throughout neurogenesis does not point to a specific locus promoting cell death, being more consistent with the concept of a quantitative threshold of CNV production beyond which cell death occurs.

Aneuploidy, as detected in metaphase spreads, is not only altered by cell death, but also prevalent in the embryonic cortex (~30% of NPC metaphase spreads) and preferentially involves DNA loss (1, 6, 7). The relationship between aneuploidy in metaphase spreads versus interphase scWGS is unknown; however, similarities between rates of metaphase spread aneuploidies and CNVs reported here along with the mutual preference for DNA loss, support an association between previously reported aneuploidies and the CNVs identified in this study. To match the reported metaphase aneuploidy rates (7, 30), nonmitotic chromosomes with ≥2% of their length affected by CNVs would need to be considered aneuploid. The relationship between the historical gold standard of metaphase spread aneuploidy and nonmitotic, TbA-identified fragmented aneuploidy is not known. However, it is conceivable that fragmented aneuploidies formed by many small CNVs as noted here could manifest as metaphase aneuploidy upon chromosome condensation. Indeed, this possibility is supported by reported chromosomal aberrations affecting ~65% of neuronal nuclei following somatic cell nuclear transfer to allow chromosomal condensation (31), implicating a linkage between fragmented aneuploidies in nonmitotic neuronal cells, as this approach provides a snapshot of how an interphase NPC could appear if instantaneously condensed to a metaphase spread.

**Fig. 6.** CNVs are generated and developmentally varied during cortical neurogenesis. (A) The frequency with which each genomic locus is affected by an amplification (above the gray line) or deletion (below the gray line) event. The Ig heavy chain and TCR α chain are indicated by open and closed arrowheads, respectively. (B) The number of CNVs per cell increases through E14.5. (C) The proportion of cells containing 0, moderate (≤5), or extreme CNV numbers. (D) Amplification and deletion events per cell showing a preference for DNA loss. (E) Heatmap of net DNA change per cell. The dashed line indicates 0, an equal quantity of DNA amplified and deleted. All error bars show SEM; sample sizes are listed in Table 1; **P < 0.05 vs. E13.5; †P < 0.01 vs. E14.5; ‡P < 0.001 vs. E11.5; §P < 0.001 vs. Lym; ¶P < 0.001 vs. deletions.
CNV prevalence peaks during midcortical neurogenesis, strongly supporting their initial, but not exclusive, generation during prenatal life. Additionally, both the deletion predominance and the apparently random genomic localization that we have observed in mouse are consistent with scWGS studies in the human cerebral cortex (12–15). Use of array-comparative genomic hybridization revealed more prevalent clonal CNVs in the elderly human brain (32), which might be explained by technical and/or biological differences. Our use of fetal mouse brain cells limited to a mean of ~20 cells per brain likely missed low-frequency clones that were initiating expansion and/or that were not sampled.

Our data identify CNVs as a universal feature of neural cells in the mammalian cerebral cortex that varies developmentally and, with aneuploidies and likely other sequence forms (2), contributes to neural GM. Although mechanisms regulating CNV generation, cell-type specificity, and their functional significance remain to be determined, our data indicate the existence of a vast range of previously unrecognized, smaller CNVs that can occur throughout the genome, a number that will surely increase as the resolution of scWGS improves. The alteration of CNVs during development supports involvement of regulatory mechanisms within the normally developing brain that could also be disrupted by environmental perturbations and disease, which likely affect specific genes.

Materials and Methods

Detailed descriptions of methods are presented in SI Appendix.

Sample Isolation. All animal protocols were approved by the Institutional Animal Care and Use Committee at The Scripps Research Institute and conform to National Institutes of Health guidelines. Embryonic cortices were collected from timed-pregnant dams, cortical neurons, and splenocytes from adult C57BL/6J mice. Nuclei isolation was performed as described previously (3, 9). Samples were sorted into sterile, low-bind, DNase/RNase-free strip tubes containing 2.5 μL sterile PBS and stored at −80 °C.

Tbang Library Preparation. All pre-PCR reagents were manufactured in a clean room using aseptic techniques. Genomic DNA released from cells was directly tagmented in the context of the cell lysate by adding Tagment DNA Buffer (5×; Illumina, Inc.) and Nextera transposomes (2 nM; Illumina, Inc.) for a total volume of 20 μL, followed by a 5-min incubation at 55 °C. After tagmentation, the reaction was stopped by adding 5 μL of 0.1% SDS and incubated for 5 min at room temperature. The library fragments were subsequently amplified by adding 15 μL PCR master mix composed of KAPA HiFi Fidelity buffer (3.3×); KAPA Biosystems), 1 mM dNTPs, and KAPA HiFi DNA Polymerase (0.03 U/μL; KAPA Biosystems) and 5 μL each of Nextera i5 and i7 PCR primers with barcodes (4 μM; Illumina, Inc.). PCR parameters were the following: 72 °C for 3 min, 98 °C for 10 s, 60 °C for 30 s, and 72 °C for 30 s, and a final 72 °C for 5 min. PCR products were purified with AMPure XP beads (0.8×; Beckman Coulter) followed by library normalization and pooling of up to 96 uniquely barcoded samples, as with Nextera XT (Illumina, Inc.).

Data Analysis. Sequencing data were processed by standard methods (10, 33) to obtain copy number profiles (25,000 bins of ~0.1 Mb). intD was calculated for each CNV call and compared with the threshold value for its size. CNV calls exceeding the cutoff were converted to integers corresponding to the threshold copy number value. As data were not normally distributed, nonparametric statistical comparisons were applied. All statistical testing was two-tailed, and multiple comparisons were adjusted with the Bonferroni–Holm method.

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