Neuronal DNA Content Variation (DCV) With Regional and Individual Differences in the Human Brain

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ABSTRACT

It is widely assumed that the human brain contains genetically identical cells through which postgenomic mechanisms contribute to its enormous diversity and complexity. The relatively recent identification of neural cells throughout the neuraxis showing somatically generated mosaic aneuploidy indicates that the vertebrate brain can be genomically heterogeneous (Rehen et al. [2001] Proc. Natl. Acad. Sci. U. S. A. 98:13361–13366; Rehen et al. [2005] J. Neurosci. 25:2176–2180; Yurov et al. [2007] PLoS ONE:e558; Westra et al. [2008] J. Comp. Neurol. 507:1944–1951). The extent of human neural aneuploidy is currently unknown because of technically limited sample sizes, but is reported to be small (Iourov et al. [2006] Int. Rev. Cytol. 249:143–191). During efforts to interrogate larger cell populations by using DNA content analyses, a surprising result was obtained: human frontal cortex brain cells were found to display “DNA content variation (DCV)” characterized by an increased range of DNA content both in cell populations and within single cells. On average, DNA content increased by ~250 megabases, often representing a substantial fraction of cells within a given sample. DCV within individual human brains showed regional variation, with increased prevalence in the frontal cortex and less variation in the cerebellum. Further, DCV varied between individual brains. These results identify DCV as a new feature of the human brain, encompassing and further extending genomic alterations produced by aneuploidy, which may contribute to neural diversity in normal and pathophysiological states, altered functions of normal and disease-linked genes, and differences among individuals. J. Comp. Neurol. 518:3981–4000, 2010.

INDEXING TERMS: aneuploidy; copy number variation; neural diversity; DNA; neuron; glia

The brain and virtually all somatic cells have been generally assumed to contain identical amounts of genomic DNA. The major exception to this assumption are lymphocytes that produce immunoglobulins or T-cell receptors, in part through DNA recombination processes (Schatz and Spanopoulou, 2005). The notion that somatic alterations in DNA sequence could also contribute to the complexity of the brain can be traced to theoretical explanations of immunoglobulin diversity by Dreyer and Bennett (1965), who proposed the existence of somatic DNA rearrangement in the immune system or, more specifically, what is now known as V(D)J recombination (Schatz and Spanopoulou, 2005). This proposal led to further speculation by Dreyer, Gray, and Hood that a similar kind of mechanism might occur within the brain, based in part upon the microscopic appearance of regenerating goldfish nerves from the retina into the optic tectum (Dreyer et al., 1967).

In the ensuing decades, plausibly rearranged genomic DNA loci have been examined, including those consisting of many homologous genes such as the protocadherins (reviewed in Chun, 1999) and olfactory receptors (reviewed in Chun, 2004); however, none have thus far shown evidence for DNA rearrangements. The absence of identified, rearranged loci contrasts with the documented

Additional Supporting Information may be found in the online version of this article.

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gene expression in the brain of multiple genes that are involved in immunological V(D)J recombination, and include RAG1 (Chun et al., 1991) as well as the nonhomologous end-joining (NHEJ) proteins Ku70, Ku80, XRCC4, and Lig4, the latter three of which were shown to disrupt normal brain development when genetically deleted (Chun and Schatz, 1999; Sekiguchi et al., 1999; Gu et al., 2000). The functions of these proteins in the brain remain unclear, but data exist to support DNA functions distinct from NHEJ (Sekiguchi et al., 2001).

Despite the failure to find bona fide DNA rearrangements in neural cells of the brain, the effects of NHEJ gene deletion on brain development suggest that other DNA alterations could be occurring. In particular, loss of NHEJ proteins like Ku80 has been associated with aneuploidy (Difilippantonio et al., 2000), which is the gain and/or loss of chromosomes to produce deviation from haploid multiples (Kingsbury et al., 2006). An extensive assessment of this possibility identified a surprisingly high degree of neural aneuploidy during mouse cerebral cortical development, as demonstrated by both chromosome counts as well as spectral karyotyping of mitotic neuroprogenitors. Consistent with these results, postmitotic neurons, as identified by fluorescence in situ hybridization (FISH) of selected chromosomal loci, could also be aneuploid (Rehen et al., 2001). During development, aneuploidy can be produced by known chromosomal segregation mechanisms (Yang et al., 2003), generating an apparently random assortment of aneuploidies that are predominantly losses (hypoploidy) along with rarer gains (hyperploidy) (Rehen et al., 2001; Yang et al., 2003; McConnell et al., 2004; Kingsbury et al., 2005; Rajendran et al., 2007; Westra et al., 2008). The functions of normally occurring neural aneuploidy are still being determined. However, it is associated with apparently physiological functions, with neurons integrated into neural circuitry (Kingsbury et al., 2005), and has also been shown to alter gene expression compared with non-aneuploid cells of the same kind (Kaushal et al., 2003).

Neural aneuploidy occurs to a similar extent in both neurons and glia (Rehen et al., 2005), is found phylogenetically from fish through humans (Rehen et al., 2001, 2005; Yurov et al., 2005; Kingsbury et al., 2006; Rajendran et al., 2007; Peterson et al., 2008), and is present throughout the neuraxis, with no obviously distinguishable neuroanatomical patterns (Rehen et al., 2001, 2005; Westra et al., 2008). Aneuploidy represents the first, proven form of DNA sequence alteration(s) present in normal neural cells, while making it likely that other phenomena associated with “genomic instability” could also be present. The creation of complex mosaics of intermixed aneuploid and euploid cells have implications for normal brain development, function, and disease (Kingsbury et al., 2006; Yurov et al., 2009).

A caveat in assessing aneuploidy for nonmitotic or postmitotic cells is the technical inability to interrogate the entire genome, i.e., all chromosomes, within single cells because there are no visible chromosomes present. Thus, formally, all studies utilizing FISH with subgenomic probes on cells without visible chromosomes leave open a range of alternative explanations for observed fluorescent signals, both legitimate and artifactual. Further complicating analyses of neural aneuploidy by FISH is its labor-intensive nature, which allows only a tiny sampling of the total number of cells within the brain to be assessed. To overcome these hybridization-based limitations, we have optimized DNA flow cytometry, particularly via use of isolated cell nuclei rather than whole cells, to assess the overall level of DNA content among cells of the brain. Here we report the methodologies, necessary controls, and results that demonstrate a new feature of human brain neurons, which, while encompassing aneuploidy, identifies distinct populations with increased range and/or amounts of DNA: we refer to this general property as “DNA content variation (DCV).”

MATERIALS AND METHODS

Human tissue samples

All human protocols were approved by the Human Subjects Committee at The Scripps Research Institute and conform to National Institutes of Health guidelines. Fresh-frozen postmortem human brain tissue from nondiseased control individuals was obtained from the Institute for Brain Aging and Dementia Tissue Repository (University of California, Irvine), the NICHD Brain and Tissue Bank for Developmental Disorders at the University of Maryland, Dr. Jeanne Loring at The Scripps Research Institute (TSRI), and the UK Parkinson’s Disease Brain Bank (Fig. 1). Human peripheral blood was obtained from healthy donors (aged 50–60 years) at TSRI Normal Blood Donor Services.

Flow cytometry (FCM) and fluorescence-activated cell sorting (FACS)

FCM and FACS were performed at TSRI Flow Cytometry Core Facility with a Becton Dickinson (San Jose, CA) LSRII and FACS-Aria, respectively. For DNA FCM, results were replicated by using three different LSRII machines. In addition, FCM results were verified and reproduced by independent researchers. Human brain nuclei were isolated as described previously (Westra et al., 2008), and fixed in ice-cold 70% ethanol for a minimum of overnight prior to FCM. Before FCM analysis, nuclei were filtered through 40-μm nylon mesh, washed in Ca2+/Mg2+-free
phosphate-buffered saline (PBS) supplemented with 2 mM EGTA (PBSE) and resuspended in PBSE with 50 mg/ml propidium iodide (PI; Sigma, St. Louis, MO), 50 μg/ml RNaseA (Sigma), and chicken erythrocyte nuclei or CEN (Biosure, Grass Valley, CA). Samples were incubated at room temperature for a minimum of 90 minutes prior to analysis. For FACS, nuclei were sorted into NeuN$^+$ and NeuN$^-$ nuclear fractions as described previously (Westra et al., 2008). For DNA content FACS, nuclei were labeled with a NeuN (1:100 dilution) primary antibody (Chemicon, Temecula, CA, cat. no. MAB377) and an Alexa Fluor 488 goat anti-mouse IgG secondary antibody (1:250 dilution, Invitrogen, Carlsbad, CA, cat. no. A1100-1).

The NeuN primary antibody used in this study has been well characterized (Lind et al., 2005; Desilva et al., 2007); it was raised in mice against purified cell nuclei from mouse brain and reacts with most neuronal cell types throughout the nervous system of mice including the cerebellum, cerebral cortex, hippocampus, thalamus, spinal cord, and neurons in the peripheral nervous system. The immunohistochemical staining is primarily in the nucleus of neurons, with light cytoplasmic staining. The few cell types not reactive with MAB377 include Purkinje and mitral cells. In immunoblotting of brain tissue from mice, this antibody recognized two to three bands in the 46-48-kDa range and possibly another band at approximately 66 kDa (manufacturer’s data). Immunohistochemical quality control experiments with the NeuN antibody revealed intense nuclear staining in sections of mouse brain, whereas staining was absent in HEK293 cell cultures.

Nuclei were stained with the DNA-binding dye DRAQ5 (Biostatus) for assessment of DNA content in NeuN-labeled nuclei. DRAQ5 exhibits high affinity and selective DNA binding and has not been shown to exhibit appreciable nonspecific binding (e.g., to RNA). Experimentally, DRAQ5 has been used to detect nuclear boundaries with high sensitivity, measure DNA content in leukocytes, and detect nuclear reorganization events (but is less sensitive than PI in detecting apoptotic nuclei by flow cytometry). As controls for flow cytometry, positive and negative selection gates were set up by using samples stained with secondary antibody alone, as well as DNA dye alone.

### Table: Human Tissue Samples

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Figure 1. Human tissues used in this study. Postmortem human brain samples from frontal cortex (FCTX) and cerebellum (CB), as well as peripheral blood samples (LYM [lymphocytes]) from nondiseased, consenting donors listed by sex (M or F), age, sample identifier, and postmortem interval (PMI). Tissues were obtained from the NICHD Brain and Tissue Bank for Developmental Disorders at the University of Maryland, Baltimore, MD (NICHD contract no. N01-HD-4-3368 and N01-HD-4-3383). The role of the NICHD Brain and Tissue Bank is to distribute tissue, and therefore it cannot endorse the studies performed or the interpretation of results. Additional samples were obtained from Dr. Jeanne Loring at The Scripps Research Institute (TSRI) and the UK Parkinson’s Disease Brain Bank. Human peripheral blood was obtained from the Normal Blood Donor Services at TSRI.
Fluorescence in situ hybridization

For quantitative FISH, we used a fluorescein-labeled PNA probe (5'-Flu-ATTGTTGAAAACGGGA-3') according to previous reports (Chen et al., 1999; Taneja et al., 2001). Briefly, sorted nuclei were dropped onto glass slides and hybridized for 1 hour at 37°C with PNA. Slides were washed in 2X SSC/50% formamide, 4X SSC/0.1% Tween-20, and 2X SSC for 2 minutes each at 45°C, and the nuclei were counterstained with DAPI. Fluorescence images were obtained by using a DeltaVision deconvolution system attached to an Olympus IX70 microscope (Applied Precision, Issaquah, WA). A series of Z-stacks at 0.5-μm intervals were taken through 5 μm of individual nuclei. The deconvolved image was projected into a single file and analyzed by using MetaMorph (Molecular Devices, Downingtown, PA) quantitation software. A minimum of 30 NeuN-labeled and NeuN-unlabeled nuclei from FACS was scored for PNA fluorescence integral intensity in four cortical brain samples. Images were saved in TIFF format, composed in Adobe Photoshop CS 8.0 (Adobe Systems, San Jose, CA), and adjusted for brightness and contrast.

Southern blotting

Genomic DNA was isolated from human postmortem brain tissue and lymphocytes by using the DNEasy Tissue Kit (Qiagen, Chatsworth, CA). For genomic samples, DNA was quantitated by reading the absorbance at 260 nm in a UV-1201 spectrophotometer (Shimadzu, Columbia, MD). Then 250 ng of genomic DNA was blotted onto a Nytran SPC nylon transfer membrane (Whatman, Clifton, NJ) and UV cross-linked for 2 minutes in a UV Stratalinker 2400 (Stratagene, La Jolla, CA) prior to probing the membrane. Membranes were hybridized with 32P-labeled DNA probes generated by random priming. Polymerase chain reaction (PCR) primers used to generate the LINE1 probe spanning ~650 bp (containing portions of ORF1 and ORF2) of the consensus human LINE1 sequence were 5'-TCAGGAAATACAGAGAACGCC-3' and 5'-TGACTTCCTTACCAGG-3'. Primers used to design the ALU probe spanning ~230 bp of the consensus human ALU sequence were 5'-GTGGCCTACGCCTGAATC-3' and 5'-GATCTC GGCTACGTCA-3'. Purified cDNA for APP and GPR84 were used as probe sequences. Probes for Southern blotting were labeled by random priming by using the Prime-It II Random Primer labeling kit (Stratagene). Signal intensity was determined by exposing 32P-hybridized membranes to a phosphor screen with subsequent scanning by using the Typhoon 8600 imager (Molecular Dynamics, Sunnyvale, CA).

Quantitative PCR

Real-time quantitative (q)PCR was used to quantify gene copy number from human brain and lymphocyte genomic DNA. The primer sets for LPA4 consisted of LPAF (5'-CACAGAAATCATGATGAAGG-3') and LPAR (5'-CTGGGTCTGAAGGCTGTTG-3'), and the primer set for SRY consisted of SRYF (5'-TTTGCAACTCTGGACCC TTTC-3') and SRYR (5'-CTGTTTCTCCGTTACAC-3'). Primers were synthesized by Integrated DNA Technologies (Coralville, IA). qPCR was performed on a Rotor-Gene RG-3000 72-well thermocycler (Corbett Research, Cambridge, UK) by using SYBR Green JumpStart Taq Ready-Mix (Sigma). All reactions were performed in triplicate. Standard curves for quantifying gene copy number were determined by using purified plasmid DNA (pGemT) containing a single copy of the gene of interest. For copy number quantitation, serial dilutions of plasmid DNA were used to construct standard curves for LPA4 (R2 = 0.9961) and SRY (R2 = 0.9972). DNA concentrations were converted to gene copy number by calculating the weight (in g/mol) of the plasmid and insert used for generating the standard curves, and converting this into copy number (g/molecule) by using Avogadro's number (mol/molecule).

Statistical analysis

For DNA content FCM, DNA indices were determined as described in the Results section. P values (for DNA indices, and coefficient of variation [CV] and “skewness” [SKW] values of cortical and cerebellar populations) were determined by using an unpaired, two-tailed t-test. For qPCR analysis, copy numbers were determined by using the standard curve method. For NeuN FACS and PNA FISH experiments, P values were determined by using an unpaired, two-tailed t-test. Differences between groups were considered statistically significant with P < 0.05.

RESULTS

To assess DNA content from cells of the human brain, a modified protocol for measuring DNA content by using flow cytometry (FCM) and fluorescence-activated cell sorting (FACS) was developed that relied on isolation of intact cellular nuclei to eliminate difficulties associated with the brain’s interconnected cells (Levitt et al., 1997). Control procedures and experiments were pursued to eliminate trivial explanations or artificial data and included consideration of nuclear RNA/RNase treatments, varying nuclei density, autofluorescence, discrimination of intentionally mixed nuclei populations, internal DNA standards, PI concentrations and staining protocols, and reproduction of the results with a chemically and mechanistically distinct DNA binding dye, DRAQ5 (Smith et al., 2000). DRAQ5 is a UV-excitable, membrane-permeable dye that binds stoichiometrically to DNA and has been used to detect DNA content in leukocytes by FCM.
Yuan et al., 2004; Swerts et al., 2007). Nuclear DNA content of isolated, fixed human lymphocyte nuclei (LYM, n = 8) stained with PI (Crissman and Steinkamp, 1973) recapitulated whole-cell FCM (data not shown), and this preparation served as a euploid, internal reference sample along with a second standard, chicken erythrocyte nuclei (CEN) (Vindelov et al., 1983b). These standards were compared with large cohorts (minimum of 10,000 nuclei per sample) of nondiseased human brain cell nuclei from the cerebellum (CB; n = 8) and frontal cortex (FCTX, n = 24) of different individuals controlled for age and sex (Fig. 1). The total number of nuclei assessed in this study represents several orders of magnitude greater coverage than any previously published study assessing genomic variation in the central nervous system (CNS).

As anticipated, lymphocyte and cerebellar samples showed nearly identical, superimposable DNA content histograms with smooth, sharp peaks and thin bases (Fig. 2a,b). Surprisingly, and in marked contrast, DNA content histograms from frontal cortical cell nuclei of different individuals (Fig. 2c) deviated in shapes, peaks, and base widths. These differences were apparent in histogram (Yuan et al., 2004; Swerts et al., 2007). Nuclear DNA content of isolated, fixed human lymphocyte nuclei (LYM, n = 8) stained with PI (Crissman and Steinkamp, 1973) recapitulated whole-cell FCM (data not shown), and this preparation served as a euploid, internal reference sample along with a second standard, chicken erythrocyte nuclei (CEN) (Vindelov et al., 1983b). These standards were compared with large cohorts (minimum of 10,000 nuclei per sample) of nondiseased human brain cell nuclei from the cerebellum (CB; n = 8) and frontal cortex (FCTX, n = 24) of different individuals controlled for age and sex 

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overlays comparing frontal cortical nuclei with lymphocyte nuclei (Fig. 2d). Moreover, cerebellar DNA content histograms, from the same brain as the examined frontal cortex, were themselves distinct, resembling lymphocyte histograms. Cortical histograms typically showed peaks with additional right-shoulders and widened bases, indicative of increased DCV. A top-down orthogonal view of the histograms (Fig. 2e) provides similar perspectives (as in Fig. 2d).

The DNA index (DI) (Darzynkiewicz and Huang, 2004) of each sample was calculated, defined as the ratio of the brain G0/G1 peak to that of the lymphocyte G0/G1 peak, normalized to an internal control (CEN). The G0/G1 peak is the main peak (Fig. 3c,g,k) containing cells (or nuclei) in the G0/G1 phase of the cell cycle. The average DI of human cortical cell nuclei was always higher than either lymphocyte or cerebellar samples (1.04 in cortical nuclei compared with 0.98 and 1.0 for cerebellar and lymphocyte samples, respectively, \( P = 0.002 \) for FCTX vs. LYM and \( P < 0.0001 \) for FCTX vs. CB; Fig. 4a,c). Cortical samples also exhibited greater CVs (Fig. 4b). Deviation of each DNA content histogram from a Gaussian distribution

![Image of DNA index calculation](image-url)
Figure 4. Quantitative analysis of DNA content in nondiseased human nuclei. a: Individual DNA indices for human lymphocyte (LYM), cortical (FCTX), and cerebellar (CB) nuclei samples. The DNA index was determined by comparing the mean DNA content value for brain nuclei with the average of the sex-matched lymphocyte samples to obtain a ratio. b: Higher coefficient of variation (CV) values were seen in human cortical samples (6.18 mean, 3.43–9.90 range) compared with cerebellar (3.86 mean, 2.20–6.69 range), and lymphocyte (2.73 mean, 2.20–3.25 range) samples ($P < 0.0001$). c: Average DI for each sample group (1.0 for lymphocytes, 0.98 for cerebellum, and 1.04 for frontal cortex). The cortical group had a higher mean DI than both lymphocyte and cerebellar sample groups ($P = 0.0002$). d: Human cortical samples show increased skewness (SKW) values (0.516 mean) compared with lymphocyte (−0.125 mean) and cerebellar samples (−0.106 mean) ($P < 0.0001$). SKW values were determined by using the formula $\text{SKW} = \frac{\text{mean DNA content}}{\text{mode DNA content}}/\text{standard deviation}$ for the “diploid” peak of each sample. Individual cortical samples with high SKW values displayed prominent right histogram shoulders (Fig. 2c). Whereas lymphocytes and cerebella SKW values were near 0 or negative, 23/24 cortical samples had a positive SKW value. e,f: Comparative autoradiography of genomic Southern blots from different, human nuclear populations. Human cortical (n = 20) and cerebellar (n = 8) genomic DNA samples probed with an L1 repeat element produced a respective 4.4- ($P < 0.05$) and 4.6-fold ($P < 0.01$) increase in the ratio of L1 repeat sequences compared with single-copy gene (GPR84, chromosome 12) probing of the same DNA samples, contrasting with human lymphocyte DNA (n = 7). A similar result was obtained by using a different single-copy gene (LPAR4, X chromosome). The similar cerebellar and cortical ratios supported a genomic origin for cortical DCV. g: Quantitative PCR (qPCR) analysis of sex chromosome loci from human lymphocyte, cerebellar, and cortical genomic DNA. The relative gene copy number of LPAR4 and SRY (on the Y chromosome) was determined from male lymphocytes (set to a value of 1.0 as a reference) and cortices (FCTX #1–7). Relative gene copy numbers of LPAR4 and SRY ranged from 0.47 (SRY in sample #2) to 1.77 (LPAR4 in sample #1) in cortical samples, with the average SRY copy number being 0.98, compared with the LPAR4 copy number of 1.25. For a–g: *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$. DNA content variation (DCV) in human neurons.
was calculated as SKW, defined as the \([\text{mean DNA content} - \text{mode DNA content}] / \text{standard deviation for each sample}\) (Aldhous et al., 1994). Whereas lymphocyte and cerebellar samples contained near-zero or negative SKW values, cortical samples showed predominantly positive SKW values indicating increased DNA content (Fig. 4d). The average increase in DNA content between the frontal cortex and lymphocytes was \(\sim 4\%\) (with a range of \(-2\%\) to \(9\%\)) of an approximately 6,000-Mb, 2N diploid genome, or \(\sim 250\) Mb. By comparison, a 3N signal would have produced a 50% increase in DNA content (an additional 3,000 Mb) which was never observed, whereas 4N peaks of 100% increase were also observed, consistent with the known presence of mitotic, non-neuronal brain cells.

Hypothesized genomic origins of the increased cortical DNA were assessed by Southern blotting of L1 (LINE1) repeat sequences, relative to two single-copy genes, \(GPR84\) on chromosome 12 and \(LPAR4\) on the X chromosome (Venter et al., 2001). L1 repeats are interspersed throughout the genome (Kazazian and Goodier, 2002). Therefore, genomic origins of DCV would be expected to maintain relative L1 copy number representation rather than reducing it by dilution with non-L1-containing DNA as would occur with viral or mitochondrial sources that do not contain L1 repeat elements. By using this approach, a consistent level of repeat sequence representation was observed for L1 in cortical samples, arguing against massive, nongenomic contributions to DCV. Interestingly, the representation of L1 sequences in cerebellar samples was indistinguishable from that of frontal cortical samples, both of which were significantly greater than lymphocyte samples, even though similar global levels of DNA content were present in cerebellar vs. lymphocyte samples as identified by FCM (Fig. 4e,f). These data further suggested qualitative differences in DCV for lymphocytes as compared with neural nuclei, based upon relative L1 representation. qPCR analysis further revealed increased single-copy gene signals within cortical samples relative to lymphocytes (Fig. 4g), supporting the presence of subgenomic increases of DNA loci relative to single-copy genes (shown for the genes \(LPAR4\) and \(SRY\) from male samples). Of note, there was no appreciable correlation between age and DI in our samples, analyzed within tissue regions, or all together (Fig. 5).

One possible artifact that could contaminate data collected by FCM/FACS is autofluorescence, particularly in analyzing aged neurons, within which lipofuscin is known to accumulate. Lipofuscin is a broad-spectrum autofluorescent pigment seen in both rhodamine and fluorescein channels that accumulates, particularly at the nuclear periphery in some aging neurons (Riga and Riga, 1995). This possible contribution of lipofuscin to FCM/FACS signals was assessed (Fig. 6). No relevant signal from unstained nuclei (Fig. 6p, unstained) was observed in the PI (or DRAQ5, data not shown) channel, demonstrating that the presence of lipofuscin does not contribute to fluorescence signals detecting DNA content in human brain nuclei. A related control examined the effect of different DNA binding dyes on DNA content (Fig. 7) by comparing PI with the DNA minor groove binding dye DRAQ5 (Smith et al., 2000). Labeling nuclei with both dyes produced similar right-hand shoulders in the frontal cortical samples while appropriately labeling internal controls. The identification of cortical DCV with DRAQ5 also further ruled out any contribution of lipofuscin autofluorescence to DNA content signals, because lipofuscin is minimally fluorescent in the far-red spectra (633 nm) that was used to excite DRAQ5 (Dowson, 1982).
Figure 6. The effect of lipofuscin autofluorescence on human brain DNA content histograms. Lipofuscin is a broad-spectrum autofluorescent pigment seen in both rhodamine and fluorescein microscope channels that accumulates at the nuclear periphery in some aging neurons and might contribute autofluorescent signals in DNA content flow cytometry experiments (Riga and Riga, 1995). This possibility was assessed on nuclei with low lipofuscin (a–d; Lipo−) vs. high lipofuscin (e–h; Lipo+) pigment levels, stained with PI (red); Lipo+ nuclei were visible in a fluorescein channel (green). Autofluorescence from lipofuscin can be minimized with the addition of the lipophilic dye Sudan Black (i–l; SB), which preferentially binds to lipofuscin granules (black particles in k and l) (Schnell et al., 1999) and quenches the autofluorescent signal, and then is visualized by brightfield microscopy. SB staining was used in initial experiments, however, further analyses demonstrated that it was not necessary (m–o); the addition of SB to human brain nuclei prior to FCM did not significantly alter DNA content histograms (SB− [red histogram], SB+ [black histogram], and overlay). Importantly, when unstained isolated nuclei were analyzed by FCM, no signal from lipofuscin alone (p, unstained) was observed in the PI (or DRAQ5; data not shown) channel, indicating that the presence of lipofuscin does not contribute to fluorescence signals in DNA content FCM of human brain nuclei. CEN, chicken erythrocyte nuclei. Scale bar = 5 μm in a–l. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
A striking difference was identified in comparing DNA content histograms between different brain regions of the same individual (Fig. 8a,b). DNA content was analyzed pair-wise in seven individuals for cortical vs. cerebellar samples. All samples were run blind to identity. Each resultant DNA content histogram was quantitatively distinct, most notably within cortical samples, with distinguishable histogram shapes that further underscored differences between the cortex and cerebellum; six of seven samples showed clearly increased cortical DNA (Fig. 8c) as marked by a broad, right-hand domain (green shading in samples #4 and #5 in Fig. 8d,f) consisting of nuclei with higher DNA content. These data demonstrated that DCV can vary between neuroanatomically and neuroembryologically different regions of the same brain.

Neurons are postmitotic and phenotypically distinct from glial/other non-neuronal cells in the brain. Initial studies using backgating analysis to determine the size of nuclei within the cortical histogram shoulders indicated that nuclei with increased DNA content were generally larger (Fig. 10), yet controls for nuclear size using intermixed large and small nuclei derived from whole human blood (Fig. 9) and a DNA content comparison of comparably sized human cortical, cerebellar, and lymphocyte nuclei (Fig. 10) demonstrated that DNA content was independent of nuclear size. These results suggested that neurons, known to have larger nuclei among cortical cells (Dombrowski et al., 2001; Knusel et al., 1973), might be enriched in populations with increased DNA content. To determine whether DCV differed in this postmitotic population, FCM and NeuN (a neuron-specific nuclear antigen) immunolabeling were combined to analyze neurons. NeuN+ nuclei were found about twice as often in populations of nuclei sorted from the right shoulders of histograms as compared with unsorted nuclei (Fig. 11). FACS of NeuN+ nuclei further identified an increased DI of NeuN+ relative to NeuN− nuclei in all samples analyzed (Fig. 12) (Mullen et al., 1992). In general, neuronal nuclei tended to be larger than non-neuronal nuclei. However, even when taking nuclear size into account by analyzing NeuN+ and NeuN− nuclei with similar dimensions, the DI of NeuN+ nuclei was still higher than that of NeuN− nuclei (Fig. 13).

To further assess a genomic origin for DCV, quantitative fluorescence of fluorescently labeled peptide nucleic

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**Figure 7.** The effect of distinct DNA binding dyes on DNA content histograms. Isolated human lymphocyte (LYM) and cortical nuclei (FCTX) spiked with chicken erythrocyte nuclei (CEN) and stained with the DNA intercalating dye propidium iodide (PI; a–c) or the DNA minor groove binding dye DRAQ5 (d–f) (Smith et al., 2000) were analyzed by FCM. A similar right-hand shoulder in the frontal cortical sample was seen by using both dyes in either an orthogonal, top-down view (orange arrowheads in b and e) or a histogram view (green shading in c and f). Similar results were seen with all other paired samples examined, indicating that similar DCV patterns are seen with distinct DNA binding dyes having different DNA binding characteristics. The identification of cortical DCV with DRAQ5 also further ruled out any contribution of lipofuscin autofluorescence to DNA content signals, because lipofuscin is minimally fluorescent in the far-red spectra (633 nm) used to excite DRAQ5 (Dowson, 1982). [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
acid (PNA) probes against human centromere repeats (CENP-B box) that are structural DNA sequences on chromosomes (Masumoto et al., 2004) was employed. NeuN+ nuclei showed higher levels of CENP-B fluorescence than non-neuronal nuclei (average of 25%, n = 4), supporting increased prevalence of chromosomal organization elements, consistent with increased, genomic DNA content (Fig. 14). These data indicated that, on average, human neurons have more genomic DNA than non-neuronal cells.

**DISCUSSION**

This study identifies a new feature of the human brain: cells with DNA content variation (DCV). It exists on a large scale and manifests most prominently as increased DNA content and range in neurons compared with
non-neuronal cells, with greater DCV in cells from the frontal cerebral cortex compared with the cerebellum of the same brain. DCV also appears to be distinct among individuals. The methodologies used to determine DCV are well established and in current use to assay DNA content in a wide range of cell lines and types, and were optimized here for the adult human brain. Classical FCM/FACS for DNA content requires single-cell suspensions (Nunez, Figure 10. DNA content analysis of similarly sized nuclei from different brain regions and lymphocytes by FCM. Backgated fractions of similarly sized (boxed regions) human nuclei from lymphocytes (LYM; green, a–c), frontal cortex (FCTX; blue, d–f), and cerebellum (CB; red, g–i) stained with propidium iodide and analyzed by FCM. Backgates were drawn from dot plots of nuclear size (forward scatter [FSC]) and granularity (side scatter [SSC]) (Brasseur et al., 1984) in a, d, and g. Lymphocytes and cerebellar nuclei show overlapping DNA content histograms, whereas similarly sized cortical nuclei show increased DNA content in the orthogonal, top-down views (red vertical line from population mode in c, f, and i) and overlay view (j). These data indicate that nuclear size does not affect DCV. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
however, the adult brain does not lend itself to such suspension, reflecting in part its high degree of complex interconnectivities that have prevented prior, extensive DNA content analyses of intact brain cells by FCM or FACS. The use of isolated cell nuclei as well as the use of chemically distinct DNA dyes (PI and DRAQ5) produces the same general results, arguing in toto against non-DNA content explanations for DCV. Critically, the multiple controls used here, which include lymphocytes and CEN, are consistent with the prior literature on

Figure 11. Backgating analysis and immunolabeling of human cortical nuclei. Histograms with a prominent DCV shoulder (a) or no-shoulder (d) from human cortical nuclei stained with propidium iodide were backgated on the basis of DNA content fluorescence into lower 60% (LO) and upper 20% (HI) fractions. HI and LO fractions were then analyzed for physical nuclear characteristics (plotted as nuclear size (FSC) against nuclear granularity (SSC) (Brasseur et al., 1984). Backgating analysis of these populations revealed that HI nuclei (b,e) were generally larger and more granular than LO nuclei (c,f) in histograms with shoulders and normal histograms (91% large/high granularity nuclei and 2% small nuclei/low granularity in HI [b] vs. 7% large/high granularity nuclei and 89% small/low granularity nuclei in low [c] for the histogram with a shoulder). Percentages refer to the percent of the backgated nuclei that fall within the large or small nuclear populations. g: FACS was used to isolate the upper 20% of the nuclei based on DNA content from propidium iodide-stained cortical nuclei. h-i: FACS-isolated nuclei were then immunolabeled for the neuronal marker NeuN (green). The percentage of NeuN-immunolabeled nuclei in the upper 20% was 1.9-fold that of the percentage of unsorted nuclei (j; n = 4, P = 0.0101). These results indicate that at the upper end of the DNA content histograms in cortical samples, there are more neurons, which tend to be large and granular. Scale bar = 10 µm in h,i. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
DNA content analyses, demonstrating that DNA content was indeed detected with the same fidelity as in previous studies (Laerum and Farsund, 1981; Vindelov et al., 1983a; Danova et al., 1987; Berger et al., 1996; Darzynkiewicz and Huang, 2004; Muehlbauer and Schuler, 2005). Moreover, assessments of repeat element hybridization, qPCR analyses, and CENP-B fluorescence, each independent of FCM/FACS, are consistent with genomic origins of increased DCV.

Figure 12. DNA content analysis of NeuN immunolabeled nuclei from the non-diseased frontal cortex. a–c: Human brain nuclei were isolated from postmortem samples, immunolabeled for NeuN, and stained with the non-spectrally overlapping DNA dye DRAQ5. All DRAQ5-positive nuclei (a) were gated on NeuN immunoreactivity (c) and analyzed for DNA content. Gates (horizontal lines in the top two histograms) for the NeuN+ and NeuN− populations were determined by using nuclei exposed to the secondary antibody alone (b; full gating parameters are shown in Fig. 3). Numbers above the gates represent the percentage of nuclei falling within the gate. d: NeuN-positive nuclei (green histogram) had higher DNA contents than NeuN-negative nuclei (blue histogram); however, both histograms fell within the un gated DRAQ5− population (red histogram). Note: A representative example is shown in a–d; n = 6 samples analyzed in total. e: Quantitative analysis of the DNA content of NeuN+ nuclei compared with NeuN− nuclei for the six samples analyzed (NeuN− samples set to 1.0 as a reference) revealed a ~33% increase in DRAQ5 intensity in NeuN+ nuclei (black bars), with every sample showing increased DNA content in neuronal nuclei relative to non-neuronal nuclei (white bars). [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]

DNA content analyses, demonstrating that DNA content was indeed detected with the same fidelity as in previous studies (Laerum and Farsund, 1981; Vindelov et al., 1983a; Danova et al., 1987; Bruner et al., 1993; Berger et al., 1996; Darzynkiewicz and Huang, 2004; Muehlbauer and Schuler, 2005). Moreover, assessments of repeat element hybridization, qPCR analyses, and CENP-B fluorescence, each independent of FCM/FACS, are consistent with genomic origins of increased DCV.

Historically, studies using less sensitive techniques have reported polyploidy in Purkinje neurons (Lapham, 1968), although modern techniques have not substantiated these early reports (Del Monte, 2006). The subgenomic increase in DNA content, averaging ~250 Mb, observed here in the frontal cortex has not been previously reported and is inconsistent with polyploid DNA content. The origin of the additional DNA in cortical nuclei appears to be genomic; although we cannot eliminate all nongenomic sources such as unidentified DNA viruses, the high viral copy numbers needed to account for the ~250 Mb that affects large populations of normal cells make this mechanism seem unlikely. In a similar vein, active L1 retrotransposition has been reported in rat neuroblasts (Muotri et al., 2005) and human neural progenitor cells (Coufal et al., 2009), which could theoretically increase DCV. However, active L1 retrotransposition has not thus far been shown to occur in human postmitotic neurons, whereas the small size of an active retrotransposon (~6 Kb), similar to a retroviral provirus, would require an additional 40,000+ copies/neuron to produce a 250-Mb DNA content increase. It is notable that different levels of L1 elements have been reported among tissues of the same individual, including increased representation in the frontal cortex compared with the cerebellum (Coufal et al., 2009), consistent with a previous report of a similar LINE-like element that showed increased brain representation over other tissues (Yokota et al., 1989), and having an
unclear relationship to active retrotransposition. Although the formal relationship between DCV and possibly active retrotransposition of L1 remains uncertain, the L1 levels determined here (Fig. 4e,f) were not distinguishable between the frontal cortex and cerebellum, despite their marked differences in DCV, which argues against L1 elements as a primary source of increased DCV in the frontal cortex.

Prior studies on the ploidy of mouse lymphocytes indicated that this population has ~10% of the aneuploidy observed in neural progenitor cells (Rehen et al., 2001), suggesting a distinction between these mitotic cell types in their ability to tolerate genomic variation involving chromosomal gains and/or losses. A related cell lineage effect may in part explain the DCV differences observed here between human peripheral blood lymphocytes and postmitotic neurons from the cerebral cortex. In view of the DCV similarities between human lymphocytes and cerebellar neurons, this difference is not likely to reflect the postmitotic state; however, postmitotic cortical neurons may be able to tolerate and presumably utilize DCV throughout the life of the cell compared with mitotic lymphocytes or the neuroembryologically distinct neurons of the cerebellum.

There have been previous, controversial reports of cell cycle reactivation with DNA synthesis in neurons of adult mammals in both normal and diseased settings (Gould et al., 1999; Yang et al., 2001); however, our data do not support classical cell cycle-mediated events (Storchova and Pellman, 2004) in view of the ~4% (250 Mb) average increase in DNA content seen in frontal cortical neurons. Furthermore, studies measuring radioactive carbon in neuronal cells from postmortem brain samples of individuals exposed during neurogenesis to atomic bomb testing did not detect an appreciable amount of de novo DNA synthesis in neurons (Spalding et al., 2005; Bhardwaj et al., 2006). However, it is notable that the frontal cortex was not examined in these studies, while the possibility of subgenomic DNA synthesis might be below the detectable limits of such approaches. Postnatal nucleotide incorporation has been reported in the primate frontal cortex and is formally consistent with subgenomic, de

Figure 13. Nuclear size and DNA content in NeuN-labeled cortical nuclei. Nuclear fractions of similar size (magenta boxes in a and b) of NeuN-negative (a magnified view of the nuclear fraction within the magenta box is shown in c and d for NeuN-negative and NeuN-positive nuclei, respectively) and NeuN-positive (e–g) DRAQ5-stained human cortical nuclei analyzed for NeuN labeling (f) and DNA content (g). Similarly sized nuclei from a and b maintained separation of the NeuN label (FITC in f), whereas NeuN-positive nuclei (green histogram in g) contained more DNA than similarly sized NeuN-negative nuclei (black histogram in g). These data argue that even in a population of similarly sized nuclei, neurons contain more genomic DNA than non-neuronal nuclei. FSC, forward scatter; SSC, side scatter. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
novo DNA synthesis in neurons that might be distinct from neurogenesis (Gould et al., 1999; Rakic, 2002). This possibility requires further assessment.

Aneuploidy, defined as a chromosome complement that deviates from haploid multiples (Rajagopalan and Lengauer, 2004), can be produced during neurogenesis.
and is prevalent in the vertebrate brain, where it might theoretically account for major DCV forms. Indeed, this is likely to be an important subset of DCV, because it exists throughout the neuraxis (Rehen et al., 2001; Westra et al., 2008). However, aneuploidy is unlikely to account for the major DCV differences producing the increased DNA content observed in the human frontal cortex because hypoploidy (chromosome and thus DNA loss) is the most common form of demonstrated neural aneuploidy (Rehen et al., 2001; Yang et al., 2003; Yurov et al., 2007), which contrasts with the observed average increase in DNA content in the frontal cortex. As notable is the existence of aneuploidy throughout the neuraxis in both neurons and glia, which contrasts with the increased representation of DCV in neurons. Moreover, neural aneuploidy has been documented in all examined vertebrates, whereas the extent of DCV in mouse tissues is not of the same magnitude as that detected by the same methodologies used to identify human DCV (Fig. 15). Therefore, the combination of increased DNA content, species variability, neuronal enrichment, and neuroanatomical variation does not support neural aneuploidy as a primary mechanism for the increased DCV observed here in the human frontal cortex.

Possible DCV functions include contributions to neural diversity and brain function by altering gene availability for transcriptional events or micro-RNAs (Rehen et al., 2001; Torres et al., 2007; Guarneri and DiLeone, 2008). We speculate that DCV within single brains may also contribute to sporadic disease processes that preferentially affect distinct brain regions, as observed in Alzheimer’s disease, in which neurodegeneration predominantly affects the hippocampus and entorhinal/frontal cortices, with relative sparing of the cerebellum (Goedert and Spillantini, 2006). It is also possible that DCV may represent cells with disease latency or potentiality that is manifested only after other conditions, e.g., injury, stress, distinct disease insults, aging, initiate or promote detectable disease through DCV elements. Alternatively, DCV might represent a reactive response to disease processes toward counteracting insults, although its prevalence in apparently normal brains might require redefining what is normal. These non-mutually exclusive possibilities remain to be experimentally assessed.

A full characterization of DCV requires further study, and thus far appears to be most notable as a gain of genomic DNA in human frontal neurons based upon the maintained representation of LINE and ALU sequences, as well as centromere signals in DCV samples, which are all indicative of genomic DNA. We speculate that this increased DNA content reflects a combination of both large-scale genome amplification (e.g., equivalent to many megabases of contiguous DNA sequence that could be duplicated one or more times) and more localized amplifications, as seen in copy number variations (CNVs) (Freeman et al., 2006). Altered functions of mutant and wild-type genes linked to brain disease might occur in cells with differing degrees of DCV, that could also produce alterations akin to CNVs (Sebat et al., 2004), yet doing so in a localized, somatic manner, particularly within single cells, distinct from germline or zygotic mechanisms. CNVs could well be represented within DCV, yet would also differ compared with current CNV models. These differences could include: 1) CNVs that would not be present in all somatic cells vs. the current understanding of CNVs (Freeman et al., 2006); 2) mosaic cell populations that would consist of intermixed CNV and non-CNV or variable CNV-containing cells; 3) lineage-specific (e.g., neuronal vs. glial) CNVs; and 4) region-specific CNVs (e.g., cortex vs. cerebellum). Whether CNVs are indeed present within cells possessing DCV requires further analyses that must also involve next-generation DNA sequencing at a single-cell level. Additionally, as a subgenomic, DNA-altering phenomenon, it would not be surprising to find related alterations within DCV sequences (rearrangements, deletions, de novo sequences) that could emerge from further analyses. Thus, future DCV studies analyzing...
DNA sequences, neuroanatomical distributions, lineage relationships, and functions may provide new insights into brain diseases (particularly sporadic forms) as well as neural and behavioral diversity within and among individuals.

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