

# Dosage-dependent phenotypes in models of 16p11.2 lesions found in autism

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Recurrent copy number variations (CNVs) of human 16p11.2 have been associated with a variety of developmental/neurocognitive syndromes. In particular, deletion of 16p11.2 is found in patients with autism, developmental delay, and obesity. Patients with deletions or duplications have a wide range of clinical features, and siblings carrying the same deletion often have diverse symptoms. To study the consequence of 16p11.2 CNVs in a systematic manner, we used chromosome engineering to generate mice harboring deletion of the chromosomal region corresponding to 16p11.2, as well as mice harboring the reciprocal duplication. These 16p11.2 CNV models have dosage-dependent changes in gene expression, viability, brain architecture, and behavior. For each phenotype, the consequence of the deletion is more severe than that of the duplication. Of particular note is that half of the 16p11.2 deletion mice die postnatally; those that survive to adulthood are healthy and fertile, but have alterations in the hypothalamus and exhibit a “behavior trap” phenotype—a specific behavior characteristic of rodents with lateral hypothalamic and nigrostriatal lesions. These findings indicate that 16p11.2 CNVs cause brain and behavioral anomalies, providing insight into human neurodevelopmental disorders.

Home-cage | stereotypic behavior | structural variation | brain MRI

Accumulating evidence suggests the importance of copy number variations (CNVs) in the etiology of neuropsychiatric disorders, including autism (1), schizophrenia (2–4), developmental delay (5), and other complex traits (6). The 16p11.2 region is particularly intriguing. Whereas deletion of 16p11.2 has been associated with autism (7–9), duplication of 16p11.2 has been associated with autism (9, 10) as well as schizophrenia (11). 16p11.2 CNVs have also been reported in patients with developmental delay, mental retardation, repetitive behaviors (12–16), and a highly penetrant form of obesity (17). A reciprocal effect of 16p11.2 dosage on head size has been noted, as deletions are associated with large head size or macrocephaly, whereas duplications are associated with microcephaly (16). These studies reveal the variability of symptoms in patients carrying the same 16p11.2 CNV, an extreme example being a family with three affected members with symptoms so heterogeneous that they were barely overlapping (18).

Mouse models allow direct assessment of CNVs while reducing variability caused by genetic and environmental factors. We and others have previously used chromosome engineering (19) to model genetic alterations found in complex human diseases including cancer (20) and genomic disorders (21–24), allowing identification of the causative gene and elucidation of the mechanism involved (20, 25–27). Here we used a similar approach to generate mouse models with deletion and duplication corresponding to those found in patients with 16p11.2 CNVs. Because of the evidence for clinical heterogeneity, we screened these models for multiple changes in brain anatomy and behavior by using a combination of high-resolution MRI (28) and a monitoring system that assesses multiple behaviors (29). We found that the deletion and the duplication affect behavior and brain anatomy in opposing ways, with deletion mice exhibiting behaviors that resemble sensorimotor deficits in rats with lateral hypothalamic and nigrostriatal lesions (30, 31).

These findings provide evidence that brain anatomy and behavior depend on dosage of the region corresponding to 16p11.2.

## Results

**Generation of Mouse Models for Human 16p11.2 CNVs.** We asked whether altered dosage of the region corresponding to 16p11.2 causes abnormalities in mice. Genes mapping to the 0.52-Mb 16p11.2 CNV in humans cluster within a 0.44-Mb region of mouse chromosome 7 (Fig. 1A). Using chromosome engineering (19) as we have previously (20, 27, 32), we generated mice with one copy [heterozygous for a deletion or deficiency (*df*) allele], as well as mice with three copies [heterozygous for a duplication (*dp*) allele] of the region corresponding to 16p11.2 (Fig. 1B and Fig. S1). Endpoints for the rearrangement were selected based on human data (1), with each gene in the interval being conserved in mouse (Dataset S1). Gene targeting constructs were generated using MICER (33), and sequential targeting in mouse ES cells resulted in integration of *loxP* sites and selection cassettes at each endpoint (Fig. 1B and Fig. S1). Cre-mediated recombination and drug selection within eight independent doubly targeted clones revealed that three clones had been targeted in *cis* and five clones had been targeted in *trans*, which generated *df/+* and *df/dp* ES cells, respectively (Fig. 1B and Figs. S1 and S2). Five independent *df/dp* clones were used for blastocyst injection, producing 40 different male chimeras that were crossed to *+/+* females. Ten of these chimeras (representing two independent ES cell clones) produced *df/+* and *dp/+* mice that were identified by PCR (Fig. 1C). This approach provides mouse models for directly assessing the consequences of both the 16p11.2 CNV losses (i.e., deletion) and gains (i.e., duplication) found in humans.

We established both *df/+* and *dp/+* mice, but at weaning we noticed that *df/+* mice were underrepresented and litter sizes were smaller than expected (Table S1). Before weaning, *df/+* mice were sometimes small (Fig. 1D), but as adults, they were essentially the same size as their siblings and appeared healthy (SI Experimental Procedures). To determine whether *df/+* mice were dying during embryogenesis, we crossed *df/+* males to *+/+* females, and harvested embryos at day 13.5 of development [i.e., embryonic day (E) 13.5] as well as just before birth (E17.5–E18.5); progeny from similar crosses using the same studs as well as their male siblings were also genotyped at weaning (Table S1). Whereas litter sizes during embryogenesis averaged 9.4 embryos and the ratio of *df/+* embryos was Mendelian, litter sizes at weaning averaged only 5.0 mice and the ratio of *df/+* mice was half that expected. In addition, litter sizes were normal and *df/+*

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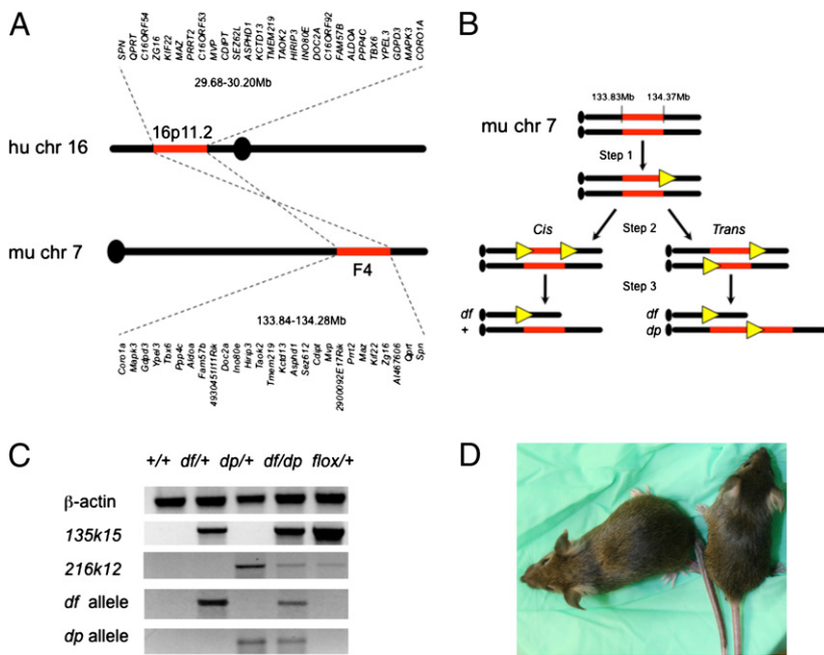
The authors declare no conflict of interest.

Data deposition: The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, [www.ncbi.nlm.nih.gov/geo](http://www.ncbi.nlm.nih.gov/geo) (accession no. GSE32012).

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**Fig. 1.** Generation of 16p11.2 models. (A) Genes mapping to human 16p11.2 CNVs are conserved in mouse. (B) Schematic of the chromosome engineering strategy used to generate mouse models of 16p11.2 CNVs. Step 1 is gene targeting at the *135k15* locus; step 2 is gene targeting at the *216k12* locus in the *135k15*-targeted ES cells; and step 3 is Cre-mediated recombination. *Cis* and *trans* indicate that *loxP* sites (yellow triangles) had integrated on the same or different chromosome homologues, respectively. (C) Molecular validation. PCR products using primers specific for the positive control ( $\beta$ -actin), targeting at the first and second endpoints (*135k15* and *216k12* loci, respectively), the *df* allele, and the *dp* allele are shown. (D) Before weaning, *df/+* mice (Right) tend to be smaller than their *+/+* siblings (Left; 8.8 and 15.4 g, respectively, for the females shown). Note the light-colored tail and ears of the *df/+*, which is a result of the presence of the *Agouti* transgene. In A and B, chromosome positions are shown in megabases. Further information is provided in Figs. S1 and S2, Dataset S1, and SI Experimental Procedures.

mice were present in expected ratios immediately after birth, whereas dead pups lacking a milk pouch were sometimes found later on. Therefore, some *df/+* mice die after birth, indicating that 16p11.2 loss can compromise survival.

**Gene Expression in Multiple Brain Regions Corresponds to 16p11.2 Dosage.**

To validate the models, we analyzed gene expression profiles in the brain and determined whether expression corresponded with dosage. We measured mRNA intensities in 37 microarray hybridizations representing four brain regions (olfactory bulbs, cortex, cerebellum, and brainstem; five samples were hybridized twice for estimation of technical errors) in two *df/+*, three *+/+*, and three *dp/+* male mice. All mice were F1 C57BL/6N:129Sv hybrids; therefore, other than the engineered CNV, their genomes were identical. A scatter plot of the gene expression intensity difference between *dp/+* and *+/+* vs. the difference between *df/+* and *+/+* indicated that genes within 16p11.2 displayed a large difference between *df/+* and *+/+* brain, and a much smaller difference between *dp/+* and *+/+* brain (Fig. S3A). Two-way ANOVA with brain region and dosage as main factors indicated that, of 33 genes in the engineered region, expression of 26 was affected directly by dosage (Dataset S2). *Gdgd3*, mapping within the engineered region, showed extreme up- and down-regulation; this reflected differences in *Gdgd3* expression in C57BL/6N vs. 129Sv strains. Further analysis indicated that expression of genes in the region was significantly altered in each of the four brain regions analyzed, and that expression was affected more by deletion than by duplication (Fig. S3B). These findings indicate that copy number dictates gene expression levels in multiple brain regions, and that loss has the largest effect.

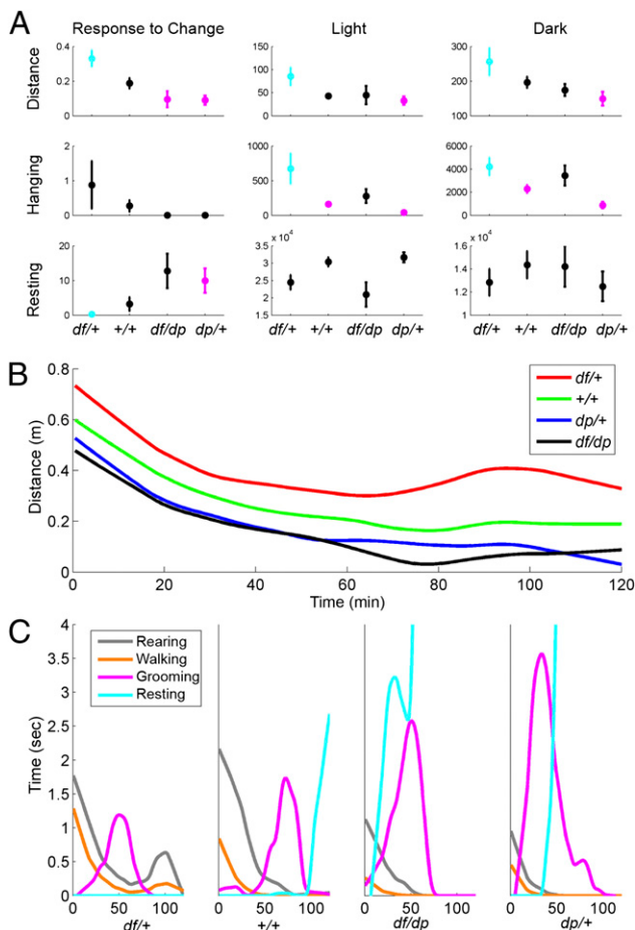
**df/+ and dp/+ Mice Have Distinctive Behavioral Phenotypes. General survey of behavior.**

The clinical evidence that patients with 16p11.2 CNVs have highly heterogeneous symptoms suggested that if the corresponding genomic alteration did in fact cause behavioral alterations in mice, the phenotypes might also be highly variable. Therefore, we believed it imperative to monitor the 16p11.2 CNV models for multiple behaviors by using as quantifiable and unbiased approaches as possible. We used HomeCageScan, a system previously used to assess behavioral alterations caused by neurodegenerative disease, neurotoxic agents, and pain (29, 34–36). We investigated behavior of a cohort of 50 male and female

mice. The mice were progeny from *df/+*  $\times$  *dp/+* crosses, and therefore included *+/+* and *df/dp* diploid controls ( $n = 15$  and  $n = 9$ , respectively), *df/+* ( $n = 13$ ), and *dp/+* ( $n = 13$ ). The parents in these crosses came from two chimeras. Thirty-nine of these were later used for MRI (as detailed later). Recording was done in cages that were significantly larger, with a ceiling that was much higher, than a standard mouse cage. The reason for using large cages is that, by minimizing the physical constraints on the animals, a rich spectrum of behaviors evolves (37) and the dynamics of the change in behavior varies between genotypes (38). In this experimental paradigm, the recording cages posed a new environment to the mice being analyzed. In particular, mice had to adapt their climbing abilities to this new environment. In each session, the behavior of four individual mice was recorded simultaneously. Multiple sessions were performed so that the behavior of each of the 50 mice was analyzed. Mice were transferred into the recording cages before the last 2 h of the light period. The recording started immediately and continued for 60 h after the onset of the first dark period (i.e., over three 12-h dark periods and two 12-h light periods). We also tested social behavior and grip strength (Fig. S4). These analyses did not show significant differences and therefore we do not discuss those data.

However, six of eight distinct behavioral measures revealed significant genotype differences. The changes were evident immediately after the mice were introduced into the new cages, as well as throughout the entire period of the test (Fig. 2A and Dataset S3). Five of these six differences were particularly interesting, as these behaviors were affected in opposite directions in *df/+* and *dp/+* mice relative to diploid (*+/+* and *df/dp*) controls. As was the case for gene expression profiles, the effect of the deletion on behavior was more severe than that of the duplication. As 16p11.2 CNV-associated syndromes sometimes have a gender bias (16), we asked whether the changes in behavior were sex-specific. Two-way ANOVA did not reveal significant interactions between 16p11.2 dosage and sex for any of the behavioral measures (Dataset S3). Later we describe in detail the behaviors affected by 16p11.2 dosage.

**Response to change in environment.** The 16p11.2 CNV models responded uniquely to environmental change: within the 2-h period after being transferred to the test cage, the distance traveled, as well as the time spent walking, lingering, and resting, depended on genotype ( $P_{\text{distance}} = 0.0016$ ,  $P_{\text{walking}} = 0.003$ ,  $P_{\text{lingering}} = 0.021$ ,  $P_{\text{resting}} = 0.025$ ; Dataset S3). Tukey's confi-

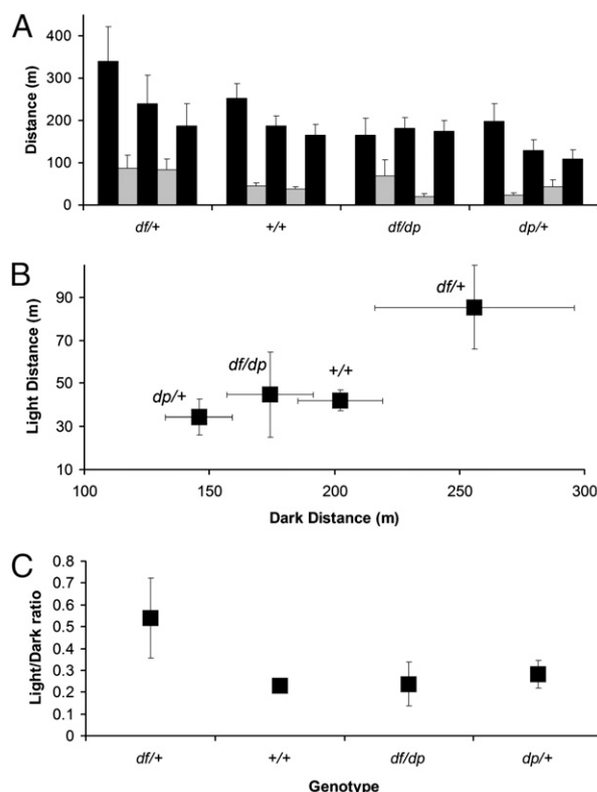


**Fig. 2.** Multiple behaviors depend on 16p11.2 dosage. (A) The relationship between 16p11.2 dosage and distance traveled (Top), hanging (Middle), and resting (Bottom) is shown during the 2-h period immediately after cage transfer (Left), the light periods (Middle), and the dark periods (Right). Averages and standard error of the mean (SEM) are displayed. Statistically significant pair-wise differences relative to the *df/+* group (determined by ANOVA followed by Tukey's confidence intervals) are depicted by the following: *df/+* is cyan when it differs significantly from at least one other cohort, but is otherwise black; cohorts that differ significantly from *df/+* are magenta, but are otherwise black. Dataset S3 shows pair-wise comparisons between genotypes. (B) Median distance traveled vs. time during the first 2 h after cage transfer. (C) Detailed behaviors during the first 2 h after cage transfer. Medians of the cumulative time of four distinct behaviors (rearing, walking, grooming, resting) vs. time elapsed from the beginning of the trial are shown for each genotype. During this period, *df/+* mice do not rest; instead, they have a second peak of activity (B and C). Further information is provided in Dataset S3 and SI Experimental Procedures.

dence intervals were calculated to determine pair-wise differences between genotypes while controlling for false discovery rate (FDR) (39). Distance traveled and time spent walking were significantly higher in *df/+* relative to other groups, and the time spent lingering was significantly higher in *df/+* relative to *dp/+* mice. Resting was lower in *df/+* and higher in *dp/+* mice relative to controls. We further analyzed the time course of these changes during the first 2 h of the test (Fig. 2 B and C). Mice of all four genotypes were most active immediately after being transferred to the test cage, with activity gradually declining during the first hour as the mice habituated to their new environment (Fig. 2B). However, the *df/+* cohort had a second burst of activity that did not occur in other groups. The response to being transferred to the test cage occurred in three sequential stages defined by the actions of the mice (Fig. 2C). For each group, the first stage was characterized by elevated walking and

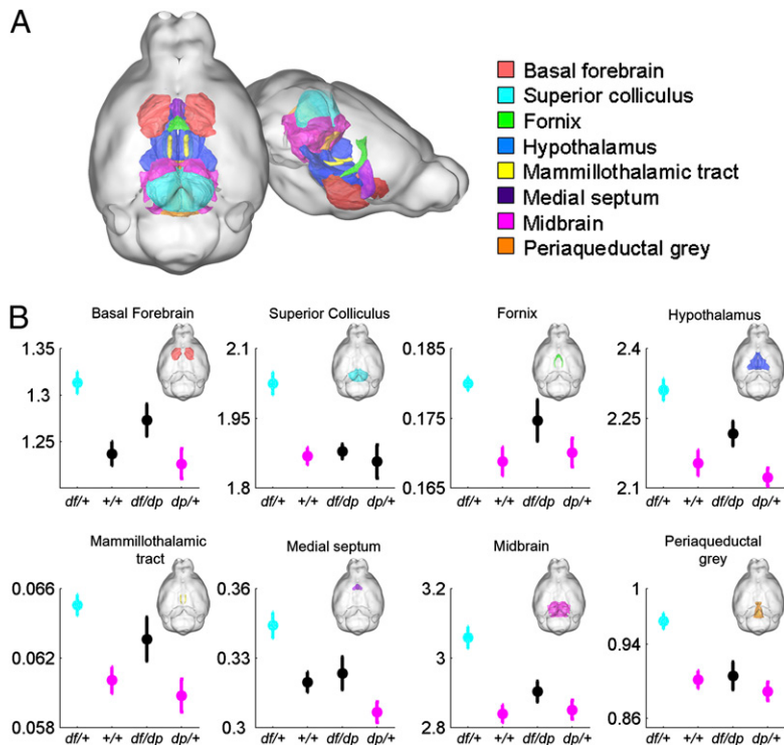
rearing, and the second stage was characterized by elevated grooming. The third stage was characterized by resting, which was significantly decreased and increased in *df/+* and *dp/+*, respectively, relative to *+/+*; indeed, *df/+* mice had a burst of walking and rearing, whereas resting was absent during this stage. Thus, the rate of certain behaviors is affected reciprocally by loss and gain of 16p11.2 dosage in response to environmental challenge. In addition, the sequence of these behaviors is disrupted in *df/+* mice. Thus, 16p11.2 CNVs affect both the rate and the timing of specific behaviors.

**Diurnal deficits.** Sleeping disorders are frequently reported in many psychiatric disorders, including autism (40). Because we recorded behavior over sequential dark/light intervals, we could assess the effect of 16p11.2 CNVs on light and dark cycling by using previously established methods (29). Mice are nocturnal, and indeed, each genotype was most active during the dark periods (Fig. 3A). The *+/+*, *df/+*, and *dp/+* mice were most active during the initial dark period, with activity decreasing in successive dark periods. Although activity of *df/+* mice was highest in the first dark period, during subsequent dark cycles, the mice adapted and had activity levels similar to controls. The activity of *df/dp* and *+/+* mice was indistinguishable in light and dark periods; *dp/+* mice were notably less active in the dark (but not light) period (Fig. 3B). In striking contrast, *df/+* mice were significantly more active than mice of other genotypes in both light and dark periods. Furthermore, *df/+* mice were unique, as they had a higher ratio of light to dark activity compared with the other



**Fig. 3.** Behavior/diurnal rhythms are affected in 16p11.2 CNV mice. Comparison of the distance traveled in cohorts of distinct genotypes (detailed in the text) during 60 h of alternating 12-h dark and 12-h light cycles (i.e., spanning three dark periods plus two light periods). (A) The distance traveled over successive dark (black bars) and light (gray bars) periods is shown in bins of 12 h. Five successive dark or light periods are shown. (B) The average distance traveled during light vs. the average distance traveled during dark. (C) The ratio of light-to-dark activity indicates that *df/+* mice are unusual in that their activity is not as restricted to the dark periods as the controls are. Average and SEM are presented in all panels.





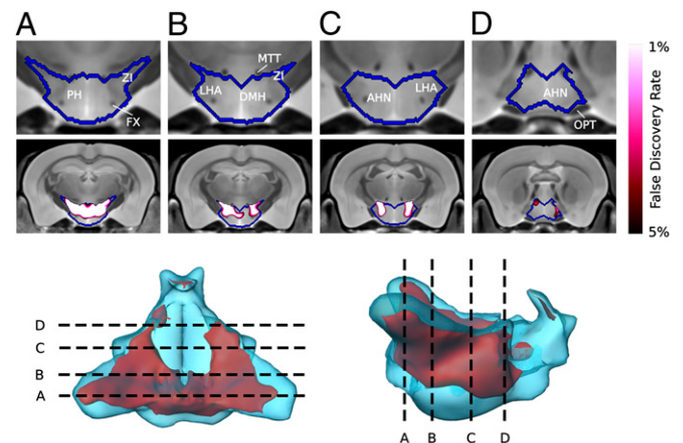
**Fig. 4.** MRI identifies structural changes in brains of 16p11.2 CNV mice. The relative volume (percentage of total brain volume) of eight brain regions is increased in *dtf/+* mice. (A) Three-dimensional representation of the mouse brain highlights eight regions (colored as in legend) affected by 16p11.2 dosage. (B) Relative volumes (shown as percentage of total brain volume) are dependent on dosage. Mean and SEM are shown. Statistically significant pairwise differences to the *dtf/+* group (determined by *t* test followed by Bonferroni–Holm procedure) are depicted as follows: cyan indicates that *dtf/+* differs from at least one other cohort, magenta indicates cohorts that differ significantly from *dtf/+*, and black indicates comparisons that do not differ significantly from *dtf/+*. Full pairwise comparisons are shown in Dataset S4.

extremely rare; in addition, patients with 16p11.2 deletions tend to be diagnosed earlier than those with duplications (16).

**16p11.2 CNVs Affect Many Brain Regions.** Changes in head circumference and abnormal brain structure have been reported in patients with 16p11.2 CNVs (14, 16). By using MRI, we find significant volumetric changes in eight different brain regions. Brains of *dtf/+* (but not *dp/+*) mice, have significant volumetric changes relative to controls, but the most extensive difference is between *dtf/+* and *dp/+* mice, emphasizing the opposing effects that 16p11.2 dosage has on brain architecture. Importantly, brains of *dtf/dp* diploid controls are not significantly different from *+/+* controls, providing genetic evidence that the structural changes in *dtf/+* and *dp/+* models are dosage-dependent.

**16p11.2 CNVs Affect Multiple Behaviors.** Several human studies compared the behavioral symptoms of patients with 16p11.2 deletions and duplications (10, 13, 16); however, to our knowledge, there is no evidence that loss and gain of 16p11.2 affect behavior in opposing ways. Even with patients harboring the same 16p11.2 lesion, there is a broad spectrum of clinical symptoms, some patients being severely affected and others highly functional. By simultaneously analyzing multiple behaviors in the context of a new environment, we identify a number of behaviors that are altered in 16p11.2 CNV mice, revealing that deletion and duplication have opposite consequences. These highly significant changes survive strict statistical analyses (37, 43). Each genotype responds to the new cage with heightened activity, but only *dtf/+* mice have a second burst of activity at a time when controls are already resting. When control mice have become accustomed to their new environment, they have a gradual increase in freedom of movement on the ceiling over the course of the trial, i.e., a mobility gradient that recapitulates the ontogeny of movement (44). In contrast, *dtf/+* mice do not show the mobility gradient: their ceiling-climbing behavior is restricted to specific locations and their movements are stereotypic. Interestingly, this ceiling-climbing behavior is similar to the behavior trap described in rats with lateral hypothalamic lesions and 6-hydroxydopamine-induced lesions (30, 31), a well characterized model of Parkinson

disease. Other phenotypes of these rats are feeding problems (45, 46), sensory neglect, and abnormal gait (30, 31, 47–49). Indeed, abnormal gait and motor delay (13, 16, 18, 50), attention deficits (13), and feeding defects (16) are common in patients with 16p11.2 deletion. Moreover, motor development problems are common in autism spectrum disorders and may serve as an indicator for early intervention, as these features appear before the core symptoms that define autism (51).



**Fig. 5.** Details of alterations in the hypothalamus detected by MRI. Three-dimensional models of the surface of the hypothalamus (Bottom), coronal images showing the regions affected (Middle), and magnification focusing on the hypothalamus (Top). Red indicates voxels that differ significantly between *dtf/+* and *dp/+* cohorts with an FDR of 0.05. The sections performed along four locations marked A–D (A, most posterior; D, most anterior). Colors indicate voxels that differ significantly between *dtf/+* and *dp/+* cohorts, with brightness indicating the significance of the difference, as specified by the FDR. AHN, anterior hypothalamic nucleus; DMH, dorsomedial hypothalamus; FX, columns of the fornix; LHA, lateral hypothalamic area; MTT, mammillothalamic tract; OPT, optic tract; PH, posterior hypothalamic nucleus; ZI, zona incerta.

**Deletion of 16p11.2 Causes Lethality in Neonates.** A major finding of this work is that approximately half of *df/+* neonates die after birth, a finding that may have relevance to autism incidence. The precise cause of death in *df/+* mice could be related to feeding deficits, but this remains to be investigated. Based on our findings, we suggest that efforts be made to determine whether 16p11.2 deletion is associated with unexplained cases of infant death. If these findings generalize to other genotypes associated with autism, they may explain puzzling aspects of the human condition. The recent increase in autism incidence (52) might be partially attributable to factors that improve pre- and/or postnatal survival. Human studies are consistent with this idea, as it is much more common for inherited rare copy number polymorphisms that affect coding regions to be duplications than deletions (53).

**Closing.** This work demonstrates the value of using mice to model CNVs found in human disorders. This approach provides functional evidence that 16p11.2 CNVs affect brain anatomy and behavior in mice, with loss and gain having opposing effects. Multiple brain regions are affected, with deletion of 16p11.2 causing profound behavioral changes such as hyperactivity, dif-

ficulty adapting to change, sleeping abnormalities, and repetitive or restricted behaviors. In addition, our findings suggest a potential link between 16p11.2 copy number alterations and infant mortality. Finally, we note a similarity in phenotype between 16p11.2 deletions and rats with lateral hypothalamic lesions. These 16p11.2 CNV models should prove valuable for elucidating the physiological basis of neurodevelopmental syndromes and for evaluating their treatments.

## Experimental Procedures

Mice carrying rearrangements corresponding to the human CNVs (Dataset S1) were established by using chromosome engineering as described previously (19, 20, 27, 32). HomeCageScan system (CleverSys) was used to analyze behavior in a cohort of 50 adult *df/+*, *+/+*, *df/dp*, and *dp/+* mice. Thirty-nine of these mice were also analyzed by MRI. Hypothesis testing was followed by correction for multiplicity (SI Experimental Procedures provides additional details).

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