Neurobiology of Disease

The Disruption of *Celf6*, a Gene Identified by Translational Profiling of Serotonergic Neurons, Results in Autism-Related Behaviors

Joseph D. Dougherty,^{1,2} Susan E. Maloney,^{1,2} David F. Wozniak,² Michael A. Rieger,^{1,2} Lisa Sonnenblick,^{3,4,5} Giovanni Coppola,⁴ Nathaniel G. Mahieu,¹ Juliet Zhang,⁶ Jinlu Cai,⁸ Gary J. Patti,¹ Brett S. Abrahams,⁸ Daniel H. Geschwind,^{3,4,5} and Nathaniel Heintz^{6,7}

Departments of ¹Genetics and ²Psychiatry, Washington University School of Medicine, St. Louis, Missouri 63110, ³UCLA Center for Autism Research and Treatment, Semel Institute for Neuroscience and Behavior, ⁴Program in Neurogenetics, Department of Neurology, and ⁵Department of Human Genetics, David Geffen School of Medicine at UCLA, Los Angeles, California 90095, ⁶Laboratory of Molecular Biology, Howard Hughes Medical Institute, and ⁷The GENSAT Project, Rockefeller University, New York, New York 10065, and ⁸Departments of Genetics and Neuroscience, Albert Einstein College of Medicine, New York, New York 10461

The immense molecular diversity of neurons challenges our ability to understand the genetic and cellular etiology of neuropsychiatric disorders. Leveraging knowledge from neurobiology may help parse the genetic complexity: identifying genes important for a circuit that mediates a particular symptom of a disease may help identify polymorphisms that contribute to risk for the disease as a whole. The serotonergic system has long been suspected in disorders that have symptoms of repetitive behaviors and resistance to change, including autism. We generated a bacTRAP mouse line to permit translational profiling of serotonergic neurons. From this, we identified several thousand serotonergic-cell expressed transcripts, of which 174 were highly enriched, including all known markers of these cells. Analysis of common variants near the corresponding genes in the AGRE collection implicated the RNA binding protein CELF6 in autism risk. Screening for rare variants in *CELF6* identified an inherited premature stop codon in one of the probands. Subsequent disruption of *Celf6* in mice resulted in animals exhibiting resistance to change and decreased ultrasonic vocalization as well as abnormal levels of serotonin in the brain. This work provides a reproducible and accurate method to profile serotonergic neurons under a variety of conditions and suggests a novel paradigm for gaining information on the etiology of psychiatric disorders.

Introduction

The CNS has remarkable cellular diversity with hundreds of distinct cell types based on morphology alone (Ramon y Cajal et al., 1899). Although nearly every one of these cell types has an identical genome, each cell only uses a subset of genes as required by

Correspondence should be addressed to Dr. Joseph D. Dougherty, Department of Genetics, Campus Box 8232, 4566 Scott Avenue, St. Louis, MO 63110-1093. E-mail: jdougherty@genetics.wustl.edu.

DOI:10.1523/JNEUROSCI.4762-12.2013

Copyright © 2013 the authors 0270-6474/13/332732-22\$15.00/0

its particular functional role. Understanding this molecular genetic diversity of cell types in the CNS may provide important insight, both for the particular roles of a given cell type as well as for the potential consequences of genetic polymorphisms to neural circuits implicated in human disease.

We have developed techniques that permit the global assessment of translation in genetically defined cell types *in vivo* (Doyle et al., 2008; Heiman et al., 2008). Here, we apply this methodology for the first time to the serotonergic system. The serotonergic system is thought to have important roles in regulation of basic physiological processes, such as breathing, thermoregulation, and sleep, as well as higher cognitive phenomena, such as mood and learning. Most importantly, its dysfunction is suspected in several neuropsychiatric diseases, including obsessive compulsive disorder and autism, among others (Veenstra-VanderWeele et al., 2000; Canli and Lesch, 2007; Deneris and Wyler, 2012).

Autism is a pervasive developmental disorder characterized by core symptoms, including impairment in social interactions and communication, as well as repetitive behaviors, restricted interests, and resistance to change (American Psychiatric Association, 2000; Fombonne, 2005). With a concordance rate reported from 60–90% for monozygotic twins, autism clearly has a remarkably

Received Oct. 4, 2012; revised Oct. 30, 2012; accepted Nov. 15, 2012.

Author contributions: J.D.D., D.F.W., G.C., G.J.P., B.S.A., D.H.G., and N.H. designed research; J.D.D., S.E.M., M.A.R., L.S., G.C., N.G.M., and J.Z. performed research; J.C. contributed unpublished reagents/analytic tools; J.D.D., S.E.M., D.F.W., M.A.R., G.C., G.J.P., and B.S.A. analyzed data; J.D.D., S.E.M., D.F.W., M.A.R., and B.S.A. wrote the paper.

This work was supported by the Howard Hughes Medical Institute (N.H.), the Simons Foundation (J.D.A.), and National Institutes of Health Grant 4R00NS067239-03 to J.D.D. and Grant P30 H0062171 to D.F.W. We thank Wenxiang Zhang, Connie Zhao, Rada Norinsky, Chingwen Yang, Sujata Bupp, Wendy Yang, Rajiv Shah, Eric Klein, Nilambari Pisat, A. Gulhan Ercan-Sencicek, and Afua Akuffo for their assistance. We thank Shiaoching Gong, Jennifer Stone, John Constantino, Tim Holy, Terra Barnes, and Don Conraf for advice. We gratefully acknowledge the resources provided by the Autism Genetic Resource Exchange (AGRE) Consortium and the participating AGRE families. AGRE is a program of Autism Speaks and National Institute of Mental Health Grant 1U24MH081810 [Clara M. Lajonchere, Principal Investigator (PI)]. We are equally grateful to all of the families at the participating Simons Simplex Collection (SSC) sites, and PIs (A. Beaudet, R. Bernier, J. Constantino, E. Cook, E. Fombonne, D. Geschwind, R. Goin-Kochel, E. Hanson, D. Grice, A. Klin, D. Ledbetter, C. Lord, C. Martin, D. Martin, R. Maxim, J. Miles, O. Ousley, K. Pelphrey, B. Peterson, J. Piggot, C. Saulnier, M. State, W. Stone, J. Sutcliffe, C. Walsh, Z. Warren, E. Wijsman).

The authors declare no competing financial interests.

strong genetic component, yet, as is the case for many psychiatric disorders, studies indicate that this genetic contribution is likely to be complex and polygenic. In autism, those genes that have been implicated thus far either explain only a small number of the cases or make relatively small contributions (Moldin and Rubenstein, 2006; Freitag, 2007; Wang et al., 2009; Weiss et al., 2009; Abrahams and Geschwind, 2010). One possible explanation for the difficulty in discovering genes contributing to complex psychiatric disorders, such as autism, would be distinct genetic causes in different groups of individuals. However, the commonality of the symptoms suggests that these distinct genes would still be impacting a common pathway or circuit in the brain. This suggests a candidate approach, focused on particular categories of genes, or genes expressed in particular cell types that are a priori suspected to contribute to a particular symptom, may increase statistical power by decreasing the number of tests. However, one risk with candidate gene studies is that choice of genes may be considered arbitrary or limited by our current knowledge of relevant genes for a specific biological process. Here, we propose that cell type specific translational profiling can be applied to provide guidance for genetic studies of the symptomatology of CNS disorders, and we test this approach in the case of serotonergic neurotransmission and autism.

To this end, we have applied the Translating Ribosome Affinity Purification (TRAP) methodology to identify the comprehensive *in vivo* suite of ribosome bound mRNA in serotonergic neurons in adult mice. Screening the TRAP-identified serotonergic genes using human data suggested polymorphisms in the *CUGBP Elav-like family member* 6 (*CELF*6) gene may contribute to autism risk. Consistent with these findings, *Celf6* mutant mice exhibit some autism-related behaviors and abnormal brain levels of serotonin.

Materials and Methods

Generation and husbandry of Slc6a4 TRAP mice

All protocols involving animals were approved by the Institutional Animal Care and Use Committee of Rockefeller University and the Animal Studies Committee of Washington University. BAC transgenic mice were generated as described previously (Gong et al., 2002), using BAC RP24-335M24. Positive founders and subsequent eGFP-L10a-positive progeny were bred to C57BL/6J and pups were genotyped with tail-DNA PCR for eGFP at each generation. Lines from two independent founders were tested for accurate expression as described below. All serotoninpositive neurons labeled with GFP in both lines. A few nonserotonergic neurons showed trace labeling with GFP antibodies in the cingulate cortex, hypothalamus, inferior colliculi, lateral nucleus of the olfactory tract in one of the two lines (JD57), and in the dorsal part of the caudal spinal trigeminal nucleus in the other line (JD60, data not shown). We used line JD60 for all subsequent experiments.

Generation and characterization of Celf6 antibodies

Peptides (QPGSDTLYNNGVSPC and AASEGRGEDRKC) from Celf6, selected for cross-species conservation, relative uniqueness across the Celf family, and hydrophobicity, were synthesized, conjugated to KLH, and injected into New Zealand white rabbits following standard protocols (Green Mountain Antibodies). ELISA was used to confirm generation of antibodies specific to each peptide. High-titer rabbits were boosted, and blood sera were affinity purified following standard protocols. Antibodies were tested for effectiveness for Western blot, and immunofluorescence, using 3T3 cells transfected with GFP-tagged and untagged isoforms of Celf6, then fixed with 4% paraformaldehyde. Antibodies against both peptides exhibited immunoreactivity by immunoblot on protein from in Celf6 overexpressing 3T3 cells. Only antibodies against QPGSDTLYNNGVSPC were effective for immunofluorescence on fixed cells. Specificity was confirmed in Celf6 knock-out (Celf6 $^{-/-}$) mouse tissue.

Immunofluorescence and microscopy

Adult mice were killed and perfused transcardially with 15 ml PBS, then 25 ml 4% paraformaldehyde in PBS. Brains were extracted and then cryoprotected in 30% sucrose PBS overnight, frozen in Neg 50 mounting media (Richard Allen Scientific), and sectioned on a cryostat to 40 μ m of thickness. Serial sections were collected in PBS with 0.1% sodium azide and stored at 4°C, protected from light, until use.

EGFP-L10a expression was examined both with and without chicken anti-GFP antibody (Abcam ab13970), and colocalization was confirmed with rabbit anti-serotonin antibody (Immunostar), mouse anti-Neun (Millipore Bioscience Research Reagents), detected with appropriate Alexa dye-conjugated secondary antibodies (Invitrogen). Images were captures on a LSM 510 Zeiss confocal microscope.

Immunohistochemistry

Mice were processed as above. For anti-GFP immunohistochemistry, brains were processed by Neuroscience Associates as described previously (Doyle et al., 2008), using custom goat anti-GFP antibodies. For Celf6 immunohistochemistry, brains were processed as above, incubated overnight with 1:1000 with purified rabbit anti-Celf6 antibody, then donkey anti-rabbit biotinylated secondary antibodies (Jackson ImmunoResearch Laboratories), and developed with the Vectashield Elite ABC kit (Vector Laboratories). Sections were digitized with a Zeiss Axioskop2 and customized macros.

Translating ribosome affinity purification

Three replicate pools of five adult mice of both sexes were killed, and brains were rapidly dissected in ice-cold dissection buffer, containing cycloheximide, to isolate the midbrain and brainstem. TRAP was conducted as described previously (Heiman et al., 2008). Briefly, each pool was homogenized for 12 strokes in a glass Teflon homogenizer on ice, in buffer (10 mM HEPES [pH 7.4], 150 mM KCl, 5 mM MgCl₂, 0.5 mM dithiothreitol, 100 µg/ml cycloheximide, protease inhibitors, and recombinant RNase inhibitors). Nuclei and debris were removed with centrifugation at 2000 \times g for 10 min at 4°C. DHPC (Avanti) and NP-40 (Ipgal-ca630, Sigma) were added to supernatant to final concentrations of 1% and 30 mM, respectively. After 5 min incubation of ice, supernatant was centrifuged for 15 min at 20,000 \times g, and pellet was discarded. Supernatant was mixed with protein G-coated magnetic beads (Invitrogen), previously conjugated with a mix of two monoclonal anti-GFP antibodies (Doyle et al., 2008), and incubated with rotation for 30 min at 4°C. Beads were washed three times with high salt wash buffer (10 mM HEPES [pH 7.4], 350 mм KCl, 5 mм MgCl₂, 1% NP-40, 0.5 mм dithiothreitol, and 100 µg/ml cycloheximide), and RNA was purified from ribosomes using Trizol (Invitrogen), following the manufacturer's protocols, followed by DNase treatment, further purification, and concentration with RNeasy min-elute columns, following the manufacturer's protocol (Qiagen). RNA was also harvested in parallel from each unbound fraction of affinity purification as a measure of total tissue RNA. RNA concentration of all samples was measured with Nanodrop spectrophotometer, and integrity was confirmed with PicoChips on the Agilent BioAnalyzer (RIN > 8).

Microarray experiments and statistical analysis

A total of 20 ng of each RNA sample was amplified with the Affymetrix Two-Cycle amplification kit following the manufacturer's instructions, and quality of labeled aRNA was assessed with Bioanalyzer. Labeled RNA from immunoprecipitated ribosomes and total tissue RNA was hybridized to separate Affymetrix Mouse Genome 430 2.0 arrays and scanned following the manufacturer's protocols. Data are available from the Gene Expression Omnibus (GSE36068).

Data were analyzed using the Bioconductor module within the statistical package R. Data quality was assessed by examining raw images of slides, boxplots, histograms, correlation coefficients, false-positive rates, and scatterplots comparing replicate experiments. Data were normalized as described previously (Dougherty et al., 2010) but using Affymetrix chip definition files. Briefly, GCRMA was used to normalize within replicates and to biotinylated spike in probes between conditions. Fold change, Specificity Index (SI), and Specificity Index statistic (pSI) were calculated for all genes with expression above background, as described enriched in serotonergic neurons for the purposes of further analysis. This resulted in 196 probesets covering 174 distinct genes.

Heirarchical clustering was conducted using Bioconductor on any genes with pSI < 0.01 in any cell type from the current data and the initial survey (Doyle et al., 2008).

Mouse genes from above were mapped to human Unigenes genes using homologene, gene symbol, and BLAT of mouse mRNA against human genome sequence, with 147 genes having clear human homologs. Custom perl scripts were used to identify alignment coordinates of human Unigenes from UCSC genome browser, along with 10 kb of flanking sequence to cover potential regulatory regions, with 136 Unigenes having clear and unique genome coordinates.

Chromosomal distribution was analyzed by comparing number of regions expected to be found on each chromosome (Wilming et al., 2008), compared with observed number of regions.

Analysis of genome-wide association studies

The Rockefeller University Institutional Review Board and the Washington University Human Research Protection Office reviewed all human subjects related work in this manuscript.

Deidentified human genotype and phenotype data were downloaded with permission from the Autism Genetic Research Exchange (AGRE) consortium (Geschwind et al., 2001) for 3787 individuals in 943 multiplex families, genotyped for 550,000 SNPs with the Illumina Hap550 platform (Wang et al., 2009). SNP genotypes were called with BeadStudio 3.1.3.0, and genotypes and phenotypes were formatted for PLINK (Purcell et al., 2007) using Perl. Individuals with missing or poor data were excluded (MIND > 0.1, Geno > 0.1). Only those individuals with clearly diagnosed as autistic by AGRE were scored as affected. Those with ambiguous diagnosis (scored NQA or broad spectrum) were excluded. We selected all available SNPs within 10 kb of the transcription start or end site of the serotonergic neuron enriched genes, filtered to remove those with MAF < 0.05 and those that were out of Hardy-Weinburg equilibrium (p < 0.01). We further removed SNPs with an $r^2 > 0.5$ as they would violate the assumption of independence for multiple testing corrections and would not provide any further information. This resulted in a total of 555 SNPs being tested with the transmission disequilibrium test. Tests were then corrected with the Bonferroni method for multiple testing. Tests were run for all probands (1142 trios); and because prior work had suggested that autism in males and females may have a separate genetic etiology (Stone et al., 2004), we further analyzed these SNPs with males (915 trios) and females (227 trios) independently.

To determine whether segregating probands by sex was meaningful, we performed an additional permutation analysis by randomly sampling 915 probands and repeating the analysis. Excluding SNPs in *C1QTNF2* and *CELF6*, corrected *p* values <0.05 were observed in only 8 of 1000 iterations, suggesting that this sex-specific difference was not spurious.

Resequencing of candidate genes

Primers were designed for resequencing following standard methods (Montgomery et al., 2008), taking care to avoid SNPs and repetitive sequences in locations for the primers. Standard M13F and M13R sequences were incorporated into the primers. Deidentified DNAs from 384 male white probands from distinct families within the AGRE sample and 384 male white controls from the NIMH Human Genetic Initiative Controls (HGIC) set were amplified with PCR.

Amplicons were then purified and Sanger sequenced using M13F or M13R primers by Agencort Beckman Coulter Genomics. Sequences were trimmed to exclude low-quality reads and analyzed with Seqman software to identify known and novel SNPs. Any putative novel coding sequence mutations were confirmed by sequencing with the opposite primer.

Screening for stop mutation in additional cases and controls

An additional 864 probands from the Simons Simplex Collection (SSC) and 4992 controls from the HGIC set were screened as follows:

HGIC samples were first screened with Sequenom. The sequence variant was run in a 29-plex SNP Genotyping Assay using Sequenom iPLEX Gold Technology. The initial PCR, performed as per the manufacturer's instructions (Sequenom). The subsequent Shrimp Alkaline Phosphatase clean, the iPLEX reaction, and the resin clean were all performed as stated by Sequenom. Spectro chip arrays were spotted using a nanodispenser, according to the manufacturer's instructions. A Sequenom MALDI-TOF spectrophotometer was used to read the array. The SpectroAquire and MassARRAY Typer Software packages (Sequenom) were used for interpretation, and Typer analyzer Version 3.4.0.18 was used to review and analyze all data.

Any positive HGIC samples and all SSC samples were then screened with allele-specific PCR (Little, 2001). Primers that specifically amplify CELF6 only with the stop codon mutation were included in a multiplex reaction with internal control primers for β -actin for a final concentration of 500 nM each with Maxima SYBR Green quantitative RT-PCR Master Mix (Fermantas), and subjected to quantitative RT-PCR in an ABI Prism 7900, followed by a melt curve. β -Actin and CELF6 amplicons were distinguished on the basis of melting temperature (83°C and 88°C, respectively). Positive control DNA with stop codon was included on each plate of PCR.

Analysis of 1000 genomes data

VCF files corresponding to exome sequence data from 1092 individuals (1000 Genomes Project Consortium, 2010) were downloaded into WASP (Dubin et al., 2010) and custom scripts used to extract all exonic variants, annotate these with regard to likelihood of being pathogenic (Sunyaev et al., 2001; Ng and Henikoff, 2002; Eddy, 2004; Adzhubei et al., 2010), and then determine minor allele frequencies. The DGV was queried in March 2012 in the UCSC genome browser using the UCSC annotated *CELF6* transcriptional start and stop as a query.

Quantitative real-time RT-PCR

TRAP was conducted as above for an additional three male and three female slc6a4 bacTRAP mice. Each sample was used to synthesize cDNA from 50 ng of total RNA using Protoscript (*NEB*), primed with Oligo dT, following the manufacturer's instructions. Quantitative RT-PCR was conducted in a Bio-Rad IQ5 with Bio-Rad 2× Sybr Green Mastermix (Bio-Rad). Data were analyzed in the IQ5 software using the ddCT method, with β -actin as a control. Melt curves were conducted to assure specificity of product, and each product was sequenced to confirm accuracy of amplification.

Generation of Celf6 knock-out mice

Long and short arms with LoxP sites flanking exon 4 of *Celf6* were cloned by PCR adjacent to an Frt flanked neomycin-resistance cassette gene. B6(Cg)-Tyr^{c-2j}/j-derived ES cells were electroporated using standard methods, and neomycin-resistant colonies were screened by PCR and southern blot for proper integration. Positive colonies were injected into C57BL/6J mouse blastocysts, and chimeric mice were bred to germline Flpe expressing C57BL/6J mice to remove neomycin selection cassette, then actin-Cre C57BL/6J mice to create germline deletions of *Celf6*. Recombination was confirmed in progeny by PCR. Heterozygous progeny were crossed to generate knock-out (*Celf6*^{-/-}) and wild-type (WT) littermate mice for behavioral and anatomical assays.

Behavioral tests

Animals and experimental design of behavioral studies. A total of 46 C57BL/6J Celf6^{-/-} (n = 23, 14 females and 9 males) and litter-matched WT (n = 23, 14 females and 9 males) mice were used for ultrasonic pup vocalization (USV) recording and analysis. The animal colony room lighting was a 12:12 h light/dark cycle; room temperature ($\sim 20^{\circ}C-22^{\circ}C$) and relative humidity (50%) were controlled automatically. Standard laboratory chow and water were available on a continuous basis.

Twenty experimentally naive male C57BL/6J *Celf6^{-/-}* (n = 11) and litter-matched WT (n = 9) mice were used for adult behavioral testing. The sequence of behavioral testing was devised to minimize "carry-over" effects across behavioral tests as much as possible. Behavioral testing began when the mice were 3–5 months of age and included a 1 h locomotor activity/exploration test, followed 1 d later by an evaluation on a battery of sensorimotor measures, to assess possible disturbances in general activity and sensorimotor functions that might affect performance on subsequent tests. One day later, the mice were tested in the Morris

water maze, which included cued (visible platform), place (hidden platform), and probe (platform removed) trials to assess potential nonassociative deficits as well as spatial learning and memory impairments. The following week, the mice were tested for their ability to learn another platform location (reversal and additional probe trials) to evaluate cognitive flexibility in terms of extinguishing one learned response and acquiring another. During the next week, sociability was assessed using the social approach test followed by an evaluation of sensorimotor reactivity and gating by quantifying acoustic startle and prepulse inhibition (PPI). Approximately 2-3 months later, anxiety-like behaviors were measured in the mice by testing them in the elevated plus maze (EPM). The following week, the mice were retested on the social approach test for further evaluation of sociability and preference for a novel conspecific (stimulus mouse). Nineteen days later, exploratory behaviors and olfactory preference were quantified in the holeboard exploration/olfactory preference test.

Ultrasonic vocalization recording and analysis. WT and Celf6^{-/-} pups were individually removed from their parents at postnatal day 8 and placed in a dark, enclosed chamber. Ultrasonic vocalizations were obtained using an Avisoft UltraSoundGate CM16 microphone, Avisoft UltraSoundGate 416H amplifier, and Avisoft Recorder software (gain = 6 dB, 16 bits, sampling rate = 250 kHz). Pups were recorded for 3 min, after which they were weighed and returned to their home cages. Frequency sonograms were prepared from recordings in MATLAB (frequency range = 40 kHz to 120 kHz, FFT size = 256, overlap = 50%), and individual syllables were identified and counted from the sonograms according to a previously published method (Holy and Guo, 2005).

One hour locomotor activity and sensorimotor battery. Locomotor activity was evaluated in the mice using transparent (47.6 \times 25.4 \times 20.6 cm high) polystyrene enclosures and computerized photobeam instrumentation as previously described (Wozniak et al., 2004). General activity variables (total ambulations and vertical rearings) along with indices of emotionality, including time spent, distance traveled, and entries made in a 33 \times 11 cm central zone and distance traveled in a 5.5 cm contiguous peripheral zone were analyzed. All mice were also evaluated on a battery of sensorimotor tests designed to assess balance (ledge and platform), strength (inverted screen), coordination (pole and inclined screens), and initiation of movement (walking initiation), as previously described (Wozniak et al., 2004; Grady et al., 2006). For the walking initiation test, the mouse was placed on a surface in the center of a 21×21 cm square marked with tape, and the time the mouse took to leave the square was recorded. During the balance tests, the time was recorded that the mouse remained on a Plexiglas ledge (0.75 cm wide) or a small circular wooden platform (3.0 cm in diameter) elevated 30 or 47 cm, respectively. The screen tests were conducted by placing a mouse head-oriented down in the middle of a mesh grid measuring 16 squares per 10 cm, elevated 47 cm, and angled at 60° or 90°. The time was recorded that the mouse required to turn 180° and climb to the top. For the inverted screen, the mouse was placed on the screen as described above; and when it appeared to have a secure grip, the screen was inverted 180°, and the time the mouse remained on to the screen was recorded. Each test had a maximum time of 60 s, except for the pole test, which had a maximum time of 120 s. The averaged time of two trials for each test was used for the analyses.

Morris water navigation. Spatial learning and memory were evaluated in the Morris water maze using a computerized tracking system (ANYmaze, Stoelting) as previously described (Wozniak et al., 2004). Cued (visible platform, variable location) and place (submerged, hidden platform, constant location) trials were conducted, and escape path length, latency, and swimming speeds were computed. To assess nonassociative dysfunctions, the cued condition involved conducting 4 trials per day (60 s maximum per trial) for 2 consecutive days with the platform being moved to a different location for each trial using a 30 min intertrial interval, and with very few distal spatial cues being present to limit spatial learning. Performance was analyzed across four blocks of trials (2 trials/ block). Three days later, place trials were conducted, which involved acquisition training to assess spatial learning where mice were required to learn the single location of a submerged platform in the presence of several salient distal spatial cues. During the 5 consecutive days of place trials, the mice received 2 blocks of 2 consecutive trials [60 s maximum for a trial; 30-s intertrial interval (spent on platform)] with each block being separated by ${\sim}2$ h and each mouse being released from a different quadrant for each trial. The place trials data were analyzed over five blocks of trials (4 trials/block) where each block represented the performance level for each of five consecutive days. A probe trial (60 s maximum) was administered \sim 1 h after the last place trial on the fifth day of training with the platform being removed and the mouse being released from the quadrant opposite to where the platform had been located. Time spent in the various pool quadrants, including the target quadrant where the platform had been placed and crossings over the exact platform location, served as the dependent variables. Three days after completing the place and probe conditions, mice were tested on reversal trials, which involved the same procedures as the place trials, except that the hidden platform was moved to a new location in the pool for all trials. Another probe trial was conducted after completing the reversal trials.

Social approach. Our protocol was adapted from methods previously described (Moy et al., 2004; Silverman et al., 2011) and involves quantifying sociability [tendency to initiate social contact with a novel conspecific (stimulus mouse) or cagemate] and preference for social novelty (tendency to initiate social contact with a novel vs a familiar stimulus mouse). The apparatus was a rectangular 3-chambered Plexiglas box (each chamber measuring 19.5 cm \times 39 cm \times 22 cm) containing Plexiglas dividing walls with rectangular openings (5 \times 8 cm) covered by sliding Plexiglas doors. A small stainless-steel withholding cage (10 cm h × 10 cm diameter; Galaxy Pencil/Utility Cup, Spectrum Diversified Designs) was used to sequester a stimulus mouse. The withholding cage consisted of vertical bars, which allowed for minimal contact between mice but prevented fighting, and one was located in each outer chamber. A digital video camera connected to a PC loaded with a tracking software program (ANY-maze, Stoelting) recorded the movement of the mouse within the apparatus and quantified time spent in each chamber and investigation zone surrounding the withholding cages. The investigation zones were 12 cm in diameter, encompassing 2 cm around the withholding cages. An entry into the chambers was defined when a chamber contained 80% of the mouse's body while only the head was tracked in the zones surrounding the withholding cages to capture investigative social behaviors. Indirect lighting illuminated the test room, and the entire apparatus was cleaned with Nolvasan solution while the withholding cages were cleaned with 75% ethanol solution between tests.

The first test sequence consisted of 3 consecutive 10 min trials. For the first trial, each mouse was placed in the middle chamber with the doors to the outer chambers shut to become acclimated to the apparatus. During the second trial, the mouse was allowed to freely investigate and habituate to all three chambers, including the empty withholding cages (Page et al., 2009; Naert et al., 2011; Pobbe et al., 2012). Neither group showed prior preference for a particular withholding cage during habituation. The third (test) trial assessed sociability exhibited toward a novel stimulus mouse versus a familiar, empty withholding cage by placing an unfamiliar, gender-matched stimulus mouse in one withholding cage while the other was left empty, and the test mouse was allowed to freely explore the apparatus and investigate the novel mouse in the withholding cage. A second test sequence conducted \sim 2 months later also consisted of 3 consecutive 10 min trials. For trial 1, each mouse was allowed to freely investigate and habituate to all three chambers, including the empty withholding cages. For the second trial, a gender-matched cagemate was placed in one withholding cage while the other remained empty, and the test mouse could freely explore the apparatus and investigate the cagemate in the withholding cage. During the third (test) trial, the familiar cagemate remained in the withholding cage, although a new, unfamiliar gender-matched stimulus mouse was now placed in the other withholding cage, and the test mouse was allowed to explore the apparatus and investigate the two mice contained in the withholding cages. The locations of the stimuli mice in the outer chambers for the first test sequence were counterbalanced within and across groups, and placement of the stimuli mice for the second test sequence was randomized. All stimuli mice were acclimated to the metal withholding cages for 10 min before beginning of testing.

Acoustic startle/prepulse inhibition. Sensorimotor reactivity and gating were also evaluated in the mice by quantifying their acoustic startle response (ASR) and PPI because PPI has been reported to be abnormal in individuals with autism (Perry et al., 2007; Yuhas et al., 2011). The ASR and PPI were tested using computerized instrumentation designed specifically for mice (StartleMonitor, Kinder Scientific). The ASR to a 120 dBA auditory stimulus pulse (40 ms broadband burst) and PPI (response to a prepulse plus the startle pulse) were measured concurrently using our previously described methods (Hartman et al., 2001; Gallitano-Mendel et al., 2008). Beginning at stimulus onset, 65, 1 ms force readings were averaged to obtain an animal's startle amplitude for a given trial. A total of 20 startle trials were presented over a 20 min test period, during which the first 5 min served as an acclimation period when no stimuli above the 65 dB white noise background were presented. The session began and ended by presenting 5 consecutive startle (120 db pulse alone) trials unaccompanied by other trial types. The middle 10 startle trials were interspersed with PPI trials, consisting of an additional 30 presentations of 120 dB startle stimuli preceded by prepulse stimuli of 4, 12, or 20 dB above background (10 trials for each PPI trial type). A percent PPI score for each trial was calculated using the following equation: %PPI = $100 \times (ASR_{startle \ pulse \ alone} - ASR_{prepulse \ + \ startle \ pulse})/ASR_{startle \ pulse \ alone}$ (see Hartman et al., 2001 for more details).

EPM. Anxiety-like behaviors were evaluated in the EPM according to our previously described procedures (Schaefer et al., 2000; Boyle et al., 2006). Our apparatus consisted of two opposing open arms and two opposing enclosed arms $(36 \times 6.1 \times 15 \text{ cm})$ that extended from a central platform $(5.5 \times 5.5 \text{ cm})$, which were constructed of black Plexiglas. The maze was equipped with pairs of photocells configured in a 16 (x-axis) \times 16 (y-axis) matrix, the output of which was recorded by a computer and interface assembly (Kinder Scientific). A system software program (MotorMonitor, Kinder Scientific) enabled the beam-break data to be recorded and analyzed to quantify time spent, distance traveled, and entries made into the open and closed arms and center area. To adjust for differences in general activity, the percentage of distance traveled, time spent, and entries made into the open arms out of the totals (open arms + closed arms) for each variable were also computed. Test sessions were conducted in a dimly lit room where light was provided by two 13 W black-light bulbs (Feit EcoBulbs) and each session began by placing a mouse in the center of the maze and allowing it to freely explore the apparatus. Each test session lasted 5 min, and the mice were tested over 3 consecutive days.

Holeboard exploration/olfactory preference test. Mice were evaluated for possible differences in exploratory behaviors and olfactory preferences using a slightly modified version of our previously published methods (Ghoshal et al., 2012) where hole poking served as the main behavioral response. Our modifications were based on procedures described by Moy et al. (2008) for Study 2 within their publication to assess shifts in olfactory preference in an autism mouse model. We used the same odorant, familiarization technique, and testing procedure, although our apparatus is slightly different from the one used by these authors. Our protocol involved the use of a computerized holeboard apparatus (41 imes 41×38.5 cm high clear plastic chamber), containing 4 corner and 4 side holes in the floor, with a side hole being equidistant between the corner holes (Learning Holeboard; MotorMonitor, Kinder Scientific). Pairs of photocells were contained within each hole (27 mm in diameter) and were used to quantify the frequency and duration of pokes, whereby a poke that was at least 35 mm in depth was required to be registered as a hole-poke. Each mouse received one 30 min habituation session in the holeboard during which no holes contained any odorants. The following day (test session 1), a 20 min session was conducted during which holepoke frequency and duration data were collected in the presence of empty and odorant-containing holes. All side holes were empty, while odorants were placed at the bottom of each of three corner holes, although access to the odorants was blocked, and one corner hole remained empty. The configuration of odorant-containing and empty corner holes (see Fig. 7C) was such that one hole contained a familiar odorant (fresh corn-cob bedding used in the home cages), which was placed in a corner hole opposite to a corner hole containing an unfamiliar odorant (woodchip bedding). Another novel odorant (chocolate chips) was placed in a corner hole opposite to the empty corner hole. This pattern was counterbalanced within and across groups. Immediately after completing test session 1, the mice were familiarized with the chocolate chips by placing 4-7 chocolate chips in each home cage. The same thing was done on the following day, and all but 1 chocolate chip were consumed before the start of the second test session. Test session 2 was conducted two days after test session 1 using the same procedures and pattern of odorant-containing and empty corner holes, except that the actual configuration used for test session 2 was opposite to the one used during test session 1 (see Fig. 8C). For example, the corner holes that contained bedding for test session 1 contained chocolate chips or were empty during test session 2, and vice versa. General exploratory behavior was evaluated by quantifying total hole-pokes as well as pokes into the side and corner holes for test sessions 1 and 2. Olfactory preferences were assessed by analyzing poke frequencies involving odorant-containing and empty corner holes within and between groups. Poke durations exhibited for the different types of holes were also analyzed to provide additional data on possible differences in the processing of olfactory stimuli.

Statistical analyses for behavioral data. ANOVA models were used to analyze the behavioral data. A two-way ANOVA model was used to analyze differences within and between genotype and sex for USV. Repeated measures (rm) ANOVA models containing one between-subjects variable (genotype) and one within-subjects (repeated measures) variable (e.g., blocks of trials) were typically used to analyze the learning and memory data. The Huynh-Feldt adjustment of α levels was used for all within-subjects effects containing more than two levels to protect against violations of sphericity/compound symmetry assumptions underlying rmANOVA models. Typically, one-way ANOVA models were used to analyze differences between groups for 1 h locomotor activity test and measures in the sensorimotor battery. Planned comparisons were conducted within ANOVA models for certain holeboard variables. In most other instances, pairwise comparisons were conducted following relevant, significant overall ANOVA effects, which were subjected to Bonferroni correction when appropriate.

Extraction for measurement of neurotransmitter levels Ten *Celf6*⁻⁷ and seven WT whole adult mouse brains frozen at -80° C were dried on a lyophilizer for 72 h. The dried brains were crushed to a powder and stored at -80° C. A total of 10 mg of each was extracted as follows: vortexed 30 s; centrifuged 13,000 RPM, 4°C, 10 min; sonicated at 50°C in a bath sonicator; stored at -4°C; liquid nitrogen bath 1 min and then allowed to thaw. Acetone (600 µl at 4°C) was added to each tube. Tubes were vortexed, placed in liquid nitrogen and sonicated for 10 min; this was repeated three times. Samples were incubated for 30 min and then centrifuged. Supernatant was poured into new microcentrifuge tubes and stored. Methanol, water, and formic acid (400 μ l at 86.5: 12.5:1) were added to the remaining pellet. Tubes were vortexed, sonicated for 15 min, and incubated for 30 min. Samples were centrifuged, and the supernatant was combined with the previous 600 μ l. The pooled supernatant was dried with a speedvac with no heat for 23 h. To each tube, 100 μ l of water was added. Tubes were sonicated for 5 min and incubated for 1 h. Samples were centrifuged and the resulting supernatant was transferred to deactivated glass autosampler vials for MS analysis.

LC/MS analysis

Targeted LC/MS analyses were done using an HPLC system (Agilent 1260 Infinity) coupled to an Agilent 6460 Triple Quadrapole MS operated in positive ESI mode. Analytes were separated using a Zorbax 300 SB-C18 (5 μ m, 150 \times 0.5 mm) column. An isocratic elution was used with 0.1% v/v formic acid in water. The injection volume was 0.8 μ l. A 30 min wash was performed between injections. Flow was set to 5 μ l/min. Source conditions were gas temperature 300°C, gas flow 6 L/min, nebulizer 15 psi, capillary 4000 V. Acquisition settings were dwell time 200 ms, MS1 resolution unit, MS2 resolution unit, polarity positive, cell accelerator voltage 7. The 6460 Triple Quadrapole was operated in MRM mode monitoring two transitions for each analyte with the most intense transition being used for analysis. MRM transitions were integrated with Agilent Mass Hunter Qualitative Analysis B.04.00. Statistical analysis was conducted in MS Excel using 2-tailed t tests, with Bonferroni correction.

Results

Slc6a4 bacTRAP mice target eGFP-L10a specifically to serotonergic neurons

Classical studies using a variety of methodologies (Hillarp et al., 1966; Steinbusch, 1981, 1984; Ishimura et al., 1988), including *in situ* hybridization studies of the presynaptic serotonin transporter *Slc6a4* (Bengel et al., 1997), have defined nine populations of serotonergic neurons, predominantly within the raphe nuclei. To enable translational profiling of these cells, we generated transgenic mouse lines using a *Slc6a4* BAC to drive the expression of the TRAP construct (eGFP fused to the ribosomal protein L10a). Immunohistochemistry with anti-GFP antibodies demonstrated robust expression in a pattern consistent with expression in all serotonergic neurons (Fig. 1*A*, *B*). Neurotransmitter phenotype was confirmed using confocal immunofluorescence with antibodies to GFP and serotonin (Fig. 1*C*). In all cells, GFP signal was seen only in the cytoplasm and nucleolus, consistent with ribosomal incorporation of the fusion protein.

Serotonergic neurons have a distinct translational profile

Using eGFP as an affinity tag, we purified the suite of ribosome bound mRNAs from serotonergic neurons (the TRAP methodology) (Doyle et al., 2008, Heiman et al., 2008) and interrogated them with microarrays. Independent biological replicates showed high reproducibility (Fig. 2A), with an average correlation >0.99. When these samples were compared with microarrays of the whole tissue (Fig. 2B), there was a clear, robust, and comprehensive enrichment of all mRNAs known to be specifically expressed in serotonergic neurons, including enzymes in the pathway for synthesizing serotonin in the CNS (Tph2, Ddc) (Goridis and Rohrer, 2002), transcription factors known to be important for specifying these neurons (Goridis and Rohrer, 2002; Pfaar et al., 2002) (Fev, Gata2, and Gata3), the serotonin autoreceptor (Htr1a; Korte et al., 1996), and the serotonin transporter itself (Slc6a4). There was also a clear depletion of mRNAs known to not be expressed by neurons, such as the glial genes Mobp, Aldh1L1, Mbp, and Plp. This suggests that we could indeed capture mRNA specifically from serotonergic neurons.

To select for all mRNAs used in serotonergic neurons, we removed those probesets with median expression <50, or a fold change below a background threshold determined by the glial genes (Dougherty et al., 2010). These normalized data represent >5000 genes clearly expressed in serotonergic neurons.

To generate a comprehensive list of those mRNAs either highly enriched or specific for serotonergic neurons, we compared this translational profile to our previous survey of cell types in the CNS (Doyle et al., 2008) using the pSI statistic (Dougherty et al., 2010). We identified 196 probesets with pSI < 0.01 (Table 1), corresponding to 174 genes that included the known serotonergic cell genes listed above. To validate this list, we compared it to the online catalog of mouse *in situ* hybridizations of the Allen Brain Atlas (Lein et al., 2007) in a blinded and unbiased manner (Dougherty et al., 2010). Compared with a random list of genes from the array, the serotonergic cell list had nearly three times as many *in situ* hybridizations with a pattern of enrichment in the raphe nuclei. Seventeen of these genes scored as specific to serotonergic neurons, compared with none from the random list (Fig. 2C).

We next compared our list of enriched and specific genes with the recently published data from flow-sorted embryonic serotonergic neurons from rostral and caudal nuclei (Wylie et al., 2010). There are numerous differences in methodology and design between the two studies, including age (embryonic vs adult), purification methodology (flow sorting vs TRAP), amplification methodology, normalization, and analytical strategy. Despite this, there was highly significant overlap between the two studies (χ^2 test, $p < 8.7 \text{E}^{-227}$), suggesting good conservation of gene expression across development for these neurons. Thus, the lines generated here provide *in vivo* access to the translational profile of the adult serotonergic system and can be used in the future to study the response of these cells to whole animal manipulations in a reproducible and accurate manner.

Using hierarchical clustering, we compared the serotonergic translational profile to the cell types from our previous survey (Doyle et al., 2008; Fig. 2D). Serotonergic neurons are clearly distinct from other cells. However, remarkably, of the surveyed cell types, serotonergic neurons are most similar to cholinergic neurons of the basal forebrain. Although these two populations come from distinct regions and have distinct transmitter phenotypes, both have similar functional roles; that is, both cell types project widely through the CNS and function as neuromodulators, influencing the physiological properties of whole circuits. This suggests that the translational profile of a cell population is more strongly influenced by its particular roles within the CNS than the region in which it is found or the particular neurotransmitter it uses.

Next, we performed a standard Gene Ontologies analysis on the 174 serotonergic genes using DAVID (Dennis et al., 2003), which indicated, unsurprisingly, a role for a significant fraction of these genes in serotonin signaling (not shown). However, many other genes were unannotated, and we suspected that these 174 genes may provide a novel list of candidate genes for neuropsychiatric disorders thought to involve the serotonergic system.

Of the 174, 147 had clear human homologues based on homologene, BLAT, and probematchDB (Wang et al., 2002). These were distributed across 136 noncontiguous regions of the genome. Chromosomal analysis revealed a remarkably nonrandom distribution of these regions (χ^2 test, $p < 3.57 \text{E}^{-7}$). In particular, 10.4% of these were found on the X chromosome, compared with the 4.5% that would be expected to be found there by chance given the size and gene density of the chromosomes. This is interesting considering the gender disparity in prevalence for autism (4:1 male to female; Fombonne, 2005) and depression (1: 1.7 male to female; Kessler et al., 1993), two of the disorders with suspected connections to the serotonergic system.

Transmission disequilibrium testing for polymorphisms in serotonergic genes in autism

We next sought to determine whether polymorphism in these 136 regions may be related to autism, using the Transmission Disequilibrium Test (Spielman and Ewens, 1996; Collins, 2007), applied to data generated by AGRE (Geschwind et al., 2001; Wang et al., 2009) from multiplex families with autism. We discovered two SNPs that met criteria for significance after multiple testing correction (Table 2): one in both males and females in the *C1QTNF2* gene and one specifically in males in the gene encoding the CELF6 RNA-binding protein.

We decided to first focus on the *CELF6* gene because another RNA-binding protein, FMRP, is disrupted in fragile X syndrome, the most common syndromic cause of autism (Verkerk et al., 1991; Chonchaiya et al., 2009). *CELF6* is a less well-studied member of an RNA binding protein family known as the CELF family (Cugbp and Etr-3 like factors) (Barreau et al., 2006). CELF proteins are known to regulate RNA in a variety of manners. In Drosophila, this family has been studied in the context of regula-

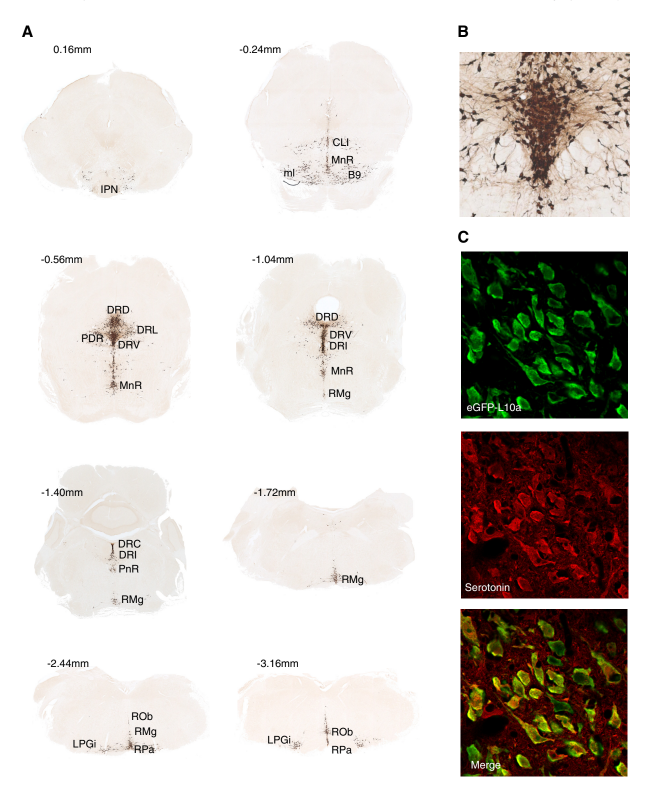


Figure 1. A bacTRAP line for serotonergic neurons. *A*, DAB immunohistochemistry with anti-GFP antibodies on Slc6a4 bacTRAP mice reveals eGFP-L10a transgene expression in the known serotonin cell-containing nuclei. *B*, Higher magnification of DRV shows neuronal morphology of eGFP-positive cells. *C*, Confocal immunofluorescence colocalization of eGFP-L10a transgene with neurotransmitter serotonin. Abbreviations following Paxinos (Franklin and Paxinos, 1997). Coordinates relative to interaural. IPN, Interpeduncular nucleus; ml, medial lemniscus; RMg, raphe magnus; R0b, raphe obscurus; RPa, raphe pallidus; LPGi, lateral paragigantocellular nucleus; DRD, dorsal raphe dorsal; DRI, dorsal raphe interfasicular; DRL, dorsal raphe lateral; DRV, dorsal raphe ventral; PDR, posterodorsal raphe; MnR, median raphe; CLi, caudal linear raphe.

tion of translation (Good et al., 2000); whereas in Xenopus, homologous proteins are implicated in mRNA localization and stability (Barreau et al., 2006). In mammals, family members are known to both positively and negatively regulate splicing of a variety of transcripts (Ladd et al., 2004), and members of the CELF family seem to be in a dynamic competition with other splicing factors to regulate cell type-specific alternative splicing (Charlet et al., 2002).

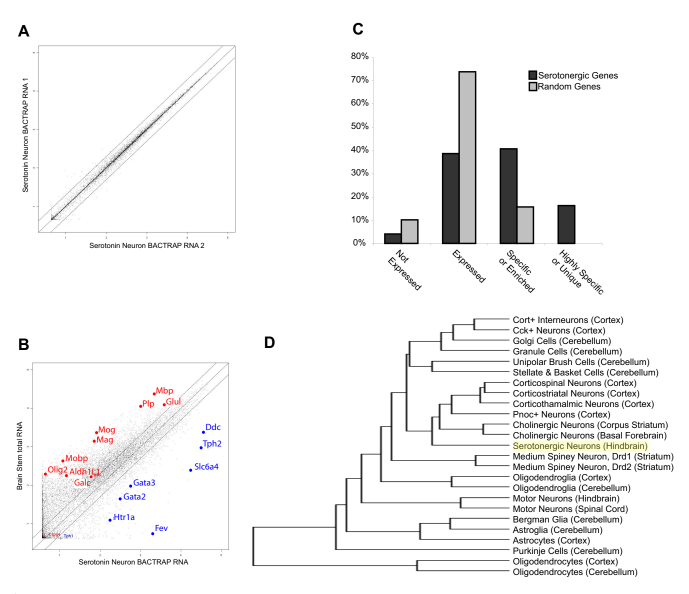


Figure 2. Reproducible and accurate polysome purification from serotonergic neurons. *A*, Scatterplot of probeset signal intensity (\log_{10}) for independent biological replicate TRAP experiments reveals extremely high reproducibility. Black lines indicate fold change = 0.5 and 2. *B*, Scatterplot of mRNA TRAPed from serotonergic neurons (average of 3 replicates, *x*-axis), compared with mRNA from hindbrain (average of 3 replicates, *y*-axis), reveals a robust enrichment for genes expressed in serotonergic neurons (blue), but not for known glial genes (red). *C*, Blinded analysis of *in situ* hybridization patterns for probesets predicted to be enriched in serotonergic neurons (light gray, n = 197), compared with randomly selected probesets (dark gray, n = 197), reveals a robust overrepresentation of raphe-specific and enriched genes in serotonergic neuron gene list ($p < 1E^{-66}$, χ^2 test). *D*, Hierarchical clustering of TRAP data for serotonergic neurons with previously characterized cell types (Doyle et al., 2008).

SNP rs2959930 is found near an alternative transcriptional start site of *CELF6*

The *CELF6* gene has 16 exons (Fig. 3A) and is highly homologous to other *CELF* genes, particularly *CELF3* (*TNRC4*), *CELF4*, and *CELF5*. Their long isoforms each contain three RNA recognition motifs (RRMs) that are nearly identical across family members, and a divergent domain, required in other family members for both splicing activity and repression (Han and Cooper, 2005). In other family members, the first two RRMs are required for RNA binding (Good et al., 2000), whereas the third may be involved in nuclear localization (Chapple et al., 2007).

The SNP associated with autism in the AGRE male probands, rs2959930, is found in the 5' UTR of an alternative first exon (Fig. *3C*). Inclusion of this exon precludes the incorporation of the first RRM (Fig. *3D*). In family member *CELF2*, truncations removing the first RRM reduce the inclusion of *GRIN1* exon 21 in a minigene splicing assay (Han and Cooper, 2005). This would

suggest that polymorphisms that impact the transcription, translation, or stability of the different isoforms of *CELF6* may have an impact on the splicing, localization, and translation of mRNA within the cell. However, it is unclear whether SNP rs2959930 is a relevant functional polymorphism or whether it is in linkage disequilibrium with a more profound variation. First, we sought to determine whether both isoforms are expressed in serotonergic neurons and to confirm that CELF6 is indeed being translated in these cells. Then, we initiated a screen for more profound polymorphisms that may be influencing this gene in autism.

Celf6 is found in serotonergic neurons

To determine which isoforms might be used in serotonergic neurons, we designed primers specific to the long and short isoforms. Quantitative RT-PCR clearly confirmed an enrichment of *Celf6* mRNA in serotonergic neurons (data not shown), with both isoforms detectable. To investigate whether Celf6 protein is also

Table 1. Serotonin cell-specific and enriched genes

Gene symbol	Probe set ID	pSI	Name		
Acsl4	1451828_a_at	6.54E-04	Acyl-CoA synthetase long-chain family member 4		
Ankrd2	1419621_at	1.72E-03	Ankyrin repeat domain 2 (stretch responsive muscle)		
Apoa2	1417950_a_at	9.57E-03	Apolipoprotein A-II		
Arhqdiq	1448660_at	4.87E-04	Rho GDP dissociation inhibitor (GDI) gamma		
lsb4	1433919_at	1.91E-06	Ankyrin repeat and SOCS box-containing protein 4		
Celf6	1429790_at	8.40E-03	Cugbp and Elav-like family member 6, RNA binding protein		
.1qtnf2	1431079_at	8.01E-03	C1q and tumor necrosis factor related protein 2		
Calcr	1418688_at	1.91E-06	Calcitonin receptor		
Calr3	1453232_at	2.20E-03	Calreticulin 3		
Cckar	1421195_at	1.43E-03	Cholecystokinin A receptor		
Cd1d1	1449131_s_at, 1449130_at	1.91E-06	CD1d1 antigen		
Ceacam10	1448573_a_at, 1417074_at	1.91E-06	CEA-related cell adhesion molecule 10		
Серб3	1436274_at	1.91E-06	Similar to protein C6orf117		
Chodl	1451440_at	3.57E-03	Chondrolectin		
Ciapin1	1438163 xat	1.97E-03	Rhomboid domain containing 2		
Cited1	1449031_at	1.91E-06	Cbp/p300-interacting transactivator with Glu/Asp-rich carboxy-terminal domain		
Col7a1	1419613_at	1.91E-06	Procollagen, type VII, alpha 1		
СраЗ	1448730 at	1.91E-06	Carboxypeptidase A3, mast cell		
Cpne8	1430521_s_at, 1430520_at	2.19 to 3.24E-03	Copine VIII		
ryba2	1419011_at	1.91E-06	Crystallin, β A2		
Cryouz Cthrc1	1452968_at	5.54E-05	Collagen triple helix repeat containing 1		
Ctla2b	1432908_at 1416811_s_at, 1448471_a_at	1.91E-06	cytotoxic T lymphocyte-associated protein 2 β		
Cxcl7		1.91E-06	Chemokine (C-X-C motif) ligand 7		
	1418480_at 1430591_at, 1426215_at				
Ddc		1.91E-06 1.91E-06	Dopa decarboxylase		
Dlk1	1449939_s_at		Delta-like 1 homolog (Drosophila)		
Dnajc12	1417441_at	3.44E-05	DnaJ (Hsp40) homolog, subfamily C, member 12		
E2f5	1434493_at	4.57E-04	E2F transcription factor 5, p130-binding		
En1	1418618_at	1.34E-05	Engrailed 1		
Esm1	1449280_at	2.87E-05	Endothelial cell-specific molecule 1		
Fbxw4 -	1442674_at	7.26E-03	F-box and WD-40 domain protein 4		
Fev	1425886_at	1.91E-06	FEV (ETS oncogene family)		
Fgd5	1460578_at	1.91E-06	FYVE, RhoGEF. and PH domain containing 5		
Foxa1	1418496_at	7.42E-03	Forkhead box A1		
Fxyd5	1418296_at	1.91E-06	FXYD domain-containing ion transport regulator 5		
Fxyd6	1417343_at	9.95E-03	FXYD domain-containing ion transport regulator 6		
Gas5	1449410_a_at	2.42E-04	Growth arrest-specific 5 (non-protein coding)		
Gata2	1428816_a_at, 1450333_a_at	1.91E-06	GATA binding protein 2		
Gata3	1448886_at	1.91E-06	GATA binding protein 3		
Gch	1429692_s_at, 1420499_at	1.91E-06	GTP cyclohydrolase 1		
Glra2	1434098_at	2.58E-04	Glycine receptor, α 2 subunit		
Gm1574	1446591_at	1.91E-06	Gene model 1574, (NCBI)		
Gnas	1421740_at	6.19E-04	GNAS (guanine nucleotide binding protein, $lpha$ stimulating) complex locus		
Gng2	1418451_at	6.11E-03	Guanine nucleotide binding protein (G protein), γ 2 subunit		
Gnpda2	1457230_at	3.19E-04	Glucosamine-6-phosphate deaminase 2		
Gpr1	1460123_at	5.89E-04	G protein-coupled receptor 1		
Gpr4	1457745_at	1.91E-06	G protein-coupled receptor 4		
Gpx3	1449106_at	1.34E-05	Glutathione peroxidase 3		
Gstm6	1422072_a_at	1.83E-04	Glutathione S-transferase, μ 1		
Guca1a	1421061_at	1.91E-06	Guanylate cyclase activator 1a (retina)		
Gus		3.05E-03	Glucuronidase, β		
H2-Q1	1418734_at	1.91E-06	histocompatibility 2, Q region locus 1		
Hcrtr1		1.91E-06	Hypocretin (orexin) receptor 1		
loxb3	1456229_at	4.80E-03	Homeo box B3		
Hs6st2	1420938_at, 1450047_at	2.05E-04 to 1.91E-06	Heparan sulfate 6-0-sulfotransferase 2		
Htr1a	1438710_at	1.91E-06	5-Hydroxytryptamine (serotonin) receptor 1A		
Htr1d	1440166_x_at, 1440741_at	4.22E-04 to 1.91E-06	5-Hydroxytryptamine (serotonin) receptor 1D		
Htr5b	1422196_at	1.91E-06	5-Hydroxytryptamine (serotonin) receptor 5B		
lqh-6	1427351_s_at, 1427329_a_at	5.75E-03 to 1.91E-06	Immunoglobulin heavy chain 6 (heavy chain of IgM)		
lgh-VJ558	1425763_x_at, 1421653_a_at	3.82 to 1.91E-06	Immunoglobulin heavy chain (J558 family)		
1911-VJSS8 1r1	1425765_X_al, 1421655_a_al 1448950_at	7.70E-04	Interleukin 1 receptor, type l		
ng1	1456857_at	2.10E-05	Inhibitor of growth family, member 1		
lrs4 VIL27	1441429_at	1.91E-06	Insulin receptor substrate 4		
Klk27	1421587_at	1.91E-06 6.65E-03	Kallikrein 1-related peptidase b27 Keratin complex 2, basic, gene 1		
Krt2—1	1422481_at				

Table 1. Continued

Gene symbol	Probe set ID	pSI	Name
Lgals8	1422661_at	4.22E-03	Lectin, galactose binding, soluble 8
Lst1	1425548_a_at	1.92E-03	Leukocyte specific transcript 1
Maob	1434354_at	1.91E-06	Monoamine oxidase B
Mbd1	1430838_x_at, 1453678_at	4.20E-03 to 7.45E-05	Methyl-CpG binding domain protein 1
Mcpt5	1449456_a_at	7.07E-05	Chymase 1, mast cell
Nanos1	1436648_at	8.47E-03	Nanos homolog 1 (Drosophila)
Nkx6-1	1425828_at	1.10E-03	NK6 transcription factor related, locus 1 (Drosophila)
Nos3	1434092_at	8.57E-03	Nitric oxide synthase 3 antisense
Nrl	1450946_at	1.91E-06	Neural retina leucine zipper gene
Oas1e	1416847_s_at	1.91E-06	2'-5' Oligoadenylate synthetase 1E
Pcbd	1418713_at	1.09E-04	Pterin 4 α carbinolamine dehydratase
Pcdha4	1424341_s_at	3.91E-03	Protocadherin α 4
Pcdha6	1451769_s_at	7.28E-03	Protocadherin α 6
Pcdhac1	1425017_at	1.91E-06	Protocadherin α subfamily C, 1
Pcsk1	1421396_at	1.91E-06	Proprotein convertase subtilisin/kexin type 1
Pcsk5		2.89E-03 to 6.66E-03	Proprotein convertase subtilisin/kexin type 5
Pdzk8	1439088_at	1.91E-06	PDZ domain containing 8
Prph1	1422530_at	3.16E-03	Peripherin 1
Ptger3	1450344_a_at	1.91E-06	Prostaglandin E receptor 3 (subtype EP3)
Resp18	1417988 at	2.03E-04	Regulated endocrine-specific protein 18
Rims3	1459042_at	1.75E-03	Regulating synaptic membrane exocytosis 3
Rpgrip1	1454231_a_at	1.91E-06	Retinitis pigmentosa GTPase regulator interacting protein 1
Rph3al	1444409_at	2.10E-05	Rabphilin 3A-like (without C2 domains)
Rpl37a	1416217_a_at	9.57E-03	Ribosomal protein L37a
Rps27	1460699_at	2.10E-05	Ribosomal protein S27
Saa1	1419075_s_at	1.51E-04	Serum amyloid A1
Scq2	1450708_at	3.12E-04	Secretogranin II
Scn9a	1442333_a_at, 1442810_x_at, 1442809_at	1.91E-06	Sodium channel, voltage-gated, type IX, α
Siat8f	1456147_at, 1456440_s_at	9.94E-05 to 1.91E-06	ST8 α -N-acetyl-neuraminide α -2,8-sialyltransferase 6
Slc18a2	1437079_at	1.91E-06	Solute carrier family 18 (vesicular monoamine), member 2
SIc22a3	1420444_at	1.91E-06	Solute carrier family 10 (vesicular monoamme), member 2 Solute carrier family 22 (organic cation transporter), member 3
SICZZUS SIC6a4	1417150_at	1.91E-06	Solute carrier family 22 (organic cation transporter), member 3 Solute carrier family 6 (neurotransmitter transporter, serotonin), member
Sicou4 Sin	1417150_at 1420884 at	3.20E-03	
	—	1.91E-06	Sarcolipin Survidain at
Sncg Stard2 ml	1417788_at		Synuclein, γ STARD3 N-terminal like
Stard3 nl	1430274_a_at	7.55E-03	
Stk32b	1431236_at	1.91E-06	Serine/threonine kinase 32B
Swam2	1449191_at	1.91E-06	WAP four-disulfide core domain 12
Tinag	1419314_at	1.91E-06	Tubulointerstitial nephritis antigen
Tmie This can	1441926_x_at, 1443964_at	1.26E-03 to 5.76E-03	Transmembrane inner ear
Tnfrsf11b	1449033_at	4.80E-03	Tumor necrosis factor receptor superfamily, member 11b (osteoprotegerin
Tph2	1435314_at	1.91E-06	Tryptophan hydroxylase 2
Trh	1418756_at	1.91E-06	Thyrotropin releasing hormone
Trpm4	1435549_at	5.66E-03	Transient receptor potential cation channel, subfamily M, member 4
Twsg1	1441302_at	8.37E-03	Twisted gastrulation homolog 1 (Drosophila)
Zar1	1434494_at	2.10E-05	Zygote arrest 1
Zfp622	1447775_x_at	8.60E-04	Zinc finger protein 622
Zwint	1444717_at	7.90E-03	ZW10 interactor

Probesets corresponding to all named genes with pSI < 0.01 are listed.

found in serotonergic neurons, we generated antibodies to Celf6. Quantitative RT-PCR of 3T3 cells suggested no endogenous expression of Celf6. Experiments with these cells transiently expressing long or short isoforms, or a GFP fusion, demonstrate that these antibodies can detect Celf6 by SDS-PAGE (Fig. 4A) or in fixed cells (Fig. 4B).

Immunofluorescence in Slc6a4 bacTRAP mice reveals colocalization of eGFP with Celf6 immunoreactivity in raphe neurons (Fig. 4*C*). Celf6 immunoreactivity is also seen in a few other neuromodulatory populations, including the noradrenergic locus ceruleus, populations of cells in the hypothalamus, including presumptive dopaminergic cells of the arcuate nucleus, and a few, scattered, dimly labeled cells in cortex (data not shown). Unlike other presumptive neuronal splicing factors, which are predominantly nuclear (Underwood et al., 2005), labeling is largely cytoplasmic, suggesting that in the brain Celf6 may have roles beyond splicing. Having confirmed Celf6 protein is found in serotonergic neurons, we next conducted a screen to identify additional variants that may be associated with human autism.

Resequencing of candidate exons in *CELF6* identifies rare premature stop codon

Toward an understanding of the extent to which *CELF6* tolerates genetic variation in humans, we used publicly available exome data from 1092 individuals, to compare SNV rates here with those in all other RefSeq protein coding genes (n = 19,032). Although tolerance for nonsynonymous variation was seen to vary, the large majority of annotated genes (n = 14,022) were observed to harbor one or more SNVs predicted to be deleterious (2+ of BLOSUM62, Polyphen2, and SIFT) (Sunyaev et al., 2001;

Table 2. Transmission disequilibrium in SNPs near two serotonergic neuron genes in human autistic probands

Chromosome	SNP	A1 (minor allele)	A2 (major allele 2)	No. of observations of minor allele transmission to individual with autism	No. of observations of untransmitted minor allele	OR	<i>p</i> value	Bonferonni- corrected p value	Gene
Male probands									
15	rs2959930	G	Α	273	378	0.72	3.87E-05	0.02521	CELF6
5	rs9313845	А	G	110	171	0.64	0.000274	0.1785	C1QTNF2
14	rs8003220	А	G	38	77	0.49	0.000276	0.18	IGH
5	rs17409286	А	G	144	203	0.71	0.001539	1	C1QTNF2
8	rs11573856	А	G	113	165	0.68	0.001816	1	TNFRSF11B
Female probands									
17	rs4470197	G	A	70	113	0.62	0.00148	0.9647	CCDC55
16	rs9929530	С	А	52	88	0.59	0.002346	1	TMEM114
23	rs4599945	А	С	6	22	0.27	0.002497	1	DGKK
5	rs3762986	A	G	93	138	0.67	0.003069	1	PCSK1
4	rs11731545	Α	G	90	56	1.61	0.004895	1	TNFRSF11B
Male and female probands									
5	rs9313845	A	G	129	202	0.64	6.01E-05	0.03918	C1QTNF2
14	rs8003220	А	G	48	92	0.52	0.0002	0.1306	IGH
15	rs2959930	G	А	352	451	0.78	0.000477	0.3107	CELF6
8	rs11573856	A	G	143	205	0.70	0.000889	0.5795	TNFRSF11B
10	rs363341	Α	G	400	498	0.80	0.001074	0.7005	SLC18A2

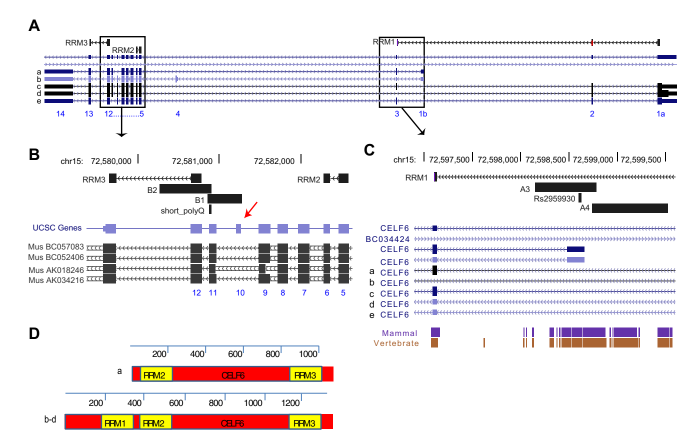


Figure 3. *CELF6* gene, exons, domains, and regions of interest. *A*, UCSC genome browser view of the *CELF6* gene in human, showing 16 exons (blue numbers), 5 putative isoforms (a– e), and alignment of the 3 RRMs. *B*, First candidate region, covered by two amplicons (B1 and B2), encompassing short poly-glutamine sequence (polyQ) and putative human-specific exon (red arrow). Human *CELF6* exons are shown condensed in blue, and aligned mouse mRNAs are shown below in black. *C*, Second candidate region, covered by two amplicons (A3 and A4), encompassing the alternative first exon, and highly conserved putative promotor region (purple, brown) and the associated SNP RS2959930. *D*, Illustration of *CELF6* protein isoforms. Isoform "a" of *CELF6*, which includes the alternative first exon (1b), does not contain the first RRM.

Ng and Henikoff, 2002; Eddy, 2004; Adzhubei et al., 2010) and present at an appreciable frequency in this population (>1%). Consistent with intolerance to nonsynonymous variation, no such events were observed within *CELF6*. Similarly, no structural variants overlapping with *CELF6* were observed in the DGV

(Iafrate et al., 2004). Although power issues currently limit a precise ranking of how tolerant different genes are to genetic variation, consideration of all coding variation put *CELF6* at the 97th percentile of all annotated genes, a result consistent with biological intolerance. Based on this result, we reasoned that a

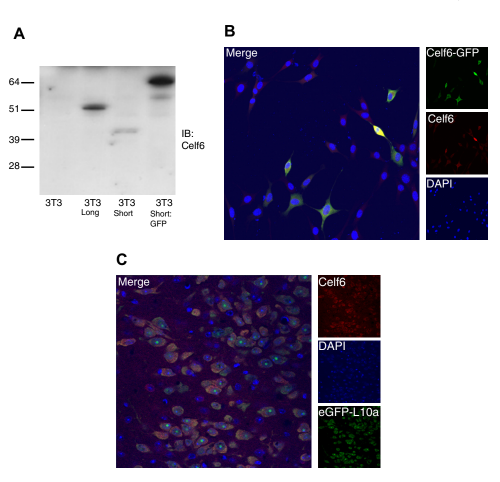


Figure 4. Celf6-like immunoreactivity in serotonergic nuclei. *A*, Celf6 immunoblot of whole-cell lysates from 3T3 fibroblasts transfected with a long isoform of Celf6, a short isoform of Celf6, or a short isoform of Celf6 fused to GFP demonstrates specificity of antibody to target. *B*, GFP and Celf6 immunofluorescence in 3T3 fibroblasts transfected with Celf6-GFP (as in *A*). *C*, Celf6 immunofluorescence in SIc6a4 bacTRAP mice demonstrates colocalization of Celf6 with eGFP-L10a in serotonergic neurons of dorsal raphe.

focused resequencing effort, enabling deep characterization of key functional regions, should be used.

Toward this end, we selected three regions for our first screen of *CELF6*. This included the area around the alternative first exon 1b, where the common variant was identified, as well as the exon 12, which contains a short CAG repeat (Fig. 3*B*). Finally, we also targeted the exon 11 because available EST data suggest that it is used exclusively by humans: no other available species show any mRNA or ESTs aligning to it (Fig. 3*B*). Of course, as human, mouse, and rat sequences dominate the EST databases, these data could represent a loss of the exon in rodents, rather than a gain in humans.

Thus, we sequenced the promoter and alternative first exon 1b, as well as exons 11–13 in 384 male white probands (randomly selected from previous AGRE probands) and 384 normal male white controls (new samples). In this semi-independent sample, we found no common alleles showing stronger association than rs2959930 (Fisher's exact test, p < 0.051), although one novel low-frequency intronic variant was seen only in controls (Table 3, p < 0.0012). We did not detect any instances of triplet repeat expansion in exon 12, although extraordinarily large repeats would not be assayable with PCR-based methods.

We did, however, discover a premature stop codon in one autistic patient (Fig. 5A) and no controls. This SNP is found in the putative human-specific exon 11 and would be predicted to lead nonsense-mediated decay, and thus is likely a loss of function allele. No coding sequence variations were identified in the 384 controls. The presence of this rare variant in an

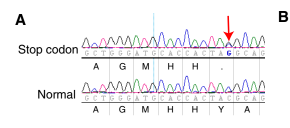
autistic patient provides further evidence implicating this gene in autism, although it clearly cannot account for the original transmission disequilibrium findings. Sequencing of the entire pedigree (AU1397) reveals that the variant was inherited from the father, suggesting incomplete penetrance if the allele is involved in autism. There is also additional strong matrilineal contribution for autism in this family, as the proband has affected twin siblings and matrilineal cousins not harboring this allele (Fig. 5*B*). Although not included in the AGRE consortium, the father's extended family also included additional individuals with intellectual disability and reportedly odd behavior.

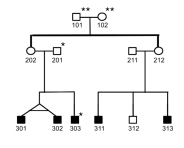
To determine whether this rare variant is found in any additional cases or controls, we designed both a primer-extension (Sequenom) assay as well as an allele-specific PCR (Fig. 5*C*, *D*). Analysis of 4992 normal individuals, as well as 864 autistic individuals from the Simons Simplex Collection, identified no additional cases or controls harboring this allele, highlighting the rarity of the variant. Likewise, this variant was not seen in the 1092 individuals above, or the ~5000 individuals currently available in the Exome Variant Server (http://evs.gs.washington.edu/ EVS/)[3/2012].

Like other cases of extremely rare variants detected in autism, such as many of the exonic deletions (Bucan et al., 2009) and other copy number variants (Pinto et al., 2010), it is difficult to amass sufficient power to draw statistical conclusions regarding

DBsnp	Amplicon	Position within amplicon	SNP	AGRE, % (no.) of samples with SNP detected, of interpretable reads	Controls, % (no.) of samples with SNP detected, of interpretable reads	<i>p</i> value (Fisher's test), deviating from expected (Controls)	Туре	Confirmed
Novel	B1	129	$C \rightarrow CG$	0.27% (1/370)	0% (0/355)	0.510	Stop	Yes ^a
Novel	B1	150	$C \rightarrow CG$	0.27% (1/370)	0% (0/355)	0.510	Intron	None
Novel	B1	247	$T \rightarrow CT$	0.81% (3/370)	0.57% (2/353)	0.522	Intron	None
Novel	B1	263	$T \rightarrow CT$	0.27% (1/370)	0.57% (2/353)	0.469	Intron	None
Novel	B2	88	$C \rightarrow CG$	0.53% (2/375)	0.29% (1/341)	0.536	Intron	None
Novel	B2	104	$T \rightarrow TC$	0% (0/375)	0.29% (1/341)	0.476	Intron	None
Novel	B2	200	$G \rightarrow AG$	0.28% (1/357)	0% (0/341)	0.511	Intron	None
Novel	B2	362	$G \rightarrow GA$	0% (0/375)	0.29% (1/341)	0.476	Intron	None
Novel	B2	391	$T \rightarrow CT$	0% (0/375)	0.29% (1/341)	0.476	Intron	None
Novel	B2	459	$G \rightarrow AG$	0% (0/375)	2.64% (9/341)	0.001	Intron	None
Novel	B2	513	$C \rightarrow CT$	0% (0/375)	0.29% (1/341)	0.476	Intron	None
rs4625684	A3	60	T→C	100% (282/282)	100% (357/357)	1.000	Promoter/intron	In dbSNP
rs2959930	A3	147	$C \rightarrow CT$	29.43% (83/282)	35.85% (128/357)	0.051	5' UTR	In dbSNP
rs2959930	A3	147	$(\rightarrow \uparrow$	6.74% (19/282)	3.64% (13/357)	0.055	5' UTR	In dbSNP
Novel	A3	151	$C \rightarrow CT$	0.71% (2/282)	1.12% (4/357)	0.459	5' UTR	None
rs74026061	A3	165	$G \rightarrow CG$	0.35% (1/282)	0% (0/357)	0.441	5' UTR	In dbSNP
Novel	A3	166	$C \rightarrow CT$	0% (0/282)	0.28% (1/357)	0.559	5' UTR	None
Novel	A3	234	$C \rightarrow CT$	0.35% (1/282)	0% (0/357)	0.441	5' UTR	None
rs2959928	A4	91	$C \rightarrow CT$	1.06% (4/377)	0% (0/363)	0.067	Promoter/intron	In dbSNP
rs2959929	A4	159	$G \rightarrow AG$	4.26% (16/376)	2.48% (9/363)	0.129	Promoter/intron	In dbSNP
Novel	A4	233	$C \rightarrow CT$	0.27% (1/376)	0% (0/363)	0.509	Promoter/intron	None
Novel	A4	50	$G \rightarrow AG$	0% (0/376)	0.28% (1/363)	0.491	Promoter/intron	None
Novel	A4	169	$C \rightarrow AC$	0% (0/376)	0.28% (1/363)	0.491	Promoter/intron	None

^aConfirmed in proband and father with bidirectional Sanger sequencing, as well as sequenome and allele-specific PCR assays.





D

Sample	Status	# Assayed	Positives
AGRE	Cases	384	1
SSC	Cases	864	0
NIMH-HGIC	Controls	4992	0

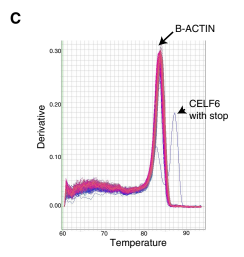


Figure 5. Premature stop codon in CELF6 in a family with autism. *A*, Novel coding SNP identified in human-specific exon. Top row, Proband 303 shows C->C/G. Bottom row, Normal DNA. Translation shown beneath each. Red arrow indicates heterozygote Y->stop SNP. *B*, Pedigree of Family AU1397. **Not sequenced. *Has stop Y-> Stop. *C*, Example of quantitative RT-PCR melt curves from 96 allele specific PCRs. 95 amplified only β-actin (genomic control). Positive control DNA (proband 303) amplified both β-actin and mutant Celf6 allele. *D*, Summary of all assays for mutant Celf6 allele.

the relevance of such an extremely rare variant. Also, similar to what is seen in deletions of NRXN1 and 16p11, this polymorphism, if contributing to causation, is not showing complete penetrance in the afflicted family (Abrahams and Geschwind, 2010).

Further statistical validation of this allele now awaits more complete resequencing studies of *CELF6*, a byproduct of ongoing large-scale autism exome sequencing efforts. However, even with these studies, the human genetics alone will still likely be under-

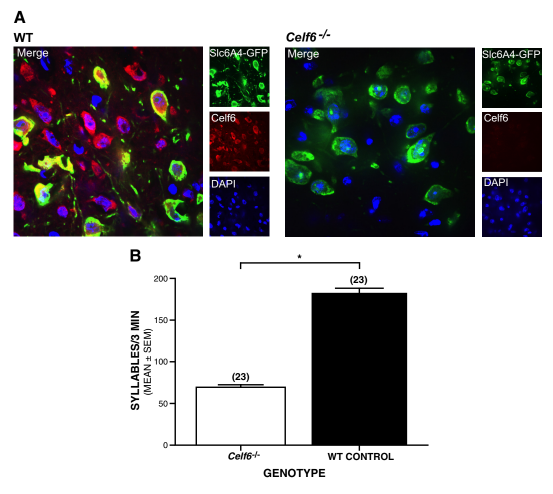


Figure 6. *Celf6* mutant mice show loss of Celf6 signal in brain, and *Celf6*^{-/-} mouse pups show deficits in early communicative behaviors. *A*, Immunofluorescent confocal microscopy of dorsal raphe neurons, labeled with GFP antibody (green), reveals loss of Celf6 antibody staining (red) in *Celf6*^{-/-} mice. *B*, *Celf6*^{-/-} mouse pups produce significantly fewer ultrasonic syllables per 3 min recording session compared with WT mice (ANOVA, *p = 0.001). Data are mean \pm SEM.

powered. In addition, reanalysis of published microarray datafrom human lymphoblasts reveals that CELF6 is not detectable in these cells (Luo et al., 2012), precluding any straightforward patient-based allelic expression imbalance analyses. Therefore, we decided to investigate this allele functionally by creating a mutation in mouse and testing for autism-like behaviors.

Characterization of Celf6 mutant mice

Autism is characterized by abnormal social interactions and language delay, as well as restricted interests and resistance to change. To determine whether mutations in Celf6 could contribute to analogous abnormalities in mouse behavior, we deleted the constitutive fourth exon of Celf6. This deletion introduces premature stop codons in all known isoforms of Celf6 and, like the human mutation, would be predicted to result in nonsensemediated decay of mRNA. PCR confirmed the deletion of the locus in the genome. Quantitative RT-PCR from mouse brain confirmed decrease of Celf6 mRNA (data not shown). Examination of the brain using nuclear stains (DAPI) and neuronal staining (NeuN) did not reveal any gross morphological differences. However, immunofluorescence using Celf6 antibodies confirms a loss of Celf6 signal from cells in all regions of the brain, including raphe neurons (Fig. 6A). Behavioral studies were then conducted to characterize Celf6^{-/-} and littermate WT control mice during both development and later in adulthood. It was our goal not only to study behaviors that may be related to autism but also to broadly characterize the phenotype of $Celf6^{-/-}$ mice also, as this is the first report of deletion in mice.

Celf6^{-/-} pups have decreased ultrasonic vocalizations

To determine whether $Celf6^{-/-}$ mice exhibited any alterations in communication-related behaviors during early development, we conducted the standard maternal isolation test, which is frequently used to evaluate mouse models of autism (Hofer et al., 2002; Chadman et al., 2008; Scattoni et al., 2008; Nakatani et al., 2009). Eight-day-old mouse pups exhibit robust stereotyped vocalizations in the ultrasonic range when separated from their dam. These calls ("syllables") (Holy and Guo, 2005) can be considered a form of social communication as they elicit a retrieval response from the dam. $Celf6^{-/-}$ pups produced 60% fewer syllables per 3 min recording session compared with WT littermates (Fig. 6*B*, ANOVA; $F_{(1,42)} = 12.794$, p = 0.001). The significant reduction in vocalization in *Celf6^{-/-}* pups was further observed between genotypes within each sex separately ($F_{(1,42)} = 5.792$, p = 0.02 for females, $F_{(1,42)} = 7.049$, p = 0.01 for males; not shown) with no significant difference between sexes. There was no difference in body weights between groups, suggesting that the vocalization effect was not the result of gross developmental delay (data not shown). Results from this assay can be interpreted as reflecting alterations in anxiety levels or communicative function (Hofer et al., 2002). However, we think that the decreased USV in $Celf6^{-/-}$ mouse pups probably reflects communication deficits because we found no evidence of abnormally high levels of anxiety-like behaviors in $Celf6^{-/-}$ mice when they were tested as adults in the elevated plus maze (EPM), at ages when this test has been well validated (see results described below for the EPM).

Adult male *Celf6^{-/-}* mice exhibit several normal behavioral/cognitive functions

Adult male Celf6^{-/-} and WT mice performed similarly on several basic behavioral assays, including a 1 h locomotor activity test and a battery of sensorimotor measures (e.g., Fig. 7A, B). For example, no differences were observed between groups in terms of general ambulatory activity (Fig. 7A) or vertical rearing (not shown) during a 1 h locomotor activity test. In addition, no performance deficits were observed in the *Celf6*^{-/-} mice on the ledge test (Fig. 7B) or on any of the other 6 sensorimotor measures that were conducted (not shown). The Celf $6^{-/-}$ mice also generally performed at control-like levels on the Morris water maze. Specifically, there were no effects involving genotype for either escape path length (Fig. 7C) or latency (not shown) during the cued trials, although the $Celf6^{-/-}$ mice swam significantly (25%) faster than the WT control group (Fig. 7D; rmANOVA-genotype effect: $F_{(1,18)} = 8.13$, p = 0.011). Importantly, no differences were observed between groups with regard to spatial learning and memory; no effects involving genotype were found for path length (Fig. 7E) or latency (not shown) during the place (hidden platform) trials, and no retention performance deficits were evident during the probe trial in terms of spatial bias (Fig. 7F), platform crossings, or time in the target quadrant (data not shown). The increased swimming speeds in the $Celf6^{-/-}$ mice during the cued trials were a transitory effect because no differences were observed between groups on this variable during the place (spatial learning) trials. Sensorimotor reactivity and gating were also evaluated in the mice, but no significant effects involving genotype were found for either the acoustic startle response (data not shown) or PPI (Fig. 7G). In summary, the above results suggest that Celf6^{-/-} mice

Dougherty et al. • Disruption of *Celf6*

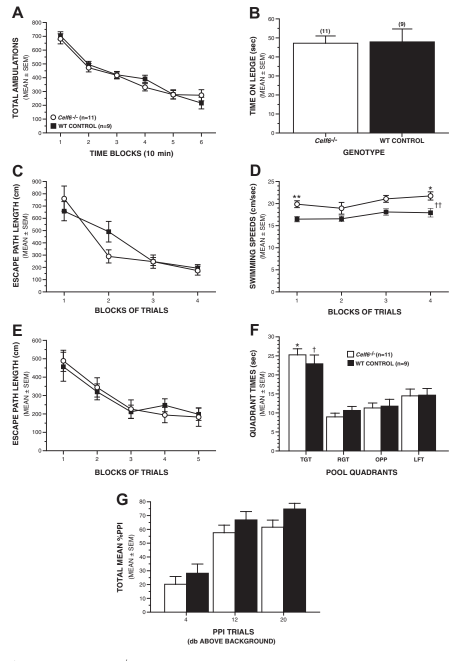


Figure 7. Adult male *Celf6*^{-/-} mice demonstrate intact sensorimotor and spatial learning functions. *A*, *B*, Adult male *Celf6*^{-/-} and WT mice performed similarly on several basic behavioral assays. *A*, Comparable levels of general ambulatory activity were observed in *Celf6*^{-/-} and WT mice during the 1 h locomotor activity test. *B*, *Celf6*^{-/-} mice did not exhibit any performance deficits on the ledge test or on any of the other 6 measures within the sensorimotor battery (data not shown). *C*-*F*, The *Celf6*^{-/-} mice also generally performed at control-like levels on the Morris water maze. *C*, *Celf6*^{-/-} and WT controls exhibited comparable path lengths to the escape platform during cued trials, indicating that the *Celf6*^{-/-} mice did not display nonassociative dysfunctions that would interfere with subsequent learning and memory performance in the water maze. *D*, *Celf6*^{-/-} mice swam significantly faster than the WT controls during the cued trials in the water maze (mANOVA–genotype effect, $\pm p = 0.011$, pairwise comparisons: ***p* = 0.004 and **p* = 0.012). *E*, WT and *Celf6*^{-/-} groups performed equally well during the place trials in the water maze, suggesting that spatial learning was not impaired. *F*, During the probe trial, both the *Celf6*^{-/-} and WT mice spent more time in the target (TGT) quadrant where the platform had previously been (mANOVA with planned comparisons, **p* < 0.003 and $\pm p < 0.025$, respectively) compared with the right (RGT), opposite (OPP), and left (LFT) quadrants, suggesting that retention was not impaired. *G*, Equivalent levels of PPI were observed in the two groups of mice, indicating normal sensorimotor gating and startle responses. Data are mean \pm SEM.

did not have demonstrable deficits in nonassociative (visual, sensorimotor, or motivational) functions and that their spatial learning and memory appeared to be intact, at least for reference memory-based capabilities. The mice were also evaluated on standard social interaction assays involving the social approach test (Moy et al., 2004; Silverman et al., 2011), the results of which showed that the $Celf6^{-/-}$ mice performed like WT controls. In the first social

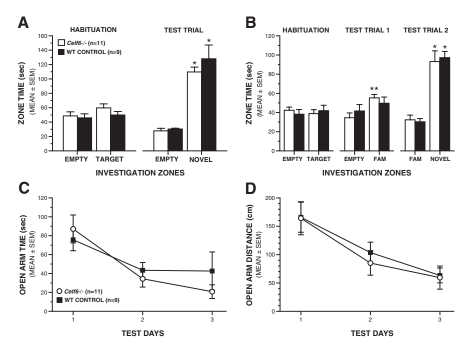


Figure 8. Adult *Celf6*^{-/-} mice exhibit control-like levels of sociability, social preference, and anxiety-like behaviors. *A*, During the first social approach test, adult *Celf6*^{-/-} and WT mice both showed normal sociability preference for a novel stimulus mouse over an empty withholding cage during the test trial (rmANOVA with planned comparisons, **p* < 0.0004). *B*, During the second social approach test, adult *Celf6*^{-/-} mice spent more time investigating the cagemate compared with the empty withholding cage during test trial 1 (***p* < 0.005), demonstrating no deficits in sociability toward a familiar cagemate. During test trial 2, adult *Celf6*^{-/-} and WT mice both displayed a preference for social novely by spending more time investigating the novel stimulus mouse compared with a familiar cagemate during test trial 2 (rmANOVA with planned comparisons, **p* < 0.0003). *C*, *D*, Adult *Celf6*^{-/-} and WT mice showed similar levels of anxiety-like behaviors as indexed by the amount of time spent (*C*) and distance traveled (*D*) in the open arms of the elevated plus maze. Data are mean ± SEM.

approach test sequence, neither Celf6^{-/-} nor WT control mice demonstrated investigation zone biases during the habituation trial (empty vs target zone, Fig. 8A), presenting no confounding issues for the interpretation of the test trial results. We assessed sociability exhibited toward a novel stimulus mouse versus an empty withholding cage during the test trial, and rmANOVAs conducted on the relevant variables did not reveal significant effects involving genotype. More importantly, planned comparisons conducted within each group showed that both groups spent significantly more time investigating a novel stimulus mouse compared with an empty withholding cage, whether measured by time in the investigation zone surrounding the cage (Fig. 8A; $F_{(1,18)} = 37.30, p = 0.000009$ for *Celf6^{-/-}*, $F_{(1,18)} = 43.45, p = 0.000003$ for WT), or by time spent in the chamber ($F_{(1,18)} = 17.30, p =$ 0.0006 for Celf6^{-/-}, $F_{(1,18)} = 25.75$, p = 0.0008 for WT not shown). A second social approach test conducted at a slightly later age involved quantifying social preference for a same-sex cagemate versus a novel stimulus mouse. Again, neither Celf6 nor WT control mice demonstrated investigation zone biases during the habituation trial (empty vs target zone, Fig. 8B). The $Celf6^{-/-}$ mice showed no deficits in sociability toward a cagemate during test trial 1. No differences were found between $Celf6^{-/-}$ and WT control mice during test trial 2 in that both groups of mice spent significantly more time in the investigation zone (Fig. 8*B*; $F_{(1,16)} = 30.46$, p = 0.00005 for *Celf6^{-/-}*, $F_{(1,16)} = 23.51$, p = 0.0002 for WT) and chamber ($F_{(1,16)} = 12.59$, p = 0.003 for *Celf6^{-/-}*, $F_{(1,16)} = 9.28$, p = 0.008 for WT) containing a novel stimulus mouse compared with the times spent in the same areas where a cagemate was contained.

Given the robust decreases in USV observed in the $Celf6^{-/-}$ mouse pups, we were interested to determine whether they

showed high levels of anxiety-like behaviors in the EPM because it might have relevance to interpreting the vocalization results. Specifically, the $Celf6^{-/-}$ mice exhibited control-like levels in terms of time spent (Fig. 8C), distance traveled (Fig. 8D), and entries made into the open arms, as well as the percentages of these variables computed with reference to totals observed in both sets of arms (not shown). Thus, analyzing the classic variables thought to represent anxiety-like behaviors in the EPM did not reveal any differences between the Celf6^{-/-} and WT control mice. These results suggest that the USV deficit was not likely the result of any consistent lifelong differences in anxiety but instead may reflect an impairment in early social communication.

Celf6^{-/-} mice show evidence of resistance to change

Another diagnostic criterion of autism involves resistance to change. Although $Celf6^{-/-}$ mice exhibited intact spatial learning and retention during in the Morris water maze, they swam faster than the WT control group when introduced into the pool during the cued trials, an effect that disappeared with extended exposure during the place trials. This initial increase in swimming speeds in the *Celf6^{-/-}* mice

suggests an altered response in these mice to a significant change in environmental conditions. To evaluate the mice more formally for resistance to change, we evaluated their performance during reversal trials in the water maze. In the first block of trials, the $Celf6^{-/-}$ mice exhibited a tendency toward considerably longer escape latencies and path lengths (Fig. 9A, B), although planned comparisons showed that these differences were not significantly different ($F_{(1,18)} = 3.88$, p = 0.065 and $F_{(1,18)} = 3.45$, p = 0.080, respectively). Consistent with the data from the place trials, the groups performed in a similar manner in the later blocks of trials, with regard to swimming speeds, and during the probe trial (data not shown).

Based on the differences observed during the first block of reversal trials, we decided that a less stressful behavioral task, which involved exploratory behavior using the primary sensory system of mice (olfaction), might provide a more sensitive test of the resistance-to-change phenotype. A type of holeboard task has been used to study behaviors in mice considered to be analogous to the resistance to change and restricted interests found in people with autism-spectrum disorder (Moy et al., 2008; Silverman et al., 2011). Our modified version of this holeboard test (Ghoshal et al., 2012) was used to evaluate both exploratory and olfactory preference behaviors and whether familiarization with the putative reward value of an odorant could alter behavior. Figure 9C shows a schematic of the holeboard apparatus as well as the placement of odorants into the corner holes for test sessions 1 and 2. The results from ANOVAs showed that, initially, $Celf6^{-/-}$ and WT mice displayed similar general hole-poke frequencies with regard to total and corner hole-pokes (Fig. 9D, E, left), as well as side (empty) hole-pokes (not shown). Within-subjects comparisons revealed that both groups also poked significantly

more often ($F_{(1,18)} = 17.83$, p = 0.0005 for *Celf6^{-/-}*, $F_{(1,18)} = 8.58$, p = 0.009 for WT) into the hole containing a familiar odorant (fresh homecage bedding) than the empty corner hole (Fig. 9*F*, left), suggesting that both *Celf6^{-/-}* and WT mice have intact olfactory abilities, and a baseline preference for familiar bedding, as do other strains of mice (Moy et al., 2008). Poke frequencies did not differ between groups for any odorant or the empty corner hole at baseline.

After test session 1, the mice were familiarized with the chocolate chips by permitting consumption over a two-day period before they were tested on the holeboard again, with the odorants being placed in the corner holes as depicted in Figure 9C (right, test session 2). Familiarization with the chocolate chips had a profound effect on exploratory hole-poking and olfactory preference behaviors in the WT control mice but had no effect on the $Celf6^{-/-}$ group. For example, the results of ANOVAs showed that, after familiarization, WT mice made significantly greater numbers of total hole-pokes (Fig. 9D, $F_{(1,18)} = 12.46$, p = 0.002), corner hole-pokes (Fig. 9E, $F_{(1,18)} = 16.71$, p = 0.0007), and side hole-pokes ($F_{(1,18)} = 6.185$, p = 0.023; data not shown) compared with the Celf6^{-/-} group. Most importantly, familiarization with the chocolate also had a major impact on differences in poke frequencies into the odorant-containing and empty corner holes (Fig. 9F, right). For example, a significant genotype by hole interaction $(F_{(3,54)} = 2.814, p =$ 0.048) for test session 2, followed by subsequent comparisons showed that the WT mice poked significantly more often into the holes containing the novel bedding $(F_{(1,18)} = 9.90, p = 0.006)$, and particularly the chocolate chips ($F_{(1,18)} = 15.91$, p = 0.0009), compared with the Celf6^{-/-} group, with large differences also observed between groups in hole-pokes into the hole containing familiar bedding $(F_{(1,18)} = 6.51, p = 0.020, Bonferroni$ corrected critical p = 0.013). Moreover, within-subjects comparisons revealed that the $Celf6^{-/-}$ mice showed no preference for any odorant over the empty corner hole during test session 2, whereas the WT group showed significant preferences $(F_{(3,16)} = 14.13, p = 0.00009)$, for the familiar bedding (p = 0.0002), novel bedding (p = 0.006), and chocolate (p = 0.006)0.00005) over the empty corner hole (Bonferroni corrected critical p = 0.017). Thus, familiarization with the chocolate had a robust effect on WT control mice in terms of increasing their general exploratory hole poking and changing their ol-

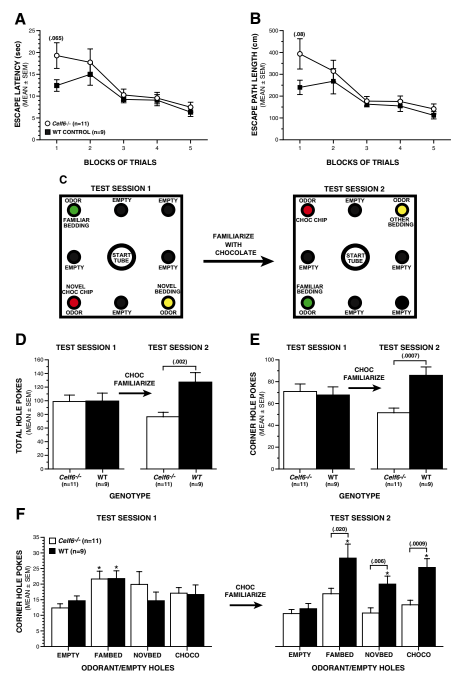


Figure 9. Adult *Celf6^{-/-}* mice showed evidence of resistance to change. *A*, *B*, There were no significant performance differences between groups during the reversal trials in the water maze, although there were nonsignificant trends for the Celf6⁻ mice to have longer escape latencies (A, rmANOVA with pairwise comparisons, p = 0.065) and path lengths (B, rmANOVA with pairwise comparisons, p = 0.080) compared with the WT group. *C–F*, Holeboard assay for resistance to change. *C*, A schematic of the holeboard showing the location of odorants and empty holes for test session 1 and test session 2 (after familiarization with chocolate). **D**, Celf6 $^{-/-}$ and WT mice showed equivalent exploratory behavior (hole-poke frequencies) during test session 1. Celf6^{-/-} mice failed to show the potentiation of exploratory behavior after familiarization with chocolate that was exhibited by the WT controls in terms of total hole-pokes (ANOVA, p = 0.002). **E**, The increase in exploratory behavior by WT mice occurred largely in the corner holes (ANOVA, p = 0.0007). **F**, Before familiarization, Celf6^{-/-} and WT mice showed equivalent exploratory behavior as both groups showed preference for holes containing familiar bedding compared with the empty corner holes (rmANOVA with planned comparisons, *p < 0.010), suggesting normal olfactory capacities for both groups (test session 1). After familiarization, WT mice poked significantly more often than the Celf6 $^{-/-}$ mice into the holes containing novel bedding (rmANOVA with planned comparisons, p = 0.006) and, particularly, chocolate chips (p =0.0009), with large differences into the hole containing familiar bedding (p = .020). Overall, the WT group also showed significant preferences for the holes containing odorants over the empty corner hole (*p < 0.007), whereas the Celf6 ⁻ mice failed to show any change in exploratory hole poking or evidence of olfactory preferences following familiarization to the chocolate. Data are mean \pm SEM.

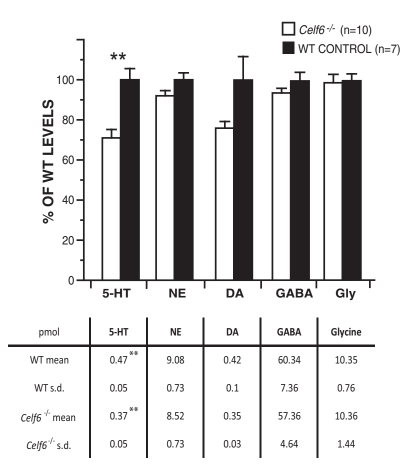


Figure 10. Celf6^{-/-} mice show decreased brain serotonin levels. LC-MS/MS measures of levels of neuromodulatory neurotransmitters serotonin (5HT), norepinephrine (NE), and dopamine (DA) reveal a significant decrease (**p < 0.002) in 5HT levels, and nonsignificant trend for NE and DA in whole brains from n > 7 WT and Celf6^{-/-} mice. There is no change in the negative control transmitters GABA and glycine. Data are plotted as a percentage of WT ion count levels; error bars represent SEM. Tables provide absolute levels in pMol.

factory preferences, whereas the $Celf6^{-/-}$ group showed a resistance to change on either of these two behavioral dimensions after exposure to the chocolate.

We also measured duration of hole-pokes to determine whether it was likely that the $Celf6^{-/-}$ and WT groups were processing the olfactory stimuli in a different manner. Our analyses showed that there were no differences between the groups with regard to the durations of total hole-pokes or for pokes made into odorant-containing or empty holes for either test session (data not shown). These findings along with the similar hole-poking performances of the groups before familiarization with the chocolate provide evidence that the groups did not differ in their abilities to discriminate odors. Rather, our holeboard results suggest that mutations in *CELF6* may contribute autism risk by influencing the propensity to modify behavior in response to a rewarding experience. More generally, our results provide evidence that *Celf6* contributes to a subset of the autism-like behaviors that were assayed in the present study.

Celf6^{-/-} mice have abnormal levels of CNS serotonin

Finally, our initial bacTRAP screen was to identify genes that may be important for the regulation of the serotonergic system in the brain (Figs. 1, 2), and we determined that Celf6 protein in mice was indeed found fairly specifically in monoamine-containing neurons within the brainstem (Fig. 4). To determine whether the changes in behavior seen might indeed be the result of perturbations of monoaminergic systems, we used mass spectrometry to measure total levels of three monoamines in the brains of Celf6^{-/-} and WT mice. We observed a $30 \pm 4\%$ (mean \pm SEM) decrease in serotonin levels (p < 0.002, Bonferroni corrected), and similar trends in norepinephrine and dopamine (p < 0.08, < 0.04, respectively, before Bonferroni corrections, not significant after) in brains of $Celf6^{-/-}$ mice (Fig. 10). There was no effect on levels of the GABA, glycine, or glutamate (not shown) neurotransmitters (*p* > 0.88, *p* > 0.23, and *p* > 0.28, respectively, no Bonferroni correction), demonstrating the specificity of the effect to neurons previously seen to be expressing Celf6.

Discussion

The paradigms of the previous era of human genetics, typified by observations of a suspicious looking variant in a few families and nothing similar in a few hundred families, are insufficient to cope with the overwhelming catalogs of variants being discovered now (Neale et al., 2012; O'Roak et al., 2012; Sanders et al., 2012). Results like those reported here are becoming standard: an inherited variant within a compelling candidate gene is observed only once in thousands of families. This highlights the need not only to increase our sample sizes, but also to find new analytical approaches to this complexity to overcome issues of power. We propose here that a broader adoption of blended strategies, drawing simultane-

ously on statistical genetics and empirical biology, are required.

There is a growing consensus that many psychiatric disorders are strongly influenced by a diverse set of rare, nearly private mutations (McClellan and King, 2010; Pinto et al., 2010). Thus, it is likely that there are multiple genetic routes to developing a particular disorder, or even individual symptoms within a disorder. Yet these diverse genetic routes must converge at some smaller number of common pathways in their molecular or cellular neurobiology to engender a common behavioral phenotype. Therefore, we have combined a unique and unbiased method for generating key biological priors in mice, with analysis of genetic variation in humans, and functional validation in murine models, to identify a gene contributing to some of the essential diagnostic features of autism. We propose that this kind of paradigm, leveraging information at multiple analytical levels, may be required to dissect complex genetic disorders. By working from the hypotheses that serotonergic abnormalities may contribute to a reasonable fraction of the cases of autism, we were able to reduce the penalties taken by looking at common variation genomewide, as well as appropriately direct our functional assays in mice. More broadly, although the evidence linking serotonergic neurons to some cases of autism is alluring, the true extent of the contribution of this cell type remains to be established. Several lines of evidence suggest that a subset of cases of autism may involve a perturbation of the serotonergic system. Multiple studies have detected abnormally high levels of serotonin in whole blood of a subset (>25%) of autistic patients (Schain and Freedman, 1961; Takahashi et al., 1976; Anderson et al., 1987; Cook, 1990). A combination of genetic investigation, neural imaging, and drug treatment have all implicated corresponding dysregulation of serotonin-related genes and pathways in affected individuals (McDougle et al., 1996b, 1997; McDougle et al., 1998; Chugani et al., 1999; Chandana et al., 2005; Coon et al., 2005; Hollander et al., 2005). For example, selective serotonin reuptake inhibitors have been shown to alleviate rigid/compulsive behaviors in some cases (Gordon et al., 1993; McDougle et al., 1996b; Hollander et al., 2011) and depletion of serotonin in autistic individuals exacerbates these symptoms (McDougle et al., 1996a). Likewise, genetic or pharmacological perturbation of the serotonergic system in rodents alters behavioral responses which characterize animal models of autism, such as altered USV, repetitive behavior, social interaction, and resistance to change (Olivier et al., 1998; Fish et al., 2000; Lopez-Rubalcava et al., 2000; Whitaker-Azmitia, 2001; McNamara et al., 2008; Nakatani et al., 2009). Most recently, Veenstra-VanderWeele et al. (2012) have shown that introduction of an autism-associated human coding variant in the serotonin transporter gene into mice resulted in a number of autism-related behaviors, such as impaired USV, decreased social dominance, and increased repetitive behaviors.

Our behavioral results suggest that, although $Celf6^{-/-}$ and WT mice performed similarly on several different tests, the Celf6^{-/-} mice exhibited certain selective functional deficits that may be analogous to the behavioral impairments that exist in a subset of people with autism. Specifically, Celf6^{-/-} and WT mice performed similarly on several behavioral assays, including: a 1-h locomotor activity test; a battery of sensorimotor measures; spatial learning and memory performance in the Morris water maze; various tests of social interaction as indexed by the social approach test; and anxiety-like behaviors quantified in the elevated plus maze. However, reduction of isolation-induced USV suggests that Celf6^{-/-} pups show early communication deficits. The measurement of USV in response to maternal isolation is an established assay that has been used in several other mouse models of autism, including the BTBR, Shank1 null, and neuroligin models (Chadman et al., 2008; Scattoni et al., 2008; Nakatani et al., 2009). Although genotypic differences in pup USV may be the result of either communication deficits or changes to anxiety, we feel the latter is unlikely as adult $Celf6^{-/-}$ mice did not exhibit differences compared with littermate WT control mice in anxiety-like behaviors measured during elevated plus maze testing. However, we acknowledge that it will be important to evaluate vocalization in $Celf6^{-/-}$ mice at later stages, such as during juvenile play and adult mating, and this is one of our future research aims. Our behavioral data also suggest that Celf6^{-/-} mice react abnormally to certain environmental alterations and may be resistant to changing their behavior under conditions that robustly alter WT mouse behavior. For example, $Celf6^{-/-}$ swam faster than controls when first introduced into the water maze during cued trials, but this effect habituated and was not present during the place condition. Also, familiarizing the mice with chocolate chips during the holeboard exploration/olfactory preference test led to robust changes in the behavioral responses of the WT control group, whereas $Celf6^{-/-}$ mice showed a resistance to such changes. Specifically, familiarization with the chocolate in WT mice led to greatly increased exploratory hole poking and distinct olfactory preferences for several odorants, whereas $Celf6^{-/-}$ showed no such changes. These results are similar to those reported by Moy et al., 2008 who used a slightly different procedure to show that BTBR mice exhibit a similar resistance to change after familiarization with chocolate. Our holeboard results provide evidence that mutations in *CELF6* may promote autism risk by modifying behavioral responses to rewarding experiences. However, it will be necessary to further characterize the resistance to change in olfactory preference in *Celf6^{-/-}* mice after exposure to various rewarding odorant stimuli, and we are conducting additional studies to this end. Our initial behavioral data suggest that *Celf6^{-/-}* mice exhibit selective deficits and may serve as a model for understanding a subset of autism-like behaviors.

Still, additional mechanistic studies are needed to inform both the Celf6 and the serotonergic contribution to autism in general. Although Celf6 is clearly enriched in these cells, and serotonin is decreased in the brains of $Celf6^{-/-}$ mice, conditional deletion and rescue studies are needed to confirm the behavioral abnormalities are cell-autonomous to the serotonergic system. Likewise, although we have provided evidence for this gene at multiple levels (a common 5'UTR variant and a rare heterozygous stop codon in humans, and homozygous deletions in mouse), each of these levels warrants additional investigation to generate a comprehensive view of the contribution of this gene. Although the common variant is linked with autism in the AGRE sample, this finding does not replicate in the Simons Simplex Collection (J. K. Lowe, personal communication), although that sample was smaller and optimized for finding idiopathic and especially de novo events rather than inherited, familial causes. The most parsimonious interpretation of our CELF6 findings thus far is that CELF6 levels may influence some behaviors relevant to autism; thus, there are alleles in this gene that contribute to risk of developing autism in a subset of patients. This would be consistent with autism being a highly heterogeneous and polygenic disorder; there are likely to be many genes with this type and magnitude of effect. This may also explain the incomplete penetrance seen in the family with the CELF6 mutation: resistance to change, in the absence of social deficits, would be insufficient to generate a diagnosis for autism. This would be consistent with recent studies, which suggest that the restricted interests and sociability components are factorially separable and partially genetically independent (Happe and Ronald, 2008; Frazier et al., 2012; Robinson et al., 2012). Thus, other genes and cell types likely contribute more to the social symptoms. An unbiased screen using this same approach systematically across many cell types may permit inferences regarding circuitry important to the neurobiology of autism.

References

- Abrahams BS, Geschwind DH (2010) Genetics of autism. In: Vogel and Motulsky's human genetics: problems and approaches, Ed 4 (Speicher MR, Antonarakis SE, Motulsky AG, eds), pp 699–714. Berlin: Springer.
- Adzhubei IA, Schmidt S, Peshkin L, Ramensky VE, Gerasimova A, Bork P, Kondrashov AS, Sunyaev SR (2010) A method and server for predicting damaging missense mutations. Nat Methods 7:248–249. CrossRef Medline
- American Psychiatric Association (2000) Diagnostic and statistical manual of mental disorders: DSM-IV-TR, Ed 4. Washington, DC: American Psychiatric Association.
- Anderson GM, Freedman DX, Cohen DJ, Volkmar FR, Hoder EL, McPhedran P, Minderaa RB, Hansen CR, Young JG (1987) Whole blood serotonin in autistic and normal subjects. J Child Psychol Psychiatry 28: 885–900. CrossRef Medline
- Barreau C, Paillard L, Méreau A, Osborne HB (2006) Mammalian CELF/ Bruno-like RNA-binding proteins: molecular characteristics and biological functions. Biochimie 88:515–525. CrossRef Medline
- Bengel D, Jöhren O, Andrews AM, Heils A, Mössner R, Sanvitto GL, Saavedra

JM, Lesch KP, Murphy DL (1997) Cellular localization and expression of the serotonin transporter in mouse brain. Brain Res 778:338–345. CrossRef Medline

- Boyle MP, Kolber BJ, Vogt SK, Wozniak DF, Muglia LJ (2006) Forebrain glucocorticoid receptors modulate anxiety-associated locomotor activation and adrenal responsiveness. J Neurosci 26:1971–1978. CrossRef Medline
- Bucan M, Abrahams BS, Wang K, Glessner JT, Herman EI, Sonnenblick LI, Alvarez Retuerto AI, Imielinski M, Hadley D, Bradfield JP, Kim C, Gidaya NB, Lindquist I, Hutman T, Sigman M, Kustanovich V, Lajonchere CM, Singleton A, Kim J, Wassink TH, et al. (2009) Genome-wide analyses of exonic copy number variants in a family-based study point to novel autism susceptibility genes. PLoS Genet 5:e1000536. CrossRef Medline
- Canli T, Lesch KP (2007) Long story short: the serotonin transporter in emotion regulation and social cognition. Nat Neurosci 10:1103–1109. CrossRef Medline
- Chadman KK, Gong S, Scattoni ML, Boltuck SE, Gandhy SU, Heintz N, Crawley JN (2008) Minimal aberrant behavioral phenotypes of neuroligin-3 R451C knockin mice. Autism Res 1:147–158. CrossRef Medline
- Chandana SR, Behen ME, Juhász C, Muzik O, Rothermel RD, Mangner TJ, Chakraborty PK, Chugani HT, Chugani DC (2005) Significance of abnormalities in developmental trajectory and asymmetry of cortical serotonin synthesis in autism. Int J Dev Neurosci 23:171–182. CrossRef Medline
- Chapple JP, Anthony K, Martin TR, Dev A, Cooper TA, Gallo JM (2007) Expression, localization and tau exon 10 splicing activity of the brain RNA-binding protein TNRC4. Hum Mol Genet 16:2760–2769. CrossRef Medline
- Charlet-B N, Logan P, Singh G, Cooper TA (2002) Dynamic antagonism between ETR-3 and PTB regulates cell type-specific alternative splicing. Mol Cell 9:649–658. CrossRef Medline
- Chonchaiya W, Schneider A, Hagerman RJ (2009) Fragile X: a family of disorders. Adv Pediatr 56:165–186. CrossRef Medline
- Chugani DC, Muzik O, Behen M, Rothermel R, Janisse JJ, Lee J, Chugani HT (1999) Developmental changes in brain serotonin synthesis capacity in autistic and nonautistic children. Ann Neurol 45:287–295. CrossRef Medline
- Collins A (2007) Linkage disequilibrium and association mapping: analysis and applications. Totowa, NJ: Humana.
- Cook EH (1990) Autism: review of neurochemical investigation. Synapse 6:292–308. CrossRef Medline
- Cook EH Jr, Courchesne R, Lord C, Cox NJ, Yan S, Lincoln A, Haas R, Courchesne E, Leventhal BL (1997) Evidence of linkage between the serotonin transporter and autistic disorder. Mol Psychiatry 2:247–250. CrossRef Medline
- Coon H, Dunn D, Lainhart J, Miller J, Hamil C, Battaglia A, Tancredi R, Leppert MF, Weiss R, McMahon W (2005) Possible association between autism and variants in the brain-expressed tryptophan hydroxylase gene (TPH2). Am J Med Genet B Neuropsychiatr Genet 135B:42–46. CrossRef Medline
- Deneris ES, Wyler SC (2012) Serotonergic transcriptional networks and potential importance to mental health. Nat Neurosci 15:519–527. CrossRef Medline
- Dennis G Jr, Sherman BT, Hosack DA, Yang J, Gao W, Lane HC, Lempicki RA (2003) DAVID: Database for Annotation, Visualization, and Integrated Discovery. Genome Biol 4:P3. CrossRef Medline
- Dougherty JD, Schmidt EF, Nakajima M, Heintz N (2010) Analytical approaches to RNA profiling data for the identification of genes enriched in specific cells. Nucleic Acids Res 38:4218–4230. CrossRef Medline
- Doyle JP, Dougherty JD, Heiman M, Schmidt EF, Stevens TR, Ma G, Bupp S, Shrestha P, Shah RD, Doughty ML, Gong S, Greengard P, Heintz N (2008) Application of a translational profiling approach for the comparative analysis of CNS cell types. Cell 135:749–762. CrossRef Medline
- Dubin R, Jing Q, O'Broin P, Calder B, McLellan A, Moskowitz D, Suzuki M, Greally JM (2010) WASP: Wiki-based Automated Sequence Processor for Epigenomics and Genomics Applications. J Biomol Techniques 21: S11.
- Eddy SR (2004) Where did the BLOSUM62 alignment score matrix come from? Nat Biotechnol 22:1035–1036. CrossRef Medline
- Fish EW, Sekinda M, Ferrari PF, Dirks A, Miczek KA (2000) Distress vocalizations in maternally separated mouse pups: modulation via 5-HT(1A),

5-HT(1B) and GABA(A) receptors. Psychopharmacology (Berl) 149: 277–285. CrossRef Medline

- Fombonne E (2005) Epidemiology of autistic disorder and other pervasive developmental disorders. J Clin Psychiatry 66(suppl 10):3–8. Medline
- Franklin KBJ, Paxinos G (1997) The mouse brain in stereotaxic coordinates. San Diego: Academic.
- Frazier TW, Youngstrom EA, Speer L, Embacher R, Law P, Constantino J, Findling RL, Hardan AY, Eng C (2012) Validation of proposed DSM-5 criteria for autism spectrum disorder. J Am Acad Child Adolesc Psychiatry 51:28–40.e3. CrossRef Medline
- Freitag CM (2007) The genetics of autistic disorders and its clinical relevance: a review of the literature. Mol Psychiatry 12:2–22. CrossRef Medline
- Gallitano-Mendel A, Wozniak DF, Pehek EA, Milbrandt J (2008) Mice lacking the immediate early gene Egr3 respond to the anti-aggressive effects of clozapine yet are relatively resistant to its sedating effects. Neuropsychopharmacology 33:1266–1275. CrossRef Medline
- 1000 Genomes Project Consortium (2010) A map of human genome variation from population-scale sequencing. Nature 467:1061–1073. CrossRef Medline
- Geschwind DH, Sowinski J, Lord C, Iversen P, Shestack J, Jones P, Ducat L, Spence SJ (2001) The autism genetic resource exchange: a resource for the study of autism and related neuropsychiatric conditions. Am J Hum Genet 69:463–466. CrossRef Medline
- Ghoshal N, Dearborn JT, Wozniak DF, Cairns NJ (2012) Core features of frontotemporal dementia recapitulated in progranulin knockout mice. Neurobiol Dis 45:395–408. CrossRef Medline
- Gong S, Yang XW, Li C, Heintz N (2002) Highly efficient modification of bacterial artificial chromosomes (BACs) using novel shuttle vectors containing the R6Kgamma origin of replication. Genome Res 12:1992–1998. CrossRef Medline
- Good PJ, Chen Q, Warner SJ, Herring DC (2000) A family of human RNAbinding proteins related to the Drosophila Bruno translational regulator. J Biol Chem 275:28583–28592. CrossRef Medline
- Gordon CT, State RC, Nelson JE, Hamburger SD, Rapoport JL (1993) A double-blind comparison of clomipramine, desipramine, and placebo in the treatment of autistic disorder. Arch Gen Psychiatry 50:441–447. CrossRef Medline
- Goridis C, Rohrer H (2002) Specification of catecholaminergic and serotonergic neurons. Nat Rev Neurosci 3:531–541. CrossRef Medline
- Grady RM, Wozniak DF, Ohlemiller KK, Sanes JR (2006) Cerebellar synaptic defects and abnormal motor behavior in mice lacking alpha- and betadystrobrevin. J Neurosci 26:2841–2851. CrossRef Medline
- Han J, Cooper TA (2005) Identification of CELF splicing activation and repression domains in vivo. Nucleic Acids Res 33:2769–2780. CrossRef Medline
- Happé F, Ronald A (2008) The 'fractionable autism triad': a review of evidence from behavioural, genetic, cognitive and neural research. Neuropsychol Rev 18:287–304. CrossRef Medline
- Hartman RE, Wozniak DF, Nardi A, Olney JW, Sartorius L, Holtzman DM (2001) Behavioral phenotyping of GFAP-apoE3 and -apoE4 transgenic mice: apoE4 mice show profound working memory impairments in the absence of Alzheimer's-like neuropathology. Exp Neurol 170:326–344. CrossRef Medline
- Heiman M, Schaefer A, Gong S, Peterson JD, Day M, Ramsey KE, Suárez-Farinas M, Schwarz C, Stephan DA, Surmeier DJ, Greengard P, Heintz N (2008) A translational profiling approach for the molecular characterization of CNS cell types. Cell 135:738–748. CrossRef Medline
- Hillarp NA, Fuxe K, Dahlström A (1966) Demonstration and mapping of central neurons containing dopamine, noradrenaline, and 5-hydroxytryptamine and their reactions to psychopharmaca. Pharmacol Rev 18: 727–741. Medline
- Hofer MA, Shair HN, Brunelli SA (2002) Ultrasonic vocalizations in rat and mouse pups. Curr Protoc Neurosci Chapter 8:Unit 8.14. CrossRef Medline
- Hollander E, Phillips A, Chaplin W, Zagursky K, Novotny S, Wasserman S, Iyengar R (2005) A placebo controlled crossover trial of liquid fluoxetine on repetitive behaviors in childhood and adolescent autism. Neuropsychopharmacology 30:582–589. CrossRef Medline
- Hollander E, Soorya L, Chaplin W, Anagnostou E, Taylor BP, Ferretti CJ, Wasserman S, Swanson E, Settipani C (2012) A double-blind placebocontrolled trial of fluoxetine for repetitive behaviors and global severity in

adult autism spectrum disorders. Am J Psychiatry 169:292–299 CrossRef Medline

- Holy TE, Guo Z (2005) Ultrasonic songs of male mice. PLoS Biol 3:e386. CrossRef Medline
- Iafrate AJ, Feuk L, Rivera MN, Listewnik ML, Donahoe PK, Qi Y, Scherer SW, Lee C (2004) Detection of large-scale variation in the human genome. Nat Genet 36:949–951. CrossRef Medline
- Ishimura K, Takeuchi Y, Fujiwara K, Tominaga M, Yoshioka H, Sawada T (1988) Quantitative analysis of the distribution of serotoninimmunoreactive cell bodies in the mouse brain. Neurosci Lett 91:265– 270. CrossRef Medline
- Kessler RC, McGonagle KA, Swartz M, Blazer DG, Nelson CB (1993) Sex and depression in the National Comorbidity Survey: I. Lifetime prevalence, chronicity and recurrence. J Affect Disord 29:85–96. CrossRef Medline
- Korte SM, Meijer OC, de Kloet ER, Buwalda B, Keijser J, Sluyter F, van Oortmerssen G, Bohus B (1996) Enhanced 5-HT1A receptor expression in forebrain regions of aggressive house mice. Brain Res 736:338–343. CrossRef Medline
- Ladd AN, Nguyen NH, Malhotra K, Cooper TA (2004) CELF6, a member of the CELF family of RNA-binding proteins, regulates muscle-specific splicing enhancer-dependent alternative splicing. J Biol Chem 279: 17756–17764. CrossRef Medline
- Lein ES, Hawrylycz MJ, Ao N, Ayres M, Bensinger A, Bernard A, Boe AF, Boguski MS, Brockway KS, Byrnes EJ, Chen L, Chen L, Chen TM, Chin MC, Chong J, Crook BE, Czaplinska A, Dang CN, Datta S, Dee NR, et al (2007) Genome-wide atlas of gene expression in the adult mouse brain. Nature 445:168–176. CrossRef Medline
- Little S (2001) Amplification-refractory mutation system (ARMS) analysis of point mutations. In: Current Protocols in Human Genetics. New York: Wiley.
- López-Rubalcava C, Hen R, Cruz SL (2000) Anxiolytic-like actions of toluene in the burying behavior and plus-maze tests: differences in sensitivity between 5-HT(1B) knockout and wild-type mice. Behav Brain Res 115: 85–94. CrossRef Medline
- Luo R, Sanders SJ, Tian Y, Voineagu I, Huang N, Chu SH, Klei L, Cai C, Ou J, Lowe JK, Hurles ME, Devlin B, State MW, Geschwind DH (2012) Genome-wide transcriptome profiling reveals the functional impact of rare de novo and recurrent CNVs in autism spectrum disorders. Am J Hum Genet 91:38–55. CrossRef Medline
- McClellan J, King MC (2010) Genetic heterogeneity in human disease. Cell 141:210–217. CrossRef Medline
- McDougle CJ, Naylor ST, Cohen DJ, Aghajanian GK, Heninger GR, Price LH (1996a) Effects of tryptophan depletion in drug-free adults with autistic disorder. Arch Gen Psychiatry 53:993–1000. CrossRef Medline
- McDougle CJ, Naylor ST, Cohen DJ, Volkmar FR, Heninger GR, Price LH (1996b) A double-blind, placebo-controlled study of fluvoxamine in adults with autistic disorder. Arch Gen Psychiatry 53:1001–1008. CrossRef Medline
- McDougle CJ, Epperson CN, Price LH, Gelernter J (1998) Evidence for linkage disequilibrium between serotonin transporter protein gene (SLC6A4) and obsessive compulsive disorder. Mol Psychiatry 3:270–273. CrossRef Medline
- McNamara IM, Borella AW, Bialowas LA, Whitaker-Azmitia PM (2008) Further studies in the developmental hyperserotonemia model (DHS) of autism: social, behavioral and peptide changes. Brain Res 1189:203–214. CrossRef Medline
- Moldin SO, Rubenstein JLR (2006) Understanding autism: from basic neuroscience to treatment. Boca Raton, FL: CRC/Taylor and Frances.
- Montgomery KT, Lartchouck O, Li L, Perera A, Yassin Y, Tamburino A, Loomis S, Kucherlapati R (2008) Mutation detection using automated fluorescence-based sequencing. In: Current protocols in human genetics. New York: Wiley.
- Moy SS, Nadler JJ, Perez A, Barbaro RP, Johns JM, Magnuson TR, Piven J, Crawley JN (2004) Sociability and preference for social novelty in five inbred strains: an approach to assess autistic-like behavior in mice. Genes Brain Behav 3:287–302. CrossRef Medline
- Moy SS, Nadler JJ, Poe MD, Nonneman RJ, Young NB, Koller BH, Crawley JN, Duncan GE, Bodfish JW (2008) Development of a mouse test for repetitive, restricted behaviors: relevance to autism. Behav Brain Res 188: 178–194. CrossRef Medline

Naert A, Callaerts-Vegh Z, D'Hooge R (2011) Nocturnal hyperactivity, in-

creased social novelty preference and delayed extinction of fear responses in post-weaning socially isolated mice. Brain Res Bull 85:354–362. CrossRef Medline

- Nakatani J, Tamada K, Hatanaka F, Ise S, Ohta H, Inoue K, Tomonaga S, Watanabe Y, Chung YJ, Banerjee R, Iwamoto K, Kato T, Okazawa M, Yamauchi K, Tanda K, Takao K, Miyakawa T, Bradley A, Takumi T (2009) Abnormal behavior in a chromosome-engineered mouse model for human 15q11–13 duplication seen in autism. Cell 137:1235–1246. CrossRef Medline
- Neale BM, Kou Y, Liu L, Ma'ayan A, Samocha KE, Sabo A, Lin CF, Stevens C, Wang LS, Makarov V, Polak P, Yoon S, Maguire J, Crawford EL, Campbell NG, Geller ET, Valladares O, Schafer C, Liu H, Zhao T, et al (2012) Patterns and rates of exonic de novo mutations in autism spectrum disorders. Nature 485:242–245. CrossRef Medline
- Ng PC, Henikoff S (2002) Accounting for human polymorphisms predicted to affect protein function. Genome Res 12:436–446. CrossRef Medline
- Olivier B, Molewijk HE, van der Heyden JA, van Oorschot R, Ronken E, Mos J, Miczek KA (1998) Ultrasonic vocalizations in rat pups: effects of serotonergic ligands. Neurosci Biobehav Rev 23:215–227. CrossRef Medline
- O'Roak BJ, Vives L, Girirajan S, Karakoc E, Krumm N, Coe BP, Levy R, Ko A, Lee C, Smith JD, Turner EH, Stanaway IB, Vernot B, Malig M, Baker C, Reilly B, Akey JM, Borenstein E, Rieder MJ, Nickerson DA, et al (2012) Sporadic autism exomes reveal a highly interconnected protein network of de novo mutations. Nature 485:246–250. CrossRef Medline
- Page DT, Kuti OJ, Sur M (2009) Computerized assessment of social approach behavior in mouse. Front Behav Neurosci 3:48. CrossRef Medline
- Perry W, Minassian A, Lopez B, Maron L, Lincoln A (2007) Sensorimotor gating deficits in adults with autism. Biol Psychiatry 61:482–486. CrossRef Medline
- Pfaar H, von Holst A, Vogt Weisenhorn DM, Brodski C, Guimera J, Wurst W (2002) mPet-1, a mouse ETS-domain transcription factor, is expressed in central serotonergic neurons. Dev Genes Evol 212:43–46. CrossRef Medline
- Pinto D, Pagnamenta AT, Klei L, Anney R, Merico D, Regan R, Conroy J, Magalhaes TR, Correia C, Abrahams BS, Almeida J, Bacchelli E, Bader GD, Bailey AJ, Baird G, Battaglia A, Berney T, Bolshakova N, Bölte S, Bolton PF, et al (2010) Functional impact of global rare copy number variation in autism spectrum disorders. Nature 466:368–372. CrossRef Medline
- Pobbe RL, Pearson BL, Defensor EB, Bolivar VJ, Young WS 3rd, Lee HJ, Blanchard DC, Blanchard RJ (2012) Oxytocin receptor knockout mice display deficits in the expression of autism-related behaviors. Horm Behav 61:436–444. CrossRef Medline
- Purcell S, Neale B, Todd-Brown K, Thomas L, Ferreira MA, Bender D, Maller J, Sklar P, de Bakker PI, Daly MJ, Sham PC (2007) PLINK: a tool set for whole-genome association and population-based linkage analyses. Am J Hum Genet 81:559–575. CrossRef Medline
- Ramon y Cajal S, Pasik P, Pasik T (1899) Texture of the nervous system of man and the vertebrates. New York: Springer.
- Robinson EB, Koenen KC, McCormick MC, Munir K, Hallett V, Happé F, Plomin R, Ronald A (2012) A multivariate twin study of autistic traits in 12-year-olds: testing the fractionable autism triad hypothesis. Behav Genet 42:245–255. CrossRef Medline
- Sanders SJ, Murtha MT, Gupta AR, Murdoch JD, Raubeson MJ, Willsey AJ, Ercan-Sencicek AG, DiLullo NM, Parikshak NN, Stein JL, Walker MF, Ober GT, Teran NA, Song Y, El-Fishawy P, Murtha RC, Choi M, Overton JD, Bjornson RD, Carriero NJ, et al (2012) De novo mutations revealed by whole-exome sequencing are strongly associated with autism. Nature 485:237–241. CrossRef Medline
- Scattoni ML, Gandhy SU, Ricceri L, Crawley JN (2008) Unusual repertoire of vocalizations in the BTBR T+tf/J mouse model of autism. PloS One 3:e3067. CrossRef Medline
- Schaefer ML, Wong ST, Wozniak DF, Muglia LM, Liauw JA, Zhuo M, Nardi A, Hartman RE, Vogt SK, Luedke CE, Storm DR, Muglia LJ (2000) Altered stress-induced anxiety in adenylyl cyclase type VIII-deficient mice. J Neurosci 20:4809–4820. Medline
- Schain RJ, Freedman DX (1961) Studies on 5-hydroxyindole metabolism in autistic and other mentally retarded children. J Pediatr 58:315–320. CrossRef Medline
- Silverman JL, Turner SM, Barkan CL, Tolu SS, Saxena R, Hung AY, Sheng M,

Crawley JN (2011) Sociability and motor functions in Shank1 mutant mice. Brain Res 1380:120–137. CrossRef Medline

- Spielman RS, Ewens WJ (1996) The TDT and other family-based tests for linkage disequilibrium and association. Am J Hum Genet 59:983–989. Medline
- Steinbusch HW (1981) Distribution of serotonin-immunoreactivity in the central nervous system of the rat-cell bodies and terminals. Neuroscience 6:557–618. CrossRef Medline
- Steinbusch HW (1984) Serotonin-immunoreactive neurons and their projections in the CNS. In: Handbook of chemical neuroanatomy, Vol 3: Classical transmitters and transmitter receptors in the CNS, Part II (Björklund A, Hökfelt T, Kuhar MJ, eds), pp 1,4,6–16,18,20. Amsterdam: Elsevier.
- Stone JL, Merriman B, Cantor RM, Yonan AL, Gilliam TC, Geschwind DH, Nelson SF (2004) Evidence for sex-specific risk alleles in autism spectrum disorder. Am J Hum Genet 75:1117–1123. CrossRef Medline
- Sunyaev S, Ramensky V, Koch I, Lathe W 3rd, Kondrashov AS, Bork P (2001) Prediction of deleterious human alleles. Hum Mol Genet 10:591–597. CrossRef Medline
- Takahashi S, Kanai H, Miyamoto Y (1976) Reassessment of elevated serotonin levels in blood platelets in early infantile autism. J Autism Child Schizophr 6:317–326. CrossRef Medline
- Underwood JG, Boutz PL, Dougherty JD, Stoilov P, Black DL (2005) Homologues of the Caenorhabditis elegans Fox-1 protein are neuronal splicing regulators in mammals. Mol Cell Biol 25:10005–10016. CrossRef Medline
- Veenstra-VanderWeele J, Anderson GM, Cook EH Jr (2000) Pharmacogenetics and the serotonin system: initial studies and future directions. Eur J Pharmacol 410:165–181. CrossRef Medline
- Veenstra-VanderWeele J, Muller CL, Iwamoto H, Sauer JE, Owens WA, Shah CR, Cohen J, Mannangatti P, Jessen T, Thompson BJ, Ye R, Kerr TM, Carneiro AM, Crawley JN, Sanders-Bush E, McMahon DG, Ramamoorthy S, Daws LC, Sutcliffe JS, Blakely RD (2012) Autism gene variant causes hyperserotonemia, serotonin receptor hypersensitivity, social impairment and repetitive behavior. Proc Natl Acad Sci U S A 109:5469–5474. CrossRef Medline

- Verkerk AJ, Pieretti M, Sutcliffe JS, Fu YH, Kuhl DP, Pizzuti A, Reiner O, Richards S, Victoria MF, Zhang FP, et al (1991) Identification of a gene (FMR-1) containing a CGG repeat coincident with a breakpoint cluster region exhibiting length variation in fragile X syndrome. Cell 65:905–914. CrossRef Medline
- Wang K, Zhang H, Ma D, Bucan M, Glessner JT, Abrahams BS, Salyakina D, Imielinski M, Bradfield JP, Sleiman PM, Kim CE, Hou C, Frackelton E, Chiavacci R, Takahashi N, Sakurai T, Rappaport E, Lajonchere CM, Munson J, Estes A, et al (2009) Common genetic variants on 5p14.1 associate with autism spectrum disorders. Nature 459: 528–533. CrossRef Medline
- Wang P, Ding F, Chiang H, Thompson RC, Watson SJ, Meng F (2002) ProbeMatchDB: a web database for finding equivalent probes across microarray platforms and species. Bioinformatics 18:488–489. CrossRef Medline
- Weiss LA, Arking DE, Daly MJ, Chakravarti A (2009) A genome-wide linkage and association scan reveals novel loci for autism. Nature 461:802– 808. CrossRef Medline
- Whitaker-Azmitia PM (2001) Serotonin and brain development: role in human developmental diseases. Brain Res Bull 56:479–485. CrossRef Medline
- Wilming LG, Gilbert JG, Howe K, Trevanion S, Hubbard T, Harrow JL (2008) The vertebrate genome annotation (Vega) database. Nucleic Acids Res 36:D753–D760. CrossRef Medline
- Wozniak DF, Hartman RE, Boyle MP, Vogt SK, Brooks AR, Tenkova T, Young C, Olney JW, Muglia LJ (2004) Apoptotic neurodegeneration induced by ethanol in neonatal mice is associated with profound learning/ memory deficits in juveniles followed by progressive functional recovery in adults. Neurobiol Dis 17:403–414. CrossRef Medline
- Wylie CJ, Hendricks TJ, Zhang B, Wang L, Lu P, Leahy P, Fox S, Maeno H, Deneris ES (2010) Distinct transcriptomes define rostral and caudal serotonin neurons. J Neurosci 30:670–684. CrossRef Medline
- Yuhas J, Cordeiro L, Tassone F, Ballinger E, Schneider A, Long JM, Ornitz EM, Hessl D (2011) Brief report: sensorimotor gating in idiopathic autism and autism associated with fragile X syndrome. J Autism Dev Disord 41:248–253. CrossRef Medline