Brain development is a dramatic process that unfolds throughout the first decades of life, gradually transforming the brain, and involving both microscopic and macroscopic changes. By far the greatest developmental changes occur by the early twenties, and frontal brain regions are among the last to fully mature; even so, many developmental processes, such as myelination, continue throughout life, only to be overtaken by degenerative changes in old age.

In the course of development, the brain undergoes a remarkable process of restructuring as it adapts to the environment and becomes more efficient in processing information. A variety of brain imaging methods can be used to probe how anatomy, connectivity, and function change in the developing brain. Here we review recent discoveries regarding these brain changes in both typically developing individuals and individuals with neurodevelopmental disorders. We begin with typical development, summarizing research on changes in regional brain volume and tissue density, cortical thickness, white matter integrity, and functional connectivity. Space limits preclude the coverage of all neurodevelopmental disorders; instead, we cover a representative selection of studies examining neural correlates of autism, attention deficit/hyperactivity disorder, Fragile X, 22q11.2 deletion syndrome, Williams syndrome, Down syndrome, and Turner syndrome. Where possible, we focus on studies that identify an age by diagnosis interaction, suggesting an altered developmental trajectory. The studies we review generally cover the developmental period from infancy to early adulthood. Great progress has been made over the last 20 years in mapping how the brain matures with MR technology. With ever-improving technology, we expect this progress to accelerate, offering a deeper understanding of brain development, and more effective interventions for neurodevelopmental disorders.
**Clinical research**

Using postmortem examinations of tissue, the age at which synaptic density peaked for a range of cortical areas was investigated by tracking changes in synaptic density at different ages. Among the last regions to mature are those responsible for higher-level cognition, which is still developing in adolescents (reviewed in ref 3). Some neuropsychiatric disorders emerge in childhood or adolescence and distinctly alter the developmental trajectory for both brain structure and function. By studying characteristic patterns of abnormalities in these disorders, many clues emerge about biological mechanisms contributing to a range of psychiatric illnesses and neurodevelopmental disorders. A more mechanistic understanding of each disorder is crucial—for more effective diagnosis, to better design interventions, and better understand treatment effects. With constantly improving technology, we can now visualize neural structures, axonal pathways, and functional connections with ever-increasing precision.

Here we review recent neuroimaging research in the fields of typical and atypical development, focusing primarily on studies from age 4 to early adulthood. There are now many studies of infancy and even fetal development with magnetic resonance imaging (MRI), but the vast majority of pediatric MRI studies evaluate children old enough to keep still for the duration of a scan, making later ages somewhat easier to study. We cover structural imaging methods, such as voxel-based and tensor-based morphometry (VBM and TBM), structural connectivity analyses, using diffusion tensor imaging and high angular resolution diffusion imaging (DTI and HARDI), and functional connectivity analyses, using graph theory, independent components analysis (ICA), and seed-based methods. We describe these approaches in the **Typical Development** section. We will not cover functional development, as many studies are task-specific and would require much more space to review. In addition to covering development of healthy individuals, we review the neuroimaging literature on a number of neurodevelopmental disorders (Table II), including autism, attention deficit-hyperactivity disorder (ADHD), Fragile X, 22q11.2 deletion syndrome, Williams syndrome, Down syndrome, and Turner syndrome. Where possible, we selected studies that examined the interaction of age and diagnosis, but in some cases we discuss studies simply addressing the effects of a disorder on the brain, as fewer studies have mapped disease effects on the entire developmental trajectory. A few other recent reviews focus on the development of brain structure, functional connectivity, or structural connectivity either in typically developing or atypically developing individuals. In this review, we address each of these topics, but readers are encouraged to refer to these reviews, in addition to the articles we cite here.

**Typical development**

An exhaustive review of all studies of typical development with various neuroimaging methods is beyond the scope of this paper, so we will highlight illustrative examples that reflect some general trends in the field (Table I).

**Structural MRI**

A vast number of methods have been used to investigate changes in brain structure. The most traditional way to measure anatomical changes in the brain is to identify the substructures of brain—often by manual tracing, or more recently by using automated computer programs to measure their volumes. By parcellating the brain into regions with different functions, such as the major lobes, the subcortical nuclei, and cortical regions, several early landmark studies generated “growth curves,” or norms, to show how the size of different brain regions increases—or decreases—with age. Around the year 2000, the first studies were published describing data from large cohorts of children scanned with MRI. Distinct and characteristic growth trajectories were found for each brain region, with some notable sex differences. A more detailed picture of the developmental trajectory emerged with the advent of voxel-based brain mapping methods. Voxel-based statistical approaches can create a color-coded 3D map of growth rates at each location in the brain, or changes in cortical thickness. Alternatively, they can simply test if the density or thick-
ness of a tissue in a brain region is affected by age, or if it relates to a clinical variable of interest. One such statistical mapping method, VBM, begins by spatially normalizing all MRI scans from a group of subjects into the same space. The scans are then segmented into gray matter, white matter, and cerebrospinal fluid (CSF), maps of each tissue are smoothed, and statistical tests are performed at each voxel—or 3D location—in the brain, to determine where age related changes occur, and what other factors affect the brain measures. The values in VBM analyses then represent the average proportion of gray matter in a small region around any given voxel. To be clear, “density” here is not intended to measure axonal or cellular packing density, but it offers a gross anatomical measure of regional tissue volumes, offering spatial detail on the pattern of tissue in the brain.

TBM is a more computationally intensive method, in which a deformation field is obtained for each subject, warping their brain to match a common brain template, and encoding the relative positions of various brain landmarks. Tensor fields, or Jacobian matrices, are then calculated from the gradient of the deformation field, at each point, representing the volume of the unit-cube after the deformation. From these, it is possible to determine the degree of regional volume expansion or shrinkage across scans taken at different times, or to

### Table I.

| Study                          | N     | Ages            | Modality          |
|-------------------------------|-------|-----------------|-------------------|
| **Structural MRI**            |       |                 |                   |
| Giedd et al, 1999             | 145 (56 f) | 4-22 overall | Brain volume      |
| Gogtay et al, 2004†           | 13 (7 F)     | 4-21 yo overall | Brain volume      |
| Sowell et al, 2002            | 35 (15 F)     | 7-16 yo        | Brain volume      |
| Sowell et al, 2003            | 176 (86 F)    | 7-87 yo        | Brain volume (GMD) |
| Sowell et al, 2004            | 45 (22 F)     | 5-10 yo baseline | Cortical thickness |
| Hua et al, 2009†              | 13 (6 F)      | 6-15 yo baseline | Brain volume      |
| Tannes et al, 2010            | 168 (87 F)    | 8-30 yo        | Brain volume, cortical thickness, and DTI |
| **Diffusion-weighted imaging**|       |                 |                   |
| Klingberg et al, 1999         | 7 children, 5 adults, all male | 8-12 and 20-31 yo | DTI - ROI         |
| Schmithorst et al, 2002       | 33 (17 F)     | 5-18 yo        | Voxel-wise DTI    |
| Barnea-Goraly et al, 2005     | 34 (16 F)     | 6-19 yo        | Voxel-wise DTI    |
| Bonekamp et al, 2007          | 40 (18 F)     | 5-19 yo        | DTI - ROI         |
| Eluvathingal et al, 2007      | 31 (16 F)     | 6-17 yo        | DTI - ROI (tract-based) |
| Qiu et al, 2008               | 1: 24 (11 F), 2: 27 (11 F), 3: 24 (13 F) | 1:6-8, 2:9-12, 3:18-26 yo | Voxel-wise DTI |
| Lebel et al, 2008†            | 202 (98 F)    | 5-30 yo        | DTI - ROI (tract based) |
| Kochunov et al, 2010          | 831 (485 F)   | 11-90 yo       | DTI - ROI (tract-based) |
| Hagmann et al, 2010           | 30 (17 F)     | 18 mo-18 yo    | DTI - graph theory |
| Dennis et al, 2013†           | 439 (267 F)   | 12, 16, 20-30 yo | HARDI - graph theory |
| Dennis et al, 2013b           | 438 (266 F)   | 12, 16, 20-30 yo | HARDI - graph theory |
| **Functional connectivity**   |       |                 |                   |
| Fair et al, 2007, 2008†       | 210 (*)       | 7-9, 10-15, 19-31 yo | Seed-based func. conn. |
| Kelly et al, 2009             | 1: 14 (4 F), 2: 12 (5 F), 3: 14 (5 F) | 1:8-13, 2:14-17, 3:20-24 yo | Seed-based func. conn. |
| Dosenbach et al, 2010         | 238 (115 F)   | 7-30 yo        | Seed-based func. conn. |
| Supek et al, 2010             | 1: 23 (13 F), 2: 22 (11 F) | 1: 7-9, 2: 19-22 yo | DTI, ICA func. conn. |
| Thomason et al, 2011          | 65 (32 F)     | 9-15 yo baseline | Seed/ICA func. conn. |
| Jolles et al, 2011            | 1: 19 (10 F), 2: 29 (16 F) | 1: 11-13, 2: 19-25 yo | ICA func. conn. |

Table I. Studies investigating typical development that are reviewed in this paper. Bold indicates study that examined age*diagnosis effect. *, no gender information; AD, autism; ADHD, attention deficit/hyperactivity disorder; PBD, pediatric bipolar disorder; FX, fragile X; DD, developmental delay; 22q, 22q11.2 deletion syndrome; WS, Williams syndrome; DS, Down syndrome; TS, Turner syndrome; TD, typically developing; (# F), number of female participants; yo, years old; mo, months old; MRI, magnetic resonance imaging; GMD, gray matter density; DTI, diffusion tensor imaging; ROI, region of interest; HARDI, high angular resolution diffusion imaging; ICA, independent components analysis; † indicates study for which we have included a figure.
### Table II. Studies investigating neurodevelopmental disorders that are reviewed in this paper. **Bold** indicates study that examined age*diagnosis effect. AD, autism disorder; TD, typically developing; DTI, diffusion tensor imaging; FX, Fragile X; DD, developmental delay; ADHD, attention deficit-hyperactivity disorder; WS, Williams syndrome; ICA, independent components analysis; * = no gender information; (# F), number of female participants. ‡ indicates study for which we have included a figure; # indicates study listed in multiple places.

| Study                          | N              | Ages          | Modality                  |
|-------------------------------|----------------|---------------|---------------------------|
| **Autism**                    |                |               |                           |
| Langen et al, 2009            | 99 AD (8 F), 89 TD (7 F) | 6-25 yo       | Brain volume              |
| Hardan et al, 2009            | 18 AD, 16 TD, all male | 8-12 yo baseline | Brain volume, cortical thickness |
| Brun et al, 2009              | 24 AD, 26 TD, all male | 6-16 yo       | Brain volume              |
| Mengotti et al, 2011          | 20 AD (2 F), 22 TD (2 F) | 4-14 yo       | Brain volume, DTI         |
| Hua et al, 2013               | 13 AD, 7 TD, all male | 6-13 yo baseline | Brain volume              |
| Barnea-Goraly et al, 2004     | 7 AD, 9 TD, all male | AD mean=14.6, TD mean=13.4 | DTI                      |
| Bashat et al, 2007            | 7 AD, 41 TD (23 F) | 1.8-3.3 AD, 4 mo-23 yo TD | DTI                      |
| Keller et al, 2007            | 34 AD, 31 TD, all male | 10-35 yo     | DTI                      |
| Barnea-Goraly et al, 2010     | 13 AD (2 F), 13 same-sex siblings, 11 TD (2 F) | 6-13 yo | DTI                      |
| Shukla et al, 2010‡           | 26 AD (1 F), 24 TD (1 F) | 9-20 yo       | DTI                      |
| Cherkesky et al, 2006         | 57 AD (4 F), 57 TD (5 F) | AD mean=24, TD mean=24 | Seed-based func. conn.  |
| Just et al, 2007              | 18 AD (1 F), 18 TD (3 F) | AD mean=27.1, TD mean=24.5 | Seed-based func. conn.  |
| Kennedy et al, 2008           | 12 AD, 12 TD, all male | 15-52 yo     | Seed-based func. conn.   |
| Monk et al, 2009              | 12 AD (1 F), 12 TD (2 F) | AD mean=26, TD mean=27 | Seed-based func. conn.   |
| Noonan et al, 2009            | 10 AD, 10 TD, all male | 14-43 yo     | Seed-based func. conn.   |
| Weng et al, 2010              | 16 AD (2 F), 15 TD (1 F) | 13-18 yo     | Seed-based func. conn.   |
| Assaf et al, 2010             | 15 AD (1 F), 15 TD (2 F) | 10-23 yo     | ICA func. conn.          |
| Rudie et al, 2012‡            | 23 AD (2 F), 25 TD (3 F) | 8-17 yo      | Seed-based func. conn.   |
| Rudie et al, 2013             | 42 AD (6 F), 37 TD (6 F) | 9-18 yo      | Graph theory func. conn. |
| **Attention-deficit hyperactivity disorder** | | | |
| Castellanos et al, 2002       | 152 ADHD (63 F), 139 TD (56 F) | 4-19 yo | Brain volume |
| Sovell et al, 2003            | 27 ADHD (11 F), 46 TD (17 F) | 8-18 yo | Brain volume |
| Carmona et al, 2005           | 25 ADHD (4 F), 25 TD (4 F) | 6-16 yo | Brain volume |
| Shaw et al, 2006              | 163 ADHD (*), 166 TD (*) | Mean baseline=8 yo | Cortical thickness |
| Shaw et al, 2007              | 223 ADHD (82 F), 223 TD (82 F) | Mean baseline=10 yo | Cortical thickness |
| Brieber et al, 2007           | 15 AD, 15 ADHD, 15 TD | ADHD mean=13.1, TD mean=13.3 | Brain volume |
| Luders et al, 2009            | 19 ADHD, 19 TD, all male | 7-16 yo | Callosal thickness |
| Kobel et al, 2010             | 14 ADHD, 12 TD, all male | 9-13 yo | Brain volume, DTI, ICA func. conn. |
| Ashhari et al, 2005           | 18 ADHD (6 F), 15 ADHD (6 F) | 7-11 yo | DTI |
| Hamilton et al, 2008          | 17 ADHD, 16 TD, all male | ADHD mean=12.0, TD mean=11.7 | DTI |
| Pavuluri et al, 2009          | 13 PBD (3 F), 13 ADHD (1 F), 13 TD (7 F) | ADHD mean=13.4, TD mean=13.7 | DTI |
| Silk et al, 2009a,b           | 15 ADHD, 15 TD, all male | 8-18 yo | DTI |
| Tamm et al, 2012‡             | 12 ADHD, 12 TD, all male | 14-18 yo | DTI |
| Lawrence et al, 2013          | 56 ADHD (17 F), 31 siblings (24 F), 17 TD (11 F) | 6-18 yo | DTI |
| Cao et al, 2006               | 29 ADHD, 27 TD, all male | 11-16.5 yo | Regional homogeneity (func. conn.) |
| Wang et al, 2009              | 19 ADHD, 20 TD, all male | ADHD mean=13.6, TD mean=13.3 | graph theory func. conn. |
| Fair et al, 2010              | 23 ADHD (7 F), 23 TD (11 F) | 7-16 yo | Seed-based func. conn. |
| Qiu et al, 2011               | 15 ADHD, 15 TD, all male | 10-15 yo | Brain volume, cortical thickness, DTI, ICA func. conn. |
| Study | N | Ages | Modality |
|-------|---|------|----------|
| **Fragile X** | | | |
| Lee et al, 2007 | 36 FX (18 F), 33 TD (17 F) | FX mean=14.7, TD mean=14.7 | Brain volume |
| Hoeft et al, 2010 | 41 FX, 28 TD, all male | 1-3 yo baseline | Brain volume |
| Hoeft et al, 2011† | 52 FX, 31 TD, all male | 1-4 yo | Brain volume |
| Hazlett et al, 2012 | 53 FX, 50 TD, all male | 18-42 mo baseline | Brain volume |
| Peng et al, 2013 | 48 FX (*), 28 DD (*), 36 TD (*) | 15-27 yo | Brain volume |
| Barnea-Goraly et al, 2003a | 10 FX, 10 TD, all female | 11-23 yo | DTI |
| Haas et al, 2009 | 17 FX, 13 TD, 8 DD, all male | 1.5-4 yo | DTI |
| Villalon et al, 2013# | 18 FX, 25 22q, 17 TS, 41 TD, all female | 7-14 yo | DTI |
| **22q11.2 Deletion syndrome** | | | |
| Simon et al, 2005 | 18 22q (11 F), 18 TD (7 F) | 7-14 yo | Brain volume |
| Campbell et al, 2006 | 39 22q (19 F), 26 siblings (10 F) | 22q mean=11, TD mean=11 | Brain volume |
| Bearden et al, 2007 | 21 22q (11 F), 13 TD (6 F) | 8-17 yo | Cortical thickness |
| Bearden et al, 2009 | 21 22q (11 F), 13 TD (6 F) | 8-17 yo | Cortical thickness |
| Schaar et al, 2009† | 59 22q (35 F), 80 TD (44 F) | 6-40 yo baseline | Cortical thickness |
| Shashi et al, 2010 | 22 22q (7 F), 16 TD (6 F) | 9-17 yo | Brain volume |
| Gothelf et al, 2011 | 19 22q (8 F), 18 TD (8 F) | 22q mean=13, TD mean=13.4 baseline | Brain volume |
| Srivastava et al, 2012 | 49 22q, 37 TD* | 6-15 yo | Cortical gyrification |
| Barnea-Goraly et al, 2003b | 19 22q (6 F), 19 TD (6 F) | 7-22 yo | DTI |
| Villalon et al, 2013# | 18 FX, 25 22q, 17 TS, 41 TD, all female | 7-14 yo | DTI |
| Debbane et al, 2012 | 27 22q (15 F), 33 TD (14 F) | 12-19 yo | ICA func. conn. |
| **Williams syndrome** | | | |
| Thompson et al, 2005 | 42 WS (23 F), 40 TD (24 F) | 12-50 yo | Cortical thickness |
| Eckert et al, 2006 | 42 WS (23 F), 40 TD (24 F) | 12-50 yo | Cortical morphology |
| Gaser et al, 2006 | 42 WS (23 F), 40 TD (24 F) | 12-50 yo | Cortical morphology |
| Boddaren et al, 2006 | 9 WS (2 F), 11 TD (5 F) | 5-15 yo | Brain volume |
| Tosun et al, 2006 | 39 WS (22 F), 39 TD (23 F) | WS mean=29.9, TD mean=27.1 | Cortical morphology |
| Chiang et al, 2007 | 41 WS (23 F), 39 TD (23 F) | 12-50 yo | Brain volume |
| Luders et al, 2007 | 12 WS (5 F), 12 TD (5 F) | 13-30 yo | Callosal thickness |
| Meda et al, 2012 | 31 WS (11 F), 50 TD (23 F) | WS mean=26.3, TD mean=28.0 | Brain volume, cortical thickness |
| Hoeft et al, 2007 | 10 WS (4 F), 10 DD (7 F), 10 TD (3 F) | WS mean=26.8, TD mean=27.8 | DTI |
| Arlinghaus et al, 2011† | 16 WS (6 F), 16 TD (8 F) | 16-33 yo | DTI |
| Jabbi et al, 2012 | 14 WS (7 F), 23 TD (10 F) | WS mean=27.8, TD mean=32.1 | Brain volume, DTI |
| Haas et al, 2012 | 20 WS (9 F), 10 DD (7 F), 10 TD (2 F) | WS mean=23.7, TD mean=27.8 | DTI |
| Haas et al, 2012 | 39 WS (24 F), 40 TD (23 F) | WS mean=26.1, TD mean=21.3 | Brain volume |
| **Down syndrome** | | | |
| Pinter et al, 2001 | 16 DS (5 F), 15 TD (*) | DS mean=11.3, TD mean=11.9 | Brain volume |
| Smigielska-Kuzia et al, 2011 | 23 DS (9 F), 26 TD (11 F) | 3-15 yo | Brain volume |
| **Turner syndrome** | | | |
| Murphy et al, 1993 | 18 TS, 19 TD, all female | TS mean=30, TD mean=27 | Brain volume |
| Reiss et al, 1995 | 30 TS, 30 TD, all female | 6-17 yo | Brain volume |
| Brown et al, 2002 | 26 TS, 26 TD, all female | TS mean=13.2, TD mean=13.4 | Brain volume |
| Kesler et al, 2004 | 30 TS, 29 TD, all female | 6-33 yo | Brain volume |
| Molko et al, 2004 | 14 TS, 14 TS, all female | 18-36 yo | DTI |
| Holzapfel et al, 2006 | 10 TS, 10 TD, all female | 7-24 yo | Brain volume, DTI |
| Yamagata et al, 2012† | 26 TS, 20 TD, all female | 3-12 yo | Brain volume, DTI |

Table II. Continued
determine anatomical differences in a set of scans. These can then be analyzed statistically, to identify characteristic diagnostic group differences, age effects, or links with clinical or cognitive measures. Cortical thickness assessments use semiautomated methods to reconstruct 3D representations of the gray matter–white matter boundary and the pial surface, and they calculate the distance between the two for every point across the cortex.\textsuperscript{22,23} Using cortical thickness maps, time-lapse movies have been created to show the shifting pattern of cortical thinning in typically developing children between ages 4 and 21,\textsuperscript{1} and in disorders such as childhood-onset schizophrenia,\textsuperscript{24,25} or before and after the onset of bipolar disorder.\textsuperscript{26}

Between age 5 and adulthood, important changes occur in higher cognitive functions that in part reflect changes in brain structure. Total brain volume increases with age, and many studies have found that the growth rate varies across the brain, and over time. Gray matter volume increases into adolescence, when it plateaus and begins to decline, but white matter volume usually increases into adulthood.\textsuperscript{23,27}

Myelination continues throughout life—even well into old age—and white matter volume reductions only begin to be observed when the balance between myelin production and degeneration tilts in favor of white matter loss.\textsuperscript{28} If this is recognized, it avoids the misconception that some of the developmental processes “stop,” when in fact they continue but are often dominated by other more dramatic changes.

The age at which gray matter volume plateaus varies across the lobes, and temporal gray matter volume tends to reach a maximum last.\textsuperscript{13} Within the lobes too, there is a great deal of variation in time to mature. In a whole-brain study, it was found that the prefrontal cortex and the posterior part of the superior temporal gyrus were shown to be the last to mature (Figure 1).\textsuperscript{1} In general, phylogenetically earlier structures—those supporting vision, hearing, and sensorimotor function—develop the most rapidly in infancy. To some extent, “ontogeny recapitulates phylogeny.” Brain areas that support speech, language comprehension, and finally executive function, tend to develop in roughly the same sequence as they emerged during human evolution. Sowell et al similarly found that the posterior temporal cortex had a more protracted development.\textsuperscript{29} For subcortical structures, they showed that as the brain grows in size, the proportion taken up by subcortical structures decreases, but at a different rate for males and females.\textsuperscript{13} Additionally, they proposed that the decrease in gray matter, while due in part to cortical pruning (ie, synapse elimination and dendritic pruning), was also due in large part to the ongoing increase in white matter. They also examined cortical thickness between ages 5 and 11.\textsuperscript{20} While large areas of cortex became thinner with age, cortical gray matter in Broca’s and Wernicke’s areas thickened.

Hua et al used TBM to show regional brain changes in a longitudinal dataset from children, finding expansion of cerebral white matter and shrinkage of parietal, temporal, and occipital gray matter (Supplementary Figure 1—Supplementary figures are available in the online version of this article).\textsuperscript{31} Using TBM, one can create a picture of the mean growth rate, for each brain region, at any age. Tamnes et al examined age-related changes in a large cohort of subjects between ages 8 and 30 with both struc-

| Study            | N            | Ages  | Modality          |
|------------------|--------------|-------|-------------------|
| Villalon et al, 2013# | 18 FX, 25 22q, 17 TS, 41 TD, all female | 7-14 yo | DTI               |
| Bray et al, 2011  | 37 TS, 18 TD, all female          | 7-13 yo | Seed-based func. conn. |
| Bray et al, 2012  | 15 TS, 14 TD, all female          | 7-14 yo | Seed-based func. conn. |

Table II. Continued
structural MRI (sMRI) and diffusion tensor imaging (DTI—described below). They found prominent cortical thinning across the parietal lobe, superior medial frontal lobe, cingulate gyrus, prefrontal cortex, and occipital cortex. The rate of thinning was greatest in the youngest subjects, after which the rate slowed down.

**Diffusion-weighted imaging**

Diffusion-weighted imaging (DWI) is a variant of MRI scanning, which allows us to visualize, at a gross level, the rate of diffusion of water along axons. It can therefore be used to visualize axonal pathways. The MR signal is reduced when water is diffusing, and it is possible to design an MRI protocol whose signals are depleted by water diffusing in a particular direction (a diffusion gradient image). By measuring diffusion in a large set of different directions (at least 6, but often as many as 30 to 256 directions), we can identify the primary directions of water diffusion in each voxel in the brain.

Diffusion tensor imaging (DTI) models water diffusion at each voxel as an ellipsoid or “tensor,” after which tractography may be used to follow and reconstruct the major white matter fiber bundles. HARDI is similar to

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**Supplementary Figure 1.** Gray matter maturation between ages 7-15. Tissue growth maps modeled by linear regression, for all subjects and males and females separately.

Reproduced from ref 31: Hua X, Leow AD, Levitt JG, Caplan R, Thompson PM, Toga AW. Detecting brain growth patterns in normal children using tensor-based morphometry. *Hum Brain Mapp.* 2009;30:209–219. Copyright © Wiley-Liss 2009.
DTI, but can map crossing fibers better, as it does not rely on the assumption that there is only one dominant fiber present in each voxel. HARDI collects diffusion information from more angles and uses orientation distribution functions (ODFs), or other spherical functions—instead of tensors—to map the probability of water diffusion in every direction, leading to more accurate tractography. Fractional anisotropy (FA), the degree to which water diffuses in one direction (along the axon), is one of the most widely used measures of axonal integrity. As a rule of thumb—which has many exceptions—higher FA and lower mean diffusivity (MD) tend to reflect more highly developed, more strongly myelinated tracts, with a higher axonal conduction speed. These measures are reproducible in children, providing reliable developmental biomarkers.

Figure 2. White matter maturation between ages 5 and 30. Age-related fractional anisotropy increases measured by tractography in 202 individuals across 11 tracts. Reproduced from ref 48: Lebel C, Walker L, Leemans A, Phillips L, Beaulieu C. Microstructural maturation of the human brain from childhood to adulthood. *NeuroImage*. 2008;40:1044-1055. Copyright © Elsevier 2008
Specifically examining the frontal lobe white matter, Klingberg et al found significantly greater white matter fractional anisotropy in adults than in children.\(^{42}\) They attributed this to a lesser degree of myelination in children; this is also consistent with visual inspection of brain MRI scans from infants, which often show limited white-matter contrast in poorly myelinated regions. Schmithorst et al expanded on earlier work, examining a range of specific tracts in subjects between 5 and 18 years old.\(^{43}\) FA increased with age in the internal capsule, corticospinal tract, left arcuate fasciculus, and right inferior longitudinal fasciculus. Similar trajectories have been reported in DTI studies of the entire lifespan.\(^{44}\) In one study, FA increased with age in the internal capsule, the white matter of the prefrontal cortex, corpus callosum, basal ganglia and thalamic pathways, and visual pathways.\(^{45}\) Several of these regions underlie cognitive functions such as memory and attention, as well as motor skills. Eluvathingal et al examined 6 specific tracts and found three patterns in the results.\(^{46}\) Various parts of the arcuate fasciculus, inferior longitudinal fasciculus, inferior fronto-occipital fasciculus, uncinate fasciculus, and corticospinal tract showed either increased in FA with decreases in other measures of diffusivity, or no detectable effect on FA and decreases in diffusivity. Only the somatosensory pathway showed no detectable age effect at all, probably because it matures very early in infancy, prior to the age range examined in that study. In a voxel-wise analysis, Qiu et al found widespread age effects on FA across the cerebellum, temporal, frontal, and parietal lobes.\(^{47}\) Additionally, they found that reading scores (in Chinese and English) were associated with higher FA in a number of regions. Lebel et al found that the developmental trajectory of measures of anisotropy and diffusivity across most tracts were best fit with an exponential curve (Figure 2).\(^{48}\) Echoing structural studies above, they found the last tracts to mature were frontotemporal connections. In one of the largest brain imaging studies to date, Kochunov et al detailed how 11 major tracts change over the lifespan (age 11 to 90) in 831 subjects.\(^{49}\) By charting the FA of these tracts across their subject pool, they reported the “age-at-peak” for each tract, as well as the rate of increase/decrease, along with sex differences, in some cases. Using DTI-based connectivity analysis, Hagmann et al used graph theory to show that the efficiency of the brain’s anatomical network increased with age—as did the number of detectable connections for each brain region.\(^{50}\) Graph theory represents the brain as a set of nodes (brain regions) and edges (the connections between them). A number of standard parameters such as path length and modularity, to name a few, are used to describe network topology.\(^{51}\) Characteristic path length measures the average path length in a network. It does not refer to the physical length of the tracts, but the number of edges, or individual “jumps,” between nodes in the network. Modularity is the degree to which a system may be subdivided into smaller networks. Graph theory can quantify more global features in brain connectivity patterns. These include network efficiency, or the degree to which the network is differentiated into modules. Using cortical connectivity matrices calculated from HARDI data, Dennis et al examined the developmental trajectory of graph theoretical measures of structural connectivity (Figure 3).\(^{52}\) Path length and modularity, among

![Figure 3. Development of structural connectivity between age 12 and age 30. Still images from videos available online from ref 55 displaying the increases and decreases in degree and fiber density between age 12 and age 30. For this image, node size is proportional to the degree (number of connections), and connection thickness is proportional to relative fiber density. The connection color is simulated to make the connections easier to see. The rate of increase or decrease for each node and connection was the regression coefficients from the age analyses for those nodes and connections. Small blue dots indicate nodes for which there was no significant age-related increase or decrease in degree. Only connections that had a significant age-related increase or decrease in fiber density are included in this image, other connections exist but are not drawn in for clarity. In this image are both weighted (fiber density) and binary (degree) measures. These images are created from the results when analyses were restricted to only connections existing in at least 95% of subjects. Reproduced from ref 52. Dennis EL, Jahanshad N, McMahon KL, et al. Development of brain structural connectivity between ages 12 and 30: a 4-Tesla diffusion imaging study in 439 adolescents and adults. NeuroImage. 2013;64:671–684. Copyright © Elsevier 2013](image-url)
other measures, decreased with age, suggesting an increase in network integration. Interestingly, the left and right intrahemispheric networks, when analyzed separately, showed opposing age trends; some parameters increased with age in the left hemisphere, but decreased in the right. If this is corroborated in the future, it could point to different developmental processes in each hemisphere, perhaps due to the known structural asymmetry of the brain, which also increases with age.\textsuperscript{15,53} The level of structural brain asymmetry is also known to relates to the morphometry of corpus callosum, the major interhemispheric commissure,\textsuperscript{54} so it may also relate to detectable differences in connectivity. Dennis et al also found differences in the structural core of the brain, as the “rich club” is restructured and strengthened.\textsuperscript{55} The “rich club” of the brain is the core of the network, made up of high-degree nodes that are highly interconnected and play an important role in network efficiency.\textsuperscript{56}

**Functional connectivity**

Resting-state fMRI (rsfMRI) is a branch of research based on the theory that distributed brain regions are functionally coupled, even if they are not directly structurally connected. In fact, the coherence (temporal correlations) in brain activity across disparate brain regions may be used to identify systems or networks in the brain that interact. Resting-state functional connectivity can be assessed through blood oxygenation level dependent (BOLD) time-courses of these distant regions, resulting in a number of intrinsic connectivity networks (ICNs) that are reliably found across individuals, and across studies. The main methods to assess functional connectivity are independent components analysis (ICA), seed-based analysis, and graph theory. ICA is a model-free approach, in which the four-dimensional resting-state data (the time-series) is decomposed into time courses and associated spatial maps, describing the temporal and spatial characteristics of the components making up the data.\textsuperscript{57} Seed-based analysis is a model-based approach in which the researcher selects a seed region of interest, and extracts the time course of that seed. They then correlate that time course with the time-course of activations in the rest of the brain, searching for those that are most similar.\textsuperscript{58} Regions whose time course is highly correlated with the seed are considered to be functionally coupled. Lastly, graph theory can also be applied to functional images, exactly as discussed in the previous section. Graph theory is applicable to functional, anatomical, or diffusion-weighted MRI—any scans that measure the relationship between brain regions in terms of correlation, coherence, mutual information, or physical measures of connectivity such as fiber density.

Focusing on regions involved in task control, Fair et al found that the period of development between 7 and 31 was marked by increases in segregation and integration, as distinct networks mature.\textsuperscript{59} In the same dataset, they examined the maturation of the default mode network, and it was found to be only sparsely connected in children (Figure 4).\textsuperscript{60} The default mode network (DMN) is a network usually including the precuneus/posterior cingulate (PCC), medial prefrontal cortex (mPFC), hippocampus, inferior parietal lobule, and lateral temporal cortex.\textsuperscript{61} These regions are more active during rest than during a task, hence the name “default mode” or “task negative” network.\textsuperscript{62} Using five seeds in distinct regions of the anterior cingulate cortex it was found that over development, local patterns of connectivity evolved from diffuse to focused, and networks changed from exhibiting mostly local connectivity to include more distant brain regions.\textsuperscript{63} Subjects’ resting state data were able to be used to predict their age—their maturational curve accounted for more than half of the variation in their data.\textsuperscript{64} Examining both structural and functional connectivity of DMN regions, it was found that the connectivity of the PCC-mPFC along the cingulum was the least mature in children.\textsuperscript{65} Some regions that were poorly connected structurally in children still had strong functional connectivity. This suggests that the saying “what fires together, wires together”\textsuperscript{66} may hold on a larger scale—the functional coupling of some brain regions may strengthen their structural connectivity over time.

In a cohort of subjects scanned multiple times—both within scan session and between sessions separated by a few years—it was demonstrated that rsfMRI can reliably map brain networks in children and adolescents.\textsuperscript{67} A study that focused less on the specific regions connected and more on the quality of the connections found that children’s functional networks tended to include more voxels and than did those of adults.\textsuperscript{68} This supports earlier hypotheses that maturation is marked by a process of refining and “focusing” of brain networks.
**Neurodevelopmental disorders**

While we cannot cover all neurodevelopmental disorders, here we review some of the more common or more commonly studied neurodevelopmental disorders (*Table II*).

**Autism**

Autism is a neurodevelopmental disorder characterized by deficits in social interaction and communication, and by repetitive behaviors. The prevalence of autism is estimated to be around 2.5%\(^69\) and is usually diagnosed by age 3.\(^70\) Autism has a partially genetic basis, although the specific mechanisms that contribute to the disorder are complex and are not expected to be the same in all children with autism.\(^71\)

**Structural MRI**

A number of studies have compared individuals of a specific age group with autism with typically developing individuals; fewer have examined changes in the developmental trajectory associated with autism. In an impressively large study (N=188), Langen et al examined...
the development of the striatum in autistic and typically developing individuals. They found opposite developmental trends in individuals with autism in the caudate and the nucleus accumbens, and an altered trajectory, albeit in the same direction, in the putamen. Hardan et al examined longitudinal changes in cortical thickness in autistic boys, finding a greater decrease with age in cortical thickness in the autistic individuals than the typically developing boys. Brun et al found that autistic boys had enlarged lobes compared with typically developing boys, but voxel-wise analyses also showed gray matter deficits in parietal, temporal, and occipital lobes. Mengotti et al examined changes in the developmental trajectory of both regional brain volume and structural connectivity in individuals with autism and found that the volume of the inferior temporal cortex, superior and inferior parietal lobule, and superior occipital lobe were larger in individuals with autism, while the volumes of the inferior frontal cortex and supplementary motor cortex were smaller. Hua et al examined longitudinal data, and the trajectory of white matter growth was slowed in autistic boys, especially in the parietal lobe. In gray matter, they found accelerated growth in the anterior cingulate cortex and putamen.

Diffusion-weighted imaging

Diffusion imaging studies of autism show widespread disruption of white matter tracts, especially between regions implicated in social behavior (Figure 5). According to one theory of autism, at least a subset of children with autism experience an initial brain “overgrowth,” after which typically developing children catch up and surpass autistic children. This is a debated hypothesis in the field, however, and it may apply to some autistic children but not others. Various findings support this. Significantly accelerated maturation of the white matter has been found in autistic children. Following this overgrowth, the autistic brain may fail to effectively prune connections, leading to disorganization. One region has been found to show an interaction of age with diagnostic group: the right posterior limb of the internal capsule decreased in FA with age in typically developing individuals, but it increased with age in individuals with autism. It has also been found that the apparent diffusion coefficient (ADC) was negatively associated with age across most of the cortex and the splenium of the corpus callosum in autistic individuals, but no detectable associations with age in typically developing individuals were found.

Functional connectivity

We were unable to find any reports of an age by diagnosis interaction effect on functional connectivity in autism. A number of studies have reported effects of autism diagnosis on intrinsic connectivity networks.
(ICNs). One set of studies supports the theory that autism results from widespread underconnectivity in the brain. Monk et al found a correlation between social functioning and the degree of connectivity between the posterior cingulate and the superior frontal gyrus; those with poorer skills exhibited weaker connectivity. They also found stronger functional connectivity in autistic subjects between the posterior cingulate and the temporal lobe and parahippocampal gyrus. One study reported generally more extensive functional connectivity in autistic individuals, challenging the underconnectivity hypothesis, but this study had fewer subjects than any of the others. Addressing the question in a different way, some have found both reduced integration within network and reduced segregation between networks in individuals with autism (Figure 6).91,92

Figure 6. Functional connectivity in autism. Bilateral amygdala connectivity. (A) The Harvard-Oxford bilateral amygdala (25% probability) used as seed region and displayed on the 1mm MNI152 T1 standard brain. (B) Typically developing (TD) within-group connectivity maps, (C) Autism spectrum disorder (ASD) group connectivity maps, and (D) direct between-group contrasts rendered on the Inflated PALS B12 brain using CARET (Computerized Anatomical Reconstruction and Editing Toolkit) and on the 1mm Montreal Neurological Institute (MNI)152 T1 standard brain using Analysis of Functional NeuroImages. Maps are thresholded at Z > 2.3 (P < 0.01) with correction for multiple comparisons applied at the cluster level (P < 0.05). Red circles highlight areas of greater positive connectivity with the seed region for the TD group. Blue circles highlight areas of greater negative connectivity with the seed region for the TD group. The original paper also details the connectivity of the right inferior frontal gyrus, pars opercularis.

Adapted from ref 91: Rudie JD, Shehzad Z, Hernandez LM, et al. Reduced functional integration and segregation of distributed neural systems underlying social and emotional information processing in autism spectrum disorders. Cereb Cortex. 2012;22:1025–1037. Copyright © Oxford University Press 2012.
Attention deficit/hyperactivity disorder

Attention deficit/hyperactivity disorder (ADHD) is a common neurodevelopmental disorder, disproportionately diagnosed in boys. It is a heterogeneous disorder with a strong familial factor.

Structural MRI

There are many studies reporting brain structural differences in individuals with ADHD, but they do not appear to have arrived at a consensus. Widespread differences in gray matter volume are reported, while other
reports discuss more specific regions of differences in volume, focusing on the parietal and temporal areas of the brain, respectively.95-98 Some studies have found a thinner corpus callosum,99 and widespread differences in cortical thickness.100 In a longitudinal study, Shaw et al reported thinner cortex in children with ADHD, across a number of areas important for attentional control.101 Additionally, while mean cortical thickness showed consistent differences across development, there was an age by diagnosis effect in the right parietal cortex. A similar longitudinal study by these authors found that the “age at peak” for cortical thickness was significantly delayed in children with ADHD, especially in the prefrontal cortex.102

Diffusion-weighted imaging

A number of studies have found disruption in white matter tracts implicated in the pathophysiology of ADHD.103-106 Silk et al examined subjects with ADHD between ages 8 and 18.107 The mean FA of the whole brain for both groups increased with age, with localized increases in FA in the ADHD subjects. Further analysis suggested that these increases were in fact due to decreased neural branching in subjects with ADHD. Looking specifically at the FA of the basal ganglia in the same subject pool, they also found differences in the developmental trajectory of the caudate.108 Similarly, Tamm et al also found greater FA and axial diffusivity in ADHD subjects specifically in frontostriatal pathways, perhaps due to decreased branching (Supplementary Figure 2—Supplementary figures are available in the online version of this article).109 Lawrence et al found that white matter disruptions in individuals with ADHD were also found to some degree in their siblings, suggesting a strong familial factor.110

Functional connectivity

A few studies have found that the functional connectivity within the DMN (default mode network) is disrupted or decreased in ADHD.108,111 Along with increases in the regional homogeneity in the occipital cortex, decreases in the regional homogeneity of the frontostriatal-cerebellar circuits were found in boys with ADHD.112 This fits with some current hypotheses regarding the pathophysiology of ADHD. Using graph theory, decreased global efficiency and increased local efficiency in ADHD were found, pointing to a shift from the typical “small-world” networks towards less biological “regular” networks.113 Small-world networks have a balance of network integration and segregation and are most efficient, while a regular or lattice network is highly segregated, a topology that is rarely found in functioning biological networks.

Neurogenetic disorders

Fragile X syndrome

Fragile X (FX) is caused by an expansion of the CGG repeat in the 5’ untranslated region of the fragile X mental retardation 1 (FMR1) gene, leading to a loss or decrease in functionality of fragile X mental retardation protein (FMRP). It is a common genetic cause of intellectual disability,114 especially in boys.

Structural MRI

In a longitudinal study, Hoeft et al found altered developmental trajectories in the gray matter volume of the orbital gyri, basal forebrain, and thalamus in young boys with FX, along with a number of differences that persisted across development.115 Differences in the white matter volume of the frontostriatal regions became more pronounced with age. Also using a longitudinal design, Hazlett et al found generalized brain overgrowth in boys with FX, especially in the temporal lobe, cerebellum, and caudate.116 Looking at a main effect of diagnosis, Lee et al found volumetric increases in the caudate and ventricles—abnormalities that correlated with the degree of reduction in the FMRP protein in females.117 Comparing boys with FX with those with AD, idiopathic developmental delay, and typically developing boys, Hoeft et al found widespread reductions in frontal and temporal gray and white matter in young boys with FX (Figure 7).118

Diffusion-weighted imaging

Studies of white matter integrity in fragile X report abnormalities mainly in the frontostriatal pathways. Decreased FA in females with FX, and increased fiber density in males with FX, have both been found in frontostriatal regions.119,120
**Functional connectivity**

We were unable to find any studies of functional connectivity in Fragile X.

**22q11.2 Deletion syndrome**

22q11.2 Deletion syndrome (22q DS), also called velocardiofacial syndrome and DiGeorge syndrome (among other names), is caused by a deletion on chromosome 22 and results in a heterogeneous spectrum of physiological, neurological, and psychological symptoms. Several of the 30 genes encoded in the deleted segment are highly expressed in the developing brain and known to affect early neuronal migration. Several neuroimaging studies have pointed to abnormal patterns of cortical thinning and white matter impairments.

**Structural MRI**

In a cross-sectional study, Schaer et al found altered developmental trajectories of cortical thickness in 22q DS, with a decreased rate of thinning in childhood followed by an increased rate of cortical thinning in late adolescence (Supplementary Figure 3—Supplementary figures are available in the online version of this article). This study built on earlier work by Bearden et al suggesting regionally specific cortical thinning in 22q DS, in superior parietal cortices and right parieto-occipital cortex, regions critical for visuospacial processing, and bilaterally in the most inferior portion of the inferior frontal gyrus (pars orbitalis), a key area for language development. A later study of the same cohort also used fractal dimension analysis to reveal altered complexity and gyrification in 22q DS, a further index of disturbed cor-

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**Figure 7.** Differences in regional brain volume in fragile X. A: Regions showing significant differences in regional gray matter (GM) volume and white matter (WM) volume between fragile X syndrome (FXS) and idiopathic autism (iAUT) (panel A), FXS and typically developing (TD) and idiopathic developmentally delayed (DD) controls (panel B), and iAUT and TD/DD controls (panel C). The left side shows the right hemisphere. The statistical threshold is set at \( P=0.01 \), familywise error cluster-level corrected. Montreal Neurological Institute coordinates. Adapted from ref 118: Hoeft F, Walter E, Lightbody A, et al. Neuroanatomical differences in toddler boys with fragile X syndrome and idiopathic autism. Arch Gen Psychiatry. 2011;68:295–305. Copyright © American Medical Association 2011

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tical development. Studies of volumetric changes in the gray matter in 22q DS have found reductions in the cerebellum and posterior areas of the posterior and occipital areas and expansions in the frontal lobes, although one group found reductions in the frontal lobe as well.\textsuperscript{127} Looking specifically at which individuals with 22q DS developed psychosis, Gothelf et al found that greater reduction of the left DLPFC predicted greater psychotic symptoms. 22q DS is a heterogeneous disorder, which predisposes individuals to a range of other psychiatric and neurological issues.\textsuperscript{128} This heterogeneity might explain some of the lack of agreement across studies. In the white matter, volume is reduced in individuals with 22q DS, across the cerebellum, internal capsule, and frontal cortex.\textsuperscript{125,126} Srivastava et al found abnormalities in the development of the cortical gyri in children with 22q DS, specifically in areas important for visuospatial, attentional, and numerical cognition tasks.\textsuperscript{129}

**Diffusion-weighted imaging**

Widespread changes in FA in 22q DS, including decreased FA in the corpus callosum, and increased FA in the cingulate and parietal lobe,\textsuperscript{125} and widespread effects, with individuals with 22q DS (velocardiofacial syndrome in the paper) having decreased FA across the frontal, parietal, and temporal white matter\textsuperscript{130} have also been found. In a more recent study, Villalon et al found that girls with 22q11.2DS showed lower fractional anisotropy (FA) than controls in the association fibers of the superior and inferior longitudinal fasciculi, the splenium of the corpus callosum, and the corticospinal tract.\textsuperscript{131}

**Functional connectivity**

Only one study, to our knowledge, has examined how individuals with 22q DS differ in functional connectiv-
Debbané et al found widespread changes in functional connectivity using ICA to compare networks between groups. Individuals with 22q DS had altered connectivity in the default mode network, with both increased connectivity between lateral frontal regions and the inferior parietal lobule and decreased connectivity between medial frontal regions and the precuneus. They also found altered connectivity in visual, sensorimotor, and visuospatial networks.

**Williams syndrome**

Williams syndrome (WS) is a disorder caused by a hemizygous deletion of chromosome 7q11.23 resulting in physiological, intellectual, and behavioral abnormalities.

**Structural MRI**

Thompson et al found a pattern of excesses and deficits in cortical thickness in WS, along with alterations in the complexity of the gyral pattern. Using a fractal dimension analysis of the cortical surface, abnormalities of gyral folding were found, consistent with reports of altered sulcal patterns, especially in perisylvian regions, in people with WS. The analysis of cortical patterns using surface-based analyses of local curvature has also revealed gyral-sulcal anomalies in WS. The shape anomalies are also found subcortically at midline, and have been characterized using mesh-based shape analysis methods. Boddart et al found reductions in the gray matter volume of the left parieto-occipital region in children with WS, which overlapped with prior findings in adults with WS. Chiang et al similarly found reductions in the parietal and occipital regions, along with subcortical structures including the basal ganglia and thalamus, and the volumes of these structures were positively correlated with IQ. Meda et al also found the basal ganglia to be significantly reduced in WS. Additionally, individuals with WS had increased cortical thickness and/or decreased surface area in a number of ROIs across parietal, occipital, and frontal regions. Even the smaller subcortical gray matter nuclei, such as the amygdala and caudate nucleus, show shape anomalies in WS, and their implications for cognition and behavior are only just beginning to be understood.

**Diffusion-weighted imaging**

DWI studies of Williams syndrome have shown increases in FA in the superior longitudinal fasciculus and inferior longitudinal fasciculus, association pathways important for language, memory, visuospatial processing, and object processing, to name a few (Figure 8). Reduced FA in the uncinate fasciculus—one of the fiber tracts connecting the limbic system—has also been found. Looking specifically at tracts related to the fusiform gyrus, Haas et al found both an increased volume of fibers and increased FA in individuals with WS. Face processing is altered in WS, and these results may explain these abnormalities.

**Functional connectivity**

We were unable to find any studies examining functional connectivity in WS.

**Chromosomal disorders**

**Down syndrome**

Down syndrome (DS), or trisomy 21, is a common chromosomal disorder and the most common cause of intellectual disability. There are surprisingly few brain imaging studies of DS in children. DS increases the risk of developing Alzheimer’s-like dementia with age, so many more studies focus on adults with DS.

**Structural MRI**

Total brain volume is decreased in DS and certain structures are disproportionately affected. Consistent with adult imaging studies, the hippocampus is reduced in DS, but there is conflicting information as to whether...
the amygdala is as well.\textsuperscript{151,152} Children with DS were found to have reduced frontal and temporal lobe volumes.\textsuperscript{152} The differences in the hippocampus are particularly intriguing given the increased risk for dementia in DS individuals.

**Diffusion-weighted imaging**

We were unable to find any studies of white matter integrity in DS in children.
**Functional connectivity**

To our knowledge, no studies have examined functional connectivity in DS.

**Turner syndrome**

Another chromosomal disorder, Turner syndrome (TS) results from the absence of one X chromosome in girls, resulting in a number of changes physically, hormonally, and neurologically.\(^{153}\)

**Structural MRI**

A number of studies have examined brain volume in TS, generally finding decreased brain volume in the parietal and occipital regions.\(^{154-156}\) The hippocampus and subcortical structures such as the thalamus and basal ganglia are also reduced in TS,\(^ {154,157}\) but the amygdala is larger.\(^ {157}\)

**Diffusion-weighted imaging**

DWI studies in TS reveal abnormalities across a large area of the white matter. Molko et al found microstructural differences in the temporal lobe, especially tracts running anterior-posterior in the temporal lobe.\(^ {158}\) Holzapfel et al found lower FA in the pallidum, internal capsule, and prefrontal cortex, as well as in the parieto-occipital region, extending into the superior longitudinal fasciculus.\(^ {159}\) Yamagata et al found lower FA in a wide array of regions implicated in visuospatial processing, face processing, and sensorimotor and social abilities, including a number of association, commissural, and projection fibers (Supplementary Figure 4—Supplementary figures are available in the online version of this article).\(^ {160}\) In one DTI study comparing TS with Fragile X syndrome and 22q DS, Villalon et al, found that girls with TS had lower FA in the inferior longitudinal fasciculus, right internal capsule and left cerebellar peduncle.\(^ {131}\) Even so, partially overlapping white matter anomalies were detected in all three neurogenetic disorders. They suggested that altered white matter integrity in the superior and inferior longitudinal fasciculi and thalamic to frontal tracts may contribute to the behavioral characteristics of all of these disorders.

**Functional connectivity**

Based on the known deficits girls with TS experience in working memory tasks, one study examined functional connectivity during a working memory task.\(^ {161}\) Reduced connectivity was found between parietal and dorsal frontal regions, which correlated with task performance. A second study examined the specific connectivity of the posterior parietal cortex, finding differential clustering in TS, which may underlie the visuospatial processing deficits in TS.\(^ {162}\)

**Conclusion**

In this paper, we have reviewed representative research over the last 20 years investigating brain development using neuroimaging techniques. We discussed both healthy development and neurodevelopmental disorders, including autism, ADHD, fragile X, 22q DS, Williams syndrome, Down syndrome, and Turner syndrome. The brain undergoes remarkable changes in structure and connectivity as it matures into adulthood. The developmental trajectory of these brain measures is important to identify for our fundamental understanding of the brain and of neurodevelopmental disorders. Disrupted brain structure or connectivity can lead to neurodevelopmental or neuropsychiatric disorders. Understanding these disorders and their developmental trajectory in greater detail should expedite the discovery and more efficient evaluation of effective interventions.

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Desarrollo cerebral clásico y atípico: una revisión de estudios de neuroimágenes

Durante el curso del desarrollo el cerebro experimenta un notable proceso de reestructuración para adaptarse al ambiente y llegar a ser más eficiente en el procesamiento de la información. Se puede emplear una variedad de métodos de imágenes cerebrales para evaluar cómo cambia la anatomía, la conectividad y el funcionamiento durante el desarrollo. Se revisan los descubrimientos recientes en relación con estos cambios cerebrales en sujetos que tienen un desarrollo típico y en quienes tienen trastornos del neurodesarrollo. El artículo comienza con el desarrollo clásico, resumiendo la investigación acerca de los cambios en el volumen regional y la densidad del tejido cerebral, el espesor cortical, la integridad de la sustancia blanca y la conectividad funcional. La limitación de espacio impide cubrir todos los trastornos del desarrollo y se aborda una selección representativa de estudios que examinan los correlatos neurales en autismo, trastorno por déficit de atención/hiperactividad, Síndrome X frágil, Síndrome de delección 22q11.2, Síndrome de Williams, Síndrome de Down y Síndrome de Turner. Cuando es posible se destacan los estudios que identifican una interacción entre la edad y el diagnóstico, lo que sugiere una alteración en el curso del desarrollo. Los estudios revisados en general cubren el periodo de desarrollo entre la infancia y la adultez inicial. En los últimos 20 años, con tecnología de resonancia magnética, se han realizado grandes progresos en el mapeo de cómo madura el cerebro. Se espera que con tecnologías cada vez mejores se acelere este progreso, se posibilite una comprensión más profunda del desarrollo cerebral y se puedan realizar intervenciones más efectivas para los trastornos del neurodesarrollo.

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