Gender Differences in Language and Motor-Related Fibers in a Population of Healthy Preterm Neonates at Term-Equivalent Age: A Diffusion Tensor and Probabilistic Tractography Study

BACKGROUND AND PURPOSE: Sex differences in white matter structure are controversial. In this MR imaging study, we aimed to investigate possible sex differences in language and motor-related tracts in healthy preterm neonates by using DTI and probabilistic tractography.

MATERIALS AND METHODS: Thirty-eight preterm neonates (19 boys and 19 girls, age-matched), healthy at term-equivalent age and at 12 months were included. TBV was measured individually. Probabilistic tractography provided tract volumes, relative tract volumes (volume normalized to TBV). FA, MD, and $\lambda_1$ in the SLF, in the TRs, and in the CSTs. Data were compared by using independent $t$ tests, and Bonferroni corrections were performed to adjust for multiple comparisons.

RESULTS: We showed that healthy preterm boys had larger TBV than girls. However, girls had statistically significantly larger relative tract volumes than boys bilaterally in the parieto-temporal SLF, and in the left CST. Moreover, in the left parieto-temporal SLF, a trend toward lower MD and $\lambda_1$ was observed in females.

CONCLUSIONS: Structural sex differences were found in preterm neonates at term-equivalent age in both sides of the parieto-temporal SLF and in the left CST. Further studies are necessary to investigate whether these structural differences are related to later sex differences in language skills and handedness or to the effect of prematurity.

ABBREVIATIONS: ATR = anterior thalamic radiation; CST = corticospinal tract; DTI = diffusion tensor tractography; FA = fractional anisotropy; GA = gestational age; MD = mean diffusivity; PTR = posterior thalamic radiation; SENSE = sensitivity encoding; SLF = superior longitudinal fasciculus; STR = superior thalamic radiation; TBV = total brain volume; TR = thalamic radiation; $\lambda_P$ = longitudinal diffusivity; $\lambda_L$ = transverse diffusivity

Received February 9, 2011; accepted after revision March 28.

From the Departments of Radiology (Y.L., T.M., J.A., D.B., P.D., F.A.), Neonatology (B.V.O.), and Pediatric Neurology (P.V.B., A.A.) and Laboratoire de Cartographie Fonctionnelle du Cerveau (P.V.B., A.A.), ULB-Hôpital Erasme, Brussels, Belgium; Imaging Diagnosis Center (Y.L.), Shanghai Children’s Medical Center, Shanghai, China; Department of Biostatistics and Medical Computer Science (V.D.De M.), Faculty of Medicine, ULB, Brussels, Belgium; and Philips Healthcare Benelux (V.D.), Brussels, Belgium.

This work was supported by grants from the Fonds Xénophila (ULB) and the Fond de la Recherche Scientifique of Belgium (grant 1.5.149.10).

Please address correspondence to Yan Liu, Department of Radiology, ULB-Hôpital Erasme, 808 Lennik St, 1070 Brussels, Belgium, e-mail: yanliu@ulb.ac.be

Indicates open access to non-subscribers at www.ajnr.org

Indicates article with supplemental on-line table.

http://dx.doi.org/10.3174/ajnr.A2690

AJNR Am J Neuroradiol 32:2011–16 | Dec 2011 | www.ajnr.org 2011

Substantial interest in sex differences in neural structures has been generated in recent years by observations of sex differences in cognitive functions. A male advantage for spatial abilities has been widely observed in humans and other animals, whereas a female advantage has been seen for verbal abilities such as verbal fluency and verbal memory in adult life. This difference also has been found in children, with girls having better language development at an early age and boys experiencing more frequent language impairments.

Therefore, postmortem pathologic and in vivo quantitative brain imaging studies have been looking for differences between males and females. In adults, several studies have shown that men have larger (by ~10%) brains than women. Interestingly, these differences are already present in children and neonates. In adults, the regional volumetric gray matter distribution patterns tend to show an enlargement in females when adjusting for brain size. In children, findings of sex differences in relative gray matter volume have shown enlargement in females, most prominently in the temporal and parietal cortices.

Studies on sex effects on global and regional WM are controversial; both significant and nonsignificant interactions have been reported. It is possible that the measured WM volumes, as determined from conventional MR imaging, reflect changes in macrostructure only, and may not be sensitive to WM microstructure. Such microstructural changes are within the reach of DTI, an MR imaging technique that allows studying the in vivo microstructure and the volume of the major WM tracts. DTI assesses and quantifies water diffusion at a microstructural level, given that water diffuses more easily in the direction of the fibers than orthogonal, diffusion indices, such as FA, MD, and $\lambda_P$ and $\lambda_L$, allow us to indirectly quantify brain microstructure. Results for sex differences in diffusion indices in adults, either global or regional, have been inconsistent. One study showed no sex difference, whereas others showed significant sex differences, but only when focusing on predefined brain regions, such as the frontal lobe or the corpus callosum. Nevertheless, it
should be noted that all these studies by using either ROI analysis or voxel-based morphometric techniques have an error related to anatomic ambiguity in the ROI definition, WM segmentation, and other postprocessing steps such as spatial normalization and smoothing. These methods focus on predefined brain regions but not on specific WM tracts. DTT provides a 3D reconstruction of specific WM tracts and is able to overcome these confounding effects. Moreover, to our knowledge, no diffusion imaging studies have yet investigated whether sex differences are present in neonates.

In this study, we investigated, by using DTT and DTT, whether sex-related differences were present in the language and motor related fibers in healthy preterm neonates at term-equivalent time.

Materials and Methods

Subjects

Among preterm neonates born between June 2005 and June 2009 who underwent brain MR imaging to detect lesions related to premature birth, 78 preterm neonates with acceptable (see below) DTTI were studied. The inclusion criteria for normality were as follows: 1) normal head circumference at birth (>5th and <95th percentiles), 2) 5-minute Apgar score >6, 3) lack of evidence for congenital infection or multiple congenital anomaly syndrome, 4) normal structural brain MR imaging as assessed by 2 board-certified neuroradiologists (D.B., P.D.), and 5) normal physical and neurologic examination at term-equivalent age and at 12 months corrected for GA as assessed by a board-certified neonatologist (A.A.). On the basis of these criteria, 28 neonates were excluded. Furthermore, 12 normal neonates were excluded to obtain sex groups of equal sample size, that were matched for GA at birth and corrected GA at the time of MR imaging. Thirty-eight healthy preterm neonates (19 boys and 19 girls) were finally included in this study (Table). The study was approved by the ethics committee of our institution (reference P2004/207 and P2009/208).

MR Imaging Data Acquisition

MR imaging data were acquired by using a 1.5T magnet (Achieva; Philips, Best, the Netherlands) equipped with an 8-channel SENSE head coil. The following sequences were acquired for all subjects: 1) sagittal 3D T1-weighted gradient-echo images, 2) coronal T2-weighted turbo-spin-echo images, 3) spin-echo echo-planar images (DTI): TR/TE = 5888/92 ms, FOV = 220 × 220 mm², 32 noncolinear diffusion-sensitizing gradient directions with diffusion sensitivity of b = 600 s/mm² and a 2 × 2 mm² in-plane resolution, acceleration factor (SENSE) of 2.2, section thickness = 2.3 mm, and the scanning time for DTI acquisition of 3 minutes 40 seconds.

No sedation was used, and the neonates were spontaneously asleep, positioned in a vacuum immobilization pillow to minimize body and head movements. Ear-muffs were placed to minimize noise exposure. Oxygen saturation and electrocardiography were monitored throughout the acquisition.

Data Postprocessing

Data analysis was performed by using FSL software. Image analysis was performed by using FSL software. Image artifacts due to eddy current distortions and head movements were minimized by registering the DTI from 32 directions to the B0 images. DTI images corresponding to directions with motion artifacts was excluded from further data processing. DTI was considered as acceptable when <5 directions had to be excluded. Extraction of the brain parenchyma from scalp and skull was performed with the FSL Brain Extraction Tool; any small errors identified in the masks were manually corrected. Maps of the diffusion indices were obtained by using FSL Diffusion Toolbox.

Probabilistic Tractography

The bundles were reconstructed in each subject by a single investigator (Y.L.) by using multisetor probabilistic tractography. Seed masks and waypoint masks were generated on color-coded FA maps, placed carefully by one radiologist (Y.L.) and checked by a second radiologist (D.B.). The SLF was separately tracked into 2 parts: the frontoparietal SLF and the parieto-temporal SLF. For the frontoparietal SLF, a seed mask covered the frontal WM and the waypoint mask covered the frontoparietal WM; for the parieto-temporal SLF, the seed mask was the same as the waypoint mask of the frontoparietal SLF, and the waypoint mask covered the temporal lobe. The TRs were studied separately in 4 subradiations; ATR, the motor and sensory STR, and the PTR. A seed mask was positioned in the bottom of the thalamus and a waypoint mask was positioned in the anterior limb of the internal capsule for the ATR; in the precentral gyrus for the motor STR; in the postcentral gyrus for the sensory STR, and in the occipital lobe for the PTRs. The CST was isolated as a whole, by using a seed mask positioned in the cerebral peduncle and a waypoint mask in the precentral gyrus.

The original tracts were normalized by the total number of samples going from the seed mask to the target mask. Finally, the obtained connectivity distributions were thresholded with a probability of 2%.

We calculated the TBV by measuring the volume of the voxels located in the brain mask. To assess tract macrostructure, tract volumes and relative tract volumes (defined as the ratio between individual tract volume and TBV) were computed. The microstructure of the tracts was evaluated with diffusion indices (FA, MD, λ1, and λ2) by using FSL maths.

Statistical Analyses

All variables were analyzed with the SPSS software (SPSS, Chicago, Illinois). A 1-sample Kolmogorov-Smirnov test was performed to detect a possible departure from normality of our variables. Sex-related differences in the TBV, the volumes, the relative volumes, and the diffusion indices (FA, MD, λ1, and λ2) of each tract were analyzed by using a t test for independent samples. Adjustment for multiple comparisons was performed by using the Bonferroni correction.
tical significance was reached when $P < .004$. A trend toward significance was reported when $P < .05$.

Results

Brain Volume
The TBV of the 38 participants ranged from 367 to 614 cm$^3$ (mean ± SD, 438 ± 52 cm$^3$). TBV values in males (461 ± 59 cm$^3$) were 10.7% larger than those in females (414 ± 30 cm$^3$; $P = .004$).

Sex Differences in Principal WM Tracts
WM tracts related to sensorimotor and language functions are shown in Fig 1.

Relative tract volumes were statistically significantly larger in females than in males (Fig 2) bilaterally in the parieto-temporal SLF (left, $P < .001$; right, $P < .001$) and in the left CST ($P < .001$). Moreover, trend toward larger tract volumes (Online Table 1) was found bilaterally in the parieto-temporal SLF (left, $P = .034$; right, $P = .011$).

A trend toward lower MD ($P = .041$) and $\lambda_\perp$ ($P = .033$) in females was observed in the left parieto-temporal SLF (Online Table).

Discussion
In this in vivo brain MR imaging study, we investigated sex differences in the TBV and WM tracts with DTI probabilistic tractography in the language and motor networks in a population of healthy preterm neonates scanned at term-equivalent age. We found, like other studies in neonates$^{19}$ and in adults,$^{14,23}$ that at term-equivalent age healthy preterm male neonates had larger TBV than females. The original findings of our study were that female neonates had larger relative tract volumes bilaterally in the parieto-temporal SLF and in the left CST, with a trend toward lower MD and $\lambda_\perp$ in the left parieto-temporal SLF after Bonferroni correction.

Previous studies have shown that the temporal cortex is larger in females than in males. This has been demonstrated in children by using structural imaging,$^{25,26}$ and also in adults through pathologic studies showing larger planum temporale$^{56}$ and Heschl gyrus,$^{57}$ and a greater attenuation of neurons$^{58}$ in females. Given that the parieto-temporal SLF is supposed to transmit auditory information from the superior temporal gyrus to the inferior parietal lobe, we suggest that our results may reflect an early established difference in favor of female neonates in the number or size of axons in these language-related regions.

The SLF is one of the slowest maturing WM tracts, being not yet myelinated at birth.$^{59,60}$ Lower MD and $\lambda_\perp$ are probably caused by a decrease in brain water content and an increase in membrane attenuation, and they suggest an advanced pre-myelination stage characterized by proliferation and maturation of oligodendrocytes.$^{61,62}$ Therefore, we propose that this
microstructural sex difference might be caused by an advanced maturation in the left parieto-temporal SLF in female neonates.

The finding of different tract relative volumes with no significant difference in diffusion indices is a feature with no straightforward interpretation. In the right parieto-temporal SLF, a larger relative tract volume associated with a trend toward larger tract volume in females was found in the absence of difference in diffusion indices: this might possibly reflect macrostructural changes (more axons at the same myelination stage). In the CST, a larger relative tract volume in females was observed together with no significant difference in either tract volume or diffusion indices, suggesting a similar maturation and number of axons in a smaller female brain. In other published series, differences in tract volumes were not always associated with differences in diffusion indices.33,48,63,64 Moreover, we used probabilistic tractography, which does not directly rely on diffusion index values, but on the uncertainty orientation of the distribution function, enabling it to progress across regions with principal direction uncertainty and through regions with crossing fibers. Therefore, in probabilistic tractography, volume measurement is not directly linked to diffusion indices.51

Language acquisition and processing have shown sex-related differences in infants as young as 2 years old.8,10,65 McCoby and Jacklin66 reported that girls outperformed boys during preschool and early years in articulation, length of sentences, verbal fluency, grammar, and spelling. In giving the California Verbal Learning Test to children between 5 and 16 years old, girls were found to use more semantic clustering, to recall and recognize more items, and to relate words together more as a recall aid than did boys.67 In addition, language impairments have been found to occur more frequently in boys than in girls.11,12 Because of its implication in language function, we suggest that the sex effect on parieto-temporal SLF relative tract volume and microstructure might explain the more rapid development of language skills hitherto reported in females.

Although we could not evidence a significant sex effect on the volume of the CST, we showed that the relative volume of the left CST is larger in females. Interestingly, studies in adults have already shown, after adjusting for the TBV, an increased volume21 and gray matter concentration14 in the precentral gyri in females. A relatively larger left CST in females might explain why meta-analyses showed more right-handed females than males in the general population.68 However, the relationship between handedness and asymmetry in the adult brain is not established, because some studies have found such relationship69,70 but others have not.71

Sex differences in the volume and microstructure of certain WM tracts, as observed in this study, might result from genetic factors as well as from effects of sex steroids on brain development, both factors being known to affect regional tissue composition.72-74

Another hypothesis is that our results may have been influenced by the effect of prematurity. Indeed, even if the normality of our preterm population was based on robust structural and clinical criteria, as in previous studies,48,52 we cannot exclude the possibility that the sex related differences observed in the language and motor networks may have been caused by subtle cerebral lesions, because certain studies seem to suggest that preterm males may be more sensitive to brain injuries
than females. Therefore, it would be of interest to investigate whether these sex differences are also present in healthy term neonates.

Because the first years of life are perhaps the most dynamic phase of postnatal brain development, with rapid development of a wide range of cognitive and motor functions, the link between structural sex differences at term-equivalent age with later functional differences should be interpreted with great caution. Longitudinal studies combining cognitive evaluation with structural and functional imaging may provide insights into the structure-function relationship in sex differences.

Another limitation of our study is that the reproducibility of mask placement was not assessed. Nevertheless, mask placements were checked by 2 radiologists and in probabilistic tractography, by using the approach of normalization, the size of seed and target masks can be ignored.51

Conclusions
In this DTI and probabilistic tractography study on healthy preterm neonates, we demonstrated that sex differences are present in language and motor-related tracts at term-equivalent age. Further studies are needed to investigate whether these structural differences are related to later sex differences in language skills and handedness or to the effect of prematurity.

Acknowledgments
We are grateful to Doni Tamblyn for assistance in language editing.

Disclosures: Vivianne De Maertelaer. Research Support (including provision of equipment or materials): University of Brussels; Vincent Demolin. Consultant: Advice on sequence parameters and data analysis.

References
1. Weissa EM, Kemmler G, Deisenhammer EA, et al. Sex differences in cognitive functions. Pers Individ Dif 2005;38:663–75
2. Kimura D. Sex and Cognition. Cambridge, Massachusetts: MIT Press; 1999
3. Kimura D, Harshman RA. Sex differences in brain organization for verbal and non-verbal functions. Prog Brain Res 1984;61:423–41
4. Jones CM, Brainthwaite V. A., Healy SD. The evolution of sex differences in spatial ability. Behav Neurosci 2003;117:703–11
5. Caplan PJ, Crawford M, Hyde JS, et al. Gender Differences in Human Cognition. New York: Oxford University Press; 1997
6. Halpern DF. Sex Differences in Cognitive Abilities. Hilldale, New Jersey: L. Erlbaum Associates; 1992
7. Hyde JS, Linn MC. Gender differences in verbal ability: a meta analysis. Psych Bull 1988;104:53–69
8. Bornstein MJ, Haynes OM. Vocabulary competence in early childhood: measurement, latent construct, and predictive validity. Child Dev 1998;69:654–71
9. Lung FW, Shu BC, Chiang TL, et al. Predictive validity of Bayley scale in language development of children at 6–36 months. Pediatr Int 2009;51:666–89
10. Reilly S, Wake M, Bavin EL, et al. Predicting language at 2 years of age: a prospective community study. Pediatrics 2007;120:1441–49
11. Law J, Boyle J, Harris F, et al. Prevalence and natural history of primary speech and language delay: findings from a systematic review of the literature. Int J Lang Commun Disord 2006;35:165–88
12. Robinson RJ. Causes and association of severe and persistent specific speech and language disorders in children. Dev Med Child Neurol 1991;33:843–62
13. Leonard C, Towler S, Welcome S, et al. Size matters: cerebral volume influences sex differences in neuroanatomy. Cereb Cortex 2008;18:2920–31
14. Luders E, Narr KL, Thompson PM, et al. Mapping cortical gray matter in the young adult brain: effects of gender. Neuroimage 2005;26:493–501
15. Peters M. Sex differences in human brain size and the general meaning of differences in brain size. Can J Psychol 1991;45:507–22
16. Giedd JN, Castellanos FX, Rajapakse JC, et al. Sexual dimorphology of the developing human brain. Prog Neuropsychopharmacol Biol Psychiatry 1997; 21:1185–201
17. Reis AL, Abrams MT, Singer HS, et al. Brain development, gender and IQ in children: a volumetric imaging study. Brain 1996;11:1763–74
18. Wilke M, Krageloh-Mann I, Holland SK. Global and local development of gray and white matter volume in normal children and adolescents. Exp Brain Res 2007;178:296–307
19. Gilmore JH, Lin W, Prastawa MW, et al. Regional gray matter growth, sexual dimorphism, and cerebral asymmetry in the neonatal brain. J Neurosci 2007;27:1255–60
20. Allen JS, Damasio H, Grabowski TJ, et al. Sexual dimorphism and asymmetries in the gray-white composition of the human cerebrum. Neuroimage 2004;21:4880–94
21. Goldstein JM, Sridman LJ, Horton NJ, et al. Normal sexual dimorphism of the adult human brain assessed in vivo with magnetic resonance imaging. Cereb Cortex 2001;11:940–97
22. Good CD, Johndrow J, Ashburner J, et al. Cerebral asymmetry and the effects of sex and handedness on brain structure: a voxel-based morphometric analysis of 465 normal adult human brains. Neuroimage 2001;14:665–70
23. Gur RC, Turetsky BJ, Matsui M, et al. Sex differences in brain gray and white matter in healthy young adults: correlations with cognitive performance. J Neurosci 1999;19:4053–60
24. Nopoulos P, Flaum M, O’Leary D, et al. Sexual dimorphism in the human brain: evaluation of tissue volume, tissue composition and surface anatomy using magnetic resonance imaging. Psychiatry Res 2000;89:1–13
25. Sowell ER, Peterson B, Kan E, et al. Sex differences in cortical thickness mapped in 176 healthy individuals between 7 and 87 years of age. Cereb Cortex 2007;17:1550–60
26. Sowell ER, Trauner DA, Gamst A, et al. Development of cortical and subcortical brain structures in childhood and adolescence: a structural MRI study. Dev Med Child Neurol 2002;44:4–16
27. Blanton RE, Levitt JG, Peterson IR, et al. Gender differences in the left inferior frontal gyrus in normal children. Neuroimage 2004;22:626–36
28. Hsu JL, Lemans A, Bai CH, et al. Gender differences and age-related white matter changes of the human brain: a diffusion tensor imaging study. Neuroimage 2008;39:566–77
29. De Bellis MD, Keshavan MS, Beers SR, et al. Sex differences in brain maturation during childhood and adolescence. Cereb Cortex 2001;11:552–57
30. Smith CD, Chebrul H, Weckstein D, et al. Age and gender effects on human brain anatomy: a voxel-based morphometric study in healthy elderly. Neurobiol Aging 2007;28:1075–87
31. Behrens TE, Johansen-Berg H, Woolrich MW, et al. Non-invasive mapping of connections between human thalamus and cortex using diffusion imaging. Nat Neurosci 2003;6:750–57
32. Dubois P, Hertz-Pannier L, Dehaene-Lambertz G, et al. Assessment of the early organization and maturation of infants’ cerebral white matter fiber bundles: a feasibility study using quantitative diffusion tensor imaging and tractography. Neuroimage 2006;30:121–32
33. Hüppi PS, Dubois I. Diffusion tensor imaging of brain development. Semin Fetal Neonatal Med 2006;11:489–97
34. Hüppi PS, Maier SE, Peled S, et al. Microstructural development of human newborn cerebral white matter assessed in vivo by diffusion tensor magnetic resonance imaging. Pediatr Res 1998;44:584–90
35. Jellison BJ, Field AS, Medow J, et al. Diffusion tensor imaging of cerebral white matter: a pictorial review of physics, fiber tract anatomy, and tumor imaging patterns. AJNR Am J Neuroradiol 2004;25:356–69
36. Berman JJ, Mukherjee P, Partridge S, et al. Quantitative diffusion tensor MRI of fiber tractography of sensory-motor white matter development in premature infants. Neuroimage 2005;27:862–71
37. Ulag AM, van Zijl PC. Orientation-independent diffusion imaging without tensor diagonalization: anisotropy definitions based on physical attributes of the diffusion ellipsoid. Magn Reson Imaging 1999;8:804–15
38. Sullivan EV, Adalsteinnson E, Hedges M, et al. Equivalent disruption of regional white matter microstructure in ageing healthy men and women. Neuroreport 2001;12:99–104
39. Schmithorst VJ, Holland SK, Dardzinski BJ. Developmental differences in white matter architecture between boys and girls. Hum Brain Mapp 2008;29:696–710
40. Silveri MM, Rehan ML, Pimentel PJ, et al. Sex differences in the relationship between white matter microstructure and impulsivity in adolescents. Magn Reson Imaging 2006;24:833–41
41. Szeszko PR, Vogel J, Ashari M, et al. Sex differences in frontal lobe white matter microstructure: a DTI study. Neuroreport 2003;14:2469–73
42. Westerhausen R, Kreuder F, Dos Santos Sequira S, et al. Effects of handedness and gender on macro- and microstructure of the corpus callosum and its subregions: a combined high-resolution and diffusion tensor MRI study. Cogn Brain Res 2004;21:418–26
43. Wiesthagen R, CD, Josephs O, et al. Optimization of 3-D MP-RAGE sequences for structural brain imaging. Neuroimage 2000;12:112–27
44. Woodward JJ, Anderson PJ, Austin NC, et al. Neonatal MRI to predict neurodevelopmental outcomes in preterm infants. N Engl J Med 2006; 17:727–29
45. Smith SM, Jenkinson M, Woolrich MW, et al. Advances in functional and structural MR image analysis and implementation as FSL. Neuroimage 2004;23 Suppl 1:S208–19
Dubois J, Dehaene-Lambertz G, Perrin M, et al. Thalamo-cortical connectivity in children born preterm: a diffusion tensor imaging and probabilistic tractography study. *Neuroimage* 2007;34:896–904.

Basu I, Risch D, Volzonne A, et al. Probabilistic diffusion tractography of the optic radiations and visual function in preterm infants at term-equivalent age. *Brain* 2008;131:573–82.

Liu Y, Balériaux D, Kavec M, et al. Structural asymmetries in motor and language networks in a population of healthy preterm neonates at term-equivalent age: a diffusion tensor imaging and probabilistic tractography study. *Neuroimage* 2010;51:783–88.

Makris N, Kennedy DJ, McKinney S, et al. Segmentation of subcomponents within the superior longitudinal fascicle in humans: a quantitative, in vivo, DT-MRI study. *Cereb Cortex* 2005;15:854–69.

Wakana S, Caprithan A, Panzenbock MM, et al. Reproducibility of quantitative tractography methods applied to cerebral white matter. *Neuroimage* 2007;36:630–44.

Johansen-Berg H, Behrens TE. Diffusion MRI. London, United Kingdom: Elsevier; 2009:333–52, 434–36.

Aebly A, Liu Y, De-Torre X, et al. Maturation of thalamic radiations between 34 and 41 weeks gestation: a combined voxel-based study and probabilistic tractography using diffusion tensor imaging. *AJNR Am J Neuroradiol* 2009;30:1780–86.

Powell HW, Parker GI, Alexander DC, et al. Hemispheric asymmetries in language-related pathways: a combined functional MRI and tractography study. *Neuroimage* 2006;32:388–99.

Choi CH, Lee JM, Koo BB, et al. Sex differences in the temporal lobe white matter and the corpus callosum: a diffusion tensor tractography study. *Neuroreport* 2010;21:73–77.

Campbell MJ, Maclin D. Medical Statistics. London, United Kingdom: John Wiley & Sons; 1999:148.

Harasty J, Double KL, Halliday GM, et al. Language-associated cortical regions are proportionally larger in the female brain. *Arch Neurol* 1995;52:171–76.

Badenmacher J, Morosan P, Schleicher A, et al. Human primary auditory cortex in women and men. *Neuroreport* 2001;12(8):1561–65.

Witelson SF, Glezer II, Kigar DL. Women have greater density of neurons in posterior cortical cortex. *J Neurosci* 1995;15:3418–28.

Kinney HC, Brody BA, Kolman AS, et al. Sequence of central nervous system myelination in human infancy: II. Patterns of myelination in autopsied infants. *J Neuropath Exp Neurol* 1988;47:217–34.

Thompson PM, Giedd JN, Woods RP, et al. Growth patterns in the developing brain detected by using continuum mechanical tensor maps. *Nature* 2000;404:190–93.

Dubois J, Dehaene-Lambertz G, Perrin M, et al. Asynchrony of the early maturation of white matter bundles in healthy infants: quantitative landmarks revealed non-invasively by diffusion tensor imaging. *Hum Brain Mapp* 2008;29:14–27.

Neil JJ, Miller J, Mukherjee P, et al. Diffusion tensor imaging of normal and injured developing human brain: a technical review. *NMR Biomed* 2002;15:543–52.

Dubois J, Hertz-Pannier L, Cachia A, et al. Structural asymmetries in the infant language and sensori-motor networks. *Cereb Cortex* 2009;19:414–23.

Thompson DK, Inder TE, Faggian N, et al. Characterization of the corpus callosum in very preterm and full-term infants utilizing MRI. *Neuroimage* 2011;55:479–90.

Hindmarsh GJ, O'Callaghan MJ, Mohay HA, et al. Gender differences in cognitive abilities at 2 years in ELBW infants. *Early Hum Dev* 2006;82:115–22.

Maccoby EE, Jacklin CN. The Psychology of Sex Differences. Stanford, California: Stanford University Press; 1974.

Kramer JH, Kaplan E, Delis DC, et al. Developmental sex differences in verbal learning. *Neuropsychology* 1997;11:577–84.

Sommer IE, Alemán A, Somers M, et al. Sex differences in handedness, symmetry of the planum temporale and functional language lateralization. *Brain Res* 2008;1206:76–88.

Hervé PY, Leonard G, Perron M, et al. Handedness, motor skills and maturation of the corticospinal tract in the adolescent brain. *Hum Brain Mapp* 2009;30:3151–62.

Bademacher J, Bürgel U, Geyer S, et al. Variability and asymmetry in the human precentral motor system. A cytoarchitectonic and myeloarchitectonic brain mapping study. *Brain* 2001;124:2232–58.

Westerhausen R, Huster RJ, Kreuder F, et al. Corticospinal tract asymmetries at the level of the internal capsule: is there an association with handedness? *Neuroimage* 2007;37:379–86.

Geschwind N, Galaburda AM. Cerebral lateralization. Biological mechanisms, associations, and pathology: III. A hypothesis and a program for research. *Arch Neurol* 1985;42:634–54.

Thompson M, Cannon TD, Narr KL, et al. Genetic influences on brain structure. *Nat Neurosci* 2001;4:1253–58.

Toga AW, Thompson PM. Genetics of brain structure and intelligence. *Annu Rev Neurosci* 2005;28:1–23.

Lauterbach MD, Raz S, Sander CJ. Neonatal hypoxic risk in preterm infants: the influence of sex and severity of respiratory distress on cognitive recovery. *Neuropsychology* 2001;15:411–20.

Nunez JN, McCarthy MM. Sex differences and hormonal effects in a model of preterm infant brain injury. *Ann NY Acad Sci* 2003;1008:281–84.

Kagan J, Herschowitz EC. *Young Mind in a Growing Brain*. Mahwah, New Jersey: Lawrence Erlbaum; 2005.