Apparent Diffusion Coefficient Determination in Normal Fetal Brain: A Prenatal MR Imaging Study

Andrea Righini, Elena Bianchini, Cecilia Parazzini, Patrizia Gementi, Luca Ramenghi, Cristina Baldoli, Umberto Nicolini, Fabio Mosca, and Fabio Triulzi

BACKGROUND AND PURPOSE: Diffusion-weighted MR imaging studies of normal brain development have focused on premature babies who were free of focal lesions on conventional MR images. The condition of prematurity, however, is dissimilar to intrauterine life. We sought to establish normal values of fetal brain apparent diffusion coefficient (ADC) to highlight its abnormal changes in pathologic conditions and to obtain information about normal brain development.

METHODS: We measured the ADC, in utero, by using an echo-planar three-axes diffusion-sensitized sequence (b factor, 0 and 600 s/mm²), in frontal and occipital white matter and basal ganglia gray matter of 15 fetuses. Their gestational ages ranged from 22 to 35 weeks, and the postnatal MR images or sonograms revealed normal brain.

RESULTS: Mean ADC value was 1.96 ± 0.1 μm²/ms (SD) in frontal white matter, 1.95 ± 0.1 μm²/ms in occipital white matter, and 1.56 ± 0.1 μm²/ms in basal ganglia. A significant negative correlation between ADC and gestational age was found for basal ganglia, whereas only a trend was present for frontal white matter.

CONCLUSION: Although moderately higher, the ADC determinations we obtained are consistent with those reported in the literature in postnatal studies performed in premature babies.

The measurement of water apparent diffusion coefficient (ADC) in the human brain by means of diffusion-weighted (DW) MR imaging has provided valuable information in adults with stroke, seizures, trauma, or white matter disorders (1–4). In premature and full-term neonates, DW imaging has allowed observations to be obtained not only in certain brain diseases, such as global hypoxia-ischemia (5–8) or periventricular leukomalacia (9), but also in normal cerebral development (10–13). Prenatal MR imaging is widely used to assess fetal brain abnormalities in cases of unclear or incomplete prenatal sonographic (US) results. In particular, prenatal US has poor sensitivity in the detection of acute lesions, such as the early stages of hypoxic-ischemic damage (14), which is one of the leading causes of fetal brain destructive processes. Prenatal DW imaging enabled detection of acute fetal brain ischemic lesions in a very recent report (15). We thought it would be worthwhile to establish the normal values of fetal brain ADC not only to highlight abnormal changes in pathologic conditions, but also to add information about normal brain development. DW imaging studies on normal brain development have focused, so far, on premature babies who were free from focal lesions noticeable on conventional MR images (11–13). Despite the high quality and high spatial resolution of such imaging data, the condition of prematurity is far from being similar to intrauterine life. For these reasons, we measured the ADC in cerebral white and gray matter of fetuses with a normal brain and whose gestational ages ranged from 22 to 35 weeks.

Methods

Subjects

Among consecutive prenatal MR imaging studies routinely performed for clinical purposes at our institution from January 2001 to March 2002, the images of 15 fetuses were included in the study. All mothers signed an informed consent form. The following inclusion criteria were adopted: 1) fetuses imaged because of clinical and US suspicion of anomalies in organs other than the fetal brain, but with normal brain at prenatal US...
and at T2-weighted single-shot fast spin-echo (SE) MR imaging (nine cases); 2) fetuses imaged with prenatal MR imaging because of the risk of possible brain damage, but with normal prenatal brain sonograms, as in twin-to-twin transfusion syndrome, and resulting in negative prenatal T2-weighted single-shot SE MR images (two cases); and 4) fetuses imaged because of suspected mild intracranial anomalies at prenatal US such as widening of cisterna magna and with otherwise normal prenatal sonograms and prenatal T2-weighted single-shot fast SE MR images (two cases). All pregnancies were free from history of maternal diabetes or hypertension. All neonates had a normal neurologic examination. In all but one case (fetus 7 in Table), postnatal sonograms or MR images were available and evaluated separately by three pediatric neuroradiologists (A.R., E.B., and C.P.), who confirmed normal brain findings, with the exception of the two cases in criterion 4, who showed widening of cisterna magna and posterior fossa arachnoid cyst at postnatal MR imaging, respectively. We decided to include these two cases because postnatal findings related to cerebellar and cerebral morphology and parenchymal signal intensity were unremarkable, as was the neurologic examination. Although it is well known that the time course of normal brain development can present some delay in twins, we also included the two cases of twin-to-twin transfusion syndrome because their ADC values were similar to the ones of the singletons of the same gestational age.

Mean maternal age was 32.3 ± 4.1 years (SD), and mean gestational age of the fetuses was 27.6 ± 4.4 weeks (range, 22–35 weeks) (Table).

**MR Imaging and Data Analysis**

All MR imaging was performed with a 1.5-T unit (Horizon LX, Echospeed; GE Medical Systems, Milwaukee, WI). No sedation of the mother or fetus was required. For conventional prenatal MR imaging, a body dedicated phased-array coil was used, and no breath holding was required of the mother. Multispectral T2-weighted single-shot fast SE 3.5-mm-thick sections were acquired with the following parameters: TR, indefinite; TE, 90 ms; field of view, 320–380 mm; matrix, 256 × 192. For DW imaging, we switched to the whole-body coil (without repositioning the mother), because on our system phased-array coils did not allow performance of echo-planar DW imaging. By using sagittal T2-weighted single-shot fast SE sections as a scout view, axial single-shot echo-planar DW imaging sections were prescribed on fetal brain. The DW sequence, which had an acquisition time of 20 seconds, was performed with breath holding by the mother. Because of frequent image motion degradation, usually two or three DW sequence acquisitions had to be performed before obtaining satisfactory images. DW imaging section thickness varied from 5 to 6 mm according to fetal head size. Other parameters were 5000/81 (TR/TE), field of view of 320 mm, and matrix of 128 × 128, leading to an in-plane resolution of 2.5 × 2.5 mm. The sensitizing diffusion gradients were applied on the three (x, y, z) orthogonal axes with two b factors (0 and 600 s/mm²) per axis.

Rotational relatively insensitive mean diffusivity maps (trace ADC maps) were then calculated by using a standard software (Func tool) running on the imager’s main console. For each case, a pediatric neuroradiologist (A.R.), after selecting the basal ganglia, traced circular regions of interest (ROIs) on the corresponding echo-planar (b = 0 s/mm², T2-weighted) image in the basal ganglia, frontal white matter, occipital white matter, and CSF of the lateral ventricles (Fig 1). ROI sizes varied from...
30 to 50 mm$^2$ and 40 to 80 mm$^2$ for the white matter and basal ganglia regions, respectively, according to the gestational age and size of the fetal brain. From the image obtained with $b = 0$ s/mm$^2$, ROIs were then automatically transferred onto the trace ADC map. The ROI values for frontal white matter were available in 13 of the 15 cases; for occipital white matter, in 14 cases; and for basal ganglia, in 13 cases. Since no statistically significant difference was found between the ROI values of the right and left hemispheres, the average between ROI values of both hemispheres was considered for final statistics. To assess statistically significant differences between the mean ADC value in the ROIs traced, a paired $t$ test was used, and $p < .05$ was considered a statistically significant difference.

**Results**

The mean ADC value was $1.96 \pm 0.1 \, \mu\text{m}^2/\text{ms}$ (SD) in frontal white matter, $1.95 \pm 0.1 \, \mu\text{m}^2/\text{ms}$ in occipital white matter, and $1.56 \pm 0.1 \, \mu\text{m}^2/\text{ms}$ in basal ganglia; the difference between basal ganglia and both frontal and occipital white matter was statistically significant ($P < .0001$, paired $t$ test). The mean ADC value in CSF was $3.1 \pm 0.1 \, \mu\text{m}^2/\text{ms}$ (Fig 2). A significant negative correlation between ADC values and gestational age could not be found for frontal and occipital white matter, although there was a trend in this sense for frontal white matter. A significant negative correlation with gestational age was found for the basal ganglia ROIs (Fig 3).

**Discussion**

DW imaging and ADC determinations are feasible in fetal brain by using prenatal MR imaging. This type of study is intrinsically more difficult than postnatal studies, since image degradation due to mother-fetal motion artifact and to aorta pulsatility effects may require more than one acquisition to be repeated to obtain satisfactory images. The signal-to-noise ratio and the spatial resolution are not as good as those obtained with postnatal imaging in infants and in adult imaging. However, the ADC determinations we obtained are consistent with those reported in postnatal studies performed in premature babies (11–13). Moreover, our ADC values for CSF are similar to what has been reported in neonates and adult ventricles (12, 16), supporting the consistency of our measurements. The mean ADC value for white matter ROIs was higher than that of the basal ganglia gray matter, as already demonstrated by previous studies in premature neonates (12, 13). The explanation for this is not totally clear, but it could be because gray matter has more cells and membranes per volume unit that unmethylated white matter, and the interstitial water is more abundant in the latter.

The average ADC values for both white and gray matter in our study were moderately higher than those reported in premature neonate DW imaging studies (range of means for white matter, approximately $1.8–1.9 \, \mu\text{m}^2/\text{ms}$; range of means for basal ganglia gray matter, approximately $1.2–1.3 \, \mu\text{m}^2/\text{ms}$) (11–13). The reason for this discrepancy is not clear. The mean gestational age of our fetuses was moder-
FIG 3. Graphs show the correlation between ADC values for the ROIs traced and gestational age. A significant negative correlation was present only for the basal ganglia ROI.
A, Basal ganglia: Y = 2.222 – 0.024X; R² = 0.471; P = .0096.
B, Frontal white matter: Y = 2.394 – 0.015X; R² = 0.182; P = .1457.
C, Occipital white matter: Y = 2.14 – 0.007X; R² = 0.028; P = .5654.
diffusion tensor imaging, their relative anisotropy index and ADC would probably be different from those of other white matter areas.

Because of the relatively coarse resolution of the echo-planar images we obtained, it is possible also that the basal ganglia ROI measurements might have been affected by partial volume problems, in particular for the inclusion of some adjacent white matter areas like the posterior limb of the internal capsule. However, previous work in the premature age range (11) has shown that ADC measurements in the posterior limb of the internal capsule tend not to vary significantly with postconceptual age, probably because of the already advanced myelination process of this structure, so this partial volume effect should have affected the ADC values of basal ganglia ROIs in our cases in a constant manner.

To obtain better signal-to-noise ratio on DW images in the present study, we used a b factor maximum value of only 600 s/mm², similar to previous studies in premature neonates (11). Recent work on neonate and adult brains (24) has confirmed the presence of a slow and a fast component of the ADC decay curve. The ADC decay curve related to the neonate brain has a significantly higher fraction of the fast diffusion component with respect to that of the adult brain. The use of relatively lower b factor values, such as in our study, may therefore be preferred in investigating normal fetal brain, since the left portion of the ADC decay curve (fast component) prevails. However, in case of pathologic ADC reduction, as in acute ischemia, when the slow component prevails, the use of lower b values might underestimate the degree of ADC reduction.

One of the main limitations of this study is that it does not deal with the important issue of white matter anisotropy and its variations along with brain maturation, as shown by previous investigations in neonates (11, 13, 25). Our ADC determination was not completely rotationally insensitive, since three-axis DW imaging acquisition cannot accurately explore the preferential diffusivity within each voxel. In the aforementioned studies in neonates, technically adequate acquisitions were based on at least six-axis diffusion gradients, to allow a complete spatial characterization of the diffusion tensor for each voxel. In these studies, brain maturation was associated with an increase in white matter anisotropy index, which was noticeable even in areas known from histologic examination to be not yet myelinated: the so-called premyelination maturation process or premyelinating state. This premyelinating state is characterized by increase in axon diameter, axonal membrane changes, increased concentration of microtubule-associated proteins, and early wrapping of axones by oligodendroglial processes (11, 26).

Technical and methodological improvements in fetal DW imaging, such as the use of surface coils, will allow better spatial resolution and signal-to-noise ratio images to be obtained. Sequences exploring the preferential water diffusivity on more than six axes and allowing diffusion tensor determination will probably provide valuable and more detailed information about white matter maturation, during both the myelinating and premyelinating processes; however, the need for breath-hold images in pregnant women remains an obstacle, as does the difficulty of performing so rapid (breath-hold compatible) diffusion tensor imaging acquisitions.

Even if MR imaging has been routinely and widely used in fetal imaging in the last 10 years (27) and the subjects we studied had already been scheduled for clinical imaging, the safety issue of prenatal MR imaging should be mentioned, in particular in relation to the use of echo-planar imaging sequences. Few studies have so far focused on the safety issue regarding the use of MR imaging in human fetal imaging. One in particular by Baker and colleagues (28) investigated the potential long-term adverse effects of echo-planar imaging in children who had been imaged in utero. Their data, however, did not demonstrate any future anomaly, including hearing damage, related to intrauterine exposure to echo-planar imaging.

The present study provides information about brain ADC values in normal fetuses of the gestational age range that usually applies also to fetuses imaged for suspected cerebral malformations or destructive lesions. It would be of interest to explore possible abnormal ADC variations in fetuses with major cerebral malformations or with severe intrauterine growth retardation.

Addendum: Phantom Study

A phantom test was conducted to assess if the ADC measurements we obtained in fetuses by using the whole-body coil were comparable to those obtained in premature neonates during routine clinical studies, which are usually performed with a standard quadrature head coil. A corn oil phantom was imaged by using the whole-body coil and the standard quadrature head coil; all the DW imaging acquisition parameters were the same as for the fetal MR imaging protocol. At 22.5°C and after 10 measurements, the mean ADC value obtained with the whole-body coil (0.46 ± 0.03 μm²/μs) was similar to that obtained with the quadrature head coil (0.43 ± 0.02 μm²/μs).

Acknowledgments

We thank Mr. Giancarlo Lembo for assistance in performing the MR imaging studies.

References


MacLaurin JC. Changes in body water distribution during the first two weeks of life. *Arch Dis Child* 1966;41:286–291


