DIFFUSION TENSOR IMAGING DETECTS ABNORMALITIES IN THE CORTICOSPINAL TRACTS OF NEONATES WITH INFANTILE KRABBE DISEASE

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Abstract

Background and Purpose—It is not possible to determine if babies diagnosed with Krabbe disease through statewide newborn screening programs will develop disease as infants, juveniles or adults. The only available treatment for this fatal neurodegenerative condition is unrelated umbilical cord transplantation but this treatment is only effective before clinical symptoms appear. Therefore, a marker of disease progression is needed. The purpose of this study was to evaluate the use of Diffusion Tensor Imaging (DTI) with fiber tracking in identifying early changes in major motor tracts of asymptomatic neonates with infantile Krabbe disease.

Methods—Six babies with infantile Krabbe disease identified because of family history had a brain MRI within the first 4 weeks of life. Six direction DTI and quantitative tractography of the corticospinal tracts were performed. Hypothesis tests, one for each hemisphere, were used to test whether the FA ratio of the babies with infantile Krabbe disease was significantly different from 45 age and gender matched controls.

Results—The average FA ratio for the Krabbe patients was 0.89 and 0.87 for left and right tracts, respectively (p= 0.002 and p<0.001). After adjusting for gestational age, gestational age at birth, birth weight, gender and race, the 6 Krabbe patients had significantly lower FA values than the controls (p<0.001).

Conclusions—DTI with quantitative tractography detected significant differences in the corticospinal tracts of asymptomatic newborns that had the early onset form of Krabbe Disease. Once standardized and validated, this tool has the potential to be used as a marker of disease progression in babies diagnosed through statewide newborn screening programs.
Krabbe Disease is an autosomal recessive disorder characterized by deficiency of the lysosomal enzyme galactocerebrosidase (GALC) which aids in the breakdown and removal of galactolipids found in myelin. The accumulation of galactolipids results in inflammation, rapid demyelination and death within 2 years.1 This is a very rare disorder with an incidence of 1 in 70–100,000.1 The phenotypic expressions of the mutations vary widely with the disease presenting as an infantile, juvenile or adult form. The infantile presentation has an onset within the first 6 months of life and is the most frequent form of presentation. Umbilical cord blood transplantation is the only available treatment, and is only effective before symptoms develop.2 Treated infants preserve cognitive function but develop some degree of motor disability that ranges from mild to severe.

With the implementation of statewide mandatory newborn screening programs, babies with no previous family history are being diagnosed. Among the babies who screen positive for Krabbe disease, it is not possible to identify with certainty which babies will develop the infantile, juvenile or adult forms because phenotype predictions cannot be made a priori from the genetic mutation or degree of enzyme deficiency. Therefore, the development of a predictive measure that identifies changes in the brain before symptoms develop could potentially clarify which newborns will develop the infantile form. This information is critical for clinicians counseling parents about the treatment of a baby who screens positive for Krabbe disease. A better understanding of markers of disease onset and expected phenotype after treatment would also help parents and clinicians make decisions about the optimal time of treatment.

Mild hyperintense signal intensity abnormalities in the internal capsule have been previously reported in untreated neonates with Krabbe disease.3 These findings coupled with reports showing rapid increase in FA with age4 and in the central regions of the internal capsule5–9 lead us to believe that quantifying these changes may help identify which asymptomatic babies should be treated because they are more likely to rapidly progress and die.

The purpose of this study is to evaluate whether diffusion tensor imaging (DTI) with quantitative tractography is able to detect early changes in major motor tracts before clinical symptoms develop. We further describe the relationship between changes in these motor tracts and degree of motor disability at 12 months of age.

**Methods**

Newborns with Krabbe Disease, identified by family history or through the New York State screening program, were referred to the Program for Neurodevelopmental Function in Rare Disorders (NFRD) at the University of North Carolina at Chapel Hill for assessment of baseline neurological function before receiving unrelated umbilical cord blood transplantation (UCBT) at Duke University Medical Center. The assessment included a detailed neurodevelopmental evaluation concurrent with a brain MRI within the first 4 weeks of life. A follow up evaluation of motor function was done at 6, 9, 12 and 24 months. Six babies with infantile Krabbe disease diagnosed prenatally or at birth because of family history were scanned without sedation on a Siemens Allegra 3T head-only MR scanner. Neonatal controls were collected as part of a separate study and were scanned on the same scanner using an identical protocol.10 The studies were approved by the IRB at the University of North Carolina.

**Neurodevelopmental Assessments**

The Peabody Developmental Motor Scales-2 (PDMS-2)11, the Mullen Scales of Early Learning12 and the Protocol for the Assessment of Neurodevelopment in Infants (PANDI) [unpublished] were used to assess cognitive and motor development in children with Krabbe Disease. Age equivalents were used to track development over time and compare across tests.
Imaging

The details of the image acquisition and processing have been reported in a previous publication. Ten main imaging sequences were employed, including a magnetization prepared rapid gradient echo (MP-RAGE) T1-weighted, a PD/T2 TSE sequence, and a single shot echo planar (EPI) DTI sequence. The imaging parameters for the DTI sequence were as follows: repetition time (TR) = 4219 ms; echo time (TE) = 92.2 ms; inplane resolution = 2×2mm2 with a slice thickness of 2mm, 6 diffusion directions and 10 averages. Seven images were acquired, one without diffusion gradient (b=0) while the remaining six with b=1000s/mm2. Prior to imaging session, neonates were fed, swaddled, fitted with ear protection and had their heads fixed in a vac-fix device. A pulse oximeter was used to monitor heart rate and oxygen saturation.

Diffusion Tensor Image Processing

The processing protocol consisted of multiple repeated DTI acquisitions that were co-registered, filtered for bad slices and finally averaged via an automatic DTI quality control tool, called DTI checker (http://www.ia.unc.edu/dev/download/dtichecker). The final DTI dataset was fed into a fiber tracking tool, which automatically computes the Fractional Anisotropy (FA) and Mean Diffusivity (MD) maps. Regions of interest were outlined on the base (b=0) image for regional FA and MD analysis, and for DTI fiber tracking using our ITK-SNAP tool. For the analysis plan, the fiber circuitries of interest were determined in collaboration with a neuroanatomist (James Fallon, University of California, Irvine). Our Fiber Tracking tool computes DTI fibers via a streamline algorithm that follows the major diffusion direction from specified source to target regions. DTI Fibers were post-processed using our Fiber Viewer tool by removing outliers, grouped into separate left and right hemispheric tracts. Correspondence along all fibers was established via a user-selected origin for each fiber-bundle and an equal step parametrization along every fiber. Based on this correspondence, the mean and standard deviation of FA and MD across fiber bundles were plotted in curves and analyzed along the DTI fiber bundles.

After the fiber tracts were defined, mean FA values were calculated at 20, 3 mm intervals along the tracts. However, because preliminary data have shown that the central and peripheral regions of the tracts show increased variability due to low signal to noise ratio only the 30 mm segment near the internal capsule was used to index FA and MD. Previous studies have shown comparisons of specific distances along the tracts (e.g. @ 30 mm or 55 mm) to be highly sensitive to the spatial alignment of the tracts between subjects; by taking the mean of this segment we increase both the precision of the measure and the power to detect differences.

Statistical methods—The FA value for Krabbe patients was scaled in terms of the expected FA in typical neonates. To calculate this ratio an expected mean value based on the age at scanning, gestational age at birth, gender, and birth weight was estimated based on a general linear model that was fit to the 45 typical neonates. The outcome variable was the mean FA value for the 30mm segment with age at scanning, gestational age, gender, and birth weight as predictors. The left and right tracks were modeled separately. The coefficients from these two models were then used to calculate the expected mean FA values for the left and right tracks of each of the 6 Krabbe patients. The actual mean FA values of the Krabbe patients were then divided by the expected value based on the predictors, producing a ratio which represents the percentage of normal FA.

To test for mean and slope differences between the patients with Krabbe and typical controls a mixed regression model with repeated measures between the two hemispheres (left/right) was fit. The model regressed group, corrected age at the scan, hemisphere (left/right), the interaction terms between group and left, and group and hemisphere, on FA ratio. The data
were centered at 40 weeks gestational age and the main effect of group was tested at this age, as this was the mean for the Krabbe patients.

**Results**

Four newborns with Krabbe Disease identified by family history and two identified through newborn screening were evaluated between December of 2004 and September of 2007. Their corrected age at the time of the MRI ranged from 1 to 3 weeks of age (38 to 42 weeks from gestation) and their birth weight ranged from 2.28 to 4.13 kg. There were 3 females and 3 males (Table 1). All patients were subsequently transplanted with unrelated umbilical cord blood transplantation. Two patients died of transplant related complications, one of veno-occlusive disease and one of persistent pulmonary hypertension. The four patients that survived now range in age from 18 months to 4 years. Krabbe patients generally weighed less at birth and were younger than controls at both birth and at the time of the scan. However, these variables were controlled for in all the statistical tests. Table 1 shows descriptive information for the 6 Krabbe patients and the 45 controls. Of the 6 patients, 5 had relatively normal neurological exams and were classified as stage 1, one patient had minimal neurological symptoms, clonus in one lower extremity and was classified as stage 2. (2)

When the fiber tracts were visualized by color coding for FA values in both controls and patients with Krabbe Disease, the normal pattern of myelination that goes from the center to the periphery is evident in both groups. However, the Krabbe group show lower FA values along most of the fiber bundle than any of the controls and lower FA values towards the periphery. (Fig 1). These images produced by our Fiber Track viewer software are indicative of a more immature pattern of myelination in the Krabbe group.

The mean FA of the central area of the corticospinal tracts in the control patients shows a clear trend of increasing FA with age (Fig 2). However, this trend was less clear in the Krabbe patients. After adjusting for gestational age, gestational age, birth weight, gender and race, the 6 Krabbe patients had significantly lower FA values than the controls (p<.001). Figure 3 represents the FA ratios for the 6 Krabbe patients and the 45 controls. The average FA ratio for the Krabbe patients was .89 and .87 for left and right tracts, respectively (p=.002 and p<.001). While it appears that the FA values become more abnormal with increasing age, this effect is not statistically significant (p=.12). There was no significant difference between the FA values of the left and right hemispheres (p = .58) or in the mean fractional anisotropy by group and location (right/left) (Table 2).

Because two of the children with Krabbe fall outside the range of our controls we reanalyzed the data excluding these two children from the analysis. Using only the children who were greater than or equal to 40 weeks at the time of scan, the differences between the Krabbe group and the controls increases to .18 and .20 (p=.002 and p<.001) for left and right hemispheres.

All four children are within the range of age appropriate development in cognitive, receptive language and fine motor skills. However, 3 of the 4 patients have gross motor delay at the last follow up visit. The neurodevelopmental trajectories in the motor area were correlated to the mean FA ratios in the 4 surviving patients. Of the 2 patients with the highest FA, one can walk and run at 4 years of age, and the other can walk with a walker at 36 months. The patient with the lowest neonatal FA ratio shows the greatest delay in the motor area not being able to walk independently at 25 months of age. Another patient with low FA is not able to walk at 18 months and has the development of a 13 month old. All patients are cognitively normal.
Discussion

We have demonstrated in a very small sample of patients that DTI with quantitative tractography methodology is able to detect significant differences in the corticospinal tracts of neonates with Krabbe Disease before clinical symptoms develop. All of the 6 patients with infantile Krabbe Disease had lower FA values across the DTI fiber bundle of the internal capsule (IC) when compared to healthy age matched controls. The average FA ratio for the Krabbe patients was 0.89 and 0.87 for left and right tracts, respectively, with a significant difference ($p<.001$) in FA values between Krabbe and control values. Interestingly, we also found a strong trend albeit non significant in FA with age, with the Krabbe group becoming progressively abnormal over a matter of weeks. Of the 4 children followed > 12 months after transplantation, the children with FA values around 0.90 (0.89 and 0.91) are performing better in standardized assessments than the children with FA values of 0.73 and .84.

A previous study has shown normal myelination and axonal integrity as measured by DTI methods in Krabbe babies. However their methodology was very different from ours. First, the authors looked at many regions that do not show early myelination and thus are less likely to show a difference. Second, when they looked at the internal capsule, they used the manual region of interest based methodology that often mixes different fibers that cross the same region. Our methodology nicely picks out only those image regions that are associated with the corticospinal tract.

The early abnormalities in FA seen in our study in the internal capsule, a major motor pathway, may explain the plateau in motor development seen in treated babies with Krabbe disease even in the presence of improved overall brain myelination. Previous studies have shown that FA appears to be dependent on tract organization. This disruption could also explain the motor impairment that may be present even before myelination is disrupted by the lack of GALC. Further studies in animal models looking at the influence of GALC in tract development would be needed to confirm this hypothesis.

This study’s greatest limitation is the small number of infants with Krabbe (n=6) but this represents a very large portion of this population and the results are statistically significant. While random sampling was not employed sequential enrollment was used in that each child who was thought to have infantile Krabbe who presented at our center was scanned. During the study period our center saw nearly all pre-symptomatic children from the U.S. who were suspected to have early-infantile Krabbe. Therefore, there is no reason to believe that the results are not similar to those from a random sample.

Another limitation is the limited number of controls who were scanned prior to 41 weeks gestational age. While a linear model was assumed to create estimates of typical FA values during the 38–41 week period it is possible that the linear relationship observed from 41–48 weeks does not hold for this early period of development. However, since the 2 youngest babies showed the least differences from our estimate of typical babies we believe our results to be a conservative estimate. Conversely, any estimates of increasing abnormality with age would be highly sensitive to any misspecification of the model during this early period. For this reason the non-significant age effect should be examined with caution.

The fact that all babies showed abnormalities in motor examination confirms that these babies were going to develop Krabbe disease since the pattern of neurological involvement has been described in previous studies and this pattern is not seen in babies transplanted for other diseases. We have previously published our ability to predict motor outcome based on the staging system developed by our group. However, this is subjective, based on the experience of the examiner and therefore more difficult to disseminate. Correlating development with previous DTI measures is more objective and the methodology easier to disseminate to sites...
with no experience in evaluating these patients. However, the prediction could be challenging due to uncontrolled environmental influences. Intervention services, length of hospitalization and medical complications may impact a child's motor development. For example, one of the patients had higher initial values of FA than the other babies but developed graft versus host disease that required high doses of steroids. The increased weight and steroid induced myopathy affected this child's motor development. Therefore, this patient may be more delayed than would be predicted by the FA at the time of transplantation.

Patients who screen positive for low GALC but have no family history may develop early onset disease, may be healthy or clinically silent until late childhood or even adulthood. An objective measure of disease progression is necessary to decide whether an asymptomatic baby should be treated since once clinical symptoms appear it is too late to derive benefits from treatment. As previously reported, conventional MRI scans and neurophysiological tools cannot be used to predict outcome or to correlate with the disease stage because of poor standardization of these tools in the neonatal period. In this study, even a small sample of patients showed significant differences between healthy controls and babies diagnosed with infantile Krabbe disease. The differences in FA in the superior and middle cerebellar peduncle may help differentiate between babies who will develop the early onset disease, those who will develop disease at a much later time, and those who will never develop disease. This approach can provide very consistent FA values and detect small regional differences and age related changes in the first 2 years of life. For this reason, DTI with quantitative tractography may prove to be an excellent technology for studying infantile Krabbe disease and other demyelinating conditions of early childhood. A large sample of healthy controls and babies with low enzyme that do not develop early infantile disease will need to be studied, for an adequate standardization and validation of this tool. Once fully standardized, this tool has the potential to be a key component of a population screening program to help determine who should receive treatment before symptoms develop.

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Fig. 1.
Visualization of DTI fiber tracts through the right hemispheric internal capsule with Fractional Anisotropy (FA) coloring of the fibers from a lateral viewpoint. Top row: 5 representative control subjects, bottom row: 4 selected Krabbe subjects. Visually, all Krabbe subjects show lower FA values along most of the fiber bundle than any of the controls.
Fig. 2.
Raw FA mean values for Infantile Krabbe patients and controls. Each point represents a child's mean FA across the central portion of the corticospinal tract. The red triangles represent the 6 Krabbe patients while the blue diamonds show the typical controls. The red and blue lines represent the linear trend related to corrected age at the time of the MRI scan.
Fig. 3.
Normalized FA ratios for Infantile Krabbe patients and controls. The FA ratio is the observed FA divided by the expected FA given the child’s gestational age at birth, corrected age at scan, birth weight and gender. The red triangles represent the 6 Krabbe patients while the blue diamonds show the typical controls. The horizontal black lines represent the standard deviations based on the controls.
Table 1
Descriptive information for infantile Krabbe patients and healthy controls.

|                        | Krabbe (n=6) | SD | Controls (n=45) | SD |
|------------------------|--------------|----|-----------------|----|
| Birth Weight (grams)   | 3.13         | .65| 3.45            | .45|
| Gestational age birth (weeks) | 38.3 | 1.5| 39.7            | 1.0|
| Gestational age at scan (weeks) | 40.1 | 1.9| 42.9            | 1.7|
| male                   | 50%          |    | 60%             |    |
| white                  | 100%         |    | 84%             |    |
Table 2
Mean fractional anisotropy by group and location (right/left)

| Variable | Krabbe Mean | SD  | Controls Mean | SD  |
|----------|-------------|-----|---------------|-----|
| Right    |             |     |               |     |
| Mean FA  | 0.36        | 0.04| 0.45          | 0.04|
| FA Ratio | 0.87        | 0.09| 1.00          | 0.08|
| Left     |             |     |               |     |
| Mean FA  | 0.36        | 0.04| 0.43          | 0.04|
| FA Ratio | 0.89        | 0.09| 1.00          | 0.08|