Neurochemistry of Hypomyelination Investigated with MR Spectroscopy

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Proton magnetic resonance spectroscopy (MRS) allows the noninvasive exploration of tissue metabolism in vivo, providing neurophysiological and neurochemical information. N-acetylaspartate (NAA) is generally considered to be a marker of neurons and axons, and many neurodegenerative disorders, including demyelinating disorders, exhibit a decrease in total NAA (tNAA). MRS in human hypomyelination disorders, such as Pelizaeus-Merzbacher disease (PMD), is characterized by normal to elevated tNAA, elevated myo-inositol and creatine (Cr), and normal to decreased choline (Cho).

MRS in the thalamus of a hypomyelinating mouse model, a myelin synthesis-deficient (msd) mouse, a model of connatal PMD with mutation of the Plp1 gene, revealed increased tNAA and Cr and decreased Cho. That of a shiverer mouse with an autosomal recessive mutation of the Mbp gene showed decreased Cho with normal tNAA and Cr. Accordingly, the reduction of Cho on MRS might be a common marker for hypomyelinating disorders.

tNAA concentrations range from normal to increased, probably depending upon the underlying pathology of oligodendrocytes. tNAA may be increased in hypomyelination with a reduced number of mature oligodendrocytes, such as PMD.

Keywords: hypomyelination, magnetic resonance spectroscopy, mouse model, myelin basic protein, proteolipid protein

Introduction

Magnetic resonance (MR) imaging has greater sensitivity than computed tomography (CT) in the detection of lesions of the white matter that result from abnormal conditions of the myelin, including demyelination, hypomyelination, white matter rarefaction, and cystic degeneration. However, its specificity is limited because all these conditions result in T2 prolongation of the white matter.

Proton MR spectroscopy (MRS) allows the noninvasive exploration of tissue metabolism in vivo, providing neurophysiological and neurochemical information. As the N-acetylaspartate (NAA) is synthesized in neuronal mitochondria and transported to axons (Fig. 1), NAA at 2.01 ppm on MRS is generally considered an important marker of viable, functioning neurons and axons. NAA is either released from the neuron or transported to oligodendrocytes, where it is catabolized by aspartoacyclase (ASPA) into acetate and aspartate. NAA is also the precursor for the synthesis of N-acetylaspartylglutamate (NAAG) (2.04 ppm on MRS) in neurons. The biosynthesis or regulatory mechanisms of NAAG are poorly understood, but it modulates glutamatergic synaptic transmission and is released along with other neurotransmitters from neuronal synapses. Extracellular NAAG is hydrolyzed into NAA and glutamate by glutamate carboxypeptidase II (GCP II) on the membrane surface of astrocytes, and the breakdown products are taken in by the astrocytes. Accordingly, NAA is purported to have signal functions for axon-oligodendrocytes and NAAG, for axon-astrocytes.

The choline (Cho) peak at 3.2 ppm likely contains various cell membrane precursors or breakdown products, such as phosphocholine, glycerophosphocholine, and phosphatidylcholine. Therefore, its elevation is seen in conditions of enhanced
membrane turnover, such as myelination, active demyelination, and tumor growth. Thus, many neurodegenerative disorders, including demyelinating disorders (with increased Cho) and neuronal degenerative disorders, exhibit a decrease of total NAA (tNAA; NAA and NAAG, which are difficult to distinguish on clinical MRS with a 1.5- or 3-T magnet).

I herein review MRS findings in hypomyelinating disorders of humans and their mouse models.

**Myelin Structure**

The myelin is a spiral membranous structure around the axons that comprises a protein-lipid-protein structure. The lipid layers are composed of a bimolecular layer of hydrocarbon chains, cholesterol, phospholipids, and glycolipids. A dark line in electron microscopy, called the major dense line, contains myelin basic protein (MBP), an intracellular protein attached to the inner surface of the cell membrane and situated mainly in the cytoplasm (Fig. 2). A less electron-dense protein line, the intraperiod line, represents proteolipid protein (PLP) in the outer portion of the cell membrane and in the extracellular space (Fig. 2). The 2 major structural proteins of myelin, PLP and MBP, constitute 50% (PLP) and 30% (MBP) by weight of myelin proteins. The deficiency of PLP
can cause hypomyelinating disorders, i.e., Pelizaeus-Merzbacher disease (PMD) in human and myelin synthesis-deficient (msd) mouse; and that of MBP can cause shiverer mouse.

MRS in Hypomyelinating Disorders in Humans

The term hypomyelination describes a permanent, substantial deficit of myelin deposition in the brain. In patients with classic-type PMD with duplication of the PLP1 gene, a representative hypomyelinating disorder, analysis of quantitative MRS with LCModel revealed increased tNAA, creatine (Cr), and myo-inositol (mIns) with normal to decreasing Cho (Fig. 3).9,10 mIns is almost exclusively located in astrocytes and is recognized as an astrocyte-specific marker.11 Astrocytes also contain relatively high Cr, compared with neurons.12 Concomitant astrogliosis observed in PMD13,14 probably results in increased mIns and Cr. It is not certain whether Cho is normal or decreased, but a reasonable decrease in Cho reflects decreased myelin synthesis and turnover, as observed in the brain in PMD.13,14 An increased concentration of Cr prevents detection of increased tNAA when the tNAA/Cr ratio is utilized. This may explain the failure in previous studies that utilized relative values to detect increased tNAA.15,16

Fig. 2. Schematic of the structure of myelin. (Reprinted from Reference 7 with permission)

Fig. 3. Magnetic resonance (MR) imaging (A) and spectroscopy (MRS) (B) of a 12-year-old patient with Pelizaeus-Merzbacher disease (PLP1 duplication) shows T2 prolongation in the white matter; increased total N-acetylaspartate (tNAA), creatine (Cr), and myo-inositol (mIns); and reduced choline (Cho) (1.01 mM, normal mean ± standard deviation [SD] 1.31 ± 0.09 mM), as analyzed with LCModel. Normal MRS (C) of the 12-year-old boy showed a Cho concentration of 1.31 mM.
In another hypomyelinating disorder, hypomyelination with atrophy of the basal ganglia and cerebellum (HABC), MRS showed normal tNAA and Cho with increased mIns and Cr.\textsuperscript{17} MRS in patients with Pelizaeus-Merzbacher-like disease (PMLD) also showed normal ratios of tNAA/Cr and Cho/Cr.\textsuperscript{18} Based on these data, MRS in hypomyelination disorders in humans is characterized by elevated to normal tNAA, elevated mIns and Cr, and normal to decreased Cho, findings distinct from those of MRS in active demyelination, which include decreased tNAA and increased Cho.\textsuperscript{6} The increase of tNAA may be explained by the higher axonal density in the absence of oligodendrocytes and myelin sheath\textsuperscript{10}; however, the exact mechanism is uncertain. Neither is it clear whether changes in the level of NAA or NAAG, or both, result in the increased tNAA.

**MRS in the Hypomyelinating Mouse Model**

I herein review findings of MRS studies and immunohistochemical analysis in 2 hypomyelinating mouse models to evaluate neurochemical derangement in hypomyelination.

*Mouse model and MRS methods*

The 2 mouse models of hypomyelination were the msd mouse, a model of connatal PMD with mutation of the *Plp1* gene, and the shiverer mouse with an autosomal recessive mutation of the *Mbp* gene.\textsuperscript{19,20} Both models were subjected to MR imaging and MRS performed on a 7T MR imaging scanner. \textit{T}2 prolongation in the white matter, thalamus, and cortex on \textit{T}2-weighted images reflected hypomyelination.\textsuperscript{19,20} For single-voxel \textsuperscript{1}H-MRS, we chose a region of interest (ROI) predominantly in the thalamus, with volumes of interest of 3.0 \times 3.0 \times 3.0 mm, because the white matter of mice is too thin for ROI placement. We combined outer volume suppression with a point-resolved spectroscopy sequence for signal acquisition (repetition time [TR]/echo time [TE], 2,500/20 ms for msd mice and 4,000/20 for shiverer mice). \textsuperscript{1}H-MRS was quantitatively analyzed using the water scaling method of LCModel, which uses the unsuppressed water signal obtained from the same ROI as an internal reference for quantification (default water proton density [PD] of 35.88 M and \(\exp(-\text{TE/T2}) = 0.7\)).\textsuperscript{21,22} The concentration was corrected by the PD and \textit{T}2 value of the ROI, as multiplied by \(R = \text{PD of ROI}/35.88 \times \exp(-20/T2 \text{ of ROI})/0.7\); however, we used the default PD because we did not measure the PD of the ROI.\textsuperscript{19–22}

**Brain metabolites measured by MRS**

Msd mouse. tNAA, Cr, glutamate, and glutamine were elevated, and Cho was decreased in msd mice (–21.8%) compared with wild-type mice (Fig. 4). There was no difference in mIns. NAAG was not measured because LCModel did not permit acceptable reliability for NAAG.\textsuperscript{19}

Shiverer mouse. Cho was decreased in the shiverer mice compared with heterozygous and wild-type mice. The degree of Cho reduction in the shiverer mice compared with that in wild-type mice (–6.6%) was much less than that in the msd mice. Other metabolites did not differ among the 3 groups. We did not measure NAAG for the same reason as in the msd mice.\textsuperscript{20}

**High performance liquid chromatography (HPLC) measurement of NAA and NAAG in msd mice**

Levels of both NAA and NAAG were higher in the thalamus of msd mice (NAA, 3.52 ± 0.37;
NAAG, 0.66 ± 0.06 µmol/g tissue) than in wild-type mice (NAA, 2.90 ± 0.43; NAAG 0.57 ± 0.07 µmol/g tissue).19

**Immunohistochemical Analysis**

Msd mouse. Immunostaining of Mbp, a marker for mature myelin sheath in msd mice, showed sparse and weak staining in the white matter, thalamus, and cortex compared with that in the wild-type mice.19 Immunostaining of glial fibrillary acidic protein (Gfap), a marker for astrocytes, of the msd mice (white matter and thalamus) showed dense and strong staining and increased positive cells compared with findings in the wild-type mice. These indicate hypomyelination and astrogliosis in the brain of the msd mice. Immunostaining of neuronal nuclear antigen (NeuN), a marker for neuronal cells, showed no obvious difference between the msd and wild-type mice.

Shiverer mouse. Luxol fast blue (LFB) staining and immunostaining of Mbp of shiverer mice revealed sparse and weak staining in the white matter, thalamus, and cortex compared with staining in heterozygous and wild-type mice,20 which indicated hypomyelination in the brains of the shiverer mice. Gfap immunostaining of the shiverer mice showed dense and strong staining and increased positive staining in the white matter, thalamus, and cortex compared with staining in the wild-type mice. Increased tNAA (both NAA and NAAG) in the msd mice (model of PMD) is compatible with MRS in human PMD,13,14 whereas the concentration of tNAA is normal in shiverer mice. NeuN immunostaining of the thalamus in both msd and shiverer mice revealed no difference compared with that in wild-type mice; therefore, the number of neuronal cells in the thalamus seems unlikely to explain the difference in tNAA. These findings suggest that an increase in tNAA is not directly related to hypomyelination.

One possible explanation for the difference in tNAA is the different patterns of oligodendrocytes in shiverer and msd mice (Table). Mutant PLPL proteins in msd mice are abnormally folded and ac-cumulated in the endoplasmic reticulum, which results in the activation of an unfolded protein response that finally leads to apoptotic death of oligodendrocytes before normal myelination occurs.23 In msd and in jimpy, another mouse model of PMD, massive apoptosis of oligodendrocytes appears to induce proliferation of oligodendrocyte progenitor cells (OPC) in the white matter,24,25 which leads to an increase in OPCs and the absence of mature oligodendrocytes (Table). On the other hand, the absence of MBP protein in shiverer mice results in the failure of oligodendrocytes to form a compact myelin sheath. OPCs in shiverer mice, unlike those in msd mice, can differentiate into oligodendrocytes, and both OPCs and mature oligodendrocytes are increased in number by as much as 2 times (Table).26

NAA is one of the most abundant free amino acids in the brain (about 10mM in human) and generally considered an important marker of neurons and axons.1,2 NAA is either released from neurons or transported to oligodendrocytes, where it is catalyzed by ASPA into acetate and aspartate,27 which are used for fatty acid and steroid synthesis, and energy production (Fig. 1). In msd mice, the absence or dysfunction of mature oligodendrocytes may either disable neuron-to-oligodendrocyte NAA transport or affect NAA catalysis in oligodendrocytes, which leads to accumulation of NAA in neurons. An elevated concentration of NAA also increases NAAG biosynthesis,2 which probably results in increased tNAA on MRS, as observed on MRS in patients with PMD.13,14 On the other hand, NAA in shiverer mice may be normally transported from neurons to oligodendrocytes and then catalyzed into acetate and aspartate, leading to normal tNAA.

Cho reduction is much less in shiverer than msd mice, which may also be explained by the different patterns of oligodendrocytes. MRS in vitro demonstrated higher concentrations of Cho in cultured oligodendrocytes per se than neurons, astrocytes, and OPCs.12 The pathological difference in oligodendrocytes may explain the differences in Cho reduction in the 2 strains of hypomyelination, that is, the increased number of oligodendrocytes with se-

**Table. Different patterns of oligodendrocytes in shiverer and msd mice**

<table>
<thead>
<tr>
<th>Oligodendrocyte progenitor cell</th>
<th>Mature oligodendrocyte</th>
<th>Myelin sheath</th>
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<tbody>
<tr>
<td>Shiverer</td>
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vere hypomyelination in shiverer mice,\textsuperscript{26} which leads to a mild reduction of Cho, and the absence of oligodendrocytes with severe hypomyelination in msd mice,\textsuperscript{24} which results in a more severe reduction of Cho.

MRS showed a normal concentration of Cr in the thalamus of shiverer mice, though increased Cr has been observed in msd mice and human patients with PMD.\textsuperscript{9,10,19} Because the elevated Cr was considered to result from the increased number of astrocytes in human PMD, the difference may result from thalamic astrogliosis, which is present in msd\textsuperscript{19} and absent in shiverer\textsuperscript{20} mice. Astrogliosis was recognized in the white matter of shiverer mice, therefore, MRS in the white matter of shiverer mice may have increased Cr. Because mIns is almost exclusively located in astrocytes (astrocyte marker), its increase would also be expected in msd mice, as observed in patients with PMD.\textsuperscript{9,10} Nevertheless, mIns was normal in msd mice compared with wild-type mice. The exact mechanism is uncertain, but markedly decreased mIns and elevated glutamine are observed in patients with hepatic encephalopathy and disorders of the urea cycle.\textsuperscript{28} This has been explained by volume-regulatory mIns release in response to ammonia-induced astrocyte glutamine accumulation. Thus, it is possible that the lack of increase of mIns observed in msd mice may be a consequence of the elevated concentration of glutamine.

Conclusions

The reduction of Cho on MRS might be a common marker for hypomyelinating disorders. Ranges of tNAA from normal to increased probably depend upon the underlying pathology of oligodendrocytes. In hypomyelination, tNAA is increased with reduced numbers of mature oligodendrocytes, such as PMD.

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