Sjögren-Larsson Syndrome: Definitive Diagnosis on Magnetic Resonance Spectroscopy

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Sjögren-Larsson Syndrome (SLS) is a rare autosomal-recessive neurocutaneous disorder comprising a triad of ichthyosis, mental retardation, and spastic diplegia or quadriplegia.1 The disorder was first described by Sjögren and Larsson2 in 1957. Early reports of SLS were mainly in white patients, with a particularly high prevalence of 8.3 cases per 100,000 individuals in the county of Västerbotten in Sweden.3 Reports of SLS in Asian and Indian populations are rare.4,5 We report a case of SLS in an Indian boy.

Case Report
A 12-year-old Indian boy born to nonconsanguineous parents after a full-term pregnancy with normal vaginal delivery presented with generalized dry scaly skin that had been present since 2 months of age. He had a history of delayed milestones (ie, facial recognition, sitting without support at 3 years of age), inability to walk, dysarthria, mental retardation). He had never attended school due to subnormal intellectual functioning. He had a single episode of a tonic-clonic seizure at 4 years of age but was not on any regular antiepileptic medication. There was a history of similar skin lesions in one male sibling of the patient and in 2 maternal uncles. None of them survived beyond early childhood, but detailed information regarding the cutaneous and neurologic manifestations in these family members was not available.

Cutaneous examination revealed lamellarlike ichthyosis on the dorsal aspects of the arms and legs (Figure 1A). Ichthyosis with lichenification was present on the neck, axillae, cubital, and popliteal fossae, and abdomen (Figure 1B). The palms and soles showed keratoderma. Neurologic examination of the arms revealed mild rigidity and brisk reflexes. Examination of the legs showed marked rigidity, brisk knee jerks, ankle clonus, extensor plantar reflex, flexion deformity with contractures, and scissor gait. A Goddard (Seguin) formboard test was performed and indicated a mental age of 4 years. The patient's IQ was in the range of 25 to 30,

FIGURE 1. Sjögren-Larsson syndrome with lamellarlike ichthyosis of the left foot (A) and ichthyosis with lichenification on the trunk and cubital fossae (B).
indicating a severe degree of subnormality in intellectual functioning. The clinical presentation suggested a diagnosis of SLS.

A skin biopsy from the ichthyotic lesion showed hyperkeratosis, acanthosis, and papillomatosis with sparse superficial perivascular lymphocytic infiltrate, thus confirming the diagnosis of lamellar ichthyosis. Fundus examination was normal. Magnetic resonance imaging (MRI) of the brain revealed confluent symmetrical signal abnormalities along the body of the lateral ventricles, white matter in the pericapsular horn, and in deep white matter of centrum semiovale (Figure 2). Magnetic resonance spectroscopy revealed a narrow lipid peak at approximately 1.3 ppm in the region of signal abnormality (Figure 3). Thus, the diagnosis of SLS was confirmed. Measurement of fatty aldehyde dehydrogenase (FALDH) activity and genetic analysis were not performed due to unavailability.

The patient was treated with topical emollients for the ichthyosis. To reduce his dietary intake of long-chain fatty acids and increase the intake of omega-3 and omega-6 fatty acids, the patient’s parents were advised to use canola, mustard, and/or coconut oil for cooking for the patient, and skim milk was recommended instead of whole milk. Neurodevelopmental techniques in the form of stretching exercises were given to maintain his range of movements. Gutter splints were given to maintain the knees in extension for physiological standing and to prevent osteoporosis. Subsequently, the patient also underwent a multilevel soft-tissue release (hip and knee joints) to relieve the contractions. These measures resulted in considerable improvement and the patient was able to walk with support.

Comment

Presentation—The characteristic clinical features of SLS begin to develop during the intrauterine period and infancy.1,6 Pathologic skin involvement can be detected as early as week 23 of gestation. Preterm births associated with SLS have commonly been described.3 Ichthyosis often is evident at birth, but colloidion membrane is uncommon. Severe pruritus is a marked feature unlike most other types of ichthyosis. The ichthyosis often is generalized with prominent involvement of the flexural areas and nape of the neck, varying from fine furfuraceous to larger lamellarlike scales. Velvety orange or brown lichenification often is a predominant feature in the flexures of the arms, legs, neck, and mid abdomen. Mental retardation, developmental delay, and spasticity usually become apparent at 1 to 2 years of age and subsequently are nonprogressive.6,7 However, patients rarely have been described with normal intellectual functioning.7 Spasticity often is more severe in the lower limbs and may lead to contractions, kyphoscoliosis, hip dislocation, and short stature. Delayed speech and dysarthria are common. Parafocal glistening white dots on the retina are a pathognomonic feature and typically appear in the first 2 years of life; however, they are seen in approximately 30% of patients and increase slightly in number with age.6,8 There may be associated decreased visual acuity, photophobia, myopia, and astigmatism. Other clinical features include enamel hypoplasia, metaphyseal dysplasia, and epilepsy.1,6

Gene Mutations—Sjögren-Larsson syndrome is caused by mutation in the aldehyde dehydrogenase 3 family member A2 gene, ALDH3A2 (17p11.2), which codes for FALDH.1,5,6 The ALDH3A2 gene is 11 exons long and gives rise to 2 protein isoforms that differ in their carboxy-terminal domains; the major isoform, composed of 485 amino acids, localizes to the endoplasmic reticulum. The minor protein isoform (FALDHv) is composed of 508 amino acids, possesses a longer carboxy-terminal, and appears to be targeted to the peroxisome. Several mutations have been reported throughout the ALDH3A2 gene, including missense mutations (most common [38% of cases of SLS]), deletions, insertions, splicing errors, and complex rearrangements. Although several of these mutations are private, several common mutations may be indicative of founder effects (ie, shared ancestry), consanguinity, or recurrent mutational events (mutation hotspots).6,7 Despite the wide spectrum of mutations, there is very little phenotypic variation, with consistently severe cutaneous and neurological involvement occurring in a majority of patients. However, Lossos et al9 described remarkable phenotypic variation in 6 siblings of an Arab family and suggested that additional unknown genetic or environmental factors may compensate for the biochemical defect.

Lipid Metabolism—Fatty aldehyde dehydrogenase is expressed in almost all cells and tissues and catalyzes the oxidation of fatty aldehydes to fatty acids (Figure 1). It also is a part of the fatty alcohol:NAD oxidoreductase (FAO) enzyme complex, which catalyzes fatty alcohol oxidation to fatty acid. Fatty aldehyde dehydrogenase deficiency leads to accumulation of long-chain alcohols (eg, hexadecanol, octadecanol, octadecenol) and diversion of fatty alcohol into alternate biosynthetic pathways such as wax esters and 1-O-alkyl-2,3-diacylglycerol.10 Other lipids that...
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**Histopathology**—The epidermal permeability barrier is critically dependent on the appropriate lipid composition of the multilamellar stratum corneum intercellular membranes, an equimolar ratio of cholesterol, ceramides, and fatty acids. Histopathology of the skin in SLS generally shows hyperkeratosis, papillomatosis, acanthosis, and mildly thickened granular layer. Ultrastructural studies of the skin reveal misshapen/empty lamellar bodies, abnormal cytoplasmic lamellar inclusions in the granular keratinocytes, lipid droplets in the stratum corneum with decreased lamellar bilayers, and lamellar/nonlamellar phase separation in the stratum corneum interstitium. These findings indicate that lipid metabolism dysfunction in SLS results in marked impairment in formation and secretion of lamellar bodies in the epidermis and consequent disorganization of the stratum corneum lamellar membranes. The resulting disruption of the skin barrier function leads to increased transepidermal water loss, resulting in ichthyosis. Another proposed mechanism for ichthyosis in SLS is disruption of the normal epidermal differentiation resulting from abnormal lipid metabolites (eFigure 2). Also, increased leukotriene B4 (LTB4) and 20-hydroxy-leukotriene B4 (20-OH-LTB4) (eFigure 1) may be responsible for the classical cutaneous, neurologic, and ophthalmologic features of SLS.

**Neurologic Changes**—Neurologic changes in SLS result from delayed and deficient myelination. Neuropathological studies have shown ballooning of myelin sheaths, extensive loss of myelin, axonal damage, and astrogliosis. The presence of lipoid material positive for periodic acid–Schiff that stains light rather than dark pink, dense distribution of round/ellipsoid bodies in the white matter of the cerebrum and brainstem positive for periodic acid–Schiff, and proliferation of perivascular macrophages containing lipofuscinlike pigments also have been described. Possibly, in the absence of FALDH, metabolism of plasmalogens (a major component of myelin) results in increased fatty aldehydes, which are either diverted to fatty alcohols or form adducts with phosphatidylethanolamine and myelin basic proteins (eFigure 1). Magnetic resonance imaging of the brain usually shows hypomyelination involving the periventricular white matter extending from the frontal to the occipital area. Mild ventricular enlargement may be an additional feature.

A useful application of MRI is the proton magnetic resonance spectroscopy, which quantifies the brain metabolites noninvasively, displaying them as a spectrum on a graph. The spectrum comprises a set of resonances/peaks distributed along an x-axis. The resonances of these metabolites are obtained after suppressing the large signals from water protons. Proton magnetic resonance spectroscopy of the normal brain shows 3 prominent peaks: (1) N-acetylaspartate (NAA) at 2.02 ppm, (2) creatine at 3.02 ppm, and (3) choline at 3.22 ppm. In SLS, cerebral proton MRI spectroscopy reveals a characteristic abnormal, prominent, and narrow lipid peak at 1.3 ppm (corresponding to hexadecanol and octadecanol) and may offer a quantitative parameter for monitoring the effects of therapeutic interventions. The most intense lipid peaks are located in the periventricular regions in the anterior and posterior tringes. An abnormal but much smaller peak may be seen at 0.8 to 0.9 ppm, corresponding to phytol. Proton magnetic resonance spectroscopy also can be used for screening of SLS heterozygotes. Lipid peaks have been described in other disorders of lipid metabolism, but they are less intense, broader, and disappear on longer echo time sequences.

Besides the characteristic parafoveal glistening white dots the retina, optical coherence tomography shows focal hyperreflectivity in the perifoveal ganglion cell layer and inner plexiform layer of the retina as well as cystoid foveal degeneration. The intraretinal deposition of lipid metabolites probably leads to Müller cell degeneration with subsequent formation of cystoid spaces and atrophic changes in the fovea.
Measurement of FALDH or FAO activity in cultured skin fibroblasts and leukocytes using fluorometric or gas chromatography mass spectrometry assays is a reliable biochemical test in cases of SLS as well as in heterozygotes. A decrease in FALDH/FAO activity also can be demonstrated by histochemical staining in skin biopsy. Pathologic urinary excretion of LTB4 and 20-OH-LTB4 also is a biochemical marker of SLS. Mutation analysis for a specific gene defect is diagnostic in cases of SLS as well as in heterozygotes. Prenatal diagnosis of SLS is possible by assessing FALDH activity or gene defects in cultured chorionic villus fibroblasts and amniocytes.

**Differential Diagnosis**—The differential diagnosis of SLS includes congenital ichthyosiform erythroderma with neurological signs (Tay syndrome, Conrad-Hünermann-Happle syndrome) and neurocutaneous disorders such as neutral lipid storage disease and multiple sulfatase deficiency; however, the nature of the ichthyosis, presence of spastic diplegia/tetraplegia, characteristic parafoveal glistening white dots on the retina, and MRI and MRTI are helpful in differentiating these disorders.

**Treatment**—Treatment of SLS mainly is palliative. Ichthyosis can be treated with topical keratolytics, emollients, calcipotriol, and oral retinoids (acitretin). Zileuton, a 5-lipoxygenase inhibitor, inhibits synthesis of LTB4 and cysteinyi leukotrienes, thereby reducing the severity of pruritus and also has been shown to improve the speed of information processing. Similarly, montelukast, a leukotriene antagonist, is helpful in relieving the agonizing pruritus. Experimental studies have shown that bezafibrate, a peroxisome proliferator-activated receptor α agonist, induces FALDH activity in fibroblasts of SLS patients that still have some residual FALDH activity; further research is required to determine whether SLS patients could benefit from treatment.

Physiotherapy helps in relieving the spasticity to some extent, such as in our case. Dietary intervention with reduced fat intake (up to 30% of total daily caloric requirement) and supplementation with omega-3 and omega-6 fatty acids has shown variable results in anecdotal reports. Gene therapy using recombinant adeno-associated virus vectors to restore FALDH has been projected as a future treatment option. Despite lack of effective treatment options, most patients of SLS survive well into adulthood.

**Conclusion**

Because ichthyosis is one of the earliest and prominent symptoms of SLS, a dermatologist can play an important role in early diagnosis. Any child with the classical pattern of ichthyosis should be thoroughly examined for early neurologic signs and investigated to rule out SLS. Proton magnetic resonance spectroscopy serves as a useful adjunct in the diagnosis of SLS by confirming the accumulation of abnormal lipids in the periventricular white matter, especially when specific enzyme analysis and genetic analysis are not available in resource-restricted settings.

**REFERENCES**

eFIGURE 1. Role of fatty aldehyde dehydrogenase (FALDH) in lipid metabolism in Sjögren-Larsson syndrome. Fatty aldehyde dehydrogenase is responsible for oxidation of fatty aldehydes derived from long-chain alcohols, branched alcohols, isoprenoid alcohols, and ether glycerolipids. Fatty aldehyde dehydrogenase also is necessary for \( \omega \)-oxidation of leukotriene B4 (LTB4) and epoxyalcohols (trioxillen A3). Lipid metabolites shown in italics are increased in Sjögren-Larsson syndrome.
**eFIGURE 2.** Lipid pathways affected in Sjögren-Larsson syndrome, consequent lipid abnormalities, and resultant pathogenic effects. HMG-CoA indicates 5-hydroxy-3-methylglutaryl-coenzyme A; PPARα, peroxisome proliferator-activated receptor α; LTB4, leukotriene B4.