**Magnetic Resonance Imaging (MRI) findings in White Matter Disease of Brain**

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**Abstract**

Demyelinating and dysmyelinating white matter diseases are important components of neurological problems. Recently, Magnetic Resonance Imaging (MRI) has played a key role in diagnoses of white matter diseases. Therefore, the purpose of the current study is to evaluate the usefulness of MRI in determining the type and frequency of white matter disease.

We studied 35 patients who visited the Radiology Department of the Aga Khan University Hospital (AKUH) for MRI with suspected demyelinating/dysmyelinating disorder from January 2003 to December 2005. Multiple Sclerosis (MS) (17; 48%) and leukodystrophies (10; 29%) were the most common diseases.

The MRI helped identify the sites and types of the lesion precisely and thereby helped made clearer distinction between various types of white matter diseases. The current study demonstrated the effective use of the imaging and clinical presentation for arriving at the correct diagnosis.

**Introduction**

Demyelinating and dysmyelinating white matter diseases are important components of neurological problems. White matter diseases include Multiple Sclerosis (MS), Leukodystrophies, Central Pontine Myelinolysis, Acute Disseminated Encephalomyelitis and Progressive Multifocal Leukoencephalopathy. MS is the commonest of all the white matter diseases. Leukodystrophies, which include Disseminated Encephalomyelitis (ADEM), Central Pontine Myelinolysis and Progressive Multifocal Leukoencephalopathy, are ranked second.

With the ever rising burden of the white matter disease, it is imperative to better understand pathophysiological, clinical and imaging findings for proper management of patients. The advent of Magnetic Resonance Imaging (MRI) has revolutionized the concept of diagnosing and understanding of white matter diseases.

Many of the white matter diseases, if detected early, can be cured and MRI plays a vital role in its early diagnosis. MRI is considered far superior to Computed Tomography (CT) and the imaging modality of choice in white matter diseases. CT does not detect subtle lesions especially in stages of clinical inactivity and is not ideal in posterior fossa imaging due to the beam hardening artifacts. The multiplanar imaging capability and very high sensitivity for demyelinating foci due to its excellent gray-white matter resolution make MRI imaging the modality of choice. Through MRI, simultaneous imaging of spinal cord and orbits can also be done. Moreover, with the advent of multi-echo sequences of MR, even subtle lesions of demyelination can be detected.

The purpose of the current study is to evaluate the frequency of white matter disease among suspected cases of demyelinating disorders in our setting. Further the study will also look into use of MR imaging features in diagnosis of white matter disease and relate it with clinical findings.

**Patients, Methods and Results**

We conducted a descriptive study of the patients who had their MRI brain done with clinical suspicion of demyelinating disorder at the Radiology Department of The Aga Khan University Hospital (AKUH) from January 2003 to December 2005.

Forty eight patients visited the department during the study period. However, thirteen were excluded due to prior diagnosis of white matter disease and/or incomplete medical records. Thirty five patients were finally included in the study. The MRI examination features of these patients were highly suggestive of demyelinating or dysmyelinating diseases.

All the patients were examined with 1.5 Tesla systems (Toshiba Medical System). Head coil was used in all the patients. A combination of T1 and T2 weighted and FLAIR sequences and post gadolinium T1 weighted was obtained in each patient.

Each MR image was interpreted by two Radiologists proficient in reporting brain MRI. In case of difference of opinion, consensus was developed through discussion between the two radiologists.

The diagnosis was confirmed by laboratory findings of CSF oligoclonal band in patients with MS. All cases of acute disseminated encephalomyelitis and leukodystrophies, the diagnosis was established through typical imaging findings and clinical course without any relapse. MRI
Our study included 35 patients of white matter diseases: 14 (40%) males and 21 (60%) females. The majority of patients suffered from MS 17 (48%): 10 (58.8%) females and 7 (41.2%) males with average ages of 32 and 31 years respectively. MRI revealed multiple lesions confined to the white matter appearing hypo-intense on T1W and hyper-intense on T2W images. In all the patients at least three or more lesions were seen. The majority of the lesions were less than one cm. in size. The commonest sites of involvement were the periventricular and pericallosal area (Figures1a and b). The other sites were centrum semiovale, deep white matter and sub cortical white matter of left frontal lobe, parietal lobe, parieto-occipital region, left temporal lobe, cervical spinal cord, rightpons and posterior part of left optic nerve.

Leukodystrophies, which include Metachromatic Leukodystrophy and Adrenoleukodystrophy, Alexander's and Niemen Pick Diseases, were the other most common group of white matter diseases. The results showed that 10 (28.6%) patients, 7 (70%) males and 3 (30%) females, suffered form leukodystrophies. Each type of leukodystrophy presented with different frequencies and MRI findings.

Three patients, all males with an average age of 4 years, suffered from metachromatic leukodystrophy. The commonest site was periventricular white matter. On the other hand, adrenoleukodystrophy was seen in three patients. The average age was three years. The abnormal signals of deep periventricular white matter specifically in trigonal area with ventriculomegaly were observed in two out of three patients.

Leigh disease was diagnosed in two cases. The MRI showed hyperintense signal on T2 weighted images in deep grey matter nuclei, tegmentum, thalami, periventricular white matter, periequiductal region and dentate nuclei.

One patient showed signs of Alexander's disease. The MRI showed periventricular hyperintensity in the frontal lobe with post contrast enhancement and a cyst in the third ventricle.

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One patient showed signs of Alexander's disease. The MRI showed periventricular hyperintensity in the frontal white matter with a cyst in the third ventricle and post contrast periventricular enhancement. Similarly, one patient had Niemen Pick Disease. MRI showed Gliosis in the frontal and temporal lobes.

The Central Pontine Myelinosis was ranked third among the white matter diseases identified. Three (8.5%) patients, two males and one female, had the disease. T2 signal changes were seen in the pons in all cases on MRI.

One male patient suffered from Progressive Multifocal Leukoencephalopathy. He was 30 years old and tested positive for Human Immunodeficiency Virus (HIV). MRI showed abnormal signal intensity in deep white matter of brain in right frontal regions, parietooccipital regions, putamen and midbrain. The signals were isointense on T1 and hyperintense on T2 weighed images.

Acute Disseminated Encephalomyelitis was identified in four patients. T2 weighted and FLAIR images...
showed asymmetrical hyperintense signals in the subcortical white matter of frontal, parietal and occipital lobe and cerebellum.

**Conclusion**

The current study presents the proportion of patients suffering from each type of white matter disease. It also demonstrates the effective use of the imaging and clinical presentation in making the right diagnosis.

White matter diseases are a heterogeneous group of disorders, where the main imaging finding is the abnormal white matter. Although a clinical diagnosis is usually present in these disorders, imaging reinforces the diagnosis in many cases. Thereby, MRI imaging could play a pivotal role in prompt treatment of patients with white matter disease.

**References**


**Short Report**

**An open study to assess the safety and efficacy of Heprovac-B vaccine 10 mcg-dose for adults**

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**Abstract**

Heprovac B is a novel recombinant vaccine. There are many vaccines available in the Pakistani market but Heprovac B claims to be immunogenic even at 10 mcg dose. Aim of the study is to determine whether using 10 mcg of Heprovac B vaccine is safe and effective in producing sufficient immunity in the Pakistani population. One hundred and twenty-five subjects, who fulfilled the inclusion criteria, were enrolled for the study. Heprovac B was administered in a three-dose regimen given at 0, 1 and 6 months and adverse events were recorded. Immunogenicity was tested by measuring hepatitis B surface antibody one month after each dose received. One month after the third dose, 98.7% of the subjects were found to be seroprotected with geometric mean titer of 488.83 mIU/1 after the third dose. Heprovac B vaccine was well tolerated with minimal reported adverse events. It is safe and 10mcg is immunogenic in producing antibodies in the Pakistani population against Hepatitis B virus.

**Introduction**

Hepatitis B virus infection is a major healthcare issue that affects children and adults worldwide. Long term infection can lead to chronic hepatitis, cirrhosis and primary hepatocellular carcinoma. In Pakistan, Hepatitis B is one of the major causes of chronic liver disease and hepatocellular carcinoma. In Pakistan, the estimate is 4.5 million carriers, with a carrier rate of 3.4%. Since, there is no definite treatment available yet, great emphasis is being placed on its prevention through immunization. Currently there are yeast derived vaccines present in Pakistan. The immunogenicity of these vaccines has been subjected to intense study since their birth.

There are three forms of HBsAg that are expressed in the hepatitis B vaccine, namely the major form consisting of the S protein alone, the large form containing the preS1 - preS2 - S components and a middle form containing preS2 - S components. The presence of these conforms to the Immunogenicity of the vaccine. The yeast cell recombinant vaccines contain only the major form i.e. only protein S; hence chances of failure of sero-conversion in these vaccines does exist.

The CHO cell derived vaccine, on the other hand,