Metabolic disease in Nepal: A perspective

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ABSTRACT
Inborn errors of metabolism or metabolic diseases, are a group of genetically determined metabolic disorders that result in mental retardation or early death. The prevalence of IEM in various countries shows a prevalence varying between 1 in 800 to 1 in 5000. As the technology for detecting metabolites has become more advanced, studies utilizing more modern methods report a higher prevalence. There have been reports of a few Inborn errors of metabolisms in Nepal, but studies to gauge the prevalence of these disorders in the Nepalese population are lacking. With conflicting statistical numbers from different sources regarding mental retardation cases in Nepalese population and a substantial rate of consanguinity and inter caste marriages, it would be prudent to initiate some pilot studies to estimate the prevalence of a group of disorders that can be diagnosed through simple laboratory tests, to be followed by a screening programme depending upon the results. The presented review discusses the need for and the possibilities of screening for these errors for early intervention in Nepal.

Key Words
consanguinity, genetic disease, inborn errors, metabolic disease, mental retardation

INTRODUCTION
Molecular basis of Inborn Errors of Metabolism (IEM) / Metabolic Disease. The term “Inborn Errors of Metabolism” (IEM), represents a group of inherited disorders that result in the impaired activity of an enzyme, a structural protein or a transporter molecule. The underlying causes of IEMs are mutations in genes that code for proteins, resulting in a dysfunctional or structurally altered protein, causing a block in a metabolic pathway. These dysfunctional proteins may be the enzymes required for the metabolism of carbohydrate, proteins or lipids, resulting in disorders associated with altered or blocked metabolism of these biomolecules. This results in the accumulation of abnormal metabolites proximal to the block or the lack of products of the reaction that is blocked. For example

A ➔ B ➔ C ➔ D

A block due to the deficiency of the enzyme converting B to C will result in excess of B and a lack of C, either of which can manifest as disease. A restriction of the precursor of B (i.e. A in the above example) or a supply of the product C, are potential therapeutic interventions. Us, about half of the over about a thousand of known metabolic disorders can be treated biochemically. The medical consequences of IEMs vary from a failure to thrive to acute illness that can lead to brain damage, coma and death. Some examples of these enzymatic deficiencies associated with common metabolic disorders are shown in Table 1.
Table 1. Selected Metabolic disorders with major associated symptoms

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Enzyme Deficiency</th>
<th>Symptoms</th>
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<tbody>
<tr>
<td>Disorders of Carbohydrate Metabolism</td>
<td></td>
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<tr>
<td>Galactosemia:</td>
<td>Galactose 1 P uridyl transferase</td>
<td>Liver failure, mental retardation</td>
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<tr>
<td>Mucopolysaccaridoses:</td>
<td>Glycosaminoglycans degrading enzymes</td>
<td>Accumulation of GAGS in tissues resulting in hepatomegaly/splenomegaly/disturbances of growth/coarse facies/mental retardation</td>
</tr>
<tr>
<td>Glycogen storage disorders</td>
<td>Enzymes of glycogen metabolism</td>
<td>Deposition of abnormal type or quantity of glycogen in tissues, liver or heart failure</td>
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<td>Disorders of amino acid metabolism</td>
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<td></td>
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<tr>
<td>Phenylketonuria</td>
<td>Phe Ala hydroxylase</td>
<td>Retarded development and neurological abnormalities</td>
</tr>
<tr>
<td>Hyperglycinemia</td>
<td>Glycine cleavage enzyme</td>
<td>Failure to suck, coma, myoclonic jerks, Ochronosis</td>
</tr>
<tr>
<td>Homocystinuria:</td>
<td>Cystathione beta synthase</td>
<td>Thrombosis, osteoporosis, dislocated lens, mental retardation</td>
</tr>
<tr>
<td>Maple syrup urine disease:</td>
<td>alpha ketoacid decarboxylase</td>
<td>Development delay, convulsions in later infancy, low IQ</td>
</tr>
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<td>Tyrosinemia:</td>
<td>Fumarylacetoacetate hydrolase</td>
<td>Diarrhoea, vomiting, “cabbage like” odor, liver failure, renal tubular dysfunction</td>
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<td>Organic acidemias</td>
<td></td>
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<tr>
<td>Methylmalonic acidurias</td>
<td>Methylmalonyl CoA mutase/ Vit. B12</td>
<td>Severe acidosis, multiple organ failure</td>
</tr>
<tr>
<td>Propionic academia</td>
<td>Propionyl CoA carboxylase</td>
<td>Metabolic acidosis, seizures, coma</td>
</tr>
<tr>
<td>Isovaleric academia:</td>
<td>Isovaleryl CoA dehydrogenase</td>
<td>“sweaty feet” odor, seizures, coma</td>
</tr>
<tr>
<td>Glutaric academia Type I</td>
<td>Glutaryl CoA dehydrogenase</td>
<td>Spasticity, dystonia, seizures, coma</td>
</tr>
<tr>
<td>Urea cycle disorders:</td>
<td>Enzymes of Urea Cycle</td>
<td>Hyperammonemia, encephalopathy, coma, death</td>
</tr>
<tr>
<td>Hyperammonemias</td>
<td>Ornithine carbamoyl transferase</td>
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<td></td>
<td>Argininosuccinate synthase</td>
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<td>Argininosuccinate lyase</td>
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<td></td>
<td>Arginase</td>
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<tr>
<td>Fatty acid oxidation disorders</td>
<td>Medium chain acyl CoA dehydrogenase</td>
<td>Acidosis, hyperammonemia, hypoketotic hypoglycemia, encephalopathy, hepatomegaly, microvesicular fatty infiltration of the viscera</td>
</tr>
<tr>
<td>Lysosomal storage disorders</td>
<td></td>
<td></td>
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<tr>
<td>Tay-sachs disease:</td>
<td>Hexosaminidase A</td>
<td>Hepatomegaly, mental and neurological deterioration, death</td>
</tr>
<tr>
<td>Pompe's disease:</td>
<td>Lysosomal alpha 1-4 or alpha 1-6 glucosidase</td>
<td>Cardiomegaly and cardiac failure</td>
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PREVALENCE

The majority of metabolic disorders have autosomal recessive inheritance, a few though, show X-linked inheritance. The frequency of a selected birth in a population with random (non-consanguineous) mating may be calculated from the frequency of carriers by using the formula:

\[
\text{Frequency of affected births} = 0.25 \times (\text{frequency of heterozygotes})^2.
\]

Frequency of a selected birth in a population with random mating can be significant differences in the incidence of individual inborn errors of metabolism. In areas where there is a high rate of consanguineous marriage, there will be higher incidence of metabolic disease overall.

Although individual inborn errors of metabolism are relatively rare conditions, as a group they represent a vast and diverse collection of diseases that are a significant cause of morbidity and mortality worldwide. Even though reports in the literature often quote a cumulative incidence varying between 1 in 1500 and 1 in 5000 live births, a recent retrospective study on an ethnically diverse population in the United Kingdom found this range to underestimate the real figure, and placed the prevalence of inherited metabolic disorders at 1 in 784 live births. Data for the prevalence of IEMs in South Asia is starting to come in. A study undertaken in India indicates an incidence of 1 in every 1000 newborns. Another study from Pakistan that tested undiagnosed children <1-10 years showing neurological symptoms, development delay and vomiting, found 26% of the children to have one of the tested IEMs. A similar study in China found 48.6% of the 4981 pediatric patients suspected of metabolic disease to have aminoacidurias, organic acidemias or fatty acid disorders. The measured incidence in a particular area depends on the methods used for screening, with older and less sensitive biochemical methods showing a much lower incidence compared to the methods employing the more specific tandem mass spectrometry.
In Nepal, Wilson's disease, an inborn error of copper metabolism has been documented twice as case reports. Although studies have been conducted on the incidence of chromosomal disorders, inborn errors of metabolism, probably because of their rarity, have not caught a mention. There has been a case report on -thalassemia along with earlier studies done in the 1960s, that found some cases of -thalassemia, a few cases of the presence of abnormal hemoglobins H and E, and Glucose 6 phosphate dehydrogenase deficiency in Nepal.

DIAGNOSTICS

In many countries, newborn screening for a few inherited metabolic diseases like hypothyroidism, phenylketonuria and sickle cell disease is a routine part of neonatal care. The primary aim of neonatal screening is the early detection and treatment of clinically important disorders in order to minimize morbidity and mortality. Infants with inborn errors of metabolism appear normal at birth as most of the accumulating abnormal metabolites can cross the placenta and can be cleared by the mother. Initial findings are usually those of lethargy and poor feeding, seen in almost any sick infant. An infant in whom these go unnoticed may come to attention because of apnea or sudden respiratory distress. Signs of Central Nervous System dysfunction, such as seizures and abnormal muscle tone, may also be noted. A clinical diversity that these disorders present makes it difficult to recognize them clinically, with specific diagnosis usually those of lethargy and poor feeding, seen in almost any sick infant. An infant in whom these go unnoticed may come to attention because of apnea or sudden respiratory distress. Signs of Central Nervous System dysfunction, such as seizures and abnormal muscle tone, may also be noted.

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IEMs in the Nepalese context

Mental Retardation

Screening of newborns for metabolic disorders is carried out in many countries in Europe, America and Asia. It includes screening for phenylketonuria, cystic fibrosis, hypothyroidism (though not strictly a metabolic disorder) and thalassemias. Technological advances in diagnosis led to identification of more of these disorders, which had escaped detection with older methods.

In Nepal, the earliest study done to detect IEMs was a part of a larger study carried out in India, Bangladesh, Pakistan, Nepal, Bhutan and Sri Lanka. A study limited itself to hemoglobinopathies and related disorders, and the Nepalese population that was tested comprised of Gurkhas and Sherpas residing outside Nepal. One in approximately 130 screened Nepalese were found to be thalassemia trait carriers. In a 2009 study done at BPKIHS genetic clinic, 10 of 30 children with mental retardation, dysmorphic features, short stature and ambiguous genitalia, were found to have chromosomal disorders. A study concluded that in the remaining 20 children, either the aberrations were undetectable by the available technology, or could have been the result of other single gene disorders. It is reasonable to speculate that some of these undiagnosed cases of mental retardation could have been the result of metabolic disorders. A survey of self-reported disability in Eastern Nepal found a rate of disability of 4.87%, of which 17.1% is reported to be a result of an “inborn syndrome”. It is logical to assume that these “inborn syndromes” are a combination of cytogenetic and metabolic disorders.

In Nepal, the country health system profile shows a prevalence of disabilities of 1.63% out of which 5.9% are mental retardation. A 1989 survey done by "Maryknoll
Father’s Project” found a prevalence of mental retardation of 4.1% amongst Nepalese population. A survey of two developing towns in western Nepal in 1998 revealed a high point prevalence (35%) of “conspicuous psychiatric morbidity”. According to Dr. Kan Tun, a former World Health Organization representative to Nepal, around 1% of the Nepalese population has severe mental illness and 10-20% milder mental health problems. A population of around 29 million hence puts the number of severely mentally ill at 290,000, and mildly mentally ill at 2,900,000 to 5,800,000. It is again logical to assume that a considerable number of these estimates might be the surviving cases of inborn errors of metabolism, most of which result in mental retardation.

Consanguinity

It is established that the incidence of inborn errors is higher in areas with higher rates of consanguinity. Anthropological studies undertaken by Fricke and his colleagues in the Timling region of Nepal (along Tibet and Burmese borders) report that cross cousin marriages are common and part of the social structure that is strongly based on kin alliance with reciprocal rights and obligations. In rural areas in Nepal where farming and manual labour are predominant occupations, cross cousin marriages also have an economic value as the boundaries of endogamy are shaped by family allegiances and reciprocal rights and obligations. Amongst the Hindus of Nepal, the common form of consanguineous marriages is patrilateral cross cousin marriages. Endogamy has been prevalent in communities like Magars, Gurungs, Tamangs and others where the members of the communities marry among their own near relations. The system was also adopted among the ruling families of Shivas and Ranas.

Caste system and recessive disease

Populations arising from “founder events” have a limited gene pool to begin with. For example, in a recent study on Indian population history, it was found that two ancient populations, genetically divergent, are ancestral to most Indians today. One the Ancestral North Indians (ANI) is genetically close to Middle Easterners, Central Asians and Europeans, whereas the other the “Ancestral South Indians” (ASI) is as distinct from ANI and East Asians as they are from each other. Allele frequency differences between groups reflect strong founder events whose signatures have been maintained for thousands of years owing to endogamy. Authors hence predicted that there will be an excess of recessive diseases in India, which should be possible to screen and map genetically.

They proposed that founder events are responsible for an even higher burden of recessive diseases in India than consanguinity. Political scientists Joshi and Rose broadly classify the Nepalese population into three major ethnic groups in terms of their origins: Indo-Nepalese, Tibeto-Nepalese and indigenous Nepalese. The genetic diversity of populations inhabiting an area is also influenced by the geographic and physical features encompassing the region. Whereas the Hindu Kush Mountains and the arid deserts in Iran have served as obstacles to gene flow, the Nile River Valley, the strait of Babel Mandeb and Beringia are examples of natural passageways for the migrations of modern humans. The Himalayan range, in addition to being a formidable barrier, provides for dramatically diverse climatic conditions on either side of it. An investigation of the genetic affinities of Newar, Tamang and people from cosmopolitan Kathmandu and Tibet suggests that the Tibetans and Nepalese are in part descendants of Tibeto-Burman speaking groups originating from Northeast Asia. With the exception of Tamang, both Newar and the people of cosmopolitan Kathmandu exhibit considerable similarities to the Indian Y haplogroup distribution.

Although the conclusion from the above can only be implied, the gene pool of the Nepalese populations residing in various geographically distinct areas was probably limited, both due to founder events as well as the presence of the Himalayas as a barrier to gene flow. These populations have over time, come to be known as different castes or Janajatis.

deletion of caste system most relevant to Nepal is “castes are ranked endogamous divisions of society in which membership is hereditary and permanent.” Nepali caste rules normally prescribe isogamy for its members. Such marriages are held lawful for the inheritance of property by the offspring and for ensuring ritual purity of a caste-member. Caste endogamy is thus held sacrosanct because heredity is basic to the concept of caste-purity. Newar have a highly structured caste system, which the Malla kings dictated over 600 years ago. Although old caste restrictions on occupation are gradually fading, the social restrictions of the caste system are still largely observed and the Newar rarely marry outside their caste. The caste system has persisted in Indian Hindu society for around 3500 years. Like the Y chromosome, caste is determined at birth, and males cannot change their caste.
predominance of a single cluster of haplotypes in India constitute the genetic isolation and drift within the Jaunpur upper castes which are likely to result from founder effects and social factors. John Burdon Sanderson Haldane (1892-1964), one of the founders of population genetics, has commented in one of his essays that if intercaste marriages in India became common, various recessive characters will become rarer.

Diagnosis. To begin with, the health community in Nepal needs to assess the importance of inborn errors of metabolism in terms of infant mortality, incidence of mental retardation due to IEMs, time and resources. An assessment of incidence of IEMs anywhere is difficult to diagnose unless a screening system is in place. As is because most IEMs are difficult to diagnose unless being evaluated by someone trained in IEM diagnosis and care, so that most cases that go undiagnosed result in irreversible mental retardation or death. In a retrospective study on known cases of inborn errors of metabolism in a pediatric intensive care unit in India, 36% of the infants expired, 45% improved and 36% progressed to sequelae. However, an initial attempt at knowing the incidence of inborn errors in Nepal can be made by screening known cases of mental retardation. is can be done by a series of simple biochemical tests on blood and urine. Such tests were developed over a period of years and have been used for diagnosis since 1962. In 1972, Guthrie and his colleagues extended their Phenylketonuria test to a multiple screening program for several inborn errors of metabolism. These tests can be carried out in a biochemical laboratory using coloring agents, paper chromatography and enzymatic methods.

Once an estimate of the number of cases of mental retardation resulting from inborn errors of metabolism is known, an informed decision can be made about whether screening newborns for IEMs to calculate the incidence per 1000 newborns would be economically and ethically feasible. It would also need to take into account the known number of cases where there were the classic symptoms of metabolic encephalopathy in the newborn followed by death. A regular screening programme to screen for metabolic disorders prevalent in Nepal will meet the criteria set by the Wilson's and Jungner's principles of screening for disease. These principles state that

- The condition sought should be an important health problem to the individual and/or the community: So far, the focus of various health organizations in Nepal has been on infectious diseases so that diseases causing disabilities have taken a backseat. The problem, when occurring in an individual is devastating to the family as well as to the child anywhere, and not just in Nepal.

- There should be an accepted treatment for patients with recognized disease. It is axiomatic that case finding should only be undertaken when the prospects for treating the condition are at least reasonable. - Most inborn errors of metabolism are treatable with changes in diet. - The knowledge and skill for treating inborn errors of metabolism is negligible in Nepal. A system to begin training individuals for tackling IEMs should be initiated as soon as possible.

- Facilities for diagnosis and treatment should be available: Diagnosis can be carried out with simple biochemical tests. However, treatment, as mentioned above, will require education and training.

- There should be an agreed policy on whom to treat as patients (to be tested)

- The costs of case finding should be economically balanced in relation to possible expenditure on medical care as a whole: Case finding using simple biochemical tests will be far cheaper than providing for and treating a retarded child, or facing the devastating effect of losing a child on a family. Case finding should be a continuous process and not a “once and for all” project. (to be tested)

TREATMENT

It is widely held in Nepal that inborn errors of metabolism are not treatable. A misconception might also be responsible for the hesitation on the part of medical professionals in pursuing the diagnosis of these conditions. Older, but quite effective methods of treatment of
IEMs require dietary management in order to restrict the formation of or accelerate the removal of the toxic metabolite. Administration of vitamins helps increase the activity of certain enzymes responsible for a particular metabolic disease. Advances in biotechnology have made it possible to produce in laboratories the actual enzymes that are deficient in order to administer them to patients. There are also new strategies in development, involving, for example, recovery of residual enzyme activity using chaperones, cell therapy and gene therapy and also combination treatments. A detailed description of treatment of different types of inborn errors is beyond the scope of this article. The reader is referred to some reviews for further reading on treatment of inborn errors of metabolism.

CONCLUSIONS

Keeping in view the incidence of mental retardation, prevalence of consanguinity in some areas of Nepal and marriage within a particular caste, it is prudent to assume that there are substantial cases of inborn errors of metabolism occurring in Nepal, which remain undiagnosed due to a lack of concern towards this group of disorders. The results of these disorders are either irreversible retardation or death, both of which are devastating to any family, rich or poor. Studies to gauge the burden of this group of disorders on Nepalese society are in order. Simultaneously, biochemists and pediatricians need to be trained in the diagnosis and treatment of inborn errors by way of preparation to be ready when such cases are encountered.

RECOMMENDATION

It is time for Nepal health professionals and authorities to consider the possibility of existence of inborn errors of metabolism in the Nepalese population. In order to gauge the prevalence of metabolic disease, existing cases of mental retardation should be tested biochemically for the presence of known metabolites of IEMs. A true scenario however can only be obtained on screening newborns all over Nepal. An initiative should be taken in Kathmandu using simple biochemical methods. For this purpose biochemists and pediatricians need to be trained in the field of metabolic disease in order to have an established system so that the possibility of early diagnosis and intervention is enhanced.
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