Study to identify the role of high performance liquid chromatography in detecting haemoglobinopathies in antenatal patients

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Abstract
Introduction: Haemoglobinopathy is a major genetic public health problems in India, responsible for significant morbidity and mortality. Individuals with trait (carriers) are healthy and unaware of their carrier status unless specifically screened. If a couple carry a clinically significant haemoglobinopathy trait there is a 1 in 4 chance with each pregnancy that their child will inherit a major haemoglobinopathy. The most effective approach to reduce the burden of the society is to reduce the incidence by implementation of a carrier screening programme.

Material and Methods: The present study was carried out in a tertiary care hospital for a period of 2 years. 500 Antenatal patients attending Obstetric-out patient department for routine check-up were included in the study. Screening of patient was done by Solubility test, Sickling test and Alkali denaturation test. Finally a very standardized method to detect HbA and HbF was done by using an automated High Performance Liquid Chromatography.

Results: Out of a total of 500 pregnant women 4% were having haemoglobinopathies, whereas 96% had normal haemoglobin. Out of 20 cases, maximum number of cases i.e. 2.2% was of β–Thalassemia trait, followed by Sickle cell trait cases i.e. 1.2%. Other haemoglobinopathies are also found i.e. sickle cell disease 0.2% δβ Thalassemia case 1.2% and double heterozygous 0.2%. Out of a total of 20 positive pregnant women only partners of 8 women were screened and out of them 37.5% were found positive.

Conclusion: Detection of carrier status using HPLC during pregnancy along with couple screening provides prospective parents with the option of testing the fetus for hemoglobinopathy.

Keywords: Antenatal, Double heterozygous, Haemoglobinopathies, High Performance Liquid Chromatography, β–Thalassemia trait.

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Introduction

Haemoglobinopathies are a complex group of red cell disorders, which constitute a major burden of genetic diseases and one of the major public health problems in India. Haemoglobinopathies have a highly variable clinical manifestations. At one end of the spectrum there is incompatibility with life and at the other, the patient under a stress, such as pregnancy, may experience some deterioration in her normal healthy state. Both the abnormal haemoglobin and the thalassaemia give rise to health problems of immense proportions.

The haemoglobinopathies are autosomal recessive inherited disorders of haemoglobin synthesis (thalassaemias) or structure (sickle cell disorder) that are responsible for significant morbidity and mortality on a worldwide scale.

Individuals with trait (carriers) are healthy and unaware of their carrier status unless specifically screened. If a couple both carry a clinically significant haemoglobinopathy trait there is a 1 in 4 chance with each pregnancy that their child will inherit a major haemoglobinopathy. The exact magnitude of the problem in India is still obscure. There is a genetic, ethnic and regional diversity of the haemoglobin variants as well as of the mutations in India which emphasizes to tackle the problem at a regional level. Most of the patients of haemoglobinopathies have a high morbidity rate, intercurrent infections being unusually common, suffer from high economic burden, terminate fatally in childhood, and have emotional and psychological trauma including the family members.

The most effective approach to reduce the burden of the society is to reduce the incidence by implementation of a carrier screening programme. There is an urgent need for making the people aware of this lethal malady. Health
education is an important component of the preventive genetic programmes.\textsuperscript{4} Thalassaemia and haemoglobinopathy, which are prevalent throughout India, is heritable, treatable, curable and preventable disorders. A joint venture of antenatal and inductive screening seems to be the most fruitful strategy for haemoglobinopathy in India. Hence initiating a preventive programme for these diseases is a necessity rather than an option.\textsuperscript{5}

**Material and Methods**

The present study was carried out in a tertiary care hospital for a period of 2 years from 2008 to 2010. 500 Antenatal patients attending Obstetric-out patient department (OPD) for routine check-up were included in study. Antenatal care (ANC) patient with clinical suspicion of anaemia, attending the ANC clinics (Routine checkup) at OBGY Department were included in study group. Institutional Ethical committee clearance was taken. Blood was collected in ethylene diammine tetra acetic acid (EDTA) bulbs from the anticubital vein, with all aseptic precaution for different types of laboratory investigations. Complete blood counts (using Mythic18, Orphee SA counter) and smears stained by using Standardised Romanowsky stain, Leishman stain for peripheral blood smears and Brilliant cresyl blue stain for reticulocyte count of all the patients were studied. Further screening of patient was done by Solubility test, Sickling test and Alkali denaturation test. Finally a very standardized method to detect HbA\textsubscript{2} and HbF and presence of some abnormal Hb was done by using an automated high performance liquid chromatography (HPLC) (using VARIANT\textsuperscript{TM} of Biorad Company) to study sickle cell and β-thalassemia syndromes. This is known as β-Thalassemia short programme. A final correlation of Solubility test, Sickling test, Alkali denaturation test and evaluation of HPLC was done in detection of sickle cell disorders and thalassaemia in Antenatal cases.

**Observations**

Out of total 500 pregnant women, 20(4\%) women are having haemoglobinopathies, whereas rest 480 women have normal haemoglobins (96\%). Out of 20 cases, maximum number of cases i.e. 11(2.2\%) were of β-Thalassemia trait, followed by Sickle cell trait cases i.e. 06 (1.2\%). Other haemoglobinopathies are also found i.e. sickle cell disease 1(0.2\%), δβ Thalassemia case 1(1.2\%), and double heterozygous 1 (0.2\%). [Table 1, Fig. 1]

| Case Distribution          | No. of cases | Percentage |
|----------------------------|--------------|------------|
| Normal Cases               | NAD          | 480        | 96\%       |
| Affected Cases             |              |            |            |
| B- Thalassemia trait       | 11           | 2.2\%      |
| Sickle cell trait          | 06           | 1.2\%      |
| Sickle cell disease        | 01           | 0.2\%      |
| δβ Thalassemia             | 01           | 0.2\%      |
| Double Heterozygous        | 01           | 0.2\%      |
| Total Cases                | 500          | 100\%      |

**Table 1: Distribution of all cases studied**

**Figure 1: Percentage distribution of affected cases (20 Cases)**
All the cases of β- Thalassemia show HbA2 values >3.5. This can also be seen in double heterozygous case. The HbF values are increased, i.e. >1 in SCD, δβ Thalassemia and double heterozygous case. Whereas the abnormal Hb i.e. HbS is seen in SCT, SCD and double heterozygous. [Table 2] It is evident from the table that a statistically significant difference is noted for values of HbA2 in normal pregnant women and β - Thalassemia trait cases. Also a statistically significant difference is noted for HbS/Abnormal Hb values in normal pregnant women and sickle cell trait cases.

It is found that sickling test is showing positivity in all the cases of sickle cell trait, SCD and double heterozygous, with none of the cases showing false positivity. [Fig. 2, 3] However solubility test is showing false positivity in 4 cases, and AD test is showing false positivity in 1 case. [Table 3] Sickling test has highest sensitivity and specificity as compared to the solubility test and AD test.

Table 2: HPLC interpretation in affected cases

| Disease                        | HbF      | HbA2  | Abnormal Hb |
|--------------------------------|----------|-------|-------------|
| Normal Pregnant women (480)    | 0.4 ± 2.1| 3.0 ± 0.5 | 0.7 ± 0.1  |
| Thalassemia trait(11) (mean±S.D)| 0.3 ± 0.5| 5.4 ± 0.4 | 0.6 ± 0.1  |
| Sickle cell trait (06) (mean±S.D)| 0.9 ± 0.7| 3.3 ± 0.4 | 34.6 ± 7.3|
| P value                        | >0.05    | >0.05* | >0.05       |
| Sick cell disease (01)          | 12.9     | 2.8    | 80.5        |
| δβ Thalassemia (01)             | 11.3     | 2.8    | 0           |
| Double Heterozygous (01)        | 18.7     | 3.9    | 48.2        |

* - Statistically significant

Figure 2: (a) b thalassemia trait; (b) Sickle cell disease

Figure 3: (a) Delta beta thalassemia; (b) Double heterozygous
Table 3: Test results of different screening tests

| Results         | Solubility test | Sickling test | AD/FF test |
|-----------------|-----------------|---------------|------------|
| True Positive   | 9               | 8             | 5          |
| True Negative   | 487             | 492           | 494        |
| False Positive  | 4               | Nil           | 1          |
| Sensitivity     | 69.20%          | 100%          | 83.33%     |
| Specificity     | 100%            | 100%          | 99.79%     |

Out of total 20 positive pregnant women only partners of 8 women were screened. Rest of the 12 partners was either not traceable or not ready to get screened. Out of total 8 partners screened, 3 (37.5%) partners are found positive. 1 screened positive for SCT and other 2 for β-Thalassemia trait. Thus 3 out of 8 couples screened are at risk of having children with haemoglobinopathies. [Table 4]

Table 4: Results for partner screening

| Result | No. of cases | Total |
|--------|--------------|-------|
| Positive | 1 (SCT) + 2 (Thal-trait) | 3     |
| Negative | 5            | 5     |
| Total     | 8            | 8     |

Discussion

The prevalence of haemoglobinopathies in present study was found to be 4%. WHO (World Health Organization) estimates that 5% of adults are carriers with 3.9% β-Thal and 2.3% SCD. Thus, prevalence rate of haemoglobinopathies in present study correlates well with study by Rowley et al. Health technology assessment in England, Al-Allawi et al. The prevalence of β-Thal trait in present study is 2.2% which is in accordance with study by Gupta et al. and Panda et al. However a little higher prevalence is found in Al-Allawi et al and Sachdev et al. studies as compared to present study. The prevalence of Sickle cell trait in present study matches that with Al-Allawi et al study i.e. 1.2%. However in all other studies the SCT prevalence is much higher than present study. The prevalence of Sickle Cell Disease in present study correlates well with Gupta et al study. According to Census 2001 the expected sufferers of sickle cell gene i.e. sickle cell disease is 0.5%. Thus these values correlate well with present study. Prevalence of δβ-thal correlates well with Al-Allawi et al study. No reference regarding studies for double heterozygous in pregnant women are found. Studies are done by various authors in double heterozygous, in general population. Mittal et al recorded a high incidence of double heterozygous i.e. 3-7%. The routine diagnostic test is Hb HPLC: these will demonstrate increase in HbA2 (i.e., >3.5% of total hemoglobin) and usually HbF (i.e., >1%). In the right clinical and ethnic context, an elevated HbA2 is considered diagnostic of β-thalassemia trait. [Table V] Thus the elevated level of HbA2 is diagnostic of β-thalassemia trait also elevated levels of HbA2 in present study matches well with other studies. Also a statistically significant difference is noted for values of HbA2 in normal pregnant women and β-Thalassemia trait cases, in present study. [Table 5]

Table 5: Comparative Study of values of HbF, HbA2 and HbS in various haemoglobinopathies.

|                      | R.S.Balgir | Amrita Panda et al. | Present Study |
|----------------------|-----------|---------------------|--------------|
| β-thal trait         |           |                     |              |
| HbF                  | 1-5       | 0.6                 | 0.3          |
| HbA2                 | 3.5-7.0   | 4.5                 | 5.4          |
| Sickle Cell trait    |           |                     |              |
| HbF                  | N         | 0.8                 | 0.9          |
| HbA2                 | 2-4       | 1.75                | 3.3          |
| HbS                  | 38-45     | 25.22               | 34.6         |
| Sickle Cell Disease  |           |                     |              |
| HbF                  | 1-20      | -                   | 12.9         |
| HbA2                 | 2-4       | -                   | 2.8          |
| HbS                  | 75-95     | -                   | 80.5         |
| Double Heterozygous  |           |                     |              |
| HbF                  | 5-30      | -                   | 18.7         |
| HbA2                 | 4-8       | -                   | 3.9          |
| HbS                  | 60-85     | -                   | 48.2         |
Comparative Study for values of HbF, HbA2 and HbS for δβ-Thal trait, sickle cell trait, sickle cell disease and double heterozygous matches very well with the reference values. [Table 5]

It is evident from present study that a difference is noted for HbS/Abnormal Hb values in normal pregnant women and Sickle cell trait/disease cases. In addition to HbS, HbF values are also significantly raised in Sickle cell disease. Hence significantly raised HbS can be reliably used to diagnose cases of Sickle cell trait cases while raised HbS and HbF can be reliably used to diagnose cases of Sickle cell disease cases in pregnancy.

In Double Heterozygous cases, significant difference is noted for all the Hb HPLC values as compared to normal pregnant women. Hence these values can be reliably used to diagnose cases of Double Heterozygous in pregnancy.

No comparative studies for values of HbF, HbA2 and HbS for δβ-Thal in pregnant women are found so far. However, difference is noted for HbF values among δβ-Thal case in pregnant women. Sickling test had highest sensitivity and specificity of 65% and 95% respectively compared with solubility test which had sensitivity and specificity of 45% and 90% respectively according to study by Andrew LivexOkwi et al. Overall sensitivity and specificity of solubility test was found to be 93.8% and 100% respectively with positive predictive value of 100% and negative predictive value of 97.4% according to study by Mukherjee et al.

In the present study it was found that sickling had 100% sensitivity and specificity as compared to solubility test which has 69.2% sensitivity and 100% specificity. Present study correlates well with above studies. In over two million automated HPLC screening tests carried out in California between 1990 and 1993, only 1 false positive and 1 false negative test have been recorded (unpublished report).

A recent study across nine US laboratories, using automated HPLC (Bio-Rad) and the same standard operating procedures for the State of California Neonatal Screening Programme, has reported a specificity of 99%, while the same programme has reported, to members of the US Guideline Panel, a sensitivity of over 99.9% for the technique. Present evidence and experience suggest that HPLC have acceptable sensitivity and specificity when properly used. Study by Sachdeva et al. states that cation exchange HPLC is a convenient, efficient, reproducible & cost effective method for screening & diagnosis of Thalassemia & other haemoglobinopathies. As compared to above studies, in present study it was found that HPLC is emerging as the most advance technique in diagnosis of Haemoglobinopathies, with acceptable sensitivity and specificity. Also a very convenient, efficient and reproducible method. Screening for hemoglobinopathy was done in which 55% partners were tested. 77 partners at risk were identified out of 810. 3 out of 591 couples were at risk of having child with hemoglobinopathy like Thal major/intermedia or 5/1000. If black women has sickle cell trait and her partner is black, 1:40 chances that she will have a child with sickle cell disorder more commonly SCA, SCD, delta-betaδ-β-Thal. In present study 3 out of 8 partners were tested for hemoglobinopathy that is 37.5% partners were positive. Hence highly significant.

Conclusion

Using HPLC method in screening for Thalassemia and sickle cell disorder in pregnant women in the region support the notion of establishing a preventive and control program of this common genetic disorder. Detection of carrier status during pregnancy along with couple screening provides prospective parents with the option of testing the fetus for a hemoglobinopathy. This will give parents the opportunity of planning a family without disease, also alleviating the health burden to society. A joint venture of antenatal and high risk couple screening seems to be the most fruitful strategy for control of haemoglobinopathy in India.

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