Can Homeopathy Bring Additional Benefits to Thalassemic Patients on Hydroxyurea Therapy? Encouraging Results of a Preliminary Study

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Several homeopathic remedies, namely, Pulsatilla Nigricans (30th potency), Ceanothus Americanus (both mother tincture and 6th potency) and Ferrum Metallicum (30th potency) selected as per similia principles were administered to 38 thalassemic patients receiving Hydroxyurea (HU) therapy for a varying period of time. Levels of serum ferritin (SF), fetal hemoglobin (HbF), hemoglobin (Hb), platelet count (PC), mean corpuscular volume (MCV), mean corpuscular hemoglobin concentration (MCHC), mean corpuscular hemoglobin (MCH), white blood cell (WBC) count, bilirubin content, alanine amino transferase (ALT), aspartate amino transferase (AST) and serum total protein content of patients were determined before and 3 months after administration of the homeopathic remedies in combination with HU to evaluate additional benefits, if any, derived by the homeopathic remedies, by comparing the data with those of 38 subjects receiving only HU therapy. Preliminary results indicated that there was a significant decrease in the SF and increase in HbF levels in the combined, treated subjects. Although the changes in other parameters were not so significant, there was a significant decrease in size of spleen in most patients with splenomegaly and improvement in general health conditions along with an increased gap between transfusions in most patients receiving the combined homeopathic treatment. The homeopathic remedies being inexpensive and without any known side-effects seem to have great potentials in bringing additional benefits to thalassemic patients; particularly in the developing world where blood transfusions suffer from inadequate screening and fall short of the stringent safety standards followed in the developed countries. Further independent studies are encouraged.

Keywords: Ceanothus Americanus – ferritin – Ferrum Metallicum – fetal hemoglobin – homeopathic remedy – Hydroxyurea – Pulsatilla Nigricans – Thalassemia

Introduction

Thalassemia is a hereditary anemia resulting from defects in hemoglobin production. It is among the most common genetic disorders worldwide. About 4.83% of the world’s
population is carrying globin variants, including 1.67% of the population that is heterozygous for α-thalassemia and β-thalassemia (1). In β-thalassemic patients, there is a decrease in the production of β-globin chains, as a consequence of which multiple organs get affected (2) and considerable mortality results. In HbE/β-thalassemia, there is a mutation on both genes for the synthesis of β-chain in adult and hemoglobin E meant for synthesis of embryonic hemoglobin. Thalassemia is caused by any of more than 200 point mutations and rarely by deletions (3). It is clinically heterogeneous because various genetic lesions variably impair globin chain synthesis. The anemia associated with thalassemia may be moderate to severe depending on the gene mutations involved. Severe anemia is generally accompanied by ineffective erythropoiesis, with bone expansion and extra medullary hematopoiesis in the liver and spleen (also at other sites) generally resulting in enlarged liver and spleen. Transfusion therapy is considered the main stay of treatment when the patients become anemic beyond a threshold with all typical sufferings of thalassemia, and treatment by the usual chelation therapy is no more effective.

However, transfusion-related infections (primarily Hepatitis B and C, or HIV) are now considered the major cause of death in developing countries where proper blood testing facility is not available. Although regular transfusion therapy can help maintain growth and development and also reduces hepto-splenomegaly due to extra medullary hematopoiesis, transfusion therapy is often taken as the last resort in the developing countries, partly for economic reasons and partly for lack of infra-structural facilities. Therefore, iron chelation therapy occupies a pivotal place in the control and management of thalassemia, because this therapy is reported to double the life expectancy of patients with thalassemia major (4,5). Of the different chelators, Deferiprone, Deferoxamine, etc., are regularly used, although they have been reported to have considerable side effects and they are generally more costly (6–10), precluding their use to a large extent by poor patients. However, a number of new oral chelators like Deferasirox (ICL 670) are presently under development (11).

It is well established that affected patients do not become anemic until the fetal (gamma) globin genes are developmentally silenced, and that patients with persistent high levels of fetal globin typically have less severe anemia, have milder clinical syndromes and are often transfusion independent (12). Therefore, another approach through augmenting the synthesis of fetal hemoglobin has been made by using drugs like 5-Azacytidine, Hydroxyurea (HU) and various butyrate derivatives (13). HU has shown substantial benefits in a subgroup of patients with sickle cell anemia (14) and in a few pediatric patients with thalassemia where transfusion requirements were generally eliminated after treatment with HU for ~20 months (15,16). However, several authors (13,17) reported the role of HU in thalassemia therapy to be rather inconsistent, some showing good responses while others did not (17,18). Further, despite the beneficial effects of HU on erythropoiesis it may be difficult to correct the anemia associated with thalassemia with the use of HU alone particularly in case of genetic predisposition, such as the presence of HbE polymorphism (19) and the type of thalassemia such as hemoglobin E-thalassemia (20). Therefore, importance of proper genetic screening has been emphasized by some authors (21,22).

Recombinant human erythropoietin was employed to achieve the benefit of increasing 'thalassemic erythropoiesis' without raising fetal hemoglobin in β-thalassemia intermedia who had undergone splenectomy (23). Recently, the treatment with Darbepoetin-α was reported to increase hemoglobin levels substantially in patients with HbE/β-thalassemia disease. However, the search for new compounds that augment fetal hemoglobin production and are non-toxic in nature is still on in in vitro system as well as on various animal models (24–26). Iron overload causes most of the mortality and morbidity associated with thalassemia, and serum ferritin (SF) is most commonly measured as an indicator of iron stores. Ferritin levels below 2500 mg/ml are associated with improved survival (1), and therefore drugs that are non-toxic but capable of reducing SF level may be useful in thalassemia.

In the present investigation, therefore, homeopathic remedies that generally agreed with the symptoms of thalassemic patients have been administered either singly or in combination depending on changeable symptoms in patients who have been on HU treatment for varying periods of time from a few months to over 6 years. In this investigation, the primary objectives were to examine if the homeopathic remedy, in combination with HU, could bring beneficial effects, as manifested by (i) diminishing SF level, (ii) augmenting HbF synthesis and (iii) reduction of spleen size in patients with splenomegaly.

Material and Methods

Subjects
This research was conducted in compliance with the protocols of Helsinki Declaration and the Institutional Ethical Committee, University of Kalyani. Informed consents were also obtained from the patients or their parents (of minors), who volunteered to undertake this study. Subjects were examined for their general health and status of their spleen enlargement periodically.

Thirty-eight patients attending the Thalassemia Foundation Clinic at F-44 Footnani Chamber, Corporation Place, Kolkata-700013 from March 2001 to September 2006 had been put on regular HU administration at rates varying
between 5 mg/kg/day to a maximum of 20 mg/kg/day depending on conditions of the patient. Most of these patients were transfusion dependent.

Test protocols used and number of subjects for each protocol

They were periodically examined for physical conditions of liver and spleen and also for other blood parameters like SF content, fetal hemoglobin (HbF) and hemoglobin (Hb) content before and after administration of HU. Tests were made on an actual cost-no-profit basis only when patient’s condition demanded, as most of them were extremely poor, many of them visiting from some other districts like Midnapur, Cooch Behar in West Bengal. The results and number of subjects who provided samples for examination before and after HU treatment totaled 38, of which 32 belonged to HbE/β type and 6 belonged to β-thalassemia major type. Of the 32 HbE/β patients, the mean values of SF content of 18, and HbF and Hb contents of all 32 were studied. Of the six β-major patients, the mean values of SF content of four, HbF level of three and Hb content of all six have been obtained. This variation was unavoidable owing to the disapproval of some patients for testing too many blood parameters. Out of 38 patients who received homeopathic remedies as well as HU, 32 belonged to HbE/β type and six to β-major type. Out of 32 patients with HbE/β thalassemia, SF level was determined from 20 patients, HbF from 10, Platelet count (PC) from 27, Hb content from 27, mean corpuscular volume (MCV), mean corpuscular hemoglobin concentration (MCHC), and white blood cell (WBC) count from 27 patients each, bilirubin from 6, aspartate amino transferase (AST) from 4, alanine amino transferase (ALT) from 4, serum total protein content from 4 and mean corpuscular hemoglobin (MCH) was analyzed from 27 patients.

Out of six β-major patients, SF and HbF level of four subjects, PC and Hb content of six, MCV, MCHC and WBC of six patients each, bilirubin, ALT, AST and serum total protein from two subjects each and MCH from six subjects were considered. Incidentally, the patients showing symptoms of liver disorders and jaundice were subjected to additional tests like AST, ALT and bilirubin.

Age and sex of patients and blood transfusion periods

The age of these 38 patients ranged between 6 and 25 years, the majority being in 4–10 years’ age group. In β-major group, there was one female and five male subjects. In HbE/β group, there were 20 males and 12 females. Except for three patients, the remaining patients had history of blood transfusion ranging from once in a week to (rarely) once in 6 months.
High-performance liquid chromatographic analysis of hemoglobin

Hemoglobin type HbF was quantified by automated high-performance liquid chromatography (VARIANT; Bio-Rad, Hercules, CA, USA) (29).

Immunoassay of Ferritin EIA

Enzyme immunoassay for the quantitative determination of immunoglobulin subclass E (IgE) in human ferritin was made in an ELISA Reader (Biorad, USA) by using the kits from Genix Technology Inc, Canada.

For Hb analysis, a Cell Counter (Medonic CA 530, E. Merck, Germany) was used by deploying standard kit from E. Merck, Germany.

Total protein, ALT, AST and Bilirubin

These were recorded by Biochemical Semi auto-analyzer, Microlab 300 (E. Merck, Germany).

Statistical Analysis

The levels of significance of differences were determined between the data of parameters obtained before the homeopathic drug administration and 3 months after administration of the remedies by Student’s ‘t’ test method.

Results

Urinary Iron Excretion Pattern

A careful analysis of the data (Fig. 1) revealed a relatively low iron content from the first day and showed a wave like curve in Cean-6-fed group. On the other hand, the initial iron contents were recorded to be more in the other drug-fed and placebo-fed series. There was also a wave like fall and rise of iron contents in these groups as well as in the placebo, except for the fact that the Puls-30- and Puls-Q-fed group showed relatively more iron excretion. Therefore, there was no clear indication of any definite trend of iron excretion except for the fact that iron excretion appeared to be marginally more in the drug-fed groups (except Cean-6-fed group) and subsequently the three placebo-fed patients were also administered homeopathic remedy as per similia principle. Since the main objective was also to examine if the homeopathic remedy could benefit the patients already on HU treatment, we did not put any further importance to the placebo-fed control as we considered the HU administration group as our main control to evaluate additional benefit, if any, derived by the administration of homeopathic remedy.

Changes in SF, HbF and Hb Levels After Drug Administration

There was a decrease of ferritin level and an increase of HbF content in both HbE/β and β-major subjects after administration of HU (Fig. 2A and B), although Hb content in HbE/β group did not appreciably decrease. There was a noticeable decrease in the Hb content of the β-major group after administration of HU for a varying period of time ranging from a therapy for 1–5 years (complete data not shown). There was a significant reduction of SF in the combined treatment (P < 0.001) in HbE/β group (Fig. 3A).

Although there was a further reduction in the β-major group, the decrease was statistically insignificant like that of only HU treatment group. There was also a significant increase (P < 0.05) in HbF level in HbE/β group but in the β-major group there was actually a decrease in HbF level (Fig. 3B). The Hb content in HbE/β group however, was very marginally reduced, while there was a little increase in Hb level in the β-major group (Fig. 3C).
Figure 2. Effect of HU on thalassemic subjects (BD = HU untreated, AD = HU treated; N = Number of subjects): (A) SF level. (B) Fetal hemoglobin (HbF) content. (C) Hemoglobin (Hb) content.

Other Blood Parameters in the Combined Treated Group

The PC in both HbE/β and β-major were decreased (Fig. 3D). The MCV of Hb E/β group remained more or less the same, while there was a decrease in β-major group (Fig. 3E), which was statistically insignificant. The MCHC in HbE/β group remained more or less the same, while in β-major there was a little increase (Fig. 3F). There was also a little increase in MCH (Fig. 3G) of the β-major group; however, the increase was statistically insignificant. The WBC count in HbE/β was slightly increased while it was marginally decreased in β-major (Fig. 3H). The bilirubin content was same in HbE/β while it was decreased marginally in β-major group (Fig. 3I). While there was a distinct reduction in the AST (SGOT) level in Hb E/β group, in β-major there was only a marginal increase noted, though the level was in the normal range (Fig. 3J). The same was true for ALT (SGPT) level (Fig. 3K), while in β-major there was palpable change in serum total protein level, there was marginal increase in serum total protein level (Fig. 3L) of Hb E/β group.

Blood Transfusion Period

In most cases dependent on blood transfusion, there was an increase in the period of their transfusion demand (ranging between 25 and 75%) in the combined treatment group. In two cases, patients used to have fever, before and after transfusion, but after administration of homeopathic remedies on similia principle for 3 months, the incidence of fever disappeared.

Spleen Size and General Health

In the 30 patients having enlarged spleen of varying lengths (total 125 cm) before administration of homeopathic remedy, there was a distinct reduction in length (total 45 cm); as many as 11 of them showed no palpable enlarged spleen after homeopathic treatment. The general health and spirits of the patients appeared to have improved after this combined treatment in the majority of the cases studied.

Discussion

In the present investigation, out of a total of 38 patients, 6 belonging to thalassemia major type (β-thalassemia) and 32 to HbE/β type, the greater majority showed positive response to combined homeopathic therapy. There was a significant increase in induction of HbF in them, which is very encouraging. In fact, augmentation of fetal hemoglobin (HbF) synthesis is highly solicited in thalassemic patients as high levels of fetal hemoglobin production can ameliorate sickle cell disease and β-thalassemia (30). Incidentally, oral administration of HU alone has been reported to increase the level of HbF in both thalassemia major and thalassemia intermedia patients, with varying degree of response, presumably because of the molecular heterogeneity of the β-globin gene in major β-thalassemia and the complexity of interacting genetic factors involved in HbF switching in each respective patient (31). In fact, among the various drugs recently proposed to improve the clinical conditions in both sickle cell disease and β-thalassemia (20,32–36), HU was stated to be the most effective. In the present study, a noticeable increase in HbF was also noted in patients receiving HU alone, but there was further increase of HbF level in patients receiving the combined homeopathic treatment, thereby showing additional beneficial effect of homeopathy. However, effect on individual patients in regard to increase in HbF was variable (data not shown) as also reported in the other study (13). In case of Hb, in the only HU-treated group, significant increase of HbF was also noted in patients receiving HU alone, but there was further increase of HbF level in patients receiving the combined homeopathic treatment, thereby showing additional beneficial effect of homeopathy. However, effect on individual patients in regard to increase in HbF was variable (data not shown) as also reported in the other study (13). In case of Hb, in the only HU-treated group, there was a considerable increase in Hb content in the β-major group, but there was actually a little increase in the combined treated group. Therefore, on the whole, there appeared to be additional benefit of homeopathic drug in terms of both HbF and Hb contents. It is suggested that HU produces fetal hemoglobin via reactivation of gamma genes through some unknown
Figure 3. Comparative efficacy of only HU (BD) treatment and HU plus homeopathic drug (AD) treatment on thalassemic subjects: (A) SF level. (B) HbF content. (C) Hb content. (D) PC. (E) MCV. (F) MCHC. (G) MCH. (H) WBC Count. (I) Bilirubin content. (J) AST level. (K) ALT level. (L) Serum total protein level.
molecular mechanism (37). These authors also reported promising results of HU therapy in thalassemia intermedia.

But there are reports in which some patients exhibited insignificant or no change in HbF levels after HU treatment (20,33). Nevertheless, HU therapy has been termed as the most promising therapy because of the oral route of administration, for being relatively inexpensive and with no or little side effects (37). Therefore, if the induction of HbF level could be further accelerated or rejuvenated, as the homeopathic therapy appeared to produce, could be of additional benefit to the patients. The advantages of the homeopathic remedies are manifold. They are also inexpensive, easily available, taken in micro doses and without any known side-effects.

How the homeopathic remedies could bring forth such positive changes remains unclear. But the present results would also lend support to the contention (38–40) that the potentized homeopathic remedies possibly acted through regulation of specific and relevant gene expression responsible for the apparently enhanced synthesis of the HbF and decrease of ferritin level. While in the advanced countries the treatment and management strategies have made significant progress in the reduction of thalassemia to a great extent, such treatment strategies are beyond reach of the poor patients inhabiting developing countries. Incidentally, homeopathic remedies also showed considerable potentials to ameliorate arsenicosis (41–43), a burning public health problem of arsenic exposure through groundwater contamination in some third world countries, because only limited success could so far be achieved with the orthodox medicine tested to cure arsenicosis. Therefore, the initial results of the present study are very encouraging and need further trial by other researchers independently to verify and confirm (or refute) our findings. Further works involving additional important parameters are presently in progress.

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