Combination of exchange transfusion treatment and hydroxyurea cause beneficial changes to laboratory parameters and clinical outcome in patients with sickle cell disease/β thalassemia compared with hydroxyurea or exchange transfusion alone

Abstract

Sickle-cell disease is one of the most common severe monogenic disorders in the world. Hemoglobin polymerization, leading to erythrocyte rigidity and vaso-occlusion, is central to the pathophysiology of the disease, although the importance of chronic anemia, haemolysis, and vasculopathy has been established. Clinical management is basic and few treatments have been established in evidence base. Hydroxyurea (HU) is considered to be the most successful drug therapy for severe sickle cell disease (SCD). Red blood cell transfusion therapy also has significantly improved the morbidity and mortality of patients with SCD and improves the quality of life among these patients. This study favored chronic transfusions and hydroxyurea compared with hydroxyurea alone or red cell exchange for the outcomes: annual crisis rate; acute chest syndrome; stroke; anemia; pulmonary hypertension; liver function. Levels of hemoglobin, fetal hemoglobin and hemoglobin S; MCV; No serious adverse effects were reported from the study. We conclude that chronic transfusions combined with hydroxyurea improve the quality of life and the laboratory parameters in adults with SCD without any adverse effects over the last 5 years.

Keywords: exchange transfusion, hydroxyurea, sickle cell, combination, hemoglobin

Abbreviations: SCD, sickle cell disease; CT, computed tomography; PME, partial manual exchange transfusion; NHLBI, national heart, lung, and blood institute; TIA, transient ischemic attack; SCIC, sickle cell intrahepatic cholestasis; RCT, randomized controlled trial; ACS, acute chest syndrome; CTT, chronic transfusion therapy; SWITC, stroke with transfusions changing to hydroxyurea; PH, pulmonary hypertension; PAH, pulmonary arterial hypertension; MPAP, mean pulmonary artery pressure; RHC, right heart catheterization

Introduction

Sickle cell disease is a multisystem disease associated with episodes of acute painful illness and progressive organ damage.1–3 The term sickle cell disease is used to refer to all the different genotypes that cause the characteristic painful clinical syndrome, such as homozygous for the βs allele, and also the co-inheritance of the βs and βc or the βs and the β-thalassemia allele.4 The high morbidity rate of SCD patients is related to vascular complications5 that include multiple chronic organ damage affecting the brain, heart, lungs, kidneys, liver, eyes, skin, and skeleton. Vaso-occlusive crises result in acute and chronic severe pain, as well as acute chest syndrome, splenic sequestration, hemolytic anemia, stroke, acute and chronic multi-system organ damage, and shortened life expectancy. Chronic transfusion6 and hydroxyurea1 are commonly used preventive treatments for sickle cell disease. Transfusion therapy is the standard of care to prevent an increased risk of stroke reducing the frequency of pain and acute chest syndrome. Hydroxyurea is a drug that used to raise fetal hemoglobin and is an attractive to transfusion for clinical complications including chronic pain, ACS, stroke and hospitalizations.

Patients and methods

The study was conducted at Thalassemia and Transfusion Department of Hippokratio Athens Hospital. The medical records of 30 patients who were in continuously follow up since 2010-2015 were retrospectively analyzed. The patients were identified who had clinical events for the use of hydroxyurea alone, exchange transfusion, or combination of chronic transfusion and hydroxyurea.

For each patient the steady state hematological parameters were obtained. SCD genotypes were established by hemoglobin analysis by HPLC (Variant II-HPLC, Board) methods. All patients were followed up every month with a complete blood count and differential, biochemistry and HbS/Hbf levels. All the important adverse effects and the reported events by the patients reviewed in every visit. Examinations were performed to evaluate cardiac function using pulsed two-dimensional M mode and color flow Doppler echocardiography. Cardiac dimensions and pulmonary parameters were measured according to the criteria of the American Society of Echocardiography. Chest Computed Tomography (CT) was performed with high resolution protocol to determine the diameter of pulmonary artery. Neurological evaluation and MRI/MRA were performed before the enrollment.
Combination of exchange transfusion treatment and hydroxyurea cause beneficial changes to laboratory parameters and clinical outcome in patients with sickle cell disease/β thalassemia compared with hydroxyurea or exchange transfusion alone

In our patients hydroxyurea was started at a dose of 15mg/kg per day in a single oral dose and the dose escalated every four weeks to the maximum tolerated dose or to a maximum dose of 30mg/kg per day. Sickle cell disease patients were prescribed transfusions every 3-4 weeks with the goal of maintaining HbS levels of ≤30%, according to the standard clinical care guidelines of the National Heart, Lung, and Blood Institute (NHLBI). Transfusion modalities included either:

i. Simple transfusion (ST) of 10-15mL/kg RBCs with the post-transfusion Hb goal of 11.5-12.0g/dL

ii. Partial manual exchange transfusion (PME): Removal of 5-10mL/kg of whole blood prior to transfusion of 10-15mL/kg RBCs with the post-transfusion Hb goal of 10-12.0g/dL or

iii. Automated RBC exchange with the post-transfusion goal of 25-30% fraction of cells remaining and Hb approximately 0-2g/dL above the pre-transfusion Hb.

All RBC units were pre-storage leuko reduced, sickle-negative, ABO/Rh compatible, C/c, E/e, and Kell matched with additional antigenic matching dependent on alloantibody identification. Regularly scheduled blood transfusion therapy or exchange transfusion involves periodic transfusion of the patient at regularly scheduled intervals, with the frequency guided by the patient’s symptoms, Hgb, and percent HgbS. A detailed deferoxamine or deferasirox protocol after 1 year of chronic transfusion therapy or exchange blood transfusion was initiated.

Patient’s characteristics

30 patients (15 male, 15 female), with S/S genotype 5/30 and S/β genotype 25/30 and mean age at the study entry 45, 2 years, were divided into three subgroups of 10, depending on the received therapy: hydroxyurea, exchange blood transfusion, and exchange transfusion and hydroxyurea combination. The indications for entering in each subgroup were: more than 5 painful hospitalizations/year 25/30, ACS 6/30, prevention of stroke 8/30, pulmonary hypertension 4/30, symptomatic or severe anemia 1/30 and liver disease 1/30. Mean HB, HBS, HBF and MCV were calculated in all groups, pre and post treatment. Mean difference of HB, HBS, HBF and MCV from baseline to post treatment were also calculated, in all groups (Pre treatment value-Post treatment value).

Table 1 Patient characteristics

<table>
<thead>
<tr>
<th>Hydroxyurea (n=10)</th>
<th>Exchange Transfusion (n=10)</th>
<th>Combined Therapy (n=10)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Enrollment</strong></td>
<td><strong>End of study</strong></td>
<td><strong>P</strong></td>
</tr>
<tr>
<td>Pain Crisis</td>
<td>7</td>
<td>3</td>
</tr>
<tr>
<td>ACS</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>PH</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Stroke</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Liver Disease</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Anemia</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Citation: Delicou S, Lavda M, Argiri V, et al. Combination of exchange transfusion treatment and hydroxyurea cause beneficial changes to laboratory parameters and clinical outcome in patients with sickle cell disease/β thalassemia compared with hydroxyurea or exchange transfusion alone. Hematol Transfus Int J. 2016;2(4):71–75. DOI: 10.15406/htij.2016.02.00042
Combination of exchange transfusion treatment and hydroxyurea cause beneficial changes to laboratory parameters and clinical outcome in patients with sickle cell disease/β thalassemia compared with hydroxyurea or exchange transfusion alone

In group 2 (exchange transfusion/transfusion) of the 2/10 patients who enrolled with a painful crisis, only 1/10 had a more than 5 painful crisis during the study. This patient, a 50-year-old male, was under exchange transfusion program every 2–3 months while on transfusions had multiple auto- and allo-antibodies and there was a need of extended red cell phenotyping and difficulty finding compatible blood. Of the 2/10 who enrolled with ACS none had a recurrent acute chest syndrome. The 4/10 patients with stroke who enrolled in exchange transfusion program one had experienced a new event of TIA presented with neurological deficits and negative MRA/MRI. Of the 2/10 patients who enrolled in this group with pulmonary hypertension, the 1/2 had a persistent hemodynamically significant pulmonary hypertension. This patient, a 76-year-old woman had also abnormal kidney function which is an important parameter in patients with sickle cell disease and PH and has been related to mortality. The same patient had also a persistent anemia, underwent to exchange transfusion after she maintained the percent HbS near 30 and the pretransfusion Hct between 25–28%.

In group 3 (combination of hydroxyurea and exchange transfusion) the 1/10 patient who enrolled with debilitating painful crises, didn’t experience any painful crisis. Of the 2/10 patient with ACS none had recurrent acute chest syndrome or developed chronic pulmonary disease. None of the 5/10 had experienced a new event included transient ischemic attack (TIA). Both the 4/10 patients who enrolled with pulmonary hypertension, had a significant reduction in pulmonary artery pressure. The 1/10 patient with liver disease, a 48-year-old woman is the same with recurrent admissions for pain crisis and pulmonary hypertension sickle cell intrahepatic cholestasis (SCIC) has stayed stable with no evidence of impaired hepatic synthetic function, liver failure or portal hypertension and ascites.

HBS, HBF, HB and MCV pre and post treatment were compared and presented at Table 2. In group 1, significant differences were observed at HBS, HBF and MCV from baseline to post treatment stage. Similarly, in group 2 and 3, significant differences were observed at HBS, HBF, MCV and HB also from baseline to post treatment stage. When we compared the mean differences of HB, HBS, HB and MCV (value at baseline value at post treatment stage) among the three groups, statistical differences was revealed. There was a significant decrease of HBS mostly in group 3 (combination therapy) vs group 1 (hydroxyurea) and group 2 (blood transfusion) (49.6±6.7% vs 9.2±2.8% and 35.3±7.4%, respectively; p<0.001). A significant increase of HB was revealed in group 2 vs group 1 (-1.97±0.79mg/dl vs -0.02±0.31mg/dl, respectively; p<0.001) but not vs group 3 (-1.97±0.79mg/dl vs -1.48±0.85, respectively; p=0.386) (Table 2).

In group 3, a significant increase of HBF was observed mostly in group 1 and group 3 vs a decrease in group 2 (-8.40±3.41% and -6.3±3.08% vs 0.77±0.77%, respectively; p<0.001). Finally, MCV increased significantly in group 1 and group 3 vs group 2 (-2.30±3.26FL and -2.50±5.16FL vs -5.80±6.06FL, respectively; p<0.001).

Table 2

<table>
<thead>
<tr>
<th>Hydroxyurea</th>
<th>Exchange transfusion/transfusion</th>
<th>Combined</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enrollment</td>
<td>End of study</td>
<td>Mean difference</td>
</tr>
<tr>
<td>Hb (mg/dl)</td>
<td>9.06±0.91</td>
<td>9.08±0.77</td>
</tr>
<tr>
<td>MCV (fl)</td>
<td>75.90±5.54</td>
<td>97.2±7.08</td>
</tr>
<tr>
<td>HBF (%)</td>
<td>2.73±0.76</td>
<td>11.1±3.42</td>
</tr>
<tr>
<td>HbS (%)</td>
<td>87.03±12</td>
<td>77.8±5.5</td>
</tr>
</tbody>
</table>

1. 1-Indicates significant difference from group 1 to group 2 (p<0.001).
2. 2-Indicates significant difference from group 1 to group 3 (p<0.001).
3. 3-Indicates significant difference from group 1 to group 2 (p<0.001).
4. 4-Indicates significant difference from group 2 to group 3 (p<0.001).

Discussion

Sickle cell disease is a life-long genetic disease that begins in childhood, affecting the structure of erythrocytes. Although SCA is genetically characterised by a single point mutation, there are various genetic modulators that affect the phenotype of this disease, and patients can manifest with varying degrees of clinical severity. Recurrent episodes of acute, severe pain are the hallmark of SCD. The pain is highly variable both within and among patients, and is the result of complex and poorly understood interactions between biological and psychosocial factors. The association between painful VOC visits and hospitalizations and mortality as reported by the CSSCD almost 20 years ago still remains significant in the contemporary era of SCD specific therapies.

Hydroxyurea (HU) was the first drug approved by the FDA for clinical use in sickle cell patients to induce HbF after its clinical effectiveness in reducing acute disease complications of painful crises and hospitalizations was demonstrated. However, clinical and laboratory response to HU is highly variable: not all patients reach a clinically significant increase in HbF even at maximum tolerated dose. Blood transfusion in SCD can serve two roles, either for therapy (typically for life-threatening SCD related complication) or for prophylaxis, to decrease the incidence of specific SCD related complications. Subsequently prospective studies have verified the efficacy and tolerability of hydroxyurea, leading to a placebo-controlled, randomized controlled trial (RCT) that demonstrated the efficacy of hydroxyurea in reducing painful vaso-occlusive crises and acute chest syndrome (ACS) in adults with SCA. Both children and adults with SCA benefited from reductions in the frequency of VOC in 11studies. In our study, the use of hydroxyurea decreased significantly the painful crises.

Citation: Delicou S, Lavda M, Argiri V, et al. Combination of exchange transfusion treatment and hydroxyurea cause beneficial changes to laboratory parameters and clinical outcome in patients with sickle cell disease/β thalassemia compared with hydroxyurea or exchange transfusion alone. Hematol Transfus Int J. 2016;2(4):71–75. DOI: 10.15406/htij.2016.02.00042
Combination of exchange transfusion treatment and hydroxyurea cause beneficial changes to laboratory parameters and clinical outcome in patients with sickle cell disease/β thalassemia compared with hydroxyurea or exchange transfusion alone

This is consistent with HBF’s ability to inhibit HBS polymerization, thereby decreasing either vaso occlusion or hemolysis. Initial MCV suggesting that cells became more spherical during treatment, perhaps because cell surface area could not continue to increase as cell volume was expanded by increasing Hb content indicating that the increase in MCV was associated with proportionally increased cell water. Although, there are no randomized trials demonstrating the optimal treatment of acute chest syndrome in adults with sickle cell disease, transfusion therapy especially exchange transfusion has remained the cornerstone of management in moderate to severe cases of acute chest syndrome in several centers. In our study there was not significant difference between the three therapeutic groups. This could be due to the advances in medical care and management strategies over the past decade and reflects the impact of advances in medical care in the given time frame.17–20

Natural history studies reveal that nearly 70% of SCA patients will suffer a recurrent stroke if left untreated. Additionally, almost 40% of SCA patients who have an overt stroke also have evidence of vasculopathy on magnetic resonance angiography placing this group of patients at the highest risk for recurrent stroke. Moreover, 45% of SCA patients who have had an overt stroke will suffer progressive neurologic damage due to both overt and silent cerebral infarctions, despite chronic transfusion therapy (CTT). Hydroxyurea was not equivalent to transfusion in the Stroke with Transfusions Changing to Hydroxyurea (SWiTCH) trial based on a composite end point; in those patients who have experienced prior stroke or at risk of initial stroke based on elevated TCD velocity, chronic blood transfusion or exchange transfusion is the standard of care to prevent recurrence or initial stroke. In our study we found that patient at high risk for stroke had significantly improvement with combination therapy of Hydroxyurea and exchange transfusions vs simple transfusions/ exchange transfusions vs hydroxyurea use. Subjects with the highest MCV values and HbF (group 3) had significantly more corroborative evidence of prevention from a new event of stroke.21–23

A similar relationship has been described in sickle cell patients with pulmonary hypertension. Another one of the most controversial complications of sickle cell disease (SCD) is pulmonary hypertension (PH). The words “pulmonary hypertension” like “anemia” does not indicate a specific diagnosis but imply a constellation of signs and symptoms that have many possible etiologies. The World Health Organization classifies PH into 5 groups which were collectively referred to as “pulmonary hypertension” with number one being pulmonary arterial hypertension (PAH). Pulmonary hypertension is defined as resting mean pulmonary artery pressure (MPAP) ≥25 mmHg determined by right heart catheterization (RHC). Moreover, the hallmark of PAH, besides the elevated MPAP, is a co-existent pulmonary-capillary wedge pressure ≤15 mmHg. About 3% of patients with SS develop PAH and the overall prevalence of all types of PH in SS is approximately 6%.

In both cases (Stroke and PH), exchange blood transfusion with hydroxyurea does more than simply raise the hemoglobin (Hgb) level for oxygen delivery; transfusion also lowers further the percentage of sickle Hgb (HgbS) and increases Hgb oxygen saturation, both of which decrease the propensity for vaso occlusion. The increased MCV may be linked to a reduce rate of sickling. The use of hydroxyurea is a mainstay in the overall management of individuals with SCD, since it reduces the incidence of hemolysis, and prolongs survival.20–23 The liver can be affected by a number of complications due to the disease itself and its treatment. In addition to the vascular complications from the sickling process, patients with SCD have often received multiple transfusions, placing them at risk for viral hepatitis, iron overload and (combined with the effects of chronic hemolysis) the development of pigment gallstones, all of which may contribute to the development of liver disease. In our study, the patient had multorgan complications from the disease.24–30

Conclusion

Prompt and effective treatment for SCA and its complications is recommended as it is likely to improve survival. Further research is required to determine the etiology, pathophysiology and the most appropriate strategies for management of SCA. The results of this study showed the combination of chronic exchange transfusions and hydroxyurea is superior to hydroxyurea alone or chronic transfusions alone.

Conflict of interest

The authors certify that they have NO affiliations with or involvement in any organization or entity with any financial interest (such as honoraria; educational grants; participation in speakers’ bureaus; membership, employment, consultancies, stock ownership, or other equity interest; and expert testimony or patent-licensing arrangements) or non-financial interest (such as personal or professional relationships, affiliations, knowledge or beliefs) in the subject matter or materials discussed in this manuscript.

Acknowledgements

Special thanks to my major advisors, Dr. Markisia Karageorgas and Prof. Athanassios Aessopos for their patience, understanding, guidance and most of all the encouragement they have given me during all this time. Sophia Delicou MD, Clinical Hematologist.

References

Combination of exchange transfusion treatment and hydroxyurea cause beneficial changes to laboratory parameters and clinical outcome in patients with sickle cell disease/thalassemia compared with hydroxyurea or exchange transfusion alone.


