Relief of Acute Chest Syndrome of Sickle Cell Anaemia by Nitric Oxide Inhalation - A Report of 2 Cases

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Abstract

Depletion of nitric oxide (NO) in the microcirculation plays a central role in the pathogenesis of sickle cell disease (SCD) and acute chest syndrome (ACS). Various other mechanisms which are responsible for the sickle-cell vaso-occlusive crisis are polymerization of sickle haemoglobin, platelet activation, endothelial damage and leucocyte adhesion. During the sickle cell crisis, the high rate of red blood cell (RBC) destruction leads to release of erythrocyte arginase (into the plasma), which thereby destroys arginine. Arginine, the substrate for NO synthase, once depleted, results in lower levels of NO in vessels, leading to vasoconstriction, pulmonary hypertension and ventilation perfusion mismatch. Increasing the availability of NO therefore has a central role in SCD, especially in cases of ACS. With the prevention of vasoconstriction by NO inhalation therapy in ACS, pain relief can be provided to the patients in sickle cell crisis. We present a report of 2 cases of sickle cell disease with ACS (Acute Chest Syndrome) in whom inhalation of NO through non-invasive ventilation (NIV) brought about pain relief and recovery from ACS.

Introduction

In patients with sickle cell disease (SCD), hypoxic conditions lead to polymerization of haemoglobin S in red blood cells (RBCs), ultimately resulting in the occlusion of blood vessels [1]. Polymerization of haemoglobin S in the RBCs leads to a change in morphology to make it a crescent or sickle shape, which is more rigid than the normal RBC. This change results in hemolysis, inflammation, adhesion of RBC and end organ ischemic reperfusion injury leading to infarct [2]. The release of haemoglobin, as a result of hemolysis, rapidly consumes nitric oxide, resulting in a whole sequence of events that inhibit blood flow. The increased rate of destruction of the deformed RBCs in patients with SCD, releases erythrocyte arginase into plasma, which thereby destroys arginine. Arginine, the substrate for NO synthase, once depleted, results in lower levels of NO in vessels, leading to vasoconstriction, pulmonary hypertension and ventilation perfusion mismatch. Increasing the availability of NO therefore has a central role in SCD, especially in cases of ACS. With the prevention of vasoconstriction by NO inhalation therapy in ACS, pain relief can be provided to the patients in sickle cell crisis. We present a report of 2 cases of sickle cell disease with ACS (Acute Chest Syndrome) in whom inhalation of NO through non-invasive ventilation (NIV) brought about pain relief and recovery from ACS.

Case Report

Case 1

A 31 year male patient of SCD by the name P.B. of SCD, was admitted to the hospital on 25th Aug, 2013 with complaints of bilateral chest pain, low grade fever, body aches and jaundice for 3-4 days. On arrival at Emergency room, he was in respiratory distress, with a respiratory rate of 40/min and on room air, oxyhaemoglobin saturation (HbSaO2) was 88-89%. Arterial blood gas analysis showed pH 7.33, PaCO2 41 mmHg, PaO2 77 mmHg (with 30% inspired oxygen concentration), HCO3 22 mEq/L and a lactate of 2.3 mmol/L. Other investigations revealed a total bilirubin of 28.2 mg/dL (direct 15.8 mg/dL), raised alkaline phosphatase (725 U/L), and raised liver enzymes (ALT 640 U/L and AST 902 U/L). The total leucocyte count (TLC) was 17,200/mm3 with haemoglobin of 8.9 gm/dL. Chest X-ray revealed lower zone homogenous opacities. He was admitted with the diagnosis of ACS and immediately given NIV with pressure support of 14 cm H2O and a positive end expiratory pressure of 5 cm H2O. He was put on antibiotics piperacillin tazobactam along with levofloxacin. After noting Verbal Analogue Score [6] (VAS) he was started on inhalational NO therapy through NIV. Each session started with NO being delivered through a face mask from a Hamilton ventilator and the NO apparatus Inosys, with the target NO level of 80 parts per million parts (ppm). A written informed consent was taken prior to administration of inhaled NO. Since NO concentration was falling below the desired level due to hyperventilation (minute ventilation >10 L/min), the patient was given midazolam intravenous boluses of 0.05 mg/kg to decrease the minute ventilation, thereby again attaining the desired NO target levels. Pain scores were monitored using verbal analogue score (VAS) of 0-10, where the patient was explained that a score of 0 meant no pain at all, and a score of 10 meant unbearable agonising pain. The patient's heart rate, blood pressure, minute ventilation, oxygen saturation were monitored

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continuously but recorded at 15 minutes interval. At the end of a 4 hour session of inhalation NO, the patient's pain score was compared to the initial pain score.

After 6 hours of inhalation of NO therapy at 80 ppm, the pain score due to ACS decreased from initial 6/10 to 1/10. He received another session of inhaled NO therapy on the third day after his admission, when the pain score decreased from 5/10 to 1/10 at the end of a 4 hour session. His repeat chest X-ray showed clearance of pulmonary infiltration in both lungs. He was discharged after 11 days of hospital stay as his condition improved.

**Case 2**

A 22 years male patient with a case of SCD (diagnosed 3 years back) presented to the Emergency department of Apollo Hospital, BBSR on 12th June, 2013 with complains of Low back pain and pain in multiple joints for 3 days, breathing difficulty and altered sensorium for 2 days. He was initially admitted in the MKCG Medical College Hospital, Berhampur (nearly 200 KM away from Bhubaneswar) and was intubated there in view of persisting respiratory distress. He was referred to us for further management. He had no history of fever/vomiting/convulsion/loss of consciousness. On arrival, he was restless, disoriented, moving all 4 limbs. GCS: E(2), V(T), M(5), BP: 140/80 mmHg, PR: 154/min., RR: 26/min, Temp: 101°F, Icterus+, Pallor+ and ABG: showed severe lactic acidosis. Investigations revealed moderate anemia (Hb: 6.9 g%), Resp. Failure with Hypeoxemia (Rt. Lower zone pneumonia in chest X-ray (A-P view), renal dysfunction: (S.Cr: 1.9 mg/dl, SK+: 7.0 mmol/L, S.Na: 123 mmol/L), hepatic dysfunction (S. Bilirubin: 8.3 mg/dl (dir: 1.7 mg/dl, indirect: 6.6 mg/dl), GGTP: 18 U/L, ALP: 476 U/L) TLC 47,800/cumm of blood, CRP: 123 mg/L, Procalcitonin >10 ng/ml, CT chest: bilateral lower lobe and one segment of right upper lobe having inhomogeneous consolidation and USG Abd: gross splenomegaly with mild hepatomegaly, globular and hypocoeic kidneys with prominent CMD suggestive of acute renal failure (ARF).

He was managed initially in ICU with mechanical ventilation and nitric oxide inhalation for 6 hours twice, extubated on 20/06/2013 after 8 days and kept on intermittent NIV support till 22nd June, 2013 along with I.V. antibiotics i.e. Inf. Meropenem, Inf. Cefoperazone + sulbactam, Inf. Targocid and Inf. Levofloxacin. He was transfused with 3 units of PRC for improving his anemia. Gradually his clinical and laboratory parameters improved, his pain subsided and he was discharged on 5th July, 2013.

**Discussion and Conclusion**

Acute chest syndrome is the most common cause of death in patients with SCD. Pain is an initial prodromal symptom of ACS and up to 50% of patients of ACS have painful crisis at presentation [7]. Dyspnea and pain in chest are the commonest symptoms of ACS in adults [8]. Therapy for ACS is mainly supportive which comprises of bronchodilators, antibiotics to prevent bacterial infections, mechanical ventilation and exchange transfusions [9].

One study has shown that sickle cell anaemia is associated with decreased NO bioactivity in both peripheral conduit as well as in resistance vessels [10]. In this study, the authors had measured the effects of intra-arterial infusion of methacholine and sodium nitroprusside (SNP) on forearm blood flow using venous occlusion plethysmography in 8 HbSS patients and 11 HbAA controls and found that the blood flow response to sodium nitroprusside was greater in HbSS patients than in controls.

It has been postulated that NO has multiple beneficial effects which enhance blood flow and suppress platelet aggregation and procoagulant substances [11-13]. In a 30 centre study involving 671 episodes of ACS in 538 patients, it was shown that nearly 50% of patients admitted with ACS had pain as the presenting symptom [14]. Patients more than 20 years of age, presence of extensive lobar pulmonary involvement and history of any cardiac disease were independent risk factors for respiration failure during hospital stay. The cause of ACS was unknown in 45.7% patients in that study with pulmonary infarct being the second leading cause (16.1%). Bacterial infections were the cause of ACS in 4.5% in patients. As pulmonary infarct was the second leading cause of ACS (apart from unknown cause contributing to the largest number of ACS cases), the significance of NO to relieve the vasospasm in pulmonary vessels (and hence pulmonary infarct) was of paramount importance as concluded in the study.

One trial has shown that there is a significant reduction with NO administration in the severity and duration of pain due to the vaso-occlusive crisis in sickle cell disease affecting children [15]. Inhaled NO reduces pulmonary artery pressures and improves ventilation-perfusion matching and oxygenation.

Some of the earlier studies have shown that NO has a beneficial effect in patients with SCD in the early resolution of vaso-occlusive crisis and acute chest syndrome [4,13]. In the study done by Reiter et al. [13] the authors had concluded that inhaled NO improves pulmonary ventilation-perfusion matching, reduces alveolar and systemic inflammation, inhibits circulating plasma haemoglobin, and therefore ultimately modulates the course of the disease. Likewise one case report mentions about successful treatment of ACS by inhaled NO in a concentration of 20 ppm for 72 hours [16]. Another case report finds that there is improvement in ACS by inhalation of NO at 80 ppm for 72 hours [17]. Likewise, yet another study revealed that inhaled NO led to lowered requirement of analgesic therapy in children [14]. Another study on twenty patients aged 10-21 years found significant reduction in pain scores in the inhaled NO group at 4 hours and less opioid requirement [4].

On the contrary, one study revealed no benefit in adults in acute pain crisis [5]. In this study with 150 patients of SCD with vaso-occlusive crisis (VOC), a lower concentration of NO with lesser ppm could have been one of the reasons for ineffective NO therapy.

The results in both the cases in which we administered inhaled NO were very good and both the patients had pain relief and complete resolution of the painful ACS. We had also reported in our earlier study the beneficial effects of NO inhalation in relieving acute pain in VOC of SCD [18]. We followed up both the patients after they were discharged from the hospital after successful NO therapy.

The advantages of inhalation NO therapy are (i) it can be easily administered in a patient via the non-invasive ventilator (ii) it leads to an early relief in painful crisis, (iii) it decreases opioid requirement and (iv) it can be given on a day-care basis i.e. after a 4-6 hours therapy, the patient can be discharged home. However, during the administration of inhalation of NO, the patient requires intensive monitoring of vitals including oxygen saturation, ventilation related parameters and NO concentration, which is dependent on patient's minute ventilation.
We recommend that there should be a multicentric clinical trial with inhaled NO in Indian patients to establish the efficacy of inhaled NO under varying conditions, its optimum concentration in inhaled mixture and its duration of exposure.

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