Review of Blood Transfusion Reactions in a Tertiary Hospital Blood Bank

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Abstract

Background: Transfusion of blood components are vital therapeutic procedures in clinical medicine. However patients may still be at risk of adverse effects of transfusions. Analysis of all untoward effects of blood transfusion must be monitored in order to correct their cause and prevent recurrence. This study was designed to analyse the incidence and spectrum of adverse effects of blood transfusion so as to initiate measures to minimize risks and improve overall transfusion safety.

Methods: In the present study, we totally reviewed data over 12 years. All the acute transfusion reactions of blood components, that were reported to the hospital blood bank were included. Reactions due to Platelets and delayed transfusion reactions were excluded. The transfusion reaction workup done for these reported cases included; verification of patient identity and clinical records, examination of blood transfusion set and bag, ABO and Rh blood grouping, cross matching (pre & post transfusion samples) and urine analysis.

Results: Of the total 293023 transfusions during the study period, 417 (0.14%) acute transfusion reactions (ATR) were reported. The commonest type of reaction noted were of the allergic type (ANHTR) (n=303; 72.6%), followed by febrile non hemolytic transfusion reactions (FNHTR) (n=104; 24.9%), 8 (1.9%) hemolytic transfusion reactions (HTR). 2 cases of the NHTRs presented with clinical suspicion of TRALI. All the HTRs were due to packed red cell (PC) transfusions. 324 NHTRs were due to red cell transfusions, 85 due to infusion of plasma.

Conclusions: The NHTRs were far more in number (esp the ANHTRs), effective leucodepletion holds the key. HTRs were completely preventable but far more dangerous clinically. A strict protocol needs to be followed not only in the blood bank, but also in other relevant procedures like; pre transfusion sampling, storage outside blood banks, bed side patient identification and monitoring of transfusion to ensure blood safety and reduce such adverse effects.

Keywords: Transfusion reactions, hemolytic, non-hemolytic.
Introduction
With the discovery of the ABO blood groups by Karl Landsteiner, the fatalities and severe adverse effects of blood transfusion reduced drastically.\(^1\) Further advances including screening for the transfusion transmitted diseases were introduced in the mid 21st century and advances in serology (including Coombs cross match) made transfusions safer. Unfortunately even, in today’s age of modern medicine; transfusion related adverse effects are still recorded. The common reasons found in literature in include alloimmunisation, Febrile, allergic reactions, volume overload, bacterial contamination and last but not the least reactions that result due to human errors.\(^2\)

Acute transfusion reactions are one of the most important factors monitored as part of Hemovigilance programs towards patients receiving transfusions. Haemovigilance is a set of surveillance procedures covering the whole transfusion chain from the collection of blood and its components to the follow-up of its recipients intended to collect and assess information on unexpected or undesirable effects resulting from the therapeutic use of labile blood products and to prevent their occurrence and recurrence. It is an important tool for improving safe blood transfusion practices in a country. The Haemovigilance Programme of India (HvPI) was launched on 10th December, 2012 in the country by under the National Institute of Biologicals (Ministry of Health and family welfare-Government of India) NIB in collaboration with the Indian Pharmacopoeia Commission (IPC). Currently, 154 centers have been enrolled in this program.\(^3\)

The need for safe blood transfusion was felt as early as 1980's and 1990's when many hemophilia patients in the developed countries contracted HCV and HIV from blood transfusions and factor concentrates.\(^4\) This dangerous example in history emphasized the need for hemovigilance. The work on hemovigilance was first initiated in France in 1991, with the setup of monitoring systems by Blood Transfusion Committees followed by the inception of Centre National d'Hemovigilance in 1992.\(^5\) Currently, on a global scale an International Hemovigilance Network (IHN) is functional, and an international database - International Surveillance of Transfusion Associated Reactions and Events has been formed to share hemovigilance data across the globe.\(^6\)

The information obtained through hemovigilance is imperative to make necessary changes in transfusion policies, for amendments in transfusion practices in hospitals and blood services, to enhance transfusion standards, to help in formulating transfusion guidelines and to improve quality and safety of entire transfusion process. As per the Ministry of Health and Family Welfare, Government of India, there are 2545 authorized blood banks in India which emphasise the need of a centralized hemovigilance system in India.\(^7\)

The scope of hemovigilance encompasses issues related to donors and blood collection, blood bank testing methods, documentation and finally patient related issues. As an initial attempt ,the present study was undertaken in order to review the transfusion protocols followed in our hospital for any possible loopholes and to document all acute transfusion reactions.

Aim
1. To analyse the incidence and spectrum of adverse effects of blood transfusion and
2. To initiate measures to minimise risks and improve transfusion safety

Materials and Methods
All recorded acute transfusion reactions from 2004 to 2016 were reviewed and included in the study. Data from 2004 to 2012 were reviewed retrospectively. Post training period (2012 onwards upto 2016) data was recorded prospectively. Repeated training sessions were held from September 2012 onwards on regular basis to the nursing staff, doctors involved with blood transfusion and blood bank technical staff.
The record of reactions reviewed included: (1) Verification of patient id, relevant clinical records, (2) Examination findings of the blood transfusion set and bag, (3) Record and review of ABO and Rh blood grouping of the blood unit, pre and post transfusion patient sample and pilot sample (collected from the donor during blood collection) (4) Analysis of Post transfusion urine sample of the patient
Reactions due to platelet transfusions and delayed transfusion reactions were excluded.

Results
Of the 293023 transfusions, in the period (2004-2016), 417 (0.14%) of Acute transfusion reactions (ATR) were recorded.

Types of reactions: While 8 were Hemolytic transfusion reactions (HTRs), the majority were NHTRs (n=409)
NHTRs were Allergic (ANHTR, n=303) and Febrile (FNHTR, n=104) types. 2 cases were clinically suspected to have TRALI.
All 8 HTRs were due to Red cell transfusions. While 278 (66.7%) NHTRs were due Packed Red cell transfusions (PC), 46 (11.03%) due to whole blood (WB) and 85 (20.9%) were due to Plasma transfusions. (Fig1)

Fig 1: Types of Blood units transfused

Correlating with prevalence of blood groups in our patient population, O blood groups were most commonly implicated (143, O Positive and 5, O Negative), followed by B group (142, B positive and 7, B Negative). The patients with Blood groups A (84, A Positive and 7, A Negative) and Group AB (28, AB Positive and 1, AB Negative) were fewer.

a) Allergic NHTRs
Allergic symptoms were classified into mild moderate and severe. 
Mild: Rashes, sweating, burning sensation, periorbital edema, abdominal distension
Moderate: chest pain, restlessness, breathlessness, vomiting, giddiness

b) Febrile NHTRs
104 cases of FNHTRs were noted with a recording of rise in temperature of > 1 deg F/C. Rigors and chills were commonly associated in FNHTRs.

c) HTRs
The 8 cases of HTR were evaluated and the cause of hemolysis is summarized in Table 1.
The most common cause was:
1. Inappropriate handling /storage after release of units from blood bank (4 cases). Use of spot light for warming blood quickly in the Neonatal ICU.
2. In 3 cases, 2 of which error were due to improper patient identification protocols being followed during collection of pre transfusion samples, sent to blood bank for cross matching. 1 was improper identification during release from blood bank.
3. 1 case was due to transfusion of a G6 PD deficient donor unit to a baby.

**Indications for transfusion were as specified in Table 2**
The most common indication for red cell transfusions was severe anemia (n=131 out of 332 units transfused) and bleeding was the most common indication for Fresh frozen plasma transfusions (n=25 out of 85 units).

**Steps taken:** Post collection of the retrospective data upto 2012, repeated training sessions with nursing staff and resident doctors were held regarding; necessary protocols for patient identification (including use of wrist bands for inpatients with barcodes), procedure for pretransfusion sampling, protocols for transfusing blood components- including monitoring and reporting of transfusion reactions. Discussions at the transfusion committee meetings with senior doctors and Nursing incharges to reinforce the right methods esp in patient identification was emphasised.

**Comparison of data between retrospective and prospective periods of study (Table 3)**
From 2013 to 2016 December, no haemolytic transfusion reactions have been reported in the hospital, indicating the importance of strictly following the above mentioned protocols.

**Table 1: Details of Hemolytic transfusion reactions**

<table>
<thead>
<tr>
<th>Patient’s group</th>
<th>ward</th>
<th>Transfusion</th>
<th>Workup results</th>
<th>Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>A Pos</td>
<td>NICU</td>
<td>A Pos 50 ml PC</td>
<td>Hemolysis in bag and Post tx** sample</td>
<td>Bag warmed in spot light</td>
</tr>
<tr>
<td>B Pos</td>
<td>PICU</td>
<td>B Pos 125 ml</td>
<td>Bag sample and Post tx** sample was hemolyed. PreTx sample-Neg*</td>
<td>Bag warmed in the spot light before transfusion</td>
</tr>
<tr>
<td>O Pos</td>
<td>NICU</td>
<td>O Pos 75ml</td>
<td>Hemolysis in Post tx** sample and urine. Bag-Neg*</td>
<td>G6PD deficient donor unit</td>
</tr>
<tr>
<td>O Pos</td>
<td>MICU</td>
<td>O Pos 200ml</td>
<td>Hemolysis in the bag</td>
<td>Transfused 21 hrs after release</td>
</tr>
<tr>
<td>O Pos</td>
<td>Pvt ward</td>
<td>AB Pos 20 ml</td>
<td>Mild Hemolysis in post tx** sample</td>
<td>Pre Sample sent from another patient for cross match</td>
</tr>
<tr>
<td>O Pos</td>
<td>FMITU</td>
<td>Given B Pos blood crossmatched for another patient with same name</td>
<td>Mild Hemolysis in post tx** sample</td>
<td>Error in blood released from blood bank</td>
</tr>
<tr>
<td>O NEG</td>
<td>Pvt wd</td>
<td>B pos 15 ml</td>
<td>No hemolysis</td>
<td>Pre Sample from another patient</td>
</tr>
<tr>
<td>O Pos</td>
<td>Outside nursing home With PPH</td>
<td>Complete unit transfused</td>
<td>Hemolysis in post tx**sample</td>
<td>Bag stored for 1 day post release in the freezer compartment</td>
</tr>
</tbody>
</table>

*Neg- No hemolysis, **tx- transfusion
Table 2: Indications for Transfusions

<table>
<thead>
<tr>
<th>Indications</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red Cells</td>
<td>332</td>
</tr>
<tr>
<td>Severe anemia</td>
<td>80</td>
</tr>
<tr>
<td>Moderate anemia</td>
<td>51</td>
</tr>
<tr>
<td>CKD</td>
<td>32</td>
</tr>
<tr>
<td>Surgery (correction)</td>
<td>80</td>
</tr>
<tr>
<td>Infections</td>
<td>14</td>
</tr>
<tr>
<td>Bleeding</td>
<td>19</td>
</tr>
<tr>
<td>Thalassemia</td>
<td>21</td>
</tr>
<tr>
<td>Malignancies</td>
<td>35</td>
</tr>
<tr>
<td>Plasma indications</td>
<td>85</td>
</tr>
</tbody>
</table>

Table 3: Comparison of data between retrospective and prospective periods of study

<table>
<thead>
<tr>
<th>Type of reaction</th>
<th>2004-2012</th>
<th>2013-2016</th>
</tr>
</thead>
<tbody>
<tr>
<td>NHTR</td>
<td>407</td>
<td>105</td>
</tr>
<tr>
<td>HTR</td>
<td>8</td>
<td>nil</td>
</tr>
</tbody>
</table>

Discussion

The overall reported rates of Acute Transfusion Reactions (ATR) range from 0.2% to 10%. In the present study, a reaction rate of 0.19% was noted, excluding platelet transfusions.

In addition, transfusion of red cells were the most common cause of ATRs in most studies. This was ratified in the present study where 79.6% of the reactions were due to red cell transfusions.

Types of reactions

a. Hemolytic Transfusion reactions:

The reported overall risks for acute HTR observed in different studies range from 0.02 – 0.07% to 3-5% per 1000 red cell units transfusions. In the Indian study by Bhattacharya et al, the reported rate was 0.23/1000 red cell units. The present study we observed a rate of 0.05 /1000 red cell units (8 cases for 151888 transfusions).

The causes of HTRs have been classified as immune and non immune. Immune causes include ABO and more commonly non ABO incompatibility eg Anti M, Anti P, Anti C, Anti K etc, as has been reported in various.

Surprisingly in the present study, all the cases were attributed to non immune causes, such as errors in Patient identification (including, both in the blood bank and Bed side), sample errors, storage errors, and after release from blood bank – storage and transport errors etc.

The other study by Bhattacharya et al also attributed 7 out of the 9 cases of HTRs to non immune causes.

It must be emphasised, to prevent these life threatening situations strict protocols must be put in place for:

1) Patient identification at the bed side. Use of bar coded wrist bands with complete patient identification has been introduced in major centres.

2) Following uniform protocols regarding handling and storage of blood products, with repeated educational reinforcement to all the concerned hospital and technical staff.

3) Monitoring blood product transfusions and prompt reporting of any untoward incidents to blood bank immediately.

b. Febrile Non haemolytic transfusion reactions

FNHTRs are defined as a transfusion reactions, observed as increase in body temperature of >1 degree C or greater unrelated to sepsis, hemolysis or other known causes of fever, that can occur during or within several hours of transfusion.

Pathophysiology: FNHTRs appear due to 3 possible underlying causes:

a) Infusion of passenger leucocytes into recipients alloimmunised against leucocytes or platelets.

b) Infusion of pyrogenic cytokines/mediators (eg IL-6, IL1 –B, aTNF alpha) that accumulate in plasma portion of the component during storage.

c) Infusion of components contaminated with bacteria/bacterial products.

d) Effective leucoreduction (esp prestorage leucoreduction) is an effective way to prevent FNHTRs.
The reported rate of FNHTR varies from 0.5 to 1%.

A comparative study on incidence of FNHTR in leucoreduced vs non leucoreduced blood components showed that the incidence was higher (0.12%) in the former group compared to the latter (0.08%) which used pre storage leucoreduced blood.

In other studies the rates of FNHTRs are 0.55 for red cell transfusions and 0.3-3% for platelets on non leucoreduced units, and 0.08-0.5% for red cells and 0.03 -0.12 % for platelets in leucoreduced units.

Our centres used buffy coat removal technique in component preparation, reducing the leucocyte load in packed red cells. In the our study FNHTR was seen for Packed red cells at the rate of 0.04% (59/149595 transfusions) and for Whole Blood transfusions at the rate of 0.25%(45/17585 units).Whole blood units showed higher rates than the red cell concentrates.

The other relevant debate found in literature was, whether pre medication with anti pyretics can be made routine for prevention of FNHTRs.

Though it is argued as an advantageous practice, one must wonder whether, the premedication may mask more severe signs of anaphylaxis, TRALI etc.

c. Allergic Non Hemolytic transfusion reactions

These reactions are type I hypersensitivity reaction in response to the proteins in the donor plasma. Most studies quote an incidence of mild allergic reactions in 0.2 to3 % of transfusions.

In the present study ANHTR was seen in 1.99/1000 units.

Severe anaphylactoid reactions characterized by respiratory distress, tachycardia or bradycardia, seizures and hypotension may be observed.

The rates for such reactions vary from 0.00212,24 in developed countries to 0.2 to 1.02 /1000 units in developing Asian countries.

In the present study 3 cases with anaphylactoid reactions were noted.

Hypotension is an important sign indicating a transfusion reaction as was noted first by Domen et al.

It was seen commonly in the study by Bhattacharya et al (50%) and the present study (36%).

Respiratory distress was another common finding and seen by Bhattacharya et al (50%) and the present study on Acute transfusion reactions had a few lacunae, as evaluation for certain other conditions like Hypovolemia, Hypocalcemia, bacterial culture for sepsis or proven cases of TRALI were not evaluated for.

On review of clinical data retrospectively, 1 patient had complained of breathlessness after transfusion, in a male general medical ward. He was kept under observation for 2 days as his oxygen saturation was <95% during that period. He was treated symptomatically and subsequently his condition improved. Retrospectively, we suspect this could have been a case of TRALI.

After the launch of the national hemovigilance program, about 765 adverse reports were submitted via hemovigil software by centers to NIB. Of 735 reports submitted between February to November 2013, 364 (49.7%) were febrile nonhemolytic transfusion reactions and 167 (22.8%) were allergic reactions. Of the reactions reported under the Hemovigilance Program of India, Not a single case of transfusion-related acute lung injury were reported which may be a result of under-diagnosis as well as under-reporting.

Despite being active, there is overall under-reporting of adverse reactions associated with blood transfusion. WHO identified that the fragmented blood transfusion systems, lack of government commitment, lack of understanding among clinicians, lack of culture of reporting, fear of punishment, lack of expertise and regulatory framework on hemovigilance, lack of computerized management system might be
challenges for the implementation of hemovigilance program in the world. Awareness among treating staff to pick up these important clinical signs and reporting it to the hemovigilance cell/blood bank must be reinforced. Proper monitoring of transfusions reporting of all such reactions will be a prerequisite for accurate calculation of ATR incidence.

Conclusions

1) Patients undergoing transfusions have to be monitored closely for any occurrence of ATRs, especially the dangerous signs of HTRs.
2) Increased awareness of treating staff and repeated reinforcement programs regarding: (i) patient identification protocols during sampling and before transfusions (ii) handling of blood components before and during transfusions (iii) monitoring of patients during and post transfusion (iv) reporting of transfusion reactions
In our experience, these interactions brought down the HTRs to nil in the past 4 years.
3) Though NHTRs are far more frequent they can be prevented effectively by measures like leucoreduction
4) The HTRs, though fewer are life threatening and therefore steps must be taken to avoid these easily preventable clinical disasters.

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