CASE REPORT

Case study of transfusion related acute lung injury in intensive care unit

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ABSTRACT

Transfusion is a common method of treatment of haemorrhagic events of patients treated in the Intensive Care Unit (ICU). Nonetheless, it involves various dangers, in many cases fatal, such as transfusion related acute lung injury (TRALI). This article refers to the case encountered during our traineeship in the ICU. It is about a 47-year-old man, who was transported to us from another tertiary acute care facility intubated due to diabetic coma. After 39 days of treatment in the ICU, the acute diabetes mellitus, the hemodynamic instability and the electrolyte disorders were regulated. However, he started to have diffuse haemorrhagic events due to intestinal necrosis and required transfusion of blood factors. After the transfusion of Platelets (PLT) he developed Acute Respiratory Distress Syndrome (ARDS), which evolved to TRALI and finally the death of the patient occurred.

Key Words: Transfusion related acute lung injury, Blood transfusion reaction, Lung Injury, Intensive Care Unit

1. INTRODUCTION

It is an unquestionable fact that nowadays science is constantly evolving. TRALI even today remains a fatal syndrome, which is often confused with ARDS. It is a rare complication following transfusion of blood and blood products and in particular whole blood, Packed Red Blood Cells (PRBC), Platelets (PLT) and Fresh Frozen Plasma (FFP).[1, 2] It is a non-cardiogenic pulmonary oedema that often occurs within the first six hours following blood components transfusion. Its frequency, according to international literature, is estimated to occur in 1:5000 transfusions with fatality 5%-10% of the total number of transfusions.[2-6] The etiology of the syndrome remains not fully understood; due to that health professionals should report the events ending to TRALI, as well as their way of treatment in order to acquire some type of “experience”. In the future, this will improve the possible approaches in order to save the patient’s life.

2. CASE REPORT

A 47-year-old man was transported to us from a tertiary acute care facility. Prior to this event the patient did not have any severe health problems. The only pathosis were arterial hypertension and hypertriglyceridaemia, which were regulated by prescribed medication. One week before hospitalisation he developed clinical symptoms of polyuria, polydipsia and difficulty in speech. Upon arrival at the first hospital he was intubated due to diabetic coma: blood glucose was 1300 mg/dl, glycated haemoglobin was 15.5%, serum creatitine 5.0 mg/dl and his blood pressure was 77/27 mmHg. He developed metabolic acidosis with pH = 7.15 and hemodynamic instability under vasopressor agents.

The first problem encountered by the patient were the dia-
The patient during the first days of hospitalisation presented clinical examination, which called for brain MRI that detected diffuse anoxaemic damage. The results of laboratory tests showed decreased white blood cells \((1500 \times 10^9/\mu l)\).

Due to persistent fever, a tracheotomy was made with the creation of temporary tracheostomy and change of central venous catheter. Moreover, a gas tube was placed due to severe diarrhoeic evacuation. However, neurologic improvement was noticed (automatic cough, slight finger movement of the upper limbs to pain stimuli, isocoria, eye movement upon command, expression on pain). The communication with the patient was gradually re-established and he was able to “contact” people surrounding him, even watch television.

During the following ten days of his hospitalisation, his fever became higher \((38.5^\circ C-39.3^\circ C)\), without corresponding to cold compresses and paracetamol administration. Due to many hypotensive episodes a blood work of FT4 \((4.8)\), FT3 \((0.9)\), FSH \(<0.3)\), LH \(<0.4)\) and TSH \(<0.186)\) was ordered that suggested Pituitary deficiency. Additionally, due to low FHS and LH a brain MRI took place that confirmed the first hypothesis. Many purulent and bloody secretions appeared in the lower respiratory system during suction. Additionally, he was transfused for two consecutive days \((13^{th}-14^{th})\) with 1 Unit of Packed Red Blood Cells (PRBC) each day, due to low haemoglobin. On the 14th day the patient was gradually released from the mechanical respiratory support (placement of T-tube) with satisfactory level of arterial blood gases.

The persistent high fever led to research of the underlying cause with the following actions:

1. Change of central venous catheter and culture of its tip
2. Interruption of Pentaglobin (immunoreaction)
3. Thorax - Abdomen CT
4. Gastroscopy and Proctoscopy

The Gastroscopy revealed several oesophageal ulcerative lesions, which were attributed to the presence of nasogastric feeding line (Levin). These actions did not reveal the etiology of persistent fever.

Transfusions of PRBC continued at regular intervals until the day he died due to seriously low haemoglobin \((Hb = 5.8 \text{ g/dl})\). The cardiac rhythm alternated between atrial fibrillation and sinus rhythm with extrasystoles.

For further and more detailed examination of the lower gastrointestinal (bloody stools), the 16th day of hospitalization anosigmoidoscopy – colonoscopy were done, which showed ischemic rectosigmoiditis.

On the 22nd day incidence of hypoxaemia, numerous haemorrhagic diarrhoeic evacuations (placement of flexi-seal) were noticed due to increased prothrombin time \((18.1 \text{ sec})\), INR \((1.75)\) and APTT \((46.2 \text{ sec})\), as well as extensive oedema.

Table 1. Objective examination of the patient on his admission to hospital

| Objective examination          |
|-------------------------------|
| Cold limbs (hypotension–tachycardia) |
| Miosis (sedation)                  |
| Rx thorax: infiltrates in both sides |
| HR \(\geq 130/\text{min})        |
| Abdomen: soft/rare intestinal sounds |
| Oliguria to anuria               |

Table 2. First assessment of the patient on his admission per system

| First Assessment   |
|--------------------|
| C.N.S.: intubated in sedation–hemodynamically unstable in SN (Sinus Node), anisocoria, lack of pupillary reflexes |
| Respiratory: probable aspiration during intubation at the previous hospital–image of ARDS on chest X-ray |
| Digestive: Levin → administers gastric - bloody content |
| Urinary: oliguria to anuria |
| Skin: pale, cyanotic, dry |
| Vital Signs         |
| Blood Pressure: 77/27 mmHg |
| Pulse: 127/min       |
| Body temperature: 37.0°C |
| \(\text{SpO}_2\): 91% |

During the first 48 hours after sedation was discontinued, the patient did not show any neurologic improvement during clinical examination, which called for brain MRI that detected diffuse anoxaemic damage. The results of laboratory tests showed decreased white blood cells \((1500 \times 10^9/\mu l)\).
The following day, a thorax–upper and lower abdomen CT was done to the patient, which showed an image of air leakage. The same day, during palpation, acute surgical abdomen was noticed and the patient was delivered to the operating room for exploratory laparotomy. Necrotic surfaces on the large intestine were detected, which were removed and a colostomy was formed.

The first day after the surgery the patient did not produce sufficient quantity of urine, he developed serious hypokalaemia and was placed under continuous extrarenal clearance. The 32nd day of hospitalization because of bowel perforation and peritonitis a second surgery took place (colon laparotomy). In that surgery local petechial lesions of the small bowel serosa were observed. Nonetheless, on the 37th day the patient underwent a third surgery, this time an exploratory laparotomy due to ischemic lesions of the colon. A total colectomy and ileostomy was performed.

On the 35th day of hospitalization the patient other than intestine bleeding showed intensive posterior epistaxis from the left nasal fossa and blood flow in the throat, therefore nasal tamponade was placed.

### Table 3. Vital signs and Arterial Blood Gases (ABG) of the last 24 hours

| Hours   | Blood Pressure (mmHg) | Temperature (°C) | Heart Rate (/min) | SaO₂ (%) | PaO₂ (%) |
|---------|-----------------------|------------------|-------------------|-----------|----------|
| 07:00   | 140/68                | 36.7             | 85                | 83.7      | 52.3     |
| 10:00   | 146/70                | 36.8             | 89                | 89.0      | 58.9     |
| 13:00   | 138/62                | 36.7             | 83                | 83.9      | 130      |
| 15:00   | 140/65                | 37.9             | 85                | 85.3      | 50.1     |
| 16:00   | 141/65                | 38.0             | 86                | 82.5      | 49.2     |
| 18:00   | 142/68                | 37.6             | 87                | 76.0      | 44.1     |
| 21:00   | 140/63                | 36.8             | 85                | 75.1      | 51.5     |
| 23:00   | 95/41                 | 36.8             | 87                | -         | -        |
| 01:00   | 85/35                 | 37.0             | 82                | 70.9      | 120      |
| 02:00   | 64/30                 | 36.7             | 38                | 67.5      | 46.1     |
| 03:00   | 51/12                 | 36.6             | 30                | 47.9      | 36.1     |
| 04:00   | 155/33                | 37.0             | 99                | 65.1      | 45.9     |
| 05:00   | 100/42                | 37.3             | 55                | -         | -        |

During the last week, because multisystemic intensive bleeding continued, transfusions with PRBC, FFP and PLT were done. At the 38th day of hospitalization at 19:00 the patient was transfused with 2 PLT. Four hours after the transfusion, as shown in Table 3, the patient showed severe hypotension and hypoxia. Symptoms during the next three hours intensified, despite fully supported mechanical ventilation and inotropic support. At 02:00 the patient had severe bradycardia with hemodynamic instability and he was administered 2mg of atropine and 1mg of adrenaline. At 03:00 he showed cardiac asystole and chest compressions along with high inotropic support started, without response. After two hours of ongoing effort the patient died.

### 3. DISCUSSION

In 1818 James Blundell performed the first blood transfusion. It has been a long time since then and procedures have become safer for patients, thus reducing the danger of transmission of diseases and complications.[5]

Only in 2004 TRALI was officially recognised during a conference held at Toronto by the National Institute of Health as a severe complication of blood transfusion and was clinically separated from Acute Lung Injury (ALI) which until then was not connected to it. The clinical features of TRALI are dyspnea, hypoxaemia and infiltrations of lungs in both sides that appear on chest X-ray.[7–11] There are no reports for a differential frequency of appearance of TRALI according to gender or the age of the patient.[2]

Pathophysiology of TRALI is ambiguous and has not been fully understood. However, its etiology is based on the well-known “two-hit” hypothesis which includes the clinical condition (pre-existing pathology – first hit) of the patient and the transfusion of blood products, especially plasma (second hit).[2,11–14]

The vast majority of cases mention that the pre-existence of HLA class I, class II and HNA antibodies in the plasma of the donor, react with the neutrophils of the already very sensitive pulmonary mucosa. In a very smaller percentage of cases, it is reported that the pre-existing heavy pathology of the patient could result in a systemic inflammation of lungs which leads to the accumulation of neutrophils in their microvasculature. These neutrophils are highly possible to react to lipids or other mediators (CD40L) accumulated in the blood cell components during its storage and contribute to the endothelial damage of the patients in severe condition, thus causing vascular leakage and pulmonary oedema. A different hypothesis claims that mtDNA (mitochondrial DNA) DAMPs (Damage-Associated Molecular Pattern molecules) appear in PRBC, FFP and PLT, possibly contributing to the creation of TRALI. However, further research is required for the proof of this hypothesis.[2,12,13,15]

In many cases TRALI is confused with other diseases. Therefore, possible causes of acute pulmonary oedema should be excluded, which may be acute haemolytic reaction, pneumonia and cardiogenic pulmonary oedema.[5] Following the exclusion of the above, the most basic and most differential diagnosis from TRALI is TACO (Transfusion-Associated
Circulatory Overload), which is characterised by circulatory overload. Despite the fact that there are not specific examinations for TRALI diagnosis, the following clinical examinations may confirm the suspicion of its presence:

- Echocardiogram
- White blood cell count (WBC)
- Brain natriuretic peptide (BNP)
- Pulmonary oedema fluid protein analysis[11]

Supportive care is the only treatment for patients suffering from TRALI, which is confirmed both by a wide literature and clinically.[5] Use of diuretics are indicated when the patient suffering from TRALI presents circulatory overload in order to improve their condition, but they should be avoided in other cases.[16,17] In the above described case there were oedemas and positive fluid balance, so the use of diuretics preceded the appearance of TRALI and continued until the end.

As there is not specialised treatment of TRALI, the health professionals should focus on prevention measures. According to scientific research, prevention is based on three main measures. The first one focuses on the strict selection of donor. In 2003 the English National Blood Service applied the exclusion of women from FFP donation and within 6 years significant reduction of TRALI patients was noticed.[2,17–20] The same measure is applied in Greece despite the lack of registered percentages. The second measure concerns the strict methods of storage of blood and its products, especially FFP. The third and last measure refers to the avoidance of unnecessary transfusions.[13,18,21,22]

The case presented in this article fully confirms the theory of the “two-hit” model. In particular, the impaired state of the patient combined with the last PLT transfusion resulted in TRALI. The vital signs and ABG of the patient as developed within the first six hours following the transfusion confirm this syndrome based on literature. The diagnosis of the syndrome could be challenged, because of the constant transfusions during hospitalization. Nevertheless, this case follows the course of development of TRALI, with intense and sudden drop in blood pressure, hypoxia-despite mechanical ventilation—and fever that was treated with antipyretic drugs.

4. CONCLUSION
TRALI is considered one of the most basic reasons of death in the ICU due to onset of non-cardiogenic pulmonary oedema, a fact that is also claimed by numerous articles, as well as by this case. The analysis and study of the case led to the conclusion that both physicians and nurses, who are responsible for the treatment of seriously ill patients should be able to recognise and proceed to differential diagnosis of TRALI correctly and on time.

In conclusion, it should be emphasized that in every blood and blood products transfusion (PRBC, FFP, PLT), regardless of the number of transfusions, health professionals should be vigilant for the possible development of TRALI.

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CONFLICTS OF INTEREST DISCLOSURE
The authors declare that there is no conflict of interest statement.

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