February 2018

Bladder distension as a cause of abdominal compartment syndrome

Muhammad Yasir  
Security Forces Hospital, Riyadh, Saudi Arab

Muhammad Qamarul Hoda  
Aga Khan University, qamar.hoda@aku.edu

Tauseef Ahmed  
Manchester Royal Infirmary Hospital, Oxford Road, Manchester

Follow this and additional works at: https://ecommons.aku.edu/pakistan_fhs_mc_anaesth

Part of the Anesthesiology Commons, and the Urology Commons

Recommended Citation
Available at: https://ecommons.aku.edu/pakistan_fhs_mc_anaesth/130
INTRODUCTION

Abdominal compartment syndrome (ACS) has been increasingly recognized in critically ill patient and its harmful effects are well documented. The disparity among the pressure, volume in abdominal cavity and its contents, results in ACS. The actual incidence of ACS is not known. However, it has been observed predominantly in patients with severe blunt and penetrating abdominal trauma, ruptured abdominal aortic aneurysms, retro- and intra-peritoneal hemorrhage, pneumoperitoneum, neoplasm, pancreatitis, ascites and multiple bone fracture. We present a case of 40-year female who underwent emergency cesarean section and developed abdominal compartment syndrome due to urinary bladder distension secondary to blockade of urinary catheter with blood clots. This is a very unusual cause of ACS.

CASE REPORT

We report a case of abdominal compartment syndrome caused by postoperative urinary bladder distension secondary to blockade of urinary catheter with blood clots in a 40-year female patient who underwent emergency cesarean section. As per our knowledge, this is the only case report of abdominal compartment syndrome due to bladder obstruction.

A 40-year female with no known comorbid condition, was planned for elective cesarean section due to type IV placenta previa, polyhydramnios and poor obstetric history. Two days earlier to her planned procedure, she presented in labour room with a history of four hours of labour pain for which emergency lower cesarean section was undertaken. Her preoperative assessment was normal and she underwent surgery under general anaesthesia.

Intraoperative course was complicated by massive blood loss (6-7 liters) for which she was given 13 units of packed red blood cell (PRBC), 11 units of fresh frozen plasma (FFP), 8 units of platelets, 8 units of cryoprecipitate, 12 litres colloid, and 3 litres of crystalloid. Hemostasis was difficult to achieved so a cesarean hysterectomy was performed as a rescue. A diagnosis of disseminated intravascular coagulopathy was made clinically; abdominal closure was difficult, therefore, the wound was left open and covered with soaked abdominal packs. She required ionotropic support with dopamine at 20 mcg/kg/min and epinephrine at 0.05 mcg/kg/min. She was electively ventilated postoperatively. Patient was kept sedated. On inotropic support with dopamine and epinephrine, she was maintaining her systolic arterial pressure between 90-110 mmHg, with a CVP of 5-7 cm H2O. Her abdominal girth was monitored hourly. She received 6 units of red cells concentrate, 10 units of cryoprecipitate, 4 units of fresh frozen plasma and six units of platelets within first
18 hours postoperatively. She became oliguric followed by an increase in the abdominal girth from a baseline of 93 cm to 101 cm. Subsequent to that, her CVP also dropped to 2 cm H$_2$O and she became severely hypotensive with a blood pressure of 56/32 mmHg and heart rate went up to 135/minute. A surgical review was sought immediately and provisional diagnosis of intra-abdominal bleeding was made. For the management of this life-threatening peri-arrest condition, she was immediately transferred to operating room. Re-exploratory laparotomy was negative for any active bleeding with minimal amount of intraperitoneal blood stained fluid. The most striking finding was an enormously distended urinary bladder, extending from pubic symphysis up to umbilicus, which was confirmed by needle aspiration. On examination, it was found that urinary catheter was blocked with blood clots which was replaced and drained about 1400 ml urine.

Following the decompression of urinary bladder, immediate improvement was noticed in her blood pressure with decreasing requirement of dopamine. Her blood pressure increased to 140/70 mmHg and heart rate came down to 110 per minute, and the urine output increased to 200 ml/hour. The peak airway pressure also dropped from 30 cm H$_2$O to 20 cm H$_2$O. Her CVP increased to 12 cm H$_2$O. It was decided to put abdominal pack and not to close the abdominal wound at this stage. Patient was transferred to post anaesthesia care unit for elective ventilation. On the very next day, her inotropes were successfully tapered off. Two days later, patient was again taken to operating room for removal of abdominal packs and wound closure. She remained stable and was extubated on the next day and shifted to high dependency unit and later discharged home after 10 days in good condition.

**DISCUSSION**

ACS is a disorder associated with significant morbidity and mortality, refers to organ dysfunction resulting from increased intra-abdominal pressure (IAP). ACS, defined by world society of abdominal compartment syndrome as intra-abdominal pressure of at least 20 mm Hg with dysfunction of at least one thoraco-abdominal organ. This increase in intra-abdominal pressure leads to organ dysfunction involving primarily heart and lungs. The reason for cardiac symptoms are multifactorial with decreased preload due to compression of both the portal vein and inferior vena cava, and an increased afterload due to increased systemic vascular resistance, leading to a decreased stroke volume and thus decreased cardiac output and cardiac arrest with pulseless electrical activity (PEA). The main reasons for respiratory compromise are the combination of cephaled displacement of the diaphragm, with resultant decrease in total and residual lung volume and lung compliance.

The standard recommended method for diagnosis and management of IAH and ACS is based on accurate and repeated measurements of its surrogate pressure; intra-vesical urinary bladder pressure. But in peri-arrest situations, when there is no time to measure IAP, clinical signs of organ dysfunctions, and hypoperfusion with low cardiac output, hypoxia, tense distended abdomen, progressive oliguria or anuria are sufficient to justify emergency decompression.

In this case, the diagnosis was also made by clinical parameters only because of deteriorating and near crash situation which developed suddenly. The intra-vesical pressure was not measured as she had to be rushed to operating room while cardiopulmonary resuscitation was actively going on.

This patient had signs and symptoms that could suggest hypovolemic/septic shock (tachycardia, severe hypotension, tachypnea, fever and decreased urine output). However, following emergency surgical decompression, the patient had a prompt normalization of her vital signs and urine output. In view of the above, a distended urinary bladder, due to catheter block, was the most likely diagnosis of her deterioration. There was no case report of ACS secondary to urinary catheter obstruction.

**REFERENCES**


---

Muhammad Yasir, Muhammad Qamarul Hoda and Tauseef Ahmed