Intravascular absorption syndrome : case report of a life-threatening complication during hysteroscopy

R. Niewenhuizen (*), A.F. Kalmar (*), N. Van Der Vekens (*), J. Heerman (*), M. Casteels (**), H. Vanoverschelde (*)

Abstract : A 40-year old, healthy woman underwent hysteroscopic endometrial ablation with a bipolar electrocautery using 0.9% saline as distension fluid. After 45 minutes of surgery, arterial oxygen saturation decreased and liquid was obstructing the laryngeal mask. With an estimated total fluid deficit of 5000 mL, fluid overload was suspected and the patient’s trachea was intubated. Furosemide was given intravenously and a urinary catheter was inserted. The intra-abdominal pressure, measured through the urinary catheter, was 28 mmHg. As ventilation became unfeasible, the patient became cyanotic and no clear pulse could be detected, cardiopulmonary resuscitation was started. In order to decrease the abdominal pressure, a laparotomy was performed and extracorporeal membrane oxygenation was started to increase the arterial oxygenation. After one week in the Intensive Care Unit, the patient was extubated and gradually recovered without further complications nor residual morbidity.

The pathophysiological aspects of the evolution to severe pulmonary edema due to massive fluid translocation during operative hysteroscopy, and the rationale behind the successful interventions are being discussed.

Close continuous monitoring of the amount of fluid deficit should be performed to avoid severe fluid overload during operative hysteroscopy. Extracorporeal membrane oxygenation can be life-saving to bridge the period of desaturation until standard treatment suffices to provide adequate oxygenation.

Key words : Intravascular absorption syndrome, volume overload, hysteroscopy, pulmonary edema, ECMO.

INTRODUCTION

Operative hysteroscopy has successfully replaced hysterectomy for multiple indications, including treatment of congenital anomalies, endometrial polyps, cervical and uterine neoplasia and submucous myomas (1). Although complications are rare (incidence <1%)(2), anesthesiologists and surgeons should be aware that, even during such a minimally invasive procedure, life-threatening complications can suddenly arise. We report a case of massive pulmonary edema caused by severe fluid overload during operative hysteroscopy.

CASE REPORT

The patient gave written consent to publish this case report, read the article and confirmed its content. A 40-year old, healthy female underwent an operative hysteroscopy for endometrial ablation under general anesthesia. She had no significant medical history and preoperative laboratory test results were normal. After induction of anesthesia with propofol and sufentanil, a laryngeal mask airway was placed, and anesthesia was maintained with sevoflurane. Monitoring consisted of electrocardiography (ECG), end-tidal carbon dioxide (EtCO₂), non-invasive blood pressure (NIBP) and pulse oximetry (SpO₂). The patient was continuously in supine position with the legs elevated and placed in the leg holders. Saline 0.9% was used as distention and irrigation medium with an irrigation pressure of 350 mmHg. A total of 12000 mL of rinsing fluid was instilled, and at the end of the 30-minute procedure, the recollected volume was 7000 mL. The endometrial resection with bipolar electrocautery necessitated coagulation of a wide area of the endometrial surface due to significant blood loss in the cavum uteri. During the total procedure, 2-3 instrument insertions were performed for specimen removal.

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After an uneventful 30 minutes of surgery, the tidal volume (TV) and EtCO₂ decreased from 230 to 150 mL and from 34 mmHg to 24 mmHg, respectively, over a period of 10 minutes, and the SPO₂ and NIBP also gradually decreased. The drapes were removed (Figure 1, arrow) from the face for visual inspection and chest auscultation, immediately showing signs of facial oedema with severely thickened and cyanotic lips. Further evaluation showed a general oedema of the neck, conjunctiva and face, and protrusion of the tongue. Chest auscultation revealed significantly dampened symmetrical ronchi and crepitations. The abdomen was tense but compressible. The auscultation and the absence of subcutaneous emphysema suggested a pneumothorax unlikely. Closer examination revealed clear liquid coming out of the laryngeal mask. Pulmonary edema due to severe fluid overload was suspected and an endotracheal tube was placed. Cardiac and abdominal echography revealed a normal ventricular function, no signs of pericardial effusion, and no evidence of ascites. Further examination revealed clear liquid coming out of the laryngeal mask. Pulmonary edema due to severe fluid overload was suspected and an endotracheal tube was placed. Cardiac and abdominal echography revealed normal ventricular function, no signs of pericardial effusion, and no evidence of ascites. Cardiac and abdominal echography revealed normal ventricular function, no signs of pericardial effusion, and no evidence of ascites.

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The pathophysiology of pulmonary edema can be understood as the net result of the startling forces across the pulmonary microvasculature, where the net driving force (NDF) is governed by tissue permeabilities, the transmural hydrostatic pressure difference and the oncotic pressure difference. The transmural hydrostatic pressure force is determined by the fluid permeability of the capillary wall (K̅h), the capillary surface area (A), and the difference between the capillary (Pc) and interstitial (Pi) hydrostatic pressure, while the oncotic pressure force is determined by the reflection coefficient for plasma proteins (η) and the difference between the capillary (xc) and interstitial (xi) oncotic pressure.

The NDF of fluid across the pulmonary capillary membrane is therefore governed by the net balance between the transcapillary hydrostatic pressure (Pc-Pi), causing fluid to flow out of the capillary, and the transcapillary protein osmotic pressure (xc-xi), which acts to retain fluid within the capillary.

As such, different pathophysiological entities of pulmonary edema exist: permeability edema, caused by a disruption of the normal alveolar-capillary barrier, and hemodynamic pressure edema, caused by an increase in the capillary hydrostatic pressure difference or a decrease in the capillary oncotic pressure. In this case of volume overload, the pulmonary edema was primarily induced by a combined increase in capillary hydrostatic pressure and a decrease in capillary oncotic pressure due to dilution of the plasma proteins with saline 0.9%. In addition, the increased hydrostatic capillary pressure and increased hydraulic flow through the capillary glycocalyx is described to deform the endothelial

### Table 1

<table>
<thead>
<tr>
<th>Laboratory Parameter</th>
<th>Normal values</th>
<th>10 min before start</th>
<th>10 min after start</th>
<th>10 min before start ECMO</th>
<th>25 min after start ECMO</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>7.35 - 7.45</td>
<td>6.90</td>
<td>6.93</td>
<td>7.04</td>
<td>7.32</td>
</tr>
<tr>
<td>PCO₂ (mmHg)</td>
<td>35 - 48</td>
<td>74</td>
<td>81</td>
<td>66</td>
<td>34</td>
</tr>
<tr>
<td>PO₂ (mmHg)</td>
<td>83 - 108</td>
<td>46</td>
<td>46</td>
<td>66</td>
<td>78</td>
</tr>
<tr>
<td>Base excess (mmol/L)</td>
<td>-2.0 - 3.0</td>
<td>-18.5</td>
<td>-19.5</td>
<td>-13.2</td>
<td>-7.8</td>
</tr>
<tr>
<td>Sodium (mEq/L)</td>
<td>136 - 145</td>
<td>141</td>
<td>141</td>
<td>143</td>
<td>143</td>
</tr>
<tr>
<td>Potassium (mEq/L)</td>
<td>3.5 - 5.1</td>
<td>1.9</td>
<td>2.2</td>
<td>1.9</td>
<td>2.2</td>
</tr>
<tr>
<td>Chloride (mEq/L)</td>
<td>98 - 117</td>
<td>117</td>
<td>115</td>
<td>115</td>
<td>115</td>
</tr>
<tr>
<td>Calcium (mEq/L)</td>
<td>1.11 - 1.23</td>
<td>0.62</td>
<td>0.88</td>
<td>0.77</td>
<td>0.94</td>
</tr>
<tr>
<td>Glucose (mg/dL)</td>
<td>74 - 106</td>
<td>219</td>
<td>206</td>
<td>216</td>
<td>205</td>
</tr>
<tr>
<td>Lactate (mg/dL)</td>
<td>2.24 - 6.75</td>
<td>29</td>
<td>26</td>
<td>24</td>
<td>25</td>
</tr>
<tr>
<td>Haemoglobin (g/dL)</td>
<td>12.0 - 15.0</td>
<td>10.6</td>
<td>11.0</td>
<td>11.0</td>
<td>11.0</td>
</tr>
</tbody>
</table>
glycosaminoglycan fibers, which activates the endothelial nitric oxide synthase, leading to barrier dysfunction with eventually additional permeability edema (11).

A fluid deficit of no more than 2500 mL fluid has been recommended, and guidelines prescribe that the surgery should be stopped if this is exceeded (4,12,13). In the presented case, because of focused attention on challenging hemorraghic control, the surgeon only became aware of the fluid deficit at the end of this relatively short procedure. Intrauterine bleeding is conventionally controlled by coagulation, while also the pressure of the distention media themselves may yield enough pressure to cause hemostasis during a procedure. In the literature, procedures with a particular risk of intraabdominal fluid are reported to be resection of fibroids or myometrial trauma that results in open uterine venous channels or unidentified perforations (14). Still, in this procedure for endometrium ablation – albeit over a large surface – very fast translocation of irrigation fluid occurred. A noteworthy consideration is that the presence of adenomyosis may have contributed to the exceptional fluid intravasation. Limiting the pressure of the irrigation fluid will correspondingly decrease the amplitude of the fluid shift. In this case, the irrigation fluid pressure was set at a level greatly exceeding recommended pressure (13). Secondly, close monitoring of the fluid balance during and after each hysteroscopic surgery is therefore, necessary to timely detect and avert dangerous complications. Whereas isotonic irrigation fluid bears a minimal risk of severe electrolyte disturbance, the risk of fluid overload still rationalizes the use of precise irrigation instruments with accurate pressure and inflow/outflow measurement. Moderating the irrigation pressure to the strictly necessary level for adequate visualisation and working conditions will both be important of moderation of the fluid distension media in operative hysteroscopy. Gynecol. Surg. 13: 289-303.

When observing the clinical deterioration as described, the anaesthesiologist should be aware of alternative causes of hemodynamic collapse. While in this case fluid overload was quickly considered most plausible, given facial swelling and echography information, anaesthesiologists and surgeons must be vigilant towards the venous air embolism (15) (VAE) or pneumothorax diagnosis, and consider the primary symptoms as observed in this case as caused by VAE until proven otherwise, in which case a completely different therapeutic approach would be mandatory. Particularly during procedures such as hysteroscopic endometrial resection with pressurised irrigation fluid, a large resection surface, electrocautery and repeated instrumentations, VAE is probably more common than anticipated by many. VAE will cause a sudden drop in ETCO2, followed by a gradual decline in SpO2. In hypervolemic pulmonary edema, the decrease in ETCO2 and SpO2 will both be gradual. Also bronchospasm, pulmonary oedema and cardiac arrest can occur after VAE. In our case, echocardiography rapidly confirmed fluid overload. We advise that echocardiography is performed swiftly in such circumstances to help establish the diagnosis: a dilated right ventricle would suggest pulmonary embolism, biventricular dilatation would suggest hypervolemia and air-bubbles in the right ventricle and pulmonary artery would suggest air embolism (15).

When life-threatening hypoxemia due to fluid overload pulmonary edema occurs, ECMO can temporarily be used until edema reduction measures are effective and conventional intensive support suffices.

Conclusion

While hysteroscopy is mostly a low-risk routine procedure, this case emphasizes the importance of moderation of the fluid pressure, continuous monitoring and vigilance for fluid overload, and of the close availability for immediate cardiopulmonary support and intensive care therapy.

References