Abdominal compartment syndrome caused by gastric distension in bulimia nervosa and fatal injury following surgical decompression - A case report -

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Background: Abdominal compartment syndrome (ACS) occurs due to increased abdominal cavity pressure, causes multiple organ damage, and leads to fatal consequences. Increased intraperitoneal pressure due to different reasons generally does not result in serious damage, due to the compliance of the abdominal wall. However, when the pressure exceeds the limit, ACS develops, thereby causing fatal damage to the organs.

Case: A patient presented with rapid stomach swelling due to excessive food intake and was known to have bulimia nervosa, which had now resulted in ACS. Mental changes, abdominal distension, color change in the legs, acute kidney injury, and acidosis were seen. The patient expired due to ischemia-reperfusion injury and disseminated intravascular coagulation, which occurred after surgical decompression.

Conclusions: Under suspected ACS conditions, we should be aware of various symptoms that can occur. Early attempts for decompression are helpful, and it is important to be prepared for reperfusion injury prior to surgical decompression attempts.

Keywords: Bulimia; Intra-abdominal hypertension; Reperfusion injury; Shock, hemorrhagic.
The patient was diagnosed with major depression and bulimia nervosa nine years ago. The patient also had a history of frequent vomiting after episodes of binge eating, along with numerous suicide attempts. Ten hours before reporting to the hospital, the patient ate a very large quantity of food, equivalent to approximately ten portions; however, the quantity of the vomit was much less, relative to the amount of food taken. The physical examination revealed abdominal pain and distension in the entire region, along with tenderness, but no rebound tenderness. Subsequent abdominal computed tomography (CT) revealed clear gastric distension. The results of various blood tests were normal. In the joint general surgery consultation that followed, it was decided that an immediate surgical treatment was not necessary; hence, the need for continuous observation of the condition was explained, and a gastroenterological or psychiatric hospitalization was advised, but both the patient and her guardians declined hospitalization. Afterward, they were warned of the possible complications, such as electrolyte imbalance, aspiration pneumonia, gastrointestinal perforation, ischemic changes, due to the pressure applied by gastrointestinal distension, and the possibility of death, in case the condition worsened. The self-discharge form was signed, and they returned home.

Seven hours after returning home, the patient re-visited the emergency room due to persistent abdominal pain, and mental confusion. At the time of the visit, the patient was unconscious. On physical examination, a more severe abdominal distension, and signs of abdominal rigidity were observed. Both legs of the patient had turned pale in color. No auscultation sounds were heard over the abdomen, and the dorsalis pedis pulse was not felt on either foot. Shock was suspected due to a 60/40 mmHg blood pressure, 160 beats/min heart rate, 22 times/minute aspiration rate, and 36.4°C temperature. Other findings included pH 7.076, PaCO₂ 22.3 mmHg, PaO₂ 122 mmHg, HCO₃⁻ 6.3 mM/L, base excess (BE) 23.7 mM/L, and lactic acid 11.1 mM/L, which were measured in the arterial blood gas analysis and, thus, metabolic acidosis was confirmed. The electrolyte levels Na⁺ 157 mM/L, K⁺ 6.2 mM/L, and Cl⁻ 130 mM/L, indicated that the electrolyte imbalance was severe (Table 1). In addition, the patient was in a severe hypoglycemic state with a blood sugar level of 15 mg/dl; because of an elevated creatinine level of 2.84 mg/dl, acute renal failure was suspected. First, to treat the metabolic acidosis, sodium bicarbonate was set at 80 mEq/h. In the abdominal X-ray, the gastrointestinal tract (GIT) was seen to be filled with a large amount of food but without any bowel gas (Fig. 1). Abdominal CT re-

![Image](image_url)

**Fig. 1.** The plain abdominal X-ray image shows markedly dilated state in stomach with abundant food materials.

<table>
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revealed a very large dilation of the stomach, esophagus, and the second part of the duodenum; it was suspected that the distended organs were pressing against the descending aorta, and the right kidney was pressed upwards (Fig. 2). One hour after the emergency room re-visit, the patient’s mental state became lethargic—communication became impossible and spontaneous respiration decreased; as SpO₂ was measured to be 88%, endotracheal intubation was performed. A Foley catheter was then inserted, but no urine was drained; an emergency hemodialysis was prepared, and for enough fluid supply, a central venous catheter was inserted into the right jugular vein. For continuous monitoring of changes in the vital signs, a conduit was placed in the right radial artery. An attempt was made to insert a nasogastric tube to relieve gastrointestinal pressure, but the food was not drained due to a blockage near the esophagus, which made further insertion impossible. An abdominal CT performed for follow-up observation revealed that the abdomen was still filled with food, and the nasogastric tube was twisted at the end of the esophagus (Fig. 3). It was also suspected that insertion was not possible due to gastroesophageal junction deviation or esophageal stenosis; however, no such sign was seen.

There was no decompression through the nasogastric tube, and as time passed, the mental state of the patient changed into a state of confusion. The patient’s vital signs became increasingly unstable. Since the symptoms and follow-up blood test results exacerbated, instead of performing a percutaneous drainage, it was decided to attempt a surgical decompression; subsequently, the patient was transferred to the operation room six hours after the re-visit. In the emergency room, a total of 3,170 ml of crystalloid fluid, in which a 5% glucose solution was included, was administered. Urine volume was not measured.

After reaching the operation room, the patient’s blood pressure was 70/46 mmHg and the heart rate was 128 beats/min. First, in order to improve the vital signs, crystalloid fluid was administered at the highest speed via the right arm’s peripheral venous catheter and via the central venous catheter of the right internal jugular vein. The in-

![Fig. 2. The abdomen-pelvis computed tomography axial (A) and coronal (B) images revealing markedly distended stomach, dilated distal esophagus and proximal duodenum.](image1)

![Fig. 3. Axial (A) and coronal (B) images of the abdomen-pelvis computed tomography (CT). CT shows coiling of L-tube tip in distal esophagus.](image2)
stillation of norepinephrine was initiated with sustained dosing at an additional rate of 0.2 μg/kg/min, and continuous infusion of vasopressin at 4–8 unit/h was also initiated. Subsequently, general anesthesia was induced using sevoflurane 1–2 vol% and rocuronium 50 mg. In the arterial blood gas analysis conducted after entering the operating room, pH 7.45, PaCO₂ 20.8 mmHg, PaO₂ 335 mmHg, Na⁺ 161 mM/L, K⁺ 3.8 mM/L, glucose 205 mg/dl, lactate 13 mM/L, hemoglobin 6.8 g/dl, and HCO₃⁻ 14.4 mM/L were measured (Table 1). Four units of erythrocytes were prescribed to correct the reduced hemoglobin, and 20 mg of furosemide was injected due to a continuous lack of urination. Furthermore, for rapid transfusion, the rapid infusion system (RIS) was connected to the central venous catheter of the right internal jugular vein, and the transfusion of four erythrocyte units was initiated.

The operating surgeon planned and performed a gastrotomy, after making a skin incision of approximately 10 cm and within 20 min of beginning the surgery. The food and 5,000 ml of body fluid present in the stomach were drained via suction. Thirty minutes after the operation, blood pressure was maintained low at 60/40 mmHg and almost all food was drained; as a large amount of continuous gastric bleeding occurred after decompression, a total of 6,000 ml was drained into the suction bottle. In the follow-up arterial blood gas analysis, results showed: pH 6.75, PaCO₂ 46 mmHg, and lactate 20 mM/L. Acidosis had worsened, the blood sugar level had significantly increased to 469 mg/dl, electrolyte imbalance had exacerbated with K⁺ 7.5 mM/L, and hemoglobin had fallen to 4.3 g/dl, suggesting a large amount of bleeding. To correct this, 60 mEq of sodium bicarbonate, and two units of insulin were administered. Further, 1.2 g of calcium chloride was administered to treat hyperkalemia. Moreover, additional 20 μg of epinephrine was injected to correct persistent hypotension.

Bleeding in the stomach continued, and despite initiating erythrocyte transfusion via RIS, hemoglobin level decreased. As a result, a large amount of internal bleeding was suspected, and additional eight units of erythrocytes, and eight units of fresh frozen plasma were prescribed. The operating surgeon tried to determine the cause and area of bleeding, but as the surgical incision was only 10 cm, abdominal organs other than the stomach could not be identified with the naked eye. To reduce bleeding, the operating surgeon considered performing a total gastrectomy or extending the abdominal incision to confirm the bleeding areas from, other organs but concluded that the bleeding was due to ischemia-reperfusion injury after surgical decompression, and DIC. Since the patient’s vital signs steadily worsened, it was decided that the abdomen be closed urgently, and the patient moved to the intensive care unit (ICU).

One hour after the start of the operation, suturing after gastrectomy was completed. When the closing of the abdomen started, non-invasive blood pressure was not measured, and flat waveforms were observed in the catheter of the right radial artery. There was no pulse detected via palpation, and after confirming pulseless electrical activity through the observation of electrocardiogram, an additional 1 mg of epinephrine was injected, and cardiopulmonary resuscitation (CPR) was initiated.

A total of 3 mg of epinephrine was injected, with cardiac rhythm monitoring at 2 min intervals; afterward, 2 units of vasopressin were injected at 2 min intervals, totaling an injection of 4 units of vasopressin. After completing the abdominal closure while performing CPR, the patient was transferred to the intensive care unit, and no urine output was seen until the patient left the operation room.

The operation was conducted for 1 h 20 min, and the anesthesia time was 1 h 50 min. During the operation, a total of 800 ml, four erythrocyte units, were transfused, and 6,200 ml of crystalloid fluid were administered. A total of 6,000 ml was drained into the suction bottle—5,000 ml of food and body fluid through the suction unit via gastrectomy, and 1000 ml due to bleeding that had occurred after decompression from the start of the operation until finish; however, as mentioned earlier, the incision area was small, and bleeding occurring from other abdominal organs could not be confirmed, making it difficult to estimate the exact estimated blood loss. Additionally, prescribed erythrocytes and fresh frozen plasma were transfused through the RIS as the patient left the operation room.

After entering the ICU, CPR was stopped due to a return of spontaneous circulation, but the existing usage of epinephrine, norepinephrine, and vasopressin was preserved, and transfusion continued. However, blood pressure was still not measured, an additional 1,000 ml of bleeding was confirmed through the nasogastric tube, and bleeding from the closed abdominal area continued. Subsequently, as 3.5 g/dl hemoglobin was measured in the follow-up test, transfusion continued, and a 113.1-second prothrombin time, 180-s activated partial thromboplastin time, and platelet count of 22,000/μl were found, confirming that the patient was in a state of DIC. Despite continuous drug administra-
Abdominal compartment syndrome and complications due to bulimia nervosa

DISCUSSION

Dozens of cases have been reported that required surgical treatment due to bulimia nervosa, mentioned the problems caused by gastrointestinal necrosis or perforation, or addressed the problems associated with the occurrence of ACS. However, such a case is not commonly reported, in which ACS occurred due to rapid abdominal distension without perforation or necrosis, due to binge eating, and surgical decompression was performed without effective prior decompression, causing hypovolemic shock due to massive hemorrhage caused by ischemia-reperfusion injury and DIC, ultimately leading to death.

Kim et al. [1] and Youm et al. [2] reported similar cases. In the case report by Kim et al. [1], while gastrointestinal distension, abdominal pain, and leg color changes due to overeating were shown before surgical decompression, vital signs and blood test results were close to normal, and no signs of other organ failures were seen, thus showing no clear symptoms of ACS. In the case report by Youm et al. [2], changes in the vital signs and mental state also occurred two hours after admission. However, in this case, acute renal failure and acidosis were present since the hospital visit, and severe ACS symptoms were seen even before decompression, including mental state changes and respiratory failure.

To understand ACS, it is important to first differentiate between ACS and intra-abdominal hypertension (IAH). The common feature of the two is that IAP increases above the normal level. IAP is defined as the pressure in the abdomen, and 5–7 mmHg is considered normal. While IAH is defined as a state in which IAP is above 12 mmHg, ACS not only means a state of increased IAP but also a state in which IAP is above 20 mmHg, accompanied by a new or ongoing organ dysfunction. When the increase in abdominal pressure (AP) exceeds the abdominal wall compliance, the supply of blood and oxygen to the organs becomes restricted, and the increased pressure causes fluids to penetrate into the various organs of the abdomen, further increasing the pressure, thereby causing organ failure [3].

IAH, which can be considered as the stage before ACS, can also cause various severe complications. The World Society of the Abdominal Compartment Syndrome (WSACS) recommends measuring IAP in patients who possess risk factors for IAH or ACS, since IAH is correlated with patients’ morbidity and mortality and, therefore, treating and preventing it are important [3]. The reported risk factors include abdominal surgery, major trauma, prone position, and intra-abdominal infection. Among these circumstances which were included as risk factors, gastrointestinal swelling, acidosis, and sepsis, were present in our case, suggesting a high possibility of ACS [3].

As mentioned above, IAP is important because through it, the patient’s present state and prognosis can be predicted, and it is correlated with mortality. According to Kirkpatrick et al. [3], IAP grades can be divided into Grade I = IAP 12 to 15 mmHg, Grade II = IAP 16 to 20 mmHg, Grade III = IAP 21 to 25 mmHg, and Grade IV = IAP > 25 mmHg. According to the study performed by Prasad et al. [4], a higher IAP grade significantly correlated with a higher mortality. In a Grade IV case, 71–85% mortality has been reported [5].

In the IAP measuring method recommended by WSACS, saline solution is injected into the empty bladder and connects the inserted Foley catheter to the pressure transducer and monitor [3]. In their study, Flores-Alvarez et al. [6] also indirectly measured IAP to indirectly obtain an early ACS diagnosis and noted that anuria and grade III–IV IAH were the most important factors impacting mortality. In the present case, however, the IAP could not be measured due to a lack of time for IAP measurement and a lack of adequate equipment. According to Torquato et al. [7], increased IAP increases the airway pressure. With reference to this finding regarding the correlation between positive end expiratory pressure (PEEP) and IAP, given that the airway pressure increased to 28 cmH₂O immediately after anesthesia and dropped to 10 cmH₂O after surgical decompression under the same ventilator setting (volume control ventilation mode, 400 ml tidal volume, 12 times/min respiration rate, 6 cmH₂O PEEP), we conjectured that the IAP in this case was in a significantly elevated state. Furthermore, Bailey and Shapiro [8] reported that from an IAP of 15–20 mmHg, urine volume starts to decrease, and above 30 mmHg IAP, anuria occurs; taking into consideration that the patient in the present case was in a state of anuria, a high possibility of the patient having been in a grade IV state with an IAP above 30 mmHg can be inferred. The aspect that could be considered responsible for causing such a decrease in kidney function is that in an increased IAP state, the inferior vena cava and portal vein are compressed.
due to pressure; venous return decreases and the intra-tho-
racic pressure increases, thereby reducing the flow of the
superior and inferior vena cava and, thus, reducing the
cardiac output. According to the report by Bailey and Sha-
piro [8], even after the restoration of cardiac output in a
state of renal failure due to ACS, the impairment of kidney
function did not recover. Through this observation, they
reported that rather than the cause of renal failure being
influenced by decreased cardiac output, if the parenchyma
of the kidney was compressed, renal vascular resistance
could increase from 500% to 1,500%, and this phenomenon
could increase the secretion of renin, antidiuretic hor-
mones, and aldosterone, further increasing the renal vas-
cular resistance and exacerbating renal dysfunction [8].
Therefore, it is necessary to anticipate that high grade IAH
and high mortality rates may be present in cases of exacer-
bation and to prepare for active treatment.

Saggi et al. [9] reported that intestinal perfusion damage
occurs at mucosal and submucosal levels in IAPs above 20
mmHg, which reduces tissue oxygen tension and increases
acidosis and free radical production. This ischemic dam-
age, increased free radicals, and endotoxins lead to mul-
tiple organ failure [10]. Recent studies have revealed a cor-
relation between increased IAPs above 10 mmHg with sep-
sis, mortality, and multiple organ failure [9].

An important factor contributing to the death of patients
with long-term failure due to ACS is ischemia-reperfusion
injury. This means that the tissue damage occurs when tis-
sues are reperfused after a period of ischemia, during
which blood is not supplied or oxygenated for any reason.
During the ischemic period, many changes occur in the
cells; if ischemia persists, the cells die, or if reperfusion oc-
curs in already changed cells, the cells generate free rad-
cicals. The resulting oxidative stress causes inflammatory
and oxidative reactions that damage tissues. When this
damage occurs in microvessels, the permeability of the
cells increases, and fluids, including blood, easily escape
the tissues [11,12]. Such tissue damage is likely to worsen
organ failure and simultaneously cause massive hemor-
rhage and induce DIC.

Reperfusion syndrome can frequently occur during de-
compression surgery and be a cause of death. Rapid dia-
lation of the abdominal and pelvic vein after decompression,
and tissue damage and increased blood vessel permeabili-
ity due to ischemia-reperfusion injury cause massive hem-
orrhage, leading to a state of low blood pressure, cardiac
atrophy and ventricular arrhythmias, and resulting in high
mortality [5]. Therefore, in order to prepare for the phe-
nomenon of ischemia- reperfusion, systemic vascular re-
sistance, and a rapid decrease of IAP after decompression,
the administration of sufficient fluid is needed before de-
compression [9].

Since the problems caused by increased IAP are impor-
tant, the treatment and prevention of IAH and isch-
emia-perfusion injury are also important. According to the
ACS guideline published by the WSACS in 2013, it was rec-
commended that anxiety and pain be reduced in patients
through sedation and anesthesia to prevent and treat IAH;
it also reported that temporary use of neuromuscular
blockers can reduce IAP by reducing the abdominal mus-
cle tension, and increasing abdominal compliance [3]. Ma-
calino et al. [13] reported that neuromuscular blockers im-
proved IAP, blood pressure, airway pressure, and urine
output volume before surgery. Moreover, in case intraperi-
toneal fluid is clearly present in the abdominal cavity, per-
cutaneous drainage was deemed to be helpful, and it was
recommended to attempt a decompression via a nasoga-
stric or rectal catheter if the GIT was enlarged. Diuretics,
albumin, and dialysis, on the other hand, were not helpful
and therefore, not recommended. While the aforemen-
tioned methods can improve the patients’ state, ultimately
a surgical method should be employed, and decompres-
sion laparotomy should be performed in order to reduce
IAP and improve the organ function. Muresan et al. [14] in-
vestigated how decompression laparotomy could reduce
mortality in patients with ACS. Treatment in this study
used a four-step therapy protocol. The first step was to in-
sert a nasogastric catheter into the patient, give an ade-
quate sedation effect, and not to supply excessive fluids. Step
two was to manage the edema of the tissue using di-
uretics and hypertonic or colloid solutions in a Trelenden-
burg position. Step three was to stop nutrition and insert a
rectosigmoid aspiration catheter or puncture for discharge
using ultrasound or CT to decompress the colon before
mechanical ventilation after endotracheal intubation. In
step four, despite the medical treatment, surgical decom-
pression was attempted if the IAP remained above 20
mmHg after 24 h after IAP elevation. After decompression,
the Vivano® Med Abdominal Kit (Paul Hartmann AG, Ger-
many) was used to continuously lower IAP and reduce the
tissue edema. The Vivano® Med Abdominal Kit used foams
and layers to keep the abdominal wound open without su-
turing and to control the intraperitoneal pressure using a
suction catheter connected to the foam. Several days later,
sutting was performed after the IAP, and the wound was stabilized. The result of the study showed that surgical decompression reduced the mortality rate by 8.7%. It was reported that the shorter time between the onset of ACS and decompression laparotomy, the better the prognosis; prognosis was poor in case the time exceeded 24 h.

As previously mentioned, ischemia-reperfusion injury and shock due to DIC are more likely to occur after decompression laparotomy. In the report by Morris et al. [15], 1 g of 0.45% normal saline was mixed with 50 g mannitol and 50 mEq sodium bicarbonate to minimize damage due to the reperfusion washout effect of the anaerobic byproducts, and providing sufficient volume prior to surgery, which helped in the reduction of ischemic reperfusion injury.

The unfortunate aspect in this case was that the wound was closed immediately after decompression laparotomy, and IAP was not controlled. Muresan et al. [14] reported that it is important to keep in mind that suturing the IAP with open wounds can improve patient condition and improve the survival rate.

In conclusion, we report a case in which bulimia nervosa increased IAP and progressed to ACS; after surgical decompression, a massive hemorrhage occurred due to ischemia-reperfusion injury and DIC, leading to a state of hypovolemic shock and ultimately death. If it is possible to measure IAP, the patient’s state and prognosis can be predicted, but in emergency rooms, measuring IAP is realistically difficult; thus, evaluating the clinical symptoms that may occur in the patient, depending on the increase of IAP and progression of ACS, may be useful for the early detection and prediction of IAP and, thus, the occurrence of ACS. If the patient’s condition is critical, AP should be relieved via surgical decompression as soon as possible, and nasogastric tube insertion and the use of neuromuscular blockers prior to surgery may be helpful in alleviating the patient’s state. Prophylactic treatment with mannitol and sodium bicarbonate prior to decompression to minimize ischemia-reperfusion injury and suturing after adjusting IAP with open wound are expected to reduce mortality.

**SUPPLEMENTARY MATERIALS**

Supplementary data containing Korean version of this article is available at https://doi.org/10.17085/apm.2020.15.2.251.

**CONFLICTS OF INTEREST**

No potential conflict of interest relevant to this article was reported.

**AUTHOR CONTRIBUTIONS**

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**REFERENCES**


