Non-occlusive mesenteric ischemia during acute stroke management: three case reports

Kazuki Fujii*, Tadahiko Seki, Yasuyuki Nakata, Kazuaki Atagi and Takeshi Matsuyama

Abstract
Background: There are many reports of non-occlusive mesenteric ischemia in patients on maintenance hemodialysis and following cardiac surgery. However, there are few reports of non-occlusive mesenteric ischemia in patients with acute stroke.

Case presentation: We report three cases of non-occlusive mesenteric ischemia with onset during treatment for acute stroke. All of the patients were undergoing strict blood-pressure control, and two patients developed NOMI soon after tracheostomy when enteral nutrition had been resumed.

Conclusion: Many stroke patients are older adults with risk factors such as arteriosclerosis. Thus, during acute stroke management, there is a possibility that patients may develop non-occlusive mesenteric ischemia due to decreased intestinal blood flow secondary to strict blood-pressure control. This case report implicates early enteral nutrition as a potential etiopathogenic factor of non-occlusive mesenteric ischemia in patients with acute stroke.

Keywords: Non-occlusive mesenteric ischemia, Enteral nutrition, Stroke

Background
Non-occlusive mesenteric ischemia (NOMI) is an acute mesenteric circulatory disorder that occurs without organic vascular occlusion and is associated with extremely high mortality [1]. Potential risk factors for NOMI include age, renal disease, diabetes mellitus, reduced cardiac output, vasopressor use, intra-aortic balloon pump, and increased inflammatory markers [2–4]. However, there are few reports of NOMI in acute stroke. This report aims to present the key features of NOMI in patients with acute stroke.

Case presentation
Among the 296 stroke patients (110, 103, and 83 patients with cerebral infarction, subarachnoid hemorrhage, and intracerebral hemorrhage, respectively) treated in the intensive care unit (ICU) of our hospital from May 2018 to January 2021, based on contrast-enhanced computed tomography (CECT) scans of the abdomen, we identified three patients with acute stroke and NOMI.

Case 1
A 60-year-old woman underwent coil embolization for subarachnoid hemorrhage due to a basilar artery aneurysm rupture, but developed abdominal distension, vomiting, and metabolic acidosis on the sixth postoperative day. Abdominal CECT showed widespread contrast defects in the small intestine along with portal and intestinal emphysema, which indicated NOMI. Emergency laparotomy was performed on the same day and revealed small intestinal and ascending colon necrosis with skip lesions (Fig. 1). Indocyanine green (ICG) fluorography to evaluate intestinal blood flow showed poor fluorescence emission in the ischemic area of the bowel. The necrotic bowel was resected, but bowel Anastomosis or stoma creation was not undertaken. In the resected specimens,
histopathology showed massive ischemic changes that reflected discontinuous necrosis of the small bowel and ascending colon without arterial or venous thrombus, which supported a diagnosis of NOMI. The patient was discharged on day 119 of hospitalization.

Case 2
A 78-year-old woman underwent a craniotomy for subcortical hemorrhage in the right parietal lobe. A tracheostomy was performed on postoperative day 6, and enteral nutrition was started on the same day. To evaluate abdominal distension, a CECT was performed on postoperative day 8 and showed poor contrast uptake in the small intestine and portal emphysema. Thus, NOMI was suspected and, on the same day, emergency laparotomy revealed small bowel and ascending colon necrosis with skip lesions. ICG fluorography to evaluate intestinal blood flow revealed the ischemic bowel segments as regions of poor fluorescence emission. The necrotic bowel was resected. Bowel anastomosis was not performed, and a stoma was created. Histopathological findings supported a diagnosis of NOMI, and the patient was discharged on day 35 of hospitalization.

Discussion
NOMI causes segmental, discontinuous intestinal vascular obstruction without organic occlusion of the mesenteric vascular trunk [1]. NOMI generally affects patients older than 50 years suffering from myocardial

| Case 1 | Case 2 | Case 3 |
|---|---|---|
| Age (years) | 60 | 78 | 82 |
| Sex | Female | Female | Female |
| Stroke | SAH | Subcortical hemorrhage | SAH |
| Stroke treatment | Coil embolization Antihypertensive therapy | Craniotomy for removal of hematoma Antihypertensive therapy | Coil embolization Antihypertensive therapy |
| Risk factors for NOMI | Hypertension | OMI | Hypertension |
| Treatment for NOMI | Small bowel resection, Right hemicolecotomy, stoma creation | Small bowel resection, stoma creation | Small bowel resection, subtotal colectomy, stoma creation |
| Period from stroke onset to NOMI (days) | 7 | 8 | 9 |
| Period from tracheostomy onset to NOMI (days) | N/A | 2 | 2 |
| Outcome | Survive | Survive | Death |
infarction, congestive heart failure, aortic insufficiency, and renal or postoperatively following cardiac surgery [5]. Only one report has discussed NOMI occurring during treatment for acute stroke [6].

The American Society for Parenteral and Enteral Nutrition and Society of Critical Care Medicine 2009 recommendations for early initiation of enteral nutrition [7] have promoted early enteral nutrition for critically ill patients worldwide. Accordingly, stroke patients have undeniably benefited from early enteral nutrition in many ways.

However, NOMI is a disadvantage of early enteral nutrition. Although early enteral nutrition is cautiously administered in patients with known risk factors, enteral nutrition is aggressively used in stroke patients. Given the increasing frequency of NOMI in stroke patients (including suspected cases and cases not treated surgically), we evaluated whether, due to several factors, stroke patients might be at higher risk for NOMI.

One of the pathogenetic mechanisms of NOMI is a mismatch in the oxygen demand-and-supply ratio in the intestine, particularly in the vulnerable superficial mucosa, that is induced by early enteral nutrition [8, 9]. Thus, early enteral nutrition, which is often started earlier in stroke patients because they do not have organic bowel problems, should be recognized as a risk factor for NOMI. Furthermore, for various reasons, stroke patients undergo tracheostomy, followed by early enteral nutrition.

We consider it is preferable to start enteral nutrition after tracheostomy with a small amount or the day after tracheostomy. Of course, pathological condition leading to tracheostomy has greater effect on intestinal peristalsis. Although tracheostomy does not directly affect intestinal movement, we think it does have an indirect effect on decreased intestinal movement. Under such circumstances, early enteral nutrition may be a risk factor for NOMI.

In addition, nicardipine, which is used for strict blood-pressure control, has an inhibitory effect on the gastrointestinal smooth muscle and may cause paralytic ileus (which is a well-known side effect that tends to be ignored). Moreover, administering early enteral nutrition in ileus may lead to bacterial overgrowth and progressive edema, thereby impairing intestinal mucosal blood supply and causing ischemic injury [10].

Many stroke patients have high vascular risks, and optimal intestinal blood flow may not be maintained during strict blood-pressure control. All of the above-mentioned cases occurred under strict blood-pressure control, and two patients developed NOMI early after tracheostomy following resumption of enteral nutrition.

For the above-mentioned reasons, patients with acute stroke should be considered to be at a high risk for NOMI.

**Conclusion**

Early enteral nutrition, in addition to its manifold benefits, poses a risk of NOMI.

The stroke patients under strict blood-pressure control may be at increased risk for NOMI, and we should especially be aware when initiating enteral nutrition.

**Abbreviations**

NOMI: Non-occlusive mesenteric ischemia; SAH: Subarachnoid hemorrhage; OMI: Old myocardial infarction.

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**Authors’ contributions**

KF wrote this manuscript. TS, YN, KA and TM helped to write the manuscript. KF and YN performed the operation. All authors read and approved the final manuscript.

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**Availability of data and materials**

All data generated or analyzed during this study are included in this published article.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Informed consent was obtained from the patient and patient’s family for publication of this case report.

**Competing interests**

The authors declare that they have no competing interests.

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**References**

1. Suzuki S, Kondo H, Furukawa A, et al. The treatment and diagnosis of non-occlusive mesenteric ischemia (NOMI). J Abdom Emerg Med. 2015;35(3):177–85 (Japanese).
2. Lim JY, Kim JB, Jung SH, et al. Risk factor analysis for nonocclusive mesenteric ischemia following cardiac surgery. Medicine (Baltimore). 2017;97(37):1–13.
3. Klotz S, Vestring T, Rötker J, et al. Diagnosis and treatment of nonocclusive mesenteric ischemia after open heart surgery. Ann Thorac Surg. 2001;72(5):1583–6.
4. Stockmann H, Roblick UJ, Kluge N, et al. Diagnosis and therapy of nonocclusive mesenteric ischemia. Zentralbl Chir. 2000;125:145–51.
5. Trompeter M, Brazda T, Remy CT, et al. Non-occlusive mesenteric ischemia: etiology, diagnosis, and interventional therapy. Eur Radiol. 2002;12:1179–87.
6. Okumura E, Jimbo H, Sunagawa S, et al. Nonocclusive mesenteric ischemia is a potentially lethal complication just after stroke onset: report of six case studies. NMC Case Rep J. 2021;8:631–5.
7. Martindale RG, McClave SA, Vanek VW, et al. Guidelines for the provision and assessment of nutrition support therapy in the adult critically ill patient: Society of Critical Care Medicine and American Society for Parenteral and Enteral Nutrition: executive summary. Crit Care Med. 2009;37:1757–61.

8. Won JG, Lee YJ, Kyoungh KH, et al. Enteral nutrition associated non-occlusive bowel ischemia. J Korean Surg Soc. 2012;83:171–4.

9. Al-Diery H, Phillips A, Evennett N, et al. The pathogenesis of nonocclusive mesenteric ischemia: implications for research and clinical practice. J Intensive Care Med. 2019;34:771–81.

10. Marvin RG, McKinley BA, McQuiggan M, et al. Nonocclusive bowel necrosis occurring in critically ill trauma patients receiving enteral nutrition manifests no reliable clinical signs for early detection. Am J Surg. 2000;179:7–12.

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