Abstract

Corrosive substance ingestion is a serious public health hazard. Mostly, children are victim due to accidental ingestion occurring commonly in less than five years. Whether acid or alkali, the ingestion may have catastrophic effects and the outcomes can vary from minimal injury to perforation and death. Esophagogastroduodenoscopy should be done 12-24 hours of ingestion in order to assess the extent of mucosal injury. Esophageal stricture remains one of the major sequel usually seen after three weeks of ingestion. Endoluminal dilatation is current recommendation for initial treatment of stricture and surgery should follow for strictures refractory to dilatation. We here present a case of a five year old child with accidental corrosive ingestion at her school which led to esophageal stricture. She underwent multiple dilatation of stricture followed by esophageal corrective surgery which again led to post-surgical stricture requiring further dilatations.

Key words: Corrosive substance, esophagogastroduodenoscopy, Esophageal Stricture, dilatation

Introduction

Corrosives are a group of chemicals that have the capacity to cause tissue injury on contact by a chemical reaction most commonly affecting gastrointestinal tract, respiratory system and eyes. Accidental ingestion of corrosive substances is more commonly seen in early childhood with children younger than five years being more prone to such ingestion. Ingested corrosives are either acids or alkali. The subsequent injury varies from minimal to severe, with perforation and death as potential complication. Esophageal burns account for most serious complication. In children, 18% to 46% of all corrosive ingestion result in esophageal injury. Esophagogastroduodenoscopy is the most efficient method for assessing UGI tract mucosa after ingestion of corrosive substance and should be performed 12-24 hours of ingestion. Esophageal injury after corrosive ingestion is endoscopically graded with a score of 0 for no injury to IIIb for significant circumferential injury with ulcers and necrosis. Esophageal stricture is main complication which occurs after 3 weeks. Current recommendation for the initial management of esophageal stricture is endoluminal dilatation performed starting...
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4-6 weeks after initial ingestion. Strictures refractory to dilatation should undergo surgery\(^1\).

**The Case**

A five years old girl from Baglung district had accidental ingestion of corrosive, namely Methyl orange at her school 16 months back. Soon after ingestion she was noted with swelling of lips and tongue followed by drooling and acute onset of epigastric pain. Immediately she was referred to tertiary hospital. Upon examination, there was note of erosion of lower lip, dorsum of tongue and floor of mouth with mild tenderness at epigastric region, with normal vitals and other systemic examination. Supportive treatment was given and was discharged after 12 days with good oral intake. However, vomiting started soon after discharge which became persistent hence was referred to tertiary hospital in Kathmandu. A barium follow through was done with suspicion of esophageal/ gastric stricture which was normal. UGI endoscopy was also performed which showed esophageal stricture and was referred to India for further management.

There in India, she was admitted under pediatric gastrointestinal unit where a repeat endoscopy was performed showing two strictures at 11cm and another at 15cm, latter showed ulceration. Subsequently guide wire was placed and NJ tube inserted for feeding. She underwent multiple esophageal dilatations of stricture after that. As there was no relief, gastrostomy was performed by pediatric surgeon. Subsequently, barium swallow was done still showing narrowing of esophagus and dilatation could not be done as planned due to presence of bleeding non healing ulcer at stricture site. Patient then underwent esophageal strictuloplasty with end to end anastomosis.Gastrografin study showed normal flow following surgery. However, after three weeks of discharge vomiting started. UGI endoscopy revealed anastomotic site narrowing, again requiring esophageal dilatation. She underwent multiple dilatations in 15 days interval. Patient was then referred back to Nepal at our center for subsequent esophageal dilatation since the facility was already available here.

Endoscopic view 9 months after the last dilatation

Slight narrowing of esophagus seen at 15 cm from incisor

No difficulty in passing scope from esophagus to stomach

Esophageal stricture dilatation

Esophageal stricture at 15 cm from incisor

Guidewire placed across the stricture

Fluoroscopic appearance of guidewire

Savary Gillard dilator passed across guidewire

Esophageal stricture post dilation
In our hospital, multiple serial dilatations were done using Savary Gillard dilator under fluoroscopic guidance, initially at 15 days interval then one month and three months intervals. Gastrostomy tube was removed subsequently and was fed orally which was tolerated. Our patient is the first to undergo esophageal dilatation for stricture in children in our country.

At present, she has good oral intake and is asymptomatic with weight gain of three kilograms.

Discussion

Children amount to 80% of ingestion injury globally. Most ingestions are due to parent’s lack of knowledge of the hazards of corrosive substances kept in the house, and the availability of chemicals in and around houses, combined with the natural curiosity of children. Acids and alkali are two primary types of agents responsible for corrosive exposures. Each produce different types of tissue damage. In general alkaline substances tend to be more palatable than acidic products resulting in ingestion of larger quantities and increased risk of serious injury. Acids cause coagulation necrosis with eschar formation that may limit substance penetration and injury depth. Conversely alkalis combine with tissue proteins and cause liquefactive necrosis, saponification of fats, dehydration of tissues and thrombosis of blood vessels resulting in deeper tissue injury.

Clinical presentations are very diverse and initial symptoms do not show deep correlation with ultimate degree of injury. Clinical manifestation may vary from no symptoms to nausea, vomiting, drooling, abdominal pain, dysphagia, odynophagia, chest pain or stridor. There may be damage in the esophagus or stomach even if oropharyngeal examination is normal or patient is asymptomatic. Fever, palpitations, shock indicate extensive damage.

Initial management involves stabilizing the patient by keeping nil per oral, vigorously replacing volume and securing airway. Imaging should be done to rule out perforation of esophagus and stomach. Esophagogastrroduodenoscopy is crucial and usually recommended in the first 12-24 hours after corrosive ingestion once perforation has been ruled out, however, some reports indicate it can be safely performed up to 96 hours after ingestion. The endoscopic grading of injuries is not only to evaluate esophageal and gastric injuries but also predict prognosis and establish a management plan. Grade 0 indicates normal mucosa; Grade I indicates slight swelling and erythema; Grade IIa indicates presence of superficial ulcers, bleeding and exudates; Grade IIb indicates local or encircling deep ulceration; Grade IIIa indicates focal necrosis; and Grade IIIb indicates extensive, circumferential necrosis.

Gastric lavage and induced emesis are contraindication. Nasogastric or Nasojejunoscopy tube should be placed under direct endoscopic vision for feeding purpose. Emergent surgery is recommended if there is evidence of perforation. Patients with grade 0 injuries can be discharged immediately. Those with grade I or IIa do not require specific treatment. Patient with grade II must be carefully observed for perforation symptoms for at least 1 to 2 weeks in intensive care unit and adequate nutritional support is required. Gastrostomy may be required for Grade IIb lesions for nutritional support. Other supportive measures include broad spectrum IV antibiotics, PPI, antacid. Although the use of systemic steroid in treatment of corrosive esophagitis remains controversial, some reports have documented the advantage of high dose systemic steroid administration for prevention of stricture formation.

Long term sequel include esophageal or gastric stricture usually after 3 weeks. The rate of stricture formation is reported to be between 2% and 63%. Pyloric stenosis and gastric outlet obstruction can also occur following corrosive ingestion. Endoluminal dilatation is the management of choice in children, which can be done endoscopically by either a bougie or balloon catheter. After initial injury, it is advisable to wait for 3 to 6 weeks prior attempting dilatation. For dilatation, progressively larger bougies are inserted over endoscopically placed guidewires. Esophageal replacement surgery becomes necessary for those refractory ones. However, balloon dilatation or endoscopic bouginage is still necessary for recurrence of stricture post surgery, like in our case presented.

Conclusion

Corrosive ingestion possesses a potential public health risk and widespread lack of knowledge regarding the harmful effects and preventive measures remains important risk factor for these injuries. Education is important to spread awareness of dangers associated, also its use should be regulated in order to reduce or prevent such injuries.
References


