CASE REPORT

Accidental aspiration of a solid tablet of sodium hypochlorite

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SUMMARY

Sodium hypochlorite is a corrosive, highly alkaline (pKₐ = 7.52) household product. Ingestion of sodium hypochlorite liquid is common, showing toxicity on the oesophagus and stomach. Nevertheless, cases of sodium hypochlorite ingestions in pellet are rare and the management of them is unknown. We report the case of a 65-year-old man who accidentally swallowed a bleach tablet of 3.5 g. Six hours later, the patient developed an aphony associated with dyspnea stage IV, motivating a nasofibroscopy showing glottis and supraglottic necrosis and oedema for which the patient received intravenous steroids, was intubated and then underwent a tracheotomy. After 2 weeks under tracheotomy, local evolution was favourable allowing a removal of the cannula and a return back home.

BACKGROUND

Caustic agent ingestion, accidental or voluntary, is a common problem with more than 5000 cases reported each year in the USA. Sodium hypochlorite is one of the agents most commonly swallowed in liquid form. Emergency care following this kind of ingestion is documented, as well as its effects on the oesophagus and stomach. However, ingestion of a soluble tablet of sodium hypochlorite is rare. The treatment, as well as possible damage on the pharynx and larynx, remains unknown. Through this report, we are going to describe the ingestion of a tablet of sodium hypochlorite, the clinical presentation, the evolution and our support.

CASE PRESENTATION

A 65-year-old man, only known, medically, for high blood pressure, accidentally swallowed a tablet of soluble bleach sodium hypochlorite. The patient had a headache and decided to take an acetaminophen tablet that was near the kitchen sink. However, confused by his pain, he took by mistake a tablet of soluble sodium hydroxide (3.5 g of sodium hypochlorite per tablet) followed by a glass of water. He described an immediate burning sensation in his mouth and throat, followed by a severe cough, allowing him to expectorate part of the tablet. He arrived at the emergency room 1 hour after ingestion.

The patient complained only of an odynophagia associated with light dysphonia. He suffered no abdominal pains and drank before coming to the emergency room. Vital signs showed an afibrile and with a blood pressure of 168/82 mm Hg. Physical examination showed no erythema on the lips, in the oral cavity or deep throat. Abdominal examination was normal, without pain during the palpation. Blood tests showed an aligned blood formula, a preserved kidney function and excluded any inflammatory syndrome.

The ear, nose and throat (ENT) specialist made a nasofibroscopy showing a soft laryngeal erythema, without oedema. After sufficient pain treatment with intravenous opioid, the patient described an improvement of the odynophagia and a partial improvement of his voice. Then, the patient was kept under medical supervision in the emergency room for 6 hours post-ingestion, in anticipation of a new ENT control before a possible return home.

After 6 hours of monitoring, the patient presented a stage IV dyspnoea associated with a severe odynophagia and aphony. A new ENT examination excluded an oral lesion, showing necrotising fibrin of supraglottis and massive bilateral arytenoid oedema significantly decreasing glottis flow by 40%. The visible real vocal folds were described as fine.

Initial management includes 250 mg of solumedrol followed by an awake nasotracheal intubation.

INVESTIGATIONS

After intubation, a pharyngoscopy was realised and confirmed superficial necrosis of dorsal epiglottis extending to vocal folds (figure 1) with transcutaneous oedema of both arytenoids (figure 2). Flexible gastroscopy showed a superficial 1 cm² ulcer of proximal oesophagus without any other lesion up to duodenum. The day after, considering the expected slow healing process and to avoid laryngeal synechias, a surgical tracheotomy was performed.

DIFFERENTIAL DIAGNOSIS

The cause of the oedema was the direct irritation of an alkaline product on the laryngeal mucosa. An allergic origin was unlikely.

TREATMENT

The patient had a tracheotomy for 2 weeks, associated with a proton pump inhibitor treatment (esomeprazol 40 mg) two times per day for 2 days and then one time per day. The patient was hospitalised in intensive care for 2 weeks, where he was given an enteral alimentation via a nasogastric tube only.

OUTCOME AND FOLLOW-UP

Hospitalisation was complicated by a pulmonary infection, for which an antibiotic treatment was initiated. After 17 days under tracheotomy, the cannula was removed. The patient was given a smooth, then...
Unusual presentation of more common disease/injury

Ingestion of caustic alkaline product, such as sodium hypochlorite, causes lesions of the mucosa, by a penetrating injury called liquefac-tive necrosis, allowing deeper tissue damage than acidic substances. By aspiration of a tablet of sodium hypochlorite in this patient’s case, the lesions remained mainly high, at the laryngeal level, causing a local necrosis and oedema.

The different guidelines describe support for lesions of the oesophagus and stomach but not for the lesions of the larynx. The consequences of such ingestion are not known before an endoscopic examination, so it is recommended to offer a similar support the first time. This recommendation is the initiation of a proton pump inhibitor, an endoscopy within 24 hours and a medical supervision. It should be noted that endoscopy is recommended only for stable patients and without clear evidence of perforation of the oesophagus. It is also best to do X-rays of the thorax and abdomen regularly to exclude an oesophageal perforation. The use of emetic agent, such as coal, or neutralising agent is contraindicated as it can cause vomiting and a new exposure to the irritant product.

In the case described above, we decided as a first step to initiate a corticosteroid therapy in view of important laryngeal oedema. Various studies show the absence of benefits from a corticosteroid therapy in the case of oesophageal lesions. There is no evidence that it is beneficial for laryngeal caustic lesions, but meta-analysis supports its beneficial effect on laryngeal oedema after intubation. It has been used in many similar situations, like burns by hot drink or food. Here, a major laryngeal oedema motivated the initiation of this particular treatment, just by a single bolus, with a good clinical evolution.

Also, it was advisable for this patient to perform a nasopharyngeal and awake intubation. This practice is not indicated in the recommenda-tions for the treatment of ingestion of caustic liquids, in view of the risk of vomiting following this intubation and thus re-exposure to the caustic product. However, in this case, the main problem being the high obstruction of the larynx by the oedema, like in cases of laryngeal oedema in burns, then visualisation was needed for this gesture and early intubation is necessary for upper airway protection. Furthermore, the risk of vomiting and re-exposure to the irritant was low, the patient having ingested it in solid form, with only local irritation and more than 6 hours before.

In this case, the patient presented with a little voice modification but with an improvement after pain killer and a quite normal first nasofibroscopy. This explains why an immediate intubation was estimated overweening. However, an earlier intubation should be discussed in case of severe dysphonia and/or oedema.

Finally, the tracheotomy is the final treatment, the intubation tube being a possible cause of exacerbating irritation and synechias, needed to discharge this area. On a public health side, this case reminds the importance of appropriate storage for medications and cleaning products.

Patient’s perspective

I was drowsy, my throat was painful and I could not really talk any more. Much later, an ENT finally came to examine me again: he left quickly and returned with four or five doctors in green clothes. I remember that I needed to urinate but it was denied to me because ‘there was no time’. I don’t have any more memories from this moment and until I got out of coma weeks later. (…)

Everyone told me: I came back from a long way. I know I have fought a survival battle, so I wanted to live.

Learning points

► Ingestion of sodium hypochlorite in undiluted pellet form is an uncommon but dangerous emergency department complain.
► In the absence of specific guidelines, management should be the same as for liquid form.
► Steroids and awake nasotracheal fibroscopic intubation should be considered for life-threatening laryngeal oedema.
Correction notice  This article has been corrected since it was published online. The substance ingested by the patient described in the case report was sodium hypochlorite (Bleach, pHK=7.52) and not sodium hydroxide (Caustic soda, pHK=14.8).

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REFERENCES
1. Day RC, Bradberry SM, Sandilands EA, et al. Toxicity resulting from exposure to oven cleaners as reported to the UK National Poisons Information Service (NPIS) from 2009 to 2015. Clin Toxicol 2017;55:645–51.
2. Palao R, Monge I, Ruiz M, et al. Chemical burns: pathophysiology and treatment. Burns 2010;36:295–304.
3. De Lusong MAA, Timbol ABG, Tuazon DJ. Management of esophageal caustic injury. World J Gastrointest Pharmacol Ther 2017;8:90–8.
4. Hawkins DB, Demeter MJ, Barnett TE. Caustic ingestion: controversies in management. A review of 214 cases. Laryngoscope 1980;90:98–109.
5. Contini S, Scarpignato C. Caustic injury of the upper gastrointestinal tract: a comprehensive review. World J Gastroenterol 2013;19:3918–30.
6. Park KS. Evaluation and management of caustic injuries from ingestion of Acid or alkaline substances. Clin Endosc 2014;47:301–7.
7. Penner GE. Acid ingestion: toxidology and treatment. Ann Emerg Med 1980;9:374–9.
8. Anderson KD, Rouse TM, Randolph JG. A controlled trial of corticosteroids in children with corrosive injury of the esophagus. N Engl J Med 1990;323:637–40.
9. Fulton JA, Hoffman RS. Steroids in second degree caustic burns of the esophagus: a systematic pooled analysis of fifty years of human data: 1956-2006. Clin Toxicol 2007;45:402–8.
10. Zhang W, Zhao G, Li L, et al. Prophylactic administration of corticosteroids for preventing postoperative complications related to tracheal intubation; a systematic review and meta-analysis of 18 randomized controlled trials. Clin Drug Investig 2016;36:235–65.
11. Iyama K, Ueki T, Yamano S, et al. An adult case of laryngopharyngeal burn by drinking hot water. Acute Med Surg 2017;4:198–201.
12. Lund T, Goodwin CW, McManus WF, et al. Upper airway sequelae in burn patients requiring endotracheal intubation or tracheostomy. Ann Surg 1985;201:374–82.
13. Rosenblatt WH. The airway approach algorithm: a decision tree for organizing preoperative airway information. J Clin Anesth 2004;16:312–6.
14. Sahu G, Behera P. Fatal laryngeal oedema in burns. J Trauma & Treatment 2015;4.
15. Hogg G, Goswamy I, Khwaaja S, et al. Laryngeal trauma following an inhalation injury: a review and case report. J Voice 2017;31:388.e27–31.
16. Rafiey M, Ghajzadeh M, Sheikhi S, et al. Caustic ingestion in children: a systematic review and meta-analysis. J Caring Sci 2016;5:251–65.