A fatal case of creosote poisoning

CLIVE E. BOWMAN
B.Sc., M.R.C.P.

MARK F. MUHLEMAN
M.R.C.P.

EWAN WALTERS
B.Sc., M.B.Ch.B.

Department of Medicine, Southmead Hospital, Westbury-on-Trym, Bristol BS10 5NB

Summary

A case of fatal creosote poisoning is described. On presentation, extensive oropharyngeal ulceration was noted and gastric lavage withheld. Post-mortem examination showed an intact oesophagus and stomach.

KEY WORDS: kidney failure, acute acidosis.

Introduction

Industrial creosote, commonly used for wood preservation, is a distillate of coal and wood tars. Phenols and cresols are responsible for its toxicity. Management of creosote poisoning is based on experience from cases poisoned by cleaning agents such as Lysol, now largely withdrawn, which are much more concentrated phenolic solutions. To our knowledge there is not a full published account of fatal creosote poisoning.

Case report

A 70-year-old man was found unconscious with a cup of creosote beside him. On admission he was unconscious responding only to painful stimuli, rectal temperature 34°C, centrally cyanosed and shocked with a pulse of 60/min, blood pressure 60/40 mmHg. His respiratory effort was poor and on auscultation widespread crackles were heard. Creosote and vomitus stained his face and clothing. There was obvious extensive ulceration of the oropharynx.

Endotracheal intubation and artificial ventilation was instituted immediately. Gastric lavage was not performed because of presumed oesophageal ulceration. Forty-five minutes following presentation and initial resuscitation (including intubation and ventilation), measurements showed pH 7.15, $P_O_2$ 204 mmHg, $P_CO_2$ 24 mmHg, bicarbonate 9.3 mmol/l, base excess -16.2 mmol/l. He was treated with gradual rewarming, intravenous amoxycillin and hydrocortisone (200 mg 4-hourly). Two litres of normal saline were infused over the first 2 hr of admission which raised the central venous pressure from −2 cmH₂O to +5 cm (using mid axillary line as zero). His blood pressure remained low but with an infusion of dopamine rose to 140/80 mmHg. Over the first 8 hr of admission an average of 30 ml/hr of smoky coloured urine was excreted. Thereafter urine output fell gradually in spite of vigorous therapy with frusemide and mannitol.

Five hours after admission arterial blood showed pH 7.09, $P_O_2$ 82 mmHg, $P_CO_2$ 22 mmHg, HCO₃⁻ 6.9. Base excess -17.8 after which bicarbonate was given. The patient became anuric and after 30 hr death occurred.

At post-mortem about 1 litre of fluid, mainly creosote, was found in the stomach. The oesophageal and gastric mucosa were stained brown but were well preserved. Petechial haemorrhages were noted over the serosal surface of the lower jejunum, ileum and also over the pericardium. Histology of oesophageal and gastric mucosa showed no ulceration. Renal histology showed acute tubular necrosis and the liver showed degeneration and necrosis of hepatocytes with minimal inflammation.

Discussion

Phenols denature and precipitate proteins; they are general cellular poisons, for which, once absorbed, there is no specific antidote.

When ingested, absorption is initially rapid and then subsequently slower probably due to local vascular damage (Locket, 1957). Early poisoning may be characterized by a respiratory alkalosis due to a salicylate-like stimulation of the respiratory centre (Dreisbach, 1977), thereafter an acidosis of mixed aetiology occurs. Methaemoglobinemia may develop. Death is a result of multi-organ failure.
Prevention of absorption by gastric lavage clearly is the most effective treatment. Addition of vegetable oils to the lavage fluid by dissolving phenols should minimize absorption (Burston, 1970). Traditionally oesophageal integrity has been assessed indirectly by the condition of the oropharynx, but in the presented case this proved misleading. It seems likely that ulceration of the oesophagus and stomach did not occur in this case because of the greater dilution of phenols and cresols in creosote compared with, for example, Lysol. Supportive evidence comes from veterinary experience, chronic ingestion of creosote in sheep leads to death from multi-organ failure, whereas purer phenol preparations cause violent corrosive effects (Clarke and Clarke, 1975). Therefore in creosote poisoning, oropharyngeal ulceration is an unreliable indicator of gastrooesophageal integrity. Expert endoscopic assessment and the possible alternative of lavage via a gastrostomy should be considered.

Acknowledgments

We thank Dr N. Malcolm for permission to report this case and Dr A. Savage for performing the post-mortem examination.

References


(Accepted 8 June 1983)
A fatal case of creosote poisoning.

C. E. Bowman, M. F. Muhleman and E. Walters

Postgrad Med J 1984 60: 499-500
doi: 10.1136/pgmj.60.705.499

Updated information and services can be found at:
http://pmj.bmj.com/content/60/705/499

Email alerting service

These include:
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/