Lead preparations in Ayurvedic medicines

Sir,

Keen et al.1 presented a typical case of hepatitis after using an Ayurvedic preparation for diabetes. The authors did not give the details of the drug dosage, trade name, pharmaceutical dosage form or source. Ayurveda lays much emphasis on the purificatory aspects of its drugs which are mainly of herbo-metallo-mineral and animal origin, some of which are highly toxic when taken in their crude or natural form. After specific purificatory processes, metals are subjected to treatments such as impregnation and trituration with other herbal drugs and are eventually subjected to different types of heat treatment, using conventional methods. The heavy metals are brought into colloidal form after incineration. For example, the lead-containing preparation, sastiputa naga bhasma is subjected to 60 cycles of incineration.

Studies were carried out at Gujarat Ayurveda University2 and Banaras Hindu University,3 in India on lead as prepared by the different methods used to make Ayurvedic medicines and their use in the treatment of different diseases.

Experimental studies showed that Pb, PbO, PbSO4, and PbS preparations available in the market are toxic. X-Ray diffraction analysis proved that the Ayurvedic preparation sastiputa naga bhasma is chemically PbS; it is nevertheless reported to be non-toxic in experimental and clinical studies.4 Depending on the preparation, the lead content varies from 14.5% to 68.14%. The lead content of sastiputa naga bhasma reported by Nagaraj was 64.24% whereas in their sample of the drug Keen et al.2 reported a lead content of 19%.

The experimental study on rats and the clinical trials conducted proved that the Ayurved lead preparations are non-toxic when the dose is below 6 mg/100 g body weight.5 The prescribed dose in the treatment of diabetes is 30–120 mg/day in case of an adult.6 The results of the clinical trials carried out with sastiputa naga bhasma and other hypoglycaemics are given in the table to get a clear picture of its action on diabetes.

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Table Results of clinical trials with sastiputa naga bhasma

<table>
<thead>
<tr>
<th>Groups</th>
<th>Fall in blood sugar level (% of cases)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sastiputa naga bhasma</td>
<td>50*</td>
</tr>
<tr>
<td>Sastiputa naga bhasma</td>
<td>65</td>
</tr>
</tbody>
</table>

*No hypoglycaemia was reported

This letter was shown to the authors who reply as follows:

Sir,

Ravinarayan and Skandhan claim that lead, present as PbS, in Ayurvedic medicines is non-toxic, because of specific purification processes which include multiple incinerations. The evidence cited in support of this claim is contained in four unpublished theses submitted for either MD or PhD degrees between 1984 and 1992. However, no treatment can alter the facts that lead sulphide is soluble in acids, including gastric secretions, and that soluble lead is toxic.

Ravinarayan and Skandhan report a prescribed dose of 30–120 mg/day of an Ayurvedic medicine for the treatment of adult diabetes. The intake of lead from this medicine would be 6–24 mg/day—assuming a lead content of 20% w/w. Sustained intakes by adults, of more than 1 mg/day will lead to increased blood-lead concentration and adverse metabolic, functional and clinical effects.4 Therefore, one would expect to see adult poisoning with prolonged intakes of the above medicines. This is what we observed and reported.2

Our patient had a limited supply of a medicine obtained indirectly from India, and had taken the preparation for a relatively short period, about six weeks. Although it was not possible to establish and document the dose taken, the similarity between the unusual lead isotopic compositions of the substance and the patient’s blood, indicated this to be the source of lead poisoning.

Our conclusion, that such medicines are not safe, remains unaltered.

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Streptococcus mitis causing epidural abscess

Sir,

We report a case of spinal epidural abscess caused by Streptococcus mitis. A 57-year-old haemophiliac man presented to accident and emergency with neck pain of six days duration. It radiated to the shoulders, was worse with movement and had been getting gradually worse. Examination revealed a temperature of 38°C. Neck movements were restricted. Muscle spasm was present and there was mild tenderness over C4–C5. Peripheral white cell was 11.8 x 109/l and the erythrocyte sedimentation rate was 70 mm/h. Blood cultures and cervical X-rays were taken. A bleed was thought to be the cause of the symptoms although the fever was difficult to explain. The X-ray showed diffuse degenerative changes of the cervical vertebrae but no soft tissue changes suggestive of haemorrhage or infection. The patient was given 1500 units of factor VIII, but despite a further several doses over 24 hours there was no improvement. His temperature continued to fluctuate, sometimes up to 39°C and he had occasional rigors.

On day three of admission both bottles of blood cultures grew Streptococcus epidermidis and an alpha-haemolytic streptococcus later identified as Streptococcus mitis. These organisms were felt to be probable contaminants and repeated cultures were performed which proved negative. The following day he developed weakness and numbness of the right arm and MRI scan revealed an anterior cervical abscess over C4–C5 vertebrae. This was drained under factors VIII and IX. The pus and a swab from the abscess site grew a heavy pure growth of Streptococcus mitis. This proved a sensitive organism and he was treated with benzylpenicillin 1.2 g iv four hourly and metronidazole 400 mg orally three times a day for two weeks followed by benzylpenicillin alone for a further 10 days when he developed a maculopapular rash. The penicillin was replaced by erythromycin 500 mg orally four times daily for two weeks. His temperature settled and he made a good recovery.

The microbiology of spinal epidural abscesses is dominated by S aureus. Standard therapy should involve surgical drainage where possible as well as prolonged antibiotic therapy (4–6 weeks). We chose benzylpenicillin and metronidazole in case there was an anaerobic organism that failed to grow. Erythromycin is a suitable alternative in cases of allergy. We believe this is the first reported case of spinal epidural abscess caused by Streptococcus mitis, and illustrates the importance of sampling at operation to obtain the correct microbiological diagnosis.

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Spinal epidural abscess

- Staphylococcus aureus 52–95%
- coagulate-negative staphylococci
- aerobic Gram-negative rods
- anaerobes
- streptococci (A, B, pneumococcus)


References

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HT Delves and PG Frost

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