Hereditary sensory and autonomic neuropathy in a male child: ‘The other side of not feeling pain’

Nripen Gaur,1 Rachna Meel,2 Shahnaz Anjum,1 Pallavi Singh1

A 1-year-old boy was brought to our outpatient department with complaints of whitish opacity in both eyes noticed 3 months ago. The mother also gave a history of self-mutilating behaviour in the child. There was a history of absence of crying during vaccination. There was no history of consanguinity and a similar disease in the family. The child was following light binocularly. On anterior segment examination, bilateral corneal scarring secondary to keratitis was seen (figure 1A). Corneal sensations were absent. An ocular ultrasonography was done for posterior segment evaluation, which revealed no abnormalities. There were signs of self-mutilation in the perioral area in form of damage to both upper and lower lips that caused as a result of repeated insults in form of tooth bite (figure 1B). Apart from this, similar injuries were present in fingers of hands (figure 1C) as well as toes. On neurological examination, the child had normal motor functions, and deep tendon reflexes were normal. No thickening of peripheral nerves was noted. However, there was no response to pain and temperature stimuli. Rest of the systemic examination was not remarkable. No abnormality was found on neuroimaging. The child was referred to the department of paediatrics where a clinical diagnosis of hereditary sensory and autonomic neuropathy (HSAN) was made. Nerve conduction study and sympathetic skin responses tests showed absent sympathetic response from the skin of both hands and both feet. However, the electromyography study was normal. Gene sequencing test revealed the presence of homozygous mutation in the PRDM12 gene, which confirmed the diagnosis of HSAN type 8 (HSA-VIII). The child was referred to the department of cornea for considering visual rehabilitation in form of keratoplasty.

HSAN is a group of disorder that is characterised by inherited autonomic and sensory neurological defect. HSAN is further classified into eight types based on the genetic mutation and clinical characteristics.1 HSAN-VIII is caused by the homozygous mutations in the PRDM12 gene. This syndrome compromises of following features: insensitivity to pain and thermal stimuli, self-mutilation behaviour, altered sweat and tear formation, the absence of corneal reflexes and presence of repeated infections of the skin and bone.2 The patients having PRDM12 mutations lack pain sensations.3 Hence, these patients are subject to multiple injuries leading to recurrent skin infections and bone deformities. The lack of corneal sensations can lead to corneal neurotrophic ulcerations and subsequent scarring. Pain insensitivity can lead to oral mutilations like dental attrition, bite wounds, premature tooth loss and ulcerations. The most common sites for self-inflicted injuries involve oral mucosa and lips. The rare nature of this disease often leads to a delayed diagnosis and development of complications. The management involves use of lubricants to prevent corneal ulcers. Frequent and careful examination of limbs to look for any fresh injuries should be done.

Contributors NG, SA and PS contributed to diagnosis, workup, writing the manuscript and performing critical revision. RM is the overall responsible for the presentation, contributed to diagnosis and performed critical revision of the manuscript.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Patient consent Parental/guardian consent obtained.

Provenance and peer review Not commissioned; externally peer reviewed.
REFERENCES

