Case Report

DOI: 10.5152/tftrd.2015.62444


Congenital Indifference to Pain: A Report of Three Cases from Pakistan

Ali Raza QURESHI, Saeed Bin AYAZ, Atif Ahmed KHAN, Zaheer Ahmed GILL

Armed Forces Institute of Rehabilitation Medicine (AFIRM), Rawalpindi, Pakistan

Abstract

Congenital indifference to pain is a rare disease defined by the failure to react to painful stimuli. Patients often present in early childhood with multiple self-inflicted injuries and damaged dentition. The response to other sensory modalities remains intact. Patients have normal nerve conduction and electromyography, as well as normal nerve biopsy. The disease is diagnosed after excluding congenital insensitivity to pain, fragile X-syndrome, syringomyelia, and pain asymbolia. Early diagnosis can reduce recurrent injuries and avoid unnecessary investigations. Here we present three patients with congenital indifference to pain who were referred to our institute for electrophysiological evaluation and found to have this disease.

Keywords: Pain, congenital indifference to pain, congenital insensitivity to pain, nerve conduction studies, electromyography, Pakistan

Introduction

Pain is generally described as an unpleasant sensation or a distressing experience, but it is actually the natural defense from physical harm. Absence of pain perception makes a person prone to repeated soft tissue damage, bone fractures, and rapidly degenerating joints. Congenital indifference to pain is a rare syndrome, which characteristically features failure to exhibit aversion to or withdrawal from painful stimuli, despite not having neurological abnormalities in pain pathways (1). Children affected by this syndrome present with unusual injuries during their early years, which are generally caused by repetitive self-inflicted damage (2). The diagnosis depends on clinical features, as well as biochemical, electrophysiological, pathological, and genetic testing. Here we report three patients who were electrophysiologically evaluated at the electrodiagnostic department of our institute and who were found to have congenital indifference to pain.

Case Reports

Case 1

A 9-month-old female infant was referred to our institute for electrophysiological evaluation. She had a history of abnormal biting and self-mutilating behavior that had damaged her tongue, teeth, lips, and fingers. She was the first-born to a consanguineous marriage and had no siblings. On examination, she was sitting comfortably in her mother’s lap and had excessive dribbling of saliva from the mouth. Her face was not dysmorphic. She had bitten her tongue and lower lip. Her two upper incisors were missing (Figure 1). There were bilateral bite marks on her fingers. She did not respond to pinprick or exposure to hot and cold stimuli. Her joint examination was normal. Muscle tone and deep tendon reflexes (DTR) were also normal.

Laboratory investigations of this patient showed normal complete blood count and serum uric acid levels. Magnetic resonance imaging (MRI) of her brain with and without contrast was also normal. Her electrodiagnostic evaluation was...
performed on a XLTEK Neuromax 1004 Electromyography (EMG) Unit using surface electrodes for nerve conduction studies (NCS) and concentric needle electrodes for EMG. Motor NCS revealed normal latencies, amplitudes, and conduction velocities in median, ulnar, and radial nerves in the upper limbs as well as the common peroneal and tibial nerves in the lower limbs. Sensory NCS revealed normal latencies, amplitudes, and conduction velocities in median, ulnar, and radial nerves in the upper limbs and sural nerves in the lower limbs. The child did not respond to electrical stimulation but started crying on EMG. EMG showed no involuntary activity and normal morphology and recruitment of motor unit action potentials in key sampled muscles of the four limbs. The patient was given an electrophysiological impression of a normal study, and a clinical correlation for a diagnosis of congenital indifference to pain was suggested. Informed consent was obtained from the mother of the patient for publication of this case report and the accompanying image.

Case 2
A 30-month-old male presented with a three month history of pressure ulcers on both knees. He had a history of repeated tooth loss because of biting very hard objects. His two siblings from a non-consanguineous marriage were normal. There was no history of similar cases in the family of the father or mother. When queried, his parents explained that the child had achieved bladder control and had normal sweating. However, he did not demonstrate a normal reaction to foul smells. On examination, he had an active pressure ulcer of grade III over the anterior aspect of the right knee and a healing pressure ulcer of grade II over the left knee (Figure 2). There was a healed wound over the left elbow (Figure 3). The skin around the right big toe was thick, hard, and rough and had a crack over the right first metatarsophalangeal joint (Figure 2). His tongue was slightly bitten, and his lower incisors and canines were missing (Figure 4). He did not show withdrawal response to pinprick yet distinguished between hot and cold stimuli. His muscle tone and DTR were normal.

The patient’s hemoglobin (9.3 g/dL; normal: 13–17 g/dL) and neutrophil percentage (37%; normal: 40–80%) were reduced. His platelet count (562 × 10^9/L; normal: 150–400 × 10^9/L), lymphocyte percentage (56%; normal: 20–40%), and erythrocyte sedimentation rate (ESR) (18 mm at the first hour; normal: 0–9 mm at the first hour) were all raised. However, his total leucocyte count and serum uric acid level were normal. His NCS showed normal motor and sensory studies, and EMG was within normal limits. The patient was advised to undergo a sural nerve biopsy that turned out to be normal. Informed consent was obtained from the mother of the patient for the publication of this case report and the accompanying images.

Case 3
A 4-year-old boy presented with a two month history of pressure ulcers that had developed over the anterior aspect of both knees. He had a history of biting his right hand. His response
to foul smells was previously abnormal. There was no history of urinary retention, incontinence, abnormal sweating, or dry eyes. His siblings from a non-consanguineous marriage were normal. On examination, he had excessive dribbling of saliva from the mouth. There were two active grade III pressure ulcers over the anterior aspect of both knees (Figure 5). Multiple abrasion marks were found over the legs (Figure 5). Major portion of the distal phalanges of his right index and middle fingers was missing (Figure 6). He had no withdrawal response to pinprick; however, he could distinguish between cold and hot stimuli. His muscle tone and DTR were normal.

The patient had a raised serum ESR (16 mm at the first hour; normal: 0–9 mm at the first hour). His uric acid level was within normal limits. MRI of his brain was also normal. He had normal motor and sensory NCS in the upper and lower limbs. EMG was normal in all sampled muscles of the four limbs. The patient was also advised to undergo a sural nerve biopsy, but his parents did not agree because of financial constraints. Informed consent was obtained from the father of the patient for the publication of this case report and the accompanying images.

Discussion

The terms congenital indifference to pain and congenital insensitivity to pain (CIP) were used interchangeably until 1970, when they were distinguished and described as two distinct syndromes (1). In patients with CIP, the pain stimulus is not properly transmitted to the central nervous system due to an alteration of the structure or function of the sensory pathways that becomes evident on NCS and nerve biopsy (1,3-6). Nevertheless, reports of CIP cases with normal NCS have been described (7-10). In patients with congenital indifference to pain, the peripheral sensory pathways are intact and sufferers correctly perceive the nature of painful stimuli but fail to react in the normal defensive manner. Rather, they demonstrate a rise in pulse and respiratory rate and an upsurge in blood pressure.
Imaging studies are required to rule out this disorder.

to the loss of the affective-emotional component of pain (13).

pain, and in the anterior cingulate and insular cortices that leads
generate deficits in the sensory-discriminative component of
asymbolia have lesions in the somatosensory cortex, which
should be considered in such patients include CIP , pain asymbo-
diagnosis by exclusion (15). Other differential diagnoses that
congenital indifference to pain (4,14).

It has been suggested that these individuals have a deficit in
their affective-emotional response to pain, rather than a deficit
in their discrimination of painful stimuli (1). A few genetic stud-
ies have provided evidence that SCN9A genetic mutations af-
ing the sodium channel [Na (v) 1.7] play the essential role in
patients discussed in the present report
SCN9A genetic mutations affecting the sodium channel [Na (v) 1.7] play the essential role in
congenital indifference to pain (4,14).

Congenital indifference to pain is generally considered a
diagnosis by exclusion (15). Other differential diagnoses that
should be considered in such patients include CIP, pain asymp-
ia, Fragile X syndrome (11,15), and syringomyelia. Patients with
pain asymbolia have lesions in the somatosensory cortex, which
generate deficits in the sensory-discriminative component of
pain, and in the anterior cingulate and insular cortices that leads
to the loss of the affective-emotional component of pain (13).
Imaging studies are required to rule out this disorder.

Fragile X syndrome is an X-linked recessive disorder char-
acterized by short stature, a long narrow face with mid-facial
hypoplasia, a large head and ears, mental retardation, and be-
behavioral abnormalities. Children demonstrate self-injurious be-
avior; they scratch to produce wounds, bang their head, and
bite themselves (11).

Syringomyelia presents with a loss of pain and temperature
sensibility while light touch, vibration and position senses re-
main intact. Painless ulcers of the hands are frequent. Edema,
hyperhidrosis, and Horner syndrome can occur due to the in-
terruption of autonomic pathways (16). Less frequently, neuro-
pathic osteoarthropathy, which affects shoulder, elbow, or wrist
joints, may develop (16,17). Impaired bowel and bladder func-
tions usually occur as a late manifestation (16).

Patients from northern Pakistan have presented with phenotypes similar to congenital indifference to pain but have been diag-
nosed as having CIP on detection of a genetic defect in Na (v) 1.7 sodium channels (14). All patients discussed in the present report
were diagnosed on the basis of clinical, electrophysiological, and
histopathological findings to be suffering from congenital indifferent-
to pain rather than CIP based on the evidence provided in
previous reports (1,3-6). The patients were brought to the institute
with complaints of self-biting, but they did not have siblings af-
fected by the syndrome. The latter two patients had active wounds
over their knees; however, they were able to differentiate hot from
cold sensations. Excessive dribbling of saliva was observed in the
first and third patients. All patients had an abnormal response to
painful stimuli and normal DTR. MRI brain scans performed in two
patients were normal. NCS and EMG in all cases were normal. Sural
erve biopsy performed in one patient was also normal.

Congenital indifference to pain is a problem that is diffi-
cult to control and impossible to treat (11). Vigilance is required
from parents to prevent recurrent injuries. An interdisciplinary
team management approach should be promoted that includes
the services of medical staff, rehabilitation physicians, psycholo-
gists, and physiotherapists.

Informed Consent: Written informed consent was obtained from
parents of the patients who participated in this case.

Peer-review: Externally peer-reviewed.

Author Contributions: Concept - A.R.Q.; Design - A.R.Q.; Supervision
ing - A.R.Q., S.B.A., A.A.K.; Analysis and/or Interpretation - S.B.A.; Literature

Conflict of Interest: No conflict of interest was declared by the
authors.

Financial Disclosure: The authors declared that this study has re-
ceived no financial support.

References

1. Nagasako EM, Oaklander AL, Dworkin RH. Congenital insensitivity
3. Rahalkar MD, Rahalkar AM, Joshi SK. Case series: Congenital insensi-
tivity to pain and anhidrosis. Indian J Radiol Imaging 2008;18:132-
4. [CrossRef]
4. Goldberg YP, MacFarlane J, MacDonald ML, Thompson J, Dube MP,
Mattice M, et al. Loss-of-function mutations in the Nav1.7 gene underlie congenital indifference to pain in multiple human popu-
5. Dyck PJ, Mellinger JF, Reagan TJ, Horowitz SJ, McDonald JW, Litchy
WJ, et al. Not ‘indifference to pain’ but varieties of hereditary senso-
ty and autonomic neuropathy. Brain 1983;106:373-90. [CrossRef]
1999;14:460-4. [CrossRef]