Congenital insensitivity to pain syndrome accompanied by neglected orthopedic traumas and complications

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Abstract

Pain is a protective mechanism. Congenital insensitivity to pain syndrome is a very rare disease in which there is no ability to feel physical pain. It has been reported that it occurs with an incidence of 1 in 125 million newborn. Patients with congenital insensitivity to pain may have various orthopedic complications such as recurrent fractures, osteomyelitis and neuropathic joints. The most frequently affected body parts are lower extremities. Besides, curvature of spine can be seen. Injuries in epiphyseal points may cause incompatibility of extremity. Charcot joints may develop, which can lead to neuropathic arthropathy as a result of insensitivity to pain. Here, we present a patient with traumatic fracture-dislocation on left hip that neglected the treatment, had a bilateral femur fracture and then had septic arthritis of knee.

Keywords: CIPA, Congenital insensitivity to pain syndrome, orthopedic complications

Introduction

Congenital insensitivity to pain syndrome (CIPA) is a very rare case in which there is no ability to feel physical pain. It has been reported that it occurs with an incidence of 1 in 125 million newborn. Therefore, very little is described in the literature about this devastating condition that both the patient and the family can suffer [1]. These patients do not feel pain in any trauma or injuries throughout their life. They have feeling of hot and cold. Pain is a protective mechanism. Because of lack of pain, bruises, bone fractures, and other health problems may go undetected in such people.

Small children with congenital insensitivity to pain may damage themselves by biting their corner of mouth or fingers. These repeated injuries often lead to a lower life expectancy. It is considered as peripheral neuropathy because it affects the peripheral nervous system.

Here, we present a patient with traumatic fracture-dislocation on left hip, who neglected the treatment, had a bilateral femur fracture and then had septic arthritis of knee.

Case report

An 8 year-old girl applied with pain and swelling in her left knee. Two years ago, the patient underwent osteosynthesis surgery with a diagnosis of right femur shaft fracture after trauma with reduction titanium elastic nail at another center. The implants were removed one year after the operation. She applied to the orthopedic physician with a complaint of
pain in the left hip after a fall again 1.5 years ago. The patient who had a traumatic hip dislocation after the acetabulum fracture, was diagnosed with congenital hip dislocation and was not intervened (Figure 1). However, the hip was in place, when we looked at the older x-rays of the patient. The patient, who had a left femur shaft fracture 1 year ago, underwent closed reduction and fixation with titanium elastic nail at another center (Figure 2).

Destruction of the proximal tibial epiphyses and premature partial occlusion occurred due to the infection after the operation (Figure 3-4). Swelling in the left knee and increase in temperature was observed in the examination of the patient. After the investigations performed, septic arthritis was seen in the left knee. Two titanium elastic nails on the left femur were removed and joint debridement was performed (Figure 4).

Fig. 1. Untreated traumatic hip dislocation which was thought as congenital hip dislocation

Fig. 2. Operated left femur fracture

Fig. 3 and 4. Destruction of the proximal tibial epiphyses and premature partial occlusion
There was no medical history in the patient's family. Her parents were healthy, and no consanguinity was present. Her APGAR score was 6 at one minute and 9 at five minutes. When she was born, there was no hypotonia, hypoxia, hypothermia or hyperthermia. When hospital records were examined, erythrocyte sedimentation rate (ESR) and C-reactive protein (CRP) values were observed to be consistently high. The family said she had had febrile seizures during infancy and they had been warned by the nurses because of not crying of the patient after the injection for the treatment. When looking at the patient's fingers, it was seen that she injured herself by biting her fingertips and some of 2, 3 and 4. Distal phalanges were missing (Figures 5, 6, and 7).

Fig. 5. Wounds due to fingers biting
Fig. 6. Finger bones
Fig. 7. The hands X-ray showing lack of distal phalanges due to fingers biting
It was understood that she has a feeling of hot and cold but it is not very effective. However, it was noticed that she perceives the temperature while drinking tea. We learned from the family that her skin is dry, she never sweats and she took her teeth out herself (Figures 8 and 9). The difference of the case we presented is that there is no mental retardation.

Fig. 8. Wounds in the corner of the mouth due to lips biting

Fig. 9. The self-extracted teeth

**Discussions**

Insensitivity to pain may have different underlying causes such as syringomyelia, diabetes mellitus, leprosy and interstitial hypertrophic neuritis [1]. Insensitivity to pain can lead to various complications such as infection, ulceration and sometimes death [2].

Inheritance is divided into five categories based on clinical features, disease progression, autonomic involvement and specific genetic abnormalities in hereditary sensory autonomic neuropathy [3, 4]. Clinical evaluation of each subtype is necessary in the treatment and prevention of long-term complications. Type 1 (sensory radicular neuropathy) is characterized by sensory disorder particularly in lower extremity in the second and fourth decades of life. Patients often apply with foot joint complications, such as plantar ulcers, soft tissue infections and osteomyelitis. Type 2 (Morvan’s disease) affects all four extremities and occurs during infancy or early childhood, a strong precursor for motor fibers. Patients usually have pathological fractures and sacral anhidrosis. Deep tendon reflexes are not present. Type 3 (Riley eDay Syndrome) occurs in the postnatal period with nutritional failure, recurrent aspiration pneumonia and severe autonomic dysfunction, often ending up with dead in premature birth. Type 5 is characterized by the beginning of insensitivity to pain and temperature, and variable autonomic involvement in childhood [3].

CIPA was first described by Dearborn in 1932 and systematically published by Swenson in 1963 [5]. This extremely rare autosomal recessively inherited disease is described as hereditary sensory autonomic neuropathy type 4. Nerve fiber endings or electrophysiological studies should be assessed microscopically in order to distinguish congenital insensitivity to pain from peripheral neuropathic pains such as diabetes [6].

Diagnosis is confirmed with the aid of genetic tests whereas it can be diagnosed clinically. Thrush et al. suggested using three basic clinical features for diagnosis: the feeling of pain does not exist after birth, the entire body is affected, and all other sensory types and deep tendon reflexes can be used [7].

In addition to clinical evaluation for CIPA diagnosis, pharmacological tests (1:10.000 histamine intradermal reaction) and
neuropathological examination in electron microscopy can also be used. In neurophysiologic examination, isolated axonal sensory polyneuropathy was determined in CIPA patient [8]. A histamine test is performed by intradermal injection of a 1/10.000 solution of 0.05 ml histamine phosphate. Generally, a hole surrounded by an erythematous flare occurs extending from 1 to 3 cm in a normal response; but in a pathological response, there is no flare and the pain is minimal.

Patients with congenital insensitivity to pain may have various orthopedic complications such as recurrent fractures, osteomyelitis and neuropathic joints. The most frequently affected body parts are lower extremities. Besides, curvature of spine can be seen. Injuries in epiphyseal points may cause incompatibility of extremity [9]. Charcot joints may develop, which can lead to neuropathic arthropathy as a result of insensitivity to pain [10].

Mostly weight-bearing joints, particularly knee and ankle joints, are affected. Despite fracture healing, arthropathy is progressive and eventually results in deformity and joint instability. In addition, cases such as auto-amputation of fingers and toes have been reported in the literature. Unexplainable fever and anhydrous, another component of CIPA, can be fatal in some children [11].

The most common complaint in CIPA after birth is recurrent fever secondary to autonomic dysfunction and anhydrous [12]. This is the reason of 20% of deaths under 3 years of age. The most important characteristic is the self-mutilation behavior which causes oral ulceration in lip, tongue and cheeks, self-extraction of teeth, and biting one’s own fingers.

Mutations in the NTRK1 (TRKA) gene affect CIPA. NTRK1 is located on chromosome 1 (1q21-q22) and encodes for autophosphorylated neurotrophic tyrosine kinase receptor type 1 in response to nerve growth factor [13]. However, these patients do not feel pain due to the mutation of the gene encoding the neurotrophic tyrosine kinase receptor.

In rodent studies, the presence of NGF was shown on osteosynthetic cells and the disruption of NGF led to the production of new hypertrophic bone [14, 15]. This can cause significant bone hypertrophy in young patients. Patients with CIPA may develop recurrent, non-healing fractures with significant periosteal reactions and excessive callus formation, and 50% of patients have osteomyelitis and destruction of underlying bone [16].

We reviewed the literature searching for CIPA cases and possibly similar cases. Syringomyelia can lead to the resemblance of pain clenching, therefore leading to differential diagnoses [17]. Cornelia de Lange syndrome (CdLS) is a rare genetic syndrome with clinical findings related to multiple organ systems such as extremities, gastrointestinal tract, skin and central nervous system [18]. Lesch-Nyhan syndrome (LNS) is a rare X-linked genetic disorder in which hypoxanthine-guanine phosphoribosyl transferase (HPRT) enzyme is absent clinically characterized by mental retardation, choreoathetosis, spasticity, hyperuricemia and cerebral palsy [19].

The other diagnoses were excluded after the genetic study which detected mutations in the NTRK1 gene responsible for CIPA in 2015.

The medical history, clinical signs of anhidrosis, pain insensitivity, genetic tests, and negative histamine flare tests of our patient sufficed to endorse the diagnosis of CIPA.

Sixteen cases about CIPA have been reported in the literature. Makari et al. described two brothers with CIPA who suffered from child abuse [20]. In recent days, two cases of CIPA patients who continue treatment have been reported [21]. Generally, effective CIPA treatment has been established within the framework of family education and patient education [22]. Early detection of the disease using techniques such as prenatal genetic screening and amniocentesis is the only way to prevent the birth of an affected child, because treatment is not available [23]. After a child is born with CIPA, prevention of joint diseases is very important. Small cracks can be treated with conservative approaches. In a highly degenerated joint, arthrodesis may be appropriate eventually, but nonunion and infection are common.
Conclusion

Currently, there is no standard treatment for CIPA patients. The most important factors are early diagnosis and being patient under the control of the family members to increase the quality of life by protecting from self-injury and other harms. Paying particular attention on patients, who has had extremity injury, can prevent complications and deformities; thus, the quality of life of the patients can be improved.

Conflict of interest

The author(s) declare that they have no competing interests.

Consent

Written informed consent was obtained from the patient's parents for publication of this case report and accompanying images.

References


