CONGENITAL INSENSITIVITY TO PAIN WITH ANHIDROSIS
REPORT OF A CASE AND REVIEW OF THE LITERATURE

S. D. THEODOROU¹, A. E. KLIMENTOPOULOU², E. PAPALOUKA¹

In a previous paper published in this journal, we reported two cases of “Congenital Sensory Neuropathy with Anhidrosis” with reference to the orthopedic complications (Theodorou et al., 1985). We now present a new typical case, under the currently used term: “Congenital Insensitivity to Pain with Anhidrosis” (CIPA) and a brief review of the literature on the incidence, etiology and problems arising in various systems. CIPA is an autosomal recessive form of sensory neuropathy manifesting with typical clinical features. Universal insensitivity to pain, anhidrosis or hypohidrosis, bouts of hyperpyrexia from very young age, self inflicted injuries, defective or absent lacrimation and mental retardation are specific diagnostic findings. Orthopedic, maxillofacial, dermatological and ophthalmologic complications are common. Counseling of the family and school personnel for the prevention of injuries is necessary. Early diagnosis is very important for the prevention and treatment of various complications. The etiology and pathogenesis of the condition is still unclear. The recent detection of a new gene, which encodes a receptor tyrosine kinase for nerve growth factor and lately of a specific point mutation associated with the gene inactivation11, may open new ways for the study and management of this disabling condition.

Key words: insensitivity, neuropathy, anhidrosis, pain absence.

Mots-clés: insensibilité, neuropathie, anhidrose, absence de douleur.

INTRODUCTION

The term “Congenital Insensitivity to Pain with Anhidrosis” (CIPA) which is in current use was introduced by Swanson (1963) in his classification of sensory neuropathies.

Previously, Gillespie and Perruca in 1960 described the case of a 3-year-old child with analgesia, hypohidrosis and mental retardation under the title “Congenital Insensitivity to Pain”. Other terms that have also been used include “Familial Sensory Neuropathy with Anhidrosis” (33), “Congenital Sensory Neuropathy with Anhidrosis” (42) and “Type IV Congenital Sensory and Autonomic Neuropathy” (6). A new typical case with six years follow-up and a review of the literature on the genetics, pathophysiology and the problems arising, are presented.

CASE REPORT

The patient is the second of dizygotic male twins born by caesarian section with a birth weight of 2800 gr. The pregnancy was uneventful except for slight bleeding at the eighth week. His 42-year-old healthy mother and his 50-year-old diabetic father were unrelated. Soon after his birth his mother noticed that he was “different” from his twin brother. He seemed weaker and there were difficulties in feeding him. He presented bouts of high fever particularly in hot weather, without sweating. Because of the hyperpyrexias and food refusal, he had to be

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admitted to the children's hospital at the age of 8 weeks and was treated for bronchopneumonia with antibiotics. After eruption of teeth at the age of 7 months, he was noticed to bite his tongue, lips and fingertips causing bleeding. Later he kept losing his teeth either by biting or self-extraction. He walked at the age of 15 months and spoke words at 17 months. He continued presenting intermittent fevers and bouts of bronchitis (postinfectious asthma), which became less frequent with age. He was and his mother said he never sweated, even in very hot weather or when running high temperatures. He was noted also to touch hot objects and receive burns without complaining. Nevertheless, superficial sensation to light touch was intact. There was some scarring on his lips and tongue, the tip of which was missing as a result of biting. For the same reason his fingertips showed scarring and absorption. Many of his teeth were missing. Bruises in several parts of his body were also noted. Corneal reflexes were absent and tear production was reduced. Tendon reflexes were hyporeactive. He was hyperactive trying to compete with his healthy twin brother, but was cooperative. Routine laboratory tests were unremarkable. Radiographs of both tibias showed increased bone density of the metaphyseal-epiphyseal region of the proximal end of the right tibia, hyperplastic bone formation and disorganization. (fig. 1-A). An above knee plaster of Paris was applied which was soon destroyed and had to be replaced repeatedly. Radiographs 4 months later showed reorganization of the proximal end of the right tibia, but periosteal reaction and increased density was noted in the metaphyseal region of the upper end of the left tibia (fig. 1-B).

Fig. 1. — A. Bone density of the proximal part of the right tibia following fracture and neuropathic knee joint.

seen for the first time at the age of 4 years for swelling of the proximal end of the right tibia. Pressure over the swelling was painless as he was walking and running. His mother said he never protested when given injection or after falls. Pressure over tendons or muscles did not produce any pain nor to the pinprick in various areas of the body. His skin was unusually dry, hard and hirsute

Fig. 1. — B. Good reconsrtuction after 4 months of immobility in plaster. The proximal part of the contralateral tibia shows densification and periosteal reaction.
An above knee plaster of Paris was applied to the left leg, which was again repeatedly destroyed and replaced (fig. 1-C) and immobilization had to be continued for another three months until satisfactory reconstruction in both tibiae was demonstrated (fig 1-C). Ulcerations of the anterior surface of the knees and a perforating ulcer in the foot were treated with debridements, aseptic dressings and efforts to prevent external pressure on these areas by various means with great improvement.

A few months later he sustained a non-displaced fracture of the right elbow, which healed with immobilization in plaster for two months. On a recent examination at the age of 9 years, the patient was still very active but more cooperative. His height was 122 cm and weight 23 kg (both at the 10th percentile). His general condition was good. Hyperpigmentation was noted in the posterior aspect of the elbows and over the knees. Tendon reflexes were diminished, abdominal and cremaster reflexes were hypoactive. Radiographs showed no abnormalities in the spine and hips. Radiographs of the knees showed complete healing but the proximal growth cartilage of both tibiae was narrow (fig. 1-D). The right elbow was normal and the left one showed neuropathic changes (fig. 2-A). The right ankle appeared normal, but the left was swollen and radiographs showed neuropathic changes (fig. 2-B). Most of his teeth were missing and scarring of the lips and the tongue in association with wasting of the maxillofacial area, gave an appearance resembling a “toothless old man” (fig. 3).

![Fig. 1. — C. In spite of immobilization in plaster for three months, bone remodeling of the proximal end of the left tibia was not satisfactory.](image)

![Fig. 1. — D. Radiograph of knees at the age of 9 years shows good remodeling but growth plates of proximal tibial epiphysis are narrowed.](image)
The fingertips of most fingers were missing and the nails were dystrophic. Ulcerations on the anterior surface of the knees and posterior surface of the elbows had recurred, but there had been no new fracture for the last 13 months. The knees and the ankles were slightly swollen and the skin of the legs below the knee was edematous with some hard nodules (fig. 3). Despite the family’s support and the help from psychologists the patient manages to escape supervision and to take part in sports like basketball and football.

Fig. 2. — A. Neuropathic left elbow joint.

Fig. 3. — Photograph of the patient at the age of 9 years shows scarring of lips and wasting of maxillofacial area. The fingertips of almost all fingers are missing and their ends are rounded. Ulcerations of knees and the feet, swelling and deformity of the left ankle are also noted.
DISCUSSION

Congenital Insensitivity to Pain with Anhidrosis (CIPA) is an autosomal recessive condition, presenting with a wide spectrum of typical clinical symptoms and signs. The classical clinical presentation in neonates and infants includes apathy, hypotonia, often refusal or inability to feed, dry and hot, non-sweating skin even in high summer temperatures. Frequent episodes of unexplained hyperpyrexia occur, particularly in hot weather during the first years of life. This may result in the death of 20% of the affected children (34). Iwanaga et al. (1996) reported magnetic resonance imaging and cerebrospinal fluid abnormalities in a 9-month old girl with CIPA and hyperpyrexia, complicated by generalized tonic convulsions and coma. Juri et al. (1997) observed encephalitis with generalized tonic convulsions in a 14-month old girl, as a result of heat stroke. Mental retardation of mild degree is usually present and motor abilities may be delayed. Deep sensation is absent and the superficial may be diminished or normal. Many children fail to thrive, the height and the weight percentiles being below average. After eruption of the teeth, biting and scarring of the lips is common and loss of the tip of the tongue results in a round or bifid appearance. Autoextraction of teeth is also common and decubital ulcers of the ventral surface of the tongue from the incisors may lead to severe problems including bleeding, infection, halitosis and malnutrition (2, 5, 23, 32). Maxillofacial problems may also require special care (32).

As a result of biting, fingertips appear scarred, smaller and rounded with dystrophic nails. Burns and self-inflicted lesions of the skin, mainly of the extremities, are common (13, 16). Lacrimation may be defective or normal. Severe opthalmologic problems may arise from corneal ulcers and introduction of foreign bodies (43).

Fractures may occur early in the first months of life (29, 36, 24) and become frequent with time as the child becomes mobile. Complicated fractures, dislocations, neuropathic joints and infection necessitated amputation in many cases (27, 18, 12). All parts of the skeleton may be affected including the spine (8). Nagayama et al. (31) reported spinal cord injury at birth in a neonate with CIPA.

Diagnosis or differential diagnosis of CIPA from other conditions presenting with absence or depression of pain can be easily established on the basis of the distinct clinical picture. The cardinal features of insensitivity to pain, anhidrosis, frequent hyperpyrexias and mental retardation are not found in any other type of neuropathy.

CIPA has been considered as a rare condition. In 1976 Lee et al. estimated the number of reported cases as 7, Theodorou et al. (1985) as 28, Rosenberg et al. (1994) as 31 and Edwards-Lee et al. (1997) as 32 (7). In reviewing the literature we found 51 cases reported in which the general, pediatric, orthopedic and neurological aspects of the condition were presented, 50 cases dealing with opthalmologic, dermatologic, maxillofacial and oral problems and 9 cases in which aspects of histology, genetics and research were discussed, a total of 110 cases.

It is probable that there is overlapping in certain cases, with the same case reported by different authors devoted to different aspects of the condition of the same patient and therefore an accurate estimation is difficult to make. Nevertheless, we believe that the increasing number of reported cases during the last few years has contributed to the earlier diagnosis, study and management of the condition.

Except for the mode of inheritance and the specific gene mutation, the pathophysiology of CIPA remains unclear. In table I, the type of investigations performed to date in patients and the corresponding findings/results are demonstrated. There are some contradictory results, a fact that can be partly attributed to the limited number of times these investigations are performed. The almost universal finding of extreme paucity of small myelinated fibers as well as the lower density of unmyelinated fibers (which convey the sensess of pain and temperature) in electron microscopy in sural nerve biopsy, appears as a pathognomonic feature along with the almost stereotyped clinical presentation (10, 29, 14, 18, 20, 39, 4, 34).

Table I also reflects the different theories on the pathophysiology of the disease. Briefly, these have
Table I. — Investigations performed in patients with CIPA with corresponding findings/results*

<table>
<thead>
<tr>
<th>Type of investigation</th>
<th>Findings/results</th>
</tr>
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<tbody>
<tr>
<td>Full blood count, Serum electrolytes, Renal function</td>
<td>Normal</td>
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<tr>
<td>Tests, Lipid profile, Uric Acid, Blood pH, Lactic acid (serum), Urine organic acids,</td>
<td>Normal/SLightly elevated (7)</td>
</tr>
<tr>
<td>Serum aminoacids, Electromyography, Cerebrospinal fluid protein, cells, glucose,</td>
<td>Normal/SGOT, LDH elevated (19)</td>
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<tr>
<td>chloride, Karyotype, Sural nerve biopsy (light microscopy), Vitamins B6, B12, C and</td>
<td>Elevated (19)</td>
</tr>
<tr>
<td>folic acid</td>
<td>Normal/IgA levels transiently decreased (41)</td>
</tr>
<tr>
<td>Ammonia</td>
<td>Normal/diffuse slowing/other non specific findings (7)</td>
</tr>
<tr>
<td>Liver function tests</td>
<td>Normal/decreased/absence of action potentials in ulnar and sural nerve (3)</td>
</tr>
<tr>
<td>Creatine kinase</td>
<td>Normal/elevation of myelin basic protein (19)</td>
</tr>
<tr>
<td>Immunologic studies</td>
<td>Decreased (30)</td>
</tr>
<tr>
<td>Electroencephalogram</td>
<td>Slightly elevated (35)</td>
</tr>
<tr>
<td>Nerve conduction tests (motor and sensory)</td>
<td>Wheal with no axonal flare</td>
</tr>
<tr>
<td>Cerebrospinal fluid electrophoresis</td>
<td>No sweat production</td>
</tr>
<tr>
<td>Cerebrospinal fluid Substance-P</td>
<td>No miosis (7)</td>
</tr>
<tr>
<td>Urine HVA/VMA** ratio</td>
<td>Normal/non specific dilatation of the third or the lateral ventricles</td>
</tr>
<tr>
<td>Histamine intradermal injection</td>
<td>Normal/changes consistent with hyperpyrexia</td>
</tr>
<tr>
<td>Pilocarpine iontophoresis</td>
<td>Normal/lack of innervation of sweat glands</td>
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<tr>
<td>Ocular Pilocarpine 0.05%</td>
<td>Lipid droplet accumulation and reduced cytochrome C oxidase</td>
</tr>
<tr>
<td>Computed tomography brain scan</td>
<td>histochemically/neurogenic and myogenic changes</td>
</tr>
<tr>
<td></td>
<td>Lack/reduction of small myelinated fibers and decreased unmyelinated fibers</td>
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<tr>
<td></td>
<td>Absence of Lissauer tract, loss of small myelinated fibers in dorsal roots and</td>
</tr>
<tr>
<td></td>
<td>spinal tract of trigeminal nerve, loss of small ganglion cells of dorsal ganglia</td>
</tr>
<tr>
<td></td>
<td>TrKA-NGF gene defect</td>
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</tbody>
</table>

* Literature references shown either when test performed only once or when findings/results differ from the ones usually reported.

** Homovanillic acid/Vanillylmandelic acid.

been focused on five main hypotheses: I) Disturbance of catecholamine metabolism (35), II) Faulty development of neural crest derivatives (18), (III) An inborn error of metabolism involving a minor pathway of tyrosine metabolism (7), (IV) Substance-P neurotransmission defect (30), V) Low biological activity of nerve growth factor (TrKA/NGF gene), in an Ecuadorian family using gene sequencing (15). Moreover, Greco et al. in January 2000 found a specific point mutation that is associated with the inactivation of this gene. Nerve Growth Factor (NGF) induces neurite overgrowth and promotes survival of embryonic sensory and sympathetic neurons. This newly detected gene encodes a receptor tyrosine kinase for NGF. This in turn suggests that defects in TrKA cause CIPA and that the NGF-TrKA system has a crucial role in the development and func-
tion of the nociceptive reception as well as establishment of thermoregulation via sweating in humans. Mardy et al. and Yotsumoto et al. (1999) have confirmed the findings mentioned above.

Early diagnosis of the condition may save the life of neonates and infants suffering from complications such as hyperpyrexia and protect children from exposure to various dangerous activities. The management of bone and joint lesions is extremely difficult. With the protecting benefit of sensation absent, patients overuse the bones and joints, resulting in fractures and dislocations. For psychological reasons affected children are hyperactive and tend to demonstrate their particular qualities and abilities resulting from the absence of pain.

Fractures are more common in the lower limbs particularly proximal to the knee and also ankle and hip joints with consequent neuropathy of the adjacent joints, but practically all bones and joints may be affected. Fractures must be suspected in every bruising or swelling especially near the metaphyses of the long bones. They should be properly immobilized as soon as possible. Usually, well-padded plaster of Paris was applied which should be constantly checked by the parents and replaced immediately if it is worn out. In our case fractures of the proximal tibias and neuropathic knee joints showed satisfactory healing with immobilization in plaster. Fixation of fracture of the lateral condyle of the elbow with Kirschner wires failed. Therefore if internal fixation is indicated, this must be rigid. In one case of an 11-week-old male infant with non-union of a fracture of the ulna, internal rigid fixation with dynamic compression plate was successful (24).

Dislocations are difficult to deal with particularly in neuropathic joints. Attempt for surgical treatment of congenital hip dislocation has failed (22).

Ulcerations in the posterior aspect of the elbows, knees, heels, hands and other parts of the body are managed by local care of the wounds, aseptic dressings and avoidance of external pressure by covering the area with thick dressings, bandaging and other means.

Of utmost importance, are the continuous information, guidance and help of the whole family by psychologists and social workers. The school personnel should also be informed of the dangers of injuries. The idea is to “steal time”. If children are not severely mentally handicapped or uncooperative, they learn with time to avoid exposing themselves to dangerous activities and excessive exertion.

REFERENCES


SAMENVATTING


De auteurs stellen een typisch geval voor van aangeboren ongevoeligheid voor pijn met anhidrose, alsmede een overzicht van de literatuur naar frequentie, etiologie en problemen veroorzaakt door deze aandoening. Het gaat om een autosomaal recessieve aandoening met een sensoriële neuropathie met volgende kenmerken: ongevoeligheid voor pijn, anhidrose of hypohydrose, koortsperiodes bij jongere patiënten, auto-traumata, geen of weinig tranen en lichte mentale achterstand. Orthopedische, maxillofaciale, dermatologische en oftalmologische complicaties zijn frequent. Een vroegtijdige diagnose is belangrijk. Alhoewel de etiopathogenese nog onbekend is, zal de ontdekking van een gen verantwoordelijk voor een factor voor de ontwikkeling en de groei van zenuwen een doorbraak betekenen in de studie en behandeling van deze aandoening.

RÉSUMÉ


Les auteurs presentent un cas typique d’«insensibilité congénitale à la douleur avec anhidrose» et aussi une revue de la littérature sur la fréquence, l’étiologie et les problèmes provoqués par la maladie. Il s’agit d’une forme autosomique récessive de la neuropathie sensorielle qui se manifeste par des signes cliniques typiques: insensibilité à la douleur, anhidrose ou hypohydrose, crises d’hyperpyrexie chez des sujets jeunes, auto-traumatismes, pas ou peu de larmes et retard mental léger. Des complications orthopédiques, maxillofaciales, dermatologiques et ophtalmiques sont fréquentes. Le diagnostic précoce est très important. Bien que la physiopathologie soit pas encore connue, la découverte d’un nouveau gène responsable du facteur déterminant le développement et la croissance des nerfs pourrait faire progresser l’étude et le traitement de la maladie.