Anorexia, a First Sign of Diagnosis in Severe Pediatric Disorders

RAMONA-MIHAELA NEDELCUTA1, GIGI CALIN1, VLAD DUMITRU BALEANU2, DRAGOS VIRGIL DAVITOIU3, DANIEL-ILIAN VOICULESCU4, CECIL SORIN MIREA5, TIBERIU STEFANITA TENEA COJAN6, BOGDAN SOCEA7, DRAGOS OVIDIU ALEXANDRU8, DIANA CLENCIU9, VICTOR GHEORMAN10, ION UDRISTOIU11, VERONICA CALBOREAN12, COSMIN ALEXANDRU CIORA12

1University of Medicine and Pharmacy of Craiova, Pediatrics Department, 2 Petru Rares Str., 200349, Craiova, Romania
2University of Medicine and Pharmacy of Craiova, Surgery Department, Clinical Emergency Hospital Sf. Pantelimon Bucharest, 340-343 Pantelimon Road, 021659, Bucharest, Romania
3University of Medicine and Pharmacy of Bucharest, Surgery Department, Clinical Emergency Hospital Sf. Pantelimon Bucharest, 340-343 Pantelimon Road, 021659, Bucharest, Romania
4University of Medicine and Pharmacy Carol Davila Bucharest, Department of Surgery University Emergency Hospital Bucharest, 169 Sf. Iuliu Indepenei, 050098, Bucharest, Romania
5University of Medicine and Pharmacy of Craiova, Surgery Department, 2 Petru Rares Str., 200349, Craiova, Romania
6University of Medicine and Pharmacy of Craiova, Department of Medical Informatics and Biostatistics, 66 1 Mai Blvd, 200638, Craiova Romania
7General Surgery Clinic, Emergency Clinical Hospital Sfantul Pantelimon, 340-342 Pantelimon Road, 021659, Bucharest, Romania
8University of Medicine and Pharmacy of Craiova, Department of Medical Informatics and Biostatistics, 66 1 Mai Blvd, 200638, Craiova Romania
9University of Medicine and Pharmacy of Craiova, Department of Metabolism and Nutrition Diseases, Filantropia Clinical Hospital of Craiova, 1 Filantropiei Str., 200143, Craiova, Romania
10University of Medicine and Pharmacy of Craiova, Psychiatry Department, Neuropsychiatry Hospital of Craiova, 24 Potelul Alley, 200473, Craiova, Romania
11University of Medicine and Pharmacy of Craiova, Cardiology Department, 2 Petru Rares Str., 200349, Craiova, Romania
12Carol Davila University of Medicine and Pharmacy, Discipline of Gastroenterology and Hepatology. 8 Eroii Revolutiei Blvd., 050474, Bucharest, Romania

Anorexia, as a symptom, is part of the complex picture of a large pediatric pathology. Through attentive observation, anorexia as a single sign, may announce the onset of serious and particular affections. In all 4 cases, anorexia was a singular sign for a period of time before the complete illness was established. Whether adenoid cyst, cerebral abscess, pontocerebellar atrophy, Wilms tumor, neuroendocrine system of hunger regulation, satiety, complex control achieved through interactions of the limbic-hypothalamus-cortex system, come into operation with the occurrence of organic anorexia, different of the psychogenic anorexia (usually common in adolescents). The study is an alarm signal on the importance of a symptom in early diagnosis, and the rapid establishment of therapy in severe life threatening conditions.

Key words: anorexia, symptom, diagnosis

Anorexia is an eating disorder with physical and emotional implications for the patient. Moreover, it can be considered a lethal psychiatric disorder with an increased risk of death (4 times higher than in major depression - in a study on adults.) Practically, that morbid condition is characterized by the loss of desire or appetite, which occurs as a symptom in the course of general illness or as a long-lasting behavior in the presence of psychomotor or neuroendocrine disorders [1,2].

Hunger and satiety are part of a complex system of behavioral regulation and integration involving the hormonal system (leptin, ghrelin, insulin and cholecystokinin - contributing in slowing gastric evacuation), nervous regulation by hypothalamus and limbic system and involves neuropeptides, glycemic homeostasis and other factors (regulation by hypothalamus and limbic system and involves neuropeptides, glycemic homeostasis and other factors) with anorexic role - glucagon like peptide GLP1[3].

Anorexia nervosa is a rarer entity in pediatric pathology, rather than in adults, especially in adolescents[4]. Possible causes

Newborn -NS (nervous system) disease, birth defects, meningo-cerebral haemorrhage, congenital bupharyngeal malformations, cardiovascular disorders, metabolic diseases.

Infant - Iron and vitamin C deficiency, chronic digestive diseases, meningo-cerebral sequelae, encephalopathies, oral antibiotics, sulphamides, vitamin D intoxication, food and care mistakes, overuse, conflicting situations, affective frustrations, infections, genetic factors are involved in an unsolved proportion.

Causality diagnosis is extremely difficult, laborious, often a diagnosis of exclusion. There are rare diseases found in current pediatric practice that may have as anorexic onset.

Experimental part

Material and method

4 cases where anorexia was the only sign of onset of the condition
Case I -O.I. 2 years old of cerebral abscess
Case II -G.V. 5 years old adenoid cyst
Case III -V.N. 6 years old PCH2 (pontocerebellar hypoplasia) syndrome
Case IV -P. K. 3,6 years old Wilms’ tumor

Case I. A 2-year-old patient (coming from a family with religious and food taboos) has a serious overall condition, anorexia for about 2 weeks, dehydration, sporadic vomiting and diarrheal stools at home, conscious, temporospatial disorientation.

* email: baleanuvlad@gmail.com; Phone: 0040765865886    # The authors contributed equally to the manuscript and share first authorship

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The patient presented 1 year and 3 months ago, 2 moments of temporary loss of consciousness, of unspecified duration, accompanied by muscle contraction, and in the last 3 months there was a regression in neuromotor function.

Clinical - poor nutritional condition, diminished adipose tissue, cutaneous and mucosal pallor, slow skin turgor response, tachycardia cardiac sounds, AV = 120 beats per minute, systolic murmur II (functional due to severe anemia), dyspnea sive materia, FR = 50resp/min., without evidence pulmonary manifestations, excavated abdomen, diarrhea stools, spontaneous urination, rare, without sniff neck, neuromotor retard.

Hydroelectrolytic rebalancing treatment with low rhythm has been established in order not to worsen functional cardiac failure, oxygen therapy, antibiotic therapy, antidiarrheal therapy.

At 2 h after admission there was an irreversible cardiorespiratory stop and death.

The anatomopathological diagnosis is:
1. Right hemisphere cerebral abscess
2. Protein-calorie malnutrition
3. Heart failure
4. Severe anemia
5. Epidemic
6. Severe psychomotor retardation
7. Neglected child-social case

Case I - Cerebral abscess multiple causes - explanations
1. Somatic impairment of cerebral structures / disorganisation ½ left hemisphere
2. Alteration of the limbic system
3. Severe retardation, mastication disorder, swallowing
4. Infectious systemic impregnation that acts as a thief calories
5. Persistent / vegetarian food taboos
6. Neglecting closely with the family socio-economic level

Case II - Rahnoid cyst

A 5-year-old patient with severe anorexia. The clinical examination is normal, the laboratory tests are in normal parameters for the child’s age, but the anamnesis reveals a c by bicycle fall 5 months ago. After CCT, the child did not experience loss of consciousness, headache or vomiting.

The imaging examination specified the diagnosis of arachnoid cyst with dimensions 4.5/5.5 cm without associated lesions localized left fronto-temporoparietal cerebral cortex. 2 months after surgery, CT scan reveals an anterior temporal mass and 10 months postoperatively the dimensions exceed the previous ones to 5.5 / 6.9 cm, with cyst-like appearance inserted in the Sylvian fissure with minimal neighborhood affection on left cerebral artery.

At 5 years post intervention, the cyst has 6.5/4.5/7.6 with left temporoparietal localization with effect on the frontal lobes, left temporoparietal inserted at the Sylvian fissure with its deformation, as well as deformation of the skull, ventricular dilation, cortical gliosis and areas of atrophy in the left fronto-temporoparietal cortex (fig. 1.2).

Atrophy has become severe, with the metabolic status of the patient being altered.

It also associates other diagnoses: severe anorexia, hypopitrophy, secondary anemic syndrome, iatrogenic epilepsy, spasmodilia, impaired seizures, transient neurologic deficit, recurring cyst.

In this case the causes of anorexia are:
Cyst compression on the neighboring structures causes stimulation on limbic, paralimbic structures involved in the regulation of hunger and satiety [5].

Case III
6-month-old baby with normal weight=3500g, Apgar 8, cesarean delivery, has approximately 5 weeks of anorexia that is accentuated in time.

At the clinical examination is found malnutrition with current weight =170 g, anemic syndrome associated with psychomotor retardation (does not respond to visual stimuli, sounds, does not catch objects, does not sit, has difficulty in supporting the head).

A gay diet is set up, the child not having a correct coordination of the act of swallowing, and sending evidence for the genetic analysis of the karyotype of both parents and the child. The results showed severe chromosomal abnormalities in parents and children: interstitial duplication of the short Y chromosome arm, interest in the p22-12 region (in both parents), interstitial deletion of the long arm of chromosome 11 (to father), interstitial deletion of the short arm of chromosome 12, in the region p12.21 (to father), an interstitial deletion of the short arm of chromosome 1 in the region p36-32 (to the mother). The child has three mutations: an interstitial duplication of the short arm of the Y, the region p22.12 (also present to father), an interstitial deletion of the short arm of chromosome 11, which interests the region q 24.2 (also present to father), and an interstitial deletion of the long arm of chromosome 12 that interests the region p11.21 (also present to father).

The diagnosis of PCH2 has been established.

Pontocerebellar atrophy directly affects the process of muscular coordination involved in the act of nutrition (mastication, swallowing), seizure, and aggravation of the retardation over time. The clinical aspect is inconsistent with survival.

Over time, severe anorexia has been associated with other diagnoses. At 1 year, the child presents severe malnutrition, severe anemic syndrome, epilepsy, severe anorexia. Severely psychomotor retard. At 1.2 years, the patient died.

Anorexia’s explanation associated with severe chromosomal abnormality results in direct damage to
muscle tone coordination, muscle atrophy and secondary malnutrition as immediate consequence.

The occurrence of paroxysmal convulsive disruptions aggravates psychomotor retardation, causes secondary cerebral atrophy, affects the superior integrative control that cerebral cortex has in the nutrition process.

Involvement of mastication and swallowing acts by cerebellum stimuli is completely impaired, the act of nutrition being disorganized (fig. 3).

2. TNFα secretion has been associated with anorexigenic effect

3. Visceral / muscular damage causes changes in hormonal homeostasis (leptin, ghrelin, insulin)

4. Cytostatic therapy with anorexic role

Results and discussions

The limbic system is the oldest phylogenetic constituent of the cerebral hemisphere. It is in the form of a ring composed of primitive cortical tissue in which the limbic and paralimbic structures are made up. The limbic structures are hippocampus, amygdala, limbic striatum with acumbens nucleus, mesolimbic dopaminergic tracts, nonspecific thalamic nuclei, hypothalamus, piriform cortex, substance called septum, midbrain tegmental area and fornix [8](fig. 6).

Fig. 3 Macroscopic aspects: swelling of the left cerebral hemisphere (left), and diffuse cerebral tissue as wall (right)

Case III - PCH2 - explanations

Pontocerebellar atrophy directly affects the process of muscular coordination involved in the act of nutrition (mastication, swallowing), seizure, and aggravation of the retardation over time. The clinical picture is inconsistent with survival (fig. 4).

Fig. 4 Karyotype

Case IV

A 3,6 years old patient with anorexia for approximately 2 weeks, with 900g weight loss in this period (12.7kg).

The medical exam does not bring pathological data. The laboratory test shows a slight increase in the non-specific parameters associated with inflammation (ESR, PCR, fibrinogen).

The abdominal echography shows the presence of a mass of approximately 4.3 / 3.4, in the upper pole of the left kidney, nonhomogeneous echographic appearance with areas of tissue necrosis that does not exceed the renal capsule but compresses the renal sinus, the diagnosis being established as Wilms tumor, diagnosis subsequently confirmed postoperatively, both macroscopically, microscopically and by immunohistochemistry [6]. The patient followed the full oncologic treatment protocol post-surgery, being declared cured at 3 years post-intervention [7]. Anorexia persisted and increased during the chemotherapy series, currently 5 years after the intervention, the child being completely recovered somatic, nutritional, neuropsychic (fig. 5).

Case V - Wilms tumor - explanations

1. Impregnation caused by oncological affectation causes increased catabolism

2. The paralimbic structures are: the parahipocampic gyrus, the cingulate gyrus, the temporal pole, the island and the caudal portion of the orbitofrontal cortex. The hippocampus participates in autonomous, neuroendocrine, memory, emotion function. It has many afferents and a single afferent (fornix) ending in the mammillary body of the hippocampus.

Amygdala is a structure with an increased number of afferent and eferent, communicating with the hypothalamus, thalamus, corpus striatum, cortical, cerebral trunk and directly intervenes in the regulation of homeostasis. Additionally, it contributes to the mediation of the final somatic limbic pathways and visceromotors.

A part of the basal lymph nodes (the caudate and lenticular nucleus) is linked to the olfactory tuber and the acumbens nucleus. The talamus is a very complex structure with vast heteromodal areas interconnected with the limbic cortex receiving signals from the anterior thalamic nuclei, from the hypothalamus, from the vagus nerve, and sends signals to the hypothalamus and the midbrain. It receives tactile, visceral, proprioceptive sensory information and signals [9].
The limbic system, in its entirety, is an interwoven region between the external environment and the internal world. It powers the upper areas of the cerebral cortex and the instinctive parts of the nervous system of the trunk and the marrow as well as the autonomous NS. The hippocampus, due to the multitude of information received, can achieve a synthesis and influence of the hypothalamus, with multiple implications in somatovisceral functions [10].

The frontal lobe is the highest level of autonomous function integration.

Experiences on it (damage / stimulation) cause changes in bladder, intestinal, temperature and sweat control.

The hunger center is made up of neurons that form the ventrolateral nuclei of the hypothalamus. Stimulation is done at the homeostatic level of hypoglycemia, destruction causes anorexia. The center of satiety is located in the ventromedial nucleus of the hypothalamus. Stimulation is achieved by fatty acid hyperglycemia [11].

The studies on the mechanism of reflex regulation of food intake are incomplete because a multitude of chemical stimuli (olfactory, taste), homeostatics (sugar, lipid, enzyme, protein) act synergistically on the hunger center (fig. 7).

In practice, an emotional component, an instinctual one, another learns through various associations, involving the multiple structures (cortex, hypothalamus, limbic system, tonsils) that contain a vast reactive inheritance underlying the act of nutrition. The starting point is profound at the level of organogenesis and ends at the behavioral level through the development of feeding rituals based on mechanisms of knowledge. The hypothalamus is the one that will achieve the neuropsychic level of superior adaptive integration [12-18] (fig. 9).

Child anorexia is an onset symptom that may indicate a severe affection. Even singular, in the initial form of debut, we must give this important sign, without being minimized.

Investigations should be complex and in-depth, starting with thorough medical history, complete clinical examination, laboratory, imaging, genetic or immunohistochemistry tests. It is a complete protocol that eliminates possible artefacts that would delay a correct diagnosis and the application of the curative treatment potential.

We are in era of developing and influencing aberrant eating behaviors, an era in which psychogenic anorexia, especially in adolescence, but also in school or preschool children, including those deprived of parental affection, is experiencing an unprecedented boom.

The path of clinician thinking may be experiencing multiple traps, and relying on stereotypes, past experiences, is and can be extremely damaging [19-26].

After removing the banal medical / digestive diseases, cancers, chronic inflammation, anosmia, loss of taste, known cerebral diseases, depression, anxiety, medication (amphetamine, antibiotics, antiinflammatory etc.) or bad eating habits, we must thoroughly investigate this cardinal signal of the body to diagnose / exclude a possible rare disorder of pediatric pathology.

Conclusions

The complexity of anatomical, nervous, hormonal, homeostatic, behavioral, mental relationships makes anorexia an extremely extensive chapter in ongoing research.

This paper is an alarm signal that shows the importance of complete investigation from sign / symptom to anamnesis and correct diagnosis.

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