IDIOPATHIC ESOPHAGEAL ACHALASIA: A STUDY OF ETIOLOGY AND PROFILE OF THE PATIENTS

Acalásia idiopática do esôfago: análise da história clínica e antecedentes na etiologia e perfil dos pacientes

Gustavo Carvalho de OLIVEIRA, Luiz Roberto LOPES, João de Souza COELHO-NETO

ABSTRACT – Background - The idiopathic esophageal achalasia is a disease of unknown etiology, characterized by esophageal aperistalsis and failure of its lower sphincter with dysphagia. Its etiology factors includes: esophageal gastric junction obstruction, degeneration of Auerbach’s plexus, virus infection, congenital origin, autoimmune affection and injury by toxic agent. The achalasia diagnosis is reached after excluding Chagas disease possibilities, which includes seronegative results for Trypanosoma cruzi, absence of megacolon and epidemiology for Chagas disease.

Aim - To characterize the disease and propose hypothesis concerning its etiology and associated factors.

Methods - Review of medical records from 78 patients operated at the Hospital de Clinicas da Unicamp obstruction between 1989 and 2005 and the subsequent interview, using directed questionnaire, reaching for common data between them and emphasizing history, possible co-morbidities and associated factors. In the group of 78 records collected it was possible to contact and interview 33 patients.

Results - The main findings of this study were: 1) presence of a triggering relevant emotional factor before the symptoms (80%) and over 30% with psychiatric and/or psychological treatment reported; 2) typical childhood infections highly prevalent (88% measles, varicella, rubella); 3) possible associations with: exposure to chemicals, especially herbicides; other diseases of the gastrointestinal tract, autoimmune diseases, genetic propensity and other changes in the nervous system highlighting the seizures.

Conclusions: The idiopathic esophageal achalasia is probably an autoimmune disease, which seems to be highly related to emotional problems.

RESUMO – Racional - Acalásia idiopática do esôfago é doença de etiologia desconhecida, caracterizada por não relaxamento do esfíncter esofágico inferior provocando disfagia. As causas estudadas incluem: obstrução na junção esofagogástrica, degeneração do plexo nervoso intramural, infecção por vírus, doença de origem congênita, afeção autoimune e lesão por agente tóxico. O diagnóstico é feito após a exclusão de doença chagásica, encontrando-se sorologias negativas, ausência de megacólon e antecedentes epidemiológicos negativos para Chagas. Objetivo - Avaliar a caracterização da doença e elaboração de hipóteses relativas à sua etiologia.

Métodos - Foram levantados prontuários de 78 pacientes operados no Hospital de Clinicas da Unicamp entre 1989 e 2008 com uso de questionário direcionado, estudando-se os dados comuns entre eles, enfatizando a história clínica, possíveis co-morbididades e fatores associados. Destes, foram entrevistados 33 doentes para obtenção e avaliação de dados.

Resultados - Os principais achados foram: 1) presença de um fator emocional desencadeante dos sintomas (cerca de 80%) e mais de 30% com tratamento psiquiátrico/psicológico declarados; 2) infecções típicas da infância muito prevalente (88% sarampo, varicela, rubéola ou outras); 3) possíveis associações a: exposição a químicos, em especial herbicidas; outras doenças do trato gastrintestinal; doenças auto-imunes; pré-disposição genética; outras alterações no sistema nervoso destacadamente as convulsões. Conclusões: A acalásia idiopática do esôfago é doença de origem provavelmente auto-imune, e associada a fatores emocionais.
INTRODUCTION

The idiopathic esophageal achalasia (IEA) is a disease of unknown etiology, characterized by esophageal aperistalsis and failure of its lower sphincter, making it difficult to swallow, causing dysphagia. The studied causes include: obstruction in the esophageal gastric junction, degeneration of Auerbach’s plexus, virus infection, congenital origin, autoimmune affection and injury by toxic agent.

The degeneration of Auerbach plexus of the esophagus determines the absence of peristalsis at the level of body of the organ as well as no aperture of its lower sphincter (achalasia) in response to swallowing. Consequently, an esophageal stasis occurs and, initially, motor incoordination and dilatation. Besides the Chagas disease (the only etiologic factor proven), its recognized that drugs may determine plexular lesion.

The congenital dilatation has been described in several dog breeds, but the proof of inherited megaesophagus in humans is thwarted by its scarcity. Among the few cases observed stand out monozygotic twins with achalasias and suggestion of transmission from father to son.

In some of the cases observed, achalasia was present in children descendant from consanguineous relations. In a more recent research, Zilberstein, et al., found nonchagasic twins developing typical symptoms of achalasia. These same authors considered idiopathic achalasia has expressions and symptoms similar to Chagas Disease.

The diagnosis of achalasia is made after excluding a possible chagasic origin of the disease, in other words, there must be seronegative results for Trypanosoma cruzi, no association what so ever to megacolon or any associated intestinal illnesses and negative epidemiological precedent for Chagas disease.

Comparatively, among idiopathic and chagasic achalasia there are similarities and differences in a few points. The destruction of the esophagus intramural nervous plexus, even though there is no consensus, is considered similar in both cases, as shown by Herbella, et al. The lower sphincter is, usually, more hypertonic in idiopathic esophageal achalasia than it is in chagasic one. On the other hand, the dilatation of the esophagus is more intense in the second.

The duration of dysphagia is a relevant differential factor between the two diseases which is more durable in chagasic patients, possibly due to the slower development. Although the incidence of cancer was quite low in all cases of both diseases, there was a higher occurrence in chagasic patients due to the duration of the dysphagia, a risk factor.

The physiopathology of idiopathic esophageal achalasia seems to indicate a previous inflammation of the myenteric nervous plexus of the esophagus developed to a posterior chronic autoimmune response in patients with larger genetic propensity which, in sequence, would raze this esophageal denervation. There are also chances of damages in the nervous system, weakening the Auerbach’s intramural plexus directly.

The inflammation that results from the autoimmune response may be due to virus infection and cases in which Guillard Barré syndrome (demyelination) previously occurred. The genetic association was reported in a few cases among which monozygotic twins, whose parents manifested consanguinity. As to the autoimmune response, besides the described sequence of chronic inflammation, there are publications that relate a higher elevation of the HLA Class II antigen in achalasia patients. These are the same molecules associated to lupus erythematosus, Sjogren syndrome and many other systemic autoimmune diseases.

Research conducted in the Hospital de Clinicas da Unicamp pointed interesting data about idiopathic achalasia. The prevalence found was of 21% (relatively high), the duration of dysphagia in years was less then half of the chagasic achalasia cases, the mean age was also lower: 37,6 (49,9 years in chagasic).

The aim of this study is to characterize the disease and propose hypothesis concerning its etiology and associated factors.

METHODS

The achievement and analysis of the clinical history and precedents from patients carrying idiopathic esophageal achalasia in this paper was approved by the Committee of Ethics in Research of the Faculty of Medical Sciences, Unicamp (179/2007).

Records from 78 operated patients suffering from idiopathic achalasia have been raised since 1989 until 2008. Among them, 33 patients were interviewed, answered a elaborated questionnaire and had their data noted, based on: age, gender, nationality, average duration of the dysphagia, serology for Chagas disease, other relevant or associated illnesses, staging of megaesophagus, applied surgery and possible complications, satisfaction after surgery, recurrence/maintenance or not of the symptoms. The classification of megaesophagus used was the one proposed by Mascarenhas (1958), considering the esophageal diameter (radiographic findings), varying from I to IV. To consider a patient with idiopathic esophageal achalasia, he must have had two negative serologies for Chagas disease and absence of positive epidemiology for Chagas.

The patient satisfaction index after the surgery(s) was graded from 1 to 5, adopting the following parameters: 1 – worse: symptoms increased; 2 – bad:
scleroderma and osteoarthritis, 5 (15%) allergic

Other recent studies point out that the amount of interstitial cells of Cajal of the esophagus - responsible for motility and called the pacemaker of the GI tract - can be greatly reduced or markedly increased by

11-15

ABCD Arq Bras Cir Dig 2010;23(1):11-15
compensatory mechanism) in patients with idiopathic achalasia, which would indicate two different mechanisms responsible for the disease. When studying the motor behavior of the esophagus, once again it is noticed the difference within the group of patients, whereas, by comparing the motor response to esophageal swallow simple and multiple swallows in 20 patients with untreated idiopathic achalasia, some patients showed increased motor activity in multiple swallows, others had inhibited motor activity and some no motor response to any swallow.

When comparing the findings between the chagasic and idiopathic achalasia, it became clear after reviewing of the literature, that the excitatory effect is larger in Chagas disease, which should justify the finding of greater pressure on lower esophageal sphincter in idiopathic achalasia. The duration of dysphagia seems to be lower in patients and idiopathic dilation of the esophagus, more pronounced in chagasic patients.

Briefly, it is likely that idiopathic achalasia is related to autoimmune disorder and that this disorder has a multifactorial etiology. There seems to be several diseases with multiple causes and unknown, many of the possibly associated factors.

The patients profile is already known: young people, usually younger than 40 years, similar prevalence between sexes, shorter duration of dysphagia accompanied by significant weight loss, incipient megaesophagus (usually grade I or II) and good surgical resolution.

The presence of previous surgery in most patients, especially abdominal and/or GI tract, could reflect attempts to correct other diseases, possibly secondary to megaesophagus, or recurrences.

The occupation does not seem to establish strong correlation, since they were very diverse. The most relevant were agricultural labor (21%) and office work (18%).

The exposure to chemicals, especially herbicides (27%), and exposition to derivatives of glyphosate resulted in complaints of epigastic pain and conjunctivitis, typical of this type of poisoning. Knowing the various negative effects that these substances can cause, it is possible to question the possibility of, in patients with predisposition to develop the idiopathic achalasia, this exposure to the herbicide act as a triggering factor. Exposure to other agents alone do not appear to be responsible for alterations.

In relation to immune system diseases, 30% of patients possibly carried some of these alterations. Idiopathic achalasia is currently regarded as an autoimmune manifestation and associated to other diseases of this same nature. It should be noted that this number may be even higher, since many of these late-onset diseases and patients operated for idiopathic achalasia are usually young.

The 18% family history of esophageal symptoms seem compatible, at least in part, to the idiopathic achalasia, and suggest genetic tendency in the family. The history of consanguinity in the families of these 18% does not seem to be of great relevance. Besides, it is difficult to obtain the percentage rate, because some people fear of having any conversation on this matter, and also the lack of information about these data within a traditional family. It should be noted that the possible genetic component, with familial occurrence, has been studied specifically in other articles.

Since achalasia is a disease that affects the intramural nervous plexus in investigation on other diseases of the nervous system may be recommended. Among all patients 15% had a history of seizures, 12% in childhood. This rate is higher than the one normally found in populations.

Smoking and alcohol consumption (39% and 42%) were not as prevalent as in cases of esophagus and stomach cancer, for example. Taking into account the multifactorial hypothesis, these types of cancer may actually be associated in some cases, for they are always potentially harmful substances. Moreover, some patients believed in the relation of these habits to the worsening of other symptoms and even opening of the disease.

The continuous use or overuse of drugs did not revealed anything particularly conclusive. The high rate of use of psychiatric medications called attention to a fact not yet appreciated nor studied in other researches: the emotional factor attached to the disease. In interviews, when asked if passed by significant emotional problems at the time prior to the onset of symptoms, 79% of patients responded positively, especially 48% reporting various family problems. Approximately 33% of patients also declared having psychiatric or psychological treatment, determining strictly that those emotional problems were serious. This number can also be underestimated because many patients do not feel comfortable to report this type of problem in one interview.

Surprisingly, for more than one time, the patient companion wanted to know details of the work with the interviewer and many were categorical in stating that the patient was going through major emotional problems before they develop the disease. In addition, there are very specific cases where the patient started reporting symptoms after the death of very close relative, parents’ divorce, separation from spouse and to long-term relationship. With this works contribution, the emotional factor precedes the disease appears to be of great relevance. Surely this still depends on further study for definitive characterization of their importance in this disease.

Finally, as found in other studies, may be association to some infections specially those from childhood. Stands the fact that many patients do not remember having these infections in childhood. It was the interviewer who gave small simple details that differentiate these diseases, in the attempt of achieving the most reliable data possible. Thus, 88%
of patients said they had suffered from at least one of the following diseases: varicella (70%), mumps (58%) and measles (52%). Among them, three patients reported having rubella, and one of them, reported the opening of dysphagia after the disease. Although not fully reliable data can be stimulating new serological studies or even DNA, evaluating whether patients with idiopathic achalasia are more frequently affected even some of these diseases than by some mechanism not yet elucidated.

CONCLUSIONS

The idiopathic esophageal achalasia is probably an autoimmune disease, which seems to be highly related to emotional problems.

AGRADECIMENTOS

Ao Programa Institucional de Bolsas de Iniciação Científica – PIBIC, Bolsa SAE/Unicamp.

REFERENCES