Gastrointestinal involvement in Chagas disease

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RESUMEN

La acalasia y el megacolon son las segundas manifestaciones más comunes de la enfermedad de Chagas (ECh), crónica en áreas endémicas de América Central y del Sur. Luego de veinte o más años desde la primoinfección, aproximadamente un tercio de las personas infectadas desarrollan alteraciones cardíacas y/o gastrointestinales típicas de la etapa crónica. Esta fase se caracteriza por daño neuronal mientérico (intrínseco) y autonómico (extrínseco). En pacientes con megaesófago y megacolon puede detectarse una densidad disminuida de células entéricas gliales, así como una pérdida de células intersticiales de Cajal. Estas lesiones afectan los complejos mecanismos neuronales, moleculares y celulares que modulan la actividad motora y otras funciones específicas del tracto digestivo. Evidencias sobre la presencia de anticuerpos séricos con capacidad para reconocer epitopes similares tanto en antígenos de Trypanosoma cruzi como del huésped sugirieron que el mimetismo molecular podría desempeñar un papel sustancial en la fisiopatología de la enfermedad de Chagas crónica. En particular, se ha encontrado una alta prevalencia de anticuerpos circulantes contra receptores muscarínicos de acetilcolina M2 (RM2) en pacientes chagásicos con acalasia y megacolon. Estos anticuerpos se unen y activan a los RM2, exhibiendo una actividad semejante a la de los agonistas muscarínicos. Los anticuerpos anti-RM2 pueden incrementar la contracción tónica tanto en el esófago distal como en el colon distal por estimulación directa o bien contrarrestar los efectos relajantes de drogas que incrementan la acumulación de AMPc (p. ej., agonistas betaadrenérgicos). Los efectos bioquímicos y funcionales de estos anticuerpos en el músculo liso esofágico y colónico podrían desempeñar un rol importante en la fisiopatología de la acalasia y el megacolon secundarios a ECh. (NeuroGastroLatam Rev. 2017;1:168-179)

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ABSTRACT

Achalasia and megacolon are the second most common manifestations of chronic Chagas disease (ChD) in endemic areas of Central and South America. Twenty or even more years after the initial infection, approximately one-third of infected people develop cardiac and/or gastrointestinal abnormalities as typical chronic damages of ChD. The chronic phase of ChD is characterized by the damage of myenteric (intrinsic) and autonomic (extrinsic) neurons. A decreased density of enteric glial cells can be detected in patients with mega-esophagus and megacolon and a loss of interstitial cells of Cajal. These lesions affect the complex mechanisms of neural, molecular, and cellular interactions that modulate the motor activity and other specific functions of the alimentary tract. Evidence for serum antibodies with the ability to recognize similar epitopes in both Trypanosoma cruzi and host antigens suggested that molecular mimicry could play a substantial role in the pathophysiology of chronic ChD. In fact, a high prevalence of circulating antibodies against M2 acetylcholine muscarinic receptor (M2R) in ChD patients with achalasia and megacolon has been found. These antibodies bind to and activate M2R, exhibiting agonist-like activity. Anti-M2R antibodies can both enhance tonic contraction in lower esophagus and distal colon by direct stimulation and also by counteracting the relaxant effect of drugs that increase cAMP accumulation (i.e. beta-adrenergic agonists). The biochemical and functional effects of these antibodies on esophageal and colon smooth muscle could play an important role in the pathophysiology of achalasia and megacolon secondary to ChD.

Key words: Chagas. Achalasia. Megacolon. Muscarinic antibodies.

CHAGAS DISEASE (ChD): GENERAL ASPECTS

The American trypanosomiasis ChD still represents a major health problem, particularly in poor and rural populations of Central and South America. ChD has three well-recognized clinical successive phases: (1) the acute infection, (2) an “indetermined” (symptomatically silent) phase, and (3) a chronic phase, typically 20 years or more after the initial infection, during which clinical manifestations appear.

As regard to cardiac abnormalities, their signs and symptoms are the most common expressions of chronic ChD: palpitations, dizziness, chest pain, and even syncope that reflect different arrhythmias, including ventricular tachycardia with sudden death, which is not at all unusual in an otherwise healthy young adult. Signs and symptoms related to arrhythmias could be present for a long period before cardiomegaly or congestive heart failure appears.

The primary aim of this article is to review the gastrointestinal involvement of chronic ChD, focusing on the most recent molecular advances to better understand the pathophysiological mechanisms of chagasic digestive damage.
Achalasia and megacolon are the second most common manifestations of chronic ChD. The gastrointestinal involvement results from denervation of the hollow viscera. Usually, esophagus and colon are affected the most; however, the duodenum and the small bowel could also be dilated. Moreover, even in patients with asymptomatic chagasic involvement of the intestine, a significant reduction in submucosal and myenteric ganglion neurons was demonstrated3.

Cardinal symptoms include gastric dilatation (gastroparesis), nausea, vomiting, and upper abdominal pain. More frequently, with duodenal and small bowel dilatations: bloating, diarrhea and/or constipation may be part of the picture; in the presence of stasis, it is possible to find signs of bacterial overgrowth4.

**CHRONIC CHAGASIC ESOPHAGEAL INVOLVEMENT**

Except for the exposure to Trypanosoma cruzi, the clinical history of patients with chagasic achalasia is indistinguishable from that of idiopathic achalasia. The symptoms are dysphagia, feeling, of fullness after eating or drinking small amounts, chest pain, and regurgitation. Weight loss and aspiration with secondary pneumonia are common findings in advanced cases. Chagasic achalasia, in early stages, shows minimal or no esophageal dilatation in contrast barium studies (Fig. 1). Various degrees of esophageal dilatation and retention of barium because of a delayed esophageal emptying could be seen in advanced stages of chagasic achalasia (megaesophagus).

Upper video endoscopy should be regularly performed to rule out obstructive - proliferating endoluminal diseases: malignancies and peptic and non-peptic (inflammatory and/or infectious) stenosis. During the chronic phase, the diagnosis of ChD is based on a history of exposure to the insect vector and a positive serologic test for specific antibodies against T. cruzi, generally an enzyme-linked immunosorbent assay (ELISA) and an indirect immunofluorescence test1.

**ESOPHAGEAL MANOMETRY: DEFINITIONS AND CONTRIBUTIONS**

Achalasia is defined as an esophageal motor disorder characterized by an impaired lower esophageal sphincter (LES) relaxation and an aperistaltic smooth muscle esophageal segment. Conceptually, identical findings have been described in patients with chagasic achalasia5. Currently, high-resolution manometry (HRM) is considered the best way of analyzing and quantifying LES relaxation and the peristaltic activity of the esophageal body, which are the crucial functions to assess to establish the diagnosis of achalasia6.
LES relaxation resting pressure is measured by means of the four second-integrated relaxation pressure (4 - IRP), which can be defined as the median LES pressure related to the gastric pressure during the 4 s of lower LES pressure within a timeframe of 10 s starting from the moment of upper esophageal sphincter relaxation. In fact, utilizing IRP, achalasia is defined more precisely than with conventional manometry. Moreover, with the criteria coming from the use of HRM and the application of the Chicago Classification, three distinct types of achalasia were described (Fig. 2A-D). The three types of achalasia are characterized by incomplete LES relaxation; they differ in pressurization patterns and the presence of premature high amplitude contractions in the type III (or spastic achalasia). The type I, or classic achalasia, has poor or absent pressurization, and the type II shows panesophageal pressurization (or bolus compression). Following this sub classification of achalasia, different studies demonstrate distinct responses after treatment. All evaluations uniformly show that the best treatment response was in achalasia type II patients and the worst therapeutic response was obtained among type III achalasia patients; this observation includes the different modalities of achalasia treatment.

Series of chagasic achalasic patients are currently being investigated, presumably, under the Chicago classification criteria, which will be of the salient clinical value to know the distribution of subcategorization of chagasic

**Figure 2.** (A) Classic achalasia or type I, non-relaxing lower esophageal sphincter (LES), aperistaltic esophageal body with an intraluminal pressure <30 mmHg. High-resolution esophageal manometry (HREM). (B) Achalasia type II, non-relaxing LES, aperistaltic esophageal body with panpressurization or compression with a 30 mmHg isobara (black contour) (HREM). (C) Achalasia type III, non-relaxing LES, aperistaltic esophageal body with high-amplitude premature contractions (spastic achalasia) and “shortening” due to a longitudinal muscle layer contraction (HREM). (D) The same achalasia type III with simultaneous impedance, the purple color indicates fluid retention in the esophageal lumen and in the gastric fundus (high-resolution esophageal/impedance manometry).
achalasia and to uniformly evaluate their therapeutic responses.

**CHRONIC CHAGASIC INTESTINAL INVOLVEMENT**

Chronic chagasic patients with megacolon suffer from infrequent bowel movements (constipation), usually associated with bloating, distention, and colicky abdominal pain. Hypoactive or “distant” bowel sounds are typical physical findings. Constipation could be severe, not infrequently, these patients refer several weeks between every bowel movement. Volvulus, obstruction, and perforation can occur as dramatic complications in chagasic patients with universal or segmental megacolon. An abdominal XR film, a barium enema with air contrast, and a computed tomography abdominal scan, characteristically, show dilated colonic loops with poor or no peristalsis and megarectum (Fig. 3). Video colonoscopy should be performed to rule out malignant diseases and other colonic and extra-colonic causes of gastrointestinal obstructions.

**FROM CLASSICAL AND RECENT MORPHOLOGIC STUDIES TO MANOMETRIC EVALUATIONS**

Classical and early morphologic studies relied on either resection specimens or necropsy material from patients undergoing resections for end-stage or advanced chagasic megaviscera. These dilated, atrophic, and “chronically non-peristaltic” hollow viscera iteratively showed the absence of myenteric ganglia and marked fibrosis.

The availability of tissue specimens from patients in earlier stages of chagasic megacolon became possible due to the wide adoption of laparoscopic surgery and the use of new diagnostic tests. The enteric nervous system (ENS) is a vast neural network along the whole alimentary tract, biliary tract, and pancreas. Based on histochemical and electrophysiological properties, the 80–100 million enteric neurons can be classified into functionally distinct subpopulations: intrinsic primary afferent neurons, interneurons, motor neurons, and secretomotor and vasomotor neurons. The ENS has the unique ability to control these fundamental gastrointestinal functions: absorption/secretion, vascular tone, and motor activity; given these important functions of the ENS, its damage results in a variety of digestive disorders.

Modern immunohistochemistry techniques have shown marked abnormalities of the ENS in patients with chagasic megacolon: (a) a decreased number of enteric neurons and enteric glial cells, (b) a mild lymphocytic infiltration of the enteric plexuses (“plexitis”), and (c) a
loss (but not absence) of interstitial cells of Cajal subtypes, although an increase of the intramuscular subtype has also been found, suggesting a possible compensative mechanism. These findings could be related to the abnormalities that have been found in chagasic patients with lower basal motility index and wave frequency of the sigmoid colon and rectum compared with normal subjects. In chronic chagasic patients, the propagation of the migrating motor complex is slower and its duration longer than in control individuals, suggesting that neuropathic damage results in alterations of the interdigestive motor cycle controlled by the ENS. As occurs with idiopathic megacolon, colonic manometry could facilitate the identification of chagasic colonic inertia characterized by a poor or absent contractile response to feeding and after a pharmacologic provocative test employing 5 mg of intracolonic bisacodyl (Fig. 4).

A lack of relaxation of the internal anal sphincter and an absence of the rectoanal inhibitory reflex (RAIR) were demonstrated in chronic chagasic patients with megacolon; this finding could be explained by the presence of aganglionosis. However, in some patients, the absence of the RAIR could be the result of megarectum rather than the expression of a true aganglionosis.

**ROLE OF THE IMMUNE SYSTEM IN THE PATHOPHYSIOLOGY OF ACHALASIA AND MEGACOLON SECONDARY TO ChD**

Over the past 20 years, several findings suggested the involvement of the immune system in the pathophysiology of achalasia and megacolon secondary to *T. cruzi* infection. The presence of inflammatory cell infiltrate in the myenteric plexus of the gastrointestinal tract as well as the detection of circulating antibodies against neural and/or smooth muscle structures in the absence of evident infection has been associated with immune activation and inflammation, both in ChD patients and *T. cruzi*-infected rodents. The inflammatory process is characterized by myositis and ganglionitis, which appear to be more frequent in organs with mega than those without enlargement and in esophagus compared to colon. Qualitatively, the inflammatory infiltration in muscular and myenteric plexus from esophagus and colon is similar, consisting predominantly of T lymphocytes (CD3+), few macrophages (CD68+), and rare B lymphocytes (CD20+) regardless of the presence of organ enlargement, although other studies have found a predominance of macrophages (CD68+) in the myenteric plexus region from patients with megaesophagus or a predominance of B lymphocytes in the submucous plexus from patients with megacolon. Quantitatively, megaorgans appear to exhibit higher numbers of inflammatory cells than those with normal dimensions.

As described above, *T. cruzi* promotes a strong immune response not only against its own antigens but also against host self-structures. For example, T cells crossreacting with *T. cruzi* antigens and nervous tissue can cause peripheral nervous tissue damage when passively transferred to naive animals. Selected anti-*T. cruzi* monoclonal antibodies have been found to recognize host antigens, and cross-reactivity of circulating antibodies from ChD patients between *T. cruzi* antigens
and host proteins muscular or nervous tissue has also been described\textsuperscript{25}.

The finding of both immunocompetent cells and circulating antibodies with the ability to recognize similar epitopes in both \textit{T. cruzi} and host antigens suggested that molecular mimicry between parasites and host antigens could play a substantial role in the pathophysiology of chronic ChD. However, it has
been difficult to demonstrate a correlation between molecular mimicry and clinical findings in most cases. We will next review the previous studies on “circulating anti-M2 muscarinic acetylcholine receptor antibodies in ChD patients” and discuss their potential physiopathological significance with respect to esophageal and colonic alterations. In particular, we will focus on our own contributions to the field.

**ANTI-M2 MUSCARINIC ACETYLCHOLINE RECEPTOR ANTIBODIES IN ChD**

During the early 90s, a series of studies were conducted to find a link between heart disease and the humoral immune response in ChD patients. The circulating IgG fraction from *T. cruzi*-infected mice was found to elicit a negative inotropic effect in isolated mouse atria preparations and trigger biochemical signals classically coupled to M2 muscarinic acetylcholine receptor (M2R) activation in atrium strips from non-infected mice, such as a decrease in cAMP accumulation. This antibody fraction was also able to inhibit the binding of a muscarinic radioligand to murine heart membranes in a non-competitive manner, suggesting a direct interaction between the IgG antibodies and the M2R. Years later, the IgG fraction of chronic ChD patients was found to mimic the pharmacologic behavior of that from *T. cruzi*-infected mouse previously reported. Moreover, the human IgG fraction from ChD patients not only decreased atrial contractility but also inhibited carbachol- and pilocarpine-mediated negative inotropic effects in a non-competitive manner. For this reason, this antibody fraction was thought to mimic the effect of muscarinic partial agonists. In addition, ChD IgG was found to inhibit L-type Ca2+ currents on binding to the M2 receptor and promote short-term regulation of the M2R by inducing desensitization and sequestration of M2R stably expressed in CHO cells.

The direct interaction of anti-M2R antibodies from ChD patients with their target receptor was demonstrated by immunoprecipitation, immunoblotting and a combination of both procedures. Moreover, an epitope within the second extracellular loop (II ECL) of the M2R was identified as the main immunogenic region interacting with the anti-M2R antibody IgG fraction. Next, a synthetic peptide with an amino acid sequence within the II ECL of the human M2R was used as an immobilized antigen to monitor the immune reactivity of circulating anti-M2R antibodies by ELISA or affinity purify monospecific anti-M2 mACh antibodies from the serum IgG fraction.

The monospecific anti-II ECL antibody fraction was found to reproduce all the pharmacological effects of the whole IgG fraction. In addition, this antibody fraction was able to induce basal bradycardia on murine cardiomyocytes and rat isolated atria, confirming the muscarinic response on functional myocardium.

The presence of circulating anti-M2R antibodies was found to be strongly associated with the evidence for cardiovascular dysautonomia in ChD patients with or without ECG alterations. Furthermore, Ribeiro et al. found that the levels of circulating anti-M2R antibodies negatively correlated with HRV index.
power, suggesting an inhibitory effect of anti-M2R antibodies on vagal function\(^37\). Other authors reported a high prevalence of anti-M2R antibodies in ChD patients with sinus node dysfunction\(^38\).

**ANTI-M2R ANTIBODIES IN CHRONIC ChD PATIENTS WITH ACHALASIA AND MEGACOLON**

Achalasia and megacolon are neurodegenerative disorders primarily not only affecting the myenteric plexus of the ENS but also vagal trunks and dorsal vagal nucleus. Actually, several studies, although not all, have provided evidence that patients with primary achalasia, whose symptoms are fairly similar as compared with those observed in the chagasic form, show functional autonomic nervous system disturbances (vagal impairment\(^39-42\)). Besides degeneration and decreased numbers of intrinsic myenteric neurons, reduced numbers of interstitial cells of Cajal\(^43,44\) and glial cells\(^22,45\) hypertrophic smooth cells, and increased fibrosis have also been reported, suggesting a complex physiopathology of chagasic esophageal and colonic disorders.

Most studies so far have focused on the integrity of pre- and post-ganglionic innervation, without considering the mechanisms of synaptic transmission at the level of effector tissues. The M2R is widely expressed in gastrointestinal smooth muscle fibers and plays an important role in smooth muscle contractility. Agonist activation of M2R can contribute to smooth muscle contractility either by promoting a direct contractile effect together with the M3 subtype\(^46-49\) or by counteracting the relaxant effect of ligands that increase cAMP (i.e. noradrenaline)\(^50\).

The previous studies had shown that (a) an anti-\textit{T. cruzi} monoclonal antibody that cross-reacts with host antigens, particularly, in esophageal and colonic smooth muscle binds to and activates M2R \textit{in vitro}\(^51-53\); (b) anti-M2R antibodies cross-react with \textit{T. cruzi}’s ribosomal P0 protein, suggesting that these antibodies could be generated as a result of an immune response to a \textit{T. cruzi} ribosomal protein through a molecular mimicry mechanism\(^34\); and (c) circulating anti-M2R antibodies with agonist-like activity are associated with parasympathetic dysautonomia in chronic ChD patients\(^27,35,37\). On the basis of these results, it was hypothesized that anti-M2R antibodies could play a physiopathological role in the gastrointestinal motility disorders in chronic ChD patients. We will next summarize the main findings of our previous reports on anti-M2 circulating antibodies in ChD patients with achalasia a megacolon\(^54,55\).

**ANTI-M2R ANTIBODIES IN CHRONIC ChD PATIENTS WITH ACHALASIA AND MEGACOLON**

In our previous studies\(^54,55\), a 24-mer peptide with an amino acid sequence corresponding to the II EC of the human M2R (pM2) was used as an immobilized antigen to screen for circulating anti-M2R antibodies in patients with/without achalasia or megacolon, either primary or secondary to ChD. These antibodies were found highly prevalent in ChD patients with achalasia (ChD+A+, 84%, \(n = 19\)) or megacolon (ChD+M+, 87%, \(n = 15\)), compared with ChD patients without gastrointestinal damage.
(ChD+A−/ChD+M−, 22%, \( n = 31 \)), non-chagasic patients with idiopathic achalasia (ChD-A+, 28%, \( n = 25 \)) or megacolon (ChD-M+, 0%, \( n = 3 \)), or healthy control patients (ChD-A−/ChD-M−, 0%, \( n = 40 \)).

It was not surprising that a few ChD+A- or ChD+M- patients showed significant titers of anti-M2R antibodies because these antibodies are frequently found in asymptomatic ChD individuals with dysautonomia\(^2\)\(^7\),\(^3\)\(^5\),\(^3\)\(^6\). Besides, it is tempting to speculate that at least some ChD patients with no motility disorders that were currently seropositive for anti-M2R antibodies might develop achalasia and/or megacolon in the future. Antibody levels were also higher in ChD+A+ and ChD+M+ patients than in the other groups. The remarkable differences in immune reactivity and frequency of anti-M2R antibodies between chagasic and idiopathic achalasia/megacolon patients suggest that these antibodies are characteristic of patients with motility disorders secondary to ChD and do not appear to play a major role in individuals with idiopathic motility disorders.

The serum IgG fractions from ChD+A+ and ChD+M+ patients elicited pharmacological effects associated with M2R activation on rat smooth muscle strips. Actually, these antibodies enhanced tonic contractions and promoted inhibition of both basal and isoproterenol-stimulated cAMP accumulation in lower esophageal and distal colonic preparations. Both the tonic contractile effect and the inhibition of cAMP accumulation were inhibited by the muscarinic peptide pM2, a selective M2 muscarinic antagonist AF-DX 116, and the Gi protein inhibitor pertussis toxin, demonstrating the M2 subtype specificity, the involvement of a Gi protein-mediated effect, and the participation of the M2R’s II ECL in the effects of anti-M2R antibodies, respectively. The monospecific anti-pM2 antibody fraction from ChD+A+ and ChD+M+ patients mimicked the contractile and biochemical effects of the crude IgG preparations, confirming the epitope-specificity of IgG antibodies.

The high frequency of circulating anti-M2R antibodies in patients with either achalasia or megacolon suggests an association between specific serum antibodies and motility disorders in ChD. If so, how can we explain the small fraction of ChD+A+ or ChD+M+ with undetectable circulating anti-M2R antibodies? Actually, a previous study showed a frequency of 100% for anti-M2R antibodies in a population of 30 ChD+M+ patients, by measuring the chronotropic effect of patient’s IgG fraction on cultured rat cardiomyocytes (\( n = 30 \))^56. The screening procedure used by these authors has been proved more sensitive than conventional ELISA against an immobilized peptide presumably because antibodies interact with a native receptor and promote a physiological amplified response. Therefore, the antibody frequency detected in our studies should probably increase if the sensitivity of our detection procedure was enhanced.

As regards the potential association between serum anti-M2R antibodies and the pathophysiology of achalasia, these findings do not explain the degenerative lesions and the loss of myenteric neurons, which are the main alterations in this disorder. However, the agonist-like muscarinic activity could contribute to LES hypertension by enhancing LES tonic contractions through M2R activation. Hence, muscarinic effects of anti-M2R antibodies.
could counteract physiological β-adrenergic LES relaxation by promoting Gi-mediated inhibition of adenylate cyclase.

Unlike idiopathic megacolon, where the excitatory innervation is preserved, megacolon secondary to ChD is characterized by a loss of cholinergic excitatory neurons. A decrease in the excitatory activity is believed to be associated with lower basal motility index and wave frequency than in normal subjects. Other alterations, such as rectum-sigmoid motor incoordination and the impairment of RAIR in response to rectal distension, can also be explained by the destruction of the excitatory motor innervation.

The contribution of circulating anti-M2R antibodies to the physiopathology of megacolon secondary to ChD has been discussed. The potential inhibition of ACh-mediated contractile response through M2R in the colonic smooth muscle by ChD anti-M2R antibodies could be explained as follows: (a) these antibodies, acting as partial muscarinic agonists, not only could certainly display muscarinic agonist-like activity but also inhibit agonist-mediated receptor activation, and (b) the chronic exposure of M2R to specific anti-receptor antibodies could promote receptor desensitization and internalization, which would result in the attenuation of the subsequent response to Ach. Indeed, both hypotheses suggest that the presence of circulating antibodies against M2R could ultimately impair the parasympathetic activity at the level of the effector M2R in smooth muscle cells. While awaiting further studies, agonist-like activity of anti-M2R antibodies could still become a factor of concern, in that the generalized activation of M2R in colonic smooth muscle could disturb the normal synchronized pattern of peristaltic activity.

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