THE PRESENCE OF ANTIAUTONOMIC MEMBRANE RECEPTOR ANTIBODIES DO NOT CORRELATE WITH BRAIN LESIONS IN CHAGAS’ DISEASE

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Abstract – We previously demonstrated correlation between parasympathetic dysfunction and brain white matter lesions in chronic chagasic patients. Objective: To correlate serum functional circulating antibodies with beta adrenergic (Ab-β), muscarinic (Ab-M) or muscarinic and beta adrenergic (Ab-Mβ) activity, the autonomic system function and brain lesions in chronic chagasic patients. Method: In fifteen consecutive chagasic patients, the autonomic nervous system was evaluated and brain magnetic resonance imaging (MRI) was performed. The sera of all patients were tested to the presence of circulating functional antibodies. Results: Sera from 11 of 15 chronic chagasic patients had some activity (Ab-β: 7; Ab-M: 1; Ab-Mβ: 3); however, there was no significant correlation between the presence of antibodies and the autonomic system function or the presence of hyperintensities in MRI. Conclusion: The mechanism involved in the genesis of hyperintense lesions seen in brain MRI of chronic chagasic patients is still unresolved, although apparently related to parasympathetic dysfunction.

KEY WORDS: Chagas’ disease, American trypanosomiasis, autonomic dysfunction, antibodies, magnetic resonance imaging, hyperintense lesions.

A presença de anticorpos anti-autonômicos não se correlaciona a lesões cerebrais na doença de Chagas

Resumo – A correlação entre disfunção parassimpática e lesões de substância branca cerebral em pacientes chagásicos já foi previamente demonstrada. Objetivo: Correlacionar a presença de anticorpos circulantes funcionais com atividade beta-adrenérgica (Ab-β), muscarínica (Ab-M) ou muscarínica e beta adrenérgica (Ab-Mβ), a presença de disautonomia e lesões de substância branca cerebral em pacientes chagásicos crônicos. Método: Em quinze pacientes chagásicos consecutivos, foram realizados a avaliação do sistema nervoso autônomo e ressonância magnética (RM) do crânio. O soro dos pacientes foi testado para a presença de anticorpos circulantes funcionais. Resultados: O soro de 11 dos 15 pacientes chagásicos apresentou alguma atividade (Ab-β: 7; Ab-M: 1; Ab-Mβ: 3); porém não houve correlação significativa entre a presença de anticorpos circulantes e disautonomia ou de hiperintensidades à RM. Conclusão: O mecanismo envolvido na gênese das lesões hiperintensas à RM do crânio dos pacientes chagásicos crônicos não está esclarecido ainda, apesar de aparentemente relacionada à disfunção parassimpática.

PALAVRAS-CHAVE: doença de Chagas, tripanossomíase americana, disfunção autonômica, anticorpos, ressonância magnética, lesões hiperintensas.

Chagas’ disease (American trypanosomiasis) is endemic in Latin America and is still an important public health problem in Brazil. It has been associated with neurological alterations ever since it was first described¹. In the chronic phase, cerebrovascular disease is a recognized cardioem-
tonomic dysfunction might be responsible for cerebrovascular disease as it may induce endothelial dysfunction leading to vasoconstriction, cerebral vascular spasms and, ultimately, a clinically defined stroke. Considering that the pathogenesis of the autonomic nervous system dysfunction has an autoimmune basis, it would be fair to postulate that autonomic dysfunction such as observed even in the early phase of Chagas’ disease might be consequent to autoreactive antibodies. These antiautonomic membrane receptor antibodies are able to activate, either, beta adrenergic-(Ab-β), muscarinic (Ab-M) or both, muscarinic and beta adrenergic (Ab-Mβ) autonomic receptors. In addition, Ribeiro et al. has shown strong correlation between the presence of Ab-M and abnormal vagal modulation in chronic chagasic patients.

White matter hyperintense lesions were previously described by the authors in 53% of brain magnetic resonance imaging (MRI) of chronic chagasic patients, even in early phase of the disease (without cardiomyopathy). These lesions significantly correlated to parasympathetic dysfunction, evaluated by sinus arrhythmia test. In contrast, white matter hyperintense lesions are described in up to 13% of general population. Other authors also found white matter lesions in brain MRI of early phase Chagas’ disease patients, associated with unspecific electroencephalographic disorders. Aging and arterial hypertension have been related to these lesions, but correlation between them and strokes are still disputed.

The objective of this study was to test the correlation between the presence of functional active circulating antibodies (Ab-β, Ab-M and Ab-Mβ), autonomic system dysfunction and white matter lesions seen in brain MRI of chronic chagasic patients.

METHOD

This is a prospective study, carried out from 2006 to 2008. The study population comprises patients evaluated at the Chagas’ disease outpatient clinic from the University Hospital of the Federal University of Rio de Janeiro, Brazil. Consecutive patients with chronic Chagas’ disease in early phase were considered for inclusion. All patients were in phase IA in the Modified Los Angeles classification, which means no cardiomyopathy or cardiac arrhythmias. The local ethics committee (Clementino Fraga Filho University Hospital of the Federal University of Rio de Janeiro) approved the protocol and written informed consent was obtained from each patient.

The diagnosis of Chagas’ disease required at least two positive standard serologic tests for antibodies against Trypanosoma cruzi (indirect immunofluorescence, indirect hemagglutination, and/or enzyme-linked immunosorbent assay).

Exclusion criteria were: age over 70 years; ischemic heart disease; association of any other heart disease; previous stroke, arterial hypertension, diabetes mellitus, renal failure; chronic obstructive pulmonary disease; smoking, alcoholism and any other important systemic disease. Patients who have received specific treatment for chagasic infection were also excluded.

Beta-blockers (as carvedilol) and calcium channel blockers were transiently discontinued (five half-lives) before blood samples were drawn.

In order to evaluate autonomic system function, cardiac frequency variability was analyzed on resting electrocardiogram (ECG). The signals were collected and stored in a personal computer (processing software Biopotentials Capture System, which allows the examiner to eliminate QRS complexes with artifacts), to measure the cardiac frequency variability in the time and frequency domains: after a 20-minute resting period in supine position, the patients were submitted to the respiratory sinus arrhythmia maneuver, with a controlled respiratory rate of 12 incursions per minute (along 2 minutes), to promote a vagal response. Variables evaluated in the time domain included the proportion of consecutive cardiac beats with a difference of 50 milliseconds or more between the consecutive beats (pNN50); and the mean square root of the sum of squares of the differences between consecutive cycles (RMSSD). Both variables give estimates of cardiac parasympathetic function. Frequency domain (spectral analysis) was evaluated by the ratio between low frequency (LF components of the cardiac cycle (representing the sympathetic component) and high frequency (HF) components (parasympathetic component): the LF/HF represents the relationship between sympathetic and parasympathetic balance. LF and HF were measured in normalized units, which represent the relative value of each power component in proportion to the total power. As there are no published parameters of normality in Chagas’ disease, cardiac frequency variability was analyzed as a continuous variable, instead of using “normal” and “abnormal” dichotomization.

MRI (Escient Prestige, 2.0 Tesla) was performed using T1, T2 and FLAIR sequences in all subjects. For this paper we considered only the presence or absence of white matter hyperintense lesions, evaluated by an expert neuroradiologist.

Isolated heart and sera characterization

The method for ECG recording from isolated rabbit hearts has been described in detail. In brief, young rabbits weighing 1.5 to 2.0 kg were killed by cervical dislocation, and their hearts were rapidly removed and cannulated through the aorta for continuous perfusion of the coronary circulation with Tyrode solution (mmol/L): NaCl, 127; KCl, 2.7; NaHCO3, 12; MgCl2, 0.5; glucose, 10; and CaCl2, 2.7; pH 7.2 at 36°C ± 0.2°C. The hearts were immersed in warmed Tyrode solution in a water-jacketed glass flask; three glass electrodes filled with 1 mol/L NaCl were positioned inside the flask to obtain optimal QRS complexes with artifacts. Two electrodes were connected to the differential input of a high gain amplifier (A-M System, USA) and the third was connected to ground. The experimental protocol consisted of control recordings for 15 to 20 min in Tyrode’s solution, a 30 min perfu-
sion with Tyrode solution containing serum (1:100 v/v) from the chagasic patients, and return to Tyrode solution (washout). The ECG was continuously recorded with the data acquisition software Axoscope (Axon Instruments, USA). Experiments were carried out only if no significant change in the ECG parameters was observed for the 15 to 20 min duration of the control recordings. The ECG analysis included RR and PR interval and presence of atrioventricular conduction block.

The sera characterization as beta-adrenergic, muscarinic, both beta and muscarinic effect or non functional (Ab-NF) was made by ECG recordings in isolated rabbit hearts. In this preparation, heart rate response to sera was assessed during spontaneous rhythm, to divide the sera of patients into four groups and to assess the functional activity of the antibodies. A given patient’s serum was characterized as Ab-β only when the serum was able to increase spontaneous heart rate by at least 10% within 30 min of initial perfusion of the isolated rabbit heart, and this effect was abolished by a beta adrenergic antagonist (1 μM atenolol). On the other hand, chronic chagasic patient’s serum was characterized as Ab-M when the serum decreased the heart rate by at least 10% within 30 min, and this effect was abolished by a muscarinic antagonist (1 μM atropine). When the sera increased together heart rate and PR interval, at least 10% within 30 min of sera perfusion, they were considered Ab-Mβ. Ab-NF was considered when the serum did not alter spontaneous heart rate.

Sera from healthy orthopedic surgery patients without Chagas infection or history of cardiac disease were also used (reference sera). Independent experiments were performed to test each serum. When necessary, to confirm the serum effect, the experiment was repeated in a new heart, in the presence of adrenergic or muscarinic antagonist, atenolol (1 μM) and atropine (1 μM) respectively.

Statistical analysis

All variables were correlated with each other using Spearman correlation coefficient and Mann-Whitney test. Statistical analysis was made with SPSS 11.0 software. Statistical significance was estimated by Student t test or one way ANOVA coupled with Newman-Keuls. Difference was considered significant at p<0.05.

RESULTS

Fifteen patients (eight men) with mean age 54.06 (±11.31) were enrolled. Eight patients (53.3%) presented white matter hyperintense lesions in brain MRI (Fig 1).

From 15 chronic chagasic patients sera tested in isolated rabbit hearts, seven showed adrenergic activity, as shown in Figure 2A. These sera have significantly and reversibly shorted the RR interval from 1257±38.8 ms in control condition to 931±23.5 ms under sera perfusion and returned to 1228±41.2 ms after washout (One-way analysis of variance p<0.0001control and washout vs Ab-β). The adrenergic sera activity, in other experiment, was
Chagas’ disease: antiautonomic antibodies
Py et al.

prevented by the perfusion of the beta-1 antagonist atenolol (Fig 2B). One serum was characterized as muscarinic (Fig 3). Three sera had both effect, beta-adrenergic and muscarinic, thus, significantly increased PR interval (control=150.2±3.5 ms, PR Ab-Mβ: 247.1±6.1 ms and washout: 155.6±3.2 ms; One-way analysis of variance p<0.0001, control and washout vs Ab-Mβ) and shortened RR intervals from 1423±43.5 ms in control to 1139±30.5 ms in Ab-Mβ and 1506±49.1 ms after washout (One-way analysis of variance p<0.0001 control and washout vs Ab-Mβ; Fig 4A). The effect of Ab-Mβ on PR interval was completely neutralized when atropine (1 μM) was present in the solution, and, as expected, the heart rate was higher than control condition, as shown in Figure 4B. To confirm the presence of the muscarinic activity in this serum, a new experiment was performed in the presence of atenolol (1 μM) to inhibit the adrenergic response. Thus, as show the Figure 4C, Ab-Mβ in the presence of atenolol diminished heart rate, increased the PR interval and evoked atrioventricular block. All of them are a characteristic of muscarinic effect.

Sera from normal subjects were tested by our group previously, without effect upon the parameters analyzed here. In addition, sera from four patients in the present study had no cardiac activity (Control: 866.7±38.2, Ab-NF: 853.7±35.7 ms and washout: 868.9±34; p>0.05) in isolated rabbit heart ECGs.

There was a significant correlation between the presence of hyperintense lesions in MRI and: pNN50 (p<0.01), rMSSD (p<0.01) and LF/HF (p<0.04), as published elsewhere. On the other hand, there was no significant correlation between pNN50, rMSSD or LF/HF and the presence of the antibodies Ab-β, Ab-M or Ab-Mβ (p>0.05).

Fig 3. Muscarinic activity of serum from chronic chagasic patient on isolated rabbit heart. Control condition is in upper panel; and the presence of serum (1:100 v/v) is in the middle, showing atrioventricular block, characterized by several non-conducting P waves (not followed by the QRS complex). PR and RR intervals longer than control condition, consequent to bradycardia induced by muscarinic activity. All of these effects were reversible in the washout condition, lower panel.

Fig 4. Muscarinic and beta adrenergic effects of serum from chronic chagasic patient on isolated rabbit heart. [A] Control and washout traces represent ECG recorded during Tyrode perfusion, and the middle trace, labeled Ab-Mβ, was obtained in the presence of a serum that induced an increase of both sinus rate and PR interval, suggesting both adrenergic and muscarinic effect, respectively. [B] The experiment was made in a new heart in the presence of muscarinic antagonist (atropine 1 μM). In the serum condition, more RR intervals appeared compared to control or washout condition, showing adrenergic effect. [C] Atenolol (1 μM) was added during this experiment. Atrioventricular block occurred, characterized by several non-conducting P waves (not followed by the QRS complex). PR and RR intervals are longer than control condition, consequent to the bradycardia induced by muscarinic activity present in the serum. All of these effects were reversible in the washout condition, lower panel.
There was no significant correlation between the presence of hyperintense lesions in MRI and the presence of the antibodies as well.

In summary, 11 of 15 tested sera had some cardiac effect. But, when we separated the patients by the presence or absence of hyperintense lesions in MRI, we did not find any significant correlation with the characterized sera.

**DISCUSSION**

We have previously demonstrated that chronic chagasic patients, even in early stages of the disease, without any cardiac dysfunction, present signs of a parasympathetic disorder that correlate significantly with brain subcortical white matter abnormalities. This was demonstrated by an inverse and significant correlation between lower pNN50 and rMSSD values (representing reduced cardiac frequency variability) and the presence and number of hyperintense lesions in MRI; and by the direct correlation between the LF/HF ratio and the presence of hyperintensities\(^1\). It shows that parasympathetic dysfunction may enhance the probability to have hyperintense lesions in brain MRI. Additionally, while 53.3\% of our patients presented hyperintensities in MRI, only 13\% subjects present similar lesions in the general population\(^5\). In agreement with our findings, other authors also demonstrated the presence of white matter lesions in brain MRI of early phase Chagas’ disease patients, possibly associated with unspecific electroencephalographic disorders\(^6\).

Once the reason for that correlation is still unclear, we speculated that Chagas’ disease, even in early stages, can promote an imbalance between sympathetic and parasympathetic systems and may promote an intense and sustained cerebral vasoconstriction that cannot be corrected by cerebral vascular autoregulation. On the other hand, we know that chronic chagasic patients often present circulating active functional antibodies that affect isolated rabbit heart, by activation of adrenergic or muscarinic receptors respectively\(^6,7,22,23\). In fact these antibodies were positively correlated with vagal dysfunction in chronic chagasic patients\(^8\). So it is reasonable to think that the presence of circulating active functional antibodies could be related to autonomic nervous system dysfunction and brain white matter lesions. In addition, even with vagal dysfunction, our patients do not have any cardiac disorder, in agreement with a previous work of Ribeiro et al.\(^24\).

Nevertheless, the results in this paper did not confirm this hypothesis. There was no significant correlation between autonomic nervous system function, measured by sinus arrhythmia test, and the presence of antiautonomic membrane receptor antibodies. There was no significant correlation between brain white matter lesions and the presence of the sera active functional antibodies as well.

We conclude that the mechanism involved in the genesis of hyperintense lesions seen in MRI of chronic chagasic patients is still unresolved, although apparently related to parasympathetic dysfunction. More studies are needed to find answers about lesion’s physiopathology, which may lead to effective treatment and preventive measures.

**REFERENCES**


