2nd Brazilian Consensus on Chagas Disease, 2015*

João Carlos Pinto Dias[1], Alberto Novaes Ramos Jr[2], Eliane Dias Gontijo[3], Alejandro Luquetti[4], Maria Aparecida Shikanai-Yasuda[5], José Rodrigues Coura[6], Rosália Morais Torres[3], José Renan da Cunha Melo[3], Eros Antonio de Almeida[7], Wilson de Oliveira Junior[8], Antônio Carlos Silveira[9], Joffre Marcondes de Rezende[10], Fabiane Scalabrini Pinto[11], Antonio Walter Ferreira[12], Anis Rassi[13], Abílio Augusto Fragata Filho[14], Andréa Silvestre de Sousa[15], Dalmo Correia[16], Ana Maria Jansen[6], Glauca Manzan Queiroz Andrade[3], Constança Felícia De Paoli de Carvalho Britto[6], Ana Yecê das Neves Pinto[17], Anis Rassi Junior[13], Dayse Elisabeth Campos[18], Fernando Abad-Branch[1], Silvana Eloi Santos[3], Egler Chiari[19], Alejandro Marcel Hasslocher-Moreno[15], Eliane Furtado Moreira[20], Divina Seila de Oliveira Marques[21], Eliane Lages Silver[22], José Antonio Marin-Neto[23], Lúcia Maria da Cunha Galvão[19], Sergio Salles Xavier[24], Sebastião Aldo da Silva Valente[17], Noêmia Barbosa Carvalho[25], Alessandra Viana Cardoso[26], Rafaela Albuquerque e Silva[26], Veruska Maia da Costa[26], Simone Monzani Vivaldini[26], Suelene Mamede Oliveira[27], Vera da Costa Valente[17], Mayara Maia Lima[26] and Renato Vieira Alves[26]


Abstract
Chagas disease is a neglected chronic condition with a high burden of morbidity and mortality. It has considerable psychological, social, and economic impacts. The disease represents a significant public health issue in Brazil, with different regional patterns. This document presents the evidence that resulted in the Brazilian Consensus on Chagas Disease. The objective was to review and standardize strategies for diagnosis, treatment, prevention, and control of Chagas disease in the country, based on the available scientific evidence. The consensus is based on the articulation and strategic contribution of renowned Brazilian experts with knowledge and experience on various aspects of the disease. It is the result of a close collaboration between the Brazilian Society of Tropical Medicine and the Ministry of Health. It is hoped that this document will strengthen the development of integrated actions against Chagas disease in the country, focusing on epidemiology, management, comprehensive care (including families and communities), communication, information, education, and research.


INTRODUCTION
General aspects of Chagas disease epidemiology, with specific focus on Brazil
Chagas disease, an infectious condition with an acute and chronic phase, is classified as a neglected disease by the World Health Organization (WHO)\(^{(1)}\)(\(^{(2)}\). As a result of human poverty, it presents high morbidity and mortality load in endemic countries, including Brazil, with focal expression in different epidemiological contexts\(^{(1)}\)(\(^{(2)}\)(\(^{(3)}\)(\(^{(4)}\). The geographic distribution of the disease is limited primarily to the American continent due to the distribution of more than 140 species of insect vectors (Triatominae, Hemiptera, Reduviidae); hence, it is also called American trypanosomiasis\(^{57}\). Gradually, however, the disease has reached non-endemic countries; this as a result of an intense process of international migration and as the mode of disease transmission has changed\(^{(5)}\)(\(^{(6)}\)(\(^{(7)}\)(\(^{(8)}\)(\(^{(9)}\)(\(^{(10)}\)(\(^{(11)}\)(\(^{(12)}\)(\(^{(13)}\)

The World Health Organization (WHO) estimates that approximately 6-7 million people are infected worldwide, mostly in Latin America\(^{(14)}\). Recent estimates for the 21 Latin American countries, based on data from 2010, showed that 5,742,167 people were infected by Trypanosoma cruzi, of which 3,581,423 (62.4%) were living in the countries of the Southern Cone Initiative against Chagas disease, especially Argentina (1,505,235), Brazil (1,156,821), Mexico (876,458), and Bolivia (607,186)\(^{(5)}\). However, these data differ from the estimates of other research groups using other methods to define T. cruzi infection in many countries; this hinders accurate establishment of the prevalence of Chagas disease in the Americas. Nevertheless, the authors agree that the number of infected individuals is still very significant in the health and social context of the continent, requiring priority and attention from the countries\(^{(15)}\)(\(^{(16)}\).

Table 1 represents changes in epidemiological parameters specific to Latin America in the recent years\(^{(5)}\)(\(^{(17)}\).

Thus, despite advances in vector control and quality assurance of blood transfusions in many of these countries, especially from intergovernmental initiatives started in the 1990s\(^{(3)}\)(\(^{(4)}\)(\(^{(10)}\)(\(^{(11)}\)(\(^{(18)}\), Chagas disease remains a relevant public health problem in Latin America, showing different regional patterns of epidemiological expression. The challenges are magnified by the lack of access to diagnosis and systematic treatment; according to 2015 estimates >80% of individuals affected by Chagas disease globally had no such access. This supports the high impact of morbidity, mortality, and the social cost of this illness.

Uncontrolled human migration, environmental degradation, climate change, high population concentrations in urban areas, and precarious socio-economic conditions (housing, education, sanitation, and income, among others) are determinants and social conditioning factors for the transmission of T. cruzi\(^{(1)}\)(\(^{(10)}\)(\(^{(11)}\)(\(^{(19)}\)(\(^{(20)}\)(\(^{(21)}\)(\(^{(22)}\)(\(^{(23)}\). Infected populations are vulnerable to varying degrees of neglect, expressed by overlapping and increased exposure to other diseases, conditions and injuries; lower coverage for preventive interventions; a higher likelihood of illness; less access to a network of health services, to secondary and tertiary health care, and to other services; poor quality of primary care services; lesser likelihood of receiving essential treatments; and greater likelihood of developing severe forms of the disease with increased risk of progression to death\(^{(24)}\).

From this perspective, it is essential to obtain greater knowledge concerning the epidemiological scenario of Chagas disease and its transmission dynamics, involving people who are

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of deaths/year</td>
<td>&gt;45,000</td>
<td>21,000</td>
<td>12,500</td>
<td>12,000</td>
</tr>
<tr>
<td>Number of infected individuals</td>
<td>30,000,000</td>
<td>18,000,000</td>
<td>15,000,000</td>
<td>5,742,167</td>
</tr>
<tr>
<td>New cases/year – vector transmission</td>
<td>700,000</td>
<td>200,000</td>
<td>41,200</td>
<td>29,925</td>
</tr>
<tr>
<td>Total population at risk</td>
<td>100,000,000</td>
<td>40,000,000</td>
<td>28,000,000</td>
<td>70,199,360</td>
</tr>
</tbody>
</table>

Source: Adapted from the Pan American Health Organization, 2006\(^{(17)}\) (TDR/WHO, PAHO, WHO); World Health Organization, 2015\(^{(5)}\).
infected or are at risk of infection, different parasite populations, vector species, and *T. cruzi* reservoirs. This integrated knowledge is central to the search for consistent and sustainable management actions, monitoring, and coherent health and social control programs that are efficient and effective[1](10)(11)(12)(13)(14)(15).

Despite a clear need for more evidence, more than 105 years since the discovery of the disease by Carlos Ribeiro Justiniano Chagas (1909) there are still important gaps in technical, scientific, and political fields that must be overcome to effectively face this widely-ignored condition.

Brazil, with its continental dimensions, is undergoing major and rapid demographic, social, and environmental change, but serious socio-economic and regional inequalities persist. Here, diseases associated with social contexts of vulnerability and neglect still afflict a considerable part of the population[16]. Gradually, however, a consistent improvement in the overall health status and life expectancy of the Brazilian population has been observed. This can be attributed to changes in the social determinants of health, advances in the control of infectious diseases, and implementation of a more comprehensive national health system with relevant social participation[17](18)(19).

Despite these social advances, it is recognized that the persistence of large social inequalities, that result in individual, programmatic and social vulnerabilities, is strongly related to Chagas disease. New health problems emerge, however, as the result of an intense urbanization process and changes in the social and environmental field; others tend to persist and coexist[20], requiring new healthcare response networks[21].

These demand specific actions, adjusted to new realities, focused on an integrated approach to the different components involved in the natural history of the disease: humans, vectors, reservoirs, and *T. cruzi*[3](8)(9)(15). The great extent and territorial diversity, with the specific ecologic, demographic, social, and economic dynamics of the regions, implies multiple epidemiological and operational clinical scenarios[3](14)(15).

In addition to the challenges, the coming years bring opportunities for shared, more concrete action in fighting Chagas disease. In 2015 the 17 new Sustainable Development Goals (SDGs) were released; these are based on the eight Millennium Development Goals (MDGs). The agenda, which was agreed to and consolidated in the framework document *Transforming Our World: The 2030 Agenda for Sustainable Development*, represents a historic and unprecedented opportunity for countries to seek new global paths[31]. In a broader sense, in order to promote prosperity and well-being for all, to protect the environment and cope with climate changes[31], these decisions will determine the global course of action to confront and eliminate poverty.

Chagas disease is part of the third objective of this document, which aims to *ensure a healthy life and promote well-being for all, at all ages*. The goal is to control the acquired immunodeficiency syndrome (AIDS) epidemic, to eradicate tuberculosis, malaria, and neglected tropical diseases and to combat hepatitis and waterborne and other transmitted diseases by 2030[31]. The progress in specific indicator composition to monitor neglected tropical diseases in the world is highlighted, from the recognition of the number of people requiring interventions against this group of diseases. In addition, Chagas disease and other neglected tropical diseases also interface with the second (zero hunger and sustainable agriculture), fifth (gender equality), and sixth (drinking water and sanitation) objectives[31].

**MORBIDITY AND MORTALITY**

*Trypanosoma cruzi* infection

In Brazil, few population-based systematic studies have been conducted. This hinders accurate estimation of the magnitude of the impact Chagas disease throughout history.

Prior to the 1950s, Chagas disease was recognized as an eminently rural endemic disease, transmitted predominantly through vectors, occurring in areas of high social vulnerability. With industrialization, the disease became modeled in a new, urban epidemiological context, increased by internal migration within the country from rural to urban areas with consequent growth of cities[32].

Between 1975 and 1980, a national serological survey in the rural population of Brazil (except São Paulo), estimated the prevalence to be 4.2%, corresponding to 6.5 million people infected by *T. cruzi*[20](32)(33). In 1996, it was estimated that the prevalence of *T. cruzi* infection in Brazil was at 3.1% in 1978 and 1.3% in 1995, amounting to an estimated 1.96 million people infected[34]. Subsequently, the Pan American Health Organization (PAHO) estimated that 21.8 million people in endemic areas in Brazil were at risk, given the estimated 1.9 million (1.019% of the population) infected individuals[17].

In 2014, the first systematic review and meta-analysis to estimate the prevalence of Chagas disease in Brazil was published. It included publications from the period between 1980 and 2012, and estimated the pooled prevalence of Chagas disease at 4.2% [95% confidence interval (95% CI): 3.1–5.7], ranging from 4.4% (95% CI: 2.3–8.3) in the 1980s to 2.4% (95% CI: 1.5–3.8) after 2000[35]. In that study, the highest prevalence was found in women (4.2%; 95% CI: 2.6–6.8), individuals over 60 years old (17.7%; 95% CI: 11.4–26.5), residents in the Northeastern (5.0%; 95% CI: 3.1–8.1) and Southeastern regions (5.0%; 95% CI: 2.4–9.9), and in mixed urban/rural areas (6.4%; 95% CI: 4.2–9.4). It was estimated that 4.6 million people (95% CI: 2.9–7.2 million) would be infected by *T. cruzi* in Brazil. Emphasizing the limitation of the findings in the literature, the authors highlight the need for new studies, with a view to obtaining truer estimates[35].

From this and other studies, the most recent estimates in Brazil regarding the number of people infected by *T. cruzi* vary from 1.9 million to 4.6 million people[35](36)(37), probably closer now to 1.0–2.4% of the population.

This epidemiological scenario brings challenges in the coming decades for the country to support control actions and establish a consistent plan in the Unified Health System [Sistema Único de Saúde (SUS)] for diagnosis, treatment, and comprehensive care of millions of people[38], combined with epidemiological surveillance adjusted to this reality[15](22). It is
presumed that up to 30% of chronically infected people are likely to develop cardiac alterations and up to 10% may experience digestive, neurologic, or mixed alterations. Given this, there is a heightened need to structure a comprehensive network of timely health care for Chagas disease in the country(39).

In order to establish estimates for the number of people infected by *T. cruzi*, the publication *Projeção da População do Brasil por Sexo e Idade para o Período 2000/2060 e Projeção da População das Unidades da Federação por Sexo e Idade para o período 2000/2030* (Projection of Brazil’s Population by Sex and Age for the Period 2000/2060 and Projection of the Federative Unit’s Population by Sex and Age for the Period 2000/2030) by the Brazilian Institute of Geography and Statistics [*Instituto Brasileiro de Geografia e Estatística (IBGE)] was taken as a population-based reference. According to the IBGE the Brazilian population will continue to grow until 2042, when it will reach 228.4 million people, mostly concentrated in urban centers. From there, it will gradually decrease and it will account for approximately 218.2 million individuals in 2060(40).

From there, it will gradually decrease and it will account for 228.4 million people, mostly concentrated in urban centers. The Population and Social Indicators. Studies Management and Analysis of 2014(35) was taken as a population-based reference. According to the IBGE the Brazilian population will continue to grow until 2042, when it will reach 228.4 million people, mostly concentrated in urban centers.

**Table 2** shows the relative projection to the estimates of the number of infected people by *T. cruzi* and the number of Chagas disease cases in the chronic phase with the cardiac and digestive form of the disease in Brazil, between 2015 and 2055, according to five-year periods. In 2015, for example, taking as a basis an estimated population of 204,450,649, it is estimated that between 1,426,994 and 3,357,633 of Brazilian individuals would be infected with *T. cruzi*. Of these, 142,699 to 335,763 would potentially have the digestive form and 428,098 to 1,007,290 would have the cardiac form of the disease. Moreover, an estimated 856,197 to 2,014,580 individuals would potentially present with *T. cruzi* infection in the indeterminate form.

In addition to internal migration in Brazil, with the urbanization of Chagas disease its occurrence in migrants from endemic areas in countries not traditionally endemic has brought about an expanded discussion on the contexts of risk and vulnerability, with a challenge to develop control actions for these countries(3) (6) (7) (10) (11) (12) (13) (16) (18) (37) (41) (42). Multiple bio-ecological, socio-cultural, and political factors have been implicated in this process(39), demanding an increased need for human and social sciences research concerning the disease(43).

In 2005, it was estimated that there were a total of 501,036 Brazilians in the United States, with a prevalence of infection by *T. cruzi* of 1.02%(17), or 5,106 cases(44). In a broader sense, it is estimated that the total of people infected by *T. cruzi* in that country could range from 300 thousand to just over 1.0 million(9), with different impacts on the health system. It is worth noting that this is a non-endemic area(9)(45).

It is estimated that approximately 72,000 people infected by *T. cruzi* live in Europe(12). From analysis of the aggregated data of a systematic review and meta-analysis investigating the prevalence of Chagas disease in Latin American people living in Europe (Spain, France, Switzerland, Italy and Germany), it is estimated that the prevalence of this infection in Brazilian migrants is 0.6% (from 0.16 to 1.12%) or 4/954(13). Migrants from Bolivia had the highest prevalence of Chagas disease (18.1%), followed by Paraguayan migrants (5.5%). The prevalence among migrants from Argentina was 2.2%; there were no cases of Chagas disease detected among migrants from Uruguay, Venezuela, Panama, Guatemala, and Mexico(13).

The occurrence of other important international migration waves within Latin America, such as from Bolivia and Paraguay to Argentina and Brazil or from Colombia to Venezuela, is

<table>
<thead>
<tr>
<th>Year</th>
<th>Estimate of the Brazilian population</th>
<th>Reference</th>
<th>Reference age range</th>
<th>Estimated number of infected people</th>
<th>Estimate of cases with the digestive form</th>
<th>Estimate of cases with the cardiac form</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(age range)</td>
<td>Population</td>
<td>%</td>
<td>infection 1.02%a</td>
<td>infection 2.4%b</td>
<td>infection 1.02%a</td>
</tr>
<tr>
<td>2000</td>
<td>173,448,346</td>
<td>≥5</td>
<td>156,133,836</td>
<td>90.0</td>
<td>1,592,565</td>
<td>3,747,212</td>
</tr>
<tr>
<td>2005</td>
<td>185,150,806</td>
<td>≥10</td>
<td>150,944,641</td>
<td>81.5</td>
<td>1,539,635</td>
<td>3,622,671</td>
</tr>
<tr>
<td>2010</td>
<td>195,497,797</td>
<td>≥15</td>
<td>145,563,676</td>
<td>74.5</td>
<td>1,484,749</td>
<td>3,493,528</td>
</tr>
<tr>
<td>2015</td>
<td>204,450,649</td>
<td>≥20</td>
<td>139,901,357</td>
<td>68.4</td>
<td>1,426,994</td>
<td>3,357,633</td>
</tr>
<tr>
<td>2020</td>
<td>212,077,375</td>
<td>≥25</td>
<td>133,880,929</td>
<td>63.1</td>
<td>1,365,585</td>
<td>3,213,142</td>
</tr>
<tr>
<td>2025</td>
<td>218,35014</td>
<td>≥30</td>
<td>127,334,466</td>
<td>58.3</td>
<td>1,298,812</td>
<td>3,056,027</td>
</tr>
<tr>
<td>2030</td>
<td>223,126,917</td>
<td>≥35</td>
<td>120,096,221</td>
<td>53.8</td>
<td>1,224,981</td>
<td>2,882,309</td>
</tr>
<tr>
<td>2035</td>
<td>226,438,916</td>
<td>≥40</td>
<td>112,013,898</td>
<td>49.5</td>
<td>1,142,542</td>
<td>2,688,334</td>
</tr>
<tr>
<td>2040</td>
<td>228,153,204</td>
<td>≥45</td>
<td>102,983,115</td>
<td>45.1</td>
<td>1,050,428</td>
<td>2,471,595</td>
</tr>
<tr>
<td>2045</td>
<td>228,116,279</td>
<td>≥50</td>
<td>92,984,144</td>
<td>40.8</td>
<td>948,438</td>
<td>2,231,619</td>
</tr>
<tr>
<td>2050</td>
<td>226,347,688</td>
<td>≥55</td>
<td>82,097,220</td>
<td>36.3</td>
<td>837,392</td>
<td>1,970,333</td>
</tr>
<tr>
<td>2055</td>
<td>222,975,532</td>
<td>≥60</td>
<td>70,485,475</td>
<td>31.6</td>
<td>718,952</td>
<td>1,691,651</td>
</tr>
</tbody>
</table>

Source: IBGE – Overall population estimates and by age group. Coordination of the Population and Social Indicators. Studies Management and Analysis of Demographic Dynamics(38). a Parameters for estimates (minimum and maximum) of the prevalence of *T. cruzi* infection - PAHO, 2006(17). Martins-Melo et al., 2014(13). b Parameters for maximum estimates of the prevalence of Chagas disease in the chronic phase with cardiac form (≥30%) and digestive form (≥10%).
recognized. On the other hand, from the second half of the twentieth century, it is especially emphasized the intensification of the movement of thousands of citizens from Latin America to countries in North America (mainly the United States, but also Canada), Europe (particularly Spain, among other nations), Asia (Japan), and Oceania (Australia), increasing the number of people with Chagas disease living in non-endemic countries. Overall, this migration process is associated with the search for improved living conditions for people from endemic countries living in conditions of serious social vulnerability.

In addition to Brazilians potentially infected by T. cruzi living abroad, a considerable number of people have migrated to Brazil from other endemic countries such as Bolivia; an estimated 80,000 to 200,000 Bolivian migrants live in the State of São Paulo. This fact has prompted discussion within the country, about public health policies consistent with the need for greater attention to this migrant population, including viewing the precarious life and work conditions to which they are subjected.

In addition to the migration context, there is a clear shift in the prevalence of infection by T. cruzi to older age groups, which brings new challenges for the SUS. With the prevalence of chronic phase Chagas disease resulting largely from infection by vectors in the past, the probability of infectious and non-infectious comorbidities has expanded. These comorbidities consist mostly of chronic degenerative conditions (especially diabetes mellitus, systemic arterial hypertension, and other heart diseases), human immunodeficiency virus (HIV) co-infection with T. cruzi, and other immunosuppressive conditions that are acquired or induced. The risks are amplified by longer survival combined with migration and the urbanization that had occurred over the past five decades.

As for other infectious diseases, T. cruzi can behave as an opportunistic microorganism in individuals with immunosuppression. Although the first case of T. cruzi and HIV co-infection was reported in the 1980s, the frequency of this co-infection and of specific reactivation; the clinical and laboratory features associated with co-infection; specific, adequate treatments for T. cruzi infection in the context of co-infection with HIV; and outcomes (survival and death) of co-infected individuals, persist as knowledge gaps at a global level.

In Brazil, we estimated the prevalence of co-infection to be 1.3%. Considering that in 2014 there were approximately 734,000 people living with HIV (without AIDS) in Brazil, corresponding to a prevalence in the general population of 0.4%, there would be an estimated 9,542 cases of T. cruzi/HIV co-infection. On the other hand, since the beginning of the AIDS epidemic in Brazil to June 2014, total of 757,042 AIDS cases were registered in the Brazilian Notifiable Diseases Information System (Sistema de Informação de Agravos de Notificação) (SINAN); it is estimated that more than 9,842 of these would be co-infected. Therefore, it is estimated that there are 19,384 cases of T. cruzi/HIV-AIDS co-infection in Brazil, based on data up to June 2014, stressing that many of these cases probably died, are undiagnosed, or unrecognized as having co-infection. In Brazil, anti-T. cruzi antibodies testing is recommended for HIV-infected patients, taking into account their epidemiological risks.

It is worth noting that, for surveillance purposes, since 2004 reactivation of Chagas disease has been added to the list of AIDS-defining illnesses, and has been an important marker, from the definitive diagnosis of Chagas myocarditis and meningoencephalitis. Generally, such HIV co-infection occurs through sexual transmission in individuals previously infected with T. cruzi. A progressive decrease in the number of co-infection cases is expected in Brazil, since Chagas is gradually becoming restricted to higher age groups.

In addition to the above co-infection, generally, despite the high morbidity and mortality of Chagas disease in Brazil, data relating to the specific epidemiological surveillance of human cases of the disease does not allow estimation of its magnitude, since it is only mandatory to report cases in the acute phase to SINAN. Furthermore, it is estimated that only 10–20% of acute Chagas disease cases are actually reported.

In the period 2000–2013 (data updated in May 2014), 1,570 cases of acute Chagas disease were reported—an average of 112 cases per year. This figure was derived from records from most Brazilian states, with the exception of Mato Grosso do Sul and Federal District in the Central-West region and Paraná in the South. Most of these cases (1,430; 91.1%) were concentrated in the North, followed by the Northeast (73; 4.7%), South (28; 0.2%), Central-West (27; 1.8%) and Southeast (12; 0.8%). It is noteworthy that the State of Pará was responsible for 75% of all cases in the country and 82% of cases in the Northern region.

With regard to the municipality of residence, there were 163 reported cases of acute Chagas disease: 97 (60%) from the Northern region, 37 (23%) from the Northeast, 14 (9%) from the Central-West and, in smaller proportions, 9 (6%) and 6 (4%) from municipalities of the Southern and Southeastern regions, respectively. This notification pattern is quite distinct from that of the 1990s, but still demarcates operational contexts of epidemiological silence in some areas.

Mortality

The burden of mortality related to Chagas disease in Brazil remains high, despite the introduction of control measures. A study conducted to quantify the mortality pattern of Chagas disease in the period 1981–1998 found that out of 68,936 deaths in Brazilian individuals with a known birthplace, 32,369 (32%) deaths occurred in states other than the deceased’s birthplace, varying from 0.3% in Rio Grande do Sul to 100% in Roraima and Amapá. Most (67%) of these deaths in migrants occurred in individuals of Minas Gerais (51%) and Bahia (16%). Mortality rates of residents within the period showed a consistent mortality pattern from that of the 1990s, but still demarcates operational contexts of epidemiological silence in some areas.

The burden of mortality related to Chagas disease in Brazil remains high, despite the introduction of control measures. A study conducted to quantify the mortality pattern of Chagas disease in the period 1981–1998 found that out of 68,936 deaths in Brazilian individuals with a known birthplace, 32,369 (32%) deaths occurred in states other than the deceased’s birthplace, varying from 0.3% in Rio Grande do Sul to 100% in Roraima and Amapá. Most (67%) of these deaths in migrants occurred in individuals of Minas Gerais (51%) and Bahia (16%). Mortality rates of residents within the period showed a consistent decrease in the Southeast, South, and Central-West, but not in the Northern and Northeastern regions, where the median age at death was comparatively low.

Later, from 1999 to 2007, Chagas disease was recorded as the cause of death on 53,924 (0.6%) of approximately 9.0 million death certificates from the Brazilian Mortality Information System (Sistema de Informações sobre Mortalidade):
for 44,537 (82.6%) it was recorded as the underlying cause and for 9,387 (17.4%) as an associated cause\(^{(59)}\). Acute Chagas disease was recorded in the statement in 2.8% of deaths. The average number of deaths per year was 5,992, with an average standardized mortality rate of 3.36/100,000 inhabitants/year, verifying a gradual reduction in the specific mortality rate of 26.4%. It is emphasized the decreasing tendency of a standardized mortality rate, especially in the Central-West region (-37.9%). The proportional mortality (for numerous reasons) was 0.6%, highest in the Central-West region (2.1%) where there was a reduction of 19.5%, with a significant decreasing tendency. However, in this study, there was a considerable increase of 38.5% in the Northeast region, with an increasing tendency. Risk factors identified in the bivariate analysis were older age groups and residence in the Central-West region. In the multivariate analysis, however, age over 30 years and residence in the States of Minas Gerais and Goiás and in the Federal District were identified as risk factors\(^{(59)}\). It should be noted that more than 40% of expected deaths associated with Chagas disease in Latin America occurred in Brazil\(^{(56)}(59)\).

Another study of the same Brazilian population base examined 53,930 deaths, recorded between 1999 and 2007, where Chagas disease was mentioned as a cause of death. When analyzed as the underlying cause, Chagas disease was the fourth leading cause of death (10.8%) among all infectious diseases and parasites\(^{(56)}\). Of the 53,930 recorded deaths, 37,800 (84.9%) were related to cardiac forms and 4,208 (9.4%) to digestive forms of Chagas disease. Acute cardiac involvement was noted in 1,097 (2.5%) deaths; other organ involvement in 1,157 (2.6%); and finally, other clinical presentations of the disease (central nervous system involvement and no cardiac involvement in the acute phase) in 281 (0.6%). The associated causes of death with Chagas disease as the underlying cause included diseases of the digestive, circulatory, and respiratory systems. Direct complications of heart disease, especially conduction disturbances/arrhythmias and heart failure were identified in more than 35% of deaths, followed by shock (15%). Among the basic causes of death in which Chagas disease was mentioned as an associated cause, the most common diseases were of the circulatory and respiratory systems (59%), followed by cancer (11.6%) and digestive system diseases (10.6%). Cerebrovascular disease, ischemic heart disease, and hypertensive diseases were the main underlying causes of death in cases where Chagas disease had been identified as an associated cause. Chronic lung diseases and, in particular, chronic obstructive pulmonary disease also represented particularly important causes of death\(^{(56)}\).

In an additional study from this same database, using different spatial analytical approaches, we identified a cluster of high mortality risk for Chagas disease involving nine States of Brazil’s Central-West region: Almost all municipalities from the States of Goiás and Minas Gerais, Federal District and some municipalities from the States of São Paulo, Mato Grosso, Mato Grosso do Sul, Tocantins, Piauí and Bahia. This area was indicated by the authors as a priority for integrated planning of actions to strengthen the network of attention focused on Chagas disease\(^{(60)}\).

A study examining regional patterns and tendencies in Brazil over a longer time period (1979–2009), found that of 27,560,043 deaths analyzed, Chagas disease was the underlying cause in 172,066 deaths\(^{(57)}\). The proportional mortality of Chagas disease was 0.62% with a crude specific death rate of 3.61 deaths/100,000 inhabitants/year and a specific mortality rate adjusted for age of 5.19 deaths/100,000 inhabitants/year. Nationally, there was significant reduction over the period, but with important inter-regional differences; there was significant reduction in the Southeast, South and Central-West regions, but significant growth in the Northeast and Northern regions\(^{(57)}\).

Another historical series, from 1980 to 2008, found that the proportion of deaths attributed to Chagas disease as the underlying cause remained stable in the period\(^{(29)}\).

In the State of São Paulo from 1985 to 2006, there were 40,002 Chagas disease-related deaths. Chagas disease was an underlying cause in 34,917 (87.3%) and an associated cause in 5,085 (12.7%). The mortality rate, according to the underlying cause, declined by 56.1%, but remained stable in terms of associated cause of death. There were 44.5% more deaths in men than in women, and 83.5% of deaths occurred in people over 45 years old. The main causes associated with those who had Chagas disease as underlying cause of death were: direct complications of cardiac impairment (conduction disorders, arrhythmias and heart failure). For Chagas disease as an associated cause, the most important underlying causes were ischemic heart disease, cerebrovascular disease, and neoplasms\(^{(61)}\).

On the other hand, from the national database on mortality related to Chagas disease as the underlying cause, from 2000 to 2010, it was found that most deaths (85.9%) occurred in males older than 60 years. These deaths were mainly caused by cardiac involvement, and the mortality rate of this clinical form of the disease decreased in all regions except in the North, where there was an increase of 1.6%. The Northeastern region had the lowest reduction, while the Central-West had the largest. It is noteworthy that the mortality rate related to the digestive form increased in all regions\(^{(61)}\).

A pattern of aging and an expansion of ages at highest age ranges for the population affected by Chagas disease has been verified, not only by previous studies of mortality, but also by other population-based studies based on cases of cohorts followed through long periods of time\(^{(63)}(64)\).

Data from the Ministry of Health on deaths from acute Chagas disease indicate that in the period 2005–2013, the national average annual mortality rate over the 14 years was 2.7% (37.9/14). In 2005, a high (20%) mortality rate was recorded. This coincided with an acute Chagas outbreak by oral transmission in Santa Catarina, at which time little was known about the etiology and management of the disease by this form of transmission. In 2006, the mortality remained high (5.9%), with subsequent reduction that has remained relatively constant (average of 1.7% from 2007 to 2013), ranging from 0.5% in 2009 to 4.7% in 2011. It is worth noting that the lack of disease suspicion and the delay in establishing the diagnosis may worsen the clinical picture and the evolution of the disease,
and may affect the prognosis. For example, deaths occurred due to lack of timely intervention\(^5\), particularly in terms of the cardiovascular forms of the disease.

In terms of mortality associated with co-infection of \(T. cruzi/\)HIV, approximately 9 million deaths were studied in the period 1999–2007. Seventy-four deaths were identified as related to \(T. cruzi/\)HIV co-infection; in 57 (77%) the underlying cause was AIDS, and in 13 (17.6%) the underlying cause was Chagas disease\(^5\). Co-infection was associated with male sex (51.4%), Caucasian ethnicity (50%), age between 40 and 49 years (29.7%), and place of residence in the Southeastern region (75.7%). The average age at death was significantly lower among co-infected (47.1 years) individuals than among those without co-infection (64.1 years)\(^5\). These data probably indicate a understimation the real impact of co-infection on mortality.

**Epidemiological profile according to transmission modes**

In more recent years (SINAN, 2000–2013), with the operational limitations as previously indicated, it was found that oral transmission was the most frequent mode of transmission every year (1,081; 68.9%), followed by vectorial transmission (100; 6.4%)\(^5\). The persistence of the latter has been observed since 2006, the year that Brazil obtained certification for interrupting transmission of \(T. cruzi\) by *Triatoma infestans*. Transmission still occurs, albeit to a lesser extent, probably due to reduced visibility of these cases. Comparatively, WHO estimates, based on data from 2010, indicated that a total of 46 cases of Chagas disease per year in Brazil originated from vectorial transmission\(^5\). Together, these data reinforce the likely underestimation of cases relating to this mode of transmission, both nationally and internationally. The mode of transmission was not identified in 372 (23.7%) reported cases; 93% (306 cases) of these were from the State of Pará. In 11 (0.7%) cases other modes of transmission (transfusion or accidental) were identified, and there were 6 (0.4%) cases of vertical transmission\(^5\). Underreporting is also clear in the later mode of transmission.

This change in the epidemiological pattern of transmission of Chagas disease is the combined result of the control measures undertaken over the last four decades and major environmental, demographic, economic, and social changes, and the concentration of the population in urban areas\(^9\)\(^15\)\(^13\)\(^65\). Thus, mechanisms directly related to the enzootic cycle of infection, such as extradomiciliary vector transmission or visitation of wild vectors to households, in addition to oral transmission (also mediated by vectors, in most cases), have to have relevance as main modes of \(T. cruzi\) transmission to human populations\(^22\)\(^27\)\(^65\).

**Vector transmission**

In 1950 and 1951, the first Prophylaxis Campaign for Chagas disease was developed in Brazil, led by the then National Service for Malaria, reaching 74 municipalities on the border of the States of Minas Gerais and São Paulo with chemical vector control activities. By 1975, control activities were conducted intermittently, with varying territorial scope. The area at risk of vector transmission of Chagas disease in Brazil in the late 1970s included 18 states, with more than 2,200 municipalities, which proved the presence of domiciled triatomine. Notably, the Amazon region was excluded from this risk area\(^6\)\(^19\)\(^20\)\(^22\)\(^66\).

Since then, with the intensification of control measures, the vectorial transmission of \(T. cruzi\) in Brazil has shown marked and systematic reduction. This was achieved by the development of systemized actions of chemical control of domiciled vector populations, introduced in 1975, with a range of coverage of the endemic area in 1983\(^20\)\(^22\)\(^66\). These actions have taken place on a regular basis since then. However, their reach has progressively declined over the past two decades due to technical and political prioritization of changes, and due to political-institutional reorganization in the country\(^22\).\(^22\)

This process, in an integrated manner, culminated with the Certification of Interruption of Transmission of the main domiciled vector, *T. infestans*, granted in 2006 by Pan American Health Organization/World Health Organization (PAHO/WHO) to countries of the Southern Cone Initiative\(^18\)\(^22\)\(^39\). The control of vector-borne transmission had considerable impact in relation to congenital and transfusion transmissions\(^4\)\(^10\)\(^18\)\(^23\)\(^29\). Despite the advances, the risk of vectorial transmission of Chagas disease persists. This has been evaluated from different perspectives\(^22\), including the existence of indigenous species of triatomine with high potential for colonization; the presence of \(T. cruzi\) reservoirs and increasing proximity of human populations to these environments; and the persistence of residual foci of *T. infestans*, still existing in some municipalities of the States of Bahia and Rio Grande do Sul\(^22\)\(^23\)\(^39\)\(^67\). The impacts of the certification process achieved by Brazil in 2006 in relation to *T. infestans* generated concern about the demobilization of society in general and the imprecise way information was obtained and disclosed to society\(^68\).

Furthermore, additional studies and analyses have sought to evaluate further the operational processes involved in different strategies to control/eliminate of indigenous vector transmission, including the certification model and potential alternatives\(^69\)\(^70\)\(^71\)\(^72\). Data from the Ministry of Health on entomological surveillance (passive or active) specific to the period 2007–2011 (Epidemiological Bulletin of Chagas Disease 2015) indicate a record catch of >770,000 triatomine in local homes and peridomestic contexts. Like human cases, considering the low coverage of entomological surveillance actions, this record probably comprises less than 10% of triatomine in or around residences. Also, according to data from the Ministry of Health, among the 62 species distributed in intradomicile and peridomicile spaces in Brazil, the following stand out as species of epidemiological relevance: *Panstrongilus geniculatus, Panstrongilus lutzi*, *Panstrongilus megistus*, *Rhodnius nasutus*, *Rhodnius neglectus*, *Rhodnius robustus*, *Rhodnius pictipes*, *Triatoma infestans*, *Triatoma brasiliensis*, *Triatoma maculata*, *Triatoma pseudomaculata*, *Triatoma rubrovaria*, *Triatoma rubrofasciata*, *Triatoma sordida* and *Triatoma viticeps*.

According to the data from the Epidemiological Bulletin of the Ministry of Health on Chagas Disease, 591,360 (76.8%)
of the captured triatomine were examined for infection with *T. cruzi*; the natural infection rate was an estimated 2.7% (15,967 triatomine). The *Triatoma vitticeps* (52%), *Rhodnius robustus* (33.3%), and *Panstrongylus lutzi* (29.4%) species had the highest rates of natural infection. According to the data from the Ministry of Health, *T. vitticeps* appears more frequently in the Southeastern States (Minas Gerais and Espirito Santo), *R. robustus* in the Northern region (Tocantins, Amazonas, Acre, and Rondônia), and *P. lutzi* in the Northeastern region (Piauí, Bahia, Sergipe, Alagoas, Paraíba, Rio Grande do Norte, Ceará and Pernambuco) (55).

In 2012, the investigation of two cases of acute Chagas disease recorded by vector transmission showed the involvement of wild species with high rates of natural infection in municipalities that were considered non-endemic for Chagas disease. One case occurred in the city of Ibitirama (Espirito Santo), where infected *T. vitticeps* adults are often found by local residents. The other case occurred in Mangaratiba (Rio de Janeiro), involving *Triatoma tibiamaculata* found to be infected with *T. cruzi*. These episodes reflect a nationwide reality; that parasite transmission cycles exist in wild environments close to human dwellings, which favors transmission (55).

For the period 2007–2011, the persistence of *Triatoma infestans* foci was identified in four municipalities of the State of Bahia (Itaguaçu da Bahia, Ibipeba, Novo Horizonte, and Tremedal) and 12 municipalities of Rio Grande do Sul (Ajarucaba Alegria, Coronel Barros, Doutor Mauricio Cardoso, Giruá, Humaitá, Ijuí, Independência, Porto Mauá, Salvador das Missões, Santo Cristo, and São José do Inhacorá) (55). Integrated actions of vector control mobilizing Federal, State, and Municipal governments have advanced towards the control of this triatoma.

Currently, the risk related to the transmission of *T. cruzi* in Brazil depends mainly on: I) the persistence of residual foci of *T. infestans*, with episodes found in some states, such as Bahia and Rio Grande do Sul; II) the existence of large number of species that were confirmed as indigenous or acting as potential vectors in the country, even though in some cases resident populations have been greatly reduced; III) the emergence of new species at risk of domiciliation (*T. rubrovaria, P. lutzi*); IV) the existence of endemic transmission in the Amazon Region, with still poorly characterized mechanisms of transmission such as domiciliary vectorial without colonization, extradomiciliary vectorial, and oral; and V) the occurrence of outbreaks and micro-epidemics of orally transmitted of *T. cruzi* infection (20, 33, 65).

The major change in the epidemiological situation of the disease in the country has generated the need for a dynamic review of the control strategies adopted. From this perspective, identifying areas with greater vulnerability to the synanthropic occurrence of triatomine has been an important tool to reorient prevention, control, and surveillance actions (39, 67), and to optimize the use of available program resources (20). On the other hand, risk stratification associated with vector transmission has been proposed as an important strategy for the guidance and support of control measures in the country and has been incorporated, in different ways, by the control programs of states and municipalities (4, 20, 22, 39, 65). This stratification is based on three different scenarios or eco-epidemiological spaces of vector-borne transmission of *T. cruzi*: (22):

- **an area first considered endemic, with registered domiciliary vectorial transmission, which was submitted for an extended period to intensive chemical control operations, resulting in the almost complete elimination of the main vector (*T. infestans*), and the risk being currently limited to a more or less focal transmission, mainly by native species. This scenario includes the States of Alagoas, Bahia, Ceará, Goiás, Mato Grosso do Sul, Minas Gerais, Paraíba, Pernambuco, Paraná, Piauí, Rio Grande do Norte, Rio Grande do Sul, São Paulo and Sergipe, as well as the Federal District;**

- **an area previously considered harmless for human Chagas disease, in which the disease is now transmitted through mechanisms that before were considered extraordinary or improbable and are still poorly characterized but able to maintain endemic transmission of relatively low intensity to sustain endemic transmission (oral, extradomiciliary and domiciliary without vector colonization). This scenario includes the region covering the States of Acre, Amazonas, Amapá, Rondônia, Roraima and Pará; and**

- **a transition area between the two previous scenarios, in which both coexist. This scenario includes the States of Maranhão, Mato Grosso, and Tocantins.**

As previously discussed, the States of Santa Catarina, Rio de Janeiro, and Espirito Santo, although not covered by the proposal of the previous risk stratification, given the historical patterns observed from the entomological surveillance of the states, have registered autochthonous cases of Chagas disease (39, 55, 71, 74, 75, 76, 77).

The evaluation and systematic monitoring of this process should incorporate the possibility of colonization of the Amazon and other agricultural fronts (extensive cattle ranching and soybean cultivation, among others) by domiciliated vectors, with epidemiological importance in the areas of human migration (22, 65).

The great diversity of vector-borne transmission of *T. cruzi* inherent to the aforementioned scenarios or eco-epidemiological spaces include (22, 65): I) the remote risk of reestablishment by introduced species, such as *Triatoma infestans*; II) focal transmission by native species known as vectors, such as *Panstrongylus megistus, Triatoma brasiliensis* and, less likely, *Triatoma pseudomaculata* and *Triatoma sordida*; III) domiciliation of some species typically considered as exclusively wild, and the risk that these then act as vectors in some areas, as in case of *Triatoma rubrovaria* in the State of Rio Grande do Sul and *Panstrongylus lutzi* in the States of the Northeastern region; and IV) the possibility of extradomiciliary transmission or triatomine house visitation (basic form of outbreak generation by oral transmission, at least in rural areas), as already observed with *Rhodnius brethesi* in the region of Alto Rio Negro, in the State of Amazonas, and with *Rhodnius pictipes* in the State of Tocantins.

Given the coexistence of areas with different degrees of risk, surveillance and vector control actions should be adjusted...
and based on established risks\textsuperscript{(22)}\textsuperscript{(65)}. In addition to considering the operational capacity of municipalities, the stratification of a large traditionally endemic area is based on a set of variables that potentially influence the process of infestation and/or reinfestation, and therefore of transmission or reintroduction of vectorial transmission of Chagas disease in the house environment\textsuperscript{(22)}.

It is recognized that the process of simplification of fauna is the result of unplanned occupation of the environment, having the following potential consequences: I) lower diversity of food supply for triatomines in different habitats; II) a greater tendency of displacement of triatomines in search of food (hematophagous activity) from their original habitat, to feed on more ecologic mammals species that remained and expanded numerically; III) the populations of the parasite will also expand from these species of mammals with this environmental eclecticism, in general large reservoirs of \textit{T. cruzi}\textsuperscript{(21)}\textsuperscript{(78)}\textsuperscript{(79)}\textsuperscript{(80)}. As these animals (such as marsupials and rodents) have a high degree of synanthropy, the epidemiological risk becomes clear\textsuperscript{(79)}\textsuperscript{(80)}.

\textit{T. cruzi} is a multi-host flagellate parasite capable of infecting tens of species of wild and domestic mammals distributed in all phytogeographic regions in Brazil; it is found in the most diverse ecological niches, contributing, in each type of ecotope, to form singular modalities of natural transmission foci\textsuperscript{(21)}\textsuperscript{(23)}\textsuperscript{(27)}\textsuperscript{(39)}\textsuperscript{(78)}\textsuperscript{(79)}\textsuperscript{(80)}. The integration of the reservoirs in the routine monitoring process should be taken into account given that domestic animals are not confined and can also act as a link between wild and domestic transmission routes\textsuperscript{(80)}\textsuperscript{(81)}. As with wild mammals, the importance of domestic animals (for example, dogs and cats) as reservoirs of \textit{T. cruzi} varies in different places, and they can be used as sentinel populations of transmission of \textit{T. cruzi} in a given area\textsuperscript{(21)}\textsuperscript{(81)}. In general, they are always exposed and their infection usually precedes human infection\textsuperscript{(78)}. Therefore, the presence of domestic animals infected with \textit{T. cruzi} in a given area indicates that transmission is occurring in areas where these animals circulate, which reinforces the need to widen control actions\textsuperscript{(81)}.

The research of entomological and reservoirs must be associated with sustained environmental surveillance actions\textsuperscript{(39)}. To this end, the understanding of the processes of habitat selection by triatomines is fundamental for the construction of epidemiological evidence, in order to plan and develop local systems of epidemiological surveillance and control, thus reinforcing the role of surveillance\textsuperscript{(22)}\textsuperscript{(39)}\textsuperscript{(82)}. However, operational failures identified in the detection of vectors can generate critical classification errors and, therefore, prevent the scope of the actions, hampering the composition of scenarios and dynamics of these insects in their habitat\textsuperscript{(70)}\textsuperscript{(71)}\textsuperscript{(82)}.

Assuming that no sampling technique is perfect, some studies in Brazil have applied methods that explicitly incorporate the failures in the detection of infestation foci in the research of the ecology and the surveillance of vectors of Chagas disease in different environments, including the Amazon and Northeast region\textsuperscript{(70)}\textsuperscript{(71)}\textsuperscript{(82)}. This approach has and will increasingly enable future improvement of the estimates of eco-epidemiological indicators and may significantly strengthen the strategies for integrated monitoring and control of vectors\textsuperscript{(70)}\textsuperscript{(71)}\textsuperscript{(83)}\textsuperscript{(84)}. The failures were observed in areas with well-established triatomine infestation, where active searches by trained and motivated agents usually detect approximately 40–60% of intradomiciliary and extradomiciliary infestation outbreaks, while undetected foci are not eliminated\textsuperscript{(70)}. In addition, from the analysis of the monitoring process by means of active searches conducted by control agents, its sensitivity is estimated at approximately 20% for locations with low intensity infestation (few outbreaks, with few triatominae) and at approximately 40% in locations with more intense infestation\textsuperscript{(71)}.

In addition to this approach, the integrated use of spatial analysis techniques to extend the analytical capability, with insertion of spatial and temporal dimensions of the transmission of Chagas disease, has shown great potential, in particular by the significant development of portable electronic equipment and software for data processing and analysis\textsuperscript{(70)}\textsuperscript{(85)}\textsuperscript{(86)}\textsuperscript{(87)}.

In the past two decades, several studies have been published describing the different degrees of resistance of triatomine populations to insecticides\textsuperscript{(88)}. However, the actual factors behind the emergence of these phenotypes are unknown and the impact of this on the development of standardized strategies for vector control are unclear\textsuperscript{(90)}\textsuperscript{(99)}. This new context will require endemic countries to build networks of cooperation between laboratories for the analysis of resistance in triatomine populations in a standardized manner, integrating this activity in the process of epidemiological surveillance\textsuperscript{(88)}.

**Oral transmission**

In Brazil, vectorial transmission of Chagas disease by its principal vector, \textit{T. infestans}, is under control. However, oral transmission of \textit{T. cruzi} has become increasingly epidemiologically relevant, particularly in the Amazon Region\textsuperscript{(11)}\textsuperscript{(23)}\textsuperscript{(39)}\textsuperscript{(76)}\textsuperscript{(91)}\textsuperscript{(92)}. Transmission by the oral route is considered a primary mechanism, in particular in the wild cycle\textsuperscript{(93)}; it will occur regardless of control actions undertaken\textsuperscript{(21)}\textsuperscript{(39)}. It presents a common feature in the early enzootic cycle of this parasite: ingestion of vectors and reservoirs of infection by susceptible mammals\textsuperscript{(21)}. In the case of humans, this transmission occurs in a sporadic and circumstantial manner, through food contaminated with the parasite, mainly from triatomines or their feces, with records since the 1960s\textsuperscript{(93)}. Therefore, the analysis of this epidemiological context refers indirectly to vectors.

In Brazil, most cases of orally transmitted Chagas disease have been reported in the Amazon as outbreaks in families or multifamily contexts\textsuperscript{(11)}\textsuperscript{(23)}\textsuperscript{(91)}\textsuperscript{(92)}. Outside the Amazon region, few events have been investigated; most were related to sugar cane juice, the likely food carrier in the oral transmission of these cases\textsuperscript{(76)}\textsuperscript{(93)}. A recent review of the subject identified records of outbreaks or micro-outbreaks in Brazil, in the States of Rio Grande do Sul, Pará, Paraíba, Santa Catarina, Bahia, and Ceará. Other countries with records included Venezuela and Colombia\textsuperscript{(76)}.

The Ministry of Health reported 112 outbreaks nationally between 2005 and 2013, involving all 35 municipalities in...
the Amazon Region. The probable source of infection was food contaminated with T. cruzi, including: açaí, bacaba, jaci (Syagrus), sugar cane juice, and palm hearts of babaçu nuts. Most of the outbreaks occurred in the States of Pará, 75.9% (85 outbreaks) and Amapá, 12.5% (14 outbreaks) and, to a lesser extent, in the States of Amazonas, 4.5% (5 outbreaks), Tocantins, 1.8% (2 outbreaks), and Bahia, 1.8% (2 outbreaks)\(^{(55)}\).

The process of surveillance of Chagas disease and acute oral transmission was potentiated by the increased sensitivity of the surveillance system. The first officially investigated outbreak of orally transmitted Chagas disease in Brazil occurred in Santa Catarina in 2005, probably linked to the intake of sugar cane juice contaminated with T. cruzi\(^{(55)}\). Between 2007 and 2013, it should be emphasized that more than 50% of orally transmitted Chagas disease cases presented with symptoms between August and November, a period that coincides with the açaí harvest in Pará\(^{(55)}\).

Available experimental evidence suggests that oral transmission may occur from trypomastigote, epimastigotes, and probably amastigotes forms of T. cruzi and cell masses, originating from mammals or contaminated vectors, as well as, accidentally, through artificial cultures of the parasite\(^{(21)}(23)(94)\). Depending on the predominance of a large group or lineage of T. cruzi transmitted, as well as of the inoculum in question, there is a diversity in pathogenicity, histiotropism, morbidity, and mortality\(^{(21)}\). In relation to the mechanism of oral transmission of T. cruzi, the following possible scenarios have been identified\(^{(21)}(39)(93)(94)\):

- ingestion of triatomine feces or urine of or even infected triatomines, on the assumption that these are processed or carried along with food, as verified in episodes investigated in which the infection was assigned to the consumption of açaí fruit, typical of the Brazilian Amazon Region\(^{(92)}\);
- intake of food or drinks contaminated with metacyclic trypomastigote forms in the secretion of the anal gland or in the urine of infected marsupials of the genus Didelphis;
- intake of suspensions of T. cruzi in pipettes in the context of research or diagnostic laboratories;
- intake of breast milk from mothers diagnosed with: acute Chagas disease, T. cruzi-HIV co-infection in the AIDS stage, documented reactivation of Chagas disease, or bleeding from fissures of the nipples.

Other speculative situations of exposure include\(^{(21)}(39)(93)(94)\):

- ingestion of raw or poorly cooked meat or blood from infected mammals, especially wild animals;
- consumption of the blood of infected animals for therapeutic purposes, as performed by some indigenous groups in the Amazon. This has been reported in some regions of Colombia, where the intake of blood of armadillos and skunks has been observed;
- contamination of equipment used in the handling of carcasses of infected mammals;
- ingestion of triatomines by primitive or exotic habits.

It is noteworthy that the feces of infected triatomines can retain infectious potential for a few hours in environments with high humidity. Therefore, they can potentially contaminate food directly or indirectly, by the feet and oral cavity of secondary carriers, like flies and cockroaches. Experimental studies have shown that the parasite can remain viable for ≥24 hours in foods such as milk or sugar cane juice if stored at room temperature. Although gastric juice is able to destroy a considerable number of parasites, some are able to escape this through chemical mechanisms of external protection, allowing the parasites to penetrate the intestinal mucosa\(^{(21)}(92)\).

Although oral transmission is generally associated with outbreaks, it can also occur in isolated cases\(^{(21)}(39)\). In fact, we can identify two main profiles of outbreaks, according to the place of occurrence: urban and rural. Urban outbreak derives from the consumption of semi-industrialized and commercialized açaí, reflecting an issue of sanitary surveillance, associated with food safety and good food handling practices. In rural outbreaks, there is the participation of various types of food prepared locally by hand, generally by family or people of the social network/community of a given family.

For research and surveillance purposes, food recall is a useful tool to identify the possibility of suspect food intake. It allows one to identify the source, place of preparation and consumption of suspect food, and to list all the people who may have ingested the suspect food\(^{(21)}(39)(92)\). The focus of surveillance actions and control should therefore include the productive chain, based on good food handling practices\(^{(21)}(39)\).

Transmission by blood transfusion and transplantation of tissues/organs

The transmission of T. cruzi in blood transfusions was amplified by the urbanization of Chagas disease in Brazil and in other endemic countries in Latin America\(^{(3)}(4)(10)(95)(96)\). The risk of transmission by blood transfusion is dependent on various factors: I) the presence of the parasite in the blood or blood component transfused; II) type and number of infected blood products transfused; III) immune status of the recipient; IV) quality of the clinical-epidemiological screening; V) level of coverage of the serologic screening of donors; and VI) sensitivity of serologic tests used to screen blood donation candidates\(^{(4)}(23)(95)(96)\).

Given that the majority of potential blood donors are aged 18–35 years—the age group with the lowest number of candidates potentially infected with T. cruzi—and that, in recent years, the obligation of clinical, epidemiological, and serologic status screening has been established in most endemic countries, the risk of transfusion transmission of Chagas disease has been largely reduced in Latin America as a whole\(^{(4)}(23)(95)(96)\). In Brazil in the 1950s, the estimated average prevalence of positive serology for T. cruzi among potential blood donors was 8.3%. This decreased to 6.9% in the 1960s and 1970s, dropping to 3.2% in the late 1980s and early 1990s\(^{(96)}\). A Health ministry report from January to June 1994 stated an estimated prevalence of Chagas disease through blood transfusion, by the public blood donation network to different geographic regions
of the country, of 0.75%\(^{(95)}\). In 2006, this estimate was 0.21%, a reduction of more than 95% compared with previous data\(^{(17)}\). Most recent WHO data from 2010, estimated this prevalence to be 0.18% in Brazil\(^{(95)}\).

In Brazil, not only the safety of blood transfusion has improved, but also the process of hemovigilance, with the establishment of a national system integrating blood banks and epidemiological and sanitary surveillance\(^{(97)}\). With this, one can classify the rare, but possible cases of Chagas disease transmitted by blood, associated with the residual blood transfusion risk and any failures within the process. The same applies for situations associated with screening of candidates for donation of tissues or organs. For both, sanitary inspection in hematology and transplantation services (hospitals or blood banks) are instituted, as is communication and integrated action with hemovigilance\(^{(97)}\), recalling that the transfusion transmission can only be determined if the infected person received blood or some other component within the 120 days before the onset of symptoms\(^{(97)}\).

A similar trend in the reduction of the risk of transmission from blood transfusions has been observed in other endemic countries, to a greater or lesser degree. For non-endemic countries, in the last decade several strategies have been adopted to prevent and control transfusion transmission of Chagas disease. Nations like the United States, Canada, Spain, France, the United Kingdom, Switzerland and Australia, for example, have already introduced strategies of universal or selective serologic screening of blood donation candidates\(^{(99)}\).

In case of organ transplants, endemicity patterns of areas of origin and residence should be considered, both for donors and recipients, in order to outline the best approach possible for each case in terms of screening and management of possible infection by \emph{T. cruzi}\(^{(100)}\).

**Vertical transmission (congenital)**

Vertical transmission of Chagas disease is still relatively important in Brazil, despite the absence of systematic preventive measures against this type of transmission in the country. To assess its impact and also the impact of vector control, between 2001 and 2008 a new national serologic survey was conducted, now having as a reference population 105,000 children aged 0–5 years\(^{(101)}\). The estimated prevalence of children infected with \emph{T. cruzi} was 0.03% (32 cases). Of these, 20 (0.02%) had concomitant maternal positivity, suggesting congenital transmission [coming from the States of Rio Grande do Sul (12), Minas Gerais (3), Paraná (1), Pernambuco (1), Alagoas (1), Bahia (1), and Acre (1)]. In the other 11 (0.01%) cases, only the child tested positive, indicating probable vector transmission [derived from the State of Piauí (2), Ceará (2), Paraíba (2), Alagoas (2), Rio Grande do Norte (1), Amazon (1) and Paraná (1)]. This study was the first to identify regional differences in the congenital transmission of Chagas disease in Brazil, possibly related to the existence of \emph{T. cruzi} TcV and TcVI\(^{(101)}\). Moreover, it points to the absence of Chagas disease transmission by a sustained domiciliary vector in the country\(^{(99)}\).

Official data from the Ministry of Health concerning epidemiological surveillance by reporting cases of acute Chagas disease indicate consistency with the findings obtained in the aforementioned survey: 50% of notifications of this form of transmission were from the State of Rio Grande do Sul\(^{(55)}\).

The overall estimate of the prevalence of \emph{T. cruzi} infection in pregnant women ranges from 1 to 40%\(^{(102)}\)\(^{(103)}\)\(^{(104)}\), and approximately 1.8 million women of childbearing age in Latin America are infected\(^{(17)}\). The latest data from WHO estimate that 1,124,930 women aged 15–44 years are infected with \emph{T. cruzi}\(^{(9)}\). A recent meta-analysis estimated the global risk for congenital \emph{T. cruzi} infection in children born to infected mothers as being 5% (range: 0–28.6%)\(^{(101)}\). In Latin America, more than 15,000 cases per year of congenital Chagas disease are estimated to occur\(^{(17)}\). More recent studies estimate that 8,668 children are vertically infected\(^{(35)}\). On the other hand, in non-endemic areas, the availability of evidence to estimate incidence is even more limited\(^{(3)}\)\(^{(17)}\)\(^{(104)}\). In North America, for example, the estimate is of 2,000 cases of congenital Chagas disease per year\(^{(45)}\).

A systematic review with meta-analysis of studies conducted in Brazil estimated the prevalence of infection in pregnant women to be 1.1% with a vertical transmission rate of 1.7%\(^{(106)}\). According to data from the Brazilian Live Birth Information System [Sistema de Informações sobre Nascidos Vivos (SINASC)], 2,861,868 live births were recorded in the country in 2010. From these data it was estimated that 34,629 pregnant women would be infected with \emph{T. cruzi} this year; the number of congenitally infected children would thus range from 312 to 1,073 (average: 589 cases)\(^{(106)}\). These data are consistent with those of an epidemiological survey conducted in the State of Minas Gerais in 1998, extrapolated to the Brazilian population\(^{(107)}\), and with the global estimate provided by the WHO of 571 Brazilian children infected by vertical transmission in 2010\(^{(4)}\). In 2006, PAHO estimated that 5,000 Brazilian children would be vertically infected\(^{(17)}\). Considering the gradual reduction in fertility rates in Brazil, it is estimated that over the next few decades, vertical transmission will be substantially reduced, with possible elimination as public health problem for the country in the next 10 to 20 years.

A recent retrospective study from a referral hospital of the Federal University of Goiás [Universidade Federal de Goiás (UFG)] analyzed 1,211 individuals born to mothers known to be infected by \emph{T. cruzi} and identified a rate of vertical transmission of 2%, consistent with previous estimates. This suggests that the presence of TcII in the study area may be associated with lower rates of transmission than TcV, which predominates in the southern regions of Brazil and other Southern Cone countries, where the vertical transmission rates are generally known\(^{(108)}\). Despite existing scientific evidence, important gaps remain in terms of the role of genetic diversity of \emph{T. cruzi} in relation to the evolution of the chronic phase of the disease, the risk of congenital transmission or reactivation, and the occurrence of oral outbreaks transmission\(^{(27)}\).

In Brazil, congenital Chagas disease is considered acute and therefore its registration is mandatory\(^{(20)}\). However, surveillance for Chagas disease specifically in pregnant women or exposed/infected children has not yet been established in the country. On the other hand, it is recognized that provision
of anti-T. cruzi treatment to non-pregnant infected women of childbearing age may be an effective strategy to reduce vertical transmission in future pregnancies\(^{(109)}\). It should also be noted that vertical transmission may be repeated in every pregnancy\(^{(104)}\).

**Accidental transmission**

Accidental transmission has been recorded in different contexts: triatomine laboratories, vector capture actions in endemic areas, experimental work with infected mammals and cultures, aerosols of infected materials, surgical infection and infection during collection of blood samples from people with acute infection. In these cases, safety deficiencies are evident in the transportation of contaminated materials, among other factors\(^{(93)}\). Risk factors include ignorance, inattention, lack or misuse of personal protective equipment, inadequate facilities and equipment, poor lighting, lack of training, non-compliance with standard precautionary measures, and non-adoptive of routine technical protocols\(^{(93)}\)\(^{(110)}\).

The process of monitoring and control must include the development of continuing education actions in health and supervision, to check the proper use of personal protective equipment and, if applied, the communication of work accidents within the demarcated processes for the health surveillance of workers\(^{(39)}\)\(^{(93)}\)\(^{(110)}\).

**Epidemiological surveillance**

Epidemiological surveillance is a set of strategic actions that provide knowledge, detection, or prevention of any change in determinants and conditioning factors of individual or collective health, in order to recommend and adopt measures for the prevention and control of diseases and disorders\(^{(39)}\). In the case of Chagas disease, it includes necessarily integrated actions involving human cases, vectors, and reservoirs, interfacing closely with the health care network, with emphasis on primary care or primary health\(^{(41)}\)\(^{(115)}\).

Systematized below are the main elements in Brazil for the development of surveillance processes for human cases and for entomological surveillance. Several questions related to this process have already been presented and previously discussed.

**Chagas disease cases surveillance**

Epidemiological surveillance actions for Chagas disease in Brazil have the following main objectives\(^{(20)}\)\(^{(39)}\):

- to detect early cases of acute Chagas disease, with the aim to treat newly diagnosed cases timely and to apply preventive measures of occurrence to new cases;
- to conduct epidemiologic investigation of all acute cases, aiming to identify the form of transmission and, consequently, to institute appropriate control measures;
- to monitor T. cruzi infection in humans, through periodic serologic surveys in key populations, and by means of adopting a national screening process of blood donation candidates in blood donation centers;
- to monitor the profile of morbidity and mortality of Chagas disease in the country, outlining scenarios for strengthening the healthcare network for infected people;
- to maintain the elimination of vector transmission by T. infestans and to monitor/control other important species involved in transmission to humans;
- to integrate health surveillance and environmental actions (regarding vectors and reservoirs) with epidemiological surveillance activities.

The change in the epidemiological patterns of the disease in the country required structuring of Chagas Epidemiological Surveillance in the Northern region due to oral transmission. There is a clear need to closely interfacing with the Health Surveillance, with the aim to improve prevention and control measures, besides the definition of reference flows for the diagnosis, treatment and monitoring of disease complications.

The high percentage of cases in the Health Ministry database with an undetermined mode of transmission suggests a weakness of the surveillance process and indicates the need for improvement in the opportunity of surveillance activities regarding the detection and investigation of suspect cases\(^{(55)}\). It also reinforces the need for quality health surveillance actions in the SUS, potentiated through continuing education activities and monitoring and evaluation by the health teams.

The occurrence of suspected cases of acute Chagas disease in Brazil requires immediate notification (within 24 hours after suspicion) using a specific instrument, the Acute Chagas Disease Research Sheet, which is standardized throughout the national territory. For epidemiologic surveillance purposes, the following case definitions have been established\(^{(39)}\):

1. **Suspected case of acute Chagas disease**

- Person with persistent fever (more than 7 days) with one or more of the following clinical manifestations: swelling of the face or limbs, rash, lymphadenopathy, hepatomegaly, splenomegaly, acute cardiomyopathy (tachycardia and signs of heart failure), hemorrhagic manifestations, jaundice, Romaña's sign, inoculation chagoma, or:
  - He/she has had a direct contact with triatomines or their excreta; or
  - He/she has received blood/blood products or transplantation of cells/tissues/organs contaminated with T. cruzi; or
  - He/she has ingested food suspected of being contaminated with T. cruzi; or
  - He/she is a newborn, from an infected mother.

2. **Confirmed case of acute Chagas disease**

**Laboratory criteria**

- Parasitologic: T. cruzi circulating in the peripheral blood identified by direct parasitologic examination.
- Serologic: suspected case with serology reactive with anti-T. cruzi IgM antibodies by the indirect immunofluorescent antibody test (IFA); or serology reactive with anti-T. cruzi immunoglobulin G (IgG) antibodies by indirect IFA, with change in IgG concentration of at least two titers after a minimum interval of 21 days in preferably paired samples; or seroconversion by any of the methods [enzyme-linked immunosorbent assay (ELISA), indirect hemagglutination (IHA) or indirect IFA].
Clinical-epidemiological criteria

Cases of acute Chagas disease should always be confirmed by laboratory diagnosis. Only in specific situations can clinical-epidemiological criteria be used for suspected cases, such as during an outbreak of the disease in the acute phase by oral transmission. Such cases have initial negative or non-reactive serologic and parasitologic test results but possess an epidemiological link to confirmed cases of acute Chagas disease by laboratory criteria.

3. Discarded case of acute Chagas disease

This refers to a suspected case with negative or non-reactive laboratory test results, or in whom another illness is diagnosed. Due to the limited clinical expression presented in many cases in the acute phase and the long, silent course of the disease, the reported case can be discarded as being a chronic case or a case of disease reactivation.

4. Definition of a case according possible forms of transmission in Brazil

Reserved for confirmation of cases of acute Chagas disease characterized according to the probable form of transmission.

Confirmed case of Chagas disease by oral transmission

Case in which other routes of transmission are excluded, with epidemiological evidence of a food as a common source of transmission, and usually the simultaneous occurrence of more than one case with an epidemiological link (origin, customs, and cultural elements).

Confirmed case of Chagas disease by vector transmission

Case in which other routes of transmission are excluded, with clinical evidence (Romãña’s sign or inoculation chagoma) and/or epidemiological evidence of triatomine occurrence at the site of the infection. Typically, this is observed as an isolated case.

Confirmed case of Chagas disease by laboratory accident

Case that had contact with T. cruzi cultures, exposure to contaminated triatomine feces or blood (human or animal cases) containing infective forms of the parasite. Usually, this occurs in laboratory scientists, health professionals, or researchers.

Confirmed case of Chagas disease by vertical transmission

Newborn from mother with positive parasitologic examination or reactive serology with T. cruzi and presenting:

- Positive parasitologic examination from birth; or
- Serologic examination reactive from 9 months of age (before that, maternal antibodies may still be present in the child) and no evidence of infection by other forms of exposure to T. cruzi.

In addition to the surveillance of current outbreaks for the specific surveillance of Chagas disease in Brazil, with the revision of the case definition for epidemiological surveillance of AIDS in the country, from January 2004 throughout the national territory, reactivation Chagas disease (myocarditis and/or meningoencephalitis) has been introduced as indicative of AIDS for SUS(49)(54).

Although so far, these are not the subject of epidemiologic surveillance in Brazil(46), individuals with the chronic form of Chagas disease should be confirmed, considering the evaluation of people with no indicative picture of febrile illness in the last 60 days and the presence of one of the following tests(39):

- Serology: Anti-T. cruzi IgG reactive by two methods based on different principles [ELISA, indirect fluorescent antibody technique (IFAT) or IHA]; or
- Xenodiagnosis (artificial or indirect): positive for T. cruzi; or
- Positive blood culture for T. cruzi in blood samples or cerebrospinal fluid; or, even from postmortem samples.

It is noteworthy that the current epidemiological context in Brazil brings to debate the expansion of the focus and approaches of epidemiological surveillance in the human population for Chagas disease in its chronic phase, considering, among other factors, the possibility of increased sensitivity of epidemiological research actions of other cases in the family, social network and communities (in the acute phase or chronic), as well as the design of spatial contexts of transmission (past or recent, active or not) in a manner integrated with vectors and reservoirs. At the same time, consideration should be given to expanding the criteria for the specific treatment of Chagas disease, considering that millions of infected Brazilian citizens would clearly benefit(15)(59). Therefore, these questions bring the need to overcome operational problems of the health care network, which should be prepared for longitudinal management of this chronic condition. There is also a need to guarantee sustainable production of medicines, given the potential increase in the demand for specific treatment, and international agreements(38). Extensive national discussion is therefore ongoing, to assess the inclusion of surveillance through mandatory reporting of human cases in the chronic phase in order to qualify the actions of both epidemiology and control, and health care.

Entomological surveillance

Entomological surveillance for Chagas disease should be implemented throughout the country, supported mainly on two pillars: passive surveillance, with the participation of the population in the notification of triatomine; and active surveillance, carried out by entomological teams of the municipalities in partnership with regional health states, without necessarily being based on prior notification by the resident(39).

The entomological surveillance has been enhanced with a community-based support, responsible for the network of Triatomine Information Posts (TIPs), which is being slowly consolidated in the country(22). The home research by the institutional technical team has been recommended in a more systematic and comprehensive form, depending on the existing risk. However, few municipalities have adopted this approach, and those that developed it, generally, have low sensitivity in active search, not detecting at least half.
of infestation foci. Systematic review addressing impacts of community participation in the entomological surveillance process found that the development of strategies to ensure such participation should be incorporated as a component of entomological surveillance process. This study emphasizes that only the standardized chemical control seems to be consistently effective in eliminating foci of infection. It was strengthened, therefore, the necessary participation of individuals, families and communities of endemic areas in all stages of this process, from the planning to actions evaluation, considering the social dimensions in which they fall, and recognizing the need for empowerment and participation of these populations.

The strategies adopted must be appropriate to the reality of each location, since passive surveillance with the participation of the population is a priority in Brazil and is suitable for the vast majority of scenarios, safeguards the necessary and prompt response by the services to the demands of the population. On the other hand, active surveillance should be carried out obligatorily in municipalities with residual foci of T. infestans, and recognizing the need for empowerment and participation of these populations.

Amplification of T. cruzi transmission, registered in areas previously considered harmless or with no risk for transmission of Chagas disease, such as the Amazon region, required greater attention to the epidemiologic surveillance teams. In several Brazilian regions, environmental changes favoring the adaptation of vectors to artificial environments, thus establishing new spaces for the appearance of disease, have been observed. Similarly, it has also been observed the domiciliation of secondary species, even in a much smaller scale to that seen previously to the control of T. infestans.

The emergence or re-emergence of Chagas disease in Brazil characterizes a new epidemiological profile that is independent of the intradomiciliary transmission by T. infestans. Thus, the control of transmission of T. cruzi, in the current scenario, should be examined under a new perspective. Surveillance attention is extremely important, especially mainly maintaining the functioning of sensitive entomological surveillance in the municipalities, in addition to health and environmental educational actions.

**Challenges for Brazil**

In Brazil, the SUS is based on core values and principles within society, such as universality, equity, integrity, social participation and control, which should be the foundation for policies and programs of public interest. The country is one of the main endemic areas of Chagas disease in the world, in various contexts of great complexity for prevention and control. Due to the high morbidity and mortality burden associated with Chagas disease and their relative invisibility in society, the government must ensure its prioritization for public health and mobilize necessary resources and capabilities with other state actors or non-governmental organizations to its confrontation, by strengthening the singular role that primary care plays for the national health system.

National estimates of millions of people infected by T. cruzi indicate the great responsibility of the country in the technical-scientific and political fields, not only for the prevention of new cases, but especially in the implementation of better decisions and benefits for patients with disease that are both acute and chronic. Brazil and the other Latin American countries have a key role in conducting this process and implementing the action commitments internationally agreed under the seals of PAHO and WHO. The emerging social movement should be valued and promoted, in order to seek the sustainability of actions for coping.

The health surveillance activities for Chagas disease, based on primary health care, should have as principles: territorialization, intersectoriality, focus on people and not the disease, formation of multidisciplinary teams, focus on the needs and expectations of the population and finally, the search quality. The qualified and ethical approach of families affected by Chagas disease should be integrated with promotion and prevention actions, health care for diagnosis and timely treatment, but also to physical, psychological and social rehabilitation. Similarly, for transversal epidemiological surveillance activities in these territories. Therefore, new pacts and agendas must be built, by inserting the Chagas disease as a relevant issue, aimed at ensuring access to inputs necessary for the diagnosis and treatment of the disease in the SUS.

Therefore, in addition to the elements presented previously, the country must continually review the current goals and focuses of epidemiological surveillance of Chagas disease, considering the accumulation of scientific evidence and successful experiences. The social and human development must be strongly linked to promote strategic research focusing on overcoming science, market and public health system failures, with a broad popular participation and the emerging social movement in Chagas disease. The establishment of the International Federation of Associations of People Affected by Chagas Disease (FINDECHAGAS), with the important participation of various representations of Brazil and other countries, strengthens the possibility of raising and catalyzing discussions and decisions from a technical and political point of view based on evidence, to reach the true control and prevention.

The Ministry of Health, in partnership with states, Federal District and municipalities, has played a key inducer role and should be guaranteed the strengthening and sustainability of the National Program for Chagas Disease Control, to enable the scientific evidence, many of which are summarized in this Consensus, in fact be applied in the different realities of the SUS. It is worth to reiterate its strategic role to ensure the sustainability of research funding for control and prevention of Chagas disease in Brazil, more adjusted to different situations and contexts in the country.

The expansion of epidemiologic surveillance activities, in addition to acute cases of Chagas disease, integrating cases of the disease in the chronic phase, has to be incorporated strategically in this perspective, expanding network access to health care for diagnosis, timely infection treatment and potential complications of the disease. The antiparasitic treatment should be guaranteed to all cases that have indication for its use, and new safe and effective treatment options must be continuously pursued.
The priority areas for the development of monitoring programs should be designed for both current and future epidemiological scenarios, encompassing the redefinition of instruments and tools with innovative character for the development of monitoring procedures, integrating environmental aspects, reservoirs, vectors, parasites and human population. The macroecologic and eco-geographic perspective should be integrated into the planning of actions in the country, in coordination with the other nations of the Southern Cone Initiative against Chagas disease.

Considering the risk contexts outlined for Brazil, it should be defined the role of other forms of surveillance and how these will be integrated into existing processes of surveillance, including: hemovigilance, technovigilance, pharmacovigilance, vertical transmission, accidents with biologic materials, transplants, HIV/AIDS infections, other immunosuppressive conditions, unusual acute events, such as outbreaks and microepidemics, among others.

It should be emphasized that the integration of Chagas disease control program to other communicable disease control programs (vectorial programs or not) is required and non-communicable chronic diseases in the country - all linked mainly to the Secretariat of Health Surveillance [Secretaria de Vigilância em Saúde (SVS)], as well as with other areas of the Ministry of Health: Secretariat of Science, Technology and Strategic Inputs (SCTIE), Secretariat of Health Care (SAS), Secretariat of Strategic and Participative Management (SGEP), Secretariat of Labor Management and Education Health (SGTES), and Special Secretariat of Indigenous Health (SESA), aiming to strengthen intersectorial actions for the effective control of Chagas disease.

Faced with such prospects, it is justified the construction of this consensus, which, in addition to focusing on the Brazilian reality, could be a reference tool, encouraging and mobilizing all actors involved in the global fight against Chagas disease.

Vertical transmission of \textit{Trypanosoma cruzi}

Vertical transmission (from mother to child) of \textit{T. cruzi} remains a reality, even with the favorable impact of vector control actions and the qualification of the transfusion process in several countries, including\textsuperscript{104-105,112,113} included in Brazil\textsuperscript{20} \textsuperscript{(106,107). The existence of infected women in childbearing age, in endemic countries or not, supports this risk, which tends to be reduced over the next two decades.}

In Brazil, in 2011, according to SINASC data were collected about 3 million live births. Based on the estimated prevalence of 1% of \textit{T. cruzi} infection in women between 25 and 44 years of age\textsuperscript{17} and a risk of vertical transmission estimated as maximum of 1% for the State of Minas Gerais, it is estimated that up to 300 children were born infected that year\textsuperscript{107}. Detection of vertical transmission is complicated in practice, since the vast majority of congenital cases is totally asymptomatic\textsuperscript{14,104,109,112,113,114,115,116}.

It is a complex issue, considering the existence of little scientific evidence of population base. In Brazil, as congenital Chagas disease is considered acute, its notification is compulsory within the epidemiological surveillance activities.

Diagnosis and management of \textit{Trypanosoma cruzi} infection during pregnancy

It is recommended to carry out a serologic assessment (screening) for \textit{T. cruzi} infection in all pregnant women living in or are from endemic areas, preferably during their first prenatal visit\textsuperscript{20,117}. The Technical Group on Prevention and Control of Congenital Transmission and Case Management of Congenital Infections (IVA) of WHO Programme on Control of Chagas disease also recommends that screening for those women who have a history of having received blood transfusions performed in endemic areas or who were born in these areas\textsuperscript{104}. Although transmission depends fundamentally by maternal parasitemia, there is no clear evidence on which women may actually be transmitting the infection, which emphasizes the importance of prenatal screening\textsuperscript{112,116,117}.

The transmission may occur at any time during pregnancy, possibly being greater in the third quarter, with increasing parasitemia\textsuperscript{116} \textsuperscript{(118). On the other hand, the timely diagnosis of infection during pregnancy allows a more qualified care to the infected pregnant women, who should be monitored throughout gestation.

Reports of accidental exposure to benznidazole in pregnant women do not indicate adverse effects on the newborn. However, given the evidence of teratogenicity demonstrated in animals, the specific antiparasitic treatment of infection by \textit{T. cruzi} is contraindicated during pregnancy and not recommended during breastfeeding\textsuperscript{20,115}, and should be given only after these periods\textsuperscript{115,116}. Accidental exposure to antiparasitic drug during pregnancy is not a criterion for interrupting the gestation\textsuperscript{115}.

Studies point to the possible benefit of antiparasitic treatment, with better evolution of Chagas disease\textsuperscript{116,119,120} \textsuperscript{(121,122) and reduction of the risk of vertical transmission in future gestations\textsuperscript{109,116}. High maternal parasitemia is associated with an increased risk of vertical transmission and abortion\textsuperscript{105,115}. Pregnant women who are in the acute phase of Chagas disease should be evaluated case by case, preferably with the participation of experts, to define the risk-benefit of indicating the antiparasitic treatment. The pregnant women with chronic Chagas disease must be accompanied in referral centers for high-risk pregnancy, as they may present the need for high-complexity care.

The evidence of \textit{T. cruzi} infection does not justify the abdominal delivery indication (Cesarean\textsuperscript{115}) since the congenital \textit{T. cruzi} infection may result in delayed uterine growth, detected in ultrasound, and premature delivery\textsuperscript{102,115,116,123}.

The importance of carrying out all the recommended evaluations during prenatal care, including anti-HIV testing is emphasized. The coinfection of \textit{T. cruzi} and HIV in pregnant women increases the risk for congenital transmission of \textit{T. cruzi}, by high parasitemia, also implying an increased morbidity and mortality\textsuperscript{20,116,120,123} \textsuperscript{(124).

After the delivery, the woman should be evaluated as to the conduct of Chagas disease, from the clinic of this disease, aiming at an appropriate specific treatment.
**Diagnosis and management in exposed children**

The risk of vertical transmission of *T. cruzi* depends on the level of parasitemia, maternal immune status, infecting strain and placental factors, and might occur at any stage of disease (41)(104)(113)(115)(116).

Although most cases are asymptomatic, in some cases prolonged fever, hepatosplenomegaly, respiratory failure, prematurity, low birth weight and stillbirth may occur. Signs of meningoencephalitis and myocarditis were observed in coinfection with HIV (102)(104)(113)(114)(115)(123). Therefore, they have been recommended as routine assessments in children with clinical signs of congenital Chagas disease: complete blood count, serum biochemistry, urinalysis, chest X-ray, electrocardiogram (ECG), echocardiogram, and cerebral and abdominal ultrasonography (115).

The strategy employed for the diagnosis of *T. cruzi* infection depends essentially on the child’s age. Children exposed to *T. cruzi*, by vertical transmission, have circulating maternal IgG antibodies that can be detected by routine serologic testing up to 9 months of age, and their detection in this period, does not necessarily characterize a congenital infection. In turn, the persistence of unaltered titer of anti-*T. cruzi* antibodies in children from 9 months old, is indicative of congenital infection, and the absence of these antibodies removes the possibility of infection in children (104)(123).

The diagnosis in suspected cases of Chagas disease by vertical transmission (excluding other forms of transmission), because it is an acute infection, should be confirmed using direct parasitological methods (fresh examination, micro-hematocrit, leukocyte cream and/or Strout method) in cord blood or of the newborn in the first 30 days of age (preferably in the first week of life), with evaluation of two or three samples in the absence of signs and symptoms to expand sensibility (20)(104)(123).

For symptomatic children or newborn from mother presenting an acute *T. cruzi* infection or reactivation in the presence of coinfection with HIV/ *T. cruzi*, these parasitological tests should be performed repeatedly and, if negative, parasitological methods of enrichment and/or molecular (in research laboratories) have been used. However, these are not accessible and standardized for use in the routine of health services (115)(125).

In case of negative parasitological/molecular tests in the first months or in the impossibility of using parasitological methods, it is recommended to search for anti-*T. cruzi* IgG antibodies after the ninth month in two tests (20)(104)(123)(125). Conventional reagent serology in children during this period is strongly indicative of congenital transmission, especially when excluding possibilities of vector transmission and transfusion (20).

In Brazil, the inclusion of serologic tests searching IgG for *T. cruzi* in the National Newborn Screening Program (heel prick blood test) is recommended, especially in endemic areas for *T. cruzi* infection, representing an useful and low cost strategy (20)(107).

The search for anti-*T. cruzi* IgM antibodies has a low sensitivity and there are still difficulties to standardize this technique and obtain controls. The use of serological methods employing recombinant antigens, such as the Shed Acute Phase Antigen (SAPA), it may be used if available. There are reports that anti-SAPA maternal antibodies disappear earlier than conventional antibodies in about three months (126).

Once the diagnosis of *T. cruzi* infection has been established, the child must be treated with benznidazole (10–15 mg/kg/day in two or three doses for 60 days); this is a well-tolerated regimen (20)(104)(115). There is a clear need to develop formulations in suspension that are more suitable for this treatment, despite the recent advance in providing pediatric tablets (12.5 mg) for these clinical situations. Treatment is mandatory in all cases of congenital infection, since it treatment is highly efficacious and safe for the vast majority of children. In addition, studies show a high cure rate (≥ 95%) when treatment is started before 1 year of age, already in the first weeks of life (104)(112)(115)(127).

Clinical and laboratory evaluation (blood count testing) should be performed at the beginning of the treatment and 30, 60, and 90 days after the use of benznidazole. In children with the clinical syndrome of Chagas disease, other complementary assessments should be carried out according to the table presented. The cure control must be performed with serologic tests every six months with titration until the child has two consecutive non-reactive serologic test results. Persistence of positive serology or evidence of positive parasitologic tests may indicate treatment failure and the child must be monitored and evaluated for possible complications of the disease. In refractory cases, benznidazole therapy may be repeated or may be substituted with nifurtimox (up to 15 mg/kg/day in two or three doses, for 60 days).

In newborns from infected mothers and asymptomatic, whenever possible, *T. cruzi* research should be performed. Those with initial negative parasitologice examination or not carried out must be submitted at nine months after birth, for the research with serologic tests of anti-*T. cruzi* IgG antibodies or conventional serology for infection by *T. cruzi* (107)(117)(127). A reactive serologic test after this age necessarily implies the need to initiate antiparasitic treatment. On the other hand, no evidence of positive serology excludes infection.

It is not recommended to suspend breastfeeding of infants whose mothers have Chagas disease in the chronic phase - except in cases where there is bleeding by nipple fissure, when the interruption of breastfeeding in the affected breast would be appropriate - or in situations of high parasitemia, such as in reactivation of Chagas disease or in women in the acute phase of the pathology (112)(115). If exposed to breast milk of mother with acute or chronic infection with the presence of nipple cracks, the infant should be monitored for the acquisition of *T. cruzi* infection during the exposure period, using the same parasitologic and/or serologic criteria described above. In some of these cases, heat treatment of milk prior to administration to infants can be considered (122).

In *T. cruzi*/HIV coinfection, it is recommended that all infected mother is advised not to breastfeed, considering that breastfeeding, regardless of the association with Chagas disease, it is associated with an additional 7–22% risk of HIV transmission. Meanwhile, in cases of acute maternal infection by HIV, breastfeeding increases the likelihood of vertical transmission of HIV to 29%. In Brazil, the mother is entitled to receive infant milk formula, at least until your child reaches 6 months of age (126). Figure 1 shows the general flow diagram for the approach of *T. cruzi* infection in the mother/child pair.
Pregnant women living in an area endemic to Chagas disease

1st prenatal consultation

Two anti-*T. cruzi* serological tests in the pregnant women - different methods [ELISA / indirect IFA / IHA]

At birth

- Confirmed infection by *T. cruzi* (according to diagnostic flowchart)\(^a\)
- Discarded infection by *T. cruzi* (according to diagnostic flowchart) \(^a, b\)

Forward to high-risk prenatal care - evaluate the commitment of the pregnant woman (clinical evaluation and ECG) \(^b, c\)

In the postpartum period

- Refer the mother for medical control of the disease and therapeutic evaluation

Assess *T. cruzi* infection in the newborn

- Absence of clinical alterations suggestive of congenital infection
- Presence of clinical alterations suggestive of congenital infection

Assessment of *T. cruzi* \(^d\)

- Positive result
  - Treatment of infection by *T. cruzi*
- Negative result
  - Assessment of *T. cruzi* again (one week after birth)

Assessment of *T. cruzi* again (one week after birth)

- Positive result with clinical alterations
  - Child serology (2 tests after 9 months age) \(^e\)
- Negative result without clinical alterations
  - Confirmed congenital infection by *T. cruzi* (according to diagnostic flowchart) \(^a\)
  - Discarded congenital infection by *T. cruzi* (according to diagnostic flowchart) \(^a\)

\(^a\) Follow flowchart to confirm the diagnosis of infection by *T. cruzi* (see section on laboratorial diagnosis of infection by *T. cruzi* in this Consensus)

\(^b\) If the complementary exams do not confirm changes, prenatal assistance can be done within the routine procedures of the Basic Health Unit (primary health care). If signs/symptoms suggestive of Chagas disease are present, one should continue prenatal assistance in a reference care unit for high risk pregnancies.

\(^c\) Evaluation of the fetus by obstetric ultrasonography; one should look for signs of restricted uterine growth and other signs common in the TORCH group (Toxoplasmosis, Other infections, Rubella, Cytomegalovirus, and Herpes).

\(^d\) Parasitological tests are recommended in the first days of life of the child.

\(^e\) After the 9th month of life use serological tests for the diagnosis of infection.

**FIGURE 1.** Flowchart for investigating and managing *T. cruzi* infection in both the mother and child.
LABORATORY DIAGNOSIS OF TRYPANOSOMA CRUZI INFECTION

In Brazil, an attempt to establish the etiologic diagnosis of Chagas disease should be made in all suspected cases, both in the acute phase and in the chronic phase. Therefore, it is essential to integrate epidemiologic, clinical, and laboratory evidence, in order to increase the degree of predictive and diagnostic accuracy. The complementary diagnosis of infection by *T. cruzi* through various laboratory techniques should follow defined criteria, depending on the stage of the disease. The following recommendations for diagnosis include the current Brazilian regulations.

**Laboratory criteria to define the diagnosis of Chagas disease**

**Acute phase**

*Parasitologic criterion*

Parasitologic examination is the most appropriate at this stage. This criterion is defined by the presence of *T. cruzi* trypomastigotes, identified by direct examination of the peripheral blood (with or without centrifugation) using microscopy (with or without staining).

Simultaneous performance of different modalities of direct parasitological examinations is recommended - fresh examination of trypanosomatids, concentration methods, or thick smear or smear stained slide. When the results of the fresh examination and concentration methods are negative in the first collection, new collections must be carried out until the case definition and/or the disappearance of the symptoms of the acute phase, or until the confirmation of another diagnostic hypothesis.

*Serologic criterion*

The serologic criterion is based on indirect methods for diagnosis that may be performed when parasitologic tests are negative but clinical suspicion persists. Such methods have complementary usefulness, and should always be performed in suspected or confirmed cases of acute Chagas disease.

When the parasite is not identified by direct observation, verifying the presence of anti-*T. cruzi* IgM antibodies in the peripheral blood is considered suggestive of the acute stage, particularly when combined with the patient’s epidemiologic context and clinical events. This represents a more complex technique and is most suitable in the late acute phase, when repeated direct examinations are negative.

Seroconversion to *T. cruzi* infection is defined by an initial non-reactive serum sample for anti-*T. cruzi* IgG antibodies, followed by a reactive sample collected 2–4 weeks later, based on an assay that includes both samples simultaneously.

Alternately, an increase of at least two titers between two reactive samples taken 2–4 weeks apart, in a clinical and epidemiologic context suggestive of acute Chagas disease, may also be considered diagnostic of acute Chagas disease.

In situations where vertical transmission is possible, the strategy used for the diagnosis of infection by *T. cruzi* would depend on the child’s age; this is discussed in a specific part of this document. It is noteworthy that every case of congenital transmission means an acute case of Chagas disease and, as such, should be reported and managed.

**Chronic phase**

*Parasitologic criterion*

Due to the low parasite burden in the blood in the chronic phase of Chagas disease, parasitologic methods of enrichment/multiplication, blood culture and xenodiagnosis have demonstrated low sensitivity, which implies the absence of diagnostic value when the result is negative.

When positive, the results are useful, mainly to monitor the specific treatment, or in unusual cases where serology presented inconclusive results.

*Serologic criterion*

In the chronic phase, the diagnosis is essentially serologic, and must be performed using a test with high sensitivity in conjunction with another having high specificity.

An individual is considered in the chronic phase of infection if anti-*T. cruzi* IgG antibodies are detected by means of two serologic tests with different principles/methods having different antigenic preparations. Other diseases (for example, visceral leishmaniasis, virchowian (lepromatous) hanseniasis, and autoimmune diseases) should be considered in the differential diagnosis.

**Principles and Guidelines for the Laboratory Diagnosis of Trypanosoma cruzi Infection**

**Parasitologic diagnosis in the acute phase**

The parasitologic diagnosis of Chagas disease at this stage is mainly based on the identification of the parasite, and its sensitivity depends on the level of parasitemia. In the acute phase, the number of parasites in the peripheral blood is high. It is recommended that, if presented with a suspected case of Chagas disease in the acute phase, different methods of direct parasitologic examination be used for immediate and repeated reading, in order to clarify the diagnosis.

The fresh search for trypanosomatids is quick and easy to perform, and is more sensitive than staining the smear. The ideal situation is to collect them from a febrile patient within 30 days of symptoms onset. The examination can be performed directly under the microscope with a drop of blood between the slide and cover slip. Blood collection should be carried out simultaneously for techniques of blood concentration.

The concentration methods are quick and inexpensive (Strout’s method, micro-hematocrit anduffy coat). They are recommended as the first choice of diagnostic test for symptomatic cases with more than 30 days of evolution, due to the decline in parasitemia with time. Blood samples should be examined within 24 hours due to possible lysis of the parasites.

Direct examination of a stained thick blood drop or blood smear can be used, but this has lower sensitivity than the methods described above. However, in the Northern region of Brazil, belonging to Legal Amazon, this method is widely
used; it is convenient and able to be integrated into malaria diagnostic actions\(^{(20)}\)\(^{(39)}\). It also represents an important method for verification and morphologic characterization, especially in geographic areas where the infection by \textit{T. rangeli} can coexist with \textit{T. cruzi}.

It is emphasized that in cases with high parasitemia—as in the acute phase of the disease, but also in transfusion transmission and in immunocompromised individuals—parasites can be incidentally found on blood smear when performing a differential leukocyte count\(^{(20)}\)\(^{(39)}\)\(^{(49)}\)\(^{(52)}\)\(^{(100)}\).

In cases where a strong clinical and epidemiological suspicion of acute phase \textit{T. cruzi} infection exists but direct parasitologic examinations are negative, molecular diagnosis using polymerase chain reaction (PCR) methods with hybridization has shown promising results. However, these are in-house tests, performed by only a few research/reference centers. The methods, protocols, and operating procedures should follow the recent recommendations for the standardization for the PCR use\(^{(143)}\)\(^{(144)}\). In Brazil, due to the absence of defined protocols and standard operating procedures, as well as of commercial kits to be routinely used in health surveillance, PCR cannot be considered an isolated diagnostic method to confirm or discard cases of acute or chronic Chagas disease\(^{(39)}\).

### Serologic diagnosis in the acute phase

In Brazil, there are currently difficulties in performing serologic tests in patients in the acute phase, due to the lack of registered commercial kits approved by the Brazilian National Health Surveillance Agency (ANVISA) and difficulty in obtaining positive controls for IgM\(^{(20)}\)\(^{(39)}\)\(^{(133)}\). Therefore, it has been traditionally recommended that indirect IFA methods are performed in reference laboratories to detect IgM\(^{(145)}\) in addition to the conventional techniques already in use, such as indirect IFA to detect IgG, IHA\(^{(146)}\), and ELISA\(^{(147)}\).

### Parasitologic diagnosis in the chronic phase

Indirect conventional methods for the isolation and identification of \textit{T. cruzi} (xenodiagnosis and blood culture) have low sensitivity; sensitivity can be increased through repetition\(^{(20)}\)\(^{(129)}\)\(^{(133)}\)\(^{(148)}\)\(^{(149)}\). A negative test does not rule out the possibility of infection, but a positive test has an absolute diagnostic value. The PCR at this stage, despite being limited by the lack of standardized protocols, is indicated when serologic tests present an indeterminate result or to determine cure after antiparasitic treatment. PCR should be performed by laboratories with recognized competence, carried out by experts in the field\(^{(150)}\).

### Serologic diagnosis in the chronic phase

Diagnosis in the chronic phase is essentially serologic, and should be performed using a test with high sensitivity (ELISA with total antigen or indirect IFA) in conjunction with another method with high specificity (IHA\(^{(129)}\)\(^{(133)}\)\(^{(134)}\)). Known conventional tests (IHA, indirect IF, and ELISA\(^{(133)}\)) can determine the diagnosis in almost 100% of cases. Unconventional tests (with recombinant antigens, for example) may preferentially be used in parallel with another conventional test\(^{(151)}\).

Performing reactions in eluates from collected blood on filter paper is not recommended to diagnose the infection, although these are usually used in the screening steps of epidemiological investigations\(^{(152)}\). The Guerreiro and Machado reaction (or complement fixation) for Chagas disease does not meet the standards currently required. Since it is not commercially available, it is also not indicated\(^{(20)}\)\(^{(129)}\).

The chemiluminescence test also allows the identification of IgG antibodies. While some kits are available on the market, the technique is still not recommended by the Ministry of Health of Brazil\(^{(133)}\).

The recent evaluation of 11 rapid test kits used for serologic diagnosis of Chagas disease indicates the potential of these methods. However, further studies should be conducted in the laboratory and in the field to confirm these data, in particular assessing their reproducibility in the context of limited resources or the use of whole blood in the real contexts of endemic and non-endemic areas\(^{(153)}\).

The following flowchart (Figure 2) summarizes the steps for the laboratory diagnosis of \textit{T. cruzi} infection in the chronic phase of the disease.

### ACUTE PHASE OF CHAGAS DISEASE

Acute Chagas disease is becoming a less frequent event in endemic countries, including Brazil, where control of vector (especially by \textit{T. infestans}) and transfusion transmission has altered the epidemiological scenario\(^{(20)}\)\(^{(39)}\). On the other hand, the occurrence of cases and outbreaks by oral transmission, transmission by domiciliary vectors without colonization and by extradomiciliary vectors, especially in Amazônia Legal, has gained epidemiological importance\(^{(20)}\)\(^{(21)}\)\(^{(39)}\)\(^{(76)}\)\(^{(154)}\)\(^{(155)}\)\(^{(156)}\).

Urbanization and globalization of Chagas disease due to large migrations, both in endemic and non-endemic countries, has contributed to making the epidemiological scenarios even more complex. In addition, reactivation episodes of Chagas disease associated with immunodeficiencies have become emerging challenges for health systems\(^{(20)}\)\(^{(39)}\)\(^{(49)}\)\(^{(157)}\)\(^{(158)}\)\(^{(159)}\). In this sense, the epidemiological investigation of the mode of transmission of Chagas disease is of particular importance, not only in order to determine actions to be implemented, but also because the different routes potentially lead to specificity in the clinical expression of the acute disease\(^{(20)}\)\(^{(39)}\)\(^{(76)}\)\(^{(132)}\).

### Clinical and complementary examination aspects

#### Acute phase of Chagas disease by vectorial transmission

The clinical picture is characterized by the appearance of a set of manifestations of variable intensity after an incubation period that is inversely proportional to the load of the inoculum and the inoculation route. There may be a lesion at the inoculation site, a sign of infection at the inoculation site, fever, subcutaneous edema, lymphadenopathy, hepatomegaly, splenomegaly, and evidence of myocarditis and meningoencephalitis—to asymptomatic and oligosymptomatic situations\(^{(20)}\)\(^{(76)}\)\(^{(91)}\)\(^{(160)}\). The incubation period may vary from 4 to 15 days.
The portal of entry is the ocular conjunctiva (Romaña sign) in 50% of cases and the skin (inoculation chagoma) in 25% of cases; it is unknown in the remaining 25% of cases. However, it is assumed that the ocular route is the most frequently diagnosed as it is easy to recognize, both by the affected individual and the healthcare team. The Romaña sign is essentially characterized by unilateral, painless, elastic edema of the eyelids with satellite lymph node reaction (mainly pre-auricular). Often, the swelling spreads to the ipsilateral side of the face; its degree is variable and can be severe enough to cause total occlusion of the palpebral fissure. The presence of the Romaña sign is an excellent marker for the diagnosis of the acute phase, allowing recognition of many cases.

In terms of the skin, the inoculation chagoma presents as a hard, erythematous, slightly painful nodule surrounded by elastic edema. It is also accompanied by satellite lymph node reaction and, sometimes, ulceration. It can be found on any region of the body, especially exposed parts.

Along with these signs of entry, general symptoms such as fever, malaise, headache, asthenia, and pyrexia appear. Fever is an initial symptom in almost all cases; it usually does not exceed 39°C; although it can be higher, especially in children. The thermal curve is not characteristic; the temperature record may show a continuous or relapsing and intermittent pattern, with significant increases in the late afternoon.

Around the second week of the disease, generalized or localized edema may appear (on the face or lower limbs), regardless of the inoculation site or presence/absence of cardiac failure. Its consistency can be elastic or soft; the elastic is observed only in young children. The pathogenesis of this edema is unclear; several hypotheses have been suggested.

Usually, lymph nodes are slightly to moderately increased in size, isolated, mobile, smooth, painless, and firm. Of those accessible to palpation, lymphadenopathy is most commonly found in the cervical, axillary, and inguinal regions; this manifestation occurs early in disease.

**FIGURE 2.** Flowchart of steps for the laboratory diagnosis of *T. cruzi* infection in suspected cases of chronic Chagas disease.
Hepatomegaly and splenomegaly, alone or in combination, are often observed. In general, the increase in liver and spleen volume is small or moderate, the consistency does not change, the edge and the surface are smooth and, sometimes, there is pain on palpation of the liver. These signs are an early part of the clinical framework (160) (163).

Although less common, other features (neurologic and cutaneous) may be present. Neurologic manifestations refer to signs of meningeoencephalitis (vomiting, agitation, convulsions, opisthotonos, neck stiffness, etc.), which is mainly observed in young children. Meningoencephalitis may present with abnormal cerebrospinal fluid (CSF) findings. The characteristic examination findings are clear CSF, increased cellularity with mild lymphocytosis (<100 cells/mL), low glucose and slightly increased protein levels; it is possible to see trypanostigide forms of T. cruzi after centrifugation of CSF, using specific staining methods (20) (160).

Cutaneous manifestations (rare in Brazil and relatively frequent in Argentina) include rashes (morbilliform, urticariform, and macular) called esquizontipanes and hematogenous chagomas (usually flat formations). These affect skin and subcutaneous tissues without changing their color; are non-adherent to the deep planes; usually painless; and of variable size, from that of a coin to large plaques (160) and, according to Lugones (2001), they are more palpable than visible (162).

The heart presents changes of greater or lesser severity; with a moderate frequency. However, the clinical, radiologic, or ECG manifestations involved are not significant, are not in keeping with the histologic findings, and may even be absent. On the other hand, it is necessary to bear in mind that serial radiologic and electrocardiographic examinations are required in the short-term, seeking signs of cardiac impairment, given the transience of some of the manifestations (20) (160) (164).

The symptoms of acute Chagas myocarditis are essentially identical to those of acute myocarditis of other etiologies, sometimes being masked by other clinical manifestations. Tachycardia is often reported and, generally, not dependent on the degree of temperature increase; in most cases this is observed early. A systolic murmur with functional blow characteristics can be detected in the mitral area. Excitability arrhythmias are only occasionally found. Heart failure, when present, is global; its symptomatology is no different from that related to other etiologies. There may be a degree of arterial hypotension (160) (164).

Already in the first weeks of infection, radiologic and/or electrocardiographic changes, of greater or lesser significance, may be observed. These tests are not always simultaneously abnormal. Practically, if one test result is abnormal it has the same diagnostic value as both being abnormal, as both, individually, reveal disorders with the same frequency. When performed in combination and serially, however, these allow more frequent demonstration of impairment (160) (164). Therefore, the study of cardiac function by echocardiography is indicated in cases with signs and symptoms of myocarditis.

Radiologically, the heart size can be normal or may be mildly, moderately, or severely increased. Cardiomegaly is global, predominantly as a result of enlargement of the ventricles, but in some cases, due to a dilation of only the left ventricle. The exudate in the pericardial cavity contributes to increase the heart shadow (160) (164).

The electrocardiographic changes most frequently encountered are sinus tachycardia, decreased QRS complex voltage, first-degree atrioventricular block (AVB), primary alteration of ventricular repolarization, and increased electrical systole. Arrhythmias, except for first-degree AVB, are only occasionally observed (160) (164).

The chronologic asynchrony between the appearance of echocardiography findings and the occurrence of acute phase is not a reason not to perform this examination. In a study conducted in Venezuela that followed 58 patients in the acute phase, the ECG findings proved to be abnormal in 27 (52%); the most important finding was pericardial effusion, present in 42% of the cases, ranging from mild to moderate intensity in 17 patients and severe in 5; in 11 (21%) patients earlier and/or apical dyskinesia was demonstrated, and in 3 (6%) patients, left ventricular dilation was observed (165). A study of 158 patients with acute Chagas disease in the Amazon (most cases due to oral transmission) revealed the presence of 108 changes, with more than one alteration present in the same individual (91). The main abnormal findings were pericardial effusion (from small to large volume), mitral or tricuspid valve regurgitation, and symmetric hypertrophy of the left ventricle. Occasionally, accentuation of sinus tachycardia was observed, in contrast to the reduction in intensity or disappearance of acute manifestations, including fever; this was previously reported by Chagas (166).

The main nonspecific laboratory abnormalities found are, in order of frequency: anemia, leukopenia, relative lymphocytosis, and an increase, from mild to moderate, of aminotransferases. Also described are thrombocytopenia and, more rarely, thrombocytosis and atypical lymphocytosis (21) (70).

The complete blood count reveals that the total leukocyte count is generally increased, although it may be normal or slightly decreased, with intense lymphocytosis, plasmacytosis, and related neutropenia. A high percentage of atypical lymphocytes and leukocytoid cells appear after the second week of the disease. As the disease evolves to chronicity, these changes disappear, and eosinophilia emerges (20) (21) (39) (160).

**Acute phase of Chagas disease by blood transmission**

In the acute phase of transfusion transmitted Chagas disease, the clinical syndrome is practically identical to that seen when transmitted by triatomines, except for the absence of an inoculation chagoma. Similarly, it must be assessed with complementary examinations. The incubation period can range from 30 to 112 days, being a little longer than that observed in vector-borne transmission, although much shorter periods (such as 8 days) or longer periods (such as 120 days) have been observed (160) (166). Chagas disease must always be considered in cases presenting with a fever of unknown origin, particularly if the patient under investigation has received a transfusion in a remote or hyperendemic region (166).

Fever is the most common symptom, affecting 80–100% of cases. It is often the only symptom found. Lymphadenopathy
and splenomegaly are also frequently observed, while anemia/pallor, perimalleolar and periorbital edema, skin rash, and hepatomegaly appear in <50% of cases.

**Acute phase of Chagas disease by oral transmission**

Oral transmission of Chagas disease has been reported in the Amazon and extra-Amazon regions with peculiar clinical characteristics when compared to vectorial transmission, and sometimes with regional differences. The major clinical differences between the descriptions of past endemic areas and the Amazon region (in which oral transmission prevails) is the clinical presentation of high morbidity, highlighting the regional epidemiological characteristics regarding the occurrence of a more efficient mode of transmission than the vectorial route. Evaluation by non-specific complementary tests is generally similar to that previously seen.

Evidence suggestive of inoculation (Romaña sign and inoculation chagoma)—typical of the disease induced by vector transmission—is rarely described in cases of oral transmission, emphasizing the clinical differential diagnosis between the two forms of transmission and the low frequency of contact between humans and non-domesticated vectors in this region. The incubation period ranges from 3 to 22 days. The main signs and symptoms recorded in acute Chagas disease are, in order of frequency: prolonged fever, headache, pallor, myalgia, skin rash, and edema of the face and lower limbs. The characteristic signs and symptoms of cardiac impairment include tachycardia without fever, palpitations, dyspnea, and chest pain.

Fever is the predominant manifestation in almost all cases. In general, it is characterized by daily high temperatures, from the morning until the late afternoon. This continues for an average of 18 days (range: 3–25 days). In individuals manifesting prolonged fever, the fever is initially high (38.7–39.0°C) and is accompanied by chills and generalized pain. After approximately 12–15 days, it begins to settle; the temperatures range between 37.0°C and 37.8°C, occur daily, and the fever usually disappears in the afternoon.

Pallor (anemia) occurs from the beginning of the febrile syndrome, in particular on days 3–5, and lasts 20–25 days. Myalgia can be intense; it is often described as an acute stabbing pain, or similar to that experienced in dengue fever (generalized and confused with polyarthralgia). Pallor appears as an important sign and is associated with abundant parasitemia. It occurs more frequently than when Chagas disease is caused by vectorial transmission. In general, it presents as a macular rash that is not itchy or painful. It occurs on around days 4–8 of disease, usually affecting the chest, back, lower limbs, and neck, and sparing the face and palmoplantar regions. It is relatively transient and most often goes unnoticed by the patient, found only on physical examination.

The edema of lower limbs generally appears after approximately 12 days of illness and lasts until day 20, coinciding with the disappearance of fever. Invariably, it has the characteristics of inflammatory edema, inelastic and slightly painful. It affects the malleolar region or the entire lower limb. Facial edema can occur in the same period; it has no specific features.

In most cases, acute myocarditis may start slightly before the disappearance of fever, on average from day 15 to 20 of the disease. The main signs and symptoms are dyspnea, palpitations, tachycardia (without fever) and, possibly, precordial chest pain simulating myocardial infarction. Myocarditis is one of the most common complications among patients in the acute phase. ECG and echocardiographic examinations should be performed immediately after diagnosis.

Among patients evaluated in the Amazon, 52.3% had altered ECG findings, consisting of diffuse changes with ventricular repolarization abnormalities (VRA), low voltage of the QRS complex, deviation of the electrical axis to the left, and sinus tachycardia, demonstrating some involvement of the cardiac electrical conduction system of the heart and, more often, diffuse signs of inflammation. Cardiac electrical conduction disorders are more evident in adults. Myopericarditis, evidenced by the presence of VRA and pericardial effusion (seen on ECG and echocardiography, respectively) is important in both adults and children. In outbreaks outside the Amazon Region, a high frequency of pericardial effusion was observed, emphasizing the need for early diagnosis in view of severe evolution if not promptly diagnosed and treated.

Pericardial and/or pleural effusion may occur in more than 50% of cases infected via the oral transmission route, suggesting that pericarditis may be more important than involvement of the cardiac electrical conduction system during the acute phase.

Painful nodules of lower limbs may be associated with edema, and have been invariably recorded in lower limbs of women. The presence of erythema nodosum has been frequently observed in cases of oral transmission. Similar descriptions in children were also recorded in an Argentinian case series, in which some skin and subcutaneous lesions were labeled as hematogenous or metastatic chagomas.

Other manifestations associated with this form of transmission can be present, such as epigastric pain, jaundice of the skin and mucous membranes, hepatomegaly, and lymphadenopathy. Lymphadenopathy is common in children, affecting especially cervical lymph nodes chains; it is not generalized. Splenomegaly and diarrhea are rare events.

It has been reported that 13.3% of cases are severe. Severity is almost always related to manifestations of acute myocarditis and severe gastrointestinal bleeding. The presence of gastrointestinal bleeding might represent the portal of entry, with the inflammatory infiltrate containing amastigotes. There are rare reports of acute meningoencephalitis. Hemorrhage represents a critical scenario of the epidemic that occurred in the City of Navegantes, in the State of Santa Catarina (Southern region of Brazil), providing differential diagnosis for severe diseases that presented a jaundice-hemorrhagic condition, such as leptospirosis, dengue and hantavirosis. The mortality rate was significant among adults aged over 50 years (5.6%).
In just over half (54.1%) of the cases in the Amazon, the diagnosis was confirmed by parasitologic examination. Therefore, in cases with negative parasitologic tests, but with strong clinical suspicion, concomitant serologic testing is recommended, considering in particular their higher sensitivity after the first month of clinical history.

**Specific treatment**

Specific and immediate treatment is indicated in the acute phase, with long-term follow-up to identify the treatment response serologically. For more details, see the specific treatment part of this Consensus.

A case series was reported of 179 patients treated with benznidazole in the acute phase of Chagas disease induced by oral transmission who were followed-up under a research protocol for an average period of 5.6 years. A serologic response to treatment was found in 26.3% of patients, most evident four years after treatment; 2.7% developed a chronic mild-to-moderate cardiomyopathy; and 73.7% had persistently reactive serology, although with a significant decrease in the levels of antibodies.

**Reactivation of Chagas disease**

The management of cases with reactivation of Chagas disease, in the context of immunosuppression, is an important emerging public health problem in all countries where people with chronic Chagas disease reside. In particular, the high congenital transmission rate (>50%) in T. cruzi/HIV coinfected pregnant women is highlighted in the literature. It is advised, therefore, that careful follow-up with clinical assessment and periodic direct search for the parasite be conducted in these women.

Reactivation of chronic Chagas disease in previously infected cases is defined as positivity of the following tests, regardless of the presence of other signs and symptoms:

1. **Parasite detected by direct microscopic examination of blood or biological secretions:** cerebrospinal fluid, pleura, pericardium, ascitic fluid, among others. In terms of sensitivity, parasite search in buffy coat or microhematocrit specimens or sediment from other materials is more sensitive than fresh direct search in biological material or smear. The thick blood smear is more difficult to perform and interpret, and can only be carried out by a trained technician.

2. **Histopathologic examination of tissue lesions** (panniculitis, myocarditis, encephalitis, enteritis, colpitis, among others), with parasite nests within the acute inflammatory infiltrate.

Reactivation of Chagas disease was first reported in patients immunocompromised by hematologic neoplasms. From the late 1980s and early 1990s, cases of reactivation of Chagas disease began to be reported in HIV-infected individuals. In the largest prospective study of T. cruzi/HIV coinfection (n = 53 patients), the rate of reactivation was 20.8%. Considering only followed-up patients and excluding hospitalized patients diagnosed with reactivation, this value was approximately 10%.

In kidney transplant recipients, the rates of transmission (from infected donors) and reactivation (in chronically infected patients submitted to transplantation) are 18.7% and 21.7%, respectively. In the context of heart transplant recipients, a reactivation rate of 26.4–40% was found, and, although use of infected donor hearts is prohibited, this has been reported in the absence of donor screening in non-endemic countries.

In Brazil, in addition to prohibiting the use of infected donor heart and intestine in transplants, it is recommended not to use other transplant organs from donors with infection; exceptional situations should be discussed on a case-by-case basis with specialized teams.

The rate of reactivation in recipients of allogeneic hematopoietic stem cell and liver transplants is 27.3%, and 18.7%, respectively. The use of corticosteroids in association with increased parasitemia and a possible effect with immunosuppressive drugs to treat mesenchymal diseases have been recorded, without a well-defined causal relationship.

**Diagnosing reactivation of Chagas disease**

Clinically, the most common manifestations of reactivation are fever, panniculitis (subcutaneous nodules), myocarditis, meningoencephalitis, stroke, and symptoms such as anorexia, myalgia, malaise, and diarrhea. Case series studies suggest that reactivation associated with HIV infection has a higher morbidity and mortality than other causes of immunosuppression, such as in transplant recipients, where the monitoring of organ recipients is suggested, with high success rate for some groups.

In reactivation associated with HIV infection, meningoencephalitis (74% of the cases), myocarditis (17.1%), and both occurrences (7.9%) have been reported. Oligosymptomatic forms were diagnosed in mothers who gave birth to children with severe Chagas disease and in patients with prolonged fever in long-term follow-up, with good therapeutic responses to benznidazole. Other rare manifestations of reactivation in immunosuppressed patients have been observed, such as erythema nodosum, myelitis, peritonitis, and colpitis.

Complementary examinations, such as computed tomography and/or magnetic resonance imaging of the brain, are fundamental to localize lesions; to differentiate other diagnostic possibilities, such as neurotoxoplasmosis and primary lymphoma of the central nervous system; and to assess whether it is possible to perform a lumbar puncture to obtain CSF for examination to confirm the diagnosis, by recognizing the presence of the parasite (and subsequent cultivation). These tests are more sensitive when performed on CSF than on peripheral blood. CSF examination reveals discrete pleocytosis (with predominance of lymphocytes and monocytes), average or slightly reduced glucose level, and a mild-to-moderately raised protein level.

In reactivation disease, myocarditis is manifested as heart failure induced by arrhythmia, cardiogenic shock, and low response to symptomatic drugs for congestive heart failure. Echocardiography and other complementary examinations to assess cardiac function are indicated.

Diagnosis is facilitated when there is a concomitant increase in parasitemia.
in peripheral blood; however, reactivation can occur in the tissue without systemic repercussions\(^{(49)}\)\(^{(159)}\).

The reactivated form of Chagas disease, in its manifestations of meningoencephalitis and/or myocarditis, is officially considered in Brazil as indicative of severe immunodeficiency in individuals over 13 years, and have been included as AIDS-defining conditions since January 2004\(^{(24)}\).

For epidemiological surveillance purposes, cases of reactivation of Chagas disease are those that present a definitive diagnosis of \(T.\ cruzi\) infection by parasitologic diagnosis through direct investigation of blood or body fluids (CSF, pericardial, or peritoneal fluid), associated with:

- **Meningoencephalitis**: Brain injury image with mass effect (magnetic resonance imaging or computed tomography, with or without injection of contrast medium, showing ring enhancement); and/or
- **Acute myocarditis**: Arrhythmias and/or heart failure diagnosed by electrocardiography and echocardiography.

**Factors associated with the reactivation of Chagas disease**

A CD4\(^+\) T lymphocyte count <200 cells/mm\(^3\) is observed in >80% of cases of reactivation\(^{(40)}\)\(^{(159)}\)\(^{(175)}\). Reactivation is rarely seen in patients with a CD4\(^+\) T lymphocyte count >350 cells per mm\(^3\)\(^{(48)}\)\(^{(49)}\). Available data do not suggest the importance of HIV viral load as prognostic factor for reactivation or as factor related to the severity of the cases. On the other hand, HIV viral load increases have been observed in the context of reactivation\(^{(49)}\)\(^{(159)}\).

Regarding the role of the protozoan, prospective studies have shown the importance of high parasitemia (observed by quantitative methods) as a predictive factor of reactivation (50% of cases). In addition, the participation of different subpopulations of \(T.\ cruzi\) in the genesis of clinical manifestations of reactivation is discussed; this remains controversial in the literature\(^{(48)}\)\(^{(49)}\)\(^{(159)}\).

In heart transplant situations, factors associated with reactivation are the number of rejection episodes, the presence of neoplasms, and tangentially, the use of mycophenolate mofetil\(^{(182)}\).

**Evolution and prognosis of reactivated Chagas disease**

If early treatment is not provided, most patients with AIDS and reactivation of Chagas disease die before or shortly after diagnosis\(^{(159)}\)\(^{(175)}\). Of patients who complete at least 30 days of treatment, about 80% survive\(^{(159)}\). The prognosis in cases of HIV infection and reactivation of Chagas disease was reserved before the advent of highly active antiretroviral therapy (HAART), with an estimated average survival of 10 days\(^{(20)}\). In immunocompromised patients, the severity depends on the degree of immunosuppression, the parasite load and the sensitivity of the isolated parasite to antiparasitic treatment. The outcomes recorded in kidney transplant recipients were favorable, due to monitoring, diagnosis, and early treatment. The clinical picture is quite severe in newborn infants with congenital infection from born to \(T.\ cruzi\)/HIV coinfectcd mothers; the morbidity and mortality is high\(^{(173)}\)\(^{(174)}\)\(^{(175)}\).

**CHRONIC INDETERMINATE FORM OF CHAGAS DISEASE**

The indeterminate chronic form (ICF) of Chagas disease is particularly relevant as it is the most prevalent clinical form of the disease. It has a benign nature and low evolutionary potential in the short and medium term\(^{(19)}\)\(^{(20)}\)\(^{(39)}\)\(^{(188)}\)\(^{(189)}\)\(^{(190)}\)\(^{(191)}\). With the reduced incidence of acute Chagas disease and improved clinical management, the duration of survival of patients with chronic Chagas disease has increased, reflecting a tendency to proportional reduction of ICF with advanced age\(^{(19)}\)\(^{(39)}\)\(^{(189)}\)\(^{(192)}\)\(^{(193)}\)\(^{(194)}\).

On the other hand, since the last Brazilian Consensus in 2005\(^{(20)}\), no applicable strategy of clinical or complementary evaluation to the classic concept of ICF has been proposed.

**Diagnostic criteria**

Patients are considered to have ICF if in a chronic phase of Chagas disease with positive serology and/or positive parasitologic examination for \(T.\ cruzi\)\(^{(188)}\)\(^{(189)}\) but no typical clinical syndrome specific of the disease; conventional ECG results; and normal radiologic examination of the chest, esophagus, and colon\(^{(188)}\). No other complementary examinations in addition to those previously listed, are required to define ICF. The expression *indetermined chronic form* was first used by Carlos Chagas and established by its use\(^{(195)}\)\(^{(196)}\)\(^{(197)}\)\(^{(198)}\). The reference of this term, mainly in the context of scientific research, was consolidated by Brazilian researchers in 1984\(^{(20)}\)\(^{(142)}\)\(^{(188)}\)\(^{(199)}\).

In field and operational studies, for asymptomatic cases with normal physical examination, conventional ECG, and chest and esophagus radiography findings, the term ICF was used without performing radiologic investigation of the colon\(^{(19)}\)\(^{(189)}\)\(^{(190)}\)\(^{(191)}\). This conduct is justified, taking into account the low availability and feasibility of these complementary tests in the real context of the national health systems in endemic regions.

**Treatment and clinical follow-up**

Specific antiparasitic treatment is indicated for all cases with ICF (see specific section on antiparasitic treatment in this Consensus\(^{(20)}\)\(^{(39)}\))

While the ECG finding is normal, the prognosis of cases with ICF of Chagas disease is similar to the general population, since serial repetition of these examinations can detect evolution to a cardiac form of Chagas disease\(^{(20)}\)\(^{(189)}\)\(^{(192)}\)\(^{(197)}\). Based on this key concept, is not recommended performing other routine complementary tests while the ECG finding is normal\(^{(19)}\)\(^{(20)}\)\(^{(189)}\)\(^{(190)}\)\(^{(191)}\).

Cases with ICF should be advised not to donate blood, and in principle, tissues or organs\(^{(20)}\)\(^{(39)}\). Due to the benign nature of ICF, the (still common) practice of requesting specific serologic exams for Chagas disease in pre-employment assessments for labor purposes and periodic examinations conducted by institutions and/or public or private companies, is not justified\(^{(192)}\)\(^{(200)}\)\(^{(201)}\)\(^{(202)}\)\(^{(203)}\).

As for other complementary tests, these may be ordered according to the specific labor activities that the individual will perform. Similarly, in these cases, temporary or permanent removal from work activity is not recommended\(^{(192)}\)\(^{(200)}\)\(^{(201)}\)\(^{(202)}\)\(^{(203)}\).

Engaging in usual, regular physical activity is not contraindicated in patients with ICF. For professional activities
involving individual and collective risk that may require high demand of physical effort and/or psychological stress, the adoption of a complementary specific cardiologic evaluation is recommended. There is no restriction of sexual activity for patients with ICF (192) (200) (201) (202).

ICF carriers must be assessed annually, preferably in the primary healthcare services, by means of medical evaluation and conventional ECG (20) (202). If disease evolution is observed, these cases should be referred to more specialized healthcare services with the aim to provide the patient with a more precise direction (to define the appropriate investigation and treatment interventions) for the integrated management with primary health care teams (up- and down-referral).

Indeterminate chronic form does not interfere with the management of associated diseases and does not justify neglecting the follow-up and treatment of comorbidities that may possibly be present (199) (197) (203) (204). If immunodeficiency (acquired or induced) occurs in patients with ICF, special attention should be given to possible reactivation of Chagas disease (see specific section on reactivation of Chagas disease) (20) (149) (159). The surgical risk for the patient with ICF does not differ from that observed in the general, uninfected population (20) (194) (200) (202) (203).

Regarding pregnant women infected with T. cruzi with this classification, attention should be paid to the possibility of vertical transmission, and opportunities should be created for adequate assessment of the newborn infant (see specific section on vertical transmission (20) (19) (202)). Women with ICF should not restrict breastfeeding, except in the presence of fissures and/or bleeding of the nipple/areola (20) (202).

It is emphasized that health professionals should avoid the adoption of any practice that may induce stigma or prejudice. Health services should provide space and resources for advice, clarification, and orientation of the population regarding the characteristics of this form of disease. It is strongly recommended that this approach is led by a multidisciplinary team (20) (189) (190) (191) (193) (202).

CARDIAC FORMS OF CHAGAS DISEASE

Chronic Chagas heart disease is the most prevalent symptomatic clinical form, responsible for the high burden of morbidity and mortality, with great impact on social and medical systems (4) (19) (20). Recently, the notion has been considered that the pathogenesis of myocardial injury in the chronic phase of Chagas disease depends mainly on parasite persistence and the unfavorable immune system response to this incessant infectious stimulus (20) (206) (207) (208). Among the most peculiar characteristics of chronic Chagas heart disease, the following are highlighted: Its inflammatory and intensely fibrosing nature, presence of complex ventricular arrhythmias in association with disturbances in atrioventricular and intraventricular formation and conduction of the electrical stimulus, high incidence of sudden death and thromboembolic events, and right ventricular dysfunction and ventricular aneurysms (209) (210) (211) (212) (213) (214).

Acute Chagas disease and consequently, acute Chagas myocarditis, was previously restricted to vectorial transmission of Chagas disease, now rare in Brazil. However, their epidemiological importance in Brazil and other countries, endemic and non-endemic, has expanded due to reactivation of the disease in immunocompromised individuals, favored by the spread and chronicity of HIV infection, as well as by the increased access to organ transplantation. In addition to this scenario, there is the challenge posed by Chagas disease in the Amazon, attributed to oral transmission. As a result of changing the epidemiological profile of disease transmission, cases of acute Chagas heart disease, which currently present clinical and specific epidemiological aspects, has again increased (76) (91) (92) (169) (170).

The importance of the composition of a resolute healthcare network for people with Chagas disease, in particular the role of primary care as the first contact and provider of secondary prevention of the disease, is emphasized. From the recognition of the context of clinical cases, this primary health care network should be integrated into the matrix/reference network for more complex situations.

Acute Chagas heart disease

1. Acute Chagas myocarditis associated with oral transmission

In Amazonía Legal, especially, a systematic record exists of cases with the acute form of Chagas disease, occurring in sporadically, in outbreaks, or in micro epidemics, where the main form of transmission is oral, involving food contaminated with triatome feces (216). The clinical presentation differs from classical acute Chagas myocarditis (vectorial) mainly by the absence of evidence of a portal of entry (inoculation chagoma), the involvement of community groups or families in outbreaks, and having no identified predilection for a specific age group or for disease severity (91) (169).

Clinical manifestations of acute disease associated with oral transmission are variable, ranging from asymptomatic cases up to those developing severe heart failure, cardiogenic shock, and even death. It can also manifest as a nonspecific infectious syndrome, with prolonged fever (usually for more than three weeks). The manifestations of acute myocarditis are described in Figure 3 (216).

Laboratory and radiologic examinations

Chest radiography may be normal or may show varying degrees of cardiomegaly and pleural effusion. The ECG readings are altered in most acute cases. The main ECG findings are non-specific changes in ventricular repolarization, followed by prolonged QTc interval, left atrial overload, low voltage of QRS complex, AVB, bundle branch block, sinus tachycardia, and atrial fibrillation (216). ECG can detect pericardial effusion, atrioventricular valve regurgitation, increase in the size of the heart chambers, presence of thrombi, and changes in left ventricular systolic function.

Treatment

Pharmacological management of acute Chagas myocarditis is even recommended for the treatment of heart failure in patients with acute myocarditis of other etiologies: Routine use
of a combination of three types of drugs [diuretics, angiotensin converting enzyme inhibitors (ACEI) or angiotensin II AT1 receptor blockers (ARBs), and beta-blockers] is always combined with specific treatment of *T. cruzi* infection (using benznidazole or nifurtimox)(203).

The natural history of the acute phase of Chagas myocarditis, induced by oral transmission of Chagas disease, is still not fully known. The most severe cases, with ventricular dysfunction, pericardial effusion, and atrial fibrillation, tend to have worse outcomes in the acute phase, requiring treatment in an intensive care environment.

2. Acute Chagas myocarditis in immunocompromised individuals

Individuals infected with *T. cruzi*, when exposed to immunosuppressive agents or who have other concomitant diseases such as cancer and other infections, particularly HIV infection, may experience reactivation of Chagas disease. The frequency of this reactivation is not fully known. However, in a prospective study in which cases were evaluated pre- and post-antiretroviral therapy, the occurrence of reactivation was observed in 20% of the cases(48).

The heart appears to be involved in approximately 30–40% of cases of reactivation of Chagas disease in individuals coinfected with HIV. However, the incidence of myocarditis alone does not seem to be common. Cardiac involvement is usually characterized by acute myocarditis, with diffuse or focal cardiac involvement. Clinically, it is characterized by signs and symptoms of heart failure (tachycardia, edema, hepatomegaly) or severe arrhythmias. In some cases, only electrocardiographic changes are observed; in others, myocarditis is confirmed only by histopathologic examination of endomyocardial biopsy material. Anatomopathological examination reveals acute myocarditis with intense inflammatory infiltrate, cardiac fiber injury with focal necrosis, and large numbers of amastigote forms of the parasite(217). In cases where there was previous cardiac involvement (for chronic Chagas heart disease), there may be overlapping reactivation Chagas disease cardiomyopathy and decompensation of preexisting Chagas heart disease. Determining if the clinical picture is solely due to reactivation of Chagas disease, myocarditis associated with HIV itself, or the overlap of the two conditions, is complex(218). Acute myocarditis due to the reactivation of *T. cruzi* infection might be also confused with the natural progression and worsening of chronic Chagas heart disease. The differential diagnosis between these two conditions is important in terms of deciding whether or not to administer the specific treatment of *T. cruzi* infection itself, as many patients do not tolerate the medication. High levels of parasitemia are indicative of reactivation. However, there is no record of patients who developed low parasitemia(219).

**Diagnosis**

Confirmation of reactivated Chagas disease in patients with immunosuppression is performed by parasite detection in peripheral blood and other body fluids (cerebrospinal fluid, cavity effusions, etc.), by direct observation methods, or in sites of organ damage (heart, skin, etc.) in individuals with co-infection(48).

**Treatment**

There are no specific treatment measures for this group. Treatment should follow evidence-based recommendations developed for the treatment of heart failure in general. Acute decompensated heart failure, resulting from the hemodynamic effects of heart failure, is associated with neurohumoral activation and symptoms of congestion and/or low cardiac output. Early recognition of this condition and the implementation of appropriate therapeutic measures for heart failure, associated with the specific treatment for *T. cruzi* infection, can reduce the high rate of mortality observed in cases of myocarditis induced by reactivation of Chagas disease in immunocompromised patients.

**Chronic Chagas cardiomyopathy**

**Definition**

Chronic Chagas cardiomyopathy (CCC) is defined as the presence of ECG changes suggestive of cardiac involvement typical of Chagas disease in symptomatic and asymptomatic
individuals. Since ECG examination is easily accessible and has high sensitivity and specificity (around 90%), the modified conventional ECG is used, *a priori*, to diagnose CCC in people chronically infected with *T. cruzi* (20). Cardiac involvement in the chronic phase of Chagas disease includes a wide spectrum of manifestations, ranging from the presence of clinically unapparent abnormalities to severe forms, such as end-stage heart failure, thromboembolic complications, refractory ventricular arrhythmias, and sudden death.

Taking as a reference only conventional 12-lead ECG, it is possible to establish, in a simplified way, the degree of cardiac involvement in routine evaluation of the patient with chronic Chagas disease, as shown in the following flowchart (Figure 4) (20).

Cases with nonspecific ECG changes would not be diagnosed with CCC, but they should be re-evaluated according to the concomitant symptoms and changes found in additional tests.

**Chronic Chagas cardiomyopathy staging**

Presence of severe heart failure [functional class III and IV of the New York Heart Association (NYHA)] and left ventricular global systolic dysfunction are the most important prognostic factors in CCC (20) (21) (22) (23).

Echocardiography allows evaluation of both the overall and segmental myocardial function. It can also identify important

---

**FIGURE 4.** Algorithm for evaluation of individuals with Chagas disease using conventional ECG.

IRBB: incomplete right bundle-branch block; LAFB: Upper anterior fascicular block; AVB: Atrioventricular block; ST-T: ST-T segment; RBBB: Complete right bundle branch block; VES: Ventricular extrasystoles; T: T-wave; HR: Heart rate; NSVT: Non-sustained ventricular tachycardia; AF: Atrial fibrillation; CAVB: Complete atrioventricular block; LBBB: Left bundle branch block; LVEF: Left ventricular ejection fraction; CHF: Congestive heart failure; ECO: Echocardiogram; HF: Heart failure.
markers for staging the disease, such as size of chambers, alterations in segmental mobility, and presence of aneurysms and mural thrombosis. For these reasons, and because it is a noninvasive and inexpensive examination, echocardiography is of great value for the initial staging of CCC, in association with the functional NYHA classification. From a prognostic and therapeutic point of view, it is possible to identify five different subgroups of CCC, as shown in Figure 5 (223) (224) (225) (226).

In addition to the impairment of myocardial function, rhythm disturbances and electrical impulse conduction abnormalities are significant changes in CCC; in some cases, arrhythmias may occur alone, without global ventricular dysfunction, or only with small regional dyskinesias (227) (228) (229) (230) (231) (232). If possible, it is obligatory to perform a 24-hour electrocardiogram (Holter) in all cases with suspected arrhythmia. Less frequently, an ergometric test may be used instead of a Holter, to show ventricular arrhythmias during standardized physical effort. When possible, both tests should be performed, as they are complementary in the evaluation of these cases.

There is also a group of cases in the chronic phase of Chagas disease (not included in the classification above) that shows normal ECG findings, but presents changes—usually discrete and not associated with an increased risk of death—in other cardiac tests, such as echocardiography, myocardial scintigraphy, and magnetic resonance imaging. However, since the risk of progression of these cases to the cardiac form, classically defined, appears to be higher than that of cases of Chagas disease without these changes, they should receive regular clinical follow-up (231) (232).

**Chronic Chagas cardiomyopathy prognosis**

In a systematic review of studies that used multivariate analysis to evaluate the prognosis of patients with CCC (233), four independent prognostic variables were identified: functional class III/IV of NYHA, cardiomegaly on chest radiograph, left ventricular systolic dysfunction, and non-sustained ventricular tachycardia (NSVT) on Holter examination. By using these four variables in an integrated manner, it was possible to develop an algorithm capable of stratifying the prognosis of cases with CCC (Figure 6).

<table>
<thead>
<tr>
<th>Stage</th>
<th>Electrocardiogram</th>
<th>Echocardiogram</th>
<th>Heart failure</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Altered</td>
<td>Normal</td>
<td>Absent</td>
</tr>
<tr>
<td>B1</td>
<td>Altered</td>
<td>Altered, LVEF ≥45%</td>
<td>Absent</td>
</tr>
<tr>
<td>B2</td>
<td>Altered</td>
<td>Altered, LVEF &lt;45%</td>
<td>Absent</td>
</tr>
<tr>
<td>C</td>
<td>Altered</td>
<td>Altered</td>
<td>Compensated</td>
</tr>
<tr>
<td>D</td>
<td>Altered</td>
<td>Altered</td>
<td>Refractory</td>
</tr>
</tbody>
</table>

**Source:** Adapted Xavier SS et al. 2005 (223)

It is important to highlight that the presence of functional class III or IV NYHA, *per se*, identifies high-risk cases, since virtually all these cases have ventricular systolic dysfunction (evident on echocardiography) and NSVT (detected by Holter). The combination of ventricular dysfunction with NSVT (regardless of functional class) already identifies a group with a risk about 15 times higher than cases without these two variables (222) (233) (234).

**Clinical manifestations of chronic Chagas cardiomyopathy**

Clinical manifestations of CCC can be grouped into three syndromes: arrhythmic, heart failure, and thromboembolic. These syndromes may occur alone or in combination in the same case, and may be associated with megaesophagus and/or megacolon (235) (236).

1. **Arrhythmic syndrome**

**Ventricular arrhythmias**

Ventricular arrhythmias are common in Chagas disease and are of various types, including ventricular extrasystoles, isolated and in pairs; NSVT; sustained ventricular tachycardia (SVT); and ventricular fibrillation (237). These commonly present in combination. Clinically, they present as palpitations, lipoptymia, syncope, and sudden death (237) (238). Syncope and other symptoms of low output in cases with CCC should be quickly investigated, due to the risk of complex ventricular arrhythmias and sudden death.

Ventricular extrasystole is the most common arrhythmia. Initially uncommon, monomorphic, and isolated, it becomes polymorphic and repetitive with the evolution of the disease. Holter and ergometric tests are the methods of choice for the detection of ventricular arrhythmias and, when possible, should be performed in all cases with CCC, regardless of the presence of symptoms (42). Ventricular tachycardia may be sustained (duration ≥30 sec or shorter than this when interrupted electrically or pharmacologically) or non-sustained (3 heartbeats or more and a duration <30 sec).

NSVT relates to the degree of ventricular dysfunction, occurring in about 40% of cases with CCC associated with
Patients with serological exam reactive to T. cruzi

Conventional electrocardiogram

Abnormal

FC III/IV (NYHA)

FC I/II (NYHA)

Chest radiograph

Normal

Echocardiogram

Normal

Holter

Without NSVT

Low risk

Without NSVT

Intermediate risk

With NSVT

High risk

Reduced LVEF

Echocardiogram

Normal

Holter

Without NSVT

Low risk

Without NSVT

Intermediate risk

With NSVT

High risk

Reduced LVEF

Holter

Without NSVT

Low risk

Without NSVT

Intermediate risk

With NSVT

High risk

Source: Adapted from Rassi Jr. A et al., 2007(233)

FC: Functional class; NYHA: New York Heart Association; LVEF: Left ventricular ejection fraction; NSVT: Non-sustained ventricular tachycardia.

FIGURE 6. Algorithm for risk stratification in patients with chronic Chagas cardiomyopathy.

regional changes in ventricular contractility, and in 90% of cases with left ventricular global systolic dysfunction associated with heart failure(222)(237). This may occur in cases with normal ventricular function(234). There is evidence that the frequency of ventricular arrhythmias is increased in heart failure due to Chagas disease, compared with other etiologies(239). SVT can occur spontaneously or be reproduced by electrophysiological study, which is indicated for cases whose history suggests this hypothetical diagnosis but for whom non-invasive methods have failed to confirm the diagnosis(240). Ventricular fibrillation is the leading cause of sudden death in CCC. It occurs more frequently in patients who have had previous episodes of ventricular tachycardia, although it may also be the first manifestation of the disease or its terminal event, especially in cases with severe ventricular dysfunction and chronic cardiac insufficiency (CCI)(238).

Treatment of ventricular arrhythmias

Treatment of the arrhythmia has two main objectives: to control symptoms and to prevent sudden death(238). Simple ventricular arrhythmias (isolated and monomorphic ventricular extrasystoles) are not associated with an increased risk of death. Hence, they should be treated only if causing symptoms of functional impairment. If needed, suitable treatment consists of normal doses of amiodarone, sotalol, or beta blockers.

In cases of life-threatening arrhythmias (NSVT, SVT, and ventricular fibrillation), the most effective antiarrhythmic,
and the safer to use, is amiodarone\(^{(241)}\). It is necessary to pay attention to the side effects of this drug — that are generally related to the cumulative dose — which can trigger serious bradyarrhythmias. In such cases, evaluation of the need for permanent pacemaker implantation is required. Extra-cardiac toxicity, thyroid dysfunction, and skin abnormalities are not uncommon, while severe pulmonary toxicity is rare\(^{(242)}\). Periodic assessment of the thyroid function is recommended in patients treated with amiodarone.

In turn, cases with SVT (a relatively common condition) and those who recover from cardiorespiratory arrest in an out-of-hospital environment (a much rarer condition) are at high risk of death and deserve a rigorous evaluation. For these cases, there are not only antiarrhythmic drugs, but also arrhythmic focus catheter ablation techniques (or, rarely, surgical techniques) and, especially, implantable cardioverter-defibrillator (ICD) devices\(^{(237)}\).

The treatment of ventricular arrhythmias in CCC is described briefly in Figure 7. Drug treatment and ICD recommendations are described in Figure 8 and Figure 9, respectively.

**Supraventricular arrhythmias**

Atrial fibrillation is the sustained supraventricular arrhythmia more frequently observed in CCC, found in 4–12% of cases\(^{(229)}\). It tends to manifest later on and is often associated with marked cardiomegaly. Treatment consists in controlling the ventricular rate, which can be obtained through the use of drugs prolonging the refractory period of the atrioventricular node. If there is an associated heart failure, preference is given to digitalis and beta-blockers (metoprolol succinate, carvedilol, or bisoprolol). If ventricular function is normal, the use of conventional beta-blockers (propranolol and atenolol) or calcium channel blockers (verapamil and diltiazem) is recommended for the initial control of the heart rate, with subsequent evaluation of the possible need for electrical or pharmacologic cardioversion. Anticoagulation is indicated when atrial fibrillation is associated with heart failure and cardiomegaly, CHA\(_2\)DS\(_2\)VASc score \(\geq 2\), or when there is evidence of intracavitary thrombosis or previous embolic episodes. The drug of choice is warfarin, in a dose sufficient to maintain the international normalization factor (INR) between 2 and 3\(^{(243)}\).

**Bradyarrhythmias**

The treatment of bradyarrhythmias in CCC does not differ from that recommended for other types of cardiomyopathies. It consists of the implantation of a permanent cardiac pacemaker in symptomatic or high-risk blockage cases. These recommendations are well defined in the Brazilian Guidelines for Implantable Cardiac Electronic Devices, published in 2007\(^{(244)}\).

The association between disorders of the heart conduction system and frequent and complex arrhythmias is common in CCC. In these cases, the effective pharmacologic antiarrhythmic

---

**FIGURE 7.** Interventions for treatment of ventricular arrhythmias in Chagas disease.
Recommendation class | Indication | Level of evidence
--- | --- | ---
I | Individualized antiarrhythmic medication for cases of isolated or paired VES<sup>a</sup>, symptomatic | C
I | Amiodarone for cases with symptomatic NSVT<sup>b</sup> | C
I | Amiodarone for recovered cases of cardiac arrest or with stable SVT<sup>c</sup> and EF<sup>d</sup> >35% | B
I | Amiodarone for cases with stable SVT<sup>c</sup> | C
I | Amiodarone for reduction of shock in appropriate cases with ICD<sup>e</sup> | C
IIa | Amiodarone for cases with asymptomatic NSVT<sup>b</sup> and LV<sup>f</sup> dysfunction | B
IIb | Routine amiodarone for cases with symptomatic SVT<sup>c</sup> who were treated with ICD<sup>e</sup> | C
IIb | Amiodarone for cases with asymptomatic NSVT and normal LV<sup>f</sup> function | C
IIb | Amiodarone for recovered cases of cardiac arrest or with unstable SVT<sup>c</sup> and EF<sup>d</sup> ≤ 35% | C
III | Antiarrhythmic drugs for cases with asymptomatic isolated or paired VES<sup>a</sup> | C

<sup>a</sup>VES: Ventricular extrasystole; <sup>b</sup>NSVT: Non-sustained ventricular tachycardia; <sup>c</sup>SVT: Sustained ventricular tachycardia; <sup>d</sup>EF: Ejection fraction; <sup>e</sup>ICD: Implantable cardioverter defibrillator; <sup>f</sup>LV: Left ventricle.

**FIGURE 8.** Recommendations and levels of evidence for the use of antiarrhythmic drugs in the treatment of ventricular arrhythmias in chronic Chagas cardiomyopathy.

Recommendation class | Indication | Level of evidence
--- | --- | ---
I | ICD<sup>a</sup> for recovered cases of cardiac arrest or with unstable SVT<sup>d</sup>, regardless of EF<sup>c</sup> | C
I | ICD<sup>a</sup> for cases with stable SVT<sup>d</sup> and LVEF<sup>g</sup> ≤35% | B
IIa | ICD<sup>a</sup> for cases with stable SVT<sup>d</sup> and EF<sup>c</sup> >35% | C
IIb | ICD<sup>a</sup> for cases with NSVT<sup>e</sup> and EF<sup>c</sup> ≤35% | C

<sup>a</sup>ICD: Implantable cardioverter defibrillators; <sup>d</sup>SVT: Sustained ventricular tachycardia; <sup>c</sup>EF: Ejection fraction; <sup>g</sup>LVEF: Left ventricular ejection fraction; <sup>e</sup>NSVT: Non-sustained ventricular tachycardia.

**FIGURE 9.** Class of recommendation and levels of evidence for the use of CDI in the treatment of ventricular arrhythmias in chronic Chagas cardiomyopathy.

drug therapy may require permanent pacemaker implantation in order to prevent a possible AVB or high-risk bradyarrhythmia induced by antiarrhythmic. The choice of the pacing mode is, to date, the subject of controversy in the literature.

**II - Heart failure syndrome**

In the acute phase of Chagas disease, heart failure (HF) can occur due to severe myocarditis. Although the outcome may be death, in most cases, it is reversible, progressing to the indeterminate form of the disease. Approximately 30–40% of cases with the indeterminate form will progress to the heart form of the disease, usually after decades of evolution<sup>42</sup>,<sup>236</sup>.

The chronic heart form, characterized by the gradual appearance of electrocardiographic changes, is caused by the slow but steady destruction of myocardial fibers, caused by a continuous chronic inflammatory process, with intense reparative fibrosis and progressive ventricular remodeling<sup>208</sup>.

In the early chronic stage of the cardiomyopathy, it is not uncommon that only the ECG findings are changed; individuals remain asymptomatic, capable of performing unrestricted, sometimes even extreme, physical activity. Therefore, they are included into NYHA class I<sup>227</sup>,<sup>230</sup>,<sup>239</sup>. Rarely, in this group of cases, the first and only manifestation of CCC is sudden death<sup>245</sup>. In most cases, a progressive reduction in the ability to perform physical activities is observed, followed by deterioration of contractile function of the left ventricle, initially in the form of regional dyskinesia and diastolic dysfunction, followed by a drop in the global systolic function of the chamber, due to several associated factors, such as the progressive destruction of cardiomyocytes, microvascular changes, the breakdown of the muscle structure and fibrosis<sup>208</sup>,<sup>236</sup>,<sup>247</sup>. It is not common for cases with Chagas disease in the chronic phase to present with acute pulmonary edema. There are also many cases with CCC in whom the initial manifestations of heart failure are fatigue and edema, with minimal dyspnea<sup>246</sup>,<sup>247</sup>. In these cases there is the early involvement of the right ventricle, with impaired systolic function. In the more advanced stages of the disease, when heart failure is completely manifested, predominant symptoms and signs of systemic congestion may arise, but with little significant pulmonary congestion. Chest teleradiography shows a marked cardiomegaly, with little engorgement of the pulmonary vasculature. It is important to identify early signs of left ventricular systolic failure, since the treatment at this early stage could, in theory, delay the deterioration of contractile cardiac function<sup>42</sup>,<sup>227</sup>. In addition, left ventricular
systolic dysfunction is also the most important indicator of risk in the chronic phase of Chagas disease-induced cardiomyopathy(222)(224)(225).

Treatment of heart failure in cases with chronic Chagas cardiomyopathy

Treatment of heart failure aims to relieve the symptoms, slow the progression of ventricular dysfunction, and prolong survival(20). Treatment of heart failure due to Chagas disease, as well as other etiologies, should be comprehensive, starting with general measures, removing the factors that contribute to worsen heart failure, and treating the underlying syndrome(248)(249).

General measures

1. Diet: Obesity should be corrected and the ideal weight should be maintained, avoiding inadequate intake of sodium chloride and, in specific cases, alleviating the symptoms of dysphagia and constipation.

2. Controlling water retention: A balanced intake of salt (3–4g/day of sodium chloride [mild to moderate disease], and ≤2g/day [severe disease]). A simple and reliable way to evaluate water retention is daily assessment of body weight. Variations higher than 1kg/day are indicative of fluid retention. A gain of rapid and constant weight (1kg/day) is an indication that heart failure is worsening. People affected should be encouraged to monitor their weight daily in the morning (after urinating and fasting). In severe heart failure with hypervolemia and/or hyponatremia, fluid restriction may be required.

3. Control of aggravating factors: The intake of alcohol and the use of anti-inflammatories should be avoided. Arterial hypertension, cardiac arrhythmias, anemia, or thyroid function disorders should be controlled. Other comorbidities may contribute to worsening heart failure, such as coronary artery disease and diabetes mellitus.

4. Individualization of the recommendations for rest or physical activity, in accordance with the degree of heart failure and the patient’s age.

5. Vaccination against influenza (annually) and pneumococcal 23-valent (a booster after five years for those cases with compromised immune systems, or adults over the age of 60 years who received the first dose before 65 years old) should be provided.

Drug treatment of heart failure caused by Chagas disease

At the time of finalization of this Consensus, the results of the BENEFIT study were published. This was a double-blinded, multicenter, clinical trial that tested the hypothesis that trypanosomicide treatment with benznidazole, compared to placebo, could modify the prognosis of patients with Chagas disease chronic cardiomyopathy(250) (251). Overall, 2,854 patients were evaluated. They were randomly assigned to receive benznidazole or placebo for up to 80 days, with a mean follow-up period of 5.4 years. A reduction in the parasite load was observed with benznidazole, but it had no effect on the progression of heart disease (estimated by complications occurring during clinical follow-up, such as death, ventricular arrhythmias, need for implanted devices, thromboembolic events, heart failure, or heart transplantation): 27.5% of patients in the benznidazole group and 29.1% of patients in the placebo group experienced complications, hazard ratio (HR) = 0.93; 95% CI, 0.81–1.07; p = 0.31)(251). This result differed from the results of previous observational studies(252)(253). These results shift the focus of attention to treatment (of patients with CCC) with conventional anti-failure methods derived from other cardiomyopathies(254).

Long-term treatment of heart failure is usually based on a combination of the following classes of drugs: diuretics; ACE inhibitors or ARBs; adrenergic blockers (BB); and the aldosterone antagonists(254). Positive inotropic drugs (such as catecholamine or milrinone) have limited applicability in intensive care settings, and slightly prolonged cardiac decompensation. Digitalis, however, may be administered chronically in individuals without bradyarrhythmias to alleviate their symptoms and prevent cardiac decompensation and hospital admission, especially in patients with atrial fibrillation with rapid ventricular response(255).

Loop and thiazide diuretics may be used separately or in combination to relieve symptoms and signs of pulmonary and systemic venous congestion. When it is imperative to use high doses of loop diuretics, such as furosemide, one should be aware of the higher likelihood of hypokalemia and/or hyponatremia, which, in turn, can exacerbate or trigger serious arrhythmias(256).

In CCC with heart failure or asymptomatic systolic dysfunction (ejection fraction <45%), the chronic administration of ACE inhibitors is indicated to reduce the morbidity, mortality or AR, when patients are intolerant to ACE inhibitors(256). With the same purpose, the use of spironolactone is recommended in cases with heart failure, in NYHA functional classes II–IV(248). Furthermore, it is thought that aldosterone antagonists can provide additional benefits when used in patients with CCC because of their antifibrotic properties (demonstrated in experimental studies). The combination of hydralazine and nitrates is a recommended alternative to treat cases in functional NYHA class II–III if a contraindication to the use of ACE inhibitors or ARBs (e.g., progressive renal failure or hyperkalemia) exists(248).

The use of adrenergic beta-blockers in addition to blockade of the renin-angiotensin-aldosterone system (with ACE inhibitors or ARBs) is recommended to reduce the morbidity and mortality associated with the disease. It is assumed that these drugs can help to prevent the worsening of the ventricular remodeling, development of malignant arrhythmias, and sudden death(257)(258). However, the use of an optimized dose of adrenergic blockers may be hindered in CCC, given bradyarrhythmias and frequent need of amiodarone administration for the treatment of tachyarrhythmias. To date, there is no consensus among experts about which drug, beta-blockers or amiodarone, should be prioritized in the management of Chagas disease.

Complementary forms of treatment for heart failure caused by Chagas disease

Heart transplantation is the treatment of choice for end-stage heart failure, despite the many limitations of medical and social nature that embargo its broader use in cases of Chagas disease.
The results obtained in a selected series of cases, compared with those observed in cases with heart failure due to other etiologies, show an even more favorable evolution in transplanted cases with Chagas disease. Among other possible factors, this is possibly due to the fact that, in this series, cases with Chagas disease were younger and had fewer comorbidities. There is no definitive indication for prophylaxis; however, active control is recommended for possible reactivation of the disease, especially in the first year after transplantation, when immunosuppression is more pronounced. Benznidazole, in a dose of 5mg/kg/day, should be promptly started and maintained for 60 days to obtain a good clinical outcome. There is no evidence to support routine indication of cardiac resynchronization therapy (CRT). In addition to the fact that there are no controlled studies exploring in a scientific and valid manner this therapeutic possibility, in CCC the complete block of the right bundle branch is frequent and, in the presence of this condition, evidence of the benefits of CRT, even in other etiologies of heart failure, are even more scarce and less convincing.

There is no evidence to support routine indication of cardiac resynchronization therapy (CRT). In addition to the fact that there are no controlled studies exploring in a scientific and valid manner this therapeutic possibility, in CCC the complete block of the right bundle branch is frequent and, in the presence of this condition, evidence of the benefits of CRT, even in other etiologies of heart failure, are even more scarce and less convincing.

Cell implant-based therapies were recently explored in a single study, properly controlled. The results did not show any benefit, even on surrogate outcomes such as the left ventricular ejection fraction.

The recommendation classes and levels of evidence for treating heart failure in patients with CCC are summarized in Figure 10.

**III - Thromboembolic syndrome**

Systemic and pulmonary thromboembolic events are common in CCC. Pulmonary thromboembolism is certainly under-diagnosed when considering the results of postmortem studies. Embolic events are favored by the combination of several factors. The most important are venous stasis, reduction in cardiac output, and intracardiac mural thrombus, favored by localized ventricular dyskinesias (e.g., apical aneurysm), dilation of the cardiac chambers, and atrial fibrillation in more advanced stages of the cardiomyopathy.

**Prevention of thromboembolic complications in chronic Chagas cardiomyopathy**

Embolic accidents in the central nervous system are the most severe form of thromboembolism in CCC and contribute to the high morbidity and mortality of the disease. The scientific basis for antithrombotic treatment in CCC is based on findings of a prospective cohort study of 1,043 cases, which assessed the risk and defined prevention strategies for cardioembolic stroke (CES). In that study, the incidence of CES was 3.0% (or 0.56% per year), and the multivariate statistical analysis allowed the composition of a risk score for CES occurrence: The presence of left ventricular systolic dysfunction contributed 2 points; while presence of apical aneurysm, primary alteration of ventricular repolarization in ECG, and age over 48 years contributed 1 point each. Warfarin may be indicated for cases with 4–5 points (in this subgroup, the incidence of stroke is 4.4% vs. 2.0% of severe bleeding per year). In the subgroup with a score of 3 points, embolism rates and bleeding with oral anticoagulants are equivalent, and acetylsalicylic acid or warfarin may be indicated. In cases with 2 points, with low incidence of CES (1.22% per year), acetylsalicylic acid or no prophylaxis is recommended. Cases with 0–1 point, with an event incidence close to zero, do not require prophylaxis. Obviously, in cases with intracavitary thrombus, atrial fibrillation associated with CHA2DS2-VASc score ≥2, and previous embolic accident, the use

<table>
<thead>
<tr>
<th>Recommendation class</th>
<th>Indication</th>
<th>Level of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>ACEI(^a) or ARB(^b) (in intolerant to ACEI(^b)) in cases with t reduced LVEF(^c), symptomatic or no</td>
<td>C</td>
</tr>
<tr>
<td>llb</td>
<td>ACEI(^b) or ARB(^b) (in intolerant to ACEI(^b)) in cases with normal LVEF(^c) and segmental dysfunction of LVd</td>
<td>C</td>
</tr>
<tr>
<td>I</td>
<td>Spironolactone in cases with LVEF(^c) &lt;35% and FC(^e) III/IV of the NYHA</td>
<td>B</td>
</tr>
<tr>
<td>Ila</td>
<td>Spironolactone in cases with LVEF(^c) &lt;35% and FC(^e) II of the NYHA</td>
<td>C</td>
</tr>
<tr>
<td>I</td>
<td>Carvedilol, metoprolol succinate or bisoprolol in cases with reduced LVEF(^c),</td>
<td>C</td>
</tr>
<tr>
<td></td>
<td>symptomatic or not, and with HR(^f)&gt;55 bpm</td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>Loop diuretics in symptomatic cases, with signs and symptoms of congestion</td>
<td>C</td>
</tr>
<tr>
<td>I</td>
<td>Combination of hydrochlorothiazide in cases resistant to loop diuretics</td>
<td>C</td>
</tr>
<tr>
<td>Ila</td>
<td>Digoxin in symptomatic cases with reduced LVEF(^c) (&lt;45%) and sinus rhythm or AF(^h),</td>
<td>C</td>
</tr>
<tr>
<td></td>
<td>despite optimized therapy with ACEI(^b) and BB(^i)</td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>Heart transplant in cases with refractory HF(^j)</td>
<td>C</td>
</tr>
</tbody>
</table>

\(^a\)ACEI: Angiotensin converting enzyme inhibitors; \(^b\)ARB: Angiotensin II AT1 receptor blockers; \(^c\)LVEF: Left ventricular ejection fraction; \(^d\)LV: Left ventricle; \(^e\)FC: Functional class; \(^f\)NYHA: New York Heart Association; \(^g\)HR: Heart rate; \(^h\)AF: Atrial fibrillation; \(^i\)BB: Adrenergic beta-blocker; \(^j\)HF: Heart failure.
of oral anticoagulants is always advisable, maintaining the INR between 2 and 3\(^{264}\). Recommendations for oral anticoagulation in CCC are described in Figure 11.

### Pregnancy approach in chronic Chagas cardiomyopathy

Chronic Chagas cardiomyopathy ranks second among the cardiomyopathies present in the pregnancy-puerperal cycle, being second only to rheumatic cardiomyopathy. Most pregnant women with Chagas disease are asymptomatic or oligosymptomatic, being carriers of indeterminate or initial cardiac forms.

The risks of pregnancy in women with Chagas cardiomyopathy depend on the cardiac functional status and the presence and severity of arrhythmias. Pregnancy should be discouraged in cases with heart failure and/or arrhythmias. Pregnant women with these conditions require monitoring and special care, due to the possibility of worsening during pregnancy. In the initial consultation of the pregnant woman with Chagas disease-induced cardiomyopathy, in addition to routine tests, ECG and echocardiography (to assess cardiac dimensions and ventricular function) and a 24-hour Holter (in order to identify conduction disorders and arrhythmias) should be requested\(^{265}\). The relative and absolute contraindications to the use of drugs acting on the cardiovascular system or with a teratogenic potential should always be observed. Pregnant women with CCC must be attended to at reference centers for high-risk pregnancy, as they may require high-complexity care. It is worth highlighting the importance of carrying out all recommended assessments during the prenatal period, including HIV testing\(^{220}\).

### Surgical risk in patients with chronic Chagas disease

The evaluation of surgical risk in patients with Chagas cardiomyopathy is based on available data related to other heart diseases. However, the unique characteristics of CCC, especially those related to autonomic dysfunction, complex arrhythmias, and intraventricular stimulus conduction disturbances, can cause different responses to surgical trauma\(^{266}\). Cases with more severe myocardial damage (NYHA class IV heart failure or LVEF <30%; moderate systolic dysfunction, but with complex arrhythmia, atrial fibrillation, significant dilation of the left ventricle, total AVB and sinus node dysfunction) are more likely to have perioperative complications.

Asymptomatic sinus bradycardia in patients undergoing surgery under general anesthesia should be monitored by ECG during the procedure, which must be carried out where there is the possibility of performing emergency pacemaker implantation. If bradycardia is symptomatic, preventive implantation of pacemaker should be considered.

Patients with atrial fibrillation with a high ventricular rate (>90 beats/min), regardless of the type of anesthesia, should be operated on under continuous ECG monitoring and prior scanning. In patients with atrial fibrillation with a ventricular rate <60 beats/min, the possibility of a marked decrease in the heart rate due to anesthetic agents must be borne in mind. In these cases, the surgical procedure should be performed under continuous ECG monitoring and in an institution able to perform emergency pacemaker implantation. The same care should be taken in relation to cases with AVB associated with bundle branch block, due to the possibility of developing full AVB during surgery due to the action of anesthetic agents\(^{266}\).

Cases with complex ventricular arrhythmias should be evaluated by Holter and operated on after institution of appropriate antiarrhythmic therapy. The surgical procedure should be performed under continuous ECG monitoring and where there are defibrillators.

Preoperative evaluation of patients with cardiac pacemaker includes knowing the arrhythmia that lead to the implant, as well as generator characteristics. In addition to the clinical history, important data can be obtained from the card that cardiac pacemaker carriers receive and identifies the main characteristics of the device, such as the date of the implant, programmed heart rate, operating mode, and manufacturer.

The main risks associated with the presence of a pacemaker during surgical or diagnostic procedures consist of changing the unit’s operating threshold (by the action of drugs, changes in serum potassium levels, and changes in thoracic impedance for pulmonary ventilation), ventricular fibrillation (conduction of the electrical current from the electrocautery by the intracardiac electrode pacemaker) and, finally, damage, inhibition or reprogramming of the system by electrocautery, cardioversion, or magnetic resonance imaging. Whenever possible, biphasic electrocautery (bipolar) should be used, since the electric current remains confined between the region of the surgery and the electrocautery, reducing the risk of leakage and interference with the device\(^{224}(26)\).

In any case, the use of electrocautery should be minimized. It should be used in short and irregular intervals, monitoring not only the ECG, but also the pulse by plethysmography or pulse oximeter. In the event of severe bradycardia or tachycardia during the use of the electrocautery, a magnet cover can be used over the pacemaker, which makes it work in its programmed

---

**FIGURE 11.** Recommendations for oral anticoagulation in chronic Chagas cardiomyopathy.
magnetic response. Attention should be paid to the fact that the placement of the magnet should be restricted to the short periods of electrocautery use. The patient should be advised postoperatively to return to the clinic for pacemaker follow-up for the assessment of its operation and eventual reprogramming of its generator\(^2\)\(^{7}\).

In patients with a CDI, the presence of technician or specialist in the operating room is recommended. The antitachycardia function should be switched off, with staff prepared to treat any possible arrhythmias, including having to perform electrical cardioversion\(^2\)\(^{4}\)\(^{1}\)(\(^2\)\(^{4}\)\(^{7}\)).

If electrical cardioversion is required in patients with pacemakers, the ideal would be to use adhesive pads in the anteroposterior chest position, as remote as possible from the generator, using as little energy as possible for resolution of the arrhythmia. Similarly, the thresholds and the functioning of the generator, using as little energy as possible for resolution of the arrhythmia. Similarly, the thresholds and the functioning of the pacemaker should ideally be assessed after the procedure\(^2\)\(^{4}\)\(^{4}\).

**Occupational health evaluation**

Chronic Chagas cardiomyopathy is an important cause of incapacity to work in endemic areas. It should be the subject of surveillance actions for workers' health, included in primary care. Careful clinical examination, with particular attention to the cardiovascular system, is the basic component of functional and occupational health evaluation in CCC\(^2\)\(^{6}\). These assessments should take into account: the degree of ventricular dysfunction; the presence, magnitude and complexity of conduction and rhythm disorders; and the thromboembolism risk. It is also important to consider the characteristics of the labor activity performed by the worker, especially the need for intense or continued physical effort in the activity, as well as the degree of professional qualification of the worker and his/her ability to enroll in a vocational rehabilitation program. The age of the worker should also be considered, as the possibility of evolution of heart disease is greater for younger workers and vice versa. In general, the worse the LVEF and the greater the ventricular diameter, the worse the prognosis and the greater the inability to work\(^2\)\(^{6}\).

The presence of cardiomegaly and significant ventricular dysfunction implies incapacity to perform activities that require physical effort, and, when vocational rehabilitation is not possible, disability. In all functional assessments of prognosis and working capacity of an individual with Chagas disease, the great clinical variability of these cases should be taken into account, even when in the same stage of any classification of cardiac impairment. Often, but not always, advanced disorders of stimulus conduction are associated with the severity of myocardial damage, leading to the need for a complementary evaluation of the degree of myocardial dysfunction, especially in cases where the work requires intense physical activity and involves personal or social risk (e.g., aviation pilots)\(^2\)\(^{7}\).

**DIGESTIVE FORM OF CHAGAS DISEASE**

The digestive form of Chagas disease can affect all organs in the gastrointestinal tract. However, from a practical point of view, it is manifested as involvement of the esophagus and large intestine, resulting in megaesophagus and megacolon, respectively\(^2\)\(^{2}\). The association between megaesophagus and megacolon in cases that have an indication for surgery is approximately 92%. The association between megaesophagus, megacolon, and heart disease is 65%\(^2\)\(^{7}\). A specific search for \textit{T. cruzi} infection should be undertaken in patients with clinical syndromes consistent with those described below, who live in at-risk contexts and/or who have increased vulnerability to Chagas disease\(^2\)\(^{2}\).

**Diagnosis**

**Digestive manifestations in the acute phase of Chagas disease**

Clinical manifestations are usually nonspecific, with practically imperceptible symptoms related to the gastrointestinal tract. However, there are reports describing the occurrence of gastrointestinal bleeding in cases of oral transmission and of dysphagia in rare cases of vectorial transmission\(^2\)\(^{2}\).

**Digestive manifestations in the chronic phase of Chagas disease**

Digestive manifestations of Chagas disease are concentrated in the esophagus and colon and are basically dysphagia and constipation, due to chronic alterations that can lead to megaesophagus and/or megacolon\(^2\)\(^{2}\). However, as the disease causes autonomic nervous system injury throughout the gastrointestinal tract, anatomical and functional abnormalities of the salivary glands, stomach, extra-hepatic bile ducts, duodenum, small intestine, large intestine, and even of organs not belonging to the gastrointestinal tract, such as the ureter, can occur. The prevalence of \textit{Helicobacter pylori} infection and of characteristic gastric endoscopic and histologic changes were similar in patients with and without Chagas disease. Furthermore, studies have shown that \textit{H. pylori} is the main cause of gastritis in patients with Chagas disease\(^2\)\(^{2}\). The most common symptoms and signs of the digestive system, in the chronic phase of Chagas disease, are described below.

**Dysphagia**

**Clinical diagnosis**

Dysphagia is the main symptom of patients with severe impairment of the esophagus. Other dysphagia-related complaints occur in cases of megaesophagus, including ptyalism, hiccup, odynophagia, regurgitation, a sense of nocturnal choking, aspiration pneumonia, and malnutrition\(^2\)\(^{2}\). In the early stages of the disease, the patient reports a feeling of obstruction at the level of the xiphoid after eating solids and, later, after fluid intake, especially cold fluids. Dysphagia progresses slowly and is well-tolerated for many years. Therefore, patients with achalasia often do not seek care in health services until the progressive dysphagia interferes with their lifestyle. It is observed that the patients themselves spontaneously change their eating habits to alleviate dysphagia. Patients with achalasia feed slowly, ingest large volumes of water to assist the food into the stomach, often bend the back, lift the chin, extend the neck, or walk to help esophageal emptying. Regurgitation of undigested food is common as the disease progresses, with a
risk of aspiration, which may lead to pneumonia, lung abscess, bronchiectasis, hemoptysis, and bronchospasm. Great distension of the dilated esophagus may lead to dyspnea, compression of the main stem of the bronchus or lung hilum.

Esophageal involvement occurs to varying degrees, with great variation in morphology and differences of esophageal motor behavior, ranging from minimal changes of esophageal transit to advanced forms of dolicomegaesophagus with an extremely prolonged esophagogastric transit time. Considering practical and therapeutic aspects, patients with megaesophagus can be classified into groups, depending on the degree of expansion of the organ, as shown in Figure 12.

Most cases of megaesophagus are found in groups II and III. The degree of megaesophagus does not necessarily reflect the duration of the disease. There are cases that rapidly progress to advanced forms, while others remain stable in the early stages. Lower esophageal sphincter dysfunction is greater in cases where the motor abnormalities of the esophagus are more evident. Although there is a correlation between the degree of megaesophagus and the severity of dysphagia, it is common to find patients with bulky megaesophagus complaining about or not even reporting dysphagia. Notably, the incidence of esophageal cancer in patients with idiopathic achalasia is 3.3%, 15 times higher than that of the general population without achalasia.

**Complementary examinations**

- **Plain radiography of the thorax** (posterior-anterior or lateral views): In cases of megaesophagus grade III or IV, great dilation of the esophagus extending to the posterior mediastinum can be observed (Figure 13 A and B).

**Contrast radiography of the esophagus:** This shows not only the degree of esophageal dilation, allowing classification of megaesophagus, but also demonstrates functional changes, such as abnormal or missing peristaltic waves and the emptying time of esophageal contents into the stomach. The radiologic characteristics of megaesophagus in Chagas disease are: increased esophageal diameter and emptying time (Figure 14); a tapered neck; numerous secondary and tertiary waves in the esophagogram (Figure 13 C).

**FIGURE 12. Radiologic classification of megaesophagus.** (A) Group I: The esophageal diameter is within normal limits, without food stasis, but with an increase in food transit time from mouth to stomach. Functional motor disturbances predominate without corresponding dilation. (B) Group II: Moderate dilation of the esophagus with loss of motor control. Secondary and tertiary waves can be seen in the esophagogram. (C) Group III: More accentuated dilation can be observed than in group II, transit time more prolonged and reduced motor activity. The esophagus behaves like an inert tube. (D) Group IV: Consists of advanced forms with large dilation and stretching of the esophagus (dolicomegaesophagus).

Source: Rezende, 1982.
FIGURE 13. Plain radiography of the thorax. (A) Posterior-anterior view and (B) lateral view, demonstrating the great dilation of the esophagus extending to the posterior mediastinum.

Source: Radiographs of a patient attended at the Hospital das Clínicas, Federal University of Minas Gerais (UFMG).

FIGURE 14. Contrast study of a patient with esophageal achalasia (group III). Images taken (A) 2 minutes and (B) 6 hours after ingestion of contrast. Note the pacemaker wires implanted in this patient due to AV block of concomitant Chagas cardiomyopathy.

Source: Radiographs of a patient attended at the Hospital das Clínicas, Federal University of Minas Gerais (UFMG).

Note: Images taken 2 minutes (A) and 6 hours after ingestion of contrast (B). Note the pacemaker wires implanted in this patient due to AV block of concomitant Chagas cardiomyopathy.

Note: Images taken 2 minutes (A) and 6 hours after ingestion of contrast (B). Note the pacemaker wires implanted in this patient due to AV block of concomitant Chagas cardiomyopathy.

Note: Images taken 2 minutes (A) and 6 hours after ingestion of contrast (B). Note the pacemaker wires implanted in this patient due to AV block of concomitant Chagas cardiomyopathy.
distal third; retention of food debris; esophageal/mediastinal stretching; and reduced gastric volume.

- **Esophageal electromanometry:** This is a useful method to distinguish between the differential diagnoses of other diseases that develop due to motor dysfunction that present with dysphagia. In patients without dilatation, or with slight dilation of the esophagus (group I and II), uncoordinated pressure waves can occur in the body of the esophagus and incomplete or atypical relaxation of the lower esophageal sphincter (LES) can be observed. These findings precede radiologically demonstrated abnormalities.

- **Endoscopy:** Although endoscopy is not essential to confirm the diagnosis of megaesophagus, it should be performed to assess the degree of mucosal inflammation and to exclude the presence of neoplastic lesions in patients with dysphagia, varicose veins, and ulcers. This examination is useful in cases requiring forced balloon dilatation of the gastric cardia and in cases requiring botulinum toxin injection.

**Colon**

**Clinical diagnosis**

The main signs and symptoms of megacolon are constipation, paradoxical diarrhea (constipation interspersed by periods of diarrhea), dyschezia, abdominal distension, and fecaloma. Anorectal manometry is useful to diagnose internal anal sphincter achalasia. Colonic dilatation located in the rectum and sigmoid is better observed by contrast radiography studies and can be detected in up to 80% of cases. However, dilatation can occur only in the rectum, only in the sigmoid, or in the entire colon. It should be emphasized that the clinical picture of severe constipation in patients with idiopathic megacolon who are not infected with *T. cruzi* is indistinguishable from that induced by Chagas disease.

**Complementary tests**

- **Plain radiography of the abdomen:** Bulky megacolons can be detected on plain radiographic films of the abdomen, especially if a fecaloma is present. Plain radiography is also useful in cases of suspected volvulus, since it can demonstrate rotation of the sigmoid and dilatation of small bowel loops caused by obstruction (Figure 15).

- **Contrast radiography of the colon (barium enema):** This is the gold standard for the diagnosis of megacolon. It shows dilation and/or stretching of the colon, presence of fecaloma, and anatomic abnormalities of the sigmoid (Figure 16).

- **Computed tomography and magnetic resonance imaging:** These more modern imaging tests occupy a prominent place in the evaluation of patients with Chagas disease and are gradually replacing conventional imaging methods. Both methods are more sensitive in evaluating pericolonic and/or perisophageal tissues; they can detect infiltration, not only visible masses and fecalomas detectable by conventional radiography. These modalities also provide greater detail than a barium enema (Figure 17A, B and C).

- **Digestive endoscopy:** This is useful in cases of megacolon, to assess the degree of mucosal inflammation, detect ulcers, wall necrosis, perforation or bleeding, and to rule out the presence of associated lesions, particularly cancer. It is particularly important in cases of suspected sigmoid volvulus and can be therapeutic, since it might allow laparotomy to be avoided. If it is not possible to reduce the sigmoid volvulus by endoscopic examination, emergency surgery is absolute indicated.

**Megaesophagus**

Considering the impossibility of restoring normal physiology due to the irreversible changes caused by denervation, the goal of treatment is to act on the lower esophageal sphincter in a patient with achalasia, by removing the functional barrier to the passage of food into the stomach. The treatment, except if specific, does not cure the disease; it seeks only improve the symptom of dysphagia.

**Clinical treatment**

This is indicated in patients with advanced age, with no history of complications, who are oligosymptomatic, are at high risk for surgical treatment, who refuse to undergo invasive treatments, or those treated in hospitals lacking adequate infrastructure to perform this type of surgery. The basis for the treatment is specified below.

![Image of patient attended at Hospital das Clinicas, UFMG.](source: Radiograph of a patient attended at the Hospital das Clinicas, UFMG.)

**FIGURE 15.** Simple radiograph of the abdomen in a case of sigmoid volvulus. Note the anomalous position of the sigmoid caused by the dilatation and the characteristic image of the small bowel loops secondary to the obstruction caused by the volvulus.
1. **Counseling and health education** related to megaesophagus, in order to improve patient safety.

2. **Adequacy of eating habits.** It is recommended that the patient chew their food well; eat small amounts at a time; and consume liquid foods, pastes, and solids appropriate to their severity of dysphagia. Irritating foods, such as very spicy or hot foods or ice cream, aggravate dysphagia and should be avoided. Due to the risk of regurgitation and aspiration, food intake should be avoided before bedtime. Balanced enteral nutrition is indicated in patients with megaesophagus in Groups III and IV, malnourished patients, and candidates for surgical treatment. If the passage of a nasoenteric tube is not possible, parenteral nutrition can be administered via a central venous catheter. In both situations, the intervention is temporary.

3. **Use of drugs:** Medicines that relax smooth muscle fibers of the lower esophageal sphincter can relieve the symptoms of dysphagia. These drugs include:
   - Isosorbide dinitrate, at a dose from 2.5mg to 5mg, administered sublingually, 15 minutes before each meal.
   - Nifedipine, at a dose of 10mg, administered sublingually, 30 minutes before each meal. The use of nifedipine should be performed with caution, considering the significant association of this condition with chronic Chagas cardiopathy; nifedipine may induce hypotension and shock.

FIGURE 16. Radiograph of a barium enema examination revealing a hypodense image typical of a fecaloma.

FIGURE 17. Abdominal computed tomography, showing a megaesophagus with an air-fluid level formed by ingested contrast (A); megacolon with a large dilatation of the air-filled colon with a fecaloma in the rectum (B); and megacolon with a predominant image of a fecaloma in the dilated sigmoid (C).

Source: Radiograph of a patient attended at the Hospital das Clínicas, UFMG.
4. **Botulinum toxin**: Botulinum toxin type A, applied locally, blocks the release of acetylcholine from pre-synaptic nerve endings and leads to chemical denervation. The treatment is effective in 70% of patients, and the average duration of symptom remission is 16 months. Patients over 50 years of age respond better to this treatment (279) (280). Botulinum toxin treatment of megaesophagus has worse outcomes than those of dilation or surgery (level IIB). Although the effect is temporary, botulinum toxin injection may be indicated in selected cases, presenting the advantage that it can be repeated several times. The dose of botulinum toxin is 20 units injected into each of the lower quadrants of the esophageal sphincter. One of the limitations of this treatment is that the procedure should be guided by endoscopy.

5. **Balloon dilatation**: Pneumatic dilatation is indicated for patients with group I megaesophagus, but can be performed in patients in groups II and III who have a contraindication to surgery. It can also be used preoperatively in patients in groups II and III who are candidates for surgical treatment of megaesophagus, aiming to improve their preoperative nutritional status. In patients in group IV, due to the risk of redundant esophageal perforation, dilatation is not recommended. The procedure is performed with a balloon coupled to gauge introduced endoscopically, which makes it the safest method. The result of megaesophagus treatment by balloon dilatation relieves dysphagia in 71% of patients (280). However, the effect is transient, worsening over time with successive dilations.

**Surgical treatment**

1. **Conventional surgery (laparotomy)**: This does not address the underlying cause of the disease, but restores the ability of the patient to swallow. Based on the knowledge gained about the pathogenesis and pathophysiology of megaesophagus, the cardiomyotomy known as Heller surgery (281) associated with some kind of partial fundoplication (282), is the most appropriate surgical procedure for cases of megaesophagus grades I to III. The response rate is about 90% (280), giving a more long-lasting and effective result than that achieved by pneumatic dilatation. For dolichomegaesophagus, there is still no consensus among surgeons; hence, different surgical techniques are still used.

2. **Laparoscopic surgery**: This technique was introduced in Brazil in the early 1990s, with simultaneous availability of modern surgical instruments, such as electronic scalpels and staplers. The experience of the surgeons with this technological innovation and its use in the treatment of cases of megaesophagus from achalasia related to Chagas disease, has reduced the morbidity and mortality associated with conventional surgery for megaesophagus induced by Chagas disease (level IA). Laparoscopic cardiomyotomy, associated with some type of antireflux valve, especially those made by partial fundoplication, has become standard surgery; it is virtually the standardized surgical procedure for patients with this condition (283).

3. **Robotic surgery**: Robotic surgery is probably the most advantageous of all the surgical options. However, the complexity of its implementation and the high associated costs hinder its routine application in practice (220). Megaesophagus treatment by myotomy associated with antireflux surgery carries a risk of esophageal perforation; this complication can be severe. Robotic surgery, by making scheduled and precise movements, will most certainly reduce the rate of this complication.

4. **Endoscopic surgery**: Technological advances and the increasing expertise of professionals has allowed myotomy to be performed as a totally endoscopic procedure. This surgery is now known by the acronym POEM (per-oral endoscopic myotomy). It uses the Heller technique, performed by transluminal endoscopic surgery through natural orifices [natural orifice transluminal endoscopic surgery (NOTES)]. This procedure was introduced by professionals from the Mayo Clinic Endoscopic division and, after testing on animals, the procedure was widely disseminated, being used in humans since 2012. The POEM technique seems to be a safe method, and is currently an effective alternative for the treatment of achalasia.

5. The procedure is performed with the person in the supine position under general anesthesia, and consists in approaching the esophageal lumen with a high-definition gastroscope with frontal view. Saline stained with indigo carmine is injected to create a safety gap between the mucosa and submucosa of the esophagus. Then, a 2 cm longitudinal incision of the mucosa is performed, at the two o’clock position, 10–15 cm proximal to the gastroesophageal junction, to expose the submucosal layer. A tunnel is dissected in the submucosa, extending from the point of section to 2–3 cm beyond the gastroesophageal junction (GEJ) (284). Myotomy of the lower esophageal sphincter is usually performed starting at 2 cm distal to the opening of the mucosa to below the GEJ. As a rule, myotomy should have a minimum length of 6 cm (on average, 8–10 cm): 2 cm in the esophagus, 2–3 cm in the lower esophageal sphincter, and 2 cm in the cardia. Dissection of the submucosal tunnel is performed in the plane before the muscle of the mucosa, and esophageal muscle section is performed only in the circular layer of the muscle, there is no need for longitudinal muscle layer section.

6. Data from 14 English language publications show that of 804 patients who underwent a POEM procedure, therapeutic success was documented in more than 80% (level A1) (285). The technique is less invasive and the patients do not experience gastroesophageal reflux, as observed in cases submitted to myotomy by conventional laparotomy or laparoscopy surgical technique. The familiarization of the endoscopist with the method enables secure closure of smaller perforations and adequate hemostasis (level B1) (280). If there is a more extensive perforation of the esophagus, or more intense bleeding, the assistance of a surgeon may be necessary.
Megacolon

Clinical treatment

1. 
Health counseling and education related to megaesophagus, in order to improve patient safety.

2. 
Adequacy of eating habits: This should focus on a higher degree of self-care. Patients evacuated two to three times a week without reporting a fecaloma should be instructed to drink more fluids, and eat more fruits and vegetables. The use of a diet rich in fiber in these patients is controversial, because there is the fear that excess fiber may favor fecal impaction and the worsening of symptoms. The diet can be the patient’s usual diet, with restriction of constipating foods. Abundant water intake is recommended – a volume of at least 2L/day – as well as foods that favor intestinal functioning.

3. 
Regular defecation: It is necessary to systematically pay attention to the urge to defecate and to create the habit to evacuate at certain hours.

4. 
Laxatives: These are indicated for patients who do not adequately respond to dietary measures. Osmotic laxatives (20% mannitol or polyethylene glycol) are the most suitable. Mineral oil is also effective.

5. 
Avoidance of potentially constipating drugs: Opioids, diuretics, antidepressants, antihistamines, anticonvulsants, antiparkinsonians, among others, should only be prescribed under medical monitoring.

6. 
Removal of fecaloma: Fecalomas, located in the rectum, can be removed manually (under anesthesia) if they do not respond to dietary and drug measures.

7. 
Intestinal wash-out: Patients who do not respond satisfactorily to drug and dietary treatment, or those with significant fecalomas, should be treated with glycerin or saline enemas. Intestinal wash-out with 500–1,000mL twice a week is recommended for patients resistant to medical treatment. In patients with fecalomas, the procedure may be repeated two to three times a day, taking care not to use more than 3.5L of solution per day. Experience with these treatments shows that, with the removal of the fecalomas and normalization of bowel movements, the diameter of the colon tends to return to normal.

Surgical treatment

1. 
Elective surgery: Surgical treatment of megacolon is indicated for patients who have large dilatation of the sigmoid, repeated episodes of volvulus, prolonged retention of feces, recurrent fecalomas, and difficulties in applying enemas at home. Similar to the surgical treatment of megaesophagus, the goal of treating the megacolon is to improve or normalize the intestinal rhythm, since the predominant symptom is constipation. Except in cases of emergency caused by volvulus, bleeding, obstruction or perforation, in which emergency procedures must be adopted (colonoscopy or surgery), megacolon surgery is elective and should be well planned. The most commonly used conventional surgical techniques are the aforementioned rectosigmoidectomy and the surgery originally proposed by DuHamel for the treatment of Hirschsprung disease\(^{287}\). With the advent and spread of laparoscopic surgery, the laparoscopic DuHamel procedure has become the gold standard for treatment of these cases\(^{288}\).

2. 
Emergency surgery: The indications for emergency surgery to treat megacolon are: sigmoid volvulus not resolved by endoscopy; and ulceration, necrosis and/or perforation, due to volvulus or endoscopy.

3. 
The surgery to be performed will depend on the patient’s condition and anatomopathological situation. The surgical procedure may vary from a simple colostomy to fixing the part affected by sigmoid resection with primary or delayed anastomosis.

ANTIPARASITIC TREATMENT OF CHAGAS DISEASE

Although there is disagreement about the cure rates of the antiparasitic treatment of Chagas disease, there is consistent evidence showing its usefulness in both acute and chronic phases of the disease and in all clinical forms of chronic infection, as the organic lesions depend exclusively (acute phase) or at least in part (chronic phase) on the presence of the parasite. In addition, there is apparent suppression of the parasitemia with the current antiparasitic treatment\(^{289}\)\(^{290}\). In turn, the rates of cure and their confirmation depend on several factors, such as the phase and the duration of disease, the age of the patient, the tests used to evaluate therapeutic efficacy, the duration of follow-up after treatment, associated conditions, and the susceptibility of the specific \textit{T. cruzi} strain to antiparasitic drugs.

Treatment in the acute phase

In the acute phase, treatment should be provided to all cases and as quickly as possible, regardless of the parasite transmission route (Class I, level of evidence B). Studies of a case series followed for over 20 years have shown cure rates above 50% at this stage of the disease\(^{291}\).

Treatment of congenital Chagas disease

Patients diagnosed with congenital Chagas disease should receive antiparasitic treatment (more details on the specific part of vertical transmission)\(^{290}\), regardless of whether the diagnosis has been made by parasitologic methods in the first weeks of life or by conventional serologic tests at least nine months after birth (Class I, level of evidence B).

Treatment of chronic phase

All children aged less than 12 years with chronic phase Chagas disease should be treated (Class I, level of evidence A). Prospective, randomized, double-blind, controlled studies with placebo, performed in asymptomatic school-aged children,
demonstrated therapeutic cure of *T. cruzi* infection in 58–62% of cases\(^{292}(293)\). In an uncontrolled study involving Latin American countries\(^{294}\), large regional differences were observed in treatment response rates: The seroconversion rate in Honduras was 92.7% for children aged 9 months to 12 years and 58.8% in a quarter of the study patients younger than 15 years. There were variations in Bolivia, from 0% in Sucre to 5.4% in Entre Rios. In the latter region this rate was higher in children younger than 5 years (24.4%) than in children aged 5–9 years (4.6%)\(^{294}\).

For adolescents (aged 13–18 years) and adults with chronic infection, when it can be established that the acute phase occurred up to 12 years before (considered as recent infection) antiparasitic treatment is usually recommended\(^{290}(294)\), although there is no consistent evidence from randomized studies justifying this (Class IIa, level of evidence C)\(^{295}\).

For individuals with Chagas disease aged 19–50 years with no recent documented infection, antiparasitic treatment should be considered on an individual basis, whether in ICF\(^{120}(252)\) (Class IIa, level of evidence B) or in the determined chronic form, without advanced cardiopathy\(^{42}(120)(252)(295)(296)(297)(298)\) (Class IIIb, level of evidence C). Specifically, treatment of chronically infected women of childbearing age, when provided before pregnancy, can reduce congenital transmission\(^{104}(109)\).

Some observational studies have shown that in this age group, antiparasitic treatment is able to prevent the onset or delay the progression of the disease in a significant proportion of cases\(^{42}(120)(252)(295)(298)\). This would lead to a reduction in the complications of the disease, especially those related to heart disease. Classical experimental studies in mice have already documented total or partial regression of early myocardial injuries and the prevention of cardiopathy with antiparasitic treatment\(^{299}(300)\). However, regression of inflammatory and fibrotic lesions, observed in experimental studies, has not been verified in the clinical context\(^{290}\).

A case series demonstrated cure in at least 20% of the cases treated in the late chronic phase\(^{101}\). It should be noted that demonstrating cure depends on various factors, including follow-up of treated cases, which should continue for more than two decades for patients treated in the late phase. Increased seroconversion of cases showing no acute illness and without cardiac failure was also assessed\(^{290}\).

For individuals older than 50 years, without advanced heart disease, there are no studies justifying antiparasitic treatment. An observational study in elderly subjects infected with *T. cruzi* demonstrated that the disease is present and active – the percentage of patients with ICF decreases but the percentage with the heart form increases\(^{100}\). Thus, these individuals should not be completely excluded from the etiological treatment plan of Chagas disease; treatment can be individualized (Class IIb, level of evidence C).

Antiparasitic treatment should not be given to patients with chronic phase Chagas disease with severe heart dysfunction, since there is no evidence of clinical benefit\(^{251}(289)(291)(295)(296)(302)(303)\) (Class III, level of evidence C).

Recently, a series results from the BENEFIT study – a prospective, multicenter, randomized, double-blind, controlled study with placebo (placebo vs. benznidazole) – was published\(^{251}\). This is the first study to evaluate the effect of a specific treatment for up to 80 days in relation to cardiac outcomes and *T. cruzi* infection. The effects of benznidazole in reducing morbidity and mortality among 2,854 patients with established chronic Chagas cardiopathy were evaluated\(^{250}(251)\). After 5.4 years of follow-up, the results showed no statistically significant difference between the two groups in relation to the primary composite outcome that included death, sustained ventricular tachycardia, recent heart failure, implantation of a cardioverter-defibrillator or pacemaker, cardiac transplantation, resuscitated cardiac arrest, stroke, or other thromboembolic event\(^{251}\). These findings, at first, seem to indicate a divergence of considerably lower rates of parasite detection by PCR in blood in the same study after treatment with benznidazole\(^{383}\).

In the BENEFIT study, the treatment with benznidazole in the protocol used probably has no significant preventive effect on the progression of heart disease in patients with advanced Chagas disease\(^{251}\) (Class III, level of evidence B). However, the severity of Chagas disease is emphasized in this study, as is the resulting need for diagnosis and timely treatment, based on qualified management, especially of heart disease. In the study, 503 patients died within 5 years (246 the in benznidazole group and 257 in the placebo group), and 63% of patients (in both treatment groups) developed a left ventricular ejection fraction <40%, one of the primary outcomes of the study\(^{251}\).

The study highlighted, therefore, the urgent need to structure a network of qualified health care to manage patients with Chagas disease comprehensively, without loss of diagnostic and treatment opportunities. Longitudinal clinical management is needed in terms of the various clinical forms of the disease, especially the advanced cardiopathy form, with protocols adjusted to local realities. In addition, the ability to create specific treatment opportunities for people with therapeutic indications based on more consistent evidence, is required\(^{303}\).

Furthermore, the study draws attention to the lower frequency of serious adverse effects to benznidazole compared with previous study findings\(^{251}(300)\). This reinforces the real possibility of including, in a more expanded form, the specific treatment of Chagas disease in the primary healthcare network (primary) not only in Brazil but also in other endemic countries.

The digestive form of Chagas disease is not a contraindication for antiparasitic treatment. Moreover, megaesophagus correction should be performed in order to ensure full transit of the drug and, subsequently, its absorption\(^{20}(302)\) (Class IIa, level of evidence C).

**Treatment of immunosuppressed cases**

Cases with immunodeficiency, including patients taking immunosuppressive drugs to treat neoplasms, transplants, and other diseases, and patients with *T. cruzi/HIV* co-infection, can develop reactivation of Chagas disease\(^{181}(184)(304)\). This is discussed, in themes, below.

**Transplantation**

In transplants in general, it is necessary to know if the donor or the recipient has Chagas disease, considering the risk.
of transmission or reactivation of the disease. Thus, there are three possible scenarios:

- donors with Chagas disease and recipient without Chagas disease;
- donors without Chagas disease and recipient with Chagas disease;
- donors and recipient with Chagas disease.

**Donor with Chagas disease and recipient without Chagas disease**

1. **Guidelines for the donor with Chagas disease:**

   - The candidate for organ donation who has positive serology for *T. cruzi* should be treated with antiparasitic drugs for 60 days before the procedure. In special situations requiring transplantation before completing treatment, the transplant can be performed, but if possible, not before 14 days of treatment has been completed (Class I, level of evidence C).

2. **Guidelines for the recipient when the donor is treated:**

   a. Consider etiological treatment soon after transplantation, maintaining treatment for 60 days by means of intense immunosuppression (Class IIa, level of evidence C)\(^{(184)}\).

   b. Do not treat, but provide sequential clinical monitoring or parasitologic evaluation as follows: weekly direct search for *T. cruzi* in the peripheral blood, until 60 days post-transplantation; indirect parasitologic methods (blood and/or xenodiagnosis) of follow-up of the recipient during the period of immunosuppression; and serologic examination 30 and 60 days post-transplant.\(^{(181)}\)\(^{(184)}\)\(^{(305)}\). In cases where monitoring is not possible, etiologic treatment is recommended. The following clinical and serologic examinations should be performed at 3, 6, 9, and 12 months of follow-up: direct/indirect parasitologic tests (blood culture and/or artificial xenodiagnosis) at 3, 6, and 12 months or PCR. Subsequently, the evaluation is repeated every 6 months, while there is persistent immunosuppression (Class IIa, level of evidence B)\(^{(181)}\)\(^{(184)}\)\(^{(305)}\). It is noteworthy that, in some reference centers, PCR, but not indirect parasitologic examination, is performed. At any time, if an acute infection is detected through parasitologic or serologic tests, the conventional etiological treatment should be instituted. Out of the usual controls, any clinical signs suspicious of infection should be investigated through parasitologic examination, and positive cases should receive conventional treatment for the acute phase. Since 2009 in Brazil, organ transplantation laws have restricted the possibility of patients with Chagas disease being accepted as organ donors\(^{(183)}\). In Brazil, Ordinance number 2,600, in 2009, which approved the Technical Regulations of the National Transplant System, states that it is mandatory to perform serologic testing for *T. cruzi* infection in the following situations\(^{(183)}\): I) for all donations, following the same algorithms used for screening blood donors; II) for registration purposes of potential organ recipients in a Single Technical Registry; III) for all potential deceased donor organs, tissues, cells, or body parts before the allocation of the grafts. It also establishes criteria for the classification of living and deceased donors as well as for potential recipients for the distribution of organs, tissues, cells, and captured parts. It does indicate that the team will decide whether the organ/tissue is acceptable for use or not regarding kidney, kidney/pancreas, pancreas, liver, and lung donations. In the order, the rejection of organs relates only to the heart. For ocular tissues, hematopoietic stem cells, musculoskeletal tissues, skin, and cardiovascular tissue, the issues related to *T. cruzi* infection are not mentioned\(^{(183)}\).

3. **Guidance for the recipient without Chagas disease:**

   In cases in which the donor with Chagas disease has not been treated or has received incomplete treatment, three possible actions may be taken in relation to the recipient:

   a. Start antiparasitic treatment soon after transplantation, for a 60-day period (class IIa, level of evidence C). This is recommended if the sequence monitoring indicated in item c below is not possible.

   b. Start treatment immediately after surgery, for an initial 14-day period. Then, proceed to sequential monitoring with clinical, parasitologic, and serologic evaluation (Class IIa, level of evidence C or without evidence). If seroconversion occurs, introduce antiparasitic treatment for the acute phase.

   c. Do not treat, but proceed to sequential clinical monitoring and parasitologic evaluation (Class IIa level of evidence B)\(^{(181)}\)\(^{(184)}\)\(^{(305)}\). Such evaluation consists of direct search for *T. cruzi* in the peripheral blood, every week, for up to 60 days, and indirect parasitologic and serologic examinations on days 30 and 60 after transplantation. Thereafter, clinical, serologic, and parasitologic (direct/indirect/PCR) examinations should be performed every two months to up to one year of follow-up; then every six months, for as long as immunosuppression persists (time dependent on the mode and type of transplant). It is worth emphasizing that in some centers PCR, rather than indirect parasitologic examination, is performed.

   d. At any time, if an acute infection is detected by parasitologic or serologic tests (IgM or seroconversion), conventional antiparasitic treatment should be started. Outside of the above-stated monitoring intervals, any clinical sign of suspected infectious condition should be investigated through parasitologic examination, and positive cases should receive conventional treatment for acute stage disease\(^{(20)}\)\(^{(302)}\).

   e. A few of cases have been reported of kidney transplant recipients without Chagas disease who were treated with antiparasitic prophylaxis for 14 days, as the organ donors had Chagas disease\(^{(308)}\). In another work, prophylaxis...
was provided for 60 days to recipients (without Chagas disease) who received livers from donors with Chagas disease\(^{300}\) as usual strategy; seroconversion was not observed in the surviving recipients, as this may not occur in about 20% of patients in the post-transplant period. Use of antiparasitic drugs in patients without documented infection (no identification of the parasite or evidence of seroconversion) is considered prophylactic use. In this text, even when the drug was administered for the same duration as the duration of treatment (60 days), the term prophylaxis has been used if the drug was used in the absence of a documented infection in the recipient. The works cited are not controlled and, in addition of having been carried out in a small number of cases in kidney and liver transplant recipients, these did not use the systematic monitoring of parasitemia, and the final results were based on the absence of seroconversion in patients who survived.

f. However, since 2009, the possibility of patients with Chagas disease being accepted as organ donors in Brazil has been restricted by transplant regulations\(^{184}\), with systematic serologic screening limiting this type of donation in individuals with \textit{T. cruzi} infection. However, these same organs can function as "marginal organs" in specific situations\(^{184}(305)(307)\).

**Donor without Chagas disease and recipient with Chagas disease**

In these cases, the expected complication is the reactivation of Chagas disease in the recipient, due to the use of immunosuppressive drugs to prevent or treat rejections. With time, pre-transplant conventional antiparasitic treatment can be instituted. However, it should not prevent the realization of transplantation, as subsequent actions can be adopted (Class IIa, level of evidence C).

1. **Prophylaxis of episodes of reactivation:** In heart transplant recipients, antiparasitic treatment may fail\(^{100}(307)\).
2. **Clinical and parasitologic monitoring of the recipient after transplantation** should be carried out, as in the case with a donor with Chagas disease and a recipient without Chagas disease, and early conventional treatment of reactivation episodes should be provided\(^{181}(184)(305)\). The monitoring during the post-transplant period in patients receiving cardiac transplantation is mandatory and must include monitoring of blood and tissues, due to the greater sensitivity of the methods in the reactivation site (myocardium).

**Donor and recipient with Chagas disease**

Both should be considered as carriers of the chronic form of Chagas disease and should be evaluated for antiparasitic treatment, as described for cases with the chronic form of the disease. After transplantation, the recipient should be monitored from a clinical and parasitologic point of view for the diagnosis of potential episodes of reactivation, such as in the case of a donor with Chagas disease and a recipient without Chagas disease. If reactivation occurs, conventional treatment for the acute phase should be provided (class IIa, level of evidence C). Episodes of reactivation can occur more than once and should be treated when documented. Hence, systematic parasitologic monitoring while immunosuppression persists, is recommended.

**Cointfection Trypanosoma cruzi/HIV**

Observational studies have shown that, in the presence of \textit{T. cruzi}/HIV cointfection without AIDS, the natural evolution of both diseases occurs, and one should follow the guidelines for antiparasitic treatment of Chagas disease in its acute and chronic forms, or monitoring of parasitemia, quantitatively if possible\(^49(159)(175)\) (Class IIa, level of evidence C) (see specific part of this Consensus). In patients with co-infection and AIDS, reactivation of Chagas disease might occur, and the following measures are recommended: Specific antiparasitic treatment is indicated for reactivation with a clinical syndrome of myocarditis, meningoencephalitis, or other; and parasitologic examination of the blood or cerebrospinal fluid yields positive results by direct microscopy; or characteristic lesions are demonstrated on histologic examination of tissue specimens\(^{20}(49)(159)\) (Class I, level of evidence C). This therapeutic approach should also be adopted in cases without evidence of reactivation but with the associated clinical syndrome and a high level of parasitemia, as defined by Sartori et al. as the presence of more than 20% of positive nymphs in the same xenodiagnostic test (indirect parasitologic examination, quantitative PCR) or sustained parasitemia\(^{49}(159)(175)(308)\) (Class IIa, level of evidence C).

**Immunosuppression in other diseases associated with Chagas disease**

Clinical conditions such as neoplasms and collagen vascular disorders in carriers of Chagas disease might be associated with reactivation. The recommendations are the same as those for reactivation occurring in organ transplantation patients and patients with HIV coinfection\(^{20}(49)(159)(175)\).

**Accidental transmission**

In accidents characterized as high risk for disease transmission, such as percutaneous injury or mucosal contact with biological material containing live parasites (samples for the culture of \textit{T. cruzi} obtained from of patients with high parasitemia or necropsy material, vectors, and animals that are already infected), primary prophylaxis should be given.
Prophylaxis should be initiated immediately after the accident and should be continued for 10 days (Class IIa, level of evidence C). Serologic tests should be performed before initiating treatment and after 20, 40, and 60 days post-treatment to monitor possible seroconversion. In the case of positive serologic results, conventional antiparasitic treatment should be provided, as described previously for the acute phase.

In minimal risk situations, such as the contact with the blood of a patient with Chagas disease in chronic phase, drug prophylaxis is not indicated. Serologic tests should be performed immediately after the accident, and 20, 40, and 60 days post-infection. If seroconversion occurs, conventional treatment for acute phase of Chagas disease should be given, and post-therapeutic monitoring should be conducted as in the acute phase. If the serology remains positive after treatment, the therapeutic gap should be documented and repeat treatment provided, either with the same drug or with an alternative drug.

**Treatment options, dosage, and type of administration**

The neglected character of Chagas disease is evidenced by the limited therapeutic options; only two antiparasitic drugs with established efficacy are available for specific treatment: benznidazole (a nitroimidazole derivative agent) and nifurtimox (a nitrofuran compound). The drug of first choice for the antiparasitic treatment of Chagas disease in all the situations previously discussed is benznidazole (20) (38) (100) (294). Currently, this medicine is produced by a Brazilian pharmaceutical company, distributed by request to the Ministry of Health of Brazil, with no distribution to the general pharmaceutical market.

Benznidazole is a well-tolerated drug. Children generally have fewer adverse effects than adults and can tolerate higher doses. In April 2013, the 19th Expert Committee on the Selection and Use of Essential Medicines of the WHO discussed and approved the addition of two new formulations of benznidazole in the essential drugs list for children (38). Benznidazole is being increasingly prescribed for the antiparasitic treatment of Chagas disease, not only the expansion of indications based on evidence, but also as a result of increased demand in endemic and non-endemic countries with better structure of their national surveillance and healthcare systems.

Benznidazole (Class 1, level of evidence b) is produced as tablets of 100mg and 50mg (adults) and 12.5mg and 50mg (children) (20) (42) (104) (109) (122) (291) (296) (299) (302) (312) (313). In Brazil, only the 100mg and 12.5mg tablets are available in the SUS network.

- **Adults:** 5mg/kg/day, orally, in two or three daily doses, for 60 days.
- **The maximum dose recommended is 300 mg/day.**

The 100mg and 50mg benznidazole tablets are distributed in Latin America by the Argentine Endocrine Laboratory (ELEA, Maprimed) and the PAHO Revolving Fund; in Spain, through the Spanish Agency for Medicines and Health Products; and the rest of the world through the ELEA and the WHO. In Brazil, it is acquired by the Ministry of Health, and distributed to State Health Secretariats (SES) upon request in the Information Strategic Procurement System (SIES). Benznidazole distribution flow (100mg) for regional and/or municipalities should be established for each SES, articulating Pharmaceutical Assistance, Epidemiological Surveillance and Primary Care. The distribution of benznidazole in doses of 12.5mg is centralized, considering the small occurrence of cases of the disease in the recommended age for use of the drug. Therefore, to apply for a pediatric formulation, the Chagas Disease Technical Group, Communicable Disease Surveillance Coordination of the Secretariat of Health, Ministry of Health should be contacted.

- **Children:** 10mg/kg/day orally in two or three daily doses, for 60 days.
- **In all cases, the dose should not exceed 300mg/day.** When the daily dose exceeds 300mg, it is recommended that the treatment duration be extended to achieve the total dose calculated for 60 days.

The main advantage of the 12.5mg tablet is that it can be used to treat infants as young as 2 years of age, allowing proper dosing without the need to fragment and manipulate parts of larger tablets. In addition, the main advantage of 50 mg tablet is that it can be used to treat the rest of the pediatric population, including teenagers and young adults.

In addition to the specific treatment for *T. cruzi* infection, antiparasitic drugs can be used for prophylaxis in specific situations. Secondary prophylaxis should be considered after conventional treatment of reactivation in patients co-infected with HIV, and with evidence of AIDS (2.5 to 5mg/kg/day, orally, three times per week), but there is still no consistent evidence to validate the use of benznidazole prophylactically in this situation.

In turn, primary prophylaxis is indicated in situations where accidental exposure to *T. cruzi* has occurred. The recommended dose is 5mg/kg/day (adult) or 10mg/kg/day (child), orally, twice a day for 10 days, not exceeding 300mg/day. In cases where the inoculum has a high parasite load (≥10^4 trypomastigotes/mL), one should use conventional treatment for acute phase of Chagas disease for at least 60 days (Class IIa, level of evidence C). In the case of children, one should use the pediatric formulation of benznidazole, that is now available.

The adverse effects and toxicity of benznidazole, together with the respective actions to be taken, are shown in Figure 18. Thus, care should be taken before and during treatment to monitor these events. Before starting treatment, clinical examination and laboratory tests (complete blood count, evaluation of liver enzymes, and renal function tests) must be performed. This must be repeated 30 and 60 days after initiating treatment.

In cases of intolerance to benznidazole, nifurtimox is a potential therapeutic option (Class 1, level of evidence B). Nifurtimox comes as a 120mg tablet (adults) and a 30mg tablet (children).

- **Adults:** 10mg/kg/day, orally in three daily doses, for 60 days.
- **Children:** 15mg/kg/day, orally in three daily doses, for 60 days.
The adverse effects and toxicity of nifurtimox are similar to those of benznidazole, except for a lower digestive tolerance, reflected in anorexia with weight loss, and psychological disturbances\(^{(20)}\). In case of treatment failure with benznidazole, nifurtimox can be used, although cross-resistance between the two drugs has been observed.

It is necessary to remember that nifurtimox is not available in the pharmaceutical market of any country to date. In situations where the use of nifurtimox is essential, the drug must be requested via the PAHO and WHO offices. The Ministry of Health of Brazil, through the SVS Chagas Disease Technical Group, has the availability of nifurtimox to be dispensed under the conditions described above of benznidazole intolerance. Nifurtimox is acquired by the Ministry of Health via PAHO donation and is dispensed directly by the Chagas disease Technical Group of SVS\(^{(39)}\). The product may be requested through a standardized protocol directed to this technical group by the Ministry of Health.

Other drugs, such as allopurinol and azole antifungals (ketoconazole, itraconazole, fluconazole, posaconazole), are recognized as suppressors of *T. cruzi* parasitemia and may be useful in some specific situations, such as reactivation of}
in immunosuppressed patients, when it is impossible to use benznidazole or nifurtimox (Class IIa, level of evidence C). However, there is undeniable clinical evidence of proven therapeutic efficacy.

The antiparasitic treatment with benznidazole should not be imposed on pregnant women, or women of childbearing age who are not making a regular use of well-known contraceptive methods. As discussed in the specific section on vertical transmission of *T. cruzi* in this Consensus, there is evidence indicating the benefit of the antiparasitic treatment, with better evolution of Chagas disease, reduced risk of vertical transmission in future pregnancies, and lower risk of vertical transmission and evolution to spontaneous abortion by reducing the parasitemia.

In the case of pregnant women for whom antiparasitic treatment is indicated, the risk-benefit ratio and available treatment options (in particular, acute phase and co-infection) should be considered in each case. The indication in patients with other serious disorders, such as hepatic and renal failure, must be carefully assessed case by case, according to the severity, as well as prior adverse events to drug components that may constitute relative contraindications.

**Evaluation of cure of Chagas disease after antiparasitic treatment**

Demonstrating serology becoming negative is considered the only method to determine cure following antiparasitic treatment of Chagas disease. The time required for this to

<table>
<thead>
<tr>
<th>Event</th>
<th>Recommendation class</th>
<th>Level of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antiparasitic treatment in the acute phase of Chagas disease</td>
<td>I</td>
<td>B</td>
</tr>
<tr>
<td>Antiparasitic treatment in congenital Chagas disease</td>
<td>I</td>
<td>B</td>
</tr>
<tr>
<td>Antiparasitic treatment in the chronic phase of Chagas disease in children aged ≤12 years</td>
<td>I</td>
<td>A</td>
</tr>
<tr>
<td>Antiparasitic treatment in the chronic phase of Chagas disease with recent infection</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Antiparasitic treatment in the chronic phase of Chagas disease with late infection, acquired in the indeterminate form</td>
<td>IIa</td>
<td>B</td>
</tr>
<tr>
<td>Antiparasitic treatment in the chronic phase of Chagas disease with late infection and cardiomyopathy without advanced disease</td>
<td>IIb</td>
<td>C</td>
</tr>
<tr>
<td>Antiparasitic treatment in the chronic phase of Chagas disease with advanced cardiac form of the disease</td>
<td>III</td>
<td>C</td>
</tr>
<tr>
<td>Digestive form does not contraindicate antiparasitic treatment of Chagas disease</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Antiparasitic treatment in an organ donor with Chagas disease</td>
<td>I</td>
<td>C</td>
</tr>
<tr>
<td>Antiparasitic treatment in an organ recipient without Chagas disease from organ donor with Chagas disease</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Monitoring of clinical and parasitologic events instead of antiparasitic treatment in an organ recipient without Chagas disease from an organ donor with Chagas disease</td>
<td>IIa</td>
<td>B</td>
</tr>
<tr>
<td>Reactivation prophylaxis in transplanted person with Chagas disease</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Monitoring of clinical and parasitologic events and prophylactic antiparasitic treatment for transplanted person with Chagas disease</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Coinfection Chagas/HIV without AIDS: antiparasitic treatment according to criteria in immunocompromised</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Reactivation antiparasitic treatment in coinfection Chagas/HIV</td>
<td>I</td>
<td>C</td>
</tr>
<tr>
<td>Antiparasitic treatment in coinfection Chagas/HIV without clinical events and high <em>T. cruzi</em> parasitemia</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Secondary prophylactic treatment after reactivation antiparasitic treatment in coinfection Chagas/HIV in individuals with CD4+ lymphocyte T count &lt;200 cells/mm³</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Primary prophylaxis in laboratory accident with high risk of transmission of Chagas disease</td>
<td>IIa</td>
<td>C</td>
</tr>
<tr>
<td>Benznidazole as first choice drug for antiparasitic treatment of Chagas disease</td>
<td>I</td>
<td>B</td>
</tr>
<tr>
<td>Nifurtimox as a therapeutic option (instead of benznidazole) for the antiparasitic treatment of Chagas disease</td>
<td>I</td>
<td>B</td>
</tr>
<tr>
<td>Allopurinol and azole antifungals as reactivation treatment in immunosuppressed patients when benznidazole and nifurtimox treatment are not possible.</td>
<td>IIa</td>
<td>C or unknown</td>
</tr>
<tr>
<td>Serology as a criterion of cure and therapeutic failure after antiparasitic treatment for Chagas disease</td>
<td>IIa</td>
<td>B</td>
</tr>
<tr>
<td>PCR as an alternative to indirect parasitologic examination (artificial xenodiagnosis and blood culture) as an indication of therapeutic failure after specific treatment for Chagas disease</td>
<td>IIa</td>
<td>B</td>
</tr>
</tbody>
</table>

**FIGURE 19. Summary of treatment situations, the recommendation class and level of evidence.**
occur is variable and depends on the stage and duration of disease, with one year for congenital infection, 3–5 years for the acute phase, 5–10 years for the recent chronic phase and >10 years (usually 20–25 years) for the chronic phase of long duration. The reduction in serologic test titers occurs gradually. The persistent and progressive decline above three dilutions of the titers of the serologic tests is suggestive that these will be negative. Parasitologic examinations are not mandatory for evidence of cure of Chagas disease, however, at any time, positive parasitologic tests indicate therapeutic failure. PCR is an alternative option to indirect methods (xenodiagnosis and blood culture) in the parasitologic evaluation as a treatment failure criterion after completing antiparasitic treatment\(^{104}\)\(^{(313)}\). However, there are still limitations on the availability and standardization of techniques in the health networks of endemic countries\(^{313}\).

**Who and where to treat**

Considering the operational specificities of national health systems, use of antiparasitic treatment for Chagas disease by doctors working in primary care who know the peculiarities of the drugs and Chagas disease\(^{(20)}\)\(^{(303)}\), is feasible, safe\(^{(20)}\)\(^{(251)}\)\(^{(294)}\)\(^{(303)}\)\(^{(312)}\), and operationally possible\(^{(294)}\). Depending on the severity of clinical conditions of each case, particularly cases with acute or reactivated and decompensated chronic forms, it may be necessary to refer certain patients to more specialized or reference units that have greater technological capability, or even for hospitalization.

Figure 19 provides an overview of potential situations for specific treatment, taking as reference the recommendation of class and the level of existing evidence. Figure 18 shows the adverse effects of benznidazole and recommended approaches for each given situation.

**Tribute**

Tribute is paid to members of the group responsible for the Brazilian Consensus on Chagas Disease in 2005 who died in the period between the consensuses: Ademir Rocha, Aluízio Rosa Prata, Antonio Carlos Silveira, Guilherme Rodrigues da Silva, Joffre Marcondes de Rezende, and Vanize de Oliveira Macedo.

**Contributions**


**Acknowledgements**

We thank Dr. Jarbas Barbosa da Silva Jr., Secretary of Surveillance in Health of the Ministry of Health in the periods 2003–2007 and 2011–2014, for supporting the construction of two issues of the Consensus on Chagas Disease, 2005 and 2015. To Dr. Habib Fraiha Neto and Dr. Ralph Lainson (in memoriam), for the contributions in the section dedicated to oral transmission. To the Brazilian Network of Attention and Studies on Trypanosoma cruzi/HIV Coinfection and other immunosuppressive conditions. We thank Dr. Pedro Albajarin-Viñas (World Health Organization - HIV/AIDS, Tuberculosis, Malaria and Neglected Diseases - Control of Neglected Tropical Diseases - Innovative & Intensified Disease Management - Chagas disease) for the support of the translation to the english version of the Brazilian Consensus on Chagas disease.

**Conflicts of Interest**

The authors declare that they have no conflicts of interest.

**Financial Support**

Technical Surveillance Unit of Vector-Borne Diseases, Secretariat of Health Surveillance, Ministry of Health of Brazil, Brasília, Distrito Federal, Brazil. The English version of the original article in Brazilian Portuguese of this Consensus was made possible through a partnership with the World Health Organization.

**REFERENCES**


Definitions of grades or classes of recommendations and levels of evidence

Classes of recommendations

- **Class I**: conditions for which there is conclusive evidence, or, in its absence, general agreement that the procedure or treatment is useful and/or effective.
- **Class II**: conditions for which there is conflicting evidence and/or divergence of views on the utility/effectiveness and safety of the procedure or treatment:
  - **Class IIa**: evidence and opinions favor the indication for the procedure or treatment; most professionals involved in specific management approve;
  - **Class IIb**: utility/effectiveness and safety is less well established by evidence, with opinion divided; procedure or treatment is considered optional.
- **Class III**: conditions for which there is conclusive evidence and/or general agreement that the procedure/treatment is not useful and/or effective, and may even be harmful in some circumstances.

Levels of Evidence

- **Level A**: data from several consistent and good quality randomized trials or robust meta-analysis of randomized clinical trials.
- **Level B**: data obtained from a single randomized clinical trial or from several non-randomized clinical studies, observational (less robust meta-analysis).
- **Level C**: data obtained from consensual opinions of specialists on the subject.

1Based on international publications and particularly endorsed in the First Latin American Guidelines for the Diagnosis and Treatment of Chagas Heart Disease.

Evidence levels ranked as B or C cannot be interpreted as weak recommendations. There are many consensus-based recommendations, so with the recommendation of class I, level of evidence C (expert opinions). On the other hand, some indications considered controversial (recommendation grade II) may be grounded in randomized controlled trials (level of evidence A).
Erratum

Revista da Sociedade Brasileira de Medicina Tropical/Journal of the Brazilian Society of Tropical Medicine
49(Supplement I): 2016 - doi: 10.1590/0037-8682-0505-2016 - Page 22 - Figure 2

Should read:

PLASMA OR SERUM SAMPLES

2 TESTS, DIFFERENT METHODS
(ELISA, INDIRECT IFA/HAI)

1 TEST REACTIVE
1 TEST NON-REACTIVE

COLLECT NEW SAMPLE

REPEAT 2 TESTS, DIFFERENT METHODS
(ELISA, INDIRECT IFA/HAI)

1 TEST REACTIVE
1 TEST NON-REACTIVE

POSITIVE RESULT

INCONCLUSIVE RESULT

NEGATIVE RESULT

FORWARD SAMPLES TO REFERENCE LABORATORY

ELISA: enzyme-linked immunosorbent assay
IFA: indirect fluorescent antibody test
HAI: indirect hemagglutination

Should read:

PLASMA OR SERUM SAMPLES

2 TESTS, DIFFERENT METHODS
(ELISA, INDIRECT IFA/HAI)

1 TEST REACTIVE
1 TEST NON-REACTIVE

COLLECT NEW SAMPLE

REPEAT 2 TESTS, DIFFERENT METHODS
(ELISA, INDIRECT IFA/HAI)

1 TEST REACTIVE
1 TEST NON-REACTIVE

POSITIVE RESULT

INCONCLUSIVE RESULT

NEGATIVE RESULT

FORWARD SAMPLES TO REFERENCE LABORATORY

ELISA: enzyme-linked immunosorbent assay
IFA: indirect fluorescent antibody test
HAI: indirect hemagglutination