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# **Review Article**

**Cite this article:** Santana Nogueira S, Cardoso Santos E, Oliveira Silva R, Vilela Gonçalves R, Lima GDA, Dias Novaes R (2022). Monotherapy and combination chemotherapy for Chagas disease treatment: a systematic review of clinical efficacy and safety based on randomized controlled trials. *Parasitology* **149**, 1679–1694. https://doi.org/10.1017/ S0031182022001081

Received: 5 May 2022 Revised: 26 July 2022 Accepted: 27 July 2022 First published online: 12 August 2022

#### Key words:

Antiparasitic chemotherapy; Chagas disease; parasitology; *Trypanosoma cruzi* 

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# Monotherapy and combination chemotherapy for Chagas disease treatment: a systematic review of clinical efficacy and safety based on randomized controlled trials

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#### Abstract

From a systematic review framework, we analysed the clinical evidence on the effectiveness and safety of monotherapy and combination chemotherapy for Chagas disease (ChD) treatment. The research protocol was based on the Preferred Reporting Items for Systematic Reviews and Meta-Analyses and patient, intervention, comparison and outcome strategy. Only randomized controlled trials (RCT) were retrieved from Embase, Medline, Scopus and Web of Science databases. Diagnostic tools, treatment protocols, seroconversion rates and adverse events were investigated. Fifteen RCT mainly concentrated in endemic countries were identified. ChD diagnosis was mainly based on haemagglutination, immunofluorescence, enzyme-linked immunosorbent assay and polymerase chain reaction. Benznidazole (BNZ), nifurtimox, fosravuconazole, posaconazole, allopurinol and thioctic acid were the identified drugs. The best negative seroconversion results (100, 96, 94 and 91.3%) were, respectively, based on BNZ (5 mg kg day<sup>-1</sup>, 200 mg day<sup>-1</sup>, 150 mg day<sup>-1</sup> and 2.5 mg kg<sup>-1</sup>) administration for 60 days. Negative seroconversion was not achieved with allopurinol  $(300 \text{ mg day}^{-1} \text{ for}$ 60 days). Adverse reactions ranged from 5 to 73% in patients receiving antiparasitic chemotherapy. Treatment discontinuation (1.5-57%) was mainly associated with gastrointestinal, cutaneous and neurological manifestations. Current RCT-based evidence indicates that BNZ is the most viable option for ChD treatment. However, new protocols need to be developed to mitigate side effects and increase patient adherence to antiparasitic chemotherapy. Therefore, shorter regimens, lower concentrations and treatments combining BNZ with posaconazole, fosravuconazole or ravuconazole may be viable to ensure comparable efficacy to BZN-based monotherapy, contributing to reduce dose- and time-dependent toxicity reactions.

#### Introduction

Chagas disease (ChD) or American trypanosomiasis is a life-threatening anthropozoonosis markedly related to poverty and caused by the protozoan parasite *Trypanosoma cruzi* (Bern, 2015; Torrico *et al.*, 2021). About 5.7–9.4 million people are infected with *T. cruzi* worldwide (Echeverría *et al.*, 2020*a*; Torrico *et al.*, 2021), especially in 21 Latin American endemic countries and 19 non-endemic countries in North America and Europe (Bivona *et al.*, 2020; Torrico *et al.*, 2021). Vector and oral transmission are the main forms of *T. cruzi* infection in endemic countries. However, donation of contaminated blood and organs from infected people, vertical transmission (e.g. mother to fetus) and laboratory accidents are the main causes of ChD spread in non-endemic areas (Nogueira *et al.*, 2018; Guhl and Ramirez, 2021).

ChD is clinically divided into acute and chronic phases (Bern, 2015). The acute phase courses with intense parasitaemia and marked cellular parasitism in multiple organs, especially the heart, skeletal muscles and cells of the phagocytic mononuclear system (Bivona *et al.*, 2020; Torrico *et al.*, 2021). However, acute infections are often asymptomatic and associated with low mortality rates (0.2–0.5%) (Pérez-Molina and Molina, 2018; Echeverría *et al.*, 2020*a*). Once parasitaemia is controlled, the disease progresses to the chronic phase, which may remain asymptomatic for decades or evolve into a symptomatic form associated with digestive, nervous and/or cardiovascular manifestations (Bern, 2015). Cardiac involvement characterizes chronic Chagas cardiomyopathy (CCC), which is the most severe form of ChD (Nogueira *et al.*, 2018; Bivona *et al.*, 2020).

Chronic cardiomyopathy is the leading cause of ChD-associated mortality (Nogueira *et al.*, 2018; Caldas *et al.*, 2019). In addition, CCC is the most common infectious cardiomyopathy

worldwide and the third most frequent cause of heart transplantation in endemic countries (Nogueira *et al.*, 2018). CCC exhibits a multifactorial and complex aetiology. It is often associated with parasite persistence, low-grade inflammation, autoimmunity, redox imbalance, thromboembolic events, myonecrosis, autonomic denervation and progressive myocardial fibrosis (Rassi *et al.*, 2017; Rodrigues *et al.*, 2017; Bonney *et al.*, 2019). Together, these events contribute to CCC progression, which often manifests electromechanical abnormalities such as conduction defects (e.g., bundle branch blocks), frequent and complex ventricular arrhythmias, and systolic ventricular dysfunction (Bonney *et al.*, 2019; Caldas *et al.*, 2019).

The specific ChD treatment is limited to the nitroheterocyclic compounds nifurtimox (NFx) and benznidazole (BNZ) (Muñoz et al., 2011; Diniz et al., 2013; Novaes et al., 2016), which was developed almost 50 years ago (Muñoz et al., 2011, Diniz et al., 2013). Due to high toxicity and serious side effects (e.g. hypersensitivity reactions, anorexia, vomiting, polyneuritis and bone marrow depression), NFx is no longer available in most endemic countries (Urbina and Docampo, 2003; Caldas et al., 2019). Thus, BNZ becomes the first-line drug for ChD treatment (Caldas et al., 2019). Despite its limitations (e.g. systemic toxicity, prolonged treatment and limited efficacy in chronic infections), the risk-benefit of BNZ-based chemotherapy is still favourable (Muñoz et al., 2011; Caldas et al., 2019), especially in acute infections where high cure rates can be obtained (Martinez et al., 2020; Caldas et al., 2019). However, side effects often dictate treatment discontinuation, negatively influencing chemotherapy effectiveness and cure rates (Santos et al., 2015; Torrico et al., 2021).

Considering the limited effectiveness of the classic BNZ-based protocol in improving cardiac function, sustainably attenuating tissue parasitism, and achieving parasitological cure in chronic ChD (Molina-Morant et al., 2020; Martín-Escolano et al., 2020); developing more efficient therapeutic regimens is an urgent need (Molina et al., 2014; Vallejo et al., 2016; Torrico et al., 2021). Thus, repositioning and combining drugs with antiparasitic effects and different mechanisms of action have emerged as potential alternatives to improve ChD treatment (Morillo et al., 2017; Mendonça et al., 2020; Torrico et al., 2021). In addition to new NFx- and BNZ-based protocols, clinical trials currently registered on ClinicalTrials.gov for ChD treatment are based on drug repositioning, which admits that: (i) distinct diseases may share similar targets, (ii) drugs act on multiple targets and (iii) identifying new targets for a drug contributes to innovative therapeutic applications (Sardana et al., 2011; Reddy and Zhang, 2013). Allied to this repositioning strategy, drug combinations have been investigated in search of synergistic or additive interactions to increase the effectiveness of anti-T. cruzi chemotherapy (Mazzeti et al., 2019; Martinez et al., 2020). As combinations can increase drug half-life, reduce dose and treatment time, minimize systemic toxicity and side effects; this strategy can also increase chemotherapy adherence and improve clinical outcomes in ChD patients (Muñoz et al., 2011; Rassi et al., 2017; Torrico et al., 2021).

Currently, evidence on the types and effectiveness of different chemotherapy protocols applied to ChD treatment is based on diffuse initiatives. Thus, it becomes difficult to identify which drugs and combinations were investigated in clinical trials to delimit the most efficient protocols and potential risks associated with drug therapy. Therefore, we used a systematic review framework to map the evidence on specific ChD treatments included in randomized controlled trials (RCT). In addition to identifying the drugs administered (pharmacological class and dosimetry parameters), the effectiveness of the available chemotherapeutic protocols to control the parasite load, achieve negative seroconversion and therapeutic cure was investigated. Considering the pharmacological safety, adverse reactions and treatment discontinuation rates were also evaluated. Finally, the methodological quality of all studies reviewed and potential sources of bias associated with current evidence were objectively characterized, contributing to the refinement of further investigations.

## Methods

#### Guiding question and protocol registration

Our research protocol was developed considering the patient, intervention, comparison and outcome (PICO) strategy (Huang *et al.*, 2006), which was used to define the following guiding question adopted in this systematic review: Could ChD patients undergoing different chemotherapy regimens exhibit better parasitological control, cardiovascular function, lower rates of adverse reactions and mortality compared to untreated control patients? The present review protocol has been registered within the PROSPERO (International Prospective Register of Systematic Reviews) database (registration number CRD42021276800).

#### Search strategy and primary studies selection

Study selection was based on the updated Preferred Reporting Items for Systematic Reviews and Meta-Analyses strategy (PRISMA) (Page et al., 2021). The retrieval of indexed studies on the aetiological treatment for ChD was operationalized from 2 complementary strategies: (i) electronic database search and (ii) indirect searches in citations and/or reference lists of all studies identified from electronic databases (Pereira et al., 2017). The direct strategy was based on advanced searches in PubMed/ Medline, Embase, Scopus and Web of Sciences databases. To identify relevant studies, search filters were initially built using the MeSH Terms, which are standardized descriptors obtained from the PubMed thesaurus (https://www.ncbi.nlm.nih.gov/ mesh/) (Felizardo et al., 2018). The search filters were developed considering 2 levels: (i) disease (ChD) and (ii) intervention (aetiological treatment). All descriptors were associated with specific algorithms [(MeSH Terms) and (TIAB)] to optimize the retrieval of relevant studies in PubMed/Medline (Altoé et al., 2019), ensuring the identification of research records indexed and in the indexing process (Souza-Silva et al., 2019). The 'human' and 'clinical trials' search limits provided by the search engine were applied to refine the quality of the PubMed/Medline search.

The same search filters built for the PubMed/Medline database were adapted for Embase, Scopus and Web of Science. Therefore, the respective syntax and algorithms required in the search engine of each database were used, such as: de,ab,ti, TITLE-ABS-KEY or TS =. To refine the quality of the Embase search, studies also indexed in Medline were automatically excluded from the Venn diagram tool (Sources tab) (Silva et al., 2018), selecting studies exclusively indexed in Embase. In addition, the standardized limits 'human', 'major clinical study' and 'article' were used in our search strategy applied to Embase. The search limits 'human' and 'AND NOT INDEX (Medline)' were applied to the Scopus database. This last limit was used to exclude duplicate Medline studies from the results. For the Web of Science database, a filter was built to select clinical studies and specific animal species (humans). No chronological or language limits were used (Silva et al., 2018). The complete search strategies used in each database can be accessed in the supplementary material (Table S1).

# Studies screening, eligibility criteria and inter-rater agreement

Only clinical studies investigating the impact of ChD aetiological treatment were included in this systematic review. All relevant

studies were selected according to the PRISMA flowchart (Page et al., 2021). Two reviewers (S.S.N. and R.O.S.) retrieved and independently applied search strategies across all databases (Altoé et al., 2019). Disagreements were resolved by arbitration in consultation with an expert researcher (R.D.N). Duplicate studies that were not directly excluded by the search algorithms were removed using Mendeley software (Mendeley Desktop Version 1.19.8). After this step, the titles and abstracts of all research records were screened and irrelevant records (not related to the investigated topic) were excluded. The remaining studies were retrieved in full text and well-defined eligibility criteria were analysed. The exclusion criteria applied in this review were: (i) grey literature (not peerreviewed and formally published), (ii) studies unavailable in full text (title and/or abstract only), (iii) studies with nonpharmacological interventions, (iv) studies with multiple interventions where it was not possible to isolate the aetiologic treatment effect, (v) secondary studies (literature reviews, comments, letters to the editor and editorials), (vi) studies without control groups and (vii) studies unrelated to parasitological control. After selecting all relevant studies from the primary search, reference lists were screened to identify additional studies (Felizardo et al., 2018; Nogueira et al., 2018). These studies were retrieved in full text and the same eligibility criteria used in the direct search were analysed in this secondary strategy. The results obtained from the primary and secondary searches were compared and the inter-rater agreement (Kappa coefficient) was calculated (McHugh, 2012).

#### **Data extraction**

Qualitative and quantitative data were extracted from all relevant studies included in the systematic review. To this end, we use standardized spreadsheets (data extraction masks) (Marques *et al.*, 2018) structured from basic methodological requirements to characterize studies at different descriptive levels, such as: (i) publication characteristics: research design, authors, year of publication and country where the study was conducted; (ii) patient characteristics: age, sex, and disease stage; (iii) treatment characteristics: drugs and dosimetry (doses, administration frequency and rout, treatment duration, and patient follow-up); (iv) primary outcomes: parasite load (parasitological cure), seroconversion and mortality rates; (v) secondary outcomes: cardiovascular function (electrocardiographic and echocardiographic data), adverse reactions and treatment discontinuation rates.

#### Reporting quality as a risk of bias

Methodological quality and potential risk of bias in all studies reviewed were analysed using the Downs and Black (D&B) checklist, which is targeted at randomized and non-randomized trials of health care interventions (Downs and Black, 1998). The scale is based on 27 questions and is structured in 5 categories or domains, such as: (i) reporting quality, (ii) external validity, (iii) bias, (iv) confounding and (v) statistical power. This scale has high test-retest reliability (r = 0.88) and internal consistency (KR20 formula = 0.89). Due to previous recommendations and high ambiguity, question 27 (statistical power) was not applied (Nogueira *et al.*, 2018). The overall result obtained from the Downs and Black checklist was expressed graphically, and the average score was calculated (Nogueira *et al.*, 2018).

### Results

#### **Publication characteristics**

In the primary search, 508 papers published between 1996 and 2021 were retrieved from PubMed, Embase, Scopus and Web of

Science. After removing duplicates and evaluating eligibility criteria, 11 relevant papers were identified. Four additional papers were identified in the secondary search. Therefore, 15 original studies were included in the systematic review. Most studies (n = 10, 66.6%) were developed in Latin American countries. Three (20%) multicentre studies were also identified, followed by 2 studies developed in Spain (13.3%). The list of all papers included and the PRISMA flowchart with the complete strategy are shown in Table S2 and Fig. 1, respectively. The Kappa coefficient obtained from our search strategy ( $\kappa = 0.842$ ) indicated substantial agreement between independent evaluators (Table S2).

#### **Patient characteristics**

From all the studies reviewed, 6 (40%) classified participants by sex, and female patients were investigated in 3 studies (26.6%). Men were exclusively recruited in 1 study (6%), and gender was not reported in 4 studies (26.6%). In general, the patients' ages ranged from 7 to 75 years, and 1 study (6%) investigated newborns. Body mass was underreported in most studies (n = 12, n)80%). Considering disease stage, 11 studies (73.3%) included patients in the indeterminate chronic phase. One study (6.6%) investigated congenital disease (acute phase), and the disease phase was not reported in 3 studies (20%). ChD diagnosis was based on indirect haemagglutination assay (IAH), indirect immunofluorescence assay (IFA), enzyme immunoassay (EIA), enzyme-linked immunosorbent assay (ELISA), or chemiluminescence immunoassay (CL-ELISA, A&T antigen) in 6 studies (40%). Polymerase chain reaction (PCR) was also used in 6 studies (40%). Xenodiagnostic alone or combined with IFA was applied in only 1 study (6%) each (Table 1).

#### Chemotherapy protocols, seroconversion and cure rates

As indicated in Table 2, BNZ was the drug mainly investigated in monotherapy in 8 studies (53.3%). Monotherapy based on NFx, allopurinol, posaconazole or the prodrug ravuconazole (E1224) was reported in 7 studies (46.6%). Drug combinations based on BNZ + thioctic acid, BNZ + posaconazole and BNZ + fosravuconazole were investigated in 3 studies (20%). Negative seroconversion rates reported in Latin American studies using BNZ were: 11.2–82.2% at 5 mg kg day<sup>-1</sup>, 58–88.7% at 7.5 mg kg day<sup>-1</sup>, 91.3% at 2.5 mg kg day<sup>-1</sup> and 94% at 150 mg day<sup>-1</sup>. Two other BNZ studies (developed in Canada and Spain), found 96%  $(200\ mg\ day^{-1})$  and 100%  $(5\ mg\ kg\ day^{-1})$  seroconversion rates, respectively. With respect to other treatments, seroconversion rates ranging from 10.9 to 28.9% were observed for E1224 (2000 and 4000 mg total<sup>-1</sup>), and 10–20% for posaconazole (100 and  $400 \text{ mg day}^{-1}$ ). Seroconversion was not achieved with allopurinol treatment. For drug combinations, a high seroconversion rate (96%) was obtained in patients treated with BNZ + posaconazole, while an 84% rate was achieved in patients receiving BNZ + fosravuconazole. This parameter was not investigated in studies with NFx and BNZ + thioctic acid. The parasitological cure was estimated from T. cruzi DNA detection in blood samples by PCR. This method was used in 6 studies (40%). Negative PCR ranged from 46.7 to 100% in BNZ-treated patients, 8.3-28.9% for E1224 treatment and 10-20% for posaconazole treatment. For drug combinations, negative PCR were obtained in 96% patients receiving BNZ + posaconazole, and 84% for patients treated with BNZ + fosravuconazole. Only 2 studies (13.3%) investigated potential changes in PCR results over time or over a year (Vallejo et al., 2016; Morillo et al., 2017). Negative PCR results were maintained in 46.7% to 100% of all patients investigated in a follow-up ranging from 1 to 5 years (Table 2).



Fig. 1. Flow diagram of the systematic review literature search results. Based on the PRISMA statement 'Preferred Reporting Items for Systematic Reviews and Meta-Analyses-http://www.prisma-statement.org/'.

Table 1. Patient characteristics, ChD stage and diagnostic method of Trypanosoma cruzi infection used in randomized clinical trials

Author and year of publication	Country	Sample size	Sex	Age /min-max (mean)	Clinical stage of disease	Methods used in ChD diagnostic
De Andrade <i>et al.</i> (1996)	Brazil	130	M/F	≥07 and $≤12$	(-)	IHA, IFA, ELISA
Coura et al. (1997)	Brazil	77	(–)	(–)	Chronic	IFA, Xenodiagnosis
Sosa Estani <i>et al.</i> (1998)	Argentina	106	(–)	<13	Chronic indeterminate	IHA, IFA, EIA
Andrade et al. (2004)	Brazil	129	M/F	$\geqslant$ 14 and $\leqslant$ 19	(-)	A&T CL-ELISA
Sosa-Estani et al. (2004)	Argentina	249	M/F	$\geqslant$ 15 and $\leqslant$ 44	Chronic indeterminate	IHA, IFA, ELISA
Rassi et al. (2007)	Brazil	35	M/F	$\geqslant$ 18 and $\leqslant$ 64	Chronic indeterminate	Xenodiagnosis
Marin-Neto et al. (2008)	Multicentre	3000	M/F	≥18 and ≤75	(-)	IHA, IFA, ELISA
Chippaux et al. (2010)	Bolivia	111	(-)	Newborn	Congenital	ELISA
Molina <i>et al.</i> (2014)	Bolivia	78	M/F	39 ± 9	Chronic	RT-PCR
Morillo <i>et al.</i> (2015)	Multicentre	2854	M/F	$\geqslant$ 18 and $\leqslant$ 75	Chronic	RT-PCR
Vallejo <i>et al.</i> (2016)	Spain	14	M/F	$\geqslant$ 26 and $\leqslant$ 57	Chronic indeterminate	RT-PCR
Morillo et al. (2017)	Canada	120	М	$\geqslant$ 18 and $\leqslant$ 54	Chronic indeterminate	RT-PCR
Torrico et al. (2018)	Bolivia	231	M/F	$\geqslant$ 18 and $\leqslant$ 50	Chronic indeterminate	RT-PCR
Molina-Morant et al. (2020)	Multicentre	240	(–)	≥ 18	Chronic	IFA, IHA, ELISA
Torrico et al. (2021)	Bolivia	210	M/F	$\geqslant$ 18 and $\leqslant$ 50	Chronic indeterminate	RT-PCR

-, data not reported; M, male; F, female; IHA, indirect haemagglutination assay; IFA, indirect immunofluorescence assay; EIA/ELISA, enzyme-linked immunosorbent assay; A&T CL-ELISA ELISA, chemiluminescent ELISA with A&T antigen; RT-PCR, real-time reverse transcription polymerase chain reaction.

# Cardiovascular and laboratory outcomes, treatment discontinuation and adverse effects

As indicated in Table 3, pre-treatment and post-treatment cardiovascular parameters were reported in 9 studies (60%). Right bundle branch block, left anterior hemiblock, atrioventricular block, ectopic rhythm, atrial fibrillation, ventricular arrhythmia, ventricular tachycardia, stroke, transient ischaemic attack, systemic embolism, pulmonary embolism, pacemaker or implantable cardioverter-defibrillator, cardiac arrest and transplantation were the main cardiovascular abnormalities/outcomes reported. No study reported significant improvement or worsening of cardiac function in patients treated with the different drugs and therapeutic regimens investigated.

All treatments were associated with a low frequency of altered liver function estimated from alanine aminotransferase levels (2.2-38%) and reduced white blood cell counts (0.1-43%). Altered liver function, estimated from serum transaminase levels,

Study	Patients / group (n)	Withdrawn <sup>a</sup>	Therapeutic schemes	Dose reported	Administration frequency and rout	Treatment (days)	Follow-up (years)	Seroconversion negative (n/%)
De Andrade et al. (1996)	64	06	BNZ	7.5 mg/kg/ day	b.d.i/oral	60	3	37/64 (58%)
	65	11	PLA	(-)	(-)			3/64 (5%)
Coura <i>et al.</i> (1997)	26	03	BNZ	5 mg/kg/day	b.d.i/oral	30	1	(?)
	27	08	NF	5 mg/kg/day				(?)
	24	02	PLA	(-)				(?)
Sosa Estani <i>et al.</i> (1998)	51	07	BNZ	5 mg/kg/day	(-)	60	2	5/44 (11.2%)
	50	06	PLA	(–)	(-)			2/44 (4.5%)
Andrade et al. (2004)	64	06	BNZ	7.5 mg/kg/day	b.d.i/oral	60	6	47/53 (88.7%)
	65	11	PLA	(–)	(–)			12/46 (26.1%)
Sosa-Estani <i>et al.</i> (2004)	62	35.5%	AT + BNZ	50 + 5 mg/kg/day	b.d.i/oral	1–37 + 5–37	0.14 (53 days)	(-)
	66	33.4%	AT + BNZ	100 + 5 mg/kg/day	b.d.i/oral	1–37 + 5–37		(–)
	59	28.8%	PLA + BNZ	(–) + 5 mg/kg/day	b.d.i/oral	1–37 + 5–37		(-)
	62	21%	PLA + AT + BNZ	(–) + 50 mg + 5 mg/kg/day	b.d.i/oral	1-4+1-37+5-37		(-)
Rassi <i>et al.</i> (2007)	23	06	ALLP	300 mg	3 daily	60	2	0/17 (0%)
	12	02	PLA	(–)	(-)			0/10 (0%)
Marin-Neto <i>et al.</i> (2008)	300	(-)	BNZ	5 mg/kg/day	(-)	40, 60 or 80	2	? (60%)
	300	(-)	PLA	(-)	(-)			? (30%)
Chippaux et al. (2010)	59	01	BNZ	2.5 mg⁄ kg	b.d.i/oral	60	1	53/58 (91.3%)
	52	02	BNZ	7.5 mg⁄ kg	o.d	30		45/50 (90%)
	68	(-)	Untreated	(-)	(-)	(–)		(-)
Molina <i>et al.</i> (2014)	26	09	BNZ	150 mg/day	b.d.i/oral	60	1	16/17 (94,1%)
	26	01	POS	400 mg/day				5/25 (20%)
	26	06	POS	100 mg/day				2/20 (10%)

Table 2. Characteristics of treatments used in the management of Chagas disease patients in randomized clinical trials.

POS, posaconazole; BNZ, benznidazole; PLA, placebo; NF, nifurtimox; ALLP, allopurinol; AT, thioctic acid; b.d.i, twice day; o.d, once a day; -, data not reported; ?, incomplete information. <sup>a</sup>For discontinuity, adverse events and other reasons.

Table 2. (Continued.)

#### Study group (n) Withdrawn<sup>a</sup> schemes Dose reported Administration frequency and rout (days) (years) negative (n/%) Morillo et al. (2015) 1431 ?/14 BNZ $300 \text{ mg day}^{-1}$ o.d 1431 ≥ 5 ? (46.7%) (–) 1423 ?/14 PLA (-) 1423 ? (33.1%) 1.5 BNZ 5 mg kg day<sup>-1</sup> b.d.i/oral Vallejo et al. (2016) 07 03 60 7/7 (100%) (-) (-) 07 01 (-) 3/7 (42.8%) Untreated Morillo et al. (2017) 32 00 POS 400 mg b.d.i/oral 60 1 ? (16%) 30 01 PLA ? (17%) 10 mg 28 09 BNZ + POS ? (96%) 200 mg + 400 mg ? (96%) 30 10 BNZ + PLA 200 mg $4000 \text{ mg total}^{-1}$ Torrico et al. (2018) 45 03 E1224 400 mg/o.d for 1-3 days followed 400 mg/once wk 65 1 13 (28.9%) for 7 wk/oral $2000 \text{ mg total}^{-1}$ 46 (-) 200 mg/o.d for 1-3 days + Placebo followed by E1224 + PLA 4 (8.3%) 200 mg/E1224 and Placebo/once wk for 7 wk/oral 48 03 $2400 \text{ mg total}^{-1}$ 400 mg/once wk for 3 wk followed by Placebo/4 wk E1224 + PLA 5 (10.9%) or Placebo/8 wk/oral 100 mg tablet<sup>-1</sup> 45 03 BNZ 5 mg/kg/day in 2 daily doses/oral 37 (82.2%) 47 01 PLA 400 mg 4 tablets o.d for 1-3 days followed for 4 tablets once 4 (8.5%) a wk for 7 wk/oral

<sup>a</sup>For discontinuity, adverse events and other reasons. E1224 (ravuconazole prodrug).

Patients /

BNZ, benznidazole; POS, posaconazole; PLA, placebo; o.d, once a day; b.d.i, twice day; wk, week; -, data not reported; ?, incomplete information.

Therapeutic

Treatment

Follow-up

Seroconversion

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Study	Patients/ group (n)	Withdrawn <sup>a</sup>	Therapeutic schemes	Dose reported	Administration frequency and rout	Treatment	Follow-up (years)	Seroconversion negative $(n/\%)$
Molina-Morant <i>et al.</i> (2020)	80	(-)	BNZ	300 mg	(-)	60 days	1	(-)
	80	(-)	BNZ	150 mg	(-)			(-)
	80	(-)	BNZ	400 mg	(-)	15 days		(-)
Torrico et al. (2021)	30	11	BNZ	$300~{ m mg~day}^{-1}$	b.d.i/oral	8 wk	1	18/23 (78%)
	30	60	BNZ	$300~{ m mg~day}^{-1}$	b.d.i/oral	4 wk		24/27 (89%)
	30	10	BNZ	$300~{ m mg~day}^{-1}$	b.d.i/oral	2 wk		22/28 (79%)
	30	90	BNZ	$150~{ m mg}~{ m day}^{-1}$	b.d.i/oral	4 wk		24/29 (83%)
	30	06	BNZ + FOS	150 + 300 mg day <sup>-1</sup>	b.d.i + o.d for 3 days followed by 300 mg once wk	4 wk		20/24 (83%)
	30	05	BNZ + FOS	300 + 300 mg day <sup>-1</sup>	b.d.i + o.d for 3 days followed by 300 mg once wk	8 wk		21/25 (84%)
	30	01	PLA	(-)	(-)	(-)		1/30 (3%)
<sup>a</sup> For discontinuity, adverse events and BNZ honznidazolo: EOC foccasuronaz	d other reasons.	od once a davi hidi	i twice day: whe week	·· - data not ronortod: 2 in	comulate information			

was identified in patients treated with BNZ (Molina et al., 2014; Morillo et al., 2015, 2017; Vallejo et al., 2016; Torrico et al., 2018, 2021), posaconazole (Molina et al., 2014; Morillo et al., 2017) and E1224 (Torrico et al., 2018) alone, as well as BNZ combined with thioctic acid (Sosa-Estani et al., 2004), posaconazole (Morillo et al., 2017) or fosravuconazole (Torrico et al., 2021). Leucopenia, neutropenia and/or lymphopenia were identified in patients treated with allopurinol, BNZ alone or combined with fosravuconazole (Rassi et al., 2007; Morillo et al., 2015; Torrico et al., 2021) (Table 3). Treatment discontinuation ranged from 1.5 to 57% in patients receiving BNZ alone or combined with thioctic acid, posaconazole or fosravuconazole. Cutaneous, gastrointestinal and neurological reactions were the most common adverse effects identified in 5-73% of all patients investigated. The mean rate of serious adverse reactions was 22.78% in BNZ-treated patients, while this rate was 2.56% in patients receiving placebo (Table 4).

### Sources of methodological bias

Based on the D&B checklist, compliance with the evaluated methodological criteria ranged from 46 to 100% (average result = 85.57%). Studies prior to 2014 were unable to meet all criteria. However, 4 studies (26.66%) after 2014 met all the methodological criteria analysed (Molina et al., 2014; Morillo et al., 2017; Torrico et al., 2018, 2021). Considering the items evaluated in the D&B checklist, criteria 8 and 10 were the least observed by the authors, namely: were the statistical tests used to assess the main outcomes appropriate? and were the main outcome measures used accurate (valid and reliable)? These criteria were met by 73.3% of the studies reviewed. Items 3 (Are the outcomes to be measured clearly described in the introduction or in the methods section?), 4 (Are the interventions of interest clearly described?) and 23 (Were the study subjects randomized to the intervention groups?) were consistently attended in all studies. The individual and overall results of bias can be accessed in Fig. 2 and Table S3.

### Discussion

In this review, we identified that the evidence provided by randomized controlled clinical trials targeting ChD treatment is based on BNZ (De Andrade et al., 1996; Sosa Estani et al., 1998; Andrade et al., 2004; Marin-Neto et al., 2008; Chippaux et al., 2010; Morillo et al., 2015; Vallejo et al., 2016; Molina-Morant et al., 2020), NFx (Coura et al., 1997), allopurinol (Rassi et al., 2007), posaconazole (Molina et al., 2014; Morillo et al., 2017), ravuconazole (Torrico et al., 2018) and fosravuconazole (Torrico et al., 2021); administered in monotherapy, as well as BNZ combined with thioctic acid (Sosa-Estani et al., 2004), posaconazole (Morillo et al., 2017) and fosravuconazole (Torrico et al., 2021). In line with the perspective of efficacy for the treatment of neglected tropical diseases, most drugs used in monotherapy and in combination showed remarkable relevance in attenuating T. cruzi infection. Accordingly, therapeutic effects were primarily associated with better parasitological control, as evidenced by negative seroconversion rates obtained from different treatment protocols. Although studies do not report a definitive parasitological cure, a low frequency of functional cardiac deterioration has been identified. However, the therapeutic regimens administered were associated with important side effects, indicating variable systemic toxicity.

Interestingly, the RCT identified were mainly concentrated in 3 Latin American countries (i.e., Brazil, Argentina and Bolivia) (De Andrade *et al.*, 1996; Coura *et al.*, 1997; Sosa Estani *et al.*, 1998; Andrade *et al.*, 2004; Sosa-Estani *et al.*, 2004; Rassi *et al.*, 2007; Chippaux *et al.*, 2010; Molina *et al.*, 2014; Torrico *et al.*, 2018,

Study	Grou treatn	ips nent	Pre-treatmer condit	nt cardiac ion	Post-tr	reatment cardiac condition	Labora	tory findings
De Andrade <i>et al.</i> BNZ (1996) 7.5 mg/kg		BNZRight bundle I7.5 mg/kg/60dblock $(n = 2)$ Left anterior I $(n = 1)$ Atrioventricula $(n = 2)$ Ectopic rhythr		oranch emiblock r block n ( <i>n</i> = 1)	nchRight bundle branch block $BNZ: n = 1 (1.7\%)$ iblockPLA: $n = 4 (6.9\%)$ lock $n = 1$		The frequency of anaemia (haemoglobin $\leq 110 \text{ g L}^{-1}$ ) was similar in the 2 groups no patient developed leucopenia (white-cell count <0.3 × 10 <sup>9</sup> /L) or neutropenia (neutrophil count <0.75 × 10 <sup>9</sup> /L)	
	PLA		Right bundle b block (n = 7)	oranch				
Coura et al. (1997) 5 mg/kg/30 NTX 5 mg/kg/30		/30d	(-)		ECG of patients did not suffer changes after treatment			(-)
		/30d	(-)		_			(-)
	PLA		(-)					(-)
Sosa Estani <i>et al.</i> (1998)	BNZ 5 mg/kg,	/60d	ECG: Left anter hemiblock or r	rior ight bundle	Changes 48 month	in ECG after is follow-up:		(-)
	PLA		branch block BNZ: <i>n</i> = ? (5%) PLA: <i>n</i> = ? (4.8%)	) %)	ventricular ectopic beats. BNZ: <i>n</i> = 1 (2.5%) PLA: <i>n</i> = 1 (2.4%)		(-)	
Author and year	of publication	Group	s treatment	Pre-treat cardiac co	ment ndition	Post-treatment cardiac condition		Laboratory findings
Andrade <i>et al.</i> (2004)		BNZ 7.5 ו	ng/kg/b.d.i	(-)		3 years follow-up ECG: complete right branch block (n = 1) 6 years follow-up No incident case of was found	bundle ECG abnormality	(-)
		PLA		(-)		3 years follow-up ECG: complete right block (n = 4) 6 years follow-up No incident case of was found	bundle branch ECG abnormality	(-)
Sosa-estani <i>et al.</i> (2004)		TA 50 mg + BNZ 5 mg	g day <sup>-1</sup> 1–37d	(–)		(–)		↑ALT ( <i>n</i> = 1)
		TA 100 mg/1–37d + BNZ 5 mg kg <sup>-1</sup> /5–37d		(–)	(-) (-)			↑ALT ( <i>n</i> = 1)
	-	PLA 1–37 BNZ 5 m	′d + g kg <sup>-1</sup> /5–37d	(-)		(-)		(–)
-		PLA 1–4c TA 50 mg BNZ 5 mg	PLA 1-4d + (- TA 50 mg/1-37d + BNZ 5 mg kg <sup>-1</sup> /5-37d			(–)		(-)
		Untreated (-)			(-)		(-)	
Rassi et al. (2007)	)	ALLP 300 mg, 3 × d/60d		(-)		(-)		Leucopenia (n = 2)
		PLA		(–)		(-)		(-)
Study	Groups treatmer	nt	Pre-treatment	t cardiac conditi	ion	Post-treatment of	cardiac condition	Laboratory findings
Molina <i>et al.</i> (2014)	BNZ 150 mg/b.d.	i	(-)		No significant effect		ts on QT interval	↑ ALT ( <i>n</i> = 5) ↑ AST ( <i>n</i> = 3) ↑ ALP ( <i>n</i> = 13)
	POS 400 mg/b.d.	i		(-)		_		↑ ALT ( <i>n</i> = 6) ↑ AST ( <i>n</i> = 4) ↑ ALP ( <i>n</i> = 11)
	POS 100 mg/b.d.	i		(-)				↑ ALT ( <i>n</i> = 10) ↑ AST ( <i>n</i> = 8) ↑ ALP ( <i>n</i> = 19)
Morillo <i>et al.</i> (2015)	BNZ 5 mg/kg/60c	d Atria Vent Resu	l fibrillation (n = ricular arrhythm iscitated cardiac	107) ia ( <i>n</i> = 221) arrest ( <i>n</i> = 19)		Deaths ( <i>n</i> = 246) Resuscitated cardia Ventricular tachyca	c arrest ( <i>n</i> = 10) rdia ( <i>n</i> = 33)	↑ ALT ( $n = 75$ ) Neutrophil < 1900/ mm <sup>3</sup> ( $n = 2$ )
-	-		-	-		-	-	

Table 3. Adverse events, cardiac function, immune response and laboratory findings in ChD patients

(Continued)

# Table 3. (Continued.)

Study	Groups treatment	Pre-treatn	nent cardiac co	ondition	Post-tr	reatment cardiac condition	Laboratory findings	
		Heart failure (n = Pacemaker (n = 2 Implantable card Stroke or transie Systemic or pulm	142) 05) ioverter-defibri nt ischaemic at nonary embolis	llator (n = 39) ttack (n = 61) m (n = 7)	Heart failure ( $n = 109$ ) Pacemaker or implantable cardioverter-defibrillator ( $n = 109$ ) Stroke or transient ischaemic attack, systemic embolism, or pulmonary embolism ( $n = 54$ ) Cardiac transplantation ( $n = 3$ )			
	PLA	Atrial fibrillation Ventricular arrhyt Resuscitated care Heart failure ( $n =$ Pacemaker ( $n = 1$ Implantable card Stroke or transie Systemic or pulm	(n = 90) thmia (n = 189) diac arrest (n = 128) 98) ioverter-defibr nt ischaemic at nonary embolis	16) illator (n = 31) ttack (n = 62) m (n = 11)	Deaths (n Resuscita Ventricula Heart fail Pacemake cardiovert Stroke or systemic o embolism Cardiac tr	= 257) ted cardiac arrest $(n = 17)$ ar tachycardia $(n = 41)$ ure $(n = 122)$ er or implantable ter-defibrillator $(n = 125)$ transient ischaemic attack, embolism, or pulmonary (n = 61) ransplantation $(n = 9)$	↑ ALT ( <i>n</i> = 28)	
Author and year of publication	Groups treatmen	Pre-treatr t cor	ment cardiac ndition	Post-treatme condit	ent cardiac tion	Laborator	y findings	
Vallejo <i>et al.</i> (2016)	BNZ 5 mg/kg/60 d	l Normal echocardi	ography	None develoj cardiomyopa	ped thy	↑ Naïve CD4 T cell, KLGF on CD4 memory cells, T TNF-α-producing CD4 T and TNFα/IL4 ratio ↑ Liver test values (n = 0	R 1 receptor expression h17, Th1; INF-γ and cells, Th1/Th2, INFγ/IL4 1)	
	Untreated	(-)		(-)		(-)		
Morillo <i>et al.</i>	POS 400 mg/b.i.d	Normal Q	QT interval	No significan	t effects al	↑ ALT ( <i>n</i> = 1)		
(2017)	PLA 10 mg/b.i.d.				at	↑ ALT ( <i>n</i> = 2)		
	BNZ 200 mg/b.i.d. POS 400 mg/b.i.d.	+				↑ ALT ( <i>n</i> = 4)		
	BNZ 200 mg+ PLA b.i.d					↑ ALT ( <i>n</i> = 2)		
Torrico <i>et al.</i>	E1224 4000 mg/8v	V				$\uparrow$ ALT and/or AST ( <i>n</i> = 5)		
(2018)	E1224 2000 mg/8v	V				(-)		
	E1224 2400 mg/4w 4 w placebo	v + Normal Q	T interval	No significan increases in (	t QT interval	(-)		
	BNZ 5 mg/kg/60d					↑ ALT ( <i>n</i> = 1)		
	PLA 8w					(-)		
Study	Groups tr	eatment	Pre-treatmer	nt cardiac conditio	on	Post-treatment cardiac condition	Laboratory findings	
Torrico <i>et al.</i> (2021) BNZ 300 mg/d/8w		v	Normal electrocardiogram			There were no meaningful safety signals from standard haematology, biochemistry and ECG	Neutropenia $(n = 8)$ Leucopenia $(n = 5)$ Lymphopenia $(n = 3)$ $\uparrow$ ALT $(n = 4)$ $\uparrow$ AST $(n = 1)$	
	BNZ 300 mg/d/4w	v				assessments.	Neutropenia $(n = 2)$ Leucopenia $(n = 2)$ lymphopenia $(n = 3)$ $\uparrow$ ALT $(n = 1)$ $\uparrow$ AST $(n = 1)$	
	BNZ 300 mg/d/2v	BNZ 300 mg/d/2w					Lymphopenia $(n = 1)$ $\uparrow$ ALT $(n = 2)$	
	BNZ 150 mg/d/4v	V					Neutropenia $(n = 2)$ Lymphopenia $(n = 1)$ $\uparrow$ ALT $(n = 4)$ $\uparrow$ AST $(n = 1)$	
	BNZ 150 mg mg/o	d/4w + FOS					Neutropenia $(n = 4)$ Leucopenia $(n = 1)$ Lymphopenia $(n = 1)$ $\uparrow$ ALT $(n = 4)$ $\uparrow$ AST $(n = 5)$	
							(Continued)	

### Table 3. (Continued.)

Study	Groups treatment	Pre-treatment cardiac condition	Post-treatment cardiac condition	Laboratory findings
	BNZ 300 mg/w/8 w + FOS			Neutropenia (n = 2) Leucopenia (n = 2) ↑ ALT (n = 3) ↑AST (n = 7)
	PLA	-		Normal: ALT and AST

BNZ, benznidazole; PLA, placebo; POS, posaconazole; TA, thioctic acid; ALLP, allopurinol; -, data not reported; ?, incomplete information; ALT, alanine aminotransferase; ECG, electrocardiogram; d, day; NTX, nifurtimox; AST, aspartate aminotransferase; ALP, alkaline phosphatase; E1224, ravuconazole prodrug; w, week; FOS, fosravuconazole; b.d.i, twice day.

Table 4. Adherence, discontinuation and adve	rse events associated with the treatm	nent administered in ChD patients.
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Study	Adherence	Discontinuation	Adverse events	Serious adverse events	Most common adverse events
De Andrade <i>et al.</i> (1996)	Completed the treatment and follow-up n = 112 (87%) BNZ: $n = 58$ PLA: $n = 54$	BNZ: <i>n</i> = 1 (1.5%)	Nausea, anorexia, headache, stomach ache, arthralgia (<5% patients). Cutaneous rash and pruritus BZN ( <i>n</i> = 8), placebo ( <i>n</i> = 2)	BNZ: Cutaneous rash and pruritus ( <i>n</i> = 1)	(-)
Coura <i>et al.</i> (1997)	Completed the treatment n = 64 (83.1%)	BNZ: <i>n</i> = 3 (11.5%) NF: <i>n</i> = 8 (29.6%)	(-)	BNZ: <i>n</i> = 3 (11.5%) NF: <i>n</i> = 8 (29.6%)	NF: gastrointestinal disorders, paraesthesia and neurological BNZ: gastrointestinal, rash, neurological
Sosa Estani <i>et al.</i> (1998)	Completed the treatment <i>n</i> = 101 (95.2%)	(-)	BNZ: Intestinal colic, cutaneous rash, headache, anorexia, vomiting, nausea, diarrhoea, dizziness, paraesthesia, hands shivering (n = <20%)	BNZ: <i>n</i> = 6 (10%)	Intestinal colic and rush were more frequent in BNZ than PLA
Sosa-Estani et al. (2004)	Completed the treatment n = 175 (70.3%)	BNZ: <i>n</i> = 74 (17.7%)	Patients affected by adverse events was similar between the 4 groups: 54.8 to 58%	Cutaneous rash, itching and fever (n = ?)	Rash 28%, pruritus 13.6%, headache 8%, epigastric pain 6.2%, fever 6.2%, asthenia 4.3%, nausea 4%, myalgia 4.3%, vomiting 3.2%, fatigue 4.3%, others 21.5%
Rassi <i>et al.</i> (2007)	Completed the treatment ALLP: <i>n</i> = 17 PLA: <i>n</i> = 10	ALLP: n=6 (26%)	ALLP: 11/23, fever ( $n$ = 5), cutaneous allergy ( $n$ = 5), pyrosis ( $n$ = 3), pruritus ( $n$ = 3), leucopenia ( $n$ = 2), lymph node enlargement ( $n$ = 1), vomiting (1), stomachache ( $n$ = 1), red eye ( $n$ = 1) PLA: 1/12, vertigo ( $n$ = 1)	(-)	(-)
Molina <i>et al.</i> (2014)	Completed the treatment and follow-up BNZ: $n = 17$ High-dose POS: n = 25 Low-dose POS: n = 20	BNZ: <i>n</i> = 5 (19%).	BNZ: Cutaneous rash $(n = 16)$ , Gastrointestinal $(n = 7)$ , Dysgeusia $(n = 2)$ , High-dose POS: Cutaneous $(n = 5)$ , gastrointestinal $(n = 4)$ , mucosal dryness $(n = 3)$ Low-dose POS: Cutaneous reaction $(n = 4)$ , gastrointestinal $(n = 3)$ , mucosal dryness $(n = 2)$	BNZ: Dermatitis ( <i>n</i> = 5), anaphylaxis with angioedema ( <i>n</i> = 1)	Headache, asthenia, sleepiness, arthralgia, dizziness

(Continued)

# Table 4. (Continued.)

Study	Adherence	Discontinuation	Adverse events	Serious adverse events	Most common adverse events
Morillo <i>et al.</i> (2015)	Completed the treatment and follow-up BZN: 1429/ 1431 (99.9%) PLA: 1422/ 1423 (99.9%)	BZN: n = 288 (20.1%) PLA: <i>n</i> = 66 (4.6%)	BNZ: Cutaneous rash ( $n = 9.59\%$ ), gastrointestinal ( $n =$ 7.84%), neurological ( $n = 3.64\%$ ) PLA: Cutaneous rash ( $n = 1.27\%$ ), gastrointestinal ( $n =$ 2.88%), Neurological ( $n = 1.34\%$ )	BNZ: Cutaneous rash (1.16%), gastrointestinal (1.82%) neurological (0.98%) PLA: Cutaneous rash ( $n = 0.14\%$ ), gastrointestinal ( $n = 0.63\%$ ), neurological ( $n = 0.42\%$ )	(-)
Vallejo <i>et al.</i> (2016)	Completed the treatment and follow-up BNZ: <i>n</i> = 3 (42%) PLA: <i>n</i> = 6 (85%)	BZN: <i>n</i> = 4 (57%) PLA: <i>n</i> = 1 (15%)	BNZ: Rash $(n = 3)$ , change in liver function $(n = 1)$ , rash and fever $(1)$	BNZ: Rash ( <i>n</i> = 2), fever ( <i>n</i> = 1)	(-)
Morillo <i>et al.</i> (2017)	Completed the treatment and follow-up POS: <i>n</i> = 32 (100%) PLA: <i>n</i> = 29 (97%) POS + BNZ: <i>n</i> = 19 (70%) BNZ + PLA: <i>n</i> = 20 (60%)	32% patients discontinue therapy due to side effects POS + BNZ: <i>n</i> = 9 (30%) BNZ + PLA: <i>n</i> = 10 (33%) PLA: <i>n</i> = 1 (3%)	POS: Cutaneous $(n = 2)$ , gastrointestinal $(n = 12)$ , neurological $(n = 4)$ PLA: Cutaneous $(n = 3)$ , gastrointestinal $(n = 5)$ , neurological $(n = 3)$ POS + BNZ: Cutaneous $(n = 12)$ , gastrointestinal $(n = 10)$ , neurological $(n = 10)$ , BNZ + PLA: Cutaneous $(n = 18)$ , gastrointestinal $(n = 8)$ , neurological $(n = 10)$	POS + BNZ: Hepatitis ( $n = 1$ ), rash ( $n = 1$ ) BNZ + PLA: Peripheral neuropathy ( $n = 1$ ), Abortion ( $n = 1$ ), cutaneous rash ( $n = 1$ ) PLA: Head injury ( $n = 1$ )	Headache (14%), nausea (10%), rash (10%)
Torrico <i>et al.</i> (2018)	Completed the treatment and follow-up n = 217 (?)	HD E1224 = 5 (11.1%) BNZ = 4 (8.9%)	81% patients affected by adverse events. BZN = 64.4%. LD E1224 = 31.3% SD E1224 = 52.2%. HD E1224 = 44.4%	Short-dose E1224: 1 (2%) High-dose E1224: 3 (7%) BNZ: 2 (4%)	(-)
Torrico et al. (2021)	Completed treatment and follow-up. <i>n</i> = 202 (96%)	BNZ 300 mg/day/ 8w: <i>n</i> = 6 (20%) BNZ 300 mg/day/ 4w: <i>n</i> = 1 (3%) BNZ 300 mg/day/ 2w: <i>n</i> = 0 (0%) BNZ 150 mg/day/ 4w: <i>n</i> = 1 (3%) BNZ 150 mg/day/ 4w + FOS: <i>n</i> = 3 (10%) BNZ 300 mg/day/ 8w + FOS: <i>n</i> = 4 (13%) PLA: <i>n</i> = 0 (0%)	70% patients affected by adverse events. BNZ 300 mg/day/8w: <i>n</i> = 23 (73%) BNZ 300 mg/day/4w: <i>n</i> = 19 (63%) BNZ 150 mg/day/2w: <i>n</i> = 19 (63%) BNZ 150 mg/day/4w: <i>n</i> = 19 (63%) BNZ 300 mg/day/8w + FOS: <i>n</i> = 22 (73%) BNZ 300 mg/day/8w + FOS: 19 (63%) PLA: <i>n</i> = 14 (47%)	BNZ 300 mg/day/8w + FOS: neutropenia $n = (1)$ , elevated AST $(n = 2)$ and GGT $(n = 1)$ , breast cancer $(n = 1)$ BNZ 300 mg/day/8w: neutropenia $(n = 2)$ , leucopenia $(n = 1)$ BNZ 150 mg/day/4w + FOS: Elevated AST (2) and GGT (1) BNZ 300 mg/day/8w: leucopenia and neutropenia $(n = 1)$ , leucopenia and neutropenia $(n = 1)$ , BNZ 300 mg/day/8w + FOS: neutropenia and leucopenia $(n = 1)$ , breast cancer $(n = 1)$ BNZ 300 mg/day/4w: pyrexia and rash $(n = 1)$ BNZ 150 mg/day/4w + FOS: acute cholecystitis and biliary polyp $(n = 1)$	BNZ 300 mg/day/8w: Headache, neutropenia and leucopenia were reported more frequently than other BZN regimens

BNZ, benznidazole; PLA, placebo; POS, posaconazole; ALLP, allopurinol; AT, thioctic acid; NF, nifurtimox; E1224, ravuconazole prodrug; LD, low-dose; SD, short-dose; HD, high-dose; -, data not reported; FOS, fosravuconazole; AST, aspartate aminotransferase; GGT, gamma-glutamyl transferase; ?, incomplete information.



**Fig. 2.** Percentage of items met in the methodological bias analysis (reporting quality) for all RCTs included in the systematic review. Bias analysis was based on the Downs & Black checklist for randomized and non-randomized studies. The dotted line indicates the average percentage of methodological criteria met (85.51%). The complete bias analysis stratified by domains and items assessed can be found in Table S3.

2021). In these endemic countries, environmental and socioeconomic factors create favourable conditions for T. cruzi infection (Mejía-Jaramillo et al., 2014; Santos et al., 2020). However, poor early diagnosis favours the transmissibility of this parasite in non-endemic areas, which is mainly linked to migratory movements of infected people (Guarner, 2019; Echeverría et al., 2020a). In the studies reviewed, ChD was rigorously confirmed by one or more diagnostic tools, including xenodiagnosis, indirect haemagglutination, indirect immunofluorescence, ELISA and PCR. Although xenodiagnosis has traditionally been proposed to confirm ChD, this method has shown a marked decline in the last 3 decades (Zingales, 2018). On the other hand, indirect haemagglutination, indirect immunofluorescence and ELISA are the most used tests for ChD diagnosis (Andrade et al., 2011; Nogueira et al., 2018; Zingales, 2018). These methods exhibit high analytical sensitivity (96% - 99%) (Castro et al., 2002; Nogueira et al., 2018), a characteristic consistent with most randomized studies published up to 2014. In the last 2 decades, PCR-based methods were frequently incorporated into clinical studies, increasing the sensitivity and specificity of parasitological diagnosis (Nogueira et al., 2018; Caldas et al., 2019; Molina-Morant et al., 2020). This method was predominant from 2014 onwards, being widely applied for ChD diagnosis, as well as for evaluating chemotherapy effectiveness. PCR is a remarkable tool for early detection of therapeutic failure (Caldas et al., 2019), which has been clearly demonstrated in patients with chronic ChD treated with BNZ during recent (Silveira et al., 2000; Solari et al., 2001) and late (Morillo et al., 2017) infections. Although PCR is the gold standard for *T. cruzi* detection, its applicability is quite restricted to clinical studies and specialized laboratories due to the higher cost, need for specialized devices and professionals with greater technical qualification (Caldas et al., 2019). Thus, expanding access to this technology can improve the monitoring of populations exposed to greater risk of T. cruzi infection and the health care of newly infected patients.

Although BNZ is the first-line treatment for acute (Caldas *et al.*, 2019), recent (Silveira *et al.*, 2000; Solari *et al.*, 2001) and late (Morillo *et al.*, 2017) chronic infections, its effectiveness in achieving parasitological cure is still controversial, especially in

chronic cases. Apparently, divergent parasitological results are influenced by parasite strain (e.g., tropism, virulence and pathogenicity), patient characteristics (e.g., age, immunological status and comorbidities) and infection phase (De Andrade *et al.*, 1992; Filardi and Brener, 1987). In addition, the toxicity profile and numerous side effects of different antiparasitic drugs have a marked impact on treatment adherence (Morillo, *et al.*, 2015, 2017). Accordingly, longer drug regimens that require high doses represent a greater clinical challenge, as dose-dependent and time-dependent toxicity favours treatment discontinuation and therapeutic failure (e.g., poor parasite control) (Vallejo *et al.*, 2016; Caldas *et al.*, 2019; Molina-Morant *et al.*, 2020).

Currently, there is an objective recommendation to administer BNZ between 5 and 8 mg kg day<sup>-1</sup>, twice a day for 60 days (Andrade et al., 2011). However, marked variations in BNZ dose and treatment period are recurrent in clinical studies (De Andrade et al., 1996, Sossa-Estani et al., 1998; Andrade et al., 2004; Marin-Neto et al., 2008; Chippaux et al., 2010, Morillo et al., 2015; Vallejo et al., 2016; Mollina-Morant et al., 2020). Accordingly, we identified that therapeutic schemes based on  $2-7.5 \text{ mg kg day}^{-1}$  for 60 or 30 days were respectively reported for newborns (Chippaux et al., 2010); while conventional dosimetry (5 mg kg day $^{-1}$  for 60 days) was investigated in children aged 13 years and older (Andrade et al., 2004), and patients aged 26-57 years (Vallejo et al., 2016). The same conventional dose was assigned for patients aged 18-75 years; however, different administration periods (40, 60 or 80 days) were evaluated (Marin-Neto et al., 2008). In addition, significant adaptations were recently incorporated into randomized studies with BNZ, which used 150 and 300 mg kg day<sup>-1</sup> by 60 days or 400 mg kg day<sup>-1</sup> by 15 days in patients aged 50-80 years (Molina-Morant et al., 2020). Objectively, these variations express the urgent need to define more efficient protocols for the reference drug, whose dose ranges and administration periods were mainly established in clinical practice and not in unequivocal evidence of therapeutic efficacy.

Regardless of the therapeutic regimen adopted, parasitological clearance, negative seroconversion, as well as low systemic toxicity are desirable clinical outcomes of drug treatment, including BNZ (Caldas *et al.*, 2019). Thus, we found that 5 mg BNZ kg<sup>-1</sup> day<sup>-1</sup> per60 days administered to patients with chronic ChC aged 26-57 years was associated with 100% negative seroconversion compared to 45% in untreated patients after 1 year follow-up (Vallejo et al., 2016). Even using similar therapeutic regimens  $(2-7.5 \text{ mg kg day}^{-1} \text{ for } 30-80 \text{ days})$ , BNZ-induced divergent negative seroconversion rates, ranging from 11.2 to 91.3% (De Andrade et al., 1996; Sosa Estani et al., 1998; Andrade et al., 2004; Chippaux et al., 2010). Thus, the best parasitological results were obtained by Vallejo et al. (2016), followed by Morillo et al. (2017), Molina et al. (2014) and Chippaux et al. (2010), who reported respectively 100, 96 and 94.1% cure rates for young adults and 91.3% cure for newborns, both after 1 year follow-up. Despite this response, clinical trials support BNZ efficacy for congenital ChD, corroborating a consistent trend of negative serology maintenance over time, an outcome that can be more easily reversed in adults (Blanco et al., 2000).

Although NFx is the second choice for ChD treatment, RCT with this drug are scarce. Thus, NFx was evaluated in only one trial identified in this review (Coura *et al.*, 1997). Unfortunately, this study reinforced the controversial effectiveness of this drug for chronic ChD. Accordingly, Coura *et al.* (1997) did not identify negative seroconversion in patients receiving 5 mg kg day<sup>-1</sup> NFx for 30 days or BNZ after 1 year follow-up. Thus, the lack of evidence of parasitological cure attributed to NFx in addition to its recognized toxicity profile (e.g., polyneuritis and bone marrow depression) makes the clinical use of this drug inadvisable when BNZ is available and well tolerated (Urbina and Docampo,

2003; Muñoz et al., 2011; Nogueira et al., 2018). Despite clinical evidence generated in recent decades supporting BNZ-based treatment for indeterminate chronic ChD (Molina et al., 2014; Morillo et al., 2017; Torrico et al., 2018), more efficient and safer drugs are still needed. Thus, several studies reviewed were based on repositioning strategies involving drugs with trypanocidal potential, such as allopurinol (Rassi et al., 2007), posaconazole (Molina et al., 2014; Morillo et al., 2014; Morillo et al., 2017), ravuconazole (Torrico et al., 2018) and fosravuconazole (Torrico et al., 2021).

Studies with allopurinol have confirmed its potent trypanostatic effect in vitro on 5 T. cruzi strains (Marr et al., 1978; Avila and Avila, 1981). Interestingly, Berens et al. (1982) confirmed that allopurinol is metabolized by bloodstream trypomastigotes and intracellular amastigotes, which can be eradicated in vitro by this drug. Contrary to expectations, allopurinol administration (300 mg kg day<sup>-1</sup> for 60 days) was not associated with negative seroconversion in patients with indeterminate chronic ChD (Rassi et al., 2007). In contrast, posaconazole and ravuconazole (e.g., antifungal drugs used in humans) have proven their trypanocidal activity in vitro and in vivo (Urbina et al., 1998; Molina et al., 2014). Accordingly, posaconazole induced marked parasitological cure compared to BNZ-based monotherapy in acute experimental ChD (Olivieri et al., 2010; Calvet et al., 2020). Thus, all T. cruzi-infected mice (100%) receiving posaconazole and 50% receiving BNZ had negative blood cultures for this parasite (Calvet et al., 2020). Surprisingly, this effect was even better in chronic T. cruzi infection, with parasitological cure rates reaching 60 and 0% when these animals were respectively treated with posaconazole and BNZ (Urbina et al., 1998; Molina-Morant et al., 2020). Conversely, BNZ (150 mg<sup>-1</sup>kg per 60 days) showed better results compared to posaconazole (100 or 400 mg<sup>-1</sup>kg per 60 days), returning a higher negative seroconversion rate (94.1%) for BNZ compared to posaconazole (10-20%) (Molina et al., 2014). Torrico et al. (2018) identified a similar response in patients with chronic ChD receiving ravuconazole. Accordingly, this drug (4000 or 2000 mg for 65 days) determined lower negative seroconversion rates (28.9% or 8.3%) compared to BNZ (82.2%,  $5 \text{ mg}^{-1}$ kg per 60 days).

Currently, available evidence indicates that negative seroconversion rates obtained with these azoles are lower than BNZ-based monotherapy (Moraes et al., 2014; Chatelain, 2015). However, combining these drugs has been suggested as a means to improve treatment efficacy, with the prospect of simultaneously interfering with multiple molecular pathways associated with T. cruzi parasitism (Bustamante et al., 2014; Morillo et al., 2017). Accordingly, Torrico et al. (2021) identified that therapeutic regimens combining BNZ (e.g., 150 or 300  $\rm mg\,kg^{-1}$  for 2, 4 or 8 weeks, respectively) and fosravuconazole  $(300 \text{ mg kg}^{-1})$ for 4 or 8 weeks) achieved better effects than BNZ-based monotherapy. Interestingly, this study supported the proposition that BNZ dose could be reduced without losing its effectiveness (Bustamante et al., 2014; Álvarez et al., 2020). In this sense, Torrico et al. (2021) demonstrated similar negative seroconversion rates and cardiac function in patients exposed to conventional and low BNZ doses when combined with fosravuconazole in a 4-week protocol. Conversely, BNZ + posaconazole (200 + 400 mg kg<sup>-1</sup> twice daily for 60 days) did not change seroconversion rates or improved cardiac function compared to BNZ-based monotherapy after 1-year follow-up (Morillo et al., 2017). A similar effect was reported combining BNZ and thioctic acid (50 and 100 + 5 mg kg<sup>-1</sup> twice daily for 1–37 days), which showed no superior benefit over reference chemotherapy after 53 days follow-up (Sosa-Estani et al., 2004). Thus, the efficacy of pharmacological combinations is not unequivocal, even considering drugs with potent antiparasitic effects in preclinical models (Diniz et al., 2013; Echeverría et al., 2020b).

Despite successful examples, evidence of therapeutic failure indicates that ChD treatment is still challenging, especially considering that the long-term prognosis may not be favourable even with negative seroconversion in up to 1-year follow-up (Caldas et al., 2019). Thus, parasite clearance may not improve or prevent cardiac deterioration in patients with chronic ChD (Marin-Neto et al., 2008; Bern et al., 2007). Although this finding is discouraging, it also indicates that negative seroconversion and parasitological cure may not be ideal indicators for estimating the full spectrum of benefits related to antiparasitic chemotherapy (Gonçalves and Novaes, 2018). From this perspective, De Andrade et al. (1996) identified a positive association between negative seroconversion rates and cardiac function in ChD patients receiving BNZ, whose electrical abnormalities (e.g., right bundle branch block) were attenuated even in the absence of parasitological cure. Although Sosa Estani et al. (1998) found low negative seroconversion rates (11.2%) in BNZ-treated children, no electrical conduction disturbances were identified after 48-months follow-up. Likewise, radiological or electrocardiographic changes were not detected in BNZ-treated adults  $(5 \text{ mg kg day}^{-1} \text{ twice a day})$ , who maintained positive serology 12 months after treatment (Coura et al., 1997). These findings indicate the need to reframe the perception of therapeutic failure and success, since preventing progression to the symptomatic chronic phase and CCC evolution may be equally or more relevant outcomes than parasitological cure (Gonçalves and Novaes, 2018).

Considering the evidence based on randomized clinical trials, BNZ clearly showed greater overall efficacy compared to NFx, allopurinol, posaconazole, ravuconazole and fosravuconazole. However, the toxicity profile of BNZ is still a serious limitation to be overcome, especially considering the negative impact on treatment adherence (Molina-Morant et al., 2020; Martín-Escolano et al., 2020). In line with clinical experience, laboratory findings such as neutropenia, leucopenia, lymphopenia (De Andrade et al., 1996; Morillo et al., 2015; Torrico et al., 2018, 2021), increased transaminases circulating levels, and upregulation in pro-inflammatory effectors (Vallejo et al., 2016) were reported in the studies reviewed. Accordingly, BNZ/placebo ratios for adverse events ranging from 1.64 (Morillo et al., 2015) to 3.85 (Torrico et al., 2021) were identified in these studies. These findings were consistent with gastrointestinal, neurological and cutaneous adverse events, which can be reported in up to 90% of patients receiving trypanocidal chemotherapy (Francisco et al., 2020). In general, treatment discontinuation is most often attributed to serious adverse effects associated with therapeutic regimens with higher doses administered for longer periods (Norman and López-Vélez, 2019; Morillo et al., 2017). However, the incidence of adverse effects does not always follow this quantitative logic, exhibiting marked variability even in similar therapeutic regimens (Torrico et al., 2018, 2021). Accordingly, adverse effects ranging from less than 20% (Sosa Estani et al., 1998) to 64.4% (Torrico et al., 2018) in patients with chronic ChD receiving the same treatment (e.g., 5 mg kg day<sup>-1</sup> BNZ for  $\cong$  60 days), reinforce the proposition that patient characteristics influence the organic tolerability to antiparasitic drugs (Salvador et al., 2015; Echeverría et al., 2020b). Apparently, sensitivity to BNZ may have an important genetic component, as a higher treatment discontinuation rate related to cutaneous adverse reactions was detected in patients carrying the HLA-B\*3505 allele (Salvador et al., 2015).

From a critical interpretation of the evidence, we identified that  $85.51\% \pm 15.49\%$  of all the basic methodological criteria investigated in the bias analysis instrument were met, with adherence scores ranging from 46% (Molina-Morant *et al.*, 2020) to 100% (Molina *et al.*, 2014; Morillo *et al.*, 2017; Torrico *et al.*, 2018, 2021). Interestingly, these findings indicate that most of

the studies reviewed presented high methodological rigour. Contrary to expectations, the quality index did not show a clearly time-dependent behaviour (influence of the publication year). Thus, the variability detected may be linked to the systematic replication of confounding factors (sources of bias) despite the advances applied to the design, operationalization and monitoring of randomized clinical trials, as well as greater availability of sensitive and specific analytical tools applicable to parasitology research. In cases of partial methodological adherence, the least met criteria were related to incomplete reporting of random variability estimates, specific delimitation of statistical probability, definition of appropriate statistical tests and precise use (validity and reliability) of the main outcome measures. Admittedly, these methodological limitations undermine the reproducibility, internal and external validity of the studies reviewed, limiting evidence reliability (Downs and Black, 1998; Torrico et al., 2018, 2021). However, it is important to consider that these quality scores do not indicate flaws in the experimental protocols, as they exclusively point out limitations in the research reports. Thus, by mapping potential bias sources in all investigated studies, this review provides objective support to delimit further clinical trials with greater methodological rigour.

Taken together, the RCT provide robust evidence that BNZ is the most viable therapeutic option for the aetiological treatment of acute and chronic ChD. However, adjustments in therapeutic protocols based on this drug are still underway in search of optimized responses to increase adherence to antiparasitic chemotherapy and therapeutic success rates (Molina-Morant et al., 2020; Morillo et al., 2017; Torrico et al., 2018, 2021). Therefore, shorter regimens or lower BNZ doses appear to be viable options to ensure similar efficacy compared to the reference protocol, reducing the incidence of adverse effects potentially linked to dose- and time-dependent toxicity reactions. In addition, combining BNZ with other antiparasitic drugs such as posaconazole, fosravuconazole or ravuconazole may be relevant to attenuate the frequency of adverse effects, despite not having a significant impact on negative seroconversion and parasitological cure compared to BNZ-based monotherapy. In order to improve the pharmacological management of ChD patients, longer clinical follow-up is required to evaluate aetiological treatment efficacy, allowing objectively characterizing the relevance of negative seroconversion and parasitological cure as primary endpoints of therapeutic success or failure. Thus, an ambitious proposal is to design robust methodological protocols incorporating more sensitive and specific diagnostic methods (e.g., PCR), allowing the reassessment of patients included in these randomized trials to clarify the relationship between parasitological cure and ChD progression.

**Supplementary material.** The supplementary material for this article can be found at https://doi.org/10.1017/S0031182022001081

Author's contributions. Silas Santana Nogueira: Investigation, visualization, data gathering and writing – original draft. Eliziária Cardoso Santos: Investigation, visualization, data gathering and writing – original draft. Roberta Oliveira Silva: Investigation, visualization, data gathering and writing – original draft. Reggiani Vilela Gonçalves: Formal analysis, writing – review and editing. Graziela Domingues Almeida Lima: Formal analysis, writing – review and editing. Rômulo Dias Novaes: Conceptualization, data gathering, formal analysis, writing – original draft, review and editing, resources, supervision.

**Financial support.** This work was supported by the Brazilian agencies: Fundação do Amparo à Pesquisa do Estado de Minas Gerais (FAPEMIG, processes PPM-00077-18 and PPM-00687-17) and Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq, processes 310331/2020-0, 423594/2018-4, 408503/2018-1 and 311105/2020-3). The Coordenação de Aperfeiçoamento de Pessoal de Nível Superior – Brasil (CAPES – Finance Code 001), partially funded this study. Conflict of interest. None.

Ethical standards. Not applicable.

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