Cardiovascular findings by echocardiography in canine model of Chagas disease immunized with DNA Trypanosoma cruzi genes.

**CURRENT STATUS:** POSTED

Olivia Rodríguez-Morales  
Instituto Nacional de Cardiologia Ignacio Chavez  

Francisco-Javier Roldán  
Instituto Nacional de Cardiologia Ignacio Chavez  

Jesús Vargas-Barrón  
Instituto Nacional de Cardiologia Ignacio Chavez  

Enrique Parra-Benítez  
Instituto Nacional de Cardiologia Ignacio Chavez  

María de Lourdes Medina-García  
Instituto Nacional de Cardiologia Ignacio Chavez  

Emilia Vergara-Bello  
Instituto Nacional de Cardiologia Ignacio Chavez  

Minerva Arce-Fonseca  
Instituto Nacional de Cardiología Ignacio Chávez  

✉ mini_arce@yahoo.com.mx  
**Corresponding Author**  
**ORCiD:** https://orcid.org/0000-0002-2706-4675

**DOI:**  
10.21203/rs.2.18247/v1

**SUBJECT AREAS**  
Small Animal Medicine

**KEYWORDS**  
Echocardiography, Chagas disease, Trypanosoma cruzi, canine model, DNA vaccine
Abstract
Background: Chagas disease (ChD) is nowadays considered as an emerging disease in the USA and Europe. pBCSP and pBCSSP4 plasmids, containing Trypanosoma cruzi genes encoding a trans-sialidase protein and an amastigote-specific glycoprotein, respectively, were tested as vaccines in canine model. Echocardiography studies for determining the prophylactic effect of these genes in experimentally infected dogs were evaluated to compare with findings obtained by other techniques performed previously. Hemodynamic parameters after DNA-immunization were performed.

Results: Low fractional-shortening values of non-vaccinated dogs suggested an impairment in general cardiac function. Low Left-Ventricular-Ejection-Fraction values found in infected dogs suggested myocardial injury regardless of whether they were vaccinated or not. Low Left-Ventricular-Diastolic/Systolic-Diameters in vaccinated dogs suggested that progressive heart damage or heart dilation could be prevented by DNA vaccination. Systolic-Peak-Time was higher in non-vaccinated groups increasing vulnerability to malignant arrhythmias and sudden death. High Left-Ventricular-Volume in infected groups suggested a decrease in wall thickness that might lead to increased size of the heart cavity, except in the pBCSP plasmid-vaccinated dogs.

Conclusions: The use of echocardiography allowed a more complete follow-up the pathological process in the living patient than with other techniques like electrocardiography, anatomopathology and histopathology, being the method of choice for characterizing the clinical stages of ChD.

Introduction
Chagas disease (ChD) is caused by the protozoan parasite Trypanosoma cruzi, which infects humans and more than 100 species of domestic and sylvatic mammals and can be transmitted by over 150 species of hemiptera insects of the subfamily Triatominae (Reduviidae). ChD has become one of the biggest public-health problems in Latin America due to its incapacitating effects and mortality rates. According to data reported by the World Health Organization, Argentina, Brazil and Mexico are the three countries with the highest estimated number of infected people. Estimated numbers of chagasic cardiopathy cases are highest in Argentina, Brazil, Colombia, Bolivia and Mexico [1]. Nevertheless, patterns of emigration from Chagas-endemic areas have drastically altered the epidemiology of this
disease in the United States, Europe, and other non-endemic regions in recent decades, making it one of the neglected tropical diseases now found in non-endemic areas of the world [2, 3].

There are two successive stages in ChD: acute and chronic phases. In the acute phase, cardiac involvement may occur in up to 90% of cases. After six to eight weeks, most patients show recovery of the clinical manifestations [3]. In the chronic phase of ChD, during which parasites are hidden in target tissues, especially the cardiac and digestive system muscles, different clinical forms may be observed: (i) the asymptomatic form; (ii) the cardiac form, which occurs in around 30% of the patients with disorders of the heart’s electrical conduction system, arrhythmia, heart-muscle disorder, heart failure (HF) or secondary embolisms; (iii) the digestive form, with localized lesions and enlargement of the oesophagus and the colon; and (iv) a mixed form (cardiac plus digestive) that affects around 10% of patients. Patients ultimately die, usually from sudden death caused by arrhythmias or HF, which often occurs in early adulthood [1]. The main causes of death associated with chronic Chagas’ cardiomyopathy (CCC) are progressive congestive HF and sudden cardiac death [4].

The diagnosis of ChD can be made by serology a few weeks after the primary infection. Large-scale epidemiological studies have consistently demonstrated that about three-quarters of seropositive patients remain asymptomatic throughout their entire life [5]. It is therefore important to detect cardiac involvement as early as possible in order to estimate the risk and prognosis before the patient becomes symptomatic [6]. Cardiac damage is suspected by ≥ 1 of the following electrocardiographic findings: right bundle-branch block, left anterior fascicular block, atrioventricular blocks, multiform ventricular beats, and sinus bradycardia [7].

In chronic Chagas heart disease, the electrocardiogram (EKG) has been the method of choice for detecting myocardial damage, especially in remote endemic areas, because of its low cost, portability and simplicity. For non-invasive detection of early myocardial damage other methods have been used, including Holter monitoring, echocardiography, nuclear scans and stress tests [6]. Two-dimensional and Tissue Doppler Imaging echocardiography are useful, complementary methods for the follow-up of patients with chronic ChD myocarditis. Both techniques provide valuable information on cardiac structure and function that complements information provided by EKG allowing the
recognition of left ventricular (LV) systolic and diastolic dysfunction, right ventricle involvement, and regional contractility abnormalities, including typical apical aneurysms. The cardiac ultrasound has utility in the diagnosis, classification, and detection of early myocardial damage and the prognostic assessment of patients with ChD [2, 7-9].

Dogs play a major role in the domestic cycle of Trypanosoma cruzi, acting as reservoirs, but they can also develop acute and chronic disease, similar to human infection; therefore, canine Chagas’ disease has become a major veterinarian concern in the Americas [10-12]. In small animals’ veterinary medicine, echocardiography is common technique performed on dogs to non-invasively assess systolic myocardial function. This technique has been shown to be repeatable and reproducible in dogs [13]. In addition, the canine model has gained wide acceptance as another experimental model to study a wide variety of conditions associated with ChD; however, the development of vaccines as a prophylactic method has not been widely addressed [14-24].

Development of anti-T. cruzi vaccines could significantly contribute to the control of ChD.

Immunological protection against experimental infection with T. cruzi has been studied since the second decade of the last century testing many types of immunogens [25]. In our previous studies, TcSP (encoding trans-sialidase protein) and TcSSP4 (encoding amastigote-specific protein) genes were evaluated prophylactically as DNA vaccines in both murine and canine model of ChD [26-30].

The clinical, serological and post-mortem parameters obtained were consistent among these studies, showing a moderate effectiveness in protection against experimental ChD. The effect of these genes, used as vaccines, on the clinical parameters was evaluated through general physical examinations as consequence of immunization and through their effect on the cardiac protection determined by electrocardiography [27]. However, these studies do not include complementary pre-mortem examinations about both structural and functional cardiac abnormalities provided by medical imaging methods such as echocardiography, which could be useful in obtaining accurate data on the development of morphological changes in the heart of experimentally infected and immunized/infected chagasic dogs.

The aim of this study is to examine the usefulness of echocardiography in determining the
prophylactic effect of the DNA vaccines on the heart damage using T. cruzi genes cloned into an
expression vector, which was intramuscularly injected to mongrel and Beagle dogs. To do so, we use
this cardiac imaging method to measure hemodynamic parameters to determine whether
echocardiography provides accurate data on the development of morphological changes in the heart
of experimentally infected and immunized/infected chagasic dogs, and if these data are consistent
with those reported previously [27–29].

Results
Vaccination ameliorated the LVEF
To correlate the prophylactic effects of DNA vaccines with T. cruzi genes reported previously on the
protection of heart damage with the reduction of the cardiac function alterations by hemodynamic
parameters, echocardiography was performed. This non-invasive method provided the useful tool to
evaluate both anatomy structure and function of the heart. The mean ± standard deviations values of
some cardiovascular parameters recorded at each group (specified in Table 1) are showed in Table 2.
Table 2
Cardiovascular parameters in T. cruzi experimentally infected dogs immunized with DNA vaccine containing the genes encoding a trans-sialidase protein (pBCSP) or an amastigote-specific glycoprotein (pBCSSP4).

| Group                     | Fractional shortening (FS) (%) | Left ventricular ejection fraction (LVEF) (%) | Left ventricular (LV) diastolic diameter (mm) | Left ventricular (LV) systolic diameter (mm) | Heart rate (bpm) | Systolic peak time (ms) | Left Ventricular Volume (mL) |
|---------------------------|--------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|------------------|------------------------|-------------------------------|
| A (healthy control)       | 39.29 ± 3.76                   | 77.60 ± 4.22                                 | 28.70 ± 2.84                                 | 17.42 ± 1.98                                  | 101.80 ± 24.14   | 212.40 ± 20.11         | 18.56 ± 3.11                  |
| B (Non-imm/Non-inf)       | 31.64 ± 8.12*                  | 50.29 ± 6.59*                                 | 34.19 ± 7.15*                                 | 23.35 ± 5.83*                                 | 102.87 ± 21.58   | 261.93 ± 31.57*        | 33.97 ± 18.62*                |
| C (P-imm/Non-inf)         | 41.19 ± 4.13                   | 78.20 ± 5.17*                                 | 26.30 ± 2.30                                 | 15.44 ± 1.46                                  | 128.60 ± 31.38   | 189.00 ± 15.51         | 16.80 ± 1.30                  |
| D (pBCSP-imm/inf)         | 33.57 ± 6.31                   | 54.62 ± 9.19*                                 | 29.90 ± 2.43                                 | 20.68 ± 2.77                                  | 93.00 ± 11.17    | 206.25 ± 15.31         | 23.73 ± 6.01                  |
| E (pBCSSP4-imm/inf)       | 33.94 ± 9.17                   | 58.45 ± 12.03*                                | 30.06 ± 1.88                                 | 20.84 ± 2.36                                  | 112.63 ± 26.38   | 193.00 ± 31.92         | 25.49 ± 5.18                  |
| F (pBK-CMV-imm/inf)       | 31.55 ± 1.13*                  | 52.33 ± 5.37*                                 | 31.70 ± 34.59*                               | 21.67 ± 2.80*                                 | 137.67 ± 30.66   | 244.00 ± 19.16         | 32.83 ± 4.66                  |
| G (SS mock-imm/inf)       | 27.32 ± 8.29*                  | 47.67 ± 4.93*                                 | 30.2 ± 1.13*                                 | 21.80 ± 4.81*                                 | 159.67 ± 37.53*  | 240.33 ± 16.74*        | 29.13 ± 9.02                  |

All data are expressed as means and standard deviations. Statistical significances (P ≤ 0.05) are assigned as * when there was a significant difference with group A control healthy dogs and/or with reference values [35]. Bold values belong to the groups which were vaccinated and did not show significant difference with group A control healthy dogs. bpm = beats per minute, imm = immunized, inf or Inf = infected, P-imm = each plasmids-immunized, pBCSP = plasmid carrying TcSP gene, pBCSSP4 = plasmid carrying TcSSP4 gene, pBK-CMV = empty cloning vector, and SS = saline solution.

The infected dogs (B group) had fractional shortening (FS %) values significantly reduced, very similar to those obtained in mock-immunized (G group) and empty cloning vector-immunized (F group) dogs suggesting an impairment in general cardiac function.

Significant differences in the LVEF values among infected and non-infected dogs could be demonstrated regardless of whether they were vaccinated or not, although the lowest values belonged to the group non-immunized with T. cruzi DNA (G group). No significant differences could be demonstrated in the LVEF values among immunized and non-immunized infected dogs, although slightly higher values in D and E groups (pBCSP-immunized/infected and pBCSSP4-immunized/infected dogs, respectively) could be detected suggesting a better effectiveness of pumping into the systemic circulation.
Plasmid DNA immunization prevented the increase in LV diastolic and systolic diameter. Both LV diastolic and systolic diameter were also significantly high in B, F and G groups (infected, empty cloning vector-immunized/infected, and SS mock-immunized/infected, respectively), while the values observed in vaccinated dogs were slightly lower (P < 0.05) than those of dogs that did not receive either vaccination, who progressive heart damage or heart dilation was diagnosed. Vaccination protected against arrhythmias and sudden death. Neither infection nor the vaccination/infection altered the heart rate. However, systolic peak time was significantly higher in infected (B group) and mock-vaccinated/infected (G group) dogs; conversely, immunization with T. cruzi genes or with the empty plasmid maintained this parameter as healthy or reference values showing a less vulnerability to malignant arrhythmias and sudden death. TcSP gene avoided left ventricular dilation. LV volume was also elevated in all infected dogs (B, E, F and G groups) suggesting a decrease in wall thickness that might lead to increased size of the heart cavity; on the other hand, values in this parameter in animals vaccinated with the plasmid that carried the TcSP gene (D group) resemble those obtained in the healthy control (A group) and immunized/non-infected (C group) dogs. Discussion: In this study, echocardiographic features found in experimental infected and DNA-immunized/infected dogs in chronic phase of the ChD were obtained. These features provided accurate data about cardiac structure and function, which complemented information given by other methods previously reported by us (Table 3) [27–29]. As expected, most of typical abnormalities found in these chagasic dogs by EKG studies, histology and post-mortem examination could be seen in non-immunized/infected dogs, and they are in concordance with the evidence from this imaging technique.
Table 3

Other findings obtained previously in the same T. cruzi experimentally infected dogs immunized with DNA vaccine containing the genes encoding a trans-sialidase protein (pBCSP) or an amastigote-specific glycoprotein (pBCSSP4).

| Group (description) | Electrocardiographic abnormalities (percentage of animals with these alterations) [27, 29, 32] | Parasitemia [27, 32] | Post-mortem macroscopic alterations (percentage of animals with these findings) [29] | Cardio- and/or Splenomegaly [29] | Histopathology |
|---------------------|-----------------------------------------------------------------------------------------------|---------------------|-------------------------------------------------------------------------------------|----------------------------------|-----------------|
| A (healthy control) | None                                                                                         | Negative            | None                                                                                 | None                             | None            |
| B (Inf)             | Ischemia and conduction block LV hypertrophy (100%)                                           | Positive at 22–55 days post-infection | Whitish areas in heart of fibrous consistency and abundant pericardial fluid (69%) | Cardiomegaly                     | Inflammatory lesions from subepicardium, myocardium and subendocardium |
| C (P-imm/Non-inf)   | None                                                                                         | Negative            | None                                                                                 | None                             | None            |
| D (pBCSP-imm/Inf)   | AV Block (33%) and LV enlargement (11%)                                                       | Positive at 32–46 days post-infection | None                                                                                 | Cardiomegaly                     | Inflammatory lesions from subepicardium, myocardium and subendocardium |
| E (pBCSSP4-imm/Inf) | MIMI (11%) and Second-degree AV block (11%)                                                   | Positive at 32–46 days post-infection | Heart with adhesions in trachea and pericardium, splenic thickened walls and whitish areas in spleen and heart (11%) | None                             | Inflammatory lesions from subepicardium |
| F (pBK-CMV-imm/Inf) | LV enlargement (75%) and Right BBB (25%)                                                      | Positive at 31–46 days post-infection | Abundant pericardial fluid and whitish areas in spleen (75%). Ascites, megaesophagus, LV hypertrophy, thinning of the RV wall and severe tricuspid endocarditis (25%) | Cardiomegaly Splenomegaly        | Inflammatory lesions from subepicardium, myocardium and subendocardium |
| G (SS mock-imm/Inf) | VPC (25%), pericardial effusion (50%), and myocardial infarction and/or pericarditis, and RV enlargement (50%) | Positive at 21–55 days post-infection | Whitish areas in heart of fibrous consistency and abundant pericardial fluid (75%) | Cardiomegaly Splenomegaly        | Inflammatory lesions from subepicardium, myocardium and subendocardium |

imm = immunized, inf or Inf = infected, P-imm = each plasmids-immunized, pBCSP = plasmid carrying TcSP gene, pBCSSP4 = plasmid carrying TcSSP4 gene, pBK-CMV = empty cloning vector, and SS = saline solution. FS = fractional shortening, LVEF = left ventricular ejection fraction, LV = left ventricular, MIMI = Microscopic intramural myocardial infarctions, N/D = not determined.

A low FS percentage was found in non-immunized dogs. This parameter is an index of general cardiac function that can provide a comprehensive evaluation of right ventricular systolic function with other recommended echocardiographic parameters. Right ventricular function is an important predictor of...
mortality and quality of life in patients with LV failure, myocardial infarction, congenital heart disease, and pulmonary hypertension. FS is affected by many factors, but the three most important are: preload, afterload and finally contractibility itself [35, 36]; according to this, we suggest that non-vaccinated dogs developed an ability reduction of the myocardial fibers to distend, which together with the diastolic dimensions suggests little contractility. It is documented that a high afterload hinders muscle contraction could show a decreased FS as an important feature by echocardiography [36], such as we can observe in those non-vaccinated/infected dogs suggesting an impairing in general cardiac function.

The LVEF percentage had a decrease in all experimental infected groups, suggesting significant myocardial injury caused by T. cruzi infection. This finding is consistent with a study performed with a group of 89 patients with CCC classified according to the presence of normal or pseudonormal ventricular filling pattern there were some with pseudonormal filling pattern who reported a significant LV systolic impairment in terms of LV dimensions, wall motion score, and LVEF [37]. Diastolic measurements were higher in the groups that were not immunized with any T.cruzi-DNA vaccine than in the those of the healthy control dogs and of the immunized dogs. Specifically, they were higher in the infected animals (B group), in those which were immunized with pBK-CMV plasmid (F group) and in SS- mock immunized (G group). This parameter assesses size of cardiac chambers; if the parameter is high it reflects a volume increase inside the chamber and therefore heart dilation [35]. However, among the groups that received the immunizations only pBCSP-immunized/infected animals had a low volume inside the chamber similar than healthy control dogs; therefore, this demonstrated that T. cruzi gene encoding TcSP trans-sialidase protein is able to protect against heart dilation or to avoid progressive heart damage if it is used as a prophylactic measure. Also, the vaccination could be effective to avoid more severe disorders such as malignant arrhythmias and sudden death, because in accordance with Biolo et al. (2010) [9], the LV systolic dysfunction in ChD may be present immediately after of an early parasympathetic dysautonomia which is a condition where autonomic derangements enhance the dependency of cardiac output increase on volume and shape modifications requiring more ventricular dilation and forceful contraction and also trigger
microcirculatory vasospasm, another important mechanism in Chagas cardiomyopathy. LV systolic diameter high values in animals of B, F and G groups in the present study, which did not receive DNA immunization as preventive action, showed a marked cardiac function damage demonstrated by a significant increase of LV systolic diameter as well as systolic peak time. The systolic measurements by echocardiography assess cardiac function as well as the increased LV chamber size during systole [35]. Two-dimensional, Tissue Doppler and Strain echocardiography have been used in dogs and cats to assess both left and right ventricular function in normal and pathological conditions to estimate intracardiac pressures and myocardial dysfunction, diagnose cardiomyopathies, and assess inter-and intraventricular synchrony [38]. The LV volume remained in reference values in those animals received the TcSP gene as DNA vaccine (D group), suggesting that pBCSP immunization may protect in developing an increase in ventricular mass and end-diastolic volume as adaption mechanism by a chronic volume overload [39] like hypertrophy and marked dilation commonly seen in CCC. Those animals that showed increased LV volume could be developing a left atrial dilation, which is an indicator of the severity of volume overload and increased cardiac pressures. This feature is one of the most important prognostic factors for humans, dogs, and cats with heart disease [38]. In addition, this finding was in accordance with our previous results [29], which showed that macroscopic cardiac alterations like whitish areas in the heart of fibrous consistency, abundant pericardial fluid, LV hypertrophy, thinning of the right ventricular wall, severe tricuspid endocarditis, and heart adherences with trachea and pericardium were absent in all dogs vaccinated with pBCSP plasmid (Table 3).

The overall evaluation of these echocardiographic parameters indicates that the hearts of non-immunized/infected dogs had volume overload with FS % below the references values, and together with the increase of the systolic dimensions can be inferred that those animals with experimental chagasic infection that did not receive any immunizations developed certain degree of HF. This is important because CCC may be detected with or without symptoms. Most investigators combine clinical and EKG findings, cardiomegaly, and systolic dysfunction observed by echocardiography into four groups of progressive heart damage. The recent American College of Cardiology/American Heart
Association staging of disease progression classifies the CCC into A (high risk of HF without structural heart disease), B (structural heart disease without HF), C (structural heart disease with prior or present HF), and D (refractory HF) [40]. Asymptomatic subjects comprise about three quarters of seropositive persons, and those with a normal EKG are referred as being in the indeterminate phase of the disease (stage A). Of 2-18% of patients had been classified in stage B, which no cardiomegaly is present, and LV systolic function is normal. Symptomatic patients with mild to moderate cardiac damage (stage C and/or D) may present arrhythmias, embolism, sudden death, and reversible HF, as well as dilated heart, abnormal LV systolic and diastolic functions, others may have LV apical and other segmental wall abnormalities [7]. Diagnosis by echocardiography in our experimental dogs was a suitable tool to detect structural heart disease that would be able to classify the chagasic patients into A or C stages for the infected ones, and into B or D for the vaccinated-infected dogs, even being asymptomatic animals. This classification in these dogs can be possible because the more severe cardiac abnormalities registered by echocardiography were in the infected-dogs than in immunized-infected dogs. Therefore, it is possible to assume that vaccination helped to avoid the highest level of damage in the heart that lead to HF.

This study, using dogs as an experimental animal model, suggests that more valuable clinical data could be obtained by echocardiography for the management of the patient than those found by electrocardiography, or by evidence of cardiac damage observed postmortem at necropsy or by histopathology. However, our echocardiography recordings were performed from 7 to 19 months after the challenge with Trypanosoma cruzi being a study limitation. Further research should consider using echocardiography at an earlier stage of the ChD in order to follow-up the pathological process, give a timely symptomatic treatment or even try the trypanocidal therapy, because although acute chagasic myocarditis is infrequent, appearing in only 1-5% of those having the acute phase (1 to 5 of every 10,000 infected subjects), there are published echocardiographic series on ChD patients who had abnormal two-dimensional echocardiograms in 52%, and pericardial effusion in 42%. In addition, mean-LVEF was normal (63%) in these patients. Apical or anterior dyskinesis was found in 21%, and only 6% had LV dilation. These findings demonstrate the need to perform echocardiograms to rule out
other cardiac conditions to lead to HF and to evaluate LV dysfunctions during the acute phase of ChD [7].

There is a growing literature demonstrating the potential of non-conventional echocardiography in clinical and subclinical cardiac conditions; speckle tracking echocardiography is capable to discriminate between different causes of LV hypertrophy by ultrasound deformation imaging [41]. However, several limitations should be acknowledged before seeking to try new techniques. The lack of some normal references ranges in veterinary echocardiography precluded the suitable interpretation of values obtained. Nevertheless, our main objective was to compare few imaging parameters among healthy dogs and immunized/infected and non-immunized/infected dogs with findings previously obtained by other diagnostic non-imaging studies.

In conclusion, it has successfully demonstrated that ChD in experimental infected dogs resulted in cardiac dilation, and plasmid DNA vaccination with T. cruzi genes induces moderate protection in immunized dogs avoiding enlargement of cardiac chambers, poor contractibility and HF, especially with pBCSP plasmid. Using echocardiography, structural and functional changes in the chagasic heart were monitored easily and without pain or discomfort to the patient. Also, this method was suitable tool to evaluate the protection of progressive heart damage or heart dilation provided by the prophylactic effect of the DNA vaccine. Transthoracic echocardiography should be the method of choice for characterizing the clinical stages of ChD, such as systolic dysfunction during the acute phase of ChD, and to rule out a rapidly treatable cause of HF (e.g. pericardial effusion).

**Methods**

**Experimental animals**

Animal handling and experimental procedures were approved by the Bioethics Committee of the Instituto Nacional de Cardiología, Ignacio Chávez and performed under the established guidelines of the International Guiding Principles for Biomedical Research involving Animals and the Norma Oficial Mexicana (NOM-0062-ZOO-1999) Technical Specifications for the Care and Use of Laboratory Animals [38]. All sections of this study adhere to the ARRIVE Guidelines for reporting animal research. A completed ARRIVE Guidelines checklist is included in Checklist S1. Fifty-four male and female dogs
(46 Beagle and 8 mongrels), aged 16 (± 2.9) months and weighing 13.79 (± 4.72) kg were used in this study. The Beagle dogs were purchased from Criadero El Atorón (Teotihuacán, Estado de México; Mexico) and the mongrels were provided by the Canine Central Service, both without distinction of sex. Animals were housed in the appropriate Animal Facility, maintained on a 16-h/8-h light-dark cycle under conditions of controlled temperature (20-22 °C) and humidity, fed a standard commercial dry food formulated for dogs and water were available ad libitum. All dogs had a completed schedule of preventive medicine. A standardized enzyme-linked immunosorbent assay (ELISA) [31] used to diagnose ChD in the dogs ruled out any previous infection by *T. cruzi*.

**Immunizations**

The *T. cruzi* genes encoding the *TcSSP4* (amastigote-specific protein) and *TcSP* (*trans*-sialidase protein) antigens were cloned in the commercially available eukaryotic expression vector pBK-CMV (Stratagene) to generate two constructs, pBCSSP4 and pBCSP, respectively, as described previously [26, 27, 39]. The dogs were immunized with doses of 500 mg DNA dissolved in 500 mL saline solution (SS) twice at 15-day intervals by intramuscular injection between the semitendinosus and semimembranosus muscles of the pelvic limbs using a 3-mL syringe with a 21 G x 32 mm needle. The dogs were separated in seven experimental groups (A, B, C, D, E, F and G) (table 1) in order to compare the disease progression after DNA-vaccine administration.

**Study Design**

Experimental groups, the DNA vaccine administered and the time when the echocardiograms were taken are shown in table 1. Fifty-four male and female dogs (46 Beagle and 8 mongrels) were randomly divided into seven groups. Group A (*n* = 6) were non-immunized/non-infected dogs as healthy control; group B (*n* = 16) were positive control, seven of these dogs with 19-month of chronic ChD and the other nine with 11-month of chronic infection; group C (*n* = 6) were non-infected dogs immunized with the three different plasmids (*n* = 2 with pBK-CMV, *n* = 2 with pBCSP and *n* = 2 with PBCSSP4), group D (*n* = 9) were pBCSP plasmid-immunized/infected dogs, group E (*n* = 9) were pBCSSP4 plasmid-immunized/infected dogs, group F (*n* = 4) were pBK-CMV empty cloning vector-immunized/infected dogs, and group G (*n* = 4) were mock-immunized with saline solution (SS) /
infected dogs.

**Trypanosoma cruzi challenge of dogs**

The dogs immunized with DNA or mock-immunized with SS, as well as three animals belonging to positive control group were challenged intraperitoneally two weeks after the last immunization with $5 \times 10^5$ metacyclic trypomastigotes per animal, which were obtained from urine and feces of triatomes and resuspended in SS of a well-characterized Mexican *T. cruzi* Ninoa strain (MHOM/MX/1994/Ninoa [*T. cruzi*]) [40, 41]. The rest of the positive control dogs ($n = 12$) were infected with an inoculum size that ranged from $50 \times 10^3$ to $2 \times 10^6$ metacyclic trypomastigotes. The amount of inoculum had no effect on the severity of the canine ChD [27-29, 31]. The experimental *T. cruzi* infection was confirmed microscopically (parasitemia) in all infected groups by examining freshly isolated blood samples collected from the brachiocephalic vein every third day, observing from 200 to 400 parasites/mL as limit of detection intermittently between day 22 and day 55 postinfection. *T. cruzi* infection was diagnosed by the enzyme-linked immunosorbent assay (ELISA) method two months post-inoculation and was confirmed by indirect immunofluorescence technique in all unimmunized/infected dogs. All animals were monitored clinically by general physical examinations and electrocardiographic studies [27].

**Echocardiography**

Transthoracic echocardiography (Phillips IE33) with a 2-5-3.5 MHz probe was performed in all dogs during the chronic stage of infection (8, and 11 or 19 months after inoculation in vaccinated or chronically infected groups, respectively) in order to detect and compare morphological changes in the dogs’ heart. Most of the animals were positioned in dorsal and right or left lateral decubitus without any chemical restriction during the study; this could be achieved with previous training through daily manipulation in examination tables in order to keep the dogs in the desired position to carry out the study. In those few cases which chemical restriction was necessary, acepromazine at light sedation dose of 2.75 mg/kg of body weight was used. Image acquisition was done by means of a long parasternal axis; two- and four- chamber apical view in bidimensional mode. Heart rate was
simultaneously calculated from the preceding R-to-R interval on the electrocardiogram. End-diastolic and end-systolic diameters as well as end-diastolic and end-systolic free wall thickness of the left ventricle were measured. LV septum walls, LV end-diastolic volume and the systolic peak time were also calculated. Parameters of LV systolic function, i.e. fractional shortening (FS %), and left ventricular ejection fraction (LVEF %) calculated from the end-diastolic and end-systolic volumes in the standard four-chamber long-axis 2D-echo views were also recorded following the formula [end-diastolic volume - end-systolic volume] / end-diastolic volume × 100.

**Euthanasia**

At the end of the echocardiographic study, at 11 or 19 months post infection, chronic chagasic dogs were euthanized to perform other studies *post-mortem* such as macroscopic evaluation and histology in order to determine cardiac damage [29]. Briefly, the euthanasia method indicated for dogs was used according to the Norma Oficial Mexicana (NOM-033-SAG/ZOO-2014) Humane Sacrificing of Domestic and Wild Animals [42], through intravenous injection of barbiturate derivatives at a dose of 30 mg/kg, and then a lethal dose of 15% potassium chloride administered intravenously. This protocol was approved by the Committee for the Care and Use of Animals of Laboratory (CICUAL, for its acronyms in Spanish) and the Bioethics Committee of the Instituto Nacional de Cardiología, Ignacio Chávez (Registration number: 08-578) and performed under the established guidelines of the International Guiding Principles for Biomedical Research involving Animals.

**Statistical analysis**

All data obtained were analyzed with commercially available software (SPSS version 22.0). One-way analysis of variance was used to examine the vaccination effect within each group for echocardiographic variables. When a significant difference was detected, the Tukey test was used to compare the means. Results were expressed as mean ± standard deviation. The level of significance in all tests was set at *P* ≤ 0.05.

**Declarations**

**Supplementary information**

**Additional file 1:** Checklist S1
Abbreviations

ChD: Chagas disease; HF: heart failure; CCC: chronic Chagas’ cardiomyopathy; EKG: electrocardiogram; pBCSP: pBK-CMV plasmid containing a trans-sialidase gene of T. cruzi, named TcSP gene; pBCSSP4: pBK-CMV plasmid containing an amastigote-specific gene of T. cruzi, named TcSSP4 gene; pBK-CMV: name of the commercially eukaryotic expression vector (Stratagene) used as cloning plasmid; ELISA: enzyme-linked immunosorbent assay; SS: saline solution; LV: left ventricular; LVEF: left ventricular ejection fraction; FS: fractional shortening.

Acknowledgments

We thank Susana Kolb, an English-speaking colleague/English-language instructor from Writing Workshop for Academic Articles in English for Graduate Tutors staff at Universidad Nacional Autónoma de México for her assistance with English editing of the manuscript.

Consent for publication

Not applicable.

Authors’ contributions

A-F and O. R-M directed the research, reviewed the data, reviewed the manuscript critically, directed revisions and provided statistical analysis. F-J R. and J. V-B recorded the echocardiographic data by the management of the echocardiographic equipment and participated in drafting the manuscript. E. P-B, ML. M-G and E. V-B handled and held all dogs, helped to their physical restraint, and created the databases from recordings obtained. All authors read and approved the final version of the manuscript.

Funding

The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors. The affiliation institute’ resources were the only ones that financed this study.

Availability of data and materials

Data are available upon request or they will be share in additional supporting files or publicly available repositories.
**Ethics approval and consent to participate**

Studies in dogs were done in accordance with the current International Guiding Principles for Biomedical Research involving Animals and the Normas Oficiales Mexicanas NOM-0062-ZOO-1999:Technical Specifications for the Care and Use of Laboratory Animals and NOM-0033-ZOO-1995: Humane Sacrificing of Domestic and Wild Animals. Animal handling and experimental procedures were approved by the Committee for the Care and Use of Animals of Laboratory (CICUAL, for its acronyms in Spanish) and the Bioethics Committee of the Instituto Nacional de Cardiología, Ignacio Chávez (Registration number: 08-578).

**Competing interests**

The authors declare that they have no competing interests.

**References**

1. World Health Organization (WHO). Weekly epidemiological record. 2015;90:33-44.
   
   Available in: https://www.who.int/wer/2015/wer9006.pdf?ua=1.

2. Machado FS, Jelicks LA, Kirchhoff LV, Shirani J, Nagajyothi F, Mukherjee S, et al. Chagas Heart Disease: Report on Recent Developments. Cardiol Rev. 2012;20(2):53-65.

3. Pereira Júnior C de B, Markman Filho B. Clinical and echocardiographic predictors of mortality in chagasic cardiomyopathy – systematic review. Arq Bras Cardiol. 2014;102(106):602-10.

4. Muratore CA, Baranchuk A. Current and emerging therapeutic options for the treatment of chronic chagasic cardiomyopathy. Vasc Health Risk Manag. 2010;6:593-601.

5. Hagar JM, Rahimtoola, SH. Chagas’ heart disease. Curr Probl Cardiol. 1995;20(12):825-928.

6. Acquatella, H. Regional diastolic dysfunction in chronic Chagas’ heart disease. Eur J
7. Acquatella, H. Echocardiography in Chagas heart disease. Circulation. 2007;115(9):1124-31.

8. Viotti RJ, Vigliano C, Laucella S, Lococo B, Petti M, Bertocchi G, et al. Value of echocardiography for diagnosis and prognosis of chronic Chagas disease cardiomyopathy without heart failure. Heart. 2004; 90(6):655-60.

9. Biolo A, Ribeiro AL, Clausell N. Chagas Cardiomyopathy - Where Do We Stand After a Hundred Years? Progr Cardiovasc Dis. 2010;52(4):300-16.

10. Gürtler RE, Cecere MC, Lauricella MA, Cardinal MV, Kitron U, Cohen JE. Domestic dogs and cats as sources of Trypanosoma cruzi infection in rural northwestern Argentina. 2007;134(Pt 1):69-82.

11. Esch KJ, Petersen CA. Transmission and epidemiology of zoonotic protozoal disease of companion animals. Clin Microbiol Rev. 2013;26(1):58-85.

12. Mendonça PHB, da Rocha RFDB, Moraes JBB, LaRocque-de-Freitas IF, Logullo J, Morrot A, et al. Canine macrophage DH82 cell line as a model to study susceptibility to Trypanosoma cruzi Front Immunol. 2017;8:604.

13. Chetboul V, Serres F, Gouni V, Tissier R, Pouchelon JL. Radial strain and strain rate by two-dimensional speckle tracking echocardiography and the tissue velocity based technique in the dog. J Vet Cardiol. 2007;9(2):69-81.

14. De Lana M, Chiari E, Tafuri WL. Experimental Chagas’ disease in dogs. Mem Inst Oswaldo Cruz. 1992;87(1):59-71.

15. Caliari MV, de Lana M, Cajá RA, Carneiro CM, Bahia MT, Santos CA, et al. Immunohistochemical studies in acute and chronic canine chagasic cardiomyopathy. Virchows Arch. 2002;441(1):69-76.

16. Coura-Vital W, Carneiro CM, Martins HR, de Lana M, Veloso VM, Teixeira-Carvalho A,
et al. *Trypanosoma cruzi*: immunoglobulin isotype profiles during the acute phase of canine experimental infection with metacyclic or blood trypomastigotes. Exp Parasitol. 2008;120(3):269-74.

17. Guedes PM, Veloso VM, Mineo TW, Santiago-Silva J, Crepalde G, Caldas IS, et al. Hematological alterations during experimental canine infection by *Trypanosoma cruzi*. Rev Bras Parasitol Vet. 2012;21(2):151-56.

18. Quijano-Hernández IA, Castro-Barcena A, Vázquez-Chagoyán JC, Bolio-González ME, Ortega-López J, Dumonteil E. Preventive and therapeutic DNA vaccination partially protect dogs against an infectious challenge with *Trypanosoma cruzi*. Vaccine 2013;31(18):2246-52.

19. Duz AL, Vieira PM, Roatt BM, Aguiar-Soares RD, Cardoso JM, Oliveira FC, et al. The Tcl and TcII *Trypanosoma cruzi* experimental infections induce distinct immune responses and cardiac fibrosis in dogs. Mem Inst Oswaldo Cruz. 2014;109(8):1005-13.

20. Hartley AN, Cooley G, Gwyn S, Orozco MM, Tarleton RL. Frequency of IFNg-producing T cells correlates with seroreactivity and activated T cells during canine *Trypanosoma cruzi* Vet Res. 2014;45:6.

21. Aparicio-Burgos JE, Zepeda-Escobar JA, de Oca-Jimenez RM, Estrada-Franco JG, Barbabosa-Pliego A, Ochoa-García L, et al. Immune protection against *Trypanosoma cruzi* induced by TcVac4 in a canine model. PLoS Negl Trop Dis. 2015;9:e0003625.

22. Floridia-Yapur N, Monje Rumi M, Ragone P, Lauthier JJ, Tomasini N, Alberti D'Amato A, et al. The TcTASV proteins are novel promising antigens to detect active *Trypanosoma cruzi* infection in dogs. Parasitology. 2016;143(11):1382-89.

23. Santos FM, Mazzeti AL, Caldas S, Gonçalves KR, Lima WG, Torres RM, et al. Chagas cardiomyopathy: The potential effect of benznidazole treatment on diastolic dysfunction and cardiac damage in dogs chronically infected with *Trypanosoma cruzi*. 
24. Caldas IS, Diniz LF, Guedes PMDM, Nascimento ÁFDSD, Galvão LMDC, Lima WG, et al. Myocarditis in different experimental models infected by Trypanosoma cruzi is correlated with the production of IgG1 isotype. Acta Trop. 2017;167:40-9.

25. Rodríguez-Morales O, Monteón-Padilla V, Carrillo-Sánchez SC, Rios-Castro M, Martínez-Cruz M, Carabarín-Lima A, et al. Experimental vaccines against Chagas disease: A journey through history. J Immunol Res. 2015;489758.

26. Arce-Fonseca M, Ramos-Ligonio A, López-Monteón A, Salgado-Jiménez B, Talamás-Rohana P, Rosales-Encina JL. A DNA vaccine encoding for TcSSP4 induces protection against acute and chronic infection in experimental Chagas disease. Int J Biol Sci. 2011;7(9):1230-8.

27. Rodríguez-Morales O, Pérez-Leyva MM, Ballinas-Verdugo MA, Carrillo-Sánchez SC, Rosales-Encina JL, Alejandre-Aguilar R, et al. Plasmid DNA immunization with Trypanosoma cruzi genes induces cardiac and clinical protection against Chagas disease in the canine model. Vet Res. 2012;43:79.

28. Arce-Fonseca M, Ballinas-Verdugo MA, Zenteno ER, Suárez-Flores D, Carrillo-Sánchez SC, Alejandre-Aguilar R, et al. Specific humoral and cellular immunity induced by Trypanosoma cruzi DNA immunization in a canine model. Vet Res. 2013;44:15.

29. Rodríguez-Morales O, Carrillo-Sánchez SC, García-Mendoza H, Aranda-Frausto A, Ballinas-Verdugo MA, Alejandre-Aguilar R, et al. Effect of the plasmid-DNA vaccination on macroscopic and microscopic damage caused by the experimental chronic Trypanosoma cruzi infection in the canine model. Biomed Res Int. 2013;2013:826570.

30. Boon JA. Two Dimensional and M-Mode Echocardiography for the Small Animal Practitioner. Jackson, WY, USA: NewMedia; 2002.
31. Rodríguez-Morales O, Ballinas-Verdugo MA, Alejandre Aguilar R, Reyes PA, Arce-Fonseca M. Trypanosoma cruzi connatal transmission in dogs with Chagas disease: Experimental case report. Vector Borne Zoonotic Dis. 2011;11(10):1365-70.

32. Allam LE, Onsy AM, Ghalib HA. Right ventricular outflow tract systolic excursion and fractional shortening: can these echocardiographic parameters be used for the assessment of right ventricular function? J Cardiovasc Echogr. 2017;27(2):52-8.

33. Barros ML, da Costa Rocha MO, Ribeiro AP, Machado FS. Tissue Doppler imaging enables the identification of diastolic dysfunction of pseudonormal pattern in Chagas' disease. J Am Soc Echocardiogr. 2001;14(5):353-9.

34. Baron Toaldo M, Guglielmini C, Diana A, Sarcinella F, Cipone M. Feasibility and reproducibility of echocardiographic assessment of regional left atrial deformation and synchrony by tissue Doppler ultrasonographic imaging in healthy dogs. Am J Vet Res. 2014;75(1):59-66.

35. Bening C, Hamouda K, Leyh R. Sex differences in volume overload in skinned fibers. BMC Cardiovasc Disord. 2016;16(1):197.

36. Hunt SA. ACC/AHA 2005 guideline update for the diagnosis and management of chronic heart failure in the adult: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines (Writing Committee to Update the 2001 Guidelines for the Evaluation and Management of Heart Failure). J Am Coll Cardiol. 2005;46(6):e1-e82.

37. García-Álvarez A, Sitges M, Regueiro A, Poyatos S, Jesus Pinazo M, Posada E, et al. Myocardial deformation analysis in Chagas heart disease with the use of speckle tracking echocardiography. J Card Fail. 2011;17(12):1028-34.

38. Norma Oficial Mexicana NOM-0062-ZOO-1999: Especificaciones para el cuidado y uso de animales de laboratorio. Mexico City: 1999, Diario Oficial de la Federación.
Available in:
https://www.gob.mx/cms/uploads/attachment/file/203498/NOM-062-ZOO-1999_220801.pdf

39. Salgado-Jiménez B, Arce-Fonseca M, Baylón-Pacheco L, Talamás-Rohana P, Rosales-Encina JL. Differential immune response in mice immunized with the A, R or C domain from TcSP protein of Trypanosoma cruzi or with the coding DNAs. Parasite Immunol. 2013;35(1):32-41.

40. Monteón VM, Furuzawa-Carballeda J, Alejandre-Aguilar R, Aranda-Frausto A, Rosales-Encina JL, Reyes PA. American trypanosomiosis: in situ and generalized features of parasitism and inflammation kinetics in a murine model. Exp Parasitol. 1996;83(3):267-74.

41. World Health Organization (WHO) Technical Report Series. Control of Chagas Disease. Second report of the WHO expert committee. Geneva: World Health Organization, 2002;1-109. Available in:
http://apps.who.int/iris/bitstream/10665/42443/1/WHO_TRS_905.pdf

42. Norma Oficial Mexicana NOM-033-ZOO-1995, Sacrificio humanitario de los animales domésticos y silvestres. Mexico City, Mexico: Diario Oficial de la Federación; 1995.

Tables
Table 1. Study Design for T. cruzi experimentally infected dogs immunized with DNA vaccine containing the genes encoding a trans-sialidase protein (pBCSP) or an amastigote-specific glycoprotein (pBCSSP4).
| Group | Group description and n | Vaccine | Post-infection (PI) vaccination (PV) time for the echocardiographic study |
|-------|-------------------------|---------|---------------------------------------------------------------------|
| A     | Non-immunized/Non-infected (healthy control) \( n = 6 \) | None | NA |
| B     | Chronically infected control \( n = 16 \) | None | 19 months \( n = 11 \) months \( n = 7 \) PI |
| C     | Each plasmids-immunized/Non-infected \( n = 6 \) | pBCSP \( n = 2 \) pBCSSP4 \( n = 2 \) pBK-CMV \( n = 2 \) | 8-11 months PI |
| D     | Immunized with \( TcSP \) gene/Infected \( n = 9 \) | pBCSP | 8 months PI |
| E     | Immunized with \( TcSSP4 \)/Infected \( n = 9 \) | pBCSSP4 | 8 months PI |
| F     | Immunized with cloning vector/Infected \( n = 4 \) | pBK-CMV | 8 months PI |
| G     | Mock-immunized with saline solution/Infected \( n = 4 \) | Saline Solution | 8 months PI |

NA = Not Applicable, because they were neither infected nor vaccinated; pBCSP = plasmid carrying \( TcSP \)
gene; pBCSSP4 = plasmid carrying TcSSP4 gene; pBK-CMV= empty cloning vector.

Table 2. Cardiovascular parameters in *T. cruzi* experimentally infected dogs immunized with DNA vaccine containing the genes encoding a *trans*-sialidase protein (pBCSP) or an amastigote-specific glycoprotein (pBCSSP4).

| Group                        | Fractional shortening (FS) (%) | Left ventricular ejection fraction (LVEF) (%) | Left ventricular (LV) diastolic diameter (mm) | Left ventricular (LV) systolic diameter (mm) |
|------------------------------|--------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| A (healthy control)          | 39.29±3.76                     | 77.60±4.22                                    | 28.70±2.84                                    | 17.42±1.98                                    |
| A (Non-imm/Non-inf)          |                                |                                               |                                               |                                               |
| B (Inf)                      | 31.64±8.12*                    | 50.29±6.59*                                   | 34.19±7.15*                                   | 23.35±5.83*                                   |
| C (P-imm/Non-inf)            | 41.19±4.13                     | 78.20±5.17                                    | 26.30±2.30                                    | 15.44±1.46                                    |
| D (pBCSP-imm/Inf)            | 33.57±6.31                     | 54.62±9.19*                                   | 29.90±2.43                                    | 20.68±2.77                                    |
| E (pBCSSP4-imm/Inf)          | 33.94±9.17                     | 58.45±12.03*                                  | 30.06±1.88                                    | 20.84±2.36                                    |
| F (pBK-CMV-imm/Inf)          | 31.55±1.13*                    | 52.33±5.37*                                   | 31.70±34.59*                                  | 21.67±2.80*                                   |
| G (SS mock-imm/Inf)          | 27.32±8.29*                    | 47.67±4.93*                                   | 30.2±1.13*                                    | 21.80±4.81*                                   |

All data are expressed as means and standard deviations. Statistical significances (*P*≤0.05) are assigned as * when there was a significant difference with group A control healthy dogs and/or with reference values [30]. Bold values belong to the groups which were vaccinated and did not show significant difference with group A control healthy dogs. bpm = beats per minute, imm = immunized, inf or Inf = infected, P-imm = each plasmids-immunized, pBCSP = plasmid carrying TcSP gene, pBCSSP4 = plasmid carrying TcSSP4 gene, pBK-CMV = empty cloning vector, and SS = saline solution.

Table 3. Other findings obtained previously in the same *T. cruzi* experimentally infected dogs immunized with DNA vaccine containing the genes encoding a *trans*-sialidase protein (pBCSP) or an amastigote-specific glycoprotein (pBCSSP4).
| Group (description) | Electrocardiographic abnormalities (percentage of animals with these alterations) [27, 29, 31] | Parasitemia [27, 31] | Post-mortem macroscopic alterations (percentage of animals with these findings) [29] | Cardio- and/or Splenomegaly [29] |
|---------------------|-------------------------------------------------------------|------------------|---------------------------------------------------------------------------------|-----------------------------|
| A (healthy control) | None (healthy control) | Negative | None | None |
| B (Inf) | Ischemia and conduction block LV hypertrophy (100%) | Positive at 22-55 days post-infection | Whitish areas in heart of fibrous consistency and abundant pericardial fluid (69%) | Cardiomegaly |
| C (P-imm/Non-inf) | None | Negative | None | None |
| D (pBCSP-imm/Inf) | AV Block (33%) and LV enlargement (11%) | Positive at 32-46 days post-infection | None | Cardiomegaly |
| E (pBCSSP4-imm/Inf) | MIMI (11%) and Second-degree AV block (11%) | Positive at 32-46 days post-infection | Heart with adhesions in trachea and pericardium, splenic thickened walls and whitish areas in spleen and heart (11%) | None |
| F (pBK-CMV-imm/Inf) | LV enlargement (75%) and Right BBB (25%) | Positive at 31-46 days post-infection | Abundant pericardial fluid and whitish areas in spleen (75%), Ascites, megaesophagus, LV hypertrophy, thinning of the RV wall and severe tricuspid endocarditis (25%) | Cardiomegaly Splenomegaly |
| G (SS mock-imm/Inf) | VPC (25%), pericardial effusion (50%), and myocardial infarction and/or pericarditis, and RV enlargement (50%) | Positive at 21-55 days post-infection | Whitish areas in heart of fibrous consistency and abundant pericardial fluid (75%) | Cardiomegaly Splenomegaly |

imm = immunized, Inf or Inf = infected, P-imm = each plasmids-immunized, pBCSP = plasmid carrying TcSP gene, pBCSSP4 = plasmid carrying TcSSP4 gene, pBK-CMV= empty cloning vector, and
SS = saline solution. FS = fractional shortening, LVEF = left ventricular ejection fraction, LV = left ventricular, MIMI = Microscopic intramural myocardial infarctions, N/D = not determined.

Supplementary Files
This is a list of supplementary files associated with this preprint. Click to download.
Checklist S1.pdf