1 **Title**

- 2 Spatial metabolomics identifies localized chemical changes in heart tissue during chronic cardiac
- 3 Chagas disease

4 Short Title

5 Spatial metabolomics of chronic cardiac Chagas Disease

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27 Abstract

28 Chagas disease (CD) is one of thirteen neglected tropical diseases caused by the parasite 29 Trypanosoma cruzi. CD is a vector-borne disease transmitted by triatomines but CD can also be 30 transmitted through blood transfusions, organ transplants and congenital transmission. While 31 endemic to Latin America, T. cruzi infects 7-8 million people worldwide and can induce severe 32 cardiac symptoms including apical aneurysms, thromboembolisms and arrhythmias during the 33 chronic stage of CD. However, these cardiac clinical manifestations and CD disease 34 pathogenesis are not fully understood. Using spatial metabolomics (chemical cartography), we 35 sought to understand the localized impact of infection on the cardiac metabolome of mice 36 chronically infected with two divergent T. cruzi strains. Our data showed chemical differences in 37 localized cardiac regions upon chronic T. cruzi infection, indicating that parasite infection 38 changes the host metabolome at select sites in chronic CD. These sites were distinct from the 39 sites of highest parasite burden. In addition, we identified acylcarnitines and phosphocholines as 40 discriminatory chemical families within each heart region, comparing infected and uninfected 41 samples. Overall, our study indicated overall and positional metabolic differences common to 42 infection with different T. cruzi strains, and identified select infection-modulated pathways. 43 These results provide further insight into CD pathogenesis and demonstrate the advantage of a 44 spatial perspective to understand infectious disease tropism.

45 Author Summary

46 Chagas disease (CD) is a tropical disease caused by the parasite *Trypanosoma cruzi*. CD 47 originated in South America; however, there are now 7-8 million people infected worldwide due 48 to population movements. CD is transmitted through a triatomine vector, organ transplants, 49 blood transfusions and congenital transmission. It occurs in two stages, an acute stage (usually 50 asymptomatic) and the chronic stage. Chronic stage CD presents with severe cardiac symptoms 51 such as heart failure, localized aneurysms and cardiomyopathy. Unfortunately, what causes 52 severe cardiac symptoms in some individuals in chronic CD is not fully understood. Therefore, 53 we used liquid chromatography-tandem mass spectrometry to analyze the heart tissue of 54 chronically T. cruzi-infected and uninfected mice, to understand the impact of infection on the 55 tissue metabolome. We identified discriminatory small molecules related to T. cruzi infection. 56 We also determined that regions with the highest parasite burden are distinct from the regions 57 with the largest changes in overall metabolite profile; these locations of high metabolic perturbation provide a molecular mechanism to why localized cardiac symptoms occur in CD. 58 59 Overall, our work gives insight to chronic cardiac CD symptom development and shapes a 60 framework for novel treatment and biomarker development.

61 Introduction

62 Chagas disease (CD) is a parasitic disease caused by the protozoan *Trypanosoma cruzi* 63 and is one of the designated "neglected tropical diseases" [1]. T. cruzi is endemic to Latin 64 America and infects 7-8 million people worldwide [1]. An estimated 300,000 infections have 65 been recorded in the United States due to a large Latin American immigrant population and 66 endemic transmission [2–4]. CD is primarily transmitted through triatomine insects of the 67 Triatoma and Rhodnius genera [2]. Non-vectorial modes of transmission involve blood 68 transfusion, transplacental transmission, and food and drink contaminated with T. cruzi [1]. The 69 T. cruzi life cycle includes three main stages: epimastigotes, trypomastigotes and amastigotes. T. 70 *cruzi* in the insect vector undergoes transformation from trypomastigotes to epimastigotes in the 71 midgut, and then migrates to the hindgut and differentiates into infective trypomastigotes [1]. 72 Upon triatomine defecation on the human host, the infective trypomastigotes enter the host 73 through scratching or rubbing of the bite wound, or through eyes and mucosal surfaces [1]. 74 Following mammalian host cell infection, trypomastigotes differentiate into amastigotes, which 75 proliferate and subsequently transform into trypomastigotes [1]. CD has two disease stages: 76 acute and chronic [1,2]. The acute stage is usually asymptomatic, or presents with non-specific 77 symptoms (fever, malaise) [1,2]. 20-30% of infected individuals will then progressively develop 78 clinical manifestations of chronic CD, including cardiomegaly, cardiac arrhythmias, apical 79 aneurysms, megacolon, and megaesophagus [2]. T. cruzi infections are treated with either 80 benznidazole or nifurtimox; however, these treatments cause significant adverse effects to the 81 point that up to 30% of treated individuals fail to complete the full treatment course [5,6]. 82 CD was previously considered to have an autoimmune etiology, but parasite persistence 83 has now conclusively been demonstrated to be required for disease pathogenesis [7]. Along with parasite persistence, chronic pro-inflammatory responses, including cytokine release and CD8+
T cell- mediated cytotoxicity, contribute to tissue damage [8]. A heterogeneity of interacting
parasite-host factors, including *T. cruzi* strain, load and tissue tropism, host genetic background,
and mode of infection, influence the clinical outcomes of the disease [9,10]. However, CD
disease pathogenesis is not yet completely understood [2]. A holistic understanding of the
molecular pathways involved in disease progression could help identify new drug development
avenues and outcome-predictive biomarkers.

91 Metabolites are the final products of mRNA and protein expression and protein activity, 92 thus providing information closely linked to phenotype [11]. Metabolic pathways are druggable. 93 They also change dynamically in response to disease [12,13]. As such, an improved 94 understanding of metabolism in CD may lead to new avenues for drug development and CD 95 patient monitoring. Acute T. cruzi-infection affects in vitro and in vivo host metabolic pathways, 96 including decreasing mitochondrial oxidative phosphorylation-mediated ATP production [8,14– 97 16]. In addition, acute T. cruzi-infected mice heart tissue and plasma showed significant up- or 98 down-regulation of certain metabolic pathways, such as glucose metabolism (glucose levels 99 elevated in heart tissue and lowered in plasma over time), tricarboxylic acid cycle (TCA) 100 (decrease in select TCA metabolites in the heart tissue and a decrease in all TCA metabolites in 101 plasma), lipid metabolism (increased long-chain fatty acids in the heart tissue and the opposite in 102 plasma), and phospholipid metabolism (high accumulation of phosphocholine precursor 103 metabolites in the heart in comparison to plasma) [14]. Prior analysis of hearts from acutely-104 infected mice also showed that cardiac metabolite profiles reflected disease severity, with 105 changes in cardiac acylcarnitines and phosphatidylcholines predictive of acute infection outcome 106 [8]. Metabolomic analysis of chronic CD has been limited to serum and gastrointestinal tract

107 samples [17][18]. Serum analysis demonstrated significant changes in amino acid and lipid 108 metabolism, particularly acylcarnitines, sphingolipids, and glycerophospholipids [17]. Analysis 109 of GI tract samples observed persistent metabolic perturbations in the oesophagus and large 110 intestine in chronic CD, including infection-induced elevation of acylcarnitines, 111 phosphatidylcholines and amino acid derivatives [18]. However, metabolic changes in the heart 112 may differ from those in the circulation or in the GI tract [14]. It is therefore essential to perform 113 metabolomic analysis of tissues collected from the heart in chronic CD. Many sudden fatalities 114 due to chronic cardiac CD are often attributed to apical aneurysms which occur at the bottom of 115 the heart [19,20]. We therefore focused on liquid chromatography-tandem mass spectrometry-116 based metabolomic analysis of horizontally-sectioned hearts from mice chronically infected with 117 T. cruzi strains CL and Sylvio X10/4. These samples had previously been analyzed in terms of 118 positional differences (heart apex vs heart base), but not in the context of metabolic changes 119 associated with chronic T. cruzi infection [8]. Overall, we observed significant localized 120 chemical differences associated with infection, with a disconnect between parasite localization 121 and overall positional metabolic perturbations. Our data also showed infection-induced 122 variations in acylcarnitine and phosphocholine chemical families.

123

124 Methods

125 Ethics statement

All vertebrate animal studies were performed in accordance with the USDA Animal
Welfare Act and the Guide for the Care and Use of Laboratory Animals of the National Institutes
of Health. The protocol was approved by the University of California San Diego Institutional
Animal Care and Use Committee (protocol \$14187).

130 In vivo experimentation

131 All *in vivo* experimentation, sample collection and qPCR analysis was conducted and

132 previously reported in [8].

133 Metabolite extraction and UHPLC-MS/MS

134 The two-step procedure for metabolite extraction was adapted from Want *et al* [21] and

135 was conducted in McCall *et al* [8], with "aqueous" and "organic" extracts referring to the

136 metabolites recovered from the first 50% methanol extraction and the second 3:1

137 dichloromethane-methanol extraction, respectively. UHPLC-MS/MS analysis was conducted

138 using a Thermo Scientific UltiMate 3000 Ultra High Performance Liquid Chromatography with

139 a C8 LC column and MS/MS detection on a Maxis Impact HD QTOF mass spectrometer (Bruker

140 Daltonics), as previously reported [8][21].

141 LC-MS/MS data analysis

142 Data processing was performed as previously reported using Optimus, July 21, 2016

143 version [8][22]. Total ion current (TIC) normalization was performed in R studio. Principal

144 coordinate analysis (PCoA) was performed on total ion current (TIC) normalized MS1 feature

145 data table using the Bray-Curtis-Faith dissimilarity metric using QIIME1 [23], for both organic

146 and aqueous extractions combined. The three-dimensional PCoA plots were visualized in

147 EMPeror [24]. Heart three-dimensional modelling was completed using 'ili' (<u>http://ili.embl.de/</u>)

148 [25]using a three dimensional heart model from 3DCADBrowser.com

149 (http://www.3dcadbrowser.com/).

Global Natural Products Social Molecular Networking (GNPS) was used to perform
molecular networking according to the parameters in Table 1 [26]. Cytoscape 3.7.0. was used to

152 visualize the molecular networks [27].

153	Table 1: Global Natural Products Social Molecular Networking (GNPS)	parameters.
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Parent mass	2.0 Da
MS/MS fragment ion tolerance	0.5 Da
Cosine score	0.7
Minimum matched fragment ions	6
Analog search against the library	not allowed
Network TopK	10
Maximum Connected Component Size	100
Minimum cluster size	2
Score threshold	0.7
Library search Min Matched Peaks	6
Filter precursor window	filter
Filter peaks in 50Da window	filter

154

Random forest analysis was performed in Jupyter Notebook using R with the number of trees set to 500. Random forest classifier cutoff was based on ranked variable importance score of differential metabolites in combination with unadjusted p-values<0.05 where 4 consecutive non-significant unadjusted p-values defined the cutoff. FDR-corrected Mann Whitney p<0.05 for all positions was also used as an alternate method to determine significant metabolite differences.

160	Venn diagrams were used to visualize the unique and common metabolites differential between
161	CL and Sylvio X10/4 infection, compared to uninfected samples, based on heart segment
162	positions, random forest classifier for all positions, and FDR-corrected Mann Whitney p<0.05
163	for all positions, using http://bioinformatics.psb.ugent.be/ webtools/Venn/.
164	
165	Data availability
166	Metabolomics data has been deposited in MassIVE (<u>http://massive.ucsd.edu/</u> , accession
167	#MSV000080450). Molecular networks can be accessed at
168	https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=f16bc44c3d5040d098c978823f50c68f (all
169	samples, Aqueous extract),
170	https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=5f8af6d62d8549358966f3896a81063a (all
171	samples, Organic extract).
172	
173	Results
174	The purpose of this study was to compare the metabolic impact of chronic T. cruzi
175	infection in the mouse heart between divergent T. cruzi strains and between cardiac regions. To
176	do so, we analyzed previously-collected positive mode LC-MS/MS data from mice chronically
177	infected with <i>T. cruzi</i> strain CL or Sylvio X10/4 [8]. While this prior study focused on positional
178	differences between uninfected samples and on the impact of acute infection on the cardiac
179	metabolite profile, here we specifically focused on the impact of chronic infection on the cardiac
180	metabolite profile.

181 We observed a clear distinction in the impact of *T. cruzi* infection on the overall
182 metabolite profile between *T. cruzi* strains by heart position (Fig 1, S1 figure, S2 figure). As

183	previously described [8], parasite burden was highest at the base of the heart (position A) for
184	strain CL and central positions (position C) for strain Sylvio X10/4 (Fig 1A). PERMANOVA
185	analysis indicated that the highest significant perturbation in the overall metabolite profile
186	occurred at central positions for strain CL infection (PERMANOVA analysis of Bray-Curtis-
187	Faith distance matrix R^2 =0.20813, p-value=0.004 at position C) and at apical positions for strain
188	Sylvio X10/4 infection (PERMANOVA analysis Bray-Curtis-Faith distance matrix R ² =0.27923,
189	p-value=0.014 at position D) (Fig 1 B). Strikingly, in both cases chemical disturbance was
190	greatest at sites distinct from the- highest parasite burden, which corroborates our observations in
191	the context of chronic gastrointestinal T. cruzi infection in mice [18]. The localization of
192	chemical disturbance also provides a molecular mechanism explaining the apical aneurysms
193	observed in CD patients.

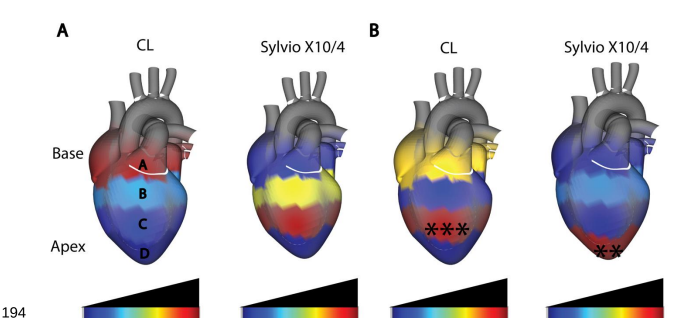
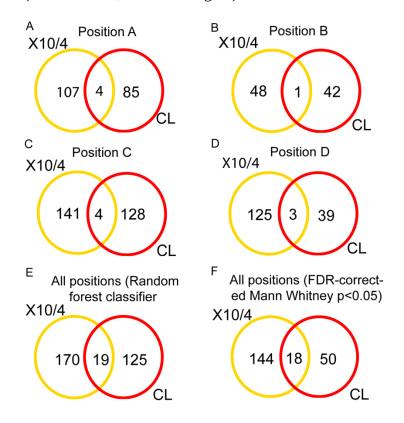


Fig 1. Disconnect between sites of parasite persistence and metabolic alterations in chroniccardiac CD.

197 (A) Median cardiac parasite burden, as determined by qPCR. Parasite burden was highest at the 198 heart base (position A) for strain CL and central heart segments (position C) for strain Sylvio X10/4, indicating parasite strain-specific differences in parasite tropism. (B) Statistically 199 200 significant perturbations in the overall metabolite profile between uninfected and strain CL-201 infected (left), and between uninfected and strain Sylvio X10/4-infected mice (right) were 202 analyzed using PERMANOVA. The highest significant metabolite perturbation was at central 203 heart segments (position C) for strain CL (***, p < 0.001 by PERMANOVA) and at the heart 204 apex (position D) for strain Sylvio X10/4 (**, p < 0.01 by PERMANOVA). 205 To identify the specific cardiac metabolites spatially perturbed by infection, initially we 206 207 built a random forest classifier for each position, each strain and each extraction method,

- comparing to uninfected matched control samples (S1-S6 tables). We first assessed the overlap
- between the top-ranked most differential metabolites by random forest for the two different

- 210 strains, as described in Methods. Limited overlap of these significant metabolites was observed
- 211 between strains (Figure 2). However, annotation of these differential metabolites using
- 212 molecular networking through the GNPS platform [26] revealed that while differing in terms of
- 213 m/z, many were part of the same chemical families, including acylcarnitines and phosphocholine
- 214 (S1 S6 Table; S3 and S4 figure).



215

216 Fig 2. Limited overlap of specific differential metabolites between strains.

Yellow and red circles represent differential metabolites between strain Sylvio X10/4-infected
and matched uninfected controls, and between strain CL-infected and matched uninfected
controls, respectively. Intersect are metabolites impacted by infection in both strains. (A-D)
Differential metabolites for each strain, at given heart positions. (E) Metabolites impacted by
infection with each strain, irrespective of position, as determined by random forest classifier,
with variable importance score cutoff as described in Methods. (F) Metabolites impacted by

infection with each strain, irrespective of heart position using FDR-corrected Mann Whitney
 p<0.05 cutoff.

225

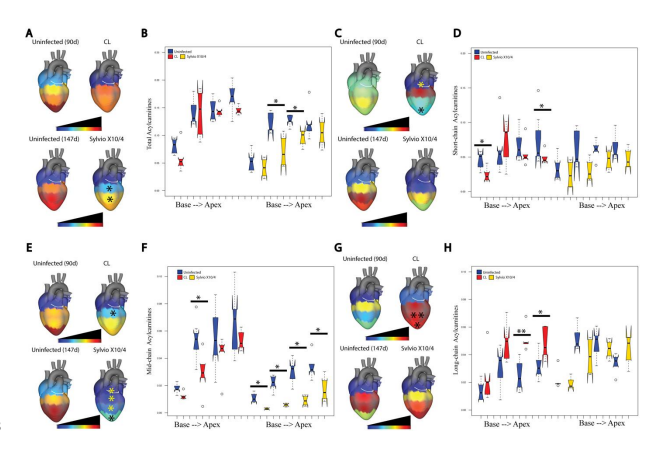
Random forest classifier identified several acylcarnitines and phosphocholines as
impacted by infection (S1 - S5 Tables). Both chemical families play a major role in several
biochemical pathways. Carnitine serves as a shuttling mechanism for fatty acids, in the form of
acylcarnitines, from the cytosol into the matrix of the mitochondria for beta-oxidation [28].
Phosphocholines are major components of lipid metabolism, cell membrane structure, and
choline production, the latter of which is essential for select amino acid and neurotransmitter
synthesis [29,30].

233 Total acylcarnitines in central positions of the heart were decreased by strain Sylvio 234 X10/4 infection compared to the uninfected group (Fig 3 A and B, Mann-Whitney p<0.05). A 235 similar trend was observed for total acylcarnitines following strain CL infection when compared 236 to matched uninfected samples, even though this difference was not statistically significant (Fig 237 3A and B). Previous studies demonstrated that acylcarnitines of different lengths were associated 238 with infection outcome in acute T. cruzi mouse models [8]. Therefore, we sought to understand 239 how different length acylcarnitines were affected by chronic infection. Acylcarnitines are 240 classified based on the number of carbons in their fatty acid chain as short- (<C4), mid- (C5 -241 C11), and long-chain (\geq C12) acylcarnitines.

Central and apical positions (positions B, C and D) had the largest abundance of mid and
long chain acylcarnitines in both CL and Sylvio X10/4 strain compared to the heart base (Mann
Whitney p<0.05) (Fig 3 C-H). In the case of CL strain infection, when compared to uninfected
samples, short chain acylcarnitines were significantly decreased at the heart base and apex
(positions A and D, p<0.05 Mann-Whitney)(Fig 3 C and D). Mid-chain acylcarnitine levels

- 247 were decreased by strain CL infection compared to uninfected samples at central positions
- 248 (position B, p<0.05 Mann-Whitney) (Fig E and F). In contrast, long chain acylcarnitines were
- significantly increased at central and apical positions (positions C and D respectively) by strain
- 250 CL infection (Mann-Whitney p<0.05) (Fig 3 G and H). Strain Sylvio X10/4 infection
- significantly decreased mid chain acylcarnitine at all positions (Mann-Whitney p<0.05)(Fig 3 E
- 252 and F).

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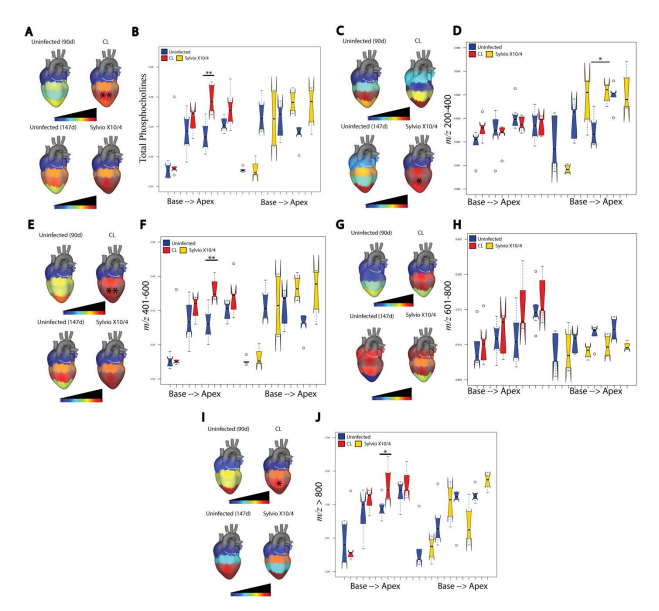
253

254 Fig 3. Spatial impact of chronic *T. cruzi* infection on cardiac acylcarnitines.

255(A) and (B) Differential total acylcarnitine distribution between uninfected and infected heart256sections for both CL and Sylvio X10/4 strains. (*, p<0.05 by Wilcoxon-Mann-Whitney). (C, D)</td>257CL-infected mice showed statistically significant decreases (*, p<0.05 by Mann-Whitney test) in</td>258short-chain acylcarnitine (\leq C4) at heart positions A and D. (E, F) Both strains of *T. cruzi*259showed statistically significant decreases in mid-chain acylcarnitines. (G, H) CL-infected mice260increased long-chain acylcarnitines (\geq C12) at position C (**, p<0.01) and D (*, p<0.05).</td>261

CL strain infection significantly increased total phosphocholines at central position C compared to uninfected samples (Mann-Whitney p<0.01), with a similar but non-significant trend for strain Sylvio X10/4 infection at the heart apex (Fig 4 A and B). Further analysis based on phosphocholine mass was performed, because previous studies showed differences in

- 266 phosphocholine mass range between fatal and non-fatal acute mouse infection [8].
- 267 Phosphocholines were categorized into four mass ranges: short (200 400 m/z), mid (401 600
- 268 m/z), long (601 800 m/z), and very long (>801 m/z). Significantly elevated short
- 269 phosphocholines were observed in central heart positions (position C) for Sylvio X10/4 infection
- 270 compared to uninfected samples (Fig 4 C and D). This same pattern was also observed for CL
- strain infection in mid and very long phosphocholines at the same position (p < 0.01 and < 0.05,
- 272 respectively), when compared to uninfected samples (Fig 4 E and F, I and J).





274 Fig 4. Spatial impact of chronic *T. cruzi* infection on cardiac phosphocholines.

275 (A, B) Statistically significant differences in total phosphocholine levels were identified in mice

276 infected with CL strain at heart position C when compared to uninfected mice (**, p<0.01 by

- 277 Wilcoxon-Mann-Whitney test). (C,D) Sylvio X10/4-infected mice showed statistically
- 278 significant differences (*, p<0.05 by Wilcoxon-Mann-Whitney test) in small phosphocholines
- 279 (200-400 m/z) at heart position C. (E, F) Only CL-infected mice showed statistically significant
- 280 differences (**, p<0.01 by Wilcoxon-Mann-Whitney test) for mid-sized phosphocholines (401-

281	600 m/z) at position C. (G, H) Large phosphocholines (601-800 m/z) were not affected by
282	infection for both strains. (I, J) CL-infected mice showed a statistically significant difference (*,
283	p<0.05 by Wilcoxon-Mann-Whitney test) in very long phosphocholines (> 801 m/z) at position
284	C.

285 **Discussion**

286 Currently, there are 7 T. cruzi discrete typing units (DTUs TcI - TcVI and Tcbat). TcI to TcVI are infectious to humans [31]. These DTUs, while still currently considered the same 287 species, nevertheless present significant genetic differences [31][32]. However, pathogenic 288 289 processes are overall similar in cardiac CD across T. cruzi strains, with accumulation of fibrosis 290 and inflammation, although timing and magnitude of symptoms may be different depending on 291 parasite and host characteristics [32][33]. These similarities are reflected in the common 292 metabolomic changes observed for strain Sylvio X10/4 (TcI) and strain CL (TcVI)-infected heart 293 tissue in this study, including chronic infection-induced increases in phosphocholines and 294 decreases in acylcarnitines.

295 Our results also highlight the importance of considering metabolic changes at the level of 296 chemical families, beyond just individual metabolites. While there was little overlap of highly 297 significant metabolite m/z at each position between strains, most differential metabolites were 298 from these two chemical families. McCall et al. described these two chemical families as 299 discriminatory compounds between fatal and non-fatal acute T. cruzi infected heart tissue [8]. 300 Considering acute stage infection progresses into chronic stage infection, it is not surprising that 301 changes in the relative abundance of these molecules are also observed in chronic CD. 302 Phosphocholines have been linked to coronary heart disease due to production of 303 lysophosphatidylcholines and choline. [29,34]. Increased acylcarnitine levels have been linked to

304 cardiovascular disease as well as cardiac symptoms in those already possessing a cardiac disease 305 [35,36]. However, our results show the opposite pattern compared to non-infectious heart 306 disease, highlighting the need to specifically study CD rather than extrapolate from other cardiac 307 conditions (Fig 3). In a study addressing gene expression differences between human CD 308 cardiomyopathy and dilated cardiomyopathy, there was an upregulation of gene expression 309 associated with lipid metabolism from heart samples of human cardiac CD patients, while the 310 opposite was seen in non-infectious dilated cardiomyopathy patient samples [37]. Higher lipid 311 metabolism would increase acylcarnitine catabolism and thus decrease overall acylcarnitine 312 abundance. Decreased carnitine palmitoyltransferase and acetyltransferase levels, as observed by 313 proteomic analysis of infected mouse heart tissue [38], may alternatively also contribute to the 314 decreased acylcarnitine levels we observed.

315 Interestingly, in a study on the effects of diet on chronic T. cruzi mouse infection, a 316 similar pattern was observed as in our study, where serum acylcarnitines were amongst the most 317 differential compounds in infected samples compared to uninfected samples, with most short-318 and mid-chain acylcarnitines decreased, and select long-chain acylcarnitines increased [17]. In 319 addition, significant acylcarnitine differences were seen in the gastrointestinal tract of acute T. 320 cruzi-infected mice [18]. Likewise, both long chain acylcarnitines and phospholipid synthesis 321 were increased in the heart tissue of acutely infected mice in prior studies [14]. Thus, our data 322 agree with and expand upon the existing T. cruzi metabolomics literature. Overall, the different 323 patterns in metabolites we observe here in contrast to other cardiac diseases is consistent with 324 differences in gene expression in humans with CD compared to other diseases.

325 Differences in pathogenesis between strains may be due to differential strain tropism.
326 Indeed, TcI strains tend to produce cardiomyopathy, while TcVI strains commonly produce

327 megacolon and megaesophagus, although cardiomyopathy can still occur [39]. Our results 328 indicate a disconnect between sites of highest parasite burden and sites of metabolic perturbation. 329 Although parasite levels were highest in central heart segments following strain Sylvio X10/4330 infection, we observed statistically significant perturbations in metabolism at the apex of the 331 heart (Fig 1). Apical aneurysms are one of the major symptoms in chronic CD patients [40]. In 332 addition, lateral heart wall damage is also common among chronic CD patients, in central 333 regions of the heart [41], and we observed significant perturbations in cardiac metabolism at 334 lower central heart positions in strain CL infection (Fig 1B) [41]. Based on these results, we 335 propose a concept of spatial disease tolerance, whereby some tissue regions are more affected by infection, while others are less functionally affected. This is likely due to a combination of host 336 337 and pathogen factors, given the differences we observe here between strain CL and strain Sylvio 338 X10/4 infection in the same C3H mouse genetic background. Importantly, the localization of 339 maximal metabolic perturbation in acute strain Sylvio X10/4 infection was also the heart apex, 340 indicating that the spatial course of disease may be set early in CD [8]. Likewise, host factors 341 likely contribute, such as the higher production of antiparasitic but tissue-damaging IFNy at the 342 heart apex or specific cardiac regions being more prone to microvasculature disruptions [8]. 343 These results set a foundation for biomarker studies and for host-directed therapeutic 344 development. CD may be particularly amenable to such treatment strategies, due to the 345 contribution of host-mediated tissue damage to CD pathogenesis [1,7]. Indeed, we have 346 previously shown that carnitine supplementation can be used to treat acute CD [18]. Our 347 observation of decreases in cardiac acylcarnitines in chronic CD indicate that this approach may 348 also be useful to treat chronic CD. Importantly the fact that acylcarnitines are affected in both

349 chronic CL and Sylvio X10/4 infection suggests broad applicability. Other studies have

350 emphasized the impact of metabolism modulators on CD progression. High fat diet reduces 351 parasite levels and increases survival in acute CD mouse models [42]. Treatment of acutely T. 352 *cruzi* infected mice with metformin (a metabolic modulator used to treat diabetic patients) also 353 led to an increase in overall survival rate and decreased p blood parasitemia [43]. 354 In addition to the need for novel treatments, several studies have highlighted the 355 importance of novel diagnostic methods for CD [44,45]. Current diagnostic methods rely on 356 serological and microscopic exams and polymerase chain reaction (PCR) [46]. In addition, 357 during the chronic stage, parasite levels decrease drastically therefore PCR techniques have to be 358 used instead of microscopy. PCR however only detects the presence of infection but not cardiac 359 damage [46]. Hence, biomarkers in the form of small molecules or chemical families, as 360 identified in this study, can aid in addressing this issue. Future work will investigate whether the 361 infection-induced perturbations observed here in the heart are also detectable in clinically-362 accessible biofluids.

363 Due to the low parasite burden in chronic Chagas disease and instrumental limits of 364 detection, we anticipate most if not all detected metabolites to be host-derived, supported by their 365 detection in uninfected tissues. As such, this study is focused on the impact of T. cruzi infection 366 on host metabolism. A further limitation is that many of the differential metabolites were not 367 annotatable, as is usual in metabolomic studies [47]. Nevertheless, we were able to annotate 368 metabolites affected by chronic infection that make up important host biochemical pathways. 369 Overall, our study highlights the importance of not only identifying overall differences 370 but also positional metabolic differences associated with multiple T. cruzi strains, and the 371 strength of systematic chemical cartography in understanding disease tropism and how it differs

372	from pathogen tropism. These results will serve as stepping stones for further CD drug
373	development and biomarker discovery, something that is urgently needed.
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383	or preparation of the manuscript.

384 **References**

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505 Supporting information

506	S1 Table. Annotated metabolites of combined extracts perturbed by infection at
507	position A, identified through random forest classifier.
508	S2 Table. Annotated metabolites of combined extracts perturbed by infection at
509	position B, identified through random forest classifier.
510	S3 Table. Annotated metabolites of combined extracts perturbed by infection at
511	position C, identified through random forest classifier.
512	S4 Table. Annotated metabolites of combined extracts perturbed by infection at
513	position D, identified through random forest classifier.
514	S5 Table. Annotated metabolites of combined extracts perturbed by infection at
515	positions A-D, identified through random forest classifier.
516	S6 Table. Annotated metabolites of combined extracts identified as perturbed by
517	infection at all positions (FDR-corrected Mann Whitney p<0.05).
518	S1 Figure. Principal coordinate analysis plot of <i>T. cruzi</i> strain CL infected (red) and
519	uninfected (blue) heart tissue samples. Statistically different clustering found in position C
520	(PERMANOVA p-value<0.05).
521	S2 Figure. Principal coordinate analysis plot of T. cruzi strain Sylvio X10/4 infected
522	(gold) and uninfected (blue) heart tissue samples. Statistically different clustering found in
523	position D (PERMANOVA p-value<0.05).

524	S3 Figure. Sub-molecular networks and mirror plot of aqueous and organic extract
525	acylcarnitines and phosphocholines. Each pie chart is one metabolite colored by MS2 spectral
526	count in CL-infected and Sylvio X10/4-infected samples where red is CL and gold is Sylvio
527	X10/4. (A) Subnetwork of aqueous extract acylcarnitines with representative acylcarnitine mirror
528	plot (acetylcarnitine, m/z -204.124). (B) Subnetwork of aqueous extract phosphocholines with
529	representative phosphocholine mirror plot (Spectral match to 1-Hexadecanoyl-2-(9Z-
530	octadecenoyl)-sn-glycero-3-phosphocholine, m/z 758.65). (C) Subnetwork of organic extract
531	acylcarnitines with representative acylcarnitine mirror plot (acetylcarnitine, m/z - 204.126). (D)
532	Subnetwork of organic extract phosphocholines with representative phosphocholine mirror plot
533	(Spectral Match to 1-Oleoyl-2-palmitoyl-sn-glycero-3-phosphocholine, m/z -760.601).
534	S4 Figure. GNPS mirror plots of annotated metabolites. (A) mirror plot of m/z
535	703.575, RT 286s (top, black) to reference library spectrum (SM(d18:1/16:0), bottom, green).
536	(B) mirror plot of m/z 454.294, RT 206s (top, black) to reference library spectrum
537	(hexadecanoyl-lysophosphatidylethanolamine, bottom, green). (C) mirror plot of m/z 377.146,
538	RT 137s (top, black) to reference library spectrum (riboflavin, bottom, green). (D) mirror plot of
539	m/z 646.614, RT 417s (top, black) to reference library spectrum (ceramide, bottom, green). (E)
540	mirror plot of m/z 716.523, RT 395s (top, black) to reference library spectrum (1-palmitoyl-2-
541	oleoyl-sn-glycero-3-phosphoethanolamine, bottom, green).