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Metabolic changes of the host-pathogen environment in a Cryptosporidium infection — Source link

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Published on: 04 Jun 2017 - bioRxiv (American Society for Microbiology)

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- Key words: *Cryptosporidium*, NMR, metabolites, taurine, mitochondria, metabolic
 scavenging
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33 Abstract

Cryptosporidium is an important gut microbe whose contributions towards infant and 34 immunocompromise patient mortality rates are steadily increasing. Current techniques for 35 diagnosing, curing or simply understanding the biology of the parasite are few and far between, 36 relying on a combination of *in-silico* predictions modelled on a varied and unique group of 37 organisms and medical reports. The development of an *in-vitro* culture system, using COLO-38 39 680N cells, has provided the Cryptosporidium community with the opportunity to expand its toolkit for investigating this disease. One area in particular that is sorely overlooked is the 40 metabolic alterations upon infection. Existing research is extremely limited and has already 41 shown that significant variation can be found between the metabolome of different infected 42 host species. Using a ¹H Nuclear Magnetic Resonance approach to metabolomics, we have 43 explored the nature of the mouse gut metabolome as well as providing the first insight into the 44 metabolome of an infected cell line. Through a combination of Partial Least Squares 45 46 Discriminant Analysis and predictive modelling, we exhibit new and potentially game changing insights into the effects of a *Cryptosporidium parvum* infection, while verifying the 47 presence of known metabolic changes. Of particular note is the potential contribution of host 48 49 derived taurine to the diuretic aspects of the disease previously attributed to a solely parasite based alteration of the gut environment. This practical and informative approach can spearhead 50 51 our understanding of the Cryptosporidium-host metabolic exchange and thus provide novel 52 targets for tackling this deadly parasite.

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55 Introduction

Cryptosporidiosis is a disease characterised by prolonged episodes of intense diarrhoea and is 56 the second largest cause of diarrhoeal disease and death in infants across Africa and South Asia 57 58 (Checkley et al. 2014, Kotloff et al. 2013, Striepen 2013, Wanyiri et al. 2014). It is also amongst one of the highest medically important diseases of the immunocompromised, especially HIV 59 positive patients who are at 75-100% risk of contracting the disease depending on the 60 geographical area (Shirley et al. 2012, Wanyiri et al 2014). The pathogens responsible are 61 parasites belonging to the apicomplexans, the Cryptosporidium species, of which C. parvum is 62 typically the more likely (Caccio 2005, Leoni et al. 2006, Widmer and Sullivan 2012, Wielinga 63 et al. 2008). Infection occurs when an individual ingests the oocysts of the parasite, often 64 swallowing a contaminated water source. Water treatment options are limited to filtering, 65 which is generally not possible at an industrial scale and UV treatment, which is both expensive 66 and rarely available prior to the outbreak. Failing this, treatment is typically rehydration, 67 although one drug has been shown to be effective, the broad spectrum anti-parasitic 68 Nitazoxanide (Doumbo et al. 1997). However, the drug is far from ideal and displays a range 69 of undesirable side effects including cytotoxicity and nausea, as well as being limited to use in 70 cases where the patients are immunocompetent (Domjahn et al. 2014, Hussien et al. 2013, 71 72 Manjunatha et al. 2016, Sparks et al. 2015).

Until recently, a significant barrier to research into cryptosporidiosis has been the absence of a combined long-term *in vivo* culturing system and comprehensive model of host parasite interactions in addition to a heavy reliance on antibody based detection both in the scientific and the medical field (Briggs et al. 2014, Checkley et al. 2014, Domjahn et al. 2014, Girouard et al. 2006, Karanis and Aldeyarbi 2011, Leitch and He 2012, Muller and Hemphill 2013, Striepen 2013). Recent papers have attempted to rectify this by proposing improved or entirely novel techniques for culturing the parasite *ex-vivo* in tissue cultures, using the cultured cancer

cells as host cells (Morada et al. 2016, Muller and Hemphill 2013). A recent study identified 80 that infections of COLO-680N cell cultures produced a longer term and higher production 81 volume culture of the parasite compared to previously existing *in-vitro* cultures (Miller et al. 82 2017). These advances have allowed higher in depth microscopy-based studies and even 83 promise to provide a solution to developing a genetic engineering platform for the parasite. 84 However, beyond microscopy and localisation studies, the knowledgebase of the host parasite 85 86 interaction remains largely undeveloped (Manjunatha, et al. 2015, Sponseller et al. 2014, Striepen 2013, Wilhelm and Yarovinsky 2014). 87

One area lacking study is metabolomics. Only two peer-reviewed publications have explored 88 the concept of the infection metabolome, one on mice and the other on human faecal samples, 89 both showing a clear relation between infection and change in metabolite levels (Ng Hublin et 90 al. 2013, Ng Hublin et al. 2012). While working on different sample sources, each identified 91 the hexadecanoic acid as a significant contributor to the change in the metabolome during 92 93 infection. Previous studies noticed a number of metabolites, mainly amino acids, decreased in relative abundance in infected mice faeces compared to an increase seen previously in humans 94 (Ng Hublin et al. 2012). This was explained to be most likely due to the inherent variation 95 between the different host species metabolomes, as highlighted by Saric et al. in 2008 and 96 97 highlights a pressing need for further and wider reaching studies into the metabolome of *Cryptosporidium* infections as well as the development and application of different techniques 98 beyond the Gas Chromatography Mass Spectrometry (GC-MS) used in those papers (Ng 99 Hublin et al. 2013, 2012, Saric et al. 2008). 100

¹H Nuclear Magnetic Resonance (NMR) metabolomics is a powerful alternative to GC-MS for 101 metabolic screening. ¹H NMR is a simpler method that allows for a comparatively lossless 102 analysis of metabolites, with fewer steps between sample recovery and analysis (Bezabeh et al. 103 2009, Hong et al. 2010, Jacobs et al. 2008, Saric, et al. 2008, Wu et al. 2010). This translates 104 to a more reliable result in terms of quantification and reproducibility. As such, NMR has 105 already seen use in analysing the profile of *Plasmodium falciparum*, although the metabolome 106 of the apicomplexan parasite as a whole is almost entirely unexplored (Sengupta et al. 2016). 107 Here, we show a novel method of analysing cryptosporidiosis-induced changes in infected mice 108 guts metabolomes, using a ¹H NMR approach. In addition, we have applied the same NMR 109 based methodology to the *in vitro* infected COLO-680N cell cultures, in order to explore the 110 similarities and differences displayed between *in-vivo* and *in-vitro* models and identify 111 potential cross-species markers of infection. 112

113

114 Materials and Methods

115 Cryptosporidium

Three isolates of *C. parvum* were used in this study. The reference strain *C. parvum* Iowa II was obtained from Bunch Grass Farm in the United States, isolated from infected calves. The human isolate *C. parvum* Weru strain was supplied courtesy of Dr Martin Kváč of the Institute of Parasitology Biology Centre CAS, Czech Republic. The Weru strain was originally isolated from an infected human patient and subsequently maintained by passing through SCID mice. The final isolate used was the human isolate of *C. hominis*, supplied courtesy of Prof. Rachel

122 Chalmers from the *Cryptosporidium* Reference Unit, Singleton Hospital of NHS Wales.

123 *Tissue culture*

124 75 cm^2 monolayers of COLO-680N were infected and maintained as per the protocols outlined 125 in Miller *et al.* 2016, using all three isolates of *Cryptosporidium*. A control group was also 126 established, following the same protocols as the infections, absent oocvsts.

127 Animals and infection

For this study, seven day old BALB/c mice were infected at the Institute of Parasitology, 128 Biology Centre CAS using pre-established protocols detailed in Meloni and Thompson, 129 totalling three mice per condition (Meloni and Thompson 1996). Three separate groups were 130 131 used, one infected with 100,000 oocysts of C. parvum Iowa II, another group was infected with 100,000 oocysts of the *C. parvum* Weru isolate and the final group were given a PBS control. 132 The groups were kept physically spate and never allowed to interact. Infection was monitored 133 from Day-1 post-infection by aniline-carbol-methyl violet staining of faecal smears staining 134 of faecal smears, in addition to an antigen based strip test (Milacek and Vitovec 1985), 135 RIDA®QUICK Cryptosporidium, supplied by R-Biopharm. At ten days post-infection, the 136 mice were euthanized by cervical dislocation and decapitation. This study was carried out in 137 accordance with Act No 246/1992 Coll. of the Czech Republic. The protocol was approved by 138 the Committee for Animal Welfare of Biology Centre Czech Academy of Science and the 139 veterinary administration authorities with regards to the animal experiments. 140

141 *Sample preparation for NMR*

Animal samples were retrieved from the contents of the ileum and surrounding intestinal structure by dissecting out the area of interest and washing through with three ml 100% ethanol at room temperature via syringe inserted into the opening, collecting the wash through.

145 Collected samples were then centrifuged for three minutes at 10,000 g, the supernatant 146 discarded and the pellet weights recorded. The samples were then suspended by vortex in two 147 ml of 75 % ethanol then transferred to a new tube and an additional five ml of 75% ethanol 148 added.

149 Two ml of two mm diameter glass beads were added to the samples and agitated by vortex for

30 seconds before incubating the samples for three minutes at 80°C. The samples were vortexed

for a further 30 seconds or until the sample was completely homogenised. Tissue culture

152 samples were collected by draining the media, adding six ml of ethanol at 80°C to the culture 153 flask and scraping the cells off the surface by cell scraper, decanting the mixture of lysed cells

- 153 into 15 ml polyethylene tubes.
 - 155 The samples were then decanted into two ml tubes, retaining the glass beads in the falcon tubes.
 - The beads were washed with an additional two ml of 80°C, 75% ethanol and again the liquid
 - 157 was decanted into sterile two ml tubes, retaining the glass beads in the tube.

Cell debris and general detritus were removed from the samples by centrifugation at 16,000 g 158 for 10 minutes and the supernatant transferred to new, sterile two ml microcentrifuge tubes. 159 The samples were then dried via Rotorvac overnight at 40°C, suspended in 330 µl double 160 distilled water and centrifuged at 2,500 g for 10 minutes. The supernatant for the samples were 161 recombined into a single 1.5 ml microcentrifuge tube per original samples and frozen at -20 °C 162 until the day before NMR analysis. Twenty-four hours prior to analysis, the sample tubes were 163 placed into a freeze drier until completely desiccated. For NMR analysis, the samples were 164 suspended in one ml of deuterated water and spiked with the sodium salt of the calibration 165 control compound 3-(Trimethylsilyl)-1-propanesulfonic acid (DSS) to a final concentration of 166 20 mM and a tested pH of 7.5. 167

168 *NMR protocol and analysis*

Samples were analysed using a 4-channel Bruker Avance III 14.1 T NMR spectrometer (600 169 MHz¹H) equipped with a 5 mm QCI-F cryoprobe. For controls: six separate, uninfected 25 170 cm² COLO-680N 100% confluent monolayer cultures were analysed in addition to three 171 uninfected BALB/c mice. Infected samples consisted of six 25 cm² COLO-680N 100% 172 confluent monolayers in addition to three Iowa infected BALB/c and three Weru infected 173 BALB/c mice. One dimension NMR datasets were acquired with a pulse repetition rate of 5 s 174 175 over 128 scans, preceded by eight equilibrating dummy scans and suppression of the residual Deuterium Oxide solvent (HDO) resonance using presaturation. Processed NMR 176 spectrographic datasets were produced by Topspin 3.2 and analysed using Chenomx NMR 177 178 Suite version 8.2. Partial Least Squares Discriminant Analysis (PLS-DA) of the Chenomx data were generated with the freely available Microsoft Excel Add-in "multi-base 2015" by 179 Numerical Dynamics, Japan (Mutlibase for Microsoft Excel 2015). Pathway predictions were 180 181 produced by the MetaboAnalyst 3.0 web tool, using a hypergeometric test and relativebetweeness centrality against Homo sapiens and Mus musculus databases for the tissue culture 182 and mouse models respectively (Xia et al. 2015). 183

184 Indirect Fluorescence Assays

COLO-680N cultures were seeded onto Lab-Tek, two well, Permanox chamber slides (Sigma 185 Aldrich, Cat No. Z734640) and allowed to reach 70% confluence before infecting, following 186 previously published protocols (Miller et al. 2017). At seven days post infection the media was 187 aspirated from the cultures and washed twice with 1 x PBS. Fresh, pre-warmed RPMI-1640 188 189 (Sigma Aldrich, Cat. No R8758) (1% Antibiotic/Antimycotic) containing 200 nM Thermofisher Mitotracker Red CMXRos (Molecular probes; Cat. No M7512), was added to 190 191 the wells and incubated in the dark at 37°C for 45 minutes. The media was removed and 192 replaced with further pre-warmed RPMI-1640 (1% Antibiotic/Antimycotic), containing 3.5% formaldehyde, for 15 minutes at 37°C as per the manufacturer's protocol. The cells were then 193 briefly permeabilised with 0.2% Triton-x100 in 1x PBS for 10 minutes, washed twice with 1x 194 PBS and four drops of SporoGlo[™] or Crypt-a-glo[™] (WATERBORNE, INC) added, with 195 incubation at 37°C for a further 45 minutes. The final sample was then washed three times with 196 PBS, dried and FluoroshieldTM with DAPI (Sigma Aldrich, Cat. No F6057) was added before 197 applying a glass coverslip and sealing. Slides were visualised by fluorescence microscopy 198 using an Olympus IX82 or Zeiss Elyra P1 confocal microscope. 199

200 *Electron microscopy images*

Aclar disks of tissue culture were infected and prepared for EM according to the protocols detailed in Miller et al. 2017.

203 *Ethics*

All animals involved in the experiments were treated and cared for by trained members of staff according to the standards set out by Directive 2010/63/EU regarding Legislation for the protection of animals used for scientific purposes.

- 207
- 208
- 209 **Results**
- 210 Mice faecal sample extractions

Faecal samples from infected and uninfected mice were monitored using aniline-carbol-methyl
violet staining (Figure 1) to determine validity of the control and progress of infection.
Samples from both control and infected mice were taken at ten days post infection.

The spectra produced by the NMR showed clear distinctions between the infected and 214 uninfected mice, as well as distinctions between the different strains of infections (Figure 2a). 215 Several metabolites were readily distinguishable prior to the metabolomics analyses, including 216 indicators of phosphorylation; creatine and creatine phosphate (Figure 2b), taurine (Figure 217 **2c**) and lactate (Figure 2d). Processing the data from the mice guts (n=9) via the Chenomx 218 Nmr Suite version 8.2 platform produced a list of 151 compounds that were extrapolated from 219 the spectra (Figure 3). Statistical analysis of the data, with freely available Microsoft Excel 220 Add-in "multi-base 2015", by Partial Least Squares Discriminant Analysis (PLS-DA) 221 determined significant separation of the three conditions, (uninfected control, C. parvum Iowa 222 II and C. parvum Weru infections), whilst maintaining group cohesion (Figure 4a). The 223 loading values of the variable compound contributions (Figure 4b), suggest certain metabolites 224 were more significant to the separation of the groups than others. The presence of L-alanine 225 and valine, two common amino acids, agrees with the previous literature and 2-oxoisocaproate 226 is a component of the valine/leucine/isoleucine biosynthetic pathways reports (Ng Hublin et al. 227 2013, Ng Hublin et al. 2012). 228

MetaboAnalyst 3.0 based analysis of the metabolites proposed that a number of amino acid 229 biosynthesis pathways could be altered during the course of an infection, such as the glycine, 230 231 valine and taurine pathways. In addition, the mice infections displayed possible changes to other metabolic pathways (Figure 5a) as those pathways furthest from the x, y axis intercept, 232 233 representing both the overall completeness of the pathways and number of contributing 234 detected metabolites respectively. The pathways identified in the manner, and the compounds discovered by the NMR demonstrated that infections caused changes in at least the valine 235 (Figure 5c), glycine (Figure 5d) and taurine amino (Figure 5e) acid biosynthetic pathways, in 236 addition to several sugar pathways (Figure 5b, f, g). 237

238 *Cell culture sample extractions*

Extrapolated NMR data from COLO-680N (n=18) metabolite extractions, demonstrated clear 239 differences between the each strain and species of *Cryptosporidium* used (Figure 6). As with 240 241 the mice samples, differences between creatine, creatine phosphate, taurine and lactate (Figures 6b-d) were readily visible in the raw spectra. Chenomx analysis produced a list of 242 161 total compounds of varying concentrations across samples (Figure 7). The PLS-DA 243 generated by the same statistical analysis as before, produced ample separation of the 244 Cryptosporidium-infected and uninfected cultures, (Figure 8a). Furthermore, the separation of 245 the individual infection groups suggests that differences between both Cryptosporidium species 246 247 and within individual strains of C. parvum, may illicit different metabolic responses in cell cultures. The loading scores plot of the PLS-DA showed a number of amino acids contributed 248 heavily to the separations, as well lactate, several fatty acid derivatives and taurine (Figure 249 250 **8b**).

Metabolic pathway fitting via MetaboAnalyst 3.0 revealed that amino-acid biosynthesis pathways for glycine, alanine and arginine were influenced by infection. These were in addition to taurine, pantothenate and CoA biosynthetic pathways as shown in **Figure 9a**. As with **Figures 5a-g**, the graph shows a combination of how much of a pathway is completed by data from the NMR, as well as simply how many metabolites were detected. Among other pathways, perhaps the most significant detections were glycine (**Figure 9b**), taurine (**Figure 9c**) alanine (Figure 9d) and arginine (Figure 9g) amino acid pathways as well as, potentially the synthesis
 and degradation of ketones (Figure 9e) and pantothenate and CoA biosynthesis (Figure 9f).

259 *Comparison of mice faecal and COLO-680N metabolome changes*

MetaboAnalyst data from Figure 5 and Figure 9, demonstrate that a number of altered 260 pathways are shared between the mice and tissue culture metabolites, particularly taurine and 261 amino acid metabolic pathways. Taurine is involved in a number of roles, including bile acid 262 conjugation, osmoregulation, membrane integrity and protection against oxidative free 263 264 radicals. Glycine synthesis was also shown to be affected to a large degree and is involved with numerous and diverse cellular functions including purine synthesis, basic protein construction 265 and provides the building blocks for porphyrins (Denis and Daignan-Fornier 1998, Marver et 266 al. 1966). All of these pathways have a direct or indirect impact on the host's mitochondrial 267 energetic activity. 268

To investigate the cellular role of host mitochondria during infection, we employed an Indirect Fluorescence Assay (IFA) approach to determine if the mitochondria of the host cells were responding to *Cryptosporidium* infection (Figure 10). Our results demonstrate that on multiple occasions, the host mitochondria were shown to congregate in larger densities near the *Cryptosporidium* infection (Figures 10). Transmission Electron Microscopy images of infected cells also show abnormal host's mitochondrial congregation around the parasitophorous vacuole (Figure 11).

276

277 Discussion

Solution-state ¹H NMR offers a novel approach to metabolomics that is especially useful where 278 sample volume sizes are particularly small (Jacobs et al 2008, Novak et al. 2006, Wu, et al. 279 2010). Although GC-MS holds an advantage for detecting low-levels of metabolites with 280 unique mass signatures, for the purpose of determining the change in metabolite quantities, 281 NMR provides a viable alternative (Bezabeh et al. 2010, Jacobs et al 2008, Saric et al. 2008, 282 Sengupta et al. 2016, Wu et al 2010). Initial analysis of our data showed a clear distinction 283 between the metabolic fingerprints of infected and uninfected samples, even between infections 284 285 of different strains of the parasite (Figure 4).

Of particular importance is the degree to which these results, both from the *in-vitro* and *in-vivo*, agree with the previous literature. Our study also demonstrates that metabolic compounds Lalanine, isoleucine and succinic acid (succinate) were detected as contributors to the variance between the sample conditions that indicated infection. Moreover, even though valine was not detected in the uninfected controls, it was visible in the infected samples and in agreement with previous reports (Ng Hublin et al. 2013, Ng Hublin et al. 2012).

The predictive fits of the metabolic pathways highlighted a remarkable selection of pathways including several involved in amino acid biosynthesis, sugar metabolism, CoA biosynthesis and taurine biosynthesis. Of the predicted metabolic pathways, which have been previously shown to be influenced by infection, there are several whose presence should have been projected, such as the amino acid biosynthesis pathways for alanine and glycine as the previous reports had already highlighted their potential involvement reports (Ng Hublin et al. 2013, Ng Hublin et al. 2012).

As a parasite, *Cryptosporidium* is dependent on host derived biosynthetic pathways for survival. For example, *C. parvum* is incapable of producing the majority of amino acids *denovo*, instead relying heavily on the import of host metabolites via active channelling

(Abrahamsen et al. 2004). The biosynthetic pathway for glycine, threonine and serine was 302 upregulated, in both cell culture and animal experimentations, with particularly high levels of 303 glycine detected. Both C. parvum and C. hominis are incapable of manufacturing these amino 304 acids de novo, instead relying on scavenging host serine and glycine, utilising serine and 305 glycine hydroxymethyltransferases to convert one to the other when needed (Abrahamsen et 306 al. 2004, Doyle et al. 1998). The reliance on host amino acids could provide a novel method 307 308 for combating the infection, based upon previous studies that identified other amino acid metabolic chains as potential targets (Clark 1999, Doyle et al. 1998). For example, glycine 309 reuptake inhibitors (GRIs) that are often used in treating schizophrenia, could be utilised to 310 311 partially starve the parasite of the metabolite.

In addition to the amino acid biosynthesis pathways, it is also apparent that taurine synthesis is 312 also implicated in the metabolic profile of the disease as shown in the presented analyses; 313 taurine has frequently been used in the past as an agent for inducing excystation for *in-vitro* 314 cultures as sodium taurochloate (Feng et al. 2006, Gold et al. 2001, Kar et al. 2011, King et al. 315 2012). In the host, taurine has a number of roles, those relevant to the cell types involved 316 include: cell membrane integrity, osmoregulation and adipose tissue regulation. Previous 317 metabolomic studies of faecal samples from Cryptosporidium-infected patients revealed 318 increased taurine concentrations, explained by the characteristic decline in gut absorption as a 319 result of villi malformation by the parasite (Goodgame et al. 1995, Kapembwa et al. 1990). 320 However, an even greater increase in taurine levels was observed in the infected COLO-680N 321 cell cultures, wherein malabsorption is not an applicable explanation. In addition to the 322 pathways and the relevant metabolites featured in Figures 5 and 9, there were also a number 323 324 of potentially important metabolites not represented. Similarly observed, was an increase in the abundance of adenosine derivatives (AMP, ADP and ATP); all showed increased abundance 325 in infected cells and mice in C. parvum Iowa II infections, along with a similar increase in 326 creatine levels in C. parvum Weru infections. This heavily implicates a role for the host 327 mitochondria in the context of infection as each species and strain used lacks the creatine kinase 328 needed to produce creatine phosphate, which typically operate in localisation with 329 mitochondria. Levels of pyruvate in C. hominis cell and pantothenate in C. parvum Iowa II 330 infections suggest a role for oxidatative phosphorylation. This is of particular interest as it has 331 been established that the C. parvum genome contains a sequence for a potential pantothenate 332 scavenging protein (Augagneur et al. 2013). Moreover, the further increase in lactate levels 333 detected in C. hominis cell cultures and C. parvum Iowa II mouse infected samples, compared 334 to the controls, indicate a strong contribution from anaerobic pathways most likely from the 335 host. This suggests that more ATP is being produced than the oxidative capacity of the host 336 337 mitochondria can maintain, producing a net increase in lactate as the oxygen debt increases. 338 This holds particular interest as a theory of *Cryptosporidium's* targets of parasitism include ATP production pathways, similarly to the intracellular rhizarian Mikrocytos mackini (Burki et 339 340 al. 2013).

These data suggest that C. parvum and C. hominis infections may be directly or indirectly 341 inducing an increase in host mitochondrial activity. If factual, this would result in a large 342 number of oxygen free radicals being produced by the metabolic machinery. Consequently, 343 cell(s) would respond with a matching increase in the synthesis of antioxidants such as taurine, 344 which also sees increases during infection (Giris et al. 2008, Green et al. 1991, Zhang et al. 345 2004). Support for this hypothesis can be seen in the way host mitochondria appear to 346 347 congregate around the Cryptosporidium infection (e.g. parasitophorous vacuole) (Figures 10 and 11). Nevertheless, taurine also plays another role within cells, for example as a diuretic. 348 Taurine is involved in the maintenance of the ionised forms of magnesium and potassium 349 within the cell, producing a diuretic effect that may contribute towards the characteristic water-350

loss of a patient with cryptosporidiosis (Kapembwa et al. 1990, Lin et al. 2016, Niggli et al. 351 1982, Yu et al. 2016). Furthermore, it has been found that taurine levels influence production 352 of short chained fatty acid, another aspect of host biology theorised to be scavenged by C. 353 parvum and C. hominis (Guo et al. 2016, Seeber and Soldati-Favre 2010, Yu et. 2016). The 354 detection of a rise in taurine levels *in-vitro* further suggest that the increase in taurine typically 355 detected in cryptosporidiosis patients' stool, is more than simply the result of the guts decrease 356 in absorptive qualities. It is likely that the intra-cellular role of taurine in this disease has been 357 overlooked and that the pathophysiology of this disease is more complicated than currently 358 understood, and extends beyond simple villi degradation. 359

Lastly, these results alone provide a promising method of determining infections via a possible comparative ¹H NMR of patient and reference biopsies. This method offers an alternative approach in the medical field, where current methods of diagnosis are reliant on separate methods to achieve the same result as NMR, with infections detected by laborious and often inaccurate microscopy and strain typing dependant on successful PCR.

In conclusion, we have demonstrated for the first time that the use of ¹H NMR in the context 365 of both medical and scientific applications is indispensable in the fight against 366 cryptosporidiosis. With the application of a more user-friendly and reproducible approach of 367 metabolomics, through the ¹H NMR methodology described in this paper, it will now be easier 368 for the *Cryptosporidium* community to further explore the remaining aspects of the disease 369 metabolome in patients' samples. Future experiments would be best approached by increasing 370 371 the number of strains analysed both in-vitro and in-vivo to test the relevant proposed hypotheses. Additionally, elucidating the more pathogenic influences of taurine biosynthesis 372 373 in the pathobiology of cryptosporidiosis is critical. With these data, a metabolomics based 374 method of diagnosing and treating the disease could become a reality.

375

376 Abbreviations

- 377 NMR: Nuclear Magnetic Resonance
- 378 DSS: 3-(Trimethylsilyl)-1-propanesulfonic acid, sodium salt
- 379 PCA: Principal component analysis
- 380 PLS-DA: Partial Least Squares Discriminant Analysis
- 381 UV: Ultraviolet
- 382 HIV: Human Immunodeficiency Virus
- 383 GC-MS: Gas Chromatography-Mass Spectrometry
- 384 HDO: Deuterium Oxide
- 385 IFA: Indirect Fluorescence Assay
- 386 PCR: Polymerase Chain Reaction
- 387 DAPI: 4',6-diamidino-2-phenylindole
- 388 PBS: Phosphate-buffered saline
- 389 EM: Electron microscopy
- 390 SCID: Severe Combined Immunodeficiency Disease

- 391 ATP: adenosine triphosphate
- 392 AMP: adenosine monophosphate
- 393 ADP: adenosine diphosphate
- 394 CoA: Coenzyme A
- 395 GRIs: glycine reuptake inhibitors
- 396

397

398 **Declarations**

The authors have declared that the research was conducted in the absence of any commercialor financial relationships that could be construed as a potential conflict of interest

Funding was provided by the BBSRC, Wellcome Trust and a Microbiology Society ResearchVisit Grant.

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404 Acknowledgements

This research was supported by BBSRC research grant (BB/M009971/1) to Dr. Anastasios 405 Tsaousis and a Wellcome Trust Equipment Grant 091163/Z/10/Z to Dr. Mark J. Howard. 406 Christopher N. Miller is supported by a GTA studentship from the School of Biosciences, 407 University of Kent and a Research Visit Grant award from the Microbiology Society. Martin 408 Kvac is supported by the Czech Science Foundation (project No. 15-01090S). We thank Dr. 409 Michelle Rowe for NMR technical support at Kent and members of the Dr. Tsaousis and Dr. 410 Kvac laboratories for their intellectual and methodological support. We would also like to 411 thank Matthew Lee and Matthew D. Badham from the University of Kent for the assistance in 412 using the confocal microscope. 413

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- 569
- 570
- 571 Figure legends

572 Figure 1: Staining of *Cryptosporidium* in faecal samples

- 573 Aniline-carbol-methyl violet stain of a faecal smear taken from a mouse in the infection group.
- 574 The abundant presence of *Cryptosporidium* (arrows) indicates that the infection has been
- 575 successful; and that the animal is producing oocysts.
- 576

577 Figure 2: NMR Spectra of mice models of infection

a. Stacked NMR Spectra produced from faecal samples of the control mice (green), or either the Iowa II (blue) or Weru (purple) groups. **b.** Direct comparisons of the spectra revealed several clearly identifiably differences, including differences in creatine and creatine phosphate
levels. c. Levels of taurine were substantially lower in the control or *C. parvum* Weru samples
compared to *C. parvum* Iowa II. d. Lactate levels were also much higher in *C. parvum* Iowa II
infected mice compared to the barely detectable levels in the control mice or *C. parvum* Weru

584 infected groups.

585

586 Figure 3: Mice Experiment Metabolites

All the metabolites identified by ¹H NMR analysis in infected and uninfected mice were explored via PLS-DA statistical analysis. The colour coded heat map represents the significance to which each individual metabolite contributed to the identity of the sample groups, with more significant contributors tending towards green and less reliable contributors tending towards red.

592

593 Figure 4: PLS-DA and loading plot of mice model NMR results

a. PLS-DA statistical analysis of the information provided by the Chenomx screening produced 594 595 clear groupings, separating the controls (green), C. parvum Iowa II infections (blue) and C. *parvum* Weru infections (purple). As the grouping areas, indicated by the areas highlighted, do 596 no overlap, it can be said that the separation between the infection conditions represent clear 597 differences in the metabolome, which correspond to the C. parvum strain. b. The loading biplot 598 of the PLS-DA analysis shows many of the compounds identified by Chenomx contributed 599 towards the separation and groupings. Those on the outer most edges, for example alanine, 600 sarcosine, lactate and lactulose, had some of the greatest influence on the amount of separation 601 as determined by the PLS-DA. 602

603

Figure 5: Metabolic pathways detected in mouse model NMR samples

a. Data analysed by MetaboAnalyst 3.0, utilising all compounds which displayed some degree 605 of change as a result of infection, produced a graph of pathways most heavily impacted (x axis) 606 and pathways containing the most amount of the given compounds (pathway impact: y-axis), 607 with statistical significance of the predicted pathways increasing as the colour ranges from 608 609 yellow (low) to red (high). Six pathways were chosen to be of particular interest by their position on the graph, with metabolites present in the experimental samples highlighted in red, 610 including: b. pentose and glucoronate interconversions, valine, c. leucine and isoleucine 611 biosynthesis, d. glycine serine and threonine metabolism, e. taurine and hypotaurine 612 613 metabolism, f. galactose metabolism and g. starch and sucrose metabolism.

614

615 Figure 6: Cell Culture infection NMR spectra

a. Stacked NMR Spectra produced from the COLO-680N control cultures (green), or either the *C. parvum* Iowa II (blue), *C. parvum* Weru (purple), or *C. hominis* groups. Direct comparisons of the spectra revealed several clearly identifiably differences, including, again, differences in creatine and creatine phosphate (**b.**), taurine (**c.**) and lactate (**d.**) levels. Noticeably, taurine

620 levels were almost undetectable in *C. hominis* or *C. parvum* Weru infections.

621

622 Figure 7: COLO-680N Experiment Metabolites

All the metabolites identified by 1H NMR analysis in infected and uninfected cells were explored via PLS-DA statistical analysis. The colour coded heat map represents the significance to which each individual metabolite contributed to the identity of the sample groups, with more significant contributors tending towards green and less reliable contributors tending towards red.

628

629 Figure 8: PLS-DA and loading plot of COLO-680N - infected cells NMR results

a. PLS-DA statistical analysis of the information provided by the Chenomx screening produced clear groupings, separating the controls (green), *C. parvum* Iowa II infections (blue), *C. parvum* Weru infections (purple) and *C. hominis* infections (red). As the grouping areas do no overlap the separation between the infection conditions again indicates that metabolome differences can be at least in part explained by different *Cryptosporidium* strains/species. **b.** The loading biplot of the PLS-DA analysis shows lactate as a significant contributor to variation, as seen before in Figure 2b in addition to tauring and myo inositel among others.

- before in **Figure 2b**, in addition to taurine and myo-inositol among others.
- 637

638 Figure 9: Metabolic pathways detected in cell cultures' NMR samples

a. Data analysed by MetaboAnalyst 3.0, utilising all compounds which displayed some degree
of change as a result of infection, produced a graph of pathways most heavily impacted (x axis)
and pathways containing the most amount of the given compounds (pathway impact: y-axis),
with statistical significance of the predicted pathways increasing as the colour ranges from
yellow (low) to red (high). Six pathways were chosen to be of particular interest by their
position on the graph, with metabolites present in the experimental samples highlighted in red,
including: glycine, serine and threonine metabolism (b.), taurine and hypotaurine metabolism

- 646 (c.), Alanine, aspartate and glutamate metabolism (d.), synthesis and degradation of ketones
- 647 (e.), pantothenate and CoA biosynthesis (f.) and arginine and proline metabolism (g.).
- 648

649 Figure 10: Indirect Fluorescence Assay of infected cell cultures

a. Fluorescence microscopy showing the staining of infected COLO-680N culture with Sporo-650 glo (green), MitoTracker CMXRos (red) and DAPI nuclear stain (blue). From the figure we 651 could observe an obvious mitochondrial "clumping" and polarisation towards areas of 652 infection, suggesting that the presence of the parasite within a host cell affects the positioning 653 of host mitochondria. **b**. Confocal microscopy showing the localisation of Crypt-a-glo (green), 654 655 MitoTracker (red) and DAPI (blue) in a 3D rendering of 31 individual, 0.16 µm thick sections, overlapping with a final representative thickness of 4.8 µm. The images are rotated around the 656 x-axis, from 0° to 80°, showing a COLO-680N cell infected with C. parvum (green). Individual 657 images of the stainings were captured in different angles, to show the infection on a three-658 dimensional level. A whole video showing a 360° rotation of the three-dimensional z-stack of 659 the image is found as an animation in Video 1. c. Confocal microscopy showing the localisation 660 of Crypt-a-glo (green), MitoTracker (red) and DAPI (blue) in a 3D rendering of 55 individual, 661 0.16 µm thick sections, overlapping with a final representative thickness of 8.6 µm. The images 662 are rotated around the x-axis, from 0° to 80° , showing a COLO-680N cell infected with C. 663 *parvum* (green). Individual images of the stainings were captured in different angles, to show 664 the infection on a three-dimensional level. A whole video showing a 360° rotation of the three-665

dimensional z-stack of the image is found as an animation in Video 2. d. Confocal microscopy 666 showing the localisation of Crypt-a-glo (green) and MitoTracker (red) in a 3D rendering of 51 667 individual, 0.16 µm thick sections, overlapping with a final representative thickness of 8.0 µm. 668 The images are rotated around the x-axis, from 0° to 70° , showing mitochondria surrounding 669 an intracellular stage of with C. parvum (green). Individual images of the stainings were 670 captured in different angles, to show the infection on a three-dimensional level. A whole video 671 showing a 360° rotation of the three-dimensional z-stack of the image is found as an animation 672 in Video 3. 673

674

675 Figure 11: Electron microscopy of *Cryptosporidum* infected host cells.

a. Infection of a host cell by *C. parvum*. Mitochondria of the host cell appear to closely associate with the parasitophorous vacuole surrounding the parasite, while cytoskeletal structures appear to be associated with the organelles. **b.** Cartoon of image a. demonstrating the presence of mitochondria, cytoskeleton, nuclear material and *Cryptosporidium*.





 PPM

	Mouse Control	Mouse Iowa	Mouse Weru		Mouse Control	Mouse Iowa	Mouse Weru	
1,3-Diaminopropane				Acetoacetate				Dimeth
1,3-Dihydroxyacetone				Acetoin				Ery
1,3-Dimethylurate				Acetone				Eth
1,6-Anhydro-β-D- glucose				Alanine				Ethyle
1,7-Dimethylxanthine				Allantoin				For
2-Aminoadipate				Alloisoleucine				Fu
2-Hydroxy-3- methylvalerate				Anserine				Gala
2-Hvdroxvbutvrate				Arabinitol				Gala
bioRxiv preprint doi: https://doi.org/10.1101/14 not certified by peer review) is the Social Content of the second	5979; this version pos ne author/funder. All ri	sted June 4, 2017. Ti ghts reserved. No re	he copyright holder fo use allowed without p	er this preprint (which was				Gluo
2-Hydroxyisovalerate				Ascorbate				Glu
2-Hydroxyvalerate				Asparagine				Gluo
2-Oxocaproate				Aspartate				Glu
2-Oxoisocaproate				Azelate				Gluc phos
2-Phosphoglycerate				Betaine				Glut
3-Aminoisobutyrate				Butanone				Glyc
3-Hydromuconate				Caffeine				Gly
3-Hydroxy-3-				Caprate				Glv
methylglutarate				Oamitina				Ohio
3-Hydroxyisovalerate				Carnitine				Giyco
3-Hydroxymandelate				Cellobiose				Gly
3-Methyladipate				Chlorogenate				Glycy
3-Methylxanthine				Cholate				Guanidir
3-Phenylpropionate				Choline				Guanid
4-Carboxyglutamate				Citraconate				Home
5,6-Dihydrothymine				Citrate				lbup
5-Aminolevulinate				Creatine				Isoc
5-Hydroxyindole-3-				Creatine				Isole
5-Hydroxylysine				Creatinine				Isopr
Acetamide				Cystathionine				La
Acetaminophen				Cysteine				Lac
Acetate				Cystine				Lac





Leucine

Maltose

Mannitol

N-

NADH

	Mouse Control	Mouse Iowa	Mouse Weru		
Pantothenate				Key:	
Phenylacetate Pimelate					-1.00 -0.90
Pyroglutamate					-0.80
Ribose Sarcosine					-0.70 -0.60
Succinate					-0.50
Succinylacetone					-0.40
Sucrose		X			-0.30
Syringate					-0.20
Tartrate					-0.10
Taurine					0.00
Theophylline					0.10
Threonine					0.20
Thymidine					0.30
Thymine					0.40
Thymol					0.50
Tiglylglycine					0.60
Trehalose					0.70
Trimethylamine					0.80
methylamine N-oxide					0.90
-N-Acetylglucosamine		1			1<
UDP-glucose					
Urea					
Valine					
Xylitol					
Xylose					
dTTP					
myo-Inositol					
sn-Glycero-3-					
phosphocholine					

a.











D

C.





	Uninfected: Control	Infected: Iowa	Infected: Weru	Infected: Hominis	
1,3-Diaminopropane					
1,3-Dihydroxyacetone					
1,3-Dimethylurate					ŀ
1,6-Anhydro-β-D-glucose					
2-Aminoadipate					A
2-Hydroxybutyrate					
2-Hydroxyglutarate					Ace
bioRxiv preprint doi: https://doi.org/10.1101/145979; this version pos 2-Hydrocectige is per review in the two of funder. All rive	sted June 4, 2017. The copyright h ghts reserved. No reuse allowed w	older for this preprint (which was ithout permission.			
2-Hydroxyisocaproate					
2-Hydroxyisovalerate					Al
2-Hydroxyvalerate					
2-Methylglutarate					8
2-Oxobutyrate					
2-Oxoisocaproate					
2-Phenylpropionate					А
3,5-Dibromotyrosine					
3-Hydroxy-3-methylglutarate					
3-Hydroxyisovalerate					
3-Methylxanthine					
3-Phenyllactate					1
4-Aminobutyrate					H
4-Hydroxybutyrate					
4-Hydroxyphenylacetate					C
4-Hydroxyphenyllactate					
4-Pyridoxate					
5,6-Dihydrothymine					Creat
5-Aminolevulinate					(
5-Hydroxylysine					
5-Methoxysalicylate					Dim
ADP					Dir

Uninfected: Infected: Control AMP ATP Acetamide Acetate cetoacetate Acetone cetylsalicylate Alanine Allantoin lloisoleucine Anserine Arabinitol Arabinose Arginine Asparagine Aspartate Azelate Betaine Biotin Butanone Carnitine Choline Citraconate Citrate Creatine atine phosphate Creatinine Cysteine methyl sulfone imethylamine







	Uninfected: Control	Infected: lowa	Infected: Weru	Infected: Hominis		Uninfected: Control	Infected: Iowa	Infected: Weru	Infected: Hominis		Uninfected: Control	Infected: Iowa	Infected: Weru	Infected: Hominis		
ndole-3-lactate					N-Methylhydantoin					Trimethylamine N-oxide					Key:	
Isocitrate					N-Nitrosodimethylamine					Tyrosine						-1
Isoleucine					N-Phenylacetylglycine					UDP-N-Acetylglucosamine						-0.9
Isopropanol					N6-Acetyllysine					UDP-glucuronate						-0.8
Lactate					NAD+					Valine						-0.7
Lactose					Na-Acetyllysine					Xylitol						-0.6
Lactulose					O-Acetylcarnitine					myo-Inositol						-0.5
Leucine					O-Acetylcholine					sn-Glycero-3-phosphocholine						-0.4
Levulinate					O-Phosphocholine					trans-4-Hydroxy-L-proline						-0.3
Malate					O-Phosphoethanolamine					β-Alanine						-0.2
Maleate					Oxypurinol					π-Methylhistidine						-0.1
Malonate					Pantothenate											0
Mandelate					Pyridoxine											0.1
Mannitol					Pyroglutamate											0.2
Methanol					Pyruvate											0.3
Methionine					Ribose											0.4
Methylamine					Sarcosine											0.5
lethylguanidine					Sebacate											0.6
/lethylsuccinate					Succinate											0.7
Dimethylformamide					Succinylacetone											0.8
I-Dimethylglycine					Sucrose											0.9
Acetylaspartate					Syringate											1
-Acetylcysteine					Tartrate											1.1<
cetylglucosamine					Taurine											
Acetylglutamate					Theophylline											
Acetylglutamine					Threonate											
N-Acetylglycine					Threonine											
-Acetylornithine					Thymine											
-Acetylserotonin					Trehalose											
arbamoyl-β-alanine					Trimethylamine											



PC1 (22.6%)







Pathway Impact

Sporo-glo

MitoTracker

