1 Genetic Susceptibility to Enteric Fever in Experimentally Challenged Human

2 Volunteers

- 3 Amber Barton^{*1,2}, Jennifer Hill¹, Sagida Bibi¹, Liye Chen³, Claire Jones¹, Elizabeth Jones¹, Susana
- 4 Camara¹, Sonu Shrestha¹, Celina Jin¹, Malick M Gibani^{1,4}, Hazel Dobinson¹, Claire Waddington^{1,5}, Thomas
- 5 C Darton^{1,6}, Christoph J Blohmke¹, Andrew J Pollard¹

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- 7 *Corresponding author, Amber.Barton@lshtm.ac.uk
- 8 1. Oxford Vaccine Group, Department of Paediatrics, University of Oxford and the NIHR Oxford
- 9 Biomedical Research Centre, Oxford OX3 7LE, UK
- 10 2. Clinical Research Department, London School of Hygiene and Tropical Medicine, Keppel Street,
- 11 London, UK
- 12 3. Nuffield Department of Orthopaedics, Rheumatology and Musculoskeletal Sciences, University of
- 13 Oxford, UK
- 14 4. Department of Infectious Disease, Imperial College London, UK
- 15 5. Department of Medicine, University of Cambridge, Cambridge, UK
- 16 6. Department of Infection, Immunity and Cardiovascular Disease and the Florey Institute for Host-
- 17 Pathogen Interactions, University of Sheffield, Sheffield UK
- 18

19 Abstract

- 20 Background: Infection with Salmonella enterica serovars Typhi and Paratyphi A cause an estimated 14
- 21 million cases of enteric fever annually. Here the controlled nature of challenge studies is exploited to
- 22 identify genetic variants associated with enteric fever susceptibility.
- 23 Methods: Human challenge participants were genotyped by Illumina OmniExpress-24 BeadChip array
- 24 (n=176) and/or transcriptionally profiled by RNA-sequencing (n=178).
- 25 Results: Two SNPs within *CAPN14* and *MIATNB* were identified with p<10⁻⁵ for association with
- 26 development of symptoms or bacteraemia following oral S. Typhi or S. Paratyphi A challenge. Imputation of
- 27 classical human leukocyte antigen (HLA) types from genomic and transcriptomic data identified HLA-
- 28 B*27:05, previously associated with non-typhoidal Salmonella-induced reactive arthritis, as the HLA type
- 29 most strongly associated with enteric fever susceptibility (p=0.012). Genes related to the unfolded protein 1

response and heat shock were over-represented in HLA-B*27:05⁺ participants following challenge (p=0.01).
Furthermore, intracellular replication of *S*. Typhi is higher in C1R cells transfected with HLA-B*27:05
(p=0.02).
Conclusion: These data suggest that activation of the unfolded protein response by HLA-B*27:05
misfolding may create an intracellular environment conducive to *S*. Typhi replication, increasing

35 susceptibility to enteric fever.

36

37 Key words

Typhoid Fever; Single Nucleotide Polymorphism; Transcriptome; Salmonella Typhi; Genomics; Unfolded
Protein Response; HLA Antigens

40

41 Background

42 Salmonella enterica serovars Typhi and Paratyphi A cause an estimated 14 million cases of enteric fever per 43 year, resulting in 135,000 deaths [1]. Several risk factors have been identified for enteric fever, including 44 poor sanitation and flooding [2]. Individual host factors also likely to contribute to disease susceptibility. 45 Human challenge models, where volunteers are deliberately exposed to a pathogen, have been developed to 46 study the biology of enteric fever and test experimental vaccines. Despite ingesting the same inoculation 47 dose of bacteria, some challenged individuals remain infection-free, while others develop bacteraemia or 48 symptoms consistent with enteric fever [3,4]. This could be explained in part by unmeasured factors such as 49 effective bacterial dose reaching the intestinal mucosa, or other random effects not amenable to control. 50 Alternatively, certain participants may have an innate resistance or susceptibility to enteric fever: in 51 unvaccinated human challenge participants undergoing homologous re-challenge with S. Typhi, those who 52 did not develop enteric fever on the first exposure were less likely to develop enteric fever on the second 53 exposure [5]. Host genetics could play a role in this resistance. Genome-wide association studies (GWAS) 54 are frequently used to find associations between genetic variants and complex non-Mendelian traits, with 55 the aim of identifying genes which may provide insight into the pathology of a disease. For example, a GWAS identified polymorphisms in the NOD2 pathway as being associated with leprosy susceptibility [6]. 56 57 NOD2 activation was later found to induce dendritic cell differentiation, which may protect against disease 58 progression [7]. In the case of Salmonella infections, GWAS have revealed the HLA-DRB1*04:05 allele as

conferring resistance against typhoid fever [8] and a locus in *STAT4* as being associated with non-typhoidal *Salmonella* bacteraemia [9].

61

In epidemiological studies, genetic heterogeneity in the pathogen is a confounder to the infected human 62 63 host's individual susceptibility to that pathogen, as illustrated by studies of tuberculosis, in which host SNPs predispose individuals to infection with a particular strain only [10]. In studies performed at our centre to 64 65 date, only three strains of Salmonella have been used as a challenge agent, which has allowed us to 66 statistically control for pathogen heterogeneity. All participants are exposed to the pathogen under highly 67 controlled conditions, whereas in the field "non-infected controls" may have avoided infection due to lack 68 of environmental exposure rather than having been exposed and resisted infection. Furthermore, prior 69 exposure modifies susceptibility to enteric fever [5], which is difficult to account for in the field as 70 Salmonella exposure is likely frequent during childhood in endemic settings. However this can be managed 71 in challenge studies through strict inclusion criteria and careful screening, including exclusion of participants who had received a typhoid vaccine or lived in a typhoid-endemic area. Despite the advantages 72 73 of human challenge studies, to our knowledge a GWAS has not previously been carried out on human 74 challenge participants. Here we exploit this unique setting, and investigate how differences in host genetics 75 relate to outcome of challenge. We identify SNPs within the genes CAPN14 and MIATNB as having $p < 10^{-5}$ 76 for association with development of enteric fever symptoms or bacteraemia following exposure. We find 77 that HLA-B*27:05 is the HLA type most strongly associated with enteric fever susceptibility, enhancing 78 intracellular replication of S. Typhi.

79

80 Methods

81 Enteric Fever Human Challenge Cohorts

Five enteric fever human challenge cohorts from studies conducted at the Centre for Clinical Vaccinology and Tropical Medicine (Churchill Hospital, Oxford, UK) were included in this analysis: a typhoid dosefinding study, a paratyphoid dose-finding study, a typhoid oral vaccine study, a typhoid Vi vaccine study, and a study investigating the role of the typhoid toxin, summarised in Table 1. All participants provided written informed consent. Following challenge, individuals with fever (sustained oral temperature $\geq 38^{\circ}$) or positive blood culture were diagnosed with enteric fever. All challenged participants were treated with

ciprofloxacin or azithromycin either at time of diagnosis in diagnosed individuals, or after completing the
14-day challenge period if undiagnosed. Peripheral blood samples from participants from five different
enteric fever human challenge cohorts were either genotyped or transcriptionally profiled, or in some cases
both (Figure 1). A subset of participants underwent longitudinal transcriptional profiling, with data available
from up to nine time points.

93

94 Genotyping

DNA was extracted from blood clots using a QIA Symphony SP. Briefly, 180ul of ATL buffer (QiagenTM) 95 96 was added to each clot and then vortexed and incubated overnight at 56°C for lysis. The following day 200ul 97 of AL buffer (QiagenTM) was added to the lysed clot and mixed before transferring 500ul of the lysate to a 98 2ml tube and run on the QIA Symphony using the QIA Symphony DSP DNA Midi kit (QiagenTM). The 99 protocol was a customised BC 400 protocol and DNA was eluted into 100ul. Samples were quantified using 100 the Qubit and Qubit BR dsDNA reagents (Invitrogen). Samples from the typhoid dose finding and typhoid 101 oral vaccine trial (total n=96) were genotyped by the Wellcome Trust Centre for Human Genetics using an 102 Illumina OmniExpress-24 v1.0 BeadChip array, while samples from the paratyphoid dose finding study and typhoid toxin study (total n=80) were genotyped by Cambridge Genomic Services using an Illumina 103 OmniExpress-24 v1.3 BeadChip array. Data cleaning for association analysis was carried out in PLINK 104 105 [14]. Data processing steps are summarised in Figure 1.

106

Association analysis was carried out using a logistic regression model in PLINK, with challenge dose,
vaccination status and principal components as covariates. The online tool SNPnexus [15] was used to
identify genes proximal to SNPs. With the HapMap CEU dataset as a reference, SNP2HLA software [16]
was used to impute single nucleotide polymorphisms in the HLA region and identify HLA alleles.

111

112 RNA-sequencing

Whole blood samples were collected in Tempus Blood RNA tubes. RNA samples from the paratyphoid dose-finding and Vi vaccine trial were poly-A selected and underwent paired-end using a HiSeq V4 at the Wellcome Trust Sanger Institute. RNA samples from the typhoid toxin study underwent poly-A selection and paired end sequencing at the Beijing Genomics Institute using an Illumina HiSeq4000. Fastq files from

117	the same sample were concatenated. Paired fastq files were aligned to a pre-built graph reference using
118	HISAT2, followed by extraction of HLA-aligning reads. HLA typing and assembly was then carried out
119	using HISAT-genotype [17].
120	
121	Correlation between HLA types imputed from different time points and by different
122	methods
123	To assess the consistency of HISAT-genotype in imputing HLA-types, we compared estimated HLA type
124	doses given for the same participant from whole blood samples collected at different timepoints. A Pearson
125	correlation analysis was carried out on raw dosages for each pairwise comparison between timepoints to
126	give a Pearson correlation coefficient (R) and p value for strength of association. We calculated both
127	whether within a participant, dosage of each HLA type was consistent between timepoints, and within a
128	HLA type, whether the dosage for each participant was consistent between timepoints.
129	
130	To assess agreement between HLA types imputed by HISAT-genotype and SNP2HLA, for 71 participants
131	where both genotyping and RNA-sequencing data were available, dosage was rounded for the closest 50%.
132	As for certain participants HISAT-genotype results were available from RNA samples taken at different
133	timepoints, the median was taken for each participant. A Pearson correlation analysis was carried out to
134	compare estimated dosages given by HISAT-genotype and SNP2HLA. As above, we calculated both
135	whether within a participant, the dosage of each HLA type was consistent between methods, and within a
136	HLA type, whether the dosage for each participant was consistent between methods.
137	
138	Association between HLA type and outcome
139	Dosages were rounded to the nearest 50%, and for participants with multiple timepoints HLA-typed by
140	HISAT-genotype, any timepoints with outlying dosages (Figure S1) were excluded and the median of the

141 remaining timepoints taken. HLA types where there was no significant correlation (p>0.05) between

- 142 timepoints were excluded. For those with both SNP2HLA and HISAT-genotype derived HLA-types the
- 143 mean dosage was then taken. HLA type data from all cohorts were then combined. A logistic regression
- 144 model was used to identify HLA types associated with outcome (1=diagnosed with enteric fever,

145 0=remained undiagnosed). Vaccination status, challenge dose and challenge strain were included as

- 146 covariates. Statistical tests were carried out in R.
- 147

148 Intracellular survival of *S*. Typhi in HLA-B*27:05⁺ cells

149 HLA-B*27:05⁺ C1R cells generated using lentiviral constructs were provided by the Bowness Group [18].

150 Transfected control and HLA-B*27:05⁺ cells were seeded in a 96 well plate at a density of 100,000 cells per

151 well. A frozen glycerol stock of 5 x 10⁸ CFU/ml S. Typhi Quailes strain was thawed and washed twice with

152 RPMI 1640 Media. Cells were inoculated at a multiplicity of infection (MOI) of 50 in triplicate. After one

hour, gentamycin was added at a concentration of 200ug/ml. 24 hours post-inoculation cells were washed

twice with RPMI then resuspended in 50ul 1% Triton-X100. After two minutes, lysates were serially diluted

- in PBS and plated onto tryptone soya agar. Colonies were counted following overnight incubation at 37°C.
- 156 A one-tailed t-test was used to assess whether the number of colonies was higher in HLA-B $*27:05^+$ cells.
- 157

158 Differences in gene expression in those with HLA-B*27:05

159 Pre-alignment quality control on sequenced samples from the paratyphoid dose-finding study and Vi

160 vaccine trial was carried out using FASTQC. As all files had high phred scores (>25) across their length, all

161 were aligned to the human genome (GRCh38 Gencode version 26) using STAR-2.6.1c [19]. Total reads per

sample ranged from 16-44 million. Reads per gene were counted using the STAR GeneCounts mode.

163 Principal component analysis was used for outlier detection, with no samples excluded on this basis. Non-

164 protein coding and haemoglobin subunit genes were excluded. Count tables were filtered to exclude genes

165 with <1 count per million (cpm) in >31 samples (the number of baseline samples in control participants

166 challenged with S. Typhi) and normalised using weighted trimmed mean of M-values scaling (edgeR). The

167 count matrix was transformed using limma voom, and a linear regression model fitted with vaccination

status, challenge strain, sequence pool and dose as covariates and participant ID as a blocking variable. At

- baseline and 12 hours post-challenge, differential gene expression analysis between those with and without
- a copy of HLA-B*27:05 was carried out, filtering to genes with average log₂(expression)>0.

171

172 Gene set enrichment analysis

173	Differences in gene expression between human challenge participants with and without a copy of HLA-
174	B*27:05 were ranked by t-statistic at both baseline and 12 hours post-challenge. The entire ranked gene list,
175	including non-significantly differentially expressed genes, were input into GSEA 4.1.0 software [20]. A
176	custom gene set was created containing genes relating to the unfolded protein response and heat shock
177	response (Table S1). An enrichment score reflecting the degree to which these genes were over-represented
178	at the top of each ranked gene list was calculated. The p value of the enrichment score was then calculated
179	by the GSEA 4.1.0 software using an empirical phenotype-based permutation test procedure [20].
180	
181	Results
182	No SNPs were significantly associated with the outcome of challenge at the genome-wide
183	level
184	A genome-wide association analysis was carried out on genotyped participants (103 cases of enteric fever,
185	68 controls following data cleaning) in order to identify any SNPs associated with development of fever,
186	symptoms or bacteraemia following S. Typhi or S. Paratyphi A challenge (Figure 2). No SNPs reached
187	genome-wide significance, with two SNPs within the genes CAPN14 and MIATNB giving a p value below
188	1x10 ⁻⁵ (Figure S2).
189	
190	HLA-B*27:05 is associated with susceptibility to enteric fever
191	Given the number of individuals was too small to identify SNPs at the genome-wide level, we then focused
192	on variation within the HLA region. HLA typing was performed either by imputation from genotyping data
193	using SNP2HLA [16], or from raw RNA-sequencing data using HISAT-genotype [17]. We found HISAT-
194	genotype to be highly consistent between RNA-sequencing samples taken from the same participant at
195	different time points (Figures 3a and 3b; Table S3). For participants with both genotyping and RNA-
196	sequencing data, HLA-typing using HISAT-genotype significantly correlated with the results of SNP2HLA
197	imputation (Figures 3c and 3d).
198	
199	The most common HLA-A, -B, -C, -DQA, -DQB1 and -DRB1 allele groups were A*02, B*07, C*07,
200	DQA*01, DQB1*06 and DRB1*15 respectively (Figure 4a; Table S4). To identify whether any HLA types
201	were associated with enteric fever, a logistic regression was carried out on HLA types at a 2-digit resolution.

202	The HLA type most associated with susceptibility was HLA-B*27 (p=0.015, odds ratio=1.04, 95%
203	confidence intervals 1.01-1.08, Figure 4b). While a small odds ratio, this finding was of particular interest
204	as HLA-B*27 has been associated with non-typhoidal Salmonella-induced reactive arthritis and ankylosing
205	spondylitis [21–24]. At 4-digit resolution this association was driven by HLA-B*27:05 (Figure 4c). Of 10
206	participants heterozygous for HLA-B*27:05, 9 were diagnosed with enteric fever (Figure 4d). While HLA-
207	B*27:05 is most common in European populations, and the cohort analysed was predominantly white
208	(Figure 5a), in the 1000 Genomes project HLA-B*27:05 was present in both the Punjabi population in
209	Pakistan and Bengali population in Bangladesh, two countries where enteric fever is endemic [1,25](Figure
210	5b).
211	
212	To investigate the mechanism by which HLA-B*27:05 may contribute to enteric fever susceptibility, C1R
213	cells transfected with HLA-B*27:05 were infected with S. Typhi in vitro for 24 hours. Compared with non-
214	transfected controls, higher numbers of viable bacteria were recovered from HLA-B*27:05 ⁺ cells (Figure
215	6a), suggesting a mechanism independent of antigen presentation. This is consistent with previous literature
216	finding that HLA-B*27:05 lowers the threshold for induction of the unfolded protein response, a pathway
217	that is induced by and enhances intracellular S. Typhimurium infection [26].
218	
219	To investigate whether differences in the unfolded protein response can be detected in human challenge
220	participants, we explored transcriptional differences between those who did and did not possess a copy of

221 HLA-B*27:05 in the paratyphoid dose finding and Vi vaccine trial studies. We hypothesised that outcome

of challenge is dependent on events occuring early after exposure and preceding development of acute

disease, and therefore focused on 12 hours post-challenge, the timepoint at which dissemination of typhoidal

224 Salmonella is thought to take place in the blood [27,28]. At 12 hours post-challenge with S. Typhi or S.

225 Paratyphi A, the most significant differentially expressed gene expressed gene between the two groups was

MICA (MHC Class I Polypeptide-Related Sequence A), encoding a ligand for NK cell activating receptor

- 227 NKG2D (Figures 6b and 6c). Expression of *MICA* is inhibited by the unfolded protein response [29] and
- 228 was expressed at lower levels by those with a copy of HLA-B*27:05 (p=0.00006 12 hours post-challenge,

229 p=0.006 at baseline, linear modelling). The gene CALR encoding the calcium-binding chaperone calreticulin

- 230 was more highly expressed in those with HLA-B*27:05 at 12 hours post-challenge but not at baseline
- 231 (p=0.04 12 hours post-challenge, p=0.8 at baseline, Figure 6c). Gene set enrichment analysis [20] was then

8

232	used to assess	whether trans	scripts encodin	g proteins	involved in	n the unfolded	protein response	e were
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enriched amongst those with HLA-B*27:05. A custom gene set containing CALR, ATF4, DDIT3, HSPA5,

234 XBP1, EDEM1, HYOU1 as well as 61 genes annotated as relating to the heat-shock response was over-

represented in those with HLA-B*27:05 at 12 hours post-challenge when ranked by t-statistic (p=0.01,

Figure 6d). This gene set was not over-represented at baseline (p=0.4).

237

238 Discussion

239 This study investigated genetic susceptibility to enteric fever in a human challenge setting. We found

240 HISAT-genotype to be a consistent tool to impute HLA types from RNA-sequencing data, with HLA

241 dosages correlating significantly with SNP2HLA dosages imputed from genotyping data. Of the HLA-types,

242 HLA-B*27:05 was most associated with susceptibility to infection (p=0.012). Although participants were

243 predominantly European, HLA-B*27:05 is also present in certain South Asian populations. HLA-B*27:05

244 mis-folds in the endoplasmic reticulum (ER), reducing the threshold for activation of the unfolded protein

response, and has been linked with both reactive arthritis following Gram-negative bacterial infection [26],

and ankylosing spondylitis [30]. When infected in vitro with S. Typhimurium, both monocyte-like U937

247 cells and epithelial HeLa cells transfected with HLA-B*27:05 exhibit higher levels of intracellular

replication [26,31]. Although the exact mechanism is unknown, the unfolded protein response appears to

create a favourable environment for S. Typhimurium, the presence of HLA-B*27:05 increasing its

expression of SPI-2 genes [32] and causing it to replicate at the cell periphery [26]. Enhanced replication of

251 S. Typhimurium was abrogated when HLA-B*27:05 was stabilised by fusion with beta-2-microglobulin

252 [26]. Pharmacological induction of ER stress by thapsigargin enhances S. Typhimurium replication, while

253 infection with S. Typhimurium stimulates the unfolded protein response by a mechanism dependent on

bacterial effector *sifA* [26]. Although *sifA* is also present in *S*. Typhi, its sequence differs to sifA in *S*.

255 Typhimurium [33]. However, we still observed enhanced replication of S. Typhi Quailes strain in C1R cells

256 (p=0.02, one-tailed t-test), suggesting this phenomenon is not serovar-specific.

257

258 The gene encoding ER chaperone calreticulin, *CALR*, was higher in HLA-B*27:05⁺ human volunteers 12

bours following enteric fever challenge but not at baseline (p=0.04 12 hours post-challenge, p=0.8 at

260 baseline). Gene set enrichment analysis [20] was then used to assess whether transcripts encoding proteins

261 involved in the unfolded protein response were enriched amongst those with HLA-B*27:05. A custom gene 262 set containing genes annotated as relating to the unfolded protein response and heat-shock response was over-represented in those with HLA-B*27:05 at 12 hours post-challenge when ranked by t-statistic (p=0.01). 263 264 However this gene set was not over-represented at baseline (p=0.4). This supports the hypothesis that HLA-265 B*27:05 reduces the threshold for unfolded protein response activation in infection. At 12 hours post-266 challenge, the most significant differentially expressed gene expressed gene between the two groups was 267 MICA, encoding a ligand for NK cell activating receptor NKG2D (p=0.00006, linear modelling). MICA is downregulated by the unfolded protein response [29], and was expressed at lower levels in participants with 268 269 HLA-B*27:05 12 hours post-challenge. In viral infections, downregulation of MICA prevents recognition by 270 NK cells [34]. Polymorphisms in MICA have been related to susceptibility to leprosy, which, in common 271 with enteric fever, infects mononuclear phagocytes [35-37]. In contrast to CALR, MICA was also 272 differentially expressed in HLA-B*27:05⁺ participants at baseline (p=0.06, linear modelling), suggesting 273 either that HLA-B*27:05 can induce certain aspects of the unfolded protein response in the absence of 274 infection, or that its decreased expression is mediated by a different mechanism.

275

276 In the absence of SNPs with very high odds ratios in our cohort, we were underpowered to detect significant 277 SNPs at a genome wide level. The SNP with the lowest p value (rs4952069, 4.2 x 10⁻⁶) falls in the intronic 278 region of CAPN14, a calcium-dependent cysteine protease regulated by Th2 cytokines IL-13 and IL-4 [38]. 279 Intronic SNPs may either be linked to a causative coding SNP, or themselves affect gene expression through 280 splicing or transcription factor binding [39]. CAPN14 is thought to play a regulatory role in the oesophageal 281 epithelium, with overexpression impairing barrier function, and SNPs in this locus having been associated 282 with susceptibility to the allergic inflammatory disease eosinophilic oesophagitis [40] and middle ear infection [41]. While the cellular response to enteric fever infection is Th1 dominated, Th2 cytokines may 283 284 be modulated by infection, with S. Typhi-specific IL-13 secretion observed in peripheral blood mononuclear 285 cells isolated during typhoid fever convalescence [45] and IL-4 secreted at the apical side of intestinal biopsies infected in vitro with S. Typhi [46]. Co-infection of mice with both S. Typhimurium and Th2-286 287 inducing hookworms impairs clearance of S. Typhimurium, suggesting that polarisation towards a Th2 288 response could be detrimental [47]. Therefore genetic variations predisposing individuals to a more Th2-289 dominant response to infection could feasibly affect susceptibility to enteric fever.

291 This is the first genetic study to investigate susceptibility to infection using samples obtained from human 292 challenge volunteers. Furthermore, while HLA-B*27:05 has been linked to non-typhoidal Salmonella 293 infections, this is the first study to find an association with enteric fever. However, we were limited by 294 several factors. Firstly, there were cases where the HLA type of a participant was ambiguous, predominantly 295 due to SNP2HLA suggesting several possible HLA types, but also incomplete agreement between 296 SNP2HLA and HISAT-genotype dosages. Secondly, due to the nature of human challenge studies, our 297 sample size was smaller than conventional GWAS. While notable GWAS with smaller samples than ours 298 have included those associating genetic variants with vitiligo and response to anti-TNF treatment, a larger 299 sample would have enabled us to detect associations with smaller effect sizes [48,49]. Only 10 participants 300 were unambiguously identified as HLA-B*27:05⁺ and the odds ratio was small in magnitude, suggesting 301 that HLA-B*27:05 explains only a small proportion of innate susceptibility to enteric fever. However, given 302 previous evidence of an association with both Salmonella-induced reactive arthritis and intracellular S. 303 Typhimurium replication, this intriguing association warrants validation by further studies. Finally, this 304 study was carried out a predominantly European cohort not previously exposed to typhoidal Salmonella. 305 While this allowed us to investigate genetic susceptibility without the confounding factor of previous 306 exposure, it is not representative of the population in an endemic setting. However, it could have 307 implications for travel medicine: for example, those with a family history of ankylosing spondylitis could be 308 strongly encouraged to undergo typhoid vaccination prior to travelling. Although reactive arthritis following 309 live oral typhoid vaccination is a rare complication [50], parenteral vaccination may be preferable in this 310 case. Furthermore, HLA-B*27:05 is present both in Punjabi and Bengali populations, suggesting this allele 311 could play a role in an endemic setting [25].

312

313 Acknowledgements

The authors would like to thank the volunteers for participating in the studies, and the High-Throughput Genomics Group at the Wellcome Trust Centre for Human Genetics for the generation of genotyping data. We would also like to thank the Data Safety and Monitoring Committee for providing safety oversight of the studies, the University of Maryland for providing *S*. Typhi Quailes challenge strain and GSK Vaccines Institute for Global Health for providing *S*. Paratyphi A NVGH308 strain. Finally, we would like to

319	acknow	wledge Paul Bowness for supervising transfection of the transgenic C1R cells, and the additional
320	clinica	al and laboratory support provided by Oxford Vaccine Group staff during the enteric fever studies.
321		
322	Foot	notes
323	Conf	lict of interest statement
324	AJP is	Chair of the UK Department of Health and Social Care's (DHSC) Joint Committee on Vaccination &
325	Immu	nisation (JCVI) and is a member of the WHO's Strategic Advisory Group of Experts. CJB is currently
326	emplo	yed by GlaxoSmithKline.
327		
328	Fund	ing statement
329	This v	vork was supported by the Wellcome Trust [grant numbers 092661/Z/10/Z awarded to AJP,
330	09053	2/Z/09/Z]; The Bill & Melinda Gates Foundation [OPP1084259; Global Health Vaccine
331	Accele	erator Platform grant OPP1113682]; European Commission FP7 grant "Advanced Immunization
332	Techn	ologies" (ADITEC) and the NIHR Oxford Biomedical Research Centre.
333		
334	Meet	ings where work was previously presented
335	BSI vi	rtual conference, December 2020, United Kingdom, Abstract ID 989
336		
337	Corre	esponding author information
338	Ambe	r J Barton
339	Ambe	r.Barton@lshtm.ac.uk
340		
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Figure and tables

Figure 1: Number of participants and samples at each stage of the analysis pipeline

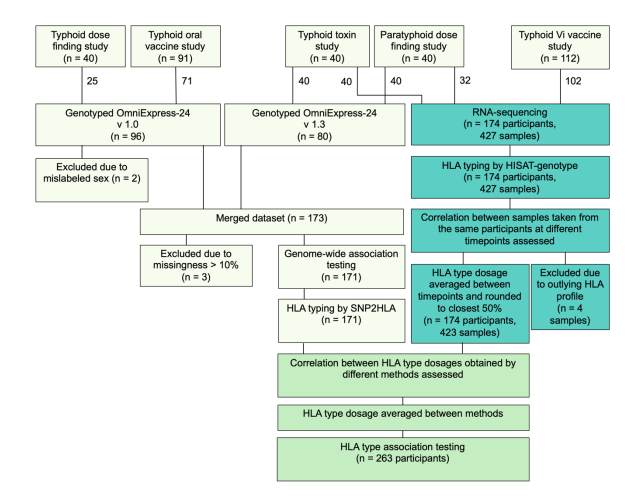


Figure 2: Manhattan plot showing the significance (-log₁₀(unadjusted p value)) of the relationship between each single nucleotide polymorphism (SNP) and development of symptoms or bacteraemia following oral *S*. Typhi or *S*. Paratyphi A challenge, for each chromosome. The dotted line indicates a suggestive p value of 10⁻⁵. The ten SNPs with the lowest p values are highlighted, with the nearest proximal gene as identified by SNPnexus indicated as well as the odds ratio (OR).

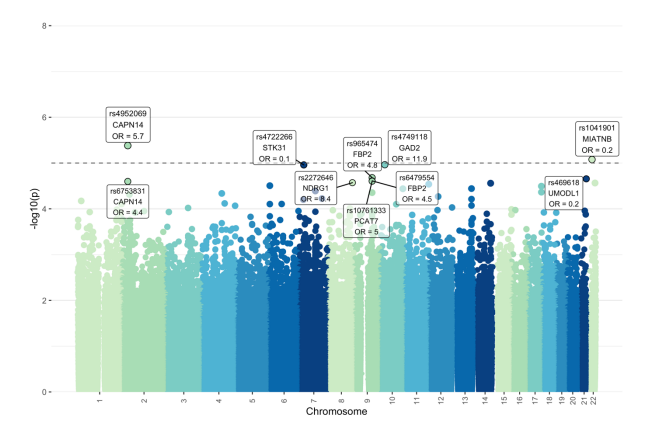


Figure 3: Distribution of squared Pearson correlation coefficients (R^2) for HLA types at a 2-digit resolution and 4-digit resolution. Quantiles are indicated by vertical lines. Points are coloured by whether in a test of whether the Pearson correlation coefficient is different to zero, the p value was below 0.05.

a. Correlation between the doses of each HLA type at different timepoints within each participant. HLA

types were profiled from RNA-sequencing samples using HISAT-genotype, giving a dose (0-100%) of each

HLA type for each participant. Each point represents one comparison; for participants where more than two timepoints were profiled, more than one point is shown per participant.

b. Correlation between the doses for each participant at different timepoints within each HLA type. HLA

types were profiled from RNA-sequencing samples using HISAT-genotype, giving a dose (0-100%) of each

HLA type for each participant. Each point represents one HLA type.

c. Correlation between the median doses of each HLA type for the same participant using either SNP2HLA imputation from genotyping data or HISAT-genotype typing from RNA-sequencing data. Each point represents one participant.

d. Correlation within each HLA type between the doses for each participant using either SNP2HLA imputation from genotyping data or HISAT-genotype typing from RNA-sequencing data. Each point represents one HLA type.

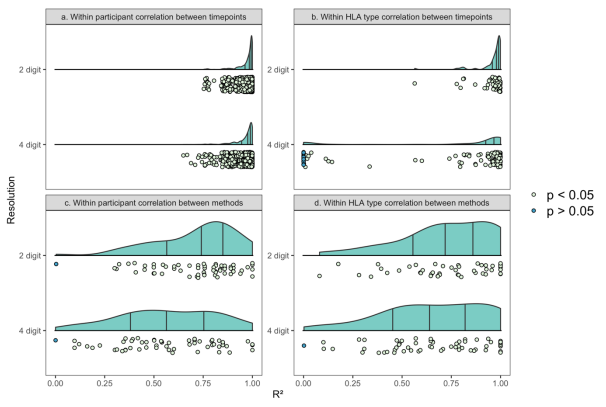


Figure 4: a. Relative frequency of each HLA type at a resolution of 2 digits for HLA-A, HLA-B, HLA-C, HLA-DQA1, HLA-DQB1 and HLA-DRB1 in the entire combined cohort, including participants from the typhoid dose finding study, typhoid oral vaccine trial, typhoid Vi vaccine trial, paratyphoid dose finding study and typhoid toxin study.

b. Odds ratios (odds ratio >1 indicates association with susceptibility and <1 with resistance) and 95% confidence intervals for the five HLA types most significantly associated with outcome of challenge at a resolution of 2 digits. P values are indicated for each.

c. Odds ratios for the two HLA-B*27 sub-types at a resolution of 4-digits with 95% confidence intervals. P values are indicated for each.

d. Percentage of participants who were diagnosed with enteric fever following challenge, stratified by the presence or absence of one copy of HLA-B*27:05. The proportion of participants diagnosed is indicated for each group.

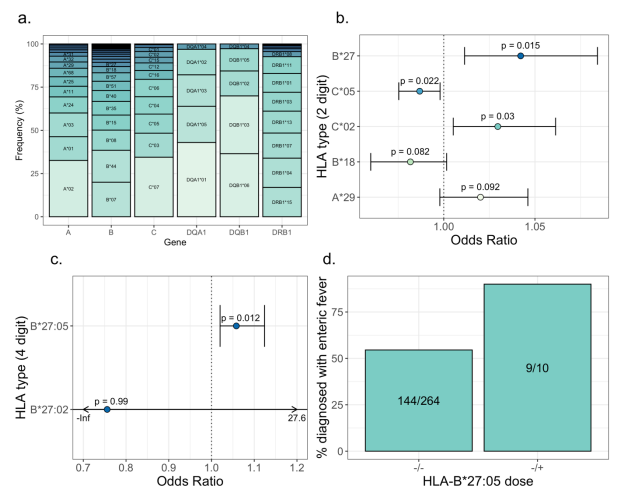
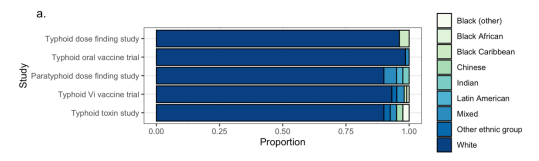
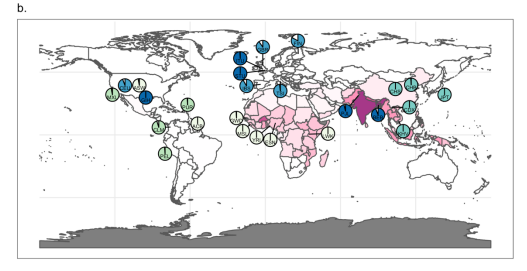


Figure 5: a. Self-reported ethnicity of participants within each study. b. The white sector of each pie chart indicates the proportion of 1000 Genome Project participants with at least one HLA-B*27:05 allele in each population. The remainder of the pie chart is coloured by ancestral continental region. Each country is coloured by enteric fever incidence rate per 100,000 as estimated by Stanaway et al. 2019. CHB = Han Chinese in Beijing, China, JPT = Japanese in Tokyo, Japan, CHS = Southern Han Chinese, CDX = Chinese Dai in Xishuangbanna, China, KHV = Kinh in Ho Chi Minh City, Vietnam, CEU = Utah Residents with Northern and Western European Ancestry, TSI = Toscani in Italia, FIN = Finnish in Finland, GBR = British in England and Scotland, IBS = Iberian Population in Spain, YRI = Yoruba in Ibadan, Nigeria, LWK = Luhya in Webuye, Kenya, GWD = Gambian in Western Divisions in the Gambia, MSL = Mende in Sierra Leone, ESN = Esan in Nigeria, ASW = Americans of African Ancestry in SW USA, ACB = African Caribbeans in Barbados, MXL = Mexican Ancestry from Los Angeles USA, PUR = Puerto Ricans from Puerto Rico, CLM = Colombians from Medellin, Colombia, PEL = Peruvians from Lima, Peru, GIH = Gujarati Indian from Houston, Texas, PJL = Punjabi from Lahore, Pakistan, BEB = Bengali from Bangladesh, STU = Sri Lankan Tamil from the UK, ITU = Indian Telugu from the UK.





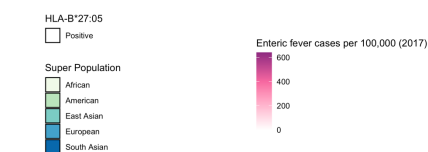


Figure 6: a. Colony forming units per ml recovered from C1R cells infected with S. Typhi Quailes strain, in the presence or absence of HLA-B*27 expression, 24 hours post-infection. Parent and HLA-B*27:05+ cells were seeded in a 96 well plate at a density of 100,000 cells per well, and infected with S. Typhi Quailes strain at an MOI of 0 or 10 in triplicate. After one hour gentamycin was added to kill extracellular bacteria. 24 hours post-inoculation cells were lysed using 1% Triton-X100, and lysates serially diluted and plated onto tryptone soya agar. Colonies were counted following overnight incubation at 37°C. A p value for a t-test is indicated. Points represent replicates within a single experiment.

b. Volcano plot showing the log_2 (Fold Difference) in gene expression between HLA-B*27:05 positive and negative participants 12 hours post-challenge against the $-log_{10}$ (p-value). A dashed line indicating where p = 0.05 is shown, and genes relating to the unfolded protein response and heat shock proteins are highlighted. Genes more highly expressed in participants who were HLA-B*27:05 positive are shown positive further to the right, and those more highly expressed in HLA-B*27:05 negative participants further to the left. RNA expression was characterised by RNA-sequencing. Data were filtered, normalised and transformed, and differential expression then assessed using the limma R package, using participant ID, sequencing pool, vaccination status, challenge strain and dose as blocking variables.

c. Expression of *MICA* and *CALR* following normalisation and transformation using the edgeR and limma packages, in HLA-B*27:05 positive and negative participants at baseline and 12 hours post-challenge. d. Running enrichment score for a custom gene set containing genes involved in the unfolded protein and heat shock response. Gene set enrichment analysis calculates an enrichment score by walking down a list of genes ranked by t statistic. When a gene within a gene set is encountered the running enrichment score is the maximum deviation from zero. The genes in the custom gene set are indicated.

