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1	A rationally designed oral vaccine induces Immunoglobulin A in the
2	murine gut that directs the evolution of attenuated Salmonella variants
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55 Introductory paragraph

56 The ability of gut bacterial pathogens to escape immunity by antigenic variation, 57 particularly via changes to surface-exposed antigens, is a major barrier to 58 immune clearance¹. However, not all variants are equally fit in all environments^{2,3}. It should therefore be possible to exploit such immune escape 59 60 mechanisms to direct an evolutionary trade-off. Here we demonstrated this 61 phenomenon using Salmonella enterica subspecies enterica serovar 62 Typhimurium (S.Tm). A dominant surface antigen of S.Tm is its O-antigen: A 63 long, repetitive glycan that can be rapidly varied by mutations in biosynthetic pathways or by phase-variation^{4,5}. We quantified the selective advantage of O-64 65 antigen variants in the presence and absence of O-antigen specific IgA and 66 identified a set of evolutionary trajectories allowing immune escape without an 67 associated fitness cost in naïve mice. Through the use of oral vaccines, we 68 rationally induced IgA responses blocking all of these trajectories, which 69 selected for Salmonella mutants carrying deletions of the O-antigen 70 polymerase wzyB. Due to their short O-antigen, these evolved mutants were 71 more susceptible to environmental stressors (detergents, complement), 72 predation (bacteriophages), and were impaired in gut colonization and 73 virulence in mice. Therefore, a rationally induced cocktail of intestinal antibodies 74 can direct an evolutionary trade-off in S.Tm. This lays the foundations for the 75 exploration of mucosal vaccines capable of setting evolutionary traps as a 76 prophylactic strategy.

77

78 Main text

79 The gut is a challenging environment for bacteria with high densities of phage, 80 bile acids, antimicrobial peptides and secretory antibodies. These interact first 81 with the outermost layer of the bacterial surface. Long, repetitive glycans, such 82 as capsular polysaccharide, teichoic acids or O-antigens are ubiquitous as the 83 outermost defense in bacteria. A particularly relevant feature of these glycan 84 structures is that small changes in the structure of the repeating units, such as 85 gain or loss of acetyl groups, when polymerized, result in major changes in 86 conformation and charge-distribution of the glycans.

87 In the case of non-Typhoidal Salmonella this outermost glycan layer is 88 predominantly made up of O-antigen: lipopolysaccharide core-linked, long, 89 repetitive heteroglycans that hide most common outer-membrane proteins (^{6,7}, 90 **Fig.ED1**). The S.Tm wild type (S.Tm^{WT}) O:4[5], 12-0 O-antigen is a polymer of 91 а triose repeating backbone (-mannose- α -(1 \rightarrow 4)-rhamnose- α -(1 \rightarrow 3)-92 galactose- α -(1 \rightarrow 2), constituting the O:12-0 epitope) with an α -(1 \rightarrow 3)-abequose 93 side-branch at the mannose (constituting the O:4 epitope, or when O-acetylated the O:5 epitope) (Fig. 1A). The S.Tm^{WT} reacts to O:5-typing antisera and O:12-94 0-typing antibodies (Fig. 1B, and C, S1-3). In the SL1344 strain of S.Tm, two 95 96 major shifts in O-antigen composition have been reported. Firstly, complete loss 97 of abequose acetylation, generating an O:4-only phenotype, occurs via loss of 98 function mutations in the abequose acetyl transferase gene $oafA^8$, (Fig. 1A and 99 **B**). Secondly, the O:12-0 epitope can be converted to an O:12-2 epitope by (α -100 $(1\rightarrow 4)$ glucosylation of the backbone galactose (**Fig. 1A and C**). This occurs 101 via expression of a glucosyl transferase gtrABC operon (STM0557-0559), 102 controlled by DAM-dependent methylation i.e. by phase variation^{4,9}. Note that 103 S.Tm strain SL1344 lacks a second common operon required for linking

104 glucose via an α -(1 \rightarrow 6) linkage to the backbone galactose, generating the O:1 105 serotype. All of these structural O-antigen variants exert only a mild fitness defect in the naïve gut (^{5,9,10}, **Fig 1D and E**). However, there is also evidence 106 107 for selection of mutants at loci coding for the O-antigen polymerases and socalled "non-typable" Salmonella strains with a single-repeat O-antigen are 108 occasionally observed amongst isolates from infected humans or animals¹¹. 109 Such strains lose outer membrane robustness, due to loss of the rigid 110 hydrophilic glycan layer¹², . and therefore have decreased fitness both in the 111 gut and in the environment 2,3,13 . 112

113 We hypothesized that the host's immune response could generate conditions 114 in which the fitness of O-antigen polymerase mutants is promoted, driving the 115 emergence of an evolutionary trade-off. Intestinal antibodies (predominantly 116 secretory IgA) are known to exert specific selective pressures on targeted species^{14–16}. In order to investigate the evolutionary consequences of vaccine-117 118 induced secretory antibody responses in the gut, without the major ecological shifts associated with live-attenuated vaccine infection^{17–19}, we made use of an 119 established high dose, inactivated oral vaccination technique^{15,20,21} that induces 120 121 intestinal IgA responses without detectable intestinal damage, inflammation or colonization by the vaccine strains²¹. Our standard vaccine ("PA-S.Tm") 122 consists of concentrated peracetic acid killed bacteria²¹. Conventional mice 123 harboring a complex microbiota (16S amplicon analysis available²²) received 124 10¹⁰ particles of PA-S.Tm orally once per week for 4 weeks. Subsequently, 125 126 these mice were antibiotic-treated to open a niche for the pathogen in the large 127 intestine, and were infected with S.Tm SL1344, which rapidly colonizes the

128 cecum, generating typhlocolitis, and invasive disease in the mesenteric lymph
 129 nodes, spleen and liver^{23,24}.

130 We first quantified the competitive fitness of S.Tm mutants genetically "locked" 131 into individual structural O-antigen compositions in vaccinated and naïve mice. Competition between S.Tm^{$\Delta oafA \Delta gtrC$} (O:4, O:12-0-locked) and S.Tm^{$\Delta gtrC$} 132 133 (O:4[5], O:12-0-locked) demonstrated no difference in fitness in naïve mice over 4 days of infection. However, in mice vaccinated either against the O:4 or 134 135 the O:4[5] variant (Fig. S4), we observed up to a 10⁷-fold outcompetition of the 136 IgA-targeted O-antigen variant within 4 days (Fig. 1D). The magnitude of the selective advantage correlated with the magnitude of the intestinal IgA 137 138 response to each O-antigen variant (Fig. 1F and G). Therefore, IgA can exert 139 a strong selective pressure on the O:4/O:4[5] O-antigen variants. Competing S.Tm^{$\Delta oafA$} (**O:12-phase-variable**, O:4) against S.Tm^{$\Delta oafA$} $\Delta gtrC$ (**O:12-locked**, 140 141 O:4) revealed a mild benefit of O:12 phase variation in naïve mice up to day 4 post-infection, in line with published data (Fig. 1E)^{4,5}. However, we observe a 142 143 major fitness benefit of phase variation in vaccinated mice in which the IgA response is highly biased to recognition of O:12-0 O-antigens (Fig. 1E, H. Red 144 145 symbols, Fig. S5). Correspondingly, vaccinated mice with an outgrowth of 146 phase-variable S.Tm also displayed initiation of intestinal inflammation, as 147 quantified by fecal Lipocalin 2 (LCN2, Fig. 11). The mechanistic basis of this selective advantage could be confirmed by complementation of the *gtrC* gene 148 in trans (Fig. S6). Therefore O:12-0-targeting IgA can exert a strong selective 149 150 pressure against S.Tm unable to phase-vary the O:12-0 part of the O-antigen. 151 As neither of these variants (O:4[5] to O:4 and O:12-0 to O:12-2) are associated 152 with a major loss-of-fitness in naïve mice (Fig. 1D and E), this implied that such

variants should be selected for during infections of vaccinated mice with wildtype *Salmonella*.

155 We therefore established whether natural emergence of these "IgA-escape"-156 S.Tm variants occurred sufficiently fast to be observed during wild type S.Tm 157 infections. For this purpose, we treated mice with a wild type PA-S.Tm oral 158 vaccine as above, or with a vehicle-only control, and then challenged these 159 animals with wild type S.Tm. Around 30% of vaccinated mice showed intestinal 160 inflammation at 18 h post infection (Fig. 2A), despite the presence of robust anti-S.Tm^{wt} intestinal IgA in all vaccinated animals (Fig. 2B). When S.Tm 161 162 clones were recovered from the cecal content of vaccinated mice with intestinal 163 inflammation, these were typically recognized less well by vaccine-induced IgA 164 than S.Tm clones from the cecum of vaccinated and protected mice (Fig. 2C). 165 In 11 of 34 mice analysed, we observed clones with complete loss-of-binding 166 to an O:5-specific polyclonal antisera within 4 days (Table S3, Fig 2D). 167 Resequencing of O:5-negative clones confirmed a 7 bp contraction of a tandem repeat in the open reading frame of *oafA*, coding for the abequose acetylase 168 (Fig. 2E, 10 different clones from three independent experiments), that is also 169 170 found in multiple NCBI deposited genomes²⁵ (Fig. ED2A). A second site of 171 microsatellite instability is present in the promoter of *oafA* suggesting a further 172 possibility for rapid inactivation (Fig. ED2B), and this gene was found to be 173 under negative selection in a recent screen of published Salmonella genomes²⁶. 174

In contrast, loss of O:12-0 staining was bimodal within individual clones (Fig.
2F), consistent with phase-variation⁴ and no reproducible mutations were
identified in these clones on genome resequencing (Table S3). Instead,

methylation analysis revealed a methylation pattern indicative of the gtrABC 178 promoter being in an "ON" conformation (Fig. 2G). Serial passage of these 179 180 clones (Fig. ED3A), as well as cultivation in microfluidic devices 181 (Supplementary videos 1 and 2) confirmed the ability of clones to switch between O:12-0-positive and negative states. The STM0557-0559 gtrABC 182 183 locus was confirmed to be essential for this observed loss of O:12-0 epitope as strains lacking gtrC remained 100% O:12-0-positive even under strong in vivo 184 185 selection (Fig. ED3B and C). This phenotype could be replicated by adoptive 186 transfer of a recombinant monoclonal IgA specific for the O:12-0 epitope (mSTA121, Fig. ED4), confirming that O:12-0-binding IgA is sufficient to drive 187 188 outgrowth of O:12-2-producing variants. Computational modeling of phase-189 variation and growth, as well as comparison of O:12-0/O12:2 switching rates of 190 *lacZ* reporter strains suggested that selection for clones expressing *gtrABC* is 191 sufficient to explain the recovery rate, without any intrinsic shift in phase 192 variation switching rates (Fig. ED5). The chemical structure of O-antigen of the 193 recovered clones was further confirmed by ¹H-NMR of purified O-antigen and 194 by high resolution magic-angle spinning NMR of O-antigen on the surface of 195 intact cells (Fig. ED6). Therefore, vaccine-induced IgA can select for the natural 196 emergence of O-antigen variants within a few days of infection with S.Tm wild 197 type, resulting in disease in vaccinated mice. This phenomenon can also be 198 observed at later time-points in IqA-competent but not IqA-deficient mice during 199 chronic infection with live-attenuated S.Tm strains (Fig. ED7A and B) i.e. IgA 200 is necessary for selection of O-antigen variants during chronic infection. 201 Correspondingly, although the inactivated oral vaccines induce a higher titre of 202 Salmonella-binding IgA than the live vaccines (Fig. ED7C), the response to

203 chronic infection binds to O:4 and O4[5]-producing S.Tm with similar titres, 204 while the response to inactivated vaccine is highly biased for the O-antigen 205 variant of the vaccine (**Fig. ED7C**). This indicates that within-host O-antigen 206 variation also occurs under the selective pressure of intestinal antibodies during 207 chronic infections, and sequential priming will include a broad IgA response 208 capable of recognizing multiple O-antigen variants.

209

210 We next investigated whether the relative fitness defect of a short O-antigen 211 mutant can be compensated for by the selective advantage from lower IgA-212 binding in the gut lumen, i.e. whether IgA could drive an evolutionary trade-off. 213 One-on-one competitions were carried out between S.Tm^{ΔoafA ΔgtrC} (O:4,12-0locked, long O-antigen) and S.Tm^{\LoafA \Left_gtrC \Left_wzyB} (O:4,12-0-locked, short O-214 215 antigen, retains just a single O-antigen repeat) in the intestine of mice with and without IgA raised against S.Tm^{$\Delta oafA \Delta gtrC$} (Fig. 3A). The single repeat O-antigen 216 217 strain was rapidly outcompeted in naive animals, in line with earlier studies^{11,27} 218 (Fig. 3A) indicating a major loss-of-fitness. However, in the gut of vaccinated 219 mice, strains with short O-antigen were dominant by day 4 (Fig. 3A). 220 Vaccinated antibody-deficient mice were indistinguishable from naive mice in 221 these experiments, verifying that IgA is necessary for the selection of short O-222 antigen strains in the gut of vaccinated mice (Fig. 3A). Introduction of day 4 223 fecal bacteria from vaccinated mice into naïve mice resulted in re-outgrowth of 224 the strain with a long O-antigen, indicating that vaccine-induced IgA, and not secondary mutations in S.Tm^{\LoafA \LogtrC \LogTWZYB}, was responsible for competition 225 226 outcome (**Fig. 3B**). The IgA titre recognizing short O-antigen-producing strains 227 was lower than that against full-length O-antigen strains, consistent with the

selective advantage in vaccinated mice (**Fig. 3C**). As the long O-antigen can have several hundred repeats of the glycan, decreased antibody binding could be driven by lower O-antigen abundance or by loss of avidity-driven interactions. Loss of long O-antigen can therefore be an advantage to *Salmonella* in the gut lumen of vaccinated mice.

233 Based on these above observations, we hypothesized that emergence of mutants with a short O-antigen could be achieved for a wild type S.Tm infection 234 235 if we could block all other IgA escape routes, effectively generating an 236 evolutionary trap. To this end, mice received an oligovalent vaccine containing 237 the **O:4[5],12** S.Tm^{$\Delta gtrC$}, **O:4,12** S.Tm^{$\Delta oafA \ \Delta gtrC$}, **O:4,12-2** S.Tm^{$\Delta oafA \ pgtrABC$}, 238 and **O:4[5],12-2** S.Tm pgtrABC strains (referred to as PA-S.Tm^{ET}). This 239 induced a broad antibody response with high avidity for all four of the known 240 long O-antigen variants present in our S.Tm SL1344 strain (Fig. 3D, Fig.S7-8). PA-S.Tm^{ET} provided subtly better protection from intestinal inflammation in 241 242 long-term infection of 129S1/SvImJ mice than the monovalent O:5,12-0 vaccine 243 (Fig. 3E, significant protection from intestinal inflammation at d9 with PA-S.Tm^{ET} but not PA-S.Tm^{$\Delta gtrC$}), as well as on mixed challenge of Balb/c mice 244 (Fig. S9, significant protection from intestinal inflammation at d4 with 245 PA-S.Tm^{ET} but not PA-S.Tm^{$\Delta gtrC$}). Moreover, our hypothesis that this vaccine 246 247 can set an evolutionary trap was supported: short O-antigen-producing clones were detected in 12 of 18 PA-S.Tm^{ET} vaccinated mice analysed across multiple 248 experiments by phenotypic characterization (anti-O5^{dim} flow cytometry staining, 249 250 Fig. 3F). The O-antigen phenotype was confirmed by gel electrophoresis of purified LPS (Fig. 3G). Sequencing of evolved short-O-antigen clones (Table 251 252 **S4**, n=5) revealed a common large deletion encompassing the wzyB gene (also termed *rfc*), encoding the O-antigen polymerase¹¹ (**Fig. 3H**, **Fig. ED8** also reported in some "non-typable" *S*.Tm isolates from broilers¹¹). This deletion is mediated by site-specific recombination between flanking direct repeats, which renders the *wzyB* locus unstable¹¹.

We have previously published that IgA responses against the surface of rough 257 258 Salmonella are identically induced by vaccination with either rough or wild type Salmonella oral vaccines²⁸. Correspondingly, including a short-O-antigen 259 mutant into our PA-S.Tm^{ET} mix does not further improve IgA titres (Fig. S10). 260 261 Note that in these experiments, we also do not observe a significant improvement of protection with PA-S.Tm^{ET}, as PA-STm^{WT} protected well out to 262 263 day 3 in n=6 of 8 mice, when the experiment was terminated for ethical reasons 264 relating to the control group. As the generation of Salmonella O-antigen variants 265 is inherently stochastic, but a prerequisite for selection by IgA and therefore 266 within-host evolution, perfect protection can be observed in a variable fraction 267 of animals that had received the monovalent vaccine up to this time-point. However, no intestinal inflammation, as quantified by fecal Lipocalin-2, was 268 observed in any of the mice receiving PA-S.Tm^{ET} (n=9) or PA-S.Tm^{ET+wzyB} 269 270 (n=4).

We finally confirmed that re-isolated *wzyB*-deletion mutants phenocopied the fitness defects of targeted *wzyB* mutations in harsh environments. Single infections with $S.Tm^{\Delta oafA \, \Delta gtrC \, \Delta wzyB}$ revealed that, in comparison to isogenic wild type counterparts, *wzyB*-deficient mutants (synthetic or evolved) are significantly less efficient at colonizing the gut of streptomycin pretreated naïve mice (**Fig. 4A**), disseminating systemically (**Fig. 4B**) and triggering inflammation (**Fig. 4C**), i.e. they have an intrinsic defect in colonization and

virulence. This attenuation can be attributed to compromised outer membrane 278 integrity¹² and also manifests as an increased sensitivity to membrane 279 280 destabilization by EDTA, bile acids and weak detergents (Fig ED9A-E) and 281 increased sensitivity to complement-mediated lysis^{11,27} (Fig. ED9F). It is also 282 well-documented that specific interactions between the tail spike fiber and Oantigen reduce the host-range of ubiquitous lytic phages^{29,30}. Correspondingly, 283 infection of the short-O-antigen strains with filtered wastewater generated 284 285 visible lysis plaques of various sizes (Fig. 4D and E, Fig. ED10A). About 10-286 fold less lysis plagues were visible in the same conditions with long O-antigen 287 strains (Fig. 4D and E, Fig. ED10A). Sequencing of phages isolated from four 288 plaques revealed four different T5-like phages (Fig. 4F). Infections with the 289 purified phage $\phi 12$ yielded more phages after infection of a short O-antigen 290 evolved clone compared to the ancestor strains (Fig. 4G). We could confirm 291 that infection was dependent on btuB, the vitamin B12 outer-membrane transporter that is normally shielded by a long O-antigen (Fig, ED10B and 292 293 C). These results confirmed that the recovered wzyB mutants were indeed 294 sensitive to diverse membrane stresses, innate immune defenses and common 295 environmental phages that would be encountered during transmission or on 296 infection of a new host. Therefore, vaccination can successfully drive evolution 297 toward fitness trade-off in vivo.

These observations revealed the overlap between host IgA driven- and phagedriven *Salmonella* evolution. Both the *oafA* gene and the *gtrABC* operon are found at bacteriophage remnant loci, indicating that *S*.Tm has co-opted functions modulating sensitivity to bacteriophage attack in order to escape adaptive immunity. Of note, this example of "coincidental evolution"^{31,32} could

be also driven by and influence how *Salmonella* escape protozoa predator
 grazing in the gut³³. As protozoa are specifically excluded from our SPF mouse
 colonies, this effect could not be investigated here.

Our data, along with previous work on O:4[5] and O:12 variation^{4,5,9,10}, clearly 306 307 indicated direct selective pressure of the host immune system for within-host 308 evolution/phase variation of the O-antigen. Nevertheless, IgA specificity is only 309 one of many strong selective pressures that can be present in the intestine of a free-living animal. Previous work^{20,32-34} indicates that inflammation, phage 310 and predation by protozoa can all contribute, and may exhibit complex 311 312 interactions. For example, inflammation induces the lytic cycle of a temperate 313 phage: a phenomenon inhibited by IgA-mediated protection from disease²⁰. 314 Inflammation is also expected to be particularly detrimental to O-antigen-315 deficient strains that are poorly resistant to antimicrobial peptides and bile acids³ (Fig. ED9). Aggregation of Salmonella by IgA may also generate 316 317 particles that are too large for protozoal grazing, further interacting with 318 bacterial predation in the gut, although this hypothesis has not been 319 experimentally tested. We hope that our work has generated a framework and 320 a set of tools that can be applied to better understand the influence of intestinal 321 adaptive immunity on within-host evolution of bacteria more comprehensively, 322 and that eventually this can be translated into better control of enteric 323 pathogens. In our case, we observed that a tailored adaptive immune response can influence the evolution of bacteriophage/bacteria interactions to the 324 325 detriment of the bacteria.

We have focused on one particular *S*.Tm strain here and it remains to be seen how far this concept can be extended. Further phage-encoded modification of

328 the O-antigen, such as the O:12-1 modification⁴ will likely be required to make robust "evolutionary traps" for Salmonella Typhimurium "in the wild". 329 330 Additionally, species capable of producing capsular polysaccharides that mask 331 the O-antigen, such as Salmonella Typhi and many E.coli strains, would require 332 additional vaccine components (typically glycoconjugates) able to induce 333 robust anti-capsule immunity. However, we expect the principle uncovered here, i.e. understanding the rapid within-host evolution of bacterial surface 334 335 structures and using this information to rationally design oligomeric vaccines, 336 to be broadly applicable. Correspondingly, our findings are consistent with 337 earlier reports of IgA-mediated selection of surface glycans in diverse 338 species^{14,35}, and an earlier report that *gtrABC*-mediated O-antigen phase-339 variation of Salmonella Typhimurium ATCC 14028 confers a colonization 340 benefit starting at day 10 post-infection (roughly the time when an IgA response 341 would be first detected)⁵. Surface variation of teichoic acids for immune evasion can also be prophage-driven in *Staphylococcus aureus*³⁶, although adaption of 342 343 antibody-based techniques for gram-positive pathogens that are masters of 344 immune evasion will likely be beyond the limits of this approach.

345 "Evolutionary trapping" of Salmonella by vaccine-induced IgA does not require 346 any effect of IgA on the intrinsic mutation rate or phase-switching rates of 347 Salmonella. Rather within-host evolution is the product of specific selective pressures (driven by IgA) on mutants and phase variants with changes in O-348 antigen structure, which are spontaneously generated at relatively high 349 350 frequencies in the course of any intestinal infection. This genetic plasticity of 351 large populations of microbes has always been the "Achilles heel" of antibiotic³⁷, phage³⁸ or CRISPR-based³⁹ treatments, leading to resistance and treatment 352

353	failure. In the complex ecological setting of the intestine, where bacterial
354	populations are large and relatively fast-growing, within-host evolution can be
355	rapid, and surprisingly predictable. Via rationally designed oral vaccines, we
356	demonstrate that this force can be harnessed to weaken pathogenicity and to
357	alter bacterial susceptibility to predation. We therefore propose that
358	understanding the most common within-host evolutionary trajectories of gut
359	pathogens holds the key to developing robust prophylactics and therapies.

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569 Author contributions

570 MD, WDH and ES designed the project and wrote the paper. MD and ES 571 designed and carried out experiments relating to vaccination and infection of mice, re-isolation of S.Tm clones, phenotyping of S.Tm clones by flow 572 573 cytometry and gel electrophoresis, characterization of human monoclonal 574 antibodies, analysis of antibody titres, and analysis of fitness of O-antigen 575 variants of S.Tm in vitro and in vivo. MvdW, BHM, CL, RM contributed to 576 experimental design / data interpretation. GZ carried out HR-MAS NMR 577 analysis, OH carried out proton NMR analysis. MA generated the mathematical 578 model for O:12 switching. JA carried out and analysed all AFM imaging. AR, NAB carried out phage-sensitivity assays. AE, FB, DW carried out Illumina 579 580 whole-genome resequencing of re-isolated S.Tm isolates. EB, VL, DH, FB, 581 KSM, SA carried out S.Tm challenge infections in vaccinated mice and analysed re-isolated clones. AH carried out microfluidic video microscopy of 582 583 O:12 switching. PV and LF carried out methylome analysis of re-isolated S.Tm 584 clones. LP, AL and BMS generated novel antibody reagents. All authors 585 critically reviewed the manuscript.

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601 **Competing Interests Statement**

M.D. W-D.H. and E.S. declare that Evolutionary Trap Vaccines are covered by
European patent application EP19177251. No other authors declare any
competing interests.

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607 Figures



Figure 1: Vaccine-induced IgA exerts a strong selective pressure on O-antigen variants during murine non-Typhoidal Salmonellosis

609 Figure 1: Vaccine-induced IgA exerts a strong selective pressure on Oantigen variants during murine non-Typhoidal Salmonellosis: A. 610 Schematic of the O-antigen of S.Tm (O:4[5],12), and its common variants 611 612 depicted using the "Symbol Nomenclature for Glycans". B and C. Overnight 613 cultures of the indicated S.Tm strains were stained for presence of O:5 (B) or O:12-0 (C) epitopes. (D-I) Naïve and vaccinated C57BL/6 mice were 614 615 streptomycin-pretreated and infected with the indicated combination of S.Tm strains. (D,F,G) Naïve (closed circles, n=5), PA-S.Tm^{ΔgtrC}-vaccinated (O:4[5]-616 vaccinated, open circles, n=5) and PA-S.Tm^{ΔgtrCΔoafA}-vaccinated (O:4-617 vaccinated, open squares, n=5) SPF mice were streptomycin-pretreated, 618 infected (10⁵ CFU, 1:1 ratio of S.Tm^{$\Delta gtrC$} and S.Tm^{$\Delta gtrC$} $\Delta oafA$ per os). **D**. 619 Competitive index (CFU S.Tm $\Delta gtrC$ /CFU S.Tm $\Delta gtrC \Delta oafA$) in feces at the indicated 620 time-points. Two-way ANOVA with Bonferroni post-tests on log-normalized 621 values, compared to naive mice. *1p=0.0443, *2p=0.0257, *1p=0.0477, 622 **p=0.0021,***p=0.0009 F and G. Correlation of the competitive index with the 623 O:4[5]-binding (**F**) and O:4-binding (**G**) intestinal IgA titre, r² values of the linear 624 625 regression of log-normalized values. Open circles: Intestinal IgA from O:4[5]vaccinated mice. Open squares: Intestinal IqA from O:4-vaccinated mice. Lines 626 indicate the best fit with 95% confidence interval. E.H. I. Naive (closed circles, 627 n=5) or PA-S.Tm ^{ΔoafA ΔgtrC} -vaccinated (O:4/O:12-0-vaccinated, open circles 628 629 and red circles, n=10) C57BL/6 mice were streptomycin-pretreated and infected (10⁵ CFU, 1:1 ratio of S.Tm^{$\Delta oafA$} (O:12-2 switching) and S.Tm^{$\Delta oafA \Delta gtrC$} (O:12-630 locked) per os). E. Competitive index (CFU S.Tm^{ΔoafA} ΔgtrC /CFU S.Tm^{ΔoafA}) in 631 feces at the indicated time-points. Red circles indicate vaccinated mice with a 632 competitive index below 10⁻² on d4 and are used to identify these animals in 633 634 panels H and I. E Effect of vaccination is not significant by 2-way ANOVA 635 considering vaccination over time. H. Correlation of the competitive index on 636 day 4 with the ratio of intestinal IgA titre against an O:12-2-locked S.Tm pgtrABC variant to the titre again an O:12-0-locked S.Tm^{GtrC} variant (linear 637 regression of log-normalized values, lines indicate the best fit with 95% 638 639 confidence interval). I. Intestinal inflammation, corresponding to mice in panel 640 E, quantified by measuring Fecal Lipocalin 2 (LCN2).

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Figure 2: O-Antigen variants rapidly emerge during wild type S.Tm infection of vaccinated mice

643 Figure 2: O-Antigen variants rapidly emerge during wild type S.Tm 644 infection of vaccinated mice: A-C : Naïve (n=22) or PA-S.Tm-vaccinated (Vaccinated, n=23) SPF C57BL/6 mice were streptomycin-pretreated, infected 645 646 (10⁵ S.Tm^{wt} Colony forming units (CFU) per os) and analyzed 18 h later. **A**. Fecal Lipocalin 2 (LCN2) to quantify intestinal inflammation, 2-tailed Mann 647 Whitney U test p<0.0001 **B**. Intestinal IgA titres against S.Tm^{wt} determined by 648 649 flow cytometry, for vaccinated mice with LCN2 values below (open symbols, 650 protected) and above (filled symbols, inflamed) 100ng/g. p=0.61 by 2-tailed Mann Whitney U test. C. Titres of intestinal lavage IgA from an "inflamed 651 652 vaccinated" mouse (red borders) or a "protected vaccinated" mouse (black 653 borders) against S.Tm clones re-isolated from the feces of the "inflamed 654 vaccinated" mouse (red filled circles) or "protected vaccinated" mouse (open circles) at day 3 post-infection. Two-way ANOVA with Bonferroni post-tests on 655 656 log-normalized data. Clones and lavages from n=1 mouse, representative of 9 657 "vaccinated but inflamed" and 13 "vaccinated protected" mice, summarized in 658 Table S4. *p=0.0156, ***p=0.0003. **D.** Flow cytometry staining of S.Tm^{wt} and 659 an evolved with anti-O:5 typing sera (gating as in Fig. S1). E. Alignment of the oafA sequence from wild type (SL1344 RS11465) and an example O:5-660 negative evolved clone showing the 7bp contraction leading to premature stop 661 codon (all four re-sequenced O:5-negative strains showed the same deletion). 662 663 **F**. Binding of an O:12-0-specific monoclonal antibody to S.Tm^{wt} and O:12^{Bimodal} evolved clones, determined by bacterial flow cytometry. (gating as in Fig. S1). 664 665 **G**. Methylation status of the gtrABC promoter region in S.Tm, and three O:12^{Bimodal} evolved clones determined by REC-seq. Heat-scale for normalized 666 read-counts, schematic diagram of promoter methylation associated with ON 667 668 and OFF phenotypes, and normalized methylation read counts for the indicated 669 strains. 670



Figure 3: Single-repeat O-antigen confers a selective advantage in the presence of broad-specificity vaccine-induced IgA

672 Figure 3: Single-repeat O-antigen confers a selective advantage in the presence of broad-specificity vaccine-induced IgA: A-C. Mock-vaccinated 673 wild type (C57BL/6, n=10), PA-S.Tm^{$\Delta oafA \Delta gtrC$} -vaccinated JH^{-/-} mice (JH^{-/-}, n=6), PA-S.Tm^{$\Delta oafA \Delta gtrC$} -vaccinated wild type (C57BL/6,-n=16) and PA-S.Tm^{$\Delta oafA \Delta gtrC$} 674 675 -vaccinated JH+/- littermate controls (JH+/-, n=5 mice) were streptomycin pre-676 treated and infected with 10^5 CFU of a 1:1 ratio S.Tm $\Delta oafA \Delta gtrC \Delta wzyB$ and 677 S.Tm^{\LoafA \LogfTC} i.e. serotype-locked, short and long O-antigen-producing strains. 678 679 A. Competitive index of S.Tm in feces on the indicated days. 2-way ANOVA with Tukey's multiple comparisons tests. *p=0.0392, ****p<0.0001. B. Feces 680 681 from the indicated mice (grey-filled circles panel A) were transferred into 682 streptomycin-pretreated C57BL/6 naive mice (one fecal pellet per mouse, n=5). Competitive index in feces over 2 days of infection. C. Intestinal IgA titre from 683 PA-S.Tm^{$\Delta oafA \Delta gtrC$} -vaccinated mice binding to S.Tm $^{\Delta oafA \Delta gtrC}$ (long O-antigen) 684 and S.Tm^{ΔoafA ΔgtrC ΔwzyB} (short O-antigen). *p=0.0078 by 2-tailed Wilcoxon 685 matched-pairs signed rank test. **D.** Intestinal IgA titre induced by PA-S.Tm^{wt} or 686 PA-S.Tm^{ET} (4-strains) in 129S1/SvImJ mice determined by bacterial flow 687 688 cytometry. Two-way ANOVA with Bonferroni multiple comparisons tests. Adjusted p values *p=0.0332, ***p=0007. (Gating Fig.S5, further data Fig. S7 689 690 and S8) E. 129S1/SvImJ Mice were vaccinated with vehicle only (Naïve, n=8), PA-S.Tm^{wt} (n=8), PA-S.Tm^{ET} (n=8). On day 28 after the first vaccination, mice 691 692 were streptomycin pre-treated and challenged with 10⁵ S.Tm^{wt} orally. Intestinal inflammation as scored by fecal Lipocalin-2 (LCN2) days 1-9 post-infection. 693 694 Dotted line = detection limit. Grey box = normal range in healthy mice. 2-way 695 repeat-measures ANOVA with Tukey's multiple comparison test. *** adjusted p 696 value=0.0002 F. Representative plot of O:5 staining in an evolved clone with 697 short O-antigen and quantification of the percentage of O:5-dim S.Tm clones re-isolated from the feces of infected SPF mice vaccinated with PBS only 698 (n=13), PA-S.Tm^{Δ gtrC} (n=9) or PA-S.Tm^{ET} (n=18). Kruskal-Wallis test with 699 Dunn's multiple comparison tests shown. **p=0.0016. (gating as Fig. S1) G. 700 Silver-stained gel of LPS from representative control and evolved S.Tm strains 701 from 2 different control and vaccinated PA-S.Tm^{ET} mice. H. Resequencing of 702 703 short O-antigen strains revealed a deletion between inverted repeats (n=5 704 clones, isolated from 2 different mice).



Figure 4: Single-repeat O-antigen mutants arising during infection of vaccinated mice have attenuated virulence, fitness and diminished resistance to phage predation

Figure 4: Single-repeat O-antigen mutants arising during infection of vaccinated mice have attenuated virulence, fitness and diminished resistance to phage predation.

710 A, B, C, Single 24h infections in streptomycin pretreated naïve C57BL/6 mice (n=14, short O-antigen, n=9 long O-antigen). Evolved and synthetic wzyB 711 mutants have reduced ability to colonize the gut (A, CFU/g feces, ***p=0.0002) 712 713 and to spread systemically (B, CFU per mesenteric lymph node (MLN), 714 ***p=0.0001). This translates into diminished propensity to trigger intestinal 715 inflammation in comparison to isogenic wild type strains (C, fecal Lipocalin 2 716 (LCN2), ****p<0.0001). Mann-Whitney U, 2-tailed tests. D. Phage plaques on a 717 lawn of ancestor S. Tm^{wt} (left) and evolved S.Tm^{$\Delta wzyB$} (right) after infection with filtered wastewater; scale=1cm. E. Quantification of the plaques from three 718 719 independent experiments (2-tailed Paired T test **p=0.0046). F. Pairwise 720 comparison matrix of de novo assembled and aligned genomes of isolated 721 bacteriophages (ϕ 12, ϕ 23, ϕ 34, ϕ 37) and a reference sequence from Enterobacteriaceae phage T5 (NC 005859). Values indicate the alignment 722 723 percentage (comparisons below diagonal) between genomes and the average nucleotide identity between the aligned parts (comparisons above diagonal, 724 green frame). This analysis shows that the four isolated bacteriophages are 725 726 different but all belong to the T5 family. G. Quantification of phage plaques 727 formed on infection of the ancestor S.Tm^{wt} (long O-antigen) and evolved S.Tm^{$\Delta wzyB$} (short O-antigen) with the isolated phage φ 12. 2-tailed Mann-728 729 Whitney U test. **p=0.0041.

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731 Materials and Methods

732 **Ethics statement**

All animal experiments were approved by the legal authorities (licenses 223/2010,
222/2013, 193/2016, 120/2019; Kantonales Veterinäramt Zürich, Switzerland). All
experiments involving animals were carried out strictly in accordance with the legal
framework and ethical guidelines.

737

738 Mice

739 Unless otherwise stated, all experiments used specific opportunistic pathogen-free 740 (SPF, containing a complete microbiota free of an extended list of opportunistic pathogens) C57BL/6 mice. IgA^{-/-40}, Balb/c, J_H^{-/-41}, Rag1^{-/-42} (all C57BL/6 background) 741 and 129S1/SvImJ, mice, were re-derived into a specific pathogen-free (SPF) foster 742 colony to normalize the microbiota and bred under full barrier conditions in 743 744 individually ventilated cages in the ETH Phenomics Center (EPIC, RCHCI), ETH 745 Zürich and were fed a standard chow diet. Low complex microbiota (LCM) mice (IgA+/- and -/-, used in Fig. ED2) are ex-germfree mice, which were colonized with a 746 naturally diversified Altered Schaedler flora in 2007¹⁴ and were bred in individually 747 748 ventilated cages or flexible-film isolators at this facility, and received identical diet. All mouse facilities were regulated to maintain constant temperature (22°C +/- 1°C) and 749 humidity (30-50%), with a 12h/12h standard dark/light cycle. Male and female mice 750

751 were included in all experimental groups, and the number of animals per group is 752 indicated in each figure legend.

753

Vaccinations and chronic infections with attenuated *Salmonella* strains in naïve mice were started between 5 and 6 weeks of age, and males and females were randomized between groups to obtain identical ratios wherever possible. Challenge infections with virulent *Salmonella* were carried out between 9 and 12 weeks of age. As strong phenotypes were expected, we adhered to standard practice of analysing at least 5 mice per group. Researchers were not blinded to group allocation.

760

761 Strains and plasmids

All strains and plasmids used in this study are listed **Table S1**.

For cultivation of bacteria, we used lysogeny broth (LB) containing appropriate
antibiotics (i.e., 50 µg/ml streptomycin (AppliChem); 6 µg/ml chloramphenicol
(AppliChem); 50 µg/ml kanamycin (AppliChem); 100 µg/ml ampicillin (AppliChem)).
Dilutions were prepared in Phosphate Buffer Saline (PBS, Difco).

- 767 In-frame deletion mutants (e.g. *gtrC::cat*) were performed by λ *red* recombination as described in⁴³. When needed, antibiotic resistance cassettes were removed using the 768 769 temperature-inducible FLP recombinase encoded on pCP20⁴³. Mutations coupled with 770 antibiotic resistance cassettes were transferred into the relevant genetic background by generalized transduction with bacteriophage P22 HT105/1 int-20144. Primers used for 771 772 genetic manipulations and verifications of the constructions are listed Table S2. 773 Deletions of gtrA and gtrC originated from in-frame deletions made in S.Tm 14028S, 774 kind gifts from Prof. Michael McClelland (University of California, Irvine), and were 775 transduced into the SB300 genetic background.
- 776

The gtrABC operon (STM0557-0559) was cloned into the pSC101 derivative plasmid 777 pM965⁴⁵ for constitutive expression. The operon gtrABC was amplified from the 778 779 chromosome of SB300 using the Phusion Polymerase (ThermoFisher Scientific) and 780 primers listed Table S2. The PCR product and pM965 were digested with PstI-HF and 781 EcoRV-HF (NEB) before kit purification (SV Gel and PCR Clean up System, Promega) 782 and ligation in presence of T4 ligase (NEB) following manufacturer recommendations. 783 The ligation product was transferred by electro-transformation in competent SB300 784 cells.

785

786 Targeted sequencing

Targeted re-sequencing by the Sanger method (Microsynth AG) was performed on kit
purified PCR products (Promega) from chromosomal DNA or expression vector
templates using pre-mixed sequencing primers listed Table S2.

790

791 Whole-genome re-sequencing of O:12^{Bimodal} isolates

The genomes of S.Tm and evolved derivatives were fully sequenced by the Miseq system (2x300bp reads, Illumina, San Diego, CA) operated at the Functional Genomic Center in Zürich. The sequence of *S*.Tm SL1344 (NC 016810.1) was used as reference.

Quality check, reads trimming, alignments, SNPs and indels calling were performedusing the bioinformatics software CLC Workbench (Qiagen).

797

Whole-genome sequencing of S.Tm isolates from "Evolutionary Trap" vaccinated mice and variant calling.

800 Nextera XT libraries were prepared for each of the samples. The barcoded libraries 801 were pooled into equimolar concentrations following manufacturer's guidelines (Illumina, San Diego, CA) using the Mid-Output Kit for paired-end sequencing (2×150 802 bp) on an Illumina NextSeq500 sequencing platform. Raw data (mean virtual coverage 803 804 361x) was demultiplexed and subsequently clipped of adapters using Trimmomatic v0.38 with default parameters⁴⁶. Quality control passing read-pairs were aligned against 805 806 reference genome/plasmids (Accession numbers: NC 016810.1, NC 017718.1, NC 017719.1, NC 017720.1) with bwa v0.7.17⁴⁷. Genomic variant were called using 807 808 Pilon v1.23⁴⁸. with the following parameters: (i) minimum coverage 10x; (ii) minimum 809 quality score = 20; (iii) minimum read mapping quality = 10. SnpEff v4.3 was used to annotate variants according to NCBI and predict their effect on genes⁴⁹. 810

811

812 **PA-STm vaccinations**

813 Peracetic acid killed vaccines were produced as previously described²⁸. Briefly, 814 bacteria were grown overnight to late stationary phase, harvested by centrifugation and re-suspended to a density of 109-1010 per ml in sterile PBS. Peracetic acid (Sigma-815 816 Aldrich) was added to a final concentration of 0.4% v/v. The suspension was mixed 817 thoroughly and incubated for 60 min at room temperature. Bacteria were washed once in 40 ml of sterile 10x PBS and subsequently three times in 50 ml sterile 1x PBS. The 818 final pellet was re-suspended to yield a density of 10^{11} particles per ml in sterile PBS 819 820 (determined by OD600) and stored at 4°C for up to three weeks. As a quality control, 821 each batch of vaccine was tested before use by inoculating 100 µl of the killed vaccine (one vaccine dose) into 300 ml LB and incubating over night at 37 °C with aeration. 822 823 Vaccine lots were released for use only when a negative enrichment culture had been confirmed. For all vaccination, 10¹⁰ particles, suspended in 100µl PBS were delivered 824 by oral gavage, once weekly for 4 weeks. Where multiple strains were combined, the 825 total number of vaccine particles remained constant, and was roughly equally divided 826 between the constituent strains. Unless otherwise stated, PA-STm vaccinated mice were 827 828 challenged orally on d28 after the first vaccination.

829

830 Adoptive transfer of recombinant mSTA121 IgA

A recombinant monoclonal dimeric murine IgA specific for the O:12-0 epitope
(described in ¹⁵) was buffer-exchanged into sterile PBS. 1 mg of antibody was injected
intravenously into mice 30 min prior to infection and again 12 h post-infection, to
maintain sufficient dimeric IgA for export into the gut by PIgR.

835

836 Chronic infection with live-attenuated vaccine strains of non-typhoidal Salmonella

- 837 6-week-old mice were orally pretreated 24 h before infection with 25 mg streptomycin.
- 838 Live-attenuated strains (sseD::aphT, $\Delta gtrC \Delta aroA$ and $\Delta oafA \Delta gtrC \Delta aroA$, Table S1,

- ⁵⁰) were cultivated overnight separately in LB containing streptomycin. Subcultures were prepared before infections by diluting overnight cultures 1:20 in fresh LB without antibiotics and incubation for 4 h at 37°C. The cells were washed in PBS, diluted, and 50 μ l of resuspended pellets were used to infect mice *per os* (5x10⁷ CFU).
- Feces were sampled at day 1, 9 and 42 post-infection, homogenized in 1 ml PBS by bead beating (3mm steel ball, 25 Hz for 1 minute in a TissueLyser (Qiagen)), and *S*.Tm strains were enumerated by selective plating on MacConkey agar supplemented with streptomycin. Samples for lipocalin-2 measurements were kept homogenized in PBS at -20 °C. Enrichment cultures for analysis of O-antigen composition were carried out by inoculating 2 μ l of fecal slurry into 5ml of fresh LB media and cultivating overnight at 37 °C.
- 850

851 Non-typhoidal *Salmonella* challenge infections

Infections were carried out as previously described ²³. In order to allow reproducible
gut colonization, 8-12 week-old SPF mice, naïve or PA-STm vaccinated, were orally
pretreated 24 h before infection with 25 mg streptomycin or 20 mg of ampicillin. Strains
were cultivated overnight separately in LB containing the appropriate antibiotics.
Subcultures were prepared before infections by diluting overnight cultures 1:20 in fresh
LB without antibiotics and incubation for 4 h at 37°C. The cells were washed in PBS,

858 diluted, and 50 μ l of resuspended pellets were used to infect mice *per os* (5×10⁵ CFU). 859 Competitions were performed by inoculating 1:1 mixtures of each competitor strain.

Feces were sampled daily, homogenized in 1 ml PBS by bead beating (3 mm steel ball,

861 25 Hz for 1 min in a TissueLyser (Qiagen)), and S.Tm strains were enumerated by 862 selective plating on MacConkey agar supplemented with the relevant antibiotics. Fecal 863 samples for lipocalin-2 measurements were kept homogenized in PBS at -20°C. At 864 endpoint, intestinal lavages were harvested by flushing the ileum content with 2 ml of 865 PBS using a cannula. The mesenteric lymph nodes, were collected, homogenized in 866 PBS Tergitol 0.05% v/v at 25 Hz for 2 min, and bacteria were enumerated by selective 867 plating.

868 Competitive indexes were calculated as the ratio of population sizes of each genotype, 869 enumerated by selective plating of the two different strains on kanamycin- and 870 chloramphenicol-containing agar, at a given time point, normalized for the ratio 871 determined by selective plating in the inoculum (which was always between 0.5 and 2).

872

873 Non-typhoidal Salmonella transmission

Donor mice were vaccinated with PA-S.Tm^{$\Delta oafA \Delta gtrC$} once per week for 5 weeks, 874 875 streptomycin pretreated (25 mg streptomycin per os), and gavaged 24 h later with 10⁵ CFU of a 1:1 mixture of S. $\text{Tm}^{\Delta oafA \Delta gtrCwzyB::cat}$ (Cm^R) and S. $\text{Tm}^{\Delta oafA \Delta gtrC \text{ Kan}}$ (Kan^R). On 876 day 4 post infection, the donor mice were euthanized, organs were harvested, and fecal 877 878 pellets were collected, weighed and homogenized in 1 ml of PBS. The re-suspended 879 feces (centrifuged for 10 s to discard large debris) were immediately used to gavage (as 880 a 50 µl volume containing the bacteria from on fecal pellet) recipient naïve mice 881 (pretreated with 25 mg streptomycin 24 hours before infection). Recipient mice were 882 euthanized and organs were collected on day 2 post transmission. In both donor and recipient mice, fecal pellets were collected daily and selective plating was used to enumerate *Salmonella* and determine the relative proportions (and consequently the competitive index) of both competing bacterial strains.

886

887 Quantification of fecal Lipocalin2

Fecal pellets collected at the indicated time-points were homogenized in PBS by beadbeating at 25 Hz, 1min. Large particles were sedimented by centrifugation at 300 g, 1
min. The resulting supernatant was then analysed in serial dilution using the mouse
Lipocalin2 ELISA duoset (R&D) according to the manufacturer's instructions.

892

893 Analysis of specific antibody titres by bacterial flow cytometry

894 Specific antibody titres in mouse intestinal washes were measured by flow cytometry as described^{15,51}. Briefly, intestinal washes were collected by flushing the small 895 896 intestine with 2 ml PBS, centrifuged at 16000 g for 30 min to clear all bacterial-sized 897 particles. Aliquots of the supernatants were stored at -20°C until analysis. Bacterial targets (antigen against which antibodies are to be titred) were grown to late stationary 898 899 phase or the required OD in 0.2µm-filtered LB, then gently pelleted for 2 min at 7000 g. The pellet was washed with 0.2µm-filtered 1% BSA/PBS before re-suspending at a 900 901 density of approximately 10⁷ bacteria per ml. After thawing, intestinal washes were centrifuged again at 16000 g for 10 min to clear. Supernatants were used to perform 902 serial dilutions. 25 µl of the dilutions were incubated with 25 µl bacterial suspension at 903 4°C for 1 h. Bacteria were washed twice with 200 µl 1% BSA/PBS by centrifugation at 904 905 7000g for 15 min, before resuspending in 25 µl of 0.2µm-filtered 1% BSA/PBS containing monoclonal FITC-anti-mouse IgA (BD Pharmingen, 10 µg/ml) or Brilliant 906 907 violet 421-anti-IgA (BD Pharmingen, 10µg/ml). After 1 h of incubation, bacteria were 908 washed once with 1% BSA/PBS as above and resuspended in 300 µl 1% BSA/PBS for 909 acquisition on LSRII or Beckman Coulter Cytoflex S using FSC and SSC parameters to threshold acquisition in logarithmic mode. Data were analysed using FloJo 910 911 (Treestar). After gating on bacterial particles, log-median fluorescence intensities 912 (MFI) were plotted against lavage dilution factor for each sample and 4-parameter 913 logistic curves were fitted using Prism (Graphpad, USA). Titers were calculated from these curves as the dilution factor giving an above-background signal (typically IgA 914 915 coating MFI=1000 – e.g. Fig. S7 and S8).

916

917 Dirty-plate ELISA analysis of intestinal lavage IgA titres specific for S.Tm. 918 Bacterial targets (antigen against which antibodies are to be titred) were grown to late 919 stationary phase in 0.2µm-filtered LB, then gently pelleted for 2 min at 7000 g. The 920 pellet was washed with 0.2µm-filtered 1% BSA/PBS before re-suspending at a density of approximately 10⁹ bacteria per ml in sterile PBS. 50µl of this bacterial suspension 921 922 was added to each well of a Nunc Immunosorb ELISA plate and was incubated 923 overnight at 4°C in a humidified chamber. The ELISA plates were then washed 3 times with PBS/0.5% Tween-20 and blocked with 200µl per well of 2% BSA in PBS for 3h. 924 925 After thawing, intestinal washes were centrifuged again at 16000 g for 10 min to clear. 926 Supernatants were used to perform serial dilutions. 50 µl of the dilutions were added to

927 each well and the plates were incubated at 4°C overnight in a humidified chamber. The 928 next morning, the plates were washed 5 times with PBS/0.5% Tween-20 and 50µl of 929 HRP-anti-mouse-IgA (Sigma-Aldrich, 1:1000) was added to each well. This was incubated for 1h at room temperature before washing again 5 times and developing the 930 931 plates with 100µl per well of ABTS ELISA substrate. Absorbance at 405nm was read 932 using a Tecan Infinite pro 200. A₄₀₅ readings were plotted against lavage dilution factor 933 for each sample and 4-parameter logistic curves were fitted using Prism (Graphpad, 934 USA). Titers were calculated from these curves as the dilution factor giving an above-935 background signal ($A_{405}=0.2 - e.g.$ Fig. S7 and S8).

936

Flow cytometry for analysis of O:5, O:4 and O:12-0 epitope abundance on Salmonella in cecal content, enrichment cultures and clonal cultures

939 1 µl of overnight cultures made in 0.2µm-filtered LB, or 1µl of fresh feces or cecal 940 content suspension (as above) was stained with 0.2µm-filtered solutions of STA5 941 (human recombinant monoclonal IgG2 anti-O:12-0, 6µg/ml¹⁵), Rabbit anti-Salmonella O:5 (Difco, 1:200) or Rabbit anti-Salmonella O:4 (Difco, 1:5). After incubation at 4°C 942 943 for 30 min, bacteria were washed twice by centrifugation at 7000g and resuspension in 944 PBS/1% BSA. Bacteria were then resuspended in 0.2µm-filtered solutions of appropriate secondary reagents (Alexa 647-anti-human IgG, Jackson Immunoresearch 945 946 1:200, Brilliant Violet 421-anti-Rabbit IgG, Biolegend 1:200). This was incubated for 947 10-60 min before cells were washed as above and resuspended for acquisition on a BD LSRII or Beckman Coulter Cytoflex S. A media-only sample was run on identical 948 949 settings to ensure that the flow cytometer was sufficiently clean to identify bacteria 950 without the need for DNA dyes. Median fluorescence intensity corresponding to O:12-951 0 or O:5 staining was calculated using FlowJo (Treestar, USA). Gates used to calculate 952 the % of "ON" and "OFF" cells were set by gating on samples with known O:5/O:4 953 (oafA-deletion) and O:12-0 (gtrC-deletion) versus O:12-2 (pgtrABC) phenotypes (Fig. 954 S2 and 3).

955

956 Live-cell immunofluorescence

957 200 uL of an overnight culture was centrifuged and resuspended in 200 µL PBS 958 containing 1 µg recombinant murine IgA clone STA121-AlexaFluor568. The cells and 959 antibodies were co-incubated for 20 min at room temperature in the dark and then 960 washed twice in 1 mL Lysogeny broth (LB). Antibody-labeled cells were pipetted into an in-house fabricated microfluidic device⁵². Cells in the microfluidic device were 961 continuously fed S.Tm-conditioned LB52 containing STA121-AlexaFluor568 (1 962 963 µg/mL). Media was flowed through the device at a flow rate of 0.2 mL/h using syringe 964 pumps (NE-300, NewEra PumpSystems). Cells in the microfluidic device were imaged 965 on an automated Olympus IX81 microscope enclosed in an incubation chamber heated 966 to 37°C. At least 10 unique positions were monitored in parallel per experiment. Phase contrast and fluorescence images were acquired every 3 min. Images were 967 deconvoluted in MatLab⁵³. Videos are compressed to 7 fps, i.e. 1 s = 21 mins. 968 969

970 HR-MAS NMR

971 S. Typhimurium cells were grown overnight (~18 h) to late stationary phase. The equivalent of 11–15 OD₆₀₀ was pelleted by centrifugation for 10 min 4 °C and 3750 g. 972 The pellet was resuspended in 10% NaN₃ in potassium phosphate buffer (PPB; 10 mM 973 pH 7.4) in D₂O and incubated at room temperature for at least 90 min. The cells were 974 975 then washed twice with PPB and resuspended in PPB to a final concentration of 0.2 976 OD_{600}/μ l in PPB containing acetone (final concentration 0.1% (v/v) as internal 977 reference). The samples were kept on ice until the NMR measurements were performed 978 - i.e. for between 1 and 8 h. The HR-MAS NMR spectra were recorded in two batches, as follows: $S.Tm^{WT}$, $S.Tm^{\Delta wbaP}$, $S.Tm^{Evolved_1}$, $S.Tm^{Evolved_2}$ were measured on 979 16.12.2016, S.Tm^{$\Delta oafA$} was measured on 26.7.2017. 980

981

982 NMR experiments on intact cells were carried out on a Bruker Biospin AVANCE III 983 spectrometer operating at 600 MHz ¹H Larmor frequency using a 4 mm HR-MAS 984 Bruker probe with 50 µl restricted-volume rotors. Spectra were collected at a temperature of 27 °C and a spinning frequency of 3 kHz except for the sample of 985 986 S.Tm^{\LoafA} (25°C, 2 kHz). The ¹H experiments were performed with a 24 ms Carr-987 Purcell-Meiboom-Gill (CPMG) pulse-sequence with rotor synchronous refocusing 988 pulses every two rotor periods before acquisition of the last echo signal to remove broad 989 lines due to solid-like material⁵⁴. The 90° pulse was set to 6.5 µs, the acquisition time 990 was 1.36 s, the spectral width to 20 ppm. The signal of HDO was attenuated using water 991 pre-saturation for 2 s. 400 scans were recorded in a total experimental time of about 30 992 minutes.

993

994 **O-Antigen purification and ¹H-NMR**

The LPS was isolated applying the hot phenol-water method⁵⁵, followed by dialysis 995 996 against distilled water until the phenol scent was gone. Then samples were treated with DNase (1mg/100 mg LPS) plus RNase (2 mg/100 mg LPS) at 37°C for 2 h, followed 997 998 by Proteinase K treatment (1 mg/100 mg LPS) at 60°C for 1 h [all enzymes from Serva, 999 Germany]. Subsequently, samples were dialyzed again for 2 more days, then freeze dried. Such LPS samples were then hydrolyzed with 1% aqueous acetic acid (100°C, 1000 1001 90 min) and ultra-centrifuged for 16 h at 4°C and 150,000 g. Resulting supernatants 1002 (the O-antigens) were dissolved in water and freeze-dried. For further purification, the 1003 crude O-antigen samples were chromatographed on TSK HW-40 eluted with 1004 pyridine/acetic acid/water (10/4/1000, by vol.), then lyophilized. On these samples, 1D 1005 and 2 D (COSY, TOCSY, HSQC, HMBC) ¹H- and ¹³C-NMR spectra were recorded with a Bruker DRX Avance 700 MHz spectrometer (¹H: 700.75 MHz; ¹³C: 176.2 MHz) 1006 1007 as described⁵⁶.

1008

1009 Atomic force microscopy

1010 The indicated *S*.Tm strains were grown to late-log phase, pelleted, washed once with 1011 distilled water to remove salt. A 20 μ l of bacterial solution was deposited onto freshly 1012 cleaved mica, adsorbed for 1 min and dried under a clean airstream. The surface of 1013 bacteria was probed using a Dimension FastScan Bio microscope (Bruker) with Bruker
1014 AFM cantilevers in tapping mode under ambient conditions. The microscope was 1015 covered with an acoustic hood to minimized vibrational noise. AFM images were 1016 analyzed using the Nanoscope Analysis 1.5 software.

1017

1018 Methylation analysis of S.Tm clones

1019 For REC-Seq (restriction enzyme cleavage-sequencing) we followed the same procedure described by Ardissone et al, 2016⁵⁷. In brief, 1 µg of genomic DNA from 1020 each S.Tm was cleaved with MboI, a blocked (5'biotinylated) specific adaptor was 1021 1022 ligated to the ends and the ligated fragments were then sheared to an average size of 1023 150-400 bp (Fasteris SA, Geneva, CH). Illumina adaptors were then ligated to the 1024 sheared ends followed by deep-sequencing using a HiSeq Illumina sequencer, the 50 1025 FastQC bp single end reads were quality controlled with v0.9 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/). To 1026 remove 1027 contaminating sequences, the reads were split according to the MboI consensus motif 1028 (5'-^GATC-3') considered as a barcode sequence using fastx toolkit v0.0.13.2 1029 --bcfile (http://hannonlab.cshl.edu/fastx toolkit/) (fastx barcode splitter.pl barcodelist.txt --bol --exact). A large part of the reads (60%) were rejected and 40% 1030 kept for remapping to the reference genomes with bwa mem⁴⁷ v0.7.15 and samtools⁵⁸ 1031 1032 v0.1.19 to generate a sorted bam file. The bam file was further filtered to remove low 1033 mapping quality reads (keeping AS ≥ 45) and split by orientation (alignmentFlag 0 or 16) with bamtools⁵⁹ v2.4.1. The reads were counted at 5' positions using Bedtools⁶⁰ 1034 1035 v2.26.0 (bedtools genomecov -d -5). Both orientation count files were combined into a 1036 bed file at each identified 5'-GATC-3' motif using PERL script (perl v5.24). The MboI positions in the bed file were associated with the closest gene using bedtools closest⁶⁰ 1037 v2.26.0 and the gff3 file of the reference genomes⁶¹. The final bed file was converted 1038 to an MS Excel sheet. The counts were loaded in RStudio v1.1.44262 with R v3.4.463 1039 and analysed with the DESeq2 v1.18.1 package⁶⁴ comparing the reference strain with 1040 the 3 evolved strains considered as replicates. The counts are analysed by genome 1041 1042 position rather than by gene. The positions are considered significantly differentially 1043 methylated upon an adjusted p-value < 0.05. Of the 2607 GATC positions, only 4 were 1044 found significantly differentially methylated and they are all located in the promoter of the gtrABC operon. 1045

1046 The first step in the reads filtering was to remove contaminant reads missing the GATC 1047 consensus motif (MboI) at the beginning of the sequence. These contaminant reads are 1048 due to random fragmentation of the genomic DNA and not to cuts of the MboI 1049 restriction enzyme. Using fastx barcode splitter.pl v0.0.13.2 about 60% of the 1050 reads were rejected because they did not start with GATC. The rest (40%) was 1051 analyzed further. Random DNA shearing and blunt-ended ligation of adaptors, 1052 combined with sequencing noise at the beginning of reads likely generates this high 1053 fraction of reads missing at GTAC sequence.

1054

1055 *gtrABC* expression analysis by blue/white screening and flow cytometry.

1056 About 200 colonies of *S*.Tm^{gtrABC-lacZ} (strain background 4/74, ⁴) were grown from an 1057 overnight culture on LB agar supplemented with X-gal (0.2 mg/ml, Sigma) in order to select for *gtrABC* ON (blue) and OFF clones (white). These colonies were then picked
to start pure overnight cultures. These cultures were diluted and plated on fresh LB agar
X-gal plate in order to enumerate the proportion of *gtrABC* ON and OFF siblings. The
proportion of O:12/O:12-2 cells was analyzed by flow cytometry.

1062

1063 In vitro growth and competitions to determine wzyB-associated fitness costs

1064 Single or 1:1 mixed LB subcultures were diluted 1000 times in 200 µl of media 1065 distributed in 96 well black side microplates (Costar). Where appropriate, wild type S.Tm carried a plasmid for constitutive expression of GFP. To measure growth and 1066 1067 competitions in stressful conditions that specifically destabilize the outer membrane of S.Tm, a mixture of Tris and EDTA (Sigma) was diluted to final concentration (4 mM 1068 1069 Tris, 0.4 mM EDTA) in LB; Sodium cholate (Sigma) and Sodium Dodecyl Sulfate 1070 (SDS) (Sigma) were used at 2% and 0.05% final concentration respectively. The lid-1071 closed microplates were incubated at 37°C with fast and continuous shaking in a 1072 microplate reader (Synergy H4, BioTek Instruments). The optical density was 1073 measured at 600 nm and the green fluorescence using 491 nm excitation and 512 nm 1074 emission filter wavelengths every 10 minutes for 18 h. Growth in presence of SDS 1075 causes aggregation when cell density reaches OD=0.3-0.4, therefore, it is only possible 1076 to compare the growth curves for about 250 minutes. The outcome of competitions was 1077 determined by calculating mean OD and fluorescence intensity measured during the last 100 min of incubation. OD and fluorescence values were corrected for the baseline 1078 1079 value measured at time 0.

1080

1081 Serum resistance

1082 Overnight LB cultures were washed three times in PBS, OD adjusted to 0.5 and 1083 incubated with anonymized pooled human serum obtained from Unispital Basel (3 vol 1084 of culture for 1 vol of serum) at 37°C for 1 h. Heat inactivated (56°C, 30 min) serum 1085 was used as control treatment. Surviving bacteria were enumerated by plating on non-1086 selective LB agar plates. For this, dilutions were prepared in PBS immediately after 1087 incubation.

1088

1089 Bacteriophage sensitivity tests:

1090 5 ml sewage water (sewage plant inflow treated with 1 % v/v chloroform; Basel Stadt, 1091 Switzerland) were mixed with 500 µl of dense bacterial culture (ancestor wild type *S*. 1092 Tm; evolved short O-antigen wzyB mutant AE860.3, *S*.Tm ^{*dgtrC oafA::cat*}, *S*.Tm ^{*dgtrC doafA*} 1093 ^{*wzyB::cat*}), incubated for 15 minutes at 37 °C. The mixtures were added to 15 ml LB 1094 containing 10 mM CaCl₂, 10 mM MgSO₄ and 0.7 % *w/v* agar, and immediately poured 1095 onto LB agar plates with the appropriate antibiotics.

1096

1097 Sensitivity to isolated phage $\varphi 12$ was quantified by calculating phage titres obtained 1098 after overnight cultures of evolved short O-antigen *wzyB* mutant AE860.3 or ancestor 1099 wild type *S*. Tm in presence of the isolated bacteriophage (MOI=10).

1100

1101 Isolation of bacteriophages and resistant clones:

- 1102 Plaques with different morphologies appearing on $S.Tm^{\Delta gtrC \ \Delta oafA \ wzyB::cat}$ plates were
- 1103 streaked on overlay plates containing $S.\text{Tm}^{AgtrC \,AoafA \,wzyB::cat}$. The resulting plaques were
- 1104 used to inoculate 200 μ l of a *S*.Tm^{Δ gtrC Δ oafA wzyB::cat} culture at OD₆₀₀=0.3 in a 96-well
- 1105 plate and optical density was measured every 10 minutes at 37 °C with shaking in a
- Synergy 2 plate-reader. Well contents after 18 hours of growth were streaked onto LB-Cm plates to isolate bacterial colonies from the regrowing population. Resistance to phage was confirmed by testing for absence of plaque formation in presence of the
- 1109 corresponding phage.
- 1110 The rest of the well contents were cleared by centrifugation and filtered $(0.45 \,\mu\text{m})$ for
- 1111 phage purification. The cleared supernatants were used to inoculate 20 ml of a $S.Tm^{\Delta gtrC}$
- 1112 $\triangle oafA wzyB::cat$ culture at OD₆₀₀=0.3 and subsequently grown at 37 °C for 5 hours. Cell
- 1113 debris was removed by centrifugation, the supernatants cleared by 0.45 μ m filtration 1114 and stored at 4 °C.
- 1115

1116 Phage genome sequencing and analysis:

- Phage DNA was isolated using the Phage DNA Isolation Kit from Norgen Biotek andsequenced at MiGS, Pittsburgh, Pennsylvania, USA. For this, Nextera libraries were
- prepared for each sample and sequenced on an Illumina NextSeq 550 sequencingplatform to generate paired end reads.
- 1121 De novo genome assembly was performed using the De Novo Assembly Algorithm of
- 1122 CLC Genomics Workbench and the resulting high coverage contigs were aligned
- using the Whole Genome Alignment Plug-In to calculate neighbor-joining trees andcorresponding pairwise comparison tables.
- Assembly of the phage genomes resulted in a single contig of 108,227 bp and 114,055 bp for φ 12 and φ 23, respectively (4,928 and 4,495-fold coverage). For φ 34 four separate contigs with more than 3000-fold coverage were identified (81,319, 12,250, 10,937, 5,594 bp), giving a total genome size of more than 100,100 bp, while for φ 37 three contigs with more than 1600-fold coverage (95,133, 14,559, 4,197 bp) gave a total genome size of at least 113,889 bp.
- For comparison, enterobacteria phage T5 has a double-stranded linear DNA genome of1132 121,750 bp.
- 1133

1134 Modeling antigen switching between O12 and O12-2

- The aim of this modeling approach is to test whether a constant switching rate between
 an O12 and an O12-2 antigen expression state can explain the experimentally observed
 bimodal populations.
- 1139 To this end, we formulated a deterministic model of population dynamics of the two 1140 phenotypic states as:
- 1141

1142
$$\frac{dO_{12}}{dt} = (\mu O_{12} - s_{\to 12-2}O_{12} + s_{\to 12}O_{12-2}) * \left(1 - \frac{(O_{12} + O_{12-2})}{K}\right)$$

1143

1144
$$\frac{dO_{12-2}}{dt} = (\mu O_{12-2} + s_{\to 12-2}O_{12} - s_{\to 12}O_{12-2}) * \left(1 - \frac{(O_{12} + O_{12-2})}{K}\right)$$

1146 where O_{12} and O_{12-2} denote the population sizes of the respective antigen variants, μ 1147 denotes the growth rate, which is assumed to be identical for the two variants, *K* the 1148 carrying capacity, and $s_{\rightarrow 12-2}$ and $s_{\rightarrow 12}$ the respective switching rates from O_{12} to 1149 O_{12-2} and from O_{12-2} to O_{12} . Growth, as well as the antigen switching rates, are scaled 1150 with population size in a logistic way, so that all processes come to a halt when carrying 1151 capacity is reached.

1152

We use the model to predict the composition of a population after growth in LB overnight, and therefore set the specific growth rate to $\mu = 2.05h^{-1}$, which corresponds to a doubling time of roughly 20min. The carrying capacity is set to K =10⁹ cells. We ran parameter scans for the switching rates $s_{\rightarrow 12}$ and $s_{\rightarrow 12-2}$, with population compositions that start either with 100% or 0% O_{12} , and measure the composition of the population after 16h of growth (**Fig. S11C**). The initial population size is set to 10^4 cells

1160

1161 Experimentally, we observe that when starting a culture with an O_{12} colony, after overnight growth the culture is composed of around 90% O_{12} and 10% O_{12-2} cells, 1162 whereas starting the culture with O_{12-2} cells yields around 50% O_{12} and 50% O_{12-2} 1163 cells after overnight growth (Fig. S11B). To explain this observation without a change 1164 1165 in switching rates, we would need a combination of values in $s_{\rightarrow 12}$ and $s_{\rightarrow 12-2}$ that 1166 yield the correct population composition for both scenarios. In Fig. S11D, we plot the values of $s_{\rightarrow 12}$ and $s_{\rightarrow 12-2}$ that yield values of 10% O_{12-2} (starting with 0% O_{12-2} , 1167 green dots) and 50% O_{12-2} (starting with 100% O_{12-2} , orange dots). The point clusters intersect at $s_{\rightarrow 12} = 0.144h^{-1}$ and $s_{\rightarrow 12-2} = 0.037h^{-1}$ (as determined by a local linear 1168 1169 regression at the intersection point). 1170

1172 We then used the thus determined switching rates to produce a population growth curve 1173 in a in a deterministic simulation, using the above equations for a cultures starting with 1174 $100\% O_{12-2}$, (Fig. S11E, Left-hand graph) and for a culture starting with $0\% O_{12-2}$ 1175 (Fig. S11E, right-hand graph).

- 1177 These switching rates are consistent with published values ⁴. Our results show that the 1178 observed phenotype distributions can be explained without a change in the rate of 1179 switching between the phenotypes.
- 1180

1176

1171

1181 Data availability

- All Plotted data and associated raw numerical data and calculations for figure 1-4,
 extended data fig. 1-10 and supplementary figures 1-10 is provided in source data tables
 (one per figure, titled accordingly). Uncropped images are provided as supplementary
 files.
- All raw flow cytometry data, ordered by figure, is publically available via the ETH
 research collection doi: 10.3929/ethz-b-000477737
- All Illumina sequencing data data is publically available at NCBI BioProjectAccession: PRJNA720270
- 1190

1191 **Code availability**

- 1192 R code used to generate the figures shown in extended data figure 5 can be freely
- 1193 downloaded from <u>https://github.com/marnoldini/evotrap</u>
- 1194



Extended Data Fig. 1: Surface phenotype of S.Tm mutants

1196 Fig. ED1: Surface phenotype of S.Tm mutants: A-C. Atomic force microscopy phase 1197 images of S.Tm^{wt}, S.Tm^{$\Delta wzyB$} (single-repeat O-antigen), and S.Tm^{$\Delta wbaP$} (rough mutant - no O-antigen) at low magnification (A, uncropped image, scale bar = $1\mu m$) and high 1198 magnification (B and C, scale bar main image = 150nm, scale bar inset = 15nm). 1199 Invaginations in the surface of $S.\text{Tm}^{\Delta wbaP}$ (dark colour, B) show a geometry and size 1200 1201 consistent with outer membrane pores⁶⁵. These are already less clearly visible on the 1202 surface of S.Tm^{$\Delta wzyB$} with a single-repeat O-antigen, and become very difficult to discern in S.Tm^{wt}. One representative image of 3 for each genotype is shown. While 1203 1204 arrows point to features with consistent size and abundance to be exposed outer 1205 membrane porins. C. Fast-Fourier transform of images shown in "B" demonstrating clear regularity on the surface of $S.\text{Tm}^{\Delta wbaP}$, which is progressively lost when short and 1206 1207 long O-antigen is present.

oafA (SL1344_RS11465)	CACACCTGGTCACTATCAGTTGAGTGGCAAT	TATCCTTTATTAGTCATTATAGTTAAAAAATTACGGTTT	CCTGTTGGACTCTCATTATCTGTAATTTTAGCCATGTCACTTGCAATTACACTTATGC(
Wild-type allele	STGTGGACCAGTGATAGTCAACTCACCGTTAAAATATAAAATA	TAGGAAATAATCAGTAATATCAATTTTTTAATGCCAAA	GGACAACCTGAGAGTAATAGACATTAAAATCGGTACAGTGAACGTTAATGTGAATACG(
CP006048.1	H T W S L S V E W Q F Y I L Y P L L V I I V K K L R F P V G L S L S V I L A M S L A I T L M F CACACCTGGTCACTATCAGTTGAGTGGCAATTTTTATAT CACACCTGGTCACTATCAGTTGGGAGGCAATTTTTATAGTCATTATAGTCATTATAGTGGTAAAAATTAGGGTCACCTGGGAGGTCATTATCTGTGAATTTTTAGCCATGTCACTTATAGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGCCATGTCACTTATGTCACTTTTTTTT				
	H T W S L S V E W Q F Y I	L Y 🔤			
В	232 <u>9</u> 725				
B oafA (SL1344_RS11465)		TTGATGTAGTTGATGTAGTTGATGTA	ACAGGTCAATTITAGGTGACGACTITTATAAAAAAGTCTATCATCCATTATC	TAATTTCGTCTTGTGTGGGGCACCTTGGAATTATAGGTAAAAAATGATCTACAAGAAATTCAGACTCGATAT	
B <i>oafA</i> (SL1344_RS11465) Wild-type allele	2329725	TTGATGTAGTTGATGTAGTTGATGTA TTTGATGTAGTTGATGTAGTTGATGTA AACTACATCAACTACATCAACTACAT	ACAGGTCAATTTTAGGTGACGACTTTTATAAAAAAGTCTATCATCCATTATC 	TAATTICGTCTTGTGTGGCACCTTGGAATTATAGGTAAAAAATGATCTACAAGAAATTCAGACTCGATAT 	

Extended Data Fig. 2: Mutations detected in the *oafA* gene sequence among several strains of S. Tm

1209 Fig. ED2: Mutations detected in the *oafA* gene sequence among several strains of

1210 **S.Tm.** A. Aligned fractions of the *oafA* ORF from a natural isolate (from chicken)

1211 presenting the same 7 bp deletion detected in mutants of *S*.Tm SL1344 emerging in

1212 vaccinated mice. S.Tm SL1344 was used a reference⁶⁶. **B**. Aligned *oafA* promoter 1213 sequences from three natural isolates of human origin (stool or cerebrospinal fluid⁶⁷)

- 1214 showing variations in the number of 9 bp direct repeats.
- 1215



Extended Data Fig. 3: Loss of O:12-0-staining is a reversible phenotype dependent on the *gtrABC* locus STM0557-0559

Fig. ED3: Loss of the O:12-0 epitope is a reversible phenotype. A. Wild type and
evolved *S*.Tm clones were picked from LB plates, cultured overnight, phenotypically
characterized by O:12-0 (left panel) and O:5 staining (right panel), plated and re-picked.
This process was repeated over 3 cycles with lines showing the descendants of each

1220 This process was repeated over 3 cycles with lines showing the descendants of each 1221 clone. B and C. Wild type 129S1/SvImJ mice were mock-vaccinated or were vaccinated with PA-S.Tm^{$\Delta oafA \Delta gtrC$} as in Fig. 1. On d28, all mice were pre-treated with 1222 1223 streptomycin, and infected with the indicated strain. B. Feces recovered at day 10 post-1224 infection, was enriched overnight by culture in streptomycin, and stained for O:12-0 1225 (human monoclonal STA5). Fraction O:12-0-low S.Tm was determined by flow 1226 cytometry. Percentage of S.Tm that are O:12-0-negative was quantified over 10 days 1227 and is plotted in panel C. Vaccination selects for S.Tm that have lost the O:12-0 epitope, 1228 only if the *gtrC* gene is intact.

1228



Extended Data Fig. 4. Selective pressure for O:12 phase-variation can be exerted by adoptive transfer of a monoclonal dimeric IgA.

1231 Fig. ED4: Loss of the O:12-0 epitope can be driven by adoptive transfer of O:12-1232 0-specific IgA. C57BL/6 SPF mice received oral streptomycin to deplete the 1233 microbiota 23.5h before an intravenous injection with saline only, or with 1mg of recombinant dimeric murine IgA specific for the O:12-0 epitope (STA121). 0.5 h later 1234 all mice were orally inoculated with $S.Tm^{\Delta oafA \ pM965}$ or $S.Tm^{\Delta oafA \ \Delta G4 \ pM965}$ (lacking 4) 1235 1236 different glucosyl transferases, including gtrC) both carrying pM965 to drive 1237 constitutive GFP production. The adoptive transfer was repeated 12h later and all animals were euthanized at 24h post-infection. A. O:12-0 expression on S.Tm enriched 1238 1239 from cecum content by overnight culture on 1:1000 dilution LB with selective 1240 antibiotics, determined by staining with the monoclonal antibody STA5. Flow cytometry plots shown have been gated on scatter only - see Fig. S1 for example. B. 1241 1242 Quantification of the O:12-0-high fraction of S.Tm from A. C. Individual clones of 1243 S.Tm of the indicated genotype were recovered from the cecal content of mice from A 1244 that had received an adoptive transfer of mSTA121 and individual clones, cultured 1245 overnight in LB were analysed as in A and B for fraction of O:12-0-high cells.



Extended Data Fig. 5: Phase-variation and selection, without a shift in switching rate, underly recovery of O:12-2-producing clones from vaccinated mice

Fig. ED5: Phase-variation and selection, without a shift in switching rate, underly 1247 1248 recovery of O:12-2 producing clones from vaccinated mice. A. Comparison of fractions of O:12-0-positive and O:12-0-negative bacteria (in fact O:12-2) determined 1249 by flow cytometry (gating - see Fig.S1) staining with typing sera and by blue-white 1250 1251 colony counts using a gtrABC-lacZ reporter strain and overnight cultures from 1252 individual clonal colonies. **B-D:** Results of a mathematical model simulating bacterial 1253 growth and antigen switching (see supplementary methods). B. Switching rates from 1254 O:12-0 to O:12-2 and from O:12-2 to O:12-0 were varied computationally, and the 1255 fraction of O:12-2 was plotted after 16 h of growth. Left-hand plot depicts the results 1256 of the deterministic model when starting with 100% O:12-2, right-hand plot depicts the 1257 results when starting with 100% O:12-0. C. depicts only the switching rates that comply 1258 with the experimentally observed antigen ratios after overnight growth (90% O:12-0 1259 when starting with O:12-0, and 50% O:12-0 when starting with O:12-2). Right-hand 1260 plot is a zoomed version showing values for switching rates between 0 - 0.2 h⁻¹ (marked 1261 by a grey rectangle). Dashed lines are linear regressions on the values in this range, and their intersection marks the switching rates used for the stochastic simulation in (D). **D**. 1262 1263 Simulation results of bacterial population growth, when starting with only O:12-2 (lefthand plot) or only O:12-0 (right-hand plot). $\mu = 2.05h^{-1}$ was kept constant in all 1264 1265 simulations; switching rates were kept constant at $s_{->12-0} = 0.144h^{-1}$ and $s_{->12-2} = 0.0365h^{-1}$ ¹; the starting populations were always individuals of the indicated phenotype; carrying 1266 capacity was always $K = 10^9$ cells. Time resolution for the simulations is 0.2*h*. 1267 1268



Extended Data Fig. 6: NMR of purified LPS and HR-MAS 1H-NMR confirms O-antigen structures in evolved clones

1269 Fig. ED6: NMR of purified LPS and HR-MAS ¹H-NMR confirms O-antigen 1270 structures in evolved clones. A. Schematic diagram of expected NMR peaks for each molecular species **B.** HR-MAS ¹H-NMR spectra. Spectra show predicted peak 1271 positions and observed spectra for C1 protons of the O-antigen sugars. C. ¹H NMR of 1272 1273 purified LPS from the indicated strains. Note that non-acetylated abequose can be 1274 observed in wild type strains due to spontaneous deacetylation at low pH in late stationary phase cultures⁵⁴. A gtrA mutant strain is used here to over-represent the O:12-1275 1276 2 O-antigen variant due to loss of regulation⁵. 1277



Extended Data Fig. 7: S.Tm O-antigen variants arise during chronic S.Tm infections, dependent on a specific IgA response. After 35 days of infection, this is weaker than the IgA titres induced by inactivated oral vaccines, but less biased to recognition of O:5.

1278 Fig ED7: S.Tm O-antigen variants arise during chronic S.Tm infections, 1279 dependent on a specific IgA response. IgA^{-/-} (A) and Rag1^{-/-} (B) and heterozygote littermate controls (C57BL/6-background) were pre-treated with streptomycin and 1280 infected with $S.Tm^{\Delta sseD}$ orally. Fecal S.Tm were enriched overnight by culturing a 1281 1282 1:2500 dilution of feces in LB plus kanamycin. These enrichment cultures were then 1283 stained for O:5 and O:12-0 and analysed by flow cytometry (gating as in Fig. S1-4). 1284 The fraction of the population that lost O:5 and O:12-0 antisera staining is shown over 1285 time, as well as the total CFU/g in feces. Both immunocompetent mouse strains show 1286 increased O:5-negative S.Tm in the fecal enrichments from day 14 post-infection: 1287 approximately when we expect to see a robust secretory IgA response developing. 1288 These changes are not observed in Rag1-deficient or IgA-deficient mice. The kinetics 1289 of O:5-loss are likely influenced by development or broader IgA responses as the 1290 chronic infection proceeds. Note that lines joining the points are to permit tracking of 1291 individual animals through the data set, and may not be representative of what occurs 1292 between the measured time-points. C. Titres of intestinal lavage IgA specific for O:4[5] (S.Tm^{wt}, O:4[5], 12-0) and O:4(S.Tm^{\Delta oafA}, O:4,12-0), presented as the dilution of 1293 1294 intestinal lavage required to give an IgA-staining MFI=1000 by bacterial flow 1295 cytometry, and the ratios of these titres. Samples: d28 post-vaccination with PA-STm^{wt} 1296 (n=12) or d35 post-colonization with live-attenuated S.Tm (n=8 S.Tm^{$\Delta aroA$} + n=8 S.Tm^{Δ sseD}), This revealed a weaker, but less biased IgA response in mice infected with 1297 the live-vaccine strain, when compared to that induced by the inactivated oral vaccine. 1298 1299 Results of 2-tailed Mann-Whitney U tests shown.

1300



Extended Data Fig. 8: Schematic diagram of O-antigen synthesis in S.Tm

1302 Fig. ED8: Schematic of *S*. Tm O-antigen synthesis (based on⁶⁸)





Extended Data Fig. 9: Synthetic and natural deletions of *wzyB* reduce the fitness of S.Tm in presence of Tris-EDTA, cholate, SDS and human serum

1304 Fig. ED9: Synthetic and natural deletions of *wzyB* reduce the fitness of S.Tm in 1305 presence of Tris-EDTA, Cholate, SDS and serum complement. The deletion of wzyB does not affect the growth of S.Tm or S.Tm^{$\Delta oafA \ \Delta gtrC$} in LB (No stress) (A) but 1306 impairs growth in presence of Tris-EDTA (B), 2% cholate (C) and 0.05% SDS (D). 1307 Dashed lines represent the range of variations between the n=4 pooled experiments. 1308 1309 (E). Relative fitness of the long versus short O-antigen in the presence of membrane stress as quantified by competitive growth of $S.Tm^{GFP}$ against $S.Tm^{\Delta oafA \, \Delta gtrC}$, $S.Tm^{\Delta oafA}$ 1310 $\Delta gtrC \Delta wzyB$ or an evolved S.Tm $\Delta wzyB$, in LB with or without Tris-EDTA. 2-tailed Mann-1311 Whitney U test. ** p=0.0013 (F) Loss of complement resistance in evolved and 1312 1313 synthetic wzyB mutants revealed by relative CFU recovery after treatment with heat-1314 inactivated and fresh human serum. Mann-Whitney U 2-tailed tests * p=0.0167



С

Clone ID	Phage ID	Position & Variation	Outcome	
b3 (MDBZ0639)	φ12	4370748 G to T	Premature stop codon at position 401 in <i>btuB</i>	
b14 (MDBZ0640)	φ23	4270040 C to T	Premature stop codon at position 165 in <i>btuB</i>	
b25 (MDBZ0641)	φ34	4370040 G to 1		
b28 (MDBZ0642)	φ37	4370312 deleted G	Frame shift leading to premature stop codon at position 258 in the <i>btuB</i> open reading frame	

Extended Data Fig. 10: Analysis of bacteriophages preferentially infecting short O-antigen S.Tm mutants.

1316 Fig. ED10: Analysis of bacteriophages preferentially infecting short O-antigen S.

1317 **Tm mutants. A.** Lysis plaques observed on lawns of \tilde{S} . Tm $\Delta gtrC \Delta oafA$ and S. Tm $\Delta gtrC$ 1318 $\Delta oafA \Delta wzyB$ isogenic mutants exposed to wastewater samples. Scale = 1 cm. This

1319 phenocopies the observation with naturally arising wzyB mutants **B**. Growth curves of

1320 S.Tm $^{AgtrC \, AoafA \, AwzyB}$ exposed to purified bacteriophages from Fig. 4D. The re-growing

1320 S. Tm clones were isolated for sequencing. The mutations identified and their effects

- 1322 are listed in the table below (C), confirming btuB as the most likely exposed outer-
- 1323 membrane receptor for these phages.
- 1324
- 1325

1326	List of Supplementary Materials:				
1327					
1328	Supplementary Table S1-4				
1329	Supplementary Movies 1 and 2				
1330	Supplementary Figures S1-10				
1331	• Source data files for Fig.1-4, Extended Data Fig. 1-10 and				
1332	Supplementary Fig. 1-10				
1333	• Uncropped image files for Fig. 3G, Fig. 4D and extended data Fig. 10A				
1334					

- 1335 Supplementary Materials
- 1336 Contents:
- 1337 Supplementary Tables and Movies
- 1338 Supplementary figures 1-10
- 1339

1340 Supplementary tables and movies

Table S1: Strains and plasmids used in this study^{4,43,45,50,69–71}

1342 **Table S2:** Details of primers used in strain construction, testing and sequencing

Table S3: Details of mutations found in resequenced O12-0 or O12 bimodal evolved
clones studied by REC-Seq as shown in Fig. 2D-G. Numbers indicate the position of
the mutation, numbers in brackets indicates the percentage of reads were the mutation

- 1346 was detected.
- 1347 **Table S4:** Details of experiments where *S*.Tm evolution was tracked, as well as

1348 further information on mice used and on clones analysed.

1349 1350

1351 Supplementary Movies A and B

Visualization of O:12 phase variation using live-cell immunofluorescence. Cells
expressing GFP (green) pre-stained with fluorescently-labeled recombinant murine IgA
specific for the O:12-0 epitope (red) were loaded into a microfluidic chip for time-lapse
microscopy. Cells were fed continuously *S*.Tm-conditioned LB containing
fluorescently-labeled recombinant murine IgA STA121 specific for the O:12-0 epitope.
(A) Loss and (B) gain of antibody reactivity (red staining) was observed, indicative of
O:12 phase variation.

1361 Supplementary Figures



Fig. S1: Difco Rabbit-polyclonal anti-O:5 and human monoclonal STA5 (specific for O:12-0) can be used to distinguish Salmonella with known O-antigen type, and can be distinguished from contaminants without DNA dyes.

1363 1364 Fig. S1: Difco Rabbit-polyclonal anti-O:5 and human monoclonal STA5 (specific 1365 for O:12-0) can be used to distinguish Salmonella with known O-antigen type, and can be distinguished from contaminants without DNA dyes. Overnight cultures of 1366 1367 the indicated S. Typhimurium strains we made in 0.2µm-filtered LB containing 1368 streptomycin (50µg) or Ampicillin (100µg/ml to select for plasmid-maintenance of pgtrABC-containing strains). 1µl of an OD₆₀₀=2 culture was stained in 0.2µm-filtered 1369 1370 PBS/0.05%Azide with 1:200 Rabbit polyclonal anti-O:5 and 6µg/ml STA5. Brilliantviolet-421-Donkey-anti-Rabbit (Biolegend) and Alexa-647-anti-human IgG (Jacksom 1371 1372 Immunolabs) were used at a 1:200 dilution, and SybrGold at 1:10'000 dilution. Samples 1373 were acquired on a Beckmann Coulter Cytoflex-S. A. Full gating is shown for each 1374 sample and the final analysis of O:5 versus O:12-0 staining is shown both for the entire 1375 population gated on Forward- and Side-scatter or only on DNA-dye-positive Salmonella. Note that the live bacteria do not stain uniformly with. SybrGold. B. 1376 1377 Quantification of the O-antigen variant distribution within each strain, when gating on the entire population of the SybrGold-positive fraction reveals no difference in the 1378 1379 analysis when DNA dyes are omitted. A sample of LB cultured overnight and treated 1380 as the samples and acquired for the same length of time as the samples ("Media only") 1381 reveals very little background noise in our flow cytometry analysis. C. Schematic diagram of the O-antigen structures present on bacteria in each quadrant on the scatter 1382 1383 plots.



Fig. S2: Controls for the specificity of Rabbit-polyclonal-anti-O:5 and STA5 staining.

1385 Fig. S2: Controls for the specificity of Rabbit-polyclonal-anti-O:5 and STA5 1386 staining. Overnight cultures of the indicated S. Typhimurium strains we made in 1387 0.2µm-filtered LB containing streptomycin (50µg) or Ampicillin (100µg/ml to select for plasmid-maintenance of pgtrABC-containing strains). 1µl of an OD₆₀₀=2 culture 1388 was stained in 0.2µm-filtered PBS/0.05%Azide with the indicated combinations of 1389 1390 1:200 Rabbit polyclonal anti-O:5, 1:200 Rabbit polyclonal anti-E.coli O:6, 6µg/ml STA5. Brilliant-violet-421-Donkey-anti-Rabbit (Biolegend) and Alexa-647-anti-1391 1392 human IgG (Jacksom Immunolabs) were used at a 1:200 dilution as secondary reagents 1393 in all stainings. Samples were acquired on a Beckmann Coulter Cytoflex-S. A. Samples 1394 were gated on Forward- and Side-scatter as in Fig. S1. This reveals good specificity of 1395 the antibodies with the exception of low level cross-reactivity of the anti-human IgG 1396 for the rabbit polyclonal antibody. However, the background generated by this staining is much lower than the real positive signal and does not alter interpretation of our data. 1397 1398 **B.** Representative histogram overlays of the above stainings indicating antibody 1399 specificity.



Fig. S3: Monoclonal antibody STA5 recognises many O:12-containing *S.enterica* O-antigens

1401 Fig. S3: Characterization of the specificity of STA5 using diverse Salmonella 1402 enterica serovars and recombinant S.Tm strains. A. Recombinant monoclonal STA5 1403 human IgG1 was used to surface stain overnight cultures of the indicated Salmonella 1404 enterica serovars. Bacterial surface binding was detected with a Dylight-647-anti 1405 human IgG secondary antibody and analysed by flow cytometry. STA5 binds to all 1406 serovars that include the O:12 epitope. Top panel shows the pre-gating strategy, which served only to remove events landing on the axes in forward-scatted and side-scatter 1407 1408 measurements 1409



Fig. S4: Bacterial flow cytometry titring of intestinal IgA, corresponding to Fig, 1F and G

1410

1411 Fig. S4: Raw data for intestinal IgA titre calculations shown in Fig. 1F and G (binding to S.Tm $\Delta gtrC$ (O:4[5],12-0) and S.Tm $\Delta oafA \Delta gtrC$ (O:4,12-0). A. Forward- and 1412 side-scatter plot showing gating based on scatter, used for all analysis. B. 1413 1414 Representative overlayed histograms of S.Tm $\Delta gtrC$ stained with intestinal lavage from a 1415 C57BL/6 mouse orally vaccinated once per week for 4 weeks with PA-STm^{$\Delta gtrC$} (left) and PA-STm^{$\Delta oafA \ \Delta gtrC$} (right). BV421-conjugated anti-mouse IgA was used as a 1416 secondary antibody to reveal IgA coating of S.Tm. Colours represent different dilutions 1417 1418 of the intestinal lavages ranging from a dilution factor of 2 (red) to 486 (pink). C. 1419 Intestinal lavage dilution factor plotted against the median fluorescence intensity of IgA 1420 staining (circles: PA-STm^{$\Delta gtrC$}-vaccinated, squares: PA-STm^{$\Delta gtrC$}-vaccinated) for all mice shown in Fig. 1F and G. Lines (black = PA-STm^{$\Delta gtrC$}-vaccinated, orange = PA-1421 $STm^{\Delta oafA \Delta gtrC}$ -vaccinated) indicate 4-parameter logisitic curves fitted to these values 1422 1423 using least-squares non-linear regression. **D**. Titres calculated from the fitted curves as 1424 the intestinal lavage dilution giving a median fluorescence intensity of staining = 1000 for each curve shown in C. Line indicates median value. P value of 2-tailed Mann-1425 Whitney U test. E. Representative overlayed histograms of S.Tm $\Delta oafA \Delta gtrC$ stained with 1426 intestinal lavage from a mouse orally vaccinated once per week for 4 weeks with PA-1427 1428 $STm^{\Delta gtrC}$ (left) and PA-STm^{$\Delta oafA \ \Delta gtrC$} (right). BV421-conjugated anti-mouse IgA was 1429 used as a secondary antibody to reveal IgA coating of S.Tm. Colours represent different 1430 dilutions of the intestinal lavages ranging from a dilution factor of 2 (red) to 486 (pink). 1431 F. Intestinal lavage dilution factor plotted against the median fluorescence intensity of 1432 IgA staining (circles: PA-STm^{$\Delta gtrC$}-vaccinated, squares: PA-STm^{$\Delta gtrC$}-vaccinated) for 1433 all mice shown in Fig. 1F and G. Lines (black = PA-STm^{$\Delta gtrC$}-vaccinated, orange = PA- $STm^{\Delta oafA \Delta gtrC}$ -vaccinated) indicate 4-parameter logisitic curves fitted to these values 1434 1435 using least-squares non-linear regression. G. Titres calculated from the fitted curves as 1436 the intestinal lavage dilution giving a median fluorescence intensity of staining = 1000 for each curve shown in F. Line indicates median value. P value of 2-tailed Mann-1437 1438 Whitney U test. All vaccinated mice were C57BL/6 and had an SPF microbiota. Note 1439 the significantly higher titres of IgA specific for the vaccination strain than the mis-1440 matched strain. 1441



Fig. S5: Bacterial flow cytometry titring of intestinal IgA, corresponding to Fig, 1H
1443 Fig. S5: Raw data for intestinal IgA titre calculations shown in Fig. 1H (binding to S.Tm *DoafA*, pgtrABC (O:4,12-2) and S.Tm^{DoafA DgtrC} (O:4,12-0). A. Forward- and side-1444 scatter plot showing gating based on scatter, used for all analysis. B. Representative 1445 overlayed histograms of S.Tm^{$\Delta oafA \ \Delta gtrC$} stained with intestinal lavage from a mouse 1446 orally vaccinated once per week for 4 weeks with PA-S $Tm^{\Delta oafA \Delta gtrC}$. BV421-1447 1448 conjugated anti-mouse IgA was used as a secondary antibody to reveal IgA coating of 1449 S.Tm. Colours represent different dilutions of the intestinal lavages ranging from a dilution factor of 2 (red) to 486 (pink). C. Intestinal lavage dilution factor plotted 1450 against the median fluorescence intensity of IgA staining for all mice shown in Fig. 1H. 1451 1452 Lines indicate 4-parameter logisitic curves fitted to these values using least-squares 1453 non-linear regression. **D**. Representative overlayed histograms of $S.Tm^{\Delta oafA \ pgtrABC}$ 1454 stained with intestinal lavage from a mouse orally vaccinated once per week for 4 weeks with PA-S Tm^{$\Delta oafA \ \Delta gtrC$}. BV421-conjugated anti-mouse IgA was used as a secondary 1455 1456 antibody to reveal IgA coating of S.Tm. Colours represent different dilutions of the intestinal lavages ranging from a dilution factor of 2 (red) to 486 (pink). F. Intestinal 1457 IgA Titres calculated from the fitted curves as the intestinal lavage dilution giving a 1458 1459 median fluorescence intensity of staining = 1000 for each curve shown in E and C. Lines link the same lavage titred against $S.Tm^{\Delta oafA \ \Delta gtrC}$ and $S.Tm^{\Delta oafA \ pgtrABC}$. Red 1460 symbols and lines correspond to samples in which a strain able to phase-vary O:12 out-1461 1462 completed the O:12-0-locked strain by more than 100-fold on day 4. In each of these 1463 mice, the IgA titre specific for S.Tm with an O:12-0 epitope was higher than the titre of IgA specific for the phase-varied O:12-2 variant, whereas in mice where the ability 1464 1465 to phase-vary O:12 did not confer a selective advantage, titres against S.Tm^{$\Delta oafA \Delta gtrC$} and S.Tm^{$\Delta oafA pgtrABC$} were similar. All vaccinated mice were C57BL/6 and had an SPF 1466 1467 microbiota.

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Fig S6: The $\Delta gtrC$ mutation can be complemented in trans

1470 **Fig. S6: The** $\Delta gtrC$ **mutation can be complemented in trans:** C57BL/6 mice were 1471 vaccinated and pre-treated as in **Fig. 1**. The inoculum contained a 1:1 ratio of *S*.Tm^{$\Delta oafA$} 1472 and *S*.Tm^{$\Delta oafA \Delta gtrC$} pgtrABC. Competitive index in feces was determined by differential 1473 selective plating over 4 days post-infection.

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Fig. S7: Intestinal Lavage IgA titres from uninfected mice, quantified by ELISA and bacterial flow cytometry

1475 Fig. S7: Intestinal lavage IgA titre calculations for uninfected C57BL/6 mice 1476 vaccinated with PA-STm $^{\Delta gtrC}$ and PA-STm ET by dirty-plate ELISA and flow cytometry. C57BL/6 mice received either PA-STm^{$\Delta gtrC$} (n=8) and PA-STm^{ET} (n=11) 1477 per os once per week for 4 weeks. On d28, mice were euthanized and intestinal lavages 1478 1479 collected and cleared by centrifugation. Overnight cultures of S.Tm^{$\Delta gtrC$}, S.Tm^{pgtrABC} S.Tm^{$\Delta oafA$} and S.Tm^{$\Delta oafA$} pgtrABC</sup> were made in 0.2µm-filtered LB containing the relevant 1480 antibiotics. Bacteria were washed twice by centrifugation at 7000g to remove debris 1481 that may have accumulated during growth and used to coat ELISA plates (50µl of 1482 1483 OD=1-0 per well) or as target for bacterial flow cytometry (10⁵ bacteria per sample). 1484 Titration curves plotting A405 (ELISA) or median fluorescence intensity (bacteria flow 1485 cytometry) as read-outs of IgA binding, against dilution factor of lavages were used to 1486 calculate titres from 4-parameter logistic curve-fits. Representative overlayed histograms of the flow cytometry read-out from one PA-STm^{ΔgtrC} and PA-STm^{ET}-1487 1488 vaccinated mouse are shown (Colours represent different 3-fold serial dilutions of the 1489 intestinal lavage: red=2, blue=6, orange=18, green=54, dark green = 162, pink = 486. Pre-gated on scatted as in Fig. S4). P-values were calculated using 2-tailed Mann 1490 1491 Whitney U tests. Flow cytometry and ELISA reveal similar results, but with flow 1492 cytometry giving a clearer read-out. This is likely due to binding of lysed bacterial 1493 components to the ELISA plate scaffold, including protein components that will be 1494 identical between our strains as well as antigenic, but which are not accessible on the 1495 surface of live cells, and are therefore irrelevant for protection. We have therefore used 1496 bacterial flow cytometry to titre intestinal IgA throughout the manuscript as it more 1497 straightforward to equate to IgA binding to the surface of whole, intact, live cells.

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Fig. S8: Intestinal Lavage IgA titres from d9 post-infection, quantified by ELISA and bacterial flow cytometry

1500 Fig. S8: Intestinal lavage IgA titre calculations for 129S1/SvImJ mice vaccinated with PA-STm^{AgtrC} and PA-STm^{ET} and infected with S.Tm^{WT} for 9 days, by dirty-1501 plate ELISA and flow cytometry. 129S1/SvImJ mice received the indicated vaccine 1502 per os once per week for 4 weeks. On d28, mice were treated with oral streptomycin 1503 1504 and were infected with S.Tm^{WT}. Nine days post-infection, all mice were euthanized and 1505 intestinal lavages collected and cleared by centrifugation. Overnight cultures of S.Tm^{$\Delta gtrC$}, S.Tm^{pgtrABC} S.Tm^{$\Delta oafA$} and S.Tm^{$\Delta oafA$} pgtrABC</sup> were made in 0.2µm-filtered LB 1506 containing the relevant antibiotics. Bacteria were washed twice by centrifugation at 1507 1508 7000g to remove debris that may have accumulated during growth and used to coat 1509 ELISA plates (50µl of OD=1-0 per well) or as target for bacterial flow cytometry (10⁵ 1510 bacteria per sample). Titration curves plotting A405 (ELISA) or median fluorescence 1511 intensity (bacteria flow cytometry) as read-outs of IgA binding, against dilution factor 1512 of lavages were used to calculate titres from 4-parameter logistic curve-fits. P-values 1513 were calculated using 2-tailed Mann Whitney U tests. Flow cytometry and ELISA 1514 reveal similar results. Note that there is some broadening of the IgA response in PA-STm^{ΔgtrC}-vaccinated mice over the 9 days of infection when compared to the data in 1515 1516 Fig. S7. 1517



Figure S9: IgA-driven selective pressure functions identically in SPF Balb/c mice

1518 Fig. S9: IgA-driven selective pressure functions identically in SPF Balb/c mice. A-1519 C. Previous work indicated that Balb/c mice may respond better to oral vaccines and produce more secretary IgA than C57BL/6 mice (Fransen et al. 2015), therefore we 1520 tested the ability of PA-STm^{ET} to protect in Balb/c mice. Naive (closed circles), PA-1521 S.Tm^{ΔgtrC}-vaccinated (open circles) and PA-S.Tm^{ET}-vaccinated (crossed-circles) SPF 1522 1523 Balb/c mice were streptomycin-pretreated, infected (10⁵ CFU, 1:1 ratio of S.Tm^{$\Delta gtrC$}) and S.Tm^{$\Delta gtrC \Delta oafA$} per os). Note that naïve Balb/c mice were euthanized on day 3 due 1524 to severe disease. A. Secretory IgA titres (intestinal lavage dilution) against O:4[5], 12-1525 1526 0, and an O:4, 12-0 S.Tm. *p=0.0159 B. Competitive index (CFU S.Tm ^{\DeltagtrC}/CFU 1527 S.Tm^{Δ gtrC Δ oafA) in feces at the indicated time-points. 2-way ANOVA with Bonferroni} 1528 post-tests on log-normalized values, compared to naive mice. *p<0.0285. O:4-only producing S.Tm outcompetes in mice vaccinated with PA-STm^{$\Delta gtrC$} but not PA-STm^{ET}. 1529 1530 C and D. Correlation of the competitive index with the O:4-specific (C) and O:4[5]-1531 specific (**D**) intestinal IgA titre, r² values of the linear regression of log-normalized 1532 values. Open circles: Intestinal IgA from PA-S.Tm $^{\Delta gtrC}$ -vaccinated mice, crossed circles: Intestinal IgA from PA-S.TmET -vaccinated mice. Lines indicate the best fit 1533 1534 with 95% confidence interval. As both vaccinated groups have similar titres against the 1535 O:4[5]-producing S.Tm, a correlation of C.I. is observed only with the O:4-specific IgA 1536 titre E. CFU of S.Tm^{$\Delta gtrC$} (black symbols) and S.Tm^{$\Delta gtrC$} (orange symbols) per gram feces at the indicated time-points. **F** and **G**. CFU of $S.\text{Tm}^{\Delta gtrC}$ (black symbols) 1537 and S.Tm^{$\Delta gtrC \Delta oafA$} (orange symbols) per organ and day 4 post-infection (vaccinated) 1538 1539 and day 3 post-infection (naïve). Kruskal-Wallis test with Dunn's multiple comparison 1540 adjusted P values are shown. *p=0.022, **p=0.0085 H. Fecal Lipocalin 2 as a marker 1541 of inflammation in the indicated groups. 2-way repeat-measures ANOVA on lognormalized data, with Bonferroni post-tests comparing to the Naïve mice. 1542 1543 ***p=0.0002,****p<0.0001 I and J. Correlation between fecal lipocalin 2 on d3 post-1544 infection and O:4 and O:4[5]-specific intestinal IgA titres. r² values of the linear regression of log-normalized values. Lines indicate the best fit with 95% confidence 1545 1546 interval. Note that lines joining the points in **B**, **E** and **H** are to permit tracking of 1547 individual animals through the data set, and may not be representative of what occurs between the measured time-points. This experiment was based on the observations 1548 made in Fransen et al⁷² that better IgA-mediated protection is achieved in Balb/c mice 1549 than in C57BL/6 mice in response to live-attenuated vaccines. However, both mouse 1550 1551 lines behave similarly in this model. 1552



Fig. S10. PA-STm^{ET} mediated effects are not improved by addition of S.Tm^{ΔwzyB} to the vaccine cocktail.

1553 1554 Fig S10: PA-STm^{ET} mediated effects are not improved by addition of S.Tm $\Delta wzyB$ to the vaccine cocktail. C57BL/6 mice were vaccinated with vehicle only (Naïve, 1555 n=10), PA-S.Tm^{wt} (n=8), PA-STm^{ET} (combined PA-S.Tm^{\Delta}gtrC, PA-S.Tm^{\Delta}gtrC, PA-1556 S.Tm pgtrABC, and PA-S.Tm^{\Delta}oafA</sup> pgtrABC, n=9) or PA-STm^{ET+wzyB} (combined PA-1557 S.Tm^{$\Delta gtrC$}, PA-S.Tm^{$\Delta oafA$} $\Delta gtrC$, PA-S.Tm pgtrABC, PA-S.Tm^{$\Delta oafA$} pgtrABC and PA-1558 S.Tm^{$\Delta oafA \Delta gtrC \Delta wzyB$}, n=4). On day 28 after the first vaccination, mice were 1559 streptomycin pre-treated and challenged with 10⁵ S.Tm^{wt} orally. Fecal Lipocalin-2 1560 (LCN2) at day 1-3 post-infection, (A) and CFU S.Tm^{wt} per gram feces on day 1-3 post 1561 1562 -infection (**B**), CFU S.Tm^{wt} per mesenteric lymph node (MLN) at day 3 post-infection 1563 (C), and CFU S.Tm^{wt} per spleen at day 3 post-infection (D). A and B, 2-way repeat-1564 measures ANOVA on log-normalized data with Bonferroni multiple comparisons-tests 1565 reveals no significant difference between the vaccinated groups at any time-point. 1566 Adjusted p values are displayed. C and D: Kruskal-Wallis analyses with Dunn's multiple comparisons-tests comparing all groups were carried out for significance. 1567 Exact adjusted p values displayed. E. IgA titres in intestinal lavage of an experiment 1568 not included in (Fig. 3D), and additionally showing the group PA-S.Tm^{$\Delta oafA \Delta gtrC \Delta wzyB$}. 1569 1570 Titres are expressed as the dilution factor of lavage required to give an MFI=1000. 2-1571 way repeat-measures ANOVA on log-normalized data with Bonferroni multiple 1572 comparisons-tests. Note that lines joining the points in A and B are to permit tracking 1573 of individual animals through the data set, and may not be representative of what occurs between the measured time-points. 1574 1575