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29 Summary

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31 Though bacteriophages (phages) are known to play a crucial role in bacterial fitness and 32 virulence, our knowledge about the genetic basis of their interaction, cross-resistance and host-33 range is sparse. Here, we employed genome-wide screens in Salmonella enterica serovar Typhimurium to discover host determinants involved in resistance to eleven diverse lytic phages 34 including 4 new phages isolated from a therapeutic phage cocktail. We uncovered 301 diverse 35 36 host factors essential in phage infection, many of which are shared between multiple phages demonstrating potential cross-resistance mechanisms. We validate many of these novel 37 38 findings and uncover the intricate interplay between RpoS, the virulence-associated general 39 stress response sigma factor and RpoN, the nitrogen starvation sigma factor in phage crossresistance. Finally, the infectivity pattern of eleven phages across a panel of 23 genome 40 41 sequenced Salmonella strains indicates that additional constraints and interactions beyond the 42 host factors uncovered here define the phage host range.

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50 Introduction

There is increasing evidence that bacteriophages are a critical feature of microbial ecology, 51 52 evolution, virulence and fitness(Abedon, 2009; Breitbart and Rohwer, 2005; Koskella and 53 Taylor, 2018; Shkoporov and Hill, 2019; Suttle, 2007). However, knowledge about the molecular and genetic determinants of host-phage interactions and how they vary across the populations 54 55 of both are sparse even in otherwise well-studied model systems(Brüssow, 2013; de Jonge et 56 al., 2019; Nobrega et al., 2018; Rostøl and Marraffini, 2019; Samson et al., 2013; Weitz et al., 57 2013; Young and Gill, 2015). This derives, in part, from the technological limitations to resolve 58 the incredible specificity and the complex suite of bacterial mechanisms that confer both 59 resistance and sensitivity to the phage(De Smet et al., 2017; Nobrega et al., 2018; Young and Gill. 2015). A given bacterial host is likely to be susceptible to multiple phages, while a given 60 61 phage may infect a specific array of hosts and their variants. There are few, if any, studies that 62 map these mechanisms across phages and hosts, their interdependencies, and how variations in these mechanisms encode tradeoffs in host and phage fitness under different 63 64 conditions(Calendar, 2012; Casjens and Hendrix, 2015; De Smet et al., 2017; Karam and Drake, 1994; Molineux, 2002). However, this information is critical to an understanding of 65 microbial ecology and possibly exploiting the predator-prey dynamics for applications(Campbell, 66 2003; Díaz-Muñoz and Koskella, 2014; Lenski, 1988; Mirzaei and Maurice, 2017). 67

Knowledge of phage susceptibility and resistance determinants underlies a number practical 68 69 applications of phages. These applications span the use of phages and their combinations as biocontrol agents to improve water quality, decontaminate food, protect agricultural yield, and 70 defend and improve human health(Gordillo Altamirano and Barr, 2019; Keen and Adhya, 2015; 71 72 Kortright et al., 2019; Pirnay and Kutter, 2020). For example, because of the apparent ubiquity 73 of lytic phage with high host specificity for nearly any known pathogenic bacterial strain, phages 74 may provide a powerful alternative or adjutant to antibiotic therapies. The development of such 75 therapeutic phage formulations is pressing due to the alarming rise of antibiotic 76 resistance(Gordillo Altamirano and Barr, 2019; Keen and Adhya, 2015; Kortright et al., 2019; 77 Young and Gill, 2015). By characterizing the genetic basis of a bacterium's susceptibility and 78 resistance to a given phage and the pattern of cross-resistance or cross-sensitivity with other 79 phages, we can uncover evolutionary trade-offs on bacteria-phage interactions. These insights could also identify knowledge-gaps in our understanding of the host-range of a phage and offer 80 therapeutic solutions to recalcitrant infections(Chan et al., 2016; Gordillo Altamirano et al., 2021; 81 82 Mangalea and Duerkop, 2020; Shin et al., 2012; Trudelle et al., 2019; Wright et al., 2018). For 83 instance, by leveraging phages that target different receptors, combinations of phages or phage cocktails can be rationally formulated to both extend the host range and limit the rate of 84 85 resistance emergence(Bai et al., 2019; Chan et al., 2013; Kortright et al., 2019; Tanji et al., 2004; Yen et al., 2017). Such strategies can be further augmented by selecting phages that 86 specifically bind to bacterial virulence or antibiotic resistance factors to benefit from evolutionary 87 88 trade-offs in rational therapeutic outcomes (Chan et al., 2013; Kortright et al., 2019).

89 Compared to other antimicrobials, characterization of infectivity and cross-resistance between a 90 panel of phages has been limited to a few model organisms and remained phenomenological until recently(Hudson et al., 1978; Lindberg and Hellerqvist, 1971; Marti et al., 2013; Samuel et 91 al., 1999; Tu et al., 2017; Wright et al., 1980). The advent of genome-wide saturated transposon 92 93 sequencing (Bohm et al., 2018; Chan and Turner, 2020; Christen et al., 2016; Cowley et al., 94 2018; Pickard et al., 2013) and the corresponding DNA bar-code based modifications has 95 enabled the high-throughput and low cost genome-scale screening for the genetic determinants of these phenomena(Carim et al., 2020; Mutalik et al., 2019, 2020; Rousset et al., 2018; 96 Wetmore et al., 2015). Since this economically permits the independent screening of many 97

98 phages against a host library, it is now possible to determine and compare the genes that affect 99 the successful infection of one or more phages. These insights further suggest possible 100 mechanisms of cross-resistance (ie single mutations that confer resistance to multiple phages) 101 and collateral-sensitivity (ie single mutations that cause resistance to one phage while 102 sensitizing to another phage) that might arise when the host is naturally exposed to different combinations of these phages in the environment. As an example of this approach, we recently 103 104 employed a high-throughput genetic screening platform to characterize the phage resistance 105 landscape in Escherichia coli (E.coli) at an unprecedented scale(Mutalik et al., 2020). However, 106 the scale and benefits of these technologies have not vet been realized outside of such model 107 organisms, where bacterial physiology and phage-host interactions can be dramatically 108 different. Here, we employ a genome-wide loss-of-function screening technology to discover the 109 genetic determinants of phage susceptibility in Salmonella enterica, a globally important 110 infectious bacteria whose variants are responsible for the vast majority of bacterial food-borne 111 infections with an annual cost of \$3.7B dollars in 2013(Maculloch et al., 2015). Though 112 Salmonella enterica serovar Typhimurium (S. Typhimurium) has been used in the past as a model host to study phage infections(Graña et al., 1985; Lee et al., 2013; Lindberg and 113 Hellerqvist, 1971; MacPhee et al., 1975; Marti et al., 2013; Schwartz, 1980; Wilkinson et al., 114 115 1972; Wright et al., 1980), most Salmonella phages including therapeutically employed phage formulations have limited characterization regarding target receptors and host resistance 116 mechanisms (Bai et al., 2019; Gao et al., 2020; Islam et al., 2019; McCallin et al., 2018; 117 118 Petsong et al., 2019; Zschach et al., 2015). With increased numbers of Salmonella infections 119 that are resistant to antibiotics and displaying increased virulence(Medalla et al., 2016; (u.s.) and Centers for Disease Control and Prevention (U.S.), 2019), it is imperative to characterize 120 121 phages and their resistance patterns to enable their rational use as diagnostic and antimicrobial 122 agents.

In this study, we use a S. enterica serovar Typhimurium (S. Typhimurium) LT2 derivative that 123 124 serves as a genetically-accessible model for a clade of Salmonella responsible for food-borne enteric infections in humans. Using a barcoded transposon mutant library, we identified bacterial 125 genes whose loss confers altered sensitivity to 11 diverse double-stranded DNA phages 126 127 including 4 new phages isolated from a therapeutic phage cocktail. Our screens identified 128 known (and proposed new) receptors and also yielded novel host factors important in phage 129 infection, some of which we validate using single-gene deletion strains. Our genetic analysis allowed high resolution mapping of phage interactions with diverse cell surface components and 130 131 the operation of specific global regulatory systems that mediate specific metabolisms (e.g. rpoN) or virulence and stress response (e.g. rpoS). The diversity of cellular processes that influence 132 133 sensitivity to phage suggest that there may be multiple routes for cross-resistance and cross-134 sensitivity to emerge due to phage exposure in the environment or when used as therapeutic 135 cocktails. Finally, to assess if phage susceptibility and phage host-range can be predicted 136 based on the genetic determinants uncovered by our screens, we measured and analyzed 137 phage infectivity against a panel of 23 S.Typhimurium strains representative of naturally 138 occurring genetic diversity and phage infectivity.

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140 Results

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142 Identifying Salmonella genes involved in resistance to 11 diverse phages

143 We previously established a high-throughput approach to assay gene fitness using genome-

- 144 wide, random barcoded transposon sequencing (RB-TnSeq) (Price et al., 2018; Wetmore et al.,
- 145 2015). To systematically characterize phage infectivity pathways in Salmonella, we first

constructed an RB-TnSeq library in S. Typhimurium LT2 derivative strain MS1868 (Graña et al., 146 1985) (Methods). As an LT2-derived strain, S. Typhimurium MS1868 benefits from a long 147 148 history of Salmonella phage-host genetic interaction studies and well-characterized phage-149 resistant genotypes (Hudson et al., 1978; Lindberg and Hellergvist, 1971; Marti et al., 2013; 150 Samuel et al., 1999; Tu et al., 2017; Wilkinson et al., 1972; Wright et al., 1980). In addition, S. Typhimurium MS1868 is a restriction-minus genetic background (Graña et al., 1985), which 151 152 would potentially help uncover additional phage resistance factors by expanding the number of 153 phages infectious to the RB-TnSeq library. After transposon mutagenesis, we obtained a 66,996 154 member pooled library consisting of transposon-mediated disruptions across 3,759 of 4,610 155 genes, with an average of 14.8 disruptions per gene (median 12) (Figure 1A). We note that our 156 library does not have sufficient coverage of some likely non-essential genes that are likely to play an 157 important role in phage infection, for example igaA(Cho et al., 2014; Mariscotti and García-del 158 Portillo, 2009; Mutalik et al., 2020). Additional details for the composition of the S. Typhimurium 159 MS1868 library and comparison to a related single-gene-deletion library(Porwollik et al., 2014) 160 can be found in Table S1 and Dataset S1.

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We collected 11 lytic, dsDNA Salmonella phages, some of which are currently employed in 162 163 therapy and diagnostics. These phages are diverse, representing 5 of the 9 major dsDNA phage 164 families currently listed by the International Committee on Taxonomy of Viruses (ICTV): 3 from Myoviridae (FelixO1, S16, and Savina_GE), 1 from Podoviridae (P22), 4 from Autographiviridae 165 (Br60, Ffm, Shishito GE, and SP6), 2 from Siphoviridae (Chi, and Reaper GE), and 1 from 166 167 Demerecviridae (Aji GE). Though the receptors for 4 of the phages investigated here, Chi, FelixO1, P22 (obligately lytic mutant), and S16 are relatively well-studied, (Graña et al., 1985; 168 169 Lee et al., 2013; Lindberg and Hellergvist, 1971; MacPhee et al., 1975; Marti et al., 2013; 170 Schwartz, 1980; Wilkinson et al., 1972; Wright et al., 1980), only P22 phage has been subjected to a genome-wide genetic screen (Bohm et al., 2018). Additionally, Br60, Ffm, and SP6 have 171 172 suspected host-factor requirements for their infectivity cycle but otherwise have not been 173 extensively studied (Gebhart et al., 2017; Lindberg and Hellerqvist, 1971; Tu et al., 2017). This 174 panel of 11 phages also consists of 4 newly isolated phages from a commercial phage-cocktail 175 preparation from the Republic of Georgia (see Methods, Dataset S7): Aji GE EIP16 (Aji GE), 176 Reaper GE 8C2 (Reaper GE), Savina GE 6H2 (Savina GE), and Shishito GE 6F2 (Shishito_GE). All 11 phages except 3 (Br60, Ffm and Shishito_GE) infect wild-type S. 177 178 Typhimurium LT2 MS1868, which has an intact O-antigen (known as 'smooth-LPS'). Br60, Ffm and Shishito GE phages were grown using a 'rough-LPS' mutant strain of Salmonella which 179 180 consists of only core LPS (see Methods). Thus, the phage panel used here contains phages that either bind to smooth or rough LPS strains and allows comparison of the key host factors 181 182 important in their infectivity cycles.

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To identify Salmonella genes important for phage infection, we challenged the S. Typhimurium mutant library with each of the 11 dsDNA lytic phages (Table 1) at multiplicities of infection \geq 2 in both planktonic and non-competitive solid plate fitness experiments, and collected the surviving phage-resistant strains post incubation (Figure 1A, Methods). From samples collected before and after phage incubation, we sequenced the 20 base pair DNA barcodes (i.e. BarSeq) associated with each transposon mutant. We then calculated strain and gene fitness scores as the relative log2-fold-change of barcode abundances before versus after phage selection, as

previously described (Mutalik et al., 2020; Price et al., 2018; Wetmore et al., 2015) (Figure 1A, 191 192 Methods). Thus, in this study, a high positive fitness score indicates loss-of-function mutants in 193 Salmonella that are resistant to phage infection. We observed very strong phage selection 194 pressures during these competitive fitness experiments, consistent with our earlier observations (Mutalik et al., 2020), and thus we mostly limited our analysis to positive fitness scores. As 195 196 expected with our MS1868 library primarily consisting of O-antigen positive mutants, the vast majority of gene disruptions in MS1868 showed no significant fitness benefit against rough-LPS 197 binding Br60, Ffm and Shishito GE phages. However, we noticed strong fitness defects in 198 199 many of the LPS and O-antigen mutants in our library (Figure 1BC, Dataset S4), consistent with 200 optimal adsorption and infection in O-antigen-defective Salmonellae.

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202 In aggregate, we performed 42 genome-wide RB-TnSeq assays across liquid and solid growth 203 formats and discovered 301 phage-gene interactions (with 184 unique gene hits) that are 204 important for phage infection across the 11 phages studied (Figure 1BC, Datasets S2 and S4). 205 Though solid plate assay results were largely consistent with planktonic assays, some resulted 206 in genes with stronger fitness effects. Across all fitness experiments, we observed at least one 207 gene with a high fitness score (except 3 phages that infect rough-LPS strains), affirming the 208 successful competitive growth of mutants under phage selection. Some Salmonella phages 209 show enrichment of strains with disruptions in multiple genes, while other phages enrich strains 210 with disruptions in a more limited number of genes (Figure 1B). For example, we observed 98 genes enriched after Chi phage challenge and 73 high-scoring genes after Felix O1 challenge. 211 212 yet only 7 high-scoring genes after S16 phage challenge. As expected in any phage selection 213 experiment, we observed enrichment of genes that encode components of the cell envelope. 214 Nonetheless, we also identified dozens of genes that encode cytoplasmic components not 215 previously associated with phage resistance. To further categorize the genetic basis of phage resistance, we manually classified all identified genes with high fitness values into broad-216 217 functional categories: core-LPS and O-antigen biosynthesis, motility, secondary messengers, 218 transcription factors and other metabolism (Figure 1C). These results demonstrate that genes 219 downstream from phage receptors are important for phage infectivity.

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221 Both receptor and non-receptor host factors are involved in phage infection

222 A key-determining step in the phage infectivity cycle is the interaction of phages with any 223 bacterial cell surface-exposed molecules or receptors. Consequently, any changes in the structure or level of these surface-exposed molecules that accompany resistance to specific 224 225 phages are usually assigned a function of phage receptor. To confirm the effectiveness of our 226 genetic screen, we looked for receptors that are known for a few of the phages used in this work (Bohm et al., 2018; Hudson et al., 1978; Lindberg and Hellerqvist, 1971; MacPhee et al., 1975; 227 228 Marti et al., 2013; Samuel et al., 1999). Indeed, in agreement with published data available for 229 FelixO1, P22, Chi, SP6 and S16 phages, we found high fitness scores for candidate receptor genes with >1,000 fold enrichment of transposon mutants. These included genes encoding 230 231 protein receptors such as ompC (outer membrane porin C) for S16 and flagellar body for Chi phage, while LPS and O-antigen biosynthesis genes for P22, SP6 and FelixO1 phages (Figure 232 233 1C). Our results are also largely consistent with a recent genome-wide screen in Salmonella against P22 infection (Bohm et al., 2018). Though O-antigen and outer core GlcNAc (the 234 235 biosynthetic product of RfaK) have been known as SP6 and as FelixO1 phage receptors 236 respectively (Hudson et al., 1978; Lindberg and Hellergvist, 1971; Marti et al., 2013; Samuel et 237 al., 1999; Tu et al., 2017; Wilkinson et al., 1972; Wright et al., 1980), our genome-wide screens 238 provided an array of additional, non-receptor genes as target loci for phage resistance selection.

A detailed description and analysis of outer membrane components such as LPS required for these phages can be found in Text S1.

241 242 In addition to the genes coding for phage receptors, our genetic screens also uncovered high-243 scoring genes that are known to be involved in the regulation of target receptors. For example, 244 deletion of the EnvZ/OmpR two component system involved in the regulation of ompC and gene products involved in the regulation of cellular motility (nusA, tolA, cyaA and guanosine 245 246 penta/tetraphosphate ((p)ppGpp) biosynthesis and metabolism all showed high fitness scores 247 in the presence of S16 and Chi phages, respectively(Graña et al., 1985; Lee et al., 2013; 248 Lindberg and Hellerqvist, 1971; MacPhee et al., 1975; Marti et al., 2013; Schwartz, 1980; 249 Wilkinson et al., 1972; Wright et al., 1980). These high-scoring gene candidates were previously not known to be associated with phage resistance in Salmonella. Other than the phages 250 251 mentioned above that bind to surface components of smooth-LPS Salmonellae, we also 252 screened Br60 and Ffm phages, which are known to strictly infect rough-LPS strains and not 253 bind to smooth WT MS1868 parental strain (as O-antigen structure probably occludes their 254 native receptor)(Gebhart et al., 2017; Lindberg and Hellerqvist, 1971; Tu et al., 2017). As our MS1868 library primarily consists of O-antigen positive mutants, the vast majority of gene 255 disruptions in MS1868 showed no significant fitness benefit against these phages. However, we 256 257 noticed strong negative scores for many of the LPS and O-antigen mutants in our library (Dataset S4), indicating these strains have rough LPS phenotype and are sensitive to Br60 and 258 259 Ffm phages. As an additional resource, we determined the specific rough-LPS requirements for 260 these phages using an O-antigen deficient library and individual mutant susceptibility assays (Text S1 - Extended Results: Uncovering Host-Factors of Rough-LPS Requiring Phages Br60, 261 262 Ffm, and Shishito_GE).

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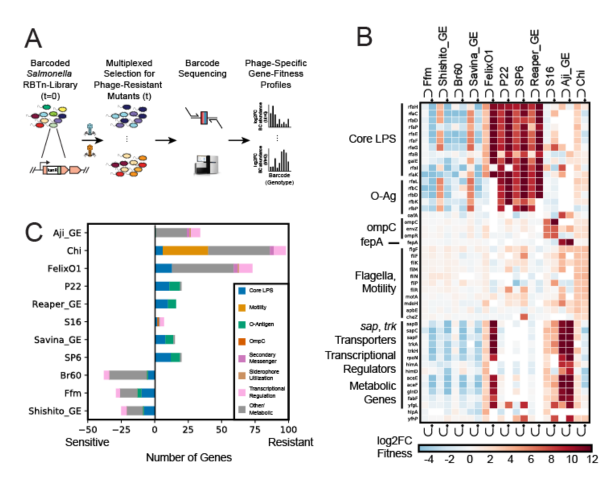
Among the four newly isolated phages (Reaper GE, Savina GE, Aji GE and Shishito GE), 264 265 Reaper GE showed strict requirements for O-antigen including a complete LPS (Figure 2B), while Savina GE primarily showed dependency on O-antigen followed by inner core mutants, 266 267 and outer core mutants (Figure 2B, Dataset S4). For T5-like phage Aji_GE, both fepA (TonBdependent enterobactin receptor) and oafA (O-antigen acyltransferase) showed high fitness 268 269 scores. OafA performs an acetylation reaction on the abequose residue to create the O5-270 antigen serotype in LT2-derived strains (Slauch et al., 1996), and probably enhances infection 271 via gaining access to the FepA-TonB complex. Related phenomena have been observed for 272 other Demerecviridae, where other O-antigen modifications facilitated increased phage 273 susceptibility (Heller and Braun, 1982; Kim and Ryu, 2012). Finally, similar to Br60 and Ffm phages isolated on rough-LPS Salmonella, Shishito_GE displayed strong host fitness defects in 274 275 many of the LPS and O-antigen mutants in our library.

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282 Figure 1. Genome-wide screen to identify host factors involved in phage infection. (A) Overview of 283 pooled fitness assays. For additional details, see methods. Briefly, for each experiment, S. Typhimurium 284 RB-TnSeq library was exposed to a high MOI of one of eleven dsDNA Salmonella phages. Strains were 285 tracked by quantifying the abundance of DNA barcodes associated with each strain by Illumina sequencing. Phage-specific gene fitness profiles were calculated by taking the log2-fold-change of 286 287 barcode abundances post- (t) to pre- (t=0) phage predation. High fitness scores indicate that loss of genetic function in Salmonella confers fitness against phage predation. (B) Heatmap of top 10 high-288 289 confidence gene scores per phage are shown (many genes are high-confidence hits to multiple phages). 290 Both planktonic and solid plate data are shown. Three rough-LPS binding phages Br60, Ffm, and 291 Shishito GE do not infect wild-type MS1868, but can infect specific MS1868 mutants, overall showing 292 negative fitness fitness values in our screen. Noncompetitive, solid agar growth experiments are marked 293 with a (*). (C) Total number of high-scoring genes per phage and their functional role. Input data for 294 Figures 1B and 1C is found in Dataset S4 and can be recreated using Supplementary Code - Figure1BC. 295

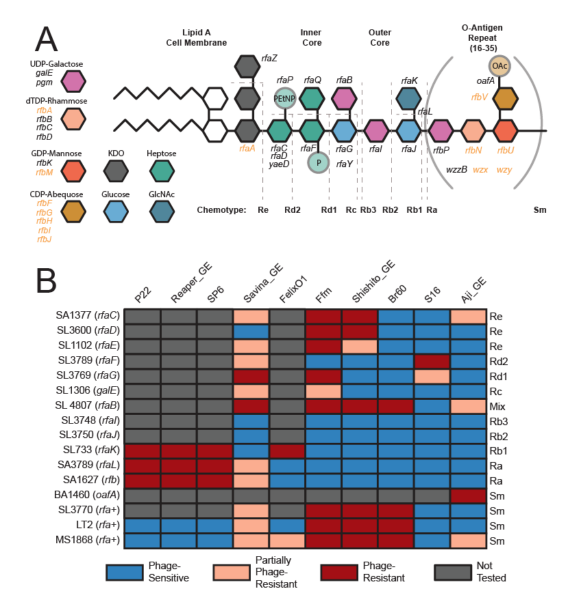
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To validate some of the top phage resistance phenotypes from our genetic screens, we used an established collection of *Salmonella* mutant strains in addition to the construction of deletion strains (Table S5)(Roantree et al., 1977; Sanderson et al., 1974). To confirm the role of OafA and TonB-dependent enterobactin receptor FepA (FepA-TonB complex) on Aji_GE infectivity, we constructed strains with deletions in *oafA*, *fepA*, and *tonB* (Methods). Aji_GE phage plaque assays on these strains confirmed the essentiality of OafA and both FepA-TonB in infection (Figures 2 and S17). For phages that showed stringent requirements of O-antigen and LPS, we

used an established chemotype-defined LPS mutant panel in a S. Typhimurium strain 305 306 background that is closely related to our LT2 derivative (Figures 2, S5-S11, and S13-S15, Table 307 S5, Methods). Our phage infectivity results on the LPS chemotype panel are in agreement with 308 earlier published data for some of the phages used in this work (Bohm et al., 2018; Lindberg 309 and Hellergvist, 1971; Marti et al., 2013; Mutalik et al., 2020; Wright et al., 1980) and consistent with our high-throughput genetic screens for all phages (Figure 1B). For example, LPS 310 311 chemotype panel data confirmed the strict requirements for O-antigen including a complete LPS 312 for Reaper_GE infectivity. We confirmed that Savina_GE most efficiently infects strains with an 313 incomplete outer core, but less so against strains without O-antigen or strains missing outer 314 core entirely (Figures 2 and S9). This result indicates Savina_GE preferentially employs LPS as 315 a receptor, but branched LPS residues such as those added by *rfaK* and O-antigen biosynthesis probably hinder efficient adsorption. Though OafA activity is important for Aji GE infection 316 317 (Figure 2, BA1460), the acetylation provided by OafA activity does not seem to be critical in the 318 absence of complete LPS and O-antigen as those mutants showed significant infection (Figures 2, S3, S11, and S17). The plaque assays of Shishito_GE on the LPS mutant panel confirmed 319 320 that, like phages Br60 and Ffm, it only infects rough-LPS strains of Salmonella (Figures 2, S4, 321 and S13-S15). As the inner and outer core LPS structure of S. Typhimurium is conserved in E. coli K-12, we confirmed these observations using data from an RB-TnSeq library of E. coli K-12 322 323 (Figure S4, Text S1, Methods). In summary, the combination of our high-throughput genetic screen and assays on single-gene deletion strains provided higher resolution mapping of O-324 325 antigen, LPS or protein receptor requirement for all 11 phages in Salmonella (a detailed 326 description for each phage is in Text S1).

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329 Figure 2: Validation of LPS-moiety requirements for Salmonella phages (A) Overview of O5 S. Typhimurium LPS and O-antigen biosynthesis. The four sugars in brackets comprise the O-antigen, which 330 331 repeats 16-35 times per LPS molecule under standard growth conditions. Key for non-essential LPS and 332 O-antigen precursor biosynthesis genes are described to the right. Genes covered in our library and used for analysis are written in black. Genes not covered in our library, and thus not analyzed in this study are 333 334 written in orange. (B) Infectivity matrix using a previously established Salmonella LPS panel (Table S5). 335 The identity of the LPS chemotype corresponding to specific mutation is presented in (A). sm stands for 336 smooth-LPS chemotype. Data for this figure is aggregated from Figures S5-S11, S13-S15, and S17. 337 338

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Discovery of Novel Cross-Resistant Genotypes Between Diverse Phages 342

Next, we looked at the number and pattern of high-fitness scoring genes against our panel of 343 344 phages to identify similarity in infectivity cycles and commonality in genetic barriers leading to 345 phage cross-resistance. The most studied mode of resistance between phages is when they 346 share a common receptor (for example, phages binding to LPS), and any modification in the 347 common receptor yields cross-resistance to those phages (Chan and Turner, 2020; Mutalik et al., 2020; Shin et al., 2012; Wright et al., 2018, 2019). Though it is possible that other host 348 349 factors are important for the infectivity cycle of different phages and can impart phage cross-350 resistance phenotypes, it remains a challenge to identify such non-receptor host factors and 351 their role in phage infection. Thus they are not widely reported, nonetheless in the context of 352 phage cross-resistance. For example, mutations in global transcriptional regulators can impart 353 broad resistance to diverse phages that bind to different receptors, but have been proposed to 354 impart higher fitness costs which probably explain their lower frequency of emergence(Betts et 355 al., 2016; Díaz-Muñoz and Koskella, 2014; Hesse et al., 2020; Mutalik et al., 2020; Wright et al., 356 2018, 2019).

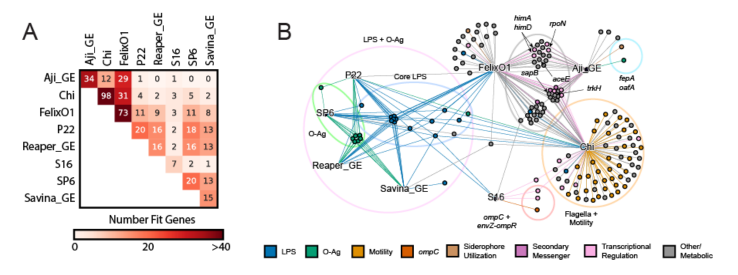
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To gain more insights into phage cross-resistance, we compared the genes that show high-358 359 fitness scores across the 8 smooth-LPS binding phages screened in our study (Figure 3AB). 360 The pair-wise comparison between any two phages indicated that, there is a wide range of shared high-fitness scoring genes. As expected, phages that bind to the same receptor shared 361 362 many high-scoring genes indicating potential cross-resistance between them. For example, P22 363 SP6 and Reaper_GE bind to O-antigen and share many common high scoring hits. Conversely, there are instances of no high-fitness scoring genes shared between phages employing 364 different receptors (for example, between Aii GE and the O-antigen requiring phages 365 366 Reaper_GE, SP6 and Savina_GE) (Figure 3A). Unexpectedly, we also observed instances of shared genes across phages that bind to different receptors, and point to a role played by the 367 368 non-receptor host factors (Figure 3). For example, Aji GE and FelixO1 have different receptors, 369 yet they share a large number of high-fitness scoring genes, indicating potential cross-370 resistance independent of their primary receptors (Figures 1 and 3). Out of 52 non-receptor 371 genes conferring resistance to FelixO1 and 32 non-receptor genes conferring resistance to Aji_GE, 29 were common to Aji_GE and FelixO1. These common non-receptor host factors 372 373 appear to play diverse roles, and the functions they encode include disruptions across central 374 metabolism (aceEF, pta, ackA, fabF), amino acid biosynthesis and regulation (rpoN, glnDLG, ptsIN, aroM), global regulation (himAD, crp, rpoN, lon, arcB), ion transport (trkAH), peptide 375 376 transport (sapABCF), secondary messenger signaling (gppA, cyaA), translation (trpS), and 377 other genes with less clear functions (nfuA, yfgL, ytfP). Some of these genes were recently 378 implicated as host-factors in phage resistance in related organisms (Cowley et al., 2018; 379 Goosen and Putte, 1995; Wright et al., 2018), though their role in phage cross-resistance and 380 mechanisms were not determined. For example, trk, sap, ace and rpoN were recently associated with diverse phage resistance in E. coli (Cowley et al., 2018; Kortright et al., 2020) 381 382 and P. aeruginosa (Wright et al., 2018). himA and himD (i.e. integration host factor subunits alpha and beta) are known to be involved in temperate phage infection pathways, though not 383 shown for obligately lytic phages such as Aji GE and FelixO1(Goosen and Putte, 1995). 384

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To investigate if these mutants indeed display cross-resistance to both Aji_GE and FelixO1, we selected a few top scoring genes to study further: *trkH*, *sapB*, *aceE*, *rpoN*, *himA*, and *himD*. For each of these 6 genes, we created individual mutants (Methods) and assessed Aji_GE and FelixO1 phage infectivity. Indeed, the *trkH*, *sapB*, *aceE*, *rpoN*, *himA* and *himD* mutants showed increased resistance to both FelixO1 and Aji_GE (Figures S19-S20). Consistent with prior reports of high fitness costs being associated with non-receptor phage cross-resistant mutants

392 (Wright et al., 2018), mutants in *aceE*, *rpoN*, and *himA* displayed significant growth defects 393 during planktonic growth, but were sufficiently fit to be uncovered in our screens. Some of these 394 genes (for example, potassium transporter Trk and nitrogen assimilation sigma factor RpoN) are 395 known to play an important ecological role in *Salmonella* virulence and fitness in infection 396 contexts (Klose and Mekalanos, 1997; Su et al., 2009), indicating these phage resistance loci 397 may exhibit an evolutionary trade-off with virulence.



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399 Figure 3: Cross-resistance is common between Salmonella phages. (A) Summary of cross-400 resistance patterns between phages observed in our screens. Heatmap color represents the total number 401 of shared gene disruptions in S.Typhimurium that yield resistance to both phages. (B) Mixed-node 402 network graph showing connections between phage nodes (text labels, black) and gene nodes (colored nodes). A gene node is connected to a phage node if disruptions in that gene gave high fitness against 403 404 that phage (Dataset S4). Gene nodes are colored by encoded function. Notable gene function groupings 405 and genes are additionally highlighted. Figures 3A and 3B are created from Dataset S4 using 406 Supplemental Code - Figure3AB.

407

408 Sigma Factor Interplay Mediates Phage Cross-Resistance in Salmonella

409 To better identify the genetic basis of the phage cross-resistance phenotype imparted by trkH, 410 sapB, rpoN, and himA mutants, we carried out RNA-Seq experiments and investigated whole-411 genome expression-level differences for each deletion compared to wild-type MS1868 (N=3 for 412 all except for himA, which was N=2). In aggregate, we observed 635 differentially expressed genes (among which 437 are unique to one of the knock-out strains) in trkH, sapB, rpoN, and 413 himA mutants compared to wild-type (Figure 4A, Dataset S6). To the best of our knowledge, 414 415 none of the differentially expressed genes were related to FelixO1's and Aji GE's suspected 416 receptors (LPS and FepA respectively). In addition, neither of the known innate immunity 417 defense mechanisms in S. Typhimurium (type I CRISPR or type I BREX), were found to be differentially expressed in any of these genetic backgrounds (Barrangou and Oost, 2014; 418 Shariat et al., 2015). Thus we suspected this mode of resistance was likely due to global 419 420 regulatory changes. We focused our analysis to trkH, sapB, and rpoN mutant backgrounds that showed upregulation of the spv virulence operon (spvABC), located on the PSLT plasmid native 421 422 to S. Typhimurium (Dataset S6). In addition to being studied for its essentiality in Salmonella virulence, the spv operon is also well-known for being regulated by RpoS, a general stress 423

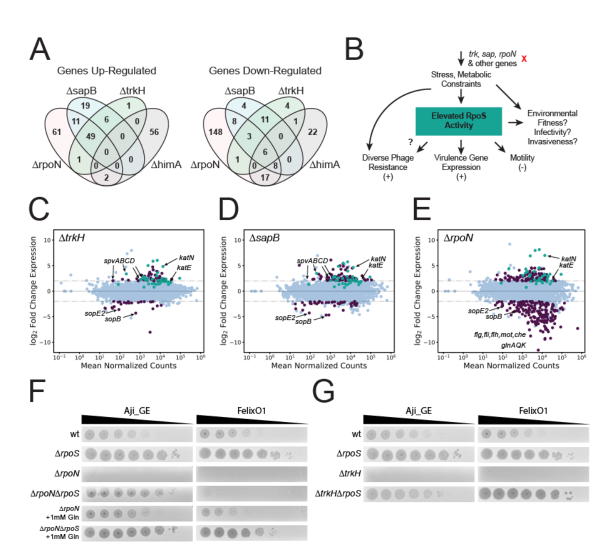
response sigma factor (Chen et al., 1995; Fang et al., 1992; Nickerson and Curtiss, 1997). As 424 the RpoS regulation is well-studied in E. coli and S. Typhimurium (Battesti et al., 2011; Hengge-425 426 Aronis, 2002; Ibanez-Ruiz et al., 2000; Lago et al., 2017; Lévi-Meyrueis et al., 2014; Lucchini et 427 al., 2009; Nickerson and Curtiss, 1997), we looked for expression changes in RpoS-dependent 428 genes in trkH, sapB, and rpoN mutant backgrounds. We found that a number of known RpoSregulated genes were significantly upregulated versus wild-type (passing thresholds of log 2FC >429 430 2, p_adj < 0.001) (Figure 4CDE, Dataset S6), further implicating RpoS involvement in resistance 431 to both Aji_GE and FelixO1 phages (Figure 4B).

432

The general stress response sigma-factor RpoS activity in Salmonella is critical for many 433 aspects of its adaptive lifestyle, including general virulence (Lévi-Meyrueis et al., 2014; Lucchini 434 et al., 2009). However, comparative studies in clinical isolates of Salmonella found decreased 435 436 RpoS activity in model strain LT2 versus related virulent strains due to a suboptimal start codon 437 (Wilmes-Riesenberg et al., 1997). As a LT2 derivative, MS1868 has this suboptimal codon 438 (Graña et al., 1985), so it is intriguing to find signatures of elevated RpoS activity and virulence-439 associated spv expression in phage resistant candidates. To confirm the impact of RpoS on 440 phage infection, we created a rpoS deletion mutant and additional double gene replacement 441 mutants of rpoS with one of trkH, sapB, or rpoN. The single rpoS deletion mutant displayed increased sensitivity to both FelixO1 and Aji GE phage (Figures 4FG and S19-S20). In addition, 442 the rpoS deletion was also sufficient to restore infectivity in trkH, sapB, and rpoN mutants to 443 444 levels observed in rpoS mutants (Figures 4FG and S19-S20). While himA mutants did not show 445 elevated levels of RpoS activity in our RNA-Seg data, we suspect that phage-resistance in 446 many mutants within the Aji GE and FelixO1 cross-resistance network emerged from RpoS 447 activity beyond these mutants. More broadly, RpoS activity likely plays a role in intermediate 448 phage-resistance phenotypes that are typically difficult to quantify in pooled fitness assays, but 449 observable for these two phages.

450

In the rpoN (encoding sigma factor-54) mutant background, the rpoS mutation was sufficient to 451 452 restore infectivity of phage Aii GE, but insufficient to restore infectivity of FelixO1 (Figures 4F 453 and S19-S20). Like RpoS, the alternate sigma factor RpoN is known to regulate a diverse set of 454 pathways involved in adaptation and survival in unfavorable environmental conditions including 455 nitrogen starvation. Because rpoN mutants decrease glutamine uptake and biosynthesis and 456 have significant growth defects, the phage resistance phenotype observed in rpoN mutants 457 potentially indicate the importance of glutamine levels on successful phage infection (Dataset 458 S4) (see also: (Aurass et al., 2018; Samuels et al., 2013)). To assess the dependence of 459 glutamine on phage resistance mechanism, we repeated phage infection supplemented with 460 glutamine in rpoN mutants. Both FelixO1 and Aji_GE were able to successfully plaque on rpoN mutants supplemented with glutamine. In the *rpoN*, *rpoS* double mutant background, additional 461 glutamine supplementation was able to nearly restore FelixO1 infectivity to the rpoS mutant's 462 463 baseline (Figures 4F and S19-S20). Thus, we propose *rpoN* loss-of-function probably manifests two avenues of phage resistance. First, nutrient limitation to the cell can "starve" phage 464 replication, such as FelixO1 but not Aii GE, during infection, Second, elevated RpoS activity 465 466 (likely induced by nutrient limitation) confers further resistance to phage infection, extending to 467 diverse phages such as FelixO1 and Aji_GE. In summary, these studies uncover intricate 468 interplay between host factors and nutritional status of the cell in phage cross-resistance 469 phenotype.



470

471 Figure 4: Cross-resistance mechanisms are mediated by RpoS. (A) Summary of genes with significant 472 up- and down-regulation relative to wild-type for sapB, trkH, rpoN, and himA mutants. Reported values 473 are genes with log2-fold changes over 2 and Bonferroni-corrected p values below 0.001. (B) Proposed model for phage cross-resistance observed in this study. Loss of function of genes such as trkH, sapB, or 474 rpoN impose stress and metabolic constraints on S. Typhimurium. In some cases, this elevates RpoS 475 476 activity and leads to multi-phage resistance. However, the environmental fitness, virulence, and 477 invasiveness implications of these mutants are not known. (CDE) MA-plots for differential expression data for (C) MS1868 Δ trkH, (D) MS1868 Δ sapB, and (E) MS1868 Δ rpoN mutants over wild-type MS1868. 478 Differentially expressed genes (abs(log2FC \geq 2), Bonferroni-corrected p values below 0.001) are shown in 479 480 purple. RpoS-regulated genes are shown in teal based on a curated list from (Lucchini et al., 2009). 481 Specific genes are highlighted for emphasis including RpoS-activity indicators katE and katN. (F) Aji GE 482 and FelixO1 phage susceptibility assays focused on *ArpoN*-mediated phage resistance. For both phages 483 supplementing with glutamine (Gln) restores phage infectivity in $\Delta rpoN$ context. A secondary deletion in 484 rpoS is sufficient to restore Aji GE infectivity in a \triangle rpoN strain. However, FelixO1 is only restored with

485 additional supplementation of glutamine. (G) Aji_GE and FelixO1 phage susceptibility assays focused on

486 *∆trkH*-mediated phage resistance. A secondary deletion in *rpoS* is sufficient to restore both Aji_GE and

487 FelixO1 infectivity in a \triangle *trkH* strain. Figures 4ACDE are created from Dataset S6 using Supplementary

488 Code - Figure4. 489

490 Investigation into phage sensitivity of natural Salmonella strain variants

491 Finally, we wondered how gene requirements uncovered in our genome-wide genetic screens 492 corresponded to naturally occurring variation in and phage sensitivity of S.Typhimurium isolates. 493 More broadly, we were interested in to what degree these gene requirements in a model strain 494 were predictive of phage sensitivity patterns in closely related strains. Though phage host range 495 determination using a panel of strains belonging to a species of bacterium is a century old practice, the genetic basis of the phage infectivity pattern has remained unresolved (Holmfeldt et 496 al., 2007; Hyman and Abedon, 2010; de Jonge et al., 2019; Moller et al., 2019; Weitz et al., 497 498 2013). For example, phage infectivity patterns using a panel of phages (phage typing) to discriminate Salmonella serovars for epidemiological investigation/surveillance is even practiced 499 today, while the infectivity pattern is not typically investigated mechanistically(Chirakadze et al., 500 2009; Rabsch, 2007). We hypothesized that the similarity and differences in genetic 501 502 determinants involved in phage resistance might be able to explain the genetic basis of phage 503 infectivity when extended to a panel of Salmonella strains. To assess the relationship between genomic content and phage sensitivity among natural strain variants, we sourced a panel of 21 504 505 S. Typhimurium strains belonging to the SARA collection (Beltran et al., 1991). We also included a model nontyphoid clinical isolate D23580 from Malawi (Canals et al., 2019) and ST4/74 strain, 506 507 originally isolated from a calf with salmonellosis(Richardson et al., 2011) as a reference. The 508 SARA collection is a set of strains of Salmonella isolated from a variety of hosts and 509 environmental sources in diverse geographic locations, classified into 17 electrophoretic types, 510 observed variation in natural populations and is reflective of much of the diversity identified in 511 panels derived from recent S. Typhimurium outbreaks(Fu et al., 2015).

512

513 We re-sequenced these 21 strains to confirm their identity and assembled their genomes as 514 described in Methods. Our analysis showed that all isolates, except for SARA7 and SARA8, 515 have a close phylogenetic relationship (>99% pairwise average nucleotide identity, ANI) in agreement with an earlier report(Fu et al., 2015). Next, we searched for the 184 unique high-516 517 scoring gene hits uncovered in this work (Table S4) across our panel of Salmonella genomes 518 and observed little variation in the sequence of genes, though there might be changes in expression and activity (Dataset S8). Among the key differences in our gene content analysis. 519 520 we observed nonsense mutations or frame shifting changes in the coding region of rfaK in 521 SARA20, ompC and rfbN in SARA6 and oafA in SARA9 compared to our reference strain 522 Salmonella LT2. Mutation in the coding region of rfaK/waaK in SARA20 yields two truncated 523 proteins, and neither of them have a complete glycosyltransferase domain. It is known that *rfaK* mutants lack the GlcNAc residue in the LPS outer core and are also unable to express O-524 525 antigen because this GlcNAc residue is essential for the recognition of core oligosaccharide 526 acceptor by the O-antigen ligase WaaL(Hoare et al., 2006). We postulated that absence of outer 527 core GlcNAc (the biosynthetic product of RfaK) in SARA20 probably alters the structure of O-528 antigen and may yield resistance to O-antigen binding by P22, SP6, Reaper GE and FelixO1 phages. Disruption in the ompC coding region in SARA6 may compromise S16 phage infectivity 529 and disruption in *oafA* coding region (in SARA9) probably interferes with efficient infection by 530 531 Aji_GE. Broadly, our analysis predicts that all 23 Salmonella isolates except the ones mentioned

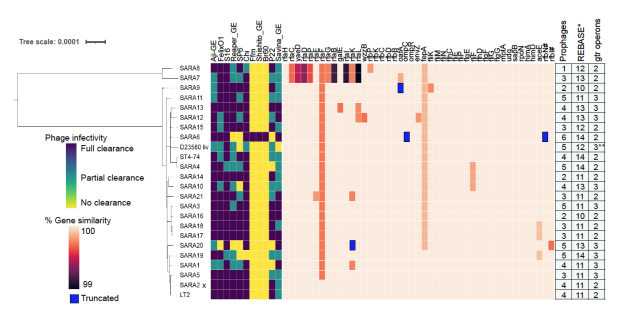
above should show similar phage infectivity patterns as compared to the laboratory strain used in our genetic screens.

535 To assess the infectivity pattern of the 11 phages against the 23 Salmonella strains, we carried 536 out standard spotting assays. Figure 5 shows the phage infectivity data and phylogenetic distance between Salmonella strains, with a phylogenetic tree built from gene sequences of 115 537 538 single-copy marker genes (Methods). In agreement with our genome-based prediction, 22 539 strains (out of 23 strains) displayed broad sensitivity to all O-antigen binding phages (except 540 Ffm, Br60 and Shashito-GE) (Figures 1-2 and 5). SARA6 was the only strain sensitive to Ffm, 541 Br60 and Shashito_GE phages and was also resistant to all phages binding O-antigen (P22, 542 SP6, Reaper_GE), indicating SARA6 may have rough-LPS phenotype. Analysis of SARA6 543 genome indicated that rfbN, a gene encoding rhamnosyltransferase important for O-antigen 544 synthesis has a mutation and that this strain would not be able to express O-antigen, in 545 agreement with its resistance to O-antigen binding phages (P22, SP6 and Reaper GE) while 546 showing sensitivity to core LPS binding phages (Ffm, Br60 and Shashito-GE). Disruption of the 547 ompC coding region in SARA6 while retaining infectivity with S16 phage indicates there is a 548 possibility of OmpC independent infectivity pathway as seen in some T4-like phages(Washizaki 549 et al., 2016). SARA20 showed no sensitivity to both smooth-LPS binding (P22, SP6, 550 Reaper GE, FelixO1) and rough-LPS binding phages (Ffm, Br60 and Shashito-GE), raising an 551 interesting question about its LPS architecture. Though mutation in the coding region of rfaK 552 and absence of O-antigen in SARA20 explains its resistance to P22, SP6, Reaper GE, FelixO1 553 phages, though the resistance showed by rough-LPS binding phages Ffm, Br60 and Shashito-554 GE indicate additional factors likely play a role. Finally, in agreement with our gene content 555 analysis (above), SARA9, with disruption in the *oafA* coding region, showed inefficient infection 556 by Aji GE.

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In addition to these broad agreements between gene content analysis and phage infectivity, the 558 genetic basis of strong phage resistance showed by SARA1 and SARA4 (for P22). SARA10 (for 559 560 SP6), D23580 (for SP6 and Savina_GE) and SARA19 (for SP6, Chi) is unclear. We also observed a partial clearance pattern in our spot tests for many phages across 23 isolates, 561 562 probably indicating inefficient infection cycles or partial inhibition characterized by turbid 563 plaques. Overall, these results indicate that phage host range is probably defined by additional 564 constraints than the host factors uncovered in our genetic screens. To look for other host factors 565 that might be playing a role in efficient phage infection, we bioinformatically searched for prophages, genes encoding O-antigen modification systems (gtrABC operon) and restriction 566 567 modification systems encoded in our panel of 23 Salmonella strains. The role played by these 568 genetic elements on phage infectivity and resistance are well appreciated (Bernheim and Sorek, 2020; Bondy-Denomy et al., 2016; Cota et al., 2015; Davies et al., 2013; Dedrick et al., 2017; 569 570 Dy et al., 2014; van Houte et al., 2016; Owen et al., 2020; Rostøl and Marraffini, 2019; Samson 571 et al., 2013; Vasu and Nagaraja, 2013; Wahl et al., 2019). Our comparative analysis provided a list of strain-specific restriction/modification and O-antigen modification genes that may 572 573 influence phage infection outcome. However, we could not identify a single genomic loci, whose presence or absence fully coincides with the strong phage resistance in the SARA1, SARA4, 574 575 SARA10, SARA19 and D23580 strains in addition to the inefficient phage infectivity pattern 576 across our panel of Salmonella strains. We postulate that a combination of genetic factors 577 rather than a single gene mutation probably drive the smaller changes in phage infective 578 efficiency (Datasets S8 and S9).



579 580

581 Figure 5. Host range of Salmonella phages and conservation of host factors involved in phage 582 infection

583 Infectivity pattern of 11 phages on a panel of 23 Salmonella strains was inferred by spotting assay. Phylogenetic relationships of the Salmonella strains were estimated by phylogenetic analysis of 115 584 585 single-copy marker genes (Methods). Gene similarity was calculated by TBLASTN search with LT2 586 proteins in SARA genomes for 184 unique gene hits uncovered in this work, and 45 (out of 184) are shown in this figure (Supplementary Datasets S4 and S8) with total number of predicted prophages, 587 588 restriction/modification proteins and gtrABC operons. Gens with premature stop codons/truncations due 589 to insertion or deletion (compared to LT2) are marked blue. Complete detail of gene similarity of 184 590 unique gene hits, prophages, restriction/modification proteins and gtrABC operons across 23 Salmonella 591 strains are given in Datasets S8 and S9 * pseudogenes and incomplete genes were excluded from the analysis, SARA2 x indicates LT2 strain. ** BTP1 prophage of D23580 contains only gtrA and gtrC genes; 592 593 # denotes genes that are not from the genetic screen. Figure 5 was created from Datasets S8 and S9.

594

595 **Discussion**

596

597 Here, we employed an unbiased RB-TnSeq loss-of-function approach to uncover the genetic determinants important in phage infection and resistance in a model enteric Salmonella species 598 599 across 11 distinct dsDNA phages, including four phages from a therapeutic formulation. In 600 addition to identifying known receptors for model Salmonella phages, our genome-wide screens 601 identify novel receptor and non-phage-receptor host factors important for a panel of dsDNA phages. We validate many of these high fitness hits via single gene deletion strains. Our results 602 indicate diverse modes of phage resistance including disruption in the phage infectivity pathway 603 downstream from phage receptors. Characterization of these non-receptor phage resistance 604 605 factors shared between two unrelated phages (indicating cross-resistance) identified an intricate interplay between alternative sigma factors pointing to how phage predation might be influenced 606 by growth and nutritional status of the cell. Finally, our host-range investigation of 11 phages 607 across a panel of 23 Salmonella strains showed differences in the infectivity pattern of some 608 609 phages despite having high conservation of top scoring hits in our genome-wide screens in a closely related model organism. Comparative analysis identified instances where sequence 610

variation in target receptor explained some of the phage susceptibility pattern, but there are additional factors and interactions both in the target host and phages that defines the host range. Overall this study highlights the importance of unbiased high-throughput genetic screens across a panel of phages in uncovering diversity of host factors important phage infection, provides insights on the genetic basis and modes of cross-resistance between sets of phages, and uncover gaps in our understanding of phage host-range across natural bacterial isolates.

617

618 Our genome-wide screens also suggest how phage selection can be used to drive beneficial tradeoffs to modulate pathogen virulence, sensitivity and fitness. For example, LPS and O-619 620 antigen play a critical role in the lifestyle of Salmonella virulence and have a myriad of effects on phage predation. Phage selection to drive truncation. loss, or reduction of LPS and O-antigen in 621 622 Salmonella could be employed to decrease virulence and increase its susceptibility to 623 antibiotics, decreased swarming motility, decreased colonization, and decreased fitness (Kong et al., 2011; Nagy et al., 2006; Toguchi et al., 2000). Our investigation into a network of cross-624 625 resistant genotypes against unrelated phages FelixO1 and Aji GE led to the role of RpoS and RpoN activity, a virulence-regulating alternative sigma factors in Salmonella sp (Chen et al., 626 627 1995; Fang et al., 1992; Wilmes-Riesenberg et al., 1997) on phage infectivity. Specifically, the association of increased RpoS activity with phage resistance raises intriguing ecological 628 questions for consideration. While RpoS activity is associated with virulence, are these phage-629 630 resistant genotypes more virulent and fit in infection contexts? Further, do these genotypes display increased RpoS activity and/or virulence in more virulent Salmonella strains? If so, the 631 selection for increased phage-resistant strains with increased virulence-associated RpoS 632 633 activity would be a deleterious outcome from therapeutic phage predation and an undesirable criterion for potentially therapeutic phages. Conversely, does the lack of phage predation 634 contribute to the neutral drift of RpoS alleles and virulence in laboratory settings (Robbe-Saule 635 et al., 1995; Wilmes-Riesenberg et al., 1997)? It is known that RpoS directly and indirectly 636 637 regulates more than 10% of all genes in E. coli (Battesti et al., 2011) and S. Typhimurium (Lago 638 et al., 2017; Lévi-Meyrueis et al., 2014), and is involved in adaptation to diverse environments and metabolic states (Battesti et al., 2011, 2015; Hengge-Aronis, 2002; Lucchini et al., 2009; 639 Nickerson and Curtiss, 1997; Robbe-Saule et al., 1995). Thus, phage resistance phenotypes 640 641 associated with RpoS activity may be acting through activation of RpoS-mediated stress response pathways rather than the direct loss of RpoS itself. In some cases, dual regulation by 642 643 genetic or nutritional factors and RpoS could lead to compensation. Some of these genotypes 644 (for example, mutants in trk, sap, ace, and rpoN) were recently associated with phage resistance in E. coli (Cowley et al., 2018; Kortright et al., 2020) and P. aeruginosa (Wright et al., 2018), but 645 646 are not yet linked to RpoS activity. Future work will explore if we see similar dependencies of alternative sigma-factors on phage resistance phenotypes in other pathogens (Dong and 647 648 Schellhorn, 2010).

649

650 Our investigation into the host-range of Salmonella phages across closely related Salmonella 651 isolates indicated that the highly fit phage resistance genotypes uncovered via genome-wide genetic screens are not complete predictors of phage infectivity and host range. Though these 652 host factors showed little variation in their sequences across our panel of Salmonella isolates, it 653 654 is possible that they vary in expression and activity, sufficient to impact phage infectivity cycle. The host-range of phages is not only defined by whether the host is susceptible to phage 655 infection, but also on how phages evade host defences and overcome barriers to efficient 656 657 infection. For example, Because O-antigen structures in Salmonella typhimurium sp. can comprise over 400 sugars per O-antigen LPS molecule(Crawford et al., 2012), it is no surprise 658 659 that many bacteriophages adsorb to this highly exposed structure. However, many

bacteriophages that adsorb to centralized outer membrane receptors can be occluded from their 660 661 native receptor by the O-antigen structure (Domínguez-Medina et al., 2020). Systematic studies exploring the form and structure of O-antigens and how they impact accessibility of phage 662 663 receptors are needed. We postulate that there are likely additional constraints affecting optimal 664 phage infectivity, and probably we might have missed uncovering these additional factors in our model strain because of highly fit phage receptor mutants. Considering differences in phage 665 666 infectivity in a panel of strains with highly conserved genetic determinants, we posit that 667 systematic study of differences in transcriptional and translation processes in these strains might provide more insights as illustrated in a few recent phage-host interaction 668 669 studies(Brandão et al., 2021; Howard-Varona et al., 2018). Future studies could also employ 670 recently developed methods (Mutalik et al., 2019; Rishi et al., 2020; Thibault et al., 2019) to provide higher resolution into phage-host interactions and may aid in filling the knowledge gaps 671 672 on phage host-range. These methods could be extended to a few closely related and 673 phylogenetically distant strains to understand the variability in host factors impacting phage 674 infectivity patterns. Finally, by combining the genetic tools developed for functional assessment of host genes with targeted or genome-wide loss-of-function mutant libraries in few model 675 676 phages, can provide additional insights into the host specificity of phages.

677

678 As high-throughput genetic screens to understand phage-host interactions grow more commonplace across diverse bacteria (Bohm et al., 2018; Christen et al., 2016; Cowley et al., 679 680 2018; Mutalik et al., 2019; Pickard et al., 2013; Price et al., 2018; Rishi et al., 2020; Rousset et al., 2018; Wetmore et al., 2015), leveraging fitness data across phages and bacterial genetic 681 diversity constitutes a major challenge and opportunity. Further screens against antibiotics, such 682 683 as those presented in earlier (Price et al., 2018), could rapidly discover collateral sensitivity 684 patterns wherein phage resistant genotypes display sensitization to antibiotics or ecologically-685 relevant conditions (for instance sera or bile salts). Such information has the potential to form the basis of successful combinations of treatments (Chan et al., 2018; Kortright et al., 2019; 686 Mangalea and Duerkop, 2020). We posit that phage-host interaction studies across diverse 687 688 bacterial isolates in a range of biotic and abiotic conditions powered with novel transcriptomics and proteomics tools can provide rich datasets for host-range predictive models and rational 689 phage cocktails formulations. 690

- 691
- 692

693 Materials and Methods

694

695 **Bacterial strains and growth conditions**

Strains, primers, and plasmids are listed in Tables S3-S5, respectively. S. Typhimurium LT2 696 derivative strain MS1868 genotype is S. Typhimurium LT2 (leuA414(Am) Fels2- hsdSB(r-697 m+))(Graña et al., 1985). In general, all Salmonella strains were grown in Luria-Bertani (LB-698 699 Lennox) broth (Sigma) at 37°C, 180 rpm unless stated otherwise. When appropriate, 50 µg/mL 700 kanamycin sulfate and/or 34 µg/mL chloramphenicol were supplemented to media. For strains 701 containing an ampR selection marker, carbenicillin was employed at 100 µg/mL, but exclusively 702 used during isolation of clonal mutants to avoid mucoidy phenotypes. All bacterial strains were 703 stored at -80°C for long term storage in 25% sterile glycerol (Sigma).

704

705 Bacteriophages and propagation

706 Bacteriophages employed in this study and sources are listed in Table 1. All phages were either 707 successively serially diluted or streaked onto 0.7% LB-agar overlays for isolation. For 708 bacteriophage Chi, 0.35% LB-agar overlays were employed. Bacteriophage Aji GE EIP16, 709 Reaper GE 8C2, Savina GE 6H2, and Shishito GE 6F2 were isolated from a commercial 710 bacteriophage formulation from Georgia. All phages isolated from this source are denoted with "_GE" (to recognize being sourced from Georgia). All other bacteriophages were re-isolated 711 from lysates provided from stock centers or gifts from other labs (Table 1). Bacteriophage 712 Aii GE EIP16, Chi, FelixO1, P22 (a strictly lytic mutant), Reaper GE 8C2, S16, and SP6 were 713 isolated and scaled on S. Typhimurium MS1868. Bacteriophage Br60, Ffm, Savina GE 6H2, 714 715 and Shishito GE 6F2 were isolated and rough-LPS mutant S. Typhimurium. SL733 (BA1256). 716 We followed standard protocols for propagating phages (Kutter and Sulakvelidze, 2004). Br60, Chi, Ffm, P22, Reaper_GE_8C2, S16, Savina_GE_6H2, Shishito_GE_6F2, and SP6 were 717 718 propagated in LB-Lennox liquid culture on their respective strains. Reaper GE 8C2, Savina_GE_6H2, and Shishito_GE_6F2 were additionally buffer-exchanged into SM-Buffer 719 720 (Teknova) via ultrafiltration (Amicron 15) and resuspension. Bacteriophage Aii GE EIP16 and FelixO1 were propagated in LB-Lennox liquid culture on their respective strains and further 721 propagated through a standard overlay method. Whenever applicable, we used SM buffer 722 723 without added salts (Tekova) as a phage resuspension or dilution buffer and routinely stored 724 phages as filter-sterilized (0.22um) lysates at 4°C.

725

726 Bacteriophages Aji GE EIP16, Reaper GE 8C2, Savina_GE_6H2, and Shishito_GE_6F2 were additionally whole-genome sequenced and assembled. Approximately 1e9 PFU of phage 727 lysate was gDNA extracted through Phage DNA Isolation Kit (Norgen, 46800) as per 728 manufacturer's instructions. Library preparation was performed by the Functional Genomics 729 730 Laboratory (FGL), a QB3-Berkeley Core Research Facility at UC Berkeley. Sequencing was 731 performed at the Vincent Coates Sequencing Center, a QB3-Berkeley Core Research Facility at UC Berkeley on a MiSeq using 75PE runs for Reaper GE 8C2, Savina GE 6H2, and 732 Shishito_GE_6F2 and using 150SR run for Aji_GE_EIP16. Phage genomes were assembled 733 using KBase (Arkin et al., 2018). Illumina reads were trimmed using Trimmomatic v0.36 (Bolger 734 735 et al., 2014) and assessed for quality using FASTQC. Trimmed reads for Aji_GE_EIP16, Reaper GE 8C2, and Shishito GE 6F2 were assembled using Spades v3.13.0 (Nurk et al., 736 2013). Trimmed reads for Savina GE 6H2 were assembled using Velvet v1.2.10 (Zerbino and 737 738 Birney, 2008). The primary, high coverage contig from these assemblies was investigated and 739 corrected for incorrect terminus assembly using PhageTerm v1.011 on CPT Galaxy (Garneau et al., 2017). In this manuscript, we limited analyses of these sequences to assessing phylogeny of 740 741 these phages, which we performed with BLASTN (Dataset S7). A detailed genomic

characterization will be published by Dr. Elizabeth Kutter. Sequences and preliminary
annotations can be found at JGI IMG under analysis projects Ga0451357, Ga0451371,
Ga0451358, and Ga0451372.

745

746 Construction of MS1868 RB-TnSeq library

747 We created the Salmonella enterica serovar Typhimurium MS1868 (MS1868_ML3) transposon 748 mutant library by conjugating with E. coli WM3064 harboring pHLL250 mariner transposon 749 vector library (strain AMD290) (Figure S1). To construct pHLL250, we used the magic pools 750 approach we outlined previously(Liu et al., 2018). Briefly, pHLL250 was assembled via Golden 751 Gate assembly using BbsI from part vectors pHLL213, pHLL216, pHLL238, pHLL215, and 752 pJW14(Liu et al., 2018). We then incorporated millions of DNA barcodes into pHLL250 with a 753 second round of Golden Gate assembly using BsmBI. Briefly, we grew S. Typhimurium LT2 754 MS1868 at 30°C to mid-log-phase and combined equal cell numbers of S. Typhimurium LT2 755 MS1868 and donor strain AMD290, conjugated them for 5 hrs at 30°C on 0.45-µm nitrocellulose 756 filters (Millipore) overlaid on LB agar plates containing diaminopimelic acid (DAP) (Sigma). The 757 conjugation mixture was then resuspended in LB and plated on LB agar plates with 50 ug/ml 758 kanamycin to select for mutants. After 1 day of growth at 30°C, we scraped the kanamycin-759 resistant colonies into 25 mL LB and processed them as detailed earlier to make multiple 1-mL -760 80°C freezer stocks. To link random DNA barcodes to transposon insertion sites, we isolated 761 the genomic DNA from cell pellets of the mutant libraries with the DNeasy kit (Qiagen) and 762 followed published protocol to generate Illumina compatible sequencing libraries(Wetmore et al., 763 2015). We then performed single-end sequencing (150 bp) with the HiSeq 2500 system (Illumina). Mapping the transposon insertion locations and the identification of their associated 764 765 DNA barcodes was performed as described previously (Price et al., 2018). In total, our 66,996 766 member pooled library consisted of transposon-mediated disruptions in 3,759 out of 4,610 767 genes, with an average of 14.8 disruptions per gene (median 12). Compared to a non-barcoded reported transposon mutant library in S. Typhimurium 14028s (Porwollik et al., 2014), we 768 769 suspect 434 of the 851 unmutated genes are likely essential, and 380 likely nonessential. We 770 abstain from interpreting essentiality of 37 additional genes due to inability to uniquely map insertions or due to gene content differences between the two libraries. Additional details for the 771 772 composition of the S. Typhimurium MS1868 library can be found in Table S1 and Dataset S1.

773

774 Liquid culture "competitive" fitness experiments

775 Competitive, phage-stress fitness experiments were performed in liquid culture, as phage 776 progeny from an infection of one genotype could subsequently infect other host genotypes. All 777 bacteriophages were tested against the MS1868 library. Bacteriophage Br60, Ffm, and 778 Shishito_GE_6F2 were additionally tested against the previously described E. coli BW25113 779 library (Wetmore et al., 2015). To avoid jackpot effects, at least two replicate experiments were 780 performed per phage-host library experiment as presented earlier (Mutalik et al., 2020). Briefly, 781 a 1 mL aliquot of RB-TnSeq library was gently thawed and used to inoculate a 25 mL of LB supplemented with kanamycin. The library culture was allowed to grow to an OD600 of ~1.0 at 782 783 37°C. From this culture we collected three, 1 mL pellets, comprising the 'Time-0' or reference samples in BarSeq analysis. The remaining cells were diluted to a starting OD600 of 0.04 in 2X 784 785 LB with kanamycin. 350 µL of cells were mixed with 350 µL phage diluted in SM buffer to a 786 predetermined MOI and transferred to a 48-well microplate (700 µL per well) (Greiner Bio-One #677102) covered with breathable film (Breathe-Easy). Phage infection progressed in Tecan 787 Infinite F200 readers with orbital shaking and OD600 readings every 15 min for 3 hours at 37°C. 788 789 At the end of the experiment, each well was collected as a pellet individually. All pellets were 790 stored at -80°C until prepared for BarSeq.

791

792 Solid agar "noncompetitive" fitness experiments

793 Noncompetitive, phage-stress fitness experiments were performed on solid-agar plate culture as 794 presented earlier (Mutalik et al., 2020). Solid plate fitness experiments were performed by 795 assaving all 11 bacteriophages against the MS1868 library. Bacteriophage Br60, Ffm, and 796 Shishito_GE_6F2 were additionally assayed on the *E. coli* BW25113 library (Wetmore et al., 797 2015). For the solid plate experiments a 1 mL aliquot of the RB-TnSeq library was gently 798 thawed and used to inoculate a 25 mL LB supplemented with kanamycin. The library culture 799 was allowed to grow to an OD600 of ~1.0 at 37°C. From this culture we collected three, 1 mL 800 pellets, comprising the 'Time-0' for data processing in BarSeq analysis. The remaining cells were diluted to a starting OD600 of 0.01 in LB with kanamycin. 75 µL of cells were mixed with 801 802 75 µL of phage diluted in SM buffer to a predetermined MOI and allowed to adsorb for 10 803 minutes. The entire culture was spread evenly over a LB agar plate with kanamycin and grown 804 overnight at 37°C. The next day, all resistant colonies were collected and suspended in 1.5 mL 805 LB media before pelleting. All pellets were then stored at -80°C until prepared for BarSeq.

806

807 BarSeq of RB-TnSeq pooled fitness assay samples

Genomic DNA was isolated from stored pellets of enriched and 'Time 0' RB-TnSeg samples 808 809 using the DNeasy Blood and Tissue kit (Qiagen). We performed 98°C BarSeg PCR protocol as described previously (Mutalik et al., 2020; Wetmore et al., 2015). BarSeq PCR in a 50 uL total 810 811 volume consisted of 20 umol of each primer and 150 to 200 ng of template genomic DNA. For the HiSeq4000 runs, we used an equimolar mixture of four common P1 oligos for BarSeq, with 812 variable lengths of random bases at the start of the sequencing reactions (2-5 nucleotides). 813 814 Equal volumes (5 uL) of the individual BarSeq PCRs were pooled, and 50 uL of the pooled PCR product was purified with the DNA Clean and Concentrator kit (Zymo Research). The final 815 816 BarSeq library was eluted in 40 uL water. The BarSeq samples were sequenced on Illumina 817 HiSeq4000 instruments with 50 SE runs. Typically, 96 BarSeq samples were sequenced per lane of HiSeq. 818

819

820 Data processing and analysis of BarSeq reads

821 Fitness data for the RB-TnSeq library was analyzed as previously described (Wetmore et al., 822 2015). Briefly, the fitness value of each strain (an individual transposon mutant) is the 823 normalized log2(strain barcode abundance at end of experiment/strain barcode abundance at 824 start of experiment). The fitness value of each gene is the weighted average of the fitness of its 825 strains. Further analysis of BarSeq data was carried out in Python3 and visualized employing 826 matplotlib and seaborn packages. For heatmap visualizations, genes with under 25 BarSeg 827 reads in the phage samples had their fitness values manually set to 0 to avoid artificially high 828 fitness scores (due to the strong selection pressure imposed by phage predation).

829

830 Due to the strong selection pressure and subsequent fitness distribution skew resulting from phage infection, a couple additional heuristics were employed during analysis. Initially, per 831 phage experiment, fitness scores were filtered for log2-fold-change thresholds, aggregated read 832 counts, and t-like-statistics. Experiments using phages Ffm, Shishito GE, and Br60 (which 833 834 cannot infect wild-type MS1868, but can infect specific MS1868 mutants) against the MS1868 835 library employed negative thresholds to identify sensitized genotypes. A summary of log2-foldchange fitness and t-like statistic thresholds are provided in the Dataset S2. Each reported hit 836 837 per phage was further processed via manual curation to minimize reporting of false-positive 838 results due to the strong phage selection pressure. Here, all individual barcodes per genotype 839 were investigated simultaneously for each experiment through both barcode-level fitness scores 840 and raw read counts. First, genotypes were analyzed for likely polar effects. If the location and 841 orientation of each fit barcode were exclusively against the orientation of transcription and/or exhibited strong fitness at the C-terminus of a gene, while being transcriptionally upstream of 842 843 another fit gene, the genotype was likely a polar effect and eliminated. Second, genotypes were 844 analyzed for jackpot fitness effects that could indicate a secondary site mutation. These cases 845 were identified by investigating consistency between individual strains within a genotype. If the 846 vast majority of reads per genotype belonged to a singular mutant (of multiple), we attributed the 847 aggregate fitness score to secondary-site mutation effects and eliminated those genotypes from 848 reported results. Genotypes where there were too few strains to make a judgment call on within 849 genotype strain consistency (ie 1-3 barcodes) were generally excluded from analysis unless 850 they were genotypes consistent with other high-scoring genotypes. Next, we investigated for 851 consistency between read counts and fitness scores at both the strain level. In general, we 852 found that strains with read counts under 25 often had inflated fitness scores under strong 853 phage selection pressure and the subsequent fitness distribution skew resulting from phage 854 infection. Cases where high fitness scores were attributed to a couple of strains with reads under 25 were eliminated as false positives as well. Finally, all genotypes were loosely curated 855 856 for consistency across liquid experiments. Cases that barely passed confidence thresholds as 857 described above that were inconsistent across replicate experiments were eliminated from 858 reporting. A summary of fit genotypes that passed automated filtering and manual curation are 859 reported in Dataset S4. No fit genotypes were added during manual analyses.

860

Network graphs were constructed using Gephi. Graph layout optimization was determined through a combination of manual placement of nodes (for instance phage nodes in Figure 3B) and layout optimization based off of equally weighted edges using the Yifan Hu algorithm. In all graphs, edges were calculated based on Dataset S4 using custom python scripts. In brief, in the mixed node graph in Figure 3B, edges were drawn with weight 1 between a phage node (fixed) and a gene node if that gene conferred resistance according to Dataset S4.

867

868 Individual Mutant Creation

All individual deletion mutants in *S*. Typhimurium were created through lambda-red mediated genetic replacement (Sawitzke et al., 2007). Per deletion, primers were designed to PCR amplify either kanamycin or ampicillin selection markers with ~30-40 bp of homology upstream and downstream of the targeted gene locus, leaving the native start and stop codons intact preserving directionality of gene expression at the native locus (Table S3). PCRs were generated and gel-purified through standard molecular biology techniques and stored at -20°C until use. All strains (including mutants) employed in this study are listed in Table S5.

876

877 Deletions were performed by incorporating the above dsDNA template into the Salmonella 878 genome through standard pSIM5-mediated recombineering methods (Sawitzke et al., 2007). 879 First temperature-sensitive recombineering vector, pSIM5, was introduced into the relevant 880 Salmonella strain through standard electroporation protocols and grown with chloramphenicol at 30°C. Recombination was performed through electroporation with an adapted pSIM5 881 recombineering protocol. Post-recombination, clonal isolates were streaked onto plates without 882 883 chloramphenicol at 37°C to cure the strain of pSIM5 vector, outgrown at 37°C and stored at -884 80°C until use. For double deletions, this process was repeated two times in series with kanamycin followed by ampicillin selection markers. Gene replacements were verified by colony 885 886 PCR followed by Sanger sequencing at the targeted locus (both loci if a double deletion mutant) 887 and 16S rDNA regions (primers provided in Table S3).

888

889 Assessing Phage Sensitivity

890 Phage-resistance and -sensitivity was assessed through efficiency of plating experiments. Bacterial hosts were grown overnight at 37°C. 100 µL of these overnight cultures were added to 891 892 5 mL of top-agar with appropriate antibiotics and allowed to solidify at room temperature. For 893 assays including supplements such as glutamine, the supplement was added directly to the top agar layer. Phages were ten-fold serially diluted in SM Buffer, two microliters spotted out on the 894 895 solidified lawn, and incubated the plates overnight at 37°C. Efficiency of plating was calculated 896 as the ratio of the average effective titer on the tested host to the titer on the propagation host. 897 For some assay strains, plagues showed diffused morphology and were difficult to count, or 898 displayed plaque phenotypes distinct from its propagation host. In all cases, representative 899 images are presented (Figures S5-S11, S13-S15). All plaquing experiments were performed 900 with at least three biological replicates, each replicate occurring on a different day from a 901 different overnight host culture.

902

903 **RNA-Seq experiments**

Samples for RNA-Seq analysis were collected and analyzed for wild-type MS1868 (BA948) 904 905 (N=3), knockout mutants for trkH (BA1124) (N=3) sapB (BA1136) (N=3), rpoN (BA1139) (N=3), and himA (BA1142) (N=2). All cultures for RNA-Seq were grown on the same day from unique 906 907 overnights and subsequent outgrowths. Strains were diluted to OD600 ~0.02 in 10 mL LB with 908 appropriate selection marker, and then grown at 30°C at 180 RPM until they reached an OD600 909 0.4-0.6. Samples were collected as follows: 400 µL of culture was added to 800 µL RNAProtect 910 (Qiagen), incubated for 5 minutes at room temperature, and centrifuged for 10 minutes at 5000xg. RNA was purified using RNeasy RNA isolation kit (Qiagen) and quantified and quality-911 912 assessed by Bioanalyzer. Library preparation was performed by the Functional Genomics 913 Laboratory (FGL), a QB3-Berkeley Core Research Facility at UC Berkeley. Illumina Ribo-Zero 914 rRNA Removal Kits were used to deplete ribosomal RNA. Subsequent library preparation steps 915 of fragmentation, adapter ligation and cDNA synthesis were performed on the depleted RNA using the KAPA RNA HyperPrep kit (KK8540). Truncated universal stub adapters were used for 916 917 ligation, and indexed primers were used during PCR amplification to complete the adapters and to enrich the libraries for adapter-ligated fragments. Samples were checked for quality on an 918 919 Agilent Fragment Analyzer, but ribosome integrity numbers were ignored. This is routine for 920 Salmonella sp., since they natively have spliced 23S rRNA (Burgin et al., 1990). Sequencing 921 was performed at the Vincent Coates Sequencing Center, a QB3-Berkeley Core Research 922 Facility at UC Berkeley on a HiSeq4000 using 100PE runs.

923

924 RNA-Seq Data Analysis

925 For all RNA-Seq experiments, analyses were performed through a combination of KBase-(Arkin 926 et al., 2018) and custom jupyter notebook-based methods. The data processing narrative in 927 KBase can be found here: https://kbase.us/n/48675/70/. StringTie and DESeg2 KBase outputs 928 are currently available in Datasets S5 and S6 (https://doi.org/10.6084/m9.figshare.12185031). Briefly, Illumina reads were trimmed using Trimmomatic v0.36 (Bolger et al., 2014) and 929 assessed for quality using FASTQC. Trimmed reads were subsequently mapped to the S. 930 931 Typhimurium LT2 along with PSLT genome (NCBI Accession: AE006468.2 and AE006471.2 932 respectively) with HISAT2 v2.1.0 (Kim et al., 2019). Alignments were quality-assessed with 933 BAMQC. From these alignments, transcripts were assembled and abundance-estimated with StringTie v1.3.3b (Pertea et al., 2015). Tests for differential expression were performed on 934 normalized gene counts by DESeq2 (negative binomial generalized linear model) (Love et al., 935 936 2014). Additional analyses for all experiments were performed in Python3 and visualized employing matplotlib and seaborn packages. Conservative thresholds were employed for 937

assessing differentially expressed genes. Conclusions were considered differentially expressed
 if they possessed a Bonferoni-corrected p-value below a threshold of 0.001 and an absolute
 log2 fold change greater than 2. Assembled transcripts from StringTie and differential
 expression from RNA-Seg analyses can be found in Datasets S5 and S6 respectively.

942

943 Genome sequencing of SARA collection

944 We sequenced the 21 reference S. Typhimurium genomes (Beltran et al., 1991) using standard 945 molecular biology protocols. Briefly, we grew up all 21 strains to stationary phase in LB media. 946 We then extracted gDNA using the DNeasy Blood and Tissue kit (Qiagen). Illumina library 947 preparation was performed by the Functional Genomics Laboratory (FGL), a QB3-Berkeley Core Research Facility at UC Berkeley. Sequencing was performed at the Vincent Coates 948 Sequencing Center, a QB3-Berkeley Core Research Facility at UC Berkeley on a HiSeq4000 949 950 using 100PE reads. We used Unicycler with default parameters(Wick et al., 2017) to do a 951 reference based assembly from closely related Salmonella strains.

952 Bioinformatic analysis of SARA collection genomes

953 Predicted genes in 24 Salmonella typhimurium genomes were classified in families of 954 homologous genes by PPanGGoLiN(Gautreau et al., 2020). Gene clusters encoding LPS core 955 oligosaccharide and O-specific antigen (OSA) biosynthetic enzymes were identified in the 956 Salmonella genomes by search for gene families containing characterized LPS and OSA 957 biosynthesis genes of LT2 strain (STM2079-STM2098, STM3710-STM3723)(Heinrichs et al., 958 1998: Seif et al., 2019), O-antigen modification genes were identified by DIAMOND similarity 959 search(Buchfink et al., 2015) with characterized LT2 proteins OpvA (STM2209), OpvB 960 (STM2208), GtrA (STM0559, STM4204), GtrB(STM0558, STM4205)(Broadbent et al., 2010) 961 using blastp command with --very-sensitive option. Restriction/modification genes were identified by DIAMOND similarity search with 78008 proteins from REBASE database(Roberts 962 963 et al., 2015). Point mutations in LPS and OSA biosynthesis enzymes were identified by running 964 a command-line application for TBLASTN search(Camacho et al., 2009) of LT2 proteins vs. 965 genome sequences of 23 Salmonella typhimurium genomes.

966 **Phylogenetic analysis**

967 To estimate phylogenetic relationships between genomes of our collection of Pseudomonas spp. strains, we identified a set of 120 bacterial marker genes with GTDB-Tk toolkit(Chaumeil et 968 969 al., 2019). Only 115 marker genes were found in single copy in each of the 24 genomes studied. 970 Gene sequences of those 115 markers were aligned by MAFFT v7.310(Katoh and Standley, 2014) with --auto option, and the resulting 88 alignments were concatenated into a single 971 multiple sequence alignment. A phylogenetic tree was reconstructed from the multiple alignment 972 973 using the maximum likelihood method and generalized time-reversible model of nucleotide 974 substitution implemented in the FastTree software v2.1.10(Price et al., 2010) and visualized 975 using the Interactive Tree of Life (iTOL) online tool(Letunic and Bork, 2019). 976

977 Prophage analysis

To determine prophage content, we submitted all contigs longer than 10kb to the PHASTER web server, culminating in 250 potential prophage regions across the 21 strains investigated(Arndt et al., 2016) (Dataset S9). These 250 identified regions were aligned against each other using nucmer(Marçais et al., 2018). Grouping and subsequent filtering of prophages was performed through network graph analysis using Gephi; prophage nodes were connected by edges representing total nucmer alignments greater than 60% alignment. Graph layout optimization was determined through layout optimization based off of equally weighted edges

using the Yifan Hu algorithm. For each "cluster" of prophages and each alone prophage, a few
representatives were investigated manually for prophage similarity to determine if a PHASTERidentified region (or "cluster") was correctly identified as a prophage, yielding 84 likely prophage
regions. Based on similarity to studied prophages, we assigned each "cluster" to one of "ST64B
(118970_sal3-like)", "Gifsy-1", "Gifsy-2", "Gifsy-3", "Fels-1", "P2-like", "P22-like", "phiKO2-like",
"SPN1S-like" classifications (Dataset S9).

991

992 Because this prophage determination was based upon a reference-based assembly(Wick et al., 993 2017), it was possible for some regions to mis-assemble depending on the reference genome 994 used. So, we further validated if these prophage regions were artifacts of assembly. For each 995 genome, we re-aligned our reads to the assembled genome using samtools and noted all 996 regions that were not covered in BAM-alignments. We noted if prophages were either (1) split 997 across contigs (common for "Gifsy-2"), (2) not covered by reads (noted 2 instances for P22-like 998 prophages), (3) partially not covered by reads (common for P22-like phages, which have known 999 mosaic sequences)(Fu et al., 2017) and (4) compared our prophage identification efforts to 1000 earlier work (Fu et al., 2017). After eliminating prophage regions that were assembly artifacts, we 1001 culminated in 74 high confidence prophage regions across the 21 SARA strains (Dataset S9).

1002

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 1016 (University of California at Berkeley). Sequencing was performed at: Vincent J. Coates
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 1018 Instrumentation Grants S10RR029668, S10RR027303, and OD018174.
- 1019

1020 Author Contributions

- 1021 B.A.A., V.K.M., and A.P.A. conceived the project.
- 1022 B.A.A. led the experimental work, analysis, and manuscript preparation.
- 1023 B.A.A., V.K.M., C.Z., A.M.D. and H.L. built and characterized the RB-TnSeq library.
- 1024 B.A.A. performed experiments, processed, and analyzed data.
- 1025 E.B.K. provided critical reagents and advice.
- 1026 T.N.N., L.M.L. assembled genomes.
- 1027 B.A.A., E.B.K., A.M.D., V.K.M., and A.P.A. wrote the paper.
- 1028

1029 **Competing Interests**

- 1030 V.K.M., A.M.D., and A.P.A. consult for and hold equity in Felix Biotechnology Inc..
- 1031
- 1032
- 1033

1034 Data Availability

- 1035 Supplementary Information can be found here:
- 1036 https://doi.org/10.6084/m9.figshare.12185001.v2. Complete Supplementary Datasets can be
- 1037 found here: https://doi.org/10.6084/m9.figshare.12185031. Supplementary Code and figure
- reproduction can be found here: https://doi.org/10.6084/m9.figshare.12412814.v2. All NGS
- 1039 reads have been deposited and made publicly accessible via the Sequence Read Archive
- 1040 (SRA) under Bioproject PRJNA638761: http://www.ncbi.nlm.nih.gov/bioproject/638761. Draft
- 1041 S.Typhimurium genome sequences for strains SARA1-SARA21 can be found under
- 1042 BioSamples SAMN17506935-SAMN17506955. Draft phage genome sequences and JGI-
- 1043 performed gene annotations can be found at JGI IMG under analysis projects Ga0451357,
- 1044 Ga0451371, Ga0451358, and Ga0451372. The RNA-Seq data processing narrative in KBase
- 1045 can be found here: https://kbase.us/n/48675/70/.

1046 Table 1. Bacteriophages employed in this study.

1047 Virus families were assigned via ICTV taxonomy release 2019. For new phages isolated in this

- study, the family of the nearest BLASTN relative was reported (in line with ICTV 2019
- 1049 standards). This information can be found in Dataset S7.

otanidaido			Databot 07:	1
Phage	Family	Established Receptor?	Source	Reference
Aji_GE_ EIP16 (Aji_GE)	Demerecvirid ae	No	Intesti Bacteriophage formulation M2-601	This study.
Br60	Autographiviri dae	"Rough LPS <i>Salmonella</i> "	Salmonella Genetic Stock Center (SGSC)	(Wilkinson et al., 1972)
Chi	Siphoviridae	Flagella	Gift from Jason Gill	(Samuel et al., 1999; Schade et al., 1967)
FelixO1	Myoviridae	Outer core LPS	Félix d'Hérelle Reference Center for Bacterial Viruses	(Gebhart et al., 2017; Lindberg and Hellerqvist, 1971; Tu et al., 2017)
Ffm	Autographiviri dae	"Rough-LPS Salmonella"	<i>Salmonella</i> Genetic Stock Center (SGSC)	(Graña et al., 1985; Lee et al., 2013; Lindberg and Hellerqvist, 1971; MacPhee et al., 1975; Marti et al., 2013; Schwartz, 1980; Wilkinson et al., 1972; Wright et al., 1980)
P22	Podoviridae	O-Antigen LPS	Gift from Richard Calendar	(Bohm et al., 2018; Steinbacher et al., 1997; Wright et al., 1980)
Reaper_ GE_8C2 (Reaper _GE)	Siphoviridae	No	Intesti Bacteriophage formulation M2-601	This study
S16	Myoviridae	OmpC	Félix d'Hérelle Reference Center for Bacterial Viruses	(Marti et al., 2013)
Savina_ GE_6H2 (Savina_ GE)	Myoviridae	No	Intesti Bacteriophage formulation M2-601	This study
Shishito _GE_6F 2 (Shishito _GE)	Autographiviri dae	No	Intesti Bacteriophage formulation M2-601	This study
SP6	Autographiviri dae	O-Antigen LPS	Gift from lan J Molineaux	(Tu et al., 2017; Wright et al., 1980)

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