1	The tryptophan salvage pathway is dynamically regulated by the
2	iron-dependent repressor YtgR in Chlamydia trachomatis
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## 24 Abstract

25 Mammalian hosts restrict cellular nutrient availability to starve invading pathogens and 26 successfully clear an infection by a process termed "nutritional immunity". For the 27 obligate intracellular pathogen Chlamydia trachomatis, nutritional immunity likely 28 encompasses the simultaneous limitation of the amino acid tryptophan and the essential 29 biometal iron. Unlike other model bacteria, C. trachomatis lacks many global stress 30 response systems – such as "stringent response" homologs – adapted to these host 31 insults. However, a physiological response by Chlamydia that is common to both stresses is the development of an aberrant, "persistent" state, suggesting that 32 33 tryptophan and iron starvation trigger a coordinated developmental program. Here, we 34 report that the trpRBA operon for tryptophan salvage in C. trachomatis serovar L2 is 35 regulated at the transcriptional level by iron. The expression of the tryptophan synthase 36 encoding genes, trpBA, is induced following iron limitation while that of the repressor 37 *trpR* is not. We show that this specific induction of *trpBA* expression initiates from a 38 novel promoter element within an intergenic region flanked by trpR and trpB. YtgR, a DtxR-homolog and the only known iron-dependent transcriptional regulator in 39 40 Chlamydia, can bind to the trpRBA intergenic region upstream of the alternative trpBA promoter to repress transcription. This binding also likely attenuates transcription from 41 the primary promoter upstream of *trpR* by blocking RNA polymerase read-through. 42 43 These data illustrate a dynamic and integrated method of regulating tryptophan biosynthesis in an iron-dependent manner, which has not been described in any other 44 45 prokaryote, underscoring the uniqueness of *Chlamydia*.

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## 47 Significance Statement

48	Genital serovars of the obligate intracellular parasite Chlamydia trachomatis are the
49	leading cause of bacterial sexually-transmitted infections globally. Proliferation of C.
50	trachomatis is likely controlled by simultaneous immunological and environmental
51	restriction of critical nutrients such as tryptophan and iron. However, our understanding
52	of the immediate chlamydial responses to these stimuli is poorly defined. We utilized
53	expression of the stress-responsive trpRBA operon in C. trachomatis L2 as a proxy for
54	regulatory integration between iron and tryptophan limitation. We identified a unique
55	iron-dependent regulatory mechanism for trpRBA in C. trachomatis, mediated by the
56	transcriptional repressor YtgR. This distinguishes Chlamydia from other bacteria which
57	regulate tryptophan biosynthesis strictly by tryptophan-dependent mechanisms,
58	highlighting a distinct evolutionary adaptation in C. trachomatis to integrate stress
59	responses.
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## 70 /body

## 71 Introduction

72 Nutrient acquisition is critical for the success of pathogenic bacteria. Many 73 pathogenic bacteria must siphon nutrients from their hosts, such as nucleotides, amino 74 acids and biometals (1–4). This common feature among pathogens renders them 75 susceptible to nutrient limitation strategies associated with the host immune response 76 (5). Counteractively, bacterial pathogens have evolved sophisticated molecular 77 mechanisms to counter nutrient deprivation, involving increasingly complex and 78 sophisticated nutrient-sensing regulatory networks. These stress response mechanisms 79 are essential for pathogens to avoid clearance by the immune system. By delineating 80 their function at the molecular level, we can better target aspects of the host-pathogen 81 interface suitable for the rapeutic manipulation. However, stress responses in the 82 obligate intracellular bacterium *Chlamydia trachomatis* are relatively poorly 83 characterized, leaving unanswered many fundamental questions about the biology of 84 this pathogen.

Chlamydia trachomatis is the leading cause of bacterial sexually transmitted 85 86 infections (STIs) and infection-derived blindness worldwide (6–8). Genital infections of 87 chlamydia are associated with serious sequelae in the female reproductive tract such as 88 tubal factor infertility (9). Chlamydiae are Gram-negative bacterial parasites that develop 89 within a pathogen-specified membrane-bound organelle termed the inclusion (10). 90 Chlamydial development is uniquely characterized by a biphasic interconversion of an infectious elementary body (EB) with a non-infectious, but replicative reticulate body 91 92 (RB) (11). An obligate intracellular lifestyle has led to reductive genome evolution

across chlamydial species; Chlamydiae have retained genes uniquely required for their 93 94 survival, but have become nutritionally dependent on their hosts by discarding many 95 metabolism-related genes (12). Of note, C. trachomatis does not possess genes 96 necessary for eliciting a stringent response to nutrient starvation (e.g. relA, spoT), 97 suggesting that this pathogen may utilize novel mechanisms to respond to stress (13). 98 It is well established, however, that in response to various stressors, Chlamydiae 99 deviate from their normal developmental program to initiate an aberrant developmental 100 state, termed "persistence" (14). This persistent state is distinguished by the presence 101 of viable, but non-cultivable, abnormally enlarged chlamydial organisms that display 102 dysregulated gene expression. Importantly, Chlamydia can be reactivated from 103 persistence by abatement of the stress condition. As such, chlamydial persistence at 104 least superficially resembles a global stress response mechanism. Yet the molecular 105 underpinnings of this phenotype are poorly understood, with most published studies 106 focusing on the molecular and metabolic character of the aberrant, persistent form. It is 107 therefore unclear to what extent primary stress responses contribute to the global 108 persistent phenotype in *Chlamydia*.

The best described inducer of persistence is the pro-inflammatory cytokine interferon-gamma (IFN- $\gamma$ ). The bacteriostatic effect of IFN- $\gamma$  has been primarily attributed to host cell tryptophan (Trp) catabolism, for which *C. trachomatis* is auxotrophic (15–17). Following IFN- $\gamma$  stimulation, infected host cells up-regulate expression of indoleamine-2,3-dioxygenase (IDO1), which catabolizes Trp to *N*formylkynurenine via cleavage of the indole ring (18). *C. trachomatis* cannot recycle kynurenines, unlike some other chlamydial species (19), and thus IFN- $\gamma$  stimulation effectively results in Trp starvation to *C. trachomatis*. The primary response to Trp
starvation in *C. trachomatis* is mediated by a TrpR ortholog, whose Trp-dependent
binding to cognate promoter elements represses transcription (20, 21). This
mechanism of regulatory control is presumably limited in *C. trachomatis*, as homologs of
genes regulated by TrpR in other bacteria (*e.g. trpF, aroH, aroL*) have not been shown
to respond to Trp limitation (22).

In many Gram-negative bacteria, such as *Escherichia coli*, trpR is monocistronic 122 123 and distal to the Trp biosynthetic operon. In *C. trachomatis*, TrpR is encoded in an operon, *trpRBA*, which also contains the Trp synthase  $\alpha$ - and  $\beta$ - subunits (TrpA and 124 125 TrpB, respectively), and possesses a 351 base-pair (bp) intergenic region (IGR) that 126 separates *trpR* from *trpBA*. The functional significance of the *trpRBA* IGR is poorly 127 characterized; while a putative operator sequence was identified overlapping an 128 alternative transcriptional origin for *trpBA* (20), the IGR was not shown to be bound by 129 TrpR (21). Based on in silico predictions, an attenuator sequence has been annotated 130 within the *trpRBA* IGR (23), but this has not been thoroughly validated experimentally. Regardless, the IGR is more than 99% conserved at the nucleotide sequence level 131 132 across ocular, genital and lymphogranuloma venereum (LGV) serovars of C. 133 trachomatis, indicating functional importance (Fig. S1). Therefore, relative to other 134 model bacteria, the regulation of the *trpRBA* operon remains poorly elucidated. 135 In evaluating alternative regulatory modes of the *trpRBA* operon, an interesting consideration is the pleiotropic effects induced by IFN-y stimulation of infected cells. 136 IFN- $\gamma$  is involved in many processes that limit iron and other essential biometals to 137 intracellular pathogens as a component of host nutritional immunity (5, 24). Chlamydia 138

139 have a strict iron dependence for normal development, evidenced by the onset of 140 persistence following prolonged iron limitation (25). Importantly, *Chlamydia* presumably 141 acquire iron via vesicular interactions between the chlamydial inclusion and slow-142 recycling transferrin (Tf)-containing endosomes (26). IFN- $\gamma$  is known to down-regulate 143 transferrin receptor (TfR) expression in both monocytes and epithelial cells with replicative consequences for resident intracellular bacteria (27–30). However, iron 144 145 homeostasis in *Chlamydia* is poorly understood, due to the lack of functionally 146 characterized homologs to iron acquisition machinery that are highly conserved in other 147 bacteria (31). Only one operon, represented by *ytqABCD*, has been clearly linked to iron 148 acquisition. The periplasmic metal-binding protein YtgA displays a specific binding 149 affinity for ferric iron over other divalent cations and likely transports iron from the outer membrane to an ABC-3 type inner membrane metal permease formed by the YtgBCD 150 151 complex (32). Intriguingly, the YtgC (CTL0323) open reading frame (ORF) encodes a N-152 terminal permease domain fused to a C-terminal DtxR-like repressor domain, annotated 153 YtgR (33, 34). YtgR is cleaved from the permease domain during infection and functions 154 as an iron-dependent transcriptional repressor to autoregulate the expression of its own 155 operon (34). YtgR represents the only identified iron-dependent transcriptional regulator 156 in Chlamydia. Whether YtgR maintains a more diverse transcriptional regulon beyond 157 the ytgABCD operon has not yet been addressed and remains an intriguing question in 158 the context of immune-mediated iron limitation to *C. trachomatis*. Crucially, 159 simultaneous IFN- $\gamma$ -mediated iron and Trp starvation raise the possibility that C. 160 trachomatis could have evolved an integrated response to multiple stresses.

161 Consistent with the highly reduced capacity of the chlamydial genome, it is likely 162 that *C. trachomatis* has a limited ability to tailor a specific response to each individual stress. In the absence of identifiable homologs for most global stress response 163 164 regulators in C. trachomatis, we hypothesized that primary stress responses to 165 pleiotropic insults may involve mechanisms of regulatory integration, whereby important 166 molecular pathways are co-regulated by stress-responsive transcription factors such that they can be utilized across multiple stress conditions simultaneously induced by the 167 168 host. Here, we report on the unique iron-dependent regulation of the trpRBA operon in 169 Chlamydia trachomatis. We propose a model of iron-dependent transcriptional 170 regulation of *trpRBA* mediated by the repressor YtqR binding specifically to the IGR, 171 which may have implications for how C. trachomatis responds to immunological and 172 environmental insults. Such a mechanism of iron-dependent regulation of Trp 173 biosynthesis has not been previously described in any other prokaryote and adds to the 174 catalog of regulatory models for Trp biosynthetic operons in bacteria. Further, it reveals 175 a highly dynamic mode of regulatory integration within the *trpRBA* operon, exemplifying 176 the importance of this pathway to chlamydial stress response.

177 Results

Brief iron limitation via 2,2-bipyridyl treatment yields iron-starved, but nonpersistent *Chlamydia trachomatis.* To identify possible instances of regulatory integration between iron and Trp starvation in *C. trachomatis*, we optimized a stress response condition that preceded the development of a characteristically persistent phenotype. We reasoned that in order to effectively identify regulatory integration, we would need to investigate the bacterium under stressed, but not aberrant, growth

184 conditions such that we could distinguish primary stress responses from abnormal growth. To specifically investigate the possible contribution of iron limitation to a broader 185 186 immunological (e.g. IFN- $\gamma$ -mediated) stress, we utilized the membrane-permeable iron chelator 2.2-bipyridyl (Bpdl), which has the advantage of rapidly and homogeneously 187 188 starving *C. trachomatis* of iron (35). We chose to starve *C. trachomatis* serovar L2 of iron starting at 12 hrs post-infection (hpi), or roughly at the beginning of mid-cycle 189 190 growth. At this point the chlamydial organisms represent a uniform population of 191 replicative RBs that are fully competent, both transcriptionally and translationally, to 192 respond to stress. We treated infected HeLa cell cultures with 100 µM Bpdl or mock for 193 either 6 or 12 hours (hrs) to determine a condition sufficient to limit iron to C. 194 trachomatis without inducing hallmark persistent phenotypes. We stained infected cells 195 seeded on glass coverslips with convalescent human sera and analyzed chlamydial 196 inclusion morphology under both Bpdl- and mock-treated conditions by laser point-197 scanning confocal microscopy (Fig. 1A). Following 6 hrs of Bpdl treatment, chlamydial 198 inclusions were largely indistinguishable from mock-treated inclusions, containing a 199 homogeneous population of larger organisms, consistent with RBs in mid-cycle growth. 200 However, by 12 hrs of Bpdl treatment, the inclusions began to display signs of aberrant 201 growth: they were perceptibly smaller, more comparable in size to 18 hpi, and contained 202 noticeably fewer organisms, perhaps indicating a delay in RB-to-EB differentiation. 203 These observations were consistent with our subsequent analysis of genome replication 204 by quantitative PCR (qPCR; Fig. 1B.) At 6 hrs of Bpdl treatment, there was no 205 statistically distinguishable difference in genome copy number when compared to the 206 equivalent mock-treated time-point. However, by 12 hrs of treatment, genome copy

207 number was significantly reduced 4.7-fold in the Bpdl-treated group relative to mock-208 treatment (p = 0.0033). We then assayed the transcript expression of two markers for 209 persistence by reverse transcription quantitative PCR (RT-qPCR): the early gene euo, 210 encoding a transcriptional repressor of late-cycle genes (Fig.1C), and the adhesin 211 omcB, which is expressed late in the developmental cycle (Fig. 1D). Characteristic 212 persistence would feature sustained high euo expression late into infection, and 213 suppressed omcB expression throughout development. We observed that at 6 hrs of 214 Bpdl treatment, there was no statistically distinguishable difference in either euo or 215 omcB expression when compared to the mock-treatment. Still at 12 hrs of Bpdl 216 treatment, euo expression was unchanged. However, omcB expression was 217 significantly induced following 12 hrs of Bpdl-treatment (p = 0.00015). This was 218 unexpected, but we note that *omcB* expression has been shown to vary between 219 chlamydial serovars and species when starved for iron (31). Collectively, these data 220 indicated that 6 hrs of Bpdl treatment was a more suitable time-point at which to monitor 221 iron-limited stress responses.

We additionally assayed these same metrics following 6 or 12 hrs of Trp 222 223 starvation by culturing cells in either Trp-replete or Trp-deplete DMEM-F12 media 224 supplemented with fetal bovine serum (FBS) pre-dialyzed to remove amino acids. We 225 observed no discernable change in inclusion morphology out to 12 hrs of Trp starvation 226 (Fig. S2A), but genome copy numbers were significantly reduced 2.7-fold at this time-227 point (p = 0.00612; Fig. S2B). The transcript expression of *euo* (Fig. S2C) and *omcB* 228 (Fig. S2D) did not significantly change at either treatment duration, but Trp-depletion did 229 result in a 2.0-fold reduction in omcB expression, consistent with a more characteristic

persistent phenotype. These data therefore also indicated that 6 hrs of treatment wouldbe ideal to monitor non-persistent responses to Trp limitation.

232 We next sought to determine whether our brief 6-hr Bpdl treatment was sufficient 233 to elicit a transcriptional iron starvation phenotype. We chose to analyze the expression 234 of three previously identified iron-regulated transcripts, ytqA (Fig. 2A), ahpC (Fig. 2B) 235 and devB (Fig. 2C), by RT-qPCR under Bpdl- and mock-treated conditions (35, 36). In 236 addition, we analyzed the expression of one non-iron-regulated transcript, dnaB (Fig. 237 2D), as a negative control (37). Following 6 hrs of Bpdl treatment, we observed that the 238 transcript expression of the periplasmic iron-binding protein ytgA was significantly 239 elevated 1.75-fold relative to the equivalent mock-treated time-point (p = 0.0052). 240 However, we did not observe induction of ytgA transcript expression relative to the 12 241 hpi time-point. We distinguish here between *elevated* and *induced* transcript expression, 242 as chlamydial gene expression is highly developmentally regulated. Thus, it can be 243 more informative to monitor longitudinal expression of genes, *i.e.* their induction, as 244 opposed to elevation relative to an equivalent control time-point, which may simply 245 represent a stall in development. While we did not observe induction of ytgA, which 246 would be more consistent with an iron-starved phenotype, we reason that this is a 247 consequence of the brief treatment period, and that longer iron starvation would 248 produce a more robust induction of iron-regulated transcripts. Note that the identification 249 of ytqA as iron-regulated has only been previously observed following extended periods 250 of iron chelation (32, 35, 38). Similarly, we observed that the transcript expression of the 251 thioredoxin ahpC was significantly elevated 2.15-fold relative to the equivalent mock-252 treated time-point (p = 0.038) but was not induced relative to the 12 hpi time-point. The

253 transcript expression of *devB*, encoding a 6-phosphogluconolactonase involved in the 254 pentose phosphate pathway, was not observed to significantly respond to our brief iron 255 limitation condition, suggesting that it is not a component of the primary iron starvation 256 stress response in C. trachomatis. As expected, the transcript expression of dnaB, a 257 replicative DNA helicase, was not altered by our iron starvation condition, consistent 258 with its presumably iron-independent regulation (37). Overall, these data confirmed that 259 our 6-hr Bpdl treatment condition was suitable to produce a mild iron starvation 260 phenotype at the transcriptional level, thus facilitating our investigation of iron-261 dependent regulatory integration. 262 Transcript expression of the *trpRBA* operon is differentially regulated by iron in 263 Chlamydia trachomatis. Upon identifying an iron limitation condition that produced a 264 relevant transcriptional phenotype while avoiding the onset of persistent development. 265 we aimed to investigate whether the immediate response to iron starvation in C. 266 trachomatis would result in the consistent induction of pathways unrelated to iron 267 utilization/acquisition, but nevertheless important for surviving immunological stress. 268 The truncated Trp biosynthetic operon, *trpRBA* (Fig. 3A), has been repeatedly linked to 269 the ability of genital and LGV serovars (D-K and L1-3, respectively) of *C. trachomatis* to 270 counter IFN- $\gamma$ -mediated stress. This is due to the capacity of the chlamydial Trp 271 synthase in these serovars to catalyze the  $\beta$  synthase reaction, *i.e.* the condensation of 272 indole to the amino acid serine to form Trp (17). In the presence of exogenous indole, 273 C. trachomatis is therefore able to biosynthesize Trp such that it can prevent the development of IFN- $\gamma$ -mediated persistence. Correspondingly, the expression of *trpRBA* 274 275 is highly induced following IFN- $\gamma$  stimulation of infected cells (39, 40). These data have

historically implicated Trp starvation as the primary mechanism by which persistence
develops in *C. trachomatis* following exposure to IFN-γ. However, these studies have
routinely depended on prolonged treatment conditions that monitor the terminal effect of
persistent development, as opposed to the immediate molecular events which may
have important roles in the developmental fate of *Chlamydia*. As such, these studies
may have missed the contribution of other IFN-γ-stimulated insults such as iron
limitation.

283 To decouple Trp limitation from iron limitation and assess their relative 284 contribution to regulating a critical pathway for responding to IFN- $\gamma$ -mediated stress, we monitored the transcript expression of the *trpRBA* operon under brief Trp or iron 285 286 starvation by RT-qPCR. When starved for Trp for 6 hrs, we observed that the 287 expression of *trpR*, *trpB* and *trpA* were all significantly induced greater than 10.5-fold 288 relative to 12 hpi (p = 0.00077, 0.025 and 9.7e-5, respectively; Fig. 3B). All three ORFs 289 were also significantly elevated relative to the equivalent mock-treated time-point (p =290 0.00076, 0.025 and 9.7e-5, respectively). This result was surprising with respect to the 291 relative immediacy of operon induction in response to our Trp starvation protocol, 292 confirming the relevant Trp-starved transcriptional phenotype. To induce Trp-deprived 293 persistence in *C. trachomatis*, many laboratories rely on compounded techniques of IFN-γ pre-treatment to deplete host Trp pools in conjunction with culturing in Trp-294 295 depleted media, among other strategies. While the phenotypic end-point differs here, it 296 is nonetheless interesting to note that only 6 hrs of media replacement is sufficient to 297 markedly up-regulate trpRBA expression. This suggests that C. trachomatis has a 298 highly attuned sensitivity to even moderate changes in Trp levels.

299 We next performed the same RT-qPCR analysis on the expression of the trpRBA 300 operon in response to 6 hrs of iron limitation via Bpdl treatment (Fig. 3C). While we 301 observed that the transcript expression of all three ORFs was significantly elevated at 302 least 2.1-fold relative to the equivalent mock-treated time-point (p = 0.015, 0.00098 and 303 0.0062, respectively), we made the intriguing observation that only the expression trpB 304 and *trpA* was significantly induced relative to 12 hpi (p = 0.00383 and 0.0195, 305 respectively). The significant induction of *trpBA* expression, but not *trpR* expression, 306 suggested that *trpBA* are specifically regulated by iron availability. This result is 307 consistent with a recent survey of the iron-regulated transcriptome in C. trachomatis by 308 RNA sequencing, which also reported that iron-starved Chlamydia specifically up-309 regulate *trpBA* expression in the absence of altered *trpR* expression (37). Our results 310 expand on this finding by providing a more detailed investigation into the specific profile 311 of this differential regulation of *trpRBA* in response to iron deprivation. Taken together, 312 these findings demonstrated that an important stress response pathway, the *trpRBA* 313 operon, is regulated by the availability of both Trp and iron, consistent with the notion 314 that the pathway may be cooperatively regulated to respond to various stress 315 conditions. Notably, iron-dependent regulation of Trp biosynthesis has not been 316 previously documented in other prokaryotes. 317 Specific iron-regulated expression of *trpBA* originates from a novel alternative 318 transcriptional start site within the trpRBA intergenic region. We hypothesized that 319 the specific iron-related induction of *trpBA* expression relative to *trpR* expression may 320 be attributable to an iron-regulated alternative transcriptional start site (alt. TSS) 321 downstream of the *trpR* ORF. Indeed, a previous study reported the presence of an alt.

322 TSS in the trpRBA IGR, located 214 nucleotides upstream of the trpB translation start 323 position (20). However, a parallel study could not identify a TrpR binding site in the 324 trpRBA IGR (21). We reasoned that a similar alt. TSS may exist in the IGR that 325 controlled the iron-dependent expression of trpBA. We therefore performed Rapid 326 Amplification of 5'-cDNA Ends (5'-RACE) on RNA isolated from C. trachomatis L2-327 infected HeLa cells using the SMARTer 5'/3' RACE Kit workflow (Takara Bio). Given the 328 low expression of the trpRBA operon during normal development, we utilized two 329 sequential gene-specific amplification steps (nested 5'-RACE) to identify 5' cDNA ends 330 in the *trpRBA* operon. These nested RACE conditions resulted in amplification that was 331 specific to infected-cells (Fig. S3A). Using this approach, we analyzed four conditions: 332 12 hpi, 18 hpi, 12 hpi + 6 hrs of Bpdl treatment, and 12 hpi + 6 hrs of Trp-depletion (Fig. 333 4A). We observed three RACE products that migrated with an apparent size of 1.5, 1.1 334 and 1.0 kilobases (kb). At 12 and 18 hpi, all three RACE products exhibited low 335 abundance, even following the nested PCR amplification. This observation was 336 consistent with the expectation that the expression of the *trpRBA* operon is very low 337 under normal, iron and Trp-replete conditions. However, we note that the 6-hr difference 338 in development did appear to alter the representation of the 5' cDNA ends, which may 339 suggest a stage-specific promoter utilization within the *trpRBA* operon. In our Trp 340 starvation condition, we observed an apparent increase in the abundance of the 1.5 kb 341 RACE product, which was therefore presumed to represent the primary TSS upstream 342 of trpR, at nucleotide position 511,389 (C. trachomatis L2 434/Bu). Interestingly, the 1.0 kb product displayed a very similar apparent enrichment following Bpdl treatment, 343 344 suggesting that this RACE product represented a specifically iron-regulated TSS. Both

the 1.5 and 1.0 kb RACE products were detectable in the Trp-depleted and iron-

346 depleted conditions, respectively, during the primary RACE amplification, consistent

347 with their induction under these conditions (Fig. S3B).

348 If iron depletion was inducing *trpBA* expression independent of *trpR*, we 349 reasoned that we would observe specific enrichment of trpB sequences in our 5'-RACE 350 cDNA samples relative to trpR sequences. We again utilized RT-qPCR to quantify the 351 abundance of *trpB* transcripts relative to *trpR* transcripts in the 5'-RACE cDNA samples 352 (Fig. 4B). In agreement with our model, only under iron starved conditions did we 353 observe a significant enrichment of trpB relative to trpR (p < 0.01). Additionally, we 354 observed that at 12 and 18 hpi in iron-replete conditions, the ratio of trpB to trpR was 355 approximately 1.0, suggesting non-preferential basal expression across the three 356 putative TSSs. Another factor contributing to this ratio is the synthesis of the full-length 357 trpRBA polycistron. In support of this, the trpB to trpR ratio remained near 1.0 under the 358 Trp-starved condition, which would be expected during transcription read-through of the 359 whole operon. The apparent lack of preferential promoter utilization as described above 360 could be attributed to the relatively low basal expression of the operon at 12 and 18 hpi 361 under Trp- and iron-replete conditions, thus precluding quantitative detection of 362 differential promoter utilization in this assay.

To determine the specific location of the 5' cDNA ends within the *trpRBA* operon, we isolated the 5'-RACE products across all conditions by gel extraction and cloned the products into the pRACE vector supplied by the manufacturer. We then sequenced the ligated inserts and BLASTed the sequences against the *C. trachomatis* L2 434/Bu genome to identify the location of the 5'-most nucleotides (Fig. 4C). These data are

displayed as a statistical approximation of the genomic regions most likely to be 368 369 represented by the respective 5'-RACE products in both histogram (semi-continuous) 370 and density plot (continuous) format (See Dataset S1 for a description of all mapped 5'-371 RACE products). As expected, the 1.5 kb product mapped in a distinct and tightly 372 grouped peak near the previously annotated *trpR* TSS, with the mean and modal 373 nucleotide being 511,388 and 511,389, respectively (Fig. S4A). Surprisingly, we found 374 that neither the 1.1 or 1.0 kb RACE product mapped to the previously reported alt. TSS 375 in the *trpRBA* IGR, at position 511,826. Instead, we observed that the 1.1 kb product 376 mapped on average to nucleotide position 511,878, with the modal nucleotide being 377 found at 511,898 (Fig. S4B). The 1.0 kb product mapped with a mean nucleotide 378 position of 512,013, with the modal nucleotide being 512,005 (Fig. S4C), only 35 bases 379 upstream of the *trpB* coding sequence. Interestingly, the 1.0 kb product mapped to a region of the *trpRBA* IGR flanked by consensus  $\sigma^{66}$  -10 and -35 promoter elements, 380 381 found at positions 512,020-5 and 511,992-7, respectively (41). These data collectively 382 pointed toward the 1.0 kb 5'-RACE product representing a novel, iron-regulated alt. TSS and *bona fide*  $\sigma^{66}$ -dependent promoter element that allows for the specific iron-383 384 dependent expression of trpBA.

YtgR specifically binds to the *trpRBA* intergenic region in an operator-dependent
manner to repress transcription of *trpBA*. As the only known iron-dependent
transcriptional regulator in *Chlamydia*, we hypothesized that YtgR may regulate the irondependent expression of *trpBA* from the putative promoter element we characterized by
5'-RACE. Using bioinformatic sequence analysis, we investigated whether the *trpRBA*IGR contained a candidate YtgR operator sequence. By local sequence alignment of

391 the putative YtqR operator sequence (33) and the *trpRBA* IGR, we identified a high-392 identity alignment (76.9% identity) covering 67% of the putative operator sequence (Fig. 393 5A). Interestingly, this alignment mapped to the previously identified palindrome 394 suspected to have operator functionality (20). By global sequence alignment of the YtgR 395 operator to the palindromic sequence, an alignment identical to the local alignment was 396 observed, which still displayed relatively high sequence identity (43.5% identity). We 397 hypothesized that this sequence functioned as an YtgR operator, despite being located 398 184 bp upstream of the *trpBA* alt. TSS.

399 To investigate the ability of YtgR to bind and repress transcription from the putative *trpBA* promoter, we implemented a heterologous two-plasmid assay that 400 401 reports on YtgR repressor activity as a function of  $\beta$ -galactosidase expression (14). In 402 brief, a candidate DNA promoter element was cloned into the pCCT expression vector 403 between an arabinose-inducible pBAD promoter and the reporter gene lacZ. This 404 plasmid was co-transformed into BL21 (DE3) E. coli along with an IPTG-inducible 405 pET151 expression vector with (pET151-YtgR) or without (pET151-EV) the C-terminal 139 amino acid residues of CTL0323 (YtgC). Note that we have previously 406 407 demonstrated that this region is a functional iron-dependent repressor domain (34). To 408 verify the functionality of this assay, we determined whether ectopic YtgR expression 409 could repress pCCT reporter gene expression in the presence of three candidate DNA 410 elements: a no-insert empty vector (pCCT-EV), the putative promoter element for C. 411 trachomatis IpdA (pCCT-IpdA), and the promoter region of the ytgABCD operon (pCCT-412 ytqABCD; Fig. 5B). As expected, from the pCCT-EV reporter construct, ectopic YtqR 413 expression did not significantly reduce the activity of  $\beta$ -galactosidase. Additionally,

414 reporter gene expression from pCCT-lpdA, containing the promoter of iron-regulated 415 lpdA (37), which is not known to be YtqR-regulated, was not affected by ectopic expression of YtgR. This demonstrated that the assay can discriminate between the 416 417 promoter elements of iron-regulated genes and *bona fide* YtgR targeted promoters. 418 Indeed, in the presence of pCCT-ytqABCD, induction of YtqR expression produced a 419 significant decrease in  $\beta$ -galactosidase activity (p = 0.03868) consistent with its 420 previously reported auto-regulation of this promoter (34). 421 Using this same assay, we then inserted into the pCCT reporter plasmid 1) the 422 *trpR* promoter element (pCCT-*trpR*), 2) the putative *trpBA* promoter element 423 represented by the IGR (pCCT-*trpBA*), and 3) the same putative *trpBA* promoter 424 element with a mutated YtgR operator sequence that was diminished for both 425 palindromicity and A-T richness, two typical features of prokaryotic promoter elements 426 (pCCT-trpBAAOperator; Fig. 5C) (42, 43). When YtgR was ectopically expressed in the pCCT-*trpR* background, we observed no statistically distinguishable change in  $\beta$ -427 428 galactosidase activity. However, in the pCCT-trpBA background, ectopic YtgR expression significantly reduced  $\beta$ -galactosidase activity at levels similar to those 429 430 observed in the pCCT-ytgABCD background (p = 0.01219). This suggested that YtgR 431 was capable of binding to the *trpBA* promoter element specifically. Interestingly, this 432 repression phenotype was abrogated in the pCCT-*trpBA* Operator background, where we observed no statistically meaningful difference in  $\beta$ -galactosidase activity. We 433 subsequently addressed whether the region of the *trpRBA* IGR containing the YtgR 434 operator site was sufficient to confer YtgR repression in this assay (Fig. S5). Therefore, 435 436 we cloned three fragments of the *trpRBA* IGR into the pCCT reporter plasmid: the first

437 fragment represented the 5'-end of the IGR containing the operator site at the 3'-end 438 (pCCT-IGR1), the second fragment represented a central region of the IGR containing the operator site at the 5'-end (pCCT-IGR2), and the third fragment represented the 3'-439 440 end of the IGR and did not contain the operator site (pCCT-IGR3). Surprisingly, we 441 observed that none of these fragments alone were capable of producing a significant 442 repression phenotype in our reporter system. This finding indicated that while the operator site was necessary for YtgR repression, it alone was not sufficient. Together, 443 444 these data indicated that YtgR could bind to the *trpBA* promoter element and that this 445 binding was dependent upon an intact AT-rich palindromic sequence, likely representing 446 an YtgR operator, but that further structural elements in the *trpRBA* IGR may be 447 necessary for repression. Nonetheless, we demonstrated the existence of a functional 448 YtgR binding site that conferred iron-dependent transcriptional regulation to trpBA. 449 independent of the major *trpR* promoter.

450 Iron limitation promotes transcription read-through at the *trpRBA* YtgR operator 451 site. If YtgR binds to the operator sequence within the intergenic region under iron-452 replete conditions, does it influence RNA polymerase (RNAP) read-through from the major *trpR* promoter to synthesize the polycistronic *trpRBA* mRNA? We hypothesized 453 454 that the presence of YtgR at the *trpRBA* operator may disadvantage the processivity of 455 RNAP initiating transcription at the upstream *trpR* promoter. Similar systems of RNAP 456 read-through blockage have been reported; the transcription factor Reb1p "roadblocks" 457 RNAPII transcription read-through in yeast by a mechanism of RNAP pausing and subsequent labelling for degradation (44). To investigate this question, we first identified 458 459 transcription termination sites (TTSs) in the *trpRBA* operon in *C. trachomatis*. We

460 utilized 3'-RACE to map the 3'-ends of transcripts using gene-specific primers within the 461 trpR CDS (Fig. 6A; bottom). We again utilized two RACE amplification cycles to generate distinct, specific bands suitable for isolation and sequencing (Fig. S6B-C). By 462 463 gel electrophoresis of the 3'-RACE products, we observed the appearance of four 464 distinct bands that migrated with an apparent size of 0.55, 0.45, 0.40 and 0.20 kb. In our Trp-depleted condition, we observed only a very weak amplification of the 2.5 - 3 kb 465 full-length trpRBA message by 3'-RACE (Fig. S6A). However, we did observe it across 466 467 all replicates. To confirm that the full-length product was relatively specific to the Trpdeplete treatment, we amplified the *trpRBA* operon by RT-PCR from the 3'-RACE cDNA 468 469 (Fig. 6A; top). As expected, only in the Trp-deplete sample did we observe robust 470 amplification of the full-length trpRBA message. We note however that image contrast 471 adjustment reveals a very weak band present in all experimental samples.

472 To identify the specific TTS locations, we gel extracted the four distinct 3'-RACE 473 bands across all conditions and cloned them into the pRACE sequencing vector as was 474 done for the 5'-RACE experiments. We then sequenced the inserted RACE products 475 and mapped them to the C. trachomatis L2 434/Bu genome (Fig. 6B). This revealed a 476 highly dynamic TTS landscape contained almost exclusively within the *trpRBA* IGR, 477 which has not previously been investigated (For a full description of mapped 3'-RACE products, see Dataset S2). The 0.20 kb RACE product mapped tightly to the 3'-end of 478 479 the *trpR* CDS, with a mean nucleotide position of 511,665 and a modal nucleotide 480 position of 511,667 (Fig. S7A). Contrastingly, the other three RACE products did not map in such a way so as to produce specific, unambiguous modal peaks. Instead, their 481 482 distribution was broader and more even, with only a few nucleotide positions mapping

483 more than once. Accordingly, the 0.45 kb product mapped with an average nucleotide 484 position of 511,889, just downstream of the 1.1 kb 5'-RACE product (Fig. S7C), while 485 the 0.55 kb product mapped with an average nucleotide position of 511,986, upstream 486 of the 1.0 kb 5'-RACE product (Fig. S7D). Interestingly, the 0.40 kb product mapped to 487 a region directly overlapping the putative YtgR operator site, with a mean nucleotide 488 position of 511,811 (Fig. S7B). We therefore reasoned that this putative TTS may have 489 an iron-dependent function.

490 We next aimed to quantitatively analyze the possibility that iron-depletion, and 491 thus dissociation of YtgR from this region, may facilitate transcription read-through at 492 the operator site. Working from the 3'-RACE generated cDNAs, we utilized RT-gPCR to 493 monitor the abundance of various amplicons across the *trpRBA* operon in relation to a 494 "read-through" normalization amplicon that should only be represented when the full-495 length *trpRBA* message is transcribed (Fig. 6C). Therefore, as each amplicon is 496 increasingly represented as a portion of the full-length, read-through transcript, the 497 representation ratio of the specific amplicon to the normalization amplicon should 498 approach 1.0. We first analyzed an amplicon from nucleotide 511,416 - 531 to monitor 499 transcript species associated with transcription initiating at the *trpR* promoter. We 500 observed that the representation of this amplicon was not significantly altered following 501 iron limitation relative to 12 hpi, suggesting that the depletion of iron was not affecting 502 initiation of transcription at the *trpR* promoter. Interestingly, at 18 hpi, the representation 503 ratio of this amplicon significantly shifted further away from 1.0 (p = 0.00358), indicating that at 18 hpi this amplicon is represented less as a component of read-through 504 505 transcription relative to 12 hpi. As expected, under Trp-deplete conditions, the

representation ratio shifted significantly closer to 1.0 (p = 0.00064), consistent with readthrough transcription of the full-length *trpRBA* message.

508 We then preformed the same analysis on an amplicon from nucleotide 511.639 -509 764, immediately upstream of the TTS at the YtgR operator site. We again observed 510 that at 18 hpi, the representation ratio was significantly increased (p = 0.01046), and 511 following Trp-depletion, the ratio was significantly decreased (p = 0.00023), as 512 expected. Notably, and consistent with our hypothesis, we observed that the 513 representation ratio of this amplicon was also significantly closer to 1.0 following iron 514 limitation (p = 0.00407), suggesting that transcription read-through was increased at this 515 site under iron limited conditions. If YtgR is dissociating from the operator site during 516 iron depletion, a greater proportion of transcripts would be expected to read-through this 517 locus. 518 Finally, we analyzed an amplicon from nucleotide 513,856 – 968, at the very 3'-end of 519 trpA to monitor terminal transcription under our experimental conditions. At 18 hpi, we 520 observed a significant increase in the representation ratio of this amplicon (p =521 0.00476), which is likely attributable to both basal levels of alternative transcription from 522 the IGR as well as poor transcription read-through of the full-length message. Following 523 6 hrs of Bpdl treatment, we also observed a significant increase in the representation ratio of this amplicon (p = 0.01510), which supports the finding that trpBA is being 524 525 preferentially transcribed under this condition, distinct from the full-length trpRBA 526 transcript. We were only able to detect a marginal decrease in the representation of this amplicon under Trp-depleted conditions (p = 0.07942), which may suggest that the very 527 528 3'-end of *trpRBA* is relatively under-represented than our normalization amplicon, which

529 falls within the middle of the operon. In fact, recent work has reported on the relatively 530 poor representation of 3'-end mRNAs in Chlamydia (45). In sum, this set of experiments 531 provides evidence that iron-depletion specifically alters the representation of particular 532 mRNA species across the *trpRBA* operon. Additionally, they implicate iron-dependent YtgR DNA-binding as the mediator of these effects. By alleviating YtgR repression via 533 iron depletion, transcription is allowed to proceed through the operator site, albeit at 534 535 basal levels. Concomitantly, transcription is specifically activated at the downstream alt. 536 TSS for *trpBA*.

537 Discussion

In this study, we report a mechanism of stress adaptation that integrates 538 539 responses to iron and Trp starvation. Specifically, we demonstrated that the trpRBA 540 operon is transcriptionally regulated by the iron-dependent repressor YtgR. We 541 determined that iron-dependent expression of *trpBA* initiates from a novel internal 542 promoter in an IGR of the *trpRBA* operon and that this IGR also contains an YtgR 543 operator site necessary to confer a transcriptional repression phenotype. We suggest that the dual promoter configuration of *trpRBA* presents the opportunity for YtgR to 544 block transcription read-through from the *trpR* promoter; transcripts terminate at the 545 546 YtgR operator site and iron depletion facilitates read-through of the operon at this locus. 547 This is the first time an iron-dependent mode of regulation has been shown to control 548 the expression of tryptophan biosynthesis in prokaryotes, which is a reflection of the 549 uniquely specialized nature of *C. trachomatis*.

550 The distance separating the YtgR operator site from the *trpBA* alt. TSS indicate 551 the involvement of a regulatory mechanism more complex than simple steric hindrance 552 of RNAP by YtgR. One possible explanation is that YtgR functions similarly to other 553 prokaryotic transcription factors that repress "at a distance" by a mechanism of DNA-554 bending (46, 47). In this scenario, YtgR binding simultaneously to an additional operator 555 site may facilitate a conformational bend in the double-helix DNA such that RNAP no 556 longer has access to the alternative *trpBA* promoter site. This would be consistent with 557 the observation that a truncated *trpRBA* IGR containing the candidate YtgR operator is 558 insufficient to confer transcriptional repression. The topological alteration induced by 559 DNA-bending could also feasibly contribute to diminished RNAP read-through from the 560 trpR promoter. In silico identification of additional putative YtgR operator sites was 561 unsuccessful, which could be due to the lack of enough validated binding sites to 562 generate a robust consensus sequence. Thus, a more unbiased approach (e.g. ChIP-563 Sequencing) will be required to identify additional YtgR binding events. We also note 564 that the YtqR operator upstream of ytqA, while only 21 bp from the predicted -35 565 element, rests within a 660-bp IGR, raising the possibility that other cryptic YtgR 566 operators are present in this sequence. Another possibility is that YtgR functions in concert with additional transcription factors more proximal to the trpBA promoter 567 568 elements. In *E. coli*, the repressor Fis binds 135 bp upstream of the *nir* promoter TSS, 569 controlling co-activation of *nir* expression by proximally-bound Fnr and NarL/NarP (48). 570 While we have no evidence to suggest other transcription factors are controlling trpBA 571 expression, this does not preclude the possibility of their involvement.

572 While we demonstrate here that iron-dependent *trpBA* expression originates from 573 a novel promoter element immediately upstream of the *trpB* CDS, this is not the first 574 description of an alt. TSS within the *trpRBA* IGR. Carlson, et al. identified an alt. TSS 575 within the IGR which they suggested was responsible for *trpBA* expression (20). 576 Interestingly, this TSS was observed to originate from within the palindromic sequence 577 that we have identified here as a functional YtgR operator site. In these studies, we 578 were unable to confirm the presence of the previously identified alt. TSS by 5'-RACE. 579 This is likely because Carlson, et al. examined the presence of transcript origins 580 following 24 hrs of Trp starvation whereas here we monitored immediate responses to 581 stress following only 6 hrs of treatment. Prolonged Trp depletion would result in a more 582 homogeneously stressed population of chlamydial organisms that may exhibit the same 583 preferential utilization of the promoter identified by Carlson, et al. Population 584 heterogeneity in response to brief stress may explain the observation of multiple 585 T(S/T)Ss across the *trpRBA* operon in our studies. However, the contribution of such a 586 Trp-dependent alt. TSS to the general stress response of *C. trachomatis* remains 587 unclear. Akers & Tan were unable to verify TrpR binding to the *trpRBA* IGR by EMSA, 588 suggesting that some other Trp-dependent mechanism may control transcription from 589 this site (21). Ultimately, our approach of investigating more immediate responses to 590 stress revealed previously unreported mechanisms functioning to regulate Trp 591 biosynthesis in *C. trachomatis*, underscoring the value of transient as opposed to 592 sustained induction of stress.

593 Another mechanism of regulation reported to control the chlamydial *trpRBA* 594 operon is Trp-dependent transcription attenuation. Based on sequence analysis, a 595 leader peptide has been annotated within the *trpRBA* IGR (23). Presumably, this 596 functions analogously to the attenuator in the *E. coli trpEDCBA* operon; Trp starvation 597 causes ribosome stalling at sites of enriched Trp codons such that specific RNA 598 secondary structures form to facilitate RNAP read-thru of downstream sequences - in 599 this case, trpBA (49). However, robust experimental evidence to support the existence 600 of attenuation in *C. trachomatis* is lacking. To date, the only experimental evidence that 601 supports this model was reported by Carlson, et al., who demonstrated that transcript expression of *trpBA* is increased following 24 hr Trp-depletion in a *trpR*-mutant strain of 602 603 *C. trachomatis*, suggesting that an additional level of Trp-dependent regulation controls 604 trpBA expression (20). However, this could be attributable to an alternative Trp-605 dependent mechanism controlling *trpBA* expression at the alt. TSS identified by 606 Carlson, et al. None of the data presented here point conclusively to the existence of a 607 Trp-dependent attenuator; while we acknowledge that the additional termination sites 608 identified in our 3'-RACE assay may represent termination events mediated by an 609 attenuator, without more specific analysis utilizing mutated sequences we cannot draw 610 definitive conclusions about the functional relevance of those termination sites. 611 Additionally, it is unlikely that we would be able to observe Trp-dependent attenuation 612 under our brief stress conditions given that attenuation has a much higher tolerance for Trp-depletion than TrpR-mediated transcriptional repression in *E. coli* (50). 613 614 Interestingly, in *Bacillus subtilis*, Trp-dependent attenuation of transcription takes 615 on a form markedly different from that in *E. coli*. Whereas attenuation functions in *cis* for 616 the E. coli trp operon, B. subtilis utilize a multimeric Tryptophan-activated RNA-binding 617 Attenuation Protein, TRAP, which functions in *trans* to bind *trp* operon RNA under Trp-618 replete conditions, promoting transcription termination and inhibiting translation (51). This interaction is antagonized by anti-TRAP in the absence of charged tRNA<sup>Trp</sup>, leading 619 620 to increased expression of TRAP regulated genes. We suggest that YtgR may

621 represent the first instance of a separate and distinct clade of attenuation mechanisms: 622 iron-dependent *trans*-attenuation. We have demonstrated that transcription terminates 623 in the *trpRBA* operon at the YtqR operator site, and that read-thru of the operon is 624 facilitated under iron-deplete conditions, which is consistent with the idea that relief of 625 iron-activated YtqR DNA-binding at this site would permit RNAP to pass through the 626 YtgR operator site. This mechanism may function independently of specific RNA 627 secondary structure, relying instead on steric blockage of RNAP processivity, but 628 ultimately producing a similar result. Possible regulation of translation remains to be 629 explored. The recent development of new genetic tools to alter chromosomal 630 sequences and generate conditional knockouts in C. trachomatis should enable a more 631 detailed analysis of *trpRBA* regulation, including possible *trans*-attenuation (52, 53). 632 As a Trp auxotroph, what might be the biological significance of iron-dependent 633 YtgR regulation of the *trpRBA* operon in *C. trachomatis*? We have already noted the 634 possibility that iron-dependent trpBA regulation in C. trachomatis may enable a 635 response to simultaneous Trp and iron starvation, such as that likely mediated by IFN- $\gamma$ . 636 However, this mechanism also presents the opportunity for C. trachomatis to respond 637 similarly to distinct sequential stresses, where a particular stress primes the pathogen to 638 better cope with subsequent stresses. To reach the female upper genital tract (UGT), 639 where most significant pathology is identified following infection with C. trachomatis, the 640 pathogen must first navigate the lower genital tract (LGT). Chlamydia infections of the 641 female LGT are associated with bacterial vaginosis (BV), which is characterized by 642 obligate and facultative anaerobe colonization, some of which catalyze the production of 643 indole (54, 55). This provides C. trachomatis with the necessary substrate to salvage

644 tryptophan via TrpBA. Interestingly, the LGT is also likely an iron-limited environment. 645 Pathogen colonization and BV both increase the concentration of mucosal lactoferrin 646 (Lf), an iron-binding glycoprotein, which can starve pathogens for iron (56, 57). Lf 647 expression is additionally hormone-regulated, and thus the LGT may normally 648 experience periods of iron limitation (58, 59). Moreover, the expression of TfR is 649 constrained to the basal cells of the LGT stratified squamous epithelium (60), which 650 likely restricts iron from C. trachomatis infecting the accessible upper layers of the 651 stratified epithelia. In fact, it was recently demonstrated that C. trachomatis 652 development is attenuated in the terminally differentiated layers of an *in vitro*-generated 653 stratified squamous epithelium (61). Collectively, LGT conditions that favor C. 654 trachomatis infection may be marked by concomitant iron limitation and indole 655 accessibility. For C. trachomatis, iron limitation may therefore serve as a critical signal in 656 the LGT, inducing the expression of *trpBA* such that Trp is stockpiled from available 657 indole, allowing the pathogen to counteract oncoming IFN- $\gamma$ -mediated Trp starvation. 658 We propose the possibility that iron limitation in the LGT may be a significant predictor 659 of pathogen colonization in the UGT. 660 Finally, and of note, the expression of the ribonucleotide diphosphate-reductase 661 encoding *nrdAB* was also recently shown to be iron-regulated in *C. trachomatis* (37). 662 The regulation of *nrdAB* is known to be mediated by the presumably 663 deoxyribonucleotide-dependent transcriptional repressor NrdR, encoded distal to the 664 nrdAB locus (62). As NrdR activity is not known to be modulated by iron availability, this 665 raises the intriguing possibility that here too a unique iron-dependent mechanism of 666 regulation may integrate the chlamydial stress response to promote a unified response

667 across various stress conditions. Future studies may require more metabolomics-based 668 approaches to thoroughly dissect the integration of these stress responses, as 669 transcriptome analyses alone often miss broader, pathway-oriented metabolic 670 coordination. Ultimately, these studies point towards a need to carefully re-evaluate the 671 molecular stress response in *Chlamydia*, using more targeted approaches to answer 672 more specific questions. We anticipate that the rapid progress of the field in recent 673 years will continue to catalyze exciting and important discoveries regarding the 674 fundamental biology of Chlamydia. 675 Materials and Methods 676 Please refer to the SI Appendix for a complete and detailed description of all 677 experimental reagents and methodology. For all infections, Chlamydia trachomatis LGV 678 serological variant type II was used to infect human cervical epithelial adenocarcinoma 679 HeLa cells at a multiplicity of infection of 2 or 5, depending on experiment. Indirect 680 immunofluorescent confocal microscopy experiments were performed on a Leica TCS 681 SP8 laser scanning confocal microscope in the Integrative Physiology and Neuroscience Advanced Imaging Center at Washington State University. RT-qPCR and 682 683 qPCR experiments were performed essentially as described (35, 37). Transcript 684 abundance was normalized to genome copy number for all RT-gPCR analyses. RACE 685 experiments were conducted using the SMARTer® RACE 5'/3' Kit (Takara Bio) with 686 minor modifications as noted in SI Materials and Methods. RACE products were 687 isolated by gel extraction using the Macherey-Nagel Nucleospin PCR/gel clean-up kit (Takara Bio) and sent to Eurofins Genomics, LLC for sequencing. Sequenced RACE 688 689 products were mapped to the *C. trachomatis* L2 434/Bu genome (NCBI Accession:

690	NC_010287) by nucleotide BLAST on the NCBI server. The E. coli YtgR reporter assay
691	was performed essentially as described (34) with minor modifications as indicated in $SI$
692	Materials and Methods. Briefly, BL21 (DE3) E. coli were co-transformed with the
693	indicated pCCT and pET vectors and clonal populations were selected on double-
694	selective media. Clones were cultured in double-selective media and recombinant YtgR
695	expression was induced by the addition of IPTG prior to induction of <i>lacZ</i> expression by
696	the addition of L-arabinose. Cell lysates were collected and $\beta$ -galactosidase activity was
697	measured by the Miller Assay (63).
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- 878 **Figure Legends**
- **Fig. 1.** Brief iron limitation via 2,2-bipyridyl treatment precedes the onset of
- characteristic chlamydial persistence. (A) C. trachomatis L2-infected HeLa cells were
- fixed and stained with convalescent human sera to image inclusion morphology by
- confocal microscopy following Bpdl treatment at the indicated times post-infection.
- 883 Arrowheads indicate inclusions with visibly fewer organisms in the 12-hour Bpdl-treated

condition. Figure shows representative experiment of three biological replicates. (*B*)

885 Genomic DNA (gDNA) was harvested from infected HeLa cells at the indicated times

post-infection under iron-replete (blue) and -deplete (red) conditions. Chlamydial

genome copy number was quantified by qPCR. Chlamydial genome replication is

stalled following 12 hours of Bpdl treatment, but not 6. N=2. (C) Total RNA was

889 harvested from infected HeLa cells at the indicated times post-infection under iron-

replete (teal) and -deplete (orange) conditions. The transcript abundance of hallmark

- 891 persistence genes *euo* and (*D*) *omcB* were quantified by RT-qPCR and normalized
- against genome copy number. Only at 12 hours of Bpdl treatment is *omcB* expression
- significantly affected. N=3 for 12+6, N=2 for 12+12. Statistical significance was
- determined by One-Way ANOVA followed by post-hoc pairwise *t*-tests with Bonferroni's

895 correction for multiple comparisons. \* = p < 0.05, \*\* = p < 0.01, \*\*\* = p < 0.005.

896

897	Fig. 2. Brief iron limitation condition produces mild iron-starved transcriptional
898	phenotype. (A) Total RNA and gDNA was harvested from infected HeLa cells at the
899	indicated times post-infection under iron-replete (teal) and -deplete (orange) conditions.
900	The transcript abundance of iron-regulated ytgA, (B) ahpC, (C) devB and (D) non-iron
901	regulated dnaB were quantified by RT-qPCR and normalized against genome copy
902	number. The transcript expression of <i>ytgA</i> and <i>ahpC</i> were significantly elevated
903	following 6-hour Bpdl treatment, indicative or iron starvation to <i>C. trachomatis</i> . N=3.
904	Statistical significance was determined by One-Way ANOVA followed by post-hoc
905	pairwise <i>t</i> -tests with Bonferroni's correction for multiple comparisons. $* = p < 0.05$ , $** = p$
906	< 0.01, *** = <i>p</i> < 0.005.

907

908 Fig. 3. Expression of the *trpRBA* operon in *C. trachomatis* is non-uniformly regulated by 909 brief iron limitation. (A) Cartoon depiction of the trpRBA operon (drawn to scale) with the 910 primary transcriptional start site upstream of trpR annotated. (B) Total RNA and gDNA 911 were harvested from infected HeLa cells at the indicated times post-infection under Trp-912 replete (black) and -deplete (red) conditions. The transcript expression of trpRBA 913 operon was quantified by RT-qPCR and normalized against genome copy number. All 914 three ORFs are significantly induced relative to 12 hpi following Trp starvation. N=2. (C) 915 Total RNA and gDNA were harvested from infected HeLa cells at the indicated times 916 post-infection under iron-replete (blue) and -deplete (red) conditions. The transcript 917 expression of *trpRBA* operon was quantified by RT-qPCR and normalized against 918 genome copy number. Only *trpB* and *trpA* expression was significantly induced relative 919 to 12 hpi. N=3. Statistical significance was determined by One-Way ANOVA followed by bioRxiv preprint doi: https://doi.org/10.1101/322586; this version posted May 15, 2018. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

920 post-hoc pairwise *t*-tests with Bonferroni's correction for multiple comparisons. \* = p <921 0.05, \*\* = p < 0.01, \*\*\* = p < 0.005.

922

923 Fig. 4. Iron-dependent induction of *trpBA* expression initiates within the *trpRBA* 924 intergenic region from a novel alternative transcriptional start site. (A) Total RNA was 925 harvested from infected HeLa cells at the indicated times post-infection to examine iron-926 dependent and Trp-dependent changes in the 5'-cDNA profile of the *trpRBA* operon by 927 Rapid Amplification of 5' cDNA Ends (5'-RACE). RACE products were separated on an 928 agarose gel, revealing three distinct and specific bands with apparent sizes of 1.5, 1.1 929 and 1.0 kb. Trp depletion led to the apparent enrichment of the 1.5 kb product, while 930 Bpdl treatment produced a similarly enriched 1.0 kb RACE product. Figure shows 931 representative experiment of three biological replicates. (B) To confirm that iron-932 dependent induction of *trpBA* could originate from alternative transcription initiation, RT-933 gPCR was performed on 5'-RACE cDNA to quantify the abundance of trpB transcripts 934 relative to *trpR*. Only under iron-limited conditions were *trpB* transcripts enriched relative 935 to trpR. N=3. Statistical significance determined by One-way ANOVA followed by posthoc pairwise *t*-tests. \* = p < 0.05, \*\* = p < 0.01, \*\*\* = p < 0.005. (C) The nucleotide 936 937 position of the 5' cDNA ends generated from RACE were mapped to the C. trachomatis 938 L2 434/Bu genome by nucleotide BLAST. Figure displays histogram (discrete; bin 939 width=20) and overlaid density plot (continuous) distribution of 5' nucleotide positions 940 generated from each 5'-RACE product. The dotted line represents the weighted mean of the distribution, as indicated by the integer value above each line. The identified alt. 941 942 TSSs are depicted on the *trpRBA* operon (drawn to scale) above the plot. N=3.

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943

944 Fig. 5. Ectopically expressed YtgR domain is capable of binding the putative trpBA 945 promoter element in an operator-specific manner and repress transcription in a 946 heterologous system. (A) Identification of putative YtgR operator sequence by local and 947 global nucleotide sequence alignment using EMBOSS Water and Needle algorithms, 948 respectively, to align the previously identified YtgR operator to both the *trpRBA* IGR and 949 palindromic candidate sequence. The palindrome was then mutated in our YtgR 950 repression assay as depicted to abolish palindromicity and AT-richness. (B) Ectopic 951 expression of YtgR significantly represses  $\beta$ -galactosidase activity only from the 952 promoter of its own operon, ytgABCD, and not from an empty vector or another iron-953 regulated but presumably non-YtgR targeted promoter, *lpdA*.  $3 \le N \ge 2$ . (*C*) Expression 954 of recombinant YtqR represses  $\beta$ -galactosidase activity from the putative trpBA 955 promoter element, but not the *trpR* promoter, and this repression is dependent on the 956 unaltered operator sequence identified in Fig. 5A.  $3 \le N \ge 2$ . Statistical significance 957 determined by two-sided unpaired Student's *t*-test with Welch's correction for unequal variance. \* = p < 0.05, \*\* = p < 0.01, \*\*\* = p < 0.005. 958

959

Fig. 6. Transcription termination at *trpRBA* YtgR operator site coincides with irondependent transcription read-through. (*A*) Total RNA was harvested from *C*. *trachomatis*-infected HeLa cells to analyze 3'-cDNA landscape downstream of *trpR*promoter. The top panel displays representative RT-PCR of full-length *trpRBA* message
across experimental conditions (NTC = No Template Control). Bottom panel depicts
electrophoresed 3'-RACE products and estimated sizes. N=3. (*B*) 3'-RACE products

966	were sequenced and mapped to C. trachomatis L2 434/Bu genome by nucleotide
967	BLAST. The 0.40 kb RACE product mapped to a region overlapping the predicted YtgR
968	operator site, as indicated by the cartoon depiction of the trpRBA operon (drawn to
969	scale) above the plot. N=3. (C) Analysis of transcription read-through by RT-qPCR was
970	performed on 3'-RACE cDNAs at three distinct loci across the trpRBA operon
971	representing upstream transcription initiation (511,416-531), YtgR operator site
972	termination (511,639-764) and terminal trpBA transcription (513,856-968). Abundance
973	of each amplicon was normalized to a region (Read-through) predicted to be transcribed
974	only as a part of the full-length product based on 5' and 3'-RACE data (511,792-
975	512,080). Thus, the ratio of each amplicon to the normalization amplicon represents the
976	proportion of that amplicon encoded as part of the full-length transcript, approaching
977	one as the two more closely coincide. At the YtgR operator termination site, iron
978	limitation reduces the ratio relative to 12 hpi, suggesting that transcription read-through
979	increases at this site under this condition. The transcription initiation ratio is unaffected
980	by iron limitation, while the terminal trpBA amplicon is increased, consistent with
981	alternative transcription from the alt. TSS. Statistical significance determined by One-
982	way ANOVA followed by post-hoc pairwise <i>t</i> -tests. * = $p < 0.05$ , ** = $p < 0.01$ , ***
983	0.005.

984

Author Contributions: N.D.P. and R.A.C. wrote the manuscript; N.D.P. and R.A.C.
designed the experiments; N.D.P. performed the experiments; N.D.P. and R.A.C.
analyzed and interpreted the data.

988











