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### **RESEARCH ARTICLE**



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# High-throughput comparison of gene fitness among related bacteria

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#### Abstract

**Background:** The contribution of a gene to the fitness of a bacterium can be assayed by whether and to what degree the bacterium tolerates transposon insertions in that gene. We use this fact to compare the fitness of syntenic homologous genes among related *Salmonella* strains and thereby reveal differences not apparent at the gene sequence level.

**Results:** A transposon Tn5 derivative was used to construct mutants in *Salmonella* Typhimurium ATCC14028 (STM1) and *Salmonella* Typhi Ty2 (STY1), which were then grown in rich media. The locations of 234,152 and 53,556 integration sites, respectively, were mapped by sequencing. These data were compared to similar data available for a different Ty2 isolate (STY2) and essential genes identified in *E. coli* K-12 (ECO). Of 277 genes considered essential in ECO, all had syntenic homologs in STM1, STY1, and STY2, and all but nine genes were either devoid of transposon insertions or had very few. For three of these nine genes, part of the annotated gene lacked transposon integrations (*yejM, ftsN* and *murB*). At least one of the other six genes, *trpS*, had a potentially functionally redundant gene encoded elsewhere in *Salmonella* but not in ECO. An additional 165 genes were almost entirely devoid of transposon integrations in all three *Salmonella* strains examined, including many genes associated with protein and DNA synthesis. Four of these genes (*STM14\_1498, STM14\_2872, STM14\_3360*, and *STM14\_5442*) are not found in *E. coli*. Notable differences in the extent of gene selection were also observed among the three different *Salmonella* isolates. Mutations in *hns*, for example, were selected against in STM1 but not in the two STY strains, which have a defect in *rpoS* rendering *hns* nonessential.

**Conclusions:** Comparisons among transposon integration profiles from different members of a species and among related species, all grown in similar conditions, identify differences in gene contributions to fitness among syntenic homologs. Further differences in fitness profiles among shared genes can be expected in other selective environments, with potential relevance for comparative systems biology.

#### Background

When a library of transposon (Tn) integrations is created in a bacterial genome, some insertions are not recovered in the resulting pool of mutants, either because the insertion is in an essential gene or because the gene is required in the media used to grow the bacterium. This fact has been exploited extensively to identify genes under selection when growth conditions are changed [1-3].

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Another potential utility of such data, that is explored here, is to compare different strains, serovars, and species to reveal apparent orthologs that have very different levels of fitness in different strains. We perform the first experiments to quantitate this phenomenon in *Salmonella*.

We used high-throughput sequencing to determine the location of tens of thousands of integration sites of a Tn5 derivative in the genome of *Salmonella enterica* serovar Typhimurium strain ATCC 14028 (STM1) and in *Salmonella enterica* serovar Typhi Ty2 (STY1) after growth in rich media (Luria Broth). Our datasets were compared to each other and to a series of other published data on the fitness of mutations in *Salmonella* [4-6] and *Escherichia coli* (ECO) [7-10], including a



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previously obtained transposon profile in a separate Ty2 isolate, STY2, which differs from STY1 by having mutations in *htrA*, *aroC* and *aroD* [3].

Differences in the selective pressure on apparent orthologs in the related genomes (STM1, STY1, STY2 and ECO) are of interest because they likely reflect differences in the systems that interact with these otherwise functionally similar genes or their products.

#### **Results and discussion**

# Profiling of a library of transposon insertions in *Salmonella* Typhimurium and Typhi

Five independent transposon libraries were constructed in *S.* Typhimurium ATCC 14028 (STM1) and two in *S.* Typhi Ty2 (STY1), using the EZ-Tn5 < KAN-2 > Promoter Insertion Kit (Epicentre Biotechnologies) (see methods), and grown in Luria broth (Additional file 1: Table S1). The genomic DNA directly adjacent to each transposon was obtained using a procedure similar to that described in Santiviago *et al.* [4] and sequenced as described in Additional file 2 and in Additional file 3: Figure S1.

We obtained a total of 16,642,379 first-strand Illumina sequencing reads of 100 bases in length. Sequences were subsequently filtered to reveal those reads that contained a complete primer including each unique barcode, followed by two bases of transposon (Tn) beyond the primer. The remainder of the sequence was mapped to the genome to determine the transposon integration site.

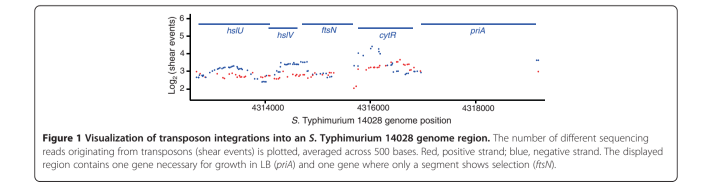
Each Tn integration site was generally represented by multiple reads, and these reads usually varied in the length of the *Salmonella* sequence due to the random DNA shearing used in the sequencing protocol. To reduce bias due to preferential PCR of some fragments, duplicate identical shear events were removed. The remaining reads for each transposon integration site were used to determine the number of different "shear events" for that transposon. This filtering resulted in the mapping of 234,152 and 53,556 Tn integration locations, with 2,827,876 and 313,585 unique shear events in STM1 and STY1, respectively. The average density of

integrations into the genome was one every 20 bases with an average of about 12 shear events per site in STM1, and one every 90 bases with an average of about 6 shear events per site in STY1.

# A genome-wide survey of permitted transposon integrations

The transposon libraries used in these experiments were constructed and grown in rich media. Those regions of the genome with rare or absent transposon integrations include regions that are essential or under strong selection in rich media. A sizeable subset of these regions should also be essential under all growth conditions. An example of a profile of transposon integrations is shown in Figure 1, which displays a region in the STM1 genome that includes a known essential gene, *priA* (primosome assembly), and a gene with an essential region, *ftsN* (involved in cell division). Regions that were essential in Luria broth are identifiable in this plot as having no transposon integrations. Near-essential regions have a lower than average number of integration sites, usually accompanied by a lower number of shear events.

Our data also provide information on the orientation of each transposon, which can be informative. For example, in Figure 1, at the beginning of cytR, adjacent to *ftsN*, the negative strand contains far more transposons; this is the strand in which the heavily expressed antibiotic resistance marker of the transposon is oriented away from *ftsN*, likely making these integrations less disruptive. Many other examples of general selection and strand-specific selection are seen in Additional file 4: Figure S2, which shows a plot of the frequency of transposon insertions across the entire STM1 genome. A dramatic example of strand-specific selection is seen in the ribosomal RNA operons, for example at position 290,000 in the genome (Additional file 4: Figure S2). In this operon transposons are only permitted in the antisense strand, perhaps because truncated sense strand transcripts produced by the strong antibiotic resistance promoter in the transposon would disrupt ribosome assembly.



Species/serovar	Strain	Essential genes/non- essential	Method (type of mutagenesis, medium)	Reference				
E. coli	K-12 MG1655	302/4477	Published literature and MD (medium-scale) and LD (large-scale) deletion mutants (targeted mutagenesis, antibiotic medium 3)	Profiling of <i>E. coli</i> chromosome (PEC) database (http://shigen.lab.nig.ac.jp/ECOli/pec/) [9,10]				
E. coli	K-12 BW25113	303/3985	Single-gene deletion mutants (targeted mutagenesis, LB)	Keio collection [7]				
E. coli	K-12 BW25113	299/3864	Single-gene deletion mutants (targeted mutagenesis, LB)	Update on the Keio collection [8]				
E. coli	K-12 W3110	299/4109	Published literature	PEC database (http://shigen.lab.nig.ac.jp/ECOli/pec/) [9,10]				
S. Typhimurium	ATCC 14028	NA/1,023	Single-gene deletion mutants (targeted mutagenesis, LB)	[4]				
S. Typhimurium	ATCC 14028	257/NA	Insertion-duplication mutagenesis (IDM) sequencing (random mutagenesis, LB)	[5]				
S. Typhimurium	LT2	144 (LB and/or M9/glc)/NA	Metabolic reconstruction ( <i>in silico</i> approach, M9/glc and LB)	[6]				
S. Typhi	Ty2 (STY2)	356/4162	Random transposon mutagenesis and two types of $\operatorname{growth}^{\operatorname{a}}$	[3]				

Table 1 Numbers of essential genes under laboratory conditions in relevant E. coli, S. Typhimurium and S. Typhi isolates

<sup>a</sup> Plating on an "aro mix" agar containing L-phe, L-trp, *p*-aminobenzoic acid and 2,3-dihydroxybenzoic acid (condition 1), six passages of growth in Luria broth (condition 2).

The transposon frequency analyses for all genes in STM1 (compared with known essentiality information for STY2 and ECO) and STY1 are presented in Additional file 5: Table S2 and Additional file 6: Table S3, respectively. Data for the *htrA<sup>-</sup> aroC<sup>-</sup>* and *aroD<sup>-</sup>* mutant STY2 were derived from two selections: a single passage on a solid medium (i) and six passages in Luria broth (ii).

#### Essential genes in E. coli

Between *E. coli, S.* Typhimurium and *S.* Typhi, over 60% of protein coding genes are syntenic and have over 95% amino acid sequence identity [11]. There are 339 of the approximately 4,000 genes in ECO that have been reported to be essential in at least one of two comprehensive studies, the PEC (Profiling of E. coli Chromosome) database and the Keio collection [7-10] (sources are summarized in Table 1). Of these potentially essential genes, 277 genes are considered essential in both databases, and all 277 genes have syntenic homologs in STM1. These genes are listed in Additional file 5: Table S2.

We ranked all *Salmonella* genes based on their density of transposons and the total number of shear events, and set a threshold of the  $15^{\text{th}}$  percentile for "highly selected" genes. Exactly 549 genes in STM1, 582 genes in STY1, and 437 genes in STY2 met these criteria. Only six of the 277 genes essential in ECO were not among these highly selected genes in STM1 (Table 2): three narrowly missed the threshold (*folK*, *yejM* and *trpS*) and three had average amounts of transposon insertions (*murB*, *ftsN* and *degS*) indicating that mutants were not under selection in LB in this isolate. Two other genes did not meet the "selected" threshold in our STY1 assay (*yrfF*, *gpsA*) and one gene, *folA*, was not found to be selected in the published STY2 data [3].

Visual inspection of Additional file 4: Figure S2 revealed that in three of the genes that are essential in ECO but seemingly not essential in STM1, part of the respective gene was, in fact, devoid of any transposon insertions: *yejM*, a putative hydrolase; *ftsN*, which encodes a cell division protein; and *murB*, a UDP-*N*-acetylenolpyruvoylglucosamine reductase. Figure 1 shows the *ftsN* gene as an example. These genes can apparently be disrupted in certain locations without losing their essential function. Thus, because of the high density of transposon integration data, we were able to reveal those cases where only part of the gene is essential.

The fourth of the six cases, *trpS*, encodes a tryptophanyl-tRNA synthetase. It may tolerate transposon insertions in STM1 and STY because of the presence of a distant paralog (*trpS2*), which does not exist in ECO. TrpS2 may substitute the TrpS function, although it is only 28% identical [12].

The fifth gene essential in ECO and not under strong selection in STM1 is *degS*, a serine endoprotease. This gene was under strong selection in STY2 [3]. That strain (but not our STY1) is an *htrA* mutant, a paralog of *degS* [13]. In our STY1 data, transposon insertion into *degS* was somewhat diminished, but not enough to qualify the gene for the "selected" category, suggesting at least some effect of the lack of a functional HtrA in STY2. However,

S. Typhimurium 14028 gene symbol	Gene name	Gene description	Best hit in S. Typhimurium LT2	Best hit in S. Typhi Ty2 <sup>a</sup>	Best hit in ECO <sup>a</sup>	STM1 transposons	STM1 reads	STY1 transposons	STY1 reads	STY2 transposons <sup>b</sup>	STY2 reads <sup>b</sup>	STY2 transposons <sup>c</sup>	STY2 reads
						Percentile ra	nk						
STM14_0106	folA	Dihydrofolate reductase	STM0087	t0090	<i>b0048</i>	4	3	8	8	19	19	46	60
STM14_0217	folK	2-amino-4-hydroxy- 6-hydroxymethyl- dihyropteridine pyrophosphokinase	STM0183	t0191	b0142	12	22	18	16	25	56	16	21
STM14_2754	уејМ	Putative hydrolase	STM2228	t0626	b2188	14	16	30	25	13	19	17	20
STM14_4041 <sup>d</sup>	degS	Serine endoprotease	STM3349	t3265	b3235	50	34	14	14	6	7	4	4
STM14_4193	trpS	Tryptophanyl-tRNA synthetase	STM3481	t4024	b3384	18	16	14	14	78	64	12	12
STM14_4208	yrfF	Intracellular growth attenuator protein	STM3495	t4011	b3398	1	4	19	19	70	42	12	27
STM14_4460	gpsA	NAD(P)H-dependent glycerol-3-phosphate dehydrogenase	STM3700	t3819	b3608	10	10	14	17	45	46	8	11
STM14_4921	ftsN	E cell division protein	STM4093	t3525	b3933	42	38	14	15	20	27	15	15
STM14_4971.J	murB	UDP-N- acetylenolpyruvoyl- glucosamine reductase	STM4136	t3489	b3972	42	48	26	22	8	24	13	21

#### Table 2 Essential genes in *E. coli* that are not as strongly selected in Typhimurium or Typhi\*

\* Details for all genes can be found in Additional file 5: Table S2. Genes were ranked by the number of transposon insertions and number of independent reads per gene. Ranks above the 15<sup>th</sup> percentile are shown in bold.

<sup>a</sup> Best hits are listed only if gene is syntenic with S. Typhimurium strain 14028 and has at least 95% sequence identity.

<sup>b</sup> Growth on L-agar with "aro mix" (L-phe, L-trp, *p*-aminobenzoic acid and 2,3-dihydroxybenzoic acid) [3].

<sup>c</sup> Growth after six passages in LB broth [3].

<sup>d</sup> Considered essential in STY2 [3].

in STM1 there is no evidence for any degree of transposon underrepresentation in *degS*, and it seems likely that another paralog, as yet undefined, can perform the proteolytic activity of DegS in this strain, if needed.

Finally, *folK* was somewhat underrepresented in transposon insertion frequency in STM1, but not to a degree that warranted inclusion in the "selected" category.

There are three other genes that are essential in ECO and STM1, but seemed not strongly selected in our STY1 survey and/or STY2. The folA gene, involved in folate biosynthesis, was strongly selected in STM1 and STY1 but did not show as much selection in STY2, especially after six passages in LB. Lastly, *yrfF* and *gpsA* are two genes essential in ECO that did not meet the "selected" threshold in our STY1 assay. Both these genes were under strong selection in STM1 and in STY2 after six passages in LB, indicating that mutations cannot be maintained for many passages in this medium. The yrfF (*igaA*) gene encodes an intracellular growth attenuator protein; and gpsA encodes a NAD(P)H-dependent glycerol-3-phosphate dehydrogenase. IgaA has been described as essential in S. Typhimurium unless there are additional mutations in the RcsCDB system, because it acts as a repressor of this system [14,15]. Expression of igaA is positively regulated by Lon and negatively modulated by Hnr (MviA) through the transcriptional regulator RpoS [16]. S. Typhi Ty2 carries a defect in the rpoS gene [17,18], which may explain the lesser degree of selection of *igaA* in STY.

# Genes that are not essential in *E. coli* but are under strong selection in Typhimurium and Typhi

The list of essential genes in ECO is stringent, generally including the inability to obtain a viable deletion mutant in rich medium. In contrast, the measure available from transposon integrations in STM1 and STY reveals genes that were under strong selection, but not necessarily essential. A group of 159 such genes that are under selection in all Salmonella (i.e. STM1, STY1 and STY2) but not essential in ECO were identified (Additional file 5: Table S2). This list included many of the genes that might be expected to be under selection, such as genes encoding parts of the ribosome and its accessory proteins, as well as some genes encoding replication components. However, there were at least 14 genes that still have an unknown or a poorly understood function (ybaB, ybeD, ybeY, phoL, ycaR, ycdC, yciM, yciS, ygfZ, yhaL, yheM, yheN, wecF, and yigP). Given the conservation of these genes between Salmonella and E. coli, these are particularly interesting targets for future studies to determine their exact function.

The approximately 900 genes shared by STM1 and STY that have no synteny in ECO (*Salmonella*-specific genes) yielded only two strongly and consistently selected genes:

*STM14\_5442* and *STM14\_2872*, which both encode putative cytoplasmic proteins. However, Santiviago *et al.* reported successful knockout mutations in both of these genes in this same strain, so they are likely not essential [4]. However, the genes may have an effect on growth, because mutants in these genes grow poorly in competitive assays (unpublished data).

# Genes under greater selection in Typhimurium than in Typhi

Genes that were under stronger selection in STM1 than in STY when the transposon libraries were grown in LB are depicted in Table 3A. The most dramatic difference was in hns, with an almost equal effect on the near adjacent gene hnr. Whereas hns is essential in Salmonella unless certain second site mutations are also present [19], no selection against insertion in this gene was found in STY in this study. The most likely explanation for this phenomenon is that S. Typhi Ty2 is known to contain a mutation in the rpoS gene [17,18]. Mutations in this gene permit second site mutations in hns to be viable [19], and hnr (mviA) is a response regulator which post-transcriptionally modulates RpoS levels [20]. Interestingly, stpA, which encodes a 53% identical paralog of hns, also showed a greater tolerance for transposons in STY than in STM1. Unlike hns, viable hnr mutants can be obtained in S. Typhimurium [21], even though this gene appears to be strongly selected in STM1. Hnr participates in RpoS stability by acting as an adaptor for degradation by the ClpXP protease [22]. Mutants in hnr show reduced growth rate because of an increased RpoS stability, which increases transcription of genes involved in growth arrest and resistance to a variety of stresses [20]. The selection against hnr mutations seen in STM1 might be due to the non-advantageous phenotype of slow cell division in these mutants when they are in competitive growth.

At least 26 additional genes appeared to be under strong selection in STM1 but not in either STY. This class includes, among others, the putrescine ABC transporter *potF*; an excisionase; a cation transport regulator *chaB*; the L,D-carboxypeptidase A *ldcA*; the phosphoribosylaminoimidazole-succinocarboxamide synthase *purC*; and *cpxP*, a periplasmic repressor of the envelope stress response pathway. Viable mutants were obtained in this same Typhimurium strain for six of these genes [4] (listed in Table 3). It is not yet known if the remaining 20 genes can be deleted but it is likely that most, if not all, are not essential.

# Genes under greater selection in Typhi than in Typhimurium

Genes that were under greater selection in STY1 and STY2 compared to STM1 are listed in Table 3B. Among the 10 genes under consistent selection in STY2 (both

S. Typhimurium 14028 gene symbol		Gene description	Best hit in S. Typhimurium strain LT2		Best hit in ECO <sup>a</sup>	STM1 transposons	STM1 reads	STY1 transposons	STY1 reads	STY2 transposons <sup>b</sup>	STY2 reads <sup>b</sup>	STY2 transposons <sup>c</sup>	STY2 reads <sup>c</sup>
						Percentile ra	nk						
A. Examples of gen	es with stron	nger selection in STM1 than in S	STY1 and STY2										
STM14_0199.RJ	yacC		STM0167	t0172	b0122	12	13	57	70	53	51	59	51
STM14_1028.R	potF	Putrescine ABC transporter putrescine-binding protein	STM0877	t2019	b0854	14	15	47	63	35	24	38	32
STM14_1092		Hypothetical protein		t1967		13	10	61	59	35	42	75	66
STM14_1127	ycbL	Putative metallo-beta- lactamase	STM0997	t1937	b0927	13	13	30	40	27	34	23	36
STM14_1141	xisW	Excisionase	STM1006	t1928		11	9	72	62	53	30	43	24
STM14_1152	dnaC	Putative replication protein	STM1015	t1917		16	16	27	36	34	31	41	32
STM14_1548	уеаМ	Putative regulatory protein	STM1279	t1159	b1790	14	17	50	59	36	25	44	38
STM14_1599	celG	Hypothetical protein	STM1317	t1196	b1733	15	17	67	75	54	46	50	48
STM14_1682	orf70	Hypothetical protein	STM1388	t1256	b1675	12	11	32	71	34	27	37	29
STM14_1877 <sup>d</sup>		Putative coiled-coil protein	STM1554	t1468		15	16	39	37	79	81	78	81
STM14_1977 <sup>d</sup>		Putative periplasmic binding protein	STM1633	t1536		13	16	34	27	86	85	86	83
STM14_1978 <sup>d</sup>		Putative ABC transporter permease component	STM1634	t1537		6	7	17	44	87	91	85	90
STM14_2116	hns	Global DNA-binding transcriptional dual regulator H-N	<i>STM1751</i> S	t1662	b1237	5	3	32	29	91	89	35	91
STM14_2119	hnr	Response regulator of RpoS	STM1753	t1664	b1235	1	1	63	58	69	59	48	31
STM14_2120	ychK	Hypothetical protein	STM1754	t1665	b1234	11	11	64	81	64	61	62	46
STM14_2140	chaB	Cation transport regulator	STM1770	t1679	b1217	12	11	30	29	63	60	73	49
STM14_2176	ldcA	L,D-carboxypeptidase A	STM1800	t1077	b1192	14	15	43	37	28	28	39	39
STM14_2186 <sup>d</sup>	gsnB	Putative cytoplasmic protein	STM1809	t1068		18	14	39	30	43	58	59	70
STM14_2278		Hypothetical protein	STM1873	t1004	b1839	14	13	37	57	48	28	54	38
STM14_2685		Putative 1,2-dioxygenase	STM2178	t0677		15	18	32	33	31	29	24	32
STM14_2708.RJ		Putative DNA-binding protein	STM2195	t0660		17	15	51	44	65	36	62	49
STM14_2745		Bicyclomycin/multidrug efflux system		t0634		16	14	65	47	66	43	73	68
STM14_3050	purC	Phosphoribosylaminoimidazole- succinocarboxamide synthase	STM2487	t0372	b2476	11	11	28	32	47	34	34	30
STM14_4596 <sup>d</sup>		Pseudogene	STM3806	t0121		14	12	78	49	53	55	36	51
STM14_4883	срхР	Periplasmic repressor	STM4060	t3559	b3913	16	16	27	46	38	41	43	34

#### Table 3 Genes displaying prominent differences in selection between Typhimurium and Typhi\*

STM14_5142 <sup>d</sup>		Putative cytoplasmic protein	STM4276	t4182		15	13	46	28	16	49	19	47
B. Examples of g	genes with st	ronger selection in STY1 and STY2	than in STM1										
STM14_0109	ksgA	Dimethyladenosine transferase	STM0090	t0093	b0051	37	28	5	5	19	13	5	5
STM14_2281	holE	DNA polymerase III subunit theta	STM1876	t1001	b1842	63	70	11	11	9	8	25	15
STM14_2498		Hypothetical protein	STM2011.1n	t0860		51	43	13	12	8	10	14	17
STM14_2665 <sup>e</sup>		Hypothetical protein	STM2161	t0694	b2128	37	53	14	13	8	7	14	11
STM14_3017	eutA	Reactivating factor for ethanolamine ammonia lyase	STM2459	t0399	b2451	26	29	16	16	10	11	16	16
STM14_3267 <sup>e</sup>		Hypothetical protein	STM2666	t2621	b2598	81	84	17	14	9	7	19	13
STM14_4041 <sup>e</sup>	degS	Serine endoprotease	STM3349	t3265	b3235	50	34	14	14	6	7	4	4
STM14_4465	yibP	Hypothetical protein	STM3705	t3814	b3613	63	65	19	18	11	11	10	11
STM14_4747	yifL	Putative outer membrane lipoprotein	STM3946	t3351	b4558	77	65	12	12	22	17	18	16
STM14_5345	treC	Trehalose-6-phosphate hydrolase	STM4453	t4488	b4239	50	49	13	13	10	8	5	9

#### Table 3 Genes displaying prominent differences in selection between Typhimurium and Typhi\* (Continued)

\* Details for all genes can be found in Additional file 5: Table 52. See Table 2 for notes a, b, and c. Ranks above the 20<sup>th</sup> percentile are shown in bold.

<sup>d</sup> Knockout mutant has been created in STM [4].

<sup>e</sup> Considered essential in STY2 [3].

after passage in LB and after growth on aro-mix agar [3]) and our own STY1 data, but not in STM1, were four genes that encode hypothetical proteins and the previously mentioned *degS*. Other genes in this class include *eutA*, involved in the ethanolamine utilization pathway, the dimethyladenosine transferase *ksgA* and *treC*, a trehalose-6-phosphate hydrolase. The gene *holE* is an interesting example that is more strongly selected in STY than in STM1. This gene encodes the theta subunit of DNA polymerase III. The STM14\_5586 protein encoded on the virulence plasmid in STM1, which is not present in STY, is a paralog that may partially substitute for *holE* [23].

#### Pseudogenes

Integrations in genes that are thought to be pseudogenes in STY and intact in STM1, or vice versa, were inspected. Of approximately 60 putative pseudogenes annotated in the *S*. Typhimurium 14028 genome, four showed strong selection in STM1 and are annotated as intact in *S*. Typhi Ty2: *STM14\_1358*, *STM14\_1498.L*, *STM14\_1778*, and *STM14\_4596*. Only one of them, *STM14\_1358*, has an ortholog in ECO (*yceQ*) and, interestingly, has been reported as essential in this species. In *STM14\_1358* and *STM14\_1498.L*, levels of selection in STM1 were similar to the levels in both STY. In the other two cases, there was no selection in at least one of the two STY isolates.

Of approximately 200 pseudogenes in *S.* Typhi Ty2, four were strongly selected in at least one of the STY: *eda*, a keto-hydroxyglutarate-aldolase/keto-deoxy-phosphogluco-nate aldolase; *astA*, an arginine succinyltransferase; *t2152* (*STM14\_0843*), a putative glycosyltransferase involved in cell wall biogenesis; and *t3548* (*STM14\_4894*), a putative cytoplasmic protein. The *eda* gene is the only case showing a strong selection in both STY1 and STY2.

In *S.* Typhi Ty2, an RNA-seq analysis of the transcriptome was recently published [24] and concluded that the vast majority of pseudogenes had low or undetected transcription. Only nine pseudogenes showed high levels of transcription, none of which correspond to our four strongly selected pseudogenes. A region annotated as a pseudogene and showing strong selection in any environmental condition suggests that a function is encoded in this region, whether it is a partial protein or a regulatory region.

# Differences between transposon mutant libraries of two strains of Typhi

In our study, 53,556 transposon integration sites for STY1 were determined. In a previous work, 370,000 insertion sites were identified in STY2. Over 100 genes showed a difference in fitness between these two strains. Some of these differences may be attributable to the growth conditions used in the two studies. Our STY1 data are from a single LB growth passage whereas the

STY2 data were from L-agar supplemented with aromatic compounds as well as from six passages in LB. Furthermore, STY2 is an attenuated strain, CVD908htrA [3], which differs from STY1 in that it carries additional mutations in aroC, aroD, and htrA [25]. Deletions in the aro genes, which encode enzymes involved in the shikimate biosynthesis pathway, render bacteria auxotrophic for the aromatic amino acids *p*-aminobenzoate (pABA) and 2,3-dihydroxybenzoate [26]. These aro mutations also result in the inability to produce ubiquinone and menaquinone, leading to respiration defects [27], and in defects in some components of the cell envelope [28], unless aromatic precursors are added to the medium. HtrA is a serine protease involved in the degradation of aberrant periplasmic proteins. An htrA mutant presents more susceptibility to oxidative stress than the wild type [29].

We found 17 transposons in *htrA* and one each in *aroC* and *aroD* in STY1. The number of transposons in *htrA* was close to the average random transposon insertion frequency (15.9) whereas *aroC* and *aroD* showed selection in both STY1 and STY2. The apparent selection of *aroC* and *aroD* in STY2 is explained by the fact that these genes were knocked out in this strain, resulting in a much smaller gene remnant as transposon target area. The *htrA* mutation may explain at least some differences between the two strains of STY, such as the selection of DsbA after six passages in LB in STY2, but not in our STY1 study. In *E. coli*, the *skp* (*hlpA* in *Salmonella*) *degP* (*htrA* in STM) double mutant is lethal [30] and the *dsbA degP* double mutant shows reduced growth [31].

STY1 showed selection in some *Salmonella* Pathogenicity Island 2 (SPI-2) genes [32,33]. Some of these were also selected in STM1 (*ssaI, ssaH, ssaR, ssaT, sifB*) but not in STY2. Some other SPI genes were interesting from a regulatory point of view. The *hilC* and *rtsA* SPI-1 regulators were selected in STY1, but not in STY2 [34]. RtsA is encoded in an operon that also includes the similarly selected *rtsB*, whose product represses the master regulator of the flagellar regulon, *flhDC* [35].

#### **Flagellar genes**

Flagellar genes show different patterns of selection comparing growth in LB broth under aeration (STM1, STY1 and STY2) versus on LB agar (STY2). Flagella and motility are highly regulated in *S*. Typhimurium and involve over 60 genes integrated in a hierarchy of controlled transcription [36]. The flagellar structure consists of three components: the basal body, the hook and the filament. The basal body consists of three rings and a rod which transverses the periplasmic space. These three components are expressed coordinately, first the hookbasal body (HBB) and later the filament, and assembled via a flagellar type III secretion apparatus [37]. The flgM gene encoding the anti-sigma 28 negative regulator of the synthesis of the flagellar filament, was selected in broth media, but not on LB agar. FlgM binds to the sigma 28 factor to prevent transcription of late flagellar genes before the completion of HBB structure [38]. In contrast, flgC, flgJ, fliI, fliK, and fliO were selected only when bacteria were grown on an agar surface. FlgC is one of the structural rod components and FlgJ is the capping protein of the rod which also possesses muramidase activity [39]. FliI is an ATPase that forms part of the flagellar type III export apparatus, although it is not essential [40]. FliK regulates the length of the hook by switching the secretion specificity from rod-hook type substrates to filament-type substrates [41]. FliO is one of the integral membrane proteins of the flagellar secretion system which seems to have a role in stabilizing another protein of this system, FliP [42]. These five proteins are components of the basal body or the flagellar export apparatus, or interact with these structures. Furthermore, *fliT*, which encodes a protein that acts as a repressor of flagella biosynthesis [43], was selected after six passages in LB in STY2 [3]. The flagellar gene *flhE* showed selection on agar growth and after six passages in LB in STY2 [3]. Although it is known that the lack of FlhE does not affect flagella biogenesis or swimming motility, these mutants are defective in swarming motility [44]. In agreement with these results, Wang et al. reported that flagellar genes were regulated in a surface-specific manner [45]. Overall, our data indicates selection for genes encoding inhibitors of flagellar biogenesis when bacteria are grown in LB broth, such as *flgM* and *fliT*; and selection for genes necessary for flagellar biosynthesis when bacteria are grown on agar.

# Comparison of fitness data with previous efforts to identify essential genes in *Salmonella*

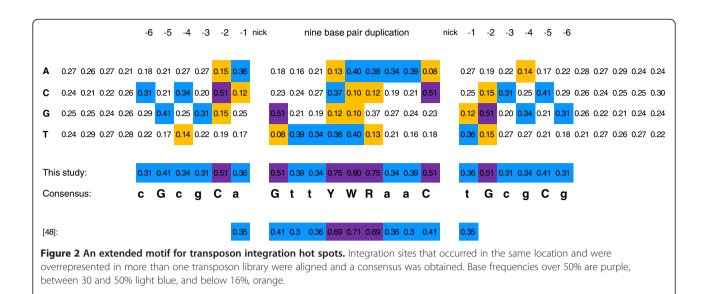
We compared our data (Additional file 5: Table S2) to previous experiments that had sought to identify essential and non-essential genes in Typhimurium (Table 1). We previously reported 1,023 genes that give viable mutants in rich media in the same strain of Typhimurium as used here [4]. At least 38 of these genes are "selected" in our study (among the 15% with the lowest density of transposon integrations and shear events). The differences may be attributable to measuring essentiality, which is absolute, versus fitness selection, which is relative. In another study, 257 genes were identified as potentially essential genes for in vitro growth in LB using an insertion-duplication mutagenesis (IDM) strategy based on a temperature-sensitive integration plasmid [5]. Of these 257 genes, only 104 were under strong selection in STM1 in our data. It was previously suggested that some of these putative essential genes might be non-essential [46]. It appears that IDM may give a high rate of false positives for essential genes. In another study, a prediction using metabolic reconstruction identified 144 genes that could be essential for growth of *S*. Typhimurium LT2 in LB, of which 71 were already known to be essential in *E. coli* [6]. Of the remaining 74 potentially novel essential phenotypes, 19 were under strong selection in STM1. Additionally, 57 genes were predicted to be essential only in minimal media. Six of these genes were under strong selection in rich media in STM1. Given the considerable discrepancy in the observed and predicted genes under selection, fitness data have the great potential to improve systems biology models in the future.

#### An extended motif for transposon integration

Tn5 transposition can occur into almost any sequence. Indeed, we were unable to find any bias in integration targeting when we used all the transposon integration data we had available and a variety of motif-finding programs (data not shown), indicating that the vast bulk of integration sites were largely randomly distributed. However, hotspots, where Tn5 integration is preferred, have been reported [47]. Goryshin et al. sequenced 198 integration sites in a plasmid and identified a short degenerate consensus palindromic motif where integrations were more frequent: a-GntYWRanC-t [48]. The dash indicates the site of cleavage of the target that is then duplicated on either side of the transposon integration. To examine whether hotspots for integration in our data might further refine this motif we filtered over 300,000 integration sites for those sites that occurred in more than one of our independent transposon libraries. Then we filtered for integration sites represented by more than the average number of total shear events. This strategy yielded 654 sites that were among the most highly preferred targets for integration. By aligning the sequences surrounding these sites, we were able to refine the originally reported motif and further extend the motif by five bases on either side, including a highly conserved C and G located two and five bases upstream of the nick in the target DNA: cGcgCa-GttYWRaaC-tGcgCg (Figure 2). The opportunity for preferred interactions along the length of this 21-base target motif might stabilize transposase-DNA pre-cleavage or cleavage complex for co-crystalization or other imaging studies.

#### Conclusions

We have identified differences in the ability to tolerate transposon integrations between *Salmonella* Typhimurium strain 14028 and two strains of *S*. Typhi Ty2. We also found potential differences in essentiality of



homologous genes between *Salmonella* and *E. coli* (summarized in Tables 2 and 3).

Sometimes, these differences in selection can be explained by the presence of an identifiable paralog, present in one genome but not another, which can take over some or all of the functions of the mutated gene. Examples include trpS and holE. In other cases, a difference in a function encoded elsewhere in the genome differentially impacts the role of orthologs. The best example of this phenomenon in the present study is hns which is essential in STM1 but not in either STY dataset. It is known that rpoS mutants permit mutations in hns in Salmonella in some circumstances [19], and STY1 and STY2 are rpoS mutants [17,18]. Similarly, hnr also showed a profound selection only in STM1. Although this gene is not essential in Salmonella, the presence of a functional copy of *rpoS* may also be the reason for this selection, because Hnr acts as a stability moderator for RpoS [22].

For some genes, such as *yejM*, *ftsN* and *murB*, we identified ORF segments that are essential whereas other regions of the gene can be freely interrupted by transposon insertions. Likely, these non-essential ORF regions exclude protein domains that are involved in critical functional modules of those genes.

Our experiments here involve comparisons among three *Salmonella* strains grown in rich media. However, the study of fitness profiles in hundreds of different strains in multiple growth conditions has the potential to reveal differences in life strategy not evident from the genome sequences alone and to contribute to understanding natural diversity. Advances in DNA sequencing and the ability to incorporate any number of different barcodes for comparison of multiple samples at the same time mean that comparative analysis of fitness among many different natural strains with different phenotypes has become practical. We speculate that as systems biology models become more refined, the fitness profiles of genomes may become useful for constraining these models.

#### Methods

#### Strains and growth conditions

The strains used in this study were *Salmonella enterica* serovar Typhimurium ATCC 14028 (STM1) and *S*. Typhi Ty2 strain JSG624 (STY1) provided by Ferric Fang (University of Washington, Seattle, WA). Bacterial cells were grown in LB medium containing 1% Bacto tryptone (Difco), 0.5% Bacto yeast extract (Becton, Dickinson and Company), and 1% NaCl, supplemented with kanamycin at 50  $\mu$ g/ml, when necessary.

# Construction of transposon integrations in *S*. Typhimurium 14028 and *S*. Typhi Ty2

Salmonella cells were made competent by standard methodology. Briefly, cells were grown in LB with shaking at 37°C to logarithmic phase, then washed three times with cold 10% glycerol and concentrated 250 fold in 10% glycerol. Transposome mixtures were prepared mixing 2  $\mu$ l glycerol, 2  $\mu$ l EZ-Tn5 < T7/KAN-2 > transposon, and 4  $\mu$ l EZ-Tn5 transposase. After 3 h of incubation, 1  $\mu$ l of this mixture was mixed with 50  $\mu$ l of competent cells and 1  $\mu$ l TypeOne restriction inhibitor. Transformation was performed at 2.5 kV using 0.2 cm electrode gap cuvettes and a Bio-Rad MicroPulser at EC2 setting. Transformed cells in each cuvette were resuspended in 1 ml of LB and incubated for 1 h at 37°C. After incubation, reaction aliquots

were joined, complexity was determined by cell counts of various dilutions on LB agar containing kanamycin, and the remainder was grown overnight at 37°C in LB broth supplemented with kanamycin. Stocks of the different transposon mutant libraries containing 20% glycerol were prepared from the overnight cultures.

#### High-throughput sequencing of transposon insertion sites

The entire procedure is illustrated in Additional file 3: Figure S1. In brief, DNA was sheared, poly(A) tailed and PCR amplified using a pair of primers, one located in the transposon and one appended to the poly(A) tail, in a manner similar to that described in Santiviago *et al.* [4]. Subsequently, Illumina sequencing primers were added by PCR and sequencing performed on a Genome Analyzer GAII.

#### Mapping of transposons to the genome

The beginning of each read primer contained a code that defined the particular transposon library used. The codes are listed in Additional file 1: Table S1. The reads were sorted into seven libraries. The sequencing primers were positioned such that the first two bases beyond the primers used for PCR would be the last two bases of the transposon. Reads that included this AG sequence were retained. Most transposons were represented by many shear events. Transposons that were represented by only one or two shear events were distributed throughout the genome, including in known essential regions. Thus, these rare reads were artifactual samples of the whole genome. These sites were eliminated from further consideration.

#### Mapping of genes between genomes

Best hits between Typhimurium 14028, Typhimurium LT2, Typhi CT18, Typhi Ty2 and *E. coli* K-12 were identified by Blast searches of annotated genes against each other in the different genomes. Synteny was determined based on at least one of the two adjacent genes also being a best hit and being adjacent in other genomes. The assigned unique gene numbers differ for all the genome annotations, so all assigned gene numbers and gene symbols in these five genomes are reported in the Additional file 5: Table S2 for the convenience of the reader.

#### Data processing for genome comparisons

Because the distribution and number of transposons is not identical between STM1 and STY1, the data were processed to express the ratio of the number of transposon integrations in each gene and intergenic regions versus the number of integrations in that region that would be expected if all the integrations were randomly distributed. The ratio of observed divided by expected numbers were then expressed as a log<sub>2</sub>. The published STY2 data [3] were recalculated to allow direct comparison.

To identify those genes that had large differences in fitness among STM1, STY1 and STY2, all log<sub>2</sub> ratios (observed number of insertion events/expected number of insertion events) among all 3,907 genes shared between these genomes were ranked from those with the lowest density of transposons to those with the highest density. Ranks in STM1 and both STYs were subsequently subtracted to identify those genes with the biggest difference in rank order of transposon density.

The percentile rank of the number of transposons per base and the number of reads per base (shear events) were calculated for each putative ortholog in each experiment in *Salmonella*. We arbitrarily defined "selected" genes as those having a sum of the two percentile ranks less than 30% (i.e., averaging less than the 15<sup>th</sup> percentile).

#### **Additional files**

Additional file 1: Table S1. Transposon libraries (EZ-Tn5 < T7/Kan-2>) assayed.
Additional file 2: Supplemental Methods. Detailed explanation of protocols.
Additional file 3: Figure S1. Diagram of sequencing protocol.
Additional file 4: Figure S2. Transposon insertion frequency across the S. Typhimurium 14028 s genome.
Additional file 5: Table S2. Essentiality surveys and transposon data.
Additional file 6: Table S3. STY1 complete transposon data.

#### Competing interests

The authors declare that they have no competing interests.

#### Authors' contributions

RC constructed the transposon libraries, and developed the high-throughput sequencing strategy. She helped to write the manuscript and prepare the tables and figures. CF performed the sample preparation for sequencing. SWC coordinated and directed the sequencing project. XQX implemented the trimming and mapping strategy for counting transposons and shear events. He implemented the plots of transposon and shear events across the STM1 genome. BA provided four of the transposon libraries used for screening. HLAP advised on some steps of the project. SP was involved in Typhimurium annotation and manuscript writing. MM devised the sequencing and mapping strategies, helped to analyze the data, and wrote the manuscript.

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