Epistatic Effect of Regulators to the Adaptive Growth of *Escherichia coli*

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Bacteria survive in the environment with three steps: a sensing environmental conditions, a responding to sensed signals, and an adaptation for proper survival in the environment. An adapting bacterial cell occurs cell division to increase the number of sister cells, termed adaptive growth. Two-component systems (TCSs), representing the main bacterial signal transduction systems, consist of a pair of one sensor kinase (SK) and one response regulator (RR), and RR genes are abundant in most bacterial genomes as part of the core genome. The OmpR gene family, a group of RR genes, is conserved in 95% of known bacterial genomes. The *Escherichia coli* genome has an estimated 34 RR genes in total, including 14 genes of OmpR family genes. To reveal the contribution of TCSs for fast growth as an adaptive growth strategy of *E. coli*, we isolated a set of gene knockout strains by using newly developed genome editing technology, the HoSeI (Homologous Sequence Integration) method, based on CRISPR-Cas9. The statistics of single cell observation show a knockout of an arbitrary pair of *phoP*, *phoB*, and *ompR* genes, stably expressed by positive feedback regulation, dramatically inhibit the optimum adaptive growth of *E. coli*. These insights suggest that the adaptive growth of bacteria is fulfilled by the optimum high intracellular level of regulators acquired during growth under environmental conditions.

Bacteria survive and then increase their population by binary cell division in a timely manner. Individual bacterial cells sense changing environmental signals, transduce those environmental signals into biological responses, and adapt to the environmental change by biological responses. Successful integration of these biological stress responses must induce adaptive growth to obtain a continuous chance for the production of offspring. Bacteria are able to survive in various environments by changing the expression pattern of their genome, which takes place by controlling the promoter recognition properties of RNA polymerase by transcription factors¹. Transcription factors (TFs) are classified as those that facilitate direct sensing and indirect sensing of environmental signals. The Lac repressor and Crp activator are good examples of direct sensing TFs and sense lactose and cAMP, respectively. A response regulator (RR) is a typical example of an indirect sensing TF and is activated by phosphorylation by the cognate sensor kinase (SK), detecting environmental signals at the membrane¹,². Direct sensing TFs temporally support bacterial growth because extracellular nutrients stimulate intracellular metabolism and produce signal metabolites that activate those TFs. Indirect sensing TFs, such as RR, are known to be important for temporal survival but are not known for aiding continuous growth.

Recent massive amounts of genomic information show the core genome, a set of species-specific genes, and pan genome, representing non-conserved genes³. Within the core genome, there is a set of genes involved in central and secondary metabolism, cell cycle, and gene expression, many of which are essential for growth⁴. In addition to genes coding for essential biological functions, regulatory genes are conserved in the core genome but are not essential for growth⁵. One of the core-genome regulatory genes is the RR gene, which is a unique signal transduction component that is only conserved among prokaryotes. Two-component systems, known as the common signal transduction systems of prokaryotes, basically consist of a pair of one SK and one RR. SK autophosphorylates upon stimulus and then transfers a phosphate group to a cognate RR. The typical phosphorylated form of a given RR is active and controls the cellular system to survive in changing environments by gene expression regulation because most response regulators harbour structural domains for DNA binding to regulate genome expression⁴. Several TFs are autoregulated during the response to environmental signals. In *Escherichia coli*, more than 50% of identified TFs in the genome are positively or negatively autoregulated⁶. Positive autoregulation leads to a fast response and impacts the response dynamics⁷,⁸. Among ~30 positively autoregulated TFs in the *E. coli* genome⁹, 10 of those are RRs of TCSs, and 34 RRs have been characterized in the *E. coli* genome. On the other hand, negatively autoregulated RRs are less common than positively autoregulated RRs in TCSs⁸. A recent report showed that coupled positive and negative feedback allowed both a fast response and optimal RR protein levels in

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the PhoB/PhoR system in E. coli\(^6\). However, the contribution of coupled autoregulation for continuous bacterial growth is poorly understood.

We found a correlation of the RR family with bacterial genome size, speculating that the main RR genes (phoP, phoB, ompR) affect E. coli growth. To gain insight into the effect of RR genes on growth, we isolated the set of all combinations of single-, double-, and triple-gene knockout strains by using the newly developed HoSeI method for the E. coli genome based on CRISPR-Cas (clustered regularly interspaced short palindromic repeats CRISPR-associated proteins)\(^9\). The differences of elapsed time until the first cell division and initial growth rate of the isolated strains were measured and analysed with statistics, indicating that a particular pair of RR genes of E. coli K-12 are required for adaptive growth via genetic epistasis.

Results
Estimation of highly conserved response regulator genes in the bacterial genome. The Clusters of Orthologous Groups of proteins (COGs) database has been designed to classify proteins on the basis of orthology\(^7\). Twelve groups are detected as two-component response regulators from the genomes of 628 bacteria species in the COG database\(^10\) as follows: OmpR family as COG0745, NarL/FixJ family as COG2197, PleD family as COG3706, NtrC family as COG2204, CheB family as COG2201, LytR/AlgR family as COG3279, AmiR/NasT family as COG3707, FixJ family as COG4566, ActR/RegA family as COG4567, YesN/AraC family as COG4753, GrlR family as COG4565, and SAPR family as COG3947. We found a correlation between the number of RR COGs and genome size, as fitted by the Gompertz curve (Fig. 1A). Bacterial genomes less than 5 Mbp in size contained the relative RR COG number of 2 RR COGs per 1 Mbp of genome, while bacterial genomes more than 5 Mbp in size contained 8 RR COGs (Fig. 1A). Bacterial RNA polymerase core enzyme consist of 2\(\alpha\), \(\beta\), \(\beta^*\), \(\alpha\)I, \(\sigma^70\), which were featured by COG0086, and COG0085, COG0085, COG0086, and COG0085, respectively (Figs. 1B and S2A). The COG0202, COG0085, and COG0086 were almost completely conserved among more than 99% of bacteria registered in COG database\(^10\).

In addition of four COGs featuring bacterial RNA polymerase core subunits, the 8 COGs of RNA polymerase subunits were similarly analysed, resulting that bacterial genomes contained at least 4 RNA polymerase COGs relating RNA polymerase core subunits and major sigma factor and bacterial genomes more than 3 Mbp in size contained 7 RNA polymerase COGs by the addition of COGs relating minor sigma factors (Fig. S1). Among the 628 species of bacteria, conservation of RR COGs is the highest in the OmpR family, at 95%, and the lowest in the SAPR family, at 8.4% (Fig. 1B), suggesting that the OmpR family is extensively conserved in bacteria as a member of the core genome.

The copy number of RR genes of the E. coli K-12 genome was evaluated in detail. The genome sequence of E. coli K-12, which is 4.6 Mbp in size, has 34 predicted response regulator genes, of which 6 are classified as RR COGs\(^12,13\). Among all 34 RR genes, OmpR family genes are the most prevalent, with 14 genes, arcA, phoB, cusR, kdpE, torR, phoP, rtaA, hprR, basR, qseB, ompR, cpxR, creB, and cbbR, and the Crr family and LrtT family have the smallest number, at 2 genes, with citsB and dcrB and with btsR and ppyR, respectively (Figs. 1B and S2A). Six RR genes of the OmpR family are known to be positively self-activated (Fig. 1C)\(^14,15\). In good agreement with the positive feedback regulation of the OmpR family RRs of E. coli K-12, OmpR, PhoB, and PhoP are abundant regulators in E. coli through the growth phase in rich and poor medium, with approximately 1,000 copies per genome\(^16\) (Fig. S2B). Taking these observations together, we suspected that OmpR, PhoB, and PhoP play important roles in the adaptive growth of fast-growing E. coli as main RR factors.

Homologous sequence integration (HoSeI) method for multi-gene knockout in the E. coli genome by introduction of a nonsense codon using CRISPR-Cas. The CRISPR-Cas system is available to knock out genes in the E. coli genome by insertion of an antibiotic resistance marker gene\(^21\). We developed a novel gene knockout system by introducing a nonsense codon, a homologous sequence integration (HoSeI) method, to overcome the limitations of antibiotic resistance marker genes. It is readily feasible for us to design and isolate multi-gene knockout strains of E. coli. The designed sgRNA (single-guide RNA) containing the recognition sequence for the E. coli genome specifically digests the genome, resulting in a dead phenotype (Fig. S3A). The designed DNA fragment containing the following sequence, however, recombines with the genome at the injury site by lambda-Red recombinase, resulting in a living phenotype (Fig. S3A). To knock out gene function, the DNA fragment is designed to introduce the stop codon, TAA, instead of the PAM sequence (Fig. S3B). For knockout of each gene, we constructed sgRNA-expressing plasmids (Table S2). None of the constructed psgRNA plasmids caused E. coli harbouring pCas to gain ampicillin resistance (data not shown). The addition of the designed DNA fragment, including the nonsense codon for the target gene, recovered the transformation of E. coli harbouring pCas by psgRNA. The proper introduction of the nonsense codon, TAA, was confirmed by Sanger sequencing in all cases (Fig. S4). We isolated a psgRNA-free transformant by IPTG-inducible sgRNA for the psgRNA plasmid from pCas as previously described\(^21\). The isolated psgRNA-free transformant was subjected to further gene knockout by repeating the HoSeI method. In addition to all single knockouts of the 14 OmpR family RRs, these experiments bred all strains of seven combinations of main RR gene knockout for PhoB, PhoP, and OmpR (Table S1).

Contribution of the optimum adaptive growth of E. coli by RR genes. Bacteria are able to grow both in liquid and solid conditions. In both cases, the number of bacteria cells similarly increases\(^22\). To measure the elapsed time until cell division and growth rate of single E. coli cells, time-lapse observation of single cells growing on solid medium was adopted in a 30 min range for 8 h with light microscopy. At the end of observation after 8 hours, 70.8% of the parent strain, W3110 typeA, cells formed micro-colonies including 4 to 8 cells, while 8.3% of the cells were not completely divided (Fig. S5a). The majority of parent cells started cell division 0.5–4.5 hours after incubation (80.2%) and showed a growth rate of 0.5–1.1 divisions/hour (75.0%) (Figs. 2 and S5a). Fourteen single OmpR family RR gene knockout strains isolated in this study were subjected to time-lapse

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observation, as was the parent strain. Most of the single RR gene knockouts, except for \( \Delta \) ompR, showed that the percentage of non-growing cells was in the range of 17% to 1.5% at the end of observation after 8 hours, with a maximum of 17% for \( \Delta \) rstA and a minimum of 1.5% for \( \Delta \) cpxR (Figs. 2A and S5). All of the \( \Delta \) ompR strain cells formed micro-colonies (Figs. 2A and S5k). For all of the single RR gene knockout strains, more than 75% of cells started cell division in the time range of 0.5 - 4.5 hours, similar to the parent strain, with several deviations (Fig. 2A). The growth rates were divided into three groups with comparison to the parent strain: the faster growth rate mutants were \( \Delta \) phoB, \( \Delta \) cusR, \( \Delta \) torR, \( \Delta \) rstA, \( \Delta \) hprR, \( \Delta \) baer, \( \Delta \) qseB, \( \Delta \) ompR, \( \Delta \) cpxR, \( \Delta \) basR, and \( \Delta \) arcA; the similar growth rate mutants were \( \Delta \) phoP and \( \Delta \) creB; and the slow growth rate mutant was \( \Delta \) kdpE (Fig. 2B).

Figure 1. Conservation of two-component system response regulator families in bacterial genomes. (A) The correlation between genome size and the number of RR families in the genomes. Twelve groups are detected as two-component response regulators from the bacterial genomes of 628 species in the COG database as follows: OmpR family as COG0745, NarL/FixJ family as COG2197, PleD family as COG3706, NtrC family as COG2204, CheB family as COG2201, LytR/AlgR family as COG3279, AmiR/NasT family as COG3707, FixJ family as COG4566, ActR/RegA family as COG4567, YesN/AreC family as COG4753, CriR family as COG4565, and SAPR family as COG3947. The number of COGs involved in the RR family (y-axis) was analysed in comparison with the genome size (x-axis). The average number of COGs (black circle) and standard deviation (SD, error bar) were calculated for each genome size of 0.5, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 12, and 15 Mb. The calculated data were fitted by the Gompertz curve \( y = a e^{bx} + c \), where \( y \) is the number of COGs and \( x \) is the genome size. The formula of the fitted Gompertz curve is shown in the graph. (B) Conservation of RR families in the bacterial genomes and gene copy number in the E. coli genome. Based on the calculated data shown in (A), the conservation (%) of each COG was calculated on all genomes of 628 species of bacteria (x-axis). The y-axis shows the gene copy number of each RR family in the E. coli K-12 genome. (C) OmpR family RR genes in the E. coli K-12 genome. E. coli K-12 W3110 has 14 OmpR family response regulator genes in its genome. Each arrow shows OmpR family response regulators (white), their cognate sensor kinases (gray), and other genes in the operon (light gray). The direction of arrows shows the direction of genes in the E. coli genome. Red letters and arrows show the genes known as positive feedback regulators.
The adaptive growth of *E. coli* defected by a knockout of an arbitrary pair of *phoP*, *phoB*, and *ompR* genes. Next, three double main RR gene knockout strains, Δ*phoP* Δ*phoB*, Δ*phoP* Δ*ompR*, and Δ*phoB* Δ*ompR*, were isolated from two single RR knockout strains, Δ*phoB* and Δ*phoP* (Table S1) and subjected to time-lapse observation to determine the elapsed time until cell division and the growth rate. Most of the cells of all of the three double RR gene knockout strains did not significantly grow at the end of observation after 8 hours (Fig. S6), resulting in a long elapsed time and slow growth rate (Figs. 3A and S6). However, all three double RR gene knockout strains formed colonies after incubation for more than 48 hours under the same conditions (data not shown). Surprisingly, the triple RR gene knockout strain showed similar behaviour to that of the parent strain. At the end of observation after 8 hours, 93.1% of Δ*phoP* Δ*phoB* Δ*ompR* strain cells formed micro-colonies, while 6.9% of them were not totally divided (Fig. S6). The majority of the Δ*phoP* Δ*phoB* Δ*ompR* strain cells started cell division at 1.5–3.5 hours after incubation (86.2%) and showed a growth rate of 0.6–1.4 divisions/hour (79.3%), as did the parent strain (Figs. 3A, 5B, and S7).

We examined to confirm gene knockout of an arbitrary pair of *phoP*, *phoB*, and *ompR* genes, defecting the adaptive growth, by genetic complement test using luciferase reporter system (Fig. 3B). The *ompC*, *pstS*, and *mgtA* promoters were employed as target promoters by EnvZ-OmpR, PhoQ-PhoP, PhoR-PhoB. The activation of *ompC* promoter requires the phosphorylated OmpR to upstream of *ompC* promoter by *ompC-lux* reporter plasmid (Fig. S5), resulting in a long elapsed time and slow growth rate (Figs. 3A and S6). However, all three double RR gene knockout strains formed colonies after incubation for more than 48 hours under the same conditions (data not shown). Surprisingly, the triple RR gene knockout strain showed similar behaviour to that of the parent strain. At the end of observation after 8 hours, 93.1% of Δ*phoP* Δ*phoB* Δ*ompR* strain cells formed micro-colonies, while 6.9% of them were not totally divided (Fig. S6). The majority of the Δ*phoP* Δ*phoB* Δ*ompR* strain cells started cell division at 1.5–3.5 hours after incubation (86.2%) and showed a growth rate of 0.6–1.4 divisions/hour (79.3%), as did the parent strain (Figs. 3A, 5B, and S7).

Taking all these observations together, OmpR, PhoP, and PhoB each slightly contribute to the growth of *E. coli*, while the disappearance of any two OmpR, PhoP, and PhoB proteins seriously disrupts the adaptive growth of fast-growing *E. coli*.

The adaptive growth of *E. coli* not defected by a knockout of an arbitrary pair of *envZ*, *phoR*, and *phoQ* genes. EnvZ, PhoR, and PhoQ are the cognate SKs of OmpR, PhoB, and PhoP, respectively. Three single SK gene knockout strains, Δ*envZ*, Δ*phoR*, and Δ*phoQ*, were subjected to time-lapse observation as described above. At the end of observation after 8 hours, the percentages of non-growing cells of Δ*envZ*, Δ*phoR*, and Δ*phoQ* were 32.0%, 20.3%, and 21.5%, respectively (Fig. S6). In the case of Δ*envZ*, 54.0% of cells started cell division 4.0–6.5 hours after incubation, and 50.0% showed a growth rate of 0.8–2.0 divisions/hour, similar to the parent strain (Figs. 3C, S6i and S7). In the case of Δ*phoQ*, 61.5% of cells started cell division 4.0–5.5 hours after incubation, and 67.7% showed a growth rate of 0.3–1.0 divisions/hour, similar to the parent strain (Figs. 3B, S6i and S7). In the case of Δ*phoR*, 62.5% of cells started cell division 1.5–4.5 hours after incubation, and 73.4%...
Figure 3. Contribution of main two-component RRs to the adaptive growth of *E. coli* cells. Time-lapse observation of single cells growing on solid medium was adopted in the 30 min range for 8 hours with a light microscope (Fig. S6). Based on the measured data (Fig. S6), the elapsed time until cell division (hours, shown on the y-axis) and growth rate of a single *E. coli* cell (divisions/hour, shown on the x-axis) were calculated and are shown as contour graphs (A,C, and D). (A) The parent strain (blue) and three of the main RR (*phoP, phoB*, and *ompR*) gene knockout strains, Δ*phoP* (light blue), Δ*phoB* (gray), Δ*ompR* (yellow), Δ*phoP* Δ*phoB* (pale blue), Δ*phoP* Δ*ompR* (yellowish green), Δ*phoB* Δ*ompR* (red), and Δ*phoP* Δ*phoB* Δ*ompR* (dark brown). (B) The constructed pLux-mgtA (a), pLux-pstS (b), and pLux-ompC (c), shown in Table S2, were transformed into each strain (a-1,b-1, and c-1) and each strains carrying either RR protein expression plasmids, pBADPhoP-FLAG (a-3), pBADPhoB-FLAG (b-3), or pBADompR-FLAG (c-3) or the empty pBAD33 vector (a-2,b-2, and c-2). Transformants were grown in the M9-glucose medium including 50 μg/mL kanamycin for pLUX plasmid and 20 μg/mL chloramphenicol for pBAD plasmid at 37 °C with shaking for overnight. And then, overnight culture was diluted 100-fold by fresh M9-glucose medium with (a-2 and 3, b-2 and 3, and c-2 and 3) and without (a-1,b-1, and c-1) 0.002% arabinose for RR protein expression from pBAD plasmid and incubated at 37 °C with shaking. OD600 and a total intensity of luminescence were measured and the ratio of luminescence to OD600 (LUX/OD) of each strain was calculated. The specific activity for each promoter was normalized with the parent strain W3110. (C) The parent strain (blue), cognate SK genes of main RR (*phoP, phoB*, and *ompR*) knockout strains Δ*phoQ* (light blue), Δ*phoR* (gray), Δ*envZ* (yellow), Δ*phoQ* Δ*phoR* (pale blue), Δ*phoQ* Δ*envZ* (yellowish green), Δ*phoR* Δ*envZ* (red), and Δ*phoQ* Δ*phoR* Δ*envZ* (dark brown), and cognate pairs of SK-RR knockout strains Δ*phoP* Δ*phoQ* (pink), Δ*phoB* Δ*phoR* (light brown), and Δ*ompR* Δ*envZ* (navy). (D) The parent strain (blue), double RR gene knockout strains Δ*phoP* Δ*phoB* (pale blue), Δ*phoP* Δ*ompR* (yellowish green), Δ*phoB* Δ*ompR* (red), Δ*KdpE* (pink), Δ*creB* (green), Δ*ompR* Δ*cpxR* (orange), and Δ*ompR* Δ*rstA* (light blue), and a triple RR gene knockout strain Δ*rstA* Δ*cusR* Δ*hprR* (purple).
showed a growth rate of 0.4–1.4 divisions/hour, similar to the parent strain (Figs. 3C, S6j and S7). Overall, three single SK gene mutants showed slightly delayed elapsed times, but their growth rate was similar to that of the parent strain.

Next, three double SK gene knockout strains, Δ phoQ Δ phoR, Δ phoQ Δ envZ, and Δ phoB Δ envZ, were subjected to time-lapse observation to determine the elapsed time until cell division and the growth rate. Most of the cells of the three double SK gene knockout strains grew significantly at the end of observation after 8 hours (Fig. S6). All three double SK gene knockout strains showed an elapsed time in the range of 2.0–3.5 hours, and the growth rate varied in the range of 0.8–2.4 divisions/hour (Figs. 3B, S6 and S7). In addition, for the triple SK gene knockout strain, 92.6% of cells formed micro-colonies (Fig. S6). The majority of the Δ phoQ Δ phoB Δ envZ strain cells started cell division 1.5–4.5 hours after incubation (77.8%) and showed a growth rate of 0.4–2.0 divisions/hour (74.1%) (Figs. 3C, S6o and S7).

Taking all these observations together, the disappearance of any two OmpR, PhoP, and PhoB proteins seriously disrupts the adaptive growth of fast-growing E. coli, while the disappearance of any two EnvZ, PhoR, and PhoQ, the cognate SKs for OmpR, PhoB, and PhoQ, proteins slightly affect the adaptive growth of fast-growing E. coli.

**A specific arbitrary pair of phoP, phoB, and ompR genes for double gene knockout inhibiting the optimum adaptive growth of E. coli.** All three double RR gene knockout strains, Δ phoP Δ phoB, Δ phoP Δ ompR, and Δ phoB Δ ompR, almost disrupted the adaptive growth. We examined whether double RR gene knockout of other combinations. Cluster analysis with all RR of E. coli K-12 showed that PhoP, PhoB, and OmpR were highly similar RRs (Fig. S2). Therefore, we knocked out a pair of similar genes and isolated four double RR knockout strains of Δ phoP Δ kdpE, Δ phoB Δ creB, Δ ompR Δ cpxR, and Δ ompR Δ rstA (see above) (Fig. S4). Most of the cells of the Δ phoP Δ kdpE, Δ phoB Δ creB, Δ ompR Δ cpxR, and Δ ompR Δ rstA strains grew significantly at the end of observation after 8 hours (Fig. S6). Three double RR gene knockout strains, Δ phoP Δ kdpE, Δ phoB Δ creB, and Δ ompR Δ cpxR, showed an elapsed time in the range of 2.5–4.0 hours, and the growth rate varied in the range of 0.75–1.5 divisions/hour (Figs. 3D, S6 and S7). On the other hand, Δ ompR Δ rstA strains showed an elapsed time in the range of 2.5–7.5 hours, and the growth rate varied in the range of 0.05–1.2 divisions/hour (Figs. 3D, S6 and S7). Furthermore, another triple RR gene knockout strain, Δ rstA Δ casR Δ hrpR, also presented micro-colony formation in 92.6% of the population, starting cell division 1.5–4.5 hours after incubation (83.9%) and showing a growth rate of 0.6–2.0 divisions/hour (69.6%), as did the Δ phoQ Δ phoB Δ envZ strain (Figs. 3C, S6w and S7).

We next examined three cognate pair double gene knockout strains, Δ phoP Δ phoQ, Δ phoB Δ phoR, and Δ ompR Δ envZ, for adaptive growth. Most of the cells of the three cognate pair double gene knockout strains grew significantly at the end of observation after 8 hours (Fig. S6). These three strains showed an elapsed time in the range of 1.0–3.5 hours, but their growth rates varied (Figs. 3B and S6, S7). The Δ phoP Δ phoQ and Δ ompR Δ envZ strains showed growth rates in the range of 1.3–2.7 divisions/hour (70%) and 1.3–2.7 divisions/hour (70%), respectively, whereas 70% of cells of the Δ phoB Δ phoR strain showed growth rates of 0.4–1.4 divisions/hour, similar to the parent strain (Figs. 3B and S7). In agreement with observations of single knockout strains as shown above, these observations showed that the roles of PhoP, PhoB, and OmpR did not completely correspond with their cognate SKs, PhoQ, PhoR, and EnvZ, respectively, in adaptive growth.

To demonstrate the genetic involvement of TCSs for adaptive growth, cluster analysis was performed with the elapsed time and the growth rate on solid medium (Fig. S8). The dendrogram of the elapsed time showed two groups, each divided into two sub-groups (Fig. S8a). The parent strain was assigned to Group-Ib, which included not only all single RR gene knockout strains but also all double SK gene knockout strains and triple RR and SK gene knockout strains (Fig. S8a). Additionally, all cognate pair gene knockout strains were assigned to Group-Ia (Fig. S8a). All of the single SK gene knockout strains were assigned to Group II, in which all of the double RR gene knockout strains were part of Group Iib (Fig. S8a). The dendrogram of the growth rate also showed two groups (Fig. S8b). The parent strain was assigned to Group II, which included most of the knockout strains, whereas all of the double RR gene knockout strains were isolated in Group I (Fig. S8b).

Cluster analysis of the elapsed time also showed that all four of the double RR gene knockout strains studied were assigned to Group-Ia but not to Group-II (Fig. S8a). Additionally, clustering for the growth rate showed that all four of the double RR gene knockout strains were not assigned to Group I (Fig. S8b). Taken together, knockout of the double RR gene among ompR, phoP, and phoB, specifically caused a long delay in the start of growth and a lower growth rate (Fig. 3).

**Discussion**

Bacteria are unicellular organisms. Individual cells are known to start growth under rich nutrient conditions by operating metabolic networks. On the other hand, bacterial cells that survive have a proper memory for the function of genetic stress response systems. Thus, multi-genetic factors function for bacterial growth. To analyse the function of several genes, we developed the CRISPR-Cas9-based HoSeI method to knock out genes without antibiotic resistance gene markers. The HoSeI method is a genetic marker-less genome editing approach and introduces base substitutions in the target sequence on the original genome by screening dead or alive cells. We repeated the HoSeI method in the Pseudomonas aeruginosa K-12 genome and isolated multi-gene knockout strains. This should be a useful experimental demonstrations of bacterial genome-wide epistatic phenomena.

An OmpR family comparison of Haemophilus influenzae Rd KW20, Corynebacterium glutamicum ATCC 13032, Bacillus subtilis subsp. subtilis str. 168, Mycobacterium tuberculosis H37Rv, Shigella dysenteriae 1617, Escherichia coli str. K-12 substr. MG1655, Salmonella enterica subsp. enterica serovar Typhimurium str. LT2, and Pseudomonas aeruginosa PAO1 reveals that the four sub-families actually divided PhoP, PhoB, and OmpR...
suggesting that PhoP, PhoB, and OmpR each basically function as a DNA-binding protein. Genome-wide binding of phoQ, single and PhoB, or OmpR. could not properly grow, if insignificant occupation of the genome by 1,000 intracellular proteins of either PhoP, PhoB, and OmpR have become associated with the genome.

Thus, PhoB, PhoP, and OmpR was not essential for adaptive growth of E. coli. PhoB, PhoP, and OmpR was not essential for adaptive growth of E. coli, H. influenzae, but also in bacteria with smaller (P. aeruginosa) genomes than that of E. coli. A genome comparison of E. coli strains reveals that the predicted pan-genome comprises 15,741 gene families and that the core genome comprises only 993 (6%) of the families, as represented in every genome. OmpR family genes are included in the E. coli core genome and increase their number in the genome. We found that the OmpR family includes conserved TCS genes withing the bacterial genome. The number of the OmpR family gene is also correlated with the size of the genome (Fig. 4A).

Additionally, OmpR family RRs are conserved not only in closely related bacteria with genome sizes similar to that of E. coli but also in bacteria with smaller (H. influenzae) or larger (P. aeruginosa) genomes than that of E. coli. UhpA - decreasing order for the intracellular response regulators: PhoP -> PhoB -> CitB -> OmpR -> NarL -> CpxR -> UhpA -> RstA -> EnvG -> KdpE -> TorR -> Uvry -> QseB -> ArcA29. Among these 14 more abundant species, 9 species of RRs belong to the OmpR family (PhoP, PhoB, OmpR, RstA, KdpE, TorR, QseB, and ArcA). We isolated all of 14 OmpR family single RR gene knockout strains by HoSeI method, indicating that none of them was essential for growth under the used conditions in good agreement with previous works30,31. Time-lapse live cell imaging with microscope indicated that 14 single RR gene knockout strains were divided into three groups, the faster growth rate mutants, the normal growth rate mutants, and the slow growth rate mutant (Fig. 2B). The difference did not appear to correlate with the intracellular level or the number of regulated genes.

As unexpected, we found that knockout of the arbitrary pair of phoP, phoB, and ompR are shown as an epistatic inhibition for E. coli adaptive growth (Fig. 3). When the species of three genes knockout was introduced into single phoP knockout strain (Table S1), only double defect with ompR gene dramatically inhibited the adaptive growth of E. coli (Fig. 3). When the species of four genes knockout was introduced into single phoB knockout strain (Table S1), only double defect with either ompR or phoB gene dramatically inhibited the adaptive growth of E. coli (Fig. 3).

Besides, the knockout for citB gene, one of NarL family RR gene of E. coli, was introduced into both single phoB and single phoP knockout strains, resulting that both double gene knockout strains, ΔcitB ΔphoB and ΔcitB ΔphoP, did not show significant inhibition of the adaptive growth of E. coli (data not shown). Despite different genetic backgrounds, the genetic defect of an arbitrary pair of phoP, phoB, and ompR largely delays initiation of growth and dramatically reduces the initial growth rate, the results of which were entirely different from not only phoP phoB, and ompR RR single knockouts but also their triple knockouts (Figs. 3 and S8). In good agreement with the most abundant RRs, PhoP, PhoB, and OmpR29, a memory of expression of genes below cytotoxic levels exists for positively autoregulated systems of the genes phoP, phoB, and ompR in E. coli. Each OmpR family member, including E. coli PhoP, PhoB, and OmpR, contains an N-terminal receiver domain (RD) and a C-terminal DNA-binding domain (DBD), which are connected by a linker. To date, the molecular structure of the truncated RD and DBD and the full-length active and inactive form of OmpR family proteins have been solved by X-ray crystallography and NMR spectroscopy24–26. Among these proteins, dimerization or higher-order multimerization is commonly observed, and at least three surfaces have been observed for intermolecules. Two surfaces, the α1α5 and the α4/35α5 faces, conserve the RD and one surface the DBD. PhoP, PhoB, and OmpR are known to bind to at least 15 promoters in vivo, as denoted in RegulonDB (http://regulondb.ccg.unam.mx/index.jsp), indicating that the intracellular level of approximately 1,000 copies of PhoP, PhoB, and OmpR each per genome is clearly too high, and thus, these proteins could play a role except in the stress response. These are in good agreement with the observation of the adaptive growth of E. coli of a knockout of an arbitrary pair of envZ, phoR, and phoQ genes, the cognate SK genes for OmpR, PhoB, and PhoP (Figs. 3C, S6w and S7). PhoB and OmpR are also known to bind to more than 100 genomic sites in vitro denoted in TEC (https://shigen.nig.ac.jp/E.coli/tec/top/), suggesting that PhoP, PhoB, and OmpR each basically function as a DNA-binding protein. Genome-wide binding profiles have indicated the recognized DNA sequences TTTAnnnTTTA as the PhoP-binding consensus37, TTTGTACAT as the OmpR-binding consensus38, and TGTnAAnAAAnTGTnA as the PhoB-binding consensus39. The tandem triplet of A or T, found as a common sequence among the genes, could be weakly recognized by every one of PhoB, PhoP, and OmpR. Thus, E. coli cells could show adaptive growth when a total of 3,000 intracellular proteins of PhoP, PhoB, and OmpR have become associated with the genome. E. coli cells, however, could not properly grow, if insignificant occupation of the genome by 1,000 intracellular proteins of either PhoP, PhoB, or OmpR.

The triple RR gene knockout strain, however, recovered the elapsed time and the growth rate to values similar to those for the parent strain (Fig. 2). This observation showed that a total of 3,000 intracellular proteins of PhoP, PhoB, and OmpR was not essential for adaptive growth of E. coli. One possible reason that normal growth of the triple RR gene knockout strain is the existence of excess intracellular RRs in place of PhoP, PhoB, and OmpR. We did not, however, find a significant difference in the profile of cellular proteins, with more than 200 proteins detected with LC-MS, between the parent strain and the triple RR gene knockout strain (data not shown).

Additionally, the promoter activities of all TCS genes in the E. coli genome were measured using the luciferase reporter system, indicating that three double RR gene knockout strains and a triple RR gene knockout strain were mostly similar to the parent strain with respect to these promoter activities (data not shown).

These findings suggest that the mode of action for inhibition of adaptive growth by knockout of the arbitrary pair of phoP, phoB, and ompR is important for the bacterial cell cycle, genome replication and/or cell division.
Figure 4. Comparison of OmpR family RRs among several bacteria. (A) OmpR family RR gene copy number of several bacteria. On the basis of the data shown in Fig. 1, the copy number of OmpR family RR genes (y-axis) in eight bacterial genomes was compared with the respective genome size (x-axis). The eight bacterial strains are Haemophilus influenzae Rd KW20, Corynebacterium glutamicum ATCC 13032, Bacillus subtilis subsp. subtilis str. 168, Mycobacterium tuberculosis H37Rv, Shigella dysenteriae 1617, Escherichia coli str. K-12 substr. MG1655, Salmonella enterica subsp. enterica serovar Typhimurium str. LT2, and Pseudomonas aeruginosa PA01. (B) Phylogenetic analysis of the amino acid sequence of OmpR family RRs. All of the amino acid sequences of the OmpR family RRs in 8 bacteria shown in (A) were compared and analysed with ClustalW software. The result divided the RRs into four sub-families. The amino acid sequence of the used OmpR family RRs was compiled in Table S4.
but not for general metabolism and stress responses. The epistatic inhibition for *E. coli* adaptive growth by an arbitrary pair of *phoP*, *phoB*, and *ompR* was obscure on LB agar (data not shown). All of the isolated knockout strains also showed a similar adaptive growth to the parent strain on LB agar (data not shown). One possible is that only either *PhoP*, *PhoB*, or *OmpR* at a high intracellular level bias genome expression profile and inhibit the optimum adaptive growth in *E. coli* grown in M9 glucose that contains less amino acids than LB broth. Another is that a high concentration of only one species of OmpR family severely affect bacterial growth by an unknown molecular mechanism, which could be suppressed by a high concentration of another species of OmpR family. These findings suggest that RRs contribute to the optical high level of the intracellular RRs, resulting in a fitness advantage for growth. The results are in good agreement with the idea of fitness advantages being conferred for both positive and negative feedback in both dynamic and stable environments. Taken together, in *E. coli*, *PhoP*, *PhoB*, and *OmpR* are autoregulated at the optimal intracellular level and balance of their intracellular level play an important role in survival for repeatable growth fitness under the evaluated experimental conditions.

**Methods**

**E. coli** strains, plasmids, and oligonucleotides. The used *E. coli* strains, plasmids, and oligonucleotides are listed in Tables S1, S2 and S3, respectively. *E. coli* K-12 W3110 typeA strain was used as a parent type.

**Growth condition of E. coli.** *E. coli* cells were grown at 30 °C in M9-glucose medium.

**Construction of sgRNA expression plasmid.** To construct the sgRNA expression vector plasmid, 227 bp of the DNA sequence of the promoter and sgRNA was referred to as the pTarget series harbouring sgRNA-NAs constructed by previous work. The designed 227 bp DNA sequence introducing the Not I recognition site immediately upstream of tracrRNA (trans-activating CRISPR RNA) was synthesized and cloned into the pEX-A2 vector by Eurofin genomics (Tokyo, Japan), resulting in the construction of psgRNA (Table S2). Insertion DNA for the target sequence was prepared by hybridization of a pair of complementary synthetic oligonucleotides in length of 50-nt each: ompR_sgRNA_N20 and ompR_sgRNA_com for the *ompR* gene; *phoP* sgRNA_N20 and *phoP*_sgRNA_com for the *phoP* gene; *envZ*_sgRNA_N20 and *envZ*_sgRNA_com for the *envZ* gene; *phoR* sgRNA_N20 and *phoR*_sgRNA_com for the *phoR* gene; *phoQ*_sgRNA_N20 and *phoQ*_sgRNA_com for the *phoQ* gene; *kdpE*_sgRNA_N20 and *kdpE*_sgRNA_com for the *kdpE* gene; *creB*_sgRNA_N20 and *creB*_sgRNA_com for the *creB* gene; *cpxR*_sgRNA_N20 and *cpxR*_sgRNA_com for the *cpxR* gene; *rstA*_sgRNA_N20 and *rstA*_sgRNA_com for the *rstA* gene; *cysR*_sgRNA_N20 and *cysR*_sgRNA_com for the *cysR* gene; *hprR*_sgRNA_N20 and *hprR*_sgRNA_com for the *hprR* gene; *arcA*_sgRNA_N20 and *arcA*_sgRNA_com for the *arcA* gene; *basR*_sgRNA_N20 and *basR*_sgRNA_com for the *basR* gene; *baeR*_sgRNA_N20 and *baeR*_sgRNA_com for the *baeR* gene; *qseB*_sgRNA_N20 and *qseB*_sgRNA_com for the *qseB* gene; and *torR*_sgRNA_N20 and *torR*_sgRNA_com for the *torR* gene (see DNA sequence shown in Table S3). The DNA was inserted into the *Not I* site of psgRNA vector by the In-Fusion system (Takara Bio, Japan). The inserted DNA sequence on the resulting plasmids was confirmed by DNA sequencing (Table S3).

**Multi-gene knockout in the E. coli genome by the HoSeI method based on CRISPR-Cas9.** The W3110 typeA strain harbouring pCas was grown in LB medium containing 1% arabinose and 50 μg/mL kanamycin to logarithmic phase and then was collected and suspended in a solution of 0.1 M CaCl₂. This suspension of *E. coli* was subjected to transformation by the sgRNA-target and DNA fragment to recover the digested site by CRISPR-Cas9. sgRNA-ompR, sgRNA-phoB, sgRNA-phoP, sgRNA-envZ, sgRNA-phoQ, sgRNA-kdpE, sgRNA-creB, sgRNA-cpxR, sgRNA-rstA, sgRNA-cysR, sgRNA-hprR, sgRNA-arcA, sgRNA-basR, sgRNA-baeR, sgRNA-qseB, and sgRNA-torR were used as sgRNA-target plasmids (Table S2). The 83-bp DNA fragments were prepared by hybridization of a pair of complementary synthetic oligonucleotides: ompR_PAMstop and ompR_PAMstop_com for the *ompR* gene; *phoP*_PAMstop and *phoP*_PAMstop_com for the *phoP* gene; envZ_PAMstop and envZ_PAMstop_com for the *envZ* gene; *phoR*_PAMstop and *phoR*_PAMstop_com for the *phoR* gene; *phoQ*_PAMstop and *phoQ*_PAMstop_com for the *phoQ* gene; *kdpE*_PAMstop and *kdpE*_PAMstop_com for the *kdpE* gene; *creB*_PAMstop and *creB*_PAMstop_com for the *creB* gene; *cpxR*_PAMstop and *cpxR*_PAMstop_com for the *cpxR* gene; *rstA*_PAMstop and *rstA*_PAMstop_com for the *rstA* gene; *cysR*_PAMstop and *cysR*_PAMstop_com for the *cysR* gene; *hprR*_PAMstop and *hprR*_PAMstop_com for the *hprR* gene; *arcA*_PAMstop and *arcA*_PAMstop_com for the *arcA* gene; *basR*_PAMstop and *basR*_PAMstop_com for the *basR* gene; *baeR*_PAMstop and *baeR*_PAMstop_com for the *baeR* gene; *qseB*_PAMstop and *qseB*_PAMstop_com for the *qseB* gene; and *torR*_PAMstop and *torR*_PAMstop_com for the *torR* gene (see DNA sequence shown in Table S3). The DNA was inserted into the *Not I* site of psgRNA vector by the In-Fusion system (Takara Bio, Japan). The inserted DNA sequence on the resulting plasmids was confirmed by DNA sequencing (Table S3).

**Time-lapse observation on microscope.** Strains were inoculated in M9 glucose medium and shaken overnight at 37 °C. The cultures were washed with M9 medium three times and diluted 10-fold into the medium. The diluted cultures were spread on an M9 glucose plate. Next, the plates were cut and placed on a slide glass. The preparation was sealed on a glass coverslip with nail polish. The cells were imaged with a microscope (IX81, Olympus) using a 100x/NA 1.4 objective lens (M Plan Apochromat MPLAPON-Oil, Olympus) and a Retiga EXi Fast1394 CCD camera (Q Imaging). The temperature was maintained at 37 °C using a closed circulation system.
Construction of RR protein expression plasmid. We constructed the OmpR, PhoB, and PhoP expression plasmids as well as arabinose-inducible expression system of NarL family RR as previously described\(^4\). In brief, the protein-coding regions of phoP, phoB, and ompR genes with an artificial SD sequence were amplified by PCR using specific primers (Table S3). Each amplified DNA fragment was inserted into the linear pBAD33 vector by In Fusion system (Takara Bio, Japan), resulting in construction of the FLAG-tagged RR protein expression plasmids, pBADPhoP-FLAG, pBADPhoB-FLAG, and pBADOmpR-FLAG (Table S2).

Luciferase reporter assay in *E. coli*. We performed luciferase reporter assay in *E. coli* to evaluate activity of TCS-target promoters in vivo as previously described\(^2,4\). In brief, the ompC, mgtA, pstS promoters were amplified by PCR using W3110 type A genome as a template, a pair of primers (see DNA sequence shown in Table S3), and Ex Taq DNA polymerase (Takara Bio, Japan). The promoter DNA was inserted into the *Bam* HI and *Xho* I sites of pLUX vector\(^4\) by the In-Fusion system (Takara Bio, Japan). The inserted DNA sequence on the resulting plasmid was confirmed by DNA sequencing (Table S2). The constructed pLUX-mgtA, pLUX-pstS, and pLUX-ompC were transformed into each strain. Transformants were grown in the M9-glucose medium at 37 °C with shaking for overnight. Then, overnight culture was transferred into fresh M9-glucose medium and OD\(_{600}\) and a total intensity of luminescence of culture was measured with a plate reader (Corona). The ratio of luminescence to OD\(_{600}\) (LUX/OD) was evaluated as a specific activity of the promoter.

Cluster analysis. Cluster analysis was performed with R software. The obtained data of the elapsed time and the growth rate were prepared for Ward’s method. A hierarchical clustering dendrogram was calculated using Euclidean distances.

Correlation of the number of COGs and the genome size with the Gompertz function. We used all of the genome sequences of 628 species of bacteria registered in the COG databank. TCS genes and RNA polymerase genes were isolated, and the number of these genes was counted for each bacterial genome. The number of COGs involved in TCSs and the RNA polymerase subunit was analysed in comparison with the genome size, and the average number of COGs was calculated for each genome size of 0.5, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 12, and 15 Mb. The calculated data were fitted by the Gompertz curve \(y = ab^x e^{-cx}\), where \(y\) is the number of COGs and \(x\) is the genome size.

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