Lethal Consequences of Overcoming Metabolic Restrictions Imposed on a Cooperative Bacterial Population

Eunhye Goo, Yongsung Kang, Jae Yun Lim, Hyeonheui Ham, Ingyu Hwang
Department of Agricultural Biotechnology, Seoul National University, Seoul, Republic of Korea

ABSTRACT
Quorum sensing (QS) controls cooperative activities in many Proteobacteria. In some species, QS-dependent specific metabolism contributes to the stability of the cooperation. However, the mechanism by which QS and metabolic networks have coevolved to support stable public good cooperation and maintenance of the cooperative group remains unknown. Here we explored the underlying mechanisms of QS-controlled central metabolism in the evolutionary aspects of cooperation. In Burkholderia glumae, the QS-dependent glyoxylate cycle plays an important role in cooperativity. A bifunctional QS-dependent transcriptional regulator, QsmR, rewired central metabolism to utilize the glyoxylate cycle rather than the tricarboxylic acid cycle. Defects in the glyoxylate cycle caused metabolic imbalance and triggered high expression of the stress-responsive chaperonin GroEL. High-level expression of GroEL in glyoxylate cycle mutants interfered with the biosynthesis of a public resource, oxalate, by physically interrupting the oxalate biosynthetic enzyme ObcA. Under such destabilized cooperativity conditions, spontaneous mutations in the qsmR gene in glyoxylate cycle mutants occurred to relieve metabolic stresses, but these mutants lost QsmR-mediated pleiotropy. Overcoming the metabolic restrictions imposed on the population of cooperators among glyoxylate cycle mutants resulted in the occurrence and selection of spontaneous qsmR mutants despite the loss of other important functions. These results provide insight into how QS bacteria have evolved to maintain stable cooperation via QS-mediated metabolic coordination.

IMPORTANCE
We address how quorum sensing (QS) has coevolved with metabolic networks to maintain bacterial sociality. We found that QS-mediated metabolic rewiring is critical for sustainable bacterial cooperation in Burkholderia glumae. The loss of the glyoxylate cycle triggered the expression of the stress-responsive molecular chaperonin GroEL. Excessive biosynthesis of GroEL physically hampered biosynthesis of a public good, oxalate. This is one good example of how molecular chaperones play critical roles in bacterial cooperation. In addition, we showed that metabolic restrictions in the glyoxylate cycle acted as a selection pressure on metabolic networks; there were spontaneous mutations in the qsmR gene to relieve such stresses. However, the presence of spontaneous qsmR mutants had tragic consequences for a cooperative population of B. glumae due to failure of qsmR-dependent activation of public good biosynthesis. These results provide a good example of a bacterial strategy for robust cooperation via QS-mediated metabolic rewiring.

The recognition of bacteria as social organisms has provided population and evolutionary biological perspectives on bacterial behavior. In Proteobacteria, acyl-homoserine lactone (AHL)-mediated quorum sensing (QS) controls social behaviors, including swarming motility, virulence, and biofilm formation (1–3). These social behaviors require cooperation, but the evolutionary stability of cooperation is precarious, because costly cooperative strategies are vulnerable to social cheating. Several studies...
have demonstrated that QS-dependent metabolism stabilizes cooperativity. In Pseudomonas aeruginosa, the production of such public resources is positively or negatively regulated in a QS-dependent manner to minimize production costs (3–5). The integration of metabolic information with QS stabilizes public good cooperation, as cells can cooperate only when they receive the appropriate nutritional resources (6).

Along with the known QS-regulatory mechanisms that support bacterial sociality, we explored whether metabolic evolution to sustain bacterial cooperativity is inherent in the structure of bacterial primary metabolic networks. To address this issue, Burkholderia glumae BGR1 was chosen as a model bacterium because of its dramatic QS-mediated metabolic fluctuations and metabolic plasticity (7, 8). In B. glumae, one LuxI-R-type QS system, TofI-R, generates N-octanoyl homoserine lactone (C8-HSL) as a major signaling molecule (9). The TofR and C8-HSL complex regulates expression of the qsmR gene, which encodes an IclR (isocitrate lyase regulator)-type transcriptional regulator (10). QsmR regulates various private and public resources, including oxalate in B. glumae (9–11). Burkholderia thailandensis E264, a nonpathogenic saprophyte, was used as a model system comparable to B. glumae BGR1. B. thailandensis contains three acyl-homoserine lactone QS circuits. The QS-1 signal (C8-HSL) is synthesized by BtaI1 complexes with BtaR1 to control aggregation, motility, and oxalate production (12). QS is essential for stationary-phase survival in both B. glumae and B. thailandensis (11). When these Burkholderia species use amino acids as a carbon source, deamination results in ammonia release, increasing the extracellular pH (11). Among QS-dependent public goods, oxalate is biosynthesized in the branched tricarboxylic acid (TCA) cycle to protect cells from ammonia-mediated alkaline toxicity during the stationary phase in a QsmR-dependent manner (Fig. 1) (11). The oxalate biosynthetic enzymes ObcA and ObcB use acetyl-coenzyme A (acetyl-CoA) and oxaloacetate as substrates (13, 14).

In this article, we report previously unknown functional roles of the glyoxylate cycle in bacterial cooperativity. The glyoxylate cycle allows an organism to bypass the CO2-generating steps in the TCA cycle (15). As a result, the glyoxylate cycle is essential for growth when acetate is the sole carbon source, as is the case for Escherichia coli (16). In addition to this known function of the glyoxylate cycle, we hypothesized that this

![QS-regulatory metabolic diagram for Burkholderia glumae](image-url)
cycle may play an important role in bacterial cooperativity. This hypothesis was based on previous QS-dependent transcriptome analyses of *B. glumae* that showed that the expression levels of two genes in the glyoxylate cycle, *aceA* and *glcB* (encoding isocitrate lyase and malate synthase, respectively), are QS dependent (11). We found that QsmR is bifunctional and rewires metabolic networks at the branch point of the glyoxylate and TCA cycles. Despite the presence of an intact QS system, a defective glyoxylate cycle generated metabolic stress and induced high expression of the stress-responsive chaperonin GroEL. The excessive levels of GroEL in the glyoxylate cycle mutants reduced the biosynthesis of oxalate by interrupting the oxalate biosynthetic enzyme ObcA. Metabolic imbalance in the glyoxylate cycle mutants exerted selection pressure, leading to spontaneous mutations in the *qsmR* gene. Such spontaneous mutations in the *qsmR* gene relieved the metabolic stress caused by the absence of a functional glyoxylate cycle, but they were accompanied by the loss of pleiotropy, including activation of *obcAB* gene expression, which subsequently disrupted cooperativity. From our results, we suggest that QS has evolved to be integrated into the central metabolism to maintain stable cooperation along with ensuring public good production.

**RESULTS**

**Metabolic rewiring by QS at the branch point of the glyoxylate and TCA cycles.**

On the basis of our previous QS-dependent transcriptome analysis, we predicted that central metabolism is QS dependent and plays an important role in *B. glumae* cooperativity. We confirmed that expression of *aceA* and *glcB* in the glyoxylate cycle is directly controlled by QsmR (Fig. 2A; see also Fig. S1A in the supplemental material). Isocitrate lyase activity was significantly higher in the wild-type strain than in the *tofI* mutant, BGS2, and the *qsmR* mutant, BGS9 (Fig. 2B). Exogenous addition of 1 µM C8-HSL to the BGS2 *tofI* mutant returned the isocitrate lyase activity to wild-type levels (Fig. 2B). Complementation strains of the *qsmR* and *aceA* mutants recovered full isocitrate lyase activity (Fig. 2B). In *B. thailandensis*, isocitrate lyase activity was also

---

**FIG 2** Rewiring of the metabolic networks by QsmR at the branch point of the glyoxylate and TCA cycles. (A) Expression of *aceA* and *glcB* genes, which encode isocitrate lyase and malate synthase, respectively, was activated by QsmR in *B. glumae*. (B) Isocitrate lyase activity was higher in the wild-type strain than in the QS mutants of *B. glumae*. (C) Isocitrate dehydrogenase (encoded by *icd1*) activity was higher in the *qsmR* mutant than in the wild-type strain of *B. glumae*. A mutation in the *aceK* gene, which encodes isocitrate dehydrogenase kinase/phosphatase, had no effect on isocitrate dehydrogenase activity in *B. glumae*. Error bars represent the error ranges of experiments performed in triplicate. ns, no significant difference; *, *P* < 0.05; **, *P* < 0.001.
The isocitrate lyase-deficient mutant was less virulent in rice panicles than the wild-type strain. (A) The wild-type strain produced severe symptoms in the panicles, including discoloration and empty heads of grain. Panicles inoculated with the isocitrate lyase-deficient mutant displayed milder symptoms. SDW, sterile distilled water. (B and C) Disease degree (B) and index (C) were evaluated as previously described (Materials and Methods). Pathogenicity assays were repeated three times in triplicate.

significantly higher in the wild-type E264 strain than in the bta1 mutant, JBT101, and the qsmR mutant, BT09539 (Fig. S1B). However, expression of the icd gene, which encodes isocitrate dehydrogenase (IDH), was controlled directly by QsmR, but isocitrate dehydrogenase activity was significantly higher in the qsmR mutants than in the wild-type strains of \( \text{B. glumae} \) and \( \text{B. thailandensis} \) (Fig. 2C; see also Fig. S1A and C). These results indicate that QsmR rewires metabolic networks to utilize the glyoxylate cycle in preference to the TCA cycle at the branch point of these two metabolic cycles in both bacterial species.

**Role of the QS-dependent glyoxylate cycle in \( \text{B. glumae} \) virulence.** Since defects in the glyoxylate cycle affected cooperativity in \( \text{B. glumae} \), we examined whether the QS-dependent glyoxylate cycle plays an important role in \( \text{B. glumae} \) virulence. Following the inoculation of rice panicles (in the flowing stage) with the wild-type strain and the aceA mutant, we observed a disease index for the aceA mutant that was 30% lower than that for the wild-type strain (Fig. 3). Our results indicated that the glyoxylate cycle-deficient mutant was less virulent than the wild-type strain in rice panicles. Inoculation with the complementation aceA strain recovered the disease index to wild-type levels. These results indicate that the QS-dependent glyoxylate cycle is important for full \( \text{B. glumae} \) virulence in rice panicles.

**Defects in the glyoxylate cycle affect cooperative behaviors.** To evaluate the biological significance of such metabolic rewiring in \( \text{B. glumae} \) and \( \text{B. thailandensis} \), we investigated whether the glyoxylate cycle plays a role in sustaining cooperativity. Growth, extracellular pH, ammonia accumulation, and oxalate production of the aceA mutants of \( \text{B. glumae} \) and \( \text{B. thailandensis} \) were monitored for 7 days in LB medium to identify any differences in known cooperative behaviors. Growth of the aceA mutants was comparable to that of the wild-type strains until the stationary phase was reached. Viable cell counts of the aceA mutants of \( \text{B. glumae} \) and \( \text{B. thailandensis} \) were decreased by approximately 10% compared to those of the wild-type population 1 to 3 days after subculture (Fig. 4; see also Fig. S2A). Extracellular pH of the aceA mutants of \( \text{B. glumae} \) and \( \text{B. thailandensis} \) remained at 7.6 to 8.2 throughout the growth period 1 day after subculture, whereas the wild-type strain exhibited slight alkalization followed by acidification and neutralization patterns (Fig. 4; see also Fig. S2A). Oxalate production by the aceA mutants of \( \text{B. glumae} \) and \( \text{B. thailandensis} \) was approximately 40 to 60% that of wild-type strain, which suggests a weak but continuous alkaline extracellular pH of the mutants (Fig. 4; see also Fig. S2A). These results indicate that the glyoxylate cycle plays pivotal roles in the persistence of cooperativity in \( \text{B. glumae} \) and \( \text{B. thailandensis} \) by influencing the biosynthesis of a public good, oxalate.

To determine the cross-activities of isocitrate lyase and methylisocitrate lyase encoded by the prpB gene in \( \text{B. glumae} \), levels of oxalate biosynthesis were measured in the prpB mutant and the aceA and prpB double mutant of \( \text{B. glumae} \). The aceA and prpB double mutant produced less oxalate than the aceA mutant; however, no significant
difference was found between the prpB mutant and the wild-type strain (see Fig. S2B). These results indicate that isocitrate lyase and methylisocitrate lyase have some cross-activities but that methylisocitrate lyase alone does not influence oxalate biosynthesis significantly in *B. glumae*.

**Interruption of ObcA by GroEL.** To identify the molecular mechanisms underlying the significant reduction in oxalate biosynthesis in the *B. glumae* aceA mutant, we measured the transcriptional and translational levels of *obcA* in the wild-type strain and aceA mutant. No difference between the wild-type strain and the aceA mutant was observed in the transcriptional or translational levels of *obcA* (Fig. 5B; see also Fig. S3). However, oxalate biosynthetic activity within the cells decreased by approximately 50% in the *B. glumae* aceA mutant strain compared to the wild-type strain (Fig. 5A). This was in agreement with the secreted oxalate concentration and extracellular pH. To determine the mechanisms involved in decreasing oxalate biosynthetic activity in the aceA

**FIG 4** Cell viability, extracellular pH, and ammonia and oxalate production of the aceA mutants of *B. glumae*. Viable cell numbers of the aceA mutant of *B. glumae* were decreased by approximately 10% compared to that of wild-type strain, and the extracellular pH of the aceA mutant was maintained at 7.6 to 7.9 throughout growth, whereas the wild-type strain showed acidification and neutralization patterns. Ammonia production by the aceA mutant was slightly lower than that by the wild-type strain, and oxalate production by the aceA mutant was ~60% that of wild-type strain.

**FIG 5** Physical interruption of the oxalate biosynthetic enzyme ObcA by metabolic stress-responsive GroEL in the aceA mutant of *B. glumae*. (A) The aceA mutant had approximately 50% of the oxalate biosynthetic activity of the wild-type strain. (B) Affinity pulldown experiments using anti-ObcA and anti-GroEL antibodies confirmed that GroEL interacts with ObcA, and the elevated levels of GroEL in the aceA mutant resulted in greater trapping of ObcA than was seen with the wild-type strain. Relative ObcA values were obtained by comparison with the wild-type values at two different time points as described previously (28). Ma, W, M, C, WB, and IP denote molecular markers, wild-type strain, aceA mutant, the complemented strain, Western blotting, and immunoprecipitation, respectively. The image of the SDS-PAGE gel stained with Coomassie brilliant blue R-250 (Bio-Rad) shows that samples were loaded equally in all lanes. (C) ObcA activity was decreased by approximately 4%, 52%, and 68% by addition of 500, 1,000, and 1,500 nM GroEL, respectively, compared to the control without GroEL *in vitro*. ObcA activity was not significantly affected by addition of 500, 1,000, and 1,500 nM BSA. Error bars represent the error ranges of experiments performed in triplicate. ns, no significant difference; *, P < 0.05; **, P < 0.001.
mutant, we performed a transcriptome sequencing (RNAseq) comparison of the wild-type strain and the aceA mutant strain. We found that expression of genes encoding stress-responsive chaperones or chaperonins, such as GroEL, was approximately 30-fold higher in the aceA mutant than in the wild-type strain (Table 1). The expression level of groEL was significantly higher in the aceA mutant than in the wild-type strain at both the transcriptional and translational levels (see Fig. S4). These results prompted us to hypothesize that highly expressed GroEL in the aceA mutant interacts with ObcA and interferes with its activity. GroEL protein was pulled down using an anti-ObcA antibody in affinity pulldown experiments in both the wild-type strain and aceA mutant (Fig. 5B), which indicated that ObcA and GroEL interact. Interestingly, even though the same amounts of ObcA were pulled down from all samples, the amount of ObcA that interacted with GroEL was higher in the aceA mutant than in the wild-type strain (Fig. 5B). The higher level of GroEL in the aceA mutant relative to the wild-type strain was in accordance with the RNAseq transcriptomic data, which showed a higher level of GroEL in the aceA mutant than in the wild-type strain (see Fig. S4). To demonstrate that the interaction between GroEL and ObcA inhibits the enzymatic activity of ObcA, we performed an ObcA activity assay in vitro. Addition of 500, 1,000, and 1,500 nM GroEL reduced ObcA activity by approximately 4%, 52%, and 68%, respectively, compared to the control without GroEL (Fig. 5C). ObcA activity was not affected by addition of the nonspecific protein bovine serum albumin (BSA) (Fig. 5C). These results suggested that the reduced activity of ObcA in the aceA mutant was due to physical interruption of ObcA by excess GroEL.

**Metabolic stress in the aceA mutant.** Based on the differential expression of multiple genes encoding chaperones and chaperonins in the aceA mutant of B. glumae, we hypothesized that the mutant may experience intracellular physiological stress. Since the aceA mutation blocks the glyoxylate cycle, we assumed that either isocitrate dehydrogenase (IDH) activity is increased to mitigate metabolic stress in the aceA mutant or isocitrate, aconitate, and citrate accumulation is greater in the mutant than in the wild-type strain. IDH activity in the aceA mutant was not significantly different from that of the wild-type strain (Fig. 2C). However, concentrations of isocitrate, aconitate, and citrate were significantly higher in the aceA mutant than in the wild-type strain (Fig. 5B). Consistent with these results, both the transcriptional level of the citA gene, which encodes citrate synthase, and its enzymatic activity were higher in the aceA mutant than in the wild-type strain (Fig. 6B; Table 1). These results indicate that the aceA mutant experiences physiological stresses due to metabolic imbalance, which is supportive of the concept of elevated expression of stress-responsive chaperones or chaperonins, including GroEL.

**Emergence of spontaneous mutations in the qsmR gene in glyoxylate cycle-defective strains.** Such continuous intracellular metabolic imbalance may not be endurable by mutants with a nonfunctional glyoxylate cycle. Since we observed morphologically distinct colonies when glyoxylate cycle-defective mutants were cultured in the preliminary experiment, colony morphology was evaluated at various culture time points. We collected samples from batch cultures in LB broth each day (days 0, 1, 2, 3, 4, 5, 6, and 7). The samples were serially diluted, spread on an LB agar

---

**TABLE 1** Expression of stress-responsive chaperone and chaperonin genes and the citA gene in the aceA mutant as analyzed by RNAseq

| Gene | Locus_ID | BGR1 (wild type) | BICL39 (aceA::EZ-Tn5) | BICL39C (aceA::EZ-Tn5/aceA) |
|------|----------|------------------|----------------------|--------------------------|
| groES | bglu_1g07140 | 657 | 23,268 | 5,352 |
| groEL | bglu_1g07150 | 893 | 26,979 | 8,264 |
| grpE | bglu_1g06330 | 75 | 359 | 95 |
| dnaK | bglu_1g06340 | 162 | 1,592 | 329 |
| dnaJ | bglu_1g06350 | 61 | 419 | 106 |
| citA | bglu_2g08280 | 736 | 3,314 | 853 |
plate, and then incubated for an additional day. When colonies were observed under a dissecting microscope, no morphologically distinct mutants were observed in samples collected before day 2 of batch culture in LB broth. However, after 3 to 4 days of batch culture in LB broth, morphologically distinct mutants of *B. glumae* appeared in the aceA mutant cultures (Fig. 7A; see also Fig. S5). The percentage of mutant colonies in the population increased to more than 90% after 7 days (Fig. 7B; see also Fig. S5). When 7-day-old cultures of the aceA mutant were subcultured in LB broth after dilution to approximately 1 × 10^7 CFU/ml and grown for 30 h, the extracellular pH reached 8.9,

---

**FIG 6** The aceA mutant experiences physiological stresses due to metabolic imbalance. (A) Concentrations of citrate, aconitate, and isocitrate were higher in the aceA mutant than in the wild-type strain. (B) Citrate synthase activity was higher in the aceA mutant than in the wild-type strain. Error bars represent the error ranges of experiments performed in triplicate. ns, no significant difference; *, P < 0.1; **, P < 0.05.

---

**FIG 7** Morphological appearance of spontaneous glyoxylate cycle mutants. (A) Colonies of the spontaneous qsmR mutants were morphologically distinct from those of qsmR-positive (qsmR^+/+) cells. Colonies were observed under a dissecting microscope at ×30 magnification. (B) The qsmR mutants appeared at day 4 after subculture and reached approximately 90% of the total viable population on day 7. (C) Among the 10 qsmR mutants characterized, 3 had an insertion of IS1418 at the 5’ end of the gene, and 4 and 3 had 112-bp and 780-bp internal deletions, respectively. Fractions at the right end indicate the frequencies of each type of mutant among the total 10 mutants. IS, SD, and LD indicate insertion, small deletion, and large deletion, respectively. (D) Isocitrate dehydrogenase activity in three types (IS, SD, and LD) of spontaneous qsmR mutants was higher than in the wild-type strain or the aceA mutant. (E) Fitness experiments comparing the wild-type strain and the spontaneous qsmR mutant_LD strain. The spontaneous qsmR mutant LD strain did not benefit from the presence of the wild type. Error bars represent the error ranges of experiments performed in triplicate. ns, no significant difference; *, P < 0.05.
and no survivors were detected (see Fig. S6). To characterize such spontaneously occurring mutants, two mutants that were siblings were subjected to whole-genome resequencing. We found that IS1418, an insertion sequence (IS) 865 bp in length that carries 15-bp inverted repeats with a target duplication of 3 bp (17), was inserted in the qsmR gene in each mutant. On the basis of whole-genome resequencing data, we amplified the qsmR gene region of another 30 mutants. PCR products representing three different mutation types, insertion, small deletion, and large deletion, were observed (Fig. 7C). We selected three insertion mutants (qsmR_IS), four small-deletion mutations (qsmR_SD), and three large-deletion mutants (qsmR_LD) to determine the nature of the mutations by sequencing (Fig. 7C). These spontaneous qsmR mutants exhibited elevated IDH activities due to derepression of icd genes (Fig. 7D), which indicates that instinctive coping mechanisms to relieve metabolic stresses are operational in the glyoxylate cycle mutants. However, these mutants consequently failed to activate oxalate biosynthesis (see Fig. S7A). Thus, due to the toxic alkaline extracellular pH, the spontaneous qsmR mutants did not survive stationary phase after they were isolated, as previously observed in the qsmR mutant, BGS9 (see Fig. S7A) (11). To determine whether the spontaneous qsmR mutants are cheaters, we monitored the population density, pH, oxalate, and ammonia in mixed culture with the wild type. Both the wild type and the spontaneous mutants failed to survive in mixed cultures (Fig. 7E; see also Fig. S7B). This population collapse was due to the alkaline condition caused by ammonia accumulation by the wild type and the spontaneous mutants and also to the level of oxalate being insufficient to neutralize it in mixed culture (Fig. 7E; see also Fig. S7B).

DISCUSSION

Positive control of the glyoxylate cycle by QS allowed us to predict that the QS-dependent glyoxylate cycle may have an important role in sustaining cooperativity. In fact, it is known that the glyoxylate cycle is positively controlled by QS in B. cepacia, Yersinia pestis, and P. aeruginosa (18–20). Such upregulation of the glyoxylate cycle by QS provides advantages for growth on certain carbon resources (19, 20). However, the biological significance of positive control of the glyoxylate cycle by QS is unknown. In this study, we explored whether the QS-dependent glyoxylate cycle plays a role in bacterial sociality.

We first showed that QsmR rewires metabolic networks at the branch point of the glyoxylate cycle and TCA cycle in B. glumae. This indicated that QsmR functions as a transcriptional activator and a repressor, which is not unusual because transcriptional regulators belonging to the IclR type are often bifunctional (21). Such metabolic rewiring by bifunctional QsmR raised questions as to what the underlying reasons are and whether it is related to bacterial cooperativity. We chose B. thailandensis E264 as a system comparable to B. glumae, because it also exhibited QmsR-dependent oxalate biosynthesis (11) as well as an 80.53% identity with B. glumae BGR1 at the genome level, as calculated by average nucleotide identity (ANI) based on BLAST parameters (22). Concerns about the possibility that QsmR-mediated metabolic rewiring is a peculiar phenomenon in B. glumae were dismissed since expression of aceA, glcB, and icd genes in B. thailandensis exhibited the same gene regulation patterns as those observed in B. glumae. This phenomenon appears to be widely distributed among the members of the Burkholderia genus. Recent QS-dependent transcriptome analysis in B. thailandensis showed that expression of both the aceA and glcB genes is not controlled by QS (12). However, this may be due to different culture conditions. For QS-dependent transcriptome analysis, the strains of B. thailandensis were grown in LB supplemented with 50 mM MOPS (morpholinepropanesulfonic acid) (12) whereas expression of both the aceA and glcB gene was estimated in B. thailandensis cells grown in LB broth without any buffer in this study.

Since methylisocitrate lyase encoded by the prpB gene has cross-activity with isocitrate lyase in B. pseudomallei and Mycobacterium tuberculosis (23, 24), we expected a similar phenomenon in B. glumae. Cross-activity of methylisocitrate lyase and isocit-
rate lyase was detectable; however, the level of the cross-activity was not high enough to substitute isocitrate lyase with methylisocitrate lyase in the aceA mutant. Thus, the activity of methylisocitrate lyase has a negligible effect on oxalate biosynthesis in the wild-type strain.

Similarly to E. coli, the glyoxylate cycle in B. glumae is essential for growth on acetate (see Fig. S8 in the supplemental material). The glyoxylate cycle is important for the survival of pathogens inside the host and provides a physiological benefit to certain pathogenic microorganisms (19, 20, 24). A similar role of the glyoxylate cycle was observed in B. glumae since the aceA mutant was less virulent than the wild-type strain in rice panicles (Fig. 3). In addition to these biological roles of glyoxylate cycle, we were interested in the biological meaning of QS dependency in the glyoxylate cycle and in QsmR-mediated metabolic rewiring.

It was interesting that the glyoxylate cycle affects biosynthesis of a public resource, oxalate. We then explored why and how the glyoxylate cycle affects oxalate biosynthesis in B. glumae. Our systematic analyses of TCA cycle intermediates and the transcriptome of glyoxylate cycle-defective mutants led us to find that blocking of the glyoxylate cycle leads to stressful conditions and triggers high-level expression of GroEL. Considering that wild-type cells have normal metabolic conditions in the presence of normal concentrations of citrate, aconitate, and isocitrate, it was clear that the aceA mutant experiences stressful metabolic conditions due to significantly high concentrations of these metabolites. Elevated expression of stress-responsive chaperones strongly supported the concept that physiological conditions in the glyoxylate cycle defectives are indeed stressful. The idea of interaction of the oxalate biosynthetic enzyme ObcA with GroEL was supported by the results of affinity pulldown experiments in both the wild-type strain and aceA mutant. In general, GroEL is a chaperonin that facilitates the proper folding of proteins, but it has been reported that an excess of certain chaperones exerts a negative effect on protein production (25). Our current results do not provide evidence that GroEL is required for full activity of ObcA. Rather, the in vitro ObcA activity assay suggested that excess GroEL hampered ObcA activity, resulting in reduced oxalate production (Fig. 5C). These findings indicated that the metabolic flow through the glyoxylate cycle is essential for producing oxalate as a public good in a QS-dependent manner in B. glumae.

In glyoxylate cycle-defective mutants, metabolic stresses were exerted as a selection pressure on the metabolic networks, resulting in the occurrence of spontaneous mutations in the qsmR gene to derepress expression of icd genes. These spontaneous mutations in the qsmR gene had a direct effect on the biosynthesis of public goods (see Fig. S7A), which is quite different from nonsocial adaptation, in which an individual’s fitness would increase by derepressing transcriptional repressor PsdR, for example, in P. aeruginosa (26). Spontaneous mutations in the qsmR gene were not observed in the wild-type B. glumae strain under the same growth conditions as those of the aceA mutant. Occasionally, QS enhancement of extracellular stress responses, such as the oxidative stress response, acts as a counterselective force for the appearance and survival of QS cheaters. However, intracellular metabolic stress acts as a pressure on individual cells to develop genomic mutations to relieve such stress, even though these mutations lost all of their attributes under the control of the QS-dependent QsmR. Therefore, intracellular metabolic homeostasis is an important issue in Burkholderia cooperative cells. It might be assumed that spontaneous qsmR mutants in the glyoxylate cycle-defective B. glumae strains are cheaters. However, these mutants are not the cheaters, because three types (IS, SD, and LD) of spontaneous qsmR mutant did not benefit from mixed culture with the wild-type oxalate producer (Fig. 7E; see also Fig. S7B). Mutations that inactivate qsmR were the fastest means of rescuing the metabolic stress but did not benefit individual cells.

In B. glumae, the QS-dependent glyoxylate cycle functions to support bacterial cooperativity, as well as to provide a metabolic bypass. Our results reveal important physiological roles of the glyoxylate cycle in cooperative bacteria and suggest that the glyoxylate cycle may represent a target for the development of chemical agents to
control bacterial social behaviors, such as the virulence of pathogens. This may also be a good example of overcoming metabolic restrictions caused by blocking a critical metabolic flow in a population of cooperators.

MATERIALS AND METHODS

Bacterial strains and growth conditions. Bacterial strains and plasmids used in this study are listed in Table S1 in the supplemental material. Strains of B. glumae and B. thailandensis were grown in Luria-Bertani (LB) broth (Affymetrix, Santa Clara, CA) (0.1% tryptone, 0.5% yeast extract, and 0.5% NaCl [all wt/vol]) at 37°C.

Quantitative reverse transcription-PCR (qRT-PCR). Total RNAs from B. glumae BGR1, BGS2 (BGR1 tofi::Δ fragment), BGS9 (BGR1 qsmR::Δ fragment), and S9NCS (BGR1 qsmR::Δ fragment) grown in LB medium at 37°C for 10 h after subculture, were extracted using RNeasy minikits (Qiagen, Venlo, Netherlands), as described by the manufacturer. Total RNA was treated with RNase-free DNase I (Ambion, Waltham, MA) to remove genomic DNA. Total RNA (1 μg) was subjected to reverse transcription into cDNA using Moloney murine leukemia virus (MMLV) reverse transcriptase (Promega, Madison, WI) and incubation for 1 h at 42°C. Primer pairs used for qRT-PCR are listed in Table S2. The 16S RNA gene served as the positive control. Transcriptional levels were determined using SsoFast Eva Green SuperMix (Bio-Rad, Hercules, CA) and a CFX96 Real-Time PCR system (Bio-Rad). The thermal cycling parameters were as follows: 95°C for 30 s, followed by 40 cycles of 95°C for 5 s and 60°C for 5 s. All PCRs were performed in triplicate, and all data were normalized to the expression levels of the 16S RNA gene using Bio-Rad CFX Manager software.

Isocitrate lyase activity assay. Isocitrate lyase activity was measured as described previously (27). Briefly, isocitrate was converted to succinate and glyoxylate by isocitrate lyase. During a reaction with phenylhydrazine, glyoxylate formed phenylhydrazine-glyoxylate, and the absorbance of this compound was measured at 324 nm every 60 s for 5 min. Enzymatic activity (in units per milliliter) was calculated as follows: ΔA324nm/min blank)/millimolar extinction coefficient of phenylhydrazine-glyoxylate at 324 nm.

Isocitrate dehydrogenase activity assay. Isocitrate dehydrogenase activity was measured using an isocitrate dehydrogenase activity assay kit (Biovision, Milpitas, CA) according to the manufacturer’s instructions. We measured the absorbance at 450 nm every 1 min in a microplate reader (PerkinElmer, Waltham, MA) and used only the data obtained within the linear range to calculate isocitrate dehydrogenase activity using the NADH standard curve, as described by the manufacturer.

Measurement of secreted oxalate and ammonia. Secreted oxalate and ammonia levels were measured as described previously (11). Overexpression and purification of GroEL. The groEL gene of B. glumae was amplified using the primers listed in Table S2. The amplified product was cloned into NdeI and HindIII restriction sites of a pET28b expression vector (Merck, Darmstadt, Germany), resulting in pET28b-groEL (see Table S1). His-GroEL was overexpressed in Escherichia coli BL21 (DE3), which was induced by adding 0.5 mM isopropyl β-D-thiogalactopyranoside, followed by additional growth for 14 h at 20°C. His-GroEL was purified in a buffer A, which contained 50 mM Tris-Cl (pH 8.0), 100 mM NaCl, and 5% (vol/vol) glycerol, using an immobilized metal affinity column (GE Healthcare, Chicago, IL) equilibrated with buffer A and then eluted with buffer A containing 500 mM imidazole. Purified His-GroEL was dialyzed against buffer A overnight at 4°C.

Oxalate biosynthetic activity assay. His-ObcA and His-Obc1* were purified, and 50 nM His-ObcA and 800 nM His-Obc1* were used for oxalate biosynthetic activity assay, as described previously (14). Oxalate biosynthetic activity and oxalate levels were measured as described previously (11, 13, 14).

Commmunoprecipitation (Co-IP) and Western blot analysis. Co-IP was performed using a Pierce Direct IP kit (Thermo Fisher Scientific, Waltham, MA) according to the manufacturer’s instructions. An anti-ObcA antibody (AbFrontier, Seoul, South Korea) (2 μg) was coupled to AminoLink Plus coupling resin (Thermo Fisher Scientific). Cell lysates from B. glumae BGR1, BICL39 (BGR1 aceA::EZ-Tn5), and BICL39C (BGR1 aceA::EZ-Tn5 aceA) containing 1 mg of protein were added to the anti-ObcA-coupled resin in a spin column and mixed at 4°C for 24 h. Immunoprecipitates were eluted, followed by the addition of 5× nonreducing lane marker sample buffer. Samples were boiled for 5 min, cooled to room temperature, and separated by sodium dodecyl sulfate-polyacylamide gel electrophoresis (SDS-PAGE). Samples were then transferred to nitrocellulose membranes. An anti-GroEL antibody (Abcam, Inc., Cambridge, United Kingdom) was used to detect GroEL protein interacting with ObcA protein. The immunoreactive bands were detected using ECL reagents (Bio-Rad) and captured by ChemiDoc XRS+ (Bio-Rad). The number of pixels per inch of each band was measured using Image Lab version 2.0.1 software (Bio-Rad). For Western blot analysis, the number of pixels measured in the wild-type strain at two different time points was set to 1, and the values for the aceA mutant and complemented strains were normalized to this value to allow ratio comparisons. For IP and Western blot analyses, the IP readings were divided by the input readings to obtain ratios (28).

Quantitative analysis of citrate, aconitate, and isocitrate. Levels of citrate, aconitate, and isocitrate in B. glumae BGR1, BICL39 (BGR1 aceA::EZ-Tn5), and BICL39C (BGR1 aceA::EZ-Tn5 aceA) were analyzed by capillary electrophoresis time of flight mass spectrometry, as described previously (7, 29, 30). All experiments were performed using at least three independent replicates.

Citrte synthase activity assay. Citrate synthase activity was measured using a citrate synthase assay kit (Sigma-Aldrich, St. Louis, MO, USA) and a multilabel plate reader (PerkinElmer) according to the instructions of the manufacturers.
RNAseq analysis. Total RNAs from B. glumae BGR1, BICL39 (BGR1 aceA::EZ-Tn5), and BICL39C (BGR1 aceA::EZ-Tn5/aceA), grown in LB medium at 37°C for 12 h after subculture, were extracted using RNAasy mini kits (Qiagen) following the manufacturer's protocols. Extracted total RNA was treated with RNase-free DNase I (Ambion) to remove DNA. The quantity and quality of the total RNA were evaluated using RNA electropherograms (Agilent 2100 Bioanalyzer, Santa Clara, CA) and by assessing the RNA integrity number (RIN). An 8-μg volume of total RNA from each sample with a RIN value greater than 8.0 was used as the starting material and was treated with a MICROBExpress mRNA enrichment kit (Invitrogen, Carlsbad, CA). The resulting mRNA samples were processed for the sequencing libraries using an Illumina mRNAseq sample preparation kit (Illumina, San Diego, CA) following the manufacturer's protocols. One lane per sample was used for sequencing by Illumina Genome Analyzer Ilx (Illumina) to generate nondirectional, single-ended, 36-bp reads. Quality-filtered reads were mapped to the B. glumae BGR1 genome sequence (NCBI RefSeq assembly accession no. GCF_000022645.2; https://www.ncbi.nlm.nih.gov/assembly/GCF_000022645.2/) using the BWA package (31). The mRNA reads were normalized to reads per kilobase per million (RPKM) mapped reads (32).

Identification of the mutations in spontaneous mutant strains. Chromosomal DNA was isolated from 10 spontaneous mutants cultured in LB broth using a method previously published by Sambrook et al. (33). To identify the mutation positions, 2 of the 10 mutants (BICL39 LY and BICL39 C) were subjected to whole-genome resequencing. Quantitation and quality assessment of the 2 samples were carried out using PicoGreen (Invitrogen) on a Victor 3 fluorometer (PerkinElmer). Illumina shotgun libraries from gDNA were prepared and sequenced on one lane of an Illumina HiSeq 2500 sequencer. Resequenced Illumina HiSeq data were trimmed using Trimmomatic software (34) and aligned to the B. glumae BGR1 genome using the BWA package (31). DNA variants, including single nucleotide polymorphisms, insertions, and deletions, between libraries were detected using SAMtools (35, 36) and FreeBayes (37). Subclonal single nucleotide variants were detected using the deepSNV package in R (38, 39). Gene mutation candidates were visualized and validated using Integrative Genomics Viewer (40). On the basis of whole-genome resequencing data, we amplified the qsmR gene region of another 30 mutants using the qsmR-F and qsmR-R primers (see Table S2). PCR products representing three different mutation types, insertions, small deletions, and large deletions, were detected. We selected three insertion mutants, four small-deletion mutants, and three large-deletion mutants to confirm the position of the mutation by sequencing. The amplified 1,791-bp fragment from 10 mutants was cloned into the pBlueScript II SK(+) vector prior to sequencing to identify the mutation sites and complete sequencing.

Plant inoculation. In a greenhouse, rice plants (Oryza sativa cv. Milyang 23) were inoculated at the flowering stage with approximately 1 × 10⁶ CFU/ml of B. glumae. Disease symptoms in the rice plants were evaluated on day 7 after inoculation. The disease index was determined as described previously (41), using the following scale: 0 = healthy panicle, 1 = panicle 0 to 20% discolored, 2 = panicle 20 to 40% discolored, 3 = panicle 40 to 60% discolored, 4 = panicle 60 to 80% discolored, and 5 = panicle 80 to 100% discolored (disease index = Σ (number of samples per score × score)/the total number of panicles). Pathogenicity assays were repeated three times with three replications.

Electrophoretic mobility shift assay. QsmR-His was purified as described previously (10). The upstream regions of putative QsmR target genes were amplified using the primers listed in Table S2. PCR products were labeled with biotin using a LightShift chemiluminescent electrophoretic mobility shift assay kit (Thermofisher Scientific), according to the manufacturer’s protocol. We used a 329-bp sequence upstream of katE as competitor DNA. This DNA fragment was amplified using primers KatE1-F and KatE1-R (see Table S2). Purified QsmR-His (10 nM or 250 nM) was incubated in binding buffer (10 mM Tris-HCl (pH 7.5), 100 mM NaCl, 0.5 mM EDTA, 1 mM dithiothreitol [DTT], 5% (vol/vol) glycerol, and 10 ng μl⁻¹ poly(dI-dC)] containing 1 nM or 2 nM biotin-labeled DNA for 15 min at 28°C. In competition analyses, unlabeled target DNA at 10-fold to 15-fold molar excess was added to each reaction mixture along with the extract before the addition of labeled DNA targets. Reaction mixtures were separated on nondenaturing 4% (wt/vol) gels and transferred to nitrocellulose membranes, followed by detection of relevant bands using streptavidin/horseradish peroxidase-derived chemiluminescence kits, as described by the manufacturer. Images were visualized using ChemiDoc XRS+ and Image Lab Software (Bio-Rad).

Construction of B. thailandensis aceA::lacZY and aceA::lacZY/aceA. The aceA (BTH_11998) gene was deleted by integration of a suicide vector, as described previously (42). Briefly, a 250-bp DNA fragment encoding only the 5’ end of the aceA gene was generated by PCR using BTaceA-F and BTaceA-R primers, cloned into pBluescript II SK(+) and termed “pBTACE1.” A 250-bp DNA fragment obtained from pBTACE1 by digestion performed with EcoRI and XbaI was ligated into a suicide vector containing lacZY genes and pVJK112 to produce pBTACE2 (see Tables S1 and S2) (42). We mutated pBTACE2 by the use of EZ-Tn5(Tp) to replace kanamycin resistance with trimethoprim (Tp) resistance. pBTACE2_Tp was mobilized from DH5α (λpir) into B. thailandensis E264 by conjugation using PRK2013 as a helper plasmid. The integrated mutant was confirmed by PCR using LacFuse and BTaceA-R primers (see Table S2). To complement the aceA mutant of B. thailandensis, we used the Tn7 site-specific transposition machinery system (9) to insert a single copy of aceA downstream of glmS.

Generation of S9NC5 and BICL39C. For genetic complementation of qsmR mutant BGS9, we mutagenized the mutant via the use of mini-TnSrescue (9) to integrate qsmR into the genome as a single copy. We confirmed that a single copy of the qsmR gene was inserted into the intergenic region between bglu_1g10240 and bglu_1g10250 in chromosome 1 of the qsmR mutant BGS9, resulting in strain S9NC5 (BGR1 qsmR::Tn::qsmR). Likewise, the aceA gene was inserted into the intergenic region between bglu_2g21870 and bglu_2g21880 in chromosome 2 of the aceA mutant BICL39, resulting in strain BICL39C (BGR1 aceA::EZ-Tn5/aceA).
**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.00042-17.

**FIG S1**, PDF file, 0.9 MB.
**FIG S2**, PDF file, 0.4 MB.
**FIG S3**, PDF file, 0.7 MB.
**FIG S4**, PDF file, 0.5 MB.
**FIG S5**, PDF file, 0.9 MB.
**FIG S6**, PDF file, 0.3 MB.
**FIG S7**, PDF file, 0.4 MB.
**FIG S8**, PDF file, 0.4 MB.
TABLE S1, DOCX file, 0.02 MB.
TABLE S2, DOCX file, 0.03 MB.
TABLE S3, DOCX file, 0.01 MB.
TABLE S4, DOCX file, 0.01 MB.
TABLE S5, DOCX file, 0.01 MB.
TABLE S6, DOCX file, 0.01 MB.
TABLE S7, DOCX file, 0.01 MB.
TABLE S8, DOCX file, 0.01 MB.
TABLE S9, DOCX file, 0.01 MB.

**REFERENCES**

1. Fuqua C, Greenberg EP. 2002. Listening in on bacteria: acyl-homoserine lactone signalling. Nat Rev Mol Cell Biol 3:685–695. https://doi.org/10.1038/nrm907.
2. Ng WL, Bassler BL. 2009. Bacterial quorum-sensing network architectures. Annu Rev Genet 43:197–222. https://doi.org/10.1146/annurev-genet-102108-134304.
3. Schuster M, Sexton DJ, Diggle SP, Greenberg EP. 2013. Acyl-homoserine lactone quorum sensing: from evolution to application. Annu Rev Microbiol 67:43–63. https://doi.org/10.1146/annurev-micro-092412-155635.
4. Pai A, Tanouchi Y, You L. 2012. Optimality and robustness in quorum sensing (Qs)-mediated regulation of a costly public good enzyme. Proc Natl Acad Sci U S A 109:19810–19815. https://doi.org/10.1073/pnas.1211072109.
5. Gupta R, Schuster M. 2013. Negative regulation of bacterial quorum sensing tunes public goods cooperation. ISME J 7:2159–2168. https://doi.org/10.1038/ismej.2013.109.
6. Boyle KE, Monaco H, van Ditmarsch D, Deforet M, Xavier JB. 2015. Integration of metabolic and quorum sensing signals governing the decision to cooperate in a bacterial social trait. PLoS Comput Biol 11:e1004279. https://doi.org/10.1371/journal.pcbi.1004279.
7. An JH, Goo E, Kim H, Seo YS, Hwang I. 2014. Bacterial quorum sensing and metabolic slowing in a cooperative population. Proc Natl Acad Sci U S A 110:19980–19985. https://doi.org/10.1073/pnas.1211072109.
8. Goo E, An JH, Kang Y, Hwang I. 2015. Control of bacterial metabolism by quorum sensing. Trends Microbiol 23:567–576. https://doi.org/10.1016/j.tim.2015.05.007.
9. Kim J, Kim JG, Kang Y, Jang JH, Kim JY, Lim JY, Kim S, Suga H, Nagamatsu T, Hwang I. 2004. Quorum sensing and the LysR-type transcriptional activator ToxR regulate toxoflavin biosynthesis and transport in Burkholderia glumae. Mol Microbiol 54:921–934. https://doi.org/10.1111/j.1365-2958.2004.03438.x.
10. Kim J, Kang Y, Choi O, Jeong Y, Jeong JE, Lim JY, Kim M, Moon JS, Suga H, Hwang I. 2007. Regulation of polar flagellar genes is mediated by quorum sensing and FhICD in Burkholderia glumae. Mol Microbiol 64: 165–179. https://doi.org/10.1111/j.1365-2958.2007.05646.x.
11. Goo E, Majerczyk CD, An JH, Chandler JR, Seo YS, Ham H, Lim JY, Kim H, Lee B, Jang MS, Greenberg EP, Hwang I. 2012. Bacterial quorum sensing, cooperativity, and anticipation of stationary-phase stress. Proc Natl Acad Sci U S A 109:19775–19780. https://doi.org/10.1073/pnas.1218092109.
12. Majerczyk CD, Brittnacher M, Jacobs M, Armour CD, Radey M, Schneider E, Phattarasokul S, Bunt R, Greenberg EP. 2014. Global analysis of the Burkholderia thailandensis quorum sensing-controlled regulon. J Bacteriol 196:1412–1424. https://doi.org/10.1128/JB.00405-13.
13. Li HQ, Matsuda I, Fujise Y, Ichiyama A. 1999. Short-chain acyl-CoA-dependent production of oxalate from oxaloacetate by Burkholderia glumae, a plant pathogen which causes grain rot and seedling rot of rice via oxalate production. J Biochem 126:243–253. https://doi.org/10.1093/oxfordjournals.jbchem.a022429.
14. Oh J, Goo E, Hwang I, Rhee S. 2014. Structural basis for bacterial quorum sensing-mediated oxalogenesis. J Biol Chem 289:11465–11475. https://doi.org/10.1074/jbc.M113.543462.
15. Cronan JE, Jr, Laporte D. 2005. Tricarboxylic acid cycle and glyoxylate bypass. Ecosal Plus 1 https://doi.org/10.1128/ecosalplus.3.5.2.
16. Cozzone AJ. 1998. Regulation of acetate metabolism by protein phosphorylation in enteric bacteria. Annu Rev Microbiol 52:127–164. https://doi.org/10.1146/annurev.micro.52.1.127.
17. Hasebe A, Iida S. 2000. The novel insertion sequences IS1417, IS1418, and IS1419 from Burkholderia glumae for hydrogen cyanide production in a cystic fibrosis isolate of Pseudomonas aeruginosa. J Bacteriol 191:6335–6339. https://doi.org/10.1128/JB.00692-09.
18. Aguilar C, Friscina A, Devescovi G, Kojic M, Venturi V. 2003. Identification of quorum-sensing-regulated genes of Burkholderia cepacia. J Bacteriol 185:6456–6462. https://doi.org/10.1128/JB.185.21.6456-6462.2003.
19. Hagins JM, Locy R, Silo-Suh L. 2009. Isoctarate lyase supplies precursors for hydrogen cyanide production in a cystic fibrosis isolate of Pseudomonas aeruginosa. J Bacteriol 191:6335–6339. https://doi.org/10.1128/JB.00692-09.
20. LaRock CN, Yu J, Horswill AR, Parsek MR, Minion FC. 2013. Transcriptome...
Metabolic Rewiring for Bacterial Cooperativity

23. van Schaik EJ, Tom M, Woods DE. 2009. *Burkholderia pseudomallei* isocitrate lyase is a persistence factor in pulmonary melioidosis: implications for the development of isocitrate lyase inhibitors as novel antimicrobials. Infect Immun 77:4275–4283. https://doi.org/10.1128/IAI.00609-09.

24. Muñoz-Elías JL, Janda AM, Cherian J, McKinney JD. 2006. Role of the methylcitrate cycle in *Mycobacterium tuberculosis* metabolism, intracellular growth, and virulence. Mol Microbiol 60:1109–1122. https://doi.org/10.1111/j.1365-2958.2006.05155.x.

25. Martínez-Alonso M, García-Fruitós E, Ferrer-Miralles N, Rinas U, Villaverde A. 2010. Side effects of chaperone gene co-expression in recombinant protein production. Microb Cell Fact 9:64. https://doi.org/10.1186/1475-2859-9-64.

26. Asfahl KL, Walsh J, Gilbert K, Schuster M. 2015. Non-social adaptation defers a tragedy of the commons in *Pseudomonas aeruginosa* quorum sensing. ISME J 9:1734–1746. https://doi.org/10.1038/ismej.2014.259.

27. Chell RM, Sundaram TK, Wilkinson AE. 1978. Isolation and characterization of isocitrate lyase from a thermophilic *Bacillus* sp. Biochem J 173:165–177. https://doi.org/10.1042/bj1730165.

28. Slanina H, Hebling S, Hauck CR, Schubert-Unkmeir A. 2012. Cell invasion by *Neisseria meningitidis* requires a functional interplay between the focal adhesion kinase, Src and cortactin. PLoS One 7:e39613. https://doi.org/10.1371/journal.pone.0039613.

29. Soga T, Ueno Y, Naraoka H, Tomita M, Nishioka T. 2003. Quantitative metabolite analysis using capillary electrophoresis mass spectrometry. J Proteome Res 2:488–494. https://doi.org/10.1021/pr034020m.

30. Ishii N, Nakahigashi K, Baba T, Robert M, Soga T, Kanai A, Hirasawa T, Naba M, Hira K, Hoque A, Ho PY, Kakazu Y, Sugawara K, Igarashi S, Masuda T, Sugiyama N, Togashi T, Hasegawa M, Takai Y, Yugi K, Arakawa K, Iwata N, Toya Y, Nakayama Y, Nishioka T, Shimizu K, Mori H, Tomita M. 2007. Multiple high-throughput analyses monitor the response of *E. coli* to perturbations. Science 316:593–597. https://doi.org/10.1126/science.1132067.

31. Li H, Durbin R. 2010. Fast and accurate long-read alignment with Burrows-Wheeler transform. Bioinformatics 26:589–595. https://doi.org/10.1093/bioinformatics/btp698.

32. Mortazavi A, Williams BA, McCue K, Schaeffer L, Wold B. 2008. Mapping and quantifying mammalian transcriptomes by RNA-Seq. Nat Methods 5:628–632. https://doi.org/10.1038/nmeth.1226.

33. Sambrook J, Fritsch EF, Maniatis T. 1989. Molecular cloning: a laboratory manual. 2nd ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.

34. Bolger AM, Lohse M, Usadel B. 2014. Trimomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 30:2114–2120. https://doi.org/10.1093/bioinformatics/btu170.

35. Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R. 1000 Genome Project Data Processing Subgroup. 2009. The sequence alignment/map format and SAMtools. Bioinformatics 25:2078–2079. https://doi.org/10.1093/bioinformatics/btp352.

36. Li H. 2011. A statistical framework for SNP calling, mutation discovery, association mapping and population genetical parameter estimation from sequencing data. Bioinformatics 27:2987–2993. https://doi.org/10.1093/bioinformatics/btr509.

37. Garrison E, Marth G. 2012. Haplotype-based variant detection from short-read sequencing. arXiv arXiv:1207.3907 [q-bio.GN].

38. Gerstung M, Beisel C, Rechsteiner M, Wild P, Schraml P, Moch H, Beer-weninkel N. 2012. Reliable detection of subclonal single-nucleotide variants in tumour cell populations. Nat Commun 3:811. https://doi.org/10.1038/ncomms1814.

39. Gerstung M, Papaemmanuil E, Campbell PJ. 2014. Subclonal variant calling with multiple samples and prior knowledge. Bioinformatics 30:1198–1204. https://doi.org/10.1093/bioinformatics/btt750.

40. Robinson JT, Thorvaldsdóttir H, Winckler W, Guttman M, Lander ES, Getz G, Mesirov JP. 2011. Integrative genomics viewer. Nat Biotechnol 29:1288–1295. https://doi.org/10.1038/nbt.2076.

41. Iiyama K, Furuya N, Takanami Y, Matsuyama N. 1995. A role of phytotoxin in virulence of *Pseudomonas glumae* Kurita et Tabei. Jpn J Phytopathol 61:470–476. https://doi.org/10.3186/jjphytopath.61.470.

42. Kalogeraki VS, Winans SC. 1997. Suicide plasmids containing promoter-less reporter genes can simultaneously disrupt and create fusions to target genes of diverse bacteria. Gene 188:69–75. https://doi.org/10.1016/S0378-1119(96)00778-0.