Imbalance in Carbon and Nitrogen Metabolism in *Comamonas testosteroni* R2 Is Caused by Negative Feedback and Rescued by L-arginine

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The collapse of *Comamonas testosteroni* R2 under chemostat conditions and the aerobic growth of strain R2 under batch conditions with phenol as the sole carbon source were investigated using physiological and transcriptomic techniques. Phenol-/catechol-degrading activities under chemostat conditions gradually decreased, suggesting that metabolites produced from strain R2 accumulated in the culture, which caused negative feedback. The competitive inhibition of phenol hydroxylase and catechol dioxygenase was observed in a crude extract of the supernatant collected from the collapsed culture. Transcriptomic analyses showed that genes related to nitrogen transport were up-regulated; the ammonium transporter *amtB* was up-regulated approximately 190-fold in the collapsed status, suggesting an increase in the concentration of ammonium in cells. The transcriptional levels of most of the genes related to gluconeogenesis, glycolysis, the pentose phosphate pathway, and the TCA and urea cycles decreased by ~0.7-fold in the stable status, whereas the activities of glutamate synthase and glutamine synthetase increased by ~2-fold. These results suggest that ammonium was assimilated into glutamate and glutamine via 2-oxoglutarate under the limited supply of carbon skeletons, whereas the synthesis of other amino acids and nucleotides was repressed by 0.6-fold. Furthermore, negative feedback appeared to cause an imbalance between carbon and nitrogen metabolism, resulting in collapse. The effects of amino acids on negative feedback were investigated. L-arginine allowed strain R2 to grow normally, even under growth-inhibiting conditions, suggesting that the imbalance was corrected by the stimulation of the urea cycle, resulting in the rescue of strain R2.

**Key words:** metabolism, negative feedback, ammonium, L-arginine, *Comamonas*

*It has been challenging to understand the principles by which microbial communities are formed (Fernandez *et al.*, 1999, 2000; Haruta *et al.*, 2013; El-Chakhtoura *et al.*, 2015). Microbial diversity, the functional stability of whole systems, and the coexistence of different microbes (Aziz *et al.*, 2015; Azwani *et al.*, 2021) are important features of microbial ecosystems and are relevant to the mechanisms by which they are formed. The functional stability of microbial communities is important for agricultural production, wastewater treatment, bioremediation, human health, and ecosystems, and is based on metabolic processes in response to changing environmental conditions.

Real microbial ecosystems are too complex to analyze because of intertwined relationships based on the functional diversity of the microbial world. Therefore, more simple and controllable systems, known as synthetic bacterial communities (SBCs), are needed. SBCs have increasing become the focus of research in recent years due to the reduced complexity of natural ecosystems and increased controllability (Haruta *et al.*, 2002; Kato *et al.*, 2005; Narisawa *et al.*, 2008; De Roy *et al.*, 2014; He *et al.*, 2014; Mee *et al.*, 2014; Aziz *et al.*, 2015; Friedman *et al.*, 2017; Haruta and Yamamoto, 2018). In our previous study, the SBC constructed with phenol-degrading bacteria, *Pseudomonas* sp. strain LAB-08, *Cupriavidus* sp. strain P-10, and *Comamonas testosteroni* strain R2, showed functional stability with coexistence under chemostat conditions for more than 800 days, and this was predicted to be dependent on a metabolic networking system (Azwani *et al.*, 2021).

Metabolic networking systems are attracting the attention of researchers interested in the mechanisms by which microbial ecosystems are formed (Freilich *et al.*, 2011; Morris *et al.*, 2013; Cao *et al.*, 2018; Hsu *et al.*, 2019). Interspecies interactions are considered to be relevant to the formation of metabolic networks. The supernatants of microbial cultures may affect the metabolic processes of other microbes (Tanaka *et al.*, 2005; Tashiro *et al.*, 2013; Inaba *et al.*, 2015), in which microbial metabolites play major roles in positive (Christensen *et al.*, 2002) and negative interactions.*

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**Article ME21050**
(Kim and Copley, 2012). In our efforts to clarify the coex-
isting mechanisms of three strains, LAB-08, P-10, and R2, we
found a unique phenomenon. A pure culture of strain
R2 suddenly collapsed under chemostat conditions supplied
with phenol as the sole carbon and energy source, even
though strain R2 has a complete set of genes relevant to
phenol-utilizing metabolism for the conversion of phenol
to acetyl-CoA (Powloski and Shingler, 1994; Azwani et al.,
2017) and grew and completely degraded phenol under
aerobic batch conditions with phenol as the sole carbon
and energy source (Watanabe et al., 1996; Futamata et al.,
2001a). These findings suggest that the three strains coex-
ist through mutualistic interactions. Therefore, analyses of
the collapsing process occurring in strain R2 will provide
insights into coexisting mechanisms, how the metabolic net-
work is formed, and how to manage microbes.

The aims of the present study were to analyze collapsing
processes using physiological and transcriptomic techniques
and to develop a method that rescues strain R2 from col-
lapse. Physiological analyses indicated that collapse was
induced by feedback growth inhibition, which had already
been initiated under stable conditions. The transcriptomic
analysis showed that the activities of primary metabolism
decreased by approximately 60 to 70% of stable conditions.
An imbalance between carbon and nitrogen metabolism
appeared to occur due to a decrease in the supply of carbon
skeletons and an increase in ammonium influx into cells.
We herein also discuss how strain R2 responded to negative
feedback, the mechanisms underlying the imbalance in car-on and nitrogen metabolism in cells, and how strain R2
was rescued from collapse.

Materials and Methods

Bacterium and culture conditions

The phenol-degrading bacterium C. testosteroni strain R2
was used in the present study. Strain R2 was isolated from activated
sludge in wastewater treatment at an oil refinery plant (Watanabe
et al., 1996). Strain R2 was precultured at 25°C in BSM medium
supplemented with phenol at 2.0 mM (BSM2.0phe medium)
under aerobic and batch conditions (Futamata et al., 2001a).
The medium contained the following (L–1): 12.5 g K2HPO4, 3.8 g
KH2PO4, 1.0 g (NH4)2SO4, 0.1 g MgSO4·7H2O, and 5 mL of
trace-element solution (pH 7.2). The trace-element solution con-
tained the following (L–1): 0.232 g H3BO3, 0.174 g ZnSO4·7H2O,
0.116 g Fe(NH4)2(SO4)2·6H2O, 0.096 g CoSO4·7H2O, 0.022 g
(NH4)6Mo7O24·4H2O, 8 mg CuSO4·5H2O, and 8 mg MnSO4·4H2O.
Cultures were harvested at the mid-exponential growth phase
and then transferred to 1.5 L of BSM medium containing
0.2 mM of phenol in a chemostat reactor (capacity of 2 L).
The initial cell density of strain R2 was adjusted to OD600 nm of
0.1. OD600 nm of 0.1 corresponded to 5.0×10⁸ cells mL⁻¹.
Strain R2 was precultured in BSM medium supplemented with
2.0 mM phenol (BSM2.0phe) and 0.3 mL of filter-sterilized supernatant
were transferred into 2.7 mL of fresh BSM2.0phe medium. The initial amount of cells
was adjusted to OD600 nm of 0.01. As the control condition, 0.3 mL
of BSM medium without phenol was added instead of the super-
натant. The growth curve was automatically measured using a
Biophotorecorder (TVS062CA, ADVANTEC). Growth parameters,
including the lag time (h), growth rate constant (μ h⁻¹), and amount of growth in the stationary phase (ODmax), were calculated
using the growth curve. We herein defined specific growth activity as surviving activity maintaining cell density at more than
1.0×10⁹ cells mL⁻¹ in a chemostat culture under the condition of
D. Therefore, 1 unit (U) of specific growth activity was calculated using the following equation: 1 U=0.105 (h⁻¹)×10⁹ (cells mL⁻¹)
under HRT of 10 days. As described above, the cell density of strain R2 was 5.0×10⁸ cells mL⁻¹ at an OD600 nm of 0.1. OD600 nm
was then converted to cell density. The unit of specific growth activity was calculated according to the following equation:
U=(μ×cell density from ODmax)/(1 U×lag time) (Aziz et al., 2015). The
proportion of U in the presence of the supernatant (Usup) to U
in the control condition (Ucont) was calculated as relative growth activity (%).

Real-time quantitative PCR (qPCR)

The population density of strain R2 was monitored using real-
time qPCR targeting the gene encoding the large subunit of phenol
hydroxylase (PH). Specific sets of primers were designed by the
alignment of genes encoding the large subunit of PH in strain R2
(Azwani et al., 2021). A specific PCR product amplified with a
specific primer set was used as the standard DNA fragment in the
qPCR analysis. To monitor strain R2, the qPCR profile consisted of preheating at 95°C for 10 min, followed by 40 cycles of denatu-
ration at 95°C for 10 s, annealing at 63°C for 5 s, and extension
at 72°C for 15 s. The fluorescence signal was detected at 72°C
in each cycle, and a melting curve was obtained by heating the
product to 95°C and cooling to 40°C. The reaction was performed
using a LightCycler FastStart DNA Master SYBR Green I kit
(Roche Molecular Biochemicals) and a LightCycler system (Roche
Diagnostics) according to the manufacturer’s instructions. The
copy number of each amplicon was calculated using LightCycler
software version 3.52. The copy number of the amplicon was equal
to the cell number because only one copy of the PH gene was
present in all strains (Azwani et al., 2017).
Kinetic analysis

A kinetic analysis was conducted to investigate the effects of metabolites on phenol- and catechol-degrading activities. Strain R2 was grown in a chemostat reactor with BSM medium and phenol as the sole carbon and energy source (R2-chemostat), and the kinetic properties of strain R2 for phenol and catechol degradation were investigated using the culture on days 18, 25, 30, and 35 according to a previously described method (Futama et al., 2001a; Haruta et al., 2013). When the accumulation of phenol and a decrease in OD_{600nm} were observed in the R2-chemostat, we considered the system to be collapsed. In total, 1.5 L of the culture was collected from the collapsed R2-chemostat and centrifuged at 5,800×g at 4°C. The supernatant was treated with the same volume of hexane, ethyl acetate, butanol, and dH₂O. Organic solvent fractions were concentrated to approximately 60 mL using a rotary evaporator (Buchi Rotavapor R3) under vacuum conditions (Buchi V-700) at 40°C. The H₂O fraction was mixed with methanol at a ratio of 1:3 and kept overnight at 4°C. The solution was then centrifuged at 5,800×g at 4°C and filtered (0.2 µm PTFE Membrane, OmnipureTM, Merck Millipore) to remove precipitates. The H₂O fraction was concentrated to approximately 60 mL by evaporating methanol using the rotary evaporator under vacuum conditions at 40°C. When the effects of the H₂O fraction on kinetic parameters were investigated, the culture of strain R2 and the H₂O fraction were mixed at a ratio of 9:1 for 30 min. Phenol- and catechol-oxidizing activities (the phenol and catechol consumption rates) were measured at various phenol and catechol concentrations, respectively, using an oxygen electrode (DO METER TD-51, Toko Chemical Lab.) after respiratory oxygen consumption had been suppressed by the addition of potassium cyanide (Watanabe et al., 1990; Haruta et al., 2013; Futama et al., 2005) using JMP statistical visualization software (SAS Institute). The apparent kinetic constants, \( K_A \) (affinity constant) and \( V_{max} \) (theoretical maximum activity) were assessed using the non-linear regression method as previously described (Azwani et al., 2021). As reported by Folsom et al. (1990), the term \( K_A \) was employed instead of \( K_m \) because activity was measured using intact cells rather than purified enzymes.

Transcriptomic analysis

A transcriptomic analysis was conducted to analyze contributing factors to the change from a stable to collapsed status in strain R2. Strain R2 was incubated in new chemostat cultures, reactors I and II, until collapse and cells were collected on days 17 (sample I-1), 22 (I-2), 30 (I-3), 31 (I-4), and 32 (I-5) from reactor I and on days 15 (sample II-1), 20 (II-2), 23 (II-3), 25 (II-4), and 28 (II-5) from reactor II (Supplementary Fig. S1). In the present study, samples I-1 and II-1 were defined as a stable status under which phenol did not accumulate and the population density was maintained, while samples I-5 and II-5 were defined as a collapsed status under which phenol accumulated. The library preparation and RNA sequencing of strain R2 were performed by Macrogen. The total RNA of strain R2 cultivated under chemostat conditions was extracted using the SV Total RNA Isolation System (Promega). The removal of ribosomal RNA molecules from total RNA was performed using the NEBNext rRNA Depletion Kit (Bacteria) (New England Biolabs) and strand-specific RNA sequencing libraries were prepared using TruSeq Stranded Total RNA Library Prep Gold (Illumina). Libraries were sequenced on NovaSeq 6000 (Illumina) to generate 2×101 bp paired-end sequence reads. Raw reads were cleaned using Trimmomatic ver. 0.36 by trimming adapter sequences, the base at the 3’t end, low-quality ends (quality score, <15), and dropping out reads of less than 75 bp (Bolger et al., 2014). The resulting high-quality reads were aligned to the genome sequence of strain R2 (GenBank accession number BDQJ00000000.1) using HISAT2 ver. 2.1.0 with options --dta and --no-spliced-alignment (Kim et al., 2019). Read counts were calculated from BAM files using featureCounts ver. 2.0.0 (Liao et al., 2014) and transcripts per million (TPM) values were calculated to normalize gene lengths and total read counts. The differentially expressed genes (DEGs) of strain R2 in the stable and collapsed statuses (I-1 vs I-5, and II-1 vs II-5) were analyzed using edgeR package ver. 3.16.4 (Robinson et al., 2010). Raw reads for genes were filtered by removing genes with low expression levels with a count per million (CPM) value of less than 2, and were normalized with scaling factors calculated for the library sizes using the Trimmed Mean of M-values (TMM) method (Robinson and Oshlack, 2010; Robinson et al., 2010). DEGs were defined by a log₂ fold-change (log₂FC) ≥ 2 (up-regulated) or ≤ –2 (down-regulated) and a false discovery rate (FDR) ≤ 0.05. Raw reads for RNA-seq analyzed in the present study have been deposited in the DDBJ Sequence Read Archive (DRA) under the accession numbers DRRR309243 to DRRR309247 and DRR309248 to DRR209252 for samples I-1 to II-5, respectively (Supplementary Table S1).

Growth of strain R2 in the presence of amino acids

We investigated whether the growth inhibition of strain R2 was rescued by the following amino acids: L-arginine, L-ornithine, L-citrulline, L-glutamate, and L-glutamine, because these amino acids are directly and indirectly related to the urea cycle. L-arginine, L-ornithine, and L-citrulline consisting of the urea cycle, and L-glutamate and L-glutamine are precursors for L-citrulline. Strain R2 cells precultured in BSM2.0phe liquid culture and 0.4 mL of filter-sterilized supernatant collected from collapsed chemostat cultures were transferred into 3.6 mL of fresh BSM2.0phe medium. The initial amount of cells was adjusted to 0.01 at OD_{600nm}. A stock solution of amino acids (200 mM) was prepared in BSM medium and the filter-sterilized stock solution was added at a final concentration of 10 mM. As the control condition, 0.4 mL of BSM medium without phenol was added instead of the supernatant and amino acid solution. The growth curve was automatically measured using a Bio-photorecorder (TVS062CA, ADVANTEC). Growth parameters, including the lag time (h), growth rate constant (µ [h⁻¹]), and the amount of growth in the stationary phase (OD_{max}), were calculated using the growth curve. As described above in the section on relative growth activity, the unit of specific growth activity (U) was calculated in the presence of phenol and an amino acid as a control (U_{amino acid}) or in the presence of phenol, the amino acid, and the supernatant (U_{amino acid+supernatant}). The negative growth effect (%) was calculated using the following equation:

\[
\text{Negative growth effect} = \left(1 - \frac{U_{amino acid+supernatant}}{U_{amino acid}}\right) \times 100.
\]

Scanning electron microscopy (SEM) observations

SEM was used to observe the morphology of C. testosteroni strain R2 on days 18 and 38 in the chemostat culture. Samples were fixed with a mixed solution of 25% (v/v) glutaraldehyde and 5% (v/v) formaldehyde for 2 h in potassium phosphate buffer (0.2 M, pH 7.0) and dehydrated with ethanol with serially increasing concentrations (30, 50, 75, 95, and 99.9% of ethanol for 15 min each) and then with 100% of butyl alcohol. The sample was mounted on an aluminium stub with double-sided carbon tape, and sputter-coated with gold under argon at a thickness of 50 Å in the Quick Auto Coater (SC-701AT, Sanyu Denshii) for 20 s. Following coating, samples were imaged at different magnifications with a field emission scanning electron microscope (Model JSM-6335F, JEOL) at an acceleration voltage of 5 kV and working distance of 5 mm.

Chemical analysis

Phenol and catechol concentrations were monitored using high-pressure liquid chromatography (Waters Japan) equipped with a column (YMC-Triart C18 [150×2 mm], YMC) and UV detector. Liquid samples collected from the chemostat culture were centri-
fuged and filtered (Millipore LG [pore size of 0.2 μm, diameter of 13 mm], Millipore). Liquid samples were eluted using 50% acetonitrile solution with 20 mM ammonium acetate delivered at 0.1 mL min⁻¹, and elutes were monitored at 210 nm. Phenol and catechol were identified according to their retention times of 7.2 and 5.2 min, respectively. Concentrations were assessed by comparing the peak area with that of the cognate standard sample.

**Statistical analysis**

DEGs were identified by the likelihood-ratio test implemented in the edgeR package. Other data were analyzed using the Student’s t-test. P=0.05 was considered to be significant.

**Results**

**Collapse of strain R2 growth under chemostat conditions**

Strain R2 grew aerobically in the presence of phenol as the sole carbon and energy source and completely utilized phenol in batch cultures (Fig. 1A). Catechol, a metabolite produced by PH, was not detected during the experiment (data not shown). In chemostat cultures, the growth amount of strain R2 reached 0.96±0.010 at OD 600 nm from days 4 to 14 and then gradually decreased (Fig. 1B). The concentration of phenol was maintained at 0.13±0.010 mM until day 32 and increased to 1.3 mM±0.010 on day 38, namely, the growth of strain R2 collapsed (Fig. 1B). Relative growth activity was stable at 74±0.64% until day 25, after which it rapidly decreased to 32±0.080% on day 38 in the collapsed status (Fig. 1C). The viability of strain R2 cells decreased from 100±0.10% on day 18 to 50±1.8% on day 38 (Fig. 1D). SEM observations showed that the morphology of strain R2 cells changed in the collapsed status (Supplementary Fig. S2).

**Effects of the chemostat supernatant on kinetic properties**

The kinetic properties of phenol and catechol degradation by strain R2 were investigated using R2-chemostat cultures on days 18, 25, 30, and 35. \( V_{\text{max}} \) for phenol and catechol were 65±6.4 and 130±13 mM [g dry cell]⁻¹ respectively, on day 18, and gradually decreased to 9.4±2.5 and 9.1±0.18 mM [g dry cell]⁻¹, respectively, on day 38 (Fig. 2A and B).

We hypothesized that certain metabolites accumulating in the chemostat culture of strain R2 may be responsible for the collapse of its growth. To clarify this, we investigated the effects of supernatant (SN) collected from chemostat cultures of strain R2 on kinetic parameters for phenol and catechol degradation. As a preliminary test, we fractionated the supernatant using H₂O and organic solvents (i.e., hexane, ethyl acetate and butanol) and found that the H₂O fraction exerted the strongest negative effects on the relative growth activity of strain R2 (Supplementary Fig. S3). The H₂O fraction of supernatant was used to establish whether the activities of PH and catechol dioxygenase were inhibited by metabolites produced from strain R2. \( V_{\text{max}} \) values were not markedly affected, whereas \( K_{S} \) values for PH and catechol dioxygenase increased from 0.70±0.050 to 1.3±0.10 μM for phenol and from 14±1.4 to 36±0.70 μM for catechol in the presence of the H₂O fraction (Fig. 2 C and D).

**Change in metabolism from the stable to collapsed status**

The results of physiological analyses indicated that the growth collapse of strain R2 under chemostat conditions was triggered by metabolic changes. To elucidate the met-
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Catechol-degradation rate

0 40 80
0 120
(B)

10¹ 10² 10³ 10⁴
Catechol concentration (μM)

Phenol-degradation rate

0 20 40 60
(A)

10¹ 10² 10³ 10⁴
Phenol concentration (μM)

Relative transcriptional levels from the stable to collapsed status slightly decreased from 0.5- to 0.7-fold in the main metabolic pathways: glycolysis, gluconeogenesis, the pentose phosphate pathway, 5-phosphate-α-D-ribose 1-diphosphate (PRPP) synthesis (Supplementary Fig. S5 and Supplementary Table S2), and nucleotide synthesis (Supplementary Fig. S6 and Table S3). The relative transcriptional levels of genes encoding enzymes in the pentose phosphate pathway decreased to 0.67±0.20-fold in the collapsed status, whereas those of genes encoding PRPP decreased to 0.44-fold and 0.47-fold in the collapsed status in reactors I and II, respectively (Supplementary Fig. S5C).

In the TCA cycle, the relative transcriptional levels of genes encoding enzymes related to the conversion of 2-oxoglutarate (2-OG) to succinate (2-OG dehydrogenase [sucA, CTR2_2972, and sucB, CTR2_2971] and succinyl-CoA synthetase [sucC, CTR2_4841, and sucD, CTR2_4842]) significantly decreased by 0.34±0.065-fold in the collapsed status (Fig. 3B and 4), whereas those of the

Relative transcriptional levels from the stable to collapsed status slightly decreased from 0.5- to 0.7-fold in the main metabolic pathways: glycolysis, gluconeogenesis, the pentose phosphate pathway, 5-phosphate-α-D-ribose 1-diphosphate (PRPP) synthesis (Supplementary Fig. S5 and Supplementary Table S2), and nucleotide synthesis (Supplementary Fig. S6 and Table S3). The relative transcriptional levels of genes encoding enzymes in the pentose phosphate pathway decreased to 0.67±0.20-fold in the collapsed status, whereas those of genes encoding PRPP decreased to 0.44-fold and 0.47-fold in the collapsed status in reactors I and II, respectively (Supplementary Fig. S5C).

On the other hand, relative transcriptional levels in some metabolic pathways were moderately repressed, maintained, or increased. In the phenol-/catechol-degrading pathways, relative transcriptional levels decreased to 0.40±0.28-fold in the collapsed status, with the exception of genes encoding the subunit of PH (dmpK, CTR2_1592) and catechol 2,3-dioxygenase (dmpB, CTR2_1599). The genes encoding PH and catechol 2,3-dioxygenase were up-regulated by approximately 30- and 3.3-fold, respectively, in reactor I, and by 15- and 1.2-fold, respectively, in reactor II (Fig. 3A and 4, Table 1, and Supplementary Table S4-1).

In the TCA cycle, the relative transcriptional levels of genes encoding enzymes related to the conversion of 2-oxoglutarate (2-OG) to succinate (2-OG dehydrogenase [sucA, CTR2_2972, and sucB, CTR2_2971] and succinyl-CoA synthetase [sucC, CTR2_4841, and sucD, CTR2_4842]) significantly decreased by 0.34±0.065-fold in the collapsed status (Fig. 3B and 4), whereas those of the
Fig. 3. Relative transcriptional levels of genes encoding enzymes in phenol/catechol degradation, the TCA cycle, glyoxylate shunt, and urea cycle in reactors I and II.

(A-I) and (A-II): Phenol/catechol degradation in reactors I and II, respectively; 1: from phenol to catechol; 2: from catechol to 2-hydroxyxuconic semialdehyde; 3: from 2-hydroxyxuconic semialdehyde to 2-hydroxyxuconate; 4: from 2-hydroxyxuconate to gamma-oxalocrotonate; 5: from gamma-oxalocrotonate to 2-oxopent-4-enoate; 6: from 2-oxopent-4-enoate to 4-hydroxy-2-oxopentanoate; 7: from 4-hydroxy-2-oxopentanoate to acetaldehyde/pyruvate; 8: from acetaldehyde/pyruvate to acetyl-CoA. (B-I) and (B-II): the TCA cycle in reactors I and II, respectively; 1: from acetyl-CoA and oxaloacetate to citrate; 2 and 3: from citrate to isocitrate; 4: from isocitrate to 2-oxoglutarate; 5: from 2-oxoglutarate to s-succinyl dihydrolipoyllysine; 6: from s-succinyl dihydrolipoylysine to succinyl-CoA; 7 and 8: from succinyl-CoA to succinate; 9, 10, 11, and 12: from succinate to fumarate; 13 and 14: from fumarate to malate; 15 and 16: from malate to oxaloacetate. (C-I) and (C-II): the glyoxylate shunt in reactors I and II, respectively; 1: from acetyl-CoA and oxaloacetate to citrate; 2 and 3: from citrate to isocitrate; 4: from isocitrate to glyoxylate; 5: from glyoxylate to malate; 6: from malate to oxaloacetate. (D-I) and (D-II): GS-GOGAT and GDH synthesis; 1: from L-glutamate to L-glutamine by glutamine synthetase [EC:6.3.1.2]; 2: from L-glutamine to L-glutamate; 3: from L-glutamine to L-glutamate; 4: from 2-OG and NH$_3$ to L-glutamate; 5: from L-glutamate to 2-OG and NH$_3$. Bold black line: stage I-I or stage II-I; black line: stage I-II or stage II-2; broken line: stage I-3 or stage II-3; gray line: stage I-4 or stage II-4; and bold gray line: stage I-5 or II-5. (E-I) and (E-II): the urea cycle in reactors I and II, respectively; 1: from L-argininosuccinate to L-arginine and fumarate; 2: L-arginine to L-ornithine and urea; 3: from L-ornithine to L-citrulline; 4: from L-citrulline and L-aspartate to L-argininosuccinate. The genes, locus tag numbers, enzymes, ko numbers, and transcripts per million values of genes are listed in Supplementary Table S4.
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Fig. 4. Schematic diagram of metabolism in strain R2 in the collapsed status. All values are shown as a percentage of the relative transcriptional levels of genes in the collapsed status to those in the stable status. a: the values in reactors I and II on the left and right sides of the slash, respectively; b: from 2-hydroxymuconic semialdehyde to acetyl-CoA; c: from pyruvate to fructose 6-phosphate; d: from fructose 6-phosphate to pyruvate; e: from acetyl-CoA and oxaloacetate to 2-oxoglutarate; f: from 2-oxoglutarate to succinate; g: from succinate to oxaloacetate; h: from isocitrate to malate via glyoxylate; i: glutamate dehydrogenase [EC:1.4.1.3]; GDH: glutamate dehydrogenase; GS: glutamine synthetase; GOGAT: glutamate synthase. Arrow sizes are scaled to relative transcriptional levels: (1) bold arrow; highly expressed in both reactors I and II (e.g., phenol/toluene 2-monooxygenase, catechol 2,3-dioxygenase, and GS), (2) middle-sized arrow; maintained or slightly decreased expression level (e.g., 60%-100%) in either reactor, (3) dotted arrow; decreased (e.g., less than 60%) in both reactors. Genes, locus tag numbers, enzymes, ko numbers, and fold values in # numbered pathways are shown in Table 1.

Other genes moderately decreased by 0.73±0.25-fold from citrate to 2-OG (citrate synthase [gltA, CTR2_1354], aconitase hydratase [acnA, CTR2_1384, and acnB, CTR2_1366], and isocitrate dehydrogenase [icd, CTR2_1985]), and to 0.80±0.23-fold from succinate to oxaloacetate (succinate dehydrogenase/fumarate reductase [sdhA/frdA, CTR2_1357, sdhB/frdB, CTR2_1356, sdhC/frdC, CTR2_1359, and sdhD/frdD, CTR2_1358], fumarate hydratase [fumA/fumB, CTR2_5301, and fumC, CTR2_5299], and malate dehydrogenase [mdh, CTR2_1362, and mqo, CTR2_1070]) (Fig. 3B and 4, and Supplementary Table S4-2). In the glyoxylate shunt, the relative transcriptional levels of all 6 genes moderately decreased from the stable to collapsed status with the exception of the gene encoding isocitrate lyase (aceA, CTR2_1688) (Fig. 3C and 4, Table 1, and Supplementary Table S4-3). The relative transcriptional levels of the gene encoding isocitrinate lyase (aceA, CTR2_1688) fluctuated in the collapsed status and became similar to or higher than those in the stable status.

The relative transcriptional levels of genes encoding enzymes in the biosynthesis of amino acids decreased to 0.63±0.12-fold in the collapsed status, with the exception of the synthesis of L-leucine, L-glutamate, and L-glutamine (Supplementary Fig. S7). The levels of glutamate dehydrogenase (GDH) (gdhA, CTR2_3645) in the collapsed status decreased to ~0.40-fold those in the stable status. The relative transcriptional levels of glutamate synthase (GOGAT) (gltD, CTR2_4073 and gltB, CTR2_4074) and glutamine synthetase (GS) (glnA, CTR2_1472) increased by approximately 2-fold in the collapsed status (Fig. 3D and 4, Table 1, and Supplementary Table S4-4).

In the urea cycle, the relative transcriptional levels of the 4 genes encoding enzymes were separated into two groups: the relative transcriptional levels of genes encoding enzymes related to the conversion of L-argininosuccinate to L-ornithine via L-arginine (argininosuccinate lyase [argH, CTR2_1809] and arginase [rocF, CTR2_0414]) moderately decreased to 0.82±0.22-fold in the collapsed status, whereas those of genes encoding enzymes related to the conversion from L-ornithine to L-arginino succinate via L-citrulline (ornithine carbamoyltransferase [argF, CTR2_1066] and argininosuccinate synthase [argG, CTR2_5281]) significantly decreased to 0.47±0.051-fold in the collapsed status (Fig. 3E and 4, Table 1, and Supplementary Table S4-5).

Effects of amino acids on growth inhibition

We investigated whether the growth inhibition of strain R2 was rescued by amino acids from the urea cycle (L-
arginine, L-ornithine, and L-citrulline), L-glutamate, and L-glutamine (Fig. 5). Lag times and μ values in the presence of phenol and amino acids from the urea cycle were similar to those in the positive control incubated in the presence of phenol only, whereas OD₉₀ max decreased to between 80 and 90% of the positive control (Table 2). Growth parameters in the presence of a supernatant (SN) collected from the collapsed culture were similar to those of the negative control, in the presence of a supernatant (SN) collected from the collapsed culture. Unexpectedly, increases were noted in the relative transcriptional levels of the genes encoding PH and catechol 2,3-dioxygenase (Fig. 4, Table 1). These genes are located in an operon regulated by the regulator protein DmpR (Azwani et al., 1998), indicating that phenol concentrations had increased in cells. Since PH and catechol 2,3-dioxygenase were competitively inhibited by the metabolites (Fig. 2), we considered a deficiency in a usable carbon source to be a trigger for collapse. Transcriptomic analyses showed that the actual condition of the collapsing mechanism was metabolically complex; metabolites induced an imbalance in carbon and nitrogen metabolism in cells, which resulted in collapse. We discussed how the imbalance occurred and why L-arginine effectively rescued it.

Discussion

The present study attempted to elucidate the collapsing mechanism induced by feedback growth inhibition in strain R2. Collapse was not observed under batch conditions, it only occurred under chemostat conditions. The results obtained revealed that growth inhibition and metabolic changes had already occurred before the collapse, which was attributed to negative feedback induced by the accumulation of hydrophilic metabolites produced from strain R2. Unexpectedly, increases were noted in the relative transcriptional levels of the genes encoding PH and catechol 2,3-dioxygenase (Fig. 4, Table 1). These genes are located in an operon regulated by the regulator protein DmpR (Azwani et al., 1998), indicating that phenol concentrations had increased in cells. Since PH and catechol 2,3-dioxygenase were competitively inhibited by the metabolites (Fig. 2), we considered a deficiency in a usable carbon source to be a trigger for collapse. Transcriptomic analyses showed that the actual condition of the collapsing mechanism was metabolically complex; metabolites induced an imbalance in carbon and nitrogen metabolism in cells, which resulted in collapse. We discussed how the imbalance occurred and why L-arginine effectively rescued it.

The transcriptomic analysis demonstrated that relative
transcriptional levels decreased in the majority of carbon metabolic pathways, indicating that the supply of carbon skeletons for nitrogen assimilation was limited under growth-inhibiting conditions. On the other hand, the relative transcriptional levels of some metabolic pathways moderately decreased, were maintained, increased, or fluctuated, which were considered to be adaptive processes to negative feedback. The relative transcriptional level of the isocitrate lyase gene (aceA, CTR2_1688) related to the TCA cycle and glyoxylate shunt fluctuated and increased under collapsed conditions (Fig. 3C), indicating that strain R2 adapted to a carbon source deficiency because carbon was not released as CO₂ in the glyoxylate shunt. The relative transcriptional levels of the metabolic pathways from citrate to 2-OG in the TCA cycle in the collapsed state were limited to approximately 0.70-fold those in the stable status, whereas the metabolic pathways from 2-OG to succinate were repressed to approximately 0.35-fold (Fig. 3B and 4, and Supplementary Table S4-2). 2-OG may be used in metabolic pathways other than the TCA cycle because it plays a major role in carbon skeletons for the biosynthesis of nitrogenous compounds via L-glutamate and L-glutamine (Merrick and Edwards, 1995). GDH and GS have been widely suggested to play a central role in responses to ammonia stress by converting ammonia to non-toxic or less toxic nitrogenous compounds (Zhang et al., 2020). In the collapsed status, the transcriptional level of GDH (gdhA, CTR2_3645) decreased (Fig. 3D and 4, and Table 1), whereas those of GS (glnA, CTR2_1472) and GOGAT (gltD, CTR2_4073 and gltB, CTR2_4074) (the GS-GOGAT pathway) increased (Fig. 3D and 4, and Table 1). Differences in activities between the GS-GOGAT and GDH pathways are considered to depend on differences in the affinity for ammonium of GS (~0.1 mM) (Sharkey and Engel, 2008) and GDH (more than 1 mM) (Alibhai and Villafranca, 1994), whereas the GS-GOGAT pathway is the main ammonium-assimilatory route even with a high ammonium concentration (10 mM) in E. coli (Yuan et al., 2009). The Kₘ values of GS and GDH for 2-OG are ~0.24 mM (Mantsala and Zalkin, 1976) and ~0.64 mM (Veronese et al., 1975; Sharkey and Engel, 2008), respectively, suggesting that strain R2 is under 2-OG deficiency conditions. The activity of Icd (CTR2_1985) producing 2-OG from isocitrate decreased in the presence of phenol (10 mM) because the excess supply of carbon skeletons enabled the transcription of the isocitrate lyase gene (aceA, CTR2_1688) for the production of 2-OG. The activity of 2-OG dehydrogenase (Icd) reduced by the collapse of chemostats and repressed by the addition of supernatant from collapsed chemostats, which may be due to negative feedback. The relative transcriptional levels of the metabolic pathways in the presence of phenol and amino acid, and the supernatant.

Table 2. Effects of amino acids on the growth of strain R2

| Condition  | Lag time (h) | Growth rate constant (μ) | OD₆₀₀nm | Specific growth activity (U) | Negative growth effect (%)  |
|------------|-------------|--------------------------|--------|----------------------------|-----------------------------|
| Positive control | 10±1.2 | 0.25±0.0064 | 0.40±0.0093 | 0.048±0.0027 | 42±0.60 |
| Negative control | 12±1.0 | 0.17±0.011 | 0.40±0.0087 | 0.027±0.0024 | 3.2±8.5 |
| Arg  | 10±0.50 | 0.29±0.0074 | 0.32±0.0064 | 0.044±0.0070 | 34±11 |
| Arg+SN  | 8.8±0.58 | 0.25±0.013 | 0.32±0.019 | 0.043±0.0037 | 40±15 |
| Orn  | 9.5±0.50 | 0.26±0.0065 | 0.31±0.039 | 0.041±0.0050 | 59±6.3 |
| Orn+SN  | 11±0.58 | 0.19±0.031 | 0.33±0.029 | 0.027±0.0030 | 27±10 |
| Ctr  | 11±0.29 | 0.26±0.0057 | 0.36±0.010 | 0.041±0.0022 | 4.2±0.60 |
| Ctr+SN  | 11±0.29 | 0.22±0.0090 | 0.27±0.083 | 0.025±0.00061 | 34±11 |
| Glu  | 4.8±0.29 | 0.35±0.0093 | 1.2±0.036 | 0.41±0.015 | 40±15 |
| Glu+SN  | 8.5±1.5 | 0.25±0.0047 | 1.2±0.035 | 0.17±0.025 | 27±10 |
| Gln  | 9.0±1.0 | 0.26±0.0040 | 1.2±0.046 | 0.17±0.021 | 4.2±0.60 |
| Gln+SN  | 11±0.56 | 0.22±0.0051 | 1.3±0.027 | 0.12±0.0033 | 34±11 |

Notes: Phenol was added as the sole carbon and energy source in all conditions.
Arg: L-arginine, Orn: L-ornithine, Ctr: L-citrulline, Glu: L-glutamate, Gln: L-glutamine, negative growth effect (%): the effect was calculated by the following equation:

\[ (1-\frac{U_{\text{amino acid}+\text{SN}}}{U_{\text{amino acid}+\text{SN}}}) \times 100 \]

where \( U_{\text{amino acid}} \) is relative growth activity in the presence of phenol and an amino acid, and \( U_{\text{amino acid}+\text{SN}} \) is relative growth activity in the presence of phenol, the amino acid, and the supernatant.
trate in the TCA cycle may be repressed because enzyme activity is positively regulated by metabolites derived from glycolysis/glucoseogenesis (Cozzone and El-Mans, 2005), which supports the hypothesis of a 2-OG pool deficiency in strain R2 in the collapsed state.

2-OG acts as a metabolic signal of small and transient fluctuations in ammonium availability in Escherichia coli, i.e., the concentration of 2-OG increases with ammonium limitations under steady-state growth conditions (Senior, 1975; Reyes-Ramirez et al., 2001) and vice versa (Yuan et al., 2009; Radchenko et al., 2010; Yan et al., 2011). A negative correlation has been reported between ammonium availability and 2-OG accumulation in other microorganisms (Muro-Pastor et al., 2001; Dodsworth et al., 2005; Brauer et al., 2006). This negative correlation contributes to the balance between carbon and nitrogen metabolism: the decrease in 2-OG and increase in L-glutamine facilitate the construction of the complex of the ammonium transporter (AmtB) and regulator protein (GlnK), resulting in the repression of NH$_4^+$ influx through the AmtB-GlnK complex (Radchenko et al., 2010), and the replenishment of the 2-OG pool is then achieved by increased transamination after the consumption of ammonium. However, the relative transcriptional levels of nitrogenous compound synthesis decreased not only in nucleotide synthesis (Supplementary Fig. S6), but also in amino acid synthesis, with the exception of L-leucine, L-glutamate, and L-glutamine (Supplementary Fig. S7). Furthermore, the gene encoding the ammonium transporter Amt family (CTR2_4687) was up-regulated by ~190-fold in the collapsed status (Supplementary Table S5-1), suggesting that cellular ammonium concentrations increased in the collapsed status. In parallel, the genes encoding nitrite reductase (nirB, CTR2_4180) and the urea ABC transporter (CTR2_0055) were up-regulated by ~155- and ~20-fold, respectively, in the collapsed status (Supplementary Table S5-1), indicating that strain R2 attempted to decrease endogenous ammonium toxicity and balance carbon and nitrogen metabolism by reducing excess ammonium. The gene encoding AmR was significantly up-regulated even under conditions of low 2-OG and high glutamine concentrations, which has yet to be resolved with the identification of growth-inhibiting metabolites. This imbalance may affect various metabolic pathways; e.g., outer membrane protein porin (CTR2_5017) and cell division protein (CTR2_0482) were down-regulated (Supplementary Table S5-3 and S5-4), whereas PilA (CTR2_4856) was up-regulated in the collapsed status (Supplementary Table S5-1 and S5-2). The imbalance in metabolism related to the cell structure may have caused the disordered cells observed in the collapsed status (Supplementary Fig. S2B). The mechanisms maintaining the balance between carbon and nitrogen metabolism in strain R2 have not yet been elucidated.

The urea cycle is an essential pathway for the disposal of ammonia in mammals, and the genes encoding enzymes in the urea cycle in mammals are evolutionarily derived from bacterial genes for the detoxification of ammonia (Walsh and Patricia, 1995). A previous study reported that the urea cycle of Helicobacter pylori acts as an effective mechanism to extrude excess nitrogen from cells (Mendz and Hazell, 1996). Therefore, we attempted to enhance the activity of the urea cycle by adding amino acids to the urea cycle because the urea cycle of strain R2 was repressed in the collapsed status (Fig. 3E and 4, and Table 1). Only L-arginine enabled the rescue of strain R2 from growth inhibition without an increase in the number of cells (Fig. 5A and Table 2), indicating that L-arginine was not used in anabolic processes, but functioned as a trigger to enhance the urea cycle. The mechanisms by which L-arginine rescues strain R2 from growth inhibition warrant further study.

Conclusion

The present study revealed the metabolic responses of strain R2 to negative feedback (Fig. 4 and Table 1). The imbalance in carbon and nitrogen metabolism was caused by both an increase in the concentration of NH$_4^+$ and a decrease in the supply of carbon skeletons, which resulted in collapse. 2-OG and PRPP are at the metabolic intersection between carbon and nitrogen metabolism, with 2-OG playing a major role as not only a master regulator (Huergo and Dixon, 2015), but also a major carbon skeleton in nitrogen-assimilatory reactions (Commichau et al., 2006). Nitrogen regulates primary and secondary metabolism in various bacteria (Merrick and Edwards, 1995); therefore, a more detailed understanding of the regulatory mechanisms of NH$_4^+$ concentrations in cells will be indispensable for managing microorganisms. The monitoring of these compound concentrations in cells will provide insights into collapse and rescue mechanisms and these analyses will contribute to the clarification of bacterial coexisting mechanisms in microbial ecosystems.

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