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Tolerance against butanol stress by disrupting succinylglutamate desuccinylase in *Escherichia coli*[†]

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Background: The four-carbon alcohol, butanol, is emerging as a promising biofuel and efforts have been undertaken to improve several microbial hosts for its production. However, most organisms have very low tolerance to *n*-butanol (up to 2% (v/v)), limiting the economic viability of butanol production. Although genomic tools (transcriptomics, proteomics, and metabolomics) have been widely used to investigate the cellular response to butanol stress, the existing knowledge of the molecular mechanisms involved in butanol tolerance is limited, and strain improvement is difficult due to the complexity of the regulatory network. Results: In this study, a butanol-tolerant Escherichia coli was constructed by disrupting gene astE (encoding succinylglutamate desuccinylase) to obtain higher butanol tolerance (increased by 34.6%). To clarify the tolerance mechanism, a metabolome analysis was also performed. As a result, a total of 73 metabolites (11 elevated and 62 decreased) were significantly changed. Most of the downregulated metabolites were mainly involved in the L-arginine degradation pathway, sulfate metabolic pathway, and 2-methylcitrate metabolic pathway. To further analyze the differential gene expression, a transcriptome was created. In total, 311 genes (113 upregulated and 198 downregulated) showed over a twofold difference and were associated with carbohydrate metabolism, energy metabolism, and ABC transporters. The integration of metabolomics and transcriptomics found that acid-activated glutaminase ybaS and the amino acid antiporter gadC were significantly up-regulated, but the levels of L-arginine and glutamate were not significantly increased and decreased. Therefore, the changes of amino acids between strains BW25113 and BW25113-AastE were measured by amino acid analysis. The ability of a mutant strain against acid stress was also measured by the growth experiment under various pH conditions in the absence of butanol. Conclusions: Based on the above experiments, it could be concluded that mutant BW25113- Δ astE mainly regulated intracellular pH-homeostasis to adapt to butanol stress, indicating the non-negligible impact of pH on microbial butanol tolerance, broadening our understanding of microbial butanol tolerance and providing a novel strategy for the rational engineering of a more robust butanol-producing host.

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Background

Recently, global environmental problems and fuel crises have drawn public attention toward the development of microbial production for fuels and chemicals, such as ethanol, higher alcohols,^{1,2} biodiesel,³ and alkanes.⁴ However, most of them are toxic to microbial cells, leading to a low fermentation titer and high cost in downstream product recovery.⁵ For example, butanol is a promising biofuel but is highly toxic to microbes⁶ (such as *Clostridia*, *Escherichia coli*, and *Saccharomyces cerevisiae*),⁷ and only 13 g L⁻¹ of butanol is sufficient to restrain cell growth, which is a major drawback in practical butanol production.⁸ Therefore, understanding the mechanisms of the response to butanol and improving microbial robustness will further ameliorate butanol toxicity, and maintain microbial stability and the dynamic homeostasis of butanol fermentation.⁹

To date, several mechanisms underlying the physiological responses to solvent have been identified in eukaryotic and prokaryotic microorganisms. The hydrophobic nature and lipid solubility of butanol could: (i) alter cell membrane fluidity; (ii) denature membrane proteins; and (iii) impair membrane-related processes (*e.g.* nutrient transport, respiration, and photosynthetic electron transport).⁸ For example, butanol can

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effectively inhibit the activities of membrane-bound ATPases, resulting in a lower internal pH and the abolishment of a pH gradient across the membrane.¹⁰ Furthermore, *n*-butanol stress also promotes the release of autolysin, which can hydrolyze bacterial components by breaking down the *b*-1,4-bond between *N*-acetylmuramic acid and *N*-acetylglucosamine molecules.¹¹ However, the butanol tolerance mechanism is complex and the existing knowledge of genes involved in butanol tolerance needs to be further expanded. The strategies for strain improvement should consider contributes synergistically to obtain a strain with the desired properties.¹²

At present, many strategies have been carried out to improve cellular robustness for butanol, including classical mutagenesis, metabolic engineering, and transcriptional engineering.13 For example, a genomic library enrichment strategy was performed for E. coli, in which approximately 270 candidate genes (enriched or deleted) were identified against n-butanol stress.¹⁴ Interestingly, the deletion of gene astE, encoding succinylglutamate desuccinylase, significantly enhanced n-butanol tolerance with increases of 48.7%,14 but the physiological mechanism has not been comprehensively elucidated (Fig. S1[†]). AstE gene encodes succinyl-glutamate desuccinylase, which catalyzes the fifth and final reaction in the ammonia-producing arginine catabolic pathway,15 can decompose succinyl-glutamate into succinic acid and glutamate, thus regulating the intracellular arginine concentration. Deletion of astE enhances tolerance to nbutanol.16 In this study, we aimed to comprehensively elucidate the tolerance mechanism for the disruption of the astE gene in E. coli. To this end, a butanol-tolerant strain was constructed by disrupting gene astE, to obtain a higher butanol tolerance (increase by 34.6%). In addition, to clarify the tolerance mechanism, metabolome and transcriptome analyses were performed to assess differential gene expression and related metabolic pathways. It was found that mutant BW25113-\DeltaastE had developed a special tolerance through regulating cellular pHhomeostasis. This expands the current knowledge on butanol tolerance and provides a novel strategy for the rational engineering of a more robust butanol-producing host.

Results

Effects of knocking-out *astE* on cellular robustness

To investigate the effect of knocking-out astE on cell growth under butanol stress, a spotting assay on agar plates containing different butanol concentrations was applied. Compared to the wild strain BW25113, the mutant BW25113-ΔastE could grow well on agar plates containing 8 g L^{-1} of butanol (Fig. S2[†]). Furthermore, cell tolerance was also evaluated in shake-flask cultures. As shown in Fig. 1a, under control conditions (0 g L^{-1} butanol), strain BW25113 and BW25113- Δ astE showed a similar growth ability, respectively, indicating that knockingout gene astE had no effect on cell growth. However, when butanol stress was increased to 8 g L⁻¹, the biomass of BW25113-ΔastE could reach 0.51 after cultivating for 24 h, which is an increase of 34.6% compared with that of BW25113 (Fig. 1b). Therefore, knocking-out gene astE could effectively improve cellular robustness, showing a higher butanol tolerance (increased by 34.6%), which agreed with a previous study.14

GC-MS based metabolomics analysis

To further investigate the tolerance achieved by knocking-out gene *astE*, the difference in metabolic responses between BW25113 and BW25113- Δ astE after exposure to 0 g L⁻¹ and 5 g L⁻¹ butanol was determined with gas chromatography-mass spectrometry (GC-MS). Metabolite profiles were generated and data matrixes were analysis by the principal component analysis (PCA) (see Fig. 2). In which the differences of 76.7% in the metabolite profiles were explained by the first principal component. Compared to the wild strain, mutant strains under the stress of 0 g L⁻¹ or 5 g L⁻¹ butanol showed relatively concentrated at PCA analysis, indicating that mutant strain has better robustness under butanol stress.

Meanwhile, under the 5 g L^{-1} butanol stress, there was a lots of differential metabolites between the wild strain and mutant strain based on the metabolomic cloud plot (Fig. S3†). The size of each bubble corresponds to the log fold change of the feature,



Fig. 1 Effects of knocking-out gene *astE* on cell growth under different butanol concentrations: (a) 0 g L⁻¹; (b) 8 g L⁻¹ (\triangle : BW25113; \square : BW25113- Δ astE).



Fig. 2 Principal component analysis of strains in the presence and absence of butanol.

the larger the bubble the bigger the log fold change. The colors of features with low *p*-values are brighter than which features with high *p*-values.

Therefore, we select the wild and mutant strain under the stress of 5 g L^{-1} butanol for detailed metabolite analysis and transcriptome analysis.

In order to get a better discrimination between BW25113 and BW25113- Δ astE after exposure to 5 g L⁻¹ butanol, orthogonal partial least square-discriminant analysis (OPLS-DA) was applied in this study (Fig. 3). PCA and OPLS-DA analysis were performed using the OmicShare tools. The OPLS-DA showed a clear separation of samples into two distinct groups, indicating that BW25113 and BW25113- Δ astE had a significantly different metabolic profile. Score plots using the first two principal components were used to present a 2D representation of variations among the spectra. From the metabolic cloud plot in Fig. S3,† after exposure to 5 g L⁻¹ butanol, a total of 73



Fig. 3 Orthogonal partial least square-discriminant analysis of strains BW25113 and BW25113- Δ astE after exposure to 5 g L⁻¹ butanol.

metabolites were significantly changed between BW25113 and BW25113- Δ astE, of these 11 were significantly elevated, including 6-carboxy-5,6,7,8-tetrahydropterin, thymidine, 2-deoxycytidine, 7,8-dihydromonapterin, L-phenylalanine, 1-myristoyl-2-palmitoleoyl phosphatidate, L-leucine, γ -butyr-obetaine, and L-valine, whereas 62 of them, such as 6-deoxy-6-sulfo-D-fructose 1-phosphate, and *N*-acetylmuramate, were significantly decreased (Table S1†).

Investigation of the tolerance mechanism by transcriptomics

To further elucidate the tolerance mechanism of knocking out gene *astE*, transcriptomics of strains BW25113 and BW25113- Δ astE were analyzed after exposure to 5 g L⁻¹ butanol stress. Reviewer/collaborator link to metadata: http://ftp://ftp-trace.ncbi.nlm.nih.gov/sra/review/SRP179828_20190117_0935-16_305cf1fb13b9539dcd317a0354c9ed61.

As shown in Fig. S4,[†] based on the corrected *P*-value < 0.05, the results showed that 311 genes were considered to have a significantly different expression in response to butanol stress, of these, 113 genes were upregulated and 198 genes were downregulated.

GO enrichment analysis of the differentially expressed genes

GO enrichment analysis was performed to cluster genes with a significant difference in transcription level. The same types of genes were then gathered in a cluster with similar biological functions to intuitively understand the relationships and discrepancy of the samples (Fig. S5†). According to sequence homology, the differentially expressed genes were categorized into three groups: biological process, cellular component, and molecular function.

In the biological process group, the enriched GO terms for downregulated genes included 17 subcategories. The maximum number of genes in one subcategory was found in the oxidationreduction process, which included 25 genes that were significantly altered. Furthermore, the fatty acid metabolic process (involving eight genes) and mono-carboxylic-acid metabolic Table 1 The significantly enriched KEGG pathways (metabolism) of differentially expressed genes between BW25113 and BW25113-ΔastE

KEGG pathway	Gene name	Description	Corrected <i>P</i> -value ^{<i>a</i>}	log 2-fold change ^b
Transporters	fecC	Ferric citrate ABC	1.70×10^{-4}	+2.3
	shn	Sulfate transporter subunit	3.10×10^{-3}	-1.2
	fecB	Ferric citrate ABC	3.13×10^{-9}	+1.8
	5002	transporter periplasmic		
	cusII	Sulfate/thiosulfate ABC	1.22×10^{-6}	_11
	0,000	transporter permease	1.22 / 10	1.1
	cysW	Sulfate/thiosulfate ABC transporter permease	1.25×10^{-6}	-1.1
	araG	L-arabinose ABC transporter ATPase	3.28×10^{-2}	-1.0
	znuC	Zinc ABC transporter ATPase	$5.11 imes 10^{-3}$	+1.0
	ddA	D% 2CD-dipeptide ABC transporter periplasmic	8.55×10^{-4}	-1.3
		binding protein		
	fadL	Long-chain fatty acid outer membrane transporter	1.51×10^{-13}	-1.2
	psuT	Putative nucleoside transporter	3.76×10^{-5}	+1.3
	ybaT	Putative amino acid	3.13×10^{-9}	+1.1
	lplT	Lysophospholipid transporter	1.90×10^{-5}	+1.0
	ynfM	Putative arabinose efflux transporter	1.17×10^{-12}	-1.2
	uhpT	Hexose phosphate transporter	$1.19 imes10^{-3}$	-1.2
	yihO	Putative sulphoquinovose	4.87×10^{-2}	-1.4
	narU	Nitrate/nitrite transporter	$6.34 imes10^{-4}$	-1.5
Membrane	yhiI	Putative membrane fusion protein (MFP) of efflux pump	1.85×10^{-4}	+1.1
	frdC	Fumarate reductase (anaerobic)% 2C membrane anchor subunit	1.79×10^{-3}	+1.3
	mdtM	Multidrug efflux system	6.48×10^{-5}	+1.5
	fecA	TonB-dependent outer membrane ferric citrate	4.57×10^{-12}	+1.5
		transporter and signal transducer% 3B ferric citrate extracellular receptor% 3B		
	yjhB	FecR-interacting protein Putative MFS transporter%	3.24×10^{-2}	-1.5
	slp	Outer membrane	3.93×10^{-10}	+1.0
	nlpA	lipoprotein Cytoplasmic membrane	1.79×10^{-4}	-1.5
	ubiI	hpoprotein-28 2-Octaprenylphenol hydroxylase% 2C FAD-	6.13×10^{-3}	+1.0
	chbC	dependent N% 2CN'-diacetylchitobiose- specific enzyme IIC	1.34×10^{-10}	-1.9
	cysN	component of PI'S Sulfate adenylyltransferase% 2C subunit 1	3.98×10^{-6}	-1.4
	yjhB	Putative MFS transporter% 2C membrane protein	3.24×10^{-2}	-1.5
Transcriptional	yebK	Putative DNA-binding transcriptional regulator	1.74×10^{-6}	-1.2

KEGG pathway	Gene name	Description	Corrected <i>P</i> -value ^{<i>a</i>}	log 2-fold change ^b
	cbl	ssuEADCB/tauABCD operon	3.91×10^{-2}	-1.1
Pyruvate metabolism	aldR	Aldebyde debydrogenase B	1.38×10^{-10}	-1.0
1 ji di dece interdati ontoni	frdC	Fumarate reductase	1.79×10^{-3}	+1 3
	Jiuo	(anaerobic)% 2C membrane	1.79 ~ 10	1.5
Propanoate metabolism	nrnF	Propionate-CoA ligase	1.33×10^{-13}	-3.0
	prpL prnD	2-Methyleitrate debydratase	1.33×10^{-7}	-5.0
	pipD nmB	2-Methylcrate denydratase	3.84×10^{-32}	-4.0
	ргрв	2-Methylisocitrate lyase	3.82×10 1.10 × 10 ⁻¹⁰	-3.0
	prpC	2-Methylchrate synthase	1.10×10	-3.8
	Juub	epimerase/delta(3)- <i>cis</i> - delta(2)- <i>trans</i> -enoyl-CoA isomerase/enoyl-CoA hydratase/3-hydroxyacyl-CoA	2.47 × 10	-1.1
		dehydrogenase		
Sulfur metabolism	cysN	Sulfate	3.98×10^{-6}	-1.4
		adenylyltransferase% 2C subunit 1		
	cysC	Adenosine 5′- phosphosulfate kinase	2.76×10^{-3}	-1.3
	cysD	Sulfate adenylyltransferase% 2C subunit 2	3.33×10^{-7}	-1.8
	cysU	Sulfate/thiosulfate ABC transporter permease	1.22×10^{-6}	-1.1
	cysW	Sulfate/thiosulfate ABC transporter permease	1.25×10^{-6}	-1.0
Nucleotide metabolism	cysH	Phosphoadenosine phosphosulfate reductase% 3B PAPS reductase% 2C this reductase dent	3.75×10^{-6}	-1.4
	cysI	Sulfite reductase% 2C beta subunit% 2C NAD(P)-	1.07×10^{-15}	-1.5
	cysJ	Sulfite reductase% 2C alpha	3.22×10^{-6}	-1.3
	chn	Sulfate transporter subunit	3.10×10^{-3}	_1 2
	sop cusN	Sulfate	3.10×10^{-6}	1.2
	USIV	adenylyltransferase% 2C subunit 1	5.56 × 10	-1.4
	cysC	Adenosine 5'- phosphosulfate kinase	2.76×10^{-3}	-1.3
	cysD	Sulfate adenylyltransferase% 2C	3.33×10^{-7}	-1.8
		subunit 2		
	рѕик	Pseudouridine kinase	$5.8/ \times 10^{\circ}$	+3./
	psuG	Pseudouridine 5'-phosphate glycosidase	4.35×10^{-10}	+2.7
Acid resistance	gadC	glutamate:gamma- aminobutyric acid antiporter	$1.49 imes 10^{-5}$	+0.9
	patD	Gamma- aminobutyraldehyde dehydrogenase	2.05×10^{-3}	-0.7
	ariR	RcsB connector protein for regulation of biofilm and acid-resistance	1.68×10^{-73}	-3.3
	<i>ybaS</i>	Glutaminase 1	$1.03 imes10^{-7}$	+1.1

 a A hypergeometric test was used for statistical analysis, and *P*-values have been corrected for multiple testing by the Benjamini and Hochberg adjustment method. A corrected *P* value of <0.05 is considered statistically significant. b log 2-fold change of differential expression; "+" means up-regulated genes, "-" means down-regulated genes.

process (involving 10 genes) were also significantly downregulated. However, only 10 subcategories were found in the enriched GO terms for upregulated genes. Of these subcategories, the oxidation-reduction process, mono-carboxylic-acid metabolic process, and glutamine metabolic process were significantly upregulated, involving 12, 3, and 2 genes, respectively.

Additionally, the cellular component category, which included five subcategories, was also found in the enriched GO terms for different regulated genes. The altered genes related to cytoplasmic parts was significantly upregulated, involving 8 genes. Interestingly, in the molecular function group, upregulated genes were found in four subcategories: rRNA binding, squalene monooxygenase activity, oxidoreductase activity, and malate dehydrogenase activity. However, downregulated genes were only identified in three subcategories: oxidoreductase activity, malate dehydrogenase activity, and acyl-CoA dehydrogenase activity, and only a few genes were involved.

KEGG enrichment analysis of differentially expressed genes

To identify the function of differentially expressed genes, an enrichment analysis for specific metabolic pathways was carried out using the KEGG database. Differentially expressed genes were mainly involved in carbohydrate metabolism, energy metabolism, amino acid metabolism, ABC transporters, and microbial metabolism in diverse environments (Fig. S6†). As shown in Table 1, to adapt to butanol stress, most genes (such as *cysH*, *cysC*, *cysN*, *cysJ*, and *cysD*) involved in sulfur metabolism were downregulated by more than twofold (Fig. S7a†). In addition, genes related to propanoate metabolism were also significantly downregulated by over 8-fold, in which the most change was found in gene *prpD* (encoding 2-methylcitrate dehydratase) of over 16-fold (Fig. S7b†).

In contrast, most genes (such as *yhiI*, *frdC*, *mdtM*, *fecA*, *fecC*, and *fecB*) involved in membrane metabolism and transporters were upregulated by more than twofold. More interestingly, genes, related to nucleotide metabolism *psuK* and *psuG*, and acid-activated glutaminase *ybaS* and the amino acid antiporter *gadC*, were significantly upregulated.

Validation of transcription results by RT-qPCR

Real-time quantitative PCR (RT-qPCR) was performed to confirm the transcription results using differentially expressed genes. A group of 7 upregulated and 13 downregulated genes involving different metabolism pathways was randomly selected for validation. As shown in Fig. 4, the RT-qPCR and transcription analysis results correlated well with each other, and with similar trends and expression levels between the transcription and RTqPCR analysis, indicating that the RNA-Seq results were credible.

Identification of significant metabolites and pathway analysis

To characterize the underlying metabolic impact of knockingout gene *astE* under butanol stress, a hierarchical clustering was conducted to assist in accurately screening for marker metabolites (Fig. S8†), and then the related metabolic processes were investigated by integrating transcriptomics and



Fig. 4 Real-time gPCR analysis of the selected genes.

metabolomics. As shown in Fig. 5, most of the differential metabolites were mainly involved in the L-arginine degradation pathway, sulfate metabolic pathway, and 2-methylcitrate metabolic pathway, and were downregulated to adapt to butanol stress. Unexpectedly, although the L-arginine degradation pathway was suppressed due to the knocking-out of gene *astE*, significantly decreasing most of the intermetabolites, the levels of L-arginine and glutamate did not appear to change, indicating that L-arginine and glutamate played an important role in improving cell robustness.

However, interestingly, the acid-activated glutaminase ybaS and the amino acid antiporter gadC, mainly involved in the glutamine metabolic process, were significantly upregulated (Fig. 6). In the glutamine metabolic process, glutamine is converted into glutamate by ybaS, while NH₃ is generated, which reacts with H⁺ to alkalize the environment. In addition, gadC, a glutamic acid y-aminobutyrate antiporter, is part of the glutamate-dependent acid resistance system 2 (AR2) which confers resistance to extreme acid conditions.17 Insertional inactivation of the gadC gene results in sensitivity to extreme acid conditions (pH 2-3) and eliminates glutamic acid enhancement of acid resistance,18 which is similar to the acid resistance system for E. coli survival under an extremely acidic environment.¹⁹ To further demonstrate the hypothesis, the effect of knocking-out astE on amino acids were also investigated using the amino acid analyzer (Table S2[†]). Compared with strain BW25113, the mutant BW25113-∆astE could significantly enhance the synthesis of some amino acids, such as Met, Val, and Tyr. It is interesting that the concentration of extracellular glutamate and free ammonia (NH₃) in mutant BW25113- Δ astE was significantly increased. Furthermore, the growth experiment under various pH condition in absence of butanol was used to confirm the ability of mutant strain against acid stress. In the acid-stress experiment (Fig. 6), when extracellular pH decreased to 3.0, cell growth effectively improved in mutant BW25113-AastE and biomass reached 0.17 after 24 h of cultivation, which is an increase of 54.5% compared with that of BW25113.

Discussion

During microbial fermentation, microorganisms are usually exposed to fluctuations in solvent concentration, osmotic



Fig. 5 Different metabolites and genes in pathways: L-arginine degradation, sulfate metabolic pathway, and 2-methylcitrate metabolic pathway.

pressure, pH, nutrient availability, and temperature.²⁰ Particularly, in acetone-butanol-ethanol fermentation, butanol is highly toxic to microorganisms due to its hydrophobicity, and severely inhibits cell growth and the production of solvent.²¹ Therefore, understanding the mechanisms involved in the *n*butanol response can help to facilitate the engineering of hosts with improved tolerance.²² In the present study, a butanol tolerance strain, with higher butanol tolerance (increased by 34.6%) than that of the control strain, was successfully engineered by the disruption of gene *astE*.

To elucidate the mechanism of tolerance against butanol stress after deleting gene *astE*, metabolic responses were determined by using gas chromatography-mass spectrometry (GC-MS). As a result, a total of 73 metabolites were significantly changed between BW25113 and BW25113- Δ astE. Most of the differential metabolites were mainly involved in the L-arginine degradation pathway, sulfate metabolic pathway, and 2-meth-ylcitrate metabolic pathway and were downregulated. Unexpectedly, although the L-arginine degradation pathway was suppressed due to the knocking-out of gene *astE*, decreasing most the intermetabolites significantly, the levels of L-arginine and glutamate did not appear to change.

To further elucidate the tolerance mechanism, a transcription analysis was also performed, and found that 311 genes (113 upregulated and 198 downregulated) showed different expression levels, which mainly involved carbohydrate metabolism, energy metabolism, nucleotide metabolism, amino acid metabolism, ABC transporters and microbial metabolism in diverse environments. For example, to adapt to butanol stress,

most genes (such as cysH, cysC, cysN, cysJ, and cysD) involved in sulfur metabolism were downregulated by more than twofold. This was maybe due to the fact that butanol stress could effectively inhibit sulfur assimilation,23 and then further weakened by cysteine biosynthesis and the formation of cytomembrane.²⁴ In contrast, most genes (such as *yhiI*, *frdC*, *mdtM*, *fecA*, fecC, and fecB) involved in membrane metabolism and transporters were upregulated by more than twofold, fecC encodes a hydrophobic inner membrane protein, increased external iron concentration and increased expression of iron transport genes (fecA, fecC, and fecB) improved E. coli butanol tolerance.²⁵ Slp is believed to take part in acid resistance as its expression increased when cells were grown at pH 4.5 and 5.5 under conditions known to induce glutamate-dependent acid resistance.²⁶ YhiI is a predicted bitopic inner membrane protein.²⁷ Overexpression of yhiI leads to abnormal biofilm architecture.28 The *MdtM* proteins a multidrug efflux protein that belongs to the major facilitator superfamily (MFS) of transporters.²⁹

In addition, genes related to transporters were also significantly upregulated by over twofold, suggesting that increased expression of the transporters was a physiological adaptation to stressful environments.³⁰ In general, transporter systems consist of different transmembrane protein components and play roles in bacterial virulence, cell growth and development, and survival under various environments.^{31,32} Genes (such as *psuK* and *psuG*) related to nucleotide metabolism were also significantly upregulated by 6- to 15-fold. In microorganisms, nucleotides are obligatory metabolites and serve an important role in the regulation of numerous cellular processes, including



Fig. 6 The upregulated genes mainly involved in glutamine metabolic process regulated the intracellular pH-homeostasis to adapt to butanol stress: L-glutamine degradation I; L-glutamate:4-aminobutyrate antiporter; impact of different pH stresses; acid resistance system.

cellular energy supply, signaling molecules, and are incorporated into cofactors (*e.g.*, NAD and coenzyme A) and precursors (*e.g.*, UDP-glucose and GDP-mannose).³³ Particularly, nucleotide synthesis is phosphate consuming, involving the regulation of phosphate uptake and utilization,³⁴ which then regulates the plasticity of the cell wall in bacteria.³⁵

More importantly, the genes of *ybaS* and *gadC* involving in glutamine metabolism were significantly up-regulated. In *E. coli, gadC*, as the amino acid transporter protein, was responsible for the exchange of extracellular L-glutamine (Gln) with intracellular L-glutamic acid (Glu), and the glutamine enzyme *ybaS* was responsible for the conversion of Gln into Glu, releasing ammonia. As a result, the free ammonia could be used to neutralize intracellular protons, and then increased the intracellular pH to resist acidic environment. More interestingly, there was a typical acid resistance system in *E. coli* that relies on L-glutamine,¹⁹ in which glutamine is converted to L-glutamate by acid-activated glutaminase *ybaS*, with

concomitant release of gaseous ammonia to neutralize protons, resulting in an elevated intracellular pH under an acidic environment. Therefore, the *ybaS* and the amino acid antiporter *gadC*, which exchanges extracellular glutamine with intracellular glutamate, together constitute an acid resistance system that is sufficient for *E. coli* survival under an extremely acidic environment, which was similar with our resistance system for *E. coli* survival under extreme butanol stress.

To further elucidate the physiological mechanism, the changes of amino acids between strains BW25113 and BW25113- Δ astE were measured by amino acid analyzer. It is interesting that the concentration of extracellular glutamate and free ammonia (NH₃) in mutant BW25113- Δ astE was significantly increased. Furthermore, the mutant BW25113- Δ astE showed the better physiological performance against acid stress. When extracellular pH decreased to 3.0, cell growth of mutant increased 54.5% compared with BW25113. With the integration of amino acids analysis and an acid tolerance

Paper

experiment, it could be concluded that, in the mutant BW25113- Δ astE, butanol can effectively inhibit the activities of membrane-bound ATPases, resulting in a lower internal pH.¹⁰ *YbaS* and *GadC* are activated by acidic pH. *GadC* exchanges intracellular L-glutamate and extracellular glutamine.¹⁹ Upon uptake into *E. coli*, glutamine (Gln) is converted to L-glutamate (Glu) by the acid-activated glutaminase *YbaS*, with concomitant release of gaseous ammonia.¹⁹ Then some of the free ammonia is transferred out of the cell, and some of the free ammonia neutralizes proton, resulting in elevated intracellular pH under acidic, thereby maintaining intracellular pH homeostasis to adapt to butanol stress.

At present, genomic tools (transcriptomics, proteomics, and metabolomics) have been widely used to investigate cellular response under butanol stress, and correspondingly, a series of strategies for improving cellular robustness was elucidated,³⁶ including (i) metabolic detoxification; (ii) heat shock proteins; (iii) the proton motive force and associated energy production; (iv) molecular efflux pumps; (v) the changes of cell membrane composition and biophysics;³⁷ and (vi) other transcriptional responses.^{38,39} However, the molecular mechanism underlying butanol tolerance is still not comprehensively understood for microorganisms, which has made strain improvement difficult due to the complexity of the regulatory network.^{40,41}

Conclusion

In this study, a mutant strain with high butanol tolerance (increased by 34.6%) was constructed by disrupting gene *astE*. To clarify the tolerance mechanism, a transcriptome and metabolome were performed to analyze the differential gene expression and characterize the underlying metabolic impact. As a result, it was found that the mutant BW25113- Δ astE had developed a special tolerance mechanism by regulating intracellular pH-homeostasis to adapt to butanol stress. Our findings indicate the non-negligible impact of pH on microbial butanol tolerance, broadening the understanding on microbial butanol tolerance and also providing a novel strategy for the rational engineering of a more robust butanol-producing host.

In the future, the correlation between cell growth and gene expression under the conditions of butanol and acid stresses would be investigated in detailed, providing some novel pathways for further improving cellular robustness and fermentation performance with metabolic engineering.

Materials and methods

Strains and plasmids

E. coli BW25113 was used as a receptor for gene deletion. More information about the plasmids and strains used in this study are presented in Table S3.[†]

Construction of engineering strains

The primers were designed using *E. coli* k-12 MG1655 as the template and are given in Table S4.† Standard bacterial transformations were performed according to the procedures

described by Sambrook.⁴² In all the cases, PCR was performed using TaKaRa PrimeSTAR® HS (TAKARA Bio Inc., Tokyo Japan). All the genes were sequenced to verify the insert prior to transformations. The engineered strains were transformed using electrical conversion method.⁴³

Culture medium and conditions

Culture medium. Luria–Bertani (LB): 10 g L⁻¹ tryptone, 5 g L⁻¹ yeast extract and 10 g L⁻¹ NaCl;

M9 medium: 6.78 g L⁻¹ Na₂HPO₄, 3.0 g L⁻¹ KH₂PO₄, 0.5 g L⁻¹ NaCl, 1.0 g L⁻¹ NH₄Cl, 4 g L⁻¹ glucose, 0.493 g L⁻¹ MgSO₄·7H₂O, 0.011 g L⁻¹ CaCl₂;

Screened on synthetic complete (SC) medium: 20 g L^{-1} glucose, 7 g L^{-1} urea, 5 g L^{-1} KH₂PO₄, 0.8 g L^{-1} MgSO₄·7H₂O, 3 g L^{-1} sodium acetate, 15 g L^{-1} agar.

Culture conditions. During construction, strains were grown in Luria–Bertani (LB). All engineered strains were screened on synthetic complete (SC) medium;

Butanol and acid tolerance curve of the BW25113 and the mutant BW25113- Δ astE: LB medium with different butanol concentrations or different PH (adjusted pH with 0.1 mol L⁻¹ citric acid and 0.1 mol L⁻¹ sodium citrate buffer) were prepared and inoculated with 10% (v/v) inoculum size of the BW25113 and the mutant BW25113- Δ astE, strains were inoculated and cultured at 37 °C for 24 h. OD₆₀₀ was determined to detect the growth of the strains.

Fermentation was carried out in M9 medium with 0 g L^{-1} and 5 g L^{-1} butanol in a shake-flask culture (200 rpm, 37 °C) with 10% (v/v) of the inoculum size.

Analytical methods

Cell growth was determined by measuring the optical density at 600 nm (OD_{600}) by using a UV-vis spectrophotometer (Beckman Coulter DU800). Cells were incubated to the exponential phase ($OD_{600} = 0.6$), different butanol or pH concentrations were added to the broth (200 rpm, 37 °C), and then the cell concentration was measured to represent cell growth.

Spotting assay for evaluating butanol tolerance

To evaluate microbial tolerance, a spotting assay was applied in the presence of butanol.⁴⁴ First, cells were incubated overnight in LB medium with shaking (200 rpm, 30 °C), and then collected by centrifugation (10 000 × g, 20 s), resuspended with sterilized water. Second, the suspension was serially diluted to an OD₆₀₀ of 1×10^{-1} , 1×10^{-2} , 1×10^{-3} , 1×10^{-4} , 1×10^{-5} , 1×10^{-6} , and then spotted (5 µL each) onto agar plates containing different butanol concentrations. The plates were sealed with vinyl plastic tape to prevent evaporation of butanol and incubated at 37 °C.

GC-MS-based metabolomics analysis

For metabolomics analysis, cells were collected after cultivating 12 h under different butanol concentration. For each sample, broth equivalent to 10^8 cells was collected by centrifugation at

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 $8000 \times g$ for 10 min at 4 °C. The pellets were immediately frozen by liquid nitrogen and then stored at -80 °C before using.

The metabolomics analysis protocol was performed according to the following. (i) Sample preparation. Cells were thawed and resuspended in 5 mL of cold 2:2:1 (v/v/v) acetonitrile/ methanol/H2O solution, and then vibrated with vortex oscillation blender for 1 min. Subsequently, cells were frozen in liquid nitrogen and thawed for five times. Supernatants were collected by centrifugation at 14 000 \times g for 3 min at 4 °C and dried by vacuum centrifugation. (ii) Sample derivatization. The samples were dissolved in 40 µL of 98% methoxyamine hydrochloride (40 mg mL⁻¹ in pyridine), after shaking at 30 °C, 180 rpm for 90 min, 180 µL of N-methyl-N-(trimethylsilyl)trifluoroacetamide (MSTFA) was added and incubated at 37 °C, 180 rpm for 30 min to trimethylsilylate the polar functional groups. The derivates were then collected by centrifugation at 1, 4000 \times g for 3 min, and the supernatant was used directly for GC/MS analysis. (iii) GC-MS analysis. The analysis was performed on an Agilent 5977E GC/MSD equipped with an HP-5MS capillary column (30 m \times 250 mm \times 0.25 µm), with 70 eV of electron impact ionization, in which 0.2 µL of sample was injected in splitless mode at 230 °C with a constant flow of 1 mL min⁻¹ helium. The temperature program started isocratic at 45 °C for 2 min, followed by temperature ramping of 2 °C min to a final temperature of 250 °C, and then held constant for an additional 2 min (solvent delay: 6 min, scanning rate: 1250, and scanning method: full scanning). The range of mass scan was m/z 38–650. (iv) Data processing and statistical analysis. To identify the compounds, the mass fragmentation spectrum was analyzed using an automated mass spectral deconvolution and identification system (AMDIS)32 and the data was matched with the Fiehn Library 41 and the mass spectral library of the National Institute of Standards and Technology (NIST). Peak areas of all identified metabolites were normalized against the internal standard and the acquired relative abundances for each identified metabolite were used for data analysis. All metabolomics profile data were first normalized by the internal control and the cell numbers of the samples and then analyzed using xcms online and metaboanalyst.

Transcriptional analysis45

The cells of BW25113 and BW25113- Δ astE were collected after cultivating for 12 h under 0 g L⁻¹ and 5 g L⁻¹ butanol condition, washed with sterile water three times, then frozen in liquid nitrogen and stored at 80 °C for further RNA-seq and RT-qPCR analyses.

RNA extraction and examination

The total RNA was extracted by grinding the cells in TRIzol reagent (TaKaRa, Japan) in liquid nitrogen, isolated with chloroform, and precipitated with isopropanol, and then the sediment was washed with 75% alcohol and dissolved in RNA-free distilled water. RNA degradation was detected on 1% agarose gels, and the purity was checked using a Nanodrop 2000C spectrophotometer. Furthermore, the concentration of RNA was measured using an RNA Assay Kit in Qubit (Life Technologies,

CA, USA). The RNA integrity was assessed using an RNA Nano 6000 Assay Kit with the Bioanalyzer 2100 system (Agilent Technologies, CA, USA).

Library preparation and transcriptome sequencing

A total amount of 3 µg RNA per sample was used as input material for the RNA sample preparations. Sequencing libraries were generated using the NEBNext Ultra™ RNA Library Prep Kit from Illumina (NEB, USA). First, the mRNA was separated from the total RNA using poly-T oligo-attached magnetic beads and fragmented using divalent cations under elevated temperature in NEBNext First-Strand Synthesis Reaction Buffer $(5\times)$. Second, the mRNA was reverse transcribed to synthesize double-stranded cDNA using random hexamer primers. Third, the base A and an adaptor with a hairpin loop structure were added to the 30 ends of the cDNA to prepare for hybridization. The library fragments were purified with the AMPure XP system (Beckman Coulter, Beverly, USA) to select fragments 150-200 bp in length. Then, PCR was performed with universal PCR primers and an Index (X) Primer. Finally, the PCR products were purified (AMPure XP system) and the library quality was assessed using the Agilent Bioanalyzer 2100 system. The library was loaded into a flow cell and the fragments hybridized to the flow cell surface. Each bound fragment was amplified into a clonal cluster through bridge amplification. Sequencing reagents including fluorescently labeled nucleotides were added and the first base was incorporated. The flow cell was imaged and the emission from each cluster was recorded. The emission wavelength and intensity were used to identify the base. This cycle was repeated 150 times to create a read length of 150 bases on an Illumina HiSeq platform at Novogene Bioinformatics Institute (Novogene, Beijing, China).

Data analysis

Differential expression analysis of BW25113 and BW25113- Δ astE (three biological replicates under 5 g L⁻¹ butanol stress) was performed using the DESeq R package (1.18.0). The resulting *P*-values were adjusted using the Benjamini and Hochberg's approach for controlling the false discovery rate. Genes with an adjusted *P*-value < 0.05 found by DESeq were assigned as differentially expressed.

GO and KEGG enrichment analysis of differentially expressed genes

GO enrichment analysis of differentially expressed genes was implemented by the GOseq R package, in which gene length bias was corrected. GO terms with corrected *P*-value less than 0.05 were considered significantly enriched in differentially expressed genes.

KEGG is a database resource for understanding high level functions and utilities of the biological system, such as the cell, the organism, and the ecosystem, from molecular-level information, especially large-scale molecular datasets generated by genome sequencing and other high-throughput experimental technologies (http://www.genome.jp/kegg/). We used KOBAS software to test the statistical enrichment of differential expression genes in KEGG pathways. We then added the differentially expressed gene accession ID to the xcms online (https://xcmsonline.scripps.edu) multiomics lookup database for integrating metabolic data to identify key differences.

Real-time quantitative PCR (RT-qPCR)

The expression levels of seven upregulated and 13 downregulated genes in two strains, which were selected randomly from different metabolism pathways, were detected by qPCR. Total RNA isolation was carried out by using an RNAprep pure Kit, and reverse transcription (cDNA synthesis) was performed according to the protocol of the PrimeScript® RT reagent kit Perfect Real Time (Takara Bio Inc, Shiga, Japan). Quantitative real-time PCR (RT-qPCR) was done using the *rrsG* gene as the internal control, and the primers used in qPCR are given in Table S5.† Each sample was tested in triplicate in a 96-well plate (Bio-Rad).

Amino acid determination

When the cells had been incubated to the exponential phase $(OD_{600} = 0.6)$, 5 g L⁻¹ of butanol was added and cultivation was proceeded for 12 h (200 rpm, 37 °C). Cells were then collected and washed three times with sterilized water. Afterward, the concentration of amino acids from the collected cells and fermentation supernate were detected by a high speed amino acid analyzer (Hitachi L-8900, Japan), in which the test parameters were: sodium ion exchange column, visible light detector, separation column temperature for 57 °C, and a reaction column temperature of 135 °C.

Consent for publication

All authors agreed to publish this article.

Authors' contributions

YG and HP conceived of the study; YG, HP and LHL drafted the manuscript; YG, BL, DWB and CHT carried out the study; YG and ZKZ data analysis. All the authors read and approved the final manuscript.

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Conflicts of interest

The authors declared that they have no competing interests.

Abbreviations

GO	Gene ontology
KEGG	Kyoto encyclopedia of genes and genomes
RT-PCR	Reverse-transcription PCR
GC-MS	Gas chromatography-mass spectrometry

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