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Douglas McCloskey, Sibei Xu, Troy E. Sandberg, Elizabeth Brunk, Ying Hefner, Richard Szubin, Adam M. Feist, Bernhard O. Palsson

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Adaptation to the coupling of glycolysis to toxic methylglyoxal production in tpiA deletion strains of *Escherichia coli* requires synchronized and counterintuitive genetic changes.\textsuperscript{1}

Douglas McCloskey\textsuperscript{1,2}, Sibei Xu\textsuperscript{1}, Troy E. Sandberg\textsuperscript{1}, Elizabeth Brunk\textsuperscript{1}, Ying Hefner\textsuperscript{1}, Richard Szubin\textsuperscript{1}, Adam M. Feist\textsuperscript{1,2}, and Bernhard O. Palsson\textsuperscript{1,2,*}

\textsuperscript{1}Department of Bioengineering, University of California - San Diego, La Jolla, CA 92093, USA.

\textsuperscript{2}Novo Nordisk Foundation Center for Biosustainability, Technical University of Denmark, 2800 Lyngby, Denmark.

*Corresponding author, Department of Bioengineering, University of California, San Diego, 9500 Gilman Drive, La Jolla, CA 92093-0412, USA. Tel.: 858-534-5668; Fax: 858-822-3120. palsson@ucsd.edu

**Abstract**

Methylglyoxal is a highly toxic metabolite that can be produced in all living organisms. Methylglyoxal was artificially elevated by removal of the tpiA gene from a growth optimized *Escherichia coli* strain. The initial response to elevated methylglyoxal and its toxicity was characterized, and detoxification mechanisms were studied using adaptive laboratory evolution. We found that: 1) Multi-omics analysis revealed biological consequences of methylglyoxal toxicity, which included attack on macromolecules including DNA and RNA and perturbation of nucleotide levels; 2) Counter-intuitive cross-talk between carbon starvation and inorganic phosphate signalling was revealed in the tpiA deletion strain that required mutations in inorganic phosphate signalling mechanisms to alleviate; and 3) The split flux through lower glycolysis depleted glycolytic intermediates requiring a host of synchronized and coordinated mutations in non-intuitive network locations in order to re-adjust the metabolic flux map to achieve optimal

\textsuperscript{1} Adaptive laboratory evolution, tpiA gene knockout, mutation analysis, multi-omics data integration, systems biology, *E. coli*
growth. Such mutations included a systematic inactivation of the Phosphotransferase System (PTS) and alterations in cell wall biosynthesis enzyme activity. This study demonstrated that deletion of major metabolic genes followed by ALE was a productive approach to gain novel insight into the systems biology underlying optimal phenotypic states.

Introduction

Methylglyoxal (mthgxl) is a highly toxic metabolite that can be produced in all living organisms, and is directly linked to sugar metabolism. Methylglyoxal is highly cytotoxic owing to keto and aldehyde groups that can readily react with nearby metabolites and exposed units of macromolecules. For example, the keto and aldehyde groups can bind covalently to macromolecules such as DNA and proteins to form advanced glycation end products (AGEs) (Basta et al., 2004; Goldin et al., 2006; Singh et al., 2001; Thomas, 2011). Methylglyoxal is also mutagenic, and has been shown to interact with DNA through direct binding of guanine residues (Kasai et al., 1982; Murata-Kamiya et al., 1999). Due to the potential for methylglyoxal generation in all living organisms and its toxic effects to the cell, a greater understanding of its toxicity and mechanisms of detoxification are needed.

Methylglyoxal can be formed as a side product of glycolysis via the methylglyoxal pathway (Saadat and Harrison, 1999; Weber et al., 2005). Methylglyoxal levels can be artificially increased by bifurcating flux through lower glycolysis by removing the tpiA gene. tpiA encodes the Triosephosphate Isomerase (TPI) that is responsible for isomerizing dihydroxyacetone phosphate (dhap) to glyceraldehyde-3-phosphate (g3p). Loss of TPI forces dhap to be metabolized by the methylglyoxal pathway instead of the enzymes of lower glycolysis, which generates methylglyoxal. Removal of tpiA has been used in biotechnology applications to increase production of 1,2-propanediol, which has high commercial utility in the production of
polymers (Shelley, 2007). Additional studies have also identified genes in potassium transport that allow for cells to better cope with exposure to methyglyoxal (Ozyamak et al., 2013). Further studies that both reveal genetic interventions for how to optimize flux through the methylglyoxal pathway and allow the cell to rapidly detoxify methylglyoxal are needed for efficient production of methylglyoxal derived compounds.

Flux through the methylglyoxal pathway for production of methylglyoxal derived compounds for biotechnology applications can be optimized by adaptive laboratory evolution (ALE) (Fong et al., 2006). ALE is an experimental approach that introduces a selection pressure (e.g., growth rate selection) in a controlled environmental setting (Dragosits and Mattanovich, 2013; Plucain et al., 2014; Tenaillon et al., 2016). Using ALE, organisms can be perturbed from their evolutionary optimized homeostatic states, and their re-adjustment to a new growth condition can be studied. Monitoring cellular changes during the course of adaptation can reveal novel and non-intuitive component functions and interactions that can then be re-engineered into a host organism. ALE of tpiA mutants found an increase in flux capacity through the methylglyoxal pathway (Fong et al., 2006), however, a systems level understanding of methylglyoxal detoxification as a result of mutations that appeared during ALE were not revealed.

In this study, the effects of methyglyoxal toxicity on cellular functions were investigated by removing the tpiA gene from a growth optimized E. coli strain (LaCroix et al., 2015). The consequences of increased level of methyglyoxal was studied in the tpiA knock-out strains, that was then evolved to tolerance using ALE. Multi-omic data analysis revealed biochemical evidence of methyglyoxal toxicity and genetic changes that upregulated mechanisms to overcome its toxic effects.
Results

tpiA KO

In order to eliminate the confounding variable of adaptation to the growth conditions of the experiment, a starting strain in which the gene KO was implemented was carefully chosen. A wild-type *E. coli* K-12 MG1655 strain previously evolved under glucose minimal media at 37°C (LaCroix et al., 2015) (denoted as “Ref”) was selected in order to minimize biological changes caused by adaption to the loss of a gene product from those caused by adaption to the growth conditions of the experiment.

TPI (*tpiA*, Triosphosphate Isomerase) was removed from Ref to generate strain uTpiA (denoted “unevolved *tpiA* knockout strain”). The loss of *tpiA* resulted in an 80% loss in growth rate (Fig. 1D). Four uTpiA independently-inoculated starting cultures were simultaneously evolved on glucose minimal media at 37°C in an automated ALE platform (LaCroix et al., 2015; Sandberg et al., 2014) to generate endpoint strains, denoted “evolved *tpiA* knockout strains” or “eTpiA”. A statistically significant increase in final fitness (Student’s t-test, pvalue<0.05) was found in all ALE endpoints of the eTpiA lineages (ave±stdev 164±7% increase in fitness) compared to uTpiA.

Loss of *tpiA* activated the glyoxylate pathways

Fructose 1,6-bisphosphate (fdp) from upper glycolysis is split into dihydroxyacetone phosphate (dhap) and glyceraldehyde-3-phosphate (g3p) by fructose-bisphosphate aldolase (FBA) (Fig. 2A). g3p can be directly processed down the enzymatic machinery of lower glycolysis, but dhap must first be isomerized by Triosephosphate Isomerase (TPI) to g3p before glyceraldehyde...
phosphate dehydrogenase (GAPD) can catalyze the first step of lower glycolysis (Fig. 2A). Loss of enzymatic interconversion by TPI thus bifurcates glycolysis into lower glycolysis and pathways that degrade dhap, that involve methylglyoxal formation.

Reaction fluxes estimated from C\textsuperscript{13} labeling experiments and estimated using Genome-scale Metabolic Flux Analysis (MFA) (McCloskey et al., 2016a, 2016b) found that upper glycolytic flux was approximately evenly split between lower glycolysis and the methylglyoxal pathway in uTpiA (Fig 2B). LC-MS/MS-based metabolomics (McCloskey et al., 2015, 2014) found that bifurcation of lower glycolysis also led to an approximate 3.55 log2 fold change buildup of dhap in uTpiA compared to Ref (Table S3). While not measured directly, LC-MS/MS analysis was able to measure the amounts of methylglyoxal break down products, which indicate an elevation in methylglyoxal (Fig. S2). An end product of the methylglyoxal pathway is D/L-Lactate (lac-DL), which can then be converted back to pyruvate (pyr), the end product of lower glycolysis, by lactate dehydrogenase (LDH) (Fig. 2C). Minimal amounts of intracellular lac-DL and no detectable lac-DL excretion coupled with overexpression of \textit{lldD} and 1.67 mmol*gDCW\textsuperscript{-1}*hr\textsuperscript{-1} flux through LDH (approximately the same amount of flux through MGSA), indicated that the majority of lac-DL produced was converted back to pyr in uTpiA (Fig. 2D-E).

\textit{E. coli} has four paths (Fig. 3A) to metabolize methylglyoxal: paths 1 and 2 lead to the production of propane 1,2-diol (12ppd-R), and paths 3 and 4 lead to the production of lac-DL. Unlike lac-DL, \textit{E. coli} has no known enzyme to process 12ppd-R and hence it can only be excreted out of the cell. RNA sequencing revealed that in uTpiA paths 1 and 2 were primarily activated with minor contribution from path 3 and no contribution from path 4 (Fig. 3B). In contrast, path 4 was highly activated in all eTpiA strains with varying degrees of path 2 and path 3 activation. Based on expression profiling and metabolomics measurements, it appears that the majority of dhap was metabolized to 12ppd-R in uTpiA compared to lac-DL in eTpiA strains.
Known mechanisms of methylglyoxal toxicity

Mechanisms of methylglyoxal toxicity are known and enable deeper interpretation of the multi-omic data set generated (discussed in the sections below). We thus describe them briefly. Methylglyoxal is capable of attacking exposed nucleotides of single stranded RNA or DNA (Krymkiewicz, 1973), particularly at guanosine bases (Broude and Budowsky, 1971) (Fig. 4B). This toxicity has been shown to severely inhibit DNA replication through methylglyoxal cross links formed between guanosine residues of DNA and lysine and/or cysteine residues of the DNA polymerase (Murata-Kamiya and Kamiya, 2001). Methylglyoxal is also capable of oxidizing amino acids of RNA or DNA via Fenton’s reaction (Imlay and Linn, 1988; Kang, 2003).

Methylglyoxal damage is not limited to nucleotides, but extends to proteins and peptides. The thiol of cysteine groups is susceptible to form methylglyoxal induced hemithioacetal adduct cross links with nearby peptide strands or glutathione (Ridderström and Mannervik, 2002) (Fig. 4C). Methylglyoxal can form lysine-lysine cross links via the Maillard Reaction (Nagaraj et al., 1996) (Fig. 4D). In addition to lysine, methylglyoxal has been shown to bind and modify arginine and cysteine residues (Lo et al., 1994). The glycation of lysine in particular has been shown to generate superoxide anions that are then capable of causing additional damage (Kang, 2003) (Fig. 4D). The mechanisms by which methylglyoxal damages proteins and peptides has been shown to be capable of inhibiting enzymatic function \textit{in vitro} and \textit{in vivo} (Kalapos, 1994).

ROS generated through Fenton Reactions involving methylglyoxal can lead to DNA strand breaks (Fig. 4E). 2-phosphoglycolyate is formed during repair of DNA strand breaks (Fig. 4F). 2-phosphoglycolyrate can then be broken down by phosphoglycolate phosphatase (Gph) to
glycolate (glyclt) and inorganic phosphate (pi). In addition, methylglyoxal forms abducts with dATP and dGTP (Frischmann et al., 2005; Papoulis et al., 1995).

Elevated methylglyoxal induced global stress responses

Global stress responders were activated in uTpiA (Fig. 4H). Expression of rpoS was significantly increased in uTpiA compared to Ref (Fig. 4H). rpoS encodes an alternative sigma factor to rpoD that acts as a master regulator of general stress response that is itself induced during times of stress (see (Landini et al., 2014) for a review). Many universal response genes were also activated. uspB, uspD, uspF, and uspG were all significantly upregulated in uTpiA compared to Ref (Fig. 4H).

Elevated methylglyoxal activated glycation responses

Evidence for methylglyoxal damage to and glycation of DNA was revealed by expression profiling (Fig. 4H). Numerous chromosomal toxin-antitoxin (TA) pairs were found to be activated in uTpiA. Chromosomal TA systems respond to DNA damage and nutrient limitation by suppressing protein synthesis and promoting cell motility and biofilm formation (Magnuson, 2007; Ramisetty and Santhosh, 2017). Activation of chromosomal TA pairs have been shown to be associated with persister phenotypes (Kim and Wood, 2010; Luidalepp et al., 2011). TA pair mqsR/mqsA was significantly upregulated in uTpiA compared to Ref (Fig. 4H). mqsR inhibits protein synthesis by cleaving mRNA at GCU sites (Yamaguchi et al., 2009). Overexpression of mqsR induces expression of cspD, which was also significantly upregulated in uTpiA (Kim et al., 2010). cspD inhibits DNA replication and increases persister cell formation (Kim and Wood, 2010). TA pair yafQ/dinJ was significantly upregulated in uTpiA compared to Ref (Fig. 4H). yafQ inhibits protein synthesis by interacting with the 50S subunit of the ribosome where it cleaves mRNAs at AAA codons (Motiejūnaitė et al., 2007; Prysak et al., 2009).
Elevated methylglyoxal activated ROS responses

Evidence for iron oxidation caused by ROS attack via Fenton’s reaction on iron-sulfur clusters was found in the transcriptomics data. Evidence included shifts in the primary ribonucleotide reductases. *E. coli* has two ribonucleotide diphosphate reductases that are encoded by *nrdAB* and *nrdEF* that operate under aerobic conditions and a ribonucleotide triphosphate reductase encoded by *nrdDG* that operates under anaerobic conditions (Brown et al., 1969; Hantke, 1988). The ribonucleotide di-/tri-phosphate reductases act to generate deoxynucleotides for DNA and RNA synthesis (Torrents, 2014). Of the aerobic ribonucleotide reductases, *nrdEF* expression is induced under oxidative stress when iron is unavailable to activate *nrdAB* (Martin and Imlay, 2011; Monje-Casas et al., 2001). The entire *nrdHIEF* operon was significantly upregulated in uTpiA compared to Ref (and eTpiA strains) (Fig. 4H), while *nrdA* was significantly downregulated in uTpiA compared to Ref. Fur appeared to be activated based on expression profiles of the small regulatory RNA *rhyB*. *rhyB* was upregulated in uTpiA compared to Ref (and eTpiA strains).

The manganese importer encoded by *mntH* was upregulated in uTpiA compared to ref (and eTpiA strains) (Fig. 4H). *mntH* is regulated by OxyR, Fur, and MntR, and its expression is induced under iron starvation (Anjem et al., 2009). All genes of the *sufABCDSE* operon except *sufC* and *sufS* were upregulated in uTpiA compared to Ref (and eTpiA strains). The genes of the *sufABCDSE* operon encode enzymes that compose an alternative Fe-S cluster biogenesis pathways that is activated in times of oxidative stress and iron starvation (Lee et al., 2004; Outten et al., 2004). In contrast, the primary Fe-S cluster biogenesis pathway encoded by the ICS system was upregulated in eTpiA strains (Fig. 4H. Table S4) (Kim et al., 2014; Roche et al., 2015; Tokumoto et al., 2002).
Elevated methylglyoxal activated nucleotide synthesis responses

Expression of nucleotide synthesis and salvage pathways were massively perturbed in uTpiA (Table S5, pvalue < 0.05, topGO, ParentChild, Fischer). The majority of nucleotide synthesis and salvage pathways were significantly down-regulated in uTpiA compared to Ref (and eTpiA strains) (Table S4). Metabolomics also revealed massive perturbations in nucleotides, nucleosides, and nucleobases. In fact, almost all nucleotides, nucleosides, and nucleobases assayed were significantly changed in uTpiA compared to Ref (and eTpiA strains) (Fig. 4G, Table S3). For example, a significant drop in ATP (-0.68 log2 fold change compared to Ref) resulted in a significant drop in the Energy Charge ratio in uTpiA (from 0.90 in Ref to 0.82) (Table S3). In addition, nucleotide triphosphates GTP, ITP, UTP, dTTP, and dUTP were significantly depleted; nucleotide diphosphates GDP, UDP, and dADP were significantly perturbed; nucleotide monophosphates CMP, dIMP, and dCMP were significantly changed; and nucleobases nucleotides Adenine, Guanosine, Hypoxanthine, Inosine, and Uridine were significantly perturbed in uTpiA compared to Ref (Fig. 4G, Table S3). These large changes in the nucleotide pools at all levels of phosphorylation could be indicative of perturbations of not just de novo synthesis and salvage, but also potentially break down from DNA, RNA, or direct attack by methylglyoxal.

Taken together, elevated methylglyoxal levels activated a number of stress responses that are consistent with reported detoxification responses. These responses were characterized in more detail here than in previous studies, and not only did they reveal mechanistic details and the global nature of the response, but their characterization provided the necessary starting point for the interpretation of the adjustments made during the ensuing ALE experiment.
ALE alleviated glyoxalase repression increasing flux through the methylglyoxal pathway

Path 4 relies on the Glyoxalase enzymes encoded by *gloAB* (Fig. 3A, Fig 5). Glyoxalase 1 binds methylglyoxal with Reduced Glutathione (gthrd) to form S-D-lactoyl-glutathione (lgt-S) (Thornalley, 2003) (Fig. 3A). Glyoxalase 2 then regenerates gthrd from lgt-S while enzymatically generating lac-D (Vander Jagt, 1993) (Fig. 3A). Expression of *gloA* is repressed by NemR (Fig. 5A-B); repression is enhanced by elevated levels of mthgxy1 (Ozyamak et al., 2013; Umezawa et al., 2008) (Fig. 5C). A mutation in the *nemR* promoter region in eTpiA03, and mutations in the *nemR* small-molecule-binding domain or tetramerization regions in eTpiA01, 02, and 04 were found to offset NemR repression, and allowed for increased expression of *gloA* (Fig. 4A-D).

The increased expression of *gloA* appeared to have provided a fitness advantage by increasing the flux capacity of the methylglyoxal pathway (Fig. 2B). The enhanced conversion of dhap to lac-DL and subsequent conversion of lac-DL to pyr (Fig. 2E) substantially increased the levels of both lac-DL and pyr in eTpiA strains. The levels of lac-DL were 8.65, 8.37, 8.34, and 8.72 log2 fold change in eTpiA strains, respectively, compared to uTpiA, while levels of pyr were 10.39, 10.12, 10.08, and 10.47 log2 fold change in eTpiA strains, respectively, compared to uTpiA. In particular, the levels of pyr were severely depleted in uTpiA due to the forced bifurcation of flux that resulted from the loss of the TPI. The levels of pyr were -4.51 log2 fold change less in uTpiA compared to Ref. Unconverted lac-DL was excreted by the eTpiA strains (Fig. 1 D). In addition, the increased activity of the Glyoxylase system also led also to an elevation in methylglyoxal derived detox products (Fig S2).
Transcription factor cross-talk induced expression of phosphate starvation pathways

Phosphate import in *E. coli* is tightly regulated by the PhoB/PhoR two component system (TCS) (Hsieh and Wanner, 2010) (Fig. 6A). Signalling by the PhoBR TCS involves the high affinity ABC phosphate transporter encoded by the *pstSCAB* operon and a regulatory protein encoded by *phoU* (Aguena et al., 2002; Gardner et al., 2014) that coordinate phosphate import with phosphate metabolism (Fig. 6A). When extracellular phosphate levels are high, phosphorylation and activation of PhoB by PhoR is inhibited by PhoU; when extracellular phosphate levels are low, PhoR phosphorylates and activates PhoB (Fig. 6A). Phosphorylated PhoB then activates numerous operons believed to be involved in phosphate metabolism (Baek and Lee, 2007; Hengge, 2009; Yang et al., 2012). Among these is the *phn* operon that encodes genes required to metabolize methylphosphonate (Kamat et al., 2011; Metcalf and Wanner, 1993), which provides a source of phosphate under conditions of phosphate starvation (Metcalf and Wanner, 1993).

The majority of phoB controlled operons were upregulated in uTpiA compared to Ref (Fig. 6H, Table S4). The genes of the *phn* operon were massively upregulated in uTpiA. A majority of other experimentally verified genes controlled by *phoB* were also significantly differentially expressed in uTpiA (Fig. 6H, Table S4). Given the tightly controlled amount of phosphate in the cultivation medium, lack of phosphate would seem improbable. However, cross-talk between other histidine kinases including CreC and KpdD have been experimentally verified (Wanner, 1993). Targets of CreB, the TF pair of the CreCB TCS, including all genes of the *malEFG* operon, *recG, spoT, trmH, recG*, and *talA* were all differentially expressed in uTpiA compared to Ref (Table S4). The only known target of KpdE, the *kdpFABC* operon was not differentially
expressed. KpdE is the TF of the KpdDE TCS. Most probable given the expression of PhoB, CreB, and KpdE gene targets was cross-talk between CreC and PhoB. CreB is activated during the fermentation of glycolytic carbon sources that produce short chain organic acids such as 12ppd-R, lac-DL, and pyr (Cariss et al., 2008).

The PhoBR two component system and PST Pi ABC transporter were mutation hotspots during ALE

A series of mutations were found in phoB, phoR, and pstA that altered expression of genes of the PhoB regulon (Fig. 6B-F). pstA was mutated in all eTpiA strains (Fig. 6F). The mutation truncated the transmembrane protein PstA from 286 to 38 residues. Given that the PstA component is an integral part of the PST ABC transporter, it would seem likely that the truncation would inhibit the ability to transport phosphate, and would disrupt normal complex formation. The PST ABC transporter is the primary phosphate transporter in E. coli (Beard et al., 2000; Harris et al., 2001; Jackson et al., 2008). Loss of any of the PST components has been shown to change interaction of PstB with PhoU, leading to changes in phosphorylation patterns of PhoB by PhoR (Chekabab et al., 2014). Deletion of the PST system has been associated with virulence in Extraintestinal pathogenic E. coli (ExPEC) ( Crépin et al., 2012; Daigle et al., 1995; Glattard et al., 2010; Lamarche and Harel, 2010).

Mutations were found in either phoB or phoR that fixed in eTpiA02, 03, and 04 strains, while lower frequency mutations were found in phoB and phoR in eTpiA01 (Fig. 6B-D). These mutations targeted the PhoB TF binding site of the phoBR operon (Fig. 6B), altered the amino acid sequence of PhoB (Fig. 6C), shortened the length of the PhoR peptide (Fig. 6D), or hit the phoB response regulatory domain (Fig. 6E). The mutations in eTpiA02 and 03 resulted in the most deactivation of PhoB regulation, while the mutations in eTpiA01 and eTpiA04 retained
similar expression profiles to that of uTpiA (Fig. 6H). Interestingly, expression patterns of CreB targets in eTpiA01 and eTpiA04 also showed similar expression profiles to that of uTpiA indicating continued cross-talk with CreC (Table S4).

Bifurcated glycolytic flux perturbed energy and redox carrier balances, which lead to a global rerouting in the flux map to regain optimal growth

While lower glycolysis generates four energy equivalents (i.e., ATP) and two redox equivalents (i.e., NADH), the methylglyoxal pathway generates a toxic intermediate (i.e., methylglyoxal) and burns zero to two redox equivalents (i.e., NADH, NADPH, Reduced Glutathione) depending upon the pathway used. Consequently, levels of ATP were significantly depleted in uTpiA (-0.74 log2 fold change compared to Ref), which lead to a significant drop in the energy charge ratio (from 0.90 in Ref to 0.82) (Table S3). In addition, lower glycolytic intermediates including 1-3/2-3 diphosphoglycerate (13/23dpg), 2/3-phosphoglycerate, (2/3pg), pep, and pyr were severely depleted in uTpiA (-1.39, -2.62, -1.47, and -4.51 log2 fold change compared to Ref) (Table S3).

In particular, the depleted levels of pep appeared to be most detrimental to growth, and required major adjustments to all reactions and subsystems that involved pep. These included mutations to the Phosphotransferase System (PTS) and lower glycolytic enzymes. Major adjustments in flux also extended to subsystems that do not directly involve pep, but contributed flux to or from the pep node. These included upper glycolysis, the pentose phosphate pathway (PPP), and the TCA cycle. These adjust are detailed below.
Systematic decoupling of pep production from glucose import and cell wall recycling occurred during ALE

A host of mutations involving *ptsG*, *galR*, and *nagA*, *nagC*, and *nagE* in the eTpiA strains were found that appeared to systematically decouple pep consumption from carbon import. Mutations that altered the expression and function of nAcGam import and catabolic genes were found in eTpiA02/04 strains (Fig. 7). The *nagBAC* and *nagE* operons are negatively regulated by the NagC TF (Fig. 7A). EIIB encoded by *nagE* of the nAcGam PTS Permease imports and phosphorylates periplasmic N-acetyl glucosamine (Fig. 7B). The phosphate is derived from the conversion of phosphoenolpyruvate (pep) to pyruvate (pyr) after a series of phosphorylation reactions along the PTS components EI, HPr, and EIIA. N-acetyl glucosamine 6-phosphate (acgam6p) is then broken down to nitrate (nh4) and fructose-6-phosphate (f6p) by N-acetylglucosamine-6-phosphate deacetylase (AGDC) encoded by *nagA* and Glucosamine-6-phosphate deaminase (G6PDA) encoded by *nagB* (Fig. 7B). The genes that compose the *nagBAC* and *nagE* operons contribute to a significant recycling of peptidoglycan (Plumbridge, 2009). All *nagC* mutations in eTpiA02/04 appeared to be deleterious resulting in truncated peptides (Fig. 7C). The *nagC* mutations relieved NagC repression of the *nagBAC* and *nagE* operons, resulting in significantly higher expression in eTpiA02/04 (Fig. 7D). Increased expression appeared to be offset by mutations that prevent import and phosphorylation of acgam by *nagE*. The *nagE* mutations occur in the PTSEIIC type-1 complex formation region and resulted in amino acid changes that could have affected activity (Table S8).

Mutations that increased expression of non-PTS glucose importers occurred in eTpiA01/03 strains (Fig. 8). The primary importer of glucose in *E. coli* is the PTS, which is similar to the nAcGam PTS imports and phosphorylates glucose at the expense of pep (Fig. 8A). Specifically,
glucose is imported and phosphorylated from the periplasm into the cytosol by EIIB encoded by \textit{ptsG}. Alternative passive (i.e., D-galactose symporter encoded by \textit{galP}) and active importers (i.e., D-galactose ABC transporter encoded by \textit{mglBAC}) utilize the proton gradient or ATP, respectively, to import glucose (Fig. 8B). Passive and active importers are regulated by the GalR and GalS TFs (Fig. 8C). Mutations in \textit{galR} in eTpiA01/03 appeared to alter activity so as to increase expression of \textit{galETKM} and passive importer encoded by \textit{galP}, but not active ATP-dependent importer encoded by \textit{mglBAC} (Fig. 8F). In addition, a \textit{galP} INS mutation occurred at the NagC binding site in eTpiA01, further indicating an adaptive preference for passive import over active import (Fig. 8D). Note that eTpiA01 had the highest level of expression of \textit{galP}.

Complementary mutations in eTpiA01/03 occurred in \textit{ptsG} in that appeared to abolish import and phosphorylation activity of the primary PTS-dependent glucose importer. The \textit{ptsG} SNP mutation in eTpiA01 introduced a stop codon that truncates the cytosolic portion of the protein including the PTS EIIB type-1 active site and phosphocysteine residue (Fig. 8E). The \textit{ptsG} INS mutation (not shown in the figure) introduced an additional codon that shifted all residues by one in the PTS EIIB active site, which could have altered activity (Table S8).

**Additional systematic decoupling of pep enzymatic reactions**

In addition to carbon import processes, other enzymatic reactions that consume pep were also targeted by mutations. Mutations were found in \textit{pykF} and \textit{murA} that potentially affected enzymatic function (Fig. 9). Pyruvate kinase (PYK) is encoded by \textit{pykF} or \textit{pykA} and converts pep to pyr while generating ATP (Fig. 9A). Mutations found in \textit{pykF} in eTpiA02, 03, and 04 were most likely deleterious (Fig. 9B). The \textit{pykF} MOB resulted in a truncated peptide that removed all features except for two potassium ion binding sites and a substrate binding residue. The \textit{pykF} DEL was located at the interface between the dimers, and resulted in a frameshift that...
shifted the stop codon beyond the normal length of the peptide that most likely generated an aggregate peptide. The pykF SNP changed the 226 residue from G to D, and was located in close proximity to the magnesium binding site residues. UDP-N-acetylglucosamine 1-carboxyvinyltransferase (UAGCVT), encoded by murA, converts UDP-N-acetyl-α-D-glucosamine (uacgam) to UDP-N-acetyl-α-D-glucosamine-enolpyruvate (uaccg) at the expense of pep in the first committed step to peptidoglycan biosynthesis (Fig. 9C). The murA DEL in eTpiA04 resulted in a non-frameshift that shortened the peptide by one amino acid (Fig. 9D). Interestingly, the mutation changed the cysteine residue at position 115, which interacts with pep. The 115 residue has been actively investigated for implications in drug discovery (Bachelier et al., 2006; Barbosa et al., 2002; Baum et al., 2001; Couce et al., 2012; Desai and Miller, 2010; Dunsmore et al., 2008; Francisco et al., 2004; Kim et al., 1996; Miller et al., 2010; Xu et al., 2006).

Multiple pathway flux adjustments during ALE balanced methylglyoxal generation with biosynthesis

Bifurcation of lower glycolysis coupled glucose metabolism to the generation of the toxic intermediate methylglyoxal. A complete rewiring of metabolic flux through central metabolism was required for the eTpiA strains to recover approximately 50% of the growth rate of Ref. Notable shifts in flux were found in the pentose phosphate pathway (PPP), lower glycolysis, flux into the TCA cycle either through the anaerobic reaction Phosphoenol Pyruvate Carboxylase (PPC) or Citrate Synthase (CS), and utilization of the glyoxylate shunt. A 5.14 log2 fold change in flux per mol glucose through the Entner–Doudoroff (ED) pathway compared to Ref in uTpiA shifted to 3.60, 2.46, 3.36, and 4.76 log2 fold change in eTpiA strains, respectively. A -1.69 log2 fold change in flux per mol glucose through PYK compared to Ref in uTpiA shifted to -5.94, -5.49, -6.36, and -4.49 log2 fold change in eTpiA strains, respectively. The relative drop in PYK
flux in many of the endpoints could in part be attributed decreased expression, but also due to mutations found in the pykF gene (discussed previously). The majority of pyr levels were recovered through the rejoining of flux through the methylglyoxal pathway instead of through PTS reactions and PYK. A 1.7 mmol*gDCW^{-1}*hr^{-1} flux in from lac-DL to pyr in uTpiA increased to 4.9, 6.2, 4.7, and 2.5 in eTpiA strains, respectively.

Overall flux into the TCA cycle per mol of glucose dropped in all eTpiA strains. An initial 90% relative boost per mole of glucose in PPC flux compared to Ref in uTpiA was lost to a -54%, -15%, -49%, and -31% relative drop in eTpiA strains, respectively. An initial 182% relative boost per mole of glucose in CS flux compared to Ref in uTpiA was reduced to 17%, 96%, 34%, and 90% in eTpiA strains, respectively. In addition, partial utilization of the glyoxylate shunt was restored in eTpiA strains. A -94% relative change per mole of glucose through the glyoxylate shunt compared to Ref in uTpiA was increased to 57%, 58%, 103%, and 52% in eTpiA strains, respectively.

The above shifts in flux appeared to simultaneously increase flux through lower glycolysis (i.e., increased flux through ED) to increase the levels of lower glycolytic intermediates and generation of ATP and redox equivalents while minimizing glycolytic flux that would commit 1 mol of glucose to methylglyoxal generation. In addition, the above shifts in flux also maximized pyr production while minimizing pep consumption resulting in an overall loss in flux through the TCA cycle. Interestingly, while many of the redox ratios were recovered, the energy charge ratio was never recovered (Table S3), indicating that the coupling of biomass production to methylglyoxal production induced an intrinsic cost to energy production.
Conclusion

Flux through glycolysis was stoichiometrically coupled to the production of the toxic metabolite methylglyoxal in *E. coli* by removing the *tpiA* gene. Production of methylglyoxal resulted in cellular damage that was overcome through efficient detoxification by the Glyoxalase enzymes. Elevation in the end-products of the methylglyoxal pathway, 12ppd-R and lac-DL, triggered an erroneous phosphate starvation response via cross talk with CreCB TCS that was dampened through mutations that affect PhoB signalling. Split flux through lower glycolysis depleted glycolytic intermediates, in particular, pep. Pep levels were restored to levels similar to Ref through mutations that decoupled pep production from carbon import. The detoxification strategy of methylglyoxal and network changes required an increase in carbon import without exacerbating methylglyoxal toxicity or depletion of glycolytic intermediates that serve as key biomass precursors. Thus, multi-omics analysis and ALE confirmed in higher resolution than previously reported methylglyoxal toxicity responses and revealed novel biochemical mechanisms that should provide researchers with valuable insights into strategies deployed for metabolic detoxification in bacteria.

Taken together, this study shows that the combination of KO, multi-omic data analysis and ALE represent a new line of biological inquiry. It reveals the full consequences of the loss of a gene product and the genetic changes needed to regain fitness. This overall conclusion provides an impetus to perform analogous studies for a series of major gene products in *E. coli*. Such studies would provide us with a new and deep understanding of the systems biology of basic regulatory and metabolic processes in bacteria.
Contributions

D.M. designed the experiments; generated the strains; conducted all aspects of the metabolomics, fluxomics, phenomics, transcriptomics, and genomics experiments; performed all multi-omics statistical, graph, and modeling analyses; and wrote the manuscript. T.E.S. ran the ALE experiments. E.B. assisted with structural analysis. R.S. processed the DNA and RNA samples. S.X. assisted with metabolomics and fluxomics data collection, sample processing, and peak integration. Y.H. assisted with fluxomics data collection and sample processing. A.M.F designed and supervised the evolution experiments, and contributed to the data analysis and the manuscript. B.O.P conceived and outlined the study, supervised the data analysis, and co-wrote the manuscript.

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Competing financial interests

The authors declare no competing financial interests.
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Fig. 1 Evolution of knockout (KO) strains from a pre-evolved (i.e., optimized) wild-type strain. A) Wild-type (wt) E. coli (MG1655 K-12) was previously evolved on glucose minimal media at 37°C (LaCroix et al., 2015). An isolate from the endpoint of the evolutionary experiment was selected as the starting strain for subsequent KO of tpiA and adaptive laboratory evolution (ALE). B) Adaptive laboratory evolution trajectories of the evolved knockout lineages. -Oms data collected included metabolomics, fluxomics, physiology, DNA resequencing, and transcriptomics. C) Triosephosphate isomerase (TPI) was disabled by the gene KO. TPI interconverts the triosephosphate dihydroxyacetone phosphate (dhap) to glyceraldehyde 3 phosphate (g3p). Additional metabolites shown for context include D-glucose (glc-D), glucose 6-phosphate (g6p), ribulose 5-phosphate (ru5p-D), methylglyoxal (mthgxl), phosphoenolpyruvate (pep), pyruvate (pyr), DL-lactate (lac-DL), acetate (ac), alpha-ketoglutarate (akg), and succinate (succ). D) Growth rate and uptake and secretion rates of D-glucose, acetate, and DL-lactate for unevolved KO (uTpiA) and evolved KOs (eTpiA).

Fig. 2 The removal of triose phosphate isomerase (TPI) forced a bifurcation of flux in glycolysis through a pathway that involved the toxic intermediate methylglyoxal (mthgxl) (panels A–E). The bifurcated flux is rejoined by the lactate dehydrogenase (LDH) reaction. A) A network
schematic of middle glycolysis. Shown along with removed TPI (shown in red) are the reactions catalyzed by fructose bisphosphate aldolase (FBA), glyceraldehyde 3-phosphate dehydrogenase (GAPD), and methylglyoxal synthase (MGSA). B) Flux levels of reactions derived from C\textsuperscript{13} labeling experiments near the gene knockout. C) Network schematic of lactate and pyruvate conversion. D) Regulatory schematic showing the feedback loop that up-regulates the conversion of D/L-Lactate to pyruvate when elevated levels of intracellular lactate were sensed by the transcription factor lldR. E) Metabolite, expression, and measured flux levels involved in the regulatory feedback loop.

Fig. 3 Optimization of methylglyoxal detoxification pathway expression in tpiA lineages. Paths 1, 2, and 3 were actively expressed in uTpiA, while path 4 was not. Through fixed mutations that affected gloA expression, path 4 was expressed in the eTpiA endpoints. A) Network diagram of the four methylglyoxal detox pathways in \textit{E. coli}. B) Expression profiles of methylglyoxal detox pathway genes. Path 1: \textit{fucO} and \textit{gldA} are up regulated in uTpiA compared to all eTpiA strains; \textit{yqhD} and \textit{dkgA} are up regulated in uTpiA compared to Ref and are up regulated in most eTpiA strains compared to uTpiA. Path 2: \textit{AldA} is up regulated in uTpiA compared to Ref and in most eTpiA strains compared to Ref, but down regulated in all eTpiA strains compared to uTpiA. Path 3: \textit{HchA} is significantly elevated in uTpiA compared to all eTpiA strains and Ref; \textit{yhbO} is up regulated in eTpiA 3 compared to Ref. Path 4: GloA is up regulated in all eTpiA strains compared to Ref and uTpiA. Up/Down regulation p-value < 0.05.

Fig. 4 Select mechanisms of methylglyoxal (mg or mthgxl) attack on cellular components. A) ball and stick diagram of methylglyoxal. Methylglyoxal contains two reactive carbonyls: an aldehyde and a ketone. Methylglyoxal exerts its toxicity through damage of cellular components with which it interacts. B) methylglyoxal cross-links single stranded RNA or DNA (shown) and/or peptides (shown in D) with other peptides between a RNA/DNA base and/or sulfur residues and lysine residues through formation of a imidazolysine group via the Maillard reaction (Broude and Budowsky, 1971; Krymkiewicz, 1973; Lo et al., 1994; Murata-Kamiya and Kamiya, 2001). C) methylglyoxal forms adducts with sulfur residues of enzymes directly that can inhibit their function (Kalapos, 1994; Ridderström and Mannervik, 2002). D) methylglyoxal participates in the formation of free radicals and reactive oxidative species (ROS) through glycation of amino acids (shown) or nucleotide bases (not shown) via Fenton’s Reaction (Cadet et al., 2003; Cooke et al., 2003; Imlay and Linn, 1988; Kang, 2003). E) The generated ROS is then capable of inducing single stranded DNA or RNA (ssRNA/ssDNA) breaks. F) ROS attacks the ssRNA or ssDNA backbone generating 2-phosphoglycolate (2pgblyc) a deoxynucleotide and a nucleotide base. G) Metabolite concentrations of nucleobases, nucleotides, and mono-, di-, and tri- nucleotide phosphates. H) RNA expression profiles of select genes that respond to cellular stress and ROS.

Fig. 5 Mutations that affect the transcription factor NemR that allowed for expression of the gloA detox pathway (Ozyamak et al., 2013). A) Crystal structure of the \textit{nemR} multimer (Gray et al., 2015). A mutation that alters NemR binding to the regulatory region are annotated in red. Chains A, B, C, and D are highlighted in gold and grey; the regulatory region is highlighted in cyan; mutations are annotated in red. B) Reorientation and zoom in on two of the \textit{nemR} multimer chains. C) Schematic of the \textit{nemRA-gloA} operon (see Fig. 6 for expression profiles of the methylglyoxal detox pathways). NemR exerts negative feedback on the operon that is enhanced by increased levels of methylglyoxal (mthgxl) (Ozyamak et al., 2013; Umezawa et al.,
2008). *gloA* is co-expressed with *nemR* and *nemA* genes. D) Mutation frequency and expression profiles of *nemR* and *nemR*-associated genes. The lineages with mutations have significantly increased expression of the *gloA* detox pathway (p-value < 0.05).

Fig. 6 A host of mutations targeted elements of the phosphate starvation response regulon, including the PhoRB two component system (TCS) and the Pst phosphate ABC transporter. A) The PhoRB two component system (TCS) senses and responds to the absence of extracellular inorganic phosphate (Pi), and is composed of a transmembrane sensory histidine kinase (PhoR) and transcription factor (PhoB). The PhoRB signalling processes are mediated by an ABC (ATP-dependent) transporter encoded by *pstS*, *pstC*, *pstA*, and *pstB*, that imports Pi as well as a chaperone-like inhibitory protein (PhoU) that physically interacts with the cytosolic portion of the Pst Pi ABC transporter (PstB) and PhoR. In the presence of extracellular Pi (Pi concentration approximately > 4 uM), the extracellular portion of the Pst Pi ABC transporter (PstS) is saturated. This leads to a signal transduction mediated by PhoU that leads to a conformational change of PhoR into an inactive state. Under conditions of phosphate limitation or in the absence of any of the Pst components, PhoR returns to its default active conformation, where it is able to phosphorylate and activate PhoB. B) Operon diagram for *phoB*. An 11 nucleotide deletion occurred at the PhoB TF binding site (*sbcD*/*phoB* (DEL) 416288 Δ11nt) in eTpiA03. C) A 1 nucleotide deletion at the 205 residue in *phoB* (*phoB* (DEL) 205Δ1nt) in eTpiA03 in 61.3% of the population resulted in a frameshift that changed all amino acids from the site of the mutation to the end of the peptide. Interestingly, the peptide length was not changed. D) *phoR* encodes the transmembrane sensory histidine kinase component of the PhoRB TCS. Two, most likely deleterious mutations, targeted *phoR*. The *phoR* deletion at the 180 residue (*phoR* (DEL) 180Δ1nt) in eTpiA01 affected 53.2% of the population, and resulted in a frameshift that truncated the peptide at the 187 residue, cleaving the phosphohistidine and histidine kinase residue and domains, respectively. The *phoR* deletion at the 358 residue (*phoR* (DEL) 358Δ1nt) in eTpiA04 resulted in a frameshift that truncated the peptide at the 368 residue, cleaving the histidine kinase domain, but leaving the phosphohistidine regulatory regions intact. E) Crystal structure of PhoB (Solá et al., 1999). PhoB is the response regulator (i.e., TF) component of the PhoRB TCS, and induces or represses operons of the Pho regulon. An SNP in *phoB* (*phoB* (SNP) A84P) occurred in eTpiA01 that affected 52.9% of the population. The mutation is located near the L105 residue that binds magnesium (mg2+), which may affect magnesium binding. The *phoB* deletion at the 39 residue (*phoB* (DEL) 39Δ17nt) in eTpiA03 resulted in a frameshift that truncated the peptide at the 89 residue. The *phoB* deletion at the 76 residue (*phoB* (DEL) 76Δ5nt) in eTpiA02 resulted in a frameshift that truncated the peptide at the 93 residue. Mutations are highlighted in red. The bound magnesium is shown in green. F)
PstA encodes a transmembrane component of the phosphate ABC transporter, which imports extracellular phosphate at the expense of ATP. A deleterious 774 nucleotide deletion in PstA (PstA (DE) 1Δ774nt) truncated the peptide to all but 38 amino acids. Loss of any of the Pst components has been shown to lead to a perpetual activation of the PhoRB TCS.

G) Regulatory diagram of operons controlled by PhoB. H) Mutation frequency and expression of genes involved in phosphate starvation response regulon. Note that the majority of genes in the Phn and Pho operon are significantly elevated in uTpiA. See Table S4 for a comprehensive list of all significant changes in gene expression.

Fig. 7 Mutations that altered expression and function of nAcGam import and catabolic genes. A) Schematic of the nagBAC (catabolic enzymes) and nagE (PTS-specific component) operons. B) A network and mechanistic schematic of the nAcGam PTS Permease. N-acetyl glucosamine is imported from the periplasm and phosphorylated by EIIB (encoded by nagE). The phosphate is derived from the conversion of phosphoenolpyruvate (pep) to pyruvate (pyr) after a series of phosphorylation reactions along the PTS components EI, HPr, and EIIA. N-acetyl glucosamine phosphate (acgam6p) is then broken down to nitrate (nh4) and fructose 6-phosphate (f6p) by N-acetylglucosamine-6-phosphate deacetylase (AGDC) encoded by nagA and Glucosamine-6-phosphate deaminase (G6PDA) encoded by nagB. C) Crystal structure of NagC. The majority of the nagC mutations appeared to be deleterious. The nagC mobile element insertion (MOB) mutation truncated the peptide from 409 AAs to 339 AAs. The nagC insertion (INS) mutation truncated the peptide from 409 to 96 AAs. The nagC SNP mutation results in a change of amino acids from D to V. Mutations are highlighted in red, orange, and yellow. The DNA binding region is highlighted in cyan. The nagE mutations (not shown) occurred in the PTSEIIIC type-1 complex formation region and resulted in amino acid changes (Table S8). The nagA SNP mutations (not shown) all resulted in amino acid changes (Table S8). The nagA MOB was most likely deleterious, and truncated the peptide from 382 to 168 AAs (Table S8). D) Mutation frequency and expression profiles of nagBAC and nagE genes. Note that nagE and nagC were both mutated in eTpiA02/04, while lower level mutations in nagA occurred in eTpiA01/03. While nagBAC and nagE are overexpressed in all eTpiA strains, eTpiA02/04 have significantly higher expression than eTpiA01/03.

Fig. 8 Mutations that increased expression of non-PTS glucose importers. Network and mechanistic schematic of the A) Glucose-specific PTS permease and B) passive and active glucose importers. Glucose is imported and phosphorylated from the periplasm into the cytosol by EIIB encoded by ptsG. C) Network schematic of operons regulated by GalR and GalS. D) Regulatory schematic of the GalP operon. The galP INS mutation occurred at the NagC binding site. Note that eTpiA01 had the highest level of expression of galP. E) Crystal structure of the cytosolic portion of EIIB encoded by ptsG. The ptsG SNP introduced a stop codon that truncated the cytosolic portion of the protein including the PTS EIIB type-1 active site and phosphocysteine residue. The mutation is highlighted in red. The crr interaction domain is highlighted with a bounding box. The ptsG INS mutation (not shown) introduced an additional codon that shifted all residues by one in the PTS EIIB active site, which could have altered activity (Table S8). F) Crystal structure of GalR. Mutations are highlighted in red. The small molecule binding domain is highlighted in cyan. The DNA binding region is highlighted in...
magenta. G) Mutation frequency and expression levels of genes that compose the glucose-specific PTS as well as the D-galactose symporter and ABC transporter. Note that The D-galactose symporter and ABC transporter shown high affinity for glucose. \(P_{ts}G\) is down regulated in \(uTpiA\) and all \(eTpiA\) strains compared to Ref. \(GalP\) is up regulated in \(eTpiA01\) and 03 strains compared to \(uTpiA\) and Ref while \(uTpiA\) is down regulated in \(uTpiA\) compared to Ref. \(GalU\) is down regulated in \(uTpiA\) compared to Ref. \(GalS\) is up regulated in \(uTpiA\) and \(eTpiA\) strains compared to Ref. \(GalETKM\) operon genes are all up regulated in \(eTpiA01\) and 03 strains. Up/Down regulation p-value < 0.05.

Fig. 9 Mutations that affect \(pykF\) and \(murA\) activity. A) Network schematic of the PYK reactions. B) Crystal structure of PykF. Mutations are annotated in red. Substrate binding site residues are shown as ball and stick. The magnesium and potassium ion binding sites are highlighted in blue. The N6-acetyllysine residues are highlighted in cyan. The \(pykF\) MOB resulted in a truncated peptide that removed all features except for two potassium ion binding sites and a substrate binding residue. The \(pykF\) DEL was located at the interface between the dimers, and resulted in a frameshift that shifted the stop codon beyond the normal length of the peptide that most likely generated an aggregate peptide. The \(pykF\) SNP changed the 226 residue from G to D, and was located in close proximity to the magnesium binding site residues. C) Network schematic of the UAGCVT reaction encoded by \(murA\). D) Crystal structure of MurA (Eschenburg and Schonbrunn, 2004). The \(murA\) DEL resulted in a non-frameshift that shortened the peptide by one amino acid. Interestingly, the mutation changed the cysteine residue at position 115, which interacts with pep. The 115 residue in particular and \(murA\) as a whole has been actively investigated for implications in drug discovery (Bachelier et al., 2006; Barbosa et al., 2002; Baum et al., 2001; Couce et al., 2012; Desai and Miller, 2010; Dunsmore et al., 2008; Francisco et al., 2004; Kim et al., 1996; Miller et al., 2010; Xu et al., 2006). The mutated residue is highlighted in yellow. E) Mutation frequency, metabolite concentrations, expression, and metabolic flux derived from \(C^{13}\) labeling experiments of PYK and associated network components.

Highlights:

- \(tpiA\) Knockout-out in \(E. coli\) and adaptive laboratory evolution (ALE)
- Consequences of methylglyoxal toxicity and mechanisms for detoxification revealed
- Cross-talk between carbon starvation and inorganic phosphate signalling was uncovered
- Counter intuitive inactivation of PTS required to rebalance the flux map