Metabolic and Transcriptional Response to Cofactor Perturbations in Escherichia coli

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Metabolic cofactors such as NADH and ATP play important roles in a large number of cellular reactions, and it is of great interest to dissect the role of these cofactors in different aspects of metabolism. Toward this goal, we overexpressed NADH oxidase and the soluble F1-ATPase in Escherichia coli to lower the level of NADH and ATP, respectively. We used a global interaction network, comprising of protein interactions, transcriptional regulation, and metabolic networks, to integrate data from transcription profiles, metabolic fluxes, and the metabolite levels. We identified high-scoring networks for the two strains. The results revealed a smaller, but denser network for perturbations of ATP level, compared with that of NADH level. The action of many global transcription factors such as ArcA, Fnr, CRP, and IHF commonly involved both NADH and ATP, whereas others responded to either ATP or NADH. Overexpressing NADH oxidase invokes response in widespread aspects of metabolism involving the redox cofactors (NADH and NADPH), whereas ATPase has a more focused response to restore ATP level by enhancing proton translocation mechanisms and repressing biosynthesis. Interestingly, NADPH played a key role in restoring redox homeostasis through the concerted activity of isocitrate dehydrogenase and UdhA transhydrogenase. We present a reconciled network of regulation that illustrates the overlapping and distinct aspects of metabolism controlled by NADH and ATP. Our study contributes to the general understanding of redox and energy metabolism and should help in developing metabolic engineering strategies in E. coli.

Microbial metabolic networks function in a coherent fashion to convert the available substrates into biomass and products. Often, intermediates of metabolism are not balanced with respect to their redox and/or energy content. Metabolic cofactors such as NADH and ATP serve in overcoming these constraints. Indeed, these cofactors rank among the most highly connected metabolites in the metabolic networks of most microorganisms (1). A direct consequence of this metabolic structure is that a small change in the concentration of these cofactors is likely to propagate to widespread aspects of metabolism. For example, the synthesis of proteins, lipids, and nucleotides is energetically expensive and would drain cellular ATP and also require NADPH (2). ATP is primarily produced in the electron transport chain, which is fueled by NADH. NADH is produced in the catabolism, relating substrate utilization to biosynthesis and product formation. Therefore, these cofactors hold the potential to serve as targets for altering cellular metabolism.

To achieve the desired metabolic changes, it is important to understand the metabolic processes that are specifically controlled by cofactors. NADPH primarily drives anabolic reactions, whereas NADH is the result of catabolism. To fulfill their distinct roles, these two redox couples are generally not in thermodynamic equilibrium. NADPH is primarily produced in the oxidative branch of the pentose phosphate pathway or by the isocitrate dehydrogenase in the tricarboxylic acid (TCA) cycle. Many organisms also have a transhydrogenase that can interconvert NADH and NADPH, and some bacteria such as Escherichia coli rely more heavily than others on transhydrogenases for NADPH (3).

Because the cofactors are intimately connected with one another via the transhydrogenases or the oxidative phosphorylation, there is no clear distinction on the role of NADH and ATP in regulating metabolism. Increasing the intracellular level of NADH in E. coli increased acetate excretion accompanied with a shift in metabolism to produce typically fermentative products such as lactate and succinate even in the presence of ample oxygen (4, 5). The converse approach of lowering NADH level decreased acetate overflow (6). On the contrary, acetate overflow increased when ATP level was lowered in E. coli (7). These two approaches are of particular interest because they involve introducing heterologous enzymes that specifically decrease NADH or ATP concentrations. This is likely to invoke an unbiased response as opposed to affecting a specific metabolic reaction, which will have localized network effects around the altered reaction. Water-forming NADH oxidase from Streptococcus pneumoniae was used to lower cellular NADH concentrations (6), while the soluble F1 part of the F0F1-ATP synthase was used to specifically hydrolyze ATP (7).

1 The on-line version of this article (available at http://www.jbc.org) contains supplemental text sections S1–S7.
2 The abbreviations used are: TCA, tricarboxylic acid; MOPS, 4-morpholinepropanesulfonic acid; TF, transcription factor; SF, sigma factor; HSN, high scoring subnetwork; PEP, phosphoenolpyruvate; OAA, oxaloacetate.

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The predominant phenotype of *E. coli* harboring these enzymes shows an increased glycolytic flux at the expense of biomass production. Overexpression of NADH oxidase decreased acetaldehyde overflow whereas overexpression of the soluble ATPase increased acetaldehyde excretion. Transcription analysis in NADH oxidase overexpressed *E. coli* revealed elegant regulation governed by ArcA that brought about these changes (6). Many global transcription factors and two-component signal transduction systems in *E. coli* are triggered by redox or energy cofactors (8) and yet, only few of these global transcription factors affect metabolic fluxes (9). From a basic research point of view, there is a gap in our understanding of the influence of these transcriptional regulators on metabolism.

The objective of this research is to determine the role of NADH and ATP in regulating metabolism. More specifically, we aim to map the overlap in the regulation and identify aspects of metabolism that are specifically controlled by NADH or ATP. Toward this goal, we quantified changes in metabolic fluxes in central carbon metabolism, the level of central carbon ATP. Toward this goal, we quantified changes in metabolic fluxes in central carbon metabolism, the level of central carbon metabolism, the level of central carbon metabolites and cofactors and genome-wide transcription changes in response to overexpressing NADH oxidase or the soluble F₁,ATPase subunit. We analyzed the response in the context of a global interaction map and identified metabolic modules that responded to the perturbations.

**EXPERIMENTAL PROCEDURES**

**Construction of Strains and Plasmids**—The *E. coli* K12 strain, MG1655, was used as the reference strain in this study. All the strains and plasmids used in this study are listed in Table 1. In this strain, NADH oxidase or the soluble ATPase were overexpressed using the *nox* gene from *Streptococcus pneumoniae* (GenBank¹¹ entry AF014458) or the native *atpAGD* operon (encoding the α-, β-, and γ-subunits of the ATPase). The *nox* gene was amplified from the pTrc99A-*nox* plasmid (6) using the forward primer containing the restriction site for XhoI (underlined), AGTTGGCTCGAGATGCNNNNNAGTTTATTCNNNTGACANNNNNNNNNNNNTGRTATAATNNNNCN- and the reverse primer containing the restriction site for Xmal (underlined), GTCAAGAGCTGAGATGAAAAATCTTGGTGGAGTCAGTCCCGGG.

The annealing portion of the primer to vary the promoter strength (10–12). These fragments were digested with XhoI and XmaI restriction enzymes and ligated into the pAK80 vector (13), which was also subjected to restriction digest with the same enzymes. The pCP41 plasmid containing the *atpAGD* operon (7), which resulted in a moderate level of ATPase activity was used. The PCR was performed using Phusion polymerase (Finnzymes, Espoo, Finland). The plasmids were introduced into *E. coli* MC1000 for screening. The plasmid that resulted in approximately the same reduction in growth rate as the pCP41 was selected for this study (pAC06). The presence of the genes was verified by an enzymatic assay as previously described (6, 7). The MG1655 strain was transformed with pAK80, pCP41 and pAC06 plasmids and the resulting strains are designated REF (containing empty vector), ATPase, and NOX, respectively.

**Media and Growth Conditions**—Strains were maintained on LB agar plates with 200 µg/ml of erythromycin when required. Growth experiments were carried out in 500-ml baffled shake-flasks with a working volume of 100 ml at 37 °C in defined MOPS medium (14) supplemented with 0.1% glucose and 200 µg/ml erythromycin. The flasks were agitated at 150 rpm to ensure aerobic conditions. Optical density was measured at 600 nm on a Shimadzu UV mini 1240 spectrophotometer. Promoters of different strengths were selected based on the extent of blue coloration on X-gal plates and quantified by a subsequent assay for β-galactosidase activity, as described previously (12). From this promoter library, a clone showing a similar decrease in the growth rate as the pCP41:*atpAGD* was selected for all further studies. The cultivations for quantification of intracellular fluxes were performed under identical conditions as described above using a mixture of 20% (wt/wt) [U-¹³C]glucose (˃99% pure, Martek Biosciences, Columbia, MD) and 80% naturally labeled glucose. Samples from different stages of exponential growth were filtered through 0.2-µm filters and analyzed by HPLC (Shimadzu, Kyoto, Japan) containing a Bio-Rad HPX-87H column (Phenomenex, Torrence, CA). The mobile phase consisted of 5 mM H₂SO₄ at a flow rate of 0.5 ml/min, and glucose, lactate, formate, and acetate were detected on the refractive index detector, RID-10A.

**Sample Extraction and Processing**—Biomass samples for RNA extraction were taken from mid-exponential growth phase. An aliquot of 25 ml was chilled on ice, centrifuged at 5000 × g at 4 °C and stored at −80 °C until subsequent analysis. Total RNA was purified using the RNeasy Mini kit (Qiagen, Carlsbad, CA) and quantified using a NanoDrop (NanoDrop products, Wilmington, DE). RNA quality was determined using an Agilent 2100 Bioanalyzer (Agilent, Santa Clara, CA). Gene expression was analyzed using the Affymetrix platform with Affymetrix *E. coli* genome 2.0 chips according to the manufacturer’s specification.

Biomass samples for determining the fluxes were centrifuged (5000 × g, 10 min at 4 °C) and harvested in 150 µl of 6 M HCl and hydrolyzed at 105 °C for 24 h before drying at 85 °C. The hydrolysate was dissolved in 50 µl of dimethyl formamide and derivatized using 30 µl of N-methyl-(tert-butyl(dimethyl)silyl)-trifluoroacetamide. The mixture was incubated at 85 °C with shaking at 550 rpm for 60 min (Orbital Shaker CSA, Thermo Scientific) before GC-MS analysis, using previously described protocols (15, 16). From the mass isotope distribu-

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**TABLE 1**

<table>
<thead>
<tr>
<th>Plasmid/strain</th>
<th>Description</th>
<th>Source/Ref.</th>
</tr>
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<tbody>
<tr>
<td>pAK80</td>
<td>Promoterless vector carrying the lacLM genes, Erm&lt;sup&gt;R&lt;/sup&gt;</td>
<td>(13)</td>
</tr>
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<td>pAC06</td>
<td>pAK80::nox (synthetic promoter), Erm&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This study</td>
</tr>
<tr>
<td>pCP41::atpAGD</td>
<td>ATPase activity, pCP41::atpAGD, Erm&lt;sup&gt;R&lt;/sup&gt;</td>
<td>(7)</td>
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</table>

**Strains**

<table>
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<tr>
<th>Strain</th>
<th>Description</th>
<th>Source</th>
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<tbody>
<tr>
<td>MC1000</td>
<td>Wild-type reference for expression analysis</td>
<td>(35)</td>
</tr>
<tr>
<td>MG1655</td>
<td>Wild-type for growth experiments</td>
<td>Lab stock</td>
</tr>
<tr>
<td>BK1052</td>
<td>ROE270 with pCP41 plasmid</td>
<td>(7)</td>
</tr>
<tr>
<td>REF</td>
<td>MG1655 transformed with pAK80</td>
<td>This study</td>
</tr>
<tr>
<td>NOX</td>
<td>MG1655 transformed with pAC06</td>
<td>This study</td>
</tr>
<tr>
<td>ATPase</td>
<td>MG1655 transformed with pCP41::atpAGD</td>
<td>This study</td>
</tr>
</tbody>
</table>
tion patterns of the proteinogenic amino acids, relative pathway contributions (metabolic flux ratios) were computed that were used as constraints to infer metabolic activity (15, 16). An aliquot of 5 ml of the culture from mid-exponential phase was quenched in hot phenol (80 °C) and subsequently frozen at −20 °C for measuring the concentration of intracellular metabolites. The samples were processed as described previously (17) and analyzed using LC-MS/MS for quantitative determination of central carbon metabolites and cofactors. The values were normalized with biomass concentrations at the time of sampling.

**Microarray Data Analysis**—All the transcriptome data used in this study can be obtained in the form of CEL files from the NCBI Gene Expression Omnibus using the accession number GSE 20374. The raw data obtained in the form of the CEL files were subjected to preprocessing using median polish summarization. Background correction was performed using GCRMA (18), and the data were normalized by non-linear, quantile normalization using qspline (19). The data obtained after preliminary processing were assessed for good reproducibility. Genes with log fold-changes for the expression of NOX and ATPase, which differed by a factor of two, relative to REF, were considered significantly changed (increased or decreased). The p value of their significance was calculated using Student’s t test. The biological role of the significantly altered gene expressions was determined by using the biological process semantics maintained by the Gene Ontology Consortium. A hypergeometric distribution was used to determine the significant biological processes in different sets of genes. We also used datasets available in public repositories to validate our observations. The context of the datasets and their accession numbers is presented in the S6 section of the supplemental text.

**Integrated Data Analysis**—The transcriptional regulatory network of *E. coli* was constructed using information from the RegulonDB database (20), which documents the known regulatory interactions between transcription factors (TFs) and sigma factors (SFs) with genes. The amount of regulation exerted by each TF or SF was quantified as a p value that was calculated as described (21). These algorithms take into account the significance of the expression of genes regulated by the different TFs and SFs. The regulators are scored after normalizing with the background (rest of the genes), and the probability that their influence is different from the background is calculated. Those TFs and SFs whose p value is less than 0.05 in either NOX or ATPase are considered in this study. To identify the most affected nodes in the metabolic network, we used the genome-scale metabolic network of *E. coli* (22) and overlaid the p values of the genes which encode for the enzymes that catalyze the different reactions in the model. Using a similar algorithm (23), metabolites around which significant transcriptional activity in their network neighborhood were identified. A protein interaction network of *E. coli* was constructed by combining data from high throughput TAP-tag pull-down experiments (24, 25) by computational curation (26, 27). A global interaction map in *E. coli* was constructed by overlaying the transcriptional regulatory network and the protein interaction network on the genome-scale metabolic network. This network resulted in 4936 cellular components with 18943 interactions, and is available in mark-up language format as a supplemental file. High-scoring networks (HSNs) within the global interaction map were identified using simulated annealing by invoking the *activemodules* algorithm (28) using the default parameters in a Cytoscape environment (29), which was also used for visualization.

**RESULTS**

We decreased the level of NADH or ATP in *E. coli* to dissect the regulatory effects of these cofactors on metabolism. We used water-forming NADH oxidase and the soluble F$_1$-ATPase to effect the perturbation. The net reactions mediated by these enzymes are given in Reaction 1.

Soluble F$_1$-ATPase: ATP → ADP + P,

NADH oxidase: NADH + (1/2)O$_2$ → NAD$^+$ + H$_2$O

**Reaction 1**

This strategy is expected to invoke an unbiased response, in contrast to deleting or overexpressing metabolic pathways directly involved in cofactor utilization, which may result in localized changes in fluxes surrounding the reaction. The strains of *E. coli* in which the heterologous genes are expressed are referred to as ATPase or NOX for convenience.

**Physiological Response to Cofactor Perturbations**—The physiological changes we observed in response to overexpressing NADH oxidase or the soluble ATPase appeared to be geared to restoring the redox or energy balance, respectively. The dominant phenotype that was common to NOX and ATPase strains was increased glucose uptake by 27 and 70%, respectively (Table 2). The specific growth rate decreased in NOX and ATPase strains by 10 and 20%, respectively. The biomass yield also decreased substantially in the NOX strain (33%) and ATPase (49%) strains relative to the reference strain (see section S1 in supplemental text for growth profiles). This interesting behavior of consuming glucose faster, but channeling it away from biomass was shared by both NOX and ATPase strains. Acetate production rate increased 3-fold in ATPase, relative to the reference strain, whereas it decreased by 40% in NOX compared with the reference strain (Table 2).

**Metabolic Flux Redistribution**—To elucidate the impact of decreasing NADH or ATP level, we obtained flux distribution in the central carbon metabolism in the reference, NOX and ATPase strains. We normalized the fluxes to the specific rate of glucose uptake and present relative flux values (Fig. 1). The values of absolute and relative fluxes are provided in section S2 in the supplemental text.

**TABLE 2**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Growth (μm/h)</th>
<th>Specific glucose uptake rate (mmol/g dry cell weight (DCW) h)</th>
<th>Specific acetate production rate (mmol/d DCW h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>REF</td>
<td>0.72 ± 0.01</td>
<td>0.43 ± 0.02</td>
<td>9.2 ± 0.11</td>
</tr>
<tr>
<td>NOX</td>
<td>0.65 ± 0.02</td>
<td>0.29 ± 0.03</td>
<td>11.7 ± 0.21</td>
</tr>
<tr>
<td>ATPase</td>
<td>0.58 ± 0.01</td>
<td>0.22 ± 0.00</td>
<td>15.6 ± 0.19</td>
</tr>
</tbody>
</table>

$^a$ Maximum specific growth rate (μm/h).

$^b$ Biomass yield from glucose (g dry cell weight/g glucose).

$^c$ Specific glucose uptake rate (mmol/g dry cell weight (DCW) h).

$^d$ Specific acetate production rate (mmol/d DCW h).
Increased glucose uptake in the mutant strains resulted in a higher flux in the glycolytic pathway. This phenotype is more prominent in ATPase strain (30%) than in NOX (12%). Concomitantly the relative flux in the pentose phosphate pathway decreased. Faster operation of glycolysis triggered related changes in the entire central carbon metabolism, which manifested in flux redistribution around key branch points, such as PEP (phosphoenolpyruvate) and pyruvate. The anaplerotic flux via PEP carboxylase decreased by 21% only in the NOX strain, whereas it remained unchanged in the ATPase strain. Interestingly, we detected a small amount of anaplerotic flux via malic enzyme (from malate to pyruvate) only in the NOX strain, while it was absent in the reference and ATPase strains. In comparison to REF, the relative TCA cycle flux increased by 78% in NOX while it increased only 21% in ATPase. Higher glycolytic flux (relative to REF) was transmitted to the TCA cycle only in NOX, whereas it was excreted as acetate in ATPase. The perturbations to reduce the level of NADH and ATP also resulted in a remarkable increase in the transhydrogenase flux, to produce NADH from NADPH. In comparison with REF, the relative transhydrogenase flux increased over 6-fold in NOX and over 5-fold in ATPase (Fig. 1).

The nature of the analysis technique also allows quantifying the origin of key metabolic intermediates such as PEP and pyruvate under the conditions studied. Based on the measurements, we calculated that about 22% of PEP in REF originated from oxaloacetate (OAA) via the PEP carboxykinase. In NOX, only 18% of PEP originated from OAA, but in ATPase, 33% of PEP originated from OAA. The remainder of PEP is derived from pentoses. This result is coherent with the decrease in the pentose phosphate flux in ATPase. We also observed substantial changes in the origin of OAA from PEP via the reverse reaction, PEP carboxylase. Relative to the REF, NOX produced 35% less OAA from PEP whereas ATPase produced 15% less (Fig. 2). The higher glycolytic flux in NOX and ATPase translated into a
higher contribution to the production of serine by 9% in NOX and by 24% in ATPase, compared with REF. Information on the origin of metabolic intermediates indicates the degree of utilization of the different pathways, and provides critical insight into regulation prevalent at the flux level. Detailed information on the drain of precursors to biomass is shown in section S2 in the supplemental text.

Response of the Central Carbon Metabolites—We measured the intracellular concentration of the central carbon metabolites and cofactors under identical conditions at which metabolic fluxes were measured. Principal component analysis separated NOX and ATPase strains from REF (see section S3 in supplemental text). The first principal component (PC1) accounted for 79% of the variance, whereas PC2 accounted for 11% of the variance. PC1 distinctly separated NOX and ATPase from REF. Given the high amount of variance that PC1 accounts for, the result indicates that the levels of the central carbon intermediates are in NOX and ATPase significantly different from that in REF. PC2 separated the strains in opposing directions from REF.

The concentration of the glycolytic metabolites and TCA cycle intermediates was higher in ATPase than in NOX. A notable exception was in the concentration of glyceraldehyde-3-phosphate (GAP), which was present at a higher concentration in NOX. The concentration of the pentose phosphates was lower in ATPase and NOX, compared with REF. The key metabolites at the divergence of metabolism from glucose metabolism, cellular respiration, and electron transport, acetyl-CoA metabolism, and the TCA cycle. The transport of metal ions, enterobactin metabolism, and ATP synthesis-coupled electron transport were induced only in ATPase (see the complete list in section S4 in supplemental text). On the other hand, genes responsible for the generation of energy from fatty acids and lipids (fatty oxidation), ATP biosynthesis and the non-oxidative branch of the pentose phosphate pathway were induced only in NOX. We observed that substantially fewer genes were repressed in ATPase and NOX strains. Only 60 genes were significantly repressed in ATPase and 70 in NOX, with an overlap of only 18 genes. Key genes in the biosynthesis of macromolecules (lipid, protein, and nucleotide) were repressed only in the NOX strain. Cellular processes related to translation and protein synthesis were commonly repressed in ATPase and NOX strains.

The genes whose expression was significantly different in the mutant strains were mapped to the transcription factors and sigma factors that control them. The network topology allowed the quantification of the transcriptional regulation exerted by the regulators on the corresponding genes. Many of the transcriptional changes observed were brought about by global TFs such as ArcA, FNR, CRP, IHF, pdhR and fur in ATPase and NOX (Fig. 4). These TFs regulate central biological processes such as respiration, catabolite repression, DNA replication, acetyl-CoA metabolism etc, indicating that these processes are commonly affected in both ATPase and NOX. Furthermore, we also identified regulators that played a significant role in only one strain whereas remaining relatively unaffected in the other. Unlike the global TFs, these regulators had a narrower realm of influence, thereby revealing the differences in regulation between the two strains at a higher resolution. Some key examples of such TFs include fadR, gadEWX, fis, kdpE, ompR, and envY, which were identified to be significant only in NOX (Fig. 4). These TFs control specific biological processes such as fatty acid metabolism, cellular respiration, and electron transport, acetyl-CoA metabolism, and the TCA cycle.
This result is in complete concurrence with our discovery that only NADH (and not ATP) appeared to be the metabolites identified by our analysis for all interaction map was used to extract significantly changed system and organic acid production. The sigma factors, translocation mechanisms mediated by formate hydrogenase (rpoS) and pius efflux, membrane integrity, etc. On the other hand, TFs such as pspF, fnr, and ATP consistently ranked among the top metabolites in NOX and ATPase, respectively.

To validate our correlation of the TFs to the specific cofactors, we overlaid the transcription response from mutants in which some of these TFs were deleted on to the metabolic network. Using transcription data from arcAΔ, furΔ or arcAΔ furΔ mutants, this converse approach identified that NADH and ATP consistently ranked among the top metabolites around which the enzymes were subjected to significant (p < 0.03) transcriptional changes, compared with an isogenic reference. The only except was the rpoSΔ mutant. Gene expression indicated that only NADH (and not ATP) appeared to be the surrounded by enzymes whose genes significantly changed. This result is in complete concurrence with our discovery that σ5 (RpoS)-mediated regulation was predominantly sensitive to NADH. The list of metabolites identified by our analysis for all the mutants is shown in section S6 of the supplemental text.

Transcriptionally Significant Metabolic Modules—A global interaction map was used to extract significantly changed metabolic modules, i.e. subnetworks of cellular components (metabolites, reactions, and genes), which were most affected by the overexpression of the soluble ATPase or NADH oxidase. Overexpressing NADH oxidase had a more widespread impact on metabolism, as indicated by the larger size of the HSN in NOX (526 nodes with 1091 interactions between them). The HSN in ATPase was much smaller (264 nodes with 631 interactions between them), but had a higher density. The clustering coefficient, which indicates the extent to which the nodes cluster together, was double for the HSN in ATPase (0.262) than that in NOX (0.135). The HSNs in ATPase and NOX are provided as markup files in the supplemental material. The top scoring metabolites in the HSN of NOX are the redox couples NAD(H), FAD(H), quinones, TCA cycle intermediates, and acetyl-CoA. The reactions involving these metabolites were identified to be transcriptionally significant in the NOX network. The reactions predominantly involved the conversion of cofactors (such as dehydrogenases in the central carbon metabolism and fatty acid oxidation) and transporters of potassium and ammonia (see section S7 in the supplemental text). The HSN of ATPase was denser and specifically involved the quinones, succinate, fumarate, oxygen, and the proton as the top scoring metabolites.

Further inspection of the HSN of NOX and ATPase indicated that proton translocation in the electron transport chain was significantly affected only in the ATPase strain (Fig. 5). Non-proton translocating NADH dehydrogenase and the TCA cycle were significantly affected only in NOX strain. Although the transhydrogenase reaction was significantly different in both NOX and ATPase, but we did not see significant changes in the gene expression (pntAB or udhA). A complete list of the significant changes is provided in supplemental material S7.

DISCUSSION

NADH and ATP effect the conversion of metabolic reactions where the products and the reactants are not balanced in terms of redox and/or energy. Because this is often the case in metabolic networks, these cofactors rank among the most highly connected metabolites in many microbial networks. Previous studies along these lines were restricted to either a single enzyme or a regulator or analyzing information from only one level of hierarchy (transcription of metabolic flux, etc). Clearly, these proteins cannot be studied in isolation nor can only one level of information comprehensively identify regulation (1). The widespread role of NADH and ATP in metabolism and cellular processes calls for a true integrative systems biology approach toward this goal.

Regulation of Metabolism by NADH and ATP—Increased glycolytic flux in ATPase or NOX raises the question of the exact control mechanism. Arguably, this is the most well studied pathway in metabolism and cellular ATP demand was shown to control glycolytic flux (7). We observed increased glycolytic flux upon lowering NADH also. It remains to be seen whether ATP and NADH share the control of glycolysis or the net outcome of the glycolytic flux is a ramification of decreased NADH lowering ATP production in the oxidative phosphorylation.

Increased glycolysis translated into increased flux in the TCA cycle in NOX and ATPase strains. When the NADH concentration decreased, the genes of the TCA cycle were induced, ensuing in increased flux. On the other hand, lowering ATP level induced only citrate synthase and succinate dehydrogenase...
ase genes. Yet the TCA cycle flux increased by a factor of 60% relative to that in REF. Thus, it can be inferred that the TCA cycle is predominantly regulated at the transcriptional level by TFs such as ArcA and also allosterically inhibited by ATP. Indeed, deleting ArcA, resulted in over 60% increase in the TCA cycle flux (9). The flux in the pentose phosphate pathway decreased in NOX and ATPase strains and did not appear to be actively regulated by cofactor availability.

Beyond the central carbon metabolism, NADH and ATP appear to work independently, and sometimes antagonistically, in regulating many other peripheral pathways. For example, decreasing ATP repressed the biosynthesis of proteins, lipids, and nucleotides and increased acetate overflow. Decreasing NADH induced the synthesis of lipids and amino acids and decreased acetate overflow. Information from S5–S7 of the supplemental text is summarized in Fig. 6, which shows the impact of NADH and ATP in various cellular processes.

**Insights from Integrated Analysis**—Perturbation in NADH level affects larger number of reactions that a perturbation in ATP level, as indicated by the size of the HSN for NOX and ATPase, respectively. A denser network (higher clustering coefficient) for ATPase indicates higher degree of transcriptional regulation by TFs and sigma factors as compared to NOX. The TCA cycle and portions of the electron transport chain are magnified from the HSNs. The interactions by regulators (transcription factors and sigma factors) and metabolites were removed from the magnified part of the figure, for clarity. Only physical interactions between proteins are retained. The genes (proteins) are shown in circles, and the reactions are shown in rectangles. The magnified portion is also shown in section S7 of supplemental text.
Systemic Response to Cofactor Perturbation

The recent progress in biotechnology did not translate into success stories in metabolic engineering because it is only now that we are able to understand the complexity of regulation. In addition to facilitating quantitative modeling, the multilevel data presented in this study will also have direct applications in metabolic engineering. The practice of metabolic engineering is moving toward engineering regulatory control (1) where hierarchical organization of global and local TFs provide a framework for the selection of appropriate targets for metabolic engineering. Recent example included the deletion of ArcA to reduce acetate overflow for recombinant protein production (33). Similarly, TFs could be deleted in appropriate combination to elicit only the desired response with minimal side effects. Our results pointed out that isocitrate dehydrogenase is a more potent producer of NADPH than the pentose phosphate pathway. Therefore, this route might be a more efficient means to increase NADPH availability. This is particularly useful in engineering metabolism of pentose sugars (34) or production of heterologous proteins or fatty acids, etc. where NADPH demand is increased.

The data sets provided in this study also open the door to understanding connections between regulatory networks, metabolic flux networks and possibly governing physicochemical states like the redox potential or the cellular energy load. Future metabolic engineering efforts for designing microbial cell factories need to take these network connectivities into account to systematically optimize cellular systems.

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REFERENCES

Systemic Response to Cofactor Perturbation