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Adaptation of anaerobic cultures of *Escherichia coli* K-12 in response to environmental trimethylamine-N-oxide

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Summary

Systematic analyses of transcriptional and metabolic changes occurring when *Escherichia coli* K-12 switches from fermentative growth to anaerobic respiratory growth with trimethylamine-N-oxide (TMAO) as the terminal electron acceptor revealed: (i) the induction of *torCAD*, but not genes encoding alternative TMAO reductases; (ii) transient expression of *frmRAB*, encoding formaldehyde dehydrogenase; and (iii) downregulation of copper resistance genes. Simultaneous inference of 167 transcription factor (TF) activities implied that transcriptional reprogramming was mediated by 20 TFs, including the transient inactivation of the two-component system ArcBA; a prediction validated by direct measurement of phosphorylated ArcA. Induction of *frmRAB*, detection of dimethylamine in culture medium and formaldehyde production when cell-free extracts were incubated with TMAO suggested the presence of TMAO demethylase activity. Accordingly, the viability of an *frmRAB* mutant was compromised upon exposure to TMAO. Downregulation of genes involved in copper resistance could be accounted for by TMAO inhibition of Cu(II) reduction. The simplest interpretation of the data is that during adaptation to the presence of environmental TMAO, anaerobic fermentative cultures of *E. coli* respond by activating the TorTSR regulatory system with consequent induction of TMAO reductase activity, resulting in net oxidation of menaquinone and inhibition of Cu(II) reduction, responses that are sensed by ArcBA and CusRS respectively.

Introduction

Trimethylamine-N-oxide (TMAO) is an osmolyte in marine organisms and is often found in anaerobic environments where it is used as a terminal electron acceptor, being reduced to trimethylamine (TMA) to support the growth of enteric bacteria (Barrett and Kwan, 1985). TMA, generated from metabolism of dietary phosphatidylcholine, L-carnitine and TMAO (from fish) by human gut microflora is transferred to the liver, where, after oxidation to TMAO by flavin monoxygenases, it promotes cardiovascular disease (Wang et al., 2011; Koeth et al., 2013). *Escherichia coli* K-12 is a metabolically versatile enteric bacterium that is able to grow in the presence and absence of O₂. Aerobic respiration, anaerobic respiration and fermentation represent its three major metabolic modes (Bock and Sawers, 1996; Gennis and Stewart, 1996; Guest et al., 1996). Anaerobic respiration with TMAO as the terminal electron acceptor conserves less energy than aerobic respiration but is more efficient than fermentation (Guest et al., 1996). TMAO is reduced to TMA by the enzyme TMAO reductase, encoded by the *torCAD* operon (Méjean et al., 1994). The expression of *torCAD* is controlled by a modified two-component system comprised of three proteins: TorT, S and R. TorT is a periplasmic TMAO-binding protein (Baraquet et al., 2006) that interacts with the membrane protein TorS to form a sensory complex capable of transmitting the signal (TMAO) across the cytoplasmic membrane to alter the behaviour of the regulator TorR (Moore and Hendrickson, 2012). Upon stimulation the TorS protein phosphorylates the regulator TorR, which then activates transcription of the *torCAD* operon (Simon et al., 1994; 1995). In the absence of TMAO, TorS acts as a TorR–P phosphatase to ensure that TMAO reductase is not produced in the absence of its substrate (Ansaidi et al., 2001). The TMAO reductase subunit (TorA) has a molybdopyranopterin cofactor and is located in the periplasm. Electrons from reduced quinones (menaquinol and dimethylmenaquinol;
Wissenbach et al., 1992) are transferred to TorA via the membrane-associated penta-heme cytochrome TorC (McCrindle et al., 2005). Thus, reduction of TMAO can be coupled to the action of various dehydrogenases that reduce the quinone pool and in this way energy is conserved chemiosmotically. Unlike other alternative electron acceptors that are used by E. coli, TMAO respiration has been reported to occur in the presence of the preferred electron acceptor O2 (Ansaldi et al., 2007).

Thus, there is a body of knowledge covering the regulation and operation of the E. coli TMAO respiratory chain, but crucially there is almost no understanding of the dynamic adaptive processes that occur during the transition from anaerobic fermentative to TMAO respiratory growth. Here, glucose-limited chemostat cultures have been used to systematically study the effects of perturbing anaerobic fermentative cultures of E. coli K-12 by addition of the terminal electron acceptor TMAO using transcript profiling, metabolite and biochemical measurements combined with probabilistic modeling of transcription factor (TF) activities to obtain a deeper understanding of the dynamics of the adaptive process.

Results and discussion

Analysis of over-metabolite production by anaerobic cultures of E. coli K-12 in the absence and presence of TMAO

Glucose-limited anaerobic fermentative growth of E. coli can be described by Eq. 1 and 2:

\[ C_6H_{12}O_6 + H_2O \rightarrow CH_3COOH + CH_3CH_2OH + 2 HCOOH \]  

(1)

\[ HCOOH \rightarrow CO_2 + H_2 \]  

(2)

In addition, other minor fermentation products (over-metabolites) such as lactate and succinate are produced. Anaerobic steady-state chemostat cultures (20 mM glucose-limited medium, dilution rate 0.2 h) of E. coli K-12 MG1655 were established, and measurement of over-metabolites present in the culture medium at steady state showed the presence of acetate, ethanol, formate, succinate, fumarate, lactate and orotate, consistent with anaerobic fermentative growth during which ~40% of the formate was converted to CO2 and H2 by formate hydrogen-lyase (Table 1; Eq. 2). The anaerobic fermentative steady state was perturbed by the introduction of TMAO to both the cultures and the medium feed. Fully anaerobic respiratory growth on glucose with TMAO as the electron acceptor is described by Eq. 3.

\[ C_6H_{12}O_6 + 4 \left(CH_3\right)3NO \rightarrow 2 CH_3COOH + 4 \left(CH_3\right)3NH_2 + 2 CO_2 + 2 H_2O \]  

(3)

| Time after addition of TMAO (min) | TMAO (mM) | Acetate (mM) | Formate (mM) | Ethanol (mM) | Lactate (mM) | Orotate (mM) | Succinate (mM) | DMA (mM) | Biomass (mg cell dry weight/ml) |
|----------------------------------|-----------|--------------|--------------|--------------|--------------|-------------|----------------|---------|-------------------------------|
| 0                                | ND        | 45.8 ± 1.4   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | ND      | 0.29 ± 0.07                   |
| 2                                | 45.0 ± 1.2 | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | ND      | 0.29 ± 0.07                   |
| 5                                | 45.0 ± 1.2 | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | ND      | 0.29 ± 0.07                   |
| 10                               | 45.0 ± 1.2 | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | ND      | 0.29 ± 0.07                   |
| 15                               | 45.0 ± 1.2 | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | ND      | 0.29 ± 0.07                   |
| 20                               | 45.0 ± 1.2 | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | ND      | 0.29 ± 0.07                   |
| 60                               | 45.0 ± 1.2 | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | 45.0 ± 1.2   | ND      | 0.29 ± 0.07                   |
| 1440                             | ND        | ND           | ND           | ND           | ND           | ND          | ND            | ND      | ND                            |

*a. The values reported for ethanol are corrected to account for losses to the gas phase.

Data are the mean values ± standard deviation (n = 3). ND, not detected.

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External milieu of the new steady state had higher acetate and orotate, but lower succinate and ethanol concentrations than those observed under fermentative conditions. The metabolite measurements indicated that TMAO was consumed and a transcriptional response had been initiated. For 60 min following the addition of TMAO, the concentration of fumarate increased, whereas the concentrations of succinate and formate decreased (Table 1). During the same period, the concentration of TMAO decreased from ∼46 mM to ∼40 mM. After 1440 min, the new TMAO-respiratory/fermentative steady state was established and TMAO was undetectable, whereas TMA reached ∼46 mM. The external milieu of the new steady state had higher acetate and orotate, but lower succinate and ethanol concentrations than those observed under fermentative conditions (Table 1). The physiological changes characterized by these metabolite measurements were accompanied by an increase in biomass (Table 1). The biomass supported by the TMAO-respiratory/fermentative steady state was twofold greater than that under fermentative conditions (0.61 g l⁻¹ compared 0.29 g l⁻¹). The dry weight of an E. coli cell is 0.28 pg (Neidhardt et al., 1990). Therefore, this increase in biomass was equivalent to an additional ∼1.2 × 10¹² bacteria l⁻¹. An electron transport chain consisting of NADH dehydrogenase I and TMAO reductase contributes 6 H⁺ to the proton motive force per NADH molecule oxidized (Keseler et al., 2013). For an H⁺/ATP ratio of 3.3 (Futai et al., 2012), 1.8 ATP molecules are produced for every TMAO molecule reduced, which equates to the production of 5 × 10²³ molecules of ATP. The Yₐₕₐₜ value for E. coli growing under anaerobic conditions at a dilution rate of 0.2 h⁻¹ is ∼5 g cell dry weight (cdw) per mol of ATP (Hempfling and Mainzer, 1975). Assuming that glucose fermentation yields 3 ATP per glucose molecule, this value is consistent with the cdw of the initial anaerobic steady-state cultures analysed here (60 mmol ATP from 20 mmol glucose yielding 0.29 g cdw equivalent to Yₐₕₐₜ = 4.8 g cdw per mol of ATP (Table 1). Therefore, the synthesis of an E. coli cell requires ∼40 × 10⁹ ATP molecules, and hence the additional ATP gained from respiration with TMAO as the terminal electron acceptor accounts for the synthesis of an additional ∼1.3 × 10¹² cells. Thus, the metabolic map shown in Fig. 1 accurately captures the physiological state of the TMAO-respiratory/fermentative steady state.

The transcriptional response of E. coli K-12 during adaptation to TMAO-respiratory conditions

Transcript profiles were obtained for E. coli K-12 cultured under anaerobic fermentative conditions and then 2, 5, 10, 15, 20 and 60 min after the addition of the electron acceptor TMAO and under the new TMAO-respiratory/fermentative steady state (1440 min). For this analysis, an operon was deemed to be significantly regulated if, at one or more of the sampling points, transcript abundance was threefold greater or smaller after the addition of TMAO compared with the initial anaerobic fermentative steady state (Table 2). In total, 34 operons were judged to be significantly regulated (29 were upregulated and 5 were downregulated).

The upregulated transcripts could be classified into six groups based on the functions of the encoded proteins (Table 2). Unsurprisingly, within 2 min of the addition of TMAO (46 mM) to the system the torCAD (TMAO reductase) operon was induced (Table 2; group A), indicating that the presence of the electron acceptor had been perceived and a transcriptional response had been initiated. The metabolite measurements indicated that TMAO reduction to TMA was initiated within 2 min (Table 1). The torCAD operon transcript exhibited maximal abundance (11.6-fold greater than in the anaerobic fermentative steady state) 10 min after initial exposure of the culture to TMAO. The enhancement in the abundance of the torCAD operon transcript was sustained into the new
Table 2. Transcripts that are altered in abundance by ≥ threefold in response to the addition of TMAO to anaerobic fermentative steady-state cultures of E. coli K-12 MG1655.

| Operon | Product                                                                 | 2 min | 5 min | 10 min | 15 min | 20 min | 60 min | TMAO respiratory steady state | Regulatory proteins |
|--------|-------------------------------------------------------------------------|-------|-------|--------|--------|--------|--------|-------------------------------|--------------------|
|        |                                                                         |       |       |        |        |        |        |                               |                    |
| **(A)** Terminal recuptase/oxidases |                                                                         |       |       |        |        |        |        |                               |                    |
| cyoA-E | Cytochrome bo' terminal oxidase                                         | 2.5   | 3.8   | 6.4    | 3.3    | 3.2    | 2.2    | 1.1                                         | ArcA (-), Cra (-), CRP (+), FNR (-), Fur (-), GadE (+), PdhR (-), NarL (-) TorR (+) |
| torCAD | TMAO reductase                                                          | 4.4   | 6.1   | 11.6   | 7.9    | 7.6    | 7.6    | 3.5                                         |                    |
| **(B)** Carbon metabolism |                                                                         |       |       |        |        |        |        |                               |                    |
| acs-yjch-actP | Acetyl-CoA synthetase, inner membrane protein, acetate transporter       | 1.7   | 3.3   | 6.1    | 3.1    | 1.8    | 1.0    | 1.2                                         | CRP (+), Fis (-), IHF (-) |
| frmRAB | Transcription regulator, formaldehyde dehydrogenase, S-formylglutathione hydrolase | 2.4   | 4.6   | 4.9    | 6.1    | 10.9   | 9.5    | 1.3                                         | FmrR (-) |
| gldAB  | Anaerobic glycerol-3-phosphate dehydrogenase                            | 2.1   | 3.5   | 2.1    | 1.3    | -1.2   | -1.4   | -1.0                                        | ArcA (-), CRP (+), Fis (+), FlhDC (+), FNR (+), Gpr (-) |
| gldC   | Aerobic glycerol-3-phosphate dehydrogenase                              | 2.9   | 4.6   | 2.8    | 1.6    | 1.2    | 1.1    | 1.2                                         | ArcA (-), CRP (+), Gpr (-) |
| pdhR-aceEF-lpd | PdhR regulator, pyruvate dehydrogenase complex                        | 1.6   | 1.7   | 2.3    | 3.1    | 2.7    | 1.6    | 1.0                                         | ArcA (-), Cra (-), CRP (+), Fis (+), FNR (+), Fur (-), NsrR (-), PdhR (-) |
| **(C)** Methionine metabolism |                                                                         |       |       |        |        |        |        |                               |                    |
| metBL  | O-Succinylhomoserine lyase, aspartate kinase                            | 1.2   | -1.3  | 1.2    | 2.4    | 5.6    | 1.2    | 1.2                                         | MetJ (-), PhoP (+), YjE (+) |
| metF   | 5, 10-Methylenetetrahydrofolate reductase                              | -1.1  | -1.5  | -1.2   | 1.5    | 3.8    | 1.1    | 1.1                                         | MetJ (-) |
| metR   | Methionine reductase                                                    | 1.0   | -1.3  | -1.2   | 1.4    | 3.2    | 1.1    | 1.0                                         | MetJ (-), MetR (-) |
| mmuPM  | S-Methyl-L-methionine transport                                         | 1.1   | 1.1   | 1.1    | 1.1    | 3.5    | 1.1    | 1.1                                         |                    |
| sbp    | Sulfite-binding protein                                                | -1.1  | 1.0   | 1.0    | 1.8    | 3.6    | 1.7    | -1.2                                        |                    |
| ybdL   | Methionine aminotransferase                                            | 1.1   | -1.6  | 1.0    | 1.8    | 4.0    | 1.0    | 1.1                                         |                    |
**Transport proteins**

| Gene   | Description                               | Transcript Abundance | Regulators |
|--------|-------------------------------------------|----------------------|------------|
| cstA   | Peptide transport                         | 2.2, 4.0, 3.6, 1.4, 1.1, 1.1, 1.6 | CRP (+)    |
| dctA   | C4-dicarboxylate/orotate : proton symport  | 2.1, 3.8, 4.3, 1.8, 1.6, 1.3, 1.6 | ArcA (-), CRP (+), DcuR (+) |
| mglBAC | Galactose transport                       | 2.1, 2.5, 3.7, 1.1, 1.0, -1.2, 2.0 | Crp (+), GalR (-), GalS (-) |
| ugpBAECQ | Glycerol-3-phosphate transport          | 1.8, 3.3, 2.4, 1.5, 1.2, 1.2, 1.2 | PhoB (+) |

**Motility and chemotaxis**

| Gene | Description | Transcript Abundance | Regulators |
|------|-------------|----------------------|------------|
| flgAMN | Flagella biosynthesis | 1.0, 1.1, -1.1, 1.1, 1.1, -1.1, 3.0 | FlhDC (+) |
| flgB-J | Flagella proteins | -1.1, -1.2, -1.1, -1.2, -1.1, 1.0, 3.3 | FlhDC (+) |
| fliAZY | Flagella regulation and biosynthesis | -1.1, -1.1, -1.3, -1.1, -1.1, 1.1, 4.0 | FlhDC (+), H-NS (+), MarA (-), NsrR (-) |
| motB-cheAW | Flagella and chemotaxis proteins | 1.1, -1.1, 1.0, 1.1, 1.0, 1.0, 3.2 | CpxR (-) |
| tar-tap-creBYZ | Chemotaxis proteins | 1.0, 1.3, 1.5, 1.1, -1.1, 1.0, 3.1 | FNR (+) |

**Miscellaneous**

| Gene | Description | Transcript Abundance | Regulators |
|------|-------------|----------------------|------------|
| betIBA | Betaine aldehyde dehydrogenase and regulatory protein | 1.8, 3.3, 2.5, 1.9, 1.6, 1.4, 1.0 | ArcA (-), BetI (-) |
| bsmA | Conserved protein | 1.7, 3.0, 1.4, 1.0, -1.2, 1.0, 1.0 |  |
| glcC | Glycolate regulator | 1.8, 3.9, 2.3, 1.4, 1.1, 1.0, 1.0 | CRA (-), CRP (+), Fis (-), GlcC, (-) |
| putA | Fused transcriptional repressor-proline dehydrogenase | 1.6, 2.8, 5.5, 2.2, 2.2, 1.7, 1.2 | BasR (-), MarA (+), PutA (-) |
| ybdH | Predicted oxidoreductase | 1.3, 1.2, 1.3, 1.9, 3.0, 1.1, 1.2 |  |

**Predominantly downregulated operons**

| Gene | Description | Transcript Abundance | Regulators |
|------|-------------|----------------------|------------|
| borD | Predicted lipoprotein | -1.8, -3.0, -1.4, 1.5, 1.8, 1.2, 1.1 | PhoP (+) |
| cusCFBA | Copper efflux | -1.1, -2.2, -3.3, -2.5, -3.0, -2.9, 1.0 | CusR (+) |
| manXYZ | Mannose PTS permease | -1.8, -1.4, -3.7, -5.0, -3.0, -2.3, -1.1 | CRP (+), CRA (-), DgsA (-), NagC (-) |
| ompF | Outer membrane protein | 1.0, -2.4, -3.1, -2.4, -1.3, 1.1, 1.7 | CpxR (-), CRP (+), EnvY (+), Fur (+), IHF (+), OmpR (+), RstA (-) |
| thrABC | Threonine and homoserine biosynthesis | -1.6, -1.6, -2.4, -3.2, -2.3, 1.0, -1.5 | DksA (+) |

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a. The data shown are for the first gene in the operon unless indicated in bold type.
b. Numbers are fold increase or decrease (by at least threefold, P ≤ 0.05, at one or more sampling points) in transcript abundance after introduction of TMAO.
c. Regulatory proteins are indicated (-) negative regulation, (+) positive regulation, (±) dual regulation.
PST, phosphotransferase system.
TMAO-respiratory/fermentative steady state but had declined to be only 3.5-fold greater than that of the initial anaerobic fermentative steady state. Nevertheless, torCAD was the only transcript, with the exception of the motility and chemotaxis transcripts discussed below, to exhibit > threefold enhanced abundance in the final TMAO-respiratory/fermentative steady-state relative to the initial anaerobic fermentative steady state (Table 2). The pattern of torCAD operon transcript abundance is consistent with an initial high expression phase of capacity building during the transition period followed by decreased expression to a level commensurate with the growth rate of the culture in the new TMAO-respiratory/fermentative steady-state.

Anaerobic conditions were maintained throughout the experiments described here, and thus it was surprising to find that the transcript encoding a terminal oxidase with low O2 affinity (cytochrome bo; Cyo) was strongly induced during the first 10 min of the transition before returning to its initial level of abundance (Table 2; group A). The assertion that the increased abundance of the cyo transcript was not caused by unintended introduction of O2 was supported by the absence of the broad transcriptional response that is observed upon transfer of E. coli cultures to micro-aerobic or aerobic growth conditions (Partridge et al., 2006; Trotter et al., 2011; Rolfe et al., 2012).

Group B consisted of seven operons involved in carbon metabolism. Like the cyoA-E operon, these were all transiently expressed during the transition to TMAO respiratory growth, returning to their initial abundances in the TMAO respiratory/fermentative steady state (Table 1; group B). These included transcripts encoding key components of central metabolism: pyruvate dehydrogenase, 2-oxo-glutarate dehydrogenase and succinate dehydrogenase, indicating that the presence of the electron acceptor TMAO invokes a shift to enhanced substrate oxidation. Also prominent in this grouping were transcripts associated with glycerol and acetate acquisition and metabolism, including both the anaerobic and aerobic glycerol-3-phosphate dehydrogenase operons and the glycerol-3-phosphate transport system (ugp operon assigned to group D). The activation of complex II (succinate dehydrogenase) and the anaerobic and aerobic glycerol-3-phosphate dehydrogenases suggests the establishment of respiratory chains with succinate and glycerol-3-phosphate as the electron donor in the presence of TMAO. Although extracellular glycerol was not detected in the NMR analysis, the decrease in extracellular succinate concentration and increase in fumarate concentration after addition of TMAO was consistent with the induction of complex II (Table 1). The activation of the betiBA operon (glycine-betaine synthesis from choline; see below) is suggestive of osmotic re-balancing in the 5 min period after addition of the osmolyte TMAO. A more sustained (> 60 min after TMAO addition) expression profile was observed for the fmmRAB operon, which encodes a regulator and enzymes involved in formaldehyde detoxification (Table 2; group B).

Group C consisted of operons associated with methionine metabolism. These transcripts were transiently increased in abundance 15–20 min following addition of TMAO to the chemostats before returning to the pre-TMAO addition level (Table 2; group C). Three of the six transiently induced operons in group C are regulated by MetJ, suggesting that during this period of enhanced biomass production the supply of S-adenosylmethionine, a molecule used in methyl transfer reactions, is restricted (Keseler et al., 2013). Group D contained peptide, C2-dicarboxylic acid/orotate, galactose and glycerol-3-phosphate transport systems (Table 2; group D).

The transcriptional profile of the new TMAO-respiratory/fermentative steady state was characterized by enhanced abundance of the flagella and chemotaxis transcripts that formed group E. The abundances of these transcripts did not change during the first 60 min of transition but were increased only in the final TMAO-respiratory/fermentative steady state (Table 2; group E).

The sixth group consisted of transiently induced transcripts encoding regulatory proteins, proteins of unknown function and proteins involved in osmotic homeostasis (e.g. betiBA).

Five operons were transiently downregulated ≥ threefold, with roles in copper resistance, threonine biosynthesis, mannose transport and outer membrane function (Table 2; group G). The decreased abundance of the cus operon, which encodes an anaerobic copper detoxification system, suggests that in the initial phases of adaptation the copper stress experienced by the bacteria was diminished.

**Simultaneous inference of TF activities**

The altered patterns of gene expression, substrate utilization and over-metabolite production described above are the result of complex interactions between signalling molecules and TFs. In addition to the changes in transcript abundance that pass an arbitrary ≥ threefold (t-test, \( P = 0.05 \)) filter (Table 2), there were widespread changes that are components of the transcriptional response that sit below this statistical filter. To capture the breadth of the transcriptional response when the anaerobic fermentative system was perturbed by introduction of the primary signal TMAO, the activities of 167 TFs that are recorded as able to directly regulate target genes in the EcoCyc database (Keseler et al., 2013) were simultaneously inferred from the transcript profiles using the open access TFINFER software (Asif et al., 2010). TFINFER is based on a state space model that employs a linear approximation (in
log space) to the dynamics of transcription and treats noise in a principled way such that the estimated TF activities are associated with confidence limits (Sanguinetti et al., 2006). Although the TFINFER model represents a simplified version of transcription, it has proved its utility in several previous studies (e.g. Partridge et al., 2007; Rolfe et al., 2012). TFs that exhibited a signal to noise ratio > 2 were deemed to have significantly altered activities following the introduction of TMAO (Fig. 2A). The inferred activities of these 20 TFs could account for many of the changes in transcript abundance shown in Table 2 because one or more TFs exhibiting altered activity are known to regulate transcription from the corresponding promoters (Table 2). A visual representation of the regulatory events (as inferred by TFINFER) during the time course is provided in Fig. 2B.

Remarkably, only two TFs (FlhDC and TorR) displayed > twofold enhanced activity in the TMAO-respiratory steady-state compared to the initial fermentative steady-state (Fig. 2A). The explanation for the enhanced FlhDC activity and consequent expression of motility and chemotaxis genes is unclear. However, the inferred activity of TorR implied that an activity ~sixfold lower than the maximal, which occurred 10 min after introduction of TMAO, is sufficient to maintain an appropriate concentration of TMAO reductase in the new steady state.

Four TFs (ArcA, CusR, CsgD and PutA) exhibited lower activity during the first 60 min of the transition before almost returning to their original activities in the TMAO-respiratory/fermentative steady state (Fig. 2A). The inferred activities of ArcA were validated by measuring the amount of phosphorylated ArcA (ArcA-P) present in the bacteria (Fig. 2B). ArcA and ArcA-P were separated by Phos-tag gel electrophoresis (Wako Pure Chemical Industries Ltd.) and detected by Western blotting using anti-ArcA serum. Quantitative densitometry of the blots was used to determine the proportion of ArcA-P in the samples. This showed that the ArcA-P decreased from the expected ~50% level in the anaerobic fermentative steady state (Rolfe et al., 2011) to 0% in the 2–20 min period after TMAO addition, before increasing to ~30% after 60 min and returning to ~50% in the TMAO-respiratory/fermentative steady state (Fig. 2B). Thus, direct measurement of the ArcA phosphorylation profile correlated well with the predicted ArcA activities inferred from the transcript profiles. The transient decrease in ArcA activity emphasizes the complexity of the ArcBA response. At least three signals have been proposed as modulators of ArcB activity: oxidized ubiquinone acts negatively; reduced menaquinone acts positively; and fermentation products, such as acetate, promote ArcB kinase activity and ArcA phosphorylation (Georgellis et al., 1999; 2001; Rodriguez et al., 2004; Bekker et al., 2010; Rolfe et al., 2011; 2012; Alvarez et al., 2013). The dynamics of ArcA activity reported here suggests that introduction of TMAO to the fermenting culture causes net oxidation of the menaquinone pool, resulting in the initial dephosphorylation of ArcA followed by restoration of ArcA activity, in the final TMAO-respiratory/fermentative steady state, perhaps mediated by enhanced production of acetate (Table 2). The modulation of ArcA activity in this way could account for the surprising transient induction of the cytochrome bo’ operon, as well as genes encoding the pyruvate, 2-oxoglutarate and succinate dehydrogenases because they are all repressed by ArcA (Table 2).

The majority (15 from 20) of responsive TFs (signal to noise ratio ≥ 2), including TorR, exhibited maximal activity during the adaptive phase of the transition before their activities returned to values close to those of the initial steady state. The greatest activities for ArgR, CRP and PurR were 5 min after the introduction of TMAO; IlvY, Lrp, MalT, MetJ and TorR after 10 min; DgsA, GalS and OxyR after 15 min; CysB, RcsAB and PhoP after 20 min; and GlpR after 60 min (Fig. 2A).

**Measurement of TorA protein and TMAO reductase activity suggests that regulation is at the level of transcription**

Measurements of the TorA protein content (Fig. 3A) and TMAO reductase activities (Fig. 3B) after the introduction of TMAO showed that the bacteria grown under fermentative conditions had low TMAO reductase activity, which had increased ~twofold after 60 min (Fig. 3B). The final ~fourfold increase in TMAO reductase enzyme activity in the new TMAO-respiratory/fermentative steady state matched the 3.5-fold increase in tor operon transcript abundance (Table 2), suggesting the absence of post-transcriptional regulation of TMAO reductase activity. Estimation of the TorA content of the bacteria by Western blotting suggested a greater overall increase in abundance than might be expected based on the increase in TMAO reductase activity, but there was a good positive correlation between enzyme activity and TorA in the sample obtained in the 10–1440 min period of the transition (Fig. 3A and B).

**Transient induction of the formaldehyde dehydrogenase FrmAB is likely to be the result of the presence of TMAO demethylase activity**

The frmRAB operon is induced by formaldehyde, and its expression is negatively regulated by FrmR, encoded by the first gene of the operon (Herring and Blattner, 2004). FrmA and FrmB proteins catalyse the glutathione-dependent conversion of formaldehyde to formate (Gutheil et al., 1992). Thus, the transient increased abundance of the frmRAB transcript upon exposure of the
A

| TF   | 0   | 2   | 5   | 10  | 15  | 20  | 60  | 1440 |
|------|-----|-----|-----|-----|-----|-----|-----|------|
| ArcA | 2.42| 0.80| 0.33| 0.26| 0.68| 1.20| 1.63| 2.25 |
| CusR | 2.96| 2.39| 0.61| 0.31| 0.46| 0.36| 0.39| 2.64 |
| CsgD | 2.13| 0.74| 0.78| 1.23| 0.51| 0.61| 0.61| 2.98 |
| PutA | 2.83| 1.20| 0.36| 0.13| 0.67| 0.88| 1.38| 1.97 |
| PurR | 0.78| 2.03| 5.60| 2.30| 0.57| 0.33| 0.37| 0.85 |
| ArqR | 1.15| 1.68| 4.48| 2.38| 0.36| 0.15| 1.08| 0.82 |
| CRP  | 0.85| 1.51| 2.86| 2.49| 0.72| 0.52| 0.48| 1.03 |
| MalT | 0.76| 2.21| 1.12| 4.53| 1.76| 0.37| 0.20| 1.33 |
| Lp   | 0.70| 1.24| 1.71| 4.17| 1.73| 0.53| 0.44| 0.57 |
| DgsA | 0.33| 0.92| 0.92| 3.28| 4.54| 2.10| 1.08| 0.40 |
| IlvY | 0.86| 1.18| 1.30| 3.20| 2.11| 0.59| 0.34| 0.70 |
| TorR | 0.19| 0.88| 1.83| 3.16| 3.04| 2.85| 2.97| 0.51 |
| MetJ | 1.36| 1.41| 2.22| 2.40| 0.72| 0.09| 0.93| 1.31 |
| GaIIS| 0.74| 0.62| 0.61| 0.88| 6.17| 1.15| 1.01| 0.60 |
| OxrR | 0.81| 0.71| 0.62| 0.92| 3.64| 1.02| 1.14| 0.75 |
| CysB | 0.79| 0.62| 0.77| 0.41| 0.99| 5.22| 2.08| 0.72 |
| RcsAB | 1.37| 0.80| 1.61| 0.85| 2.07| 2.70| 0.77| 0.16 |
| GipR | 2.04| 0.36| 0.13| 0.25| 1.17| 2.65| 3.30| 1.56 |
| PhoP | 1.07| 0.59| 0.24| 0.71| 2.01| 2.47| 1.51| 1.16 |
| FhNDC| 0.87| 0.57| 0.72| 0.58| 0.89| 0.64| 0.85| 11.51|

B

In fold-change

-2.0  0  2.0
0  5  10  15  20  25  30  35  40  45  50  55  60
time after TMAO addition (min)

C

ArcA~P
ArcA
cultures to TMAO was unexpected (Table 2). Therefore, confirmation of frmRAB induction by TMAO was sought by quantitative reverse transcription polymerase chain reaction (qRT-PCR) analyses of RNA isolated from batch cultures of E. coli MG1655 after 60 min exposure to different concentrations (0–40 mM) of TMAO. The data showed that the abundance of the frmRAB transcript was significantly increased when the bacteria were exposed to ≥5 mM TMAO (Fig. 4). Hence, these experiments confirmed the transcriptional response detected by the transcript profiling and showed that this adaptation occurred in both continuous and batch cultures and that frmRAB induction occurred at TMAO concentrations ≥5 mM (Table 2; Fig. 4). Incubation of E. coli K-12 cell-free extracts with TMAO (40 mM) allowed the detection of formaldehyde using the Nash reagent (Fig. 5A). Control reactions lacking cell-free extract or TMAO failed to accumulate formaldehyde.

Furthermore, closer inspection of the extracellular metabolite profile of the chemostat cultures revealed the presence of up to 0.23 mM dimethylamine (DMA) after the addition of TMAO (Fig. 4). These observations suggest that E. coli K-12 possesses TMAO demethylase activity, resulting in the formation of sub-millimolar DMA and formaldehyde, the latter accounting for the induction of frmRAB expression, which is known to be induced when E. coli cultures are exposed to 0.25 mM formaldehyde (Herring and Blattner, 2004); a concentration very similar to the amount of DMA detected in the cultures analysed here (Table 1).

The detection of formaldehyde when cell-free extracts were incubated with TMAO and the induction of the frmRAB operon during transition to TMAO respiratory growth suggested that the inability of an frmRAB mutant to detoxify formaldehyde would compromise its fitness when exposed to TMAO. Therefore, an isogenic E. coli MG1655 frmRAB mutant (JRG6703) was created along with a plasmid (pGS2486) to express frmRAB under the control of its own promoter to complement the mutant. The numbers of colony-forming units (cfu) present in anaerobic cultures of the parent, mutant and complemented mutant before and after incubation with TMAO were measured. The results showed that the cfu values of the parent and complemented mutant cultures increased

Fig. 2. Changes in TF activities in response to perturbation of anaerobic fermentative cultures of E. coli by switching to TMAO-respiratory/fermentative metabolism. (A) The output from the TFINFER software (Asif et al., 2010) at the indicated times (minutes) after TMAO (46 mM) addition to anaerobic fermentative steady-state cultures. Each cell is coloured to provide a visual representation of the TF activities inferred from the complete transcript profile dataset as follows: TF activity 0–1, blue; 1–2, green; 2–3, yellow; 3–4, orange; >4, red. In all cases, the signal to noise ratio was ≥2. (B) Graphical representation of regulatory dynamics. The figure shows the time course of all 1381 genes used in the TFINFER analysis; time is re-scaled to have equal spacing between time points. The panels above and below the time course show the three TFs exhibiting the greatest fold increase (red) or decrease (blue) in activity compared with previous time point as inferred by TFINFER, providing a depiction of the main regulatory events. (C) Validation of inferred ArcA activity by measurement of ArcA–P. A typical immunoblot of a Phos-tag gel developed with anti-ArcA serum. Lane 1, His-tagged ArcA (1.2 μg) phosphorylated in vitro by incubation with carbamoyl phosphate (50 mM); lane 2, His-tagged dephosphorylated ArcA; lanes 3–10, whole cell samples taken at t = 0, 2, 5, 10, 15, 30, 60 and 1440 min after the addition of TMAO to the anaerobic fermentative steady-state culture. The locations of ArcA and ArcA–P are indicated.

Fig. 3. The amount of TorA protein and TMAO reductase activity during adaptation to TMAO-respiratory growth correlate with the changes in tor operon transcription. (A) Typical Western blot of whole cell samples taken from anaerobic fermentative chemostat cultures 0, 2, 5, 10, 15, 30, 60 and 1440 min after the addition of TMAO developed with anti-TorA serum. (B) Quantitative densitometric analysis of the Western blot shown in A to measure the increase in the amount of TorA protein present in the bacteria after TMAO addition (open bars). The amounts of TorA shown are relative to the level in the final sample (100%). TMAO reductase activities (filled bars) were measured for duplicate samples obtained at the same time points used for quantification of TorA protein (see above). The TMAO reductase activity data are also shown relative to the activity in the final sample (100%).

Fig. 4. The frmRAB operon is induced after exposure of E. coli MG1655 to concentrations of TMAO ≥5 mM. Anaerobic batch cultures were exposed to different concentrations (0–40 mM) of TMAO. After 30 min, total RNA was isolated for qRT-PCR of the frmR mRNA. The data shown and the mean and standard deviation for the fold increase relative to the 0 mM TMAO culture and are typical of three independent experiments.
after 60 min exposure to 40 mM TMAO, but the cfu value of the strain did not (Fig. 5B). This result is consistent with production of potentially toxic concentrations of formaldehyde, which requires a regulatory response to prevent inhibition of growth during the transition to anaerobic respiration with TMAO as the terminal electron acceptor.

TMAO/TMAO reductase competes with Cu(II) for electrons from the anaerobic electron transport chain

The inferred decrease in activity of the TF CusS implied that after introduction of TMAO, the exposure of the E. coli cultures to Cu(I) was diminished because the signal for the cognate sensor CusS is Cu(I). The trace element Cu(II) can be reduced to Cu(I) (E_m + 340 mV) by E. coli using electrons transferred to ubiquinone by NADH dehydrogenase II (E_m NAD+/NADH, −320 mV; ubiquinone/ubiquinol + 110 mV) under aerobic conditions (Thauer et al., 1977; Volentini et al., 2011). Under anaerobic conditions, menaquinone (E_m menaquinone/menaquinol, −74 mV) is the electron donor for the TMAO reductase (Wissenbach et al., 1992) but like ubiquinone has the potential to reduce Cu(II). Therefore, the capacity of E. coli K-12 MG1655 to reduce Cu(II) was tested under anaerobic conditions in the presence and absence of TMAO. Under both conditions Cu(II) reduction was detected, but this was diminished ~1.6-fold in the presence of TMAO (Fig. 6). In combination with the data described above, the Cu(II) reduction experiments suggest that under anaerobic fermentative conditions there is a relatively low flux of electrons from the menaquinone pool to fumarate reductase to reduce fumarate to succinate (Table 1), allowing electrons to leak directly or indirectly from reduced quinones to periplasmic Cu(II) resulting in the production of Cu(I) and activation of the CusSR transcriptional response (Table 2; Fig. 7A). Supplying the electron acceptor TMAO to the culture offers an additional outlet for the electrons in the menaquinone pool resulting in decreased reduction of Cu(II) to Cu(I) by the electron transport chain during the initial adaptation phase (Fig. 6; Fig. 7B). In this way, CusR is inactivated resulting in decreased abundance of the cus operon transcript (Fig. 2A; Table 2). Furthermore, a shift towards a more oxidized menaquinone pool in the presence of excess TMAO would contribute to the inactivation of ArcA and hence de-repression of the cyaA-E, pdhR-aceEF-lpd and sdhCDAB-sucA-D operons (see above; Fig. 2, Table 2).
Fig. 7. Excess TMAO activates TorTSR, inhibits Cu(II) reduction by the electron transport chain and permits activation of ArcA. (A) In the absence of a non-endogenous electron acceptor (anaerobic fermentation), fumarate is reduced to succinate in the reductive arm of the anaerobic Krebs cycle and the trace element Cu(II) is reduced to Cu(I) both at the expense of menaquinol (MQH2). The Cu(I) generated by this process is sensed by the CusSR two-component system, and the copper efflux system CusFCBA is activated. Flux through the electron transport chain is relatively low, and the MQH2 : MQ ratio is high resulting in the phosphorylation of ArcA (ArcA-P) and repression of the cyoA-E, pdhR-aceEF-lpd, and sdhCDAB-sucA-D operons. (B) During the acute phase of adaptation to the presence of excess TMAO is sensed by the TorTSR sensor-regulator system, which activates expression of the torCAD operon encoding TMAO reductase. TMAO reductase effectively competes for the electrons of the menaquinone pool such that the MQH2 : MQ ratio is low, which decreases phosphorylation of ArcA resulting in de-repression of the cyoA-E, pdhR-aceEF-lpd and sdhCDAB-sucA-D operons. This could permit the establishment of a respiratory chain with succinate as the electron donor and TMAO as the electron acceptor, consistent with the observed transcriptional profile and the concentrations of succinate and fumarate in the culture medium. The flux of electrons to TMAO lowers the capacity of the bacteria to reduce Cu(II) to Cu(I), and hence the TF CusR is switched off. The width of the regulatory arrows is an indication of the relative rate of each step, and font size is used to indicate the NADH : NAD+ and MQH2 : MQ ratios.

Conclusion

The experiments described above have provided new insights into the dynamic adaptive processes that occur when E. coli switches metabolic mode from anaerobic fermentative growth to the energetically more efficient anaerobic respiratory growth with TMAO as the terminal electron acceptor (Fig. 7). As well as predictable changes in the transcript and metabolite profiles, such as the induction of the torCAD operon and the reduction of TMAO to TMA, some unexpected components of the adaptive process were revealed. It will be of interest to perform similar experiments with mutants lacking some of the key regulators (e.g. ArcA, FhDC, FrmR and MetJ) identified here to further characterize the roles played by their respective regulons in adaptation to TMAO exposure.

The detection of DMA and formaldehyde suggests that E. coli possesses TMAO demethylase activity that produces physiologically significant amounts of formaldehyde in environments in which TMAO is abundant. The formaldehyde generated by this activity is sufficient to induce the expression of the formaldehyde detoxifying FrmAB enzyme when TMAO is in excess. Work is ongoing to attempt to identify the source of the TMAO demethylase activity. The transient inactivation of ArcA predicted by probabilistic modelling of the transcript profiling data was validated by direct measurement of ArcA-P and provided an explanation for the surprising induction of the cyoA-E transcript that encodes a low O2-affinity terminal oxidase despite maintaining anaerobic culture conditions throughout the experiments. In addition to cyoA-E, the induction of several other ArcA-P-repressed operons could be accounted for by the temporary inactivation of ArcA. As stated above, the ArcBA two-component system acts as an indirect sensor of O2 by ArcB monitoring the redox state of the electron transport chain and the production of fermentation products. Thus, ArcBA activity is sensitive to cellular NADH/NAD+ ratios (Holm et al., 2010). It has been shown that glucose-limited chemostat cultures of E. coli are affected by the availability of electron acceptors fumarate, nitrate and oxygen, which act as effective NADH sinks. The greatest NADH/NAD+ ratios (∼0.75) were observed during anaerobic fermentative growth, and lower ratios were measured when an external environmental electron acceptor (∼0.3 with 7.5 mM nitrate; ∼0.4 with 70 mM fumarate; and ∼0.08 under fully aerobic conditions) was available (de Graef et al., 1999; Alexeeva, 2000). Addition of excess TMAO to the anaerobic fermentative culture would be expected to result in a decreased NADH/NAD+ ratio by providing an outlet for the electrons held in the menaquinone pool during the initial phase of adaptation, resulting in transient dampening of ArcBA activity, which is then counterbalanced by increased production of acetate as TMAO-respiratory metabolism is established. Competition for electrons from the menaquinone pool could also account for the downregulation of copper resistance genes in the presence of excess TMAO. Previous work has shown that electrons from the E. coli respiratory chain can reduce Cu(I) to Cu(II) (Volentini et al., 2011). The demonstration that TMAO was able to inhibit reduction of Cu(II) by E. coli suggests that TMAO reductase/TMAO are effective competitors for electrons that might otherwise have contributed to the reduction of Cu(II). The lower production of Cu(I) would result in the inactivation of CusR and downregulation of copper resistance genes. Thus,
systematic analysis of E. coli cultures has provided new insights into the dynamics of transcriptional and physiological changes required to successfully adapt to anaerobic environments containing TMAO.

Experimental procedures

Bacterial strains and growth conditions

Escherichia coli K-12 MG1655 and the isogenic frmRAB mutant (JRG6703 created using the red recombineering; Datsenko and Wanner, 2000), along with strains transformed with either the plasmid pGS2486 to express frmRAB under the control of its own promoter or the corresponding vector control (pBR322), were used in this study.

Glucose-limited steady-state continuous cultures of E. coli MG1655 were established in a 2 l Labfors chemostat (Infors-HT, Bottmingen, Switzerland) at 37°C, a 950 ml working volume, a stirring rate of 400 r.p.m., a dilution rate of 0.2 h⁻¹, maintained at 6.9 by titration with 1 M NaOH. Steady-state fermentative cultures were perturbed by addition of 1.5 mM TMAO to the medium feed and directly into the chemostat. Samples were removed from the chemostat at stated times (in minutes) after the addition of TMAO. For some experiments, E. coli strains were grown as batch cultures in rich medium [Luria–Bertani (LB) – yeast extract, 5 g l⁻¹; tryptone, 10 g l⁻¹; NaCl, 5 g l⁻¹] with supplements as indicated.

NMR metabolite measurements

Extracellular carbon metabolite concentrations were determined as described previously (Rolfe et al., 2011; Trotter et al., 2011). Bacteria were removed by passing chemostat culture samples through 0.22 μm pore size filters. The cell-free samples were analysed in 5 mm NMR tubes containing 450 μl supernatant, 50 μl D₂O and 1 mM (final concentration) trimethylsilyl propanoic acid (TSP) as an internal standard. Spectra were acquired on a Bruker DRX500 spectrometer (Bruker UK) at 298 K with 5 mm TXI probe, tuned to 1H at 500 MHz. The H₂O signal was reduced by pre-saturation for 2 s applied during the recycle time while carbon decoupling was applied during acquisition to suppress 13C satellites. Spectra were processed and peaks quantified by integration using Topspin (Bruker UK) while chemical shifts and metabolite concentrations were established by reference to the TSP peak.

Transcriptional profiling

Transcriptional profiling was carried out in a reference style (Rolfe et al., 2011) for three biological replicates. Briefly, samples were removed from the chemostat directly into RNA protect (Qiagen) before purification of total RNA using the RNeasy mini kit (Qiagen). Genomic DNA (2.0 μg) was converted to Cy5-labelled cDNA, and purified RNA (16 μg) (from each time point) was converted to Cy3-labelled cDNA. These were combined and hybridized overnight to an oligonucleotide microarray obtained from Agilent Technologies (AMADID 029412). Quantification of cDNA samples, hybridization to microarrays, microarray processing and scanning were carried out as described in the Fairplay III labelling kits (Agilent Technologies, 252009, Version 1.1) and scanned with a high-resolution microarray scanner (Agilent Technologies). GENESPRING GX v7.3 (Agilent Technologies) was used for data normalization and analysis. Transcripts exhibiting > threefold change in abundance after t-test with a P-value of < 0.05 at one or more of the time points analysed were deemed to be differentially regulated. The data are available in ArrayExpress E-MTAB-2784. The relative activities of TFs were inferred using the software package TFINFER as previously described (Asif et al., 2010).

qRT-PCR

Anaerobic cultures were grown overnight at 37°C in anaerobic Evans medium buffered with 50 mM phosphate buffer, pH 7.0, and supplemented with 20 mM glucose. The amounts of frmR mRNA in samples of total RNA (100 ng; isolated as described above) after exposure of these cultures to different concentrations of TMAO for 30 min were determined on an RT-PCR plate in a Mx3005P Thermocycler (Agilent Technologies) using Brilliant III Ultra-Fast SYBR Green qRT-PCR Master Mix kit (Agilent Technologies) according to the manufacturer’s instructions. The housekeeping gene gyrA was used for normalization, and a genomic DNA dilution series was used to correct for differences in primer amplification efficiencies. The sequences of the primer sets were: frmR CTGGAAACGGTGCCTGGAG, ATCAGCCATTGCGCG; gyrA ACCCTGGCGAGAAATTACACC, AATGACCGACAT CGCATAATC.

Measurement of ArcA phosphorylation

The degree of ArcA phosphorylation was measured using Phos-tag-acrylamide gel electrophoresis and subsequent Western immunoblotting. Chemostat culture samples were collected directly into a final concentration of 1 M formic acid as previously described (Rolfe et al., 2011). Samples were then processed as described elsewhere (Barbieri and Stock, 2008) using OD₆₀₀ measurements to calculate volumes of sample to be resolved on a 37.5 μM Phos-tag-10% acrylamide gel. Standard wet transfer to a Hybond-C membrane (Amersham) was followed by immunodetection using rabbit ArcA antiserum and the ECL detection system (GE Healthcare). ArcA protein was purified and phosphorylated in vitro as previously described (Bekker et al., 2010) with the exception of using 100 mM carbamoyl phosphate.

Detection of TorA protein by Western immunoblotting

Chemostat culture samples were collected directly into a final concentration of 50 mg ml⁻¹ chloramphenicol. Aliquots of
suspension were pelleted by centrifugation (10 000 r.p.m. in a microcentrifuge for 3 min) and pellets stored at −80°C until required. Samples were standardized with respect to their OD_{600} value by dilution with water. The samples were boiled in SDS-PAGE loading buffer for 15 min prior to being resolved on a 10% acrylamide gel followed by electroblotting onto Hybond-C membrane (Amersham). TorA protein was detected using rabbit TorA antiserum (1:1000) and visualized using the ECL system (GE Healthcare) according to manufacturer’s instructions.

**TMAO reductase assays**

Methyl viologen-linked reductase assays were carried out with intact cell samples from the chemostat. Cell samples were washed and re-suspended in 25 mM sodium phosphate buffer pH 7.0 to OD_{600} ~6. Reductase assay measurements were carried out as previously described (Hitchcock et al., 2010), except 0.2 mM methyl viologen was used. Duplicate assays were carried out in screw-topped glass cuvettes fitted with silicon seals in a total volume of 1 ml in a CARY 50 UV-visible spectrophotometer (Agilent Technologies) pre-warmed to 37°C. All buffers and reagents were made anaerobic plus 40 mM TMAO (in TMAO-containing samples) in a 1 ml total volume of citrate phosphate buffer, pH 7.0, at 37°C. Control assays with no cells and no CuSO_{4} were carried out in parallel. At the indicated times, the assays were stopped by pelleting the cells and 0.1 mM bathocuproine disulphonate (Sigma-Aldrich) was added to the resulting supernatant. The absorbance at 412 nm for the supernatant was measured and the molar extinction coefficient for the Cu(I) complex (12 220 M cm⁻¹) used to determine Cu (II) reduction by the samples per OD_{600} unit.

**Measurement of formaldehyde production**

Formaldehyde was measured using acetylacetone reagent (Nash, 1953). Preparation of cell samples and setting up of assays were performed in an anaerobic workstation. Anaerobic cultures of *E. coli* MG1655 were grown overnight with stirring at 37°C in LB supplemented with 0.2% glucose. Cells were pelleted, washed in 25 mM sodium phosphate buffer pH 7.0 and re-suspended in a final volume of 2 ml anaerobic cell breakage buffer. An oxygen scavenger (10% glucose, 10 μg ml⁻¹ catalase and 5 μg ml⁻¹ glucose oxidase), 1 ml, was added to the re-suspended cells before sonication (15 s, 15 micron amplitude). Any unbroken cells were removed by centrifugation at 5500 r.p.m. in a benchtop microcentrifuge for 1 min. Cell-free extracts were added to a 1 ml total volume of 25 mM sodium phosphate buffer and 40 mM TMAO (in TMAO-containing samples only). Assays were incubated anaerobically at 37°C and at the indicated times were stopped by precipitation using 5% trichloroacetic acid and centrifugation. Resulting supernatants were treated as previously described (Fu et al., 2006). Formaldehyde content was calculated using the extinction coefficient at 412 nm (8000 M cm⁻¹). Protein content of cell-free extracts was determined using the Bio-Rad assay.

**Cell viability in the presence of TMAO**

Anaerobic cultures were grown overnight at 37°C in anaerobic Evans medium buffered with 50 mM phosphate buffer, pH 7.0, and supplemented with 20 mM glucose in 7 ml bijoux. After 18 h growth, a 10% inoculum of the culture was added to 7 ml Evans medium in an anaerobic workstation, and pre-treatment viable counts were measured by serial dilution in medium. The 7 ml culture was split in half (2 × 3.5 ml), and 40 mM TMAO was added to one of the subcultures. The cultures were incubated in parallel for a further 1 h at 37°C before post-treatment viable counts were measured. Plated cells were grown overnight at 37°C on LB agar and cfu per millilitre calculated.

**Copper (II) reduction assays**

*Escherichia coli* MG1655 cells were grown anaerobically overnight at 37°C in Evans medium (pH 7.0) supplemented with 20 mM glucose and 40 mM TMAO. After 18 h, cells were pelleted and washed three times in anaerobic Evans medium, pH 7.0. Cells were re-suspended in Evans medium plus 20 mM glucose to an OD_{600} ~25. The assays were prepared as previously described (Volentini et al., 2011) in an anaerobic workstation. Briefly, 100 μl of washed cells were incubated with 0.05 mM CuSO_{4}, 40 mM TMAO (in TMAO-containing samples) in a 1 ml total volume of citrate phosphate buffer, pH 7.0, at 37°C. Control assays with no cells and no CuSO_{4} were carried out in parallel. At the indicated times, the assays were stopped by pelleting the cells and 0.1 mM bathocuproine disulphonate (Sigma-Aldrich) was added to the resulting supernatant. The absorbance at 412 nm for the supernatant was measured and the molar extinction coefficient for the Cu(I) complex (12 220 M cm⁻¹) used to determine Cu (II) reduction by the samples per OD_{600} unit.

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