C₄-dicarboxylate metabolons: Interaction of C₄-dicarboxylate transporters of

Escherichia coli with cytosolic enzymes and regulators

Christopher Schubert and Gottfried Unden

Institute for Molecular Physiology (IMP), Microbiology and Wine Research; Johannes Gutenberg-University, Mainz, Germany

Address for correspondence:

Gottfried Unden

Institute for Molecular Physiology (IMP), Microbiology and Wine Research

Hanns-Dieter-Hüsch-Weg 17, BZII

55128 Mainz, Germany

unden@uni-mainz.de

Phone +49-6131-3923550
**Abstract**

Metabolons represent the structural organization of proteins for metabolic or regulatory pathways. Here the interaction of enzymes fumarase FumB and aspartase AspA with the C4-DC transporters DcuA and DcuB of *Escherichia coli* was tested by a bacterial two-hybrid (BACTH) assay *in situ*, or by co-chromatography (mSPINE). DcuB interacted strongly with FumB and AspA, and DcuA with AspA. The *fumB-dcuB* and the *dcuA-aspA* genes encoding the respective proteins are known for their colocalization on the genome and the production of co-transcripts. The data consistently suggest the formation of DcuB/FumB, DcuB/AspA and DcuA/AspA metabolons in fumarate respiration for the uptake of L-malate, or L-aspartate, conversion to fumarate and excretion of succinate after reduction. The DcuA/AspA metabolon catalyzes L-Asp uptake and fumarate excretion in concerted action also to provide ammonia for nitrogen assimilation. The aerobic C4-DC transporter DctA interacted with the regulator EIIA\textsuperscript{Glc} of the *E. coli* glucose phosphotransferase system. It is suggested that EIIA\textsuperscript{Glc} inhibits C4-DC uptake by DctA in the presence of the preferred substrate glucose.

**Introduction**

Versatile bacteria such as *Escherichia coli* co-ordinate metabolism and cellular physiology on various levels including gene expression and post-translational regulation for optimal adaption to current growth conditions. For stabilization and improvement of linear pathways, the enzymes and proteins can be organized in metabolons. Metabolons represent weak complexes of enzymes catalyzing consecutive reactions in order to transfer intermediates between the enzymes of pathways. Channeling prevents the release of labile or toxic intermediates, supports higher local concentration of the intermediates and prevents drifting to other pathways. This organization is able to increase metabolic efficiency and provide an opportunity to control the flux. High-throughput approaches have been used to identify complexes of weakly interacting proteins, or ‘interactomes’, in microorganisms, including proteins linking membrane processes to cytoplasmic reactions (Butland *et al.* 2005; Hu *et al.* 2009). However, the general approaches provide limited information on complexes when partner proteins are present in low concentrations or are membrane-bound. Thus, established regulatory complexes of the membrane-integral C4-dicarboxylic acids (C4-DC) sensor kinase DcuS with the coregulatory C4-DC carriers DctA, DcuB or DauA (Kleefeld *et al.* 2009b; Witan *et al.* 2012; Karinou *et al.* 2013; Unden *et al.* 2016) and others were not identified.
Alternatively, (Moraes and Reithmeier 2012) suggested an alternative approach for identifying gene clusters of *E. coli* for metabolons consisting of membrane integral transporters together with cytosolic enzymes for the subsequent pathway. The overview (Moraes and Reithmeier 2012) lists polycistronic operons encoding on the one hand transporters for the uptake of sugars, sugar derivatives, amino acids, C4-DC, and other substrates, and on the other hand, enzymes or regulators for subsequent pathways. For some systems interaction of the proteins has been demonstrated, such as for AmtB-GlnK (ammonia transporter and PII-type regulator GlnK) (van Heeswijk *et al.* 1996; Coutts *et al.* 2002). Remarkably, for most of the genetically co-localized systems interaction of the transporter/enzyme pairs is hypothetical.

The list of potential metabolons (Moraes and Reithmeier 2012) also includes the operons *fumB-dcuB* encoding fumarase FumB and the C4-dicarboxylate transporters DcuB, *dcuA-aspA* encoding the C4-dicarboxylate transporters DcuA and aspartase AspA (Six *et al.* 1994; Golby *et al.* 1998b), and *ttdAB-ttdT* encoding tartrate dehydratase TtdAB and the L-tartrate transporter TtdT (Kim and Unden 2007). Both *dcuA* and *dcuB* are transcribed independently from *fumB* and *aspA*, respectively, but also as *fumB-dcuB* and *dcuA-aspA* cotranscripts (Golby *et al.* 1998b). DcuB is produced only under anaerobic conditions (Zientz *et al.* 1998b; Golby *et al.* 1999) and catalyzes the uptake of L-malate which is then dehydrated by FumB to fumarate (Engel *et al.* 1994; Six *et al.* 1994) (Fig. 1A). Fumarate is reduced to succinate by fumarate reductase in fumarate respiration. Succinate is not further catabolized anaerobically and excreted by DcuB via a L-malate/succinate antiport. DcuB also catalyzes the uptake of fumarate which is used directly in fumarate respiration. In the same way, L-Asp is a substrate for transport by DcuB and used after deamination by AspA to fumarate for fumarate respiration. In any case, the succinate is excreted by C4-DC/succinate antiport with C4-DC indicating any of the C4-DCs L-malate, fumarate, or L-aspartate (Engel *et al.* 1994; Six *et al.* 1994; Janausch *et al.* 2002; Schubert *et al.* 2021).

DcuA on the other hand is expressed constitutively under aerobic and anaerobic conditions (Golby *et al.* 1998b). DcuA is able to substitute DcuB in anaerobic growth (Six *et al.* 1994) and catalyze an C4-DC/succinate antiport for fumarate respiration (Fig. 1A). L-Asp, fumarate or L-malate are accepted as the C4-DCs, but with a clear preference for L-Asp over L-malate or fumarate (Strecker *et al.* 2018). In addition to its role in catabolism and fumarate respiration, DcuA catalyzes the uptake of L-aspartate for anabolism (Strecker *et al.* 2018; Schubert *et al.* 2020). Under these conditions (Fig. 1B) L-aspartate is deaminated in bacteria by AspA to
fumarate and ammonia, which can saturate the complete nitrogen requirement for growth. The uptake of L-Asp depends on DcuA and is coupled to the excretion of fumarate (and some L-malate). The combined action of DcuA and AspA results in the net uptake of ammonia, which is able to supply all the nitrogen for cell synthesis without consuming the carbon, provided that other carbon sources are available.

L-Tartrate that is taken up by TtdT for tartrate fermentation (Fig. 1A), has to be dehydrated by dehydratase TtdAB to oxaloacetate (Reaney et al. 1993; Kim and Unden 2007). After conversion to fumarate, it is used in fumarate respiration, and the succinate is excreted by TtdT in antiport against L-tartrate.

In the metabolic pathways, DcuA, DcuB and TtdT are closely linked to AspA in case of DcuA and DcuB, to FumB in the case of DcuB, and to TtdT in the case of TtdT. To test the metabolon hypothesis for the C4-dicarboxylate metabolism, interaction of DcuA and DcuB with AspA and FumB was tested. Interaction was tested in vivo by the adenylate cyclase-based bacterial two-hybrid (BACTH) system which has been established as a reliable assay for protein-protein interactions involving membrane-integral proteins, or in vitro by co-purification of the proteins.

Fig. 1: Pathways of (A) fumarate respiration of *E. coli* using fumarate, L-malate, and L-aspartate as the electron acceptors, (B) L-tartrate fermentation and (C) nitrogen assimilation from L-aspartate. For details on (A) and (B) see (Unden et al. 2016) and on (C) (Schubert et al. 2020). Abbreviations: AspA, aspartase; DcuA, C4-DC transporter DcuA; DcuB, C4-DC transporter DcuB; ET, electron transport; FrdABCD, fumarate reductase; Fum, fumarate; FumB, fumarase B; L-Asp, L-aspartate; L-Mal, L-malate; MKH₂, menaquinol; Succ, succinate; TtdT, L-tartrate transporter.
Results and Discussion

DcuB interacts with AspA and FumB.

The adenylate cyclase-based bacterial two hybrid (BACTH) system (Karimova et al. 1998; Karimova et al. 2001) was used to investigate the interaction of AspA or of FumB with DcuB. The BACTH system relies on the restoration of Bordetella pertussis adenylate cyclase (AC) that has been separated genetically in two domains T18 and T25. When the domains are fused to interacting proteins, local vicinity of the domains allows restoration of AC activity due to high flexibility of domain arrangement of B. pertussis AC. The activity is tested in a reporter strain deficient of the E. coli AC CyaA by expression of the cAMP-CRP dependent β-galactosidase LacZ. The N- and C-termini of DcuB (similar to those of DcuA and DcuC) are located in the periplasm, which excludes the use of the BACTH system under standard conditions since cAMP-CRP resides in the cytoplasm. Therefore, the T25 fragment of the AC was fused for the interaction assays into a cytosolic loop of DcuB (or the other Dcu transporters) (Bauer et al. 2011; Witan et al. 2012; Wörner et al. 2016; Strecker et al. 2018). DcuB with the sandwich (SW) T25 domain (DcuBSWT25) is membrane-integral and active in co-regulating DcuS (Wörner et al. 2016) allowing interaction studies. Bacteria producing the pairs DcuBSWT25 with FumBT18, and DcuBSWT25 with AspAT18 showed high β-galactosidase activity (Fig. 2) which exceeded the negative control represented by the non-interacting proteins T18AspA and T25Zip. The interaction was independent of the presence of transport substrate molecules L-malate and L-aspartate (not shown). On the other hand, bacteria producing the sandwich construct DcuCST25 and FumBT18, or AspAT18 respectively, showed only low β-galactosidase activity, which exceeded the background activity only slightly (Fig. 2). DcuC serves as a C4-DC efflux transporter (Zientz et al. 1996; Zientz et al. 1999; Janausch et al. 2002) for succinate derived from fermentation metabolism and not from externally supplied C4-DCs. The data indicates specific protein-protein interaction of DcuB with FumB and with AspA which is not observed for the DcuC efflux transporter that has a different metabolic task.
**DcuA interacts with AspA.**

The role of DcuA in L-Asp uptake for fumarate respiration in catabolism, and as a source of nitrogen suggests a close metabolic link to aspartase AspA. C- and N-termini of DcuA are located in the periplasm (Golby *et al.* 1998a) which prevents their use for fusion to the T18 and T25 AC domains for the BACTH system. Insertion of the T25 domain into a cytosolic loop of DcuA transporter in a sandwich fusion (DcuASWT25) produced a fusion protein in analogy to DcuBSWT25. The fusion protein was located in the membrane but was not active in transport (Strecker *et al.* 2018) or any other biological activity, which makes its application in the BACTH assay questionable. In an alternative approach, DcuA was fused to the bacterial alkaline phosphatase PhoA. PhoA-tagged membrane protein like DcuB often retain biological activity (Bauer *et al.* 2011) and the tag can be used for immunodetection with PhoA antisera. In the

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**Fig. 2: Interaction of DcuB and DcuC with AspA and FumB tested in the BACTH system.** *E. coli* BTH101(ΔcyaA) was co-transformed pairwise with plasmids encoding T25 sandwich constructs with DcuB (DcuBSWT25) or DcuC (DcuCSWT25) and fusions of FumB or AspA to T18 (FumB18, and AspA18). The combinations are shown on the x-axis. The leucine zipper pair T18Zip and T25Zip are applied as positive control (Karimova *et al.* 1998; Karimova *et al.* 2001). The pair T18Zip/T25AspA represents a negative control for background β-galactosidase activity. The corresponding plasmids are derivatives of pUT18 (FumB18 and AspA18) and pKNT25 (DcuBSWT25) (Table 1). Bacteria were grown anaerobically in LB medium. β-Galactosidase activities were quantified in Miller-Units MU (Miller 1992).
experiment, *E. coli* strain C43 was co-transformed with plasmids encoding AspA-Strep and DcuA-PhoA fusions. Proteins were simultaneously overexpressed and the bacteria were treated with formaldehyde after expression to crosslink proteins that are located in close proximity (Sutherland et al. 2008). Detergent extracts of the membranes were then applied to a Strep-Tactin column. After a wash step, the specifically bound proteins were eluted with desthiobiotin. A column treated with the extracts from bacteria that produced DcuA-PhoA and AspA-Strep eluted DcuA-PhoA and AspA-Strep from the column upon addition of desthiobiotin (Fig. 3). DcuA-PhoA was missing, however, in the eluate when either AspA-Strep or DcuA-PhoA were not produced by the bacteria. The experiment demonstrates that interaction with AspA-Strep is required to retard DcuA to the Strep-Tactin column. In addition, binding of DcuA to AspA-Strep depended on formaldehyde cross-linking in the cell homogenate (not shown), supporting the notion that the interaction is too weak to survive column chromatography without stabilization by crosslinking.

![Fig. 3: Copurification of DcuA-PhoA with AspA-Strep on a Strep-Tactin column.](https://example.com/image)

<table>
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<th>Protein</th>
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<th>Anti-Strep</th>
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**Function of DcuA/AspA, DcuB/FumB and DcuB/AspA metabolons?**

The experiments show specific interaction of the C4-DC transporters DcuA with AspA, and of DcuB with FumB and AspA (Fig. 4). Under anaerobic conditions fumarate represents the substrate for fumarate respiration. DcuB is a major carrier for C4-DC/succinate antiport in fumarate respiration (Six et al. 1994; Golby et al. 1998b; Janausch et al. 2002; Unden et al. 2002).
The major substrate for DcuB is L-malate, but due to the high levels of L-aspartate in the intestine of animals (Bertin et al. 2018; Schubert et al. 2021), L-Asp is also transported by DcuB (Six et al. 1994; Golby et al. 1998b). The interaction of DcuB with AspA and FumB suggests a coordination of the C4-DC uptake with the conversion to fumarate, and the excretion of the succinate after fumarate reduction, which indicates in the presence of DcuB/FumB and DcuB/AspA metabolons (Fig. 4A). A very similar situation applies to the role for a DcuA/AspA metabolon in fumarate respiration (Fig. 4A). Overall, DcuB/FumB, DcuB/AspA and DcuA/AspA metabolons are consistently indicated for fumarate respiration by interaction of the proteins, the colocalization of the corresponding genes in the genome and the physiological relevance of a coordination (Fig. 4A). The metabolons await, however, direct proof of metabolic coupling and biochemical backgrounds of the coupling.

![Diagram](https://example.com/diagram.png)

**Fig. 4:** Tentative scheme for the DcuA/AspA, DcuB/AspA and DcuB/FumB complexes and the coordinated reactions in fumarate respiration and nitrogen assimilation from L-Asp. Abbreviations as in Fig. 1.

When L-Asp functions as the nitrogen source for bacterial growth, it can be used as the sole source of nitrogen (Strecker et al. 2018). The ammonia released by AspA feeds the route for nitrogen assimilation via glutamine synthetase GS and glutamine 2-oxoglutarate amino transferase GOGAT (Strecker et al. 2018). The deaminating activity of AspA is further
stimulated by the regulator GlnB (or PII), which responds to the nitrogen status of the bacteria (Schubert et al. 2020). In consequence, from the L-Asp only the ammonia used under conditions of nitrogen assimilation, and fumarate with some L-malate is released by DcuA in near stoichiometric amounts to L-aspartate uptake (Strecker et al. 2018). Thus, coordination of the pathway with DcuA and AspA in the DcuA/AspA metabolon (Fig. 4B) is also highly indicated by genetic colocalization of the genes, protein interaction and the coordination of the L-Asp/Fum antiport to the ammonia supply.

Interaction of DctA with the regulator EIIAGlc of the phosphotransferase system PTS.

Bacteria producing the aerobic C4-DC transporter DctA and the protein EIIAGlc from the glucose phosphotransferase system were tested for interaction using the bacterial two-hybrid system BACTH. DctA represents the main transporter for the uptake of C4-DCs under aerobic condition, and EIIAGlc a component of the glucose-specific phosphotransferase system (PTS) of E. coli. EIIAGlc transfers the phosphoryl group to the glucose transporter EIIBC (Deutscher et al. 2006). EIIAGlc has in addition a regulatory role in glucose repression by transmitting the status for glucose availability to the AC CyaA and to some transporters for the uptake of carbon substrates. The N-and C-termini of DctA have cytoplasmic location (Witan et al. 2012), and fusions of DctA to the T18 and T25 fragments of B. pertussis AC were produced in all combinations, as well as to the N- and C-termini of EIIAGlc. DctA and EIIAGlc fusions showed high β-galactosidase activity at the same level as the positive control or higher than the background activity, indicating an interaction of DctA with EIIAGlc (Fig. 5). Although interaction of DctA by EIIAGlc is strong and evident, the role and physiological consequence is not clear. The interaction is suggested to inhibit DctA and C4-DC uptake in the presence of the preferred substrate glucose (i.e. by dephosphorylated EIIAGlc) and to exclude C4-DCs as alternative substrates from the cell. Inhibition of C4-DC uptake does not result in inducer exclusion, since C4-dicarboxylates are sensed in the periplasm by the two-component system DcuS-DcuR (Pappalardo et al. 2003; Unden et al. 2016). The supposed model suggesting ‘substrate exclusion’ is at variance with classical inducer exclusion, exemplified by the inhibition of lactose permease LacY or the maltose ABC transporter (Dills et al. 1982; Misko et al. 1987; Dean et al. 1990). Classical inducer exclusion inhibits uptake of substrates that serve also as inducers in the cytoplasm, thereby inhibiting the induction of alternative metabolic pathways (Deutscher et al. 2006). The interaction between DctA and EIIAGlc indicates a further level of
glucose regulation on DctA, in addition to the transcriptional regulation of dctA by cAMP-CRP (Davies et al. 1999).

\[ \text{Interaction [MU]} \]

\[ \text{T18 Zip} \quad \text{DctA}_{T25} \quad \text{T18 Zip} \quad \text{T25 Zip} \quad \text{EIIA}_Glc \quad \text{DctA}_{T18} \quad \text{EIIA}_Glc \quad \text{T18 Zip} \quad \text{T25 Zip} \quad \text{DctA}_{T18} \quad \text{DctA}_{T25} \]

**Fig. 5:** DctA-EIIA$^{Glc}$ interaction in vivo using the BACTH system. *E. coli* BTH101(ΔcyaA) was co-transformed pairwise with plasmids encoding fusions of T25 to DctA/EIIA$^{Glc}$ ($\text{T25}_{\text{DctA}}$) and fusions of T18 to DctA/EIIA$^{Glc}$ ($\text{T18}_{\text{DctA}}$). The combinations are shown on the x-axis. The leucine zipper pair T18Zip and T25Zip are applied as positive control (Karimova et al. 1998; Karimova et al. 2001), the pair T18Zip/T25DctA as the negative control for background β-galactosidase activity. The corresponding plasmids are derivatives of pUT18C ($\text{T18}_{\text{DctA}}$), pUT18 ($\text{DctA}_{\text{T18}}$), pKT25 ($\text{T25}_{\text{DctA}}$), and pKNT25 (DctA$\text{T25}$) (Tab. 1). The strains were grown aerobically in LB medium at 30°C to the exponential phase (OD$_{578}$ = 0.5). β-Galactosidase activities were quantified in Miller-Units (MU).

**Materials and Methods**

**Bacterial strains and growth conditions.** The *E. coli* K12 strains and the plasmids are listed in Table 1. All molecular methods, including cloning, DNA isolation, and manipulations were performed according to standard procedures (Jones and Gunsalus 1987; Sambrook et al. 1989; Miller 1992; Zientz et al. 1998a; Kleefeld et al. 2009a). The oligonucleotide primers are shown in Table 2. Bacteria were grown aerobically and anaerobically at 37°C in lysogeny broth (LB).

**β-Galactosidase assay.** Cells for the BACTH measurements were grown anaerobically in LB medium. Interactions in the BACTH system were measured in terms of the β-galactosidase activity (Miller 1992). BACTH experiments were conducted as described previously (Monzel et
al. 2013) with slight modifications (Wörner et al. 2016). Activities are the mean of at least two independent experiments and four replicates each.

**Membrane protein interaction (mSPINE).** The mSPINE assay for protein interactions was performed as described previously (Graf et al. 2014; Wörner et al. 2016). *E. coli* strain C43 was co-transformed with plasmids, pMW3049 and pMW577, which encoded the fusion proteins AspA-Strep and DcuA-PhoA, respectively. The bacteria were grown aerobically in 400 ml LB medium to an OD$_{578}$ of 0.5. Protein expression was induced for 3 h with 100 µM L-arabinose and 100 µM anhydrotetracycline (AHT). To crosslink the proteins in vivo, the bacteria were incubated with formaldehyde (0.6% w/v) for 15 min at 37°C with shaking. Next, the bacteria were harvested with centrifugation (11300 × g for 10 min), and the sediment was washed in Tris-HCl buffer (pH 8, 50 mM). For Strep-Tactin purification, cells were disrupted with a French Press (1260 psi) in a buffer that contained lauryldimethylamine oxide (LDAO, 0.05% (v/v)). The cell homogenates were clarified by centrifugation (39100 × g for 30 min at 4°C) and subjected to Strep-Tactin chromatography, 10% SDS-PAGE, and Western blotting. Western blots were probed with anti-Strep and anti-PhoA antisera (IBA Lifesciences, Sigma-Aldrich) (Scheu et al. 2010; Graf et al. 2014; Wörner et al. 2016).

**SDS-PAGE, Western Blot, and Immunostaining.** SDS-PAGE and Western blots were performed according to published procedures (Graf et al. 2014; Wörner et al. 2016). Immunostaining was performed with horse-radish peroxidase (HRP)-coupled anti-His-HRP, anti-Strep-HRP, and anti-IgG-mouse-HRP polyclonal antiserum (Sigma-Aldrich) or with anti-PhoA antibodies (produced in mouse, Sigma-Aldrich). For visualization chemiluminescent substrate HRP (Merck Millipore) was used and the blots exposed on X-ray film (Advansta).

**Acknowledgments.** We are grateful to Dr. A. Strecker (Mainz) for preparing the plasmids with the DcuASWT25 and DcuC$_{SWT25}$ constructs. Financial support from Deutsche Forschungsgemeinschaft (grant UN 49/19-1) is gratefully acknowledged.

**References**


Schubert, C., Zedler, S., Strecker, A. and Unden, G. 2020 L-Aspartate as a high-quality nitrogen source in Escherichia coli: Regulation of L-aspartase by the nitrogen regulatory system and interaction of L-aspartase with GlnB. *Molecular Microbiology*.


### Table 1. Strains of *E. coli* and plasmids used

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<th>Strain</th>
<th>Genotype</th>
<th>Reference</th>
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<th>Plasmid</th>
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pBAD18  | Expression vector; pBR322 ori, arabinose-inducible pBAD promoter, KanR  | (Guzman et al. 1995)
pMW577  | pBAD18 with *dcuA-phoA*, KanR  | (Bauer et al. 2011)

Table 2: Oligonucleotide primers. Primer for PCR amplification for AspA, FumB and EIIA. Restriction sites and amino acid exchange positions are printed in red.

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<td>KpnI</td>
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<td>BACTH_pKT25_EIIA_for</td>
<td>GCGCGCGGATCCGATGTTTTTGTGCTGATGAAC</td>
<td>BamHI</td>
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<tr>
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<td>GCTAGCCTACGGCGACTTCTTGATGC</td>
<td>KpnI</td>
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