Aspartate transport in *Rhizobium meliloti*

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Aspartate transport in *Rhizobium meliloti* was found to be mediated by at least two transport systems. High rates of aspartate uptake, necessary for growth on aspartate as a carbon source, required the dicarboxylate transport (Dct) system, which also transports succinate, fumarate and malate. The apparent $K_m$ for aspartate transport by this system was about 10 mM, compared to 15 $\mu$M for succinate. This difference in affinity was also apparent in competitive inhibition studies, which showed that succinate effectively inhibits aspartate transport. Although aspartate was not a preferred substrate, it was a very efficient inducer of the Dct system. Both the Dct system and a second aspartate transport system were capable of supplying aspartate for use as a nitrogen source. The second system had a lower apparent $K_m$ for aspartate transport (1.5 mM), and was competitively inhibited by glutamate. This aspartate-glutamate system was regulated independently from the Dct system, since it functioned in mutants lacking the Dct system regulatory genes dctB and dctD, and its induction did not coactivate the Dct system. Uptake kinetics in cultures growing on aspartate as nitrogen source showed rapid substrate exchange between extracellular and internal aspartate. *R. meliloti* was shown to be able to selectively activate the two uptake systems, and also regulated its metabolism as required to utilize aspartate as either carbon or nitrogen source.

Introduction

*Rhizobium meliloti* is a nitrogen-fixing bacterium which forms a symbiotic association with alfalfa (Verma & Long, 1983; Haaker, 1988). The bacteria elicit the formation of root nodules, in which they reside and provide an important source of nitrogen for the plant. In turn, the bacteria benefit from the association since they are provided with nutrients. They are induced to grow to high titres within the nodules, in which they differentiate to forms known as bacteroids contained within host plant cells.

The primary energy source for nitrogen fixation is photosynthesis. Organic compounds formed by this process, principally sucrose, are transported to the root and nodules where they are converted to substrates supplied to the bacteroids (McDermott et al., 1989; Streeter, 1991). Studies of C$_4$-dicarboxylate transport (dct) mutants have provided evidence that dicarboxylates are the major carbon and energy source for *R. meliloti* bacteroids in the nodule (Ronson *et al.*, 1981; Bolton *et al.*, 1986; Engelke *et al.*, 1987; Watson *et al.*, 1988). Mutants unable to transport dicarboxylates into the cell due to mutation of dctA, the structural transport gene, produced nodules which were symbiotically ineffective and contained an underdeveloped symbiotic zone (Watson *et al.*, 1988; Engelke *et al.*, 1989). Bacteroids were found to senesce prematurely within the host cells, such that a senescent zone predominated in the nodule. These studies demonstrated that C$_4$-dicarboxylates are essential for symbiotic nitrogen fixation, and are probably provided to the bacteroids within the host cells.

The Dct system is encoded by three genes, dctA, dctB and dctD, located on the exo megaplasmid. The dctB and dctD genes encode regulatory proteins which sense dicarboxylates outside the cell and respond by activating the dctA gene (Ronson *et al.*, 1987; Jiang *et al.*, 1989; Yarosh *et al.*, 1989; Watson, 1990). In *R. meliloti* the dctA gene encodes a highly hydrophobic protein which is thought to reside in the inner membrane and transport dicarboxylates into the cell (Jiang *et al.*, 1989; Engelke *et al.*, 1989; Watson, 1990). Substrates for the Dct system include the TCA cycle intermediates succinate, fumarate and malate, and aspartate. Succinate, fumarate or malate have been considered the most likely of these compounds...
to serve as carbon and energy substrates for bacteroids, since these have been reported to be taken up most readily by either free-living cells or purified bacteroids (Watson et al., 1988; McRae et al., 1989; Miller et al., 1988). Nevertheless, aspartate remains as a possible substrate supplied to R. meliloti within the nodule. Recently, the pathway for aspartate catabolism for use as a carbon and energy source was found to be deamination to oxaloacetate by an aspartate aminotransferase (Rastogi & Watson, 1991). A mutant of R. meliloti lacking this aminotransferase activity was symbiotically defective, implying that this pathway, or its reverse, is in some way necessary for the nitrogen fixation process. It has also been proposed that in bacteroids aspartate is exported during the operation of an aspartate-malate shuttle which would generate reductant, NADH, and thus provide energy for nitrogen fixation (Kahn et al., 1985; Appels & Haaker, 1991; Kouchi et al., 1991).

To study the metabolic interactions between the host plant and the bacteria, it is first necessary to have a basic understanding of the transport systems involved in the uptake of compounds which may be transferred between them. In this report we describe studies of aspartate transport in R. meliloti and define its relationship to the transport of succinate, fumarate and malate. We found that aspartate may be transported by at least two separate systems, depending upon culture conditions, and characterized the two systems.

Methods

Bacterial strains. R. meliloti JJ1c10 is a symbiotically effective strain used as the wild-type parent in these studies. It is a rifampicin-resistant derivative of an alfalfa nodule isolate, IZ450, initially obtained from the Institute Zimotecnio, Sao Paulo, Brazil. JJ1c10 also lacks a 200 kb plasmid band which is present in R. meliloti IZ450 (Adachi et al., 1983). The dct derivatives of this strain used in this work are shown in Table 1. Transductions were done using phage &M12 as described by Finan et al. (1984).

Table 1. Bacterial strains

<table>
<thead>
<tr>
<th>R. meliloti strain</th>
<th>Relevant characteristics</th>
<th>Source/reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>JJ1c10</td>
<td>Wild-type parent; Det' Fix' Rif'</td>
<td>Selvaraj et al. (1987)</td>
</tr>
<tr>
<td>4F6</td>
<td>dctA4 mutant derived from JJ1c10 by Tn5 mutagenesis; Det' Fix' Rif' Km'</td>
<td>Watson et al. (1988)</td>
</tr>
<tr>
<td>R647</td>
<td>JJ1c10 dctA::Tn5-35, Tn5 introduced by recombination, carries plasmid pH11J; Det' Fix' Rif' Km' Gmi'</td>
<td>Watson (1990)</td>
</tr>
<tr>
<td>R684</td>
<td>JJ1c10 dctA::Tn5-51, Tn5 transduced from R646; Det' Fix' Rif' Km'</td>
<td>Watson (1990); this study</td>
</tr>
<tr>
<td>R716</td>
<td>JJ1c10 dciB::Tn5-33, Tn5 transduced from R639; Det' Fix' Rif' Km'</td>
<td>Watson (1990); this study</td>
</tr>
<tr>
<td>R717</td>
<td>JJ1c10 dciD::Tn5-39, Tn5 transduced from R649; Det' Fix' Rif' Km'</td>
<td>Watson (1990); this study</td>
</tr>
</tbody>
</table>

Medium and culture conditions. Minimal medium used for the growth of R. meliloti was M9, containing (1-1): 7 g Na2HPO4, 3 g KH2PO4, 1 g NH4Cl and 1 g NaCl, with 10 ml each of sterile 0-1 mM MgSO4 and 0-1 mM CaCl2 added after autoclaving. The medium was supplemented with 20 mM carbon source and 5 ml of 1 % yeast extract (Difco) per litre. Yeast extract was found to be required for continued exponential growth of R. meliloti in this medium. For experiments with aspartate as nitrogen source, NH4Cl was omitted from the M9 and the medium supplemented with 5 mM-aspartate.

Cultures were grown from single colonies inoculated in a culture tube with 5 ml of medium, grown with shaking at 30 °C for 24-48 h. This culture was used to inoculate 50 ml of medium in a baffled culture flask which was shaken overnight to obtain an exponentially growing culture. Growth was monitored by measuring optical density at 620 nm (OD620). A second subculture was then made by appropriate dilution in 50 ml of fresh medium to obtain a mid-exponential-phase culture (OD620 = 0-5-0-8) early the next day.

Transport assays. To induce transport, cells were either cultured directly in medium containing the inducer, or washed and incubated for 1 h in medium containing inducer. To assay transport the cells were chilled, washed twice in M9, and resuspended in M9. An aliquot of cells was added to a vial shaking at 30 °C and warmed for 1 min. Labelled substrates were added, and 100 µl aliquots sampled at intervals. Uptake was measured using 50 or 100 µM-succinate containing 1 µCi ml-1 [2,3-14C]succinic acid (37 kBq ml-1) (New England Nuclear) or 5 mM-aspartate containing 2-5 µCi ml-1 [U-14C]Aspartic acid (92-5 kBq ml-1) (New England Nuclear). Samples were filtered through 0-45 µm nitrocellulose membrane filters (Schleicher & Schuell), washed twice with 5 ml M9, dried, and their radioactivity measured in a scintillation counter. Also, a 20 µl aliquot was spotted on a dry filter and counted to measure total radioactivity in the assay culture. Uptake rates were determined from plots of data using six sampling times, or at least two sampling times when a series of substrate concentrations was being assayed. Experiments have been repeated at least twice.

The loss of 14CO2 from cultures was determined as follows. A culture of R. meliloti grown in M9 medium with mannitol was washed, resuspended in M9 containing 5 mM-aspartate, and induced by incubation for 1 h. Assay of uptake was initiated by addition of [14C]succinate to an aliquot of cells to give a final concentration of 50 µM. Under these conditions, uptake was linear for 5 min, but the substrate was depleted within 5-10 min. Loss of 14CO2 was monitored by withdrawing 30 µl samples at 1 min intervals and depositing them directly onto membrane filters moistened with 10 % (w/v) trichloroacetic acid in scintillation vials. The samples were dried in a vacuum oven at 80 °C for 1 h, 10 ml of scintillation fluid was added, and the
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Fig. 1. Determination of apparent $K_m$ values for aspartate and succinate uptake. Lower plots show uptake rates ($V$) obtained with different substrate concentrations ($S$). Above each is a plot of $V'/S'$ used to calculate apparent $K_m$ values. (a) R. meliloti JJ1c10 succinate uptake by the Dct system. $\bullet$, Aspartate induced; $\blacktriangle$, succinate induced. (b) $\bullet$, R. meliloti JJ1c10 aspartate uptake by the Dct system. (c) Aspartate uptake by the aspartate-glutamate system. $\bullet$, R. meliloti JJ1c10; $\bigcirc$, dctB mutant R716.

Results

Aspartate and succinate transport by the dicarboxylate transport system

In R. meliloti, aspartate is taken up by the dicarboxylate transport system when it is provided as the sole source of carbon. A comparison of succinate and aspartate transport at different substrate concentrations is shown in Fig. 1(a and b). Although succinate at micromolar concentrations was transported readily, aspartate was taken up at a measurable rate only when its concentration was increased above 1 mM. The apparent $K_m$ for aspartate transport was 10 mM, compared to 15 $\mu$M for succinate.

The relatively high apparent $K_m$ for aspartate transport was consistent with comparisons of growth rates of R. meliloti on aspartate as carbon source when provided at different concentrations. A concentration of 4 mM aspartate or more was required to achieve the maximum growth rate, a doubling time of about 6 h. At concentrations below 4 mM, growth rate was approxi-
mately proportional to aspartate concentration. The doubling time in minimal medium with 1.0 mM-aspartate was about 24 h. Strains mutated in the dctA gene showed no growth on aspartate, as described previously (Watson et al., 1988). These data show that the Dct system is the only system in *R. meliloti* capable of aspartate transport for growth on this compound as a carbon source, and that below 4 mM growth is limited by the high apparent *K*ₘ for its transport.

Aspartate transport was rapidly inhibited (> 90%) by either 2 mM-dinitrophenol or 10 mM-sodium azide. The same result was obtained for succinate transport, as reported by others for *R. leguminosarum* and *R. meliloti* (Finan et al., 1981; Engelke et al., 1987). These results indicate that both aspartate and succinate transport by the Dct system are active transport processes.

Inhibition studies were done to compare the relative affinities of the Dct system for aspartate, succinate, fumarate and malate. As shown in Table 2, succinate transport was stoichiometrically inhibited by the presence of fumarate, malate or oxaloacetate. In contrast, aspartate produced little inhibition of succinate transport, even at concentrations 100-fold higher than that of succinate. Equivalent studies of the effect of dicarboxylic acids on aspartate transport showed that succinate, fumarate and malate all drastically inhibit its transport, even when added at one-tenth the concentration of succinate. These inhibition results are in agreement with the differences in apparent *K*ₘ values for aspartate compared to succinate.

Under our standard assay conditions, uptake by the Dct system is linear for 5 min, as expected if this represents the initial rates of transport by the system. However, Salminen & Streeter (1991) showed that uptake measurements using 14C-labelled substrates may underestimate the true rate due to evolution of 14CO₂ during metabolism. To provide an estimate of the magnitude of this effect in our experiments, we measured 14CO₂ production as the decrease of total radioactivity from *R. meliloti* cultures taking up 14C-succinate, as described in Methods. These experiments showed that the radioactivity retained in cells collected on membrane filters represented only 70% of the total uptake, taking into account the measured loss of 14CO₂. The evolution of 14CO₂ was detected within 1 min, and paralleled the uptake measured using filtration (data not shown).

Table 3 shows comparisons of the relative degrees of induction of the Dct system by different compounds. Only C₂-dicarboxylic acids induced the system to a high rate of uptake. There was a low but significant uptake by cells grown with other substrates or by cells starved for a carbon source. Glucose-grown cells characteristically showed the lowest rate of transport, but this was still significantly higher than uptake rates obtained using dct mutants [< 1 nmol min⁻¹ (mg protein⁻¹)]. Despite its high apparent *K*ₘ for transport, aspartate was a very efficient inducer, consistently causing initial succinate uptake rates 1.5–2 times higher than succinate-induced cells. Even at 1 mM, a concentration which does not support significant growth of *R. meliloti*, aspartate induced the Dct system to a higher degree than 20 mM-succinate.

### Table 2. Inhibition of succinate and aspartate uptake in *R. meliloti* by various carbon substrates

<table>
<thead>
<tr>
<th>Carbon substrate</th>
<th>Substrate concn</th>
<th>Succinate*</th>
<th>Aspartate†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>50 µM 500 µM 5000 µM 500 µM 5000 µM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glucose</td>
<td>ND 14 8 14 23</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Succinate</td>
<td>ND ND ND 89 92</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fumarate</td>
<td>57 90 99 88 91</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Malate</td>
<td>50 88 98 87 90</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aspartate</td>
<td>2 16 12 ND ND</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxaloacetate</td>
<td>34 85 ND ND ND</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glutamate</td>
<td>ND 3 6 5 13</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

ND, Not determined.

* Cultures were grown in M9 medium with 20 mM-succinate as carbon source. Assays were performed using 50 µM-14C-succinate with 50 µM, 500 µM or 5000 µM carbon substrate tested as inhibitor. The uptake rate in the absence of additional carbon substrate averaged 42 nmol min⁻¹ (mg protein⁻¹).

† Cultures were grown in M9 medium with 20 mM-aspartate as carbon source. Assays were performed using 5000 µM-14C-aspartate with 500 µM or 5000 µM carbon substrate tested as inhibitor. The uptake rate in the absence of inhibitors averaged 40 nmol min⁻¹ (mg protein⁻¹).

### Table 3. Induction of succinate uptake by different carbon sources in *R. meliloti*

<table>
<thead>
<tr>
<th>Carbon source</th>
<th>Succinate</th>
<th>Aspartate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mannitol</td>
<td>7</td>
<td>ND</td>
</tr>
<tr>
<td>Glucose</td>
<td>4</td>
<td>9</td>
</tr>
<tr>
<td>Succinate</td>
<td>45</td>
<td>18</td>
</tr>
<tr>
<td>Fumarate</td>
<td>33</td>
<td>23</td>
</tr>
<tr>
<td>Malate</td>
<td>68</td>
<td>34</td>
</tr>
<tr>
<td>Aspartate</td>
<td>92</td>
<td>40</td>
</tr>
<tr>
<td>Oxaloacetate</td>
<td>19</td>
<td>ND</td>
</tr>
<tr>
<td>Glutamate</td>
<td>10</td>
<td>ND</td>
</tr>
<tr>
<td>2-Oxoglutarate</td>
<td>8</td>
<td>ND</td>
</tr>
<tr>
<td>None</td>
<td>14</td>
<td>ND</td>
</tr>
</tbody>
</table>
Aspartate transport in R. meliloti

Aspartate transport as a nitrogen source

*R. meliloti* derivatives mutated in *dctA*, *dctB*, *dctD* or *ntrA* have been found to grow as well as the wild-type on minimal media in which aspartate is provided as the sole source of nitrogen, and containing a separate carbon source such as glucose or mannitol (Watson, 1990). These results imply the presence of a second aspartate transport system. To examine this second system and its relationship to the Dct system, we have studied the uptake of aspartate for use as a nitrogen source in Dct+ and Dct− *R. meliloti*.

Fig. 2 shows the growth and [14C]aspartate uptake of cultures of wild-type *R. meliloti* JJ1c10 and its *dctA* derivative 4F6 during 10 h growth in three minimal media with aspartate as sole nitrogen source. The highest uptake occurred when aspartate was supplied to the wild-type strain, JJ1c10, as both carbon and nitrogen sources. Aspartate is transported exclusively by the Dct system under these growth conditions. When succinate was also provided, the rate of [14C]aspartate incorporation in the JJ1c10 culture was less, about 30% of the rate attained by the wild-type culture growing with aspartate alone, demonstrating that succinate is transported for use as a carbon source in preference to aspartate. In this case a separate system for the transport of aspartate for use as a nitrogen source must be active, since succinate is a potent competitive inhibitor of aspartate transport by the Dct system. When mannitol was present as a possible carbon source, aspartate was incorporated at about the same rate as in the succinate plus aspartate culture. Apparently, mannitol was also preferred over aspartate as a carbon source, but aspartate uptake continued at a lower rate to supply nitrogen.

As expected, little growth or uptake of aspartate occurred in the 4F6 (*dctA*) culture when either aspartate or succinate were the only carbon sources. However, the culture grew well when mannitol was present as a carbon source and aspartate as sole nitrogen source, and [14C]aspartate was incorporated at about the same rate as in the wild-type.

These results demonstrated that *R. meliloti* is able to regulate the rate of aspartate utilization in response to the availability of carbon and energy substrates and its requirement for nitrogen. This regulation involves the activation of an aspartate transport system other than Dct as required to provide the cells with aspartate as a nitrogen source. Here, we will refer to this second system...
Aspartate uptake

medium without NH₄Cl with 20 mm-mannitol and 5 mm-aspartate. Aspartate uptake (a) was measured after addition of [¹⁴C]aspartate; succinate uptake ability (b) was measured after addition of [¹⁴C]succinate. ○, Wild-type R. meliloti; ○, dctA mutant 4F6; ▲, dctB mutant R716; Δ, dctD mutant R717.

as the aspartate-glutamate system, since, as described below, it also transports glutamate.

The aspartate-glutamate transport system

Fig. 3 shows assays of aspartate transport in cultures growing with mannitol as carbon source and aspartate as nitrogen source. Several dct mutants were used to discriminate between aspartate transport by the Dct system and that of the aspartate-glutamate system. In these experiments samples of each culture were also assayed for [¹⁴C]succinate uptake as an indication of Dct activity.

Uptake of [¹⁴C]aspartate as a nitrogen source did not show the linear kinetics characteristic of its uptake for use as a carbon source. The initial rates of uptake were over 10-fold greater than those characteristic of aspartate or succinate uptake by the Dct system for use as a carbon source. The rates then decreased to a constant, lower value. Since these cultures were in steady-state growth, and were not manipulated other than to add a trace of radioactive aspartate to the 5 mm-aspartate already present, the rapid, initial uptake of label must be due to equilibration of aspartate inside the cells with that outside by an exchange reaction. Therefore, net aspartate uptake by these cultures must be represented by the plateau rates rather than initial rates. This phenomenon also indicates the presence of internal aspartate pools which were largest in the wild-type cells.

Activity of the aspartate-glutamate system alone was demonstrated by [¹⁴C]aspartate uptake by the cultures of R716 (dctB) and R717 (dctD). These cultures were unable to take up succinate, nor was the aspartate uptake by these mutants inhibited by succinate. Instead, it was found that aspartate uptake was inhibited by glutamate, with aspartate and glutamate serving as equivalent substrates for the aspartate-glutamate system (Table 4).

Wild-type R. meliloti showed activity of both the Dct and aspartate-glutamate systems. The presence of an active Dct system was indicated by the ability of the culture to take up succinate (Fig. 3b). This demonstrated that Dct contributes to the uptake of aspartate for use as a nitrogen source, and is induced by aspartate under these conditions. The aspartate-glutamate system was also operating, since addition of succinate as an inhibitor during [¹⁴C]aspartate uptake resulted in only a 66% decrease in the initial aspartate uptake rate (Table 4). The uptake kinetics of the wild-type strain in the presence of succinate corresponded with those shown for aspartate-glutamate system transport by R716 and R717 in Fig. 3. Addition of glutamate resulted in a 40% decrease in [¹⁴C]aspartate uptake, an amount consistent with competitive inhibition of its aspartate-glutamate system component.

Unexpectedly, the dctA mutant 4F6 exhibited aspartate uptake kinetics intermediate between those of the wild-type and the dctB and dctD mutants. It was also found to take up succinate at about 20% the rate of the wild-type and this uptake was inhibited by malate (Table 4). This mutant carries a Tn5 insertion near the 5' end of the dctA gene, but these results showed that it still exhibited residual activity of the Dct system. We have found that this phenotype is not characteristic of all dctA mutants; analysis of two other independent dctA::Tn5 mutants, R647 and R684, showed that they exhibit as little succinate uptake as R716 (dctB) and R717 (dctD). Therefore, the Tn5 insertion in 4F6 is unusual in that a low level of transport function was still detectable. This level of transport appears sufficient to maintain an aspartate pool in the cell as large as that in the wild-type, but the initial [¹⁴C]aspartate equilibration by exchange was slower.

Determination of the $K_m$ for aspartate transport by the aspartate-glutamate system using wild-type R. meliloti and R716 (dctB) is shown in Fig. 1(c). The $K_m$ was found to be 1.5 mM. At low aspartate concentrations
Table 4. Inhibition of succinate and aspartate uptake in R. meliloti grown with aspartate as nitrogen source

Cultures were grown with 20 mM-mannitol and 5 mM-aspartate. Succinate uptake was measured by addition of 100 μM-[14C]succinate with or without 20 mM-malate as inhibitor. Uptake was sampled for 10 min as in the experiments shown in Fig. 2. Aspartate uptake was measured by addition of [14C]aspartate with or without 20 mM-succinate or glutamate as inhibitors. Uptake was sampled for 10 min as in the experiments shown in Fig. 2.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype</th>
<th>Succinate uptake [nmol min⁻¹ (mg protein)⁻¹]</th>
<th>Aspartate uptake [nmol min⁻¹ (mg protein)⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>(% inhibition in parentheses)</td>
<td>(% inhibition in parentheses)</td>
</tr>
<tr>
<td>JJ1c10</td>
<td>wt</td>
<td>88</td>
<td>5.3 (88)</td>
</tr>
<tr>
<td>4F6</td>
<td>dctA</td>
<td>15</td>
<td>1.0 (93)</td>
</tr>
<tr>
<td>R716</td>
<td>dctB</td>
<td>0.5</td>
<td>ND –</td>
</tr>
<tr>
<td>R717</td>
<td>dctD</td>
<td>0.2</td>
<td>ND –</td>
</tr>
</tbody>
</table>

(< 100 μM) there remained a low rate of uptake which was about equal in both strains, and which did not fit a straight-line double-reciprocal plot (data not shown). This indicates that there might be a third aspartate transport system which operates under these growth conditions, but it appears to be a high-affinity system which contributes little to aspartate uptake at concentrations above 1 mM compared to the other two systems.

Discussion

In this study, two aspartate transport systems have been characterized in R. meliloti. The Dct system is well known from other studies, but its role in aspartate transport has not previously been examined. An aspartate-glutamate transport system, active during growth on aspartate as a nitrogen source, has also been identified.

Metabolites identified as substrates for the Dct system are aspartate and the TCA cycle intermediates succinate, fumarate, malate and oxaloacetate. A large difference in the affinity of the Dct system was found for aspartate compared to succinate, fumarate and malate. The approximately 600-fold higher affinity makes succinate or the other TCA cycle intermediates highly inhibitory to aspartate transport, and cells exposed to a mixture will preferentially take up the other dicarboxylates before aspartate. Although the Dct system is known in other bacteria, a marked difference in substrate affinities has not been noted (Kay et al., 1987).

The Dct system was found to be inducible by its substrates. Despite the low affinity of the Dct system for aspartate as a substrate, it was a very effective inducer. The difference is consistent with a model in which transport and induction are mediated by separate components of the Dct system. In the current model for dct gene regulation, DctB protein is thought to be the sensor which detects dicarboxylates outside the cell membrane, and, through the DctD protein, activates the structural gene, dctA (Ronson et al., 1987b). A refinement of this model suggests that DctB protein does not sense dicarboxylates directly, but instead is proximal to DctA protein in the membrane, and senses dicarboxylates by monitoring activity of this protein (Ronson et al., 1987b; Yarosh et al., 1989). Our observations of induction of the system by aspartate do not favour this latter possibility, since we observed high levels of induction by aspartate concentrations which were severely limiting its transport as a substrate.

The Dct system is also induced when aspartate is present as a nitrogen source, and functions actively in the transport of aspartate for this purpose. The involvement of the Dct system in nitrogen utilization is consistent with the fact that the dctA gene is transcribed from a σ⁶⁴ type RNA polymerase promoter (Ronson et al., 1987b; Jiang et al., 1989; Watson, 1990). This characteristic is usually, but not universally, associated with nitrogen regulation (ntr) genes and genes involved in nitrogen metabolism (Magasanik, 1982; Ronson et al., 1987a).

The aspartate–glutamate system transports aspartate for use as a nitrogen source and recognizes aspartate and glutamate as substrates. It appears to be regulated, possibly partly in response to a nitrogen requirement, but is not directly induced by aspartate. It has a relatively low apparent $K_m$ for aspartate, about 1.5 mM compared to 10 mM for the Dct system. The difference in apparent
$K_m$ values for the two systems is such that their relative contributions to aspartate transport vary with extracellular aspartate concentration. At concentrations below about 5 mM the proportion of aspartate transported by the Dct system decreased in favour of transport by the aspartate–glutamate system. In addition to functioning as a higher-affinity system for aspartate transport for use as a nitrogen source, this second system is also necessary for aspartate transport when a dicarboxylate preferred by the Dct system, such as succinate, is present as a carbon source.

*R. meliloti* is able to regulate the rate of aspartate utilization in response to availability of carbon and energy substrates and its requirement for nitrogen. This regulation involves the activation of the aspartate–glutamate transport system in addition to the Dct system. However, the utilization of aspartate as a nitrogen source appears to be primarily regulated at the level of its metabolism. For example, when 5 mM-aspartate must serve as both carbon and nitrogen source, the Dct system alone is able to supply about 40 nmol aspartate min$^{-1}$ (mg protein)$^{-1}$. At the same aspartate concentration, but with mannitol also present, the cells choose mannitol as main source of carbon instead of aspartate. The rate of aspartate uptake is then only about 10 nmol min$^{-1}$ (mg protein)$^{-1}$, even though the Dct system is fully induced, as measured by its capacity for succinate uptake, and the aspartate–glutamate system is also operating. The lower rate of aspartate metabolism in the presence of these two active uptake systems apparently results in a marked increase in size of internal aspartate pools, as evidenced by the initial rapid exchange of exogenous and endogenous aspartate in uptake assays.

Streeter & Salminen (1990) have found that in *Bradyrhizobium japonicum* bacteroids a periplasmic aspartate aminotransferase activity, released from cells by dilute detergent, is present which can mediate transfer of nitrogen from aspartate to 2-oxoglutarate. They suggested that in our dctA mutant, *R. meliloti* 4F6, the ability to utilize aspartate as a nitrogen source could be accounted for by a similar transamination and subsequent uptake of the glutamate into the cytoplasm. In support of this idea, they subsequently demonstrated the presence of aspartate aminotransferase activity which could be released from this mutant by mild detergent treatment. In the case of the studies done here, it is clear that this mechanism of nitrogen transfer is not the major route for aspartate nitrogen uptake, since aspartate utilization was accompanied by proportional incorporation of the $^{14}$C-radiolabel.

Interestingly, the dctB and dctD genes are not required for dctA function in the nodule, since mutants in these genes are nearly as effective symbiotically as wild-type *R. meliloti* (Watson et al., 1988; Engelke et al., 1989; Yarosh et al., 1989). This suggests that an activator other than DctD protein turns on dctA in the nodule. Since both the Dct and aspartate–glutamate systems are activated when aspartate is present as a nitrogen source, our experiments tested the possibility that an aspartate–glutamate system activator may also turn on the dctA gene. However, since no Dct function was detected in cultures of dctB or dctD mutants growing with aspartate as nitrogen source, cross-talk between the two systems cannot account for dctA gene activation in the nodule.

The Dct system is required for symbiotic nitrogen fixation at the stage of the symbiosis in which the bacteria have proliferated within the host cells. It is unlikely that the Dct system requirement solely involves transport of aspartate for use as a nitrogen source, since the aspartate–glutamate system is also able to provide this function. As the aspartate–glutamate system was found not to operate in free-living cells when aspartate is both carbon and nitrogen source, it remains possible that aspartate serves both functions in the nodule.

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References


Aspartate transport in R. meliloti


