Generation of *Azotobacter vinelandii* strains defective in siderophore production and characterization of a strain unable to produce known siderophores

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Siderophore-negative mutants of *Azotobacter vinelandii* were generated by insertional mutagenesis with a Tn5 construct containing a promoterless *luxAB* fusion. The use of this construct, delivered on a suicide plasmid by conjugation, allowed the selection of mutations in iron-repressible genes by virtue of the expression of iron-regulated bioluminescence. Although many iron-regulated mutants were selected, only a few could be easily identified as defective in siderophore production. These included a non-fluorescent azotobactin-negative phenotype (strain D27), and strain F196, which had lost the ability to produce the catechol siderophores azotochelin and aminochelin as well as the lower-affinity chelator 2,3-dihydroxybenzoic acid. Strain D27 had normal production of catechol siderophores, while strain F196 produced 2.5 times as much azotobactin as the wild-type. Two other mutants demonstrated normal catechol levels and either low or relatively unrepressed azotobactin levels. Transformation of the DNA from strain F196 into another spontaneously obtained azotobactin-negative strain (UA1) resulted in strain P100, which was unable to produce the known siderophores. Unlike the wild-type and other siderophore-deficient mutants, this strain was unable to grow in the presence of the iron chelator ethylenediamine di-(o-hydroxyphenylacetic acid) (EDDHA; 50 µg ml⁻¹) unless stored iron was carried over in the inoculum. Strain P100 did grow on iron-limited medium containing EDDHA when the catechol or azotobactin siderophores were provided exogenously. However, strain P100 gave a positive result in the chrome azurol-S assay (CAS), a non-specific assay for siderophores. The CAS activity was iron-repressible and strain P100 was able to grow and accumulate more iron than the insoluble iron minerals Fe₅, vivianite and Fe₃O₄, than was available by simple diffusion or exchange. Therefore, it appears that iron-limited *A. vinelandii* produces an as yet unidentified low-affinity non-conventional (non-catechol, non-hydroxamate) siderophore.

Introduction

Almost all organisms require iron for survival. However, under aerobic, neutral pH growth conditions soluble iron is rapidly oxidized and forms extremely insoluble oxides and oxyhydroxides (Neilands, 1981). Most aerobic and facultative organisms must scavenge trace amounts of soluble iron and have to transport it into the cell by the production of small, soluble, high-affinity Fe(III)-coordinating ligands, the siderophores (Neilands, 1981). The production of siderophores and the specific transport of iron into the ligand-producing cell is an important adaptation for survival and success in a variety of iron-limited environments (Cox, 1989; Neilands, 1982).

*Azotobacter vinelandii* is an obligate aerobe that is capable of aerobic nitrogen fixation. The organism uses its extremely high respiratory rate to consume oxygen and protect the oxygen-labile nitrogenase (Robson & Postgate, 1980). Such a high respiratory activity generates toxic oxygen radicals that are destroyed by active catalase and superoxide dismutase (Jurtshuk et al., 1984). All of these activities are dependent on an adequate supply of iron. Upon growth in iron-limited medium, *A. vinelandii* produces a pyoverdin-type siderophore, azotobactin (Demange et al., 1986; Page et al., 1991) and two catechol-type siderophores, azotochelin (Corbin & Bulen, 1969) and aminochelin (Page & von Tigerstrom, 1988). Furthermore, 2,3-dihydroxybenzoic acid (2,3-DHBA) appears to be produced as a low-affinity iron

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Abbreviations: CAS, chrome azurol-S; 2,3-DHBA, 2,3-dihydroxybenzoic acid; EDDHA, ethylenediamine di-(o-hydroxyphenylacetic acid); IROMP, iron-regulated outer-membrane protein.
chelator that solubilizes iron and facilitates low-affinity constitutive iron uptake (Page & Huyer, 1984).

The production of siderophores by *A. vinelandii* follows a sequential induction pattern. As iron becomes limiting (≈7 μM-Fe²⁺) the catechol siderophores are produced coordinately (Page & von Tigerstrom, 1988). If iron continues to be limiting (≤3 μM-Fe²⁺), then azotobactin is formed (Page & von Tigerstrom, 1988). When iron is supplied to the cells, azotobactin is repressed first, followed by the catechols, until 2,3-DHBA remains as the sole ligand (Page & Huyer, 1984). In enteric bacteria, iron repression is mediated by the association of a repressor protein (Fur) with intracellular Fe²⁺ and specific promoter sequences (de Lorenzo et al., 1987). The pattern of sequential siderophore regulation is not seen in enteric bacteria (McIntosh & Earhardt, 1977), which suggests that in *A. vinelandii* iron-repressible genes may be regulated by a different mechanism or that the interaction of the Fur–Fe²⁺ complex with iron-repressible promoters may be affected by promoter affinity or other factors (Page & Patrick, 1988).

A first step in studies of iron regulation in *A. vinelandii* is the generation of mutants defective in iron-regulated gene activity. Transposon mutagenesis, using a derivative of Tn5 containing a promoterless *luxAB* fusion, has been used in this study to generate *A. vinelandii* strains defective in siderophore production. Tn5 was delivered on a suicide vector (pTnSlux), so that Tn5 would only persist in cells where transposition into the chromosome had occurred and bioluminescence (*Lux*) activity would persist in cells where transposition into the chromosome was defective in siderophore production. Tn5 was delivered with 50 pg ml⁻¹ to further limit iron availability. Growth of *A. vinelandii* strains under aerobic nitrogen-fixing conditions was carried out in 500 ml Erlenmeyer flasks (each flask contained four radial indentations in its base) containing 100 ml of medium. The flasks were incubated with shaking at 300 r.p.m. The inoculum was prepared in 0FeBBGN, to minimize the carryover of stored iron (see Results).

*Escherichia coli* strain S17-1 (prov*Res*; Mod*) containing the suicide plasmid pTn5luxAB (a gift of A. Szalay, Department of Plant Science, Faculty of Agriculture, University of Alberta; Berg et al., 1989), and E. coli strain HB101 (supE44 hsdS20 (r·m·) recA13 ara-14 proA2 lacY1 galK2 rpsL20 xyl-5 mtl-l) containing the helper plasmid PRR2013 (Figurski & Helsinki, 1979) were grown at 37 °C in TYE medium (tryptone, 15 g l⁻¹; yeast extract, 10 g l⁻¹; NaCl, 5 g l⁻¹; agar, 18 g l⁻¹) with 50 μg kanamycin ml⁻¹ for strain HB101 and 40 μg ampicillin ml⁻¹ for strain S17-1.

Mineral iron enclosed in dialysis bags and determination of iron concentration. In order to study the ability of *A. vinelandii* strains to solubilize iron from insoluble minerals and to have a mineral-free cell suspension for further analysis, crushed mineral (50 mg, about 200 mesh) was first added to a 1 cm diameter, 10–14 kDa molecular mass cut-off dialysis tubing sac containing 2 ml 0FeBBGN, then the sealed bag was placed in 100 ml 0FeBBGN. Each mineral was prepared in duplicate flasks and sterilized by autoclaving. One flask was prepared 24 h in advance, incubated with shaking overnight, then the dialysis bag was aseptically removed. This flask served as a control for freely exchangeable iron (Page, 1987). The second flask, containing sterile medium and the dialysis bag, and the freely exchangeable iron control flask were inoculated with an iron-limited medium, pregrown overnight in 0FeBBGN. After 24 h incubation at 30 °C with shaking at 225 r.p.m., 75 ml cell suspension was harvested by centrifugation and washed with 10 mM-sodium citrate (pH 7.2) to remove surface-bound iron. The cell pellet was suspended in 2.5 ml 7% (w/v) perchloric acid and extracted overnight at room temperature and for 4 h at 80 °C prior to iron determination using a 2,3-bipyrilid acid assay (Page & Huyer, 1984). All glassware for these studies was acid washed (Collinson et al., 1987). Cell protein was determined as described by Page & Huyer (1984).

Introduction of pTn5lux by conjugation and selection of transconjugants. Exponentially growing *A. vinelandii* and *E. coli* cells were mixed in approximately equal proportions after washing three times with iron-limited Burk’s buffer (0FeBB). Matings were carried out directly on the surface of BBGN plates at 30 °C without collecting the cells on a filter. At 24 h intervals the mixed culture was scraped from the plate, resuspended in 1–3 ml 0FeBB, and spread onto BBGN plates containing 10 μg kanamycin ml⁻¹ for the selection of transconjugants. Isolated Kan* Azobacter* colonies were purified by streaking on BBGN plates containing kanamycin.

Detection of iron-repressible bioluminescence. Bioluminescence was detected by Petri plate contact printing using Kodak XAR-5 X-ray film. In comparisons of the bioluminescence of cells grown on iron-limited (Burk’s medium containing 1 μM-ferric citrate) and iron-sufficient (Burk’s medium containing 300 μM-ferric citrate) medium on a single Petri plate, the bacteria were spotted (about 5–7 mm diameter) in duplicate onto the surface of the respective agar media contained in a Petri dish split in half by a plastic divider. One drop (10 μl) of n-decanal was spread on the lid of the Petri dish and the complete dish was placed agar-side-down on the film for 15 min to allow the bioluminescent spots to expose the film.

Detection of siderophores. Siderophores were detected in acidified (HCl, pH 1–8) culture supernatants by measuring absorption at 310 nm for catechols and 380 nm for azotobactin (Page & Huyer, 1984).
using a Hitachi U-2000 recording spectrophotometer. Azotobactin contains a hydroxamate group contributed by N6-hydroxyornithine (Demange et al., 1986), which was quantified by the method of Csaky (1948), with the hydrolysis step conducted at 100°C for 6 h, as recommended, or for 1 h in the autoclave (121°C, 1-1 kg cm⁻² pressure). Catechols were quantified by the methods of Barnum (1977). Citric acid was assayed enzymically with a kit obtained from Boehringer. Samples were concentrated 10-fold by freeze-drying prior to the citrate assay.

Siderophore production also was detected using the chrome azurol-S (CAS) plating and CAS-shuttle colorimetric assays described by Schwyn & Neilands (1987), except that Burk's medium was substituted for the MM9 salts/casamino acids medium. The CAS assay is a universal assay for the detection of siderophores and operates independently of the siderophore structure. The presence of an iron-chelator (presumably a siderophore) is indicated by the decolorization of a blue-coloured ferric-CAS complex, resulting in a yellow-gold halo around colonies growing on CAS medium or as a result of mixing culture fluids in a tube assay. The shuttle assay contains 4 mm-sulphosalicylic acid added to the reaction mixture to destabilize the ferric-CAS complex and accelerate ferric exchange to the siderophore (Schwyn & Neilands, 1987). The CAS shuttle assay was initiated by the addition of 0.5 ml culture supernatant fluid to 0.5 ml of the CAS shuttle solution. This was immediately mixed and placed in a Hitachi U-2000 spectrophotometer and the decrease in absorption at 630 nm was recorded over ≥ 2 min. The end-point of the reaction was the time at which no further decolorization of the CAS complex was observed. Total siderophore activity (TSA) was calculated from the initial rate of decolorization, where one unit of activity was a decrease in A₆₃₀ of 0.001 s⁻¹. The total siderophore specific activity was calculated as the TSA units (mg cell protein)⁻¹ present in 0.5 ml culture.

Catechol siderophores in the acidified culture supernatant fluid were extracted into ethyl acetate and visualized by thin-layer chromatography on silica gel G (Brinkmann), using a benzene/acetic acid/water (15:2:3 by vol.) solvent system and detection with 2,2'-dipyridyl-ferric chloride (Kreb's et al., 1969). Aminochelin was demonstrated as residual material absorbing at 310 nm that could be extracted with butanol (Page & von Tigerstrom, 1988).

When the siderophore activity of ethyl-acetate-extracted culture fluids was tested by the CAS-shuttle assay, the ethyl acetate fraction was air dried and dissolved in distilled water and the ethyl acetate in the aqueous phase was allowed to evaporate overnight. All fluids were adjusted to pH 6-0 before the CAS-shuttle assay. Ethyl acetate and ethanol were found to give false positive results in this assay, but 10-fold concentrated iron-limited medium did not produce interference.

Demonstration of iron-repressible outer membrane proteins (IRMPs). A. vinelandii was grown overnight in iron-limited (1 μM-ferric citrate) and iron-sufficient (50 μM-ferric citrate) Burk's medium. The cells were harvested by centrifugation and broken by sonication as described by Page & Huyer (1984). Outer membranes were prepared by extraction with Sarcoyl (Filip et al., 1973; Page & Huyer, 1984) and examined by SDS-polyacrylamide gel electrophoresis as described previously (Page & von Tigerstrom, 1982). The outer membrane proteins in the gel (20 μg per lane) were visualized with the silver stain of Wray et al. (1981).

Transformation of A. vinelandii. Chromosomal DNA was purified as described by Wilson (1990). Transformation and competence development conditions for A. vinelandii were as described by Page & von Tigerstrom (1979), except that the incubation time of 20 h for enrichment of the phenotype was not necessary, as previously noted (Doran et al., 1987).

**Results**

pTn5luxAB mutagenesis of A. vinelandii

The plasmid pTn5luxAB does not encode transfer functions and would normally be mobilized by a helper plasmid in heterologous matings. However, transconjugants were not obtained in short-term matings (up to 24 h) using traditional procedures (Selveraj & Iyer, 1983). A few transconjugants were observed after 2-3 d of mating when triparental matings were conducted directly on the surface of the BBGN plates. However, the frequency of plasmid transfer was increased by mixing only the donor E. coli with the recipient A. vinelandii, without the helper plasmid. Clearly pTn5luxAB was not being transferred by normal conjugation. The frequency of plasmid transfer could not be accurately determined, because many of the Kan' A. vinelandii isolates were undoubtedly identical clones which arose during the long incubation period.

Over 500 Kan' strains were generated by repeated transfers of pTn5luxAB to A. vinelandii UW. Because the plasmid was not maintained in A. vinelandii, stable Kan' strains were assumed to be the result of Tn5 transposition into the chromosome. Screening the strains for iron-repressible bioluminescence categorized the strains into four groups: (i) those that were not bioluminescent (very frequent); (ii) those that expressed constitutive strong bioluminescence (rare); (iii) those that expressed moderately repressible bioluminescence (very frequent); and (iv) those that expressed strongly regulated bioluminescence (frequent). All the Kan' strains were stable and did not lose antibiotic resistance or Lux activity, even after transfer under nonselective conditions.

The Kan' strains expressing iron-regulated bioluminescence (about 100 strains including some identical clones) were immediately tested on CAS medium plates in an attempt to detect siderophore-negative strains, siderophore overproduction, or siderophore transport deficiencies as described by Schwyn & Neilands (1987). In all cases, the haloes around the spots of bacterial growth (indicating deferration of the ferric complex in the medium) were normal (within the limits of experimental error).

However, since A. vinelandii normally produced multiple siderophores, CAS screening was likely to be too imprecise to detect a deficiency in a single siderophore. Therefore, the culture supernatant fluids of cultures grown overnight in iron-limited (0 and 1 μM-ferric citrate) and iron-sufficient (10 and 50 μM-ferric citrate) medium were scanned spectrophotometrically to detect catechols (A₄₁₀) and azotobactin (A₃₈₀). The vast majority of the strains demonstrated absorption per cell protein values identical to the wild-type (within the
Table 1. Characterization of siderophore-negative strains

All strains were grown for 24 h at 30 °C in iron-limited (1 μM-ferric citrate) medium. All values are means of at least two duplicates.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Protein (μg ml⁻¹)</th>
<th>Iron-repressible Lux activity</th>
<th>Hydroxamate</th>
<th>Catechol</th>
<th>Cellular iron [ng (mg protein)⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>UW</td>
<td>389</td>
<td>NA</td>
<td>A₃₈₀*</td>
<td>A₃₁₀*</td>
<td>3.24</td>
</tr>
<tr>
<td>D27</td>
<td>310</td>
<td>No</td>
<td>0</td>
<td>0</td>
<td>2.75</td>
</tr>
<tr>
<td>UA1</td>
<td>402</td>
<td>NA</td>
<td>0</td>
<td>0</td>
<td>3.31</td>
</tr>
<tr>
<td>F196</td>
<td>348</td>
<td>Yes</td>
<td>6.96</td>
<td>4.06</td>
<td>0</td>
</tr>
<tr>
<td>P100</td>
<td>229</td>
<td>Yes</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

NA, Not applicable.

* Absorption of acidified supernatant fluids per mg cell protein present per ml culture.
† Hydrolysis was conducted for 1 h in the autoclave or for 6 h at 100 °C (values in parentheses). Units are μg hydroxylamine equivalents ml⁻¹.

limits of experimental error). However, three strains (F196, F174, F124; probably identical clones) did not form any catechols and two other strains demonstrated normal catechol formation, but low (E21) or relatively unpressed (F130) azotobactin formation. In addition, a strain that was unable to form azotobactin (D27) was detected amongst the non-bioluminescent strains by its failure to show fluorescence under 366nm UV illumination. No differences were detected in the production of IROMPs by the wild-type strain and these mutant strains.

Strain F196 was grown in iron-limited medium containing 500 μM-2,3-DHBA and the resulting culture supernatant fluid was acidified and extracted with ethyl acetate. All of the catechol was extracted into ethyl acetate and all of this was determined to be 2,3-DHBA by thin-layer chromatography. Therefore, it appears that the failure of strain F196 to form the catechol siderophores cannot be reversed by the addition of 2,3-DHBA as a precursor.

Generation of the pyoverdin-negative, catechol-negative strain P100

A siderophore-negative strain of A. vinelandii could be a very useful tool for the study of siderophore-mediated iron-uptake, for use in a bioassay to detect mutant siderophores and to examine the importance of iron accumulation on cell activities. Such a strain was constructed by transfer of chromosomal DNA from strain F196 to the spontaneously generated azotobactin-negative strain UA1 by transformation and the selection of Kan' colonies. Almost all of the transformants were phenotypically identical (failure to show fluorescence under UV light, whitish colony colour, small size). One transformant was picked and designated strain P100. This strain showed no trace of azotobactin or catechols in spectrophotometric scanning.

Siderophore production by siderophore-negative strains

The production of catechols and hydroxamates by the strains was determined quantitatively (Table 1). The azotobactin-negative strains, D27 and UA1, did not produce hydroxamate and produced normal amounts of catechol. On the other hand, the catechol-negative strain, F196 (and its identical clones, data not shown), produced 2.5 times as much of the hydroxamate azotobactin as the parent strain. The low level of catechol detected in the normal or 10-fold concentrated culture fluids of strains F196 and P100 could not be extracted with ethyl acetate, was not detectable by thin-layer chromatography, and could not be extracted with butanol. This amount of catechol corresponded to an absorbance value of 0.004, essentially at the limit of detection and within background values. Strains that lacked a single type of siderophore were not greatly impaired in their ability to grow, and accumulated iron to an extent comparable to the wild-type strain (Table 1). Strain P100 grew to about 59% of the wild-type value, but accumulated about 94% of the iron per cell protein found in the wild-type strain. Thus strain P100 appeared capable of iron accumulation from iron-limited medium, but this was less efficient than that of the other strains, hence overall growth was limited.

Inhibition of growth of strain P100 with EDDHA

Strains UW, D27, F196 and UA1 grew equally well in the iron-sufficient BBGN medium alone or containing
A. vinelandii defective in siderophore production

Fig. 1. Growth of the A. vinelandii wild-type strain UW (a) and strain P100 (b, c) in iron-sufficient BBGN alone (○) or containing 50 μg EDDHA ml⁻¹ (■). (a, b) Inoculant cells pregrown in BBGN medium; (c) inoculant cells pregrown in 0FeBBGN.

The inhibitory effect of EDDHA on the growth of strain P100 was also demonstrated on EDDHA-containing plates (Fig. 2). However, on solid medium the growth of strain P100 was only inhibited by EDDHA when the plating medium was 0FeBBGN. When the plating medium contained 25 μM-ferric citrate, strain P100 was able to grow even when the inoculum had been pregrown under iron-limited conditions. Strain P100 could be used in a bioassay for Azotobacter siderophores, as demonstrated by cross-feeding with the other strains on 0FeBBGN medium containing EDDHA (Fig. 2). The inhibition of the growth of strain P100 caused by EDDHA was completely reversed by azotobactin, by the catechol siderophores secreted by the mutants and by the mixture of siderophores secreted by the wild-type strain, but was not reversed by products excreted by strain P100.

Fig. 2. Reversal of EDDHA inhibition of the growth of strain P100 by siderophores from A. vinelandii strains. A plate of 0FeBBGN containing 50 μg EDDHA ml⁻¹ was spotted with 2 μl of iron-limited culture of strain P100 (upper spot in each pair) and the siderophore-producing strains or P100 as indicated. The plate was incubated for 8 d at 30 °C.

The inhibitory effect of EDDHA on the growth of strain P100 was also demonstrated on EDDHA-containing plates (Fig. 2). However, on solid medium the growth of strain P100 was only inhibited by EDDHA when the plating medium was 0FeBBGN. When the plating medium contained 25 μM-ferric citrate, strain P100 was able to grow even when the inoculum had been pregrown under iron-limited conditions. Strain P100 could be used in a bioassay for Azotobacter siderophores, as demonstrated by cross-feeding with the other strains on 0FeBBGN medium containing EDDHA (Fig. 2). The inhibition of the growth of strain P100 caused by EDDHA was completely reversed by azotobactin, by the catechol siderophores secreted by the mutants and by the mixture of siderophores secreted by the wild-type strain, but was not reversed by products excreted by strain P100.

Reaction of strain P100 in the CAS assay

When strain P100 was plated on the CAS medium, a zone of clearing was clearly evident after 24 h incubation (Fig. 3). This result was unexpected because all the other data had demonstrated that strain P100 was apparently devoid of known siderophore production. The CAS result with strain P100 was possibly due to a non-

the strong chelator of iron EDDHA (50 μg ml⁻¹). In the presence of EDDHA, the growth rate of strain P100 was decreased (Fig. 1b). The growth of strain P100 was dependent on the iron content of the medium used to grow the inoculum; when it was pregrown in iron-limited medium (0–1 μM-ferric citrate) there was essentially no subsequent growth in the medium containing EDDHA (Fig. 1c). The growth of the other strains in this medium was not adversely affected by prior iron starvation.
Fig. 3. Siderophore production by wild-type and siderophore-deficient *A. vinelandii* strains on CAS medium. The plate was spotted with 2 µl of inoculum pregrown in BBGN and incubated for 1 d at 30°C.

Table 2. Total siderophore activity of strains as estimated by the CAS-shuttle assay

<table>
<thead>
<tr>
<th>Strain</th>
<th>Total siderophore activity</th>
<th>Specific activity</th>
<th>End-point (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>UW</td>
<td>23</td>
<td>118</td>
<td>1-2</td>
</tr>
<tr>
<td>D27</td>
<td>22</td>
<td>142</td>
<td>1-2</td>
</tr>
<tr>
<td>UA1</td>
<td>21</td>
<td>104</td>
<td>1-2</td>
</tr>
<tr>
<td>F196</td>
<td>13</td>
<td>75</td>
<td>5-6</td>
</tr>
<tr>
<td>P100</td>
<td>7-1</td>
<td>61</td>
<td>≥15</td>
</tr>
<tr>
<td>EDDHA</td>
<td>9-1</td>
<td>NA</td>
<td>10-15</td>
</tr>
</tbody>
</table>

NA, Not applicable.
* Total siderophore activity, where one unit is a ΔA₆₃₀ of 0.001 s⁻¹.
† Total siderophore specific activity, as units (mg cell protein)⁻¹ present in 0.5 ml of culture.
‡ EDDHA concentration was 50 µg ml⁻¹ in unincubated medium.

The plating results were confirmed using the spectrophotometric CAS-shuttle assay. The ability of iron-limited culture supernatant fluids to decolorize the ferric-CAS complex was followed by measuring the ΔA₆₃₀ at 10 s intervals until no further decolorization occurred (Table 2). These data showed that strains UW, D27 and UA1 had about the same total siderophore activity in this assay. The catechol-negative strain F196 had about 60% of the wild-type activity and strain P100 had about 30% of the wild-type activity (Table 2). The catechol-positive strains UW, D27 and UA1 recovered ferric iron from the CAS complex very rapidly, whereas the azotobactin-positive strain F196 recovered iron relatively slowly. Strain P100 was the slowest. The deferrating capacity of the putative siderophore in the culture fluid of strain P100 was similar to, but less than that of 50 µg EDDHA ml⁻¹ (Table 2); hence EDDHA at this concentration would inhibit the growth of strain P100 (Figs 1 and 2).

The culture fluid of strain P100 was examined for the presence of citric acid, because this has been reported to act as a simple siderophore in *Azotobacter salinus* (Page, 1987; Page & Shivprasad, 1991). However, no detectable citric acid was found in 10-fold concentrated culture fluid of iron-limited strain P100.

Function of the putative siderophore of strain P100

Strain P100 was incubated in the presence of insoluble iron minerals to determine if the putative siderophore would promote iron uptake. The mineral was contained inside a dialysis bag to eliminate the possibility of mineral adsorption to the cell surface. By definition (Neilands, 1981) a siderophore will be small enough to exchange through the dialysis bag, mobilize the insoluble iron and make it available to the cell. Strain P100 was unable to grow better in the presence of 50 mg ml⁻¹ olivine ([Mg,Fe₄SiO₄]), haematite (Fe₂O₃), siderite (FeCO₃) or goethite (FeO(OH)) than in OFeBBGN alone, whereas growth of strain UW was enhanced in the presence of these insoluble iron sources (data not shown; Page & Huyer, 1984). However, growth of strain P100 was promoted in the presence of the minerals FeS, vivianite [Fe₃(PO₄)₂·8H₂O], and to a lesser extent Fe₃O₄ (Table 3). This growth promotion was paralleled by an increase in cellular iron which was greater than that obtained from the mineral by simple diffusion. Strain P100 was notably less efficient than strain UW in removing iron from these minerals.

Production of the putative siderophore by strain P100 was repressed by iron. The total siderophore activity of strain P100 in the CAS-shuttle assay was decreased from 59 units (mg protein)⁻¹ to 2.8 units (mg protein)⁻¹ when the iron content of the medium was increased from 1 µM to 25 µM. Under the same conditions, the total siderophore activity of strain UW decreased from 118 units (mg protein)⁻¹ to 3.5 units (mg protein)⁻¹. Addition of ferric iron to an iron-limited culture supernatant fluid of strain P100 did not result in the formation of a coloured ferric complex or any detectable spectral shift.
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Table 3. Solubilization of iron from iron-containing minerals included in dialysis bags

The data are means ± standard deviation of two or more duplicates.

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Control* With mineral</th>
<th>Control* With mineral</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>9.6 ± 0.3</td>
<td>11.6 ± 1.9</td>
</tr>
<tr>
<td>Fe₃O₄</td>
<td>10.0 ± 0.2</td>
<td>113 ± 7.4</td>
</tr>
<tr>
<td>Vivianite†</td>
<td>16.4 ± 0.8</td>
<td>255 ± 1.4</td>
</tr>
<tr>
<td>FeS</td>
<td>22.0 ± 1.4</td>
<td>651 ± 7.8</td>
</tr>
</tbody>
</table>

* Freely exchangeable iron control culture.
† Cellular iron accumulation.
‡ Fe₃(PO₄)₂·8H₂O.

Table 4. Comparison of the nitrogen-fixing growth of siderophore-negative strains at high aeration

Cultures were incubated at 28 °C with shaking at 300 r.p.m. in baffle flasks, in Burk’s nitrogen-free medium containing 0, 1 and 25 μM-iron. Growth rates were determined from plots of turbidity (OD₆₂₀) versus time during the exponential growth phase from 0 to 18 h.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Doubling time (h)</th>
<th>Siderophores in iron-sufficient culture*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 μM-Fe</td>
<td>1 μM-Fe</td>
</tr>
<tr>
<td>UW</td>
<td>3.1</td>
<td>2.7</td>
</tr>
<tr>
<td>D27</td>
<td>4.0</td>
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<tr>
<td>UA1</td>
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<tr>
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<tr>
<td>P100</td>
<td>4.6</td>
<td>3.4</td>
</tr>
</tbody>
</table>

* Iron-sufficient medium contained 25 μM-iron and was harvested at 24 h.
† Hydroxamate was determined as the A₃₈₀ of acidified culture fluid.
‡ Catechol was determined as the concentration of catechol (μM) according to the assay of Barnum (1977).
§ CAS assay results using the aqueous phase after ethyl acetate extraction (CAS-AQ) or the ethyl acetate extract (CAS-EA) of 10-fold concentrated iron-sufficient culture supernatant fluids. A positive result was indicated by a decrease in A₆₃₀ of ≥ 5 units (according to Table 2) which continued for at least 120 s. A negative result was no net decrease in A₆₃₀ over 120 s.

Aerobic nitrogen-fixing growth of strain P100

The growth of strain P100 was significantly impaired compared to strain UW under highly aerobic nitrogen-fixing conditions (Table 4). In 0FeBBG, the growth of strain P100 was the slowest, followed by the azotobactin-negative strains. As the iron content of the medium was increased to 1 μM, the growth rate of all the strains increased, but the pattern of growth rates remained the same. A further increase in iron concentration to 25 μM did not increase the growth rate of strain P100 to a value comparable with the wild-type strain. In this iron-sufficient medium the catechol-producing strains released about 11 μM-2,3-DHBA into the culture fluid and strain F196 always released a small amount of azoto-bactin. Examination of 10-fold concentrated culture supernatant fluids revealed that all of the strains produced siderophore activity according to the CAS-shuttle assay. This was attributed to small amounts of catechol siderophores produced by all catechol-positive strains, a small amount of azotobactin produced by the wild-type, definite azotobactin production by strain F196 and non-conventional siderophore production by strain P100. 2,3-DHBA alone (0-5 ml of 500 μM) did not give a positive result in the CAS-shuttle assay. The final amount of cell protein (μg ml⁻¹) produced by strains F196, D27, UA1 and P100 in this iron-sufficient medium was 80, 75, 75 and 72% of that produced by the wild-type strain.
Discussion

It appeared that *A. vinelandii* produces another siderophore in addition to the known catechol and pyoverdin siderophores. This compound functioned as a siderophore: it promoted iron accumulation from insoluble iron sources, it promoted cell growth under iron-limited conditions and its production was repressed by iron. The siderophore appeared to be a non-conventional one, in that it lacked hydroxamate or catechol groups and was only detectable in the CAS assay of Schwyn & Neilands (1987). Other non-conventional siderophores have been detected using the CAS assay, for example citric acid produced by *Bradyrhizobium japonicum* (Guerinot et al., 1990), rhizobactin produced by *Rhizobium melloti* (Schwyn & Neilands, 1987), and staphyloferrin produced by *Staphylococcus* spp. (Meiwes et al., 1990), which all coordinate iron by carboxylic acid groups. The non-conventional *A. vinelandii* siderophore is not citric acid and citric acid does not promote iron uptake into *A. vinelandii* (Knos et al., 1984). While the structure of the *A. vinelandii* non-conventional siderophore is unknown, it did appear to be a relatively low-affinity chelator that remained in the aqueous phase after ethyl acetate extraction. The siderophore did not reverse the inhibition caused by 50 μg EDDHA ml⁻¹ and only solubilized iron from the most soluble minerals tested. Previous studies have shown that the iron in FeS and vivianite can be solubilized by 2,3-DHBA alone at neutral pH, but utilization of the iron from Fe₂O₄ requires the induction of the catechol siderophores (Page & Huyer, 1984). The catechol siderophores of *A. vinelandii* are low-affinity chelators capable of bidentate or tetradentate coordination of iron (Page & von Tigerstrom, 1988). It is interesting that attempts to reconstruct the total siderophore activity of the iron-limited *A. vinelandii* culture supernatant fluid, by mixing appropriate portions of catechol and azotobactin siderophores, has never resulted in 100% of the original activity (Knope et al., 1984; Page & von Tigerstrom, 1988; W. J. Page & M. von Tigerstrom, unpublished data). It would now appear that this was because not all the siderophores had been identified.

The success of the growth of strain P100 under iron-stress conditions was affected by the iron status of the medium used for the production of the inoculum. *A. vinelandii* is known to produce bacterioferritin (Juudi et al., 1990; Steifel & Watt, 1979) and this iron reserve was apparently sufficient to promote the growth of strain P100 under conditions where exogenous iron uptake was minimal. Iron-starved cells of strain P100 were unable to grow well under nitrogen-fixing conditions at high aeration, presumably because the low-affinity non-conventional siderophore was unable to scavenge the small amount of iron present in this medium. The growth rate of strain P100 did increase when 1 μM-iron was available in the medium, presumably as a result of iron mobilization by the unconventional siderophore, but the growth rate did not increase further when 25 μM-iron was present. Both strain P100 and the wild-type should demonstrate the same siderophore-independent growth rate under these iron-sufficient conditions, as was observed with the other siderophore-deficient strains. The results showed that all the strains produced very low levels of siderophores under these conditions, which indicated that the iron in the medium must not have been freely available. Thus growth of strain P100 would be retarded according to the affinity of its siderophore for iron under these iron-stress conditions. Low-level siderophore production in B6 medium containing 25 μM-FeSO₄ was also observed by Fekete et al. (1983), when *A. vinelandii* was grown in continuous culture at high aeration. It has been shown that 50 μM-nitrilotriacetate, 10-fold more than used by Fekete et al. (1983), has to be added to B6 medium to keep the iron soluble under aerobic chemostat conditions (Partridge & Yates, 1982).

Very little has been published concerning the genetic organization and regulation of the gene products responsible for the synthesis of *A. vinelandii* siderophores. While this present study did not generate a great variety of easily identifiable phenotypes, it does demonstrate several interesting points. A single Tn5 insertion in strain F196 has inactivated the production of all known catechol siderophores, including 2,3-DHBA. This suggests that the biosynthetic genes for catechol siderophores may be organized in an operon. Alternatively, the biosynthesis of 2,3-DHBA may be a prerequisite for catechol siderophore synthesis, analogous to the requirement for the *entA*, *entB* and *entC* gene products in enterobactin biosynthesis (Ozenberger et al., 1989). However, exogenous 2,3-DHBA did not overcome the mutation in strain F196 and promote catechol siderophore biosynthesis. It also appeared that 2,3-DHBA synthesis was iron-repressible. Thus, the apparent constitutive production of 2,3-DHBA (Page & Huyer, 1984) must have been a result of the difficulty in maintaining freely soluble iron in highly aerated medium, not due to unregulated transcription.

Glick et al. (1988) used chemical mutagenesis to generate *A. vinelandii* mutants defective in siderophore synthesis. They found that strains defective in azotobactin synthesis (Flu⁻) were common, but most of these mutants were also defective in catechol siderophore synthesis and some also lacked IROMPs. In addition, mutant strains that lacked both azotobactin and azotocelin production were siderophore-negative in the CAS assay. Glick et al. (1988) concluded that siderophore synthesis in *A. vinelandii* was highly integrated and that
the synthesis of azotobactin and azotochelin was functionally coupled. The results of the present study showed that azotobactin-negative and catechol-negative mutants can be generated independently and that the loss of azotobactin does not affect the level of catechol siderophore production.

Obtaining and maintaining mutants of A. vinelandii has been a problem for years (Kennedy & Toukdarian, 1987), due to the extreme polyploidy of this prokaryote. The organism has ≥ 40 copies of its chromosome (Punita et al., 1989; Sadoff et al., 1979), which segregate randomly during cell division (Phadnis et al., 1988). Thus the selection of stable recessive mutations requires many generations to ensure that a homozygous cell is produced, at the price of losing mutations by repair or through dilution (Phadnis et al., 1988). Transposon mutagenesis provides a means of introducing single insertions and a selective pressure (antibiotic resistance), to promote the formation of a homozygous mutant cell (Kennedy & Toukdarian, 1987; Phadnis et al., 1988). It is highly likely that the chemical mutagenesis used by Glick et al. (1988) generated multiple mutations and that their conclusions are thus not valid. However, in this study it was observed that a loss of catechol siderophores did increase the level of azotobactin produced. This may relate to the functional role of these siderophores, rather than to linked transcription. The lower-affinity catechol siderophores appear to be efficient solubilizers of mineral iron sources, while azotobactin appears to be a scavenger of soluble iron (Page & Huyer, 1984; Page & von Tigerstrom, 1988). Only when the amount of catechol siderophores produced in 0FeBBGN fails to liberate iron from insoluble minerals, is azotobactin hyperproduced. Thus catechol-siderophore-negative cells may sense that they are iron-limited, even though mineral iron is present. Under these conditions, azotobactin is produced, soluble iron is scavenged and iron-limited growth is promoted, resulting in enhanced azotobactin production.

It is clear that A. vinelandii has developed elaborate systems for iron acquisition, attesting to the importance of this micronutrient. However, the majority of the iron-repressible mutants generated in this study were not identified as being defective in siderophore production or iron uptake. Iron limitation of A. vinelandii is known to enhance capsule formation (Jarman et al., 1978), to enhance encystment (Layne & Johnson, 1964) and to induce genetic transformation competence (Page & von Tigerstrom, 1978). Further characterization of these mutant strains and identification of the non-conventional siderophore is in progress.

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References


