# Sequence analysis of a DNA fragment from Sinorhizobium fredii USDA257 which extends the nitrogen fixation host range of Rhizobium species NGR234 to soybean, Glycine max (L.) Merr cultivar Peking

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#### Abstract

A fragment of DNA (pBTBX) from the genome of Sinorhizobium fredii USDA257 was sequenced by shotgun strategy to identify the potential genes which enabled the Rhizobium species NGR234 to fix nitrogen on soybean, Glycine max (L.) Merr cv. Peking. The total length of the cosmid is 32,824 base pairs with a GC content of 61%. A 29 open reading frames (ORF) were identified representing 71.8% (23,574 bp) of the cosmid. Out of these ORF, 96.5% (22,749 bp) were identical and similar to reported and hypothetical genes and proteins. The remaining 3.5% (825 bp) had no apparent similarity to any genes in the data base. Gene and gene products found on the DNA fragment include those involved in the synthesis of Fe-Mo component of nitrogenase, regulation of nitrogen fixation, transport of amino acids and sugars, chemotaxis and transcriptional regulation.

Keywords: Sequence analysis, DNA fragment, Sinorhizobium fredii, Rhizobium species NGR234, soybean Peking

## 1. Introduction

Generally, legumes fix nitrogen in symbiotic association with compatible bacteria collectively known as rhizobia. Rhizobia are grouped majorly into Rhizobium. Bradyrhizobium, Azorhizobium, Mesorhizobium Sinorhizobium. During the symbiosis, signals to and from both the macro- and micro-symbionts are released (Fischer, 1994). This exchange of chemical signals between soil bacteria (rhizobia) and legumes termed a molecular dialogue involves two main groups of molecules: nod geneinducing flavonoids from plants and the mitogenic lipochito-oligosaccharide Nod factors (NFs) of rhizobia. The NFs synthesized by rhizobia elicit, at very low concentrations and in a specific manner, various symbiotic

responses on the roots of the legume hosts (Debellé et al., 2001; Shaw et al., 2006; Steinkellner et al., 2007). This is because rhizobia respond to chemoattractants and growthenhancing compounds in root exudates, and several plant non-flavonoids possess nod gene-inducing properties (Cooper, 2007). A number of nodulation genes which specify the synthesis of NFs have been identified. All rhizobia, in spite of their diversity, possess conserved nodABC genes responsible for the synthesis of the Nacylated oligosaccharide core of NFs, which suggests that these genes are of a monophyletic origin. Other genes, the host specific nod genes, specify the substitutions of NFs (Debellé et al., 2001). Entry into the plant is restricted to bacteria that have the "keys" to a succession of legume "doors". Some symbionts intimately associate with many different partners (and are thus promiscuous), while others are more selective and have a narrow host range. This is related to difference in type of NF produced by rhizobia

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Table 1. Predicted genes and proteins encoded by pBTBX of S. fredii USDA257 which confers on NGR234 the ability to fix nitrogen on G. max cv. Peking.

Orf	Base posit. in plasmid		Frame	RBS' and start codon	Homolog gene or amino acid position in plasmid	name	Gene base/ amino aci length	Organism d	Protein access no.	Identity (%)	Similarity (%)	/ Description
bt1	1-213		-3	gGGAGc-T-4ATG	34-186	BJORF	1447	BJ	M17635	67	67	DNA region with 3
					72-165	YBIO	786	E. coli	P75783	37	59	ORF complete cds Hypothetical 86.8 kD protein in Ding-GLNQ inter-
bt2	294-539	cat	-2	cGGcGGc-9-ATG	330-509	SCRSCAT	C 212	S. cerevisiae	M12865	64	64	genic region CAT repetitive element, clone pYCAT8
					369-522	NPT2	68	Mus musculus	Q62111	34	59	Na+.PO4 cotrans- porter type III fragment
ot3	1296-1691	gpmB	-3	cGccGGc-5-GTG	1357-1614	GPMB	349	Syn. sp.	P72649	35	54	Phosphoglycerate mutase
ot4	1631-4093		-1	cGGAGGT-4-ATC	i 1691-4079	?	821	Syn. sp	P74690	61	74	Hypothetical 92.4 kD protein
ot5	4491-4615		1	gcGAGaa-5-TTG	4495-4614	U650AH	55	Myc. leprae	Q50135	35	47	Hypothetical 5.8 kD protein
ot6	4672-5394		-3	AGGAccT-4-ATG	4708-5368	?	248	Sal. typhi	P74886	24	44	High affinity peri- plasmic glutamine binding protein
bt7	5397-7775	nifL	-2	gtGtGGT-10-ATG	6393-7689	NifL	840	Syn. sp.	P72843	36	55	Nitrogen fixation positive activator protein, NifL
ot8	8046-8954	gcvA	-2	AGGaaa-7-ATG	8076-8946	GCVA	305	E. coli	P32064	37	60	Glycine cleavage system trans- criptional activator
ot9	9108-9767		3	AGGcGGc-6-ATG	9164-9671	SGKSACP	G 5364	Str. griseus	X77865	62		Ketosynthase, Acyl carrier protein, ketoreductase, cyclase and dehydrase
t10	10162 -10896	nodG lar	1	gtGAGGT-13-ATC	9108-9753 G 10219 -10696	NODG RNU87960	245 2479	R. meliloti Ratus norvegicus	P06234 U87960	33 62	62	NODG Leukcoyte common antigen receptor (LAR) gene trans- spliced alternative untranslated exon
ot l l	10969 -11562		1	AGGAGGg-3-ATC	G 11006 -11445	MCU60315	190,289	Virus	U60315	61	61	Mollusum contagiosum virus subtype, complete
t12	11660		2	gGGAGGa-4-ATG		RBSA	493	H.	P44735	37	59	genome cds Ribose transport
bt13	-13156 13173 -14237	mxa	3	gGGAGaa-4-ATG	-13136 13218 -14216	AF017434	4815	influenzae Mt. extorquens	AF01743	4 58	58	ATP-binding protein Methanol oxidation genes mxaE, mxaH, mxaB and pmi-like genes, complete genome cds
					13290 -14187	Y4mJ	333	NGR234	P55569	30	54	Predicted ABC trans- porter ATP-binding protein Y4mJ
t14	14285 -15385	тср	2	gGGAGGa-7-ATG	14285 -15197	RCMCPAE	5186	Rh. capsulatus	L48927	59	59	Methyl-accepting chemoreceptors (mcpA and mcpB) genes complete genome cds
					14444 -15305	Y4mL	324	NGR234	P55568	24		Predicted ABC trans- porter periplasmic binding protein Y4m precursor
ot15	15431 -16450	gntR	-1	AGGAGGa-4-GTC	5 15489 -16445	GNTR	331	E. coli	P46860	31		GNTR utilization system GNT-1 trans- criptional repressor

Table 1. Continued.

-	2	1 2	AGGAGaa-3-ATG cGGAGtc-3-ATG	16754 -17545 16774 -17383 17672 -18284 17663 -18284 18373 -18943	AVINIFE Y4tE Y4tF Y4tG	1509 300 238 231	Az. vinelandii NGR234 NGR234 NGR234	X07293 P55659 P55660 P55661	56 25 30 28	56 47 49 50	nijE gene  Predicted ABC transporter periplasmic binding protein Y4tE precursor Predicted ABC transporter periplasmic binding permease Y4tF Predicted ABC transporter periplasmic binding permease Y4tG
512-1831 313-1897 953-1968	2	1		16774 -17383 17672 -18284 17663 -18284 18373 -18943	Y4tF Y4tG	238	NGR234 NGR234 NGR234	P55660 P55661	30 28	49	porter periplasmic binding protein Y4tE precursor Predicted ABC trans- porter periplasmic binding permease Y4tF Predicted ABC trans- porter periplasmic binding permease Y4tG
313-1897 353-1968	2	1		-18284 17663 -18284 18373 -18943	Y4tG	231	NGR234	P55661	28	50	Predicted ABC trans- porter periplasmic binding permease Y4tF Predicted ABC trans- porter periplasmic binding permease Y4tG
953-1968			cGGAGGT-6-ATG	-18284 18373 -18943							porter periplasmic binding permease Y4tG
953-1968			cGGAGGT-6-ATG	-18943	Y4tF	238	NGR234	P55660	30	51	-
-	4	2		18331						٠.	Predicted ABC trans- porter permease protein Y4tF
-	4	2		-18952	Y4tG	231	NGR234	P55661	28	49	Predicted ABC trans- porter permease protein Y4tG
	t19 18953-19684		AGGAGGc-3-ATG	18965 -19673	Y4tH	257	NGR234	P55662	48	65	Predicted ABC trans- porter ATP-binding
20 20003-20914		2	tatcGGT-5-ATG	20078 -20417	MJ0604	100	M. jannaschi	Q58021	34	50	protein Y4tH Hypothetical protein MJ0604
487-2191 162-2234		1 -3	cGGAGat-5-ATG AGGctGa-6-TTG	N.o. 22186 -22338	N.o. YDJA	N.o. 183	N.o. E. coli	N.o. P24250	N.o. 40	N.o. 53	Hypothetical 16.2 kD Hypothetical 20.1 kD protein in SELD- SPPA intergenic region (ORF 183)
124 599	snaC	-1	cGGAGac-7-ATG	22746 -23509	SP21216	770	SPS	U21216	60	60	NADH-FMN oxido- reductase (snaC) gene, complete cds
				22835 -23597	?	351	Ps KHP41	005599	34	45	Integrase-like protein
				22772 -23594	RECR	343	BPP1	P06956	26	46	Recombinase CRE
124 23670-23948			_	23709 -23937	REPC	314	Staph. aureus	P03064	24	49	Replication initiation protein (protein A)
t25 23991-24377				23991 -24369	MTCY20 H10.05	131	Myc. tuber	P96914	61	75	Hypothetical 14.4 kD protein
26 24925-25317			-	N.o.	N.o.	N.o.	N.o.	N.o.	N.o.	N.o.	Hypothetical 14.7 kD protein
27 25428-25991			-	25428 -25977	•					82	Hypothetical 36.1 kD protein Y4jO
988-2709	1	-1	AtGAttc-9-ATG	26687 -27086					61	75	Hypothetical 36.1 kD protein Y4jP
		3	AGccGGT-7-TTG	25994 -26588 31465	•	262 99			63	73 42	Hypothetical 29.5 kD protein Y4jP MCP protein
92	91-2437 25-2531 28-2599 88-2709	01-24377 25-25317 28-25991 38-27091 32 mcp	25-25317 -2 28-25991 -2 38-27091 -1	21-24377 -2 AcGgGcT-9-ATG 25-25317 -3 tGGcGGg-10-GTG 28-25991 -2 tGGActg-8-GTG 28-27091 -1 AtGAttc-9-ATG	-23594 -23594 -23709 -23937 -23937 -23937 -23937 -24369 -24369 -25-25317 -3 tGGcGGg-10-GTG N.o.  -28-25991 -2 tGGActg-8-GTG -25977 -38-27091 -1 AtGAttc-9-ATG -27086 -25994 -26588 -26588 -27090 -26588 -27086 -26588 -27086 -26588 -27086 -26588 -27086 -26588 -27086 -26588 -27086 -26588 -27086 -26588 -27086 -26588 -27086 -26588 -27086	70-23948 3 AGGgGGa-8-ATG 23709 REPC -23937 -23937 -23937 -24377 -2 AcGgGcT-9-ATG 23991 MTCY20 -24369 H10.05 -25-25317 -3 tGGcGGg-10-GTG N.o. N.o.  28-25991 -2 tGGActg-8-GTG 25428 Y4jO -25977 -25977 -27086 -27086 -25994 Y4jP -26588 -26588 -262 mcp 3 AGccGGT-7-TTG 31465 MCP	70-23948 3 AGGgGGa-8-ATG 23709 REPC 314 -23937 -23937 -23937 -2 AcGgGcT-9-ATG 23991 MTCY20 131 -24369 H10.05 -25-25317 -3 tGGcGGg-10-GTG N.o. N.o. N.o. N.o28-25991 -2 tGGActg-8-GTG 25428 Y4jO 321 -25977 -25977 -26687 Y4jP 321 -27086 -27086 -27086 -27994 Y4jP 262 -26588 -262 mcp 3 AGccGGT-7-TTG 31465 MCP 99	-23594 70-23948 3 AGGgGGa-8-ATG 23709 REPC 314 Staph23937 aureus 71-24377 -2 AcGgGcT-9-ATG 23991 MTCY20 131 Myc. tuber -24369 H10.05 75-25317 -3 tGGcGGg-10-GTG N.o. N.o. N.o. N.o. 78-25991 -2 tGGActg-8-GTG 25428 Y4jO 321 NGR234 -25977 78-27086 25994 Y4jP 321 NGR234 -27086 -25994 Y4jP 262 NGR234 -26588 70-23948 3 AGccGGT-7-TTG 31465 MCP 99 Myc. tuber	70-23948 3 AGGgGGa-8-ATG 23709 REPC 314 Staph. P03064 aureus 01-24377 -2 AcGgGcT-9-ATG 23991 MTCY20 131 Myc. tuber P96914 -24369 H10.05 15-25317 -3 tGGcGGg-10-GTG N.o. N.o. N.o. N.o. N.o. N.o. 128-25991 -2 tGGActg-8-GTG 25428 Y4jO 321 NGR234 P55515 -25977 26687 Y4jP 321 NGR234 P55515 -27086 25994 Y4jP 262 NGR234 P55516 -26588 12 mcp 3 AGccGGT-7-TTG 31465 MCP 99 Myc. tuber E1191436	-23594 70-23948 3 AGGgGGa-8-ATG 23709 REPC 314 Staph. p03064 24 23937 aureus 70-24377 -2 AcGgGcT-9-ATG 23991 MTCY20 131 Myc. tuber P96914 61 25-25317 -3 tGGcGGg-10-GTG N.o. N.o. N.o. N.o. N.o. N.o. N.o. N.o	70-23948 3 AGGGGGa-8-ATG 23709 REPC 314 Staph. p03064 24 49 -23937 aureus  101-24377 -2 AcGGGCT-9-ATG 23991 MTCY20 131 Myc. tuber P96914 61 75 -24369 H10.05 N.o. N.o. N.o. N.o. N.o. N.o. N.o. N.o.

Posit., position; funct., functional; RBS, ribosome binding site; the number between RBS and start codon (for example 4 in gGGAGcT-4-ATG) indicates the position of RBS relative to the translation initiation start site. Az, Azotobacter; Bj, Bradyrhizobium japonicum; BPPI, Bacteriophage P1; E, Escherichia; H, Haemophilus; M, Methanococcus; Mt, Methylobacterium; Myc. tuber, Mycobacterium tuberculosis; NGR234, Rhizobium species NGR234; Ps, Pseudomonas; R, Rhizobium; Rh, Rhodobacter; S, Saccharomyces; Sal typhi, Salmonella typhimurium; SPS, Streptomyces pristinaespiralis; Staph, Staphylococcus; Str, Streptomyces; Syn, Synechocystis; N.o., None obvious.

(Perret et al., 2000; Debellé et al., 2001; Shaw et al., 2006; Steinkellner et al., 2007). At the onset of the association, exudates such as flavonoids (the strongest *nod* genes inducers) are secreted by plant roots into the rhizosphere.

Following root hairs colonization by the bacteria,

common *nod* genes (*nod*ABC) are expressed in the microbial cells under the control of the regulatory *nod* genes (*nod*D, *syr*M or *nod*UVM) (Fellay et al., 1995; Hanin et al., 1998). This results in the biosynthesis of the lipooligosaccharide backbone of Nod factors which are

decorated by the products of host-specific *nod* genes; the action of which causes root hairs to deform, branch, curl, form infection threads.

Rhizobium species NGR234 and Sinorhizobium fredii USDA257 are both closely allied broad-host-range rhizobial strains. The former bacterium distinctly nodulates more than 110 genera of legumes as well as the non-legume Parasponia andersonii but induces non-nitrogen-fixing nodules on soybeans (Trinick, 1980; Lewin et al., 1987; Relic et al., 1994; Pueppke and Broughton, 1999). In contrast, USDA257 forms effective nodules on many cultivars of soybeans including cv. Peking (Keyser et al., 1982; Balatti and Pueppke, 1992).

Recently, we screened a genomic DNA library of USDA257 and identified a clone containing a DNA fragment, pBTBX which endows NGR234 with the ability to fix nitrogen on *Glycine max* (L.) Merr cv. Peking. Nodule-like structures formed by the wild-type NGR234 lack rhizobial cells in comparison with bacteroids-containing nodules produced by the transconjugant, NGR234(pBTBX) on the soybean. In order to identify and study the molecular basis of the genetic locus/loci on the pBTBX which confers this capability on NGR234, we have shotgun-sequenced the DNA fragment and complemented NGR234 with a gene segment from the pBTBX. Here, sequence analysis of the DNA fragment is presented.

1kb

#### 2. Materials and Methods

Construction of shotgun library

Procedures employed in this work were according to Hanahan (1983) and Sambrook et al. (1989). Cosmid DNA (pBTBX) was prepared from a late log-phase grown cells of the clone containing the DNA fragment (obtained from Sinorhizobium fredii USDA257 which enabled NGR234 to fix Nitrogen). The DNA (pBTBX) was sonicated and ends of fragments were filled-in with nucleotides in the presence of T4 DNA polymerase. DNA fragments ranging in size from 1.6 to 3.0 kb were ligated to Sma1-cleaved M13mp18 at 16°C. Aliquots of ligated DNA were used to transform competent cells of DH5α. Recombinant phages were prepared from the transformed DH5α and used for sequencing.

Sequencing and nucleotides assembly

This sequence analysis was carried out at the Institut für Molekulare Biotechnologie, Abteilung Genomanalyse, Jena in Germany. Sequencing was carried out using Dye Terminator/Thermo Sequenase sequencing methods essentially as reported by Freiberg et al. (1996).

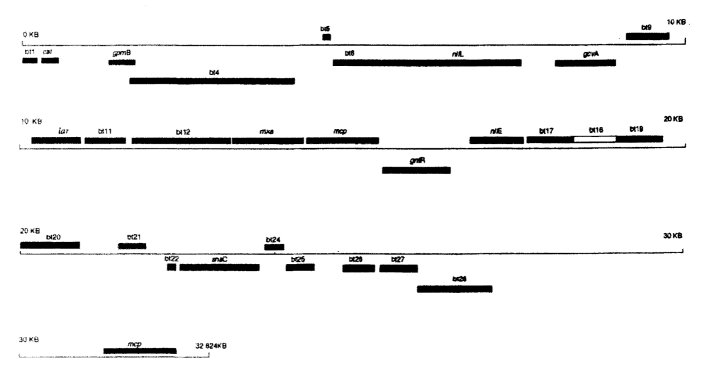


Figure 1. Genetic map of pBTBX. Open and shaded rectangles denote genes. ORF names or names of genes (for example bt1 or *cat*) correspond to those in Table 1. Genes positioned on top and below each line are transcribed from the forward and complementary DNA strands, respectively.

ORF bt27	* ** ** ** * * *
bt27: (1) y4jO: (137) Identity:	VIVSDDAGQFRVANHALCWVHTERLLQKLMPATPKE QRLVTT TRDLVWRFYKALKVWKQQ VIVSDDAGQFRVGNHALCWVYAERLLQKLMPATPRQVRQVEAVRDLVWRFYRALKS V KRK VIVSDDAGQFRV NHALCWV ERLLQKLMPATP R V RDLVWRFY ALK K
bt27: y4jO: Identity:	PSPQL I NGFRRFEQIFARRTGYAA LDKLLLRLHRRKAELLKVLEHPYIPLHTNASEND IRS PPPGLAAAFRKRFARIFS LRTGYE DLDKLLARLSRRKD ELLKVLERPDIPLHTNASENDLRS P P L FR RF IF RTGY LDKLL RL RRK ELLKVLE P IPLHTNASEND RS
bt27: y4jO: Identity:	* **
ORF bt28 bt27h	* *** ** ** **
bt27h: (233) y4jO: (1) Identity:	VLHAGLV SAPFITVDDTGARHNRRNAFTTQIGGERF STFRTSLSKSRLNFLSVLRAGHQGYVLNDE MLHAGLVSAPYITVDDTGARHARDSFHTTQIGAEHFTAFRTTASKSRLNFLS+LRG SYQDYVLNDA LHAGLV SAP ITVDDTGARH R TTQIG E F FRT SKSRLNFLS LR Q YVLND
bt27h: y4jO: Identity:	* * * * * * * * * * * * * * * * * * *
bt28	
bt28: (2) y4jP: (17) Identity:	KKRLPSVEH I ETLSLLAMRRLVGGLVEELQALKAEVATLR S ENEALREDNAQLRLDNARLKAENQQL KKRLASPEHADTLSLKAL RVLVTGLVDEVKELSAEVT TLHAENAALREDNEAL RLENT RLKVENQLL KKRL S EH TLSL A R LV GLV E L AEV TL EN ALREDN L RL N RLK ENQ L
bt28: y4jP: Identity:	** * ** ** * * * * * * * * * * * * * *
bt28: y4jP: Identity:	* * * * * * * * * * * * * * * * * * *

Figure 2. Alignments of protein sequences encoded by open reading frames bt27 and bt28 with y4j0 and y4jP of R. species NGR234. bt27h is bt27 homolog which exist in the same ORF with ORF bt28. Amino acid numbers in the protein corresponding to the first and last residues in the alignments shown are presented in parentheses. Amino acids identical in both proteins are shown below the alignments. Conserved leucine, alanine, glycine, proline and isoleucine are indicated with asterisks.

Nucleotides were compiled and edited after those for M13mp18 were removed. Gaps between contigs were closed with additional sequences obtained from dye primer and big-dye terminator sequencing methods.

# Analysis of nucleotides

Open reading frames and intergenic regions were looked for in the nucleotides by using *Rhizobium meliloti* matrix of GeneMark prediction program version 2.3 (Borofsky, 1995). Search for similar and identical protein and gene sequences was conducted with the GCS computation program using the BLAST Network Service.

## 3. Results and Discussion

The DNA fragment (pBTBX) is 32,824 base pairs long with a GC content of 61%. Twenty nine Open Reading Frames were detected in the pBTBX (Table 1, Fig. 1). The initiation codon is ATG for most of the genes except for seven cases where GTG and TTG are used. The putative ribosome binding sites are generally rich in purines and showed variable locations in the sequence. Collectively, 72% (23,574 bp) of the pBTBX are potentially coding (Fig. 1). Two and half percent (825 bp) of the sequence show no significant similarity to any known genes or gene products in the database bank. Nucleotide content of the non-coding regions is essentially the same as that of the coding parts.

-35

ATGACGATCTACCAAGTGGaggagaaACCATGACGATTTCG

-10

btnifE:

btnifE:

AzvnifE:

btnife: (13) AACTTTGTGCGGACAATGGT-GGC-A-GCCATCGGAATAGCG-GCG-GCCGCCTCGTT Azvnife: (451)ACCTGCGTGCCGGCGCTGATCGGCGACGACGTCGACGCACTGTGCAAAGCCGCCGCCGCCA-GCGGCTGCGGAGGCGCAGGCCGCCACGCTGAACGACATCATCTCGCGCGGTACCGTGCGC AzvnifE: btnifE: ATCGGAGTCCTGACCGCCGCCCCCCATGGGAATGGTCGACGAACAGGGCAACC-CAA- $\mathtt{CTCGGCAACCGCATCGCCGGTGAGGCCATGCTCAAGTACGT} - \mathtt{GATC} - - \mathtt{GGCACCCGCGAG}$ AzvnifE: btnifE: CCGGCTACGACGTCGACGTGGCCAATCTGATTG-CCGGCTATCTGTC-GCT-GCCGGTCG  $\underline{\mathtt{CCCGATCCG}}\mathtt{CT}\underline{\mathtt{CCC}} - \underline{\mathtt{CGT}}\mathtt{C}\underline{\mathtt{GGCAG}} - \underline{\mathtt{C}} - \underline{\mathtt{GA}}\mathtt{GC}\underline{\mathtt{GT}}\underline{\mathtt{CCGG}}\mathtt{GC}\underline{\mathtt{ATC}}\mathtt{C}\underline{\mathtt{GCGT}}\underline{\mathtt{GCACGACG}} - \mathtt{TCA}$ AzvnifE: AGCTCGT-GCCGCTGACGCCGCCGGCT-CGCATTCCCGCTT-TGCAGAC-CGGCAAGGTC btnifE: AzvnifE:  $\underline{\mathtt{ACCT}} \mathtt{GATCGGCGAGT} \underline{\mathtt{ACATCGCCGGCGAGTTC}} \mathtt{TGCATGTCCTGCCGCTGCTCGA-C}$ btnifE: GATTTCCTCGT-CGCCACC-CTGGCGCC-GACCGGGGA-GCGCGCCAAGACGGTGATGTT AzvnifE:  $\underline{GAACT}GGGCCTGCGGGTGCTCTGCACCCTGGCCGGCGATGCGGCGCTACCGCGAGG-\underline{TG}CA$ CACCCAGCCCTACAGCGCCTTCAACATGGACATCATCTCCG-GTCCCGAC-CA--GAAAT btnifE: AzvnifE:  $\underline{GACCATGCACCGCGCCGAAGTGAACATGATGGTCTGCTCCAAGGCCATGCTCAATGTCGC}$ TTGCAAAGCTTGCCG-ATCTCGAAGGCAAGCGCGTCGC-CGTCAACCGTGGCT-CGTCGbtnifE: AzvnifE: TCGCAA-GCT-GCAGGAAACCTACGGCACGCCC-TGGTTCGAGGGCAGCTTCTACGGCAT CAG-GAGACGCCCTGCGCAAGGCGGCAGTTC-CCGGCCTGGAAATCGTCG-TCTACGAG btnifE: AzvnifE: btnifE: GATGATTCCACCAGCGCACAGGCGCTGATCGCCGGCCAGGTCGATGCGGTC-GCGCTGCC AzvnifE:  ${\tt CTCGA-CGGTCG-GTGAGGCGATCATCAAG-CAGCGCCCAGATGCCGGTCTGCAG-GTTG}$ btnifE: AzvnifE:  $\underline{\mathtt{CTCGAACCCTGGCGTGCG}} - \underline{\mathtt{CG}} - \underline{\mathtt{TCTGGAGGGCAA}} - \underline{\mathtt{GCGC}} - \underline{\mathtt{G}} - \underline{\mathtt{TGCTGCTCTACACCCGGCG}}$ btnifE: GCTTCACCTTCTTCCAGCAGGGCAATTCGATGGCG-ACCCGGATGGAGGACTTTGAGATC AzvnifE:  $\tt GCGTGAAGTCCTGGTCGGTGGTTTCCCCCCTGCAGGACCTGG--GCATGAAGGTG-\underline{G}-\underline{TC}$ btnifE: CGCCAGTGGCTCAACACCGCCATCTACCTGATGAAGATCTCCGGCG-ATCT-CGACAAGA -GCCACCGGCACCA-A-GAAGTCCACC-GAGGAAGACAAGGCACGCATCCGCGAACTGA AzvnifE:

Figure 3. Nucleotide sequence of the *nif*E homologue (btl6*nif*E) and the homologous gene from *Azotobacter vinelandii* (Azv*nif*E). First and last nucleotides in the alignments shown are indicated in parenthesis. Positions of identical nucleotides are underlined. Putative ribosome binding site is shown in bold lower-case letters; hypothetical translation initiation codon is asterisked. Introduced gaps to give the best alignments are indicated with hyphens. Presumed Pribnow boxes are overlined.

TCG-CGACGAAGTGGACCGGC-CGCCCGATGCCGACGCTTCCGTCCTTCTGA (804)

TGGGCGACGTCAAGATGCTCGAC-GAGGGCAATGCGCGGGTGCTGCTGA (1243)

The highest similarities found were with proteins of the closely related NGR234 (Table 1). The hypothetical proteins of ORFs bt27 and bt28 show strong identities of 71% and 63% and, similarities of 82% and 73%, respectively to the hypothetical y4jO and y4jP proteins of NGR234. Generally, the proteins are hydrophobic (Fig. 2). Thus, they may be important in the formation of cell structural components involve in transport systems. Other ORFs that have homology to transporting genes lie at 294 to 539bp (ORF bt2), 4672 to 5394bp (ORF bt6) and between bases 11660 and 19684bp (ORFs bt12, bt13, bt14, bt16, bt17, bt18 and bt19; Table 1). These are probably ATP binding cassette (ABC) type transporters except for bt2. The bt6, bt16, bt17, bt18 and bt19 show

similarities ranging from 44 to 65% to the periplasmic glutamine binding protein of Salmonella typhimurium, y4tE, y4tF and y4tG, the ABC transporter binding protein and permeases of NGR234. The bt12, bt13 and bt14 show similarity matches to the ribose transport ATP-binding and ABC periplasmic binding proteins. The ABC superfamily is generally made up of multicomponent primary active transporters, capable of transporting both small and to ATP hydrolysis macromolecules in response (Paulsen et al., 1997). These transporters may influence nitrogen fixation considerably. Two genes (prsDE) coding for protein secretion which showed Fix defects in nodule bacteroids formed by R. leguminosarum by. viciae and R. leguminosarum bv. trifolii in pea and vetch appeared to

bt7NIFL:	(332):	LAVMTDVT-ELKTAEQRNHVQAITDHLTGLLNRPGFEMALDAAIRQTAEDGGELACLLID
SynNIFL:	(390):	VAVKEDITKEKQQAEALFH-QAHYDHLTGLPNRILAKDRLQQAIESALRQKHIFGLMFLD
bt7NIFL:		LDRFKQINDNLGHAAGDEVLRQIAGRIRAQVRGEDKVGRLGGDEFVVLIPASKAQNAALQ
SynNIFL:		LDNFKKVNDTLGHDAGDQLLVEVSERLQRALRQTDTVARLGGDEFLIILDQVSHSRKLMA
bt7NIFL:		ISERIAAACAEPVIVDGNTLSLSASIGIALYPIQAITAAELLQKSDMAMYARKHNGKNGA
SynNIFL:		IAQRLLRVMRQPVNLQGLEFFVHGSIGITVFPDDGFHADVLLRNADTAMYAAKLAGRNMF
bt7NIFL:		KLFDPRMASLAQERLKIDTYIEEGLRQDWFDVHLQPIVDLKVGRIAGFEALMRLNHPEHG
SynNIFL:		RFFTPHMNQAAQQRMAIESELRQGLSRQEFQILYQPIVSLESGQIVGAEALMRWHNRLLG
bt7NIFL:		VLPPADIIRVAEETGAILRIGERIFEKAVAHLARLTSVPGLENAYLAVNFSPLQFCPK-L
SynNIFL:		TVPPDQFIPIAEEVGLIVELGEWLLDNVCCQAAHWHSALGEQTFWVSVNVSPRQLKDSYF
bt7NIFL:		PASTVSTLMKWGITPSRIVIEITEAVLMHHSPDIRDVLGALSSAGMKIALDDFGTGYSSL
SynNIFL:		VAILQGFLQRYQVRPEWLELEITENLILEENGDLLKNLSDLEEENIALSLDDFGTGYSSL
bt7NIFL:		SYLVHFPVNIIKIDQAFTRSLTDESEMVRRRVRKLVAGIHTVAKELNCQVVAEGIETEEQ
SynNIFL:		NYLRKFNFNSLKIDRSFVELLPHDNNTVGL-VRAIIAMAHHLELKVIAEGIETPEQ
bt7NIFL:		LNALLSLDVNSGQGY (764)
SynNIFL:		WNFLRLQGCDYGQGY (818)

Figure 4. Comparison of amino acid sequence of protein bt7NIFL with the homologous protein (SynNIFL) from the nitrogen-fixing cyanobacterium, *Synechocystis* species. Amino acid numbers in the protein corresponding to the first and the last residues in the alignments shown are presented in parentheses. Identical residues are in upper-case letters.

be involved in the formation of an ABC-type transporter (Finnie et al., 1997; Kröl and Skorupska, 1997).

The bt2 is another type of transporter having 59% homology with Saccharomyces cerevisiae Na<sup>+</sup>-PO<sub>4</sub> cotransporter type III fragment (NPT2) of sodium-dependent phosphate transport integral membrane protein. The Na<sup>+</sup>-PO<sub>4</sub> cotransporter (NPT2) play a role in active transport of phosphate into cells via sodium cotransport which belongs to the anion-cation symporters (ACS) family. These ACS are widely distributed in nature occurring in gram-negative and gram-positive bacteria, and in both animal and fungal eukaryotic kingdoms (Stephanie et al., 1998). From the sequence data obtained in this study, it is suspected that this cotransporter may be involved in the exchange of sodium and/or phosphate between the bacteroids and the cells of cv. Peking.

The ORF bt16 possesses 56% similarity to the *nif*E gene of *Azotobacter vinelandii* as against 47% similarity it shows to ABC transporter, Y4tE. This ORF bt16 and ORF bt7 may be directly involved in nitrogen fixation. The *nif*E gene (ORF bt16) lies within the Fix<sup>+</sup> segment (Fig. 3). The *nif*E is involved in the synthesis of nitrogenase molybdenumiron (MoFe) cofactor. The MoFe component of nitrogenase (dinitrogenase) is directly involved in the final step of dinitrogen reduction to ammonia. Since NGR234 possesses a copy of this gene on its symbiotic plasmid (Freiberg et al., 1997), the dinitrogenase on the pBTBX may com-

plements that of the wild-type NGR234 to establish an effective symbiosis with G. max cv. Peking.

The ORF bt7 encodes nitrogen fixation positive activator protein. This protein exhibits a similarity to the nifL of Synechocystis species, a nitrogen fixing cyanobacterium (Fig. 4). The nifL is a nif-specific regulatory gene which with nifA is involved in the global nitrogen regulatory (Ntr) system which controls nif genes expression in Klebsiella pneumoniae in response to environmental oxygen (Roelvinic and van den Bos, 1989; Fischer, 1994). Generally in rhizobial species aerobiosis directly interferes with NifA activity, the nifA is sensitive to oxygen in vivo. In K. pneumoniae, NifL is required for this control. The nifL also regulates the activity of NifA in response to nitrogen conditions in K. pneumoniae, a control mechanism that is not found in rhizobia (Fischer, 1994). It is assumed that, in addition to the activity of nifA in NGR234, the regulation of nitrogen fixation in the transconjugant, NGR234(pBTBX) is at least partially under the control of NifL. It is also suspected that the nifL homologue on the pBTBX renders the transconjugant NGR234(pBTBX) insensitive to nitrogen nutrient supply to the bacteroids by the cv. Peking cells, thereby enabling the bacterium to fix nitrogen effectively.

A gcvA (ORF bt8) homologue is located on the DNA. A comparison of the deduced amino acid sequence of ORF bt8 revealed homology to GCVA, a transcriptional

LPPLNALRMFESSARHLNFRVASEELGVTOGAVAOOVRALEOHLNTRLFDRRARGLELTEAGRRYMKFLORA ht8: LPPLNALRVFDAAARHLSFTRAAEELFVTQAAVSHQIKSLEDFLGLKLFRRRNRSLLLTEEGQSYFLDIKEI GcvA: MAAVNLRHIEIFHAVMTAGSLTEAAHLLHTSQPTVSRELARFEKVIGLKLFERVRGRLH-PTVQGLRLFEEVQR LysR: MRFKGLDLNLLVALDALMTERKLTAAARSINLSQPAMSAAIGRLRAYFNDELFLMQQRRL-VPTPRAEALAPAVRE NodD: MTRSYLPLNSLRAFEAAARHLSFTHAAIELNVTHSAISQHVKTLEQHLNCQLFVRVSRGLM-LTTEGENLLPVLND AmpR: 354 43 4 4 3 333 33 4 5 4 3 4 3 3 4 4 3 4 3 55 4 4 5 4 3 3 3 PLN LR FA ARHLSFT AA EL VTQ AVS LE N LFR R L L T LP Consensus: LSLISDATQEL--KPATTVVTISVTPSFAAKWLIPRLGKFSDSNADTQVQVLAGNGLANFQSDGVDLAVRQTKPPF bt8: FSOLTEATRKLOARSAKGALTVSLLPSFAIHWLVPRLSSFNSAYPGIDVRIQAVDRQEDKLADDVDVAIFYGRGNW GcvA: LvsR: SWYGLDRIVSAAESLREFRQGELSIACLPVFSQSFLPQLLQPFLAR-YPDVSLNIVPQESPLLEEWLSAQRHDLGL ALHIOLSVIANDPLVPAESDRRFRIVLSDFMTLVFFEKVIKRVAREAPGVSFEL-LHVNDDPDERLRSGDLDFLIL NodD: AmpR: SFDRIAGMLDRFANH--RAQEKLKIGVVGTFATGVLFSQLEDFRRG-YPHIDLQLSTHNNRVDPAAEG---LDYTI 3 3 3 3 Consensus: GPGLTGELLFPMRFIAVCSPTIVA-ESQIRTVEDLTRHVLLHDAHGM-WPVFLERAGVAADVRTLKSLRFSHSSLA bt8: -PGLRVEKLYAEYLLPVCSPLLLTGEKPLKTPEDLAKHTLLHDASRRDWQTYTRQLGLN-HINVQQGPIFSHSAMV GcvA: TETLHTPAGTERTELLSLDEVCV-LPPGHPLAVKKVLTPDDFQGENYISLSRTDSYRQLLDQLFTEHQVKRRMIVE LysR: PDQFMSATHPSAKLFEDKLVCVGCPSNQQLRGKLSLKRFMSMGHVAAMFGRTLKPSIEQW---LLLEHGFKRRVEI NodD: RYGGGAWHGTEAEFLCHAPLAPLCTPDDIAASLHSPADILRFTLLRSYRRDEWTAWMQAAG----EHPPSPTHR AmpR: 3 33 33 3 33 3 GL ADVL L Consensus: IDAAVSGOGIALATEALVEDDLAAGRLCRPLDFAVTDELGFYIVHPRSPHKAGHVRSMRDWLLAQSHGSRQ (291) Bt8: GcvA: LOAAIHGQGVALANNVMAQSEIEAGRLVCPFNDVLVSKNAFYLVCHDSQAELGKIAAFRQWILAKAAAEOE (305) THSAASVCAMVRAGVGISVVNPLTALDYAASGLVVRRFSIAVPFTVSLIRPHRPSSALVQA ......(300) LysR: NodD: VVPGFNSIPMLLQGTNRIATLPLLLVRHFEPTIPLQIVDHPLPPLSFTEALQWPLLHNSDP......(300) AmpR: VMVFDSSVTMLEAAQAGVGIAIAPVDMFTHLLASERIVQPFATQIELGSYWLTRLQSRAETPA.....(291) 3 3 3 33 Consensus: A S M LA

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Figure 5. Multiple alignments of the amino acid sequence encoded by ORF bt8 with some LysR family proteins. GcvA, LysR, NodD and AmpR sequences are from E. coli, E. coli, Bradyrhizobia, Enterobacter cloacae respectively. Residue number corresponding to the last residue in the protein shown is presented in parenthesis. Numbers below the alignments represent the number of identical residues at a position where three or more matches are found. Helix-turn-helix residue position (asterisked) was identified based on the consensus motif (LTAAARALHLSQPAISRAIA; Henikoff et al., 1988) for LysR family proteins. Gaps introduced to make the best alignments are indicated with hyphens.

regulator. The GCVA functions as both a positive (by glycine) and negative (by purine, inosine) regulator of the glycine cleavage operon in *E. coli* (Wilson and Stauffer, 1994). It belongs to the LysR family. Transcriptional regulators, NodD and SyrM commonly found in rhizobia belong to the LysR family protein. This protein encoded by ORF bt8 has 291 amino acid residues containing a putative HTH DNA binding motif (Fig. 5). The hypothetical protein encoded by the ORF may therefore be involved in the transcriptional regulation of the genes upstream and/or downstream. Transcriptional regulators are host-range determinants in rhizobia-legume interaction.

The role played by the gcvA homologue located on the pBTBX in the interaction of NGR234(pBTBX) with cv. Peking is possibly similar to that of NodD and SyrM in regulating the biosynthesis of specific Nod factor required for host recognition and efficient nodulation of the soybean. This is substantiated by the fact that nodG (ORF bt9) homologue lies adjacent to the gcvA homologue (ORF bt8)

on the pBTBX. The nodD genes control the first level of host specificity. The role played by nodG has been shown in certain rhizobial strains. Tn5 insertion mutation in nodG gene of R. meliloti resulted to a reduction in the number of nodules and a delay in nodule appearance (Horvath et al., 1986). Cloutier et al. (1997) reported reduced number and size of nodules on Astragalus cicer and Onobrychis vicifolia when inoculated with Tn5 nodG mutants of R. spp. strain N33, a Canadian high arctic rhizobial species. Therefore, ORF bt9, a nodG homologue is proposed to be the synthesis of Nod involved in NGR234(pBTBX); possibly specifically in reduction reaction involved in the metabolism of fatty acids. The bt9 encoded a protein which might endowed NGR234(pBTBX) with the ability to efficiently nodulate cv. Peking relative to the wild-type NGR234 which formed root-like structures. Another transcriptional regulator found on this DNA fragment has homology to GNTR transcriptional repressor of Escherichia coli. This protein

belongs to Lac1 family of transcriptional regulators. It negatively controls the induction of gluconate genes (gntRKU) of the Gnt1 system (Tong et al., 1996).

Methyl-accepting chemotaxis proteins (MCP) similar to the MCPs of *Rhodobacter capsulatus* and *Mycobacterium tuberculosis* are potentially encoded by ORFs bt14 and bt29. The MCPs are involved in motility. Motility confers a selective advantage on rhizobia in competition against nonmotile strains (Ames and Bergman, 1981; Caetano-Anollès et al., 1988). The presence of *mcp* homologues on DNA in addition to the chemoreceptor genes in NGR234 may confer a selective advantage on the NGR234(pBTBX) transconjugant over other strains during its symbiotic interaction with soybean cv. Peking.

Recombinase, replication initiation protein, NADH-**FMN** oxidoreductase (encoded by snaC) phosphoglycerate mutase are encoded by the DNA fragment. Recombinase (Cre) is part of a two components (Lox-Cre) of the bacteriophage P1 site-specific recombination system. It is essential for establishment of prophage and viral vegetative growth in a recA host (Sternberg et al., 1986). Phosphoglycerate mutase reversibly catalyzes the migration of phosphate ester from C3 to C2 in the conversion of 3-phosphoglycerate to 2phosphoglycerate in glycolysis (Brock et al., 1984), a reaction which may contribute to the building up of sugar subunits during Nod factor synthesis important in the early stage of the symbiosis or in ATP generation coupled to the reduction of nitrogen.

This DNA sequence has shown that *R. fredii* USDA257 contains many genes encoding a large amount of theoretical proteins with potential roles assigned to them. This can be used to tackle problems of great importance in biotechnology, particularly where the manipulation of certain proteins may be of specific interest to legume yields. This is of special relevance in the case of *R.* species NGR234 that is promiscus, nodulating and/or fixing nitrogen on many legumes across the world. The use of this rhizobium in the development of inoculant will require the increase in its ability to nodulate and fix nitrogen on many more legumes; hence the significance of this sequencing to harness necessary genes.

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