The Beluga Genome: The Legacy of Aurora and Qila
Identification of *Bradyrhizobium elkanii* Genes Involved in Incompatibility with *Vigna radiata*

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**Abstract:** The establishment of a root nodule symbiosis between a leguminous plant and a rhizobium requires complex molecular interactions between the two partners. Compatible interactions lead to the formation of nitrogen-fixing nodules, however, some legumes exhibit incompatibility with specific rhizobial strains and restrict nodulation by the strains. *Bradyrhizobium elkanii* USDA61 is incompatible with mung bean (*Vigna radiata* cv. KPS1) and soybean cultivars carrying the *Rj4* allele. Here, we explored genetic loci in *B. elkanii* that determine incompatibility with *V. radiata* KPS1. We identified five novel *B. elkanii* genes that contribute to this incompatibility. Four of these genes also control incompatibility with soybean cultivars carrying the *Rj4* allele, suggesting that a common mechanism underlies nodulation restriction in both legumes. The fifth gene encodes a hypothetical protein that contains a *tts* box in its promoter region. The *tts* box is conserved in genes encoding the type III secretion system (T3SS), which is known for its delivery of virulence effectors by pathogenic bacteria. These findings revealed both common and unique genes that are involved in the incompatibility of *B. elkanii* with mung bean and soybean. Of particular interest is the novel T3SS-related gene, which causes incompatibility specifically with mung bean cv. KPS1.

**Keywords:** symbiosis; nodulation; *Bradyrhizobium elkanii*; *Vigna radiata*; *Rj4* soybean

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1. **Introduction**

Symbiotic relationships between leguminous plants and soil bacteria, collectively termed rhizobia, are characterized by the formation of root nodules, which are specialized plant organs that accommodate the rhizobia. In the root nodules, rhizobia fix atmospheric dinitrogen into ammonia, which is utilized by the host legumes. This symbiotic association is of critical importance in agricultural production and the nitrogen cycle. The interactions between legumes and rhizobia exhibit high levels of specificity, which is determined by the exchange of signal molecules. Legume roots excrete flavonoid compounds that interact specifically with the transcriptional regulator NodD in rhizobia [1,2]. The activation of NodD induces the expression of nodulation genes, which are required for the synthesis of lipo-chitooligosaccharides called Nod-factors [1,2]. The Nod factors function as signals for host receptors, which activate signal pathways leading to rhizobial infection and nodule formation [1,2]. In addition, rhizobial factors, such as exopolysaccharides, lipopolysaccharides, and secreted proteins, have also been reported to affect nodulation and host specificity [3,4].

The rhizobium *Bradyrhizobium elkanii* forms symbiotic relationships with a number of legumes, including soybean, mung bean (*Vigna radiata*), and groundnuts (*Arachis hypogaea*). In Brazil, *B. elkanii* is used as an inoculant for soybean cultivation [5]. The formation of symbiotic root nodules in soybean
is controlled by several loci, which are referred to as the *rj* or *Rj* loci [6]. Soybean varieties carrying the dominant *Rj4* allele, such as the cultivars Hill and BARC2, are incapable of nodulation with *B. elkanii* USDA61, but are compatible with *Bradyrhizobium diazoefficiens* USDA110 [7]. These studies revealed that *Rj4* incompatibility is strain-specific and cultivar-specific, and appears to resemble gene-for-gene resistance in plant–pathogen interactions [8]. The recent identification of genes that are encoded by the *Rj2* and *Rj4* genes supports this idea. *Rj2* encodes a member of the Toll-interleukin receptor/nucleotide binding site/leucine-rich repeat (TIR-NBS-LRR) class of proteins [9], while *Rj4* encodes a thaumatin-like protein [10]. Both of these classes of protein are involved in plant resistance to pathogens.

Mung bean is an important leguminous crop in tropical and subtropical regions. Growth and seed production by *V. radiata* are highly dependent on symbiosis with nitrogen-fixing rhizobia [11]. The *V. radiata* cv. KPS1 cultivar is incompatible with *B. elkanii* USDA61, resulting in ineffective nodulation [12]. In a previous study [12], we mutated two *B. elkanii* genes, *rhcC2* and *rhcJ*, which are well-conserved components of the type III secretion system (T3SS). The T3SS is a complex protein transport system that is involved in plant and animal pathogenesis by Gram-negative bacteria, and it is also found in many rhizobia [13–17]. The *B. elkanii* T3SS mutants formed functional nodules on *V. radiata* cv. KPS1 and *Glycine max* cv. Hill plants [12]. These results implicated the T3SS in the control of incompatibility between rhizobia and their plant hosts, however, genetic elements involved in the incompatibility between *B. elkanii* USDA61 and *V. radiata* cv. KPS1 have not been fully elucidated. Here, we identified five novel *B. elkanii* genes that contribute to this incompatibility. Four of these genes also control nodulation restriction in *Rj4* soybean, suggesting a common mechanism that is underlying both incompatible interactions. The fifth gene encodes a T3SS-related protein with a host-specific function in KPS1. These findings shed light on the diverse and complicated mechanisms underlying the incompatible symbiosis, and suggest the involvement of a novel T3SS-related gene in the nodulation restriction of mung bean.

### 2. Materials and Methods

#### 2.1. Bacterial Strains and Growth Conditions

The bacterial strains and plasmids that were used in this study are listed in Table 1. *B. elkanii* strains were grown at 28 °C in arabinose–gluconate medium [18], supplemented with 50 µg mL⁻¹ polymyxin and 200 µg mL⁻¹ kanamycin. *Escherichia coli* strains were grown at 37 °C in Luria-Bertani medium (LB) [19] supplemented with 50 µg mL⁻¹ kanamycin.

<table>
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<th>Reference or Source</th>
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<tr>
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<tr>
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<tr>
<td>pRK2013</td>
<td>Helper plasmid, ColE1 replicon carrying RK2 transfer genes; Km&lt;r, tra</td>
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<td>InnBECorR-intR</td>
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*Pol<r: Polymyxin resistant; Km<r: Kanamycin resistant; Sm<r: Streptomycin resistant; Sp<r: Spectinomycin resistant; Tc<r: Tetracycline resistant; Ap<r: Ampicillin resistant; inn: incompatible nodulation. b United States Department of Agriculture, Beltsville, MD, USA.

2.2. Transposon Mutagenesis of *B. elkanii* USDA61

The transposon mini-Tn5 was introduced into the *B. elkanii* USDA61 strain by conjugation, as described by Krause et al. [23], using *E. coli* strain S17-1 [20] carrying the plasmid pUTKm [21], and *E. coli* HB101 (pRK2013) [22]. Strains containing transposons were selected on arabinose–gluconate agar plates, supplemented with 50 µg mL<sup>-1</sup> polymyxin and 200 µg mL<sup>-1</sup> kanamycin.

2.3. Plant Cultivation, Mutant Screening, and Inoculation Tests

Seeds of *V. radiata* cv. KPS1 and *G. max* cv. BARC2 were surface sterilized and germinated on wet sterile paper, then the seedlings were transferred to sterile vermiculite, using procedures described previously [8]. One day after transplantation, 216 *V. radiata* KPS1 seedlings were inoculated with approximately 36,000 Tn5 insertion strains of *B. elkanii* USDA61 (1 mL of 10<sup>7</sup> cells per mL on each plant). Plants were cultivated in a plant growth cabinet (LPH-410SP; NK Systems Co. Ltd., Osaka, Japan) at 25 °C and 70% humidity under a day/night regimen of 16/8 h. After 30 days of cultivation, nodules that formed on the *V. radiata* KPS1 plants were sampled, and the *B. elkanii* mutants were isolated, as described previously [8]. These isolates were used to re-inoculate *V. radiata* KPS1 plants in order to confirm the nodulation phenotypes. *G. max* cv. BARC2 seedlings were also inoculated with the identified mutants in order to determine if the mutants could form nodules on this *G. max* cultivar. Inoculation tests were performed, as described previously [8].

2.4. Nucleotide Sequence Analysis of Tn5-Flanking Regions

Genomic DNA was isolated from *B. elkanii* strains using the Wizard Genomic DNA Purification Kit (Promega, Madison, WI, USA). Transposon-flanking sequences were amplified using the Y-linker method [24]. The genomic DNA was digested with NlaIII or SpI, and ligated with the Y-linker. PCR was carried out using the GoTaq Green Master Mix (Promega), Y-linker primer and Tn5 primer (Table 1). The thermal cycling conditions were as described previously [8]. The PCR fragments were sequenced.
directly using the Tn5 or Y-linker primers. Homology searches were carried out using BLASTN at the National Center for Biotechnology Information and an in-house database for the B. elkanii USDA61 genome [25]. For the identification of tts box (TCGTCAG[CGT]TTNTCG[TA][CA]AGCTN(8)[TC]A) motif, the 2 kb upstream sequences of each gene were searched with the program “fuzznuc” of the EMBOSS package [26]. The consensus sequences from Rhizobium sp. strain NGR234 [27] and B. diazoefficiens USDA110 [23] were compared to select the tts box motif sequence (data not shown).

In order to confirm that the observed phenotypes of the mutants were caused by transposon insertion and not by a secondary mutation independent of the transposon, innB identified in BE53 was mutated via single-crossover recombination. The internal region of innB was amplified by PCR using the primers InnBXbaI-intF and InnBEcoRI-intR (Table 1). The PCR product was cloned into the vector pSUPSCAKm and was mobilized into B. elkanii USDA61 via conjugation, as described previously [12]. The mutant generated by integration of the plasmids was selected on PSY medium containing polymyxin and kanamycin. Integration of the plasmids was confirmed by PCR using the primer set Psuppol-F and Psuppol-R (Table 1).

2.5. Microscopy

Nodules were excised from V. radiata cv. KPS1 and G. max cv. BARC2 and were fixed, sectioned, and photographed under a stereoscopic microscope, as described previously [8].

2.6. Nucleotide Sequence Accession Numbers

The sequences of the innA, innB, innC, innD, and innE genes have been submitted to GenBank and may be found under accession numbers KX499540, KX499541, KX499542, KX499543, and KX499544, respectively.

3. Results

3.1. Isolation of the Transposon Mutants of B. elkanii

To investigate the molecular mechanisms underlying the incompatibility of V. radiata cv. KPS1 with B. elkanii USDA61, we created about 36,000 Tn5 transposon insertion mutants of USDA61, and then inoculated 216 KPS1 seedlings with the mutants. Tn5 mutants that formed nodules on KPS1 were isolated and used to re-inoculate KPS1 plants to confirm the nodulation phenotypes. Five mutants (BE5, BE53, BE85, BE103, and BE168) were confirmed to form nodules on KPS1 (Figure 1). The wild-type strain USDA61 formed only a small number of nodules on KPS1 plants, and these plants exhibited nitrogen deficiency symptoms (Figure 1A,H). In contrast, all of the Tn5 mutants were able to form mature nodules and promote the growth of KPS1, indicating efficient nitrogen-fixing activity (Figure 1B–F,I–M). The B. elkanii strain BErhcJ, a mutant defective in the type III protein secretion system [12], also formed nodules and promoted growth of KPS1 plants (Figure 1G,N). Nodules formed by all the plants showed a red color in the central tissue due to the presence of leghemoglobin, which is an indicator of nitrogenase activity (Figure 1O–U).
3.2. Symbiotic Phenotypes of the Transposon Mutants on V. radiata cv. KPS1 Plants

To characterize the symbiotic phenotypes of the mutants, we measured the nodule numbers, nodule weights, and plant fresh weights of inoculated KPS1 plants at 40 days after inoculation (Figure 2). For this analysis, we classified the nodules into two types by size: <2 mm and ≥2 mm in diameter. The Tn5 mutants exhibited significantly higher nodule numbers (about 18 to 114 nodules per plant) than the wild-type USDA61 strain (1.8 nodules per plant) (Figure 2A). The number of nodules and the total weight of nodules per plant were highest in plants that were inoculated with BE53 (Figure 2A,B). These plants also showed the highest fresh weights. Interestingly, BE53 induced many small nodules: 80% of total nodules (Figure 2A). In contrast, the mutants BE5, BE85, BE103, and BE168 formed mainly large nodules, and the numbers of large nodules and nodule weights per plant were similar to those of BErhcJ (Figure 2A,B). Apart from plants that were inoculated with BE53, the fresh weights of plants inoculated with the other mutants were similar to one another, and correlated with the numbers of large nodules and total nodule weights. These correlations were probably due to similar nitrogen fixation levels. As we reported previously, BErhcJ formed large nodules on KPS1 plants and promoted plant growth (Figure 2C).

3.3. Symbiotic Phenotypes of the KPS1-Nodulating Mutants on Rj4 Soybean

We also tested the symbiotic phenotypes of the Tn5 mutants on soybean cultivar BARC2 (Rj4/Rj4), which is known to restrict nodulation by B. elkanii [8,12]. Wild-type USDA61 induced only a few nodules on BARC2 plants, and the plants exhibited retarded growth due to nitrogen deficiency (Figure 3A,H). Interestingly, the four mutants BE5, BE85, BE103, and BE168 induced numerous nodules on the BARC2 roots (Figure 3B,I–D–F–K–M). More than 75% of the nodules that were formed by these mutants were large (Figure 4A,B), and they enhanced plant growth (Figure 4C). In contrast, BE53, which formed numerous nodules on V. radiata KPS1, formed only a few nodules on soybean BARC2 (Figure 3C,J). The nodules formed by BE53 were small and irregular in shape, and were similar to those that were formed by USDA61 (Figure 3Q). The BARC2 plants inoculated with BE53 showed nitrogen deficiency symptoms, indicating that the small nodules were ineffective.
Figure 2. Symbiotic properties of *V. radiata* (L.) cv. KPS1 inoculated with *B. elkanii* USDA61, Tn5 mutants and type III secretion system-deficient mutant BErhcJ. (A) Nodule number, (B) nodule weight and (C) plant weight measured at 40 days post inoculation. Open bars and closed bars in (A) show the numbers of small nodules (<2 mm) and large nodules (≥2 mm), respectively. Nodulation tests were performed at least five times, the values are means of seven mung bean plants, and the error bars indicate standard deviations. Statistical analysis (Fisher’s method) was performed comparing the total nodule numbers induced by USDA61, BErhcJ and Tn5 mutants. Means followed by the same letters are not significantly different at 5% level of significance by Fisher’s test. N.D.: Not determined.
Figure 3. *Glycine max* cv. BARC2 (Rj4Rj4) plants, roots, and nodules inoculated with wild-type *B. elkanii* USDA61, Tn5 mutants and type III secretion system-deficient mutant BErhcJ. Plants were photographed at 30 days after inoculation. Scale bars: (A–G) 10 cm, (H–N) 1 cm, and (O–U) 1 mm.

Figure 4. Symbiotic properties of *G. max* cv. BARC2 (Rj4Rj4) inoculated with *B. elkanii* USDA61, Tn5 mutants and type III secretion system-deficient mutant BErhcJ. (A) Nodule number, (B) nodule weight and (C) plant weight measured at 30 days post inoculation. Open bars and closed bars in (A) show the numbers of small nodules (<2 mm) and large nodules (≥2 mm), respectively. Nodulation tests were performed at least two times, the values are means of six soybean plants, and the error bars indicate standard deviations. Statistical analysis (Fisher’s method) was performed comparing the total nodule numbers induced by USDA61, BErhcJ and Tn5 mutants. Means followed by the same letters are not significantly different at 5% level of significance by Fisher’s test. N.D.: Not determined.
 whose homologues were well conserved among rhizobia as well as species of *Afipia*, *Xanthomonas campestris*, *Azospirillum*, and *TerB*. The gene products are similar with those in other rhizobia and pathogenic bacteria. BE103 (GenBank accession No. BAB52639.1), is located in the cluster of genes that are encoding the T3SS [29].

**3.4. *B. elkanii* Genes Responsible for the Symbiotic Incompatibility**

To identify the genes that were disrupted by the Tn5 insertions, Y-linker PCR was performed using genomic DNA from the Tn5 mutants. All of the mutant genomes contain the Tn5 fragment, and the PCR fragment sizes differ among strains, indicating that they are independent mutations. The genome of *B. elkanii* USDA61 has been completely sequenced [25], and the Tn5 flanking regions all contain sequences that are found in the genome (Table 2). The location map and genome context of the identified genes indicated that these genes are located outside T3SS gene cluster (Figures S1 and S2). BE5 contains a Tn5 insertion in a 1212 bp open reading frame (ORF), designated as *innA*, that encodes a cytosine deaminase, which is similar to those in rhizobia and other Gram-negative bacteria. BE85 contains a Tn5 insertion in a 504 bp ORF (designated as *innC*) that encodes a tellurite resistance protein (TerB). The gene products are similar with those in other rhizobia and pathogenic bacteria. BE103 has an insertion in a 1089 bp ORF (*innD*) that encodes an ABC-transporter substrate-binding protein, whose homologues were well conserved among rhizobia as well as species of *Afipia*, *Rhodopseudomonas*, *Azospirillum*, and *Agrobacterium*. One of its homologues in *B. diazoefficiens* USDA110, bsr3743, has been reported as a secreted protein [28]. BE168 has an insertion in a 2292 bp ORF (designated as *innE*) encoding GTP pyrophosphokinase (Table 2). The *innE* product has 88% and 60% identity with the products of *relA* genes in rhizobia and *spoT* genes in other bacteria including *Rhodopseudomonas palustris*, *Xanthomonas campestris*, *Pseudomonas fluorescens*, and *Salmonella enterica*.

BE53 mutant carries a Tn5 insertion in the 2280 bp ORF (*innB*), which encodes a hypothetical protein that is conserved among rhizobia as well as *Xanthomonas* and *Rhodopseudomonas*. Notably, a *tts* box motif (5′-TATGGGACCTAGCTTGCAGAAAAAGCTGACGA-3′) is present at 96 bp upstream of the *innB* ORF. The *tts* box motifs are often located in the promoter regions of genes encoding structural components of the T3SS and secreted proteins. Furthermore, a homologue of *innB* in *Mesorhizobium loti*, mlr6327 (GenBank accession No. BAB572639.1), is located in the cluster of genes that are encoding the T3SS [29].

In order to confirm that the observed phenotypes of the mutants were caused by transposon insertion and not by a secondary mutation independent of the transposon, we reintroduced the mutation into the wild-type background. The *innB* gene was mutated via single-crossover recombination in wildtype USDA61 and the newly constructed mutant, designated BE53S, was used to inoculate *V. radiata* cv. KPS1 plants, and its symbiotic phenotypes were examined. As a result, the BE53S formed numerous mature nodules similar to the transposon mutant (BE53) (Figure 2A–C), confirming that transposon insertion in BE53 was responsible for the altered nodulation phenotype.
Table 2. Results of BLAST and \textit{tts} box searches of Tn5-flanking sequences.

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<th>Designated Symbol/GenBank Accession No.</th>
<th>Length (bp)</th>
<th>Deduced Gene Products</th>
<th>(\text{tts} \text{ Box})</th>
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<td>\textit{innA}/KX499540</td>
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<td>Cytosine deaminase</td>
<td>-</td>
<td>\text{Bradyrhizobium pachyrhizi} AOQ73_15990 (KRQ04415.1, 95%), \text{Afipia} \text{clevelandensis} ATCC 49720 HM4PREF9696, 04200 (EK531979.1, 77%), \text{Agrobacterium rhizogenes} CN09_18315 (KAE04550.1, 61%), \text{Bradyrhizobium diazoefficiens} NK6 Nk6_7958 (BAR10109.1, 59%), \text{Bradyrhizobium sp. BTAi1} BBa7_2704 (ABQ30984.1, 59%), \text{Nitrosospira sp. NaPv SQ1_05400} (KK049569.1, 58%), \text{Mesorhizobium loti} NITP2014 A8146_24615 (OBQ733313.1, 56%), \text{Azorhizobium caulinarum} ORS571 AZC_1945 (BAF587943.1, 49%), \text{Sinorhizobium meliloti} 1021 Smc02420 (CAC47175.1, 40%), \text{Pseudomonas fluorescens} C3 VC34_22545 (KJZ38903.1, 43%)</td>
</tr>
<tr>
<td>BE53</td>
<td>-</td>
<td>\textit{innB}/KX499541</td>
<td>2280</td>
<td>Hypothetical protein</td>
<td>+</td>
<td>\text{Bradyrhizobium yanomami} BR3267 AOQ72_03805 (KRP85897.1, 73%), \text{B. japonicum} USDA6 Bj6t_78540 (BAL13101.1, 71%), \text{B. diazoefficiens} USDA110 Olr1998 (BAR57109.1, 70%), \text{B. japonicum} USDA 6 Bj6t_78530 (BAL13091.1, 66%), \text{B. diazoefficiens} USDA110 b1l1877 (BAR47142.1, 46%), \text{M. loti} MAF303099 mlr6327 (BAB52639.1, 45%)</td>
</tr>
<tr>
<td>BE85</td>
<td>+</td>
<td>\textit{innC}/KX499542</td>
<td>504</td>
<td>TerB family tellurite resistance protein</td>
<td>-</td>
<td>\text{Bradyrhizobium erythrophilum} MT12 SAMN05444164_5326 (SED40490.1, 97%), \text{B. diazoefficiens} NK6 Nk6_3937 (BAR57108.1, 81%), \text{B. japonicum} Is-34 MA20_25200 (KGT77838.1, 80%), \text{B. japonicum} USDA6 Bj6t_05250 (BAL05822.1, 79%), \text{A. clevelandensis} ATCC 49720 HM4PREF9696_03100 (EK533980.1, 77%), \text{Proteobacteria bacterium} 5C_2bin9 A4S14_11410 (OQW55616.1, 114%), \text{Rhodopseudomonas sp. AAP120} IP86_13420 (KPF97560.1, 73%), \text{Nitrobacter hamburgensis} X14 Nhar_0005 (ABE61949.1, 72%)</td>
</tr>
<tr>
<td>BE103</td>
<td>+</td>
<td>\textit{innD}/KX499543</td>
<td>1089</td>
<td>ABC transporter substrate-binding protein</td>
<td>-</td>
<td>\text{B. pachyrhizi} BR3262 AOQ73_18465 (KRQ01332.1, 99%), \text{B. diazoefficiens} USDA110 bhr7816 (BAR53081.1, 87%), \text{Afipia sp. P52-10} X566_18245 (EET47616.1, 62%), \text{Rhodopseudomonas palustris} BisB5 RPD_2049 (ABE39284.1, 61%), \text{Azospirillum brasilense} Sp7 AMK58_19775 (ALJ36795.1, 61%), \text{B. japonicum} USDA6 Bj6t_61480 (BAL11399.1, 60%), \text{Bosea vacuolata} SD260 AEs18_12875 (KPH80033.1, 61%), \text{Agrobacterium tumefaciens} KCJ17 A7J57_24755 (OAE45812.1, 59%), \text{B. diazoefficiens} USDA110 bhr3743 (BAC49008.1, 59%)</td>
</tr>
<tr>
<td>BE168</td>
<td>+</td>
<td>\textit{innE}/KX499544</td>
<td>2292</td>
<td>GTP pyrophosphokinase</td>
<td>-</td>
<td>\text{B. pachyrhizi} BR3262 AOQ73_36790 (KRQ06030.1, 99%), \text{Bradyrhizobium jacunae} PAC68 CQ12_18875 (KRQ98944.1, 94%), \text{B. diazoefficiens} USDA110 blr5069 (BAC50300.1, 88%), \text{B. japonicum} USDA 6 Bj6t_46480 (BAL09914.1, 87%), \text{Rhodopseudomonas palustris} BisB5 RPD_2049 (ABE39284.1, 61%), \text{Azospirillum brasilense} Sp7 AMK58_17855 (ALJ37338.1, 52%), \text{Frankia alni} ACN14a FRAL2148 (CAJ60797.1, 40%)</td>
</tr>
</tbody>
</table>

\(a\) Nodulation on \textit{G. max} cv. BACR2 (Rj4/Rj4) soybean; +: Positive; −: Negative. \(b\) \textit{inn}: incompatible nodulation. \(c\) Presence of \textit{tts} box in upstream regions of the genes; +: Present; −: Absent.
4. Discussion

In fields, the application of high nitrogen-fixing rhizobia to their target legume crops often fails to increase the crop yields. One of the major causes of this phenomenon is that more competitive indigenous rhizobial strains, with low nitrogen fixation activity, occupy most of the nodules. This problem is known as the competition problem, and it is very important in the cultivation of legumes. Besides the competitiveness of different rhizobial strains, compatibility between leguminous plants and rhizobia also affect nodule occupancy. In soybean, Rj genes control nodulation: plants carrying specific Rj genes cannot form nodules when inoculated with specific rhizobial strains [6]. This compatibility/incompatibility might be reminiscent of host–pathogen interactions. Consistent with this observation, the Rj2 and Rj4 genes encode TIR-NBS-LRR and thaumatin-like proteins, respectively [10], and these proteins are involved in plant resistance to pathogens.

Similarly, host-controlled nodulation occurs in mung beans, and the compatibility is dependent on both the host cultivar and the rhizobial strain. In a previous report, we showed that two genes that encode components of the T3SS, rhcC2 and rchJ, are necessary for the incompatibility between V. radiata cv. KPS1 and B. elkanii USDA61 [12]. The T3SS is involved in plant and animal pathogenesis, and many rhizobial strains possess a functional T3SS [13–17]. The secreted proteins, called nodulation outer proteins (Nops), affect symbiosis positively or negatively, depending on the host plant [23,30,31]. The expression of the rhizobial T3SS is controlled by flavonoids that are derived from the host legume. Flavonoids perceived by the NodD protein activate the expression of ttsI, a transcriptional activator of the type III secretion gene cluster (tts). TtsI subsequently activates the ttsI gene cluster, and proteins are secreted via the T3SS. A conserved motif, called the tts box, is located in the promoter region of genes encoding structural components and secreted effector proteins [23,32,33].

In the present study, we explored additional genetic determinants of B. elkanii, which are involved in the nodulation incompatibility between V. radiata KPS1 and B. elkanii USDA61. We performed Tn5 transposon mutagenesis and identified five Tn5 insertion mutants of B. elkanii USDA61, which were able to form functional nodules on V. radiata KPS1 and enhance the growth of the plants.

The BE5 mutant contains a Tn5 insertion in the innA gene, which encodes cytosine deaminase. Cytosine deaminase deaminates cytosine to form uracil and ammonia [34]. Although the mechanism for the involvement of cytosine deaminase in nodulation restriction remains unclear, the BE5 mutant efficiently nodulated and promoted the growth of both KPS1 and Rj4 plants. The disruption of innA might cause the inactivation of the T3SS, which abolishes the secretion of effector proteins that are responsible for nodulation restriction in both Rj4 soybeans and V. radiata KPS1.

The BE53 mutant carries transposon insertion in the innB gene, which encodes a hypothetical protein with an unknown function. The innB gene is preceded by the consensus tts box, and the innB homologue in M. loti, mrl6327, occurs in the cluster of genes encoding the T3SS [28]. Intriguingly, the innB-deficient mutant formed nodules on V. radiata KPS1 plants, but not on Rj4 soybean. This suggests that innB encodes a protein with a host-specific function in KPS1. One possibility is that innB encodes an effector protein that is recognized specifically in V. radiata KPS1. It is likely that the rhizobial effector proteins that are recognized by KPS1 and Rj4 soybean differ. The InnB might be recognized specifically by a resistance (R) protein in KPS1, causing nodulation restriction in KPS1, but not in Rj4 soybean.

The BE85 mutant contains a Tn5 insertion in innC, encoding TerB family tellurite resistance protein that is conserved among the Bradyrhizobiaceae. It has been reported that the Tellurium resistance proteins of Frankia spp. are induced by seed phenolic compounds that are produced by their actinorhizal host plants [35]. Considering that the disruption of innC induced nodulation on both KPS1 and Rj4 plants, the gene might be involved in the T3SS. However, the function of these proteins in rhizobial symbiosis remains unknown.

The BE103 mutant carries a Tn5 insertion in the gene innD, which encodes a substrate-binding protein in the ABC transporter family. These substrate-binding proteins are located in the periplasm, and are involved in the active transport of substrates into the cytoplasm [36–38]. Although the function of the innD protein remains unclear, its homologue in B. diazoefficiens USDA110 (blr3743, BAC49008.1)
is secreted [28], and is upregulated in bacteroids in nodules [39,40]. Clearly, functional analyses are needed to determine the contribution of these transporters to the compatibility/incompatibility between rhizobia and their hosts. Since the inactivation of the innD gene by the Tn5 insertion canceled the nodulation restriction in both KPS1 and Rj4 plants, the innD product might be involved in the T3SS. Further molecular experiments will be performed to explore this possibility.

The BE168 mutant contains a Tn5 insertion in innE, which encodes a GTP pyrophosphokinase (ppGpp synthetase) that is conserved among various rhizobia and pathogenic bacteria. innE shares similarity with the relA genes of B. diazoefficiens USDA110 (BAC50330.1) and other bacteria. It has been proposed that ppGpp synthetase binds to, and alters the ability of, RNA polymerase to initiate and elongate transcription. ppGpp synthetase activates genes that function in the biosynthesis of amino acids and transcription factors that are responsible for stress responses [41,42]. In addition, ppGpp synthetase plays important roles in the virulence and the persistence of phytopathogens within host plants [43]. In Rhizobium elli, the relA gene regulates gene expression in bacteroid cells [44]. A relA mutant of Sinorhizobium meliloti shows a non-nodulation phenotype on alfalfa [45]. These findings suggest that the mutation in innE affects cellular ppGpp synthetase levels and leads to changes in cell functions that affect the strain specific incompatibility between B. elkanii USDA61 and mung bean and soybean. It should be noted that, except for the innB gene, which was confirmed by the newly constructed mutant BE53S, the other genes also need to be confirmed by mutational and/or complementation analysis to unequivocally associate the gene with the phenotype.

In addition to the genes and gene products discussed above, rhizobial factors have been reported to be involved in incompatible symbioses, including the type IV secretion system [46], extracellular polysaccharides [47], and peptidase that cleaves host-derived signaling peptides [48]. T3SSs in other rhizobia involved in incompatible symbioses have been reported for the following interactions: Rhizobium sp. NGR234 with Crotalaria juncea [49], M. loti MAFF303099 with Lotus halophilus [28], Sinorhizobium fredii USDA257 with G. max cv. McCall [15], B. elkanii USDA61 with Rj4 soybean [8], Bradyrhizobium japonicum USDA122 with Rj2 soybean [50], and B. diazoefficiens USDA110 with V. radiata KPS2 [51]. The type III-secreted proteins NopE1 and NopE2 of B. diazoefficiens USDA110 were reported as negative effectors of compatibility with V. radiata KPS2 [51]. In our preliminary results, B. diazoefficiens USDA110 formed nodules on V. radiata KPS1 and promoted plant growth efficiently (data not shown), indicating that NopE1 and NopE2 do not control incompatibility with KPS1, and that B. elkanii might possess other effectors that are recognized by KPS1. These findings suggest that the molecular mechanisms that are controlling incompatibility between various rhizobia and legumes differ, although the T3SS appears to underlie at least three types of nodulation restriction. Clearly, further study is necessary to define the roles of the T3SS components in both innate-immunity and rhizobial nodulation.

5. Concluding Remarks

In conclusion, we have identified novel genes of B. elkanii that are involved in incompatibility with V. radiata cv. KPS1. Four of the genes (innA, innC, innD, and innE) are also involved in incompatibility with Rj4 soybean, suggesting that they participate in a common mechanism underlying nodulation restriction in both of the legumes. In contrast, innB is involved in nodulation restriction with V. radiata cv. KPS1, but not with Rj4 soybean. The innB gene is preceded by a tts box, which is a promoter element in type III secretion system-related genes, suggesting its T3SS-related function. These findings shed light on the diverse and complicated mechanisms underlying the incompatible symbiosis, and suggest the involvement of a novel T3SS-related gene in the nodulation restriction of mung bean.

Supplementary Materials: The following are available online at www.mdpi.com/2073-4425/8/12/374/s1. Figure S1: The location map of the identified genes innA, innB, innC, innD, and innE in B. elkanii USDA61 genome, Figure S2: Genome context of the identified genes innA, innB, innC, innD, and innE.

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Conflicts of Interest: The authors declare no conflict of interest.

References


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