The title:
Genetic diversity and geographical distribution of indigenous soybean-nodulating bradyrhizobia in
the United States

The running title:
Genetic diversity of American bradyrhizobia

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ABSTRACT

We investigated the relationship between the genetic diversity of indigenous soybean-nodulating bradyrhizobia and their geographical distribution in the United States using nine soil isolates from eight states. The bradyrhizobia were inoculated on three soybean $R_j$ genotypes (non-$R_j$, $R_j2R_j$, and $R_j4$). We analyzed their genetic diversity and community structure by means of restriction-fragment length polymorphisms of PCR amplicon to target the 16S–23S rRNA gene internal transcribed spacer region, using 11 USDA *Bradyrhizobium* strains as reference strains. We also performed diversity analysis, multi-dimensional scaling analysis based on the Bray-Curtis index, and polar ordination analysis to describe the structure and geographical distribution of the soybean-nodulating bradyrhizobial community. The major clusters were *Bradyrhizobium japonicum* Bj123, in the northern U.S., and *Bradyrhizobium elkanii* in the middle to southern regions. Dominance of bradyrhizobia in a community was generally larger for the cluster belonging to *B. elkanii* than for the cluster belonging to *B. japonicum*. The indigenous American soybean-nodulating bradyrhizobial community structure was strongly correlated with latitude. Our results suggest that this community varies geographically.

Keywords:

- genetic diversity
- geographical distribution
- soybean $R_j$-genotype
- multi-dimensional scaling
- polar ordination
INTRODUCTION

The United States is the world’s largest producer and consumer of soybeans (*Glycine max.* L. Merr.), which are utilized for food products, feedstuff and biofuel feedstock. Soybean is a legume that forms root nodules after infection with soybean-nodulating rhizobia, which perform symbiotic nitrogen fixation by taking up atmospheric nitrogen (as ammonia) through the root nodules. The major soybean-nodulating rhizobia are *Bradyrhizobium japonicum*, *Bradyrhizobium elkanii*, and *Sinorhizobium (Ensifer) fredii* (5, 12, 16, 34, 51). In addition, *Bradyrhizobium yuanmingense*, *Bradyrhizobium liaoningense*, *Sinorhizobium xinjiangense*, and *Mesorhizobium tianshanense* have been classified as soybean-nodulating rhizobia (4, 24, 38, 45, 49, 50).

Inoculation of soybean with bradyrhizobia can improve nitrogen fixation, resulting in increased soybean yield. However, the efficiency of the inoculum may be poor if the inoculum cannot compete with indigenous soybean-nodulating rhizobia in the soil or cannot establish an efficient symbiosis with the host plants. To solve this problem, it will be necessary to understand the ecology of indigenous soybean-nodulating rhizobia in terms of their genetic diversity, geographical distribution, compatibility with the host soybean, and the environmental factors associated with the localization and dominance of the rhizobial strains in the soil.

Saeki et al. (28) investigated the genetic diversity and geographical distribution of indigenous soybean-nodulating rhizobia isolates from five sites in Japan (Hokkaido, Fukushima, Kyoto, Miyazaki, and Okinawa) by analyzing polymerase chain reaction restriction-fragment length polymorphisms (PCR-RFLP) of the 16S–23S rRNA gene internal transcribed spacer (ITS) region. They reported that a geographical distribution of indigenous bradyrhizobia varied from northern to
southern Japan. Furthermore, Saeki et al. (31) reported that the distribution of soybean-nodulating rhizobia in Japan was strongly correlated with latitude ($r^2 = 0.924$). The representative clusters of the isolated bradyrhizobia changed from those of $B. japonicum$ strains USDA 123, 110, and $6^T$ to $B. elkanii$ strain USDA $76^T$ moving from northern to southern Japan (26). Adhikari et al. (1) revealed the genetic diversity of soybean-nodulating bradyrhizobia in relation to climate depending on altitude and soil properties such as soil pH in Nepal. Other researchers found that $S. fredii$ strains were dominant in the alkaline soils of Vietnam and Okinawa, Japan (30, 37). These results suggest that a relationship exists between the geographic distribution of indigenous soybean-nodulating rhizobia, soil temperature (and its variations due to latitude and altitude), and soil pH. The ability of a given soybean accession to bradyrhizobia and form nodules depends on the characteristics of nodulation regulatory genes ($Rj$ genes), and $Rj$ genotypes of $rj_1$, $Rj_2$, $Rj_3$, $Rj_4$ and non-$Rj$ which lack these genetical phenotypes have been confirmed to exist in nature (3, 42, 43, 48). Previous experimental results have also demonstrated that the community structure of soybean-nodulating bradyrhizobia depends on the host soybean $Rj$-genotype and on the soybean cultivar, and varies with cultivation temperature even in an identical soil sample (20, 36). The United States is the world’s biggest soybean producer, and soybean cultivars are grown at latitudes similar to those of the soybean production areas in Japan. Understanding the geographical distribution of soybean-nodulating rhizobia in the United States would therefore provide important knowledge about bradyrhizobial ecology and insights into appropriate inoculation techniques for soybean-nodulating rhizobia with high nitrogen-fixation ability. Thus, in this study, we examined the relationship between the genetic diversity and geographical distribution of indigenous soybean-nodulating bradyrhizobia by isolating these organisms using three $Rj$ genotypes soybean.
cultivars from nine soil samples of American soybean fields. Our objectives were to investigate the genetic diversity, community structure, and geographic distribution of the bradyrhizobia by means of PCR-RFLP analysis of the 16S–23S rRNA gene ITS region. In addition, we described the community structure and geographical distribution of the bradyrhizobia using several tools from mathematical ecology.

MATERIALS AND METHODS

Soil samples

We obtained soil samples for isolation of soybean-nodulating bradyrhizobia from nine experimental fields and farm fields in eight American states (Michigan, Ohio, Kentucky, North Carolina, Alabama, Georgia, Florida, and Louisiana) in August 2010 (Table 1). Three samples (each at least 200 cm³) were obtained in each field, to a depth of 10 cm after removing the surface litter. The samples were homogenized to produce a single composite sample. The Alabama soil sample was collected from two separate soybean fields, which we designated Alabama1 and Alabama2. Table 1 summarizes the location, soil pH, electrical conductivity (EC), total carbon (C) and total nitrogen (N) contents, and C/N ratio at these sites.

Isolation of the indigenous bradyrhizobia

To isolate indigenous soybean-nodulating bradyrhizobia, we used three soybean cultivars of three Rj genotypes, Bragg as non-\(R_j\), CNS as \(R_jR_j\), and Hill as \(R_j\), and planted each soybean cultivar in 1-L culture pots (\(n = 3\) plants per cultivar). The culture pots were filled with vermiculite containing N-free nutrient solution (29) at 40% (v/v) water content, and were then autoclaved at 121°C for 20 min. Soybean seeds were sterilized by soaking them in 70% ethanol for 30 s and in a
dilute sodium hypochlorite solution (0.25% available chlorine) for 3 min. They were then washed with sterile distilled water. A soil sample (2 to 3 g) was placed in the vermiculite at a depth of 2 to 3 cm, and the soybean seeds were sown on the soil and then the pot weight was measured. The plants were grown for 4 weeks in a growth chamber (day, 28˚C for 16 h; night, 23˚C for 8 h) with a weekly supply of sterile distilled water until the measured pot weight.

After 4 weeks, 23 to 24 nodules were randomly collected from the soybean roots and sterilized by soaking them in 70% ethanol for 3 min and in a diluted sodium hypochlorite solution (0.25% available chlorine) for 30 min; they were then washed with sterile distilled water. Each nodule was homogenized in sterile distilled water, and then streaked onto a yeast extract–mannitol agar (YMA; 44) plate medium, and incubated for 5 to 7 days in the dark at 28˚C. To determine the genus of the isolates, a single colony was streaked onto YMA plate medium containing 0.002% (w/w) bromothymol blue (13) and incubated as described above. After incubation, each isolate was maintained on YMA slant medium at 4˚C for further analysis. A total of 69 to 72 isolates per soil sample were used to represent the soybean-nodulating bradyrhizobial community; we obtained a total of 645 isolates from the nine soybean fields, and used them in the diversity analysis and multi-dimensional scaling (MDS) analysis described later in the Methods section.

Representative isolates in each OTUs of dendrogram were confirmed their nodulation capability on host soybean by inoculation test. Each isolate was cultured in yeast extract–mannitol broth culture (42) for 6 days at 28˚C, and the cultures were then diluted with sterile distilled water to approximately 10^6 cells mL^-1. The soybean seeds were sown into 500-mL prepared culture pots without soil, as described above, and inoculated with a 1-mL aliquot of each isolate per seed with two or three replicates. We assessed nodule formation after 3 weeks in a growth chamber under the
conditions described above.

**PCR-RFLP analysis of the 16S–23S rRNA gene ITS region**

For DNA extraction, we cultured each isolate in 1.5 mL of HEPES-MES medium (6) supplemented with 0.1% L-arabinose (33) for 5 days at 28°C. Total DNA was extracted from the isolates using BL extraction buffer, as described previously (20) based on the method reported by Hiraishi et al. (11). As reference strains, we used *B. japonicum* USDA strains 4, 6ᵀ, 38, 110, 115, 123, 124, and 135, and *B. elkanii* USDA strains 46, 76ᵀ, and 94 (27). Total DNAs of the reference strains were extracted by means of the same procedure used for the isolates.

The PCR amplification for the 16S–23S rRNA gene ITS region was carried out using *Ex Taq* DNA polymerase (TaKaRa Bio, Otsu, Shiga, Japan) and a previously developed ITS primer set (BraITS-F, 5’-GACTGGGGTGAAGTCGTAAC-3’; BraITS-R, 5’-ACGTCCTTCATCGCCTC-3’; 28). The PCR cycle consisted of a pre-run at 94°C for 5 min, followed by 30 cycles of denaturation at 94°C for 30 s followed by annealing at 55°C for 30 s and extension at 72°C for 1 min, with a final post-run extension at 72°C for 10 min.

The restriction-fragment length polymorphism (RFLP) analysis of the ITS region was investigated using the restriction enzymes *Hae*Ⅲ, *Hha*I, *Msp*I, and *Xsp*I (TaKaRa Bio). A 5-µL aliquot of the PCR product was digested with the restriction enzymes at 37°C for 16 h in a 20-µL reaction mixture. The restriction fragments were separated on 3% or 4% agarose gels by means of electrophoresis and were visualized with ethidium bromide.

**Cluster analysis of the indigenous soybean-nodulating rhizobia**

The fragment sizes on the electrophoresis gels were measured using a 50-bp reference-ladder marker (TaKaRa Bio) and the fragment sizes from the sequences of the reference strains. All
reproducible fragments longer than 50 bp were used for the cluster analysis (described later in this section), and some irreproducible fragments were excluded. The genetic distance between pairs of isolates \( D \) was determined using the following equation:

\[
D_{AB} = 1 - \left[ \frac{2N_{AB}}{N_A + N_B} \right]
\]

where \( N_{AB} \) represents the number of RFLP bands shared by strains A and B, and \( N_A \) and \( N_B \) represent the numbers of RFLP bands found only in strains A and B, respectively (22, 32). The cluster analysis was carried out using the unweighted pair group method with an arithmetic average (UPGMA) method. The dendrograms were constructed using version 3.69 of the PHYLIP software (J. Felsenstein, University of Washington, Seattle, WA; http://evolution.genetics.washington.edu/phylip.html).

**Diversity analysis for the bradyrhizobial communities**

To estimate the diversity of the bradyrhizobial communities in the United States that we isolated from the host soybeans, we used the Shannon-Wiener diversity index (17, 25, 31):

\[
H' = -\sum P_i \ln P_i
\]

where \( P_i \) is the dominance of dendrogram cluster \( i \), defined as \( n_i/N \), where \( N \) is the total number of isolates \( (n = 69 \) or \( 72 \)) and \( n_i \) is the total number of tested isolates belonging to dendrogram cluster \( i \). We also calculated the alpha diversity \( (H'_{\alpha}) \), beta diversity \( (H'_{\beta}) \), and gamma diversity \( (H'_{\gamma}) \) to estimate the differences in the bradyrhizobial communities between pairs of soil samples (15, 47).
The $H'_\alpha$ index represents a weighted average of the diversity indices of two bradyrhizobial communities, the $H'_\beta$ index represents the differences between the bradyrhizobial communities from two soil samples (i.e., differences between sites), and the $H'_\gamma$ index represents the diversity of the total isolate communities from the two soil samples ($n = 141$ or 144). The relationship among these indices is:

$$H'_\beta = H'_\gamma - H'_\alpha$$

We also estimated the differences among the compositions of the bradyrhizobial communities by comparing the ratio of beta to gamma diversity ($H'_\beta/H'_\gamma$), taking into consideration the difference in gamma diversity in each pairwise comparison of bradyrhizobial communities.

**Multi-dimensional scaling analysis, cluster analysis, and polar ordination analysis**

To describe the characteristics of the bradyrhizobial communities and their differences among the field sample sites, we performed an MDS analysis based on the Bray-Curtis similarity measure. The Bray-Curtis similarity measure has a robust monotonic relationship with ecological distance, and a robust linear relationship with ecological distance until large values of the distance. Thus, the Bray-Curtis similarity measure is one of the indices that best reflects the properties between communities (9). The Bray-Curtis similarity measure ($BC$) was calculated using the following equation:

$$BC_{AB} = \sum |n_A - n_B| / \sum (n_A + n_B)$$
where $BC_{AB}$ is the dissimilarity between communities A and B, and $n_A$ and $n_B$ represent the total number of strains in a particular cluster for communities A and B, respectively (2, 19). The three-dimensional MDS analysis and the UPGMA analysis based on the Bray-Curtis similarity measure were conducted using version 2.15.1 of the R software (http://www.r-project.org/).

Furthermore, UPGMA analysis was conducted as described above for all communities as OTUs with the Bray-Curtis similarity to obtain the objective index for the clustering of MDS plots.

To determine the relative distances among the diversities of the bradyrhizobial communities based on the three-dimensional MDS plots as a function of latitude (°N), we calculated the Euclidean distances between the bradyrhizobial communities. The distances between the MDS plot were calculated using the coordinates on the $x$-, $y$-, and $z$-axes as the Euclidean distance ($Ed$) using the following equation:

$Ed_{AB} = (|X_A - X_B|^2 + |Y_A - Y_B|^2 + |Z_A - Z_B|^2)^{1/2}$

where $Ed_{AB}$ is the linear distance between communities A and B in the MDS plot, and $X_A$ and $X_B$, $Y_A$ and $Y_B$, and $Z_A$ and $Z_B$ represent the $x$, $y$, and $z$ coordinates of communities A and B, respectively.

The distances from each pole were converted into percentage differences, $D_1$ and $D_2$, from the two polar communities (i.e., the Michigan and Louisiana sites, which were considered to have a 100% difference).

Polar ordination was conducted to determine the coordinates of communities on the axis between communities of the Michigan and Louisiana sites using Pythagorean theorem (15, 31, 46).

We constructed simultaneous equations from the trigonometric figure using the Pythagorean theorem.
Theorem, as described previously (31). Parameter $P_d$ represents the polar difference (%) from the 0% pole (the bradyrhizobial community of Louisiana), and is calculated as follows (46):

$$P_d = \frac{(L^2 + D_1^2 - D_2^2)}{2L}$$

where $D_1$ and $D_2$ are the percentage differences between a particular bradyrhizobial community and the communities at Louisiana and Michigan, respectively. The parameter $L$ represents the total length of the polar axis (i.e., 100%).

RESULTS

Isolation of indigenous soybean-nodulating bradyrhizobia

We obtained 23 or 24 isolates from each $R_j$ soybean genotype, and a total of 645 indigenous soybean-nodulating bradyrhizobia that could be used for further analysis. The indigenous bradyrhizobia isolated from each soil sample plus host soybean cultivar combination were labeled using a combination of the site abbreviation, followed by an abbreviation for the three cultivars (B, Bragg; C, CNS; and H, Hill) and the number of the isolate (1–23 or 1–24) (e.g., Michigan (MI), MIB1–23, MIC1–23, and MIH1–23). The YMA cultures of all isolates turned blue as a result of the presence of bromothymol blue, indicating that all isolates belonged to the genus *Bradyrhizobium* (12). Representative isolates indicated their nodulation capability on the host soybean by the inoculation test.

PCR-RFLP analysis of the 16S–23S rRNA gene ITS region

PCR products of the amplified 16S–23S rRNA gene ITS region of isolates were digested using
Figure 1 provides a schematic representation of the restriction-fragment patterns for each enzyme. The dendrogram was constructed based on the differences in fragment size and pattern shown in Figure 1. Figure 2 presents the results of the cluster analysis based on PCR-RFLP data. We detected a total of 24 operational taxonomic units (OTUs) containing 11 reference strains (Fig. 2). The maximum similarity between the OTUs and the reference strains was 90%, and occurred between OTUs USDA 76 and USDA 94. These results were then applied as the criterion for distinguishing clusters in the dendrogram, which produced 12 clusters, 11 of which included one of the reference strains. All indigenous bradyrhizobia were classified into 9 clusters: Bj6, Bj38, Bj110, Bj123, Bj124, Be46, Be76, Be94, and BeOH (Fig. 2). Three of the clusters included only a single reference strain and no indigenous bradyrhizobia. Bj6, Bj38, Bj110, Bj123, and Bj124 showed identical or similar RFLP patterns to those of *B. japonicum* strains USDA 6T, 38, 110, 115, 123, and 124, respectively. Be46, Be76, and Be94 showed identical or similar RFLP patterns to *B. elkanii* strains USDA 46, 76T, and 94, respectively. Cluster BeOH consisted of isolates from five sites, was independent from those of the reference strains.

Figure 3 shows the geographical distribution of the soybean-nodulating bradyrhizobial isolates that belonged to each cluster. The isolates belonging to Bj123 were isolated from northern regions (Michigan and Ohio), whereas those belonging to Bj6 were found in the middle regions (Kentucky, North Carolina, and Alabama), and those belonging to Be46 were found in southern regions (Georgia, Florida, and Louisiana). Furthermore, the southern sites were dominated by *B. elkanii* isolates belonging to clusters Be46, Be76, and Be94. The dominance of *B. elkanii* exceeded the dominance of *B. japonicum* in the middle to southern regions (Fig. 2 and 3).
Diversity analysis for the indigenous bradyrhizobial communities

The differences in the indigenous bradyrhizobial communities among the nine sample sites were estimated based on the $H'/H_\gamma$ ratio. The diversity index ($H'$) values were indicated in the bottom of Figure 2, and the beta diversity ratio ($H'/H_\gamma$) at each field site were shown in Figure 4, respectively. As shown in Figure 2, in the indigenous American bradyrhizobial communities, Alabama2 had the highest $H'$ value (1.56), whereas Michigan had the lowest diversity index (0.25). The values of $H'/H_\gamma$ were largest for comparisons of the northern regions (Michigan and Ohio) with the other field sites (Fig. 4).

Multi-dimensional scaling analysis, UPGMA analysis, and polar ordination analysis

Figure 5 shows the results of the MDS analysis. The MDS plots showed three groups of bradyrhizobial communities. Based on UPGMA analysis, group I comprised Michigan and Ohio, group II-1 comprised North Carolina, Alabama2, Georgia, and Louisiana and group II-2 comprised Kentucky, Alabama1, and Florida. Based on the Figure 3, the dominant region of the $B. japonicum$ cluster belonged to group I, the region with high diversity in the $B. elkanii$ cluster belonged to group II-1, and the dominant region for the Be76 cluster belonged to group II-2.

Figure 6 shows the results of the polar ordination analysis based on the percentage differences of the bradyrhizobial communities from each pole and the differences in the latitudes of the sample sites. The results of the polar ordination indicate that the transition in the composition of the indigenous soybean-nodulating bradyrhizobial communities was strongly and significantly related to latitude ($r^2 = 0.815, F = 30.9, p < 0.001$).
DISCUSSION

The American soils from which we isolated soybean-nodulating bradyrhizobia in the present study were acidic to slightly alkaline (pH 5.18 to 7.68, Table 1), and the cluster analysis based on PCR-RFLP data shows that strains of both *B. japonicum* and *B. elkanii* were isolated from nine field soils in the USA (Fig. 2). The major clusters from the sample soils were Bj123 in the northern region, versus Be46, Be76, and Be94 in the middle to southern regions (Fig. 3). Bj6 and Bj110 were moderately dominant in the middle regions. On the whole, the number of clusters of *B. elkanii* strains was larger than the number of clusters of *B. japonicum* strains (Fig. 2 and 3). Cluster Be76 was especially dominant in most sample soils in the middle to southern regions. Keyser *et al.* (14) investigated the distribution of *Bradyrhizobium* serogroups, which were determined using rabbit antisera prepared against 15 serotype strains, in 12 states (Arkansas, Delaware, Florida, Kansas, Louisiana, Minnesota, Mississippi, New Jersey, North Carolina, Pennsylvania, South Carolina, and South Dakota) and found that the dominant serogroup was serogroup 31, which belongs to *B. elkanii* USDA 31 (21.5%; percentage of total isolate), followed by serogroup 123, which belongs to *B. japonicum* USDA 123 (13.6%), and serogroup 76, which belongs to *B. elkanii* USDA 76\(^T\) (10.2%). Interestingly, this report also indicated that *B. elkanii* was more dominant than *B. japonicum*, which confirms the results of the PCR-RFLP analysis of the 16S–23S rRNA gene ITS region in the present study; in addition, the distribution of the major serogroups generally agreed with the distribution of the major clusters in the present study. Additionally, Fuhrmann (10) investigated the diversity and symbiotic effectiveness of indigenous soybean-nodulating bradyrhizobia isolated from 18 locations in Delaware using serological, morphological, rhizobitoxine and hydrogenase phenotypes, and revealed that the dominant serogroup of indigenous
bradyrhizobia was serogroup 94, which belongs to *B. elkanii* USDA 94 (17.5%; percentage of total isolate), followed by serogroup 6, which belongs to *B. japonicum* USDA 6\(^T\) (10.3%), serogroup 122, which belongs to *B. japonicum* USDA 122 (8.6%), and serogroup 76, which belongs to *B. elkanii* USDA 76\(^T\) (5.8%). On the other hand, grouping of the *Bradyrhizobium* USDA strains by sequence analysis and PCR-RFLP targeted to the 16S–23S rRNA gene ITS region used in this study revealed that *B. japonicum* USDA 110 and USDA 122, and *B. elkanii* USDA 31 and USDA 76\(^T\) belong to the same clusters (27, 41). Therefore, to distinguish between *B. japonicum* USDA 110 and USDA 122, and between *B. elkanii* USDA 31 and USDA 76\(^T\), a characterization method based on gene markers such as those for housekeeping genes and/or single nucleotide polymorphism of ITS region will be necessary, because some of the isolates belonging to the Bj110 and Be76 cluster in the present study may be included in the serogroup of the *B. japonicum* USDA 122 and the *B. elkanii* USDA 31.

Our diversity analysis (Fig. 2) revealed that the sampling site with the highest diversity index (\(H^\prime\)) was Alabama2 (1.56), and that Michigan had the lowest diversity index (0.25). The value of \(H^\prime_\beta/H^\prime_\gamma\) was largest for the comparison of the two northern regions (Michigan and Ohio) with the other regions (Fig. 4). This indicates that the bradyrhizobial communities of soybean-nodulating bradyrhizobia differed greatly between the northern region and the other regions (Table 2 and Fig. 4). The Bj123 cluster was only dominant in the northern region (Michigan and Ohio), and the BeOH cluster that was dominant in Ohio may have affected the difference among the communities.

The results of the geographical distribution analysis using polar ordination (Fig. 6) show a strong correlation between the latitude of the samples and the community structure (\(r^2 = 0.815\)). However, the correlation between the community structure in Japan and latitude was higher (\(r^2 = 0.815\)).
The major clusters of indigenous soybean-nodulating bradyrhizobia in Japan were regularly distributed, with clusters in the order of *B. japonicum* strains USDA 123, USDA 110, and USDA 6T and *B. elkanii* strain USDA 76T moving from northern to southern regions, and was generally dominated by *B. japonicum* strains (28). On the other hand, in the geographical distribution of the American soybean-nodulating bradyrhizobia, the *B. elkanii* clusters were more dominant than the *B. japonicum* clusters (Fig. 3), and the dominance of the Bj110 cluster in the middle regions was lower than in Japan. Furthermore, the American and Japanese agricultural systems differ greatly, including different climates and soil types; most American agriculture is irrigated aerobic cultivation, whereas major Japanese agriculture is flooded rice cultivation on alluvial soils, though the aerobic cultivation is also major in soils such as Andosol. The difference in dominance of the Bj110 cluster between the United States and Japan might be affected by these factors. These results are one reason why the coefficient of determination for the relationship between community structure and latitude was lower in the United States than in Japan. However, since the transition of Bj123, Bj110, Bj6, and Be clusters are in common with Japan, the present results clearly indicate that the indigenous soybean-nodulating bradyrhizobial community in the United States varies with latitude.

In the United States, the dominance of localized *B. elkanii* strains in the soil was high. Minamisawa *et al.* (21) investigated the preferential nodulation of soybean cultivars, a wild soybean progenitor (*Glycine soja*), and siratro (*Macroptilium atropurpureum*) by *B. japonicum* and *B. elkanii* strains, and found that *B. japonicum* and *B. elkanii* preferentially nodulated *G. max* and *M. atropurpureum*, respectively, whereas both bradyrhizobial species formed nodules on *G. soja* with similar efficiency. Marr *et al.* (18) reported that *Amphicarpaea bracteata* performed nodule
formation with *B. japonicum* and *B. elkanii*, and performed nitrogen fixation with *B. elkanii*, thought *A. bracteata* will not be the original host for *B. elkanii*. Furthermore, Tang et al. (39) investigated the microevolution and origins of *Bradyrhizobium* populations in east North America associated with soybean and native legumes (*A. bracteata* and *Desmodium canadense*) using genetic characterization by multilocus sequence typing of six core (housekeeping) gene sequences and two symbiotic gene sequences, and the result suggested that soybean-nodulating bacteria associated with native legumes represent a novel source of ecologically adapted bacteria for soybean inoculation. In addition, the *Rj* soybean genotype is known to produce weak nodulation with certain *B. elkanii* strains (e.g., USDA 61 and USDA 94; 7, 8). In fact, infection by bradyrhizobia belonging to the Be94 cluster isolated from Bragg (non-*Rj*) and CNS (*Rj2*/*Rj3) cultivars in Ohio and North Carolina, which produced high dominance of *B. japonicum*, was inhibited by the soybean cultivar with the *Rj* genotype (Fig. 2). These results suggest that the interaction between host plants and rhizobia involved both host specificity and host preference. Further analysis for isolates about phenotypic traits such as compatibility and preference with *Rj* genotypes and genetic diversity of other genes such as nodulation genes and house-keeping genes of isolates must be conducted for elucidation of ecologically, evolutionally and phylogenetically relationship.

Siratro is a major pasture legume that is cultivated in the tropics and subtropics, including parts of Australia, South and Central America, and some Pacific Islands (35). Additionally, *A. bracteata* are an annual legume that distributes widely in eastern North America (23, 40). The presence of these legumes that have compatibility for nodulation with *B. elkanii* might contribute strongly to the high dominance of *B. elkanii* in North America. We expect that further research on the competition
between inoculated and indigenous bradyrhizobia and the environmental adaptability of
bradyrhizobia will lead to the establishment of improved inoculation techniques based on the use of
the most effective bradyrhizobium for a given host and region. The knowledge of the geographical
distribution of indigenous soybean-nodulating bradyrhizobia provided by the present study will help
to guide that future research.

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Table footnotes

TABLE 1. Soil sample sites for analysisa

aIdentification of soil group referred to website of the following:


TABLE 2. Alpha, beta and gamma diversity indices for each soil sample site pair
Figure legends

FIG. 1. Schematic representation of RFLP patterns of the 16S-23S rRNA gene ITS region. Reference strains and representative isolates are indicated as follows: USDA 4, 6, 38, 110, 115, 123, 124 and 135 represent *Bradyrhizobium japonicum* USDA 4, 6, 38, 110, 115, 123, 124 and 135, respectively, and USDA 46, 76 and 94 represent *Bradyrhizobium elkanii* USDA 46, 76 and 94, respectively. MIC, MIH, NHB, NCB, NCC, NCH, GAH, FLB, FLC and LAB represent Michigan-CNS, Michigan-Hill, Ohio-Bragg, North Carolina-Bragg, North Carolina-CNS, North Carolina-Hill, Georgia-Hill, Florida-Bragg, Florida-CNS and Louisiana-Bragg, respectively.

FIG. 2. Dendrogram of the 16S-23S rRNA gene ITS region of indigenous soybean-nodulating bradyrhizobia and *Bradyrhizobium* USDA reference strains. The similarity between *Bradyrhizobium elkanii* USDA 76 and 94 (which was 90%) was applied as the criterion to differentiate the clusters. Clusters were designated as indicated on the right. The diversity index \( H' \) was calculated using the following equation: 

\[
H' = -\sum P_i \ln P_i.
\]

FIG. 3. Distribution of clusters and the population ratio of indigenous soybean-nodulating bradyrhizobia in USA (map from CraftMAP [http://www.craftmap.box-i.net]).
FIG. 4. The proportion of beta diversity to gamma diversity ($H'_{\beta}/H'_{\gamma}$) among pair of the soil sampling site.

FIG. 5. Plots of indigenous soybean-nodulating bradyrhizobial community of the soil sampling site by 3-dimensional multidimensional scaling analysis based on Bray-Curtis index and dendrogram from UPGMA analysis. Dendrogram was applied as the objective index for grouping of MDS plots.

FIG. 6. Relationship between indigenous soybean-nodulating bradyrhizobial community and latitude of the soil-sampling site.
TABLE 1. Soil sample sites for analysis

<table>
<thead>
<tr>
<th>Sampling sites</th>
<th>Soil group</th>
<th>Latitude, Longitude</th>
<th>pH (H₂O)</th>
<th>EC (dS m⁻¹)</th>
<th>C (%)</th>
<th>N (%)</th>
<th>C/N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Michigan (MI)</td>
<td>Loam</td>
<td>43.05° N, -82.53° W</td>
<td>12.84</td>
<td>7.68</td>
<td>0.15</td>
<td>2.03</td>
<td>11.3</td>
</tr>
<tr>
<td>Ohio (OH)</td>
<td>Silt Loam</td>
<td>40.78° N, -81.93° W</td>
<td>10.56</td>
<td>6.34</td>
<td>0.10</td>
<td>0.97</td>
<td>10.9</td>
</tr>
<tr>
<td>Kentucky (KY)</td>
<td>Silt Loam</td>
<td>38.93° N, -86.47° W</td>
<td>6.71</td>
<td>6.14</td>
<td>0.10</td>
<td>2.05</td>
<td>11.1</td>
</tr>
<tr>
<td>North Carolina (NC)</td>
<td>Sandy Loam</td>
<td>35.79° N, -78.69° W</td>
<td>5.57</td>
<td>5.23</td>
<td>0.06</td>
<td>1.37</td>
<td>11.9</td>
</tr>
<tr>
<td>Alabama1 (AL1)</td>
<td>Loamy Sand</td>
<td>32.59° N, -85.49° W</td>
<td>2.37</td>
<td>5.77</td>
<td>0.07</td>
<td>1.16</td>
<td>0.90</td>
</tr>
<tr>
<td>Alabama2 (AL2)</td>
<td>Loamy Sand</td>
<td>32.59° N, -85.48° W</td>
<td>2.37</td>
<td>5.18</td>
<td>0.04</td>
<td>0.91</td>
<td>15.2</td>
</tr>
<tr>
<td>Georgia (GA)</td>
<td>Loamy Sand</td>
<td>31.48° N, -83.52° W</td>
<td>1.26</td>
<td>5.70</td>
<td>0.03</td>
<td>0.45</td>
<td>0.03</td>
</tr>
<tr>
<td>Florida (FL)</td>
<td>Course Sand</td>
<td>30.68° N, -85.31° W</td>
<td>0.46</td>
<td>5.58</td>
<td>0.02</td>
<td>0.40</td>
<td>0.01</td>
</tr>
<tr>
<td>Louisiana (LA)</td>
<td>Clay</td>
<td>30.32° N, -91.16° W</td>
<td>0.00</td>
<td>5.52</td>
<td>0.05</td>
<td>0.94</td>
<td>0.09</td>
</tr>
</tbody>
</table>

*aIdentification of soil group referred to website of the following:

TABLE 2. Alpha, beta and gamma diversity indices for each soil sample site pair

<table>
<thead>
<tr>
<th>Index</th>
<th>Michigan</th>
<th>Ohio</th>
<th>Kentucky</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MI-OH</td>
<td>MI-KY</td>
<td>MI-NC</td>
</tr>
<tr>
<td>α diversity ($H_\alpha$)</td>
<td>0.71</td>
<td>0.60</td>
<td>0.28</td>
</tr>
<tr>
<td>β diversity ($H_\beta$)</td>
<td>0.24</td>
<td>0.20</td>
<td>0.22</td>
</tr>
<tr>
<td>γ diversity ($H_\gamma$)</td>
<td>0.95</td>
<td>2.08</td>
<td>1.80</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Index</th>
<th>North Carolina</th>
<th>Alabama1</th>
<th>Alabama2</th>
<th>Georgia</th>
<th>Florida</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NC-NC1</td>
<td>NC-NC2</td>
<td>NC-SC</td>
<td>NC-SC</td>
<td>AL-AL1</td>
</tr>
<tr>
<td>α diversity ($H_\alpha$)</td>
<td>0.01</td>
<td>0.11</td>
<td>0.11</td>
<td>0.20</td>
<td>0.10</td>
</tr>
<tr>
<td>β diversity ($H_\beta$)</td>
<td>0.24</td>
<td>0.20</td>
<td>0.20</td>
<td>0.20</td>
<td>0.20</td>
</tr>
<tr>
<td>γ diversity ($H_\gamma$)</td>
<td>1.32</td>
<td>1.37</td>
<td>1.37</td>
<td>1.37</td>
<td>1.37</td>
</tr>
</tbody>
</table>
FIG. 1. Schematic representation of RFLP patterns of the 16S-23S rRNA gene ITS region.
FIG. 2. Dendrogram of the 16S-23S rRNA gene ITS region of indigenous soybean-nodulating bradyrhizobia and *Bradyrhizobium* USDA reference strains.

<table>
<thead>
<tr>
<th>Cluster</th>
<th>Percentage Similarity</th>
<th>Michigan</th>
<th>Ohio</th>
<th>Kentucky</th>
<th>North Carolina</th>
<th>Alabama</th>
<th>Georgia</th>
<th>Florida</th>
<th>Louisiana</th>
</tr>
</thead>
<tbody>
<tr>
<td>BJ4</td>
<td>1.56</td>
<td>3.86</td>
<td>0.31</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>BJ6</td>
<td>1.42</td>
<td>3.76</td>
<td>0.25</td>
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<tr>
<td>BJ123</td>
<td>1.17</td>
<td>3.44</td>
<td>0.20</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>BJ124</td>
<td>0.77</td>
<td>2.91</td>
<td>0.15</td>
<td></td>
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<tr>
<td>BJ135</td>
<td>0.25</td>
<td>2.32</td>
<td>0.10</td>
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</tr>
<tr>
<td>BJ110</td>
<td>1.15</td>
<td>2.60</td>
<td>0.15</td>
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<tr>
<td>BJ38</td>
<td>1.08</td>
<td>2.55</td>
<td>0.15</td>
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<tr>
<td>BJ115</td>
<td>1.39</td>
<td>2.44</td>
<td>0.15</td>
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<td>BJ46</td>
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<td>1.82</td>
<td>0.15</td>
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<tr>
<td>BJ94</td>
<td>1.15</td>
<td>1.70</td>
<td>0.15</td>
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</tr>
<tr>
<td>Be46</td>
<td>1.08</td>
<td>1.65</td>
<td>0.15</td>
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<td></td>
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<td></td>
</tr>
<tr>
<td>Be76</td>
<td>1.39</td>
<td>1.52</td>
<td>0.15</td>
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<td></td>
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<td></td>
</tr>
<tr>
<td>Be94</td>
<td>0.77</td>
<td>1.21</td>
<td>0.15</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BeOH</td>
<td>1.15</td>
<td>1.14</td>
<td>0.15</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

B. japonicum

B. elkanii

Similarity (%)
FIG. 3. Distribution of clusters and the population ratio of indigenous soybean-nodulating bradyrhizobia in USA.
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FIG. 6. Relationship between indigenous soybean-nodulating bradyrhizobial community and latitude of the soil-sampling site.

\[ y = 7.3662x - 5.9087 \]

\[ r^2 = 0.81531 \ (p < 0.001) \]

FIG. 6. Relationship between indigenous soybean-nodulating bradyrhizobial community and latitude of the soil-sampling site.