Differentiation of Dextran-Producing *Leuconostoc* Strains by a Modified Randomly Amplified Polymorphic DNA Protocol

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Seven dextran-producing *Leuconostoc* strains were differentiated by using a modified randomly amplified polymorphic DNA (RAPD) protocol that incorporated specific primers designed from conserved regions of dextranucrase genes. RAPD profiles showed intraspecies differences among the *Leuconostoc mesenteroides* strains tested. This modified RAPD protocol will aid in the differentiation of polymer-producing *leuconostocs*, which are currently distinguished by time-consuming analyses of the dextrans they synthesize.

Leuconostocs are heterofermentative lactic acid bacteria that can produce extracellular polymers such as alternan, dextran, and levans from sucrose metabolism (2, 4). Dextrans are high-molecular-weight homopolymers composed of α-1,6-glucose with predominantly α-(1→6) linkages and have been used in a variety of commercial applications (2). Dextrans are structurally diverse and are characterized according to the percentage, nature, and distribution of their non-α-(1→6) linkages (5). The exact molecular structure of a dextran is determined by the specific *Leuconostoc* strain that synthesizes the polymer (2, 5). Many *Leuconostoc* strains can produce more than one type of dextran, and the proportions of each polymer synthesized can vary depending on the culture conditions used (5, 19). Since dextran-producing leuconostocs are physiologically similar (4), differentiation of specific strains is often accomplished through unreliable and elaborate methods such as gas chromatography-mass spectrometry of methylated dextran derivatives (15), 1H and 13C nuclear magnetic resonance spectrometry (14), or Fourier-transform infrared spectroscopy (13). The development of a convenient method to aid differentiation and identification of dextran-producing *Leuconostoc* strains would assist both basic and applied research.

Several DNA-based methods for the differentiation of *Leuconostoc mesenteroides* strains have been developed (6, 17); however, the protocols used either were time-consuming or did not significantly differentiate between strains. Random amplification of polymorphic DNA (RAPD) is a PCR method that incorporates a single arbitrarily designed oligonucleotide primer in the amplification reaction to generate DNA fragment polymorphisms (18, 21). The DNA fragment polymorphisms, or RAPD profiles, have been used to distinguish between closely related bacterial species (1, 11). The RAPD technique is simple and quick, and thus it would serve as a convenient alternative protocol to complement the methods currently used to distinguish dextran-producing *Leuconostoc* strains.

The objective of this study was to determine if dextran-producing *Leuconostoc* strains could be differentiated by a modified RAPD protocol using primers designed from conserved regions of dextranucrase genes.

All *Leuconostoc* strains used in this study (NRRL B-512F, B-742, B-1118, B-1142, B-1149, B-1299, and B-1355) were obtained from the Agricultural Research Service culture collection (Peoria, Ill.) and are designated *L. mesenteroides* except for B-742, which is considered *L. citreum* (16). These *Leuconostoc* strains were chosen for this study because they are commonly used in research and for commercial applications (2, 7, 8, 10). *Leuconostoc* strains were grown in MRS (3) broth containing 2% glucose, and the genomic DNA was isolated according to the method of Pitcher et al. (12). Three oligonucleotide primers were used for RAPD analyses and were designed from conserved sequences of dextranucrase genes from *Leuconostoc* B-512F and B-1299 (9, 20). Two primers were designed from the N-terminal region of the B-512F dextranucrase gene (9, 20) and were designated 512Fa (5'-GATGCGA GTGAATATGTTGGGTGCT-3') and 512Fb (5'-GTCAAA GGATCCGTGATGAATCGGAAT-3') (9). Primers 512Fa and 512Fb were originally designed and used by Monchois et al. (9) to clone a dextranucrase gene from *L. mesenteroides* NRRL B-1299. One primer was designed from the C-terminal region of the B-1299 dextranucrase gene (9) and was designated 1299 [5'- (AGCT)CC(AG)TC(TGAGCT)CC(AG)AA(AG) TA(AGCT)ACCCA-3']. (Degenerate positions within primer 1299 are indicated by parentheses.) Primer 1299 was designed degenerate in order to increase the number of DNA fragments detected. The typical 20-μl reaction mixture for RAPD PCR analysis of the *Leuconostoc* strains contained 1× *Pfu* buffer (Stratagene, La Jolla, Calif.), 250 μM (each) dATP, dCTP, dGTP, and dTTP (Stratagene), 1 μg of template DNA, 0.25 μg of a single oligonucleotide primer, and 1.5 U of *Pfu* DNA polymerase (Stratagene). A denaturation step of 95°C for 1 min was performed with the reaction mixture before PCR cycling. The RAPD PCR cycling program consisted of 40 cycles of denaturation at 95°C for 1 min, annealing at 30°C for 1 min, and primer extension at 72°C for 5 min. After the cycling program was finished, the samples were held at 72°C for 10 min to complete the extension of products. PCR amplification was performed in a Progene thermocycler (Techne Incorporated, Princeton, N.J.). The DNA banding patterns were examined by
using 1.5% agarose gel electrophoresis, and a DNA ladder (0.25 to 12 kb) was used as size standards (Stratagene).

Results from the modified RAPD analysis using each primer are illustrated in Fig. 1 through 3. All seven *Leuconostoc* strains that were examined by RAPD analysis could be differentiated based on DNA banding patterns when all three oligonucleotide primers were used in separate tests. Many of the strains shared common DNA band sizes (for example, strains B-512F, B-1118, and B-1149 in Fig. 1, strains B-1142 and B-1299 in Fig. 2, and strains B-742 and B-1142 in Fig. 3), which is not unexpected, since all the strains except B-742 are designated *L. mesenteroides*. All the strains tested showed reproducible RAPD profiles based on DNA band size, number, and intensity after three trials.

All the dextran-producing *Leuconostoc* strains tested could be differentiated from each other by RAPD PCR amplification using a combination of primers that were designed from conserved regions of dextransucrase genes. RAPD analysis showed intraspecies differences between the dextran-producing strains examined. Use of all three primers for RAPD analysis would ensure accurate differentiation of dextran-producing *Leuconostoc* strains when control strains are used. It is not clear why long (24-mer), specific primers produced polymorphic DNA banding patterns, since short, arbitrary primers are usually used in RAPD PCRs. Only two *Leuconostoc* dextransucrase genes have been cloned (from B-512F [20] and B-1299 [9]), and these displayed 55% amino acid sequence similarity (9). Evidence has indicated that many polymer-producing *Leuconostoc* strains produce more than one type of dextransucrase (2), which may account for some of the DNA polymorphisms observed during RAPD analysis. *L. mesenteroides* B-512F, however, has only one type of dextransucrase (2) but still displays DNA polymorphism when the modified RAPD protocol is followed. The presence of multiple dextransucrase genes may not account for all the DNA polymorphism observed when the modified RAPD protocol is used. Although the exact molecular mechanism defining the modified RAPD protocol is unknown, this should not preclude its use as a tool to differentiate polymer-producing *Leuconostoc* strains.

RAPD analysis was proven to be a simple and reproducible method for differentiation of the dextran-producing *Leuconostoc* strains compared to other strain characterization methods which involve polymer structural analyses. This study provides a basis for molecular differentiation of dextran-producing *Leuconostoc* strains that are used in basic and applied research.

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REFERENCES


