

# Complete Sequences of the *S*-genes, *Sd*- and *Sh*-RNase cDNA in Apple

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**Abstract.** The *S*-locus genes in the pistil (*S*-RNases) were cloned from the apple (*Malus × domestica* Borkh.) cultivar Akane (*S*-genotype *SdSh* from pollination analysis). The *Sd*- and *Sh*-RNase corresponded to *S7*- and *S24*-RNase, which have been cloned from 'Idared' and 'Braeburn', respectively. *Sh*-RNase was very similar to *Sf*- and *Sg*-RNases at the deduced amino acid-sequence levels (93%). We developed an *S*-allele specific polymerase chain reaction (PCR)-restriction fragment length polymorphism (RFLP) analysis method for distinguishing the *Sh* from *Sf* and *Sg*, and the *Sh*-alleles of 'Akane', 'Touhoku 2', 'Vista Bella', and 'Worcester Pearmain' were identified. We also identified the *S*-allele genotypes of 16 apple cultivars.

Gametophytic self-incompatibility in the apple is controlled by a single, multiallelic locus, the *S*-locus. In this system, pollen tube growth is inhibited when pollen shares the same *S*-allele with the pistil on which the pollen landed.

The *S*-locus genes in the pistil have been cloned from apple cultivars. As the *S*-genes in the pistil encode ribonuclease, they are called *S*-RNases. *Sc*-, *Sf*-, and *Sg*-RNases have been cloned from apple cultivars in Japan (Kitahara et al., 1999; Sassa et al., 1996), and *S2*-, *S3*-, *S5*-, *S7*-, *S9*-, *S24*-, *S26*-, and *S27*-RNases from apple cultivars in Europe (Brothoerts et al., 1995; Janssens et al., 1995; Verdoodt et al., 1998). *Sc*- and *S9*-RNase are identical at the deduced amino acid sequence level.

Komori et al. (1998) established *Sa*-, *Sb*-, *Sc*-, *Sd*-, *Se*-, and *Sf*-allele genotypes by polli-

nation and progeny analysis of apple cultivars in Japan. Using the molecular method of *S*-allele identification based on polymerase chain reaction (PCR)-restricted fragment length polymorphism (RFLP) analysis, developed by Janssens et al. (1995) and modified by Sakurai et al. (1997) and Matsumoto et al. (1999a, 1999b), to analyze apple cultivars in Japan, it was confirmed that *Sa*, *Sb*, and *Sd* correspond to *S2*, *S3*, and *S7*, respectively. However, *Se*, *Sf*, and *Sg* did not correspond to *S5*, *S24*, *S26*, or *S27*.

Previously, the *S*-allele genotype of 'Akane' was identified as *S7S* (unknown) using the PCR-RFLP method (Sakurai et al., 1997). *S7*-allele corresponds to the *Sd*-allele, and the *S*-allele genotype of 'Akane' was identified as *SdSh* by pollination and progeny analysis (S. Komori, personal communication). In this study, the cDNA sequences encoding *Sd*- and *Sh*-RNase were determined. We also identified the *S*-allele genotypes of 16 apple cultivars by *S*-allele-specific PCR-RFLP analyses.

## Materials and Methods

**Plant material.** Most of the plants used in this study were from collections at the Apple Research Center of the National Institute of Fruit Tree Science, Japan. 'Akibae' and 'Chouka 8' were obtained from the Nagano Fruit Tree Experiment Station, Japan. 'Akane'

flowers were collected at the popcorn stage, and the pistils were removed and immediately frozen in liquid nitrogen and stored at  $-80^{\circ}\text{C}$ . Pistils of other cultivars were obtained in the same way. Young leaves were collected and stored at  $-80^{\circ}\text{C}$  until used.

***Sh*-RNase cDNA cloning.** Total RNAs of the mature pistils were extracted essentially as described by Chang et al. (1993). Reverse transcription (RT) was performed in a total volume of 20  $\mu\text{L}$  containing 1  $\mu\text{L}$  ( $\approx 1.0\ \mu\text{g}$ ) of RNA, 0.125  $\mu\text{M}$  Oligo dT-Adaptor Primer (Takara Shuzo Co., Kusatsu, Shiga, Japan), 0.25 U AMV Reverse Transcriptase XL (Takara Shuzo Co.), 20 U RNase Inhibitor (Takara Shuzo Co.), 1 mM dNTPs, 5 mM  $\text{MgCl}_2$ , and 1 $\times$  RNA PCR buffer (10 mM Tris-HCl pH 8.3, 50 mM KCl). The analysis was programmed in a thermal cycler (GeneAmp 2400 apparatus; Perkin-Elmer Co., Urayasu, Chiba, Japan) and conducted under the following conditions: 60 min at  $42^{\circ}\text{C}$ , 5 min at  $99^{\circ}\text{C}$ , and cooling to  $4^{\circ}\text{C}$ . A PCR was carried out in a total volume of 100  $\mu\text{L}$  containing 20  $\mu\text{L}$  of the RT reaction mixture; 0.2  $\mu\text{M}$  OWB134 primer (Brothoerts et al., 1995), 0.2  $\mu\text{M}$  M13 Primer M4 (Takara Shuzo Co.), which was complementary to the adaptor region of the OligodT-Adaptor primer; 2.5 mM  $\text{MgCl}_2$ , 1 $\times$  RNA PCR buffer; and 2.5 U *Taq* DNA polymerase (TaKaRa *Taq*<sup>TM</sup>; Takara Shuzo Co.). The amplification program consisted of 2 min at  $94^{\circ}\text{C}$  for initial denaturation, 45 cycles of 30 s at  $94^{\circ}\text{C}$ , 30 s at  $50^{\circ}\text{C}$ , 1.5 min at  $72^{\circ}\text{C}$ , and a final extension of 7 min at  $72^{\circ}\text{C}$ .

The cDNAs obtained were excised from a 1.0% (w/v) agarose gel in TAE (40 mM Tris-acetate, 1 mM EDTA pH 8.0), purified by the GeneClean kit (BIO 101 Co., Calif.) and subcloned into a pCRII cloning vector with the TA cloning kits (Invitrogen, Co., Hongo bunkyo-ku, Tokyo). The nucleotide sequences of three clones named *Sh*-1, -2, and -3 were determined by dideoxy chain termination on an ABI PRISM<sup>TM</sup> 377 DNA sequencer (Perkin-Elmer, Co.) using dRhodamine Terminator Cycle Sequencing Kits (Perkin-Elmer Co.).

***Sd*-RNase cDNA cloning.** OWB 126 primer (Janssens et al., 1995) was used as the gene-specific primer for rapid amplification of cDNA 3' ends (3' RACE) (Frohman et al., 1988). A PCR was carried out in a total volume of 100  $\mu\text{L}$  containing 20  $\mu\text{L}$  of the RT reaction mixture, 0.2  $\mu\text{M}$  OWB 126 primer, 0.2  $\mu\text{M}$  M13 Primer M4 (Takara Shuzo Co.), 2.5 mM  $\text{MgCl}_2$ , 1 $\times$  RNA PCR buffer, and 2.5 U *Taq* DNA polymerase (TaKaRa *Taq*<sup>TM</sup>; Takara Shuzo Co.). The amplification program consisted of 2 min at  $94^{\circ}\text{C}$  for initial denaturation, 45 cycles of 30 s at  $94^{\circ}\text{C}$ , 30 s at  $52^{\circ}\text{C}$ , 1.5 min at  $72^{\circ}\text{C}$ , and a final extension of 7 min at  $72^{\circ}\text{C}$ .

PCR products were separated on a 1.0% (w/v) agarose gel in TAE (40 mM Tris-acetate, 1 mM EDTA, pH 8.0), and bands of ca-600 bp were purified by the GeneClean Kit (BIO 101 Co.), then subcloned with the pCRII cloning vector. Two cDNA 3' end clones named *Sd*-3'-1 and -2 were obtained.

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To obtain the cDNA 5' ends of the cDNA 3' end clones, rapid amplification of cDNA 5' ends (5' RACE) was carried out using a 5'-Full RACE Core Set (Takara Shuzo Co.). First, the reaction mixtures were prepared according to the manufacturer's instructions (Takara Shuzo Co.). These mixtures were for first strand cDNA synthesis by RT using the cDNA 3' end clone specific 5'-phosphorylated RT-primer, Sd-PF1 (5'-P-TCA-GTCACCTGGTTA-3'; nucleotides 624-638 in Fig. 2), which is specific for Sd-3'-1 and -2. The analysis was conducted under the following conditions: 10 min at 30 °C, 60 min at 50 °C, and 2 min at 80 °C, followed by cooling to 4 °C.

After the RT reaction, the hybrid RNA of the cDNA first strand was degraded by RNaseH, then the cDNA first strand was self-ligated by T4 RNA ligase according to the manufacturer's instructions (Takara Shuzo Co.).

For the first and second PCRs of the self-ligated cDNA first strand with Sd-PF1 primer, two sets of the oligonucleotide primers were designed. For the first PCR the primers were Sd-5S1 (5'-AAACATGGCACCTGTGGACA-3'; nucleotides 409-428 in Fig. 2) and Sd-5A1 (5'-ACTGAGTTGTCAGATTTCTC-3'; nucleotides 321-340 in Fig. 2). For the second PCR the primers were Sd-5S2 (5'-ATCCCA-CATAATGAACGAC-3'; nucleotides 428-447 in Fig. 2) and Sd-5A2 (5'-AGCATCCA-CGGTTGATTTAC-3'; nucleotides 293-312 in Fig. 2). The reaction mixtures of the first and second PCRs were prepared according to the manufacturer's instructions (Takara Shuzo Co.). The amplification program for the first PCR consisted of 3 min at 94 °C for initial denaturation, 25 cycles of 30 s at 94 °C, 30 s at 55 °C, 1.5 min at 72 °C, and a final extension of 7 min at 72 °C. The program for the second PCR consisted of 35 cycles of 30 s at 94 °C, 30 s at 55 °C, 1 min at 72 °C, and a final extension of 7 min at 72 °C. After the second PCR, the PCR products were separated as described above, and they were then subcloned with the pCRII-cloning vector. Two cDNA clones named Sd-5'-1 and -2, corresponding to Sd-3' clones, were obtained. The nucleotide sequences of Sd-3' and Sd-5' clones were determined as described above.

**S-allele specific PCR analysis of DNA.** Total DNA from leaves of individual plants was isolated as described by Thomas et al. (1993). S2-, S3-, S5-, S7-, S9-, Sf-, and Sg-allele-specific PCR amplification and digestion were carried out essentially as described by Matsumoto et al. (1999a, 1999b). For Sh-allele identification, PCR was conducted using sense (5'-ACGATCATGAAGCTT-CTGGCG-3'; Sf-sense 3) and antisense (5'-TTGGTGGGGCAGAAATTCC-3'; Sf-antisense 2 from Matsumoto et al., 1999a) primers, with the total DNA of each cultivar being expected to possess the Sf- or Sg- or Sh-allele. Each 35-µL PCR mixture contained 1.0 µM of each primer, 200 µM deoxynucleotides, 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 1.5 mM MgCl<sub>2</sub>, 0.001% (w/v) gelatin, 2.5 U *Taq* polymerase, and 50 ng template DNA. The analy-

sis was programmed in a thermal cycler (GeneAmp 2400 apparatus; Perkin-Elmer Co.) and conducted under the following conditions: 3 min at 94 °C for preheating, 1 min at 94 °C, 1 min at 56 °C, and 1 min at 72 °C for 30 cycles, followed by an extension for 10 min at 72 °C. After PCR, the amplified fragments were digested by *Sna*BI. The amplified and digested fragments were separated on a 4% polyacrylamide gel in TAE (40 mM Tris-acetate, 1 mM EDTA, pH 8.0) at 50-70 V for 1-1.5 h.

## Results and Discussion

The OWB134 primer anneals to a conserved region in all known S-RNases, and its 3'-nucleotide triplet corresponds with translation initiation codon ATG (Verdoodt et al., 1998). An RT-PCR product of ≈840 bp from 'Akane' apple was subcloned for further analysis.

We sequenced three independent clones; all cDNA inserts represented an S-locus gene (Fig. 1). The nucleotide sequences of the clones termed Sh-1, -2, and -3 were matched with those of the S24-RNase from 'Braeburn' (Verdoodt et al., 1998) except for two nucleotides in 3' untranslated region (nucleotide positions 828 from T to C, and 832 from T to A in Fig. 1). The protein sequences of Sh- and S24-RNase are identical, as are those of Sc- and S9-RNase. We could not obtain the cDNA clone corresponding to Sd-RNase by using the OWB134 primer, because the primer might not anneal to the conserved region of the Sd-RNase. Because the Sd-RNase corresponds to S7-RNase, the Sd-RNase cDNA 3' end clone should be obtained by 3' RACE methods with the primer OWB 126 designed from S7-RNase by Janssens et al. (1995).

We obtained two 3' end cDNA clones termed Sd-3'-1 and -2. The two clones were identical except for the site of a poly (A) addition (nucleotide position 798 for Sd-3'-1

and 840 for Sd-3'-2 in Fig. 2). The nucleotide sequences of Sd-3' clones were matched with those of the S7-RNase from 'Idared' except for the one nucleotide at nucleotide position 540 from A to G, and the site of a poly (A) addition (nucleotide position 780 for S7-RNase cDNA in Fig. 2). This difference represented in the third letter of triplet codon (GGA for S7-RNase and GGG for Sd-3'), and both of the codons representing one amino acid, glycine. Two 5' end cDNA clones, designated Sd-5'-1 and -2 corresponding to the Sd-3' clones were then obtained by 5' RACE method. The putative initiation codon ATG at position 67-69 is preceded by a stop codon TAA at position 55-57 in-frame, and the sequence surrounding the first ATG codon (ATTCATG) was well conserved in the cDNAs of the apple S-genes. The nucleotide sequence of Sd-5' was identical to that of the S7-RNase from 'Idared' except for the 5'-end of the untranslated region and the coding region encoding the first 49 amino acids (nucleotides 1-212 in Fig. 2), which are uncloned in the cDNA clone of S7-RNase (Fig. 2). From these results, the protein sequences of Sd- and S7-RNase seem to be identical. Although Sakurai et al. (1997) identified the S-allele genotype of 'Akane' as S7S (unknown), we designated the S7-RNase in S7-allele of 'Akane' as the Sd-RNase in Sd-allele. Because the nucleotide sequence of S7-RNase isolated from 'Idared' differed from that in 'Akane', we could not confirm whether or not the first 49 amino acids of the S7-RNase in 'Idared' were exactly the same as those of the Sd-RNase in 'Akane'.

The cloned cDNA inserts of Sd (851 bp) and Sh (840 bp) contain 684 bp (228 amino acids) and 678 bp (226 amino acids) of coding regions, respectively. In apple, S2-, S3-, S9-, Sf-, Sg-, S24-, S26-, and S27-RNase code for polypeptides containing 226-228 amino acids. The termination codon TAA is present at positions 751-753 (Sd) and 679-681 (Sh), followed by the 3' untranslated

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ATGGGGACCGGGATGATATATATGGTTATGATGGTATTTTCACTAATTTTATTAATATTG      60
M G T G M I Y M V M M V F S L I L L I L      20
CCTTCGTCACCGGTGGGATTCGATTAATATCAATTTACGCAGCAATATCAGCCGGCTGTC      120
P S S T V G F D Y Y Q F T Q Q Y Q P A V      40
TGCAACTCTAATCCAACCTCTGTAAGGATCCTACTGACAAGTTGTTTACTGTTTCACGGT      180
C N S N P T P C K D P T D K L F T V H G      60
TTGTGGCCTTCAAACCTCGAATGGAAATGACCCAAAATATGCAACGCACAGCAATATCAG      240
L W P S N S N G N D P K Y C N A Q Q Y Q      80
ACGATGAAAATACTCGAACCCAGTTGGTAATTTATTTGGCCGAACTACTCAATCGAAAC      300
T M K I L E P Q L V I I W P N V L N R N      100
GATCATGAAGGCTTTCGGCTAAACAGTGGGAGAAACATGGCTCCTGTGGCTTTCCTCCCA      360
D H E G F W R K Q W E K H G S C A S S P      120
ATACAGAACCCAGAAGCATTACTTTGATACAGTAATCAAAATGTACCAACCCGAGAACAA      420
I Q N Q K H Y F D T V I K M Y T T Q K Q      140
AACATCTCTGAATCCTCTCAAAGGCGAATATAAAACCGGGTAGGAAAACAGGACACTG      480
N I S E I L S K A N I K P G R K N R T L      160
GTGGATATTGAAAATGCCATACGTAATGTTATCAACAAATATGACACCAATTCAGTGC      540
V D I E N A I R N N V I N N M T P F C K C      180
CAAAGAATACTAGGACATCATTGACTGAATTTGGTTGAGGTCGGTCTTTTCAGCGATAGC      600
Q K N T R T S L T E L V E V G L C S D S      200
AACTTAACCGCACTTCATAAAATGGCCCCGCCCATTTCCACGAGGATCAGCGTATTTCTCG      660
N L T Q F T I A N C P R P F P R G S R Y F C      720
CCCACAAATATTCAGTATTAAGATCGCGGCTAGCTAGTGTGTGTGTCTGTATGTATACAC      780
P T N I Q Y *
ATATAAAATAATAGTAGTGGCTTACACCGGATGACGATTTCCATATATAGGATGAGAA      840
TGTGATGTAATTCFAACGAACTTGGCTTAATTAATTAATAATAGTACGAAAAA          840
AAA

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Fig. 1. Nucleotide and deduced amino acid sequences of the Sh-RNase cDNA from 'Akane' apple. The first three nucleotides were from the 3'-nucleotide triplet of the OWB 134 primer. Asterisk indicates stop codon. The sequence of the Sh-RNase cDNA was deposited under the DDBJ accession number AB032247.

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CCACCACCACTTCGAATCGATCAAATTAGTAATTAATCTGCCTCGCTCTTCGACTAATAT 60
TATTC AATGGGGATTACGGGGATGATATATATGGTTACGATAGTATTTTCATTAATPGTT 120
      M G I T G M I Y M V T I V F S L I V 18
TTACTATTGCTTCGTCCTCCGGCGGAGATACGATTATTTTCAATTTACGCAGCAGTATCAG 180
L L L S S S A A R Y D Y F Q F T Q Q Y Q 38
TTGGCTGCCTGCAACTCTAAACCTATTCCTTGTAAAGGATCCCTCCTGACAAGTTGTTTACG 240
L A A C N S K P I P C K D P P D K L F T 58
GFTCCAGGTTTGTGGCCTTCAGACTCGAATGGACATGACCCAGTAAATTCGAGTAAATCA 300
V H G L W P S D S N G H D P V N C S K S 78
ACCCTGGATGCTCAGAACTTAGGAAATCTGACAACCTCAGTTGGAAATAATTTGGCCGAAC 360
T V D A Q K L G N L T T Q L E I I W P N 98
GTATACAATCGAACCAGTATATAAGCTTCTGGGATAAACAGTGGAAACAACATGGCACC 420
V Y N R T D H I S F W D K Q W N K H G T 118
TGTGGACATCCCAATAATGAACGACATTCATTACTTTCAAACAGTAAATCAAATGTAC 480
C G H P T I M N D I H Y F Q T V I K M Y 138
ATAACCAGAAACAAACGCTCTCAAATCCTCTCAAGGCGGAAGATTGAACCGGAGGG 540
I T Q K Q N V S K I L S R A K I E P E G 158
AAACCCAGGAAACAGGTAGATATTGTAATGCCATACGCCAAAGGTACAAACGATAAGGAA 600
K P R K Q V D I V N A I R K G T N D K E 178
CCAAACTCAAGTCCAAAAGAAATAACCAGGTGACTGAATGGTTGAGTCACTCTTTGC 660
P K L K C Q K N N Q V T E L V E V T L C 198
AGTAATCGCAACCTAACGGGTTTCATAAATGCCCCCGCATATTCCAAATGGATCCAGCA 720
S N R N L T G F I N C P R H I P N G S R 218
TATTCCTGCCCAAAAATATCTGTATTAAGAGCGCTGCTAGCTATATATATAGAGCG 780
Y S C P T K N I L Y * 228
GGTGACCATTTCCATATAATGAATAACTTGGCTTAATTAATTAACATCATGTGAATATGA 840
AAAAAAAAAAAA 851
    
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region. The deduced amino acid sequence of Sh-RNase showed a high similarity to both Sf- (92.5%) and Sg-RNase (93.4%). The similarities ranged from 92.5% to 93.8% among Sf-, Sg-, and Sh-RNase, and from 57.0% to 73.7% for S2-, S3-, S5-, Sd-, S9-, S26-, and S27-RNase.

We identified the S2-, S3-, S5-, Sd- (=S7-), S9-, and Sf-alleles of 'Akane', 'Akibae', 'Ambitious', 'Benihazuki', 'Chouka 8', 'Chouka 17', 'Cox's Orange Pippin', 'Hac 9', 'Homei', 'Kidd's Orange Red', 'Michinoku', 'Natsumidori', 'North Queen', 'Shinsekai', 'Worcester Pearmain', and 'Yataka' using the S-allele specific PCR-RFLP analysis method described by Matsumoto et al. (1999a, 1999b) (Table 1). These cultivars are commercially less important than those examined by Sakurai et al. (1997) and Matsumoto et al. (1999a, 1999b). However, as these cultivars might be parents of new cultivars in the future, knowledge of their S-genotypes may be useful in selecting compatible parents in a breeding program.

Although the method of S24-allele specific PCR analysis was developed by Verdoodt et al. (1998), the method cannot distinguish S24 from Sf and Sg. From the sequence data of Sf-RNase of 'Fuji' (Sassa et al., 1996), Sg-RNase of 'Indo' (Kitahara et al., 1999) and Sh-RNase of 'Akane' (this paper), we developed a method to distinguish Sh- (= S24) from Sf- and Sg-alleles using restriction enzyme analyses. A 370-bp genomic fragment was obtained from 'Indo', of which the S-genotype was S7Sg (Matsumoto et al., 1999a), 'Iwakami', of which the S-genotype was S7Sf (Matsumoto et al., 1999b) and from 'Akane', using the Sf-, Sg- and Sh-allele-specific primers, Sf-sense 3 and Sf-antisense 2 (Fig. 3). No fragment was obtained from 'Golden Delicious', of which the S-genotype was S2S3 (Janssens et al., 1995), 'Kidd's Orange Red' (S5S9), 'Chouka 8' (S3S7), or 'Baskatong', of which the S-

Fig. 2. Nucleotide and deduced amino acid sequences of the Sd-RNase cDNA from 'Akane' apple. Asterisk indicates stop codon. The sequence of the Sd-RNase cDNA was deposited under the DDBJ accession number AB032246.

Table 1. S-allele genotypes of apple cultivars.

Cultivars	Parentage	S-allele genotypes
Akane	Jonathan x Worcester Pearmain	SdSh
Akibae	Senshu x Tsugaru	S3Sf
Ambitious	Toukou x ? <sup>z</sup>	S2S9
Benihazuki	Tsugaru x ? <sup>z</sup>	S2S3
Chouka 8	Akane x Tsugaru	S3Sd
Chouka 17	Golden Delicious x Senshu	S2Sd
Cox's Orange Pippin		S5S9
Hac 9	Fuji x Tsugaru	S3S9Sf
Homei	mutant of Tsugaru	S3Sd
Kidd's Orange Red		S5S9
Michinoku	Kitakami x Tsugaru	S3S9
Natsumidori	Kitakami x Meku 10	S3S9
North Queen	Fuji x Tsugaru	S3Sf
Shinsekai	Fuji x Akagi	S3Sf
Worcester Pearmain		S2S3
Yataka	sport of Fuji	S9Sf

<sup>z</sup>?Pollen parent is unknown.

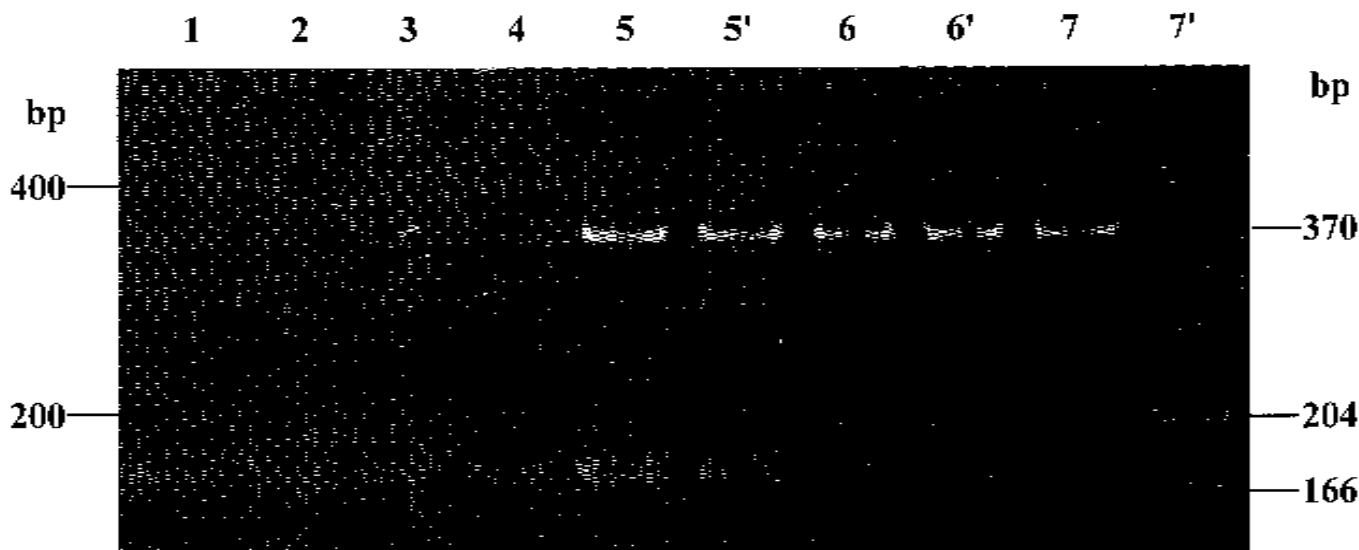


Fig. 3. Sh-allele specific PCR-RFLP analysis. PCR products obtained by using 'Golden Delicious' (S2S3) (lane 1), 'Kidd's Orange Red' (S5S9) (lane 2), 'Chouka 8' (S3S7) (lane 3), 'Baskatong' (S26S27) (lane 4), 'Indo' (S7Sg) (lane 5, 5'), 'Iwakami' (S7Sf) (lane 6, 6'), and 'Akane' (SdSh) (lane 7, 7') genomic DNA as templates, and Sf-sense 3 and Sf-antisense 2 primer. The products were digested by SnaBI (lane 5', 6', 7').

genotype was *S26S27* (Verdoodt et al., 1998) (Fig. 3). By *Sna*BI digestion, the 370-bp genomic fragment of 'Akane' (*Sh*) was separated into 204-bp and 166-bp fragments, while the fragments of 'Iwakami' (*Sf*) and 'Indo' (*Sg*) were not (Fig. 3). Using this method, the *Sh*-allele in 'Worcester Pearmain' was identified (Table 1). The *Sh*-allele was also present in 'Vista Bella' and 'Tohoku 2' ('McIntosh' x 'Worcester Pearmain').

Previously, we developed a method to distinguish the *Sf*- and *Sg*-allele by PCR-RFLP (Matsumoto et al., 1999a). In this method, the 1047 bp (*Sf*) or 1021 bp (*Sg*) of amplified fragments obtained by using *Sf*-sense 2 and *Sf*-antisense 2 primers were digested by *Rsa*I or *Sau*3AI, then each allele was identified by specific fragments for the *Sf*- or *Sg*-allele (Matsumoto et al., 1999a). Although about a 1040-bp genomic fragment was obtained from *Sh* using the primers *Sf*-sense 2 and *Sf*-antisense 2, neither the 337-bp fragment specific for *Sf* nor the 122-bp and the 78-bp fragments specific for *Sg* obtained by *Rsa*I digestion (Matsumoto et al., 1999a) were detected from *Sh* (data not shown). This result indicates that the method to distinguish between the *Sf*- and *Sg*-allele could be useful for identifying those

alleles within *SfSh* or *SgSh* genotypes. The cDNA sequences of the S-RNases in the *Se*-allele and in 'McIntosh' (*S*-allele genotype unknown) are under investigation.

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