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# Microsatellite markers spanning the apple (Malus x domestica Borkh.) genome 

Received: 19 December 2005 / Revised: 27 April 2006 / Accepted: 18 May 2006 / Published online: 9 August 2006
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#### Abstract

A new set of 148 apple microsatellite markers has been developed and mapped on the apple reference linkage map Fiesta $x$ Discovery. One-hundred and seventeen markers were developed from genomic

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libraries enriched with the repeats GA, GT, AAG, AAC and ATC; 31 were developed from EST sequences. Markers derived from sequences containing dinucleotide repeats were generally more polymorphic than sequences containing trinucleotide repeats. Additional eight SSRs from published apple, pear, and Sorbus torminalis SSRs, whose position on the apple genome was unknown, have also been mapped. The transferability of SSRs across Maloideae species resulted in being efficient with $41 \%$ of the markers successfully transferred. For all 156 SSRs, the primer sequences, repeat type, map position, and quality of the amplification products are reported. Also presented are allele sizes, ranges, and number of SSRs found in a set of nine cultivars. All this information and those of the previous CH-SSR series can be searched at the apple SSR database (http://www. hidras.unimi.it) to which updates and comments can be added. A large number of apple ESTs containing SSR repeats are available and should be used for the development of new apple SSRs. The apple SSR database is also meant to become an international platform for coordinating this effort. The increased coverage of the apple genome with SSRs allowed the selection of a set of 86 reliable, highly polymorphic, and overall the apple genome well-scattered SSRs. These SSRs cover about $85 \%$ of the genome with an average distance of one marker per 15 cM .

Keywords SSR • Genetic mapping •
Simple sequence repeat

## Introduction

In the last few years, apple genetics has made significant progresses, partly due to the increasing availability of multi-allelic SSR markers. These markers proved to be extremely useful for integrating mapping results from independent studies and in the development of innovative procedures for assessing marker-gene associations. Their high level of transferability, general high level of poly-
morphism, and the relative ease by which they are generated, being PCR-based, makes them the marker of choice for alignments among linkage maps of apple. As a result, recent maps have been based on a backbone of multi-allelic SSR markers embedded in RAPD and/or AFLP markers, which can be produced in large amounts in a relatively short time (Liebhard et al. 2002, 2003b; Kenis and Keulemans 2005). Alignments can be made for the identification and orientation of corresponding homologous linkage groups as well as for relative positions of specific loci.

SSR markers are, therefore, extremely valuable for building integrated genetic maps comprising genes, confidence intervals of QTLs, and other loci gathered from multiple maps. SSR-based maps have been essential for identification and positional comparison of major genes and QTLs for scab, powdery mildew, and fire blight resistance (Durel et al. 2000, 2003; Evans and James 2003; Liebhard et al. 2003c; Calenge et al. 2004; Gygax et al. 2004; James et al. 2004; Patocchi et al. 2004; Tartarini et al. 2004; Vinatzer et al. 2004; Calenge et al. 2005; Calenge and Durel 2005; Khan et al. 2006) as well as morphological or physiological traits (Conner et al. 1998; King et al. 2000; Liebhard et al. 2003a; Costa et al. 2005). Recognition of SSR associations led to the identification of clusters of resistance genes (e.g., Bus et al. 2004) and to the first integrated map of apple that reviews many papers with regard to the location of scab resistance genes (Durel et al. 2004). The recent discovery of many new gene-SSR marker associations, made necessary a new update of this integrated map with recently mapped scab resistance genes (Bus et al. 2005a,b; Patocchi et al. 2005), thus continuously improving our understanding of the organization of the apple genome.

Technically, SSR markers are highly suitable for direct genotyping of any new individual, being easily transferable, multi-allelic, and having a known map position. They are also the most cost effective marker for directed genome-wide genotyping approaches, as with known map positions only a relatively low number of wellselected markers have to be tested to obtain a good coverage. The possibility of multiplexing several SSR markers in the same PCR reaction allows an additional reduction of the costs of genotyping.

A good coverage of the apple genome with SSR markers is the prerequisite for two innovative techniques for assessment of molecular-marker trait associations. Firstly, the Genome Scanning Approach (GSA, Patocchi and Gessler 2003) allows efficient mapping of major genes. This procedure was successfully applied to map the apple scab resistance genes Vr2, Vm (Patocchi et al. 2004; Patocchi et al. 2005), and $V b$ (Erdin et al. 2006). Secondly, the Pedigree Genotyping concept was developed (Van de Weg et al. 2004), which allows the exploitation of breeding material in the assessment of marker-trait associations and in allele mining by using multiple pedigreed plant populations, which can be any combination of crosses, cultivars, and breeding lines. This concept makes use of directed genotyping and the so-called Identity By Descent
(IBD) concept. It forms the base of the EU-HiDRAS project (Gianfranceschi and Soglio 2004) aimed at a proof of concept for Pedigree Genotyping and at the identification of molecular markers for fruit quality and disease resistance.

The major disadvantage in the use of SSR markers is the considerable initial investment needed to develop and map them. Although around 160 SSRs have been developed for the apple (Guilford et al. 1997; Gianfranceschi et al. 1998; Hokanson et al. 1998; Liebhard et al. 2002; Hemmat et al. 2003; Vinatzer et al. 2004), their distribution within the genome is not homogenous. Almost all linkage groups contain regions with large gaps between two successive SSRs (Liebhard et al. 2003b). The development of new SSRs may solve this problem. Thus far, the most widely used method to produce SSRs is based on the cloning and sequencing of genomic fragments enriched for a repeated sequence and the designing of upstream and downstream primers (Tenzer et al. 1999; Gautschi et al. 2000). Additionally, some new SSR markers have been obtained by selecting the transferable SSR markers from closely related species (Yamamoto et al. 2004).

Recently, apple genomic projects have made thousands of apple EST sequences available, which can now be searched for SSR repeats and used for the development of new SSR markers (Crowhurst et al. 2005; Korban et al. 2005). This approach has several advantages: 1) no enriched genomic library has to be constructed; 2) extensive sequencing is not necessary, thus reducing the cost of the development of the SSR markers; 3) it is possible to develop SSR markers for which it is difficult to construct enriched libraries (e.g., AT repeats), and last but not least, 4) markers are developed from coding sequences.

In this paper, we present a new, extensive set of apple SSRs, developed within the framework of the HiDRAS European project (Gianfranceschi and Soglio 2004), from genomic libraries, publicly available EST sequences, and SSR markers of other species closely related to Malus. All these SSRs are also tested for their level of polymorphism and are positioned on a molecular marker linkage map. These SSRs, together with those already published, have been used to select a set of 86 highly polymorph SSRs well-scattered on the apple genome.

## Materials and methods

## Plant material and DNA extraction

Cultivars Elstar, Golden Delicious, and Florina were used to construct the SSR libraries. A series of nine diploid cultivars (Fiesta, Discovery, Florina, Nova Easygro, TN108, Durello di Forlì, Prima, Mondial Gala, and Fuji) was used to estimate the level of polymorphism of the new markers. Forty-four progeny plants of the Fiesta $\times$ Discovery cross, which is a subset of the 251 plants used by Liebhard et al. (2003b) to generate "the reference map", were used to map the new SSRs. Three other mapping populations, Discovery $\times$ TN10-8 (149 plants),

Durello di Forlì $\times$ Fiesta (subset of 60 plants), or Fuji $\times$ Mondial Gala (subset of 60 plants), were used to map the SSRs that could not be mapped in the cross Fiesta $\times$ Discovery. DNA was extracted according to Koller et al. (2000), gel quantified and diluted to $1 \mathrm{ng} / \mu \mathrm{l}$.

SSR development

## Genomic libraries

SSR-enriched libraries from Elstar were developed at Plant Research International. The procedure for microsatellite enrichment by selective hybridization was modified from Karagyozov et al. (1993) by Van de Wiel et al. (1999) and Van der Schoot et al. (2000). DNA was digested with TaqI and size-fractionated by agarose gel electrophoresis. Fragments between 300 and $1,000 \mathrm{bp}$ were recovered by electro-elution, enriched by hybridization to five oligonucleotides (GA, GT, ATC, AAG, ACC), ligated in pGEM-T (Promega) or pCRII-TOPO (Invitrogen) and transformed to competent TOP10 F' (Invitrogen). Colonies were transferred onto Hybond $\mathrm{N}+$ membranes and hybridized with the appropriately labeled oligonucleotides. Positive clones were sequenced with the primers $\mathrm{Sp6}$ and T 7 by Greenomics ${ }^{\mathrm{TM}}$ (Wageningen, the Netherlands). The enriched libraries of Golden Delicious (ATC) and Florina (AAG and AAC) were developed by Ecogenics GmbH (Zürich, Switzerland) from size-selected digested (MboI for AAG and AAC libraries and Tsp509I for the ATC library) genomic DNA ligated to adaptors and enriched by magnetic bead selection with biotin-labeled corresponding oligonucleotide repeats (Gautschi et al. 2000). DNA fragments were PCR-amplified with the corresponding primer (Table 1). The PCR products of the ATC library were cloned into the vector Торо® (Invitrogen) and transformed in the TOP10 F' competent cells (Invitrogen), while the AAC and AAG libraries were cloned in the vector pDrive (Qiagen) and transformed in the EZ cells (Qiagen). Recombinant cells were spotted over nylon membranes and hybridized with the corresponding SSR repeat. Positive clones were sequenced with the primer M13 reverse by Synergene Biotech GmbH (Zürich, Switzerland).

SSRs from publicly available ESTs
Malus sequences from the NCBI database (September 2003) (http://www.ncbi.nlm.nih.gov/) were examined for microsatellite repeats using the software Tandem Repeat Finder v 3.21 (Benson 1999). From these ESTs, a subset of sequences was selected that contained microsatellite repeats and in which the repeat was sufficiently far from the edge of the sequence to allow design of both forward and reverse PCR primers.

## SSRs from the literature

Eight apple SSRs with unknown map position (GD12, -15, $-96,-100,-103,-142,-147$, and -162 ; Hokanson et al. 1998) have been tested for polymorphism with and between Fiesta and Discovery as well as with the parents of the other mapping populations available. Polymorphic markers were screened over a segregating mapping population and mapped. In addition, the following 17 SSRs from a map of pear (Yamamoto et al. 2004) were examined: NB102a, NB106a, NB111a, NH020a, NH023a, NH029a, NH025a, NB113a, KA4b, BGT23b, HGA8b, NH002b, NH009b, NH004a, NH015a, NH033b, and MSS6 (Yamamoto et al. 2002a-c; Oddou-Muratorio et al. 2001). This SSRs were selected because they map at positions for which the homologous regions of the apple genome lacked or had only a few SSRs. After verification that they generated amplicons in apple, they were mapped in the Fiesta $\times$ Discovery population using a range of annealing temperatures.

## Primer design and PCR conditions

Primer pairs flanking the SSR sequence were designed with the program Primer3 (Rozen and Skaletsky 2000) publicly available at http://fokker.wi.mit.edu/primer3/. The ideal annealing temperature ( Tm ) of the primers was set at $60^{\circ} \mathrm{C}$. Some primers were pig-tailed (Brownstein et al. 1996), whereby a variable number of nucleotides was added to the $5^{\prime}$ end of the reverse primer to obtain the sequence GTTT. Primers were synthesized at Microsynth (Balgach, Switzerland). PCR amplification and tests of primers were performed, as described by Gianfranceschi et al. (1998), with the following modifications: the PCR

Table 1 Restriction enzymes, adaptor sequences, and primers used for the construction of the AAG, AAC, and ATC SSR libraries

| Library | Restr. <br> enzyme | Adaptor (seq1) | Adaptor seq 2 (rev complementary of seq1) | PCR primer |
| :--- | :--- | :--- | :--- | :--- | :--- |
| AAG | MboI | SAULA: 5'GCGGTACCCGGGAAGCTTGG3' | SAULB: 5'GATCCCAAGCTTCCCGGGTACCGC3' | SAULA |
| AAC | MboI | SAULA: 5'GCGGTACCCGGGAAGCTTGG3' | SAULB: 5'GATCCCAAGCTTCCCGGGTACCGC3' | SAULA |
| ATC | Tsp509I | TSPAdShort | TSPAdLong | TSPAdShort |
|  |  | 5'TCGGAATTCTGGACTCAGTGCCAATT3' | 5'AATTGGCACTGAGTCCAGAATTCCGA3' $^{\prime}$ |  |

volume was reduced in some cases from 15 to $10 \mu \mathrm{l}$; $0.07 \mathrm{U} / \mu \mathrm{l}$ reaction of $T a q$ polymerase (New England BioLabs) was used, and the amplification profile was simplified to an initial denaturation at $94^{\circ} \mathrm{C}$ for 2 min 30 s followed by 35 cycles of $94^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 60^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 72^{\circ} \mathrm{C}$ for 1 min , both for regular (preliminary test of the primers) and with ${ }^{33} \mathrm{P}$-labeled forward primers. Before loading, radiolabeled PCR products were denatured by the addition of one volume of denaturing gel-loading buffer (Sambrook et al. 1989) and heating at $94^{\circ} \mathrm{C}$ for 5 min .

SSR alleles were analyzed by running ${ }^{33} \mathrm{P}$-labeled PCR products on a $6 \%$ denaturing sequencing gel (National Diagnostic, Atlanta, USA) in $1 \times$ TBE buffer using a IBI STS45i DNA sequencing unit. Data scoring was performed, as described by Liebhard et al. (2002), by comparison of the allele size of the nine cultivars with the ${ }^{33} \mathrm{P}$-labeled size standard 30-330 bp (Invitrogen).

## Nomenclature for SSR markers

SSR markers developed from enriched genomic libraries were prefixed with 'Hi' (indicating the HiDRAS project, Gianfranceschi and Soglio 2004) followed by a combination of two two-digit numbers, separated by a lower-case letter. The first two-digit number indicates the number of the sequencing plate, the letter and and second two-digit number indicate the well of this plate from which the DNA sequence was obtained. SSR markers developed from EST sequences were designated with the GenBank accession number followed by the subscript "SSR". Names of the SSRs from the literature were not changed.

## Marker quality

The new SSR markers were divided into four quality classes based on the type and number of additional, non-SSR containing amplicons that appeared on the gels: (1) clean: no extra band, (2) complementary band (s): all additional bands are complementary; this is at a constant distance to (specific) SSR bands, (3) extra bands: non-complementary bands are present that may hamper multiplexing, (4) dirty: many additional, non-complementary bands that hamper scoring of the true SSR bands, not suitable for use in multiplex reactions.

## Genetic mapping

The segregation data for the 44 progeny plants genotyped with the new markers were added to the data previously used to develop the "reference maps" of Fiesta and Discovery (Liebhard et al. 2003b). As these reference maps were based on 251 progeny plants, missing data were assigned to the 207 progeny plants not analyzed with the new SSRs. Mapping of the SSRs was performed with JoinMap ${ }^{\text {TM }}$ version 2.0 (Stam and Van Ooijen 1995) in
connection with JMDesk 3.6 provided by Dr. B. Koller (Ecogenics GmbH, Switzerland). A LOD score of 5 was used to assign markers to linkage groups. Mapping of the SSRs was considered correct (also third-round maps) if the introduction of the new data did not change, or only slightly changed, distances between markers or orders among the flanking markers. Drawings of the linkage maps were generated with MapChart (Voorrips 2001). Some SSR markers, which showed no polymorphism in the Fiesta $\times$ Discovery progeny could be mapped in other crosses. One of the following three crosses was used in such cases: Discovery $\times$ TN10-8, Durello di Forlì $\times$ Fiesta, or Fuji $\times$ Mondial Gala, and the map position was estimated by manual alignment.

## Results

One hundred and forty-eight new SSR markers have been developed and mapped, 117 from genomic libraries (65 from dinucleotide repeat libraries, 52 from trinucleotide repeat libraries) and 31 from Malus sequences from GenBank ( 24 containing dinucleotide repeats and seven with larger repeats). Moreover, it was possible to determine the location of previously published SSRs of apple (GD 147), pear (HGA8b, KA4b, NB102a, NH009b, NH029a, and NH033b) and of a Sorbus torminalis SSR (MSS6). For all 156 SSRs details of the forward and reverse primers, nucleotides added to the reverse primer to build a pigtail, repeat sequence, repeat type, size-range, number of alleles for the set of nine cultivars (polymorphism level), map position (linkage group), and the quality of the markers is presented (Table 2). For the EST sequences used to develop EST-SSRs, their deduced functions and origins (tissues) have also been indicated (Table 3).

In addition, the allele composition of nine cultivars has been determined to allow the estimation of the level of polymorphism of the marker (Table 4). However, in the presence of only a single allele, it usually remained unclear whether this was due to the presence of an allele at the homozygous state or to the presence of one amplified allele and a null allele (Table 4). These two options could be distinguished for Fiesta and Discovery based on the segregation patterns in the mapping population. For other cultivars, this has still to be clarified and so a single value has been entered (Table 4). For some SSRs, mainly containing trinucleotide repeats, it was not always possible to distinguish SSR amplicons from other PCR products due to lack of stutter bands. In such cases, the size of all amplicons is reported in Table 4. This may have led to an overestimation of the level of polymorphism of these SSRs.

## Efficiency of SSR development

The repeat type of the new SSR sequences (2 nt or more) greatly affected the success of development of polymorphic markers that could be mapped. Sixty-one to sixtythree percent of 2-nt repeats and only $33-40 \%$ of the SSRs
Table 2 Names, primers, pigtail sequence, repeat type, repeat sequence, range and number of alleles found in nine diploid cultivars, type of marker, map position and quality of the new set of SSR markers developed

| SSR name | Forward primer | Reverse primer | Pigtail $s e q^{a}$ | Repeat type ${ }^{\text {b }}$ | Repeat seq | Allele range | No. of alleles | Type of marker ${ }^{\mathrm{c}}$ |  | Quality ${ }^{\text {d }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hi01a03 | CGAATGAAATGTCTAAACAGGC | AAGCTACAGGCTTGTTGATAACG | - | Perf | AAG | 168-193 | 3 | SL | 10 | Clean |
| Hi01a08 | AAGTCCAATCGCACTCACG | CGTAGCTCTCTCCCGATACG | - | Comp | AAG-GA | 177-177 | 2 | SL | 16 | Clean |
| Hi01b01 | GCTACAGGCTTGTTGATAACGC | ACGAATGAAATGTCTAAACAGGC | - | Perf | AAG | 153-189 | 5 | SL | 10 | Compl. b |
| Hi01c04 | GCTGCCGTTGACGTTAGAG | GTTTGTAGAAGTGGCGTTTGAGG | GTT | Imp | GA | 214-232 | 4 | SL | 5 | Compl. b |
| Hi01c06 | TTAGCCCGTATTTGGACCAG | GTTTCACCTACACACACGCATGG | GTTT | Imp | GT | 128-144 | 3 | PML | 15 | Extra bands |
| Hi01c09 | AAAGGCGAGGGATAAGAAGC | GTTTGCACATTTGAGCTGTCAAGC | GTTT | Perf | GA | 214-218 | 3 | SL | 14 | Clean |
| Hi01c11 | TTGGGCCACTTCACAACAG | GTTTGAGTTTGATCTCCAACATTAC | GTTT | Imp | GT | 138-260 | 17 | ML | 8/16 | Clean |
| Hi01d01 | CTGAAATGGAAGGCTTGGAG | GTTTACCAATTAGGACTTAAAGCTG | GTTT | Comp | GA-GT | 191-222 | 5 | PML | 9 | Extra bands |
| Hi01d05 | GGTATCCTCTTCATCGCCTG | TTAGATTGACGTTCCGACCC | - | Imp | GA | 210->330 | 16 | PML | 7 | Extra bands |
| Hi01d06 | GGAGAGTTCCTGGGTTCCAC | AAGTGCACCCACACCCTTAC | - | Imp | GA | 115-165 | 11 | ML | 11/16 | Extra bands |
| Hi01e10 | TGGGCTTGTTTAGTGTGTCAG | GTTTGGCTAGTGATGGTGGAGGTG | GTTT | Perf | GA | 126-224 | 8 | SL | 4 | Compl. b |
| Hi02a03 | GACATGTGGTAGAACTCATCG | GTTTAGTGCGATTCATTTCCAAGG | GTTT | Perf | GA | 168-198 | 9 | PML | 5 | Extra bands |
| Hi02a07 | TTGAAGCTAGCATTTGCCTGT | TAGATTGCCCAAAGACTGGG | - | Imp | GA | 254-312 | 4 | SL | 2 | Extra bands |
| Hi02a09 | ATCTCTAAGGGCAGGCAGAC | CTGACTCTTTGGGAAGGGC | - | Imp | GA | 138-158 | 4 | SL | 11 | Clean |
| Hi02b07 | TGTGAGCCTCTCCTATTGGG | TGGCAGTCATCTAACCTCCC | - | Imp | GA | 204-216 | 4 | SL | 12 | Clean |
| Hi02b10 | TGTCTCAAGAACACAGCTATCACC | GTTTCTTGGAGGCAGTAGTGCAG | GTT | Perf | GA | 200-254 | 8 | PML | 16 | Clean |
| Hi02c06 | AGCAAGCGGTTGGAGAGA | GTTTGCAACAGGTGGACTTGCTCT | GTTT | Perf | GA | 208-252 | 8 | SL | 11 | Clean |
| Hi02c07 | AGAGCTACGGGGATCCAAAT | GTTTAAGCATCCCGATTGAAAGG | GTT | Perf | GA | 108-150 | 5 | PML | 1 | Clean |
| Hi02d02 | TTCCTAGGCTACCCGAAATATG | GTTTCTGGCATGGACATTCAACC | GTTT | Comp | GA-GT | 152-194 | 5 | SL | 15 | Clean |
| Hi02d04 | TGCTGAGTTGGCTAGAAGAGC | GTTTAAGTTCGCCAACATCGTCTC | GTTT | Perf | GA | 224-250 | 10 | SL | 10 | Clean |
| Hi02d05 | GAGGGAGAATCGGTGCATAG | CATCCCTCAGACCCTCATTG | - | Perf | GA | 153-205 | 7 | SL | 12 | Extra bands |
| Hi02d11 | GCAATGTTGTGGGTGACAAG | GTTTGCAGAATCAAAACCAAGCAAG | GTTT | Imp | GA | 198-262 | 8 | SL | 14 | Extra bands |
| Hi02f06 | TAAATACGAGTGCCTCGGTG | GCAGTTGAAGCTGGGATTG | GTTT | Perf | GA | 204-228 | 6 | SL | 15 | Clean |
| Hi02f12 | ACATGGCCGAAGACAATGAC | GTTTCAACCTTTATCCCTCCATCTTTC | GTTT | Perf | GA | 130-150 | 6 | SL | 17 | Clean |
| Hi02g06 | AGATAGGTTTCACCGTCTCAGC | GACCTCTTTGGTGCGTCTG | - | Comp | GA-CAC | 149-163 | 4 | SL | 15 | Clean |
| Hi02h08 | GCCACTCATACCCATCGTATTG | GTTTGGCTGGGAATATATGATCAGGTG | GTTT | Comp | GT-GA | 170-200 | 6 | SL | 16 | Clean |
| Hi03a03 | ACACTTCCGGATTTCTGCTC | GTTTGTTGCTGTTGGATTATGCC | GTT | Perf | GA | 160-228 | 10 | PML | 6 | Clean |
| Hi03a06 | TGGTGAGAGAAGGTGACAGG | GTTTAAGGCCGGGATTATTAGTCG | GTTT | Imp | GA | 158-197 | 5 | PML | 15 | Dirty |
| Hi03a10 | GGACCTGCTTCCCCTTATTC | GTTTCAGGGAACTTGTTTGATGG | GT | Imp | GA | 206-290 | 6 | SL | 7 | Clean |
| Hi03b03 | TGAATTGAGTTTGAGAATGGAATG | GTTTGTCAGGACGGGTAATCAAGG | GTTT | Perf | GA | 196-212 | 7 | SL | 12 | Clean |
| Hi03c04 | CGTAAATAGCGAATCCGATACC | GTTTCAACATCTGTGGGTCATTGC | GTTT | Perf | GA | 169-257 | 6 | SL | 10 | Clean |
| Hi03c05 | GAAGAGAGAGGCCATGATAC | GTTTAACTGAAACTTCAATCTAGG | GTT | Imp | GA | 179-221 | 8 | SL | 17 | Clean |
| Hi03d06 | TCATGGATCATTTCGGCTAA | GTTTGCCAATTTTATCCAGGTTGC | GTTT | Perf | GA | 115-169 | 8 | SL | 3 | Clean |
| Hi03e03 | ACGGGTGAGACTCCTTGTTG | GTTTAACAGCGGGAGATCAAGAAC | GTTT | Perf | GA | 187-199 | 6 | SL | 3 | Clean |
| Hi03e04 | CTTCACACCGTTTGGACCTC | GTTTCATATCCCACCACCACAGAAG | GTTT | Imp | GA | 132-160 | 6 | SL | 13 | Clean |
| Hi03f06 | ACGATTTGGTGATCCGATTC | GTTTCGTCGCATTGTGCTTCAC | GTTT | Perf | GA | 153-217 | 9 | PML | 10 | Extra bands |
| Hi03g06 | TGCCAATACTCCCTCATTTACC | GTTTAAACAGAACTGCACCACATCC | GTTT | Perf | GA | 182-204 | 5 | SL | 15 | Clean |
| Hi04a02 | TTCGTGGAAACCTAATTGCAG | GTTTCCTCTGCTTCTTCATCTTTGC | GTTT | Perf | GA | 82-104 | 6 | SL | 13 | Clean |

Table 2 (continued)

| SSR name | Forward primer | Reverse primer | Pigtail seq $^{a}$ | Repeat type ${ }^{\text {b }}$ | Repeat seq | Allele range | No. of alleles | Type of marker ${ }^{\text {c }}$ |  | Quality ${ }^{\text {d }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hi04a05 | GGCAGCAGGGATGTATTCTG | GTTTCATGTCAAATCCGATCATCAC | GTTT | Perf | GA | 194-222 | 8 | SL | 9 | Clean |
| 88 | TTGAAGGAGTTTCCGGTTTG | GTTTCACTCTGTGCTGGATTATGC | GTT | Perf | GA | 211-250 | 7 | SL | 5 | Clean |
| Hi04b12 | CCCAAACTCCCAACAAAGC | GTTTGAGCAGAGGTTGCTGTTGC | GTTT | Perf | GA | 140-156 | 5 | SL | 8 | Clean |
| Hi04c05 | AGGATGCTCTGCCTGTCTTC | GTTTCTCACTCGCCTGCTCTATCC | GTTT | Perf | GA | 179-183 | 3 | SL | 15 | Clean |
| Hi04c10 | TGCGCATTTGATAGAGAGAGAA | GTTTAACAAAGAACGACCCACCTG | GTTT | Perf | GA | 172-238 | 11 | ML | 3/4 | Dirty |
| Hi04d02 | TTCGTGGCTGAGAAAGGAGT | GTTTGTACGGTGCATTGTGAAAG | GT | Perf | GA | 176-238 | 15 | PML | 5 | Extra bands |
| Hi04d10 | AAATTCCCACTCCTCCCTGT | GTTTGAGACGGATTGGGGTAG | G | Perf | GA | 164-182 | 4 | SL | 6 | Extra bands |
| Hi04e04 | GACCACGAAGCGCTGTTAAG | GTTTCGGTAATTCCTTCCATCTTG | GTT | Perf | GA | 216-246 | 6 | SL | 16 | Clean |
| Hi04e05 | AAGGGTGTTTGCGGAGTTAG | GGTGCGCTGTCTTCCATAAA | - | Perf | GA | 144-144 | 2 | SL | 8 | Clean |
| Hi0408 | CGTGAAAACTCTAACTCTCC | GTTTGAAAAGCGCATCAAAGTTCC | GTTT | Perf | GA | 218-226 | 4 | SL | 10 | Clean |
| Hi0409 | ACTGGGTGGCTTGATTTGAG | GTTTCAACTCACACCCTCTACATGC | GTTT | Imp | GA | 222-260 | 11 | PML | 13 | Compl. b |
| Hi04g05 | CTGAAACAGGAAACCAATGC | GTtTCGTAGAAGCATCGTTGCAG | GTT | Perf | GA | 190-258 | 9 | PML | 13 | Clean |
| Hi04g11 | CAGAGGATTATCAATTGGACGC | AAACTATCTCCAGTTATCCTGCTTC | - | Perf | GA | 118-164 | 5 | SL | 11 | Clean |
| Hi05b02 | GATGCGGTTTGACTTGCTTC | GTTTCTCCAGCTCCCATAGATTGC | GTT | Perf | GA | 120-178 | 7 | PML | 10 | Clean |
| Hi05b09 | AAACCCAACCCAAAGAGTGG | GTTTCTAACGTGCGCCTAACGTG | GTTT | Perf | GA | 136-144 | 3 | SL | 7 | Clean |
| Hi05c06 | tgcgighatg ittgattita | TGTTTTCTTTGGTTTTAGTTGGTG | - | Comp | GA-GT | 136-142 | 5 | PML | 17 | Dirty |
| Hi05d10 | AATGGGTGGTTTGGGCTTA | GTTTCTTTGGCTATTAGGCCTGC | GTT | Imp | GT | 212-212 | 2 | SL | 6 | Dirty |
| Hi05e07 | CCCAAGTCCCTATCCCTCTC | GTTTATGGTGATGGTGTGAACGTG | GTTT | Perf | GT | 214-234 | 7 | SL | 9 | Extra bands |
| Hi05f12 | TTTGGGTTTGGGTAGGTTAGG | GTTTGTGCAGCGCATGCTAATG | GTTT | Comp | TA-CA | 157-177 | 5 | ML | 12/3 | Dirty |
| Hi05g12 | TCTCTAGCATCCATTGCTTCTG | GTTTGTGTGTTCTCTCATCGGATTC | GTTT | Imp | GT | 208-288 | 10 | PML | 2 | Extra bands |
| Hi06b06 | GGTGGGATTGTGGTTACTGG | GTTTCATCGTCGGCAAGAACTAGAG | GTTT | Imp | GT | 236-262 | 4 | SL | 11 | Extra bands |
| Hi06f09 | AACCAAGGAACCCACATCAG | GTTTCACTTACACACGCACACACG | GTTT | Imp | GT | 272-288 | 4 | PML | 15 | Dirty |
| Hi07b02 | ATtTGGGGTtTCAACAATGG | GTTTCGGACATCAAACAAATGTGC | GTTT | Imp | GT | 212-218 | 5 | PML | 4 | Clean |
| Hi07b06 | AGCTGCAGGTAGAGTTCCAAG | GTTTCATTACCATTACACGTACAGC | GTTT | Imp | GT | 220-226 | 4 | SL | 6 | Clean |
| Hi07d08 | TGACATGCTTTTAGAGGTGGAC | GTTTGAGGGGTGTCCGTACAAG | GT | Perf | CA | 222-232 | 3 | SL | 1 | Extra bands |
| Hi07d11 | CCTTAGGGCCTTTGTGGTAAG | GTTTGAGCCGATTAGGGTTTAGGG | GTTT | Imp | GT | 200-234 | 11 | ML | 11/16 | Clean |
| Hi07d12 | GGAATGAGGGAGAAGGAAGTG | GTTTCCTCTTCACGTGGGATGTACC | GTTT | Imp | GT | 184-250 | 8 | ML | 2/7 | Clean |
| Hi07e08 | TTCGTGCTAGGGAGTTGTAGC | GTTTGCCTCCATAGGATTATTTGAC | GTT | Perf | GT | 208-241 | 9 | ML | 8/3 | Clean |
| Hi07f01 | GGAGGGCTTTAGTTGGGAAC | GTTTGAGCTCCACTTCCAACTCC | GTT | Comp | AT-GT | 204-220 | 5 | SL | 12 | Extra bands |
| Hi07g10 | TATTGGGTtTTGGGTTTGGA | GTTTCAACCCTTTTGGTTGTGAGG | GTTT | Imp | GT | 126-128 | 3 | PML | 11 | Dirty |
| Hi07h02 | CAAATTGGCAACTGGGTCTG | GTTTAGGTGGAGGTGAAGGGATG | GTT | Perf | GT | 246-276 | 10 | SL | 17 | Clean |
| Hi08a04 | TTGTCCTTCTGTGGTTGCAG | GTTTGAAGGTAAGGGCATTGTGG | GTT | Comp | GAA-GT | 246-254 | 4 | SL | 5 | Extra bands |
| Hi08c05 | TCATATAGCCGACCCCACTTAG | GTTTCACACTCCAAGATTGCATACG | GTTT | Perf | AAC | 230-240 | 3 | PML | 14 | Extra bands |
| Hi08d09 | AACGGCTTCTTGTCAACACC | GTTTACTGCATCCCTTACCACCAC | GTTT | Perf | AAC | 183-186 | 2 | SL | 16 | Clean |
| Hi08e04 | GCATGGTGGCCTTTCTAAG | GTTTACCCTCTGACTCAACCCAAC | GTTT | Perf | AAG | 201-234 | 6 | PML | 4 | Extra bands |
| Hi08e06 | GCAATGGCGTTCTAGGATTC | GTTTGGCTGCTTGGAGATGTG | GT | Perf | AAC | 134-138 | 4 | SL | 13 | Extra bands |
| Hi08f05 | GTGTGGGCGATTCTAACTGC | GTtTCCTTTATTCTAAACATGC | GTTT | Perf | AAG | 165-165 | 2 | SL | 2 | Clean |
|  |  | CACGTC |  |  |  |  |  |  |  |  |

Table 2 (continued)
$\left.\begin{array}{llllllllllll}\hline \text { SSR name } & \text { Forward primer } & \text { Reverse primer } & \begin{array}{l}\text { Pigtail } \\ \text { seq }^{\text {a }}\end{array} & \begin{array}{l}\text { Repeat } \\ \text { type }\end{array} & \text { Repeat } & \text { seq } & \begin{array}{l}\text { Allele } \\ \text { range }\end{array} & \begin{array}{l}\text { No. of } \\ \text { alleles }\end{array} & \begin{array}{l}\text { Type of } \\ \text { marker }\end{array} \\ & & & & \text { Ls) } \\ \text { (s) }\end{array}\right]$
Table 2 (continued)

| SSR name | Forward primer | Reverse primer | Pigtail $s e q^{\text {a }}$ | Repeat type ${ }^{\text {b }}$ | Repeat seq | Allele range | No. of alleles | Type of marker ${ }^{\text {c }}$ |  | Quality ${ }^{\text {d }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hi23g02 | TTTTCCAGGATATACTACCCTTCC | GTTTCTTCGAGGTCAGGGTTTG | GT | Perf | AAC | 230-257 | 6 | SL | 4 | Compl. b |
| Hi23g08 | AGCCGTTTCCCTCCGTTT | GTTTGTGGATGAGAAGCACAGTCA | GTT | Perf | CTT | 211-220 | 3 | SL | 4 | Clean |
| Hi23g12 | CCCTTCCCTACCAAATGGAC | GTTTAAAGGGGCCCACAAAGTG | GTTT | Comp | $\begin{aligned} & \text { CAA-AGC- } \\ & \text { CCA } \end{aligned}$ | 223-241 | 4 | PML | 8 | Clean |
| Hi24f04 | CCGACGGCTCAAAGACAAC | TGAAAAGTGAAGGGAATGGAAG | - | Imp | AAG | 144-153 | 5 | SL | 2 | Clean |
| AF057134 ${ }_{\text {SSR }}$ | ACTACCCAATGCCCACAAAG | TATCCTCGCCCAAAAGACTG | - | Perf | GA | 202-224 | 7 | SL | 10 | Compl. b |
| AF527800 ${ }_{\text {SSR }}$ | TGGAAAGGGTTGATTGACCT | AACAGCGGGTGGTAAATCTC | - | Perf | GA | 168-194 | 5 | SL | 17 | Compl. b |
| AJ000761a $\mathrm{SSSR}^{\text {e }}$ | CTGGGTGGATGCTTTGACTT | TCAATGACATTAATTCAACTTACAAAA | - | Imp | TA | 260-272 | 5 | SL | 14 | Clean |
| AJ000761b ${ }_{\text {SSR }}$ | CCCTAAACACACAGCCTCCT | GTTTCAGCATCGCAGAGAACTGAG | GTTT | Perf | GA | 210-208 | 8 | SL | 14 | Clean |
| AJ001681 ${ }_{\text {SSR }}$ | CCTGAGGTTATTGACCCAAAA | CACTCAGTTGGAAAACCCTACA | - | Perf | GA | 169-195 | 6 | SL | 17 | Clean |
| AJ251116 ${ }_{\text {SSR }}$ | GATCAGAAAATTGCTAGGAAAAGG | AGAGAACGGTGAGCTCCTGA | - | Perf | GA | 165-167 | 2 | SL | 2 | Compl. b |
| AJ320188 ${ }_{\text {SSR }}$ | AACGATGCTTGAGGAAGAACA | GCTTAACAGAAACATCGCTGA | - | Perf | GA | 191-245 | 8 | SL | 9 | Clean |
| AT000174 ${ }_{\text {SSR }}$ | CGGAGGCCGCTATAATTAGG | CCTGGAAAGAAAGTAAAAGGACA | - | Comp | TA-GTA-GT | 178-200 | 6 | PML | 17 | Extra bands |
| AT000400 ${ }_{\text {SSR }}$ | CTCCCTTTGCTCCCTCTCTT | AGGATGTCAGGGTTGTACGG | - | Imp | CAG | 198-232 | 7 | SL | 2 | Extra bands |
| AT000420 $0_{\text {SSR }}$ | TTGGACCAATTATCTCTGCTATT | GATGTGGTCAGGGAGAGGAG | - | Imp | GA | 189-209 | 5 | SL | 4 | Extra bands |
| AU223486 ${ }_{\text {SSR }}$ | TGACTCCATGGTTTCAGACG | AGCAATTCCTCCTCCTCCTC | - | Comp | GAA-GA | 205-217 | 3 | SL | 13 | Extra bands |
| AU223548 ${ }_{\text {SSR }}$ | ACCACCACTGCAGAGACTCA | GACGCACCCATTCATCTTTT | - | Perf | GGA | 262-278 | 4 | SL | 10 | Extra bands |
| AU223657 ${ }_{\text {SSR }}$ | TTCTCCGTCCCCTTCAACTA | CACCTTGAGGCCTCTGTAGC | - | Imp | GA | 219-233 | 6 | SL | 3 | Clean |
| AU223670 SSR | GGACTCAATGCCTTTTCTGG | AGGATGGCAGCAATCTTGAA | - | Perf | ACC | 194-202 | 3 | PML | 5 | Extra bands |
| AU301431 ${ }_{\text {SSR }}$ | TСТTССТССТССТССТССТС | TCTTTTTCTTGGGGTCTTGG | - | Perf | AAG | 213-216 | 2 | SL | 16 | Extra bands |
| AY187627 ${ }_{\text {SSR }}$ | GAGGACTGAATTGGTTGAGGTC | GTTTCTCACCCGTATATAGGCCAAC | GTTT | Comp | GT-TA | 300-300 | 2 | SL | 17 | Clean |
| CN444542 ${ }_{\text {SSR }}$ | ATAAGCCAGGCCACCAAATC | GTTTGCAGTGGATTGATGTTCC | GT | Perf | GA | 110-156 | 8 | PML | 9 | Clean |
| CN444636 ${ }_{\text {SSR }}$ | CACCACTTGAGTAATCGTAAGAGC | GTTTGCCAGTTAAGGACCACAAGG | GTTT | Comp | AT-GT | 239-243 | 3 | SL | 2 | Clean |
| CN444794 ${ }_{\text {SSR }}$ | CATGGCAGGTGCTAAACTTG | GTTTGCAACTCACACAATGCAAC | GTT | Perf | TA | 230-306 | 8 | PML | 7 | Extra bands |
| CN445290 ${ }_{\text {SSR }}$ | TCTCAGTTGCTCTGGCTTTG | GTTTGCAATCAATGCCACTCTTC | GTT | Perf | GA | 230-242 | 3 | SL | 6 | Clean |
| CN445599 ${ }_{\text {SSR }}$ | TCAAATGGGTTCGATCTTCAC | GTTTGCCTGGCTGTAACTGTTTGG | GTTT | Perf | TA | 130-176 | 11 | PML | 5 | Extra bands |
| CN491050 ${ }_{\text {SSR }}$ | CGCTGATGCGATAATCAATG | GTTTCACCCACAGAATCACCAGA | GTT | Perf | GA | >330->330 | 3 | SL | 11 | Clean |
| CN493139 ${ }_{\text {SSR }}$ | CACGACCTCCAAACCTATGC | GTTTATGAAAGTACGGCACCCATC | GTTT | Perf | TA | 124-162 | 10 | PML | 2 | Clean |
| CN496002 ${ }_{\text {SSR }}$ | TCAGAATCTCAAGCAAGATCCTC | GTTTGATTGATCGTGGCGATATG | GTT | Comp | TA-CTT | 243-261 | 4 | SL | 5 | Clean |
| CN496913 ${ }_{\text {SSR }}$ | TGCCTTTGAGAATCGAAATG | TGTTTGTCAATTTCTTGGAACTC | - | Perf | TA | 236-278 | 5 | SL | 12 | Clean |
| CN581493 ${ }_{\text {SSR }}$ | GCTTTTCATGGTGGAAAAACTG | GTTTGACTCTCCGCTCTGATGGAC | GTTT | Perf | TA | 184-228 | 8 | SL | 2 | Clean |
| U78948 ${ }_{\text {SSR }}$ | GATCGTCCGCCACCTTAAT | AGGGTTTTCATCATGCACATT | - | Imp | TA | 178-190 | 7 | SL | 14 | Extra bands |
| U78949 SSR | TTTGTCTACCTCTGATCTTAACCAA | CAGCATCGCAGAGAACTGAG | - | Perf | GA | 172-225 | 12 | ML | 6/14 | Extra bands |
| Z38126 ${ }_{\text {SSR }}$ | AAGAGGGTGTTCCCAGATCC | TGTTCGATGTGACTTCAATGC | - | Perf | TA | 214-240 | 3 | SL | 7 | Clean |
| Z71980 ${ }_{\text {SSR }}$ | TCTTTCTCTGAAGCTCTCATCTTTC | GGACATGGATGAAGAATTGGA | - | Imp | TA | 170-172 | 2 | SL | 8 | Clean |
| Z71981 $1_{\text {SSR }}$ | GCACTTACCTTTGTTGGGTCA | CCGGCATTCCAAATGTAACT | - | Perf | AAG | 212-232 | 5 | SL | 15 | Clean |
| GD147 | TCCCGCCATTTCTCTGC | GTTTAAACCGCTGCTGCTGAAC | GTTT | u | GA | 135-155 | 6 | SL | 13 | Dirty |

Table 2 (continued)

| SSR name | Forward primer | Reverse primer | Pigtail $s e q^{a}$ | Repeat type ${ }^{\text {b }}$ | Repeat seq | Allele range | No. of alleles | Type of marker ${ }^{\mathrm{c}}$ | $\begin{aligned} & \text { LG } \\ & \text { (s) } \end{aligned}$ | Quality ${ }^{\text {d }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| HGA8b | AACAAGCAAAGGCAGAACAA | CATAGAGAAAGCAAAGCAAA (Tm $55^{\circ} \mathrm{C}$ ) | - | Comp | GA-GCTTT | 133-164 | 6 | ML | 3/11 | Dirty |
| KA4b | AAAGGTCTCTCTCACTGTCT | CCTCAGCCCAACTCAAAGCC $\left(\operatorname{Tm} 55^{\circ} \mathrm{C}\right)$ | - | Imp | GT | 137-141 | 3 | SL | 1 | Compl. b |
| NB102a | TGTTATCACCTGAGCTACTGCC | CTTCCTCTTTATTTGCCGTCTT | - | Perf | GA | 181-183 | 3 | PML | 16 | Extra bands |
| NH009b | CCGAGCACTACCATTGA | CGTCTGTTTACCGCTTCT | - | Perf | GA | 138-162 | 6 | SL | 13 | Extra bands |
| NH029a | GAAGAAAACCAGAGCAGGGCA | CCTCCCGTCTCCCACCATATTAG $\left(\operatorname{Tm} 55^{\circ} \mathrm{C}\right)$ | - | Perf | GA | 91-99 | 5 | SL | 9 | Compl. b |
| NH033b | GTCTGAAACAAAAAGCATCGCAA | CTGCCTCGTCTTCCTCCTTATCTCC | - | Perf | GA | 163-189 | 7 | SL | 2 | Clean |
| MSS6 | CGAAACTCAAAAACGAAATCAA | ACGGGAGAGAAACTCAAGACC | - | u | GT | 273-279 | 3 | SL | 4 | Extra bands |

[^0]with 3-nt repeats (or more) could be mapped (Table 5). The origin of the sequence (enriched library or GenBank accession) did not affect the success rate (Table 5). The repeat type similarly affected the observed allelic diversity of the markers, being highest for the 2-nt repeats (Table 5). However, if the average number of repeats found in the sequence used for the development of the SSRs is considered, it can be affirmed that the higher allelic variation observed for the 2-nt containing SSRs is probably due to the higher number of repeats. In fact, sequences from 2-nt repeats genomic libraries and ESTs contain on average 28 and 14 repeats, respectively, while 3-nt repeat genomic libraries and ESTs contain nine and six repeats, respectively (Table 5).

A relatively high number of SSRs (41\%) developed for pear and Sorbus torminalis were transferred to the apple. Surprisingly, only one out of eight previously published apple SSRs (GD series) could be mapped (Table 5).

To be efficiently used in genotyping projects, SSRs need to be sufficiently polymorphic and easy to score. If the threshold to declare a single-locus SSR as sufficiently polymorphic within our current set of nine cultivars is set at five alleles, $62 \%$ of the $2-n t$ repeat SSRs and $24 \% \geq 3-\mathrm{nt}$ repeat SSRs can be considered (Table 6). The two most polymorphic single-locus SSRs were $\mathrm{Hi02d04}$ and Hi07h02, both being 2-nt repeat SSR with ten alleles in the set of nine cultivars.

Approximately 58 and $10 \%$ of the new SSRs have been classified as being "clean" or showing "complementary bands" (amplification of amplicons at a constant distance from the SSR allele), respectively. On the other hand, 26 and $6 \%$ of the new markers have been classified as "extra bands" or "dirty", respectively. Under our PCR conditions, these SSRs amplify several additional non-SSR amplification products, which make them unsuitable for highthroughput genotyping. The SSR amplicons of the markers classified as "extra bands" are clearly visible, while those of class "dirty" may be difficult to recognize. Improvement of these SSRs by the design and testing of alternative primers was not pursued unless they are shown to be located in regions of high interest in which no other, high quality SSRs are located.

## Genetic mapping

One hundred and forty-eight SSRs out of 156 have been mapped on the reference map derived from the cross Fiesta $\times$ Discovery (Liebhard et al. 2003b). The remaining eight were mapped in other crosses: four in Discovery $\times$ TN10-8 (Hi23b12, Hi01c09, Hi09f01, Hi08f05), three in Durello di Forlì $\times$ Fiesta (Hi03c05, Hi08d09, Z71980 ${ }_{\text {SSR }}$ ) and one in Fuji $\times$ Mondial Gala (Hi11a01) (Fig. 1).

The 156 new SSR primer pairs enriched the reference map with 168 new loci ( 12 primer pairs amplified two loci that could both be mapped). The linkage groups (LGs) with highest increase in loci are LG 16 with 16 loci and LGs 10 and 5 with 14 loci each, followed by LGs 2, 11, and 15 with 13 loci each. The LG with the lowest increase of loci is LG

Table 3 Description and putative functions of the mapped ESTs as found at NCBI GenBank

| SSR name ${ }^{\text {a }}$ | Definition |
| :---: | :---: |
| AF057134-SSR | Malus domestica NADP-dependent sorbitol 6-phosphate dehydrogenase (S6PDH) gene, complete cds |
| AF527800-SSR | Malus x domestica expansin 3 (EXP03) mRNA, complete cds |
| AJ000761a,b-SSR | Malus domestica mRNA for MADS-box protein, MADS7 |
| AJ001681-SSR | Malus domestica mRNA for MADS box protein MdMADS8 |
| AJ251116-SSR | Malus domestica mRNA for B-type MADS box protein (mads 13 gene) |
| AJ320188-SSR | Malus domestica mRNA for MADS box protein (MADS12A gene) |
| AT000174-SSR | AT000174 Apple young fruit cDNA library Malus x domestica cDNA clone af180, mRNA sequence |
| AT000400-SSR | AT000400 Apple peel cDNA library Malus x domestica cDNA clone ap 189, mRNA sequence |
| AT000420-SSR | AT000420 Apple peel cDNA library Malus x domestica cDNA clone ap 212 , mRNA sequence |
| AU223486-SSR | AU223486 Apple shoot cDNA library Malus x domestica cDNA clone S0016, mRNA sequence |
| AU223548-SSR | AU223548 Apple shoot cDNA library Malus x domestica cDNA clone S0279, mRNA sequence |
| AU223657-SSR | AU223657 Apple shoot cDNA library Malus x domestica cDNA clone S 0159 , mRNA sequence |
| AU223670-SSR | AU223670 Apple shoot cDNA library Malus x domestica cDNA clone S0086, mRNA sequence |
| AU301431-SSR | AU301431 Apple shoot cDNA library Malus x domestica cDNA clone S1069, mRNA sequence |
| AY187627-SSR | Malus x domestica S-RNase (S) gene, S9 allele, partial cds |
| CN444542-SSR | Mdfw2003g22.x1 Mdfw Malus x domestica cDNA clone Mdfw2003g22 3-similar to TR:Q9SSL1 Q9SSL1 F15H11.6 Hypothetical protein At1g70810 |
| CN444636-SSR | Mdfw2003101.x1 Mdfw Malus x domestica cDNA clone Mdfw2003101 3-similar to TR:O81077 O81077 PUTATIVE CYTOCHROME P450, mRNA sequence |
| CN444794-SSR | Mdfw2001i05.y1 Mdfw Malus x domestica cDNA clone Mdfw2001i05 5-, mRNA sequence |
| CN445290-SSR | Mdfw2002h21.y1 Mdfw Malus x domestica cDNA clone Mdfw 2002 h 215 -, mRNA sequence |
| CN445599-SSR | Mdfw2003f11.y1 Mdfw Malus x domestica cDNA clone Mdfw2003f11 5-similar to TR:O81062 O81062 T18E12.21 Hypothetical protein At2g03120 |
| CN491050-SSR | Mdfw2008p11.y1 Mdfw Malus x domestica cDNA clone Mdfw2008p11 5-, mRNA sequence |
| CN493139-SSR | Mdfw2012f06.y1 Mdfw Malus x domestica cDNA clone Mdfw2012f06 5-similar to TR:O81808 O81808 HYPOTHETICAL 62.6 KD PROTEIN, mRNA sequence |
| CN496002-SSR | Mdfw2021d09.y1 Mdfw Malus x domestica cDNA clone Mdfw2021d09 5-similar to TR:O23131 O23131 HYPOTHETICAL 37.1 KD PROTEIN, mRNA sequence |
| CN496913-SSR | Mdfw2023a24.y1 Mdfw Malus x domestica cDNA clone Mdfw2023a24 5-, mRNA sequence |
| CN581493-SSR | Mdfw2039o14.y1 Mdfw Malus x domestica cDNA clone Mdfw2039o14 5-similar to TR:Q9ZQF5 Q9ZQF5 PUTATIVE RING-H2 FINGER PROTEIN, mRNA sequence |
| U78948-SSR | Malus domestica MADS-box protein 2 mRNA , complete cds |
| U78949-SSR | Malus domestica MADS-box protein 3 mRNA , complete cds |
| Z38126-SSR | M. domestica gene for calmodulin-binding protein kinase |
| Z71980-SSR | M. domestica mRNA for knotted1-like homeobox protein |
| Z71981-SSR | M. domestica partial gene for kn1-like protein |

${ }^{\text {a }}$ The first part of the SSR name corresponds to sequence accession number

1 with only five new loci. The largest distances between two flanking SSRs are on Discovery $6(36.2 \mathrm{cM})$ and Fiesta $7(37.5 \mathrm{cM})$. The maps of Fiesta and Discovery have been enriched by 99 and 115 loci, respectively ( 54 loci in common). The maps now span a total of $1,145.3 \mathrm{cM}$ (Fiesta) and $1,417.1 \mathrm{cM}$ (Discovery). This corresponds to an increase of 1.5 cM for the map of Fiesta and a decrease of 37.5 cM for the map of Discovery compared with the maps of Liebhard et al. (2003b). Most of the reduction of the total length of the map of Discovery is due to the splitting of its LG3 (reduction of 10 cM ). Also, the average chromosome lengths of Fiesta and Discovery did not change substantially, being 67.4 cM (previously 67.4 cM ) and 83.35 cM (previously 85.6 cM ), respectively. The fact that no substantial changes in the total length of the
parental maps are observed is an indication that the genome coverage is close to completion.

Selection of a genome covering set of apple SSRs
To facilitate efficient genome-wide mapping approaches, we aimed at developing a set of 100 SSRs that cover the entire apple genome. The high number of mapped SSRs in Fiesta $\times$ Discovery as well as in various other available mapping populations (data not shown) allowed the first design of such a set. Eighty-six SSRs were selected that span around $85 \%$ of the apple genome and that have an average distance between markers of 15 cM (Fig. 2). Regrettably, SSRs for 16 regions are not yet available. Out
Table 4 Allele composition of nine diploid cultivars

| SSR name | Fiesta | Discovery | Florina | Nova Easaygro | TN10-8 | Durello di Forlì | Prima | Mondial Gala | Fuji |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hi01a03 | 168:193 | 178:193 | 168 | nd | 193 nd | 193 nd | nd | 193 | nd |
| Hi01a08 | null | 177:null | null | null | null | 177 | null | 177 | null |
| Hi01b01 | 165:189 | 174:189 | 165 | 162153 | 153189 | 162189 | 153162 | 189 | nd |
| Hi01c04 | 218:222 | 222:232 | 218 | 214232 | 214218 | 214218222 | 218 | nd | nd |
| Hi01c06 | 142:142 | 128:142 | nd | 128 | 142 | 128142 | 144 | 142 | 142 |
| Hi01c09 | 216:218 | 216:218 | 216218 | 216 | 216 | 214216 | 216 | 216218 | 216218 |
| Hi01c11 | 216:220 ${ }^{\text {x }} 260: 260^{\text {a }}$ | null:null ${ }^{\text {a }} 234: 240^{\text {y }}$ | 138152202206 | 138146152206 | 138144146204 | 138148200204 | 138148176 | 138144152216 | 138148152 |
|  | 138200 | 138204 | 220 | 220 | 234240 | 240 | 220 |  | 218 |
| Hi01d01 | 199:220 | 195:220 | 199220 | 195 | 191195 | 195222 | 191 | 195199 | nd |
| Hi01d05 | $320:>330 \mathrm{a}$ | 320:null 210222 | 210222 | 212226242 | $210>330 \mathrm{~b}$ | 210242318 | 210222328 | 212222242300 | 210242 |
|  | 210242 | $244300>330$ b | > $330 \mathrm{a} / \mathrm{b}$ | >330 a/b |  |  |  | $320>330 \mathrm{c} / \mathrm{d}$ | $322>330 \mathrm{a} / \mathrm{b}$ |
| Hi01d06 | 125:133 | 124:131 | 155163 | 135165 | 137 | 137 | 137131124 | 115155 | 129155 |
| Hi01e10 | 220:220 | 214:126 | 220224 | 224 | 204208 | 182208 | 204212 | 224 | nd |
| Hi02a03 | 176:186 | 178:186 | 170176 | 168 | 170198 | 184188 | 176 | 174186 | 176186 |
| Hi02a07 | 254:282 | 280:280 | 312 | 312 | 254312 | 280 | 280 | 280 | 282 |
| Hi02a09 | 138 | 138:158 | 158 | 138 | 138158 | 138152 | 138 | 148 | nd |
| Hi02b07 | 214:null | null | 216 | 216 | 204216 | nd | 204216 | 204 | nd |
| Hi02b 10 | 228:228 ${ }^{\text {a }} 202: 202^{\text {a }}$ | 226:null 200:200 ${ }^{\text {a }}$ | 200226 | 200224 | 200226 | 226240 | 200202226 | 224226254 | 202224226 |
| Hi02c06 | 230:244 | 230:242 | 252 | 208242 | 242252 | 232 | 227242 | 212242 | nd |
| Hi02c07 | 116:150 | 150:150 | 116118 | 114118 | 116118 | 116 | 108118 | 116 | nd |
| Hi02d02 | null:null | 152:null | nd | 154194 | 156194 | 152 | 194 | nd | nd |
| Hi02d04 | 218:250 | 226:246 | 234 | 230240 | 234250 | 224232 | 240244 | 224 | nd |
| Hi02d05 | 153:197 | 153:205 | 153205 | 153175 | 153175 | 153173 | 153175 | 153191 | 195205 |
| Hi02d11 | 234:244 | 244:198 | 234258 | 254258 | 262 | 244198 | 254198 | 246258 | 248262 |
| Hi02f06 | 216:224 | 208:214 | 216228 | 208214 | 204228 | 214216 | 208228 | 216228 | nd |
| Hi02f12 | 130:132 | 140:140 | 134150 | 132 | 130150 | 138150 | nd | 132150 | nd |
| Hi02g06 | 161:161 | 161:163 | 161 | 161 | 161163 | 149155161 | 161163 | 161 | 161163 |
| Hi02h08 | 172:172 | 170:178 | 172 | 172 | 172 | 174200 | 172180 | 172 | 172 |
| Hi03a03 | 188:192 224 | 172:196 224228 | 160228 | 160224228 | 160172228 | 186194218228 | $\begin{aligned} & 186194218 \\ & 228 \end{aligned}$ | nd | nd |
| Hi03a06 | 158:160 178197 | 158:197 | 160178197 | 160178197 | 158160197 | nd | $\begin{aligned} & 160178183 \\ & 197 \end{aligned}$ | 158197 | nd |
| Hi03a10 | 216:240 | 216:284 | 216240 | 216290 | nd | 206226 | nd | nd | nd |
| Hi03b03 | 210:212 | 204:210 | 206210 | 200206 | 200210 | 196208 | 200210 | 212 | 210 |
| Hi03c04 | 169:null | null | 169201 | 201 | 175201 | 257 | 201 | 201 | 169201 |
| Hi03c05 | 205:221 | 205:221 | 179195 | 191193 | 215 | 209 | nd | nd | nd |
| Hi03d06 | 117:117 | 143:169 | 133143 | 143 | 117145 | 115147 | 117 | 117 | nd |
| Hi03e03 | 193:199 | 193:197 | 199 | 189199 | 193199 | 189198 | 187199 | 199 | nd |
| Hi03e04 | 148:160 | 132:148 | 132 | 132150 | 146 | 146160 | 134160 | 150160 | 132 |

Table 4 (continued)

| SSR name | Fiesta | Discovery | Florina | Nova Easaygro | TN10-8 | Durello di Forli | Prima | Mondial Gala | Fuji |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hi03f06 | 153:177 | 153:179 | 153161179 | 153211 | 153217 | 153176205211 | 153211 | 153177212 | nd |
| Hi03g06 | 198:200 | 182:198 | 182196 | 182196 | 182 | nd | 204 | 196204 | 196 |
| Hi04a02 | 82:102 | 90:104 | 102 | 102 | 88 | 100102 | 88102 | 82102 | 102 |
| Hi04a05 | 194:204 | 194:198 | 194204 | 204222 | 198208 | 194202210222 | 194204216 | nd | nd |
| Hi04a08 | 226:246 | 216:250 | 216218 | 212216 | 212216 | 212216 | 212216 | 216226 | nd |
| Hi04b12 | 150:156 | 140:156 | 156 | 142156 | 150156 | 146150 | 156 | 142156 | 142156 |
| Hi04c05 | 181:181 | 183:181 | 181 | 181 | 181 | 181183 | 179183 | 179183 | 181 |
| Hi04c10 | 230:null ${ }^{\text {x }}$ 206:206 ${ }^{\text {y }}$ | 228:238 ${ }^{\text {x }}$ 200:206 ${ }^{\text {y }}$ | 206234 | 206234 | 204234 | 172 | 172202234 | 178202234 | 172202234 |
|  | 176 |  |  |  |  |  |  |  |  |
| Hi04d02 | 208:222 | 238:null 180194 | 176182204 | 182190204 | 190198222 | 186204214218 | 190204214 | 176192222 | 186204218 |
|  |  |  |  |  |  |  |  |  | 222 |
| Hi04d10 | 164:164 | null | 164182 | 180 | 164 | 164 | 182 | nd | 164180 |
| Hi04e04 | 226:246 | 222:228 | 228246 | 216246 | 216222 | 230 | 226246 | 228246 | 216226 |
| Hi04e05 | 144:null | null | null | null | nd | null | null | 144 | null |
| Hi04f08 | 202:218 | 218:226 | 226 | 226 | 220 | 218 | 220226 | 226 | 218 |
| Hi04f09 | 222244254 | $230^{\text {b }} 224244$ | 222224254 | 224226252 | 224256 | 224;254 | 224226244 | 222224244254 | nd |
|  |  |  | 260 | 258 |  |  | 260 |  |  |
| Hi04g05 | null:null 230 | 190:194 230 | 194258 | 232258 | 240254 | 230 | 230 | 230 | 232248 |
| Hi04g11 | null:null 166 | 118:null | 118 | 158 | 158 | 140 | 158 | 164 | 118 |
| Hi05b02 | 126:126 | 120:122 | 126178 | 122178 | 134162 | 120144162178 | 122126 | 126178 | 126134 |
| Hi05b09 | 140:144 | 136:136 | 136 | 136144 | 136140 | 136 | 140144 | nd | nd |
| Hi05c06 | 139:138 | 136:136 | 138141 | 138 | 138141 | 141142 | 138 | 138139 | 136138 |
| Hi05d10 | 212:null | null | 212 | nd | 212 | 212 | nd | 212 | 212 |
| Hi05e07 | 212:214 | 216:230 | 214230 | null | 214234 | null | 214230 | 214 | 228 |
| Hi05f12 | 173: null ${ }^{\text {x }}$ 169:169 ${ }^{\text {y }}$ | 173:177 ${ }^{\text {x }} 157: 169^{\text {y }}$ | 157173 | 157171 | 157173 | 157173 | 157173 | 157173177 | 157173 |
| Hi05g12 | $230^{\text {b }} 208216238$ | 216 | 216238 | 216.192 | 216238 | 222242 | 208216230 | 208216220228 | 216 |
|  |  |  |  |  |  |  | 238 |  |  |
| Hi06b06 | 242:262 | 236:262 | 260 | 260 | 260 | 236260 | 260 | 262 | 242260 |
| Hi06f09 | 280:288 | 272:274 | 280288 | 272288 | 272288 | 274288 | 288 | 272280 | 272288 |
| Hi07b02 | 214:218 | 214:null | 212216 | 214216 | nd | 212 | nd | 216 | 216 |
| Hi07b06 | 210:220 | 220:222 | 226 | 226 | 226 | 226 | 222226 | 222226 | 222226 |
| Hi07d08 | 222:232 | 222:226 | nd | nd | nd | nd | nd | nd | nd |
| Hi07d11 | 214:218 ${ }^{\text {x }} 222: 234^{\text {y }}$ | 216:226 ${ }^{\text {x }}$ 234:null ${ }^{\text {y }}$ | 214220 | 220 | 216232 | 228 | 200218226 | 200220 | 212220 |
|  |  | 200:200 |  |  |  |  |  |  |  |
| Hi07d12 | 246:null ${ }^{\text {x }}$ 184:null ${ }^{\text {y }}$ | 194:218* 246250 | 248 | 248 | 198246 | 198 | 246 | 184206 | 250 |
| Hi07e08 | 208:208 ${ }^{\text {x }}$ 233:235 ${ }^{\text {y }}$ | 208:212 ${ }^{\text {x }} 231: 241^{\text {y }}$ | 208222231 | 208233 | 208233234 | 208233234 | 210222231 | nd | nd |
|  |  |  | 233 |  |  |  |  |  |  |
| Hi07f01 | 204:214 | 204:210 | 206220 | 210 | 210 | 210220 | 210220 | 210 | 206210 |
| Hi07g10 | 126:null | 128.null | nd | nd | nd | nd | nd | nd | nd |

Table 4 (continued)

| SSR name | Fiesta | Discovery | Florina | Nova Easaygro | TN10-8 | Durello di Forlì | Prima | Mondial Gala | Fuji |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hi07h02 | 256:264 | 270:276 | 246256 | 276 | 256268 | 266 | 248274 | 248264 | 246260 |
| Hi08a04 | 246:254 | 246:250 | 246248 | 246 | 248250 | 246254 | 246250 | 246254 | 246254 |
| Hi08c05 | 231:231 | 231:240 | 231240 | 231234240 | 231240 | 231234 | 231234 | 231234240 | 231234240 |
| Hi08d09 | 183 | 183 | 186 | 183 | 183 | 183186 | nd | nd | 183186 |
| Hi08e04 | 207:216 | 201:null | 216 | 207 | 201216 | 216225 | 216 | 207234 | 207 |
| Hi08e06 | 134:138 | 134:null | 134 | 134 | 134 | 134137 | 134 | 134138 | 134 |
| Hi08f05 | 165 | 165 | 165 | nd | 165157 | 165 | nd | nd | nd |
| Hi08f06 | 227:230 | 224:242 | 230 | 230 | 224227 | 230233 | 227230 | 227230 | 230 |
| Hi08f12 | 131:218 | 116:131 | 116220 | 116218 | 206 | 131206 | 158218 | 218220 | 129220 |
| Hi08g03 | 113:116 104 | 116:116 104 | 104113116 | nd | nd | 104113 | 104113 | 104113116 | 104113116 |
| Hi08g06 | 192:198 | 192:192 | 192198 | 192198 | 192 | 192 | 192198 | nd | 192 |
| Hi08g12 | 188:188 | 188:194 | 188194 | 191194 | 188194 | 188197 | 188 | 188191 | 188191 |
| Hi08h03 | 155:155 | 155:158 | 155 | 155 | 155 | nd | nd | 155 | nd |
| Hi08h08 | 236:236 | 236:239 | 236242 | 236 | 236242 | 236239 | 236239 | 236239 | 236 |
| Hi08h12 | 157:203 | 163:169 | 169203 | 151203 | 157163 | 151 | 169203 | 151172 | 157172 |
| Hi09a01 | 183,186 | 186:192 | 183186 | 180,183 | 183,186 | 183,192 | 183,186 | 186 | 183,192 |
| Hi09b04 | 230:227 | 278:278 | 242 | 227269 | 227242 | 227242 | 230242 | 242269 | 230242 |
| Hi09f01 | 257:260 | 257:266 | 260 | 260 | 257260 | 257266 | 257266 | 260 | 260 |
| Hilla01 | 214:217 | 214:217 | 217 | 217 | 217 | 214217 | 214223 | 214223 | 217 |
| Hilla03 | 141:141 | 141:144 | 141 | 141 | 141144 | 141144 | 141 | 141 | 141 |
| Hi12a02 | 249:255 | 255:255 | 249255 | 249255 | 249255 | 249255 | 255 | 255 | 255 |
| Hi12c02 | 190:178 | 169:null | 178 | 169 | 178 | 169178 | 169 | 169 | 190178 |
| Hi12f04 | 184:184 | 184:187 | 187 | 184 | 184 | 184 | 184 | 184187 | 184 |
| Hi15al3 | 220:232 | 220:220 | 232 | 234 | 232 | 220 | 220 | 232 | 232 |
| Hi15b02 | 196:199 | 199:199 | 202 | null | 202 | 202 | 199 | 199 | 202 |
| Hi15c07 | 210:210 | 204:210 | 204210 | 210 | 210 | 204210 | 210 | 204210 | 204210 |
| Hi15e04 | 209:209 | 209:212 | 209 | 209212 | 209 | 209 | 209212 | 209 | 209212 |
| Hi15g11 | 160:163 | 160:160 | 163 | 163 | nd | 160 | 160163 | 160163 | 163 |
| Hi15h12 | 222:222 | 222:225 | 222 | 222 | 222 | 222228 | 222225 | 222225 | 222 |
| Hi16d02 | 144:144 | 144:147 | 144165 | 144 | 144165 | 144177 | 144 | 144 | 144153 |
| Hi20b03 | 220:238 | 220:229 | nd | nd | nd | 220244 | 229 | 220244 | nd |
| Hi21c08 | 230:230 | 230:227 | 230227 | 230227 | 230227 | 230227 | 230 | 230227 | nd |
| Hi21e04 | 134:134 149156 | 134:136 149156 | 134136149 | 136149156 | 134149158161 | 134138148149 | 134136149 | 136138149 | 134138149 |
|  | 158161 | 158161 | 158 | 158161 |  | 158161 | 158161 | 156161 | 156161 |
| Hi21f08 | 242:246 280282 | 242:242 234244 | 242248280 | 242248266 | 242250280 | 242250282 | 250266272 | 244248272 | 244248282 |
|  |  | 250272282 | 282 | 280 |  |  | 280 | 280282 |  |
| Hi21g05 | 158:164 | 155:158:164 | 155158 | 155 | 155158164 | 158 | 155158164 | 155158164 | 155158 |
| Hi22a07 | 192:198 ${ }^{\text {x }} 153: 189^{\mathrm{y}}$ : | 196:196 ${ }^{\text {a }}$ null:null ${ }^{\text {a }}$ | 192202 | 153196202 | 153189202 | 153202 | 189192196 | 189198202 | 189198 |
| Hi22d06 | 129:129 | 126:129 | 132 | 129135 | 129135 | 132 | 126129 | 129135 | 129135 |

Table 4 (continued)

| SSR name | Fiesta | Discovery | Florina | Nova Easaygro | TN10-8 | Durello di Forlì | Prima | Mondial Gala | Fuji |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hi22f04 | 138:147 | 135:138 | 141147 | 141 | 138 | 138 | 138141 | 138141 | 138 |
| Hi22f06 | 240:246 | 243:246 | 240246 | 240246 | 246 | 243 | 246 | 240246 | 240246 |
| Hi22f12 | 211:null | 211:217 | 214217 | 205 | 217 | 211217 | 205 | 211 | 217 |
| Hi22g06 | 240:249 | 240:240 | 240 | 240 | 240 | 240 | 240 | 249 | 240 |
| Hi23b12 | 154:169 | 154:169 | 154169 | 142169 | 142169 | 169 | 142154 | 154169 | 142 |
| Hi23d02 | 160:166 | 160:160 | 157160 | 160 | 157160 | 160 | 160 | 160 | 157160 |
| Hi23d06 | 161:161 | 158:161 170173 | 161170 | 161 | 161 | 161 | 161170 | 161188 | 158188 |
| Hi23d11b | 181:184 177 | 178:184 177 | 177184 | 177184 | 177178184 | 177178181 | 177184 | 177178184 | 177184 |
| Hi23g02 | 248:257 | 254:257 | 239 | 239255 | 239248 | 255 | 239 | 255 | 230248 |
| Hi23g08 | 220:220 | 214:220 211 | 211220 | 220 | 211220 | 211220 | 214220 | 220 | 211220 |
| Hi23g12 | 223:223 241 | 223:226 241 | 226241 | 226235241 | 223226235241 | 223226235241 | $\begin{aligned} & 223226235 \\ & 241 \end{aligned}$ | 223235241 | 223235241 |
| Hi24f04 | 144:150 | 147:153 | 153 | 153 | 150153 | 153 | 150153 | 144150 | 152153 |
| AF057134 ${ }_{\text {SSR }}$ | 210:216 | 208:224 | 208 | 216 | 216 | 214220 | 216 | 208216 | 202216 |
| AF527800 ${ }_{\text {SSR }}$ | 178:null | 168:184 | 168 | 168 | 168 | 168 | 168194 | 168178 | nd |
| AJ000761a SSR | 262:262 | 262:266 | 262264 | 260262 | 262264 | 260266 | 264272 | nd | nd |
| AJ000761b ${ }_{\text {SSR }}$ | 248:248 | 210:248 | 244248 | 210252 | 246248 | 208226 | 210250 | 226244 | 244248 |
| AJ001681 ${ }_{\text {SSR }}$ | 169:191 | 189:191 | 169187 | 169187 | 183 | 189195 | 191 | 169 | 189 |
| AJ251116SSR | 165:167 | 167:167 | 167 | 167 | 167 | 165 nd | nd | 167 | nd |
| AJ320188sSR | 213:null | null:null 199 | 199213 | 203207 | 219 | 191245 | 199 | 199 | 199 |
| AT000174 ${ }_{\text {SSR }}$ | 186:194 | 186:192 | 178186 | 178186 | 186188 | 178 nd | 178200 | 188194 | 178 |
| AT000400 SSR | 198:216 226 | 216:216 | 210224232 | 210224232 | 198210216226 | 216224 | 210216232 | 216224232 | 214224 |
|  |  |  |  |  | 232 |  |  |  |  |
| AT000420 ${ }_{\text {SSR }}$ | 201:201 | 203:205 | 201205 | 201 | 203209 | 201189 | 201205 | 201 | 201205 |
| AU223486ssR | 205:205 | 205:217 | 205208 | 208 | 205217 | 205208 | 205208 | 205208 | 208 |
| AU223548ssR | 270:278 | 270:270 | 262278 | 262270 | 274 | 270278 | 262270 | 262270 | 262278 |
| AU223657 ${ }_{\text {SSR }}$ | 225:225 | 221:225 | 225231 | 219225 | 231 | 233223 | 231 | 231 | nd |
| AU223670 SSR | 194:202 | 194:196 202 | 194196202 | 194196202 | 194202 | 194202 | 194196202 | nd | nd |
| AU301431 ${ }_{\text {SSR }}$ | 213:216 | 216:216 | 213216 | 213216 | 213216 | 216 | 213216 | 216 | 213216 |
| AY187627 ${ }_{\text {SSR }}$ | 300:null | null:null | 300 | 300 | 300 | 300 | null | null | null |
| CN444542 SSR | 136:146 110:110 | $\begin{aligned} & 136: 156 ~ 110: 110 \\ & 128 \end{aligned}$ | 110124136 | 110124136 | 110126146 | 110124140 | 110136 | 110136146 | 110124136 |
| CN444636ssR | 241:241 | 239:243 | nd | nd | 239241 | nd | 239241 | 241 | 239 |
| CN444794 ${ }_{\text {SSR }}$ | 260:306 | 230:298 | 270 | 260 | 230 | 254270 | 256272 | 254270 | 256 |
| CN445290 ${ }_{\text {SSR }}$ | 230:230 | 230:236 | 230 | 230 | 230 | 236 | 230242 | 242 | 236242 |
| CN445599 ${ }_{\text {SSR }}$ | 154:164 131 | 153:176 130132 | 130146176 | 131136 | 131132150154 | 130131152 | 131136154 | $\begin{aligned} & 130132146 \\ & 154 \end{aligned}$ | 130131146 |
| CN491050 ${ }_{\text {SSR }}$ | $>330 \mathrm{c}: \mathrm{c}$ | >330 a:c | nd | nd | >330 c | $>330 \mathrm{a}$ | >330 b | $>330 \mathrm{bc}$ | $>330 \mathrm{bc}$ |

Table 4 (continued)

| SSR name | Fiesta | Discovery | Florina | Nova Easaygro | TN10-8 | Durello di Forlì | Prima | Mondial Gala | Fuji |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| CN493139 ${ }_{\text {SSR }}$ | 144:156 124 | 136:150 124142 | 126140142 | 124126140 | 124142146 | 124142 | 124126142 | 140144162 | 124140156 |
|  |  |  | 162 | 162 |  |  | 144 |  | 162 |
| CN496002 ${ }_{\text {SSR }}$ | 243:257 | 243:259 | 257261 | nd | 243261 | 243257 | 243257 | 243259 | 243257 |
| CN496913 ${ }_{\text {SSR }}$ | 236:236 | 236:242 | 236 | 236 | 236 | 236278 | 236244 | 236252 | 236244 |
| CN581493 ${ }_{\text {SSR }}$ | 194:194 | 184:194 | 228 | 194218 | 192220 | 194228 | 194226 | 194218 | 194218 |
| U78948 ${ }_{\text {SSR }}$ | 188:190 | 180:186 | nd | 178182 | nd | 184188 | 180186 | nd | nd |
| U78949 SSR $^{\text {b }}$ | $215^{\text {x }} 176209225$ | $221^{\mathrm{x}} 174^{\mathrm{y}} 209225$ | 176205219 | 174219 | 176207219 | $172176190211$ | 174215 | 205 | 190205 |
| Z38126 ${ }_{\text {SSR }}$ | 216:216 | 214 | 214216 | 214 | 214 | 216240 | 214 | 214 | 214240 |
| Z71980 ${ }_{\text {SSR }}$ | 170 | 170 | 172 | nd | nd | 172 | nd | 170 | nd |
| Z71981 ${ }_{\text {SSR }}$ | 222:224 | 224:232 | 212224 | 222224 | 224 | 222 | nd | 216222 | 222232 |
| GD147 | 147:151 | 141:153 | 141135 | 155151 | 147 | 141153 | 135153 | 141153 | 141153 |
| HGA8b | 160:164 ${ }^{\text {x }} 135: 151^{y}$ | 156:164 ${ }^{\mathrm{x}} 151: 151^{\mathrm{y}}$ | 156160164 | 133156160 | 151156160164 | 156164 | 135156160 | 151156160 | 151156160 |
|  |  | 135160 |  |  |  |  | 164 | 164 | 164 |
| KA4b | 141:137 | 139:141 | 141 | 137 | 141 | 137141 | 141 | 137 | 137141 |
| NB102a | 181:null | 181:183 | null | 183 | 183 | null | 181 | null | 183 |
| NH009b | 148:162 | 148 | 144148 | 144158 | 138148 | 144152 | 144148158 | 158162 | 148162 |
|  |  |  |  |  |  |  | 162 |  |  |
| NH029a | 91:91 | 91:93 | 91 | 99 | 97 | 95 | 9395 | nd | 9395 |
| NH033b | 189:189 | 175:183 | 163177 | 163177 | 175183 | 179183 | 183189 | 177189 | 177187 |
| MSS6 | 279:279 | 277:273 | 279 | 279 | 273 | 273 | 279 | 279 | 279 |

For the alleles of Fiesta and Discovery, the numbers separated by ":" indicate the estimated size of the alleles of the same locus, while the other numbers indicate the size of undefined loci. For multiloci SSRs, ${ }^{\mathrm{x}}$ and ${ }^{\mathrm{y}}$ indicate the alleles assigned to locus $x$ and $y$, respectively nd allele size not determined
${ }^{\text {a }}$ Amplicons that could be alleles of the locus $x$ as well as of locus $y$. For the other seven cultivars, the size of the amplicons are indicated; single alleles can indicate homozygosity or the
${ }^{\mathrm{b}}$ Due to the complex pattern of alleles amplified of this SSR instead of pair of alleles only an allele per locus was identified and is presented in the table (allele that has been mapped as a dominant marker)

Table 5 Statistics on SSR development

| SSR library/origin of the sequence | Sequenced clones $^{\text {a }}$ | Sequences used for primer design ${ }^{\text {a }}$ | Markers mapped $^{\text {b }}$ | Average no. of alleles per origin of the SSRs | Average no. of repeats per origin of the $\mathrm{SSRs}^{\mathrm{c}}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| GA/GT libraries | 571 | 103 | 65 (63\%) | 6.6 | 28 |
| $\begin{aligned} & \mathrm{AAG}^{\mathrm{d}} / \mathrm{AAC} / \mathrm{ATC}^{\mathrm{d}} \\ & \text { libraries } \end{aligned}$ | 587 | 131 | 52 (40\%) | 4.0 | 9 |
| GenBank 2nt |  | 39 | 24 (61\%) | 5.8 | 14 |
| $\geq 3 \mathrm{nt}$ |  | 21 | 7 (33\%) | 4.3 | 6 |
| Pear and S. torminalis ${ }^{\text {e }}$ |  | 17 | 7 (41\%) | 4.7 | na |
| Apple SSRs from literature ${ }^{\mathrm{f}}$ |  | 8 | 1 (13\%) | $13^{\text {g }}$ | na |
| TOTAL | 1,158 | 319 | 157 (49\%) |  |  |

$n a$ : not analyzable; sequences not available
${ }^{\text {a }}$ The difference between the number of sequenced clones and sequences used for primer design is due to redundant sequences, absence
of a SSR repeat, or a too-short-sequence stretch before or after the SSR repeat for primer design
${ }^{\mathrm{b}}$ In Fiesta $\times$ Discovery, or, alternatively in Durello di Forlì $\times$ Fiesta, Discovery $\times$ TN10-8, or Fuji $\times$ Mondial Gala
${ }^{\mathrm{c}}$ Found in the sequence used for primer designing. Compound SSRs with repeats of different length (e.g., GT-CAA) were not considered. Compound SSR with different repeats but with the same number of nucleotides composing the repeat (e.g., GT-TA) were considered and the length of the two repeats was summed
${ }^{\mathrm{d}}$ Positive clones of these two libraries were screened by PCR for the presence of highly redundant fragments. The number of clones sequenced after this check are reported
${ }^{\mathrm{e}}$ Yamamoto et al. (2002a,b); Oddou-Muratorio et al. (2001)
${ }^{\mathrm{f}}$ Hokanson et al. (1998)
${ }^{\mathrm{g}}$ The number of alleles may be overestimated due to low quality of the amplifications
of these 86 SSRs, 24 (28\%) were developed during this study. Some of the selected markers showed a low level of polymorphism in our set of reference cultivars. They were, nevertheless, included lacking more polymorphic alternatives for these specific genomic regions.

## Discussion

The aim of our research was to obtain a set of highly polymorphic SSR markers that cover the entire apple genome, to enable directed genotyping approaches. Directed genotyping is a target-directed, cost- and timeefficient approach for the genome-wide genotyping of new crosses, cultivars, and breeding lines. By facilitating this method of genotyping, assessments of new molecular marker-trait associations, allele mining and validation of candidate genes will also be enhanced. This paper reports the generation and mapping of a large new set of apple SSR
markers. These SSRs have allowed the enrichment of the reference map of the apple with 168 new loci. This almost doubled the number of mapped SSR markers.

## Efficiency of SSR development

The newly mapped SSRs have been obtained from different sources: genomic libraries, publicly available EST sequences, the literature, and from SSRs developed for other Maloideae species. The efficiency by which each of these sources gave new SSR markers is evaluated in Table 5. Only about $20 \%$ of the library sequences were unique (not redundant), contained a microsatellite repeat, and were suitable for designing compatible primers in the regions flanking the repeat. When exploiting published EST sequences, there is no need to generate enriched libraries, or to sequence, and considerable amounts of money and time can be saved.

Table 6 Frequency distribution (numbers, percentages, and percentage cumulative) of the number of alleles assessed in a set of nine diploid cultivars of single-locus (SL) SSRs divided by the length of the SSR repeat (two nucleotides or more than two nucleotide repeats)

| Absolute no. of SL SSRs |  | No. of alleles |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | Total |
| No. of nt repeats | 2 | 6 | 11 | 8 | 10 | 13 | 6 | 9 | 0 | 2 | 65 |
|  | $\geq 3$ | 13 | 9 | 13 | 7 | 1 | 3 |  |  |  | 46 |
| \% |  |  |  |  |  |  |  |  |  |  |  |
| No. of nt repeats | 2 | 9 | 17 | 12 | 15 | 20 | 9 | 14 | 0 | 3 | 100 |
|  | $\geq 3$ | 28 | 20 | 28 | 15 | 2 | 7 | 0 | 0 | 0 | 100 |
| Cumulative \% |  |  |  |  |  |  |  |  |  |  |  |
| No. of nt repeats | 2 | 9 | 26 | 38 | 54 | 74 | 83 | 96 | 96 | 100 |  |
|  | $\geq 3$ | 28 | 48 | 76 | 91 | 93 | 100 | 100 | 100 | 100 |  |



Liebhard et al. 2002, 2003b. SSRs are indicated in bold. SSRs of the new set have been additionally underlined and their map position is also in bold

Fig. 1 (continued)



Considering the relative high percentage of ESTs containing SSR repeats (about 5\%, data not shown), the efficiency by which SSR containing EST sequences lead to

SSR markers and the continuous increase of publicly available EST sequences (Korban et al. 2005; Crowhurst et al. 2005), it is clear that the next SSR makers should be

4Fig. 2 Set of 102 SSR primer pairs for global coverage of the apple genome. Map positions (in cM ) are aligned to the Discovery maps of Fig. 1. Gray filled bar segments indicate the linkage group segments covered by the Fiesta and Discovery maps of Fig. 1. Open bar segments indicate linkage group segments not covered by the maps of Fig. 1, but which were revealed by other, unpublished linkage maps. CH markers mapped by visual alignment that were initially mapped in other mapping populations are: CH 04 c 06 z [mapped in Prima $\times$ Fiesta and Jonathan $\times$ Prima $(\mathrm{J} \times \mathrm{P})$ ] and CH 02 g 01 (Discovery $\times \mathrm{TN} 10-8$ and $\mathrm{J} \times \mathrm{P}$ ). For 16 loci, indicated with the symbol ?, no primer pairs are publicly available yet. The symbol ?* marks positions of unpublished SSR markers, which are expected to become available in the near future. Underlined SSRs have been developed in the present work
developed from ESTs. Sequences containing long dinucleotide repeats are preferable because of their higher level of polymorphism (Table 5), making them more valuable in directed genotyping approaches. Other new SSRs may be located from the mapping of candidate genes that have SSRs in their non-coding region (e.g., Gao et al. 2005a,b; Costa et al., in preparation), or by the sequencing of previously mapped markers (Gao and Van de Weg 2006).

The approach of transferring SSRs developed in other Maloideae species was efficient (41\%). The high syntheny between apple and pear was used (Yamamoto et al. 2004; Dondini et al. 2004) to select SSRs, which, from the map position in pear, could fill the gaps between SSRs mapped in apple or could enrich regions with few SSRs. This strategy proved to be successful. One of the few new SSR markers mapped on LG1 is from pear (KA4b) and some regions (e.g., LG2, LG9, LG16) of the apple genome, which, at the beginning of the project did not contain a high density of SSRs, were enriched.

The efficiency of mapping previously developed apple GD-SSRs is probably underestimated due to our stringent PCR conditions (see later). Indeed, Hemmat et al. (2003) were able to map all 18 published GD-SSR markers except GD15, which showed very low polymorphism (Hokanson et al. 1998). Ten of these markers could be aligned to our map (data not shown), while seven (those tested in this work) could not be unequivocally aligned due to a low number of markers in common.

## Level of polymorphism

The screening of a newly developed SSR over a restricted number of cultivars (nine in our case) allows an estimation of its true level of polymorphism. Although the number of alleles identified is limited compared to what is present in the apple germplasm, more extensive tests support a rough correlation between the allele numbers identified in a small set of cultivars and the number in germplasm collections (Coart et al. 2003). This information is, thus, useful for a first selection of markers for genotyping projects as well as for genetic diversity studies.

New SSRs designed to meet high-throughput criteria
During the development of new SSRs, variable annealing temperature (Tm) could be used to improve the amplification profile of individual markers. As our aim was to develop SSR markers suitable for high-throughput testing, we, therefore, decided to design all primers with a single $\mathrm{Tm}\left(60^{\circ} \mathrm{C}\right)$, which should enable multiplex PCRs. The relatively high Tm used was set to improve specificity, avoiding amplification of additional, non-SSR 'bands' that could hamper both scoring and multiplexing. On the other hand, some SSR-containing sequences may not have delivered an SSR marker, although primers for lower Tm might have been feasible. Tm other than $60^{\circ} \mathrm{C}$ were accepted only for SSR markers developed and mapped in other Maloideae and that had a good chance of filling gaps in the SSR linkage map.

Differences between our standard PCR conditions and profiles may be the reason why most GD-SSRs could not be mapped by us because of very complex band patterns that rendered impossible the recognition of the SSR alleles, despite giving good results in other studies (Hokanson et al. 1998, 2001; Hemmat et al. 2003; Van de Weg, unpublished).

## Determination of allele sizes

Problems occurring in assessing the size of the amplified amplicons of a cultivar, using ${ }^{33} \mathrm{P}$-labeled primers and polyacrylamide sequencing gel electrophoresis, have been already discussed by Liebhard et al. (2002). The authors stated that 1) the absolute fragment size could be determined only with an accuracy of $\pm 1$ base, 2) differences in size estimation between replications may occur, but relative size differences among amplicons of tested cultivars are constant. These statements were confirmed in this study. Marker assessments with other technology platforms (fluorescently labeled primers, automated sequencers) will also lead to different absolute values, though size differences should remain constant (This et al. 2004). To allow comparisons among studies, we propose to include two or three universal reference cultivars, for which we propose Fiesta, Discovery, and Prima. We find these cultivars suitable because they have been tested with most apple SSR markers and have been involved in many genetic studies, being parental cultivars of various mapping populations in Europe.

## Accuracy of map positions

Although the mapping of the new SSRs is based on the analysis of 44 progeny plants, their map position can be considered to be sufficiently accurate for our purposes. The order of the SSRs was usually identical for both the parental Fiesta and Discovery, thus confirming the validity of their relative position (Fig. 1). In only few cases is the
order of flanking SSRs inverted. In all these cases, the SSRs involved maps very close together. Relative positions of tightly linked markers are usually uncertain due to the effects of missing values and to differences in segregation information among markers (Maliepaard et al. 1997). The data do allow assessment of approximate map positions of the new SSRs, which is sufficient for our current purpose to fill in the gaps in previous maps. In only one case, Z71981 ${ }_{\text {SSR }}$ (LG15), is the approximate map position still to be determined. Some SSRs of the Hi set have already been mapped in other crosses than Fiesta $\times$ Discovery, and their map position relative to other SSRs was confirmed (Patocchi et al. 2005; Gardiner et al. 2006; Erdin et al. 2006; Durel, unpublished).

## Genome covering set of apple SSRs

Currently, around 300 SSRs are mapped on the apple genome, mostly in Fiesta $\times$ Discovery (Fig. 1). Soon there will be sufficient SSR markers to enable an initial genomewide genotyping with a set of 100 SSR markers that have an average inter-marker distance of 15 cM (Fig. 2). Our goal is to develop such a set applying the following criteria: (1) distance between successive markers generally not larger than 20 cM , though occasionally allowed to be up to 25 cM ; (2) most proximal and most distal marker of a linkage group preferably within 10 cM from the linkage group ends; (3) cleanness of the amplifications to allow easy scoring and multiplexing; (4) high level of polymorphism; (5) (un)suitability for multiplexing; (6) range of the allele sizes in view of multiplexing markers from the same LG; (7) preference of CH- over Hi-SSR markers because of the generally wider experience with the former; and (8) preference of single-locus markers over multi-locus markers for ease of data interpretation.

Applying these criteria, we came up with a set of 86 primer pairs, 24 of which ( $28 \%$ ) were developed in the current research. This set still lacks markers for 16 chromosome segments (Fig. 2); however, in four segments, new SSRs have already been identified in other projects (unpublished data). To fill the remaining gaps, RAPD or AFLP markers previously mapped in these regions could be transformed into SCAR markers, which could be used as probes to screen apple BAC libraries. From the positive BACs, sequences containing SSR repeats can be obtained and used for the development of SSR markers. PCR-based methods for the "extraction" of sequences containing SSR repeats from BAC clones are available (Vinatzer et al. 2004).

The current set of 86 SSR markers covers around $85 \%$ of the genome. This set will be used to genotype 350 cultivars and breeding selections as well as 1,400 descendants of 24 crosses within the framework of the HiDRAS project (Gianfranceschi and Soglio 2004). This work will supply additional information on the level of polymorphism of these markers and their compatibility in multiplexed PCRs.

SSR database online
Information on the currently available apple SSR markers is scattered over various publications, and published information remains limited to the initial experiences with these markers. In the course of their use internationally, new information concerning SSR markers will arise with regard to level of polymorphism, range of allele sizes, number of loci, suitability for multiplexing, etc. Worldwide, various groups are also developing new SSR markers from the continuously increasing amount of EST data. For efficient genotyping of cultivars and breeding lines, and for an efficient generation of maps from new crosses, markers should be sorted according to their map position, polymorphism, and/or quality. To facilitate searches for and updates of SSR information, and to concentrate worldwide efforts in the development of new SSR markers, an on-line apple SSR database was constructed (http://www.hidras. unimi.it). This includes information on all SSR markers of the CH (Liebhard et al. 2002), Hi (this paper), and NZ (Guilford et al. 1997; Liebhard et al. 2002) series, and, once available, will include information about the SSRs of the GD series (Hokanson et al. 1998; Hemmat et al. 2003).

This database also allows advanced searches, e.g., a list of SSRs mapped on a specific linkage group, among which SSRs of a certain quality can be selected. For SSR markers with amplification profiles of insufficient quality, new primers can be designed and tested as the sequences of the clones from which the SSR markers were derived can be downloaded for all CH and Hi markers. Updates and comments can be added to each of the SSRs. Researchers are encouraged to share their experiences, especially in the reporting of improvements on problematic markers.

Although more than 300 apple SSRs have been mapped, there are regions of the genome not sufficiently covered with SSRs (Fig. 2). Saturation of these regions is required to allow the construction of maps based solely on SSRs. In addition to the method previously proposed, this could be achieved by the further mapping of EST sequences containing SSR repeats, as hundreds of such sequences are available. Such a work could be performed most efficiently by coordinating action among groups working with apple genetic maps all over the world. The current database is the first step towards the creation of such a worldwide platform. A form is available for announcing that certain EST sequences are under investigation, so that duplication of work could be avoided. Contact addresses are also available so that different research groups can get in touch and, if desired, exchange information.

Acknowledgements The authors thank Davide Gobbin and Vicente Martinez for the technical assistance. This project is carried out with the financial support from the Commission of the European Communities (Contract No. QLK5-CT-2002-01492), DirectorateGeneral Research-Quality of Life and Management of Living Resources Program. This manuscript does not necessarily reflect the Commission's views and in no way anticipates its future policy in this area. Its content is the sole responsibility of the publishers. The Swiss partner has been financed by BBW No. 020053.

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[^0]:    ${ }^{\mathrm{a}}$ Nucleotides added to the primer to create a pigtail (Brownstein et al. 1996) Perfect (Perf), imperfect (Imp), compound (Comp) (Weber 1990)

    Single locus (SL): for each sample, a maximum of only two alleles is amplified and the marker has been mapped to a single locus; multilocus (ML): more than two alleles for each sample is amplified and the alleles amplified with the marker have been mapped to
    ${ }^{\text {d }}$ Clean: no extra bands; complementary band(s) (Compl. b.): the only extra band(s) visible are at a constant distance from the SSR allele; Extra bands: the SSR alleles are clearly visible but other amplicons (few or many) are present; multiplexing may be difficult; Dirty: many non SSR-like amplicons visible, scoring of the SSR difficult; multiplexing not advised ${ }^{\text {e }}$ For this EST, two pairs of primers have been developed, AJ000761a and b

