

A *FRUITFULL*-like gene is associated with genetic variation for fruit flesh firmness in apple (*Malus domestica* Borkh.).

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Abstract

The *FRUITFULL* (*FUL*) and *SHATTERPROOF* (*SHP*) genes are involved in regulating fruit development and dehiscence in Arabidopsis. We tested the hypothesis that this class of genes are also involved in regulating the development of fleshy fruits, by exploring genetic and phenotypic variation within the apple (*Malus domestica*) gene pool. We isolated and characterised the genomic sequences of two candidate orthologous *FUL*-like genes, *MdMADS2.1* and *MdMADS2.2*. These were mapped using the reference population ‘Prima x Fiesta’ to loci on *Malus* linkage groups LG14 and LG06, respectively. An additional MADS-box gene, *MdMADS14* shares high amino acid identity with the Arabidopsis *SHATTERPROOF1/2* genes and was mapped to *Malus* linkage group LG09. Association analysis between quantitative fruit flesh firmness estimates of ‘Prima x Fiesta’ progeny and the *MdMADS2.1*, *MdMADS2.2* and *MdMADS14* loci was carried out using a Mixed Model Analysis of Variance. This revealed a significant association ($p < 0.01$) between *MdMADS2.1* and fruit firmness (FFF-1). Further evidence for the association between *MdMADS2.1* and fruit flesh firmness was obtained using a case-control population-based genetic association approach. For this, a polymorphic repeat, (AT)_n, in the 3’ UTR of *MdMADS2.1* was used as a locus-specific marker to screen 168 apple accessions for which historical assessments of fruit texture attributes were available. This analysis revealed a significant association between the *MdMADS2.1* and fruit flesh firmness at both allelic ($\chi^2=34$, 9 df, $P < 0.001$) and genotypic ($\chi^2=57$, 32 df, $P < 0.01$) levels.

Introduction

The regulatory network controlling dehiscence in the dry fruits of *Arabidopsis* has been the subject of intensive study. Several genes have been identified that are necessary for normal silique development. These include members of the MADS-domain transcription factor family *SHATTERPROOF* (*SHP*) and *FRUITFULL* (*FUL*) (Liljegren et al. 2000; Ferrandiz et al. 2000). The *SHP* genes (*SHP1* and *SHP2*) are required both for differentiation of the separation layer and to promote lignification of the valve margins. When both *SHP* genes are knocked out the silique fails to open and the seeds are trapped inside. *FUL* negatively regulates valve margin development and *ful* mutants show ‘ectopic dehiscence zone formation’ throughout the valve tissue (Ferrandiz 2002; Liljegren et al. 2004). Other dry fruited species also appear to share a similar regulatory mechanism. *FUL* is a member of the *SQUAMOSA* (*SQUA*) family of MADS-box transcription factors and genes with strong homology to *FUL* are present in the ‘core eudicots’ which contain the *Rosaceae* and *Solanaceae* families respectively (Litt and Irish 2003). This is likely to reflect a role for these genes in floral evolution where members of the *SQUA* family are central to floral development. Genes related to *FUL* could also play a role in fruit development throughout at least the core eudicots. A central question is whether this regulatory mechanism is confined to just the dry fruited species such as *Arabidopsis* and *Antirrhinum*. During Angiosperm evolution, fleshy fruits are thought to have arisen from dry forms. Phylogenetic evidence indicates that in the *Solanaceae* fleshy berries are derived from dry types such as capsules (Knapp 2002). We postulate that genes regulating the development and ripening of dry fruits may have been recruited to new, but related functions during the evolution of fleshy fruits (Seymour et al. 2008).

In tomato the gene at the *rin* locus is a MADS-box transcription factor and has been shown to be essential for fruit ripening (Vrebalov et al. 2002). Examination of possible *FUL* orthologues in tomato (*TDR4*, *LeFUL2*) reveals that these genes are highly expressed in carpel tissue and show up-regulation during fruit ripening (Eriksson et al. 2004; also see TIGR Tomato Gene Index, www.tigr.org). In the non-ripening tomato mutants, *nor*, *rin* and *Cnr*, there are low levels of *TDR4* expression (Eriksson et al. 2004) and this again suggests a role for this potential *FUL* orthologue in ripening in a fleshy fruit. In *Malus*, a possible *FUL* orthologue, *MdMADS2*, has already been partially described (Sung et al. 1999), but its role in fruit ripening, if any, is not known. Previously significant QTLs for fruit firmness and flesh texture were identified on apple linkage groups L01, L03, L08, L10, L15 and L16 (King et al. 2000; Maliepaard et al. 2001 and Seymour et al. 2002). Recently, candidate gene approaches placed the *Md-ACO1* on the linkage group L10 within the 5% interval border of an important QTL for fruit firmness (Costa et al. 2005), and showed that allelic variation of an SSR within the *Md-Exp7* gene, which mapped on L01, was associated with fruit softening (Costa et al. 2008).

In this paper we describe the isolation and identification of apple *MdMADS2* / *FUL*-like genomic sequences and demonstrate a genetic linkage with fruit flesh firmness through the use of genetic association studies.

Materials and Methods

Plant material and DNA isolation

DNA samples from the reference population ‘Prima x Fiesta’ (Maliepaard et al. 1998), were obtained from the plantation at Warwick HRI, Wellesbourne, and used for the marker and trait linkage analysis. DNA for the population-based association analysis was obtained for a ‘diversity set’ derived from apple variety specimen trees held in the National Top Fruit Collection at Brogdale, Kent, UK (Supplementary Table 1). For the mapping population, total DNA was isolated from young leaf tissue using a cetyltrimethylammonium bromide (CTAB) miniprep method (Doyle and Doyle 1990). DNA was isolated from the apple diversity set using the Nucleospin multi-96 Plant DNA extraction Kit (Macherey-Nagel, Germany) following manufacturer’s instructions.

Trait assessment

Fruit Flesh firmness (FFF-1) measurements for the reference population ‘Prima x Fiesta’ were described previously (King et al. 2000). The firmness data for the apple varieties were obtained from Smith (1971). Data were scanned by optical character recognition software directly from the entries of the varieties, then hand-edited and curated within a relational database. Smith categorised each variety according to a series of multistate trait characters, with restricted vocabulary. These included the following relating to fruit texture included firmness, coarseness, crispness and hardness (Table S1).

Identification of *FRUITFUL* and *SHATTERPROOF* homologous sequences in *Malus*

Identification and characterisation of the possible *AtFUL/AGL8* orthologue *MdMADS2* (accession no. U78948) in *Malus* has already been described (Sung et al. 1999). The primers 5'-ACTGGAAGCTGGACTCTGGAACATG-3' (forward) and 5'-CTACTGCGTTCTCCTTCTCCTTCAC-3' (reverse) were designed to amplify the 277 bp 3' region of *MdMADS2* (accession no. U78948) and used for PCR amplification on first strand cDNA synthesised from 4 week old, mature green and ripe fruits using Omniscript first-strand cDNA synthesis Kit (Qiagen) following manufacturer's instructions. For PCRs following conditions were used: 50 ng of first strand cDNA was amplified by PCR in a 25 µl final reaction volume containing 100 mM Tris-HCl (pH 8.3), 500 mM KCl, 2.5 mM MgCl₂, 200 µM of each dNTP, 400 nM of each primer and 1.25 U Taq DNA polymerase. The PCR conditions consisted of a preliminary denaturation step of 3 min at 94°C, followed by 30 cycles of 30 s at 94°C, 30 s at 55°C and 1 min at 72°C, with a final extension for 5 min at 72°C. The PCR products were analysed on a 2% agarose gel in 1x TBE (45 mM Tris-base, 45 mM boric acid, 1 mM EDTA, pH 8.0), and bands of the expected sizes were excised and gel-purified with the QIAquick Gel Extraction kit (Qiagen, UK) and the fragments were subjected to sequencing. Sequence analyses revealed that a fragment identical to that of the *MdMADS2* gene had been amplified. The fragment was then radioactively labelled with α-[³²P]-dCTP and used to hybridise the *Malus* BAC library constructed from the apple variety Florina (Vinatzer et al. 1998). Following identification and subsequent confirmation of the positive BAC clones with PCR analysis, the clones were digested with the restriction enzyme *Hind* III and the digestion patterns obtained were used to

assign the BAC clones into contigs. The sequences flanking the *MdMADS2* gene were obtained by the direct BAC sequencing approach. BAC DNA was purified using a Large Construct Purification kit (Qiagen, UK) following manufacturer's instructions. The BAC sequencing reaction consisted of 1 µg of BAC DNA, 4 µl BigDye™ terminator mix (Applied Bioscience), 4 µl 5x buffer (400mM Tris-HCl, pH 9.0, 10mM MgCl₂) and 10µM primer. The cycle conditions consisted of a preliminary denaturation step of 5 min at 95°C, followed by 45-75 cycles of 30 s at 95°C, 10 s at 50°C and 4 min at 60°C. After obtaining 250-400 bp sequence, a new primer was designed and used for subsequent sequencing reactions until approximately 1 kb additional sequence from the initiation and stop codons of the gene were obtained.

Both nucleotide and amino acid sequences of the Arabidopsis *SHATTERPROOF1/2* (Liljegen et al. 2000) were used to identify the homologue *Malus* sequence(s) using the tBlastx algorithm. This analysis identified *MdMADS14* (accession no. AJ251117) with 68% overall (81% positives) amino acid identity as a possible *SHP* orthologue in the *Malus* genome.

Genetic mapping of *MdMADS2.1*, *MdMADS2.2* and *MdMADS14* genes in the *Malus* genome

Following identification of two possible *MdMADS2* paralogues, specific primer pairs were designed for *MdMADS2.1* (5'-GATAGGTATTGTGTTTCCATCCATGC-3' as forward and 5'-TGCTATTTGCAAGTGTTAGTGTGTGG-3' as reverse) and *MdMADS2.2* (5'-AAGAAGGTAAGCCTGAGCTAGAAGG-3' as forward and 5'-CCTCCTCCTTTCAACCATACTTTCC-3' as reverse). Sequence analysis of

MdMADS2.1 amplicons from both parents identified a SNP on one of the alleles present in the female parent Prima. This corresponded to the *Tsp45* I restriction site and enabled the development of a cleaved amplified polymorphic sequence (CAPS) assay marker. A SNP present on one of the Prima alleles was identified following comparison of *MdMADS2.2* sequences of both parents. This corresponded to a restriction site for the enzyme *Ava*II and enabled the development of a CAPS assay.

MdMADS14 amplification was carried out with the primers MADS14B-F (5'-TAACCATTACCCTCGCCAAG-3') and MADS14B-R (5'-GAATTGGGGTCCATCATCAC-3') designed specific to 3' end of the gene using DNA from 'Prima' and 'Fiesta'. The amplified samples were then sequenced. One of the SNPs identified was specific to Fiesta allele and corresponded to the restriction enzyme *HpyCH4III* recognition site. This enabled the development of a CAPS assay.

Genetic linkage analysis was performed using JoinMap V.3.0 (Van Ooijen and Voorrips 2001). Distances (in centi-Morgans) were calculated with the Kosambi (1944) function.

Association analysis using mixed model analysis of variance (ANOVA)

In order to test if the genes *MdMADS2.1*, *MdMADS2.2* and *MdMADS14* were associated with the fruit flesh firmness (FFF-1), previously reported by King et al. 2000; 2001, in the reference population 'Prima x Fiesta', a mixed model analysis of variance (ANOVA) was carried out. For this, *MdMADS2.1*, *MdMADS2.2* and *MdMADS14* were used as fixed effects and the molecular markers corresponding to the previously identified fruit firmness QTL regions (i.e. the markers OPAM-19-2200 and OPC-09-0900 on LG01, OPA092000 on LG03, OPAC110780 on LG08, ACO1 on LG10,

NZ02b1 and LY26a on LG15 (King et al. 2000; 2001)) as random effects using the REML estimates of the reference population 'Prima x Fiesta' fruit flesh firmness data (FFF-1) (King et al. 2000; 2001). Post-hoc comparison between the alleles of the *MdMADS2.1* was carried out using Tukey's test. All statistical tests were carried out using SAS v8.2 (SAS Institute).

Population-based association analysis

A polymorphic repeat (AT)_n in the 3' UTR of the *MdMADS2.1* was identified and used as a marker to screen 192 (96 Firm and 96 Soft) apple accessions obtained from the National Top Fruit Collection at Brogdale, Kent, UK. The PCR amplification was carried out using a fluorescently tagged forward primer 5'-FAM-GATGCTCCGCCACCTTAAT-3' and 5'-AGGGTTTTTCATCATGCACATT-3' as reverse primer. Twelve un-linked SSR markers (Supplementary Table 2) distributed on 12 linkage groups, which do not correspond to previously identified fruit flesh firmness QTL regions (King et al. 2000; 2001), were used for the assessment of possible sub-structure within the apple accessions.

The 10 µl PCR reaction mixture consisted of 20 ng of genomic DNA, 100 mM Tris-HCl (pH 8.3), 500 mM KCl, 2.5 mM MgCl₂, 200 µM of each dNTP, 400 nM of each primer and 1.25 U Taq DNA polymerase. The PCR conditions consisted of a preliminary denaturation step of 3 min at 94°C, followed by 30 cycles of 30 s at 94°C, 30 s at 50°C and 30 s at 72°C, with a final extension for 7 min at 72°C. The amplified fragments were prepared for analysis on an ABI Prism 3100 Genetic Analyser (Applied Biosystem) by mixing 1 µl of a 1:10 dilution of the PCR products with 10 µl of Hi-Di Formamide (Applied Biosystems) and 0.15 µl a ROX GS500 internal standard (Applied

Biosystems). GeneMarker Version 1.1 (SoftGenetics LLC) software was then used to analyze the PCR products.

The genetic structure of the apple accessions was investigated with the Bayesian model-based clustering algorithm implemented in STRUCTURE v. 2.1 (Pritchard et al. 2000; Falush et al. 2003). Each analysis was performed without any prior population assignment with a burn-in of 100,000 and a run length of 1000,000 iterations. We carried out five runs of clusters from K=1 to K=8 allowing for the basic admixture model with uncorrelated allele frequencies. The run showing the highest posterior probability of data was considered for each K value. Basic statistics, including the number of alleles, observed heterozygosity and gene diversity (or expected heterozygosity) for the markers were calculated using PowerMarker V3.23 software (Liu 2002).

Allele and genotype distributions between the soft and firm populations were then compared using the chi-square method calculated by PowerMarker V3.23 software (Liu 2002).

Results

Characterisation of *MdMADS2.1* and *MdMADS2.2* genomic sequences

Hybridisation experiments and subsequent PCR analysis identified 11 candidate BAC clones containing *MdMADS2* sequences. Further fingerprinting of these BAC clones revealed two distinct contigs. This suggested that there were two possible paralogous sequences and loci within the *Malus* genome. We named these sequences *MdMADS2.1* (accession no.DQ205652) and *MdMADS2.2* (accession no.DQ205651). Using the direct BAC sequencing and primer walking approaches, we obtained 8,250 bp

and 6,167 bp of sequence for *MdMADS2.1* and *MdMADS2.2*, respectively (Fig. 1b). Alignment of the predicted amino acid sequence of the *MdMADS2.1* differs from that of the previously identified *MdMADS2* (U78948) cDNA sequence by only two amino acids (99% identity). These are therefore likely to be two different allelic forms of the same gene. However, the predicted amino acid sequence of *MdMADS2.2* showed only 84% identity with *MdMADS2* suggesting that *MdMADS2.2* is a possible paralogue of the *MdMADS2.1* (Fig. 1a). A schematic diagram of the predicted introns and exons of the *MdMADS2.1* and *MdMADS2.2*, as well as examples of the nucleotide differences between these two genes are given in Fig. 1b.

Linkage mapping of the *MdMADS2.1*, *MdMADS2.2* and *MdMADS14*

The *MdMADS2.1*, *MdMADS2.2* and *MdMADS14* specific primers were used to amplify sequences from Prima and Fiesta genomic DNA. For each of these genes only a single product was amplified from genomic DNA of both parents. Sequence analysis of *MdMADS2.1* amplicons from both parents identified a SNP in one of the alleles present in the female parent Prima. This enabled the development of a CAPS assay to map *MdMADS2.1* to 41cM on linkage group LG14 (LOD>10), located between the markers 7B9a and E35/M47/P4. (Fig. 2)

A SNP for the sequenced region of the *MdMads2.2* gene was identified. The SNP was used to design a CAPS assay, which allowed mapping of the *MdMADS2.2* gene to 20 cM on linkage group LG06 (LOD>10) between the markers 7B9b and UBC219-1000 (Fig. 2).

A SNP specific to one of the Fiesta alleles of *MdMADS14* was used to develop a CAPS assay which allowed the *MdMADS14* gene to be mapped to 14 cM on linkage

group LG09 (LOD>10) (Fig. 2) between the markers OPAB-14-0800 and OPAE-01-1190.

Association analysis using mixed model ANOVA

The three genes were located on linkage groups, which had not been associated with significant QTL accounting for variation in fruit flesh firmness (FFF-1) previously identified in the Prima x Fiesta population (King et al, 2000; 2001). In order to determine whether weak effects existed between the gene loci *MdMADS2.1*, *MdMADS2.2* and *MdMADS14* and fruit flesh firmness, we reanalysed the segregating trait data using a mixed model ANOVA. In order to reduce the number of parameters, first all the genes and the markers were included in the test, and then the genes and markers with a *P* value of higher than 0.20 were removed in subsequent analyses in order to simplify the model. At this stage *MdMADS2.2* was also excluded as the probability value was higher than the threshold value of 0.20 (*P*= 0.60). *MdMADS2.1* and *MdMADS14* were then used as fixed effects with the molecular markers corresponding to the previously identified fruit firmness QTL regions (i.e. markers on the linkage groups LG01, LG08, LG10, LG15, LG03) as random effects. In this analysis, *ACO1* (LG10), *MdMADS2.1* (LG14), OPC-09-0900 (LG01) and LY26a (LG15) all had highly significant effects on fruit flesh firmness (*P*<0.01) (Table 1). The model used explained 50 % of the variation in fruit flesh firmness in the reference population 'Prima x Fiesta (Table 1). Prima was shown to produce fruit with softer texture than Fiesta in a previous study (King et al. 2001). The progeny with the Prima allele of *MdMADS2.1* had significantly lower fruit flesh firmness.

Population-based association analysis

In order to further test the contribution of the *MdMADS2.1* gene to apple fruit flesh firmness, a population-based association analysis was carried out (Mackay and Powell, 2007). 192 apple accessions (96 Soft and 96 Firm) with fruit firmness data were selected from the National Top Fruit Collection at Brogdale, Kent, UK. In order to avoid spurious association as a result of population sub-structure, twelve unlinked SSR markers (Supplementary Table 2) were used to investigate population structure in the apple diversity set using the software STRUCTURE v. 2.1 (Pritchard et al. 2000) implementing a Bayesian clustering approach. This analysis revealed that there was no significant population structure in the overall sample. Under the model tested, the smallest probability is associated with a prior K of 1 (Posterior probability of 1). The proportion of individuals assigned to each cluster is approximately the same (Supplementary Fig. 1). Therefore, a possible effect of population structure on the subsequent association analysis was found to be unlikely.

The microsatellite sequence (TA)_n located within the 3' UTR region of the *MdMADS2.1* gene was used as a molecular marker in a fluorescence-based genotyping of the apple accessions. Good quality data were obtained from 168 apple accessions. A total of 10 alleles and 33 different genotypes were observed, with the allele sizes ranging from 174 to 196 bp. The distributions and frequency of the alleles obtained within firm and soft apple accessions are given in Fig. 3. Using this data set, a case-control population-based association analysis was carried out. This analysis identified a significant association between the sensory firmness data and *MdMADS2.1* 3'UTR SSR marker both at allelic ($\chi^2=34$, 9 df, $P<0.001$) and genotypic level ($\chi^2=57$, 32 df,

$P < 0.01$) (Table 2). We conclude from this that there is a significant association between the fruit flesh firmness and the *MdMADS2.1* gene.

Discussion

The *FUL* and *SHP* genes have been shown to be involved in development and ‘ripening’ in the dry fruits of *Arabidopsis* (Dinneny and Yanofsky 2005). Related sequences are also found to be expressed in fleshy fruits including tomato and peach (Seymour et al., 2008). In this study we identified genomic sequences of two *FUL*-like genes, *MdMADS2.1* and *MdMADS2.2*, present in the *Malus* genome. These two paralogous genes are highly similar in their predicted coding sequences (Fig. 1a) but show divergence in their non-coding regions (Fig. 1b). In a previous study (Sung et al. 1999) expression analysis revealed that *MdMADS2* (corresponding to *MdMADS2.1*) was transcribed through all stages of flower development and, its transcription could be detected in the four floral organs. In our study, gene specific primers were used to identify the expression patterns of *MdMADS2.1* and *MdMADS2.2* in developing apple fruit cortex. Both *MdMADS2.1* and *MdMADS2.2* transcripts could be detected in 4 week-old, mature green and ripe apple fruits, with *MdMADS2.1* being more highly expressed than *MdMADS2.2* (data not shown). We also identified *MdMADS14* as a likely apple *SHP* orthologue based on sequence homology.

To test whether these genes were associated with fruit texture we adopted two strategies. The first involved evaluating the association between cortex texture and *MdMADS2.1*, *MdMADS2.2* and *MdMADS14* in the progeny obtained from the cross of ‘Prima’ and ‘Fiesta’ (Malieepard et al. 1998). This analysis located the *MdMADS2.1*

gene on LG14 and the *MdMADS2.2* and *MdMADS14* on, LG06 and LG09, respectively. However, none of these linkage groups coincided with the previously identified fruit firmness QTL regions (King et al. 2000; 2001). In a recent study, Liebhard et al. (2003) carried out QTL analyses for a number of physiological traits in apple using a progeny derived from ‘Fiesta’ and ‘Discovery’. In this study, fruit flesh firmness QTLs were also identified on the LG06 and LG14. Recently, the SSR corresponding to *MdMads2* (U78948) was also mapped using the Fiesta x Discovery cross (Silfverberg-Dilworth et al. 2006). This marker was placed on LG14 at 45.6 cM, which shows close proximity to the QTL region (at 47 cM) identified by Liebhard et al. (2003). The QTL located on LG06 however, does not coincide with the location of the *MdMADS2.2*. Many fruit quality traits are affected by the inheritance of alleles at multiple loci and their possible interactions. The identification of these loci is still a key challenge of modern fruit genetic research. Conventional QTL analysis could not detect any association between the fruit flesh firmness and the *MdMADS2.1*, *MdMADS2.2* and *MdMADS14* genes in the ‘Prima x Fiesta’ population. This might be because the sizes of individual locus effects are modest, owing to genetic interactions between loci as suggested by Carlborg and Haley (2004) or the different alleles inherited from the two parents do not show significant functional difference. A range of approaches can be used to explore genetic networks for various QTLs (Kulwal et al. 2005; Li et al. 1997; Mei et al. 2003; Xing et al. 2002; Yu et al. 1997; Zhuang et al. 2002; Paran and Zamir 2003; Schadt et al. 2005; Brem et al. 2005). In this study we used mixed model ANOVA to test the association between the fruit flesh firmness and the genes of interest. This was carried out by controlling for the effect of previously identified QTLs (King et al. 2000; 2001). We obtained a significant association between the *MdMADS2.1* and the fruit flesh firmness

trait using mixed model ANOVA approach. Although there seems to be some correspondence in map location between the association and QTL studies, care has to be taken over the levels of significance associated with the latter, especially where QTL have been identified as putative. The association we obtained between the *MdMADS2.1* and the fruit flesh firmness should therefore be further tested by additional genetic as well as functional studies. The effect of the *MdMADS14*, a possible *SHP* orthologue, was close to significance. There are likely to be paralogues of *MdMADS14* in the *Malus* genome, some of which could be functionally redundant, as is the case for the *SHP* genes in *Arabidopsis* (Pinyopich et al. 2003).

We obtained further evidence for the association between *MdMADS2.1* and fruit flesh firmness in apple using a population-based association approach. Population-based genetic association studies have been extensively used to identify genes or genetic networks underlying complex phenotypes in medical genetics (Risch 2000; Hirschhorn and Daly 2005; Cordell and Clayton 2005). Such approaches have also been successful for the identification of genes/alleles affecting various plant traits (Neale and Savolainen 2004; Krutovsky and Neale 2005; Breseghello and Sorrells 2006; Andersen et al. 2005; Rafalski 2002; Gupta et al. 2005). In population-based association studies, spurious association could arise from subdivisions in the sample, referred to as population structure (Pritchard et al. 2000). Therefore, we investigated the genetic structure of the apple accessions used in this study with the Bayesian model-based clustering algorithm implemented in STRUCTURE (Pritchard et al. 2000). This analysis showed no evidence for significant population substructure that may affect our association analysis.

The data showed that specific alleles of *MdMADS2.1* were significantly associated with qualitative assessments of fruit texture. Although our studies link the *MdMADS2.1* locus with aspects of texture characteristics in apple, the biological basis of these effects is still to be resolved. Functional and genetic analysis in other species may, however, provide some important clues.

In tomato the *TDR4* gene shares significant homology with *MdMADS2* and *FUL*, it is up-regulated in wild type fruits and its expression is restricted in non-ripening mutants. This supports a role for these types of genes in the ripening of fleshy fruits. In *Arabidopsis*, the documented mode of action of *FUL* involves the repression or negative regulation of the genes controlling valve-margin development (Ferrandiz et al. 2000; Liljegren et al. 2004). Therefore, the *FUL* orthologues in other plant species with fleshy fruits could possibly play an analogous role to *FUL* in *Arabidopsis* and may repress cell separation and prevent a total loss of cell-to-cell adhesion, which could in turn lead to firm/crisp fruit phenotypes. Recently, Tani et al. 2007 compared the differential expression of peach *FUL*-like gene (*PpMADS6*), which is 83 % identical to *MdMads2.1*, between a split-pit sensitive and a split-pit resistant peach, and established an association between the expression level and the split-pit phenotype. In addition, Xu et al. 2008 found that constitutive expression of *PpMADS6* in *Arabidopsis* could prevent the pod shattering.

Previously, it has been shown that there is likely to be a high level of synteny between the *Prunus* genome and the two component genomes of apple (Dirlewanger et al. 2004). A likely orthologue of the *Malus MdMADS2.1* (also known as *MdMADS2*) gene in *Prunus* is EST *PrpAPI* (BU039475) (Silva et al. 2005). This gene maps to linkage group 'G05' of the 'Texas'x'Earlygold' *Prunus* reference map, only 1 cM away

from the molecular marker AG108A. Interestingly, in another study AG108A locus was found to harbour QTL acting on fruit size (Quilot et al. 2004), but no measurements of fruit firmness were reported. Fruit size could have an effect on texture characteristics depending on density of cells and cell wall material within the tissue.

Fruit ripening is the process of biochemical and physiological changes involving complex gene interactions. Functional analyses of genes in woody perennials such as apple are, however, very hard to pursue because of long generation times and space requirements for experimental material. Therefore, genetic studies are still the choice for the identification of genetic components underlying agronomic traits. The work described here demonstrates that information on the regulatory network controlling fruit development in the siliques of the model plant *Arabidopsis* can be used to nominate gene candidates involved in the texture of apple fruits.

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Table 1 Association analysis of *MdMADS2.1* and *MdMADS14* genes with the apple fruit flesh firmness in the reference population ‘Prima x Fiesta’ using mixed model analysis of variance

Factors	Effect	Degree of freedom	SS	MS	<i>F</i>	<i>P</i> -value
<i>MdMADS14</i>	Fixed	1	0.96	0.96	3.33	0.0716
<i>MdMADS2.1</i>	Fixed	1	2.80	2.80	9.68	0.0026
<i>ACO1</i>	Random	1	5.060	5.06	17.50	<0.0001
OPAC-110-780	Random	1	1.14	1.14	3.94	0.0504
OPC-09-0900	Random	1	4.814	4.81	16.65	0.0001
LY26a	Random	2	5.46	2.73	9.45	0.0002
Error		82	23.71	0.29		
R^2	0.50					

Table 2 Comparison of allele and genotype distributions between the soft and firm apple accessions using the chi-square method for the determination of population-based association between *MdMADS2.1* gene and sensory fruit firmness

Marker	Allele ChiSquare	Genotype ChiSquare	Allele df	Genotype df	Allele <i>P</i> -Value	Genotype <i>P</i> -Value
<i>MdMADS2.1</i> SSR 3'UTR	34	57	9	32	<0.001	<0.01

Legend to Figures

Fig. 1 (a) Alignment of the predicted amino acid sequences of MdMADS2 (accession no U78948), MdMADS2.1 (accession no DQ205652) and MdMADS2.2 (accession no DQ205651). The asterisks represent amino acid residues identical to the corresponding residues in MdMADS2 **(b)** Schematic diagram of the predicted exons and introns of the *MdMADS2.1* and *MdMADS2.2* genes. Solid boxes indicate the coding regions. The lines connecting the solid boxes represent the seven introns and the 5'- and 3'-flanking regions. Nucleotide differences between *MdMADS2.1* and *MdMADS2.2* genes are shown in panels in the lower part of the figure and the different nucleotide sequences were highlighted. Positions of nucleotide deletions are shown as (a) totalling over 1 kb, (b) 1 bp, (c) 7 bp, (d) 12 bp, (e) 9 bp, (f) 22 bp and (g) 190 bp

(a)

```

MdMADS2      MGRGRVQLKRIENKINRQVTF SKRRSGLMKAHEISVLCDAEVALIIFST
MdMADS2.1    *****
MdMADS2.2    *****T***L*****

MdMADS2      KGKLF EYSNDSCMERILERYERYSYTERQL LANDNESTGSWTLEHAKLKA
MdMADS2.1    *****
MdMADS2.2    *****HA*****N*****

MdMADS2      RVEVLQRNQRHYMGEDLQSLSLKELQNL EQQLDSALKHIRSRKNQVMYES
MdMADS2.1    *****
MdMADS2.2    *****I**F*****RR*****

MdMADS2      ISELQKKDKALQEQNLLAKKVKEKENAVAQQAQLEHVQEQLN SSSSLL
MdMADS2.1    *****
MdMADS2.2    *****N*****K**TS****D*A*K*S*D**T**

MdMADS2      PRALQSLNFGSGSNY-QAIRSSEGI PGDNQQYGDETPTPHRPNMLLPAWI
MdMADS2.1    *****M
MdMADS2.2    *QE**Y***-*R***H*****N**S*****D****A*****T***P*M

MdMADS2      VRHLNE
MdMADS2.1    L*****
MdMADS2.2    L*****

```

(b)

MdMADS2.1

MdMADS2.2

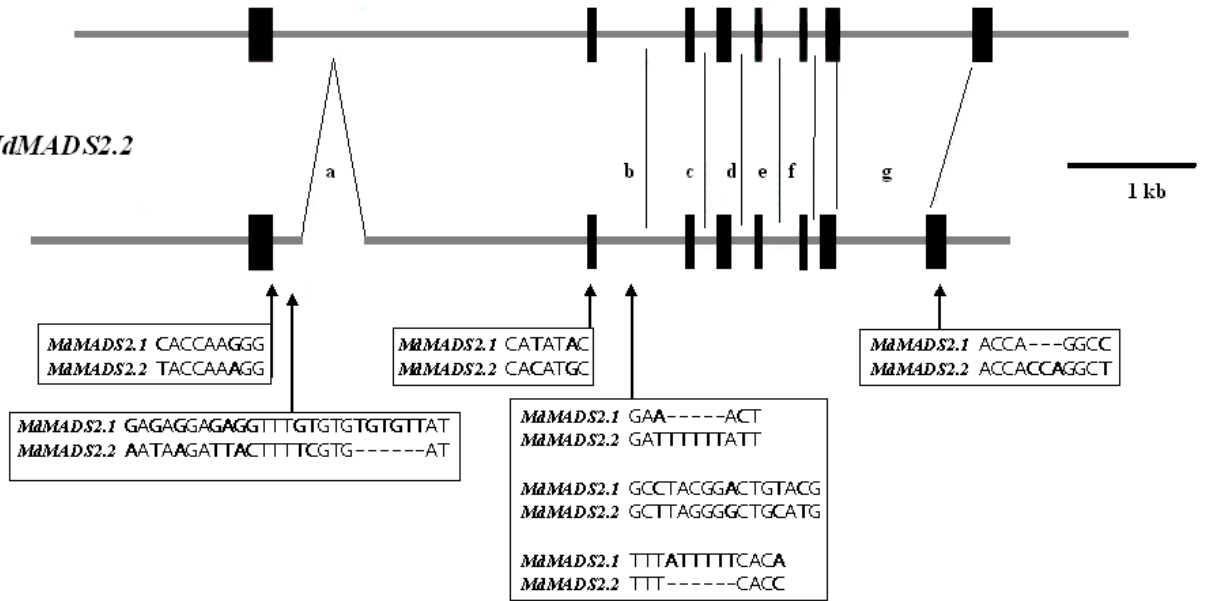


Fig. 2 Genetic map positions of *MdMADS2.1*, *MdMADS2.2* and *MdMADS14*

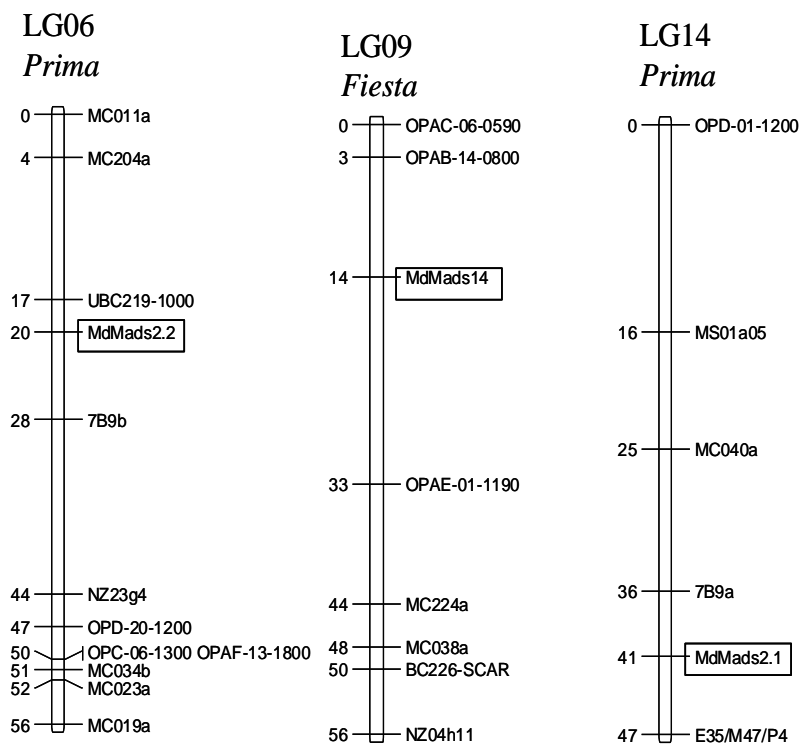


Fig. 3 Allele frequency distributions for (TA)_n repeat polymorphism in the 3'UTR of the *MdMADS2.1* gene in 88 Soft and 80 Firm apple accessions

