

Structural genomics and transcriptional characterization of the Dormancy-Associated MADS-box genes during bud dormancy progression in apple

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Abstract The molecular control of bud dormancy establishment and release is still not well understood, although some genes have already been demonstrated to play important roles in this process. The dormancy-associated MADS-box (*DAM*) genes were first identified in the peach *EVERGROWING* locus and are considered the main regulators of bud dormancy control. In this work, the apple (*Malus × domestica* Borkh.), a perennial plant adapted to temperate climates that displays cycles of growth and bud dormancy, was screened for the presence of *DAM* genes. The candidate genes retrieved were characterized in comparison to *DAM* genes from other species. Four of them (*MdDAMI-4*) are structurally very similar to the reported

DAM genes. When apple genomic segments containing these candidates were compared to the peach *EVERGROWING* locus, a highly conserved noncoding region was detected inside their largest intron. Similar sequences were also identified inside introns of apricot and pear *DAM* genes. Organ expression patterns revealed that *MdDAMI-4* are mainly expressed in dormant buds and seeds, with low transcript accumulation in vegetative structures. In addition, the *MdDAM* genes showed seasonally oscillating patterns of steady-state messenger RNA (mRNA) levels and were down-regulated by artificial chilling. Motif analyses in the promoter and in the intronic conserved region of the *MdDAM* genes disclosed some clues to the regulation of the expression patterns observed. Possible roles for the conserved intronic sequence in dormancy regulation are discussed.

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Introduction

Temperate fruit trees developed a cycle of alternating growth and arrest periods as an adaptation to survive winter conditions. Growth cessation usually involves the formation of winter-hard, cold-tolerant specialized buds and the establishment of dormancy, a developmental program defined by the failure to resume growth from meristems under favorable conditions (Rohde and Bhalerao 2007). The timing of the onset and the length of the dormancy period are modulated by both heritable determinants and environmental cues. Dormancy entrance may be elicited by short photoperiods and/or low temperatures, and dormancy release requires a genetically controlled period of exposure to cold

temperatures, which is called chilling requirement (CR; Falavigna et al. 2015).

The molecular control of development in plants and most other eukaryotes is largely attributed to the MADS-box family of transcription factors (Kaufmann et al. 2005; Dornelas et al. 2011). The MIK^C-type proteins belong to a plant-specific MADS-box subfamily and play several roles in the formation of reproductive organs specific to seed plants (Kaufmann et al. 2005). Genetic mapping of the *EVERGROWING* (*EVG*) locus in peach, whose deletion renders the plant incapable of entering dormancy, revealed six MIK^C-type MADS-box genes (Bielenberg et al. 2008). These genes were named dormancy-associated MADS-box (*DAM*) genes, and two of them (*PpDAM5* and *PpDAM6*) are progressively more expressed in dormant buds during winter and peak at the winter solstice, at which point their transcript levels start to decrease until budbreak (Li et al. 2009). *DAM* genes belong to the StMADS11/SVP-clade, represented in *Arabidopsis thaliana* by *SHORT VEGETATIVE PHASE* (*SVP*) and *AGAMOUS-LIKE 24* (*AGL24*). Both genes act antagonistically on the regulation of the flowering time, with *SVP* expression being modulated by cold exposure (Lee et al. 2007). Homologs of the *DAM* genes are also found in the rosaceous fruit crops pear (Ubi et al. 2010) and apricot (Sasaki et al. 2011) and also in the less-related weed leafy spurge (Horvath et al. 2010). In the kiwifruit, four *SVP*-like genes were characterized, and their possible roles in bud dormancy were described (Wu et al. 2012), although the ectopic expression of one of these members (*SVP3*) in kiwifruit and tobacco did not affect growth and dormancy (Wu et al. 2014). The expression of pear and apricot *DAM* genes is associated with dormancy progression in a similar way to *PpDAM5* and *PpDAM6* (Ubi et al. 2010; Sasaki et al. 2011). The overexpression of the apricot *PmDAM6* in poplar caused growth retardation and precocious bud set, providing further evidence for the role of *DAM* genes in bud development and dormancy (Sasaki et al. 2011).

The apple (*Malus × domestica* Borkh.) is a deciduous tree with well-defined growth and dormancy cycles and with variable CRs across cultivars (Labuschagné et al. 2002). Farmers from mild winter climates must resort to cultivars with a low CR and/or to dormancy-breaking agents to compensate for the insufficient chilling exposure. Despite the importance of dormancy traits for apple production and the availability of many genetic and genomic resources for apple research, no functional genomics approach was reported specifically for the identification of candidate genes for apple dormancy control until very recently (Falavigna et al. 2014; Porto et al. 2015; Wisniewski et al. 2015; Mimida et al. 2015).

With the aim of identifying apple *DAM* genes, the apple genome was searched for putative *PpDAM* orthologs. The sequences of six candidate apple *DAM* genes were thoroughly characterized in silico by comparison to the reported rosaceous *DAM* genes, and their transcription profiles were

determined in several organs and in buds during various dormancy stages and under artificial chilling. In addition, we monitored the transcript accumulation of the candidate apple *DAM* genes in three genetic backgrounds under field conditions. The results show a strongly conserved dormancy-related expression among the different genotypes for four out of the six candidates. Strikingly, these four genes, as well as all the rosaceous *DAM* genes analyzed, showed a highly conserved noncoding sequence inside their largest intron containing potential regulatory elements.

Results

Genomic structure and phylogenetic analysis of *DAM* genes in apple

To find and investigate the *DAM* candidate genes in apple, the deduced amino acid sequences of all six peach *DAM* genes (Bielenberg et al. 2008) were used as queries for BlastP analyses in the apple genome (<http://www.rosaceae.org>, version v1.0), and six predicted proteins repeatedly appeared (E value $<10^{-40}$). Comparisons with the recent literature yielded matches for five out of these six predicted gene models identified (Table 1, see the “Discussion” section for the proposed nomenclature for the apple *DAM* genes). One gene, *MdDAM1*, had been already identified by our group as dormancy-related using suppressive subtraction hybridization (Falavigna et al. 2014). Surprisingly, recent literature revealed that two of the six predicted proteins with high amino acid sequence similarity to peach *DAM* genes are in fact apple homologs of *JOINTLESS*, a transcription factor that plays roles in the abscission zone development (Table 1, Nakano et al. 2015).

Because *MdDAM2–4* are located in a narrow region in apple chromosome 8 (Table 1), in a similar trend to the localization of *DAM* genes in peach chromosome 1 (the *EVG* locus, Bielenberg et al. 2008), we evaluated if the apple genome presents a counterpart to the *EVG* locus. Syntenic relationships were inferred between peach chromosome 1 and all 17 apple chromosomes by employing a collinearity approach (Wang et al. 2012). The overall collinearity result shows a strong syntenic relation between peach chromosome 1 and apple chromosomes 8, 13, 15, and 16 (Fig. 1, gray lines). Additionally, a pair of collinear blocks containing *DAM* genes was identified between the apple chromosome 8 and the peach *EVG* region (Fig. 1, black line). In addition, *MdDAM1*, located in chromosome 16, is embedded in a small block presenting six genes collinear to the *EVG* locus. However, the collinearity algorithm only considers a gene as part of a collinear block if it participates in the sequence of pairs of reciprocally similar genes. *MdDAM1* was not included as one of these genes, but another *DAM*-like gene model in the same block, MDP0000255146 (Table 2), paired with

Table 1 Identification of *DAM* genes in apple

Genome accession	Chromosomal location	GenBank accession	Wisniewski et al. (2015)	Mimida et al. (2015)	Nakano et al. (2015)	Proposed nomenclature
MDP0000322567	chr16:17757588–17766911	KT582786	<i>MdDAM1</i>	<i>MdDAMa</i>		<i>MdDAM1</i>
MDP0000259294	chr8:23859318–23868749	KT582787	<i>MdDAM2</i>	<i>MdDAMc</i>		<i>MdDAM2</i>
MDP0000527190	chr8:23937057–23948304		pseudogene	<i>MdDAMd</i>		<i>MdDAM3</i>
MDP0000232313	chr8:24062517–24065077	KT582789				<i>MdDAM4</i>
MDP0000233948	chr11:18052613–18056301	KT582788		<i>MdSVPa</i>	<i>MdJa</i>	<i>MdJa</i>
MDP0000209705	chr4:414903–418577	LC004730	<i>MdDAM3</i>	<i>MdSVPb</i>	<i>MdJb</i>	<i>MdJb</i>
MDP0000259296	chr8:23891996–23894262					<i>SVP-like/DAM-like</i>
MDP0000285270	chr8:24040116–24041264					<i>SVP-like/DAM-like</i>
MDP0000255146	chr16:17722192–17733590					<i>SVP-like/DAM-like</i>

Names were attributed following a literature review. Genome and GenBank accession codes are provided by the “*Malus × domestica* Genome” (<http://rosaceae.org/>) and NCBI (<http://www.ncbi.nlm.nih.gov/>) databases, respectively. GenBank accession codes for *MdJb* were provided by Nakano et al. (2015). Chromosomal localization is shown

PpDAM1. The details about the collinear blocks including *DAM* genes are shown in the Online Resource 1.

To confirm and check the apple genome annotation, specific primers were designed for the amplification and sequencing of the entire coding regions for all apple *DAM* candidates, using 5' and 3' rapid amplification of complementary DNA (cDNA) ends (RACE; Online Resource 2). We were able to sequence the full coding region of *MdDAM1*, *MdDAM2*, *MdDAM4*, and *MdJa* (GenBank KT582786 to KT582789, Table 1). Alignments revealed some mispredictions in these four gene models disclosed in the

apple genome (Online Resource 3). For *MdJb*, we were able to obtain ~70 % of the coding sequence (data not shown), while only a small fragment of *MdDAM3* could be amplified and sequenced (Online Resource 4).

Phylogenetic analysis with DAM- and SVP-like proteins from seven different species (Online Resource 5) indicated that the majority of the proteins are grouped into three major clusters of orthologs and paralogs, with the apple members distributed into two of them (Fig. 2a). All DAM proteins from peach and Japanese apricot grouped in one distinct clade as orthologs. Following the same pattern, *MdDAM1–4* formed orthologous pairs with pear DAM proteins in a second clade. Finally, the last clade was composed of *MdJa*, *MdJb*, leafy spurge DAM proteins, *Arabidopsis* SVP, and kiwifruit SVP-like proteins *AdeSVP1* and *AdeSVP4* (Fig. 2a). Putatively conserved protein domains for all candidate *MdDAM* genes as well as DAM protein sequences from other species were identified using the software MEME (Bailey et al. 2009). The three characteristic domains of the MIKC^C subgroup of MADS box, the MADS (red and purple), I (orange) and K (blue) domains, were found (Fig. 2b). All sequences presented these domains, except *EeDAM1* and the out-group *Arabidopsis* FLOWERING LOCUS C (FLC). Additionally, both sequences presented a short MADS-box domain in relation to the other sequences analyzed (absence of purple box in Fig. 2b). Finally, only the sequences presented in the clade composed of *MdJa*, *MdJb*, *EeDAM1*, *EeDAM2*, *AtSVP*, *AdeSVP1*, and *AdeSVP4* do not show a conserved C-terminal domain (green boxes in Fig. 2b).

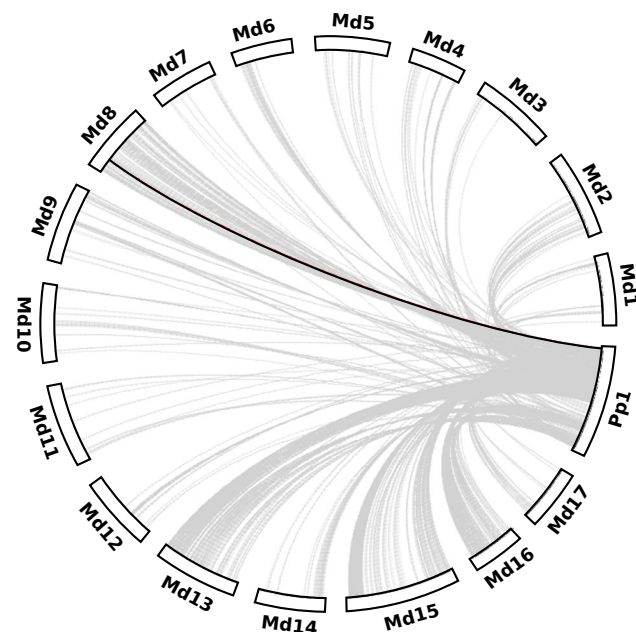


Fig. 1 Collinearity analysis between peach chromosome 1 (*Pp1*) and the apple chromosomes (*Md1* to *Md17*). Collinear blocks containing *DAM* genes are linked by a black line, and the remaining collinear blocks are connected by gray lines

Characterization of the conserved intronic region of *DAM* genes

To further investigate the sequence and structural similarities of the *DAM* candidates, we compared apple genome contigs

Table 2 Results of BlastN searches using the conserved intronic sequence as a query in rosaceous genomes

Species	<i>E</i> value	Predicted gene or genome scaffold	Feature location	Annotation
Apple	1e-39	MDP0000322567	chr16:17757588-17766911	MdDAM1
Apple	4e-21	MDP0000259294	chr8:23859318-23868749	MdDAM2
Apple	4e-21	MDP0000259296	chr8:23891996-23894262	SVP-like/DAM ^a
Apple	6e-20	MDC027822.23	chr8:24030046-24032204	–
Apple	6e-20	MDP0000285270 ^b	chr8:24040116-24041264	SVP-like/DAM ^a
Apple	6e-20	MDP0000255146	chr16:17722192-17733590	SVP-like/DAM ^a
Apple	4e-15	MDC009761.283	chr8:24030669-24045548	MdDAM4 ^c
Apple	5e-14	MDP0000527190	chr8:23937057-23948304	MdDAM3
Apple	5e-14	MDC009761.249	chr8:23901009-23910020	–
Apricot	5e-23	Pm004418	chr2:4838595-4847509	PmDAM3
Apricot	5e-23	Pm004417	chr2:4829851-4837429	PmDAM4
Apricot	2e-21	Pm004419	chr2:4853803-4865323	PmDAM2
Apricot	3e-20	Pm004416	chr2:4820532-4828577	PmDAM5
Apricot	1e-11	Pm004415	chr2:4810970-4819434	PmDAM6
Apricot	1e-06	Pm004420	chr2:4867318-4875401	PmDAM1
Pear	4e-35	Pbr038022	Not available	AGL24-like/DAM ^a
Pear	9e-24	Pbr019340	Not available	AGL24-like/DAM ^a
Pear	5e-19	scaffold293.0	Not available	–
Pear	1e-16	scaffold293.0	Not available	–
Pear	3e-14	scaffold293.0	Not available	–
Pear	1e-13	scaffold293.0	Not available	–
Peach	1e-17	ppa011123m	chr1:46353152-46360953	PpDAM4
Peach	4e-15	ppb017585m	chr1:46383267-46389606	PpDAM2
Peach	9e-13	ppa010822m	chr1:46342827-46351747	PpDAM5
Peach	1e-11	ppa010758m	chr1:46361693-46371335	PpDAM3
Peach	1e-08	ppa018667m	chr1:46390695-46398550	PpDAM1
Peach	5e-05	ppa010714m	chr1:46335129-46342354	PpDAM6

When the hit localizes inside a predicted gene, its annotation is shown in the right column. All significant hits ($E < 1e-04$) are shown

^a Overall results after BlastP of conceptual translations at NCBI's non-redundant protein database

^b Transcript model located 5 kb from the hit

^c Region containing the hit annotated as belonging to *MdDAM4* after the construction of the metacontig. See Fig. 3

containing *MdDAM* genes to the peach *EVG* locus using graphic alignment (Nix and Eisen 2005). As expected, the regions containing exons displayed the highest level of conservation (Fig. 3a, Online Resource 6). Interestingly, the second exon predicted in the genome for *MdDAM1* is not located in a region with high conservation. However, the alignment is in agreement with the *MdDAM1* structure inferred from our cDNA sequencing data, which does not predict a protein-coding exon region in this location. On the other hand, a window of highly conserved sequences could be identified between the intronic segments of apple and peach genes (Fig. 3a). This similarity still appears when the pairwise comparisons are shown with a maximum E value $< 10^{-15}$, a stringency cutoff that removes most of the coding region similarity results (Fig. 3b, arrows).

The genomic prediction for *MdDAM4* was only similar to the 3' end of the peach *DAM* genes (Fig. 3c). A closer look at the predicted structure suggests that the gene is truncated in the contig MDC027822.64. To determine if the remaining sequences for *MdDAM4* are available in another contig, we searched for *MdDAM4* segments in the apple genome and found a contig (MDC009761.283) that contains the 5' end of this gene. A manual concatenation of both contigs created a "metacontig" (see Online Resource 7 for the alignment between the ends of both contigs) that was graphically compared to the peach *EVG* locus and revealed the missing 5' portion of *MdDAM4*, including the conserved intronic sequence (Fig. 3d). The results for *MdDAM2* and *MdDAM3* resemble those found for *MdDAM1*, while *MdJa* and *MdJb* were less similar to the peach *DAM* genes (Online Resource 6).

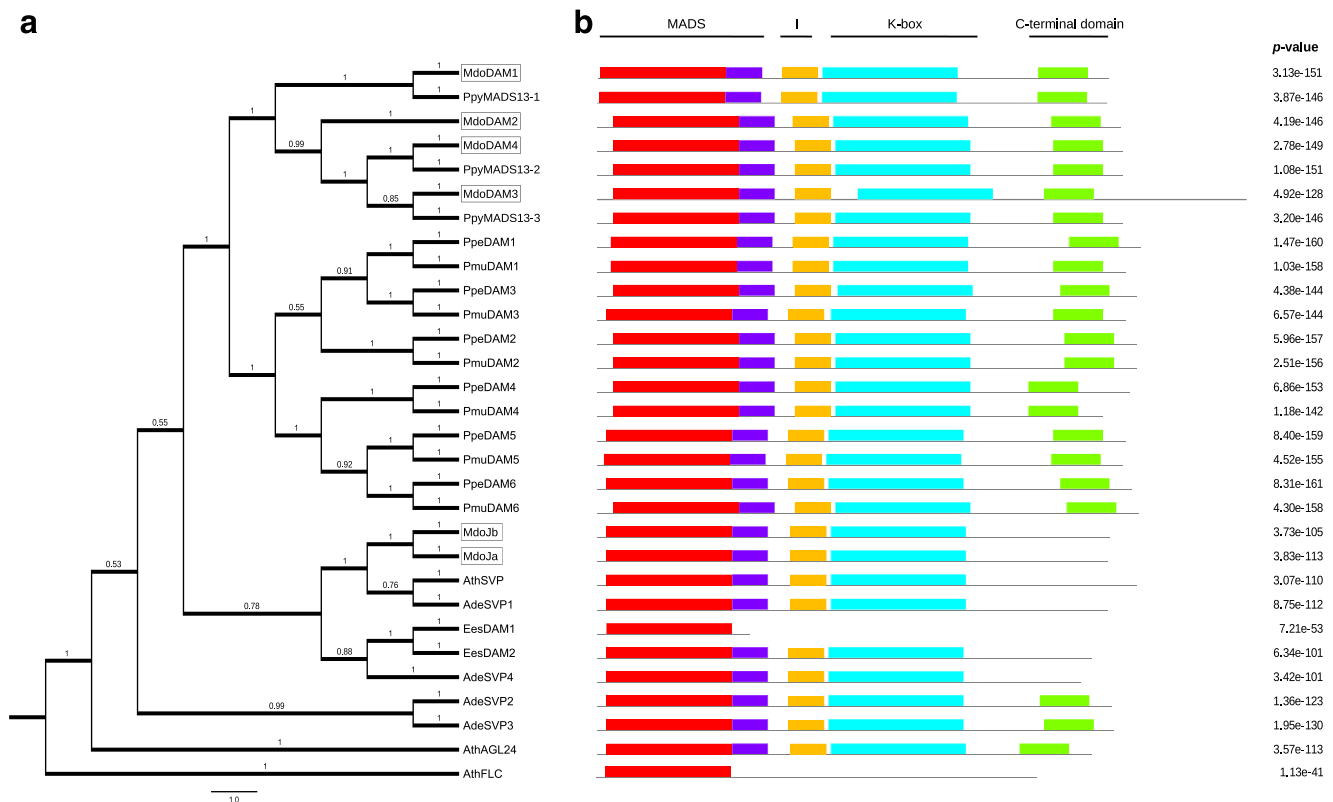


Fig. 2 Phylogenetic analysis and schematic motif structure of DAM proteins and *Arabidopsis* SVP and AGL24 protein sequences. **a** Phylogenetic relationships of full-length amino acid sequences. The *Arabidopsis* FLC was added for tree rooting. Nodal support values are given by the a posteriori probability values shown above the corresponding nodes (when >0.50), and the bar indicates the length for which the number of expected changes per site is equal to 1. Apple DAM

candidates (Mdo) are shown *inside rectangles*. **b** Schematic view of the conserved domains presented by their deduced protein sequences. Red and purple boxes, MADS domain; orange boxes, I domain; blue boxes, K domain; green boxes, C-terminal domain. Gray lines represent non-conserved sequences. Accession codes used in both analyses are depicted in OR4. *Ath Arabidopsis*, *Ade* kiwifruit, *Ees* leafy spurge, *Pmu* apricot, *Ppe* peach, *Ppy* pear

Given that a highly conserved sequence was identified between the intronic segments of apple and peach *DAM* genes, we examined the occurrence of this sequence in the apple and peach genomes. The conserved intronic sequence was used as a query for BlastN searches in apple contigs and peach scaffolds. Only six significant hits (E value $<10^{-4}$) were found in peach, all of them inside *DAM* introns. For apple, nine hits were retrieved and distributed in two small genomic windows in chromosomes 8 and 16. These hits were located inside the introns of *MdDAM1–4* as well as in other three uncharacterized gene models. BlastP searches at NCBI using these three genes retrieved *DAM*, *SVP-like*, and *AGL24-like* as best hits (Table 2). The conserved sequence was also found inside introns of the *DAM* genes of apricot and pear (Table 2), but the strawberry genome did not yield significant hits (data not shown). To evaluate the total extent of sequence conservation in the *DAM* intronic sequences, we performed multiple alignments using 500 bp up- and downstream of the conserved intronic segments from the apple and peach sequences. The alignment revealed that the segment is located in one of two highly conserved sequence intervals, with approximately 145 and 75 bp, respectively (Online

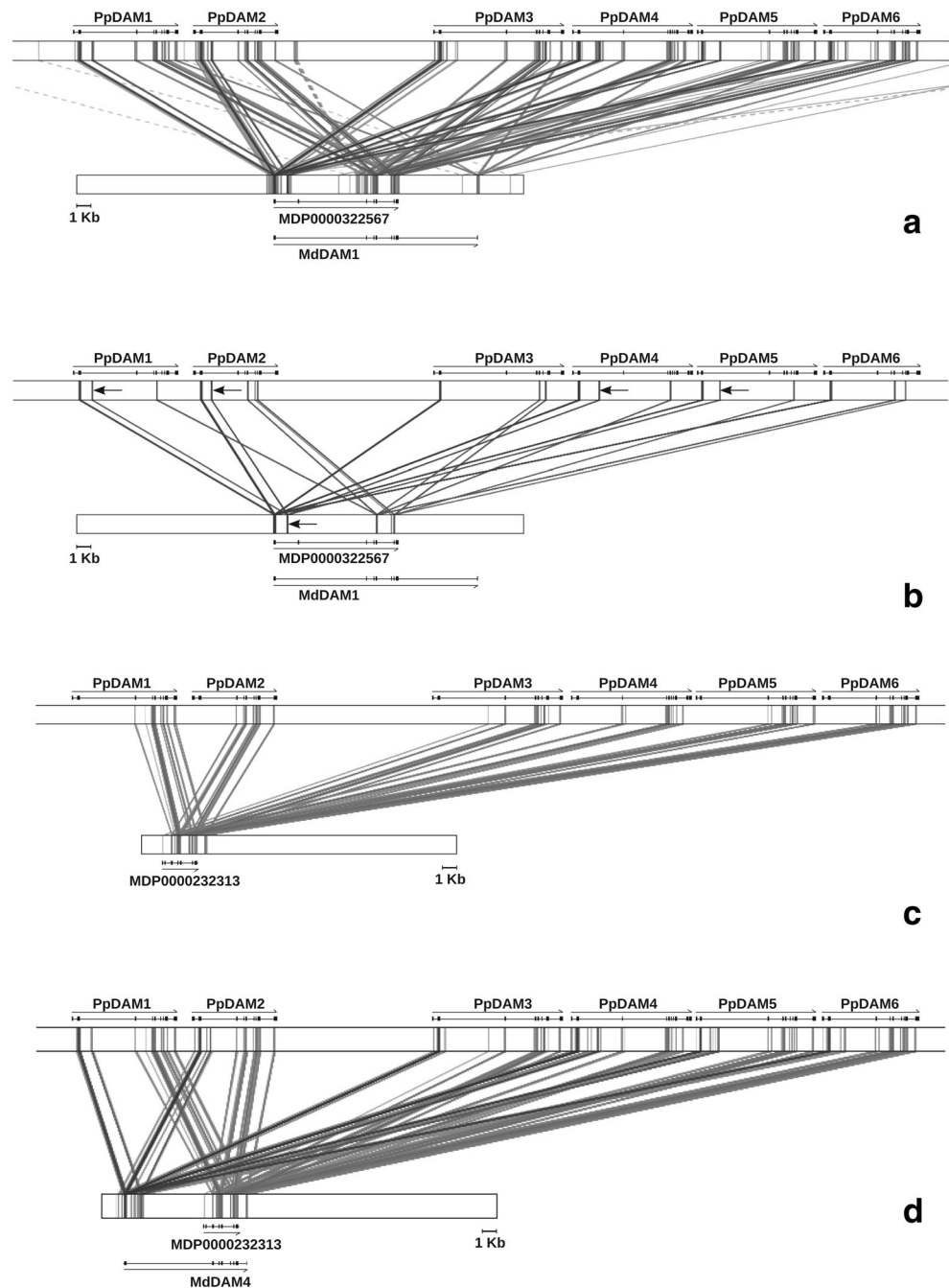
Resource 8, green boxes). They are separated by a less conserved nearly 89-bp-long sequence. The genomic locations of the conserved intronic sequences in apple and peach are shown below the alignment in the Online Resource 8.

MdDAM gene expression patterns

The transcript accumulation of apple *MdDAM* genes was quantified in several apple organs. The results show that *MdDAM1*, *MdDAM2*, *MdDAM3*, and *MdDAM4* were mainly expressed in dormant buds and seeds (Online Resource 9, samples A and S, respectively), with low transcript accumulation in vegetative organs. Interestingly, *MdDAM2* expression was 10 times higher in seeds than in dormant buds. Conversely, *MdJa* and *MdJb* were expressed in a similar fashion in dormant buds, growing buds, leaves, and seeds, with lower levels in fruits (Online Resource 9).

To evaluate the seasonal oscillation of *MdDAM* gene expression, messenger RNA (mRNA) levels were quantified in bud samples of three cultivars harvested over a growth-dormancy cycle. These genotypes were selected considering their contrasting CRs, where ‘Fuji Standard’

Fig. 3 Graphical alignment between the peach *EVG* locus (upper white bars, reference) and apple genome contigs (lower white bars) containing putative *DAM* genes. Solid lines indicate a significant similarity of 80-bp sequence windows for a maximum *E* value of $1e10^{-5}$, except for **b**, where the significance cutoff is $1e10^{-15}$. Dashed lines also represent the significant sequence similarity but denote the inverted sequences in relation to the reference. The predicted apple gene models (MDP0000322567 and MDP0000232313) and the sequenced coding regions (*MdDAM1* and *MdDAM4*) are represented below apple contig bars, where boxes and lines indicate exons and introns, respectively. Peach *DAM* genes (*PpDAM1* to *PpDAM6*) are represented in the same way above the reference bars. The apple genome contigs analyzed were the following: **a**, **b** MDC020688.360, **c** MDC027822.64, and **d** metacontig composed of the fusion of MDC027822.64 and MDC009761.283. Arrows point to the conserved intronic regions



and ‘Royal Gala’ need higher amounts of cold exposure compared to ‘Castel Gala’ (see the “Methods” section and Denardi and Seccon 2005; Botelho et al. 2006; Anzanello et al. 2014a). *MdDAM1*, *MdDAM3*, and *MdDAM4* showed peaks of transcript accumulation during winter (June and July) and low transcript levels during spring and summer. Interestingly, ‘Castel Gala’ samples showed an anticipated decrease in their transcript levels in comparison to both high CR cultivars. *MdDAM2* transcription peaked in the summer (February), with less expression in the course of the autumn and winter. Finally, *MdJa* and *MdJb* presented

less variation in their transcript amounts during the year (Fig. 4).

To investigate the effect of the controlled chilling exposure on the transcript accumulation of *MdDAM* genes and to compare this effect between contrasting CR cultivars, cuttings (brindles with terminal buds) from ‘Royal Gala’ and ‘Castel Gala’ trees were kept in a dark chamber at 6 °C for 1 or 5 weeks. At the time of harvesting, the ‘Royal Gala’ terminal buds were in deep dormancy, while the ‘Castel Gala’ buds were already in ecodormancy, as observed by budbreak rates under forcing conditions (Fig. 5). Extended cold exposure

increased the budbreak rate of ‘Royal Gala’ terminal buds but had little effect on the budbreak rate of ‘Castel Gala’ samples. Extended cold exposure decreased the gene expression of all *MdDAM* genes and *MdJa* in both cultivars, but not *MdJb*. The transcript level downregulation was more abrupt for *MdDAM2* and *MdDAM4*, which were barely detectable by real-time PCR after 5 weeks of cold treatment in both cultivars. Additionally, ‘Royal Gala’ and ‘Castel Gala’ buds showed very similar transcript accumulation levels for all genes. Finally, *MdJb* showed a small variation in the transcript levels during the cold treatment (Fig. 5).

In silico analyses of cis-elements in the promoter and in the intronic region of *MdDAM* genes

The promoter analyses of all six candidate *MdDAM* genes (~1.5 kb upstream of start codon) revealed the presence of

both unique and consensus *cis*-elements (Online Resource 10). Interestingly, *MdDAM1-4* showed two CARG-box (binding sites for MADS-box factors) motifs separated by less than 100 bp. In addition, we analyzed the whole largest intron of *MdDAM1-4* and, again, unique as well as consensus motifs were found (Online Resource 11). Consensus *cis*-elements include sites found in the promoters of the homeotic gene *AGAMOUS* (AG and AGAMOUS), response to dehydration and low temperature (ABRE-like), and CARG-box motifs (CARGCW8GAT). Once again, two CARG-box motifs were found within a window of less than 400 bp in *MdDAM1*, *MdDAM3*, and *MdDAM4*, although outside the conserved intronic region.

Finally, we further characterized the promoter and the conserved intronic region of *MdDAM1*, which presents high expression in endodormant buds (Falavigna et al. 2014, this work). Four unique *cis*-elements were found

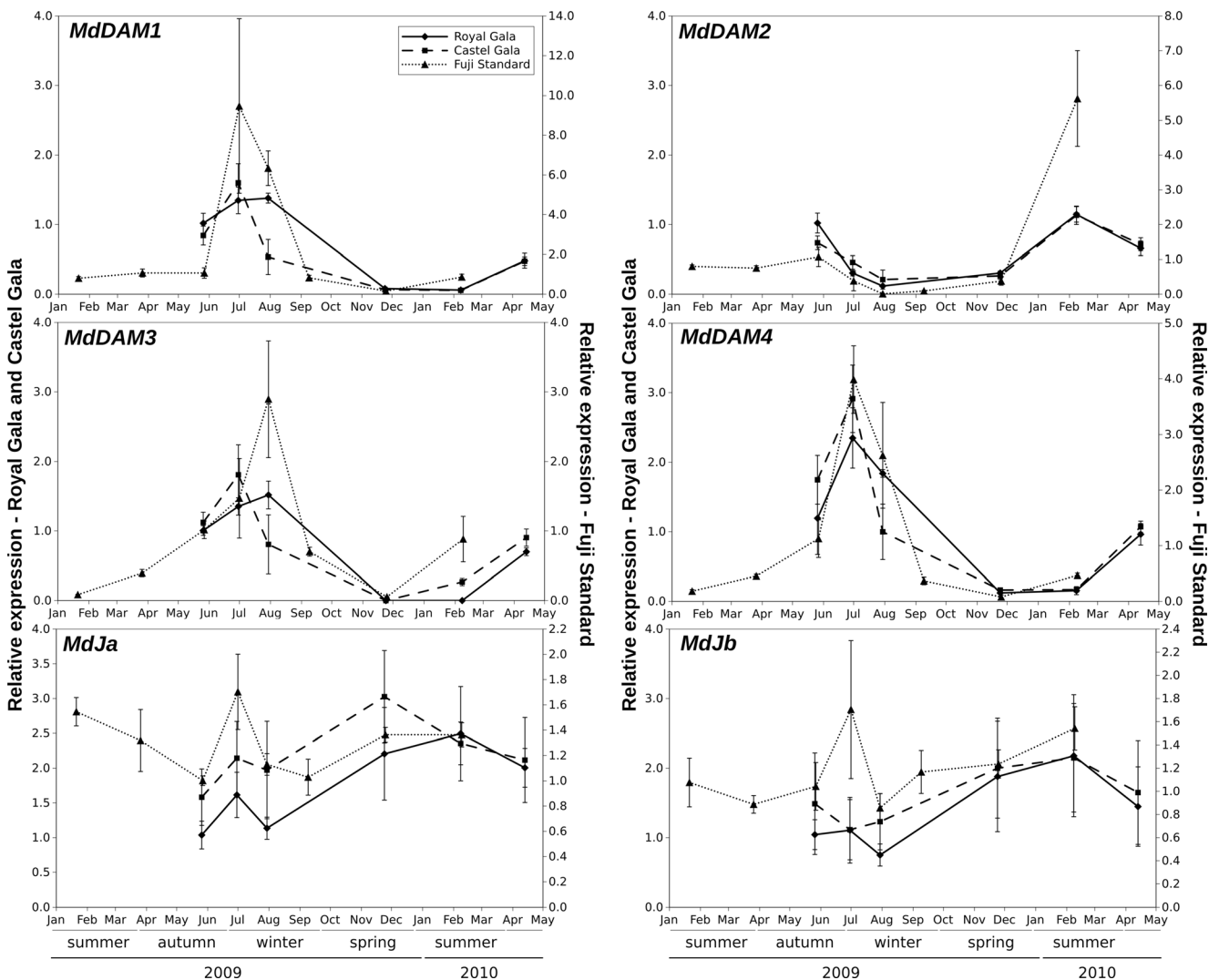


Fig. 4 Transcript accumulation of *MdDAM* genes in closed terminal buds of ‘Royal Gala,’ ‘Castel Gala,’ and ‘Fuji Standard’ over one growth-dormancy cycle. Months are

represented by their initial letters on the *x*-axis. Transcript levels are relative to May 2009 samples. Bars are standard errors of the mean (*n*=3)

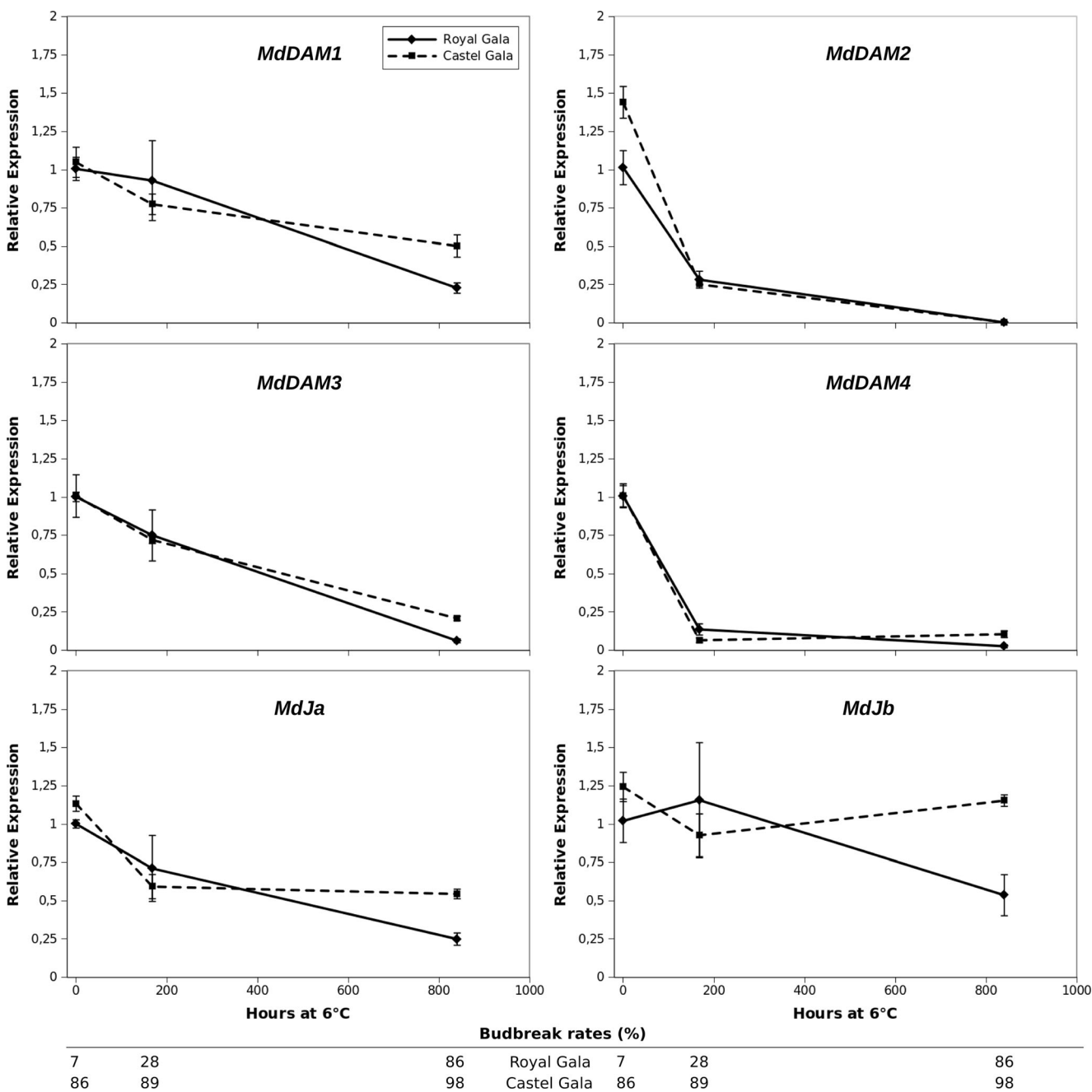


Fig. 5 Transcript accumulation of apple *DAM* candidates in closed terminal buds of two contrasting CR cultivars exposed to 0 week, 1 week (168 h), and 5 weeks (840 h) of chilling (6 °C). Transcript

levels are relative to ‘Royal Gala’ initial samples. Budbreak rates of samples analyzed are shown at the *bottom*. Bars are standard errors of the mean (*n*=3)

in its promoter region, including a CBF1 motif (specific binding of transcription factor CBF1-C-repeat binding factor/DREB1) and a LTREATLTI78 motif (a putative low-temperature responsive element; Fig. 6). A search for these motifs in the promoter regions of all predicted apple genes available (36,991 predicted genes; see the “Methods” section) revealed that only 154 CBF1 (0.44 % occurrence) and 1050 LTREATLTI78 (2.38 % occurrence) motifs were found (Online Resources 12 and 13). In the same way, the

highly conserved region of the largest intron from *MdDAM1* was screened (90 base pairs; Online Resource 8), revealing five consensus motifs (Online Resource 11). We searched for *cis*-elements in the conserved intronic sequences of peach and apricot *DAM* genes and identified only four out of these five motifs (data not shown). Interestingly, *Arabidopsis* response regulator 10 (ARR10), a binding site of the GARP family of plant Myb-related transcription factors, is only

present in some apple and pear *DAM* genes (Fig. 6). Moreover, only 641 apple genes (1.73 %) presented ARR10 in their promoter region (Online Resource 14). A gene ontology (GO) search was performed to identify biological processes related for the predicted genes that contain each of these three motifs (Online Resources 12 to 14). As a result, several similar enriched terms were recognized, with the “binding” term being the most general one (Fig. 6).

Discussion

The MADS-box family of transcription factors is known to comprise important regulators of growth and development in plants (Kaufmann et al. 2005; Dornelas et al. 2011). The sequencing of the apple genome revealed a relatively high number of MADS-box genes, possibly due to a recent genome duplication, and many of these are speculated to have roles in the development of the pome fruit, including genes coding for

the StMADS11/SVP-clade MADS-box transcription factors (Velasco et al. 2010). Recently, four reports identified genes belonging to this subfamily in apple (Falavigna et al. 2014; Wisniewski et al. 2015; Mimida et al. 2015; Nakano et al. 2015). The first gene, *MdDAM1*, was identified by our group in a suppression subtractive hybridization assay between endodormant and ecodormant buds (Falavigna et al. 2014). In later studies, two additional apple *DAM* genes were identified, and their expression patterns were associated with dormancy onset and release (Wisniewski et al. 2015; Mimida et al. 2015). Finally, Nakano et al. (2015) performed a functional analysis of two other members of this subfamily and showed that the ectopic expression of *MdJb* and, to a lesser extent, *MdJa* can restore the *JOINTLESS*-deficient tomato mutant. This finding shows that *MdJa* and *MdJb* may be involved in the development of functional pedicel abscission zones rather than with dormancy. In this work, we provide evidence that at least four out of these six apple StMADS11/SVP-clade MADS-box genes (*MdDAM1*, *MdDAM2*, *MdDAM3*, and *MdDAM4*) participate in dormancy regulation,

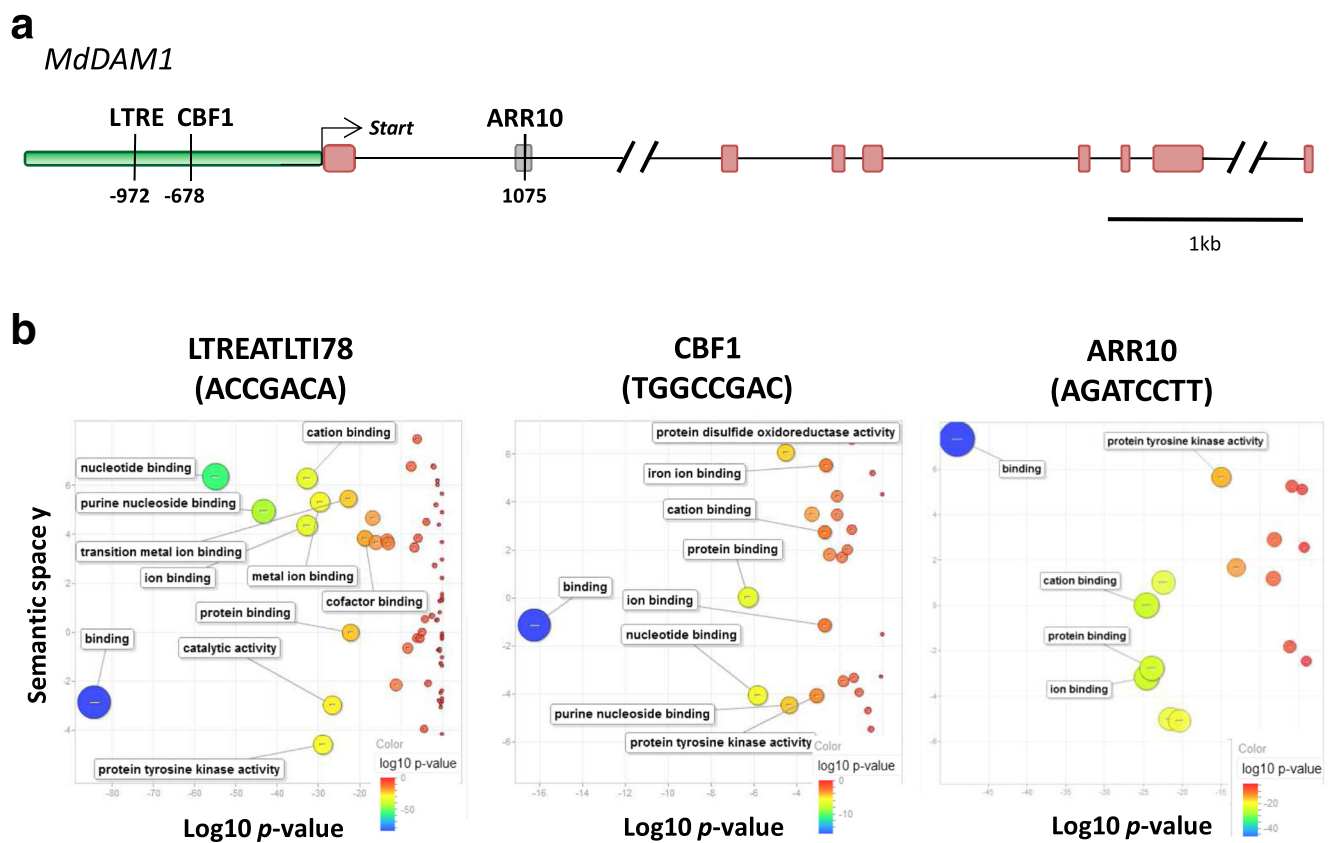


Fig. 6 Gene structure and *cis*-element analysis of *MdDAM1*. **a** The representative image is composed by the promoter region (up to 1.5 kb upstream from the start codon; green box), exons (red boxes), and introns (black lines). The gray box comprehends the conserved intronic region (see OR7). The three *cis*-elements characterized are shown by a vertical black line, along with the distance from the start codon. **b** GO analysis of the genes containing CBF1, LTREATLT178, and ARR10 *cis*-elements in

their promoter regions. The scatterplot shows the cluster representatives (i.e., the terms that remained following redundancy reduction) in a two-dimensional space derived by applying multidimensional scaling to a matrix of the GO term semantic similarities. The bubble color indicates the AgriGO-provided *p* value (see the legend in the upper right-hand corner); the size indicates the frequency of the GO term in the underlying GO analysis. The bubbles are larger for the more general terms

whereas the data collected for *MdJa* and *MdJb* corroborates the hypothesis that these genes may not be acting on dormancy control. Therefore, based on our findings and reports from the literature, we propose a unified nomenclature for *DAM* genes in apple (Table 1).

To gain knowledge of the possible roles of the six gene models identified, we quantified their transcript accumulation in different apple organs. Interestingly, *MdDAM1*, *MdDAM2*, *MdDAM3*, and *MdDAM4* expression was enriched in dormant buds and seeds (Online Resource 9). In peach, the expression of some bud dormancy-related genes is also detected in seeds, including *PpDAM1*, *PpDAM5*, and *PpDAM6*. Additionally, the mRNA levels of these *PpDAM* genes declined in the seeds after cold stratification (Leida et al. 2012). Our results are in agreement with the ones with peach, suggesting common mechanisms of bud and seed dormancy regulation.

Dormant buds of three distinct cultivars were sampled during a full year and were used to analyze the expression of apple *DAM* genes (Fig. 4). *MdDAM1*, *MdDAM3*, and *MdDAM4* showed a seasonal oscillating pattern of transcript accumulation, with a remarkable repression near budbreak (Fig. 4). The same peak of expression for *MdDAM1* was also observed in a standard dwarfing apple rootstock (Wisniewski et al. 2015) and in fruit-bearing terminal buds of apple cultivars ‘Jonathan’ and ‘Fuji’ (labeled as *MdDAMa* in Mimida et al. 2015). *MdDAM2* (labeled *MdDAMc* in Mimida et al. 2015) was more expressed during summer and autumn, as was observed in our data (Fig. 4). Similarly, peach and pear *DAM* genes exhibit clear seasonal patterns of transcript accumulation, with most genes showing increased mRNA levels during autumn and the beginning of the winter, followed by a decrease as the buds are exposed to low temperatures (Li et al. 2009; Saito et al. 2013; Ubi et al. 2010). Wisniewski et al. (2015) observed that *MdJb* (labeled by the authors as *MdDAM3*) presents a low fluctuation over the year.

Our results showed a clear repression of transcript levels during the late winter in field-harvested samples for *MdDAM1*, *MdDAM3*, and *MdDAM4* (Fig. 4). To investigate if these responses are an effect of chilling exposure, an experiment using controlled temperature treatment was carried out. When chilled under controlled conditions, buds from both contrasting CR cultivars showed a decrease of transcript accumulation for all tested genes, except *MdJb* (Fig. 5). Interestingly, *MdJb* also did not present a clear seasonal variation of mRNA levels (Fig. 4). The steady-state mRNA levels of apple *DAM* genes, as well as the rate of the transcript level repression by cold exposure, did not differ significantly between the cultivars tested, despite the clear differences of budbreak rates at the onset of the experiment (Fig. 5). This result was surprising, given that in field-harvested samples, *MdDAM1* and *MdDAM3* expression in ‘Castel Gala’ buds declined more rapidly during winter than

in ‘Royal Gala’ (Fig. 4). However, this result has to be considered with caution, because, in this experiment, the two apple cultivars were not at the same developmental stage and therefore cannot be strictly compared. In apricot, the expression of three (*PmDAM1*, *PmDAM2*, and *PmDAM3*) out of the six *DAM* genes under controlled cold exposure was not different between the two contrasting CR cultivars tested, despite the higher budbreak rates of the low CR buds (Sasaki et al. 2011). However, *PmDAM1* and *PmDAM3* expressions were different between the cultivars in field-harvested buds during winter (Sasaki et al. 2011), as was observed with *MdDAM1* and *MdDAM3* (Fig. 4; Falavigna et al. 2014). Similarly, peach *PpDAM3* was differentially expressed in the field but not under controlled conditions (Jiménez et al. 2010). Genotype-related differences of mRNA level repression rates during winter in some *DAM* genes may be dependent on other environmental cues in addition to cold exposure.

With the aim of better describing the *MdDAM* gene structures, we fully characterized them by an in silico analysis approach. We observed that, in addition to the high sequence similarity (Fig. 2), the genomic localization of several *MdDAM* genes found in this study provides good evidence for their putative role in dormancy. Three of them (*MdDAM2*, *MdDAM3*, and *MdDAM4*) are physically located very close to each other in chromosome 8 (in a 200-kb interval). Two other *DAM*-like predicted genes were also found inside this interval when the conserved intronic sequence (Online Resource 8) was used as a query in Blast searches against the apple genome (Table 2). These genes were not included in the analyses because they lack one or more typical MIKCC domains (data not shown). Apple chromosome 8 is highly syntenic to the end of peach chromosome 1, where the *DAM* genes were first mapped (Fig. 1; Bielenberg et al. 2008; Jung et al. 2012). This configuration of the apple *DAM* genes nearly in tandem resembles the peach *EVG* locus and may be a vestige remaining from an ancestor chromosome of this particular genomic region. *MdDAM1* seems to be an exception to this trend, given that it presents the conserved intron and expression profiles similar to those of *MdDAM2*, *MdDAM3*, and *MdDAM4*, although it is located in apple chromosome 16. *MdDAM1* is embedded in a block collinear to the *EVG* locus but was not included in the sequence of pairs of apple and peach genes that define the collinearity. Instead, a closely located *DAM*-like gene model (MDP0000255146) participates as a counterpart to *PpDAM1*. This predicted gene could be a functional apple *DAM*, although its sequence and structure are not as similar to the peach *DAM* genes as the ones characterized in this work. The presence of both *MdDAM1* and *DAM*-like genes in a block collinear to the *EVG* locus indicates a syntenic relationship between them and the peach *DAM* genes.

Our promoter analysis identified many unique and consensus motifs in the promoter region of all six candidate *MdDAM* genes (Online Resource 10). Interestingly, *MdDAM1–4* promoters displayed at least two CARG-box variants separated by less than 100 bp (Online Resource 10). The distance between the two *cis*-elements is essential, given that some MADS-box factors exert their transcriptional regulation by forming dimer complexes that bind to two CARG boxes separated by up to 400 bp (Mendes et al. 2013). This finding suggests that apple *DAM* genes can be directly regulated by themselves or by other MADS-box factors, although further investigation needs to be conducted to unveil their functional roles. We decided to deeply investigate the *MdDAM1* promoter, which presents high expression in endodormant buds (Falavigna et al. 2014). We found four unique *cis*-elements when compared to the other *MdDAM* members (Online Resource 10), including a low-temperature responsive element (LTRE) and the *cis*-element for the CBF1 transcription factor. In fact, a peach CBF1 was shown to regulate *MdDAM* genes in apple (Wisniewski et al. 2015). Additionally, a transient reporter assay in pear showed that the pear CBF2 can interact with the promoter of the pear *DAM* ortholog *PpMADS13-1* (Saito et al. 2015), as has already been suggested for *DAM* genes in other species (Horvath et al. 2010; Sasaki et al. 2011; Yamane et al. 2011). Taken together, these findings strongly suggest that some *DAM* genes integrate the cold regulon.

When the genomic regions containing *DAM* (or *DAM*-like) genes of apple and peach were compared, a strikingly conserved noncoding region was found inside their intronic regions (Fig. 3, Online Resource 6). As species diverge, the sequences of their orthologous genes accumulate differences. However, regions whose changes will likely compromise protein function, such as exons, remain more conserved (D'haeseleer 2006). Thus, the fact that this intronic region is more conserved than most *DAM* exonic sequences suggests that an important functional role is involved in its evolutionary maintenance. Interestingly, this sequence is not found in strawberry, a rosaceous species with well-delimited syntenic correlations with apple and peach that does not undergo a typical dormancy cycle (Illa et al. 2011; Jung et al. 2012).

In peach, changes in the CRs of cultivars have been observed in parallel to genomic insertions in the largest intron of *DAM* genes (Yamane et al. 2011; Zhebentyayeva et al. 2014). Our analysis showed at least two CARG-box variants in the largest introns of *MdDAM1*, *MdDAM3*, and *MdDAM4*, separated within a distance of 350 bp (Online Resource 11). MADS-box proteins could interact with these motifs to regulate their expression. One hypothesis for the changes in the CRs of the cultivars could be the alteration of *DAM* expression caused by modification of the distance between CARG boxes by genomic insertions (Mendes et al. 2013).

However, several pear genotypes with distinct CRs were screened for the presence of insertions in their largest intron, and no relation to dormancy could be drawn (Saito et al. 2013). Furthermore, the insertions in the *DAM5* intron described as correlated to the peach CR in the work of Zhebentyayeva et al. (2014) were not located inside the conserved sequence found in this work (data not shown). Thus, the precise effect of these insertions on *DAM* gene expression regulation remains unknown.

Finally, we analyzed the conserved intronic region presented by rosaceous *DAM* genes and identified several *cis*-elements related to light regulation (Online Resource 11). It is intriguing that some apple and pear *DAM* genes present a unique *cis*-element called ARR10 (Fig. 6), which does not appear in the conserved intron of *Prunus* species (Online Resource 8). This motif is the binding site of at least four type-B ARRs (ARR1, AAR2, AAR10, and ARR12), which have significant roles in modulating cytokinin signaling and the control of meristem cell division (Hosoda et al. 2002; Hill et al. 2013). Interestingly, although the ARR10 motif rarely appears (1.73 %) in the apple gene promoters analyzed, it is commonly presented in the promoter and in the whole intronic sequences of *MdDAM* genes (Online Resources 10 and 11). This finding suggests that cytokinin levels can indirectly repress apple *DAM* genes, given that this hormone is related to cell division and growth.

The role of intronic regions in the control of transcript accumulation is well documented in the *Arabidopsis AGAMOUS (AtAG)* gene. Reporter gene expression under the sole control of the *AtAG* promoter did not match the expected transcriptional pattern. However, this pattern is recovered when the reporter is controlled by the promoter and the largest intron (Sieburth and Meyerowitz 1997). Indeed, the presence of *cis*-elements acting as docking sites for other transcription factors in this intron is essential for the spatial and temporal control of *AtAG* expression (Hong et al. 2003). Functional analyses are needed to evaluate possible roles of conserved elements in introns of *DAM* genes over their expression, particularly the sequences found in this work for rosaceous homologs.

We identified and characterized four *DAM* genes in apple through extensive *in silico* analyses of their genomic structure. In addition, *MdDAM1*, *MdDAM2*, *MdDAM3*, and *MdDAM4* presented expression profiles with close resemblance to the ones reported for *DAM* genes of peach (Li et al. 2009), pear (Ubi et al. 2010; Saito et al. 2013), and apricot (Sasaki et al. 2011). These four genes also shared a highly conserved sequence in their largest intron with all the reported rosaceous *DAM* genes. This conserved sequence is a promising new feature of *DAM* genes. Combined, our results provide additional support to help unveil the conserved role of *DAM* genes in apple regarding endodormancy establishment and maintenance.

Methods

Plant material

Field-grown ‘Royal Gala’ and ‘Castel Gala’ plants were located in a commercial orchard (Papanduva, SC, Brazil; -26.434870 , -50.106103 , and 788 m altitude). Six-year-old ‘Royal Gala’ trees were cultivated on Marubakaido rootstock with M9 as an interstock. ‘Castel Gala’ samples were from 3-year-old trees grafted on M9 rootstock. ‘Fuji Standard’ trees were from an experimental field at Empresa de Pesquisa Agropecuária e Extensão Rural de Santa Catarina (Caçador, SC, Brazil; -26.836971 , -50.975246 , and 935 m altitude). The plants studied were 7-year-old trees grafted on M7.

Both ‘Royal Gala,’ ‘Gala Baigent,’ and ‘Castel Gala’ are phenologically stable cultivars derived from spontaneous bud sport mutations from trees of the original cv. Gala (‘Gala Standard’). ‘Royal Gala’ and ‘Gala Baigent’ trees have the same CR of ‘Gala Standard’ (around 600 CH), while their fruit skins show more concentration of red pigments. ‘Castel Gala’ fruits are identical in appearance to the ‘Gala Standard’ fruits, but the CR is approximately half the CH needed to break the ‘Gala Standard’ bud dormancy (Anzanello et al. 2014a). ‘Fuji Standard’ buds require over 1000 CH to break bud dormancy in Southern Brazil (Petri and Stucker 1988).

All bud samples were terminal buds in phenological stage A (closed) according to the Fleckinger scale (EPPO 1984). Sampling was always performed at 9:00 am for ‘Royal Gala’ and ‘Castel Gala’ and at 11:00 a.m. for ‘Fuji Standard.’ Field-harvested buds were immediately frozen in liquid N₂ and stored at -80 °C until analysis. Cold treatments were performed as previously described (Anzanello et al. 2014a, b). Briefly, 20–30 cm stem cuttings with terminal buds were cut and kept moistened until processing. Cuttings were surface-sterilized, air-dried, wrapped in plastic bags, and maintained at 6 °C in a BOD-type chamber in a vertical position in the dark. After 1 and 5 weeks, a batch of samples was either forced to budbreak (40 cuttings) or harvested for bud sampling (20 cuttings) as above. Forcing was done for 56 days in a growth chamber at 25 °C, with a 12-h photoperiod and 70 % relative humidity. The percent maximum budbreak was calculated by dividing the number of apical buds showing green tissue by the total number of viable apical buds.

The plant material for tissue/organ analysis was harvested from ‘Gala Baigent’ trees located at an experimental station of Empresa Brasileira de Pesquisa Agropecuária in Vacaria, RS, Brazil (-28.513777 , -50.881465 , and 972 m altitude). The trees were 3 years old and grafted on Marubakaido rootstocks with M9 as an interstock. Sample stages were selected following the Fleckinger scale (EPPO 1984): closed terminal buds (A stage), buds with green tips (C stage), pre-anthesis flowers and young leaves (E2 stage), whole-set fruits ~ 10 mm

in diameter and leaves (I stage), and unripe fruits ~ 40 mm in diameter that were divided into pulp and skin (J stage). We also sampled mature fruits ~ 70 mm in diameter, partitioning them into pulp, seeds, and skin (named M).

RNA extraction, cDNA synthesis, sequencing, and real-time PCR

The total RNA was extracted from samples composed of 10–20 buds by LiCl precipitation, as described by Zeng and Yang (2002), following the modifications of Reid et al. (2006). The integrity of the RNA samples was checked on 1 % agarose/ethidium bromide gels. RNase inhibitor (RNaseOUT, Invitrogen, Waltham, MA, USA) was added to samples (1–2 μ l) before storing at -80 °C. Contaminating DNAs were removed with the Turbo DNA-free Kit (Applied Biosystems). Synthesis of cDNA was done from 1 μ g DNA-free total RNA using the GeneAmp RNA PCR Core Kit (Applied Biosystems) with oligo dT₁₆ (included in the kit), according to the manufacturer’s instructions. Assessment of genomic DNA contamination was done by PCR amplification of cDNAs with a primer pair for a WD40 domain containing gene (Perini et al. 2014) spanning an intron.

Real-time PCR analysis was performed as in Falavigna et al. (2014). The only difference is that the annealing temperature for *MdDAM3* was set to 55 °C for all experiments except the tissue/organ assays, performed as described and with a different set of primers. Amplification efficiencies were determined using LinRegPCR v. 2012.0 (Ruijter et al. 2009). Relative expression levels were calculated from amplification data using Pfaffl mathematics (Pfaffl 2001). The reference genes used were chosen among the most stable for apple buds (Perini et al. 2014). The reference gene adopted for samples exposed to cold under controlled conditions was *ACCUMULATION AND REPLICATION OF CHLOROPLAST (ARC5)*, and all other analyses employed *MALATE DEHYDROGENASE (MDH)* as the reference gene.

For the sequencing of apple *DAM* candidates, the oligonucleotide sequences designed for real-time PCR analyses were used as gene-specific primers for RACE using the SMARTer RACE cDNA Amplification Kit (Clontech, Mountain View, USA), according to the manufacturer’s instructions. RACE products were sequenced by Sanger’s method using an *ABI-PRISM 3100 Genetic Analyzer* and associated chemistry (Applied Biosystems).

Phylogenetic analysis and conserved protein motif analyses

Complete amino acid sequences of DAM proteins from apple, peach, apricot, pear, leafy spurge, kiwi, and StMADS11/SVP-clade MIKCC proteins from *A. thaliana*, with *A. thaliana* FLC as the out-group, were aligned using Muscle (Edgar 2004).

The accession numbers of all sequences used are presented in Online Resource 5. The phylogenetic tree was inferred using MrBayes version 3.2.1, employing the mixed amino acid substitution model in default settings (Ronquist et al. 2012). We performed a 1,000,000 generation run, sampled every 100 generations and the first 25 % of the trees were discarded as burn-in. The remaining ones were summarized in a consensus tree that was visualized and edited using FigTree v.1.4 (<http://tree.bio.ed.ac.uk/software/figtree/>).

Protein sequences from the translations of resequenced coding regions for MdDAM1, MdDAM2, MdDAM4, and MdJa, as well as translations from apple genome versions of MdDAM3 and MdJb (genome codes MDP0000527190 and MDP0000209705, respectively), were searched for conserved protein motif discovery using the MEME software version 4.10.1 (Bailey et al. 2009). Along with the apple DAM-like translations, DAM protein sequences from the peach, apricot, pear, kiwi, and leafy spurge, as well as the StMADS11/SVP-clade proteins SVP and AGL24 from *Arabidopsis*, were used as inputs. The software parameters were set to the defaults, except for the following: motif distribution parameter set to any number of repetitions, maximum number of motifs equal to 5, presented in at least 19 sequences, and motif width between 12 and 100 amino acids.

Sequence alignment

Contigs (codes MDC020948.189, MDC020688.360, MDC008471.150, MDC017035.448, MDC016474.222, and MDC027822.64, available at <http://www.rosaceae.org>) bearing candidate *MdDAM* genes were aligned to the *EVG* locus using the GATAligner software (Nix and Eisen 2005). The parameters used were the defaults, except for the adoption of an 80-bp sequence window. Approximately 1000 bp of each apple and peach intron sequence that contains the conserved regions identified were aligned using ClustalW (Larkin et al. 2007), and the resulting alignment was plotted using JalView (Waterhouse et al. 2009). For the collinearity analyses, all the proteins mapped to peach chromosome 1 were used as queries for BlastP against the predicted apple proteins. Only the top five matches from the results of each query were used for a maximum *E* value $<1e^{-10}$. These results and the gene positions were used as inputs for the MCScanX algorithm (Wang et al. 2012). The “family circle plotter” utility from the same software was used for the plot construction. The resulting image was converted to grayscale and improved for chromosome drawing and labeling in a vector graphics editor.

Cis-element discovery and gene ontology analyses

All *cis*-element searches were performed using the Web tool PlantPAN version 1.0 (*Plant Promoter Analysis Navigator*;

<http://plantpan.mbc.nctu.edu.tw>). For the *MdDAM* promoter region analysis, ~1.5 kb upstream of the start codon was used. Additionally, the sequence of the largest *MdDAM* intron or only 90 bp of the highly conserved intronic region of *MdDAM1* was used in the *cis*-element motif analysis. Only the given strand was considered, and only *cis*-elements larger than six nucleotides were used in further analyses. CBF1, LRTEATLTI78, and ARR10 motifs were used in a genome-wide search in the promoter regions of all apple genes predicted in the 1.0 version of the genome. Promoter sequences (1 kb upstream of predicted start codons) were extracted from genome data using BEDTools (Quinlan and Hall 2010) and in-house Perl scripts. The sequences extracted (36,991) were screened for conserved *cis*-elements using FIMO (Grant et al. 2011), available at <http://meme.nbcr.net/meme/tools/fimo>, with a match *p* value of less than e^{-4} and the given strand only. Genes with these motifs were scanned for enrichment of GO terms using agriGO (Du et al. 2010), available at <http://bioinfo.cau.edu.cn/agriGO/>, and REVIGO (Supek et al. 2011), available at <http://revigo.irb.hr/>. Only terms with a *p* value lower than $\log 10^{-1}$ were analyzed. Consensus and unique motifs were analyzed using online software available at <http://bioinformatics.psb.ugent.be/webtools/Venn/>.

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Authors' contributions DDP, VSF, PP, RA, FBF, HPS, and LFR performed the samplings in the field. PP and RA harvested the samples under controlled conditions. DDP, VSF, PP, and VB isolated the mRNA and performed the real-time PCR. VSF and PP searched the apple genome for *DAM* orthologs. DDP generated and sequenced the RACE products and performed the graphic alignments and collinearity analyses. VSF constructed the phylogenetic trees. RAA performed the *cis*-element and gene ontology analyses. DDP, VSF, RAA, and LFR wrote the manuscript. HPS, FBF, PRDO, and LFR conceived of and designed the experiments. All authors read and approved the final manuscript.

Data archiving statement The nucleotide sequences generated in this work were submitted to the GenBank database under accession numbers KT582786, KT582787, KT582788, and KT582789 (Online Resource 5).

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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