

Conserved MYC transcription factors play a key role in jasmonate signaling both in tomato and *Arabidopsis*

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Jasmonates (JA) are important regulators of plant defense responses that activate expression of many wound-induced genes including the tomato proteinase inhibitor II (*pin2*) and leucine aminopeptidase (*LAP*) genes. Elements required for JA induction of the *LAP* gene are all present in the -317 to -78 proximal promoter region. Using yeast one-hybrid screening, we have identified the bHLH-leu zipper JAMYC2 and JAMYC10 proteins, specifically recognizing a T/G-box AACGTG motif in this promoter fragment. Mutation of the G-box element decreases JA-responsive *LAP* promoter expression. Expression of *JAMYC2* and *JAMYC10* is induced by JA, with a kinetics that precedes that of the *LAP* or *pin2* transcripts. JAMYC overexpression enhanced JA-induced expression of these defense genes in potato, but did not result in constitutive transcript accumulation. Using footprinting assays, an additional protected element was identified, located directly adjacent to the T/G-box motif. Mutation of this element abolishes JA response, showing that recognition of this duplicated element is also required for gene expression. Knockout mutants in the *AtMYC2* homolog gene of *Arabidopsis* are insensitive to JA and exhibit a decreased activation of the JA-responsive genes *AtVSP* and *JR1*. Activation of the *PDF1.2* and *b-CHI*, ethylene/JA-responsive genes, is, however, increased in these mutants. These results show that the JAMYC/AtMYC2 transcription factors function as members of a MYC-based regulatory system conserved in dicotyledonous plants with a key role in JA-induced defense gene activation.

[Keywords: *pin2*; *LAP*; wounding; herbivore attack; JA cross-talk]

Received January 22, 2004; revised version accepted May 5, 2004.

Plants respond to insect attack and wounding by activating a set of genes involved in herbivore deterrence, wound-healing, and defense against pathogen infection (Ryan 2000; León et al. 2001; Li et al. 2002; Turner et al. 2002). Activation of wound-induced defense genes involves signal transduction pathways that operate both locally at the site of wounding and systemically in undamaged leaves (Ryan 2000). Mechanical wounding or herbivore attack to tomato or potato leaflets results in systemic accumulation of multiple defense-related proteins, including proteinase inhibitors (PI), exopeptidases like leucine aminopeptidase (*LAP*), and components of the wound signal pathway (Ryan 2000). Wound-induced expression of these defense genes is controlled by the jasmonate family of signaling molecules (Ryan 2000; Turner et al. 2002). Genetic analysis in tomato has indicated that the 18-amino-acid peptide systemin and its precursor protein, prosystemin, are upstream components of the signaling cascade involving systemic induc-

tion of jasmonates (Howe and Ryan 1999; Li et al. 2001, 2002). Recognition of systemin by its 160-kDa receptor LRR-receptor-like kinase (Scheer and Ryan 2002) activates a cascade of intracellular signaling events that leads to the release of linolenic acid from membrane lipids and the synthesis of jasmonates via the octadecanoid pathway (Schaller 2001; Howe and Schilmiller 2002).

A role for jasmonates in intercellular signaling is supported by the fact that application of JA to one leaf induces PI expression in distal untreated leaves (Ryan 2000). Evidence for a function of JA as a long-distance signal for systemic wound-activation has also been obtained from grafting experiments using mutants deficient in JA biosynthesis or JA perception. These studies indicated that the peptide systemin might not function as a long-distance signal, although it is required for production or translocation of the systemic signal to unwounded leaves (Li et al. 2002; Lee and Howe 2003).

In *Arabidopsis*, wounding activates independent signaling pathways regulating different sets of target genes either at the wound site or in distal leaves (Rojo et al. 1999). JA induces systemic expression of wound-responsive genes such as *VSP*, *JR1*, or *Thi1.2* (Berger et al. 1995; Titarenko et al. 1997; Bohlmann et al. 1998), but induc-

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Article and publication are at <http://www.genesdev.org/cgi/doi/10.1101/gad.297704>.

tion of these genes in the wounded leaves is negatively regulated by local synthesis of ethylene (Rojo et al. 1999). JA and ethylene, however, synergistically cooperate to activate expression of basic PR proteins such as *b-CHI*, *PR3*, and *PDF1.2* (Xu et al. 1994; Penninckx et al. 1998). Pathogen-induced expression of *b-CHI* and *PDF1.2* is blocked in mutants affected in their response to JA (*coi1-1* and *jar1*) or ethylene (*ein2-1* and *etr1-1*), these mutants being more susceptible to the attack by different fungal pathogens (Staswick et al. 1998; Thomma et al. 1999). These differences in signaling may reflect different mechanisms that evolved in each plant to optimize spatial and temporal defense-gene expression, although a better characterization of the mechanisms integrating JA and ethylene signaling is still needed.

Despite the vast amount of information available for the primary signalling components (for review, see Schaller 2001; Turner et al. 2002), limited data exist on the latter signaling steps leading to defense gene activation in response to JA. Mutant screens in *Arabidopsis* for insensitivity to JA and coronatine or altered JA-regulated gene expression have identified only a few genes, including *coi1* (Feys et al. 1994), *jar1* (Staswick et al. 2002), *jin1* and *jin4* (allelic to *jar1*; Berger et al. 1996), *cev1* (Ellis and Turner 2001), and the *jasmonate underexpressing* mutations *jue1*, *jue2*, and *jue3* (Jensen et al. 2002).

Upstream regions required for JA-induced gene expression have been identified in the promoters of several JA-regulated genes (Kim et al. 1992; Ishikawa et al. 1994; Samach et al. 1995; Ruíz-Rivero and Prat 1998), but the transcription factors binding to these *cis*-elements have not yet been isolated. JA-responsive elements have also been identified in the promoter regions of the barley *LOX1* (Rouster et al. 1997) and the soybean *VspB* genes (Mason et al. 1993), although a consensus element could not be deduced from the identified sequences. Whereas a G-box element mediates JA-regulated expression of the potato *pin2* and soybean *VspB* genes, a TGACG-related motif was identified in the barley *LOX1* gene (Rouster et al. 1997) or the *nos*/minimal 35S promoters (Xiang et al. 1996).

In *Arabidopsis*, constitutive expression of the downstream component of the ethylene/JA-signaling pathway *ethylene response factor 1* (*ERF1*; Solano et al. 1998) increases resistance to several necrotrophic pathogens (Berrocal-Lobo et al. 2002). Overexpression of the AP2 domain ERF1 factor rescues the defense response defects of the *coi1* and *ein2* mutations and leads to constitutive activation of a large number of ethylene/JA-induced genes, including the *PDF1.2* and *b-CHI* genes (Lorenzo et al. 2003). Interestingly, a GCC-box-like element like that recognized by ERF1 (Fujimoto et al. 2000) was also identified as a JA- and elicitor-responsive element involved in regulation of the terpenoid indole alkaloid (TIA) biosynthetic genes in *Catharantus roseus* (Menke et al. 1999). This element is bound by the elicitor- and JA-induced AP2 transcription factors ORCA2 and ORCA3 (Menke et al. 1999; van der Fits and Memelink 2000). Overexpression of ORCA3 leads to increased accumulation of terpenoid indole alkaloids (van der Fits

and Memelink 2000, 2001), therefore suggesting a conserved role of these AP2/ERF-transcription factors in ethylene (elicitor)/JA-dependent gene activation.

Here we describe the isolation of two MYC regulatory proteins that bind to JA-responsive elements in the tomato *LAP* and *pin2* promoters. Elements required for JA induction of the *LAP* gene are present in the -317 to -78 proximal promoter region (Ruíz-Rivero and Prat 1998). Using yeast one-hybrid interaction assays, we have isolated two cDNA clones, *JAMYC2* and *JAMYC10*, encoding bHLH-Leu zipper DNA-binding proteins that specifically recognize a T/G-box motif in this promoter region. We have used transgenic potatoes to analyze the function of the T/G-box *cis*-acting element and the *trans*-activation ability of these JAMYC transcription factors, because of the faster generation of transformants in this plant species. Potato plants are, indeed, suitable for these studies, as they are genetically very close to tomato, and systemic wound- and JA-induced expression of the *LAP*, *pin2*, as well as other defense-response genes is highly conserved in these two species. *Arabidopsis* lines with T-DNA insertions in the *AtMYC2* gene, encoding the closest homolog to the tomato JAMYC regulatory proteins, are insensitive to JA and exhibit an altered pattern of activation of the *VSP*, *PDF1.2*, and *b-CHI* genes. These findings demonstrate a role of JAMYC/AtMYC2 in JA-induced gene activation, these proteins functioning as conserved master switch regulators that activate expression of differentially JA-regulated genes while repressing expression of genes regulated by the ethylene/JA pathway.

Results

Isolation of cDNAs encoding LAP promoter DNA-binding proteins

Tomato *LAP* is induced in response to wounding and JA treatment and is thought to play a role in protein turnover during defense gene activation (Hildmann et al. 1992; Pautot et al. 1993). This exopeptidase is encoded by a gene family comprised of at least three genes (Ruíz-Rivero and Prat 1998). Previous work in our laboratory has led to the isolation of two genomic clones encoding *LAP* that share nearly identical promoter sequences up to position -317 relative to the ATG. Transcriptional promoter fusions to the *GUS* reporter gene showed that these were active gene copies, driving constitutive *GUS* expression in flowers and wound- or MeJA-induced expression in leaves (Ruíz-Rivero and Prat 1998). A promoter deletion down to this conserved region (-317*LAP*) was still able to direct MeJA-induced *GUS* expression, indicating that elements required for JA-responsive gene expression are present in this region. A -317 to -78 fragment fused to the minimal CaMV 35S promoter was also able to confer JA responsiveness to this minimal promoter (Ruíz-Rivero and Prat 1998), this region therefore used as bait in a yeast one-hybrid screening to search for transcription factors that mediate JA response of the *LAP* gene. A yeast YM4271 strain containing this promoter

fragment fused to the *HIS3* and β -*GAL* selection markers was used to screen for specific DNA-binding proteins in a tomato JA-induced leaf cDNA library fused to the GAL4 activation domain. Screening of 2 million transformants yielded 18 positive clones. Of these, 17 encoded proteins with a basic-helix-loop-helix (bHLH) cMYC DNA-binding domain and one encoded a transcription factor of the WRKY family, with a zinc finger DNA-binding domain exclusive to plants (Eulgem et al. 2000). Alignment of the MYC-deduced amino sequences revealed that these clones correspond to two related genes, designated as *JAMYC2* and *JAMYC10*. The deduced JAMYC amino acid sequences shared highest homology with *LEJA3* of tomato (AF011557), a partial cDNA isolated in one-hybrid screening studies using the *threonine deaminase (Tda)* promoter; with the PG1 and PG2 (*Phaseolin G-box-binding proteins*) regulatory factors from pea (Kawagoe and Murai 1996), and with the *AtMYC2* gene from *Arabidopsis*, reported to function in drought and ABA signaling (Abe et al. 2003).

Full-length clones (Fig. 1A) were isolated by screening a λ -ZAP library. *JAMYC2* and *JAMYC10* encode proteins

with a molecular mass of 75 kDa and 70 kDa, respectively, and a DNA-binding/dimerization domain consisting of a bHLH region followed by a Leu-zipper protein-protein interaction domain located at the C-terminal part of the protein. A conserved acidic region postulated to function as the transcriptional activation domain (Abe et al. 1997) is located in the N-terminal region.

JAMYC2 and *JAMYC10* expression is induced by wounding and JA treatment

JAMYC mRNA levels were analyzed after wound induction and JA application. As shown in Figure 1B, basal levels of *JAMYC2* and *JAMYC10* mRNAs were detected in control untreated leaves, but a rapid induction of these transcripts was observed after wounding or JA treatment. High levels of expression of these transcripts were observed as early as 30 min after JA addition, indicating that these transcription factors correspond to early jasmonate-responsive genes. A biphasic induction pattern in response to both wounding or JA application (less mRNA is detected at 2 h as compared with 30 min

A

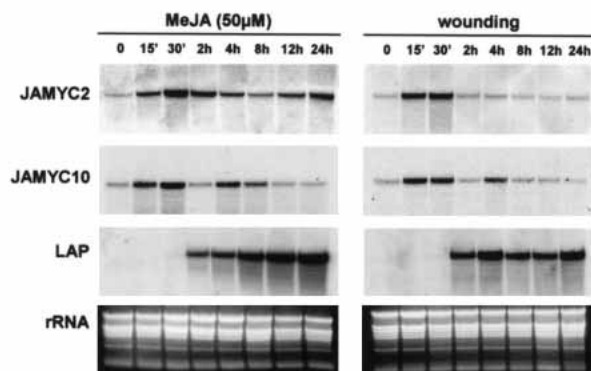
JAMYC2

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YAIWFQSSVDFSSPVLGWDGYYKGEEDKAKRKLAVSSPAYIAEQEHRKVLRE
LNSLISGAPAGTDDAVDEVTDTEWFFLISMTQSFVNGSGLPGQALYSSSPIWVAG
TEKLAASHCERVRAQGGFLQITVICIPISANGVVELGSTEIVLIVSSDLMNKRVLFPN
FNDLGGSGWAVQPESDPSALWLETPSSSGMEVRESLNTVQTNSVPSNSNKQIAY
ANENNHQSGNGQSCYNLQQQNNPPQQQTQGFRTRELNFSFEGFDGSSNRNGNASL
SCKPESGEIILNFGDSTKKSASSANVNLFTQQSQFQAVEENNNKKNKRSATSRGNS
EEGMLSFVSGTLPSSQMKSGGGGDESEHSDLEASVVKEDSSRVVEPEKRPKR
GRKPANGREEPLNHVEAERQRREKLNQRFYALRAVVPNVSRMDKASLLGDAISYIN
ELKSKLQNTESDKEDLRSQIEDLKESRRPGPPPPNQLKIGGKIVDVIDVKIIG
WDAMIGIQCNKKNHPAARLMAALMELDLVHHSVSVVNDLMIQQATVKMGSRHYT
EEQLRVALKSKIAETPLESR*
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JAMYC10

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MTDYRLWNSNTTNTSDDNMMDAFLSSDPSSFPNRTSISPTPVNGGVGETMPPFF
NQESLQORLQALIDGARESWAYAIWFQSSSTDFATPVLGWDGYYKGEENKNRK
RASSTNFVAEQEHRKVLRELNLSLISGQATGAGSGDDAVDEVTDTEWFFLI
SMTQSFANGNGLPGLAMYSSSPIWVTGTEKLAGSQCERARQAQGGFLQITVICIPSA
NGVVELGSTEIVLIVSSDLMNKVKYLFNFDIMGSVTSGSGSCAVHPEPDPALWL
TDPSSVVEAKDSLINSRRDVLVFNENENGTQNHQSQQTQGFRTKELNFSG
YFGDSSSTRNKNNGSSISCKPETREIILNFGDSSKKSGLFSGQSQFPGTGLGLME
ENKNNKRSLSLRGNNEKMLSFVSGVILPTSTMGKSGGGGNFDSLEASVVK
AIVEPERKPRKGRKPANGREEPLNHVEAERQRREKLNQRFYALRAVVPNVSRMDK
ASLLGDAIAYINELKSKVQNSDLDKEELRSQIESLRKELANKGSSNYSSPNSQD
LKIVDMIDVKVIGWDAMIRIQCSKKNHPAARLMAALKDLDLVHHSVSVVNDLMI
IQQATVKMGSRLYAQELTIALTSKFAESR*
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B



C

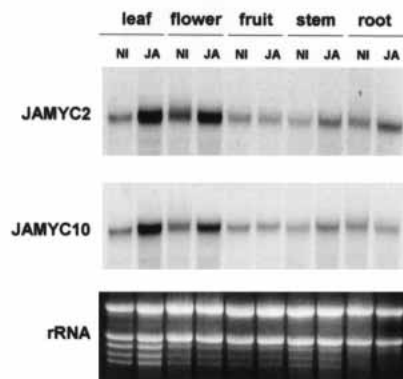


Figure 1. Amino acid sequences and patterns of expression of the JAMYC bHLH-Zip transcription factors. (A) *JAMYC2* and *JAMYC10* deduced amino acid sequences. The bHLH-Leu Zip domain is shown as a gray box, and the acidic region is underlined. The start point of the partial clones isolated by one hybrid screening is indicated as an arrowhead. (B) Northern analysis of tomato leaves induced by wounding or application of MeJA. (C) Pattern of tissue-specific expression at 30 min of MeJA treatment. Each lane was loaded with 30 μ g of total RNA. Blots were hybridized with probes corresponding to the 3'-noncoding regions of *JAMYC2* or *JAMYC10*, or to the *LAP* cDNA clone as indicated. Equal loading was verified by EtBr staining of the gel.

or 4 h) was consistently detected for transcript *JAMYC10*, with levels of this transcript returned to control levels by 24 h after induction. Hybridization of the same blots with an *LAP* cDNA probe showed that induction of the *LAP* transcript starts by 2 h of treatment, with maximal levels of mRNA observed after 12–24 h of exposure to JA (Fig. 1B), peaks of *JAMYC2* and *JAMYC10* expression therefore preceding maximal induction of the *LAP* gene. These results are consistent with a possible function of these transcription factors in *LAP* gene regulation.

Analysis of the tissue-specific pattern of expression of these genes showed elevated levels of mRNA in flowers. JA application induced accumulation of these transcripts in leaves and flowers and to a lower extent in stem, but did not modify basal levels of *JAMYC* mRNAs in fruit (Fig. 1C).

The JAMYC proteins recognize a T/G-box motif in the LAP promoter and a G-box motif required for JA induction of the pin2 promoter

To map the recognition site for these transcription factors, hydroxyl radical interference experiments (Hayes and Tullius 1989) of the –125 *LAP* fragment were performed with His-tagged fusions of the *JAMYC2* and *JAMYC10* proteins. As shown in Figure 2A, seven nucleotides were found to be protected from hydroxylation in these assays, with identical protected windows observed for both slower and faster migrating complexes. This footprinted region identifies an AAACGTG element (T/G-box motif) as the binding site for the *JAMYC* transcription factors. Noteworthy, G-box (CACGTG) sequence motifs are preferential targets for cMYC bHLH DNA-binding proteins (Toledo-Ortiz et al. 2003). These results were further confirmed by mobility shift (EMSA) studies using oligonucleotide probes with intact or mutagenized AAACGTG motifs. As shown in Figure 2B, oligonucleotides containing an intact AAACGTG element (*LAP*^{T/G-BOX}) efficiently competed for complex formation. Competition was totally abolished by replacement of the C and T residues within the AAACGTG motif (*LAP*^{t/g-box}), indicating that the AAACGTG element is critical for *JAMYC* recognition.

JAMYC recognition of the T/G-box motif prompted us to investigate whether these transcription factors would also bind a G-box (CACGTG) element present in the *pin2* promoter and required for JA-induced expression of this gene (Kim et al. 1992). EMSA assays were performed with pairs of oligonucleotides complementary to this promoter segment, containing an intact (*PIN2*^{G-BOX}) or mutagenized (*PIN2*^{g-box}) copy of the G-box element. Binding of the *JAMYC* factors to the AAACGTG *LAP* motif (*LAP*^{T/G-BOX}) was strongly competed by incubation with the wild-type *pin2* promoter segment (*PIN2*^{G-BOX}), and this competition was completely abolished by mutation of the G-box element (*PIN2*^{g-box}; Fig. 2C). Identical results were obtained when these oligonucleotide pairs were used as probes (data not shown), demonstrat-

ing a specific recognition of the tomato *pin2* promoter G-box element by these transcription factors.

Site-directed mutagenesis of the LAP promoter T/G-box motif results in reduced JA activation

To determine whether the T/G-box motif functions as a JA-responsive element, transgenic potato lines were obtained bearing a mutated version of the –317 to –3 *LAP* promoter region fused to the *GUS* reporter gene (Fig. 2D). Mutation of the T/G-box element caused a strong decrease in the levels of JA-induced activation of this promoter (cf. levels of *GUS* activity measured in the leaves of the –317*LAP* or the –317*LAP*^{t/g-box} lines treated with JA). These results demonstrate a role of the AAACGTG motif in JA-induced activation of the *LAP* promoter, although an additional element appears to be present in this proximal promoter region, this motif being responsible for the residual inducible activity detected in the –317*LAP*^{t/g-box} transformants.

T/G-Box dependent activation of the *LAP* promoter by the *JAMYC* transcription factors was further investigated by transient overexpression of these proteins in BY2 cells. Effector constructs expressing the *JAMYC2*/*JAMYC10* transcription factors under control of the 35S promoter were cotransfected with the –317*LAP* or the –317*LAP*^{t/g-box}–*GUS* reporter fusions (Fig. 2E). Expression of *JAMYC2*, *JAMYC10*, or both factors together was able to enhance expression of the –317*LAP* reporter fusion but did not result in any activation of the –317*LAP*^{t/g-box} construct. Basal levels of *GUS* activity obtained for this latter construct were lower than those obtained for –317*LAP*, consistent with a function of the T/G-box motif in JA-induced promoter expression (wound signaling is activated in bombarded cells). Overexpression of both transcription factors led to higher levels of activation than expression of each protein alone (Fig. 2E). This observation indicates that these factors can heterodimerize, the heterodimer exhibiting higher transactivation ability than the respective homodimers.

Overexpression of the JAMYC transcription factors enhances JA-induced expression of the pin2 and LAP genes

The ability of the *JAMYC* regulatory proteins to activate expression of the *LAP* and *pin2* genes was also examined in transgenic potato plants that constitutively expressed the *JAMYC2* or *JAMYC10* transcription factors under control of the 35S promoter (35S:*JAMYC2* and 35S:*JAMYC10* lines). Individual transformants were analyzed for high levels of expression of the *JAMYC* transcripts, and the two lines exhibiting a higher level of expression of the transgene were selected for further analysis. Plants were treated with increasing doses of JA, and expression of the *LAP* and *pin2* genes was analyzed after 6 h of treatment. Similar steady-state levels of *pin2* and *LAP* mRNAs were detected in the noninduced transformants and wild-type controls (Fig. 3). A marked increase in the

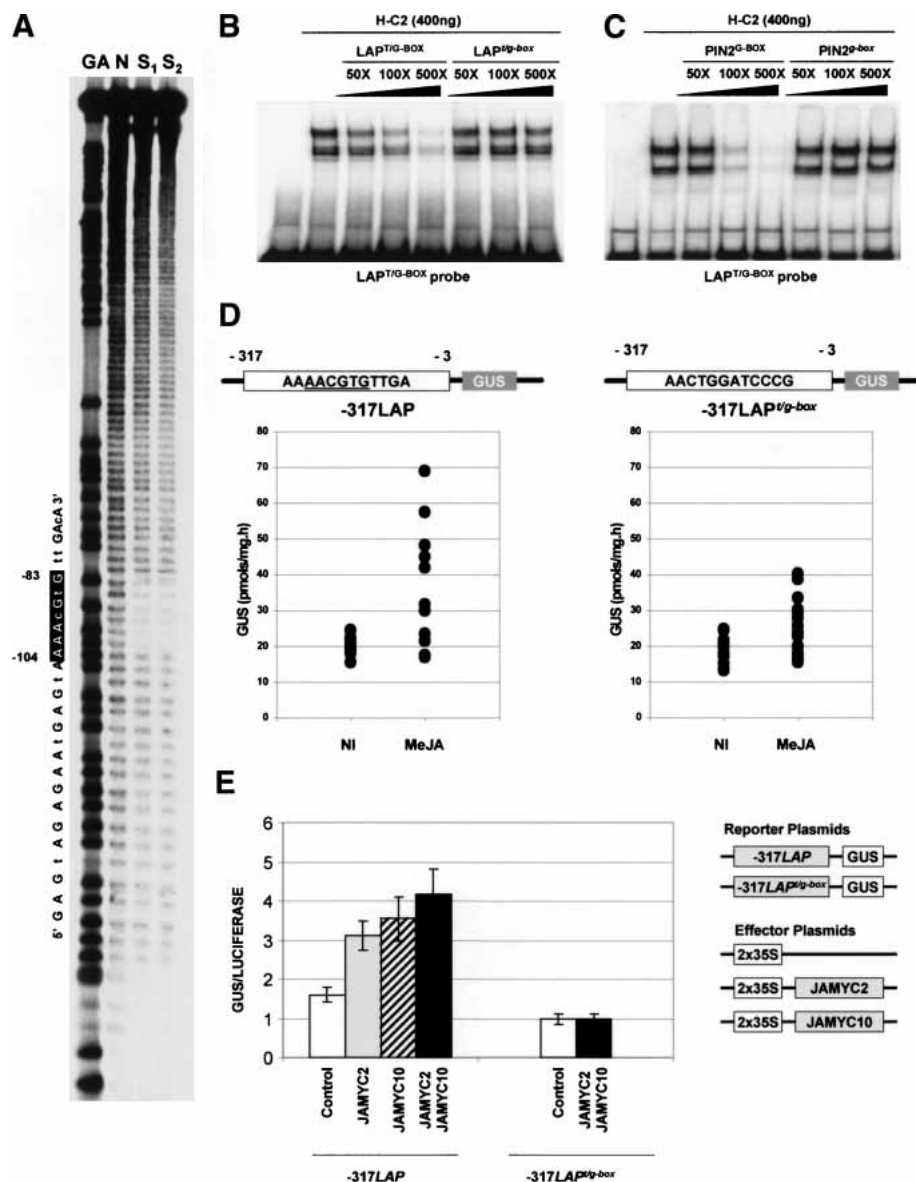


Figure 2. Characterization of the JAMYC DNA recognition motif. (A) Hydroxyl radical interference assays identify a T/G-box element in the $-125LAP$ promoter fragment as the recognition site for the His-C2 protein. The $-125LAP$ probe was asymmetrically labeled with ^{32}P -dCTP and treated with hydroxyl radical before incubation with the purified protein. Free probe (N) and retarded bands (S1 and S2) were purified by PAGE and separated on a sequencing gel. A G + A Maxam reaction (GA) on the same fragment was loaded as marker. Protected residues are shown as a black box on the nucleotide sequence on the left. Identical results were obtained for both His-C2 and His-C10 proteins. (B) Binding activity of His-C2 protein in gel retardation assays. End-labeled $LAP^{T/G-BOX}$ double-stranded oligonucleotides containing an intact T/G-box motif were used as probe. The oligonucleotide pairs $LAP^{T/G-BOX}$, with an intact motif, or $LAP^{t/g-box}$, in which the T/G-box motif was mutagenized, were used as competitors for binding. Oligonucleotides were added in 50- to 500-fold excess for competition. (C) JAMYC2 recognizes a G-box motif required for JA induction of the *pin2* promoter. Binding of the His-C2 protein to the $LAP^{T/G-BOX}$ probe was competed by incubation with a 50- to 500-fold excess of the unlabeled $PIN2^{G-BOX}$ (intact G-box) or $PIN2^{g-box}$ (mutated G-box) oligonucleotide pairs. (D) Site-directed mutagenesis of the LAP promoter T/G-box motif. Transgenic potato plants carrying either the $-317LAP$ or the $-317LAP^{t/g-box}$ constructs were grown on soil for 4 wk, and GUS activity was measured in noninduced leaves or in leaves treated for 24 h with 50 μ M MeJA. Dots represent the values of GUS activity measured in each individual transformant. (E) DNA binding and *trans*-activation activity of the JAMYC2 and JAMYC10 transcription factors in BY2 cells. The reporter constructs $-317LAP$:GUS and $-317LAP^{t/g-box}$:GUS, including intact or mutagenized copies of the T/G-box motif, were cotransfected with the effector plasmids 2x35S:JAMYC2 (JAMYC2) and 2x35S:JAMYC10 (JAMYC10), expressing the JAMYC proteins under control of the 2x35S promoter. The same vector without insert (2x35S) was used as a control. A 35S:Luciferase plasmid was also used as internal control to normalize cotransfection efficiency. Histograms represent the mean GUS/LUC values for each set of replicates.

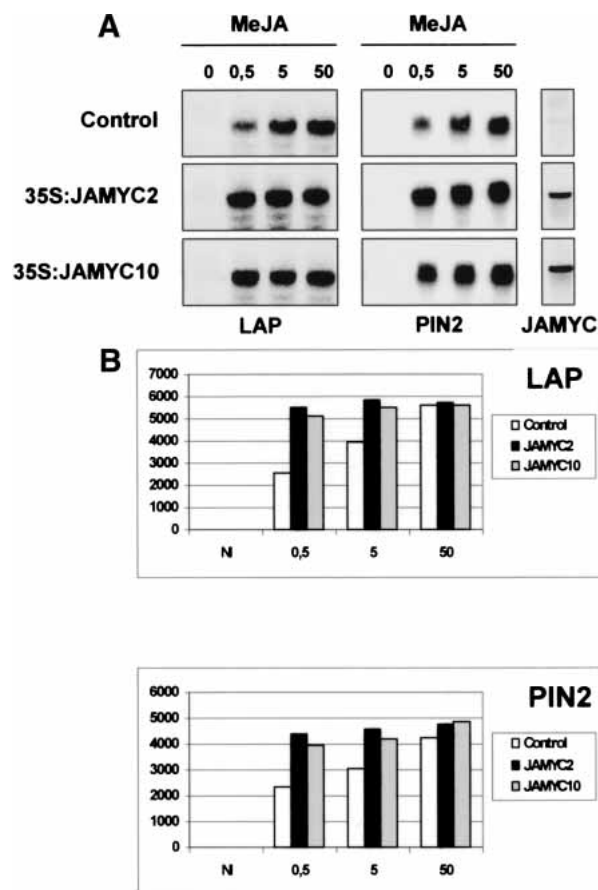


Figure 3. Potato JAMYC2 and JAMYC10 overexpressers show an enhanced induction of the *LAP* and *pin2* genes upon MeJA treatment. (A) *LAP* and *pin2* gene expression in wild-type controls (control) and the *35S:JAMYC2* and *35S:JAMYC10* transgenic potato plants after 6 h of treatment with increasing concentrations of MeJA. Each lane was loaded with 20 μ g of total RNA. Blots were hybridized with the *LAP*, *pin2*, and *JAMYC2* or *JAMYC10* probes as indicated. Equal loading was verified by EtBr staining of the gel. (B) Histograms represent the levels of expression of *LAP* and *pin2* as quantified by densitometric scanning of the blot.

levels of accumulation of these mRNAs was, however, observed in the JAMYC transformants after treatment with subsaturating doses of JA (cf. the levels of *pin2* and *LAP* transcripts detected in the *35S:JAMYC2* or *35S:JAMYC10* lines and in the controls, treated with 0.5 μ M or 5 μ M JA). At higher JA concentrations, differences between overexpresser lines and the controls were less evident, likely because of response saturation.

These results denote an enhanced JA-induced activation of the *pin2* and *LAP* transcripts in the overexpresser lines and demonstrate a role of both JAMYC2 and JAMYC10 tomato factors in regulated expression of these target genes. Activation of these genes is only observed after JA application, suggesting that JA-induced posttranscriptional modification of these transcription factors, or further induction of an additional regulatory activity, is required for gene expression.

In vivo footprinting of the *LAP* promoter region

To identify other motifs involved in JA-regulated expression of the -317 to -78 *LAP* promoter fragment, *in vivo* footprinting analysis was performed (Busk et al. 1997). Two G residues (positions -96 and -107) were identified on the top DNA strand that were partially protected from methylation in JA-treated leaves (Fig. 4A). Upstream of these two protected nucleotides, an additional G residue (position -118) exhibited enhanced reactivity to DMS treatment (see Fig. 4A). A correlation was observed between the inducible footprint and the time course of *LAP* mRNA accumulation. No further significant differences in the pattern of methylation were detected within the 300-bp promoter. Differences in methylation were also not observed on the lower DNA strand, owing to the lack of G residues within the protected region. A CGCGG sequence is identified by the hypersensitive G residue (Fig. 4C), whereas a GAGTA duplicated element, between positions -109 and -94 , is identified by the protected G residues. A duplicated GAGTA motif similar to the one identified here is present in the *cathepsin D inhibitor (CDI)* promoter, in a region that has been reported to be required for JA response of this gene (Ishikawa et al. 1994). These findings suggest a function of this duplicated motif in JA-regulated expression of these genes.

Protection of the AACGTG JAMYC target motif was not detected in these assays. Transcripts encoding these regulatory proteins are relatively abundant in noninduced plants. Therefore, it is possible that in noninduced conditions, this element is partially occupied by JAMYC in a transcriptionally inactive state.

Mutations of the protected G residues in the *LAP* promoter abolish the JA response

To determine whether these footprinted elements are required for JA induction, two progressive 5'-promoter deletions down to position -125 ($-125LAP$) and to position -78 ($-78LAP$) were obtained and introduced into transgenic potato plants. Both CGCGG sequence and GAGTA repeats are present in the $-125LAP$ deletion, whereas these elements were removed in the $-78LAP$ construct. Incubation with JA induced a substantial increase in the levels of GUS activity in the plants transformed with the $-125LAP$ construct (Fig. 4B). In contrast, a complete loss in JA responsiveness was observed in the $-78LAP$ transgenic lines. Contribution of the repeated GAGTA element to JA-induced expression was further analyzed by mutation of this duplicated motif in the context of the -317 to -3 *LAP* promoter-GUS fusion ($-317LAP^{GAGTA}$ construct; Fig. 4C). As shown in Figure 4B, mutation of the GAGTA element completely abolished JA response of the -317 promoter deletion. An intact GAGTA repeat is, therefore, required for JA response of the *LAP* promoter, indicating that this duplicated motif along with the JAMYC binding T/G-box element form a bifactorial JA-responsive (JARE) element that mediates JA-induced expression of the *LAP* genes.

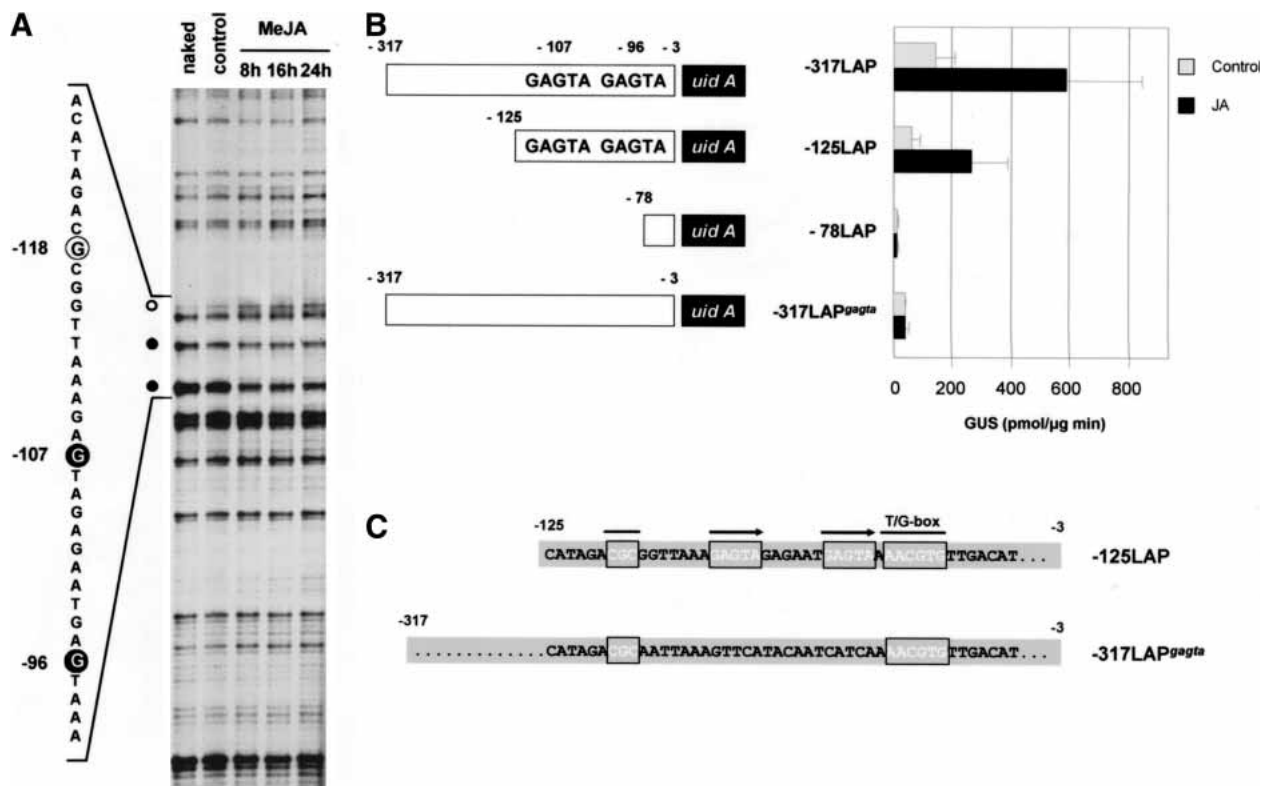


Figure 4. DMS in vivo footprinting of the proximal *LAP* promoter region identified two G residues in a GAGTA repeated motif that are protected from methylation and shown to be required for JA response. (A) DMS in vivo footprinting of the -270 to -3 *LAP* promoter region. Control tomato leaves (control) and leaves of plants treated with 50 μ M MeJA (8, 16, and 24 h) were directly incubated with DMS. DNA was extracted from these leaves, cleaved with piperidine, and subjected to LmPCR. Amplification of in vitro methylated DNA (naked) is also included as control. Protected G residues are indicated by filled circles and residues with enhanced reactivity (hypersensitivity) by open circles. The position of both protected and hyperreactive residues is indicated on the left. (B) Deletion analysis and site-directed mutagenesis of the proximal *LAP* promoter region. Transgenic potato plants carrying the -317*LAP*, -125*LAP*, -78*LAP*, and -317*LAP*^{gagta} constructs fused to the *uidA* gene were grown on soil for 4 wk, and GUS activity was measured in noninduced leaves or in leaves treated for 24 h with 50 μ M MeJA. The histogram shows the average values of GUS activity detected in nontreated and MeJA-treated plants. (C) Nucleotide sequence of the proximal *LAP* promoter. Hypermethylated G residues, GAGTA repeats, and the T/G-box motif are shown in the -125*LAP* promoter context. Mutations introduced to yield construct -317*LAP*^{gagta} are indicated.

The Arabidopsis AtMYC2 gene shares strong homology to JAMYC2 and JAMYC10 and is induced by JA treatment

To further assess the function of JAMYC2/JAMYC10 in regulated expression of the *pin2* and *LAP* genes, transgenic lines were generated with down-regulated levels of expression of these transcripts by antisense inhibition. However, only a small reduction in the levels of expression of the *JAMYC* genes was observed in some antisense lines. As an alternative to the inefficient down-regulation results obtained in these transformants, we searched for gene orthologs in *Arabidopsis*, to identify loss-of-function mutations in these genes. Comparison of the *JAMYC2* and *JAMYC10* coding regions to the complete *Arabidopsis* sequence identified three genes with significant similarity (60% identity at the amino acid level) to the tomato regulatory proteins. Phylogenetic analysis revealed that gene *RD22BP1* or *AtMYC2* (Abe et al. 1997) encodes the closest homolog to the tomato

JAMYC proteins (Fig. 5A). This gene had been reported to be induced by drought and ABA, and to play a role in transcriptional activation of the *rd22* promoter through cooperative interaction with the MYB-related protein AtMYB2 (Urao et al. 1993; Abe et al. 1997). RNA profiling analysis of AtMYC2/AtMYB2 overexpressers shows a constitutive activation of several drought and osmotic stress response genes, including the *rd22* target, but also increased levels of some JA- and pathogen-induced transcripts, like *VSP2* (Abe et al. 2003).

As a first approach to investigate whether AtMYC2 corresponds to a functional homolog of the JAMYC2/JAMYC10 factors, we investigated if expression of *AtMYC2* is induced in response to JA treatment. JA application resulted in rapid activation of the *AtMYC2* gene, with high levels of transcript observed as early as 30 min after JA treatment (Fig. 5B). *AtMYC2* thus corresponds to an early JA-responsive gene, which agrees with a role of this transcription factor in JA-regulated expression.

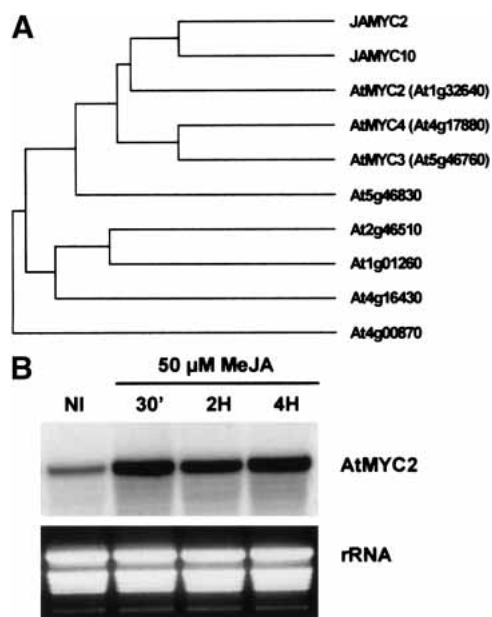


Figure 5. The *Arabidopsis* *AtMYC2* gene shares strong homology with *JAMYC2* and *JAMYC10* and is rapidly induced by JA. (A) Phylogenetic tree generated by CLUSTALW alignment of the tomato *JAMYC2* and *JAMYC10* proteins and related bHLH proteins from *Arabidopsis*. (B) RNA blot analysis of the induction of *AtMYC2* by MeJA. Total RNA was isolated from 3-week-old Col-0 plants grown in soil after different times of MeJA application, and 30 μ g was loaded per line. Equal loading was verified by rRNA visualization after EtBr staining.

Arabidopsis lines carrying an insertion within the *AtMYC2* gene are insensitive to JA

A search for *AtMYC2* insertion mutants in the T-DNA SALK collection (Alonso et al. 2003) retrieved mutations SALK_040500 (*atmyc2-1*) and SALK_083483 (*atmyc2-2*) with insertions at nucleotides 57 (Asn 18) and 1237 (Asp 411), respectively, within the protein coding region. RNA blot analysis of homozygous plants for these insertions revealed transcripts of either smaller or larger sizes, indicative of a loss of gene function (Fig. 6A). In *Arabidopsis*, JA induces strong inhibition of root growth (Staswick et al. 1992; Feys et al. 1994; Berger et al. 1996). Thus, we first investigated whether JA regulation was affected in these mutants by analyzing JA-induced root growth inhibition. Seeds of the *atmyc2-1* and *atmyc2-2* mutants or wild-type plants were germinated on JA plates, and roots were scored 2–3 wk later for growth. Normal growth and similar root lengths were observed for wild-type controls and *atmyc2* mutants in plates without JA (Fig. 6B). Increasing concentrations of JA resulted in severe root growth inhibition in wild-type plants, but did not affect root growth in the *atmyc2-1* and *atmyc2-2* mutants (Fig. 6B). Only a small reduction in root length was observed in these insertion lines at high (100 μ M) JA concentrations, these mutants therefore being insensitive to JA.

Ratios of germination and growth retardation of the *atmyc2-1* and *atmyc2-2* mutants germinated on ABA

plates were comparable to those of the controls (data not shown). Therefore, these lines apparently do not exhibit a reduced sensitivity to ABA like that reported for a Ds-insertion mutant in the *AtMYC2* gene between residues 107 and 108 of the protein (Abe et al. 2003).

The *AtMYC2* insertion lines exhibit altered JA-regulated gene expression

To further study the effects of these mutations on JA-regulated gene expression, we analyzed steady-state levels of the JA-regulated transcripts *VSP* and *JR1* (Berger et al. 1995; Titarenko et al. 1997) and the JA/ethylene (ET)-regulated transcripts *PDF1.2* and *b-CHI* (Penninckx et al. 1998; Lorenzo et al. 2003) in the *atmyc2-1* and *atmyc2-2* insertion lines. RNA blot analysis showed low levels of mRNA for these transcripts in untreated wild-type controls or the *atmyc2* mutants (Fig. 6C). However, whereas JA application induced normal levels of accumulation of these genes in wild-type seedlings, induction of the JA/wound-responsive transcripts *VSP* and *JR1* was strongly reduced in the *atmyc2* mutants (reduction was more evident for the *VSP* transcript). JA treatment, in contrast, induced much greater levels of expression of the pathogen defense-related transcripts *PDF1.2* and *b-CHI* in the *atmyc2* mutants than in wild-type seedlings, suggesting a role of *AtMYC2* in repression of these JA/ET-responsive genes. Together, these findings indicate a function of *AtMYC2* in JA-induced gene expression, this transcription factor having a role in activation of the JA/wound-responsive genes *VSP* and *JR1*, but also in repression of the JA/ET-responsive genes *PDF1.2* or *b-CHI*.

Expression of tomato *JAMYC2* and *JAMYC10* complements the JA-related phenotype of the *AtMYC2* mutants

Evidence for a functional homolog activity of the tomato *JAMYC* and the *Arabidopsis* *AtMYC2* transcription factors was further obtained by complementation of the *atmyc2-2* mutation with the tomato proteins. Mutant plants expressing the *JAMYC* overexpression constructs were generated and examined for JA-induced inhibition of root growth. Overexpression of the tomato *JAMYC* transcription factors was able to recover the root growth phenotype of the *atmyc2-2* mutation as a severe root growth inhibition was observed on JA plates in transgenic *JAMYC/atmyc2-2* seedlings compared with the *atmyc2-2* mutant controls (Fig. 6D). Such root growth differences were not observed in plates without JA (data not shown), thus evidencing JA mediation of this growth effect. JA-induced expression of the *VSP* and *PDF1.2* genes was also restored in these transgenic lines (Fig. 6E). JA induction of the *VSP* transcript was increased to levels comparable to those of wild-type plants in the *JAMYC* overexpressers. JA-induced expression of the *PDF1.2* gene was also sensibly reduced in these lines, to levels similar to those observed in transgenic *atmyc2-2* mutants complemented with the endogenous *Arabidopsis* gene (see Fig. 6E). These results demonstrate that

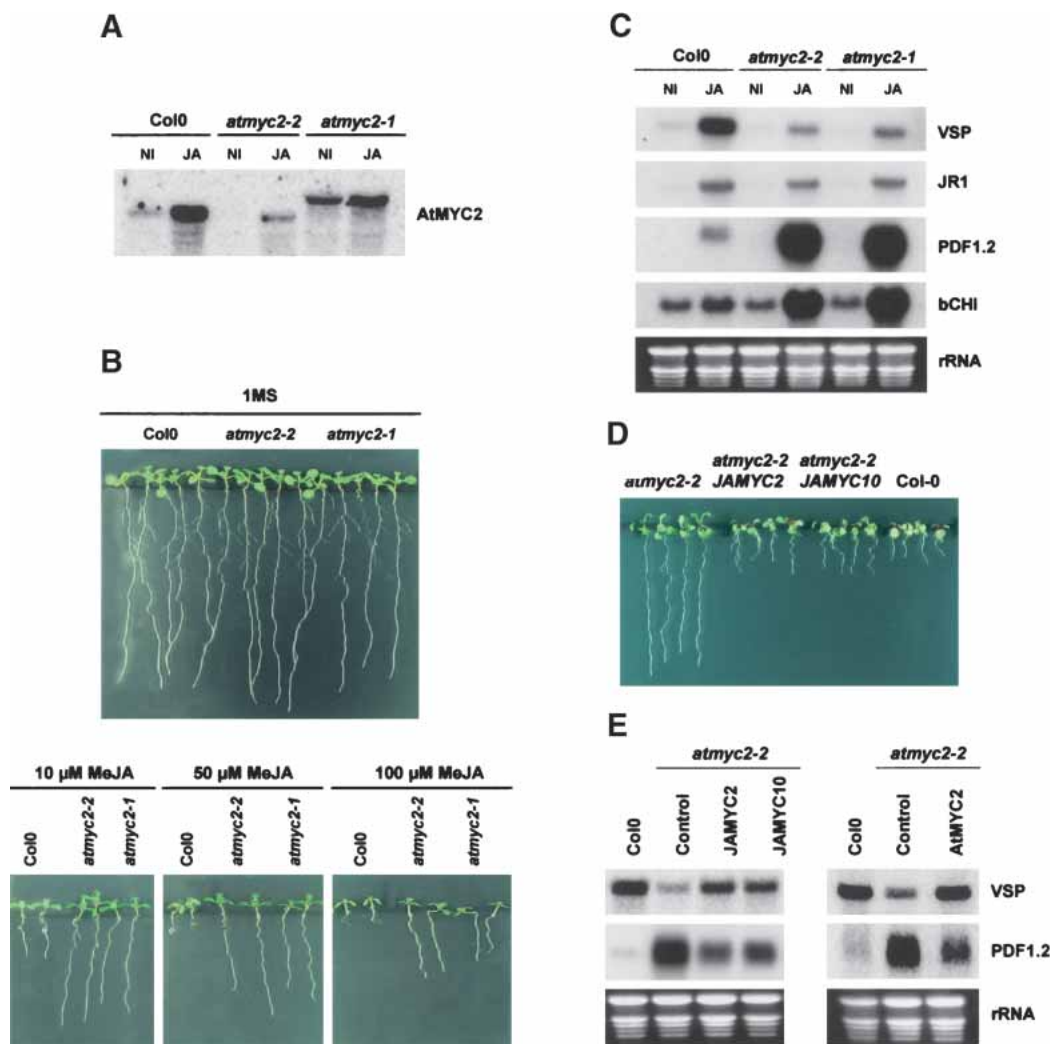


Figure 6. *Arabidopsis* lines carrying an insertion within the *AtMYC2* gene are insensitive to JA and exhibit altered JA-regulated gene expression. (A) Northern analysis of the *atmyc2-1* (SALK_040500) and *atmyc2-2* (SALK_083483) mutants. Fifteen micrograms of total RNA from Col-0 and *atmyc2* mutant seedlings noninduced or treated for 8 h with 50 μ M MeJA was loaded per lane and hybridized with the *AtMYC2* probe. (B) MeJA inhibition of root growth in the *atmyc2* mutants and Col-0 seedlings grown for 10 d on MS plates, or MS plates with 10, 50, and 100 μ M MeJA. (C) MeJA induction of the *VSP*, *JR1*, *PDF1.2*, and *b-CHI* transcripts in Col-0 and the *atmyc2* mutants. Fifteen micrograms of total RNA from noninduced seedlings (NI) or seedlings treated for 8 h with 50 μ M MeJA (JA) was loaded per lane and hybridized with the indicated probes. (D) Complementation of the *atmyc2-2* JA-insensitive root growth phenotype by overexpression of the JAMYC2 and JAMYC10 proteins (JAMYC/*atmyc2-2* lines). Seeds were grown for 10 d on MS medium containing 50 μ M MeJA. (E) Molecular complementation of the *atmyc2-2* phenotype by expression of the *Arabidopsis AtMYC2* gene or the tomato JAMYC factors. Fifteen micrograms of total RNA from seedlings treated for 8 h with 50 μ M MeJA was loaded per lane and hybridized with the *VSP* and *PDF1.2* probes.

overexpression of the tomato JAMYC transcription factors complements the JA-insensitive phenotype of the *atmyc2-2* mutation, these transcription factors therefore corresponding to the functional homologs of AtMYC2.

Discussion

Isolation of cDNA clones encoding the JAMYC factors

Using yeast one-hybrid screening, we have identified two transcription factors binding the *cis*-elements in the

proximal promoter region of the tomato *LAP* genes. A -317 to -78 promoter region shown to be sufficient for JA-regulated gene expression was used as bait for screening, to yield clones JAMYC2 and JAMYC10 with >80% overall identity, encoding proteins with a basic-helix-loop-helix leucine zipper (bHLH-ZIP) DNA-recognition domain. JAMYC2 and JAMYC10 transcripts are rapidly induced in leaves by wounding and JA treatment, with a kinetics that precedes that of the *LAP* gene, supporting a function of these transcription factors in JA-induced *LAP* gene expression.

Hydroxyl radical in vitro footprinting studies iden-

tified a 7-bp AAACGTG motif as the binding site for the JAMYC factors. Notably, members of the bHLH-ZIP (MYC-like) family of transcription factors were found to bind preferentially to canonical G-box (CACGTG) elements, although they can also recognize T/G-box sequences like the AACGTG motif identified here (Blackwell et al. 1993). EMSA studies showed that a G-box element required for JA-induced expression of the *pin2* gene (Kim et al. 1992) is able to compete efficiently for JAMYC binding to the AAACGTG motif. These proteins can bind a *pin2* promoter fragment including this G-box motif, indicating that *pin2* is a likely target for JAMYC-regulated transcriptional activation. *JAMYC2*, on the other hand, shares near-complete nucleotide sequence identity with *LEJA3*, a partial tomato cDNA clone identified in one-hybrid screening studies with the promoter region of the tomato *threonine deaminase* (*Tda*) gene (L. Broday and E. Lifschitz, pers. comm.). Notably, *pin2*, *Tda*, and *LAP* are coordinately induced in response to wounding and JA application (Hildmann et al. 1992; Samach et al. 1995; Ryan 2000). Consistent with this coordinated pattern of expression, *in vitro* studies showed that the JAMYC factors bind to these promoters and suggest a role for these proteins in JA-regulated transcriptional activation.

JAMYC-regulated gene expression

Mutation of the AAACGTG element in the -317 to -3 *LAP* promoter resulted in reduced JA activation of the -317*LAP*^{*tg-box*} construct. A residual level of activity was, however, observed in the transformants containing this construct, suggesting the presence of additional JA-responsive motifs in this promoter region. In line with this observation, overexpression of the *JAMYC2* and *JAMYC10* proteins did not lead to constitutive activation of the target *pin2* or *LAP* genes, although enhanced expression of the *pin2* and *LAP* transcripts was observed in the *JAMYC*-OE lines after treatment with JA. Therefore, it is possible that JA is required for activation of the *JAMYC* factors through phosphorylation or other types of posttranslational modification of these proteins or, alternatively, that JA induces expression of other rate-limiting transcription factors required for defense gene activation. Consistent with this latter hypothesis, animal and plant MYC-like proteins were often found to require interaction with partner proteins to activate target gene expression (Roth et al. 1991; Kretzner et al. 1992; Payne et al. 2000). Supporting evidence for interaction of additional regulatory proteins has, indeed, been obtained from *in vivo* footprinting analyses, where a GAGTA repeat situated directly adjacent to the T/G-box motif was found to be protected from methylation in induced leaves. Mutation of this duplicated sequence (-317*LAP*^{*gagta*} construct) abolishes JA response of the -317 promoter deletion, indicating a requirement of this repeated element for JA-induced expression. No apparent sequence homology was observed between this duplicated motif and other JA-responsive *cis*-elements, with the exception of the potato *CDI* promoter (Ishikawa

et al. 1994), which also has a duplicated GAGTA motif in a region required for JA-regulated expression of the gene.

Conservation of MYC-based JA signaling in Arabidopsis

Amino acid sequence comparisons showed elevated overall identity of the tomato *JAMYC* proteins with the *Arabidopsis AtMYC2* gene, with a reported function in ABA-regulated responses to drought stress (Abe et al. 2003). In RNA blot analysis, we observed that JA induces a rapid accumulation of this transcript to even greater levels than those obtained in response to ABA application (data not shown), thus suggesting a role of this gene as functional homolog of the tomato *JAMYC* factors. Consistent with this observation, knockout mutations in this gene (*atmyc2-1* and *atmyc2-2* mutants) rendered plants insensitive to root growth inhibition by JA, indicating an impaired JA response in these plants. These insertion lines, on the other hand, were male fertile, which indicates that *AtMYC2* is not required for normal stamen and pollen development.

JA-induced expression of *VSP* and *JR1* is blocked in the *atmyc2* lines, these mutants exhibiting in response to JA treatment much greater levels of activation of the *PDF1.2* and *b-CHI* genes. These results are indicative of a function of *AtMYC2* in transcriptional activation of JA-regulated genes but also in repression of JA/ET-regulated gene expression.

Expression of the tomato transcription factors rescued the root elongation phenotype of the *Arabidopsis atmyc2* mutants and recovered JA-induced expression of both JA- and JA/ET-responsive genes, thus confirming that *AtMYC2* and the *JAMYC* genes are functional homologs. Overexpression of the *JAMYC* or *AtMYC2* factors did not lead to constitutive activation of the *VSP* gene, indicating that, as observed in tomato, activation of an as-yet-unidentified regulatory protein is also required for transcriptional activation of the JA-responsive genes.

Cross-talk regulation of the JA- and JA/ET-signaling pathways

In *Arabidopsis*, JA regulates defense responses against both herbivore (McConn et al. 1997) and necrotrophic pathogen attack (Pieterse et al. 1998; Thomma et al. 1999). Activation of these two responses involves interaction of both JA- and ethylene (ET)-signaling pathways. A synergistic interaction of these pathways is involved in activation of pathogen-related defense genes like *PR5*, *PDF1.2*, and *b-CHI* (Xu et al. 1994; Penninckx et al. 1998; Ellis and Turner 2001), whereas a negative interaction would repress expression in the directly damaged tissues of JA/wound-related genes like *VSP* or *Thi2.1* (Berger et al. 1995; Vignutelli et al. 1998; Rojo et al. 1999).

JA/ET-dependent induction of pathogen-related genes is regulated by the *ethylene response factor* (*ERF1*) regu-

latory protein (Lorenzo et al. 2003). Constitutive expression of ERF1 rescues the defense response defects of the *coi1* and *ein2* mutants (Berrocal-Lobo et al. 2002) and leads to constitutive activation of JA/ET-regulated genes (Lorenzo et al. 2003). Interestingly, a down-regulated expression of the *VSP* and *Thi2.1* genes was observed in RNA profiling analysis of ERF1 plants (Lorenzo et al. 2003), indicative of a negative regulatory function of ERF1 on JA-responsive gene expression.

In this work, we have shown that knockout mutations in the *AtMYC2* gene have much reduced levels of expression of *VSP* and *JR1* in response to JA treatment, but accumulate higher levels of the JA/ET-regulated transcripts *PDF1.2* and *b-CHI*. A model of our current view of the mechanism of action of these regulatory proteins is shown in Figure 7. Herbivore or pathogen attack induces both systemic accumulation of JA and local synthesis of ET (Penninckx et al. 1998; Rojo et al. 1999). JA activates expression of *VSP*, *JR1*, and *Thi2.1*, which accumulate systemically in response to mechanical wounding or herbivore attack. Expression of these genes is repressed at the wound site by the local synthesis of ET. A positive interaction between JA and ET, in turn, is responsible for induction of *PDF1.2* and *b-CHI* in response to necrotrophic pathogen attack. Activation of these genes is mediated by ERF1 (Lorenzo et al. 2003) and

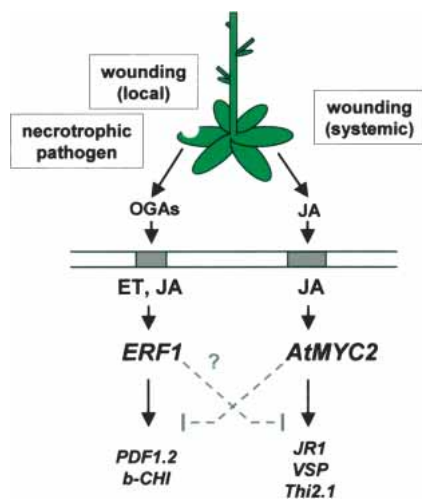


Figure 7. Model for the proposed function of AtMYC2 and ERF1 in cross-talk regulation of the JA and JA/ET pathways in *Arabidopsis*. Herbivore and pathogenic attack promote a transient increase of both JA and ethylene. JA is involved in systemic activation of wound-responsive genes, and ethylene prevents local expression of these genes. A positive interaction between JA and ethylene is involved in activation of the JA/ET-pathway regulated genes in response to necrotrophic pathogen attack. Whereas ERF1 activates JA/ET-regulated gene expression (Lorenzo et al. 2003), AtMYC2 is involved in activation of JA-regulated genes and also in repression of JA/ET-induced gene expression. Arrows and bars indicate positive and negative interactions, respectively. A possible function of ERF1 in repression of the JA-regulated pathway is indicated by a question mark. (OGAs) Oligogalacturonides; (JA) jasmonic acid; (ET) ethylene.

AtMYC2 (this work). ERF1 activates JA/ET-regulated gene expression and possibly plays a role in repression of genes differentially regulated by JA (Lorenzo et al. 2003). AtMYC2, in turn, is involved in JA-regulated gene expression and repression of the JA/ET pathway. Interaction between these two signaling pathways therefore occurs downstream in the signaling process, likely at the level of gene-specific promoters.

ERF transcription factors bind a GCCGCC (GCC-box) element present in the promoter region of several *PR* genes (Fujimoto et al. 2000; Ohme-Takagi et al. 2000; Gu et al. 2002). Deletion analysis of the *PDF1.2* promoter has recently identified a GCC-box element associated with JA/ET-responsive expression of this gene (Brown et al. 2003). This GCC-box element is directly followed by a G-box motif (-255 to -250), which might be a functional binding site for AtMYC2. Noteworthy, close proximity of G-box and GCC-box elements is a common feature of many tobacco *PR* genes (Sessa et al. 1995; Buttner and Singh 1997), binding of AtMYC2 to the G-box motif being likely to compete for binding of ERF1 to the GCC-box element. Hence, it is possible that competitive binding of these TFs mediates negative cross-talk regulation of the JA and JA/ET pathways.

A JA-responsive element in the *AtVSP1* promoter has also been recently identified (Guerineau et al. 2003). This regulatory element is comprised of an inverted repeat containing a G-box-like element, located 150 bp upstream of the TATA-box. Mutation of the G-box-like element or the left part of the inverted repeat both have a strong effect on JA-regulated gene expression (Guerineau et al. 2003). GCC-box elements indicative of a direct binding of ERF1, to repress expression of this gene, are not observed in the *AtVSP1* promoter, suggesting that negative cross-talk regulation by ERF1 might involve a different regulatory mechanism.

JA-pathway regulation in tomato

Important differences were observed in JA signaling between *Arabidopsis* and tomato. *Arabidopsis* mutants defective in JA synthesis or perception are male sterile (Feys et al. 1994), whereas tomato mutants were reported to be male fertile (Howe et al. 1996). Systemic induction of JA responses in tomato requires the 18-amino-acid peptide systemin (Ryan 2000), but evidence for a similar pathway in *Arabidopsis* has not been observed. Also, ethylene signaling is required for wound induction of *pin2* in tomato (O'Donnell et al. 1996), whereas in *Arabidopsis* it seems to suppress JA-dependent gene expression in the wounded tissues (Rojo et al. 1999). Recent advances, however, suggest that these differences are mainly due to gaps in knowledge, rather than actual discrepancies in the signaling pathways in these two species. Grafting experiments using mutants deficient in either JA biosynthesis or JA perception, for example, have questioned the role of systemin as a systemic wound signal and pointed to JA as the long-distance moving signal (Li et al. 2002). Pollen examination of the tomato *jasmonic acid insensitive-1* (*jai1*) mutant

showed that it has reduced pollen viability and germination and that JA is also required for viable pollen formation in tomato (Li et al. 2004). Defects in this mutant are caused by a lesion in the *LeCOI1* gene, encoding the tomato homolog of COI1 (Li et al. 2004). Like the *Arabidopsis coi1* mutant, *jai1* plants are insensitive to coronatine and highly resistant to coronatine-producing strains of *Pseudomonas syringae* (Zhao et al. 2003). Microarray analysis has shown that expression of all JA up-regulated genes is blocked in the *jai-1* plants, thus demonstrating a similar function of COI1 in *Arabidopsis* and tomato.

The tomato ERF1-related factors Pti4, Pti5, and Pti6 were identified in a yeast two-hybrid screen with the *Pto* resistance gene (Zhou et al. 1997; Kim et al. 2002). A role of these tomato regulatory factors in plant defense activation was proved by expression in *Arabidopsis*. Expression of *Pti4* causes the activation of several *Arabidopsis* pathogenesis-related genes, including *PDF1.2*, *b-CHI*, and *PR4*, and results in resistance to fungal pathogens and increased tolerance to the bacterial pathogen *P. syringae* (Gu et al. 2002; Wu et al. 2002), indicating that Pti4 might correspond to a tomato homolog of ERF1.

In this report, we have presented evidence for a functional homolog function of the tomato JAMYC and *Arabidopsis* AtMYC2 proteins. These factors have a main function in JA-regulated gene expression and cross-talk repression of the JA/ET-signaling pathway. Conservation of all these signaling components (COI1, Pti4/ERF1, and JAMYC/AtMYC2) thus suggests a similar regulatory network in these plants, although target genes are different in these two species (i.e., genes that encode PIs are not present in *Arabidopsis*). Noteworthy, wound/JA-regulated gene expression has been largely studied in tomato, whereas identification of the regulatory mechanisms underlying JA/ET-regulated gene activation derive mainly from studies in *Arabidopsis*. Therefore, further study of the complementary pathways will be required to confirm identical pathways for defense gene regulation in these species.

In conclusion, we have identified a conserved component of the JA-signaling pathway with a key function in JA-regulated expression and cross-talk inhibition of JA/ET signaling. Besides a function in JA-regulated transcriptional activation, JAMYC/AtMYC2 serve as cross-talk or integration points of both the JA- and JA/ET-signal outputs, concerted action between these MYC proteins and the ERF1/Pti4 playing a key role in orchestrating activation of different sets of defense-related genes in response to both herbivore or necrotrophic pathogen challenges.

Materials and methods

Plant materials and treatments

Tomato (*Lycopersicon esculentum* cv *Moneymaker*) and potato (*Solanum tuberosum* var *Désirée*) plants were grown in soil in the greenhouse under a 16 h light/8 h dark regime. For in vitro conditions, plants were grown in MS media supplemented with 2% sucrose. For Northern blot experiments, plants were

grown in soil for 4–5 wk and sprayed with a 50 μ M MeJA solution or wounded by midrib cutting of the leaves. Leaves (second to fourth from the apex) were collected at different times after treatment. JAMYC2 and JAMYC10 overexpressers were obtained by transformation of *Solanum tuberosum* var. *Désirée* with the 35S:JAMYC2 or 35S:JAMYC10 constructs in pBin19.

Arabidopsis thaliana (ecotype Columbia-0) and mutants SALK_040500 (*atmyc2-1*) and SALK_083483 (*atmyc2-2*) were grown in soil in the greenhouse under a 12 h light/12 h dark regime. For in vitro conditions, seeds were surface-sterilized and sown in MS supplemented with 1% sucrose. For Northern blot experiments, seedlings were grown on plates for 2 wk and incubated in a 50 μ M MeJA solution for 8 h during the light period. For JA-induced root growth inhibition, seeds were sown on MS plates containing 0, 10, 50, or 100 μ M MeJA, and cultured in a vertical position.

atmyc2-2/JAMYC2 and *atmyc2-2/JAMYC10* plants were obtained by transformation of the *atmyc2-2* homozygous mutants with the 35S:JAMYC2 or 35S:JAMYC10 constructs.

Yeast one-hybrid screening

The –308/–78 wild-type EcoRI/HindIII fragment of the *LAP17.1* promoter was inserted into the EcoRI/filled XbaI site of the pHSi vector (–308LAP:HIS3) or the EcoRI/SmaI site of the pLacZi vector (–308LAP:LacZ). Plasmid –308LAP:LacZ was linearized with NcoI and transformed into the yeast strain YM4271 (Clontech). Recombinants were selected on SD-Ura medium and retransformed with the –308LAP:HIS3 plasmid linearized with XhoI to obtain the double reporter yeast strain –308LAP:HIS3/–308LAP:LacZ. Recombinants were selected on SD-His-Ura and single recombination events were verified by Southern blot. Poly(A)⁺ RNA was prepared from tomato leaves treated with 50 μ M MeJA and used for cDNA synthesis using the Stratagene cDNA synthesis kit. This cDNA was directionally cloned into the EcoRI/XhoI sites of the HybriZap vector and packaged to obtain a primary library into this vector. This library was amplified and in vivo excised to a pAD-GAL4 plasmid vector as described by the manufacturer (Stratagene). The cDNA library was screened with the double reporter yeast strain in the presence of 30 mM 3-AT, as described by the manufacturers (Matchmaker One-Hybrid system; Clontech). A λ -Zap cDNA library obtained from leaves was screened with the partial cDNA clones *C2* and *C10* identified in the one-hybrid screening, to isolate the full-length clones *JAMYC2* and *JAMYC10*.

Hydroxyl Radical Interference

Hydroxyl radical interference experiments were performed as described by Hayes and Tullius (1989). A –125LAP fragment, asymmetrically labelled with ³²P-dCTP by fill-in with the Klenow polymerase fragment, was used as probe for JAMYC binding and subsequent hydroxyl reaction.

Electrophoretic mobility shift assays (EMSA)

The double-stranded oligonucleotides *LAP^{T1/G-BOX}* (5'-GGG AATGACATAGACGCGGTTAAAGAGTAGAGAATGAGTA AAACGTGTTGACATGC-3'; 5'-TTGCATGTCAACACGTTT TACTCATTCTCTACTCTTTAACCGCGTCTATGTCAC-3'), *LAP^{T1/gbox}* (5'-GGGAATGACATAGACGCGGTTAAAGAGTA GAGAATGAGTAAATCGGGTTGACATGC-3'; 5'-TTGCAT GTCAACCCGATTTACTCATTCTCTACTCTTTAACCGCGTCTATGTCAC-3'), *PIN2^{G-BOX}* (5'-GGGAGTTCCAAC TAATTATCACGTGGACTTATAAGAAACCGA-3'; 5'-GGT CGGTTTCTTATAAGTCCACGTGATAATTAAGTTGGAA CTC-3'), and *PIN2^{g-box}* (5'-GGGAGTTCCAACCTAATTATC

TCGGGACTTATAAGAAACCGA-3'; 5'-GGTCGGTTCT TATAAGTCCCCGAGATAATTAAGTTGGAACCTC-3') were used as probes and competitors. The double-stranded oligonucleotides were end-labeled with α -³²P-dATP/dCTP by fill-in with the Klenow polymerase fragment, and purified on NAP5 columns (Pharmacia) according to the manufacturer's instructions.

The EcoRI/SalI fragment corresponding to *C2* was cloned into the pET28a vector (Novagen) and transformed into *Escherichia coli* BL21 cells. Production and purification of the His-Tag fusion protein (His-C2) was performed according to the manufacturer's instructions. The radioactive probe was incubated with 300 ng of the purified protein in 20 μ L of 1 \times binding buffer (20 mM HEPES at pH 7.8, 40 mM KCl, 0.5 mM EDTA, 5 mM MgCl₂, 1 mM DTT, 1 μ g/ μ L BSA, 0.01% Triton X-100, 10% glycerol) and 100 ng poly(dI-dC), in the presence or absence of nonradiolabeled competitor for 30 min on ice, before loading on a 1 \times TBE, 5% polyacrylamide gel.

DNA constructs and plant transformation

The *JAMYC2* and *JAMYC10* coding regions were amplified by PCR using synthetic primers (*J2D*: 5'-GTGTTTATGGAATG AC-3'; *J2R*: 5'-GACGATTCTATCTAC-3' for *JAMYC2* and *J10D*: 5'-GATTGAATGACGGAC-3'; *J10R*: 5'-GATAATTTCATCGCG-3' for *JAMYC10*) and cloned into the *Sma* site of pUC18. Constructs 35S:*JAMYC2* and 35S:*JAMYC10* were obtained by excising the XbaI/SacI *JAMYC2* and *JAMYC10* fragments from pUC18 and cloning into the pBin19-35S:nosT vector. Constructs -317LAP^{g-box}:GUS and -317LAP^{gagta}:GUS were obtained by inverse PCR reactions on the -317LAP-pGUS plasmid (Ruiz-Rivero and Prat 1998), using synthetic primers with a 10-bp sequence including a BamHI site in their 5'-ends that replaces the original T/G-box site (*LS8A*: 5'-CTGGATCC CGCATGCAAATATCTTGTTTC-3'; *LS8B*: 5'-CGGGATCCAG TTACTCATTCTCTACTC-3') for the first construct, or using the oligonucleotides *FPLA*: 5'-CAATCATCAAAACGTGTTG ACATGC-3' and *FPLB*: 5'-TATGAACCTTAAATTGCGTCTATGTCATTCA-3' for the second one. Constructs -317LAP:GUS, -317LAP^{g-box}:GUS, and -317LAP^{gagta}:GUS were obtained by EcoRI/HindIII digestion of the corresponding pGUS plasmid constructs and cloning into the pBin19 vector.

Transformation of *S. tuberosum* var *Désirée* plants was performed as previously described (Carrera et al. 2000). Transformation of *A. thaliana* Col-0 or the *atmyc2-2* mutant was performed as described by Bechtold and Pelletier (1998).

Transient expression assays

Tobacco BY2 cells freshly subcultured for 5 d were spread on filter paper 1 d before bombardment and incubated on NT medium (5 g/L Murashige and Skoog salts, 1 mg/L thiamine, 200 mg/L KH₂PO₄, 0.2 mg/L 2,4-D, 100 mg/L myo-inositol, 30 g/L sucrose, 2 g/L gelrite) overnight at 26°C in the dark. Four hours before bombardment, cells were moved to NT medium with 200 mM mannitol. DNA absorption to gold particles and bombardment using a helium-driven particle accelerator (PDS-1000; Bio-Rad) was performed according to the manufacturer's recommendations. Cells were transformed with 1 μ g of the pUC18 -317LAP:GUS or -317LAP^{g-box}:GUS reporter plasmids, 1 μ g of the pUC18 35S:LUC plasmid as internal standard, and up to 2 μ g of either the pUC18 35S:*JAMYC2* or 35S:*JAMYC10* constructs or the 35S:nosT vector without insert (control). After bombardment, cells were moved to fresh NT medium and incubated for 22 h at 26°C in the dark, before freezing in liquid nitrogen.

For activity assays, samples were thawed and homogenized

on ice in a buffer containing 25 mM Tris (pH 7.8), 2 mM CDTA, 2 mM DTT, 10% glycerol, and 1% Triton X-100, and cleared by centrifugation at 12,000g for 5 min. For luciferase (LUC) and GUS assays, 60–100 μ g of protein extract was used. LUC activity was determined using the Luciferase Assay System kit (Promega), according to the manufacturer's instructions. GUS activity was measured by fluorometric assay as described by Jefferson (1987).

Analysis of gene expression

Total RNA was prepared as described by Logemann et al. (1987). Hybridizations were performed at 42°C with washes at 65°C as described by Amasino (1986). 3'-Noncoding ends corresponding to *C2* and *C10* were used as probes for hybridizations in tomato. For analysis of the potato transgenic lines, total cDNAs corresponding to *JAMYC2*, *JAMYC10*, *LAP*, and *pin2* were used as probes. Total cDNAs corresponding to *AtMYC2*, *VSP*, *JR1*, *PDF1.2*, and *b-CHI* were used as probes for Northern blot analysis in *Arabidopsis*. GUS activity of leaf protein extracts from the -317LAP:GUS, -317LAP^{g-box}:GUS, and -317LAP^{gagta}:GUS transgenic plants was measured by fluorometric assay as previously described (Jefferson 1987).

LmPCR in vivo footprinting

In vivo footprinting was performed from leaves of *Lycopersicon esculentum* cv *MoneyMaker* untreated or treated with MeJA, and directly incubated with dimethyl sulphate (DMS). DMS treatment and DNA purification and cleavage at modified guanines were performed as described previously (Busk et al. 1997).

Ligation-mediated polymerase chain reaction (LmPCR) conditions were as previously described (Busk et al. 1997). Oligonucleotides used for LmPCR amplification of the sense strand were: *Lap12*: 5'-AGAATTGAAGTGTCTAAGCGAAATTT CG-3'; *Lap13*: 5'-GCCAAATTTTCGTGCCAACTTTAGGTTGG-3'; and *Lap14*: 5'-CGTGCCAACCTTTAGGTGGTTACCGAT AGTTAGACC-3'; and those for the antisense strand: *Lap16*: 5'-CTGAAGGGTTAGAATGCAAAGATG-3'; *Lap17*: 5'-GAT GAAGAAGAAGCAAACAATGAAGAAACTC-3'; and *Lap18*: 5'-GAAGAAGAAGCAAACAATGAAGAAACTCTTAGTGT TGCCATTG-3'.

Acknowledgments

We thank Javier Paz-Ares, José Sánchez-Serrano, and David Hannapel for critical reading of the manuscript. We also thank Pilar Fontanet and Carme Badia for excellent plant care. This research was supported by grants BIO99-0961 and BIO2002-02045 from the Spanish MCyT and by predoctoral grants from the Catalan CERBA and Spanish AECl, respectively, to M.B. and A.A.

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