

## Characterization and Induction of Potato HMGR genes in Relation to Antimicrobial Isoprenoid Synthesis

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### ABSTRACT

Induction of HMG-CoA reductase(HMGR) is essential for the biosynthesis of sesquiterpenoid phytoalexins and steroid derivatives in Solanaceous plants following wounding and pathogen infection. To better understand this complex step in stress-responsive isoprenoid synthesis, three classes of cDNAs for HMGR (hmg1, hmg2, & hmg3) were isolated from a potato tuber library. The potato cDNAs had extensive homology in open reading frames but had low homology in the 3'-untranslated regions. RNA gel blot analysis using gene-specific probes revealed that hmg1 is induced by wounding but wound induction is strongly suppressed by arachidonic acid or by inoculation with *Phytophthora infestans*. In contrast, hmg2 and hmg3 are slightly induced by wounding and strongly enhanced by arachidonic acid or inoculation. The induction and suppression of HMGR genes parallel the suppression of steroid and stimulation of sesquiterpenoid accumulations observed in earlier investigations. Treatment of the tuber disks with a low concentration of methyl-jasmonate doubled the wound induced accumulation of hmg1 transcripts and steroid-glycoalkaloid accumulation, but did not affect the abundance of transcripts for hmg2 or hmg3 nor induce phytoalexins. High concentration of methyl-jasmonate suppressed hmg1 mRNA and steroid-glycoalkaloid accumulation, induced hmg3 mRNA, and did not elicit phytoalexins. Lipoxygenase inhibitors suppressed the accumulation of hmg1 transcripts and steroid-glycoalkaloids, which were restored by exogenous methyl-jasmonate. Methyl-jasmonate applied together with arachidonic acid enhanced the elicitor induced accumulation of sesquiterpenes and sustained steroid-glycoalkaloid levels with transcript levels for the various HMGR mRNAs

equal to or greater than wound-only treatment. These results demonstrate that the consequences of wound- and pathogen-responses of plants are different at the levels of gene expression and associated secondary metabolism.

## INTRODUCTION

The acetate-mevalonate pathway in plants provides a diverse array of isoprenoid compounds that have a vital role in cell structure and function (2, 3, 14, 18). These include phytosterols, dolichols, ubiquinones, phytols, carotenoids, and phytohormones such as ABA and gibberellins. This pathway also produces a variety of stress-induced metabolites such as steroid-glycoalkaloids and phytoalexins that may contribute to defense against insects and pathogens. The basic isoprene unit also is present in a number of other important compounds such as the cytokinines. Recent research suggests that isoprenylation of proteins is an important regulatory mechanism in animal cell, and similar mechanisms may occur in plants(15). In spite of the significance and vital role of these compounds in plants, the organization and regulation of the isoprenoid biosynthetic pathway is poorly understood. HMGR catalyzes the conversion of HMG CoA to mevalonate. HMGR is the rate limiting step in sterol biosynthesis in yeast and mammalian cells and is also considered a key regulatory enzyme in plant isoprenoid biosynthesis. Biochemical and molecular studies with animal and yeast HMGR revealed that regulation of this enzyme is extremely complex and includes controls at the levels of transcription and translation, as well as post-translational modification of the protein(15). Plants have the capacity to synthesize a more complex array of isoprenoid compounds than animals or yeast, suggesting that the regulation of the key enzymes in the pathway may also reflect this diversity and complexity (14). Indeed, earlier biochemical studies of plant HMGR suggest functional compartmentation at the subcellular level and also indicate that multiple forms of HMGR are present(26, 24). The first plant HMGR gene was cloned from Arabidopsis by Learned and Fink(22). Following this work, plant HMGR genes were cloned from potato(7, 8), tomato(23), rubber plant (9), and radish. Using these cloned genes and different plant organs, studies have progressed rapidly on the regulation of HMGR gene expression in different developmental and stress-related contexts. Isoprenoid metabolism is strongly activated in solanaceous plants following mechanical injury or infection(20, 29, 32, 28). Earlier biochemical studies in potato revealed that wounding of the plant induced the rapid accumulation of steroid glycoalkaloids. When the wounded tissues were inocula-

ted with the incompatible race of the fungal pathogen *Phytophthora infestans* or treated with arachidonic acid, an elicitor produced by this fungus, steroid glycoalkaloid biosynthesis was strongly and rapidly inhibited and sesquiterpenoid phytoalexins were accumulated(29, 28). Subsequently, Vogeli and Chappell(33) reported that squalene synthetase and sesquiterpene cyclase, the first enzymes in the branches leading to sterols and sesquiterpenes, respectively, responded inversely following elicitor treatment of tobacco cells. This inverse response also was observed in potato tissues following elicitor treatment or inoculation with incompatible races of *P. infestans* or fungi non-pathogenic on potato that elicit a hypersensitive response (HR; 36, see figure 1). The accumulation of phytoalexins often is associated with the tissue necrosis characteristic of the HR. Although phytoalexins are considered a possible defense response against microbial invasion, definitive proof for a causal role of these compounds in restricting pathogen ingress and colonization is lacking. The identification and cloning of genes for phytoalexin detoxifying enzymes from pathogenic fungi has strengthened the case for phytoalexins as defense compounds (30, 21). However, at this time there is no direct evidence that phytoalexins are crucial for disease resistance in any given interaction. The phytoalexin hypothesis could be tested with phytoalexin-deficient mutants, this approach, however, requires a more detailed understanding of phytoalexin biosynthesis. Accumulation of phytoalexins has been demonstrated in many plant families. Generally, species within a family of plant primarily synthesize phytoalexins of similar structure and biosynthetic origin. For example, in the leguminosae there are more than 100 different phytoalexins that have been characterized and 80% of these are isoflavonoid derivatives. Similarly, the majority of the phytoalexins that have been described in solanaceous plants are terpenoid derivatives(20). Examples of the latter are the sesquiterpenoids rishitin in potato and tomato, and capsidiol in tobacco and pepper. The research presented in this proceeding represents a contribution towards understanding the organization and regulation of the isoprenoid phytoalexin pathway in plants. It is anticipated that this information will be useful for designing strategies to better define the role of these compounds in plant growth and development and in responses to pathogens and other environmental stresses.

## **MATERIALS AND METHODS**

### **Plant and Fungal Materials**

Certified seed grade potatoes (*Solanum tuberosum* L. cv Kennebec) were obtained

from Herman Tim (Department of Vegetable Crop, UC-Davis) and stored at 40°C until 24 hour prior to use when they were placed in darkness at room temperature. To avoid the light effect, manipulation of potato tubers were carried out under a green safelight comprised of two green fluorescent tubes (Sylvania F40-G) wrapped with one sheet each of Rosecolene filter No. 874 (medium blue green) and No. 874 (medium green). Tubers were washed, surface sterilized in 70% ethanol for 3 minutes, then disks (22x5-7 mm) were cut from the central parenchymous tissue with a sterile cork borer. The disks were washed 3 times with sterile distilled water and five were placed on sterile filter paper in petri dish and excess water was removed using sterile filter papers. All treatments and inoculations were applied to upper surface of disk in 50 ul volume and incubated at 20°C. RNAs from other than tuber were extracted from green house grown plant tissue which is directly froze in liquid N<sub>2</sub>. Sporangial inoculums were prepared from *Phytophthora infestans* (Mont.) deBary race 0 and 1 grown on lima bean agar or rye seed media at 20°C. Sporangial suspensions containing 10<sup>5</sup> sporangia were applied to tuber disk aged 20 hours after preparation.

### **Chemicals**

Arachidonic acid (AA) and other chemicals were purchased from Sigma. Immediately before application, AA solution was dried under N<sub>2</sub> gas and resuspended in sterile water by brief sonication. Total 50 ug of AA was applied in 50 ul volume to each tuber disk which show approximately 50% of the maximum response for sesquiterpenoid phytoalexin accumulation (5). Methyl-jasmonate (MJ), Salicylate (SA), Acetylsalicylate (ASA), Salicylhydroxamate (SHAM), and n-Propylgalate (nPG) were treated similar way as AA.

### **cDNA Library Screening**

670,000 plaques from potato tuber cDNA library (obtained from Okita) in lambda gt 11 were transferred to Nylon filters (1). About 800 nucleotides long cDNA probe was constructed from the highly conserved region of Arabidopsis HMGR cDNA (obtained from Learned and Fink, 22) by digestion with NheI. The filters were hybridized with radiolabeled probe under low stringency hybridization condition (5X SSC, 5X Denhardt's, 10% formamide, 100 ug/ml denatured salmon sperm DNA at 42°C) and washed same condition for 2 hours. Forty-seven positive clones were obtained and 11 of them were subcloned on pBluescript II KS(+) (Stratagene) and partially mapped using 8 different restriction endonucleases. Based on the maps, probe were constructed from each end region of cDNA and hybridized with elicitor

induced and uninduced potato RNA. As a results, we selected 3 different cDNA clones which are expressed in potato tuber.

### Nuclear DNA isolation and genomic blot

Nuclei were isolated from potato leaves by the method of Timberlake(27) except that the extraction buffer was pH9.5 (12) and Triton X-100 was added to a final concentration of 0.5% to lyse chloroplasts. Nuclei were lysed in 1% sarkosyl, 0.05M Tris, 0.02M EDTA(pH 9.5), centrifuged, the DNA was precipitated with 0.5 volume of cold isopropanol, and further purified by CsCl/ethidium bromide density gradient centrifugation. DNA were digested 5x recommended units of restriction enzyme and 10 ug/lane were fractionated by electrophoresis in 0.7% agarose gels in TAE buffer and transferred to a Nytran membrane. The DNA blots were hybridized with random-prime labeled probes using standard procedures. Two copy reconstructions were calculated based on a monoploid potato genome size of 0.896 pg, in accord with the classification of the cultivated potato as a tetraploid species (2N=4X=48 chromosome).

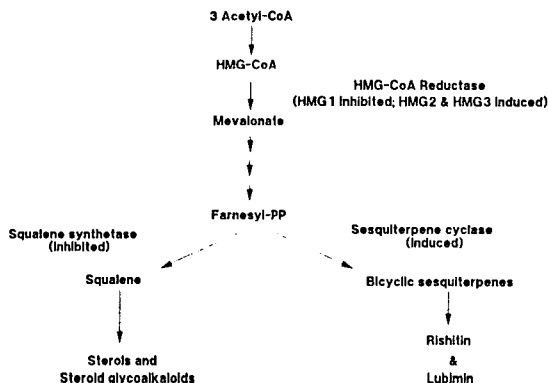


Fig. 1. Outline of stress-responsive isoprenoid biosynthesis in Solanaceous plants. During expression of the hypersensitive response, wound induced sterol/steroid glycoalkaloid synthesis is inhibited in favor of sesquiterpenoid phytoalexin synthesis. HMGR, squalene synthetase, and sesquiterpene cyclase are key enzymatic steps in the pathway affected by elicitor treatment or inoculation with fungi. The subcellular compartmentation of the various steps in the pathway has not been resolved.

### Sequencing and Probe Construction

cDNAs subcloned in pBluscript II KS(+) were digested with proper restriction endonucleases. Nested deletion sets of cDNAs were obtained by Exonuclease III and Mungbean nuclease digestion according to manufacturers guide (Stratagene). DNA sequences were determined by chain termination method (25) using T7 and

T3 universal primers and the Sequanase enzyme according to the manufacturers instructions (United States Biochemicals). Synthetic primers were also designed and used for sequencing unoverlapped regions. DNA sequences were analyzed using personal computer program Microgeni. From the sequences of potato HMGR cDNA variable regions were selected for gene specific probe untranslated region and N-terminal coding region for HMGR 1, untranslated region for HMGR 2 and 3, see 7, 8 and Fig. 2).

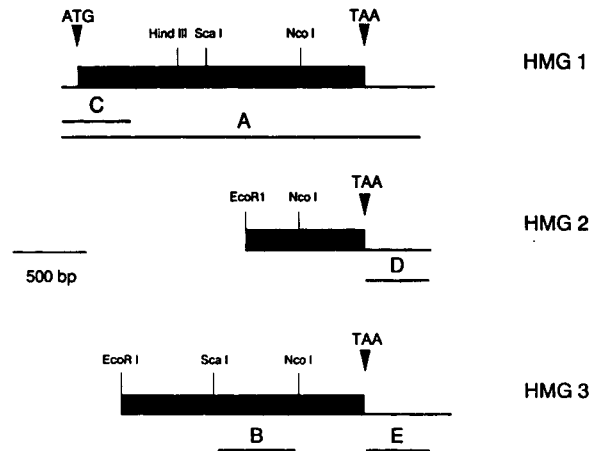


Fig. 2. Gene-specific and conserved region probes used in this study. Regions used for DNA probes in the RNA and DNA gel blots are indicated by corresponding bars with the following designations: A) probe containing entire ORF of hmgl; B) ScaI-NcoI fragment containing a region that is highly conserved among HMGR genes; C) gene-specific probe of hmgl; D) gene-specific probe of hmg2; E) gene-specific probe of hmg3

### RNA Isolation and Northern Blot

Total RNAs were prepared from potato tuber and other tissues by a modification of the method of Parish and Kirby (1966) to include precipitation in 2 M LiCl. Total RNA (20 ug/lane) or polyA+ RNA (1 ug/lane) was fractionated by electrophoresis through 1% agarose gels containing formaldehyde, and transferred to Nytran membranes (S & S inc.). Hybridization of cDNA probes to RNA blots routinely carried out at the same stringency using standard procedures. The amount of labeled probe hybridized to each RNA sample was estimated with a 2-D radioimaging system (Ambis system Inc.).

### Determination of SGA and phytoalexins

SGA were extracted from potato tuber disks by a modification of the procedure of Tjamos and Kuc(29). The amount of SGA was determined by the spectropho-

tometric assay and calculated based on the standard curve of  $\alpha$ -solanine. The sesquiterpenoid phytoalexins were quantified by a semimicro method(17). Concentration of rishtin and lubimin the major sesquiterpenoid phytoalexins in potato, were determined by gas chromatography using standard rishtin and lubimin with methyl-arachidate as an internal standard.

## RESULTS

### Potato HMGRs are encoded by small gene family and expressed in various organs.

To determine the number of genes encoding potato HMGR, nuclear DNA was hybridized with conserved region and gene specific probes. As shown in figure 3, when nuclear DNA was digested with EcoRI and hybridized with conserved region probe, which comes from C-terminal conserved coding region of hmg3, 5.3 kb, 4.8 kb, 3.0 kb, and 2.7 kb DNA fragment were detected. In contrast, when same

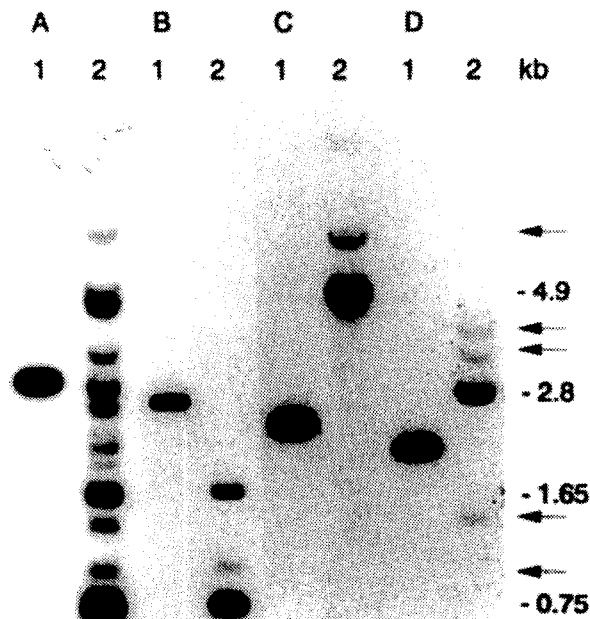


Fig. 3. Southern blot hybridization of potato nuclear DNA with potato HMGR cDNA probes. Nuclear DNA (10 ug/lane) from potato leaves were digested with SacI and HindIII and fractionated by electrophoresis in 0.7 % agarose gel. Blots were hybridized with conserved region and gene specific probes. Panel A; conserved region probe, B;hmg1, C;hmg2, D;hmg3 specific probe respectively. Lane 1 in each panel is two copy reconstruction of HMGR genes.

DNA was hybridized with gene specific probe of *hmg3*, which comes from 3' untranslated region, only 3.0 kb, and 2.7 kb fragments were detected. When DNA was digested with HindIII, conserved region probe detect 16 kb, 6.5 kb, 5.6 kb, 4.8 kb, 2.4 kb, and 2.2 kb but *hmg3* specific probe can detect only 16 and 5.6 kb fragments. These results mean that the potato HMGRs are encoded by small gene family and our gene specific probe detect subset of DNA fragments which were hybridized with conserved region probe. Potato cultivars are known as allote-tetraploid and organization of certain gene in this plant genome is quite complicate. We also obtain nuclear DNA southern blot results using *hmg2* specific probe and replicated same DNA blot. *Hmg2* specific probe detect subset of fragments which is not hybridized with *hmg3* gene specific probe (data not shown).

Total RNA was isolated from uninduced plant organs including roots, tubers, etiolated sprouts, stems, leaves, flower buds, and blossoms to determine expression of HMGR mRNAs (Fig. 4). Total HMGR mRNA which is hybridized with conserved region probe show extremely high level expression in blossoms. This can not be explained by adding up of *hmg1*, 2, and 3 mRNA expression in this organ therefore possibly another gene(s) for HMGR is present and expressed in this organ other than *hmg1*, 2, or 3. High level HMGR mRNA was also detected in etiolated sprouts, stems, and flower buds. Apparently lower level was expressed in roots, tubers, and leaves. Amount of HMGR mRNA expressed in each organs was correlated with the sum of *hmg1*, 2, and 3 except in blossom and each gene is differentially expressed in different organs. *Hmg1* is expressed high level in etiolated sprouts, stems, and flower buds and lower level expression was detected in roots, tuber, leaves, and blossoms. *Hmg2* is expressed in roots, etiolated sprouts, flower buds, and blossoms but low level expression were detected in tubers, stems, and leaves. In contrast, *hmg3* is only expressed in blossoms and in other organs the expression was below detection level with our condition.

#### **Expression of HMGR genes following wounding and elicitor treatments**

To determine the effect of elicitor(arachidonic acid) on the expression of HMGR genes in potato tuber, gene specific probes and conserved region probe were subjected to northern blot with potato RNA extracted from AA-induced and wounded potato slices. Total HMGR mRNA, which is hybridized with conserved region probe was dramatically increased right after wounding and elicitor treatment (Fig. 5-A). The relative abundance of total HMGR mRNA was about 3 times more in elicitor treatment and this induced level stayed until 96 hour after treatment. The amount and time of transcripts increase is correlated with the increase of p105,000g



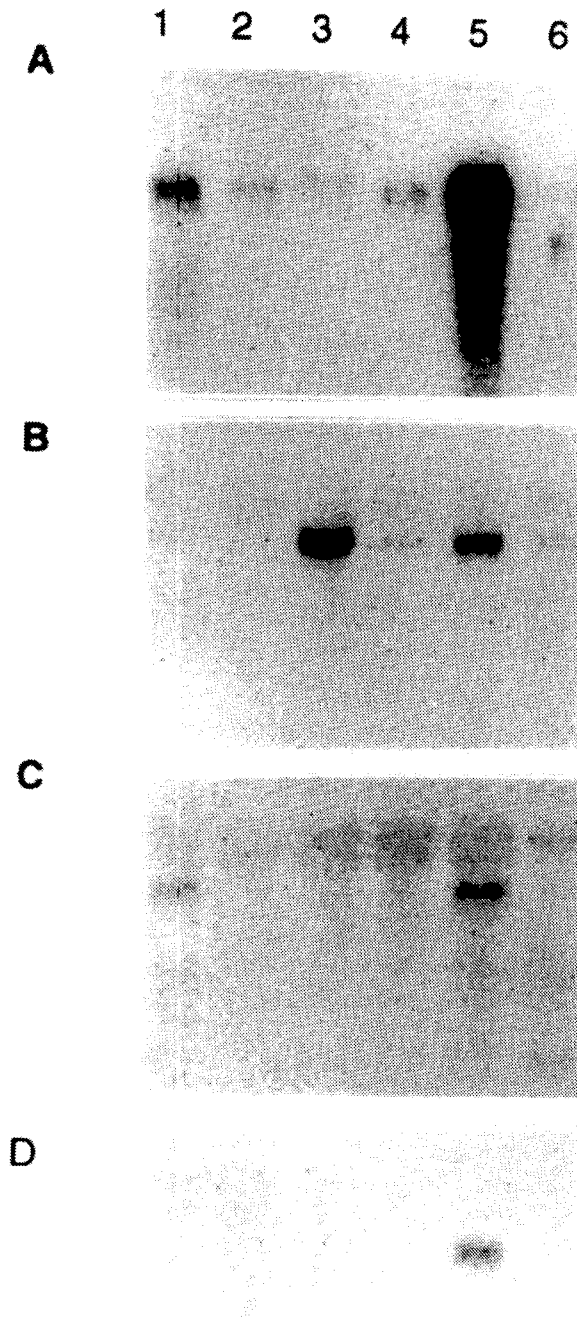


Fig. 4. Constitutive expression of HMGR genes in various organs of the potato plants. Total RNA(20 ug) from roots(lane 1), stems(lane 2), flower primordia(lane 3), petals(lane 4),anthers(lane 5), and pistils(lane 6) were hybridized with A) conserved region probe, B) hmg1 specific probe, C) hmg2 specific probe, D) hmg3 specific probe. These probes detected transcripts of approximately 2.5 kb.

HMGR activity after elicitor treatment (26). In contrast to the HMGR enzyme activity is decreased 24 hours after treatment and stay background level 48 hour, the induced mRNA stayed until 96 hour which is the last data point of our experiment. This results imply that some post-transcriptional level regulation is also involved in HMGR enzyme activity expression. In contrast to the total HMGR mRNA expression, Hmg1 transcripts(2.6 kb) were dramatically increased by wounding but this wound induction is almost completely shot down (Fig. 5-B) by wounding and simultaneous elicitor treatment. Elicitor effect on the expression of potato hmg1 gene expression is not super-imposed on the wound effect. Hmg2 and hmg3 gene specific probes detected similar sizes mRNA which is dramatically induced by elicitor but slightly induced by wounding(Fig. 5-C, D). Expression profiles of these genes were closely correlated with the total HMGR mRNA expression profile.

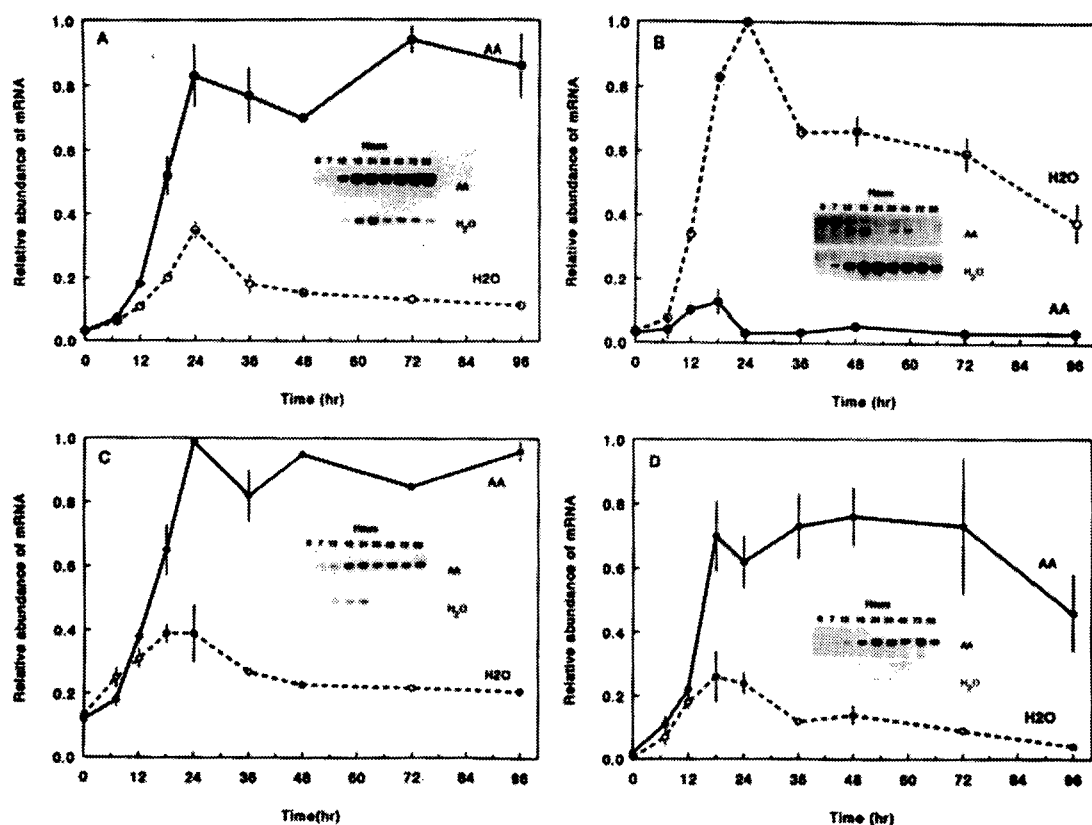


Fig. 5. Differential expression of potato HMGR genes following wounding and arachidonic acid treatment. A; expression of total HMGR mRNA which is hybridized with conserved region probe following wounding and elicitor treatment. B; expression of HMG1, C; expression of HMG2, C; expression of HMG3. RNA samples were extracted following AA application (50 ug in 50 ul H<sub>2</sub>O per potato disk) and wound control (50 ul H<sub>2</sub>O).

### Pathogen challenging and expression of HMGR genes

To determine the expression of potato HMGR genes after inoculation of pathogens, potato disks were prepared and incubated 20 hours at 20°C then inoculated with zoospores of compatible or incompatible races of *Phytophthora infestans*. RNA samples were extracted from these potato disks and hybridized with gene specific and conserved region probes. Total HMGR mRNA which is hybridized with conserved region probe was dramatically increased within 3 hour after inoculation and reached to peak at 6 to 12 hours after inoculation with race 0 (incompatible) and race 1, 2, 3, 4 (compatible) respectively (Fig. 6-A). Abundance of HMGR mRNA is decreased after peak but still present higher level than water control until 48 hours after inoculation. Water treatment show only 10 to 15% of HMGR mRNA accumulation compare with maximum level. Induction of total HMGR mRNA which detected by conserved region probe does not show significant difference between

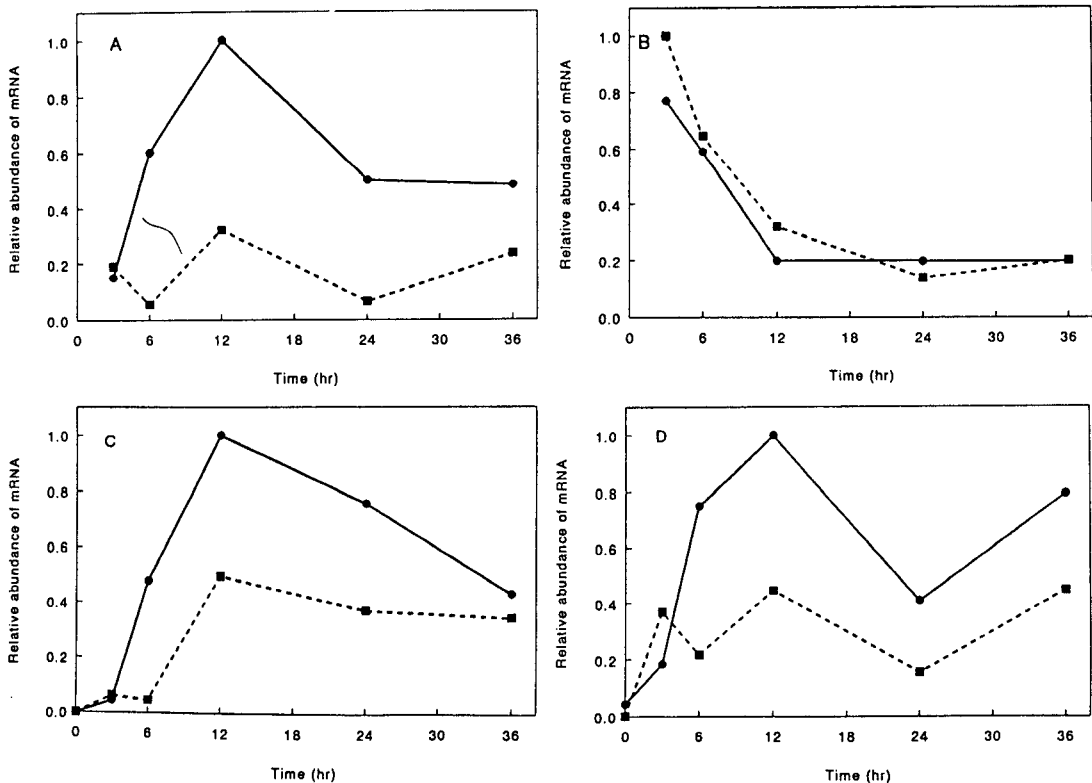


Fig. 6. Differential expression of potato HMGR genes following inoculation with *Phytophthora infestans*. A.; B.; C.; and D; same as fig. 5. *Phytophthora infestans* race 0 is incompatible to the potato cultivar kennebec which we used in this experiment and race 1, 2, 3, 4 is compatible to this potato cultivar. Pathogens were inoculated with  $10^6$  sporangia per disk in 50 ul water 20 hours after preparation of potato disks. As a control 50 ul water was applied.

compatible and incompatible races of pathogen inoculation. Hmg1 mRNA expressed high level before inoculation (wound inducible and peak at 24 hours after wounding) but after inoculation with both pathogens the abundance of hmg1 mRNA dropped down to background level within 6 hours (Fig. 6-B) and stay at this level until 48 hours. Usually disease symptom caused by *P. infestans* is appear 48 to 72 hour after inoculation and the suppression of hmg1 mRNA during this period is surprising. Hmg2 mRNA is induced by both pathogen races and peaked at 12 hours after inoculation (Fig. 6-C). Water control stay 20 % of maximum mRNA level. Hmg3 mRNA is also induced by both pathogen (Fig. 6-D) but the induction was rather rapid and earlier in incompatible pathogen than compatible pathogen inoculation. In water control, hmg3 mRNA is only expressed at about detection level with our condition (0-5 % of maximum).

#### **Differential expression following MJ treatment**

Because the maximum level transcripts were detected 24 hr after wounding and AA treatment, HMGR mRNAs were monitored at this time point in the experiments described here. RNA samples that were hybridized with conserved region probe yielded a single size transcript of approximately 2.5 kb that were strongly induced following treatment of potato disks with high concentration of MJ(2 or 4 umole per disk, Figure not shown). The abundance of total HMGR mRNA transcripts did not correspond with SGA accumulation(Fig. 7). Wound induced transcript levels of hmg1 were strongly enhanced by treatment of the tissues with low concentration of MJ(400 pmole per disk). The increase in hmg1 mRNA levels following MJ treatments were concentration dependent, and high concentration of MJ (2 umole per disk or more) completely abolished hmg1 mRNA accumulation (Fig. 7). The maximum level of hmg1 mRNA accumulation was detected following treatment with 0.4 nmole MJ per disk, and enhancement was detected as low as 40 pmole per disk. The maximum level of SGA accumulation was detected when 4 nmole of MJ was applied on a potato disk and a significant increase in SGA accumulation was detected even when 0.4 pmole MJ was applied on a disk(Fig. 7). The abundance of hmg1 transcripts corresponded closely with SGA accumulation. In contrast to the expression pattern for hmg1, hmg2 and hmg3 mRNA levels appears to be unrelated to SGA accumulation following treatment of the disks with MJ (data not shown). Hmg3 mRNA abundance was strongly enhanced following treatment with high concentrations of MJ which suppressed SGA accumulation.

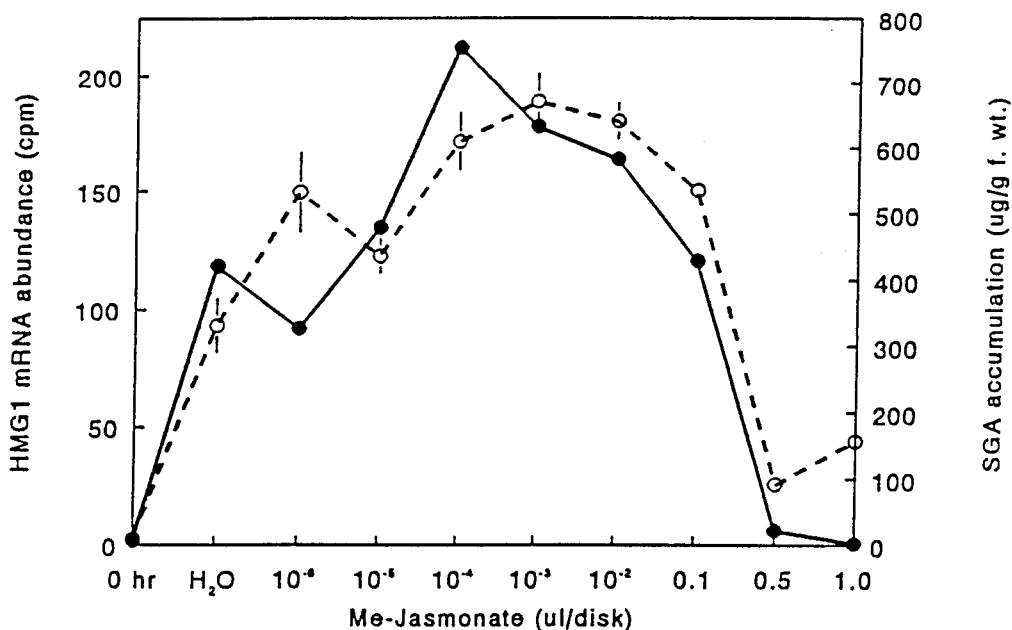


Fig. 7. Effects of different MJ concentrations on hmg1 mRNA and SGA accumulation. An RNA blot was hybridized with hmg1 specific probe and counted (solid line) with an Ambis 2-D radioisotope imaging system. Steroid-glycoalkaloid accumulation (broken line) was determined 96 hrs after treatment are indicated. MJ 0.1 ul is correspond to 0.4 umole.

#### Effects of lipoxygenase inhibitors on HMGR mRNA and SGA

Because of the effect of exogenous MJ on hmg1 mRNA and SGA accumulation, we tested the hypothesis that endogeneous jasmonate is responsible for wound induction of hmg1 and SGA accumulation. SHAM and nPG were used to inhibit lipoxygenase activity which is required for the biosynthesis of jasmonate in plants (31). The wound induction of total HMGR mRNA levels were not significantly affected by SHAM or nPG treatment (data not shown). However, wound induced hmg1 mRNA levels were reduced to 38% and 15% of the control levels by SHAM and nPG treatment respectively (Fig. 8). There was a corresponding reduction in the accumulation of SGA (Fig. 8). The reduction of hmg1 mRNA and SGA accumulation by these inhibitors was completely overcome by treatment of the tissue with MJ (40 nmole per disk) (Fig. 8). SHAM at the levels that were effective in suppressing the hmg1 and SGA response did not inhibit the expression of hmg2, hmg3, or total HMGR mRNAs (data not shown). In contrast, and unlike earlier studies wherein SHAM and nPG partially suppressed AA phytoalexin elicitor activity, these inhibitors did not suppress the AA effect on the levels of hmg2 and hmg3 mRNAs (data not shown). These results show that levels of hmg1 mRNA are closely correlated

with SGA accumulation, and that the levels of *hmg2* and *hmg3* mRNAs are not related to SGA accumulation in potato tuber.

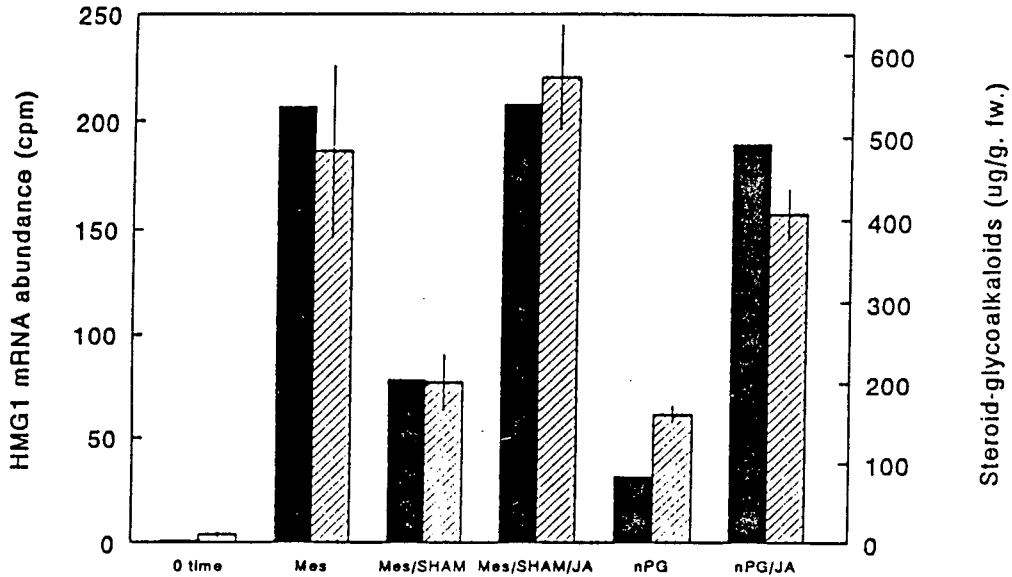


Fig. 8. Effect of lipoxygenase inhibitors, MJ, and AA on *hmg1* mRNA and SGA accumulation. *Hmg1* mRNA level (solid bar) and steroid-glycoalkaloid accumulation (dashed bars) are indicated. Five mM solution (SHAM and nPG) and fifty nmole MJ was treated.

#### Specificity of AA and MJ effects on HMGR gene expression

To further examine the specificity of the response of HMGR genes, several compounds known to induce disease resistance and to activate defense related genes in plants were tested for their effects on potato *hmg1* gene expression. SA and ASA induce pathogenesis related proteins in plants and SA may function as a signal for systemic acquired resistance in some species of plants. In potato disk, both SA and ASA inhibited wound induction of *hmg1* gene expression and suppressed SGA accumulation (Fig. 9). The concentration of SA and ASA used in this experiment did not affect the expression of *hmg2* and *hmg3* (data not shown). The high concentration of MJ (2 umole per disk) used in this set of experiments induce *hmg3* gene expression but inhibited *hmg1* gene expression and did not elicit SGA or sesquiterpenoid phytoalexins.

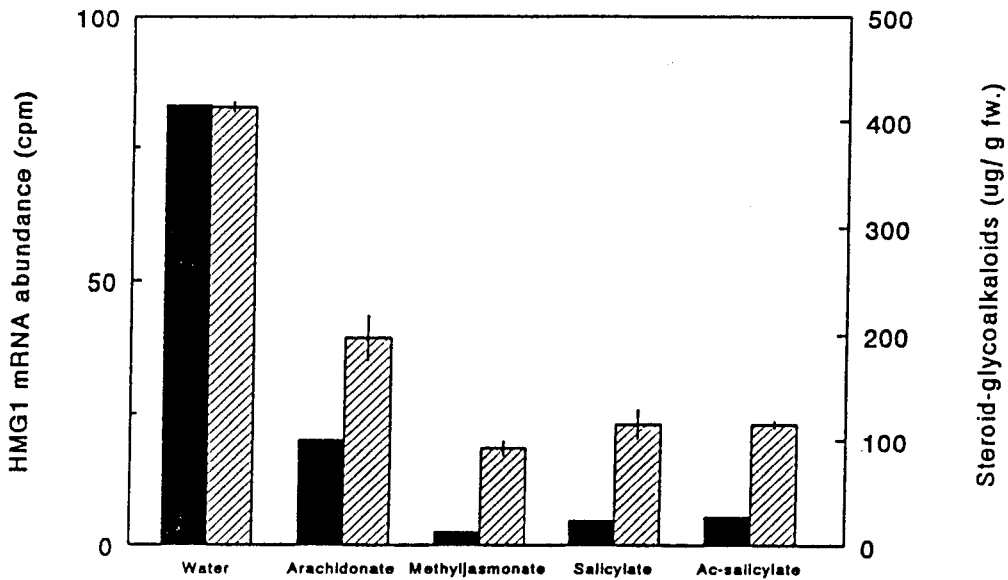


Fig. 9. Effects of chemicals which inhibit wound-inducible hmg1 mRNA levels on SGA accumulation. RNA samples were extracted 24 hrs after treatment of the disks and hybridized with hmg1 specific probe(solid bars). Accumulation of SGA(dashed bars) were analyzed 96 hrs after treatment. Values for SGA are the means and standard deviations from 3 replicated samples. Results for hmg1 mRNA are representative of at least two different experiments. AA(0.16 umole/disk), MJ(2 umole/disk), SA(5 mM solution), and ASA(5 mM solution) were tested.

## DISCUSSION

In this paper we demonstrated cloning of three genes for potato HMGR and expression of these genes following wounding, elicitor treatment, and pathogen inoculation. At least three genes are expressed in potato tuber, one is wound inducible (HMG1) but the wound induction of this gene expression is suppressed by elicitor or pathogen treatment and the others (hmg2, and 3) are elicitor and pathogen inducible which are also slightly induced by wounding. This is the first report that a members of same gene family in plant is undergoing reverse regulation by same environmental signal.

### Special Feature of Potato HMGR genes

Sequences of potato HMGR genes have high level homology both in amino acids

and nucleotides with other cloned plant HMGR genes (22, 23, 9). Potato HMGR cDNA show highly conserved amino acid sequences in catalytic domain. Furthermore, each potato HMGR gene has higher homology with corresponding tomato HMGR genes (Narita and Grisse personal communication) than homology among potato HMGR genes. Extremely high nucleotide homology was also extended to 3'-untranslated region of each pairs of gene in potato and tomato which is much higher than homology within same plant. It is not known that why these genes from different plants have more homology even in 3'-untranslated region is not clear but we assume that each pair of corresponding HMGR genes in potato and tomato may have specialized role in plant isoprenoids biosynthesis. It is known that yeast and mammalian HMGR have 7 membrane spanning domain in amino terminal region of HMGR protein (13) which is involved in sterol mediated degradation of HMGR protein (15). In plant, only 2 membrane spanning domains are known in Arabidopsis (22) and tomato (23 and unpublished results). Potato hmg1 has identical pattern of hydrophathy with other known plant HMGR protein but the role of this domain in protein degradation or posttranslational regulation is unknown.

N-terminal amino acid sequences of plant HMGR is highly conserved among plant (MDxRRRPxKP...) but the role of this conserved amino acids is not clear.

### **Role HMGR genes in Plant Isoprenoid Biosynthesis**

HMGR is known as the rate-limiting step enzyme in sterol biosynthesis in mammalian system but the essential role of HMGR in plant isoprenoid biosynthesis is still controversial (3). Recent studies in tobacco cell suspension culture (6) and potato tuber (26) show that the elicitor inducible HMGR activity is required for the accumulation of sesquiterpenoid phytoalexins in Solanaceous plants. Biochemical studies of isoprenoid metabolism also show that Sesquiterpene Cyclase a branching point enzyme of sesquiterpene biosynthesis from farnesyl-PP is induced by either elicitor treatment or pathogen inoculation (33, 36,). In contrast, Squalene Synthetase a branching point enzyme from farnesyl-PP to sterols was dramatically suppressed by elicitor or pathogen (32, 36). These data are closely correlated with the accumulation of end-products following wounding and elicitor treatment (19, 28). As a overall feature, accumulation of sterols by wounding is rapidly shot down and carbon flow is redirected toward sesquiterpene biosynthesis by elicitor treatment or pathogen inoculation. Our results, suppression of wound inducible hmg1 mRNA and induction of hmg2 and 3 mRNA after elicitor treatment, have close relationship with the branching step regulation of sterols and sesquiterpenes biosynthe-



sis. Coordinate regulation of *hmg1* mRNA expression and SS activity by wounding and elicitor treatment suggest us *hmg1* may have role in sterol biosynthesis. *Hmg2* and *3* mRNA is also coordinately regulated with SC activity which lead to accumulation of sesquiterpene phytoalexins. These results may imply that there maybe organizational channels for biosynthesis of certain species of isoprenoids and those enzymes which involve in same channels are coordinately regulated in planta.

**Lipid-derived signals discriminate wound- and elicitor-responsive pathways in potato.**

Jasmonic acid and MJ are potent inducers of proteinase inhibitors in potato and tomato(11). Because proteinase inhibitors are also induced by several other types of compounds, such as oligogalacturonides, peptide systemin, and the fungal elicitor chitosan, a model was suggested that jasmonic acid is an intermediate involved in the activation of defense responses to insects and pathogens(11). The notion that jasmonate is involved in plant defense against pathogen attack is also extended by Gundlach, et. al. (16), who demonstrated that MJ and jasmonic acid accumulate in plant cell suspension cultures following treatment with a fungal elicitor. Furthermore, MJ alone functioned as an elicitor and induced phenylalanine ammonia lyase and phenolic alkaloid accumulation. At present, however, there is no evidence that either proteinase inhibitors or alkaloid accumulation are defense mechanism against pathogens. It is unlikely that proteinase inhibitors contribute significantly to disease resistance since wounding, a treatment that induces high levels of the inhibitors in potato and tomato does not induce resistance. The proposed role of jasmonate as an inducer of disease resistance is premature, at best, and an oversimplification. For the potato responses examined in the present study, MJ affected only wound-inducible *hmg1* gene expression and SGA accumulation. No phytoalexin accumulation or induced resistance to *P. infestans* was detected after treatment of the disks with of the MJ concentrations tested. An earlier study(10) demonstrated that the ASA, SA and related hydroxybenzoic acids inhibit wound responses, including induction of proteinase inhibitors in tomato plants. Subsequently, it was reported that SA is induced systemically following inoculation of tobacco and cucumber plants with various necrotic pathogens, and induces systemic acquired resistance, perhaps, inpart, its elicitation of pathogenesis-related proteins(34, 35). Wounding does not induce most of the pathogenesis-related proteins in tobacco and does not induce systemic resistance. In the present study, SA and ASA inhibited wound inducible *hmg1* gene expression and SGA accumulation. Hence, there are similarities in the effects of the hydroxybenzoates on tobacco and potato with the effects of AA on potato; specially, SA and its analogs suppress certain wound

responses. However, unlike AA, SA does not induce phytoalexin accumulation or resistance to *P. infestans* in potato, a results that indicates that the signal transduction pathways connecting AA or SA treatment and the plant response are different. In conclusion, the results of the study show that two stress-responsive isoprenoid pathways are regulated by different lipid-derived signals present during the plant-pathogen interaction. Thus far, these studies are consistant with the hypothesis that MJ is an endogeneous signal that participate in the regulation of wound-healing programs. These experiments clearly demonstrate that the consequences of wounding and elicitor treatment are different at the level of gene expression and associated secondary metabolism. Collectively, these results and those of other studies reveal that plants discriminate wounding and pathogen attack, and indicate that the responses observed during hypersensitivity expression are not simply an enhancement of wound-responses.

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