An in planta induced gene of Phytophthora infestans codes for ubiquitin

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Abstract

An in planta induced gene of Phytophthora infestans (the causal organism of potato late blight) was selected from a genomic library by differential hybridization using labelled cDNA derived from poly(A) + RNA of P. infestans grown in vitro and labelled cDNA made from potato-P. infestans interaction poly(A) + RNA as probes. Sequence analysis showed that the gene codes for ubiquitin, a highly conserved protein which plays an important role in several cellular processes. The structure of the polyubiquitin gene (designated ubi3R) is consistent with the structure of other known polyubiquitin genes. It consists of three repeats in a head-to-tail arrangement without intervening sequences, each encoding a ubiquitin unit of 76 amino acids. The last ubiquitin unit is followed by an extra asparagine residue at the carboxy-terminal end. Northern and Southern blot analyses revealed that the polyubiquitin gene is a member of a multigene family, all genes of which show induced expression in planta.

Introduction

Potato late blight caused by the fungus Phytophthora infestans (Mont.) de Bary (Oomycetes) is one of the most important diseases of potato. Leaves and tubers of susceptible cultivars become readily infected by this pathogen. The fungus spreads rapidly through the plant tissue causing a destructive necrosis. Resistance to late blight in potato can be based on either a single gene or multiple genes. In general, single-gene-based resistance, governed by the so-called R genes, is characterized by a hypersensitive response resulting in a rapid cell death of invaded cells. Multiple-gene-based resistance is characterized by a low infection efficiency, slow tissue colonization and low sporulation rate. R-gene-mediated resistance is commercially not attractive because it becomes rapidly ineffective due to the appearance of new virulent strains of the fungus. Breeding efforts, therefore, aim at introducing durable multiple-gene-based resistance into commercial potato cultivars. The physiological basis of this type of resistance is hardly understood and, as a

The nucleotide sequence data reported will appear in the EMBL, GenBank and DDBJ Nucleotide Sequence Databases under the accession number X55717.
consequence, breeders have to rely on empirical methods.

It can be assumed that the establishment of a pathogenic relation between the potato plant and *P. infestans* involves the mutual interference in cellular processes of each partner. Defence responses of host tissue being colonized (e.g. accumulation of pathogenesis-related (PR) proteins and induction of enzymes of the phenylpropanoid pathway) are relatively well studied [9, 10, 16, 25], but nothing is known about changes in metabolism of pathogen in response to the host. In other plant-microbe interactions – pathogenic (e.g. dicots-Agrrobacterium tumefaciens and pea-Fusarium solani f.sp. pisi) as well as symbiotic interactions (legumes-Rhizobium spp.) – signal molecules originating from the host have been characterized which induce expression of genes involved in establishing the interaction [24, 41, 48]. It is therefore tempting to suggest also in the potato-*P. infestans* interaction that host factors induce physiological responses in the pathogen which are necessary for pathogenesis and which are mediated by pathogenicity genes. In view of this, both defence responses of the host and the degree to which pathogenicity genes are activated in the pathogen ultimately determine speed and efficiency of the infection and colonization process and the intensity of sporulation. The result of these processes is reflected in the degree of durable resistance. The identification and characterization of *in planta* induced genes of *P. infestans* therefore will lead to a better understanding of the molecular basis of both pathogenicity of the fungus and durable resistance of the host.

The present study concerns the characterization of a *P. infestans* gene which was isolated from a genomic library by differential screening. The probes used were cDNA made from poly(A)+ RNA of *P. infestans* grown *in vitro* and cDNA synthesized on poly(A)+ RNA isolated from potato leaves infected with a compatible *P. infestans* race. This gene, designated *ubi3R*, codes for ubiquitin, one of the most conserved proteins known to date. The significance of the *in planta* induced expression of this gene will be discussed.

### Materials and methods

**Culturing of Phytophthora infestans and inoculation of potato leaves**

Strain 88069 of *P. infestans* (Mont.) de Bary (Al-mating type, race 1.3.4.7) was isolated from a naturally infected tomato plant in the summer of 1988 and was maintained on rye-agar medium containing 2% sucrose [8]. Liquid cultures in 25 ml Henniger synthetic medium [21] were initiated from zoospores obtained from 2-week-old cultures on rye-sucrose agar at a concentration of 2 x 10^7 zoospores/ml. Cultures were incubated for 14 days at 20 °C to obtain mycelium for DNA and RNA extractions.

Leaves of the potato cultivar ‘Ajax’, which carries the R3 gene for late blight resistance, were inserted in florist’s foam oases saturated with water and inoculated with *P. infestans* by spraying a sporangial suspension (5 x 10^5 sporangia/ml) on the axial side of the leaflets. After inoculation, the leaves were incubated at 18 °C at 100% RH in plastic boxes with a transparent lid under cool fluorescent light for 16 h per day. As a control, uninoculated leaves were treated in the same way. Under these conditions the symptoms of the late blight disease develop from small necrotic lesions, which are first visible 24 h after inoculation, to completely ‘water-soaked’ leaves on day 3. During this period leaves were collected for RNA isolation. Four days after inoculation the fungus sporulates extensively. Secondary infections by necrotrophic bacteria prohibit the isolation of good-quality RNA in this stage.

**Isolation of genomic DNA and construction of a genomic library**

Genomic DNA of *P. infestans* was isolated from mycelium grown in liquid culture. Mycelium (10 g fresh weight) was ground in liquid nitrogen to a fine powder and mixed in 5 ml of extraction buffer (200 mM Tris-HCl pH 8.5, 250 mM NaCl, 50 mM EGTA, 0.8% (w/v) tri-isopropylphthalene sulphonic acid (TNS) and 0.48% (w/v) 4-amino- salicylic acid) per gramme of mycelium and incubated at 55 °C for 5 min. The mixture was
extracted once with 0.6 volume of water-saturated phenol (55 °C). The water phase was then extracted once with an equal volume of phenol/chloroform/isoamyl alcohol (25:24:1) and once with an equal volume of chloroform/isoamyl alcohol (24:1)). The DNA was precipitated with 0.6 volume of 2-propanol, dissolved in Tris-HCl pH 8.0, 1 mM EDTA) and further purified by CsCl gradient centrifugation [28].

A genomic library was constructed in the replacement vector λEMBL3 [15], according to Frischauf [14] with minor modifications. High-molecular-weight DNA was partially digested with Sau3AI and fragments of 15 to 23 kb were isolated after centrifugation on a 10–40% sucrose gradient. Of these fragments 300 ng was ligated to 250 ng of λEMBL3 Bam HI arms (Promega). Packaging of the ligated DNA, using the Packagene in vitro packaging system of Promega, was performed according to the manufacturer’s instructions.

*Isolation of RNA and poly(A)⁺ RNA and preparation of labelled cDNA probes*

Total RNA was prepared from mycelium of *P. infestans* grown in liquid cultures and from colonized leaflets (interaction RNA) using the guanidine hydrochloride RNA extraction method as described by Logemann et al. [27]. To check for integrity of RNA, the RNA was electrophoresed in a 1.5% agarose-TBE (89 mM Tris-HCl, 89 mM boric acid, 2 mM EDTA) gel containing 0.5 μg/ml ethidium bromide. Poly(A)⁺ RNA was isolated by oligo(dT)-cellulose chromatography as described by Maniatis et al. [28]. First-strand cDNA with a specific activity of ca. 1–2 × 10⁸ cpm/μg was prepared from 1 μg poly(A)⁺ RNA as described by Sargent [39] using M-MLV reverse transcriptase (Gibco BRL) and oligo(dT) to prime the synthesis reaction.

*Differential screening, subcloning and sequencing*

The genomic library was plated on a recombinant deficient *Escherichia coli* host, strain MB406 (supE, recB21, recC22, sbcB15, hflA, hflB, hsdR⁻), which enables the propagation of a non-biased genomic library [49]. Four replica filters of the genomic library were made on Hybond-N⁺ membrane (Amersham) according to the instructions of the manufacturer. The library was differentially hybridized in duplicate at high stringency in hybridization mix containing 5 × SSC (750 mM NaCl, 75 mM sodium citrate), 5 × Denhardt’s solution (0.1% (w/v) Ficoll, 0.1% (w/v) polyvinylpyrrolidone, 0.1% (w/v) BSA (fraction V)), 0.5% SDS and 100 μg/ml calf thymus DNA at 65 °C for 16 h using as probes labelled cDNA (10⁷ cpm) derived from poly(A)⁺ RNA of the fungus grown *in vitro*, and labelled cDNA (10⁷ cpm) derived from interaction poly(A)⁺ RNA (3 days after inoculation). The filters were washed in 2 × SSC/0.5% SDS at 65 °C and exposed to Kodak X-Omat S film for 2–7 days at −80 °C. The selected plaques were purified by a second round of differential hybridization as described above.

DNA from the selected lambda clone was digested with several restriction enzymes and separated on a 0.8% agarose-TBE gel. Following electrophoresis, the DNA was transferred to Hybond-N⁺ membrane (Amersham) by capillary transfer [28] and hybridized at high stringency (65 °C) as described above using ³²P-labelled interaction cDNA as probe. The membrane was washed in 2 × SSC/0.5% SDS at 65 °C and exposed to Kodak X-AR film for 24 h at −80 °C. The hybridizing *Ssr I* fragment was subcloned into pTZ19U using standard techniques [28] which resulted in the plasmid pUB-S (Fig. 1B). Both strands of overlapping subclones of pUB-S were sequenced by the dideoxy chain termination method [38] using a sequencing system (Promega) for sequencing on double-stranded DNA. For analysing the sequence data and to screen the EMBL Data Library [20] and the GenBank databank [5], the Sequence Analysis Software Package, Version 6.0, of the Genetics Computer Group (GGG) of the University of Wisconsin was used.
**Southern blot analysis of genomic Phytophthora infestans DNA**

Genomic DNA (10 μg) of *P. infestans* was digested with *Kpn I*, *Eco RI*, *Bam HI*, *Hind III*, and *Sst I* and size-separated on a 0.7% agarose-TBE gel. Following electrophoresis, the DNA was transferred to Hybond-N*+* membrane (Amersham) by capillary transfer and hybridized at high stringency (65 °C) as described above to the two 32P-labelled 228 bp *Pvu II* fragments from the *ubi3R* coding region (Fig. 1). Probe was made by random primer labelling using Promega's Primera-Gene labelling kit. The blot was washed in 0.5× SSC/0.1% SDS at 65 °C and exposed to Kodak X-Omat S film for 16 h at −80 °C.

**Northern blot analysis and primer extension of poly(A)+ RNA**

For northern blot analysis, 15 μg of total RNA was denatured and electrophoresed on a 1.5% agarose-formaldehyde denaturing gel as described by Maniatis *et al.* [28]. Following electrophoresis, the RNA was transferred to Hybond-N*+* membrane (Amersham) by capillary transfer and hybridized at high stringency (65 °C) as described above using the two 32P-labelled 228 bp *Pvu II* fragments from the coding region of *ubi3R* as probe (Fig. 5A). The blot was washed in 0.5× SSC/0.1% SDS at 65 °C and exposed to Kodak X-Omat S film for 2 days at −80 °C. Probe was then removed by immersing the membrane in a solution of boiling 0.1% SDS for 5 min. The blot was rehybridized as described above using a gene-specific probe from the upstream region of the *ubi3R* gene (Fig. 5B). The gene-specific upstream DNA fragment was generated from pUB-S by PCR using a primer (5′-GGTTGCTCCTGTTTATG-3′) complementary to the sequence at position −2 to −19 relative to the ATG start codon in the *ubi3R* gene and the pUC/M13 sequencing primer (5′-GTTTTCCAGGTCACGAC-3′) complementary to a sequence in the vector. The reaction mixture (100 μl) contained 10ng pUB-S, 120 ng of each primer, 0.2 mM dNTPs and 2 units of AmpliTaq.
DNA polymerase (Perkin Elmer Cetus) in Taq polymerase buffer (10 mM Tris-HCl pH 8.3, 50 mM KCl, 1.5 mM MgCl₂, 0.001% gelatin). Amplification was performed in a DNA Thermal Cycler in 18 cycles of 30 s at 94 °C, 30 s at 45 °C and 90 s at 72 °C. The 799 bp PCR product was labelled with $^{32}$P-dATP by random primer labelling as described above. After hybridization and exposure, the blot was again deprobed and rehybridized, now using as probe the 2.8 kb Pst I insert from pSTA31 containing the constitutively expressed actin (actA) gene of P. infestans [46] (Fig. 5C).

To map the 5' end of the ubi3R transcript, an oligonucleotide (5'-GGTGCTCGGTAT-TAG-3') complementary to the sequence at position -2 to -19 relative to the ATG start codon was labelled with $^{32}$P at its 5' end using T<sub>4</sub> poly-nucleotide kinase (Promega) according to the manufacturer's instructions. Labelled primer (5 ng) was annealed to 3 μg of poly(A)<sup>T</sup> RNA isolated from in vitro grown P. infestans mycelium and from un inoculated potato leaves as a control. Extension of the primer was performed in a reaction mixture (25 μl) containing 400 units M-MLV reverse transcriptase (Gibco BRL), 50 mM Tris-HCl pH 8.3, 75 mM KCl, 10 mM DDT, 3 mM MgCl₂, 0.1 μg/μl nuclease-free BSA, 200 μM of each dNTP as 37 °C for 45 min. The primer extension products were analysed by electrophoresis on a polyacrylamide gel (89 mM Tris-HCl, 89 mM boric acid, 2 mM EDTA, 8 M urea, 5.7% acrylamide, 0.3% bisacrylamide) and detected by autoradiography.

Results

Selection of in planta induced Phytophthora infestans genes from a genomic library and mapping of one of these genes on the selected lambda clone

As calculated from microfluorometric determinations of nuclear DNA content [45], the haploid genome length of P. infestans is approximately 2.7 × 10<sup>9</sup> bp. In order to achieve a 99% probability of having any DNA sequence represented in the library, 80,000 recombinant plaques were screened. Differential screening of the genomic library with fungal and interaction cDNA is performed under non-saturating conditions. Under these conditions the intensity of each obtained signal corresponds with the abundance of a particular cDNA in the probe. A stronger signal obtained after hybridization with interaction cDNA as compared to the signal obtained after hybridization with fungal cDNA indicates a higher abundance of those cDNAs which are complementary to the DNA in the hybridizing lambda clone. Since the filters are hybridized with equal amounts of labelled cDNA, the procedure even underestimates the difference in abundance because the quantity of fungus-derived cDNA present in the interaction cDNA probe is much less than in the fungal cDNA probe. Thirteen plaques gave a relatively strong signal after hybridization with labelled interaction cDNA and a relatively weak or no signal after hybridization with labelled cDNA of the fungus grown in vitro. Approximately fifty additional plaques gave comparable signals after hybridization with both cDNA probes. One of the strongly differential hybridizing plaques (DHCl) was purified after a second round of differential hybridization. Southern analysis of various restriction fragments of DHCl with interaction cDNA as probe revealed a strongly hybridizing Sst I fragment of 3.2 kb. This fragment containing a putative in planta induced gene was subcloned in pTZ19U and from the obtained plasmid pUB-S a restriction map was constructed. The approximate location of the coding region of the differentially expressed gene (closed bar in Fig. 1) was determined by Southern blot analysis using interaction cDNA as probe.

Differentially expressed gene codes for polyubiquitin

Several fragments of the 3.2 kb Sst I insert of pUB-S were subcloned and the sequence of 1632 nt (nucleotides) was determined by dideoxy sequencing on both strands of various overlapping clones (Fig. 2). The sequencing strategy is
Fig. 2. Nucleotide sequence of the Phytophthora infestans polyubiquitin gene (ubi3R) and corresponding amino acid sequence of the encoded protein. The transcription start site as determined by primer extension is indicated by an asterisk. Numbered methionine residues in the amino acid sequence represent the start of the three ubiquitin units. Underlined nucleotides in the 5' non-coding region match the consensus sequence for heat shock promoter elements (CNGAANNTTCNNG [4]). Overlined DNA sequence (AATTAAAA) in the 3'-flanking region of the polyubiquitin gene represents a possible polyadenylation signal.
summarized in Fig. 1. An open reading frame of 687 nt containing three almost identical 228 nt repeats was found in the 1632 nt sequence (Fig. 2). Comparison of the 687 nt sequence with the sequence databank revealed that the sequence codes for polyubiquitin. The coding region of the polyubiquitin gene encodes three ubiquitin units with a length of 76 amino acids in a head-to-tail arrangement followed by an extra asparagine residue at the carboxy-terminal end. Although the ubiquitin-coding repeats within the gene differ by up to 8 out of 228 bp, they code for identical amino acid sequences (Fig. 2)). Only 4 or 5 amino acids of the 76 amino acid ubiquitin sequence differ from the sequence of barley, human, chicken and yeast ubiquitin (Fig. 3). Because of the three ubiquitin repeated units we have designated the gene *ubi3R*.

In order to map the 5' end of the *ubi3R* transcript, primer extension was performed on poly(A)*+ RNA isolated from the fungus grown *in vitro* and from uninfected potato leaves using an oligonucleotide complementary to the *ubi3R* sequence at positions −2 to −19 relative to the ATG start codon to prime the synthesis reaction. The primer extension products were electrophoresed next to the products of a sequencing reaction using the same oligonucleotide as primer and pUB-S as template DNA. Figure 4 shows a primer extension product of 51 nucleotides indicating that the transcription initiation site of *ubi3R* is located at position −52 relative to the ATG start codon.

As in most fungal genes, no typical ‘TATAA’ or ‘CAAT’ boxes are present in the promoter region. Several CT-rich regions, commonly found in the vicinity of the initiation codons of fungal genes, are present upstream of the ATG start codon. Similarities matching the consensus sequence for heat shock promoter elements [4] which have been shown to be present in polyubiquitin genes from a number of organisms, were also found in the 5' non-coding region of the *ubi3R* gene (underlined in Fig. 2). In the 3' flanking region of the gene, a possible polyadenylation recognition sequence (AATTTAA) was found at position +797 to +804, 107 nucleotides downstream of the TAA stop codon (overlined in Fig. 2).

*The identified ubiquitin gene is a member of a multigen family*

Northern blot analysis of RNA isolated from the *in vitro* grown mycelium, using the two 228 bp

| *P. infestans* | M Q I F V K T L T G K T I T L D V E P S D G I D N V |
| * Yeast | E S T |
| * Barley | E S T |
| * Chicken | E T E |
| * Human | E T E |

| *P. infestans* | K Q K I Q D K E G I P P D Q Q R L I F A G K Q L E D |
| * Yeast | S |
| * Barley | A |
| * Chicken | A |
| * Human | A |

| *P. infestans* | G R T L S D Y N I Q K E S T L H L V L R L R G G N * |
| * Yeast | |
| * Barley | K * |
| * Chicken | Y * |
| * Human | C * |

*Fig. 3. Amino acid sequence comparison between the last ubiquitin encoding unit (77 amino acids) of *Phytophthora infestans* polyubiquitin and the last ubiquitin-encoding unit of polyubiquitin of yeast [35], barley [17], chicken [6] and human [2]. Only amino acid residues which differ from the *P. infestans* sequence are shown in the yeast, barley, chicken and human amino acid sequence.*
Fig. 4. Primer extension of 3 μg of poly(A)^+ RNA isolated from *in vitro* grown *Phytophthora infestans* mycelium (lane 1) and non-infected potato leaves (lane 2) using a primer (5’-GTGCCCTCGTITATGAG-3’) complementary to the *ubi3R* sequence at position -2 to -19.

*Pvu II* fragments from the coding region of *ubi3R* as probe, shows four bands of ca. 850, 1100, 1350 and 2350 nucleotides (Fig. 5A, lane 5). Longer exposure reveals a fifth band of circa 1850 nucleotides just above the plant ubiquitin mRNA. This suggests that the genome of *P. infestans* contains multiple copies of ubiquitin-encoding genes of different lengths. Using the gene-specific probe from the *ubi3R* promotor region, it was shown that the 850 nucleotide transcript corresponds to the *ubi3R* gene (Fig. 5B). Southern blot analysis of digested genomic DNA in which the two 228 bp *Pvu II* fragments from the *ubi3R* coding region were used as a probe (Fig. 6), confirmed that the identified ubiquitin gene belongs to a multigene family. In each digest analysed, multiple bands hybridize. Among those a 2.3 kb *Hind III* band, a 3.2 kb *Sat I* band and a 5.3 kb *Kpn I* band correspond to identical restriction fragments in the DNA of *Phaeohyphomycetes* (Fig. 1). The *Eco R1* and the *Bam HI* fragment on the Southern blot differ in size with those in the phage DHCl because only a part of these fragments is linked to vector DNA. Since the coding region of *ubi3R* does not have any internal *Kpn I*, *Eco R1*, *Bam HI*, *Hind III* and *Sat I* sites and no introns it is evident that the *P. infestans* genome contains multiple copies of ubiquitin-encoding genes.
Fig. 6. Southern blot analysis of Phytophthora infestans DNA (10 μg/lane) digested with the restriction enzymes indicated. The two 228 bp Pvu II fragments from the coding region of ubi3R were used as probe.

In planta induced expression of Phytophthora infestans ubiquitin genes

The differential expression of the characterized ubiquitin gene was confirmed in two ways. In the first procedure interaction RNA, i.e. RNA isolated from inoculated leaves, and RNA isolated from mycelium grown in vitro was subjected to northern blot analysis (Fig. 5). Equal amounts of interaction RNA and fungal RNA were applied to the gel. The hybridization signals of the ubiquitin probe with interaction RNA (Fig. 5A, lane 4, 3 days after inoculation) are 2–3 fold stronger in comparison with the signals obtained with RNA isolated from the fungus grown in vitro (Fig. 5A, lane 5) indicating that the relative amounts of all fungal ubiquitin transcripts is higher in interaction RNA. The 1800 nt transcript present in lanes 1–4 of Fig. 5A is a potato ubiquitin messenger. Due to the highly conserved sequence of ubiquitin, the potato ubiquitin transcript is cross-hybridizing with the fungal ubiquitin probe. To follow the progression of fungal growth during the development of the disease, the proportion of fungal RNA in the total RNA population isolated from infected leaves was determined. To this end a probe of the constitutively expressed actin (act4) gene of P. infestans [46] was used for hybridization of the northern blot (Fig. 5C). One day after inoculation the proportion of fungal RNA in the interaction RNA is very low. Due to the increasing amount of fungal biomass during colonization of the leaf tissue, the proportion of fungal RNA in the interaction RNA mixture increases rapidly to ca. 50%, three days after inoculation. Taking this into consideration, the results of the northern blot analyses show that the expression of the ubiquitin encoding genes during growth of the fungus in planta increases 4–6-fold. Using a ubi3R gene-specific probe, the induction of expression of the ubi3R gene seems to be even higher (Fig. 5B).

In the second procedure the difference in relative abundance of P. infestans ubiquitin mRNAs in planta and in vitro was analysed by comparative hybridization of four replica filters of the P. infestans genomic library with (1) the random primer labelled 228 bp Pvu II fragments from the ubi3R coding region (Fig. 7-U), (2) labelled interaction cDNA derived from poly(A) mRNA of the interaction, 3 days after inoculation (Fig. 7-I), (3) labelled fungal cDNA made from poly(A) mRNA of the fungus grown in vitro (Fig. 7-F) and (4) labelled cDNA derived from poly(A) mRNA of uninfected potato leaves (Fig. 7-P) as a control. Lambda clones containing ubiquitin encoding sequences (indicated by arrows in Fig. 7) gave rise to a relatively strong signal when hybridized with labelled interaction cDNA (Fig. 7-I). Hybridization of a control replica filter with labelled cDNA derived from poly(A) mRNA of uninfected potato leaves (Fig. 7-P) shows very weak or no signals indicating that the signals in Fig. 7-I are predominantly due to hybridization with fungal cDNA. Since the intensity of the signal is positively correlated with the abundance of the corresponding mRNAs, the ubiquitin mRNAs can be classified as highly abundant in the P. infestans mRNA population of the interaction. Hybridization of a replica filter with labelled fungal cDNA (Fig. 7-F) shows a relatively weak signal of the same clones
indicating that ubiquitin mRNAs are several-fold less abundant in the mRNA population of in vitro grown mycelium than in the mRNA population of in planta produced mycelium. This procedure even underestimates the difference in abundance of the ubiquitin mRNAs because the filters were hybridized with equal amounts of labelled cDNA whereas the proportion of fungal cDNA in the interaction cDNA probe is approximately half of that in the fungal cDNA probe.

**Discussion**

Differential hybridization of cDNA libraries has been proved to be a useful tool for the isolation of genes which are differentially expressed under distinct physiological conditions, in particular for the isolation of developmentally regulated genes [43, 44]. Differential hybridization of genomic libraries is not often utilized since large vector inserts and low specificity of the labelled cDNA probes make it difficult to map and isolate the genes of interest. Isolation of in planta induced genes from P. infestans can only be done by differential screening of a genomic library since the use of a cDNA library made from interaction poly(A)+ RNA will yield both differentially expressed plant and fungal genes which cannot directly be distinguished. In developing the procedure using a genomic library it was of utmost importance to use cDNA probes with a high specific activity (> 10^8 cpm/µg cDNA) to be able to isolate putative in planta induced genes from the P. infestans genomic library.

Using the procedure described above a differentially expressed P. infestans polyubiquitin was isolated from a genomic library. The structure of the selected ubi3R gene is completely consistent with the unique structure of known polyubiquitin genes from other organisms [23, 30]. The characterized gene consists of three repeats each coding for the 76 amino acid ubiquitin peptide. Southern and northern blot analyses showed that the characterized polyubiquitin gene is a member of a multigene family. The ubi3R gene codes for
the 850 nt transcript as shown in Fig. 5B. The sizes of the other transcripts (1100, 1350, 1850 and 2350 nt) indicate that they may code for polyubiquitin genes containing up to 4, 5, 7 and 9 ubiquitin units which is a common feature for polyubiquitin gene families.

Ubiquitin is one of the most conserved proteins known to date (for reviews see [13, 22, 23, 30, 36]). The 76 amino acid protein occurs in all eukaryotic cells, either free or covalently attached to proteins in the cytosol, plasma membrane or to chromosomal histones. Ubiquitin has been shown to play a key role in several important cellular processes such as the selective degradation of intracellular proteins [22, 23, 36], maintenance of chromatin structure [29, 31], regulation of gene expression [3, 18] and modification of cell surface receptors [26, 40, 50]. Genes encoding ubiquitin have been characterized for a variety of organisms such as yeast [35], man [47], chicken [6], Xenopus laevis [11], Drosophila melanogaster [1], Tripanosoma cruzi [42] and Caenorhabditis elegans [19]. In each case, ubiquitin is encoded by one or more polyubiquitin genes which consist of direct repeats of the 76 amino acid coding units. The least repeat at the 3′ end of the polyubiquitin gene is usually followed by an extra amino acid residue which is not conserved among different species. In the P. infestans ubi3R gene this appears to be as an asparagine. Although the unique structure of polyubiquitin genes has been conserved in evolution, considerable variation exists in the number of repeats within each polyubiquitin gene and the number of polyubiquitin encoding loci in the genome.

Recently another class of genes which contain the ubiquitin-coding sequence has been identified. In these genes the ubiquitin coding region is fused in frame to the 5′ end of a coding sequence of an unrelated polypeptide [33, 35, 37]. Their products are called ubiquitin carboxyl extension proteins (UbcCEPs) and have been found in several eukaryotic organisms. The carboxyl extension proteins (CEPs) are reported to be ribosomal proteins whose fusion to ubiquitin is shown to facilitate ribosomal biogenesis [12]. Like the ubiquitin-encoding sequence, the CEP-encoding sequence of these natural gene fusions is conserved among species.

The expression of both classes of ubiquitin genes is found to be differentially regulated under conditions of stress, rapid cell growth or during development [17, 34]. Induced expression of polyubiquitin genes by heat shock or other types of stress has been observed in a number of organisms [7, 32, 35]. This facilitates an increased production of ubiquitin monomers for the ubiquitin-mediated degradation of abnormal proteins which arise during stress. The expression of UbCEP-encoding genes has been demonstrated to be induced under conditions of rapid cell growth and cell division [30, 35, 42]. As protein synthesis is a major cellular process during cell growth and CEPs are identified as ribosomal proteins, the significance of induced expression of UbCEP encoding genes is obvious.

The induced expression of ubiquitin-encoding genes in P. infestans during colonization of potato leaves may reflect the highly active metabolic state of the mycelium in the host tissue. During exponential growth in vitro on synthetic Henniger medium and on rich rye-sucrose agar medium, however, the mycelium is also in a highly active metabolic state but the ubiquitin-encoding genes show a 4-6-fold lower level of expression. This implies that the host environment specifically induces the expression of the ubiquitin-encoding genes. Which factors or conditions are due to the in planta induced expression of the P. infestans ubiquitin genes needs to be investigated. Also the significance of induced expression of the ubiquitin-encoding genes during pathogenesis remains to be resolved. In view of the regulatory function of ubiquitin in gene expression [3, 18] it is tempting to speculate that induction of the characterized ubi3R gene may have an effect on the regulation of the expression of genes involved in pathogenicity. The other genes containing the ubiquitin-encoding sequence may code for polyubiquitin as well and may have a similar function, but it cannot be excluded that they encode UbCEPs and are involved in biogenesis of ribosomes.

Expression studies should reveal whether the
ubiquitin genes are inducible in vitro by heat shock and stress or by particular plant factors in order to gain more insight into the significance of the induced expression of the P. infestans ubiquitin genes in planta. The ubi3R gene is one of the first genes isolated from P. infestans and is highly expressed in vitro. The promoter region of the gene will be used for the construction of vectors for transformation of P. infestans, a necessary tool for the accomplishment of our goal, the isolation and characterization of pathogenicity genes of P. infestans.

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