A novel root-specific gene, MIC-3, with increased expression in nematode-resistant cotton (Gossypium hirsutum L.) after root-knot nematode infection

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Abstract

A full-length cDNA, MIC-3, has been identified from a \textsuperscript{NZAPII} cDNA library constructed from the mRNA of nematode-resistant cotton (Gossypium hirsutum L.) roots after infection with root-knot nematode (Meloidogyne incognita). The putative open reading frame of MIC-3 encoded a protein of 141 amino acids with a calculated molecular mass of 15.3 kDa. Seven alternative polyadenylation sites have been identified for the MIC-3 transcripts, and the major transcripts are the longest ones. The MIC-3 gene contains a single intron within its coding region and belongs to a novel, multi-gene family containing up to six members. Expression of MIC-3 is root localized and specifically enhanced in the nematode induced, immature galls of resistant cotton line M-249, suggesting that MIC-3 may play a critical role in the resistance response to root-knot nematode. Published by Elsevier Science B.V.

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Root-knot nematodes (RKN) (Meloidogyne spp.) are obligate sedentary endoparasites and cause severe losses in many crops worldwide [1]. The second-stage nematode juveniles (J2) penetrate the root, migrate to the vascular tissue, and induce the formation of multinucleate "giant cells" as permanent feeding sites [2]. During giant cell formation, nearby cells expand and divide forming enlarged galls or root-knots. In resistant tomato plants, J2 stage RKN penetrate roots as in susceptible plants, but a localized cell death, involving a hypersensitive response, inhibits giant cell development at the site where feeding is initiated [3]. Some cotton (Gossypium hirsutum L.) lines exhibit high levels of RKN resistance [4–6], which is conferred by at least two unidentified genes [7–10]. The size and index of newly formed galls were not significantly different between the resistant and susceptible lines during early stages of infection (6–10 days after inoculation, DAI); however, RKN reproduction was inhibited (≥98%) in resistant plants by the end of the normal reproductive cycle for RKN (60–80 DAI) [10]. In contrast to susceptible lines, the further development of J2 stage juveniles is arrested in the resistant cotton lines by 8–10 DAI, but little is known about the alterations in plant gene expression associated with this resistance response [10]. The genes with increased expression specifically in the resistant plant roots after nematode infection are likely to be critical for the establishment of resistance mediated by the nematode resistance gene(s), whereas genes with elevated expression in both resistant and susceptible roots might be involved in general defense mechanisms [11].

Our previous studies identified a 14-kDa protein, which was induced in the immature root galls of a RKN-resistant cotton isolate soon after inoculation (8–10 DAI) [12]. Based on limited internal amino acid sequence (MKVDGYGY) of the 14-kDa protein [12], a degenerate reverse primer,
R1 (5′-CCRTARTANCCRTCYTTCAT-3′, nucleotide (nt) positions 169–191 in Fig. 1) (Y = C/T, R = A/G, N = A + C + G + T), was designed to amplify the cDNA sequence encoding the 14-kDa protein. A cotton AZAPII cDNA library was constructed from mRNA of excised immature galls of resistant cotton (M-249) roots at 10 DAI with RKN [Meloidogyne incognita (Kofoid and White) Chitwood, race 3]. Using the gene-specific primer R1 and AZAPII vector primer T3, a DNA fragment was amplified by polymerase chain reaction (PCR) from the cDNA library stock culture. This fragment was cloned into pGEM-T Easy vector (Promega) and sequenced with ABI PRISM 310 DNA Genetic Analyzer (Perkin-Elmer). Based on the identified 5′ end cDNA sequence, a forward primer, F1 (5′-TTAAGATGGCTTGTGCTCCACTAATCAGATTCCA-3′, nt positions 16–47 in Fig. 1) was designed and then used together with the vector primer T7 to amplify the 3′ end cDNA sequence by PCR. The amplified 3′ cDNA fragment was cloned and then sequenced as described. Another reverse primer, R2 (5′-AATTGCAACCGCTCACATGGATTTCT-3′, nt positions 493–522 in Fig. 1), was designed based on the identified 3′ end cDNA sequences and paired with T3 primer for PCR amplification to further confirm the 5′ end cDNA sequence. The two primers, F1 and R2, were used to amplify the genomic DNA sequences from the resistant line. In all PCR reactions, Pfu DNA polymerase (Stratagene) possessing the 3′ to 5′ proofreading exonuclease activity was used for accurate sequence determination.

The final assembled nucleotide sequence for MIC-3 (Meloidogyne induced cotton-gene) has an open reading frame of 423 bp with the start codon at nt position 22, which is in the context of AAAATGG that fulfills the criterion of a Kozak sequence (A/GNNATGG) [13] (Fig. 1). The MIC-3 gene encodes a putative protein of 141 amino acids with an estimated molecular weight of 15,357 and isoelectric point of 5.5. A single intron of 77 bp contains the conserved GT and AG sequences as the 5′ and 3′ splicing sites, respectively. Seven MIC-3 transcripts, differing only in length at the 3′ regions, and a poly(A) sequence of at least 20 nt were amplified from the cDNA library by primers F1 and T7. The alternative polyadenylation sites are located at nt positions 679, 681, 713, 715, 716, 718, and 803 (Fig. 1). The predominant transcripts were the longest ones present in more than half of the amplified cDNAs. Three putative polyadenylation signals (AAUUAUA, AUUAAA, AAUAAA) [14] are at nt positions 651, 688, and 782. The canonical signal AAUAAA, which is processed more efficiently than the variant signals AAUUAUA and AUUAAA, is generally located at the 3′-most terminal site, and the predominant form of alternatively polyadenylated transcripts is normally the longest one [14]. The two alternative polyadenylation

![Fig. 1. Nucleotide and derived amino acid sequences of the cotton MIC-3 gene. The sequence of the 77 bp intron is shown in lowercase letters, and the conserved intron splicing sequences GT and AG are underlined. The stop codon is indicated by an asterisk. The three putative polyadenylation signals are double-underlined and the seven alternative polyadenylation sites are indicated by open arrowheads.](image-url)
sites at nt positions 679 and 681 are 28 and 30 nt downstream of the AAUAUA signal, and the four alternative polyadenylation sites at nt positions 713, 715, 716, and 718 are 25, 27, 28, and 30 nt downstream of the AUUAAA signal. The major polyadenylation site at nt position 803 is 21 nt downstream of the AAUAAA signal. The percentage of genes whose mRNA transcripts use two or more polyadenylation cleavage sites downstream from a single polyadenylation signal is considerably high in mammals, with 44% in human, 22% in mouse, and 22% in rat [27]. Alternative polyadenylation is frequently associated with tissue- or disease-specific expression [28,29]. Interestingly, the MIC-3 gene and predicted protein showed no significant homology to known sequences in any of the available databases.

Genomic Southern blot analysis using $^{32}$P-labeled MIC-3 cDNA probe revealed at least six hybridizing bands dependent on the restriction enzyme employed, suggesting that MIC-3 gene belonged to a multi-gene family (Fig. 2). Northern analysis was used to determine the expression pattern of the MIC-3 gene. Seedlings of RKN-resistant (M-249) and recurrent susceptible parent (ST213) were inoculated with J2 RKN [9]. The small immature galls at 10 DAI were excised from roots of both resistant and susceptible plants [12]. Equivalent sections of roots were also excised from non-inoculated controls under the same conditions.

Fig. 2. Genomic Southern blot analysis of MIC-3 gene family in cotton. Genomic DNA (10 µg) extracted from young leaves of M-249 (RKN-resistant isolate) was digested with restriction enzymes EcoRI, EcoRV, and HindII, respectively, separated by agarose gel electrophoresis, blotted, and hybridized with $^{32}$P-labeled MIC-3 cDNA probe. The membrane was washed under high-stringency (twice for 10 min at room temperature in 1 x SSC plus 0.1% SDS, once for 50 min at 65 °C in 1 x SSC plus 0.1% SDS, and once for 20 min at 65 °C in 0.2 x SSC plus 0.1% SDS) and visualized by autoradiography at –80 °C in the presence of an intensifying screen.

Aliquots of 10 µg total RNA isolated from the root samples, 15 DPA (days post-anthesis) fibers, yellow flowers, and young leaves were fractionated by gel electrophoresis, blotted, hybridized with a $^{32}$P-labeled MIC-3 cDNA probe, and washed under high stringency conditions. A 0.75-kb transcript was detected in the root samples but not in fibers, flowers, or leaves; the transcript level of MIC-3 was at least six times higher in the immature root galls of the resistant plants (Fig. 3). The estimated size (0.75 kb) of the MIC-3 transcript was very close to the length of its longest cDNA sequence of 747 bp (including the 21 nt poly-A sequence), indicating that the cDNA sequence reported was full-length. Rehybridization of the same blot with a control probe GhUBC2 (cotton class I ubiquitin-conjugating enzyme gene) known to be expressed in all cotton tissues [15].
the observed root-specific and induced expression of the MIC-3 gene in the resistant inoculated plants (Fig. 3).

The expression pattern for the MIC-3 gene, size of the predicted protein, and calculated pI were consistent with our previous data on expression of the 14-kDa protein [12]. These results suggest that the MIC-3 gene may encode the 14-kDa protein. The deduced MIC-3 amino acid sequence includes (MMVDGLYG) (50–57) (Fig. 1) homologous to the short internal sequence (MKVDGGYYG) obtained from the gel-purified 14 kDa protein [12]. We suspect that the two amino acid differences reflect less than homogeneous purification of the internal fragment of the 14 kDa, leading to some inaccuracy in the protein sequencing. The MIC-3 protein has been successfully expressed in E. coli at the expected size (data not shown) and will be used as antigen to develop antiserum to the protein. Western blot analysis of the cotton 14 kDa protein using the MIC-3 antiserum will further verify their relationship.

To our knowledge, this is the first report of a root-specific gene with increased expression specifically in nematode-resistant plants following nematode infection. Eight cDNAs have been identified with increased expression in both resistant and susceptible tomato after RKN inoculation, suggesting that they are likely to play a role in the more general, root-defense system [11]. Many genes which are up-regulated in nematode-induced feeding cells have been isolated [16–25], and their elevated expression is likely to be important for the establishment and maintenance of the nematode-induced feeding structures or for the common defense response in susceptible plants [11,16]. Tomato RKN resistance is conferred by a single, dominant gene, Mi-1.2, the only cloned RKN resistance gene in plants [3]. The signal transduction pathway leading to localized cell death may be initiated by the interaction between Mi-1.2 and a plant protein induced by the nematode infection [26]. As a gene whose expression is specifically enhanced in RKN-induced, immature galls of resistant cotton line M-249, MIC-3 is likely involved in the resistance mechanism mediated by the cotton RKN resistance genes. Characterization of the MIC gene family members and their expression in other cotton backgrounds with the same source of resistance genes will be needed to further substantiate a role in the resistance mechanism. The over-expression of MIC-3 gene in RKN susceptible cotton roots and the reduction or elimination of MIC-3 product by antisense mRNA expression in resistant plants will more directly test the hypothesis.

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References


