# Expression and Regulation of the Arabidopsis thaliana Cel1 Endo 1,4 $\boldsymbol{\beta}$ Glucanase Gene During Compatible Plant-Nematode Interactions 

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#### Abstract

The root-knot nematode Meloidogyne incognita is an obligate endoparasite of plant roots and stimulates elaborate modifications of selected root vascular cells to form giant cells for feeding. An Arabidopsis thaliana endoglucanase (Atcel1) promoter is activated in giant cells that were formed in Atcell:: UidA transgenic tobacco and Arabidopsis plants. Activity of the full-length Atcel1 promoter was detected in root and shoot elongation zones and in the lateral root primordia. Different 5' and internal deletions of regions of the $1,673 \mathrm{bp}$ Atcel1 promoter were each fused to the UidA reporter gene and transformed in tobacco, and roots of the transformants were inoculated with M. incognita to assay for GUS expression in giant cells and noninfected plant tissues. Comparison of the Atcel1 promoter deletion constructs showed that the region between $-1,673$ and $-1,171$ (fragment 1 ) was essential for Atcel1 promoter activity in giant cells and roots. Fragment 1 alone, however, was not sufficient for Atcell expression in giant cells or roots, suggesting that cis-acting elements in fragment 1 may function in consort with other elements within the Atcell promoter. Root-knot nematodes and giant cells developed normally within roots of Arabidopsis that expressed a functional antisense construct to Atcell, suggesting that a functional redundancy in endoglucanase activity may represent another level of regulatory control of cell wall-modifying activity within nematode feeding cells.


Key words: cellulase, cis-acting elements, giant cells, Nicotiana tabacum, parasitism, regulatory motif.

The root-knot nematode, Meloidogyne incognita, is an obligate endoparasite of plant roots that has evolved a complex feeding relationship with its host (reviewed in Davis et al., 2004). The structure of giant cells induced by root-knot nematodes includes extensive cell wall modifications acting as specialized feeding sites to allow cellular expansion and solute uptake (Jones and Northcot, 1972; Hussey and Grundler, 1998; Goellner et al., 2001; Gheysen and Fenoll, 2002; Vercauteren et al., 2002). The complex morphological and physiological changes during the establishment of giant cells and other nematode feeding sites (NFS) are reflected by altered gene expression in affected root cells (Wilson et al., 1994; Gheysen et al., 1996; Williamson and Hussey, 1996; Favery et al., 1998). The coordinated temporal expression and localization of cell wall-modifying enzymes that promote wall loosening and expansion in giant cells, and concomitantly promote cell-wall thickening and extensive ingrowths at the interface of neighboring vascular elements, likely represent augmentation of regulatory machinery active during normal plant cell wall growth and maturation.

The plant cell wall is a network of cellulose microfibrils, hemicellulose, pectin, and proteins that undergoes extensive changes in architecture during plant

[^0]growth and development (reviewed in Carpita and Gibeaut, 1993). During the growth process, plant cells respond to multiple internal and external signals. In many cases, the response is translated into the loosening of the cell wall to enable turgor-driven cell expansion (Crosgrove, 1999; Rose and Bennett, 1999). Targeted enzymatic digestion of cellulose microfibrils by endogenous plant endoglucanases (EGases) is a tightly regulated process that is one primary component of cell-wall loosening (Fry, 1995; Rose and Bennett, 1999). The plant EGase genes identified to date are usually expressed within different developmental stages of the plant such as elongation, ripening and abscission (Lashbrook et al., 1994; Shani et al., 1997; del Campillo, 1999; Levy et al., 2002).

The Arabidopsis thaliana EGase gene, Cel1 (Atcel1), is essential for normal plant cell growth and elongation, as it plays a role in cell wall deposition, cell differentiation, and cytogenesis (Tsabary et al., 2003). The primary activity of Atcell has been observed during cell elongation and within fast growing tissues (Shani et al., 1997; Nicol et al., 1998; del Campillo, 1999; Shani et al., 2000). Overexpression of Atcell resulted in accelerated growth of transgenic tobacco and poplar plants (Levy et al., 2002; Shani et al., 2004). Conversely, A. thaliana plants expressing Atcel1 antisense exhibit shorter stems and roots, a corrugated cell wall surface, and fewer $x y-$ lem elements per bundle, and both xylem elements and the interfascicular fibers are significantly less lignified than in the wild type (Tsabary et al., 2003).

Recently published evidence shows that elevated plant EGase activity localized in NFS may be, in part, responsible for the dramatic cell wall modifications observed within NFS (Goellner et al., 2001; Mitchum et al., 2004). Increased expression of five Nicotiana tabacum EGase (Ntcel) genes was detected within NFS induced in tobacco roots by both root-knot and cyst nematodes, and differential expression levels of each of
the upregulated Ntcel genes in NFS were observed by semi-quantitative PCR (Goellner et al., 2001). The fulllength promoter of Atcel1 fused to GUS (Shani et al., 1997) was upregulated in giant cells induced by rootknot nematodes, but not within syncytia induced by cyst nematodes in either $N$. tabacum or A. thaliana host plants (Mitchum et al., 2004). The data suggest that differential expression of plant EGases gives rise to different NFS, but it is not clear how this activity may be regulated or which plant EGases are essential for proper formation of a given NFS. To this end, the activity of different deletions of the Atcell promoter upon plant infection by root-knot nematodes and the response of plants expressing antisense to the Atcell gene (Tsabary et al., 2003) to nematode infection have been investigated.

## Materials and Methods

Atcel1 constructs and transgenic plants: Transgenic tobacco (N. tabacum SR1) and A. thaliana (Col-0 Ecotype) plants containing the full-length Atcel1 promoter fused to the $\beta$-glucuronidase (GUS) reporter gene were developed previously (Shani et al., 1997; Tsabary, 2003). The construct was a transcriptional fusion between 1.6 kilobases (kb) of the putative Atcell promoter region (bases 5-1,618; Genbank accession X98543) and the 5' end of the $\beta$-glucuronidase gene (UidA) (Jefferson, 1987; Shani et al., 1997). The Atcel1 promoter was divided into four 382 to 468 bp fragments (Fig. 1A), and six promoter::GUS constructs containing one to three fragments each were developed.

To facilitate subcloning, each promoter segment was amplified using primers that contain restriction sites for the enzymes Hind III, Nde I and Sal I. The primers used were as follows: Fragment 1) 5'-AAAAAAGCTT-ACCTGCAGGTCAACGG-3' and 5'-AAAACATATGT-TCATTTAGTATATAACAAAATTCG-3'; Fragment 2) 5'-ATTTAAGCTTACACCATATGAAATGAACATTTG-CTCTGATTTGG-3 and 5'-AAAACATATGATTAT-TATATACTTTTTTTTTTATAAAAG-3'; Fragment 3) 5'-AAAAAAGCTTAAAACATATGTATATAATAATTT-ACACTCGAATC-3' and 5'- TGTGCATATGCTCAAT-AGTTGATTTTTGGAGG-3'; Fragment 4) 5'-AAAAAG-TTAAATCATATGGAGATCAAAACACGTGTCGC-3' and 5'-CCCCGTCGACGTCTCTTCTTTCTTGTGC-3'. The PCR reactions were performed using thermal cycling conditions of $94^{\circ} \mathrm{C}$ for $4 \mathrm{~min}, 30$ cycles of $94^{\circ} \mathrm{C}$ for $10 \mathrm{sec}, 55^{\circ} \mathrm{C}$ for 10 sec , and $72^{\circ} \mathrm{C}$ for 10 sec , and $72^{\circ} \mathrm{C}$ for 4 min using a buffer containing 1 Unit of DeepVent Taq polymerase (New England Biolabs, Inc., Beverly, MA), 20 mM Tris- $\mathrm{HCl} \mathrm{pH} 8.8,10 \mathrm{mM} \mathrm{KCl}, 10 \mathrm{mM}$ $\left(\mathrm{NH}_{4}\right)_{2} \mathrm{SO}_{4}, 2 \mathrm{mM} \mathrm{MgSO} 4,0.1 \%$ Triton X-100, $200 \mu \mathrm{M}$ dNTPs, 10 pmol primer and 1 ng DNA template. The plasmid pUC18, containing a Sal I/Eco RI genomic fragment harboring the Atcell promoter (Shani et al., 1997), was used as the template.


Fig. 1. Schematic representation of Atcel1 promoter-UidA constructs in transformed tobacco plants analyzed for tissue expression and response to nematode infection. A) The full-length $1,673 \mathrm{bp}$ Atcel1 promoter, B-F) 5' Atcel1 promoter deletion constructs harboring different lengths of the promoter (serial B and C , internal $\mathrm{D}, \mathrm{E}$, and F). Numbers indicate the length in bp of the respective promoter regions. Coding region of the $\beta$-glucuronidase gene. TSP transcription starting point, 1 to 4 indicate respective excised promoter regions.

The fragments were purified, digested with different digestion enzymes (Fermentas Inc., Hanover, MD) and cloned, in various combinations, into pUC18 to create the promoter segment combinations shown in Figure 1. The promoter constructs were then digested with Hind III and Sal I and cloned into the binary vector pBinPlus, which contained the UidA.

The vectors were introduced into Agrobacterium tumefaciens using the freeze-thaw method (Erbert et al., 1988) and then transformed into tobacco. The constructs were then transformed into N. tabaccum-SR1 plants using the leaf-disc transformation as described previously (Horsch et al., 1985). Kanamycin-resistant plants were regenerated and confirmed by PCR. T2 homozygote plants were selected for further analysis.

Cel1 antisense Arabidopsis plants were previously developed and characterized by Tsabary et al. (2003). The construct contains bases 1 to 403 bp of the Atcel1 coding region inserted in reverse orientation into the vector pBI101.1 containing the CaMV 35S RNA promoter and the octopine polyadenylation site (Tsabary et al., 2003).

Seeds of transgenic Cel1-UidA tobacco and Arabidopsis, Cel1 antisense Arabidopsis, and nontransformed seeds were surface disinfected with $2.5 \% \mathrm{NaOCl}$ and
$0.5 \%$ sodium dodecyl sulfate (SDS) for 10 min , rinsed four times with sterile water, and then germinated and grown monoxenically in petri plates containing $0.8 \%$ Noble agar (Fisher Scientific, Pittsburgh, PA) supplemented with MS basal medium (Murashige and Skoog, 1962), pH 5.8, sucrose ( $30 \mathrm{~g} /$ liter), and kanamycin (50 $\mathrm{ug} / \mathrm{ml}$ ). Tobacco and Arabidopsis seedlings were grown in a controlled temperature growth chamber at $25^{\circ} \mathrm{C}$ with a 16 -hr photoperiod. At least five independent kanamycin-resistant lines were analyzed for each transgenic construct.

Nematode Infection: The root-knot nematode, M. incognita, was propagated on roots of greenhouse-grown tomato (Lycopersicon esculentum cv. Rutgers). Meloidogyne incognita eggs were isolated from egg masses on tomato roots with $0.5 \% \mathrm{NaOCl}$ (Hussey and Barker, 1973), surface disinfected in a solution of $0.02 \%$ sodium azide for 30 min , rinsed with water on a $25-\mu \mathrm{m}$-opening sieve, and hatched in water at $28^{\circ} \mathrm{C}$ on a Baermann pan (Mitchum et al., 2004). Hatched M. incognita J2 were surface sterilized in $0.002 \% \mathrm{HgCl}_{2}, 0.002 \% \mathrm{NaN}_{3}$, and $0.001 \%$ triton X-100 for 10 min , followed by five washes with sterile water. Surface-sterilized J2 were resuspended in $50 \mu \mathrm{l}$ of 2 mM penicillin-G and $950 \mu \mathrm{l}$ of $0.1 \%$ water agar immediately prior to inoculation of roots of plants grown on sterile nutrient agar. Five-microliter aliquots of $M$. incognita J2 were used to inoculate 10 - to 12 -dayold tobacco root tips and 10-day-old Arabidopsis root tips grown in monoxenic culture at a concentration of 15 $\mathrm{J} 2 / \mu \mathrm{l}$ and $100 \mathrm{~J} 2 / 5 \mu \mathrm{l}$, respectively. Penetration of roots by J2 was monitored using an inverted light microscope. Infected and noninfected transgenic root tissues were excised from petri dishes at specific time points after penetration of roots by J2. For all the time points examined, at least 100 infected and 30 uninfected transgenic roots were assayed for nematode infection and GUS expression. Promoter activity was also monitored in the root elongation zone and giant cells of control plants harboring the $\Delta 0.6$ TobRB7 promoterUidA construct (Yamamoto et al., 1991) treated similarly as a positive control.

Histochemical GUS analysis: $\beta$-glucuronidase (GUS) activity was monitored by the method of Jefferson (1987) with some modifications (Yamaguchi et al., 2001). Fresh, excised root pieces were vacuum-infiltrated for 5 min with GUS substrate ( 2 mM 5 -bromo-4-chloro-3indolyl $\beta$-D-glucuronide [X-Gluc], 100 mM Tris, pH $7.0,50 \mathrm{mM} \mathrm{NaCl}, 0.06 \%$ Triton X-100) and incubated 12 hr at $37^{\circ} \mathrm{C}$. In leaf samples excised to confirm construct (GUS) expression in experimental lines, chlorophyll pigmentation was removed by incubation of the samples for approximately 1 hr in $90 \%$ (v/v) ethanol. Samples stained for GUS activity were placed in $70 \%$ ethanol for long-term storage at $4^{\circ} \mathrm{C}$.

Tobacco and Arabidopsis plants harboring the Atcel: GUS construct were analyzed for GUS expression following infection of host roots by M. incognita at 3, 4, 7,

14,21 and 28 dpi in five to seven independent transformed lines for which 10 to 15 seedlings were assayed.

Histology of nematode feeding cells: Prior to sectioning, stained root pieces were fixed at $4^{\circ} \mathrm{C}$ in $4 \%$ paraformaldehyde for 16 hr for Arabidopsis and 3 d for tobacco, washed twice in PBS for 15 min after fixation, dehydrated in a graded ethanol series ( 30 min each), incubated sequentially in Histoclear (National Diagnosis, Atlanta, GA):ethanol 25:75, 50:50, 75:25, and then in $100 \%$ Histoclear twice for 30 min each time (Goellner et al., 2001). The root pieces were incubated in Histoclear:Paraplast Plus (Fisher Scientific) 75:25 overnight at $60^{\circ} \mathrm{C}$ and then overnight in pure Paraplast at $60^{\circ} \mathrm{C}$. The Paraplast-embedded root pieces were sectioned to a thickness of $30 \mu \mathrm{~m}$ for tobacco and $10 \mu \mathrm{~m}$ for Arabidopsis using an a rotary microtome (American Optical, Buffalo, NY) and adhered to Superfrost Plus microscope slides (Fisher Scientific) overnight at $40^{\circ} \mathrm{C}$ on a slide warmer. Three 15 min incubations in Histoclear were used to remove the Paraplast from sections adhered to slides, followed by rehydration in a graded ethanol series to water prior to mounting with Permount (Fisher Scientific). For each time point, 15 to 30 infected roots were analyzed for GUS staining, and an equal number of uninoculated roots were analyzed for comparison.

Computational analyses of the Atcel1 promoter sequence: For putative motif analysis of the Atcell promoter, we utilized the results of the Plant-CARE (Lescot et al., 2002) (http://oberon.fvms.ugent.be:8080/ PlantCARE/index.html) and PLACE (Higo et al., 1999) (http://www.dna.affrc.go.jp/htdocs/PLACE/ signalscan.html) algorithms as described by Rombauts et al. (2003). To detect novel common regulatory elements in multiple promoters, the MOTIF SAMPLER algorithm (Thijs et al., 2001) was used (http://www .esat.kuleuven.ac.be/~thijs/Work/MotifSampler .html). Consensus motifs identified using MOTIF SAMPLER were subsequently compared with the regulatory sites described in the Plant-CARE and PLACE databases.

Cel1 antisense A. thaliana: The development of M. incognita females in roots of A. thaliana expressing the antisense Atcell construct (Tsabary et al., 2003) were compared to M. incognita development in roots of wildtype A. thaliana. Gross shoot and root morphology was compared to published descriptions to confirm the reported Atcel1 phenotype (Tsabary et al., 2003) in test plants. The cellular morphology of infection sites in nematode-infected antisense roots as compared to wildtype was also evaluated using the fixation, embedding, and sectioning procedures described above. Sections were stained using Johansen's safranin/fast green protocol (Johansen, 1940) with some modifications (Ruzin, 1999) to enhance observable differences among cells and tissues. Photomicrographs of specimens were taken using a Nikon eclipse E600 microscope (Nikon

Instruments, Melville, NY) equipped with RT-color SPOT camera (Diagnostic Instruments, Sterling Heights, MI).

## Results

Impact of Atcel1 promoter deletions on tissue-specific and nematode-induced expression: Tobacco plants transformed with the full-length or a truncated Atcell promoter-UidA constructs were analyzed for promoter activity in uninfected plant tissue and within giant cells (Table 1). Five to seven independent transformed lines and at least 10 plants per transformed line were tested for each promoter construct. At least 100 tissue samples, including samples from each independent transformed line, were assayed for each promoter construct. Histochemical assays of $\beta$-glucuronidase (GUS) expression were used to analyze the temporal and spatial characteristics of the Atcell promoter activity. During plant development, the expression of the full-length Atcell promoter-UidA construct was observed in shoot and root elongation zones of infected and uninfected plants (Construct A, Figs.1,2A-B, Table 1). The expression of the full-length Atcell promoter-UidA construct was not induced by mechanical wounding of the roots or leaves (data not shown). Low expression of construct B (Figs. 1,2C-D, Table 1), harboring a $502 \mathrm{bp} 5^{\prime}$ deletion (promoter fragment 1), was observed in tobacco shoot elongation zones, but not in the roots. There was no obvious variation in GUS expression patterns among the transformed lines containing the same promoter construct, although slight variations in expression intensity could be observed. GUS activity was never observed in plants with constructs C to F, even though constructs D to F contained promoter fragment 1 (Fig. 1, Table 1). GUS activity was observed in the root elongation zone and giant cells of control plants harboring the $\Delta 0.6$ TobRB7 promoter-UidA construct (Opperman et al., 1994), treated similarly as a positive control (data not shown).

We previously demonstrated that the full-length At-

Table 1. Activity of the Atcel1 promoter-UidA constructs in roots, shoot, leaves and root-knot nematode feeding sites (NFS).

|  | GUS Expression $^{\mathrm{b}}$ |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| Construct $^{\mathrm{a}}$ | Root | Shoot/leaves | NFS | No. of GUS + NFS/ <br> total no. NFS |
| A | + | + | + | $175 / 205$ |
| B | - | + | - | $0 / 313$ |
| C | - | - | - | $0 / 258$ |
| D | - | - | - | $0 / 136$ |
| E | - | - | - | $0 / 147$ |
| F | - | - | - | $0 / 100$ |

[^1]

Fig. 2. Histochemical staining for GUS activity in transgenic tobacco plants containing the Atcell promoter infected by the root-knot nematode, Meloidogyne incognita. A) Atcel1-driven GUS expression in a 7-day-old uninfected tobacco root. B) Atcel1-driven GUS expression in a shoot tip of a young uninfected tobacco seedling. C) Atcel1 promoter deletion construct B (harboring a 502 bp 5 , deletion, promoter fragment 1)-driven GUS expression in the shoot elongation zone of an uninfected tobacco plant. No activity is detectable in the roots. D) Whole-mount histochemical GUS assay of a Cel1-transgenic tobacco plant infected with M. incognita. Atcel1 activity is confined to the nematode feeding cells (not shown) and the plant elongation/ differentiation zones. $\mathrm{A}=$ shoot meristem (Construct B shown). E ) Atcell-driven GUS expression within M. incognita-induced giant cells four days post-inoculation of nematodes to an Atcell-GUS transgenic tobacco root (construct A, Fig. 1). GUS activity is confined to the central region of the developing gall tissue. F) Sections ( $30 \mu \mathrm{~m}$ thick) through M. incognita-infected Atcel1-GUS tobacco roots after GUS staining. GUS expression is restricted to the giant cells induced by the nematode. $\mathrm{N}=$ nematode, $\mathrm{GC}=$ giant-cells,
cell promoter could drive UidA gene expression within giant-cells three days after nematode infection of Arabidopsis plants by root-knot nematodes (Mitchum et al., 2004). Upregulation of the full-length Atcel1-UidA construct (construct A) was observed within giant cells induced by root-knot nematodes in tobacco plants (Table 1, Fig. 2E-F) between 4 to 28 dpi. Similar results were obtained for all the time points used in this study. Frequently, full-length Atcell-driven GUS activity was also visible early in the lateral root primordia, even in roots distant from the nematode feeding site. In contrast to full-length Atcel1, in all deletions in the Atcell promoter assayed, including construct B , no detectable activity within the giant cells induced by $M$. incognita was observed (Fig. 1, Table 1).

Deletion of fragment $1(-1,171$ to $-1,673)$ of the Atcel1 promoter abolished expression in both giant cells and uninfected roots in all constructs examined. To determine whether all the motifs needed for Atcell expres-


Fig. 3. Putative cis-acting elements of the Atcell promoter as predicted by the Plant-CARE (Lescot et al., 2002), PLACE (Higo et al., 1999; Rombauts et al., 2003), and MOTIF SAMPLER (Thijs et al., 2001) algorithms. The transcription start point (TSP) is indicated with +1 : transcription start point. Distances in bp are relative to the translation start codon.
sion in roots and in NFS were in fragment 1 of the Atcel1 promoter (Fig. 1), several internal deletions between fragments 1 and 4 were examined (Fig. 1D-F; Table 1). No internal Atcell promoter deletions that included fragment 1 were sufficient for GUS expression in NFS or roots.

Identification of conserved sequence motifs in the Atcell promoter: Because the Atcel1 promoter is upregulated within giant cells induced by root-knot nematode, we analyzed the promoter for specific sequences that act as nematode responsive cis-acting elements that could be responsible for the observed GUS gene expression. Plant-CARE, PLACE, and MOTIF SAMPLER analyses of the $1,673 \mathrm{bp}$ Atcel1 promoter sequence revealed a number of predicted functional motifs found in most eukaryotic promoters, in addition to several potential regulatory elements that have been shown to be functional in other plant promoters (Fig. 3). A typical TATA box was identified at position -31 , and a CAAT box-like sequence was found at position -50 relative to its transcription start point (TSP), respectively. The upstream sequence relative to the TSP of the Atcell has several regions with over $80 \% \mathrm{~A} / \mathrm{T}$ content (data not shown).

To evaluate sequence motifs that may be common among promoters of NFS-expressed genes, we compared motifs found in the Atcell promoter to those of other genes known to be upregulated in NFS. Included were characterized promoters of five other Arabidopsis genes, in addition to Atcell, that are known to be upregulated in NFS, as well as three NFS-responsive promoters from other plant species (Table 2). In particular, the 300 bp 'nematode box' from the tobacco $T o b R B 7$ promoter and a 246 bp fragment from the Hahsp17.7G4 sunflower promoter were included because functional studies have definitively linked these minimal nematode-inducible sequences with NFSspecific expression. Two Arabidopsis promoters reported to be down regulated in NFS were included as negative controls.

Analysis of the promoter regions using MOTIF SAMPLER revealed the presence of several putative regulatory elements that were previously reported in NFS upregulated genes, including E-BOX, AUX-RR, ROOT-MOTIF, and W-BOX (Fenoll et al., 1997; Escobar et al., 1999; Puzio et al., 2000; Mazarei et al., 2002; Thurau et al., 2003). All of these sequences were present in one or both of the promoter sequences from the NFS downregulated genes, suggesting that they do not, in fact, represent NFS-specific transcription factor binding sites. Four motifs (EIRE, ERE, P-BOX, and WUNMOTIF) were present in the NFS upregulated promoter sequences but not in the downregulated ones (Table 2).

Nematode infection of antisense Atcel1 plants: To further investigate the potential function of Atcell during rootknot nematode infection, three Arabidopsis lines expressing an antisense Cell construct (Tsabary et al., 2003) were infected with root-knot nematode. Samples of infected and uninfected plant roots were collected at $3,4,7,14,21$ and 28 dpi . Samples were fixed, sectioned,

Table 2. Putative regulatory elements common between the Atcell promoter (accession X98543) and other plant promoters activated and downregulated in feeding cells induced by root-knot nematodes (RKN) and cyst nematodes (CN).

| Gene construct ${ }^{\text {c }}$ | Accession | Induced by | Element ${ }^{\text {a }}$ |  |  |  |  |  | Reference |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | $\begin{gathered} \text { E-BOX } \\ \text { CANNTG } \end{gathered}$ | EIRE <br> TTCGacc | ERE <br> ATTTcaaa | $\begin{aligned} & \text { P-BOX } \\ & \text { CCTTtg } \end{aligned}$ | $\text { W-BOX }{ }^{\text {b }}$ TTGACC | WUN-MOTIF aAATTtcct |  |
| ATCEL1-FRAG1-GUS | X98543 | RKN | X | X | X | X | X | X | Mitchum et al., 2004 |
| ATPYK20-GUS | AJ249204 | RKN, CN | X | X | X | X | X | X | Puzio et al., 2000 |
| AT\#25.1-GUS | A91914 | RKN,TCN | X | X | X | X | X | X | Ohl et al., 1997 |
| AT\#1164-GUS | A79355 | RKN, CN | X |  | X |  | X |  | Ohl et al., 1997 |
| ATSUC2-GUS | X79702 | RKN, CN | X |  | X | X | X | X | Juergensen et al., 2003 |
| ATATAO1-GUS ${ }^{\text {c }}$ | AF034579 | RKN, CN | X |  |  |  |  | X | Moller et al., 1998 |
| LELEMMI9-GUS | S45406 | RKN | X | X | X | X | X | X | Escobar et al., 1999 |
| NTTOBRB7-0.3-GUS | S45406 | RKN | X |  | X |  |  | X | Opperman et al., 1994 |
| HAHSP17.7G4_83-GUS | U46545 | RKN |  | X |  |  |  | X | Escobar et al., 2003 |
| ATPAL1-GUS ${ }^{\text {d }}$ | X62747 | RKN, CN | X |  |  |  | X |  | Goddijin et al., 1993 |
| AT-TIP-GUS ${ }^{\text {d }}$ | X63552 | RKN, CN |  |  |  |  |  |  | Goddijin et al., 1993 |

[^2]stained, and examined for general root morphology as well as the ability for $M$. incognita to induce giant-cell development. Uninfected A. thaliana plants containing the Atcel1 antisense construct exhibited the same alterations in shoot and root morphology as previously reported (Tsabary et al., 2003). The Atcell antisense plants had shorter stems and roots relative to the wildtype plants (Fig. 4A), indicating that the antisense construct was actively expressed.

Microscopic examination of stained root sections revealed a typical pattern of M. incognita infection and development, as well as typical giant cell formation (Fig. 4B,C). Similar patterns of nematode and giant-cell development were observed in both the wild-type and


Fig. 4. Constitutive expression of antisense Atcel1 in transgenic A. thaliana and infection of these plant roots with the root-knot nematode, M. incognita. A) Both shoot and root development of Atcel1 antisense (ANTI) plants are compromised as compared to wild-type (WT) Arabidopsis. B) Giant cells (GC) form normally around the head of a developing root-knot nematode (N) in a ( $10-\mu$ m-thick) crosssection of a wild-type Arabidopsis root. C) Giant cell and nematode development in antisense Atcel1 Arabidopsis progress normally even as root and shoot development are compromised. Abbreviations are defined in the text and in Table 2.
in the Atcel1 antisense constructs. Similar numbers of NFS developed in wild-type and in the Atcell antisense infected plants ( 1,151 galls $/ 50$ plants in wild type vs. 1,185 galls/ 50 plants in antisense plants; data from two repetitions). Because sections from both wild-type Arabidopsis (Fig. 4B) and Atcel1 antisense Arabidopsis (Fig. 4C) revealed that $M$. incognita J2 penetrated roots of both plant types equally and that giant cell and nematode development were comparable, we did not count numbers of adult female nematodes. However, we did observe that the nematodes completed their life cycle in the antisense plants.

## Discussion

The observed upregulation of the Atcel1 promoter within giant cells induced in roots by root-knot nematodes and the lack of this activity within the feeding sites of cyst nematodes suggested potential transcriptional regulation of Atcell expression upon nematode infection (Mitchum et al., 2004). In this study, a comparative analysis of Atcel1 promoter deletion constructs demonstrates that the region between $-1,673$ and $-1,171$ (fragment 1) was essential to provide specificity of Atcell promoter expression in roots and giant cells. It is unclear if elements within fragment 1 of the Atcel1 promoter that are required for expression within roots are also required for expression within giant cells. Some analyses of gene expression within giant cells (Wilson et al., 1994; Gheysen and Fenoll, 2002) indicate that wild-type expression in roots was not a prerequisite for plant genes recruited during giant-cell formation. The expression of Atcell promoter construct B (containing a deletion of fragment 1) in shoots observed here, however, was uncoupled from expression in giant cells and roots. It has been demonstrated with promoter deletions of the $\operatorname{Tob} R B 7$ gene of tobacco that elements that drive root-specific expression can be uncoupled from elements that can drive expression specifically within giant cells (Opperman et al., 1994). Similarly, the -83 to +163 region of the Hahsp17.7G4 gene in sunflower that contains heat shock element core sequences was sufficient to drive expression within giant cells (Escobar et al., 2003). Inclusion of fragment 1 of the Atcel1 promoter in the absence of internal regions within the Atcell promoter constructs analyzed here, however, indicates that the presence of a fragment 1 alone within Atcell is not sufficient to drive expression within giant cells. The effects may be due simply to the relative change in distance and conformation between upstream and downstream elements. In general, the function of a regulatory region is complex, involving a multiprotein complex interacting with the transcription factors bound to neighboring DNA sites. A supplementary layer of complexity is added by bringing the transcription factors together on the promoter and by adopting a three-dimensional configuration, enabling the interaction with other parts to activate the
basal transcription machinery (Buratowski, 2000; Rombauts et al., 2003).

Alternatively, the activity of other functional elements within the Atcell promoter may also be required for both the root and giant-cell expression. A number of conserved sequence motifs that may represent transcription factor binding sites were found within the Atcell promoter as well as in promoter regions of other nematode induced genes. Interestingly, these motifs include WUN-motif (Van de Loecht et al., 1990; Washida, et al., 1999) and EIRE (Shah and Klessig, 1996; Fuduka, 1997), both of which have been reported in genes that are transcriptionally activated in response to pathogenderived elicitors. The ethylene responsive element (ERE) is found in many pathogenesis-related (PR) genes that are activated during nematode infection (Schwechheimer et al., 1998; Mazarei et al., 2002). The ERE motif was also found in within the Atcell promoter, but not in the promoter regions of nematode-repressed genes. Whether these motifs have specific roles in transcriptional modulation of Atcell remains to be determined; however, they represent candidates for further functional analyses using directed mutagenesis and/or deletions.

Regulation of plant endoglucanase expression at the transcriptional level may be only one level of control of cell wall-modifying activity induced by nematodes within feeding cells. The inability of a functional Atcel1 antisense to affect giant-cell or nematode development here suggests that Atcell activity may not be essential for proper giant-cell formation and/or that functional redundancy in induced endoglucanase activity within NFS exists. In Arabidopsis, the EGase gene family comprises more than 20 members (del Campillo, 1999; Tsabary et al., 2003). Potential functional redundancy is supported by the upregulation of at least five tobacco endoglucanase genes within the feeding cells of both root-knot and cyst nematodes (Goellner et al., 2001). The tobacco endoglucanases upregulated in NFS are phylogenetically distinct, however, and may represent functional differences in both normal plant development and activity within NFS. It remains to be investigated whether an endoglucanase essential to the formation of NFS can be identified.

These results indicate that expression of the Atcel1 promoter in NFS is regulated by the combinatorial interactions of cis-acting regulatory elements in the promoter, including essential element in the distal region of the promoter. The multiple putative cis-acting elements (Fig. 3) of the Atcell promoter accommodate the argument that they may act as coupling elements that may function in different combinations to confer a diversity of tissue-specific, developmental, and stressregulated patterns. The promoter deletions examined did not result in restricting activity to giant cells as observed with the $\Delta 0.3$ TobRB7 element (Opperman et al., 1994). Further work, including the generation of a fine-
scale series of 5 ' promoter deletions within fragment 1 in combination with linker scanning and or/sitedirected mutagenesis will be required to precisely define, if possible, given cis-elements within the Atcell promoter that convey a specific response in NFS. This potential has important implications for strategies to engineer nematode resistance. Nematode-responsive promoters may be used to localize the expression of anti-nematode constructs that interfere with the development of the feeding site and/or nematode specifically to nematode infection sites (Atkinson, 2003). When targeting plant endoglucanase genes for inhibition within NFS, one must consider potential functional redundancy of endoglucanase activity within NFS as well as whether the target gene is essential to the success of NFS formation.

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[^1]:    ${ }^{\text {a }}$ A denotes the full-length $1,673 \mathrm{bp}$ Atcell promoter; B-F are 5' Atcell promoter deletion constructs (serial B and C, internal D, E, and F).
    ${ }^{\mathrm{b}}$ The activity of the reporter gene in roots, shoots, leaves and in NFS as visually determinate are indicated as follows: + and - indicate presence or absence of GUS activity, respectively. Number of nematode feeding sites were collected between 3 to 28 dpi. Similar results were obtained for all the time points used in this study.

[^2]:    ${ }^{\text {a }}$ Regulatory motifs predicted by the Plant-CARE (Lescot et al., 2002), PLACE (Higo et al., 1999; Rombauts et al., 2003), and MOTIF SAMPLER (Thijs et al., 2001) algorithms.
    ${ }^{\mathrm{b}}$ Motif was present in the ATCEL1 promoter but not in the ATCEL1-FRAG.1.
    ${ }^{\text {c }}$ AT: Arabidopsis thaliana, LE: Lycopersicon esculentum, NT: Nicotiana tabacum.
    ${ }^{\mathrm{d}}$ Gene constructs that are downregulated in response to nematode infection.

