The C-terminal propeptide of a plant defensin confers cytotoprotection and subcellular targeting functions

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Abstract

Background: Plant defensins are small (~5 kDa, 45–54 amino acids), basic, cysteine-rich proteins that have a major role in innate immunity in plants. Many defensins are potent antifungal molecules and are being evaluated for their potential to create crop plants with sustainable disease resistance. Defensins are produced as precursor molecules which are directed into the secretory pathway and are divided into two classes based on the absence (class I) or presence (class II) of an acidic C-terminal propeptide (CTPP) of about 33 amino acids. The function of this CTPP had not been defined.

Results: By transgenically expressing the class II plant defensin NaD1 with and without its cognate CTPP we have demonstrated that NaD1 is phytotoxic to cotton plants when expressed without its CTPP. Transgenic cotton plants expressing constructs encoding the NaD1 precursor with the CTPP had the same morphology as non-transgenic plants but expression of NaD1 without the CTPP led to plants that were stunted, had crinkled leaves and were less viable. Immunofluorescence microscopy and transient expression of a green fluorescent protein (GFP)-CTPP chimera were used to confirm that the CTPP is sufficient for vacuolar targeting. Finally circular dichroism and NMR spectroscopy were used to show that the CTPP adopts a helical confirmation.

Conclusions: In this report we have described the role of the CTPP on NaD1, a class II defensin from Nicotiana alata flowers. The CTPP of NaD1 is sufficient for vacuolar targeting and plays an important role in detoxification of the defensin as it moves through the plant secretory pathway. This work may have important implications for the use of defensins for disease protection in transgenic crops.

Keywords: Plant defensin, Phytotoxicity, Vacuolar targeting, Transgenic cotton

Background

Plant defensins are a family of small (~5 kDa, 45–54 amino acids), basic, cysteine-rich proteins that are widespread in the plant kingdom and have been described in various tissues and species [1]. Many, but not all plant defensins, have potent antifungal activity and thus have generated interest in their application for control of fungal disease in crop plants [2].

Plant defensins are divided into two classes based on precursor proteins predicted from cDNA clones [1,3].

Class I defensin precursors have an endoplasmic reticulum (ER) signal peptide and a defensin domain. Defensins from this class enter the ER where they are folded and subsequently secreted via the default pathway into the extracellular space [4,5]. Many of the well-characterised seed defensins belong to this class [1].

Class II defensins have an additional C-terminal propeptide (CTPP) of about 33 amino acids. Most members of this class are produced by solanaceous species where they are expressed constitutively in floral tissues and fruit, and are induced in leaves during salt-stress (reviewed in Lay and Anderson [1]). However, a small number of defensins with CTPPs have been discovered outside the Solanaceae such as ZmESR-6 (Zea mays

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Embryo Surrounding Region-6 from maize (Poaceae) [6]. The CTPPs from NaD1 (from the flowers of Nicotiana alata), PhD1 and PhD2 (from the flowers of Petunia hybrida) are not present on the mature, biologically active defensins [7]. This led to our hypothesis that the CTPP could be a vacuolar targeting signal (VTS) since VTs are removed from vacuolar proteins during transit to, or deposition in the vacuole and unlike the extracellular class I defensins [4,5] the NaD1 defensin is deposited in the vacuole [3].

The CTPPs of solanaceous class II defensins carry an overall negative charge that closely matches and counter-balances the positive charge of the defensin domain [3]. This led us to ask whether the CTPP could have an additional role in protein folding and/or protein detoxification (i.e. preventing phytotoxicity) [3] like the propeptides on plant α-β-thionins and mammalian defensins that share an analogous charge disparity between their pro- and mature protein domains [7].

In a recent paper [8], we described the production and performance of transgenic cotton plants expressing the natural NaD1 precursor consisting of the mature 47 amino acid defensin domain flanked by an ER signal peptide and the 33 amino acid CTPP. These plants were phenotypically indistinguishable from the non-transgenic parents and exhibited enhanced resistance to both Fusarium and Verticillium wilt over three years of field trials.

In this paper, we investigated the role of the 33 amino acid CTPP of NaD1 by generating transgenic cotton plants expressing NaD1 without the CTPP to determine its effect on vacuolar targeting and potential phytotoxicity. Furthermore, we examined whether the defensin CTPP was sufficient to direct cytosolic green fluorescent protein to the vacuole.

**Results**

**Production of transgenic cotton**

Twelve PCR-positive primary transgenic plants (T₀) representing 10 transgenic consisting were produced with the pHEX22 (SP-NaD1ΔCTPP) construct (Table 1 and Figure 1A). All of these plants had detectable levels of NaD1 in the leaves as determined by ELISA (data not shown). An immunoreactive protein of ~5 kDa protein was observed on protein blots of leaf extracts from lines 78.131.1 and 83.68.1 consistent with the size of mature NaD1 (Figure 1B). In contrast, immunoreactive proteins of ~5 kDa and ~9 kDa were present in leaf extracts from the transgenic cotton line D1 (pHEX3: SP-NaD1-CTPP), consistent with the sizes of mature NaD1 and proNaD1 (NaD1-CTPP), respectively (Figures 1A and B). Fifty-eight primary transgenic plants representing 38 transgenic events were produced with the pHEX3 (SP-NaD1-CTPP) construct. Ten plants (17%), all representing different events, were either infertile and/or had distorted or small leaves. The remaining plants had a normal phenotype and were fertile. In contrast, 10 of the 12 primary pHEX22 transgenic plants (83%), representing nine events, exhibited distorted or small leaves and/or short internodes (Table 1). Furthermore, seven of these 10 primary transformants were infertile (Table 1). Two of the primary pHEX22 transformants (from two events) had a normal phenotype.

**Characterisation of segregating T₁ transgenic plants**

To determine if the abnormal phenotype of the pHEX22 transformed cotton plants segregated with the NaD1 gene, seed from the four fertile transgenic events (lines 78.131.1, 83.23.2, 83.68.1 and 83.102.1) were planted in the greenhouse and the germinated T₁ plants were assayed for NaD1 accumulation, number of transgene inserts and phenotype. PCR screening of the T₁ plants indicated that the primary transgenic plants (T₀) of lines 78.131.1 and 83.102.1 had only one copy of the transgene NaD1 (Table 2). Real-time PCR revealed the distribution of homozygous, hemizygous and null plants expected for a single insertion (Table 2) and Southern blot analysis confirmed single inserts (data not shown). There were very few null plants in the segregating T₁ populations of lines 83.68.1 and 83.23.2 which is consistent with the presence of more than one transgene insertion in the primary transgenic plants (Table 2).

Homozygous plants of line 78.131.1 had the highest levels of NaD1 of the pHEX22 transgens with an average of 6.8 ppm in their leaves (Table 3). Hemizygous plants of this line had about half the NaD1 levels at about 2.6 ppm (Table 3). Line 83.102.1 had less NaD1 than line 78.131.1, with an average of 1.7 ppm and 0.9 ppm in the leaves of homozygous and hemizygous plants, respectively (Table 3). Lines 83.68.1 and 83.23.2 had lower levels of NaD1 at about 1.1 ppm and 0.6 ppm, respectively (Table 3). In comparison, the three

| Event | Plant no. | Phenotype | Fertile |
|-------|-----------|-----------|---------|
| 78.131 | 1         | Distorted leaves | Yes |
| 2     |           | Distorted leaves | Yes |
| 83.23  | 1         | Slightly distorted leaves | No |
| 2     |           | Normal | Yes |
| 83.54  | 1         | Distorted leaves, short internodes | No |
| 83.67  | 1         | Distorted leaves, short internodes | No |
| 83.68  | 1         | Normal | Yes |
| 83.96  | 1         | Distorted leaves, short internodes | No |
| 83.102 | 1         | Small leaves | Yes |
| 83.111 | 1         | Normal leaves, short internodes | No |
| 83.166 | 1         | Distorted leaves, short internodes | No |
| 83.182 | 1         | Distorted leaves, spindly habit | No |
homozygous line D1 plants (pHEX3: SP-NaD1-CTPP) that were sown at the same time as the pHEX22 events had about 3.5 ppm NaD1 in their leaves.

Phytotoxicity was observed in the pHEX22 (SP-NaD1ΔCTPP) segregating T₁ plants that was correlated with the level of NaD1 in the leaves. All the leaves on line 78.131.1 homozygous plants, which had the highest levels of NaD1, were severely distorted (Table 3, Figures 2A and C) and these plants were shorter than the null plants. Furthermore, line 78.131.1 homozygous plants were either infertile or produced fewer bolls that were smaller and had less seed than the null plants (Table 3, Figure 2D). All null plants had a normal phenotype (Figures 2B and E) whereas the hemizygous 78.131.1 plants had a phenotype that was intermediate between that of the homozygous and null plants. That is, there was moderate distortion of the leaves (Table 3, Figure 2C) and some reduction in seed production. In comparison, line D1 homozygous plants (pHEX3: SP-NaD1-CTPP) had a normal phenotype (Table 3).

**Figure 1 Production of NaD1 in leaves of transgenic cotton transformed with pHEx22 or pHEx3. (A)** Diagram of the defensin precursors encoded by pHEx3 and pHEx22. Both precursors have ER signal peptides that direct the protein into the secretory pathway. pHEx22 encodes the mature NaD1 defensin of 5.3 kDa and pHEx3 encodes NaD1 with a C-terminal propeptide. This propeptide is removed from the ~9 kDa intermediate during transit to or storage in the vacuole where the mature 5.3 kDa defensin accumulates. Processing sites are indicated by black arrows. (B) Immunoblot of leaf tissue using an anti-NaD1 antibody. Lane 1: 50 ng purified NaD1 from *N. alata* flowers, lane 2: 150 ng purified NaD1 from *N. alata* flowers, lane 3: line D1 (homozygous T₃), lane 4: line 78.131.1 (T₁), lane 5: line 83.68.1 (T₁), lane 6: 83.68.1 (T₁), lane 7: non-transgenic Coker 315. Size markers are SeeBlue Plus2 (Invitrogen).

| Line       | No. plants | PCR -ve | PCR + ve | Homozygous | Hemizygous |
|------------|------------|---------|----------|------------|------------|
| 78.131.1   | 45         | 8 (18%) | 37 (82%) | 14 (31%)   | 23 (51%)   |
| 83.102.1   | 36         | 10 (28%)| 26 (72%) | 8 (22%)    | 18 (50%)   |
| 83.68.1    | 29         | 2 (7%)  | 27 (93%) | MC         | MC         |
| 83.23.2    | 21         | 1 (5%)  | 20 (95%) | MC         | MC         |

The presence of NaD1 was determined by PCR. Lines 73.131 and 83.102 plants were identified as homozygous or hemizygous by real time PCR. Values in parentheses indicate the percentage of the tested plants. Lines 83.68.1 and 83.23.2 had more than one DNA insertion so zygosity was not determined. MC = multicopy lines.
Homozygous plants of line 83.102.1 exhibited some phytotoxicity but it was much less obvious than for line 78.131.1 and was correlated with lower NaD1 expression levels in leaf tissue. There was no difference in leaf morphology between line 83.102.1 transgenic plants and the nulls in the first few weeks, but as the plants matured, some leaf distortion was observed (Table 3). At 87 days after sowing, when plants were flowering, the

| Line   | Zygosity* | Plant no. | NaD1 ppm | Leaf distortion | Fertile | Bolls | Seed | Germination % |
|--------|-----------|-----------|----------|-----------------|---------|-------|------|----------------|
| 78.131.1 Hom 10 | 5.0       | Severe    | Yes      | 1               | 11      | 55    |      |                |
|        23 | 5.9       | Severe    | No       | 0               |         |       |      |                |
|        34 | 7.9       | Severe    | No       | 0               |         |       |      |                |
|        42 | 8.3       | Severe    | Yes      | 1               | 7       | 57    |      |                |
| Ave.   6.8 |           |           |          |                 |         |       |      |                |
| 78.131.1 Hem 48 | 2.7       | Moderate  | Yes      | 2               | 20      | 70    |      |                |
|        12 | 3.0       | Moderate  | Yes      | 1               | 17      | 100   |      |                |
|        44 | 2.4       | Moderate  | Yes      | 2               | 38      | 80    |      |                |
|        47 | 2.2       | Moderate  | Yes      | 2               | 16      | 80    |      |                |
| Ave.   2.6 |           |           |          |                 |         |       |      |                |
| 83.102.1 Hom 7  | 1.5       | Mild      | Yes      | 2               | 25      | 90    |      |                |
|        20 | 2.0       | Mild      | Yes      | 2               | 40      | 80    |      |                |
|        21 | 0.6       | Mild      | Yes      | 4               | 39      | 100   |      |                |
|        27 | 2.6       | Mild      | Yes      | 3               | 33      | 100   |      |                |
| Ave.   1.7 |           |           |          |                 |         |       |      |                |
| 83.102.1 Hem 15 | 0.8       | Slight    | Yes      | 4               | 31      | 90    |      |                |
|        16 | 0.7       | Slight    | Yes      | 4               | 43      | 80    |      |                |
|        25 | 0.9       | Slight    | Yes      | 2               | 49      | 100   |      |                |
|        29 | 1.0       | Slight    | Yes      | 3               | 32      | 80    |      |                |
| Ave.   0.9 |           |           |          |                 |         |       |      |                |
| 83.68.1 MC 1  | 0.6       | None      | Yes      |                 |         |       |      |                |
|        2  | 0.7       | None      | Yes      |                 |         |       |      |                |
|        3  | 1.1       | None      | Yes      |                 |         |       |      |                |
|        4  | 1.0       | None      | Yes      |                 |         |       |      |                |
|        5  | 1.0       | None      | Yes      |                 |         |       |      |                |
|        6  | 2.1       | None      | Yes      |                 |         |       |      |                |
| Ave.   1.1 |           |           |          |                 |         |       |      |                |
| 83.23.2 MC 2  | 0.2       | None      | Yes      |                 |         |       |      |                |
|        3  | 0         | None      | Yes      |                 |         |       |      |                |
|        4  | 3.0       | None      | Yes      |                 |         |       |      |                |
|        5  | 0         | None      | Yes      |                 |         |       |      |                |
|        6  | 0.1       | None      | Yes      |                 |         |       |      |                |
|        7  | 0         | None      | Yes      |                 |         |       |      |                |
| Ave.   0.6 |           |           |          |                 |         |       |      |                |
| D1 Hom 1  | 2.7       | None      | Yes      |                 |         |       |      |                |
|        2  | 3.6       | None      | Yes      |                 |         |       |      |                |
|        3  | 4.1       | None      | Yes      |                 |         |       |      |                |
| Ave.   3.5 |           |           |          |                 |         |       |      |                |

*Hom = homozygous, Hem = hemizygous, MC = multicopy lines, zygosity not determined. NaD1 levels were determined 6 weeks after sowing. ppm = ng NaD1/mg wet weight tissue. 
homozygous plants were shorter and more spindly than the null plants (Figure 3). Hemizygous line 83.102.1 plants had an intermediate phenotype between homozygous and null plants (Figure 3). All line 83.102.1 transgenic plants were fertile although some bolls were smaller than the bolls from null plants (Table 3). The T₁ line 83.68.1 and 83.23.2 plants had, on average, less NaD₁ at 1.1 and 0.6 ppm than the other two transgenic lines, respectively. They had normal phenotype and were fertile (Table 3).
**Cellular structure of leaves from transgenic cotton plants**

Leaves from the non-transgenic Coker control and line D1 (transformed with pHEX3) plants presented characteristic, well-ordered arrangements of upper epidermal, palisade mesophyll, spongy mesophyll and lower epidermal cells (Figure 4). In contrast, cells in the leaves of homozygous line 78.131.1 plants were more irregular in shape and the cell layers were disordered. This observation may explain the altered (‘wrinkly’) leaf morphology presented in Figures 2 and 3.

**Subcellular location of NaD1 in transgenic cotton plants**

Sections from leaves taken from equivalent positions on the homozygous line D1 (pHEX3) and line 78.131.1 (pHEX22) transgenic plants and non-transgenic Coker control plants were probed with anti-NaD1 antibodies and Alex Fluor 568-labelled secondary antibodies to examine the subcellular location of the expressed NaD1. NaD1 was present in the vacuoles of mesophyll cells from line D1 (Figure 5A) but was confined to the periphery of cells in sections from line 78.131.1 (Figure 5B). The antibody did not bind to the leaf sections from the non-transgenic control plants (Figure 5C).

**Examination of subcellular targeting of the GFP chimeras**

In order to assess whether the CTPP from NaD1 was sufficient to direct an otherwise cytosolic protein to the plant vacuole, constructs encoding green fluorescent
protein (GFP) linked to the CTPP from NaD1 or the known C-terminal vacuolar sorting determinant (VSD) of NaPI were expressed transiently in *N. benthamiana* leaves. Micrographs showing the location of the GFP (left panels) produced during transient expression of the various constructs in *N. benthamiana* leaves are presented in Figure 6. The middle panels are equivalent sections showing membranes highlighted by the FM5-95 dye while the right panels are an overlay of the two images. Spectral analysis of emitted fluorescence was used to confirm the specificity of the signals for GFP and the FM5-95 dye [9,10].

GFP was located in the vacuole of cells expressing SP-GFP-CTPP<sub>NaD1</sub> (Figure 6A). In these cells, the junction between the vacuole and the cytoplasm was highlighted by FM5-95 dye which stained both the plasma membrane and the tonoplast. A similar result was obtained with the positive control SP-GFP-VSD<sub>NaPI</sub> construct (Figure 6B). In cells expressing the GFP construct with an ER signal peptide (SP-GFP) but no CTPP or VSD, GFP was located in the extracellular space, with fluorescence concentrated around the periphery of the cell (Figure 6C). This contrasted with the GFP construct lacking both the ER signal (SP) and CTPP/VSD sequences where the expressed GFP was present in the cytoplasm, which was outlined by the fluorescently labelled plasma and tonoplast membranes (Figure 6D).

**Structure of the NaD1 CTPP**
Circular dichroism (CD) and NMR spectroscopy were conducted to determine whether a 33 amino acid synthetic peptide corresponding to the CTPP from NaD1 adopts a helical structure as reported for several other C-terminal vacuolar sorting determinants (VSDs) [11]. The CD spectra indicated a propensity for helical structure that was stabilised in the presence of trifluoroethanol.

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**Figure 6 Location of GFP in *N. benthamiana* cells after transient expression of various GFP-chimera constructs.** Micrographs showing the location of GFP produced during transient expression of (A) SP-GFP-CTPP<sub>NaD1</sub>, (B) SP-GFP-VSD<sub>NaPI</sub>, (C) SP-GFP or (D) GFP constructs in the leaves of *N. benthamiana*. The left panels show GFP fluorescence, the middle panels reveal fluorescence from the FM5-95 dye and the right panels are merge images (c, cytoplasm; ec, extracellular; pm, plasma membrane; t, tonoplast; v, vacuole).
(TFE), as shown in Figure 7A. The NMR data were consistent with this interpretation and analysis of the secondary shifts, shown in Figure 7B, indicated that residues 10–29 form a helix in solution. The secondary shifts for these residues became more negative on addition of TFE, again indicating that the helical structure was stabilised in the presence of TFE (Figure 7B).

**Discussion**

We have discovered that the class II defensin, NaD1, can be phytotoxic in transgenic cotton plants when expressed without its 33 amino acid CTPP. This phytotoxicity was manifested in the leaves of transgenic plants as distortions in the cellular architecture that presented as a wrinkly phenotype at the macroscopic level. The plants were stunted and had short internodes, but more critically, were often infertile or produced small bolls with reduced seed number. The phytotoxicity was most obvious in plants that had accumulated about 7 ppm NaD1 in their leaves and was not detectable at NaD1 levels of about 1 ppm or less. The phenotype segregated with NaD1 in the T1 generation, demonstrating that it had been caused by expression of NaD1 and was not an artefact that had been introduced during tissue culture. This is supported by the observation that 10 out of the 12 primary transformants produced with the SP-NaD1ΔCTPP construct had an abnormal phenotype. This is distinguished from the low number of plants (less than 20%) with an unusual phenotype that normally arise from the cotton embryogenesis regeneration method used to produce the transgenic cotton (unpublished data). This effect was most likely responsible for the 10 plants out of 58 primary transformants (17%) with the SP-NaD1-CTPP construct that were either infertile and/or exhibited distorted or small leaves. Thus, phytotoxicity was not evident in plants when NaD1 was expressed with its cognate CTPP.

Line D1 plants have been assessed in greenhouse bioassays for resistance to *Fusarium oxysporum* f. sp. *vasinfectum* (Fov) and subsequently in field trials over three growing seasons in fields naturally infested with Fov and in two growing seasons in fields naturally infested with *V. dahliae*. In both Fov or *V. dahliae* infested soils, line D1 plants had higher survival rates, greater tolerance to

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**Figure 7** Structural analysis of the NaD1 CTPP. (A) CD spectrum of the 33 amino acid CTPP of NaD1 in 50 mM phosphate buffer pH 7.0 (dashes), 10% TFE (dots), 20% TFE (grey) and 30% TFE (black). (B) Secondary shift analysis of the CTPP in aqueous solution (black bars) and in 20% TFE (white bars). The secondary shifts were calculated by subtracting random coil Hα shifts [12] from the Hα shifts.
the pathogens and 2- to 4-fold increases in lint yield compared to the non-transgenic control plants [8]. Line D1 plants typically accumulated about 3–6 ppm NaD1 in their older leaves and did not exhibit any detrimental agronomic features such as short internodes, decreased height or fertility relative to the non-transgenic parent. Additionally, when the plants were grown in non-diseased soil there was no yield penalty or difference in lint quality when compared to the non-transgenic control plants [8].

Almost 20 unique plant defensins have been transformed into more than 10 different plant species ranging from Arabidopsis to monocot and dicot crop species in an attempt to enhance resistance to a variety of fungal diseases [13]. Given the number of plant defensins tested, the number of crop species transformed and reports of enhanced disease resistance, it is interesting that none have been developed into a commercial trait. One of the most promising reports was from Gao and co-workers [4] who demonstrated that constitutive expression of the class I defensin alfAFP in potatoes conferred enhanced resistance to Verticillium dahliae. This enhanced resistance was maintained in the laboratory and under field conditions over several years at different geographical sites. Much later they reported that plants expressing alfAFP had smaller tubers [14].

To date, the majority of these studies have used class I defensins. In reports where it was explicitly stated, the plants had no obvious deleterious effects from defensin expression. However yield was not reported in any of the glasshouse or field trials and thus, the effects of constitutive expression of class I plant defensins have not been fully evaluated. As reported here, removal of the CTPP from the class II defensin NaD1 to make it more like a class I defensin resulted in abnormal morphology and loss of fecundity. Likewise, Ghag and co-workers have recently demonstrated that the class II petunia defensins [15] are not toxic when expressed in bananas but become phytotoxic when expressed without their cognate CTPPs [16]. Transformation experiments using these defensins without their CTPP produced significantly fewer embryos and these did not develop into mature transgenic banana plants [16].

It is interesting to contemplate why class I defensins are not phytotoxic to their own host plants when they are naturally expressed without a CTPP. Either they do not accumulate to the same high levels as the class II defensins or they have a different mechanism of action. Examination of the three-dimensional structures of the mature domains of various defensins does not indicate any obvious differences between those of class I or class II defensins [1,17,18]. They all adopt comparable and superimposable folds centred on the cysteine-stabilized αβ motif comprising an α-helix and a triple stranded antiparallel β-sheet that is stabilized by a network of four disulfide bonds [7,17,18].

However, although the three-dimensional structure is conserved, the amino acid sequence of plant defensins is highly variable apart from the eight highly conserved cysteine residues, serine 8, an aromatic residue at position 11, glycine 13 and glutamates at position 29 and 34 (numbering relative to RsAFP2) [1]. This has led to the hypothesis that defensins have a conserved scaffold that presents surface loops that are highly variable in sequence and that this variability accounts for the diverse range of biological activities that have been ascribed to defensins [1,19]. NaD1 and other class II defensins from the Solanaceae are thus likely to have a different mechanism of antifungal activity to antifungal class I defensins [13,19]. NaD1, for example, interacts with a specific target on the fungal cell wall before it accesses and permeabilises the plasma membrane and enters the cytoplasm where it may interact with a specific target in the cytoplasm and initiate cell death [20,21]. Psd1, the class I defensin from peas, also enters the fungal cell and travels to the nucleus where it inhibits cell cycle [22]. In contrast, the class I defensins from radish (RsAFP2), dahlia (DmAMP1) and alfalfa (MsDef1) do not enter the cell but interact with specific sphingolipids in the plasma membrane and cell wall leading to membrane disruption and cell death [23-26]. In another mechanism, plant defensins such as MsDef1 and ZmES4 have been reported to interact with ion channels [27,28].

We have previously suggested that the difference in net charge between the mature defensin domain and the CTPP of class II defensins may have evolved as a strategy to prevent deleterious effects on the host cell by making the defensin less reactive (possibly mediated via electrostatic and hydrophobic interactions) [1,3]. This could reduce the likelihood of defensin aggregation or undesirable interactions with other proteins and membranes in the cell during folding and transport through the endomembrane system. In effect, the CTPP could act as an intramolecular chaperone and impart transport competency to the prodefensin.

It is likely that part of the cytoprotective function that is mediated by the CTPP in NaD1 (and probably other class II defensins) is inextricably linked to its vacuolar targeting function. In this paper, we showed by two approaches that the CTPP of NaD1 is necessary and sufficient for vacuolar targeting. Immunofluorescence microscopy on the stably transformed cotton plants revealed a different location for NaD1 expressed with or without its CTPP. When the CTPP was present, NaD1 was targeted to the vacuole and when it was absent, NaD1 was directed to the periphery of the cell. This is consistent with the observations for the well-characterised class I defensins such as RsAFP2 from
radish and MsDef1 from alfalfa, which have an extracellular location [4,5]. The observation that the NaD1 CTPP can direct cytosolic GFP to the vacuoles of *N. benthamiana* plants in transient expression studies demonstrates its sufficiency as a vacuolar sorting determinant.

The sequence of the CTPPs from NaD1 and other class II plant defensins (such as PhD1) were compared to other well characterised vacuolar sorting determinants (VSDs) from other plant proteins (Figure 8). The defensin CTPPs are longer, that is, about 33 amino acids in length compared to 15 amino acids for the C-terminal VSD of barley lectin [29] or seven for tobacco chitinase [30]. Furthermore, while no consensus motif has been identified for C-terminal VSDs, they are all enriched in hydrophobic and acidic amino acids (Figure 8). The first half of the NaD1 CTPP also shares two tetrapeptide motifs with the barley lectin VSD: VFDE and ILAE in the NaD1 CTPP and the motifs VFAE and LVAE in the barley lectin VSD (underlined in Figure 8). Others have reported that as few as three or four amino acids can function as vacuolar sorting signals in plants [31-34], including the VFAE or LVAE motifs of barley lectin [31]. The defensin CTPPs may be longer than normal vacuolar sorting motifs because of their dual function in targeting and detoxification.

We demonstrated that the NaD1 CTPP forms a helical structure in solution like other C-terminal VSDs [11]. From a structural context, it would be interesting to examine the presentation of the CTPP with respect to the mature NaD1 defensin domain. Such information could provide details of possible contacts that could be mediated between the two domains and may form the molecular basis for the protein folding and/or detoxification hypothesis. Alternatively, such information could provide clues as to whether the CTPP is indeed helical in the context of proNaD1 and whether its helical structure is important for its vacuolar targeting function as has been proposed for the NaPI C-terminal VSD [11]. Nielsen et al. [11] suggested that the α-helical structure of the NaPI VSD may be a necessary requirement for the sorting signal to be relatively exposed to interact with the sorting receptor. Indeed, when the slightly curved amphipathic α-helical structure of the NaPI VSD was combined by molecular modelling with the structure of one of the adjoining proteinase inhibitor (PI) domains, the VSD seems to protrude from an otherwise compact PI domain [11].

**Conclusions**

In conclusion, the class II defensin NaD1 is phytotoxic in transgenic cotton plants when expressed without its CTPP. These plants had distortions in their cellular architecture that presented as a wrinkly leaf phenotype, were typically stunted and often infertile or produced small bolls with reduced seed number. We demonstrated by two approaches that the CTPP of NaD1 is necessary and sufficient for vacuolar targeting and when the CTPP was absent, NaD1 was directed to the periphery of the cell. Thus the CTPP plays an important role in targeting and detoxification of the defensin as it moves through the plant secretory pathway. The CTPP is crucial for the transgenic expression of class II defensins to avoid a yield penalty from phytotoxic side-effects.

**Methods**

**Binary vector constructs**

The construct encoding SP-NaD1ΔCTPP (ER signal peptide and mature defensin domain without the 33 amino acid CTPP) was generated by PCR using the pHEX3 plasmid DNA [8] as a template with the forward oligonucleotide primer DEF1 (5′-CCG AAT TCTG CTC GCT CCT TGT G-3′) and reverse oligonucleotide primer DEF3 (5′-GCG GTC GAC TTA ACA TGG CTT AGT ACA TAG G-3′). Primers DEF1 and DEF3 introduced BamHI and SalI sites (underlined), respectively. The PCR product was subcloned into the pCR2.1-TOPO vector (Invitrogen, Carlsbad, CA), excised with BamHI and SalI, and cloned between the CaMV 35S promoter

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**Figure 8** Comparison of the CTPP of NaD1 and PhD1 with C-terminal vacuolar sorting determinants from other plant proteins. The acidic amino acids are boxed in grey and the hydrophobic amino acids are boxed in black. NaD1 and barley lectin have two related tetrapeptide motifs that are underlined.
and terminator sequences of the pAM9 vector [37]. The resultant expression cassette was excised with EcoRI and ligated into pBIN19 binary vector [38]. This construct was named pHEX22. The preparation of pHEX3 is described in [8]. pHEX3 has DNA encoding the entire sequence of the NaD1 precursor (ER signal peptide, defensin domain and CTPP) (GenBank accession no. AF509566).

Production of transgenic cotton
The pHEX22 construct was transferred to Agrobacterium tumefaciens strain LBA4404 and used to infect hypocotyl sections of Gossypium hirsutum L. cv. Coker 315 cotton plants. Embryogenic callus was selected on kanamycin at 35 mg/L and embryos were germinated and plantlets transferred to soil as described in [8]. Plantlets were screened by PCR using primers for the CaMV 35S promoter (forward: 5′-CCT TCC TGT ATA TAA GGA AGT-3′, reverse: 5′-GAT AGA TTT GTA GAG AGA GAC-3′). NaD1 protein levels were determined by ELISA using polyclonal anti-NaD1 antibodies [21] as described in Gaspar et al. [8]. Immunoblot analysis was conducted using polyclonal anti-6H.proNaD1 antibodies [3] as described in Gaspar et al. [8]. The transgene copy number was determined by quantitative real-time PCR as described in Yi et al. [39]. Transgenic cotton plants expressing pHEX3 (including line D1) were prepared and tested as described in Gaspar et al. [8].

Subcellular location of NaD1 in transgenic plants
Leaf segments from non-transgenic, line D1 (transformed with pHEX3) and line 78.131.1 (transformed with pHEX22) plants were fixed in 4% (v/v) paraformaldehyde before paraffin embedding and sectioning. The sections were probed with antibodies raised to hexahistidine-tagged proNaD1 (anti-6H.proNaD1; [3]) (50 μg IgG/mL in blocking solution; 0.2% (v/v) Triton X-100, 1 mg/mL BSA in PBS) for 60 min. The slides were then washed three times with PBS before application of Alex Fluor®-labelled secondary antibodies (Molecular Probes, Carlsbad, CA), diluted 1:200 in blocking solution. The sections were visualised on an Olympus BX50 microscope and images were captured with a Spot RT software (version 3.4, Diagnostic Instruments, Sterling Heights, MI). At the time of fixation, samples were taken from the same leaves for ELISA determination of NaD1 levels as outlined above.

Generation of GFP-CTPP chimera constructs
Specific oligonucleotide primers and PCR was used to generate a number of green fluorescent protein (GFP) chimera expression cassettes that encoded soluble modified GFP (GenBank accession no. U70495, [40]) in combination with DNA encoding the ER signal peptide (SP) from NaPI (GenBank accession no. U08219, nucleotides 10-97) and the CTPP of NaD1 or the C-terminal vacuolar sorting determinant (VSD) sequence of NaPI (GenBank accession no. U08219, nucleotides 1126-1203). These constructs were referred to as SP-GFP-CTPPNaD1 and SP-GFP-VSDNaPI respectively. Two control constructs, SP-GFP and GFP alone, were also made. The PCR amplified products, which incorporated BamHI and SalI sites, were subcloned into the pCR2.1-TOPO vector before they were cloned between the CaMV 35S promoter and terminator sequences of the pAM9 vector and ligated into the pBIN19 binary vector as described for pHEX22.

Transient expression of GFP chimeras in Nicotiana benthamiana leaves and subcellular location of the expressed GFP
Reporter GFP constructs in Agrobacterium were infiltrated into leaves and production and subcellular location of the expressed GFP was monitored by confocal microscopy as described in Conlan et al. [41].

Subcellular location of GFP produced from GFP chimeras
Leaf peels from infiltrated leaf sections were mounted in water on glass slides and glass coverslips were placed on top and secured with melted wax. Samples were examined under a Leica TCS SP2 confocal microscope with a HCX APO 63× W Corr/0.17 CS objective. GFP and FM5-95 fluorescence were monitored at 509 nm and 573 nm, respectively. Ten fields of view were scanned from each leaf section and were captured with confocal LCS 3D software (Leica, Wetzlar, Germany) and Image J (developed by Wayne Rasband, National Institutes of Health, Bethesda, MD; http://rsb.info.nih.gov/ij/). Images are representative of at least three independent experiments.

Peptide synthesis
Boc-L-amino acids were obtained from Merck, 2-(1H-Benzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (HBTU) was obtained from Richelieu Biotechnologies (Quebec, Canada). Trifluoroacetic acid (TFA), N,N-diisopropylethylamine (DIEA), and N,N-dimethylformamide (DMF), all of peptide synthesis grade, were purchased from Auspep (Melbourne, Australia).

The 33 amino acid peptide corresponding to the NaD1 CTPP was assembled using manual solid phase peptide synthesis with Boc chemistry on a 0.5 mmole scale. PAM resin (Applied Biosystems, Foster City, CA) was used and amino acids were added to the resin using HBTU with in situ neutralisation [42]. Cleavage of the peptide from the resin was achieved using hydrogen fluoride (HF) with cresol and thiocresol as scavengers (HF:cresol:thiocresol; 9:1:1 v/v). The reaction was allowed to proceed at −5−0°C for 1 h. Following cleavage, the peptide was dissolved in 50% acetonitrile/0.1%
TFA and lyophilised. The crude peptide was purified using preparative reverse-phase HPLC (RP-HPLC) on a Vydac C18 column. Gradients of 0.1% aqueous TFA and 90% acetonitrile/0.09% TFA were employed with a flow rate of 8 mL/min and the eluant monitored at 230 nm. Mass analysis was performed on a Sciex (Thornhill, Ontario) triple quadrupole mass spectrometer using electrospray sample ionisation.

Circular dichroism spectroscopy
Circular dichroism (CD) spectra were measured on a JASCO J810 spectropolarimeter, installed with a standard analysis program. The temperature was maintained at 25°C. Spectra were recorded with a quartz cell of 0.1-cm path length, using an acquisition time of 2 s/nm, with a 1-nm spectral bandwidth, over the wavelength range from 185 to 260 nm. The samples were prepared by dissolving lyophilised peptide in 50 mM sodium phosphate with the pH adjusted to 7.0. Data are represented in molar ellipticities (θ) (max. degree · cm² · dmol⁻¹). The CD spectra represent an average of at least four accumulations and were corrected by subtracting the buffer baseline.

Nuclear magnetic resonance spectroscopy
Samples for nuclear magnetic resonance (NMR) analysis consisted of ~1 mM peptide in 50 mM sodium phosphate, 90% H2O/10% D2O at pH 5.5 or 7.0. TOCSY, NOESY and DQF-COSY spectra were acquired on either a Bruker 600 MHz or AVANCE 750 MHz spectrometer. Spectra were acquired at 283 and 298 K. TOCSY spectra were acquired with an 80 ms mixing time and NOESY spectra with 150 or 250 ms mixing times. Spectra were acquired with 4096 data points in F2 and 512 in F1, and multiplied with squared sine bell window functions shifted by 90°.

Abbreviations
NaD1: *Nicotiana alata* defensin 1; CTPP: C-terminal propeptide; VSD: Vacuolar sorting determinant; EUSA: Enzyme-linked immunosorbent assay; CD: Circular dichroism; NMR: Nuclear magnetic resonance; SP: Signal peptide.

Competing interests
Hexima Limited funded production of the transgenic cotton plants and hold a patent regarding targeting of plant defensins. All the other work was funded by the ARC. Hexima Limited will not in any way gain or lose financially from the publication of this manuscript. Patent WO/2008/128289, funded by the ARC. Hexima Limited will not in any way gain or lose financially from the publication of this manuscript. Patent WO/2008/128289, funded by the ARC.

Authors’ contributions
FTL, SP, JAM, RLH and MAA participated in study design, coordination and drafting the manuscript, SP designed and built constructs, AAC and JLF carried out plant molecular analysis, BLB performed microscopy, NLD and DJC conducted the CD and NMR and BSM grew and conducted the plant segregation and phenotypic analysis. All authors read and approved the final manuscript.

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