Possible consequences of the overlap between the CaMV 35S promoter regions in plant transformation vectors used and the viral gene VI in transgenic plants

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Introduction

Cauliflower mosaic virus (CaMV) is a pararetrovirus (Caulimoviridae) that infects members of the Brassicaceae family. CaMV was one of first plant DNA viruses to be studied, and its double-stranded circular DNA genome (~8 kg base (kb)) has been completely sequenced. The genome encodes seven genes and has a large (~700 bp) and a small (~150 bp) intergenic region that contain regulatory sequences and single-stranded interruptions. The coding sequences are either separated or overlap by several nucleotides, except for gene VI, which lies between the two intergenic regions. CaMV DNA is transcribed from two promoters in the intergenic regions into two major capped and polyadenylated transcripts, the 19S and 35S RNAs.

The regulatory elements of CaMV have been used since the 1980s to express novel genes in plants; specifically, the 35S promoter (P35S) and terminator are widely used in research and plant biotechnology. The P35S is a strong constitutive promoter, generating high levels of gene expression in dicotyledonous plants. Of the 86 single transgenic plant events that have been authorised in the United States, 54 contain one or more copies of the CaMV P35S.

Odell et al. demonstrated that a P35S that contains 350 bp (-343 to +8, with +1 as the transcriptional start site) is sufficient to obtain constitutive expression, which is due to different domains. The CaMV genome in the region of the P35S region contains multiple overlapping domains, with colinearity between regulatory regions and protein-encoding sequences. The 3' end of P35S overlaps with CaMV polyadenylation regions. The 5' end of P35S overlaps with the 3' end of the coding sequence of gene VI.

The product of gene VI is a multifunctional protein (P6, 62 kDa) that harbours nuclear targeting and export signals and ssRNA-, dsRNA-, and protein-binding domains. Considerable effort has been devoted to determine the various functions of P6.

Bioinformatic tools are increasingly being used in the evaluation of transgenic crops. Guidelines, proposed by WHO/FAO and EFSA, include the use of bioinformatics screening to assess the risk of potential allergenicity and toxicity. With this aim, the EFSA GMO Panel has updated its guidance for the risk assessment of GM plants and proposed to identify all new ORFs due to the transformation event. New ORFs are defined as strings of codons uninterrupted by the presence of a stop codon at the insert genomic DNA junction and within the insert. The putative translation products of these ORFs are then screened for similarities with known toxins and allergens.
vary in length between -1329 to +45 and -300 to +8 (relative position to CAP).

Figure 1 shows a representation of the overlapping elements between the P35S, gene VI, and the 35S terminator, illustrating that the 5' ends of the -300 and -343 P35S variants overlap with domain 4 of P6. The -941 P35S variant overlaps with domains 3 and 4, and, in part domain 2 of P6. The -1329 P35S variant overlaps with domains 2–4 of P6.

Variants that contain one or more duplications of the 35S enhancer have also been created. Kay et al. fused to the -343 to +9 P35S to the -343 to -90 enhancer. These enhancers overlap with domain 4 of P6.

Determine if ORFs within P35S show similarity to allergenic proteins.

Although information is available on the elements important for promoter activity and the functional domains of the overlapping gene VI this information has not been combined to investigate the possible impact of this overlap. In this article, we discuss the possible consequences of the overlap between gene VI and the 35S promoter, when variants of this promoter are introduced into plant nuclear genomes using stable transformation technology. More specifically, we address whether potential expression of the ORFs contained by the P35S promoter overlapping with gene VI: (1) may affect the plant phenotype, and (2) show similarity to known allergenic and toxic proteins.

**Results**

Identification of CaMV 35S promoter variants. The similarity searches against the Patent division of GenBank and information from the literature indicated that different variants of the CaMV P35S are used by plant biotechnologists. These 35S promoters vary in length between -1329 to +45 and -300 to +8 (relative position to CAP).

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Determine if ORFs within P35S show similarity to allergenic proteins. The strategy used to search for similarities with toxic and allergenic proteins is in line with current risk assessment requirements in the European Union.

The DNA sequences of two variants of the P35S were translated and used to search against allergen databases: (1) the -1329 to +60 P35S variant and
and some 35S promoters contain more than one copy of the 35S enhancer. A version that contained the -343 to +4 P35S with a duplicate enhancer (-343 to -90) that has been used in T-DNA vectors, such as pCAMBIA (www.cambia.org/daisy/cambia/585.html).

Multiple allergen databases and search algorithms, described in the EFSA GMO Panel opinion, were used to determine if any of the translated ORFs in the two selected P35S sequences showed similarity to known allergens (Table 1). As described in the Materials and Methods section, the search algorithms recommended by the FAO/WHO 2001 expert panel were used in combination with the FARRP, Allermatch™ allergen and Allergome database. The ADFS was used with the sliding window, word match, and the MEME motif-based method. In addition to these databases, combined with routinely used tools based on percentage identity, the AlgPred database and all provided web tools were used.

None of the searches identified similarities to known allergens. The AlgPred also allows the use of algorithms based on statistical and optimising theory. The vector support machines (SVM) in AlgPred indicated on the basis of the dipeptide composition that the ORF that encoded part of P6 might have some allergenic properties. The sensitivity and specificity of this method is 88.87% and 81.86% respectively and should therefore always be used in combination with other tools. Further analysis of the P6 protein using the SVM method suggested that the potential allergenicity was spread along the protein, except in domain D1 (data not shown).

**Determine if ORFs within P35S show similarity to toxic proteins.** The toxin database was obtained by selecting a subset of sequences from the GenBank non-redundant protein database. No significant hits were obtained to the toxin database using the DNA sequences of the two 35S promoters; all hits had e-values higher than 0.6 (Table 1).

**Discussion**

Multiple variants of the P35S have been constructed and are being used, the lengths of which vary between 1400 to 300 bp, and some 35S promoters contain more than one copy of the 35S enhancer.

Assessing the allergenicity of a transgenic plant is a complex task, and there have been several consensus documents and scientific opinions regarding such assessments of allergenicity. Here, two P35S variants (the -1329 and the short double enhancer variant, respectively) were screened for the presence of ORFs that possibly encode allergenic and toxic proteins. Different databases and search algorithms were employed. No
similarities were shown to known allergens using the different algorithms. The AlgPred SMV algorithms indicated that the ORF-encoding portion of the P6 yields a possible allergen. AlgPred is based on dipeptide composition and calculates the frequency of all possible dipeptide combinations. This approach is theoretical and needs to be used in combination with other methods. As no scientific literature has been reported on any allergenic properties of CaMV and no similarities have been shown to know allergens, it can be concluded that the P6 protein is most likely not an allergen. In addition, a toxin database was constructed, and no significant sequence similarity with the P35S variants was detected. These data suggest that the P35S variants do not contain ORFs that encode for proteins that have allergenic or toxic properties.

Clearly, the longer the P35S, the greater the overlap with the coding sequence of gene VI encoding P6 will be. Our literature survey shows that short versions of the P35S (up to position -522 relative to the CAP) overlap only with domain D4 of P6. This domain, when mutated, deleted, or inverted, reduces the rate of viral movement and influences viral host range.10,29-31 Thus, the D4 domain appears to be partially dispensable. For short P35S sequences that overlap only with the D4 domain of P6 and for promoters that harbour an additional 35S enhancer that overlaps only with the D4 domain, it is unlikely that chimeric proteins will have unintended effects.

The longest identified version of the P35S (-1329) overlaps with all P6 domains except domain D1. The P6 protein that lacks domain D1 localizes exclusively to the nucleus, because D1 contains residues that are required for P6-P6 intermolecular interactions and viroplasm formation.15 At least one of P6’s nuclear functions is to suppress RNA silencing,12 and various abnormalities that are associated with overexpression of P6 have been suggested to correlate with inhibition of tasiRNA processing.33 Variants in which the D1 domain has been deleted inhibit replication of the genome in single cells,12 and De Tapia et al.34 observed that this deleted protein transactivates translation of a polycistronic transcript. Therefore, it is clear that the D1 deletion variant of the P6 protein retains several functions. If a chimeric P6 that contains domains D2–D4 is generated in transgenic plants, it might suppress RNA silencing, affect viral infection through its transactivation activity, or result in an aberrant phenotype. Some of the phenotypes described are leaf chlorosis, vein clearing, plant stunting, late flowering and reduced fertility.30,35-38

Although the P35S overlaps partially with gene VI, the likelihood of unintended effects occurring will depend on whether the partial gene VI is transcribed. We believe that if P35S is embedded in a transformation construct with another gene cassette at its 5’ flank, it is unlikely that the partial gene VI will be transcribed. In contrast, when the P35S is inserted adjacent to plant genomic DNA, transcription from an endogenous plant promoter might take place and create a chimeric protein that contains part of P6. To assess these additional aspects a flowchart has been constructed in Figure 2 to identify the potential unintended effects due to the overlap between the P35S and gene VI. The assessment begins with information on which variant has been used and considers the position of the P35S in constructs and the insertion site. The impact of the insertion site can be determined, based on the phenotype of the transgenic plant and bioinformatic analyses. In case characteristics attributed to the expression of the P6 gene are observed it should be analyzed if the ORF is expressed.

In conclusion, different P35S variants are in use to express proteins in transgenic plants. Here, we detailed the overlap of P35S with the coding sequence of gene VI. Our bioinformatic analyses indicated that no ORFs are present in the P35S that are similar to known toxic and allergenic proteins. Possible unintended effects that are linked to the use of extended versions of the P35S have been determined. The -343 variant, identified by Odell and colleagues,22 contains all of the necessary elements for full promoter activity and does not appear to result in the presence of an ORF with functional domains, rendering it and its related variants the most appropriate promoter variants for avoiding unintended effects.

Materials and Methods

Identification of P35S variants. Similarity searches were performed against the Patent division of GenBank (on 04/08/2010: 15619638 sequences) using the BLASTn algorithm using default parameters. The search was conducted using sequences from the CaMV genome (GenBank accession number V00140.1) from position 6001 to 7500 to retrieve the largest fragment used as P35S.

Allergenicity assessment by sequence analysis. DNA sequences of the P35S variants were translated using the ExPASy tool (expasy.org). The translated sequences were studied in all six frames to determine the similarity to known allergenic proteins using several databases and the search algorithms that were provided.

Similarity searches were performed against: (1) the Food Allergy Research and Resource Program database (FARRP; www.allergenonline.org Version 12 - February 2012: 1603 sequence entries). The 80-amino acid (AA) sliding window method using the criterion of >35% identity as recommended by the FAO/WHO 2001 expert panel, was employed.19,25 In addition, potential identities of 8-AA stretches of identity were investigated; (2) the combined AllerMatch database (December 2005, 792 sequences entries) using the 80-AA sliding window approach and full FASTA, searching for 8-AA exact word matches (allermatch.org); (3) the Allergome database (performed 06/09/2010: 1844 sequence entries) using a full FASTA search (www.allergome.org); (4) the Allergen Database for Food Safety database (ADFS February 2010 version: 1285 sequences, 91 epitopes, 77 structures, and 88 sugar attached entries) using the 80-AA sliding window approach, searching for 8-AA exact word matches and using the MEME motif discovery tool;23 (5) the AlgPred database using all of the provided search tools: (a) IgE epitope and PID search for IgE epitopes, (b) MEME/MAST motif, (c) SVM method based on AA composition, (d) SVM method based on dipeptide composition, and (e) BLAST search of representative allergen peptides.24
Sequence similarity to known toxic proteins. The toxin database was obtained by retrieving a subset of sequences from the GenBank non-redundant protein database (extracted 04/10/2010) using a string search (toxin and toxic). It contains 140774 sequences and was searched using BLASTx with default settings.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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References


