



TMV-Gate vectors: Gateway compatible tobacco mosaic virus based expression vectors for functional analysis of proteins

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Plant viral expression vectors are advantageous for high-throughput functional characterization studies of genes due to their capability for rapid, high-level transient expression of proteins. We have constructed a series of tobacco mosaic virus (TMV) based vectors that are compatible with Gateway technology to enable rapid assembly of expression constructs and exploitation of ORFeome collections. In addition to the potential of producing recombinant protein at grams per kilogram FW of leaf tissue, these vectors facilitate either N- or C-terminal fusions to a broad series of epitope tag(s) and fluorescent proteins. We demonstrate the utility of these vectors in affinity purification, immunodetection and subcellular localisation studies. We also apply the vectors to characterize protein-protein interactions and demonstrate their utility in screening plant pathogen effectors. Given its broad utility in defining protein properties, this vector series will serve as a useful resource to expedite gene characterization efforts.

With the completion or near completion of genome sequencing of many plant species in the last decade, there is a growing need to functionally characterize the large number of predicted proteins and define their physical interaction networks. Consequently, the requirement for vectors that facilitate high-throughput gene cloning and expression of native proteins or derivatives fused with epitope tags and/or fluorescent proteins has become increasingly important. In plants, protein expression can be achieved either by a transient expression approach or by developing stably transformed lines. Transient expression has emerged as an appealing option compared to stable transgenic approaches due to its significant advantages in terms of short production time lines and low cost.

Virus-based expression systems in plants are particularly attractive versus alternative transient expression systems due to the high level of gene multiplication and concomitant elevated levels of expression achievable within a short period of time while minimizing impairment of host activities^{1,2}. Tobacco mosaic virus (TMV) is one of the most extensively studied plant viruses and has thus become a natural choice for vector development. TMV-based vectors have led to recombinant protein yield as high as 80% of total soluble protein^{3,4}. Agroinfection is inexpensive and reproducible, making it a preferred method of delivering viral expression vectors into plant tissues as part of the T-DNA of a binary vector carried by *Agrobacterium tumefaciens*^{5,6}. First-generation TMV-derived vectors that utilized the entire virus genome were limited by low agroinfection efficiency and expression capability⁷. Significant improvements to agroinfection capability has resulted from extensive modification of the TMV genome, including removal of transcript splicing sites, altering codon usage of the RNA-dependent RNA polymerase (RdRP) and introduction of introns within RdRP and movement protein coding sequences⁸. Expression capability has been enhanced by co-expression of *P19*, a RNA silencing suppressor gene of Tomato Bushy Stunt Virus, leading to 10–25 fold increases in recombinant protein accumulation⁹. Additionally, expression capability has been increased multi-fold by positioning the open reading frame (ORF) of the target gene closer to the 3' end of the TMV RNA¹⁰. This has been exploited recently to develop the TRBO (TMV RNA-based overexpression) vector that lacks the coat protein coding sequence, enabling placement of a foreign gene insert close to the 3' end of the viral RNA¹¹. TRBO vector can produce 100-fold more recombinant protein in plants than a *P19*-enhanced transient expression system⁹, and can be easily delivered into plants by agroinfection without the need for co-delivery of a *P19* expression construct¹¹. TRBO vector was capable of producing 3.3 to 5.5 g of recombinant GFP/kg of infiltrated *Nicotiana benthamiana* tissue¹¹.

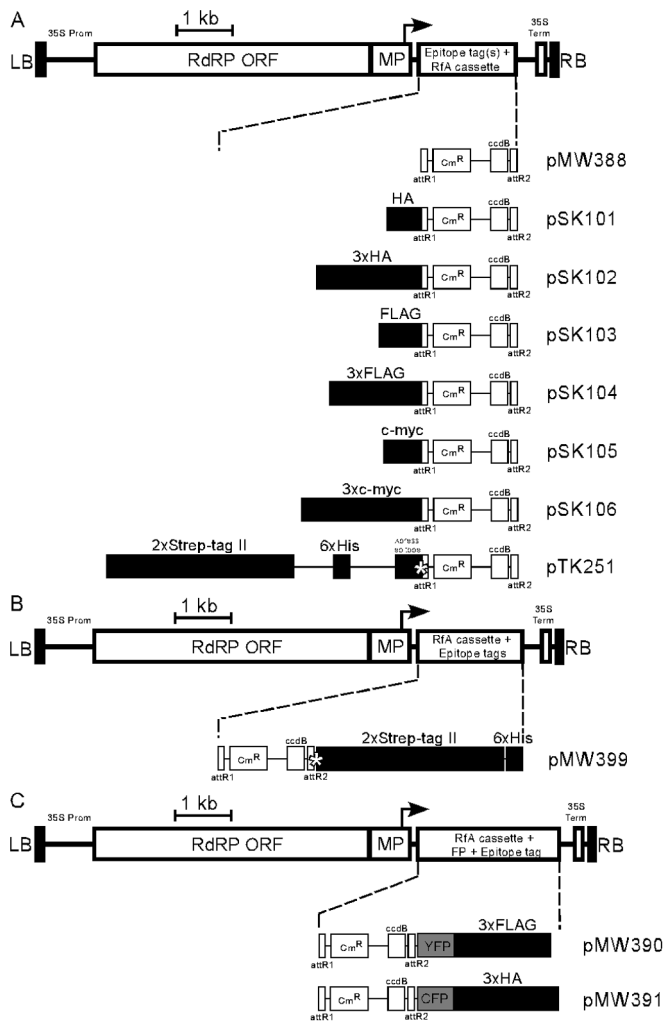


Figure 1 | Schematic illustration of the TMV-Gate vectors. (A) Expression vectors facilitating N-terminal tagging of target protein with various epitope tags, including single- or triple-repeats of HA, FLAG or c-myc, as well as 2xStrep-tag II-6xHis-Xpress epitope. (B) Expression vector facilitating C-terminal tagging of target protein with 2xStrep-tag II-6xHis epitope tags. (C) Expression vectors facilitating C-terminal tagging of target protein with epitope tagged fluorescent proteins, including YFP-3xFLAG or CFP-3xHA. The illustrated vector features include: LB, T-DNA left border; RB, T-DNA right border; 35S Prom, CaMV 35S promoter; 35S Term, CaMV 35S terminator; RdRP, RNA-dependent RNA polymerase; MP, movement protein; bent arrow, coat protein promoter; attR1 and attR2, Gateway recombination sites; Cm^R, chloramphenicol resistance marker; ccdB, *E. coli* DNA gyrase inhibitor; asterisk in pTK251, enterokinase protease recognition site; asterisk in pMW399, 3C protease recognition site; YFP, yellow fluorescent protein; CFP, cyan fluorescent protein. The epitope tag regions in the map are enlarged 1:20 for readability. DNA sequences for the vectors have been deposited at GenBank with accession numbers as follows: pMW388, JX971627; pSK101, JX971619; pSK102, JX971620; pSK103, JX971621; pSK104, JX971622; pSK105, JX971623; pSK106, JX971624; pTK251, JX971625; pMW399, JX971626; pMW390, JX971628; and pMW391, JX971629.

Thus, TRBO vector has great potential for use in high-throughput plant proteomic studies. A major drawback, however, is that it relies on traditional restriction digestion and ligation based cloning for target gene insertion, hampering rapid gene cloning for high throughput functional analysis of proteins. Additionally, the curtailed polylinker region of TRBO vector with only three unique restriction enzyme sites (PacI, NotI, and AvrII) poses another hurdle

in devising cloning strategies for large cDNAs, genomic DNA inserts or inserts carrying translational fusions. To overcome the limitations of restriction and ligation-based cloning, a modified version of the TRBO vector was developed¹² to exploit ligation-independent cloning (LIC)¹³; however, an obvious downside of the LIC strategy is that sub-cloning or re-cloning is not efficient due to the inherent need for PCR re-amplification and, thus, sequencing of the inserts after each procedure¹⁴. A strict requirement for correct T4 DNA polymerase treatment of the insert and vector is another disadvantage of the LIC method.

The time requirements and limitations of traditional cloning procedures have been significantly reduced through development of the Gateway cloning system¹⁵ which employs site-specific recombination to enable inter-molecular transfer of DNA fragments between plasmid vectors. Gateway 'entry' clones of full-length ORFs for several organisms are currently available through public repositories and commercial suppliers. Alternatively, they can be assembled by topoisomerase-mediated cloning¹⁶ or conventional cloning procedures, both technically amenable due to the generally small size of entry vectors. To complement application of ORFs, the genomic coding regions of genes, encompassing the complete repertoire of potential splice variants thereby enabling broader analysis of gene function than possible using an ORF, can also be rapidly incorporated into the Gateway system using a bacteriophage-based homologous recombination system (recombineering)¹⁷. The Gateway vector system has been employed in large-scale plant genomic projects and compatible vectors have been developed for many applications, including ectopic overexpression, promoter-reporter fusions, RNA interference, complementation analysis, gene stacking, and affinity purification using epitope tags^{17–21}. Epitope tagging has proved to be an indispensable way for analyzing protein functions^{22–24} by enabling specific detection of recombinant proteins using commercially available antibodies, thereby eliminating the need for generating antibodies for each protein to be studied. While several non-viral Gateway compatible binary expression vectors incorporating epitope tags have been described in the literature^{18,20}, they are limited by lacking the autonomous replication and high transient expression capability of viral expression vectors. Examples of incorporating Gateway recombination sites into TMV, potato virus X and tobacco rattle virus based expression vectors have been reported^{25–27}; however, with the single exception of an N-terminal GFP fusion variant²⁶, the design of these vectors do not support fusion of target proteins with epitope tags or multiple fluorescent proteins, thus limiting their utility in comprehensive functional analysis studies.

Here we describe construction and functional validation of a unique set of viral expression vectors that combine the favourable attributes of the TRBO vector with Gateway technology, epitope-tagging and fluorescent protein-fusion systems to enable rapid and efficient cloning of gene inserts and production of native or recombinant proteins at very high yields within a short time frame. We further demonstrate the utility of these vectors for protein production, affinity purification, immunodetection, subcellular localisation and analysis of protein–protein interactions as well as plant pathogenic effector screening.

Results

Vector design and features. A major caveat with currently available viral expression vectors which depend on traditional *in vitro* cloning procedures is the inefficiency and laboriousness of cloning ORFs of gene candidates in multiple formats (e.g. epitope-tag variants). To enable rapid cloning of gene inserts and achievement of high level protein expression in a short time, we constructed a set of TMV-based Gateway compatible (TMV-Gate) expression vectors that facilitate production of native as well as epitope- or fluorescent protein-tagged variants (Figure 1). The backbone of these



Table 1 | Recombinant expression constructs generated using TMV-Gate vectors to evaluate their functionality and utility

LR recombination		Recombinant clone	Recombinant protein
TMV-Gate vector	Entry clone		
For ectopic expression			
pMW388	pMW350	pSK133	mGFP5
pTK251	pSK6	pSK154	2xStrep-tag II-6xHis-GusPlus
For ectopic expression, affinity purification and analysis of protein-protein interactions			
pSK101	pMW350	pSK143	HA-mGFP5
pSK102	pMW350	pSK144	3xHA-mGFP5
pSK103	pMW350	pSK145	FLAG-mGFP5
pSK104	pMW350	pSK146	3xFLAG-mGFP5
pSK105	pMW350	pSK147	c-myc-mGFP5
pSK106	pMW350	pSK148	3xc-myc-mGFP5
pTK251	pMW350	pSK149	2xStrep-tag II-6xHis-mGFP5
pSK104	pSW45	pSK84	3xFLAG-GusPlus
pSK101	pTK135	pSK109	HA-AtDMC1 ^{G138D}
pSK102	pTK135	pSK110	3xHA-AtDMC1 ^{G138D}
pSK103	pTK135	pSK111	FLAG-AtDMC1 ^{G138D}
pSK104	pTK135	pSK112	3xFLAG-AtDMC1 ^{G138D}
pSK105	pTK135	pSK113	c-myc-AtDMC1 ^{G138D}
pSK106	pTK135	pSK114	3xc-myc-AtDMC1 ^{G138D}
pTK251	pTK135	pTK255	2xStrep-tag II-6xHis-AtDMC1 ^{G138D}
pMW399	pMW389	pMW403	AtDMC1 ^{G138D} -2xStrep-tag II-6xHis
For analysis of subcellular localisation			
pMW388	pMK6	pMW412	NLS-mCherry
pMW390	pSK212	pSK215	NLS-mCherry-YFP-3xFLAG
pMW391	pSK212	pSK216	NLS-mCherry-CFP-3xHA
For pathogen effector screening			
pSK103	pSU73	pSU101	FLAG-XopD

Destination vectors is derived from TRBO vector¹¹ which includes a full-length cDNA clone of TMV U1 strain modified to lack the coat protein (CP) coding sequence and which is placed adjacent to the cauliflower mosaic virus (CaMV) 35S promoter in a T-DNA binary vector. All of the Destination vectors described in this study use the attR1/R2 RfA Gateway cassette which is linked in-frame to various epitope tags and/or fluorescent proteins and is cloned adjacent to the coat protein promoter of the TRBO vector. These assemblies were confirmed by restriction digests (data not shown) followed by DNA sequencing of the junctions between the TRBO vector, RfA cassette and epitope tag/fluorescent proteins.

The structure of the T-DNA region of various TMV-Gate expression vectors is illustrated in Figure 1. *In planta* expression of the viral replicons in all eleven TMV-Gate vectors is driven by the CaMV 35S promoter (Figure 1). The TMV-Gate vector that allows expression of native proteins is designated as pMW388. TMV-Gate vectors that allow addition of N-terminal epitope tags, such as HA, 3xHA, FLAG, 3xFLAG, c-myc, 3xc-myc and 2xStrep-tag II-6xHis-Xpress epitope tag are designated as pSK101–pSK106 and pTK251, respectively. The TMV-Gate vector that allows addition of a C-terminal 2xStrep-tag II-6xHis epitope tag is designated as pMW399. pTK251 and pMW399 further encode an enterokinase cleavage site or 3C protease site, respectively, that permits proteolytic removal of the epitope tags from the recombinant protein. Another set of TMV-Gate vectors, designated as pMW390 and pMW391, allow generation of C-terminal fusions of target proteins to YFP with a 3xFLAG tag or CFP with 3xHA tag, respectively. All TMV-Gate vectors encode a kanamycin-resistance (Km^R) bacterial selectable marker. Conventional pENTR Gateway vectors also utilise a Km^R marker which may interfere in selection of recombinant clones following the ‘LR’ Gateway insert transfer reaction. To overcome this problem, the entry clone encoding the gene of interest needs to be linearized by a restriction enzyme that uniquely cleaves the backbone of the vector outside the attL sites prior to performing LR recombination reactions. Alternatively, a derivative of the Gateway Entry vector carrying a

different bacterial selectable marker may be used. We have previously described a pENTR2B derivative vector, pJM1 (GenBank: FJ391469) and a pENTR1A derivative, pTK172 (GenBank: FJ410920) in which the Km^R marker is inactivated by insertion of functional tetracycline and zeocin resistance genes, respectively¹⁷.

***In planta* evaluation and utility of TMV-Gate vectors for functional analysis of proteins.** TRBO vector from which TMV-Gate vectors are derived has several favourable attributes, including high agroinfection rates, high protein expression levels, ease of scale-up and bio-containment¹¹. TMV-Gate vectors should perform very similarly to the TRBO vector since none of the TRBO features have been modified during vector assembly. Functionality of TMV-Gate vectors was evaluated for transient expression of native and epitope-tagged proteins, affinity purification, immunodetection, analysis of protein-protein interaction, subcellular localisation and pathogen effector screening. Additionally, the agroinfectivity and expression capability of a subset of the TMV-Gate vectors were analyzed and compared with that of the TRBO vector. To enable these analyses, entry clones encoding ORFs of reporter genes such as mGFP5²⁸, GusPlus and NLS-mCherry as well as AtDMC1 (*Arabidopsis thaliana* disrupted meiotic cDNA 1), an orthologue of a RecA-like protein in yeast that facilitates meiotic homologous recombination²⁹, and XopD (a type III effector protein from *Xanthomonas campestris*)³⁰ were recombined into various TMV-Gate vectors as listed in Table 1.

Agroinfectivity of TMV-Gate vectors. To test the agroinfectivity of TMV-Gate vectors *A. tumefaciens* cells carrying a subset of expression constructs encoding native or epitope-tagged mGFP5 (pSK133 and pSK143–149; Table 1) or 2xStrep-tag II-6xHis-GusPlus (pSK154; Table 1) were infiltrated into *N. benthamiana* leaves and expression of mGFP5 and GusPlus was monitored for 6 days post infiltration (DPI). Expression of these reporter genes was detected at 3 DPI and increased gradually over the next 2–3 days reaching a maximum at

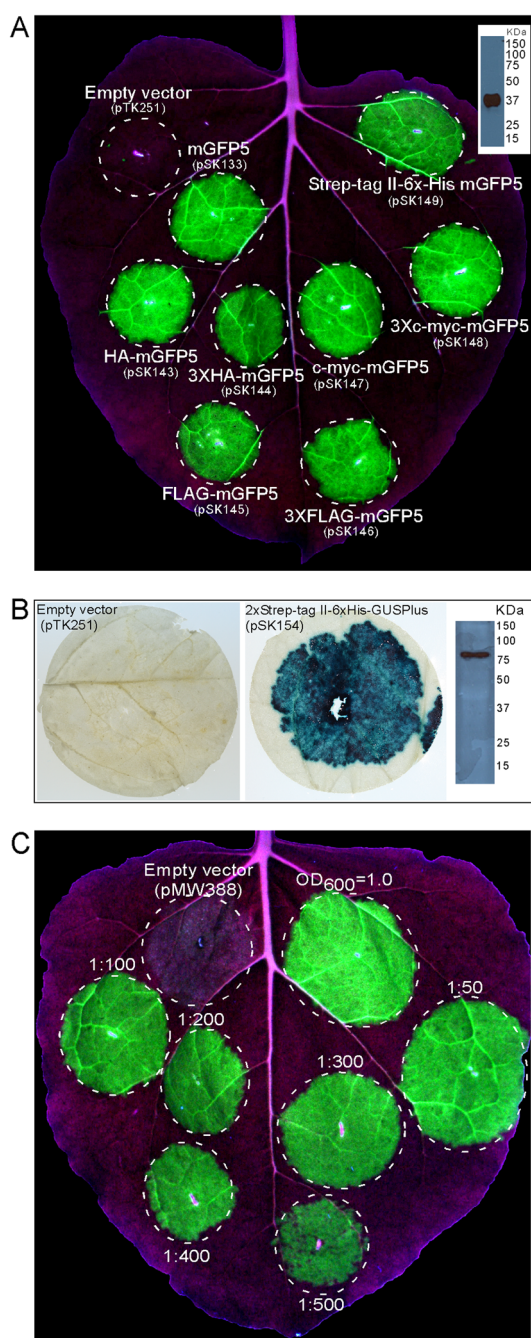


Figure 2 | Transient expression of mGFP5 and GusPlus in *N. benthamiana* leaves by using TMV-Gate vectors. (A) *N. benthamiana* leaf infiltrated with *A. tumefaciens* carrying empty vector or viral expression vector encoding mGFP5 with or without translational fusion to either HA, 3xHA, FLAG, 3xFLAG, c-myc, 3xc-myc or 2xStrep-tag II-6xHis-Xpress epitope tag(s) was illuminated under UV light at 5 DPI and photographed. Inset shows immunoblot detection of 2xStrep-tag II-6xHis-Xpress epitope-tagged mGFP5 (*in silico* predicted size of 37.8 kDa) in crude extract using anti-6xHis antibody. (B) *N. benthamiana* leaf discs from the area infiltrated with *A. tumefaciens* carrying empty vector (pTK251) or viral expression vector encoding 2xStrep-tag II-6xHis-GusPlus (pSK154) were excised 5 DPI and stained for GUS activity. Inset in right panel shows immunoblot detection of 2xStrep-tag II-6xHis-Xpress epitope-tagged GusPlus (*in silico* predicted size of 81.5 kDa) in crude leaf extract using anti-6xHis antibody. (C) *N. benthamiana* leaf infiltrated with various dilutions of a cell suspension of *A. tumefaciens* at OD_{600} of 1.0 carrying pSK133 (pMW388 encoding mGFP5) and photographed with UV illumination at 5 DPI.

5–6 DPI. At 5 DPI, most of the cells in the infiltrated leaf area were found to be expressing mGFP5 (Figure 2A) as well as GusPlus (Figure 2B), thus confirming the high agroinfection capability and expression potential of TMV-Gate vectors.

The CP gene is required for systemic movement of virus particles^{31,32}. Thus, TMV CP deletion vectors, such as TRBO¹¹ and those developed by ICON Genetics³, cannot form virions and are unable to move systemically in plants, desirable attributes for bio-containment. TMV-Gate replicons share the feature of non-systemic movement as evident from the infiltration zone-restricted expression of mGFP5 and GusPlus (Figure 2). This feature of TMV-Gate vectors thus possesses the desirable attribute of bio-containment by reducing plant to plant transmission and persistence in the environment.

Lindbo¹¹ demonstrated that *N. benthamiana* leaves can be efficiently agroinoculated with TRBO-GFP replicon over a wide range of *A. tumefaciens* cell densities (upto 1:300 dilution from a starting optical density at 600 nm (OD_{600}) of 1.0). To determine whether dilute suspensions of TMV-Gate vectors also can be used to efficiently inoculate *N. benthamiana* leaves, we evaluated various dilute suspensions (upto 1:500 dilution from a starting OD_{600} of 1.0; Figure 1C) of *A. tumefaciens* carrying pSK133 (pMW388 encoding mGFP5; Table I) and the expression of mGFP5 was monitored over the next five days. Expression of mGFP5 in all treatments began to be visible at 3 DPI and nearly all cells in the leaf zone infiltrated with *A. tumefaciens* cell suspension diluted upto 1:400 expressed mGFP5 at 5 DPI, suggesting that, similar to the TRBO vector, the agroinfectivity of TMV-Gate vectors remains high at a wide range of cell densities of inoculum. A low density of *A. tumefaciens* cells [dilution of 1:500 (Figure 2C) and beyond (data not shown)] resulted in reduced mGFP5 expression, while higher *A. tumefaciens* cell density ($OD_{600} > 1.0$) occasionally resulted in yellowing and necrotic spots in the infiltration zone (data not shown), probably due to host cell death triggered as a defence mechanism in response to the high titre of inoculum. These results illustrate the inoculum cell density and the length of infection by *A. tumefaciens* cells carrying TMV-Gate-based expression constructs may be varied to achieve optimal levels of recombinant protein production for different applications.

Expression of native and fusion proteins using TMV-Gate vectors.

To determine the protein production efficiency of TMV-Gate vectors, mGFP5 accumulation in *N. benthamiana* leaves infiltrated with *A. tumefaciens* cells carrying empty vector (pMW388), TRBO-GFP¹¹, pSK133 (pMW388 encoding mGFP5) and pSK143 (pSK101 encoding HA-tagged mGFP5) was quantified using an ExperionTM automated electrophoresis system. Functionality of the constructs was first confirmed by assessing GFP fluorescence in agroinfiltrated *N. benthamiana* leaves (Figure 3A). Protein extracts from infiltrated leaf sectors were prepared at 5 DPI and were resolved using an Experion Pro260 protein chip. A virtual gel image of the proteins detectable in each extract is shown in Figure 3B. As expected, the sample from the empty viral replicon extract did not possess a protein representing GFP, whereas distinct bands for GFP (~27 kDa) and HA-GFP (~30 kDa) were seen in protein extracts from *N. benthamiana* leaves agroinfected with TRBO-GFP or pSK133 and pSK143, respectively. Quantification of resolved proteins indicated that TRBO-GFP, pSK133 and pSK143 produced 4.7, 2.5 and 2.8 mg of GFP g^{-1} FW, respectively (Figure 3C). This GFP production by TRBO-GFP is in agreement with values (3.3 to 5.5 mg GFP g^{-1} FW) reported earlier¹¹. It has been previously demonstrated that the level of recombinant protein production increases proportionately with the proximity of the corresponding gene insert to the viral 3' terminus^{10,11}. Thus, the relatively lower levels of protein produced by TMV-Gate replicons (pSK133 and pSK143) may be due to the incorporation of ~70 bp attR sequence between the GFP coding region and the 3' terminus of the TMV RNA genome. Nevertheless, the levels of protein produced by the TMV-Gate

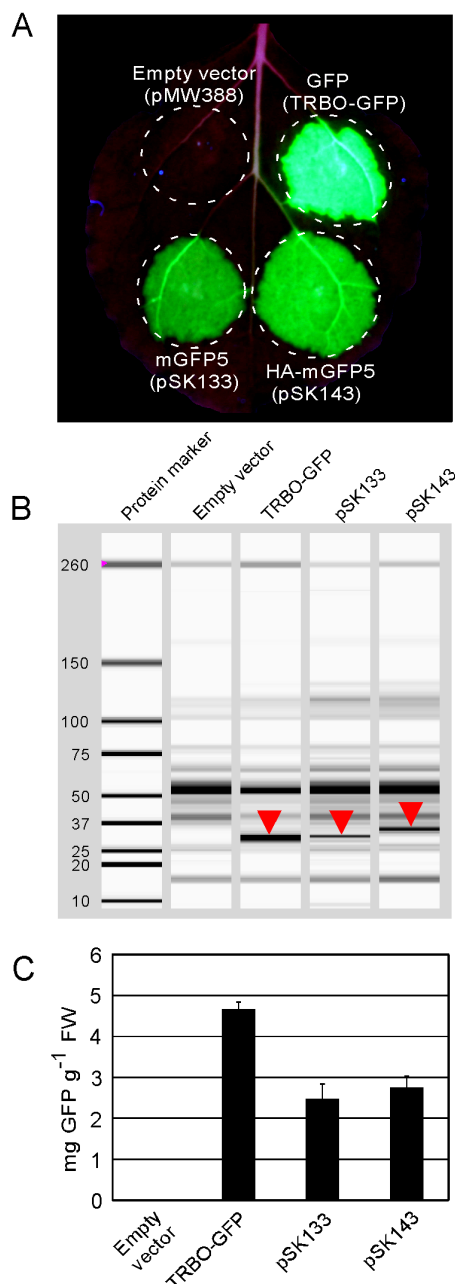


Figure 3 | Quantitative analysis of transient GFP production in *N. benthamiana* using TRBO or TMV-Gate vectors. (A) Fluorescence emitted from GFP transiently produced in a *N. benthamiana* leaf by infiltration with *A. tumefaciens* carrying TRBO-GFP (Lindbo, 2007b) or TMV-Gate constructs pSK133 encoding mGFP5, pSK143 encoding HA-mGFP5 or pMW388 (empty vector) and photographed with UV illumination at 5 DPI. (B) Assessment of GFP accumulation by automated capillary electrophoresis analysis. *N. benthamiana* leaf discs from the area outlined in (A) were excised at 5 DPI and corresponding protein extracts were resolved and GFP identified using the Experion capillary electrophoresis system. Position of the GFP band in each sample is indicated by downward pointing triangles in the simulated gel view produced by the Experion software. The shift in the molecular weight of GFP in pSK143 is due to the presence of an HA tag at the N-terminus. (C) Quantification of GFP resolved by automated capillary electrophoresis analysis using the Experion system. Error bars represent standard deviation. FW, fresh weight.

replicons is much higher than the amount of GFP reported¹¹ to be produced by conventional non-TMV based agroinfiltration vectors (0.03–0.04 mg GFP g⁻¹ FW), full-length TMV genome based vectors (1.5–1.7 mg GFP g⁻¹ FW) or pEAQ series of Cowpea mosaic virus-derived binary vectors (0.7–1.25 mg GFP g⁻¹ FW)³³.

Affinity purification and immunodetection of epitope-tagged proteins expressed using TMV-Gate vectors. TMV-Gate vectors designed to express epitope-tagged proteins allow addition of single or triple-repeats of several commonly used epitope tags, such as HA, FLAG, c-myc, Strep-tag II, 6xHis, and Xpress epitope, to either the N- or C-terminus of target proteins (Figure 1). To demonstrate the utility of these vectors in expression, detection and purification of recombinant proteins, the ORF of an AtDMC1 variant with a defect in its ATPase motif (AtDMC1^{G138D}; see below) was recombined into TMV-Gate vectors, including pSK101–106, pTK251 and pMW399, resulting in expression constructs pSK109–114, pTK255 and pMW403, respectively (Table I). Leaf tissues encompassing zones agroinfiltrated with these constructs were collected at 5 DPI, total protein extracted and the epitope tagged proteins affinity purified by using either agarose resins possessing epitope tag-specific antibody or Strep-Tactin matrix columns. Immunodetection of various epitope-tagged versions of AtDMC1^{G138D} expressed from TMV-Gate vectors is shown in Figure 4. The anti-HA, -FLAG, -c-myc, -Strep-tag II and -6xHis antibodies effectively recognised both the single- (molecular mass of ~42 kDa) and triple-repeat (molecular mass of ~44 kDa) epitope tagged AtDMC1^{G138D} fusion proteins from the input as well as affinity purified fractions. These results demonstrate the functionality of the epitope and affinity purification modules of the TMV-Gate vectors.

Epitope tagged proteins are most commonly eluted from affinity matrices using SDS-PAGE sample buffer. This elution method typically results in denaturation and loss of activity of the purified proteins. Proteins can be eluted under native conditions by competitive elution using epitope tag-specific synthetic peptides or counter ligands. To test this possibility, the single or triple-repeat FLAG-tagged AtDMC1^{G138D} expressed using TMV-Gate vectors and immobilized on anti-FLAG agarose affinity resin was eluted using 3xFLAG peptide. As shown in Figure 4B, the 3xFLAG peptide efficiently eluted the triple-repeat FLAG-tagged AtDMC1^{G138D}; however, as suggested by the peptide manufacturer, it failed to elute the single-repeat FLAG-tagged AtDMC1^{G138D}. Similarly, the 2xStrep-tag II-6xHis-Xpress epitope-tagged AtDMC1^{G138D} immobilized on Strep-Tactin matrix was eluted using desthiobiotin as a counter ligand. Experion-based analysis of the purified AtDMC1^{G138D} revealed the presence of a single band at ~50 kDa (Figure 4F), the predicted size of the fusion protein, indicating high purity of the product. Subsequently, the epitope tags from the purified fusion protein were cleaved off by treatment with enterokinase and the resultant AtDMC1^{G138D} was successfully used for raising antibody in a rabbit (data not shown). Taken together, the results of these experiments establish the suitability of TMV-Gate vectors for expression of recombinant proteins, their immunodetection and affinity purification to obtain native proteins of high purity.

Utility of TMV-Gate vectors for analysis of protein-protein interaction. Protein-protein interactions play a central role in the maintenance and control of cellular and organismal processes. The study of protein-protein interactions has been greatly enhanced by epitope-tagging technology facilitating ‘pull-down’ experiments that can be used to detect and validate protein interactions. In order to establish the utility of epitope-tagging TMV-Gate vectors for analysing protein-protein interactions we employed AtDMC1. The yeast and human DMC1 orthologues have been shown to physically interact homotypically by yeast two-hybrid (Y2H) analysis and, in the case of human DMC1, by immunoprecipitation³⁴. DMC1 DNA strand exchange activity *in vitro* is dependent upon ATP binding

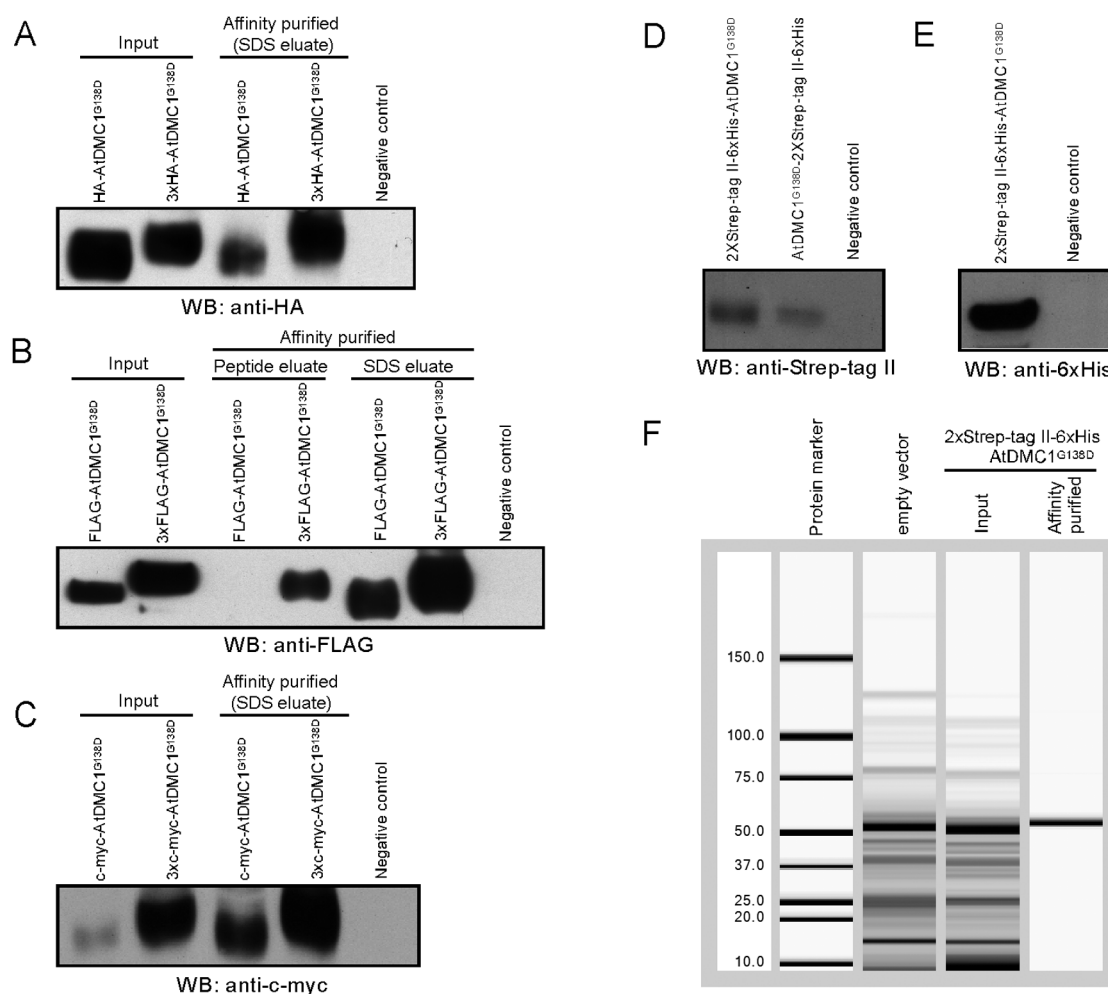


Figure 4 | Affinity purification and detection of epitope-tagged AtDMC1^{G138D} transiently expressed in *N. benthamiana* leaves by using TMV-Gate vectors. Epitope-tagged recombinant AtDMC1^{G138D} immobilized on anti-HA, anti-FLAG or anti-c-myc antibody-conjugated agarose beads were eluted either by boiling the beads in SDS-PAGE sample buffer (SDS eluate) or by incubating the beads with 3xFLAG peptide (peptide eluate). The Strep-tag II-tagged AtDMC1^{G138D} immobilized on Strep-Tactin matrix was eluted using desthiobiotin. Equal volume aliquots of the input protein extract and eluted AtDMC1^{G138D} proteins were subjected to SDS-PAGE, transferred onto PVDF membrane then subjected to immunoblot analysis. WB, western blot. (A) Immunoblot detection of HA- or 3xHA-tagged AtDMC1^{G138D} protein affinity captured using anti-HA agarose affinity gel. (B) Immunoblot detection of FLAG- or 3xFLAG-tagged AtDMC1^{G138D} protein affinity captured using anti-FLAG agarose affinity gel. (C) Immunoblot detection of c-myc- or 3xmyc-tagged AtDMC1^{G138D} protein affinity captured using anti-c-myc conjugated agarose beads. (D) Immunoblot detection of 2xStrep-tag II-6xHis-Xpress epitope-tagged AtDMC1^{G138D} in crude extract using anti-Strep-tag II antibody. (E) Immunoblot detection of 2xStrep-tag II-6xHis-Xpress epitope-tagged AtDMC1^{G138D} in crude extract using anti-6xHis antibody. (F) Capillary electrophoresis analysis using the Experion system of 2xStrep-tag II-6xHis-Xpress epitope-tagged AtDMC1^{G138D} protein affinity purified by using Strep-Tactin matrix.

and hydrolysis mediated by an ATPase motif highly conserved between prokaryote and eukaryote RecA-like proteins³⁵, including AtDMC1. Notably, mutating the ATP-binding motif in yeast DMC1 (yDMC1^{G126D}) results in defective meiotic homologous recombination; however, homotypic interactions of yDMC1^{G126D} have not been demonstrated because of instability of the Y2H plasmid expressing this protein³⁴. In characterising AtDMC1 we created a variant (AtDMC1^{G138D}) analogous to yDMC1^{G126D} and sought to assess if it could interact homotypically. Self-interaction of AtDMC1^{G138D} was initially analysed by Y2H assay wherein interaction of the bait and prey fusion proteins results in expression of a *HIS3* reporter gene enabling growth of *his3* auxotrophic yeast cells on a selective medium lacking histidine. When the AtDMC1^{G138D} bait was expressed with the non-interacting empty prey vector no growth of yeast cells was detected in histidine-deficient plates containing 15 mM 3-amino-1,2,4-triazole (3-AT) which was added to suppress autoactivation of the reporter gene by the AtDMC1^{G138D} bait. In contrast, co-expression of AtDMC1^{G138D} as bait and prey enabled growth of

yeast cells, indicating that AtDMC1^{G138D} can interact homotypically (Figure 5A) and, as proposed for yDMC1^{G126D}³⁴, given the substitution of glycine by the relatively bulky and negatively charged aspartate likely does so independently of ATP binding.

To validate the results of the Y2H assay, we performed pull-down experiments with various versions of epitope tagged-AtDMC1^{G138D} expressed using TMV-Gate vectors. FLAG-tagged AtDMC1^{G138D} was first immobilized on anti-FLAG agarose beads and then incubated with crude protein extracts containing c-myc-tagged AtDMC1^{G138D} or c-myc-tagged mGFP5 to allow protein complex formation. Western blot analysis of the proteins eluted off the anti-FLAG beads after thorough washing revealed the presence of c-myc-tagged AtDMC1^{G138D}, with expected molecular mass of 42 kDa, but not the c-myc-tagged mGFP5 (Figure 5B, left panel), demonstrating that FLAG-tagged AtDMC1^{G138D} interacted with c-myc-tagged AtDMC1^{G138D} but not with c-myc-tagged mGFP5. In a similar manner, 3xHA-tagged AtDMC1^{G138D} was immobilised on anti-HA antibody conjugated beads and incubated with crude protein extract of

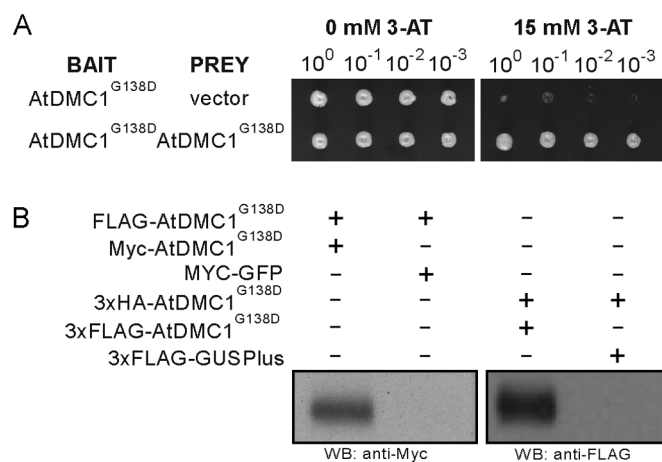


Figure 5 | Protein-protein interaction of AtDMC1^{G138D} expressed using TMV-Gate vectors. (A) Yeast two-hybrid analysis demonstrating self-interaction of AtDMC1^{G138D}. (B) Confirmation of AtDMC1^{G138D} self-interaction by pull-down experiment. The FLAG-tagged AtDMC1^{G138D} and 3xHA-tagged AtDMC1^{G138D} immobilized on anti-FLAG and anti-HA agarose beads, respectively, were incubated with total protein extract of *N. benthamiana* leaf discs transiently expressing c-myc-tagged AtDMC1^{G138D} or c-myc-tagged mGFP5 (negative control) and 3xFLAG-tagged AtDMC1^{G138D} or 3xFLAG-tagged GusPlus (negative control), respectively; corresponding TMV-Gate vectors and recombinant plasmids are listed in Table I. After washing the affinity matrices, proteins were eluted using SDS-PAGE sample buffer, resolved by SDS-PAGE and visualized by immunoblotting with antibodies as indicated. WB, western blot.

3xFLAG-tagged AtDMC1^{G138D} or 3xFLAG-tagged GusPlus then eluted proteins were probed with anti-FLAG antibody. As predicted, 3xHA-tagged AtDMC1^{G138D} interacted with 3xFLAG-tagged AtDMC1^{G138D} but not with 3xFLAG-tagged GusPlus (negative control; Figure 5B, right panel). These results, apart from confirming the ability of AtDMC1^{G138D} to interact homotypically and validating the results of the Y2H assay, demonstrate the applicability of TMV-Gate vectors for the analysis of protein-protein interactions.

Utility of TMV-Gate vectors for analysis of subcellular localisation of proteins. In recent years, translational fusion of candidate proteins to a fluorescent protein combined with microscopy has become a powerful tool for defining protein function through determining its subcellular localisation. TMV-Gate vectors designed to express C-terminal YFP or CFP fusion proteins include an in-frame 3xFLAG or 3xHA epitope tag, respectively (Figure 1), and thus enable simultaneous analyses, including (i) *in planta* evaluation of subcellular localisation of fluorescent protein fusions and (ii) immunodetection, affinity purification and analysis of protein-protein interaction by virtue of the epitope tags attached at the C-terminal ends of the fusion proteins. To test the suitability of these vectors for studying subcellular localisation of proteins, the mCherry red fluorescent protein linked to a nuclear localisation signal derived from the SV40 large T-antigen (NLS-mCherry) was recombined with pMW388; additionally, a variant lacking the stop codon (NLS-mCherry^{ΔSTOP}) was recombined with the TMV-Gate vectors pMW390 or pMW391 to generate in-frame fusions with YFP-3xFLAG or CFP-3xHA, respectively (Figure 6A). The resultant constructs, designated as pMW412, pSK215 and pSK216, respectively (Table I), were agroinfiltrated into *N. benthamiana* leaves and subcellular localisation of the fusion proteins assessed by fluorescent microscopy. The subcellular fluorescence signal emitted by NLS-mCherry overlapped with the DAPI (4',6-diamidino-2-phenylindole; a nuclear stain) signal (Figure 6B), demonstrating that NLS-mCherry localizes to the nucleus and can be utilized as a positive

control in nuclear localisation experiments. As expected, both NLS-mCherry-YFP-3xFLAG and NLS-mCherry-CFP-3xHA fusion proteins localized to the nucleus, as deduced by comparing the fluorescence signal with the DAPI signal (Figure 6B). These results demonstrate pMW390 and pMW391 TMV-Gate vectors can be utilized for detection of subcellular localisation of proteins *in planta*.

Utility of TMV-Gate vectors in pathogen effector protein screening. Effectors are pathogen-derived molecules that manipulate host cell structure and function, thereby facilitating infection and/or triggering disease resistance, culminating in a hypersensitive response (HR)-based programmed cell death^{36,37}. In recent years, the combination of genome sequencing of plant pathogens with computational data mining has dramatically accelerated the identification of effector candidates. Developing high-throughput vector platforms for functional screening of candidate effectors thus would provide an important complement to the bioinformatics approach of identifying effectors. Ectopic expression of effector proteins in plant cells often leads to phenotypic effects, such as induction of HR-like necrosis³⁸. To test the suitability of TMV-Gate vectors for the expression and functional screening of effector proteins we selected XopD which has been shown to elicit HR-like tissue chlorosis and necrosis symptoms in *N. benthamiana*³⁹. The ORF of XopD was recombined in-frame with the FLAG epitope tag of pSK103, resulting in pSU101. This and pSK145 encoding mGFP were agroinfiltrated into *N. benthamiana* leaves at 1:5 dilution from the initial OD₆₀₀ of 1.0 (optimised to overcome the necrosis potentially induced by *A. tumefaciens*). Transient expression of FLAG-XopD resulted in localized tissue chlorosis by 5–6 DPI and necrosis by 9–10 DPI (Figure 7). In contrast, transient expression of FLAG-mGFP5 did not elicit such symptoms. These data, in agreement with a previous report by Kim et al.³⁹, illustrate that XopD is capable of eliciting an HR response in *N. benthamiana* and also confirm the suitability of TMV-Gate vectors for effector screening.

Discussion

In this study, we describe construction and validation of a viral vector system for expedited production and characterisation of recombinant proteins via transient expression in plant leaves. The vectors are derived from the TRBO variant of TMV previously reported¹¹ as enabling high-level production of recombinant proteins but which is severely constrained in practical applications by requiring reliance on traditional DNA cloning procedures. The vectors described in this study are unique from other plant virus vectors integrating Gateway technology^{25–27} by further incorporating a very broad series of epitope, affinity purification and fluorescent protein tags to more readily exploit the attributes of TMV for rapid production and analysis of protein attributes regarding their biochemical and cell biology properties. The compatibility of this expression system with Gateway technology enhances its suitability for high-throughput functional analysis studies. Gateway technology has been widely adapted by the research community to facilitate high-throughput proteomic research. We anticipate that this vector system will complement the on-going and future efforts to clone and functionally characterize ORFeomes using Gateway pENTR vectors⁴⁰.

We demonstrate the utility of the vector system through examples of protein-protein interaction analysis and protein sub-cellular localisation. Effective protein expression using the system was demonstrated by immunoblot analysis of AtDMC1 and GFP with eight different tags or GusPlus with 2xStrep-tag II-6xHis tag revealing no truncated recombinant proteins; furthermore, in all of these cases the proteins were extracted without a denaturation step, collectively supporting their accumulation as intact soluble cellular constituents and not as truncated proteins or insoluble aggregates. Because of the capability to generate grams of recombinant protein per kg of fresh weight biomass within a short time frame, this vector set has great

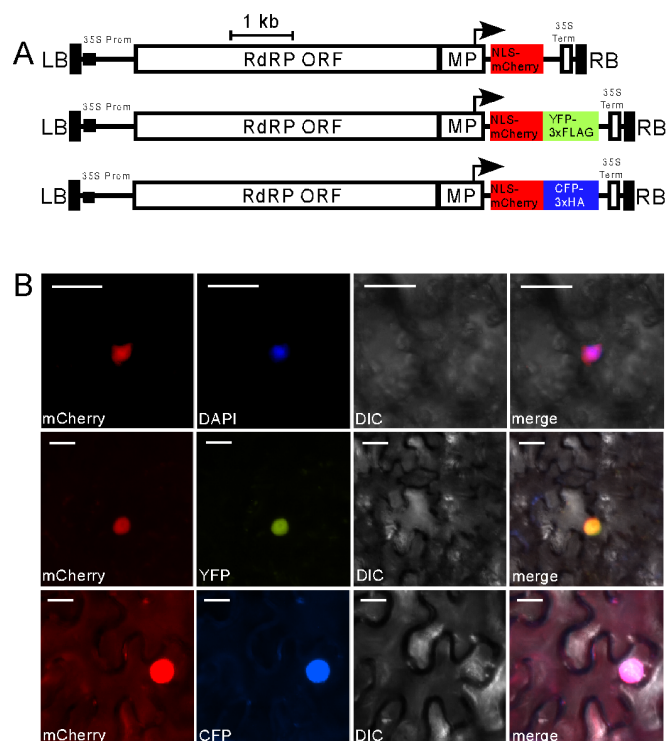


Figure 6 | Nuclear localisation of NLS-mCherry expressed using TMV-Gate vectors. (A) Schematic diagram showing constructs used for analysis of subcellular localisation of NLS-mCherry. mCherry carrying a nuclear localization signal (NLS) was introduced unmodified into pMW388. A NLS-mCherry variant without translational stop codon was linked in-frame with the coding sequence of YFP-3xFLAG or CFP-3xHA in pMW390 and pMW391, respectively. Annotated features as per Figure 1. (B) Epifluorescent microscopy images showing nuclear localisation of NLS-mCherry (upper panel), NLS-mCherry linked to YFP-3xFLAG (middle panel) or NLS-mCherry linked to CFP-3xHA in cells of *N. benthamiana* leaves infiltrated with *A. tumefaciens* carrying the constructs described in (A) and imaged 3 DPI. The nucleus was detected by staining cells with DAPI. Images were taken using YFP, CFP, mCherry or DAPI filter sets as indicated. Scale bar, 20 μ m. DIC, differential interference contrast.

potential to be utilized in large scale protein production applications. Additionally, the mid- to high-level of protein production inherent to TMV-Gate vectors combined with the flexibility of modulating protein accumulation through altering *A. tumefaciens* cell density and length of infection has utility in experiments involving toxic proteins which when expressed at very high levels may confer detrimental effects on host cells. Recombinant proteins produced using this system can be easily separated from cellular material using anti-epitope tag affinity matrices and be readily detected by either using anti-epitope tag antibodies or by fluorescence microscopy. Such capabilities can be useful for functional genomic studies by defining protein properties, including analysis of protein-protein interactions and subcellular localisation. Functionality of the system for subcellular localisation studies was illustrated using NLS-mCherry which maintained its ability to localise to the nucleus when expressed in fusion with fluorescent proteins. Similarly, AtSAP18 (At2g45640), an established corepressor involved in chromatin remodelling and previously demonstrated to localize to the nucleus⁴¹, when expressed in fusion with CFP using pMW391 reproducibly localised to the nucleus (Supplementary Figure S1), thereby further validating the utility of the TMV-Gate vectors in cell biology applications. It is evident from these results that TMV-Gate driven high-level expres-

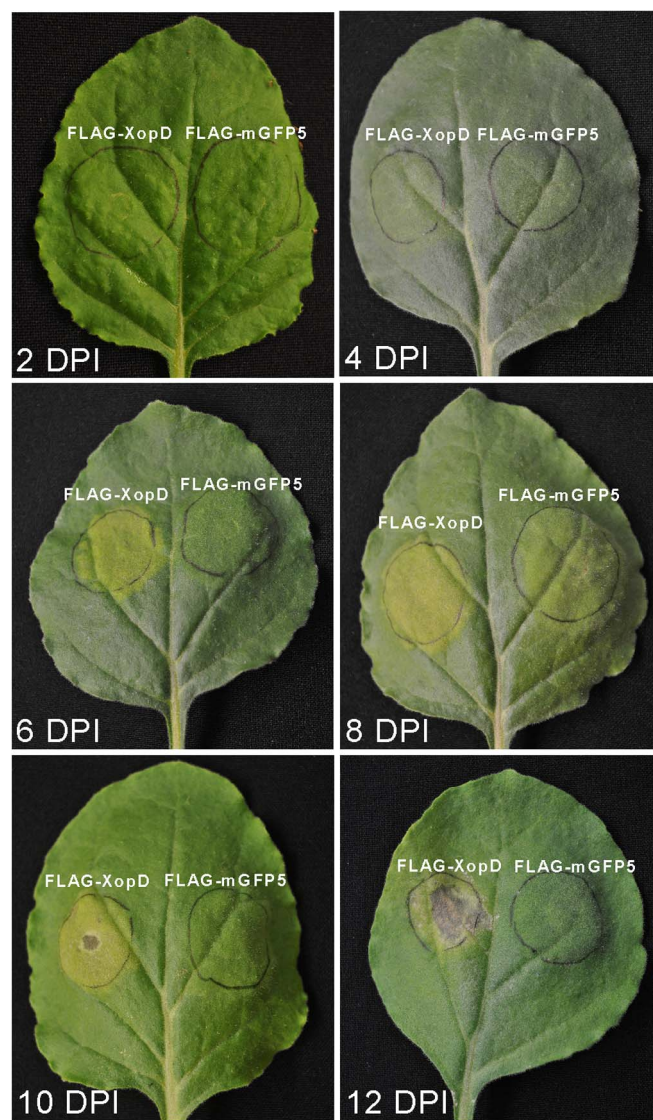


Figure 7 | TMV-Gate vector based expression of XopD effector protein in *N. benthamiana* leaves. *N. benthamiana* leaves were infiltrated with *A. tumefaciens* strains expressing FLAG-tagged XopD (pSU101) or mGFP5 (negative control; pSK145) and the phenotypes were photographed at 2, 4, 6, 8, 10 and 12 DPI.

sion of proteins does not affect their integrity, solubility or localisation. To-date, this system has been used to successfully express 45 native or fusion proteins, including: fluorescent protein, GusPlus, AtDMC1 and XopD variants as listed in Table I (with insert sizes ranging from 741–2358 bp); four transcriptional regulators (654–939 bp; data not shown); a corepressor involved in chromatin remodelling (1422 bp; data not shown); and 17 pathogen effectors (234–1335 bps; data not shown) from *Albugo candida*⁴². In the past, several viral-based expression vector systems have been utilized for functional screening of effector proteins^{43–45}; however, TMV-Gate vectors, by virtue of their ability to produce native as well as various epitope- or fluorescent protein-tagged recombinant proteins, provide a more complete and integrated platform for functional evaluation of effector candidates in *N. benthamiana*. Collectively, this illustrates the robustness and broad utility of the system.

In summary, the vector system described here allows rapid and versatile assembly of expression constructs for production of intact, soluble and functionally active recombinant proteins at a very high yield within a short time-frame and overcomes some limitations of



prokaryotic expression systems, thereby providing an invaluable tool to expedite plant functional genomics and proteomics research. While this study employs plant proteins as examples, it is expected the vector system will be attractive for applications with proteins from animal or other species given the combination of tags incorporated and the potential for rapid, high-level production of heterologous proteins.

Methods

Plant material. *N. benthamiana* plants were grown for 4–6 weeks in a greenhouse at ~25 °C on a 16 h light and 8 h dark cycle before infiltration with *A. tumefaciens* cultures carrying various expression constructs.

Construction of Destination vectors. Cloning techniques were performed using standard protocols⁴⁶. Nucleotide sequences of the primers (Integrated DNA Technologies) used in this study are listed in Supplementary Table S1. Resynthesised DNA cassettes were obtained from the National Research Council (Saskatoon, SK). Gateway reagents and vectors were obtained from Invitrogen. PCR amplifications were performed using PicoMaxx polymerase (Stratagene). All destination vector assemblies as described below were transformed into One Shot ccdB Survival 2 T1 Phage-Resistant *Escherichia coli* cells (Invitrogen), and positive transformants were selected in the presence of kanamycin (50 mg/L) and chloramphenicol (30 mg/L).

Vectors encoding RfA-cassette with or without N-terminal epitope tag(s). The RfA-cassette encoding a chloramphenicol resistance marker and *ccdB* flanked by attR1 and attR2 was amplified by PCR using primers Dest-5'-SnaMfePac and Dest-3'-SphPspAvr. Coding sequences for the RfA-cassette linked to the HA (YPYDVPDYA) or 3xHA (YPYDVPDYAGYPYDVPDYAGSYYPYDVPDYA) epitope tags were amplified by PCR from pEarleyGate201²⁰ using sense primers pEG201-5'-BspPac or pEG201-5'-BspPac+2xHA, respectively, and a common antisense primer DEST-3'AvrPsm. Similarly, the FLAG (DYKDDDDK) or 3xFLAG (DYKDDDDKDYKDDDDKDYKDDDDK) tags linked to the RfA-cassette were amplified from pEarleyGate202²⁰ using the primers pEG202-5'-BspPac or pEG202-5'-BspPac+2xFLAG, respectively, as sense primers and the common antisense primer DEST-3'AvrPsm. To amplify c-myc (EQKLISEEDLN) or 3xc-myc (EQKLISEEDLNGEQKLISEEDLNGEQKLISEEDL) tags linked to the RfA-cassette, pEarleyGate203²⁰ was used as template and the primers pEG203-5'-BspPac or pEG203-5'-BspPac+2xMyc, respectively, were used as sense primers along with the common antisense primer DEST-3'AvrPsm. The 2xStrep-tag II-6xHis-Xpress epitope-linked RfA-cassette was amplified using as template pTK249, a pEXP1-Dest (Invitrogen) derivative encoding a 2xStrep-tag II-6xHis-Xpress epitope-linked RfA-cassette (see Supplementary Methods), and primers Strep-5'-PciNdePac and Exp1-Dest-3'-PspAvr. The PCR amplicons (~1.8 kb) encoding the RfA-cassettes with or without epitope tags from the above PCR reactions were digested with PacI and AvrII then ligated into the TRBO vector digested with the same enzymes resulting in the respective TMV-Gate vectors pMW388, pSK101, pSK102, pSK103, pSK104, pSK105, pSK106 and pTK251, as depicted in Figure 1.

Vector encoding RfA-cassette with C-terminal epitope tag. The 2xStrep-tag II-6xHis epitope assembly at the 3' end of the RfA cassette in pMW394, a pEXP2-Dest (Invitrogen) derivative (see Supplementary Methods), was amplified by PCR using the primers EXP2-Dest-5'-XhoSnaPac and EXP2-Dest-3'-SacPspAvr. The 1.95 kb amplicon was digested with PacI and AvrII then ligated into the TRBO vector digested with the same enzymes, resulting in pMW399 (Figure 1).

Vectors encoding RfA-cassette with C-terminal epitope-tagged fluorescent protein. Plasmids pMW397 and pMW396 encoding assemblies of the RfA-cassette linked to C-terminal YFP or CFP and 3x concatamers of the FLAG or HA tag, respectively, were derived from pEarleyGate101 and pEarleyGate102²⁰ (see Supplementary Methods). The RfA-YFP-3xFLAG and RfA-CFP-3xHA cassettes were isolated by digestion of pMW397 and pMW396 (Supplementary Methods), respectively, with PacI and AvrII then ligated into the TRBO vector digested with the same enzymes, resulting in pMW390 and pMW391 (Figure 1).

Gateway Entry clones. The Gateway Entry clones pMK6, pMW350, pSK6, pSW45, pSU73 and pTK135 (Table I) were assembled as described in the Supporting Information (see Supplementary Methods). These plasmids were digested with PvuI, uniquely cutting the *Km^R* gene, gel-purified and recombined with Destination vectors pSK101-106, pTK251, pMW388, pMW390-391, or pMW399 (Table I) using LR clonase II Invitrogen.

Agroinfection. Expression constructs (Table I) encoding mGFP5, GusPlus, AtDMC1^{G138D}, NLS-mCherry and XopD were introduced into *A. tumefaciens* GV3101::pMP90⁴⁷ by electroporation. Agroinfection of *N. benthamiana* leaves was carried out as described previously¹¹ and summarised in the Supporting Information (see Supplementary Methods).

Visualization of mGFP5 and GusPlus activity. Five days after infiltration of *N. benthamiana* leaves with *A. tumefaciens* carrying various TMV-Gate expression vectors encoding GFP and GusPlus reporter genes, leaf material encompassing the

infiltrated area was excised. Leaves expressing GFP were photographed under UV illumination generated by a transilluminator (VWR Scientific). GusPlus activity was visualized by histochemical staining⁴⁸ with overnight incubation at 37 °C.

Yeast two-hybrid assay. The GAL4-based ProQuest (Invitrogen) Y2H system was used. To assemble the bait and prey constructs, the ORF encoding AtDMC1^{G138D} was transferred from pTK135 to pDEST32 (bait vector) and pDEST22 (prey vector) using the LR clonase reaction, resulting in pTK149 and pTK145, respectively. Yeast strain MaV203 (Invitrogen) was transformed with plasmid combinations listed in Figure 5A according to the method of Gietz and Woods⁴⁹. Assays were conducted on plates containing minimal medium lacking leucine, tryptophan and histidine supplemented with 15 mM 3-AT (Sigma-Aldrich).

Extraction of leaf proteins. Proteins from agroinfected *N. benthamiana* leaves were extracted by grinding tissue in the presence of liquid nitrogen then suspending in 4–5 volumes of extraction buffer [50 mM Tris HCl, pH 8.0, 150 mM NaCl, 1 mM EDTA, 1% Triton-X-100, 1 mM PMSF and 1x plant protease inhibitor mix (Sigma-Aldrich)]. The homogenate was centrifuged twice at 14,000 rpm for 15 min at 4 °C. Protein concentration was determined by the Bradford method⁵⁰ (BioRad).

Experion automated electrophoresis analysis. Samples for Experion Pro260 (BioRad) analysis were prepared by mixing 2 µl of sample buffer with 4 µl samples of protein extracts then loaded onto chips as recommended by the manufacturer. Electrophoresis runs were analyzed using the Experion software (BioRad) as summarised in Supporting Information (Supplementary Methods).

Affinity purification and pull-down experiment. Protein extracts prepared as described above were either incubated with 50 µl (bed volume) of pre-washed EZview red agarose beads (Sigma-Aldrich) conjugated with anti-HA, anti-FLAG or anti-c-myc antibodies, or passed through Strep-Tactin matrix (IBA GmbH). Bound protein was then washed three times with extraction buffer. Tagged proteins affinity captured on the beads were eluted either by boiling the beads in SDS-PAGE sample buffer or, in the case of FLAG-based purification, by mixing with antibody-specific peptide (3xFLAG peptide; Sigma-Aldrich) or for the Strep-Tactin matrix eluted by addition of 2.5 mM desthiobiotin. Eluted proteins were analysed by SDS-PAGE followed by immunoblotting using antibodies specific to each of the epitope tags (see Supplementary Methods).

For pull-down experiments, FLAG or 3xHA-tagged AtDMC1^{G138D} fusions immobilized on anti-FLAG or anti-HA-antibody-conjugated agarose beads, respectively, were washed three times with 500 µl of extraction buffer, then incubated for ~90 min with total protein extract prepared from *N. benthamiana* leaves expressing c-myc- or 3xFLAG-tagged AtDMC1^{G138D}, mGFP5 (negative control) or GusPlus (negative control). The beads were then washed three times with 500 µl of extraction buffer. Proteins retained on the beads were eluted using SDS-PAGE sample buffer and analyzed by SDS-PAGE followed by immunoblotting with anti-c-myc or anti-FLAG antibodies (see Supplementary Methods).

Epifluorescence microscopy. Epifluorescent images of leaf discs expressing NLS-mCherry, NLS-mCherry-YFP-3xFLAG and NLS-mCherry-CFP-3xHA were obtained by using a Zeiss Axio Imager.Z1 microscope equipped with an EC Plan-Neofluar 20X/0.50 M27 objective and an AxioCam HRm camera and using AxioVision software (Carl Zeiss Canada Ltd.). The light source was a broad spectrum X-cite 120 series mercury bulb (Lumen Dynamics Group Inc.). Nuclei were detected by staining with 4',6-diamidino-2-phenylindole (DAPI). Filter sets (Carl Zeiss Canada Ltd.) include HE DsRed (excitation: 538–562; emission: 570–640), YFP (excitation: 490–510; emission: 520–550), HE Cyan (excitation: 424–448; emission: 460–500) and DAPI (excitation: 335–383; emission: 420–470). Exposure times were between 0.2 to 0.9 seconds.

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Author contributions

KR conceived the study. KR and SK designed and coordinated the study, and wrote the manuscript. SK developed and established protocols, and assembled constructs in concert with MW and TB. SK performed transient expression, purification and characterization of recombinant protein experiments. WY performed Y2H experiments. SU and MHB designed and performed pathogen effector screening experiments. All authors read and approved the final manuscript.

Additional information

Supplementary information accompanies this paper at <http://www.nature.com/scientificreports>

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