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#### **RESEARCH PAPER**



# Phosphorylation of TGB1 by protein kinase CK2 promotes barley stripe mosaic virus movement in monocots and dicots

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### Abstract

The barley stripe mosaic virus (BSMV) triple gene block 1 (TGB1) protein is required for virus cell-to-cell movement. However, little information is available about how these activities are regulated by post-translational modifications. In this study, we showed that the BSMV Xinjiang strain TGB1 (<sub>XJ</sub>TGB1) is phosphorylated *in vivo* and *in vitro* by protein kinase CK2 from barley and *Nicotiana benthamiana*. Liquid chromatography tandem mass spectrometry analysis and *in vitro* phosphorylation assays demonstrated that Thr-401 is the major phosphorylation site of the <sub>XJ</sub>TGB1 protein, and suggested that a Thr-395 kinase docking site supports Thr-401 phosphorylation. Substitution of Thr-395 with alanine (T395A) only moderately impaired virus cell-to-cell movement and systemic infection. In contrast, the Thr-401 alanine (T401A) virus mutant was unable to systemically infect *N. benthamiana* but had only minor effects in monocot hosts. Substitution of Thr-395 or Thr-401 with aspartic acid interfered with monocot and dicot cell-to-cell movement and the plants failed to develop systemic infections. However, virus derivatives with single glutamic acid substitutions at Thr-395 and Thr-401 developed nearly normal systemic infections in the monocot hosts but were unable to infect *N. benthamiana* systemically, and none of the double mutants was able to infect dicot and monocot hosts. The mutant x<sub>J</sub>TGB1<sub>T395A/T401A</sub> weakened *in vitro* interactions between x<sub>J</sub>TGB1 and x<sub>J</sub>TGB3 proteins but had little effect on x<sub>J</sub>TGB1 RNA-binding ability. Taken together, our results support a critical role of CK2 phosphorylation in the movement of BSMV in monocots and dicots, and provide new insights into the roles of phosphorylation in TGB protein functions.

Key words: Barley stripe mosaic virus, triple gene block 1 (TGB1) protein, phosphorylation, protein kinase CK2, promotion, viral movement.

### Introduction

*Barley stripe mosaic virus* is the type species of the genus *Hordeivirus*. Barley stripe mosaic virus (BSMV) infects barley, wheat, and oats under natural conditions, and numerous

other monocots and dicots by artificial inoculation (Bragg *et al.*, 2008; Jackson *et al.*, 2009). BSMV has recently been isolated from 750-year-old barley grains found near the Nile

Abbreviations: BSMV, barley stripe mosaic virus; CP, coat protein; DIG, digoxygenin; dpi, d post-inoculation; EMSA, electrophoretic mobility shift assays; GFP, green fluorescent protein; LC-MS/MS, liquid chromatography tandem mass spectrometry; MP, movement protein; ORF, open reading frame; PD, plasmodesmata; PSLV, Poa semilatent virus; RdRp, RNA-dependent RNA polymerase; RFP, red fluorescent protein; RT-PCR, reverse transcription-PCR; TGB, triple gene block; TMV, tobacco mosaic virus.

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River (Smith *et al.*, 2014), and consists of a large number of strains that were early subjects of phenotypic and hostrange studies (McKinney and Greeley, 1965). BSMV has a positive-sense tripartite RNA genome (RNA $\alpha$ , - $\beta$ , and - $\gamma$ ) that encodes seven major proteins (Bragg *et al.*, 2008; Jackson *et al.*, 2009). RNA $\alpha$  directs synthesis of the  $\alpha$ a protein, which functions as the methyltransferase/helicase subunit of the replicase [RNA-dependent RNA polymerase (RdRp)]. RNA $\beta$ encodes  $\beta$ a [coat protein (CP)], and an overlapping triple gene block (TGB) sequence encoding three major movement proteins (TGB1, TGB2, and TGB3) that are expressed from two subgenomic RNAs. RNA $\gamma$  serves as the mRNA for the  $\gamma$ a RdRp polymerase subunit, and the  $\gamma$ b protein, which functions as a suppressor of RNA silencing and a modulator of host defences (Jackson *et al.*, 2009).

The BSMV TGB1 protein is a multifunctional 58 kDa protein with RNA-binding, RNA helicase, and ATPase activities (Donald et al., 1997; Verchot-Lubicz et al., 2010). The C-terminal region contains seven conserved helicase motifs, and mutations within one or more of these motifs have been shown to be involved in enzymatic and movement functions and RNA-binding activities (Lawrence and Jackson, 2001; Lim et al., 2008). TGB2 and TGB3 are trans-membrane proteins that are integrated in the endoplasmic reticulum bilayer (Morozov and Solovyev, 2003). Cell-to-cell movement of BSMV does not require the CP, and this property has permitted isolation of nucleoprotein complexes composed of the TGB1 protein and viral genomic and subgenomic RNAs (Lim et al., 2008). The TGB1 protein interacts directly with the TGB3 protein and indirectly with the TGB2 protein to form heterologous complexes required for co-localization of the TGB proteins at the plasmodesmata (PD) and BSMV cell-tocell movement (Lawrence and Jackson, 2001; Lim et al., 2008; Jackson et al., 2009; Lim et al., 2009). Our recent results have shown that TGB1 proteins function in eliciting resistance to BSMV strains that are unable to infect Brachypodium distach*yon* inbred lines containing the *Bsr1* resistance gene. In the case of the North Dakota 18 (ND18) strain TGB1 protein, amino acid residues at positions 390 and 392 are critical for TGB1 protein genetic interactions with *Bsr1* and for inducing resistance responses (Cui et al., 2012; Lee et al., 2012).

Although the studies mentioned above provide valuable insights into BSMV cell-to-cell movement processes, little information is available about post-translational biochemical events, such as phosphorylation, that may function in regulating intercellular macromolecular trafficking. Movement protein (MP) phosphorylation was first demonstrated to be important for virus cell-to-cell movement during investigations with the tobacco mosaic virus (TMV) 30kDa MP, and this study provided important approaches for subsequent MP analyses (Citovsky et al., 1993). Several proteins may function in TMV movement, because a cell-wall-associated kinase (Citovsky et al., 1993), a PD-associated protein kinase (PAPK1) (Lee *et al.*, 2005), an endoplasmic reticulum-associated kinase (Karger et al., 2003), and protein kinase CK2 (formerly known as casein kinase II; Ivanov et al., 2003) have been shown to be involved in in vitro and in vivo phosphorylation of the 30kDa protein (Citovsky et al., 1993). Moreover, mimicking MP phosphorylation by negatively charged amino acids inhibited MP transport through PD and delayed TMV and potyvirus spread in *Nicotiana tabacum* (Waigmann *et al.*, 2000; Karger *et al.*, 2003). More recently, phosphorylation of the MPs of other viruses, such as tomato mosaic virus (Kawakami *et al.*, 2003; Matsushita *et al.*, 2003), potato leafroll virus (Link *et al.*, 2011), Abutilon mosaic virus (Kleinow *et al.*, 2009), brome mosaic virus (Akamatsu *et al.*, 2007), apple chlorotic leaf spot virus (Sato *et al.*, 1995), and cucumber mosaic virus (Matsushita *et al.*, 2002), has been shown to either enhance or inhibit virus movement during infection.

Although little information is available about the roles of phosphorylation in the movement processes of TGB MPs, potato virus X TGB1 protein is efficiently phosphorylated by N. tabacum protein kinase CK2 (Módena et al., 2008). Furthermore, tyrosine phosphorylation regulates the functions of potato mop-top virus TGB3 protein because substitution of tyrosine residues in two phosphorylation domains enhances interactions between the TGB3 and TGB2 proteins and inhibits virus cell-to-cell movement (Samuilova et al., 2013). In addition, the N-terminal half of the TGB1 protein of Poa semilatent virus (PSLV), a hordeivirus closely related to BSMV, has been reported to be phosphorylated in vitro by casein kinase 1 (CK1), protein kinase A (PKA), and protein kinase C (PKC)-like kinases present in N. benthamiana cellwall fractions. In the case of PSLV, phosphorylation of an internal domain decreases RNA-binding activity and homologous protein-protein interactions, but experiments to determine whether these activities affect movement have not been conducted (Makarov et al., 2012). Here, we present the first evidence that BSMV TGB1 protein is phosphorylated in vitro and in vivo by the host protein kinase CK2. Our biochemical and molecular approaches demonstrated that Thr-401 in the TGB1 C-terminal region is a major phosphorylation site of the TGB1 protein, and that the Thr-395 residue serves as a CK2 docking domain. Mutational analyses of these residues indicate that a phosphorylation-dependent mechanism is involved in BSMV local and systemic infections in monocots and dicots.

#### Materials and methods

#### Plant growth conditions

*N. benthamiana* plants were grown in a climate chamber at 23 to 25 °C with a 14/10h light (~75 mmol m<sup>-2</sup> s<sup>-1</sup>)/dark photoperiod as described previously (Yuan *et al.*, 2011). Barley (Yangpi 8), wheat (Yangmai 11), and *B. distachyon* Bd21 plants were grown in a greenhouse until the two-leaf stage, and then inoculated and transferred to a climate chamber at the same temperature and light regimen as above until evaluated at 5 to 12 d post-inoculation (dpi).

# Construction of infectious clones of BSMV Xinjiang $(x_{,i}BSMV)$ strain

Genomic (g) RNAs of <sub>XJ</sub>BSMV strain were extracted from purified virus with Trizol (Life Technologies) and used to prime reverse transcription of the gRNAs with primer BS32 as described previously (Yuan *et al.*, 2011; Lee *et al.*, 2012). BSMV  $\alpha$ ,  $\beta$ , and  $\gamma$  cDNAs were amplified with the primer pairs XJ-1/BS32, XJ-2/BS32, and XJ-3/BS32, respectively (Supplementary Table S1, available at *JXB* online), and inserted into the pMD20-T vector (Takara) to generate pT7- $\alpha_{XJ}$ , pT7- $\beta_{XJ}$ , and pT7- $\gamma_{XJ}$ . Site-specific mutagenesis was carried out with a QuikChange Site-Directed Mutagenesis kit (Agilent Technologies) to make alanine (A), aspartic acid (D), and glutamic acid (E) substitutions for <sub>XJ</sub>TGB1 protein residues 395 and 401 in pT7- $\beta_{XJ}$  with the corresponding primer pairs (Supplementary Table S2, available at *JXB* online). These clones and all those described below were verified by DNA sequencing (Tsingke Biotech, Beijing).

To engineer <sub>XJ</sub>BSMV derivatives for agroinfiltration, full-length cDNAs were amplified from pT7- $\alpha_{XJ}$ , pT7- $\beta_{XJ}$  (or the site-specific pT7- $\beta_{XJ}$  TGB1 mutants), and pT7- $\gamma_{XJ}$  clones with the primer pairs CH-10/BS-26, CH-11/BS-26, and CH-12/BS-26, respectively (Supplementary Table S1). The cDNAs then were inserted between the *Stu*I and *Bam*HI sites of pCass4-Rz (Annamalai and Rao, 2005), and the resulting clones and site-specific mutants with the 395 and 401 residue substitutions were designated pCa- $\alpha_{XJ}$ , pCa- $\beta_{XJ}$ , rGB1 mutants.

# Mechanical inoculation of in vitro transcripts and agroinfiltration of BSMV derivatives

The pT7- $\alpha_{XJ}$ , pT7- $\beta_{XJ}$ , and pT7- $\gamma_{XJ}$  plasmids were linearized with *SpeI* or *Bam*HI and used as templates for T7 RNA polymerase (Promega) *in vitro* transcription of capped infectious RNAs (Petty *et al.*, 1989). The  $\alpha$ ,  $\beta$ , and  $\gamma$  *in vitro* transcripts were mixed in equal amounts with FES inoculation buffer (0.1 M glycine, 0.06 M potassium phosphate, 1% sodium pyrophosphate decahydrate, 1% bentonite, 1% celite, pH 8.5) and used immediately for inoculation of 7- to 10-d-old barley, wheat, and *B. distachyon* Bd21. Plasmids pCa- $\alpha_{XJ}$ , pCa- $\beta_{XJ}$ , and pCa- $\gamma_{XJ}$  were maintained in *Agrobacterium tumefaciens* strain EHA105 and infiltrated into the lower side of *N. benthamiana* leaves as described previously (Yuan *et al.*, 2011).

#### Immunoprecipitation

Immunoprecipitation assays were carried out with minor modifications of a published protocol (Rubio *et al.*, 2005). *N. benthamiana* leaf sections were harvested at 5 d after agroinfiltration and proteins were extracted in 2 vols (w/v) of GTEN buffer [10% (v/v) glycerol, 25mM Tris/HCl (pH 7.5), 1mM EDTA, 150mM NaCl, 10mM dithiothreitol, 2% (w/v) polyvinylpolypyrrolidone, 1% Protease Inhibitor Cocktail (Roche)]. Extracted complexes were stirred for 30min and centrifuged at 12 000g for 30min at 4 °C. The supernatants were first incubated with TGB1 protein antibody at 4 °C for 6h, and then mixed with protein G–agarose (Millipore) beads to enrich the <sub>XJ</sub>TGB1 protein. The immunoprecipitation products were washed five times with immunoprecipitation buffer [GTEN buffer with 0.15% (v/v) NP-40, 0.5mM dithiothreitol] and evaluated by Western blotting with anti-TGB1 and anti-phosphothreonine antibody ( $\alpha$ -pT; Millipore) at 1:500 dilutions.

# Construction of protein expression vectors for in vitro phosphorylation, glutathione S-transferase (GST), and Histagged pull-down assays

To engineer  $_{XJ}$ TGB1 C-terminal His-tagged ( $_{XJ}$ TGB1-6His) fusion proteins, the  $_{XJ}$ TGB1 open reading frame (ORF) was amplified from the plasmid pT7- $\beta_{XJ}$  with the primer pair TGB1-NdeIF/TGB1-XhoIR (Supplementary Table S1), and integrated into the *NdeI* and *XhoI* sites of the pET-30a(+) expression vector (Novagen) to generate pET- $_{XJ}$ TGB1. The TGB1 mutant derivatives of the Thr-395 and Thr-401 residues were engineered using a QuikChange Site-Directed Mutagenesis kit with corresponding primer pairs for the TGB1 ORFs (Supplementary Table S2).

The  $\alpha$ -catalytic subunit of CK2 from *N. benthamiana* (NbCK2 $\alpha$ , GenBank accession no. KJ748371) (Bombarely *et al.*, 2012) and barley (HvCK2 $\alpha$ , GenBank accession no. AB252049) (Kato *et al.*, 2008)

was isolated from host RNA by reverse transcription (RT)-PCR amplification with the primer pairs NbCK2-NdeIF/NbCK2-XhoIR and HvCK2 $\alpha$ -NdeIF/HvCK2 $\alpha$ -SalIR, respectively (Supplementary Table S1). Primers for amplifying NbCK2 $\alpha$  were based on the flanking sequence of the *N. tabacum* CK2 $\alpha$  ORF (GenBank accession no. AF374474) (Salinas *et al.*, 2001). The PCR products of the NbCK2 $\alpha$  and HvCK2 $\alpha$  genes were inserted into the *NdeI/XhoI*, and *NdeI/SalI* sites of pET-30a(+), respectively, to generate pET-NbCK2 $\alpha$  and pET-HvCK2 $\alpha$ .

For GST pull-down of <sub>XJ</sub>TGB1 and <sub>XJ</sub>TGB3 proteins, the <sub>XJ</sub>TGB3 ORF was amplified via PCR and ligated in frame to the C terminus of the GST ORF at the *Bam*HI and *Xho*I restriction sites of the plasmid pGEX-KG (Guan and Dixon, 1991) to yield the pGEX-<sub>XJ</sub>TGB3 plasmid.

# Expression and purification of recombinant proteins from Escherichia coli *BL21* cells

The pET plasmids designed for expression of the C-terminal Histagged <sub>XJ</sub>TGB1 proteins, the <sub>XJ</sub>TGB1 protein mutants, and the HvCK2 $\alpha$  and NbCK2 $\alpha$  proteins were transformed into *E. coli* strain BL21(DE3) by standard procedures (Sambrook and Russell, 2001). Each transformant was grown overnight at 37 °C in 3 ml of LB medium containing kanamycin (100 µg ml<sup>-1</sup>), transferred into 1 litre of fresh LB/kanamycin medium and grown to a density of 0.4 OD<sub>600</sub>. Recombinant protein expression was induced by the addition of 0.2 mM isopropyl β-D-1-thiogalactopyranoside and the cells were shaken for 16 additional hours at 18 °C. Recombinant proteins were extracted from the bacteria and purified by Ni-affinity chromatography according to the manufacturer's instructions (Bio-Rad) and evaluated by 12.5% SDS-PAGE.

#### In vitro phosphorylation assays

Soluble cytoplasmic protein extracts of healthy N. benthamiana leaves were used for in vitro kinase assays according to the protocol described by Hung et al. (2014) and Vijayapalani et al. (2012). Phosphorylation reactions were performed with the N. benthamiana soluble protein extracts or with purified recombinant NbCK2a and HvCK2a subunits. Assays were performed with 1 µg of N. bentha*miana* soluble protein extracts, or 0.1  $\mu$ g of recombinant CK2 $\alpha$ , and 1 µg of purified xITGB1 protein or its mutants in a final volume of 10 µl of 25 mM Tris/HCl (pH 7.4), 10 mM MgCl<sub>2</sub>, and 1 µl [γ-<sup>32</sup>P]ATP or GTP (10 µCi, ~3000 Ci mmol<sup>-1</sup>; Perkin Elmer). Selected reactions were carried out in the presence or absence of heparin, and various concentrations of unlabelled ATP or GTP. Negative controls contained no TGB1 protein or 500 and 1000 ng of bovine serum albumin. After incubation at 30 °C for 30 min, the reactions were terminated by addition of 2.5  $\mu$ l of 5× SDS buffer, and the samples were subjected to 12.5% SDS-PAGE. The gels were dried with a Model 583 Gel Dryer (Bio-Rad) and phosphorylated proteins were visualized by autoradiography.

#### Mass spectrometry analysis

Phosphorylated <sub>XJ</sub>TGB1 and unphosphorylated <sub>XJ</sub>TGB1 proteins were digested with trypsin at 37 °C overnight. The digested peptides were then analysed by Q-Exactive liquid chromatography tandem mass spectrometry (LC-MS/MS) (Thermo Scientific) at the Mass Spectrometry Facility at China Agricultural University. The data were searched against the NCBI database using Mascot software with a 1% false discovery rate.

#### Fluorescence and confocal microscopy

Green fluorescent protein (GFP) or red fluorescent protein (RFP) fluorescence in epidermal cells of *N. benthamiana* was observed with an Olympus confocol FV1000 microscope. GFP and RFP were excited at 488 or 546 nm, respectively, with an argon laser. Images

were recorded with an Olympus camera and processed using an Olympus Fluoview version 3.0 Viewer. In addition, cell-to-cell movement assays in epidermal cells of barley and *N. benthamiana* were observed with a BX53+DP72 fluorescence microscope (Olympus) and images were manipulated with the cellSens Entry programs.

#### Electrophoretic mobility shift assays (EMSA)

RNAs for binding assays were transcribed in vitro in the presence of digoxigenin (DIG)-UTP (Roche) and the DIG-labelled transcripts were purified to remove the DNA templates (Wu et al., 2013). Phosphorylation reactions were performed in a final volume of  $5 \mu l$ containing phosphorylation assay buffer and purified NbCK2a, and different amounts of recombinant xJTGB1 protein as described above. Negative controls consisted of samples lacking recombinant xITGB1 protein or 500 and 1000 ng of bovine serum albumin. EMSA binding comparisons were performed by adding increasing amounts of protein to 50 ng of purified RNA in binding buffer [50 mM Tris/ HCl (pH 7.5), 10mM MgCl<sub>2</sub>, 1mM EDTA] in a final volume of 20 µl. The binding reaction mixtures were incubated on ice for 30 min and subjected to electrophoresis on 1% (w/v) non-denaturing agarose gels in 0.5× TBE buffer. RNA-protein complexes were transferred to a Hybond N<sup>+</sup> nylon membrane (GE Healthcare) via a pump suction filter, and the RNA was cross-linked to the membrane by two 60 s cycles at 0.12 J in a Bio-Link crosslinker (Vilber Lourmat). Mobility shifts of the DIG-labelled RNAs were detected with DIG- alkaline phosphatasealkaline phosphatase Fab fragments (Roche), and the blots were developed with a 5-bromo-4-chloro-3-indolyl-phosphate/ nitro blue tetrazolium chloride substrate solution (Amresco).

#### GST pull-down

For co-expression of  $GST_{XJ}TGB3$  with  $_{XJ}TGB1$ –His or the  $_{XJ}TGB1_{T395A/T401A}$ –His fusion proteins, relevant plasmids were co-transformed into *E. coli* BL21(DE3). Cells were harvested by low-speed centrifugation and disrupted by vortexing in TB buffer [20 mM Tris/HCl (pH 7.3), 500 mM NaCl] in the presence of glass beads. The GST fusions and bound TGB proteins were purified by glutathione–Sepharose affinity chromatography and elution with glutathione (Pharmacia).

#### Results

#### Construction and sequence analysis of infectious clones of the BSMV Xinjiang strain

Several BSMV field strains from China collected in our laboratory have broader host ranges than the more extensively studied NDBSMV and Type BSMV (TYBSMV) strains. To evaluate the diversity of the more virulent BSMV strains (Lee et al., 2012), we constructed infectious clones of the  $_{XJ}BSMV$  strain (Xie et al., 1981) under the bacteriophage T7 or double cauliflower mosaic virus 35S promoters (Petty et al., 1989; Yuan et al., 2011) (Fig. 1A). The infectivity of in vitro transcripts synthesized from linearized pT7- $\alpha_{XJ}$ , pT7- $\beta_{XJ}$ , and pT7- $\gamma_{XJ}$  plasmids was tested by mechanical inoculation to barley, wheat, and B. distachyon Bd21. Inoculated plants consistently developed chlorotic stripes and mosaic symptoms typical of BSMV infections on upper uninoculated leaves by 6-7 dpi (Fig. 1B) and the efficiency of infectivity in barley, wheat, and B. distachyon Bd21 was 70-80, 80-90, and 50-60%, respectively (also see Supplementary Table S6). Agroinfiltration was used to initiate infections of N. benthamiana because only 10-30% of the plants became infected when using in vitro transcripts as inocula. Agrobacterium harbouring the plasmids pCa- $\alpha_{XJ}$ , pCa- $\beta_{XJ}$ , and pCa- $\gamma_{XJ}$  were infiltrated into the basal sides of the leaves. Newly emerging leaves developed mild mottling symptoms at 7–9 d after agroinfiltration (Fig. 1C), and the efficiency of infectivity was increased to approximate 90%. RT-PCR and Western blot analysis verified the infectivity of the <sub>XJ</sub>BSMV infectious clones in the monocot hosts (Fig. 1B) and in *N. benthamiana* (Fig. 1C).

For comparisons of <sub>XJ</sub>BSMV genomic variation with other published BSMV strains, the pT7- $\alpha_{XJ}$ , pT7- $\beta_{XJ}$  and pT7- $\gamma_{XJ}$ cDNAs were sequenced. The results indicated that <sub>XJ</sub>RNA $\alpha$ (GenBank accession no. KJ746471), <sub>XJ</sub>RNA $\beta$  (GenBank accession no. KJ746472), and <sub>XJ</sub>RNA $\gamma$  (GenBank accession no. KJ746473) consisted of 3789, 3222, and 2793 nt, respectively, and shared nucleotide identities of 95.2–99.8% (Supplementary Table S3, available at *JXB* online), 94.1–99.2% (Supplementary Table S4, available at *JXB* online), and 87.9– 98.8% (Supplementary Table S5, available at *JXB* online) with RNAs  $\alpha$ ,  $\beta$ , and  $\gamma$  of other BSMV strains. These results suggest that substantial diversity may exist among BSMV strains.

# The $_{XJ}TGB1$ protein is phosphorylated in vitro and in vivo

In order to explore phosphorylation *in vitro*, the full-length <sub>XJ</sub>TGB1 protein was expressed as a C-terminal His-tagged



**Fig. 1.** Diagram of <sub>XJ</sub>BSMV strain infectious clones and cereal and dicot host infectivity test results. (A) Illustration of <sub>XJ</sub>BSMV infectious clones under the T7 promoter or double cauliflower mosaic virus 35S promoter as described previously (Yuan *et al.*, 2011; Lee *et al.*, 2012). (B) Infectivity assays with *in vitro*-synthesized gRNAs of barley, wheat, and *B. distachyon* Bd21. (C) Agroinfiltration was used to initiate infections of *N. benthamiana*. Typical chlorotic stripes and mosaic symptoms (top) of BSMV appeared on emerging uninoculated leaves by 7–9 dpi. Upper uninoculated leaf tissue was harvested at 12 dpi, and the relative BSMV RNA and CP amounts were evaluated by RT-PCR (middle) and Western blotting with the antibody against BSMV CP (bottom). BSMV RNAγ was detected by RT-PCR with the primer pair BS-10/BS-32 (Supplementary Table S1). (This figure is available in colour at *JXB* online.)

fusion protein and purified to near homogeneity by Ni-affinity chromatography (Fig. 2A). A soluble protein kinase is known to be present in tobacco species (Hung et al., 2014), and hence the purified xITGB1 protein was first assayed for phosphorylation using N. benthamiana protein extracts as a kinase source. The first control reaction containing  $[\gamma^{-32}P]ATP$  and cytoplasmic extracts without the xITGB1 protein resulted in no distinct labelled products (Fig. 2B, lane 1). The corresponding control with the x<sub>J</sub>TGB1 protein and  $[\gamma^{-32}P]ATP$ alone suggested that the xITGB1 protein was not autophosphorylated in vitro (Fig. 2B, lane 2). In contrast, when both the cytoplasmic extracts and the XITGB1 protein were present, a radioactive phosphorylated product co-migrated with the XJTGB1 protein (Fig. 2B, lane 3). These results provide evidence that the xJTGB1 protein is phosphorylated in vitro with  $[\gamma^{-32}P]$ ATP by a soluble kinase in the *N*. benthamiana extracts. The vast majority of protein kinases use ATP as an exclusive



Fig. 2. Phosphorylation of the xITGB1 protein in vitro and in vivo. (A) Coomasie Brilliant Blue (CBB) staining of recombinant xJTGB1 protein purified from E. coli cells. Molecular weight markers (Fermentas) are indicated on the left side of the gel. (B) In vitro phosphorylation of purified xJTGB1 protein by cellular kinases present in healthy N. benthamiana extracts in the absence or presence of  $[\gamma^{-32}P]ATP$  or  $[\gamma^{-32}P]GTP$ . After the phosphorylation reactions, the TGB1 proteins were separated by 12.5% SDS-PAGE and the incorporated radioactivity was analysed by autoradiography. Reaction mixtures lacking x, TGB1 protein or N. benthamiana protein extracts served as negative controls. The CBB staining in the lower panel indicates that similar amounts of the xJTGB1 protein were present in each in vitro phosphorylation reaction. (C) In vivo phosphorylation of xJTGB1 protein in N. benthamiana by Western blotting with  $\alpha$ -TGB1 polyclonal antibodies and  $\alpha$ -threonine antibodies. A mock agroinfiltration lacking xIRNAB was used as a negative control and molecular weight markers (Thermo Scientific) were used to estimate the size of the xJTGB1 protein. (D) In vivo phosphorylation of xJTGB1 protein immunoprecipitated (IP) from N. benthamiana was analysed as in Fig. 2C. (This figure is available in colour at JXB online.)

phosphate donor, whereas CK2 can effectively use either ATP or GTP (Matsushita *et al.*, 2000); hence we carried out phosphorylation comparisons with GTP to obtain clues about the identity of the kinase involved in <sub>XJ</sub>TGB1 phosphorylation. Autoradiography of the phosphorylated products revealed a single intense radiolabelled band when either  $[\gamma^{-32}P]$ ATP or  $[\gamma^{-32}P]$ GTP was used as a phosphoryl donor (Fig. 2B, lanes 3 and 4), implying that <sub>XJ</sub>TGB1 is phosphorylated by a CK2-like kinase in *N. benthamiana*.

To determine whether the x<sub>J</sub>TGB1 protein is phosphorylated in vivo, XIBSMV-infiltrated N. benthamiana leaves were harvested, concentrated by immunoprecipitation, and subjected to Western blot analysis with an anti-phosphothreonine ( $\alpha$ -pT) antibody. The  $\alpha$ -pT results revealed a labeleld protein with an electrophoretic mobility corresponding to the 58 kDa <sub>XJ</sub>TGB1 protein (Fig. 2C and D, top, lane 3). Similarsized bands were not observed in mock-inoculated leaves, or leaves infiltrated with only pCa- $\alpha_{XJ}$  and pCa- $\gamma_{XJ}$  (Fig. 2C and D, top, lanes 1 and 2). Western blot analyses using the <sub>XI</sub>TGB1 protein antibody confirmed that the  $\alpha$ -pT-reacting protein co-migrated with xITGB1, and revealed a 45kDa immunospecific band that we suspect is a degradation product (Fig. 2C and D, bottom, lane 3). Collectively, these experiments provide convincing evidence that the XITGB1 protein is phosphorylated by a soluble CK2-like kinase in vitro and in vivo.

#### CK2 is able to phosphorylate the <sub>x,1</sub>TGB1 protein

To predict potential xITGB1 protein phosphorylation kinases and sites, we analysed the xITGB1 protein sequence with the GPS 2.1 program tool (Xue et al., 2011) and the Scansite Motif Scanner online server (http://scansite.mit.edu) (Obenauer, 2003). The GPS 2.1 program was set at a HIGH threshold (false-positive rates of 2% for Ser/Thr kinases) and the Scansite was set at a high stringency level to predict potential phosphorylation targets in the xJTGB1 protein. According to the conserved CK2 phosphorylation site motif [(S/T)-X-X-(D/E); Meggio and Pinna, 2003], the GPS 2.1 predictions suggested that three residues in the xITGB1 protein (Ser-69, Thr-395, and Thr-401) are potential CK2 phosphorylation sites, whereas the Scanner only indicated that the Thr-401 site is phosphorylated by CK2 (Supplementary Fig. S1, available at JXB online). For comparison with ND18, the x1TGB1 amino acids Ser-69, Thr-395, and Thr-401 correspond to the ND18 TGB1 (NDTGB1) Ser-70, Thr-396, and Thr-402 residues, respectively.

Based on the GTP results and phosphorylation predictions, we suspected that a CK2-like kinase might be responsible for phosphorylation of the <sub>XJ</sub>TGB1 protein. Therefore, the CK2 $\alpha$  subunits from *N. benthamiana* (NbCK2 $\alpha$ ) and barley (HvCK2 $\alpha$ ) were cloned, and the C-terminal His-tagged fusion proteins were purified from *E. coli* cells (Fig. 3A). Subsequent *in vitro* phosphorylation assays revealed that the purified NbCK2 $\alpha$  and HvCK2 $\alpha$  proteins efficiently phosphorylated the recombinant <sub>XJ</sub>TGB1 protein in the presence of [ $\gamma$ -<sup>32</sup>P] ATP (Fig. 3B). CK2 is highly sensitive to heparin inhibition (Matsushita *et al.*, 2003), and our results confirmed that the



**Fig. 3.** *In vitro* phosphorylation of  $_{XJ}$ TGB1 protein by recombinant CK2 kinase. (A) SDS-PAGE analysis of NbCK2 $\alpha$  and HvCK2 $\alpha$  purified from *E. coli* BL21 cells. (B) *In vitro* phosphorylation of  $_{XJ}$ TGB1 protein with the NbCK2 $\alpha$  and HvCK2 $\alpha$  recombinant proteins and negative controls lacking the kinases. (C) Effects of heparin on *in vitro* phosphorylation of  $_{XJ}$ TGB1 protein. Phosphorylation levels were reduced with increasing amount of heparin. (D) Ability of NbCK2 $\alpha$  to use both ATP and GTP as phosphate donors. (E) Divalent metal ion specificity of NbCK2 $\alpha$  and the TMV-MP (P30) proteins. The CBB-stained proteins at the bottom of panels (B)–(E) are as indicated as in Fig. 2B. (F) Co-localization of the GFP:<sub>XJ</sub>TGB1 and DsRed:NbCK2 $\alpha$  proteins in *N. benthamiana* leaf cells. Single localization of GFP:<sub>XJ</sub>TGB1 and DsRed:NbCK2 $\alpha$  proteins are indicated at the top of the panels. Bars, 50 µm. (This figure is available in colour at *JXB* online.)

levels of phosphorylation were reduced proportionally with increasing heparin concentrations (Fig. 3C). To further test the kinase specificity, we used NbCK2 $\alpha$  to compare <sub>XJ</sub>TGB1 and TMV P30 MP (Ivanov *et al.*, 2003) protein phosphorylation in the presence of [ $\gamma$ -<sup>32</sup>P]ATP and [ $\gamma$ -<sup>32</sup>P]GTP (Fig. 3D). In both cases, the NbCK2 $\alpha$  phosphorylation assays resulted in the presence of highly intense bands that co-migrated with the <sub>XJ</sub>TGB1 and P30 proteins, and the kinase exhibited similar

activities in the presence of both ATP and GTP (Fig. 3D, lanes 3 and 4, and 6 and 7). Moreover, the absence of radioactive bands in reactions lacking the NbCK2 $\alpha$  protein confirmed that the 58 kDa <sub>XJ</sub>TGB1 and the P30 proteins are not autophosphorylated and the lack of radioactivity in the reactions without substrate proteins also indicated that the NbCK2 $\alpha$  protein is not self-phosphorylated (Fig. 3D, lanes 2 and 5). In addition, tests were carried out in the presence of  $Mn^{2+}$ ,  $Mg^{2+}$ , and  $Ca^{2+}$  to assess the cation specificity of CK2 phosphorylation (Niefind *et al.*, 1999). These results revealed that NbCK2 $\alpha$  exhibited similar phosphorylation intensities for <sub>XJ</sub>TGB1 and P30 in the presence of either Mn<sup>2+</sup> or Mg<sup>2+</sup> (Fig. 3E, lanes 2 and 3, and 7 and 8), and that <sup>32</sup>P incorporation was negligible in reactions containing Ca<sup>2+</sup> (Fig. 3E, lanes 4 and 9). All of the *in vitro* phosphorylation data with the TGB1 protein were consistent with several published biochemical properties of CK2 (Ivanov *et al.*, 2003; Hung *et al.*, 2014).

Furthermore, to explore the potential co-localization of the  $_{XJ}TGB1$  protein and NbCK2 $\alpha$  *in planta*, we conducted co-expression experiments with GFP:<sub>XJ</sub>TGB1 and DsRed:NbCK2 $\alpha$  fusion proteins in *N. benthamiana* cells. Confocal microscopy revealed that GFP:<sub>XJ</sub>TGB1 was distributed in both the cytoplasm and nucleus, whereas DsRed:NbCK2 $\alpha$  was present primarily in the nucleus. The co-localization of the two proteins indicated that the <sub>XJ</sub>TGB1 protein and NbCK2 $\alpha$  interact in both the nucleus and cytoplasm in some manner (Fig. 3F, overlay channel). Taken together, the presented data demonstrate that XJTGB1 is phosphorylated by CK2 *in vitro* and *in planta*.

# Thr-401 is the major $_{XJ}$ TGB1 protein site for CK2 phosphorylation

To identify the phosphorylation sites of CK2, the purified  $_{XJ}TGB1$  protein was phosphorylated by NbCK2 $\alpha$  *in vitro* with unlabelled ATP, and the gel-purified phosphorylated and unphosphorylated  $_{XJ}TGB1$  proteins were separated by PAGE and digested with trypsin. The trypsin digestion products were analysed by Q-Exactive LC-MS/MS. The analysis showed that 87.9% of the TGB1 protein amino acid sequence was covered, and revealed that the phosphorylated and unphosphorylated proteins differed in a unique monophosphorylated peptide (<sup>399</sup>GETDETEKNIAFTVDTVR<sup>416</sup>) with a 2103.9362 *m*/*z* peak corresponding to a neutral precursor ion lacking phosphoric acid (97.9769 Da). Based on the observed masses of the phosphorylated and unphosphorylated soft the phosphorylated and unphosphorylated  $_{Y16}$  fragment ions (Fig. 4A), we conclude that the phosphorylated  $_{X1}$ TGB1 residue is located at Thr-401.

To verify the GPS 2.1 and Scanner predictions (Supplementary Fig. S1) and LC-MS/MS analysis of x1TGB1 protein, we replaced one or both of the Thr-395 and Thr-401 residues with alanine residues to produce  $_{XI}TGB1_{T395A}$ , <sub>XJ</sub>TGB1<sub>T401A</sub>, and <sub>XJ</sub>TGB1<sub>T395A/T401A</sub> phosphorylation-deficient mutants. To mimic the phosphorylation state of the XJTGB1 protein, Thr-395 and Thr-401 residues were substituted with aspartic acid (D) or glutamic acid (E) residues to produce the XJTGB1T395D, XJTGB1T395E, XJTGB1T401D, and XJTGB1T401E mutants. In vitro phosphorylation comparisons of the wild-type (wt) <sub>XJ</sub>TGB1 protein, and the Thr-395 and Thr-401 mutants were performed with HvCK2 $\alpha$ and NbCK2 $\alpha$ , respectively. Compared with the wt xITGB1 protein (Fig. 4B, lane 1), the phosphorylation level of the XJTGB1<sub>T395A</sub> mutant protein was reduced partially (Fig. 4B, lane 2). However, both the XJTGB1T395D and the XJTGB1T395E mutant proteins incorporated slightly larger amounts of <sup>32</sup>P label than the wt <sub>XJ</sub>TGB1 protein (Fig. 4B, compare lane 1 with lanes 3 and 4), suggesting that the positive charges imparted by the aspartic acid and glutamic acid residues increased the kinase efficiency. As anticipated, the phosphorylation intensities of the <sub>XJ</sub>TGB1<sub>T401A</sub> (Fig. 4B, lane 5) and <sub>XJ</sub>TGB1<sub>T401E</sub> (Fig. 4B, lane 7) proteins only showed faint shadows. We also observed similar reductions during incorporation into the <sub>XJ</sub>TGB1<sub>T401D</sub> protein (Fig. 4B, lane 6). We believe that Thr-395 may have been phosphorylated, and this is supported by negligible incorporation into the three double mutants (Fig 4B, lane 8; Fig. 4C, lanes 1–3). Thus, our interpretation of these results is that Thr-401 is a major phosphorylation site and that Thr-395 is a minor phosphorylation target.

Based on the data above and previous analyses of 308 CK2 phosphorylation sites for other proteins (Meggio and Pinna, 2003), we propose that Thr-395 is a docking site for CK2 because phosphorylation of this residue enhanced kinase activity at Thr-401. To evaluate this hypothesis, we used a double mutant, XITGB1T395A/T401A, for phosphorylation. The results revealed extremely low, if any, <sup>32</sup>P incorporation into other TGB1 residues (Fig. 4B, lane 8; Fig 4C, lane 3). To determine whether there was 'off-site' targeting of other residues within x1TGB1 at the proposed Thr-395 docking site, we engineered the XJTGB1T395D/T401A and XJTGB1T395E/T401A double mutants, which could not be phosphorylated at Thr-401, and both double mutants had negligible levels of <sup>32</sup>P incorporation (Fig. 4C, lanes 1 and 2). In summary, these results support the hypothesis that Thr-395 functions as a docking residue and that Thr-401 is the major phosphorylation site within the kinase motif of the xITGB1 protein.

#### Mutations of the <sub>XJ</sub>TGB1 protein phosphorylation site affect BSMV local and systemic infections of dicots and monocots

To determine whether the  $_{XJ}TGB1$  mutants affected systemic movement in *N. benthamiana*, leaves were infiltrated with *Agrobacterium* harbouring the pCa- $\alpha_{XJ}$ , wt pCa- $\beta_{XJ}$ , or individual TGB1 mutant derivatives, and the pCa- $\gamma_{XJ}$  clones. Three independent experiments revealed that only wt  $_{XJ}TGB1$  and the  $\beta_{XJ}$ -TGB1 $_{T395A}$  mutant, which exhibited slightly lower phosphorylation levels than the wt  $_{XJ}TGB1$  protein, were able to establish systemic infections at 10 dpi (Fig. 5A, lanes 2 and 3). None of the remaining infiltrations containing single or double  $_{XJ}TGB1$  mutants developed mosaic symptoms or invaded the upper uninoculated leaves as assessed by the absence of CP accumulation (Fig. 5A, lanes 4–11).

For systemic infectivity on monocots, barley and wheat leaves were co-inoculated with *in vitro* transcripts of pT7- $\alpha_{XJ}$ , pT7- $\beta_{XJ}$  (wt or mutant derivatives), and pT7- $\gamma_{XJ}$ . Visual observations, ELISA and RT-PCR analyses demonstrated that the barley and wheat plants had similar systemic infections on upper uninoculated leaves at 14 dpi after wt and mutant<sub>XJ</sub>TGB1 inoculations (Fig. 5B, C). In the case of  $\beta_{XJ}$ -TGB1<sub>T395A</sub>, a milder infection phenotype was noted on the systemically infected cereal leaves, compared with those of the wt  $\beta_{XJ}$ -TGB1<sub>T395D</sub> was unable to infect the plants



**Fig. 4.** Thr-401 is the major  $_{xJ}$ TGB1 protein site for CK2 phosphorylation. (A) LC-MS/MS analysis of  $_{xJ}$ TGB1 protein phosphorylation by NbCK2 $\alpha$ . The absence of phosphoric acid (97.9769 Da) on the  $y_{16}$  ion fragment demonstrates that Thr-401 is a phosphorylation site for CK2 kinase. (B) Identification of the phosphorylation sites in  $_{xJ}$ TGB1 protein mutants by *in vitro* phosphorylation with HvCK2 $\alpha$  and NbCK2 $\alpha$ . The radioactive intensities of the  $_{xJ}$ TGB1 protein mutants indicate the extent of radiolabelling with [ $\gamma_{-32}$ P]ATP. CBB-stained proteins at the bottom of the panels (B) and (C) are as indicated in Fig. 2B. (C) Phosphorylation comparisons of selected  $_{xJ}$ TGB1 protein mutants with wt  $_{xJ}$ TGB1 protein to confirm that Thr-401 is the major phosphorylated residue. (This figure is available in colour at *JXB* online.)

systemically (Fig. 5B, C, lane 4). In contrast, when inoculated with the  $\beta_{XJ}$ -TGB1<sub>T395E</sub> mutant, which more effectively mimics phosphorylation of threonine residues, the cereal plants exhibited CP accumulation levels that were similar to wt BSMV (Fig. 5B, C, lane 5). Moreover, the  $\beta_{XJ}$ -TGB1<sub>T401E</sub> mutant elicited systemic infections, albeit with slightly lower CP levels (Fig. 5B, C, lane 8), but the levels were higher than those with  $\beta_{XJ}$ -TGB1<sub>T401A</sub> (Supplementary Table S6, available at *JXB* online). However, the  $\beta_{XJ}$ -TGB1<sub>T395A/T401A</sub>,

 $\beta_{XJ}$ -TGB1<sub>T395D/T401A</sub>, and  $\beta_{XJ}$ -TGB1<sub>T395E/T401A</sub> double mutants were unable to infect either barley or wheat (Fig. 5B, lanes 9–11).

Taken together, these results demonstrated that the  $_{XJ}TGB1$  phosphorylation site mutations generally reduced the infection efficiencies in dicots and monocots, and that the mutants had more dramatically compromised systemic movement phenotypes in *N. benthamiana*. To summarize, the T395A, T395E, T401A, and T401E mutants exhibited systemic movement in



**Fig. 5.** Mutants affecting phosphorylation of the  $_{X,J}$ TGB1 protein have host-specific effects on systemic infectivity. (A) Symptoms of *N. benthamiana* elicited after infiltration with an *Agrobacterium* mixture harbouring pCa- $\alpha_{X,J}$ , pCa- $\gamma_{X,J}$ , and pCa- $\beta_{X,J}$  or its phosphorylation site mutants. Upper uninfiltrated leaf tissues were harvested and photographed at 10 dpi (top). CP ELISA (middle) and RNA $\gamma$  RT-PCR amplification (bottom) were monitored to estimate the infectivity levels. (B, C) Systemic symptoms appearing in barley (B) and wheat (C) after inoculation with pT7- $\alpha_{X,J}$  and pT7- $\gamma_{X,J}$  *in vitro* transcripts mixed with pT7- $\beta_{X,J}$  and various phosphorylation mutant transcripts. Leaves were photographed at 14 dpi (top) and all experiments were repeated three times. (This figure is available in colour at *JXB* online.)

barley and wheat, whereas only the T395A mutant is able to establish systemic infections in *N. benthamiana*.

We next conducted experiments to evaluate the cell-to-cell movement profiles of the Thr-395 and Thr-401 mutants in *N. benthamiana* and barley. For this purpose, *N. benthamiana* 

leaves were infiltrated with *Agrobacterium* containing pCa- $\alpha_{XJ}$ , pCa- $\beta_{XJ}$  and its mutant derivatives, and a pCa- $\gamma_{XJ:GFP}$  construct that harbours a  $\gamma$ b:GFP reporter gene to assess cell-to-cell movement. In barley, leaves were co-inoculated with T7 transcripts of the RNA $\alpha$ , RNA $\beta$  derivatives, and

RNA $\gamma$ - $\gamma_{XJ:GFP}$  transcripts. Epidermal cells of the leaves were observed by confocal microscopy at 3 dpi and compared with inoculated controls lacking the RNA $\beta$ .

The localized movement in infiltrated *N. benthamiana* leaves generally reflected the systemic infection phenotypes elicited by the mutants (Fig. 6A). In *N. benthamiana*, the wt  $\beta_{XJ}$  and  $\beta_{XJ}$ -TGB1<sub>T395A</sub> mutant both exhibited cell-to-cell movement encompassing several cells at 3 dpi (Fig. 6A), as expected due to their ability to elicit systemic infections. The remaining mutants usually developed fluorescence in a single cell or rarely in two to three adjacent cells (Fig. 6A). Hence, the localized movements of the mutants correlated reasonably well with their systemic movement patterns in *N. benthamiana*. In barley leaves, most of the fluorescence at 3 dpi appeared in mesophyll cells, but in this case, the virus had to traverse only two to three cell layers before encountering the closely aligned parallel vasculature (Fig. 6B). Hence, the  $\beta_{XJ}$ -TGB1<sub>T395A</sub>,



Bar = 500 µm

**Fig. 6.** Effects of  $_{X_J}$ TGB1 protein phosphorylation on  $_{X_J}$ BSMV cellto-cell movement in *N. benthamiana* and barley. (A) Fluorescence in *N. benthamiana* leaves at 3 dpi with an *Agrobacterium* mixture of pCa- $\alpha_{ND}$ , pCa- $\gamma_{ND:GFP}$ , and pCa- $\beta_{XJ}$  or the pCa- $\beta_{XJ}$  mutant derivatives. The total bacterial concentrations for infiltration were OD<sub>600</sub> of 0.08. (B) Fluorescence in barley leaves at 3 dpi with *in vitro* transcripts of RNA $\alpha$  and RNA $\gamma$ :GFP plus wt  $_{XJ}$ RNA $\beta$  or the  $_{XJ}$ RNA $\beta$  phosphorylation site mutant derivatives. Bars represent 500  $\mu$ m. (This figure is available in colour at *JXB* online.)

 $\beta_{XJ}$ -TGB1<sub>T395E</sub>, and  $\beta_{XJ}$ -TGB1<sub>T401A</sub> mutants that established systemic infections in barley and wheat needed to negotiate only a limited number of mesophyll cells to reach the vascular elements for systemic spread, whereas movement through a larger number of cells was required to reach the dicot vasculature. The other mutants,  $\beta_{XJ}$ -TGB1<sub>T395D</sub>,  $\beta_{XJ}$ -TGB1<sub>T401D</sub>, and  $\beta_{XJ}$ -TGB1<sub>T395A/T401A</sub>, exhibited more limited cell-to-cell movement compared with  $\beta_{XJ}$ -TGB1<sub>T395A</sub>,  $\beta_{XJ}$ -TGB1<sub>T395E</sub>, and  $\beta_{XJ}$ -TGB1<sub>T401A</sub>, but could spread to a few adjacent cells. However, these three mutants were unable to invade the upper cereal leaves. Hence, these results demonstrate that phosphorylation activities at Thr-395 and Thr-401 differentially affecte systemic movement in monocot versus dicot hosts and suggest that at least some of the host-specific results may be a consequence of the vasculature architecture of these hosts.

# Phosphorylation promotes virus infection of <sub>xJ</sub>BSMV by enhancing TGB1 and TGB3 protein interactions

Previous studies have shown that BSMV spreads from cell to cell through the co-ordinated actions of TGB proteins, which co-localize at the cell wall in close association with PD, during cell-to-cell movement in monocots and dicots (Lim *et al.*, 2008). Our results shown above demonstrated that interference with CK2 phosphorylation at the  $_{XJ}$ TGB1 Thr-395 and Thr-401 sites affected  $_{XJ}$ BSMV local and systemic movement. TGB1 is a multifunctional protein that engages in homologous interactions and formation of a ribonucleoprotein complex containing viral genomic and subgenomic RNAs (Lawrence and Jackson, 2001). Therefore, we used three approaches to identify  $_{XJ}$ TGB1 protein functions affected by CK2 phosphorylation.

To determine whether the RNA-binding affinity of the xITGB1 protein changed upon phosphorylation, we first used purified wt XJTGB1 and the double mutant XJTGB1T395A/ T401A proteins in EMSA with DIG-labelled RNA transcripts (Fig. 7A). The results clearly showed that both proteins bound almost all of the available RNA at 250 ng, indicating that the mutant protein did not affect RNA-binding activities (Fig. 7A, panels 1 and 2). Next, to determine phosphorylation effects directly, the XJTGB1 protein was incubated in phosphorylation assay buffer containing unlabelled ATP, with and without the addition of purified NbCK2 $\alpha$ , and EMSA assays were performed to compare the abilities of the non-phosphorylated and phosphorylated x1TGB1 proteins to bind the labelled RNA transcripts (Fig. 7A, panels 3 and 4). In addition, the RNA-binding activities of the phosphorylated wt xJTGB1 and mutant xJTGB1T395A/T401A proteins were compared, but the XJTGB1T395A/T401A protein was found to retain almost the same level of RNA binding as the wt <sub>XJ</sub>TGB1 protein (Fig. 7A, panels 5 and 6). Hence, phosphorylation appeared to have little, if any, effect on the RNAbinding activities of the xJTGB1 protein.

To evaluate the possible role of phosphorylation in heterologous interactions of  $_{XJ}TGB1$  and  $_{XJ}TGB3$  proteins, experiments were carried out with His-tagged  $_{XJ}TGB1$  and its mutant  $_{XJ}TGB1_{T395A/T401A}$  fusion proteins in co-expressions with the GST: $_{XJ}TGB3$  protein in *E. coli* BL21 cells. Both



**Fig. 7.** Effect of  $_{XJ}$ TGB1 protein phosphorylation mutants on its functions. (A) Comparison of RNA binding by the phosphorylated native  $_{XJ}$ TGB1 and double-mutant  $_{XJ}$ TGB1 $_{T395A/T401A}$  proteins. (B) GST affinity chromatography comparisons of the  $_{XJ}$ TGB1 and double-mutant  $_{XJ}$ TGB1 $_{T395A/T401A}$  proteins with the GST: $_{XJ}$ TGB3 protein. The concentrations of the TGB proteins were similar in the experiments, but the  $_{XJ}$ TGB1 $_{T395A/T401A}$  protein had approximate 40% TGB3 protein-binding efficiency compared with the wt  $_{XJ}$ TGB1 protein. The illustrated binding result is typical of three independent experiments. (C) Co-localization of TGB proteins. Confocal laser-scanning microscopy observation of *N. benthamiana* leaf epidermal cells co-infiltrated with mixtures of *Agrobacterium* harbouring GFP:\_{XJ}TGB1 or the GFP:\_{XJ}TGB1\_{T395A/T401A} mutant derivatives and the pGD-TGB2 and RFP:TGB3 plasmids. Bars, 50  $\mu$ m. (This figure is available in colour at *JXB* online.)

 $_{XJ}TGB1$  and  $_{XJ}TGB1_{T395A/T401A}$  proteins were retained to some extent on the affinity columns by the GST: $_{XJ}TGB3$  protein (Fig. 7B), but our three experiments consistently showed that the  $_{XJ}TGB1_{T395A/T401A}$  protein bound the TGB3 protein approximate 40% less effectively than the  $_{XJ}TGB1$  protein. These results thus suggested that the mutant  $_{XJ}TGB1_{T395A/}$  $_{T401A}$  may weaken TGB1:TGB3 protein interactions and result in impaired cell-to-cell movement functions of  $_{XJ}BSMV$ .

In additional attempts to ascertain whether the compromised TGB1:TGB3 protein interactions or CK2 phosphorylation affected localization of TGB proteins in plant cells, co-localization assays were performed by transient expression of the three TGB proteins via agroinfiltration in *N. benthamiana*, and GFP and RFP localizations were evaluated at 2 dpi by confocal laser-scanning microscopy. The results revealed that TGB2 and TGB3, the  $_{XJ}$ TGB1 $_{T395A/T401A}$  protein, and the wt  $_{XJ}$ TGB1 protein had similar TGB localization patterns (Fig. 7C). Taken together, we conclude that phosphorylation promotes  $_{XJ}$ BSMV infection by enhancing the interactions between the  $_{XJ}$ TGB1 and  $_{XJ}$ TGB3 proteins, although this effect was not sufficient to substantially alter the TGB localization patterns visible by confocal microscopy.

### Discussion

Reversible phosphorylation and dephosphorylation of proteins have regulatory roles in a wide range of cellular processes, including cell signalling transduction (Moreno-Romero et al., 2011), protein subcellular localization (Nardozzi et al., 2010), and protein–protein (Trott et al., 2001) and protein– nucleic acid interactions (Schuck et al., 2013). Numerous proteins with distinct phosphorylation sites have been investigated, and protein kinases affecting a wide range of cellular responses have been characterized (Peck, 2006; Ubersax and Ferrell, 2007; Bond et al., 2011; Dissmeyer and Schnittger, 2011). A growing body of evidence now shows that viral proteins with different functions are phosphorylated by various protein kinases during infection. These include CK2, PKA, PKC, and CK1 protein kinases (Lee and Lucas, 2001; Link et al., 2011; Makarov et al., 2012; Hung et al., 2014), and among these kinases, CK2 phosphorylation effects on infectivity have been most extensively studied.

Protein kinase CK2, a conserved Ser/Thr kinase existing in almost all eukaryotes, phosphorylates proteins with a consensus phosphorylation site motif (S/T-D/E-X-E/D, where X is any residue) (Ubersax and Ferrell, 2007). Increasing evidence indicates that CK2 protein phosphorylation has important roles in plant growth and development (Mulekar et al., 2012), and that CK2 also regulates virus infection processes, including virion assembly, cell-to-cell and long-distance movement, and interactions between viral proteins and other host proteins (Jakubiec and Jupin, 2007; Nardozzi et al., 2010). In addition to the MP phosphorylation effects mentioned in the Introduction, phosphorylation of cucumber mosaic virus, cucumber necrosis virus, and turnip yellow mosaic virus RdRp has substantial effects on virus replication (Kim et al., 2002; Shapka et al., 2005; Stork et al., 2005; Jakubiec et al., 2006; Jakubiec and Jupin, 2007). Ser/Thr phosphorylation has also been suggested to affect the CP functions of several plant viruses. For example, phosphorylation of potato virus A (PVA) CP by host CK2 inhibits viral RNA binding in vitro, and mutation of a major phosphorylation CP site generates a PVA variant that is defective in cell-tocell and long-distance movement (Ivanov et al., 2001, 2003). In addition, phosphorylation of the cauliflower mosaic virus CP precursor at several sites by CK2 is important for virus infectivity and symptom development (Chapdelaine *et al.*, 2002). Phosphorylation of the bamboo mosaic virus CP by CK2 also regulates cell-to-cell movement by modulating RNA binding (Hung *et al.*, 2014).

Although phosphorylation of the PSLV N-terminal portion of the TGB1 protein has been reported (Makarov et al., 2012), but the results were not extended to evaluate the roles of phosphorylation in PSLV movement processes. Our results now demonstrate that the BSMV TGB1 protein is phosphorylated by CK2 in vitro and in planta, and that the phosphorylation events affect virus movement. Although the prediction programs we used suggest that the x<sub>1</sub>TGB1Thr-395 residue in the <sup>395</sup>TDYD<sup>398</sup> site is more likely to be a conserved CK2 phosphorylation site than the Thr-401 (<sup>401</sup><u>TDET</u><sup>404</sup>) site (Meggio and Pinna, 2003), it is noteworthy that Thr-401 localizes within an acidic residue-rich region (<sup>399</sup>GETDETEK<sup>406</sup>) that may be more favourable for phosphorylation (Battistutta et al., 2000; Riera et al., 2001) than the Thr-395 residue. Thus, based on the in vitro phosphorylation assays of the TGB1 mutant derivatives, as well as results derived from LC-MS/MS analysis, we conclude that Thr-401 is the major TGB1 phosphorylation site and that Thr-395 functions as a CK2 docking site and has a more limited phosphorylation role. To the best of our knowledge, this is the first report showing that a plant viral protein, which can be phosphorylated by CK2, has a CK2 docking site adjacent to the phosphorylation site.

To determine the effects of XITGB1 Thr-395 and Thr-401 phosphorylation on virulence, point mutations were introduced into the XJBSMV clone. Infectivity results in the monocot and dicot hosts revealed that the mutant derivatives differed in their systemic movement phenotypes. For example, the T395A mutant was the only derivative able to infect N. benthamiana, barley, and wheat systemically, but the T395E, T401A, and T401E mutants also systemically infected the monocot hosts (Fig. 5). Our results suggest that the mutants may be compromised by partial disruption of phosphorylation and dephosphorylation dynamics in TGB1 in ways that contribute to diminished cell-to-cell movement. However, the amino acid structures of the T395A, T395D, T395E, T401D, and T401E substitutions are not entirely consistent with this simplistic model, as it is obvious that the substituted amino acids differ in the sizes of their side chains and their charges. Moreover, we cannot exclude the possibility that 'off-site' phosphorylation of Thr-395 or phosphorylation by kinases other than CK2 may be elicited by the substitutions and that these events may contribute to protein modifications that affect local and vascular movement.

Hordeivirus TGB1 proteins are multifunctional and contain two positively charged regions rich in lysine (K) and arginine (R) residues at the N-terminal half of the protein and a C-terminal region consisting of seven conserved motifs (I, IA, II, III, IV, V, and VI) (Jackson *et al.*, 2009). The Thr-395 and Thr-401 sites are located between domain IV and V of the TGB1 protein and are not included in the most highly conserved regions. The hordeivirus TGB1 proteins have multiple ssRNA- and dsRNA-binding sites, and hence mutagenesis of single or closely associated <sub>XJ</sub>TGB1 protein sites may not have obvious effects on RNA-binding activities *in vitro*.

We have shown previously that the BSMV TGB1 protein is the major protein component of ribonucleoprotein complexes involved in BSMV cell-to-cell movement. TGB1 also participates in interactions of TGB1 and TGB3 proteins during intra- and intercellular virus movement, and functions in TGB1:TGB3 interactions that recruit the TGB2 protein during transport through PD to adjacent cells (Jackson et al., 2009). These interactions are critical for movement because TGB3 serves as a bridge to direct TGB co-localization at the cell wall and to establish close associations with the PD (Lim et al., 2009). Even though both CK2 phosphorylation sites (T395A/T401A) in <sub>XJ</sub>TGB1 were mutated simultaneously, the mutant TGB1 protein did not elicit obvious changes in subcellular localization patterns (Fig. 7C). However, the XJTGB1T395A/T401A protein did reduce binding affinity to the TGB3 protein compared with the wt TGB1 protein. Therefore, our results provide evidence that CK2 phosphorylation of TGB1 affects BSMV movement by modulating TGB1:TGB3 protein interactions.

In summary, our results shown here provide evidence showing that phosphorylation of the BSMV x1TGB1 protein by CK2 at C-terminal residues affects cell-to-cell movement and the systemic infection phenotype. Moreover, the mutant results are compatible with a model whereby modulation of TGB1:TGB3 interactions contribute to phenotypic differences with in BSMV movement in monocot and dicot hosts. Compared with the TGB1 proteins of other hordeiviruses (Supplementary Fig. S2A, available at JXB online), Thr-401 but not Thr-395 is conserved in PSLV TGB1 and BSMV TGB1. This implies that the proposed docking site function of Thr-395 may be unique for BSMV phosphorylation. Furthermore, multiple sequence alignments of TGB1 proteins (Supplementary Fig. S2B) showed that the Thr-395 and Thr-401 sites are highly conserved among six sequenced BSMV strains, suggesting that phosphorylation of the TGB1 protein is required during infection of all BSMV strains.

#### Supplementary data

Supplementary data are available at JXB online.

Supplementary Fig. S1. Phosphorylation predictions of the  $_{XJ}TGB1$  protein by the GPS 2.1 program (A) and the Scansite Motif Scanner online server (B).

Supplementary Fig. S2. Alignment of the TGB1 proteins of the hordeiviruses (A) and among six sequenced BSMV strains (B).

Supplementary Table S1. Primers used in construction and analysis of biologically active BSMV Xinjiang cDNA clones.

Supplementary Table S2. Primers used for site-specific mutagenesis of Xinjiang RNA $\beta$  clones.

Supplementary Table S3. Sequence alignment of Xinjiang strain RNA $\alpha$  with different BSMV strains.

Supplementary Table S4. Sequence alignment of Xinjiang strain RNAβ with different BSMV strains.

Supplementary Table S5. Sequence alignment of Xinjiang strain RNAγ with different BSMV strains.

Supplementary Table S6. Systemic infectivity efficiency of <sub>XJ</sub>BSMV TGB1 phosphorylation mutants on dicot and monocot hosts.

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