A Plant Virus Movement Protein Regulates the Gcn2p Kinase in Budding Yeast

Frederic Aparicio, Rafael Aparicio-Sanchis, José Gadea, Jesús Ángel Sánchez-Navarro, Vicente Pallás, José Ramón Murguía*

Department of Stress Biology, Instituto de Biología Molecular y Celular de Plantas (UPV-CSIC), Valencia, Spain

Abstract

Virus life cycle heavily depends on their ability to command the host machinery in order to translate their genomes. Animal viruses have been shown to interfere with host translation machinery by expressing viral proteins that either maintain or inhibit eIF2α function by phosphorylation. However, this interference mechanism has not been described for any plant virus yet. *Prunus necrotic ring spot virus* (PNRSV) is a serious pathogen of cultivated stone fruit trees. The movement protein (MP) of PNRSV is necessary for the cell-to-cell movement of the virus. By using a yeast-based approach we have found that over-expression of the PNRSV MP caused a severe growth defect in yeast cells. cDNA microarrays analysis carried out to characterise the molecular level the growth interference phenotype reported the induction of genes related to amino acid deprivation suggesting that expression of MP activates the GCN pathway in yeast cells. Accordingly, PNRSV MP triggered activation of the Gcn2p kinase, as judged by increased eIF2α phosphorylation. Activation of Gcn2p by MP expression required a functional Tor1p kinase, since rapamycin treatment alleviated the yeast cell growth defect and blocked eIF2α phosphorylation triggered by MP expression. Overall, these findings uncover a previously uncharacterised function for PNRSV MP viral protein, and point out at Tor1p and Gcn2p kinases as candidate susceptibility factors for plant viral infections.

Introduction

*Saccharomyces cerevisiae* cells transiently inhibit initiation of protein synthesis under environmental stress to avoid misfolding of proteins which could compromise cell viability. Inhibition of translation is achieved by phosphorylation of the alpha subunit of the eukaryotic translation initiation factor-2 (eIF2α). The sole eIF2α kinase present in budding yeast is encoded by the *GCN2* gene. The general control non-repressible 2 protein (Gcn2p) regulates the selective translation of the Gcn4p transcription factor upon nutrient deprivation conditions. This regulation is exerted by four uORFs located in the 5’ untranslated region of the *GCN4* mRNA, which makes it hypersensitive to eIF2 levels. When there is no amino acid limitation, the levels of active eIF2 are high and the uORFs block translation of *GCN4* mRNA. In cells under amino acid starvation there is an accumulation of uncharged tRNAs that stimulates Gcn2p kinase activity. Phosphorylation of eIF2α diminishes the levels of active eIF2 and alleviates inhibition by uORFs, thus favouring *GCN4* mRNA translation as well as blocking general translation [1]. High levels of Gcn4p then activate the expression of genes encoding amino acid biosynthesis pathways. Increased eIF2α phosphorylation also occurs by starvation for purines, glucose, growth on ethanol, high salinity in the growth medium and treatment with methyl methanesulphonate [2,3]. Thus, Gcn2p seems to be a master regulator of gene expression in yeast and in response to various kinds of stresses. The fundamental nature of the GCN pathway is reflected by the fact that Gcn2p function is conserved throughout evolution. In mammals four protein kinases are known to phosphorylate eIF2α, namely GCN2, PKR, HRI and PERK. In plants, *Arabidopsis* and rice genome analysis revealed that plants apparently encode a single eIF2α kinase orthologue of yeast GCN2 [4,5].

Virus life cycle heavily depends on their ability to command the host machinery in order to translate their genomes. This implies that viral RNAs must compete with cellular mRNAs for the host translation apparatus. To achieve this, viruses have devised strategies to inhibit cellular protein synthesis while ensuring efficient translation of their cognate RNAs. Indeed, different viral proteins have been shown to mimic host nutrient/growth signals which activate signaling pathways that ensure viral replication. For example, adenoviruses have evolved proteins that activate the mTOR pathway, irrespective of the cellular microenvironment [6]. mTOR integrates nutrient and growth factor signals to regulate the translation initiation of mRNAs important for cell growth [7]. Another strategy consists of regulating eIF2 function by phosphorylation/dephosphorylation. Thus, the g34.5 protein from HSV-1 complexes with the cellular protein phosphatase 1a (PP1a) and directs dephosphorylation of eIF2α [8]. Vaccinia virus K3L polypeptide reduces the level of eIF2α phosphorylation by GCN2 [9]. Rotaviruses cause eIF2α phosphorylation depending on the synthesis of three viral proteins, VP2, NSP2, and NSP5...


**Results**

**Overexpression of PNRSV MP negatively affects growth in yeast**

The PNRSV MP (MP\textsubscript{pnrsv}) protein is 282 amino acids long, and contains the characteristic hydrophobic region preceded by an RNA-binding domain. Both regions are present in all members of the ilarovirus genus and are necessary for the cell-to-cell viral transport \cite{28,33,34}. We used a yeast-based approach to decipher how MP\textsubscript{pnrsv} affects cellular or modulates host factors. HA-tagged versions of MP\textsubscript{pnrsv} and MP\textsubscript{pnrsv\textsubscript{AHR}}, a non-functional MP\textsubscript{pnrsv} mutant lacking its hydrophobic region \cite{34}, were over-expressed in wild type (WT) yeast cells under the control of a yeast plasmid with a doxycycline-repressible promoter, and yeast cell growth was monitored in the absence/presence of doxycycline. As expected, doxycycline treatment did not affect yeast cell growth (Fig. 1A). Interestingly, MP\textsubscript{pnrsv}-expressing yeast cells exhibited a severe growth defect in a doxycycline-dependent manner (Fig. 1B). Expression of MP\textsubscript{pnrsv\textsubscript{AHR}} also induced a slight growth defect, but not to such extent as MP\textsubscript{pnrsv} (Figs. 1C, D). Similar growth defects have been reported in yeasts expressing foreign proteins \cite{37}. It is possible that a similar scenario occurs when expressing MP\textsubscript{pnrsv\textsubscript{AHR}} in yeast, although we cannot rule out that other mechanisms may be involved. Differences in growth were not due to expression levels as both MP\textsubscript{pnrsv} and MP\textsubscript{pnrsv\textsubscript{AHR}} proteins accumulated similarly (Fig. 1F). Therefore, the strong growth interference phenotype required a functional MP\textsubscript{pnrsv}. To further analyze the specific MP\textsubscript{pnrsv}-induced growth interference phenotype, we explored the effect on yeast cell growth of other MPs from differentially related viruses as Brome mosaic virus (BMV) and Cucumber mosaic virus (CMV) and two non-related viruses as Tobacco mosaic virus (TMV) and Grapevine fanleaf virus (GFLV) respectively. HA-tagged versions of these MPs were also expressed in yeast under the control of the yeast plasmid doxycycline-repressible promoter. Measurement of the yeast cell growth at the exponential phase with or without doxycycline showed that BMV, CMV and TMV MPs expression elicited a growth interference phenotype that was totally absent in the case of GFLV MP (Fig. 1E). Western blot analysis confirmed that all MPs accumulated at similar levels in yeast cells (Fig. 1G). Interestingly, all the MPs analyzed, except the GFLV MP, are associated to the membrane of the endoplasmic reticulum (ER) suggesting that the growth defect observed in yeast was somehow linked to MPs that move through an ER associated pathway.

**MP\textsubscript{pnrsv} overexpression triggers growth defect in yeast cells by activating the GCN pathway**

To further characterize at the molecular level the growth interference phenotype induced by MP\textsubscript{pnrsv} expression, we compared the gene expression profile of the MP\textsubscript{pnrsv}-expressing strain with that of the wild type (WT) strain using cDNA microarrays. We used Significance Analysis of Microarrays to identify differentially expressed genes between the WT and MP\textsubscript{pnrsv} strains. Any given gene was considered to be differentially regulated when its expression was at least 2 fold higher or lower in the MP\textsubscript{pnrsv} strain than in the WT strain (FDR <6%). Following these criteria, 324 genes were induced by MP\textsubscript{pnrsv} over-expression. Surprisingly, no gene was found to be differentially repressed. We performed a functional classification of induced genes according to Gene Ontology (GO) terms using the GO Term Finder tool. Interestingly, functional categories such as glutamine and nitrogen metabolism were significantly enriched in the set of induced genes (see Table S1). Nutrient deprivation
activates glutamine and nitrogen metabolism gene expression in budding yeast. Nutrient deprivation conditions trigger the GCN pathway, a nutrient-sensing pathway that transiently inhibits translation initiation and simultaneously favors selective translation of the GCN4 mRNA. GCN4 encodes a transcriptional activator of amino acid biosynthesis genes. The major transducer of the GCN pathway is the Gcn2p kinase which phosphorylates eIF2α. To explore whether over expression of MPpnrsv was able to activate the GCN pathway we measured the phosphorylation status of eIF2α in WT yeast cells by immunoblot analysis using a commercially available polyclonal antibody that specifically recognizes eIF2α phosphorylated at serine 51 (see methods). As shown in Figure 2A, overexpression of MPpnrsv, but not the mutant MPpnrsvΔHR, incremented eIF2α phosphorylation in WT yeast cells, thus showing that MPpnrsv expression activated the Gcn2p kinase. Furthermore, the growth interference phenotype associated with MPpnrsv overexpression was alleviated in the gcn2Δ mutant (see Fig. S1), these data indicated that MPpnrsv expression triggered a growth defect in yeast cells likely by activating the GCN pathway.

To gain further insights into the mechanism by which MPpnrsv activated the GCN pathway, we performed a microarray experiment to identify differentially expressed genes between the MPpnrsv and the mutant MPpnrsvΔHR overexpressing strains. Data analysis using the GO Term Finder tool revealed that functional categories such as ribosome biogenesis and initiation of protein synthesis were enriched in the set of differentially expressed genes (see Table S2 and S3). These functional categories are mainly regulated by the TOR1 nutrient sensing kinase. Therefore, we explored the involvement of TOR1 in MPpnrsv-dependent yeast growth defect using rapamycin, a highly selective inhibitor of the Torp1 kinase.

Yeast cells expressing MPpnrsv were grown in minimal medium with or without doxycycline and transformed with either pCM (lane 1) or with plasmids expressing MPpnrsv and MPpnrsvΔHR plasmids. (C) Rapamycin treatment alleviates the growth defect caused by MPpnrsv expression in yeast. Yeast cells expressing MPpnrsv were grown in SD medium with or without rapamycin, the same medium with 200 μg/ml cycloheximide (C200), and with 100 μg/ml (R100) or 200 μg/ml (R200) Rapamycin respectively. Data in B and C represent the average +/- s.e. of at least three independent experiments, each one done in triplicate. (D) Rapamycin treatment blocks phosphorylation of yeast eIF-2α triggered by MPpnrsv expression. Immunodetection of phospho-eIF-2α levels in protein extracts from yeast cells transformed with pCM and MPpnrsv expressing plasmids, grown in SD medium with or without (+R) or without (-R) 200 μg/ml Rapamycin for three hours. Coomassie Blue (CB) stained gel used as loading control.

doi:10.1371/journal.pone.0027409.g001

doi:10.1371/journal.pone.0027409.g002

Figure 1. Expression of viral MPs of the 30 K superfamily interferes with yeast cell growth. Growth kinetics of WT yeast cells cultured in SD medium +/- Doxycycline (DOX) and transformed with empty vector (pCM) (A), a full-length MPpnrsv expressing plasmid (B) or a truncated MPpnrsvΔHR expressing plasmid (C). (D) Quantitation of cell growth in SD medium +/- DOX of yeast cultures transformed with vector alone or with expressing MPpnrsv and MPpnrsvΔHR plasmids. (E) Quantitation of cell growth in SD medium +/- DOX, of WT yeasts transformed with either pCM (lane 1) or with plasmids expressing PNRSV, BMV, CMV, TMV and GLFV MPs (lanes 2, 3, 4, 5 and 6, respectively). Data in D and E, represent the average +/- standard error (s.e.) of at least three independent experiments, each one done in triplicate (p<0.001). (F) Immunodetection of MPpnrsv (lane 3) and MPpnrsvΔHR (lane 1) in protein extracts from yeast cells transformed with the corresponding expression vectors or pCM as negative control (lane 2). (G) The same as in (F), but in protein extracts from yeast cells transformed with pCM (lane 1) or plasmids expressing PNRSV, BMV, CMV, TMV or GLFV MPs (lanes 2, 3, 4, 5 and 6, respectively) Coomassie Blue (CB) stained gel used as loading control.

doi:10.1371/journal.pone.0027409.g001

Figure 2. MPpnrsv expression activates the GCN pathway in a TOR pathway-dependent manner. (A) Immunodetection of phospho-eIF-2α levels in protein extracts from yeast cells transformed with pCM (lane 1), MPpnrsvΔHR (lane 2) and MPpnrsv (lane 3) expressing plasmids. Yeast cells were grown for 3 hours in SD medium without DOX. Even loading of gels was confirmed by CB staining of membranes after transfer. (B) Disruption of the GCN pathway alleviates the growth defect caused by MPpnrsv expression in yeast. Quantitation of cell growth in SD medium +/- DOX, of WT, gcn2Δ and gcn4Δ yeast strains expressing the MPpnrsv. (C) Rapamycin treatment alleviates the growth defect caused by MPpnrsv expression in yeast. Yeast cells expressing MPpnrsv were grown in SD medium +/- DOX (SD), the same medium with 200 μg/ml cycloheximide (C200), and with 100 μg/ml (R100) or 200 μg/ml (R200) Rapamycin respectively. Data in B and C represent the average +/- s.e. of at least three independent experiments, each one done in triplicate. (p<0.005) (D) Rapamycin treatment blocks phosphorylation of yeast eIF-2α triggered by MPpnrsv expression. Immunodetection of phospho-eIF-2α levels in protein extracts from yeast cells transformed with pCM and MPpnrsv expressing plasmids, grown in SD medium with (+R) or without (-R) 200 μg/ml Rapamycin for three hours. Coomassie Blue (CB) stained gel used as loading control.

doi:10.1371/journal.pone.0027409.g002
effect was specific as cycloheximide treatment blocked yeast growth regardless the presence of MPpnrsv. Most remarkably, rapamycin treatment blocked eIF2α phosphorylation triggered by MPpnrsv expression (Fig. 2D). Hence, activation of Gcn2p by MPpnrsv expression required a functional Tor1p kinase, thus uncovering a functional link between both nutrient-sensing pathways.

Discussion

By using a yeast-based approach we have identified that the PNRSV MP is able to specifically activate the nutrient regulated kinase Gcn2p. Gcn2p activation seemed to be dependent on the nutrient regulated kinase Tor1p, as rapamycin treatment alleviated the growth defects and eIF2α phosphorylation in a dose-dependent manner. Therefore, the data here suggest that MPpnrsv has the ability to activate Gcn2p and Tor1p nutrient-regulated pathways. These findings uncover a previously uncharacterized function of MPpnrsv, and highlight Tor1p and Gcn2p kinases as candidate susceptibility factors for plant viral infections.

It is widely accepted that the main function of MPs is to assist virus in cell-to-cell movement across plant tissues [17]. In addition, plant virus MPs have been shown to be pathogenicity determinants in some virus-host interactions and in some instances cause developmental and metabolic alterations [26]. The increasing number of host virus factors interacting with viral MPs [20,21] indicate their relevant role in the viral infection process. The demonstration that the MPpnrsv is able to increase eIF2α phosphorylation in a Tor1p-dependent manner might help to uncover an intriguing function of plant virus MP in this relevant cellular route. Our data indicate that MPpnrsv expression increases eIF2α phosphorylation by activating the Gcn2p kinase (Figure 2A, B). How Gcn2p is activated by MPpnrsv remains to be determined. Interestingly, the hydrophobic region (HR) of the protein seems to be required for Gcn2p activation. It is worth noting that the HR of the PNRSV MP has been shown to be essential for a complete viral life cycle [34]. Gcn2p activation was shown to be dependent on the nutrient-regulated kinase Tor1p. Tor1p integrates nutrient and growth factor signals to regulate the translation initiation of mRNAs important for cell growth and proliferation. Although still controversial, Tor1p seems to favour selective translation of TOP mRNAs (a class of mRNAs bearing 5'-terminal oligopyrimidine tracts which encode components of the translational machinery) by signaling through ribosomal protein S6 (rpS6) kinase (S6K1) to rpS6. [38,39]. As genomic PNRSV RNAs could be considered as TOP-like RNAs [30,40], it is tempting to speculate that MPpnrsv could enhance selective viral mRNA expression and viral replication by activating the Tor1p pathway, similarly as was recently described for adenoviral E4-ORF1 and E4-ORF4 proteins [6]. Along the same line, the plant viral reinitiation factor transactivator–viroplasmin (TAV) has been shown to recruit the TOR/S6K1 signaling complex, with TAV–TOR binding being critical for both translation re-initiation and viral fitness [41].

The functional linkage that connects Tor1p with Gen2p activation by MPpnrsv expression remains unanswered. Recent studies in yeast have found that the GCN pathway is a major effector of the TOR pathway [42]. Under nutrient stress, Gcn2p was induced by the release of TOR repression through a mechanism involving Sit4p protein phosphatase. This mechanism might well be triggered by expression of MPpnrsv in yeast. Indeed, the functional categories enriched in transcriptome of yeast expressing MPpnrsv resembled those of yeast cells subjected to nutrient stress [42]. Alternatively, constitutive hyper stimulation of Tor1p by MPpnrsv expression regardless the nutrient status, would favor precocious cell entry into S-phase thus leading to replicative stress. It is well established that Gcn2p negatively regulates cell cycle progression in response to replicative stress [43,44]. The growth interference phenotype elicited by MPpnrsv, could then be interpreted as a consequence of the cell cycle delay triggered by Gcn2p activation. Indeed, exponential yeast cultures overexpressing MPpnrsv exhibited an increased number of un budded cells, thus reflecting a delay at the G1/S transition (Table S4). In addition, the fact that rapamycin treatment relieved the MPpnrsv-dependent growth defect in yeast is also consistent with this hypothesis. We cannot discard as well, that the activation of the two opposite pathways, Tor1p and Gcn2p, could be considered a compensative effect derived from the cell location of the viral MPs. It is well known that all eukaryotes respond to ER stress through a set of pathways known as the unfolded protein response (UPR) [45,46] which include the eIF2α phosphorylation. The up-regulation of the Tor1p pathway could be response to mitigate the UPR triggered by the viral MPs that are associated to the membrane of ER, avoiding the transient decrease in global translation. In this sense, we do not expect a growth interference phenotype using viral MP that are not located to the ER as we observed with the MPpnrsvAH or the MP of GFLV.

Taken together, our results point out at Tor1p and Gcn2p kinases as candidate susceptibility host factors during plant viral infections. Considering that both kinases are conserved throughout evolution, experimental validation of these candidate factors in a plant model of viral infection would further support this possibility, as recently described for other viral factors[41]. To test this hypothesis, we are currently analyzing the phosphorylation status of Gcn2p/eIF2α and TOR/S6K1 signaling proteins in transgenic A. thaliana plants overexpressing MPpnrsv. Successful completion of these experiments would definitively open a new research area on viral MPs functions and plant-virus interactions.

Methods

Strains, plasmids, media and general methods

Specific primers were used to amplify the MP of PNRSV, mutant lacking its hydrophobic domain (MPpnrsvΔHR) and MPs of BMV, CMV, TMV and GFLV (see Table S5). PCR products were inserted between Pme1–PstI sites of the centromeric plasmid pCM253 to express the corresponding proteins in yeast cells. In all these constructs, ORF is in frame with three copies of the HA plus a histidine epitope. Plasmids were confirmed by DNA sequencing. All yeast strains used in this study are listed in Table S6. Standard methods for yeast culture and manipulations were used (minimal medium (SD) contained 2% glucose, 0.67% yeast nitrogen base and the appropriate amino acid supplements to maintain selection of URA3 and cultures were routinely grown at 30°C). In growth curve analyses, cells were grown in microtiter plates using the Bioscreen C system (Thermo) with or without Doxycycline. For cycloheximide and rapamycin experiments, cells were grown in presence of 100 and 200 μg/ml of each drug, respectively.

Immunoblotting analysis

Yeast strains were grown in minimal medium without Doxycycline. At the indicated times cells were centrifuged. Cell pellets were then resuspended in 5X Laemmlı buffer and boiled for 10 min. Total cellular protein was subjected to SDS Polyacryl- amide Gel Electrophoresis (SDS-PAGE) and transferred to PVDF membranes (Amersham). Viral proteins were detected with anti HA-antibody (ROCHE) whereas a specific phospho-eIF2α (Ser51)
antibody (Cell Signalling) was used to analyze the phosphorylation state of eIF2α. In all cases, western-blots were carried out following manufacturer’s recommendations. Immunocomplexes were visualized by enhanced chemiluminescence detection (ECL Amersham) using anti-Rabbit IgG HRP conjugated (Amersham). Uniform gel loading was confirmed by Coomassie Blue staining of membranes after immunoblot studies.

### Microarray assay

Yeast cell cultures were grown 24 h in minimal medium with doxycycline (100 mg/ml). These cultures were then pelleted and diluted to an OD600 nm of 0.1 into fresh minimal medium without doxycycline. At 6 h total RNA was extracted from yeast cells transformed with the empty pCM262 vector (pCM construct), pCM:MPpnrsv and pCM:MPpnrsvAHR. RNA samples were amplified using Message Amp II amplification kit (Ambion). 7.5 μg of UTP-aminomally-templated RNA (aRNA) were labelled using Cy3or Cy5 dye (GE Healthcare), purified using MegaClear columns (Ambion), and quantified using Nanodrop spectrophotometer. 150 pmol of labeled-aRNA were dried and resuspended in hybridization buffer containing 3xSSC, 0.1% SDS, 0.1% salmon sperm DNA and 50% formamide. Microarray hybridization was performed manually using Telechem Hybridization Chambers, following Corning instructions. Slides were scanned using a GenePix 4000B scanner and analyzed using GenePix 6.0 software (Molecular Devices). Data were normalized by mean global intensity and Lowess correction. Significance analysis of microarrays (SAM) [47] was performed on the pre-processed and normalized data sets to identify differentially expressed genes. A 5% false discovery rate (FDR) and 2-fold expression cut off was considered to determine up-regulated and down-regulated genes. A functional category analysis of the up-regulated or down-regulated genes was carried out using The GO Term Finder utility (www.http://go.princeton.edu/cgi-bin/GOTermFinder). Enriched categories with p-values smaller than 0.05 were further considered. The raw data discussed in this publication is MIAME compliant has been deposited in NCBI’s Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo/) and are accessible through the GEO Series accession numbers GSE26119 and GSE26120, respectively.

### References

1. Hinnebusch, AG (2005) Translational regulation of Gcn4 and the general amino acid control of yeast. Ann Rev of Microbiology 59: 407–450.

2. Kolos RJ, Hauschka AG (1993) Translation of the yeast transcriptional activator GCN4 is stimulated by purine limitation: Implications for activation of the protein kinase GCN2. Mol Cell Biol 13: 5099–111.

3. Natarajan K, Meyer MR, Jackson BM, Slade D, Roberts C, et al. (2001) Transcriptional profiling shows that Gcr1p is a master regulator of gene expression during amino acid starvation in yeast. Mol Cell Biol 21: 4347–68.

4. Zhang Y, Dickinson JR, Paul MJ, Halford NG (2003) Molecular cloning of an Arabidopsis homologue of GCN2, a protein kinase involved in co-ordinated response to amino acid starvation. Planta 217: 660–673.

5. Halford, NG (2006) Regulation of carbon and amino acid metabolism, roles of sucrose nonfermenting-1-related protein kinase-1 and general control non- derepressible-2-related protein kinase. Ad Bot Res 43: 93–142.

6. O’Shea CK, Khosch S, Choi B, Bagus C, Soria J, et al. (2005) Adenoviral proteins mimic nutrient/growth signals to activate the mTOR pathway for viral replication. EMBO J 24: 1211–1221.

7. Proud GG (2003) eIF2 and the control of cell physiology. Semin Cell Dev Biol 14: 3–12.

8. He B, Gross M, Roisman B (1997) The c 1345 protein of herpes simplex virus 1 complexes with protein phosphatase 1a to dephosphorylate the alpha subunit of the eukaryotic translation initiation factor 2 and preclude the shutdown of protein synthesis by double-stranded RNA activated protein kinase. Proc Natl Acad Sci U S A 94: 443–446.

9. Qian W, Zhu S, Sobolev AY, Wek RC (1996) Expression of Vaccinia Virus K3L Protein in Yeast Inhibits Eukaryotic Initiation Factor-2 Kinase Gcn2 and the General Amino Acid Control Pathway. J Biol Chem 271: 13202–13207.

10. Montero H, Rojas M, Arias CA, Lopez S (2006) Rotavirus Infection Induces the Phosphorylation of eIF2a but Prevents the Formation of Stress Granules. J Virol 80: 1496–1504.

11. Rojas M, Arias CF, Lopez S (2010) Protein Kinase R Is Responsible for the Phosphorylation of eIF2a in Rotavirus Infection. J Virol 84: 10457–10466.

12. Lagrix S, Lamer E, Pouch-Pélissier MN, Espagnol MC, Robaglia C, et al. (2008) Arabidopsis eIF2α kinase GCN2 is essential for growth in stress conditions and is activated by wounding. BMC Plant Biol 8: 134.

13. Zhang Y, Wang Y, Kanyuka K, Parry MAJ, Powers JS, et al. (2008) GCN2-dependent phosphorylation of eukaryotic translation initiation factor-2α in Arabidopsis. J Exp Botany 59: 3131–3141.

14. Bilgin DD, Liu Y, Schiff M, Dinesh-Kumar SP (2005) P53IPK, a plant ortholog of double-stranded RNA-dependent protein kinase PKR inhibitor, functions in viral pathogenesis. Dev Cell 4: 651–661.

15. Fernandez-Calvino L, Faulkner C, Maule A (2011) Plasmodesmata as Active Conduits for Virus Cell-to-Cell Movement. In: Recent Advances in Plant Virology Caranta C, Aranda MA, Teplier M, Lopez-Moya JJ, eds. Caister Academic Press. ISBN: 978-1-904455-75-2. 470 p.
16. Pallas V, Genoves A, Sánchez-Pina MA, Navarro JA (2011) Systemic movement of viruses via the plant phloem. In: Recent Advances in Plant Virology Caranta C, Aranda MA, Teferri M, Lopez-Moya JJ, eds. Caister Academic Press. ISBN: 978-1-904455-75-2. 470 p.
17. Wainmann E, Ueki S, Trumyeva K, Citovsky V (2004) The ins and outs of nondestructive cell-to-cell and systemic movement of plant viruses. Crit Rev Plant Sci 23: 193–250.
18. Melcher U (2000) The “30K” superfamily of viral movement proteins. J Gen Virol 81: 257–266.
19. Whitham WJ (2006) Plant viral movement proteins: Agents for cell-to-cell trafficking of viral genomes. Virology 344: 169–184.
20. Curin M, Ojangu EL, Trutnyeva K, Llau B, Truve E, et al. (2007) MPB2C, a microtubule-associated plant factor, is required for microtubular accumulation of tobacco mosaic virus movement protein in plants. Plant Physiol 143: 801–811.
21. Kaido M, Inoue Y, Takeda Y, Takeda A, Mori M, et al. (2007) Downregulation of the NtNACa1 gene encoding a movement-protein-interacting protein reduces cell-to-cell movement of brome mosaic virus in Nicotiana benthamiana. Mol Plant-Microbe Interact 20: 671–681.
22. Lucas WJ (2006) Plant viral movement proteins: Agents for cell-to-cell trafficking of viral genomes. Virology 344: 169–184.
23. Melcher U (2000) The “30K” superfamily of viral movement proteins. J Gen Virol 81: 257–266.
24. Sasaki N, Ogata T, Deguchi M, Nagai S, Tamai A, et al. (2009) Over-expression of putative transcriptional coactivator KELP interferes with Tomato mosaic virus cell-to-cell movement. Mol Plant Pathol 10: 161–173.
25. Shinizu T, Yoshii A, Sakurai K, Hamada K, Yamaji Y, et al. (2009) Identification of a novel tobacco DnaJ-like protein that interacts with the movement protein of tobacco mosaic virus. Arch Virol 154: 959–967.
26. Curin M, Ojangu EL, Trutnyeva K, Llau B, Truve E, et al. (2007) MPB2C, a microtubule-associated plant factor, is required for microtubular accumulation of tobacco mosaic virus movement protein in plants. Plant Physiol 143: 801–811.
27. Kaido M, Inoue Y, Takeda Y, Takeda A, Mori M, et al. (2007) Downregulation of the NtNACa1 gene encoding a movement-protein-interacting protein reduces cell-to-cell movement of brome mosaic virus in Nicotiana benthamiana. Mol Plant-Microbe Interact 20: 671–681.
28. Aparicio F, Vilar M, Perez-Paya E, Pallas V (2003) The coat protein of Prunus necrotic ringspot virus. Arch Virol 153: 909–919.
29. Aparicio F, Vilar M, Perez-Paya E, Pallas V (2003) The coat protein of Prunus necrotic ringspot virus. Arch Virol 153: 909–919.
30. Aparicio F, Pallas V (2002) Molecular variability analysis of the RNA 3 of fifteen isolates of Prunus necrotic ringspot virus. Arch Virol 154: 959–967.
31. Herranz MC, Pallas V (2004) RNA-binding properties and mapping of RNA-binding domain from the movement protein of Prunus necrotic ringspot virus. J Gen Virol 85: 761–768.
32. Fiore N, Fajardo TVM, Prodan S, Herranz MC, Aparicio F, et al. (2008) Genetic diversity of the movement and coat protein genes of South American isolates of Prunus necrotic ringspot virus. Arch Virol 153: 909–919.
33. Herranz MC, Sanchez-Navarro JA, Sauri A, Mingarro I, Pallas V (2005) Mutational analysis of the RNA-binding domain of the Prunus necrotic ringspot virus (PNRSV) movement protein reveals its requirement for cell-to-cell movement. Virology 339: 31–41.
34. Martinez-Gil L, Sanchez-Navarro JA, Cruz A, Pallas V, Perez-Gil J, et al. (2009) Plant Virus Cell-to-Cell Movement Is Not Dependent on the Transmembrane Disposition of Its Movement Protein. J Virol 83: 5535–5543.
35. Alves-Rodrigues I, Galao RP, Meyerhans A, Diez J (2006) Saccharomyces cerevisiae: A useful model host to study fundamental biology of viral replication. Virus Research 120: 49–56.
36. Nagy PD (2008) Yeast as a Model Host to Explore Plant Virus-Host Interactions. Annu Rev Phytopathol 46: 217–42.
37. Steffensen L, Pedersen PA (2006) Heterologous expression of membrane and soluble proteins derepresses GCN4 mRNA translation in the yeast Saccharo-
myces cerevisiae. Eukaryot Cell 5: 248–61.
38. Duhrer A, Thomas, G (1999) Ribosomal S6 kinase signaling and the control of translation. Exp Cell Res 253: 100–109.
39. Reiter AK, Anthony TG, Anthony JG, Jefferson LS, Kimball SR (2004) The mTOR signaling pathway mediates control of ribosomal protein mRNA translation in rat liver. Int J Biochem Cell Biol 36: 2169–79.
40. Di Terlizzi B, Skrzeczkowski L, Mink GI, Scott SW, Zimmerman MT (2001) The RNA 5 of Prunus necrotic ringspot virus is a biologically inactive copy of the 3’-UTR of the genomic RNA 3. Arch Virol 146: 825–833.
41. Schepetilnikov M, Kobayashi K, Geldreich E, Curin M, Obbard G, et al. (2011) Viral factor TAV recruits TOR/S6K1 signalling to activate remittance after long ORF translation. EMBO J 30: 1343–56.
42. Staschke KA, Dey S, Zaborske JM, Palam LR, McClintick JN, et al. (2010) Integration of general amino acid control and target of rapamycin (TOR) regulatory pathways in nitrogen assimilation in yeast. J Biol Chem 285: 14595–911.
43. Menacho-Marquez M, Perez-Valle J, Arino J, Gadea J, Murguia JR (2007) Gen2p regulates a G1/S cell cycle checkpoint in response to DNA damage. Cell Cycle 6: 2302–2305.
44. Tvegard T, Solheim H, Skjelberg HC, Krohn M, Nilsen EA, et al. (2007) A novel checkpoint mechanism regulating the G1/S transition. Genes Dev 21: 649–54.
45. Kaufman RJ (1999) Stress signaling from the lumen of the endoplasmic reticulum: coordination of gene transcriptional and translational controls. Genes Dev 13: 1211–1233.
46. Harding HP, Calfon M, Urano F, Novoa I, Ron D (2002) Transcriptional and translational control in the Mammalian unfolded protein response. Annu Rev Cell Dev Biol 18: 575–599.
47. Schepetilnikov M, Kobayashi K, Geldreich A, Curin M, Obbard G, et al. (2011) Viral factor TAV recruits TOR/S6K1 signalling to activate remittance after long ORF translation. EMBO J 30: 1343–56.
48. Tusher VG, Tibshirani R, Chu G (2001) Significance analysis of microarrays applied to the ionizing radiation response. Proc Natl Acad Sci U S A 98: 5116–5121.