Co-opting ATP-generating glycolytic enzyme PGK1 phosphoglycerate kinase facilitates the assembly of viral replicase complexes

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Abstract

The intricate interactions between viruses and hosts include exploitation of host cells for viral replication by using many cellular resources, metabolites and energy. Tomato bushy stunt virus (TBSV), similar to other (+)RNA viruses, induces major changes in infected cells that lead to the formation of large replication compartments consisting of aggregated peroxisomal and ER membranes. Yet, it is not known how TBSV obtains the energy to fuel these energy-consuming processes. In the current work, the authors discovered that TBSV co-opts the glycolytic ATP-generating Pgk1 phosphoglycerate kinase to facilitate the assembly of new viral replicase complexes. The recruitment of Pgk1 into the viral replication compartment is through direct interaction with the viral replication proteins. Altogether, we provide evidence that the ATP generated locally within the replication compartment by the co-opted Pgk1 is used to fuel the ATP requirement of co-opted heat shock protein 70 (Hsp70) chaperone, which is essential for the assembly of new viral replicase complexes and the activation of functional viral RNA-dependent RNA polymerase. The advantage of direct recruitment of Pgk1 into the virus replication compartment could be that the virus replicase assembly does not need to intensively compete with cellular processes for access to ATP. In addition, local production of ATP within the replication compartment could greatly facilitate the efficiency of Hsp70-driven replicase assembly by providing high ATP concentration within the replication compartment.

Author summary

Positive-strand (+)RNA viruses build replication organelles with the help of many co-opted cellular factors and by usurping cellular metabolites and energy, which frequently lead to disease states in plants, animals and humans. The authors discovered that tomato bushy stunt virus (TBSV) co-opts the glycolytic ATP-generating Pgk1 phosphoglycerate kinase to facilitate the intracellular assembly of new viral replicase complexes. The ATP generated by co-opted Pgk1 within the replication compartment is used to fuel the ATP requirement of co-opted cellular heat shock protein 70 chaperone, which is essential for
the assembly of new TBSV replicase complexes. Direct recruitment of Pgk1 by TBSV might provide easy access to ATP. Metabolic reprogramming of the glycolytic pathway might be a widespread phenomenon during various viral infections.

Introduction

Positive-strand (+)RNA viruses build robust viral replication machineries, called replication organelles, with the help of many co-opted cellular factors. In addition, viruses also obtain metabolites and energy from the infected cells. Overall, (+)RNA viruses induce major metabolic and structural changes in the infected cells, which frequently lead to disease states [1,2,3,4,5].

One of the best characterized (+)RNA virus is TBSV, which induces large replication compartments consisting of aggregated peroxisomal and ER membranes. The replication compartments contain numerous vesicle-like structures in the limiting membrane of peroxisomes, which harbor the viral replicase [6,7,8]. TBSV also manipulates the cytoskeleton and endosomal trafficking [9,10]. TBSV co-opts numerous host proteins to support various viral functions, including the heat shock protein 70 (Hsp70), the endosomal sorting complex required for transport (ESCRT) machinery, translation factors and DEAD-box RNA helicases [11,12,13,14,15,16,17]. TBSV also induces enrichment of sterols at the viral replication sites via co-opting oxysterol-binding proteins and membrane contact sites and retargets phosphatidylethanolamine to the replication sites to build suitable membranous subcellular environment for replication [8,9,18]. TBSV-induced major changes in the cell require energy, but how TBSV can obtain the energy to fuel these energy-demanding processes is not known.

Glycolysis is an essential and highly conserved energy-producing pathway in the cytosol. Glucose is converted into pyruvate by a ten-enzyme catalyzed reaction that produces ATP and NADPH. The two ATP-generating enzymes in the glycolytic pathway are Pgk1 phosphoglycerate kinase and Cdc19 pyruvate kinase (PKM2/PKLR in humans). Pgk1 produces ATP from ADP and DPG using substrate-level phosphorylation in the cytosol [19]. Pgk1 is highly conserved and present in every organism. In humans, Pgk1 is also important for tumor growth and DNA replication/repair and mutations in Pgk1 are associated with myopathy, mental disorders and hemolytic anemia [20,21,22]. Pgk1 is also important for the activation of nucleotide-based anti-HIV drugs [23].

Previous genome-wide screens in yeast model host have identified several components of the glycolytic pathway that affected TBSV replication [24,25,26]. Proteomic screens also identified Pgk1 phosphoglycerate kinase, which was pulled-down with the tombusviral p92 RdRp and interacted with p33 replication protein [27,28]. Therefore, we decided to characterize the putative role of Pgk1 in tombusvirus replication in this work. We discovered that TBSV co-opts the glycolytic Pgk1 through direct interaction with the viral replication proteins. Altogether, we provide evidence that the ATP generated locally in the viral replication compartment by the co-opted Pgk1 is used to fuel the ATP requirement of the co-opted cellular Hsp70 molecular chaperones. The functions of co-opted Hsp70 are to facilitate the assembly of new viral replicase complexes and the activation of the viral RNA-dependent RNA polymerase. The local production of ATP by Pgk1 within the replication compartment could greatly facilitate the efficiency of Hsp70-driven replicase assembly by providing high ATP concentration within the replication compartment.
Results
Recruitment of the cytosolic Pgk1 phosphoglycerate kinase into the tombusvirus replication compartment in yeast and plant cells

Among the surprises from high-throughput screens with TBSV and yeast model host were the identification of the glycolytic Pgk1 as a putative co-opted host factor in TBSV replication [24,25,26,29]. To demonstrate interaction between Pgk1 and the viral replication proteins, first we performed co-purification experiments from yeast replicating TBSV repRNA. The isolated membrane fraction was solubilized with detergent and the FLAG-tagged p33 and FLAG-p92pol replication proteins were affinity-purified, followed by Western-blot analysis. We found that the His6-tagged Pgk1 expressed from a plasmid was co-purified with the tombusvirus replicase (Fig 1A, lane 2). In contrast, Pgk1 was not co-purified with Flag-GFP control from yeast (S1 Fig). To examine if Pgk1 is a permanent component of the tombusvirus replicase, we halted the formation of new tombusvirus replicase complexes by shutting down p33 and p92pol expression and stopping ribosomal translation by adding cycloheximide [7]. FLAG-affinity-purification of the tombusvirus replicase from the membrane fraction of yeast at various time-points showed the rapid release of Pgk1 from the replicase (Fig 1B, lanes 3–4 versus

Fig 1. Interaction between p33 and p92pol replication proteins and yeast cytosolic Pgk1. (A) Co-purification of Pgk1 with the viral replicase. Top panel: Western blot analysis of co-purified His6-tagged Pgk1 with FLAG-affinity purified FLAG-p33 and FLAG-p92pol from membrane fraction of yeast. Pgk1 was detected with anti-His antibody. The negative control was His6-tagged p33 and His6-p92 purified from yeast extracts using a FLAG-affinity column. Middle panel: Western blot analysis of purified FLAG-p33 detected with anti-FLAG antibody. Asterisk marks the SDS-resistant p33 homodimer. Bottom panel: Western blot of His6-tagged Pgk1 and His6-p33 (lane 1) proteins in the total yeast extracts using anti-His antibody. (B) Testing the presence of Pgk1 in the membrane-bound viral replicase after blocking cellular translation by cycloheximide. Top panel: Western blot analysis shows the co-purified His6-tagged Pgk1 with the viral replicase isolated from membrane fraction at the shown time points. See further details in panel A. Middle panel: Western blot analysis of the purified FLAG-p33 with anti-FLAG antibody. Bottom panel: Western blot analysis of His6-Pgk1 in the total yeast lysates with anti-His antibody. Each experiment was repeated three times.

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2). Thus, Pgk1 seems to be a temporarily co-opted factor in the tombusvirus replicase complex.

To obtain additional evidence of subversion of Pgk1 into the tombusvirus replication compartment, we performed confocal laser microscopy experiments in yeast expressing GFP-p33 and BFP-Pgk1. Interestingly, we observed the robust recruitment of the cytosolic Pgk1 into the tombusvirus replication compartment, which was marked by Pex13-RFP peroxisomal marker protein (Fig 2A). Similar experiments with ectopically expressed BFP-NbPgk1 and p92-YFP showed the enrichment of Pgk1 in the tombusvirus replication compartment in *Nicotiana benthamiana* cells replicating the TBSV repRNA (Fig 2B). In addition, bimolecular fluorescence complementation (BiFC) assay further confirmed the recruitment of the glycolytic NbPgk1 via interaction with p33 into the peroxisomal TBSV replication compartment (Fig 2C). Similarly, we observed that the interaction between TBSV p92pol replication protein and the glycolytic NbPgk1 takes place in the large replication compartment, which was marked by RFP-SKL peroxisomal luminal marker protein (Fig 2C, lower panels). Therefore, we conclude that TBSV recruits the cytosolic Pgk1 through direct interaction with the viral replication proteins into the viral replication compartment in both yeast and plant cells.

**Pgk1 acts as a pro-viral host factor in yeast**

To test if the recruitment of Pgk1 into the viral replication compartment plays a role in TBSV replication, we have made a haploid yeast strain, in which the wt *PGK1* gene was replaced with the HA-tagged *PGK1*. In addition, expression of HA-*PGK1* was placed under the regulation of GALS promoter, thus allowing induction by the addition of galactose and repression by addition of glucose to the culture media [30]. Induction of TBSV repRNA replication in GALS::*PGK1* yeast with suppressed *PGK1* expression showed ~4-fold reduction in TBSV repRNA replication (Fig 3A, lanes 7–9 versus 10–12, and S2 Fig). Expression of Pgk1 from a plasmid in GALS::*PGK1* yeast with suppressed Pgk1 expression from chromosome increased TBSV repRNA replication by ~2-fold in glucose-containing media, while did not increase it on the non-fermentable glycerol-media (Fig 3B, lanes 7–10). These data indicate that Pgk1 plays a pro-viral function during TBSV replication in yeast.

Over-expression of either yeast Pgk1 (Fig 3C) or the *N. benthamiana* cytosolic NbPgk1 (Fig 3D) led to ~3-fold and a ~2.5-fold increase, respectively, in TBSV repRNA accumulation in wt yeast, confirming a pro-viral function for Pgk1 in TBSV replication.

**Pgk1 also acts as a pro-viral host factor in plant cells**

Estimation of *PGK1* mRNA levels in TBSV-infected *N. benthamiana* leaves indicated ~2-fold increased level of expression (Fig 4A). Similarly, Western blotting demonstrated a ~50% increase in accumulation of Pgk1 protein in yeast cells replicating TBSV repRNA (Fig 4B), suggesting that a tombusvirus can mildly induce the accumulation of a glycolytic enzyme in cells.

To confirm the importance of Pgk1 in TBSV replication in a plant host, we knocked-down via virus-induced gene silencing (VIGS) the expression of the cytosolic Pgk1 in *N. benthamiana* and the Pgk1-silenced leaves were inoculated with the full-length infectious TBSV. TBSV genomic RNA accumulation was inhibited by ~4-fold in these Pgk1 knock-down plants (Fig 4C). Inoculation of Pgk1-silenced *N. benthamiana* with the similar peroxisome-replicating cucumber necrosis virus (CNV) (S3 Fig) and the mitochondrial-replicating carnation Italian ringspot virus (CIRV) showed a ~6-fold decrease in the accumulation of genomic RNAs (Fig 4D and 4E). Thus, the cytosolic Pgk1 is a critical pro-viral host factor for various tombusviruses in a plant host.
Glycolytic enzyme facilitates the assembly of viral replicase complexe

A

B

C. BiFC

p33-cYFP+ nYFP-Pgk1

RFP-SKL

merged

DIC

p33-cYFP+ nYFP-MBP

RFP-SKL

merged

DIC

p92-cYFP+ nYFP-Pgk1

RFP-SKL

merged

DIC

p92-cYFP+ nYFP-MBP

RFP-SKL

merged

DIC
Over-expression of NbPgk1 in N. benthamiana replicating TBSV RNA showed a close to 3-fold enhanced TBSV accumulation (Fig 4F), suggesting that Pgk1 level affects tombusvirus replication. Altogether, these data have confirmed the pro-viral function of Pgk1 in tombusvirus replication in plants.

Knock-down of Pgk1 level decreases the local accumulation of ATP within the tombusvirus replication compartment

Based on the above results, we assumed that the co-opted Pgk1 likely generates ATP within the viral replication compartment. Therefore, we estimated the local ATP level within the replication compartment by using the p33 replication protein tagged with ATeam, a cellular ATP-sensor module. ATeam module can measure ATP levels via FRET due to the conformational change in the enhanced ATP-binding domain of the bacterial ATP synthase upon binding to ATP without ATP hydrolysis (Fig 5A) [31]. In the ATP-bound form, the ATP-sensor module brings the CFP and YFP fluorescent tags into proximity, increasing FRET, which can be detected by confocal laser microscopy. In the ATP-free stage, the extended conformation of the ATP-sensor module places CFP and YFP at distal position, leading to low FRET signal. The ATeam-tagged p33 localizes to the aggregated peroxisomes that represent the sites of replication (Fig 5B).

We found by intracellular expression of p33-ATeam that the local ATP level within the TBSV replication compartment was decreased by ~6-fold in Pgk1 knock-down in N. benthamiana leaves in comparison with control leaves (Fig 5B). Interestingly, the mitochondria-replicating CIRV also accumulated a ~10-fold lower level ATP within the replication compartment in Pgk1 knock-down leaves than in control leaves (Fig 5C). Therefore, we conclude that Pgk1 is recruited to the sites of tombusvirus replication to generate high concentration of ATP within the replication compartment.

The co-opted Pgk1 supports viral replicase complex assembly within the tombusvirus replication compartment

To decipher the function of the co-opted Pgk1 during TBSV replication, we tested viral RNA levels in yeast replicating TBSV repRNA. Down-regulation of Pgk1 expression reduced both (+)- and (-)RNA levels by ~4-fold (Fig 6A). These data suggest that Pgk1 likely plays an early role, prior to viral RNA synthesis, possibly during replicase assembly steps.

To test this scenario, we performed various in vitro assays. First, to further examine if Pgk1 was required for viral RNA replication, we obtained cell-free extracts (CFE) from wt yeast and
GALS::PGK1 yeast with suppressed Pgk1 expression to reconstitute the TBSV replicase in vitro based on purified recombinant TBSV proteins. The yeast CFE-based assay, which supports a single full cycle of viral RNA replication [32], showed ~7-to-10-fold reduced repRNA production when low level of Pgk1 was produced in GALS::PGK1 yeast (Fig 6B, lanes 3 versus 1 and 4, also S4 Fig). Interestingly, both (+)RNA product (Fig 6B) and dsRNA replication intermediate obtained with CFE from GALS::PGK1 yeast with depleted Pgk1 in comparison with the wt yeast CFE were decreased by ~7-fold (Fig 6C, lanes 7–8 versus 3–4). Therefore, the in vitro replication results support the important role of Pgk1 in viral RNA replication.
We also obtained purified replicate preparations from GALS::PGK1 yeast with suppressed Pgk1 expression versus comparable preparations from wt yeast, which were programmed with viral RNA transcripts [33]. In spite of having comparable amounts of replication proteins, the

Fig 4. The pro-viral role of cytosolic Pgk1 in tombusvirus replication in *N. benthamiana*. (A) Up-regulation of Pgk1 expression in TBSV-infected *N. benthamiana* leaves. The mRNA levels for the cytosolic Pgk1 were estimated by semi-quantitative RT-PCR (25 and 28 cycles, the latter is shown) in total RNA samples obtained from either TBSV or mock-infected *N. benthamiana* leaves. Tubulin mRNA was used as a control (second panel). The bottom panel shows an ethidium-bromide stained agarose gel of total RNA samples with ribosomal RNA and TBSV genomic (g)RNA. (B) Western blot analysis of cytosolic Pgk1 level in total protein samples obtained from yeast either supporting TBSV repRNA accumulation or TBSV-free using anti-Pgk1 antibody. (C-E) Knock-down of PGK1 mRNA level by VIGS inhibits the accumulation of tombusvirus RNA in *N. benthamiana*. Top panels: Total RNA samples obtained from *N. benthamiana* leaves silenced as shown were analyzed by Northern blotting, which shows the accumulation of TBSV gRNA and sgRNAs in panel C, the closely-related cucumber necrosis virus (CNV) RNAs in panel D and the mitochondria-replicating CIRV RNAs in panel E. Bottom images: ethidium-bromide stained gels show ribosomal RNA level. We chose the 12th day after VIGS to inoculate the upper, systemically-silenced leaves with TBSV virions, or agroinfiltrate with pGD-CN V or pGD-CIRV. Samples for RNA extractions were taken 1 day (TBSV) and 2.5 days (CNV or CIRV) post inoculation from the inoculated leaves. The control experiments included the TRV2-cGFP vector. Each experiment was performed three times. (F) Over-expression of the cytosolic NbPgk1 was done in *N. benthamiana* leaves by agroinfiltration. The same leaves, which were first agroinfiltrated with pGD-NbPgk1 (expressing NbPgk1 from the 35S promoter), were also inoculated with TBSV virions 2.5 days later. Then, total RNA samples were obtained the subsequent day (1 dpi). The control samples were obtained from leaves agroinfiltrated with pGD empty vector (not expressing proteins) (lanes 1–6). Northern blotting was used to detect the accumulation of TBSV RNAs in total RNA samples obtained from *N. benthamiana* leaves. The ribosomal RNA (rRNA) was used as a loading control and shown in agarose gel stained with ethidium-bromide (bottom panel). The bottom image shows a representative Western blot-based detection of His<sub>6</sub>-tagged NbPgk1 using anti-His antibody. Each experiment was performed three times.

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Fig 5. The cytosolic Pgk1 affects ATP accumulation within the tombusvirus replication compartment in *N. benthamiana*. (A) A scheme of the FRET-based detection of cellular ATP within the replication compartment. The enhanced ATP biosensor, ATeam<sup>YEMK</sup> was fused to TBSV p33 replication protein. (B) Knock-down of *PGK1* mRNA level by VIGS in *N. benthamiana* was
purified replicase preparations from Pgk1 depleted yeast showed ~3-fold reduced activity in comparison with the similar preparations obtained from wt yeast (Fig 6D).

Since the activity of the purified replicase preparations mostly depends on the efficiency of replication assembly and replication protein activation, we compared the efficiency of p92pol activation by the supernatant fraction of CFEs prepared from Pgk1 depleted or wt yeasts. These assay revealed ~3-fold less RdRp activity of the purified recombinant p92pol mutant by the preparation from Pgk1 depleted than from wt yeasts (compare lanes 1–2 and 5–6 in Fig 6E). The major active component of the supernatant fraction of CFEs is Hsp70 molecular chaperone, which is essential for the activation of p92pol RdRp in vitro [29]. Accordingly, addition of purified Ssa1 (yeast Hsp70) to the WT fraction has increased the efficiency of p92pol RdRp mutant activation by ~5.5-fold, whereas the comparable assay with Ssa1 involving the preparation from Pgk1 depleted yeast showed only ~2-fold enhancement (compare lanes 3–4 and 7–8 in Fig 6E). Therefore, the in vitro results from multiple assays support the model that Pgk1 has a major role in tombusvirus RNA replication and Pgk1 likely fuels the energy requirements of co-opted Hsp70 molecular chaperones to facilitate the efficient replicase assembly and replication protein activation within the viral replication compartment.

TBSV co-opts Vps4p AAA+ ATPase, which is an ATP-dependent host protein, into the viral replicase complex to facilitate replicase assembly [7]. Vps4p is an ESCRT protein involved in membrane deformation, which is necessary for formation of TBSV-induced individual spherules (vesicle-like structures) supporting viral RNA replication [7]. The spherule formation protects the viral dsRNA replication intermediate from the innate RNAi machinery during infection [34]. To test if Pgk1-produced ATP might support Vps4p ATPase function during replicase assembly, we utilized a recently developed intracellular probe based on a reconstituted RNAi system in yeast (Saccharomyces cerevisiae), which lacks the RNAi machinery during infection [34]. We found that the induction of RNAi activity had only slightly more inhibitory effect on TBSV RNA accumulation in Pgk1-depleted yeast than in wt yeast (Fig 6F). Based on these results, the structure of the assembled tombusvirus replicase in Pgk1 depleted yeast might not be different from those assembled in wt yeast. Therefore, likely the co-opted ESCRT machinery and the Vps4p ATPase activity might not be affected when Pgk1 is depleted. Alternatively, the residual Pgk1 still present in GAL5::PGK1 yeast could provide enough ATP for Vps4p ATPase activity during the replicase assembly process.

Discussion

TBSV replication, similar to other (+)RNA viruses, is a very intensive and robust process that likely depends on consumption of a large amount of ATP. By recruiting the glycolytic Pgk1 into the virus replication compartment, Pgk1 could efficiently supply ATP to facilitate various steps in the replication process, including the assembly of the viral replicase complex. Accordingly, we show that TBSV actively recruits the cytosolic Pgk1 to the sites of viral replication...
Fig 6. The co-opted function of cytosolic Pgk1 is to provide ATP for viral replicase complex assembly. (A) Northern blot analysis shows decreased TBSV (+) and (-)repRNA accumulation in a yeast strain (GALS::PGK1) when Pgk1 was depleted (raffinose) in comparison when Pgk1 expression is induced (Gal+Raf). See further details in Fig 3, panel A. (B) Inefficient activities of the tombusvirus replicase assembled in vitro in CFEs prepared from a yeast strain (GALS::PGK1) with reduced level of Pgk1 expression. Purified recombinant p33 and p92 replication proteins of TBSV and in vitro transcribed TBSV DI-72 (+)repRNA were added to the CFEs prepared from the shown yeast strains. Top panel: denaturing PAGE analysis of in vitro tombusvirus replicase activity in the CFEs. The 32P-labeled TBSV repRNA products of the reconstituted replicases are shown. Note that the full cycle of replicase activity of the in vitro reconstituted TBSV replicase depends on Pgk1 levels in the CFEs. The CFEs contained the same amounts of total yeast proteins. (C) Reduced production of 32P-labeled dsRNA intermediate, consisting of complementary (+) and (-)repRNA strands, by the in vitro reconstituted TBSV replicase when the CFEs were obtained from GALS::PGK1 yeast when Pgk1 was depleted (yeast was cultured in glucose media for 4
32P-labeled RNA products synthesized by the purified tombusvirus replicase in vitro. The in vitro assays were performed with R/I/III (-)-repRNA, and they also contained ATP/CTP/GTP and 32P-UTP. Note that the original viral template RNA in the replicase from yeast is removed during replicase solubilization/purification. (E) In efficient in vitro activation of the RdRp function of the N-terminally truncated p92rep replication protein. The soluble fraction of CFEs were obtained from GALS::Pgk1 yeast when Pgk1 was depleted (yeast was cultured in glucose media for 4 h) in comparison when Pgk1 expression is induced (yeast was cultured in galactose media for 4 h). Denaturing PAGE analysis of the 32P-labeled RNA products obtained in an in vitro assay with recombinant p92-A167N RdRp. The samples contained or lacked affinity-purified yeast Ssa1p Hsp70 protein (7 pmol). The faster migrating RNA represents a prematurely terminated product, whereas the slower migrating RNA is terminated at the 5' end of the template. Each experiment was performed three times. (F) Co-expression of S. castellii AGO1 and DCR1 in GALS::Pgk1 yeast (BY4741 background) when Pgk1 was depleted, reduces TBSV repRNA accumulation to a similar extent as in wt yeast (BY4741). Top panel: Replication of the TBSV repRNA was measured by Northern blotting 32 h after initiation of TBSV replication. The accumulation level of repRNA was normalized based on the ribosomal (r)RNA. Each sample is obtained from different yeast colonies. Yeast strain not expressing RNAi components is taken as 100% in each experiment. Average value and standard deviation is calculated from all the biological repeats. Each experiment was repeated twice. Ribosomal RNA is shown as a loading control. (G) A model on the role of the co-opted glycolytic Pgk1 in tombusvirus replication. Direct interaction of the cytosolic Pgk1 with the viral p33 and p92 replication proteins leads to recruitment of Pgk1 into the membranous replication compartment, where Pgk1 generates ATP to fuel the function of the co-opted cellular Hsp70 molecular chaperone. Hsp70 (Ssa1p and Ssa2p in yeast) has been shown to facilitate the assembly of the viral replicase complex, insertion of the replication proteins into peroxisomal membranes and activation of the RdRp function of p92rep. It seems that at least a portion of co-opted Pgk1 is released from the active viral replicase complex.

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through direct interaction with the viral replication proteins in both yeast and plant cells. It seems that a portion of recruited Pgk1 is getting released from the sites of replication when the formation of new viral replicases is halted via inhibition of translation. This observation suggests that the ATP produced by the co-opted Pgk1 is likely used up locally to fuel early steps in virus replication, such as viral replicase assembly.

The proposed role of the co-opted Pgk1 during the early steps in virus replication is further supported by additional observations. For example, down regulation of Pgk1 level in yeast affected both (-) and (+)-strand RNA levels and also decreased the in vitro activity of the purified replicase preparations obtained from yeast. Moreover, the in vitro assembly of functional TBSV replicase in CFEs from yeast with depleted Pgk1 level was inefficient, resulting in reduced production of both dsRNA intermediate and new (+)RNA progeny. All these results point at deficiency in viral replicase assembly when Pgk1 is depleted. We have shown previously that the assembly of the tombusvirus replicase requires the co-opted cellular Hsp70 that uses ATP for its molecular chaperone function [32,36,37]. Based on the above observations, we propose that the ATP generated by the recruited Pgk1 serves the energy need of co-opted Hsp70 to drive efficient replicase complex assembly within the elaborate replication compartment (Fig 6G).

In addition, we obtained evidence that the functional activation of the TBSV p92 RdRp, which also requires Hsp70 molecular chaperone, depends on the ATP generated by the recruited Pgk1. The initially inactive p92 RdRp becomes functional during replicase assembly in the presence of p33 replication protein, the viral (+)RNA, ER membrane and the ATP-dependent Hsp70 [27,29]. In a simplified RdRp activation assay, we found that CFEs obtained from yeast with depleted Pgk1 were inefficient in promoting p92 RdRp activity. The addition of purified recombinant Hsp70 to the above assay could stimulate p92 RdRp activity to lesser extent in case of depleted Pgk1 than in the presence of wt yeast CFEs. Therefore, we propose that a major function of the Pgk1-generated ATP is to provide fuel to the co-opted Hsp70 during p92 RdRp activation and the assembly of viral replicase complexes (Fig 6G). The advantage of co-opting Pgk1 to the replicase complex could be that the energy hungry virus replicase assembly process does not need to intensively compete with cellular processes for access to plentiful ATP. Also, local production of ATP within the replication compartment could greatly facilitate the efficiency of Hsp70-driven replicase assembly by providing high ATP concentration within the replication compartment. There is previous evidence that...
Pgk1 could provide ATP to enhance the chaperone activity of Hsp90 that leads to multi-stress resistance [21].

In addition to Hsp70 molecular chaperone, TBSV also co-opts additional ATP-dependent host proteins into the viral replicase complex, including Vps4p AAA+ ATPase, involved in replicase assembly [7], and two DEAD-box helicases [38,39]. Using a reconstituted RNAi-based molecular probe, we found only a slightly less RNA protection in Pgk1-depleted yeast than in wt yeast (Fig 6F), which is in contrast with the poor TBSV RNA protection provided by VRCs assembled in the absence of ESCRT components [34]. Therefore, we propose that Vps4p AAA ATPase still functions efficiently enough in VRC assembly in yeast with depleted Pgk1 and ATP levels. On the other hand, the DEAD-box helicases, however, selectively affect (+)-strand synthesis, while Pgk1 affects both (-) and (+)-strand synthesis in CFE-based assay and in yeast to a similar extent. TBSV also affects the actin network, which requires ATP for functions, but it is currently on open question if the co-opted Pgk1 supplies ATP for the subverted actin network. Therefore, based on the collected data, we propose that the ATP-generated by Pgk1 within the replication compartment is primarily used by tombusviruses to provide ATP for the co-opted Hsp70 chaperone to support efficient assembly of the numerous viral replicase complexes.

Similar to our current findings, cancer cells also use glycolysis to efficiently generate ATP, even in the presence of oxygen [22]. Up-regulation of glycolytic machinery promotes rapidly proliferating cancer cells survival. Up-regulation of glycolytic pathway also takes place in effector T cells and it is required for immune cells activation [20,40]. Apparently, the relatively inefficient glycolytic pathway can provide enough ATP when up-regulated in these cells.

Several viruses are also known to reprogram the glycolytic pathway during infections based on metabolomic profiling [41,42,43], which unraveled enhanced glucose uptake into the infected cells. The hexokinase activity is increased in hepatitis C virus (HCV) or Dengue virus-infected cells [41,44]. ATP was shown to accumulate at the sites of HCV replication, likely to satisfy the energy demand of virus replication [45]. Whereas TBSV exploits glycolytic enzymes, such as PGK1 (this work) and Glyceraldehyde 3-phosphate dehydrogenase (GPDH, called Tdh2p and Tdh3p in yeast) for pro-viral functions [13,15], replication of a plant potexvirus is inhibited by GAPDH due to its RNA-binding function [46]. Altogether, metabolic reprogramming of the glycolytic pathway might be a widespread phenomenon during various viral infections.

In addition to its role in ATP production during glycolysis, Pgk1 seems to have additional roles in virus replication. For example, Sendai virus, a negative-strand RNA virus, co-opts Pgk1 to promote viral mRNA synthesis, albeit its enzymatic activity is not required for the stimulation of RNA synthesis [47]. A partial/recessive resistance in Arabidopsis against potyviruses is due to a single amino acid mutation in the conserved N-terminal portion of the chloroplast phosphoglycerate kinase (cPGK2), which is a nuclear gene encoding cPgk2 targeted to the chloroplast [48,49]. In addition, cPgk1 was found to bind to the 3'UTR of the viral RNA and it is involved in the localization of potexvirus RNA to the chloroplasts, which is important for virus accumulation [50,51].

**Materials and methods**

**Northern blot analysis of tombusvirus RNA accumulation**

TBSV RNA replication in yeasts and plants was analyzed after total RNA extraction with Northern blot analyses as described previously [52]. Briefly, BY4741 and GalS::PGK yeast strains were co-transformed with pHisGBK-CUP1- Hisp33/ADH-DI-72 and pGAD-CUP1-His92 [24]. The transformed yeast strains were grown at 23°C in SC-HL - (synthetic...
complete medium without histidine and leucine) media supplemented with 2% raffinose with or without 2% galactose and BCS for 24 h at 23°C. Then, yeast cultures were re-suspended in SC-HL− medium supplemented with 2% raffinose and with or without 2% galactose and 50 μM CuSO₄. Yeasts were grown at 23°C for 16 h before being collected for total RNA extraction.

**Protein analysis by Western blotting**

Yeast cells were grown as described above for Northern analysis. Total proteins were isolated by the NaOH method as described previously [53]. The total protein samples were analyzed by SDS-PAGE and Western blotting with anti-His and anti-PGK antibodies, followed by alkaline phosphatase-conjugated anti-mouse secondary antibody (Sigma) as described previously [54].

**Confocal microscopy in plants**

To examine the subcellular localization of Pgk1 in plants, *N. benthamiana* leaves were co-infiltrated with Agrobacterium carrying plasmids pGD-p92-YFP and pGD-BFP-PGK (OD₆₀₀ of 0.5, each) together with pGD-35S::p19, pGD-DI-72 and pGD-p33. After 48 h, the agroinfiltrated leaves were subjected to confocal microscopy (Olympus America FV1000) using 405 nm laser for BFP and 488 nm laser for YFP. Images were taken successively and merged using Olympus FLUOVIEW 1.5 software.

To identify interactions between NbPgk1 and TBSV p33 replication proteins, the plasmids pGD-p33-cYFP, pGD-nYFP-PGK and pGD-nYFP-MBP were transformed to Agrobacterium strain C58C1. These Agrobacterium transformants were used to co-agroinfiltrate the leaves of four-week-old *N. benthamiana* plants. Transformed leaves were subjected to confocal laser microscopy after 48 h using Olympus FV1000 microscope as described previously [13].

**Visualization and measurement of ATP levels in plants**

To detect the ATP levels within the tombusvirus replication compartments in plant cells, in the case of TBSV, PGK-silenced plants or control plants were agroinfiltrated with plasmids pGD-p33-ATEamYEMK, pGD-DI-72, pGD-35S::RFP-SKL, pGD-35S::p19 and pGD-p92. In the case of CIRV, the leaves were agroinfiltrated with pGD-p36-ATEamYEMK, pGD-35S::AtTim21-RFP, pGD-35S::p19, pGD-DI-72 and pGD-p95. The images were taken at 2.5 or 3.5 days post-agroinfiltration and analyzed with the method described previously [31]. Confocal FRET images were obtained with an Olympus FV1000 microscope (Olympus America). Cells were excited by a 405 nm laser diode, and CFP and Venus were detected at 480–500 nm and 515–615 nm wavelength ranges, respectively. Each YFP/CFP ratio was calculated by dividing pixel-by-pixel a Venus image with a CFP image using Olympus FLUOVIEW software or ImageJ software.

Additional standard experimental procedures are presented in the supporting information S1 Text.

**Supporting information**

*S1 Fig. Co-purification of Pgk1 with the p33 viral replication protein.* Top panel: Western blot analysis of co-purified His₆-tagged Pgk1 with FLAG-affinity purified FLAG-p33 from the membrane fraction of yeast. Pgk1 was detected with anti-His antibody. The negative control was FLAG-tagged GFP purified from the soluble (S) and membrane-fractions (M) of yeast extracts using a FLAG-affinity column. Second panel: Western blot of purified FLAG-p33 and FLAG-GFP detected with anti-FLAG antibody. Third panel: Western blot of His₆-tagged Pgk1
and His<sub>6</sub>-p33 (lane 1) proteins in the total yeast extracts using anti-His antibody. Fourth panel: Western blot analysis of FLAG-p33 and FLAG-GFP in the total yeast extracts with anti-FLAG antibody. Bottom panel: Coomassie Blue-stained SDS-PAGE with total protein extracts is shown as a loading control. Each experiment was repeated two times.

(S2 Fig) Reduced TBSV repRNA accumulation in yeast with depleted Pgk1 level at a late time point. Northern blot analysis shows decreased TBSV repRNA accumulation in a yeast strain (GALS::PGK1) when Pgk1 was depleted. To launch TBSV repRNA replication, we expressed His<sub>6</sub>-p33 and His<sub>6</sub>-p92<sup>pol</sup> from the copper-inducible CUP1 promoter, and DI-72(+) repRNA from the ADH1 promoter in the parental (BY4741) and in GALS::PGK1 yeast strains. Note that GALS::PGK1 yeast strain expresses HA-tagged Pgk1 from the galactose-inducible GAL8 promoter from chromosomal location (i.e., HA-Pgk1 replaces the wt Pgk1 in this haploid yeast strain). The yeast cells were cultured for 24 hours at 29°C in either 2% galactose +2% raffinose [(Raf+Gal), inducing condition for HA-Pgk1] or 2% raffinose (lack of induction of HA-Pgk1 expression) SC minimal media supplemented with 50 μM CuSO<sub>4</sub>. The accumulation level of DI-72(+) repRNA was normalized based on 18S rRNA levels (second panel, Northern blot).

(S3 Fig) Knock-down of Pgk1 level by VIGS reduces the symptoms caused by CNV. N. benthamiana plants were VIGS treated for 9 days, followed by inoculation with CNV. The pictures were taken 10 days after post infection. The control plants were VIGS treated with TRV-cGFP (the C-terminal half of the GFP sequence).

(S4 Fig) Reduced activity of the TBSV replicase assembled in vitro in CFEs prepared from a yeast strain (GALS::PGK1) with reduced level of Pgk1 expression. Purified recombinant MBP-p33 and MBP-p92<sup>pol</sup> replication proteins of TBSV and in vitro transcribed TBSV DI-72 (+) repRNA were added to the CFEs prepared from the shown yeast strains. Top panel: denaturing PAGE analysis of in vitro tombusvirus replicase activity in the CFEs. The <sup>32</sup>P-labeled TBSV repRNA products of the reconstituted replicases were detected by denaturing PAGE. Middle panel: The amount of Pgk1 was estimated by Western blotting using anti-Pgk1 antibody in yeast lysates. The sample from GALS::PGK1 yeast strain is shown on the right. Bottom panel: The CFEs contained the same amounts of total yeast proteins as demonstrated by Coomassie blue-staining of SDS-PAGE.

(S1 Text) Additional materials and methods used in this study.

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References

1. Fernandez de Castro I, Tenorio R, Risco C (2016) Virus assembly factories in a lipid world. Curr Opin Virol 18: 20–28. https://doi.org/10.1016/j.coviro.2016.02.009 PMID: 26985879

2. Wang A (2015) Dissecting the molecular network of virus-plant interactions: the complex roles of host factors. Annu Rev Phytopathol 53: 45–66. https://doi.org/10.1146/annurev-phoyo-080614-120001 PMID: 25938276

3. den Boon JA, Ahlquist P (2010) Organelle-like membrane compartmentalization of positive-strand RNA virus replication factories. Annu Rev Microbiol 64: 241–256. https://doi.org/10.1146/annurev.micro.112408.134012 PMID: 20825348

4. Paul D, Bartenschlager R (2015) Flaviviridae Replication Organelles: Oh, What a Tangled Web We Weave. Annu Rev Virol 2: 289–310. https://doi.org/10.1146/annurev-virology-100114-055007 PMID: 26958917

5. Nagy PD, Pogany J (2012) The dependence of viral RNA replication on co-opted host factors. Nature Reviews Microbiology 10: 137–149.

6. Rochon D, Singh B, Reade R, Theilmann J, Ghoshal K, et al. (2014) The p33 auxiliary replicase protein of Cucumber necrosis virus targets peroxisomes and infection induces de novo peroxisome formation from the endoplasmic reticulum. Virology 452–453: 133–142. https://doi.org/10.1016/j.virol.2013.12.035 PMID: 24606690

7. Barajas D, Martin IF, Pogany J, Risco C, Nagy PD (2014) Noncanonical Role for the Host Vps4 AAA+ ATPase ESCRT Protein in the Formation of Tomato Bushy Stunt Virus Replicase. PLoS Pathog 10: e1004087. https://doi.org/10.1371/journal.ppat.1004087 PMID: 24763736

8. Fernandez de Castro I, Fernandez JJ, Barajas D, Nagy PD, Risco C (2017) Three-dimensional imaging of the intracellular assembly of a functional viral RNA replicase complex. J Cell Sci 130: 260–268. https://doi.org/10.1242/jcs.181586 PMID: 27026525

9. Xu K, Nagy PD (2016) Enrichment of Phosphatidyethanolamine in Viral Replication Compartments via Co-opting the Endosomal Rab5 Small GTPase by a Positive-Strand RNA Virus. PLoS Biol 14: e2000128. https://doi.org/10.1371/journal.pbio.2000128 PMID: 27760328

10. Nawaz-Ul-Rehman MS, Prasanth KR, Xu K, Sassvari Z, Kovalev N, et al. (2016) Viral Replication Protein Inhibits Cellular Cofilin Actin Depolymerization Factor to Regulate the Actin Network and Promote Viral Replicase Assembly. PLoS Pathog 12: e1005440. https://doi.org/10.1371/journal.ppat.1005440 PMID: 26863541

11. Nagy PD (2008) Yeast as a model host to explore plant virus-host interactions. Annu Rev Phytopathol 46: 217–242. https://doi.org/10.1146/annurev.phoyo.121407.093958 PMID: 18422427

12. Serva S, Nagy PD (2006) Proteomics analysis of the tombusvirus replicase: Hsp70 molecular chaperone is associated with the replicase and enhances viral RNA replication. J Virol 80: 2162–2169. https://doi.org/10.1128/JVI.80.5.2162-2169.2006 PMID: 16474124

13. Wang RY, Nagy PD (2008) Tomato bushy stunt virus co-opts the RNA-binding function of a host metabolic enzyme for viral genomic RNA synthesis. Cell Host Microbe 3: 178–187. https://doi.org/10.1016/j.chom.2008.02.005 PMID: 18329617
14. Sasvari Z, Izotova L, Kinzy TG, Nagy PD (2011) Synergistic Roles of Eukaryotic Translation Elongation Factors 1Bgamma and 1A in Stimulation of Tombusvirus Minus-Strand Synthesis. PLoS Pathog 7: e1002438. https://doi.org/10.1371/journal.ppat.1002438 PMID: 22194867

15. Huang TS, Nagy PD (2011) Direct inhibition of tombusvirus plus-strand RNA synthesis by a dominant negative mutant of a host metabolic enzyme, glyceraldehyde-3-phosphate dehydrogenase, in yeast and plants. J Virol 85: 9090–9102. https://doi.org/10.1128/JVI.00666-11 PMID: 21697488

16. Li Z, Pogany J, Tumpan S, Esposito AM, Kinzy TG, et al. (2010) Translation elongation factor 1A facilitates the assembly of the tombusvirus replicase and stimulates minus-strand synthesis. PLoS Pathog 6: e1001175. https://doi.org/10.1371/journal.ppat.1001175 PMID: 21079685

17. Barajas D, Jiang Y, Nagy PD (2009) A unique role for the host ESCRT proteins in replication of Tomato bushy stunt virus. PLoS Pathog 5: e1000705. https://doi.org/10.1371/journal.ppat.1000705 PMID: 20041173

18. Barajas D, Xu K, de Castro Martin IF, Sasvari Z, Brandizzi F, et al. (2014) Co-opted Oxysterol-Binding ORP and VAP Proteins Channel Sterols to RNA Virus Replication Sites via Membrane Contact Sites. PLoS Pathog 10: e1004388. https://doi.org/10.1371/journal.ppat.1004388 PMID: 25329172

19. Wang S, Jiang B, Zhang T, Liu L, Wang Y, et al. (2015) Insulin and mTOR Pathway Regulate HDAC3-Mediated Deacetylation and Activation of PGK1. PLoS Biol 13: e1002243. https://doi.org/10.1371/journal.pbio.1002243 PMID: 26356530

20. Finlay DK (2015) Metabolic regulation of natural killer cells. Biochem Soc Trans 43: 758–762. https://doi.org/10.1042/BST20150116 PMID: 26551725

21. Chen X, Zhao C, Li X, Wang T, Li Y, et al. (2015) Terazosin activates Pgk1 and Hsp90 to promote stress resistance. Nat Chem Biol 11: 19–25. https://doi.org/10.1038/nchembio.1657 PMID: 25383758

22. Stine ZE, Dang CV (2013) Stress eating and tuning out: cancer cells re-wire metabolism to counter resistance. Crit Rev Biochem Mol Biol 48: 609–619. https://doi.org/10.3109/10409238.2013.844093 PMID: 24099138

23. Krishnan P, Gullen EA, Lam W, Dutschman GE, Grill SP, et al. (2003) Novel role of 3-phosphoglycerate kinase, a glycolytic enzyme, in the activation of L-nucleoside analogs, a new class of anticancer and antiviral agents. J Biol Chem 278: 36726–36732. https://doi.org/10.1074/jbc.M307052200 PMID: 12869554

24. Shah Nawaz-Ul-Rehman M, Reddisiva Prasanth K, Baker J, Nagy PD (2013) Yeast screens for host factors in positive-strand RNA virus replication based on a library of temperature-sensitive mutants. Methods 59: 207–216. https://doi.org/10.1016/j.ymeth.2012.11.001 PMID: 23147170

25. Jiang Y, Serviene E, Gal J, Panavas T, Nagy PD (2006) Identification of essential host factors affecting tombusvirus RNA replication based on the yeast Tet promoters Hughes Collection. J Virol 80: 7394–7404. https://doi.org/10.1128/JVI.02686-05 PMID: 16840320

26. Panavas T, Serviene E, Brasher J, Nagy PD (2005) Yeast genome-wide screen reveals dissimilar sets of host genes affecting replication of RNA viruses. Proc Natl Acad Sci U S A 102: 7326–7331. https://doi.org/10.1073/pnas.0502604102 PMID: 15883361

27. Pogany J, Nagy PD (2015) Activation of Tomato Bushy Stunt Virus RNA-Dependent RNA Polymerase by Cellular Heat Shock Protein 70 Is Enhanced by Phospholipids In Vitro. J Virol 89: 5714–5723. https://doi.org/10.1128/JVI.03711-14 PMID: 25762742

28. Mendu V, Chiu M, Barajas D, Li Z, Nagy PD (2010) Cpr1 cyclophilin and Ess1 parvulin prolyl isomerases interact with the tombusvirus replication protein and inhibit viral replication in yeast model host. Virology 406: 342–351. https://doi.org/10.1016/j.virol.2010.07.022 PMID: 20709345

29. Pogany J, Nagy PD (2012) p33-Independent Activation of a Truncated p92 RNA-Dependent RNA Polymerase of Tomato Bushy Stunt Virus in Yeast Cell-Free Extract. J Virol 86: 12025–12038. https://doi.org/10.1128/JVI.01303-12 PMID: 22933278

30. Janke C, Magiera MM, Rathfelder N, Taxis C, Reber S, et al. (2004) A versatile toolbox for PCR-based tagging of yeast genes: new fluorescent proteins, more markers and promoter substitution cassettes. Yeast 21: 947–962. https://doi.org/10.1002/yea.1142 PMID: 15334558

31. Imamura H, Nhat KP, Togawa H, Saito K, Iino R, et al. (2009) Visualization of ATP levels inside single living cells with fluorescence resonance energy transfer-based genetically encoded indicators. Proc Natl Acad Sci U S A 106: 15651–15656. https://doi.org/10.1073/pnas.0904764106 PMID: 19720993

32. Pogany J, Stork J, Li Z, Nagy PD (2008) In vitro assembly of the Tomato bushy stunt virus replicase requires the host Heat shock protein 70. Proc Natl Acad Sci U S A 105: 19956–19961. https://doi.org/10.1073/pnas.0810851105 PMID: 19060219

33. Panaviene Z, Panavas T, Serva S, Nagy PD (2004) Purification of the cucumber necrosis virus replicase from yeast cells: role of coexpressed viral RNA in stimulation of replicase activity. J Virol 78: 8254–8263. https://doi.org/10.1128/JVI.78.15.8254-8263.2004 PMID: 15254197
34. Kovalev N, Inaba JI, Li Z, Nagy PD (2017) The role of co-opted ESCRT proteins and lipid factors in protection of tombusviral double-stranded RNA replication intermediate against reconstituted RNAi in yeast. PLoS Pathog 13: e1006520. https://doi.org/10.1371/journal.ppat.1006520 PMID: 28759634

35. Drinnenberg IA, Weinberg DE, Xie KT, Mower JP, Wolfe KH, et al. (2009) RNAi in budding yeast. Science 326: 544–550. https://doi.org/10.1126/science.1176945 PMID: 19745116

36. Wang RY, Stork J, Pogany J, Nagy PD (2009) A temperature sensitive mutant of heat shock protein 70 reveals an essential role during the early steps of tombusvirus replication. Virology 394: 28–38. https://doi.org/10.1016/j.virol.2009.08.003 PMID: 19748649

37. Wang RY, Stork J, Nagy PD (2009) A key role for heat shock protein 70 in the localization and insertion of tombusvirus replication proteins to intracellular membranes. J Virol 83: 3276–3287. https://doi.org/10.1128/JVI.02313-08 PMID: 19153242

38. Kovalev N, Nagy PD (2014) The Expanding Functions of Cellular Helicases: The Tombusvirus RNA Replication Enhancer Co-opts the Plant eIF4AIII-Like AtRH2 and the DDX5-Like AtRH5 DEAD-Box RNA Helicases to Promote Viral Asymmetric RNA Replication. PLoS Pathog 10: e1004051. https://doi.org/10.1371/journal.ppat.1004051 PMID: 24743583

39. Kovalev N, Pogany J, Nagy PD (2012) A Co-Opted DEAD-Box RNA Helicase Enhances Tombusvirus Plus-Strand Synthesis. PLoS Pathog 8: e1002537. https://doi.org/10.1371/journal.ppat.1002537 PMID: 22359508

40. Peng M, Yin N, Chhangawala S, Xu K, Leslie CS, et al. (2016) Aerobic glycolysis promotes T helper 1 cell differentiation through an epigenetic mechanism. Science 354: 481–484. https://doi.org/10.1126/science.aaf6284 PMID: 27708054

41. Fontaine KA, Sanchez EL, Camardra R, Lagunoff M (2015) Dengue virus induces and requires glycolysis for optimal replication. J Virol 89: 2358–2366. https://doi.org/10.1128/JVI.02309-14 PMID: 25505078

42. Su MA, Huang YT, Chen IT, Lee DY, Hsieh YC, et al. (2014) An invertebrate Warburg effect: a shrimp glycolytic enzyme facilitates the assembly of viral replicase complexe

43. Prasanth KR, Huang YW, Liou MR, Wang RY, Hu CC, et al. (2011) Glyceraldehyde-3-phosphate dehydrogenase negatively regulates the replication of Bamboo mosaic virus and its associated satellite RNA. J Virol 85: 8829–8840. https://doi.org/10.1128/JVI.00556-11 PMID: 22175476

44. Ogino T, Iwama M, Kinouchi J, Shibagaki Y, Tsukamoto T, et al. (1999) Involvement of a cellular glycolytic enzyme, phosphoglycerate kinase, in Sendai virus transcription. J Biol Chem 274: 35999–36008. PMID: 10585490

45. Lin JW, Ding MP, Hsu YH, Tsai CH (2007) Chloroplast phosphoglycerate kinase, a gluconeogenic enzyme, is required for efficient accumulation of Bamboo mosaic virus. Nucleic Acids Res 35: 424–432. https://doi.org/10.1093/nar/gkl061 PMID: 17169994

46. Poque S, Pagny G, Ouibrahim L, Chague A, Eyquard JP, et al. (2015) Allelic variation at the rpv1 locus controls partial resistance to Plum pox virus infection in Arabidopsis thaliana. BMC Plant Biol 15: 159. https://doi.org/10.1186/s12870-015-0559-5 PMID: 26109391

47. Ouibrahim L, Mazier M, Estevan J, Pagny G, Decroocq V, et al. (2014) Cloning of the Arabidopsis rwm1 gene for resistance to Watermelon mosaic virus points to a new function for natural virus resistance genes. Plant J 79: 705–716. https://doi.org/10.1111/tpj.12586 PMID: 24930633

48. Lin JW, Ding MP, Hsu YH, Tsai CH (2007) Chloroplast phosphoglycerate kinase, a gluconeogenic enzyme, is required for efficient accumulation of Bamboo mosaic virus. Nucleic Acids Res 35: 424–432. https://doi.org/10.1093/nar/gkl061 PMID: 17169994

49. Cheng SF, Huang YP, Chen LH, Hsu YH, Tsai CH (2013) Chloroplast phosphoglycerate kinase is involved in the targeting of Bamboo mosaic virus to chloroplasts in Nicotiana benthamiana plants. Plant Physiol 163: 1598–1608. https://doi.org/10.1104/pp.113.229666 PMID: 24154620

50. Cheng CP, Jiao HM, Jonczyk M, Serviene E, Nagy PD (2007) Expression of the Arabidopsis Xnr4p 5'-3' exoribonuclease facilitates degradation of tombusvirus RNA and promotes rapid emergence of viral variants in plants. Virology 368: 238–248. https://doi.org/10.1016/j.virol.2007.07.001 PMID: 17688902

51. Panavas T, Nagy PD (2003) Yeast as a model host to study replication and recombination of defective interfering RNA of Tomato bushy stunt virus. Virology 314: 315–325. PMID: 14517084
54. Li Z, Barajas D, Panavas T, Herbst DA, Nagy PD (2008) Cdc34p ubiquitin-conjugating enzyme is a component of the tombusvirus replicase complex and ubiquitinites p33 replication protein. J Virol 82: 6911–6926. https://doi.org/10.1128/JVI.00702-08 PMID: 18463149