RESEARCH ARTICLE

Rapamycin-insensitive mTORC1 activity controls eIF4E:4E-BP1 binding [version 1; peer review: 3 approved]

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Abstract
The recent development of mammalian target of rapamycin (mTOR) kinase domain inhibitors and genetic dissection of rapamycin-sensitive and -insensitive mTOR protein complexes (mTORC1 and mTORC2) have revealed that phosphorylation of the mTOR substrate 4E-BP1 on amino acids Thr37 and/or Thr46 represents a rapamycin-insensitive activity of mTORC1. Despite numerous previous reports utilizing serine (Ser)-to-alanine (Ala) and threonine (Thr)-to-Ala phosphorylation site mutants of 4E-BP1 to assess which post-translational modification(s) directly regulate binding to eIF4E, an ambiguous understanding persists. This manuscript demonstrates that the initial, rapamycin-insensitive phosphorylation event at Thr46 is sufficient to prevent eIF4E:4E-BP1 binding. This finding is relevant, particularly as mTOR kinase domain inhibitors continue to be assessed for clinical efficacy, since it clarifies a difference between the action of these second-generation mTOR inhibitors and those of rapamycin analogues.
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Introduction

The mammalian target of rapamycin (mTOR) protein is an atypical Ser/Thr protein kinase named for its well-characterized inhibition by the natural product rapamycin. Rapamycin-sensitive orthologues of mTOR exist in eukaryotes from yeast to man and are required for growth and proliferation of perhaps all eukaryotic cells. As such, rapamycin has been classified as an anti-fungal agent and is clinically approved as an immunosuppressant and cancer therapy.[1-3]. Well-characterized in vivo substrates of rapamycin-sensitive mTOR activity include the 70 kDa ribosomal protein S6 kinase (S6K1) and the eukaryotic [translation] initiation factor 4E (eIF4E)-binding protein 1 (4E-BP1). The mTOR-dependent phosphorylation site on S6K1, Thr389, is required for kinase activity, explaining rapamycin’s inhibition of S6K1 activity[4]. 4E-BP1, on the other hand, is subject to multisite phosphorylation culminating in the release of bound eIF4E, leading to an ambiguous understanding of which phosphorylation site(s) regulate(s) eIF4E binding[5].

Mammalian 4E-BP1 is subject to an ordered phosphorylation on at least 5 major amino acid residues in response to serum-stimulation, as has been demonstrated by two-dimensional (isoelectric focusing and SDS-PAGE) electrophoresis (2DE). This approach of separating post-translationally modified forms of a protein based on charge and apparent molecular weight has proved to be particularly useful when combined with phosphorylation-specific antibodies. Due to amino acid sequence similarity, phospho-specific anti-Thr37/46 antibodies did not allow determination of whether the initial phosphorylation event is at Thr37 or Thr46 using this technique, although priming phosphorylation at both of these sites is thought to be required for subsequent phosphorylation at Thr70, followed by phosphorylation at an unidentified site, and finally at Ser65[6,7]. Given their positions flanking the amino acids responsible for eIF4E binding (amino acid residues 53–59), it is conceivable that Thr46 and Ser65 are responsible for the phosphorylation-mediated modulation of eIF4E binding occurring in response to mTOR activity. Indeed, a significant body of work supports the role of phosphorylation at Thr46 in regulating eIF4E:4E-BP1 binding[8-12]. Detailed analyses have led, however, to conflicting results regarding the importance of Ser65 phosphorylation in preventing this protein:protein interaction[6,11,13-15].

While rapamycin is effective in blocking phosphorylation at Thr70 and Ser65, phospho-specific antibodies to Thr37/46 show that at least one of these sites is largely rapamycin-insensitive[16,17]. This residual rapamycin-insensitive phosphorylation is sensitive to serum starvation, amino acid withdrawal, and non-specific phosphatidylinositol 3-kinase (PI3K) and PI3K-like kinase (PIKK) inhibitors[16-19]. Furthermore, the use of mTOR kinase domain inhibitors (Torin1 and PP242) in combination with mTOR complex 2 (mTORC2)-deficient cells, has allowed the determination that Thr37/46 phosphorylation represents a rapamycin-insensitive function of mTOR complex 1 (mTORC1)[20,21].

In vivo studies addressing the relative importance of 4E-BP1 phosphorylation sites have been hampered by its ordered phosphorylation, wherein Thr-to-Ala mutation of Thr37 or Thr46 will block subsequent phosphorylation at Thr70 and Ser65. Mounting circumstantial evidence supports the notion that phosphorylation of Thr37/Thr46 alone is the key event regulating eIF4E:4E-BP1 binding in vivo. Intracellular co-localization of endogenous 4E-BP1 and eIF4E best correlates with dephosphorylation at Thr37/46[22], 7-methyl-GTP (cap-column) pull down of eIF4E:4E-BP1 complexes is enhanced by mTOR kinase domain inhibitors more than it is by rapamycin[11]. Most importantly, however, mTOR active site inhibitors capable of blocking phosphorylation at Thr37/46 (and not rapamycin) induce 4E-BP-dependent phenotypes in cells[23].

This manuscript describes new data demonstrating that 4E-BP1 phosphorylation at the initial, mTORC1-dependent, rapamycin-insensitive phosphorylation site is alone in regulating eIF4E binding. Furthermore, this work suggests that Thr46, and not Thr37, is the key phosphorylation site. Given the recent push for pharmaceutical development of kinase inhibitors that block both the rapamycin-sensitive and rapamycin-insensitive activities of mTOR[24], a thorough understanding of the importance of rapamycin-insensitive mTORC1 activity is crucial. This manuscript supports the idea that clinically used mTOR kinase domain inhibitors will reduce eIF4E availability much more profoundly than have clinically approved rapamycin analogs.

Materials and methods

Isoelectric focusing combined with SDS-PAGE based two-dimensional electrophoresis was performed as previously described[25]. Far western analyses were performed as follows using a 32P-labelled eIF4E protein with an N-terminal substrate peptide for heart muscle kinase (HMK). One hundred units of bovine HMK was suspended in 10 µl of 40 mM DTT and allowed to stand for 10 minutes. Five micrograms of HMK-eIF4E protein was mixed with 3 µl 10X HMK Buffer (200 mM Tris, pH 7.5, 10 mM DTT, 1 M NaCl, 120 mM MgCl2), 5 µl [γ-32P] ATP 3000 Ci/mmol, 1 µl HMK (10U), and water (to 30 µl) and incubated for 45 minutes at 4°C. Probe purification was performed using Pharmacia Nick Column Sephadex G-50 DNA grade. Membranes were subjected to pre-hybridization (25 mM HEPES-KOH, pH 7.7, 25 mM NaCl, 5 mM MgCl2, 1 mM DTT, 0.1% NP40, 5% skim milk) for 5 hours at 4°C. Probe hybridization was performed in buffer (20 mM HEPES-KOH, pH 7.7; 75 mM KCl, 2.5 mM MgCl2, 0.1 mM EDTA, 1 mM DTT, 0.1% NP40, 1% skim milk) with 250,000 cpm/ml of radiolabelled probe for 10 minutes at 4°C. Membranes were washed with hybridization buffer 3 times, 15 minutes prior to exposure to film (BIOMAX MS, Kodak). Following far western analysis, membranes were probed sequentially with antibodies for Phospho-Ser65, Thr70, Thr37/46, and Total 4E-BP1 (Cell Signaling Technology). Cap column based fractionation of cell lysates was performed as previously described[26]. HeLa S3 and 293 HEK cells were treated with PP242 (2.5 µM, 30 min) or Rapamycin (10 nM, 30 min) unless otherwise indicated. Stable HeLa S3 cell lines expressing wild-type and mutant HA-4E-BP1 proteins were generated using previously described mammalian expression constructs[27] and G418 selection. Control siRNA and siRNA for mTOR was from Cell Signaling Technology and delivered using Lipofectamine2000 according to manufacturer’s instructions (Invitrogen). All antibodies were from Cell Signaling Technology and were used according to manufacturer’s instructions. Compounds used were rapamycin (LC Labs), PP242 (Intellikine), torin1 (Tocris), PI-103 (EMD), etoposide and nocodazole (Sigma). Standard laboratory practices were used to control bias and unwanted sources of variability in this study. The primary limitation of the
datasets presented in this manuscript is that they represent single biological replicates of an experimental procedure.

Results

4E-BP1 Thr37/46 Phosphorylation is sufficient to block elf4E binding

Far western blot analysis, using radiolabeled HMK-eIF4E as a probe, is an effective measure of elf4E-binding activity\(^1\). Using this approach, we demonstrate that the treatment of cells with the mTOR kinase inhibitor PP242 results in the following: (i) dramatic increases in the elf4E-binding competent pool of 4E-BP1, (ii) reduction of 4E-BP1Thr37/46 and Ser65 phosphorylation, and (iii) unaffected binding to bands corresponding in molecular weight to 4E-T and elf4G (Figure 1A). The mTOR-dependent modulation of elf4E-binding activity is also apparent under physiological conditions (serum starvation vs. serum stimulation) and can be blocked by siRNA knockdown of mTOR in HeLa S3 cells (Figure 1B). Notably, western blot-based detection of non-phosphorylated (Thr46; denoted NP-Thr46) 4E-BP1 using a rabbit monoclonal antibody (clone: 87D12) recapitulates HMK-eIF4E binding to 4E-BP1.

To more precisely elucidate the molecular modifications of 4E-BP1 induced by PP242 that induce elf4E binding, whole cell lysates were prepared from HEK293 cells that were subjected to short (30 min) treatment with PP242 or rapamycin followed by 2DE (isoelectric focusing and SDS-PAGE) and western blot analyses (Figure 1C). Under control (DMSO-treated) conditions, the previously described hierarchical, multi-site phosphorylation of 4E-BP1 is observed. Here, six differentially phosphorylated forms (labeled A-F) are detected by the total 4E-BP1 antibody, with forms B-F phosphorylated at Thr37 and/or Thr46, D-F phosphorylated at Thr70, and F phosphorylated at Ser65. Far western blot analysis demonstrates that only spot A is competent to bind HMK-eIF4E under control conditions suggesting that the modification responsible for spot B (Thr37 or Thr46) disrupts this interaction. Upon inhibition of mTOR with rapamycin or PP242, the predicted decrease in hyper-phosphorylated 4E-BP1 forms is observed with an increase in hypo-phosphorylated 4E-BP1. While the identity of the phosphorylation event responsible for spot E remains undetermined, these data show that this phosphorylation site is resistant to mTOR inhibition, as rapamycin and PP242-resistant phospho-forms emanating from spot E appear above spots B-D. It is of note that the PP242-induced spot above spot B, which is not phosphorylated at Thr37 or Thr46, represents mono-phosphorylated 4E-BP1 (at the site responsible for spot E) and retains elf4E-binding ability. The most likely candidates for this site are Thr84, which has shown to be responsible for spot E) and retains eIF4E-binding ability. The most likely candidates for this site are Thr84, which has shown to be responsible for a similarly slow SDS-PAGE migration\(^1\), and Ser101, which has been shown to promote 4E-BP1/Raptor binding\(^2\). The notion that Ser101 is responsible for spot E is particularly appealing, as this would provide a sound explanation for the hierarchical ordering of this phosphorylation event prior to Ser65 (spot F). That is, it is reasonable to believe that strong 4E-BP1/Raptor binding is required for complete 4E-BP1 phosphorylation, including at Ser65.

4E-BP1 Thr46 is phosphorylated prior to Thr37 under normal growth conditions and prevents association with cap-bound elf4E

Next, to determine whether Thr37 or Thr46 phosphorylation accounts for “spot B”, which is impaired for elf4E binding, HeLa cell lines stably expressing wild-type or mutant HA-4E-BP1 proteins were generated. Given the ordered phosphorylation of 4E-BP1, mutation of the primary phosphorylation site should block mutation of that of the subsequent phosphorylation site. For this reason, we analyzed Thr37Ala and Thr46Ala mutants in vivo using the phospho-4E-BP1 Thr37/46 antibody to determine whether the preclusion of phosphorylation at one site blocks phosphorylation at the other (Figure 2A). While endogenous 4E-BP1 was detected by the phospho-4E-BP1 Thr37/46 antibody in lysates from all stably selected cell lines, exogenous HA-tagged 4E-BP1 was poorly detected in the Thr46Ala mutant sample, suggesting that Thr46 phosphorylation is required for subsequent Thr37 phosphorylation under normal growth conditions. These results indicate that Thr46 is the initial phosphorylation site responsible for the shift from
spot A to spot B, thus phosphorylation at this site alone may be sufficient to prevent 4E-BP1 binding to eIF4E (in Figure 1C). This conclusion, that Thr46 phosphorylation precedes Thr37 phosphorylation, has previously been reached by another group 29. This model, wherein 4E-BP1 phosphorylation at the initial phosphorylation site (Thr46) is sufficient to prevent eIF4E:4E-BP1 binding, is also supported by 7-methyl-GTP (cap-column) pull down data (Figure 2B). Here the cap-column serves as a molecular mimic for the mRNA 5’-cap allowing eIF4E and associated binding proteins to be isolated from cell lysates. As a chemically induced pseudo-mitotic state has previously been shown to dramatically modulate the phosphorylation of 4E-BP1 26,30, nocodazole treatment was employed to potentially increase the diversity of 4E-BP1 phospho-forms present within our lysates. Control and nocodazole-blocked HeLa S3 cells were subjected to cap-column pull-down of eIF4E and associated 4E-BP1. This technique allowed detectable binding of only the fastest SDS-PAGE migrating forms of 4E-BP1, indicating differential binding between hypophosphorylated and hyper-phosphorylated 4E-BP1 had occurred. The use of phospho-specific antibodies demonstrates that Thr37/Thr46 phosphorylated 4E-BP1 is not detectably present in this cap-column bound eIF4E fraction. Trace amounts of Thr70 phosphorylated 4E-BP1 are detected in these lanes, suggesting that mono-phosphorylated (at Thr70) 4E-BP1 exists and is eIF4E binding competent. A similar conclusion was recently reached by another group 31. It should be noted that the total 4E-BP1 antibody detects a doublet band in the cap-column bound eIF4E fraction. The upper band of this doublet likely represents 4E-BP1 phosphorylated at the site responsible for the above-described “spot E”; a phospho-form which retains eIF4E-binding ability.

**Existence of alternative 4E-BP1 phosphorylation patterns**

To further explore the potential existence of 4E-BP1 phospho-forms failing to adhere to the strict hierarchical phosphorylation pattern Thr37/46->Thr70->Thr37/46Ala double point mutant proteins, the Thr37Ala protein is still recognized suggesting that Thr37 is not required for Thr46 phosphorylation. Endogenous 4E-BP1 phosphorylated at Thr37/46 is shown as a control. B) Untreated and Nocodazole-blocked HeLa S3 cells were subjected to cap-column pull down of eIF4E and associated proteins. eIF4E was eluted with m7-GDP followed by SDS, and fractions were analyzed by western blot using eIF4E and 4E-BP1 antibodies and phospho-4E-BP1 antibodies.

**Figure 2.** Thr46 phosphorylation is required for Thr37 phosphorylation and is sufficient to prevent eIF4E:4E-BP1 binding. A) HeLa S3 cells stably expressing 4E-BP1 mutant proteins were subjected to western blotting using anti-HA antibody (upper), and phospho-4E-BP1 (Thr37/46) antibody (middle and lower panels). While phospho-4E-BP1 (Thr37/46) antibodies fail to detect the Thr46Ala single and Thr46/46Ala double point mutant proteins, the Thr37Ala protein is still recognized suggesting that Thr37 is not required for Thr46 phosphorylation. Endogenous 4E-BP1 phosphorylated at Thr37/46 is shown as a control. B) Untreated and Nocodazole-blocked HeLa S3 cells were subjected to cap-column pull down of eIF4E and associated proteins. eIF4E was eluted with m7-GDP followed by SDS, and fractions were analyzed by western blot using eIF4E and 4E-BP1 antibodies and phospho-4E-BP1 antibodies.

HeLa S3 Control and Nocodazole-Blocked Mixture

**Figure 3.** 4E-BP1 phosphorylation is not strictly ordered. Untreated and nocodazole-blocked HeLa S3 cells lysed, pooled and subjected to 2D-E prior to analysis with phospho-specific and total 4E-BP1 antibodies (left panels). In addition to the standard ordered phosphorylation (Thr37 or Thr46, then “spot B”, then Thr70, then “spot E”, then Ser65, then “spot G”), 4E-BP1 singly phosphorylated at Thr70 is also observed indicating that this species can exist in vivo. To facilitate interpretation, the same images have been overlaid with a grid of circles corresponding to spots visible with the total 4E-BP1 antibody (right panels).
emerged. Here, in addition to the normal hierarchical phosphorylation described above, alternative mono-phosphorylated species were present, as was an additional phosphorylation site causing the appearance of “spot G”. Although, this chemical-induced atypical phosphorylation pattern may not exist under physiological conditions, these data provide support for the existence of these alternative 4E-BP1 phospho-forms.

**PP242 inhibits etoposide-induced phosphorylation of Ser/Thr-Gln motifs**

Recently it has been reported that among mTOR kinase domain inhibitors, PP242 exhibits remarkably low specificity compared with Torin1, KU63794 and WYE35410. Notably, both PP242 and Torin1 were shown to bind multiple PI3K-related kinases (PIKKs), including three key DNA damage-activated kinases ATM, ATR and DNAPK, although cell-based assays failed to show inhibition of these kinases. To assess the specificity of PP242 in our system, we employed a broadly reactive phospho-specific antibody capable of recognizing multiple phospho-Ser/Thr-Gln substrates of ATM, ATR and DNAPK10,11. Indeed, with this antibody, we observed multiple etoposide-induced bands by western blot analysis (Figure 4), including bands corresponding to VCP phosphorylated at Ser784, a hallmark of the mTOR/PI3K inhibitors tested, only PP242 visibly reduced the etoposide-induced phosphorylation of this PIKK substrates (Figure 4A). As the concentration of PP242 (2.5 µM) was higher than that used for the other inhibitors (wortmannin 0.1 µM, PI-103 1.0 µM, Torin1 0.25 µM and Rapamycin 0.01 µM), PP242 pretreatment was also assessed at 0.34, 0.68 and 1.25 µM, and gave similar results (Figure 4B).

**Discussion**

Although, our conclusion that phosphorylation at Thr46 is the key event regulating 4E-BP1:elf4E binding has been suggested previously, none of these previous studies unambiguously established that Thr46 alone is the key important site. These studies either: (i) employed in vivo phosphorylation of a Thr46Ala point mutant also block subsequent phosphorylation events12,13; (ii) failed to identify a single important phosphorylation site10,11; or (iii) employed in vitro phosphorylation of Thr46Ala point mutant using a non-physiological kinase10,12.

Importantly, equally credible work has reported that phosphorylation on Thr46 is unimportant in the regulation of 4E-BP1:elf4E binding14,15. It is interesting to note that both of these studies evaluate the capacity of in vitro phosphorylation at Thr46 to disrupt pre-existing 4E-BP1:elf4E complexes and utilize cap column purification to isolate elf4E-bound 4E-BP1. This experimental detail is particularly relevant, as it has been shown that the RNA cap can exert an allosteric effect stabilizing 4E-BP1:elf4E binding16-18. Taken together, our data and these previous reports could suggest that Thr46 phosphorylation is sufficient to block the initial binding between elf4E and 4E-BP1 as observed by far western analyses, but that hyperphosphorylation of 4E-BP1, including at Ser65, is required to disrupt existing 4E-BP1:elf4E complexes.

The far western analysis of 2DE separated 4E-BP1 phospho-forms presented in Figure 1C is apparently at the limit of its useful range of detection. While this technique allows a comparison of binding efficiencies of spots A and B to evaluate the impact of Thr46 phosphorylation, it is not useful to compare the relative binding abilities of spots E and F to allow a similar assessment of the importance of Ser65 phosphorylation. Perhaps with a more sensitive assay we would have observed a similar decrease in elf4E binding upon Ser65 phosphorylation. Perhaps with a more sensitive assay we would have observed a similar decrease in elf4E binding upon Ser65 phosphorylation. This would be evidence that Thr46 phosphorylation is sufficient to block the initial binding of 4E-BP1 to elf4E when both proteins are present at low physiological concentrations, but that phosphorylation at multiple sites culminating at Ser65 is required to prevent/disrupt binding when the two proteins are present at high concentrations/local concentrations.

The data presented in Figure 2B at first seem to be at odds with this theory that 4E-BP1 phosphorylated at Thr37 or Thr46 could be pre-associated with mRNA 5’ cap-bound elf4E in cells. This technique assesses, however, the de novo association of elf4E (4E-BP1 associated or not) with an mRNA 5’ cap analog, therefore any pre-existing complexes must dissociate from cellular mRNA caps prior to isolation. These data do tell us that if there is some pool of 4E-BP1 phosphorylated on Thr46 associated with elf4E, this complex is not able to efficiently bind the mRNA cap analog. This assay may also evaluate the sequential binding of elf4E to the cap followed by 4E-BP1 binding to cap-associated elf4E and indicate in agreement with the far western data in Figure 1C that de novo binding is blocked by Thr46 phosphorylation.

Figure 2B and Figure 3 demonstrate that blocking cells in a pseudo-mitotic state with nocodazole results in phosphorylation of 4E-BP1 at up to 6 distinct phosphorylation sites, a finding at odds with previous work demonstrating hypophosphorylation of 4E-BP1 in
mitosis\textsuperscript{38}. The polyclonal antibody, referred to as 11208 and used in this previous study, however, exhibits selectivity for the non-phosphorylated forms of the protein after nocodazole treatment (ML, unpublished). A similar faulty antibody-based discrepancy regarding the phosphorylation state of 4E-BP1 in mitotic oocytes was resolved\textsuperscript{11}. While we observed at most 6 different phospho-forms of 4E-BP1, the PhosphoSitePlus database\textsuperscript{18} indicates that 20 distinct phosphorylation sites have been described in the literature and/or mass spectrometry-based datasets. This discrepancy suggests that many of these 20 phosphorylation events are either mutually exclusive, or occur only rarely, not at all, in response to specific stimuli, or in specific cell types.

Finally, the demonstration presented within this manuscript that PP242 exhibits poor specificity for mTOR in a cell-based assay for PIKK off-target effects likely does not change the conclusions of this manuscript; however, it serves as a reminder that any new mTOR inhibitor may have unanticipated effects. While the inhibition of mTORC2 by mTOR kinase domain inhibitors is expected to have profound effects, the evidence presented in this manuscript suggests that the rapamycin-insensitive activity of mTORC1 towards 4E-BP1 will also be quite important as the clinical safety and efficacy of mTOR kinase domain inhibitors is assessed.

Author contributions ML conceived the study, designed the experiments, and prepared the first draft of the manuscript. ML and MB carried out the research, were involved in the revision of the draft manuscript, and have approved the final content.

Competing interests No relevant competing interests declared.

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References


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This submission is quite nice for experts, but not particularly well-written for those outside of the field of 4E-BP1 phosphorylation, who will have difficulty reading it; hence my suggested clarifications below. The article is important as it demonstrates that the use of mTOR-kinase domain inhibitors would be more efficacious than the use of rapamycin analogs as therapeutics.

The work clarifies that phosphorylation of 4E-BP1 at Ser65 by mTOR is not required to inhibit 4E-BP1 binding to eIF4E and that rapamycin-insensitive mTOR-mediated phosphorylation at 4E-BP1 Thr46 is sufficient to inhibit 4E-BP1 binding to eIF4E. Phosphorylation at 4E-BP1 Thr37 is also shown to be rapamycin-insensitive. Notably, evidence that phosphorylation at Thr 37 or Thr 46 appears to be required for phosphorylation at Thr 70 and Ser 65 supported the notion that the earlier phosphorylation events were key to promoting translation.

Improvement Suggestions:

- Page 2, first paragraph, last sentence. The meaning of “4E-BP-dependent phenotypes” is not particularly clear.
- Page 2, line four of Materials and Methods. Shouldn’t “32″ be superscript?
- Page 2, last paragraph. The statement “Here, six differentially phosphorylated forms (A-F) ...” is confusing. Is form A really phosphorylated? Earlier in the manuscript, the authors mentioned 5 major phosphorylation sites.
- In Figure 1, the assay used in this Figure requires much better description in the text. Part A of the legend requires specification of the experimental approach. Also, the title of Figure 1 could be more useful if it were informative/conclusive rather than descriptive.
- Page 3, right column, second paragraph. The sentence “While endogenous... , suggesting that Thr46 phosphorylation is required for subsequent The37 phosphorylation under normal...”
growth conditions.” is misleading since 4E-BP1 mutant (Thr46Ala) is clearly phosphorylated at Thr37 in Fig. 2A middle panel. With the data presented here the reader is left wondering whether the differences in detecting HA-4E-BP1 (T37A) or HA-4E-BP1 (T46A) might be simply due to different affinities of the (Thr37/46) antibody used.

○ What is the difference between the middle and the lower panel? Are the antibodies used to probe the middle and the lower panel (e.g. phosho-HA-4E-BP1 (Thr37/46) vs. phosho-4E-BP1 (Thr37/46)) different or the same? When using cell lysates of stable cell lines and a phospho-4E-BP1 (Thr37/46) antibody, shouldn’t be HA 4E-BP1 wildtype and mutant forms be detectable in the lower panel as well?

○ Page 3, last paragraph, line 8. “modulate” would be clearer if described as the specific modulations that occur – in the literature a state of 4E-BP1 hypophosphorylation. The author understands that the changes are complicated and clarified from these authors' perspective in Figure 3 and in the Discussion. However, possibly, the reader could be better informed at this point of the manuscript.

○ Page 4, second paragraph, right-hand column. Please clarify the effects and usefulness of etoposide and the meaning of VCP and, in the legend, VCP/Chk2.

○ In Figure 2B, the authors should provide a control western blot of an unrelated protein which does not belong to the eIF4E-associated proteins under normal conditions to show the unspecific background of this assay. It is not clear to this reader why mono-phosphorylated 4E-BP1 (at position Thr70) is detectable in the non-Nocodazole treated cell samples in Fig. 2B (lane 3 and 5) when in Fig. 3 they show that mono-phosphorylation of 4E-BP1 at position Thr70 is triggered by a Nocodazol-induced block of Hela cells but not detectable under normal conditions.

○ In Figure 3, why did the authors mix Control and Nocodazole-blocked Hela cell lysates instead of showing them side-by-side? Also, why are there additional spots in the interpretation panels for Phos-Thr70 and Phos-Thr37/46 that have no counterpart in the respective western blot panels? This figure is already difficult, so the authors should be careful with the overlays.

○ In Figure 4, all results in this Figure are not adequately described. For example, the authors need to explain the analysis of ubiquitin and its significance to their studies.

○ The sentence “...rapamycin-insensitive phosphorylation site is alone in regulating eIF4E binding.” should read something like “...rapamycin-insensitive phosphorylation site is alone responsible in regulating eIF4E binding”.

**Competing Interests:** No competing interests were disclosed.

We confirm that we have read this submission and believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 30 July 2012

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Through binding to eIF4E, the 4EBP proteins (4EBP1, 2, 3) interfere with formation of the eIF4F cap-binding complex. A key function of the mTOR serine-threonine kinase is to phosphorylate 4EBPs on multiple sites, which blocks their ability to bind to eIF4E thus potentiating cap-dependent translation.

Recently it has become clear that ATP-competitive mTOR kinase inhibitors are more effective than the allosteric compound rapamycin at blocking mTOR-dependent 4EBP phosphorylation, particularly on sites T37 and T46. This greater effect of mTOR kinase inhibitors on 4EBP phosphorylation correlates with increased anti-proliferative and pro-apoptotic effects. The purpose of the current study is to determine the order of phosphorylation of sites T37 and T46, and which site is most important for regulating binding of 4EBP1 to eIF4E. This is a fairly narrow question to address but the authors argue that it is important for understanding drug mechanism and that previous approaches have not addressed this question adequately.

Overall the data support the conclusion that T46 phosphorylation is the initial, rapamycin-insensitive event that blocks 4EBP1-eIF4E binding. The main approach used is two-dimensional electrophoresis (IEF followed by SDS-PAGE), then far-western blotting with labeled eIF4E to determine which spots are capable of eIF4E binding. Figure 1A provides a 1-D blot that shows convincingly that the far-western technique can reveal increased binding of eIF4E to 4EBP1 in cells treated with the mTOR kinase inhibitor PP242. Figure 1B shows that 4EBP1 phosphorylation and release from eIF4E are induced by serum in a mTOR-dependent manner. Figure 1C presents the 2D data illustrating hierarchical phosphorylation and distinct effects of rapamycin and PP242. The data support the conclusion that only a small subset of 4EBP forms are capable of binding eIF4E, that these forms lack T37/46 phosphorylation, and that PP242 increases this pool to a greater extent than rapamycin. To distinguish the roles of T37 and T46 they express HA-tagged 4EBP1 and alanine substitution mutants in HeLa cells. The data in Figure 2A show that T46A mutation, but not T37A mutation, blocks detection by the phospho-T37/46 antibody. This suggests that T46 phosphorylation precedes and is
required for T37 phosphorylation. A cap binding assay in Figure 2B supports the conclusion that T37/46-phosphorylated 4EBP1 cannot bind eIF4E in the cap-binding complex, but this is not novel and the technique does not add further evidence for initial T46 phosphorylation.

The rest of the data seem added on and superfluous to the main message of the paper. The experiment in Figure 3 shows 2D far western blotting data on lysates of nocodazole-treated cells, which artificially increases the diversity of 4EBP1 spots. As the authors point out, these might not exist under physiological conditions so the relevance is unclear. Figure 4 addresses an entirely different problem, namely the lack of selectivity of the PP242 compound. While it is of some interest that PP242 differs from other compounds of this class in its apparent inhibition of DNA-PK and related kinases, this observation is too peripheral to the main study and should be developed further before publication.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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# Comments on this article

**Version 1**

Reader Comment 20 Jul 2012

**Juming Jan**, State Key Laboratory of Phytochemistry and Plant Resources in West China, Kunming Institute of Botany, Chinese Academy of Sciences, China

The paper clearly elucidate the functionally important role of the phosphorylated Thr46 of 4EBP1 in the course of 4EBP1 binding with eIF4E. the design of experiments is very persuading, especially the 4EBP1 point mutant and pull down of eIF4E give a strong proof to unravel the crucial role of Thr46 phosphorylation in binding eIF4E. Thank the authors for providing the powerful evidences to make me under the mechanism of the interaction between 4EBP1 and eIF4E.

The eIF4E are released from the 4EBP1 in many malignant tumor due to the hyperphosphorylation of 4EBP1. It will be very helpful to discover some new mTOR signal inhibitors and to explain the mechanism, if the processes of 4EBP1 dephosphorylation are clarified.

**Competing Interests:** No competing interests were disclosed.
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