Target of Rapamycin Complex 2 Regulates Actin Polarization and Endocytosis via Multiple Pathways*§

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Background: TORC2/Ypk1 regulates actin polarization and endocytosis via unknown effectors.

Results: Pharmacological inhibition of TORC2 reveals that flippase kinases and biophysical properties of the plasma membrane are major effectors of TORC2.

Conclusion: TORC2 regulates actin and endocytosis via multiple pathways, each with different signaling kinetics.

Significance: Elucidation of TORC2 effector pathways in yeast will inform future studies in higher eukaryotes.

Target of rapamycin is a Ser/Thr kinase that operates in two conserved multiprotein complexes, TORC1 and TORC2. Unlike TORC1, TORC2 is insensitive to rapamycin, and its functional characterization is less advanced. Previous genetic studies demonstrated that TORC2 depletion leads to loss of actin polarization and loss of endocytosis. To determine how TORC2 regulates these readouts, we engineered a yeast strain in which TORC2 can be specifically and acutely inhibited by the imidazoquinoline NVP-BHS345. Kinetic analyses following inhibition of TORC2, supported with quantitative phosphoproteomics, revealed that TORC2 regulates these readouts via distinct pathways as follows: rapidly through direct protein phosphorylation cascades and slowly through indirect changes in the tensile properties of the plasma membrane. The rapid signaling events are mediated in large part through the phospholipid flippase kinases Fpk1 and Fpk2, whereas the slow signaling pathway involves increased plasma membrane tension resulting from a gradual depletion of sphingolipids. Additional hits in our phosphoproteomic screens highlight the intricate control TORC2 exerts over diverse aspects of eukaryote cell physiology.

The target of rapamycin (TOR)§ is a serine/threonine (Ser/Thr) kinase conserved in nearly all eukaryotes (1). Missense mutations in TOR can confer resistance to the growth-inhibitory properties of the natural product macrolide rapamycin, and this property led to the discovery of TOR in a selection of rapamycin-resistant Saccharomyces cerevisiae mutants (2–5). Metazoans typically encode a single TOR gene (mTOR in mammals), whereas the yeast S. cerevisiae encodes two TOR genes (TOR1 and TOR2).

TOR proteins form at least two distinct multiprotein complexes known as TOR complex 1 (TORC1) and TOR complex 2 (TORC2), and each performs one or more essential functions in both S. cerevisiae and mammals (6). In S. cerevisiae, there are two variants of TORC1; the main variant, TORC1-A, contains Tor1 as the catalytic subunit, and the minor variant, TORC1-B, contains Tor2 as the catalytic subunit (Fig. 1B) (7). The kinase activities of TORC1-A, TORC1-B, and mammalian TORC1 (mTORC1) are all acutely sensitive to rapamycin. In contrast, neither TORC2 nor mTORC2 is sensitive to acute exposure to rapamycin (7–9), although prolonged exposure can inhibit mTORC2, at least in certain cell types (10).

Rapamycin has been an incredibly valuable tool to dissect TORC1 functions, particularly in S. cerevisiae where large scale phosphoproteomics studies have identified dozens of proximal and distal effectors downstream of this kinase complex (11, 12). These data establish the groundwork for a molecular understanding of how TORC1 regulates many diverse aspects of eukaryote physiology, including anabolism, autophagy, and longevity (13). Lacking a specific small molecule inhibitor, it has been more challenging to characterize the effectors downstream of TORC2. Recently, the phosphoproteomes of mammalian cells treated with ATP-competitive small molecules that inhibit both mTORC1 and mTORC2 have been obtained (14–

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5 The abbreviations used are: TOR, target of rapamycin; CWI, cell wall integrity; Fpk, flippase kinase; SC, synthetic complete; SPT, serine palmitoyltransferase; mTOR, mammalian TOR.
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16). Although these studies represent a significant increase in our understanding of the mTOR-dependent phosphoproteome, coverage is an issue, and it remains a challenge to partition these phosphorylation events downstream of mTORC1 versus mTORC2.

Dissection of the TORC2 function in S. cerevisiae has relied heavily on the use of temperature-sensitive alleles of TOR2 or genes encoding other essential components of TORC2. These tools led to the discovery that Ypk1 and Ypk2, a pair of AGC family Ser/Thr kinases, are the essential substrates of TORC2 (17, 18). Indeed, alleles of YPK1 or YPK2 that encode hyperactive kinases rescue the inviability of tor2 mutants (17, 18).

Through Ypk1 (the more prominent of these two Ypk2), TORC2 regulates polarization of the actin cytoskeleton (22, 23), and sphingolipid biosynthesis (24). TORC2/Ypk1 signals additionally regulate the activity of the protein phosphatase calcineurin (18, 25), and the protein kinase C/cell wall integrity (CWI) MAPK pathway (26). However, temperature-sensitive and other conditional alleles (e.g. gene expression driven by glucose-repressible promoters) are not optimal tools to probe TORC2 functions as environmental stresses, including temperature shifts and changes in carbon source per se, affect TORC2 activity and thus complicate interpretation of observations made from these experimental paradigms (27). Moreover, an additional shortcoming of using these conditional alleles is the long time (often hours) required to inactivate TORC2.

The biochemistry of sphingolipid biosynthesis is relatively well understood, and the mechanism by which TORC2 regulates its other described readouts directly or indirectly.

Like sphingolipids, phospholipids also affect many functions in a cell (38) and are downstream targets of TORC2/Ypk1. Specifically, Ypk1 was recently shown to phosphorylate and inhibit the flipase protein kinase Fpk1 and its paralog Fpk2 (39). Fpks are activators of aminophospholipid flipases Dnf1 and Dnf2 (Drse2 Neo1 family) that regulate phospholipid asymmetry in the plasma membrane. Additional targets of Fpks are suspected but have not yet been reported (40).

In this study, we wished to understand better how TORC2/Ypk1 regulates actin polarization and endocytosis. Specifically, we wished to determine whether these processes are directly targeted through phosphorylation events or are indirectly regulated through lipid messengers. To address this problem, we employed a reverse chemical genetic approach where we engineered TORC1 to be resistant to the ATP-competitive small molecule TOR inhibitor NVP-BHS345. Addition of NVP-BHS345 to these cells triggered a specific and acute inhibition of TORC2. Our subsequent observations indicate that TORC2 regulates actin polarization and endocytosis via distinct pathways involving the following: (i) the Fpks; (ii) Fpk-independent phosphorylation of proteins involved in endocytosis; and (iii) slower effects due to changes in membrane tension.

Experimental Procedures

Yeast Strains and Plasmids—Plasmids and strains used in this study are summarized in Tables 1 and 2. Yeast strains were generated through crosses or by transformation with the indicated plasmids or PCR fragments for homologous recombination. Tagged proteins are expressed from their own promoters. Unless indicated otherwise, yeast cultures were grown in synthetic complete (SC) medium lacking appropriate amino acids required for plasmid selection.

Drugs and Chemicals—Rapamycin (LC Laboratories) was dissolved in 90% ethanol, 10% Tween® 20 at 1 mg/ml and used at a final concentration of 200 nM. BHS345 (Novartis) was dissolved in DMSO at 10 mM and used at final concentrations ranging from 10 to 200 μM. Myriocin (Sigma) was dissolved in methanol at 2.5 mM and used at a final concentration of 2.5 μM. Acreosabin A (Takara) stocks were prepared in ethanol at 5 mM and used at 2.5 μM. 1NM-PP1 (Calbiochem®) was dissolved in DMSO at 1 mM and used at a final concentration of 500 nM. Chlorpromazine hydrochloride was used at 500 μM (Sigma) and dissolved in water.

Antibodies—The following antibodies were used in this study: goat anti-Ypk1 1:1000 (Cell Signaling); rabbit anti-phospho-Ypk1 Thr-662 1:500 (28); rabbit anti-phospho-Sch9 Thr-737 1:250 (41); mouse anti-HA 1:20,000 (Sigma); rabbit anti-phospho-Akt substrate (RXR(S/T)) 1:2000 (Cell Signaling); mouse anti-GFP 1:700 (Santa Cruz Biotechnology); rabbit anti-Hog1 1:10,000 (Santa Cruz Biotechnology), and the appropriate infrared dye-coupled secondary antibodies (all Alexa Fluor 680-conjugated secondary antibodies from Life Technologies, Inc., and all IRDye® 800 conjugated secondary antibodies from Rockland Bioconcept).

Protein Extraction and Western Blotting—Yeast cells were grown at 30 °C to an A600 of 0.6–0.8, followed by drug or mock treatment. After treatment, protein extracts for immunodetection were prepared essentially as described previously (42).
For immunoprecipitation experiments, cells in urea buffer (25 mM Tris, pH 6.8, 6 M urea, 1% SDS) were lysed using a FastPrep bead beater (Lucerna-Chem) and glass beads (RetseTM, Fisher Scientific). Extracts were diluted in IP buffer (PBS, 10% (v/v) glycerol, 0.5% Tween 20) and glass beads (RetseTM, Fisher Scientific). Extracts were diluted in IP buffer (PBS, 10% (v/v) glycerol, 0.5% Tween 20) and glass beads (RetseTM, Fisher Scientific).

**Table 1**

| Strain       | Genotype                        | Source or Ref. |
|--------------|---------------------------------|---------------|
| TB50         | MATa leu2-3,112 ura3-52 trp1 his3 rme1 HMLa | Lab stock     |
| MS183        | TB50a 3HA-TOR1M2282T             | This study     |
| MS184        | TB50a 3HA-TOR1M2226T             | This study     |
| MS185        | TB50a 3HA-TOR1M2226T             | This study     |
| MS186        | TB50a 3HA-TOR1M2226T             | This study     |
| RL230-2b     | TB20a tor2::KanMX4 YEp352 YPK2::HIS3-4A -HA | This study     |
| DR4AY        | TB50a orml1::HIS3MX6 orml2::TRPI | This study     |
| DR5Y1        | TB50a orml1::HIS3MX6 orml2::TRPI | This study     |
| RL276-3D     | TB50a HIS3 TRPI                 | This study     |
| DR5Y9        | TB50a/tor2::KanMX4 HIS3/his3 TRP1/trp1 | This study     |
| DR550        | TB50a/tor2::KanMX6 orml1::HIS3/his3 orml2::TRP1/trp1 | This study     |
| SE13-01      | TB50a 3HA-TOR1M2226T             | This study     |
| SE14-15      | TB50a 3HA-TOR1M2226T             | This study     |
| SE14-18      | TB50a 3HA-TOR1M2226T             | This study     |
| SE14-20      | TB50a 3HA-TOR1M2226T             | This study     |
| SE15-11      | TB50a 3HA-TOR1M2226T             | This study     |
| SE15-13      | TB50a 3HA-TOR1M2226T             | This study     |
| SE15-46      | TB50a 3HA-TOR1M2226T             | This study     |
| SE15-49      | TB50a 3HA-TOR1M2226T             | This study     |

**Table 2**

| Name         | Description                        | Source or Ref. |
|--------------|------------------------------------|---------------|
| YEp352      | Lab stock                           |               |
| pRS415      | Lab stock                           |               |
| pRS16       | Lab stock                           |               |
| pMS080      | pRS415_ENT1-5HA-TADH               | This study     |
| pMS123      | pRS415_PAN1-GFP-TADH               | This study     |
| pUC52p      | YEp352_YPK2::HIS3-4A             | 17            |
| pRS416      | pRS416_YPK2::HIS3-4A             | This study     |
| pUC6A       | pUC57_ENT1-T27A S88A T160A S175A S401A T450A | GenScript     |
| pUC6D       | pUC57_ENT1-T27D S88D T160D S175D S401D T450D | GenScript     |
| pSE15-01    | pRS415_ENT1-T160A S175A-TADH     | This study     |
| pSE15-07    | pRS415_ENT1-T27A S88A T160A S175A S401A T450A-5HA-TADH | This study     |
| pSE15-08    | pRS415_ENT1-T27D S88D T160D S175D S401D T450D-5HA-TADH | This study     |

**Phospho-proteomics Sample Preparation and Analysis by Label-free Quantification**—For each of the three sample replicates, 3 mg of protein was digested with trypsin (1:125 w/w) and cleaned by reverse phase chromatography. Phosphopeptide enrichment was performed with titanium dioxide resin (1.25 mg of resin for each sample). Isolated phosphopeptides were analyzed by an LTQ Orbitrap XL (Thermo) (43). The mass spectrometry data were searched against an S. cerevisiae genome database decoy database for yeast proteins using Sequest (44). OpenMS version 1.8 (45) was used both to detect MS1 features and to align them between the different mass spectrometry runs. By using a decoy database (46), a Peptide Prophet’s probability threshold was computed to achieve a false discovery rate below 1% and was used to filter OpenMS results. Phosphopeptide features with identical sequence and phosphorylation state but different charge states were merged together. Statistical analysis was performed by MSstat R package (47), which fits the data to an analysis of variance model.

**Analysis of MS Data Normalization**—The MS intensity of each peptide in each condition was recorded; data were log2 transformed to comply with linear regression requirements. In situations where the recorded values are null, we will assume that they correspond to situations where the peptide was not identified.

**Quality Control and Principal Component Analysis**—Sample quality was assessed using principal component analysis as follows: based on data from all peptides, the sample labeled...
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Area_6 was removed from experiment 1, and samples labeled Cond2 and Cond3.2 were removed from experiment 2. We then removed all peptides that added any missing value in the control group of interest to the analysis (DMSO) or the treated groups of interest (WT|BHS345, WT|rapamycin). Peptides that had all values below the 2.5% quantile in one group of samples, but no missing values in the other group, were rescued for analysis. Once the bad samples and missing values were removed, the remaining samples were well separated according to treatment for both experiments 1 and 2.

Model Building and Analysis—For both experiments, the intensity of the remaining peptides was modeled after the interaction of treatment (BHS345 at the doses noted or rapamycin at the doses noted). The contrasts were calculated to provide a log fold change and a p value, and the data are available in the supplemental Tables S1 and S2.

Microscopy—Cells were grown in synthetic media at 30 °C to an 

Acute Inhibition of TORC2 Rapidly Arrestrs Endocytosis and Triggers Actin Depolarization—Previous genetic studies demonstrated that gradual depletion of TORC2 eventually (after several hours) triggers inhibition of endocytosis and depolarization of the actin cytoskeleton. To begin to characterize the signaling outputs of TORC2 that impinge upon these distal readouts, we first determined the time required for BHS345-induced TORC2 inhibition to block endocytosis and depolarize actin.

Fluid-phase endocytosis is conveniently monitored by the internalization of the fluorescent compound Lucifer Yellow (52). Inhibition of TORC2 with BHS345 arrested endocytosis within 1 h (Fig. 2A). Consistent results were obtained using the lipophilic styryl dye MM4-64 which in WT cells, but not endocytosis-deficient cells, accumulates in the vacuolar membrane (53) as follows: inhibition of TORC2 abrogated the vacuolar membrane accumulation of MM4-64, which instead partially accumulated in cytoplasmic structures (Fig. 2B). These results confirm that, like depletion of TORC2 activity through condi-
BHS345, an ATP-competitive TOR inhibitor as a tool to study TORC2 function. A, structures of BHS345, used here, and the related PI3K/mTOR inhibitor BEZ235. B, schematic of the TOR complexes in S. cerevisiae. Rapamycin-sensitive TORC1 is found in two variants, one with Tor1 as the catalytic component (TORC1-A) and the other with Tor2 as the catalytic component (TORC1-B). Tor2, but not Tor1, functions in rapamycin-insensitive TORC2. TORC1-A, TORC1-B, and TORC2 are all inhibited by BHS345. C, BHS345 dose-response curves of WT, TOR1^MT (TOR1^M2282T), TOR2^MT (TOR2^M2286T), and TOR1^MT TOR2^MT strains grown in SC. A_{50} was determined after 30 h of incubation at 30 °C. D, TORC1 and TORC2 activity status, respectively, assayed by Western blot detecting Sch9^{Thr717} and Ypk1^{Thr662} phosphorylation, in the indicated strains before and after treatment with rapamycin (200 nM, 20 min) or BHS345 (10 μM, 12 min).

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The effect on endocytosis and depolarization of actin structures observed upon TORC2 inhibition was suppressed in cells expressing hyperactive Ypk1^{D242A} (Fig. 2, D–G) (18). These phenomena were also suppressed in cells expressing a BHS345-resistant allele of TOR2 (data not shown) and recapitulated upon inhibition of analog-sensitive Ypk1 with 1NM-PP1 (data not shown (18)). Together, these observations confirm that regulation of endocytosis and actin polarization are Ypk-dependent outputs of TORC2 and exclude the possibility of off-target effects of BHS345. The relatively fast kinetics suggest that TORC2 signals to these effectors directly, and we next set out to characterize the relevant TORC2 effector pathways.

Essential Function of TORC2 Is to Maintain Sphingolipid Biosynthesis—TORC2, via Ypk1, stimulates sphingolipid synthesis by inactivating the SPT inhibitors Orm1 and Orm2 (Fig. 3A), and we wondered to what extent the sphingolipids influence TORC2 readouts. Initially, we asked whether the lethality of TOR2 deletion is due to insufficient levels of sphingolipids. This appears to be the case as the growth of tor2 cells can be rescued by double deletion of ORM1 and ORM2 (Fig. 3B and C). However, further experiments with orm1 orm2 cells were not possible, as the slow growth phenotype of these cells was rapidly suppressed. Instead, we made use of two low molecular weight compounds known to specifically reduce sphingolipid biosynthesis; myriocin inhibits SPT, and aureobasidin A blocks conversion of phytoceramide into inositol phosphoceramide. We treated WT cells to assess whether these treatments mimic BHS345-induced readouts. Indeed, consistent with previous work (37, 55, 56), we found that block of complex sphingolipid biosynthesis with myriocin and aureobasidin A treatments decreased the uptake of Lucifer Yellow (Fig. 3D), led to aberrant vacuolar staining with MM4-64 (Fig. 3E), and triggered actin depolarization (Fig. 3F).

The above observations are consistent with the idea that TORC2 regulates these readouts via sphingolipid intermediates. However, several observations suggest that this is not the primary mechanism by which TORC2-mediated signals impinge upon these readouts. Both myriocin and aureobasidin...
A inhibited endocytosis and depolarization of the actin cytoskeleton, but fewer cells within the population were affected as compared with BHS345 treatment. Furthermore, the time required to achieve actin depolarization in 50% of the cells was ~90 min for myriocin/aureobasidin A but only ~30 min for BHS345 (Figs. 2C and 3F). These significant penetrant and temporal differences suggest that despite being essential targets of TORC2 (Fig. 3, B and C), sphingolipids play only a partial and slow role in coupling TORC2 signals to the actin polarization and the endocytic machinery.

We wished to determine the mechanism by which sphingolipid depletion inhibits endocytosis and depolarizes actin. Sphingolipids are a major component of the yeast plasma membrane, and we have previously suggested that their depletion could increase the line tension of the plasma membrane (28). Prior studies have linked plasma membrane stress to actin depolarization (57). We confirmed in our strain background that treatments affecting the tensile properties of the plasma membrane, such as hyp-osmotic shock, heat shock, and addition of chlorpromazine (an antipsychotic compound known for its propensity to intercalate into lipid bilayers (58)), all elicit actin depolarization and presumably plasma membrane tension as readout by increased phosphorylation of Ypk1Thr-662 (Fig. 4, A and B). These observations support the notion that sphingolipid depletion blocks endocytosis and depolarizes actin by altering the physical properties of the plasma mem-
brane. To test this further, we asked whether inclusion of the osmotic stabilizer sorbitol in the growth medium would suppress actin depolarization induced by myriocin and aureobasidin A treatments. Inclusion of sorbitol in the media prevented actin depolarization normally induced by myriocin and aureobasidin A (Fig. 4C), but it did not block actin depolarization induced by BHS345-mediated TORC2 inhibition (Fig. 4D). From these observations, we conclude that sphingolipid depletion upon TORC2 inhibition gradually increases membrane tension that subsequently blocks actin polarization and endocytosis. The observation that sorbitol does not suppress actin depolarization induced by TORC2 inhibition confirms that effectors other than sphingolipids must couple TORC2 to the actin cytoskeleton.

**TORC2 Does Not Signal to the Actin Cytoskeleton via the CWI Pathway**—Previous work has demonstrated that hyperactivation of the CWI pathway restores actin organization and growth to cells possessing reduced TORC2 function (19, 20, 59). The CWI pathway is composed of the cell-surface sensors Wsc1, -2, -3, Mid2, and Mtl1, which activate the Rho1 GTPase via the guanine nucleotide exchange factors Rom1 and Rom2. One of Rho1’s many targets is Pkc1, which in turn activates the MAPK cascade consisting of Bck1, Mkk1/2, and Slt2. It is believed that TORC2 acts upstream of the CWI pathway by stimulating the exchange activity of Rom2 (26). Based on these previous reports, it seemed a likely possibility that TORC2 regulates actin polarization via activation of the CWI pathway. This, however, appears not to be the case as acute inhibition of

**FIGURE 3. Sphingolipids are major mediators of TORC2 signaling.** A, schematic of sphingolipid biosynthesis in yeast and its regulation by TORC2/Ypk1 via ORM1/2. Myriocin and aureobasidin A antagonize serine palmitoyltransferase and inositol phosphoceramide synthase, respectively. LCB, long-chain base; LCB-1P, long-chain base 1-phosphate; PHC, phytoceramide; IPC, inositol phosphoceramide; MIPC, mannosyl-inositol phosphoceramide; M2IP2C, mannosyl-diinositol phosphoceramide. B and C, tor2 lethality is suppressed by deletion of ORM1 and ORM2. Spore colonies obtained upon dissection of TOR2/tor2 (DRY49) and TOR2/tor2 ORM1/orm1 ORM2/orm2 (DRY50) diploids. No viable tor2 cells were obtained from 10 dissected ascii obtained from TOR2/tor2 diploids (left panel). In contrast, viable tor2 cells (marked with red circles, right panel) were readily observed from dissected ascii obtained from TOR2/tor2 ORM1/orm1 ORM2/orm2 diploids. These tor2 haploids all additionally harbored the orm1 and orm2 deletions. C, Om2 double deletion suppresses lethality of tor2, demonstrated by the ability of tor2 orm1 orm2 cells to lose a URA3-marked plasmid containing YPK2D239A on 5-fluoroorotic acid (5-FOA) medium. D, micrographs of TOR1MT cells treated for 90 min with Lucifer Yellow (LY) and either myriocin (Myr) or aureobasidin A (AbA) as indicated. D, right panel, quantification of the vacuolar versus cytoplasmic accumulation of Lucifer Yellow after the indicated treatments. Statistically significant differences are indicated (Student’s t test) with $p < 0.001 (***)$. E, micrographs showing MM4-64 accumulation in TOR1MT cells after the indicated treatments. Numbers represent the percentage of cells with vacuolar labeling. F, micrographs of rhodamine phalloidin stained TOR1MT cells after the indicated treatments. D–F, cells were treated with 2.5 μm myriocin or 2.5 μm aureobasidin A.
TORC2 with BHS345 results in activation of the CWI pathway as readout by increased phosphorylation of Slt2 (Fig. 5A). From these results, we conclude that actin depolarization induced by TORC2 inhibition is not caused by inactivation of the CWI pathway.

**Fpk1/2 Hyperactivation upon TORC2 Inhibition Is a Major Contributor to Actin Depolarization and Inhibition of Endocytosis**—Like Orm1 and Orm2, Fpk1 and Fpk2 are inhibited through direct phosphorylation by Ypk1, and, like orm1 orm2 mutants, fpk1 fpk2 mutants also restore growth of ypk1 hypomorphic strains and ypk1ΔΔypk2 (39). Consistently, FPK1/2 deletion improves the growth of cells in which TORC2 is inhibited with BHS345 (Fig. 5B), but unlike ORM1, ORM2 deletion does not restore growth of TOR2 deletion (data not shown). Building on these observations, we found that actin polarization and endocytosis were largely unaffected by TORC2 inhibition (data not shown). A quadruple deletion of dnf2 dnf1 dnf2 dnf3 is lethal (61), and therefore, we cannot exclude the possibility that Fpks indeed regulate actin polarization and endocytosis redundantly through the activity of these phospholipid translocases. However, it seems more likely that other yet-to-be described effectors of Fpk1/2 serve to couple TORC2 to actin and endocytosis.

**Quantitative Mass Spectrometry Identifies New Effectors Downstream of TORC2**—To better understand how TORC2 signals impinge upon its distal readouts in general and actin polarization and endocytosis in particular, we performed quantitative phosphoproteomics using BHS345. We previously used quantitative mass spectrometry to define the rapamycin-sensitive phosphoproteome of *S. cerevisiae* cells and identified many novel TORC1 effectors (11, 62). Having a yeast-permeant TORC2 inhibitor allowed us to similarly ascertain the TORC2-dependent phosphoproteome by comparing the rapamycin-sensitive phosphoproteome to the BHS345-sensitive phosphoproteome. As a prelude to this experiment, we first determined the length of time required for rapamycin and BHS345 to efficiently inhibit TORC1 as assessed by loss of phosphorylation on Thr-737 in the hydrophobic motif of Sch9. We found that the effect of BHS345 on TORC1 was detectable more rapidly than the effect due to rapamycin treatment; Sch9 was dephosphorylated after 1 min of BHS345 treatment (51) compared with 8 min required for rapamycin treatment (data not shown). Based network (61)). Similar results (i.e. only slight suppression) were obtained with lem3, dnf1 dnf2, dnf1 dnf3, dnf1 dnf2 lem3, dnf1 dnf2 dnf3 drs2, and dnf3 drs2 cells (data not shown).
on this observation, we designed experiments with short compound exposure time (≤30 min) to capture relatively early events downstream of TORC1 and TORC2 inhibition. We performed two separate experiments according to the flowchart in Fig. 6A, to cover different time points and inhibitor concentrations. Each experimental dataset was analyzed independently to retain phosphopeptides with a greater than 2-fold change and with a \( p \) value of ≤0.05 for compound versus vehicle treatment (supplemental Tables 1 and 2). In the first experiment, 123 peptides were found to be dephosphorylated and 44 hyperphosphorylated by BHS345 while remaining unaffected by rapamycin. In the second experiment, 58 and 36 peptides were similarly affected, respectively. These peptides are presumably regulated downstream of TORC2 and not TORC1. Enriched among these peptides were “RXXS” and “SP” motifs (where S is the phosphorylated residue). RXXS is a minimal recognition sequence for AGC kinases, and its enrichment could suggest that many of the TORC2-dependent phosphoproteins that we identified are direct targets of Ypk1.

The union of affected genes from the two experiments provided us with a working list of 117 proteins that are hypophosphorylated (Table 3) and 47 proteins that are hyperphosphorylated (Table 4) upon inhibition of TORC2. Having identified these putative novel effectors downstream of TORC2, we attempted to validate a subset of them.

Molecular Links between TORC2, Actin Organization, and Endocytosis—Our phosphoproteomic experiments identified several proteins implicated in the regulation of actin organization and endocytosis (Tables 3 and 4) (63, 64). Highly conserved, clathrin-mediated endocytosis involves internalization and pinching off of the plasma membrane to generate an endocytic vesicle (Fig. 6B). Strikingly, we observed that BHS345, but not rapamycin, induced changes in phosphorylation of both coat proteins as well as regulators of actin polymerization (Fig. 6B and supplemental Tables 1 and 2). Coat disassembly is mediated by the Ark family proteins, which, in S. cerevisiae, is made up of Ark1, Prk1, and the lesser studied Akl1 (65). Suggestively, we found Prk1 and Akl1 to be hypophosphorylated upon BHS345 treatment. Known substrates of these kinases include the coat proteins Pan1, Sla1, End3, Ent1, and Ent2, and indeed, these were all found to be hypophosphorylated upon BHS345 treatment. In the cases of Pan1, and Ent2, the residues found to be hypophosphorylated included residues in Ark consensus motifs ((L/I/V)XX(Q/N/T))XTG). In low throughput assays, we confirmed that Ent1 and Pan1 are dephosphorylated after BHS345 or 1NM-PP1-mediated inhibition of Ypk1 but not rapamycin treatment (Fig. 6, C and D and data not shown), demonstrating that they are TORC2/Ypk1 effectors. BHS345-induced dephosphorylation was suppressed in cells expressing Ypk1D242A (Fig. 6, C and D), and by TOR2MT (data not shown).
FIGURE 6. TORC2 regulates phosphorylation of endocytic proteins independently of sphingolipids and Fpk proteins. A, experimental flowchart for the two independent phosphoproteomic experiments, each performed with biological triplicates. B, schematic of a budding vesicle with endocytic proteins. Red Ps demark proteins that become hypophosphorylated upon TORC2 inhibition. C, TOR1\textsuperscript{MT} cells expressing Ent1–5HA and either an empty vector or a vector expressing hyperactive Ypk1\textsuperscript{D242A} were treated as indicated. Denatured protein extracts were subsequently made, and RX(x/S/T) motif phosphorylation of immunoprecipitated Ent1–5HA was assessed by Western blotting. D, TOR1\textsuperscript{MT} cells expressing Pan1-GFP and either an empty vector, or a vector expressing hyperactive Ypk1\textsuperscript{D242A}, were treated as indicated. The extent of Pan1-GFP phosphorylation was assessed by SDS-PAGE migration shifts. Rap, rapamycin. D and E, deletion of FPK1 and FPK2 does not block BHS345-induced Ent1 and Pan1 hypophosphorylation. C–E, BHS345 (BHS) was used at 15\% H9262 Myr at 2.5\% H9262 and aureobasidin A (AbA) at 2.5 \mu M.

TABLE 3

| TORC2-dependent hypophosphorylated proteins | Genes |
|---------------------------------------------|-------|
| **Biological process** | **Genes** |
| Cytoskeleton organization | AKL1, AVO1, BBC1, BUD44, BZZ1, CDC41, CLA4, ENT1, ENT2, GIN4, KAR3, NAP1, PAN1, PEA2, PRK1, SAC7, SLA1, SLM2, VHS2, VRP1, YSC4 |
| Transcription from RNA polymerase II promoter | ABF1, CAF120, CYC8, DCP2, DOT6, FHL1, FHK2, MKS1, MSN4, NAP1, NRD1, PRR1, ROX1, RSC30, SFT5, STR1, SWI5, TOP1, WHI5, YTA7 |
| Mitotic cell cycle | CDC14, CLA4, FHK2, GIN4, KAR3, MCK1, PAN1, PEA2, PIN4, PTK2, SIS2, SMM1, STB1, SWI5, TOP1, VHS2, VRP1, WHI5, ZDS1 |
| Protein phosphorylation | AKL1, CBK1, CLA4, GIN4, HAL5, LRE1, MCK1, PRK1, PRR1, PTK2, SKY1, YPK1 |
| Cellular response to DNA damage stimulus | ABF1, BDF1, CBK1, CYC8, MCK1, NUP60, PIN4, RFA1, RSC30, SFT5 |
| Regulation of organelle organization | BDF1, BUD44, BZZ1, CDC41, CLA4, ENO1, KAR3, NAP1, PEAC, SAC7 |
| Signaling | AVO1, BOI2, GIN4, LCB3, MDS3, MSN4, PXL1, SAC7, SLM2, TIP41 |
| Regulation of cell cycle | CDC14, CLA4, GIN4, KAR3, SIS2, SMM1, VRP1, ZDS1 |
| Chromatin organization | ABF1, BDF1, CYC8, FHK2, MSN4, NAP1, RSC30, TOP1 |
| Lipid metabolic process | CYC8, IST2, LAG1, LCB3, MSU4, PAH1, SUI1, YPK1 |
| Nuclear transport | CLA4, LTV1, NAPI, NIP2, NUP60, SKY1, SRF40, ZDS1 |
| Regulation of DNA metabolic | BDF1, FHK2, HST2, SPT5, STB1, SWI5, TOP1, VTS1 |
| Organelle fission | CDC14, CLA4, KAR3, MCK1, MSC3, RFA1, SWI5, TOP1 |
| Response to chemical | CLA4, LTV1, MSN4, PEAC, PRR1, RAS2, SKY1 |
| DNA repair | ABF1, BDF1, MCK1, NUP60, RFA1, RSC30, SFT5 |
| Protein targeting | APL6, ATG20, CLA4, NAPI, NIP2, NUP60, SKY1 |
| Protein complex biogenesis | BUD44, BZZ1, DCP2, GIN4, NAP1, SEC16, SLA1 |
| Cell budding | BOI1, BOI2, GIN4, NAPI, PAN1, PEA2, VRP1 |
| Meiotic cell cycle | CDC41, KAR3, MCK1, MSC3, MS54, RAS2, RFA1 |
| Cell wall organization or biogenesis | AVO1, LRE1, SAC7, SLA1, SM1, SSD1, ZEO1 |
| Cytokinesis | BOI1, BOI2, PAN1, PEA2, VHS2, VRP1, ZDS1 |
| Other functions | AC4, BSC6, CAN1, CTR1, DBP10, EFR3, FL33, GIP3, GLC8, GPD1, GSY1, GSY2, HHK1, IMH1, LS83, MAK11, MAK5, MAM3, MRH1, MSLS, MUK1, NIP1, NOP12, OM45, PHB2, PTK2, PIF2, RBS1, RPN13, SER33, SKI3, THS1, YEF3 |
TABLE 4

| TORC2-dependent hyperphosphorylated proteins | Genes |
|---------------------------------------------|-------|
| Response to chemical                        | MIG1, MLF3, RCK2, SCH9, SCP160, ST20, YCK2 |
| Protein phosphorylation                     | ALK1, HIP1, KSP1, RCK2, SCH9, ST20, YCK2 |
| Transcription from RNA polymerase II promoter | GSI1, HPc2, MIG1, REG1, RPH1, SCH9 |
| Carbohydrate metabolic process              | NUS1, PFK2, REG1, TSL1, VAN1 |
| Mitotic cell cycle                          | ALK1, HPC2, REI1, ST20 |
| Ion transport                               | AVT1, CCC1, PFK2, YBT1 |
| Lipid metabolic process                     | GSI1, PCT1, PK1, SCH9 |
| Organelle fission                           | ALK1, RCK2, SCP160, ST20 |
| Signaling                                   | KSP1, RCK2, SCP160, ST20 |
| Response to starvation                      | GSI1, KSP1, MIG1, REG1 |
| Proteolysis involved in cellular protein catabolic | DIA2, RAD23, RPN8 |
| Chromatin organization                      | GSI1, HPc2, RPH1 |
| DNA replication                             | DIA2, HEK2, ORC6 |
| Endocytosis                                 | ALY2, SWH1, YCK2 |
| Ribosomal large subunit biogenesis          | NOP8, REI1, RRP15 |
| Response to osmotic stress                  | RCK2, SCH9, ST20 |
| Other functions                             | BUL1, DED1, EAP1, FEN1, GLY1, JSN1, LYS21, NGR1, SEC31, SKG1, SMY2, SOLL, TIF3, YTA6 |

TORC2 regulates actin and endocytosis via multiple pathways. The TORC2-regulated Pan1 phosphorylation sites were found in Ark consensus motifs. Previous work demonstrated that alanine replacement of threonines within the 15 (L/i/V)XX(Q/N/T)XTG motifs in Pan1 (Pan1–15TA) causes severe defects in actin organization and uptake of Lucifer Yellow (66). TORC2 inhibition also affects phosphorylation of Ark consensus motifs in Ent1. It has been proposed that phosphorylation at these sites alters the actin-binding properties of Ent1 (67). It is possible that TORC2 regulates actin organization and endocytosis via regulation of Ark1/Prk1 activity. Alternatively, Ark1/Prk1 activity may be affected indirectly as a consequence of actin depolarization.

We additionally found that TORC2 inhibition affects Ent1 phosphorylation within RXX(S/T) motifs favored by ACG family kinases (Fig. 6) suggesting that these sites may be directly targeted by Ypk1. Indeed, Ent1 dephosphorylation upon TORC2 inhibition was very fast, with a significant loss of phosphorylation after 7.5 min of Ypk1 inhibition (data not shown). However, we were unable to observe direct phosphorylation of Ent1 by Ypk1 in vitro kinase assays (data not shown). Thus, it remains unclear whether Ent1 is directly phosphorylated by Ypk1.

To our knowledge, a function for the RXX(S/T) phosphorylation sites in Ent1 has not been investigated. Ent1 has six RXX(S/T) motifs, two of which, Thr-160 and Ser-175, are known to be phosphorylated (S. cerevisiae genome database). Thr-160 was also identified as a TORC2 effector in our present phosphoproteomics study. Conversion of Thr-160 and Ser-175 to alanine reduced but did not eliminate TORC2-dependent phosphorylation on RXX(S/T) motifs (Fig. 7A). Therefore, we mutated the serines and threonines within all six RXX(S/T) motifs into alanine or aspartate. In these mutants (Ent1–6A and Ent1–6D) phosphorylation within RXX(S/T) motifs was no longer detectable (Fig. 7A).

To assess the functional importance of Ent1 RXX(S/T) phosphorylation, we expressed Ent1–2A (data not shown), Ent1–6A, and Ent1–6D in ent1 ent2 cells. With the exception of Ent1–6D, which slightly reduced basal Lucifer Yellow uptake, expression of none of these constructs affected bulk endocytosis or altered the BHS345-induced endocytic arrest (Fig. 7, B–D).

Interestingly, actin was moderately depolarized in cells expressing Ent1–6D but not the Ent1–2A (data not shown) or Ent1–6A constructs (Fig. 7E). Thus, it appears that Ent1 phosphorylation within the RXX(S/T) motifs indeed plays some role in TORC2-dependent polarization of the actin cytoskeleton.

**Discussion**

Using BHS345, a novel, small molecular inhibitor of TOR signaling in yeast, we have investigated the readouts downstream of TORC2. In particular, we have exploited the ability to acutely inhibit TORC2 kinase activity to temporally dissect these readouts, something that was not previously possible using chronic genetic approaches that slowly deplete TORC2 from cells. Although the minor TORC1-B containing Tor2 is also inhibited in our experimental setup, we could confirm that the BHS345-induced phenomena were due to TORC2 inhibition by showing the following: 1) they are not induced by rapamycin; 2) they are rescued in cells expressing a hyperactive allele of Ypk1 (Ypk1D242); and/or 3) they are recapitulated upon inhibition of analog-sensitive Ypk1. This work complements a recent study by Shokat and co-workers (50) using a related mTOR inhibitor BEZ235 (Fig. 1A) and an “analog-sensitive” allele of TOR2 to globally characterize TORC2 signaling by systematic genetic means in yeast. In this study, we have also begun to systemically define TORC2 signaling by using quantitative phosphoproteomics. Focusing on known distal effectors, we found that TORC2 regulates actin polarization and endocytosis through at least three distinct signaling pathways (Fig. 8) as follows: a slow pathway involving plasma membrane stress induced by depletion of complex sphingolipids, and two fast pathways, one of which is mediated by the flippase kinases.

**Slow Pathway**—Sphingolipids, particularly phosphorylated long chain bases, have been suggested to signal to the endocytosis and actin polarization machineries (68). Stimulation of
sphingolipid production through direct phosphorylation of the Orm proteins by Ypk1 appears to be the essential function of TORC2 (Fig. 3). Thus, we explored the possibility that TORC2 regulates actin and endocytosis via its control of sphingolipid synthesis. This hypothesis appears to be correct, albeit not in the manner we anticipated. Blocking sphingolipid synthesis...
with either myriocin or aureobasidin A depolarizes actin and blocks endocytosis and thus mimics TORC2 inhibition (Fig. 3). However, a specific role for long chain bases, or their phosphorylated derivatives, in this signaling is negated by the observation that both myriocin and aureobasidin A trigger these readouts equally. Also curious is the striking difference in time required for myriocin/aureobasidin A versus BHS345-induced TORC2 inhibition to trigger these readouts. Based on this difference, we speculated that sphingolipid species per se are not acting as signaling molecules as we had envisioned but that chronic depletion of sphingolipids could be causing a plasma membrane stress that consequently triggers signaling events that impinge upon actin polarization and endocytosis. A prediction of this model is that manipulations that cause plasma membrane stress should also trigger depolarization of the actin cytoskeleton. Indeed, three orthogonal treatments, each targeting the tensile properties of the plasma membrane, similarly caused depolarization of the actin cytoskeleton with relatively rapid kinetics. How plasma membrane stress triggers actin depolarization remains to be determined.

**Fast Pathways—**There is substantial previously published genetic data suggesting that the CWI pathway functions downstream of TORC2 (20, 26, 56). Therefore, we were surprised to see that acute inhibition of TORC2 triggers hyperphosphorylation of Slt2 and thus presumably hyperactivation of the CWI pathway. This hyperactivation of the CWI pathway may be due to residual inhibition of TORC1 (TORC1-B) in our experimental setup (30, 69). Regardless, it is clear from these results that the CWI pathway is not inhibited upon BHS345 treatment and thus inactivation of this pathway cannot be what leads to actin depolarization and cessation of endocytosis.
TORC2 Regulates Actin and Endocytosis via Multiple Pathways

To better understand the mechanism(s) leading to actin depolarization and endocytosis arrest, we set out to identify the TORC2-dependent phosphoproteome (Tables 3 and 4). Our phosphoproteomic dataset demonstrates that TORC2, like TORC1, influences many aspects of cellular physiology. To our knowledge, this is the first time that a TORC2-specific phosphoproteome has been described. Future characterization of these TORC2 effectors promises to bring insight into the regulation of many important processes ranging from DNA repair and transcription to RNA and amino acid metabolism. Among the TORC2-regulated phosphoproteins were many components of the endocytosis machinery (Fig. 6B) whose phosphorylation status can be observed to change in less than 15 min following TORC2 inhibition. The critical targets in terms of linking TORC2 to actin polarization and endocytosis remain to be identified and characterized. We studied the regulation of the two proteins Pan1 and Ent1, and we confirmed a TORC2-dependent phosphorylation in low throughput assays. In addition, we showed that RX(S/T) sites in Ent1 are linked to the organization of actin. Future studies will be aimed at defining the nature of this link. Both Pan1 and Ent1/2 are essential for endocytosis and as such are prime candidates to serve as key regulatory nodes.

Finally, we also found that Fpk1 and Fpk2 are important players in the fast coupling of TORC2 signals to actin and endocytosis (Fig. 5). While this work was in progress, Niles and Powers (56), using an ATP analog-sensitive Ypk1, also reported that Fpks perform a major function in coupling Ypk1 signals to the actin cytoskeleton. This observation fits well with this work. However, deletion of FPK1/2 did not completely suppress actin depolymerization induced by TORC2 inhibition suggesting that additional effectors downstream of Ypk1 could be involved. Consistent with this notion is our observation that deletion of FPK1 and FPK2 does not prevent BHS345-induced Ent1 and Pan1 dephosphorylation.

What could be the role of Fpk1 and Fpk2 in actin polarization and endocytosis? It has been reported that the phospholipid translocases are the relevant targets of the Fpks in signaling to actin, a result that is seemingly discordant with our data. We do not presently understand how our two studies arrive at these different conclusions. Niles and Powers (56) subsequently present compelling data that reactive oxygen species generated downstream of hyperactive phospholipid translocases trigger actin depolarization. Future work will be needed to determine whether these reactive oxygen species are generated by flipase activity or other Fpk targets, as suggested by this work.

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Note Added in Proof—The images shown for mock-treated and BHS345-treated TOR1M2282T cells in Fig. 2B mistakenly duplicated the images of mock-treated and BHS345-treated TOR1M2282T cells containing an empty vector in Fig. 2F in the version of this article that was published on April 16, 2015, as a Paper in Press. The correct images of TOR1M2282T cells not expressing an empty vector are now shown. This mistake does not affect the interpretation of the results or the conclusions.

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