A Calcineurin-dependent Switch Controls the Trafficking Function of α-Arrestin Aly1/Art6

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Background: In response to nutrient signals, α-arrestins selectively regulate trafficking of membrane transporters.

Results: Aly1 is a substrate of the phosphatase calcineurin, and dephosphorylation triggers Aly1-dependent internalization of the permease Dip5.

Conclusion: Endocytic function of α-arrestins is stimulated by removal of inhibitory phosphorylation.

Significance: These insights define a molecular mechanism controlling the function of an α-arrestin in endocytosis, which is critical for cellular adaptation.

Proper regulation of plasma membrane protein endocytosis by external stimuli is required for cell growth and survival. In yeast, excess levels of certain nutrients induce endocytosis of the cognate permeases to prevent toxic accumulation of metabolites. The α-arrestins, a family of trafficking adaptors, stimulate ubiquitin-dependent and clathrin-mediated endocytosis by interacting with both a client permease and the ubiquitin ligase Rsp5. However, the molecular mechanisms that control α-arrestin function are not well understood. Here, we show that α-arrestin Aly1/Art6 is a phosphoprotein that specifically interacts with and is dephosphorylated by the Ca2+- and calmodulin-dependent phosphatase calcineurin/PP2B. Dephosphorylation of Aly1 by calcineurin at a subset of phospho-sites is required for Aly1-mediated trafficking of the aspartic acid and glutamic acid transporter Dip5 to the vacuole, but it does not alter Rsp5 binding, ubiquitinylation, or stability of Aly1. In addition, dephosphorylation of Aly1 by calcineurin does not regulate the ability of Aly1 to promote the intracellular sorting of the general amino acid permease Gap1. These results suggest that phosphorylation of Aly1 inhibits its vacuolar trafficking function and, conversely, that dephosphorylation of Aly1 by calcineurin serves as a regulatory switch to promote Aly1-mediated trafficking to the vacuole.

Cellular adaptation to environmental changes requires tight regulation of cell surface proteins. Specific ligands, excess nutrients, or stress factors induce endocytosis of plasma membrane receptors and permeases, thereby impacting nearly every aspect of cell physiology. Thus, a dynamic interplay exists between such extrinsic signals and endocytosis of specific membrane proteins, whose removal from the cell surface can alter intracellular signaling (1). One of the clearest examples of this interplay is the regulation of G-protein-coupled receptors (GPCRs) by the β-arrestin class of trafficking adaptors. In mammalian cells, agonist-stimulated GPCRs initiate intracellular signaling that leads to feedback phosphorylation of the receptor (2, 3) and, in several cases, dephosphorylation of β-arrestins (4–6). Dephosphorylated β-arrestins associate with the plasma membrane and bind both GPCRs and clathrin to stimulate GPCR endocytosis (4–10). GPCR removal from the plasma membrane dampens signaling. Although β-arrestin function is clearly regulated by phosphorylation, in many cases the kinases and phosphatases responsible for this regulation have not been identified.

A related protein family, the α-arrestins (also known as arrestin-related trafficking adaptors in Saccharomyces cerevisiae or arrestin domain-containing proteins (ArrDCs) in mammalian cells) are conserved across eukaryotes, with 14 members in yeast and at least 6 in mammals (11–14). Hallmarks of these proteins include N-terminal arrestin-fold domains and a C-terminal tail containing dispersed copies of a sequence motif, PXXY (and variants thereof), which are sites for interaction with a HECT domain ubiquitin ligase (Rsp5 in yeast and Nedd4 in mammals) (15, 16). The yeast α-arrestins regulate signal-induced endocytosis (12, 17–21) and intracellular sorting of nutrient permeases (22–25). Thus, α-arrestins are adaptors.

* This work was supported, in whole or in part, by National Institutes of Health Grants R01 GM-48728 (to M. S. C.), R01 GM-21841 (to J. T.), and R01 DA014204-11 (to A. Sorkin, University of Pittsburgh). This work was also supported by Agilent Technologies foundation grant (to A. F. O. and M. S. C.).

1 This article contains supplemental Tables 1 and 2.
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4 The abbreviations used are: GPCR, G-protein-coupled receptor; AzC, azetidine 2-carboxylic acid; CN, calcineurin/PP2B; Can, canavanine; Gal4-DBD, Gal4 DNA-binding domain; Gal4-TAD, Gal4 transcriptional activation domain; MIN, minimal medium; PM, plasma membrane; SC, synthetic complete medium; WCE, whole cell extract; ANOVA, analysis of variance; CHX, cycloheximide; G6PD, glucose-6-phosphate dehydrogenase; CIP, calf intestinal alkaline phosphatase.
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that link Rsp5 to specific membrane proteins and also directly interact with the trafficking machinery, such as clathrin adaptor complexes, to promote cargo trafficking (15, 24). In mammals, \( \alpha \)-arrestins associate with Nedd4, contribute to down-regulation of GPCR signaling, and impact the capacity for nutrient uptake (specifically glucose) (26–30). Recent reports also implicate ArrDC3 and Nedd4 in endocytosis of \( \beta 4 \) integrin in breast cancer cells and demonstrate that another mammalian ArrDC TXNIP (also known as VDUP1 for vitamin D3 up-regulated protein-1) promotes clathrin-mediated endocytosis of the glucose transporter GLUT1 (31, 32). Therefore, regulation of endocytosis is likely a conserved function of \( \alpha \)-arrestins.

Importantly, \( \alpha \)-arrestin-mediated trafficking is both signal- and cargo-selective. Some transporters seem to associate with a single dedicated \( \alpha \)-arrestin, whereas others can recruit multiple arrestins. Many \( \alpha \)-arrestins regulate several different transporters, and the \( \alpha \)-arrestin-cargo association can change depending on the signal inducing internalization (12, 20, 33). For example, in yeast \( \alpha \)-arrestin Ldb19/Art1 regulates endocytosis of the lysine permease Lyp1 in response to excess lysine, whereas \( \alpha \)-arrestin Ecm21/Art2 regulates Lyp1 internalization in response to general stress (20).

How do \( \alpha \)-arrestins achieve this signal-induced cargo selectivity? Recent reports suggest that phosphorylation blocks \( \alpha \)-arrestin-mediated endocytosis and, conversely, that dephosphorylation releases this inhibition (17, 34, 35). For example, Art1-mediated endocytosis of the arginine permease Can1 is impaired when Art1 is phosphorylated by Npr1 (34), a kinase that is activated during nitrogen starvation, a condition known to inhibit permease internalization (35–37). Similarly, in response to nitrogen starvation, Npr1-dependent phosphorylation of Bul1 and Bul2, recently identified as yeast \( \alpha \)-arrestins, impairs Bul1-mediated endocytosis of the general amino acid permease Gap1 (35). In addition, phosphorylation of \( \alpha \)-arrestin Rod1/Art4 by Snf1 (mammalian ortholog is the AMP- and ADP-activated protein kinase) blocks its ability to internalize the lactic acid permease Jen1, whereas dephosphorylation of Rod1 promotes endocytosis of Jen1 (17). In mammalian cells similar phospho-inhibition was recently demonstrated; phosphorylation of \( \alpha \)-arrestin TXNIP by AMP- and ADP-activated protein kinase induces \( \alpha \)-arrestin degradation, thereby impeding endocytic turnover of the glucose transporter GLUT1 (32). Hence, identification of the kinases and phosphatases that modify \( \alpha \)-arrestins provides important mechanistic insights into the regulation of \( \alpha \)-arrestin-mediated trafficking. Although dephosphorylation of at least some \( \alpha \)-arrestins appears to be required for their function in permease endocytosis, direct dephosphorylation by a specific phosphatase has not yet been shown for any \( \alpha \)-arrestin.

One signal-regulated phosphatase that is a good candidate for \( \alpha \)-arrestin regulation is calcineurin (also called phospho-protein phosphatase 2B or PP2B), a calcium and calmodulin-dependent phosphoprotein phosphatase conserved across eukaryotes and the target of the immunosuppressant drugs, FK506 and cyclosporin A. In mammals, calcineurin is abundant in the brain, where it regulates synaptic vesicle endocytosis, neural outgrowth, and synaptic plasticity; in other tissues, it impacts a wide range of processes, playing critical roles in T-cell differentiation and muscle development (38, 39). Although not essential in yeast under standard growth conditions, calcineurin is required for survival under various stressful conditions, such as in the presence of high salt, at alkaline pH, or in the presence of cell wall perturbants (40). Calcineurin promotes survival under stress conditions by binding to substrates that contain a conserved docking motif (PXIXIT and variants thereof) (39, 41) and dephosphorylating these targets to facilitate their function in stress adaptation. For example, dephosphorylation of the transcription factor Crz1 in response to cell wall stress induces expression of cell wall maintenance genes (42), whereas dephosphorylation of Hph1, an endoplasmic reticulum resident protein involved in protein translocation, assists in adaptation to alkaline pH (43, 44). In this regard, the first indication that calcineurin might also regulate endocytosis in yeast was the observation that calcineurin dephosphorylates two plasma membrane-localized phosphatidylinositol 4,5-bisphosphate-binding proteins (Slm1 and Slm2) to promote heat-induced internalization of the uracil permease Fur4 (45).

Here, we show that \( \alpha \)-arrestin Aly1/Art6 is a new substrate of calcineurin and demonstrate that dephosphorylation of Aly1 by calcineurin is required for Aly1-mediated internalization and delivery to the vacuole of a nutrient permease. Aly1/Art6 and its closely related paralog Aly2/Art3 regulate intracellular sorting of the general amino acid permease Gap1 in response to nitrogen supply (24). Aly2 also promotes endocytosis of the aspartic acid/glutamic acid permease Dip5, a function that Aly1 was suggested, but not previously shown, to share (18). We located 22 phospho-sites in Aly1, identified a subset of these as regulated by calcineurin, and delineated the specific PXIXIT-docking motif in Aly1 needed for its interaction with and dephosphorylation by calcineurin. We further show that Aly1 mutants that cannot be dephosphorylated by calcineurin or that mimic persistent phosphorylation are unable to reduce cell surface levels of Dip5. Thus, these studies identify a new role for calcineurin in membrane trafficking; calcineurin promotes vacuolar trafficking of Dip5 by dephosphorylating Aly1, thereby stimulating its endocytic function. By contrast, we found that dephosphorylation of Aly1 is not required for its role in the intracellular sorting of Gap1. Our data add to the growing body of evidence that \( \alpha \)-arrestin-mediated trafficking is strictly controlled by a phosphorylation-dependent switch wherein phosphorylation blocks and dephosphorylation promotes the function of an \( \alpha \)-arrestin in endocytosis. This study further identifies the first phosphatase responsible for direct regulation of an \( \alpha \)-arrestin.

**EXPERIMENTAL PROCEDURES**

**Yeast Strains and Growth Conditions**—Yeast strains used in this study and their construction are described in supplemental Table 1 (84, 85). Synthetic complete (SC) medium was prepared as in Refs. 24, 46 with 2.5 g/liter \((NH_4)_2SO_4\) used routinely as the nitrogen source. Minimal (MIN) medium was made in the same way as SC, except that only the amino acids required for growth of auxotrophic strains were provided. Cells were grown at 30 °C unless otherwise indicated. For growth assays on agar plates, 5-fold serial dilutions of stationary phase cultures with a starting density of \( \sim 1 \times 10^7 \) cells/ml were plated onto the indi-
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cated medium and grown for 3–6 days at 30 °C. Rapamycin (LC Laboratories, Woburn, MA), aminoglutethimide (A2-C), and canavanine (Can) (Sigma) were added to either SC or MIN + 0.5% (NH4)2SO4 at the concentrations indicated. Yeast two-hybrid analyses employed yeast strain PJ69-4a (47) containing pGBT9-derived plasmids bearing Gal4 DNA-binding domain (DBD) fusions and pACT2-derived plasmids bearing Gal4 transcriptional-activation domain (TAD) fusions. Transformants were plated as serial dilutions onto SC medium lacking leucine and tryptophan, as a positive control for cell growth, or additionally lacking histidine or adenine, where growth is a read-out of GAL1prom-HIS3 or GAL2prom-ADE2 reporter activation, respectively. A competitive inhibitor of His3, 3-aminotriazole (Sigma), was added where indicated to increase the amount of HIS3 reporter expression needed to allow growth. Yeast cells were transformed using the lithium-acetate method (48).

Plasmids and DNA Manipulations—Plasmids used in this study and their construction, where applicable, are described in supplemental Table 2 (86–90). Plasmids were generated using standard recombinant DNA methods (49) and propagated in Escherichia coli strain DH5α. Site-directed mutagenesis, used to generate Aly1 plasmids with altered CN-binding sites or mutated serines/threonines, was performed with either the QuikChange II site-directed mutagenesis kit (Agilent Technologies) for 30 min to inhibit CN or with 200 mM calcium chloride for 10 min to stimulate CN-mediated dephosphorylation. Cells were then lysed using sodium hydroxide, and proteins were precipitated using trichloroacetic acid (TCA) (51). Precipitated protein was solubilized by resuspension in SDS-/urea sample buffer (40 mM Tris (pH 6.8), 0.1 mM EDTA, 5% SDS, 8 M urea, and 1% β-mercaptoethanol) and heating to 37 °C for 15 min. An equal amount of extract was resolved by SDS-PAGE and specific proteins identified by immunoblotting (for a list of antibodies used see below). Immunoblots with anti-Pgk1, anti-Pma1, or anti-Zwf1 (glucose-6-phosphate dehydrogenase, hereafter referred to as G6PD) were detected using an Odyssey™ infrared imaging system (Li-Cor Biosciences) and quantified using ImageJ (National Institutes of Health), and the per-  

Yeast Protein Extraction, Purification, and Immunoblotting—
Yeast protein extracts were generated by growing cells to mid-exponential phase at 30 °C (A600 nm = 0.5–1.0) and harvesting an equal number of cells by centrifugation. In some cases, cell cultures were pretreated with either 2 µg/ml FK506 (LC Laboratories) for 30 min to inhibit CN or with 200 mM calcium chloride for 10 min to stimulate CN-mediated dephosphorylation. Cells were then lysed using sodium hydroxide, and proteins were precipitated using trichloroacetic acid (TCA) (51). Precipitated protein was solubilized by resuspension in SDS-/urea sample buffer (40 mM Tris (pH 6.8), 0.1 mM EDTA, 5% SDS, 8 M urea, and 1% β-mercaptoethanol) and heating to 37 °C for 15 min. An equal amount of extract was resolved by SDS-PAGE and specific proteins identified by immunoblotting (for a list of antibodies used see below). Immunoblots with anti-Pgk1, anti-Pma1, or anti-Zwf1 (glucose-6-phosphate dehydrogenase, hereafter referred to as G6PD) were used to assess protein loading.

To assess Dip5-GFP stability, MKY1800 (aly1Δ aly2Δ dip5-GFP) cells containing pRS315, pRS315-Aly1, pRS315-Aly1Δplkin, pRS315-Aly2, pRS315-Aly1ΔSE, or pRS315-Aly1ΔSA were grown to mid-exponential phase at 30 °C in MIN + 0.5% (NH4)2SO4 and treated with 200 µg/ml aspartic acid and glutamic acid to trigger Dip5 endocytosis. Cell samples were taken at times indicated post-aspartic/glutamic acid addition, and total protein was extracted using the sodium hydroxide lysis and TCA precipitation method described above. Dip5-GFP signal intensity, normalized for loading using G6PD, was quantified using ImageJ (National Institutes of Health), and the percentage of Dip5 remaining at each time point post-aspartic acid/glutamic acid addition was plotted. A representative data set from at least three independent experiments is shown.

Yeast extracts for pulldowns using GST fusion proteins were prepared by growing either strain BJ5459 or JR11 containing the pKK212-derived Aly1 or Aly2 expression plasmid indicated to mid-exponential phase in SC medium lacking tryptophan and inducing expression of CUP1 promoter-driven GST or GST-Aly fusions with 200 µM CuSO4 for 60 min. Where indicated, cells were also treated with either FK506 or CaCl2 as described above. Cells were harvested by centrifugation, washed, frozen in liquid N2, and stored at −80 °C. Cell pellets were then resuspended in co-IP buffer (50 mM Tris-HCl (pH 7.4), 15 mM EGTA, 100 mM NaCl, 0.2% Triton X-100, 5 mM N-ethylmaleimide, with phosphatase inhibitors (5 mM NaF, 5 mM Na3VO4, 5 mM EDTA, 5 mM EGTA, 1 mM sodium orthovanadate, 2.5 mM β-glycerophosphate, 10 mM sodium pyrophosphate, and 0.4 mM sodium metavanadate), and protease inhibitors (Complete protease inhibitor mixture tablets, Roche Applied Science) and lyzed at 4 °C using glass beads and vigorous vortexing. GST fusion proteins were purified from equal concentrations of clarified lysates by incubation with glutathione-Sepharose beads (GE Healthcare). Pulldowns were washed three times in 500 µl of co-IP buffer, aspirated to dryness, eluted in Laemmli buffer (52), resolved by SDS-PAGE, and assessed by immunoblotting. GST-Aly1 proteins treated with calf intestinal alkaline phosphatase (CIP) (New England Biolabs) were purified as described above, washed with CIP buffer, and incubated with 30 units of CIP for 30 min at 37 °C prior to elution in Laemmli buffer (52) and further analyses.

Where indicated, immunoblots were probed with the following: rabbit polyclonal anti-GST (Sigma); mouse monoclonal anti-GFP (Covance, Emeryville, CA); rabbit polyclonal anti-GFP (Molecular Probes, Carlsbad, CA); rabbit polyclonal anti-HA (Covance); mouse monoclonal anti-ubiquitin (gift from Richard Gardner, University of Washington, Seattle); rabbit polyclonal anti-Bmh (recognizes yeast 14-3-3 proteins Bmh1 and Bmh2; gift from Sandra Lemmon, University of Miami, FL); rabbit polyclonal anti-Rsp5 (gift from Linda Hicke, University of Texas, Austin); rabbit polyclonal anti-Pma1 (gift from Amy Chang, University of Michigan, Ann Arbor); rabbit polyclonal anti-Zwf1/G6PD (Sigma); and rabbit polyclonal anti-Pgk1 (53). Antibodies against Pma1, G6PD, or Pgk1 were examined to ensure equivalent loading of extracts. For all immunoblot data presented, anti-mouse and anti-rabbit secondary antibodies conjugated to IRDye-800 or IRDye-680 (Li-Cor Biosciences, Lincoln, NE) were detected using an Odyssey™ infrared imaging system (Li-Cor Biosciences). Quantification of immunoblots was performed using ImageJ 1.39u software (National Institutes of Health, Bethesda).

In Vitro Dephosphorylation Assays—GST-fused Aly1, Aly1Δplkin, Aly2, and Crz1 were purified on glutathione-Sepharose beads from BJ5459 cell extracts as described above. Under these conditions, GST-arpresins purify with several
interacting proteins (24), including protein kinases. To allow in vitro phosphorylation (and radiolabeling) of arrestins by these copurifying protein kinases, bead-bound proteins were washed and resuspended in “kinase” buffer (50 mM Tris-HCl (pH 7.5), 10 mM MgCl₂, 0.1 mM DTT, 0.1 mM unlabeled ATP, uponitin, and leupeptin) and incubated at 30 °C for 60 min in the presence of 75 nM [γ-32P]ATP (PerkinElmer Life Sciences). Unincorporated 32P and copurifying proteins were removed from the glutathione-immobilized arrestins by repeated washing of beads with co-IP buffer containing an additional 650 mM NaCl, 2 mM EDTA, and 0.8% Triton X-100. Bead-bound proteins were then resuspended in kinase buffer (listed above) with either λ-phosphatase (New England Biolabs) or recombinant CN-trunc, a mutant version of calcineurin that lacks its autoinhibitory domain (which is constitutively active in the absence of calcium and calmodulin), and incubated at 30 °C. Samples were removed at the times indicated, beads were aspirated to dryness, and bound proteins were eluted in Laemmli buffer (52) and resolved by SDS-PAGE. Gels were stained with Gel Code Blue (Thermo Scientific), dried, exposed to a PhosphorImager™ screen, and imaged with a Typhoon scanner (GE Healthcare). Image software was used to quantify changes in 32P signal intensity (normalized for protein loading using quantification of Gelcode Blue staining) after incubation with phosphatase. Data were quantified from three replicates (representative data show), and the mean percentage of the original phosphorylation signal is plotted ± S.E.

**Mass Spectroscopy Analysis**—GST-fused Aly1 or Aly1APILKIN was purified from BJ5459 cells that were treated with either CaCl₂ (to stimulate calcineurin-mediated dephosphorylation) or FK506 (to block calcineurin-mediated dephosphorylation), as described above. Samples were resolved by SDS-PAGE and stained with Gelcode Blue (Thermo Scientific), and the bands corresponding to Aly1 or Aly1APILKIN were excised. Proteins in each gel slice were then trypsinized and extracted as described in the “Enzymatic Digestion of Proteins from Gel Bands” protocol provided by the California Institute for Quantitative Biosciences. Mass spectrometry of extracted peptides was performed by the Vincent J. Coates Proteomics/ Mass Spectrometry Laboratory at University of California at Berkeley. A nano-LC column was packed in a 100-mum inner diameter glass capillary with an emitter tip. The column consisted of 10 cm of Polaris c18 5 μm packing material (Varian), followed by 4 cm of Partisphere 5 SCX (Whatman). The column was loaded by use of a pressure bomb and washed extensively with buffer A (see below). The column was then directly coupled to an electrospary ionization source mounted on a Thermo-Fisher LTQ XL linear ion trap mass spectrometer. Data collection was programmed so that neutral loss of phosphate would trigger the collection of an MS3 spectrum of the neutral loss peak. An Agilent 1200 HPLC equipped with a split line so as to deliver a flow rate of 30 n/min was used for chromatography. Peptides were eluted using a four-step MudPIT procedure (54). Buffer A was 5% acetonitrile, 0.02% heptfluorobutyric acid; buffer B was 80% acetonitrile, 0.02% heptfluorobutyric acid. Buffer C was 250 mM ammonium acetate, 5% acetonitrile, 0.02% heptfluorobutyric acid; buffer D was same as buffer C but with 500 mM ammonium acetate. The programs SEQUEST and DTASELECT were used to identify peptides and proteins from a database composed of the Aly1 sequence and common contaminant proteins. A minimum XCORR of 1.5, 2.2, and 3.5 was required for charge states 1, 2, and 3, respectively. To assess enrichment of phosphorylated peptides in FK506-treated versus Ca2+-treated samples, two-tailed Z-tests were performed, and p values are reported in Table 1.

**Uptake of 14C-Labeled Amino Acids**—Citrulline uptake assays were performed as described previously (24, 55). In brief, BY4741 cells were made prototrophic by transformation with pCK283 (56) and either pRS426, pRS426-Aly1, pRS426-Aly2, or pRS426-Aly1APILKIN. Cells were grown to early exponential phase (∼3–4 × 10⁸ cells/ml) in MIN + 0.5% (NH₄)₂SO₄ medium, collected, washed by filtration, and resuspended in nitrogen-free medium, and 20 μM [14C]citrulline (PerkinElmer Life Sciences) was added to cells to initiate uptake assays. Cell aliquots were removed every 30 s over four time points, collected, washed by filtration, and [14C]citrulline incorporation was measured using a Beckman Coulter LS6500 Multipurpose Scintillation Counter (Indianapolis, IN). Multiple replicate assays were performed (minimum of three), and the rate of citrulline uptake (dpm/min/μg protein) was determined. Aspartic acid uptake assays were performed as described for the citrulline uptake assays with the exception that 40 μM [14C]aspartic acid (PerkinElmer Life Sciences) was employed to initiate the uptake assays. In each case, plotted data represent the rate of amino acid uptake relative to the wild-type control.

**Fluorescence Microscopy**—Cell imaging of MKY1800 (aly1Δ aly2Δ Dip5-GFP) containing pRS315, pRS315-Aly1, pRS315-Aly1APILKIN, pRS315-Aly2, pRS315-Aly1SΔ, or pRS315-Aly1SΔA was performed at the Center for Biological Imaging (Pittsburgh, PA) using an Olympus IZB1 inverted microscope (Olympus America Inc., Center Valley, PA) with a 100 × 1.40 NA objective, and images were captured using a QImaging Retiga EXi CCD camera (QImaging, Surrey, British Columbia, Canada) and MetaMorph 7.5 Imaging System software (Molecular Devices, Sunnyvale, CA). Exposure times and microscope settings were kept constant during Dip5-GFP imaging. All Dip5-GFP images presented were processed equivalently using Adobe Photoshop software where intensity levels were adjusted and an unsharp mask filter applied (radius of 5 pixels). Pixel intensity for Dip5-GFP at the plasma membrane (PM) was measured by outlining PMs manually using a 3-pixel wide stroke with ImageJ software (National Institutes of Health) for a minimum of 140 cells per sample. Mean background fluorescence for each image was subtracted from the mean pixel intensity for each region of interest. The mean PM intensity (in arbitrary units) for each sample is plotted ± S.E., and statistical significance of changes in PM intensity was assessed using one-way ANOVA and Tukey’s post hoc test using Prism software (GraphPad Software Inc., San Diego). To assess the ratio of PM to vacuolar fluorescence, PM intensities were determined as defined above, and each cell was assigned a specific number identifier. Vacuolar regions of interest were assigned using the corresponding differential interference contrast images and

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then overlaid onto the fluorescent images. Only cells where clear vacuolar morphology was evident on the differential interference contrast image were used in subsequent analyses. A minimum of 40 cells was assessed this way per sample, and the mean PM/vacuolar fluorescence intensity is presented. Error bars represent S.E. and statistical significance of changes in PM/vacuolar fluorescence intensity ratio assessed using one-way ANOVA and Tukey’s post hoc test in Prism software.

RESULTS

C-terminal PXIXIT Motif in Aly1 Mediates Its Association with Calcineurin—Yeast two-hybrid analysis successfully identified several calcineurin substrates (43, 45, 57, 58). To identify additional potential substrates, we conducted a yeast two-hybrid screen using a Gal4-DBD fusion to Cna1, a catalytic subunit of calcineurin, as the bait. The plasmid that showed the strongest interaction with this bait encoded the C-terminal half (amino acids 493–915) of Aly1 fused in-frame to the Gal4-TAD (Fig. 1A). Further yeast two-hybrid analysis using the PJ69-4a reporter strain (47) revealed that full-length Aly1 interacts with yeast Cna1 (Fig. 1, A and B) and even with the catalytic subunit (CNA) of Homo sapiens calcineurin (Fig. 1B). Neither yeast Cna1 nor human CNA interacted with Aly2, the Aly1 paralog (39% identity; 64% similarity), or with any other yeast -arrestin (Fig. 1B), even though every -arrestin tested was expressed in the yeast two-hybrid strain (Fig. 1C)

The four serines and one threonine mutated to alanine to generate Aly1 or glutamic acid to generate Aly1 are underlined. * This denotes Ser-568, which was not identified as phosphorylated in our dataset but is close to other calcineurin-regulated sites and is phosphorylated in PhosphoPep and so was mutated in our analyses.
TAD-fusion) at a level equivalent to the wild-type Aly1 fusion (Fig. 1C).

To confirm that the interaction detected by the yeast two-hybrid method is physiologically relevant, we overexpressed either GST alone or fused to Aly1 or Aly1\textsuperscript{ΔPILKIN} from the copper-inducible promoter \textit{CUP1} in yeast cells coexpressing GFP-tagged Cna1 from its chromosomal locus. Reassuringly, Cna1-GFP strongly associated with Aly1 but did not associate with Aly1\textsuperscript{ΔPILKIN} at a level any higher than seen with the negative control (GST alone) (Fig. 1D), demonstrating that Aly1 interacts with calcineurin \textit{in vivo} and that this interaction requires the PILKIN motif near the C-terminal end of Aly1.

\textit{Aly1 Is a Substrate of Calcineurin}—Next, we tested whether Aly1 was dephosphorylated by calcineurin \textit{in vitro}. For many calcineurin substrates, loss of the PXIXIT-docking motif abrogates phosphatase binding and prevents dephosphorylation (43, 45, 59, 60). To prepare phosphorylated versions of Aly1, Aly1\textsuperscript{ΔPILKIN} (lacks the calcineurin-binding site), and Aly2 (does not bind calcineurin), we purified these proteins (as GST fusions) from yeast under low stringency conditions to retain associated protein kinases. Indeed, when these preparations were incubated with [\(γ^3P\)]ATP, radioactivity was readily incorporated into all three proteins but not into the GST alone control purified in the same fashion (data not shown). After stringent washing to remove the kinases and unincorporated [\(γ^3P\)]ATP, the radiolabeled proteins were then incubated with either purified calcineurin or λ-phosphatase (Fig. 2A, \(t = 0\) lanes). Recombinant Crz1, a known calcineurin substrate (60, 61), radiolabeled by phosphorylation with protein kinase Hrr25 (63), was used as a positive control. Upon incubation with calcineurin, both Aly1 and Crz1, but neither Aly1\textsuperscript{ΔPILKIN} nor Aly2, exhibited the increased electrophoretic mobility and decreased \(3^P\) label indicative of dephosphorylation (Fig. 2, A and B), whereas all four substrates were dephosphorylated by λ-phosphatase (Fig. 2A). Thus, Aly1, but not Aly2, is a direct substrate for calcineurin \textit{in vitro}, and the PILKIN sequence, which represents the PXIXIT motif in Aly1, is required for calcineurin to interact with and dephosphorylate Aly1.

\textit{Calcineurin Dephosphorylates a Specific Subset of Phosphosites in Aly1}—We noted that calcineurin-mediated dephosphorylation of Crz1, a substrate with a high affinity for calcineurin, was more rapid and complete than dephosphorylation of Aly1 (Fig. 2A; compare \(t = 30\) min for Aly1 and Crz1). Incubation of Crz1 with either calcineurin or λ-phosphatase resulted in a similar shift in electrophoretic mobility and equivalent loss of \(3^P\) signal. In contrast, Aly1 was not dephosphorylated as extensively by calcineurin as it was by λ-phosphatase (Fig. 2A), suggesting that only a subset of Aly1 phospho-sites are regulated by calcineurin.

To assess whether Aly1 is a calcineurin substrate \textit{in vivo}, we examined the electrophoretic mobility of Aly1 or Aly1\textsuperscript{ΔPILKIN} extracted from yeast treated with the following: (a) FK506, which inhibits calcineurin (64); (b) Ca\(^{2+}\), which stimulates calcineurin; or (c) a combination of FK506 followed by Ca\(^{2+}\), to control for any Ca\(^{2+}\)-stimulated calcineurin-independent effects. Similar analyses were conducted with Aly1\textsuperscript{PVIVIT}, a mutant in which PILKIN was converted to a known PXIXIT variant that displays high affinity calcineurin binding (59, 65), and with Aly1\textsuperscript{AAAAAA}, in which the PILKIN sequence was abrogated by six Ala substitution mutations. Upon calcineurin activation, only Aly1 and Aly1\textsuperscript{PVIVIT} migrated as a discrete doublet with the faster migrating band (below the 150-kDa marker) distinctly more prevalent than the slower migrating species (above the 150-kDa marker) (Fig. 2C). In contrast, Aly1\textsuperscript{ΔPILKIN} and Aly1\textsuperscript{AAAAAA} exhibited a diffuse banding pattern even when calcineurin was activated (Fig. 2C, compare Ca\(^{2+}\)-treated lanes). The same diffuse banding pattern was displayed by all versions of Aly1 when calcineurin was inhibited by treating cells with FK506 (Fig. 2C), consistent with a lack of dephosphorylation and a concomitant increase in phospho-Aly1 isoforms.

In agreement with this conclusion, a diffuse Aly1 migration pattern was also observed in cells where calcineurin activity was abolished by other means, specifically through loss of its regulatory subunit (in a \textit{cnb1}Δ mutant) or loss of both of its catalytic subunits (in \textit{cna1Δ cna2} double mutant) (Fig. 2D). Moreover, in contrast to the effect observed in wild-type cells, in the absence of functional calcineurin the addition of Ca\(^{2+}\) failed to generate a defined Aly1 doublet with a prominent faster migrating species (Fig. 2D, compare \textit{Ca-treated lanes}).

To confirm that the diffusely migrating species represent phosphorylated isoforms, purified GST-Aly1, Aly1\textsuperscript{ΔPILKIN}, Aly1\textsuperscript{PVIVIT}, and Aly1\textsuperscript{AAAAAA} were treated with CIP. In all cases, treatment with CIP collapsed Aly1 species into a crisp doublet in which the fastest migrating band was markedly more prominent (Fig. 2E), demonstrating that the diffuse migration of the Aly1 doublet is due to phosphorylation. As observed \textit{in vitro} (Fig. 2A), \textit{in vivo} activation of calcineurin did not sharpen the Aly1 doublet to the same extent as incubation with CIP (Fig. 2E, compare Ca\(^{2+}\)-treated Aly1 or Aly1\textsuperscript{PVIVIT} (+/+ CIP), further suggesting that calcineurin is responsible for dephosphorylation of only a subset of phospho-sites in Aly1.

\textit{Calcineurin-mediated Dephosphorylation Does Not Influence Aly1 Ubiquitinylination or Stability}—Because Aly1 was identified as an \textit{in vitro} substrate for the ubiquitin ligase Rsp5 (66), the GST-Aly1 species isolated from cell extracts was probed with an anti-GST antibody and an anti-ubiquitin antibody to allow simultaneous detection of ubiquitinylated and nonubiquitinylated forms on immunoblots. This analysis revealed that the slower migrating band in the Aly1 doublet is ubiquitinylated (band above 150-kDa marker; Aly1-Ub, Fig. 2, C–E). As noted above, both bands in the Aly1 doublet were sharpened upon addition of CIP (Fig. 2E), suggesting that both

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**FIGURE 1.** Aly1 interacts with the calcytic subunit of CN from both yeast and humans. A and B, serial dilutions of PJ69-4a containing the indicated Gal4-TAD and Gal4-DBD fusions were plated on SC medium lacking the amino acids indicated and grown for 4 days at 30°C. A, schematic depicting the region of Aly1 expressed as Gal4-TAD fusion is provided where the N-terminal arrestin-fold (N Arr), C-terminal arrestin-fold (C Arr), C-terminal tail (C Tail), and PXIXIT motif are shown in red, green, blue and yellow, respectively. B, red box helps to highlight the comparison between the full-length Aly1 and Aly1 missing the PXIXIT motif (PILKIN). C, expression of the indicated Gal4-TAD fusions in the two-hybrid reporter strain PJ69-4a was assessed by immunoblotting with anti-HA (Covance) after resolving TCA-extracted whole cell lysates (WCE) by SDS-PAGE. Red dots denote the full-length α-arrestin-Gal4-TAD fusion. White lines indicate removal of redundant replicate lanes. D, copurification of the calcytic subunit of CN, Cna1-GFP, with GST, GST-Aly1, or GST-Aly1\textsuperscript{ΔPILKIN} (expressed from pKK212-derived plasmids) from extracts of BJ5459 cells was assessed by immunoblotting. 1% of the whole cell lysates used as input for the pulldowns is shown.
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FIGURE 2. Aly1, but not Aly1\textsuperscript{ΔPILKIN}, is a CN substrate in vitro and in vivo. (A) GST-tagged Aly1, Aly1\textsuperscript{ΔPILKIN}, Aly1, and Crz1 were purified on glutathione-Sepharose beads yeast extracts, incubated with [γ\textsuperscript{32}P]ATP, and phosphorylated by copurifying kinases. Glutathione-bound proteins were washed to scumb further phosphorylation, incubated with CN-trunc or λ-phosphatase for the indicated times, and assessed by SDS-PAGE. Gels were stained for total protein or imaged to detect \textsuperscript{32}P. Representative data from one of four replicates are shown.


Aly1 and Aly1-Ub are phosphorylated. Importantly, there was no apparent defect in ubiquitylation of Aly1 when the calcineurin-binding site was mutated or in response to altered calcineurin activity (Fig. 2E).

It has been reported that the phosphorylation status of yeast α-arrestins Bul1, Bul2, and Rod1 impacts their ubiquitylation and/or interaction with Rsp5 (17, 35) and that phosphorylation regulates the stability of the mammalian α-arrestin, TXNIP (32). We therefore tested whether dephosphorylation of Aly1 by calcineurin did not alter Aly1 stability; in cells treated with Ca\textsuperscript{2+} to activate calcineurin and cycloheximide (CHX) to inhibit protein synthesis, the degradation profiles of Aly1 and Aly1\textsuperscript{ΔPILKIN} were indistinguishable (Fig. 3, A and B). Furthermore, neither ubiquitylation of Aly1, nor its interaction with Rsp5 appeared to be regulated by calcineurin. Aly1 and Aly1\textsuperscript{ΔPILKIN} were both ubiquitylated (Fig. 2E), and equivalent amounts of Rsp5 copurified with Aly1 and Aly1\textsuperscript{ΔPILKIN} from yeast extracts (Fig. 3C). As a control in these experiments, we included mutants (Aly1\textsuperscript{Y686G} or Aly2\textsuperscript{Y703G}) in which puta-
tive Rsp5-binding sites in Aly1 or Aly2 (66) were changed from a canonical PPXY motif to PPXG. Introducing these mutations into Aly1, Aly1^ΔPILKIN, or Aly2 reduced the association of each protein with Rsp5 to the background levels observed with the GST control (Fig. 3C). In addition, introducing this mutation into Aly1 resulted in loss of ubiquitinylation and an increased level of this protein (Fig. 3, C and D). However, in contrast to Aly1, Aly1^ΔPILKIN, and Aly2, whose overexpression confers...
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resistance to rapamycin, a TORC1 inhibitor that mimics nitrogen starvation (Fig. 3E) (24, 67), overexpression of PPXG mutant Aly alleles failed to improve growth under these conditions, suggesting that these mutations abrogate the function of these proteins in vivo (Fig. 3E). Thus, ubiquitinylation of Aly1 resulted in an observed, slower migrating form of the protein, and neither the ubiquitinylation, stability, nor ability of Aly1 to bind Rsp5 was altered by its calcineurin-mediated dephosphorylation.

Mapping of Calcineurin-regulated Phospho-sites in Aly1—To determine which residues of Aly1 are phosphorylated and which are susceptible to dephosphorylation by calcineurin, we performed LC-MS² analysis of purified GST-tagged Aly1 and Aly1ΔPILKIN extracted from cells in which calcineurin was activated (by treatment with Ca²⁺) or in which calcineurin was inhibited (by treatment with FK506). Peptides covering >50% of the protein were recovered (Fig. 4A), and a total of 22 phosphorylated residues in Aly1 were identified (Fig. 4B and Table 1), several of which were also earmarked as phospho-sites in various yeast global phosphoproteomic analyses (Table 1) (68, 69). Statistical analysis showed that of these 22 phospho-sites, two (Ser-252 and Ser-573) were significantly enriched in FK506-treated Aly1 (and also identified in FK506-treated Aly1ΔPILKIN), making them good candidate sites for regulation by calcineurin (Fig. 4B and Table 1). Two other sites (Thr-250 and Ser-569) near Ser-252 and Ser-573, respectively, occasionally showed a modest enrichment, and a fifth site, Ser-568, near Ser-573, although not detected in our work, was reported to be an in vivo phospho-site in PhosphoPep (66). All these sites (except Ser-568) fit an -(S/T)P- consensus, and all lie in predicted solvent-exposed regions. Thus, based on these analyses, we mutated five candidate sites (Thr-250, Ser-252, Ser-568, Ser-569, and Ser-573) to Ala (generating mutant Aly1Δ5A). The Aly1Δ5A mutant should mimic Aly1 that has been permanently dephosphorylated by calcineurin. We also generated Aly1Δ5E to mimic Aly1 persistently phosphorylated at these same sites.

Phenotypic Effects of Calcineurin Regulation of Aly1—We showed previously that Aly1 overexpression confers an increase in resistance to rapamycin (Fig. 3E) (24). Although Aly1 and all the Aly1 phospho-mutants tested elevated resistance to rapamycin (Fig. 4C), we noted that cells expressing the Aly1 variants capable of being dephosphorylated by calcineurin (Aly1) or that mimic the dephosphorylated state (Aly1Δ5A) grew slightly better on rapamycin than cells expressing Aly1 variants that cannot be dephosphorylated by calcineurin (Aly1ΔPILKIN) or that mimic the persistently phosphorylated state (Aly1Δ5E) (Fig. 4C). This modest difference in phenotype was not due to differences in protein level as all of these GST fusions were expressed equivalently to wild-type Aly1 (Figs. 3C, 4D, and 7A) and behaved very similarly to wild-type Aly1 in other assays (Fig. 5, A and E, and data not shown).

In response to activation or inhibition of calcineurin activity, the electrophoretic migrations of Aly1Δ5A and Aly1Δ5E were indistinguishable from that of wild-type Aly1 (Fig. 4D), suggesting that additional calcineurin-regulated phospho-sites exist in Aly1 that were not pinpointed by our MS analyses (Fig. 4, A and B). However, further phenotypic characterization of Aly1Δ5A and Aly1Δ5E indicated (see last section under “Results”) that the sites altered in these mutants are functionally important (Figs. 4C and 6, B–G).

Regulation by Calcineurin Is Not Required for Aly1-mediated Gap1 Trafficking—Phosphorylation of α-arrestins regulates arrestin-mediated trafficking of nutrient permeases (17, 24, 34, 35). Therefore, we examined whether dephosphorylation of Aly1 by calcineurin modulates either of the two trafficking functions described for Aly1 to date as follows: 1) regulation of Gap1 (general amino acid permease) intracellular sorting and 2) regulation of Dip5 (aspartic/glutamic acid permease) endocytosis (18, 24).

When nitrogen is limiting, Gap1 mediates entry of a wide range of amino acids, including proline (70). Uptake of two amino acid analogs, AzC, a toxic proline mimetic, and citrulline, an arginine mimetic taken up exclusively through Gap1, serve as readouts for Gap1 activity at the plasma membrane (55, 70). We showed previously that overexpression of either Aly1 or Aly2 results in hypersensitivity to AzC and an increase in the rate of citrulline uptake (24). Increased Gap1 activity at the cell surface is due to increased Gap1 retrieval from endosomes (likely via an endosome-to-Golgi route); impeding Gap1 trafficking to the vacuole (a degradative organelle) increases Gap1 levels in the cell and leads, ultimately, to an increased level of Gap1 at the plasma membrane (24). Thus, we tested whether the phosphorylation status of Aly1 influences Gap1 trafficking. We found that overexpression of Aly1, Aly1ΔPILKIN, Aly1Δ5A, or Aly1Δ5E each conferred an equivalent increase in sensitivity to AzC (Fig. 5A) and Gap1 protein levels (Fig. 5C). In addition, overexpression of either Aly1 or Aly1ΔPILKIN caused a similar increase in the rate of citrulline uptake (Fig. 5D). To ensure that overexpression of these ALY1 alleles was not masking subtle phenotypic differences between them, we examined the ability of Aly1 phospho-mutants expressed from low copy plasmids to complement the aly1Δ aly2Δ AzC resistance phenotype, and we found that all alleles restored AzC sensitivity equally (Fig. 5B).

Our earlier studies demonstrated that Aly2, but not Aly1, requires a number of factors to promote Gap1 recycling, including the following: Npr1, a nitrogen-regulated protein kinase that promotes Gap1 trafficking to the plasma membrane (35–37); Lst4, a factor known to stimulate nutrient-dependent recycling of Gap1 (22, 55), and AP-1, an adaptor complex that recruits clathrin to endosome-derived vesicles trafficking to the Golgi (24, 71, 72). Deletion of any of these factors cripples Aly2-stimulated Gap1 recycling and may thereby create a sensitized assay to detect subtle defects in Aly1-mediated Gap1 trafficking that might arise from loss of calcineurin regulation (24). By every metric employed (rate of citrulline uptake, growth using citrulline as sole nitrogen source, and sensitivity to AzC), overexpression of Aly1, Aly1ΔPILKIN, Aly1Δ5A, and Aly1Δ5E caused similar increases in Gap1 activity at the plasma membrane irrespective of the genetic background used (npr1Δ, lst4Δ, apl2Δ) (Fig. 5, E and F, and data not shown). Together, these findings indicate that dephosphorylation of Aly1 by calcineurin does not regulate Aly1-mediated Gap1 recycling.

Dephosphorylation of Aly1 by Calcineurin Is Required for Aly1-mediated Endocytosis of Dip5—A previous study by another group demonstrated that Aly2 promotes endocytosis...
Calcineurin Regulates α-Arrestins

A Aly1 peptide coverage in MS analysis

MLQFNTENDTVAPVPEMQDINAAADAVPVQLVTTTLQFVKLAEPVFLLKFGFTNGLSEIAPSILGGSLIRVLPKPKL1KSISITFTHKSRTFWPEGPPKREFsDVTWVNH7WPYQADDGMNFTLHHSNNSNNRPMS5DDEYLEKSSAVYIPPITAEPFPKDNSLSLDAERNSLSSDNKSNPSSVDVHDDSKLAIKTPLSSRPGSVFANFSGNSLSPHTFISDLFTKTFNSGATPSQEDNLYLPSDSKEVFIYFRGDFYYSFQPIQSYEPASIKANFGSV

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by calcineurin might alter its ability to stimulate Dip5 trafficking to the vacuole. We found that the absence of either Aly1 or Aly2 caused a detectable increase in the steady-state level of Dip5, and an even more pronounced increase was observed in cells lacking both Aly1 and Aly2 (Fig. 6A), as was observed previously (18). This is consistent with the idea that each of these

FIGURE 4. Identification of a subset of Aly1 phospho-sites regulated by calcineurin. A, map of Aly1 peptides sequences identified in the MS analysis. The single letter code for amino acids encoding Aly1 is shown with regions of the protein for which peptides were identified in each of the four MS analyses performed shown in boldface green type. The PILKIN calcineurin-docking motif is shown in red; no peptides containing the PILKIN sequence were identified in Aly1 alone, as was observed previously (18). This is consistent with the idea that each of these

of Dip5 in response to excess amounts of the amino acids transported by this permease (aspartic acid or glutamic acid), but it did not address directly whether Aly1 may also play a role in this process (18). Therefore, we tested whether regulation of Aly1 by calcineurin might alter its ability to stimulate Dip5 trafficking to the vacuole. We found that the absence of either Aly1 or Aly2 caused a detectable increase in the steady-state level of Dip5, and an even more pronounced increase was observed in cells lacking both Aly1 and Aly2 (Fig. 6A), as was observed previously (18). This is consistent with the idea that each of these

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Calcineurin Regulates α-Arrestins

**A**

Vector | MIN | + 50 μg/ml AzC
--- | --- | ---
WT | | |
Aly1 | | |
Aly1::PILKIN | | |
Aly2 | | |
Aly1SA | | |
Aly1SE

**B**

Vector | MIN | + 100 μg/ml AzC
--- | --- | ---
aaly1Δ, aly2Δ | | |
Aly1::PILKIN | | |
Aly2 | | |
Aly1SA | | |
Aly1SE

**C**

Vector | Gap1-Arrestins | a-GFP | a-G6PD
--- | --- | --- | ---
WT | | | |
Aly1 | | | |
Aly1::PILKIN | | | |
Aly1SA | | | |
Aly1SE

**D**

Citricline uptake (% of WT)

**E**

Vector | npr1Δ | MIN | + 50 μg/ml AzC
--- | --- | --- | ---
WT | | | |
Aly1 | | | |
Aly1::PILKIN | | | |
Aly2 | | | |
Aly1SA | | | |
Aly1SE

α-arrestins contributes to the endocytosis of Dip5. Therefore, as an independent means to assess the impact of Aly2 and Aly1 variants on Dip5 activity and trafficking, all subsequent analyses were performed in an aly1Δ aly2Δ background containing either a vector control or a low copy plasmid expressing the ALY allele of interest from its endogenous promoter. We measured Dip5 function at the cell surface by monitoring the rate of radiolabeled aspartic acid uptake and found that expression of Aly1, Aly1SA, or Aly2 each significantly reduced the rate of aspartic acid uptake compared with the vector control (Fig. 6B), consistent with diminished Dip5 level and/or activity at the PM in these cells. By contrast, significantly higher Dip5 activity and/or levels were observed in cells expressing Aly1::PILKIN or Aly1SA, which maintains Aly1 in a hyperphosphorylated or phospho-mimetic state, compared with cells expressing either Aly1 or Aly1SA (Fig. 6B), their dephosphorylated counterparts. These data support a role for Aly1 and Aly2 in basal turnover of Dip5 as the aspartic acid uptake assays are done over a very short time course (2 min) and so likely reflect differences in the steady-state levels of Dip5 operating at the plasma membrane.

To determine whether the observed effects were due to alterations in the level of Dip5 at the cell surface, steady-state localization of Dip5-GFP was examined in cells expressing various ALY alleles. For these experiments, cells containing Dip5-GFP integrated at its chromosomal locus were grown under conditions equivalent to those used for the aspartic acid uptake assays. Images (Fig. 6C) were quantified, and reported differences in Dip5-GFP fluorescence intensity were assessed statistically, both as the pixel count at the PM (Fig. 6D) and as the ratio of the pixel count at the PM to that in the vacuole of the same cell (Fig. 6E; see also “Experimental Procedures”). Despite some cell-to-cell variation in these cell populations, these analyses showed that expression of Aly2 (as a positive control), as well as Aly1 and Aly1SA, significantly reduced the amount of Dip5 at the PM and increased the amount of Dip5 in the vacuole, compared with the aly1Δ aly2Δ controls cells carrying the vector alone (Fig. 6, C–E). In marked contrast, expression of Aly1::PILKIN or Aly1SE in the aly1Δ aly2Δ cells neither reduced the amount of Dip5 at the PM nor increased the amount of Dip5 in the vacuole (Fig. 6, C–E). It should be noted, however, that due to background cytoplasmic fluorescence the PM-to-vacuolar ratios presented in Fig. 6E are lower than expected for the vector control (and likely for the Aly1::PILKIN and Aly1SE alleles as well) based on the micrographs presented in Fig. 6C. These findings are consistent with the interpretation that phosphorylation blocks the ability of Aly1 to stimulate Dip5 internalization because Aly1 (which can be dephosphorylated by CN) and the Aly1SA variant (which mimics its dephosphorylated state)

pRS426-Aly1::PILKIN, pRS426-Aly1SA, pRS426-Aly1SE, or pRS426-Aly2) and pCK8230 (Gap1-GFP) were grown in MIN 0.5% (NH₄)₂SO₄. WCEs were resolved using SDS-PAGE, and Gap1 levels were assessed by immunoblotting. Gap1 levels relative to the vector control extract are presented below the immunoblot. D and F, prototrophic BY4741 (WT) or npr1Δ cells with pCK283 and pRS426-Aly1, pRS426-Aly1::PILKIN, or pRS426-Aly2 were assayed for [³⁵S]citricline uptake. The mean uptake rate for seven replicates (in C) or three replicates (in E) is shown as a percentage relative to the WT containing vector, and error bars represent ± S.D. E, growth of serial dilutions of npr1Δ cells with pRS425, pRS425-Aly1, pRS425-Aly1::PILKIN, pRS425-Aly2, pRS425-Aly1SA, or pRS425-Aly1SE on MIN 0.5% (NH₄)₂SO₄ ± AzC.

**FIGURE 5. CN regulation is not required for Aly1-mediated Gap1 recycling.** A, growth of serial dilutions of BY4741 cells containing pRS426 (vector), pRS426-Ally1, pRS426-Ally1::PILKIN, pRS426-Ally2, pRS426-Ally1SA, or pRS426-Ally1SE on MIN 0.5% (NH₄)₂SO₄ ± AzC. B, growth of serial dilutions of aly1Δ aly2Δ (D2–6A) cells containing pRS315 (vector), pRS315-Ally1, pRS315-Ally1::PILKIN, pRS315-Ally2, pRS315-Ally1SA, or pRS315-Ally1SE on MIN 0.5% (NH₄)₂SO₄ ± AzC. C, BY4743 cells containing pRS426 (vector, pRS426-Ally1 or pRS426-Ally2) and pCK8230 (Gap1-GFP) were grown in MIN 0.5% (NH₄)₂SO₄. WCEs were resolved using SDS-PAGE, and Gap1 levels were assessed by immunoblotting. Gap1 levels relative to the vector control extract are presented below the immunoblot. D and F, prototrophic BY4741 (WT) or npr1Δ cells with pCK283 and pRS426-Ally1, pRS426-Ally1::PILKIN, or pRS426-Ally2 were assayed for [³⁵S]citricline uptake. The mean uptake rate for seven replicates (in C) or three replicates (in E) is shown as a percentage relative to the WT containing vector, and error bars represent ± S.D. E, growth of serial dilutions of npr1Δ cells with pRS425, pRS425-Ally1, pRS425-Ally1::PILKIN, pRS425-Ally2, pRS425-Ally1SA, or pRS425-Ally1SE on MIN 0.5% (NH₄)₂SO₄ ± AzC.
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both promoted basal endocytosis of Dip5 in aly1Δ aly2Δ cells, whereas the Aly1Δ PILKIN variant (which cannot bind CN) and the Aly1SE phospho-mimetic mutant were unable to promote Dip5 endocytosis. Because these assays were done in the absence of substrate (no aspartic acid/glutamic acid ligand), the differences in Dip5 cell surface levels represent changes in the basal trafficking of Dip5 to the vacuole in the presence of these ALY alleles.

We also carried out kinetic analysis of Dip5 stability in response to exogenous aspartic acid and glutamic acid, which induces endocytosis of the Dip5 permease. After addition of aspartic acid and glutamic acid, degradation of Dip5-GFP was appreciably faster in cells containing Aly2, Aly1, or Aly1SE than in cells expressing Aly1Δ PILKIN, Aly1SE, or the control vector (Fig. 6, F and G). Moreover, even at the zero time point, cells expressing Aly2, Aly1, and Aly1SE contained less Dip5 than cells containing Aly1Δ PILKIN, Aly1SE, or a control vector (Fig. 6F), consistent with faster turnover of Dip5 when Aly2 or dephosphorylated Aly1 are present. Together, this data demonstrate that the phospho-mimetic Aly1SE protein and Aly1Δ PILKIN, which cannot be dephosphorylated by calcineurin, fail to internalize and degrade Dip5 as effectively as wild-type Aly1 or the dephosphorylation-mimetic Aly1SE. Thus, we conclude that dephosphorylation of Aly1 by calcineurin is required to promote Aly1-mediated trafficking of Dip5 to the vacuole.

Our data for Aly1 are consistent with recent findings that α-arestins Rod1 and Bul1 must be dephosphorylated to promote nutrient-induced endocytosis of the Jen1 and Gap1 permeases, respectively. Dephosphorylation of Rod1 and Bul1 by unidentified phosphatases results in loss of binding to the yeast 14-3-3 proteins that may be responsible for inhibiting their endocytic function (17, 35). Because previous MS analyses revealed enrichment of Bmh1 and Bmh2 peptides in purified preparations of Aly1 and Aly2,5 we examined whether calcineurin-mediated dephosphorylation of Aly1 alters its interaction with Bmh1 and Bmh2. We found that Bmh1 and Bmh2 copurified from yeast extracts with GST-Aly1 and GST-Aly2 but not with the GST control (Fig. 7), confirming our previous MS analyses. Although significantly higher levels of Bmh1 and Bmh2 copurified with Aly2 than with Aly1, the amount of Bmh1 and Bmh2 associated with Aly1, Aly1Δ PILKIN,

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**FIGURE 6.** Calcineurin-mediated dephosphorylation of Aly1 is required for Aly1-dependent trafficking of Dip5 to the vacuole. A, WCEs from BY4741, aly1Δ, aly2Δ, or aly1Δ aly2Δ cells containing a chromosomally integrated Dip5-GFP grown in MIN + 0.5% (NH4)2SO4 were resolved by SDS-PAGE, and Dip5 bands were quantified by densitometry and normalized for alterations in the G6PD loading control. Dip5-GFP band intensities from three replicate experiments (representative panel in Fig. 7) were measured, normalized to the G6PD loading control, and the mean percentage of Dip5 remaining post Asp/Glu addition (%) was plotted. S.E. is plotted. B, Dip5-GFP band intensities from three replicate experiments (representative panel in Fig. 7) were measured, normalized to the G6PD loading control, and the mean percentage of Dip5 remaining post Asp/Glu addition (%) was plotted. DIC, differential interference contrast.

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**FIGURE 7.** CN-regulation of Aly1 does not alter Aly1-association with 14-3-3 proteins. GST, GST-Aly1, GST-Aly1Δ PILKIN GST-Aly1SE, GST-Aly2, GST-Aly2 (expressed from pKK212-derived plasmids) were extracted and purified from BJ5459 cells using glutathione-Sepharose. Copurification of endogenous 14-3-3 proteins (Bmh1 is lower band and Bmh2 is upper band in doublet) was assessed by immunoblotting. 2% of the WCE used as input for pulldowns is shown.
Aly1G5A, or Aly1G5E was not detectably different (Fig. 7). These findings suggest that phosphorylation of Aly1 at the CN-sensitive sites delineated in this study is not the major determinant of the association with Bmh1 and Bmh2. Indeed, there are two sites in Aly1 that fit the predicted 14-3-3-binding sites consensus, each of which lies in peptides not recovered in our MS analyses and neither of which map to any calcineurin-regulated phospho-site identified here (Fig. 4A). Thus, although dephosphorylation of other α-arrestins may serve as a regulatory mechanism to release these adaptors from the grasp of 14-3-3 proteins, and thereby promote α-arrestin-mediated endocytosis of permeases, that does not appear to be the underlying mechanism by which CN-mediated dephosphorylation stimulates the trafficking function of Aly1.

**DISCUSSION**

**Phospho-regulation of Aly1-mediated Trafficking**—We show here that α-arrestin Aly1 interacts with calcineurin through a C-terminal docking motif and identified up to five possible CN-targeted phospho-sites in Aly1. Furthermore, by monitoring the amino acid permease Dip5 as its cargo, we demonstrated that Aly1 must be dephosphorylated by calcineurin to mediate Dip5 trafficking to the vacuole (Fig. 8). This study is the first to identify a phosphatase that directly regulates α-arrestin function. Although phospho-sites have been identified for various α-arrestins in global proteomic screens (68, 69), aside from our work on Aly1, a comprehensive phospho-map exists for only one other α-arrestin, namely Ldb19/Art1 (34). Given that only a handful of the 22 phospho-residues detected in Aly1 appear to be regulated by calcineurin (and there appear to be additional calcineurin-regulated sites in Aly1 not identified in our study), Aly1 action may be under the control of multiple protein kinases and phosphatases. Indeed, other α-arrestins are highly modified, many with >10 phospho-sites identified to date (68, 69), underscoring the potential complexity of their regulation.

Both Aly1 and the related Aly2 promote retrieval of the general amino acid permease Gap1 from endosomes, and Npr1-dependent phosphorylation of Aly2 is needed for its function in this Gap1 recycling (24). We show here that dephosphorylation of Aly1 by calcineurin is not required for such Gap1 recycling, suggesting that different subsets of phospho-sites may regulate distinct α-arrestin trafficking functions. It was demonstrated that Aly2 is necessary for efficient endocytosis of the Dip5 transporter (18), but whether Aly1 might function semi-redundantly in this process was never directly tested. We show here that Aly1, when dephosphorylated by calcineurin, does stimulate Dip5 internalization and turnover, providing evidence that Aly1, like other α-arrestins, mediates the endocytosis of nutrient transporters.

Thus, our results indicate that Aly1 action is under control of a phosphorylation-dephosphorylation switch; calcineurin-mediated dephosphorylation stimulates Aly1 function in trafficking at least one membrane permease to the vacuole (Fig. 8), and phosphorylation, by an as yet unidentified protein kinase, inhibits it. Because excess aspartate or glutamate stimulates Dip5 internalization, and Aly1 can only mediate Dip5 down-regulation when dephosphorylated by calcineurin, influx of these acidic amino acids may stimulate calcineurin-dependent signaling through unknown mechanisms. In turn, dephosphorylated Aly1 promotes Dip5 endocytosis to the vacuole (Fig. 8), perhaps to prevent the cytotoxicity that an excess of these amino acids can cause (73). Indeed, in mammalian cells, amino acid overload increases the level of Ca2+-calmodulin (74), a requirement for calcineurin activation.

**Arrestin Dephosphorylation Is a Conserved Mechanism That Promotes Arrestin-mediated Endocytosis**—Our data and other recent findings suggest a general model for phospho-regulation of α-arrestins (Fig. 8) (75). As shown here for Aly1/Art6, phosphorylation of Rod1/Art4 (17), Ldb19/Art1 (34), and two newly recognized α-arrestins Bull1 and Bull2 (35) blocks their ability to mediate endocytosis of specific nutrient permeases. In cells grown on lactate, the protein kinase Snf1 phosphorylates Rod1 to retain the lactic acid permease Jen1 at the plasma membrane; however, upon glucose addition, which inactivates Snf1, endocytosis of Jen1 ensues (17). Rod1 dephosphorylation appears to be PP1-dependent, but direct dephosphorylation of Rod1 by PP1 has not been demonstrated. Moreover, because PP1 action (Glc7-Reg1 complex) is necessary for inactivation of Snf1, the lack of apparent Rod1 dephosphorylation in reg1Δ cells may arise indirectly from lack of Snf1 down-regulation. For Ldb19, phosphorylation by the protein kinase Npr1 impairs Ldb19-mediated endocytosis of the arginine permease Can1, whereas the dephosphorylated protein associates with the plasma membrane and promotes Can1 endocytosis (34). However, the phosphatase responsible for Ldb19 dephosphorylation is not known. For Bul1 and Bul2, phosphorylation by Npr1, which is activated on poor nitrogen sources (e.g. proline), inhibits Bul-mediated endocytosis of Gap1. When NH4+ is the nitrogen source, Npr1 is inactive, and the Bul proteins are dephosphorylated in a man-
ner that depends on the phosphatase Sit4, promoting Gap1 internalization (35). However, direct dephosphorylation of Bul1 or Bul2 by Sit4 has not been demonstrated.

Thus, in the case of Aly1 and three other yeast α-arrestins, phosphorylation blocks and dephosphorylation promotes their endocytic function. Indeed, this phospho-inhibition of α-arrestin-mediated endocytosis is conserved in mammals. For example, phosphorylation of α-arrestin TNXIP by AMPK results in reduced stability of this α-arrestin, diminishing endocytic turnover of the GLUT1 glucose transporter regulated by TXNIP (32). However, the evidence to date indicates that phosphorylation can modulate α-arrestin function through distinct mechanisms. First, in the case of yeast Rod1, dephosphorylation of the α-arrestin promotes its endocytic function and is required for its ubiquitinylation (17). In contrast, dephosphorylation does not seem to regulate the ubiquitinylation of either Aly1, as we have shown here, or Bul1 (35). Similarly, under conditions that do not promote Ldb19-mediated internalization of Can1, this α-arrestin is ubiquitinylated, further supporting the idea that α-arrestin ubiquitinylation and its endocytic function need not be coupled (20, 34). Second, phosphorylation reportedly regulates the interactions of Rod1, Bul1, and Bul2 with cytoplasmic 14-3-3 proteins, resulting in sequestration of the α-arrestin (or the α-arrestin-Rsp5 complex) in the cytosol, and thereby inhibiting α-arrestin activity at and/or recruitment to the plasma membrane (17, 32, 35). Other α-arrestins are among the 271 proteins demonstrated to associate with Bmh1 and Bmh2 in a phosphorylation-dependent manner (76); therefore, interaction with 14-3-3 proteins may broadly regulate α-arrestin function. However, neither of these mechanisms is adequate to explain phospho-regulation of Aly1, because we demonstrated here that the calcineurin-dependent dephosphorylation of Aly1 does not alter its ubiquitinylation, stability, or interaction with either Rsp5 or 14-3-3 proteins. Dephosphorylation of mammalian β-arrestins is required for their association with clathrin and AP-2 and subsequent stimulation of endocytosis (6, 10). Both Aly1 and Aly2 have been shown to interact with clathrin adaptors (24); therefore, phosphorylation might similarly regulate the endocytic function of Aly1 by altering its interaction with these components of the trafficking machinery.

Conserved Interaction of Calcineurin with α-Arrestins and Its Implications—A Caenorhabditis elegans α-arrestin, CNP-1/ArrD-17, interacts with the catalytic subunit of calcineurin and is a calcineurin substrate in vitro (57). Although no calcineurin-docking site was identified in the nematode protein, we note that CNP-1/ArrD-17 contains near its C terminus a variant PIIXIT motif (PIVIGS) that is conserved in its mammalian α-arrestin relatives ARRDC2–4 and TNXIP (as PIVIGS or PLVIGT). Interestingly, calcineurin regulation of ArrD-17 in C. elegans also impacts response to starvation, although a link between specific α-arrestins and trafficking of any nutrient transporter has not yet been established.

To our knowledge, our work is the first demonstration that calcineurin dephosphorylates an α-arrestin to regulate the trafficking function of this class of adaptor proteins. However, calcineurin action influences membrane protein trafficking in other ways. For example, heat stress-induced calcineurin-mediated dephosphorylation of the TORC2- and eisosome-associated components Sm1 and Sm2 stimulates internalization of the uracil permease Fur4 (45, 62, 77). Studies of synaptic vesicle retrieval in mammalian cells indicate that calcineurin dephosphorylates a suite of endocytic regulators, including amphiphysins, syntaptojins, epsins, and dynamin (78, 79). Calcineurin activation also stimulates receptor-mediated endocytosis of AMPA and transferrin (80, 81). Collectively, and combined with our demonstration here that calcineurin regulates α-arrestin-mediated trafficking to the vacuole in yeast, it is clear that calcineurin has an important physiological function as a global regulator of membrane protein trafficking.

Acknowledgments—We gratefully acknowledge Marko Kaksonen (EMBL, Heidelberg, Germany) for providing the MKY yeast strains prior to their publication; Sandra Lemmon (University of Miami, FL) for providing the anti-Bmh antibody; and colleagues Alexander Sorlin, Jeffrey Brodsky, and Adam Kwiatkowski (University of Pittsburgh) for use of reagents and equipment. We thank Catherine Baty and Jenny Karlsson at the Center for Biological Imaging for assistance with cell imaging. We thank Lori Kohlstaedt (supported by National Institutes of Health Grant S10 RR025622-01) and the Vincent J. Coates Proteomics/Mass Spectrometry Laboratory at the University of California, Berkeley, for generating and analyzing the MS data.

REFERENCES

Calcineurin Regulates α-Arrestins

14. Calcineurin Regulates α-Arrestins


Calcineurin Regulates \( \alpha \)-Arrestins

a novel substrate of calcineurin, is critical for modulation of egg-laying and locomotion in response to food and insulin sensation in \textit{Caenorhabditis elegans}. \textit{J. Mol. Biol.} \textbf{417}, 165–178


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## Table 2. Plasmids used in this study

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<td>$GAP1\text{prom-}GAP1\text{-}sGFP$, CEN, URA3</td>
<td>(22)</td>
</tr>
<tr>
<td>pACT1-GEV</td>
<td>$ACT1\text{prom- Estradiol receptor-}GAL4$ in pRS40NAT</td>
<td>(87)</td>
</tr>
<tr>
<td>pACT2</td>
<td>$ADH1\text{prom-}GAL4\text{-TAD}$, 2µ, LEU2</td>
<td>(88)</td>
</tr>
<tr>
<td>Aly1-TAD (1-915)</td>
<td>$ADH1\text{prom-}GAL4\text{-TAD-}ALY1$, 2µ, LEU2</td>
<td>$ALY1$ coding sequence was PCR amplified from pKK212-$ALY1$ with primers containing $Xmal$ and $Xhol$ restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Aly1-TAD (1-609)</td>
<td>$ADH1\text{prom-}GAL4\text{-TAD-}ALY1(1-609)$, 2µ, LEU2</td>
<td>$ALY1$ sequence encoding amino acids 1-609 was PCR amplified from pKK212-$ALY1$ with primers containing $Xmal$ and $Xhol$ restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Plasmid</td>
<td>Genotype</td>
<td>Description (Reference)</td>
</tr>
<tr>
<td>-------------</td>
<td>--------------------------------</td>
<td>-----------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Aly1-TAD</td>
<td>(ADH1prom-GAL4-TAD-ALY1(610-915), 2\mu, LEU2)</td>
<td>(ALY1) sequence encoding amino acids 610-915 was PCR amplified from pKK212-(ALY1) with primers containing (XmaI) and (XhoI) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>(610-915)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aly1-TAD</td>
<td>(ADH1prom-GAL4-TAD-ALY1(493-915), 2\mu, LEU2)</td>
<td>(ALY1) sequence encoding amino acids 493-915 was PCR amplified from pKK212-(ALY1) with primers containing (XmaI) and (XhoI) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>(493-915)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aly1-TAD</td>
<td>(ADH1prom-GAL4-TAD-ALY1(725-915), 2\mu, LEU2)</td>
<td>(ALY1) sequence encoding amino acids 725-915 was PCR amplified from pKK212-(ALY1) with primers containing (XmaI) and (XhoI) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>(725-915)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aly1-TAD</td>
<td>(ADH1prom-GAL4-TAD-ALY1(610-751), 2\mu, LEU2)</td>
<td>(ALY1) sequence encoding amino acids 610-751 was PCR amplified from pKK212-(ALY1) with primers containing (XmaI) and (XhoI) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>(610-751)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aly1-TAD</td>
<td>(ADH1prom-GAL4-TAD-ALY1(610-831), 2\mu, LEU2)</td>
<td>(ALY1) sequence encoding amino acids 610-831 was PCR amplified from pKK212-(ALY1) with primers containing (XmaI) and (XhoI) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>(610-831)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aly1-TAD</td>
<td>(ADH1prom-GAL4-TAD-ALY1(838-915), 2\mu, LEU2)</td>
<td>(ALY1) sequence encoding amino acids 838-915 was PCR amplified from pKK212-(ALY1) with primers containing (XmaI) and (XhoI) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>(838-915)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aly1-TAD</td>
<td>(ADH1prom-GAL4-TAD-ALY1(822-915), 2\mu, LEU2)</td>
<td>(ALY1) sequence encoding amino acids 822-915 was PCR amplified from pKK212-(ALY1) with primers containing (XmaI) and (XhoI) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>(822-915)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aly1\textsuperscript{APL, KIN}-TAD</td>
<td>(ADH1prom-GAL4-TAD-ALY1\textsuperscript{APL, KIN}(610-915), 2\mu, LEU2)</td>
<td>(ALY1\textsuperscript{APL, KIN}) sequence was PCR amplified from pKK212- (ALY1\textsuperscript{APL, KIN}) with primers containing (XmaI) and (XhoI) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Plasmid</td>
<td>Genotype</td>
<td>Description (Reference)</td>
</tr>
<tr>
<td>---------</td>
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<td>---------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Aly2-TAD</td>
<td>( ADH1prom-GAL4-TAD-ALY2, 2\mu, LEU2 )</td>
<td>( ALY2 ) coding sequence was PCR amplified from pKK212-( ALY2 ) with primers containing ( XmaI ) and ( XhoI ) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Rod1-TAD</td>
<td>( ADH1prom-GAL4-TAD-ROD1, 2\mu, LEU2 )</td>
<td>( ROD1 ) coding sequence was PCR amplified from genomic DNA with primers containing ( XmaI ) and ( SacI ) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Rog3-TAD</td>
<td>( ADH1prom-GAL4-TAD-ROG3, 2\mu, LEU2 )</td>
<td>( ROG3 ) coding sequence was PCR amplified from genomic DNA with primers containing ( XmaI ) and ( XhoI ) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Ldb19-TAD</td>
<td>( ADH1prom-GAL4-TAD-LDB19, 2\mu, LEU2 )</td>
<td>( LDB19 ) coding sequence was PCR amplified from genomic DNA with primers containing ( XmaI ) and ( XhoI ) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Ecm21-TAD</td>
<td>( ADH1prom-GAL4-TAD-ECM21, 2\mu, LEU2 )</td>
<td>( ECM21 ) coding sequence was PCR amplified from genomic DNA with primers containing ( XmaI ) and ( XhoI ) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Csr2-TAD</td>
<td>( ADH1prom-GAL4-TAD-CSR2, 2\mu, LEU2 )</td>
<td>( CSR2 ) coding sequence was PCR amplified from genomic DNA with primers containing ( XmaI ) and ( XhoI ) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Art10-TAD</td>
<td>( ADH1prom-GAL4-TAD-ART10, 2\mu, LEU2 )</td>
<td>( ART10 ) coding sequence was PCR amplified from genomic DNA with primers containing ( XmaI ) and ( XhoI ) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
<tr>
<td>Art5-TAD</td>
<td>( ADH1prom-GAL4-TAD-ART5, 2\mu, LEU2 )</td>
<td>( ART5 ) coding sequence was PCR amplified from genomic DNA with primers containing ( XmaI ) and ( XhoI ) restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)</td>
</tr>
</tbody>
</table>
### Plasmid | Genotype | Description (Reference)  
--- | --- | ---  
Bul1-TAD | *ADH1prom-GAL4-TAD-BUL1, 2μ, LEU2* | *BUL1* coding sequence was PCR amplified from genomic DNA with primers containing *XmaI* and *SacI* restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)  
Bul2-TAD | *ADH1prom-GAL4-TAD-BUL2, 2μ, LEU2* | *BUL2* coding sequence was PCR amplified from genomic DNA with primers containing *XmaI* and *XhoI* restriction site adaptors. The PCR product was then subcloned into pACT2. (This study)  
CNB-TAD (human) | *ADH1prom-GAL4-TAD-CNB, 2μ, LEU2* | The calcineurin regulatory subunit (CNB) from *Homo sapiens* was PCR amplified from a clone obtained from Open Biosystems with primers containing *BamHI* and *XhoI* restriction site adaptors. The PCR product was cloned into pACT2. (This study)  
pGBT9 | *ADH1prom-GAL4-DBD, 2μ, TRP1* | (89)  
DBD-Cna1 (yeast) | *ADH1prom-GAL4-DBD-CNA1, 2μ, TRP1* | (90)  
DBD-CNA (human) | *ADH1prom-GAL4-DBD-CNA isoform α, 2μ, TRP1* | The calcineurin catalytic subunit (CNA) isoform α from *Homo sapiens* was PCR amplified from a clone obtained from Open Biosystems with primers containing *BamHI* restriction site adaptors. The PCR product was cloned into pGBT9. (This study)