Mammalian RAFT1 kinase domain provides rapamycin-sensitive TOR function in yeast

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In complex with the prolyl isomerase FKBP12, the natural product rapamycin blocks signal transduction in organisms as diverse as yeast and man. The yeast targets of FKBP12-rapamycin, TOR1 and TOR2, are large proteins with homology to lipid and protein kinases. A mammalian FKBP12-rapamycin binding protein, RAFT1, shares 39% and 43% identity with TOR1 and TOR2 proteins, respectively but has not been linked to rapamycin action in vivo. We find that when expressed in yeast, neither wild-type nor mutant RAFT1 complemented tor2 mutations or conferred rapamycin resistance. In contrast, TOR2-RAFT1 and TOR1-RAFT1 hybrid proteins containing the carboxy-terminal RAFT1 kinase domain complemented tor2 and tor1 mutant strains, respectively. Moreover, TOR2-RAFT1 and TOR1-RAFT1 hybrid proteins mutated at the position corresponding to rapamycin-resistant TOR mutants (S2035I) conferred rapamycin resistance. Like the TOR2 protein, the TOR2-RAFT1 proteins were stably expressed, localized to the vacuolar surface, and associated with a phosphatidylinositol-4 kinase activity. These findings directly link the mammalian TOR homolog RAFT1 to rapamycin action in vivo and indicate that the TOR/RAFT1 kinase domain has been functionally conserved from yeast to man.

[Key Words: Signal transduction; immunosuppression; prolyl isomerases; RAFT1 kinase domain; TOR; yeast; rapamycin]

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The natural product rapamycin has potent antifungal and immunosuppressive activities through its ability to form toxic complexes with the FKBP12 prolyl isomerase [for review, see Schreiber 1991; Cardenas et al. 1994]. The FKBP12–rapamycin complex inhibits signal transduction events required for G1 to S-phase cell cycle progression in yeast, in T lymphocytes stimulated with interleukin 2 [IL-2], and in additional mammalian cell types stimulated by a variety of different growth factors [Dumont et al. 1990; Heitman et al. 1991a; Albers et al. 1993; Morice et al. 1993; de Groot et al. 1994]. Rapamycin is currently in clinical trials as an immunsuppressant in organ transplant recipients.

Although rapamycin is structurally related to FK506, another immunosuppressive antifungal compound that also binds FKBP12, rapamycin does not inhibit the protein phosphatase calcineurin, which is the target of the FKBP12–FK506 complex [Liu et al. 1991]. The targets of the FKBP12–rapamycin complex were first identified in yeast as TOR1 and TOR2 [Heitman et al. 1991a], ~245-kD proteins that share 67% identity and have a carboxy-terminal domain with similarity to lipid and protein kinases [Cafferkey et al. 1993; Kunz et al. 1993; Helliwell et al. 1994]. TOR2 is an essential protein [Kunz et al. 1993], whereas TOR1 is not [Cafferkey et al. 1993; Helliwell et al. 1994], suggesting that inhibition of TOR2 function by FKBP12–rapamycin is responsible for rapamycin toxicity. However, cells depleted of TOR2 arrest randomly in the cell cycle, whereas those depleted of both TOR1 and TOR2 arrest in G1, indicating that the G1 cell cycle arrest imposed by rapamycin involves inhibition of both TOR proteins [Kunz et al. 1993].

The observation of nonallelic noncomplementation between TOR1, TOR2, and FKBP12 mutants led to the model that FKBP12–rapamycin binds to and inhibits the TOR proteins [Heitman et al. 1991a], which has now been confirmed both in vitro and in vivo [Stan et al. 1994; Cardenas and Heitman 1995; Lorenz and Heitman 1995; Zheng et al. 1995]. The observations that rapamycin caused a similar cell cycle arrest in yeast and man, that yeast and human FKBP12 are 54% identical and have virtually superimposable tertiary structures [Van Duyne et al. 1991; Rotonda et al. 1993], and that human FKBP12 is functional in yeast [Koltin et al. 1991] all suggested that the TOR proteins would be conserved from yeast to man. An FKBP12–rapamycin binding protein (RAFT1, FRAP, RAPT1, SEP, or mTOR) with marked similarity to TOR1 and TOR2 has been identified [Brown et al. 1994; Chiu et al. 1994; Sabatini et al. 1994; Chen et al. 1995; Sabers et al. 1995]. However, evidence linking this mammalian FKBP12–rapamycin binding protein to rapamycin action in vivo has been limited.

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We demonstrate here that whereas full-length RAFT1 failed to provide TOR function in yeast, hybrid TOR2-RAFT1 proteins containing the amino-terminal two-thirds of TOR2 fused to the RAFT1 carboxy-terminal kinase domain function as TOR2 in yeast. Moreover, S203SI TOR2-mRAFT1 and S203SI TOR1-mRAFT1 mutant proteins rendered yeast rapamycin-resistant; importantly, this mutation corresponds to previously identified rapamycin-resistant TOR1 and TOR2 mutations. Finally, we show that the TOR2-RAFT1 fusion proteins are stably expressed and, as is the case with a TOR2, associated with the surface of the vacuole and PI-4 kinase activity. Our findings directly link the mammalian RAFT1 protein to rapamycin-sensitive growth control in vivo and suggest that RAFT1 performs similar functions in mammalian cells.

Results

Full-length RAFT1 does not provide TOR2 or TOR1 function

The mammalian FKBP12-rapamycin binding protein, RAFT1, shares 39% and 43% overall identity with the yeast TOR1 and TOR2 proteins, respectively (Sabatini et al. 1995). We tested whether RAFT1 functions as TOR2 or TOR1 in yeast. A plasmid expressing RAFT1 from the yeast GAL1 promoter (pGAL-RAFT1) was introduced into a diploid yeast strain in which one copy of the TOR2 gene had been disrupted (TOR2/αtor2::LEU2). Tetrad analysis revealed 2 viable: 2 inviable meiotic segregants in all of 60 tetrads examined; in all cases viable segregants were Leu−, indicating that the tor2::LEU2 mutation is not complemented by RAFT1 (data not shown; summarized in Fig. 1). In contrast, expression of the TOR2 gene from the GAL1 promoter restored viability to tor2::LEU2 meiotic segregants (data not shown). These findings suggest that RAFT1 does not provide TOR2 function in yeast.

Although TOR1 is not essential in many yeast strains (Cafferkey et al. 1993; Helliwell et al. 1994), we find that tor1 null mutations are conditionally lethal in srk1 mutant yeast strains at 39°C (D. Fiorentino, pers. comm.; M. Lorenz and J. Heitman, unpubl.). We therefore tested whether expression of RAFT1 from the yeast GAL1 promoter would complement a tor1 Δsrk1 mutant strain (GAY1). Whereas expression of wild-type TOR1 complemented and restored growth of the tor1 Δsrk1 mutant strain at 39°C, RAFT1 did not (data not shown; summarized in Fig. 1). These findings suggest that RAFT1 does not provide TOR1 function in yeast.

Attempts to detect the RAFT1 and mRAFT1 proteins in these complementation assays by Western blot anal-
TOR1 and TOR2 proteins are highly related (67% identity), neither complements for the function in yeast. A failure of RAFT1 to provide TOR1 may be unstable or unable to provide TOR1 or TOR2 function in yeast. A failure of RAFT1 to provide TOR function in yeast would not be unanticipated, given that although the yeast TOR1 and TOR2 proteins are highly related (67% identity), neither complements for the other under standard conditions (Helliwell et al. 1994). Even when overexpressed, only TOR2 can provide TOR1 function, whereas TOR1 does not provide TOR2 function (Helliwell et al. 1994).

**TOR2–RAFT1 hybrids provide TOR2 function in yeast**

The highest degree of identity (~60%) between RAFT1 and the TOR proteins is in a carboxy-terminal domain similar to the catalytic domains of phosphatidylinositol [PI] and protein kinases. Previous studies demonstrated that the kinase domains of TOR1 and TOR2 are functionally interchangeable (Helliwell et al. 1994). Therefore, we tested whether the kinase domain of RAFT1 could functionally replace those of the yeast TOR proteins. For this purpose, hybrid genes were constructed in a CEN LEU2 plasmid in which the promoter region and the amino-terminal two-thirds of TOR2 (residues 1–1687) were fused to the carboxy-terminal one-third of TOR1 (residues 1681–2470) or RAFT1 (residues 1721–2549), and the promoter region and amino terminal two-thirds of TOR1 (residues 1–1680) were fused to the carboxy-terminal one-third of RAFT1 (residues 1721–2549). In addition, TOR2–mRAFT1 and TOR1–mRAFT1 mutant hybrid genes were also constructed in which Ser2035 of RAFT1 was mutated to isoleucine, corresponding to TOR1 and TOR2 mutations that confer rapamycin resistance. The structures of these hybrid proteins are depicted in Figure 1.

Wild-type TOR2 and TOR1, and the TOR2–TOR1, TOR2–RAFT1, TOR2–mRAFT1, TOR1–RAFT1, and TOR1–mRAFT1 S2035I hybrids (Fig. 1) were tested for their ability to complement and restore viability in a tor2 mutant strain. Low-copy-number centromeric plasmids expressing these proteins were introduced into a tor2 mutant strain containing the wild-type TOR2 gene on a URA3 plasmid [Δtor2::ADE2 Δade2 [p2μ-TOR2–URA3] strain MH346-1a]. Leu + transformants were subjected to a plasmid shuffle on 5-fluoro-orotic acid (5-FOA) medium to eject the resident wild-type TOR2 plasmid and reveal the phenotype conferred by the introduced plasmid. In this assay only those cells expressing a functional TOR2 gene from the introduced plasmid survive. As shown in Figure 3B, wild-type TOR2, and the TOR2–TOR1, TOR2–RAFT1, and TOR2–mRAFT1 hybrid proteins were functional and complemented the tor2 deleted strain; TOR1, TOR1–RAFT1, and TOR1–mRAFT1 did not. tor2 mutant cells expressing TOR2–TOR1 or TOR2–RAFT1 fusion proteins grew at the same rate as isogenic wild-type TOR2 cells, were not cold or heat sensitive [data not shown], and were fully rapamycin sensitive [see below, Fig. 3D], indicating that the hybrid proteins provide TOR2 activity that, by these measures, is indistinguishable from wild type. Furthermore, the rapamycin sensitivity of tor2 mutant cells expressing TOR2–RAFT1 required FKBP12 [data not shown].

**Figure 2.** Northern blot analysis of RAFT1 and mRAFT1 expression in yeast. Total RNA was extracted from the TOR1 TOR2 wild-type strain JK9-3d containing the control plasmid pSEYC68 (vector), or the pGAL–RAFT1 [RAFT1] and the pGAL–S2035I–mRAFT1 [mRAFT1] expression plasmids and grown under repressing (−, glucose) or inducing (+, galactose) conditions. RNA was electrophoresed in a 1% gel and transferred to a nitrocellulose filter and hybridized with a 32P-radio-labeled probe derived from the RAFT1 gene (A) [see Materials and methods] or stained with ethidium bromide to reveal rRNA (B). Numbers at left designate the migration position of molecular weight size standards; the arrow at right indicates the position of migration of the RAFT1 mRNA.
with previous studies in which increased expression of wild-type TOR2, or of a TOR2–TOR1 hybrid protein, complemented the slow growth phenotype of a tor1 mutant (Helliwell et al. 1994). In contrast, the TOR1–RAFT1 and TOR2–RAFT1 hybrid proteins failed to complement the tor1 mutation when expressed from a centromeric plasmid (Fig. 3C), even though the TOR2–RAFT1 hybrid proteins clearly have TOR2 function (Fig. 3B) and are expressed at equivalent levels compared to endogenous TOR2 based on Western blot analysis (Fig. 5, below). In the case of the TOR1–RAFT1 hybrid, expression from a multicopy 2µ plasmid partially complemented the conditional lethality of a tor1 Δsrk1 mutant, indicating that the TOR1–RAFT1 protein provides only partial TOR1 function (Fig. 1; data not shown). The failure of the TOR1–RAFT1 hybrid to provide full TOR1 function could be attributable, at least in part, to the significantly lower level of expression of the hybrid protein compared with the wild-type TOR1 protein (Fig. 4B). In summary, we conclude that the RAFT1 kinase domain provides TOR2 activity when fused to TOR2 and provides partial TOR1 activity when fused to TOR1.

S203SI TOR2–RAFT1 and S203SI TOR1–RAFT1 hybrid proteins confer dominant rapamycin resistance

The FKBP12–rapamycin complex is toxic in yeast and mutations in a conserved serine residue of TOR1 or TOR2 confer rapamycin resistance (Heitman et al. 1991a; Cafferkey et al. 1993; Helliwell et al. 1994). This serine residue is also conserved in RAFT1 (S2035); therefore, we investigated whether the corresponding RAFT1 mutation would confer rapamycin resistance in yeast. Expression of the full-length S2035 mRAFT1 protein failed to confer rapamycin resistance (Fig. 1). We therefore introduced the S2035I mutation into the TOR2–RAFT1 and TOR1–RAFT1 hybrid proteins. As tested by the tor2 complementation assay, the TOR2–mRAFT1 S2035I hybrid protein fully restored TOR2 activity to tor2 deleted cells (Fig. 3B). Whereas growth of wild-type TOR1 TOR2 yeast cells expressing full-length RAFT1, or the TOR1–RAFT1 or TOR2–RAFT1 hybrid proteins, was inhibited by rapamycin, the TOR2–mRAFT1 S2035I mutant protein conferred resistance up to 10 μg/ml of rapamycin (Fig. 3D). Moreover, because rapamycin resistance was observed in a yeast strain also expressing wild-type TOR2, the TOR2–mRAFT1 hybrid protein confers dominant drug resistance. Finally, the TOR2–mRAFT1 protein conferred rapamycin resistance at 24, 30, 37, or

Figure 3. TOR2 and TOR1 activity and rapamycin resistance conferred by TOR–RAFT hybrid proteins. [A] Isogenic strains expressing wild-type TOR2 and TOR1, and the TOR2–TOR1, TOR2–RAFT1, TOR2–mRAFT S2035I, TOR1–RAFT1, and TOR1–mRAFT S2035I hybrid proteins, were grown on medium as indicated and assayed for TOR2 function, TOR1 function, and rapamycin resistance. [B] Complementation of TOR2 function. Strain MH346-1a, a Δtor2::ADE2 strain hosting a TOR2 CEN URA3 plasmid, was transformed with the CEN LEU2 plasmids expressing wild-type and mutant TOR and TOR–RAFT1 hybrid proteins depicted in A. Transformants were grown on medium without (– FOA) or with (+ FOA) 5-FOA to select against the TOR2 URA3 plasmid and determine whether TOR2 function is provided by the wild-type or hybrid gene introduced. Survival on FOA indicates TOR2 function. [C] Complementation of TOR1 function. A tor1 Δsrk1 yeast strain (CAY1) transformed with centromeric plasmid-borne wild-type and mutant TOR and TOR–RAFT1 hybrid genes was grown for 72 hr at 30°C and 39°C. Cells with TOR1 function survive at 39°C, whereas cells lacking TOR1 function do not. [D] Rapamycin effects on TOR–RAFT1 hybrids. A TOR1 TOR2 wild-type yeast strain (JK9-3da) transformed with centromeric plasmids expressing wild-type TOR and TOR–RAFT1 hybrid proteins was grown for 72 hr at 30°C on medium without (– RAP) or with (+ RAP) 10 μg/ml of rapamycin.
TOR kinase domain conserved from yeast to man

Figure 4. Rapamycin resistance and expression of TOR1–RAFT hybrid proteins. (A) Rapamycin effects on TOR1–RAFT1 hybrid proteins. A tor1 TOR2 yeast strain [MLY10a] lacking TOR1 (vector) or expressing TOR1 (TOR1), or the TOR1–RAFT1 wild-type protein [TOR1–RAFT1] or the S2035I TOR1–RAFT1 mutant protein [TOR1–mRAFT1] from the 2μ URA3 vector pRS306–2μ were grown for 72 hr at 39°C on medium without (0) or with 0.01, 0.1, or 10 μg/ml of rapamycin. (B) Expression of TOR1–RAFT1 hybrid proteins. Total cell extracts from strain MLY10a [tor1::LEU2 TOR2] transformed with the 2μ, URA3 plasmid pRS306–2μ (lane 1) or with the same plasmid bearing the wild-type TOR1 gene (lane 2, TOR1) or the TOR1 hybrid genes, TOR1–RAFT1 (lane 3) and S2035I TOR1–mRAFT1 (lane 4) were prepared. Western blot analysis with 60 μg of protein was performed using the anti-TOR1 antiserum. Note that the specificity of the anti-TOR1 antiserum is shown by the absence of cross-reacting material in the tor1 strain transformed with the control plasmid alone (lane 1). In C, 30 μg of protein from the same extracts in B was analyzed by Western blot for FKBP12 to verify that equal amounts of protein were loaded. In B and C, numbers at left indicate molecular mass (in kD) and the migration positions of TOR1, TOR1–RAFT1, and FKBP12 are indicated.

39°C, indicating that the mutant fusion protein is not temperature sensitive (data not shown).

Similarly, expression of the TOR1–mRAFT1 mutant hybrid protein conferred rapamycin resistance in a tor1 TOR2 strain [Fig. 4A] or in a TOR1 TOR2 strain [data not shown]. The ability of the TOR1–mRAFT1 mutant protein to confer rapamycin resistance was observed only with expression from a multicopy 2μ plasmid but not from a low-copy-number centromeric plasmid [Figs. 3D and 4A]. Rapamycin resistance was observed when cells were grown at 24°C–39°C [Fig. 4A], indicating that at least some of the mutant protein must be expressed at these temperatures. Western blot analysis with a TOR1–specific antiserum revealed that in comparison to wild-type TOR1, a smaller proportion of the TOR1–RAFT1 and TOR1–mRAFT1 hybrid proteins were expressed as full-length protein [Fig. 4B]. Thus, instability of the TOR1–RAFT1 hybrid proteins is a plausible explanation for the failure of the proteins to provide complete TOR1 function; the fact that rapamycin resistance required a higher level of expression of the mutant protein supports this view. Taken together, these findings directly link the activity of the mammalian RAFT1 protein to FKBP12–rapamycin-sensitive growth regulation in vivo.

TOR2–RAFT1 hybrid proteins are expressed as ~240-kD proteins

We have generated a polyclonal antiserum against a TOR2 carboxy-terminal fragment encompassing amino acids 2270–2474. Because TOR1, TOR2, and RAFT1 share a high degree of identity in this region [residues 2270–2370 of TOR2 and the corresponding regions of RAFT1 and TOR1 share 80% and 97% identity, respectively], we considered it likely that the anti-TOR2 antiserum would cross-react with TOR1 and RAFT1 to allow detection of TOR2–RAFT1 and TOR2–TOR1 hybrid proteins by Western blot analysis. Protein extracts were prepared from cells in which the chromosomal TOR1 and TOR2 genes had been deleted and a plasmid-borne TOR2 gene was ejected from the cell and replaced, via plasmid shuffle, with plasmids encoding TOR2 [as a control] or the TOR2–TOR1, TOR2–RAFT1, and TOR2–mARAFT1 hybrids. As shown in Figure 5, the TOR2–TOR1, TOR2–RAFT1, and TOR2–mRAFT1 hybrid proteins were readily detectable as proteins of ~240 kD that were expressed at a level comparable to the endogenous chromosomal TOR2 gene [Fig. 5, cf. lane 1 with 2–5]. This finding is in full accord with our in vivo observa-
Recognizes an amino-terminal TOR2 peptide revealed brid proteins were similarly localized to the surface of man 1995) and RAFT1 (Sabatini et al. 1995) contain PI-4 kinase activity. Immunoprecipitates of both TOR2 (Cardenas and Heitman 1995) and RAFT1 (Sabatini et al. 1995) contain PI-4 kinase activity. To determine whether TOR2-RAFT1 hybrid proteins have a similar associated activity, wild-type TOR2 and the TOR2-TOR1, TOR2-RAFT1, and TOR2-mRAFT1 hybrid proteins were immunoprecipitated with the anti-TOR2 carboxy-terminal antiserum that effectively cross-reacts with TOR2, TOR1, and RAFT1 [Fig. 5]. In addition, the proteins were obtained from cells in which the chromosomal TOR1 and TOR2 genes had been disrupted and only a plasmid-borne TOR2, TOR2-TOR1, or TOR2-RAFT1 hybrid gene was present. The resulting immunoprecipitates were assayed for PI kinase activity by incubation with [γ-^32P]ATP. PI kinase reactions were resolved by thin layer chromatography in the borate system (Walsh et al. 1991) to resolve PI3P and PI4P. As described previously for the TOR2 and RAFT1 proteins (Cardenas and Heitman 1995; Sabatini et al. 1995), a readily detectable PI-4 kinase activity was present in immunoprecipitates of the TOR2-TOR1, TOR2-RAFT1, and TOR2-mRAFT1 fusion proteins [Fig. 6]. Moreover, the PI-4 kinase activity detected was at a level comparable to that associated with the endogenous TOR2 protein immunoprecipitated in parallel. These findings reveal that a PI-4 kinase activity is tightly associated with and coimmunoprecipitates with the TOR2-TOR1 and TOR2-RAFT1 hybrid proteins.

**Discussion**

When bound to the peptidyl-prolyl isomerase FKBP12, the immunosuppressive antifungal natural product rapamycin inhibits signaling cascades required for growth and cell cycle progression, which are conserved from yeast to man. Genetic studies in yeast first implicated TOR1 and TOR2 as targets of the toxic FKBP12-rapamycin complex (Heitman et al. 1991a; Kunz et al. 1993). Both wild-type TOR1 and TOR2 directly interact with FKBP12-rapamycin whereas rapamycin-resistant TOR1 and TOR2 mutant proteins do not (Stan et al. 1994; Cardenas and Heitman 1995; Lorenz and Heitman 1995; Zheng et al. 1995). The resulting model is that the essential yeast TOR2 protein is required for events necessary for G1 to S-phase progression and cell division, and FKBP12-rapamycin binding to TOR2 inhibits some function necessary to execute these events properly. A mammalian FKBP12-rapamycin-binding protein with marked identity to the yeast TOR proteins has been identified and is variously referred to as RAFT1, FRAP, RAFT1, SEP, or mTOR; however, evidence linking this protein to rapamycin action in vivo has been limited and indirect (Brown et al. 1994; Chiu et al. 1994; Sabatini et al. 1994; Chen et al. 1995; Sabers et al. 1995). First, the decreased affinity of two FKBP12-rapamycin analog complexes [16-keto-rap and 25,26-iso-rap] for the FRAP protein is correlated with decreased in vivo activity (Brown et al. 1994). Second, introduction of mutations corresponding to rapamycin-resistant yeast TOR1 and TOR2 mutations reduces FKBP12-rapamycin binding to fragments of FRAP in vitro (Chen et al. 1995) or to a small portion of RAFT1 or RAFT1 in the two-hybrid in vivo assay (Chiu et al. 1994; Lorenz and Heitman 1995).
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in this case, the role of P70 S6 kinase, if any, in growth factor-dependent cellular proliferation remains to be established. An additional concern is that of the two FKBP12–rapamycin-binding proteins identified in yeast, TOR1 is not essential for growth and is therefore an unlikely target for the major FKBP12–rapamycin toxic effects. Thus, the mammalian FKBP12–rapamycin binding protein that has been identified might be a TOR1 homolog that does not play an essential role in rapamycin-sensitive signaling cascades in mammalian cells.

We have assessed the ability of the mammalian FKBP12–rapamycin-binding protein RAFT1 to function in FKBP12–rapamycin sensitive, TOR-dependent cell growth by heterologous expression in the yeast Saccharomyces cerevisiae. Expression of full-length RAFT1 failed to provide either TOR1 or TOR2 function in yeast. Similarly, expression of an S2035I RAFT1 mutant protein, corresponding to rapamycin-resistant TOR mutants, failed to confer rapamycin resistance in yeast. With the caveat that although we detect the RAFT1 mRNA in these cells we have been unable to detect the RAFT1 protein, these findings suggest that full-length RAFT1 lacks some activity required for TOR function in yeast. For example, RAFT1 might fail to associate properly with yeast proteins required for TOR function and cellular localization or might not be properly folded, post-translationally modified, or stably expressed in yeast.

As an alternative approach to assess RAFT1 function in yeast, we constructed TOR2–RAFT1 and TOR1–RAFT1 hybrid proteins in which the carboxy-terminal one-third of the TOR proteins, encompassing the kinase domain, was replaced with the corresponding domain of RAFT1. These experiments were based on the earlier observation that the kinase domains of TOR1 and TOR2 are functionally interchangeable (Helliwell et al. 1994). Remarkably, we find that a TOR2–RAFT1 hybrid protein can functionally replace the yeast TOR2 protein. The TOR2–RAFT1 hybrid protein supports growth that, by several measures, is indistinguishable from that of wild-type cells, including sensitivity to FKBP12–rapamycin and normal growth under a variety of different physiological conditions. Moreover, although some conditions that compromise TOR2 function render TOR1 essential (Cardenas and Heitman 1995), TOR1 is not essential in the TOR2–RAFT1 strain. One distinction is that when expressed from a centromeric plasmid, wild-type TOR2 can substitute for TOR1 (Fig. 3C), but the TOR2–RAFT1 fusion protein does not, even though this fusion protein clearly provides TOR2 function. In addition, the TOR1–RAFT1 fusion protein provided partial TOR1 function in a TOR1-dependent yeast strain when expressed from a multicopy 2μ plasmid. As a second measure of RAFT1 function in yeast, we demonstrated that S2035I mutant TOR2–RAFT1 and S2035I TOR1–RAFT1 hybrid proteins confer dominant rapamycin resistance in yeast (Figs. 3D and 4A). Finally, we found that like the wild-type TOR2 protein, the TOR2–TOR1 and TOR2–RAFT1 hybrid proteins were both localized to the vacuolar surface and associated with a PI-4 kinase activ-

Third, no binding of the mTOR protein to FKBP12–rapamycin affinity matrices was observed with extracts from rapamycin-resistant Jurkat T-cell lines (Sabers et al. 1995); however, in this case it is not known whether these mutant cell lines express mTOR, and no mutations in the mTOR locus have been identified. Finally, after this manuscript had been submitted, it was reported that FRAP regulates p70 S6 kinase in human T cells in a rapamycin-sensitive fashion, linking FRAP to rapamycin action in vivo (Brown et al. 1995). However,
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ity. Taken together with previous studies [Cardenas and Heitman 1995], these findings reveal a further correlation suggesting that TOR2 activity in vivo requires both proper localization and an associated lipid kinase activity. We note that Sabatini and colleagues have also detected PI-4 kinase activity associated with RAFT1 [Sabatini et al. 1995].

Several recent reports document that kinase inactive mutants of TOR1 [Zheng et al. 1995; Cardenas and Heitman, unpubl.], TOR2 [Cardenas and Heitman 1995], and FRAP [Brown et al. 1995] fail to complement tor1 or tor2 mutations, to activate P70 S6 kinase in vivo, or to confer rapamycin resistance. These findings indicate that the activity of the kinase domain of TOR proteins is required for biological function. Our studies reveal that the kinase domain of TOR2 and RAFT1 has been functionally conserved from yeast to man and demonstrate that this domain of RAFT1 can function in rapamycin-sensitive, TOR2-dependent signaling cascades in yeast. Moreover, our findings demonstrate further that the RAFT1 kinase domain is functionally related to TOR2, mitigating the concerns that RAFT1 might be a nonessential TOR homolog in mammals. Finally, our finding that a mutation corresponding to rapamycin-resistant TOR1 and TOR2 mutations similarly renders TOR2-RAFT1 and TOR1-RAFT1 hybrid proteins rapamycin resistant directly links the RAFT1 protein to FKBP12-rapamycin action in vivo. Further studies will be required to test whether the corresponding RAFT1 mutant protein confers rapamycin resistance in mammalian cells responding to growth factors, such as IL-2, and to determine the roles of RAFT1 in rapamycin-inhibitable mammalian signal transduction cascades.

Materials and methods

Yeast strains

Yeast strains that were employed in this study were all isogenic derivatives of strain JKg-3d (MATa, MATa, or MATa/a ura3-52 leu2-3,112 trpl-1 his4 rmel HMLa) (Heitman 1993) for biological function. Our studies reveal that the kinase inactive mutants of TOR1 [Zheng et al. 1995; Cardenas and Heitman, unpubl.], TOR2 [Cardenas and Heitman 1995], and FRAP [Brown et al. 1995] fail to complement tor1 or tor2 mutations, to activate P70 S6 kinase in vivo, or to confer rapamycin resistance. These findings indicate that the activity of the kinase domain of TOR proteins is required for biological function. Our studies reveal that the kinase domain of TOR2 and RAFT1 has been functionally conserved from yeast to man and demonstrate that this domain of RAFT1 can function in rapamycin-sensitive, TOR2-dependent signaling cascades in yeast. Moreover, our findings demonstrate further that the RAFT1 kinase domain is functionally related to TOR2, mitigating the concerns that RAFT1 might be a nonessential TOR homolog in mammals. Finally, our finding that a mutation corresponding to rapamycin-resistant TOR1 and TOR2 mutations similarly renders TOR2-RAFT1 and TOR1-RAFT1 hybrid proteins rapamycin resistant directly links the RAFT1 protein to FKBP12-rapamycin action in vivo. Further studies will be required to test whether the corresponding RAFT1 mutant protein confers rapamycin resistance in mammalian cells responding to growth factors, such as IL-2, and to determine the roles of RAFT1 in rapamycin-inhibitable mammalian signal transduction cascades.

Plasmids

Plasmids pGAL-RAFT1 (CEN LEU2) and pGAL-mRAFT1 S2035I (CEN URA3) express RAFT1 from the GALI promoter of plasmid pSEYC68. RAFT1 was PCR amplified from a RAFT1 cDNA [Sabatini et al. 1994] with primers 5′-GCTCTAGAC-GAACTCAGGGCAAGATGCT and 5′-GCCACAGCTCC-TTGTCTCTGACAGTGT containing XbaI and HindIII sites, respectively, in bold. The ~8-kb PCR product was XbaI-HindIII cleaved and cloned into pSEYC68. mRAFT1 S2035I was created by PCR overlap mutagenesis [Ho et al. 1989] using mutagenic primers 5′-CAAGCGAATGGCCTCTFCTAGGCC-CT and 5′-GGTGAAGGCCATACAGATTGA-GAG (hybridizes 5′ to the unique KpnI site in RAFT1) and 5′-GCCACAGCTCC-TTGTCTCTGACAGTGT. This PCR product was KpnI-HindIII cleaved and used to replace the wild-type fragment in pGAL-RAFT1.

Plasmids expressing TOR2 or TOR1 hybrid proteins were constructed as follows. pTOR2–TOR1 was constructed by cleaving plasmid pML40 (a derivative of CEN LEU2 plasmid pRS315 [Sikorski and Hieter 1989] bearing an 8-kb NotI fragment containing the TOR2 gene [Lorenz and Heitman 1995]) with BamHI, and replacing the excised region with a 2.6-kb BamHI fragment from plasmid pCNTOR1 [a derivative of CEN URA3 plasmid pRS316 [Sikorski and Hieter 1989] bearing an 8-kb NotI fragment containing theTOR1 gene [Muir and Heitman, unpubl.]]

The TOR2–TOR1 hybrid containing one-third TOR2 and two-thirds TOR1 was constructed by PCR amplification of a segment of TOR1 using pCNTOR1 as a template and primers 5′-GGTCACGGTATCTTAAAGATTGTTTCTCCTTAA and 5′-GCCGTCGAGTACCGTGGCCATACATCCAAT [Muil and Sall sites shown in bold]. The ~5.3-kb PCR product was cloned in the TA cloning system [Invitrogen], excised with MluI–SalI, and subcloned into gapped pML40 that had been digested with MluI–SalI.

pTOR2–RAFT1 was constructed by PCR amplification of the RAFT1 kinase domain with primers 5′-CUGGATCCGGTG-CAGACCTATGCGACAGCATGACG [467] and 5′-GCCACAGCTTC-TCTTTGTTTCTAGCTTGCT [411]. The ~2.6-kb PCR product was cloned into the TA cloning system [Invitrogen], excised with MluI–SalI, and cloned in the pML40 site of pML40. pTOR2–mRAFT1 S2035I was constructed as pTOR2–RAFT1 except pGAL–mRAFT1 S2035I was used as PCR template. The TOR2–RAFT1 hybrids containing one-third TOR2 and two-thirds TOR1 were constructed similarly to the TOR2–TOR1 hybrid described above using RAFT1 cDNA or pGAL–mRAFT1 as templates and primers 5′-CCGGACGGTCGTTGGTGGAC-GAG (hybridizes 5′ to the unique KpnI site in RAFT1) and 5′-CCACGGTGCGACCATCAGCTGGTCTCTTGCAGCTGCT.

pTOR1–RAFT1 and pTOR1–mRAFT1 S2035I were constructed by partially digesting plasmid pCNOTOR1 with BamHI, purifying the linearized plasmid (12.5 kb), digesting with XbaI, and purifying the BamHI to XbaI-gapped plasmid (10 kb). RAFT1 and mRAFT1 S2035I kinase domains were excised from the TA cloning vector as BamHI–XbaI fragments and subcloned into the 10-kb pCNOTOR1 gapped plasmid. In addition, TOR1, TOR1–RAFT1, and TOR1–mRAFT1 were subcloned as NotI cassettes into the CEN LEU2 and 2μ URA3 plasmids pRS315 and pRS306-2μ, respectively [Sikorski and Hieter 1989].

Northern blot analysis

Total yeast RNA was isolated by the hot phenol method according to Schmitt et al. [1990]. Total RNA was separated on a 1% agarose gel containing formaldehyde and transferred to a nitrocellulose membrane by the capillary method. Hybridization was performed as described [Sambrook et al. 1989]. A RAFT1 PCR fragment using oligonucleotides 467 and 411 [described
above] was gel purified and radiolabeled using a random priming kit (Boehringer-Mannheim).

**Antisera**

Generation and characterization of polyclonal antisera against the TrpE–TOR2 carboxy-terminal fusion protein [used here for Western blots and immunoprecipitation] and amino- and carboxy-terminal TOR2 peptides [used for immunofluorescence] have been described [Cardenas and Heitman 1995].

An anti-TOR1 antisera was generated as follows. An amino-terminal fragment of the TOR1 gene encoding amino acids 1–100 was PCR amplified with oligonucleotides 5'-ATATGGATCCCATGGAACCGCATGAGGAGCAG (313) and 5'-CGGCCGCTGCAGCTACTATAGATCAAATAATGAAATAC-TGGCC (304). The resulting PCR product was purified, cleaved with BamHI–PstI, and cloned into the BamHI–PstI sites of the TrpE fusion vector pATH22 [Koerner et al. 1991]. The TOR1–TrpE fusion protein was overexpressed and recovered from inclusion bodies as described [Koerner et al. 1991]. For rabbit immunization, the TOR1–TrpE protein was gel purified and electroeluted. Rabbits were injected, boosted, and bled by standard procedures [Harlow and Lane 1988]. The anti-FKBP12 polyclonal antisera has been described previously [Cardenas and Heitman 1995]. Antisera and affinity-purified antibodies directed against RAFT1 peptides corresponding to residues 64–79 (α-64–RAFT1), 782–797 (α-782–RAFT1), and 2528–2549 (α-carboxy-term–RAFT1) have been described [Sabatini et al. 1995] and were generously provided by David Sabatini and Solomon Snyder (Johns Hopkins University, Baltimore, MD).

**Yeast cell extracts**

Preparation of yeast cell extracts for Western blot and immunoprecipitation were as described previously [Cardenas and Heitman 1995].

**Immunological techniques**

Western blot, immunofluorescence, and immunoprecipitations were performed exactly as described previously [Cardenas and Heitman 1995].

**PI kinase assays**

TOR2 immunoprecipitates were assayed for PI kinase activity as described previously [Cardenas and Heitman 1995].

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