Genetic evidence links the ASTRA protein chaperone component Tti2 to the SAGA transcription factor Tra1

Julie Genereaux¹, Stephanie Kvas¹, Dominik Dobransky¹, Jim Karagiannis²
Gregory B. Gloor¹ and Christopher J. Brandl¹ *

¹Department of Biochemistry, Schulich School of Medicine & Dentistry, The University of Western Ontario, London Canada N6A5C1

²Department of Biology, The University of Western Ontario, London Canada N6A 5B7
Running title: TTI2 alleles suppress tra1

Keywords: Tra1; Tti2, FATC domain; PIKK proteins; gene regulation

*Corresponding author: Dr. Christopher J. Brandl
Department of Biochemistry,
Schulich School of Medicine & Dentistry
University of Western Ontario,
London, CANADA N6A 5C1
519-661-2111 ext 86857
Fax: 519-661-3175,
cbrandl@uwo.ca
ABSTRACT

Tra1 is a 3744-residue component of the *Saccharomyces cerevisiae* SAGA, NuA4 and ASTRA complexes. Tra1 contains essential C-terminal PI3K and FATC domains, but unlike other PIKK (phosphoinositide three-kinase-related kinase) family members, lacks kinase activity. To analyze functions of the FATC domain we selected for suppressors of *tra1-F3744A*, an allele that results in slow growth under numerous conditions of stress. Two alleles of *TTI2*, *tti2-F328S* and *tti2-I336F*, acted in a partially dominant fashion to suppress the growth related phenotypes associated with *tra1-F3744A* as well as its resulting defects in transcription. *tti2-F328S* suppressed an additional FATC domain mutation (*tra1-L3733A*), but not a mutation in the PI3K domain or deletions of SAGA or NuA4 components. We find eGFP-tagged Tti2 distributed throughout the cell. Tti2 is a component of the ASTRA complex, and in mammalian cells associates with molecular chaperones in complex with Tti1 and Tel2. Consistent with this Tra1 levels are reduced in a strain with a temperature sensitive allele of *tel2*. Further agreeing with a possible role for Tti2 in the folding or stabilization of Tra1, *tra1-F3744A* was mislocalized to the cytoplasm, particularly under conditions of stress. Since an intragenic mutation of *tra1-R3590I* also suppressed F3744A, we propose that Tti2 is required for the folding/stability of the C-terminal FATC and PI3K domains of Tra1 into their functionally active form.

Tra1 and its human homolog TRRAP are members of the PIKK (phosphoinositide three-kinase-related kinase) family of proteins. The family also includes the key cellular regulators ATM (ataxia telangiectasia mutated), ATR (ATM and Rad3-related), DNA-PKcs
(DNA-dependent protein kinase catalytic subunit), mTOR (mammalian target of rapamycin), and SMG-1 (suppressor with morphological effect on genitalia family member), many with yeast equivalents (Abraham et al., 2004; Lovejoy and Cortex, 2009). Tra1 is essential for viability in *S. cerevisiae* and a major constituent of the multisubunit SAGA and NuA4 transcriptional regulatory complexes (Grant et al., 1998; Saleh et al., 1998; Allard et al., 1999). SAGA and NuA4 have been well studied with regard to their functions, playing roles in multiple aspects of gene regulation and DNA repair (Doyon and Cote, 2004; Rodriguez-Navarro, 2009; Koutelou et al., 2010). Both possess histone acetyltransferase activity, the catalytic subunits being Gcn5 and Esa1 for SAGA and NuA4, respectively (reviewed in Roth et al., 2001). SAGA contains additional modules, critical for regulation, that function in the deubiquitylation of histone H2B (Henry et al., 2003) and interaction with the TATA binding protein (Dudley et al., 1999; Mohibullah and Hahn, 2008). Structural data indicates that Tra1 comprises a separate module in SAGA and NuA4 (Wu et al. 2004; Chittuluro et al., 2011). The position of Tra1 in these complexes is not indicative of a scaffold function, a result consistent with Tra1 not being required for stability of *S. pombe* SAGA (Helmlinger et al., 2011). This function is primarily ascribed to Spt7 for SAGA and Eaf1 for NuA4 (Sterner et al., 1999; Auger et al., 2008). TRRAP was first identified through its interactions with the transcription factors c-myc and E2F (McMahon et al., 1998). Similarly Tra1 interacts with yeast transcription factors, targeting SAGA and NuA4 to specific promoters (Bhaumik et al. 2004; Brown et al. 2001; Fishburn et al. 2005; Reeves and Hahn 2005). Independent functions for Tra1 likely also exist since Helmlinger et al. (2011) found that in *S. pombe* deletion of the SAGA specific molecule spTra1 results in changes in gene expression distinct from loss of other SAGA subunits.
Based on the association of the members in a series of immunoprecipitation experiments, Shevchenko et al. (2008) identified Tra1 in a novel complex they termed ASTRA (ASsembly of Tel, Rvb and Atm-like kinase). The components of ASTRA are encoded by essential genes in S. cerevisiae and include: Tra1, Rvb1, Rvb2, Tel2, Tti1, Tti2 and Asa1. In S. pombe, Tra1 but not Tra2, is found in ASTRA (Helmlinger et al., 2011). Rvb1 and Rvb2 are components of multiple regulatory complexes (in S. cerevisiae the Ino80, Swr1 and R2TP complexes), likely in multimeric form (Jha and Dutta, 2009, Huen et al. 2010). Both belong to the AAA+ (ATPase Associated with diverse cellular Activities) family of ATP binding proteins. Lustig and Petes (1986) identified TEL2 in S. cerevisiae through mutations that result in generation dependent telomere shortening. The association of mammalian and fission yeast Tel2 with several PIKK family members suggested a broader role (Takai et al. 2007; Hayashi et al., 2007). Tti2 and Tti1 were implicated in Tel2-dependent processes based upon their mutual association in yeast and mammalian cells (Hayashi et al., 2007; Takai et al., 2007, 2010; Hurov et al., 2010; Kaizuka et al., 2010). At least one role for Tel2, Tti1 and Tti2 (TTT complex, Huroy et al., 2010) is likely in the folding/maturation of PIKK proteins since they affect the steady-state levels and bind newly synthesized PIKK molecules (Takai et al., 2007, 2010; Horejsí et al., 2010; Hurov et al., 2010; Kaizuka et al., 2010). Furthermore, the TTT complex is functionally associated with Hsp90, Hsp70, Hsp40 and the R2TP/prefoldin-like complex (Takai et al., 2010; Horejsí et al., 2010). In a recent study of essential genes, tti1 and asa1 were implicated in chromosome instability (Stirling et al., 2011). Stirling et al. (2011) also demonstrate that knock-down mutants of these genes, as well as tti2 and tel2, result in telomere shortening and a reduction in Tor1 protein levels.
The PIKK family members are large proteins (3744 residues for Tra1), characterized by a C-terminally positioned domain that resembles the phosphatidylinositol-3-kinases (PI3K; Lempiäinen and Halazonetis, 2009). Unlike the other PIKK family members, Tra1/TRRAP lack kinase activity (McMahon et al., 1998; Saleh et al., 1998). Despite this, altering residues that parallel key regions of the kinase members affect Tra1 function (Mutiu et al., 2007). Interestingly one of these mutations, \textit{tra1-SRR3413} results in age-dependent telomere shortening.

On the N-terminal side of the PI3K domain is a FAT (FRAP-ATM-TRRAP) domain that consists largely of HEAT (Huntington, Elongation Factor 3, PR65/A, TOR) and TPR (tetratricopeptide) repeats; indeed much of the protein is likely helical (Bosotti et al., 2000; Perry and Kleckner, 2003; Sibandi et al., 2010; Knutson and Hahn, 2011). To the C-terminal side of the PI3K domain is the less highly conserved PRD (PIKK regulatory domain), the site of acetylation of ATM by TIP60 (Sun et al., 2005; for a linear schematic of the domains see Lempiäinen and Halazonetis, 2009).

At the extreme C-terminus of the PIKK molecules is the 30-35 residue FATC domain (FAT C-terminal; Bosotti et al., 2000); the conservation of the domain is evident from the finding that some FATC domains can be exchanged without loss of function (Jiang et al. 2006). Addition of as little as a single glycine to the C-terminus of Tra1 results in loss of viability (Hoke et al., 2010). Other FATC mutations in Tra1 cause growth defects, such as temperature sensitivity and slow growth on media containing ethanol, Calcofluor white or rapamycin. The FATC domain is similarly important for the other PIKK family members (Beamish et al. 2000; Priestly et al. 1998; Sun et al. 2005; Takahashi et al. 2000). Of note the parallel mutation to L3733A of Tra1 results in a dramatic loss in the kinase activity of SMG-1 (Morita et al. 2007).
The structure of the isolated FATC domain of *S. cerevisiae* Tor1 has been determined (Dames *et al.*, 2005). It is largely helical in structure with a C-terminal loop held in place by a disulphide linkage. While the helical structure is likely conserved, the absence of the cysteines in the other FATC domains suggests the loop is not. FATC domains are proposed to serve as the target site for protein interactions. In a two-hybrid analysis the FATC domain of Mec1, the *S. cerevisiae* homolog of ATR, was required for association with the RPA components Rfa1 and Rfa2 (Nakada *et al.*, 2005). Similarly the FATC domain of ATM was shown to interact with Tip60 (Sun *et al.*, 2005); however, a recent report suggests that this is indirect (Sun *et al.*, 2010).

Lempiäinen and Halazonetis (2009) speculate that the FATC and PRD domains regulate the kinase domain through interactions with the activation loop similar to the helical domains of the PI3-kinases.

We selected for suppressors of the temperature and ethanol sensitivity of *tra1-F3744A* with the goal of identifying roles for the FATC domain. Two mutations in *tti2* were found in independent selections as partially dominant suppressors of the growth related phenotypes and transcriptional changes caused by *tra1-F3744A*. *tti2-F328S* suppressed a second FATC mutation but not a mutation within the PI3K domain or deletion of SAGA or NuA4 components. Consistent with the documented role of the TTT complex (Takai *et al.*, 2007, 2010; Horejsí *et al.*, 2010; Hurov *et al.*, 2010; Kaizuka *et al.*, 2010), Tra1 levels were reduced in a strain temperature sensitive for Tel2, and an increased level of cytoplasmic Tra1-F3744A was reversed by *tti2-F328S*. We predict that the basis for the suppression results from Tti2 having a role in the formation, stability and/or localization of active Tra1. Our finding that an intragenic mutation of arginine 3590 to isoleucine within the putative activation loop of Tra1 also suppresses the
F3744A mutation supports the models of Lempiäinen and Halazonetis (2009) and Sturgill and Hall (2009) that folding of the PIKK molecules involves the interaction of FATC and PI3K domains.
MATERIALS AND METHODS

**Yeast strains and growth:** Yeast strains, listed in Table 1, are derivatives of BY4741 and BY4742 (Winzeler and Davis, 1997). *HIS3*-linked *tral* strains, wild-type *TRA1* (CY4353), *tral*-L3733A (CY4057, CY4103) and *tral*-F3744A (CY4350, CY4351), are described in Hoke *et al.* (2010). The *HIS3* allele marking these alleles is placed at the *BstBI* site of YHR100C/1*GEP4*, 11 codons from the C-terminus. A copy of YHR100C on YCPlac111 or YCplac33 was present in each of these strains to ensure its functionality. Diploid strains containing one Flag5-tagged *TRA1* allele marked with *URA3* (CY4398) containing *TRA1*-HIS3 (CY4419) or *tral*-F3744A-HIS3 (CY4421) were similarly constructed as were N-terminally eGFP tagged *tral* strains. The haploid *tral*-F3744A strain (CY5524) containing 5’-*URA3*-Flag5 and linked to 3’ *HIS3* was obtained after sporulation. CY5639 and CY5640 containing Flag5-*tral-R3590I* and Flag5-*tral-3590I, F3744A were also generated after integration into CY4398 and sporulation. The *tral*-F3744A *tti2*-F328S (CY5567 and CY5669) and *tral*-F3744A *tti2*-I336F (CY5843) double mutant strains were obtained in the selection process described below. The *tti2*-F328S strain CY5665 was obtained after crossing CY5667 with BY4742. Double mutants with *ada2Δ0* (CY5876), *eaf3Δ0* (CY5916), and *eaf7Δ0* (CY5917) were generated after mating CY5667 with the consortium knockout strains (BY4842, BY7143 and BY4940 obtained from Open Biosystems) and screening Kanr resistant spore colonies for the *tti2* allele by sequencing of PCR products. The *tti2*-F328S double mutant with *tral*-L3733A was obtained after a cross with CY4103. The *spt7::LEU2* deletion strain used for double mutant analysis (CY5873) was created after five backcrosses of FY1093 (kindly provided by Fred Winston) with BY4741. RFP-tagged strains in the EY0987 background (*MATα his3Δ1 lys2Δ0 ura3Δ0*; Huh *et al.*, 2003) were kindly
provided by Peter Arvidson. Heterozygous diploid strains containing an RFP-tagged allele and
eGFP-tra1-F3744A were made by mating the EY0987 derivatives with CY6018. CY6146 and
CY6148 containing tra1-F3744A and either tti2-1 (Stirling et al., 2011) or tel2-15 (Grandin et
al., 2012), both temperature sensitive alleles, were produced after mating of CY4350 with
PSY561 or PSY42 (kindly provided by Peter Stirling), respectively, and sporulation. CY6141
with Flag5-tagged Tra1 and tel2-15 were produced from a cross of PSY43 with CY5919.
CY6137 was derived after a cross of CY2222 and CY5665.

Growth comparisons were performed on YP media containing 2% glucose (YPD) selective
plates after 3-5 days at 30° unless stated otherwise. Standard concentrations used for the
selections were: 0.03% methyl methanesulfonate (Sigma-Aldrich), 7.5 μg/mL Calcofluor white
(Sigma-Aldrich), 1.0 μg/mL tunicamycin (Sigma-Aldrich), 6% ethanol, 60 μg/mL brefeldin (LC
Laboratories, Woburn MA), 1.2 M NaCl, 1.0 M sorbitol, 20 μg/mL geneticin (Sigma-Aldrich),
1.0 μg/mL staurosporine (LC Laboratories, Woburn MA), 1.0 μg/mL phleomycin (Sigma-
Aldrich), 1 ng/ml rapamycin (LC Laboratories).

**DNA molecules:** Promoter-lacZ fusions cloned as his3-lacZ fusions into the LEU2 centromeric
plasmid YCp87 and have been described (Brandl et al., 1993; Mutiu et al., 2007; Hoke et al.,
2010). Molecules for integrating tra1 alleles linked to HIS3 are described in Hoke et al. (2010).
Integrative vectors to generate Flag5 and eGFP fusions of Tra1 were constructed in cassettes
using as the base a molecule synthesized by Integrative DNA Technologies. This molecule
pCB2143 (see supplemental Figure 1), cloned into pTZ19 (lacking the polylinker HindIII site) as
a SphI-BamHI fragment contains an SphI site at -549 relative to the translational start site of
TRA1, a HindIII site inserted at -351, the sequence AAAAAATGTCAGGATCC surrounding the
translational start and a BamHI-SalI fragment encoding a Flag5 tag followed by a NotI site in the alanine frame, to which the coding region of TRA1 to the Kpn1 site is added (Saleh et al., 1998). Inserted into this is the 1.1 kbp HindIII genomic fragment encoding URA3. The integrative plasmid was switched to eGFP by the replacement of the Flag cassette with a BamHI-Not1 cassette encoding eGFP (Hoke et al, 2008). Myc9-tagged TTI2 and tti2-F328S were expressed from the DED1 promoter in YCplac111 by inserting a NotI-SstI fragment amplified from genomic DNA using oligonucleotides 5693-1 and 5693-2 (Table 2) downstream of the DED1 promoter-myc9 cassette (Hoke et al., 2010). N-terminally eGFP-tagged TTI2 was expressed from the DED1 promoter on YCplac33 (Hoke et al, 2008). pYHR100C/GEP4 genomic DNA was inserted into YCplac111 or YCplac33 as a BamHI-EcoRI fragment after PCR with oligonucleotides 4966-1 and 4966-2.

Selection of suppressor strains: CY4350 containing YCplac111-YHR100C was grown to stationary phase in YPD. 10 µL of culture, approximately 2 million cells, was plated onto YPD and UV irradiated at a wavelength of 302 nm for 10 s. Survival was approximately 10%. Colonies growing at 37° were colony purified under nonselective conditions and reanalyzed for growth at 37° and on YPD plates containing 6% ethanol. The suppressor strains were crossed with the URA3-tagged tra1-F3744A strain, CY5928, to determine linkage of the suppressor with tra1-F3744A. One strain had an unlinked suppressor mutation that segregated in a 2:2 fashion. A MATa spore colony was backcrossed with CY4350 nine-times at each stage selecting for temperature and ethanol resistant spore colonies. The final isolate was named CY5667. This was sent for genomic sequencing as described below. The tti2 mutations were verified after isolation of genomic DNA, PCR with oligonucleotides 6061-1 and 5693-2 and sequencing of the PCR
product using 6061-1 as primer. The selection was repeated independently on 3 plates using an initial selection at 35° with 4% ethanol. Again one unlinked suppressor mutation was obtained. This was backcrossed seven times with CY4351 with the resulting strain called CY5843. The ttii2 allele was sequenced after PCR with oligonucleotides 5693-1 and 5693-2 using PWO polymerase (Roche Applied Science) and cloning of the resulting PCR product in pUC18.

**Genomic sequence analysis:** Three genomic DNAs were prepared, two strains containing the suppressor from independent tetrads and a control strain lacking the suppressor from the same tetrad as CY5667. Genomic DNA was prepared from 10 mL of lyticase treated cells (Ausubel et al., 1998). Removal of RNA was verified by gel electrophoresis. Approximately 5 µg of DNA from each sample was sent to the Centre for Applied Genomics (Toronto, Ontario) for DNA library construction and next-generation sequencing using paired-end reads with the Applied Biosystems SOLiD 4.0 platform. The *Saccharomyces cerevisiae* genome sequence was downloaded from the *Saccharomyces* Genome Database (SGD; http://www.yeastgenome.org) on March 24, 2011. Custom bash and Perl scripts were written for the sequencing analysis. The program Bowtie (Langmead et al., 2009), allowing up to three mismatches per read, was used to map the colorspace reads to each chromosome of the yeast genome and obtain mapped reads in SAM format (Sequence Alignment/Map; Li et al., 2009). The VCF (variant call format) from SAMtools (Li et al., 2009) was used to obtain a raw list of polymorphisms from the mapped reads. Those reads with a Phred quality score below 20 were eliminated to obtain a filtered list of polymorphisms. A custom Perl script was written to account for the polymorphisms found in wild-type samples. We do note that approximately five percent of the TTI2 reads for the suppressor strains were wild-type, not containing the F328S codon. We believe these represent
errors in the reading or synthesis of the bar-codes on the 14 other yeast sequences in the lane.

**β-galactosidase assays:** Yeast strains containing lacZ-promoter fusions were grown to stationary phase in media lacking leucine. Assays with PHO5-lacZ in media depleted of phosphate, and HIS4-lacZ in media lacking histidine were performed as described in Mutiu et al (2007); GAL10-lacZ in media containing 2% galactose as the carbon source in Brandl et al. (1993), and for RPL35a-lacZ in YPD as described by Hoke et al. (2010). o-nitrophenol-β-D-galactosidase was used as substrate and values normalized to cell density.

**Western blotting:** Western blotting was performed using PVDF membranes and anti-Flag (M2; Sigma-Aldrich) or anti-Myc (9E10; Sigma-Aldrich) antibodies as described by Mutiu et al. (2007) or Hoke et al. (2010).

**Chromatin immunoprecipitation assays:** ChIP assays using anti-H3 (Abcam, Inc ab1791), anti-AcH4 K8 (ab1760) and anti-AcH3 K18 (ab1191) were performed as described by Mutiu et al. (2007). Input chromatin for the immunoprecipitations was prepared from yeast strains CY4353, CY4350 and CY5667, and normalized by PCR analysis of serial dilutions using oligonucleotides 5583-1 and 5583-2 to the PHO5 promoter. Agarose gels were stained with ethidium bromide and bands quantified using AlphalImager 3400 software (Alpha Innotech, San Leandro, CA). Background from a mock immunoprecipitation was subtracted. Values presented are the mean percentages relative to wild-type (CY4353; TRA1 TTI2) of the ratio of acetylated histone (AcH3 or AcH4) to total histone H3 for 2 independent experiments. PGK1 promoter primers were 2927-1 and 2927-2.
Fluorescence Microscopy: Yeast cells expressing eGFP and/or RFP fusions were grown in synthetic complete media then diluted 1:4 into synthetic complete media with or without 8% ethanol. Growth in ethanol was for 18 hr. Prior to visualization cells were concentrated 10-fold and 4’6-diamidino-2-phenylindole (DAPI) added to 0.02 mg/mL. Fluorescent images were obtained using a Zeiss Axioskop 2 microscope driven by ImageJ 1.41 software (National Institutes of Health) and a Scion CFW Monochrome CCD Firewire Camera (Scion Corporation, Frederick Maryland) using DAPI, RFP and GFP filter sets. Quantification of GFP-signal intensity was performed using ImageJ software (version 1.45). The freehand selections tool was used to trace each whole cell and nucleus separately. The measure function output the signal intensity per unit area for each selection. To correct for background, the average intensity of three background selections adjacent to each cell was subtracted. The corrected nuclear intensity was then divided by the corrected whole cell intensity to give a nuclear to cell intensity ratio. The mean from 20 cells was calculated plus/minus a standard deviation.
RESULTS

**Isolation of suppressors of tra1-F3744A:** We previously demonstrated that the extreme C-terminus of Tra1, the FATC domain, plays a key role in the protein’s function (Hoke *et al.*, 2010). Altering the terminal Phe of Tra1 results in temperature sensitivity and slow growth in media containing 6% ethanol, Calcofluor white or rapamycin. As shown in Figure 1A, growth of a strain containing the *tra1-F3744A* allele is impaired in a variety of other stress conditions including media depleted of phosphate and containing galactose as the carbon source. The allele does not result in sensitivity to all stress, as the strain was relatively resistant to high concentrations of sodium chloride or sorbitol. To address the role of this region of Tra1, we initiated a genetic screen to identify suppressors of the *tra1-F3744A* allele. We plated approximately two million cells and subjected them to UV radiation at a dose that allowed 10% viability. Three colonies were isolated based on their fast growth in YPD at 37°C. Each of these isolated strains also grew in media containing 6% ethanol. To determine if the suppressor mutations were linked to *tra1-F3744A*, the suppressor strains were crossed with a strain containing *tra1-F3744A* positioned downstream of a *URA3* marker (CY5928). Analysis of the resulting tetrads revealed that each of the suppressors segregated as a single mutation, and one of the three (*SUP3*) was not closely linked to *tra1-F3744A*. A second independent screen was performed starting with approximately 10 million cells and selecting for growth at 35°C on media containing 4% ethanol. Again one unlinked suppressor (*SUPB*) was obtained. Growth of a wild-type strain (CY4353), the *tra1-F3744A* strain (CY4350) and the two suppressor strains on YPD at 30°C and 37°C, and on YPD containing 6% ethanol is shown in Figure 1B.
We analyzed the expression of Tra1-F3744A in wild-type and SUP3-containing strain backgrounds to address whether the suppressor altered the expression of Tra1. Diploid strains with one copy of Flag5-Tra1 or Flag5-Tra1-F3744A were grown in YPD media. As shown in Figure 1C the amount of Flag5-tagged Tra1-F3744A (lanes 3 and 6) is similar to wild-type Tra1 (lanes 2 and 5). This level is not altered by the SUP3 mutation (lanes 1 and 4), indicating that suppression is not mediated through a change in Tra1 expression. The protein shown in Figure 1C was prepared by lysis with glass beads to minimize the processing time. We have seen changes in the profiles when extracts are prepared by grinding in liquid nitrogen and subjected to a further hour-long centrifugation. Figure 1D shows a representative profile for Flag5-tagged Tra1 or Tra1-F3744A in haploid strains grown in YPD media. We observe a pronounced proteolytic product of approximately 300 kDa for Flag5-Tra1-F3744A (arrow, lanes 3-5) that is minimal for wild-type Flag5-Tra1 (lanes 1 and 2), suggesting that the F3744A change may alter the conformation of Tra1. This proteolytic fragment was reduced in a strain containing a plasmid copy of SUP3 (lanes 6-8).

Identification of SUP3 and SUPB as alleles of TTI2: To identify the SUP3 mutation, the suppressor containing strain was backcrossed with the parent CY4350 (or CY4351) nine times, each time selecting for spore colonies that grew in ethanol and at 37°. After the ninth backcross, genomic DNA was isolated from three strains: an ethanol/37° resistant spore colony and a sensitive spore colony from the same tetrad and an ethanol/37° resistant spore colony from a distinct tetrad. Libraries were prepared and genomic sequencing performed using the ABI SOLiD 4.0 platform at the Centre for Applied Genomics at The Hospital for Sick Children (Toronto, Canada). The sequencing was performed in a single lane, multiplexing with 12
additional unrelated samples. Approximately 50 million reads were obtained for each sample, 60% of which aligned to the reference genome from the *Saccharomyces* Genome Database. Four single nucleotide differences were shared between the 37°C/ethanol resistant strains and not found in the sensitive strain. Only one of these differences, a T-C change within codon 328 of *TTI2*, was found upon manual inspection of individual sequence reads. This T-C transition converts amino acid residue Phe328 to Ser (Figure 2A), a position highly conserved in the fungal Tti2 proteins (Figure 2B; a closely related region is not apparent in human Tti2, C8orf41). As shown by the western blot in Figure 2C, the F328S mutation did not alter expression of Tti2.

Two approaches were used to demonstrate that suppression of *tra1-F3744A* was the result of *tti2-F328S*. As shown in Figure 3A, *SUP3* is partially dominant, with suppression being slightly less in a heterozygous diploid as compared to a haploid strain. We compared this effect with the addition of wild-type *TTI2* to CY4350 (*tra1-F3744A*) on a centromeric plasmid (Figure 3B). As was found for the heterozygous diploid, addition of the *TTI2* containing plasmid partially, but not completely, reversed the effect of *tti2-F328S*. Second, we analyzed eight independent spore colonies, 6 resistant and 2 sensitive, from a cross of CY5667 (*tti2-F328 tra1-F3744A*) and CY4350 (*TTI2 tra1-F3744A*). The *TTI2* allele from each spore colony was isolated by PCR and sequenced. For each of the eight alleles, *tti2-F328S* was found in all of the resistant strains and none of the sensitive strains. As the characteristics of the independently derived *SUPB* and *SUP3* strains were highly similar, having identified *tti2-F328S* as *SUP3*, we sequenced the *TTI2* allele in a derivative of the original *SUPB* strain that had been backcrossed versus its parent seven-times. One mutation was observed, an A-T at the first position of codon 336, converting isoleucine 336 to phenylalanine. The *TTI2* allele from four spore colonies was
sequenced; the *tti2-I336F* mutation segregated 2:2 with the suppressor phenotype.

Tti2 is a component of the Tra1-containing ASTRA complex and associates with Tel1 and Tor1 in part of what may be a chaperone complex with Tel2 and Tti1 (Shevchenko *et al.*, 2008; Takai *et al.*, 2010; Hurov *et al.*, 2010; Helmlinger *et al.*, 2011). To better understand the interactions of the *tti2* mutations with *TRA1*, we examined the growth of single and double mutant strains under a variety of conditions (Figure 4; also see Supplemental Figure 2). Both the *tti2-F328S* and *I336F* mutations suppressed the effects of *tra1-F3744A* under all of the conditions examined. Interestingly, this included reversal of the enhanced growth resulting from *tra1-F3744A* seen in media containing geneticin (G418), and the slow growth due to the DNA damaging agents MMS and phleomycin. Also of note the *tti2-F328S* allele in isolation resulted in minimal phenotypes; the *tti2-F328S* strain had a very slight reduction in growth in media depleted of phosphate.

**tti2-F328S suppresses transcriptional defects due to *tra1-F3744A***: To determine if the *tti2-F328S* allele restores the transcriptional competency of strains containing *tra1-F3744A*, we performed β-galactosidase assays with a *PHO5-lacZ* promoter fusion (Figure 5A). The *tra1-F3744A* allele decreases expression of *PHO5-lacZ* approximately 5-fold as compared to wild-type *TRA1*. *tti2-F328S* suppressed this effect, resulting in expression that was 80% of wild-type. *tti2-F328S* in the context of otherwise wild-type *TRA1* had only a marginal effect on *PHO5-lacZ* expression. *GAL10, HIS4*, and *RPL35a lacZ*-reporter fusions were also analyzed (Figure 5B). Similar to *PHO5, tra1-F3744A* decreased activated expression of *GAL10-lacZ* and expression of *HIS4-lacZ* approximately three-fold. In both cases *tti2-F328S* restored expression to
approximately wild-type levels. In contrast, the \textit{RPL35a} promoter was relatively unaffected by \textit{tra1-F3744A} or \textit{tti2-F328S}.

We examined the effects of \textit{Tra1-F3744A} on histone H4 acetylation (K8) as a ratio of the total histone H3 at the \textit{PHO5} and \textit{PGK1} promoters. Figure 6A shows the AcH4/H3 ratio for yeast strains CY4350 (\textit{tra1-F3744A TTI2}) and CY5667 (\textit{tra1-F3744A tti2-F328S}) as a percentage of that found for the wild-type strain CY4353 (\textit{TRA1 TTI2}). Since histone H4 acetylation of \textit{PHO5} is required prior to induction (Nourani \textit{et al.}, 2004), the chromatin immunoprecipitation was performed for cells grown in YPD media. The ratio of acetylated histone H4 to total histone H3 at \textit{PHO5} was reduced to ~40\% in the \textit{tra1-F3744A} strain. In comparison the ratio was ~90\% of wild-type at \textit{PGK1}. The ~40\% decrease for \textit{PHO5} is somewhat less than the 4-fold reduction seen upon deletion of the NuA4 component Eaf1 (Auger \textit{et al.}, 2008). Histone H4 acetylation returned to near wild-type levels in the presence of \textit{tti2-F328S}. Figure 6B shows the analysis for acetylated histone H3 (K18) at \textit{PHO5} for cells grown in low phosphate media (Nourani \textit{et al.}, 2004). In contrast to histone H4 acetylation, \textit{Tra1-F3744A} had virtually no effect on histone H3 acetylation (AcH3/total H3). Under the same conditions, deletion of Ada2, a direct regulator of Gcn5, reduced histone H3 acetylation by approximately 20-fold.

\textbf{Allele specificity of \textit{tti2-F328S}:} We examined the ability of \textit{tti2-F328S} to suppress deletions within the genes of other components of the SAGA and NuA4 complexes. Similar to the effect of \textit{tra1-F3744A}, deletions of \textit{ada2} or \textit{spt7} result in slow growth on media containing ethanol. However, unlike \textit{tra1-F3744A}, slow growth caused by \textit{ada2} and \textit{spt7} was not suppressed by
tti2-F328S (Figure 7A). Disruption of the gene encoding the NuA4 component Eaf3 results in slow growth in media containing Calcofluor white plus staurosporine (Figure 7B). Slow growth of an eaf7 disruption is observed for cells grown in 6% ethanol at 35°C. Neither of these phenotypes was suppressed by tti2-F328S (Figure 7B). As a strain with a disruption of YNG2 was not available in an isogenic background, suppression by tti2-F328S was analyzed by taking advantage of its dominant nature. YCplac111-tti2-F328S was transformed into QY202 (yng2Δ; kindly supplied by Jacques Côtè) and QY202 (YNG2). As shown in Figure 7C, the plasmid copy of tti2-F328S did not suppress slow growth of the yng2Δ strain at 30°C. We also analyzed the ability of tti2-F328S to suppress a second tra1 FATC domain mutation, tra1-L3733A, as well as a triple alanine scanning mutation of residues 3413-3315 (tra1-SRR3413; Mutiu et al., 2007) within the PI3K domain. As shown in Figure 7D, tti2-F328S partially suppressed the temperature sensitivity and slow growth in ethanol seen for the tra1-L3733A strain, though not as efficiently as for tra1-F3744A (compare to Figure 7A). In contrast, tti2-F328S did not suppress the slow growth of the tra1-SRR3413 strain at 37°C, but rather augmented the slow growth phenotype (Figure 7E).

**Localization of Tti2 and Tti2-F328S:** We expressed N-terminally eGFP-tagged Tti2 and Tti2-F328S in BY4741 to determine if the F328S mutation altered its localization (Figure 8). The wild-type molecule was found throughout the cell with both cytoplasmic and nuclear localization. When stressed with 6% ethanol, the distribution was relatively unchanged though some foci were observed. The proximity to the vacuole suggests that these may be late endosomes. The localization of eGFP-Tti2-F328S was almost identical to the wild-type protein. In media containing 6% ethanol there was a slight reduction in vacuolar proximal foci, but this
was variable.

**Localization of Tra1 and Tra1-F3744A:** We engineered yeast strains containing N-terminally eGFP-tagged Tra1 and Tra1-F3744A to examine their localization (Figure 9A). To avoid complications of slow growth due to the *tra1-F3744A* allele, the analysis was performed in heterozygous diploid strains with an untagged wild-type copy of Tra1. When grown in synthetic complete media wild-type eGFP-Tra1 was almost exclusively in the nucleus. Tra1-F3744A was found in the nucleus, but punctate fluorescence was also apparent in the cytoplasm. The amount of the cytoplasmic eGFP-Tra1-F3744A was reduced in the heterozygous *tti2-F328S/TTI2* strain. In media containing 6% ethanol (Figure 9B) Tra1 was more disperse but the majority of the protein remained in foci, which we suggest are nuclear (DAPI staining was ineffective in this media). In ethanol containing media, eGFP-Tra1-F3744A was diffusively distributed throughout the cytoplasm. Again, this appeared partially reversed in the *tti2-F328S/TTI2* strain. We used imaging software to quantify the fluorescent intensity of GFP-Tra1 and GFP-Tra1-F3744A per unit area in the nucleus as compared to the whole cell (Table 3). For cells grown in ethanol we assumed that the most pronounced focus was the nucleus. This quantification agrees with the visual conclusions drawn from the images of Figure 9 that Tra1-F3744A is more pronounced in the cytoplasm and partially relocalized to the nucleus by *tti2-F328S*.

To address the nature of the foci in which Tra1-F3744A is found, we visualized eGFP-Tra1-F3744A in strains containing RFP-tagged membrane constituents (Huh et al., 2003; strains kindly provided by Peter Arvidson). Figure 10A shows the analysis with RFP-tagged Anp1 (Golgi apparatus), Sec13 (ER to Golgi vesicles) and Nic96 (nuclear periphery) in diploid strains
(eGFG-tra1-F3744A/TRA1), when the cells were grown in media containing 6% ethanol. Of these the closest overlap was seen with RFP-Anp1; however, precise colocalization with any one type of membrane was not evident, including additional analyses with Cop1, Pex3 and Snf7 (Supplemental Figure 3). For comparison in Figure 10B we show the localization of eGFP-Tra1-F3744A with the RFP-tagged proteins in synthetic complete media (no ethanol). As shown above, eGFP-Tra1-F3744A was more evident in the nucleus than in the presence of ethanol, but cytoplasmic eGFP-Tra1-F3744A was still apparent.

If Tti2 has a role in the folding and/or stabilization of Tra1, strains with defects in the TTT complex may have reduced levels of Tra1. We introduced a tel2-15 temperature sensitive allele (Stirling et al., 2011; Grandin et al., 2012) into a strain expressing Flag5-Tra1, and examined Tra1 levels after growth at 30°, 35° and 37° (Figure 11A). The level of Flag5-Tra1 was unchanged by tel2-15 at 30°, but substantially reduced at the two elevated temperatures. We also examined the phenotypes of strains containing tel2-15, a temperature sensitive tti2 allele (tti2-1, Stirling et al., 2011), or double mutations of these alleles in combination with tra1-F3744A (Figure 11B). The recessive tel2-15 and tti2-1, alleles in an otherwise wild-type background resulted in slow growth on media containing 6% ethanol at 30°. Under all of the conditions, tel2-15 resulted in synthetic slow growth in combination with tra1-F3744A. A slight synthetic slow growth phenotype was evident for the tti2-1 tra1-F3744A strain at 30°.

**tra1-R3590I suppresses tra1-F3744A**: Two recent analyses of PIKK structure suggest that the FATC domain may interact with and regulate the kinase domain (Lempiäinen and Halazonetis, 2009; Sturgill and Hall, 2009). To perhaps provide support for such a model, we sequenced the
tra1 allele 3' of base 9730 in one of the intragenic suppressors of tra1-F3744A. A transversion of G to T at base 10769 that converts arginine 3590 to isoleucine was found. Alignments position R3590 in the putative activation loop, between β sheet 10 and α helix 7 corresponding to PI3K-γ (Figure 12A; Walker et al., 1999). To verify that R3590I is responsible for the suppression, the mutation was integrated into CY4398 and a haploid spore colony isolated after sporulation. The growth of this strain (CY5640; tra1-R3590I-F3744A) at 37°, and on media containing ethanol or rapamycin confirmed that the R3590I mutation conferred suppression (Figure 12B). Similar to tti2-F328S, the F3590I mutation in isolation had no apparent phenotype.
DISCUSSION

The FATC domain is essential for the function of Tra1 and other PIKK family members (Beamish et al. 2000; Hoke et al. 2010; Priestly et al. 1998; Sun et al. 2005; Takahashi et al. 2000). For Tra1 this is apparent from the fact that a protein containing an additional C-terminal glycine residue will not support viability (Hoke et al., 2010). Altering the terminal phenylalanine of Tra1 to alanine is less severe, but results in slow growth in rich media at 30° and under conditions of stress. We have shown that alleles of TTI2 suppress tra1-F3744A. tti2-F328S restored all of the measured properties of the strains to ~80% of the wild-type level. Suppression by tti2-F328S was specific for mutations in the FATC domain of Tra1; tra1-L3733A was suppressed, whereas alleles altering the PI3K domain, or other SAGA or NuA4 components were not. The suppression by alleles of TTI2, whose product with Tel2 and Tti1 is proposed to act as a chaperone (Horejsi et al., 2010; Hurov et al., 2010; Kaizuka et al., 2010; Takai et al., 2010), the reduced levels of Tra1 in the tel2-15 strain, and the increased number of proteolytic products seen after western blotting Tra1-F3744A lead us to suggest that the FATC domain is important for Tra1 to acquire or stabilize a fully functional conformation. The finding that the F3744A mutation increased levels of cytoplasmic Tra1 is consistent with this model, or alternatively for roles of the FATC domain and Tti2 in protein trafficking. Loss of any of these possible roles would deplete functional Tra1, and would be expected to act broadly given the importance of independent Tra1 (Helmlinger et al., 2011), as well as the SAGA and NuA4 complexes; indeed we find numerous phenotypic consequences of tra1-F3744A.
The eGFP-Tra1-F3744A present in the cytoplasm was not uniformly distributed, but appeared in foci. Though not specific for any one membrane type, we propose that these foci represent Tra1-F3744A associated with membranes, and that the altered FATC domain potentially traps these molecules on the membranes. In turn this finding predicts that the folding of Tra1 and perhaps the formation of some of its multisubunit complexes may occur on membranes. This is appealing because the membrane would provide a platform for the process to occur, and perhaps protect the C-terminal domains from proteolysis. A requirement for membrane interactions provides a rationale for the large number of synthetic interactions observed between membrane trafficking components and either tral-SRR\textsubscript{3413} or deletions of NuA4 component genes (Hoke \textit{et al.}, 2008a; Mitchell \textit{et al.}, 2008). Membrane interactions are also consistent with the lipid binding properties of some of the SAGA components (Hoke \textit{et al.}, 2008b). In the event of reduced complex formation the molecules could be targeted to the vacuole, perhaps providing an explanation for the Pep4 dependent cleavage of Spt7 (Spedale \textit{et al.}, 2010). Interestingly, Han and Emr (2011) have recently shown that Cti1 and Tup1 assemble with Cyc8 on late endosomal membranes, mediated through their binding of phosphatidylinositol-3,5-diphosphate. Lipid binding is required for nuclear import of Cti8-Tup1-Cyc8, interaction with SAGA and activation of galactose-regulated genes. Membranes are inherently sensitive to many environmental cues. As Han and Emr (2011) suggest the membrane assembly of transcriptional complexes provides a tight link with the environmental state.

Our results in combination with the association of Tra1 and Tti2 determined by Shevchenko \textit{et al.} (2008) clearly indicate a functional relationship between these molecules. The connection between Tel2, Tti2 and Tti1 demonstrated in mammalian cells suggests that the TTT
complex is also functionally associated with Tra1 (Hayashi et al., 2007; Takai et al., 2007, 2010; Hurov et al., 2010; Kaizuka et al., 2010). How this relates to other components of the ASTRA complex is less clear. In that it contains Rvb1 and Rvb2, ASTRA resembles an assembly of the R2TP (Huen et al., 2009) and TTT complexes with Tra1, similar to that seen for mTOR (Horejsi et al., 2010). The potential transient nature of ASTRA and its possible role in the folding/stability of Tra1 agrees with it not yet being isolated as an intact biochemical entity. Alternatively the suppression by Tti2 may take place in the context of an independent TTT complex, with ASTRA required for additional functions.

tti2-F328S acts in a partially dominant fashion to suppress tra1-F3744A. The suppression was most notable with tti2-F328S as the sole copy of the gene, but was still apparent in the context of the wild-type allele. With the specific mechanism of the TTT complex unknown, we can only speculate on how Tti2-F328S and Tti2-I336S act. A strict gain of function is possible, but perhaps less so given the two alleles and the partial dominance. Alternatively, the two mutations may disrupt an interaction or property of Tti2 that otherwise results in its inhibition. The dominant nature of the allele also suggested that the FATC domain of Tra1 might interact closely with the region of Tti2 surrounding F328. Tra1-F3744A may be unable to interact, and suppression result from the restored interaction with Tti2-F328S. The possibility of a direct interaction was attractive given that a hydrophobic contact for the wild-type proteins could be replaced by a hydrogen bond between the serine of Tti2-F328S and the C-terminus of Tra1. However, additional experiments were inconsistent with a direct contact. First this model would not easily explain suppression of tra1-L3733A, or the ability of tti2-I336F to suppress. Second, if the domain of Tti2 were to directly contact the FATC domain, one might expect negative effects
on the other FATC domain containing proteins. The only discernable phenotype of *tti2-F328* in isolation was a slight slow growth in media depleted of phosphate. Tti2-F328S did not lead to sensitivity to the DNA damaging agents MMS or phleomycin, suggesting a minimal effect on Mec1 and Tel1. Finally, we were unable to detect an interaction between a fragment of Tra1 containing the PI3K and FATC domains with the C-terminal half of Tti2 using recombinant proteins. We conclude that Tti2-F328S enhances the activity of Tra1-F3744A, likely by affecting folding, through a mechanism that does not restore interaction between the molecules nor involve increased levels of Tti2. We note also that *tti2-F328S* does not suppress a mutation converting the terminal tryptophan of Mec1 to alanine (Supplemental Figure 4). This suggests either that folding of the FATC domain of Mec1 does not require Tti2 function or that the change in function of Tti2-F328S is specific for Tra1.

Expression of the NuA4 (Nourani et al., 2004) and SAGA-regulated (Gregory et al., 1998) *PHO5* promoter was reduced approximately 5-fold by *tra1-F3744A*. Consistent with an effect of this mutation on NuA4 function, Tra1-F3744A reduced histone H4 acetylation of the *PHO5* promoter. In contrast, and despite Gcn5 being required for activated expression, the F3744A mutation had little effect on histone H3 acetylation at the *PHO5* promoter. Since the breadth of the phenotypes attributable to *tra1-F3744A* suggest that SAGA function is altered, we propose that the lack of change in histone H3 acetylation is due to the ability of the Ada complex (Eberharter et al., 1999), including Gcn5, Ada2 and Ngg1, to act independently of SAGA. These results with Tra1-F3744A are similar to what we observe at *PHO5* upon deletion of Spt7: partially reduced acetylation, significantly decreased expression (DD and CJB, in preparation). The lack of correlation between the importance of these molecules to *PHO5* expression and their
effect on acetylation suggests that specific targeting of *PHO5* acetylation by SAGA is required for expression.

Our study is a direct demonstration of a functional link between Tra1 and Tti2. This link is likely the result of a role for Tti2, as part of the TTT complex, in the folding/maturation of Tra1 as has been found for other PIKK proteins (Takai et al., 2007, 2010; Horejsí et al., 2010; Hurov et al., 2010; Kaizuka et al., 2010; Stirling et al., 2011). In addition, our study points to a putative role for the FATC domain in the regulated folding/maturation of Tra1 in the cytoplasm. We can not exclude that there are additional roles for the FATC domain. Recent models for the structure of the C-terminal domains of the PIKK family members predict that helical FATC domain interacts and regulate the kinase domain (Lempiäinen and Halazonetis, 2009; Sturgill and Hall, 2009). The suppression of *tra1-F3744A* by mutation of arginine 3590 to isoleucine in the putative activation loop supports such an interaction. Interestingly, the models by Lempiäinen and Halazonetis (2009) and Sturgill and Hall (2009) place the C-terminus in proximity to the active site where the FATC domain could have a role in catalysis.
ACKNOWLEDGEMENTS

We would like to thank Megan Davey and Stephen Hoke for critically reading the manuscript; Aaron Simkovich, Lance DaSilva, Esther Rosenthal and Brittany Rasmussen for technical assistance, Peter Arvidson for providing RFP tagged strains; Bernard Duncker, Philip Hieter, Peter Stirling, Fred Winston and Jacques Côtè for yeast strains; Patricia Kane for helpful suggestions; Joe Mymryk for yeast plasmids; and Ivan Sadowski for sharing sequencing lanes. This work was supported by CIHR grant MOP10845 to CJB. DD was supported by an Ontario Graduate Studentship. SK was supported by a National Science and Engineering Research Council of Canada Studentship.
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FIGURE 1. Isolation of extragenic suppressors of tra1-F3744A. A. Phenotypes resulting from tra1-F3744A. Yeast strains CY4350 (tra1-F3744A) and CY4353 (TRA1) containing YCplac33-YHR100C (included since the HIS3 allele used to select for TRA1 integration disrupts the terminal 11 codons of YHR100C) were grown to stationary phase in media lacking uracil. Ten-fold serial dilutions were spotted onto YP containing 2% glucose (YPD) or 2% galactose (galactose), or YPD with 1.0 μg/mL tunicamycin, 1.2 M NaCl, 60 μg/ml brefeldin, or 1.0 M sorbitol. YPD media was also depleted of phosphate (Han et al., 1988) or made to pH 8.0 with the addition of sodium hydroxide. Plates were grown at 30°. B. Suppression of tra1-F3774A by alleles of tti2. Cultures of yeast strains CY4353 (TRA1), CY4350 (tra1-F3744A), CY5667 (tra1-F3744A tti2-F328S; SUP3), and CY5842 (tra1-F3744A tti2-I336F; SUPB) containing YCplac33-YHR100C were serially diluted and spotted onto a YPD plate or YPD containing 6% ethanol and grown at 30° or 37°. Note that SUP3 and SUPB are tti2-F328S and tti2-I336F, respectively. C. Yeast strains CY5912 (lanes 1 and 4; Flag5-tra1-F3744A/tra1-F37443A tti2-F328S/tti2-F328S; SUP3 mutant), CY4419 (lanes 2 and 5; Flag5-TRA1/TRA1 TTI2/TTI2; SUP3 WT) and CY4421 (lanes 3 and 6; Flag5-tra1-F3744A/TRA1 TTI2/TTI2; SUP3 WT) were grown to stationary phase in minimal media depleted of uracil, then diluted 1/20 into YPD and grown for 8 hr at 30°. 50 μg or 15 μg of protein extract prepared by grinding with glass beads was separated by SDS-PAGE (5%) and Western blotted with anti-Flag antibody (top panel) or stained with Coomassie Brilliant Blue (CBB stain, bottom panel). D. Yeast strains CY6072 (lanes 1 and 2; Flag5-TRA1 YCplac111-TTI2), CY6074 (lanes 3-5; Flag5-tra1-F3744A YCplac111-TTI2), and CY6083 (lanes 6-8; Flag5-tra1-F3744A YCplac111-tti2-F328S) were
grown in YPD to an optical density at 600 nm of ~2.0. Protein extracts were prepared by grinding in liquid nitrogen and clarified by centrifugation. Serial dilutions of each were separated by electrophoresis, and Flag5-tagged Tra1 or Tra1-F3744A detected by Western blotting. The proteolytic product of ~300 kD is indicated by an arrow.

FIGURE 2. Alleles of Tti2 suppress tra1-F3744A. A. Sequence of the suppressor tti2 alleles. B. Multiple sequence alignment of Tti2 from a variety of fungal species. The two alterations found in the suppressor strains are indicated by asterisks. The alignment was performed with MUSCLE (Edgar, 2004). In order the proteins are: *Schizosaccharomyces pombe*, NP_596623.3; *Penicillium chrysogenum*, XP_002568898.1; *Candida glabrata*, XP_449362.2; *Nakaseomyces delphensis*, CAO98781.1; *Kluyveromyces lactis*, XP_451713.1; *Lachancea* thermotolerans, XP_002555374.1; *Vanderwaltozyma polyspora*, XP_001643571.1; *Zygosaccharomyces rouxii*, XP_002495325.1; *Ashbya gossypii*, NP_982872.1; *Saccharomyces cerevisiae*, NP_012670.1. C. Yeast strain BY4743 containing YCplac111 (lane 1), YCplac111-myc9-TTI2 (lanes 2 and 3), or YCplac111-myc9-tti2-F328S (lanes 4 and 5) were grown to stationary in minimal media depleted of leucine, then diluted 1/20 into YPD and grown for 8 hr at 30°. 20 or 50 μg of protein from each were separated by SDS-PAGE (10%) and Western blotted with anti-myc antibody (top panel) or stained with Coomassie Brilliant Blue (CBB, bottom panel).

FIGURE 3. tti2-F328S acts as a partial dominant allele. A. Genomic complementation. The haploid yeast strains CY4350 (tra1-F3744A TTI2) and CY5667 (tra1-F3744A TTI2-F328S), and diploid strains homozygous for tra1-F3744A and either homozygous or heterozygous for TTI2-F328S were grown to stationary phase and serial dilutions spotted onto YPD plates and grown at
30 or 37°, or YPD containing 6% ethanol and grown at 30°. B. Plasmid complementation. CY5667 (1 and 3) and CY4350 (2 and 4) were transformed with YCplac111-myc<sup>9</sup>-TTI2 or vector alone. Serial dilutions of the strains were spotted on the plates indicated.

FIGURE 4. Suppression of tra1-F3744A phenotypes by tti2-F328S and tti2-I336F. Yeast strains CY4353 (TRA1 TTI2), CY4350 (tra1-F3744A TTI2), CY5667 (tra1-F3744A tti2-F328S), CY5843 (tra1-F3744A tti2-I336F), CY5665 (TRA1 tti2-F328S), or a mec2-1 strain (Weinert et al., 1994; included to verify the phleomycin and MMS plates) were grown to stationary phase diluted 1/10<sup>4</sup> and spotted onto selection plates as follows: YPD at 30°, YPD at 37°; YPD at 30° containing 0.03% methyl methanesulfonate 0.03% (MMS), 1.0 μg/mL phleomycin (Phleo), YPD depleted of phosphate (Phosph-), 7.5 μg/mL Calcofluor White (CW), 6% ethanol, 1.0 μg/mL tunicamycin, or 20 μg/mL geneticin (G418); YP containing 2% galactose, and YPD at pH 8.0. *Note that the plating on G418 was performed with no dilution of the cells.

FIGURE 5. Expression of PHO5, GAL10, HIS4 and RPL35a promoter fusions in tti2-F328S strains. A. Yeast strains CY5667, CY4350, CY4353 and CY5665 with the indicated TRA1 and TTI2 alleles, were transformed with a LEU2 centromeric plasmid containing a PHO5-lacZ fusion, grown to stationary phase in media depleted of leucine, washed three-times with water, diluted into YPD media depleted of phosphate, grown for 16 hr at 30° and β-galactosidase activity determined, normalizing to cell density. The error bars indicated represent one standard deviation from the mean. B. GAL10, HIS4, and RPL35a promoters were analyzed as above in strains CY5667, CY4350 and CY4353. For GAL10, initial cultures were grown in raffinose containing media and shifted to 2% galactose. HIS4 analysis was performed in media depleted of
histidine. *RPL35a* was analyzed in YPD media.

**FIGURE 6. Histone H3 and H4 acetylation at the PHO5 promoter.** A. Histone H4 acetylation. Yeast strains CY4353 (*TRA1*), CY4350 (*tra1-F3744A*) and CY5667 (*tra1-F37744A tti2-F328S*) were grown to $A_{600}=1.5$ in YPD media. ChIP analysis was performed using antibodies against acetylated K8 of histone H4 and against histone H3. Levels of immunoprecipitated *PHO5* and *PGK1* promoters were determined after PCR, separation of products by electrophoresis and ethidium bromide staining. Serial dilutions were examined to ensure analysis in a linear range. The histogram shows the ratio of acetylated histone to total histone H3 (the mean for 2 independent experiments) for CY4350 and CY5667 as a percentage of the ratio found for the wild-type strain CY4353 (values for CY4353 are set at 100% and are not shown). B. Histone H3 acetylation. Yeast strains CY4353 (*TRA1*), CY4350 (*tra1-F3744A*), CY5667 (*tra1-F37744A tti2-F328S*), and BY4282 (*ada2Δ*) were grown to $A_{600}=1.5$ in YPD media depleted of phosphate. ChIP analysis was performed using antibodies against acetylated K18 of histone H3 and against histone H3. The histogram shows the average ratio of acetylated histone to total histone H3 for CY4350 and CY5667 as a percentage of the wild-type (CY4353) for an experiment performed in duplicate.

**FIGURE 7. Allele specificity of tti2-F328S suppression.** A. SAGA components. Serial dilutions of yeast strains CY4353 (*TRA1 TTI2*), CY4350 (*tra1-F3744A TTI2*), CY5667 (*tra1-F37744A tti2-F328S*), BY4282 (*ada2Δ TTI2*), CY5876 (*ada2Δ tti2-F328S*), CY5873 (*spt7Δ TTI2*) and CY5915 (*spt7Δ tti2-F328S*) were spotted onto YPD (grown at 37°C) or YPD with 6% ethanol (grown at 30°C). The *spt7* deletion strains were grown on a separate plate and for an additional
day. 

B. NuA4 components. Yeast strains BY4742 (WT), BY7143 (eaf3Δ TTI2), CY5916 (eaf3Δ tti2-F328S), BY2940 (eaf7Δ TTI2) and CY5917 (eaf7Δ tti2-F328S) were spotted onto either YPD containing 1 μg/mL staurosoprine plus 7.5 μg/mL Calcofluor White (grown at 30°) or 6% ethanol (grown at 35°). 

C. yng2Δ. QY204 (YNG2) and QY202 (yng2Δ0) were transformed with YCplac111-tti2-F328S or YCplac111. Cultures were grown in media lacking leucine, and serial dilutions spotted onto YPD plates at 30°. 

D. tra1-L3733A. Yeast strains CY4353, CY4350, CY4057 (tral-L3733A TTI2) and CY5658 (tral-L3733A tti2-F328S) were spotted onto YPD at 37° and YPD containing 6% ethanol at 30°. 

E. tra1-SRR3413. Yeast strains CY4353, CY2222 (tral-SRR3413 TTI2), CY5665 (TRA1 tti2-F328S), and CY6137 (tral-SRR3413 tti2-F328S) were spotted onto YPD at 30° and 37°.

FIGURE 8. Localization of Tti2 and Tti2-F328S. 

A. BY4741 containing YCplac111-eGFP-TTI2 or eGFP-tti2-F328S were grown in synthetic complete (SC) media to late-log phase, stained with DAPI and visualized by fluorescence microscopy. A 10 μm scale bar is shown in the bottom right. 

B. The above strains were grown to stationary phase in SC media, diluted 1:4 in SC media containing 8% ethanol, grown a further 18 hr, and visualized by fluorescence microscopy. BF, bright field.

FIGURE 9. Localization of Tra1 and Tra1-F3744A. 

A. Yeast strains CY6029 (eGFP-TRA1/TRA1 TTI2/TTI2), CY6025 (eGFP-tra1-F3744A/TRA1 TTI2/TTI2) and CY6063 (eGFP-tra1-F3744A/TRA1 tti2-F328S/TTI2) were grown in synthetic complete media to mid-log phase stained with DAPI and visualized by fluorescence microscopy (SC). 

B. The two rightmost panels
are strains grown in SC containing 6% ethanol. BF, bright field. A 10 μm scale bar is shown in the bottom right.

FIGURE 10. Localization of eGFP-Tra1-F3744A with RFP-tagged Anp1, Sec13 and Nic9. A. Localization in SC media containing ethanol. Diploid strains containing a single copy of each tagged allele were grown to stationary phase in SC media, diluted 1:4 in SC containing 8% ethanol, grown a further 18 hr, and visualized by fluorescence microscopy. A 10 μm scale bar is shown in the bottom right. B. Localization in SC media. Columns three and 5 from the left are merged images between eGFP (green) and RFP or DAPI, respectively. A 10 μm scale bar is shown in the bottom right.

FIGURE 11. Interaction of tra1-F3744A with temperature sensitive alleles of components of the TTT complex. A. Tel2 is required for the expression or stability of Tra1. Strains CY5919 (Flag5-TRAI TEL2, WT) and CY6141 (Flag5-TRAI tel2-15, ts) were grown to stationary phase at 30°, then diluted 20-fold into YPD and grown for 8 hours at 30°, 35°, or 37°. Whole cell extracts were prepared by bead lysis, The indicated amount of extract (60 or 20 μg) was separated by SDS PAGE (5%). The upper half of the gel was Western blotted with anti-Flag antibody; the lower half was stained with Coomassie Brilliant Blue (CBB). B. Growth of tra1-F3744A and tel2-15 or tti2-1 double mutant strains. CY4353 (wild-type), CY4350 (tra1-F3744A), PSY42 (tel2-15), PSY561 (tti2-1), CY6148 (tra1-F3744A tel2-15) and CY6146 (tra1-F3744A tti2-1) were grown to stationary phase at 25°, then serial dilutions plated onto YPD at 25° or 30°, or
YPD plus 6% ethanol at 30°.

FIGURE 12. *tra1-R3590I* suppresses *tra1-F3744A*. A. PI3K domain sequences of Tra1 (top) and porcine PI3K-γ (bottom) are aligned (SMART, Ponting *et al.*, 1999) with structural features of porcine PI3K-γ (Walker *et al.*, 1999) shown below. Arginine 3590 is underlined.

B. Cultures of yeast strains CY5920 (*TRA1*), CY5828 (*tra1-F3744A*), CY5640 (*tra1-R3590I, F3744A*), and CY5639 (*tra1-R3590I*) were serially diluted and spotted onto YPD (grown at 30° or 37°, 1 day), or YPD containing 6% ethanol or 1 nM rapamycin (2 days).
SUPPLEMENTAL FIGURE LEGENDS

Supplemental Figure 1. Sequence of the molecule to integrate Flag\textsuperscript{5}-tag TRA1. The ATG translational start preceding the tag is in bold. A genomic Hind\textsuperscript{III} fragment encoding \textit{URA3} (~1.1 kbp) was inserted at the underlined Hind\textsuperscript{III} site. TRA1 sequences in frame with the \textit{NotI} site (Saleh et al., 1998) were cloned 3’ to the \textit{NotI}.

Supplemental Figure 2. Suppression of \textit{tra1-F3744A} phenotypes by \textit{tti2-F328S} and \textit{tti2-I336F}. Yeast strains CY4353 (\textit{TRA1 TTI2}), CY4350 (\textit{tra1-F3744A TTI2}), CY5667 (\textit{tra1-F3744A tti2-F328S}), CY5843 (\textit{tra1-F3744A tti2-I336F}), CY5665 (\textit{TRA1 tti2-F328S}), or a \textit{mec2-1} strain (Weinert \textit{et al.}, 1994) were grown to stationary phase diluted 1/10\texttextsuperscript{4} and serial dilutions spotted onto selection plates as follows: YPD at 30°, YPD at 37°; YPD at 30° containing 0.03% methyl methanesulfonate 0.03% (MMS), 1.0 μg/mL phleomycin, YPD depleted of phosphate, 7.5 μg/mL Calcofluor white, 6% ethanol, or 1.0 μg/mL tunicamycin; YP containing 2% galactose, and YPD at pH 8.0. Note that some images are composites from two otherwise identical plates.

Supplemental Figure 3: Localization of eGFP-Tra1-F3744A with RFP-tagged \textit{Cop1}, \textit{Snf7} and \textit{pex3} in SC media containing ethanol. Diploid strains containing a single copy of each tagged allele were grown to stationary phase in SC media, diluted 1:4 in SC containing 8% ethanol, grown a further 18 hr, and visualized by fluorescence microscopy. A 10 μm scale bar is shown in the bottom right.

Supplemental Figure 4: \textit{tti2-F328S} does not suppress the temperature sensitivity of \textit{mec1-}
W2368A. A. pCB2317 for converting tryptophan 2368 of Mec1 to alanine. The altered codon is underlined. *HIS3* was inserted at the *BamHI* site. A *BclI* site was placed downstream of the stop codon to identify the allele. B. *mec1-W2368A-HIS3* was integrated into a diploid strain heterozygous for *tti2-F328S* (CY6045). The *TTI2* allele of spore colonies growing on media depleted of histidine were sequenced. CY6071 and CY6072 were *TTI2*, CY6078 was *tti2-F328S*. The strains were streaked onto a YPD plate and grown at 37° for 3 days.
Table 1. Strains used in this study.

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<th>Genotype</th>
<th>Reference</th>
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<td><strong>MATα his3Δ1/his3Δ1 leu2Δ0/leu2Δ0 LYS2/lys2Δ0 met15Δ0/MET15 ura3Δ0/ura3Δ0</strong></td>
<td>Winzeler and Davis (1997)</td>
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<td>Winzeler and Davis (1997)</td>
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<td>Winzeler and Davis (1997)</td>
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<td>BY4282</td>
<td>Isogenic to BY4741 except ada2::KanMX</td>
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<td>Winzeler and Davis (1997)</td>
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<td>FY1093</td>
<td><strong>MATα spt7-402::LEU2 his4-917- lys2-173R2 leu2-1 ura3-52 trp1-63 ade8</strong></td>
<td>Gansheroff et al. (1995)</td>
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<td>CY2222</td>
<td><strong>MATα can1Δ::STE2pr-SpHIS5 lyp1Δ his3Δ1 leu2Δ0 ura3Δ0 met10 LYS2+ tra1-SRR3413-URA3</strong></td>
<td>Hoke et al. (2008a)</td>
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<td>Hoke et al. (2010)</td>
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<td>CY5912</td>
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CY5917  MATa ura3Δ0 his3Δ0 leu2Δ0 tti2-F328S eaf7::KanMX  This work
CY5919  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-Flag2::TRA1  This work
CY5920  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-Flag2::TRA1::HIS3  This work
CY5928  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-Flag2::tra1-F3744A::HIS3  This work
CY5998  Isogenic to BY4742 except URA3-eGFP-TRA1  This work
CY5999  MATa ura3Δ0 his3Δ0 leu2Δ0 tti2-F328S URA3-eGFP-tra1-F3744A::HIS3  This work
CY6016  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-eGFP-tra1-F3744A::HIS3  This work
CY6018  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-eGFP-tra1-F3744A::HIS3  This work
CY6025  Diploid of CY6016 x BY4741  This work
CY6029  Diploid of CY5998 x BY4741  This work
CY6063  Diploid of CY5999 x BY4741  This work
CY6072  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-Flag2::TRA1
       YCplac111-TTI2  This work
CY6074  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-Flag2::tra1-F3744A::HIS3
       YCplac111-TTI2  This work
CY6083  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-Flag2::tra1-F3744A::HIS3
       YCplac111-tti2-F328S  This work
CY6137  MATa his3Δ1 leu2Δ0 ura3Δ0 tra1-SRR3413-URA3
       tti2-F328S  This work
CY6141  MATa ura3Δ0 his3Δ0 leu2Δ0 URA3-Flag2::TRA1 tel2-15::KanMX
       YCplac111-TTI2  This work
QY204   MATa his3Δ200 trp1Δ63 ura3-52 leu2Δ1 lys2-1265  Nourani et al. (2001)
QY202   MATa his3Δ200 trp1Δ63 ura3-52 leu2Δ1 lys2-1265
       yng2:: KanMX  Nourani et al. (2001)
PSY36   MATa leu2Δ0 his3Δ0 lys2Δ0 asa1-1::URA3  Stirling et al. (2011)
PSY42   MATa ura3Δ0 leu2Δ0 his3Δ0 lys2Δ0 tel2-15::KanMX  Stirling et al. (2011)
PSY43   MATa ura3Δ0 leu2Δ0 his3Δ0 lys2Δ0 tel2-15::KanMX  Stirling et al. (2011)
PSY561  MATa leu2Δ0 his3Δ0 tti2-1::URA3  Stirling et al. (2011)
PSY625  MATa leu2Δ0 his3Δ0 lys2Δ0 tti1-1::URA3  Stirling et al. (2011)
Table 2. Oligonucleotides used in this study.

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<td>5927-2</td>
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Table 3. Relative concentrations (fluorescence intensity per unit area) of eGFP-Tra1 (wild-type or F3744A) in the nucleus versus total cell.

<table>
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<th>Tra1/Tti2 (strain)</th>
<th>[Nuclear eGFP-Tra1] / [Total Cell eGFP-Tra1]¹</th>
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<tbody>
<tr>
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<td>SC Media</td>
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<tr>
<td>WT / WT (CY6029)</td>
<td>4.0 ± 0.6</td>
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<tr>
<td>F3744A / WT (CY6025)</td>
<td>2.7 ± 0.5</td>
</tr>
<tr>
<td>F3744A / F328S (CY6063)</td>
<td>3.5 ± 0.4</td>
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¹ Concentrations represent the intensity of eGFP fluorescence per unit area in the nucleus divided by the eGFP fluorescence per unit area for the cell (including the nucleus). The numbers represent the average for 20 cells. As DAPI staining was ineffective for the ethanol grown cells, the most intense focus was assigned as the nucleus.
FIGURE 1.
FIGURE 2.

S. pombe 352 NVSYLPYLTLTLDELCRIFQONPQIIQPSLSSLFLEILTKLLE
P. chrys 438 TTKYLOELIPVLFT----TLSNPFCTAHPOLLLAAATQAVIDK
C. glabr 306 STIYQRIVYDGOX----LKDPFTAFDKLIFEQTELVEAIG
N. delph 300 SIALLQRIDMGQX----LKDPFTAFEPMLHEVTCTIETYTIQ
K. lactis 316 TTYLORVIYVLGEF----YFRNAFTMLQMPILHKCLDLILD
L. thermo 308 SVVHLORVFIFGIDX----ITRNAFLFMPFVQVLSTLTHLVS
V. polyp 300 SVIIHLSLIFNLDY----VIRNPFTAFPSIMNETMLRIQTLID
Z. rouxi 306 SVYHLRMIYVLWGY----LVRNPFHTAFDSIMDLTLSLHMTLV
A. gossi 308 SVYHLQRIYVLWGY----IVRNPFHTLFPVLVEKTLDTAKLAE
S. cerev 308 SVYHLQRIYVLWGY----IVRNPFHTFPKLISKTLSVSTK

C
myc^a-Ti2       -              wt             F328S

\( \mu g \)
\( \alpha \) myc

\( \text{CBB} \)

-75 kD
FIGURE 3.
FIGURE 4.
FIGURE 5.
FIGURE 6.
FIGURE 7.
FIGURE 8.
FIGURE 9.
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<td>RFP-Sec13</td>
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FIGURE 10
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<td>Flag^5</td>
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**B**

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**FIGURE 11**
A  
Tra1 3578 SGNVFTEMLPSRFPYERVKPLKKNHDLSPDBSPHNNNEPVPRFLTPNIQ 3629
PI3Kγ 957 TGNLFHDFGhilGNKSFQLGI-------------NEPVPRFLTPDFG 996

\[ \beta_{10} \quad \text{Activation loop} \quad \alpha_7 \]

B  

<table>
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FIGURE 12.