Yeast *src* Homology Region 3 Domain-binding Proteins Involved in Bud Formation

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Abstract. The yeast protein Bem1p, which bears two src homology region 3 (SH3) domains, is involved in cell polarization. A Rho-type GTPase, Rho3p, is involved in the maintenance of cell polarity for bud formation, and the rho3 defect is suppressed by a high dose of BEM1. Mutational analysis revealed that the second SH3 domain from the NH₂ terminus (SH3-2) of Bem1p is important for the functions of Bem1p in bud formation and in the suppression of the rho3 defect. Boi2p, which bound to SH3-2 of Bem1p, was identified using the two-hybrid system. Boi2p has a proline-rich sequence that is critical for displaying the Boi2p-Bem1p two-hybrid interaction, an SH3 domain in its NH₂-terminal half, and a pleckstrin homology domain in its COOH-terminal half. A BOI2 homologue, BOI1, was identified as a gene whose overexpression inhibited cell growth. Cells overexpressing either BOI1 or BOI2

URING bud formation in the yeast Saccharomyces cerevisiae, cell polarity is established for the initiation of bud emergence and it is maintained during bud growth. Patches of actin filaments become concentrated at the bud site, towards which the transport of secretory vesicles is directed for the construction of the daughter cell (Tkacz and Lampen, 1972; Field and Schekman, 1980; Pringle and Hartwell, 1981; Cabib et al., 1982; Adams and Pringle, 1984; Kilmartin and Adams, 1984; Novick and Botstein, 1985; Pringle et al., 1986; Drubin, 1991). The establishment and maintenance of cell polarity require the functions of Rho-type GTPases Cdc42p, Rho3p, and Rho4p, which belong to the Ras superfamily (Johnson and Pringle, 1990; Matsui and Toh-e, 1992a,b; Imai et al., 1996). GTPases of the Ras superfamily act as molecular switches through their conformational change between the GTP-bound active form and GDP-bound inactive form (Barbacid, 1987; Bourne et al., 1991; Boguski were arrested as large, round, and unbudded cells, indicating that the Boi proteins affect cell polarization. Genetic analysis revealed that BOI1 and BOI2 are functionally redundant and important for cell growth. $\Delta boil$ $\Delta boi2$ cells became large round cells or lysed with buds, displaying defects in bud formation and in the maintenance of cell polarity. Analysis using several truncated versions of BOI2 revealed that the COOH-terminal half, which contains the pleckstrin homology domain, is essential for the function of Boi2p in cell growth, while the NH₂-terminal half is not, and the NH₂-terminal half might be required for modulating the function of Bem1p. Overproduction of either Rho3p or the Rho3prelated GTPase Rho4p suppressed the boi defect. These results demonstrate that Rho3p GTPases and Boi proteins function in the maintenance of cell polarity for bud formation.

and McCormick, 1993). Defects in either *CDC42* or *CDC24*, which encodes a GTP-GDP exchange factor for Cdc42p, disrupt the asymmetric localization of actin filaments and cause cells to become unbudded, large, and round, an indication that Cdc42p and Cdc24p are essential for the establishment of cell polarity (Sloat and Pringle, 1978; Sloat et al., 1981; Adams et al., 1990; Johnson and Pringle, 1990; Zheng et al., 1994).

Defects in *RHO3* cause severe growth defects. Disruption of *RHO4*, which encodes a Rho3p-related GTPase, enhances the growth defect of $\Delta rho3$ cells (Matsui and Toh-e, 1992a). Temperature-sensitive *rho3* mutant cells lose cell polarity at nonpermissive temperatures: the asymmetric localization of actin filaments is disrupted in the *rho3* cells, and the *rho3* cells are arrested as large, round cells, although, in contrast to *cdc42* mutant cells, not all of these cells are arrested as unbudded cells (Imai et al., 1996). Depletion of both Rho3p and Rho4p results in lysis of cells that have small buds (Matsui and Toh-e, 1992b). These observations strongly suggest that Rho3p is required for the maintenance of cell polarity for bud growth.

The rho3 defect is suppressed by the overexpression of *BEM1*, an indication that Bem1p has functions that affect

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the Rho3 pathway (Matsui and Toh-e, 1992b). BEM1 encodes a protein with two copies of the src homology region 3 (SH3)¹ domain (SH3-1 and SH3-2; see Fig. 1 A). Disruption of BEM1 results in temperature-sensitive growth. At nonpermissive temperatures, $\Delta bem1$ cells become unbudded, large, and round, with the loss of actin polarization, indicating that Bem1p is involved in cell polarization (Bender and Pringle, 1991; Chenevert et al., 1992). Bem1p can bind to Cdc24p independently of its SH3 domains (Peterson et al., 1994). Cdc24p binds to the Ras-type GTPase Rsr1p/Bud1p, which is required for the proper selection of bud sites (Chant and Herskowitz, 1991; Bender and Pringle, 1989; Zheng et al., 1995). These findings lead to the possibility that Bem1p is a component of the protein complex that is needed for bud emergence. Since SH3 domains mediate protein-protein interactions (e.g., Cicchetti et al., 1992), it is possible that Bem1p recruits another component to the complex via its SH3 domains.

To clarify the role of the SH3 domains of Bem1p, we characterized several *bem1* mutants that were defective in these domains. Using the two-hybrid system, we identified Boi2p as a protein that bound to the SH3 domain and was important for the function of Bem1p. In addition, Boi1p, which was functionally redundant with Boi2p, was identified by its inhibitory effect on bud emergence. Our genetic and morphological studies indicate that the function of Boi protein is related to the Rho3 pathway and is important for bud growth. We present a model in which Boi proteins and Rho3p are involved in the modulation of the Bem1p-containing complex for bud growth.

Materials and Methods

Microbiological Techniques

Rich medium containing glucose (YPD), synthetic minimal medium (SD), and synthetic complete medium (SC) are as described (Sherman et al., 1986). YPGal and SCGal are YPD and SC, respectively, except that 2% glucose is replaced with 5% galactose and 0.3% sucrose. YPGal0.1 is YPGal, except that the concentration of galactose is 0.1%. SC-U and SCGal-U are SC and SCGal, respectively, without uracil. SC-L and SCGal-L are SC and SCGal, respectively, without leucine. SC-UT and SCGal-UT are SC and SCGal, respectively, without uracil and tryptophan. Yeast transformations were performed by the method of Ito et al. (1983).

Strains and Plasmids

The yeast strains used are listed in Table I. Plasmid pBTM116 is a high copy number plasmid that harbors the 2- μ m DNA origin *TRP1* and the sequence for the lexA DNA-binding domain. pGAD424 is a high copy number plasmid that harbors the 2- μ m DNA origin *LEU2* and the sequence for the Gal4 *trans*-activation (Gal4 TA) domain (Fields and Sternglanz, 1994). Plasmid pRS316-RHO3 carries a 1.8-kb KpnI-XhoI fragment that encompasses the *RHO3* gene on pRS316, a low copy number plasmid harboring *URA3* (Sikorski and Hieter, 1989). Plasmid pKT10 is a high copy number plasmid that carries the 2- μ m DNA origin *URA3*, the *TDH3* promoter before a unique EcoRI site, termination codons downstream of the EcoRI site, and the *TDH3* terminator (Tanaka et al., 1988). pOPR3 and pOPR1 are pKT10-based plasmids that carry the coding region of *RHO3* and the coding region of *RHO1*, respectively, under the control of the TDH3 promoter (Matsui and Toh-e, 1992b). Plasmid pOPR4 is a high copy number plasmid that carries the coding region of

RHO4 under the control of the PYK1 promoter (Matsui and Toh-e, 1992a). Plasmid pKT10mycN was constructed by inserting the sequence for the initiator methionine, a myc epitope tag (EQKLISEEDL), and a multicloning site into the EcoRI site of pKT10. The KpnI-KpnI, BamHI-KpnI, HpaI-KpnI, and KpnI-Dral fragments carrying the BEM1 sequence (see Fig. 1 A) were inserted in frame into the multicloning site of pKT10mycN to generate pBEM1KK, pBEM1 Δ 1, pBEM1 Δ 1 Δ 2, and pBEM1 C, respectively. The BamHI-Sall fragment carrying the sequence for SH3-2 was removed from pBEM1KK and pBEM1 ΔC to create pBEM1 $\Delta 2$ and pBEM1 $\Delta 2\Delta C$, respectively, and was religated in frame after blunting the overhangs. BEM1 in pBEM1KK lacked the sequence for the 45 NH2-terminal amino acids but was able to serve as a multicopy suppressor of rho3 (see Fig. 1 B, sector 2). The DNA sequences of the fragments, which were derived from the PCR (Saiki et al., 1988) or oligonucleotide-directed mutagenesis and used for generating plasmids, were determined to confirm precise replication during each procedure. Nucleotide sequences were determined by the method of Sanger et al. (1977).

Construction of bem1 Mutants with Mutations in the SH3 Domain

The 2.3-kb fragment, derived from pSRO1 (Matsui and Toh-e, 1992b), from the BamHI site in the BEMI coding region to the BamHI site in the 3' noncoding region, was inserted into a derivative of pBluescriptII that had been constructed from pBluescriptII KS⁺ (Stratagene, La Jolla, CA) by removal of the sequence between the EcoRV and XhoI sites. The 1.1-kb HindIII-HindIII fragment carrying URA3 was inserted into the HindIII site (in the 3' noncoding region of BEMI) of the resultant plasmid to generate YIpUBEM1C. The 0.6-kb fragment, derived from pSRO1, from the Smal site in the 5'-coding region of BEM1 to the Sall site in the sequence for SH3-2, was inserted between the SmaI site and SaII site (in the sequence for SH3-2) of YIpUBEM1C to generate YIpUBEM1. Proline 123 in SH3-1 and proline 208 in SH3-2 were replaced with leucine to generate *bem1*^{Leu123} and *bem1*^{Leu208}, respectively, with an oligonucleotide-directed in vitro mutagenesis system (Amersham Corp., Arlington Heights, IL) using the primers 5'-AAAATAGGTCTTAAGGACAACACC (for bem1^{Leu123}), 5'-AAACCCAACTAGTACAAGGCCG (for bem1^{Leu208}), and YIpUBEM1 as the template. For the construction of *bem1-\DeltaSH3s*, a truncated version of bem1 without the sequence for both SH3-1 and SH3-2, the fragment from the SmaI site in the 5' noncoding region to the KpnI site in the coding region was inserted in frame between the SmaI and SaII sites of YIPUBEM1C after blunting the overhangs to generate YIPUBEM1\DeltaSH3s, which lacked the sequence between the KpnI and SalI sites of YIpUBEM1. For the construction of *bem1-\DeltaC*, the 1.0-kb SalI-HindIII fragment from pBEM1 Δ C, which contained the BEM1 sequence between the SalI and Dral sites, the termination codons after the Dral cleavage site, and the TDH3 terminator, was inserted between the Sall and HpaI sites of YIPUBEM1 after blunting the overhangs of the HindIII cleavage site. For the replacement of the wild-type BEM1 allele with a mutant allele, each derivative of YIpUBEM1, digested with SmaI and XbaI, was introduced into the cells. Replacement was confirmed by PCR.

Isolation of BOI2 and Assay of the Two-hybrid Interaction

The construction of plasmids for the lexA DNA-binding domain fused with Bem1p (lexA-Bem1p) was performed as follows. An EcoRI site was introduced before the initiator methionine codon of BEM1 by oligonucleotide-directed in vitro mutagenesis. The EcoRI-HpaI fragment of BEM1 was inserted into the multicloning site of pBTM116 in frame to generate plexSHs. The SalI-SalI fragment that contained the sequence for the COOH-terminal Bem1p and the 3' noncoding region of BEM1 from pBEM1KK was inserted between the Sall sites of plexSHs to create plexBEM1. Yeast strain L40, with two reporter genes (lexA-lacZ and lexA-HIS3), was transformed with plexBEM1 and then with a yeast genomic library in which yeast genomic DNA was expressed as fusion proteins with the Gal4 TA domain (Chien et al., 1991). The transformants were streaked on SD+3AT plates, which are SD plates containing 100 µg/ ml adenine sulfate and 40 mM 3-aminotriazole, an inhibitor of imidazoleglycerolphosphate dehydratase (His3p). Plasmids were recovered from colonies that formed on plates after incubation at 25°C for 1 wk. The recovered plasmids were reintroduced into the L40 strain harboring plexBEM1 to examine whether the plasmids could induce the expression of the reporter genes. HIS3 expression was assessed by the formation of L40 cell colonies on SD+3AT plates after incubation at 25°C for 1 wk.

^{1.} Abbreviations used in this paper: a.a., amino acid; GST, glutathione S-transferase; PH, pleckstrin homology; SC, synthetic complete medium; SD, synthetic minimal medium; SH3, *src* homology region 3; YPD, rich medium containing glucose.

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Table I. Yeast Strains Used in this Study

Strain	Genotype	Reference or source
W303	MATa/MATa ura3/ura3 leu2/leu2 his3/his3 trp1/trp1 lys2/lys2 ade2/ade2	Sutton et al. (1991)
W303-1A	MATa ura3 leu2 his3 trp1 lys2 ade2	segregant of W303
YMR505	MATa ura3 leu2 trp1 lys2 ade2 Δrho3::LEU2 pGAL7:RHO4	Matsui and Toh-e (1992b)
YMS1194	MATa ura3 leu2 his3 trp1 lys2 ade2 bem1 ^{Leu123}	This study
YMS1199	MATa ura3 leu2 his3 trp1 lys2 ade2 bem1 ^{Leu208}	This study
YMS1190	MATa ura3 leu2 his3 trp1 lys2 ade2 bem1 ^{Leu123, Leu208}	This study
YMS1201	MATa ura3 leu2 his3 trp1 lys2 ade2 bem1- ΔC	This study
YMS1183	MATa ura3 leu2 his3 trp1 lys2 ade2 bem1-ΔSH3s	This study
YMB101	MATa ura3 leu2 his3 trp1 lys2 ade2 Δboi1::HIS3	This study
YMB102	MAT α ura3 leu2 his3 trp1 lys2 ade2 Δ boi1::HIS3	This study
YMB201	MATa ura3 leu2 his3 trp1 lys2 ade2 Δboi2::URA3	This study
YMB202	MATa ura3 leu2 his3 trp1 lys2 ade2 Δ boi2::LEU2	YMB202 transfromed with a <i>ura3</i> -disruption plasmid
YMB1201	MATa/MATα ura3/ura3 leu2/leu2 his3/his3 trp1/trp1 lys2/lys2 ade2/ade2 Δboi1::HIS3/+ +/Δboi2::LEU2	$YMB102 \times YMB202$
YMB1207	MATa leu2 his3 trp1 lys2 ade2 Δboi1::HIS3 Δboi2::LEU2 pGAL7:BOI2	segregant of YMB1201 transformed with YIpUGAL7BOI2.
YMB1202	MATa leu2 his3 trp1 lys2 ade2 Δboi1::HIS3 Δboi2::LEU2 BOI2ΔN:URA3	segregant of YMB1201 transformed with pRS306-BOI2ΔN
DRA4	MATa/MAT $lpha$ leu2/leu2 ura3/ura3 trp1/+ +/ade6 can1/+ cyh/+ GAL ⁺ /GAL ⁺	derived from siblings of a cross between X2180-1B (from Yeast Genetic Stock Center) and AM1205-9B (Mitchell and Herskowitz, 1986)
KA31	MATa ura3 leu2 his3 trp1 lys2	Irie et al. (1991)
L40	MATa ade2 his3 leu2 trp1 lexA-lacZ:URA3 lexA-HIS3:LYS2 gal4 gal80	R. Sternglanz (Stony Brook)

All of the strains listed above, with the exception of YMR505, DRA4, KA31, and L40, are in the W303 background.

lacZ expression was examined by measuring the activity of β -galactosidase in L40 cells with the method described (Miller, 1972). DNA clones carrying *BO12* were isolated from a yeast genomic library (Matsui and Toh-e, 1992a) in λ ZAPII (Stratagene) by hybridization using the *BO12* fragment isolated by the two-hybrid method as a probe. pSKBO12 was a plasmid derived from the yeast genomic library and carried a 6-kb fragment encompassing *BO12* in pBluescript SK⁻ (Stratagene).

Construction of Plasmids for the Two-hybrid Interaction Assay

Plasmids for truncated versions of lexA-Bem1p were constructed as follows. plex Δ SHs was created by removing the EcoRI-SalI (in the sequence for SH3-2) fragment of plexBEM1, with partial digestion by SalI. plex ΔC was constructed by removing the PstI-PstI fragment of plexBEM1. plexs2, plex-s1, and plex-s12 were constructed by inserting BanI-PstI fragments carrying bem1^{Leu208}, bem1^{Leu123}, and bem1^{Leu123, Leu208}, respectively, into the multicloning site of pBTM116. To create plexBEM1 Δ C, the 1.0-kb Sall-HindIII fragment from pBEM1 DC was inserted between the Sall site (in the BEM1 coding region) and the PstI site (in the multicloning site of pBTM116) of plexSHs after blunting the overhangs of PstI and HindIII cleavage sites. Plasmids for truncated versions of Gal4 TA fused with Boi2p (Gal4TA-Boi2p) were constructed as follows. MluI-BglII, MluI-EcoRI, MluI-AatII, MluI-BanII, and MluI-XbaI fragments from pGADBOI2, a primary isolate carrying BO12 (see below), were inserted between the MluI site and the multicloning site of pGAD424 (Fields and Sternglanz, 1994) to create pGADBG2, pGADR1, pGADAA1, pGADBA2, and pGADXB1, respectively. BstEII-AatII and BanII-EcoRI fragments from pGADBOI2 were inserted into the multicloning site of pGAD424 in frame to create pGADE2A2 and pGADB2R1, respectively. pGADΔPRO was constructed as follows. An AatII site was introduced downstream of the 463th codon of BOI2 in pSKBOI2 by oligonucleotide-directed in vitro mutagenesis, and the resultant plasmid was digested with AatII and religated in frame to generate pSKBOI2 Δ PRO. The BOI2 gene in pSKBOI2 Δ PRO lacked the sequence between the introduced AatII site and the original AatII site for the region (a.a. 436-464) that contained the proline-rich sequence (see below). The 2.6-kb XbaI-SalI fragment carrying the BOI2 coding region from pSKBOI2ΔPRO was inserted between the XbaI and Sall sites of pGADR1 to create pGADAPRO. The 1.8-kb EcoRI-BamHI fragment carrying the sequence for the COOH-terminal half of Cdc24p (a.a. 167-end) was inserted between the EcoRI and BamHI sites of pGAD424 to create pGADCDC24.

In Vitro Binding Assay

The NciI-EcoRI fragment encoding Boi2p (amino acids [a.a.] 67-545) was inserted into the multicloning site downstream of the sequence for glutathione S-transferase (GST) in pGEXKG. Boi2p (a.a. 67-545) fused with GST (GST-Boi2p) was produced by use of this construct and affinity purified with glutathione-agarose beads (Sigma Immunochemicals, St. Louis, MO) as described (Shirayama et al., 1994). About 10⁸ yeast cells (wildtype strain KA31) producing myc-tagged Bem1p were washed twice in PBS. The cell pellet was disrupted by blending with glass beads in 400 μl of lysis buffer (50 mM Tris-HCl, pH 7.5, 5 mM EGTA, 2% [vol/vol] Triton X-100, 75 mM NaCl, 0.5 mM PMSF) and was clarified by centrifugation at 14,000 g for 15 min at 4°C. 30 µl of the supernatant (as the total cell lysate) was resolved by SDS-PAGE, and 160 µl of the supernatant was mixed with 50 µl GSH-agarose beads with 2 µg immobilized GST-Boi2p (or GST for control) and incubated at 4°C for 3 h. The beads were then washed extensively in washing buffer (50 mM Tris-HCl, pH 7.5, 0.1% [vol/ vol] Triton X-100, 10% [vol/vol] glycerol). Bound proteins were eluted with 100 µl elution buffer (50 mM Tris-HCl, pH 9.0, 20 mM glutathione) and 20 µl of the eluate were resolved by SDS-PAGE.

Isolation of BOI1

YEp51B, which carries *LEU2*, the 2- μ m DNA origin, and the *GAL10* promoter, was constructed by removing a 0.27-kb BamHI-Sall fragment adjacent to the *GAL10* promoter from YEp51 (Broach et al., 1983). Diploid strain DRA4 was transformed with a yeast genomic DNA library, based on YEp51B, in which yeast genomic DNAs were expressed under the control of the *GAL10* promoter. The resultant transformants that grew well on SC-L, but not on SCGal-L (which contained 2% galactose, instead of 5% galactose and 0.3% sucrose), were selected at 28°C, and the plasmids were recovered (details of the screening will be described elsewhere). pGA68, one of the isolates, carried *BO11* downstream of the *GAL10* promoter. The *BO11* gene in pGA68 was expressed from the internal methionine at position 378, as judged from a comparison of the nucleotide sequences of pGA68 and *BO11* (Bender et al., 1996).

BOI2 Expression Plasmids and Disruption of BOI1 and BOI2

YIPUGAL7 is a YIP plasmid that carries the GAL7 promoter, URA3, and the BgIII and Sall sites downstream of the GAL7 promoter (Matsui and

Toh-e, 1992b). A BamHI site was introduced before the initiator methionine codon of BOI2 by oligonucleotide-directed in vitro mutagenesis, and the BglII site in the 3' noncoding region of BOI2 was replaced with a SalI site by insertion of a Sall linker (Takara, Tokyo, Japan). The BamHI-Sall fragment carrying the complete BOI2 coding region was inserted between the BglII and SalI sites of YIpUGAL7 to create YIpUGAL7BOI2. In this construct, designated pGAL7:BOI2, BOI2 was expressed under the control of the GAL7 promoter. YIpUGAL7BOI2 was digested with BgIII and religated after blunting the overhangs to create YIpUGAL7BOI2 ΔC . From this construct, designated pGAL7:BOI24C, Boi2p without its COOH-terminal half (a.a. 697-end) was produced under the control of the GAL7 promoter. Synthetic oligonucleotides containing the initiator methionine codon and the XhoI and BgIII sites were inserted into the BgIII site of YIpUGAL7 to create YIpUGAL7f. The 2.1-kb AatII-SalI fragment containing the sequence for the COOH-terminal half of Boi2p (a.a. 466-end) was inserted between the XhoI and SalI sites of YIpUGAL7f after blunting the overhangs of the XhoI and AatII cleavage sites to create YIpUGAL7BOI2ΔN. From this construct, designated pGAL7:BOI2 ΔN, Boi2p without its NH2-terminal half was produced under the control of the GAL7 promoter. An ~6-kb SacII-SalI fragment carrying BOI2 from pSKBOI2 was inserted into pYO324, a high copy number plasmid that carries 2-µm DNA origin and TRP1 (Ohya et al., 1991), and into pRS314, a low copy number plasmid that carries CEN6 and TRP1 (Sikorski and Hieter, 1989), to create pYO324-BOI2 and pRS314-BOI2, respectively. An AatII site was introduced downstream of the 9th codon of BOI2 in pSKBOI2 by oligonucleotide-directed in vitro mutagenesis, and the resultant plasmid was digested with AatII and religated in frame to create pSKBOI2ΔN. This construct (BOI2ΔN) encodes Boi2p without its NH₂-terminal half (a.a. 10-464). An ~5-kb SacII-SalI fragment carrying BOI2 AN was inserted into pYO324 and pRS306, a YIp plasmid carrying URA3 (Sikorski and Hieter, 1989), to create pYO324-BOI2ΔN and pRS306-BOI2ΔN, respectively. pYO324-BOI2 and pRS314-BOI2 were digested with BgIII and religated after blunting the overhangs to create pYO324-BOI2 Δ C and pRS314-BOI2 Δ C, respectively. From these constructs, designated BOI2AC, Boi2p without its COOH-terminal half (a.a. 697-end) was produced. The YIpUGAL7- and pRS306-derived plasmids were digested with StuI and then introduced into cells for targeted integration at the ura3 locus.

The BOI2 sequence between the PmII site (in the 5' noncoding region) and the SpeI site of pSKBOI2 was replaced with the 1.1-kb URA3 fragment to create pBOI2 Δ . In this construct, the sequence from the position 443 bp upstream of the initiation codon to the 829th codon of BOI2 was deleted. The plasmid for the disruption of BOI1 was constructed as follows. The 2.8-kb SalI-HindIII fragment from the insert of pGA68 was inserted into pBluescriptII KS⁺ to create pKSBOI1. To create pRS306-BOI1, the 2.2-kb BamHI-BamHI fragment that carried the 3'-half of the BOII gene from pGA68 was inserted into the BamHI site of pRS306. To clone the 5'-half of BOII, the yeast genomic DNA in which pRS306-BOI1 had been integrated at the BOII locus was digested with EcoRV, religated, and introduced into Escherichia coli. With the recovered plasmid as a template, a 840-bp fragment carrying the 5' noncoding region of BOII and the sequence for the NH2-terminal portion (a.a. 1-87) of Boi1p was amplified by PCR. In the PCR, the primer was designed to replace the 87th codon of BOII in the amplified fragment with a stop codon, TAG. The amplified fragment was inserted into the BamHI site of pKSBOI1 to create pKSBOI1-2, which carried a 3.3-kb insert encompassing a version of BOII in which the sequences for a.a. 91-374 were deleted. A 1.8-kb BamHI fragment carrying HIS3 was inserted into the BglII site in the BOII coding region of pKSBOII-2 to create pBOI1Δ. In this construct, the BOII gene was disrupted with the termination codon at the 87th codon, with the deletion of the sequence for a.a. 91-374 and with the insertion of HIS3. pBOI1Δ and pBOI2Δ were digested with PvuII and introduced into the cells by replacement transformation to disrupt BOI1 and BOI2, respectively. These replacements did not disrupt any open reading frames other than BOII and BOI2, as judging from nucleotide sequences. The disruptions were confirmed by Southern analysis and PCR.

Cell Lysis Assay

Cell lysis was assayed by monitoring the leakage of alkaline phosphatase, a yeast intracellular protein, into the culture medium by the method described (Paravicini et al., 1992). Cells were streaked on YPD plates. After incubation, the plates were overlaid with a solution of BCIP (10 mM 5-bromo-4-chloro-3-indolyl phosphate [Sigma], 0.1 M Tris-HCl, pH 9.5, and 1% agar) and incubated for 1 h at 37°C.

Detection of myc Epitope-tagged Protein

Proteins containing the myc epitope tag were detected by Western blotting analysis with anti-myc antibodies as described (Yoko-o et al., 1995).

Morphological Observations

Cells were stained with rhodamine-phalloidin to reveal actin filaments as described (Pringle et al., 1989). Cells were fixed with 5% formaldehyde for 10 min, washed with PBS, stained with 1:50 diluted rhodamine-phalloidin solution (Molecular Probes, Inc., Eugene, OR) for 2 h, and washed five times with PBS. The sample was then mounted in *p*-phenylenediamine (1 mg/ml in 90% glycerol) and observed under an epifluorophotomicroscope (BH-2; Olympus, Tokyo, Japan). Dead cells were stained with methylene blue (Rose, 1975). A solution of methylene blue (0.02% methylene blue, 2% sodium citrate) was mixed with an equal volume of cell culture, and the cells were observed immediately with Nomarski optics.

Results

Bem 1p Domains Required for Suppression of the $\Delta rho3$ Defect

Overexpression of BEM1 suppresses the growth defect caused by the disruption of RHO3 (Matsui and Toh-e, 1992b; Fig. 1 B, sector 2). To examine whether the SH3 domains of Bem1p are necessary for the suppression or not, we constructed truncated BEM1 sequences (shown diagrammatically in Fig. 1A) and analyzed the suppression activities of the various constructs. Arho3 cells that overexpressed BEM1 without the sequence for SH3-1 grew as well as $\Delta rho3$ cells that overexpressed *BEM1* (Fig. 1 *B*, right, sectors 2 and 3), indicating that SH3-1 is not required for the ability of BEM1 to serve as a multicopy suppressor of *rho3*. However, $\Delta rho3$ cells that overexpressed BEM1 without the sequence for SH3-2 (Fig. 1 B, right, sectors 4, 5, and 7) or the COOH terminus of 35 a.a. (Fig. 1 B, right, sectors 6 and 7) formed colonies as tiny as those of $\Delta rho3$ cells that harbored a control plasmid (Fig. 1 B; right, sector 8), indicating that these truncated versions of BEM1 did not suppress the Arho3 defect. The deletion of SH3-2 or of the COOH-terminal sequence did not reduce the amount of the myc-tagged Bem1p, as judged by Western analysis (data not shown). These results indicate that SH3-2 and the COOH-terminal sequence are critical if BEM1 is to function as a multicopy suppressor of rho3.

The Effects of Substitutions in the SH3 Domains and of Deletion of the COOH-terminal Region of Bem1p on Cell Growth

The replacement of the conserved proline residue by leucine destroys the function of an SH3 domain in the sem5 protein of *Caenorhabditis elegans* (Rozakis-Adcock et al., 1992). We introduced the corresponding mutation into SH3-1 (*bem1*^{Leu123}) and SH3-2 (*bem1*^{Leu208}). The *bem1*^{Leu208} cells and *bem1*^{Leu123}, Leu208</sup> cells displayed temperature-sensitive growth (Fig. 1 C, sectors 3 and 4). At 37°C, the cells were arrested as large, round, unbudded cells, and the asymmetric organization of actin filaments was disrupted in these cells (data not shown). These phenotypes are similar to those of the *Δbem1* cells. By contrast, *bem1*^{Leu123} cells grew as well as wild-type cells, even at elevated temperatures (Fig. 1 C, sector 2). These results indicate that SH3-2 is important for the function of Bem1p in bud emergence.



B





Figure 1. The importance of SH3-2 in the functions of Bem1p. (A) The constructs used for overexpression of the various versions of BEM1 are shown schematically below the map of BEM1. Open box, the coding region of BEM1; hatched boxes, the regions encoding the SH3 domains (SH3-1 and SH3-2); black boxes, the sequence that encodes the myc epitope tag; lines, the BEM1 sequences that are under the control of the TDH3 promoter (narrow lines indicate deleted regions). The ability of each plasmid to suppress the growth defect of $\Delta rho3$ cells (shown in B) is indicated. K, KpnI; B, BamHI; S, SalI; H, HpaI; and D, DraI. (B) Cells of the Arho3 pGAL7:RHO4 strain YMR505 carrying the indicated plasmids were incubated on YPD (right) and SCGal-U (left) plates at 30°C for 3 d. The Arho3 strain YMR505, which carries pGAL7:RHO4, RHO4 under the control of the GAL7 promoter, grows very poorly on glucosecontaining medium but grows well on galactose-containing medium because RHO4 can serve as a multicopy suppressor of rho3 (Matsui and Toh-e, 1992a,b). Cells harbored pRS316-RHO3, a centromeric plasmid carrying RHO3 (sector 1), pBEM1KK (sector 2), pBEM1 Δ 1 (sector 3), pBEM1 Δ 2 (sector 4), pBEM1 $\Delta 1\Delta 2$ (sector 5), pBEM1 Δ C (sector 6), pBEM1 Δ 2 Δ C (sector 7), and a dummy plasmid, pKT10mycN (sector 8) as a negative control. (C) Wildtype cells (strain W303-1A, sector 1), bem1^{Leu123} cells (strain YMS1194, sector 2), bem1^{Leu208} cells (strain YMS1199, sector 3), bem1^{Leu123,Leu208} cells (strain YMS1190, sector 4), and bem1- ΔC (strain YMS1201, sector 5) were streaked and incubated on YPD plates for 3 d at 25°C (left) and at 37°C (right).

The COOH-terminal 35 a.a. of Bem1p is essential for the suppression of the $\Delta rho3$ defect (Fig. 1 B). We examined whether the COOH-terminal 35 a.a. is also important for the function of Bem1p in cell growth or not. We replaced the wild-type *BEM1* allele with a truncated allele that lacked the sequence for the COOH-terminal 35 a.a. (*bem1-* ΔC). The *bem1-* ΔC cells displayed temperaturesensitive growth (Fig. 1 C, sector 5) and were arrested as large, round, unbudded cells at 37°C (data not shown). These results indicate that the COOH-terminal 35 a.a. is important for the function of Bem1p in bud emergence. It has been reported that the COOH-terminal half of Bem1p interacts with the COOH-terminal half of Cdc24p (Peterson et al., 1994). We examined whether the COOH-terminal 35 a.a. of Bem1p is required for the Cdc24p-Bem1p interaction using two-hybrid system (Fields and Sternglanz, 1994). The lexA-Bem1p fusion (from plexBEM1) increased the level of expression of *lacZ* in L40 cells (*lexA-lacZ* strain), with the increase depending on the presence of the Gal4TA-Cdc24p fusion from pGADCDC24. The lexA-fused Bem1p without the COOH terminus of 35 a.a. (from plexBEM1 Δ C), however, did not (Table II). These results indicate that the

Table II. Assay of Two-hybrid Interaction between Cdc24p and Bem1p

	pGAD424	pGADCDC24
pBTM116	0.4	0.7
plexBEM1	7	54
plexBEM∆C	4	0.5

 $\beta\text{-}Galactosidase$ activities (units) of L40 cells that harbored plasmids in the indicated combinations are shown.

COOH-terminal 35 a.a. of Bem1p is required for the twohybrid interaction between Cdc24p and Bem1p.

Isolation of BOI2

A screening was made for genes that encode the Bem1pbinding protein using the two-hybrid system. Plasmid plexBEM1, carrying the sequence encoding the lexA-Bem1p fusion and the yeast S. cerevisiae genomic library expressed as fusion proteins with the Gal4 TA domain (Chien et al., 1991), were introduced into L40 cells that carried lexA-lacZ and lexA-HIS3. The plasmids were then recovered from the colonies that formed on selective plates (SD+3AT). Among the plasmids recovered from the His⁺ transformants, only plasmid pGADBOI2 increased the level of expression of both HIS3 and lacZ in L40 cells in a plexBEM1-dependent manner. DNA clones were isolated from the yeast genomic library (Matsui and Toh-e, 1992a) with a fragment from pGADBOI2 as a probe. The nucleotide sequence of the clones revealed a gene that encoded a 1,040-a.a. protein with an SH3 domain, a proline-rich sequence, and a pleckstrin homology (PH) domain (Musacchio et al., 1993; Fig. 2). A homology search of the Genbank database using the FASTA program revealed a homologue with 38% identity. This homologous gene was identified on the basis of its twohybrid interaction with Bem1p (Bender et al., 1996). Both groups designated these genes BOI1 and BOI2 (as bem one interacting), and the gene on pGADBOI2 was designated BO12. Four domains of Boi2p, namely, domain I (residues 39-113) including an SH3 domain (residues 50-102), domain II (residues 266-331), domain III (residues 436-462) including a proline-rich sequence (residues 438-458), and domain IV (residues 731-943) including a PH domain (residues 767-891), were highly homologous to those of Boi1p (Fig. 2 B), with the extent of identity being 71, 65, 78, and 69%, respectively.

Domains That Are Required for the Two-hybrid Interaction between Bem1p and Boi2p

To identify the sequence responsible for the two-hybrid interaction between Bem1p and Boi2p, several constructs that encoded lexA-Bem1p fused proteins with truncation and/or mutations (shown diagrammatically in Fig. 3 A) were introduced into L40 cells that harbored pGADBOI2, and the two-hybrid interaction was examined. The *BEM1* sequence for the NH₂-terminal half (a.a. 1–325), which contained both SH3 domains, was sufficient for the interaction (Fig. 3 A, plex ΔC). Introduction of the bem1^{Leu208} mutation into this *BEM1* sequence, however, abolished the increase in the expression level of the reporter genes (Fig. 3 A, plex-s2 and plex-s12). By contrast, the plasmid

carrying only the $bem1^{Leu123}$ mutation increased the expression (Fig. 3 A, plex-s1). These results indicate that SH3-2 is essential for the Bem1p-Boi2p interaction.

To identify the Boi2p domain that participates in the two-hybrid interaction, we constructed several plasmids that encoded Gal4TA-Boi2p fused proteins, as shown diagramatically in Fig. 3 B. In the original isolate, pGADBOI2, the BOI2 sequence was ligated with the sequence for Gal4TA at the Sau3AI site that was located in the sequence for the SH3 domain of Boi2p, and the SH3 domain in the fusion was disrupted (see Figs. 2 A and 3 B). Thus, the SH3 domain of Boi2p was dispensable for the Boi2p-Bem1p interaction. The deletion of the COOH-terminal sequence of Boi2p (threonine 465-end) did not reduce the interaction (Fig. 3 B, pGADAA1), an indication that the NH₂-terminal half of Boi2p is sufficient for the interaction. More extensive deletion of the COOH-terminal sequence (alanine 442-end including the proline-rich sequence), however, abolished the interaction (Fig. 3 B, pGADBA2). The deletion of the sequence for a.a. 436-464, including the proline-rich sequence, also abolished the interaction (Fig. 3 B, pGADAPRO). Proline-rich sequences are reported to be the motif of SH3 domain-binding sites (e.g., Cicchetti et al., 1992). These results suggest that the interaction between Boi2p and Bem1p is mediated by the interaction between the proline-rich sequence of Boi2p and SH3-2 of Bem1p. Deletion of the NH₂-terminal sequence (a.a. 1-266), however, also abolished the interaction (Fig. 3 B, pGADE2A2). It is likely that the NH₂-terminal sequence affects the conformation of Boi2p, allowing the proline-rich sequence to bind efficiently or, alternatively, the NH2-terminal sequence might stabilize the Gal4-Boi2p fusion protein.

The Interaction of Bem1p and Boi2p In Vitro

The interaction of Boi2p with Bem1p was examined in vitro with purified GST-Boi2p. Yeast cells were transformed with the plasmids for myc-tagged Bem1p, shown diagrammatically in Fig. 1 A. The myc-tagged Bem1p in the lysates was detected with anti-myc antibodies (Fig. 3 C, *left*). The lysates containing myc-tagged Bem1p were incubated with GST-Boi2p that had been bound to GSH beads. The versions of Bem1p that possessed SH3-2 were coprecipitated with GST-Boi2p (Fig. 3 C, *right*, lanes 1, 2, and 5), whereas the versions of Bem1p that lacked SH3-2 only coprecipitated at very low levels, if at all, with GST-Boi2p (Fig. 3 C, *right*, lanes 3, 4, and 6). Both the two-hybrid experiments and the in vitro binding assays indicate that SH3-2 is critical for the Boi2p–Bem1p interaction and that SH3-1 is not essential for this interaction.

Phenotypes of Cells that Overexpressed BOI1 and BOI2

BO11 was identified during a screening for genes whose expression, under the control of the GAL10 promoter, was lethal to yeast cells. The isolated BO11 was fused to the GAL10 promoter at the Sau3AI site located in the sequence for codons 374–375 of BO11. In this fused gene, the 378th methionine codon is expected to be the initiation codon. Cells that carried the multicopy plasmid that harbored the BO11 sequence under the control of the GAL10promoter did not grow on SCGal-L, a galactose-containing selective medium, and they were arrested as large, Å



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IV are underlined.

Boi2p

Boilp

Boilp

Boi2p

Boilp

Boi2p 765:

656:

659:

773:

885:

893:

OOFTDNA

ATIDIDTSVE

Boi2p 1005: PGVARNSSMRGTEKKGKFSTEEDYFGDNSKHKTDKI

PEENPIT<mark>SMPSE</mark>

ATETIP

KNSSPIVDKKSSKKSRSKRRS

TDTRERGLIDITAHR

DEKERGLIDITAHEV

NEEDEDQF

--DAEEEEGRDQFGWDDT?

Figure 2. Amino acid sequence of Boi2p. (A) The nucleotide and deduced amino acid sequences of BOI2. The nucleotide sequence of BOI2 has been deposited in DDBJ/EMBL/Genbank under the accession No. D38310. The Sau3AI restriction site, namely, the site of the junction with the sequence for Gal4 TA in pGADBOI2, is indicated by a box. The restriction sites for BstEII, XbaI, HincII, BanII, AatII, EcoRI, and BgIII, which were used for the construction of truncated versions of BOI2 (shown schematically in Fig. 3 B) are underlined. (B) Comparison of the amino acid sequences of Boi2p and Boi1p. Identical residues are boxed in black. Domains I, II, III, and

SKSNIALAHSETPTSSNNKEAVSOPSEGKHKHKHKHKSKHKHKNSSSKDGSSEEKSKKKLFSSTKESFVGSKEFKRSPSE

TIHGTRLSYPTNT

SKACTL EEARLOTOLR

AKEAEIF

τv

-ETWKDDKNKR

AIKGE

SLYAASUGKGKYCFKLVPPOPGSKKGLTFT

NKRNSNYPIEQDQFETSD

WEQQELQQQQHDNNQGQADRTTSASTQR<mark>TSD</mark>EDNTISTPNLSSANNTTIGSNGF<mark>SS</mark>E

CFKLLPPQPGSKKGLTFT

ROMT

TKSTLP

HYFAV

HYFAU



round, and unbudded cells (Fig. 4 Aa). Overexpression of *BOI2* from the *GAL7* promoter (*pGAL7:BOI2*) also inhibited cell growth (Fig. 4 *B*, sector 4), and cells overexpressing *BOI2* were arrested as large, round, unbudded cells (Fig. 4 Ab). The asymmetric organization of actin filaments was disrupted in cells that overexpressed *BOI1* and in cells that overexpressed *BOI2* (data not shown). These results indicate that overexpression of *BOI1* and *BOI2* inhibits bud emergence, and that the Boi proteins affect cell polarization.

To identify the domain that is important for the inhibitory effect, we constructed two truncated versions of *BOI2*, namely $pGAL7:BOI2\Delta N$ and $pGAL7:BOI2\Delta C$. The overproduction of Boi2p without its NH₂-terminal half from $pGAL7:BOI2\Delta N$ inhibited cell growth (Fig. 4 *B*, sector 3) and cells expressing $pGAL7:BOI2\Delta N$ were arrested as large, round, unbudded cells (data not shown). The overproduction of Boi2p without its COOH-terminal

Figure 3. Domains required for the Bem1p-Boi2p interaction. (A) The BEM1 sequences, expressed as fusion proteins with lexA, are represented by lines below the map of BEM1. The site of the mutations in $bem1^{Leu123}$ and $bem1^{Leu208}$ are indicated by triangles. Plasmids indicated on the left were introduced into L40 cells that harbored either pGADBOI2 or pGAD424 (as a control), and the activities of β -galactosidase (units) and the His3 phenotypes were assayed. +, growth; -. no growth. K, KpnI and BanI; S, SalI; H, HpaI; and P, PstI. (B) The BOI2 sequences expressed as fusion proteins with Gal4TA are represented by lines below the map of BOI2. The BOI2-coding region is indicated by an open box. The SH3 domain (hatched), the proline-rich sequence (black), and the PH domain (cross-hatched) are indicated. Plasmids indicated on the left were introduced into L40 cells that harbored either plexBEM1 or pBTM116 (cont.), and the expression of the reporter genes was examined. E, BstEII; X, XbaI; H, HincII; B, BanII; A, AatII; R, EcoRI; G, BgIII. (C) Cell lysates were prepared from the wild-type strain KA31 that harbored plasmids pBEM1KK (lanes 1 and 8), pBEM1 Δ 1 (lane 2), pBEM1 Δ 2 (lane 3), pBEM1 Δ 1 Δ 2 (lane 4), pBEM1 Δ C (lane 5), pBEM1 Δ 2 Δ C (lane 6), or a dummy plasmid pKT10mycN (lane 7). The plasmids are shown schematically in Fig. 1 A. Total cell lysates were incubated with GST-Boi2p beads (right panel, lanes 1-7) or GST beads (right panel, lane 8) as a control. Total cell lysates (left panel), proteins coprecipitated with GST-Boi2p (right panel, lanes 1-7), and proteins coprecipitated with GST (right panel, lane 8) were resolved by SDS-PAGE, and myc-tagged Bem1p was detected by Western blotting analysis. The positions of the various forms of myctagged Bem1p, produced from pBEM1KK (open circle), pBEM1 Δ 1 (open square), pBEM1 Δ 2 (open triangle), pBEM1 Δ 1 Δ 2 (closed circle), and pBEM1 Δ C (closed square), pBEM1 Δ 2 Δ C (closed triangle) are indicated.

half from $pGAL7:BOI2\Delta C$, however, did not have any inhibitory effect on cell growth (Fig. 4 *B*, sector 2). These results indicate that the COOH-terminal half that contains domain IV is important for inhibition of bud emergence, while the NH₂-terminal half that contains domains I, II, and III is not essential for the inhibition.

The finding that the NH₂-terminal half is not essential for the inhibition suggests that the Bem1p-Boi2p interaction is not critical for the inhibitory effect. To clarify this, we introduced both constructs for the overexpression of *BOI* genes (*pGA68 and pGAL7:BOI2*) into *bem1* mutant cells in which *bem1* lacked the sequence for both SH3 domains (*bem1*- Δ SH3s). The overexpression of either *BOI1* or *BOI2* inhibited the growth of the *bem1*- Δ SH3s cells (data not shown). These results indicate that the proteinprotein interaction between Bem1p and the Boi proteins is not critical for the inhibitory activity of the overexpressed *BOI* gene. Α



В



Figure 4. Overexpression of BOI. (A) The morphology of BOIoverexpressing cells (a and b). (a) Cells (strain W303-1A) carrying pGA68 and cultured in SC-L were shifted to SCGal-L, and (b) cells (strain W303-1A) carrying pGAL7:BOI2 and cultured in YPD were shifted to YPGal. Cells were harvested 8 h after the shift. (B) Cells (strain W303-1A) carrying a dummy plasmid YIpUGAL7 (sector 1), pGAL7:BOI2 ΔC (sector 2), pGAL7: BOI2 ΔN (sector 3), or pGAL7:BOI2 (sector 4) were streaked on an SCGal-U plate and incubated at 30°C for 2 d.

Phenotypes of the boi-disrupted Cells

Disruption of neither *BO11* nor *BO12* alone resulted in a cell growth defect (Fig. 5 A, sectors 2 and 3). However, all but one of ~200 $\Delta boi1 \ \Delta boi2$ spores isolated from $\Delta boi1/+ \Delta boi2/+$ heterozygous diploid cells failed to form colonies. The single viable $\Delta boi1 \ \Delta boi2$ spore formed a tiny colony that grew very poorly. $\Delta boi1 \ \Delta boi2$ cells with *BO12* under the control of the *GAL7* promoter (*pGAL7:BO12*) grew on medium that contained 0.1% galactose, which induced the expression of *pGAL7:BO12* with low efficiency. $\Delta boi1 \ \Delta boi2 \ pGAL7:BO12$ cells dramatically reduced the growth

rate 24 h after a shift to glucose-containing medium, which repressed the expression of pGAL7:BOI2 and then grew very poorly (Fig. 5 A, sector 4). $\Delta boil \Delta boi2 pGAL7:BOI2$ cells grew very poorly on glucose-containing medium at all temperatures tested (20, 25, 30, and 37°C). These results indicate that *BOI1* and *BOI2* are functionally redundant, and that the Boi proteins are important for cell growth.

About 24 h after the shift to glucose-containing medium, $\Delta boi1 \ \Delta boi2 \ pGAL7:BOI2$ cells began to stop growing and lysed, as judged from the leakage of alkaline phosphatase into the medium (Fig. 5 B). From the staining of

Α



Β



Figure 5. Disruption of BOI. (A) Strains to be tested were streaked on a YPD plate and incubated at 30°C for 2 d as follows. Wild-type strain W303-1A (sector 1); $\Delta boi1$ strain YMB101 (sector 2); $\Delta boi2$ strain YMB202 (sector 3); $\Delta boi1 \Delta boi2 pGAL7$: BOI2 strain YMB1207 (sector 4); $\Delta boi1 \Delta boi2 BOI2\Delta N$ strain YMB1202 (sector 5); $\Delta boi1 \Delta boi2 pGAL7$:BOI2 strain YMB1207 carrying pRS314-BOI2 ΔC (sector 6); and $\Delta boi1 \Delta boi2 pGAL7$: BOI2 strain YMB1207 carrying pYO324-BOI2 ΔC (sector 7). (B) Wild-type strain W303-1A (1), $\Delta boi1 \Delta boi2 pGAL7$:BOI2 strain YMB1207 (2), and $\Delta rho3 pGAL7$:RHO4 strain YMR505 (3) were streaked on a YPD plate, incubated for 40 h at 30°C, and overlaid with a solution of BCIP. Black regions in the photograph were blue and indicated cell lysis. dead cells with methylene blue (Rose, 1975), it appeared that \sim 30% of all the cells in the culture 24 h after the shift were dead, and $\sim 95\%$ of these dead cells were budded (Fig. 6 a). Within 48 h after the shift, the fraction of dead budded cells increased to $\sim 60\%$ of the total cells. The remaining cells were large and round (Fig. 6a), and in these large round cells, the asymmetric organization of actin filament (as observed in wild-type cells; Fig. 6 b) was disrupted (Fig. 6 d). A majority of $\Delta boil \Delta boil$ cells lysed with buds, and the death of these $\Delta boil \Delta boil$ cells was not suppressed by the addition of an osmotic stabilizer into the medium (data not shown). In the presence of an osmotic stabilizer that partially prevented cell lysis, however, most of the $\Delta boil \Delta boil$ cells were uniformly observed as large and round cells in the glucose-containing medium rather than as dead budded cells. Furthermore, the asymmetric organization of actin filaments was disrupted in these large round cells (Fig. 6, e and f), indicating that these $\Delta boil \ \Delta boi2$ cells have lost cell polarity. These morphological observations strongly suggest that cells without the Boi function failed to maintain cell polarity, and that the loss of cell polarity caused a defect in bud growth and subsequent cell lysis.

The Boi2p Domain That Is Required for Cell Growth

To identify the domain that is critical for the Boi function, we constructed two truncated versions of BOI2: $BOI2\Delta N$, which lacked the sequence for domains I, II, and III; and $BOI2\Delta C$, which lacked the sequence for domain IV. $\Delta boi1$ $\Delta boi2$ cells carrying $BOI2\Delta N$ grew as well as wild-type cells (Fig. 5 A, sector 5), but $\Delta boi1$ $\Delta boi2$ cells carrying $BOI2\Delta C$ on either a low copy number plasmid or a high copy number plasmid grew as poorly as $\Delta boi1$ $\Delta boi2$ cells (Fig. 5 A, sectors 6 and 7). These results indicate that the COOH-terminal half is essential and sufficient for cell growth while the NH₂-terminal half is dispensable.

The Effect of BOI2 on ∆rho3 Cells

Boi2p interacted with SH3-2, which was required for suppression by Bem1p of the $\Delta rho3$ defect. This result may suggest the involvement of Boi2p in the suppression. We examined whether or not multiple copies of BOI2 could suppress the $\Delta rho3$ defect. Although the overexpression of BOI2 from pGAL7:BOI2 inhibited cell growth, the introduction of pYO324-BOI2, a high copy number plasmid that harbored BOI2, into cells did not inhibit the growth of wild-type cells or $\Delta rho3$ cells that had been rescued by overexpression of RHO4 (data not shown; Fig. 7, left). The growth of $\Delta rho3$ cells that carried pYO324-BOI2 was as poor as that of $\Delta rho3$ cells without pYO324-BOI2 (Fig. 7, right, sectors 1 and 2), indicating that BOI2 can not serve as a multicopy suppressor of the $\Delta rho3$ defect. Conversely, multiple copies of BOI2 inhibited the growth of $\Delta rho3$ cells that overexpressed BEM1. The growth defect of $\Delta rho3$ cells was rescued by overexpression of *BEM1* (Fig. 7, right, sector 3), but in the presence of pYO324-BOI2, the growth of the $\Delta rho3$ cells that overexpressed BEM1 was as poor as that of Arho3 cells without the BEM1 overexpression plasmid (Fig. 7, right, sector 4). However, neither BOI2 ΔN nor BOI2 ΔC on a high copy number plasmid did not show the inhibitory effect (Fig. 7, right, sectors



Figure 6. Morphology of $\Delta boil \ \Delta boil \ D bo$

5 and 6). These results indicate that the NH₂-terminal half and the COOH-terminal half of Boi2p is critical for the inhibitory activity. It might be possible that the *BEM1*-overexpressing $\Delta rho3$ cells are hypersensitive to the inhibitory effect of *BOI2* on bud emergence, which was observed in case of *pGAL7:BOI2*. Such was not the case, however, since in addition to the difference in required domains, the *BEM1*-overexpressing $\Delta rho3$ cells that harbored pYO324-BOI2 showed morphology similar to that of $\Delta rho3$ cells without the *BEM1* overexpression plasmid (showing an increase of the proportion of lysed cells with buds) instead of becoming large, round, unbudded cells (data not shown). These results suggest that Boi2p inhibits the function of Bem1p in the suppression of the $\Delta rho3$ defect.

Genetic Interactions among the BOI Genes, RHO3, and RHO4

Since (a) Boi2p interacts with Bem1p, (b) BEM1 interacts genetically with RHO3, and (c) the morphological pheno-



Figure 7. Effect of multiple copies of BOI2 on the growth of Δrho3 cells. Δrho3 pGAL7:RHO4 strain YMR505, carrying the indicated plasmids, was incubated on SCGal-UT (left) and SC-UT (right) plates at 30°C for 4 d. The Δrho3 strain YMR505 grew very poorly on glucose-containing medium but grew well on galactosecontaining medium as a result of the overexpression of RHO4 from pGAL7:RHO4 (Matsui and Toh-e, 1992b). Plasmids were as follows: pYO324 and pKT10, dummy plasmids (sector 1); pYO324-BOI2, a high copy number plasmid that harbored BOI2, and pKT10 (sector 2); pBEM1KK, a BEM1-overexpressing plasmid, and pYO324 (sector 3), pBEM1KK and pYO324-BOI2

(sector 4); pBEM1KK and pYO324-BOI2 Δ N, a high copy number plasmid that harbored *BOI2\DeltaN* (sector 5); pBEM1KK and pYO324-BOI2 Δ C, a high copy number plasmid that harbored *BOI2\DeltaC* (sector 6).

type of the $\Delta boil \Delta boil$ cells is similar to that of $\Delta rho3$ cells (Matsui and Toh-e, 1992b), we examined whether BEM1, RHO3, or RHO3-related genes could serve as multicopy suppressors of the *boi* defect. Plasmids were introduced into $\Delta boi1/+\Delta boi2/+$ heterozygous diploid cells, and the transformants were sporulated and dissected. After incubation at 25°C for 3 d, Aboil Aboil segregants carrying plasmids that overexpressed either RHO3 or RHO4 (i.e., plasmids pOPR3 or pOPR4) formed visible colonies. They grew as well as wild-type cells and much better than $\Delta boil \Delta boil$ cells without pOPR3 and pOPR4 (Fig. 8). These results indicate that both RHO3 and RHO4 can serve as a multicopy suppressor of boi. Plasmids overexpressing either BEM1 or RHO1, encoding a Rho-type GTPase, (i.e., pBEM1KK or pOPR1) did not rescue the growth defect of $\Delta boil \Delta boil$ segregants (data not shown).

Discussion

Functions of the Boi Proteins

The phenotypes of $\Delta boil \Delta boil$ cells resemble those of cells depleted of both Rho3p and Rho4p. Arho3 Arho4 cells carrying pGAL7:RHO4, RHO4 under the control of the GAL7 promoter cease to grow in a glucose-containing medium, and the Rho3p- and Rho4p-depleted cells lyse with buds (Matsui and Toh-e, 1992b; Fig. 5 B). In the presence of an osmotic stabilizer, Rho3p- and Rho4p-depleted cells become large and round, and the asymmetric organization of actin filaments is disrupted (Matsui and Toh-e, 1992b). The phenotypes of the Boi protein-depleted cells (Figs. 5 B and 6) were quite similar to those of Rho3p- and Rho4p-depleted cells, and they strongly suggest that these cells are defective in the maintenance of cell polarity for bud growth. Moreover, both RHO3 and RHO4 can serve as a multicopy suppressor of the boi defect. Both the strong genetic interactions and the phenotypes of mutants strongly suggest that the Boi proteins, Rho3p, and Rho4p are all involved in the same process that maintains cell polarity for bud growth.

Domains of Boi Proteins

Boi2p possesses four domains that are highly conserved in Boi1p (Fig. 2 B), and the high degree of conservation suggests that the domains play an important role in the function of the Boi proteins. Domain III contains a proline-



Figure 8. Suppression of the boi defect. Wild-type cells (strain W303-1A, sector 1), $\Delta boi1 \ \Delta boi2 \ pGAL7:BOI2$ cells (strain YMR1207, sector 2), a $\Delta boi1 \ \Delta boi2$ segregant carrying pOPR3 (sector 3), and a $\Delta boi1 \ \Delta boi2$ segregant carrying pOPR4 (sector 4) were streaked on a YPD plate and incubated for 2 d at 30°C.

rich sequence that is required for displaying the two-hybrid interaction with SH3-2 of Bem1p (Fig. 3). Domain I contains an SH3 domain that can interact with an SH3 domain-binding protein. Domain IV contains a PH domain. We analyzed the role of the domains using truncated versions of Boi2p, and the results are summarized in Fig. 9. BOI2 ΔN , which lacked the sequence for domains I-III, complemented $\Delta boil \Delta boi2$, while $BOI2\Delta C$, which lacked domain IV, did not, an indication that the COOH-terminal half, including domain IV, of Boi2p is essential and sufficient for the function of Boi2p in cell growth, while the NH₂-terminal half is dispensable. The role of a PH domain is still obscure, but it has been reported that PH domains participate in interactions with lipid moiety and proteins (Musacchio et al., 1993; Harlan et al., 1994). It is possible that Boi proteins interact with proteins other than Bem1p via the PH domain. In this context, it is of interest to recall that the overexpression of BOI genes under the control of the galactose-dependent promoter inhibited bud emergence (Fig. 4). As in the case of the function of Boi2p in cell growth, the COOH-terminal half of Boi2p (from $pGAL7:BOI2\Delta N$) is essential and sufficient for the inhibitory effect, and the interaction of Bem1p and the Boi proteins is not required. These findings suggest that the COOH-terminal half of Boi2p interacts with a factor(s) involved in bud emergence. It might be possible that the interaction between the COOH-terminal half and the component(s) for bud emergence can replace, in part, the role of the Bem1p-Boi2p interaction and can allow cells to grow well without the NH2-terminal half of the Boi proteins.

The Role of the Bem1p-Boi2p Interaction

Although the NH₂-terminal half of Boi2p is not essential for either cell growth or inhibition of bud emergence, the NH₂-terminal half of Boi2p might be required to inhibit the suppression of the growth defect of $\Delta rho3$ cells by Bem1p. These findings strongly suggest that the Boi proteins possess the potential to modulate the function of Bem1p and that the NH₂- and COOH-terminal portions of the Boi proteins play a critical role in modulating the function of Bem1p. The NH₂-terminal portion of Boi2p interacts with SH3-2 of Bem1p, and SH3-2 is critical for suppression of the $\Delta rho3$ defect. Thus, it is likely that the Bem1p-Boi2p interaction plays a role in the modulating activity.

The role of the Bem1p-Boi2 interaction in the suppression of the $\Delta rho3$ defect is suggested to be negative; multiple copies of BOI2 did not suppress the $\Delta rho3$ defect, but they inhibited the growth of $\Delta rho3$ cells that had been rescued by the overproduction of Bem1p (Fig. 7). From these results, we cannot exclude the possibility that the intrinsic amount of Boi2p might play a positive role in the suppression of the $\Delta rho3$ defect. It is possible, however, that another protein that interacts with SH3-2 might play an important role in suppressing the $\Delta rho3$ defect, acting in concert with or independently of Boi2p, and that Boi2p might modulate the function of Bem1p by interacting competitively with SH3-2. It was reported very recently that Bem1p interacts with Ste20p and Ste5p, two components of the pheromone-responsive mitogen-activated protein kinase cascade, as well as with actin. The Ste20p protein kinase requires both SH3-2 and the COOH-terminal half of Bem1p for the Ste20p-Bem1p interaction (Leeuw et al., 1995). Ste20p is therefore a candidate for a protein that is critical for the suppression of the $\Delta rho3$ defect. At this time, however, we have no experimental evidence to suggest the involvement of Ste20p.

Protein Complex for Bud Formation

The COOH-terminal 35 a.a. of Bem1p is required for bud emergence at elevated temperatures and for the suppression of the rho3 defect (Fig. 1). Thus, it appears possible that protein-protein interaction at the COOH terminus of Bem1p is critical for the functions of Bem1p. One of the candidates for an interacting protein is Cdc24p (Peterson et al., 1994). Indeed, the COOH-terminal 35.a.a. was required for the two-hybrid interaction between Bem1p and Cdc24p (Table II). Cdc24p also interacts with Cdc42p as a GTP-GDP exchange factor and with Rsr1p/Bud1p, which is needed for determination of the bud site (Bender and Pringle, 1989; Chant and Herskowitz, 1991; Zheng et al., 1994, 1995). The finding that Bem1p required the Cdc24pinteracting COOH terminus for suppression of the $\Delta rho3$ defect suggests the possibility that a protein complex containing Bem1p and Cdc24p might play a role in the process that involves Rho3p. It is plausible that the protein complex for bud-site selection is developed by the association of Cdc42p, Cdc24p, and Bem1p for the polarization of cells. After the initiation of bud emergence, the protein complex for cell polarization should be developed and/or rearranged to terminate the process of initiation of bud emergence and for the continuation of bud growth. In this context, it is of great interest that factors involved in bud growth exhibit genetic and protein-protein interactions with factors that are involved in bud emergence and, moreover, that Boi2p might be able to modulate the function of Bem1p and to interact with a factor(s) other than Bem1p that is involved in cell polarization. Our present working hypothesis is that for the regulation of cell mor-



Figure 9. Summary of the domain analysis of BO12. The BO12 sequence of each allele is indicated by a thick line below the coding region of BO12 (open box). The SH3 domain (hatched), the proline-rich sequence (black), and the PH domain (cross-hatched) are indicated. The regions of domains I, II, III, and IV are indicated by narrow lines. The abilities of each allele

to complement $\Delta boil \Delta boi2$, to inhibit bud emergence of wild-type cells when overexpressed under the control of the GAL7 promoter, and to inhibit the growth of the BEM1-overexpressing $\Delta rho3$ cells are indicated.

phogenesis, Rho3p and Boi proteins control the development of the protein complex for bud growth, and the Boi proteins mediate the connections between the protein complex for cell polarization and the machinery for bud growth via its affinities for Bem1p and for the factor(s) involved in cell polarization.

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