Cell Polarity in Saccharomyces cerevisiae Depends on Proper Localization of the Bud9 Landmark Protein by the EKC/KEOPS Complex

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ABSTRACT In diploid Saccharomyces cerevisiae cells, bud-site selection is determined by two cortical landmarks, [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) at the distal and proximal poles, respectively. Their localizations depend on the multigenerational proteins [Rax1p/](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000005827)[Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074). Many genes involved in bud-site selection were identified previously by genome-wide screening of deletion mutants, which identified [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) that causes a random budding in diploid cells. [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) is an atypical kinase involved in a signaling cascade of [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) kinase, the yeast homolog of Akt/PKB, and a component of the EKC/KEOPS (endopeptidase-like, kinase, chromatin-associated/kinase, putative endopeptidase, and other proteins of small size) complex that functions in telomere maintenance and transcriptional regulation. However, its role in bipolar budding has remained unclear. In this report, we show that the [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) kinase cascade does not affect bipolar budding but that the EKC/KEOPS complex regulates the localization of [Bud9p.](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) The kinase activity of [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494), which is essential for the functions of the EKC/KEOPS complex but is not necessary for the [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) signaling cascade, is required for bipolar bud-site selection. [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is necessary for random budding in each deletion mutant of EKC/KEOPS components, and [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) is genetically upstream of EKC/KEOPS genes for the regulation of bipolar budding. The asymmetric localization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) was dependent on the complex, but [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) were not. We concluded that the EKC/KEOPS complex is specifically involved in the regulation of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization downstream of [Rax1p/](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000005827)[Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074).

THE generation of cell polarity is important for the func-
tion of many cell types and underlies various processes such as cell division, differentiation, cell migration, cell–cell signaling, and fertilization (Freifelder 1960; Mooseker 1985; Hyman and White 1987; Bedinger et al. 1994; Chant and Pringle 1995; Kraut et al. 1996). In the budding yeast Saccharomyces cerevisiae, such polarization induces asym-

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metric growth to form a bud that becomes the daughter cell. Yeast cells polarize and divide by budding in two patterns: the axial pattern of haploid a/α -cells and the bipolar pattern of diploid a/ α -cells. Axial budding in the haploid a/ α -cells forms chain-like buds adjacent to the immediately preceding bud site, while bipolar budding in diploid a/α -cells forms cluster-like buds at either the proximal pole or its opposite distal pole. Alternate bud-site selections between proximal and distal poles usually occur in bipolar budding.

The establishment of yeast cell polarity undergoes three basic steps (Drubin and Nelson 1996). First, site selection occurs on the cell surface, a spatial landmark where cells will polarize. The bud sites during axial budding are marked with the axial landmarks [Axl1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000006326), [Axl2p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001402) [Bud3p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000520) and [Bud4p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003852) (Fujita et al. 1994; Chant et al. 1995; Halme et al. 1996; Sanders and Herskowitz 1996). The bipolar landmarks [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) of the distal and proximal poles, respectively, determine the polarization axis of budding in diploid cells (Harkins et al. 2001). These cortical landmarks provide recognition sites for the [Rsr1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003384)/[Bud2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001575)[/Bud5p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000634) GTPase

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signaling module (Kang et al. 2001; Park and Bi 2007). Next, the GTPase module transmits positional information from the axial and bipolar cortical markers to the protein [Cdc42p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004219) GTPase and its guanine nucleotide-exchange factor [Cdc24p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000039) for polarity establishment. Finally, [Cdc42p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004219) recruits the machinery that organizes and polymerizes actin, actinassociated proteins, and septins to the selected site of growth.

Numerous proteins have been found to be involved in bud-site selection (Park and Bi 2007). A genome-wide screening of homozygous deletion mutants identified 127 mutants representing three different bud-site phenotypes: unipolar, axial-like, and random (Ni and Snyder 2001). Among them, 112 mutants displayed strong or weak random budding phenotypes. The homozygous deletion mutant of the gene [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) displayed a random budding pattern (Ni and Snyder 2001). [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) was originally found as a gene encoding an atypical protein kinase that belongs to the piD261 family of atypical Ser/Thr protein kinases found in virtually all eukaryotic and archaeal organisms (Stocchetto et al. 1997). Recently, two independent studies of haploid cells have shown that [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) is a component of the KEOPS (kinase, putative endopeptidase, and other proteins of small size) or the EKC (endopeptidase-like, kinase, chromatinassociated) complexes with functions of telomere maintenance and transcriptional regulation, respectively (Downey et al. 2006; Kisseleva-Romanova et al. 2006). [Cgi121p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500), a component of the KEOPS complex, was identified by a genome-wide screen as a suppressor of $cdc13-1$ $cdc13-1$, an allele of the gene encoding the telomere-capping protein [Cdc13p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000002379) (Downey et al. 2006). The components of the KEOPS complex containing [Cgi121p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) are the protein kinase [Bud32p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) the putative peptidase [Kae1p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001746) and the uncharacterized protein [Gon7p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720). Deletion of [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) or genes encoding other KEOPS components produced phenotypes with short telomeres and failed to synthesize de novo telomeres to DNA doublestranded breaks (Downey et al. 2006). The EKC complex contains a homolog of cancer-testis antigens ([Pcc1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000028512)) and four additional proteins found in the KEOPS complex. [Pcc1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000028512) is a transcription factor, and its deletion affects the expression of several genes regulated by α -factor and galactose (Kisseleva-Romanova et al. 2006). [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) is phosphorylated by [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) kinase, a yeast member of the Akt/PKB subfamily. The phosphorylation of [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) positively regulates its ability to interact with [Grx4p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000976) and to phosphorylate it (Peggion et al. 2008). The signaling pathway from Akt/ PKB (homolog of [Sch9p\)](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) to PRPK (homolog of [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)) is evolutionarily conserved. [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) kinase regulates the nutrient signaling pathway in yeast (Roosen et al. 2005). The function of [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) in this pathway is independent of the role in the EKC/KEOPS complex (Peggion et al. 2008).

Genes that are specifically linked to the random budding of diploid cells encode factors important for the localization of bipolar cortical landmarks (Casamayor and Snyder 2002). Diploid cells select two poles for bipolar budding at the poles proximal or distal to the birth scar, which are determined by two cortical landmarks, [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273), respectively. [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) were originally identified as specific genes essential for normal bipolar budding. The $bud8\Delta$ $bud8\Delta$ and $bud9\Delta$ $bud9\Delta$ mutants bud at the proximal and distal poles, respectively (Harkins et al. 2001). In addition, [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) are localized at the distal and proximal poles, respectively (Harkins et al. 2001). [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is also localized at the distal pole, whose localization may inhibit the function of [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) through an interaction between the two (Taheri et al. 2000). Previous study suggests that many of the genes that specifically cause random budding, when deleted, are associated with the localization of [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) (Ni and Snyder 2001). [Rax1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000005827) and [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) are interdependently related in the maintenance of bipolar budding (Kang et al. 2004). The genes encoding these proteins were identified by screening genes that restored bipolar budding patterns in the haploid $ax11\Delta$ mutant (Fujita et al. 1994). [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) interact with and depend on [Rax1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000005827) and [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074), respectively, for their localization (Kang et al. 2004). The delivery of [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) to the presumptive bud site and mother-bud neck, respectively, is dependent on actin. The delivery of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is also dependent on septin (Schenkman et al. 2002).

In this article, we report that the [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) kinase cascade does not associate with bipolar budding, but that the EKC/ KEOPS complex is involved in bipolar budding. The deletion mutants of each EKC/KOEPS component displayed a random budding pattern, and the phenotype was suppressed by the deletion of [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273), suggesting that [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is a cause of random budding in these mutants. Consistent with phenotypic analysis, the asymmetric localization of GFP-[Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) was dependent on the EKC/KEOPS complex, but GFP-[Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) was not. The localization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is also regulated interdependently by [Rax1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000005827)[/Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074). The phenotype of double mutants of each EKC/KEOPS gene and [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) was changed to one similar to the $rax2\Delta$ $rax2\Delta$ mutant, which showed a random budding pattern specifically in diploid cells. The localization of [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) was normal even in $bud32\Delta$ $bud32\Delta$ and other mutant cells with deleted EKC/KEOPS components. These results suggest that the EKC/KEOPS complex is required for the localization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) and acts as a downstream factor of [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074).

Materials and Methods

Yeast strains, culture medium, plasmids, and recombinant DNA methods

Homozygous deletion mutants were generated by mating MATa cells with MAT α cells or by transformation of the plasmid expressing the [HO](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000002386) (Homothallic switching) endonuclease in haploid cells, which is expected to cause matingtype switching of MATa cells to MAT α or MAT α cells to MATa and hence subsequent mating between MATa and $MAT\alpha$ cells to form a homozygous diploid. In the [supporting](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/1) [information](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/1), [Table S1](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/6) summarizes the yeast strains used in this study. Yeast cells were grown in yeast extract peptone dextrose [YPD: 2% (w/v) bacto-peptone, 1% (w/v) yeast extract, and 2% (w/v) glucose] or in synthetic defined (SD) medium [0.2% (w/v) yeast nitrogen base without amino acids and ammonium sulfate, 0.5% (w/v) ammonium sulfate, and 2% (w/v) glucose] or in synthetic complete (SC) medium [0.2% (w/v) yeast nitrogen base without amino acids and ammonium sulfate, 0.5% (w/v) ammonium sulfate, 2% (w/v) glucose, and complete amino acids] lacking appropriate amino acids to maintain various plasmids. Plasmid preparation was conducted with Escherichia coli XL10 GOLD (Stratagene, La Jolla, CA) grown in Luria– Bertani medium [0.5% (w/v) NaCl, 1% (w/v) yeast extract, and 1% (w/v) tryptone] containing 50 μ g·ml⁻¹ ampicillin. Solid media were made with 2% agar. [Table S2](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/7) summarizes the plasmids used in this study. Standard methods of yeast genetics and DNA manipulation (Gietz et al. 1992; Ausubel et al. 1995; Longtine et al. 1998) were used, except where noted. Polymerase chain reaction (PCR) was performed using KOD FX DNA polymerase (Nacalai Tesque, Kyoto, Japan). Oligonucleotide primers shown in [Table S3](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/8) were synthesized and purchased from Invitrogen (Tokyo).

Plasmids

The pYC01 was created with pRS425 (Christianson et al. 1992) by inserting a fragment containing an upstream sequence of 808 bp, an open reading frame of [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345), and a downstream sequence of 517 bp. PCR was performed to amplify the [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) region with the primer set BUD8a/BUD8b and the genomic DNA of the BY4743 yeast strain as a template. The resulting XhoI/SpeI fragment was cloned into XhoI/SpeI-digested pRS425. Detailed procedures for the preparation of pYC03 to insert epitope tags using a MluI cleavable site $(5'-ACGGCT-3')$ just downstream of the start codon of [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) have been reported (Kato et al. 2009). For green fluorescent protein (GFP) tagging, PCR was performed using the primer set GFPa/GFPb and pFA6a-GFPS65T-kanMX6 (Longtine et al. 1998) as a template. The resulting MluI fragment was incubated with T4 polynucleotide kinase (T4 PNK) (Takara Shuzo, Kyoto, Japan) and then ligated with EcoRV-digested pBluescript (SK–) to create pYC05. MluI-digested pYC05 was separated using 1.5% agarose gel electrophoresis and gel-extracted. The resulting insert was ligated with MluI-digested pYC03 to generate pYC14. The GFP[-Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) fragment was cloned into pRS426 (Christianson et al. 1992) using the same procedure to generate the pYC06.

The pYC09 was constructed to delete [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720). PCR was first performed to amplify the [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) region with the primer set GON7a/GON7b, with genomic DNA of the BY4743 yeast strain as the template. The resulting fragment containing an upstream 570 bp, an open reading frame of [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720), and a downstream 519 bp was phosphorylated with T4 PNK and then cloned in EcoRV-digested pBluescript (SK–) to create pYC07. Deletion of the open reading frame of [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) and creation of a MluI restriction site to generate pYC08 were performed by PCR using the primer set GON7c/GON7d and pYC07 as a template. For cloning the [URA3](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000747) region as a selective marker, PCR was next performed using the primer set

URA3a/URA3b and genomic DNA of the B8032 strain as the template. The resulting MluI fragment containing an upstream 871 bp, an open reading frame of [URA3](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000747), and a downstream 1000 bp was digested with MluI and then cloned into MluI-digested pYC08 to create pYC09.

For FLAG tagging, PCR was performed using a primer set, FLAGa/FLAGb, and pTM55 as the template. The resulting fragment was digested with MluI and then ligated with MluI-digested pYC04 to create pYC10.

The pYC11 was created with pBluescript $(SK-)$ by inserting a fragment containing an upstream sequence of 520 bp, an open reading frame of [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494), and a downstream sequence of 789 bp. PCR was performed to amplify the [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) region with the primer set BUD32a/BUD32b and the genomic DNA of the BY4743 yeast strain as a template. The resulting SpeI/XhoI fragment was cloned into SpeI/XhoIdigested pBluescript $(SK-)$ to generate pYC11. Site-directed mutagenesis was performed to insert a MluI cleavable site (5'-ACGCGT-3') just upstream of the stop codon of [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) with primer set BUD32c/BUD32d and pYC11 as a template to generate pYC12.

Plasmids pYC13 and pYC15 were created with pBluescript (SK-) by replacing base pairs of $BUD32$ to mutate to alanine at Lys52 and Ser258 of [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494), respectively. PCR was performed to replace base pairs with the primer sets BUD32e/BUD32f, BUD32g/BUD32h, and pYC12 as a template to generate pYC13 and pYC15, respectively. The pYC13 was then digested with SpeI/XhoI, and the insert of pYC13 was transferred to pRS306 to generate pYC16.

MluI-digested pYC05 was separated using 1.5% agarose gel electrophoresis and gel-extracted. The resulting insert was ligated with MluI-digested pYC12, pYC13, and pYC15 to generate pYC17, pYC18, and pYC19, respectively. The resulting plasmids were then digested with SpeI/XhoI, and the inserts of these plasmids were transferred to pRS306 to generate pYC20, pYC21 and pYC22, respectively. All sequences were verified using an ABI 3010 sequencer (Applied Biosystems, Foster City, CA).

Strains

Strains expressing [Rax2p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)GFP, CCY032, and CCY033 were created by a one-step tagging method using PCR product (Longtine et al. 1998). A cassette of GFP^{S65T}-kanMX6 just upstream of the stop codon of [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) using the primer set [RAX2-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)GFPa[/RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)-GFPb and pFA6a-GFP^{S65T}-kanMX6 as a template was amplified by PCR. The BY4741 and BY4742 strains were transformed with the resulting fragments to create strains CCY032 and CCY033, respectively. The transformants were incubated on YPD plates at 30° for 1 day and then selected to YPD plates with G418 (100 μ g·ml⁻¹). The correct insertion of GFPS65T-kanMX6 was verified by PCR using the genomic DNA of candidates as the template. The haploid strains CCY032 and CCY033 were mated on YPD plates for 1 day at 30° and then selected on SC plates not containing Lys and Met for 2 days at 30° to obtain the diploid strain expressing [Rax2p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)GFP (CCY034).

The [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) deletion strains (CCY015, CCY016, CCY 022, and CCY025) were created by homologous recombination with PCR products from pYC09. PCR was performed using the primer set GON7a/GON7b and pYC09 as the template. The resulting fragment was transformed in $bud32\Delta$ $bud32\Delta$ (CCY001), [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ (CCY004), [bud8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) Δ (CCY010), and [bud9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) Δ (CCY012) mutants, respectively, and then the transformants were grown on $SC-Ura$ for 3 days at 25° . Alternatively, the one-step disruption method using PCR products was used to delete [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) in [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)-GFP (CCY034) or $rax2\Delta$ $rax2\Delta$ (CCY044) strains. PCR was performed using the primer set GON7e/ GON7f and pFA6a-His3MX6 as the template. The resulting fragment was transformed in the diploid strains CCY034 and CCY044, and then the transformants were grown in $SC-His$ at 25 \degree for 3 days. The resulting heterozygous strains CCY039 and CCY049 were then confirmed by PCR using these genomic DNAs as templates to verify disruption of [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720). After dissection and selection of spores from these strains, PCR was used to confirm the deletion of [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) and [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) using genomic DNA of these segregates as the template to obtain the haploid strains CCY040 and CCY050, respectively. MATa or $MAT\alpha$ was identified by mating with reference strains on SD plates for 3 days at 25°.

The strain expressing [Bud32p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)GFP (CCY052) was created by homologous recombination with a linearized fragment of pYC20, respectively. The pYC20 fragment was digested with SphI and purified by a gel extraction. The resulting linearized fragments were transformed in BY4741 strain. Transformants were incubated on YPD plate at 25° for 3 days and then transferred to an SC plate with 5-FOA $(1 \text{ mg} \cdot \text{ml}^{-1})$ at 25° for 3 days. After the single isolation of colonies on YPD plate, the resulting strain (CCY052) was confirmed by PCR using these genomic DNAs as templates to verify the correct insertion of [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)-GFP. The diploid [BUD32-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)GFP strain (CCY053) was generated by transformation of the plasmid expressing the [HO](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000002386) endonuclease in the CCY052 strain. The growth rate of the resulting strain (CCY053) was confirmed to be identical to that of the BY4743 strain by a spotting assay.

Strains expressing [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) (K52A) (CCY064), [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) (K52A)-GFP (CCY054 and CCY055), and [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) (S258A)-GFP (CCY067) were created by homologous recombination with a linearized fragment of pYC16, pYC21, and pYC22, respectively, using the same procedure to generate CCY052. The diploid [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) (K52A)-GFP strain (CCY056) was generated by mating CCY054 with CCY055, and the diploid [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) (S258A)-GFP strain (CCY068) was generated by transformation of the plasmid expressing the [HO](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000002386) endonuclease in CCY067.

Strains expressing [Bud32p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)13myc (CCY062) and [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) (K52A)-13myc (CCY065) were created by the one-step tagging method using PCR product (Longtine et al. 1998). A cassette of 13Myc-kanMX6 just upstream of the stop codon of [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) using the primer set BUD32i/BUD32j and pFA6a-13Myc-kanMX6 as a template was amplified by PCR. The BY4741 and CCY064 strains were transformed with the resulting fragment to generate strains CCY062 and CCY065, respectively. The transformants were incubated on YPD plates at 30° for 1 day and then selected to YPD plates with G418 (100 μ g·ml⁻¹). The correct insertion of 13Myc-kanMX6 was verified by PCR using genomic DNA of candidates as the template. Diploid strains (CCY063 and CCY066) were generated by transformation of the plasmid expressing the [HO](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000002386) endonuclease in CCY062 and CCY065, respectively. The growth rate of resulting the strain (CCY063) was confirmed to be identical to that of the BY4743 strain by the spotting assay.

Strains expressing [Cdc11p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003837)GFP (CCY071 and CCY072) were created by the one-step tagging method using PCR product (Longtine et al. 1998). A cassette of GFPS65TkanMX6 just upstream of the stop codon of [CDC11](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003837) using the primer set CDC11a/CDC11b and pFA6a-GFPS65TkanMX6 as a template was amplified by PCR. The BY4741 and BY4742 strains were transformed with the resulting fragments to generate strains CCY071 and CCY072, respectively. The transformants were incubated on YPD plates at 30° C for 1 day, and then selected to YPD plates with G418 (100 μ g·ml⁻¹). The correct insertion of GFP^{S65T}-kanMX6 was verified by PCR using genomic DNA of candidates as the template. Diploid strain (CCY073) was generated by mating CCY071 with CCY072. The growth rate of resulting strain (CCY073) was confirmed to be identical to that of BY4743 strain by a spotting assay.

Analysis of budding patterns

Cells grown to late log phase were harvested by centrifugation at 3000 \times g for 5 min, washed twice, and resuspended in water. Cells were fixed with a paraformaldehyde solution to a final concentration of 3.7% with gentle agitation for 30 min at room temperature. The fixed cells were washed twice and resuspended in water and then incubated with a final concentration of 5 μ g·ml⁻¹ calcofluor white for 20 min to visualize bud and birth scars. After washing twice with water, the fluorescent image of chitin rings was observed with an ECLIPSE 80t fluorescence microscope (Nikon, Tokyo) using a UV filter set.

Budding patterns were analyzed according to Chen et al. (2000). All analyses were performed in homozygous diploids. The bud-scar positions of daughter and mother cells were separately counted and analyzed. The daughter cells with the first bud scar were classified in relation to the bud position to the birth scar as "proximal" when one bud scar was at the proximal pole, as "distal" when one bud scar was at the distal pole, or as "random" when one bud scar was around equatorial region. The mother cells that had more than three bud scars were used (Chant and Pringle 1995) and classified on the basis of the relation of the bud positions to the birth scar as "bipolar" when one or more bud scars were at both poles, as "proximal" when all bud scars were at the proximal pole, as "distal" when all bud scars were at the distal pole, as "axial-like" when all bud scars were connected in a chain with at least one touching the birth scar, or as "random" when at least one or more bud scars were in the

Figure 1 Budding pattern of the deletion mutants of each EKC/KEOPS component in diploid cells. (A) Representative image of budding patterns in both diploid daughter and mother cells. Daughter cells showed three budding patterns in the wild type and mutants. The positions of the first buds on diploid daughter cells were scored as being at the pole proximal to the birth scar (a), at the pole distal to the birth scar (b), or at a random site (c) on the cell. Mother cells showed five budding patterns in the wild type and mutants. Each diploid mother cell budding pattern was determined by counting cells with more than three bud scars. Bud scars were concentrated at the two poles (d, as bipolar), only at the pole proximal [e and g (as chain-like bud)] or distal (f) to the birth scar, and at the random sites (h). (B) Budding patterns of the diploid deletion mutants. Strains used were diploid wild type (BY4743) and bud32D (CCY003), cgi121D (CCY006), gon7Δ (CCY009), bud32Δcgi121Δ (CCY017), bud32Agon7A (CCY018), and cgi121Agon7A (CCY019). Budding positions are classified as in the budding patterns of diploid cells in Figure 1A. At least 150 cells were scored for each bud scar pattern from both daughter and mother cells (the percentages are indicated). In daughter cells, the solid, open, and dotted boxes indicate cells with buds at only the proximal pole, at only the distal pole, and at a random site, respectively. In mother cells, the shaded, solid, open, slashed, and dotted boxes indicate cells with buds at two poles, at only the proximal pole, at only the distal pole, at the axial-like, and at random sites, respectively.

equatorial region. A representative image of each pattern in both daughter and mother cells is shown in Figure 1A. Over 150 cells of each mother and daughter cell were counted for the classification. For each strain, the average value from two independent experiments is shown.

Fluorescence microscopy

Cells expressing GFP-fused protein were grown for 12–16 h at 25 \degree and then harvested by centrifugation at 3000 \times g for 5 min, stained with calcofluor white, washed twice, and resuspended in water. The fluorescent images of the GFPfused proteins and bud scars in cells were observed under a fluorescence microscope (BZ-9000; Keyence, Osaka, Japan) using GFP and UV filter sets, respectively.

Western blotting

Yeast cell lysates were prepared as previously reported (Krappmann *et al.* 2007). Briefly, the same OD_{600} units of cells were resuspended in an 80 - μ l extraction buffer [50 mm Tris–HCl (pH 7.5), 50 mm DTT, 1 mm EDTA, and inhibitor cocktail; Roche, Basel, Switzerland], vortexed with glass beads for 10 min at 4 \degree followed by an additional 20 μ l of membrane extraction buffer [50 mm Tris–HCl (pH 7.5), 50 mM DTT, 1 mM EDTA, 10% SDS, and 5% Triton X-100] and then vortexed at 4° for 1 min. The protein concentration of the resulting extracts was determined by the Bradford method (Kruger 1994). Proteins were subjected to SDS-PAGE with 15% low-bis gels (Hirano 1989). The separated

proteins were electroblotted onto polyvinylidene fluoride membranes as described in Hirano (1989). Membrane blocking was performed using the method of Kawasaki et al. (2008). Monoclonal mouse anti-FLAG (Sigma-Aldrich, St. Louis) and peroxidase-coupled goat anti-mouse immunoglobulin G (Santa Cruz Biotechnology, Santa Cruz, CA) were used as primary and secondary antibodies, respectively.

Results

Complex made with Bud32p, Cgi121p, and Gon7p regulates bipolar bud-site selection

The EKC/KEOPS complex consists of [Kae1p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001746) [Bud32p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) [Cgi121p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500), [Gon7p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720), and [Pcc1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000028512) in haploid cells (Downey et al. 2006; Kisseleva-Romanova et al. 2006). We investigated whether the EKC/KEOPS complex is involved in bipolar bud-site selection. By using tandem affinity purification (TAP), we confirmed that the components of the complex pulled down with [Bud32p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)TAP or [Gon7p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720)TAP from diploid cells are the same as those of the haploid EKC/KEOPS complex (Y. Kato, H. Kawasaki, and H. Hirano, unpublished results). The budding patterns of the deletion mutants of each EKC/KEOPS component were analyzed, except those of [KAE1](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001746) and [PCC1](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000028512), whose deletion showed a severe growth defect. The budding patterns of diploid cells were classified as shown in Figure 1A. The first buds in daughter diploid cells of these mutants were at the distal pole, as observed in

Each value indicates the percentage of total cells.

the wild-type diploid cells (Figure 1B). The deletion of each component of the EKC/KEOPS complex did not affect the budding pattern of daughter diploid cells, while the mother diploid cells of these mutants showed a severe budding defect (Figure 1B and Table 1). About half of the mother cells showed a random budding pattern in these mutants, although the $gon7\Delta$ $gon7\Delta$ mutant displayed a slightly different phenotype with a higher distal budding ratio. We also examined the phenotype of several double mutants, in which two genes of the complex component were disrupted. The phenotypes of [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , bud32 Δ [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ , and cgi121 Δ gon7 Δ double mutants were very similar to those of the single mutants. (Figure 1B and Table 1). These results suggest that [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494), [CGI121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500), and [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) act in the same process as a complex.

Random budding of bud32 Δ , cgi121 Δ , and gon7 Δ is suppressed by deletion of BUD9 but not BUD8

[Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) are cortical landmarks in diploid cells for the distal and proximal poles, respectively (Harkins et al. 2001), because each deletion of these landmarks displays unipolar budding at the distal or the proximal pole, respectively. The first buds of daughter cells in $bud32\Delta$ $bud32\Delta$, [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , or $gon7\Delta$ $gon7\Delta$ mutants formed at the distal pole as in wild-type cells (Figure 1B and Table 1). This indicates that [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494), [Cgi121p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500), and [Gon7p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) may not be related to marking of the distal pole with [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345). We observed the effect of the de-letion of [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) or [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) in the [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ mutant cells. In the daughter cells of the [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ [bud8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) Δ double mutant, the buds were observed largely at the proximal pole (\sim 60%), but some were at the distal pole (20%)

and at the equatorial sites (\sim 20%) (Figure 2A and Table 1). The budding pattern of daughter cells in the [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ [bud8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) Δ double mutant was similar to that of the [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ [bud8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) Δ double mutant. In contrast, that of daughter cells in the [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ [bud8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) Δ double mutant was similar to that of the bud8 Δ mutant in that the first bud of $>90\%$ of cells shows the proximal budding. Because the proximal budding observed in the $bud8\Delta$ $bud8\Delta$ mutant would be dependent on [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273), Bud9p seems to act as the proximal marker in daughter cells of these double mutants. However, \sim 30–50% of cells in [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ [bud8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) Δ and [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ bud8 Δ mutants budded at the distal pole or at the equatorial site. Thus, the proximal marker may be somewhat impaired in these mutants.

In the [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ [bud9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) Δ double mutant, the first and subsequent buds were formed at the distal pole, as in the $bud9\Delta$ $bud9\Delta$ mutant (Figure 2A and Table 1). Similar results were obtained for the [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ [bud9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) Δ and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ bud9 Δ double mutants. Because distal budding of the $bud9\Delta$ $bud9\Delta$ mutant would be dependent on [Bud8p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) these observations suggest that [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) in daughter cells of these mutants still acts as the distal landmark.

As shown in Figure 1B, \sim 45–60% of [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and $gon7\Delta$ $gon7\Delta$ mutant mother cells showed a random budding pattern. Similarly, \sim 40–60% of [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ [bud8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) Δ and [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ bud8 Δ double-mutant mother cells showed a random budding pattern (Figure 2B). A distal budding pattern, which was observed in \sim 15% of [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ mutant cells, was almost completely suppressed and replaced with a proximal budding pattern (Figure 1B and Figure 2B). However, the majority of the double mutants showed a random budding pattern, although the ratio of the proximal budding pattern increased slightly.

Figure 2 Genetic interaction between BUD8 or BUD9 and EKC/KEOPS genes. (A and B) Budding pattern of daughter (A) and mother (B) cells with BUD8 or BUD9 deletion in the b ud32 Δ , cgi121 Δ , and gon7 Δ mutant backgrounds. Strains used were the diploid wild type (BY4743), bud8 Δ $(CCY011)$, bud9 Δ $(CCY013)$, bud32 Δ $(CCY003)$, bud32Δbud8Δ (CCY026), cgi121Δbud8Δ (CCY027), gon7Δbud8Δ (CCY028), bud32Δbud9Δ (CCY029), cgi121 Δ bud9 Δ (CCY030), and gon7 Δ bud9 Δ (CCY031). Budding positions are classified as in the budding pattern of diploid cells in Figure 1A. At least 150 cells were scored for each bud scar pattern from both daughter and mother cells (the percentages are indicated). The solid, open, and dotted boxes in A indicate daughter cells budded at proximal, distal, and equatorial sites, respectively. The shaded, solid, open, and dotted boxes in B indicate mother cells with buds at two poles, at only the proximal pole, at only the distal pole, and at the random site, respectively.

In mother cells of [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ [bud9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) Δ double mutants, the random budding pattern was replaced by a distal budding pattern, although bipolar budding observed in the $bud32\Delta$ $bud32\Delta$ mutant was still seen in the double mutant (Figure 2B and Table 1). The random budding of mother cells of $cgi121\Delta$ $cgi121\Delta$ and $gon7\Delta$ $gon7\Delta$ mutants was also suppressed by $BUD9$ deletion. In the absence of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273), a random budding pattern clearly replaced the distal budding pattern in which [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) was marked at the distal pole. These results suggest that the proximal marking by [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is regulated by [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) and other components of the EKC/KEOPS complex.

Localization of the bipolar landmarks Bud8p and Bud9p

We showed that [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273), but not [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345), is required for random budding of [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ mutants. Therefore, we observed the localization of these bipolar landmarks in the wild type, and these mutant cells in GFPtagged [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) or [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) expressed with a high-copy vector with fluorescence microscopy. The expression of GFP[-Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) or GFP-[Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) in diploid $bud8\Delta$ $bud8\Delta$ or $bud9\Delta$ $bud9\Delta$ mutant cells complemented the budding defects, suggesting that these tagged proteins provide a normal function. In the wild-type cells, GFP[-Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) was localized at the distal pole in 36% of unbudded cells ($n = 439$) and at the proximal pole in 17% of such cells (Figure 3A, cells 1 and 2, and Table 2). The GFP-[Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) signal in mother cells appeared to be weaker than that in daughter cells. It was localized at the bud tip

in 17% of small-/large-budded cells $(n = 529)$ and at the bud tip/neck in 22% of such cells (Figure 3A, cells 3–7, and Table 2). Localization only to the bud neck was also observed in 7% of such cells (Figure 3A, cell 8, and Table 2). GFP-[Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) was localized at the proximal pole in 17% of unbudded cells ($n = 200$), and the signal at the distal pole was observed in only 5% of such wild-type mother cells (Figure 3B, cells 1 and 2, and Table 3). In addition, GFP-Bud9p was also localized at the proximal pole (10%), bud neck (7%), and bud neck/proximal pole (9%) in small-/largebudded cells ($n = 254$; Figure 3B, cells 3–8, and Table 3).

In the [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ mutants, the localization patterns of GFP[-Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) were essentially identical to that of the wild type, but its GFP signal appeared to be very diffused around the poles, tips, and bud necks (Figure 3C, cells 1–5, and Table 2; GFP[-Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) localization patterns in the [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ mutant are those of the representative mutant). In addition, GFP-Bud8p was also observed at the bud tip and bud neck in cells that had bud scars at the equatorial position (Figure 3C, cells 5–8).

However, the localization to both poles and the bud neck of the GFP[-Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) signal dramatically decreased compared to those of the wild-type cells (\sim 3% of all cells) (Table 3). No GFP signal was observed in most cells (Figure 3D, cells 1–4; localization patterns of GFP[-Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) in the [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ mutant are those of the representative mutant), but a few cells were also present with the GFP signal in which normal

Figure 3 Effect on the localization of GFP-Bud8p and GFP-Bud9p due to loss of EKC/KEOPS components. (A and B) Localization patterns of GFP-Bud8p and GFP-Bud9p in the wild type. Wild type (BY4743) expressing the full-length GFP-Bud8p or GFP-Bud9p from its own promoter in pYC14 or pYC06, respectively, was grown for 12-16 h at 25° in synthetic complete (SC) -Leu or -Ura liquid medium, stained with calcofluor white, and then suspended in water for observation. GFP-Bud8p or GFP-Bud9p and bud scars were observed with a fluorescence microscope with GFP and UV filter sets, respectively. (C and D) Localization of GFP-Bud8p and GFP-Bud9p in the cgi121 Δ mutant. The cgi121 Δ (CCY006) expressing full-length GFP-Bud8p or GFP-Bud9p from their own promoters in pYC14 or pYC06 was grown for 12-16 h at 25 \degree in SC-Leu or -Ura liquid medium, stained with calcofluor white, and then suspended in water for observation. GFP-Bud8p or GFP-Bud9p and bud scars were observed with a fluorescence microscope with GFP and UV filter sets, respectively. (E) Expression level of Bud9p in the absence of Cgi121p. Diploid wild type (BY4743) and $cqi121\Delta$ mutant (CCY006), both expressing the FLAG₆-Bud9p in pYC10, were analyzed by Western blotting. The wild type (BY4743) was considered the control, which carried the empty pRS426. Cells were grown overnight to log phase in SC medium lacking uracil. The total cell extract from the same OD_{600} units of cells was analyzed by immunoblotting using an anti-FLAG antibody (top) and stained with Ponceau-S as a loading control (bottom).

[Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization was observed (Figure 3D, cells 5–8, and Table 3). Another visible GFP signal was observed primarily as a dot-like structure along the cell periphery of both daughter and mother cells (Figure 3D, cells 9–15). Because no obvious difference was seen at the [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) expression levels of mRNA and protein between the wild-type and mutant cells, we concluded that mislocalization of the GFP signal in these mutants was not due to a difference in [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) expression (Figure 3E and real-time PCR; Y. Kato, H. Kawsaki, and H. Hirano, unpublished data). These results also suggest that [Bud32p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) [Cgi121p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500), and [Gon7p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) are specifically related in the localization of [Bud9p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) but not [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345).

Bud32p kinase activity is required for Bud9p localization

We generated a kinase dead mutant ([bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)-K52A) by replacing the 52nd lysine residue with alanine in [Bud32p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) which is the binding site for the α - and β -phosphates of ATP (Mao *et al.*) 2008), to investigate whether kinase activity is required for the regulation of bipolar bud-site selection. The mutant displayed a random budding pattern, as observed in the $bud32\Delta$ $bud32\Delta$ mutant ([Figure S1](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/2)A and Table 1). These results suggest that kinase activity is also required for regulation of bipolar budsite selection. We observed the effect of deleting [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) or [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) in the kinase dead mutant ([bud32-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)K52A); the deletion of [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) suppressed random budding in the kinase dead mutant, but deletion of [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) did not [\(Figure S1,](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/2) B and C, and Table 1). Finally, we examined the localization of GFP[-Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and GFP[-Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) in the mutant [\(Figure S1D](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/2)). In the kinase dead mutant, the GFP[-Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) localization pattern was similar to that of wild-type cells, whereas the GFP[-Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) signal was not observed in the kinase dead mutant, as in the $bud32\Delta$ $bud32\Delta$ mutant [\(Figure S1](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/2)D). These results indicate that kinase activity is required for proper localization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273), but not [Bud8p.](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345)

Mutation at Ser258 of Bud32p did not affect bipolar bud-site selection

A recent report has suggested that [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) is involved in the [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) signaling pathway (Peggion et al. 2008). We investigated whether this signaling pathway is involved in bipolar bud-site selection. [Sch9p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) a homolog of yeast Akt, phosphorylates [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) at Ser258. This phosphorylation of [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) positively regulates its ability to interact with [Grx4p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000976) and to phosphorylate it. We constructed a mutant [bud32-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494)S258A in which Ser258 on [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) was replaced with Ala. We observed the budding patterns of $sch9\Delta$ $sch9\Delta$ and $bud32-S258A$ $bud32-S258A$. The budding patterns of these two mutants were similar to those of wild-type daughter and mother cells [\(Figure S2](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/3) and Table 1). This result suggests that the [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) signaling cascade is not related to bipolar bud-site selection.

Genetic interaction between RAX2 and EKC/KEOPS genes

[Rax1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000005827) and [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) are interdependently localized and also involved in [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization via interactions (Kang et al. 2004). In daughter cells of the $rax2\Delta$ $rax2\Delta$ mutant, about half of the first buds were at the distal pole, and the rest were at the

Each value indicates the percentage of total cells.

proximal pole. Mother cells of $rax2\Delta$ $rax2\Delta$ mutants largely showed a random budding pattern, as in the $bud32\Delta$ $bud32\Delta$, [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and $gon7\Delta$ $gon7\Delta$ mutants. However, the phenotype differed between $rax2\Delta$ $rax2\Delta$ and EKC/KEOPS mutants although the ratio of random budding in both mutants was almost the same. A small number of $rax2\Delta$ $rax2\Delta$ mutant cells showed axial-like budding, while a few [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ mutant cells showed distal budding (Figure 4A). We deleted [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) in the [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ mutant cells and observed the budding pattern of these mutants (Figure 4A). In both daughter and mother cells of the double mutants, the phenotype was changed to one similar to the $rax2\Delta$ $rax2\Delta$ mutant (Figure 4A and Table 1). The distal budding pattern of the $bud32\Delta$ $bud32\Delta$ mutant disappeared following [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) deletion and was replaced with an axial-like budding pattern. The axial budding pattern was never observed in the single mutants [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ .

We observed [Rax2p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)GFP in wild-type and mutant cells (Figure 4, B and C, and [Figure S3](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/4)). Cells chromosomally expressing [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)-GFP also displayed a bipolar budding pattern, as in wild-type cells, suggesting that [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)-GFP is functional. In wild-type cells, [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074)-GFP localized mainly to bud scars and birth scars in unbudded cells and sometimes to bud tips in budded cells. Similar results were obtained from [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ single mutants (Figure 4B and [Figure S3A](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/4)). In addition, the localization of [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) at bud scars could be observed even in equatorial bud scars, suggesting that the regulation for localization of [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) is normal in the mutants (Figure 4C and [Figure S3B](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/4)).

Discussion

EKC/KEOPS complex is required for bipolar bud-site selection

[BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) was originally found as a gene encoding an atypical protein kinase found in virtually all eukaryotic and archaeal organisms (Stocchetto et al. 1997). It was also identified as a gene that regulates bud-site selection of diploid cells through a genome-wide analysis (Ni and Snyder 2001). [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) is involved in the signaling pathway of [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) kinase (Peggion et al. 2008) and is also involved in telomere or transcriptional regulation as a component of the EKC/ KEOPS complex (Downey et al. 2006; Kisseleva-Romanova et al. 2006). In an analysis with gel-filtration chromatography in haploid cells, the subcomplex containing only [Kae1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001746) and [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) was observed in yeast cells although it could have been a preparation artifact (Kisseleva-Romanova et al. 2006). No other subcomplex has been reported. We purified the EKC/KEOPS complex from diploid cells using a TAP pulldown. The diploid EKC/KEOPS complex was composed of [Bud32p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) [Cgi121p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500), [Kae1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001746), [Gon7p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) and [Pcc1p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000028512), which is identical to the haploid complex. Of the genes encoding these components, [CGI121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) and [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) were identified as new genes required for bipolar bud-site selection (Figure 1B). The EKC/KEOPS complex regulates bipolar bud-site selection, as deletion of the components showed a very similar phenotype with random budding. We also revealed that the kinase activity of [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) is required for bipolar bud-site selection ([Figure S1\)](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/2). This activity is essential for the functions of the EKC/KEOPS complex, such as transcriptional regulation and telomere maintenance, and it is also contributed to, but was not absolutely essential for, the [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) signaling cascade (Peggion et al. 2008). In addition, we observed that phosphorylation at Ser258 on [Bud32p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) by [Sch9p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) the modification of which did not alter its kinase activity, was not necessary for bipolar bud-site selection ([Fig](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/3)[ure S2](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/3)). Therefore, we concluded that the EKC/KEOPS complex is involved in bipolar bud-site selection with a signaling pathway different from the [Sch9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001248) kinase.

EKC/KEOPS complex is involved in the regulation of Bud9p localization

We showed that [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) deletion in the [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and $gon7\Delta$ $gon7\Delta$ mutants suppressed random budding and increased budding at the distal pole. In concordance with these results,

Each value indicates the percentage of total cells.

Figure 4 Genetic interaction between RAX2 and EKC/KEOPS genes. (A) Budding pattern of cells with deletion of RAX2 in the bud32 Δ , cgi121 Δ , and gon7 Δ mutant backgrounds. Strains used were the diploid wild type (BY4743), rax2Δ (CCY044), bud32 Δ rax2 Δ (CCY046), cgi121 Δ rax2 Δ (CCY048), $gon7\Delta$ rax2 Δ (CCY051), and bud32 Δ (CCY003). Budding positions are classified as in the budding pattern of diploid cells in Figure 1A. At least 150 cells were scored for each bud scar pattern from both daughter and mother cells (the percentages are indicated). In daughter cells, the solid, open, and dotted boxes indicate cells with buds at only the proximal pole, at only the distal pole, and at a random site, respectively. In mother cells, the shaded, solid, open, slashed, and dotted boxes indicate cells with buds at two poles, only the proximal pole, only the distal pole, axial-like and at random sites, respectively. (B) Diploid wild type (CCY034) and cgi121 Δ

(CCY038) expressing Rax2p-GFP from their own promoter at their chromosomal locus were grown overnight to log phase in YPD liquid medium, stained with calcofluor white, and then suspended in water for observation. Rax2p-GFP and bud scars were observed with a fluorescence microscope with GFP and UV filter sets, respectively. Bar, 5 μ m. (C) Bud scars at the equatorial position of random budding sites were also marked by Rax2p-GFP. The arrows indicate Rax2p-GFP localization at random positions. Bar, 5 μ m.

[Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization was affected by deletion of [BUD32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494), [CGI121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500), and [GON7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) (Figure 3D and Table 3). The deletion of [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) did not suppress random budding of $bud32\Delta$ $bud32\Delta$, $cgi121\Delta$ $cgi121\Delta$, and gon 7Δ mutants although proximal budding increased in the double mutants in place of the distal budding observed in the single mutants (Figure 2B). The proximal budding in the mutants might be due to the accidental localization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) at the proximal pole because [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is expressed in a normal amount in these cells (Figure 3E).

Our data suggest that mislocalization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) by loss of the EKC/KEOPS components caused random budding.

Proper localization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is also dependent on the timing of [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) expression that was observed in the G1 phase (Schenkman et al. 2002). A possibility exists that the EKC/ KEOPS complex might be related to the timing of [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) expression because the complex controls transcription of several genes (Kisseleva-Romanova et al. 2006). In a promoter swapping assay, GFP[-Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) expression from the [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) or [CLB2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000006323) promoter specifically at the G2/M phase failed to rescue unipolar proximal budding in a $bud9\Delta-1$ $bud9\Delta-1$ mutant, indicating inefficient delivery or poor stability of [Bud9p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) or both (Schenkman et al. 2002). Furthermore,

Figure 5 Regulation of the localization of the bipolar markers Bud8p and Bud9p by the EKC/KEOPS complex and Rax1p/Rax2p. The schematic illustration is based on figure 3 in Casamayor and Snyder (2002). For details, see the Discussion.

virtually no random budding was seen in these cases. Local-ization of GFP-[Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) in [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ mutants was observed in a dot-like structure at cell periphery, as shown in Figure 3D. We confirmed that the stability and expression level of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) were the same as that of the wild type (Figure 3E). GFP[-Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) from the [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) promoter in the $bud9\Delta-1$ $bud9\Delta-1$ mutant appeared to localize in internal vesicles, and the mutant showed mainly unipolar distal budding (Schenkman et al. 2002). If the time lag for [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) expression occurs in the [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ , it should largely be unipolar distal budding not random budding. Therefore, we conclude that random budding in the mutants is caused by mislocalization at the cell periphery after delivery of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) to the plasma membrane, but not through a time lag in [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) expression. Support for this conclusion was also provided by our observation that GFP-[Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) localization patterns in the mutant deleted of each EKC/KEOPS component were similar to those of the wild type because the delivery of both [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) to the presumptive bud site is dependent on actin (Schenkman et al. 2002).

The deletion of [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) in diploid cells induced random budding and axial-like budding, although \sim 20% of cells showed a bipolar budding pattern. Axial budding was suppressed in diploid cells, as [Axl1p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000006326) on which axial budding is dependent, is transcriptionally repressed in diploid cells (Fujita et al. 1994). However, the allele mutant of both [BUD8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [BUD9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) uses axial budding cues (Zahner et al. 1996). Several components of the axial budding system, such as [Axl2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000001402) and [Bud3p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000520), are also expressed in diploid cells. Indeed, [BUD3](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000000520) deletion in the [bud8](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) Δ [bud9](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) Δ double mutant essentially induces a random budding pattern (Harkins et al. 2001). Axial budding cues can be recognized with less efficiency in diploid cells when both bipolar markers are impaired. However, we never observed an axial-like budding pattern in the deletion mutant of EKC/KEOPS components (Figure 1 and Figure 2). This indicates that the axial budding system is essentially inactive in these cells although a very small part of the bud at the proximal pole might use such a cue.

We observed that deletion of [RAX2](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) in [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , and $gon7\Delta$ $gon7\Delta$ mutants changed the phenotype to one identical with $rax2\Delta$ $rax2\Delta$ (Figure 4A). In concordance with this observation, [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) distribution did not differ between the wild type and [bud32](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003494) Δ , [cgi121](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004500) Δ , or [gon7](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003720) Δ (Figure 4, B and C, and [Figure S3](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/4)). We also confirmed that regulation of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization by the EKC/KEOPS complex is independent of septin by observing the localization of [Cdc11p-](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003837)GFP, one of the septin subunits, in the deletion mutant of the EKC/ KEOPS component ([Figure S4\)](http://www.genetics.org/cgi/data/genetics.111.128231/DC1/5). These results suggest that the EKC/KEOPS complex acts to regulate [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization as a downstream factor of [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) and septin.

Regulation of bipolar bud-site selection by the EKC/KEOPS complex

We concluded that the EKC/KEOPS complex in diploid cells acts as a [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) downstream factor and that the complex maintains bipolar budding through regulation of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) lo-

calization (Figure 5). The localization of the cortical markers [Bud8p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) and [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is regulated by several proteins as shown in Figure 5. The EKC/KEOPS complex is the factor regulating the localization of only [Bud9p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) downstream of [Rax2p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004074) and septins. Recent reports suggest that the complex functions in telomere maintenance, transcriptional regulation, and $t⁶A$ modification (Downey et al. 2006; Kisseleva-Romanova et al. 2006; Daugeron et al. 2011; Srinivasan et al. 2011), although direct target of the complex is still elusive. The complex localized mainly to the nucleus, and the minor portion of the EKC/ KEOPS complex localized to cytosol (Y. Kato, H. Kawsaki, and H. Hirano, unpublished data), whereas [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localized to the cell surface. Although the complex in the cytosol might be directly linked to [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273), we could not observe a physical interaction between these proteins. These results suggest that the regulation of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization by the complex may not occur at the cell surface and that the localization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) might be regulated by factor(s) under the transcriptional control by the EKC/KEOPS complex. Further experiments, such as suppressor screening that can rescue [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization in the deletion mutants of each EKC/KEOPS component, might be able to identify the factors that are important for establishing and maintaining spatial cues in cells.

Our results suggest that the correct localization of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) is essential for bipolar budding. Although many of the genes that specifically caused random budding when deleted are associated with the localization of [Bud8p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000004345) so far no genes have been reported that specifically regulate only [Bud9p,](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) except for the septins (Ni and Snyder 2001; Schenkman et al. 2002). Our findings revealed that the EKC/KEOPS complex is specifically involved in the regulation of [Bud9p](http://www.yeastgenome.org/cgi-bin/locus.fpl?dbid=S000003273) localization. Further analysis of this regulation system will be an important step toward understanding asymmetric development from spatial cues in diploid cells.

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Cell Polarity in Saccharomyces cerevisiae Depends on Proper Localization of the Bud9 Landmark Protein by the EKC/KEOPS Complex

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Figure S1 Phenotype of the kinase dead mutant (bud32-K52). (A) Budding patterns of the kinase dead mutant. Strains used were diploid BY4743, the bud32 Δ (CCY003), the bud32-K52A (CCY056). (B) Budding pattern of cells with BUD8 deletion in the kinse dead mutant backgrounds. Diploid strains used were the bud32 Δ (CCY003), bud32Abud8A (CCY026), K52Abud8A (CCY058) and bud8A (CCY011). (C) Budding pattern of cells with BUD9 deletion in the kinse dead mutant backgrounds. Diploid strains used were the bud32 Δ (CCY003), bud32 Δ bud9 Δ (CCY029), K52Abud9A (CCY060) and bud9A (CCY013). Budding positions are classified as in the budding pattern of diploid cells

in Figure 1A. At least 150 cells were scored for each bud scar pattern from both daughter and mother cells; the percentages are indicated. The black, white, and dotted boxes indicate daughter cells with bud at only the proximal pole, only the distal pole, or the random site, respectively. The gray, black, white, and dotted boxes indicate mother cells with buds at two poles, only the proximal pole, only the distal pole, or the random site, respectively. (D) Localization of GFP-Bud8p and GFP-Bud9p in the kinase dead mutant. The bud32-K52A (CCY065) mutant expressing full-length GFP-Bud8p and GFP-Bud9p from its own promoter in pYC14 or pYC06, respectively, was grown for 12-16h at 25°C in synthetic complete (SC)-Leu or -Ura liquid medium, stained with calcofluor white, and then suspended in water for observation. GFP-Bud8p/GFP-Bud9p and bud scars were observed with a fluorescence microscope with GFP and UV filter sets, respectively.

Figure S2 Budding patterns of bud32-S258A and sch9D mutant. Strains used were diploid BY4743, sch9 Δ (CCY070) and bud32-S258A (CCY068). Budding positions are classified as in the budding pattern of diploid cells in Figure 1A. At least 150 cells were scored for each bud scar pattern from both daughter and mother cells; the percentages are indicated. The black, white, and dotted boxes indicate daughter cells with bud at only the proximal pole, only the distal pole, or the random site, respectively. The gray, black, white, and dotted boxes indicate mother cells with buds at two poles, only the proximal pole, only the distal pole, or the random site, respectively.

B.

Figure S3 Localization of Rax2p-GFP in bud32 Δ and gon7 Δ mutants. (A) Diploid bud32 Δ (CCY036) and gon7 Δ (CCY041) expressing Rax2p-GFP from their own promoter at their chromosomal locus were grown overnight to log phase in YPD liquid medium, stained with calcofluor white, and then suspended in water for observation. Rax2p-GFP and bud scars were observed with a fluorescence microscope using GFP and UV filter sets, respectively. Scale bar indicates 5 um. (B) Bud scars at the equatorial position of random budding sites were also marked by Rax2p-GFP in diploid bud32 Δ (CCY036) and gon7 Δ (CCY041) mutants. The arrows indicate Rax2p-GFP localization at random positions. Scale bar indicates 5 µm.

Figure S4 Effect on the licalization of a septin component, Cdc11p-GFP. (A) Localization of Cdc11p-GFP in cgi1211 mutants and wild type cells. Diploid wild-type (CCY073) and cgi121 Δ (CCY075) mutant cells expressing Cdc11p-GFP at its chromosomal loci were grown overnight to log phase in YPD liquid medium and the suspended in water for observation. Cdc11p-GFP and bud scars were observed with fluorescence microscope using a GFP and UV filter sets, respectively. Scale bar indicates 5 µm. (B) The septin ring is formed in the mutant cells at the equatorial position. Scale bar indicates 5 µm.

7 SI

8 SI

 $9S1$

10 SI

a All strains are congenic to BY4743.

b The strain was generated by a precise replacement of the GON7 open reading frame by URA3.

Table S2 Plasmids used in this study

Table S3 Primers used in this study

