The mother–daughter mating type switching asymmetry of budding yeast is not conferred by the segregation of parental HO gene DNA strands

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The Saccharomyces cerevisiae HO gene, which encodes a site-specific endonuclease, is transcribed in the parent (mother) cell but not in the daughter (bud) cell. Consequently mother cells can switch their mating type whereas bud cells cannot. Whether or not the different capabilities of these cells are due to a nonrandom segregation of parental HO gene DNA strands to progeny cells is tested here by assaying the pattern of switching in cells where the HO gene is inverted in the chromosome. In an inverted HO strain, as is the case with the normal HO arrangement strains, only the mother cells were found to switch. Thus, the possibility of asymmetric segregation of parental HO gene DNA strands does not regulate the mother–daughter asymmetry of switching.

[Key Words: Saccharomyces cerevisiae; HO gene; mating type switching]

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A fundamental question in developmental biology is how the progeny of a given cell division can acquire different developmental fates. The interconversion of the mating cell types of the budding yeast Saccharomyces cerevisiae has been used as an experimental system to address the question of developmental asymmetry between daughter cells. The alternate alleles of the mating type locus (MAT), called MATa and MATa, interchange by a transposition–substitution event whereby a copy of either the storage loci HML (a locus that usually contains unexpressed \( \alpha \) information) or HMR (a locus that usually contains unexpressed \( \alpha \) information) is transmitted to the expressed MAT locus, resulting in a cell type change (for a recent review of the system, see Klar et al. 1984). The recombination event is initiated by a site-specific, double-stranded break found at the MAT locus (Strathern et al. 1982, Kostriken et al. 1983). The HO gene, which is located on chromosome IV, encodes the site-specific “Y/Z endonuclease” which catalyzes the double-stranded break (Kostriken and Heffron 1984).

An interesting finding is that among a pair of sister cells only the older (parental) cell produces a pair of switched progeny in about 73% of cell divisions, whereas the newly born daughter (bud) cell never switches (Hicks and Herskowitz 1976, Strathern and Herskowitz 1979). This pattern of switching is due to the expression of the HO gene in mother cells and its failure to be expressed in daughter cells (Nasmyth 1983). Why the HO gene is expressed in mother cells and not in daughter cells is the subject of this study.

Regulation of the HO gene is complex. At least three separate regulatory systems act on the HO gene such that it is only transcribed [1] in both \( \alpha \) and \( \alpha \) but not in \( \alpha /\alpha \) diploid cells (Jensen et al. 1983); [2] only late in the G1 phase of the cell cycle, and [3] only in mother cells and not in daughter cells (Nasmyth 1983). At least six trans-acting, positive regulatory genes called SWI (for switching), have been implicated in HO regulation. Recent studies indicate that some of the SWI products may activate HO by antagonizing negative regulatory activities, of which there are at least two, in addition to the \( \alpha /\alpha \) control (Jensen et al. 1983; Breeden and Nasmyth 1987; Nasmyth et al. 1987a,b; Sternberg et al. 1987; for review, see Klar 1987b). Over 1400 bp of DNA upstream of the transcription site are required for correct expression and regulation of the HO gene (Nasmyth 1985).

One testable model that can account for restriction of HO gene expression to mother cells is that the chromatid inheriting the parental DNA strand containing a transcriptionally competent HO gene is specifically segregated to the mother cell. This competence may be acquired by assembling a transcription complex on one of the HO strands. In contrast, incompetent HO is specifically segregated to the daughter cell (Nasmyth 1983, Brown 1984, Klar 1987a,b; Nasmyth et al. 1987b). Completion of the assembly process may require progression through the daughter's entire cell cycle, in which case the negative and positive controls discussed above could be required for precise regulation of the assembly process. This mechanism generates two nonequivalent sister chromatids. A precedent for such a model is provided by the recent studies with the distantly related fission yeast Schizosaccharomyces pombe. In this case it is argued that the potential for mating type switching is
Due to a strand-specific “imprinting” event of the recombinational substrate such that only one member of a pair of progeny cells can acquire the competence for switching. Consequently, it is suggested that two sister cells are developmentally nonequivalent since they inherit different strands of parental DNA [Klar 1987a].

Recent studies have demonstrated that a positive regulator of HO, called SWI5, is probably the key determinant for conferring mother–daughter asymmetry in budding yeast. Several observations are relevant. First, the SWI5 gene is expressed in a cell cycle-regulated fashion in both mother and daughter cells, but interestingly, this positive regulator is transcribed in the S phase well after the window in which HO is transcribed in late G1 (Nasmyth et al. 1987b). Therefore, the mother cell must asymmetrically inherit the SWI5 gene product from the previous cell cycle to express HO. Second, Nasmyth et al. (1987b) have demonstrated that when the SWI5 gene is expressed throughout the cell cycle by placing it under the inducible GAL1 promoter, then both mother and daughter cells can switch efficiently. Third, the SWI5 protein binds in vitro to the HO promoter [D. Stillman and K. Nasmyth, per. comm.] and thereby blocks the action of the negative regulator(s) of HO called SIN3 (Sternberg et al. 1987) and SD11 [Nasmyth et al. 1987a]. Fourth, stationary-phase mother cells, all of which are G1 arrested, are capable of transcribing HO when they undergo “START” following inoculation into fresh medium, that is, G1 arrested mothers “remember” being mothers even after the block. Interestingly, a deletion of HO upstream regulatory region between –788 and –1160 destroys its ability to be transcribed following G1 arrest without affecting HO activity in subsequent cycles [Nasmyth et al. 1987b]. An explanation proposed by Nasmyth et al. (1987b) to account for these findings is that SWI5 may be inherited by mother cells only as a stable transcription complex whose stability is reduced in the case of the deleted promoter. Specifically, SWI5 may maintain a stable transcriptional complex at HO on only one chromatid following DNA replication. In this model the chromatid containing the competent HO segregates to the mother cell pole at cell division.

A critical test of the strand segregation model is that the daughter cells should switch in those strains containing an inverted HO gene, whereas the mother cells should not. The results presented here rule out a strand segregation model involving HO for conferring developmental asymmetry between mother and daughter cells.

Results

Installing HO in either orientation in chromosome IV

The HO gene was inserted next to the CDC36 locus located in linkage group IV by the following manipulations. The plasmid p519 contains the HO gene on a 4.3-kb Sau3A–EcoRI fragment cloned between the BamHI and EcoRI sites of pBR322 [see Fig. 1]. This clone contains over 1700 bp of sequences upstream from the transcription start site and contains the known regulatory sequences required for proper HO regulation [Nasmyth 1985]. A 2.2-kb SalI–Xhol fragment containing the selectable yeast LEU2 gene was cloned into the SalI site of p519 to generate the plasmid pAK32. A 2.7-kb SacI fragment, which contains a portion of the CDC36 gene, was derived from the plasmid YRp7–CDC36.1 [Breter et al. 1983] and ligated into the unique SacI site of pAK32. This site is present at position –1468 upstream from the HO transcription start [Nasmyth 1985]. The resulting plasmid pAK37 contains the CDC36 SacI fragment in arbitrarily designated orientation I, while pAK36 contains the inverted orientation II. Both plasmids are 13.2 kb in size.

To place these constructs near CDC36 in the chromosome, we used the observation that cleaved DNA molecules when introduced into yeast promote efficient homologous recombination at the cleaved ends [Orr-Weaver et al. 1981]. Therefore, to target the insertion next to CDC36, plasmids pAK36 and pAK37 were cleaved at the unique SalI site present in the CDC36 SacI fragment [see Fig. 1]. The linear DNA was used to transform strain DC150 (MATa leu2), and Leu+ transformants were selected. Required transformants should integrate HO, LEU2, and vector sequences, flanked by a single CDC36 SacI fragment on either side, into chromosome IV. In addition, since they contain an active HO gene, these transformants should generate MATa/MATα diploid clones as a result of switching mating type and subsequent mating between cells of opposite type. The MATa/MATα cells, unlike the parental strain, fail to mate and also can undergo meiosis and sporulation when transferred onto sporulation media. We found that most of the transformants indeed were stable Leu+ and had acquired the HO gene since they had diploidized.

A Southern analysis of DNA isolated from several transformants demonstrated that the required genomic construction was obtained. Figure 2 demonstrates such an analysis of the parental strain DC150 and two transformants. The strain K734 [see Table I for complete genotype] was obtained from transformation with pAK37 DNA and strain K735 was derived from transformation with pAK36 DNA. The desired transformants can be identified by the pattern of hybridization to a genomic Southern to the YRP7–CDC36.1 [Breter et al. 1983] probe. When DNA is digested with BglII, the parental 7.0-kb fragment, which contains the SacI fragment of the CDC36 gene, should disappear, and instead two fragments of sizes 11.3 kb and 6.9 kb from orientation I [pAK37 transformation] and of sizes 14.0 kb and 4.2 kb from orientation II [pAK36 transformation] should be generated. [7.0 kb + 13.2 kb – 2.0 kb = 11.3 kb + 6.9 kb [orientation I] = 14.0 kb + 4.2 kb [orientation II]; the 2.0-kb BglII fragment contained within the HO gene is not detected because the probe does not contain HO.] Predicted fragments were obtained (Fig. 2). In such a transformation only about one in three transformants showed a pattern consistent with the described construction. The rest of the transformants also resulted
from homologous recombination but contained multiple insertions [data not shown]. Figure 3 diagrams the HO inserts used for these experiments.

The HO orientation does not affect the mother–daughter switching asymmetry

The asci obtained from strains K734 and K735 were digested with gluculase and the resulting spores and their progeny were subjected to pedigree analysis [see Experimental procedures]. Only α to a switches were tested. As presented in Table 2, mother cells of strain K734 switched in 35.2% of cell divisions while those of strain K735 switched with an efficiency of 27.3%. Neither strain switched both their spores and daughter cells, and, therefore, followed the pattern observed with wild type HO-containing strains [Hicks and Herskowitz 1976; Strathern and Herskowitz 1979].

Both strains K734 and K735 that were used have the natural “inactive” allele of HO [i.e., ho]. To rule out the possibility that ho contributes some component essential for switching, which may regulate the mother–daughter asymmetry, the pattern of switching of strains K736 an K737 was determined. Both the latter strains contain the hoA::TRPI allele in which most of the 5’ coding region and part of the HO promoter region are deleted [Nasmyth 1985]. Again, results presented in Table 2 demonstrate that only mother cells of these strains switched regardless of the HO orientation.

Furthermore, strains K738 and K739, both of which contain HO at both CDC36 and its indigenous location, also exhibit the wild-type pattern of switching [Table 2].

Taken together, these results demonstrate that the pattern of mating type switching, and thus the pattern of HO expression, remains unaffected by [1] the HO orientation in chromosome IV, [2] the particular location of HO in the chromosome, as well as [3] the presence of more than one copy of the HO gene in the cell.

Discussion

This paper addresses the question of the means by which
developmental asymmetry is generated between mother and daughter cells of yeast. The lineages of yeast cells are analogous to stem cell lineages; each cell division produces one cell incapable of switching and a second cell that has the competence to switch. Specifically, I tested whether this asymmetry is due to a strand-specific “imprinting” event at the HO gene (which confers to it the competence to express) and whether such a marked strand is always segregated to the mother cell. Apparently the SWI5 function is a key element in mother—daughter asymmetry and one suggestion for the mother cell specificity of HO transcription is the asymmetric inheritance of the SWI5 product (Nasmyth et al. 1987b; Sternberg et al. 1987). Here it was tested genetically whether the SWI5 gene product, or some other determinant, is distributed nonequivalently to progeny cells forming a stable transcription complex to the HO promoter by constructing strains in which the HO gene, along with its upstream regulatory sequences which are known to bind in vitro to the SWI5 product, are inverted in the chromosome. In such a strain the mother—daughter switching asymmetry remained unaltered. Thus, it is concluded that the competence for HO gene expression is not segregated with a particular strand of the HO gene.

The observed efficiency of switching of mother cells in strains where HO is placed next to CDC36 gene is reduced significantly as compared with those containing HO at its original location. Although a model invoking a position effect control may be entertained to explain the reduced efficiency of switching, it is also possible that HO and ho may compete for some regulator(s) whose concentration may be limiting, so that at low concentrations there is a reduced efficiency of switching. Consistent with this interpretation is the finding that those strains containing between two to three inserts of HO exhibit increased frequency of switching (nearly 2×). Interestingly, a further increase in their number adversely affects the efficiency (data not shown). Alternatively, sequences 5' to the SacI site may also be required for efficient transcription, because it is known that an insertion of a TRP1 fragment into the SacI site at HO reduces HO transcription (K. Nasmyth, pers. comm.). Nevertheless, these results show for the first time that the SacI—EcoRI fragment contains sequences that are sufficient for the whole repertoire of HO control, though possibly not for full-level transcription.

One important point regarding the conclusion drawn needs to be addressed here. An alternate model may be considered in which the hypothesized strand segregation machinery may operate by binding directly to the HO gene sequence. Specifically, such binding, for example, to the HO sense strand, may align chromosome IV on the division plate so that the chromatid containing the competent HO gene will always segregate to the mother cell. In this case inversion of the HO gene is not expected to change the mother—daughter asymmetry. However, such a model is unlikely for the following reason. Some of the strains we used to test the model also contain the naturally occurring HO on the same chromosome. Should the hypothesized strand segregation mech-

| Strain | Orientation of installed HO | Genotype |
|--------|-----------------------------|-----------|
| DC150  | —                           | ho MATa leu2 trp1 ura3 cry1 ade8 his-       |
| K734   | I                           | ho/ho MATa/MATa leu2/leu2 trp1/ trp1 ura3/ura3 cry1/1 cry1 ade8/ ade8 his- /his- |
| K735   | II                          | ho/ho MATa/MATa leu2/leu2 trp1/ trp1 ura3/ura3 cry1/1 cry1 ade8/ ade8 his- /his- |
| K736   | I                           | hoΔ : : TRP1/hoΔ : : TRP1 MATa/ MATa leu2/leu2 trp1/1 trp1 his- / his- ade-/ade- ura3/ura3 |
| K737   | II                          | hoΔ : : TRP1/hoΔ : : TRP1 MATa/ MATa leu2/leu2 trp1/1 trp1 ade-/ ade- ura3/ura3 can1/can1 |
| K738   | I                           | HO/ho MATa/MATa trp1/1 trp1 leu2/leu2/1 his-/his- ade-/ade- ura-/ura- cry1/cry1 |
| K739   | II                          | HO/ho MATa/MATa trp1/1 trp1 leu2/leu2/1 his-/his- ade-/ade- cry1/cry1 |

Figure 2. Southern blot analysis of BgIII-digested DNA from parental strain DC150 and two transformants [K734 and K735] that contain the HO gene inserted near the CDC36 locus (lanes a, b, and c, respectively). The probe consisted of the radio-labeled plasmid YRP7-CDC36.1 DNA (Breter et al. 1983). The small difference in the sizes of 7.0-kb fragment in lane a and that of 6.9 kb in lane b is not apparent in this figure, but the difference is clear when the DNA is separated by electrophoresis for a longer time [data not shown]. The fragment corresponding to the TRP1 sequence present in the YRP7-CDC36.1 probe was run off the gel because of its small size. The numbers indicate DNA fragment size in kilobase pairs.
Thus, one of these inserts must be oriented opposite to that of the naturally occurring HO gene. On the basis of meiotic recombination mating-type switching however, are conducted on mitotically growing cells that undergo very low levels of recombination. Consequently, for our purpose these elements are tightly linked. The centromere is represented by the circle.

Anism operate by directly binding to the HO sequences, then in a strain in which HO is placed near CDC36 in an opposite orientation to that of naturally occurring HO, both chromosome IV sister chromatids should be delivered to the mother cell. This constraint should lead to death of daughter cells. No such lethality was noticed during the pedigree analysis. In such strains both HO genes express only in mother cells, as indicated by their increased efficiency of switching (Table 2) compared with strains that contain HO only at CDC36.

To explain the mother–daughter asymmetry, two other types of models have been proposed by others. The first involves asymmetric distribution of a regulatory protein (such as SWI5), the other involving some other difference between mother and daughter cells, such as the difference in length of the G1 phase of the cell cycle [Nasmyth et al. 1987b; Sternberg et al. 1987]. Future studies should help us to differentiate between these and other possibilities.

**Experimental procedures**

**Yeast strains**

All strains used are of *S. cerevisiae*. Construction of K734 and K735 is described in the text. The hoΔ::TRPI allele was described earlier by Nasmyth (1985). The HO orientation refers to the HO inserted near the CDC36 locus.

**Table 2. The effect of HO orientation on the pattern of mating-type switching**

| Strain | HO orientation | Switches per cell division |   |   |
|--------|----------------|----------------------------|---|---|
|        |                | mothers | daughters | spores |
| K734   | I              | 37/105  | 0/111     | 0/31 |
| K735   | II             | 78/286  | 0/286     | 0/41 |
| K736   | I              | 113/202 | 0/223     | 0/46 |
| K737   | II             | 27/63   | 0/72      | 0/13 |
| K738   | I              | 91/105  | 0/97      | 0/34 |
| K739   | II             | 54/75   | 0/70      | 0/33 |

**Genetic analyses**

All media for growth and sporulation and techniques for micro-manipulation and tetrad analysis have been described [Mortimer and Hawthorne 1969]. Pedigree analysis was conducted by a procedure described by Hicks and Herskowitz (1976). This procedure consists of testing the ability of cells to grow in the presence of an α-factor: the α cells grow while the α cells stop their growth and acquire a characteristic "shmoo" shaped structure. Pedigrees of α spores were followed for several generations by separating mothers from daughters by micromanipulation and observing the mating type by response to α-factor.

**Biochemical analysis**

DNA from yeast cells was isolated by the procedure of Cryer et al. (1975). Standard procedures for manipulating DNA in vitro were used [Maniatis et al. 1982]. For Southern blotting, 1 μg of DNA was digested with the restriction enzymes and then subjected to electrophoresis through a 0.8% agarose gel. Transfer of DNA from the gel to nitrocellulose paper was done essentially as described by Southern [1975]. The probes were nick-translated in the presence of α32P-labeled deoxyribonucleotides and then hybridized as described [Maniatis et al. 1982]. The DNA-mediated transformation was accomplished by the gusulase-generated spheroplast technique [Beggs 1978; Hinnen et al. 1978].

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**Note added in proof**

When HO contained in pAK37 construct was integrated at LEU2 in chromosome III, again only mother cells switched...
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[37% confirming that the mother-daughter asymmetry is indeed determined by sequences contained within the SacI–EcoRI HO fragment.

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