**Introduction**

The ends of eukaryotic chromosomes (telomeres) consist of simple DNA repeats associated with proteins and play important roles in replication, malignant transformation, and cellular aging (for reviews, see Blackburn, 1991; Zakian, 1995; de Lange, 1998a; Shore, 1998). Besides the vital functions telomeres perform in vegetative cells, they have been implicated as key players in the chromosome pairing process during meiosis (Moses, 1968; Loidl, 1990; Dernburg et al., 1995; Roeder, 1997; de Lange, 1998b; Zickler and Kleckner, 1998). The pairing of homologues and their recombination during meiotic prophase is critical for reduction of DNA content from diploid to haploid before fertilization (von Wettstein et al., 1984; Hawley, 1988). In many organisms, the premeiotic (vegetative) telomeres fail to cluster their telomeres at any prophase stage. Ndj1p is the first protein shown to be required for bouquet formation in a synaptic organism. Analysis of homologue pairing by two-color fluorescence in situ hybridization with cosmid probes revealed that disruption of NDJ1 is associated with a significant delay (>2 h) of homologue pairing. An increased and persistent fraction of ndj1Δ meiocytes with Zip1p polycytes suggests that chromosome polarization is important for synopsis progression. Thus, our observations support the hypothesis that meiotic telomere clustering contributes to efficient homologue alignment and synaptic pairing. Under naturally occurring conditions, bouquet formation may allow for rapid sporulation and confer a selective advantage.

Key words: bouquet fluorescence in situ hybridization • ndj1 • chromosome pairing • meiosis • telomeres
see Bélar, 1928; Dernburg et al., 1995; Hiraoka, 1998). Genetic and cytological analyses of the meiotic prophase of Saccharomyces cerevisiae have suggested that the formation of a chromosomal bouquet in the synaptic meiosis of budding yeast (Dresser and Giroux, 1988; Trelles-Sticken et al., 1999) occurs independent of homologous recombination and synopsis (Rockmill and Roeder, 1998; Trelles-Sticken et al., 1999). In the vegetative (premeiotic) yeast nucleus, telomeres associate in a few aggregates at the nuclear periphery (Klein et al., 1992; Gotta et al., 1996; Laroche et al., 1998), while centromeres form a single cluster near the SPB (Hayashi et al., 1998; Jin et al., 1998). During the leptotene/zygotene equivalent stage of yeast meiosis, centromeres disperse from the SPB (Hayashi et al., 1998; Jin et al., 1998) and telomeres cluster de novo at this location (Trelles-Sticken et al., 1999). Hence, the bouquet stage of yeast resembles the classical bouquet arrangement formed during prophase I of multicellular eukaryotes (Gelei, 1921; Moens, 1974; Scherthan et al., 1996; Bass et al., 1997).

Genetic analysis of disomic S. cerevisiae strains has indicated that telomeres play an important role for homologue search and alignment in synaptic meiosis (Rockmill and Roeder, 1998). The only telomeric protein that has been shown to be involved exclusively in meiosis of budding yeast is Ndj1p/Tam1p (Chua and Roeder, 1997; Conrad et al., 1997). Ndj1p is expressed specifically after induction of meiosis. Ndj1p-deficient cells show reduced efficiency in homologue disjunction, increased occurrence of nonrecombinant chromosomes (a defect in crossover interference), and a delayed progression through meiotic prophase (Chua and Roeder, 1997; Conrad et al., 1997). Furthermore, spread meiocytes show a disordered distribution of telomeric Rap1p (Conrad et al., 1997). In the present study, we investigate the extent to which bouquet formation, telomere positioning, and chromosome pairing are affected in ndj1Δ meiosis and cytologically identify Ndj1p as the first telomeric protein demonstrated to be required for bouquet formation in a synaptic organism.

Materials and Methods

Yeast Strains

To exploit the relatively synchronous progress of SK1 meiosis (Padmore et al., 1991; Trelles-Sticken et al., 1999), the NDJ1 locus was disrupted in the S. cerevisiae SK1 strain (Kane and Roth, 1974) background by replacing the sequence coding for amino acids 13–252 with the KanMX4 cassette (Wach et al., 1994). SK1 yeast transformants resistant to G418 were isolated by A. Goldman and M. Lichten and were assigned the strain name MDY1490 (MATα ura3 lys2 leu2 his3::LYS2 ndj1::KanMX6 and MDY 1493 (MATa ura3 lys2 trp1 his3::LYS2 ndj1::KanMX6). These strains were mated and diploids were selected on the appropriate media (trp–,leu–, ura–, his3–). Wild-type SK1 haploids (MDY 1484: MATα ura3 lys2 leu2 his3::LYS2, MDY 1487: MATα ura3 lys2 trp1 arg4 his3::LYS2) were mated to result in a diploid wild-type control strain. These strains were used in all fluorescence in situ hybridization (FISH) experiments.

Immunolocalization of telomeres by immunofluorescence to hemagglutinin (HA)-tagged Ndj1p was done in a diploid strain derived from MDY 431 and MDY 433 (Dresser et al., 1997).

Cell Culture and Preparation

For nuclear preparation, cultures were grown in precopulation medium to a density of 2 × 10^7 cells/ml, and then transferred to sporulation medium (2% KAc) at a density of 4 × 10^7 cells/ml (Roth and Halvorson, 1969). Aliquots from the sporulating cultures were obtained during time course experiments at induction of meiosis (transfer to sporulation medium = 0 min) and from 180 to 420 min or 20-min intervals. Aliquots were immediately transferred to tubes on ice containing 1/10 vol acid-free 37% formaldehyde (Merek). After 30 min, cells were removed from fixative, washed with 1× SSC and spheroplasted with Zymolyase 100T (100 μg/ml; Seikagaku) in 0.8 M sorbitol, 2% KAc, and 10 m M dithiothreitol. Spheroplasts were terminated by adding 10 vol ice-cold M sorbitol. To allow for the delay in meiotic prophase progression in ndj1Δ, we additionally sampled at 5, 6, and 7 h. Spheroplasts were subdivided in two aliquots and subjected either to nuclear spreading (Loidl et al., 1998) or to preparation of structurally preserved nuclei. The latter were obtained as described by Trelles-Sticken et al., 1999 and subjected to FISH.

DNA Probes and Labeling

A composite pancentromeric DNA probe was used to delineate all yeast centromeres (Jin et al., 1998). One plasmid containing a conserved core fragment of the subtelomeric X element and one containing the Y′ element (Louis et al., 1998) were used to probe all yeast telomeres (Gotta et al., 1996; Trelles-Sticken et al., 1999). The following cosmid probes (Fig. 1) were used to determine the pairing of homologous chromosome regions: the telomere-adjacent region on the right arm of XI was probed with a cosmid (cos) located at 628.5–665.8 (cos l; pUKG066). The smaller chromosomes IX and III were tagged with cosmid probes hybridizing internally on the left arm of chromosomes IX (cos p; ATCC 70095) and at HIR1, near the left telomeres of chromosomes III (cos m; ATCC 70884). An internal chromosome XI cosmid (pEKG151, cos f; Thierry et al., 1995) mapping 231.8–264.9 on the left arm of chromosomes XI was used to determine meiotic pairing at a telomere-distant chromosomal region (Trelles-Sticken et al., 1999). Chromosome condensation was measured by determining the distance between cosmid b (pUKG 040) and cos h (pEKG 011) on the left arm of XI (Fig. 1; Thierry et al., 1995). Chromosome morphology was monitored by FISH with a painting probe for chromosomes XI (Trelles-Sticken et al., 1999), which covers 338 of the 666 kbp large chromosomes XI (Dujon et al., 1994).

Probes were labeled either with dig-11-dUTP (Roche Biochemicals) or with biotin-14-dCTP (Life Technologies) using a nick translation kit, according to the supplier’s instructions (Life Technologies).

Fluorescence In Situ Hybridization

All preparations were subjected to two-color FISH as described previously (Scherthan et al., 1992; Trelles-Sticken et al., 1999). The hybridization solution contained various differentially labeled probe combinations. It contained the yeast pan-telomere probe, which delineates all telomeres in the SK1 strain (2n = 32) investigated (Jin et al., 1998) and one of the following probes: (a) chromosome-specific cosmid probes (not shown), (b) a chromosomes XI paint probe, and (c) a pan-centromeric DNA probe (see Jin et al., 1998). Analysis of pairing of homologous regions was done by two-color FISH to spread nuclei using pairwise combinations of differentially labeled cosmid probes. Immunofluorescent detection of hybrid molecules was carried out with Avidin-FITC (Sigma-Aldrich) and rhodamine-conjugated sheep anti–dig Fab fragments (Roche Biochemicals) (for details, see Scherthan et al., 1992). Before microscopic inspection, preparations were embedded in antifade medium (Vector Laboratories) containing 0.5 μg/ml DAPI (4′,6-diamidino-2-phenylindole) as DNA-specific counterstain.

Immunostaining

A polyclonal antisera against a S. cerevisiae spindle pole body component (Tub4; Marshall et al., 1996) was used to stain the SPB in conjunction with telomeres (for details, see Trelles-Sticken et al., 1999). A rabbit antisera against Zip1p transverse filament protein (Sym et al., 1993; a gift of S. Roeder, Yale University, New Haven, CT) of the yeast synaptonemal complex was applied to identify nuclei with synopsis in progress. Ndj1p was stained in freshly prepared, mildly spread nuclei obtained from a strain that expresses HA-tagged Ndj1p (Conrad et al., 1997) using a monoclonal anti-HA–tag antibody (Biotec Santa Cruz) and secondary anti-mouse Cy3-conjugated antibodies (Dianova).

Rap1p immunostaining of wild-type and ndj1Δ diploid cells in the MDY strain background (Dresser et al., 1994) was performed as follows: wild-type and mutant cells were harvested at 4, 7, and 12 h after the shift into sporulation, fixed, spheroplasted, adhered to poly-L-lysine–coated coverslips, and then prepared for immunolabeling by blocking with PBS/
0.05% Tween 20%/4% nonfat dry milk. Preparations of three-dimensionally preserved cells were labeled with a rabbit anti–Rap1p serum (gift of J. Berman, University of Minnesota, St. Paul, MN) diluted 1/10,000, followed by Oregon green-labeled anti–rabbit secondary antibodies (Jackson ImmunoResearch Laboratories), and then mounted with Citifluor anti-fade solution (Ted Pella, Inc.) containing 0.5 μg/ml DAPI under a coverslip.

**Microscopic Evaluation**

Preparations were evaluated using an epifluorescence microscope (Axioskop; Carl Zeiss, Inc.) equipped with a double band-pass filter for simultaneous excitation of red and green fluorescence, and single band-pass filters for excitation of red, green, and blue (Chroma Technologies). To allow for the evaluation of a large number of nuclei from the time course experiments, investigation of structurally preserved in situ-hybridized nuclei was performed by careful focusing through the nuclei using a 100× plan-neofluor lens (Carl Zeiss, Inc.). Signal patterns in spread nuclei were investigated with the same setup. Digital images were obtained using a cooled grey-scale CCD camera (Hamamatsu) controlled by the ISIS fluorescence image analysis system (MetaSystems). More than 100 nuclei were scored for each time point and probe combination. Fluorescence signal patterns were analyzed in nuclei with an undisturbed, homogeneous appearance in the DAPI-image.

**Confocal Microscopy and Quantitative 3-D Distribution Analysis of Telomeres**

Three-dimensionally preserved, Rap1p-stained nuclei were subjected to 3-D microscopy using a Meridian Ultima Z laser-scanning confocal microscope equipped with lasers to image DAPI and fluorescein. Simultaneous two-color scans were made of 10 randomly selected meiotic nuclei from each time point, without regard to the distribution of Rap1p signal, at a pixel and slice step-size of 0.1 μm. An aperture of 100 μm was used with short exposure times to minimize fluorescence fading. Appropriate wavelength scans were made of four fluorescent point source beads (Molecular Probes, Inc.) using the same settings. To avoid any influence of nonrandom telomere clustering that occurs during the bouquet stage, nuclei with less than five signal spots upon visual inspection were excluded from scoring. Averaged point source images were used to deconvolve the experimental images using the freeware xcosm v2.1 EM routine (Conchello et al., 1994), mainly to reduce noise in the images, before image analysis.

Image analysis employed Pascal macros running Image (The National Institutes of Health, Bethesda, MD) and free-standing C++ routines written by M.E. Dresser. To measure the distances between the Rap1p spots and the DAPI-defined nuclear periphery: (a) the nuclear periphery was defined by thresholding the DAPI images over a range of different values, where increasingly large values exclude an increasing number of Rap1p spots from the inclusion volume, (b) the coordinates of each Rap1p spot were defined as lying at the center of each distinct signal spot (scored manually using Image and blinded with respect to the location of the nuclear periphery), and (c) the distance between each spot and the threshold-defined periphery was determined by a program that scores the radius of the smallest sphere, centered on the spot, which intersects an “interior” voxel for spots included in the volume, or which intersects an “interior” voxel for excluded spots; if adjacent, the distance is defined as 0.

To unequivocally determine the relation of the Rap1-FITC signals with the nuclear periphery, we determined the nearest distance from Rap1-FITC signal centers and the nuclear boundary. The latter was determined from the optical section of the DAPI-stained nucleus from the same image plane. All signal spots analyzed were requested to lie with their center in the equatorial plane (within ±0.1 μm of best focus, as determined from the confocal image stacks of sectioned nuclei). The distance of signal gravity centers to the nearest edge of the nucleus was measured using a dedicated program (M.E. Dresser, unpublished data).

**Results**

Induction of meiosis leads to a dramatic reorganization of yeast nuclear architecture. Centromeres that are tightly clustered at the SPB during vegetative growth disperse throughout the nuclear volume (Hayashi et al., 1998; Trelles-Sticken et al., 1999), while telomeres accumulate de novo at this location (Trelles-Sticken et al., 1999). To determine whether this dramatic reorganization of the early meiotic nucleus requires the Ndj1 protein (Chua and Roeder 1997; Conrad et al., 1997), an Ndj1 null-mutation (ndj1Δ::kanMX4) was introduced into the SK1 strain background because the high synchrony in this strain background facilitates bouquet analysis (Trelles-Sticken et al., 1999). Telomere clustering, centromere distribution, and chromosome pairing were studied by FISH with telomere-, centromere-, and chromosome-specific probes in detergent spread and in three-dimensionally preserved nuclei from meiotic time-course experiments of diploid wild-type and ndj1Δ strains. Additionally, using anti–Rap1p signals as marker for meiotic telomeres (Klein et al., 1992; Conrad et al., 1997), telomere positioning with respect to the nuclear periphery was examined by confocal microscopy in the MDY strain background. All experiments analyzed displayed sporulation rates ≥85%.

**Dissolution of Premeiotic Centromere and Telomere Clusters Does Not Require Ndj1p**

It has been shown that the nuclear organization of vegetative/premeiotic yeast cells is dominated by centromere clustering at the SPB (Fig. 2, a and b), which resolves during the onset of meiotic prophase (Fig. 2, c and d) (Hayashi et al., 1998; Jin et al., 1998). When we determined centromere distribution by FISH to nuclei of SK1 strains, it was found that the frequency of nuclei with one centromere cluster diminished at similar rates after induction of meiosis, both in wild-type and in ndj1Δ meiotic time courses (Fig. 3, and not shown). This suggests that dissolution of the centromere cluster is not affected by the absence of Ndj1p.

Premeiotic (vegetative) yeast nuclei usually contain few (two to eight) perinuclear telomere clusters (Klein et al., 1992; Gotta et al., 1996) that are resolved at the onset of meiotic prophase (Fig. 2, e and d) (Hayashi et al., 1998; Trelles-Sticken et al., 1999). To monitor whether the absence of Ndj1p influences dissolution of vegetative telomere clusters, we determined the fraction of spread nuclei with two to eight telomere clusters at transfer to sporulation medium (0 min) and from 180–420 min, and in undisrupted nuclei at 0 and 180–480 min. Nuclei were also sampled at 530 and 590 min, following the shift into meiosis. It was found that the frequency of spreads with premeiotic telomere distribution (we consider nuclear topology at $t = 0$ to
represent premeiotic/vegetative nuclear organization) diminished at similar rates in wild-type and ndj1Δ meioses (Fig. 3). Similar results were obtained in repeated time courses and with undisrupted nuclei (not shown). These data suggest the dissolution of premeiotic nuclear architecture during the onset of sporulation occurs at wild-type rates in the absence of Ndj1p. To further determine whether the ndj1Δ mutation affects telomere distribution in vegetative nuclei, we compared telomere FISH signal numbers in 20 randomly selected nuclear spreads obtained during logarithmic vegetative growth in YPD. Spreading of vegetative nuclei dissociates the few clusters seen in intact nuclei to a mean of $22 \pm 5$ (SD) and to $21 \pm 3$ telomere signal spots/nucleoid in wild-type and ndj1Δ cells, respectively. Similar telomere signal numbers in spread premeiotic wild-type and ndj1Δ cells suggest that telomere distribution in vegetative cells is not affected by the ndj1Δ mutation, which contrasts with the situation in meiocytes (see below).

**NDJ1 Is Required for Meiosis-specific Telomere Distribution**

The induction of meiosis leads to the repositioning of telomeres over the nuclear periphery seen as peripheral dispersion of RAP1p-GFP foci in live cells (Hayashi et al., 1998) and by telomere FISH signals in fixed nuclei (Trelles-Sticken et al., 1999). To determine whether the telomere distribution patterns obtained by FISH are representative for meiocytes and to see whether dispersion of meiotic telomeres over the nuclear periphery occurs during sporulation of our strains, we determined the telomere distribution patterns in spread and undisrupted nuclei from time-course experiments after induction of meiosis by FISH with the XY9 telomere probe and performed immunofluorescence (IF) staining of HA-tagged Ndj1p in spread nuclei at $t = 240$ min.

Figure 2. FISH with a pan-centromere probe (fluorescein, green) and a pan-telomere probe (rhodamine, red) to undisrupted diploid nuclei (DAPI, blue) of wild-type and ndj1Δ SK1 cells. (a) Vegetative wild-type nucleus before induction of meiosis ($t = 0$ min) displays clustered centromeres that form a single green signal, while telomeres form four perinuclear clusters. (b) Premeiotic ndj1Δ nucleus with one centromere cluster and three large telomere clusters at the opposite pole of the nucleus, thereby resembling a Rabl orientation. (c) WT and (d) ndj1Δ meiocyte nuclei from two later time points (200 and 260 min, respectively) displaying centromere and telomere signals distributed throughout the nuclei. Few centromere signals are dissociated from the centromere cluster in nucleus (c), while in nucleus (d) exhibits dispersed centromere signals. Bar, 5 μm.

Figure 4. (a–d) Double immunolabeling of SPB components (fluorescein, green) and of meiotic telomeres with antibodies against HA-tagged Ndj1p (rhodamine, red) reveals meiosis-specific telomere distribution patterns in mildly spread diploid wild-type SK1 nuclei. (a) Peripheral rim-like distribution of telomeres during early meiosis (140 min). (b and c) Meiocytes with telomeres accumulated at the SPB (bouquet arrangement). (d) Meiocyte nucleus from a later time point (240 min), which shows an SPB and dispersed telomere signals. (e–h) Mildly spread meiocyte nuclei from an independent FISH experiment with the XY9 repeat probe (see Materials and Methods) reveals telomere patterns similar to the ones obtained by Ndj1p IF. (e) Rim-like telomere distribution. (f and g) Bouquet nuclei with clustered telomere signals. (h) Advanced meiocyte from a later time point displays a scattered telomere distribution. Bar, 5 μm (applies to all details). The inset shows colocalization of Ndj1-HA IF signals (red) and XY9 telo-FISH signals (green) in a wild-type pachytene nucleus. Most of the Ndj1-HA and XY9 signals show significant overlap at chromosome ends (e.g., arrowheads). Ndj1 fluorescence is often seen beyond the telo-FISH signals and/or extends between telomere signals. Fewer IF signals in the Ndj1 channel may relate to loss of some epitopes during the FISH procedure.
Ndj1p is expressed only during meiosis (Chua and Roeder, 1997; Conrad et al., 1997); therefore, the distribution patterns detected with this probe exclusively represent meiotic telomere arrangements. The signal patterns obtained by Ndj1p IF matched the telomere distribution patterns revealed by telo-FISH (Fig. 4). Costaining experiments revealed colocalization of HA-tagged Ndj1p and telomere-FISH signals, with the Ndj1-HA signals often extending beyond the XY’ FISH signals (Fig. 4, inset). This could relate to the abundance of Ndj1p at chromosome ends during meiosis and/or to slight swelling of the epitope-bearing chromatin during the FISH procedure. The lack of IF at some telo-FISH signals may relate to loss of protein during the denaturation and hybridization procedure. In any case, we observed that meiotic telomeres, as marked by FISH as well as with anti-HA-Ndj1p, do adopt a peripheral dispersed arrangement in early meiocytes, which is seen as a rim-like fluorescent signal in mild spreads (Fig. 4, a and c) and at the equatorial focus plane of undisturbed nuclei (not shown). At later time points, telomeres congregate to form a single large signal cluster (bouquet arrangement; Fig. 4, f and g) at the SPB (Fig. 4, b and c). This clustering is resolved as cells enter pachytene with telomere signals, again becoming dispersed (Fig. 4 d) (Trelles-Sticken et al., 1999).

Nuclei with a rim-like peripheral telomere signal distribution (Fig. 4 a) are rarely encountered at the induction of sporulation, but increase early after induction of meiosis in the wild type (Fig. 5). In contrast, the frequency of spread ndj1Δ meiocytes with such a preleptotene-like perinuclear telomere distribution (see Scherthan et al., 1996) essentially remained at a vegetative background level during sporulation, well below the level seen in the wild type (Fig. 5). Similar observations were obtained in time courses of undisturbed ndj1Δ nuclei (not shown).

**Perinuclear Distribution of Telomeres Is Altered in ndj1Δ Meiocytes**

To test whether the ndj1Δ mutation affects the meiosis-specific three-dimensional telomere distribution (i.e., peripheral location of dispersed telomeres in the meiocyte nucleus), we performed Rap1p staining with undisturbed nuclei and investigated the three-dimensional spot distribution by laser scanning microscopy and digital image analysis. Telomeric Rap1p signal spots and the corre-
sponding DAPI-stained nuclei were obtained by light optical serial sectioning (Fig. 6). Because nuclei in meiotic prophase are nonspherical and nuclear pore staining failed to reveal the typical rim staining seen in mitotic interphase (not shown), we determined the nuclear boundary in light optical sections of DAPI-stained nuclei. DAPI highly selectively stains DNA and renders the nuclear boundary at a high contrast. The number and relation of Rap1p signal centers to the nearest sector of the nuclear boundary was computed from 30 nuclei of uninucleate cells, 10 each at 4, 7, and 12 h after induction of meiosis from wild type and ndj1Δ. Bouquet nuclei, which rarely exceed 5–10% at any time point in this strain background, were excluded from this analysis (see Materials and Methods). The 30 prophase nuclei from wild-type and ndj1Δ cells were analyzed as a single class, where totals of 519 and of 729 Rap1p spots were scored, respectively. The difference in the number of total spots is consistent with the smaller number of larger Rap1p signals seen in spread meiotic prophase nuclei of wild type, compared with ndj1Δ cells (Conrad et al., 1997).

The relation of the Rap1p telомер signals to the nuclear periphery was determined in the light optical section of the equator of each nucleus. The distance from Rap1/ FITC signal centers to the nearest nuclear boundary of the DAPI image was calculated by a dedicated program. This two-dimensional procedure accounts for the small dimensions of the yeast nucleus and its irregular shape at meiosis (see Zickler and Olson, 1975), which precludes applying a distribution analysis of signal spots from a geometrically defined center. The data obtained are displayed in a bar graph according to frequency of signals at distance units of of 0.2 μm from the nearest nuclear boundary segment (Fig. 7). In wild-type cells, the vast majority of telomeric spots were found near the nuclear periphery (Figs. 6 and 7). A deviation from this peripheral telomeric Rap1p distribution in wild-type meiocyte nuclei was apparent in ndj1Δ nuclei. In the latter, a significant fraction of signals (97.5% level by G test) located remote from the nuclear boundary (i.e., 34% of signal spots were located 0.6–1 μm from the nearest nuclear edge), while in the wild type only 18% of spots were found in this more internal nuclear compartment (Fig. 7). These results suggest that the perinuclear localization of meiotic telomeres is disrupted in ndj1Δ meiosis.

Ndj1p Is Essential for Formation of the Bouquet

In the SK1 background, we observed that the premeiotic telomere distribution is resolved in diploid ndj1Δ cells at wild-type rates. However, a rim-like perinuclear telomere distribution was detected only at insignificant rates (Fig. 5). This prompted us to also determine the frequency of meiocytes with a single telomere-FISH signal cluster, a telomere distribution diagnostic for a chromosomal bouquet (Fig. 4, f and g) that transiently forms at the leptotene/zygotene transition during prophase I of S. cerevisiae (Trelles-Sticken et al., 1999). Induction of meiosis in the wild type led to a 22-fold increase in the frequency of mildly spread nuclei with a single telomere cluster over premeiotic background at t = 240 min (Fig. 5). In undisrupted nuclei, a 5.3-fold increase at 260 min (not shown) was noted, which is well in agreement with earlier observations in undisrupted wild-type SK1 meiocytes (Trelles-Sticken et al., 1999).

A severe deviation from the wild-type situation was observed in ndj1Δ meiosis. The frequency of nuclei with a single telomere cluster never exceeded premeiotic levels, even in time courses with prolonged duration and in time-course experiments where we performed FISH to intact meiocyte nuclei (Fig. 5, and not shown). The defect in telomere clustering was also evident when telomere FISH signal numbers were compared in 20 randomly selected wild-type and mutant nuclei. Spread ndj1Δ meiocytes (obtained 300 min after induction of meiosis) contained a significantly increased spot number/nucleus as compared with wild-type meiocytes that were sampled at 200 min to compensate for the delay in mutant prophase I (Fig. 8). Similar results were obtained using anti–Rap1p IF and confocal microscopy (data not shown).

Since formation of a true bouquet involves telomere clustering at the SPB during the leptotene/zygotene transition stage (Trelles-Sticken et al., 1999), we determined whether SPB/telomere cluster association is absent in ndj1Δ meiocytes. To this end, we simultaneously immunostained for spindle pole body and telomeres by FISH in intact nuclei at 240 min in wild-type and mutant meiosis. We failed to detect a physical association of the SPB signal and XY’ telomeric signal accumulations seen in a few ndj1Δ nuclei (not shown). This and the constant and negligible frequency of nuclei with a single telomere-FISH signal cluster over mutant time courses corroborates that bouquet formation is impaired in ndj1Δ meiosis. To further investigate whether telomere clustering is defective during the leptotene/zygotene transition stage, we immunostained spread meiocytes with antibodies to Zip1p, a

Figure 7. Spatial analysis of Rap1p telomere signal distribution with respect to the nearest nuclear boundary segment. The frequency (%) of occurrence of Rap1/FITC signal centers situated at particular distances (given in 0.2-μm intervals) from the nearest nuclear periphery segment as determined at the equatorial plane from the corresponding confocal DAPI images. The values were derived from 146 signal spots for ndj1Δ (ndj1), and 86 for wild type (WT). In the wild type, most signal spot centers locate in the immediate vicinity of the nuclear periphery (open bars). An altered spatial distribution of telomeric Rap1p signals in ndj1Δ nuclei is reflected by a higher portion of telomeric signals being more distant from the nuclear border (solid bars).
component of transverse filaments of the synaptonemal complex (Sym et al., 1993) and determined the telomere distribution by FISH in SK1 meiocytes, obtained at \( t = 200 \) min with an incomplete speckled Zip1p distribution, which represent nuclei with synapsis in progress. It was found that telomere signals were scattered throughout mildly spread \( ndj1^{-} \) nuclei (\( n = 168; \) obtained at 300 min) with a fragmented Zip1p distribution (zygotene-equivalent stage; Fig. 9, a–c), while in the wild type 20% of zygotene cells (\( n = 166 \)) contained a single telomere cluster (see also Trelles-Sticken et al., 1999). These data suggest that true bouquet formation is disrupted in \( ndj1^{-} \) meiosis.

Zip1p IF disclosed furthermore that \( ndj1^{-} \) nuclei contained an unusually high frequency of intensely staining Zip1-positive structures that have been called “polycomplexes” (Sym and Roeder, 1995). At 210 min after induction of meiosis, 81% of \( ndj1^{-} \) nuclei contained predominantly one Zip1 polycomplex (PC) in the form of a brightly stained rod (Fig. 9, a and b), while at this time point only 28% of wild-type nuclei contained a PC. This represents an approximately threefold increase of PCs in the mutant. At 330 min, 60% of \( ndj1^{-} \) meiocytes still contained PCs, while these were absent in the wild type. In yeast meiosis, formation of Zip1 polycomplexes has been observed in a variety of conditions and strain backgrounds (see Sym and Roeder, 1995). An increased frequency of PCs is often seen in recombination mutants, which generally show defects in synapsis (e.g., Alani et al., 1990; Loidl et al., 1994; Sym and Roeder 1995; Grushcow et al., 1999). Taken together, these data suggest that normal synaptic progression requires Ndj1p and bouquet formation.

**Chromosome Pairing Is Impeded in \( ndj1^{-} \) Meiosis**

Since a mutant with disrupted bouquet formation offers the possibility to test for the impact of meiotic telomere clustering for the homologue pairing process, we investigated homologue pairing by FISH with cosmid probes to regions on chromosomes III, IX, and XI (Fig. 1) during wild-type and mutant sporulation. While chromosomes XI represents a large chromosome that is capable spanning the entire nuclear volume, IX is of intermediate size, and III is among the smallest yeast chromosomes. Due to the clustering of vegetative centromeres at the spindle pole body (Hayashi et al., 1998, Jin et al., 1998) and other functions, the yeast nucleus displays a substantial amount of premeiotic homologue association (Loidl et al., 1994; 1999).
With estimations reported by others (see Burgess et al., 1999). To adjust for accidental contacts between cosmid signals, which may be influenced by a number of parameters (see above), we used centromere distant probes and furthermore subtracted 50% of the obtained fraction showing heterologous contacts from the obtained pairing values, since in two cosmid FISH experiments a signal has a two- to fourfold higher probability to be associated with a heterologous than with a homologous signal. It should be noted that the results obtained with and without such a correction remained essentially the same (not shown).

In wild-type and ndjlΔ time courses, the fraction of nucleoids with FISH signals homologously paired increased well above premeiotic values (Fig. 11). In ndjlΔ meiosis, all homologous regions investigated reached nearly wild-type levels of homologous signal pairing, but with a 2–3-h delay (Fig. 11). This slowed progression of homologous pairing is mirrored by a 2-h delay in the appearance of anaphases I and II in ndjlΔ meiosis as compared with the wild type, where these first appeared at 300 and 320 min, respectively (not shown, consistent with earlier reports). In wild-type meiosis, the homologous region near the left telomere of the small chromosomes III paired most rapidly (Table I). However, in the absence of Ndj1p and bouquet formation, the telomeric region of chromosomes III showed the most prominent retardation of homologous pairing, expressed as the difference in time required in wild-type and ndjlΔ meiosis to reach peak values in signal pairing (Δt = 160 min, Fig. 11). Pairing of the telomeric region of the right arm of the large chromosomes XI (cos I), in contrast, showed only a delay of 100 min in ndjlΔ meiosis. Pairing of large chro-

![Figure 11. FISH analysis of homologous pairing during meiotic time courses of wild-type and ndjlΔ SK1 strains. Cosmid probe combinations m/p and f/l were hybridized to spread preparations obtained at the respective time points (minutes). Pairing values were obtained by determining the fraction of nuclei containing cosmid signals of the same color that touched each other or showed an enlarged coalesced signal. More than 200 FISH signal pairs were scored per time point, probe combination, and strain. Values were corrected for accidental heterologous contacts by subtracting 5% for cosmid combination m/p and 4% for probe combination U/l (see text). (A) Frequencies of nuclei with paired cos f signals (XI internal; Fig. 1) in wild-type (WT) and ndjlΔ time courses. (B) Frequencies (%) of nuclei with paired cos m signals (III, HML) in wild-type and ndjlΔ time course. At all loci probed, the frequencies of nuclei with paired signals increase more gradually in the absence of Ndj1p, reaching nearly wild-type frequencies with a 2–3-h delay. In the wild type, signal patterns are displayed only until 320 min, since meiotic divisions (anaphases) appeared after 300 min and complicated the signal analysis.
mosomes may thus be less dependent on a catalytic action of telomere clustering on homologue pairing, since these may span the entire nucleus and therefore have a higher probability for chance encounters with their homologues in the absence of bouquet formation.

Condensation of Chromosome Territories Occurs in the Absence of Ndj1p

Previous chromosome painting studies have shown that yeast chromosomes extend, pair, and condense during the course of meiotic prophase (Scherthan et al., 1992; Trelles-Sticken et al., 1999). To test whether the ndj1Δ mutation influences this morphological change of yeast chromosomes during meiosis, we painted chromosomes XI in conjunction with telomere FISH in wild-type and ndj1Δ cells at 180, 240, 360, and 420 min after induction of meiosis. Nuclei with all aspects of meiotic chromosome morphology were detected in wild-type and ndj1Δ cells (Fig. 12). The painting of yeast chromosomes also allows chromatin condensation to be assessed. The fraction of uniconnected meiocytes that exhibit clear and compact FISH signals represents the fraction of cells in which chromosomes are condensed (pachytene; Scherthan et al., 1992; Loidl et al., 1994; Nag et al., 1995). The frequency of condensed XI pachytene bivalents was determined in uninucleate meiocytes obtained at 0 min in wild type and 240 min in ndj1Δ nuclei.

### Table 1. Retardation of Homologue Pairing if ndj1Δ Meiosis

| Cosmid probe | Maximal signal pairing in WT at t | Maximal signal pairing in ndj1Δ meiosis at t | Δt |
|--------------|----------------------------------|-------------------------------------------|----|
|              | min                              | min                                       | min |
| Cos m (III) HML | 240                              | 400                                       | 160 |
| Cos p (IX, right arm) | 300                              | 400                                       | 100 |
| Cos f (XI, interstitial) | 280                              | 400                                       | 120 |
| Cos f (XII, right telo) | 280                              | 380                                       | 100 |

Time taken in wild-type and ndj1Δ meiosis to reach peak values in homologous FISH-signal pairing (minutes). Δt, time difference between the peak homologue pairing values for a given cosmid probe in wild-type and ndj1Δ meiotic time courses.

The painting of yeast chromosomes also allows chromatin condensation per se to be assessed. The difference between the two data sets did not differ significantly (P = 0.23 at 0.01; Student’s t test). This further suggests that chromosome condensation is not affected by the absence of Ndj1p.

### Discussion

**Ndj1p May Be Required for Peripheral Localization of Meiotic Telomeres**

Chromosome topology of the vegetatively growing yeast is dominated by tightly clustered centromeres next to the spindle body (Hayashi et al., 1998; Jin et al., 1998) and by telomeres forming a few aggregates at the nuclear periphery (Klein et al., 1992; Gotta et al., 1996). Upon induction of meiosis, nuclear topology is reorganized. Centromeres become dispersed throughout the nucleus, and telomeres spread over the nuclear periphery (Hayashi et al., 1998), except during an intermediate stage when telomeres form a tight cluster in the vicinity of the spindle pole body; i.e., at the bouquet stage (Trelles-Sticken et al., 1999).

Two of the early steps in the meiotic nuclear reorganization, dispersion of centromeres and telomeres, apparently are unaffected by deletion of NDJ1. However, this close proximity of dispersed telomeres to the nuclear periphery, which is evident in wild-type cells (Hayashi et al., 1998, and this report) does appear defective in ndj1Δ. During earliest meiotic prophase, telomeres dissociate from the vegetative aggregates and line the nuclear periphery (Fig. 14), giving rise to a rim-like staining in ∼20% of wild-type nuclei, which resembles the telomere distribution in wild-type nuclei with scattered chromosome XI territories and several telomere clusters. (b, wt) WT meiocyte nucleus with extended chromosome XI signal tracks and a rim-like distribution of telomere signals. (b, ndj1) Meiocyte with scattered telomere signals, while

![Figure 12](image-url). Representative images of FISH signal patterns obtained with a painting probe for chromosomes XI (red) and the pan-telomere probe (green) in mildly spread nuclei from meiotic time courses of diploid wild type (wt) and ndj1Δ (ndj1) SK1 strains (a, 0 min; b and c, 260 min; d, 380 min). Images are aligned according to meiosis-specific changes in chromosome morphology during prophase I (see Trelles-Sticken et al., 1999). (a) Premiotic wild-type and ndj1Δ nuclei (0 min) display separated variably shaped chromosomes XI territories and several telomere clusters. (b, wt) WT meiocyte nucleus with extended chromosome XI signal tracks and a rim-like distribution of telomere signals. (b, ndj1) Meiocyte with scattered telomere signals, while

- **XI** signal tracks extend across the nucleus and touch at one end. (c, wt) Nucleus with one large telomere signal cluster (bouquet arrangement). Chromosomes XI form one outstretched signal track along the telomere cluster. (c, ndj1) ndj1Δ meiocyte with scattered telomeres. Painted chromosomes XI are seen as a single extended signal track. (d) Wild-type and mutant pachytene nuclei both exhibit a condensed XI bivalent and scattered telomeres. Bar, 5 μm.
seen in the late preleptotene/leptotene stage of mamma- 
lian prophase I (Scherthan et al., 1996). In mammals, the 
switch from premeiotic to meiotic telomere distribution is 
a two-step process, with telomeres first attaching at scat-
tered points over the nuclear envelope, and then congre-
gating in the bouquet to a limited sector of the nuclear 
envelope (Rasmussen and Holm, 1978; Boiko, 1983; 
Scherthan et al., 1996). In

Distribution of recombination events (crossover interfer-
ce), chromosome segregation, and distributive disjunc-
tion (Chua and Roeder, 1997; Conrad et al., 1997) are all 
defects are relatively mild and may arise from a telomere-
related delay in meiotic prophase (Chua and Roeder, 
1997). The delay in onset and completion of synopsis 
(Conrad et al., 1997) potentially is due to a defect in some 
aspect of chromosome pairing that normally brings homo-
logues into close apposition for the initiation of homosyn-
apsis. FISH in conjunction with Zip1p immunostaining 
showed that telomeres fail to cluster during the leptotene/ 
zygotene-equivalent stages, when bouquet formation nor-
mally occurs (Dernburg et al., 1995; Zickler and Kleckner, 
1998). Moreover, the meiotic SPB failed to show an associ-
ation with a telomere accumulation occasionally seen in 
Ndj1-deficient nuclei. This suggests that loss of bouquet 
formation is the underlying defect of the retarded synapsis 
progression and chromosome pairing in the

When we investigated homologue pairing at four re-
gions on three chromosomes by two-color cosmid FISH, a 
highly significant reduction in chromosome pairing was 
noted at time points in mutant meiosis when, in the wild 
type, signal pairing had reached its maximum. Prolonged 
time courses in the

cluster at the spindle pole body, while centromeres are dispersed. (d) While premeiotic telomere topology is not affected, the early steps of meiotic telomere redistribution appear defective in the absence of Ndj1p, which leads to scattering of both telomeres and cen-
tromeres throughout the ndj1Δ meiocyte nucleus.
2–3-h delay in chromosome pairing. A similar delay was noted for the compaction of chromosomes XI bivalents, but, ultimately, chromosome condensation as measured by cosmid FISH appeared normal. When we compared the time required for the homologous regions assayed to reach meiosis-specific values in wild-type and ndj1Δ meiosis, it appeared that the pairing of the telomeric regions of the relatively small chromosomes III was most severely delayed. In Ndj1p-deficient meiosis, the left telomere region on chromosomes III took 60 min longer to reach peak-pairing values than the telomeric region of the right arm of the large chromosomes XI and on chromosomes IX. This may indicate that smaller chromosomes are more dependent on the action of telomere clustering to instigate homologue alignment and pairing. However, after a delay of ~2–3 h, all homologous regions investigated reached approximately wild-type pairing values, which shows that telomere-independent routes of homologue pairing exist. Altogether, it appears that bouquet formation is not required for homologue recognition and synopsis per se, but mediates an important catalytic action on homologue pairing, which shortens the duration of sporulation, and thus may, in turn, confer selective advantages in a natural environment. This conclusion is in agreement with long-standing hypotheses that telomere clustering during the bouquet stage facilitates homologue interactions by aligning chromosome ends (for reviews, see Rhoades, 1961; Moses, 1968; Scherthan, 1997; de Lange, 1998b; Zickler and Kleckner, 1998).

Furthermore, there is supporting experimental evidence from S. pombe (Chikashige et al., 1997) where, in analogy to the situation in budding yeast ndj1Δ mutants, absence of the telomere protein Taz1p(+) abrogates telomere clustering and leads to reduced recombination and increased meiotic nondisjunction in fission yeast (Cooper et al., 1998; Nimmo et al., 1998). Thus, the primary, important consequence of deletion of NDJ1 is likely to be the failure of bouquet formation, with the subsequent defects arising from distorted telomere distribution and the absence of bouquet-facilitated homologue alignment and pairing, as suggested based on genetic data from disomic synapsis meiosis (Rockmill and Roeder, 1998). The ndj1Δ-related delay may result from overloading of the homology testing machinery due to disordered chromosome distribution in the absence of meiotic telomere clustering. Dissolution of premeiotic/vegetative centromere and telomere aggregates at the onset of meiosis may still create sufficient chromosome movement in the Ndj1p-deficient prophase nucleus for portions of chromosomes to make contact and, once partially aligned, to commence with homosynapsis. Particularly, larger-sized chromosomes may benefit from an increased probability of encountering a portion of its homologue and to initiate stable pairing earlier than smaller ones. Synapsis, which may initiate at a few interstitial sites (Dresser and Giroux, 1988; Rockmill et al., 1995; Chua and Roeder, 1997; Trelles-Sticken et al., 1999) seems to proceed at a slower rate in the ndj1Δ mutant, since Zip1p polycomplexes resolve more gradually. Jumbled telomere localization and nonsynchronized homologue pairing may be the cause for defective crossover interference in the absence of Ndj1p (see Chua and Roeder, 1997). However, at the present state, it cannot be excluded that the ndj1Δ phenotypes result from other so far unknown effects besides the telomeric ones.

An interesting possibility is that Ndj1p may link meiotic telomeres to motor proteins at the nuclear envelope, the action of which is thought to bring about telomere aggregation during the bouquet stage (see Bascom-Slack and Dawson, 1997; Zickler and Kleckner, 1998). Future research will have to show whether perinuclear filaments, like the Mlp proteins (Strambia-de-Castillia et al., 1999), which are involved in tethering vegetative telomeres to nuclear pore complexes (Galy et al., 2000), also play a role in telomere clustering during meiotic prophase.

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