Heterochromatin of maize chromosomes: structure and genetic effects

Margarida L.R. de Aguiar-Perecin1, Antonio Fluminhan2, Janay A. dos Santos-Serejo1, José R. Gardingo1,3, Mônica R. Bertão1,4, Maria Juliana U. Decico1 and Mateus Mondin1

1Departamento de Genética, ESALQ, Universidade de São Paulo, 13400-970 Piracicaba, SP, Brasil. 2Universidade do Oeste Paulista, Rodovia Raposo Tavares, km 572, 19067-175 Presidente Prudente, SP, Brasil. 3Universidade Estadual de Ponta Grossa-PR, Praça Santos Andrade S/N, 84010-790 Ponta Grossa, PR, Brasil. 4Universidade Estadual de Santa Cruz, Rodovia Ilhéus-Itabuna, km 16, 45650-000 Ilhéus, BA, Brasil.

Send correspondence to M.L.R.A.P. E-mail: mlrapere@carpa.ciagri.usp.br

CLASSES OF MAIZE HETEROCHROMATIN

Maize (Zea mays L.) chromosomes were first characterized by McClintock (1929) from studies of the first pollen mitosis. Later, McClintock (1930, 1931, 1933) found that pachytenic chromosomes showed better morphological details for cytological studies. The pachytenic chromosomes were identified on the basis of their length, centromere position, prominent chromomeres and heterochromatic knobs (Longley, 1938; Rhoades, 1955; McClintock et al., 1981). Other heterochromatic regions found in a number of different chromosome regions have also been described: heterochromatin adjacent to the centromeres, heterochromatin of the nucleolar organizer region (NOR), B chromosomes and the abnormal chromosome 10 (Rhoades, 1978).

The examination of chromosomes of several races of maize has revealed that the number, size and position of knobs are variable and that they are found in 23 locations on the ten maize chromosomes (McClintock, 1978a; McClintock et al., 1981).

The identification of mitotic chromosomes using conventional staining was described by Chen (1969) and Filion and Walden (1973), while C-banding procedures have shown the presence of stained distal bands in mitotic chromosomes (Hadlaczky and Kálmán, 1975; Ward, 1980; Aguiar-Perecin, 1985; Rayburn et al., 1985; Jewell and Islam-Faridi, 1994). Ward (1980) and Aguiar-Perecin (1985) described differential staining of the classes of maize heterochromatin: heavily stained bands on mitotic chromosomes correspond to knobs, while staining of centric heterochromatin is hardly observed in well-condensed metaphases. NOR-heterochromatin appears differentially stained but with a lower degree of staining than knobs. No differential staining was observed in B chromosomes, but Ward (1980) observed staining of the B centric heterochromatin. Furthermore, the analysis of C-banded mitotic metaphases of maize races with different knob constitutions showed that large bands corresponding to large and medium knobs alter the arm lengths of mitotic chromosomes (Aguiar-Perecin and Vosa, 1985).

The different classes of maize heterochromatin can also be differentiated by their times of replication in the mitotic cycle, for knobs replicate later than other heterochromatic regions (Pryor et al., 1980). Also, Peacock et al. (1981) found that a repeating unit of 180 base pairs is the major component of knob heterochromatin. A certain level of polymorphism has been detected among copies of this 180-bp sequence (Dennis and Peacock, 1984), while Viotti et al. (1985) reported that this 180-bp repeat can be found at some euchromatic sites of pachytenic chromosomes. A strict correspondence between the size of the C-bands and the signal for the in situ hybridization of the 180-bp sequence has been observed along with new evidence of increase of arm length by large bands on metaphase chromosomes (Bertão, 1998).

The recovery of maize chromosome addition lines of oat (Avena sativa L.) from oat x maize crosses, each possessing an individual maize chromosome, has provided a unique opportunity to study knob DNA structure (Ananiev et al., 1998). The analysis of an oat-maize chromosome 9 addition line has revealed that blocks of tandemly arranged 180-bp repeating units are interrupted by insertions of other repeated DNA sequences, mostly represented by individual full-size copies of retrotransposable elements.

GENETIC EFFECTS OF HETEROCHROMATIN OF MAIZE CHROMOSOMES

Some genetic effects have been reported to be associated with the heterochromatin of B chromosomes, abnormal chromosome 10, knobs and chromatin adjacent to the centromeres (Rhoades, 1978).

Among the effects caused by B chromosomes is the loss of terminal chromosomal segments from knobbed A chromosomes at the second microspore mitosis in spores possessing two or more B chromosomes (Rhoades and Dempsey, 1973). B chromosome nondisjunction also occurs at the second microspore division and Rhoades and Dempsey (1973) proposed a mechanism to explain both events by postulating that delayed replication of centric B heterochromatin and knobs of A chromosomes results in B chromosome nondisjunction and formation of bridges as the two sister centromeres of A chromosomes move to opposite poles, resulting in the loss of segments of the A chromosomes.
Some interesting effects have been associated with the large heterochromatic block of abnormal chromosome 10 (K10), including preferential segregation during meiotocenes of abnormal chromosome 10 (K10), including preferential segregation during meiosporogenesis and neocentromere formation (Rhoades, 1978). Preferential segregation and neocentromere formation involve the interaction of K10 with knobs on other chromosomes at specific meiotic stages. Neocentromeres occur in homozygotes and heterozygotes for K10, but not in homozygotes for normal chromosome 10, and are found on knobbed arms of chromosomes at anaphase I and anaphase II, where the distal neocentric regions undergo a precocious movement, pulling the ends of the chromosomes to the poles. Rhoades (1978) suggested that neocentromere formation accounts for preferential segregation: at the second meiotic division of meiosporogenesis the basal spore preferentially receives a knobbed chromatid.

Studies of the variability of knob size and numbers among maize races have emphasized the importance of knowledge of knob contents in helping to understand problems associated with the origin of maize and its present-day races (McClyntock, 1978a; McClintock et al., 1981). Knobs have also been associated with differences in recombination in particular regions of the chromosome complement (Rhoades, 1978). Chughthai and Steffensen (1987) reported a positive correlation between the presence of some knobs and later flowering time while Aguiar-Perecin and Fluminhan, 1992; Fluminhan, 1992; Fluminhan et al., 1996). Mitotic instability was investigated in Feulgen-stained preparations of embryogenic calli. Bridges resulting from delayed separation of sister chromatids as well as typical bridges (Figure 1a and b), broken bridges and fragments were observed. The examination of C-banded anaphases showed that sister chromatids were held together at knob sites (primary event), an observation which gives support to the hypothesis previously proposed by Lee and Phillips (1987) to explain the high frequency of knobs in chromosome arms involved in rearrangements. Also, typical bridges with and without bands corresponding to knobs were observed. These events were interpreted as evidence of the occurrence of breakage-fusion-bridge (BFB) cycles initiated by chromosome arms broken during the primary event. These cycles would be similar to the chromatid type of BFB cycle described by McClintock (1939). Additional

![Figure 1 - Feulgen-stained anaphases of a callus culture](image-url)

**Figure 1** - Feulgen-stained anaphases of a callus culture. (a) Bridge resulting from delayed separation of sister chromatids; (b) two typical bridges, probably involved in a breakage-fusion-bridge cycle. Bar = 10 μm.
Heterochromatin of maize chromosomes

evidence for this mechanism was the presence of gross aberrations involving chromosome 7, interpreted as duplication-deficiencies.

Further studies involving a detailed analysis of C-banded metaphases of 2-3-year-old cultures (Santos, 1995) and 2-4-month-old cultures (Gardingo, 1998) have provided new evidence for the occurrence of BFB cycles and have shown that chromosome 7 is the most affected. The genotypes used were lines highly related to the ones studied by Fluminhan et al. (1996). Figure 2a shows a C-banded metaphase cell with normal chromosome 7 possessing a knob on the short arm (K7S) and the long arm (K7L). Figure 2b and c shows one of the alterations observed, that is, an amplification of K7L and if one assumes that this alteration was derived from a BFB cycle, then it must be concluded that this chromosome has a deficiency of the distal euchromatic segment (from the knob to chromosome end) on the long arm.

These observations have raised interesting questions regarding the effect of culture age on the frequency of anaphase bridges and the occurrence of chromosome healing, similar to that described by McClintock (1941, 1942, 1978b). The analysis of C-banded metaphases in 2-3-year old calli (Santos, 1995; Santos and Aguiar-Perecin, 1995) gave strong evidence for the occurrence of healing of broken chromosome ends of chromosomes 7 and 9, after BFB cycles: certain chromosome rearrangements were observed unaltered in several samples of most cultures.

Recently, Santos-Serejo, Mondin and Aguiar-Perecin found plants homozygous for an amplification at K7L, in R1 progenies derived from regenerated plants heterozygous for this aberration. In this specific case, we speculate that this amplification of K7L was originated through a mechanism other than the BFB cycle. Gardingo (1998) detected the occurrence of unequal sister chromatid exchange in chromosome 7, resulting in amplification and deficiency at K7L. This event was observed in some cultures and must be another mechanism of chromosome alteration in vitro. In this case, chromosome 7 would not be deficient for the distal segment of the long arm.

Among other aspects investigated in our laboratory is that of the correlation between knob content and the frequency of anaphase bridges in callus cultures. If delayed chromatid separation is the primary event causing bridge formation, then we could expect a lower frequency of bridges in genotypes with low knob content. This was investigated using 5-6-month-old cultures derived from four families of related inbred lines differing in their knob contents (Fluminhan and Aguiar-Perecin, 1998). The lines used were homozygous for knobs at 6L, 6L+, 7L and 8L, and differed for the presence or absence of K2L, K3L, K7S and K9S (references in Aguiar-Perecin and Decico, 1998;
Deciço, M.J.U. (1991). Differences in embryogenic response were observed between these families of inbreds. Only one family formed highly embryogenic type II calli. Mitotic instability was investigated in most cultures and delayed separating chromatids, typical bridges and fragments were observed. The frequency of both types of bridges was not strictly correlated with the knob content of the genotypes analyzed, suggesting that knobs may undergo alterations leading to mitotic disturbance and that this response may be genotype dependent.

This research project represents a contribution to the study of stress caused by the tissue culture environment on chromosome behavior and response dependent on genotype and heterochromatin content, as well as provides information for the selection of genotypes yielding a high frequency of embryogenic cultures with a low level of chromosome instability (Gardingo, 1998; Santos-Serejo, 1999).

ACKNOWLEDGMENTS

Publication supported by FAPESP.

REFERENCES

Rhoades, M.M. (1978). Genetic effects of heterochromatin in maize. In:


