

TECHNICAL ARTICLE

Homoeologous chromosome pairing between the A and B genomes of *Musa* spp. revealed by genomic in situ hybridization

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- Background and Aims Most cooking banana and several desert bananas are interspecific triploid hybrids between Musa acuminata (A genome) and Musa balbisiana (B genome). In addition, M. balbisiana has agronomical characteristics such as resistance to biotic and abiotic stresses that could be useful to improve monospecific acuminata cultivars. To develop efficient breeding strategies for improving Musa cultivars, it is therefore important to understand the possibility of chromosome exchange between these two species.
- Methods A protocol was developed to prepare chromosome at meiosis metaphase I suitable for genomic in situ hybridization. A series of technical challenges were encountered, the main ones being the hardness of the cell wall and the density of the microsporocyte's cytoplasm, which hampers accessibility of the probes to the chromosomes. Key parameters in solving these problems were addition of macerozyme in the enzyme mix, the duration of digestion and temperature during the spreading phase.
- Results and Conclusions This method was applied to analyse chromosome pairing in metaphase from triploid interspecific cultivars, and it was clearly demonstrated that interspecific recombinations between M. acuminata and M. balbisiana chromosomes do occur and may be frequent in triploid hybrids. These results provide new insight into Musa cultivar evolution and have important implications for breeding.

Key words: Musa, banana, genomic in situ hybridization, meiosis, homoeologous chromosomes pairing, interspecific hybrid, polyploidy.

INTRODUCTION

Cultivated bananas (bananas and plantains) are seedless parthenocarpic clones, selected by early farmers in South-East Asia and maintained by vegetative propagation (Simmonds, 1962). They represent the fourth most important crop in developing countries (Lassois et al., 2009). They are important as an export crop, providing a vital source of income for many countries and also playing a major role in local food security (Chalmin, 2009). Cultivars are derived from natural intra- and interspecific hybridization between wild fertile diploid Musa accessions. Most cultivars are triploid, some are diploid and a few are tetraploid. The main species involved are Musa acuminata Colla (A genome, 2n = 2x = 22) and Musa balbisiana Colla (B genome, 2n =2x = 22) (Cheesman, 1947; Simmonds and Shepherd, 1955). Cultivars were classified based on morphological characters and chromosome number into genome groups AA, AB, AAA, AAB or ABB (Cheesman, 1947; Simmonds and Shepherd, 1955). More recently, molecular markers confirmed and refined this classification (Ude et al., 2002; Carreel et al., 2002; Creste et al., 2004; Perrier et al., 2009; Hřibová et al., 2011). The small size of Musa genomes, i.e. 500-600 Mbp (1C) (Dolezel et al., 1994; Lysak et al., 1999; Bartos et al., 2005) and thus of *Musa* chromosomes $(1.5-3.5 \mu m)$ makes

conventional karyotypic analysis difficult (Isobe and Hashimoto, 1994; Osuji et al., 1996a, b). However, many conventional cytogenetic studies have been performed in Musa, in particular to study chromosome pairing during meiosis (Wilson, 1945; Simmonds, 1962; Dessauw, 1987; Fauré et al., 1993; Shepherd, 1999; Therdsak et al., 2010). They suggested that M. accuminata subspecies differed in their chromosome structure due to rearrangements, particularly translocations (Dodds, 1943; Fauré et al., 1993; Shepherd, 1999). These variations in chromosome structures disrupt meiosis in the hybrids and are believed to contribute to their sterility. This character as well as parthenocarpy was selected by early farmers for the production of edible fruits.

Different production areas currently suffer from new emerging diseases and have to face an ever-increasing range of pests and diseases (Pennisi, 2010). In the absence of locally adapted resistant varieties, the crop requires extensive use of pesticides, which threatens the sustainability of the crop and environment. Despite its economic importance, the banana export industry has been affected, as it relies on monoculture of genetically extremely closely related clones of the Cavendish subgroup (sterile triploids, AAA). There is thus an urgent need for a wider diversity of genetically improved banana cultivars with more robust disease resistance, increased productivity and better adaptability to a wide large range of growing conditions. However, breeding

programmes are facing serious difficulties, mainly due to the low level of fertility, structural heterozygosity and triploidy of cultivated banana and to the absence of knowledge on the genetic factors involved in important agricultural traits. *Musa balbisiana* has important potential for breeding, such as conferring good ratooning ability, a strong root system and more generally resistance to biotic and abiotic stresses (Bakry *et al.*, 2009). However, there is a serious lack of information regarding chromosome segregation in interspecific hybrids between *M. accuminata* and *M. balbisiana* and the possibility of chromosome exchange between these two genomes. This is due to the fact that chromosomes from these two species cannot be differentiated based on their size and/or morphology.

Genomic in situ hybridization (GISH) is a powerful tool to differentiate chromosomes from parental species in interspecific hybrids. It was developed by Schwarzacher et al. (1989) since when it has been applied to mitotic chromosomes from many plants resulting from interspecific hybridization (Jiang and Gill, 1994; D'Hont et al., 1996; Lashermes et al., 1999; Tanguy et al., 2005; Konnan et al., 2009; Mason et al., 2010). Likewise, GISH was successfully applied to differentiate the A and B chromosome of Musa on mitotic chromosome spreads prepared from root tips (Osuji et al., 1997; D'Hont et al., 2000). However, application of GISH on meiotic chromosomes is challenging and has been reported in just a few species, such as Solanum (Zhong et al., 1996), Hordeum (Anamthawat-Jonsson et al., 1993), Medicago tranculata (Kulikova et al., 2001), Beta vulgaris (Desel et al., 2001), Arabidopsis thaliana (Armstrong et al., 1998), Alstroemeria hybrids (Kamstra et al., 2004), Brassica (Armstrong et al., 1998; Kun et al., 2006; Nicolas et al., 2007), Lilium (Zhou et al., 2008), Lolium (Kopecky et al., 2008) and Brachiaria (Souza-Kaneshima et al., 2010). In Musa, a first report of fluorescence in situ hybridization (FISH) on meiotic chromosomes at pachytene stage was published recently (De Capdeville et al., 2009) but no report exists on GISH on meiotic chromosomes.

Here we describe a protocol developed to perform GISH on meiosis preparations at metaphase I from pollen mother cells and detail the technical challenges faced. We applied this protocol to study chromosome pairing in two interspecific triploid cultivated clones. The results demonstrated for the first time that pairing between chromosomes A and B occurs and may be frequent. Finally, we discuss the potential of these findings on *Musa* cultivar evolution and for banana breeding.

MATERIALS AND METHODS

Plant material

Two interspecific triploid banana clones involving *Musa acuminata* and *M. balbisiana* and belonging to two different banana groups (AAB and ABB) were analysed: 'Figue Pomme' (AAB, 2n = 33) of the 'Silk' subgroup and 'Praha' (ABB, 2n = 33) of the 'Pisang Awak' subgroup. These two clones were provided from the African Centre for Research on Banana and Plantain in Cameroon.

Meiotic chromosome preparation

Young anthers containing meiotic chromosomes were selected according to Fauré et al. (1993). The stage of

development was determined via a acetocarmine squash preparation using a single anther from a flower. If at metaphase I. the remaining anthers were fixed directly in ethanol/acetic acid (3:1) and could be stored in 70% ethanol, 4°C for a few months. Flowers were harvested from 45-d-old inflorescences and the appropriate stages were found within buds in flower clusters (= 'hand') number 30 for 'Figue Pomme' and number 35 for 'Praha' (flower size 0.5-1 cm). Flower cluster number 0 corresponds to the latest mature flowers that are located under the latest opened bract. Fixed anthers were rinsed twice in deionized water, then in citrate buffer (10 mm, pH 4.5) and then incubated for 6 h at 37 °C in a mixture of pectolytic enzymes containing 0.3 % (w/v) cytohelicase (Duchefa, www.duchefa.com), 0.3 % cellulase 'Onozuka' RS (Duchefa), 0.3 % pectolyase Y-23 (Duchefa) completed with variable concentrations of macerozyme R-10 (Duchefa) depending on cultivars in 10 mm citrate buffer, pH 4.5. For 'Figue Pomme' and 'Praha' accessions, best results were obtained with 6 and 7 % macerozyme, respectively. After washes in deionized water, the pollen mother cells were dissected out of the anthers into a watch glass using finemounted needles, taking care to remove as much as possible of the supporting tissues.

A 3 μL droplet of the cell suspension was then carefully transferred into grease-free slides, 15–30 μL of 60% acetic acid was added and the pollen mother cells were left for 3 min on a hot plate at 50 °C for 'Figue Pomme' or 65 °C for 'Praha'. A ring of freshly prepared ice-cold fixative (3:1) was added around the droplet containing the meiotic cells. Shortly after the fixative had mixed with the cell suspension, the cells were spread on the slide.

The quality of the slides was controlled by microscopic observation under phase contrast optics. Slides were used directly for *in situ* hybridization or were stored at $-20\,^{\circ}\text{C}$ until needed.

Genomic in-situ hybridization

Before hybridization, the slides were fixed for 1 h in an oven at 65 °C, treated with RNase (Sigma, www.sigmaaldrich.com; 1 $\mu g \text{ mL}^{-1}$) in a moist chamber at 37 °C for 45 min and washed twice with 2× sodium saline citrate (SSC). The hybridization mixture (30 µL per slide) consisted of 50 % formamide, 10 % dextran sulphate, $2 \times$ SSC, 0.66 % sodium dodecyl sulphate and 6 ng μL^{-1} of each parental total genomic DNA probe. Genomic DNA of 'Pisang Klutuk (M. balbisiana, B genome) and 'Pahang' (M. acuminata, A genome) were labelled by random priming biotin-14-dUTP (Invitrogen Life Technology, www.invitrogen. com) and digoxigenin-11-dUTP (High Prime DNA Labeling Kit, Roche, www.roche.com), respectively. The hybridization mixture was denatured for 10 min in boiling water and dispensed on the slides. The chromosomes (with the hybridization mixture) were then denatured in a moist chamber placed on a water bath equilibrated at 80 °C for 2 min. Hybridization were performed overnight in a moist chamber at 37 °C. After hybridization, slides were successively washed for 10 min in $2 \times$, $0.5 \times$ and $0.1 \times$ SSC at 42 °C. Biotinylated probes were immunodetected with Texas-Red-conjugated avidin antibodies (ABCYS, www.abcysonline.com). Digoxigenin-labelled probes were detected with digoxigenin antibody conjugated with

fluorescein isothiocyanate (FITC) (ABCYS). The chromosomes were mounted in Vectashield antifade solution (Vector Laboratories, www.vectorlabs.com) containing 2.5 μg mL $^{-1}$ 4′,6-diamidino-2-phenylindole (DAPI) as counterstaining. Fluorescence images were captured separately using a cooled high-resolution black and white CCD camera (Orca Hamamatsu, www.hamamatsucameras.com) and a Leica (www.leica-microsystems.com) DMRAX2 fluorescence microscope. The camera was interfaced to a PC running Volocity software (Perkin Elmer, www.perkinelmer.com).

RESULTS AND DISCUSSION

Development of GISH on Musa meiotic chromosomes from pollen mother cells

GISH on mitotic chromosomes obtained from root tips has been reported on many species, including *Musa* (Osuji *et al.*, 1997; D'Hont *et al.*, 2000), with quite similar protocols. By contrast, GISH on meiotic chromosomes in plants is quite challenging and the protocols used are complex and highly variable depending on the species. Accordingly, we encountered a series of technical challenges in *Musa* for preparing meiotic chromosome spreads suitable for GISH. The main challenge was the hardness of the cell wall and the density of the microsporocyte's cytoplasm, which hamper the accessibility of the probes to the chromosomes and generate higher levels of background, as already reported for *Musa* by De Capdeville *et al.* (2009).

We compared and tested the digestion protocols used in several species which vary in particular for the composition and concentration of the following enzyme cocktail: cellulase, pectinase, cytohelicase and pectolyase (Armstrong *et al.*, 1998; Desel *et al.*, 2001; Kulikova *et al.*, 2001; Kun *et al.*, 2006; De Capdeville *et al.*, 2009). However, in our hands and with our plant material they did not provide acceptable results. We performed staining experiments on cells with calcofluor, propidium iodide and ruthenium red, which revealed that chromosomes were still embedded in an outer crust of pectin. Finally, we managed to digest those pectins by adding macerozyme to 0·3 % cytohelicase, pectolyase and cellulase. Macerozyme is used for protoplast production in various plants such as banana (Assani *et al.*, 2001).

The addition of macerozyme in the enzyme mix, the duration of digestion and the temperature of the hot plate during the spreading phase were the key parameters in the success of our protocol. Note that these three parameters have to be adapted slightly for each clone and stage of meiosis. With this protocol, we were able to obtain meiotic chromosome spreads suitable for FISH for various accessions of AA, AB and AABB genomic constitution in addition to those reported in the present paper and from different stages of meiosis (prophase I, metaphase I and anaphase I) (data not shown). This protocol requires fresh material (fixed for no more than a few months). For both triploid clones, 'Figue Pomme' and 'Praha', the development of microsporocytes was not synchronized. This phenomena was more important in 'Figue Pomme', for which we observed on the same slide (from a single anther) early meiotic stages still in prophase I together with microsporocytes that had already reached tetrad stage. Clones with highly asynchronous

microsporocyte development required much more effort to get a reasonable rate of suitable chromosome spreads.

We then optimized our classical GISH protocol (D'Hont *et al.*, 2000) by including the following changes: probes were labelled by random priming, slides with chromosome preparation were fixed for 1 h at 65 °C, chromosome denaturation was performed on a floating moist chamber over a water bath (Leflon *et al.*, 2006) and the washes were softened.

Homoeologous chromosome pairing between the A and B genome

This protocol was used to perform GISH on meiotic metaphases of interspecific hybrids involving M. acuminata and M. balbisiana with genomic DNA of these species. The differential labelling obtained for A versus B chromosomes was excellent, so it was possible for the first time to attribute an A or B origin to the chromosomes involved in various pairing configuration (Figs 1 and 2). Labelling was essentially located on the centromeric and pericentromeric part of the chromosome leaving most of the chromosome arms unlabelled. This partial labelling was observed by D'Hont et al. (2000) when analysing Musa mitotic chromosomes but is much clearer on meiotic chromosomes. This uneven labelling observed on species with small chromosomes (Barre et al., 1998; Lashermes et al., 1999; Ollitrault et al., 2000; Ali Hoda et al., 2004) most probably results from the lower repeated sequence content in these genomes (D'Hont, 2005) and to the fact that the repeated sequences are typically more abundant in centromeric and pericentromeric parts of the chromosomes.

It is also interesting to note that the labelling obtained with the B genomic DNA was relatively homogeneous on all centromeric regions of all chromosomes (Fig. 1B) in contrast to the labelling obtained with the A genomic DNA, which was variable depending on chromosomes (Fig. 1C). This may be related to the fact that B genomes are slightly smaller than A genomes (Dolezel *et al.*, 1994; Lysak *et al.*, 1999; Bartos *et al.*, 2005) and thus species-specific repeated sequences appeared more numerous in *M. acuminata* than in *M. balbisiana* (Baurens, 1997).

Two triploid interspecific accessions were analysed involving a different balance of A and B genomes: accession 'Figue Pomme' with an AAB genomic constitution (Fig. 1) and accession 'Praha' with an ABB genomic constitution (Fig. 2). Classical cytogeneticists have reported that it is particularly difficult to obtain good spreading at meiosis metaphase for interspecific triploid *Musa* clones (Wilson, 1945; Simmonds, 1962). Accordingly, for most of the cell only part of the pairing configuration could be analysed, the other chromosomes being not sufficiently spread. We observed that chromosome associations were quite variable between the different metaphases of the same clone. This has been noted in interspecific hybrids from *Musa* (reviewed by Shepherd, 1999) and in other genera such as *Lilium* (Zhou *et al.*, 2008).

For both genotypes, we identified univalents, bivalents and multivalents (Figs 1 and 2) in accordance with previous results based on acetocarmin staining (Shepherd, 1999). In addition, for the first time, we were able to visualize homoeologous pairing. Homoeologous bivalents were observed in all analysed cells with two to five homoeologous bivalents for 'Figue

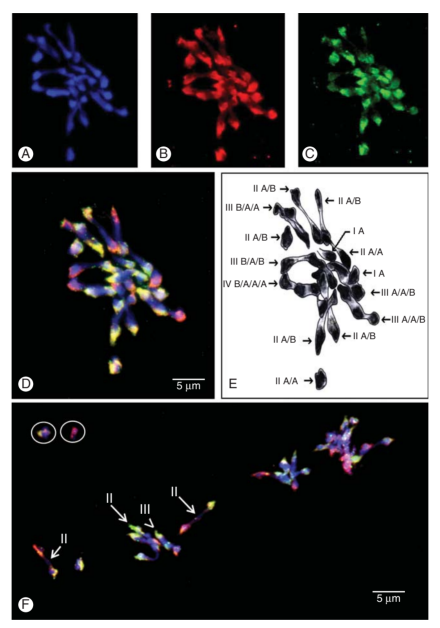


Fig. 1. GISH on meiotic metaphase I of interspecific triploid *Musa* cultivar 'Figue pomme' (genome AAB). (A) DAPI, (B) B genomic DNA revealed in red (Texas Red), (C) A genomic DNA revealed in green (FITC) and (D, F) after superimposition of the three colours. (E) Schematic interpretation of (D). Circles indicate chromosomes that were outside of the presented field. Arrows indicate homoeologous bivalents and arrowsheads indicate homoeologous multivalents.

Pomme' (Fig. 1) and one to four for 'Praha' (Fig. 2). Moreover, all multivalents (trivalents and tetravalents) observed involved homoeologous chromosomes (Figs 1 and 2).

Information regarding the possibility of homoeologous chromosome pairing have tentatively been inferred from classical chromosome pairing studies. In diploid AB interspecific hybrids, bivalents were observed in variable proportions from three to 11, suggesting partial homoeologous chromosome pairing (Dodds and Simmonds, 1946; Shepherd, 1999). In polyploid interspecific hybrids (AAB, ABB), interpretations were also based on the numbers of bivalents and multivalents observed. However, as multivalents are observed in monospecific clones (AA and AAA) (reviewed by Shepherd 1999),

these studies were largely speculative and sometimes contradictory; for example, Shepherd (1999) concluded that there was low affinity between the chromosomes of the two species whereas Dessauw (1987) concluded the opposite. GISH on mitotic chromosomes was unable to resolve this issue as the partial labelling obtained prevented the detection of interspecific recombinant chromosomes (Osuji *et al.*, 1997; D'Hont *et al.*, 2000).

Here for the first time, we clearly demonstrated that interspecific recombination between A and B chromosomes does exist and may be frequent. This information has important implications for the domestication and improvement of interspecific cultivars through breeding.

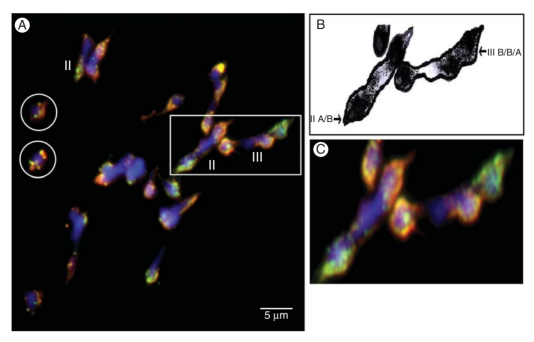


Fig. 2. (A) GISH on meiotic metaphase I of interspecific triploid *Musa* cultivar 'Praha' (genome ABB) after superimposition of DAPI, Texas Red and FITC (cf. Fig. 1). (B) Schematic interpretation of part of (A). (C) Focus on part of (A). Circle indicates chromosomes that were outside of the presented field. Arrows indicates homoeologous bivalents and arrowsheads indicate homoeologous multivalents.

Implications for Musa cultivar evolution and for banana breeding

Interspecific triploid cultivars derived from M. acuminata and M. balbisiana are believed to result from one or two steps of combination between the parental species, featuring 2n gametes derived from AA and AB genotypes (Cheesman, 1947; Simmonds and Shepherd, 1955; Perrier et al., 2009). De Langhe et al. (2010) emphasized that additional steps of combination may have occurred. The possibility of A/B pairing revealed in our study implies that chromosome re-assortments and exchanges of chromosome segments may have occurred between the two genomes, leading to unbalanced genome transmission with respect to the parental species. The 8A + 25B chromosome constitution of the triploid interspecific cultivar 'Pelipita' revealed by GISH (D'Hont et al., 2000) constitutes the first accurately documented example of important unbalanced genome constitution; note that this reveals essentially the origin of the centromeric regions and may overlook chromosome arm recombinants. Further work to understand the impact of interspecific chromosome pairing on cultivar chromosome constitution will require the analysis of interspecific cultivars and/or progenies with numerous A- and B-specific molecular marker alleles. The ongoing Musa genome sequencing project will further facilitate development of markers for future high-throughput characterizations (http://www.genoscope.cns.fr/spip/September-8th-2009-Banana-genome.html).

Production of edible banana requires sterile clones or clones with only residual fertility; consequently, developing new cultivars requires complex breeding strategies. The main strategy used so far to create new interspecific cultivars is based on exploitation of residual fertility of triploid cultivars in combination with fertile diploid accessions (Menendez and

Shepherd, 1975; Bakry and Horry, 1992; Rowe and Rosales, 1993; Vuylsteke et al., 1993). The tetraploid progeny resulting from un-reduced 2n gametes from the triploid parent and ngametes from the diploid parent are then selected. We are currently testing in Guadeloupe (French West Indies) another strategy which consists in doubling AB hybrids and crossing them with AA or BB diploids (Bakry et al., 2009). The occurrence of interspecific chromosome pairing demonstrated in the present study opens new perspectives for this latter strategy as it implies that a much wider range of gametic genotypes can be obtained from an AABB genotype compared with a pure disomic inheritance. In addition, the possibility of introgressing only fragments of B chromosomes opens new perspectives to exploit the B genomes for improving resistance and rusticity of monospecific acuminata cultivars. This should allow us to introgress only B chromosome segments bearing a target character and avoid introgression of B chromosome fragments harboring endogenous banana streak virus (eBSV) sequences. eBSV sequences are able to release infectious virions and such activations are known to be favoured in an intergenomic context (Lheureux et al., 2003; Gayral et al., 2008). Recent results suggested that eBSV sequences are present at only a few loci in the B genome (Iskra-Caruana et al., 2010).

Finally, homoeologous pairing and thus interspecific chromosome exchanges will most probably vary depending on genotypes, through genic control (Griffiths *et al.*, 2006; Kopecky *et al.*, 2008; Nicolas *et al.*, 2009), and environment. The possibility of testing the extent of homoeologous pairing, demonstrated in this paper using GISH, in various types of parental materials and crosses should help to define new efficient breeding strategies.

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