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A change of developmental program induces the remodeling of the interchromatin domain during microspore embryogenesis in *Brassica napus* L.

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Abstract

After a stress treatment, the in vitro-cultured pollen changes its normal gametophytic developmental pathway towards embryogenesis producing multicellular embryos from which, finally, haploid and double haploid plants develop. The architecture of the well-organized nuclear functional domains changes in response to DNA replication, RNA transcription, processing and transport dynamics. A number of subnuclear structures present in the interchromatin region (IR, the nuclear domain between chromosome territories) have been shown as involved, either directly or indirectly, in transcriptional regulation. These structures include the interchromatin granule clusters (IGCs), perichromatin fibrils (PFs), Cajal bodies (CBs) and perichromatin granules (PGs). In this work, we present a cytochemical, immunocytochemical, quantitative and morphometric analysis at the light, confocal and electron microscopy levels to characterize the changes in the functional architecture of the nuclear interchromatin domain during two developmental programs followed by the microspore: differentiation to mature pollen grains (transcriptionally inactive), and microspore embryogenesis involving proliferation in the first stages (highly engaged in transcription). Our results revealed characteristic changes in size, shape and distribution of the different interchromatin structures as a consequence of the reprogramming of the microspore, allowing us to relate the remodeling of the interchromatin domain to the variations in transcriptional activities during proliferation and differentiation events and suggesting that RNA associated structures could be a regulatory mechanism in the process. In addition, we document the presence of two structurally different types of CBs, and of IGC and CB-associated regions, similar to those present in animal cells, and not yet described in plants.
Introduction

The microspore or immature pollen grain normally follows the gametophytic program and differentiates to form the mature pollen, a process that can also be reproduced in vitro. Upon the application of a stress treatment it can be deviated towards a proliferation process leading to embryogenesis and plant regeneration, the so-called microspore embryogenesis (Seguí-Simarro and Nuez 2008), a process which represents an important tool in plant breeding to obtain double-haploid plants. The entry into proliferation of differentiating cells is a key event in plant development and in organogenic and morphogenetic processes. Since the cell nucleus governs the cell activity, the know-how of the dynamics of the nuclear domains during the activation of such processes will give new insights into the mechanisms that regulate them. Pollen developmental programs represent model systems to analyze the cellular changes which accompany the change of developmental program and the switch to proliferation from plant differentiating cells.

The nucleus is the cell factory where DNA replication and RNA transcription and processing take place. It is a highly dynamic compartment where different activities are confined to specific domains whose structure is markedly dependent on their function (Spector, 1993a). Nuclear chromatin is divided in chromosome territories, which encompass (1) condensed chromatin (heterochromatin, transcriptionally inactive) and (2) decondensed chromatin (euchromatin) where transcriptionally active genes are located (Cremer and Cremer, 2001). These different levels of condensation, together with other remodeling and epigenetic events, are considered as a first, pretranscriptional level of regulation of gene expression (Jarillo et al., 2009). Indeed, rearrangements in the position
and condensation of chromosomes are known to play an important role in the establishment
of the different developmental expression patterns by exposing and allowing access of the
transcriptional machinery to specific gene sequences. From a molecular point of view, the
different regulatory steps of transcription and pre-mRNA processing in eukaryotes are well
known processes. There is a wealth of data on how they proceed and the different
molecular players involved. Indeed, in vitro transcription is possible for long, which has
boosted the emergence of a pleiade of molecular biology techniques. However, there is less
information on where transcription-related processes take place in vivo, and how the
different structures involved change their architecture in response to cell requirements
during different developmental programs.

The nuclear space separating chromosome territories is the interchromatin region
(IR). At the boundaries of the IR with the condensed chromatin, decondensed chromatin
fibers are being transcribed to RNA in the form of perichromatin fibrils (PFs). PFs are short
fibers of 3-5 nm in diameter, present in a dispersed pattern through the perichromatin
region. They are thought to be the visible evidence of nascent pre-mRNA coupled to the co-
transcriptional splicing machinery (Monneron and Bernhard, 1969; Testillano et al., 1993;
Cmarko et al., 1999; Fakan, 2004). In addition, the IR houses the different structures
involved on post-translational regulatory steps such as RNA processing and nuclear protein
modification (Cremer and Cremer, 2001).

Once the pre-mRNA (hnRNA) sequence is generated in the form of a PF, three
subdomains are proposed as directly or indirectly involved in pre-mRNA splicing,
processing and transport: interchromatin granules, Cajal bodies and perichromatin
granules (Spector, 1993a; Raska, 1995; Lamond and Spector, 2003; Fakan, 2004).

Interchromatin granule clusters (IGCs) were found in mammalian (Puvion et al., 1984) and plant nuclei (Testillano et al., 1993), and they are seen as ‘speckles’ under the fluorescence microscope. They are interchromatin structures composed of 20-25 nm particles embedded in a thin fibrillar matrix. Although different pre-mRNA splicing factors, including snRNPs and SR proteins are present in IGCs, no DNA or transcriptional activity has consistently been detected in these structures (Thiry, 1995; Cmarko et al., 1999). Thus, ICGs are currently considered as storage, assembly and modification compartments for the delivery of transcription factors to active transcription sites (Lamond and Spector, 2003). Cajal bodies (CBs) are universal, multifunctional, nearly round, convoluted and highly dynamic structures frequently found associated to the nucleolus. snRNPs, the p80 coilin protein, the U2AF splicing factor and diverse snRNAs have been detected in CBs, along with several nucleolar proteins such as fibrillarin (reviewed in Gall, 2000; Lafarga et al., 2009). However, poly-A RNA, rRNA, nascent mRNA transcripts, the SC-35 protein, DNA or RNA polymerase II are not found in CBs, indicating that they are not directly involved in transcription. Instead, it is believed that CBs are involved in the assembly and modification of the machinery for processing of pre-mRNA, pre-rRNA and histone pre-mRNA (Gall, 2000; Matera, 2003; Shaw and Brown, 2004). Different studies have related the abundance of CBs to cell activity (Andrade et al., 1993), type (Raska et al., 1991) and developmental stage (Boudonck et al., 1998). Although smaller and less conspicuous, the perichromatin granules (PGs) are also integral components of the eukaryotic nucleus. RNA processing, altered transport, storage and/or degradation of specific RNAs is thought to be mediated by PGs (Puvion and Lange, 1980; Vazquez-Nin et al., 1997).
As evidenced in most of the literature above cited, the knowledge on *in vivo* functional organization of the different subnuclear structures involved in transcriptional regulation has traditionally been derived, to a great extent, from studies on animal nuclei. Comparatively, the plant nucleus has been less studied. In order to contribute to the knowledge on the relationship between plant nuclear remodeling and expression patterns, in this work we analyzed the changes in nuclear architecture during two different developmental programs, involving proliferation and differentiation. We studied *Brassica napus* embryogenesis-induced microspores and maturing pollen grains. In microspores, we induced an embryogenic developmental program defined by an initial, proliferative phase with increased transcriptional activity (reviewed in (Seguí-Simarro and Nuez, 2008). In parallel, we studied pollen differentiation, characterized by the formation of a moderately active vegetative cell, and a generative cell transcriptionally silent (Bednarska, 1984).

The comparative study of microspores and pollen grains committed to proliferative and differentiation events constitutes a good model to study cellular changes related to a developmental reprogramming (Seguí-Simarro et al., 2003, 2005; Barany et al., 2010). In this study, we focused on the changes pertaining to the nuclear structures of the interchromatin domain (perichromatin fibrils, interchromatin granules, Cajal bodies and perichromatin granules). Our results indicated that a remodeling of the interchromatin structures accompanied the change of developmental program of the microspores, being those changes related to proliferation and differentiation events. The results also reveal two novel structural domains, not previously described in plants, and point to the notion that the different availability and abundance of these structures may serve as a regulatory mechanism operating at a post-transcriptional level.
Material and methods

Plant material

Brassica napus L. cv Topas donor plants were grown as previously described (Seguí-Simarro et al., 2003). Microspore cultures, embryogenesis induction, and sampling at specific culture stages for light and electron microscopy, immunofluorescence, and immuno-gold labelling were performed essentially according to (Seguí-Simarro et al., 2003), with some modifications to promote the suspensor-bearing embryogenic route as described (Supena et al., 2008). For the study of pollen development, anthers at different developmental stages were selected and excised from plants.

Electron microscopy and ultrastructural cytochemistry

Samples to be observed for electron microscopy were fixed in Karnovsky fixative (4% formaldehyde + 5% glutaraldehyde in 0.025M cacodilate buffer, pH 6.7), post-fixed in 2% OsO₄, dehydrated in a methanol series for 3 days and slowly embedded in Epon resin for 2 days. Epon blocks were polymerized at 60ºC for 2 days. ~80 nm-thick sections were collected on 75-mesh copper grids, counterstained with uranyl acetate and lead citrate and observed in a JEOL 1010 TEM operating at 80 kV. For the study of the different structures present in the interchromatin region, the methylation-acetylation (MA) and the EDTA cytochemical methods were used, either separately or combined. For the preferential staining of DNA, the NAMA-Ur (Testillano et al. 1991) ultrastructural cytochemistry was used. The MA method is a preembedding technique that highlights nucleic acid-containing structures (fibrillar and granular) by a selective blockage of the protein ability to react with uranyl acetate (Testillano et al., 1995). For this method, fixed samples were dehydrated in a
methanol series and incubated in a freshly made methanol:acetic anhydride mixture (5:1 v:v) overnight at room temperature, prior to embedding. After three washings with methanol, samples were embedded in Epon as described. For NAMA-Ur cytochemical method, fixed samples were subjected to a mild alkaline hydrolisis with NaOH to eliminate RNA, followed by MA treatment and Epon embedding; subsequent uranyl staining of the sections result in a preferential staining of DNA containing structures (Testillano et al. 1991). The EDTA regressive staining (Bernhard, 1969) is used over EM sections to provide RNP-containing structures a preferential staining over the bleached masses of condensed chromatin. Samples fixed in 4% formaldehyde, dehydrated by the PLT (Progressive Lowering of Temperature) method in a Leica AFS automatic system, and embedded in Lowicryl K4M resin were used for this cytochemical method.

Cryoprocessing of samples for cryomicrotomy and immunofluorescence

Microspore and microspore-derived embryo cultures were cryoprocessed for cryomicrotomy and cryosectioned as previously described (Seguí-Simarro et al., 2003). Microspore cultures were slightly prefixed in 4% formaldehyde at 4°C and embedded in gelatin. Small (1 mm³) pieces of gelatin were cryoprotected with increasing concentrations of sucrose: 0.1M (1 h), 1M (3 h), and 2.3M (overnight), placed on an aluminum pin and cryofixed in liquid propane using a KF80 unit (Reichert, Vienna) at -170°C. Thin (1 µm thick) cryosections were obtained using an Ultracut E (Reichert) ultramicrotome coupled with a FC4 unit stabilized at -75°C. Cryosections were collected and placed on multiwell glass slides for immunofluorescence. Immunofluorescence was also performed essentially as described (Seguí-Simarro et al., 2003), with the primary mouse monoclonal antibody anti-2,2,7-tri-methyl-guanosine, TMG (Calbiochem, clone K121) diluted 1/100 and
incubated for 1 h at room temperature, and the secondary antibody (anti-mouse IgG-Cy3) diluted 1:25 in 1% BSA in PBS, and incubated for 45 min. Sections were additionally stained with DAPI prior to observation. Controls were performed excluding the primary antibody.

**Immunoelectron microscopy**

Microspores and haploid embryos at different stages were prefixed in formaldehyde, cryoprotected in 2.3M sucrose, cryofixed in liquid nitrogen, and cryoprocessed as described (Seguí-Simarro et al., 2003). Samples were freeze-substituted in methanol + 0.5% uranyl acetate at -80°C for 3 days, infiltrated in Lowicryl K4M, and polymerized at -30°C under UV light in a Leica AFS system. Ultrathin (80 nm) sections were placed on nickel grids, blocked with 5% BSA in PBS, and incubated with anti 2,2,7-TMG diluted 1/100 for 1 h at room temperature. Then, the grids were incubated with a secondary antibody (anti-mouse IgG-gold, 10 nm; Biocell) diluted 1:25 in 1% BSA, for 45 min at room temperature, washed, air dried, counterstained, and observed in a JEOL 1010 EM at 80 kV. Controls were performed excluding the primary antibody.

**Morphometric and quantitative analysis**

Sampling was carried out over selected samples on each grid. The number of micrographs to be taken was determined by the progressive mean test (Williams, 1977), with a maximum confidence limit of a=0.05. Digital images of cells showing one or more CBs were processed with the ImageJ software (http://rsbweb.nih.gov/ij/) for the morphometric analysis of numbers, areas, and distances. Data were exported to a spreadsheet where the final calculations and chart displays were performed.
Results

The first step of the study was the selection of the most representative stages of microspore embryogenesis and pollen development to be processed and analyzed. For microspore-derived embryogenesis, we chose the earliest stages, immediately before and after embryogenesis induction, where the most dramatic rearrangements at the structural, ultrastructural and molecular levels have been described (Maraschin et al., 2005; Seguí-Simarro and Nuez, 2008). These stages comprise: (1) vacuolate microspores and young bicellular pollen just extracted from the anther but prior to exposure to the inductive heat shock (Figure 1A, 2A) and (2) just after induction (Figure 1B), where the first sporophytic divisions are seen still within the exine coat (inset in Figure 1B); (3) induced, early microspore-derived embryos (MDEs; Figure 1C), once released from the exine coat and with two clearly differentiated suspensor and embryo proper domains; (4) globular MDEs (Figure 1D), where the first signs of tissue differentiation are visible in the embryo proper domain (inset in Figure 1D); and (5) heart-shaped MDEs (Figure 1E, 2B), where the embryo pattern of symmetry changes from radial to bilateral, hypocotyl and cotyledon begin to elongate and differentiation of internal tissues takes place (Tykarska, 1980). For pollen development, we selected (1) the vacuolate microspore, before the first pollen mitosis, (2) the young pollen grain (Figure 3A) immediately after the first pollen mitosis, (3) the mid pollen grain (Figure 3B), when the generative cell displaces towards the center of the vegetative cell, and (4) the late, mature pollen grain (Figure 3C), when sperm cells are formed after the second pollen mitosis, still within the anther locule.
Nuclear size, chromatin condensation pattern and nucleolar architecture

The nucleus of vacuolate microspores was typically round or slightly oval, large and off-centered (Figures 1A inset, 2A) due to the presence of a large central vacuole. Chromatin displayed the decondensed pattern typical of this species, with only few, large masses of condensed chromatin often associated to the inner side of the nuclear envelope (Figure 2A), allowing for an abundant interchromatin region. In induced microspores, the first embryogenic division was symmetric and gave rise to two similar nuclei, equivalent in size (Figure 6C), rounded and centrally positioned. Subsequent divisions of the embryo proper cells produced progressively smaller nuclei (Figure 4A). This nuclear morphology was observed, with slight differences, in interphasic nuclei of microspores and MDEs through the early stages of embryogenesis. The chromatin pattern in these cells could be defined as dark, electron dense masses in electron microscopy sections of samples treated with the nucleic acid-contrasting MA cytochemistry (Figures 2A, 2B), as clear, bleached masses (Figure 5A) in sections treated with the EDTA regressive cytochemical staining, or as intense, peripheral bright spots in nuclei stained with the fluorescent, DNA-specific DAPI staining (Figures 6A-D).

This pattern strikingly contrasted with that observed in developing pollen under non-inductive conditions (Figure 3). Immediately after the first pollen mitosis (Figure 3A), a large, vegetative (VN) and a smaller, generative nucleus (GN) appeared within the vegetative and generative cells. As pollen maturation proceeded, the VN retained an architecture comparable to that described for vacuolate microspore nuclei (compare Figures 3B-B’, 2A). However, as the generative cell moved from the periphery to the center of the
vegetative cell, the size of the GN dramatically decreased (Figure 3B; 4B) and chromatin adopted a highly condensed status, appearing as large bleached masses after EDTA staining (Figure 3B’). In the mature pollen grain, the VN showed signs of increased chromatin condensation (Figure 3C’), while the nuclear envelope adopted irregular, lobed morphologies (Figure 3C). After the second pollen mitosis, the two newly formed sperm nuclei occupied nearly all of the volume of the sperm cells (Figure 3C). Chromatin condensation reached the highest levels, appearing in MA-treated samples as large, densely stained masses (Figure 3C) and as bleached areas in EDTA samples (Figure 3C’’).

It is well established that changes in the transcriptional status of the cell are accompanied by significant extensive changes in nucleolar volume and morphology. (Raska et al., 1983; Goessens, 1984). Thus, in this work we used the nucleolar organization as a reliable marker to identify cells at different transcriptional status. At the vacuolate microspore stage, the nucleolus was round, centered, large with respect to the total nuclear volume (Figure 2A), and displaying a torus-shaped profile typical of highly active cells: a large central vacuole with ribonucleoprotein particles (RNPs), and abundant granular component (GC) intermingled with dense fibrillar component (DFC). In the DCF, many small and homogeneous fibrillar centers (FCs) were observed (Figure 2A). In MDEs, the nucleolar architecture appeared more variable. Indeed, besides the torus-shaped profiles indicative of highly active nucleoli, profiles indicative of moderate and low activity were also observed in other cells. Profiles of moderate activity were defined by the presence of many small, homogeneous FCs within a DFC with abundant GC (data not shown). Low activity profiles included few, large and heterogeneous FCs, and scarce GC, usually located at the periphery of the DFC (Figure 2B). During pollen development, the nucleolus of the
generative cell frequently showed low activity profiles just after the first pollen mitosis (Figure 3A), whereas profiles of moderate activity were evident in the vegetative nucleus up to the mid pollen stage (Figure 3B). In mature pollen grains, the nucleolus was reduced to a compact, dense and inactive structure in the generative nucleus and in sperm cells (Figure 3C). In the vegetative nucleus, nucleoli were visible up to the mid pollen stage.

Cytochemical analysis of the interchromatin region

In MA-treated samples, condensed chromatin masses showed higher contrast than the nucleolus and interchromatin RNP structures (Figure 5A), being the chromatin patches clearly defined. EDTA nuclear staining revealed some interesting and novel features of the plant IR. At a first visual inspection, the areas of EDTA-bleached, condensed chromatin masses (Figure 5B) were found apparently larger than their equivalents in MA (non-EDTA) treated samples (Figure 5A). To verify this observation, we quantitatively evaluated the areas of large chromatin masses in 100 randomly chosen MDE cells from EDTA and MA treated sections. We found that the average area of a bleached chromatin region in EDTA-stained samples was $0.47 \pm 0.29 \mu m^2$, whereas in MA stained samples it was $0.37 \pm 0.23 \mu m^2$. This difference (~21%) indicated that EDTA bleaches not only chromatin masses, but also the adjacent, perichromatin region where probably abundant chromatin fibres of different thickness are preferentially located. The use of the NAMA-Ur cytochemical method, preferential for DNA (Testillano et al. 1991), clearly revealed stained numerous chromatin fibres of different thicknesses at the periphery of the chromatin masses (Figure 5C, arrows). The comparison of these cytochemical techniques suggested the existence of a perichromatin region, rich in decondensed chromatin.
EDTA also revealed the presence of a dense fibrillo-granular RNP network filling the nucleoplasmic space between chromatin masses. In all of the embryogenic and gametophytic stages studied, we observed the presence of perichromatin fibrils (9.9±1.24 nm wide), either emerging from the mass borders or as isolated threads embedded in the IR (Figure 6B). Isolated granules, 37.8±5.96 nm in diameter, were observed dispersed through the IR, mostly in induced microspores and MDEs. They were frequently seen at the nuclear periphery, close to the nuclear envelope (Figures 6D, E). Their size, spherical shape, staining properties and principally the presence of a defined halo around them, allowed us to identify them as perichromatin granules according to (Monneron and Bernhard, 1969).

Smaller (22.3±5.53 nm in diameter) spherical granules were also found in the IR (Figures 6A, C), both isolated and grouped into clusters over a fibrillar matrix (Figure 5C). According to their distribution, these granules were likely interchromatin granules (IG). Correspondingly, clustered IGs would be interchromatin granule clusters (IGCs), the electron microscopy equivalents of the nuclear speckles observed by fluorescence microscopy (Lamond and Spector, 2003). In some sections we were able to identify a clearer, mostly fibrillar region devoid of IGs but always adjacent to IG clusters (Figure 6C, dotted lines). The structure of these regions highly resembled the “interchromatin granule-associated zone” previously described in mammalian cells (Visa et al., 1993).

IGCs were evident throughout microspore embryogenesis. First in dividing microspores, and to a lower extent in MDEs. During pollen development, IGCs were observed soon after the first pollen mitosis (Figures 3A-A”). However, their presence was progressively reduced during pollen maturation. Isolated IGs were observed as dark, well
defined, isolated particles through the progressively reduced IR of the early (Figures 3A- A''), mid (Figures 3B-B'') and finally mature pollen grain (Figures 3C-C''). This was paralleled by a progressive decrease in nuclear size and a remarkable increase in the condensation status of chromatin. The intensity of the transformations was higher in the generative nuclei, GNs (compare Figures 3B-B''), and in sperm cells (Figure 3C''), and with less intensity in the vegetative nuclei, VNs. In summary, these results indicated a relationship between the increase in chromatin condensation, the IR reduction and a change in the distribution pattern of IGs from clustered to isolated, during pollen maturation.

Nuclear immunolabeling with anti 2,2,7 – trimethylguanosine

Further characterization of the IR substructures involved immunolabeling with antibodies against 2,2,7-trimethylguanosine (TMG), the specific cap present in the RNA of the small nuclear RNPs (snRNPs). SnRNPs are major components of the splicing machinery (Lührmann, 1988). Anti-TMG signal was observed in all of the embryogenic stages studied during MDE development. Immunofluorescence signal (Figure 7) was located at the nucleus in the form of a faint signal and some discrete, round and intensely bright foci. These foci never colocalized with those observed for DAPI staining (compare Figures 7A-E and 7A'-E'), indicating that chromatin is excluded from them. The nucleolus always appeared devoid of fluorescence as well, indicating that these structures were confined to the IR and most probably corresponded to nuclear bodies. During pollen maturation, the faint fluorescence was negligible, and very few foci could be observed after the first pollen mitosis, mainly in the VN (Figure 7B'). As the chromatin of the pollen nuclei condensed, the TMG-labeled structures were less evident, becoming no longer
detectable in mature pollen (Figure 3C). Therefore, we focused on the study of these structures during MDE development, where intense foci were present (Figure 7C-E’).

Anti-TMG immunogold labeling confirmed a faint labeling throughout different regions of the IR. At the boundaries between the chromatin masses and the IR, significant labeling was found (Figure 8C, arrowheads), possibly detecting snRNPs associated to nascent transcripts. The nucleolus was nearly devoid of labeling. Labeling of the IR appeared either as isolated particles or as clusters of few (2-6) particles (Figures 8A-C). By combining immunogold labeling with EDTA staining it was shown that particle clusters decorated EDTA-positive, RNP-containing IR regions (Figure 8D, arrows), in a pattern similar to that described for IGCs. Thus, the clusters of gold particles of the IR would actually correspond to the granular IGC regions observed in Epon-embedded samples, although with a different appearance, due to the different processing, as described previously in other studies of the IR (Testillano et al., 1993).

In addition to small clusters, immunogold particles concentrated massively over dense, coiled-looking bodies (Figures 8A-E) that according to their size and position corresponded to the bright foci observed with immunofluorescence. These bodies frequently exhibited a nearly circular section, variable in size, and a thick fibrillar texture embedded into a light matrix continuous with the IR. Immunogold labeling decorated RNP-positive, particle-like structures contained within the matrix of the body (Figure 8D). The structure, size, nature and position of these bodies, together with the labeling with anti TMG, indicative of presence of RNPs, are characteristics of Cajal Bodies (CBs; (Fakan et al., 1984; Spector et al., 1991; Spector, 1993a; Seguí-Simarro et al., 2006). In addition,
where the sectional plane was appropriate, we could also observe a clearer, homogeneous
and scarcely labeled region associated to the CB (cb-az in Figure 8E), and structurally
different from other subnuclear structures. The structural characteristics of these novel
regions, not described to date in plants, resembled the *cleavage bodies* described as
associated to CBs in animal cells (Schul et al., 1996; Gall, 2000).

**Qualitative and quantitative analysis of Cajal bodies**

According to the criteria exposed above, we identified CBs in all of the
embryogenic stages studied. Interestingly, in freshly isolated microspores we noticed a
relatively lower electron density of CBs compared to CBs of other stages (compare Figures
8A and 8B). Most of the CBs in just induced microspores (not shown) and early globular
MDEs (Figure 8B) presented in general a similar appearance in terms of electron density
(high) and circularity (reduced). In late globular and heart-shaped MDEs, circular, electron
light CBs were more abundant.

For the quantitative analysis of CBs, we first estimated the number (Figure 9A) and
area (Figure 9B) of CBs at each stage in order to have a clear view of their abundance
during MDE development. Given the differences in nuclear size at different stages (Figure
2C), we normalized the nuclear density and size of CBs expressing them as number or area
of CBs per nuclear µm². Figure 9A shows that microspores prior to induction presented
comparatively less CBs than any other stage. Commitment of microspores to
embryogenesis was paralleled by an increase in the number of CBs, peaking in early
globular MDEs. In late globular and heart-shaped MDEs, when tissue differentiation starts,
the number of CBs showed a similar, reduced value, slightly higher than in microspores. A completely opposite trend was reflected in mean CB area calculations (Figure 9B). After induction, the size of CBs was drastically reduced, increasing progressively through the following stages of embryogenesis. Thus, an inverse relationship between CB size and number follow was evident (compare the trend lines of Figures 9A and 9B). After measuring the total area occupied by CBs at each stage (Figure 9C), we could confirm this relationship by observing that the total area was nearly constant except for one stage. In induced microspores, the total CB area was remarkably lower.

An additional unexpected finding arose when we compared the error bars of the different stages. Since the measured area of a nearly spherical object like a CB depends on the sectional plane, one would expect an error proportional to the measured area, and similar for all of the stages. However, in freshly isolated microspores, the error was comparatively smaller than in the other stages, suggesting that CBs in this stage are larger and more uniform in size. To check this out, we analyzed the frequency distribution of the areas of all of the CBs from the different stages considered altogether (Figure 9D). We found that areas did not exhibited a gaussian distribution, as expected for a uniform size. Instead, they showed a bimodal pattern, with measurements congregating around two peaks at 0.15 and 0.35 µm², which would correspond to estimated diameters of 0.44 and 0.67 µm, respectively. Thus, it appeared that CBs may have two different sizes, large and small, which according to our micrograph analysis correspond to the electron light (Figure 9E) and electron dense (Figure 9F) types above described, respectively. CBs of freshly isolated microspores would belong almost exclusively to the large and electron light type, whereas after induction both types may coexist, being the small more frequent in early post-
inductive stages and the large more frequent in later embryogenic stages. Once established
two different CB categories, we reexamined the presence of the CB-associated zones, and
interestingly, they were exclusively found associated with CBs of the small and dense type.
In summary, it appeared that CBs change their size, number, shape, and position in a
defined manner during early MDE development, whereas they dissapear during pollen
maturation. These data would be indicating that the described changes are a reflection of
the change in developmental program.
Discussion

The comparison between the gametophytic and sporophytic pathways followed by the microspore permitted us to analyse the remodelling of the nuclear domains in plant differentiating cells when switched to proliferation. The change of the developmental program and the activation of the proliferative activity affected the functional organization of the interchromatin domain, which accordingly changed their architecture and functional state, and redistributed its components. A summary of the main nuclear changes observed is shown in figure 10.

In this work, we present an ultrastructural, cytochemical and immunocytochemical characterization of the changes in nuclear architecture, with special attention to the IR-housed structures, during two different developmental processes with the same starting point: differentiation of microspores into mature, gamete-harboring pollen grains, transcriptionally inactive (Bednarska, 1984), and proliferation of embryogenesis-induced microspores during the first embryogenic stages, highly engaged in transcription (Joosen et al., 2007; Malik et al., 2008). The comparison of the changes undergone by the structures of the IR during these two processes allowed us (1) to relate the remodeling of the interchromatin domain to the change of developmental program and to variations in transcriptional activities, and (2) to reveal novel features of the plant nucleus, not yet described.
Changes in chromatin pattern parallel the change of developmental program

Numerous studies have shown a direct relationship between nuclear architecture and the level of cellular activity (Risueño and Testillano, 1994; González-Melendi et al., 1998; Testillano et al., 2000, 2005; Raska et al., 2004). In this work, we showed that the chromatin pattern is also affected by the induction of microspore embryogenesis in B. napus, constituting a good marker of developmental fate. In B. napus microspores committed to embryogenesis, all nuclei show a similar decondensed chromatin pattern, allowing for an abundant IR. The scarce chromatin masses are present mostly associated to the nuclear envelope. This pattern, similar to that of somatic cells, could be considered as a variation of the chromomeric condensation pattern (Jordan et al. 1980). This pattern changes in pollen cells, where a strikingly different, highly condensed chromatin pattern is observed. The embryogenic switch originates in the microspore and MDEs a characteristic organization with less condensed chromatin pattern the chromatin, typical of proliferating cells, and markedly different from developing microspores not committed to embryogenesis.

Changes in IR structures associated to induction of embryogenesis

In this work, we have shown evidences of a remodeling of the IR-housed structures as a consequence of the developmental switch of the induced microspore. IGs change from grouped as IGCs mostly in dividing microspores (transcriptionally active), to isolated in maturing pollen and gametes (transcriptionally inactive). In mammalian cells, various
reports showed that a transcription blockage resulted in the accumulation of splicing factors in larger speckles/IGCs (Spector et al., 1983, 1991; Melcak et al., 2000; Docquier et al., 2004). Nevertheless, in plant cells the organization of the IGs slightly differs. IGs have been observed as isolated granules in many plant cell systems (Testillano et al. 1993), being the splicing factors distribution more spread throughout the IR, as revealed by immunofluorescence assays. In the generative and sperm cells, the nuclear size progressively decreases as the chromatin adopts an extremely condensed pattern, which dramatically reduces the space available for the IR to a minimal amount (Figures 3B””, C’”). This lack of space may likely be affecting IGC nucleation. In Arabidopsis trichome cells, with extremely high endoreduplication levels, a transcription blockage causes the appearance of thousands of microspeckles, instead of the expected larger speckles (Fang et al., 2004). In parallel, the inconspicuous presence of IGs in dividing microspores could likely be related to the application of a heat shock treatment to induce the developmental switch. Indeed, heat shock has been shown to promote the accumulation of plant splicing factors in larger speckles/IGCs (Docquier et al., 2004).

Other features of the IR, which were associated with the change of developmental program, affected PGs and PFs. The increased number of PGs observed after the embryogenesis-inductive heat-shock would support a similar role for plant PGs on the accumulation of heat shock-damaged pre-mRNA molecules than in mammalian cells (Puvion and Lange 1980). In HeLa cells (Chiodi et al., 2000) showed the formation of PG clusters upon heat shock exposure. However, we could not identify such a clustered pattern in any of our plant samples. EDTA staining revealed a perichromatin region of decondensed chromatin, rich in decondensed chromatin which would correspond to active
genes ready for transcription and/or replication. The RNP fibrils (PFs) were observed further away from the chromatin masses, and would correspond to the different stages of hnRNA transcription and processing. Moreover, we identified a distinct region of ribonucleoproteic nature, structurally similar to the IG-associated zone (IG-AZ) of HeLa cells (Visa et al., 1993). The IG-AZ is considered as a part of the nuclear matrix that contains p80-coilin and U1 snRNA (but not U2), and is thus thought to be a reservoir of spliceosome components (Puvion-Dutilleul et al., 1995). These zones have never been described in plants before, and their presence in certain cells could be indicating a higher transcriptional activity. Nevertheless, further structural and immunochemical studies would be necessary to characterize the functional role of this nuclear region in plants.

Novel features of plant Cajal Bodies

The combination of anti-TMG immunolabeling with ultrastructural cytochemistry allowed us to identify several novel features of the plant CB. One of them is the identification of two ultrastructurally different types of CBs in B. napus. Despite of their differences, our structural and quantitative data reveal a number of similarities between them highly suggestive of a similar nature for both. Indeed, they may actually be the divided and fused forms of a same subnuclear structure, since it is known that CBs are highly dynamic structures, undergoing frequent movements from/to the nucleolus, and also fission and fusion between them (Lafarga et al., 1998; Boudonck et al., 1999; Platani et al., 2000; Navascués et al., 2004). This plasticity may be the origin of the heterogeneity in CB structure found in the B. napus nucleus, as well as in other plant cell types such as maize.
root cells (Docquier et al., 2004), heterogeneity also observed in some animal cells (Liu et al. 2006).

We have identified a clearer, large CB-associated zone (Figure 7E) always contacting CBs of the small, dense type. These regions are structurally similar to those described in animal cells as CB-associated cleavage bodies. due to the presence of, among others, factors for the cleavage of the 3’-end of polyadenilated mRNAs (Schul et al., 1996; Gall, 2000). A role for cleavage bodies was proposed related to the processing of specific pre-mRNA subsets not found in their associated CBs (Schul et al., 1996). Since the functional role of CBs is similar in plant and animal cells, it is likely that these plant CB-associated zones could have a similar role as well, even though more studies, combining structural characterization with in situ molecular localization, are needed to further characterize this novel compartment. Ultrastructural studies have found CBs and IGs associated in proliferating plant cells (Moreno-Díaa de la Espina et al. 1982); since CBs are mobile bodies and we have found similar AZs associated to IGs, the possibility that the three structures, CBs, IGs and AZ were associated cannot be excluded, even though it is very difficult to find the three in the same section.

It is known that the number of CBs in a cell is developmentally and cell cycle-regulated (reviewed in (Shaw and Brown, 2004), cell type-specific (Raska et al., 1991) and tightly related to the changes in transcriptional and metabolic activity (Carmo-Fonseca et al., 1992; Spector, 1993b). Our results showed that the number and size of CBs are tightly related to proliferation, in a direct manner for number, and inversely for size. The opposite scenario would apply for the relationship between CB size and number during
differentiation events. The only exception to this trend was observed in induced microspores exiting from a recently applied heat shock, where clear deviations were observed in individual and total CB size (Figs. 9B, C). Indeed, exposure to a heat shock induces the formation of smaller CBs in different animal cells (Handwerger et al., 2002; Carmo-Fonseca et al., 1992; 1993). Similar evidences have also been observed in HeLa cells, where exposure to high temperatures produced smaller CBs, even losing their contents in snRNPs.

It has been proposed in sugarcane root primordial cells a model where the number of CBs per nucleus would be fixed for a given species and cell type, independent of its activity (Acevedo et al., 2002). Our results would be in agreement with this notion, provided that what remains constant is the total “mass” of CBs, not only their number, and under non-stressing conditions. Taken together, these results would be indicating that the total CB “mass” within a nucleoplasm remains nearly constant during MDE development in the absence of stress. Upon stressing the cell, CBs would fragment into mini-CBs and dismantle, but in a reversible manner, recovering the total CB “mass” the normal levels after stress.

Concluding remarks

In this work we have shown that the different structures involved in RNA processing change their shape and distribution in a characteristic manner as a consequence of the reprogramming of the microspore from the natural, gamete-producing route to the in vitro, embryo-producing route. Some features of the IR were heat shock-derived
consequences. After heat shock CBs reduce considerably their individual size and their
total volume, although in a reversible manner. The increased occurrence of IGCs in induced
microspores could also be a heat shock effect.

Other IR changes were associated with the commitment to proliferation (in MDEs)
or to differentiation (in maturing pollen). The change of IR volume, the distribution pattern
of IGs, and the number and size of CBs were related to a change in the transcriptional status
of the cell. Moreover, novel features of plant IR-housed structures have been found, such as
the presence of two IR zones specifically associated to IGCs and CBs, the existence of
heterogeneity in the structure of the plant CB, with a smaller and denser type, and a larger
and lighter type. A significant remodelling of the microspore transcriptome has been
recently described as a essential part of the microspore switch to embryogenesis (Hosp et
al., 2007a, 2007b; Joosen et al., 2007; Malik et al., 2007). While chromosome remodelling
during reprogramming (Hosp et al., 2007a) would act as a pre-translational mechanism for
regulation of gene expression, the changes described in the present work of the structures
associated to RNA processing machinery would serve as a first post-translational regulatory
mechanism.

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References


Figure legends

Figure 1: Early stages of microspore embryogenesis. A: Freshly isolated microspores and young bicellular pollen. B: Induced microspores. C: Early (octant) MDE. D: Globular MDE. E: Heart-shaped MDE. Insets in figures A-D are sections of microspores and MDEs at the corresponding stages. Bars in A, B: 10 μm. C-E:: 50 μm.

Figure 2: Overview of the nuclear architecture during microspore embryogenesis: Vacuolate microspore (A) and heart-shaped MDE nuclei (B) in MA-treated sections. Note the remarkably different organization of the different components of the interchromatic region (ir) in both cell types. cb: Cajal body; chr: condensed chromatin; dfc: nucleolar dense fibrillar component; ex: exine coat; fc: nucleolar fibrillar center; gc: nucleolar granular component; ne: nuclear envelope; nu: nucleolus; nv: nucleolar vacuole; v: vacuole. Bars: 500 nm.

Figure 3: Changes in nuclear architecture during pollen maturation. The A, B and C series of figures cover the changes in nuclear organization undergone by young bicellular pollen, mid bicellular pollen and mature, tricellular pollen respectively. The A’B’C’ and A’’B’’ columns show the IR of the vegetative and generative nucleus respectively. C’’ correspond to a sperm nucleus of tricellular pollen. All figures but C are from EDTA-treated sections. See text for further details. Bars in A-C: 5 μm. A’-C’’: 500 nm.

Figure 4: Morphometric analysis of nuclear size. Estimation by the average nuclear area during microspore embryogenesis (A) and pollen maturation (B). Bars
represent the average nuclear area over TEM micrographs measured in μm². Mic: isolated microspores; Ind: induced microspores; EG: early globular MDE; LG: late globular MDE; HS: heart-shaped MDE; Nv: vegetative nucleus; Ng: generative nucleus. See text for further details.

Figure 5: The condensed chromatin and the perichromatin region after cytochemical techniques for nucleic acids (MA), RNPs (EDTA) and DNA (NAMA-Ur) in globular MDEs. Nuclear regions showing a chromatin mass. A: MA staining for nucleic acids; condensed chromatin (chr) displays higher contrast than the nucleolus (nu) and interchromatin RNP structures. B: EDTA staining for RNPs; condensed chromatin (chr) appears bleached whereas the nucleolus (nu) and RNP fibers and granules of the IR show contrast. C: NAMA-Ur staining for DNA; only condensed chromatin masses (chr) and chromatin fibres at the perichromatin region (arrows) and in the IR show high contrast, whereas the nucleolus (nu) and cytoplasm (ct) show no contrast at all. Bars: 300 nm.

Figure 6: The interchromatin region in EDTA-treated sections of globular MDE cells. A: nuclear overview showing the nucleolus (nu), the bleached chromatin masses (chr) and the interchromatin region (ir). B, C: magnifications of the regions boxed in A. Arrowheads in B: perichromatin fibrils. Dotted line in C: interchromatin granule-associated zone. D: interchromatin region with perichromatin granules. E: magnification of the area boxed in D. Arrows point to perichromatin granules. ne: nuclear envelope. Bars in A: 500 nm. B-E: 200 nm.
Figure 7: Anti-TMG immunofluorescence over cryosections during gametophytic and embryogenic development. A-E: DAPI staining. A’-E’: anti-TMG. Yellow arrowheads point to CBs. Note the absence of DNA in the corresponding regions in DAPI-stained images (compare white arrows in A and A’). The asterisk in A marks the massive vacuole of the microspore. Bars: 10 μm.

Figure 8: Anti-TMG immunogold labelling in microspores and MDEs. A: interchromatin region (ir) of a vacuolate microspore showing an intensely labelled Cajal body (cb). B: Interchromatin region of an early globular MDE showing abundant labelling over one electron-light (centrally located) and two electron-dense CBs (at both sides). C: Overview of the interchromatin region of a heart-shaped MDE, showing a large, round and circular CB between the nucleolus (nu) and a mass of condensed chromatin (chr). Arrows indicate clusters of gold particles over the IR, whereas arrowheads point to particles decorating the borders of chromatin masses. D-E: EDTA-treated sections where CB-marking gold particles concentrate into small clusters on the IR (arrows), over a EDTA-positive (RNP-containing) CB (outlined in D), and also over a CB-associated zone (cb-az). ct: cytoplasm; ex: exine coat; ne: nuclear envelope. Bars: 200 nm.

Figure 9: Quantitative analysis of Cajal bodies during microspore embryogenesis. A: mean CB number per nuclear μm². B: mean individual CB area per nuclear μm². C: mean total CB area per nuclear μm². D: Frequency distribution, expressed in percentage, of the areas of the different CBs observed. See text for further details. E: example of a large, electron light CB. F: example of small, electron dense CB. Mic:
isolated microspores; Ind: induced microspores; EG: early globular MDE; LG: late globular MDE; HS: heart-shaped MDE. Bars: 200 nm.

Figure 10: Summary of the changes in the chromatin condensation and IR structures in both pollen programs: pollen development (differentiation process) and microspore embryogenesis (proliferation process). IGs: Interchromatin granules, CBs: Cajal bodies. –: absence, +/-: low amount, +: present, ++: abundant, +++: high abundant.
Figure 7

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Figure 10