Periodic expression of Sm proteins parallels formation of nuclear Cajal bodies and cytoplasmic snRNP-rich bodies

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Abstract Small nuclear ribonucleoproteins (snRNPs) play a fundamental role in pre-mRNA processing in the nucleus. The biogenesis of snRNPs involves a sequence of events that occurs in both the nucleus and cytoplasm. Despite the wealth of biochemical information about the cytoplasmic assembly of snRNPs, little is known about the spatial organization of snRNPs in the cytoplasm. In the cytoplasm of larch microsporocytes, a cyclic appearance of bodies containing small nuclear RNA (snRNA) and Sm proteins was observed during anther meiosis. We observed a correlation between the occurrence of cytoplasmic snRNP bodies, the levels of Sm proteins, and the dynamic formation of Cajal bodies. Larch microsporocytes were used for these studies. This model is characterized by natural fluctuations in the level of RNA metabolism, in which periods of high transcriptional activity are separated from periods of low transcriptional activity. In designing experiments, the authors considered the differences between the nuclear and cytoplasmic phases of snRNP maturation and generated a hypothesis about the direct participation of Sm proteins in a molecular switch triggering the formation of Cajal bodies.

Keywords SnRNP ∙ Cytoplasmic bodies ∙ Cytoplasmic granules ∙ Nuage ∙ Meiosis ∙ Plant cell

Introduction

Eukaryote nuclei contain dynamic structures called Cajal bodies. Cajal first observed these nuclear bodies in rat neurons stained with silver salts using the Ag-NOR technique (Cajal 1903). By using electron microscopy, the Cajal bodies were revealed to consist of coiled fibrils that resemble a coil of twisted thread. For this reason, these nuclear domains have frequently been called “coiled bodies” (CB, Monneron and Bernhard 1969). Until the 1970s and 1980s, it was not known that Cajal bodies are also present in plants (Gorska-Brylass and Wrobel 1978; Moreno Diaz de la Espina et al. 1982). Currently, we know that CBs are evolutionarily conserved structures, which are found in the majority of animal and plant cell types (for review, see Shaw and Brown 2004; Cioce and Lamond 2005). Recently, plant Cajal bodies have been shown to also contain Atcoilin (Collier et al. 2006; Koroleva et al. 2009), which is a homolog of the CB marker protein coilin in animal cells. This finding suggests that CBs have a basic role in nuclear function in all eukaryotic cells; however, Cajal bodies in plants have been implicated as sites of siRNA biogenesis, whereas in animals siRNA dicing...
occurs in the cytoplasm (for review, see Pontes and Pikaard 2008). Recently, we have demonstrated the presence of poly(A) RNA in Cajal bodies (Kolowerzo et al. 2009). Such poly(A) RNA localization had not yet been observed in animal cells (Visa et al. 1993; Huang et al. 1994). Further study (Smolińska and Kolowerzo 2011) demonstrated the presence of several housekeeping gene transcripts in CBs. While CBs in all organisms contain components of the pre-mRNA splicing machinery, other machineries vary in CB localization. In contrast to the amphibian CBs (Wu and Gall 1993) and in HeLa cell nuclei CBs (Frey and Matera 1995), recent studies in Drosophila showed that the U7 snRNP is not concentrated in CBs but in a separate structure, the histone locus body (HLB) (Liu et al. 2006b).

One characteristic trait of Cajal bodies is the dynamic changes in their size and number in different species, individual tissues, and cells (Boudonck et al. 1998; Acevedo et al. 2002). Moreover, it has been shown that the number of CBs in plant cell changes during the cell cycle and differentiation (Boudonck et al. 1998; Straatman and Schel 2001; Seguí-Simarro et al. 2006; Zienkiewicz and Bednarska 2009).

It has been experimentally established that the number of Cajal bodies is related to the metabolic activity of a cell. A positive correlation between the formation of Cajal bodies and transcriptional activity was revealed in HeLa cells (Ferreira et al. 1994), in hamster embryos (Ferreira and Carmo-Fonseca 1996), in oocytes (Chouinard 1975; Parfenov et al. 2003) and during microspore embryogenic development (Seguí-Simarro et al. 2006), and pollen development (Zienkiewicz and Bednarska 2009) in plant cells. However, there are exceptions. In metabolically active cells like single-cell mouse embryos, CBs appear 16 h after fertilization, preceding the initiation of transcription that usually occurs in two-cell embryos (Ferreira and Carmo-Fonseca 1995). Cajal bodies are present in insect and mammalian oocytes, despite chromatin condensation in partially or completely inactive nuclei (Bogolyubov and Parfenov 2001; Parfenov et al. 2003; Batalova et al. 2005). Also a prominent Cajal body (endobody) was present during most of prophase, attached to the karyosome (Liu et al. 2006a) in the germinal vesicle of the Drosophila oocyte transcriptionally quiescent during the latter part of the first meiotic prophase. Moreover, an increase in the number of CBs was found in cells of hibernating dormice (Malatesta et al. 1994). The visualization of CBs in living plant cells suggests that significant changes in the number of Cajal bodies may be related to their ability to fuse to form larger CBs or divide into smaller ones (Boudonck et al. 1999).

Another strategy that cells use to increase the number of Cajal bodies is de novo formation. This complex process is strictly correlated with the small nuclear ribonucleoproteins (snRNPs) cycle (Navascues et al. 2004). snRNP biogenesis includes a sequence of events that occur in both the nucleus and cytoplasm. The first stage of this cycle is the synthesis of small nuclear RNA (snRNA) molecules in the nucleus. The U1, U2, U4, and U5 molecules are transcribed by polymerase II and acquire a 7-methylguanosine (m7G) cap at the 5’ end and additional nucleotides at the 3’ end as a result of the co-transcription process. The cap at the 5’ end is recognized by the cap-binding complex (CBC). This complex is necessary to transport the snRNAs to the cytoplasm. The PHAX protein (phosphorylated adapter for RNA export) is the second factor that promotes the export of snRNPs from the nucleus (Navascues et al. 2004). First, the binding of Sm proteins to the snRNA and the maturation of the 3’ end occur in the cytoplasm. Next, the snRNA acquires a 2,2,7-trimethylguanosine (m3G) cap at the 5’ end (Mattaj 1986; Dickmanns and Ficner 2005).

Despite the wealth of biochemical information, little is known about the spatial organization of snRNPs in the cytoplasm. We have used Larix microsporocytes as a model system to examine the organization of cellular snRNPs. In the cytoplasm of the larch microsporocytes, a cyclical detection pattern of bodies containing snRNA and Sm proteins was observed during anther meiosis. Here, we show that cyclic changes in the levels and distribution of Sm proteins are accompanied by the presence of bodies containing snRNA and Sm proteins in the cytoplasm and then an accumulation of snRNP occurs in the nuclei, leading to the formation of new Cajal bodies.

The mechanism responsible for managing the formation of Cajal bodies is coupled to the snRNP pathway. Such a conclusion is reached on the basis of the results from experiments using heterokaryons obtained by the fusion of primary human foreskin fibroblasts (DFSF1) with HeLa cells (Sleeman et al. 2001). During normal growth, DFSF1 cells are characterized by a lack of Cajal bodies. However, Cajal bodies form in the nuclei soon after fusion with HeLa cells. It was also determined that the exogenous overexpression of the SmB protein was sufficient for periodic induction of Cajal bodies in primary fibroblast cells (DFSF1).

To determine whether other snRNP factors observed in the Cajal bodies could induce de novo CB formation in larch, a U6 snRNA analysis was performed. Contrary to the other four “spliceosomal RNAs”, U6 snRNA is synthesized by RNA polymerase III (Kiss et al. 1987). In addition, U6 snRNA does not possess the trimethylguanosine cap at the 5’ end (Ro-Choi 1999). However, like the other snRNAs, its maturation and binding to other spliceosomal factors occurs in the Cajal bodies (Stanek and Neugabauer 2004, 2006). The nucleoli represent the first compartment to which U6 snRNA travels after its synthesis (Lange and Gerbi 2000). The inner nucleotides undergo methylation and
pseudouridylation with the participation of snoRNA (Tycowski et al. 1998; Ganot et al. 1999). Further in vitro studies have demonstrated that after several hours, the U6 snRNA is found in the CB (Gerbi and Lange 2002), where binding to U4 and U5 snRNA takes place and where the formation of the functional tricomplex occurs (Stanek et al. 2008).

In this study, we have investigated in a natural model the hypothesis that high Sm protein expression is responsible for managing the formation of Cajal bodies. In this model, CB formation is coupled to the snRNP pathway, which has not been previously demonstrated in plants.

In our model, larch microsporocytes exhibit natural fluctuations in levels of RNA metabolism. In the larch, meiosis is characterized by a 6-month-long period and a high anther synchronization level (above 95%). In these cells, the synchronous development and “timely and fast switching on and off” of RNA and protein synthesis allows the tracking of changes that occur during snRNA synthesis and maturation under “natural conditions”.

Materials and methods

Plant material

Anthers of Larix decidua Mill. were taken from the same tree in successive G2 and meiotic prophase stages, from leptotene to late pachytene, to ensure constant experimental conditions.

Antibodies

mAb 7.13 was a gift from Prof. F. Ramaekers, Department of Pathology, University of Nijmegen, The Netherlands, which was kindly provided by Dr. JHS Scheel, University of Wageningen, The Netherlands. The human polyclonal antibody AF-ANA recognizing the larch Sm proteins (Niedojadło and Górska-Brylass 2003) was a gift from Dr. Pombo, University of Oxford. mAb Y12 was a gift from Dr. K. Neugebauer, Max Planck Institute, Dresden, Germany and was kindly provided by Dr. J Niedojadło, N. Copernicus University, Toruń, Poland. The mouse monoclonal 4G3 antibody (Acris Antibodies, Herford, Germany) recognizing the spliceosomal protein U2B” (Habets et al. 1989; Boudonck et al. 1998) was used. The mouse anti-2,2′,7′ trimethyl guanosine (m3G) antibody from Calbiochem (clone K121) was used. This antibody cross-reacts with the specific 5′ cap of small nuclear RNAs (snRNAs).

Immunoprecipitation

Immunoprecipitation assays were carried out using nuclear protein extract from larch anthers. Fresh material was ground to a fine powder in liquid nitrogen. Nuclei were isolated using the CellLytic Plant Nuclei Isolation/Extraction Kit (Sigma, St. Louis, MO, USA), according to the manufacturer’s instructions.

Purified nuclei were resuspended in a buffer containing 40 mM Tris–HCl, pH 8.0, 420 mM NaCl, 1.5 mM MgCl2, 0.2 mM EDTA, 0.5 mM DTT, and 0.5 mM PMSF. The nuclei were subsequently shaken for 30 min and centrifuged (800g) for 5 min, and the nuclear protein extracts were isolated. For the immunoprecipitation procedures, we used the mouse anti-SmD 7.13 antibody. Briefly, the primary antibody (anti-Smd) and 1 mM PMSF, 1% Triton X-100, and 0.5% Nonidet P-40 were added to the nuclear protein extract and incubated for 4 h at 4°C. The sample was gently mixed using an orbital shaker. The antibodies were then bound to Protein A-Sepharose beads (Sigma) and incubated for 2 h at 4°C with 60% protein A-Sepharose in TBS and NP-40. The samples were centrifuged at 16,000g for 15 min at 4°C. The pellet was then washed with 1 ml washing buffer (TBS, 1% Triton X-100, and 0.5% Nonidet P-40). This step was repeated three times. Finally, the resulting pellet was resuspended in 50 µl Laemmli buffer and used for the SDS-polyacrylamide gel electrophoresis (PAGE) and western blotting procedures. SDS-PAGE was performed using 12% acrylamide gels with 4.5% stacking gels, using the Bio-Rad Mini-Protean equipment. Antigen–antibody complexes and total protein extracts were fractionated by SDS-PAGE (12%) and transferred to PVDF membranes. The membranes were incubated overnight at 4°C with the appropriate primary antibodies, specifically, anti-SmD diluted 1:5,000 or mouse anti-TMG, diluted 1:500. The membranes were washed with TBS (three times), probed with an anti-mouse secondary antibody (1:1,000) for 1 h, and developed using enhanced chemiluminescence (ECL system, Amersham).

Y12 Ab specificity assay

Total proteins were isolated from larch anthers. The ground tissue was resuspended in extraction buffer (100 mM Tris–Cl, pH 7.5, 1 mM EDTA, 1 mM DTT, 1 mM phenylmethylsulfonyl fluoride, and protease inhibitor cocktail, Sigma), gently shaken, and clarified by centrifugation at 16,000g for 30 min at 4°C. The supernatant was used for SDS-PAGE and immunoblotting using Y12 Ab (diluted 1:500). The membrane was then washed with TBS (three times), probed with an anti-mouse secondary antibody (1:1,000) for 1 h, and developed using enhanced chemiluminescence (ECL system, Amersham).

Standard transmission electron microscopy

Samples were processed for electron microscopy by conventional techniques (paraformaldehyde-osmium fixation.
and lead citrate-uranyl acetate staining (Niedojadlo and Görská-Brylass 2003).

Isolation of meiotic protoplasts

Anthers were fixed in 4% paraformaldehyde in PBS (pH 7.2) for 6 h and crushed to obtain free meiocytes. The cells were then centrifuged and resuspended in hydrolyzing medium, consisting of 4% cellulase Onozuka R10 (Serva, Heidelberg, Germany) and 27 U/ml pectinase (Sigma) in 0.01 M citric buffer (pH 4.8) for 45 min at 37°C. The suspended protoplast matter was spread on gelatin-coated slides that were placed on dry ice and air-dried. These protoplasts were rinsed with 0.1% Triton X-100 in PBS and for 10 min and air-dried. Next, the protoplasts were rinsed with 0.1% Triton X-100 in PBS (pH 7.2) for 10 min and air-dried. These protoplasts were then used for immunodetection of Sm, m3G snRNA, and U2B° and for fluorescence in situ hybridization of SmD1 mRNA, poly(A) RNA, U1 snRNA, U2 snRNA, and U6 snRNA.

Design of double-labeling reactions

Several double-labeling immunofluorescent-fluorescent in situ hybridization (FISH) or high-resolution immunogold in situ hybridization (ISH) reactions (Sm-U1, U2, and U6 snRNA; Sm-U2B°; BrU-poly(A) RNA; m3G snRNA-U2 snRNA) were performed as described below. In all experiments, immunocytochemistry always preceded in situ hybridization because immunofluorescent signals were very weak when ISH was applied first. In the double-labeling reactions, the obtained data were corrected for background autofluorescence determined by negative control signal intensities. For the image processing and analysis, the EZ Viewer software package (Nikon Europe BV, Badhoevedorp, The Netherlands) was used. The analysis was performed using Lucia General software (Laboratory Imaging, Prague, Czech Republic) compatible with a Nikon PCM 2000 confocal microscope and Nikon C1 confocal microscope. To test differences among multiple samples (groups, i.e., Sm level in different stages), a Kruskal–Wallis ANOVA test was used.

FISH in double-labeling reactions

For hybridization, the probe was resuspended in hybridization buffer (30% v/v formamide, 4 × SSC, 5 × Denhardt’s buffer, 1 mM EDTA, and 50 mM phosphate buffer) at a concentration of 50 pmol/ml. Hybridization was performed overnight at 37°C. The following DNA oligonucleotides were used: antisense U2 snRNA—5' Alexa 594 ATATTTAACTGATAAGAAGACTACTAGCTGGT 3', sense U2 snRNA—5' Alexa 594 CAAGTGTAGTATCTGTTGTTTAAATAT (IBB PAN, Warsaw, Poland), antisense U1 snRNA—5' Alexa 594 ACAGCAAACAGTTATGATGTTGC 3', sense U1 snRNA—5' Alexa 594 GAACCTACATCATAATTTGTTGCTGGT 3' (IBB PAN, Warsaw, Poland), antisense U6 snRNA—5' Alexa 594 CTAATCTTCTCTATGATATCATTCCAATTATTTAT 3', sense U6 snRNA—5' Alexa 594 TAAAATTGGAAC 3' (IBB PAN, Warsaw, Poland)

ISH immunogold monolabeling of U2 snRNA and double labeling with Sm proteins

After fixation in 4% PFA in PBS overnight, anthers were rinsed in PBS and dehydrated and mounted in LRGold resin (Sigma), according to the Majewska-Sawka and...
Rodriguez-Garcia (1996) schedule. Ultrathin sections were collected on Formvar-coated nickel or gold grids. Ultrathin sections were pretreated with pre-hybridization buffer (hybridization buffer without probe) for 1 h at room temperature and then incubated with the antisense U2 snRNA probe (5′- ATATTAAACTGATAAGAACAGATACTACACTTG 3′) in hybridization buffer (the probe was labeled at the 3′ end with digoxigenin nucleotides by terminal deoxynucleotidyl transferase) (Roche). Hybridization was performed in a sealed, humidified chamber for 20 h at room temperature. Post-hybridization washing was performed as described by Smoliński et al. (2007). Non-specific antigens were blocked with PBS buffer containing 0.05% acetylated BSA for 0.5 h. DNA–RNA hybrids were localized by incubation with sheep anti-digoxigenin antibody coupled to 10-nm-diameter colloidal gold particles (Orion Diagnostica, Espoo, Finland) diluted 1:30 in PBS containing 0.02% acetylated BSA for 1 h at room temperature. Grids were then rinsed and contrasted according to Smoliński et al. (2007).

In the double-labeling reactions, immunogold localization of Sm proteins was performed prior to ISH immunogold labeling, using the primary AF-ANA human anti-Sm protein B/B′, D1 antibody in 0.2% acetylated BSA in PBS (1:100) at 8°C in a humidified chamber overnight. Samples were incubated with secondary anti-human antibody coupled to 15-nm-diameter colloidal gold particles (BioCell, Cardiff, UK) and rinsed and contrasted (1% phosphotungstic acid and 2.5% uranyl acetate).

Microscopy

The results were registered with Nikon C1 and Nikon PCM 2000 confocal microscopes using an argon–ion laser emitting light with a wavelength of 488 nm (blue excitation and green fluorescence), a He–neon laser emitting light with a wavelength of 543 nm (green excitation and red fluorescence), and a He–neon laser emitting light with a wavelength of 594 nm (red fluorescence). A mid pinhole, long exposure time (75 µs) and a 60× (numerical aperture, 1.4) Plan Apochromat DIC H oil immersion lens was used. Pairs of images were collected simultaneously in the green (Alexa 488 or FITC fluorescence) and red (Cy3 or Alexa 594) channels. To minimize bleed-through between the fluorescence channels, we applied low laser power (3–10% of maximum power) and single-channel collection. For the bleed-through analysis and control experiments, Lucia G software was used (Laboratory Imaging, Prague, Czech Republic). For DAPI staining, an inverted Nikon Eclipse TE 2000 fluorescence microscope equipped with a mercury lamp, a UV-2EC UV narrow-band filter, and a DXM 1200 FX digital camera were then used.

The material was examined at the ultrastructural level, and photomicrographs were taken, using a Jeol 1010 (Japan) electron microscope at 80 kV.

Control reactions

For in situ hybridizations (high resolution) and immunofluorescence methods, control treatments consisting of incubations without the primary antibody were performed. For the in situ hybridizations—high-resolution and fluorescence analyses, sense-labeled probes and ribonuclease-treated samples were used as additional controls. All control reactions produced negative results, or the results of the control reactions were very low compared with the standard reactions.

Results

Determination of Sm and TMG snRNA antiserum reactivity

We used the mouse mAb 7.13, which recognizes the SmD protein of the Ul, U2, U4, U5, and U6 snRNPs in plants (Billings et al. 1982, 1985; Pálfi et al. 1989; Testillano et al. 1993; Straatman and Schel 2001). Immunoprecipitation and western blot analyses demonstrated that the mouse monoclonal anti-SmD antibody recognized a band with a molecular mass of about 55 kDa (Fig. S1). As a control, we used a primary monoclonal anti-SmD antibody without sample adding and we tested its direct cross-reactivity with the secondary Ab following the IP procedure.

Three bands were visible after immunoprecipitation with the anti-SmD antibody and immunoblotting with the SmD and anti-TMG cap Abs. Two prominent bands (60 and 28 kDa) represent the cross-reactivity after immunoblotting with primary anti-SmD and secondary antibody. These two prominent bands represent the heavy (upper band) and light (lower band) chains of the antibody, respectively. The weaker band cross-reacts with the primary anti-Sm Ab and the secondary antibody and represents the immunoprecipitated protein.

Similar results were obtained with an anti-TMG snRNA antibody, which recognized the same complex with a molecular mass of 55 kDa. This result indicated that the 55-kDa moiety was a snRNP complex containing SmD proteins and snRNA.

In the case of the anti-TMG cap Ab, two prominent bands also represent the heavy (upper band) and light (lower band) chains of the SmD antibody, which cross-react with secondary antibody. However, the weaker band cross-reacts with the primary anti-TMG Ab and represents...
the immunoprecipitated Sm proteins complexed with snRNA containing the TMG cap structure.

Moreover, because the two primary antibodies (SmD and anti-TMG) were obtained from mouse, the cross-reactivity observed in lane 3 also represents the heavy and light chains after probing with secondary (anti-mouse) Ab labeled with horseradish peroxidase.

The two bands are more prominent because the concentration of the SmD antibody used for immunoprecipitation was much higher than the cellular concentration of immunoprecipitated antigen in the larch anthers. A western immunoblot analysis was performed to examine the specificity of the mouse Y12 antibody, in *Larix decidua* which recognizes Sm B'/B and D1 proteins in plants (Hirakata et al. 1993; Echeverría et al. 2007). As shown in Fig. S2, the mouse Y12 antibody recognized proteins of about 30 and 14 kDa, which appear to represent the methylated SmB/B' and SmD1 proteins, respectively.

Localization of splicing factors during cyclic fluctuations in Sm protein quantities

In the analyzed stages, i.e., from G2 through pachytene, we could distinguish three cycles in which the levels and distribution of Sm proteins differed. Four A–D stages were distinguished in all three of the cycles. We named them A1–D1 in the first (G2) cycle, A2–D2 in the second (leptotene) cycle, and A3–D3 in the third (zygotene) cycle. The characteristic changes in the distribution of Sm proteins during the four stages of each cycle were demonstrated with the example of the first cycle of the G2 phase of premeiotic interphase. Two simultaneous analyses of the colocalization Sm proteins with m3G snRNA and with U6 snRNA were performed. Similar changes in snRNP distribution were observed in the two other prophase cycles (leptotene and zygotene one).

In the first stage (A1), the levels of Sm proteins were very low. In the cytoplasm and nucleoplasm, Sm proteins were present in a dispersed form (Figs. 1a, S3a). In some cases, small quantities of Sm proteins, in the form of a dispersed signal, were also observed in the nucleolus. The levels of m3G snRNA were also low at this stage (Fig. S3b), and the signal was dispersed in the nucleoplasm (Fig. S3b). The levels of U6 snRNA were also low, and U6 snRNA was mainly detected in the nucleoli (Fig. 1b).

In the next stage (B1), a significant increase in the level of Sm proteins was observed, and fine Sm bodies began to appear in the cytoplasm, some of which contained not only Sm proteins (Figs. 1c, S3e) but also m3G snRNA (Fig. S3f). The oval clusters of Sm proteins occurring in the cytoplasm of larch microsporocytes were called cytoplasmic snRNP bodies (CsBs). Many small, fine clusters of Sm proteins or dispersed Sm signals, which did not contain m3G snRNA, formed in the cytoplasm as well (Fig. S3g). Accumulation of Sm proteins (Fig. S3e) and m3G snRNA (Fig. S3f) was observed at the border between the nucleus and the cytoplasm or at the periphery of the cell nucleus (Fig. 1c) during this phase. A significant increase in levels of U6 snRNA was also observed in the nucleoli. In the nucleolus, the U6 snRNA exhibited a dispersed pattern (Fig. 1f).

In the third stage (C1), significant quantities of Sm and m3G snRNA were still observed in the cytoplasm in the form of large, CsBs (Figs. 1i, S3i–k), and Sm proteins reached their highest level in the nucleus (Fig. 1i). Sm proteins were present in the whole nucleus. Nuclear bodies were present in high numbers at stage C (Fig. 1i). The levels of U6 in the nucleolus were still significantly higher than in the nucleoplasm (Fig. 1j). We did not observe any accumulation of U6 snRNA in the nuclear bodies containing Sm proteins (Fig. 1j, k).

In the last stage (D1), there was a significant decrease in the levels of Sm proteins and m3G snRNA in the cytoplasm (Figs. 1m, S3m–o). In the cytoplasm, only single spots containing both Sm proteins and m3G snRNA were visible (Fig. S3m–o). In the nucleus, besides the evenly dispersed signal in the nucleoplasm, the Sm proteins accumulated mainly in numerous nuclear bodies (Figs. 1m, S3g). The maximal number and size of the nuclear bodies was reached in stage D1. U6 snRNA was present at similar levels in the nucleolus and the nucleoplasm; however, the highest levels were observed in the nuclear bodies (Fig. 1n). At this stage, a high level of colocalization of Sm proteins with U6 snRNA occurred in the nuclear bodies and in the nucleoplasm (Fig. 1o).

Cytoplasmic distribution of Sm proteins in larch microsporocytes

An analysis using the double immunofluorescence/FISH labeling confirmed that CsBs could contain individual types of snRNA. The colocalization of Sm proteins with U2 snRNA (Fig. 2a–d) and with U1 snRNA (Fig. 2e–h), demonstrated that some of CsB bodies containing Sm proteins also contained detectable levels of individual analyzed snRNAs. In addition to CsB localization, U1 and U2 snRNA were present as dispersed signals in the cytoplasm (Fig. 2b, f).

The distribution of Sm proteins exhibited a characteristic pattern in the cytoplasm of the microsporocytes. During periods when Sm protein levels in the microsporocyte were high (stages B and C), Sm proteins were frequently observed in the form of CsBs. Such bodies were detected during late G2 phase (Fig. 3a), late leptotene (Fig. 3e), and middle zygotene (Fig. 3i). An analysis using the double immunofluorescence labeling method
Fig. 1 Localization of Sm proteins (mAb Y12) and U6 snRNA during premeiotic interphase. Initially, very low levels of the Sm protein were observed (a). U6 snRNA was clearly visible only in the nucleoli (b, c). In the subsequent stage, cytoplasmic bodies containing Sm proteins appeared (e). In this period, an increase in the levels of these proteins was observed in the nucleus (e, g). The amount of U6 snRNA increased considerably in the nucleoli, but a significant amount was also detected in a dispersed form in the nucleus (f). In the next stage, when cytoplasmic bodies containing Sm proteins could still be observed in the cytoplasm, high Sm protein levels were observed in the nucleoplasm and in the nuclear bodies (i, k). Colocalization of U6 in the nuclear bodies was not observed. The levels of U6 in the nucleolus were still much higher than in the nucleoplasm (j). In the last stage, only very small, single Sm bodies were visible in the cytoplasm (arrow head). In the nucleus, strong Sm protein (m) and U6 snRNA (n) signals could be observed in the nuclear bodies (arrows), whereas the signal was uniform and dispersed in the nucleoplasm (o). The corresponding DAPI images were collected using widefield fluorescence and deconvolution software (d, h, l, p). Bars 10 μm.
demonstrated that many Sm protein-containing cytoplasmic bodies also contained small nuclear RNA. On the basis of the colocalization of Sm proteins with m3G snRNA, we determined that mature snRNAs can be found in most cytoplasmic Sm clusters (Fig. 3). In the cytoplasm, in addition to large CsB complexes, snRNA was also detected in the form of small grains or spots, but a dispersed localization of m3G snRNA was not observed (Fig. 3b, f, j).

An electron microscopy analysis allowed us to understand the ultrastructural characteristics of the CsBs that were observed in larch microsporocytes. Immunogold localization with the Y12 anti-Sm antibodies revealed that these oval bodies are the sites of the selective accumulation of Sm proteins (Fig. 4a). Therefore, these oval bodies correspond to the CsBs that were observed by confocal microscopy. In addition to the large bodies, the gold particles labeling the Sm proteins were also found in a dispersed form and in sparse clusters in the cytoplasm (Fig. 4a). Cytoplasmic bodies were also strongly labeled with the anti-m3G snRNA antibodies (Fig. 4b). These bodies were the main area of m3G snRNA accumulation in the microsporocyte cytoplasm. No scattered m3G snRNA signals were observed, whereas scattered signals were visible in the case of Sm proteins (Fig. 4a, b).

The presence of snRNA in the analyzed Sm-containing CsBs was confirmed through the use of a colocalization immunogold/ISH immunogold technique for Sm (Y12) and U2 snRNA (Fig. 4c). Furthermore, the cytoplasmic colocalization of Sm proteins and U2 snRNA was also observed in fine clusters (Fig. 4c).

Nuclear localization of Sm proteins in Cajal bodies

In microsporocyte nuclei, the distribution of Sm proteins often had a dispersed character (phase C, Figs. 1i, 3i), or an accumulation of Sm-snRNP occurred in the nucleus (phase D, Figs. 1m, S3m). Oval nuclear bodies (0.5–3 µm diameter) containing the Sm and U2B0 proteins and other splicing factors were often observed (Fig. 5a, b). Nuclear bodies with high concentrations of Sm proteins and snRNA were detected during stages from the G2 phase of premeiotic interphase through the prophase of the first meiotic division.

Nuclear bodies consisting of coiled fibrils, resembling a coil of twisted thread with dense coiled fibrils (with a diameter of 15–25 nm), were observed in larch microsporocytes (Fig. 5c). This substructure is a characteristic trait of the Cajal bodies found in plants and animals. The application of the immunogold technique at the ultrastructural level demonstrated that Cajal bodies were the place of Sm protein accumulation (Fig. 5d). Therefore, Cajal bodies corresponded to the oval clusters of signal that were observed by confocal microscopy. CBs were very often localized to various parts of the nucleus close to the

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**Fig. 2** Colocalization of U2 snRNA (a–d) and U1 snRNA (e–h) with Sm proteins (mAb 7.13) during premeiotic interphase (immunofluorescence/FISH method). Many of cytoplasmic structures contained Sm proteins; some CsBs possess only a small amount of snRNA or are devoid of snRNA (g). The corresponding DAPI images were collected using widefield fluorescence and deconvolution software (d, h) Bars 10 µm
nuclear envelope or near clusters of interchromatin granules (IGs) (Fig. 5d). The presence of U2 snRNA, m3G snRNA, and Sm proteins was demonstrated at the ultrastructural level in Cajal bodies using a double-label immunogold/ISH immunogold method. Large spherical cytoplasmic bodies (arrows) containing both Sm proteins and snRNA can be observed. During this period, numerous smaller clusters of Sm proteins were also present in the cytoplasm (arrowheads), but not always together with snRNA. The corresponding DAPI images were collected using widefield fluorescence and deconvolution software (d, h, l). Bars 10 μm

Fig. 3 Immunolocalization of Sm (AF-ANA) proteins and m3G snRNA in the G2 (a–d), leptotene (e–h) and zygotene (i–l) phases using the double-label immunofluorescence method. Large spherical cytoplasmic bodies (arrows) containing both Sm proteins and snRNA can be observed. During this period, numerous smaller clusters of Sm proteins were also present in the cytoplasm (arrowheads), but not always together with snRNA. The corresponding DAPI images were collected using widefield fluorescence and deconvolution software (d, h, l). Bars 10 μm

ultrastructural level (Fig. 5d–f). We found that Sm proteins and U2 snRNA were distributed preferentially in the CB (412.6 ± 96 and 149.5 ± 8.1 gold grains/μm², respectively), at significantly higher levels ($p < 0.05$) when compared with nucleoplasm areas (19.5 ± 7.2 grains/μm² and 6.7 ± 0.9 gold grains/μm²) during the G2 stage. We obtained similar results from a quantitative evaluation of Sm proteins and U2 snRNA distribution during the leptotene and zygotene cycles.
Levels of Sm proteins during the G2 phase of premeiotic interphase and early prophase during the first meiotic division

In this study, we performed a quantitative analysis of the emitted fluorescence from the snRNPs complexes. Immunofluorescence using anti-Sm Y12-Cy3 was employed to demonstrate that the levels of Sm proteins undergo significant fluctuations during the premeiotic G2 phase of interphase through the later pachytene in larch microsporocytes (Fig. 6). The stages of premeiotic interphase and meiosis were recognized on the basis of chromatin structure (DAPI staining) and measurements of nuclear volume and whole cell volume (during the studied period, the whole nuclear volume increased by over fourfold and the whole cell volume increased by more than fivefold).

High levels of Sm proteins were observed during three phases of microsporocyte development. The first of these phases occurred in interphase, at the end of the G2 phase. The other two phases were observed during late leptotene and middle zygotene (Fig. 6).

Discussion

Studies have revealed that during anther meiosis in the larch, spherical structures, which are not separated by a membrane, periodically appear in the cytoplasm. The absence of a membrane surrounding these cytoplasmic bodies suggests that a special mechanism must exist that facilitates their formation and protects them from the cellular autolytic system. Our studies have demonstrated that cytoplasmic bodies, which contain Sm proteins and snRNA, exist in the cytoplasm of larch meiocytes. These structures were named “cytoplasmic snRNP bodies” (CsBs). Until now, there have been no reports on similar phenomena in higher plants.

In animal germ cells, cytoplasmic granules that possess similar structural properties are called nuage (Chuma et al. 2003), P bodies (Eulalio et al. 2007) or P-granules (Schisa et al. 2001), and some of them may contain Sm proteins (Biliński et al. 2004).

In Xenopus oocytes, nuage contain not only the spliceosomal Sm proteins but also Xcat2 mRNA. Other components of Cajal bodies or the splicing machinery, such as coilin, SMN protein, and snRNAs were absent from the nuage (Biliński et al. 2004). The authors suggested that Xenopus Sm proteins have adapted to a role independent of pre-mRNA splicing and that instead of binding to their traditional spliceosomal partner, such as snRNA, they bind mRNAs that are components of the germinal granules (i.e. Xcat2 mRNA) and facilitate the transport of these mRNAs.
from the nucleus to the nuage, which are precursors of germinal granules.

Studies in *C. elegans* show that Sm proteins are components of P-granules that are required for germ-granule (P-granules) localization (Barbee et al. 2002; Barbee and Evans 2006). Non-spliceosomal roles for Sm proteins were also postulated during the maturation of spermatocytes and for germ-cell specification in *Drosophila* (Gonsalvez et al. 2006). Finally, Gonsalvez et al. (2010) have demonstrated that SmB and SmD3 are specific components of the oskar messenger ribonucleoprotein (mRNP) in *Drosophila*.

In the nuclei of *Drosophila* meiocytes Liu and Gall (2007) have observed that snRNA, Sm, and SMN proteins accumulate occasionally in structures that the authors called U bodies, which are located near P bodies. These structures could correspond to CsBs in the larch. The authors suggest that U bodies may be the sites for some steps of snRNP assembly or that they may serve primarily for the storage of snRNPs after assembly but before import into the nucleus. Further studies by the same authors (Lee et al. 2009) have demonstrated a relationship between SMN proteins that participate in snRNP complex maturation and the integrity of both U bodies, which contain this type of snRNP, and P bodies, which participate in the regulation and degradation of other types of RNAs in the cytoplasm. Cytoplasmic bodies that exist in the cytoplasm of larch meiocytes contain snRNA in addition to Sm proteins. The presence of Sm–snRNP in larch CsB could be in agreement
with the sequence of molecular events in the snRNP cellular cycle. In the cytoplasm, a cortex molecule comprised of Sm proteins attaches to snRNA, the cap is hypermethylated, and the 3' end matures (Will and Luhrmann 2001; Massenet et al. 2002).

Cytoplasmic snRNP bodies could serve as potential platforms for the formation of this complex in larch meiocytes. The colocalization of Sm proteins and snRNA in the larch cytoplasmic bodies supports such a hypothesis. In addition to bodies that contained both molecules, foci were also found that contained only Sm proteins. These small foci could represent compartments where the Sm cortex molecule is stored, or they could be a possible place for intensive synthesis of the Sm cortex molecule. The localization of SmD1 mRNA proved its local accumulation in the cytoplasm, which could correspond to the translation site of Sm proteins (data not shown). Using the hybridization method, we demonstrated that the U1 and U2 snRNAs exist in a dispersed form and in CsB clusters, where it almost always colocalizes with the Sm proteins. Slightly different patterns of localization were observed for snRNA detected by antibodies against the m3G cap. In this method, trimethylated snRNAs were observed in the cytoplasm. In the cytoplasm, using the colocalization method, m3G snRNA was found mainly in CsB clusters and always with Sm proteins. Almost no m3G snRNA was observed in the cytoplasm in a dispersed form or without colocalized Sm proteins. These observations indicate that after export to the cytoplasm snRNAs can travel in a dispersed form and then bind to small clusters of Sm proteins and form CsBs. After dimethylation, which presumably takes place in the CsB, snRNP complexes can move toward the nucleus.

Increase of Sm proteins and snRNA content and the formation of CsBs in all three cycles always preceded the appearance of Cajal bodies. Such a positive correlation indicates that the occurrence of both phenomena is not a random coincidence. To determine whether other snRNP factors observed in the Cajal bodies could induce de novo CB formation in larch, a U6 snRNA analysis was performed. Our observations have shown that in the early stages of the analyzed cycles, the U6 snRNA was detected in the nucleoli at the same time that the Sm-containing CsBs appeared. In the subsequent stages of the cycle, when the first Sm-containing CBs and m3G snRNA began to appear in the nucleoplasm, the U6 snRNA is localized, in addition to in the nucleolus, in a dispersed form in the nucleoplasm, but not in the CBs. The accumulation of U6 snRNA in the CBs was observed only in the later part of the cycle, when numerous and large CBs had already appeared in the nuclei. Contrary to the other spliceosomal RNAs, U6 snRNA is not necessary in the initial step of Cajal body formation in larch.

The formation of CsBs always preceded the appearance of Cajal bodies. Such a positive correlation indicates that the occurrence of both phenomena is not a random coincidence. High expression of Sm proteins, demonstrated by the appearance of foci containing this protein in the cytoplasm, induces the assembly of Sm–snRNP complexes in the cytoplasmic snRNP complex, which is where snRNA dimethylation likely occurs. In the next stage most likely cytoplasmic bodies transport to the nucleus membrane occur, which is evidenced by the accumulation of Sm proteins and CsB at the nucleus–cytoplasm border. A similar “bodies like” pathway of transport is also pointed
out by the ultrastructural images of *Xenopus* oocytes containing nuage (Biliński et al. 2004). In larch, this type of transport is also supported by the lack of a dispersed localization of m3G snRNA in the cytoplasm. If the cytoplasmic/nuclear Sm–RNP stays in a dynamic balance, a high level of Sm protein synthesis in the cytoplasm may cause an elevated accumulation of Sm–snRNP in the nucleus. A morphological manifestation of this process seems to be observed as the accumulation of diffusively dispersed Sm–snRNP in the sub-membrane space to the meiocyte nuclei. Exceeding the Sm–snRNP concentration over the threshold can trigger the interaction of these complexes with coilin homologues, whose presence has been confirmed in plants (Collier et al. 2006; Koroleva et al. 2009).

The interaction between the snRNP cycle and Cajal body formation has also been observed in the earlier studies in HeLa cells, where blockade of hTGS1, SMN, and PHAX protein expression was achieved by RNAi (Lemm et al. 2006). The knockdown of these three proteins disturbed the maturation of UsnRNP before its re-importation into the nucleus and caused a decrease in Cajal body formation. A similar decrease in the quantity of CBs has also been observed in HeLa cells exposed to leptomycin B (Sleeman et al. 2001). Leptomycin B is a compound that blocks snRNP export from the nucleus to the cytoplasm as well as its re-importation into the nucleus. This decrease in snRNP levels in the nucleus has been attributed to the inhibition of CB formation or to CB disorganization.

Lemm et al. (2006) have suggested the following model for the formation of Cajal bodies: (1) during the first stage, snRNP core molecules, which consist of Sm proteins and snRNA, interact with coilin in the nucleoplasm. (2) In the next stage, the self-oligomerization of coilin induces the joining of various factors that are related to snRNPs splicing, and (3) in the final stage of assembly, other domains such as snoRNP are also built, which finally results in the formation of CBs.

Recently, Strzelecka et al. (2010a) showed that coilin and snRNPs are concentrated in numerous CBs in zebrafish embryos before, during, and after activation of the zygotic genome. They suggested that CB number is regulated during development to respond to the demands of gene expression in a rapidly growing embryo. Following zygotic genome activation, snRNP biogenesis was required for CB assembly and maintenance, suggesting a self-assembly process that determines CB numbers in embryos.

In our embryological model, larch microsporocytes also exhibit natural fluctuations in levels of mRNA and rRNA metabolism (Smoliński et al. 2007; Smoliński and Kolowerzo 2011). In the larch, the synchronous development and “timely and fast switching on and off” of RNA and protein synthesis allows the tracking of changes that occur during snRNA synthesis and maturation snRNP cycle, and Cajal body formation has also been observed. Further study (Strzelecka et al. 2010b) showed that also coilin is required during vertebrate embryogenesis to maintain the supply of assembled spliceosomal snRNPs sufficient for expression of zygotic transcripts. Depletion of coilin in zebrafish embryos leads to CB dispersal, deficits in snRNP biogenesis and expression of spliced mRNA. snRNAs were necessary but not sufficient for rescue, showing that only assembled snRNPs can bypass the requirement for coilin. Thus, coilin’s essential function in embryos is to promote macromolecular assembly of snRNPs, likely by concentrating snRNP components in CBs to overcome rate-limiting assembly steps (Strzelecka et al. 2010b).

It seems that our results agree with the recently postulated hypothesis about the self-organization of compartments such as Cajal bodies, which is due to ability to selfdimerize such molecules like coilin, Sm proteins, and SMN (Misteli 2007). Kaiser et al. (2008) have artificially induced the de novo formation of Cajal bodies on chromatin after the immobilization of structural and functional components of the CB, such as coilin, SmD1, SmE and SmG proteins, and other factors related to snRNA and snoRNA. The authors have suggested that Cajal bodies originate as self-organizing structures, which are formed as local concentrations of macromolecules, but not through a “hierarchical assembly pathway”. Finally, tethering non-CB components to the lac operator array failed to nucleate CB formation, whereas tethering of PML body components resulted in formation of de novo PML bodies (Kaiser et al. 2008). Taken together, these data strongly support a stochastic assembly model and argue against an ordered or hierarchical nuclear body assembly pathway (Matera et al. 2009). Currently, it has been suggested that other nuclear compartments related to snRNA metabolism function in a similar way to CBs. Speckles, which represent another dynamic nuclear compartment, have their own aggregation-diffusion model of organization (Carrero et al. 2005). It cannot be excluded that also cytoplasmatic compartments related to snRNA metabolism as CsBs function in a similar way to nuclear compartments.

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