# Core spliceosomal Sm proteins as constituents of cytoplasmic mRNPs in plants.

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**Abstract:** In light of recent studies, many of the cytoplasmic posttranscriptional mRNA processing steps take place in highly specialized microdomains referred to as cytoplasmic bodies. These evolutionarily conserved microdomains are sites of regulation for both mRNA translation and degradation. It has been shown that in the larch microsporocyte cytoplasm, there is a significant pool of Sm proteins not related to snRNP complexes. These Sm proteins accumulate within distinct cytoplasmic bodies (S-bodies) that also contain mRNA. Sm proteins constitute an evolutionarily ancient family of small RNA-binding proteins. In eukaryotic cells, these molecules are involved in pre-mRNA splicing. The latest research indicates that in addition to this well-known function, Sm proteins could also have an impact on mRNA at subsequent stages of its life cycle. The aim of this work was to verify the hypothesis that canonical Sm proteins are part of the cytoplasmic mRNP complex and thus function in the posttranscriptional regulation of gene expression in plants.

# Introduction:

Posttranscriptional regulation of gene expression plays an essential role in all aspects of the cell life cycle, including cell development, differentiation, survival, homeostasis, adaptation to stress and response to environmental signals. Various mechanisms that enable efficient transcription, depending on actual need, have evolved in eukaryotic cells. Once mRNA is transcribed, it is bound by RNA-binding proteins (RBPs), key regulators of both mRNA stability and translation <sup>1</sup>. RBPs accompany mRNAs throughout their whole life cycle and guide the transcripts at every step of their maturation and turnover <sup>2,3</sup>. Due to the complexity and dynamic nature of mRNP complexes, determination of the full composition of a single mRNP complex remains a challenge. It has been shown that RNA binding proteins constitute 3 to 11% of bacterial and eukaryotic proteomes <sup>4</sup>.

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The cytoplasm plays the key role in the regulation of gene expression by controlling the level of mRNA translation and decay. The transport, localization and degradation of mRNA within this compartment are highly ordered processes. This complexity is not surprising considering the high content of mRNA in the cytoplasm (up to 150 000 molecules in mammalian cells <sup>5</sup>), where two major, counteracting processes of mRNA translation and decay must occur at the same time. As stated in recent literature, a vast majority of mRNA posttranscriptional regulation occurs within the specialized microdomains referred to as cytoplasmic bodies <sup>6,7</sup>. These nonmembrane bound, highly dynamic structures enriched in ribonucleoproteins are evolutionarily conserved, as their occurrence has been confirmed in yeasts <sup>7,8</sup>, protozoans <sup>9</sup>, nematodes <sup>10,11</sup>, insects <sup>12,13</sup>, amphibians <sup>14</sup>, mammals <sup>15,16</sup>, and plants <sup>17–20</sup>. Several cytoplasmic bodies have been described to date, including P-bodies, stress granules, neuronal granules, germinal granules and nuage, among others.

The core cytoplasmic bodies in eukaryotic cells are P-bodies, which are sites of mRNA regulation, including 5'–3' deadenylation-dependent degradation <sup>8,21</sup>, miRNA-mediated decay <sup>16</sup>, and mRNA stabilization and sequestration <sup>17</sup>, as well as nonsense-mediated decay (NMD) <sup>22</sup>, and translational regulation <sup>23–26</sup>. Due to their composition, including proteins involved in both mRNA stabilization and turnover, P-bodies constitute centers of mRNA sorting where transcripts are directed to either translation or degradation <sup>7</sup>. It has been revealed that the components of P-bodies accumulate transiently and can shuttle between the cytoplasm and other cytoplasmic granules <sup>6,24,25</sup>. Moreover, there is evidence that molecules such as GW182 protein, AGO2-miRNP and mRNA itself can be loaded into specific microvesicles, referred to as extracellular exosomes, and transferred between cells in fully functional form <sup>27</sup>. Thus, the mRNA life cycle involves constant dynamic changes in mRNP composition and subcellular localization, where mRNPs can shuttle between polysomes and cytoplasmic bodies, depending on external stimuli and the current metabolic needs of the cell. Despite growing research on the posttranscriptional regulation of mRNA utilization and its spatial organization within the cytoplasm, the full mechanisms and functional roles of mRNP assembly into higher order microdomains need to be elucidated.

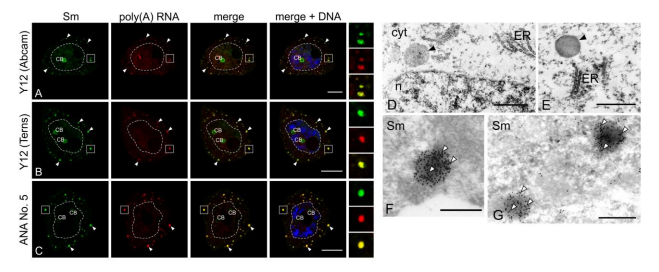
Here, we show that canonical core spliceosomal proteins (Sm proteins) constitute a part of the cytoplasmic mRNP complex in European larch (*Larix decidua* L.) microsporocytes. Sm proteins constitute an evolutionarily ancient family of small RNA-binding proteins. In eukaryotic cells, these molecules are primarily involved in pre-mRNA splicing. Only a few studies have indicated that, in addition to this well-known function, Sm proteins could also impact mRNA at subsequent stages of its life cycle. In our research, using larch microsporocytes as a model system, we show that there is a significant pool of Sm proteins accumulated within distinct cytoplasmic bodies that also contain polyadenylated RNA (poly(A) RNA). A correlation between the cyclic occurrence of

the cytoplasmic bodies and the metabolic activity of the cell was shown. A total of 222 mRNAs were identified as cytoplasmic partners for Sm proteins, and they comprise three major categories of Sm protein-associated mRNAs that code for proteins related to (i) ribosomes/translation, (ii) mitochondria/energy metabolism and (iii) chloroplasts/photosynthesis. Moreover, the distribution analysis of P-body markers (i.e., LSm4, DCP2, AGO1 and RS28) showed that Sm protein-mRNA cytoplasmic bodies are not functionally related to P-bodies. Based on the results obtained in this work and studies on other model eukaryotic cells, we propose that Sm protein-mRNA bodies constitute newly described cytoplasmic domains, as implicated in the posttranscriptional regulation of highly expressed transcripts, particularly in cells in which mRNA synthesis occurs in transcriptional bursts.

#### Results

# S-bodies: cytoplasmic bodies rich in the Sm proteins and polyadenylated mRNA

A double labeling assay of the core spliceosomal Sm proteins and poly(A) RNA revealed a remarkable distribution of these molecules within the cytoplasm (Fig. 1, Supplementary Fig. 1).





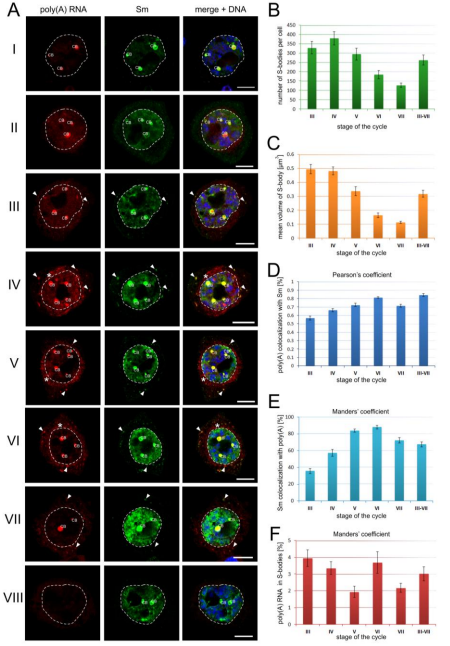
A- C, Localization of Sm proteins and poly(A) RNA in microsporocytes during diplotene. Visible numerous accumulation of Sm proteins in the colocalization of poly(A) RNA (arrowheads). The right panel represents the magnification of the cytoplasm, which is marked with a square. CB – Cajal body. Bar -15  $\mu$ m. D-E, Ultrastructural analysis of microsporocytes. Visible nonmembrane bound cytoplasmic structures (arrowheads).n - nucleus, cyt - cytoplasm, ER - endoplasmic reticulum. Bar - 1  $\mu$ m. F-G, Location of the Sm proteins in the cytoplasm of the microsporocytes as determined by the immunogold method. Visible accumulations of gold grains (20 nm) in isolated cytoplasmic clusters (arrowheads). Bar - 0.5  $\mu$ m.

Distinct  $0.5 - 1 \mu m$  cytoplasmic foci were observed, in which a significant amount of polyadenylated transcripts had accumulated, in colocalization with Sm proteins. To verify the specificity of the observed labeling, three different antibodies were used for Sm protein

localization, namely, commercially available Y12 MAbs (Fig. 1 A), Y12 MAbs from Matera's laboratory (Fig. 1 B) and the ANA No. 5 human reference anti-serum (Fig. 1 C). All the antibodies used showed a similar pattern of labeling with a well-established, characteristic nuclear pool of Sm proteins present in the nucleoplasm and Cajal bodies (CBs). Moreover, numerous discrete foci of Sm protein accumulation were observed in the cytoplasm and colocalized with poly(A) RNA (Supplementary Fig. 3). The ultrastructural analysis revealed spherical, nonmembrane bound microdomains with a diameter of  $0.4 - 1 \mu m$  (Fig. 1 D, E). The immunogold assay confirmed that a noticeable portion of the Sm proteins accumulated within distinct cytoplasmic structures (Fig. 1 F, G). Based on these observations and taking into consideration the general nomenclature used for this type of microdomains, we propose a name of S-bodies for these structures.

#### S-bodies are formed during the cytoplasmic export of poly(A) RNA

Larch microsporocytes exhibit a rigorously regulated pattern of metabolic activity. During the diplotene stage of meiosis, five bursts of *de novo* synthesis of poly(A) RNA occur, separated by periods of transcriptional silencing  $3^{2}$ . Each of these cellular poly(A) RNA turnover cycles comprises subsequent stages of nuclear synthesis, cytoplasmic export and degradation of mRNA (Fig. 2, Kołowerzo et al., unpublished). A detailed microscopic analysis showed that the S-bodies emerge periodically in each of the five cycles of poly(A) RNA turnover. Based on the transcription rate and localization of the poly(A) transcripts, the longest and most intensive fourth cycle could be divided into seven stages. Shortly after intensive synthesis in the nucleus (stages I and II, Fig. 2 A), poly(A) RNAs are transported to the cytoplasm, as indicated by an increasing level of labeling in this compartment (stages III through VII; Fig. 2 A). A remarkable ring-like accumulation of poly(A) RNA was observed around the nucleus, reflecting the pool of newly exported transcripts. Additionally, starting at stage III of the cycle, distinct foci of the poly(A) RNA and Sm proteins were observed and represented the S-bodies. At stage IV, the number of bodies was the highest in the whole cycle ( $380 \pm 35$  bodies per cell, Fig. 2 B). During the subsequent stages of cytoplasmic RNA localization, this number was consequently decreased (stages V through VII, Fig. 2 B). It should be emphasized, however, that throughout the whole period when S-bodies were observed, there was a significant colocalization of both poly(A) RNA and Sm proteins (PCC =  $0.69 \pm 0.02$ , Fig. 2 D) in the cytoplasmic pool. Furthermore, the Manders' overlap calculation demonstrated that a substantial fraction of Sm proteins in the cytoplasmic pool had colocalized with poly(A) RNA  $(67,43 \pm 2,90\%, Fig. 2 E).$ 



# Fig. 2: The longest and most intensive fourth cycle of poly(A) RNA turnover during diplotene stage of the first meiotic prophase.

A, The spatial and temporal distribution of the Sm proteins in the fourth cycle of poly(A) RNA turnover. I - VIII - successive stages of the cycles of poly(A) RNA turnover. A detailed description is in the text. CB - Cajal body. Bar -15  $\mu$ m. **B**, The mean number of S-bodies per cell in subsequent stages of the cycles of poly(A) RNA turnover. Error bars indicate the standard error of the mean (SEM). p <0.05. **C**, The mean volume of S-bodies in subsequent stages of the cycles of poly(A) RNA turnover. **D**-**E**, Analysis of the colocalization of poly(A) RNA and Sm proteins in the cytoplasm at subsequent stages of the poly(A) RNA cycle in the cell. The values are shown as the mean  $\pm$  standard error of the mean (SEM). PC - Pearson colocalization coefficient. **F**, The percentages were calculated on the basis of the Manders' colocation coefficient, in which the value of 1 was converted to 100% colocalization. Error bars indicate the standard error of the mean (SEM). a-b p <0.01, a-c and b-c p <0.001.

The overlap coefficient gradually increased as the cytoplasmic export of mRNAs proceeded: from  $35.76 \pm 3.09\%$  at stage III to  $88.03 \pm 2.13\%$  of the total cytoplasmic Sm protein pool at stage VI (Fig. 2 E). We also estimated the percentage of the cytoplasmic poly(A) RNA pool that was in S-bodies during the fourth cycle of transcriptional activity at the diplotene stage. The analysis revealed that  $3.08 \pm 0.42\%$  of the total cytoplasmic pool of poly(A) RNA had accumulated in cytoplasmic bodies, and this accumulation was consistent throughout the cytoplasmic phase of the RNA cell cycle (differences between stages were not statistically significant, ANOVA, p>0.05, Fig. 2 F).

To test the relationship between the occurrence of cytoplasmic bodies and the metabolic state of the cell, a quantitative analysis of the marker for transcriptional activity (active RNA pol II) was performed in the 3 cycles of the most intensive poly(A) RNA turnover (Fig. 3). During the stages of RNA cytoplasmic export (stages III-VII, see Fig. 2 A), i.e., when the Sm proteins and polyadenylated transcripts accumulate within cytoplasmic bodies, the transcriptional activity was considerably decreased compared to the transcriptional activity in the nuclear stages of mRNA synthesis (stages I-II, see Fig. 2 A; Fig. 3 A). In contrast, the level of the marker for potential translational activity (5S rRNA) showed no correlation (Fig. 3 B). These results indicate that the formation of S-bodies in the cytoplasm is slightly postponed with respect to bursts of nuclear mRNA synthesis and that these microdomains appear in periods of decreased transcriptional activity.

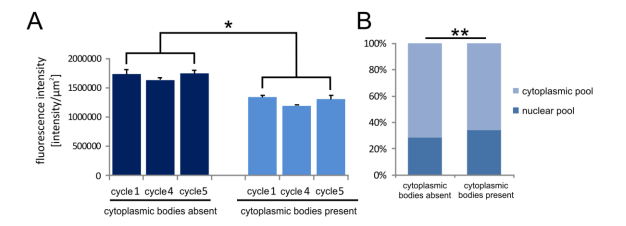


Fig. 3: Quantitative analysis of the marker for transcriptional activity.

**A**, Quantitative analysis of the active level of RNA polymerase II in the different cycles of poly(A) RNA synthesis in the presence and absence of Sm protein/poly(A) RNA bodies in the cytoplasm of the cell. The error bars indicate the standard error of the mean (SEM). \* p < 0.05. **B**, analysis of the nucleus:cytoplasm ratio of 5S rRNA in cells containing and not containing Sm protein/poly (A) RNA bodies. \*\* p < 0.01.

# S-bodies are not sites of spliceosomal particle accumulation

The well-known and broadly documented role of Sm proteins in the nucleus is their participation in pre-mRNA splicing. Sm proteins bind uridine-rich small nuclear RNAs (U snRNAs) in the form of a heptameric ring around the transcript, creating the core spliceosomal ribonucleoparticles U1, U2, U4 and U5 snRNPs . The assembly of U snRNP involves a highly dynamic cytoplasmic stage at which the Sm proteins are loaded onto the U snRNA transcripts and the 5' trimethyl-guanosine (m3G) cap is formed . Moreover, our previous research revealed that, during larch meiosis, the cytoplasmic stage of U snRNP assembly can occur in two spatially distinct manners: it may be diffused throughout the cytoplasm or localized within snRNP-rich cytoplasmic bodies (CsBs). The assembly mode depends on the rate of U snRNP *de novo* formation and the amount of U snRNP in the nucleus . Thus, it could not be precluded that the S-bodies are related to the assembly and/or sequestration of spliceosomal elements.

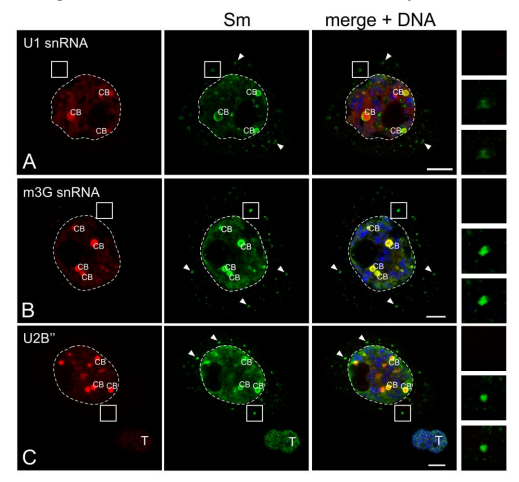
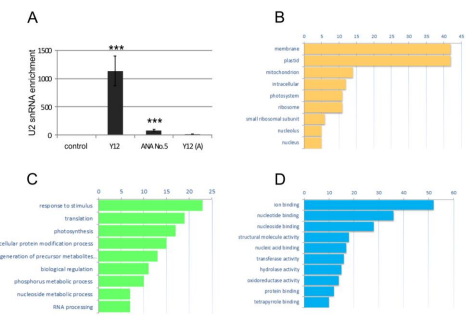


Fig. 4: S-bodies are not sites of the U snRNPs assembly.

Analysis of the Sm protein distribution in the colocalization with selected splicing elements. There was no accumulation of U2 snRNA (A), m3G cap (B) or U2B" proteins (C) in the S-bodies (arrowheads). The right panel represents the magnification of the cytoplasm, which is marked with a square. CB - Cajal body. T - anther tapetum cell. Scale of 10  $\mu$ m.

To test this possibility, the cellular distribution of both the nucleotide (U1, U2, U4, U5 snRNA, and m3G cap) and protein (U2B'') components of the U snRNPs were analyzed in relation to the S-bodies (Fig. 4, Supplementary Fig. 2). All the spliceosomal components were specifically localized within the nucleus, with significant accumulation in Cajal bodies (CBs). In contrast, no signal in the form of distinct clusters of U snRNA, m3G cap or U2B'' protein was observed in the cytoplasm, indicating that the formation of S-bodies does not coincide with the stage of U snRNP assembly in the cytoplasm. The lack of any U snRNP components and the presence Sm proteins in the S-bodies demonstrates that the bodies represent the cytoplasmic pool of canonical Sm proteins acting outside of the spliceosome, and that this localization is presumably related to mRNA.**Identification of Sm protein-associated cytoplasmic mRNA transcripts** 

To identify the mRNA transcripts that accumulate within S-bodies, we employed an RNAimmunoprecipitation (RIP) approach against the Sm proteins from the larch microsporocyte cytoplasm, followed by high-throughput sequencing of the immunopurified RNAs (RIP-seq).



# Fig. 5: Identification of mRNA transcripts

**A**, Determination of the efficiency of U2 snRNA immunoprecipitation using different anti-Sm protein antibodies (Y12, Ana No. 5, and Y12 (A)) by real-time (RT)-PCR. The graph shows the enrichment values relative to the control (for the control, the value was 1), normalized to the level of 5S rRNA. The error bars indicate the standard error of the mean (SEM). \*\*\* p <0.001. D-F The functional annotation of cytoplasmic mRNA precipitating with Sm proteins: biological process (**B**), molecular function (**C**), and cell compartment (**D**).

Since all three anti-Sm antibodies used for the assays *in situ* could label the cytoplasmic bodies with comparable specificity and strength (see Fig. 1), an additional validation of the results from the RNA immunoprecipitation analyses was performed to select the most efficient antibody to use for further analysis (Fig. 5 A).

The enrichment of U2 snRNA — a canonical cellular partner of the Sm proteins was estimated, and the results were normalized to 5S rRNA (which does not interact with Sm proteins). The experiment revealed that the Y12 MAbs obtained from Matera's laboratory was the most effective for the RIP assay, precipitating over 1130 times more U2 snRNA than the control (mock IP; no antibody added). This was significantly different than the 84-fold and 12-fold enrichment obtained from using ANA No. 5 and the commercially available Y12, respectively. The Sm protein-associated transcripts are listed in Supplementary Data 1. A total of 245 transcripts were identified as cytoplasmic partners of Sm proteins, of which 222 were classified as mRNAs. Most of the transcripts were homologous to *Pinaceae*. The Gene Ontology (GO) functional annotation revealed that the cytoplasmic Sm interactome consisted of mRNAs encoding a vast range of proteins, localizing to many cell compartments and linking a number of metabolic processes. A total of 141 sequences (58%) were annotated with 728 GO terms. This functional analysis revealed three major categories of Sm protein-associated mRNAs, and they coded for proteins related to (i) ribosomes/translation, (ii) mitochondria/energy metabolism and (iii) chloroplasts/photosynthesis (Fig. 5 B-D).

To test whether the immunoprecipitated mRNAs are indeed accumulated within S-bodies, the spatial distribution of the transcripts was analyzed *in situ*. From the 222 Sm protein-associated mRNAs 16 were selected, representing different cellular functions and localization. A double labeling assay of the larch microsporocytes was performed for the Sm proteins and each of the selected mRNAs (Fig. 6, Supplementary Fig. 3).

The microscopic analysis confirmed the specific locations of the 13 mRNAs within the cytoplasmic bodies. This specific distribution pattern was observed for mRNAs that encoded diverse proteins, including those involved in translation (Fig. 6 A, Supplementary Fig. 3 A, C, D), energy metabolism (Fig. 6 B, Supplementary Fig. 3 F) and photosynthesis (Fig. 6 C, Supplementary Fig. 3 I), as well as structural proteins of the cytoskeleton (Fig. 6 D, Supplementary Fig. 3 B). Additionally, S-bodies were enriched in several mRNAs related to protein folding (Supplementary Fig. 35 G, H), proteasomal degradation (Supplementary Fig. 3 J) and miRNA biogenesis (Supplementary Fig. 3 E). A different pattern of localization was observed for only 3 mRNAs (Supplementary Fig. 3 L) showed a diffuse pattern of cytoplasmic distribution, with no distinct concentration in any particular area.

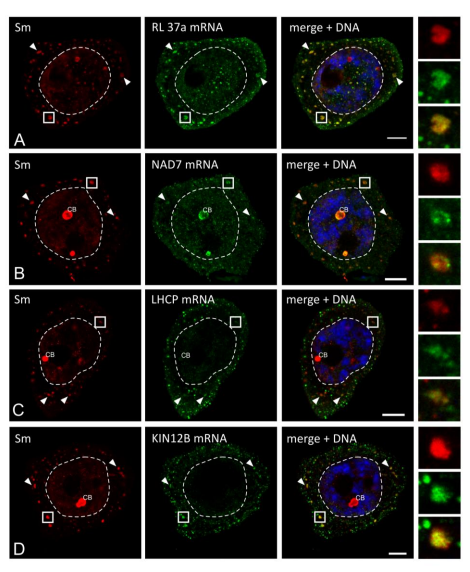


Fig. 6: In situ localization of mRNA precipitating with Sm proteins.

The results represent the stages of the IV - VI poly(A) RNA turnover. Numerous clusters of mRNA that colocalized with the Sm proteins (arrowheads) are visible in the cytoplasm. The right panel represents the magnification of the cytoplasm, which is marked with a square. **A**, RL 37a - large ribosome subunit protein 37a; **B**, NAD7 - NADH 7 dehydrogenase subunit; **C** - LHCP - protein binding chlorophyll a-b; **D**, KIN12b - kinesin 12b-like protein, CB - Cajal body. Bar -10  $\mu$ m.

Double labeling of these mRNAs with Sm proteins revealed that both of them were found in the cytoplasmic bodies, although without noticeable accumulation (i.e., there were no differences between the level of staining within the cytoplasmic bodies and in the surrounding cytoplasm). For the mRNA that encodes a light-independent protochlorophyllide oxidoreductase (LI-POR), the level of staining was below the limit of detection (Supplementary Fig. 3 M).

In the next step, we performed an additional analysis of the 14 mRNAs that were identified in the cytoplasmic transcriptome (unpublished data) of the larch microsporocytes, but not enriched in the Sm protein-immunoprecipitated pool of transcripts obtained via RIP-seq assay. Surprisingly, the

accumulation within the S-bodies was observed for 11 of the 14 mRNAs (Supplementary Fig. 4). The pattern of distribution was comparable to the RIP-seq identified transcripts. In addition to the mRNAs encoding proteins of the plastid (Supplementary Fig. 46 A) and mitochondria (Supplementary Fig. 4 E), those encoding nuclear proteins (Supplementary Fig. 4 F-H), as well as those related to Golgi apparatus (Supplementary Fig. 4 C) and ER (Supplementary Fig. 4 I) were enriched within the cytoplasmic bodies. Moreover, the S-bodies had also accumulated mRNAs encoding peroxidase (Supplementary Fig. 4B) and pectate lyase (Supplementary Fig. 4 K) – extracellular enzymes involved in cell wall metabolism and the response to environmental stress. The lack of cytoplasmic accumulation within these microdomains was observed solely in the case of mRNAs encoding SmD1, SmD2 and SmE proteins (Supplementary Fig. 46 L). This finding demonstrates that the S-bodies are not sites of *de novo* Sm protein synthesis.

#### The cytoplasmic Sm proteins interactome.

The next aim was to examine whether the S-bodies are functionally related to other microdomains, e.g., P-bodies and/or stress granules. To select potential markers for these structures, the cytoplasmic Sm protein interactome was identified. The cytoplasmic fraction of the larch anther cells was subjected to anti-Sm immunoprecipitation, which was subsequently assessed by mass spectrometry. A total of 118 proteins were detected (Supplementary Table 1, 2). Among the potential markers for the cytoplasmic bodies, LSm2, LSm3 and LSm4 were identified for P-bodies, and the ribosomal proteins eIF2/IF5, RL10e, RS1, RS3 and RS28 were characteristic for the stress granules (Supplementary Table 2).

Similar to the Sm protein RNA interactome, a significant fraction of the proteic interactome included proteins related to (i) ribosomes/translation (eIF2/IF5, RL10e, RS1, RS3, RS28, and OVA2), (ii) mitochondria/energy metabolism (SHM1 and SHM2 methyltransferases, IDH dehydrogenases, and ATM1 ABC transporters) and (iii) plastids/photosynthesis (RCA RuBisCO activase and TKL1 transketolase) (Supplementary Table 1, 2). Furthermore, two RNA-binding protein families were identified as coprecipitating with Sm proteins: Alba (*Acetylation Lowers Binding Affinity*) and ECT (*evolutionarily conserved C-terminal region*). Alba refers to a broad group of RNA- and DNA-binding proteins implicated in chromatin organization, regulation of transcription and translation, and RNA metabolism . It has been shown that these proteins are involved in the regulation of plant cell development where they perform essential roles in the sperm cell specification in *A. thaliana* . Fewer ECT proteins have been investigated compared to mRNA-binding proteins. The scarce ECT research indicates the role of ECT1 and ECT2 in calcium signal transduction from the cytoplasm to the nucleus that affects gene expression <sup>4</sup>.Based on the

IP-MS analysis and taking into consideration the availability of specific antibodies, three markers of protein degradation (LSm4, DCP2, and AGO1) and one translation regulator (RS28) were selected for the determination of *in situ* localization. All the proteins were distributed throughout the microsporocyte cytoplasm, with RS28 accumulated additionally in the nucleolus (Fig. 7). None of the investigated proteins colocalized with the S-bodies, indicating that these cytoplasmic bodies are not sites of mRNA degradation and are not involved in translational repression. Furthermore, no distinct foci for LSm4 (Fig. 7 A-D), DCP2 (Fig. 7 E-H), AGO1 (Fig. 7 I-N) and RS28 (Fig. 7 O-S) were detected in the cytoplasm, suggesting the lack of P-bodies or stress granules at this stage of meiosis in larch microsporocytes.

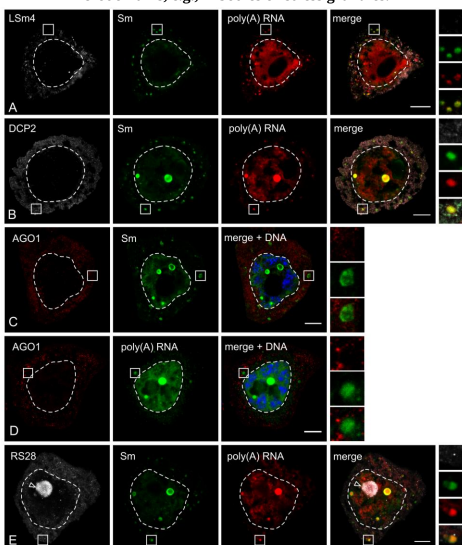


Fig. 7: S-bodies are not functionally related to other cytoplasmic microdomains, e.g., P-bodies or stress granules.

Localization of proteins related to degradation (A- D) and translation regulation (E) in *Larix decidua* microsporocytes. The results represent the stages of the IV - VI poly(A) RNA turnover. The right panel represents the magnification of the cytoplasm, which is marked with a square. Visible diffused fluorescence in the cytoplasm, without colocalization of the tested antigens for S-bodies and poly(A) RNA. Bar - 10  $\mu$ m.

#### Discussion

#### S-bodies — novel cytoplasmic mRNA domains in plant cells

Thus far, no distinct Sm proteins and mRNA accumulation has been observed in the cytoplasm of plant cells. However, a growing body of evidence shows a specific localization of Sm proteins and particular mRNAs in animal germ granules, which has implications for transcript localization, translational control and mRNA turnover. It was demonstrated that SmE and SmG, localizing to Pgranules in Caenorhabditis. elegans, are involved in transcriptional quiescence in germline precursors, and that this coordination of germline differentiation is splicing-independent <sup>10,4</sup>. Pgranules are structurally and functionally related to nuages in Xenopus oocytes, where the Sm proteins, but not other splicing elements, are localized. It was proposed that Sm proteins facilitate the transport of the mRNAs specific to germ granules (e.g. Xcat2) from the nucleus to the cytoplasm<sup>14</sup>. In mouse spermatocytes, Sm proteins accumulate in the RNP-rich chromatoid body <sup>4954</sup> and forms a complex with a marker protein of these domains – MTR1<sup>15</sup>. In *Drosophila* melanogaster, the accumulation of SmB and SmD3 at the posterior of the developing oocyte is directly linked to the unique localization and polarization of oskar mRNA. The proper localization of this transcript is crucial for germ cell specification during early embryogenesis <sup>5441,5452</sup>. Germline and early embryonic cells are highly metabolically active, with a spatiotemporally ordered pattern of expression for distinct groups of developmental genes. The European larch microsporocytes are male germline precursors that have been shown to be highly active periodically during diplotene. In this context, the S-bodies are similar to the germ granules observed in animals and might be considered, to some extent, as their plant counterparts.

#### Sm proteins as a part of the cytoplasmic mRNP complex

A growing body of evidence suggests additional roles for the canonical Sm proteins outside of the spliceosome in the processing, localization and translational control of mRNPs. However, little is known about which mRNPs in the pool of all transcripts are regulated by Sm proteins and what is the mechanism of their interaction. To determine which mRNAs are accumulated within the Sbodies, we performed a transcriptomic analysis of cytoplasmic Sm protein-mRNP complexes, followed by in situ detection analysis. The functional annotation showed that three major classes of Sm protein-associated mRNAs could be distinguished, namely, those encoding proteins related to (i) ribosomes/translation, (ii) mitochondria/energy metabolism and (iii) chloroplasts/photosynthesis. This finding is in agreement with the results obtained for Drosophila ovaries and HeLa cells. Lu et al. demonstrated that among 72 Sm protein-associated, fully spliced, polyadenylated mRNAs for Drosophila and 30 for HeLa cells, a significant portion encoded ribosomal and mitochondrial proteins. Some of the mRNAs show an overlap in all the Sm protein interactomes studied to date — plant, insect and mammalian, including those that encode small and large ribosomal subunit proteins, translation initiation factor eIF2B $\alpha$ , NADH dehydrogenase and cytochrome oxidases. Furthermore, Sm proteins were also associated with the mRNA products of intronless genes (e.g., histone H2A mRNAs in human cells , 60S ribosomal L37 mRNA in larch cells), proving that this interaction is completely independent of pre-mRNA splicing. However, the results from the animal cells do not clarify whether the Sm protein-associated mRNAs are localized to the nucleus or the cytoplasm, as the RIP-seq was performed on whole cell extracts. Nevertheless, together with the above-mentioned reports, our results indicate that the splicing-independent Sm proteins association with mRNA is conserved in invertebrates, vertebrates and plants.

Mutual posttranscriptional coordination of mRNA subpopulations has been previously described by Keene and Tenenbaum<sup>47</sup>. The following hypothesis was proposed based on eukaryotic posttranscriptional regulons (or operons): structurally and functionally related genes are regulated in groups by specific RNA-binding proteins to facilitate the organization of gene expression during cell growth and development <sup>354</sup>. The most acknowledged RBPs involved in this type of regulation are Pumilio proteins (PUF, Puf) that participate in the posttranscriptional silencing of particular groups of genes in yeasts, Drosophila, mammals and plants. The posttranscriptional regulons also operate in a number of developmental processes and stress responses, including the immunological response in mammals. In HeLa cells, the spliceosomal proteins U2AF (U2 auxiliary factor) and PTB (polypyrimidine tract binding protein) bind distinct groups of mature mRNAs in both the nucleus and the cytoplasm <sup>6</sup>. While U2AF interacts predominantly with transcriptional factors and cell cycle regulator mRNAs, PTB bound transcripts encode proteins related to intracellular and extracellular trafficking and apoptosis. Moreover, SR (Ser-Arg-rich) spliceosomal proteins are linked to cytoplasmic export of intronless histone mRNAs <sup>68,6</sup>, stimulation of translation and cytoplasmic degradation of particular transcripts. Recently, the involvement of SmD1 in posttranscriptional gene silencing (PTGS) was revealed in A. thaliana . Based on the results from a genetic screening of PTGS-defective mutants, it was proposed that, in addition to its role in pre-mRNA splicing, SmD1 facilities cytoplasmic PTGS. By protecting transgene-derived aberrant RNAs from degradation by the RNA quality control pathway (RQC) in the nucleus, SmD1 presumably promotes cytoplasmic export of transcripts to siRNA bodies, where PTGS occurs. Thus, the involvement of the evolutionarily conserved canonical Sm proteins in splicing-independent posttranscriptional control of mRNA metabolism is emerging as another regulatory component of gene expression. Taken together, these findings suggest that Sm proteins may represent a novel example of RNA-binding proteins that regulate functionally related groups

of mRNAs within the cytoplasm. In metabolically active microsporocytes (which enlarge six-fold during diplotene), the most abundant groups of transcripts are those related to efficient translation, energy production and photosynthesis, which aligns with the presented results.

It is worth noting that the S-bodies accumulated a number of mRNAs that were not enriched in the RIP-seq assay. Since RNA immunoprecipitation does not prove direct RNA-protein interaction and is highly dependent on details of technical procedures, it could not be ruled out that a portion of transiently and/or weakly associated mRNAs dissociated from the Sm complex during the experiment. Additionally, the cross-linking methods broadly used for the protein-RNA complex stabilization were previously shown to be inefficient for Sm proteins , as pointed out by Lu et al. . Therefore, it might be assumed that the Sm protein-mRNA interaction within a cell is a ubiquitous process, applying to a vast portion of transcripts, and that the number of mRNAs identified as cellular Sm partners remains underestimated.

# The mechanism of Sm protein-mRNA interaction

An open question concerns the mechanism of Sm proteins selectivity toward certain types of mRNAs and mechanism of their interaction. The only Sm proteins-binding elements described to date are the Sm-site present in U snRNAs and yeast telomerase RNA (TLC1 and TER1), and CAB box found in H/ACA Cajal-body specific RNAs (U85, U87, and U89 scaRNAs) and human telomerase RNA (hTR)<sup>7</sup>. Sm-sites or CAB box domain have not been found in mRNA to date. Furthermore, the specificity of scaRNA binding by Sm proteins is not fully understood. In both Drosophila and HeLa cells, Sm proteins co-precipitated only a fraction of the scaRNAs with no apparent structural features that could explain this selectivity. Several studies have demonstrated the participation of spliceosomal U1 snRNPs in splicing-independent posttranscriptional regulation of gene expression, including nascent mRNA protection from premature cleavage and polyadenylation at cryptic sites <sup>7987</sup>, regulation of polyadenylation of retroviral RNA as well as control of RNA pol II promoter directionality and promotion of sense mRNA transcription . Recently, it was revealed that, in plants, U1 snRNP regulates miRNA biogenesis and mediates the interactions between the microprocessor, spliceosome and polyadenylation machinery <sup>87,8</sup>. Moreover, it was shown that, in animal cells, the interaction between Sm proteins and mRNAs is mediated by U1 and presumably other snRNPs. Twelve-nucleotide putative binding sites for the 5' end of the U1 snRNPs were identified in Sm protein-associated mature mRNAs within the coding sequence far from intron-exon boundaries. It was thus proposed that U1 snRNPs bind mature mRNAs to promote the recruitment of RNA processing factors, thereby affecting mRNA localization, translation and/or turnover. In the case of the larch microsporocytes, the Sm protein-

mRNA interaction is apparently not mediated by U snRNPs, since there was no enrichment in U1 snRNA in the RIP-seq analysis. Additionally, the in situ localization of splicing elements demonstrated that the U snRNAs and m3G caps did not colocalize within the S-bodies, suggesting a different mechanism of Sm proteins-mRNA interaction in these cells. Studies on Drosophila and C. elegans have revealed that only some Sm proteins associate with mature mRNAs <sup>10541,8</sup>. Sm proteins constitute an evolutionarily conserved family and are classified into canonical (spliceosomal) Sm and noncanonical LSm (Sm-like) proteins. In Archaea, LSm proteins participate in tRNA maturation and ribosome synthesis . In Eubacteria, the Hfq homolog of the Sm protein regulates mRNA translation . The diversification of the ancestral Sm proteins in eukaryotes enabled their involvement in various processes. The LSm 2-8 ring is found in U6 and U6 atac snRNPs (spliceosomal subunits), as well as in the U8 snoRNPs involved in 5.8S and 28S rRNA processing . LSm 1-7 form a cytoplasmic complex that promotes mRNA decapping in P-bodies <sup>97,9</sup>. Another LSm ring variant, LSm 2-7, binds snR5 snoRNA, which is essential for rRNA pseudouridylation <sup>9</sup>. The unique LSm 10 and LSm 11 proteins are part of the U7 snRNP involved in the 3' end maturation of histone mRNA in animal cells. The Arabidopsis genome contains 42 Sm protein and LSM genes, many of which have not yet been characterized. With these findings considered, it could not be precluded that, in larch microsporocytes, the Sm proteins-mRNA association is driven by another noncanonical Sm/LSm ring or is indirectly mediated by other RBPs/RNPs. This hypothesis is supported by an IP-MS analysis that revealed a significant enrichment of the Alba proteins within the cytoplasmic Sm interactome. Alba proteins constitute an evolutionary ancient family of dimeric nucleic acid-binding proteins present in all domains of life . With their broad functional plasticity, Alba proteins are implicated in genome organization, transcription regulation, RNA metabolism, cell differentiation and stress response. Structurally, plant Alba paralogs display extensive diversity. Their association with target RNA is sequence-independent, and the formation of the Alba-RNA complex facilitates the accessibility of the transcript for RNA processing factors . It has been shown that the Alba interactome comprises a vast range of different RNAs and RBPs, including mRNA regulators such as PABP, ribosomal subunits and translational factors . Furthermore, the formation of a particular complex within a cell depends on Alba protein abundance and local concentration, the composition of the dimer (homo or hetero) and posttranslational modifications, including arginine methylation. Interestingly, several studies point to a cytoplasmic fraction of the Alba protein within the cell. In *Plasmodium falciparum*, PfAlba 1, 2 and 4 are localized as distinct cytoplasmic foci during proliferation <sup>10</sup>. Cytoplasmic localization of OsAlba1 was confirmed in rice (Oryza sativa), and its level of expression was increased in response to dehydration stress <sup>10</sup>. PAlba 1-3 from *P. berghei* coprecipitate with P granule

components PABP and eIF4E<sup>10</sup>. Additionally, the association of TbAlba 1-4 with cytoplasmic mRNAs was demonstrated in *Trypanosoma brucei*, in which it was localized to stress granules upon starvation, indicating their role in spatiotemporal translational regulation. Similarly, the larch microsporocyte cytoplasmic Sm protein interactome was enriched in both the Alba proteins and the ribosomal and translation-related proteins.

# Sm/poly(A) cytoplasmic bodies and the metabolic activity of the microsporocyte

The presented results unambiguously show that the S-bodies accumulate a large pool of mRNAs encoding different proteins. What would be the functional role of this kind of microdomain? The cytoplasm is a highly dynamic cell compartment where the antagonistic processes of mRNA translation and decay are strictly regulated <sup>10</sup>. The spatial separation of the mRNA processing, translational and degradation machinery in distinct microdomains facilitates the efficiency of posttranscriptional regulation and protects the transcripts from uncontrolled translation or degradation, depending on recent needs of the cell <sup>10</sup>. These studies demonstrate that the S-bodies are not sites of mRNA degradation and therefore are not analogs of P-bodies. This finding is in agreement with previous observations, showing that the mRNAs that accumulate within P-bodies are deadenylated <sup>8,10</sup>, which is not the case with S-bodies.

In many animal and plant species, the diplotene stage is characterized by significant chromatin relaxation, referred to as the diffuse stage of meiosis <sup>104–11</sup>. It has been demonstrated that in *Larix* species, the chromatin is subject to substantial rearrangements, and the microsporocyte is highly metabolically active, increasing its volume 6-fold during this stage. In Larix decidua, metabolic activity is strictly driven by cyclic chromatin contraction and diffusion. Five periods of chromatin relaxation (diffuse stages) separated by four periods of genome contraction are distinguished during diplotene. As a result, RNA synthesis is not a continuous process but occurs in transcriptional bursts — the diffuse stages are accompanied by large increases of *de novo* transcription of both mRNAs and rRNAs, as well as the expression and assembly of splicing subunits <sup>18325,11</sup>. The Sbodies were localized after transcriptional bursts, indicating that these microdomains are formed as a result of pulsed, extensive mRNA synthesis in the diffuse stage. The accumulation of a vast range of mRNAs in spatially separated cytoplasmic bodies is presumably a result of the effective organization of highly expressed transcripts during periods of greatly increased export of these molecules to the cytoplasm. Since S-bodies do not contain markers of degradation, the mRNAs are most likely sequestered to cope with the large number of molecules. This kind of regulation might act as a cellular strategy to control the accessibility of transcripts and their protein products during periods of genome contraction, at which time transcriptional regulation is not possible. This system

enables quick expression of proteins essential for normal cell growth and development (preparing the cell for division) and additionally provides the capability to respond to environmental changes and external stimuli. Recent studies have revealed that larch microsporocytes employ an unusual mechanism of nuclear mRNA retention with a putative role for Cajal bodies. It was demonstrated that a significant pool of newly transcribed poly(A) RNAs is temporarily sequestered in the nucleoplasm and CBs before being exported to the cytoplasm (Kołowerzo et al., unpublished). This functional connection between nuclear mRNA retention, accumulation in Cajal bodies and cytoplasmic localization is further supported by the observation that as many as 20 of the 29 mRNAs investigated as cytoplasmic partners for Sm proteins were also localized to the CBs (see Fig. 6, Supplementary Fig. 3, 4).

Conclusion: In summary, S-bodies represent newly described intracellular microdomains that participate in mRNA posttranscriptional regulation. Through the temporal storage of mRNAs and their spatial separation from degradation and translation factors, these domains presumably play a role in the coordination of the ongoing, frequently overlapping and competing processes of transcript maturation, translation and degradation. This strict control of mRNA fate is particularly essential in cells such as microsporocytes, in which the chromatin is periodically switched on and off and gene expression occurs in bursts. We propose a model in which the accumulation of significant amounts of different types of mRNA in cytoplasmic bodies provides a mechanism for the temporal retention and storage of highly expressed transcripts until they are needed for further translation or turnover within the cytoplasm, according to the needs of the cell. The results obtained in this work support the hypothesis that evolutionarily conserved Sm proteins have been adapted to perform a plethora of functions related to the posttranscriptional regulation of gene expression in Eukaryota. This multipurpose capacity presumably enabled them to coordinate the interdependent processes of splicing element assembly, mRNA maturation and processing, and mRNA translation regulation and degradation.

# Methods

# **Plant material**

For immunofluorescence/FISH assays, anthers of the European larch (*Larix decidua* Mill.) were collected between November and March at weekly intervals from the same tree in successive meiotic prophase (diplotene) stages to ensure consistent experimental conditions. The anthers were fixed in 4% paraformaldehyde in phosphate-buffered saline (PBS), pH 7.2, for 12 h and squashed to obtain free meiocytes. Meiotic protoplasts were isolated from these cells according to the

method of Kołowerzo et al. <sup>118</sup>. Isolated protoplasts were next subjected to immunostaining, FISH and immuno-FISH assays.

For transmission electron microscopy (TEM) assays, the anthers were fixed in 4% paraformaldehyde and 0.25% glutaraldehyde GA in 0.1 M PIPES, pH 7.2, for 12 h. The samples dedicated for the ultrastructural analysis were additionally fixed with 2% OsO<sub>4</sub> for 12 h at 4°C. Next, the samples were dehydrated in alcohol and embedded in LR Gold resin (Sigma, St. Louis, Mo., USA). The material was sectioned using a Leica Ultracut UCT ultramicrotome, and the ultrathin sections were placed on nickel-formvar-coated grids. For observations of the ultrastructure, the samples were contrasted with 2.5% uranyl acetate for 30 min and then incubated with 2.5% lead citrate for 15 min.

For immunoprecipitation (IP) and RNA immunoprecipitation (RIP), the larch anthers were flash-frozen in liquid nitrogen and stored at -80°C for further analysis.

# Immunogold labeling of Sm proteins

Ultrathin sections were preincubated with 1% BSA in PBS pH 7.2 for 30 min, followed by incubation with anti-Sm antibody (ANA No. 5, Centers for Disease Control and Prevention, Atlanta, GA 30333, USA) at 1:800 dilution and 4°C overnight. After washing with PBS ( $5 \times 10$  min), the samples were incubated with secondary anti-human antibody coupled to 20-nm-diameter colloidal gold particles (BioCell, Cardiff, UK; 1:20 dilution in 1% BSA in PBS, pH 7.2) for 1 h at 37°C, rinsed in PBS and water and then contrasted with 1% phosphotungstic acid (PTA) and 2.5% uranyl acetate. The ultrastructural and immunogold analyses were performed with the use of a JEOL EM 1010 transmission electron microscope.

# Design of the multi-labeling reactions

Prior to the assay, the cells were treated with 0.1% Triton X-100 in PBS for 25 min to induce cell membrane permeabilization. After labeling, the samples were stained for DNA detection with Hoechst (1:2000) and mounted in ProLong Gold Antifade reagent (Life Technologies, Carlsbad, CA, USA). The sequences for all the oligo probes used in this work are summarized in Supplementary Fig. 1.

# Double labeling of the poly(A) RNA/U snRNA/5S rRNA and Sm proteins

The samples were incubated with Cy3-labeled probes: oligo(dT), U1 snRNA, U2 snRNA, U4 snRNA, U5 snRNA and 5S rRNA. The 100  $\mu$ M stock solutions with the probes were diluted 1:500 in hybridization buffer (30% v/v formamide, 4× SSC, 5× Denhardt's buffer, 1 mM EDTA, and 50 mM phosphate buffer) and incubated for 4 h (oligo(dT)) or overnight (other probes) at 27°C. After

washing with  $4 \times SSC$  (5 × 1 min), 2× SSC (5 × 1 min) and PBS (1 × 3 min), the samples were treated with the following antibodies for the detection of the Sm proteins: human anti-Sm ANA No. 5 (1:300 in 0.2% acBSA in PBS, pH 7.2), mouse anti-Sm Y12 (Abcam; 1:100 in 0.01% acBSA in PBS, pH 7.2), mouse anti-Sm Y12 (a kind gift from Michael P. Terns, University of Georgia, Athens, GA 30602, USA; 1:500 in 0.01% acBSA in PBS, pH 7.2). The samples were then rinsed with PBS (5 × 3 min) and incubated for 1 h at 37°C with the following secondary antibodies: anti-human Alexa 488 (Thermo Fisher, Waltham, MA, USA; 1:750) or anti-mouse Alexa 488 (Thermo Fisher; 1:100 in 0.01% acBSA in PBS, pH 7.2).

# Double labeling of the mRNA and Sm proteins

Immunodetection of Sm proteins with ANA No. 5 was performed first to detect the mRNA and Sm proteins according to the procedure described above. The secondary anti-human TRITC antibody (Thermo Fisher; 1:200 in 0.01% acBSA in PBS, pH 7.2) was used in this assay. Next, the samples were incubated overnight at 27°C with DIG-labeled probes that are complementary to the distinct mRNAs (Supplementary Fig. 5) diluted 1:200 in hybridization buffer. After washing with 4× SSC ( $5 \times 1 \text{ min}$ ),  $2 \times SSC$  ( $5 \times 1 \text{ min}$ ),  $1 \times SSC$  ( $1 \times 10 \text{ min}$ ) and PBS ( $1 \times 3 \text{ min}$ ), the slides were incubated for 5 h at 8°C with mouse anti-DIG antibody (Roche, Basel, Basel, Switzerland; 1:200 in 0.05% acBSA in PBS, pH 7.2) and rinsed in PBS. The samples were then incubated for 1 h at 37°C with secondary anti-mouse Alexa 488 at 1:1000 in 0.01% acBSA in PBS, pH 7.2.

# **Double localization of the proteins**

Samples were incubated overnight at 4°C with the following primary antibodies: mouse anti-m3G (Santa Cruz Biotechnology, Dallas, TX; 1:50), mouse anti-U2B'' (LifeSpan Bioscience, Seattle, WA, USA; 1:20), rat anti-RNA pol II (Chromotek, Planegg-Martinsried, Germany; elongated form — hyperphosphorylated Ser2 of the CTD; 1:100) and rabbit anti-AGO1 (Agrisera, Vännäs, SWEDEN; 1:200) in 0.01% acBSA in PBS, pH 7.2. After rinsing with PBS, the samples were incubated for 1 h at 37°C with the following secondary antibodies: anti-mouse Alexa 546 (Thermo Fisher), anti-rat Alexa 488 (Thermo Fisher) or anti-rabbit Alexa 488 (Thermo Fisher) 1:500 in 0.01% acBSA in PBS, pH 7.2. Next, the slides were rinsed with PBS, preincubated with 2% BSA in PBS, pH 7.2, for 15 min, and incubated overnight at 4°C with the anti-Sm ANA No. 5 antibody diluted 1:300 in 0.2% acBSA in PBS, pH 7.2. After rinsing with PBS, the samples were incubated for 1 h at 37°C with the appropriate respective anti-human antibody: Alexa 488 at 1:750 or TRITC, at 1:200 in 0.01% acBSA in PBS, pH 7.2.

# Triple labeling of the poly(A) RNA and proteins

For the triple-labeling assays, the oligo(dT) probe was first hybridized according to the method described above (overnight incubation). Next, the samples were incubated overnight at 4°C with the following primary antibodies: mouse anti-DCP2 (Sigma, Saint Louis, MI, USA; 1:20), rabbit anti-LSm4 (Sigma; 1:100) and rabbit anti-RS28 (Sigma; 1:100) in 0.01% acBSA in PBS, pH 7.2. After washing with PBS, the slides were incubated for 1 h at 37°C with anti-mouse Alexa 488 (1:1000) or anti-rabbit Alexa 488 (1:500) in 0.01% acBSA in PBS, pH 7.2. The Sm proteins were labeled as described above with ANA No. 5 primary and anti-human Alexa 633 (Thermo Fisher; 1:200) secondary antibodies.

# Confocal microscopy and image analysis

The images were captured with a Leica TCS SP8 confocal microscope at wavelengths of 405, 488, 543, 561 and 633 nm. The optimized pinhole, long exposure time (400 kHz) and 63X (numerical aperture 1.4) Plan Apochromat DIC H oil immersion lens were used. Images were acquired sequentially in the blue (DAPI), green (Alexa 488), red (TRITC, Cy3, Alexa 546) and/or far red (Alexa 633) channels. The optical sections were collected at 0.5  $\mu$ m intervals. For the bleed-through analysis and control experiments, LAS AF software was used. For image processing and analysis, ImageJ software was used <sup>119</sup>.

For the quantitative measurements, each experiment was performed using consistent temperatures, incubation times, and concentrations of probes and antibodies. The images were collected under consistent conditions of acquisition (laser power, emission band, gain, and resolution) to ensure comparable results. Before quantification of the fluorescence intensity, the background was eliminated by adjusting the threshold according to autofluorescence based on the negative control. Between 10 and 30 cells were analyzed for each stage (experimental variant), depending on the experiment. The level of fluorescence was expressed in arbitrary units (as the mean intensity per  $\mu m^2$ ).

The statistical analysis was performed using SigmaPlot 11.0 software. To compare two groups, Student's t-tests were used. To compare between several groups, statistical significance was determined by the Kruskal-Wallis test. The significance level was set at p < 0.05 for all statistical tests.

Colocalization analysis was performed with the ImageJ plugin JACoP<sup>120</sup>. Between 10 and 25 cells were analyzed for each stage (experimental variant). Only the cytoplasmic area of the cells was subject to all colocalization analyses. The colocalization coefficients were calculated (PCC,

Pearson's correlation coefficient, and  $M_1$  and  $M_2$ , Manders' overlap coefficients) for each stage and evaluated for statistical significance with Costes' randomization test (p < 0.05).

The number and volume of the cytoplasmic bodies per cell were measured with ImageJ plugin 3D Objects Counter <sup>120</sup>. Between 10 and 25 cells for each stage (experimental variant) were analyzed. The statistical significance was determined by one-way ANOVA followed by multiple comparisons using the Holm-Sidak *post hoc* test.

#### Immunoprecipitation assays

For proteomic analysis of immunoprecipitates (IP-LC-MS/MS), contents from the cytoplasmic fraction of the cells were immunoprecipitated according to a modified protocol based on Oliva et al. <sup>121</sup>. Briefly, the larch anthers were ground in liquid nitrogen and homogenized in protein extraction buffer at 3 mL per 1 g of plant material. The homogenates were then centrifuged twice at 14 000  $\times$  g, followed by centrifugation at 16 000  $\times$  g for 30 min at 4°C to obtain native cytosolic protein extract. The supernatant (cytosolic fraction) was precleared with MagnaChiP A/G magnetic beads (Merck) for 1 h at 4°C and incubated with anti-Sm Y12 antibody (a kind gift from Michael P. Terns, University of Georgia, Athens, GA 30602, USA) overnight at 4°C. Next, a fresh set of magnetic beads was added, and the mixture was incubated for 1 h at 4°C. The beads were precipitated by immobilization on a magnetic separator (Thermo Fisher), washed four times with PBST (0.1% Tween 20) and resuspended in water. The experiment was performed in quadruplicate for the IP assay and triplicate for the mock-IP assay (no antibody added). The LC-MS/MS analysis was performed by the Mass Spectrometry Laboratory (IBB PAS, Warsaw) with the use of an Orbitrap mass spectrometer (Thermo Fisher). The proteins were identified with MASCOT software (http://www.matrixscience.com/) and the TAIR10 database. Next, the samples were analyzed with (http://proteom.ibb.waw.pl/mscan/index.html). list MScan software The of Sm proteinimmunoprecipitated proteins was corrected with the results from the mock-IP experiments. The protein was considered Sm protein-immunoprecipitated when it was present in at least 3 IP samples and not present in at least 2 control (mock-IP) samples.

For the RNA immunoprecipitation (RIP) from the cytoplasmic fraction, the assays were performed according to a modified protocol from Sorenson and Bailey-Serres <sup>122</sup>. Briefly, the larch anthers were ground in liquid nitrogen and RNP extraction buffer (200 mM Tris-HCl, pH 9.0; 110 mM potassium acetate; 0.5% (v/v) Triton X-100; 0.1% (v/v) Tween 20; 2.5 mM DTT; cOmplete protease inhibitor (Roche); and 0.04 U/µl RNase inhibitor) was added at 10 mL per 1.5 g of tissue. After thawing on ice, the material was filtered through Miracloth (Merck Millipore, Burlington, MA, USA; ø of 22-25 µm) and centrifuged at 1500 x g for 2 min at 4°C. The supernatant (10 mL per sample) containing the

cytoplasmic fraction (Supplementary Fig. 6) of the anther cells was precleared with MagnaChiP A/G magnetic beads for 1 h at 4°C and incubated with anti-Sm antibodies (ANA No. 5 — 15  $\mu$ l per sample, Y12 (Abcam) — 1  $\mu$ g per sample, Y12 (Terns) — 10  $\mu$ l per sample, mock-RIP — no antibody) for 2 h at 4°C. Next, a fresh set of magnetic beads was added and incubated for 1 h at 4°C. The beads were precipitated on a magnetic separator (Thermo Fisher), washed four times with wash buffer (200 mM Tris-HCl, pH 9.0; 110 mM potassium acetate; 0.5% (v/v) Triton X-100; 0.1% (v/v) Tween 20; and 2.5 mM DTT), resuspended in 100  $\mu$ l of cold wash buffer and subjected to RNA purification and cDNA library preparation.

#### cDNA library preparation and sequencing

RNA was purified with a TRIzol:chloroform extraction and incubated with TURBO DNA-free DNase (Thermo Fisher) according to the manufacturer's protocol. Next, the RNA samples were further purified by phenol:chloroform (Sigma; 25:24) extraction. The RNA pellet was resuspended in 20 µl of RNase-free water.

For qPCR, the RNA was reverse-transcribed with Superscript III (Invitrogen, Carlsbad, CA, USA) enzyme using random hexamers according to the manufacturer's protocol. The qPCR reaction mix contained 1  $\mu$ l of template cDNA, 5  $\mu$ l of 2× concentrated PowerSYBR Green PCR MasterMix (Thermo Fisher), 2 pmoles of each primer and water up to 10  $\mu$ l. The primers specific for U2 snRNA or 5S rRNA (Supl. 1) were used. The assays were carried out in a 7900HT Fast Real-Time PCR system (Applied Biosystems, Foster City, CA, USA), and the cycling conditions were as follows: 95°C for 10 min; 95°C for 15 s, and 40 cycles of 60°C for 1 min. The relative expression level of the U2 snRNA was calculated via the  $\Delta\Delta$ Ct method. The results were normalized to the 5S rRNA expression level and compared to the control (RNA isolated from the mock-RIP experiment), for which a value of 1 was assigned.

After qualitative (Bioanalyzer 2100; Agilent) and quantitative (NanoDrop 2000, Qubit; Thermo Fisher) evaluation, the RNA was used for cDNA library preparation with TruSeq Stranded Total RNA with a Ribo-Zero Plant kit (Illumina, San Diego, CA, USA) according to the manufacturer's protocol. Due to the extremely low RNA amount for the input, 30 cycles of library amplification were performed. The libraries were run on 1.5% agarose gels, and the 250-350 bp fragments were excised, eluted from the gel (GelElute gel extraction kit; Sigma) and quantified via qPCR with a KAPA Library Quantification Kit for Illumina (KAPA Biosystems) according to the manufacturer's protocol. The libraries were next sequenced on MiSeq (Illumina) with MiSeq Reagent Kit v3 (150 cycles). The reads were filtered in terms of quality via PRINSEQ-lite <sup>123</sup>. The sequence stretches with an average quality <20 over a window of 20 nt were removed, and the reads were cut immediately before the first incidence of a degenerated base. Furthermore, trimmed reads shorter than 36 nt were removed. The reads from all

the libraries were pooled and assembled *de novo* with Trinity v.2.8.2 at default settings <sup>124</sup>. The transcript sequences were next analyzed with BLASTX (for putative mRNAs) and BLASTN (for other RNAs) with the use of the NCBI nr and nt databases, respectively. Consensus functions and ontologies were found by LCA (least common ancestor) methodologies implemented in Blast2GO v.4.1 <sup>125</sup>.

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