Characterization of polarity development through 2- and 3-D imaging during the initial phase of microspore embryogenesis in *Brassica napus* L

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**Abstract** Isolated microspores of *B. napus* in culture change their developmental pathway from gametophytic to sporophytic and form embryo-like structures (ELS) upon prolonged heat shock treatment (5 days at 32 °C). ELS express polarity during the initial days of endosporic development. In this study, we focussed on the analysis of polarity development of ELS without suspensor. Fluorescence microscopy and 3-D confocal laser scanning microscopy (CLSM) without tissue interfering enabled us to get a good insight in the distribution of nuclei, mitochondria and endoplasmic reticulum (ER), the architecture of microtubular (MT) cytoskeleton and the places of 5-bromo-2′-deoxy-uridine (BrdU) incorporation in successive stages of microspore embryogenesis. Scanning electron microscopy (SEM) analysis revealed, for the first time, the appearance of a fibrillar extracellular matrix-like structure (ECM-like structure) in androgenic embryos without suspensor. Two types of endosporic development were distinguished based upon the initial location of the microspore nucleus. The polarity of dividing and growing cells was recognized by the differential distributions of organelles, by the organization of the MT cytoskeleton and by the visualization of DNA synthesis in the cell cycle. The directional location of nuclei, ER, mitochondria and starch grains in relation to the MTs configurations were early polarity indicators. Both exine rupture and ECM-like structure on the outer surfaces of ELS are supposed to stabilize ELS's morphological polarity. As the role of cell polarity during early endosporic microspore embryogenesis in apical–basal cell fate determination remains unclear, microspore culture system provides a powerful in vitro tool for studying the developmental processes that take place during the earliest stages of plant embryogenesis.

**Keywords** *Brassica napus* · Extracellular matrix-like structure · Microspore embryogenesis · Microtubular cytoskeleton · Nuclear DNA synthesis · Polarity

**Abbreviations**

| Acronym | Definition |
|---------|------------|
| BrdU | 5-Bromo-2′-deoxy-uridine |
| CPD | Critical point drying |
| CLSM | Confocal laser scanning microscopy |
| CMTs | Cortical microtubules |
| 2-D | Two-dimensional |
| 3-D | Three-dimensional |
| DIC | Differential interference contrast |
| DAPI | 4′,6-Diamidino-2-phenylindole × 2HCl |
| DiOC<sub>6</sub> | 3,3-Dihexylxocarbocyanine iodide |
| DNA | Deoxyribonucleic acid |
| DH | Doubled haploid |
Microspore embryogenesis, also called androgenesis, is an intriguing developmental process, in which male gametophytic cells are triggered to sporophytic development, resulting in the production of haploid/doubled haploid (DH) plants (Smýkal 2000; Wędzony et al. 2009). DHs are of high value in basic research as well as in commercial application in plant breeding (Forster et al. 2007; Wędzony et al. 2013). Since the 1980s, the *B. napus* microspore embryogenesis system has been intensively studied in many disciplines, on the physiological, biochemical, cytological and molecular levels. Numerous papers described endogenous and exogenous factors affecting the developmental fate of *B. napus* microspores in culture (Custers 2003; Ferrie 2003; Malik et al. 2008). A rather long heat treatment in the range of 32–40 °C effectively induces *B. napus* microspores or pollen to initiate androgenic development and embryo formation (Pechan and Keller 1988; Telmer et al. 1992; Binarova et al. 1997; Custers et al. 1994; Joosen et al. 2007; Supena et al. 2008; Dubas et al. 2011). During the induction period, changes in expression of genes and proteins (Cordewener et al. 1995, 2000; Boutilier et al. 2002; Malik et al. 2008), nuclear DNA synthesis (Binarova et al. 1993, 1997) and cytoskeletal aberrations (Hause et al. 1993; Simmonds 1994; Simmonds and Keller 1999) were observed.

*B. napus* androgenic embryos are commonly used as an alternative to the study of zygotic embryogenesis. Mild heat stress-induced microspore embryogenesis is the most powerful androgenic pathway to produce embryos with established polarity (Supena et al. 2008; Dubas et al. 2011). However, the procedure to obtain embryos mimicking zygotic embryos with suspensor in a big scale is limited to the DH 4079 line derived from *B. napus* Topas. For a wider range of plant species, only the long-term heat-induced ‘classical’ microspore embryogenesis could be applied, and it was used extensively (Zaki and Dickinson 1990, 1991; Binarova et al. 1993; Hause et al. 1993; Simmonds and Keller 1999; Custers 2003). Therefore, we investigated the classical system to collect more information about the induction of microspore embryogenesis under prolonged heat stress and focussed on the cytological aspects. We analyzed the process of androgenic embryo development in cases where structures resembling suspensors were absent. We took advantage of the easy access to huge numbers of embryos and the manipulation options using the ‘whole mount’ immunocytochemical technology for 3-D visualization of the MT cytoskeleton and spatial distribution of cell divisions.

The induction of microspore embryogenesis involves a set of profound changes in cell morphology (Custers 2003), in cell cycle (Binarova et al. 1993), in the pattern of cytokinesis (Gervais et al. 2000), cytoskeletal configurations (Simmonds and Keller 1999) and new cell wall formation (Simmonds and Keller 1999). Changes in the cytoskeleton as a cell polarity key factor altered polarity of dividing and growing cells (Wasteneys and Galway 2003). When the nucleus moved to the cell centre, the large vacuole fragmented, metabolic activity changed and the embryogenic development started (Hosp et al. 2007).

Ultrastructural, fluorescence and immunocytochemical studies have shown the significance of symmetrical nuclear division that coincided with rearrangements of the cytoskeleton upon heat stress (Zaki and Dickinson 1990, 1991; Hause et al. 1993; Binarova et al. 1993; Dubas et al. 2011) as compared to divisions and cytoskeletal organization observed in control microspores and pollen cultured at 18 °C (Hause et al. 1993; Dubas et al. 2012) or collected from anthers growing in planta (Hause et al. 1992). As heat-treated microspores prepared to embryogenesis exhibited changes in the cytoskeleton organization (Hause et al. 1993; Simmonds and Keller 1999; Dubas et al. 2011), we analysed the changes in microtubule (MTs) configurations. Although MTs can maintain cell polarity, endoplasmic reticulum (ER), mitochondria and starch grains were investigated in some developmental stages, too, and in addition, this paper focuses on the visualization of sites of nuclear DNA synthesis after the application of 5-bromo-2'-deoxy-uridine (BrdU).

As the extracellular matrix (ECM) is an important modulator of cell polarity and a potential structural marker for embryogenic competence acquisition (Berthiaume et al. 1996; Namasiyavam et al. 2006), we were interested in its presence in multicellular embryo-like structure (ELS) at the globular stage. For that reason, scanning electron microscopy (SEM) analysis was performed. We were interested if there is a link between the deposition of the ECM-like structure and stabilization of morphological polarity.

**Materials and methods**

Donor plants

Plants of *B. napus* L. cv. ‘Topas’ line DH 4079 were grown year-round under greenhouse conditions at 20/18 °C day/night.
temperature. Additional light (Philips SON-T lamps) was given from mid-September to the end of April, which also served to obtain a photoperiod of 16 h. At the beginning of bolting, the plants were transferred to a growth chamber and kept at 10 °C with 16 h illumination provided by 150 \( \mu \text{Em}^{-2} \text{s}^{-1} \) HPI (Philips) lamps. Plants were watered twice a week with N:P:K=15:15:18 soluble fertilizer. Inflorescences were harvested at least after 2 weeks of cold treatment, and flower buds of 3.1–3.3 mm in length were used for the microspore isolation.

Microspore isolation and culture

Microspores were isolated following the protocol of Custers (2003) with modifications by Joosen et al. (2007). Microspore suspensions were cultured in incubators (Marius Instruments or ST4/B/40; Industrial Research Institute for Automation and Measurement) at 32 °C for 5 days continuously, and then transferred to 25 °C in NLN-13 modified Lichter (1982) medium with 13 % sucrose (w/v), without potato extract and growth regulators in 6-cm Petri dishes. The culture resulted in the development of androgenic ELS.

Progress of microspore development in embryogenic culture was monitored daily from the isolation day to the tenth day of culture, when globular embryos were found. At least 40 microspores or microspore-derived embryos were studied each day of culture.

BrdU labelling

To analyze DNA synthesis and cell proliferation under embryogenic conditions, a pulse labelling with the thymidine substitute, 5-bromo-2′-deoxy-uridine (BrdU), was applied on the second day of culture. The BrdU labelling solution (RPN201, Amersham Biosciences) was added to the cultures at final dilution of 1:500. Cells were labelled for 24 h with BrdU, post-incubated in medium without BrdU during the next 24 h and then fixed. Morphological development of induced microspores was monitored in cultures without BrdU labelling.

All experiments were repeated at least four times.

Immunolocalization of microtubules and places of DNA synthesis

The whole-mount immunolocalization protocol with improved preservation of cell structures and improved penetration of antibodies was according to Friml et al. (2003) modified by Dubas et al. (2011, 2012).

Sample collection, prefixation, fixation, microspore and pollen wall permeabilization to antibodies, washing and incubation with proper antibodies were done manually on hand-made sieves prepared from ‘blue’ pipette tips (5 mm in diameter) with nylon mesh (11 \( \mu \text{m} \), Millipore). For immunostaining, the primary monoclonal antibodies [anti-\( \alpha \)-tubulin clone DM1A raised in mouse, dilution 1:1,000, Sigma T-9026, or anti-BrdU (Bromodeoxyuridine) monoclonal antibody, dilution 1:1000, Amersham Bioscience RP202] as well as the secondary antibody GaM/IgG/Alexa 488 (Molecular Probes A-11001, dilution 1:100) were applied. Control experiments were performed by omitting the first antibody. For DNA counterstaining, the samples were incubated in 0.1 % propidium iodide (PI, Sigma-Aldrich, P-4170) for 15 min, washed in PBS and embedded on slides in Citifluor-glycerol (Citifluor Ltd. in glycerol, AF2, Enfield Cloisters).

Visualization of cell components

DAPI (4′,6-diamidino-2-phenylindole×2HCl; Sigma-Aldrich, D-9564) was used to visualize chromatin in whole-mount preparations of the suspension culture according to Custers et al. (1994).

Cellulose was visualized with 0.01 % Calcofluor White ST (4,4′-bis [4-anilino-6-bis (2-hydroxyethyl) amino-s-trizin-2-ylamino]-2,2′-stilbenedisulfonic acid) (BioChemika, Fluka 18909) in water (absorption 365 nm, emission 420 nm) according to Benziman et al. (1980). Mitochondria and ER were shown with 0.0001 %, DiOC6 (3,3-dihexyloxacarbocyanine iodide; Molecular Probes; D-273; absorption 483 nm, emission 501 nm) in water (Terasaki 1994).

All the above-mentioned dyes and reagents, i.e. DAPI, PI, Calcofluor White ST or DiOC6, were added directly on the slide to an equal volume of fresh sample of suspension. Slides were kept in a humid chamber in darkness for at least 15 min before microscopic analysis.

Starch grains were stained with 0.0001 %, I/KI (Lugol's solution). To this end, 50 \( \mu \text{l} \) of I/KI solution was added directly to the same volume of fresh sample of microspore suspension culture. Staining was done on microscope slides kept in a humid chamber in darkness for about 15 min before analysis.

Microscopic observation

Confocal laser scanning microscopy (CLSM) and fluorescence microscopy were performed with a CELL MAP IC Bio-Rad (Microscience Division, Hemel Hempstead, UK) CLSM, mounted on a Nikon Eclipse TE 2000-S inverted microscope. Fluorescence was examined under filters EX 470-490/DM 510 BA/515 EF (Alexa 488) and EX 510-560/DM 580 BA/590 EF (PI). CLSM images were collected by a Kalman averaging of four full scans. Three-dimensional images and z-projections of the cells were obtained by collecting a series of approximately 30–60 optical sections in the z-axis, each section 0.5–1 \( \mu \text{m} \) thick. Images were acquired and processed using software programs, including
Lasersharp 2000 software (Microscience Division, Hemel Hempstead, UK), Confocal Assistant 4.02 (written by Todd Clark Brelje), CorelDRAW® ESSENTIALS 9.0 and Adobe Photoshop 5.0 (Adobe Systems, Mountain View, CA, USA).

Scanning electron microscopy

Material for SEM was prefixed in 5 % GA for 2 h at 4 °C. After rinsing with water, samples were post-fixed with 1 % OsO₄ in water for 2 h and rinsed again. Next to it, material was dehydrated through a graded ethanol series and dried with a CPD 020 (CO₂ critical point drying), coated with 3 nm platinum using an Oxford CT 1500 HF sputter coating unit and finally observed in a system JEOL JSM 6300 F scanning electron microscope.

Results

Morphogenesis after long heat shock treatment

When isolated microspores were cultured at 32 °C during 5 days and then at 25 °C, they underwent structural reorganization and developed to multicellular ELSs (Figs. 1, 2, 3 and 4).

When isolated from anthers, the majority (at least 60 %) of the microspores were at the late uninnucleate stage of development, each having a big central vacuole occupying most of the lumen of the cell, and the nucleus was located close to the exine opposite one of the three colpi (Fig. 1, a, insert). Mitochondria and ER gave strong fluorescence after DiOC₆ staining, particularly in the vicinity of the nucleus and in the peripheral cytoplasm (Fig. 1, a).

Based on the analysis of organelle distribution and configurations of the MT cytoskeleton within the cell, two pathways of embryogenesis were distinguished. Type I embryos (60 %), derived from microspores with their nucleus located close to the microspore cell wall (Fig. 1, a), and type II embryos (40 %), derived from microspores with a centrally located nucleus (Fig. 1, b, c, insert). Type II microspores, with ‘star-like’ internal morphology, had a big fragmented vacuole and cytoplasmic strands passing from the cell centre through the vacuole to the subcortical cytoplasm (Fig. 1, b1, b2, c, insert, Online Resource 1).

Mitochondria and ER gave strong fluorescence after DiOC₆ staining in cytoplasmic strands in both the vicinity of the nucleus and in the peripheral cytoplasm (Fig. 1, c). At that stage of development, a dense network of thick and long bundles of cortical MTs (CMTs) adjacent to the microspore cell wall was observed (Fig. 1, b1); some exhibited parallel arrangements, while other CMTs had no preferential orientation. Numerous endoplasmic MTs (EMTs) were closely associated with the nucleus (Fig. 1, b2).

The nucleus, derived from type I microspore, often remained close to the exine, when it entered S-phase (Fig. 1, d). Opposite in type II microspores, the S-phase-entered nucleus was located centrally which resulted in similar, central position of mitotic chromosomes (Fig. 1, e) and mitotic division did not depend on the initial

![Fig. 1](image-url) Microspores of Brassica napus induced to sporophytic development by prolonged heat stress treatment. b, e, f, h, k show fluorescent MTs; inserts in c and i, l show fluorescent DNA as a result of BrdU incorporation or labelling with DAPI. (a) Uni-nucleate microspore on the first day of culture. Mitochondria and ER, stained with DiOC₆ (green), are around the nucleus and in the vicinity of the exine. Insert on the right hand shows the nucleus (N) after PI labelling (red). (b1, b2, c) Type II microspore with its nucleus in the cell centre. Insert in b1 on the right hand shows the nucleolus (Nu) after PI labelling (red). (b1) Dense network of thick and long bundles of CMTs in parallel orientation close to the exine. (b2) Multiple EMTs closely associated with the nucleus (N). The same microspore as in b1. (c) Microspore with a big vacuole transected by cytoplasmic strands passing from the cell centre to the cortical cytoplasm. Mitochondria and ER, stained with DiOC₆ (green), are around the nucleus and in the vicinity of the exine. Insert on the right hand shows microspore with nucleus (N) in the cell centre. Microspore visualized in differential interference contrast (DIC)/UV light. (d) Type I microspore with its nucleus close to the exine. Microspore with green fluorescent nucleus indicating that it has entered S-phase. DNA synthesis is traced by BrdU incorporation and immunocytochemical labelling with Alexa 488 (green). Insert shows the nucleus (N) after PI labelling (red). (e) Dividing microspore with green fluorescent spindle MTs. Note that the orientation of the division plane turned 90° with respect to non-induced microspores. Insert shows the chromosomes in the equatorial plane after PI labelling. (f,g) 2-Nucleate endosporic structures derived from type I microspores on the second day of culture. (f) Structure with a network of densely packed cortical MTs without preferential orientation. Insert shows two symmetric nuclei (N) located close to exine after PI staining (red). (g1) Structure with DiOC₆-stained mitochondria and ER in the vicinity of nuclei and in the peripheral cytoplasm. (g2) Structure with symmetric nuclei (N) after PI staining. Insert on the right hand shows the structure in DIC. (h) Two-celled type II structure. (h) Structure with long MTs oriented parallel to each other and aligning the plasma membrane in the equatorial plane where the new cell wall will be formed. Insert shows the symmetric position of two spherical nuclei with well visible nuclei (Nu) after PI staining (red). (i) Structure with mitochondria and ER after DiOC₆ staining in the peripheral cytoplasm of both cells. Insert on the right hand shows symmetric nuclei (N). An arrow indicates faint fluorescence of cellulose cell wall after Calcofluor White staining. (j) Two-nucleate endosporic structure in which one nucleus entered S-phase. Visualization of DNA synthesis (in green) by BrdU incorporation and immunocytochemical labelling with Alexa 488. The second nucleus is not stained and located below; only the nucleoli are visible (arrows). Insert on the right hand shows the structure in DIC. (k1, k2) Three-celled structure with one dividing nucleus within the exine. (k1) Spindle MTs are only observed at the right hand side of the structure and demonstrates asynchronous division. (k2) The same structure as on k1 after PI staining. Note chromosomes (arrow) and nuclei (N) simultaneously. (l) 4-Nucleate structure still enclosed in the exine visualized in DIC/UV on the fourth day of culture. Insert on the right hand shows the mitochondria and ER after DiOC₆ staining in the vicinity of nuclei and in the peripheral cytoplasm close to the sporoderm. Bar=20 μm.
nuclear position. The orientation of the division plane turned 90° (Fig. 1, e), compared to the orientation that was observed in pollen cultured in vitro at 18 °C (Dubas et al. 2012) in the case of type I microspores. MTs formed a mitotic spindle with pole-to-pole MTs and chromosomes-to-pole MTs (Fig. 1, e, insert).

After the first mitosis, again the distribution of mitochondria and ER and the configuration of MTs differed between type I and type II embryos. In type I, two-nucleate embryos, mitochondria and ER gave strong fluorescence after DiOC₆ staining in the vicinity of nuclei and in the peripheral cytoplasm (Fig. 1, g1, g2, insert). Moreover, MTs were not observed at the equatorial plane of cell division, and cellulose cell wall was not detected either. In the sub-cortical layer of these structures, MTs formed a dense network without
Fig. 2 Morphogenesis of *Brassica napus* ELS. *b, f, i, j, m* show fluorescent MTs; *a, b insert, d, m2 and o* show fluorescent DNA as a result of BrdU incorporation, or labelling with DAPI or PI. *e, l, o* show fluorescent cellulose as a result of labelling with Calcofluor White. *h* shows starch grains as a result of labelling I/KI. *c, g, k, n* show fluorescent mitochondria and ER as a result of labelling with DiOC6.

(a) Microspore-derived structure with six labelled nuclei. Labelling in all cells might indicate synchronous S-phase. Visualization of DNA synthesis by BrdU incorporation and immunocytochemical labelling with Alexa 488 (green) indicating that all nuclei had incorporated BrdU as a result of DNA synthesis during the pulse labelling. (b) Structure with a dense network of CMTs (green) without preferential orientation, close to the enclosing exine. *Insert* shows six round nuclei after PI staining (red). (c) Multicellular type I embryo with DiOC6-stained mitochondria and ER in the peripheral cytoplasm of cells. (d) 8-Nucleate structure on the fourth day of culture. Structure still enclosed in the exine observed in UV after DAPI staining. (e) Multicellular ELS stained with Calcofluor White observed with UV showing cellulose cell walls (blue) of the outer cells. *Insert* shows the same structure which is partly released from the exine. (f) Globular ELS on the seventh day of culture. ELS with mitotic spindle (arrow), phragmoplast and CMTs close to the plasma membrane of the outer cells. *Insert* shows the place of exine attachment in DIC. (g) Globular ELS with DiOC6-stained mitochondria and ER in the peripheral cytoplasm of the outer cells. *Insert* shows the place of exine attachment in DIC. (h) ELS with starch grains after I/KI staining observed in DIC. Some cells accumulate starch grains at one pole. (*i–l*) Phragmoplasts and CMTs without preferential orientation in the outer cells of globular ELS, which released from the exine. (*j–l*) Globular ELSs at dermatogen stage with exine remnants at one pole. (*j*) Pattern of CMTs in protoderm cells. *Insert* shows the same structure in DIC. (*k*) Mitochondria and ER after DiOC6 staining in the peripheral cytoplasm of the outer cells. (*l*) Cellulose deposits in the outer cells after Calcofluor White labelling. *Insert* shows the same structure in DIC. (*m–o*) Multi-cellular ELS released from the exine showing two domains. (*m1*) Phragmoplasts and cortical MTs are only intensively labelled in the domain with the small cells. (*m2*) The small isodiametric cells exhibit intensively labelled nuclei after PI staining. The bigger cells show numerous small vacuoles and faint labelling of their nuclei. *Insert* shows the same structure in DIC. (*n*) The small isodiametric cells exhibit intensively labelled mitochondria and ER after DiOC6 staining. (*o*) The small isodiametric cells exhibit labelled cellulose deposits after Calcofluor White staining. *Bar* = 20 μm
preferential orientation (Fig. 1, f). Contrary, in two-celled embryos of type II, multiple long MTs in parallel arrays and parallel to the cell wall were found at the equatorial plane of cell division (Fig. 1, h).

Symmetric nuclear division produced two equally condensed daughter nuclei (Fig. 1, h, i insert) with a well visible nucleolus in each of them (Fig. 1, h insert). Arrays of MTs formed a ring-shaped phragmoplast configuration (Fig. 1, h and Online Resource 2). Mitochondria and ER gave strong fluorescence after DiOC₆ staining in the peripheral cytoplasm of both cells (Fig. 1, i). More divisions followed within a period of 3 days, resulting in three to four celled structures, of which the nuclei were still close to the original wall of the microspore (Fig. 1, k, l). In type I structures, mitochondria–ER gave strong fluorescence after DiOC₆ staining in the vicinity of the nuclei and in the peripheral cytoplasm close to the sporoderm (Fig. 1, l insert). Calcofluor White staining did not reveal cellulose deposits. Fluorescence caused by the incorporation of BrdU, Alexa 488, PI and DAPI labelling demonstrated the absence (Fig. 1, j) or presence (Fig. 2, a) of synchronization of cell cycle in both types of embryos. In dividing structures (types I and II), BrdU got incorporated into DNA, as a result of DNA synthesis during the pulse labelling of 24 h, when at least one of the ELS nuclei/cells was dividing. Alexa 488 marked MTs in mitotic spindles (types I and II) and phragmoplasts (only type II). The cortical surface cell layer of multicellular type I embryos was covered by a dense network of CMTs. These CMTs exhibited no preferential orientation (Fig. 2, b). Mitochondria and ER gave strong fluorescence after DiOC₆ staining in the peripheral cytoplasm of the cells (Fig. 2, c). As a consequence of proceeding through the cell divisions in both types of embryos, the number of nuclei (types I and II) increased. On day 4, there were structures with four to ten nuclei enclosed by the exine (Fig. 2, a–d).

Multicellular structures (derived from the types I and II embryos) emerged from the exine by day 5. In these structures, Calcofluor White stained the cellulosic component of the newly synthesized cell walls. At about day 7, they showed features of globular embryos, which developed into heart stage embryos without suspensors in 9–10-days-old cultures (Fig. 2, e, e insert).

Anticlinal and periclinal cell divisions of the outer cells of globular types I and II ELSs resulted in the development of protoderm, sub-epidermal cells and the enlargement of the embryo, which typically was devoid of a suspensor (Fig. 2, a–d).
the one pole. Two-domain ELSs released from the exine staining before day 6 of culture. Globular ELS with the exine remnant at embryogenesis, cellulose cell walls are not detected by Calcofluor White types I and II structures lead to multicellular ELS formation. In type I microspores with a centrally located nucleus. Symmetrical divisions in located close to the microspore cell wall. Type II embryos derive from pathways. Type I embryos derive from microspores with their nucleus

![Diagram](image)

**Fig. 4** Scheme of type I and type II B. napus microspore embryogenesis pathways. Type I embryos derive from microspores with their nucleus located close to the microspore cell wall. Type II embryos derive from microspores with a centrally located nucleus. Symmetrical divisions in types I and II structures lead to multicellular ELS formation. In type I embryogenesis, cellulose cell walls are not detected by Calcofluor White staining before day 6 of culture. Globular ELS with the exine remnant at the one pole. Two-domain ELSs released from the exine

e–l). There was again no overall net orientation of the CMTs during ELS development at the globular stage. Preprophase bands (PPBs), mitotic spindles, phragmoplasts and arrays of CMTs were regularly found in all cells of the ELSs (Fig. 2, f, i, j). The frequent occurrence of these forms of MT arrays in many globular embryos indicated that the cells of the globular ELS divided intensively already within the exine. The unruptured globular structures finally consisted of a mass of relatively small cells. These structures did not exhibit differentiation in shape. Therefore, they were analyzed for the expression of internal polarity by visualizing the distribution of mitochondria and ER with DiOC₆ and starch grains with I/KI. Mitochondria and ER gave strong fluorescence in the peripheral cytoplasm of all cells with cellulose deposits in the cell walls (Fig. 2, g, k–l). Cells that accumulated starch grains were only present at one pole of the ELS (Fig. 2, h). It has been observed at the stage of few cells in ELS that early exine rupturing resulted in the release of early ELS from the microspore cell wall, and those in ELS exhibited cell volume increase. The exine often remained attached to the structure at the only one pole, marking external polarity (Fig. 2, e–g, j–l). Once ELSs had released from the exine, they sometimes had bigger cells between a mass of small cells (Fig. 2, m1, m2). The MT cytoskeleton was visible in all small cells, whereas no MTs were observed in the big cells (Fig. 2, m1, m2, insert). The big cells had higher vacuolization and fainter fluorescence of DNA in nuclei after PI labelling (Fig. 2, m2, insert), fainter fluorescence of mitochondria and ER after DiOC₆ staining (Fig. 2, n) and fainter fluorescence of cellulose deposits (Fig. 2, o), while in the same cells, RNA in nucleoli resulted in intensive red fluorescence (Fig. 2, m2).

SEM analysis revealed that proembryos at the early globular stage show a compact fibrillar network on the surface of cells (Fig. 3, a) and between those cells forming net-like bridges (Fig. 3, b, b1 insert). Some of the fibrils were thicker, perhaps because they were linked to other fibrils forming bundles. This fibrillar network of ECM-like structure was less apparent in mature embryos with differentiated protoderm cell layer at the late globular stage and further (Fig. 3, c, d).

**Discussion**

Microspore embryogenesis in a conventional microspore culture of B. napus under prolonged heat stress (5 days at 32 °C and then at 25 °C) results in ELS formation without suspensor (Binarova et al. 1993; Hause et al. 1994; Custers et al. 1994; Cordewener et al. 1995).

Exposure of microspores to heat causes changes in the pathway of development from gametophytic to sporophytic. Sporophytic development in microspores is accompanied by swelling, autophagy of the cytoplasm, cytoskeletal and cytoplasmic rearrangement leading to changes in cell polarity, cell cycle, cell wall formation and gene expression (Zoriniants et al. 2005).

Male gametogenesis critically depends upon cell polarization, which contributes to a highly asymmetric cell division (PMI; first pollen mitosis). In microspores at the vacuolate stage, the nucleus is at the G1 phase and is located close to exine. Displacement continues throughout the mitotic cycle (Tanaka et al. 1979), and the polar position of the nucleus is maintained by the MT cytoskeleton (Tanaka et al. 1998; Dubas et al. 2012).

Similar to that, we have found that in some heat-induced microspores, the polar, eccentric position (type I) of the nucleus was maintained by the MT cytoskeleton. As a second result of our experiments, it was found that the nucleus remained located in the centre of the cell (type II), surrounded by cytoplasm strands in a ‘star-like’ morphology. Interestingly, microspores of rice (Raina and Irfan 1998), rapeseed (Zaki and Dickinson 1991), tobacco (Touraev et al. 1997), wheat (Indrianto et al. 2001) or triticale (Zur et al. 2008) also acquire a ‘star-like’ morphology. Stress might break a connection between the plasma membrane–actin–MT–nucleus complex and allow the nucleus to take a position at the centre of the cell (Pauls et al. 2006).

Because the majority of heat-treated microspores of B. napus exhibited nuclei remaining close to the exine (Hause et al. 1993), we suppose that in these microspores, the organization of MTs remained undisturbed. In such cases,
the position of the nucleus within the microspore expresses
the polar organization of the microspore from the beginning
of microspore embryogenesis.

We also observed that despite this difference in the ex-
pression of polarity, the nucleus entered S-phase, the mitotic
spindle changed orientation and symmetric division oc-
curred in both types of microspores (types I and II). Finally,
two daughter nuclei of similar size were observed within the
microspore parental wall (Zaki and Dickinson 1990; Telmer
et al. 1992, 1995). Successive mitotic divisions led to the
formation of multicellular or multinucleate structures from
types I and II microspores, respectively. However, to clarify
these observations, tracking of the entire process of embryo-
genesis from single selected microspores throughout the
whole culture period should be performed. In case of types
I and II microspores, embryogenesis proceeded in the well-
known, classical pathway (Zaki and Dickinson 1991; Hause
et al. 1993; Custers et al. 1994). Contrary to our results, Zaki
and Dickinson (1991) noticed that the majority of ELS
derived from the type II microspores.

Previous investigations in B. napus and triticate micro-
spore cultures indicated that changes in the MT organization
during the embryogenesis induction would lead to appear-
ance of a preprophase band (PPB), which might determine
division symmetry (Simmonds 1994; Simmonds and Keller
1999; Dubas et al. 2010, 2011).

We confirmed that microspores, induced to androgenesis,
underwent cell divisions with the change of division plane
(Simmonds 1994; Simmonds and Keller 1999), but were not
able to monitor PPB formation at this stage. Changes in the
MT cytoskeleton configuration clearly coincided with succes-
nive nuclear divisions, which led finally to multicellular ELSs
formation (Simmonds and Keller 1999; Dubas et al. 2010).

MTs and actin filaments, the two key components of the
plant cytoskeleton, play an essential role in cell wall forma-
tion and for cell-to-cell communication through plasmodes-
mata, which coordinate cellular behaviour (McLean et al.
1997). Staining with Calcofluor White revealed newly
formed cellulose cell walls, while immunodetection of MTs
revealed MTs close to the plasma membranes of multicellu-
lar ELS. Based on the observations of places of cellulose
formation and MTs occurrence, MTs close to the cell mem-
branes might function in cellulose microfibril deposition in
the new cell walls (Vesk et al. 1996; Kost et al. 2002). MTs
are known to predict the location of secondary wall deposi-
tion and influence the spatial arrangement of newly synthe-
sized cellulose microfibrils (Cyr 1994).

In type I embryos, cellulose deposition in cell walls could
not be detected after Calcofluor White staining. We assume
the primary cell wall of ELS's cells was poorly developed, or
in vitro conditions with the heat treatment affected cellulose
synthesis (Sato et al. 2001) or cellulose synthase upregulation
(Pauls et al. 2006). Apparently, multinucleate structures existed
in suspension culture for a certain period of time. However, the
question about the failure of deposition of cell wall matrix
polysaccharide components in the new developing cell plates
has no clear answer. It could be that the cellulose is not present
in ELSs cells, but the pectic middle lamella is present, and it
cements the cell walls of two adjoining cells together in ELS
before the bursting of the exine happens.

When the exine was broken, new cell wall formation was
initiated in ELS. Finally, multicellular structures at the globu-
lar stage were found in suspension.

We have noticed that polarity in multicellular structures
was expressed by the accumulation of starch grains in a
specific region, marking the future root pole. Similar finding
was reported previously by Hause et al. (1994) in Brassica.
In microspore-derived embryos of wheat, starch granules
accumulated in a zone opposite to the place of exine rapture
(Indrianto et al. 2001).

Our results show that the distribution of ER, mitochondria
and starch grains coincided with the specific arrangements of
the MT cytoskeleton, or could maintain cell polarity at some
stages of microspore embryogenesis. It is known that in
mammals ER, mitochondria exploit cytoskeletal elements
as tracks for their directional movements by using a special-
ized molecular machinery (Anesti and Scorrano 2006). In
plants, actin filaments function as such.

In addition, the polarity of ELSs was also expressed by
the size of cells, their vacuolization, MT assembly and in-
tensity of labelling of nuclei after PI staining. These facts
illustrate the presence of two types of ELS with two different
domains in suspension culture. Evidence for MT involve-
ment in cell polarity development and maintenance during
plant cell morphogenesis, especially in trichome formation,
is mounting (Smith and Oppenheimer 2005).

A similar phenomenon of the presence of two-domain
structures in maize microspore-derived embryos was ob-
served by Testillano et al. (2002). One domain contained
cells with one nucleus, a compact nucleolus, dense cyto-
plasm with abundant ribosomes, ER, plastids, mitochondria
and other organelles. The ultrastructural organization of
large cells was quite different having one or more big nuclei,
numerous and large vacuoles, few ribosomes and big plastids,
among them numerous amyloplasts. Both domains dif-
fered also in the intensity of anti-β tubulin signal. Two-
domain embryos attributed only to monocots microspore
embryogenesis so far.

The significance of the presence of a fibrillar ECM-like
structure on the surface of few-celled early proembryo that is
releasing from the exine is unclear. Is this structure really
associated with embryogenic competence and could it be an
early structural marker for polar growth of spring oilseed
rape microspore embryos? To test this hypothesis, additional
studies on the ultrastructural levels should be performed.
Physical state and local mechanical properties of the ECM

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were described in animal and plant multicellular organisms (Baluška et al. 2003). The ECM was considered as required to attain a polar shape. ECM was discovered in different types of in vitro cultures of different species, mainly in those where embryo formation was induced (Dubois et al. 1992; Šamaj et al. 1995; Namasiyam et al. 2006; Popielarska-Konieczna et al. 2010). So far, no ECM-like structure was found in direct androgenic cultures of any specimen. Moreover, for the first time, we show that the polarity of embryos was expressed by the appearance of ECM-like structure at the place where the exine was broken, playing probably a crucial role in cell adhesion, communication and integration during further embryo development.

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**Conflicts of interest** The authors declare that they have no conflict of interest.

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