Rape embryogenesis. VI. Formation of protein bodies

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Abstract

The storage protein synthesis starts in Brassica napus var. Górczański embryo at the final embryogenesis stage, i.e. in green seeds. Storage protein accumulate in selected zones adjacent to big vacuoles. These vacuoles, as well as surrounding protein zones, are subject to fragmentation. Young aleurone grains originate. They grow occupying sites of declining vacuoles. In mature rape embryo two kinds of protein bodies occur: aleuron grains, well-stainable with protein-specific dyes, and myrosin grains weakly-stainable with them but PAS-positive. Myrosin grains occur earlier than aleuron grains in special cortex and cytoleodn cells. Although the first aleuron grains form in outer cells of lateral cap parts and in cortex cells at the hypocotyl-root boundary, they originate most rapidly in endodermis. In the embryo axis aleuron grains form so rapidly that at the beginning of browning of the seed coat most of them are already formed. Aleuron grains developed in all embryo cells accept in those of the youngest columella layers and differentiated procambial strands. The accumulation of storage protein lasts till the end of seed maturation.

Key words: rape embryogenesis, storage material, protein bodies

INTRODUCTION

Protein bodies are commonly known storage reserves occuring mainly in seeds. In rape both lipid and protein bodies form storage material. Their distribution in mature embryo has been already studied by Kuraś (1984). He has presented their hystological diversity in single organs and within cells. The origin and transformation of lipid bodies in developing embryo have been presented in the previous paper from the "Rape embryogenesis" series
(Tykarska 1987). The formation of the first aggregates of storage protein and their transformation into aleuron grains with a large number of globoids in the rape embryogenesis have been studied by Kavetskaya (1960). Rest and Vaughan (1972) have distinguished two kinds of protein bodies in *Sinapis alba* (a species closely related to rape). These are: aleuron grains occurring in most embryo cells, and myrosin grains forming only in few myrosin idioblasts. The latter are homogeneous and develop earlier. They easily stain with dyes nonspecific to protein and are Schiff-positive (Goding et al. 1970). Myrosin cells are scattered over cytoledon parenchyma and embryo cortex (Guignard 1890). Iversen et al. (1979) have analysed the distribution of myrosin cells in germinating *Brassicaceae* seeds.

According to Crouch and Sussex (1981) main storage protein are not recognized in the first phase of rape embryo growth when cell divisions take place and embryo organs form. They occur in the second seed developmental phase and rapidly accumulate with the increase in embryo dry mass (Norton and Harris 1975, Crouch and Sussex 1981).

The present paper is a continuation of studies on starch (Tykarska 1982) and lipid (Tykarska 1987) accumulation in rape embryo. Also developmental changes in the embryo axis during formation of protein bodies have been taken into account. The studies are concerned with final embryogenesis stages, from green to black seeds.

**MATERIAL AND METHODS**

Cotyledon-less embryo axes or whole embryos were prepared out of rape (var. Górczański) seeds during last mitoses and at the successive maturation stages. They were fixed in 10% acrolein, 0.1-10 uranyl-acrolein and 0.25-0.5-10 CrAF (chrom-acet-formalin). Parafin sections were stained with proteinspecific dyes like: fast green in citrate-phosphate or phosphate (pH 3-8) buffers (Alfert and Geschwint 1953), brom-phenol blue (mercuric brom-phenol blue according to Mazia et al. 1953), acid fuchsin (Gerlach 1969) and by PAS method.

1 mm embryo axis apices were fixed in 2% glutaraldehyde and next in 1% osm and embedded in epon. Semi-thin sections were stained with toluidine blue.

**RESULTS**

**EMBRYOGENESIS TERMINATION**

Embryo is finally developed. It has large cotyledons, six-layered columella and leaf primordium. However, it is smaller than a mature one and loosely fits to the seed coat. Both embryo and seed coat are bright green. At this
stage the last mitoses take place in the embryo, while starch and lipid bodies rapidly accumulate (Tykarska 1982, 1987).

From the present studies it may be inferred that just in this time the formation of protein bodies occurs. First, myrosin grains form (Fig. 1). They are spherical, homogeneous, stainable by PAS method and slightly stained with fast green and brom-phenol blue. They originate in special cells of primary cortex and cotyledons. These cells are usually bigger than their surrounding cells. In hypocotyl the myrosin cells resemble their neighbours; in root they are oval (Fig. 15), scarce, usually 6-8 in a single central axis section.

When myrosin grains are visible, the cytoplasm of embryo cortex cells becomes slightly stainable with fast green, brom-phenol blue, and acid fuchsin. Soon, in the embryo axis two areas with strongly staining cells differentiate: cortex of lower hypocotyl and root top, as well as outer layers of lateral cap parts (Fig. 6). Storage protein accumulated in aleuron grains is responsible for the staining. At the same time in slightly stained cortex cells of the upper hypocotyl aleuron grains start to form. Thus, around large vacuoles slightly stained areas appear that are distinct from colourless vacuoles and cytoplasm (Fig. 2). The areas extend in time and their affinity to protein dyes increases.

**SEED MATURATION**

Early green seeds ("green a" Tykarska 1980)

Seeds are green and fairly hard. Although embryos are more massive than at the previous stage, they still loosely fit to the seed coat.

In embryo cortex cells storage protein still rapidly accumulates. The accumulation areas surround large vacuoles. In a densening substance light patches-forming globoids, are still better visible (Fig. 3). The both large vacuoles and neighbouring storage protein zones undergo fragmentation (Fig. 4). Young aleuron grains form (Fig. 5). They gradually fill up with protein and grow replacing now declining vacuoles.

Aleuron grains first originate in lateral cap parts, lower hypocotyl cortex at its boundary with root, as well as in endodermis (the quickest) and pericycle (Fig. 8). Later the process extends along and across nearly whole embryo axis. Figure 17 illustrates the directions of aleuron grain formation.

In cotyledons protein bodies originate later than in the embryo axis. At this stage they occur only in the outer cotyledon. In the inner cotyledon cytoplasm is so lightly stained that any changes are difficult to observe.

Protein bodies form quicker at the inner side of embryo curve.

Late green seeds ("green b" Tykarska 1980)

Green seeds are hard. Their colour becomes dimmer and gets brown. Massive embryo is tightly covered with seed coat and difficult to prepare
PLATE I

Formation of protein bodies. Fig. 1. Cortex fragment of upper hypocotyl from the embryo at the initial phase of protein formation, stained with fast green at pH 7.8 and by PAS, ×700. In myrosin cell strongly stained myrosin grains, elsewhere only starch is stained.

Fig. 2. Enlarged fragment of embryo cortex from Fig. 6 (in a frame) in the initial phase of aleuron grain formation: storage protein accumulation zones are distinct from the background of light vacuoles, ×1500. Fig. 3. Cortex cell with forming aleuron grains. In protein accumulation zones small, light patches, i.e. globoids, are visible, ×1250. Fig. 4. Simultaneous fragmentation of vacuoles and storage protein accumulation zones, ×1500. Fig. 5. Young aleuron grains, light vacuoles still visible, ×1250. Figs. 2-5 — stained with fast green.

PLATE II

Embryo axes at the formation of aleuron grains, fixed in 10% acrolein, ×80. Fig. 6. Termination of embryogenesis. Storage protein start to accumulate. In outer cap layers, and in periblem at hypocotyl-root boundary dark cells visible. Stained with fast green at pH 4 (framed fragment presented in Fig. 2). Fig. 7. Beginning of embryo maturation (green seed). Young aleuron grains are already present in dark-coloured periblem andplerom cells. Some very dark cortex cells, especially in subepidermis, are myrosin cells. Stained with fast green at pH 8.

PLATE III

Seed maturation. Fig. 8. Early green seed. Fragment of central cylinder and primary cortex from the central part of hypocotyl. Protein bodies visible in endodermis cells. Fixed in CrAF, stained with brom-phenol blue, ×1500. Fig. 9. Cortex and epidermis fragment from late green seeds, fixed in GA, stained by PAS. In myrosin cell large, dark-coloured myrosin grains. In the whole section starch grains are more strongly stained. In other cells slightly coloured aleuron grains are visible. Fig. 10. Root apex from late green seeds. Protein bodies already formed in outer columella layer. Fixed in 10% acrolein, stained with fast green at pH 7.4, ×500. Fig. 11. Brown seeds. Fragment of hypocotyl-root boundary. Large protein bodies in subepidermis, tiny — in epidermis. Fixed in 10% acrolein, stained with fast green at pH 4, ×1500.

PLATE IV

Protein bodies in black seeds. Fig. 12. Shoot apex. Aleuron grains in all meristematic cells. Fixed in 10% acrolein, stained with fast green at pH 3.6, ×700. Fig. 13. Procambium fragment in a strand entering in cotyledons. No protein bodies in metaxylem cells. Fixed in 0.1-10 AU, stained with fast green at pH 8, ×1500. Fig. 14. Epidermis and subepidermis fragment in upper hypocotyl. Fixed in UA, stained with acid fuchsin, ×1500. Fig. 15. Periblem fragment in upper root part. Myrosin cell surrounded by aleuron cells. Fixed in GA+OsO₄, ×1500. Fig. 16. Central cylinder and primary cortex in upper embryo part. Fixed in UA, stained with acid fuchsin, ×1500.

out. Its root is whitish, hypocotyl light green while cotyledons bright green.

Dense aleuron bodies with distinct globoids are almost in the whole embryo (Fig. 7). They are absent in the shoot apical meristem, inner cotyledon, some epidermis cells of lower hypocotyl, inner cells of lower central cylinder, outer cortex of root apex and in inner columella layers (Figs. 10, 18).
Fig. 17. Formation of aleuron grains in rape seeds. Dots indicate zones of first aleuron grain formation. Arrows show directions of aleuron grain formation in other embryo axis parts.

Fig. 18. Embryo axis from late green seeds. Dots indicate areas with already formed protein bodies.

From this stage in cortex sections cytoplasm stops to stain first at the outer side of embryo curve, while aleuron grains are well stainable there. Cytoplasm stains well merely in cells devoid of aleuron grains.

Green-brown seeds

Embryo axis whitish, cotyledons light green. Only shoot apical meristem, apical root part, inner columella layers, and epidermis at the hypocotyl-root boundary lack aleuron grains. Except those of columella, the cytoplasm of these cells is well stained.

Globoids are not evenly distributed inside aleuron grains. Aleuron grains in hypocotyl periblum contain most globoids, those of central cylinder and epidermis (Figs. 14, 16) — much less, whereas those of columella do not have them at all.
Brown seeds

Embryo axis whitish-creamy, outer cotyledon slightly green. In brown seed tiny aleuron grains form in epidermis at the hypocotyl-root boundary (Fig. 11). Upper, in epidermis cells they are much bigger (Fig. 14). Also in inner cotyledon aleuron grains develop.

Black seeds falling out of siliques

Embryo axis creamy-yellowish, cotyledons slightly lighter. The last aleuron grains with clearly visible globoids form in all shoot apex cells in most embryos (Fig. 12). Sometimes they may also develop in root apex promeristems. Small, homogeneous grains can be observed in cells of II-III layers of columella. However they are never visible in the youngest layers, as well as in procambium strands which enter into cotyledons (Fig. 13).

Three staining methods used i.e. with brom-phenol blue, with fast green at variable pH (3-8), and with acid fuchsin provide the same results. Sometimes, especially during formation of protein bodies, myrosin grains stain better than aleuron grains (Fig. 7). In material fixed in aldehydes and stained by PAS method aleuron grains are pinkish (Fig. 9).

DISCUSSION

Protein bodies occur when cell divisions terminate. Their formation takes place very fast. These results are comparable with those by Crouch and Sussex (1981). At the initial stages of embryogenesis most protein bodies are already formed. Only in shoot and root apices they are still absent.

The studies show that in rape, as in other Cruciferae, two kinds of protein bodies are formed: aleuron grains — in most embryo cells, and myrosin grains — in a few idioblasts. The latter form earlier, as in the case of Sinapis (Rest and Vaughan 1972).

Aleuron grains are well stainable with fast green in a wide pH range, with mercuric brom-phenol blue and with acid fuchsin. Myrosin grains stain weaker with these dyes but they have, as in the case of Brassica campestris (Stanley et al. 1976), positive PAS reaction, which means they contain protein and carbohydrates.

From Aschton’s (1976) studies it may be inferred that most storage proteins in plants (except in grasses) are globulins. Goding et al. (1970) have found that globulin 12S, glycoprotein with positive PAS reaction, is one of the main (50%) rape soluble proteins. If it is true, a question arises: why scarce myrosin grains, and not aleuron grains, are PAS-positive? The question has been asked by Stanley et al. (1976) in their studies on the microstructure of Brassica campestris seeds.
The studies of Crouch and Sussex (1981) indicate that glycoprotein 12S accumulates simultaneously with an increase in embryo dry mass till the seeds are mature. During this time the formation of still new aleuron grains is observed (the present studies). Pearse (1957) has stated that proteins stain by PAS method only when they contain more than 1% of carbohydrates. Thus it may be concluded that myrosin grains contain more carbohydrate components than glycoprotein 12S — in all probability a component of aleuron grains. However this problem requires further elucidation.

Protein bodies are formed differently in various plants. Many authors have found that protein accumulate in small vacuoles originated from a large vacuole by fragmentation (Dangeard 1921) — in Ricinus communis, Rest and Vaughan (1972) — in Sinapis alba, Adams et al. (1985) — in Glycine max). From the present studies it has been found that in rape hypocotyl cortex cells protein originate in special, close to central vacuole, zones. In densening storage protein globoids become visible. Shortly, still in green seeds, vacuoles with protein accumulation zones undergo fragmentation. Young aleuron grains form and grow while vacuoles decline. It might be possible that the areas with storage protein accumulation constitute protein vacuole that originated from endoplasmic reticulum. This requires further ultrastructural analyses.

In rape embryo some tendency in distribution of aleuron grains may be observed. First they occur in cap lateral parts and in lower hypocotyl cortex cells at its boundary with the root. Their formation is the quickest in endodermis. It is from these areas that the formation process of aleuron grains expands along, from the hypocotyl-root boundary to cotyledons and root apex, and across embryo axis, from endodermis inwards and outwards.

Hence the structure of endodermis at the hypocotyl-root boundary has to be considered. Maybe specific, meristematic structure of dermatogen cells at hypocotyl-root boundary constitutes a special zone through which nutrients flowing from outside enter (at the formation of first protein bodies there is still the residue of endosperm in an ovule). Moreover, rapid metabolic transformations may be stimulated by already green cotyledons.

From these series of studies it may be concluded that in the embryo axis two activity zones of storage material synthesis differentiate: primary cortex at the hypocotyl-root boundary here large starch grains occur the first and exist the longest; here also the first aleuron grains are formed and columnella (the biggest starch grains throughout embryo development exist and only a few lipid bodies and usually no, except outer layer, protein bodies are found). These are also the zones where rape embryo cell activation takes place during seed germination (Kuraś 1986).
REFERENCES:


Embriogeneza rzepaku. VI. Tworzenie ciał białkowych

Streszczenie

Syntéza białek zapasowych w zarodku Brassica napus var. Górckański rozpoczyna się w końcowym etapie embriogenezy, w zielonych nasionach. Białko zapasowe gromadzi się w wydzielonych strefach, przelegających do dużych wakuoli. Wakuole te i otaczające je
strefy białkowe ulegają fragmentacji. Powstają młode ziarna aleuronowe, które rosną, zajmując miejsce zanikających wakuoli. W dojrzałym zarodku rzepaku są dwa rodzaje ciał białkowych: ziarna aleuronowe, barwiące się intensywnie barwnikami specyficzными dla białek i zwykle słabo barwiące się nimi ziarna myrozynowe, dające pozytywną reakcję PAS. Ziarna myrozynowe tworzą się przed ziarnami aleuronowymi w nielicznych, specjalnych komórkach kory i liścieni. Pierwsze ziarna aleuronowe pojawiają się w zewnętrznych komórkach bocznych części czapeczki i w komórkach kory na pograniczu hypokotyla i korzenia, ale najszybciej formują się w endodermie. Ziarna aleuronowe w osi zarodka powstają bardzo szybko, tak, że w początkach brzuszewienia łupiny nasiennej większość ciał białkowych jest już wykształcona. Ziarna aleuronowe tworzą się we wszystkich komórkach zarodka, prócz najmłodszych warstw kolumelli i wyróżnionych pasm pramiażgi. Akumulacja białek zapasowych trwa do końca dojrzewania nasion.