

Purification and protein composition of oil bodies from *Brassica napus* seeds

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Abstract: Seed oil bodies are intracellular particles to store lipids as food reserves in oleaginous plants. Description of oil body-associated proteins of *Arabidopsis thaliana* has been recently reported whereas only few data are available in the case of rapeseed. Oil bodies have been prepared from two double-low varieties of *Brassica napus* seeds, a standard variety (Explus) and an oleic variety (Cabriolet). Oil bodies have been purified using floatation technique in the successive presence of high salt concentration, detergent or urea in order to remove non-specifically trapped proteins. The integrity of the oil bodies has been verified and their size estimated. Their protein and fatty acid contents have been determined. The proteins composing these organelles were extracted, separated by denaturing gel electrophoresis, digested by trypsin and their peptides were subsequently analyzed by liquid chromatography-tandem mass spectrometry. Protein identification was performed using *Arabidopsis thaliana* protein sequence database and a collection of Expressed Sequence Tag (EST) of *Brassica napus* generated from the framework of the French plant genomics programme "Genoplante". This led to the identification of a limited number of proteins: eight oleosins showing a high similarity each other and representing up to 75% of oil body proteins, a 11 β hydroxysteroid dehydrogenase-like protein highly homologous to the same protein from *A. thaliana*, and only few contaminating proteins associated with myrosinase activity.

Key words: *Brassica napus*, *Arabidopsis thaliana*, oil bodies, oleosins, protein composition

Introduction

Oleo proteaginous plants store lipids in oil bodies, composed of a core of triacylglycerols surrounded by a monolayer of phospholipids in which different proteins are inserted. Although the lipid body is a relative simple organelle from a structural point of view, the mechanisms of its biogenesis and degradation remain largely unknown [2]. The increase of rapeseed economical value relies in improving oil extraction while preserving the quality of protein by-products. In fact, efficient oil extraction from rapeseed is rather difficult, by comparison to other seeds (soybean, sunflower) despite similar oil content than sunflower. The process involves high temperature treatments and use of organic solvent. It is important to identify molecular and cellular factors involved in biogenesis of storage seed oil bodies to identify key factors for the stability of these organelles and to develop milder methods for extraction of rapeseed oil. Oil bodies must withstand extremes of desiccation, rehydration, heating and cooling for months before the storage oil can be mobilized following seed germination. Proteins embedded in the phospholipid membrane form an effective resistant surface for dormant and germinating seeds during the environmental extremes [3].

Description of oil body-associated proteins of *Arabidopsis thaliana* has been recently published [1]. The most abundant proteins found in *A. thaliana* are (i) four oleosins (S1 to S4) playing a structural role for stabilizing oil bodies, (ii) one caleosin with an hypothetic role in oil body maturation by calcium mediated fusion of microbodies, (iii) one sterol dehydrogenase. In the case of rape, only few data are available. An average diameter of 654 nm has been reported for oil bodies isolated from the mature seeds of *Brassica napus* [4]. Two, then three, then four oleosins have been described by Murphy's team showing a higher similarity each other [5-7]. Their transport from the site of synthesis on ribosomes via the endoplasmic reticulum prior to their accumulation on oil bodies has been also reported [8]. More recently, an oil body-associated caleosin isoform has been reported in rape seeds [9].

In this work, we describe the purification and characterization of oil bodies from two double-low varieties of rape seeds, a standard variety (Explus) and an oleic variety (Cabriolet). Their protein complement has been analyzed the most exhaustively as possible.

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Material and methods

Seeds

Mature seeds of *Brassica napus* (hybrid Explus and variety Cabriolet) were a generous gift of Monsanto (Saint-Louis, Missouri, USA). They were characterized by their fresh weight, dry weight and apparent volume estimated by counting the number of seeds filling the same volume of 15 mL.

Oil body purification and characterization

Purification

Oil bodies were purified as described by Tzen *et al.* [10] by floatation using six successive centrifugation steps (10,000 × *g*, 4 °C, 30 min) in a Kontron Ultracentrifuge equipped with a swinging-bucket rotor. In a typical oil body preparation, 300 mg of fresh weight of seeds were ground 3 times for 30 sec in 5 mL of 10 mM sodium phosphate buffer pH 7.5 containing 0.6 M sucrose (buffer 1) with a glass Potter and a teflon plunger driven by a Heidolph motor (rate 7). The sample was cooled on ice between each grinding cycle, and the potter was rinsed by 5 mL of buffer 1. The homogenate was overlaid by one volume of 10 mM sodium phosphate buffer pH 7.5 containing 0.4 M sucrose (buffer 2) and spun. The oil pad on top was collected and dispersed in 10 mL of 5 mM sodium phosphate buffer pH 7.5 containing 0.2 M sucrose and 0.1% (v/v) Tween 20. The emulsion was overlaid by one volume of 10 mM sodium phosphate buffer pH 7.5 and spun. The oil body fraction was resuspended in 10 mL of buffer 1 additionally containing 2 M sodium chloride, overlaid by one volume of 10 mM sodium phosphate buffer containing 0.25 M sucrose and 2 M sodium chloride and spun. The oil body fraction was resuspended in 5 mL of 7 M urea and left on a shaker (60 rpm) at 20 °C for 10 min. Then the suspension was placed in centrifuge tubes, overlaid by one volume of 10 mM sodium phosphate buffer pH 7.5 and spun. The oil body fraction was resuspended in 5 mL of buffer 1 and then mixed for 10 min at 20 °C with 5 mL of hexane. After centrifugation and removal of the upper hexane layer, the oil body fraction was collected and resuspended in 5 mL of buffer 1. In a last step, the resuspension was overlaid by one volume of buffer 2 and centrifuged. The oil body fraction on top was collected (OB), resuspended in a minimal volume of buffer 1 and stored at 4 °C till further use.

Microscopy

Nile red (Molecular Bioprobe, Montluçon, France, solution at 1 mg mL⁻¹ in acetone) was added to an aliquot of oil body suspension (1/10, v/v). After 1 h incubation at 20 °C, oil bodies were observed at 1000× focus through WIG or WB filters (for fluorescence) with an Olympus BX 51 light microscope equipped with 100 X oil immersion objective. Images were recorded using the Photometrics Cool SNAP software. Oil bodies were also dropped on a copper grid covered with a carbon film after negative staining using 3% phosphotungstic acid and observed by electronic microscopy (JEOL JEM-100S equipment operating at 80 kV under low illumination).

Light scattering

Oil bodies were diluted in water (generally 10 µL of purified fraction in 500 µL), and their hydrodynamic diameter was determined using a Malvern HPPS particle sizer. We used a lipid particle refraction index equal to 1.46, as determined by Michalski *et al.* [11] for milk fat globules. Experiments were performed at 20 °C in a quartz cuvette. Measurements were repeated 3 to 10 times. Standard latex particles (200 nm, Nanospheres, Duke Scientific, Palo Alto, USA) were dissolved in Milli Q grade ultrapure water (Millipore Corp, Molsheim, France) and used to check measurement accuracy.

Lipid extraction and quantification of seeds and oil bodies

Overall fatty acid composition and quantity were determined by gas chromatography. The method is based on the transmethylation of acylated and free fatty acids [12].

One seed (near 5 mg) was ground in 1 mL of 2.5% (v/v) sulfuric acid in methanol containing 100 µg of heptadecanoic acid. The sample was heated for 90 min at 80 °C making sure that no evaporation of solvent occurs. After the addition of 450 µL of hexane and 1.5 mL of water, fatty acid methyl esters were extracted into the organic phase by vigorous shaking. The tubes were centrifuged at low speed (500 *g* for 5 min) to help the two-phases separation. The organic upper phase was diluted (1/10) and 1 µL was analyzed by gas chromatography using a Girdel 30 chromatograph with a moving needle-type injector at 250 °C and a bonded silica capillary column with a stationary phase of 90% biscyanopropyl – 10% cyanopropyl siloxane (30 m × 0.25 mm I.D., 0.25 µm film thickness, SP-2380 from Supelco, Saint-Quentin Fallavier, France). The carrier gas was helium at an inlet pressure of 1 bar. The column temperature program started at 120 °C, ramping to 210 °C at 3 °C/min. The flame ionization detector was at 270 °C. Identification of fatty acid methyl ester (FAME) peaks was based upon retention times obtained for standards (Supelco). The total amount of fatty acids was calculated from the ratio between the sum of FAME peak areas and the heptadecanoic acid methyl ester peak area. Lipids from oil bodies fraction were extracted according to Folch *et al.* [13], after addition of heptadecanoic acid as internal standard and subjected to methanolysis.

Protein analysis

Electrophoresis

Proteins were quantitated with the Folin Ciocalteu reagent [14] using bovine serum albumin as standard. Protein aliquots from oil body fraction were precipitated with 3 volumes of cold acetone at -20 °C for 2 h or overnight. The pellet was dried and resuspended in a dissociation buffer consisting of 62.5 mM Tris-HCl (pH 6.8), 10% (v/v) glycerol, 5% (v/v) 2-mercaptoethanol, 2% (w/v) sodium dodecyl sulfate (SDS) and 0.02% (w/v) bromophenol blue. SDS-PAGE (polyacrylamide gel electrophoresis) of proteins was carried out according to Laemmli [15], using 12% ready to use Nu PAGE polyacrylamide gels (Novex, San Diego). Electrophoresis was run under 100 V for 180 min using 50 mM MES (2-[N-morpholino]ethane sulfonic acid) NuPAGE buffer (pH 7.3). Gel was stained with Coomassie blue (G-250) according to Neuhoff and Harold [16]. Molecular weights were estimated with Mark 12™ standard from Novex.

Gels were scanned (300 dpi) using an EPSON Perfection 1200 PHOTO scanner, and the TIFF resulting gels were analyzed using the Image Quant (version 4.2a) software (Molecular Dynamics).

Identification of oil bodies proteins

Protein bands stained with Coomassie blue were excised from the polyacrylamide gel and stored at -20 °C. Before trypsin digestion, gel slices were washed for 5 min with water, dehydrated for 15 min with acetonitrile and dried by vacuum centrifugation. Proteins were reduced for 30 min with 150 µL of 10 mM dithiothreitol – 0.1 M ammonium bicarbonate at 56 °C and alkylated in the dark with 100 µL of 55 mM iodoacetamide – 0.1 M ammonium bicarbonate for 20 min at 20 °C. The gel pieces were washed in 0.1 M ammonium bicarbonate, dehydrated with acetonitrile and vacuum dried. Then proteins were digested overnight at 37 °C with sequencing grade trypsin (EC 3.4.21.4, Roche Diagnostics, Meylan, France) at the concentration of 12.5 mg L⁻¹ in the presence of 25 mM ammonium bicarbonate and 5 mM calcium chloride. The resulting peptides were extracted successively with 5% formic acid (v/v), acetonitrile/water (50/50, v/v) and acetonitrile. Combined

extracts were dried and samples were dissolved in 1% formic acid before liquid chromatography-mass spectrometry analysis.

High performance liquid chromatography was carried out with a Spectra System equipment (Thermo Separation Products, Riviera Beach, USA) comprising a SCM1000 vacuum membrane degasser, P4000 gradient pumps and a manual injector. Volumes of 10 μL of samples were loaded onto a reversed-phase BioBasic-18 column (1 \times 150 mm, 300 \AA pore size, 5 μm film thickness, Thermo Electron Corporation). The column was eluted at a flow-rate of 0.1 mL min^{-1} at 20 $^{\circ}\text{C}$ with 5% of solvent B (acetonitrile + 0.1% formic acid) in A (water + 0.1% formic acid) for 2 min and then with a linear gradient of B in A from 5 to 45% over 40 min then 45 to 95% over 5 min before re-equilibration. Eluant from the column was introduced in the electrospray ionisation source of a Thermo Electron LCQ Deca ion-trap mass spectrometer operating in positive ion mode. Instrumental parameters were capillary temperature 280 $^{\circ}\text{C}$, capillary voltage 30 V, spray voltage 4.5 kV, sheath gas flow 80 a.u., auxiliary gas flow 5 a.u. Mass spectra were acquired scanning from m/z 200 to 2000. Ion fragmentation was carried out using a normalized collision energy of 35 arbitrary unit. Peptide ions were analyzed using the data-dependent "triple-play" method as follows: (i) full mass spectrum scan, (ii) ZoomScan (scan of the major ions with higher resolution to determine their charge), (iii) fragmentation of these ions. Protein identification was performed with Bioworks 3.1TM software using *Arabidopsis thaliana* and *Brassica napus* protein sequence databases extracted from non-redundant database downloaded from the National Center for Biotechnology Information (NCBI) FTP site. A collection of *Brassica napus* ESTs that has been generated from developing seeds (10, 15, 20, 25, 40, 45 days after pollination) and anthers in the framework of the French plant genomics programme "Genoplante" (<http://www.genoplante.com>) was also used during the progress of this work. The collection consists of 39 373 ESTs that have been arranged into 13 083 unique sequences after clustering and contigging [17]. The unigene set obtained after version 1 of contigging was used for similarity searches. No enzyme specificity was set for the query. The database-searching algorithm SequestTM uses a cross-correlation (Xcorr) and delta correlation (dCN) functions to assess the quality of the match between a tandem mass spectrum and amino acid sequence information in a database. The output data were evaluated in term of (i) trypsin nature of peptides, (ii) Xcorr magnitude up to 1.7, 2.2 and 3.3 for respectively mono-, di- and tri-charged peptides to minimize false positives, (iii) dCN higher than 0.1.

Results

Characterization of rape seeds and purified oil bodies

Explus and Cabriolet seeds were characterized only by little differences (table 1) in weight and size. On the contrary, water content of Cabriolet seeds was slightly higher. Oil bodies were purified from mature seeds and observed under fluorescence or electronic microscopy (figure 1). They proved to be constituted of spheres able to emit fluorescence light upon incubation in the presence of Nile red. The size of oil bodies was estimated using dynamic light scattering or electronic microscopy. Oil bodies from Cabriolet seeds appeared systematically smaller. Protein and fatty acid contents were measured showing a ratio fatty acid/total protein near 1 in the mature seeds and a little enrichment of oil bodies in fatty acid ($\times 2$ to 3). Protein and fat contents of seeds and oil bodies were slightly higher in the case of the standard Explus variety.

Fatty acid composition was determined in seeds and in purified oil bodies by gas chromatography of fatty acid methyl esters using heptadecanoic acid methyl ester as a standard (table 2). It was verified that the seeds of Cabriolet variety were enriched in oleic acid and diminished in linoleic

Table 1. Characteristics of rape seeds and purified oil bodies.

		Explus rape	Cabriolet rape
Seed	Fresh weight (mg)	5.03	5.10
	Estimated volume (mm^3)	7.9	8.9
	Water content (%)	8.7	12.0
	Protein content (mg/mg)	0.45 \pm 0.04	0.41 \pm 0.02
	Fat content (mg/mg)	0.43 \pm 0.05	0.39 \pm 0.02
	Oil body	Hydrodynamic diameter ^a (nm)	1095 \pm 212
	Microscopy diameter ^b (nm)	1328 \pm 404	812 \pm 322
	Protein content ($\mu\text{g}/\text{mg}$ seed)	38.8 \pm 3.5	33.8 \pm 7.4
	Fat content ($\mu\text{g}/\text{mg}$ seed)	100.3 \pm 2.3	95.7 \pm 2.5

^a hydrodynamic diameter was estimated using laser dynamic light scattering.

^b average size was estimated using electronic microscopy by measurements on near 60 oil bodies.

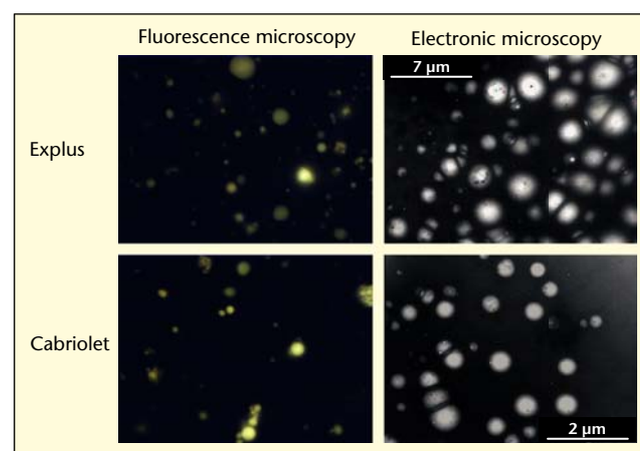


Figure 1. Oil bodies purified from rape seeds were inspected by fluorescence microscopy after staining with the lipophilic dye Nile Red or by electronic microscopy.

Table 2. Fatty acid composition of rape seeds and purified oil bodies.

Fatty acid	Explus		Cabriolet	
	Seed	Oil body	Seed	Oil body
C16:0	4.87	4.62	3.83	3.80
C16:1 (n-7)	0.29	0.20	0.18	0.18
C18:0	1.82	1.60	1.79	1.58
C18:1 (n-9)	61.47	61.82	73.48	73.87
C18:2 (n-6)	20.52	20.44	9.28	9.12
C18:3 (n-3)	9.61	9.79	9.91	9.97
C20:0	0.40	0.46	0.41	0.40
C20:1 (n-9)	1.02	1.07	1.12	1.08

Average relative error was near 1.8%.

acid in regards to standard Explus seeds. The same fatty acid composition was recovered in seeds and oil bodies.

Protein composition of purified oil bodies

The proteins contained in the oil body fraction obtained through the last step of preparation (see Material and Methods) were analyzed by denaturing polyacrylamide gel electrophoresis (figure 2). The two protein patterns were similar for Explus or Cabriolet seeds. Only six different protein bands were clearly visible within the 15-70 kDa range after Coomassie blue staining. Upon scanning of gels and image analysis, an

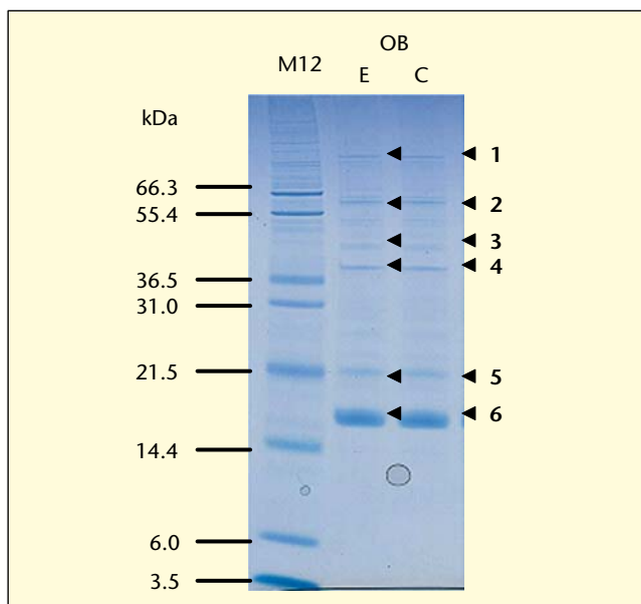


Figure 2. SDS-PAGE of proteins from oil body fraction (OB) purified from Explus (E, 10 μ g) or Cabriolet (C, 10 μ g) seeds. Molecular mass marker was Mark 12 (M12) from Novex. Protein bands were numbered as in table 3.

approximate quantification on the basis of band intensity was obtained (table 3). The protein bands were identified through the analysis of their trypsin peptides with liquid chromatography-tandem mass spectrometry. As *Brassica napus* genome was not completely sequenced and accessible as public databases, they were identified in part through homology with *Arabidopsis thaliana* proteins. All the proteins were identified without ambiguity and with a high coverage of the protein sequence. The major proteins present in rape oil bodies were very homologous to integral proteins previously described in *A. thaliana* oil bodies [1]. Interrogation of ESTs database from *B. napus* immature seeds (Genoplante cDNA library) gave the possibility to determine the real protein sequences in *B. napus*. Eight oleosins were identified in rape oil bodies. One could be distinguished from the others due to its slightly higher apparent molecular mass of 22 kDa (band 5). This oleosin (contig accession number 155968 in the Genoplante database) corresponded only to a partial protein sequence which presented 61% identity and

67% similarity with the N-terminal part of oleosin S4 from *A. thaliana* and possessed 17 supplementary amino acids. Protein band 6 showing an apparent molecular mass of 17 kDa was the major band and appeared to contain 7 oleosins the molecular mass of which was comprised between 19.3 and 21.5 kDa. Four oleosins (BnIII, BnV, napII and contig 155846) presented 82.5 to 86.7% identity with *A. thaliana* oleosin S3, one oleosin (contig 156118) presented 80.8% identity with *A. thaliana* oleosin S1 and two oleosins (contig 156532 and contig 156533) presented 77.1 to 80.3% identity with *A. thaliana* oleosin S2.

Besides oleosins, a protein highly similar to the 11- β -hydroxysteroid dehydrogenase-like protein described in *A. thaliana* was identified. However, only a partial sequence was recovered from ESTs database (235 amino acids against 349 amino acids for *A. thaliana* protein). This partial sequence corresponding to the N-terminal part of the protein presented 89.8% identity and 96.2% similarity with *A. thaliana* protein.

Finally, the presence of myrosinase-binding protein, myrosinase-associated MyAP5 and β -glucosidase indicated a low contamination of oil body fraction with protein body. However, no storage protein (cruciferins) was identified.

Discussion

The oil body preparation used repeated and numerous floatation steps. Proteins non-specifically associated with oil bodies were removed by detergent washing, ionic elution and finally urea treatment. Triglycerides from defective oil bodies were removed by hexane extraction. So, only few proteins were visible from oil body fraction and the contamination with proteins involved in the glucosinolate metabolism was relatively low. Seeds and oil body fraction had relatively high protein contents compared to other results. Murphy and Cummins [3] have reported a ratio fatty acid/total protein near 5.7 for oil bodies but they have noticed that total protein was assayed by three different methods and that protein recoveries were relatively low and variable. In fact, they have carried out protein determination on delipidated oil bodies and we have verified in that case a poor protein recovery.

Eight oleosins were identified in purified oil bodies from rape. These oleosins show a high similarity with *A. thaliana* oleosins. However, oppositely to *A. thaliana* oleosins, they were not resolved under SDS-PAGE due to their very similar molecular mass, except for rape oleosin S4. Rape oleosins were always heavier than the *A. thaliana* corresponding ones except for oleosin S2 the molecular mass of which was similar for *B. napus* and *A. thaliana*. Only four oleosins were described in rape by

Table 3. Identified proteins in oil bodies purified from *Brassica napus* mature seeds.

Band	Relative intensity (%)	Identified protein	Apparent molecular weight (kDa)	Sequence molecular weight (kDa)	Sequence coverage (%)	pI	<i>A. thaliana</i> orthologues
1	2.6	Myrosinase-binding protein	88	99.404	6.8	5.5	
2	6.4	β -glucosidase	62	56.283	5.8	6.2	At3g03640
3	4.4	Myrosinase-associated MyAP5	46	41.797	31.3	8.5	
4	9.4	11- β -hydroxysteroid dehydrogenase-like	40	40	22.1	6.1	At5g50600
5	10.7	contig 155968	22	23.024	28.6	9.1	At5g40420/S4
6	64.0	BnV	17	20.286	49.7	9.2	At4g25140/S3
		BnIII		21.540	19.0	9.3	At4g25140/S3
		NapII		19.349	53.7	9.6	At4g25140/S3
		contig 155846		19.515	47.2	9.2	At4g25140/S3
		contig 156118		20.777	23.8	8.1	At3g01570/S1
		contig 156532		20.002	46.3	9.1	At3g27670/S2
		contig 156533		19.880	25.5	6.9	At3g27670/S2

Murphy's team [5, 18-20]. BnV, BnII and napII were recovered in our experiments and it appeared that EST 156533 corresponded to napI described only partially by Murphy [18]. Then we observed four supplementary oleosins. *Brassica napus* is a natural hybrid between *B. oleracea* and *B. campestris* and in consequence, it is not surprising that *B. napus* contains many more oleosins than *A. thaliana* the genome of which is the simplest in the cruciferae family.

Oleosins contain three distinct structural domains, a central hydrophobic anchoring domain, highly conserved and containing no cleavage site for trypsin digestion, and two N- and C- terminal amphipathic domains. Recovered peptides through liquid chromatography-tandem mass spectrometry analysis belonged to the two amphipathic domains of the oleosins. The N-terminal part of *B. napus* oleosin S4 was enriched in glycine: 20 glycine residues, among 64 amino acids for N-terminal part, instead of 11 glycine residues, among 47 amino acids for N-terminal part in *A. thaliana* oleosin S4. This fact increases the hydrophobic character of N-terminal domain of rape oleosin S4.

A protein highly homologous to sterol dehydrogenase from *A. thaliana* was identified. It is also highly similar to *Sesamum indicum* sterol dehydrogenase sop2 (65% identity and 83% similarity) and at a minor degree with sop3 (46% identity and 71% similarity) [21]. This protein was never described in rape. It will be interesting to characterize the kinetic properties of *B. napus* sterol dehydrogenase, determine its specificity using labeled steroids as model substrates and classify this protein among the sterol dehydrogenases family.

Oleosins, the major proteins of oil bodies (75%), were poor candidates for bidimensional (2D) electrophoresis due to their very alkaline pI and their low solubility in aqueous medium. However, oleosin corresponding to contig 156533 has a lower pI. So, it will be interesting to attempt to separate oleosins using 2D electrophoresis. To date, no caleosin was identified in our experiments using proteomic technique. However, ESTs database reveals the presence of one gene encoding at least one protein very homologous to *A. thaliana* caleosin. The use of specific antibody could be considered to verify this hypothesis.

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