Storage Sites in Seeds of *Caesalpinia echinata* and *C. ferrea* (Leguminosae) with Considerations on Nutrients Flow

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**ABSTRACT**

The seeds of *Caesalpinia echinata* and *C. ferrea* behaved as typical endospermic seeds, despite their different morphological classification (exendospermic seeds were described for *C. echinata* and endospermic seeds for *C. ferrea*). Then, the aim of this work was to compare, under ultrastructural and histochemical terms, the nature of the storage substances and their accumulation sites, as well as the nutrient flow in seeds of these species. Cotyledons in *C. echinata* accumulate carbohydrates, lipids and proteins, which are mobilized from the outer to the inner parts as revealed by the position of plasmodesmata. Endosperm in *C. ferrea* accumulates carbohydrates and in *C. echinata* accumulates substances during the initial embryogenic phases. Such tissue develops a chalazal haustorium that is responsible for the transport of substances into the endosperm itself and from it into the embryo, confirmed by the presence of transference cells.

**Key words**: Cotyledon, embryo, endosperm, histochemistry, reserve substances, ultrastructure

**INTRODUCTION**

The accumulation of storage in seeds is one of the most important post-fertilization events. Growth, embryo differentiation and seed germination depend upon the breakdown of lipids, carbohydrates and proteins. Higher the nutrient supplies in the seed, better will be the chances for the seedlings to get established more vigorously (Kozlowski and Gunn, 1972). Lipids occur as oily bodies in the endosperm and embryo. Carbohydrates are stored either as starch grains or hemicellulosic wall thickening, which are common in the endosperm and cotyledons. Proteins occur as proteic bodies in the embryo and endosperm and build a matrix differentiated by the presence of certain minerals (Bhatnagar and Sawhney, 1981; Boesewinkel and Bouman, 1984). Other tissues, such as seed coat, can stack storage substances in the mature seed as well as during the first embryogenic phases (Boesewinkel and Bouman, 1984). Seeds can reveal growth substances such as auxines, cytokines and gibberellins, besides carbohydrates, lipids and proteins, which are stored in the endosperm. This shows that embryonary growth and differentiation are controlled by such tissue (Vijayaraghavan and Prabhakar, 1984). In most plants, endosperm persists until seed maturation and becomes a storage tissue (endospermic or albuminous seeds), but it can be completely absorbed throughout the embryogenesis (exendospermic or exalbuminous seeds).
seeds) (Johri et al., 1992). In exendospermic seeds, both embryo and endosperm development happens at the same time, and embryo growth occurs because of the endosperm that absorbs nutrients from the embryo sac. Storage substances of the mature seed are present in the cotyledons that have their photosynthetic activity transferred to other tissues. In endospermic seeds, embryo growth is limited and begins long after endosperm grows. Endosperm is not thoroughly consumed and is responsible for storage until germination; cotyledons, which are foliaceous in this case, will synthesize nutrients after they emerge from the soil (Kozlowski and Gunn, 1972; Boesewinkel and Bouman, 1984; Johri et al., 1992; Richards, 1997). The genus *Caesalpinia* s. l. (Leguminosae, Caesalpinioideae) presents species that show both seed kinds. Therefore, it was chosen for the present study. In a former study on the ontogenesis of the seeds of *C. echinata* Lam. (brazilwood) and *C. ferrea* Mart. ex Tul. var. *ferrea* (brazilian ironwood), embryos of *C. echinata* (exendospermic seeds) presented a resting phase, even with a complete absorption of the endosperm during the embryogenesis, similarly to what happened with *C. ferrea* (endospermic seeds), i.e., the seeds of both species behave as endospermic seeds (Teixeira et al., 2004). Such observations brought other comparative questions into consideration and discussed in the present article: Which kind of storage substances can be found in seeds? Which are the tissues that really present a storage function in seeds? How does the nutrient flow happen inside seeds? These questions are considered here under both histochemical and ultrastructural points of view and aim to contribute with the knowledge on the germination of seeds and establishment of seedlings. Moreover, according to Borges et al. (2005), the analysis of storage substances in embryo could contribute to understand the process of seedlings development from seeds at different stages of maturation. Our data also intend to favor the conservation of species under extinction risk and found in highly advanced degradation ecosystems, such as the brasil wood tree of the Atlantic Rainforest.

**MATERIAL AND METHODS**

**Studied species**

*Caesalpinia echinata* and *C. ferrea* var. *ferrea* are two Brazilian tree species that occur preferentially in the northeastern Atlantic rainforest (Polhill and Vidal, 1981). *C. echinata* has been almost exterminated as a result of its use as a source of red dye for fabrics and ink and the current total size of natural stands of this species is low (Cardoso et al., 1998). *C. ferrea*, a species used in Brazilian folk medicine, has been investigated for its pharmacological properties (Carvalho et al., 1996; Nakamura et al., 2002a, b).

**Plant material**

Samples of *C. echinata* were collected during the spring of 2001 from six plants cultivated in Campinas (22°54’S - 47°06’W) and on the Fazenda Campininha, Moji-Guáçu (22°18’S - 47°11’W), São Paulo state, Brazil. The *C. ferrea* samples were collected from five plants cultivated in Campinas (22°54’S - 47°06’W) and Botucatu (22°52’S – 48°30’W), São Paulo state, Brazil, during the summer and autumn of 2002. Voucher specimens were deposited in the herbarium of the Universidade Estadual de Campinas (UEC), São Paulo state, Brazil, under the accession numbers 26368 and 51731 for *C. echinata* and 60661 and 66698 for *C. ferrea*.

Seeds were taken from *C. echinata* fruits (66.5 mm and 80.0 mm long) and *C. ferrea* fruits (55.0 mm and 75.0 mm long). At least five seeds were collected at each stage of maturity per individual per species. Seeds inside shorter fruits presented embryos that started to protrude cotyledons (immature seeds) and those ones inside longer fruits presented embryos with completely expanded cotyledons (mature seeds), according to Teixeira et al. (2004).

**Ultrastructure**

For ultrastructural studies, small pieces of immature and mature seeds were fixed immediately after harvest with Karnovsky’s solution (0.075 M in phosphate buffer, pH 7.2-7.4, for 4 h) (Karnovsky, 1965), postfixed with osmium tetroxide (1% in the same buffer for 1 h), dehydrated in an acetate series and embedded in Araldite. Ultrathin sections were stained with 2% uranyl acetate for 15 min (Watson, 1958) and with lead citrate for 15 min (Reynolds, 1963) and then examined using a Philips EM 301 electron microscope.
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**Histochemical tests**

Histochemical studies were carried out on the immature and mature seeds, which were fixed immediately after harvest, embedded in paraffin, sectioned (8-10 µm) and mounted in synthetic resin. To detect lipids, material were fixed in buffered formalin and stained with Sudan Black B (Jensen, 1962) and Nile Blue (O’Brien and McCully, 1981). To detect proteins and polysaccharides, material were fixed in FAA 50 (formalin: acetic acid: alcohol 50%), and stained with Aniline Blue Black (Fisher, 1968) and PAS Reagent (O’Brien and McCully, 1981), respectively. Standard control procedures were carried out simultaneously. Photomicrographs were taken using a Leica model DM 4500 microscope.

**RESULTS**

Both immature and mature seeds were given the descriptions mentioned above; no differences in tissue types were observed.

**Ultrastructure**

Epidermal and parenchymatic cells of *C. echinata* cotyledons (Fig. 1) presented thick walls stained with PAS reagent and numerous plasmodesmata (Figs 1, 5). In the epidermal cells, plasmodesmata occurred in the anticlinal walls (Fig. 1), whereas in the parenchymatic cells plasmodesmata occurred in the anticlinal and periclinal walls. The nucleus was central and spherical in the epidermal cells (Fig. 1), but lobed in the subepidermal cells, with a conspicuous nucleolus (Fig. 3). The cytoplasm was abundant and electron-dense, rich in free ribosomes, globular mitochondriae with well-developed cristae and plastids with ellipsoid starch grains (Figs 2, 4, 6). Rough endoplasmic reticulum and protein bodies were distributed sparsely in the cytoplasm. Numerous electron-opaque oil bodies were concentrated near the cell wall (Figs 1, 2, 5); in the subepidermal cells, they were also found around plastids (Figs 3, 4). A larger quantity of plastids containing starch grains (Fig. 4) was found in the central cotyledonal cells. Cells of the hypocotyl-radicle axis differed from cotyledonal cells because of their trapezoidal nucleus (Fig. 7) and lack of endoplasmic reticulum. Uncertain organelles containing a thin granular matrix, as well as strongly electron-dense material undergoing degradation were found in the cytoplasm of the cotyledons (Fig. 4) and of the hypocotyl-radicle cells (Figs 7, 8).

In *C. ferrea*, the mycropilar endosperm cells had lobate nucleus with a prominent nucleolus (Fig. 9), a dense mitochondria-rich cytoplasm, endoplasmic reticulum, plastids with an electron-dense matrix, oil drops, plastids containing starch grains and hyperactive dictyosomes (Figs 10, 12). The cell wall stained with PAS reagent, became thick towards the embryo and formed numerous ingrowths inside the cell (Figs 10, 11). These kind of cell was only reported in *C. ferrea* mycropilar endosperm (compare figs 23, 24). The chalazal endosperm cells had thick PAS-positive walls with numerous plasmodesmata (Fig. 13); their cytoplasm was rich in plastids containing starch grains and oil droplets, mitochondriae, rough endoplasmic reticulum and dictyosomes (Figs 14-16). The remaining nucellar cells had vacuoles and cytoplasm containing mitochondriae, extensive rough endoplasmic reticulum, plastids containing oil droplets, and dictyosomes (Fig. 17). They were connected with the endosperm cells (Fig. 18) by numerous plasmodesmata.

**Histochemical tests**

Results confirmed that the main storage seed sites were the cotyledons in *C. echinata* and the endosperm in *C. ferrea* (Table 1). The storage substances found in the cotyledons of *C. echinata* were acid and saturated lipids (Figs. 1-6, 19), proteins (Fig. 20), starch grains (Fig. 4), and neutral polysaccharides. The endosperm of *C. ferrea* contained acid polysaccharides (Fig. 22) and starch grains (Fig. 12), and became gelatinous when hydrated. The foliar cotyledons of *C. ferrea* accumulated starch grains (Fig. 21) and numerous chloroplasts; the cell walls were made of neutral polysaccharides (Fig. 21).

**DISCUSSION**

Results found in the present work partly corroborated the hypothesis that had been proposed by Kozlowski and Gunn (1972). The main seed storage tissues were the cotyledons in *C. echinata* and the endosperm in *C. ferrea*, and that was expected for exendospermic and endospermic seeds, respectively (Boesewinkel and Bouman, 1984; Johri et al., 1992; Richards, 1997).
However, endosperm in *C. echinata* also accumulated substances throughout seed development and the photosynthesizing embryo in *C. ferrea* also accumulated starch grains. Moreover, endosperm, in both the species, presented mitochondriae with developed cristae, well-developed dictyosomes and rough endoplasmic reticulum, organelles that were commonly found in synthesizing tissues and substance conveyors (Bhatnagar and Sawhney, 1981). The beginning of the embryogenesis of these two species coincided with the beginning of the endosperm action towards the transport of substances and the development of an aggressive chalazal haustorium that went into nucellar cells up to the inner tegument cells (Teixeira et al., 2004), as well as labyrinth wall cells in *C. ferrea*. The labyrinth wall cells found in the mycropilar region of the endosperm in *C. ferrea* corresponded to the transference cells cited by Gunning and Pate (1969). According to DeWitt et al. (1999) and Thompson et al. (2001), the number of folds in such cell walls is responsible for the increasing of plasmalema surfaces and leads to the transport of solutes in the seeds.

In Leguminosae, labyrinth wall cells were observed in the inner layers of the seed testa and in the cotyledon epidermis of the seeds in *Vicia faba* (Talbot et al., 2001; Thompson et al., 2001) and not in the endosperm, as in *C. ferrea*. Before germination, the position of the labyrinth face in the cell showed that in the mature seeds of *C. ferrea*, the transport of nutrients was polarized towards the embryo, which was confirmed by the presence of a large amount of starch grains in the embryonary tissue. Still, the hydrophilic contents (acid and neutral polysaccharides) of the endosperm cells that were persistent in *C. ferrea* must be related to seed soaking and its resistant testa allowed the seed to remain in soil during several months (Lorenzi, 1998).

In *C. echinata*, the nutrient flow inside seeds occurred mainly through plasmodesmata, once transference cells were not observed. The cotyledon epidermic cells contacted both the nucellar cells (immature seed) and the other cotyledon, and embryonary axis cells through plasmodesmata. The final accumulator tissue, mainly starch, was made of cells from the cotyledon central region and showed that the substance mobilization got started from the outer parts.

Table 1 - Histochemical tests employed in the storage sites in seeds of *Caesalpinia echinata* and *C. ferrea* (The symbol + indicates positive reaction, i.e., presence of storage substance in seed sites, and the symbol – indicates negative reaction; * seed with cotyledonary stage of the embryo; after this stage the endosperm was completely consumed).

<table>
<thead>
<tr>
<th>Storage sites</th>
<th>Species</th>
<th>Sudan Black B (total lipids)</th>
<th>Nile Blue (acid lipids)</th>
<th>Aniline Blue-Black (proteins)</th>
<th>PAS + Toluidine Blue (polysaccharides)</th>
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<tbody>
<tr>
<td>endosperm</td>
<td><em>C. echinata</em></td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td><em>C. ferrea</em></td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>cotyledon</td>
<td><em>C. echinata</em></td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td><em>C. ferrea</em></td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Embryonary</td>
<td><em>C. echinata</em></td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>axis</td>
<td><em>C. ferrea</em></td>
<td>-</td>
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</table>
Figures 1-8 - Electron micrographs of *Caesalpinia echinata* mature embryo. 1-2. Epidermic cells of cotyledons. Note the central and spheric nucleus (n) and numerous oil bodies concentrated near the cell wall (cw) in 1, and the cytoplasm rich in mitochondriae (m), oil bodies (ob), and rough endoplasmic reticula (rer) in 2. 3-6. Parenchymatic cells of cotyledons. 3. Lobed nucleus (n) with a conspicuous nucleolus (nu). 4. Cytoplasm rich in oil bodies, plastids (p) containing starch grains, and organelles (uo) containing a thin granular matrix and electron-dense material undergoing degradation. 5. Cell wall (cw) with plasmodesmata, surrounded by numerous oil bodies and organelles containing a thin granular matrix (uo). 6. Detail of mitochondriae with well-developed cristae and of electron-opaque oil bodies. 7-8. Cells of hypocotyl-radicle axis. Note the trapezoidal nucleus (n) with a conspicuous nucleolus (nu) and the cytoplasm rich in organelles containing a thin granular matrix (uo) in 7, and plasmodesmata in the cell wall (cw) and oil bodies in 8. Scale bars: 1, 7 = 0.35 µm; 2, 6 = 0.05 µm; 3, 4 = 0.2 µm, 5 = 0.15 µm; 8 = 0.12 µm.
Figures 9-12 - Electron micrographs of the transfer cells of *Caesalpinia ferrea* mycropilar endosperm. 9. Note the lobed nucleus (n) with a conspicuous nucleolus (nu). 10. Note the cytoplasm rich in mitochondriae (m) and the cell wall (cw) with numerous ingrowths inside the cell. 11. Detail of the cell wall (cw) with numerous plasmodesmata (pl). 12. Cytoplasm with plastids (p) containing starch grains, mitochondriae (m), endoplasmic reticula (er) and dictyosomes (d). Scale bars: 9 = 1.02 µm; 10 = 0.75 µm; 11 = 0.24 µm; 12 = 0.43 µm.
Figures 13-18 - Electron micrographs of *Caesalpinia ferrea* chalazal endosperm (13-16) and remaining nucellar cells (17-18). 13. Cell wall (cw) with plasmodesmata and oil droplets. 14. Cytoplasm rich in extensive rough endoplasmic reticula (rer) and mitochondriae (m). 15. Detail of plastids (p) containing starch grains, and of mitochondriae (m). 16. Note the numerous dictyosomes (d) in the cytoplasm and numerous vacuoles. 17. Vacuoles (v) and cytoplasm containing mitochondriae (m), extensive rough endoplasmic reticulum (rer), plastids (p) containing oil droplets, and dictyosomes (d). 18. Cell walls of a nucellar cell (ncw) and of an endosperm cell (ecw). Scale bars: 13, 15, 16, 18 = 0.32 µm; 14 = 0.75 µm; 15 = 1.3 µm.
Figures 19-22 - Storage substances found in Caesalpinia echinata (19-20) and C. ferrea seed sites (21-22). 19. Cotyledonar cell contents reacted with Nile Blue indicating the presence of acid lipids. 20. Cotyledonar cell contents reacted with Aniline Blue-Black indicating the presence of proteins. 21. Epidermic (ep) and subepidermic (s) cells of cotyledons reacted with PAS Reagent and Toluidine Blue indicating the presence of starch grains in the cytoplasm and neutral polysaccharides in the cell walls. 22. Endosperm (e) reacted with Toluidine blue indicating the presence of acid polysaccharides. Scale bars: 19, 20 = 15 µm; 21 = 22 µm; 22 = 43 µm.

Figures 23-24 - Mature seed morphology of Caesalpinia echinata (23) and C. ferrea (24). Note the endosperm in C. ferrea seed containing transfer cells (dotted) in the mycropilar region. e – endosperm, c – cotyledon, hra – hypocotyl-radicle axis, sc – seed coat. Scale bars: 23 = 5 mm; 24 = 2 mm.
By comparing the seeds of *C. echinata* and *C. ferrea*, more storage substances were identified in the seeds of the former species, although a big amount of starch grains was found in the seeds of both the species; such grains presented glucose which could be promptly incorporated so that metabolic energy was generated (Buckeridge et al., 2000). Ultrastructural and histochemical evidence of proteins could be found in cotyledon cells of mature seeds in *C. echinata*, which showed numerous free ribosomes, rough endoplasmic reticulum, mitochondriae with developed cristae and strongly Aniline Black-stained contents in cells. According to Silva et al. (2000), proteins which were largely stored in sub cellular compartments and were not mobilized during the seed germination could be lecithins, these have been observed in the cotyledons of other species of Leguminosae, mainly Papilionoideae. Lecithins would act as defense proteins against predators and pathogens and would protect cotyledons while mobilization of the storage substances is being carried out (Cavada et al., 1993). Histophysiological studies on post-germination phases in the seeds of *C. echinata* would be necessary so that the function of the proteins observed in this work could be explained. Storage substances (neutral polysaccharides) have also been observed in the cell walls of tissues that make up the testa of seeds in both studied species. In the leguminous species, such polysaccharides have been identified as galactomannans, xiloglucans and galactans (Buckeridge et al., 2000); likewise starch, they could be the main storage carbohydrate in seeds (Buckeridge and Reid, 1996).

Finally, results presented in this work and previous seed developmental study obtained for the same species (Teixeira et al., 2004) have shown that the classification of seeds, based upon morphological data from only one development phase (usually the final phase), not always corresponded to their behavior during development. It is necessary to know ontogenesis, substances and their storage sites and the substance flow inside the seeds.

**REFERENCES**


**ACKNOWLEDGEMENTS**

We thank FAPESP (scholarship number 01/07124-1 and grant number 00/06422-4) for financial support, Cláudio José Barbedo and Rodrigo Augusto Santinelo Pereira for critically reading of the manuscript and Fernando Santiago dos Santos for English revision. The electron microscopy work was carried out at Centro de Microscopia Eletrônica of Universidade Estadual Paulista (UNESP), Botucatu, SP, Brazil.