Histochemical detection of acetogenins and storage molecules in the endosperm of *Annona macroprophyllata* Donn Sm. seeds

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Abstract

Acetogenins (ACGs) are bioactive compounds with cytotoxic properties in different cell lines. They are antitumoural, antiparasitic, antimalarial, insecticidal, antimicrobial, antifungal and antibacterial. These secondary metabolites function in plant defence and are found in specific organelles and specific cells, thereby preventing toxicity to the plant itself and permitting site-specific defence. The aim of this work was to histochemically determine the *in situ* localisation of ACGs in the endosperm of *Annona macroprophyllata* seeds using Kedde’s reagent. Additionally, the co-localisation of ACGs with other storage molecules was analysed. The seeds were analysed after 6 and 10 days of imbibition, when 1 or 2 cm of the radicle had emerged and metabolism was fully established. The seeds were then transversally cut in half at the midline and processed using different histological and histochemical techniques. Positive reactions with Kedde’s reagent were only observed in fresh, unfixed sections that were preserved in water, and staining was found only in the large cells (the idioblasts) at the periphery of the endosperm. The ACGs’ positive reaction with Sudan III corroborated their lipid nature. Paraflin sections stained with Naphthol Blue Black showed reactions in the endosperm parenchyma cells and stained the proteoplasts blue, indicating that they might correspond to storage sites for albumin-like proteins. Lugol’s iodine, which is similar in chemical composition to Wagner’s reagent, caused a golden brown reaction product in the cytoplasm of the idioblasts, which may indicate the presence of alkaloids.

Based on these results, we propose that Kedde’s reagent is an appropriate histochemical stain for detecting ACGs *in situ* in idioblasts and that idioblasts store ACGs and probably alkaloids. ACGs that are located in idioblasts found in restricted, peripheral areas of the endosperm could serve as a barrier that protects the seeds against insects and pathogen attack.

Introduction

Species in the *Annonaceae* have endospermic seeds with minute embryos. The ruminant endosperm occupies almost the entire volume of the seed and contains diverse storage molecules, including lipids, albumin-type proteins, and polysaccharides, such as starch and galactomannans.1-5 These storage substances are used during embryonic development, germination and seedling establishment, which occurs almost a month after the onset of germination.6 The endosperm of the mature seeds also contains secondary metabolites, such as alkaloids, polyphenols, essential oils, terpenes and acetogenins (ACGs), which are typical of these plants. Chemically, ACGs are C_{35}-C_{38} compounds of apparent polyketide origin that possess one or two tetrahydropurpuran rings and a β-lactone (either saturated or unsaturated), and they usually contain aliphatic regions that are variously hydroxylated, acetylated, or ketonised.7-10 The chromatographic separation of hexane extracts of *A. macroprophyllata* seeds led to the isolation of the ACGs laherradurine, cherimoline-2, rolliniastatin-2 and isorolliniastatin-2, which correspond to 0.1% of the endosperm’s weight.11-13 These ACGs are pharmacologically important because they are potent inhibitors of the mitochondrial NADH:ubiquinone-oxidoreductase (complex I of the respiratory chain). Laherradurine inhibits tumour cells but does not affect normal cells, and it could be used in the future as an antitumor drug.14,15 These secondary metabolites function in plant defence and occur in specific organelles and specific cells, thereby preventing toxicity to the plant itself and permitting site-specific defence. Staining with Kedde’s reagent is a well-established technique used to identify ACGs in extracts from leaves, seeds and other plant structures by thin layer chromatography; it reacts with the β-unsaturated-γ-lactone moiety commonly found in annonaceous ACGs.8,17,18

The aim of this work was to histochemically determine the *in situ* localisation of ACGs in sections of *A. macroprophyllata* seeds using Kedde’s reagent, as well as to analyse the co-localisation of ACGs with other storage molecules.

Materials and Methods

Plant materials

*A. macroprophyllata* seeds were collected in San Lucas, Chiapas, Mexico. The herborised sample was collected in June 2013 and was deposited in the Eizi Matuda Herbarium of the Universidad de Ciencias y Artes de Chiapas, Tuxtla Gutiérrez, Chiapas, Mexico under reference voucher #354.

Germination conditions

The seeds were germinated in Conviron environmental chambers that were maintained at 24-28°C with 12 h light/dark cycles and a relative humidity of approximately 65-75%. Light was delivered by cool white ultra-high-output fluorescent lamps at an intensity of 500 mW m⁻² s⁻¹.

Growth test

A total of 50 uncoated seeds were rehydrated on a paper sheet under controlled temperature and light conditions.19 Consistent with previous results,20 65% of the seeds germinated, and the seedlings were selected after 10 days of imbib-
tion, when the radicle was 1 or 2 cm in length and metabolism was fully established. The seeds were then cut in half at the midline and processed as illustrated in Figure 1.

**Seeds used for fresh sections**

Some of the seed halves were stored in water until sectioning on a cryostat. Others were stored in 70% alcohol, and others were fixed in FAA (10% formaldehyde/50% alcohol/5% acetic acid/35% water) for 72 h or in AGF (1.5% acrolein/3% glutaraldehyde/1.5% formaldehyde in 0.1 M sodium cacodylate buffer, pH 7.4) (Electron Microscopy Sciences, Hatfield, PA, USA) for 12 h. Then, the seed halves were washed with distilled water. Some halves were processed for paraffin embedding, and others remained in water until sectioning.

**Histochemistry on fresh sections**

The seeds that were stored in water, whether fixed or not, and the seeds that were stored in 70% alcohol were cut into 20-µm cross sections using a Leica CM1510S cryostat at between -14 and -18°C. Once on slides, the fresh sections were washed with distilled water three times for 5 min per wash to eliminate the Tissue Tech gel. The excess water near the section was absorbed with filter paper, and then the appropriate reagent was added. Kedde’s reagent (10% dinitrobenzoic acid and 2 N KOH in MeOH) is specific for ACGs, and a positive reaction yields a pink/magenta color. Lugol’s iodine solution (0.5 g iodine and 2.0 g KI in 100 mL distilled water) yields a dark purple color when it reacts with starch. Sudan III (0.1% in EtOH) yields a reddish orange color in the presence of fats. All analyses were performed at room temperature (approximately 23°C). After the reagents were applied, the samples were immediately observed through a microscope for ten to thirty minutes and photographed.

**Histochemistry on paraffin sections**

The paraffin-embedded samples were cut into 8-µm sections using a Microm HM 340 E microtome (Thermo Fisher Scientific, Inc., Waltham, MA, USA), deparaffinised, rehydrated, stained for 10 min with Kedde’s reagent to visualise the ACGs, and observed under a microscope. Other sections were rehydrated in 70% EtOH and stained with Naphthol Blue Black (1.0% in 50% EtOH) for protein analysis, and others were rehydrated in water, exposed to periodic acid (0.6%) for 1 min, and rinsed with distilled water. Then, Schiff reagent (Sigma-Aldrich 3952016; S. Louis, MO, USA) was added, incubated for 20 min and rinsed with distilled water. Then, the sample was dehydrated with xylene and mounted with resin. With this treatment, the polysaccharides stained a fuchsia color. Johansen’s quadruple stain was added to sections that had been rehydrated in 70% EtOH. According to Johansen, although the mixtures are rather complicated, the procedure is simple; differentiation is automatic, and little is left to personal judgment.
Microphotography

Photographs and idioblast measurements were taken with a Canon PowerShot A640 digital camera coupled to a Zeiss AxioStar Microscope with AxioVision software (Carl Zeiss Imaging Systems, Jena, Germany). Microscopic observations were made with A-Plan 10x/0.25 and A-Plan 40x/0.65 Ph2 objectives.

Results and Discussion

Positive reactions with Kedde’s reagent were only observed in fresh sections that were either unfixed or had been fixed in AGF and preserved in water, and staining was found only in the large cells (56×38 µm) at the periphery of the endosperm next to the tegmen; these cells are idioblasts (Figure 2A). The lipid nature of the ACGs was confirmed by their positive reaction with Sudan III in both fresh sections and sections that had been fixed in AGF (Figure 2B). Therefore, we propose that histochemical staining is appropriate for detecting ACGs and that the large cells are idioblasts that store ACGs.

Due to the use of solvents such as ethanol and xylene, none of the sections that had been fixed in FAA, stored in 70% alcohol, or embedded in paraffin showed staining in the idioblasts (Figure 2C). Naphthol Blue Black stained the cell walls blue, and the idioblast cytoplasm appeared secretory and lacked any special organelles; in the remaining endosperm cells, blue-stained proteoplasts were detected (Figure 2D) and might correspond to storage sites for albumin-type proteins, which have been reported to be stored in the endosperm of A. muricata and A. macroprophyllata seeds. Johansen’s quadruple stain turned these organelles purple (Figure 3A), confirming the presence of protein reserves.

The presence of oils,21 which act as reserve substances in the endosperm, was evident based on their reactivity with Sudan III (Figure 2B) and, indirectly, by the spaces observed in the cells in the paraffin-embedded material (Figure 3B). The material fixed in AGF, which had been preserved in water and sectioned on a cryostat, still contained ACGs in the idioblasts. The plastids were not stained by Schiff reagent (Figure 3C), demonstrating that they do not contain starch; their reaction with the crystal violet in Johansen’s quadruple stain indirectly indicates that they contain proteins (Figure 2D).

Staining with Lugol’s iodine (Figure 3D) did not indicate the presence of starch, but the reagent generated a golden brown color in the idioblasts in the same place as the reaction with Kedde’s reagent, indicating the presence of dextrins.5 However, the content appeared more like a drop-shaped liquid than the expected solid particles. This golden brown reaction product between the cytoplasmic content of the idioblasts and Lugol’s iodine, which is similar in chemical composition to Wagner reagent,24 may indicate the presence of alkaloids. Therefore, this reaction deserves further histochemical study; the biosynthesis of liriodenine, an aporphinic alkaloid, begins during the early stages of development of seedlings of Annona macroprophyllata Syn A. diversifolia Saff.4

ACGs are bioactive compounds with cytotoxic properties in multiple cell lines, and they have antitumour, antiparasitic, antimarial, insecticidal, antimicrobial, antifungal and antibacterial effects.17 In particular, the acetogenin laherradurin might act as an autoinducer or quorum-sensing signalling molecule that affects the expression of genes involved in biofilm formation in Pseudomonas plecoglossicida 326.18 Since ACGs are located in idioblasts that are found only in restricted areas in the periphery of the endosperm, despite the large amount of protein and oil reserves contained in the endosperm of mature seeds,25 ACGs may act as a barrier that protects seeds against insects and pathogens. In addition, the presence of alkaloids in the idioblasts might support this idea, because alkaloids are also strongly insect repellent and toxic to microorganisms.17

Villamil et al. reported amylloid granules in A. muricata endosperm.4 However, in A. macroprophyllata, amylloids are not present because do not show blue or purple staining with Lugol’s iodine in the endosperm cell walls, as reported by Kooiman2 for 25 species of Annonaceae. Some endosperm cells in A. macroprophyllata stain brown, which may correspond to dextrins. As described by Kooiman,2 the term amylloid refers to a group of polysaccharides with similar structures, particularly those reported by Schleiden in 1838 (in 2) and by Vogel and Schleiden in 1839 (in 2), who first used the term. Using Mitchell reagent, which has the same composition as Lugol’s iodine, Kooiman detected cellulose and hemicellulose via blue staining in the presence of 75% sulphuric acid. Consistent with these ideas, we suggest that the term amylloid should no longer be used to refer to the group of compounds that can be detected with Schiff
reagent, as the term is now used to refer to a beta-fibrillar protein of animal origin, the amyloid beta-protein.

This new histochemical method for detecting ACUs in situ in seed sections using Kedde’s reagent can test for the presence of these substances in developing tissue or mature organs. It can also differentiate between various substances in the same structures, as occurred using Warner reagent to detect alkaloids in idioblasts. This work histochemically demonstrates the compartmentalisation of ACGs in the endosperm, suggesting that its aforementioned characteristics are part of a protective strategy that allows selective defence for the sexual propagation of the species.

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