A Variation of C₄ Leaf Anatomy in *Arundinella hirta* (Gramineae)¹

R. KENT CROOKSTON² AND DALE N. MOSS

Department of Agronomy and Plant Genetics, University of Minnesota, St. Paul, Minnesota 55101

Received for publication May 14, 1973

ABSTRACT

The species *Arundinella hirta* L. possesses a striking variation of the leaf anatomy that is characteristic of C₄ grasses. In addition to a sheath of large, bright green cells around the vascular bundles, there are strands of large parenchyma cells which appear identical to the bundle sheath cells and which run parallel to the vascular bundles, but which are not associated with any vascular tissue. This species may be useful for studying the cellular compartmentalization associated with the C₄ pathway and should provide interesting material for determining the role of translocation in the functioning of the C₄ system.

A report exists, however, of a species which we believed had the C₃ pathway for photosynthesis but which appeared to have "bundle sheath-type" cells not associated directly with vascular bundles. Brown (2) reported that "in the genus *Arundinella* cells which are similar to parenchyma sheath cells and form starch are scattered in the chlorenchyma." Tregunna et al. (17) reported that *Arundinella hirta* was a C₃ species.

We confirmed that the C₃ system functions in *A. hirta* and attempted to characterize its specialized starch-storing leaf cells which were not associated with vascular tissue.

MATERIALS AND METHODS

Plants were grown from seed that was sown in a 2:1 mixture of loam and peat moss in 13-cm clay pots. These were placed in growth chambers with 27 to 21 C day-night temperatures, a 16-hr day length, and a light intensity from incandescent and fluorescent lamps of 0.1 cal cm⁻² min⁻¹ (400–700 nm).

CO₂ compensation measurements were made using leaves from plants that were approximately 6 weeks old. The CO₂ compensation system has been described in a previous report (13).

Fully expanded leaves were used for all anatomical observations. Freshly cut cross sections were stained with I₂-KI to observe starch storage patterns. Leaf tissue was also prepared for examination in the electron microscope. Leaves were cut into segments approximately 1 mm square and fixed for 2 hr in a 2.5% glutaraldehyde solution freshly treated with BaCO₃ and buffered with 0.05 M sodium phosphate buffer (pH 6.8). The segments were then washed in buffer, postfixed in OsO₄ (1% aqueous solution) for 1 hr, dehydrated with a water-acetone series ending in dry acetone, and embedded in Spurr's (16) embedding mixture. Silver sections were cut with a diamond knife and a Sorvall MT-2 ultramicrotome. The sections were mounted on copper grids, stained in uranyl acetate (10 min) and lead citrate (3 min), and viewed with a Phillips 300 electron microscope operated at 80 kv.

Thicker sections (1 μ) of the same material were also cut with glass knives for light microscopy. These sections were dried onto glass slides, stained with 0.05% toluidine blue for 30 sec (80 C), rinsed and dried again, and mounted under cover slips with a permanent mounting medium. They were photographed with a Zeiss research microscope.

Leaves were also cleared and stained for paradermal viewing. Fresh leaf material was placed in boiling 80% (v/v) ethanol until the chlorophyll had been extracted. It was then placed in 10% aqueous NaOH solution and left until it had cleared. The cleared material was rinsed in distilled water and then stained with 1% I₂-KI solution. Stained tissue was placed on a glass slide in water, covered with a cover glass, and examined with the light microscope.

¹ Minnesota Agricultural Experiment Station Journal Series Paper 8177. This work was supported in part by Rockefeller Foundation Grant Ga Agr 7035.

² Present address: Department of Vegetable Crops, Cornell University, Ithaca, New York 14850.
RESULTS AND DISCUSSION

As measured in our system, the CO₂ compensation point of *Arundinella hirta* was less than 5 μl CO₂/liter. The leaves had vascular bundles surrounded with prominent thick-walled starch-storing sheaths. We therefore concluded (3) that *A. hirta* was a C₄ species.

Figure 1 is a cross-sectional view of an *A. hirta* leaf, illus-

---

**Fig. 1.** A cross-sectional view of an *Arundinella hirta* leaf. Two vascular bundles are shown enclosed by sheaths of prominent cells containing numerous dark chloroplasts. Between the vascular bundles are individual prominent cells which resemble the bundle sheath cells (i.e., contain numerous dark chloroplasts), but which are not associated with vascular tissue. × 200.

**Fig. 2.** A longitudinal section through a vascular bundle of *A. hirta*. × 200.

**Fig. 3.** A longitudinal section through a row of specialized parenchyma cells of *A. hirta*. × 200.

**Fig. 4.** A cross-sectional view of some vascular bundle sheath and mesophyll cells of *A. hirta*. × 650.

**Fig. 5.** A cross-sectional view of two specialized parenchyma cells and some mesophyll cells of *A. hirta*. × 650.
trating the irregular arrangement of bundle sheath-type leaf cells as reported by Brown (2). In addition to the regular sheath of large, bright green cells around the vascular bundle, there are single large parenchyma cells which appear identical to the bundle sheath cells, but which are not associated with vascular tissue. Hereafter, we will refer to these cells as “specialized parenchyma cells.” Figure 2 shows a longitudinal section through a vascular bundle of *A. hirta*. Figure 3 shows a longitudinal section through the specialized parenchyma cell area and illustrates the continuous, end-to-end nature of the arrangement of these cells. Figures 4 and 5 are close-ups of portions of Figure 1 and show some bundle sheath cells and two of the specialized parenchyma cells at a higher magnification. The similarity between the two cells (bundle sheath and specialized parenchyma) is clear. The chloroplasts of both are large, dark, and closely packed and contain numerous starch grains (black spots). They contrast markedly with the smaller chloroplasts of the mesophyll cells which are lighter, less crowded, and relatively free of starch. The chloroplasts of both the bundle sheath and specialized parenchyma cells have a centrifugal arrangement (toward the surrounding mesophyll). The specialized parenchyma cells are not associated with any vascular tissue, however.

Figure 6 shows a paradermal view of an *A. hirta* leaf that has been cleared with NaOH and stained with I$_2$-KI. The large dark rows are vascular bundles with sheaths of surrounding parenchyma cells specialized for starch storage. Several rows of specialized parenchyma cells can be seen running in single strands parallel to the vascular bundles. These strands were generally continuous (end to end) throughout all leaf segments. Occasionally, however, one row ended and another began, or became double (extreme lower right of Fig. 6). The lack of association of the specialized parenchyma cells with vascular tissue is again apparent.

The specialized parenchyma cells did attach to vascular tissue under certain conditions, however. Figure 7 is a segment of a cleared *A. hirta* leaf showing a cross-vein which runs from one vascular bundle to another (lower left to upper right). As the cross-vein passes the rows of specialized parenchyma cells, some of the cells become appressed to the cross-vein, much like a normal bundle sheath. This happened with all the traversing veins which we observed. These cross-veins were few in number, however, and provided a vascular-bundle association for only an insignificant proportion of the specialized parenchyma cells. Figure 8 illustrates the same type of association of specialized parenchyma cells with vascular tissue as is seen in Figure 7, except the view is cross-sectional.

Figure 9 is an electron micrograph of bundle sheath and

---

Fig. 6. A paradermal view of a portion of an *A. hirta* leaf that has been cleared and stained with I$_2$-KI. The starch-containing vascular bundle sheath and rows of specialized cells show clearly. × 125.

Fig. 7. A paradermal view of a portion of an *A. hirta* leaf showing a cross-vein (arrow) traversing three rows of specialized cells. Some of the specialized cells have become appressed to the cross-vein. × 300.

Fig. 8. A cross-sectional view of an *A. hirta* leaf showing specialized parenchyma cells appressed to a cross-vein. × 750.
Fig. 9. An electron micrograph of the bundle sheath region of *A. hirta*. Portions of two bundle sheath cells (upper left) are shown in relation to adjoining mesophyll cells. The large chloroplasts of the bundle sheath cells are agranal and contain numerous starch grains. The chloroplasts of the mesophyll cells are free of starch and have well developed grana. × 6400.

Fig. 10. An electron micrograph of a specialized parenchyma cell. The ultrastructure of the specialized cell appears identical to that of the bundle sheath cells (Fig. 10). An electron-opaque band, which is thickened over the numerous pit fields containing plasmodesmata, is located within the cell wall of both the specialized parenchyma and vascular bundle sheath cells. × 6400.
mesophyll cells. Figure 10 is an electron micrograph of a specialized parenchyma cell. The chloroplasts of both the bundle sheath and specialized parenchyma cells contain numerous starch grains and have underdeveloped grana. The smaller mesophyll chloroplasts contain very little starch but have well developed grana. An electron-opaque band is found in the thick wall of both cells. This band forms a cylinder around the strand of specialized cells much like the cylinder around the vascular bundle in the outer walls of the bundle sheath. Prominent pits with plasmodesmata can be seen in the walls of both cells. It is apparent that the ultrastructural traits of the specialized parenchyma cells are identical to those of the vascular bundle sheath cells. The fact that the chloroplasts of the specialized parenchyma cells of *A. hirta* are rich in starch indicates that these cells have functional as well as structural similarities to bundle sheath cells.

It is interesting to consider the peculiar location of starch-storing parenchyma cells in *A. hirta* in terms of the role of mesophyll and bundle sheath cells in the operation of the C₄ pathway. The proposed pathway (8) requires the close association of bundle sheath and mesophyll cells. Vascular bundles are so frequent in most C₄ grasses that almost all of the mesophyll cells are in direct contact with a bundle sheath cell; *i.e.*, vascular bundle sheaths are usually spaced at intervals only two mesophyll cells apart (4). In contrast, the vascular bundles of *A. hirta* are spaced at an average of seven mesophyll cells apart. The location of specialized parenchyma cells in mid-mesophyll positions in *A. hirta*, therefore, provides a cell which appears to be identical to a bundle sheath cell (except that it lacks a vascular connection) at intervals no greater than two mesophyll cells apart. This restores, in part, the normal C₄ cell pattern and would allow the C₄ pathway of photosynthesis to operate in all the regions of *A. hirta* leaves.

The question arises, however, concerning the role of translocation in the functioning of the C₄ system. That the strands of specialized parenchyma cells do not have vein-like translocation properties is apparent from Figure 11. The cell wall region between two specialized cells is shown (longitudinal view). The view is representative of all such walls observed and shows the impermeable appearance of the connection between these cells. No pit fields or plasmodesmata could be found connecting two specialized cells, although they were plentiful in the walls shared with the surrounding mesophyll.

The presence of specialized parenchyma cells in *A. hirta* might not permit the functioning of the C₄ system at a maximal rate for any extended period, therefore, unless the starch-storing capacity of these cells is sufficient to accommodate the entire day's synthesis of carbon from adjacent mesophyll cells. Hydrolysis and translocation via the vascular bundles during the night would then be required to make way for the production of the next day.

The segments of *A. hirta* leaves between vascular bundles are sufficiently large that they could probably be dissected from the leaf, allowing the recovery of tissue containing only normal and specialized parenchyma. Such tissue could be useful in studying both the cellular compartmentalization of the C₄ syndrome and of the role of translocation in the functioning of the C₄ system. It may also be possible to learn more about the differentiation of the leaf anatomy that is characteristic of C₄ plants by studying this interesting species.

Acknowledgments—The authors are grateful to J. R. Stander for his technical assistance in obtaining the electron micrographs used in this study.

**LITERATURE CITED**