Celery Breeding Program at the Department of Vegetable Crops, University of California, Davis

Celery (Apium graveolens L.) is an important vegetable in California and is grown mostly in the coastal areas of Ventura, Monterey, San Luis Obispo, and Santa Barbara counties. According to 1989 statistics, 8900 ha of celery, valued at about $164 million, was harvested that year. California produces 50% to 60% of the total celery grown in the United States—enough to supply its own demand and export the remainder to other parts of the country. Recognizing the high value and importance of this crop, California growers established the California Celery Research Advisory Board in Oct. 1976 to promote celery research. The board has supported our research program since its initiation.

Apium graveolens is used not only for its stalks, but also for its seeds, leaves, and roots. Variety dulce commonly is cultivated in the United States for its leaf stalks. The leaves of secalinum (smallage) are used as a garnish, flavoring, or medicine, and rapaceum (celeriac) is used for its enlarged root/hypocotyl and is grown predominantly in Europe.

The Dept. of Vegetable Crops, Univ. of California (UC), Davis, is one of the few institutions in the United States and abroad currently conducting research in celery breeding and genetics. Our leading role in this area of research matches the California celery industry’s leadership as the world’s main celery producer. The main objective of our program at UC Davis is to develop disease- and pest-resistant lines that include resistance to fusarium yellows, late blight, developing resistant varieties, 'Tall Utah 52-70R', 'T.U. 52-70HK', and 'T.U. 52-75'. As a product of this program, we have released the next generation of fusarium-resistant lines, UC8-1, UC 10-1, and UC26-1, which are described on p. 351. Our goal is to advance this material further for additional releases in the future. Typical trimmed and untrimmed plants of these lines are depicted on the front cover (UC26-1 and UC10-1) and in Fig. 1 (UC8-1).

Late blight, caused by the pathogen Septoria apicola Speg., is another important disease affecting celery. Although there are means to control late blight, developing resistant varieties will reduce production costs and fungicide applications. Although our search for resis

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Front cover: UC26-1 and UC 10-1 celery plants. Photographs by Donald C. Edwards, principal photographer, Dept. of Pomology, Univ. of California, Davis.

Fig. 1. Trimmed and untrimmed plants of line UC8-1.

(continued on p. 359)
Despite resistance in cultivated accessions was unsuccessful, we found resistance in the wild species *A. chilense* Zoll. and *A. nodiflorum* (L.) Lag. (Ochoa and Quiros, 1989). *Apium chilense* hybridized readily to celery, but the F₁ hybrids had poor fertility due to the presence of chromosomal rearrangements. Although it was not possible to obtain F₂ seed, backcross seed was obtained, and we are trying to produce late-blight-resistant lines from this material.

Leafminer (*Lyriomiza trifoli* Burgess) is a major pest of celery and currently is being controlled by routine insecticide spraying. Developing leafminer-resistant celery varieties will reduce hazardous chemical use and environmental risks linked to their use. No resistance to this pest was observed in the cultivated collection; however, we found an accession of *A. prostratum* Vent. from Australia to be practically immune in no-choice greenhouse tests. No feeding or oviposition was observed in this species (Trumble and Quiros, 1988). We have hybridized *A. prostratum* to celery and derived BC₁ progeny in an attempt to develop leafminer-resistant lines. Chemical analysis of the foliage is being done in conduction with the breeding program to ensure the resistance is not based on noxious compounds (Trumble et al., 1990).

**Developing celery genetics**

Developing the genetics of this crop will expedite breeding. We are creating chromosome markers and a genetic map based on isozymes, nonspecific proteins, morphological traits (Quiros et al., 1987), and DNA-based markers. The latter includes restriction fragment length polymorphism and amplified DNA sequences by the polymerase chain reaction. Through the genetic map, we can tag useful genes, such as those determining fusarium resistance and male sterility. We are also interested in using male sterility to develop disease-resistant F₁ hybrids (Quiros et al., 1986). Male-sterile plants were found in P1229526, a land race from Iran.

Finally, another research component is to adapt new genetic engineering techniques that would be useful to breeders. This includes developing transformation procedures for gene transfer. We accomplished this task using *Agrobacterium tumefaciens* as a vector (Catlin et al., 1988).

In summary, we are confident that the celery genetics program at UC Davis, with its broad germplasm base and comprehensive genetic information, will serve to adapt and solve present and future problems faced by celery growers.

**Literature Cited**


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