The genus *Rudbeckia* consists of ≈25 annual, biennial, or perennial herbaceous species native to most of North America, except for the extreme northern and southern regions. Most species are concentrated east of the Rocky Mountains (Liberty Hyde Bailey Hortorium, 1976). This genus, a member of the Helianthus tribe within the Asteraceae, was named by Linnaeus after father and son Swedish botanists Olaus and Johan Rudbeck (Gledhill, 1989; Kochankov and Chailakhyan, 1986). *Rudbeckia* species are used widely as ornamentals but have also served as forage, silage, and nectar sources (Kochankov and Chailakhyan, 1986).

The genus *Rudbeckia* is generally characterized by floral heads composed of disk and ray florets. The ray florets are commonly yellow or orange and sometimes have red markings. The chaffy receptacle can be conical or convex with fertile disk florets and divided styles; it is subtended by more than 50 involucral bracts. The epidermis is usually rough, and the leaves are alternate, undivided, and lobed or pinnatifid (Wofford, 1989). *Rudbeckia* are pollinated by insects and, in at least three species, the ultraviolet (UV) reflection patterns are different enough to allow for pollinator discrimination (Abrahamson and McCrea, 1977). The apomictic species, *R. triloba* and *R. laciniata*, have limited intra-population variation in UV nectar guide patterns, while *R. hirta*, an obligate outbreeder, has greater variation (Abrahamson and McCrea, 1977). Little evidence of interspecific hybridization exists even though colonies of two or more species may frequently occur in close proximity (Perdue, 1957).

*Rudbeckia hirta*'s in florescence has a prominent central cone of disk florets, normally purplish black, with acute bracts and elongated, subulate styles. The ray florets are bright yellow or yellow with rust markings, giving rise to the common name “black-eyed Susan” or “coneflower.” The stems and leaves are coarsely hairy, and the species has been considered an annual, biennial, or short-lived perennial depending on location and genetic background. While native to North America, *R. hirta* is one of the most common species of the genus and is frequently observed throughout the world as a result of its successful naturalization abilities.

As cultivated plants, *R. hirta* selections have a nonuniform appearance; bloom from July through August; are not consistently available; and perform well in gardens, wildflower meadow mixes, and as a species component of highway beautification projects. They make excellent cut flowers, but, to our knowledge, their postharvest characteristics have not been researched.

Many forms of *R. hirta* are known, which has led to a plethora of varietal names (Gray Herbarium, 1968). Clute (1913) reported one variation in ray floret coloration as a form of *R. speciosa*, then known as *R. bicolor*. The Gray Herbarium lists *R. speciosa* as a variety of *R. fulgida*, and *R. bicolor* as a variety of *R. hirta* (Gray Herbarium, 1968). In his work with color variation in *Rudbeckia*, Clute (1913) divided the three main color forms into *R. hirta* var. *bicolor*, *R. hirta* var. *rubra*, and *R. hirta* var. *annulata*, yet he also concluded that these forms were unstable. Later, Clute (1924) reported that the form of *R. hirta* with a red blotch at the base of each ray floret was *R. hirta* var. *pulcherrima*, the same color form he previously listed as *R. hirta* var. *bicolor*. To further confuse the taxonomic listing of this species, Fernand and Schubert (1948) identified the species *R. hirta* var. *pulcherrima* as a varietal form of *R. serotina*. This conclusion was based on its strong tendency to sport, whereas *R. hirta*, originating from an older ecological region, was reportedly stable. Perdue (1957) reported *R. bicolor* to be synonymous with *R. hirta* var. *pulcherrima* as first listed by Farwell (1904). The Gray Herbarium later listed *R. bicolor* as a variety of *R. hirta*, separate from *R. pulcherrima*, and reduced *R. serotina* to a variety of *R. hirta*. Most recent classification places *R. bicolor* synonymous with *R. hirta* var. *pulcherrima* Farw. (Liberty Hyde Bailey Hortorium, 1976) or just *R. hirta* L. (Huxley et al., 1992), yet it is likely that all these plants are marketed for ornamental use simply as *R. hirta*.

As cultivated plants, *R. hirta* selections may be loosely distinguished based on three features: flowering plant height, ray floret color, and single vs. double inflorescences (number of ray florets). ‘Becky’, a true di-ocious standout, is one of the shortest cultivars, growing no taller than 20 to 25 cm. Plant stature appears to be the main breeding objective in developing ‘Becky’, because its flower size is variable and inflorescence colors range from pure gold to heavily flushed with brown.

Taller cultivars that reliably produce brown- or maroon-marked golden flowers include, but are not limited to, ‘Autumn Forest’, ‘Sputnik’, ‘Nutmeg’, ‘My Joy’, ‘Rustic Colors’, and ‘Rustic Dwarfs’. Unmarked, yellow-gold flowers are generally produced by ‘Golden Daisy’, ‘Goldilocks’, ‘Marmalade’, and ‘Pinwheel’, all of which exhibit some degree of doubleness. In this group, ‘Goldilocks’ is the shortest and is similar in size to ‘Becky’. ‘Double Gold’ and ‘Double Daisy’ are largely what their names imply—monochromatic inflorescences composed of abundant ray florets, resulting in dramatic double flowers. One of the newest cultivars, ‘Indian Summer’, is a 1995 All-American Selection winner. Each ‘Indian Summer’ inflorescence reaches 15 to 22 cm in diameter and has golden-yellow rays and dark disk florets. In our opinion, not all *R. hirta* cultivars are distinctly different, leading to the suspicion that some cultivars may be synonymous, depending on the seed source. ‘Irish Eyes’, however, is a uniquely beautiful cultivar that possesses a greenish rather than brown cone of disk florets. This trait has been incorporated into a recent bedding plant release of *Sanvitalia procumbens* Lam., also named ‘Irish Eyes’ (K. Sahin, Zaden B. V., Rijn, The Netherlands) and a close relative of the genus *Rudbeckia*. ‘Gloriosa Daisy’ and ‘Double Gloriosa Daisy’ are tetraploid strains tending to be more reliably perennial than other cultivars but still blooming the first year from seed as do their diploid counterparts.

**Research history**

In the 1920s, Garner and Allard (1925) first described *R. hirta* as a long-day (LD) plant. Since then, it has been used as a model for LD plant photoperiodicity and for flowering research (Austin, 1941; Greulach, 1942; Mumeek, 1936, 1940). These studies focused on the effects of temperature and critical daylength on flowering (Kochankov and Chailakhyan, 1986; Tanamoto and Harada, 1985). Following World War II, European research teams also used *R. hirta* to study flowering. These efforts focused on the physiological effects of plant growth substances, predominantly retardants, on stem growth and flowering (Kochankov and Chailakhyan, 1986; Orvos and Lyons, 1989). Selected effects of applied gibberellins and cytokinins are reported in an article on p. 141 of this *HortScience* issue (Harkess and Lyons, 1994).

As previously reported by Mumeek (1940) and more recently expanded by Orvos and Lyons (1989), plant height in *R. hirta* can be controlled by photoperiodic manipulation of its bolting behavior. While in the vegetative stage, the species maintains a flat rosette of leaves. The onset of flower initiation, however, is easily detected by a noticeable arching of the foliar canopy upward and away from the soil surface. The central stem elongates shortly thereafter and may eventually attain a length of 1 m at the time of terminal inflorescence anthesis, depending on the cultivar. Flowering (continued on p. 227)
and stem elongation are not inextricably linked, yet they do respond to the same external LD stimulus. Stem elongation ceases under short days (SD), while floral development, once initiated, does not. In fact, there is no period of total inability to flower in *R. hirta*. Sensitivity to photoperiodic induction increases with age, with the peak sensitivity occurring after the plant has produced ≈14 expanded leaves (Harkess and Lyons, 1993a; Orvos and Lyons, 1989). This vegetative stage signifies maturity in *R. hirta*. In florescence initiation depends on LD; however, once involucral bracts have appeared, a return to SD does not cause development to cease or revert (Harkess and Lyons, 1993b). As few as eight LDs can initiate flowering in some plants, but usually at least 14 LDs are required for 100% of the plants to flower. The axillary meristems are affected similarly, but initiate only after terminal meristem initiation (Harkess and Lyons, 1993b). This characteristic cannot be ignored when limited photoperiodic treatments are used specifically for height control. Although fewer LD will minimize plant height, the potential number of inflorescences will also be reduced because of a lack of axillary inflorescence production.

Recently, histological examination of the terminal meristem has outlined the major anatomical changes occurring in *R. hirta* during floral initiation and development (Harkess and Lyons, 1993a). Fluctuations in polyamides have also been correlated to these anatomical changes. Putrescine and spermidine increase during floral initiation and decrease during subsequent development (Harkess et al., 1992). This response is opposite to that observed in SD plants and is similar to results previously reported with cytokinins (Hamasaki and Galston, 1990; Krekule et al., 1989).

Additional research has indicated that, as in other species, ecotype origin has a profound effect on the flowering response of *R. hirta*. An ecotype native to Texas had a weak LD response when compared with the Texas ecotype *Orthosiphon aristatus* (Fernald, 1937). Ecotypes originating from different latitudes. MS thesis, University of Georgia Press, Athens.

**Literature Cited**


**RICHARD L. HARKESS** and **ROBERT E. LYONS**

Dept. of Horticulture Virginia Polytechnic Institute and State Univ. Blacksburg, VA 24061-0327