

Sex Change in Plants: Old and New Observations and New Hypotheses

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Summary. Evidence is presented that individuals of a large number of dioecious and subdioecious plant species are able to alter their sexual state in response to changes in the ambient environment and/or changes in size or age. We suggest that lability of sexual expression probably has survival value where a significant portion of the females must otherwise bear the cost of fruit production in unfavorable environments. We demonstrate that in patchy environments of the proper scale and variability in quality, labile sexual expression will enhance an individual's genetic contribution to the next generation.

Introduction

The idea that the sex of an individual of a dioecious plant species is fixed throughout life and is determined solely by the genetic composition of the individual is prevalent in current literature despite the existence of a large literature to the contrary. An examination of introductory biology (Wilson et al. 1978) botany, (Haney 1978, Fuller et al. 1972; Fuller and Ritchie 1967) and mid level genetic texts (Garber 1972; Srb et al. 1965; Burns 1969; Strickberger 1976; Grant 1975 and Gardener 1975) demonstrates that the view most often presented is one in which sex is controlled completely by a chromosomal or genic system. Rarely does one find a discussion of the role of environment in the sex determination of individuals of dioecious plant species, although Haney (1978), Strickberger (1976) and Srb et al. (1965) do present some discussion of this phenomenon. Thus because other possibilities are not usually presented, readers may draw the conclusion that sexual expression of individuals of all dioecious species is genetically fixed and not responsive to environmental influences. However, workers seeking to establish the genetic basis of sex have recognized for over 60 years that hermaphrodites are common in the prevailingly dioecious plant species they have studied and that the sexual state of individuals sometimes changed (Yampolsky 1919; Heslop-Harrison 1924; Correns 1928). Indeed such a recognition lead Hartman (1956) to formulate his so-called "law of bisexual potentiality of both sexes" (also see Smith 1963).

The inconstancy of the sexual state of individuals of higher plants has been well documented for a number of taxa described as dioecious, subdioecious or sequential hermaphrodites by professional botanists. The labile nature of sex expression by individuals of many dioecious and sequentially hermaphroditic plant species is documented from the literature in Table 1. Our literature search shows that 25 families and over 50 species include unisexed individuals that have functioned as male at one time and females at another or have produced hermaphroditic offspring (the reason for including those which produce hermaphroditic offspring will be discussed later). Several of the species noted in Table l have been reported to have sex chromosomes. Of the 13 species Westergaard (1958) listed as having well differentiated sex chromosomes, individuals of 3 species *(Cannabis sativa, Humulusjaponicus* and *Rumex hastatulus)* are known to exhibit sex change and/or to produce hermaphroditic offspring. In Table 1, we designate those species which are reported to have sex chromosomes to underscore the fact that sex change is a phenomenon of at least some dioecious plant species.

A large body of literature documents the fact that a variety of environmental factors are capable of altering the sexual expression of plant individuals that normally display but one sex at a time (Table 2). The ratio of male-to-female flowers on individuals of selected monoecious species and of male-to-female organs in flowers of perfect flowered taxa has likewise proven to be manipulatable through alteration of the plant's environment. As documented in Table 2, age, injury and disease have all been shown to alter the sexual expression of individuals of some species. Similarly, physical and chemical characteristics of the environment (e.g., light intensity, day length, temperature, soil fertility, relative availability of soil moisture and air chemistry) are reported to affect the sexual expression of a variety of species. Finally, plant physiologists have repeatedly demonstrated that the sexual expression of developing floral buds can be altered in predictable ways through the application of several plant hormones in different amounts and proportions (Table 2; for greater detail see Heslop-Harrison 1972 and Chailakhyan 1979).

It will be noted in Table 2 that some environmental variables predispose individual plants of labile sexual expression toward femaleness while other variables favor maleness. Some variables such as dry soils or high tissue levels of cytokinin seem to consistently favor one sex over the other, while other variables (e.g., gibberellins or trauma) sometimes favor maleness and at other times femaleness. Response in the latter case at least seems to be dependent-upon the species and/or the physiological state of the individual.

Some workers reserve the term dioecious for plant species which consist of males and females only (no hermaphrodites). Such workers also exclude species in which sexual expression of individuals changes with time. Strict adherence to that definition would greatly reduce the number of species now recognized as dioecious in the world's floras. Regardless of definitions, lability of sexual expression of individuals and the occurrence of occasional hermaphrodites in prevailingly dioecious plant species are phenomena which deserve scientific investigation since they may have considerable evolutionary and ecologicai consequences

Table 1. Species that are commonly described as dioecious but which include individuals known to display different sexual states (i.e., male or female) during their life or to produce hermaphroditic offspring. We have included species which taxonomists describe as monoecious to dioecious only when such species have been reported to have sex chromosomes (e.g., *Mercurialis annua, Itex* sp., and *Empetrum nigrum)* or when the monoecious state appears to be related to the sex reversal phenomenon (as for *Arisaema triphyllum, A. dracontium, A. japonicus* and some Orchids). An asterisk designates species reported by Westergaard to have well-established heteromorphic sex chromosomes (Westergaard 1958). Nomenclature follows Bailey and Bailey (1976)

Table 1 (continued)

¹ *Ilex serrata* has been reported to have sex chromosomes (Nakajima 1942)

Table 2. Some factors known to modify the sexual expression of vascular plants under controlled conditions. Where more than one worker has shown altered sexual expression for a given species under a given kind of treatment, we have cited pertinent review articles in order to minimize the table's length. Morphological changes in which organs of one sex or the other were induced or caused to abort were observed by all authors, but most of the references do not provide proof that the organs produced viable gametes. No distinction is made between ultimate and proximal factors since in several cases the distinction is often described as either obscure or artificial. An asterisk indicates dioecious species. Species not asterisked are monoecious or perfect flowered taxa on which the ratio of male/female flowers is significantly altered by the environmental factor considered. Nomenclature follows Bailey and Bailey (1976)

Table 2 (continued)

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The data in Table 2 reveal a strong tendency for environmental stresses to induce maleness. For example, low soil fertility, dry soils, extreme temperatures, and low light intensity all tend to incline the sex ratio toward males. For species in which the cost (in either increased mortality or decreased competitive ability) of reproducing as a female significantly exceeds that of reproducing as a male, a system of labile sexual expression that favors maleness in times of stress should have survival value.¹ Such a system would be particularly advantageous in environments that are highly variable from year-to-year (as in deserts) or that have

favorable and unfavorable patches intermixed at such a scale that each seed has a roughly equal chance of falling into good or poor habitat (as along streams in arid regions). Individuals capable of responding to such environmental patchiness with endogenous hormonal levels capable of inducing the sex that could most successfully pass on genes to the next generation from a given patch should be favored.² As we will show in succeeding paragraphs,

Several studies have demonstrated that the female sexual functions do indeed place females of several species under greater stress than males. For example, Lysova and Khizhnyak (1975) have shown that males and females of *Acer negundo* and *Fraxinus lanceolata* on similar sites grow at equal rates until the females matured the first seed crop after which females grew slower than males. The differential growth rate of males and females seemed to increasingly diverge with the number of seed crops produced. Stark (1970) has shown that females of *Atriplex hymenelytra* (Torr.) S. Wats, are often under greater water stress than males on common sites. Hikmat et al. (1972) have shown similar results for male and female plants of the desert shrub, *Simmondsia chinensis* (Link) C.K. Schneid. Freeman and McArthur (in manuscript) have likewise shown that fruiting females of three dioecious species *(Atriplex canescens, A. confertifolia* and *A. tridentata)* are under greater water stress than associated males

Several research groups working in widely separated geographical zones and dealing with diverse taxa have independently shown that the dioecious species they investigated) exhibit partial niche differentiation, that is the sexes were found to segregate along environment gradients. Spatial segregation of males and females of dioecious species has been reported for the following families: Myricaceae (Davey and Gibson, 1917), Orchidaceae (Dodson 1962; Gregg i973), Aceraceae and Oleaceae (Lysova and Khizhnyak 1975). Rosaceae (Richards 1975), Chenopodiaceae, Ephedraceae, (Freeman et al. 1976 and Waser 1980, in manuscript). Poaceae, and Ranunculaceae (Freeman et al. 1976), Buxaceae (Cole I979; Waser 1980, in manuscript). All of the foregoing workers have reported a similar distribution of males and females in space (i.e., males are more commonly encountered in harsh environments, while females are most often found in favorable sites), although Bawa and Opler (1977) could not detect a segregation of male and female trees along a moisture gradient in Costa Rica. It thus appears that some species in all the foregoing taxa do perceive their environment as patchy with respect to the success of males and females on favorable and marginal sites

some evidence already exists to support the hypothesis that natural selection has equipped at least some plants with environmental monitoring systems that permit the individual to adjust sexual expression and thus increase the likelihood that genetic material will be transmitted to another generation.

Catarino (1964, cited in Heslop-Harrison, 1972) showed that applications of cytokinins to *Kalanchoe crenulata* resulted in the regression of stamens and the enlargement of the ovary. Negi and Olmo (1966, 1972) were able to feminize male *Vitis vinifera* with exogeneous cytokinins. Moreover, when Skene (1972) grew the same clone of *Vitis vinifera* on three different root stocks, he found that the differences in fruit yield were correlated with differences in the cytokinin concentration of the sap. More recently, Louis and Durand (1978) have demonstrated that the male sterile gene of *Mercurialis annua* is the gene(s) controlling production of cytokinins. Chailakhyan (1979) has recently shown that the ratio of cytokinin to gibberillin controls the sex of *Spinacia oleracea* L. and *Cannabis sativa* individuals. When gibberillins exceed cytokinins, the individual is most likely to mature as a male: if the reverse is true, most individuals develop as females. In addition to cytokinins and gibberillins, auxin, ethylene and abscisic acid have each been implicated in control of sexual expression (see Table 2 and Heslop-Harrison, 1972, Friedlander et al. 1977 and Vince-Prue 1975 for further discussion and references).

There is good evidence that environment has a pronounced effect on hormonal levels. For example, under water stress there is reduced transport of cytokinins from the root (the site of biosynthesis) to the shoot (Itai and Vaadia 1965, 1970). Increases in salinity also reduce the transport of cytokinins (Itai et al. 1968, 1973). In addition, drought brings about an increase in the levels of abscisic acid in the shoot (Kaldeway et al. 1974). The reduction in cytokinin would be expected to bring about a shift towards maleness (Table 2, and Chailakhyan 1979). Shifts towards maleness on sites with dry soils have been reported by Pickett (1915), Freeman et al. (1976) Freeman and McArthur (in manuscript) and McArthur and Freeman (in manuscript). Drought is not the only environmental condition which influences the levels of various hormones: flooding (Burrows and Carr 1969), heat (Itai and Ben-Zioni 1974; Itai et al. (1973) and salinity (Itai et al. 1968, 1973) all influence phytohormone levels. Thus, it appears possible that external environment can modify the sexual expression of individual plants through alteration of endogenous hormone levels. This possibility has broad implications and merits careful evaluation by plant physiologists and developmental morphologists.

Recent works by Freeman and Kalled (unpublished data) demonstrates the role of water stress on sex ratio of spinach populations. They transplanted spinach seedlings (from a common seed source and grown in a common environment) into wet and dry soil environments. The wet environment received $5 \times$ the water applied to the dry environment. They obtained ratios of 82 females to 42 males in wet environment and 46 females to 79 males in dry environment. Since only 3 plants died in each treatment and all plants flowered, the results are best explained by the hypothesis of labile sexual expression.

Despite numerous reports discussing the effect of environment on the sexual state of unisexual individuals and production of hermaphroditic offspring by unisexual parents, such phenomena are rarely city by either geneticists or evolutionists. However, Vasek (1968) and Freeman and McArthur (in manuscript) have noted the frequency of sex change in nature. In his studies on Juniper, Vasek found that 7.3% of the individuals of *J. australis* and 24.5% of *J. osteosperma* individuals changed sex over a 2 to 5 year period. In their study Freeman and McArthur marked over 1,400

individuals representing seven species growing in natural populations in Utah. They found that 9% of the *Atriplex canescens,* 13% of the *A. confertifolia,* 22% of the *A. corrugata,* 11.5% of the *A. cuneata,* 30% of the *A. lentiformis,* 20% of the *A. tridentata* and 2.5% of the *Ephedra viridis* individuals changed sex between 1978 and 1979. Hermaphrodites were observed in all species, and hermaphrodites displayed a larger percentage change in sexual expression between the two years than was recorded for either males or females.

In other recent work, McArthur (1977), and McArthur and Freeman (in manuscript) have demonstrated that environmental factors are strongly correlated with the sexual expression of individuals of the subdioecious species *Atriplex canescens* (Pursh) Nutt., to alter their sexual state, They document sex switching following three environmental stresses: 1) an unusually cold winter, 2) drought and 3) heavy seed set by females. During the flowering period following each of the stressing events, there was a pronounced shift towards maleness in the population of over 600 individuals. When each stressing condition was lifted, the proportion of females in the population increased. Over 40% of the plants involved in the study changed their sexual state during the seven years of observation. Over 200 plants displayed the monoecious state during some period of the study, but only one individual remained monoecious in all seven years (McArthur and Freeman, in manuscript). Zuk (1970) has also observed that monoecious individuals of *Urtica dioica* L. may become dioecious and dioecious individuals may become monoecious. He was able to demonstrate that there was no apparent genetic distinction between monoecious and dioecious individuals. Since few researchers have examined the same individual plant year-after-year, it is impossible to determine whether the hermaphrodites reported by most observers maintained that condition throughout their life. The results of Zuk (1970), McArthur (1977) and McArthur and Freeman (in manuscript) have prompted us to include species in Table 1 that are normally regarded as dioecious but which occasionally produce hermaphroditic offspring.

In the remainder of this paper, we will present a hypothesis concerning conditions under which individuals should be expected to alter their sexual expression. We also review ecological conditions under which individuals are known to have displayed different sexual states. Finally, we will suggest new avenues of research.

A Simple Model

A body of theory has been developed recently concerning the conditions under which natural selection will favor the ability of an individual to allocate or reallocate resources to the male or female sex function (Charnov and Bull 1977). The theory applies to plants in the following way.

Suppose a dioecious species with labile sex inhabits an environment consisting of two kinds of patches (call them wet and dry). Assume a wet-patch individual reproducting as a female will produce w_2 as many seeds as a dry-patch individual reproducing as a female. Thus, w_2 is the relative fitness of the female function in the wet versus the dry patch. Likewise, let w_1 be the average number of ovules a wet-patch male will fertilize relative to a dry-patch individual reproducing as a male. Finally, let P be the proportion of the environment made up of dry patches. P could also be considered the proportion of seeds which fall in dry patches (or if mortality differs among patches, P would be the proportion of the reproductives which are in dry patches). Let r_1 equal the probability that an individual in the dry patch becomes a male

 $(r_2$ is the same for wet patches). Our question now becomes: What r_2 and r_2 probabilities form an Evolutionarily Stable Strate- 0.5 gy (ESS) (Maynard-Smith 1976)? We seek an r_1 and r_2 such that any mutant individuals, which alter their values to \hat{r}_1 and \hat{r}_2 , are selected against. Using a population genetic argument based on the assumption of random mating and random dispersal, Char- $1-r_1$ nov 1979a; also see Charnov, et al. 1978 has shown that mutants are selected against providing the following inequality is satisfied:

$$
(2 \geq) \frac{P\hat{r}_1 + (1 - P)\hat{r}_2 w_1}{P r_1 + (1 - P)r_2 w_1} + \frac{P(1 - \hat{r}_1) + (1 - P)(1 - \hat{r}_2) w_2}{P(1 - r_1) + (1 - P)(1 - r_2) w_2}.
$$

The game is now to find r_1 and r_2 such that the inequality is satisfied for all possible \hat{r}_1 and \hat{r}_2 .

By considering (1) to be a function of \hat{r}_1 and \hat{r}_2 (for fixed r_1 and r_2) and taking derivatives with respect to \hat{r}_1 and \hat{r}_2 , it is easy to show that the r_1 and r_2 we seek are those which maximize the *product of the male fitness times the female fitness* or:

$$
(Pr1 + (1 - P)r2W1) \times (P(1 - r1) + (1 - P)(1 - r2)W2).
$$
\n(2)

If $w_2 > w_1$ then *an individual gains more in terms of female reproductive success when in a wet patch.* The solution to this maximization is as follows:

$$
1 - r_1 = \begin{bmatrix} \frac{1}{2} \left(1 - w_2 \frac{1 - P}{P} \right) & w_2 > \frac{1 - P}{P} \\ 0 & w_2 < \frac{1 - P}{P} \end{bmatrix}
$$
 (3)

$$
r_2 = \begin{bmatrix} \frac{1}{2} \left(1 - 1/w_1 \times \frac{P}{1 - P} \right) & \frac{P}{1 - P} < w_1 \\ 0 & \frac{P}{1 - P} > w_1 \end{bmatrix}
$$
 (3)

These results are shown in Fig. 1. This solution makes at least two important predictions: 1) the sex ratio in the wet patch is always biased towards females, while that in the dry goes toward males and 2), the bias strongly depends upon the fraction of the environment that is made up of each kind of patch and so may change from time-to-time or place-to-place. Thus, the expression of sex is a relative character. It is not possible to answer whether an individual in the wet patch should be a male or female, or the probability of one or the other, without knowing what the entire breeding environment is like (i.e., what is P ?).

The model proposed above is simple, but we believe that it captures a basic point – that selection can favor "choice of sex" providing the environment is patchy relative to the individual reproductive-success value of being a male or a female (Charnov and Bull 1977).

There are at least two other ways in which an environment can be "patchy" relative to the value of being a male or a female. 1) Reproductive success could be dependent upon age or, more likely, some measure of size that is loosely correlated with age. This may give rise to sequential hermaphroditism or sex change, a condition in which an individual reproduces early in life as one sex and then changes to the other for the remainder of its life. 2) The reproductive success of an individual could be dependent upon the sexual state of neighboring individuals. Such social effects have been suggested for higher plants (Lloyd 1974a, Charnov and Bull 1977) and are extremely well documented for lower forms (Van der Ende 1976, Charnov et al. 1978).

Ghiselin (1969) suggested that natural selection should favor sequential hermaphroditism over fixed sex when an individual's

Fig. 1. Patchy environment sex ratios. a As the ratio of wet to dry patches increases, the proportion of individuals in the dry patch reproducing as females declines from 0.5 to 0. b As the ratio of dry to wet patches increases, the proportion of individuals in the wet patch reproducing as males declines from 0.5 to $0.$ P is the proportion of the environment made up of dry patches; r_1 is the probability that an individual in a dry patch becomes male, r_2 is the same for wet patches. W_1 is the relative fitness of wet patch male to a dry patch male, and w_2 is the relative fitness of a wet patch female compared to a dry patch female

reproductive success as a male or a female is closely related to age or size, and where the relationship is different for each sex. For example, if large size increases the egg output of a female, but does not aid males in competition to fertilize eggs, selection would favor genes which cause an individual to operate as a male while small, but as a female at a later age or larger size. The reverse order of sex change would, of course, be favored if large size or older age were relatively more important for male reproductive success. The genetic theory of sex change which specifies the ESS, sex order and time of change, has been developed by several people (Warner et al. 1975; Leigh et al. 1976; Charnov 1979 a, b) and applied to reef fish (Warner et al. 1975) and Pandalid shrimp (Charnov 1979b).

Warner et al. (1975) noted in their studies of the blue headed wrasse that individuals changed from female to male when they became large enough to defend a breeding territory. The equations used by Warner et al. (1975) and Leigh et al. (1976) dealt with the age of individuals. The age criterion was used for convenience, but the salient parameter was size. The individual changed sex when it could defend a given resource or territory. Sequential hermaphroditism in plants is probably superimposed upon environmental sex determination (our patchy environment model) since plant size is rarely a simple function of age.

Some Observations on Sex Change

Individuals of *Arisaema triphyllum* (Shaffner 1925 a, b) **, A. japonicus* (Maekawa 1929), *Eucornmia ulmoides, Castilloa elastica* (Dzhaparidze 1963), *Ilex opaca* Ait. (Clark and Orton, 1967) and *Metasequoia glyptostrobodies* (H.G. Baker personal communication) normally produce male flowers first and female flowers only after a delay of several seasons. The percentage of female flowers apparently increases with age. Moreover, sex and size of the pseudocorm (a storage structure) are related in several species of orchid (Gregg 1973). Individuals with large pseudocorms are generally female, while the male's pseudocorms are smaller. Gregg's results are suggestive of sequential hermaphroditism, since the orchid species considered exhibit sex change (Gregg 1973). We have observed a similar correlation between size and sex in *Atriplex confertifolia* and (Torr. & Frem.) S. Wars *A. canescens* (Pursh) Nutt. (i.e., the smaller individuals are usually male). Again this is particularly suggestive of sequential hermaphroditism inasmuch as individuals of both species are known to change sex (McArthur 1977; Freeman and McArthur, in manuscript).

There are, of course, substantial differences between the sequential hermaphroditism exhibited by the blue headed wrasse (Warner et al. 1975) and that suspected for plants. The fish becomes male for life when it reaches a given size and defends a breeding territory that is a relatively stable resource. In contrast, the resources important to plants may fluctuate strongly and frequently, particularly with respect to water in arid and semiarid environments. If sexual expression in plants is indeed correlated with the resource state, it would be advantageous for the sexual expression to fluctuate in synchrony with variable environments as has been observed in *Atriplex eanescens* (McArthur 1977 and McArthur and Freeman, in manuscript, and Freeman and McArthur in manuscript see also Barker et al. in manuscript).

Schaffner (1922, 1925a, 1926) and others listed below have shown that the sexual expression of individuals of some dioecious and sequential hermaphroditic species is related to their physiological state. Schaffner found that removal of a portion of the corm of a female of *Arisaema triphyllum* caused the plant to exhibit one of three sexual states when it flowered again depending on the amount of corm removed. The plant assumed the monoecious habit if a relatively small portion of the corm was removed; removal of a greater portion resulted in the production of male flowers only, and removal of a still greater part of the corm resulted in nonflowering. As the plant recovered, it passed though the stages in reverse order (i.e., if a sufficient amount of corm was removed to cause a female to become male, the plant produced male flowers first, then both male and female, and finally only female flowers).

Other cases are known in which stressed plants revert to the male state. McArthur (1977) found that, in the flowering season following an unusually severe winter, 86 females of a population of about 600 *Atriplex canescens* individuals became either male or monoecious, 10 monoecious individuals became male, and only 11 male and monoecious individuals became female. Obviously, stress triggered a shift toward maleness. In the more normal year which followed, there was a shift back towards femaleness. Frost damage has been reported to trigger a similar shift from femaleness to maleness in cycads (Menniger 1967). As previously noted, McArthur and Freeman (in manuscript) have shown significant retreat from the female habit by saltbush plants stressed by drought and heavy seed set. Barker et al. (in manuscript) have made similar observations for the polygamous tree, Acer grandi*dentatum.* Minina (1952) has shown that, under conditions of low soil moisture or low relative humidity, cucumbers have only male flowers. She also reports that the female sex organs of wheat are more susceptible to water stress than are the male organs. Schaffner (1922) has shown that well-fertilized plants of *Arisaema triphyllum* tend to be female. This fact led him to conclude that the burden of seed production is commonly borne by the most vigorous individuals. Other authors (Giard 1898; Davey and Gibson 1917; Mukerji 1936; Dzhaparidize 1963; Brubaker 1969; Gregg 1973) have come to a similar conclusion.

Data also exist which suggest that environmental variables other than available moisture, nutrients, and storage reserves can induce sex change. Dodson (1962) and Gregg (1973) have shown that sex change is related to light intensity in several orchid species that are sequentially hermaphroditic or dioecious. Shaded plants which are male often become monoecious or female when placed in the sun. Likewise, females often become monoecious or male when placed in the shade. Males in nature are normally most abundant in the shade and females in the sun. Thus, the orchids show a differential sex ratio along a gradient of light intensity. Dodson's (1962) and Gregg's (1973) work suggests that other dioecious species of tropical forests should be investigated for differential sex ratios along strong light gradients.

Most botanical references that we have consulted treat sex change as an unusual event of intellectual interest only. However, the data of McArthur (1977), McArthur and Freeman (in manuscript), Freeman and McArthur (in manuscript), Gregg (1973), Vasek (1968), Schaffner (1922, 1925a, 1926), and Pickett (1915) suggest that the phenomenon must be considered to have great ecological importance. Unfortunately, only the data of McArthur (1977), McArthur and Freeman (in manuscript) and Freeman and McArthur (in manuscript) are based upon large populations in field situations. Clearly similar studies with other species are needed before the broad ecological and evolutionary significance of sex change among individuals of dioecious and sequentially hermaphroditic plants can be fully understood.

Conclusion

Many researchers have shown that sexual expression can be changed in numerous dioecious and sequentially hermaphroditic species. Even some dioecous species reported to have sex chromosomes have the potential of producing monoecious offspring and/ or of changing sex. We have shown that dioecious individuals with the capability of sex change will have a selective advantage over individuals lacking this ability in patchy or changing environments. We suggest that variable sexual expression may be common among species that produce unisexual flowers and that such species should possess a reproductive advantage in environments that are heterogeneous in time and/or space.

References

See also Schaffner 1918, 1919, 1921, 1922, 1923a, b, c, 1925, 1926, 1927, 1928, 1930, 1931, 1933 and 1935

Allen CE (1940) The genotypic basis of sex expression in angiosperms. Bot Rev 6:227-300

Bailey LH, Bailey EZ (1976) Hortus third. MacMillian New York

- Barker P, Freeman DC, Harper KT Sexual flexibility in *Acer grandidentatum.* In manuscript
- Bawa KW, Opler PA (1977) Spatial relationships between staminate and pistillate plants of dioecious tropical forest trees. Evol 31:64-68
- Bemis WP, Wilson GB (1953) A new hypothesis explaining the genetics of sex determination in *Spinacia oleracea* LJ Heredity 44:91-95
- Blackburn KB (1938) On the occurrence of a hermaphrodite plant of *Ernpetrum nigrum L.* Jour Bot 76:306-307
- Bosse GG (1935) Iskusstvennoe izmenenie pola u eikommii (Artificial conversion of sex in *Eucommia*). Soretskie Subtropiki 7:64-68
- Brantley BB, Warren GF (1960) Sex expression and growth in muskmelon. Plant Physiol 35:741-745
- Brubaker MM (1969) Feminine mystique. Orchidata 8:130-133
- Burns GW (I969) The science of genetics: An introduction to Heredity. Macmillan Co New York p 399
- Burrows WJ, Carr DJ (1969). Effects of flooding the root system of sunflower plants on the cytokinin content in xylem sap. Physiol Plant 22 : 1105-1112
- Catarino FM (1964) Some effects of kinetin on sex expression in *Bryophyllum crenaturn* Bak. Port Acta Biol, Ser A8 : 267-284
- Chailakhyan MKh (1979) Genetic and hormonal regulation of growth, flowering, and sex expression in plants. American Journal of Botany 66 : 717-736
- Charnov EL (1979a) The genetical evolution of patterns of sexuality: Darwinian fitness. Amer Natur 113 : 465-480
- Charnov EL (1979b) Natural selection and sex change in Pandalid shrimp: test of a life history theory. Amer Natur 113:715-734
- Charnov EL, Bull J (1977) When is sex environmentally determined? Nature 266 : 828-830
- Charnov EL, Gotshall DW, Robinson JG (1978) Sex ratio: adaptive response to population fluctuations in Pandalid shrimp. Science 200 : 200-206
- Clark RB, Orton ER Jr (1967) Sex ratio in *Ilex opaea* Air. Hortscience 2:115
- Cole SL (1979) Aberrant sex ratios in Joyoba associated with environmental factors. Desert Plants $1(1):8-11$
- Correns C (1928) Bestimmung, Vererbung und Verteilung des Geschlechtes bei den höheren Pflanzen
- Davey AJ, Gibson CM (1917) Note on the distribution of the sexes in *Myrica gale.* New Phytol 16:147
- DeJong AW, Bruinsma J (1974a) Pistil development in *Cleome* flowers: III. Effects of growth-regulating substances on flower buds of *Cleome iberidella* Wellw. ex Oliv. Z Pflanzenphysiol 73:142-151
- DeJong AW, Bruinsma J (1974b) Pistil development in *Cleome* flowers: IV. Effects of growth-regulating substances on female abortion in *Cleome spinosa* Jacq. Z Pflanzenphysiol 73:152-159
- Dodson CH (1962) Pollination and variation in the subtribe Catasetinae (Orchidaceae). Annals of the Missouri Botanical Garden 49:35-57
- Dzhaparidze LI (1963) Sex in plants. Akkademiya Nauk Gruzinskoi SSR. Institut Botaniki. Translated from Russian for NSF
- Erlanson EW, Hermann FJ (1927) The morphology and cytology of perfect flowers in *Populus tremuloides* Michx. Papers Mich Acad Sci Arts Lett 8:97-110
- Frankel R, Galun E (1977) Pollination mechanisms, reproduction and plant breeding. Springer-Verlag New York
- Friedlander M, Atson D, Galun E (1977). Sexual Differentiation in cucumber: The effects of abscisic acid and other growth regulators on various sex genotypes. Plant and Cell Physiol 18:261-269
- Freeman DC, McArthur ED, Harper KT, Blauer C (In press) The influence of environment on the floral sex ratio of monoecious plant. Evolution
- Freeman DC, Klikoff LG, Harper KT (1976) Differential resource utilization by the sexes of dioecious plants, Science 193 : 597-599
- Freeman DC, McArthur ED Sex reversal by individuals in natural populations of seven dioecious plant species of the eastern Great Basin. In manuscript
- Fuller HJ, Ritchie DD (1967) General Botany. Barnes and Noble Inc. New York, p 232
- Fuller HJ, Carothers ZB, Payne WW, Balbach MK (1972) The plant world. Holt, Rinehart and Winston, Inc New York, p 553
- Gabe DR (1939) Inheritance of sex in *Mercurialis annua* in relation to cytoplasmatic theory of sex-inheritance. Compt Rend (Doklady) Acad Sci S.S.R. ns 23:477-480
- Garber ED (1972) Cytogenetics: an introduction. McGraw-Hill New York, p 259
- Gardner EJ (1975) Principles of Genetics. John Wiley and Sons Inc New York, p 622
- Ghiselin M (1969) Evolution of hermaphroditism among animals. Quarterly Rev of Biol 44:108-189
- Grant Verne (1975) Genetics of Flowering Plants. Columbia University Press New York and London, p 514
- Gregg KB (1973) Studies on the control of sex expression in the genera *Cycnoches* and *Catasetum,* subtribe Catasetinae, Orchidaceae. Unpublished PhD dissertation, University of Miami, Coral Gables, Florida
- Giard A (1898) Les variations de la sexualite chez les vegetaux. CR Soc Biol Paris 10:730
- Hall WC (1949) Effects of photoperiod and nitrogen supply on growth and reproduction in the gherkin. Plant Physiol 24:753
- Halevy AH, Rudich Y (1967) Modification of sex expression in muskmelon by treatment with the growth retardant B995. PhysioI Plant 20:1052-1058
- Haney Alan W (1978) Plants and Life. Macmillan Publishing Co. Inc New York, p 486
- Hartmann M (1956) "Die Sexualität", Vols 2. Fischer Stuttgart
- Hedrick VP, Anthony RD (1915) Inheritance of certain characters of grapes. J Agr Res 4:315-330
- Herich RK (I956) Problemu pohlavnej determinacie a difcrenciacie restlin. I. Posobenie boru na priebeh poblavney diferenciacie konopi *(Cannabis sativa L.)* Acta Fac Rerum Natur Univ Comenianae, Bot 1:419-425
- Heslop-Harrison J (1956) Auxin and sexuality in *Cannabis sativa.* Physiol Plant 9:588-597
- Heslop-Harrison J (1957) The experimental modification of sex expression in flowering pIants. Biol Rev 32:38-90
- Heslop-Harrison J (1964) The control of flower differentiation and sex expression. Pages 649-664 *in* JP Nitsch, ed. Regulateurs naturels de la croissance vegetale. CNRS Paris
- Heslop-Harrison J (1972) Sexuality of angiosperms. Page 113-289 *in* FC Steward, Ed. Physiology of development: from seeds to sexuality. Academic Press NY
- Heslop-Harrison JW (1924) Sex in the Salicacaceae and its modification by eriophyid mites and other influences. Brit J Exp Biol 1:445-472

Hirata K (1924) Sex reversal in hemp. J Soc Agr and For 16:145-168

- Hikmat AAA, Strain BR, Mooney HA (1972) The physiological ecology of diverse populations of the desert shrubs, *Simmondsia chinensis.* J Ecol 60:41-57
- Hofmeyr JDJ (1938) Genetical studies of *Carica papaya* L. I-II. Union S. Africa, Dept Agr and Forestry Sci Bull 187:5~64
- Hofmeyr JDJ (1939) Some suggestions on the mechanism of sex determination in *Carica papaya.* S African J Sci 36:288-290
- Huhnke W, Jordan C, Neuer H, von Sengbusch R (1950) Grundlagen für die Zfichtung eines monozischen Hanfes. Z Pflanzenzucht 29:55-75
- Ikeno S (1937) Zur Kenntnis des Erbverhaltens einer gynodiozischen Pflanze, *Petasitesjaponicus* Miq. Cytologia Fugii Jub vol 888-896
- Itai C, Vaadia Y (1965) Kinetin-like activity in root exudate of waterstressed sunflower plants. Physiologia PI 18 : 941-944
- Itai C, Richmond A, Vaadia Y (1968) The role of root cytokinins during water and salinity stress. Israel J of Botany 17:187-195
- Itai C, Vaadia Y (1970) Cytokinin activity in water-stressed shoots. Plant Physiol 47 : 87-90
- Itai C, Ben-Zioni A (1974) Regulations of plant response to high temperature. R Soc N Z Bull 12:477-482
- Itai C, Ben-Zioni A, Ordin L (1973) Correlative changes in endogenous hormone levels and shoot growth induced by short heat treatments to the root. Physiol Plant 29:355-360
- Kaldeway H, Ginkel V, Wawczyniak G (1974) Auxin transport and water-stress in pea. *Pisum satvum L.* Ber Dtsch Bot Ges 87(3) 563-576
- Keen R, Chadwick LD (1954) Warn propagators to watch for sex reversal in *Taxus.* American Nurseryman 100(6):13-14
- Kuhn E (t939) Selbstbestaubungen subdiocischer Blutenpflanzen, ein neuer Beweis ffir die Genetische Theorie der Geschlechtsbestimmung. Planta 30:457-470
- Lange AH (1961) Factors affecting sex changes in the flowers of *Carica papaya.* Proc Am Soc Hort Sci 77:252-264
- Leigh EG Jr, Charnov EL, Warner R (1976) Sex ratio, sex change, and natural selection. Proc Natl Acad Sci 73 10 3656-3660
- Lloyd DG (1974) Female predominant sex ratios in angiosperms. Heredity 32 : 35-44
- Louis JP, Durand B (1978) Studies with the dioecious angiosperm, *Mercurialis annua* L. $2n = 16$. Correlation between genic and cytoplasmic male sterility, sex segregation and feminizing hormones (cytokinins). Molec Gen Genet 165:309-322
- Lysova NV, Khizhnyak NI (1975) Sex differences in trees in the dry steppe. The Soviet J of Ecol 6:522-527
- Matzke EB (1938) Inflorescence patterns and sexual expression in *Begonia semperflorens.* Amer J Bot 25:465-478
- Maynard-Smith S (1976) Evolution and the theory of games. Am Sci 64:41-45
- McArthur ED (1977) Environmentally induced changes of sex expression in *Atriplex canescens.* Heredity 38 : 97-103
- McArthnr ED, Freeman DC Sex reversal in *Atriplex canescens.* In manuscript
- McPhee HC (1924) The influence of environment on sex in hemp, *Cannabis sativa.* L, J Agr Res 28:1067-1080
- Menninger Edwin A (1967) Fantastic trees. Viking Press. New York
- Minina EG, Tylkina JG (1947) Physiological study of the effect of gases upon sex differentiation in plants. CR Acad Sci SSSR (Doklady), 55:165
- Minina EG (1952) Sex shifts in plants induced by environmental factors. Moscow Acad Sci. USSR Publishing House, Moscow, (Translated from the Russian)
- Mukerji SK (1936) Contributions to the auteocology of *Mercurialis perennis.* J Ecol 24:38
- Murneek AE (1927) Physiology of reproduction in horticultural plants. II. The physiological basis of intermittent sterility with special reference to the spider flower. Miss, Agr Exp Sta, Res Bull 106
- Nakajima G (1942) Cytological studies in some flowering plants, with special reference to the sex chromosomes. Cytologia 12 : 262-270
- Negi SS, Olmo HP (1966) Sex modification in grapes by cytokinin. Sci 152:1624-1625
- Negi SS, Olmo HP (1972) Certain embryological and biochemical aspects of cytokinin. S.D. 8339 in converting sex of a male *Vitis vinifera* sylvestris. Am J Bot 59:851-857
- Negodi G (1929) Ricerche sulla distribuzione e transmissione dei sessi in *"'Utrica caudata "'* Bahl. Nuovo Giorn Bot ltal 36:60-126
- Negodi G (1931) Ricerche sulla distribuzione e transmissione dei sessi in "Utrica cannabina L" Annali Bot 19:264-277
- Negru AM (1936) Variabilitat und Vererbung des Geschlechts bei der Rebe. Gartenbauwiss 10:215-231
- Newton WCF (1931) Genetical experiments with *Silene olites* and related species. J Genet 24:109-120
- Nilsson HN (1918) Experimentelle Studien fiber Variabilitat, Spaltung, Artbildung und Evolution in der Gattung *Salix.* Lnnds Univ Arsskv Nof Avd 2:14 No 28
- Pickett FL (1915) A contribution to our knowledge of *Arisaema triphyl*lum. Torrey Botanical Club Memoirs 16:1-55
- Quagliotti L (1972) Relazioni tra concimazione azotata, habitus vegetativo ed espressione sessuale nello spinaco "riccio d' asti" sementi. Elette; 18(2):5-19
- Raino AJ (1927) Über die Intersexualitat bei der Gattung Salix. Ann Soc Zoo-Bot Fenn vanano 5:165-275
- Richards AJ (1975) Notes on the sex and age of *Potentilla fruticosa* L. in upper Teasdale. Trans of the Natural History Soc of Northumbria 42(3): 85-92
- Rick CM, Hanna GC (1943) Determination of sex in *Asparagus offininale* L Am J Botany 30:711-714
- Rudich J, Halevy AH, Kedar N (1969) Increase in femaleness of three cucurbits by treatment with ethrel, an ethylene-releasing compound. Planta 86:69-76
- Sansome FW (1938) Sex determination in *Silene otites* and related species. J Genet 35:387-396
- Schaffner JH (1918) The expression of sexual dimorphism in heterosporous sporophytes. Ohio J Sci 18:101-125
- Schaffner JH (1919) Dioeciousness in *Thalictrum dasycarpum.* Ohio J Sci 20 : 25
- Schaffner JH (1921) Influence of environment on sexual expression in hemp. Bot Gaz 71:197-219
- Schaffner JH (1922) Control of sexual state in *Arisaema triphyllun* and *A. dracontium.* Am J Bot 9:72-78
- Schaffner JH (1923a) The influence of relative length of daylight on the reversal of sex in hemp. Ecol 4:323-334
- Schaffner JH (1923b) Sex reversal in the Japanese hop. Bull Torrey Bot Cl 50:73-79
- Schaffner JH (1923c) Observations on the sexual state of various plants. Ohio J Sci 23:149
- Schaffner JH (1925a) Experiments with various plants to produce change of sex in the individual. Bull Torrey Bot C1 52:35-47
- Schaffner JH (1925b) Sex determination and sex differentiation in the higher plants. Amer Nat 59:115-127
- Schaffner JH (1925c) The influence of the substratum on the percentage of sex reversal in winter grown hemp. Ohio J Sci 25:172-176
- Schaffner JH (1926) Siamese twins of Arisaema triphyllum of opposite sex experimentally induced. Ohio J Sci 26:276-280
- Schaffner JH (1927) Control of sex reversal in the tassel of Indian corn. Bot Gaz 84:440-449
- Schaffner JH (1928) Further experiments in repeated rejuvenations in hemp and their bearing on the general problem of sex. Am J Bot 15:77-85
- Schaffner JH (1930) Sex reversal and the experimental production of neutral tassels in *Zea mays.* Bot Gaz 90:279-298
- Schaffner JH (1931) The fluctuation curve of sex reversal in staminate hemp plants induced by photoperiodicity. Am J Bot 18:424-430
- Schaffner JH (1933) The production of vestigal and sterile sex organs through sex reversal and neutral sex states. Bull Torrey Bot Cl 60:89-97
- Schaffner JH (1935) Observations and experiments on sex in plants. Bull Torrey Bot C1 62:387-400
- Sengbusch R yon (1952) Ein weiterer Beitrag zur Vererbung des Geschlechts bei Hanf als Grundlage für die Züchtung eines monozischen Hanfes. Z Pflanzenzucht 31:319-338
- Shifriss O (1961) Gibberellins as sex regulator in *Ricinus eommunis.* Science 133:2061-2062
- Schull GH (1911) Reversible sex mutants in *Lychnis dioica.* Bot. Gaz 52:329-368
- Smith BW (1963) The mechanism of sex determination in Rumex hastatulus. Genetics 48 : 1265-1288
- Srb AM, Owen RD, Edgar RS (1965) General Genetics. WH Freeman and Company, San Francisco. p 557
- Stark N (1970) Water balance of some warm desert plants in a wet year. J of Hydrology 10:113-126
- Storey WB (1953) Genetics of the papaya. J Heredity 44:70-78
- Strasburger E (1910) Sexuelle und apogame F ortpflanzung bei Urticaceen. Jahrb Wiss Bot 47:245-288
- Strickberger MW (1976) Genetics. MacMillan Co., New York. p 914
- Thompson AE (1955) Methods of producing first generation hybrid seed in spinach. Cornell Univ Agr Exp Sta Mem 336
- Tothill JC, Knox RB (1968) Reproduction in *Heteropogon contortus* I. Photoperiodic effects on flowering and sex expression. Aust J Agr Res 19 : 869-878
- Tournois J (1911) Anomalies florales du *Houblunjaponais* ed du chanure determinees par des semis hatifs. CR Acad Sci 153:1017-1020
- Tournois J (1912) Influence de la lumiere la floraison du *Houblon japonais* et du chanvre. CR Acad Sci 155:297-300
- Tournois J (1914) Etudes sur la sexualite du *Houblon,* Ann Sci Nat Bot Biol Veg 19:49-191
- Ubisch G yon (1936) Genetic studies on the nature of hermaphroditic plants in *Antennaria dioica* (L.) Gaertn Genetics 21:282-294
- Van der Ende H (1976) Sexual interactions in plants. The role of specific substances in sexual reproduction. Academic Press New York
- Vasek F (1966) The distribution and taxonomy of three western junipers. Brittonia 18 : 350-372
- Vince-Prue D (1975) Photoperiodism in plants. McGraw-Hill New York
- Warner P, Robertson D, Leigh EG Jr (1975) Sex change and sexual selection. Science 190:633-638
- Waser NW Sex ratio variation in populations of two desert perennials. (submitted to Ecology)
- Westergaard M (1958) The mechanism of sex determination in dioecious flowering plants. Adv in Genetics 9:217-281
- Wilson EO, Eisher T, Briggs WR. Dickerson RE, Metzenberg RL, O'Brien RD, Susman M, Boggs WE (1978) Life on Earth. Sinauer Associates, Inc Sunderland, Mass, p 846
- Worsdell WC (1916) The "Principles of Teratology in Plants." vol II Ray Society London
- Yampolsky C (1919) Inheritanace of sex in *MercuriaIis annua.* Am J Bot 6:410-442
- Zhukovskii PM (1940) Botanika (Botany). Second edition, Moskva Sel'khozgiz p 295
- Zuk J (1970) Structure and function of sex chromosomes in *Rumex thyrsiflorus*. Acta Societatis Botanicorum Poloniae. 39:539-564

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