### Examples of Variation Patterns within Species and Genera

CHAPTER II

number of sharply discontinuous and distinct populations or greatly restricting mating and gene interchange has produced a various individuals or populations constituting the pattern, or owing to the more or less free interchange of genes between the and, second, that involving the different species of a genus or whether the presence of isolating mechanisms preventing or pattern is essentially continuous or only partly discontinuous. the distinction between these two levels is that of whether the larger grouping. As is discussed in greater detail in Chapter VI, population systems. These exist on two different levels; first, that within the species EFORE DISCUSSING the individual factors responsible for evolution, it seems desirable to describe some of the patterns of variation which are the outcome of these processes.

#### THE ECOTYPE CONCEPT

emphasized (1936) the fact that differentiation into ecotypes is of wide-ranging Eurasian species, mostly perennial herbs. He has son, in a long series of publications (1922a, 1922b, 1925, 1927 sponse of an ecospecies or species to a particular habitat." Tures of the ecotype. This term was originally defined by Turesson major emphasis has been placed in recent years on the concept nection with adaptation to ecological conditions. For this reason, evolution is that showing certain regularities, particularly in conmuch more likely to be found in common, widespread species (1922a) as "the product arising as a result of the genotypical re-1940 for complete list), has described ecotypes in a large number 1931a, 1936, etc.; see Clausen, Keck, and Hiesey 1940 or Hiesey The type of variation within species which is most important in

> than in rare, local, or endemic ones. In this country, the studies and Miller (1941)? These two questions will be considered in modern zoological systematists like Rensch (1939), Mayr (1942), of polytypic species, or Rassenkreise, as it has been developed by the relation between the ecotype concept in plants and the concept what extent do they form a continuous series? Second, what is aggregates which may be recognized as distinct ecotypes, and to different biotypes of a species grouped into partly discontinuous connection with the ecotype concept. First, to what extent are the exist in most wide-ranging plant species. Two questions arise in like those which Turesson has recognized as ecotypes undoubtedly variation correlated with habitat differences. Groups of biotypes various other workers have demonstrated intraspecific genetic condition in several species of western North America, while of Clausen, Keck, and Hiesey (1940, 1947) have shown a similar

nearly always overlap, and even if they do not, a series could be slight discontinuity in the variation pattern in northern Scandicontinuous type within Pinus sylvestris of Europe, although a comparison of biotypes taken from a relatively small number of out." Faegri (1937) has pointed out that the apparent distinctness arranged so that there could be continuous variation throughabout many of the examples given by Clausen, Keck, and Hiesey widely separated localities. The same comment may be made of ecotypes in many of the species studied by Turesson results from within populations. The ranges of these in different populations are . . . many quantitative characters which vary continuously Davey, and Lang (1936) found that in Plantago maritima "there navia permits the recognition of a separate ecotype or subspecies and Langlet (1936) showed the presence of much variation of a of a more or less continuous series of morphologically and ecowell-marked groups of genetic variants because of the presence ness of ecotypes, other authors have found difficulty in recognizing Keck, and Hiesey (1940) have tended to emphasize the distinctbasis of Langlet's preliminary and incomplete data. Gregor, for the pines of Lapland, as suggested by Turesson (1936) on the logically intermediate populations. Engler (1913), Burger (1941), Although both Turesson (1936) and to a lesser extent Clausen,

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(1940), as was suggested by Turrill (1942b). The validity of Turrill's criticism is evident from a comparison of the discussion of Achillea given in the above-mentioned work with the later, more complete study by the authors of the same group (Clausen, Keck, and Hiesey 1948). There is no doubt that in plants, as in animals, many species may be divided into races or groups of genetic types which are adapted to the different ecological conditions found in different parts of their ranges, and that these subdivisions are separated from each other by partial discontinuities in the variation pattern. But in addition, many widespread species possess a considerable amount of ecotypic, that is, directly adaptive, genetic variation which because of its continuous nature does not permit the recognition of distinct ecotypes.

### ECOTYPIC AND CLINAL VARIATION

For this reason, much of the variation within certain species is best portrayed by the use of the "auxiliary taxonomic principle" defined by Huxley (1938, 1939) as the *cline*, or character gradient. Clines are probably common in plant species, but the ordinary methods of systematics, which deal with combinations of characters and are aimed at detecting character correlations and discontinuities, are not likely to reveal them.

Among the best examples of clines within plant species are

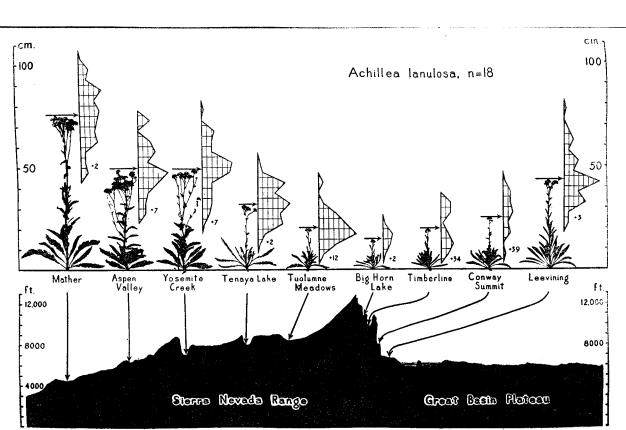


Fig. 5. Representatives of populations of Achillea lanulosa as grown in a uniform garden at Stanford. These originated in the localities shown in the profile below of a transect across east central California at approximately 38° N. latitude.

grama grass (Bouteloua curtipendula) obtained from different

latitudes in the Great Plains. Clinal trends were found by Böcher

(1943, 1944) in Plantago lanceolata and Veronica officinalis. It is

and compared under uniform cultural conditions. In A. lanulosa, the tallest genetic types were from the lowest altitudes, and the decrease in height was more or less continuous with increasing altitude (Fig. 5). Olmstead (1944) found clinal variation in vigor and in reaction to photoperiodism in strains of side-oats

those described by Langlet (1936) in *Pinus sylvestris* for genetic variation in chlorophyll content, length of mature leaves, hardiness, and rapidity of shoot development in the spring. Clausen, Keck, and Hiesey (1948a) found within the ecotypes of *Achillea* 

lanulosa and A. borealis clines for height of plant when grown

found to possess clines for the "physiological" characteristics

likely that most species with a continuous range that includes

more than one latitudinal or altitudinal climatic belt will be

The plants are herbarium specimens, each representing a population of approximately 60 individuals. The frequency diagrams show variation in height

within each population: the horizontal lines separate class intervals of 5 cm according to the marginal scale, and the distance between vertical lines represents two individuals. The numbers to the right of some frequency diagrams indicate the nonflowering plants. The specimens represent plants of average height, and the arrows point to mean heights. From Clausen, Keck, and Hiesey 1948.

adapting them to the conditions prevailing in the different parts of their range.

Clines have also been reported for the frequency of occurrence in a population of a single qualitative character, such as pubescence. Fassett (1942) found that the pubescent form (var. hypomalaca) of Diervilla lonicera in the region north of Lake Huron shows a cline ranging from 80 percent frequency at Espanola, Ontario, to 0 percent at Callander, about 100 miles farther east. The data of McClintock and Epling (1946) indicate that in Teucrium canadense there is a cline for the frequency of plants with glandular hairs on the calyx ranging from a high of 100 percent in the Rocky Mountains southeast to 0 percent in southern Florida.

ecotypes (see p. 52). spots on the leaves. Ecoclines, on the other hand, are clines cally in the same region according to whether the plants are growspike, habit of growth, and length of scape, which vary genetisimilar to the geographic or regional clines described above for two types of clines, topoclines and ecoclines. The former are forms which are the ends of these ecoclines constitute edaphic vailing higher up and on maritime rocks and cliffs. The extreme logged coastal mud or under the better drainage conditions pre-Gregor 1944). They occur in characteristics such as density of related to ecological gradients within a restricted area (see also in the frequency of such characters as pubescence and purple ters, such as the length/width index of bracts and sepals, and other genera and include variation both in quantitative characing under the poor drainage conditions of a salt marsh or water-Gregor (1939), in his studies of Plantago maritima, recognized

Treatment of intraspecific variation by clines has advantages as well as disadvantages over the more usual method of dealing in terms of the character combinations which distinguish races or subspecies. One advantage is that it makes possible the analysis of the individual characters of these combinations and is the first step toward the causal analysis of these differences in terms of selection or any other factor. It also focuses attention on the continuous variation in quantitative characteristics which is present in many wide-ranging species and is of great importance in their adaptation to the environment, but which because of its very

continuity is difficult to use as a basis for classification. One disadvantage of the use of clines is that each individual cline can be recognized only after analysis of extensive samples from many localities. This makes it impossible to describe all the clines existing in a species, so that the method of necessity places particular emphasis on certain individual characters, which may or may not be the most important ones from the standpoint of the biology and evolution of the species as a whole. Furthermore, since selection in populations is based on character combinations (see Chapter IV), too much emphasis on gradients in single characters may serve to confuse rather than clarify the whole picture of variation.

The relation of clines to subspecies depends on the character of the clines. If they are continuous over long distances, and if the extreme types are relatively limited in distribution, as with the clines in *Pinus sylvestris*, then they cannot form the basis for classification. But if a number of clines run parallel and are partly discontinuous, with a steep gradient in some regions and a moderate one or a constant level in others, then the different levels of variation in the characters forming the clines may constitute part of the basis of local races or subspecies. These are the intergroup clines of Huxley (1939).

and seed dispersal. In species occupying an area like the eastern and the size of their population units in terms of cross-breeding show continuous genetic variation. In them the interbreeding Turesson (1936) has pointed out, species with obligate crossspecies, and therefore of more distinct ecotypes. In addition, as of more distinct, easily recognized groups of biotypes within the tinuity of the available habitats will promote the differentiation ticularly prevalent. On the other hand, diversity and discontinuously, continuous or clinal ecotypic variation will be parperature and length of growing season, varies gradually and concharacteristics and where a single set of factors, such as tem-United States, which is comparatively uniform in many climatic tinctness of their ecotypes, depending on the regions they occupy pollinated trees of temperate regions, in which the pollen may pollination, particularly those like pines and other windbe carried through the air for many miles, are most likely to Various species undoubtedly differ from each other in the dis-

ternal, which promote continuity or partial discontinuity in the another and depends on various factors, both external and incan be recognized probably varies greatly from one species to dence justifies the following tentative answer to the first question really conclusive generalizations can be made, the present evi-Although much more experimental work is needed before any always between neighboring individuals, and particularly those gradient. On the other hand, species in which pollination is nearly adaptive) nature is found in nearly all species with a wide ecodistinct ecotypes are likely to be recognized with greater ease. to show more differences between colonies. In them, therefore, are selected from a great store of genetic variants. The chances logical distribution; but the ease with which distinct ecotypes with a relatively high proportion of self-pollination, will tend to exactly the continuity or discontinuity of the environmental closely adapted to their environment and will reflect more or less are therefore particularly favorable that the survivors will be population is relatively large, and the seedlings which survive posed above. Intraspecific variation of an ecotypic (that is, strictly be more uniform within colonies of closely adjacent plants and

From the foregoing discussion, the fact should be clear that clines and ecotypes are not mutually exclusive concepts, but merely express different ways of approaching the same problem. Clinal variation may occur in the characters which determine the nature of adaptation, and therefore form the basis of the ecotypes, or it can also be found in characters of no apparent adaptive value. Correspondingly, ecotypic variation may consist of a series of clines, running either parallel to each other or in opposite directions, or it may have such well-marked discontinuities or be of such an irregular type that constant gradients are not apparent. As Gregor (1939) has emphasized, both these approaches are valuable aids to an understanding of the variation within species.

### ECOTYPES AND SUBSPECIES

There are two differences between the ecotype concept and that of the polytypic species, or *Rassenkreise*. In the first place, subspecies are based primarily on recognizable differences, while ecotypes are distinguished primarily by their reaction to the

of a subspecies." A specific example of this relationship may be species." The relation between ecotypes and subspecies is exsometimes necessary to include more than one ecotype in a subterm)." And on the same page they state specifically that "it is recognize ecotypes in the field or the herbarium is due to the strongly affected by the environment, so that the inability to time of flowering, are clearly among the characteristics most other is subalpine and may be distinguished in garden cultures by dwarf, early-flowering alpine that occurs above 2600 m., while the as follows (1940, p. 43): "It consists of two ecotypes: one is a cited in their discussion of Potentilla glandulosa subsp. nevadensis, limits of one or a group of several ecotypes (an experimental field. Clausen, Keck, and Hiesey consider that (1939, p. 106) logical differences which enable them to be recognized in the environment, and may or may not possess well-marked morphomasking of genetic differences by environmental modification. ences between the two ecotypes of this subspecies, stature and included in one subspecies." In this example the genetic differthe two are not sufficiently distinct to be recognized in the field pressed briefly as follows (Clausen, Keck, and Hiesey 1940, p. 33), "limits of subspecies (a morphologic term) correspond to the only rarely masks racial differences, zoologists do not need a purely such situations in plants probably results from the fact that ecotype. It permits the generalization that the frequency of of those in which a single subspecies contains more than one under uniform conditions. This example is probably typical or in the herbarium with certainty, the two ecotypes are here its taller stature and later flowering. Since the differences between "we consider a morphologically distinguishable ecotype the basis and adaptational concept, while the subspecies is primarily a genetic-ecological term in addition to a primarily systematic one. modification in animals is much less frequent than in plants, and the most important genetic adaptive characteristics are often This effect can be eliminated only by growing the two ecotypes morphological, geographical, and historical one. Gregor (1944) that of subspecies is that the ecotype is primarily an ecological graphically describes the origin of ecotypes as follows, "The paralleled by environmental modifications. Since environmental The second difference between the concept of ecotypes and

possess similar ecotypes. of any ecotypic adaptation." In this species, different subspecies tion of its taxonomic variants appear to be entirely independent through partial isolation from other subspecies. Lawrence (1945) of similar environmental factors, but also to a large extent tive morphological characteristics partly through the influence group of populations with a common origin and a more or less originated independently in response to the same selective forces. "dune ecotype" are therefore aggregates of races which have land ecotype of H. umbellatum. The "cliff ecotype" and the different from those of the dune ecotype found at Sandhammar cliff and inland types occurring in the same region and are very of map) has leaves which in many respects resemble those of the species in the province of Scania, in southern Sweden. Turesson Figure 6 shows the distribution of the different coast types of this also shown by Turesson's studies of Hieracium umbellatum. of the environment on a heterozygous population, the same ecowherever several ecological habitats available to the species occur. are best fitted to survive." Ecotypes correspond with subspecies among the constituents of a population those genotypes which habitat environment is likened to a sieve which sorts out from found that in Deschampsia caespitosa "the evolution and distributo have originated independently from the more widespread intively at Stenshufvud on the east coast and at Kullen or Hofs points out that the dune ecotype found at Torekov (upper left type may originate independently in different localities. This is more, since the ecotype is produced by the direct selective action fields, one of sea cliffs, and one of interior woodlands. Furtherintegral geographic distribution, which has acquired its distinc-The subspecies, on the other hand, is usually conceived of as a (1939) in *Plantago maritima* have undoubtedly arisen repeatedly. The cliff ecotype and the salt marsh ecotype described by Gregor Hallar on the west coast. Each of these cliff and dune races seems bellatum; one typical of shifting sand dunes, one of arenaceous Thus, within a relatively small section of southern Sweden, the same species is expected in a single geographic region logical conditions. But the presence of two or more ecotypes of in so far as different geographical regions possess different eco-(lower right). The same is true of the cliff types found respec-Turesson (1922b) described four ecotypes of Hieracium um-

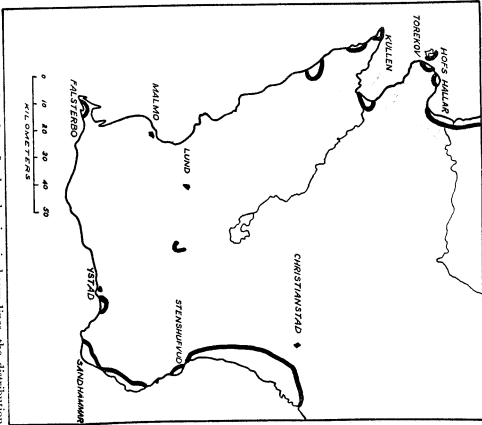


Fig. 6. Map of southern Sweden showing, in heavy lines, the distribution of the maritime ecotype of *Hieracium umbellatum*. Simplified, from Turcsson 1922b.

The ecotype and subspecies have been interpreted by some authors in a relatively broad sense and by others in a relatively narrow one, but this has been true to a considerably greater degree in regard to the ecotype. This is because the concept of what constitutes an ecological difference is largely relative. As Turrill (1946) points out, the ecotypes of Turesson include both regional groupings and small local groupings of biotypes, while in the work of Clausen, Keck, and Hiesey emphasis is placed on regional or

addition to discontinuous variation. Lawrence (1945) presented ecological groupings and the presence of continuous variation in a different sort of ecotype and proposed a system of nomenclature climatic adaptation. Sinskaia (1931a) and Gregor (1939) have another classification of ecotypes, based on the factors responsible for subspecific units which took into account both regional and restricted the term ecotype to local ecological variants within a for their segregation, as follows: units corresponding to the usual subspecies. In other publications (1931, 1942, 1944), Gregor recognized the geographic groupings as (Sinskaia) and topotype (Gregor) — for the geographic-ecological particular geographic region and use another term—climatype

- 1. Climatic ecotype (Turesson 1925) Synonym: climatype (Sinskaia 1928, 1931a)
- 2. Edaphic ecotype (Sinskaia 1928; Gregor 1942)
- 3. Biotic ecotype (Sinskaia 1931a)
- a. Synecotype (Sinskaia 1931a)
- b. Agroecotype (Gregor 1938b)

4. Geographic ecotype

Synonyms: seclusion type (Turesson 1927) geoecotype (Gregor 1931)

method of expressing the actual biological situation agreement as to what constitutes the most convenient and efficient among all scientists interested in the subject and by mutual specific categories will be solved only by impartial discussion tion. Like all other problems of classification, that of the infratype may be considered a special type of edaphic ecotype adapted origin of geoecotypes. As recognized by Sinskaia, the biotic ecoto the type of plant competition existing under human cultivamechanically isolated areas," as postulated by Gregor for the climatic factors and the effects of "chance introduction into ment of a species in any region usually includes both selection by most always have different climates, and the process of establishpossible in most cases because different geographical regions al-The distinction between geographic and climatic ecotypes is im-Some parts of this classification may prove difficult to apply.

VARIATION ON THE LEVEL OF THE SPECIES AND GENUS

In the case of variation patterns involving species - that is,

greatly restrict interbreeding and the exchange of genes-the populations or population systems separated from each other by on the basis of ease of crossing and the fertility of the hybrids of isolating mechanisms separating them from other groups. These species groups according to the degree of development of the geneticists have adopted a series of terms characterizing species or populations which they isolate. For this reason, some plant these barriers are developed and the size and diversity of the nature of the pattern is largely determined by the degree to which physiological or genetic isolating mechanisms which prevent or crossing between members of different cenospecies either is ungenes among themselves to a limited extent through hybridizaconsists of "all the ecospecies so related that they may exchange closely to the usual taxonomic species (Clausen, Keck, and Hiesey first two terms, both developed by Turesson (1922b), are defined terms are the ecospecies, the cenospecies, and the comparium. The freely without loss of fertility or vigor in the offspring" (Clausen, tions or ecotypes "so related that they are able to exchange genes the  $F_1$  and later generations. The ecospecies is a system of populaor it may comprise a whole genus, as in Aquilegia. The comof a section or subgenus, as in Pinus, Quercus, and Ceanothus, may correspond exactly with the ecospecies, but often it consists successful or yields completely sterile hybrids. The cenospecies it approaches the size of a genus, and in some plant groups, such term and includes (Clausen, Keck, and Hiesey 1945a, p. vi) "all parium (Danser 1929) is a more inclusive and strictly genetical limited extent and form at least partially fertile hybrids, but the various ecospecies composing a cenospecies can hybridize to a tion" (Clausen, Keck, and Hiesey 1945a, p. vi). This means that 1939). It is discussed more fully in Chapter VI. The cenospecies Keck, and Hiesey 1945a, p. vi). This unit corresponds most recognized genera. as the grasses and the orchids, it may include a whole series of in isolated instances consist of only one ecospecies, but usually directly or through intermediaries." Like the cenospecies, it may the cenospecies between which hybridization is possible, either

cussed and the factors of evolution which will be reviewed in the remaining chapters of the present volume all apply to the great Although the terms and concepts which have already been disof genetic patterns on the diploid level system may be considered premature rather than altogether inapsystem like that of Camp and Gilly may prove valuable in orientother geographically. Most genera have variation patterns someing the thinking of botanists on problems of evolution; their the genetic bases of variation patterns increases, the adoption of a and not always soundly based on genetic phenomena, so that it where intermediate between these extremes. Camp and Gilly sense of Mayr (1942, Chaps. V and VI); that is, they consist of a treme are genera in which all the species are polytypic in the genera, the species are genetically homogeneous, have relatively and ecotypic diversity contained within the species. In some which isolating barriers are developed and the amount of genetic will be presented which is intended to give an idea of the diversity propriate. In the remainder of this chapter, a series of examples has not been generally accepted. Nevertheless, as knowledge of found in plants, but their terminology has proved cumbersome logically recognizable subspecies or ecotypes which replace each greater or lesser number of partially discontinuous, morphorestricted ranges, and have no close relatives. At the other exfertilizing may differ greatly from each other in the degree to groups of which all the members are diploid, sexual, and crossas is brought out in Chapters V and VIII-XI. Nevertheless, even as self-fertilization, polyploidy, apomixis, and structural hybridity is due to the sporadic or frequent occurrence of such phenomena tween different sections of the same genus. Part of this diversity may differ not only between closely related genera but even beand families a great diversity of variation patterns. These patterns (1943) have attempted a classification of the types of species between evolutionary factors have produced in various genera majority of plant groups, nevertheless different interrelationships

### PATTERNS IN THE FAMILY RANUNCULACEAE

In the Ranales, the most primitive order of angiosperms, three genera have been studied by experimental as well as by descriptive systematic methods. These are *Paeonia*, *Delphinium*, and *Aquilegia*. In *Paeonia* the two New World species are complicated by structural hybridity and are discussed in Chapter XI. In the Old World there are ten to twelve diploid species plus a

are probably so. P. delavayi is very polymorphic in nature, but geneous in nature, and two others, P. albiflora and P. intermedia, series, are characteristic polytypic species or species groups. Experonica), and the P. daurica-mlokosewitschii-broteri-cambessedesii maining species groups, P. anomala (including P. veitchi, P. nevertheless reveal that it is a polytypic species. The three regeographic segregation of its variants. Careful field study may emodi, P. tenuifolia, and P. cretica, are fairly certainly homonumber of polyploids which will not be discussed here (Stebbins some of the units of the last-mentioned complex are also geneticarated from each other by strongly developed genetic barriers imental evidence on most of these has shown that they are sepwoodwardii, and P. beresowskii), P. obovata (including P. japthe evidence from herbarium specimens does not indicate any and polytypic species occur with about equal frequency. ally isolated from each other. In Paeonia, therefore, homogeneous 1939, Stern 1946). Four of these diploid species, P. suffruticosa, P. (Saunders and Stebbins 1938, Stebbins 1938a). It is possible that

specific hybridization (Epling and Lewis 1946 and unpublished). all the species native to California, and many of these have been one of which (Ewan 1945) explored thoroughly the North subspecies. A group of "species" which probably form a single and further field studies, as well as genetic analyses, may show American species. In addition, the chromosomes are known of series." These forms replace each other geographically, and Ewan of D. decorum, D. patens, D. menziesii, D. nuttallianum, and their that some of the species which he recognizes are no more than described subspecies in 23 of the 79 species recognized by him. tum, and D. gypsophilum, contain related tetraploid forms. Ewan further analyzed by means of transplants, progeny tests, and intercites examples of intermediate and intergrading populations. relatives, including 13 species placed by Ewan in his "tuberiform polytypic species (Epling, oral communication) is that consisting These are all diploid, except that three, D. hanseni, D. variega-The genus Delphinium has been treated in two monographs,

In another species group are found seven species which have been extensively observed in the field and intercrossed, namely, D. variegatum, D. hesperium, D. hanseni, D. gypsophilum, D. parryi, and D. amabile. These simulate the subspecies of a poly-

typic species in that they in general replace each other geographically or ecologically, and they are obviously closely interrelated. But they all overlap with one or more of their relatives
in a considerable proportion of their ranges, and in these regions
Drs. Mehlquist, Epling, Ewan, Lewis, and the writer have repeatedly and independently seen two or three species growing
side by side with no signs of intergradation. The isolating mechanisms are, however, of a somewhat puzzling nature, and they
will be discussed further in Chapter VI. Five of these species are
themselves polytypic, with two to four subspecies each. Two
California species, D. uliginosum and D. purpusii, are sharply
distinct relict endemics. In Delphinium, as in Paeonia, the pattern is that of closely related homogeneous species more or less
sharply isolated from each other, as well as of polytypic species,
the two types occurring with about equal frequency.

parently is not uncommon. A. flavescens, has been recorded by Anderson (1931) and ap-Natural hybridization between two such species, A. formosa and are relatively remotely related to each other, are highly fertile. that a large proportion of F1 hybrids, even between species that shape, proportions, and color of the sepals and petals. It has tween them in stamens and seed follicles are relatively slight. tative habit and essentially similar leaves, and the differences bemore than one climatic zone. All the species have the same vegeregions which because of their great topographic relief include are almost entirely allopatric; in a few places two species grow long been known among gardeners that they hybridize freely and The species are based almost entirely on differences in the size, in the same region, but rarely, if ever, more than two, if we except distributed through temperate Eurasia and North America. They monograph of the genus, that of Munz (1946), lists 67 species, isolating barriers between species are weak or absent. The latest Aquilegia has become a classic example of a genus in which

The genetic barriers between some of the species have been studied by Skalinska (1928a,b). She found that the F<sub>1</sub> hybrid between the European A. vulgaris and the rather remotely related A. chrysantha of western North America has about 50 percent normal pollen and 20 percent normal seed setting, regardless of which way the cross is made. On the other hand, if the Japanese

with A. chrysantha, the F1 is highly sterile and resembles the species, A. flabellata nana, a close relative of A. vulgaris, is crossed Since according to Munz (1946) A. californica is merely a synwhich direction the cross is made (Skalinska 1929). In a later hybrid is normal and has almost normal fertility no matter in with another North American species, A. californica, the F1 the pistillate parent, is normal and rather highly fertile (Skalinska  ${
m F}_2$  generation. But the reciprocal hybrid, with A. chrysantha as furthermore it tends to produce matroclinous segregates in the mother when A. flabellata nana is the pistillate parent, and strong genetic barriers to gene exchange between the Old World havior of the F1 hybrid differs according to the strain of the onym of A. truncata, the evidence of Skalinska would suggest that different relatives in North America. species centering about A. vulgaris and their morphologically very parental species used in the cross. At any rate, there are no in this group of Aquilegia species the sterility and the genetic behavior and high sterility in A. flabellata nana $\times A$ . truncata. paper (Skalinska 1931) the author reported matroclinous be-1929, 1935). On the other hand, if A. flabellata nana is crossed

Evidence of a different type of genetic isolating mechanism between species of Aquilegia was found by Anderson and Schafer (1933). They grew a plant of A. vulgaris which was homozygous for two recessive genetic factors in a garden beside plants of the same species which had the corresponding dominants, and in addition plants of three other rather distantly related species, A. pyrenaica, A. skinneri, and A. caerulea. Seed harvested from open pollination gave 16 percent of offspring which were the result of outcrossing to other plants of A. vulgaris, but no interspecific hybrids were found.

It is evident, therefore, that while many of the described species of Aquilegia are only subspecies from the standpoint of their genetic isolation from each other, nevertheless the whole genus cannot be viewed as one large polytypic species. Munz, however, was unable to divide the genus into natural species groups even on the basis of the most careful study of herbarium specimens and living plants in the garden, and he pointed out great divergences between previous groupings of the species. The true boundaries of the species, therefore, cannot be determined

are merely morphologically distinguishable ecotypes or subof only a few ecospecies. Probably most of the recognized 'species' somal apparatus, has remained comparatively stable. spects is primitive even for that family. In Aquilegia, therefore, of the north temperate regions (see Chapter X). Furthermore, it chiefly in the old floristic communities of the mesophytic forests youthful stage experienced by many other, now mature genera" ably another rather complex polytypic species. The suggestion of canadensis and the western A. formosa and their relatives, is probclosely related to these. A. vulgaris, therefore, is probably a polyspecies." The largest of these ecospecies is undoubtedly the Euro-Hiesey (1945a) that "Aquilegia is one huge cenospecies composed we may now safely accept the statement of Clausen, Keck, and after a careful and systematic series of hybridizations. However, On the other hand, the rest of the plant, as well as the chromo tions, along with great variability of certain organs of the flower. the development of complex polytypic species has been accombelongs to the primitive family Ranunculaceae and in some reyouthful genus in terms of chronological time. It is distributed interspecific barriers themselves. But Aquilegia is probably not a is reasonable if the reference is confined to the evolution of the Clausen, Keck, and Hiesey, that Aquilegia "possibly represents a legias, the red-flowered series, consisting of the eastern Aquilegia latus among the vertebrates. Among the North American Aquitypic species which in its complexity rivals Peromyscus manicuvarieties or subspecies of A. vulgaris, and 12 more are obviously Munz have been recognized by at least one other botanist as pean A. vulgaris. Twenty-nine of the 67 species recognized by by a study of the specimens alone, but will become apparent only panied by much geographic isolation of segments of its popula-

# VARIATION PATTERNS IN THE GENUS Potentilla

The second family of angiosperms to be considered, the Rosaceae, is noted for the frequency of polyploidy, both within and between genera, as will be discussed in Chapter VIII. Phylogenetically it is usually placed in an intermediate position. The genus that will be considered here, *Potentilla*, contains at least three different types of variation pattern. Most of the genus consists of one or a few large polyploid or agamic complexes and will be

discussed in Chapters VIII–X. The subgenus Drymocallis, however, consists entirely of diploid species, with the somatic number of 14 chromosomes. Its most widespread and complex species is P. glandulosa, which contains at least 11 subspecies (Clausen, Keck, and Hiesey 1940). These are adapted to regions as widely diverse as the warm-temperate, semiarid coast of Southern California and the arctic-alpine meadows of the high Sierra Nevada. Diagnostic characters for distinguishing the subspecies are found in the stolons, leaves, stems, inflorescences, sepals, petals, and seeds, and 16 such characters are tabulated for the 11 subspecies by Clausen, Keck, and Hiesey. In addition, transplant experiments revealed great physiological differences in rhythm of growth and other characteristics, and an important biological distinction is that some of the subspecies are self-incompatible and normally cross-pollinated, while others are self-compatible and largely self-fertilized.

Within *P. glandulosa*, therefore, we find all the types of morphological and physiological differences that often separate good, distinct species. If the extreme forms, such as subspp. *typica* and *nevadensis*, should become completely isolated from each other geographically and should develop genetic sterility barriers, they would become amply distinct species even without any further divergence in morphological and physiological characteristics. What keeps the species a single unit is the fact that every subspecies is at some locality or localities in contact with at least one other subspecies, and where these contacts occur, hybrid swarms are regularly found.

Two other species of the subgenus *Drymocallis* occur in the United States, *P. arguta*, which has two subspecies, and *P. fissa*, which is confined to high altitudes in the central Rocky Mountains and is homogeneous. Intermediates between these two and *P. glandulosa* are reported from herbarium specimens, but apparently have not been observed in the field. In the Medicine Bow Mountains *P. fissa* and *P. glandulosa* subsp. *glabrata* occur sympatrically, but no information is available on the extent and manner of their isolation from each other in that region. The uncertainty in regard to these species, which belong to one of the groups best known from the experimental point of view, is clear evidence of how much work is necessary before the true nature of species even in a relatively simple group can be made clear.

nized in animals. differ from the strictly geographic subspecies commonly recogrelatively small chance of receiving pollen from subsp. hansen. maintain themselves as fairly distinct. This separation is aided by Park, hybrid swarms are found, but elsewhere the subspecies moist meadows and subsp. reflexa on dry slopes. Where these two ecologically isolated from each other, subsp. hanseni occurring in graphic range of subsp. reflexa. The two subspecies are, however, teristics. In the first place, it is included entirely within the geothey are usually understood in animals, has two unusual characreflexa, hanseni, and nevadensis, all best known from the Sierra are typica, most characteristic of coastal California, and subspp. within this species is desirable. Its four best-known subspecies mental taxonomy, a further examination of the variation pattern These two subspecies, therefore, represent edaphic ecotypes and the fact that subsp. reflexa is self-pollinating and therefore has a habitats come together, as at Aspen Valley in Yosemite National Nevada. Subspecies hanseni, when compared with subspecies as flowering plants in North America from the standpoint of experi-Since Potentilla glandulosa is one of the best-known species of

The second peculiarity about subsp. hanseni is that in morphological characteristics and ecological preferences it is intermediate between subsp. reflexa and subsp. nevadensis. Furthermore, all the moist meadows which it occupies have resulted from the filling up of postglacial lakes and are therefore one of the most recent habitats in the region. The suggestion of Clausen, Keck, and Hiesey (1940, p. 44) that subsp. hanseni has resulted from hybridization between subspp. nevadensis and reflexa, with subsequent stabilization of a group of segregates adapted to meadow conditions, is very plausible. It may be true that this type of origin of new subspecies in plants is not infrequent.

The other seven subspecies of *Potentilla glandulosa* are much less well known, but the fact that four subspecies occur together in another region, namely, the Siskiyou mountain area of northwestern California, suggests that here also edaphic factors rather than climatic or geographic factors are maintaining the isolation between subspecies. The subgenus *Drymocallis* contains, in addition to the three American species mentioned, six in Eurasia. One of these, *P. rupestris*, is widespread, while the other five are

endemic to mountain areas in southwestern Asia, from Asia Minor and the Crimea to the Himalaya. The interrelationships between these species are relatively poorly known, but available evidence indicates that some of them may be subspecies of *P. rupestris*. Attempts to produce hybrids between *P. rupestris* and *P. arguta* have failed, suggesting that the American and the Eurasian species of the subgenus *Drymocallis* are well isolated from each other and may represent different cenospecies.

The subgenus *Drymocallis* of *Potentilla*, therefore, contains two widespread, polytypic species, one in each hemisphere of the Holarctic region, and several less complex species about equally distributed in each hemisphere.

#### THE GENUS Quercus

The next variation pattern to be considered is that of the oaks, a typical genus of angiospermous trees of the north temperate regions. According to present systematic treatments the genus Quercus, of the family Fagaceae, contains between 250 and 300 species, but it is very likely that many of these actually are subspecies of a smaller number of polytypic species. Cytologically all species of Quercus investigated, as well as all other genera of the Fagaceae, have the same chromosome number, n = 12, and only slight differences exist between them in size and in morphology of the chromosomes (Hoeg 1929, Ghimpu 1929, 1930, Jaretzky 1930, H. J. Sax 1930, Natividade 1937a,b, Duffield 1940). Natural hybrids between generally recognized species are not uncommon, and both cytological and genetic studies indicate that most are fertile (Sax 1930, Yarnell 1933, Wolf 1938, 1944).

The usual treatments in the various regional floras do not describe geographic varieties or subspecies within most of the oak species, although *Q. borealis*, the common red oak of the eastern United States, and three species of live oaks of California, *Q. wislizenii*, *Q. agrifolia*, and *Q. dumosa*, are commonly recognized to be polytypic. Nevertheless, the experience of the writer and of most other observers with whom he has spoken indicates that a much larger number of species will be found to possess geographic variation when their intraspecific variation patterns are studied more carefully. Furthermore, hybrid swarms are often found in regions where the ranges of markedly different allo-

similar hybridization apparently occurs at the southern end of garryana and Q. douglasii, mentioned in Chapter I. Furthermore, often in mixed stands throughout most of their ranges, also show of forms is equally great. This situation is further complicated acorn cups and other reproductive structures within this series ceous, evergreen, unlobed leaves; and the range of differences in broad, deciduous, lobed leaves to a shrub with very small, coriachange between a chain of forms ranging from a large tree with patric species overlap. A conspicuous example of this is Quercus amount of intergradation that can occur between them. ter of the tree, shows relatively little evidence of hybridization the counterpart of Q. garryana both in habitat and in the characby the fact that the valley oak of California, Q. lobata, which is its var., turbinella. There is thus good evidence of gene interthe range of Q. douglasii, where it overlaps with Q. dumosa and California, therefore, the magnitude of the visible differences little evidence of intergradation with each other. In the oaks of lobata and Q. douglasii, although they occur sympatrically and between species seems to bear no direct relationship to the in localities occupied by these two species. Furthermore,  $Q_{\cdot}$ 

of the pine-barren area and in general remain quite distinct from equal to that found in a series of specimens representing the sample of Q. marilandica from Cliffwood, New Jersey, showed a usually well separated from each other geographically or ecospecies occur together chiefly on the coastal plain of New Jersey, were described as natural hybrids more than fifty years ago (Daviss each other. But shrubs intermediate between them do occur and entire geographic range of the species from Pennsylvania to range of variation in six important diagnostic characteristics ranges, as is evidenced from the fact that a single population logically. Both species are relatively uniform throughout their throbalanus (black oak) in the eastern United States. These marilandica and Q. ilicifolia, two species of the subgenus Ery-New Jersey, the two species occur sympatrically over a large part by Stebbins, Matzke, and Epling (1947), with the aid of Ander-Kansas and Arkansas (Stebbins, Matzke, and Epling 1947). In 1892). Study of a population sample from one of these localities Long Island, and Staten Island, New York, but elsewhere are A relationship of a somewhat different type is shown by Q.

son's hybrid index method (see Chapter VII), revealed that eight out of the sample of 86 specimens were sufficiently intermediate between the two species so that they could be considered F<sub>1</sub> hybrids or their derivatives, while about 35 percent of the specimens showed some evidence of admixture of genes derived from the two species. Most of them were introgressive types of Q. marilandica, containing a few genes from Q. ilicifolia.

species may be connected by both first-generation hybrids and eastern United States, where in many regions as many as ten or a series of hybrid types under the name X Q. jolonensis has an example of this. Although they are distinct nearly everywhere, other intergrading types which are the result of segregation or showing any signs of intergradation, but in other areas these same the fact that species may occur together in many regions without margined leaves, and Q. velutina, the black oak, which has much are Q. imbricaria, the shingle oak, which has elliptic entiretype are more numerous. Two very distinct species of this region eleven species of oaks may occur sympatrically, examples of this been described from a small area in central California. In the backcrossing. Q. douglasii and Q. lobata, mentioned above, are sembles Q. velutina more closely than does Q. imbricaria, and occasional isolated tree, and in some regions considerable numbark, and other characteristics. The easily recognizable hybrid, broader, deeply lobed leaves and differs greatly in acorn cups, although the two species are sufficiently distinct over much of which bears the name  $\times Q$ . leana, has often been noted as an elsewhere the populations of Q. velutina contain genes from are almost as frequent as trees typical of one or the other species. their range, in some areas, such as the Atlantic coastal plain of Camp, oral communication). Quercus coccinea, the scarlet oak, rebers of hybrid and hybrid derivative trees have been seen (W. H. resulted from hybridization and introgression of genes from other most of the geographic and other variability of Q. velutina has Long Island and southeastern Massachusetts, intermediate types an even more marked degree of intergradation has been noted by species, as is discussed in Chapter VII. Turning to the white oaks, Furthermore, in the coastal plain of New Jersey and probably Q. marilandica, the blackjack oak. It is likely, therefore, that The variation pattern in Quercus is further complicated by

Hampton (Trelease 1924) and by E. Anderson (oral communication) between Q. bicolor, the swamp white oak, and Q. macrocarpa, the bur oak, both of the central United States. It is remarkable that these two species, although they occur sympatrically over most of their geographic ranges and inhabit rather similar ecological sites, should be able to intergrade so freely and nevertheless retain their identity.

sympatrically in parts of the Balkan Peninsula. The three most species recognized by Ascherson and Graebner, at least 12 occur species (Ascherson and Graebner 1913, pp. 445-544). Of the 15 ging univalent chromosomes and tetrads with extra nuclei. two types of soil meet. The same hybrid was studied in Denmark are different, and hybrid trees are frequent along the zone where ships between Q. robur and Q. sessiliflora in Hertfordshire, Salisbury (1940) has given an account of the ecological relationand by others as merely aberrant forms of the parental species been considered by some specialists to be of hybrid derivation species are frequent in many places, so that many forms have more, forms intermediate between the hybrids and their parental that triple hybrids occur, involving all three species. Furtherbotanists from many parts of Europe, and there is some evidence sessilis (Q. sessiliflora), and Q. lanuginosa. Hybrids between all variable species are the closely interrelated group, Q. robur, Q. cannot be segregated into well-marked geographical races or subental species, but in several trees of an intermediate type Hoeg meiosis showed perfectly regular, normal conditions in the parhybrid and its parents. Studies of chromosome behavior in by Hoeg (1929), who noted also trees intermediate between the England. In this region, the soil preferences of the two species three of these species have been reported as frequent by numerous found irregularities typical of interspecific hybrids, such as lag-In Europe, the species of Quercus, though extremely variable

One interesting feature of the progeny of both artificial and natural oak hybrids is that in respect to vegetative characteristics they usually segregate so sharply that even among a relatively small number of individuals the parental types can be recovered (MacDougal 1907, Ness 1927, Coker and Totten 1934, Allard 1942, Wolf 1938, 1944, Yarnell 1933, and Stebbins, unpublished). This is in striking contrast to the behavior of interracial and inter-

specific hybrids in most other plant groups, in which the number of genetic factors controlling the differences between them is so large that it is relatively difficult to recover the parental types (Müntzing 1930a, Goodwin 1937, Winge 1938, Jenkins 1939, etc.). The evidence available suggests that the number of genes by which species of oaks differ from each other is considerably smaller than it is in the case of most other plant groups.

oaks of the United States, compiled from the extensive references often found near the edge of the geographic range of a species, although hybrids may be expected between almost any two species concluded that this rarity is due, not to the difficulty of producwhere it is rare and related species are more common. He has very small percentage of the total population. They are most belonging to the same section, nevertheless they always form a in nature over a period of many years. He has pointed out that in the literature, but based also on his own experience with them ents when pollinated with their own species. Pjatnitzky was even species. A noteworthy fact is that some intersectional crosses, such was also true in pollinations between members of the same tained in every case a rather low percentage of acorns set, but this artificial cross-pollination of several different species. He obout by the results of Pjatnitzky (1946a,b) from experiments on hybrids have of becoming established. This conclusion is borne ing interspecific hybrids, but to the small chance which the one between the subgenera Lepidobalanus and Erythrobalanus able to obtain a type of hybrid not known at all in nature, namely, as Q. macranthera  $\times$  robur and Q. macranthera  $\times$  alba, gave yet reached maturity, and no data are available on their fertility. maxima). These seedlings, though vigorous, have apparently not (Q. robur imes borealis maxima and Q. macranthera imes borealishigher yields of acorns and seedlings than did the maternal par-Palmer (1948) has presented a complete list of the hybrid

In *Quercus*, therefore, we have a rather large number of species, many of them polytypic, which are often capable of exchanging genes with other, morphologically distinct populations occurring in the same region and in similar habitats. These sympatric populations have always been kept apart as distinct species, but some of them, such as *Q. bicolor* and *Q. macrocarpa*, behave genetically more like subspecies. The correct systematic treat-

ment of such a variation pattern is a difficult problem, and it may be remarked that other genera, for instance, Ceanothus (McMinn 1944), and probably Vitis, are essentially similar to Quercus. The great evolutionary possibilities of such situations will be discussed in Chapter VII.

### PATTERNS IN THE FAMILY COMPOSITAE

a pattern of mostly homogeneous, sharply differentiated species each other. This situation is not uncommon in the higher plants. which form occasional hybrids when they come in contact with in range lands overgrazed by cattle. Wyethia, therefore, presents patric, like W. helianthoides and W. amplexicaulis, they comhybrid swarms, and even when two species are completely symrarely produce enough offspring in nature to form extensive hybrids between them are found. Although these are fertile, they subgenus. When two such species occur together, occasional is wholly or partly sympatric with at least one other of the same and markedly different, as well as distinct from each other. In x = 19. Only one, W. scabra, is recognized as polytypic, but in species, all deep-rooted and large-flowered perennials, native to found have probably resulted from recent spreading of the species pletely retain their identity. Such small groups of hybrids as are the other two subgenera, Alarçonia and Euwyethia, each species three subgenera, subg. Agnorhiza, the six species are allopatric presence of variant forms is noted. In the most primitive of the three others, W. angustifolia, W. ovata, and W. arizonica, the quantitative and analytical methods. The first is the genus cussed, all of which have recently been studied with the use of four species counted have the basic haploid chromosome number the arid and semiarid regions of the western United States. The Wyethia, monographed by Weber (1946). This consists of 14 In the great family of Compositae, three genera will be dis-

The next genus to be discussed, Layia, is one of the most thoroughly investigated cytologically and genetically, as well as systematically, of any of the plant kingdom, but only part of these investigations have been published (Clausen, Keck, and Hiesey 1941, 1945a, 1947). It contains 13 diploid and one tetraploid species. Of the diploids five — L. platyglossa, L. chrysanthemoides, L. gaillardioides, L. pentachaeta, and L. glandulosa — are

polytypic, while the others are genetically rather homogeneous and of limited geographic distribution. Interspecific hybridization, which has been attempted in nearly every possible combination, has in many cases been completely unsuccessful and in others has yielded hybrids of various degrees of sterility and reduction in chromosome pairing. The results of these hybridizations are summarized in Fig. 7.

The pattern of species relationships in Layia is in some respects

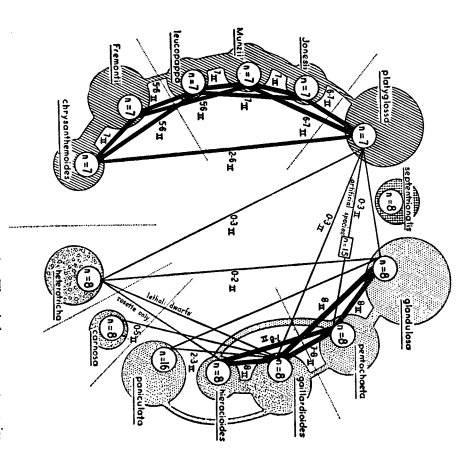


Fig. 7. Relationships in the genus Layia. The circles represent species, with chromosome numbers indicated; shaded connections show degrees of genetic affinity, and width of solid black lines represents degree of chromosome pairing in interspecific hybrids. The dotted lines indicate major morphological breaks in the genus. From Clausen, Keck, and Hiesey 1941.

different from that found by Clausen, Keck, and Hiesey (unpublished) in other genera of Madinae, but it is probably similar to that in other genera of annuals which make up a large proportion of the California flora. The presence of closely related species in adjacent areas and occupying habitats known to be geologically recent, along with genetic barriers in various stages of development, suggests that *Layia* is in an active state of relatively rapid evolution

The genus *Crepis* is the most thoroughly studied of the larger genera of angiosperms. The monumental work of Babcock (1947) on this genus shows a remarkable insight into the genetic evolutionary processes that have been operating in it. With the exception of the polyploid, apomictic species of North America, the bulk of the species are diploid and form an aneuploid series with the numbers x = 7, 6, 5, 4, and 3. Of the 196 species recognized by Babcock, 97 are known to be diploid, and 78 are probably so. Of these 175 diploid species, only 18 are recognizably polytypic. The best known of these is *C. foetida* (Babcock and Cave 1938), which has three subspecies, within some of which local varieties and forms have been recognized.

The possibility is remote that any appreciable number of the species of *Crepis* are actually allopatric subspecies of more comprehensive species, as Mayr (1942, p. 122) has suggested may be true of plant genera. In 22 of the 27 sections into which Babcock has divided the genus, the majority of the species are either sympatric or with overlapping distributions, and natural hybrids between them are rare or unknown. Of the five sections with predominantly allopatric species, the one which has been investigated genetically possesses complete barriers of genetic isolation separating its species. The species of the four other sections are morphologically at least as sharply distinct from each other as are species of *Crepis* known to be separated by reproductive isolating mechanisms; hence, it is unlikely that they represent subspecies under any criterion.

Jenkins (1939) has described an example of four allopatric, insular endemic species of *Crepis* which are separated by imperfectly developed barriers of hybrid sterility and might therefore be considered to be species in the making. It is very likely that there are other similar examples in the genus, but both mor-

phological and genetical evidence indicates that these are uncommon. In another example, *C. pulchra*, *C. palaestina*, and *C. pterothecoides*, all of which occur sympatrically in Palestine and Lebanon, were found to form fertile or only partly sterile hybrids (Babcock 1947, p. 150). These are apparently separated from each other by barriers of ecological or seasonal isolation (see Chapter V).

In *Crepis*, therefore, the great majority of the diploid species, though somewhat variable, are not polytypic in the sense that they consist of well-marked subspecies which replace each other geographically or ecologically. And when studied, they have been found to be separated from each other by more or less well-developed barriers of reproductive isolation, which differ in character according to the group investigated. The processes which have given rise to these barriers, and to the alteration in chromosome number and morphology which is one of the most distinctive features of the genus, will be discussed in later chapters.

### THE FAMILY GRAMINEAE

angiosperms. The only genus containing more than a few species subspecies than are found in any North American species. The and its relatives apparently form a polytypic species with more Graebner, and Komarov indicate that the variation pattern in species in such standard floras as those of Hegi, Ascherson and closely related species, M. imperfecta, M. torreyana, and M. calirelated species. Of the 18 North American species, all of which which is strictly diploid is Melica. This genus was studied cytohas the highest proportion of polyploid species of any family of similar. M. papilionacea is probably polytypic, but few, if any, of South American Melicas are less well known, but apparently them is similar to that in North America, except that M. ciliata impossible or nearly so. The systematic treatments of the Eurasian polytypic, and these contain only two subspecies each. Three very have the somatic chromosome number 2n = 18, only two are (1944) has investigated representative hybrids between closely logically and systematically by Boyle (1945), while Joranson fornica, have been artificially crossed. They form almost completely sterile hybrids, so that gene exchange between them is The fifth family to be discussed here, the Gramineae, probably

genera of grasses, consists mostly of homogeneous or only slightly variable species which are sharply separated by barriers of genetic the others are. Melica, therefore, in contrast with most other

species were it not for the fact that hybrids have been made conregions of overlap. They might be taken for well-marked subgaris, and B. suksdorfii - appear to be homogeneous. Morphowestern North America mostly of diploid species. Two of these, in characteristics of external morphology. to that in Melica, except that the species are less sharply defined variation pattern in this section of Bromus, therefore, is similar pletely or almost completely sterile (Stebbins, unpublished). The necting the majority of them, and all these hybrids are comthey replace each other geographically except for smaller or larger logically these species are rather similar, and for the most part the others - B. kalmii, B. porteri, B. laevipes, B. grandis, B. vul-B. ciliatus and B. orcuttianus, have at least two subspecies, but In another genus, Bromus, the subgenus Bromopsis consists in

#### GENERAL CONCLUSIONS

side under uniform conditions and occupying somewhat different sists of two or more geographically separated subspecies. In a majority of homogeneous or variable, but not polytypic, species, geographic range of any extent. The recognized species, whenphylogeny may be summarized as follows. Four of themexists in the form of variants distinguishable when grown side by Crepis, Melica, Bromus, and probably Wyethia, genetic diversity F<sub>1</sub> hybrids is the most common type. barriers of reproductive isolation. These barriers are of a diverse ever investigated, have been found to be separated by well-marked distinguished by morphological differences recognizable over a but they cannot be recognized as subspecies because they are not but in every case a larger or smaller minority of the species connature, but the partial or complete sterility of pollen or seed in habitats. These are ecotypes in the original sense of Turesson, Wyethia, Crepis, Melica, and Bromus subg. Bromopsis — contain families scattered throughout the system of classification and This survey of ten genera of seed plants belonging to five

Of the remaining six genera three, Paeonia, Delphinium, and

and Potentilla subg. Drymocallis, are the majority of the species rather weak. In only three of the ten genera, Quercus, Aquilegia, species are very well developed, but in Delphinium they are proportions. In Paeonia, barriers of hybrid sterility between the and Diplacus, indicates that the pattern found in Quercus is Layia, contain polytypic and homogeneous species in about equal ecological nature and operate through the failure of most hybrid regions, such as Acer, Fraxinus, Populus, Vaccinium, Ceanothus, of this type. What is known of the woody genera of temperate derivatives to become established. F<sub>1</sub> hybrids. The isolating mechanisms best developed are of an having ranges that coincide almost completely, often form fertile ranging and polytypic; in addition the good species, even those this pattern is not only the fact that the species are mostly wide typical of such genera (see Chapter VI). A notable feature of polytypic. But it is noteworthy that the woody genus discussed is

cannot apply uncritically the criteria of species which have been have very different modes of life. This generalization applies with developed in one group to the situations existing in another, parfrom the additional ones given in Chapters V to XI, is that we ciples of systematics, as well as of evolution, must be based on as their yardsticks of species on animal material. General prinwhich have been recognized by botanists, or of the latter to use ticularly if the groups are distantly related to each other and broad a knowledge as possible of different groups of plants and particular force to attempts of zoologists to reinterpret the species The generalization emerging from these examples, as well as