CHAPTER X

Apomixis in Relation to Variation and Evolution

and the Coelenterata parthenogenesis is the only possible method sexual reproduction and allogamy. value in them of genetic systems which deviate from continuous their tendency for indeterminate growth, increases the selective relatively simple structure and development of plants, as well as accordance with the principle, developed in Chapter V, that the but is by no means as widespread as it is in plants. This is in animals, such as rotifers, crustaceans, Hymenoptera, and aphids, of regular asexual reproduction. It appears in various groups of plants. In most animals higher than the Protozoa, the Porifera, applies to only a part of the apomictic phenomena found in term parthenogenesis, though better known to most biologists, HE TERM APOMIXIS is a general one, covering all types of substitutes for the sexual method. The more familian asexual reproduction which tend to replace or to act as

material included, and for more detailed information on every Gustafsson (1946, 1947b,c). This chapter, therefore, will be thorough, complete, and discerning review and analysis of origin are both multiform and complex. The earlier reviews of more, have shown that numerous methods of apomixis exist in the reader is advised to consult Gustafsson's work phase of this subject, as well as for a complete list of literature, the based chiefly on this review, particularly in respect to the factual (Stebbins 1941a) have now been superseded by the vastly more the literature on apomixis by Rosenberg (1930) and the writer plant kingdom, and that their genetic nature and evolutionary plant genera, and more or less casual surveys of apomixis in many Careful analyses of the apomictic phenomena in a number of

METHODS OF APOMICTIC REPRODUCTION

most botanists, so that many terms were used uncritically and various parts of the normal sexual cycle were fully understood by the morphological nature and the genetic significance of the encountered in organisms reproducing by normal sexual methods and in many ways quite different from any which are regularly for this. In the first place, the phenomena themselves are complex specialists as to the correct usage of terms. There are two reasons well as the differences of opinion which have arisen among has been the highly complex terminology it has developed, as chief barriers to an understanding of the apomictic phenomena a single substitute for both of these processes or a separate subessential processes, meiosis and fertilization, which are separated of the higher plants includes two entirely different and equally stressed the all-important point that since the normal sexual cycle to corresponding sexual processes. In particular, Winkler first the apomictic phenomena which show correctly their relationship the phenomena they purported to describe. We owe chiefly to were defined in ways that included erroneous interpretations of Secondly, the early research on apomixis was carried out before stitute for each of them, with coordination of these two substiof the gametophyte, these two processes can be expected to be chromosomes at meiosis and has produced an embryo sac and egg without fertilization is not possible unless previously some process which lead to diploid gametophytes and egg cells does not ensure cell with the diploid chromosome number. On the other hand, has occurred which has circumvented the reduction of the tutes. Thus, parthenogenesis through development of the egg cell Every harmonious apomictic cycle must therefore provide either influenced by entirely different environmental and genetic factors. from each other by the entire period of growth and development Winkler (1920, 1934) the first classifications and terminology of the occurrence of parthenogenesis and apomictic reproduction, as the mere occurrence of various altered developmental processes has been shown in a number of organisms. On the part of botanists not specializing in this field one of the

archesporial cell in the ovule to a diploid embryo sac have not since some of the modifications of meiosis which lead from an Nevertheless, the terminology of Winkler is not complete,

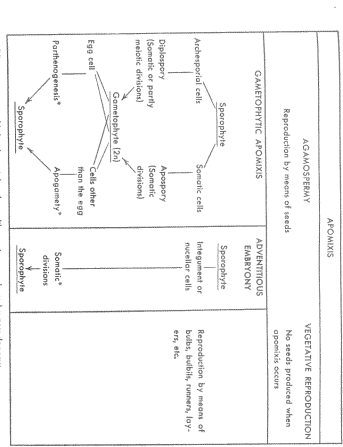
a relatively simple terminology is sufficient for all practical clarity of his lengthy exposition of the apomixis phenomena that embryo sac. Gustafsson (1946) has pointed out that Fagerlind's recognized modifications of meiosis. In addition, he shows by the numerous intermediate situations which exist between the various different ways of circumventing meiosis and producing a diploid karyological classification is difficult to apply because of the terminologies, in which names were given to as many as tour been recognized. For these reasons, Stebbins and Jenkins (1939), Fagerlind (1940a), and Stebbins (1941a) proposed amplified

animals. Since these occurrences are known only in laboratory Fig. 1) may therefore be simplified to that extent (Fig. 36). is correct, and the chart presented by the writer (Stebbins 1941a, cultures and are without evolutionary significance, this decision which have been reported from time to time in plants as well as in phenomena those isolated occurrences of haploid parthenogenesis Gustafsson does not include in his classification of the apomixis

sexual organisms. Gustafsson cites a number of species, such as such conditions, structures like stolons, rhizomes, and winter various members of the family Lemnaceae which in northern species under these conditions is essentially apomictic, although buds, which normally act only as accessory methods of reproducactivity. This may come about through a variety of causes, both sexual processes are not functioning or are greatly reduced in apomixis, vegetative reproduction and agamospermy. The are normally sexual Europe reproduce wholly by such asexual means, but elsewhere Elodea canadensis, Stratiotes aloides, Hydrilla verticillata, and the same genetic types may under other conditions be normal tion, may assume the entire reproductive function, so that the phenotypic and genotypic, as was discussed in Chapter V. Under former may be considered as apomixis whenever the normal The classification of Gustafsson includes two principal types-ot

non, often termed vivipary, is well known in such genera as methods of vegetative reproduction in which the propagules occur Polygonum (P. viviparum), Saxifraga, Allium, Agave, and some within the inflorescence and replace the flowers. This phenome More striking, and perhaps of broader significance, are those

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i.e., under the influence of pollen tubes or endosperm development. *The processes at this level can take place either autonomously or by pseudogamy,

tound in the higher plants. Modified from Gustafsson 1946. Fig. 36. Chart showing the interrelationships of the processes of apomixis

and therefore from the genetic point of view may be quite comwith this mode of reproduction may fail entirely to produce seed, genera of grasses, particularly Poa and Festuca. Species or races parable to the obligate apomicts which reproduce by seed.

which develops in the seeds is usually identical with its maternal which embryos and seeds are formed by asexual means. The simplest method is adventitious embryony, in which embryos essential feature of all types of agamospermy is that meiosis and species of the genus Citrus, but it is also found in Alchornea pletely omitted. This phenomenon is best known in various nucellus or ovule integument, and the gametophyte stage is comdevelop directly from the diploid, sporophytic tissue of the parent in both chromosome number and genic content. The fertilization are circumvented, so that the embryo (or embryos) Agamospermy includes all types of apomictic reproduction in

warm temperate or tropical climates. embryony seems to be relatively frequent in species native to Spiranthes (Orchidaceae). References are given by Stebbins rulata (Ochnaceae), in Eugenia jumbos (Myrtaceae), in Sarcococca ilicifolia and Euphorbia dulcis (Euphorbiaceae), in Ochna ser-(1941a) and Gustafsson (1946). For some reason, adventitious Hosta and Allium (Liliaceae) and of Nigritella, Zygopetalum, and pruniformis (Buxaceae), in Opuntia aurantiaca, and in species of

that is, apomixis via a morphological gametophyte, which some suggestion is here made that the term gametophytic apomixis, thenogenesis." Since a collective term is obviously needed, the and awkward phrase "apospory and diplospory followed by parspermy which are not adventitious embryony by the rather long illogical. He himself refers frequently to those types of agamosubstitute term apogumogony, which Gustafsson rejected as in a totally different sense. Fagerlind (1944b) then suggested the as Gustafsson pointed out, had been used previously by Hartmann gony by Fagerlind (1940a) and Stebbins (1941a), a term which, circumventing meiosis. These were collectively termed agamowhich diploid gametophytes arise as a result of some method of tophyte-sporophyte cycle of alternation of generations, but in times can function sexually, is both descriptive and appropriate. from the morphological viewpoint a complete sporophyte-game-More common are methods of agamospermy which include

entirely omitted or are so modified that pairing and reduction of cell of the archesporium, but in which meiotic divisions are either divisions; and diplospory, in which the embryo sac arises from a of the nucellus or of the inner integument by a series of somatic pory, in which a diploid embryo sac is formed directly from a cell evolution. Whatever the method of formation of diploid gamematter outside of the present discussion of apomixis in relation to to the plant in which they were first found (Antennaria scheme, the omission of meiosis have frequently been designated according the chromosomes does not occur. These various modifications and whether any special terms are needed for these phenomena, is a Which of these methods of terminology is the more desirable, or (1941a), and Battaglia (1945) have used special terms for them. Taraxacum scheme, etc.), but Fagerlind (1940a), Stebbins There are two principal ways of circumventing meiosis: apos-

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essentially the same. Figure 36 shows a modification of the simplitophytes, the end result, from the evolutionary point of view, is nomena included in agamospermy, or apomixis with seed formafied terminology recognized by Gustafsson to describe the phe-

others require pollination and the development of pollen tubes. may give rise to embryos through the multiplication of either the as adventitious embryony, are in some plants autonomous, but in plants, but apogamety is frequent in ferns. Both processes, as well ciously, but the endosperm will not develop unless it is fertilized they will not seed unless pollination has occurred, but the plants Such apomicts are said to be pseudogamous. Like sexual forms, former phenomenon is by far the most common one in the seed egg cell (parthenogenesis) or some other cell (apogamety). The pollination appears to vary with the species concerned. In most formed from such seed are maternal in character. The effect of opment of the embryo, the endosperm, or both are very often so some examples, particularly the species of Ranunculus investicontinued growth of the embryo. Gustafsson nevertheless cites the embryo begins development autonomously and even precopseudogamous species, such as Allium odorum (Modilewski 1930), has already started. are ready to receive pollen the formation of the embryo and seed precocious that at the time when the flowers open and the stigmas does not take place. When parthenogenesis is autonomous, devel $_{
m gated}$ by Häfliger, in which fertilization of the endosperm possibly Pollination and partial fertilization are therefore essential for Potentilla collina (Gentcheff and Gustafsson 1940), Poa spp. Håkansson 1943a, 1944a), and Parthenium incanum (Esau 1946). Diploid gametophytes formed by apospory or by diplospory

or characterized by slight abnormalities, such as a small number them. In plants possessing apospory- ("somatic apospory" of means or by adventitious embryony, but it is not characteristic of is present occasionally in plants which reproduce by vegetative genesis in the ovules and microsporogenesis in the anthers. This is the disturbance of the meiotic divisions of both megasporoof univalent and multivalent chromosomes, in both microsporo-Fagerlind and of Stebbins 1941a), meiosis is likewise either normal One of the most conspicuous features of many apomictic plants

of embryos and seeds, the archesporial tissue of the anthers ment. On the other hand, in some of the American species of of the aposporous apomicts have pseudogamous embryo developcomparative regularity may be connected with the fact that most sacs which become replaced by the products of apospory. This genesis and the formation of the megaspore tetrads and embryo before they reach the stage of meiosis (Stebbins and Jenkins develops so abnormally that the pollen mother cells degenerate Crepus, which combine apospory with autonomous development

abnormalities may be a direct or indirect result of past hybridizahas made such an opinion untenable, although some of the hybridization. Further knowledge of these phenomena, however, are of hybrid origin, and therefore that apomixis is caused by thirty years ago by Rosenberg (1917), were considered by Ernst of the meiotic divisions. These abnormalities, which were found traction of the chromosomes, retardation of meiosis, and precocity abnormalities are failure of chromosome pairing, failure of conmore carefully by Gustafsson (1947b). The most frequent genera. This work was reviewed by Stebbins (1941a) and even acum, Antennaria, Calamagrostis, and some species of other abnormalities have been intensively studied in Hieracium, Taraxwith autonomous development of embryos and seeds. These diplospory (including "generative apospory" of Stebbins 1941a) genesis are found in those plants which combine some form of tion between species. (1918) as strong evidence for his hypothesis that apomictic forms The most striking abnormalities of both micro- and megasporo-

to do with the lack of homology between dissimilar chromosomes example, it is quite clear that the failure of pairing has nothing entirely omitted or may in some ovules occur in a very abnormal contracted chromosomes. But in the ovules, meiosis is either regular, with a high degree of bivalent formation and normally in nature have microspore meiotic divisions which are essentially instance (Stebbins 1932b), the occasional male intersexes found female organs of the same species. In Antennaria fallax, for they are often, and perhaps usually, different in the male and the fashion, with practically no chromosome pairing. In such an The most conspicuous feature of these abnormalities is that

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of various Hieracium apomicts has been studied by several ment of the ovular and archesporial tissue. Meiosis in the anthers but is connected with some disturbance in the normal developmitotic, while those of the megaspore mother cells are more like since the divisions of the pollen mother cells are essentially are essentially mitotic; but in C. chalybaea, the reverse is the case, varying number of univalents, while in the ovules the divisions the anthers, although these are irregular and characterized by a found that in Calamagrostis lapponica meiotic divisions occur in these are not duplicated in the ovules. Finally, Nygren (1946) has workers, and a series of abnormalities has been described; but meiosis, although the formation of restitution nuclei causes them accompanied by great precocity in the time when they begin, the reversion toward mitosis in the pollen mother-cell divisions is to produce díploid megaspores and gametophytes. In Hieracium, while the same change in the ovules is associated with a delay in their onset or with precocity. in the pollen mother cells may be associated either with a delay in the onset of the divisions. In Calamagrostis, mitosislike divisions

meiosis, as well as the differences between anthers and ovules of occurrence of a series of developmental processes, and that it can normal course of meiosis requires the carefully synchronized the same plant, can be explained only if we assume that the still has by no means been exploited to its full extent. and genetically controlled disturbances of one or more of these be upset in a number of different ways by various environmentally differences between mitosis and meiosis and their causes, which processes. They provide a wealth of material for studies of the The complexity and multiformity of these disturbances of

a plant may produce diploid embryo sacs through failure of will therefore produce some completely maternal progeny while the latter will not form seeds unless fertilized. Such plants of parthenogenetic development, usually through pseudogamy: and egg cells. The former ovules contain diploid egg cells capable pletion of the meiotic divisions will yield haploid embryo sacs meiosis, while in other ovules of the same plant successful comabnormalities in apomicts is that not infrequently some ovules of through apomixis and other offspring by the normal sexual One of the results of this diversity in the nature of meiotic

embryo sac either does not develop at all or develops so slowly apomixis may also occur because in certain ovules the aposporio that it does not crowd out the haploid one derived from the megaform diploid embryo sacs by means of apospory, facultative process. They are termed facultative apomicts. In plants which

GENETIC BASIS OF APOMICTIC PHENOMENA

single dominant gene, although the expression of the character plex, and apomixis is recessive to sexuality. to the suggestion that in this group the inheritance is more comthe inheritance of adventitious embryony in Citrus has led him mental factors. Gustafsson's brief analysis of the meager data on may be greatly modified by other genes and by various environbulbil production in the former species is largely controlled by a carmatum and the sexual species A. pulchellum showed that bulbils which reproduce it vegetatively. A cross between A carinatum, a species in which the flowers are largely replaced by by Levan (1937b) in the predominantly apomictic Allium the various groups studied. The simplest situation is that found cally controlled, but the basis of inheritance has been different in has shown in every case that the apomixis of the species is genetisame genus. Segregation in later generations from these crosses sexual types and related facultative or obligate apomicts of the A number of hybridizations have now been performed between

sexuality (Müntzing 1940). Furthermore, since progeny from dominance of sexual plants, showing that apomixis is recessive to well as between different facultative apomicts, yield a great precrosses between facultatively apomictic and sexual biotypes, as genetic basis in both is complex, as revealed through several of Potentilla) autonomous development of the embryos. The phyte development followed by pseudogamous or (in some species apomixis is aposporous or diplosporous (in Poa alpina) gametothe genus Potentilla. In both of these groups the mechanism of obtained in the complexes of Poa pratensis and P. alpina and in (see Gustafsson 1947b,c, for complete list of references). In Poa, Müntzing, Christoff and Papasova, and Rutishauser on Potentilla publications by Müntzing and by Akerberg on Poa, and by The most extensive genetic data on apomixis have been

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such crosses yield some plants which possess apospory, but not controlled by different genes, and successful apomictic reproducgeny through pseudogamous parthenogenesis (Håkansson 1943a). have little apospory, but yield a high percentage of haploid proregular pseudogamous embryo development, and others which tion is "due to a delicate genetic balance" between a series of the various processes of the apomictic cycle are in this genus factors.

apomict and a sexual form with a low chromosome number yield number are crossed with sexual forms. On the other hand, crosses crossed with each other or when apomicts with a low chromosome apomicts give sexual progeny. sexuality. In Rubus, as in Poa, crosses between two facultative may in the case of Potentilla dominate over a smaller number for mostly apomictic progeny. Several genes for apomixis, therefore, between two high polyploid apomicts or between a polyploid The results in Potentilla are similar when diploid apomicts are

triploids. The parental plant has 72, the progeny, 108 chromogeny consist of 65 to 90 percent of morphologically aberrant, dwarf as pollination. As a result, the offspring of this plant and its proof these gametophytes cannot develop without fertilization as well embryo sacs are formed in 90 percent of the ovules, but the eggs A42163, described by Powers (1945). Meiosis fails and unreduced without fertilization - are genetically independent of each other. phytes with the unreduced chromosome number and of embryos two essential processes of apomixis-development of gametoyet been reported, evidence has been obtained to indicate that the Although no hybrids between sexual and apomictic biotypes have crude language, this line polyploidizes itself out of existence. some numbers (about 140) and more aberrant morphology. In somes. These triploids, in their turn, produce only unreduced gametophytes and egg cells, and progeny with still higher chromo-This evidence consists of the behavior of a particular strain, No. In the genus Parthenium, the apomicts are pseudogamous.

cium by Ostenfeld (1910) and Christoff (1942), but their evidence $H.~auricula \times aurantiacum$, involving a diploid sexual and a polyis considered doubtful by Gustafsson, since it deals with a cross, ploid apomictic species. In general, the sum of evidence from Dominant factors for apomixis have been reported in Hiera-

apomicts are likely to yield apomictic progeny. Potentilla, in which crosses between two polyploid facultative Hieracium suggests a genetic basis for apomixis similar to that in

and adventitious embryony, that simple genetic behavior can be relatively simple types of apomixis, like vegetative reproduction sterility, or some abnormal genetic behavior. It is only in the of these gene combinations and therefore a reversion to sexuality, combinations controlling their apomixis, will cause a breaking up and a sexual one, or between two apomicts having different gene action of many genes, so that any cross between an apomictic form number often increases its tendency toward dominance. However the fact that a successful apomictic cycle is produced by the interthis recessiveness is not usually that of a single gene, but is due to is recessive to sexuality, although increasing the chromosome Summing up, we may say that as a rule the apomictic condition

APOMIXIS, HYBRIDIZATION, AND POLYPLOIDY

of apomictic reproduction, even though their parental species and other genera have in no instance shown any clear indications sexual species of Rubus (Peitersen 1921), Antennaria (Stebbins may be closely related to known apomictic forms. tion by itself can induce apomixis. Hybrids between different apomicts. Nevertheless, there is no evidence at all that hybridizaevidence has been obtained for the hybrid origin of many of the them. The latter situation has now been confirmed in many mictic groups and the difficulties of delimiting species in most of evidence by itself is by no means conclusive. Supporting evidence specific hybrids. As was pointed out earlier in this chapter, this groups besides those known to Ernst, and in some of them, such as produced by Ernst centered in the great polymorphism in apoforms, and often simulate the same phenomena tound in interdisturbances of meiosis and of other developmental processes in (Poddubnaja-Arnoldi 1939a,b, Koroleva 1939, Gustafsson 1947b), 1932a), Potentilla (Christoff and Papasova 1943), Taraxacum Rubus, Poa, Crepis, Parthenium, and Antennaria, convincing the development of the sex cells are widespread in apomictic between species. His evidence consisted mainly of the fact that Ernst (1918), who believed that it is caused by hybridization The first hypothesis developed to explain apomixis was that of

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apomixis and polyploidy. Many groups of plants are known in viviparum, Ranunculus ficaria, Cardamine bulbifera, Saxifraga apomictic polyploids occur with equal or greater frequency. In a in Allium, Agave, and Lilium, although in all these genera occur in diploid forms. Diploid species with vegetative apomixis plete; all of the different types of apomixis are now known to ploids, largely apomictic. This correlation is not, however, comwhich the diploid species are exclusively sexual and the polyspp., various species of Festuca, Poa, Deschampsia, and other larger number of genera with this type of apomixis (Polygonum by means of bulbils or similar structures ("vivipary") are known Gramineae), all, or nearly all, the apomicts are polyploid. A similar situation exists in respect to the relationship between

is the classic example of this situation, but others are Nothoscorembryony contain a relatively high proportion of diploids. Citrus which are not known to have diploid relatives also possessing it. nigra are the only polyploid species with this type of apomixis tious embryony occur also, but Ochna serrulata and Nigritella species of Eugenia. In the latter genus, polyploids with adventidum bivalve, Alnus rugosa, Sarcococca pruniformis, and some Among agamospermous apomicts, those with adventitious

apomicts are exclusively polyploid. Additional groups of this given by Stebbins (1941a) includes 24 of this type, in which the number the diploid ones, and polyploid apomicts also occur in forms are diploid. These are Potentilla, particularly P. argentea apomixis only three are known in which some of the apomictic nature which may now be cited are Parthenium (Rollins, In both Potentilla and Hieracium, the polyploid apomicts far out-Hieracium umbellatum (Gentscheff, cited by Gustafsson 1947b). (Häflinger 1943, cited by Gustaffsson 1947b), and a form of b), although the latter genus is badly in need of careful study. Catcheside, and Gerstel 1947, Esau 1944, 1946, Stebbins and (A. and G. Müntzing 1941), the Ranunculus auricomus complex Kodani 1944), Rudbeckia (Battaglia 1946a,b, Fagerlind 1946), Ranunculus. On the other hand, the list of apomictic groups Paspalum (Burton 1948), and probably Crataegus (Camp 1942a Among the numerous groups of apomicts with gametophytic

unlikely the assumption that polyploidy directly initiates apo-On the other hand, evidence obtained recently makes very

related to T. kok-saghyz and live in the same region are apomictic all of the triploid and tetraploid "species" which are closely mictic reproduction. Autopolyploids of Taraxacum kok-saghyz same species are apomictic. Parthenium argentatum, although closely related strains of the (1946) found complete sexuality in a high polyploid form of (Poddubnaja-Arnoldi and Dianova 1934). Similarly, Gardner by the writer (unpublished) show no signs of apomixis, although produced by Kostoff and Tiber (1939), by Warmke (1945), and

would be likely to survive in natural populations only if they to be controlled by the same genes. Since the action of any of the closely integrated with each other, and which might be expected occur regularly only in the triple recessive homozygote, aabbcc. cious) development of embryo and endosperm. Apomixis will and failure of fertilization; while C vs. c control failure of the egg B vs. b control fertilization of the egg (whether haploid or diploid) spectively, normal meiosis and failure of chromosome reduction; be essential to put together such gene combinations. were recessive. Either inter- or intraspecific hybridization would the individual unless the other two were present, such alleles three alleles a, b, or c would be lethal or strongly deleterious to Gustafsson points out that pairs B, b and C, c control processes to develop without fertilization vs. autonomous (usually precothree pairs of genes, as follows: Gene pairs A vs. a control, respecies. A genetical model illustrating how this could happen is either between species or between different forms of the same cycle are probably put together most easily by hybridization, and polyploidy, interspecific hybridization, and polymorphy, on presented by Powers (1945). He conceives of a minimum of place, the combinations of genes necessary to initiate an apomictic basis. Hybridization promotes apomixis in two ways. In the first the other, must be explained on an indirect rather than a direct The usual close association between apomixis, on the one hand,

as isolated instances in a few plant species. In Scolopendrium phytes by apospory, but these cannot produce new sporophytes the single recessive homozygotes postulated by Powers are known 1935b) recessive mutant forms are known which produce gameto-(Andersson-Kotto 1932) and Leontodon hispidus (Bergman Conditions similar to those which would be expected in two of

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without fertilization. According to the scheme of Powers, these are the occasional aposporous, but not parthenogenetic, individonly triploid offspring is probably also of this constitution, as argentatum, described in the previous section, which produces uals which have been found in Oxyria digyna, Antennaria dioica, would have the combination aaBBCC. The form of Parthenium can be considered to possess the combination AABBcc. arise, due to precocious division of the egg cell. Both these forms and he found in it a tendency to produce up to 15 percent of from a cross between a sexual biotype and an apomictic biotype, Håkansson (1943a) studied a sexual plant of Poa alpina derived Coreopsis bicolor, and Picris hieracioides (cf. Stebbins 1941a). haploid embryos by pseudogamous parthenogenesis. Hagerup (1944) found that in Orchis maculata haploid embryos frequently

situation in many apomicts may be even more complex. Each of in some instances be affected by more than one gene. the separate processes, and particularly the failure of meiosis, may developed for the genetic explanation of apomixis, although the Powers's hypothesis thus seems to be the most plausible one yet

selective advantage of a mechanism tending to preserve and reecotypes in Potentilla glandulosa (see page 108). The immediate ampla, as well as in hybrids between widely different sexual in the case of such crosses as Poa pratensis \times P. lapponica or P. logical conditions possessed by hybrid genotypes of which the through the greater vigor and tolerance of a wide range of ecoproduce such well-adapted genotypes is obvious. been demonstrated clearly by Clausen, Keck, and Hiesey (1945b) parents differ greatly in their ecological preferences. This has The second way in which hybridization promotes apomixis is

although apomixis can be induced in diploids by favorable gene apomixis, Gustafsson has reached the plausible conclusion that, work of Heilbronn (1932) on Polypodium aureum. The sporois stronger on the polyploid level than it is on the diploid level mutations, the action of many of these apomixis-inducing genes gametophytes by means of regular meiosis. Such normal gametophyte of this species normally produces haploid spores and The best evidence for this is obtained from ferns, particularly the phytes, like those of most ferns, cannot develop further unless the After reviewing all the evidence in relation to polyploidy and

similar tendency has been described by Springer (1935) in the organs, and they reproduce regularly by apogamety. A somewhat sporophytes. But in addition they can under certain environ moss Phascum cuspidatum. from tetraploid sporophytes, are almost incapable of producing sex gram, Fig. 36). Tetraploid gametophytes, produced aposporously their vegetative tissue, a process known as apogamety (see diamental conditions produce diploid sporophytes directly from eggs which are capable of fertilization and yield tetraploid pieces of leaves. These, like their haploid counterparts, produce phytes may also be formed aposporously by regeneration from egg is tertilized to form a new sporophyte. But diploid gameto-

normally sexual and quite different from those of Phascum gametophytes of Funaria, Bryum, and other genera of mosses are studies of Beyerle (1932), which have demonstrated that sporo cuspidatum. tion found that the tetraploid, octoploid, and even 16-ploid their capability for regeneration. Wettstein (1927) has in addiphytes of different fern species differ greatly from each other in of polyploidy is not a general one is evident from the comparative That this property of increasing apomixis with higher levels

vivipary or other forms of vegetative reproduction. reinforce the action of genes favoring such types of apomixis as vivipary. There is good evidence, therefore, that polyploidy can octoploid, and even decaploid forms exist which show no sign of such as F. rubra, F. californica, and F. arundinacea, hexaploid, crease of the chromosome number, but in other species of Festuca, noted that in Festuca ovina the degree of vivipary increases with inrectly responsible for the bulbillifery. Turesson (1930, 1931b) paniculatum, and he is of the opinion that the polyploidy was diby crossing two varieties of the diploid, nonbulbilliferous A ploid form similar to the bulbilliferous species Allium oleraceum In the flowering plants, Levan (1937b) has produced a poly-

of gametophytic apomixis, we should expect a particularly strong reinforcing action of polyploidy on genes for its component procapomixis would not be favored by polyploidy. But, in the case should expect, therefore, that the action of genes for this type of to be more common in diploid than in polyploid plants. We Adventitious embryony is the one type of apomixis which seems

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esses. There is, however, relatively little evidence for this from all apomicts of Hieracium subg. Pilosella it produces unreduced after artificial doubling of the chromosome number. The form archesporial cell degenerate and are replaced by embryo sacs and the doubled form, the products of meiotic division of the bryological studies showed that in both typical H. hoppeanum formed, and eggs from these develop parthenogenetically. Embryo sacs with the reduced chromosome number have been mosomes, which resemble the undoubled form. Apparently emfrom colchicine treatment also reproduces in this manner to a gametophytes by apospory. A form with 90 chromosomes obtained investigated is a pentaploid with the somatic number 45, and like have obtained evidence for a tendency away from nonreduction tendency for nonreduction and the formation of diploid gametothe angiosperms of the reinforcement by polyploidy of a genetic the flowering plants. Gustafsson does not cite any examples from meiosis, thereby producing embryo sacs with the reduced chroform, however, some of the nucellar cells expand and go through derived from the integument and the nucellus. In the doubled large extent, but it yields in addition offspring with 45 chrophytes. In Hieracium hoppeanum, Christoff and Christoff (1948) mosome number.

tained hybrids between a sexual race of this species with 2n = 24was produced by Håkansson (1943a, 1944) in Poa alpina. He obgenesis may be induced by increasing the chromosome number of the parental forms were not parthenogenetic at all. Håkansson complement of 30 chromosomes obtained from haploid gametes chromosomes and an apomict having 2n = 38. Some of the F₁ through parthenogenesis. Their sister plants with the normal same number of genes from their apomictic parent as do those receive twice as many genes from their sexual parent, but the on a genic basis, since the F1 offspring capable of parthenogenesis and Gustafsson assumed that this situation could not be explained ber, nevertheless yielded a high proportion of haploid offspring duced by meiosis embryo sacs with the reduced chromosome num-41 to 43 chromosomes. These plants, although they always proplants, resulting from the fertilization of unreduced egg cells, had which are purely sexual. Evidence that a tendency toward pseudogamous partheno-

In artificial autotetraploids of some purely sexual angiosperms eggs with the reduced chromosome number may develop parthenogenetically. This was found by Randolph and Fischer (1939) in maize and by Warmke (1945) in *Taraxacum kok-saghyz*. In every instance, the number of parthenogenetic diploids was very small, but nevertheless the fact is demonstrated that polyploidy may promote the autonomous development of the egg cell into an embryo.

Fagerlind (1944a) has suggested that in some, perhaps the majority of, agamic complexes the apomixis first arose on the diploid level. The present association between polyploidy and apomixis is believed to be secondary, and due to the fact that polyploids, particularly of unbalanced chromosomal types, can reproduce themselves more efficiently when apomictic than when sexual. This may be a partial explanation of the correlation in some groups, but it seems hardly applicable to such complexes as *Antennaria*, *Rubus*, *Potentilla*, and *Poa*, which contain a considerable number of sexual polyploids.

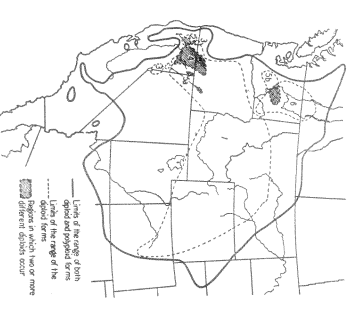
SOME TYPICAL AGAMIC COMPLEXES

of view, and variation patterns have been revealed which deviate considerably from those found in Crepis. The best conception of Crepis from the cytogenetic as well as from the taxonomic point tions of them have been studied much more intensively than studied in its entirety from both the systematic and the cytological Crepis. This complex remains the only one which has been characteristic variation pattern, not unlike that described in which these three sets of processes have been operating possess a Potentilla, Rubus, Parthenium, Taraxacum, and Hieracium, porbasis of their study of a typical example, the American species of fewer discontinuities between entities which can be recognized as apomicts and their closest sexual relatives. Species groups in point of view. But in other agamic complexes, particularly Poa, species. Such groups were characterized by Babcock and Stebbins Chapter VIII for polyploid complexes, but more intricate, and with the observation of anyone who studies thoroughly a group of hybridization, and polyploidy is so intimate that it cannot escape (1938, Stebbins and Babcock 1939) as agamic complexes, on the Whatever may be its causes, the connection between apomixis,

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the variation pattern in such complexes can therefore be obtained by summarizing first the situation in *Crepis*, and then indicating the ways in which various other agamic complexes depart from it.

The species of Crepis native to North America form a polyploid series based on the haploid number x = 11. Eight of them are diploid, but one, C. runcinata, is very different from the others and has no polyploid apomictic relatives. The agamic complex, therefore, is based on seven primary diploid species, C. pleuro-carpa, C. monticola, C. bakeri, C. occidentalis, C. modocensis, C. atribarba ("C. exilis"), and C. acuminata. With the exception of the last two, these are all much restricted in distribution, and appear as relicts, as shown on the map, Fig. 37. Although their ranges overlap to a considerable extent, they have very different



Ftc. 37. The relative areas of distribution of diploid sexual forms of *Crepis* species in western North America and of their polyploid apomictic derivatives. From Babcock and Stebbins 1938. Base map copyrighted by Rand McNally Company. Reproduced by special permission.

other hand, the highly restricted diploids, C. occidentalis and C. only species of which the nearest apomictic relatives have a narmost of these seem to contain some genes from C. acuminata, C modocensis, possess polyploid apomictic derivatives which are rower geographic distribution than the diploid form. On the the wide range of one diploid species, C. acuminata. This is the of their diploid ancestors, but not to a great extent, because of hypothesis is also supported by the fact that whenever the ecocytology, so far as it is known (Stebbins and Jenkins 1939). This to be allopolyploids, an assumption which is borne out by their of the characteristics of two or more diploids. They thus appear majority show in their external morphology various combinations which resemble closely each of the seven diploids, but the large numbers of 33, 44, 55, 77, and 88. All these polyploids which have numerous are the polyploids, which have somatic chromosome total population of Crepts in western North America. Far more geographic distribution, the apomicts as a whole exceed the range between which the apomict is intermediate morphologically. In they have been found to fall between those of the diploid species facultative apomicts. They include a certain percentage of forms bakeri, or both. found throughout most of the range of the complex, although logical and climatic preferences of an apomict have been analyzed, been tested prove to be apomictic, although some of them are These sexual diploids comprise only a minute fraction of the

those found in the diploid species, plus the effects of polyploidy. istics which cannot be explained as a result of recombinations of other diploid species, none of the polyploids possesses charactercomplex as a whole is that while each of the seven diploids possesses certain distinctive morphological characteristics not found in ficial resemblances caused the grouping together in the same systhe different apomicts could be perceived only dimly, and super-Until all the diploids were recognized, the relationships between The most important point about the variation pattern in this

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and recombinations of their characteristics. sections. They are connected by a much larger superstructure of with Old World species which are placed by Babcock in different other and so widely different that they appear to have affinities seven pillars, which by themselves are sharply distinct from each except that it is more complex. The diploids can be thought of as the different characteristics of various diploids. There thus apeither as apparent autopolyploids or as allopolyploids combining diploids permitted the grouping of the apomicts around them, ecological preferences. But recognition of the seven primary tematic category of forms with very different relationships and polyploid apomicts, which represent all sorts of intergradations peared a variation pattern which resembles that shown in Fig. 34

separate colony seems to possess its own cluster of distinctive but much less in the peripheral areas where only apomicts are diversity is very great in the regions occupied by the sexual species, apomicts. The apomicts are distributed in just the manner which hundreds of miles, while in the vicinity of the sexual types each clone can often be recognized over distances of scores or even found. In the latter regions, the same biotype or apomictic would be expected if they had been formed by hybridization and from the areas which the latter occupy. polyploidy involving the sexual species and had radiated outward Another feature of this pattern as observed in Crepis is that its

cent Canada. This makes it much easier to comprehend as a single arid transition zone of the northwestern United States and adja-Crepis has two advantages over most of the known agamic comunit than each of the widespread Holarctic or even bipolar comgraphic area and life zone, namely, the semiarid cold steppe or plexes. The total range of its complex is confined to one georise to it are all still in existence and are growing in the same acium. Secondly, it is so recent that the diploid species which gave found to disappear when these are as fully known. Crepis in the variation pattern of other agamic complexes will be In this writer's opinion, many of the apparent divergences from regions as their nearest relatives among the polyploid apomicts. plexes, such as Poa, Rubus, Antennaria, Taraxacum, and Hier-From the systematic and the phytogeographic points of view,

Although many agamic complexes have variation patterns essen-

species as well as diploid apomictic species. species; third, the relative frequency of facultative as compared to obligate apomixis; and, fourth, the presence of polyploid sexual species on which the complex is based; second, the extent to which certain important respects. These are, first, the number of sexual tially similar to that in Crepis, others deviate from this genus in the range of variation in the complex is covered by modern sexual

sexual species, but the range of variation in both of them is very and the former two are certainly larger than Grepis. The two has been undertaken in the case of Hieracium and Taraxacum thorough search had not been made for them. No such search Crepis are of this nature and would not have been found if a hardly justified, since five out of seven of the sexual species of because the sexual relatives of the apomicts are considered belong to a different type of agamic complex from that of Crepis tafsson (1947c) has considered that Hieracium and Taraxacum logical variability, as well as in its geographical distribution. Gusthree times as extensive as Crepis in the range of its morphodiploids undoubtedly exist. Taraxacum, therefore, is two or complex are either lacking or are very superficial, and many more diploid and sexual and to have close relatives among the apomicts. great. In Taraxacum, no less than twelve species are known to be one in subg. Pilosella, have been insufficiently explored as to their complexes of Hieracrum, one in the subgenus Archieracrum and diversity in the complex. The latter three are probably smaller, number of sexual species, and therefore in the size and degree of most nearly related to Crepis, namely, Taraxacum, Hieracium, (p. 254) to be "depauperate and relict." This distinction seems But cytological explorations over large parts of the range of this Chondrilla, Youngia, and Ixeris, differ from it chiefly in the So far as is known, the agamic complexes found in those genera

jority are obligate apomicts, but the high frequency in certain by diplospory and by autonomous parthenogenesis. The great maapomicts, which are mostly tetraploid and hexaploid, reproduce the one studied cytologically has fairly regular meiosis. The diploids have been found in nature (Stebbins 1932a, 1935), and different sexual diploid species. Hybrids involving three of these that of Taraxacum, since it is based on at least twelve to fifteen The agamic complex of Antennaria is nearly as extensive as

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of polyploid sexual species. Bergman (1935c) found 2n = 42 in many of the apomicts are intermediate between two or more species (A. parlinii, A. fallax), suggests that facultative apomicts areas of male plants in apparently apomictic forms of some of the staminate plants from an apparently sexual colony. both sexual and apomictic biotypes, the writer found n=28 in apomicts of arctic North America. In A. media, which possesses Antennaria carpathica, which may be related to some of the ther complication which may exist in this complex is the presence tinctions, just as in Crepis, Taraxacum, and Hieracium. A fursexual diploids, and they tend to obscure the original species distheir close relatives among the apomicts. In external morphology, occur in some regions occupied jointly by sexual diploids and

apomicts with numbers higher than 2n = 90 are dwarf aberrants. argentatum and P. incanum, and the polyploids include forms graphic range, but they occupy different habitats and nearly everyand P. incanum are sympatric throughout most of their geotypes which appear to be autopolyploid. Parthenium argentatum taking work of Rollins (1944, 1945a, 1946), are less common than tween them, although clearly identified by the brilliant and pains characteristics of the two sexual diploids, and intermediates beare grouped around two modes representing the morphological Crepis in the variation pattern of this complex is that the apomicts all the apomicts are facultative. The conspicuous divergence from diplospory followed by pseudogamous parthenogenesis, and nearly which are rare or absent in nature. The method of apomixis is with somatic numbers as high as 144, although nearly all the is possessed by the relatively restricted sexual forms of both P. entered into it. The basic haploid chromosome number, x = 18, area, and probably only two original sexual diploid species have its smaller size, since its total range is confined to a single desert where are distinct enough so that they can be recognized even by which neither of the parental species is found. people without botanical training. The largest stands of apomicts intermediate between these two species are in parts of Texas in Parthenium differs from all the complexes mentioned above in

similar to *Crepis* are those found in other genera of the family Compositae. The best-known agamic complexes in other plant The facts presented above suggest that the complexes most

published, cited by Gustafsson) has confirmed this supposition. sexual, but the cytological studies of Longley (1924) and Einset that many of the North American species are entirely or chiefly ments of Brainerd and Peitersen (1919, Peitersen 1921) indicate tion, with one series of species in Europe and western Asia and European members of this section. The hybridization experiplus a single species in Japan. Apomixis is predominant in the the other in eastern North America, from Canada to Guatemala, Moriferi of subg. Eubatus. This section has a disjunct distribufound chiefly in one of the six sections of a single subgenus, sect. several of the twelve subgenera of this large genus, apomixis is (1947b) suggest that apomictic forms also exist, and Einset (un-Gustafsson (1942, 1943a). Although polyploidy is known in Crepis in most or all of the four characteristics mentioned above. families, namely, those of Rubus, Potentilla, and Poa, differ from The best known of these is Rubus, due to the exhaustive study by

pollen that they frequently serve as the parents of hybrids are facultative, while even the obligate apomicts have such good occupied by the apomicts. Rubus differs also in the presence of allopolyploid of which the diploid ancestors are completely unseries centers about the anomalous species R. caesius, an apomictic reveal a sexual diploid with the characteristics necessary to exrepresented by the Glandulosi, has its center of variation in the existed in Europe at some time in the past. The second series, American sexual diploid R. allegheniensis, and it could be exchiefly by the group Suberecti, varies in the direction of the North of the sexual species in three directions. One series, represented apomicts. The range of variation in these forms goes beyond that count for the amount of variation represented by the polyploid extinct, but may have close relatives still existing in regions not the sexual diploid ancestors of its agamic complex appear to be known. Rubus, therefore, differs from Crepis in that several of plain the morphological characteristics of this group. The third Caucasus Mountains, and further exploration in this area may plained on the assumption that a close relative of this species the European Eubati, but shows that these will by no means acpolyploid sexual species, and in the fact that most of its apomicts Gustafsson recognizes five primary diploid sexual species among

The wide Holarctic range and disjunct distribution of the

section to which the agamic complex belongs indicates a greater Apomixis in Relation to Variation and Evolution 403

group, the Corylifolii, appear to be derived from recent (postover a much wider area than it is in Crepis. In fact, one whole tion between apomicts is taking place on a much larger scale and facultative apomixis, the origin of new forms through hybridizathough other sexual ancestors are apparently extinct. Due to the tributing genes on a large scale to its apomictic members, alplex have remained common, aggressive, and capable of conbut nevertheless some of the sexual members of the Rubus comgeological age than that of the North American section of Crepis, only half of the original sexual ancestors are now living within longing to other sections. Thus, although the total range of morhuman) hybridization between Rubus caesius and apomicts beturbed by man, has kept the complex of Rubus in a more "youththe range of the apomicts, the greater ability for hybridization, Rubus is about the same as that in Crepis, and although perhaps phological and ecological variability in the agamic complex of ful" self-perpetuating condition than is that of Crepts. plus the vigor of the plants and their adaptability to areas dis-

and pseudogamy among two diploid species complexes, those of unique in the great development of apomixis through apospory likewise the greatest known in any agamic complex. Potentilla is to 2n = 109 (the latter in the western American P. gracilis), is any other, and the range of chromosome numbers, from 2n = 14hensive treatment by a single author. The diploid species of the complexes, those of Potentilla and Poa, has been given compre-P. arguta (Popoff 1935) and P. argentea (A. and G. Müntzing Potentilla complex may prove to be more numerous than those in facultative, and hybridization between them is common in nature. 1941). Like the apomicts of Rubus, those of Potentilla are largely cise relationships between them have not been worked out, so species of Potentilla related to the apomicts are known, the prehybridization between two other groups. Although sexual diploid Gustafsson suggests that one whole group of apomicts, the that the structure and history of this agamic complex is likewise Collinae, like the Corylifoliae of Rubus, has arisen through recent Neither of the two remaining well-known and extensive agamic

The genus Poa may contain one or several agamic complexes.

over two world-wide genera. is not inconceivable, therefore, that this agamic complex is spread Puccinellia, which also contains species that may be apomictic. It Scabrellae, resemble in some respects the neighboring genus more, two North American sections, the Nevadenses and the would be undoubtedly the largest in the plant kingdom. Furtherconnected and to form a single gigantic agamic complex. This and productive of apomictic seed (Clausen, Keck, and Hiese) each other, and wide intersectional hybrids prove to be vigorous genus, and since these sections are very difficult to separate from 1945b), all the apomicts of Poa may eventually prove to be inter-Apomixis is known or strongly suspected in seven sections of this

a large extent in geographic distribution. Although additional diploid sexual species of Poa will almost certainly be discovered ably the highest. Poa, therefore, has more in common with Potenmost of those which contributed originally to the agamic complex tilla than with any other agamic complex, and agrees with it to 2n = 42 to 2n = 104, and a mode at 84 (Hartung 1946), is probtion. That in *P. alpina* is probably the lowest; that in the Scabrel. genesis. The range of chromosome numbers varies with the section); these must be obligate apomicts with autonomous partheno-Several species of western North America, such as P. nervosa and species are viviparous; the best-known method, found in P. apomixis present varies from one group to another. Several series, those of P. alpina and of P. pratensis. In both of them and a sexual diploid of P. alpina reported by Christoff (1943). some of its apomicts. These are Poa trivialis (Kiellander 1942) only two sexual diploids are known which are likely ancestors for lae of western North America, with somatic numbers from pollen over large sections of their range (Keck, oral communica-P. epilis, consist entirely of plants with abortive anthers and no parthenogenesis; and other mechanisms are likely to be found. pratensis and P. alpina, is apospory followed by pseudogamous 31, 33, and 38 in European strains of P. alpina. The type of lar, Müntzing (1933) has found somatic numbers of 22, 24, 25, there is the additional unusual feature of aneuploidy. In particu-Furthermore, sexual polyploids form a conspicuous feature in two plex of Alchemilla (Gustafsson 1947c, p. 254), is distinctive in that Poa, along with the apparently similar but little-known com-

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are probably extinct. Furthermore, many polyploid sexual species of Poa are known, and these may have contributed very largely to and which contains in addition obligate apomicts. Like Rubus which is in part purely sexual, in part facultatively apomictic, may have to be regarded as a single huge polyploid complex, the agamic complex. In fact, when this genus is better known, it activity of the facultative apomicts. youthfulness and its ability to produce new forms through the and Potentilla, the bulk of this agamic complex has retained its

a single one would seem to be rather difficult to make in many incomplex." The distinction between two merging complexes and more different complexes which merge into one another," while that (Gustafsson 1947c, p. 235) "The apomicts belong to two or apomictic complexes." They are characterized by the statement tafsson in the second of his two series of "apomictic and amphi seem to the present writer rather heterogeneous, and many of stances. The groups which Gustafsson places in his first series in the first series (p. 222) "the apomicts are included in a single them are relatively little known. Some, like Ranunculus ficaria, agamic complexes similar to the larger ones described above. Hypericum perforatum, may be relatively small and simple Allium carinatum-oleraceum, Saxifraga stellaris-foliolosa, and cernua, and Gagea spathacea, may be apomictically or asexually crassipes, Polygonum viviparum, Cardamine bulbifera, Saxifraga caespitosa and D. alpina, Nigritella nigra, Festuca ovina, Stellaria ships have not been well studied. Still others, like Deschampsia genera, and may appear to be isolated only because their relationportions of the main agamic complex found in each of these Others, like Rubus idaeus, Poa bulbosa, and P. compressa, may be reproducing units of otherwise sexual polyploid complexes. as known is an isolated apomictic species without living sexual Finally, there is the example of Houltuynia cordata, which so far relatives, and may be the last relict of an ancient agamic complex (Babcock and Stebbins 1938). The agamic complexes just discussed are all classified by Gus-

CAUSES OF VARIATION IN AGAMIC COMPLEXES

sible an explanation of an apparent paradox mentioned by several The information reviewed in the preceding section makes pos-

complex; second, hybridization between facultative apomicts allopolyploidy between the original sexual ancestors of the agamic species; third, chromosomal and genic changes within the apomic with the resulting segregation, or between apomicts and sexua morphy may be summarized as follows: first, hybridization and group in which this phenomenon occurs. The causes of this polyreproduction results in a great increase in polymorphy of the authors, namely, that partial or complete abandonment of sexua tic clones themselves.

a sexual group. It would crystallize an initial polymorphism some natural habitat. Thus, the complex would be likely to poscross types, many of which could be vigorous and well adapted to to most agamic complexes are rather closely related to each other, arrival of this process, would have disappeared which existed before the onset of apomixis and which, but for the genetic potentialities for apomixis, this process would tend to perselective value. Therefore, in any group which possesses the tially sterile hybrid derivatives. In such types gene combinations sess a high degree of polymorphism, particularly among the parparents, and so to produce a strongly segregating series of backticularly when fertilized by pollen from one or another of their their F₁ hybrids might be expected to set occasional seeds, parfavoring apomixis. Nevertheless, since the sexual species ancestral this initial hybridization were undoubtedly highly sterile, except in the past between the ancestral sexual species. The products of known agamic complexes extensive hybridization has occurred is no denying the fact that in the great majority, if not all, of the and apomixis are still to a certain extent matters for debate, there petuate types which because of their sterility would not persist in bridization with apomictic forms, would have a particularly high producing apomixis, whether acquired by mutation or by hyfor those which happened to acquire a combination of genes Although the causal relationships of hybridization, polyploidy,

summarized a series of hybridization experiments in these general apomicts are facultative. Gustafsson (1947b, pp. 141-146) has and sexual forms, or between different facultative apomicts, has ticularly in Rubus, Potentilla, and Poa, in which nearly all of the been demonstrated experimentally as a cause of variability, par-The second process, hybridization between facultative apomicts

> all of which have had essentially similar results. Hybridization Apomixis in Relation to Variation and Evolution 407

a great range of segregant types in the F2 generation. Many of may be restored. Such plants, if they find an ecological niche to these are sterile, inviable, or both, but others are vigorous, and between two apomictic forms usually yields sexual F1 plants and clones. Facultative apomixis may therefore give a pattern of which they are adapted, may be the progenitors of new apomictic in a few of the individuals in the F2 or later generations apomixis species the modes of variation represented by genetically isolated types of reproduction is that in predominantly self-fertilized apomictic clones, like that of pure lines, is periodically interrupted that found in self-fertilized organisms. The constancy of such variation and a type of evolution which is in some ways similar to reproductive ability of sexually sterile hybrid derivatives tends to species are usually maintained, while in agamic complexes the The major difference between the patterns created by these two by bursts of new variability and selection, caused by hybridization.

obscure or to obliterate the original species boundaries. obligate apomicts. In the genus Hieracium, the classic hybridization experiments of Mendel, as well as the more recent ones by obligate apomixis has set in, new variability through hybridizaapomicts in later generations. It is obvious, however, that when others may segregate further and produce still more variant apomictic and may give rise immediately to new clones, while great variety of segregant types in the F_1 generation, because of the tilization of a sexual species by pollen from an apomict yields a Ostenfeld (1910) and by Christoff (1942), have shown that ferheterozygosity of the parents. Some of these are wholly or partly species occur and to those apomicts which still have functional tion and segregation is confined to the regions in which sexual and probably Taraxacum. tilla, and Poa, while the latter is the case in Crepis, Antennaria, species. The former situation appears to be true in Rubus, Potenwould be highly concentrated in those regions containing sexual out the geographic range of the complex, while in the latter it In the former, variability would be rather evenly spread throughhaving facultative as compared to those with obligate apomixis. difference between the variation patterns of agamic complexes pollen. For this reason, we should expect to find a conspicuous Variability resulting from hybridization is still possible in some

naria, Taraxacum, Hieracium, Calamagrostis, and other comof Stebbins (1941a, p. 531). There is good reason, therefore, for normal meiosis. However, precisely this situation has been found same plant with the formation of reduced megaspores through plexes with obligate apomixis was produced in an earlier stage of believing that much of the polymorphism now found in Antenin Parthenium argentatum (Esau 1946), fulfilling the prediction apospory, and obligate apomixis with diplospory. He maintained Hieracium" and that this type of division cannot occur in the peculiar kind of parthenogenesis occurring in Antennaria and that "an entirely new type of division had to be introduced in the the observation that facultative apomixis is usually associated with greater in complexes containing facultative apomicts as compared their evolution, when facultative apomixis was still widespread directly in Taraxacum and Hieracium. He based this belief on But Gustafsson (1935a) believed that the obligate apomicts arise bins (1938, Stebbins 1941a) on the basis of their study of Crepis. facultative apomicts. This was postulated by Babcock and Stebligate apomicts do not arise directly from sexual forms, but from the case. The probable explanation of this situation is that obto those with obligate apomicts, but this does not appear to be The total amount of polymorphism would be expected to be

which will cause the egg to have one or more chromosomes less apomicts. Second, chromosomal aberrations may occur, particucalled the phenomenon autosegregation and discussed several of restitution nuclei or by some other means. Crane and Thomas than the normal number. The best example of this is the series of larly in the divisions leading to the diplosporous embryo sacs, cytological mechanisms, none of which is known to occur in these apomictic progeny of Parthenium. Gustafsson (1943, 1947c) in Rubus, but do not present cytological evidence. Rollins diploid chromosome number in the egg through the formation megaspore mother cells, with subsequent restoration of the within the apomicts. The first is that postulated by Darlington (1945b) has suggested a similar explanation for variation in the (1937) as a result of meiotic pairing and crossing over in the (1939) postulate this mechanism for the origin of certain variants Various types of chromosomal and genic changes can occur

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monosomics found by Sörenson and Gudjónsson (1946) in Taraxacum. In Taraxacum, as in most other genera, however, these chromosomal aberrants are less viable than euploid plants, and for this reason do not become established in nature. Only in Poa and in Potentilla are aneuploid apomicts abundant. Finally, apomictic clones may produce new variants by means of somatic mutations. These have been described as occasional occurrences in Rubus (Gustafsson 1943, p. 78), but they are better known in Citrus (Frost 1926). Because of the genetic heterozygosity of most apomicts, such mutations will usually affect the appearance of the phenotype, and if they occur in the tissue of the inflorescence or flower, they will produce mutant offspring.

SPECIES CONCEPTS IN AGAMIC COMPLEXES

have pointed out, such a species concept cannot be applied to genes. As Dobzhansky (1941) and Babcock and Stebbins (1938) truly biological concept of species, centers about the possibility the concepts maintained by the majority of those who desire a such genera as Taraxacum, Hieracium, Antennaria, Rubus, Poand which without apomixis would not be able to persist because have combined the genes of previously isolated sexual species, while the origin of many apomictic clones is from genotypes which is prevented by the very nature of their types of reproduction, agamic complexes. Free interchange of genes between apomicts the separation of different species by barriers to the exchange of for exchange of genes between members of the same species and tists have not been able to agree on the boundaries of species in of their sexual sterility. It is not strange, therefore, that systematentilla, Poa, and Calamagrostis. In attempting to set up species cism is that the number of apomicts in any well-developed comand Turesson (1931), by Fernald (1933), by Turrill (1938c), by better concepts of the variation patterns in the genera concerned. clone of Taraxacum, Hieracium, or Rubus likely to provide any which in the biological sense are not there. Nor are those like those found in sexual groups, they are looking for entities Criticisms of this method have been given by Müntzing, Tedin, "splitters" who make a separate species out of every apomictic Babcock and Stebbins (1938), and by Stebbins (1941a). One criti-The species concept developed in Chapter VI, which resembles

sister individuals of any sexual progeny (see diagram, Fig. 38). other in precisely those characteristics which segregate among the ternal morphology from their sexual ancestors, differ from each generation. A reductio ad absurdum would be reached by the only partly or facultatively apomictic. From time to time they apomicts, and in nearly all complexes at least some of them, are Crepis acuminata. Here the apomicts, indistinguishable in extreatment as separate species of the apomicts in an entity like parable amount of genetic variation occurs regularly in each dividual. Such forms may have sexual relatives in which a comclones, or "species," may arise in the offspring of a single inreproduce sexually, and on such occasions a whole series of new plex is so large that recognizing them as separate species makes the the fact that in many complexes the great majority of the comprehension of the group as a whole difficult or impossible; "one cannot see the forest for the trees." But more important is

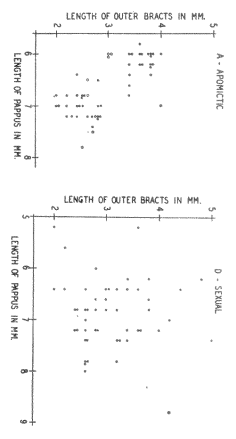


Fig. 38. Diagrams showing the range of variation in respect to length of outer involucral bracts and of pappus on the mature fruits within a colony of apomictic and of sexual *Crepis acuminata*. From Babcock and Stebbins 1938.

As Babcock and Stebbins (1938) have pointed out, the needs of classification and the clearest conception of the structure, interrelationships, and phylogeny of the agamic complex are obtained by drawing arbitrary species boundaries which are related to the original boundaries of the sexual species ancestral to the

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than the number of species recognized should not be smaller than the number of sexual ancestors which are known to exist or can be reasonably inferred to have existed in the past. In most complexes, furthermore, the number of species should be larger, since groups of apomicts combining the characteristics of sexual species which are widely different from each other should be recognized as agamospecies (Turesson 1929). The boundaries between these species must be drawn where most convenient, usually on the basis of one or a few easily recognized "key characters" which are known to separate the ancestral sexual species.

If, as in *Rubus* and in *Poa*, the sexual ancestors of the complex are mostly extinct, this method is difficult or impossible to apply. Here the method, suggested by various European systematists and followed by Gustafsson (1943, 1947c, p. 245), of recognizing as "circle microspecies" groups of apomicts morphologically similar to each other, with each species centered about one or a few particularly widespread and common apomicts, is perhaps the only feasible one. Consciously or unconsciously, it has been regularly used in practice. Nevertheless, the use of such a method must be recognized simply as a means of bringing some type of order to a situation which from the biological point of view is incapable of resolution. Disputes between systematists not fully conscious of the biological situation in these groups are particularly academic and futile.

AGAMIC COMPLEXES AND PLANT GEOGRAPHY

There are some features of agamic complexes which make them even better tools for analyses of plant distribution than are sexual polyploid complexes (Gustafsson 1935b, 1947c; Turrill 1938c,d; Babcock and Stebbins 1938; Stebbins 1941a). In the first place, the apomicts must have arisen from sexual ancestors and cannot have diverged very far from them in morphological characteristics. The identification of these ancestors should in most instances be possible by careful morphological and cytogenetical studies. If the apomicts are of allopolyploid origin, the same inferences may be drawn about past distributions of the sexual diploid members of the complex as is possible with sexual polyploids (see page 350). Secondly, the relative constancy of apomicts through hundreds or even thousands of generations makes them valuable indicators of

ecological conditions, both in their present habitats and in past migration routes.

In the case of agamic complexes like that of *Crepis*, in which all of the sexual ancestors still exist and are concentrated in one or a few geographic areas, these regions must be regarded as primary centers of differentiation, from which migrants have traveled to the peripheral portions of the range of the complex. Obviously, one method of ascertaining centers of migration for entire floras would be to analyze the distribution of the sexual members of all the agamic complexes found in them. This might be particularly successful in floras like those of the arctic regions, which contain many agamic complexes, and in which present distribution patterns are the result of relatively recent migrations. Two suggestions are already possible in this connection.

anywhere else, the apomicts of the A. alpina complex are best exfined to Alaska, and A. monocephala is more abundant there than derivatives of these two sexual species. Since A. alaskana is conplained as radiants from the Beringian center of Hultén (1937). the arctic American "species" related to it are allopolyploid these three makes rather likely the hypothesis that A. alpina and to A. alpina and its relatives are A. monocephala and A. alaskana dance of male plants) which are morphologically most similar alpina group, of which the sexual members are confined to the (Malte 1934). Comparison by the present writer of specimens of New World. The probable sexual species (judging by the abunthe most common apomicts in the arctic regions belong to the A. radiated in postglacial times. In Antennaria, on the other hand therefore the most likely center from which the arctic Taraxaca particularly the Thian Shan Mountains, studied by Poddubnajarelatives among the sexual species in those from central Asia, regions, belonging to the section Ceratophora, have their closest Arnoldi and Dianova (1934). This central Asiatic refugium is In Taraxacum, the most numerous apomicts of the arctic

As described above, the agamic complex of *Rubus* is one in which certain ancestral diploid sexual species are no longer living in the region occupied by most of the apomicts, but occur in North America. On these grounds, Gustafsson has made the plausible inference that at some time in the past, probably during the latter part of the Tertiary period, some species of *Rubus*,

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morphologically and ecologically similar to the modern North American R. allegheniensis or R. argutus, existed in Europe, so as to give rise to the Suberecti and related sections. This evidence is in line with much of that from paleontology which indicates that the Tertiary floras of the Old and the New Worlds had considerably more in common than do the modern ones, and it runs parallel to the evidence from Oryzopsis allopolyploids (see page 353) that Eurasian species of this genus were once represented in North America.

complexes, has any general scientific value, it is in this direction. of Crepis in a similar manner in connection with the flora of the solve certain problems of local distribution in Scandinavia, while indicators of past migration routes of floras. Gustafsson (1935b, many of them may be determined, makes them particularly good the endless "microspecies," or apomictic clones of the agamic western United States. If the detailed study and classification of 1947c) used apomicts of Taraxacum, Hieracium, and Poa to help species in the temperate zones of the northern and southern apomicts may be valuable, is that of the disjunct ranges of many Babcock and Stebbins (1938, pp. 39-44) have used the apomicts subarctic regions of the Northern Hemisphere, indicates that naria occur in temperate and subantarctic South America, while hemispheres - the "bipolar" distributions discussed by DuRietz A broader problem of this nature, toward the solution of which North America. such regions, from which they are at present completely absent could be obtained by suitable experiments on growing them in their ability to cross the tropics along the crests of mountain ranges dispersed southward by long-distance migration. Some idea of similar to those found in their present habitat, unless they were followed a route which had ecological conditions essentially these apomicts must somehow have migrated across the tropics their possible sexual ancestors occur only in the temperate and (1940). The fact that apomicts of both Taraxacum and Antenpossessed by most apomictic clones, we must suppose that they Because of the relatively narrow limits of ecological tolerance nating and puzzling problem of plant geography is open to experimental attack. By the use of apomicts, therefore, at least one phase of this fasci-The constancy of apomicts, and the fact that the ancestry of

denly made available. They represent evolutionary opportunism cently been subjected to alternations of glacial and temperate are prominent in the arctic and subarctic floras which have resuch genera as Taraxacum, Antennaria, Hieracium, Rubus, and carried to its limit. It is no accident, therefore, that apomicts of species are still further increased, so that agamic complexes are the colonization of newly available habitats. With apomixis, the mits the "pooling" of the genetic resources of two or more species to survive were it not for this process. In a group of strictly Crataegus have "weedy" tendencies, and that agamic complexes the rapid evolution of new genotypes adapted to new habitats sudeven better equipped than are sexual polyploid complexes for possibilities for pooling the genetic resources of several distinct better equipped than a group of comparable diploid species for fertile and constant, and to that extent a polyploid complex is to an extent limited by the factors which render allopolyploids barriers which bound the different species. Allopolyploidy pertion to new habitats must be formed from a series of separate diploid sexual organisms, new combinations of genes for adaptaings, but which because of their sexual sterility would be unable genotypes which are vigorous and well adapted to their surroundoccurs. In the first place, it makes possible the survival of many "gene funds," each of which is largely limited by the isolating Apomixis has three principal effects on the groups in which it

Thus, the concept of Darlington (1939), that apomixis is an "escape from sterility," can be taken only in the relatively limited sense that it permits the survival of well-adapted, but sexually sterile, genotypes. Gustafsson (1947b, p. 173) has pointed out that many apomicts are still able to produce viable and fertile sexual offspring, so that the selective value of apomixis cannot be confined to this function.

The second effect of apomixis is similar to that of self-fertilization; it permits the building up of large populations of genetically similar individuals for the rapid colonization of newly available habitats. Thus the "opportunism" of agamic complexes results both from the increase in the number of adaptive genotypes available and from the ease with which such genotypes may be per-

reproduction. Over a short period of time, agamic complexes are capable of a rapid burst of evolution in terms of the production

and establishment of new genotypes. But once an agamic com-

plex has come to consist chiefly of apomicts, its evolutionary future is decidedly limited. If obligate apomixis has become predominant, new evolutionary progress is bound up with the dis-

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petuated. The fact is doubtless significant that apomixis and self-fertilization appear as mutually exclusive genetic systems. So far as the present writer is aware, the sexual ancestors of all known agamic complexes are self-incompatible, dioecious, or possess other mechanisms which promote allogamy. Apomixis has not developed in self-fertilizing plants.

which promotes self-fertilization. Data presented in Chapter V habit which favors apomixis is apparently the opposite of that annuals and in short-lived perennials, and is rare or absent in (see Table 4) show that self-fertilization is predominant in is usually associated with sexual sterility, either through interspewhile the acquisition of an agamospermous method of apomixis methods of vegetative reproduction, such as rhizomes or stolons. are nearly all perennials, and many of them possess accessory phasized the fact that the sexual ancestors of agamic complexes reproduction. On the other hand, Gustafsson (1948a) has emperennials with rhizomes or other effective means of vegetative cific hybridization or through genetic factors causing a disturbance genetic mechanism which does not necessarily affect fertility, from cross- to self-fertilization can be accomplished by a simple reproduction preadapt plants to these conditions. It is probof partial sterility, and efficient vegetative means of growth and of meiosis. So, just as in the case of polyploidy, the evolutionary self-fertilization the constancy which helps them in colonization. plants of such habitats are mostly annuals, and acquire through among the weedy flora of cultivated fields and roadsides. The to promote weedy tendencies, apomicts are not conspicuous ably for this reason that, although apomixis would be expected line which is acquiring apomixis must pass through a bottleneck This situation is probably explained by the fact that the change bility imposed on those plants which have adopted this method of One probable reason for this situation is that the type of growth The third effect of apomixis is the limitation of genetic varia-

of its sexual ancestors. As long as recombinations of these ances dominant obligate apomixis is a closed system which can produce aggressive, weedy tendencies are evident in those apomicts of ually decline and die out. genetic recombinations are suited, the agamic complex will grad has started to encroach on the environments to which these to some available environment, evolution can continue, but when little or no variation beyond that already present in the sum total carpa and C. monticola are themselves relictual types, showing Crepis related to the widespread sexual C. acuminata, while the tribution and potentialities of the remaining sexual species. If climatic changes or the evolution of more efficient competitors tral characteristics will yield gene combinations which are adapted little or no tendency to spread. An agamic complex with preapomicts related to the highly restricted sexual species *C. pleuro*logically, the evolutionary plasticity of the complex is lost, and these species become much restricted geographically and eco-Babcock and Stebbins (1938, pp. 61-62) have pointed out that it can no longer evolve in response to changing environments.

apomixis there are some groups of apomicts which appear as "confoundland and eastern Quebec (Fernald 1933). above and certain forms of Antennaria and Taraxacum in Newservative," relictual types, such as the Crepis apomicts mentioned their aggressive tendencies, while in most complexes with obligate elsewhere on the earth. Furthermore, the apomicts of these comsingle hemisphere or to one hemisphere plus arctic-alpine regions gate apomixis show their relative youth by their restriction to a ter XIV). On the other hand, most of the complexes with oblitribution along with the mid-Tertiary Holarctic flora (see Chapboth hemispheres and appear to have acquired their initial disold, since they are well represented in the temperate regions of tion, the complexes of Rubus, Potentilla, and Poa are relatively contain chiefly facultative apomicts, the potentiality for new variajusted balance between apomictic and sexual reproduction (Gusplexes have a longer lease on life, acquired by a favorably adtion is independent of the fate of the sexual species. Such complexes with facultative apomixis have for the most part retained tafsson 1947b, pp. 173-178). Judging from their present distribu-In complexes like those of Rubus, Poa, and Potentilla, which

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Nevertheless, even in complexes with facultative apomixis, there is little evidence that they can give rise to anything except new recombinations of the genes present in the original sexual ancestors. There is no evidence that apomicts have ever been able to evolve a new genus or even a subgenus. In this sense, all agamic complexes are closed systems and evolutionary "blind alleys." It is true, as Gustafsson (1947b, p. 178) has maintained, that both sexual and apomictic groups are born, live their span upon the earth, and sooner or later die out. But other things being equal, the life expectancy of agamic complexes is shorter than that of sexual groups. Furthermore, while sexual species may, during the course of their existence, give rise to entirely new types by means of progressive mutation and gene recombination, agamic complexes are destined to produce only new variations on an old

APOMIXIS AND PLANT BREEDING

crop plants are the various species of Citrus, but apomixis is to studies of this phenomenon. The most important apomictic valuable crop plants has rendered a more than academic interest known or suspected in other cultivated fruits, such as the mango should include among their preliminary explorations studies de if it belongs to the Gramineae, the Rosaceae, or the Compositae, a program on some group unknown cytogenetically, particularly from those on sexual plants, and plant breeders who are planning breeding programs on such groups must be conducted differently of ornamental shrubs, particularly Malus (crabapples, Dermen also in Paspalum (Burton 1948), while it occurs in some groups forage grasses, it is conspicuous in Poa pratensis and apparently berries (see above). It occurs in the rubber-bearing plant guayule 1937), the mangosteen (Sprecher 1919, Horn 1940), and black-(H. J. Webber 1931, Juliano and Cuevas 1932, Juliano 1934, signed to determine whether or not apomixis is present. 1936b) and Eugenia (Pijl 1934, Johnson 1936). It is obvious that Taraxacum related to another rubber plant, $T.\ koksaghyz.$ Among (Parthenium argentatum, see above), as well as in species of The fact that apomixis occurs in a number of economically

As is evident from the discussion in this chapter, the presence or absence of apomixis in a group cannot be determined simply by

ceding section, these two processes do not usually occur in the and if the resulting progeny are uniform, self-fertilization rather should be examined for growing pollen tubes. If this is found, strictly maternal individuals, apomixis can be suspected. If the scale, and the stigmas of the emasculated flowers should be polthan apomixis is to be suspected; since, as mentioned in the pretions of the styles and the stigmas of flowers bagged at anthesis emasculation is possible, this should be carried out on a large most groups provide reasonably decisive indirect evidence. If conducted breeding tests on a sufficiently large scale should in same group. the microscope to see whether it stains well, and stained dissecis impracticable, the pollen should at least be examined through flowers and anthers of the species are so small that emasculation gene, as was done by Burton (1948) in Paspalum. If cross-polpollen from a different genotype bearing a dominant marker megaspore, embryo sac, and embryo development. But properly linations of this type repeatedly yield a consistent percentage of linated with either a foreign species pollen or, preferably, with be obtained only from laborious and time-consuming studies of Positive evidence for the presence or absence of apomixis can

In groups known to be apomictic, the degree of apomixis should be determined by appropriate progeny tests (Powers and Rollins 1945), since facultative apomicts can be treated entirely differently from obligate ones. Fortunately, all the economically important groups mentioned above are, so far as is known, facultatively apomictic, and interspecific hybrids have been obtained in many of them. If any economic group is found to contain mostly obligate apomicts, the genus should be explored for sexual species, and pollen from these should be employed for breeding whenever

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possible. Even in groups containing mostly facultative apomicts, the sexual species may contain a supply of genes for disease resistance and other valuable properties which can by no means be ignored. Obviously, a study of the method of inheritance of apomixis in such groups is one of the next essential steps, so that estimates can be made of the extent to which apomixis will be recovered in the progeny of hybrids.

sexual ones. Tinney and Aamodt (1940) in Poa pratensis and viously be conducted on an entirely different basis from that in of normally apomictic 72-chromosome plants. These haploids inmosome "haploids" which occasionally develop from reduced eggs useful source of new genetic variability, namely, the 36-chroing could be carried out as in Poa. But here there is a particularly sexual plants found in most strains. In Parthenium, variety testprocess of establishing numerous clonal divisions from a single possible the testing of some varieties without the time-consuming tatum have made some suggestions as to how this can be done. In Rollins, Catcheside, and Gerstel (1947) in Parthenium argenmosomes. Since genetic segregation has occurred both in the duced gametes; therefore, when pollinated by 72-chromosome plant. Selection of new strains could be made from the occasional Poa, the genetic constancy of seed produced by apomixis makes plants they produce some offspring with this number of chroherit from their mothers the tendency to produce chiefly unrevariation can be expected among the progeny of haploids polthe pollen grains which fertilize its eggs, considerable genetic intensive studies of the fundamental nature of apomixis in other with similar properties. Nevertheless, there is little doubt that needed for testing, increase, and commercial use of a variety may taining both the variability needed for selection and the constancy will breed true through apomixis. This ingenious scheme for oblinated by diploids. But each individual of this variant progeny production of the egg which gave rise to the haploid plant and in groups of economically important plants will be a valuable and be applicable only to guayule and to a few other apomictic species perhaps essential prelude to the breeding of improved varieties Selection and progeny testing in apomictic groups must ob-