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A GENETIC STUDY OF SYMBIOTIC NITROGEN FIXATION
IN THE TROPICAL LEGUME DESMODIUM

by



BRIAN ROY PINCHBECK

A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH
IN PARTIAL FULFILMENT OF THE REQUIREMENTS FOR THE DEGREE
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ABSTRACT

A genetic study was initiated in the tropical legume Desmodium to establish whether there existed genetic variation in the symbiotic relationship between Desmodium and Rhizobium. Particular emphasis was placed upon effective vs. ineffective nodulation and quantitative variation for the estimates of nitrogen fixed.

Analysis of F_2 and backcross populations from two interspecific crosses suggested that a strain-specific ineffective nodulation interaction was conditioned by a dominant allele at a single locus.

Significant genetic variation was found among genotypes for all nitrogen fixation estimates. The major portion of the variation among genotypes was non-additive variation and was attributable to a significant Varieties vs. Crosses genetic component. The mean of the F_1 crosses was found to be 80% larger than the mean of the inbred parents indicating that most F_1 crosses exceeded their larger parent for the quantitative estimation of nitrogen fixation. Since inbreeding depression for forage yield is generally considered to be absent in the presence of adequate nitrogen, the F_1 hybrid vigor for the nitrogen fixation estimates was due solely to these differences when the plants were grown in a nitrogen limiting environment. The significance of these results was discussed.

Large cotyledons and juvenile leaves appeared to be controlled by dominant genes at different loci. Plants with large cotyledons or juvenile leaves fixed significantly more nitrogen than plants with small ones. Adjusting the quantitative estimates for differences due to cotyledon and juvenile leaf size produced a totally additive model for the inheritance of the estimates of nitrogen fixation. The non-additive variance for the unadjusted estimates could be divided into two components: (a) additive variation due to plant vigor, and (b) additive variation due to 'efficiency' of nitrogen fixation. A breeding program utilizing these additive components was discussed.

Genotype x environment interactions were of significant importance suggesting that study concerning quantitative variation for amount of nitrogen fixed should be conducted in different environments with various Rhizobium strains.

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INTRODUCTION

The development of tropical agriculture has recently commanded much interest in an effort to solve pending problems of world food shortages. The improvement of tropical legumes can contribute to this development through the improved symbiotic fixation of atmospheric nitrogen. It has been conceded that the amounts of nitrogen fixed by tropical legumes were generally less than for temperate legumes (Vincent, 1974). Since temperate and tropical legume species differed in the amount of nitrogen fixed, it would be pertinent to inquire as to whether genotypes within a species also differed. If genotypes differed significantly, can this variation within the host for symbiotic nitrogen fixation be exploited in a breeding program for the improvement of tropical legume species? It would be of particular interest to initiate a genetical study of the inheritance of symbiotic nitrogen fixation in the tropical legume Desmodium because of its great variation in the morphological differentiation of the numerous species (Bryan, 1966) and its potential as a forage legume (Bryan, 1969).

Norris (1956) postulated that the legume-Rhizobium symbiosis originated in tropical rain forests. As the legume adapted to the subtropical and temperate regions, the symbiotic interaction appeared to become more specific. Since the Rhizobium which nodulates Desmodium is a member of the "Cow-pea" group, a promiscuous interaction group of

Rhizobium, the Desmodium - Rhizobium symbiotic interaction may represent a more primitive form of symbiosis. However, Bryan (1966) reported that this interaction appeared to be specific as measured by numerous ineffective nodulations.

A. Generalized description of Desmodium.

Desmodium is a self-fertile tropical legume with a somatic chromosome number of $2n=22$ (Rotar and Urata, 1967). Rotar and Urata (1967) reported collections of many different species from tropical countries including two species from Canada. McWhirter (1969) suggested that the center of origin of Desmodium was in South America. The taxonomic identity of this genus is uncertain (McWhirter, 1969) and voucher specimens have been deposited by some Desmodium researchers (Rotar and Urata, 1967; McWhirter, pers. com.) at the Harvard University Herbarium for identifications by Dr. Bernice Shubert, Arnold Arboretum, Harvard University.

Four Desmodium species have attracted attention as forage legumes. These species are D. sandwicense E. Mey, D. canum (Gmel), D. aparines (Link) D.C.* and D. uncinatum (Jacq) D.C.. D. aparines and D. uncinatum have been released as the commercial cultivars Greenleaf and Silverleaf desmodium, respectively, in Australia (Imrie, 1973) as they are much more productive and robust (stem length is 5 to 20

*Also called D. intortum (Mill) Urb.

meters) than either *D. canum* or *D. sandwicense*. *D. canum* has attracted attention in Hawaii in pasture swards (Younge, et al., 1964). *D. sandwicense* is the least robust of these four species with the stem length rarely exceeding two meters. McWhirter (1969) reported that F_1 heterosis for forage yield was not present in inter or intra-specific *Desmodium* crosses while Rotar and Chow (1971) indicated that the internode length of a two-species F_1 cross was intermediate to its two parents. These data suggest that *Desmodium* is not affected by inbreeding depression for forage yield.

Than Aung (1970) described *D. sandwicense* as a prostrate herb whereas Rotar and Chow (1971) observed an erect growth habit. The stem color has been reported by Rotar et al., (1967) to be either red-brown or green. The segregation of this character is 3 red-brown to 1 green (Hutton and Gray, 1967). Than Aung (1970) found the stems to be green with red pigmentation at the nodes. The stem was found to be round (Rotar, et al., 1967) or trisulcate (Than Aung, 1970) in cross section. The stems are pubescent with slightly hispid hairs (Than Aung, 1970) or glandular (Rotar et al., 1967). Rotar et al. (1967) found the internode length varied from 2.9 to 3.2 cm which was shorter than *D. uncinatum* or *D. aparines*. These authors believe that the internode length is controlled by multiple genes since the interspecific cross with *D. aparines* was intermediate to the parental phenotypes.

The leaves of *Desmodium* are alternate, entire and

trifoliate with the terminal leaflet being larger. McWhirter (1969) reported a dominant polyphyllous leaf gene. The leaf form of *D. sandwicense* is lanceolate with acute apices (Rotar et al., 1967) or ovate and cordate at the base (Than Aung, 1970). Rotar et al. (1967) found the length/width ratio of the sixth leaflet from the flower bud to range from 1.79 to 1.89 and the leaf area (length x width) to be from 8 to 12 sq cm. Than Aung (1970) found the leaf area to vary from 8 to 37 sq cm for all terminal leaflets. Hutton and Gray (1967) reported that the leaflets of *D. sandwicense* and *D. uncinatum* contain a silver-grey marking along the midrib. This character is dominant to the plain leaf and segregated as a single locus. The spotted leaf character common in *D. sparines* is occasionally found in *D. sandwicense* (McWhirter, pers. com.). Hutton and Gray (1967) reported that this character was dominant to the plain leaf and segregated with a 3:1 ratio in the F₂ generation. McWhirter (pers. com.), however, suggested that the spotted leaf character is a complementary dominant gene system. The spotted leaf was found to be closely linked with the stem color (Rotar et al., 1967).

The flowering habit of *D. sandwicense* is relatively indeterminant and differs from the *D. uncinatum* and *D. sparines* which flower in response to short days (Rotar, et al., 1967). The above authors also report that *D. sandwicense* will outcross at a rate of 18%. McWhirter (pers. com.) suggested that *Desmodium* is highly self pollinated in

the wild and estimates the outcrossing to be about 20%. He indicated that most collections give an impression of being rather highly inbred.

The flowers of Desmodium are borne on terminal or axillary racemes. The basal flowers of the raceme are first to open and flowering usually continues for eight days (Rotar et al., 1967) with about five flowers opening each day. Preceding the opening of the flowers, the raceme elongates rapidly separating the flowers. The petals elongate beyond the sepals the day before the flower opens. The anthers dehise at this time and the pollen is mature. The female parts of the flower mature the following morning when the standard becomes erect. The pistil is held between the keel florets and when tripped it halts midway to the standard without striking it. The pollen is discharged in a cloud about the stigma. Hutton (1960) reported that D. sandwicense trips spontaneously while Rotar et al. (1967) state that it will pollinate without tripping. McWhirter (pers. com.) reported that the pistil is 1 to 2 mm longer than the stamens and that pollination will not occur until tripped or disturbed. Following pollination, the flower wilts rapidly with the standard folding over the stigma.

The flower color in Desmodium is polymorphic varying from white to purple (Than Aung, 1970). The purple flower color is dominant to white and this character shows a complementary dominant gene inheritance in crosses of certain white flowered species (D. sandwicense and D.

aparines) (McWhirter, pers. com.).

The seed pods of Desmodium are distinctly articulated and contain an average of 6.3 seeds per pod (Rotar et al., 1967). The pods are covered with short hairs and stick readily to clothing or fur. The seeds are kidney-shaped and light brown in color. The 1000-kernel weight of D. sandwicense was 3.5 gm falling between D. uncinatum (4.0 gm) and D. aparines (1.8 gm) (Rotar and Urata, 1966).

The germination for machine harvested seed is 99% but germination is reduced when hand harvested. Hand harvested seeds may be scarified by chipping the testa (McWhirter, pers. com.) or immersion in concentrated sulfuric acid for 15 min (B. Groff, pers. com.). Following scarification, germination is normal.

McWhirter (1963) first reported successful crosses between D. sandwicense and D. aparines. Hutton and Gray (1967) reported successful crosses between D. sandwicense and D. uncinatum and D. uncinatum and D. aparines. They found no gross cytological barriers; however, incompatibilities were found by McWhirter (pers. com.) when specific lines were used. McWhirter (pers. com.) suggested that D. aparines be used as a bridging species.

Rotar and Chow (1971) found that pod formation was highest when D. sandwicense was the female parent but much reduced when it was the male parent. McWhirter (pers. com.) also found a reduced pod set when D. sandwicense was the male parent. Rotar and Chow (1971) found crosses between D.

uncinatum and D. aparines to be low for pod set and they concluded that physiological factors may be influential. Pollen abortion was highest in the D. sandwicense x D. aparines crosses and was lowest in the D. sandwicense x D. uncinatum crosses (Rotar and Chow, 1971). Rotar and Chow (1971) concluded that D. sandwicense was more closely related to D. uncinatum than to D. aparines. They reported that pod set was negatively correlated with percentage pollen abortion ($r = .74$, $df = 21$). McWhirter (pers. com.) reported that morphological differentiation was correlated with genetic divergence. He concluded that D. uncinatum was morphologically distinct from D. sandwicense and D. aparines.

Chow and Crowder (1972) report a cross between D. canum and D. uncinatum. They conclude that D. canum is genetically separate from D. sandwicense, D. uncinatum and D. aparines.

B. Genetic studies of symbiotic nitrogen fixation.

The study of symbiotic nitrogen fixation has been of considerable interest to agriculture for numerous years and this area has recently been comprehensively reviewed (Quispel, 1974; Hardy and Gibson, 1977). The limitation of genetical studies of the inheritance of symbiotic nitrogen fixation was of major concern (Dart, 1974). The majority of the genetical studies involved the study of major genes conditioning non-nodulating or ineffective nodulation interactions. Few genetical studies have been reported

giving results for plant variation for total nitrogen fixed.

Major genes conditioning strain-specific nodulation have been reported in alfalfa (Medicago sativa L.) (Burton and Wilson, 1939; Gibson, 1962), red clover (Trifolium pratense L.) (Nutman, 1954), subterranean clover (T. subterraneum L.) (Gibson, 1964), soybean (Glycine max L. Merril) (Williams and Lynch, 1954; Caldwell, 1966; Vest, 1970; Vest and Caldwell, 1972), and peas (Pisum sativum) (Holl, 1975; Lie et al., 1976). The major genes appeared to be of two types: (a) recessive genes with modifiers, or (b) dominant genes. The investigations in Trifolium suggested that strain-specific ineffective nodulation was controlled by a single recessive gene modified by recessive suppressor genes which restored effectiveness (Nutman, 1954; Gibson, 1964). Single dominant genes (Rj_2 , Rj_3 and Rj_4) were reported in soybeans conditioning strain-specific ineffective nodulation (Caldwell, 1966; Vest, 1970; Vest and Caldwell, 1972) while the recessive gene (rj_1)* conditioned a universal non-nodulation interaction in soybeans (Williams and Lynch, 1954).

Vest et al. (1973) classified the soybean plants homozygous for the RJ alleles (cultivars - Hardee, Hill) as nodulated-ineffective since the plants exhibited small, white cortical proliferations at the node sites. They concluded that infection had taken place. However, nitrogen

*Designated no by Williams and Lynch (1954).

was not being fixed. Vest *et al.* (1973) reported that RJ_2 and RJ_3 appeared to be linked yet both were independent of rj_1 . Although RJ_2 and RJ_3 were similar phenotypically, producing numerous small, white cortical proliferations, they appeared to differ in the serological properties of the ineffective strains. RJ_2 was ineffective with most strains of the cl and 122 serogroups while RJ_3 was ineffective with only one particular strain. Caldwell *et al.* (1966) performed reciprocal grafts of the cultivars Hardee (RJ_2 , RJ_2) and Lee (rj_2 , rj_2) and reported that the root genotype dictated the nodulation interaction.

The ineffective response for the RJ_4 gene (Hill x strain 61) appeared to be a specific response in the sense of the RJ_3 gene. The RJ_4 gene produced only a few small, white cortical proliferations since nodule abortion probably occurred at a very early stage of development (Vest and Caldwell, 1972).

A universal non-nodulation gene has also been reported for peas. Holl (1975) and Holl and LaRue (1975) studying crosses of an Afghanistan cultivar (non-nodulating) reported two genes affecting nodulation and fixation. Holl (1975) reported that the Sym_2 locus (nodulation) and the Sym_3 locus (fixation) interacted to give four phenotypes for fixation. Plants with the genotype Sym_2 --, Sym_3 -- were classed as nodulating and fixing while plants with the genotype sym_2 , sym_2 , -- -- were non-nodulating and non-fixing. However, when the Sym_2 gene was combined with the homozygous

recessive condition for the Sym_3 gene the resulting combination conditioned differing interactions. If the Sym_2 gene was heterozygous the resulting interaction was nodulation but no fixation, whereas, a homozygous dominant condition for the Sym_2 gene resulted in nodulation and low fixation. The Sym_2 locus conditioned nodulation and segregated 3 nodulating to 1 non-nodulating phenotypes.

Lie *et al.* (1976) reported a strain-specific, temperature-sensitive non-nodulation interaction in the pea cultivar, Iran. The authors indicated that nodulation occurred only when the root was subjected to favorable nodulation temperatures (26 C) for the second and third day following inoculation. Adverse temperatures during this period reduced nodulation whereas adverse temperatures before or after the critical period had no effect. Lie *et al.* (1976) reported that this strain-specific, temperature-sensitive ineffective nodulation interaction was conditioned by a dominant allele at a single locus.

The study of symbiotic nitrogen fixation has generally dealt with qualitative interactions relating to nodulation vs. non-nodulation. However, genetical studies have been reported for quantitative variation for estimates of nitrogen fixation. Burton and Wilson (1939) reported that varieties of alfalfa differed significantly for total nitrogen fixed when inoculated by several strains of *R. meliloti*. Conversely the authors also noted significant differences when strains were compared as well as specific

host-strain interactions. These data suggested quantitative differences among hosts and among strains. Burton and Wilson (1939) also noted that the results tended to be erratic and attributed these deviations to environmental factors. Later work in alfalfa by Gibson (1962) confirmed genetic variation of the host-strain interaction. He noted that the Canadian cultivar Rambler was inefficient when inoculated with Australian strains of *R. meliloti* yet was efficient when inoculated with Canadian strains. Gibson (1962) suggested that the *M. falcata* parentage of Rambler may be, in part, responsible for this inefficiency. Recently Seetin and Barnes (1977) reported genetic variation in alfalfa for quantitative nitrogen fixation estimated by the acetylene reduction assay and dry weight accumulation. These authors indicated that the alfalfa clones expressed F₁ heterosis and inbreeding effects for the estimates of nitrogen fixation. Duhigg et al. (1978) reported that F₁ progeny from *Mesilla* alfalfa plants selected for high acetylene reduction values expressed significantly greater acetylene reduction values than the parental population. Significant results were not reported for the low selection line.

In recent research with field beans (*Vicia faba*), El-Sherbeeny et al. (1977a,b) reported genetic variation among *Rhizobium leguminosarum* strains and among field bean lines, respectively. Mytton et al. (1977) indicated that the genetic variation for lines or strains was found to be minor in comparison to the line by strain interactions. Mytton et

al. (1977) partitioned the genetic variance into three components; namely, (a) the additive genetic effect of the host genotype, (b) the additive genetic effect of the strain genotype, and (c) the non-additive effect of the host by strain interactions. They indicated that 73% of the phenotypic variation could be directly attributed to host strain interactions and concluded that simultaneous selection for a specific host-strain interaction would be most effective.

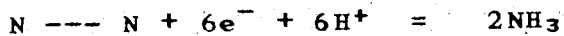
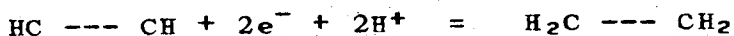
Mytton (1975), studying white clover (*Trifolium repens*) symbiosis, found that host variation for nitrogen fixation was larger than for differences among strains. He concluded that only effective strains had been collected thereby reducing genetic variation. However, Mytton (1975) did report significant host x strain interactions. Strains which were selected from one plant generally entered into a more productive interaction with relatives of that plant than with non-related plants suggesting selective hosts.

Minchin et al. (1978) also found genetic variation among cultivars of cowpea (*Vigna unguiculata*-U) but not among Rhizobium strains. The above authors, unlike Mytton et al. (1977), concluded that any breeding program should concentrate on the macrosymbiont.

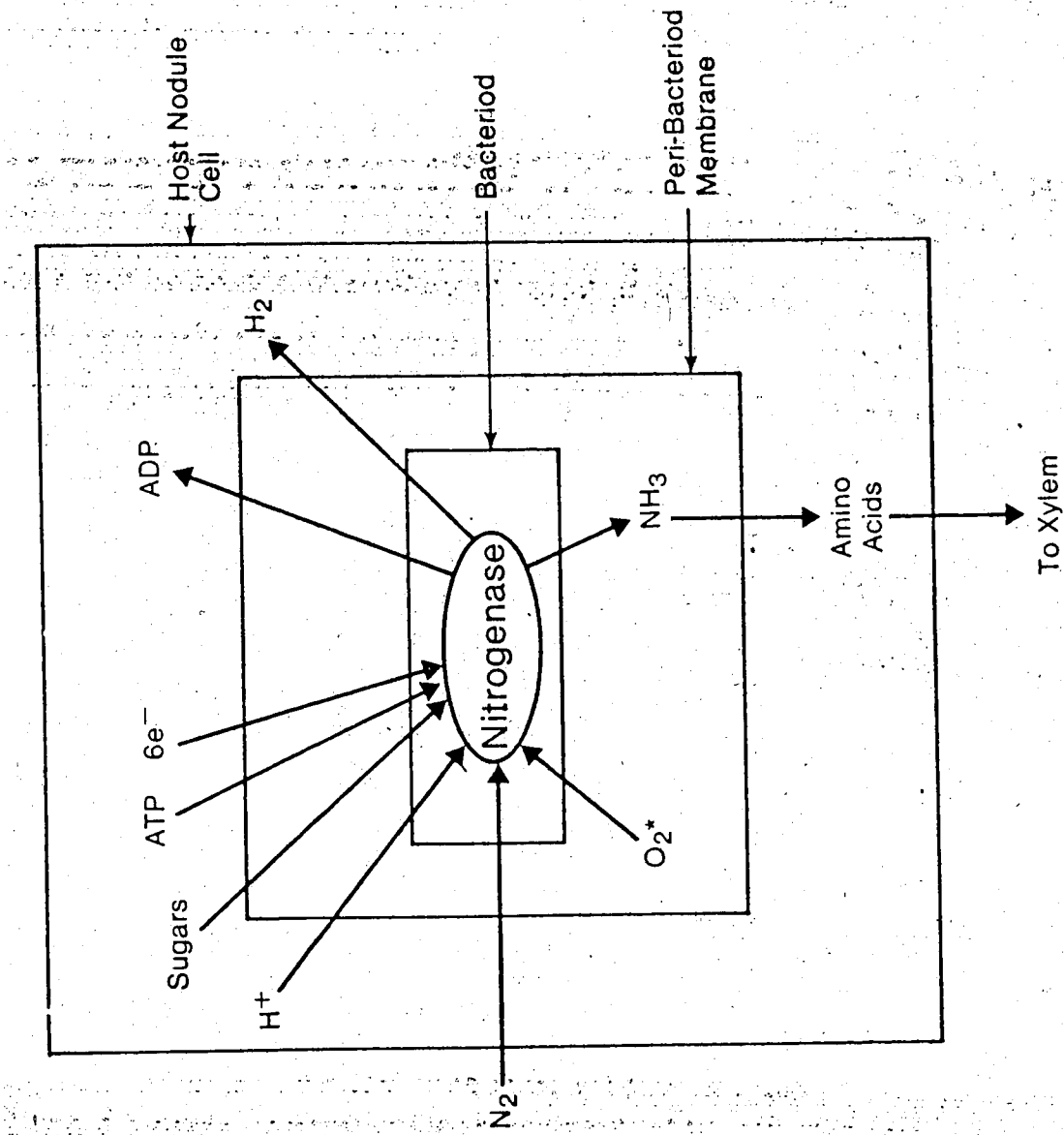
Quantitative assessments of the amounts of nitrogen fixed had been measured in earlier experiments by total dry weight accumulation of the shoots when plants were grown in nitrogen free containers (Gibson, 1962; Vincent, 1970). The

^{15}N method (Burris and Wilson, 1957) and, recently, the acetylene reduction assay (Hardy et al.; 1968) have been developed as assays of the activity of the enzyme nitrogenase (Figure 1). Of these two assays, the acetylene reduction assay has been most popular because it was simple, rapid and more sensitive than the ^{15}N method; but it was an indirect approximation (Burris, 1974). Since its introduction, the acetylene reduction assay has appeared in many publications measuring nitrogenase activity in intact nodules, nodule prepared cell-free extracts, field experiments, as well as marine and fresh-water habitats (Quispel, 1974).

The reduction of acetylene to ethylene by nitrogenase was independently discovered by Schollhorn and Burris (1966) (reviewed by Burris, 1974) and Dilworth (1966). Its usefulness as an index of nitrogenase activity was immediately realized by Hardy et al. (1968) who developed a method of measuring nitrogenase activity for field experiments. The relationship between acetylene and N_2 reduction may be represented as:



These formulae suggested that the conversion rate for N_2 to NH_3 was one-third that of C_2H_2 to C_2H_4 since acetylene reduction required two electrons while N_2 reduction required six. Burris (1974) reported that the conversion factor of 1/3 was not generally found but was much closer to 1/4 and



*Leghemoglobin Transported

Figure 1. A diagrammatic representation of the symbiotic nitrogen fixation pathway in the host nodule cell.

Burris suggested a comparison of the acetylene reduction assay with the ^{15}N method to determine an exact conversion factor when required for a particular experiment.

Recent studies with the acetylene reduction assay in soybeans suggested that nodule activity (C_2H_4) was determined by the physiological stage of the plant (Lawn et al., 1974). Lawn and Brun (1974) found that the nodule fresh weight and acetylene reduction activity for soybeans reached a maximum near the termination of flowering and declined rapidly during the onset of the pod-filling stage. The authors attributed this decline to a competition by the pods for available assimilate. Additional studies with reciprocal grafts in soybeans indicated there were significant differences among shoot and root genotypes as well as for sampling dates as measured by the acetylene reduction assay (Lawn and Brun, 1974). Significant shoot genotype x sampling date interactions were found for all characters studied while root genotype x sampling date interactions were not significant with the exception of the photosynthetic rate. There were significant shoot genotype x root genotype interactions for nodule number and nodule size. Nitrogenase activity (C_2H_4) for the shoot genotype means was significantly correlated with nodule fresh weight, root dry weight, shoot:root ratio, photosynthesis and shoot dry weight. However nitrogenase activity (C_2H_4) for root genotype means was not significantly correlated with any of the above characters. Lawn et al. (1974) concluded that root

genotypes may differ in their ability to efficiently utilize the photosynthate assimilated during symbiotic nitrogen fixation. They also suggested that the root genotype controlled nodule initiation and nodule size which were found to be significantly negatively correlated with the root genotype effect on nodule number.

Dart (1974) reported responses to selection in several Trifolium species for number of nodules per plant, time taken to form the first nodule and effectiveness of the nodules. These characteristics were also affected by the Rhizobium strain used. Dart (1974) concluded that plant breeding for increased nodulation was of significant potential.

C. Objectives.

It was the objective of this study of the inheritance of symbiotic nitrogen fixation in Desmodium to establish whether or not there exists genetic variation in the processes of the symbiotic relationship between host plant and Rhizobium with particular emphasis upon effective vs ineffective nodulation and the quantitative variation in total symbiotic nitrogen fixed per plant. It was also of interest to determine the genetic basis for such differences, if disclosed. The dry weight accumulation in a bottle jar (Leonard jar assembly) and the acetylene reduction assay were assessed to determine their potential as plant breeding tools.

MATERIALS AND METHODS

A. Outline.

In this study of the inheritance of symbiotic nitrogen fixation in Desmodium, experimentation was conducted at three levels. These levels, in increasing order of interest, have been broadly defined as methodology, qualitative genetic variation, and quantitative genetic variation (Table 1). The methodological studies encompassed the background

Table 1. Levels of study of the inheritance of symbiotic nitrogen fixation in the host

Comments:	
Level 1 Methodology	Preliminary studies How? What? Where? Improvements? When?
Level 2 - Qualitative Genetic Variation	
Host-strain interactions* effective ineffective	Which? (Major genes?)
Level 3 Quantitive genetic variation	
Fixation "efficiency"** efficient "inefficient	How much? (Minor genes?)
Ultimate goal	Why? Selection Super-fixing inbred lines or hybrids

*May also be modified by strain manipulation.

studies necessary for the development and improvement of the technique of studying symbiotic nitrogen fixation. Experiments in qualitative genetic variation were conducted to determine the genetic basis for ineffective/effective

nodulation interactions. The experiments involved the testing of the observed segregation ratios in F_2 and backcross populations for a goodness of fit to an expected segregation ratio.

The experiments involving quantitative genetic variation constituted the major interest in this study and relate to the investigation of the inheritance of total symbiotic nitrogen fixed per plant. The experimental designs which were chosen to establish and examine the genetic variation were: (1) a five-parent diallel cross, and (2) a cross between two inbred lines. The diallel cross consisted of inbred lines of *D. sandwicense* which were not selected for the characters under study. The cross between two inbred lines constituted crosses of *D. sandwicense* with two other species of *Desmodium* which were conspicuously different from *D. sandwicense* as well as a within *D. sandwicense* cross.

B. Materials.

1. *Rhizobium* strains.

To study one component of the host-strain interaction of symbiotic nitrogen fixation, it was necessary to hold the other component constant. Since the variation within the population created from crosses among *Desmodium* lines was of major interest, the *Rhizobium* component within each experiment has been held constant and was considered to be an environmental component. The two pure strains of

Rhizobium selected were strain A (CB627)* and strain B (CB1789)* which were isolated from D. parvipes and D. sandwicense, respectively. These strains differed in growth properties. Both strains are classed as slow growing; however, strain A is considered to be relatively faster growing than strain B. Both strains are members of the "Cowpea" group of Rhizobium (Vincent, pers. com.) and strain A has been released in Australia as a commercial inoculum for Desmodium.

2. Desmodium lines.

The Desmodium lines used in this study are described in Table 2. D. sandwicense or its crosses were used in most experiments because it: (a) is diploid, self-fertile and readily crossed, (b) is relatively insensitive to photoperiodism and is easily maintained under glasshouse conditions, (c) has a robust seedling (desirable for culture in a bottle jar assembly), and (d) can be hybridized easily with other conspicuously variable and commercially important species of Desmodium (Than Aung, 1970). Prolonged inbreeding does not adversely effect plant vigor and inbreeding depression is generally considered to be absent (McWhirter, 1969; Rotar and Chow, 1971). These characteristics made D. sandwicense an excellent model organism for nitrogen fixation studies.

*Commonwealth of Australia identification number.

Table 2. Description of the *Desmodium* lines involved in this study.

Study Identifi- cation	Accession or Line Ident.	Species	Source
Ds1	D. sandwicense E. Mey	CPI 18156	Dr. K.S. McWhirter, U. of Sydney
Ds2	D. sandwicense E. Mey	CPI 18163	Dr. K.S. McWhirter, U. of Sydney
Ds3*	D. sandwicense E. Mey	N993	Dr. K.S. McWhirter, U. of Sydney
Ds4	D. sandwicense E. Mey	CSL	Dr. J. Lotero C., I.C.A., Colombia
Ds5	D. sandwicense E. Mey	CWF	Dr. J. Lotero C., I.C.A., Colombia
Da1	D. aparines+(Link)D.C.	CPI 23189	Dr. K.S. McWhirter, U. of Sydney
Da2	D. aparines+(Link)D.C.	Q8382/56	Dr. K.S. McWhirter, U. of Sydney
Da3	D. aparines+(Link)D.C.	CPI 18010	Dr. K.S. McWhirter, U. of Sydney
Da4	D. aparines+(Link)D.C.	Desica-1	Dr. J. Lotero C., I.C.A., Colombia
Da5	D. aparines+(Link)D.C.	CPI 33814	Dr. B. Imrie, C.S.I.R.O., Brisbane
Dp	D. oroboides S.Watson	CPI 38722	Dr. K.S. McWhirter, U. of Sydney
Du	D. uncinatum (Jacq.) D.C.	CPI 8980	Dr. K.S. McWhirter, U. of Sydney
Dsp	?	Ecuador	Dr. B. Grof, C.I.A.T., Colombia
Dse	D. sericeophyllum (Schlecht)	CPI 33005	Dr. K.S. McWhirter, U. of Sydney

*(((Ds1 X Da1) X Ds1) X Ds1) - selfed 3 cycles until homozygosity
for the spotted leaf character.

+also called D. uniorum (Mill) Urb.

Five inbred lines of D. sandwicense (Ds1 to Ds5) were crossed, in all possible combinations including reciprocal crosses to create a diallel cross. Although the inbred lines were not selected for the characters studied, the lines have been considered as a fixed sample of D. sandwicense lines. The six populations (P_1 , P_2 , F_1 , F_2 , BC_1 , and BC_2) derived from a cross between two inbred lines were made for each of the following crosses: Ds1 x Ds1, Ds2 x Ds2, and Ds2 x Ds5. The diallel cross and the three crosses between two inbred lines constituted the basic material to determine whether genetic variation among differing populations existed for symbiotic nitrogen fixation in Desmodium.

C. Experimental Procedures.

1. Crossing technique.

Two crossing methods have been used in Desmodium, the emasculation method (Rotar et al., 1967) and the tripping method (McWhirter, pers. com.). The emasculation method is achieved by opening the flower bud the night before maturity and gently removing the anthers. The racemes with the emasculated flowers are then covered with a glassine bag containing a moist ball of cotton. During the morning hours, the flowers are pollinated with the desirable pollen using a sandpaper-tipped toothpick.

In the tripping method the mature flower is gently tripped and the stigma is examined with a 10X hand lens. Contaminated flowers are discarded. The remaining flowers

are pollinated and following pod formation glassine bags are placed over the raceme to avoid seed contamination.

Rotar *et al.* (1967) found the emasculation technique to be superior but more difficult than the tripping method. They reported that the emasculation-wet cotton method produced 60% crossed seed from a 10% flower set compared to the tripping method which yielded 40% crossed seed from a 13% flower set. The authors only brushed the stigma in the tripping method which resulted in the low percentage crossed seed. Rotar *et al.* (1967) and Hutton (1960) both noted that crossing was most successful when the temperature was cool and the relative humidity was high. Rotar and Chow (1971) report that percentage pod set and the monthly average temperatures were negatively correlated ($r = -.99$, $df = 3$).

The crossing method used in this study was a modification of the tripping method. Each flower was grasped by the thumb and forefinger at the calyx and tripped by applying pressure at the base of the standard. Following tripping, the anthers and pollen were brushed away. Beginning with the top flowers, the stigmas of the tripped flowers were thoroughly washed with a 2.5% (by weight) sugar solution and immediately pollinated with the desired pollen. The pollen was either taken directly from the pollinating flower which was prepared by holding the flower with the thumb and forefinger at the calyx and gently removing the petals without disturbing the pollen or by tripping the pollinating flower directly onto a fingertip. The finger was

dipped into alcohol between pollinations. Following pod formation the raceme was covered with a glassine bag to avoid seed contamination. Crosses were made at the Colombian Institute of Agriculture, Medellin, Colombia and at the University of Sydney, Sydney, Australia. Study of the progeny from some of these crosses showed that 95% of the F₁ crosses were hybrids for the spotted leaf character (Appendix A).

2. Bottle Jars.

The bottle jars (Leonard jar assembly), for nitrogen-free growth containers, were constructed from various inexpensive and readily available materials. The bottle jar which conformed to the description of Vincent (1970) is illustrated in Figure 2. During this study, the bottle jar was modified to facilitate the removal of the plant roots intact for the acetylene reduction analysis. The modified bottle jar is illustrated in Figure 3. After filling the spirits bottle with sand and compacting the sand about the wick, the sand was wetted and the reservoir was filled with a nutrient solution composed of the following:

CaHPO ₄	1.0 g
K ₂ HPO ₄	.2 g
MgSO ₄ · 7H ₂ O	.2 g
NaCl	.2 g
FeCl ₃	.1 g
Tap water	5.0 l

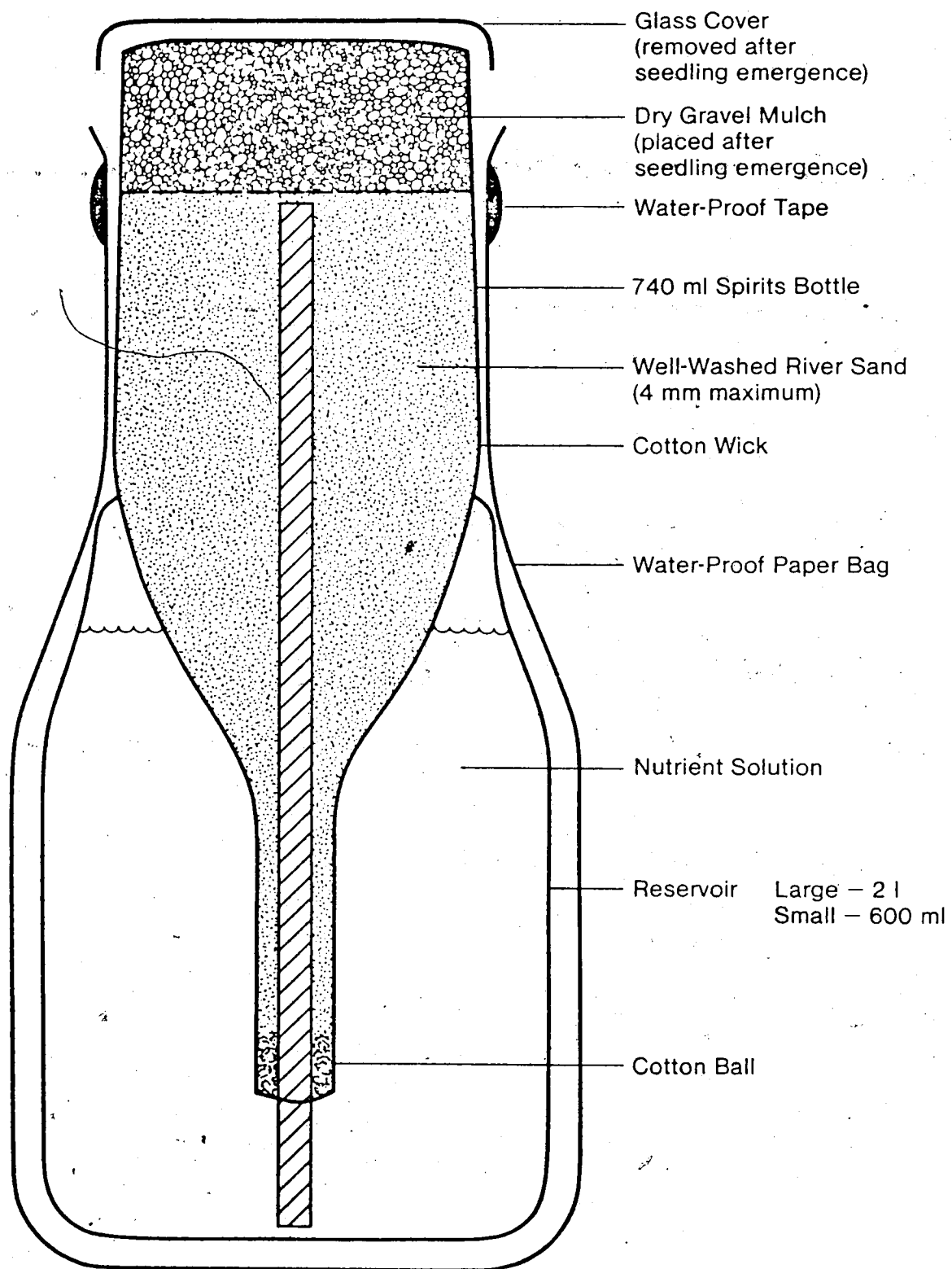


Figure 2. Initial bottle jar design (Vincent, 1970).

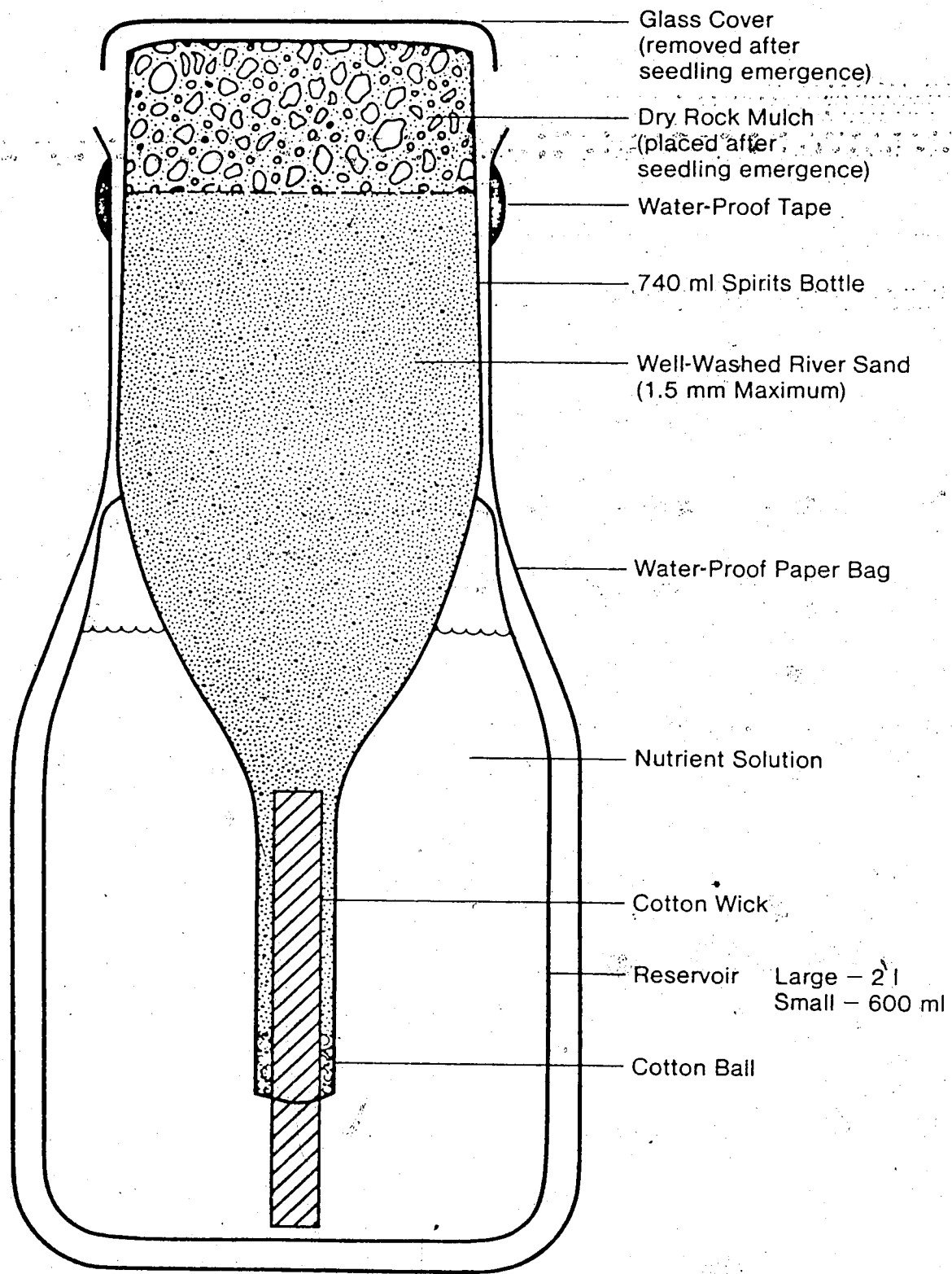


Figure 3. Modified bottle jar design.

The bottle jar was covered with a water-proof paper bag which was secured with water-proof tape, capped with half of a petri-dish, and autoclaved for 2 hours at 15 p.s.i... After cooling, the bottle jars were planted with surface sterilized seeds. The seeds were sterilized in an acidified 0.2% $HgCl_2$ solution (Vincent, 1970) and washed with six changes of sterile water. The testa of the seeds had been previously chipped to facilitate germination. Following germination and inoculation the bottle jars were covered with a pebble mulch (Vincent, 1970) and left undisturbed until harvest.

3. Rhizobium culture.

Stock cultures of the Rhizobium strains were maintained as agar slopes and stored at 2 C in McCartney bottles. Inoculum for the experiments was grown as broth cultures in 350 ml flats. The agar slopes and broth cultures were made from yeast-mannitol medium (Vincent, 1970) which consisted of the following ingredients:

K_2HPO_4	0.5 g
$MgSO_4 \cdot 7H_2O$	0.2 g
NaCl	0.1 g
$CaCO_3$	3 g
Mannitol	10 g
Yeast water	100 ml
Distilled water	900 ml
Agar (when required)	15 g

Fifty ml of the above medium was placed in each flat and autoclaved at 15 p.s.i. (120 C) for 15 min. The sterile

flats were stored at 2°C until required. Yeast water was freshly prepared by mixing 100 g of fresh compressed baker's yeast in one liter of cool water. The mixture was allowed to stand for 1 to 2 hours at room temperature before autoclaving for 50 min. The suspended material was allowed to settle and the clear supernatant was used as the yeast water.

Depending on the strain of Rhizobium, either ten to fourteen days prior to the inoculation time of the plants, a flat was removed from storage and inoculated with the desired strain using observed bacteriological precautions. The inoculated flat was placed in a growth cabinet at 26°C for the required time to facilitate the growth of the Rhizobium. For the inoculation of the experiments, a suspension of the Rhizobium was made by filling the flat with sterile water and shaking vigorously to disperse the Rhizobium. Five ml of this suspension was applied to the base of each plant 3 to 4 days following emergence, which was approximately 1 week after planting.

4. Quantitative variables.

(a) Acetylene reduction assay.

The acetylene reduction assay of Hardy et al. (1968) was used to assess the nitrogenase activity of the nodulated roots. Individual plants were harvested from the bottle jars and the root section from each plant was placed in a 13.5 ml vial which was plugged with an air tight seal. The vials

were immediately placed on a vacuum system, refilled with argon and injected with 2.8 cc of oxygen and 3.5 cc of acetylene (ethylene free). After 60 min, a 200 microliter sample was removed from each vial and injected into a F-11 Perkin-Elmer gas chromatograph with a hydrogen flame ionization detector. The detector was attached to a Tohshin electron chart recorder. Nitrogen was used as the carrier gas and the column was a stainless steel tube of 30 cm in length and .32 cm in diameter and filled with Porapack T packing material. The column temperature was held at 60°C giving a retention time of 15 sec for ethylene and 30 sec for acetylene. At periodic intervals, duplicate samples were processed to check the accuracy of sampling. An acetylene reduction estimate (nanomolar(nM)/pl/hr) was calculated from peak height using the following equation:

$$\text{Acetylene reduced} = \frac{E_s(E_f - E_o)}{A_s(A_f - A_o)} \times A_c \times S_t$$

where E_f and A_f were the final ethylene and acetylene peak heights, respectively, E_o and A_o were the origin values for ethylene and acetylene, and E_s and A_s were the scales of measurement for ethylene and acetylene. A_c was the acetylene constant which was determined as the mean of the acetylene peak heights. S_t was the correction for a standard known sample of ethylene.

Acetylene reduction assay conducted in Experiment VI

generally followed the procedure outlined above, except that a 35 ml vial was used with no argon. The ethylene peak was used as an estimate of acetylene reduced.

(b) Dry weight accumulation.

Dry weight accumulation data were collected for individual plants grown in bottle jar assemblies. The harvested plants were clipped at the cotyledonary node and the shoot portion was retained as the dry weight accumulation sample. The leaves dropped during the course of the experiment were collected and included in the sample. The samples were oven-dried at 105°C for 24 hours and left at room temperature for at least one week before weighing to the nearest milligram.

(c) Shoot length.

The shoot length was measured either *in situ* or on the harvested plants before oven drying with measurements being recorded to the nearest millimeter for the primary shoot length and the two secondary shoots originating at the cotyledonary node. Total shoot length was calculated by summing the primary and the two secondary shoot lengths.

D. Experimental designs and analyses.

1. Methodological studies.

(a) Selection for leaf size.

Prior to the commencement of the nitrogen fixation studies,

studies were conducted at the Colombian Agricultural Institute, Medellin, Colombia to become familiar with Desmodium growing in its natural habitat. A preliminary selection study was conducted with D. sandwicense for larger and smaller eighth node terminal leaflet area. Four hundred plants of a wild D. sandwicense population were selected as the base population. Several characters relating to the eighth node leaf were measured; namely, the 8th leaflet length and width, the petiole length, the internode length (between the 7th and 8th nodes), and the shoot length. From these characters the leaf area, the length/width ratio, the petiole/internode ratio, and the average internode length were computed.

Having defined the selection criterion (8th node terminal leaflet area) and the related characters, the selection was initiated by selecting five plants from the high and low 'tails' of selection criterion distribution and randomly mating the plants within each line. For the first cycle of selection, duplicate data for leaf size were recorded for the tenth leaf and were included to determine their response to a selection pressure applied to the eighth node terminal leaflet. One hundred plants were grown for each selection cycle and selecting five plants from the 'tails' of the selection criterion distribution. The data for all characters for cycles 1 and 2 were analysed with a one-way analysis of variance to determine whether the characters were responding to selection.

(b) Growth patterns.

To determine whether the bottle jar was an effective container for the study of nitrogen fixation in Desmodium, a study was initiated to observe the growth of Desmodium in a bottle jar. The fifteen diallel genotypes were planted, two plants of one genotype per bottle jar, along with five (inbred parents) uninoculated controls to determine a mean growth curve. The experiment was conducted during the winter season and the primary shoot length was recorded every twenty days for 100 days. From the mean growth curve, a sampling point for the nitrogen fixation studies would be determined.

(c) Linearity of acetylene reduction.

Hardy et al. (1968) reported that following injection of acetylene into a vial with nodulated soybean roots, acetylene was reduced to ethylene at a linear rate until approximately 90 minutes following injection. To verify that acetylene reduction by nodulated Desmodium roots occurred at a linear rate, four Desmodium species (Ds1, Da1, Dp, Du; Table 2) of plant breeding interest were inoculated with strain A and grown for 8 weeks. Two plants for each line were bulk harvested and prepared for the acetylene reduction assay. Samples, 50 microlitres each, for each vial were extracted and processed at 10 min. intervals.

The five diallel lines were also prepared for the acetylene reduction assay and a 50 microlitre sample was

measured every 10 min. to determine if fixation was linear.

(d) Bottle jar design.

A test of the bottle jar design was initiated by screening river sand into the following four sizes:

- (a) < 4.0 mm (equivalent to that of Vincent (1970))
- (b) 1.0 - 3.0 mm
- (c) 1.0 - 1.5 mm
- (d) < 1.0 mm

The sand sizes a to c were placed in the initial bottle jar design (Figure 2) and in the modified design (Figure 3) for size d. The sand in all bottle jars was compacted about the wick to ensure capillary movement of the nutrient solution.

The bottle jars were arranged into a randomized block experiment with six blocks, planted with the Ds1 line and inoculated with strain B. Three seeds were planted in each bottle jar and after the rock mulch was applied, each cell was thinned to the two most vigorous plants. The experiment was harvested after eight weeks and the data for dry weight and shoot length were recorded for each cell as: (a) the mean of the two plants, and (b) the difference of the two plants in the same bottle (within-cell difference). The cell mean data were analysed with a two-way analysis of variance to determine significant differences for dry weight accumulation among sand sizes.

2. Strain-specific ineffective nodulation interaction.

Strain-specific ineffective nodulation interactions between some lines of *D. aparines* and strain B were found in this study. To determine the mode of inheritance of these interactions, *D. sandwicense* (Ds1) X *D. aparines* (Da1 and Da2) cross plants were scored for green or yellow-chlorotic color in the F₂ and backcross (to Ds1) generations to determine the segregation ratios. The scored results were tested with the chi-square goodness of fit to an expected segregation ratio.

Two hundred and fifty F₂ seeds, 150 backcross seeds and 10 seeds each for parental controls were planted in bottle jars and inoculated with 5 ml of strain B to each seed. At eight weeks the plants were scored for green or yellow color and a random sample of seventy ineffectively nodulated F₂ plants were removed from the bottle jars and inspected for nodulation characteristics. Following inspection these plants were planted in sand and fertilized with a source of nitrogen to determine if nitrogen was, in fact, the limiting criterion.

Since some lines of *D. aparines* display the spotted leaf character, all the plants were also scored for this character and tested for a chi-square goodness of fit to an expected segregation ratio. Joint segregation tests were determined for ineffective/effective nodulation and spotted/plain leaves.

3. Experiment I.

The diallel cross was arranged in Experiment I as a randomized block design with four blocks (replicates) and fifteen bottle jars per block. The bottle jars were randomly assigned the fifteen diallel genotypes and were randomized on a 30 cm grid within each block. Similar bottle jars (original design, Figure 2) were used within each block with blocks 1 and 4 having similar bottle shapes while blocks 2 and 3 were different. Each bottle jar was planted with two surface sterilized seeds and each plant was inoculated with a 5 ml suspension of strain B following emergence. Blocks were planted at differing planting dates due to space limitations and to facilitate the acetylene reduction analysis. Experiment I was conducted during the winter season at the University of Sydney; therefore, the average glasshouse temperature was near 21 C.

Each block was harvested after a growth period of 10 weeks, and data relating to shoot length, dry weight accumulation and the acetylene reduction assay were recorded as the mean of the two plants in each bottle jar.

Although the inbred lines were not selected for the characters studied, the lines have been considered as a fixed sample of *D. sandwicense* lines. The data were analysed under the assumptions of the model I, method 2 (Griffing, 1956) and Analysis II (Gardner and Eberhart, 1966) diallel design to establish the combining ability effects (Griffing, 1956) and genetic constants (Gardner and Eberhart, 1966),

respectively.

4. Experiment II.

The Experiment II diallel cross was arranged as a randomized block design with four blocks (replicates) and fifteen bottle jars per block. The bottle jars were randomly assigned the fifteen diallel genotypes and were randomized on a 30 cm grid within each block. Similar bottle jars (original design, Figure 2) were used within each block with blocks 1 and 2 having one bottle shape, whereas blocks 3 and 4 had another. Each bottle jar was planted with two surface sterilized seeds and each plant was inoculated with a 5 ml suspension of strain A following emergence. Blocks were planted at different planting dates due to space limitations and to facilitate the acetylene reduction analysis. Experiment II was conducted during the spring season at the University of Sydney; therefore, the average glasshouse temperature was near 24 C.

Each block was harvested after a growth period of 10 weeks and data relating to dry weight accumulation and the acetylene reduction assay were recorded for each plant. The data were analysed under the assumptions of the model I, method 2 (Griffing, 1956) and Analysis II (Gardner and Eberhart, 1966) diallel design to establish the combining ability effects and genetic constants.

5. Experiment III.

Experiment III was grown as a factorial experiment with two strains (A and B), two blocks for each strain, and fifteen bottle jars per block with three plants for one genotype planted in each bottle jar. The fifteen diallel genotypes were randomly assigned to bottle jars which were randomized on a 30 cm grid within each block. Each block was randomly assigned a bench location and blocks inoculated with different strains were separated on the bench to avoid contamination. The total experiment was planted on one day at the University of Sydney during the summer season. The mean glasshouse temperature for this experiment was approximately 26 C.

Similar types of bottle jars (modified design, Figure 3) were autoclaved in two groups such that autoclaving has been confounded with blocks.

During the early juvenile leaf stage, Experiment III was invaded by insect larvae. Since leaf loss resulted, the experiment was sprayed with an insecticide and each plant was clipped back to the cotyledonary node. After 10 weeks, all plants were harvested and measurements for the acetylene reduction assay were taken for strain A on one day and for strain B the following day. One block for each strain was analysed during the late morning with the other block analysed during the early afternoon. Dry weight accumulation were also recorded for the shoot portion of each plant. Due to the death of several plants, the two plants with the

largest dry weight in each bottle jar were retained for diallel analysis. The data were analysed under the assumptions of the model I, method 2 (Griffing, 1956) and Analysis II (Gardner and Eberhart, 1966) diallel design to establish the combining ability effects and genetic constants.

6. Combined analysis.

The same diallel cross was used in the preceding diallel experiments; therefore, the data may be pooled for a comprehensive analysis. Data for individual plants were available for block 4 in Experiment I and all blocks in Experiments II and III giving nine blocks in total. Three of these blocks were inoculated with strain B while the remaining six were inoculated with strain A. Since strains have been considered as environmental effects for this study, strains have been pooled with blocks for the combined analysis. The data were analysed under the assumptions of the model I, method 2 (Griffing, 1956) and Analysis II (Gardner and Eberhart, 1966) diallel design to establish the combining ability effects and genetic constants.

7. Experiment IV.

A cross between the two inbred lines, Ds1 x Da1 (Table 2) was made creating the six populations: ($P_1(Ds1)$, $P_2(Da1)$, F_1 , F_2 , $BC_1 (P_1 \times F_1)$, and $BC_2 (P_2 \times F_1)$). Experiment IV was grown during the spring season at the University of Sydney

as a randomized block design with seven blocks and twelve bottle jars per block. All bottle jars were of the original design (Figure 2) and blocks 1 to 5 had 600 ml reservoirs (Figure 2) while blocks 6 and 7 had 2 l reservoirs. The six populations were allotted to the twelve bottle jars such that the P_1 , P_2 and F_1 populations received one bottle jar each, the F_2 population received five, and the backcross populations received two each. The bottle jars were randomized within each block and all blocks were planted on the same day with surface sterilized seeds. After emergence, each plant was inoculated with 5 ml of a suspension of strain A. The experiment was harvested after seven weeks growth with dry weight accumulation and the acetylene reduction assay being recorded for each plant. The analysis of variance of population means (Hayman, 1958) was conducted as a two-way analysis of variance with the population means within each block constituting the data which was analysed as outlined in Section E2. Although Hayman did not partition epistatic effects, a study was initiated to investigate the feasibility of such a partition (see Results and Discussion sec. A5).

8. Experiment V.

The six populations of the $Ds2 \times Dsp$ (Table 2) cross were grown in Experiment V as a randomized block design with five blocks and fifteen bottle jars per block. All bottle jars were of the modified design (Figure 3) and all had 600

ml reservoirs (Figure 3). The six populations were allotted to the fifteen bottle jars such that the P_1 , P_2 and F_1 populations received one bottle jar each, the F_2 population received six, and the backcross populations received three each. The bottle jars were randomized within each block and all blocks were planted on the same day. After emergence of each plant was inoculated with a 5 ml suspension of strain B. The experiment was harvested after ten weeks growth.

Dry weight accumulation and the acetylene reduction assay were recorded for each plant and the analysis of variance of population means (Hayman, 1958) was conducted as a two-way analysis of variance with the population means within each block constituting the data which were analysed as outlined in Section E2.

9. Experiment VI.

Experiment VI consisted of the six populations from the cross between the inbred lines Ds2 and Ds5 (Table 2) and was grown in a growth chamber at the University of Alberta. Three plants of one population were grown in a ten cm pot and inoculated at planting with five ml of a suspension of strain A. The pots were arranged into four blocks; two of which had sand of very fine (.15mm) particle size, one with fine (.25mm) sand, while the remaining block was a 50% mixture of each sand size. Within each block the populations were allotted such that 5 pots each were allotted to the P_1 (Ds2), P_2 (Ds5), and F_1 populations while 20 pots were

allotted to the the F₂ population and 10 pots each to the backcross populations. All pots were wetted with the nutrient solution prior to planting. The experiment was maintained at a temperature of 23 C and harvested after 11 weeks. Data recorded for each plant were the acetylene reduction assay, shoot fresh and dry weight, root fresh weight, the cotyledon length and width (first node), and the width of the juvenile leaf (second node) since the juvenile leaf is nearly circular. The data were analysed on an individual plant basis as outlined in Section E2.

E. Genetical analyses.

1. Diallel cross.

A diallel cross is a set of all possible crosses among a set of inbred lines and the diallel table is a square matrix of dimension equal to the number of parents. The main diagonal represents the inbred lines while the data above and below the main diagonal represents the F₁ and reciprocal F₁ crosses, respectively. In the absence of reciprocal effects the matrix is symmetrical. A diallel table may be formed by four methods from the components of the square matrix (Griffing, 1956).

The diallel analysis of variance of the data in the diallel table has been reviewed and discussed by several workers (Sprague and Tatum, 1942; Hayman, 1954; Griffing, 1956; Cockerham, 1963; Gardner and Eberhart, 1966) and the relationship among the four methods and the analyses for

each method reviewed by Baker, 1978.

The interpretation of the diallel analysis of variance depends upon the model under investigation. Griffing (1956) reported four differing models underlying the analysis of variance. The models are:

Model I. Fixed treatment and block effects

Model II. Random treatment and block effects

Model A. Random treatment and fixed block effects

Model B. Fixed treatment and random block effects.

The basic difference between these models relates to the difference in the interpretation of the fixed and random treatment (genotype) effects. The fixed model assumes that the inbred lines and their F_1 crosses constitute the total population about which inferences are to be made and estimates combining ability effects and variances for the inbred lines. The random model assumes that the inbred lines are a random sample from an ancestral population about which inferences are to be made. The random model estimates the genetic and non-genetic components of the population variance.

Hayman (1954) presented several assumptions underlying a diallel analysis which must be satisfied before the conclusions may be considered reliable. These assumptions are:

- (a) that the parents are homozygous,
- (b) there is diploid segregation,
- (c) there are no reciprocal effects,

- (d) there is no multiple allelism,
- (e) the genes are independently distributed, and
- (f) there is independent action of non-allelic genes
(epistatic effects).

Griffing (1956) reported that the independent action of the non-allelic genes was the assumption of major importance. The diallel cross among inbred lines of *D. sandwicense* used in this study satisfied assumptions (a) and (b) while assumptions (c) and (d) were considered to be of minor importance. Assumptions (e) and (f) concerning linkage and epistatic effects, respectively, were of major concern in the present investigation since *Desmodium* has never been tested for such effects.

The analysis of the acetylene reduction assay, the dry weight accumulation characters and other characters were based on a Model I, method 2 diallel analysis as outlined by Griffing (1956) and modified to include Analysis II of Gardner and Eberhart (1966). The Model I, method 2 diallel analysis used in this study implies a statistical rather than a genetic interpretation (Baker, 1978) to partition the genetic component sum of squares and to estimate genetic constants (Gardner and Eberhart, 1966) and combining ability effects (Griffing, 1956). A listing of the Fortran computer program employed in this study for the diallel analysis of the data is presented in Appendix B.

2. Cross between two inbred lines.

An analysis of a cross between two inbred lines as outlined by Mather and Jinks (1971) may consist of six or more populations. The six populations of the minimum design are: parent 1 (P_1), parent 2 (P_2), the initial cross (F_1), the segregating generation (F_2), and the backcrosses of the F_1 to both parents (B_1 and B_2). For this fixed model, estimates of the type of gene action may be obtained from first degree statistics (Mather and Jinks, 1971).

Estimates of the magnitude and significance of the genetic parameters were calculated from the population means with the formulae as presented by Mather and Jinks (1971). The equations for the calculation of the genetic parameters are:

$$m = .5\bar{P}_1 + .5\bar{P}_2 + 4\bar{F}_2 - 2\bar{B}_1 - 2\bar{B}_2$$

$$a = .5\bar{P}_1 - .5\bar{P}_2 + \bar{B}_1 - \bar{B}_2$$

$$d = 1.5\bar{P}_1 - 1.5\bar{P}_2 - \bar{F}_1 - 8\bar{F}_2 + 6\bar{B}_1 + 6\bar{B}_2$$

$$aa = -4\bar{F}_2 + 2\bar{B}_1 + 2\bar{B}_2$$

$$ad = -\bar{P}_1 + \bar{P}_2 + 2\bar{B}_1 - 2\bar{B}_2$$

$$dd = \bar{P}_1 + \bar{P}_2 + 2\bar{F}_1 + 4\bar{F}_2 - 4\bar{B}_1 - 4\bar{B}_2$$

where

m = F_2 mean

a = additive effects

d = dominance effects

aa = additive X additive interaction

ad = additive X dominance interaction

dd = dominance X dominance interaction.

These equations allow for an estimate of the magnitude of the parameters to be made with the significance of each being tested by the t-test of the estimate to its standard deviation (Mather and Jinks, 1971). The variance of each parameter is determined from the variances of the six populations. For example, an estimate of the variance of d is obtained from the following equation:

$$Vd = 2.25VP_1 + 2.25VP_2 + VF_1 + 64VF_2 + 36VB_1 + 36VB_2$$

where Vd = the variance of the dominance deviations, VP_1 = the variance of P_1 , VP_2 = the variance of P_2 , etc.

An alternate analysis of population means is the analysis of variance of the population means (Hayman, 1958; Voigt *et al.*, 1966). To estimate the genetic parameters of the model a coded matrix was constructed (Mather and Jinks, 1971). The coded matrix is as follows:

Mean	a	d	aa	ad	dd
\bar{P}_1	1	0	1	0	0
\bar{P}_2	-1	0	1	0	0
\bar{F}_1	0	1	0	0	1
\bar{F}_2	0	1/2	0	0	1/4
\bar{B}_1	1/2	1/2	1/4	1/4	1/4
\bar{B}_2	-1/2	1/2	1/4	-1/4	1/4

The matrix may be weighted by dividing each row by the standard error of the population mean. The coded matrix was subjected to a stepwise least squares analysis (Draper and Smith, 1966) which has been restricted in this investigation. The genetic parameters of the model were

estimated on the assumption that within locus effects are of more importance than between loci interactions in a genetic model; i.e., a simple model. Therefore the variance among the population means was first analysed for additive and dominance effects before estimating the interaction parameters (see Results and Discussion, Section A5). The significance of the parameter mean squares was tested by the F-ratio with the error term from the analysis of variance of population means as the denominator (Voigt et al., 1966).

Properly coding the genotypes, replications, etc. prior to date entry allows the usage of the SPSS statistical analysis package (Nie, et al., 1975) for analysing the data for a cross between two inbred lines. SPSS[®] is particularly suited to the recording of the data or the creation of new variables to assist in the complete analysis and interpretation of the data. The partitioning of the population sum of squares into its genetic components is not available in SPSS, however, an interactive Fortran computer program, listed in Appendix C, was developed for this analysis. Program HAYMAN follows the analytical procedure for the analysis of variance of population means as defined by Hayman (1958) and earlier discussed. HAYMAN is suited for genetic analysis of any type of statistical design whether a one-way or a multi-factorial analysis of variance or covariance since the population sum of squares is partitioned into genetic components by the correlation form of the stepwise regression analysis. This regression

analysis is independent of scale or design (Draper and Smith, 1966).

(3) Principal component transformations.

Total nitrogen fixation has been estimated in this study by many measures and correlations among the measures would be expected. When variables are highly correlated the communal variation within the correlation matrix can be summarized with a principal components transformation (Winer, 1971) to construct "variables" which estimate the variation along the axes of principal variation. This transformation is also called principal components analysis (Overall and Klett, 1972) where the primary emphasis is to define "composite or factor-score variates that have certain desirable statistical properties". Principal components analysis may be divided into three stages of analysis. Firstly the analysis begins with the calculation of a correlation or a covariance matrix. Then the correlation or covariance matrix is mathematically reduced to eigenvectors (factor loadings) and eigenvalues (the amount of variance accounted by each eigenvector). Generally a correlation matrix with unity (total variances) along the main diagonal is structured because this matrix is scale independent (Overall and Klett, 1972). Finally, factor score coefficients are calculated from the factor loading matrix with a straightforward mathematical procedure analogous to a regression of an unknown dependent variable on the

original variables. A composite variable is a linear combination of factor score coefficients of the variables. The first composite variable accounts for greatest amount of common variation in the data. The second composite variable is orthogonal to the first and accounts for the next largest portion of the common residual variation. The remaining composite variable accounts for the remaining variation. Principal components analysis, being primarily concerned with construction of composite variable scores, normalizes the eigenvectors by converting the eigenvector matrix to one of correlations among the original variables and the derived factors. Such a normalization is of interest in determining what the principal axes represent. From the normalized eigenvectors, composite variables are defined which maximize existing differences among individual cases.

Principal components analysis has been used in agronomy by Denis and Adams (1978) to relate plant variables to yield of *Phaseolus vulgaris* L. (common bean) and by Hussaini, et al. (1977) to define major groupings within the world collection of *Eleusine coracana* L. (finger millet). However, principal components analysis has not been employed as a method of analysing differences among the population means. Since the principal axis of variation is expected to be nitrogen fixation, it would be pertinent to define this axis and investigate its variation.

The principal axis of variation in this study was scaled to a distribution with a mean and standard deviation

of 1/3 that of the acetylene reduction assay. This scale was used because nitrogenase theoretically reduces three times the amount of acetylene than N₂ (Burris, 1974). Following scaling, the composite N₂ fixation estimate was treated as another character for the analyses. The interpretation of the principal axis of variation will be restricted to the investigation of the differences among population means rather than attempting to estimate the amount of nitrogen fixed.

RESULTS AND DISCUSSION

A. Methodology

To achieve the ultimate goal of lines or hybrids with greater total nitrogen fixation it is necessary to attain an understanding of the genetic mechanisms operating in the host. In this study, the area of genetic mechanisms of nitrogen fixation has been divided into three levels (Table 1). In this generalized section of methodology several studies were undertaken both initially and during experimentation to establish or to investigate various aspects of methodology.

1. Selection for leaf size.

A preliminary study was initiated in Desmodium to become familiar with this species growing in its natural habitat. This study involved applying a selection pressure to the eighth node terminal leaflet area to determine its response to selection. The frequency distribution of the eighth node terminal leaflet sizes for the base population are presented in Figure 4. The distribution indicated that considerable variation was present in the wild population. Principal components analysis of the base population characters suggested that most characters were related to one factor i.e. a leaf size factor (Factor 1, Table 3) while the ratio characters are individually related to other factors.

Data collected for the tenth node leaf for a Dsp (Table

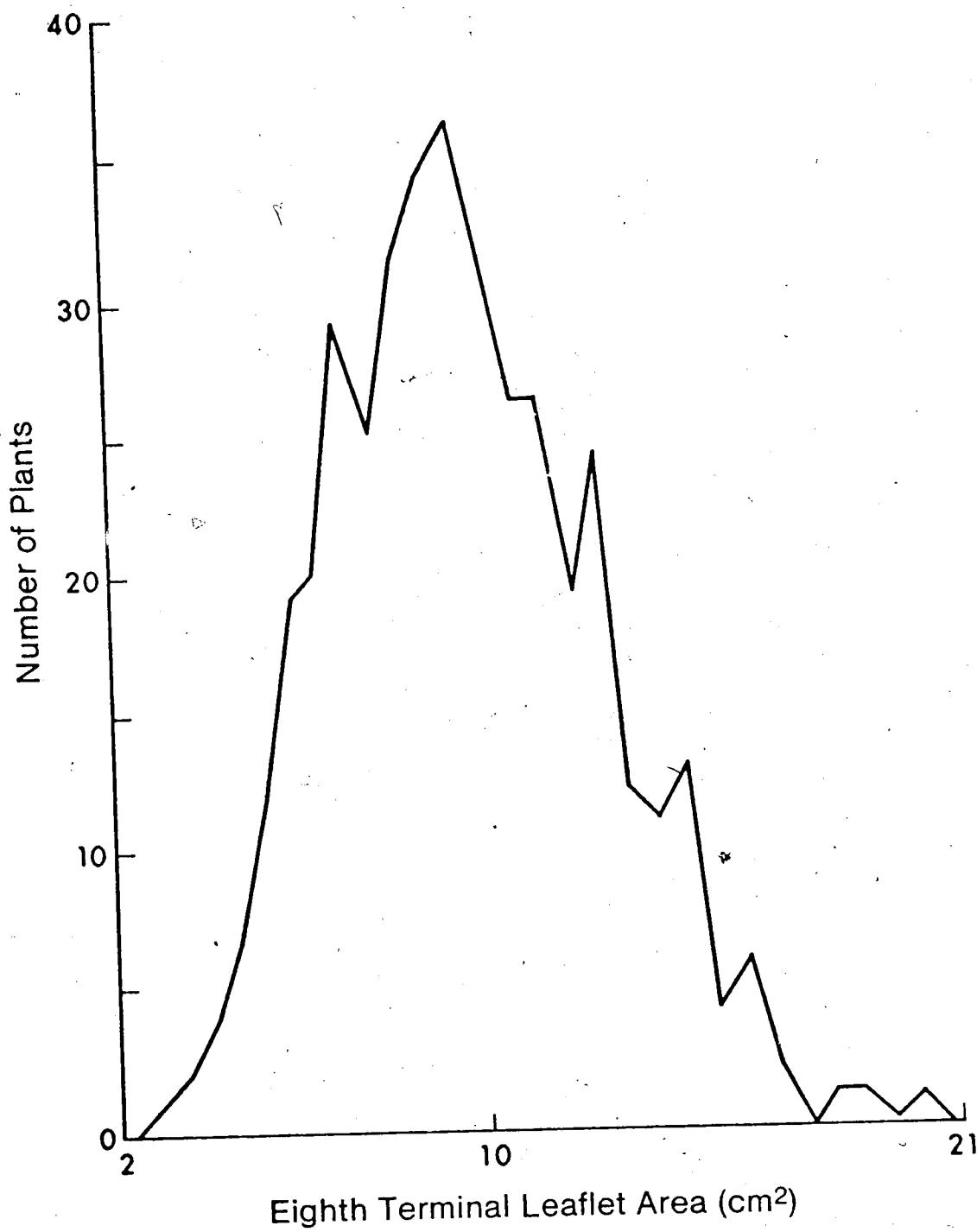


Figure 4. Frequency distribution of the eighth node terminal leaflet area for the base population.

Table 3. Eigenvectors for factors from the base population.

Character	Factor		
	1	2	3
Terminal leaflet length(8*)	0.94	0.07	0.01
Terminal leaflet width(8)	0.91	0.05	-0.34
Terminal leaflet area(8)	0.94	0.06	-0.17
Length/width ratio(8)	-0.04	0.02	0.95
Petiole length(8)	0.85	0.28	0.11
Internode length(7-8)	0.74	-0.54	0.11
Petiole/internode(8)	0.10	0.96	0.04
Shoot length	0.81	-0.33	0.25
Average internode length	0.81	-0.41	0.25

*node number.

2) population indicated that leaf measurements were related to one factor while plant weights related to another (Table 4). Ratio characters again were related each to a single factor.

Principal components analysis of the cycle 1 data still suggested leaf size characters were related to the first factor while ratio characters were related to another (Table 5). The means and standard deviations for each character for cycle 1 are presented in Table 6. The base population was found to have a significantly larger eighth terminal leaflet area than either the high or low selection lines with the exception of the ratio characters where the reverse was true. Means and standard deviations for the factor score variables indicated the same patterns as the selection criterion (Table 7).

Cycle 2 data also indicated the same pattern between the characters and the factors (Table 8). The response to selection by the high line was found to be negative for cycle 1; however, the high line mean was found to be significantly larger than the base mean suggesting a response to selection (Table 9). The low line mean was found to be significantly lower than the base mean while the factor score means also indicated that the response to selection away from the base population was significant (Table 10). The analyses suggested that the leaf size characters were highly related as indicated by their factor loadings to factor 1 and similarly with which they

Table 4. Eigenvectors for factors from the Dsp population.

Character	Factor			
	1	2	3	4
Terminal leaflet length(10*)	-.04	0.94	-.07	0.28
Terminal leaflet length(10)	-.06	0.97	-.07	-.12
Terminal leaflet area(10)	-.06	0.97	-.10	0.10
Length/width ratio(10)	0.05	0.18	0.00	0.96
Petiole length (10)	0.14	0.88	0.15	0.11
Internode length (9-10)	-.06	0.73	-.57	0.07
Petiole/internode ratio(10)	0.19	-.06	0.93	0.01
Shoot length	0.91	0.03	-.21	0.17
SQRT(no. of nodes)	0.76	-.36	0.16	-.10
Average internode	0.73	0.26	-.32	0.28
SQRT(no. of tillers)	0.76	-.12	0.40	0.00
Shoot weight	0.95	0.01	0.17	-.03
Leaf weight	0.88	-.07	0.19	0.01
Total weight	0.93	-.04	0.19	0.00
Leaf/shoot ratio	-.72	-.14	0.01	0.11

*node number.

SQRT = square root.

Table 5. Eigenvectors for factors from cycle 1.

Character	Factor			
	1	2	3	4
Terminal leaflet length(8*)	0.88	-.27	-.01	0.22
Terminal leaflet width(8)	0.89	-.26	-.04	-.20
Terminal leaflet area(8)	0.91	-.28	-.04	0.02
Length/width ratio(8)	-.04	-.03	0.07	0.84
Petiole length (8)	0.74	0.42	0.18	0.11
Internode length (7-8)	0.36	-.76	0.28	0.10
Petiole/internode ratio(8)	0.14	0.92	-.15	-.03
Terminal leaflet length(10)	0.91	0.17	0.12	0.17
Terminal leaflet width(10)	0.88	0.20	0.05	-.31
Terminal leaflet area(10)	0.93	0.20	0.08	-.07
Length/width ratio(10)	0.01	-.05	0.10	0.85
Petiole length(10)	0.56	0.33	-.39	-.01
Internode length(9-10)	0.46	0.02	0.75	0.09
Petiole/internode ratio(10)	0.10	0.27	-.93	-.07

*node number.

Table 6. Means and standard deviations from a one-way analysis of variance of the selection variables for cycle 1.

		Eighth node					
		Terminal leaflet					
	n	Length (cm)	Width (cm)	Area* (cm ²)	Petiole/ width ratio	Length/ width ratio (cm)	Petiole/ length ratio
High	100	5.0±.39 ^{a†}	3.8±.29b	19.2±2.8b	1.3±.06a	6.6±.81a	5.7±0.74b 1.2±.20a
Base	100	5.0±.47 ^a	4.0±.33a	20.2±3.3a	1.1±.07b	6.4±.74b	6.4±1.39a 1.0±.25b
Low	100	4.4±.32b	3.5±.29c	15.4±2.3c	1.3±.06a	5.8±.53c	5.6±0.85b 1.1±.17b

*Selection criterion.

†means followed by the same letter are not significantly different at the 5% level for the new multiple range test.

Table 6 (cont.). Means and standard deviations from a one-way analysis of variance of the selection variables for cycle 1.

Selection line	n	Tenth node			
		Length (cm)	Width (cm)	Area (cm ²)	Petiole/width ratio
Terminal leaflet					
High	100	6.0±.49b†	4.5±.38a	27.0±4.3a	1.3±.07b
Base	100	6.1±.50a	4.4±.34a	27.3±3.9a	1.4±.08a
Low	100	5.4±.38c	4.0±.38b	21.5±3.3b	1.3±.08b
					5.6±0.72b
					7.6±1.07b
					0.74±.11a

*means followed by the same letter are not significantly different at the 5% level for the new multiple range test.

Table 7. Means and standard deviations from a one-way analysis of variance of the factors for cycle 1.

Selection line	n	Factor			
		1	2	3	4
High	100	0.28±.88b*	0.36±0.88a	0.34±1.14a	-0.09±0.92a
Base	100	0.53±.81a	-0.36±1.19c	-0.02±1.02b	0.12±1.10a
Low	100	-0.82±.71c	-0.02±0.75b	-0.33±0.69c	-0.03±0.96a

*means followed by the same letter are not significantly different at the 5% level for the new multiple range test.

Table 8. Eigenvectors for factors from cycle 2.

Character	Factor		
	1	2	3
Terminal leaflet length(8*)	0.95	-.12	0.05*
Terminal leaflet length(8)	0.92	-.09	-.24
Terminal leaflet area(8)	0.94	-.12	-.10
Length/width ratio(8)	-.17	-.03	0.84
Petiole length (8)	0.91	-.06	-.11
Internode length (7-8)	0.56	-.75	-.11
Petiole/internode ratio(8)	0.16	0.89	0.04
Terminal leaflet length(10)	0.96	0.02	0.06
Terminal leaflet width(10)	0.93	0.01	-.30
Terminal leaflet area(10)	0.96	0.00	-.13
Length/width ratio(10)	-.14	0.05	0.86
Petiole length (10)	0.84	0.21	-.20
Internode length (9-10)	0.64	-.63	-.10
Petiole/internode ratio(10)	0.11	0.90	-.11

*node number.

Table 9. Means and standard deviations from a one-way analysis of variance of the selection variables for cycle 2.

Selection line	n	Terminal leaflet		Eighth node	
		Length (cm)	Width (cm)	Area* (cm ²)	Petiole length/width ratio
High	95	4.5±.66 ^{a†}	3.4±.53 ^a	15.6±4.4 ^a	1.3±.07 ^b
Base	99	3.9±.64 ^b	3.0±.49 ^b	12.2±3.7 ^b	1.3±.08 ^b
Low	95	3.4±.46 ^c	2.5±.38 ^c	8.6±2.2 ^c	1.4±.10 ^a
					4.6±0.77 ^c
					3.6±0.71 ^c
					1.3±.22 ^a

*selection criterion.

†means followed by the same letter are not significantly different at the 5% level for the new multiple range test.

Table 9 (cont'd). Means and standard deviations from a one-way analysis of variance of the selection variables for cycle 2.

Selection line	tenth node		terminal leaflet		petiole length/width ratio	internode petiole/internode length ratio
	n	length (cm)	width (cm)	area (cm^2)		
High	95	4.8±.82a†	3.9±.65a	20.8±6.2a	1.4±.09b	6.1±1.31a 8.4±1.44a 7.3±.17b
Base	99	4.8±.66a	3.5±.48b	17.1±4.3b	1.4±.11b	5.2±0.89b 6.6±1.56b 8.3±.22a
Low	95	4.2±.54b	2.9±.36c	12.0±2.8c	1.5±.13a	4.5±0.81c 6.0±1.20c 7.7±.15b

means followed by the same letter are not significantly different at the 5% level for the new multiple range test.

Table 10. Means and standard deviations from a one-way analysis of variance of the factors for cycle 2.

Selection line n		Factor		
		1	2	3
High	95	0.74±.98a*	-0.47±0.97b	-0.20±0.80b
Base	99	0.00±.79b	0.28±1.12a	-0.23±0.97b
Low	95	-0.74±.59c	0.17±0.70e	0.44±1.06a

*means followed by the same letter are not significantly different at the 5% level for the new multiple range test.

responded to a selection pressure upon one of them. The selection response for the eighth node terminal leaflet and the first principal factor over two cycles has been summarized in Figure 5.

2. Growth patterns.

An initial study was conducted with the crosses among D. sandwicense lines inoculated with strain B to observe the growth of Desmodium in a bottle jar and to observe the general growth pattern over a period of time. The mean growth patterns of the inoculated and uninoculated plants are presented in Figure 6. During the winter season, the transition from lag to log growth phase occurred at approximately the 10th week for the inoculated plants. Since this transition occurred later in the winter season, the 10th week was chosen as an appropriate sampling time for a study of the relationship between dry weight accumulation and the acetylene reduction assay. The uninoculated controls were yellow and chlorotic and their height did not increase beyond the height at juvenile stage. This chlorotic condition suggested that the bottle jar was an effective assembly for the study of symbiotic nitrogen fixation in Desmodium.

Diallel tables for several characters measured on the inoculated plants of the above study were constructed and partitioned into genetic component mean squares which are expressed as a percentage of the genotype mean square and

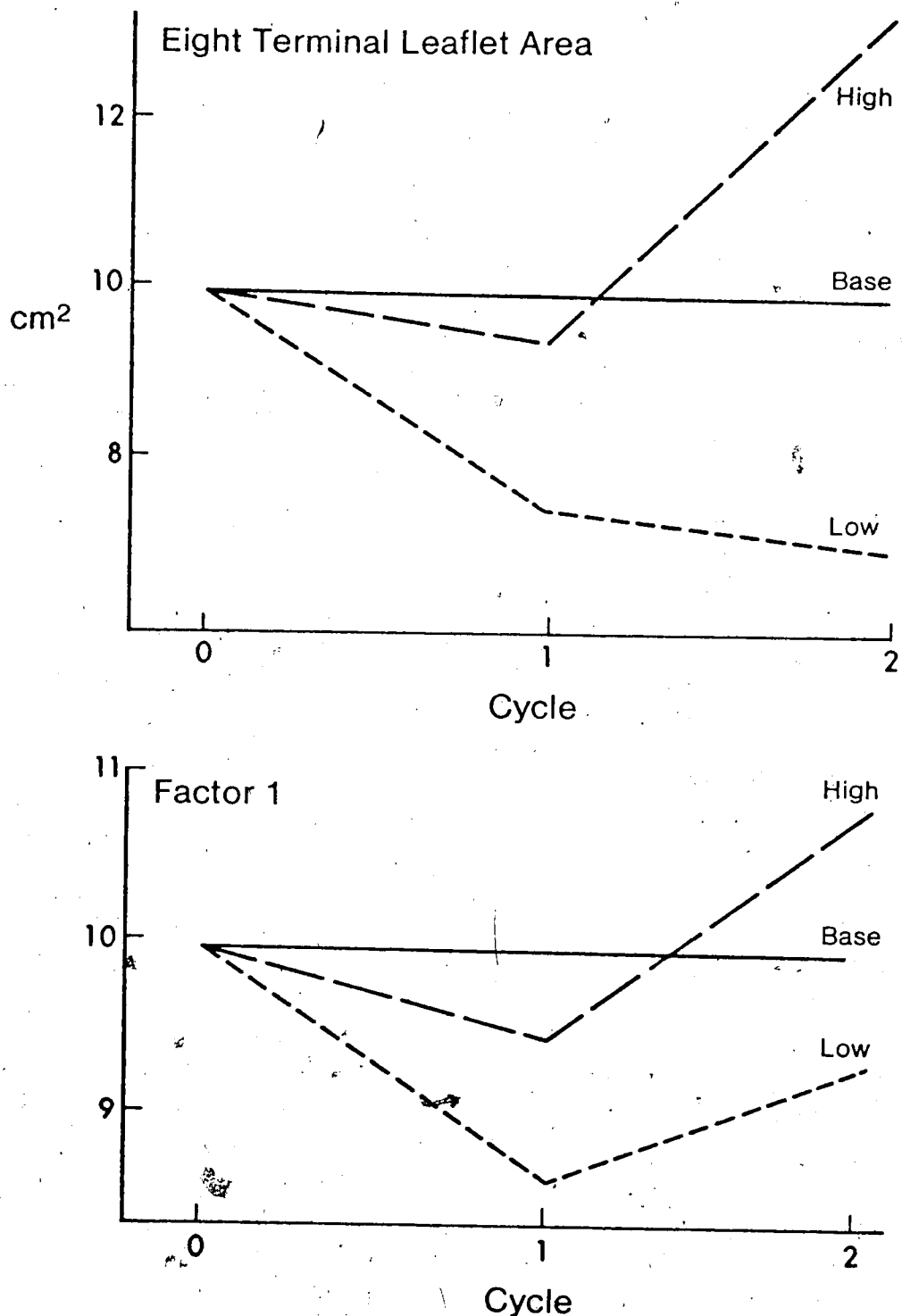


Figure 5. Selection response of the high and low lines for the selection variable and the first principal factor.

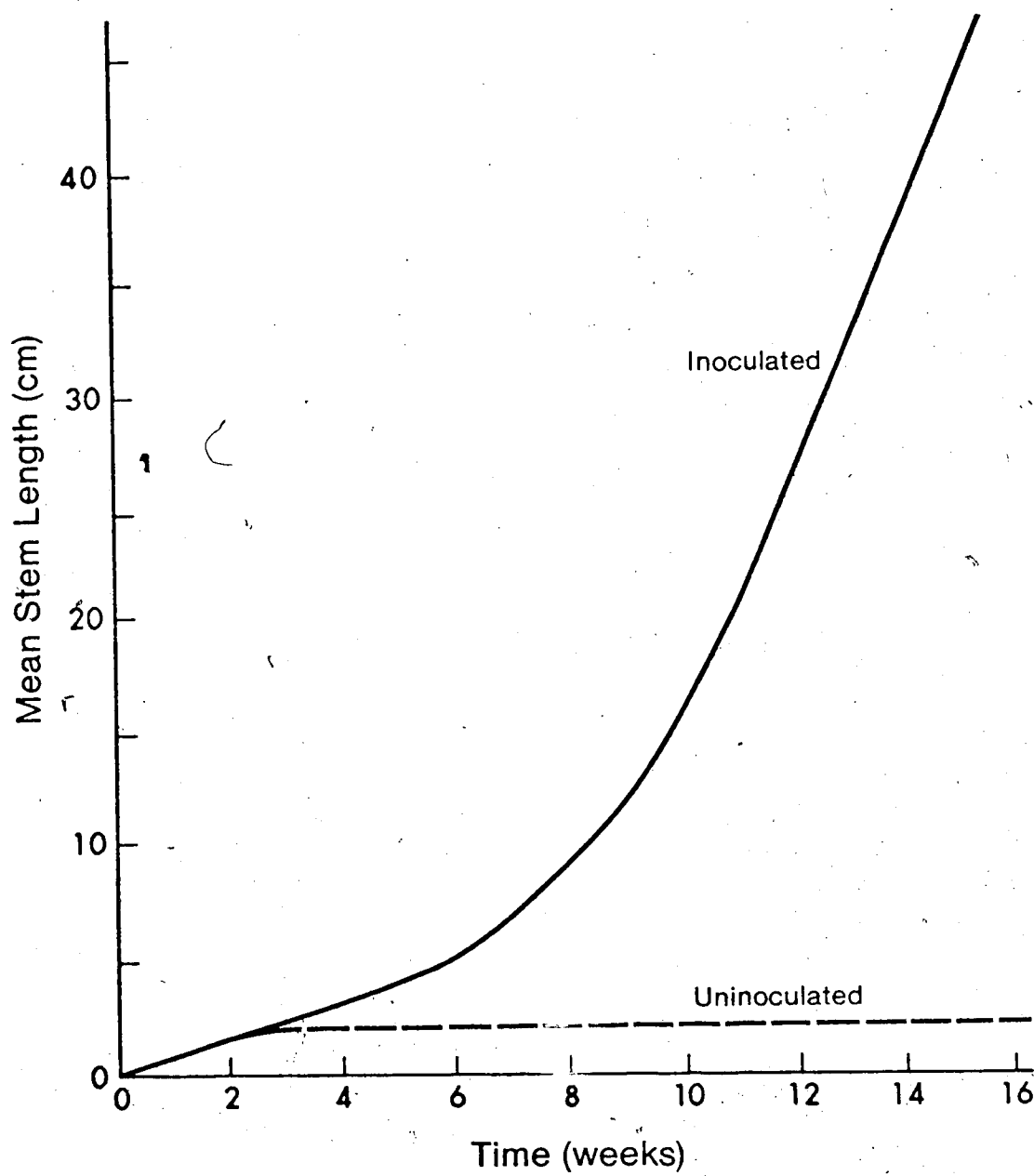


Figure 6. Mean growth curve for the fifteen genotypes of *D. sandwicense* inoculated with Rhizobium strain B and the uninoculated controls.

presented in Table 11. The pattern which emerged was the consistency with which general effects and average heterosis values exceeded the mean square for genotypes while the other components were less. These data suggested the presence of heterosis and additive genetic variance.

It was observed in the above study that the parental lines appeared to differ in growth before the initiation of the log growth phase and remained different until the study terminated. To confirm these results the primary shoot length was recorded weekly for the parental plants in block 1 of Experiment I. The growth curves for the five inbred lines are presented in Figure 7. The differences between these lines became evident as early as the sixth week after planting.

Hybrid vigor for forage yield has been generally considered to be absent in *Desmodium* when adequately supplied with nitrogen (McWhirter, 1969; Rotar and Chow, 1971). The heterosis present in the estimates of symbiotic nitrogen fixation must be due solely to the processes of nitrogen fixation because the non-nodulated controls did not accumulate dry matter. Therefore, the bottle jar is an effective tool for creating a nitrogen limiting environment for expressing differences in nitrogen fixing potential among population genotypes.

3. Linearity of acetylene reduction.

Acetylene reduction samples were prepared to determine

Table 11. Mean squares expressed as a percent of the genotype mean square of the diallel table genetic components for the initial study.

Component	Primary shoot length				Leaf areas				Total
	40 day	60 day	80 day	100 day	2nd node	5th node	8th node	10th node	
Genotypes									
General	100	100	100	100	100	100	100	100	100
Heterosis	125	180	206	160	164	107	108	92	141
Average	90	68	58	76	74	97	96	103	84
Line	242	295	152	270	296	493	131	208	175
Specific	54	54	73	99	5	10	96	63	73

*at 100 days.

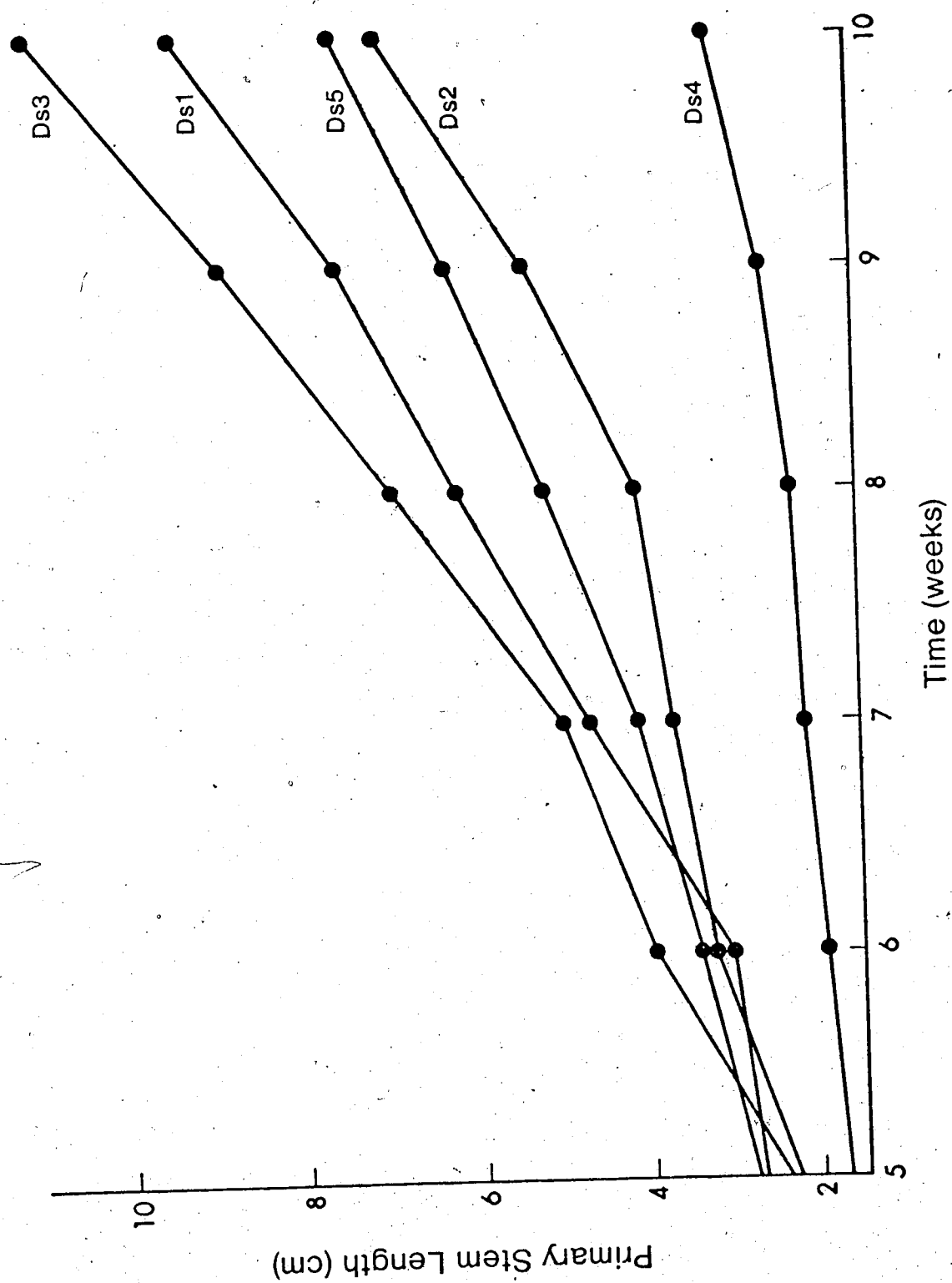


Figure 7. Growth curves for the D. sandwicense diallel lines inoculated with Rhizobium strain B.

if the reduction of acetylene by *Desmodium* was linear. The acetylene reduction diagrams for four lines are presented in Figure 8. All samples indicate linear reduction and a sample taken at one hour should adequately represent the amount of acetylene reduced in a one hour period. It should be cautioned that Figure 8 is intended to refer to the linear rate of acetylene reduction only and any comparison among genotypes should be avoided. The acetylene reduction curves for the five diallel lines are presented in Figure 9. These lines also present a linear reduction rate.

The above results suggested that symbiotic nitrogen fixation resulted from nodulated roots; however, the studies did not indicate whether the nodule was the site of nitrogen fixation. Nodules were removed from plant roots and placed separately in one vial and the roots in another. The pairs of vials were prepared and analysed by the acetylene reduction assay. Only the nodule sample indicated any acetylene reduction. It may be concluded that the nodules are the site for nitrogen fixation in *Desmodium*.

4. Bottle jar design.

During Experiment II in the quantitative genetic variation section (Level 3, Table 1), two shortcomings of the bottle jar design became apparent. Firstly, the plant roots would grow into the wick and it was difficult to separate the nodulated roots intact. It was also observed that the environmental variation in the quantitative genetic

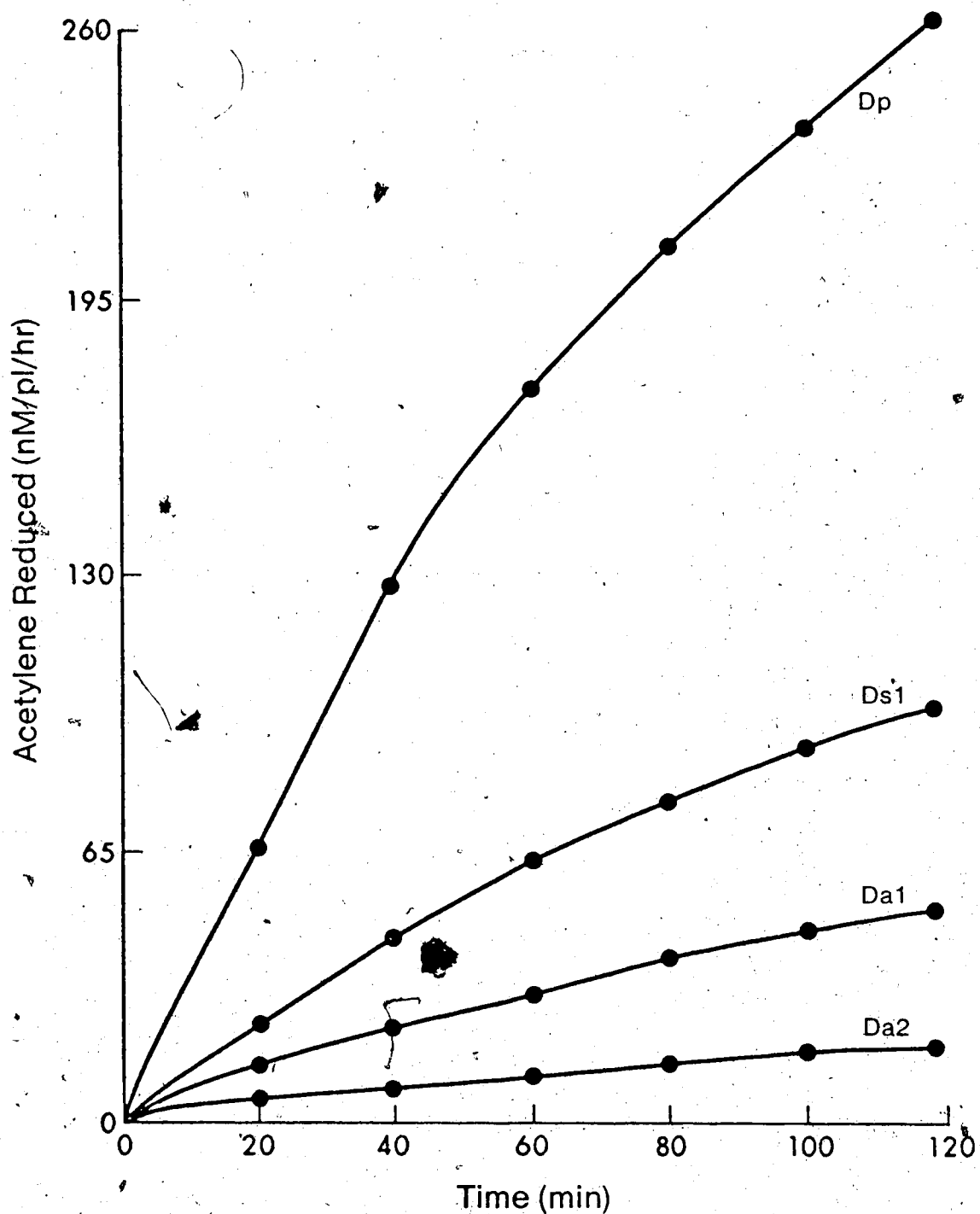


Figure 8. Acetylene reduction curves for several conspicuously different Desmodium species inoculated with Rhizobium strain A.

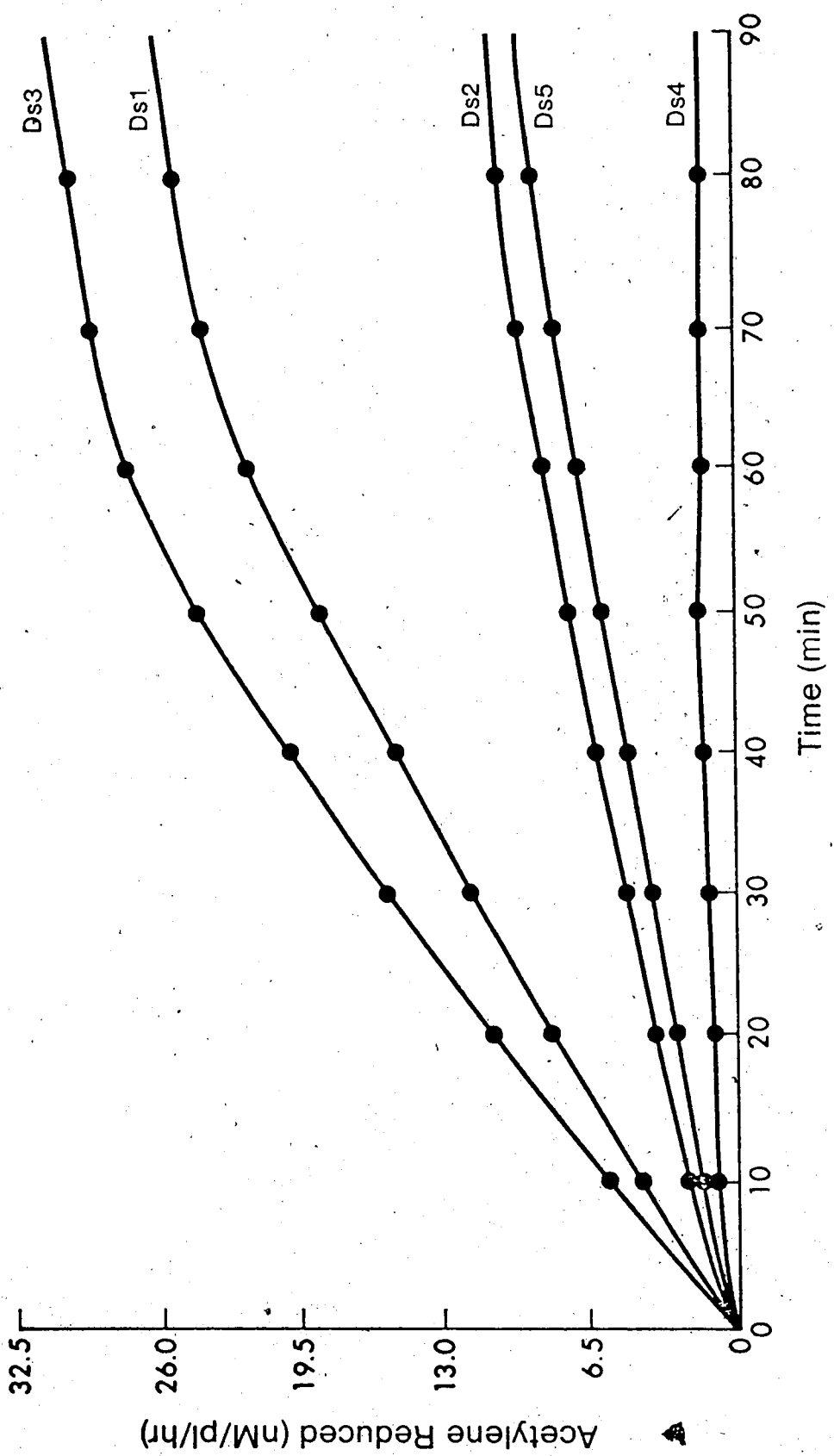


Figure 9. Acetylene reduction curves for the five D. sandwicense diallel lines inoculated with Rhizobium strain B.

studies was excessively large and appeared to be confined largely to within blocks which suggested that the bottle jar design might be manipulated in an effort to reduce this variation. It was suspected that the sand particle size and the distribution of the particles within the bottle jar may contribute to this environmental variation. Within bottle jar variation might be reduced if the sand particle size was more constant.

The analysis of variance for sand sizes is presented in Table 12. Significant differences were recorded for sand size means. The cell means and within-cell differences indicated that some means and differences had larger variances than others (Table 13). The < 4.0 mm sand size means were found to be less for dry weight and shoot length while the < 1.0 mm size had the lowest difference and variance for the within-cell difference between the two plants. Sand sizes with large inter-particle spaces appeared to have larger within cell differences suggesting that perhaps small sand grains assist capillary movement of the nutrient solution.

The germination of the plants in the above experiment was of considerable interest. It was noted that the < 1.0 mm size was the quickest to germinate while the 1.0 - 3.0 mm size emerged 2-3 days later. The other two sizes were intermediate with the < 4.0 mm size being slightly earlier. The sand surface of the 1.0 - 3.0 mm and 1.0 - 1.5 mm sizes dried before the rock mulch could be applied while the plant

Table 12. Mean squares from the analysis of variance of the sand size experiment.

Source	df	Dry weight (mg/pl)	Shoot length (cm/pl)
Blocks	5	455.80	2.57
Sand size	3	1718.92*	35.15**
Interaction	15	1191.83	16.99*
Error	23	568.70	5.62

*significant at the 5% level of probability.

**significant at the 1% level of probability.

Table 13. Means and standard deviations and within-cell differences and standard deviations for the sand size experiment.

Sand Size (mm)	Dry wt. (mg/pl)		Shoot length (cm/pl)	
	Mean	Wcd [†]	Mean	Wcd [†]
< 4.0	42.4±11.3	15.2± 9.4	11.9±1.3	1.5±1.1
1.0-3.0	63.4±41.7	37.2±43.3	15.0±4.4	3.6±3.9
1.0-1.5	67.7±26.5	26.0± 9.5	15.5±3.0	2.8±1.1
< 1.0	62.4±13.7	7.2± 4.9	14.2±1.9	0.9±0.5

[†]within-cell difference.

roots of the 1.0 - 3.0 mm size were found to be predominantly confined to the wick area. This information suggested that seeds planted in the finer sand sizes imbibed water more readily. These results may be due to a uniform capillary movement of the nutrient solution within the < 1.0 mm sand size.

The reduction of the within bottle jar variation would reduce environmental variation; therefore the modified bottle jar was accepted as a less variable container.

5. Analysis of variance of population means for individual epistatic effects.

Voigt *et al.* (1966) stated that an analysis of population means from a cross between two inbred lines "explains as much of the variance among means as possible in terms of additive gene action". Meredith and Laster (1975) report that "the estimates of A and D are independent and uncorrelated only if E is not significant". To investigate these seemingly contradictory statements a coded matrix for the genetic parameters (Mather and Jinks, 1971) was constructed and three types of least squares analysis of this matrix were investigated. The three types of analyses, differing in order of parameter input, were (a) the forward solution least squares analysis which assumed a simple model and which explained the maximum amount of the epistatic effects in terms of additive gene action, (b) a restricted stepwise least squares analysis which assumed a simple model

by forcing into the analysis the additive and dominance effect vectors and then explained the remaining variance among the population means by entering the coded interaction vector with the largest partial correlation. The remaining vectors were entered by the same criterion until the variance of the population means was explained. And (c) a free stepwise least squares analysis where the coded parameter vectors were entered in terms of the largest partial correlation. This analysis assumed no model.

The three analyses of variance of the population means are presented in Table 14. For this study the restricted stepwise least squares analysis has been assumed to be the superior method due to the free partition of the epistatic effects. If the epistatic effects were not partitioned, the restricted stepwise least squares analysis was identical to the forward solution least squares analysis for order of parameter input.

According to Meredith and Lester (1975) the significance of the epistatic effects caused the loss of independence between the additive and dominance effects. To present the relationships between the genetic parameters and the population means path diagrams (Li, 1975) were constructed for the above analysis. The path diagrams are presented in Figure 10. The additive and dominance effects remain orthogonal for both the direct and the indirect pathways. The non-orthogonality was between the main effects and the interaction effects. Because of the large

Table 14. Mean squares from the genetic analysis of variance of population means for the acetylene reduced (nM/pl/hr) for Experiment IV.

Source	df	Forward solution		Stepwise least squares	
		least squares	Restricted	Free	Free
Blocks	7	556.4		556.4	556.4
Populations	5	3689.3**		3689.3**	3689.3**
Additive	1	1923.2* (1+)		1923.2* (1)	7.2 (5)
Dominance	1	8494.4**(2)		8494.4**(2)	8494.4**(1)
Epistasis	3	2676.1		2676.1	3314.9**
Add - Add	1	4995.6**(3)		185.5 (5)	185.5 (4)
Add - Dom	1	359.1 (4)		359.1 (4)	2275.2**(3)
Dom - Dom	1	2676.1**(5)		7484.6**(3)	7484.6**(2)
Error	35	401.8		401.8	401.8

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

+Order of entry into the regression.

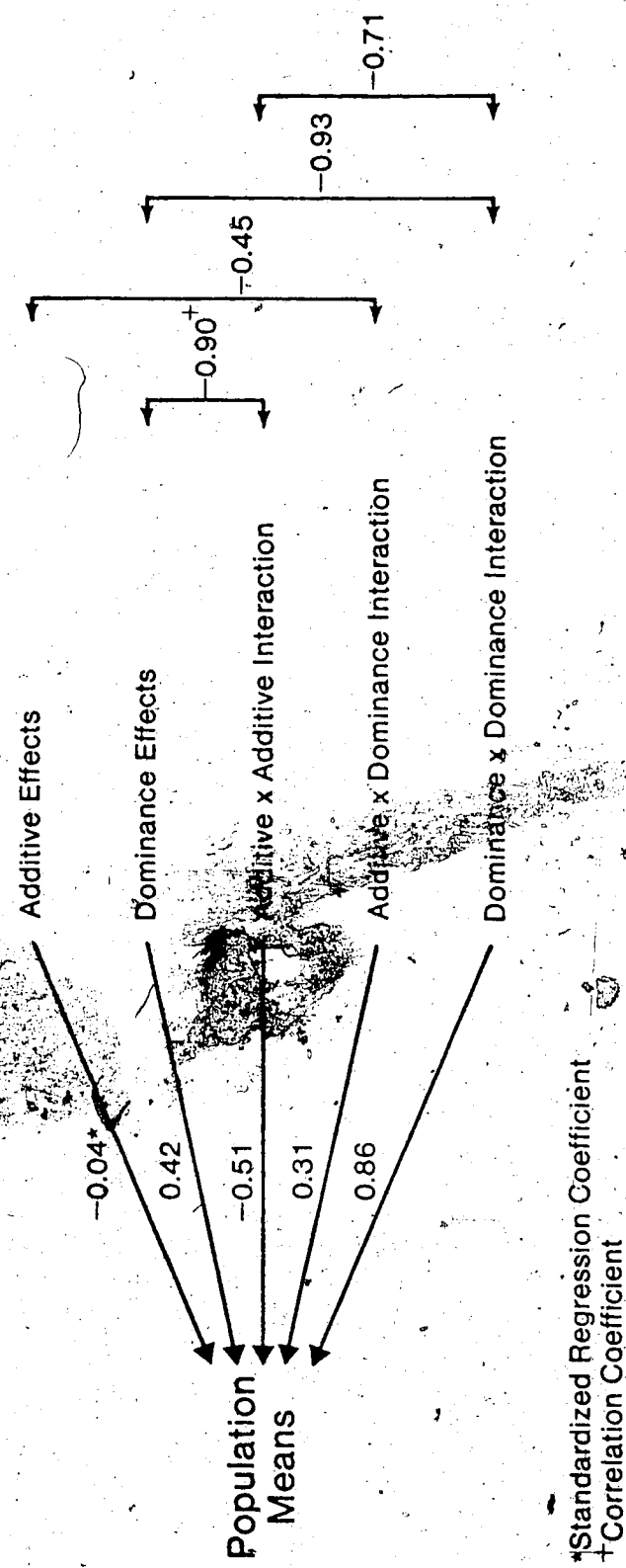


Figure 10. Path diagrams of the analysis of population means for acetylene reduced (nM/p/hr) for Experiment IV.

coefficients of determination (r^2) between the main effects and the interaction effects the analysis of variance of population means should be interpreted with caution, since a large proportion of the main effects may be explained as interaction effects. However, the analysis is of some value because the main effects were orthogonal even in the presence of significant epistatic effects and also provided an estimate of the significance of the interaction parameters.

6. Strain-specific ineffective nodulation interaction.

Several lines of conspicuously different *Desmodium* species were tested with the selected strains to determine the type of host-strain interaction. The interactions are presented in Table 15. *D. aparines* lines Da1, Da2 and Da3 were found to be ineffectively nodulated by strain B. These lines also exhibited small seed size and all originated from Costa Rica or Guatemala (Imrie, 1973). The *D. aparines* lines Da4 and Da5 exhibited effective nodulation interactions with strain B as well as large seeds. These lines were originally collected from Colombia and Mexico, respectively.

B. Qualitative genetic variation.

The area of study of the inheritance of nitrogen fixation which has commanded the greatest interest has been the study of non-nodulating or strain-specific ineffective nodulation. These conditions were found to be produced by

Table 15. Strain-specific interactions of *Desmodium* lines with *Rhizobium* strains.

Lines	Rhizobium Strain	
	A	B
Ds1	E	E
Ds2	E	E
Ds3	E	E
Ds4	E	E
Ds5	E	E
Da1	E	I
Da2	E	I
Da3	E	I
Da4	E	E
Da5	E	E
Dp	E	E
Du	E	E
Dsp	E	E*
Dse	I?	E*

E - effective nodulation (green plant).

I - ineffective nodulation (yellow-chlorotic plant).

? - not verified.

* - appears to be pH sensitive.

major genes. In preliminary experiments of this study in Desmodium it was noted that a strain-specific ineffective nodulation interaction was found between some D. canariensis lines when inoculated with strain B. The numbers of observations for each of the scores of the F₂ plants for nodulation interaction and spotted leaf character are presented in Table 16. The statistical analyses of the F₂ plants suggested that inheritance of this strain-specific ineffective nodulation interaction was conditioned by a dominant allele at a single locus.

The analysis of the leaf flecking character indicated the presence of complementary gene interaction in the Ds1 X Da2 cross. The Ds1 X Da1 cross appeared to be segregating at a single locus as the data conformed to a 3:1 segregation ratio. The joint segregation of the ineffective/effective nodulation and spotted/plain leaf character for the Ds1 X Da cross suggested that the two characters were independent in inheritance.

To test the hypothesis that a single dominant gene conditioned the strain-specific ineffective nodulation interaction between some D. canariensis lines and strain B, the backcrosses to Ds1 were grown and the plants scored for green vs yellow-chlorotic color and for spotted vs plain leaves. The character scores for the backcross plants are presented in Table 17. These data for nodulation conformed to the expectations of a single gene hypothesis thereby suggesting the single gene hypothesis for the strain-

Table 16. Observed nodulation and leaf fleck scores and Chi-square probabilities for the F₂ Plants of *Dermodius* crosses inoculated with strain B.

Cross	Nodulation		Leaf fleck		Joint segregation			
	Ineff.	Eff.	Chi-sq.	Chi-sq.	prob.	Plain prob.	Chi-sq.	probability
	(I)	(E)	(3:1)	(S)	(P)	(3:1)	I/S	I/P E/S E/P 9:3:3:1
Ds1 X Da1	88	36	.80-.70	.33	.91	.80-.70	.61	.27 .30 .6 .20-.10
Ds1 X Da2	91	24	.80-.70	.68	.47	.70-.60*	.38	.51 .7 .17 -----

--calculation not available.

*complementary gene interaction (9:7).

Table 17. Observed nodulation and leaf fleck scores and Chi-square probabilities for the backcross plants of the Desmodium crosses inoculated with strain-B.

Cross	Ineff. (I)	Eff. (E)	Chi-sq. prob. (1:1)	Leaf fleck			Joint segregation		
				Chi-sq. prob. (1:1)	Plain (P)	Chi-sq. prob. (1:1)	I/S	I/P	E/S
						Chi-sq. prob. (1:1:1)			
Ds1 X Da1	35	36	.98-.95	42	29	.30-.20	21	14	21
Ds1 X Da2	41	32	.50-.40	32	41	.50-.40*	15	26	17
Dev. Chi-sq.	76	68	.70-.60	74	70	.90-.80*	36	40	38
Heterogeneity Chi-sq.			.60-.50			.20-.10			.20-.10

*actual segregation ratio (1:1:1:1).

specific ineffective nodulation in *Desmodium*. These results have been published by Pinchbeck (1978) (Appendix D) and are in accord with the published results for soybean where three single dominant genes were reported for strain-specific ineffective interactions (Caldwell, 1966; Vest, 1970; Vest and Caldwell, 1972).

Seventy randomly selected ineffectively nodulated F₂ plants were inspected for nodulation characteristics and re-planted in pots and supplied with nitrogen. All ineffectively nodulated plants responded to nitrogen fertilizer. The roots of the ineffectively nodulated plants exhibited various degrees of nodule formation. The majority of nodules were small whereas a few were near normal size. All nodules lacked pink pigmentation, a sign of an active nodule (Vincent, 1970). These nodules appeared to be similar to those reported by Caldwell (1966) and Vest (1970) for the Rj₂ and Rj₃ genes in soybean.

The symbols N_i/n_i are suggested for the designation of the *D. saperines* strain-specific ineffective gene. *D. sandwicense* has the genotype n_i, n_i while the ineffectively nodulated *D. saperines* lines have the genotype N_i, N_i.

Similar phenotypes for ineffective nodulation interaction were exhibited by Da1 and Da2 lines. However, it is not known whether these phenotypes were conditioned by similar or different genes. To confirm this question a standard test of functional allelism (Srb, Owen and Edgar, 1965) would need to be applied to the F₂ generation of a Da1

X Da2 cross.

Experimentation for the relationship between the Ni gene and seed size was not pursued in this study as a single gene influencing seed size has not been reported. This avenue would be best examined by exploring the fertile D. parinense X D. uncinatum cross since both species exhibit the extremes for seed size and have different genotypes for the Ni gene.

C. Quantitative genetic variation.

A great need in international agriculture has been to increase total symbiotic nitrogen fixation. The traditional method of increasing total fixation has been manipulation of the Rhizobium strains used for inoculum (Vincent, 1974). An alternate research avenue is to establish and exploit the variability within the host. It was an objective of this study to investigate the hypothesis that there is variation within Desmodium for total symbiotic nitrogen fixation and that this variation has a heritable basis. The diallel cross and a cross between two inbred lines have been employed as quantitative genetic designs to establish this genetic variation.

1. Diallel cross.

a. Experiment I.

The analysis of variance* for two characters from

Experiment I is presented in Table 18. Tukey's test for non-additivity of the data (Snedecor and Cochran, 1967) indicated that a transformation of the data was necessary. Vincent (1970) suggested that dry weight accumulation data should be analysed in the logarithmic scale. Non-significance for genotypes was recorded for dry weight and the cause appeared to be block 2 which was significantly different from the remainder (Table 19). This block may have differed for three reasons: (a) due to the mechanical failure of the gas chromatograph this block was harvested one week later than the remainder, (b) block 2 may have been inoculated with a contaminated broth of *Rhizobium* since the values of parent 4 and some of its crosses are greatly increased, or (c) that the bottle type used in block 2 created a different environment thereby affecting the dry weight accumulation results. Block 2 was deleted from further analyses.

Principal components analysis of the collected characters indicated that the characters were highly correlated and related to one principal factor (Table 20).

The genotype means for the characters measured (transformed to \log_{10}) are presented in Table 21. The diallel genotypes revealed that most crosses exhibited heterosis suggesting that heterosis may be a characteristic

*Due to a mechanical failure of the gas chromatograph, the acetylene reduction data were not collected for blocks 2 and 3 of this study. Therefore acetylene reduction character was not reported for Experiment I.

Table 18. Mean squares of two original characters for Experiment I.

Source	d.f.	Total shoot length (cm/pl)	Total dry weight (mg/pl)
Blocks	3	2084**	48750**
Genotypes	14	109*	2542
Error	42	44	1873
Non-additivity	1	610**	23439**
Residual	41	31	1347

*significant at the 5% level of probability.

**significant at the 1% level of probability.

Table 19. Dry weight (mg/pl) of the diallel genotypes for Experiment I.

Genotype	Block			
	1	2	3	4
Ds1	70	136	51	111
Ds2	41	63	31	35
Ds3	91	118	43	39
Ds4	11	281	51	37
Ds5	45	142	49	65
Ds1 x Ds2	64	109	107	113
Ds1 x Ds3	115	163	74	102
Ds1 x Ds4	48	316	117	120
Ds1 x Ds5	60	137	81	112
Ds2 x Ds3	73	194	123	54
Ds2 x Ds4	42	157	34	112
Ds2 x Ds5	33	214	66	122
Ds3 x Ds4	52	300	65	96
Ds3 x Ds5	55	220	48	141
Ds4 x Ds5	40	170	51	136
Block means	56	181	66	93

Table 20. Correlations and principal eigenvector loadings for Experiment I.

Character	Correlations							
	1	2	3	4	5	6		
Main shoot length	1	1.00				.86		
Secondary shoot length	2	.43	1.00			.77		
Total shoot length	3	.91	.71	1.00		.97		
Leaf dry weight	4	.73	.75	.87	1.00	.96		
Shoot, dry weight	5	.86	.68	.96	.94	1.00	.99	
Total dry weight	6	.78	.74	.90	.99	.97	1.00	.98

Table 21. Logio means and standard error of the difference between two means for Experiment I.

Genotype	Main shoot length (cm/pl)	Secondary shoot length (cm/pl)	Total shoot length (cm/pl)	Leaf weight (mg/pl)
Ds1	1.090	.768	1.740	1.271
Ds2	0.907	.417	1.408	0.962
Ds3	1.052	.258	1.601	1.125
Ds4	0.826	.067	1.331	0.778
Ds5	1.024	.371	1.591	1.123
Ds1 x Ds2	1.146	.939	1.825	1.379
Ds1 x Ds3	1.128	.820	1.849	1.385
Ds1 x Ds4	1.010	.735	1.813	1.349
Ds1 x Ds5	1.083	.759	1.795	1.249
Ds2 x Ds3	1.102	.736	1.756	1.319
Ds2 x Ds4	0.985	.645	1.638	1.124
Ds2 x Ds5	1.056	.723	1.685	1.198
Ds3 x Ds4	1.097	.364	1.711	1.234
Ds3 x Ds5	1.102	.565	1.724	1.276
Ds4 x Ds5	1.055	.576	1.701	1.184
Se of d.	0.084	0.385	0.141	0.161

Table 21 (cont.). Log₁₀ means and standard error of the difference between two means for Experiment I.

Genotype	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Nitrogen fixation "factor" (mg/pl)
Ds1	1.261	1.866	0.366
Ds2	1.011	1.549	-0.083
Ds3	1.089	1.728	-0.404
Ds4	0.840	1.439	-1.706
Ds5	1.078	1.719	-0.362
Ds1 x Ds2	1.346	1.960	0.784
Ds1 x Ds3	1.290	1.980	0.725
Ds1 x Ds4	1.236	1.943	0.542
Ds1 x Ds5	1.246	1.912	0.419
Ds2 x Ds3	1.249	1.895	0.438
Ds2 x Ds4	1.120	1.735	-0.206
Ds2 x Ds5	1.206	1.808	0.121
Ds3 x Ds4	1.139	1.837	0.041
Ds3 x Ds5	1.197	1.856	0.222
Ds4 x Ds5	1.165	1.814	0.004
se of d.	0.105	0.148	0.581

in Desmodium for total symbiotic nitrogen fixed.

The diallel analyses of variance for the logio transformed characters in Experiment I are presented in Table 22. All characters including the "Nitrogen fixation" factor were noted to have a similar analysis pattern. The significant mean squares for genotypes suggested that genetic variation for the characters existed within the D. sandwicense population. Significant general effects indicated that a portion of this variation was heritable. Although heterosis was not significant, average heterosis was shown to be highly significant indicating that the F₁ progeny means were significantly different from the parental means.

The estimates of the G.C.A. effects and variety constants are presented in Table 23. Line Ds1 exhibited positive G.C.A. effects and variety constants for total dry weight while line Ds4 was found to have a negative G.C.A. effect and variety constant.

Estimates of the heterosis constants revealed that, in general, the line and specific heterosis constants were small in magnitude in comparison to the average heterosis constant (Table 24). These results were expected since line and specific heterosis effects were not significant in the diallel analysis of variance.

The estimates of the S.C.A. effects failed to define any hybrid which was superior to the remainder (Table 25). The negative S.C.A. estimates for inbred lines suggested

Table 22. Mean squares from the diallel analysis of variance for Experiment I.

Source	Main shoot length (cm/pl)	Secondary shoot length (cm/pl)	Total shoot length (cm/pl)	Leaf dry weight (mg/pl)	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Nitrogen fixation "factor"
Blocks	2	.437**	.483**	.143*	.455**	.455**	.194*
Genotypes	14	.023*	.610**	.065*	.079	.048*	.068*
General	4	.041*	1.105**	.099*	.128*	.091**	.110*
Heterosis	10	.015	.413	.051	.059	.031	.052
Average	1	.114**	3.593**	.465**	.473**	.268**	.457**
Line	4	.005	.108	.011	.020	.006	.010
Specific	5	.004	.020	.002	.008	.003	.004
Error	28	.011	.222	.030	.039	.016	.033
Non-additivity	1	.001	1.508**	.000	.003	.004	.001
Residual	27	.010	.175	.031	.040	.017	.034

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

Table 23. Estimates of G.C.A. effects and variety constants for Experiment I.

	Main shoot length (cm/pl)	Secondary shoot length (cm/pl)	Total shoot length (cm/pl)	Leaf dry weight (mg/pl)	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Nitrogen fixation "factor"
<u>Griffing's G.C.A. effects</u>							
$g(Ds1)$.049	.316	.093	.089	.103	.101	.457
$g(Ds2)$	-.029	.084	-.007	-.049	-.034	-.046	-.119
$g(Ds3)$.033	-.104	.009	.025	.040	.030	.088
$g(Ds4)$	-.058	-.305	-.093	-.078	-.105	-.087	-.433
$g(Ds5)$	-.006	.009	-.003	-.003	-.004	-.002	.006
$se[g(1)]$.020	.092	.025	.034	.039	.035	.139
$se[g(i)-g(j)]$.030	.146	.039	.054	.061	.056	.220
<u>Gardner's variety constants</u>							
mean	.980	-.064	1.534	1.052	1.056	1.660	.618
$v(Ds1)$.111	.720	.206	.219	.205	.206	.618
$v(Ds2)$	-.073	.046	-.126	-.090	-.044	-.111	-.365
$v(Ds3)$.072	-.201	.067	.073	.033	.068	.214
$v(Ds4)$	-.154	-.735	-.203	-.274	-.216	-.221	-.089
$v(Ds5)$.044	.170	.057	.071	.023	.059	.256

Table 24. Estimates of Gardner's average, line and specific heterosis constants for Experiment I.

	Main Heterosis Constant	Secondary shoot length (cm/pl)	Total shoot length (cm/pl)	Leaf dry weight (mg/pl)	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Nitrogen fixation "factor"
$h(\text{ave})$.107	.310	.215	.218	.164	.214	.927
$h(Ds1)$	-.015	-.027	-.009	-.015	-.022	-.004	-.081
$h(Ds2)$.018	.079	.031	.025	.037	.023	.150
$h(Ds3)$	-.008	-.028	-.020	-.008	-.018	-.010	-.044
$h(Ds4)$.044	.013	.056	.074	.035	.055	.260
$h(Ds5)$	-.039	-.038	-.060	-.093	-.032	-.064	-.285
$h(Ds1, Ds2)$.038	-.016	.012	.034	.031	.020	.097
$h(Ds1, Ds3)$	-.027	.051	-.009	-.024	-.008	-.018	-.059
$h(Ds1, Ds4)$.016	.021	.014	.048	.009	.025	.106
$h(Ds1, Ds5)$	-.027	-.056	-.018	-.058	-.032	-.027	-.145
$h(Ds2, Ds3)$.007	.037	.024	.024	.016	.030	.098
$h(Ds2, Ds4)$	-.050	.001	-.035	-.064	-.041	-.051	-.197
$h(Ds2, Ds5)$.005	-.022	-.002	-.006	-.007	.002	.002
$h(Ds3, Ds4)$.016	-.094	-.007	-.018	-.007	-.005	-.046
$h(Ds3, Ds5)$.004	.006	-.009	.018	.000	-.006	.006
$h(Ds4, Ds5)$.018	.072	.028	.034	.039	.031	.137

Table 25. Estimates of Griffing's S.C.A. effects for Experiment I.

S.C.A. effect	Main shoot length (cm/pl)	Secondary shoot length (cm/pl)	Total shoot length (cm/pl)	Leaf dry weight (mg/pl)
s(Ds1)	-.059	-.311	-.090	-.136
s(Ds2)	-.086	-.521	-.140	-.171
s(Ds3)	-.064	-.393	-.094	-.127
s(Ds4)	-.109	-.525	-.139	-.139
s(Ds5)	-.038	-.247	-.081	-.092
se[s(i)]	.041	.188	.051	.069
se[s(i)-s(j)]	.055	.252	.068	.093
s(Ds1, Ds2)	.076	.146	.095	.097
s(Ds1, Ds3)	-.004	.195	.023	.047
s(Ds1, Ds4)	.068	.283	.071	.113
s(Ds1, Ds5)	-.022	-.002	-.009	.015
s(Ds2, Ds3)	.048	.331	.081	.103
s(Ds2, Ds4)	.021	.390	.055	.087
s(Ds2, Ds5)	.028	.176	.050	.054
s(Ds3, Ds4)	.072	.158	.058	.086
s(Ds3, Ds5)	.013	.102	.026	.018
s(Ds4, Ds5)	.057	.218	.095	.097
se[s(i,j)]	.052	.238	.064	.088
se[s(i,j)-s(i,k)]	.078	.356	.096	.132
se[s(i,j)-s(k,l)]	.071	.325	.088	.120

Table 25 (cont.). Estimates of Griffing's S.C.A. effects for Experiment I.

S.C.A. effect	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Nitrogen fixation "factor"
s(Ds1)	-.132	-.140	-.549
s(Ds2)	-.167	-.162	-.747
s(Ds3)	-.152	-.134	-.580
s(Ds4)	-.192	-.190	-.840
s(Ds5)	-.065	-.088	-.374
se[s(i)]	.079	.072	.284
se[s(i)-s(j)]	.106	.097	.381
s(Ds1, Ds2)	.113	.102	.446
s(Ds1, Ds3)	.044	.046	.179
s(Ds1, Ds4)	.154	.126	.518
s(Ds1, Ds5)	-.047	.006	-.044
s(Ds2, Ds3)	.115	.108	.468
s(Ds2, Ds4)	.066	.065	.346
s(Ds2, Ds5)	.039	.049	.234
s(Ds3, Ds4)	.102	.092	.386
s(Ds3, Ds5)	.043	.023	.127
s(Ds4, Ds5)	.096	.097	.431
se[s(i,j)]	.100	.091	.359
se[s(i,j)-s(i,k)]	.149	.137	.538
se[s(i,j)-s(k,l)]	.136	.125	.491

that homozygous lines may be exhibiting inbreeding depression for total symbiotic nitrogen fixed.

b. Experiment II.

The data from Experiment I indicated that total symbiotic nitrogen fixed appeared to be heritable. To further investigate the hypothesis that there exists genetic variation for nitrogen fixation, the diallel cross was grown with Rhizobium strain, A to determine if the mode of inheritance was consistent for a different strain.

Principal components analysis of the characters measured for Experiment II revealed that all characters were highly correlated and related to one factor (Table 26).

The genotype means from the log₁₀ transformed characters are presented in Table 27. An examination of the individual genotype means revealed that every cross exhibited heterobeltiosis except cross Ds2 x Ds3 which portrayed only complete heterosis for the acetylene reduction assay.

The diallel analyses of variance for the log₁₀ transformed characters for Experiment II are given in Table 28. There were significant differences among genotypes which confirmed that genetic variation was present in the diallel population. General effects, heterosis and average heterosis components were significant while line and specific heterosis were non-significant. The genetic variance was confined to among parental lines and between the progeny and

Table 26. Correlations, coefficients and principal eigenvector loadings for Experiment II.

Character	Correlations				Eigenvector
	1	2	3	4	
Leaf dry weight	1	.99			
Shoot dry weight		2	.97	1.00	.99
Total dry weight			3	.99	.99
Acetylene reduced				4	.93

Table 27. Logio means and standard error of the difference between two means for Experiment II.

Genotype	Leaf dry weight (mg/pl)	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)
Ds1	1.627	1.056	1.732
Ds2	1.660	1.147	1.777
Ds3	1.838	1.355	1.962
Ds4	1.749	1.127	1.842
Ds5	1.795	1.272	1.909
Ds1 x Ds2	1.842	1.308	1.954
Ds1 x Ds3	1.948	1.452	2.070
Ds1 x Ds4	2.003	1.466	2.115
Ds1 x Ds5	1.950	1.457	2.073
Ds2 x Ds3	1.914	1.425	2.037
Ds2 x Ds4	2.035	1.494	2.145
Ds2 x Ds5	2.036	1.525	2.153
Ds3 x Ds4	2.214	1.683	2.327
Ds3 x Ds5	2.141	1.640	2.261
Ds4 x Ds5	1.941	1.346	2.042
se of d.	0.164	0.199	0.172

Table 27 (cont.). Log₁₀ means and standard error of the difference between two means for Experiment II.

Genotype	Acetylene reduced (nmoles/pl/hr)	Nitrogen fixation "factor"
Ds1	1.648	1.831
Ds2	1.488	1.831
Ds3	1.864	1.847
Ds4	1.506	1.835
Ds5	1.726	1.842
Ds1 x Ds2	1.759	1.845
Ds1 x Ds3	1.880	1.853
Ds1 x Ds4	1.835	1.855
Ds1 x Ds5	1.996	1.855
Ds2 x Ds3	1.863	1.851
Ds2 x Ds4	1.898	1.857
Ds2 x Ds5	2.122	1.861
Ds3 x Ds4	2.083	1.870
Ds3 x Ds5	2.030	1.866
Ds4 x Ds5	1.732	1.849
se of d.	0.206	0.460

Table 28. Mean squares from the diallel analysis of variance for Experiment II.

Source	df	Leaf dry weight (mg/pl)	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Acetylene reduced (nM/pl/hr)	Nitrogen fixation "factor"
Blocks	3	2.137**	2.401**	2.209**	4.778**	17.427**
Genotypes	14	.218**	.261**	.225*	.290**	1.559**
General	4	.169*	.212*	.172*	.241*	1.150*
Heterosis	10	.237**	.280**	.246**	.310**	1.722**
Average	1	1.923**	2.214**	1.989**	1.999**	13.622**
Line	4	.018	.017	.017	.046	.084
Specific	5	.076	.103	.081	.183	.652
Blocks x Genotypes	42	.065	.090	.070	.138*	.530
Blocks x general	12	.061	.062	.060	.118*	.423
Blocks x heterosis	30	.067	.102	.074	.146*	.572
Blocks x average	3	.095	.162	.108	.037	.657
Blocks x line	12	.055	.089	.062	.157	.517
Blocks x specific	15	.072	.100	.077	.159*	.599
Error	56	.054	.079	.059	.085	.423

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

parental means. Block x genotype interactions were not significant for dry weight characters but significant for the acetylene reduction assay. This component was distorted because of a problem of the separation of the nodulated roots from the lamp wick (Figure 2) in the preparation of the plants for the assay. As a result of this problem the bottle jar design was modified (Figure 3).

The estimates of the G.C.A. effects and variety constants are presented in Table 29 and line Ds3 was shown to have the greatest G.C.A. effect and variety constant for all characters. Line Ds1, however, exhibited a negative G.C.A. effect and variety constant which is the reverse of Experiment I. These results were of interest since line Ds3 was a Ds1 backcross line from an original Ds1 x *D. narinines* (Dai) cross. Since strain A was isolated from *D. narinines*, a strain x genotype interaction may be suggested.

The estimates of the heterosis constants again revealed that average heterosis was much larger than either the estimates of the line or specific heterosis constants (Table 30).

The S.C.A. effect estimates revealed inbreeding depression in the inbred lines for all characters (Table 31). The cross Ds3 x Ds4 showed considerable S.C.A. effect for the dry weight characters. However, the cross Ds2 x Ds5 exhibited a superior S.C.A. effect for the acetylene reduction assay. The cross Ds2 x Ds5 also had a larger acetylene reduction assay S.C.A. value relative to its dry

Table 29. Estimates of G.C.A. effects and variety constants for Experiment II.

	Leaf dry weight (mg/pl)	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Acetylene reduced "factor" (nm/pl/hr)	Nitrogen fixation "factor" (nm/pl/hr)
<u>Griiffing's G.C.A. effects</u>					
$g(Ds1)$	-.069	-.072	-.069	-.030	-.160
$g(Ds2)$	-.047	-.036	-.045	-.051	-.117
$g(Ds3)$.060	.087	.066	.088	.190
$g(Ds4)$.030	-.008	.022	-.059	.005
$g(Ds5)$.026	.030	.027	.051	.083
$se[g(i)]$.028	.034	.029	.035	.078
$se[g(i)-g(j)]$.044	.053	.046	.055	.123
<u>Gardner's variety constants</u>					
mean	1.734	1.191	1.845	1.646	.476
$v(Ds1)$	-.106	-.136	-.113	.001	-.244
$v(Ds2)$	-.074	-.044	-.068	-.158	-.211
$v(Ds3)$.104	.164	.118	.218	.373
$v(Ds4)$.015	-.065	-.002	-.140	-.098
$v(Ds5)$.061	.081	.065	.080	.180

Table 30. Estimates of Gardner's average, line and specific heterosis constants for Experiment II.

Heterosis constant (mg/pl)	Leaf dry weight (mg/pl)	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Acetylene reduced (nm/pl/hr.)	Nitrogen fixation "factor"
h(ave)	.269	.288	.273	.274	.715
h(Ds1)	-.036	-.011	-.030	-.070	-.088
h(Ds2)	-.024	-.033	-.027	.067	-.028
h(Ds3)	.017	.012	.016	-.050	.007
h(Ds4)	.053	.056	.054	.027	.125
h(Ds5)	-.011	-.024	-.013	.027	-.016
h(Ds1, Ds2)	-.011	-.038	-.017	-.079	-.085
h(Ds1, Ds3)	-.035	-.044	-.037	-.029	-.094
h(Ds1, Ds4)	.028	.042	.031	.029	.085
h(Ds1, Ds5)	.017	.040	.023	.079	.094
h(Ds2, Ds3)	-.097	-.093	-.095	-.103	-.246
h(Ds2, Ds4)	.032	.047	.035	.034	.094
h(Ds2, Ds5)	.075	.084	.077	.148	.237
h(Ds3, Ds4)	.082	.086	.082	.148	.246
h(Ds3, Ds5)	.050	.050	.049	-.016	.093
h(Ds4, Ds5)	-.142	-.174	-.149	-.211	-.425

Table 31. Estimates of Griffing's S.C.A. effects for Experiment II.

S.C.A. effect	Leaf dry weight (mg/pl)	Shoot dry weight (mg/pl)	Total dry weight (mg/pl)	Acetylene reduced ("factor") (nm/pl/hr)	Nitrogen fixation "factor" (nm/pl/hr)
s(Ds1)	-.149	.183	-.157	-.122	-.401
s(Ds2)	-.158	-.163	-.159	-.240	-.453
s(Ds3)	-.194	-.202	-.196	-.140	-.483
s(Ds4)	-.225	-.240	-.228	-.205	-.584
s(Ds5)	-.170	-.172	-.171	-.206	-.463
se[s(1)]	.057	.069	.059	.071	.159
se[s(1)-s(j)]	.076	.092	.079	.095	.213
s(Ds1, Ds2)	.045	.033	.042	.011	.087
s(Ds1, Ds3)	.044	.053	.047	-.006	.098
s(Ds1, Ds4)	.128	.163	.136	.095	.344
s(Ds1, Ds5)	.080	.116	.089	.145	.273
s(Ds2, Ds3)	-.011	-.009	-.010	-.002	-.020
s(Ds2, Ds4)	.139	.155	.141	.179	.388
s(Ds2, Ds5)	.145	.148	.145	.293	.450
s(Ds3, Ds4)	.211	.221	.213	.226	.560
s(Ds3, Ds5)	.143	.140	.142	.062	.327
s(Ds4, Ds5)	-.028	-.060	-.034	-.089	-.124
se[s(1, j)]	.072	.087	.075	.090	.201
se[s(1, j)-s(1, k)]	.108	.130	.112	.135	.301
se[s(1, j)-s(k, l)]	.099	.119	.102	.123	.245

weight accumulation S.C.A. value, whereas, the cross Ds3 x Ds4 showed equal dry weight accumulation and acetylene reduction S.C.A. values. These results suggested that dry weight accumulation and the acetylene reduction assay, although assumed to measure symbiotic nitrogen fixation, may measure closely related components of a complex character.

c. Experiment III.

Since both strain and/or environmental effects appeared to be of importance in the preceding studies, Experiment III was initiated to determine if strain X genotype interactions were of significant importance in a *D. sandwicense* population.

A review of the original and \log_{10} transformed means for the within-strain and overall analyses for Experiment III illustrated that heterosis was of the approximate magnitude of 1.6 (Tables 32, 33 and 34; respectively). The low mean values for the acetylene reduction analysis in strain A (Table 32) appeared to be due to the nodules of that strain being in a state of senescence. Nodule senescence may have been due to the harsh "environmental" conditions imposed upon this experiment; yet strain B appeared unaffected. Since strain A was faster growing than strain B the nodules of strain A may have been sampled at a later physiological stage. The compatibility of the dry weight means may be explained by the releasing of nitrogen from the decaying nodules for plant growth. Further

Table 32. Means and standard errors of the difference between two means for the Rhizobium strain A inoculated blocks for Experiment III.

Genotype	Dry weight (mg/pl)		Acetylene reduced (nm/pl/hr)	
	original	log ₁₀	original	log ₁₀
Ds1	139	2.045	4.3	0.436
Ds2	223	2.284	5.7	0.650
Ds3	167	2.187	6.3	0.714
Ds4	180	2.115	9.9	0.476
Ds5	225	2.277	11.2	0.845
Ds1 x Ds2	249	2.396	6.7	0.774
Ds1 x Ds3	309	2.481	10.0	0.982
Ds1 x Ds4	211	2.314	5.9	0.740
Ds1 x Ds5	363	2.556	16.5	1.214
Ds2 x Ds3	82	1.876	1.7	0.188
Ds2 x Ds4	302	2.454	8.1	0.877
Ds2 x Ds5	248	2.361	8.3	0.712
Ds3 x Ds4	173	2.226	3.5	0.497
Ds3 x Ds5	375	2.556	15.6	1.067
Ds4 x Ds5	415	2.572	22.2	1.124
se of d.	75	0.167	3.8	0.207

Table 33. Means and standard error of the difference between two means for the Rhizobium strain B inoculated blocks for Experiment III.

Genotype	Dry weight (mg/pl)		Acetylene reduced (nM/pl/hr)	
	original	log ₁₀	original	log ₁₀
Ds1	205	2.270	18.0	1.219
Ds2	223	2.335	15.9	1.197
Ds3	177	2.111	12.5	0.819
Ds4	161	2.184	9.5	0.923
Ds5	169	2.187	13.1	1.021
Ds1 x Ds2	260	2.404	24.1	1.340
Ds1 x Ds3	493	2.690	36.5	1.551
Ds1 x Ds4	182	2.235	15.0	1.160
Ds1 x Ds5	347	2.500	25.1	1.326
Ds2 x Ds3	351	2.542	29.6	1.453
Ds2 x Ds4	161	2.121	15.1	1.037
Ds2 x Ds5	325	2.491	31.3	1.468
Ds3 x Ds4	302	2.473	21.1	1.317
Ds3 x Ds5	287	2.439	19.4	1.267
Ds4 x Ds5	329	2.493	12.8	1.099
se of d.	67	0.155	6.9	0.203

Table 34. Means and standard error of the difference between two means from the overall analysis for Experiment III.

Genotype	Dry weight (mg/pl)		Acetylene reduced (nm/pl/hr)	
	original	Log ₁₀	original	Log ₁₀
Ds 1	172	2.158	11.1	0.827
Ds 2	223	2.310	10.8	0.923
Ds 3	172	2.149	9.4	0.767
Ds 4	170	2.150	9.7	0.699
Ds 5	197	2.232	12.1	0.933
Ds 1 x Ds 2	254	3.400	15.4	1.057
Ds 1 x Ds 3	401	2.585	23.2	1.266
Ds 1 x Ds 4	196	2.274	10.5	0.950
Ds 1 x Ds 5	355	2.528	20.8	1.270
Ds 2 x Ds 3	216	2.209	15.7	0.820
Ds 2 x Ds 4	231	2.288	11.6	0.957
Ds 2 x Ds 5	287	2.426	19.8	1.090
Ds 3 x Ds 4	238	2.349	12.3	0.907
Ds 3 x Ds 5	331	2.498	17.5	1.167
Ds 4 x Ds 5	372	2.532	17.5	1.112
se of d.	50	0.114	3.9	0.145

experimentation would be necessary to clarify these results.

The within-strain diallel analyses of variance for the normal characters are presented in Table 35 and for comparison the log₁₀ transformed diallel analyses of variance is given in Table 36. The similarity of the two scales suggested that scale effects had been removed from this experiment through improved experimental and bottle jar design.

Significant variation was again found among the genotypes and the majority of this variation was due to average heterosis. Block X genotype interactions were also found to be significant for most genetic components suggesting genetic component X environment interactions. These results were expected since Vincent (1965) reported that the environment influenced symbiotic nitrogen fixation.

Factorial diallel analyses of variance for original and log₁₀ transformed characters are presented in Table 37. Strains were not significantly different for dry weight accumulation yet highly significant for the acetylene reduction assay for both scales due to nodule senescence in Strain A. Genotypes were highly significant for all characters and scales suggesting that genetic variation was present in this population and that this variation was scale independent.

A partition of the genotype variance suggested that a predominant portion of this variance was due to dominance deviations since general effects were significant for the

Table 35. Mean squares from the diallel analysis of variance for the original characters for Experiment III.

Source	Rhizobium Strain-A			Rhizobium Strain-B		
	Dry weight (mg/pl)	Acetylene reduced (nM/pl/hr)	(mg/pl)	Dry weight (mg/pl)	Acetylene reduced (nM/pl/hr)	(mg/pl)
Blocks	1	88167**	835**	31236*	858**	
Genotypes	14	34999**	125**	36155**	250**	
General Heterosis	4	29706**	212**	20009**	282**	
Line	10	37116**	90**	42614**	237**	
Average	1	98155**	67*	181352**	988**	
Specific	4	30939**	98**	25967**	154*	
Blocks X genotypes	5	29850**	88**	28183**	154*	
Blocks X general	4	17608*	171**	16847**	88*	
Blocks X heterosis	10	20946**	91**	22153**	106*	
Blocks X average	1	38593*	26	914	4	
Blocks X line	4	8820	145**	26901**	157*	
Blocks X specific	5	27119**	148**	22605**	47	
Error	29	5671	14	4478	47	

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

Table 36. Mean squares from the diallel analysis of variance for the \log_{10} transformed characters for Experiment III.

Source	Rhizobium Strain A			Rhizobium Strain B		
	Dry weight df (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight reduced (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight reduced (mg/pl)	Acetylene reduced (nm/pl/hr)
Blocks	1	.410**	2.943**	.204*	.897**	
Genotypes	14	.163**	.313**	.121**	.176**	
General	4	.102*	.348**	.047	.170**	
Heterosis	10	.187**	.299**	.150**	.178**	
Average	1	.520**	.468**	.653**	.838**	
Line	4	.161**	.323**	.121**	.132*	
Specific	5	.141**	.248**	.073*	.082	
Blocks X genotypes	14	.075*	.233**	.072**	.108*	
Blocks X general	4	.034	.152*	.039	.129*	
Blocks X heterosis	10	.090**	.265**	.085**	.099*	
Blocks X average	1	.473**	.703**	.001	.093	
Blocks X line	4	.046	.228**	.146**	.178**	
Blocks X specific	5	.049	.207**	.053	.037	
Error	29	.028	.043	.024	.041	

*significant at the 5% level of probability.
**significant at the 1% level of probability.

Table 37. Mean squares from a factorial diallel analysis of variance for Experiment III.

Source	Original		Log ₁₀	
	Dry weight df. (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight reduced (mg/pl)	Acetylene reduced (nm/pl/hr)
Strains	1 12523	342690**	.080	6.314**
Blocks	1 112180**	169254**	.596**	3.546**
Strains X blocks	1 7223	8	.018	.295
Genotypes	14 47483**	14407**	.178**	.227**
General	4 21909	11281	.079	.219*
Heterosis	10 57714**	15675**	.217**	.230**
Average	1 273652**	78416**	1.168**	1.279**
Line	4 36878**	9337	.147*	.156
Specific	5 31195**	8154	.083	.080
Strains X genotypes	14 23803*	23069**	.105**	.262**
Strains X general	4 27904*	38025*	.068	.298*
Strains X heterosis	10 22163*	17027**	.120**	.247**
Strains X average	1 6263	26956*	.004	.027
Strains X line	4 20118	15886*	.135*	.298*
Strains X specific	5 26978*	16055*	.131*	.250*
Error	86 9420	5324	.041	.084

*significant at the 5% level of probability.

**significant at the 1% level of probability.

logio transformed acetylene reduction assay only. The additive effects which were present in the within-strain analyses may be cancelled in a factorial analysis (Comstock, R.E., pers. comm.). Average heterosis was highly significant for all analyses while the significance of line and specific heterosis was character and scale dependent.

Strain by genotype interactions were found to be significant in this experiment. Strain X heterosis and strain X specific heterosis interactions were significant for all analyses while there appeared to be little strain X average heterosis interaction.

A limitation to this analysis is the limited number of degrees of freedom for the interaction terms. Since strains have been considered to be an environmental component in this study, the strains and blocks were confounded and analysed as a four block experiment so that direct comparisons could be made with Experiment II (Table 28). The overall diallel analyses of variance for the original and logio transformed characters are presented in Table 38. Genotypes were found to be significantly different for both scales as were all components with the exception of specific heterosis for the logio transformed acetylene reduction assay. All interaction terms were significant with the exception of blocks/strains X general which was non-significant for the logio transformed dry weight accumulation. The significance of the interaction terms suggested that genetic studies for total symbiotic nitrogen

Table 38. Mean squares from the overall diallel analysis of variance for Experiment III.

Source	Original			Log10		
	Dry weight df (mg/pl)	Acetylene reduced (nM/pl/hr)	weight reduced (mg/pl)	Dry Acetylene reduced (nM/pl/hr)	Acetylene reduced (nM/pl/hr)	Acetylene reduced (nM/pl/hr)
Blocks/strains (B/s)	3	44024**	1793**	.231**	3.385**	
Genotypes	14	47406**	156**	.178**	.227**	
General	4	21800**	128**	.079*	.219**	
Heterosis	10	57648**	168**	.217**	.230**	
Average	1	273173**	893**	1.169**	1.279**	
Line	4	36721**	106*	.147**	.156**	
Specific	5	31286**	76*	.083*	.080	
B/s x genotypes	42	20196**	143**	.084**	.201**	
B/s x general	12	16336*	203**	.047	.193**	
B/s x heterosis	30	21768**	120**	.098**	.204**	
B/s x average	3	15280*	113**	.159**	.274**	
B/s x line	12	18635**	124**	.169**	.234**	
B/s x specific	15	25491**	117**	.078**	.165**	
Error	58	5075	30	.026	.042	

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

fixed should be tested with different strains and in different environments if the experimental results are to be reliable.

The error mean squares for dry weight accumulation and the acetylene reduction assay in Experiment III (Table 38) have been reduced by approximately one-half those of Experiment II (Table 28). This reduction is most likely due to the modification of the bottle jar design (Figure 3) which reduced within-cell variation. Once the variation due to bottle jar design was removed, the \log_{10} transformation was not necessary as the original analyses were similar to the \log_{10} analyses. These data suggest that further investigation of the design of bottle jars or sand size would be beneficial.

The estimates of the variety constants and G.C.A. effects for the original and \log_{10} transformed characters from all analyses are presented in Tables 39 and 40, respectively. The estimates of the G.C.A. effects for the \log_{10} transformed characters for strain B (Table 39) were found to be similar to the results of Experiment I. Line Ds1 was found to have the largest G.C.A. effect for both characters while Ds4 showed the largest negative G.C.A. effect. Line Ds2 showed the largest heterosis constant for dry weight but Ds1 had the larger constant for the acetylene reduction assay. The G.C.A. effects for strain A (Table 39) were not similar to those reported for Experiment II. As earlier discussed the nodules of the plants of strain A were

Table 39. Estimates of G.C.A. effects and variety constants for the original characters for Experiment III.

	Rhizobium Strain A	Rhizobium Strain B	Overall
	Dry Acetylene	Dry Acetylene	Acetylene reduced
	weight reduced (mg/pl)	weight reduced (mg/pl)	weight reduced (mg/pl)
<u>Griining's G.C.A. effects</u>			
$g(Ds1)$	-8	-1.0	1.5
$g(Ds2)$	-20	-2.6	-7
$g(Ds3)$	-27	-1.6	28
$g(Ds4)$	-1	.7	-42
$g(Ds5)$	55	4.4	6
$se[g(1)]$	1.3	.6	1.1
$se[g(1)-g(j)]$	20	1.0	18
$se[g(1)-g(j)]$			1.8
<u>Gardner's variety constants</u>			
mean	187	13.8	187
$v(Ds1)$	-48	-3.2	18
$v(Ds2)$	36	-1.8	36
$v(Ds3)$	-20	-1.2	-10
$v(Ds4)$	-7	2.4	-26
$v(Ds5)$	38	3.7	-18
			-0.7
			10
			1.5

Table 40. Estimates of G.C.A. effects and variety constants for the log₁₀ transformed characters for Experiment III.

	Rhizobium Strain A		Rhizobium Strain B		Overall			
	Dry weight (mg/pl)	Acetylene reduced (mg/pl)						
<u>Grieffing's G.C.A. effects</u>								
$g(Ds1)$	-0.006	0.009	-0.026	0.077	-0.010	0.043		
$g(Ds2)$	-0.032	-0.095	-0.006	0.059	-0.013	-0.018		
$g(Ds3)$	-0.052	-0.051	-0.025	-0.008	-0.014	-0.029		
$g(Ds4)$	-0.012	-0.047	-0.071	-0.117	-0.042	-0.082		
$g(Ds5)$	-0.103	-0.184	-0.015	-0.011	-0.059	-0.087		
$se[g(1)]$	0.028	0.036	0.026	0.032	0.019	0.024		
$se[g(1)-g(j)]$	0.044	0.058	0.042	0.050	0.030	0.038		
<u>Gardner's variety constants</u>								
mean	2.182	0.624	2.217	1.036	2.200	0.830		
$v(Ds1)$	-0.137	-0.188	-0.053	0.163	-0.042	-0.003		
$v(Ds2)$	-0.102	0.026	-0.118	0.162	-0.110	-0.094		
$v(Ds3)$	-0.006	-0.090	-0.107	-0.217	-0.051	-0.063		
$v(Ds4)$	-0.067	-0.148	-0.033	-0.113	-0.050	-0.131		
$v(Ds5)$	-0.096	0.221	-0.031	-0.015	-0.032	-0.103		

in a state of senescence and this physiological state may have affected the results for the acetylene reduction analysis. However the G.C.A. effects for dry weight accumulation were similar, therefore nodule senescence may not explain the divergence from the G.C.A. effects of Experiment II. Line Ds5 presented the largest G.C.A. effect for strain A. Its crosses had larger means than the other crosses (Table 32). Line Ds5 and its crosses may have been resistant to nodule senescence or responded more quickly to clipping of the juvenile plants. Line Ds5 also has the largest G.C.A. effect for both characters for the overall analysis (Table 39).

The estimates of the heterosis constants and the S.C.A. effects derived from the original and the log₁₀ characters in the various analyses are presented in Tables 41 to 44, respectively.

d. Combined analysis.

The preceding experiments were conducted with the same diallel cross but differed in environmental and strain effects. Since strains have been initially considered as an environmental effect, it would be pertinent to pool the above experiments for a combined analysis. Blocks 1, 2 and 3 of Experiment I have been deleted from the combined analysis because subsamples within cells were not available for block 1 and the acetylene reduction assay was not recorded for blocks 2 and 3. The remaining nine blocks, six inoculated

Table 41. Estimates of Gardner's average, line and specific heterosis constants for the original characters for Experiment III.

Heterosis constant (mg/pl)	Rhizobium Strain A		Rhizobium Strain B		Overall		
	Dry weight reduced (nM/pl/hr)	Acetylene reduced (nM/pl/hr)	Dry weight reduced (mg/pl)	Acetylene reduced (mg/pl)	Dry weight reduced (nM/pl/hr)	Acetylene reduced (nM/pl/hr)	
	h(ave)	8.6	2.4	11.7	9.2	10.1	5.8
h(Ds1)	38	1.5	1.3	0.8	2.6	1.2	
h(Ds2)	-88	-4.0	-5.7	1.6	7.3	-1.2	
h(Ds3)	-41	-2.3	7.8	5.5	19	1.6	
h(Ds4)	7	-1.1	-6.7	-7.2	-30	-4.2	
h(Ds5)	85	5.9	3.4	-0.8	59	2.5	
h(Ds1, Ds2)	33	1.8	-2.7	-4.5	3	-1.4	
h(Ds1, Ds3)	73	3.1	94	5.7	84	4.4	
h(Ds1, Ds4)	-79	-4.0	-64	-1.6	-71	-2.8	
h(Ds1, Ds5)	-27	-1.0	-3	0.3	-15	-0.3	
h(Ds2, Ds3)	71	-0.4	14	-1.0	-28	-0.7	
h(Ds2, Ds4)	96	3.0	-23	-1.2	36	0.9	
h(Ds2, Ds5)	-58	-4.5	36	6.7	-11	1.1	
h(Ds3, Ds4)	-52	-3.6	6	2.5	-23	-0.5	
h(Ds3, Ds5)	49	0.9	-114	-7.3	-32	-3.6	
h(Ds4, Ds5)	36	4.5	81	0.3	58	2.4	

Table 42. Estimates of Gardner's average, line and specific heterosis constants for the Log₁₀ transformed characters for Experiment III.

	Rhizobium Strain A			Rhizobium Strain B			Overall		
	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	reduced weight (mg/pl)	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	reduced weight (mg/pl)	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	reduced weight (mg/pl)
h(ave)	.198	.193	.221	.266	.209	.230			
h(Ds1)	.145	.241	-.002	-.035	.072	.103			
h(Ds2)	-.194	-.253	-.125	-.050	-.160	-.151			
h(Ds3)	-.128	-.224	.183	.235	.027	.006			
h(Ds4)	.050	.064	-.128	-.142	-.039	-.039			
h(Ds5)	.128	.172	.071	-.008	.100	.082			
h(Ds1, Ds2)	.083	.050	.006	-.049	.045	.000			
h(Ds1, Ds3)	.151	.197	.097	.066	.124	.131			
h(Ds1, Ds4)	-.158	-.214	-.084	.000	-.121	-.107			
h(Ds1, Ds5)	-.076	-.032	-.020	-.016	-.048	-.024			
h(Ds2, Ds3)	-.234	-.211	.039	-.006	-.097	-.109			
h(Ds2, Ds4)	.202	.310	-.108	-.097	.047	.107			
h(Ds2, Ds5)	.051	-.148	.063	.152	.006	.002			
h(Ds3, Ds4)	-.044	-.131	.049	.087	.003	.022			
h(Ds3, Ds5)	.127	.146	-.186	-.146	-.029	.000			
h(Ds4, Ds5)	.000	.035	.143	.011	.071	.023			

Table 43. Estimates of Griffing's S.C.A. effects for the original characters for Experiment III.

S.C.A. effect	Rhizobium Strain A		Rhizobium Strain B		Overall	
	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)
$s(Ds1)$	-89	-2.9	-89	-6.8	-89	-4.9
$s(Ds2)$	1.9	1.8	-29	-7.6	-5	-2.9
$s(Ds3)$	-2.2	.4	-144	-10.9	-83	-5.3
$s(Ds4)$	-6.3	-.6	-20	0.0	-41	-.3
$s(Ds5)$	-130	-6.6	-107	-5.5	-118	-6.0
$se[s(1)]$	26	1.3	23	2.3	1.7	1.3
$se[s(1)-s(j)]$	35	1.7	31	3.1	2.3	1.8
$s(Ds1, Ds2)$	32	1.2	-13	-0.1	10	0.6
$s(Ds1, Ds3)$	100	3.4	185	12.4	143	7.9
$s(Ds1, Ds4)$	-25	-3.0	-56	-2.1	-41	-2.5
$s(Ds1, Ds5)$	72	4.1	62	3.4	67	3.7
$s(Ds2, Ds3)$	-116	-3.2	64	6.2	-26	1.5
$s(Ds2, Ds4)$	78	0.9	-55	-1.3	11	-0.2
$s(Ds2, Ds5)$	-31	-2.6	62	10.3	15	3.9
$s(Ds3, Ds4)$	-43	-4.8	51	4.7	4	-0.1
$s(Ds3, Ds5)$	103	3.7	-11	-1.5	46	1.1
$s(Ds4, Ds5)$	116	8.0	100	-1.2	108	3.4
$se[s(1, j)]$	33	1.7	29	2.9	22	1.7
$se[s(1, j)+s(i, k)]$	49	2.5	44	4.4	33	2.5
$se[s(1, j)-s(k, l)]$	45	2.3	40	4.0	30	2.3

Table 44. Estimates of Griffing's S.C.A. effects for the \log_{10} transformed characters for Experiment III.

S.C.A. effect	Rhizobium Strain A			Rhizobium Strain B			Overall		
	Dry weight (mg/pl)	Acetylene reduced (nM/pl/hr)							
$s(Ds1)$	-.256	-.335	-.146	-.147	-.201	-.241			
$s(Ds2)$.035	.088	-.041	-.134	-.003	-.023			
$s(Ds3)$.022	.063	-.304	-.379	-.163	-.158			
$s(Ds4)$	-.175	-.183	-.038	-.056	-.106	-.120			
$s(Ds5)$	-.242	-.276	-.209	-.171	-.225	-.223			
$se[s(i)]$.057	.074	.054	.065	.039	.049			
$se[s(i)-s(j)]$.077	.100	.072	.087	.053	.066			
$s(Ds1, Ds2)$.121	.107	.008	-.009	.064	.049			
$s(Ds1, Ds3)$.226	.271	.275	.269	.250	.027			
$s(Ds1, Ds4)$.019	.025	-.084	-.013	-.033	.006			
$s(Ds1, Ds5)$.146	.268	-.094	.048	.120	.158			
$s(Ds2, Ds3)$	-.353	-.419	.146	.188	-.103	-.115			
$s(Ds2, Ds4)$.185	.266	-.178	-.118	.003	.074			
$s(Ds2, Ds5)$	-.023	-.130	.106	.207	.041	.039			
$s(Ds3, Ds4)$	-.023	-.158	.154	.229	.066	.035			
$s(Ds3, Ds5)$.193	.181	.034	.072	.113	.126			
$s(Ds4, Ds5)$.168	.234	.184	.014	.176	.124			
$se[s(i,j)]$.072	.094	.068	.082	.050	.063			
$se[s(i,j)-s(i,k)]$.109	.141	.102	.124	.075	.094			
$se[s(i,j)-s(k,l)]$.099	.129	.093	.113	.068	.086			

with strain A and three with strain B, were pooled such that block and strain effects were confounded.

The original and \log_{10} means confirmed that the mean of the F_1 genotypes was much larger than the means of their inbred parents (Table 45). The mean heterosis for all characters appeared to be approximately 1.8.

The diallel analyses of variance for Gardner's Analysis II of the original and \log_{10} transformed characters of the combined analysis are presented in Table 46 and for comparison Gardner's Analysis III is provided in Table 47. Both analyses are compatible; however, the general mean square was larger for Analysis III than for Analysis II suggesting that heritable effects are given more weighting by the Analysis III. Gardner and Eberhart (1966) report that Analysis III may be superior when average heterosis is highly significant as in the present data; however, either analysis is applicable.

Significant differences among genotypes were found for all characters except the original acetylene reduction assay. These results were not unexpected since the acetylene reduction assay values in the earlier experiments were affected by the difficulty of removing the root intact from the bottle jar wick. Highly significant heterosis and average heterosis effects were also recorded. All interaction effects for the \log_{10} transformed characters were significant with the exception of block/strains \times average heterosis for the \log_{10} transformed acetylene

Table 45. Means and standard error of the difference between two means for the combined diallel analysis.

Genotype	Original		Log ₁₀	
	Dry weight (mg/pl)	Acetylene reduced (nM/pl/hr)	Dry weight (mg/pl)	Acetylene reduced (nM/pl/hr)
Ds1	118	47	1.953	1.267
Ds2	137	25	1.986	1.214
Ds3	126	49	2.001	1.309
Ds4	126	44	1.947	1.122
Ds5	135	37	2.042	1.343
Ds1 x Ds2	181	55	2.162	1.440
Ds1 x Ds3	262	77	2.288	1.556
Ds1 x Ds4	204	72	2.182	1.419
Ds1 x Ds5	248	96	2.273	1.636
Ds2 x Ds3	160	55	2.077	1.352
Ds2 x Ds4	193	59	2.197	1.450
Ds2 x Ds5	218	101	2.264	1.622
Ds3 x Ds4	228	97	2.298	1.507
Ds3 x Ds5	262	86	2.353	1.619
Ds4 x Ds5	271	97	2.270	1.457
se of d.	42	35	0.093	0.117

Table 46. Mean squares from Gardner's Analysis II for the combined diallel analysis.

Source	Original		Log ₁₀	
	Dry weight df (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight reduced (mg/pl)	Acetylene reduced (nm/pl/hr)
Blocks/Strains (B/s)	8	237264**	267858**	1.891**
Genotypes	14	55338**	10882*	.352**
General	4	19363**	5999	.124*
Heterosis	10	69730**	12843**	.443**
Average	1	535941**	91299**	3.761**
Line	4	28983**	5229	.071
Specific	5	9092	3236	.077
B/s x genotypes	112	16338**	6722	.076**
B/s x general	32	12392	4133	.069**
B/s x heterosis	80	17915**	7754	.079**
B/s x average	8	22822**	23354**	.112*
B/s x line	32	14566*	6269	.080**
B/s x specific	40	19615**	5834	.071**
Error	129	8099	5604	.039
				.062

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

Table 47. Mean squares from Gardner's Analysis III for the combined diallel analysis.

Source	df	Original		Ratio	
		Dry weight (mg/pl)	Acetylene reduced (nM/pl/hr)	Dry weight (mg/pl)	Acetylene reduced (nM/pl/hr)
Blocks/Strains (B/s)	8	237264**	267858**	1.891**	8.486**
Genotypes	14	55338**	10882*	.352**	.428**
Varieties	4	1068	1652	.028	.135
Varieties vs crosses	1	535941**	91299**	3.761**	3.900**
Crosses	9	26057**	6049	.118**	.173**
General	4	47263**	9578	.167**	.238**
Specific	5	9092	3236	.077	.121
B/s x genotypes	112	16338**	6722	.076**	.152**
B/s x varieties	32	5308	2135	.061*	.152**
B/s x varieties vs crosses	8	22822**	23354**	.112**	.115
B/s x crosses	72	20520**	6913	.078**	.157**
B/s x general	32	21650**	8260	.087**	.175**
B/s x specific	40	19615**	5834	.072**	.141**
Error	129	8099	5604	.039	.062

*significant at the 5% level of probability.
**significant at the 1% level of probability.

reduction assay. These results suggested that genotype x environment interactions were of significant importance in the study of total symbiotic nitrogen fixed. Significant general effects for the log₁₀ transformed characters for both diallel analyses suggested that heritable genetic variation was present.

The combining ability analysis for G.C.A. effects and variety constants are presented in Table 48. As indicated by the genotype means, line Ds5 was shown to have the largest G.C.A. effect and variety constant for the log₁₀ transformed characters. Lines Ds2 and Ds4, on the other hand, had negative G.C.A. effects. Heterosis constants and S.C.A. effects for the two scales are presented in Tables 49 and 50, respectively. The average heterosis constant was much larger than the other heterosis constants suggesting that the major portion of the genetic variance is due to F₁ heterosis (Table 49). On the other hand, all inbred lines exhibited negative S.C.A. effects suggesting inbreeding depression (Table 50). Further studies would have to be undertaken to determine whether the large average heterosis effects were due to F₁ heterosis or inbreeding depression exhibited by the inbred lines when grown in a nitrogen limiting environment.

The combined analysis indicated that total symbiotic nitrogen fixed had both heritable and non-heritable components of the genetic variance. The larger proportion of variance was contributed by the non-heritable portion

Table 48. Estimates of G.C.A. effects and variety constants for the combined analysis.

	Original	Logio		
	Dry Acetylene weight (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight reduced (mg/pl)	Acetylene (nm/pl/hr)
<u>Griffing's G.C.A. effects</u>				
$g(Ds1)$	-2.3	-0.8	-0.015	.008
$g(Ds2)$	-1.7	-11.1	-0.035	-.033
$g(Ds3)$	3	2.1	0.015	.018
$g(Ds4)$	0	2.1	-0.011	-.064
$g(Ds5)$	1.7	7.6	0.647	.071
$se[g(i)]$	7	6.0	0.016	.020
$se[g(i)-g(j)]$	11	9.4	0.025	.031
<u>Gardner's variety constants</u>				
mean	12.8	40.9	1.986	1.251
$v(Ds1)$	-1.1	6.1	-0.033	.016
$v(Ds2)$	9	-14.9	0.00	-.037
$v(Ds3)$	-2	8.7	0.016	.058
$v(Ds4)$	-2	3.9	-0.039	-.129
$v(Ds5)$	6	-3.9	0.056	.092

Table 49. Estimates of Gardner's average, line and specific heterosis constants for the combined analysis.

Heterosis constant	Original		Log10	
	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)
$h(\text{ave})$	95	39.0	.250	.255
$h(Ds1)$	6	-9.0	.003	.001
$h(Ds2)$	-51	-8.4	-.082	-.035
$h(Ds3)$	8	-5.4	.016	-.025
$h(Ds4)$	3	0.4	.020	.001
$h(Ds5)$	33	22.3	.044	.058
$h(Ds1, Ds2)$	3	-2.6	.210	-.022
$h(Ds1, Ds3)$	31	4.6	.415	.038
$h(Ds1, Ds4)$	-22	-3.5	-.041	-.033
$h(Ds1, Ds5)$	-12	1.5	-.022	.017
$h(Ds2, Ds3)$	-24	-7.6	-.101	-.104
$h(Ds2, Ds4)$	14	-6.9	.042	.061
$h(Ds2, Ds5)$	6	17.1	.038	.066
$h(Ds3, Ds4)$	-3	16.0	.038	.060
$h(Ds3, Ds5)$	-4	-13.0	.022	.006
$h(Ds4, Ds5)$	10	-5.6	-.039	-.089

Table 50. Estimates of Griffing's S.C.A. effects for the combined analysis.

S.C.A. effect	original		log ₁₀	
	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)
s(Ds1)	-68	-18.3	-.169	-.172
s(Ds2)	-20	-18.8	-.097	-.140
s(Ds3)	-70	-21.4	-.181	-.148
s(Ds4)	-66	-26.3	-.184	-.170
s(Ds5)	-91	-45.1	-.204	-.219
se[s(1)]	15	12.2	.032	.040
se[s(i)-s(j)]	20	16.3	.043	.054
s(Ds1, Ds2)	9	.5	.059	.045
s(Ds1, Ds3)	71	9.4	.135	.110
s(Ds1, Ds4)	15	4.6	.055	.053
s(Ds1, Ds5)	42	22.2	.089	.136
s(Ds2, Ds3)	-16	-2.4	-.056	-.053
s(Ds2, Ds4)	19	1.5	.090	.125
s(Ds2, Ds5)	27	38.0	.100	.163
s(Ds3, Ds4)	35	26.2	.142	.131
s(Ds3, Ds5)	51	9.7	.139	.109
s(Ds4, Ds5)	62	20.4	.081	.031
se[s(1, j)]	19	15.4	.041	.051
se[s(i, j)-s(i, k)]	28	23.1	.061	.077
se[s(i, j)-s(k, l)]	25	21.1	.056	.070

because of the expression of average heterosis. Crosses Ds4 x Ds5, Ds1 x Ds3 and/or Ds2 x Ds5 would be of significant interest for an intensive study since these crosses exhibited high mean yield and specific combining ability effects for at least one of the two characters.

e. Correlation between dry weight accumulation and the acetylene reduction assay.

A secondary objective of this study was to relate dry weight accumulation and the acetylene reduction assay as quantitative measure for total symbiotic nitrogen fixation.

The correlation coefficients for these characters for the diallel experiments are presented in Table 51. The correlation coefficients were found to be highly significant with the exception of the unadjusted overall correlation.

These data suggest that the acetylene reduction assay is compatible with dry weight accumulation for juvenile plants grown in bottle jars. A similarly large correlation ($r = .89$) was found by Lawn et al. (1974) for total nodule activity and shoot dry weight for ~~selected~~ genotype means.

It may be concluded that the acetylene reduction assay would be suitable as a plant breeding tool for screening of juvenile genotypes into various classes for total nitrogen fixed. Since the acetylene reduction assay does not require nitrogen-free growth conditions this assay would be effective and reliable for screening plants grown under field conditions.

Table 51. Unadjusted and block adjusted correlation coefficients between dry weight (mg/pl) and acetylene reduced (nM/pl/hr) for the diallel experiments.

Experiment	Unadjusted "r"	Block Adjusted "r"*
I	.92	.86
II	.92	.83
III	.76	.83
Combined	.43	.85

*all block means in each character are adjusted to the grand mean for that character.

2. Cross between two inbred lines.

A second approach to the study of quantitative genetic variance is provided by the analysis of populations produced by a cross between two inbred lines. Through the analysis of first degree statistics, the cross between two inbred lines will provide information about epistatic interactions which cannot be obtained from a diallel cross. In this study of the inheritance of symbiotic nitrogen fixation the cross between two inbred lines has been employed as an alternate quantitative genetic design for the establishment and the partition of the genetic variance and to investigate the influence of epistasis in the expression of heterosis. Three experiments were conducted in this section.

a. Experiment IV.

Seven blocks were grown in this experiment, three blocks were deleted due to their uniqueness. Blocks 7 and 8 were grown in large bottom (2l) bottle jars while the remainder had small bottles (600ml). Block 5 was missing many of the B₁ population plants and was not considered to be complete. Because of poor germination within the B₁ population, six B₁ plants from the deleted blocks were fitted into the remaining blocks so as not to appreciably change the within block mean.

The original and log₁₀ means revealed F₁ heterosis with the F₂ mean falling very near the midparent value suggesting the random segregation of alleles (Table 52). The backcross

Table 52. Means, standard deviations and coefficients of variation for Experiment IV.

Genotype n	original	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	$\log_{10}(X+1)$
		original	$\log_{10}(X)$	
P ₁ (D ₈₁) 22	42.5±17.5(41)	1.60±.17(11)	45.1±26.1(58)	1.67±.04(2)
P ₂ (D ₈₁) 28	17.8± 7.6(43)	1.22±.17(14)	10.9± 7.9(72)	1.20±.19(16)
F ₁	16 58.5±33.3(57)	1.82±.19(10)	52.0±27.7(53)	1.72±.20(12)
F ₂	123 29.8±20.1(67)	1.38±.30(22)	30.6±29.4(96)	1.45±.32(22)
B ₁	22 16.3± 9.8(60)	1.12±.33(29)	15.3±10.3(67)	1.29±.20(16)
B ₂	37 23.1±12.4(54)	1.30±.25(19)	18.5±17.6(95)	1.32±.25(19)

means do not appear to fit any pattern.

The mean squares of the analysis of variance of population means for the original and log₁₀ transformed data are presented in Table 53. The analysis of variance of the population means indicated the blocks were not significantly different while populations were highly significantly different. Additive, dominance and epistatic effects were also significant with dominance by dominance interactions accounting for the major epistatic variation. The test for non-additivity revealed that the log₁₀ scale was the most appropriate due to the effects of the bottle jar design (discussed earlier in the diallel section).

b. Experiment V.

The means of the six populations created from a cross between Ds2 and Dsp (Table 1) revealed that the F₁ mean was larger than the other means indicating the presence of F₁ heterosis in this population (Table 54). The backcross population means tended to exceed the F₁ mean which is unexpected for an additive model. These results suggest epistatic interactions. The mean for Dsp was found to be much less than expected. Observing the F₂ population distribution for the original and log₁₀ dry weight accumulation it was noted that the log₁₀ data was skewed toward the upper end of the distribution (Figure 11). These distributions may be interpreted as a confounding of two distributions (Figure 12). The phenomenon producing the low

Table 53. Mean squares from the genetic analysis of variance of population means for Experiment IV.

Source	df	Dry weight (mg/pl)		Acetylene reduced (nM/pl/hr)	
		original	log ₁₀ (X)	original	log ₁₀ (X+1)
Blocks	4	155.1	.007	104.8	.003
Populations	5	1370.5**	.260**	1467.8**	.227**
Additive	1	1033.0**	.186**	2339.4*	.434**
Dominance	1	1451.3**	.149**	1034.3*	.151*
Epistasis	3	1456.0**	.3222**	1322.0**	.183**
Add-Add	1	330.9	.087*	600.0	.066
Add-Dom	1	731.9*	.2622**	826.8*	.131*
Dom-Dom	1	3305.2**	.616**	2539.6**	.353**
Error	20	126.2	.014	187.6	.018
Non-additivity	1	1418.5**	.067	1267.5**	.002
Residual	19	58.2	.012	131.0	.019

*significant at the 5% level of probability.

**significant at the 1% level of probability.

Table 54. Means, standard deviations and coefficients of variation for Experiment V.

Genotype n	original	Dry weight (mg/pl)	Acetylene reduced (nm/pl/hr)	$\log_{10}(X+1)$	$\log_{10}(X+1)$
		original	original		
P ₁ (Ds2)	15 108±34(31)	2.01±.15(8)	53.6±21.1(39)	1.74±.17(10)	
P ₂ (Dsp)	14 8± 7(88)	0.72±.34(47)	2.0± 2.0(99)	0.92±.08(9)	
F ₁	15 119±32(27)	2.06±.14(7)	71.1±19.6(28)	1.87±.11(6)	
F ₂	88 112±57(51)	1.96±.35(18)	64.0±41.2(64)	1.76±.29(16)	
B ₁	43 144±51(35)	2.12±.18(8)	86.1±35.1(41)	1.92±.19(10)	
B ₂	31 142±73(51)	2.04±.40(20)	78.1±45.0(58)	1.83±.32(17)	

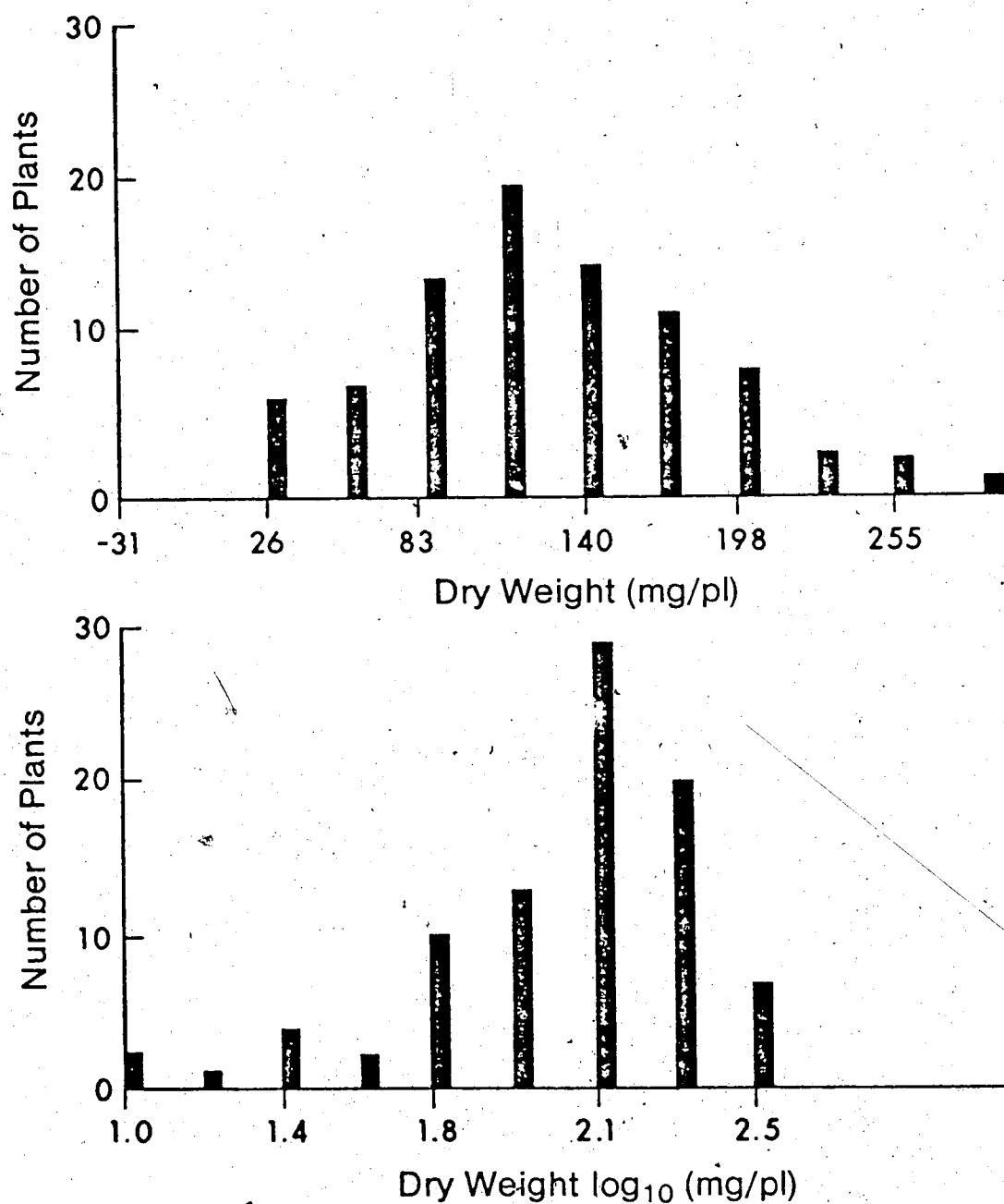


Figure 11. Normal and \log_{10} distributions of the F_2 dry weight values for Experiment V.

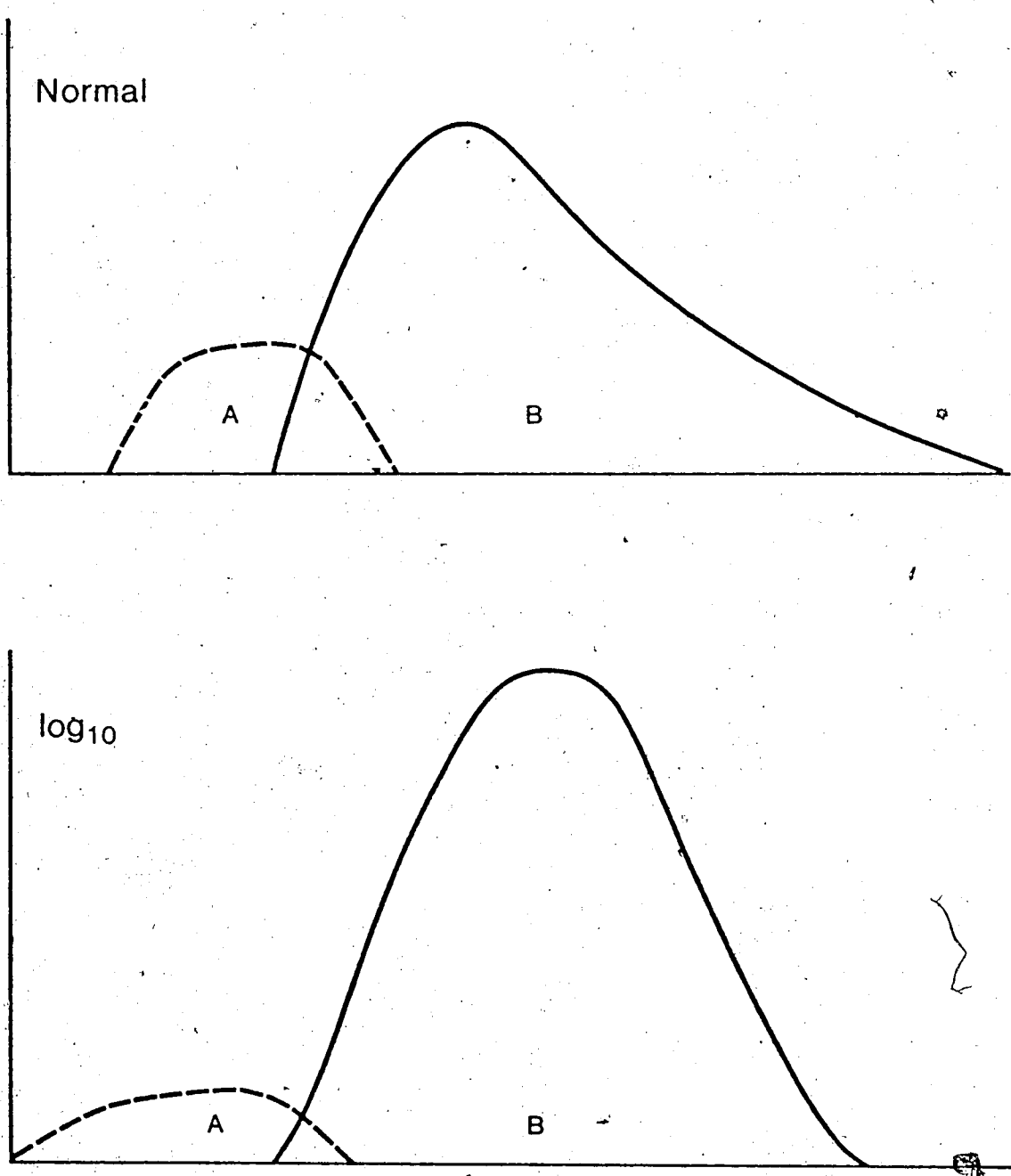


Figure 12. Normal and \log_{10} distributions suggesting the distributions of segregating factors A and B for Experiment V.

Dsp, F₂ and B₂ scores appeared to be a gene segregating for pH effects since the Dsp line grew normally at pH 6.2±.2. Further investigation would be necessary to determine the exact nature of the low Dsp scores.

The six populations of a cross between Ds2 and Dsp when analysed for the analysis of variance of population means showed highly significant population differences (Table 55). Additive, dominance and epistatic effects were highly significant. Dominance X dominance and additive X dominance interactions contributed to epistatic variation. These results were compatible with those of Experiment IV.

c. Experiment VI.

The combined analysis of the diallel suggested that three crosses showed high mean yield and specific combining ability for the characters studied. Cross Ds2 x Ds5 was the cross which had the highest mean yield for the acetylene reduction assay so in Experiment VI the six populations of the Ds2 x Ds5 cross were created for intensive study.

The correlation coefficients for the characters studied are presented in Table 56. The estimates of N₂ fixation were found to be highly correlated among themselves and partially correlated with the leaf characters. Applying the principal components transformation to the fixation estimate characters yielded one factor (Table 57). The estimate variables were highly correlated with the first factor indicating that all estimates measure the principal axis of

Table 55. Mean squares from a genetic analysis of variance of population means for Experiment V.

Source	df	Dry weight (mg/pl)		Acetylene reduced (nM/pl/hr)	
		original	log ₁₀ (X)	original	log ₁₀ (X+1)
Blocks	4	1307*	.008	989	.017
Populations	5	12810**	1.450**	4690**	.708**
Additive	1	20664**	3.471**	5509**	1.482**
Dominance	1	21831**	2.295**	10567**	1.384**
Epistasis	3	7186**	.494**	2455**	.226**
Add-Add	1	2945*	.054	1424	.054
Add-Dom	1	4892**	.651**	1010	.222**
Dom-Dom	1	13721**	.776**	4926**	.402**
Error	20	384	.026	414	.015
Non-additivity	1	1080	.019	1246	.026
Residual	19	384	.026	372	.015

*significant at the 5% level of probability.

**significant at the 1% level of probability.

Table 16. Correlations for Experiment VI.

Character	Correlations							
	1	2	3	4	5	6	7	8
Acetylene reduced	1							
Shoot fresh weight	2	0.83	1.00					
Root fresh weight	3	0.73	0.82	1.00				
Shoot dry weight	4	0.84	0.98	0.83	1.00			
Cotyledon length	5	0.19	0.29	0.36	0.28	1.00		
Cotyledon width	6	0.20	0.27	0.32	0.26	0.75	1.00	
Cotyledon area	7	0.21	0.31	0.37	0.29	0.93	0.94	1.00
Juvenile leaf width	8	0.19	0.28	0.37	0.27	0.56	0.45	0.54

Table 57. Eigenvectors for factors from Experiment VI.

Character	Factor			
	1	2	3	4
Acetylene reduced	0.90	-.35	0.24	0.00
Top fresh weight	0.97	-.01	-.22	0.11
Root fresh weight	0.90	0.38	0.20	0.00
Top dry weight	0.97	-.02	-.20	-.11
Eigenvalue	3.51	0.27	0.19	0.02
% variation explained	88	7	5	0

variation; namely N_2 fixation. Shoot fresh and dry weight appeared to be the better estimates of N_2 fixation since they are most highly correlated with the principal axis. Since scores for the principal axis can be defined with a principal components transformation, the "factor" scores were calculated for the principal axis from the eigenvectors. The new derived 'character' has been termed the "Nitrogen fixation" factor and differences among population means are of importance rather than the amount of N_2 fixed. The "Nitrogen fixation" factor has been given a distribution of 1/3 the mean and standard deviation of the acetylene reduction assay since nitrogenase is assumed to theoretically reduce 1/3 the amount of N_2 than acetylene (Burris 1974). The "Nitrogen fixation" factor was added to the data set and treated as another character for the remaining analyses in Experiment VI.

The means and standard deviations for the original and log₁₀ transformed data are presented in Tables 58 and 59, respectively. The F_1 mean was found to be greater than either parent for all scales and characters while the F_2 and backcross means were not significantly different from the parents. The $Ds2$ mean was found to be significantly smaller than the B_1 mean for the cotyledon characters (Table 58). Both the $Ds2$ and B_1 means were significantly less than the other population means suggesting a genetic basis to the difference. A similar pattern has also been shown for the juvenile leaf character.

Table 58. Means, standard deviations, and coefficients of variation for the original characters for Experiment VI.

	Population n	Acetylene reduced (nM/pl/hr)	Shoot fresh weight (mg/pl)	Root fresh weight (mg/pl)	Nitrogen fixation "factor"	Shoot dry weight (mg/pl)	Nitrogen fixation
P ₁ (Ds2)	58	65±37(57)ab ⁺	525±284(54)b	1.37±0.86(63)d	137±78(57)b	19±12(64)bc	
P ₂ (Ds5)	55	51±22(43)b	475±170(36)b	1.66±0.63(38)b-d	121±47(39)b	17±7(43)c	
F ₁	59	72±33(45)a	652±232(36)e	1.92±0.82(43)ab	173±64(37)a	24±10(41)a	
F ₂	229	59±30(52)b	537±219(41)b	1.73±0.85(49)bc	138±62(45)b	20±10(50)bc	
B ₁	108	58±30(52)b	522±243(47)b	1.46±0.82(55)c-d	138±70(51)b	19±11(57)bc	
B ₂	111	63±35(56)ab	574±266(46)b	2.06±1.14(55)a	149±73(49)b	22±12(55)ab	

⁺means followed by the same letter are not significantly different for the Student-Newman-Keul range test.

Table 58 (cont.). Means, standard deviations, and coefficients of variation for the original characters for Experiment VI.

Population n	Cotyledon			Juvenile leaf		
	Length (mm)	Width (mm)	Area (mm ²)	width (mm)	Area (mm ²)	
P ₁ (Ds2)	58	6.8±.5(7)c+	4.3±.5(12)c	29±5(17)c	5.7±0.6(11)d	
P ₂ (Ds5)	55	8.0±.7(9)a	5.0±.5(10)a	40±7(17)a	7.4±0.9(12)a	
F ₁	59	8.2±.6(7)a	5.0±.4(8)a	41±5(12)a	7.1±0.6(8)ab	
F ₂	229	8.0±.7(9)a	4.9±.5(10)a	39±7(18)a	7.0±0.7(10)b	
B ₁	108	7.4±.7(9)b	4.5±.5(11)b	34±6(18)b	6.1±0.9(15)c	
B ₂	111	8.1±.8(11)a	5.0±.6(12)a	41±5(22)a	7.3±1.3(18)a	

*means followed by the same letter are not significantly different for the Student-Newman-Keul range test.

Table 5g. Means, standard deviations, and coefficients of variation for the \log_{10} transformed characters for Experiment VI.

Population n	Acetylene reduced (nM/pl/hr)	Shoot		Root		Nitrogen fixation "factor" (mg/pl)
		fresh weight (mg/pl)	dry weight (g/pl)	fresh weight (mg/pl)	dry weight (g/pl)	
P ₁ (D82)	58	1.738±.273(10)ab†	2.663±.225(8)b	0.070±.245(350)c	1.068±.248(12)b	1.190±.040(34)b
P ₂ (D85)	55	1.666±.205(8)b	2.646±.174(7)b	0.182±.202(111)ab	1.046±.190(9)b	1.182±.251(21)b
F ₁	59	1.811±.212(7)a	2.782±.179(6)a	0.241±.196(81)a	1.202±.168(8)a	1.345±.212(16)a
F ₂	229	1.713±.234(9)b	2.693±.186(7)b	0.187±.211(113)a	1.086±.203(10)b	1.242±.236(19)b
B ₁	108	1.692±.275(10)b	2.673±.199(7)b	0.112±.231(206)bc	1.084±.223(11)b	1.203±.264(22)b
B ₂	111	1.729±.268(10)ab	2.707±.226(8)b	0.243±.261(107)a	1.111±.248(12)b	1.265±.292(23)ab

means followed by the same letter are not significantly different for the Student-Newman-Keul range test.

The analyses of variance of the population means for the original and the log₁₀ transformed characters are given in Tables 60 and 61, respectively. Populations and dominance effects were highly significant for all characters in both scales; however, additive effects were significant for root, fresh weight. Epistasis appeared to be absent for most characters. These data suggest a completely non-heritable model for the estimates of total symbiotic nitrogen fixation.

The analyses of variance of the population means for the cotyledon and juvenile leaf characters indicated that an additive-dominance model adequately explains the distribution of the means. These data suggest that large cotyledons and juvenile leaves appear to be conditioned by a dominant allele at a single locus or a block of loci segregating as a single unit. To determine whether the cotyledon and juvenile leaf size expressed pleiotropism the cotyledon area and juvenile leaf width characters were subdivided into groups. The crosstabulation of the groups indicated that plants with small cotyledons and large juvenile leaves and vice versa were found implying that the cotyledon and juvenile leaf sizes are controlled by different genes (Table 62).

Having defined the cotyledon area and juvenile leaf width groups, it would be of interest to determine if plants with larger cotyledons and juvenile leaves would fix more nitrogen. The means, standard deviations, and coefficients

Table 60. Mean squares for the original characters for Experiment VI.

Source	df	Acetylene reduced (nm/pl/hr)	Shoot fresh weight (mg/pl)	Root fresh weight (g/pl)	Nitrogen fixation "factor"	
					Shoot dry weight (mg/pl)	Nitrogen fixation "factor"
Populations	5	3245**	227926**	5.58**	18687**	455**
Additive	1	3067	14997	11.23**	2648	2
Dominance	1	6757**	878971**	10.21**	73462**	1688**
Epistasis	3	2133	81914	2.16*	5772	110
Add-Add	1	98	4936	0.11	8490	195
Add-Dom	1	3994	152147	6.16**	8683	473
Add-Dom	1	2305	88657	0.21	143	1
Error	614	999	55668	0.79	4364	111

*significant at the 5% level of probability.

**significant at the 1% level of probability.

Table 60 (cont). Mean squares for the original characters for Experiment VI.

Source	df	Cotyledonary leaf			Juvenile leaf		
		Length (mm)	Width (mm)	Area (mm ²)	Length (mm)	Width (mm)	Area (mm ²)
Populations	5	21.3**	6.4**	1668**	36.2**		
Additive	1	73.2**	26.4**	6349**	159.2**		
Dominance	1	29.6**	5.0**	1746**	15.4**		
Epistasis	3	1.2	0.3	81	2.3*		
Add-Add	1	1.4	0.1	73	0.8		
Add-Dom	1	0.4	0.4	92	3.2*		
Add-Dom	1	1.6	0.3	79	2.1		
Error	614	0.5	0.3	47	0.8		

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

Table 61. Mean squares for the log₁₀ transformed characters for Experiment VI.

Source	df	Acetylene reduced (nM/pl/hr)	Shoot fresh weight (mg/pl)	Root fresh weight (g/pl)	Shoot dry weight (mg/pl)	Nitrogen fixation "factor"
Populations	5	.155*	.143**	.376**	.178**	.239**
Additive	1	.072	.000	.983**	.002	.012
Dominance	1	.390*	.602**	.720**	.762**	.997**
Epi-stasis	3	.105	.036	.055	.042	.062
Add-Add	1	.000	.000	.006	.000	.002
Add-Dom	1	.132	.045	.170	.034	.111
Add-Dom	1	.182	.063	.001	.091	.074
Error	614	.061	.039	.051	.047	.066

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

Table 62. Numbers of plants for cotyledon area by juvenile leaf width crosstabulation for Experiment VI.

Cotyledon area	Juvenile leaf width			Row total
	< 6 mm	6-7 mm	> 7 mm	
< 32 mm ²	I I I	I 48 I	I 12 I	I 156
32-39 mm ²	I I I	I 57 I	I 21 I	I 130
39-42 mm ²	I I I	I 91 I	I 49 I	I 175
> 42 mm ²	I I I	I 56 I	I 96 I	I 162
Column total	193	252	178	623

of variation for the nitrogen fixation estimates are presented in Table 63. Plants with larger cotyledons and juvenile leaves fixed significantly more nitrogen. It was also noted that plants growing in fine sand also fixed more nitrogen than did plants growing in very fine sand (Table 63). Combining the above groupings with populations, a factorial analysis of variance was performed on the estimates of nitrogen fixation to determine whether population interactions existed. These analyses of the original and log₁₀ transformed data are presented in Tables 64 and 65, respectively. All main effects were highly significant for all estimates except root fresh weight for populations when adjusted for the other main effects. The only interaction of significance was population by sand size indicating that changing sand size in an experiment may affect the estimates of nitrogen fixation. Since cotyledon area and juvenile leaf width groups show significant differences for the total fixation estimates, it would be of interest to treat cotyledon area and juvenile leaf width as covariates to adjust the estimates for effects due to cotyledon or juvenile leaf size (juvenile plant vigor) because these leaves are functional during the onset of nodulation. Because plants with larger cotyledons and juvenile leaves may fix more nitrogen (Mytton, 1973), this advantage should be removed from the analysis to understand the mechanisms of N₂ fixation. The analysis of covariance for the estimates for the original and log₁₀ transformed

Table 63. Means, standard deviations and coefficients of variation for plant groupings for Experiment VI.

Grouping	n	Acetylene reduced (nM / pl / hr)	Shoot fresh weight (mg / pl)	Root fresh weight (g / pl)	Shoot dry weight (mg / pl)	Nitrogen fixation "factor"
<u>Cotyledon area</u>						
< 32 mm ²	157	56±33(60) ^{b+}	473±259(35) ^c	1.37±0.80(38) ^c	1.22±73(60) ^c	17±11(66) ^c
32-39 mm ²	128	55±24(44) ^b	487±192(39) ^c	1.50±0.68(45) ^c	1.26±54(43) ^c	18±8(46) ^c
39-42 mm ²	175	62±32(53) ^b	563±221(39) ^b	1.80±0.86(48) ^b	1.46±64(44) ^b	21±10(48) ^b
> 42 mm ²	159	69±34(49) ^a	644±235(36) ^a	2.17±1.02(47) ^a	1.68±65(39) ^a	25±11(44) ^a
<u>Juvenile leaf width</u>						
< 6 mm	193	55±31(56) ^{b+}	474±227(48) ^c	1.38±0.74(54) ^c	1.23±65(53) ^c	17±10(58) ^c
6-7 mm	250	59±31(53) ^b	546±239(44) ^b	1.69±0.83(49) ^b	1.41±67(48) ^b	20±10(52) ^b
> 7 mm	176	69±32(47) ^a	624±226(36) ^a	2.15±1.00(47) ^a	1.63±63(39) ^a	24±10(43) ^a
<u>Sand size</u>						
Very fine(< 15mm)	313	56±27(47) ^{b+}	505±204(40) ^b	1.70±0.91(54) ^{ab}	1.32±60(45) ^b	19±10(50) ^b
Mixed 1/2 each	155	68±38(55) ^a	563±273(48) ^a	1.87±1.03(55) ^a	1.49±77(52) ^a	22±12(57) ^a
Fine(.25mm)	153	62±34(55) ^a	609±251(41) ^a	1.62±0.76(47) ^b	1.52±68(45) ^a	21±11(50) ^a

*means followed by the same letter are not significantly different for the Student-Newman-Keul range test.

Table 64. Mean squares from a factorial analysis of variance of the original characters for Experiment VI.

Source	df	Acetylene reduced (nM/pl/hr)	Shoot fresh weight (mg/pl)	Root weight (g/pl)	Nitrogen fixation factor (mg/pl)	Shoot dry weight (mg/pl)	Root dry weight (mg/pl)	Nitrogen fixation
Populations (P)	5	4997**	286721**	1.39	27477**	485**		
Cotyledon group (C)	3	4235**	463163**	7.62**	36089**	930**		
Juvenile leaf group (J)	2	4341**	395225**	6.37**	28700**	794**		
Sand size (S)	2	7932**	516164**	4.93**	23475**	589**		
P x C	14	753	35585	0.66	2538	75		
P x J	9	1042	69065	1.29*	5481	144		
P x S	10	1803*	131303**	1.30*	10174**	243**		
C x J	6	503	43294	0.71	3574	86		
C x S	6	1126	62469	0.91	3442	125		
J x S	4	639	37881	0.45	2473	62		
Error	556	885	45784	0.65	3678	92		

*significant at the 5% level of probability.

**significant at the 1% level of probability.

Table 65. Mean squares from a factorial analysis of variance of the log₁₀ transformed characters for Experiment VI.

Source	df	Acetylene reduced (nm/pl/hr)	Shoot fresh weight (mg/pl)	Root fresh weight (g/pl)	Shoot dry weight (mg/pl)	Nitrogen fixation "factor"
Populations (P)	5	0.257**	0.198**	0.043	0.283**	0.269**
Cotyledon group (C)	3	0.236**	0.401**	0.534**	0.483**	0.672**
Juvenile leaf group (J)	2	0.334**	0.336**	0.464**	0.387**	0.555**
Sand size (S)	2	0.238**	0.268**	0.197**	0.158*	0.157
P x C	14	0.083	0.032	0.052	0.038	0.066
P x J	9	0.065	0.049	0.078	0.054	0.084
P x S	10	0.125*	0.099**	0.101**	0.118**	0.156**
C x J	6	0.005	0.017	0.038	0.018	0.024
C x S	6	0.051	0.044	0.051	0.038	0.067
J x S	4	0.010	0.014	0.010	0.011	0.009
Error	556	0.054	0.031	0.041	0.039	0.054

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

data are presented in Tables 66 and 67, respectively. As expected the covariates were highly significant as were differences among the populations (see Table 66). However, dominance deviations were no longer significant whereas additive effects have become highly significant. These data suggest that the "efficiency" (amount of fixation following nodule formation) of nitrogen fixation appears to be heritable. Examining the population means following adjustment for the covariates revealed that the Ds2 mean was adjusted upward (Table 68) as expected from earlier data (Table 60).

Since Ds2 was shown to be similar to Ds5 for the unadjusted estimate means and significantly different for cotyledon and juvenile leaf size it may be assumed that Ds2 is more "efficient" for total fixation per unit area of cotyledon or juvenile leaf than is Ds5.

A method to investigate this "efficiency" of fixation would be to divide the nitrogen fixation estimates by either the cotyledon area or juvenile leaf width to create characters of ratios of fixation per unit of cotyledon area or juvenile leaf width. The population means for the estimate ratios have been given in Tables 69 and 70 for cotyledon area and juvenile leaf width adjustment, respectively. For the estimate ratios, Ds2 was again found to be significantly more "efficient" than Ds5 for all estimates except root fresh weight. The analysis of variance for the estimate ratios revealed that total genetic effects

Table 66. Mean squares from an analysis of covariance of the original characters for Experiment VI.

Source	df	Acetylene reduced (nm/pl/hr)	Shoot fresh weight (mg/pl)	Root fresh weight (g/pl)	Shoot dry weight (mg/pl)	Nitrogen fixation "factor"
Covariates	2	15799**	1926033**	44.5**	132383**	4058**
Cotyledon area	1	10589**	1209806**	18.0**	80914**	2293***
Juvenile leaf width	1	4570*	619485**	22.5**	44374**	1517***
Populations	5	5831**	357361**	1.5*	33075**	605***
Additive	1	24088**	1469771**	3.7*	133226**	2441***
Dominance	1	628	136462	0.6	17143*	212
EpiStasis	3	1480	60186	1.1	5002	125
Add-Add	1	2949	144609	1.0	11979	217
Add-Dom	1	1490	30737	2.2	2154	131
Dom-Dom	1	2	5212	0.1	872	26
Error	608	916	48042	0.7	3789	95

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

Table 67. Mean squares from an analysis of covariance of the log₁₀ transformed characters for Experiment VI.

Source	df	Acetylene reduced (nM/pl/hr)	Acetylene fresh weight (mg/pl)	Root fresh weight (g/pl)	Shoot dry weight (mg/pl)	Nitrogen fixation "factor" (mg/pl)
Covariates	2	1.134**	1.666**	3.331**	1.850**	2.906**
Cotyledon area	1	0.788**	1.079**	1.683**	1.215**	1.839**
Juvenile leaf width	1	0.308*	0.511**	1.426**	0.554**	0.924**
Populations	5	0.303**	0.240**	0.041	0.334**	0.308**
Additive	1	1.266**	1.060**	0.164*	1.392**	1.345**
Dominance	1	0.010	0.059	0.020	0.131	0.053
Epistasis	3	0.080	0.017	0.007	0.049	0.047
Add-Add	1	0.008	0.000	0.015	0.144	0.139
Add-Dom	1	0.026	0.050	0.006	0.002	0.002
Dom-Dom	1	0.205	0.000	0.000	0.000	0.000
Error	608	0.055	0.032	0.041	0.039	0.052

*Significant at the 5% level of probability.

**Significant at the 1% level of probability.

Table 68. Adjusted means from an analysis of covariance for Experiment VI.

Population	n	Acetylene	Shoot fresh weight	Root fresh weight	Shoot dry weight	Nitrogen fixation "factor"
		(nm/pl/hr)	(mg/pl)	(g/pl)	(mg/pl)	
original						
P ₁ (D ₈₂)	58	78.0	657	1.88	173	24.9
P ₂ (D ₈₅)	55	47.2	432	1.47	106	15.4
F ₁	59	68.7	617	1.78	164	23.0
F ₂	229	57.1	519	1.67	134	19.0
B ₁	108	65.0	596	1.77	158	22.1
B ₂	111	58.1	521	1.85	135	20.3
log ₁₀						
P ₁ (D ₈₂)	58	1.85	2.79	0.21	2.20	1.35
P ₂ (D ₈₅)	55	1.64	2.61	0.14	2.00	1.15
F ₁	59	1.78	2.75	0.20	2.17	1.30
F ₂	229	1.70	2.68	0.17	2.08	1.22
B ₁	108	1.75	2.75	0.19	2.15	1.29
B ₂	111	1.68	2.66	0.19	2.06	1.20

Table 69. Means, standard deviations, and coefficients of variation for N fixation estimates/cotyledon area for Experiment VI.

Population	n	Acetylene reduced ratio		Shoot fresh weight ratio	Root fresh weight ratio	Shoot dry weight ratio	Nitrogen fixation "factor" ratio
		Shoot fresh weight ratio	Root fresh weight ratio	Shoot dry weight ratio	Root dry weight ratio	Shoot dry weight ratio	Nitrogen fixation "factor" ratio
P ₁ (DS2)	58	2.25±1.35(60) ^a	18.2±10.2(56) ^a	.047±.030(64) ^a	4.74±2.82(59) ^a	.659±.437(66) ^a	
P ₂ (DS5)	55	1.29±0.57(44) ^c	11.7± 3.8(32) ^c	.041±.015(37) ^a	2.98±1.04(35) ^c	.429±.176(41) ^c	
F ₁	59	1.78±0.80(45) ^b	16.0± 5.6(40) ^{ab}	.047±.021(45) ^a	4.25±1.52(36) ^{ab}	.602±.243(40) ^{ab}	
F ₂	229	1.52±0.76(50) ^{bc}	13.8± 5.6(41) ^{bc}	.044±.021(48) ^a	3.54±1.57(44) ^c	.509±.249(49) ^{bc}	
B ₁	108	1.75±0.97(55) ^b	15.9± 8.0(50) ^{ab}	.045±.025(56) ^a	4.18±2.22(53) ^{ab}	.570±.333(58) ^{ab}	
B ₂	111	1.53±0.77(50) ^{bc}	13.9± 5.7(41) ^{bc}	.049±.024(49) ^a	3.59±1.62(45) ^{bc}	.529±.260(49) ^{bc}	

means followed by the same letter are not significantly different for the Student-Newman-Keul range test.

Table 70. Means, standard deviations, and coefficients of variation for N fixation estimates/juvenile leaf width for Experiment VI.

Population n	Acetylene reduced ratio	Shoot fresh weight ratio	Root fresh weight ratio	Shoot dry weight ratio	Nitrogen fixation "factor" ratio
P ₁ (D ₈₂)	58	11.4±6.2(54)a	91.8±45.8(50)a	.24±.15(63)a	23.9±12.9(54)a
P ₂ (D ₈₅)	55	7.0±2.7(39)c	64.7±19.4(30)c	.23±.08(35)a	16.4± 5.4(33)d
F ₁	59	10.2±4.7(46)ab	91.8±31.2(35)a	.27±.11(41)a	24.3± 8.6(35)a
F ₂	229	8.5±4.4(52)bc	77.2±31.5(41)b	.25±.12(48)a	19.8± 8.8(44)c
B ₁	108	9.5±4.9(52)b	85.7±38.3(45)ab	.24±.13(54)a	22.6±11.0(49)ab
B ₂	111	8.6±4.5(52)bc	78.0±33.6(43)ab	.27±.14(52)a	20.2± 9.5(47)bc

*means followed by the same letter are not significantly different for the Student-Newman-Keul range test.

were additive for cotyledon adjusted data (Table 71) whereas dominance effects were shown to be present for shoot weights and the "Nitrogen fixation" factor for the juvenile leaf adjusted data (Table 72). Grouping the shoot dry weight estimate ratios and crosstabulating with either cotyledon area or juvenile leaf groups, revealed the presence of plants with large cotyledons or juvenile leaves with "efficiency" of fixation and vice versa (Tables 73 and 74). It would appear to be possible to select plants with vigorous seedlings which were "efficient" for fixation.

D. Host genes for symbiotic nitrogen fixation in Desmodium.

The genes controlling the formation of nitrogenase are found on the Rhizobium genome (Bergelson, 1974). Since each experiment was inoculated with what was assumed to be a homogeneous Rhizobium culture then the nitrogenase enzyme should be uniform within each experiment. Significant genotype effects may indicate that the host plants differed in their ability to support nitrogen fixation physiologically. Nitrogenase would not be expected to be the limiting factor since it was expected to be the same for the "efficient" as for the "inefficient" genotypes. It must be concluded that there are genes within Desmodium which may indirectly regulate nitrogen fixation. Increasing the fixation rate of the Desmodium - Rhizobium interaction may not be completely effective by solely manipulating the Rhizobium strains which has been the traditional practice.

Table 71. Mean squares for the N fixation estimates/cotyledon area for Experiment VI.

Source	df	Shoot fresh weight ratio	Root fresh weight ratio	Shoot dry weight ratio	Nitrogen fixation "factor" ratio
Populations	5	7.08**	349**	.0007	27.4**
Additive	1	29.95**	1533**	.0006	117.5**
Dominance	1	0.12	25	.0006	4.1
Epistasis	3	1.77	62	.0608	5.2
Add-Add	1	3.55	140	.0005	12.6
Add-Dom	1	1.76	41	.0017	2.3
Dom-Dom	1	0.00	5	.0000	0.6
Error	614	0.75	42	.0005	3.3
					0.081

*significant at the 5% level of probability.

**significant at the 1% level of probability.

Table 72. Mean squares for the N fixation estimates/juvenile leaf width for Experiment VI.

Source	df	Acetylene reduced ratio	Shoot fresh weight ratio	Root fresh weight ratio	Shoot dry weight ratio	Nitrogen fixation "factor" ratio
Populations	5	143.0**	7017**	.0256	593**	9.07**
Additive	1	573.9**	24726**	.0005	1594**	25.38**
Dominance	1	24.1	6500*	.6740*	633***	12.61*
Epistasis	3	39.0	1286	.0178	113	2.45
Add-Add	1	75.6	2930	.0080	283	3.78
Add-Dom	1	41.3	685	.0437	48	3.50
Dom-Dom	1	.1	.44	.0017	9	0.07
Error	614	21.3	1140	.0151	91	2.26

*significant at the 5% level of probability.

**significant at the 1% level of probability.

Table 73. Numbers of plants for shoot dry weight/cotyledon area ratio group by cotyledon area or juvenile leaf width crosstabulation for Experiment VI.

		Shoot dry weight/cotyledon area ratio				
		< 3.5	3.5-7.0	> 7.0		
Cotyledon area						Row total
< 32 mm ²	I	I	I	I	I	
	I	66	I	67	I	24
	I		I		I	
32-39 mm ²	I	I	I	I	I	
	I	74	I	52	I	4
	I		I		I	
39-42 mm ²	I	I	I	I	I	
	I	95	I	73	I	7
	I		I		I	
> 42 mm ²	I	I	I	I	I	
	I	82	I	76	I	4
	I		I		I	
Juvenile leaf width						Row total
< 6 mm	I	I	I	I	I	
	I	103	I	73	I	17
	I		I		I	
6-7 mm	I	I	I	I	I	
	I	133	I	103	I	16
	I		I		I	
> 7 mm	I	I	I	I	I	
	I	80	I	92	I	6
	I		I		I	
Column total		317	268	39	624	

Table 74. Numbers of plants for shoot dry weight/juvenile leaf width ratio group by cotyledon area or juvenile leaf width crosstabulation for Experiment VI.

		Shoot dry weight/juvenile leaf width ratio						
		< 20	20-40	> 40				
Cotyledon area						Row total		
< 32 mm ²	I	I	I	I	I			
	I	88	I	57	I	10	I	155
	I		I		I		I	
32-39 mm ²	I	I	I	I	I	I		
	I	88	I	37	I	4	I	129
	I		I		I		I	
39-42 mm ²	I	I	I	I	I	I		
	I	91	I	77	I	7	I	175
	I		I		I		I	
> 42 mm ²	I	I	I	I	I	I		
	I	74	I	83	I	5	I	162
	I		I		I		I	
Juvenile leaf width						Row total		
< 6 mm	I	I	I	I	I			
	I	101	I	80	I	11	I	192
	I		I		I		I	
6-7 mm	I	I	I	I	I	I		
	I	143	I	98	I	11	I	252
	I		I		I		I	
> 7 mm	I	I	I	I	I	I		
	I	98	I	76	I	4	I	178
	I		I		I		I	
Column total		342	254	26		622		

All analyses of the unadjusted estimates of nitrogen fixation revealed that all F_1 crosses exhibited heterosis when the plants were grown in a nitrogen limiting environment. However, when nitrogen is supplied in adequate amounts, there appears to be no difference between the F_1 or parental plants (McWhirter, 1969; Rotar and Chow, 1971). The heterosis in this study may be simply the expression of parental inbreeding depression when the plants are grown in a nitrogen limiting environment. These conclusions are supported by the negative estimates of the parental S.C.A. effects from the diallel studies (Table 50).

E. Breeding procedures for nitrogen fixation in Desmodium.

The analyses of this study revealed two major points concerning the inheritance of symbiotic nitrogen fixation in Desmodium. These hypotheses are: (a) that there was a major dominant gene conditioning strain-specific ineffective nodulation, and (b) that there was significant genetic variation for total symbiotic nitrogen fixation among effectively nodulated Desmodium genotypes and that this variation appears to have a heritable basis. Partitioning this genetic variance suggested that additive genetic variation and dominance variance were present with dominance variance being much larger when the estimates of nitrogen are not adjusted for juvenile plant vigor. Adjusting the estimates of nitrogen fixation for juvenile plant vigor revealed an almost totally additive (heritable) basis to the

genetic variation.

1. Strain-specific ineffective nodulation genes.

The presence of strain-specific ineffective nodulation genes may be construed as a disadvantage when breeding for increased nitrogen fixation in Desmodium. However, these genes may also be used to advantage. If a Desmodium line or hybrid were to be released as a commercial cultivar it would be necessary to eliminate major genes causing ineffective nodulation. A rapid screening program with recurrent selection for the recessive genotype (effective nodulation) during early generation breeding trials would quickly remove ineffective nodulation genes from the potential cultivars.

The ineffective nodulation genes may also be used by a microbiologist as a means of selecting for nodulation competitiveness among strains of Rhizobium (Vincent, 1974). Having selected a series of Desmodium tester lines which carry at least one strain-specific ineffective nodulation gene, a microbiologist may initiate competition studies to rank the strains from most to least competitive for the nodulation sites. The most competitive strain if desirable in all other characteristics would be released as the commercial inoculum. The plant breeder having available a desirable strain of Rhizobium would begin to select or breed for a Desmodium hybrid or inbred line which would maximize the total fixation by the Desmodium line - commercial strain interaction.

2. Increasing total symbiotic nitrogen fixation.

The outstanding feature of the data from the quantitative experiments was the expression of heterosis. Those data clearly indicated that heterozygous genotypes of Desmodium were significantly superior to inbred lines for total symbiotic nitrogen fixation. A hybrid breeding program exploiting this heterosis would be desirable.

The reporting of cytoplasmic-genetic male-sterility in Desmodium (McWhirter, 1969) greatly assists the initiation of a hybrid breeding program. The male-sterile factor was found in line Ds1 and fertility was partially restored in crosses with line Da1. Male-sterile factors were also found in lines Ds2 and Da2. However fertility restoration was poor (McWhirter, pers. com.). Before initiating a hybrid breeding program for total fixation, complete fertility restoration would have to be established.

However, with the presence of significant average heterosis the accumulation of efficient inbred lines should be accompanied by a hybrid testing program. Such a breeding procedure has been referred to as reciprocal recurrent selection (Comstock et al., 1949). Base populations for use in a reciprocal recurrent selection program could involve the phenotypically distinct yet fully fertile D. sandwicense by D. aparines or D. aparines by D. uncinatum crosses.

Inbred lines necessary for this program could be developed either by mass selection, i.e. preservation of additive effects or by single seed descent (Goulden, 1939).

3. Nitrogen fixation components.

Heterosis or dominance genetic effects appear to be the rule when the nitrogen fixation estimates are analysed in their totality. However, when the total variation for each estimate is partitioned into components the dominance effects disappear. The genetic variation of each part is almost exclusively additive (heritable). In Experiment VI, two components of total nitrogen fixed were defined as; (a) juvenile plant vigor; estimated by cotyledon area and juvenile leaf width, and (b) "efficiency" of nitrogen fixation as estimated by the ratio of total nitrogen fixed to juvenile plant vigor. Large cotyledon and juvenile leaf size appeared to be conditioned by a dominant allele at a single locus. Plants with large cotyledons were shown to fix significantly more nitrogen so it would be advantageous at the early stages of a breeding program to select for large cotyledons or juvenile leaves. These plants would probably be more vigorous and therefore nodulate more quickly or form nodules at more sites. These results are similar to the results reported by Mytton (1973) where large seed weight assisted nodulation. It would be expected that selection for large cotyledons and juvenile leaves in *Desmodium* would rapidly respond to selection since a single gene appears to control these characters. Also achieved with this selection would be plants with increased seedling vigor which would establish a sward and begin fixing nitrogen and dry matter more quickly. Having selected a population of seedlings with

large cotyledons or juvenile leaves the plants would enter the second phase of the breeding program; namely selection for "efficiency" of nitrogen fixation. Highly efficient plants would be tested against the selected *Rhizobium* strains (discussed in part 1 of this section) to determine the most favorable interaction which would be competitive for nodulation sites under field conditions. Such superior interactions could be selected for superior general adaptability or superior specific adaptability to a particular location which exhibits unique environmental conditions. If an interaction was selected for general adaptability, this interaction would have to be tested under many locations, sites, and/or years since environmental conditions greatly effect total nitrogen fixation.

At this early stage in the development of the study of host variation for nitrogen fixation, the best avenue to pursue to achieve maximum return for research input is not clear. A hybrid breeding program working directly with heterosis may be more effective where capital and facilities are not limiting since the heritable variation is maintained under reciprocal recurrent selection. The limitation of the breeding program would be the numbers of individual plants which would have to be maintained to insure that the program was effective.

Initiating a breeding program, emphasising the components of total nitrogen fixation would be more advantageous for laboratories with limiting resources. After

selection for vigorous juvenile plants the populations could be effectively reduced by the elimination of the non-vigorous plants. The researcher could concentrate his resources to a smaller number of plants because the genetic basis of the components appears to be heritable and a selection response for greater "efficiency" would be expected.

It may be easy to speculate about the expected results of a breeding program employing either approach but the ultimate criterion judging the success of any research for increased nitrogen fixation would be a selection response culminating in a inbred variety or the creation of a superior hybrid which fixes a maximum amount of nitrogen under field conditions. Only when the products of research achieve practical results under agricultural conditions is the research program of merit in achieving its goals.

F. Relevance to tropical agriculture.

The importance of Desmodium in tropical agriculture is two-fold. Firstly, Desmodium being a forage legume would provide increased forage protein for animal consumption thereby effectively increasing availability of protein for human consumption. Secondly, Desmodium as a legume, fixes atmospheric nitrogen which assists non-leguminous plants to produce increased yield. If the total fixation could be increased Desmodium would be valuable in the tropics as a green manure crop and a soil conditioner as well as a forage

legume.

The pending population increases in the tropics will dictate that the food chains be made as efficient as possible, i.e direct plant to human pathways for protein. D. sandwicense may be used in this context as a model organism for legume studies because of its many attributes for glasshouse growth. Principles of inheritance found in D. sandwicense may be directly applicable to other self-fertile grain legumes which are used for human consumption. The future of the need for adequate amounts of protein for human consumption cannot be underestimated and it is in this area where legumes may make a great contribution.

The secondary contribution of legumes would be as a rotational crop with cereals to buffer against catastrophes. In short, it would be optimal to develop annual grain legumes which fix more total nitrogen and increases protein production and rotate these legumes with the cereal crops which have been bred to produce maximum energy.

SUMMARY AND CONCLUSIONS

The genetic study of symbiotic nitrogen fixation in the tropical legume Desmodium was divided into three sections; namely, methodology, qualitative genetic variation, and quantitative genetic variation. The initial studies in the methodology section suggested that the bottle jar was effective for studying nitrogen fixation in Desmodium. Uninoculated or ineffectively nodulated control plants were yellow and chlorotic suggesting the lack of nitrogen. Green plants were taken from the bottle jars and analysed with the acetylene reduction assay. Dry weight accumulation was also recorded. Reduction of acetylene to ethylene proceeded at a linear rate for approximately 90 minutes. Linear reduction indicated that senescence of harvested Desmodium plants was not a limiting factor for the acetylene reduction analysis.

The within-bottle jar variation among similar genotypes suggested that the design of the bottle jar could be modified to reduce within-bottle jar variation. Modifications to the design and sand particle size greatly reduced this variation.

Initial nodulation studies indicated that some D. aparines lines were ineffectively nodulated by strain B. Studies of F_2 and backcross populations from D. sandwicense x D. aparines crosses implied that the strain-specific ineffective nodulation interaction between some D. aparines lines and strain B was conditioned by a dominant allele at a single locus.

The analysis of quantitative genetic variation suggested that several points were of importance. Firstly, the correlation between dry weight accumulation and the acetylene reduction assay was highly significant and there appeared to be a consistent pattern for both characters within an experiment. The acetylene reduction may be used as a plant breeding tool to screen juvenile *Desmodium* plants.

The most significant aspect illustrated by all quantitative experiments was the expression of F_1 heterosis. F_1 hybrids were significantly superior to inbred lines for total nitrogen fixed. A breeding program incorporating reciprocal recurrent selection would maximize the effect of F_1 heterosis. Although F_1 heterosis was significant, a portion of the genetic variance also appeared to be heritable and analysis of components of total nitrogen fixation predominantly expressed additive (heritable) genetic variation. Selecting inbred lines for these heritable effects would be beneficial since these lines could become the basis for recurrent selection programs or released as cultivars.

Since genes controlling the formation of nitrogenase are found on the *Rhizobium* genome, quantitative variation in total nitrogen fixed may be due to physiological differences among host plants. The host plant may indirectly control symbiotic nitrogen fixation.

Genotype by environment interactions were of significant importance suggesting that studies concerning

quantitative variation for symbiotic nitrogen fixation should be conducted under differing environments, locations, and/or years.

Breeding programs utilizing qualitative and quantitative genetic variation were outlined. Such a program would be useful for the production of highly efficient varieties. The relevance of such varieties to tropical agriculture was also discussed.

The preceding analysis suggested that the null hypothesis is to be rejected. The null hypothesis was that there was no genetic variation in the processes of the symbiotic relationship between Desmodium and Rhizobium. Two hypotheses may be postulated. These are: (a) that strain-specific ineffective nodulation interaction between strain B and some D. uncinatum lines was conditioned by a dominant allele at a single locus, and (b) that there was genetic variation in symbiotic nitrogen fixation in Desmodium.

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APPENDICES

Appendix 'A': Crossing results from Colombia, S.A.

Table 1. Hybridization and emasculation observations using various washing solutions and flower preparations.

Treatment		Number of flowers manipulated	Percent pod set	Avg. no. seeds per pod
Distilled water	- partial flower†	53	50	1.7
	- entire flower	32	84	3.9
	- emasculated	45	0	
2.5% sugar solution	- partial flower	28	60	3.4
	- entire flower	45	95	5.8
	- emasculated	63	5	2.3
5.0% sugar solution	- partial flower	43	46	1.8
	- emasculated	18	0	

†the tip half of the petals removed.

Table 2. The percent pod set and the average number of seeds per pod obtained during a 5x5 diallel cross.

Method	Number of flowers manipulated	Percent pod set	Avg. no. seeds per pod	Number of seeds tested	Percent hybrids
Pollinated	387	8.2	4.3	104	95
Emasculated	60	6.6	2.5		

Table 3. The correlations between environmental factors and hybridization results.

	Percent pod set	Av. no. of seeds
Max. daily temp.	0.15	-.39
Min. daily temp.	0.30	-.19
Mean daily temp.	0.24	-.41
Temp. at pollination	.0.07	-.17
SQRT(Daily precipitation)	-.24	0.19

SQRT = square root

Appendix B. Diallel analysis computer program listing.

1: DIALL subroutine instructions.	187
2: DIALL subroutine listing.	191
3: DIALL sample output.	206

Acknowledgment: A model I, method 1 diallel program was obtained from Mr. P. Hsu, Dept. of Plant Science, The Univ. of Alberta. This program was extensively edited, expanded and corrected by the author to produce an all model and method diallel program.

Subroutine name: DIALL
 Subroutine title: Diallel analysis
 Programmer: B. Pinchbeck

Program Description

Source Language: Fortran IV

Purpose: The analysis of a diallel cross for models I, II, and A; methods 1 to 4.

Description: DIALL is a algorithm which computes the diallel analysis of variance, components of genetic variance, Griffing's combining ability effects, or Gardner-Eberhart's constants. The subroutine generally follows the diallel analysis as outlined by Griffing (1956) yet is modified to include the analysis of variance of Hayman (1954) and Gardner and Eberhart (1966) for methods 1 and 2, respectively. The diallel table ($p \times p$) or within-block diallel tables ($p \times p \times b$) are calculated from the formulae:

$$X_{ij} = \frac{1}{\text{SQRT}(bs)} \sum Z_{ijkl} \quad (\text{for a two-way randomized block design with interactions})$$

or

$$X_{ij} = \frac{1}{\text{SQRT}(b)} \sum Z_{ijk} \quad (\text{for a two-way randomized block design})$$

and

$$(X_{ij})_k = \frac{1}{\text{SQRT}(s)} \sum (Z_{ij})_k ; k=1,b$$

and

$$S_{bg} = \sum_k (S_g)_k - S_g$$

where

p = number of diallel parents

b = number of blocks (replications, locations, years, etc.)

s = number of subsamples per cell

X = diallel table entry

Z = individual cell entries

X_{ij} = the ijth entry in the overall diallel table

- $(x_{ij})_k$ = the ijth entry in the diallel table of the kth block.
- S_{bg} = the sum of squares of the block x genetic component interaction
- $(S_g)_k$ = the sum of squares for the genetic component in the kth block.
- S_g = the sum of squares for the genetic component for the overall diallel table.

The above adjustments were incorporated so that the sum of the sum of squares of the main effect genetic components would equal to the treatment sum of squares for the two-way analysis of variance.

Griffing's combining ability effects and Gardner-Eberhart's constants are calculated from a diallel table of genotype means as outlined by Griffing (1956), and Gardner and Eberhart (1966), respectively.

Components of genetic variance are calculated from the diallel analysis of variance such that:

$$\hat{V}_A = 2\hat{V}_g \text{ (inbred parents)}$$

$$\hat{V}_A = 4\hat{V}_g \text{ (noninbred parents)}$$

$$\hat{V}_D = \hat{V}_h + \hat{V}_r \text{ (methods 1 and 3; inbred parents)}$$

$$\hat{V}_D = 4\hat{V}_h + \hat{V}_r \text{ (methods 1 and 3; non-inbred parents)}$$

$$\hat{V}_D = \hat{V}_h \text{ (methods 2 and 4; inbred parents)}$$

$$\hat{V}_D = 4\hat{V}_h \text{ (methods 2 and 4; non-inbred parents)}$$

$$\hat{V}_G = \hat{V}_A + \hat{V}_D$$

$$\hat{V}_P = \hat{V}_G + \hat{V}_E \text{ (model A)}$$

$$\hat{V}_P = \hat{V}_G + \hat{V}_{BG} + \hat{V}_E \text{ (model II)}$$

and where the estimated variance of the components are adjusted for block and subsamples. (for example)

$$\hat{V}_g = \frac{1}{bs(p+2)} (M_g - M_h) \text{ (for model II, method 2)}$$

whereas Griffing (1956) presents the formula:

$$\hat{V}_g = \frac{1}{p+2} (M_g - M_h)$$

and where

\hat{V}_g = the general (G.C.A.) variance component

\hat{V}_h = the heterosis (S.C.A.) variance component

\hat{V}_r = the reciprocal variance component

\hat{V}_A = the additive genetic variance component

\hat{V}_D = the dominance genetic variance component

\hat{V}_G = the genetic variance component

\hat{V}_P = the phenotypic variance component

\hat{V}_{BG} = the block x genotype variance component

\hat{V}_E = the environmental variance component (error mean square)

Restrictions:

The subroutine is limited to an input diallel table matrix of genotype means of 10 parents x 10 parents x 10 blocks.

Program Usage

Calling Sequence: CALL DIALL (NP, NQ, NR, NS, MO, ME, NA, CH, IE, EMS, A, IW, IP)

Parameters:

Name	Purpose
NP -	number of parents
NQ -	rank of the third dimension of the diallel table
NR -	number of blocks in the original design
NS -	the number of subsamples per cell in the original design
MO -	diallel model number 0 - fixed model I - Gardner-Eberhart constants 1 - fixed model I - Griffing combining ability effects 2 - random model II - components of variation 3 - mixed model A - components of variation
ME -	diallel method number 1 = parents, F_1 's and reciprocal F_1 's

2 = parents and F_1 's
 3 = F_1 's and reciprocal F_1 's
 4 = F_1 's
 0 = parents and F_1 's (Gardner-Eberhart Analysis III)

NA - a vector describing the experiment (20A4)
 CH - a vector describing the analysed character (10A4)
 IE - error degrees of freedom
 EMS - error mean square
 A - matrix of block sums if $NP \times NP$ diallel table or
 subsample means if $NP \times NP \times NQ$ diallel table
 IW - logical I/O unit for output statements WRITE (IW, FMT)...
 IP - 0 - if parents are inbred
 1 - if parents are not inbred

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SUBROUTINE DIALL(NP,NQ,NR,NS,MO,ME,NA,CH,IE,EMS,A,IW,IP)
 DIALLEL ANALYSIS SUBROUTINE
 VERSION 1.0, JUNE 1979
 C
 NP = NUMBER OF PARENTS (MAX. 10)
 NQ = NUMBER OF DIMENSIONS OF DIALLEL TABLE (MAX. 10)
 (I.E. THE RANK OF THE THIRD DIMENSION OF 'A')
 NR = NUMBER OF REPLICATIONS (BLOCKS, LOCATIONS, YEARS)
 IN THE ORIGINAL DESIGN
 NS = NUMBER OF SUBSAMPLES PER CELL IN THE ORIGINAL DESIGN
 MO = DIALLEL MODEL NUMBER
 0 = FIXED MODEL I - GARDNER-EBERHART CONSTANTS
 1 = FIXED MODEL I - GRIFFING'S COMBINING ABILITY EFFECTS
 2 = RANDOM MODEL II - COMPONENTS OF VARIANCE
 3 = MIXED MODEL A - COMPONENTS OF VARIANCE
 ME = DIALLEL METHOD NUMBER
 0 = PARENTS AND F1'S (GARDNER-EBERHART ANALYSIS III)
 1 = PARENTS, F1'S, AND RECIPROCAL F1'S
 2 = PARENTS AND F1'S
 3 = F1'S AND RECIPROCAL F1'S
 4 = F1'S
 NA = VECTOR DESCRIBING THE EXPERIMENT (20A4)
 CH = VECTOR DESCRIBING THE CHARACTER BEING ANALYSED (10A4)
 IE = ERROR DEGREES OF FREEDOM
 EMS = ERROR MEAN SQUARE
 A = MATRIX OF BLOCK SUMS IF NP X NP DIALLEL TABLE OR
 SUBSAMPLE MEANS IF NP X NP X NQ DIALLEL TABLE
 IW = LOGICAL I/O UNIT FOR OUTPUT STATEMENTS : WRITE(IW,FMT)...
 IP = 0 IF PARENTS ARE INBRED
 1 IF PARENTS ARE NOT INBRED

PROGRAMMED BY BRIAN R. PINCHBECK

```

DIMENSION A(10,10,10),NA(20),CH(10)
DIMENSION S(10,10),R(10,10),X(10),Y(10),G(10),VS(10),VG(10)
DOUBLE PRECISION RN,SN,A,S,R,X,Y,W,G,VS,VG,BT,BA,ED,BDD,BPD,BCD,BR
&,BMR,BOR,SST,SSA,SSD,SSDD,SSPD,SSCD,SSR,SSMR,SSOR,TOT
COMMON W(10,10)
IA=NP-1
ID=NP*IA/2
ICD=NP*(NP-3)/2
IOR=(NP-1)*(NP-2)/2
IF (NS.LE.0) NS=1
IF (ME.EQ.0) ID=ID-1
PP=2.0
PU=1.0
PPP=1.0
IF (IP.EQ.1) PP=4.0
IF (IP.EQ.1) PPP=16.0
IF (IP.EQ.1) PU=4.0
IB=NR-1
IBA=IB*IA
IBD=IB*ID
IBCD=IB*ICD
IBOR=IB*IOR
AI=IA
DI=ID
CDI=ICD
ORI=IOR
BI=IB
BAI=IBA
BDI=IBD
BCDI=IBCD
BORI=IBOR
SN=NS
RN=NR
PN=NP
BT=0.0
BA=0.0
BD=0.0
BDD=0.0
BPD=0.0
BCD=0.0
BR=0.0
BMR=0.0
BOR=0.0
BMT=0.0
BT1=0.0
TOT=0.0
ESS=EMS*IE

```

```

LME=ME+1
GO TO (190,10,190,360,550),LME
10 CONTINUE
C METHOD 1 ANALYSIS OF VARIANCE
IT=NP•2-1
TI=IT
IF (NQ.LE.1) GO TO 40
IBT=IT•IB
BTI=IBT
DO 30 K=1,NQ
DO 20 I=1,NP
DO 20 J=1,NP
W(I,J)=A(I,J,K)*DSQRT(SN)
20 CONTINUE
CALL DAV1(NP,SST,SSA,SSD,SSDD,SSPD,SSCD,SSR,SSMR,SSOR)
BT=BT+SST
BA=BA+SSA
BD=BD+SSD
BDD=BDD+SSDD
BPD=BPD+SSPD
BCD=BCD+SSCD
BR=BR+SSR
BMR=BMR+SSMR
BOR=BOR+SSOR
30 CONTINUE
40 CONTINUE
DO 50 I=1,NP
DO 50 J=1,NP
W(I,J)=0.0
DO 50 K=1,NQ
W(I,J)=W(I,J)+A(I,J,K)*DSQRT(SN)/DSQRT(RN)
50 CONTINUE
CALL DAV1(NP,SST,SSA,SSD,SSDD,SSPD,SSCD,SSR,SSMR,SSOR)
BT=BT-SST
BA=BA-SSA
BD=BD-SSD
BDD=BDD-SSDD
BPD=BPD-SSPD
BCD=BCD-SSCD
BR=BR-SSR
BMR=BMR-SSMR
BOR=BOR-SSOR
SMT=SST/TI
SMA=SSA/AI
SMD=SSD/DI
SMPD=SSPD/AI
SMCD=SSCD/CDI
SMR=SSR/DI
SMMR=SSMR/AI
SMOR=SSOR/QRI
FAT=SMT/EMS
FBT=SMT/BMT
FAA=SMA/EMS
FAD=SMD/EMS
FADD=SSDD/EMS
FAPD=SMPD/EMS
FACD=SMCD/EMS
FAR=SMR/EMS
FAMR=SMMR/EMS
FAOR=SMOR/EMS
PAT=FPROB(IT,IE,FAT)
PAA=FPROB(IA,IE,FAA)
PAD=FPROB(ID,IE,FAD)
PADD=FPROB(1,IE,SSDD)
PAPD=FPROB(IA,IE,FAPD)
PACD=FPROB(ICD,IE,FACD)
PAR=FPROB(ID,IE,FAR)
PAMR=FPROB(IA,IE,FAMR)
PAOR=FPROB(IOR,IE,FAOR)
IF (NQ.LE.1) GO TO 60
BMT=BT/BTI
BMA=BA/BAI
BMD=BD/BDI
BMDD=BDD/BI
BMPD=BPD/BAI
BMCD=BCD/BCDI
BRM=BR/BDI
BMRM=BMR/BAI
BORM=BOR/BORI
FABT=BMT/EMS

```

```

FABA=BMA/EMS
FABD=BMD/EMS
FABDD=BMDD/EMS
FABPD=BMPD/EMS
FABCD=BMCD/EMS
FABR=BRM/EMS
FABMR=BMRM/EMS
FABOR=BORM/EMS
PABT=FPROB(IBT, IE, FABT)
PABA=FPROB(IBA, IE, FABA)
PABD=FPROB(IBD, IE, FABD)
PABDD=FPROB(IB, IE, FABDD)
PABPD=FPROB(IBA, IE, FABPD)
PABCD=FPROB(BCD, IE, FABCD)
PABR=FPROB(IBD, IE, FABR)
PABMR=FPROB(IBA, IE, FABMR)
PABOR=FPROB(BCR, IE, FABOR)

60 CONTINUE
    WRITE(IW,730) ME,NA,CH
    WRITE(IW,720)
    WRITE(IW,740)
    WRITE(IW,1020) IT,SST,SMT,FAT,PAT
    WRITE(IW,750) IA,SSA,SMA,FAA,PAA
    WRITE(IW,760) ID,SSD,SMD,FAD,PAD
    WRITE(IW,770) SSDD,SSDD,FADD,PADD
    WRITE(IW,780) IA,SSPD,SMPD,FAPD,PAPD
    WRITE(IW,790) ICD,SSCD,SMCD,FACD,PACD
    WRITE(IW,800) ID,SSR,SMR,FAR,PAR
    WRITE(IW,810) IA,SSMR,SMMR,FAMR,PAMR
    WRITE(IW,820) IOR,SSOR,SMOR,FAOR,PAOR
    IF(NQ.LE.1) GO TO 70
    WRITE(IW,830) IBT,BT,BMT,FABT,PABT
    WRITE(IW,840) IBA,BA,BMA,FABA,PABA
    WRITE(IW,850) IBD,BD,BMD,FABD,PABD
    WRITE(IW,860) IB,BDD,BMDD,FABDD,PABDD
    WRITE(IW,870) IBA,BPD,BMPD,FABPD,PABPD
    WRITE(IW,880) IBCD,BCD,BMCD,FABCD,PABCD
    WRITE(IW,890) IBD,BR,BRM,FABR,PABR
    WRITE(IW,900) IBA,BMR,BMRM,FABMR,PABMR
    WRITE(IW,910) IBOR,BOR,BORM,FABOR,PABOR

70 CONTINUE
    WRITE(IW,920) IE,ESS,EMS
    WRITE(IW,730) ME,NA,CH
    WRITE(IW,1160) (I,I=1,NP)
    DO 90 I=1,NP
        X(I)=0.0
        Y(I)=0.0
        DO 80 J=1,NP
            W(I,J)=W(I,J)/DSQRT(SN)/DSQRT(RN)
            X(I)=X(I)+W(I,J)
            Y(I)=Y(I)+W(J,I)
            TOT=TOT+W(I,J)
        80 CONTINUE
        WRITE(IW,1080) I,(W(I,J),J=1,NP)
    90 CONTINUE
    XMU=TOT/PN**2
    SE1=SQRT(EMS/PN**2)
    WRITE(IW,1130) XMU,SE1

C METHOD 1 COMPONENTS OF VARIANCE
    IF(MO.LT.2) GO TO 100
    WRITE(IW,730) ME,NA,CH
    CALL COV(NP,NR,NS,MO,SMA,SMD,SMR,BMT,EMS,AI,DI,BTI,IE,PP,PU,PPP)
    RETURN

100 CONTINUE
    IF(MO.NE.1) GO TO 170
C METHOD 1 GRIFFING'S COMBINING ABILITY ANALYSIS
    EMS=EMS/RN/SN
    DO 120 I=1,NP
        R(I,I)=0.0
        VS(I)=0.
        G(I)=((X(I)+Y(I))/(2.*PN))-XMU
        DO 110 J=1,NP
            S(I,J)=(W(I,J)+W(J,I))/2.-(X(I)+X(J)+Y(I)+Y(J))/(2.*PN)+XMU
            S(J,I)=S(I,J)
            IF(I-J).LT.110,120,
        110 CONTINUE
        VS(I)=VS(I)+S(I,J)**2
        R(I,J)=(W(I,J)-W(J,I))/2.
        R(J,I)=-R(I,J)
    120 CONTINUE

```

```

DO 130 I=1,NP
VG(I)=G(I)*G(I)-(PN-1.)/PN/PN/2.*EMS
VS(I)=VS(I)/PN/2.-(PN-1.)/PN/2.*EMS
130 CONTINUE
SE2=SQRT((EMS*(PN-1.))/2./PN/PN)
SE3=SQRT(EMS/PN)
SE4=SQRT((EMS*(PN-1.)*(PN-1.))/PN/PN)
SE5=SQRT((EMS*(PN*PN-2.*PN+2.))/2./PN/PN)
SE6=SQRT(EMS*2.*PN-2.)/PN)
SE7=SQRT(EMS*(PN-1.)/PN)
SE8=SQRT(EMS*(PN-2.)/PN)
SE9=SQRT(EMS*(3.*PN-2.)/2./PN)
SE10=SQRT(EMS*1.5*(PN-2.)/PN)
SE11=SQRT(EMS/2.)
SE12=SQRT(EMS)
WRITE (IW,730) ME,NA,CH
WRITE (IW,1030)
WRITE (IW,1040)
DO 140 I=1,NP
WRITE (IW,1050) I,G(I),VG(I),VS(I)
140 CONTINUE
WRITE (IW,1060) SE2,SE3
WRITE (IW,1070) (I,I=1,NP)
DO 150 I=1,NP
WRITE (IW,1080) I,(S(I,J),J=1,I)
150 CONTINUE
WRITE (IW,1090) SE4,SE5,SE6,SE7,SE8
WRITE (IW,1100) SE9,SE10
WRITE (IW,1110) (I,I=1,NP)
DO 160 I=1,NP
WRITE (IW,1080) I,(R(I,J),J=1,NP)
160 CONTINUE
WRITE (IW,1120) SE11,SE12
RETURN
170 CONTINUE
DO 180 I=1,NP
DO 180 J=I,NP
W(I,J)=(W(I,J)+W(J,I))/2.0
W(J,I)=W(I,J)
180 CONTINUE
C GARDNER-EBERHART CONSTANTS
WRITE (IW,730) ME,NA,CH
CALL GARDCO(NP,ME,IW)
RETURN
190 CONTINUE
C METHOD 2 ANALYSIS OF VARIANCE
IT=NP*(NP+1)/2-1
TI=IT
IF (NQ.LE.1) GO TO 220
IBT=IT*IB
BTI=IBT
DO 210 K=1,NQ
DO 200 I=1,NP
DO 200 J=I,NP
W(I,J)=A(I,J,K)*DSQRT(SN)
W(J,I)=W(I,J)
200 CONTINUE
CALL DAV2(NP,ME,SST,SSA,SSD,SSDD,SSPD,SSCD)
BT=BT+SST
BA=BA+SSA
BD=BD+SSD
BDD=BDD+SSDD
BPD=BPD+SSPD
BCD=BCD+SSCD
210 CONTINUE
220 CONTINUE
DO 230 I=1,NP
DO 230 J=I,NP
W(I,J)=0.0
DO 230 K=1,NQ
W(I,J)=W(I,J)+A(I,J,K)*DSQRT(SN)/DSQRT(RN)
W(J,I)=W(I,J)
230 CONTINUE
CALL DAV2(NP,ME,SST,SSA,SSD,SSDD,SSPD,SSCD)
BT=BT-SST
BD=BD-SSD
BA=BA-SSA
BDD=BDD-SSDD
BPD=BPD-SSPD
BCD=BCD-SSCD

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SMT=SST/TI
SMA=SSA/AI
SMD=SSD/DI
SMPD=SSPD/AI
SMCD=SSCD/CDI
FAT=SMT/EMS
FAA=SMA/EMS
FAD=SMD/EMS
FADD=SSDD/EMS
FAPD=SMPD/EMS
FACD=SMCD/EMS
PAT=FPROB(IT, IE, FAT)
PAA=FPROB(IA, IE, FAA)
PAD=FPROB(ID, IE, FAD)
PADD=FFROB(1, IE, FADD)
PAPD=FPROB(IT, IE, FAPD)
PACD=FPROB(ICD, IE, FACD)
IF (NQ.LE.1) GO TO 240
BMT=BT/BTI
BMA=BA/BAI
BMD=BD/BDI
BMDD=BDD/BI
BMPD=BPD/BAI
BMCD=BCD/BCDI
FABT=BMT/EMS
FABA=BMA/EMS
FABD=BMD/EMS
FABDD=BMDD/EMS
FABPD=BMPD/FMS
FABCD=BMCD/EMS
PABT=FPROB(IBT, IE, FABT)
PABA=FPROB(IBA, IE, FABA)
PABD=FPROB(IBD, IE, FABD)
PABDD=FPROB(IB, IE, FABDD)
PABPD=FPROB(IBA, IE, FABPD)
PABCD=FPROB(ICD, IE, FABCD)
240 CONTINUE
WRITE (IW,730) ME,NA,CH
WRITE (IW,720)
WRITE (IW,740)
WRITE (IW,1020) IT,SST,SMT,FAT,PAT
IF (ME.EQ.2) GO TO 250
WRITE (IW,950) IA,SSPD,SMPD,FAPD,PAPD
WRITE (IW,940) SSDD,SSDD,FADD,PADD
WRITE (IW,930) ID,SSD,SMD,FAD,PAD
WRITE (IW,960) IA,SSA,SMA,FAA,PAA
WRITE (IW,980) ICD,SSCD,SMCD,FACD,PACD
IF (NQ.LE.1) GO TO 260
WRITE (IW,830) IBT,BT,BMT,FABT,PABT
WRITE (IW,1010) IBA,BPD,BMPD,FABPD,PABPD
WRITE (IW,1000) IB,BDD,BMDD,FABDD,PABDD
WRITE (IW,990) IBD,BD,BMD,FABD,PABD
WRITE (IW,970) IBA,BA,BMA,FABA,PABA
WRITE (IW,880) IBCD,BCD,BMCD,FABCD,PABCD
GO TO 260
250 WRITE (IW,750) IA,SSA,SMA,FAA,PAA
WRITE (IW,760) ID,SSD,SMD,FAD,PAD
WRITE (IW,770) SSDD,SSDD,FADD,PADD
WRITE (IW,780) IA,SSPD,SMPD,FAPD,PAPD
WRITE (IW,790) ICD,SSCD,SMCD,FACD,PACD
IF (NQ.LE.1) GO TO 260
WRITE (IW,830) IBT,BT,BMT,FABT,PABT
WRITE (IW,840) IBA,BA,BMA,FABA,PABA
WRITE (IW,850) IBD,BD,BMD,FABD,PABD
WRITE (IW,860) IB,BDD,BMDD,FABDD,PABDD
WRITE (IW,870) IBA,BPD,BMPD,FABPD,PABPD
WRITE (IW,880) IBCD,BCD,BMCD,FABCD,PABCD
260 CONTINUE
WRITE (IW,920) IE,ESS,EMS
WRITE (IW,730) ME,NA,CH
WRITE (IW,1170) (I,I=1,NP)
DO 290 I=1,NP
X(I)=0.0
DO 270 J=1,NP
W(I,J)=W(I,J)/DSQRT(SN)/DSQRT(RN)
X(I)=X(I)+W(I,J)
270 CONTINUE
DO 280 J=1,I
TOT=TOT+W(I,J)
280 CONTINUE

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      WRITE (IW,1080) I,(W(I,J),J=1,I)
290 CONTINUE
      XMU=TOT*2./PN/(PN+1.)
      SE1=SQRT(EMS*2./PN/(PN+1.))
      WRITE (IW,1130) XMU,SE1
C METHOD 2 COMPONENTS OF VARIANCE
      IF (MO.LT.2) GO TO 300
      WRITE (IW,730) ME,NA,CH
      CALL COV(NP,NR,NS,MO,SMA,SMD,0.0,BMT,EMS,AI,DI,BTI,IE,PP,PU,PPP)
      RETURN
300 CONTINUE
C GARDNER-EBERHART CONSTANTS
      IF (MO.NE.0) GO TO 310
      WRITE (IW,730) ME,NA,CH
      CALL GARDCO(NP,ME,IW)
      RETURN
310 CONTINUE
C METHOD 2 GRIFFING'S COMBINING ABILITY ANALYSIS
      EMS=EMS/RN/SN
      DO 320 I=1,NP
      VS(I)=0.0
      G(I)=((X(I)+W(I,I))-(2.*TOT)/PN)/(PN+2.)
      DO 320 J=I,NP
      S(I,J)=W(I,J)-((X(I)+X(J)+W(I,I)+W(J,J))/(PN+2.))+((2.*TOT)/((PN+
      &1.)*(PN+2.)))
      S(J,I)=S(I,J)
      IF (I.EQ.J) GO TO 320
      VS(I)=VS(I)+S(I,J)**2
320 CONTINUE
      DO 330 I=1,NP
      VG(I)=G(I)**2-(((PN-1.)/(PN*(PN-2.)))*EMS)
      VS(I)=(VS(I)/(PN-2.))-(((PN-3.)/(PN-2.))*EMS)
330 CONTINUE
      SE2=SQRT(((PN-1.)/(PN*(PN+2.)))*EMS)
      SE3=SQRT((2.*EMS)/(PN+2.))
      SE4=SQRT(((PN*(PN-1.))/((PN+1.)*(PN+2.)))*EMS)
      SE5=SQRT(((PN**2+PN+2.)/((PN+1.)*(PN+2.)))*EMS)
      SE6=SQRT(((2.*(PN-2.))/(PN+2.))*EMS)
      SE7=SQRT(((2.*(PN+1.))/(PN+2.))*EMS)
      SE8=SQRT(((2.*PN)/(PN+2.))*EMS)
      WRITE (IW,730) ME,NA,CH
      WRITE (IW,1030)
      WRITE (IW,1040)
      DO 340 I=1,NP
      WRITE (IW,1050) I,G(I),VG(I),VS(I)
340 CONTINUE
      WRITE (IW,1060) SE2,SE3
      WRITE (IW,1070) (I,I=1,NP)
      DO 350 I=1,NP
      WRITE (IW,1080) I,(S(I,J),J=1,I)
350 CONTINUE
      WRITE (IW,1090) SE4,SE5,SE6,SE7,SE8
      RETURN
360 CONTINUE
C METHOD 3 ANALYSIS OF VARIANCE
      IT=NP**2-NP-1
      TI=IT
      IF (NQ.LE.1) GO TO 390
      IBT=IT*IB
      BTI=IBT
      DO 380 K=1,NQ
      DO 370 I=1,NP
      DO 370 J=1,NP
      W(I,J)=A(I,J,K)*DSQRT(SN)
370 CONTINUE
      CALL DAV3(NP,SST,SSA,SSD,SSR,SSMR,SSOR)
      BT=BT+SST
      BA=BA+SSA
      BD=BD+SSD
      BR=BR+SSR
      BMR=BMR+SSMR
      BOR=BOR+SSOR
380 CONTINUE
390 CONTINUE
      DO 400 I=1,NP
      DO 400 J=1,NP
      W(I,J)=0.0
      DO 400 K=1,NQ
      W(I,J)=W(I,J)+A(I,J,K)*DSQRT(SN)/DSQRT(RN)
400 CONTINUE

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CALL DAV3(NP,SST,SSA,SSD,SSR,SSMR,SSOR)
 BT=BT-SST
 BA=BA-SSA
 BD=BD-SSD
 BR=BR-SSR
 BMR=BMR-SSMR
 BOR=BOR-SSOR
 SMT=SST/TI
 SMA=SSA/AI
 SMD=SSD/CDI
 SMR=SSR/DI
 SMMR=SSMR/AI
 SMOR=SSOR/ORI
 FAT=SMT/EMS
 FAA=SMA/EMS
 FAD=SMD/EMS
 FAR=SMR/EMS
 FAMR=SMMR/EMS
 FAOR=SMOR/EMS
 PAT=FPROB(IT,IE,FAT)
 PAA=FPROB(IA,IE,FAA)
 PAD=FPROB(ID,IE,FAD)
 PAR=FPROB(ID,IE,FAR)
 PAMR=FPROB(IA,IE,FAMR)
 PAOR=FPROB(IOR,IE,FAOR)
 IF (NQ.LE.1) GO TO 410
 BMT=BT/BTI
 BMA=BA/BAI
 BMD=BD/BCDI
 BRM=BR/BDI
 BMMR=BMR/BAI
 BMOR=BOR/BORI
 FABT=BMT/EMS
 FABA=BMA/EMS
 FABD=BMD/EMS
 FABR=BRM/EMS
 FABMR=BMMR/EMS
 FABOR=BMOR/EMS
 PABT=FPROB(IBT,IE,FABT)
 PABA=FPROB(IBA,IE,FABA)
 PABD=FPROB(IBD,IE,FABD)
 PABR=FPROB(IBD,IE,FABR)
 PABMR=FPROB(IBA,IE,FABMR)
 PABOR=FPROB(IBCR,IE,FABOR)

410 CONTINUE
 WRITE(IW,730) ME,NA,CH
 WRITE(IW,720)
 WRITE(IW,740)
 WRITE(IW,1020) IT,SST,SMT,FAT,PAT
 WRITE(IW,750) IA,SSA,SMA,FAA,PAA
 WRITE(IW,760) ICD,SSD,SMD,FAD,PAD
 WRITE(IW,800) ID,SSR,SMR,FAR,PAR
 WRITE(IW,810) IA,SSMR,SMMR,FAMR,PAMR
 WRITE(IW,820) IOR,SSOR,SMOR,FAOR,PAOR
 IF (NQ.LE.1) GO TO 420
 WRITE(IW,830) IBT,BT,BMT,FAET,PABT
 WRITE(IW,840) IBA,BA,BMA,FABA,PABA
 WRITE(IW,850) IBCD,BD,BMD,FABD,PABD
 WRITE(IW,890) IBD,BR,BRM,FABR,PAER
 WRITE(IW,900) IBA,BMR,BMMR,FABMR,PABMR
 WRITE(IW,910) IBOR,BOR,BMOR,FABOR,PABOR

420 CONTINUE
 WRITE(IW,920) IE,ESS,EMS
 EMS=EMS/RN/SN
 WRITE(IW,730) ME,NA,CH
 WRITE(IW,1160) (I,I=1,NP)
 DO 440 I=1,NP
 W(I,I)=0.0
 X(I)=0.0
 Y(I)=0.0
 DO 430 J=1,NP
 W(I,J)=W(I,J)/DSQRT(SN)/DSQRT(RN)
 X(I)=X(I)+W(I,J)
 Y(I)=Y(I)+W(J,I)
 TOT=TOT+W(I,J)
 430 CONTINUE
 WRITE(IW,1080) I,(W(I,J),J=1,NP)
 440 CONTINUE
 XMU=TOT/PN/(PN-1.)
 SE1=SQRT(EMS/PN/(PN-1.))

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        WRITE (IW, 1130) XMU, SE1
C  METHOD 3 COMPONENTS OF VARIANCE
  IF (MO.LT.2) GO TO 450
  WRITE (IW, 730) ME, NA, CH
  CALL COV(NP, NR, NS, MO, SMA, SMD, SMR, BMT, EMS, AI, DI, BTI, IE, PP, PU, PPP)
  RETURN
450 CONTINUE
  IF (MO.NE.1) GO TO 530
C  METHOD 3 GRIFFING'S COMBINING ABILITY ANALYSIS
  DO 460 I=1,NP
    R(I,I)=0.0
    S(I,I)=0.0
    VS(I)=0.0
460 CONTINUE
  IG=NP-1
  DO 470 I=1,IG
    K=I+1
    DO 470 J=K,NP
      S(I,J)=(W(I,J)+W(J,I))/2.-(X(I)+X(J)+Y(I)+Y(J))/2./((PN-2.))+TOT/(PN
      & -1.)/(PN-2.)
      R(I,J)=(W(I,J)-W(J,I))/2.
      S(J,I)=S(I,J)
      RCJ=-R(I,J)
470 CONTINUE
  DO 490 I=1,NP
  DO 480 J=1,NP
    VS(I)=VS(I)+S(I,J)**2
480 CONTINUE
  G(I)=(PN*(X(I)+Y(I))-2.*TOT)/2./PN/(PN-2.)
  VG(I)=G(I)**2-EMS*(PN-1.)/PN/(PN-2.)/2.
  VS(I)=VS(I)/(PN-2.)/2.-EMS*(PN-3.)/(PN-2.)/2.
490 CONTINUE
  SE2=SQRT(EMS*(PN-1.)/2./PN/(PN-2.))
  SE3=SQRT(EMS/(PN-2.))
  SE4=SQRT(EMS*(PN-3.)/2./PN-1.))
  SE5=SQRT(EMS*(PN-3.)/(PN-2.))
  SEL=SQRT(EMS*(PN-4.)/(PN-2.))
  SE7=SQRT(EMS/2.)
  WRITE (IW, 730) ME, NA, CH
  WRITE (IW, 1030)
  WRITE (IW, 1040)
  DO 500 I=1,NP
    WRITE (IW, 1050) I, G(I), VG(I), VS(I)
500 CONTINUE
  WRITE (IW, 1060) SE2, SE3
  WRITE (IW, 1070) (I, I=1, NP)
  DO 510 I=1,NP
    WRITE (IW, 1080) I, (S(I,J), J=1, I)
510 CONTINUE
  WRITE (IW, 1140) SE4, SE5, SE6
  WRITE (IW, 1110) (I, I=1, NP)
  DO 520 I=1,NP
    WRITE (IW, 1080) I, (R(I,J), J=1, NP)
520 CONTINUE
  WRITE (IW, 1150) SE7
  RETURN
530 CONTINUE
  DO 540 I=1,NP
  DO 540 J=I, NP
    W(I,J)=-(W(I,J)+W(J,I))/2.0
    W(J,I)=W(I,J)
540 CONTINUE
C  GARDNER-EBERHART CONSTANTS
  WRITE (IW, 730) ME, NA, CH
  CALL GARDCO(NP, ME, IW)
  RETURN
C  METHOD 4 ANALYSIS OF VARIANCE
550 CONTINUE
  IT=NP*(NP-1)/2-1
  TI=IT
  IF (NQ.LE.1) GO TO 580
  IBT=IT*IB
  BTI=IBT
  DO 570 K=1,NQ
  DO 560 Y=1,NP
  DO 560 J=1,NP
    W(I,J)=A(I,J,K)*DSQRT(SN)
    W(J,I)=W(I,J)
560 CONTINUE
  CALL DAV4(NP, SST, SSA, SSD)

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BT=BT+SST
BA=BA+SSA
BD=BD+SSD
570 CONTINUE
580 CONTINUE
DO 590 I=1,NP
DO 590 J=I,NP
W(I,J)=0.0
DO 590 K=1,NQ
W(I,J)=W(I,J)+A(I,J,K)*DSQRT(SN)/DSQRT(RN)
W(J,I)=W(I,J)
590 CONTINUE
CALL DAV4(NP,SST,SSA,SSD)
BT=BT-SST
BA=BA-SSA
BD=BD-SSD
SMT=SST/TI
SMA=SST/AI
SMD=SSD/CDI
FAT=SMT/EMS
FAA=SMA/EMS
FAD=SMD/EMS
PAT=FPROB(IT,IE,FAT)
PAA=FPROB(IA,IE,FAA)
PAD=FPROB(ID,IE,FAD)
IF (NQ.LE.1) GO TO 600
BMT=BT/BTI
BMA=BA/BAI
BMD=BD/BCDI
FABT=BMT/EMS
FABA=BMA/EMS
FABD=BMD/EMS
PABT=FPROB(IBT,IE,FABT)
PABA=FPROB(IBA,IE,FABA)
PABD=FPROB(IBD,IE,FABD)
600 CONTINUE
WRITE (IW,730) ME,NA,CH
WRITE (IW,720)
WRITE (IW,740)
WRITE (IW,1020) IT,SST,SMT,FAT,PAT
WRITE (IW,750) IA,SSA,SMA,FAA,PAA
WRITE (IW,760) ICD,SSD,SMD,FAD,PAD
IF (NQ.LE.1) GO TO 610
WRITE (IW,830) IBT,BT,BMT,FABT,PABT
WRITE (IW,840) IBA,BA,BMA,FABA,PABA
WRITE (IW,850) IBCD,BD,BMD,FABD,PABD
610 CONTINUE
WRITE (IW,920) IE,ESS,EMS
WRITE (IW,730) ME,NA,CH
WRITE (IW,1170) (I,I=1,NP)
DO 640 I=1,NP
W(I,I)=0.0
X(I)=0.0
DO 620 J=1,NP
W(I,J)=W(I,J)/DSQRT(SN)/DSQRT(RN)
X(I)=X(I)+W(I,J)
620 CONTINUE
DO 630 J=I,NP
TOT=TOT+W(I,J)
630 CONTINUE
WRITE (IW,1080) I,(W(I,J),J=1,I)
640 CONTINUE
SE1=SQRT(EMS*2./PN/(PN-1.))
XMU=2.*TOT/PN/(PN-1.)
WRITE (IW,1130) XMU,SE1
C METHOD 4 COMPONENTS OF VARIANCE
IF (MO.LT.2) GO TO 650
WRITE (IW,730) ME,NA,CH
CALL COV(NP,NR,NS,MO,SMA,SMD,0.0,BMT,EMS,AI,DI,BTI,IE,PP,PU,PPP)
RETURN
650 CONTINUE
C GARDNER-EBERHART CONSTANTS
IF (MO.NE.0) GO TO 660
WRITE (IW,730) ME,NA,CH
CALL GARDCO(NP,ME,IW)
RETURN
660 CONTINUE
C METHOD 4 GRIFFING'S COMBINING ABILITY ANALYSIS
EMS=EMS/RN/SN
DO 670 I=1,NP

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S(I,I)=0.0
DO 670 J=I,NP
IF (I.EQ.J) GO TO 670
S(I,J)=W(I,J)-(X(I)+X(J))/(PN-2.)+2.*TOT/(PN-1.)/(PN-2.)
S(J,I)=S(I,J)
670 CONTINUE
DO 690 I=1,NP
DO 680 J=1,NP
VS(I)=VS(I)+S(I,J)
680 CONTINUE
G(I)=(PN*X(I)-2.*TOT)/PN/(PN-2.)
VG(I)=G(I)**2-EMS*(PN-1.)/PN/(PN-2.)
VS(I)=VS(I)/(PN-2.)-EMS*(PN-3.)/(PN-2.)
690 CONTINUE
SE2=SQRT(EMS*(PN-1.)/PN/(PN-2.))
SE3=SQRT(EMS*2.)/(PN-2.)
SE4=SQRT(EMS*(PN-3.)/(PN-1.))
SE5=SQRT(EMS*2.*PN-3.)/(PN-2.)
SE6=SQRT(EMS*2.*PN-4.)/(PN-2.)
WRITE (IW,730) ME,NA,CH
WRITE (IW,1030)
WRITE (IW,1040)
DO 700 I=1,NP
WRITE (IW,1050) I,G(I),VG(I),VS(I)
700 CONTINUE
WRITE (IW,1060) SE2,SE3
WRITE (IW,1070) (I,I=1,NP)
DO 710 I=1,NP
WRITE (IW,1080) I,(S(I,J),J=1,I)
710 CONTINUE
WRITE (IW,1140) SE4,SE5,SE6
RETURN
720 FORMAT (' ', 'ANALYSIS OF VARIANCE')
730 FORMAT ('1', 'METHOD', I2, ' DIALLEL ANALYSIS'//1X,20A4//1X,10A4//)
740 FORMAT ('0', 'SOURCE OF VARIATION', 12X, 'DF', 8X, 'SUM OF SQUARES', 8X,
& 'MEAN SQUARES', 18X, 'F', 14X, 'P')
750 FORMAT ('0', 2X, 'GENERAL (GCA)', 15X, I3, 2F20.4, 10X, 2F15.4)
760 FORMAT ('0', 2X, 'HETEROSESIS (SCA)', 13X, I3, 2F20.4, 10X, 2F15.4)
770 FORMAT ('0', 4X, 'AVERAGE HETEROSESIS', 11X, '1', 2F20.4, 10X, 2F15.4)
780 FORMAT ('0', 4X, 'VARIETY HETEROSESIS', 9X, I3, 2F20.4, 10X, 2F15.4)
790 FORMAT ('0', 4X, 'SPECIFIC HETEROSESIS', 8X, I3, 2F20.4, 10X, 2F15.4)
800 FORMAT ('0', 2X, 'RECIPROCAL', 18X, I3, 2F20.4, 10X, 2F15.4)
810 FORMAT ('0', 4X, 'MATERNAL RECIPROCAL', 7X, I3, 2F20.4, 10X, 2F15.4)
820 FORMAT ('0', 4X, 'NON-MATERNAL RECIPROCAL', 3X, I3, 2F20.4, 10X, 2F15.4)
830 FORMAT ('0', 'GENOTYPE INTERACTION', 10X, I3, 2F20.4, 10X, 2F15.4)
840 FORMAT ('0', 2X, 'GENERAL INTERACTION', 9X, I3, 2F20.4, 10X, 2F15.4)
850 FORMAT ('0', 2X, 'HETEROSESIS INTERACTION', 7X, I3, 2F20.4, 10X, 2F15.4)
860 FORMAT ('0', 4X, 'AVERAGE INTERACTION', 7X, I3, 2F20.4, 10X, 2F15.4)
870 FORMAT ('0', 4X, 'VARIETY INTERACTION', 7X, I3, 2F20.4, 10X, 2F15.4)
880 FORMAT ('0', 4X, 'SPECIFIC INTERACTION', 6X, I3, 2F20.4, 10X, 2F15.4)
890 FORMAT ('0', 2X, 'RECIPROCAL INTERACTION', 6X, I3, 2F20.4, 10X, 2F15.4)
900 FORMAT ('0', 4X, 'MATERNAL INTERACTION', 6X, I3, 2F20.4, 10X, 2F15.4)
910 FORMAT ('0', 4X, 'NON-MATERNAL INTERACTION', 2X, I3, 2F20.4, 10X, 2F15.4)
920 FORMAT ('0', 'ERROR', 24X, I4, 2F20.4//)
930 FORMAT ('0', 2X, 'CROSSES', 21X, I3, 2F20.4, 10X, 2F15.4)
940 FORMAT ('0', 2X, 'VARIETIES VS CROSSES', 10X, '1', 2F20.4, 10X, 2F15.4)
950 FORMAT ('0', 2X, 'VARIETIES', 19X, I3, 2F20.4, 10X, 2F15.4)
960 FORMAT ('0', 4X, 'GENERAL (GCA)', 13X, I3, 2F20.4, 10X, 2F15.4)
970 FORMAT ('0', 4X, 'GENERAL INTERACTION', 7X, I3, 2F20.4, 10X, 2F15.4)
980 FORMAT ('0', 4X, 'SPECIFIC (SCA)', 12X, I3, 2F20.4, 10X, 2F15.4)
990 FORMAT ('0', 2X, 'CROSSES INTERACTION', 9X, I3, 2F20.4, 10X, 2F15.4)
1000 FORMAT ('0', 2X, 'VARIETIES VS CROSSES INT.', 3X, I3, 2F20.4, 10X, 2F15.4)
&
1010 FORMAT ('0', 2X, 'VARIETIES INTERACTION', 7X, I3, 2F20.4, 10X, 2F15.4)
1020 FORMAT ('0', 'GENOTYPES', 21X, I3, 2F20.4, 10X, 2F15.4)
1030 FORMAT ('0', 'GRIFFING'S COMBINING ABILITY ANALYSIS (MODEL 1//)
1040 FORMAT ('0', 10X, 'PARENT', 5X, 'G.C.A. EFFECTS', 7X, 'G.C.A. VARIANCE',
& 7X, 'S.C.A. VARIANCE')
1050 FORMAT (' ', 12X, I2, 9X, F10.4, 2(10X, F12.4))
1060 FORMAT ('0', 10X, 'S.E. OF THE G.C.A. EFFECTS -', 20X, F10.4/11X, 'S.E.
& OF THE DIFFERENCES BETWEEN G.C.A. EFFECTS -', F10.4//)
1070 FORMAT ('0', 10X, 'S.C.A. EFFECTS'//6X, 'PARENTS', 4X, 15, 9110)
1080 FORMAT ('0', 11X, I2, 2X, 10F10.4)
1090 FORMAT ('0', 10X, 'S.E. OF THE PARENTAL S.C.A. EFFECTS -', 35X, F10.4/
& 11X, 'S.E. OF THE CROSS S.C.A. EFFECTS -', 38X, F10.4/11X, 'S.E. OF THE
& PARENTAL DIFFERENCES (S.C.A. EFFECTS) -', 21X, F10.4/11X, 'S.E. OF THE
& COMMON PARENT CROSS DIFFERENCES (S.C.A. EFFECTS) -', 10X, F10.4/
& 11X, 'S.E. OF THE DIFFERENT PARENT CROSS DIFFERENCES (S.C.A. EFFECT
& S) -', 7X, F10.4)
1100 FORMAT (' ', 10X, 'S.E. OF THE PARENT MINUS PROGENY DIFFERENCES (S.C.

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& A. EFFECTS) - ', 9X, F10.4/11X, 'S.E. OF THE PARENT MINUS NON-PROGENY
 & DIFFERENCES (S.C.A. EFFECTS) - ', 5X, F10.4)
 1110 FORMAT ('0', 10X, 'RECIPROCAL EFFECTS'//25X, 'MALE PARENT'// 'FEMALE
 & PARENT', 18, 9110)
 1120 FORMAT ('0', 10X, 'S.E. OF THE RECIPROCAL EFFECTS - ', 40X, F10.4/11X, '
 & S.E. OF THE DIFFERENCES BETWEEN RECIPROCAL EFFECTS - ', 20X, F10.4//)
 1130 FORMAT ('0', 'POPULATION MEAN - ', F10.4, 5X, 'S.E. - ', F10.4//)
 1140 FORMAT ('0', 10X, 'S.E. OF S.C.A. EFFECTS - ', 48X, F10.4/11X, 'S.E. OF
 & THE COMMON PARENT CROSS DIFFERENCES (S.C.A. EFFECTS) - ', 10X,
 & F10.4/11X, 'S.E. OF THE DIFFERENT PARENT CROSS DIFFERENCES (S.C.A.
 & EFFECTS) - ', 7X, F10.4)
 1150 FORMAT ('0', 'S.E. OF THE RECIPROCAL EFFECTS - ', 40X, F10.4)
 1160 FORMAT ('0', 'DIALLEL MEANS'//25X, 'MALE PARENT'// 'FEMALE PARENT', 18
 &, 9110)
 1170 FORMAT ('0', 'DIALLEL MEANS'//6X, 'PARENTS', 4X, 15, 9110)
 END
 SUBROUTINE DAV1(NP, SST, SSA, SSD, SSDD, SSPD, SSCD, SSR, SSMR, SSOR)
 DIMENSION X(10), Y(10)
 DOUBLE PRECISION PN, X, Y, W, SST, SSA, SSD, SSDD, SSPD, SSCD, SSR, SSMR, SSOR
 &, X1, X2, X4, X6, X8, H2, H4, H6, T0, SP, SSP
 COMMON W(10, 10)
 PN=NP
 IA=NP-1
 X1=0.0
 X2=0.0
 X4=0.0
 X6=0.0
 X8=0.0
 H2=0.0
 H4=0.0
 H6=0.0
 T0=0.0
 SF=0.0
 SSP=0.0
 DO 10 I=1, NP
 X(I)=0.0
 Y(I)=0.0
 DO 10 J=1, NP
 X1=X1+W(I, J)**2
 X(I)=X(I)+W(J, I)
 Y(I)=Y(I)+W(I, J)
 10 CONTINUE
 DO 20 I=1, NP
 X2=X2+(X(I)+Y(I))**2
 X4=X4+(X(I)-Y(I))**2
 X6=X6+(X(I)+Y(I)-2.*W(I, I))**2
 X8=X8+(X(I)+Y(I)-PN*W(I, I))**2
 T0=T0+X(I)
 SP=SP+W(I, I)
 SSP=SSP+W(I, I)**2
 DO 20 J=1, NP
 H2=H2+(W(I, J)+W(J, I))**2
 20 CONTINUE
 DO 30 I=1, IA
 DO 30 K=I, IA
 J=K+1
 H4=H4+(W(I, J)-W(J, I))**2
 H6=H6+(W(I, J)+W(J, I))**2
 30 CONTINUE
 SST=X1-T0**2/PN/PN
 SSA=(X2/2./PN)-(2.*T0**2/PN/PN)
 SSD=(H2/4.)-(X2/2./PN)+(T0**2/PN/PN)
 SSDD=(T0-(PN*SP))**2/(PN-1.)/PN**2
 SSPD=(X8-((2.*T0-PN*SP)**2)/PN))/PN/(PN-2.)
 SSCD=(H6/2.)-(X6/2./(PN-2.))+((T0-SP)**2/(PN-2.)/(PN-1.))
 SSR=H4/2.
 SSMR=X4/2./PN
 SSOR=SSR-SSMR
 RETURN
 END
 SUBROUTINE DAV2(NP, ME, SST, SSA, SSD, SSDD, SSPD, SSCD)
 DIMENSION X(10)
 DOUBLE PRECISION PN, X, W, SST, SSA, SSD, SSDD, SSPD, SSCD, SSA, SP, T0, X2,
 & X4, X6, X8, H2, SSP
 COMMON W(10, 10)
 PN=NP
 SP=0.0
 T0=0.0
 X2=0.0
 X4=0.0

```

X6=0.0
X8=0.0
H2=0.0
SSP=0.0
DO 20 I=1,NP
  X(I)=0.0
  SP=SP+W(I,I)
  SSP=SSP+W(I,I)**2
DO 10 J=1,NP
  X(I)=X(I)+W(I,J)
10 CONTINUE
  X2=X2+(X(I)+W(I,I))**2
  X4=X4+(X(I)-W(I,I))**2
  X6=X6+(X(I)+W(I,I))-((PN+2.)/2.)*W(I,I)**2
  DO 20 J=1,NP
    H2=H2+W(I,J)**2
    TO=TO+W(I,J)
20 CONTINUE
  SST=H2-2.*TO**2/PN/(PN+1.)
  SSA=(X2-4.*TO**2/PN)/(PN+2.)
  SSAA=X4/(PN-2.)-4.*TO-SP)**2/PN/(PN-2.)
  SSD=H2-X2/(PN+2.)+2.*TO**2/(PN+1.)/(PN+2.)
  SSDD=(TO-(PN+1.)/2.*SP)**2*4./PN/(PN**2-1.)
  SSPD=4.*((X6-((2.*TO-(PN+2.)/2.*SP)**2/PN))/(PN**2-4.))
  SSCD=(H2-SSP)-X4/(PN-2.)+2.*TO-SP)**2/(PN-1.)/(PN-2.)
  IF (ME.NE.0) RETURN
  SSPD=SSPD+SSA-SSAA
  SSA=SSAA
  SSD=SSA+SSCD
  RETURN
END
SUBROUTINE DAV3(NP,SST,SSA,SSD,SSR,SSMR,SSOR)
DIMENSION X(10),Y(10)
DOUBLE PRECISION PN,X,Y,W,TO,SST,SSA,SSD,SSR,SSMR,SSOR,X1,X2,X4,H2
&,H4
COMMON W(10,10)
PN=NP
TO=0.0
X1=0.0
X2=0.0
X4=0.0
H2=0.0
H4=0.0
DO 20 I=1,NP
  W(I,I)=0.0
  X(I)=0.0
  Y(I)=0.0
DO 10 J=1,NP
  X1=X1+W(I,J)**2
  X(I)=X(I)+W(J,I)
  Y(I)=Y(I)+W(I,J)
10 CONTINUE
  TO=TO+X(I)
  X2=X2+(X(I)+Y(I))**2
  X4=X4+(X(I)-Y(I))**2
  DO 20 J=1,NP
    H2=H2+(W(I,J)+W(J,I))**2
    H4=H4+(W(I,J)-W(J,I))**2
20 CONTINUE
  SST=X1-TO**2/PN/(PN-1.)
  SSA=X2/2./(PN-2.)-2.*TO**2/PN/(PN-2.)
  SSD=H2/2.-X2/2./(PN-2.)+TO**2/(PN-1.)/(PN-2.)
  SSR=H4/2.
  SSMR=X4/2./PN
  SSOR=SSR-SSMR
  RETURN
END
SUBROUTINE DAV4(NP,SST,SSA,SSD)
DIMENSION X(10)
DOUBLE PRECISION PN,X,W,SST,SSA,SSD,X2,H2,TO
COMMON W(10,10)
PN=NP
X2=0.0
H2=0.0
TO=0.0
DO 20 I=1,NP
  W(I,I)=0.0
  X(I)=0.0
DO 10 J=1,NP
  X(I)=X(I)+W(I,J)

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10 CONTINUE
X2=X2+X(I)**2
DO 20 J=I,NP
H2=H2+W(I,J)**2
T0=T0+W(I,J)
20 CONTINUE
SST=H2-2.*T0**2/PN/(PN-1.)
SSA=X2/(PN-2.)-4.*T0**2/PN/(PN-2.)
SSD=H2-X2/(PN-2.)*2.*T0**2/(PN-1.)/(PN-2.)
RETURN
END
SUBROUTINE GARDCO(NP,ME,IW)
C. GARDNER-EBERHART DIALLEL CONSTANTS SUBROUTINE
C. NP = NUMBER OF PARENTS
C. ME = DIALLEL METHOD NUMBER
C.   0 = PARENTS AND F1'S (GARDNER-EBERHART ANALYSIS III)
C.   1 = PARENTS, F1'S, AND RECIPROCAL F1'S
C.   2 = PARENTS AND F1'S
C.   3 = F1'S AND RECIPROCAL F1'S
C.   4 = F1'S
C. IW = LOGICAL I/O UNIT FOR OUTPUT STATEMENTS : WRITE(IW,FMT)...
C. W = (INPUT) DIALLEL TABLE (NP X NP)
C
DIMENSION X(10),Y(10)
DOUBLE PRECISION VM,CM,AH,X,Y,W
COMMON W(10,10)
NA=0
IF (ME.EQ.0) NA=1
C NA = TYPE OF ANALYSIS
C   0 = ANALYSIS II
C   1 = ANALYSIS III
WRITE (IW,90)
IF (NA.EQ.0) WRITE (IW,160)
IF (NA.NE.0) WRITE (IW,170)
VM=0.0
CM=0.0
DO 10 I=1,NP
VM=VM+W(I,I)/FLOAT(NP)
DO 10 J=1,NP
IF (I.EQ.J) GO TO 10
CM=CM+W(I,J)/FLOAT(NP*(NP-1))
10 CONTINUE
AH=CM-VM
IF (ME.GT.2) GO TO 30
WRITE (IW,100) VM,CM,AH
DO 20 I=1,NP
X(I)=W(I,I)-VM
WRITE (IW,120) I,X(I)
20 CONTINUE
GO TO 50
30 CONTINUE
VM=0.0
AH=0.0
WRITE (IW,110) CM
DO 40 I=1,NP
40 X(I)=0.0
50 CONTINUE
DO 60 I=1,NP
Y(I)=0.0
DO 60 J=1,NP
IF (I.EQ.J) GO TO 60
Y(I)=Y(I)+(W(I,J)-CM-(X(I)+X(J))/2.)/FLOAT(NP-1)
60 CONTINUE
WRITE (IW,130)
DO 70 I=1,NP
Y(I)=Y(I)+Y(I)/FLOAT(NP-2)
VH=Y(I)
IF (NA.NE.0) VH=Y(I)+X(I)/2.
WRITE (IW,120) I,VH
70 CONTINUE
WRITE (IW,140)
DO 80 I=1,NP
DO 80 J=I,NP
IF (I.EQ.J) GO TO 80
W(I,J)=W(I,J)-CM-(X(I)+X(J))/2.-Y(I)-Y(J)
WRITE (IW,150) I,J,W(I,J)
80 CONTINUE
RETURN
90 FORMAT ('GARDNER-EBERHART CONSTANTS')
100 FORMAT ('VARIETY MEAN',F15.4/' CROSS MEAN',F17.4/' AVERAGE HETERO'

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  &SIS',F10.4//' VARIETY CONSTANTS'/' PARENT',SX,'CONSTANT')
110 FORMAT ('OCROSS MEAN',F17.4)
120 FORMAT (' ',15,F14.4)
130 FORMAT ('Ovariety Heterosis Constants (II) or General (III)'/' PAR
  &ENT',SX,'CONSTANT')
140 FORMAT ('OSpecific Heterosis Constants'/' CROSS',SX,'CONSTANT')
150 FORMAT (' ',I2,' X ',I2,F13.4)
160 FORMAT ('OAnalysis II'/' )
170 FORMAT ('OAnalysis III'/' )
END
SUBROUTINE COV(NP,NR,NS,MO,SMA,SMD,SMR,BMT,EMS,AI,DI,BTI,IE,PP,PU,
&PPP)
PN=NP
RN=NR
SN=NS
C MODEL A DESIGN COMPONENTS OF VARIATION
CVA=0.0
CVD=0.0
CVR=0.0
VCA=0.0
VCD=0.0
VCR=0.0
HF=0.0
IF (SMA.GT.EMS) CVA=(SMA-EMS)/2./PN/RN/SN
IF (SMD.GT.EMS) CVD=(SMD-EMS)*PN**2/2./((PN**2-PN+1.)/RN/SN)
IF (SMR.GT.EMS) CVR=(SMR-EMS)/RN/SN/2.
VE=SQRT(2.*EMS**2/IE)
IF (SMA.GT.EMS) VCA=SQRT(2.*CVA**2/AI)
IF (SMD.GT.EMS) VCD=SQRT(2.*CVD**2/DI)
IF (SMR.GT.EMS) VCR=SQRT(2.*CVR**2/DI)
AV=PP*CVA
DV=PU*CVD+CVR
GV=DV+AV
PV=GV+EMS
VAV=PP*PP*VCA
VDV=PPP*VCD+VCR
VGV=VDV+VAV
VPV=VGV+VE
IHN=AV/PV*100
IHB=GV/PV*100
IF (DV.GT.0.0.AND.AV.GT.0.0) HF=SQRT(DV/AV)
IF (MO.EQ.2) GO TO 10
WRITE (IW,20)
WRITE (IW,30) CVA,VCA,CVD,VCD
WRITE (IW,40) CVR,VCR
WRITE (IW,50) EMS,VE,AV,VAV,DV,VDV,GV,VGV,PV,VPV,EMS,VE,IHN,IHB,HF
RETURN
10 CONTINUE
C MODEL II DESIGN COMPONENTS OF VARIATION
CVA=0.0
VCA=0.0
CVI=0.0
VCVI=0.0
HF=0.0
IF (SMA.GT.SMD) CVA=(SMA-SMD)/2./PN/RN/SN
IF (SMA.GT.SMD) VCA=SQRT(2.*CVA**2/AI)
IF (BMT.GT.EMS.AND.NR.GT.1.AND.NS.GT.1) CVI=(BMT-EMS)/SN
IF (BMT.GT.EMS.AND.NR.GT.1.AND.NS.GT.1) VCVI=SQRT(2.*CVI**2/BTI)
AV=PP*CVA
GV=DV+AV
PV=GV+EMS+CVI
VAV=PP*PP*VCA
VGV=VDV+VAV
VPV=VGV+VE+VCVI
IHN=AV/PV*100
IHB=GV/PV*100
IF (DV.GT.0.0.AND.AV.GT.0.0) HF=SQRT(DV/AV)
WRITE (IW,60)
WRITE (IW,30) CVA,VCA,CVD,VCD
WRITE (IW,40) CVR,VCR
WRITE (IW,50) EMS,VE,AV,VAV,DV,VDV,GV,VGV,PV,VPV,EMS,VE,IHN,IHB,HF
RETURN
20 FORMAT ('0','COMPONENTS OF GENETIC VARIANCE (MODEL A)'')
30 FORMAT ('0','9X,'GENERAL',F20.4,'+-',F15.4//10X,'HETEROsis',F18.4,
  &'+-',F15.4//)
40 FORMAT ('0','9X,'RECIPROCAL',F17.4,'+-',F15.4//)
50 FORMAT ('0','9X,'ERROR',F22.4,'+-',F15.4//10X,'ADDITIVE',F18.4,'
  &'+-',F15.4//10X,'DOMINANCE',F17.4,'+-',F15.4//10X,'GENETIC',F19.4
  &'+-',F15.4//10X,'PHENOTYPIC',F16.4,'+-',F15.4//10X,'ENVIRONMENT
  &AL',F13.4,'+-',F15.4//10X,'HERITABILITY'//15X,'NARROW',I10,'%'/)

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&/15X,'BROAD',111,'%'///10X,'HETEROsis FACTOR (SQRT(DOMINANCE/ADDITIONAL VARIANCE))',F10.4)
60 FORMAT ('0','COMPONENTS OF GENETIC VARIANCE (MODEL II) ')
END
FUNCTION FPROB(M,N,X)
C COMPUTES PROBABILITY OF F-STATISTIC X WITH M AND N DEGREES OF
C FREEDOM IN NUMERATOR AND DENOMINATOR RESPECTIVELY.
INTEGER A,B
FPROB=1.0
IF (X.LE.0..OR.M.LE.0.OR.N.LE.0) RETURN
A=2*(M/2)-M+2
B=2*(N/2)-N+2
W=X*M/N
Z=1./(1.+W)
IF (A.NE.1) GO TO 20
IF (B.NE.1) GO TO 10
P=SQRT(W)
Y=0.3183099
D=Y*Z/P
P=2.*Y*ATAN(P)
GO TO 40
10 P=SQRT(W*Z)
D=0.5*P*Z/W
GO TO 40
20 IF (B.NE.1) GO TO 30
P=SQRT(Z)
D=0.5*Z*P
P=1.-P
GO TO 40
30 D=Z*Z
P=W*Z
40 Y=2.*W/Z
J=B+2
IF (J.GT.N) GO TO 60
DO 50 K=J,N,2
D=(1.+A/(K-2.))*D*Z
IF (A.EQ.1) P=P+D*Y/(K-1)
IF (A.NE.1) P=(P+W)*Z
50 CONTINUE
60 Y=W*Z
Z=2./Z
B=N-2
I=A+2
IF (I.GT.M) GO TO 80
DO 70 K=I,M,2
J=K+B
D=Y*D*J/(K-2)
P=P-Z*D/J
70 CONTINUE
80 FPROB=1.-P
RETURN
END

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METHOD O DIALLEL ANALYSIS
 DESMODIUM DIALLEL ALL BLOCK ANALYSIS
 DRY WEIGHT IN A LEONARD JAR (MG/PL)

ANALYSIS OF VARIANCE

SOURCE OF VARIATION	DF	SUM OF SQUARES	MEAN SQUARES	F	P
GENOTYPES	14	4.9265	0.3519	8.9547	0.0008
VARIETIES	4	0.1062	0.0266	0.6759	0.6099
VARIETIES VS CROSSES	1	3.7614	3.7614	95.7158	0.0000
CROSSES	9	1.0589	0.1177	2.9941	0.0028
GENERAL (GCA)	4	0.6776	0.1694	4.3106	0.0026
SPECIFIC (SCA)	5	0.3814	0.0763	1.9409	0.0919
GENOTYPE INTERACTION	112	8.5038	0.0759	1.9321	0.0002
VARIETIES INTERACTION	32	1.9619	0.0613	1.5601	0.0435
VARIETIES VS CROSSES INT.	8	0.8969	0.1121	2.8530	0.0059
CROSSES INTERACTION	72	5.6450	0.0784	1.9951	0.0003
GENERAL INTERACTION	32	2.7854	0.0870	2.2150	0.0009
SPECIFIC INTERACTION	40	2.8596	0.0715	1.8192	0.0064
ERROR	129	5.0694	0.0393		

METHOD 2 DIALLEL ANALYSIS

DESIGNUM DIALLEL ALL BLOCK ANALYSIS
DRY WEIGHT IN A LEONARD JAR (OMG/PL)

ANALYSIS OF VARIANCE

SOURCE OF VARIATION	DF	SUM OF SQUARES	MEAN SQUARES	F	P
GENOTYPES	14	4.9265	0.3519	8.9547	0.0000
GENERAL (GCA)	4	0.4973	0.1243	3.1638	0.0162
HETEROSESIS (SCA)	10	4.4292	0.4429	11.2711	0.0000
AVERAGE HETEROSESIS	1	3.7614	3.7614	95.7158	0.0000
VARIETY HETEROSESIS	4	0.2865	0.0716	1.8226	0.1283
SPECIFIC HETEROSESIS	5	0.3814	0.0763	1.9409	0.0919
GENOTYPE INTERACTION	112	8.5038	0.0759	1.9321	0.0002
GENERAL INTERACTION	32	2.1933	0.0685	1.7441	0.0159
HETEROSESIS INTERACTION	80	6.3106	0.0789	2.0073	0.0002
AVERAGE INTERACTION	8	0.8969	0.1121	2.8530	0.0059
VARIETY INTERACTION	32	2.5540	0.0798	2.0310	0.0029
SPECIFIC INTERACTION	40	2.8596	0.0715	1.8192	0.0064
ERROR	129	5.0694	0.0393		

METHOD 0: DIALLEL ANALYSIS
DESMODIUM DIALLEL ALL BLOCK ANALYSIS
DRY WEIGHT IN A LEONARD JAR (MG/PL)

GARDNER-EBERHART CONSTANTS

ANALYSIS III

DRY WEIGHT IN A LEONARD JAR (MG/PL)

DIALLEL MEANS

PARENTS 1 2 3 4 5

1 1.9534

2 2.1616 1.9861

3 2.2876 2.0767 2.0014

4 2.1821 2.1972 2.2983 1.9470

5 2.2726 2.2637 2.3532 2.2695 2.0415

POPULATION MEAN - 2.1528

S.E. - 0.0512

VARIETY	MEAN	1.9859
CROSS	MEAN	2.2363
AVERAGE	HETEROsis	0.2504
VARIETY	CONSTANTS	
PARENT	CONSTANT	
1	-0.0325	
2	0.0002	
3	0.0155	
4	-0.0369	
5	0.0556	

VARIETY	HETEROsis	CONSTANTS (II) OR GENERAL (III)
PARENT	CONSTANT	
1	-0.0137	
2	-0.0819	
3	0.0236	
4	0.0097	
5	0.0714	

SPECIFIC	HETEROsis	CONSTANTS
CROSS	CONSTANT	
1 X 2	0.0210	
1 X 3	0.0415	
1 X 4	-0.0412	
1 X 5	-0.0213	
2 X 3	-0.1012	
2 X 4	0.0422	
2 X 5	0.0380	
3 X 4	0.0377	
3 X 5	0.0220	
4 X 5	-0.0387	

METHOD 2 DIALLEL ANALYSIS

DESMODIUM DIALEL ALL BLOCK ANALYSIS

DRY WEIGHT IN A LEONARD JAR (MG/PL)

GRIFFING'S COMBINING ABILITY ANALYSIS (MODEL I)

PARENT G.C.A. EFFECTS G.C.A. VARIANCE

		S.C.A. VARIANCE
1	-0.0152	0.0094
2	-0.0351	0.0056
3	0.0145	0.0004
4	-0.0108	-0.0005
5	0.0465	0.0016

S.E. OF THE G.C.A. EFFECTS -
S.E. OF THE DIFFERENCES BETWEEN G.C.A. EFFECTS -

0.0158
0.0250

S.C.A. EFFECTS

PARENTS	1	2	3	4	5
1	-0.1691				
2	0.0590	-0.0966			
3	0.1354	-0.0556	-0.1805		
4	0.0552	0.0903	0.1418	-0.1842	
5	0.0885	0.0995	0.1394	0.0811	-0.2042

S.E. OF THE PARENTAL S.C.A. EFFECTS -

S.E. OF THE CROSS S.C.A. EFFECTS -

S.E. OF THE PARENTAL DIFFERENCES (S.C.A. EFFECTS) -

S.E. OF THE COMMON PARENT CROSS DIFFERENCES (S.C.A. EFFECTS) -

S.E. OF THE DIFFERENT PARENT CROSS DIFFERENCES (S.C.A. EFFECTS) -

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Appendix C. Hayman's analysis of population means computer subroutine.

1: HAYMAN subroutine listing.

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2: HAYMAN sample output.

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SUBROUTINE HAPM(IW,X,V,PMS,IE,EMS,NAME,CH,NC,NV)

C HAYMAN'S ANALYSIS OF POPULATION MEANS

C VERSION 1.0, JUNE 1979

C IW - I/O UNIT FOR WRITE CONTROL (I.E. WRITE(IW,...))

C X - VECTOR OF MEANS (ORDER = P1,P2,F1,F2,B1,B2)

C V - VECTOR OF ST. DEVIATIONS (ORDER = P1,P2,F1,F2,B1,B2)

C PMS - POPULATIONS MEAN SQUARE

C IE - ERROR DEGREES OF FREEDOM

C EMS - ERROR MEAN SQUARE

C NAME - NAME OF THE EXPERIMENT (ALPHANUMERIC 20A4)

C CH - CHARACTER IDENTIFICATION (ALPHANUMERIC 20A4)

C NC - MAXIMUM NUMBER OF 'WORDS' IN 'NAME' AND 'CH'

C NV - 0 = UNWEIGHTED, 1 = VARIANCE WEIGHTED

C REQUIRES SUBROGRAM - FPROB

DIMENSION X(1),V(1),NAME(1),CH(1)

DIMENSION A(6,6),Q(6),P(6),R(6),Y(6)

REAL B(6,6)/1.0,1.0,0.0,1.0,0.0,0.0,1.0,-1.0,0.0,1.0,0.0,0.0,0.0,1.0,
&0.0,1.0,0.0,0.0,1.0,1.0,0.0,0.5,0.0,0.0,0.25,1.0,0.5,0.5,0.25,0.25
&,0.25,1.0,-0.5,0.5,0.25,-0.25,0.25/

REAL POP(6)/*P1', 'P2', 'F1', 'F2', 'BC1', 'BC2'/

PSS=PMS*5.

ESS=EMS*IE

DO 10 I=1,6

10 B(1,I)=X(I)

C WEIGHTING BY STANDARD ERROR

IF (NV.EQ.0) GO TO 30

DO 20 J=1,6

DO 20 I=1,6

B(I,J)=B(I,J)/V(J)

20 CONTINUE

30 CONTINUE

C COMPUTE CORRELATIONS

DO 50 I=1,6

DO 50 J=1,6

TX=0.0

TY=0.0

SX=0.0

SY=0.0

SXY=0.0

A(I,J)=0.0

DO 40 K=1,6

TX=TX+B(I,K)

TY=TY+B(J,K)

SX=SX+B(I,K)**2

SY=SY+B(J,K)**2

SXY=SXY+B(I,K)*B(J,K)

40 CONTINUE

SMX=SX-TX**2/6.

SMY=SY-TY**2/6.

SMXY=SXY-TX*TY/6.

A(I,J)=SMXY/SQRT(SMX*SMY)

A(J,I)=A(I,J)

50 CONTINUE

C COMPUTE MULTIPLE-R SQUARED AND SUMS OF SQUARES

AUP=1.0

R(1)=1.0

DO 110 L=2,6

K=L

IF (L.LT.4) GO TO 70

C FINDS AND ENTERS THE INTERACTION COMPONENT WITH THE

C LARGEST PARTIAL CORRELATION WHEREAS THE MAIN EFFECTS

C ARE ENTERED FIRST DIRECTLY

AM=0.0

DO 60 M=4,6

VC=(A(1,M)*A(M,1))/A(M,M)

IF (VC.LE.AM) GO TO 60

K=M

AM=VC

60 CONTINUE

70 PE=A(K,K)

DO 80 I=1,6

Q(I)=A(I,K)

P(I)=A(K,I)

80 CONTINUE

DO 90 I=1,6

DO 90 J=1,6

A(I,J)=A(I,J)-(Q(I)*P(J))/PE

90 CONTINUE

DO 100 I=1,6

A(K,I)=P(I)/PE

```

100 CONTINUE
R(K)=AUP-A(1,1)
Y(K)=PSS*R(K)
AUP=A(1,1)
110 CONTINUE
C COMPUTE ANOVA COMPONENTS
EPSS=Y(4)+Y(5)+Y(6)
ERSQ=R(4)+R(5)+R(6)
EPMS=EPSS/3
PF=PMS/EMS
PP=FPROB(S,IE,PF)
AF=Y(2)/EMS
AP=FPROB(1,IE,AF)
DF=Y(3)/EMS
DP=FPROB(1,IE,DF)
EPF=EPMS/EMS
EPP=FPROB(3,IE,EPF)
AAF=Y(4)/EMS
AAP=FPROB(1,IE,AAF)
ADF=Y(5)/EMS
ADP=FPROB(1,IE,ADF)
DDF=Y(6)/EMS
DDP=FPROB(1,IE,DDF)
WRITE(IW,130)
WRITE(IW,140) (NAME(I), I=1,NC)
WRITE(IW,140) (CH(I), I=1,NC)
IF (NV.EQ.1) WRITE(IW,150)
C PRINT MEANS
WRITE(IW,160)
DO 120 I=1,6
WRITE(IW,170) POP(I),X(I)
IF (NV.EQ.0) GO TO 120
WRITE(IW,180) V(I)
120 CONTINUE
C PRINT ANOVA TABLE
WRITE(IW,190)
WRITE(IW,200) PSS,PMS,PF,PP,R(1)
WRITE(IW,210) Y(2),Y(2),AF,AP,R(2)
WRITE(IW,220) Y(3),Y(3),DF,DP,R(3)
WRITE(IW,230) EPSS,EPMS,EPF,EPP,ERSQ
WRITE(IW,240) Y(4),Y(4),AAF,AAP,R(4)
WRITE(IW,250) Y(5),Y(5),ADF,ADP,R(5)
WRITE(IW,260) Y(6),Y(6),DDF,DDP,R(6)
WRITE(IW,270) IE,ESS,EMS
RETURN
130 FORMAT ('HAYMAN'S ANALYSIS OF POPULATION MEANS'// '(ADJUSTED FOR M
&EAN EFFECTS)')
140 FORMAT ('0',20A4)
150 FORMAT ('WEIGHTED (STANDARD ERROR) ANALYSIS')
160 FORMAT ('POPULATION MEANS')
170 FORMAT (' ',3X,A4,F10.4)
180 FORMAT ('+',20X,'+',F10.4)
190 FORMAT ('-ANALYSIS OF VARIANCE'// ' SOURCE',8X,'DF',14X,'SS',18X,'
&MS',19X,'F',19X,'P',18X,'RSQ')
200 FORMAT ('0POPULATIONS',4X,'S',SF20.4)
210 FORMAT ('0 ADDITIVE',5X,'1',SF20.4)
220 FORMAT ('0 DOMINANCE',4X,'1',SF20.4)
230 FORMAT ('0 EPISTASIS',4X,'3',SF20.4)
240 FORMAT ('0 ADD-ADD',4X,'1',SF20.4)
250 FORMAT ('0 ADD-DOM',4X,'1',SF20.4)
260 FORMAT ('0 DOM-DOM',4X,'1',SF20.4)
270 FORMAT ('0ERROR',6X,15,2F20.4)
END
FUNCTION FPROB(M,N,X)
C COMPUTES PROBABILITY OF F-STATISTIC X WITH M AND N DEGREES OF
C FREEDOM IN NUMERATOR AND DENOMINATOR RESPECTIVELY.
INTEGER A,B
FPROB=1.0
IF (X.LE.0..OR.M.LE.0..OR.N.LE.0) RETURN
A=2*(M/2)-M+2
B=2*(N/2)-N+2
W=X*M/N
Z=1./(1.+W)
IF (A.NE.1) GO TO 20
IF (B.NE.1) GO TO 10
P=SQRT(W)
Y=0.3183099
D=Y*Z/P
P=2.*Y*ATAN(P)
GO TO 40

```

```
10 P=SQRT(W*Z)
D=0.5*P*Z/W
GO TO 40
20 IF (B.NE.1) GO TO 30
P=SQRT(Z)
D=0.5*Z*P
P=1.-P
GO TO 40
30 D=Z*Z
P=W*Z
40 Y=2.*W/Z
J=B+2
IF (J.GT.N) GO TO 60
DO 50 K=J,N,2
D=(1.+A/(K-2.))*D*Z
IF (A.EQ.1) P=P+D*Y/(K-1)
IF (A.NE.1) P=(P+W)*Z
50 CONTINUE
60 Y=W*Z
Z=2./Z
B=N-2
I=A+2
IF (I.GT.M) GO TO 80
DO 70 K=I,M,2
J=K+B
D=Y*D*J/(K-2)
P=P-Z*D/J
70 CONTINUE
80 FPROB=1.-P
RETURN
END
```

HAYMAN'S ANALYSIS OF POPULATION MEANS
(ADJUSTED FOR MEAN EFFECTS)

EXPERIMENT 1, 1978

COTYLEDON AREA (MM SQUARED)

VARIANCE WEIGHTED ANALYSIS

POPULATION MEANS

P1	29.000	+-	5.0000
P2	40.000	+-	7.0000
F1	41.000	+-	5.0000
F2	39.000	+-	7.0000
B1	34.000	+-	6.0000
B2	41.000	+-	9.0000

ANALYSIS OF VARIANCE

SOURCE	DF	SS	F	MS	F	P	ESQ
POPULATIONS	5	8340.0000	1668.0000	35.4893	0.0000	1.0000	
ADITIVE	1	886.6338	886.6338	18.8645	0.0000	0.1063	
DOMINANCE	1	5008.3594	5008.3594	106.5608	0.0000	0.6005	
EPISTASIS	3	2279.1465	759.7153	16.1642	0.0000	0.2733	
ADD-ADD	1	2152.1638	2152.1638	45.7907	0.0000	0.2581	
ADD-DOM	1	94.0675	94.0675	2.0014	0.1577	0.0113	
DOM-DOM	1	32.9154	32.9154	0.7003	0.4030		
ERROR	614	28858.0000	47.0000		0.0039		

HAYMANS ANALYSIS OF POPULATION MEANS
(ADJUSTED FOR MEAN EFFECTS)

EXPERIMENT 1, 1978

COTYLEDON AREA (MM SQUARED)

POPULATION MEANS

P1	29.0000
P2	40.0000
F1	41.0000
F2	39.0000
B1	34.0000
B2	41.0000

ANALYSIS OF VARIANCE

SOURCE	DF	SS	MS	F	P
POPULATIONS	5	8340.0000	1668.0000	35.4893	0.0000
ADDITIVE	1	5977.6563	5977.6563	127.1842	0.0000
DOMINANCE	1	2185.3013	2185.3013	46.4958	0.0000
EPISTESIS	3	176.8449	58.9483	1.2542	0.2893
ADD-ADD	1	48.3907	48.3907	1.0296	0.3107
ADD-DOM	1	63.9702	63.9702	1.3611	0.2438
DOM-DOM	1	64.4840	64.4840	1.3720	0.2420
ERROR	614	28858.0000	47.0000		

Appendix D. Publications.

STRAIN-SPECIFIC INEFFECTIVE NODULATION OF THE TROPICAL LEGUME DESMODIUM BY RHIZOBIUM¹

B. R. Pinchbeck²

ABSTRACT

The *Rhizobium* strain CB1789 originally isolated from *Desmodium sandwicense* E. Mey produces a strain-specific ineffective nodulation response with several lines of *D. aparines* (Link) D. C. Study of *F*₁ and backcross populations from *D. sandwicense* × *D. aparines* hybrids provided evidence for segregation of a dominant gene at a single locus conditioning ineffective nodulation in *D. aparines*. The *D. aparines* lines expressing ineffective nodulation are used in commerce.

Additional index words: Genetics, Inoculation, N fixation.

INEFFECTIVE nodulation of soybeans [*Glycine max* (L.) Merr.] by *Rhizobium japonicum* (Kirchner) Buchanan has been shown to be controlled by the three dominant genes Rj₂, Rj₃, and Rj₄ (2, 7, 8). Major genes conditioning strain-specific nodulation

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*Commonwealth of Australia identification number.

Table 1. Qualitative interactions of *Desmodium* lines with two *Rhizobium* strains.

Species†	Effectiveness‡	
	CB627	CB1789
<i>D. sandwicense</i> (CPI18156)	E	E
<i>D. sandwicense</i> (CPI18163)	E	E
<i>D. sandwicense</i> (N993)	E	E
<i>D. sandwicense</i> (CSL)	E	E
<i>D. sandwicense</i> (CWF)	E	E
<i>D. sandwicense</i> (CLL)	E	E
<i>D. aparines</i> (CPI23189)	E	I
<i>D. aparines</i> (Q8382)	E	I
<i>D. aparines</i> (CPI18010)	E	I
<i>D. aparines</i> (Desica-1)	E	E
<i>D. aparines</i> (CPI33814)	E	E
<i>D. pringlei</i> (CPI38722)	E	E
<i>D. uncinatum</i> (CPI8990)	E	E
<i>D. sp.</i> (Ecuador)	E	E§

†Accession or line identification. ‡E—effective nodulation (green), I—ineffective nodulation (yellow-chlorotic). §Strain induced chlorotic reaction that appears to be pH sensitive.

have been reported in *Trifolium pratense* L. (6) and *T. subterraneum* L. (3). In preliminary studies with the tropical legume, *Desmodium*, an ineffective nodulation interaction was found for several lines of *D. aparines* (Link) D. C. [*D. intortum* (Mill.) Urb.] when inoculated with *Rhizobium* strain CB1789³ (Table 1). Strain CB627, referred to in Table 1, is a commercial inoculum for *Desmodium* and effectively induced nodulation of all lines studied. Strain CB1789 was isolated from *D. sandwicense* E. Mey and strain CB627 was isolated from *D. aparines*. Strains CB627 and CB1789, members of the cowpea group of *Rhizobium*, are slow-growing; however, strain CB627 is relatively faster growing than strain CB1789 (J.M. Vincent, personal communication).

This study was undertaken to determine the mode of inheritance of the ineffective nodulation interaction between *D. aparines* and Strain CB1789.

MATERIALS AND METHODS

*F*₁ populations were produced from a *D. sandwicense* male-sterile line derived from CPI18156 (5) which was crossed to the *D. aparines* lines CPI23189 and Q8382. Backcrosses were obtained by crossing the *F*₁ hybrid onto the male-sterile line. Crosses between *D. sandwicense* and *D. aparines* produce fully fertile progeny and are considered to be closely allied species (4).

The *F*₂ and backcross plants were grown successively in "modified" Leonard jar assemblies (9) under N-free growth conditions and inoculated with 5 ml of a suspension of strain CB1789. Uninoculated controls and parental genotypes were placed adjacent to the experiment. Eight weeks following planting, the plants were scored for green vs. yellow-chlorotic color induced by N deficiency. Seventy randomly chosen *F*₂ plants were removed from the Leonard jars, inspected for nodulation response and replanted in pots to check their response to N fertilizer.

RESULTS AND DISCUSSION

The nodulation scores of the *F*₂ and backcross plants are presented in Table 2. The ineffectively nodulated plants were extremely stunted and chlorotic at 8 weeks and were similar in appearance to the uninoculated and ineffectively nodulated controls. The roots of the ineffectively nodulated plants exhibited various degrees of nodule formation. The majority of nodules were small, whereas a few were of normal size. All nodules sampled lacked pink pigmentation. When these ineffectively nodulated plants were transferred to pots, all responded to N fertilization.

Chi-square analysis of the segregation data support the hypothesis that the strain-specific ineffective nodulation interaction between lines of *D. aparines* and strain CB1789 is produced by a dominant gene at a

Table 2. Observed nodulation scores and Chi-square probability of *F*₂ and backcross plants inoculated with strain CB1789.

Cross	<i>F</i> ₂		Backcross			
	Nodulation		Chi-square probability 3:1	Nodulation		Chi-square probability 1:1
	Ineffective	Effective		Ineffective	Effective	
<i>D. sandwicense</i> (CPI18156) × <i>D. aparines</i> (CPI23189)	88	36	0.80-0.70	35	36	0.95-0.90
<i>D. sandwicense</i> (CPI18156) × <i>D. aparines</i> (Q8382)	91	24	0.80-0.70	41	32	0.50-0.40
Deviation χ^2	179	60	0.99-0.95	76	68	0.70-0.60
Heterogeneity χ^2			0.30-0.20			0.60-0.50

single locus. These results are similar to those reported for soybeans (2, 7, 8).

The symbols N_1/n_1 are suggested for the designation of this ineffective nodulation gene. The *Desmodium* lines which are effectively nodulated have the genotype n_1n_1 . The ineffectively nodulated lines would be expected to have the genotype N_1N_1 ; however, further experimentation would be required to test for functional allelism between the two *D. aparines* lines studied. The ineffectively nodulated *D. aparines* lines have small seeds while effectively nodulated *D. aparines* lines have larger seeds. The linkage of the N_1 allele with small seed size has not been studied.

One ineffectively nodulated line of *D. aparines* has attracted interest as the commercial cultivar - 'Greenleaf' (1). Ineffective nodulation alleles carried by this cultivar may be at a disadvantage if strain CB1789 were to attain a dominant status in the field.

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