

Full Length Research Paper

Inheritance of fresh seed dormancy in Spanish-type peanut (*Arachis hypogaea* L.): bias introduced by inadvertent selfed flowers as revealed by microsatellite markers control

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Production and seed quality in peanut (*Arachis hypogaea* L.) can be reduced substantially by *in situ* germination under unpredictable rainfed environments. Inheritance of fresh seed dormancy in Spanish x Spanish crosses was studied with two sets of segregating populations, an F₂ population derived from true F₁ hybrids identified with peanut microsatellites markers and other populations (F₂, BC₁P_{1s} and BC₁P_{2s}) from randomly-selected F₁ individuals. In the F₂ population developed with true F₁ hybrids, the chi square test was not significant for the deviation from the expected 3:1 (dormant: non-dormant) ratio. In addition, the bimodal frequency distribution curve with the F₂ population gave more evidence that fresh seed dormancy is controlled by a single dominant gene. The average frequency (48%) of true F₁ hybrids give evidence that deviations from expected ratios in the populations (F₂ and BC₁P_{1s}) developed from non-tested F₁ individuals, is most likely due to inadvertent selfs. This study emphasized the need to identify with molecular markers the cross progenies in self-pollinated crops as peanut before testing for any trait.

Key words: Peanut, true F₁ hybrids, fresh seed dormancy, SSR markers.

INTRODUCTION

More than 94% of world peanut (*Arachis hypogaea* L.) production comes from the rainfed crop grown largely by resource-poor farmers (Dwivedi et al., 2003). In such dry areas, the end of the rainy season is variable and late rains that may occur after peanut maturity can cause *in*

situ germination in peanut. Gautreau (1984) reported significant (20%) pod yield losses with the variety 55 - 437 in field experimentation in Senegal. Martin (1999) found that *in situ* germination may cause more susceptibility to aflatoxin contamination in seeds thus, reducing the seed quality.

The species *A. hypogaea* L. has been divided into two subspecies: *A. hypogaea subsp. hypogaea* and *A. hypogaea subsp. fastigiata*. In the subspecies *A. hypogaea subsp. hypogaea var. hypogaea* (Virginia and Runner

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market types) and *var. hursita*, varieties have long duration cycle and seeds are dormant. While in subspecies *fastigiata* involving *var. fastigiata* (Valencia market class) and *var. vulgaris* (Spanish market type), varieties are early-maturing but generally lack fresh seed dormancy (Krapovickas and Gregory, 1994). Spanish and Valencia varieties are currently the most commonly cultivated peanut varieties in dry areas, particularly in Africa and Asia where the shortening of the rainy season is a paramount constraint. However, these early-maturing varieties lack generally fresh seed dormancy and are prone to *in situ* germination. The growing trend of areas occupied by early-maturing varieties will still increase during the next coming decades since drought is now a worldwide abiotic constraint for peanut production. There is a need to develop short duration peanut varieties having fresh seed dormancy to prevent yield losses due to field sprouting in unpredictable rainfall environments.

During the last decade, a few studies on the inheritance of fresh seed dormancy among Spanish type varieties were carried out. Depending on whether epistasis was detected or not in inheritance of the trait, breeders suggested different strategies of selection. In fact, the conclusions of these investigations were not consistent. Khalfaoui (1991) indicated that duplicated epistasis controls fresh seed dormancy. Nautiyal et al. (1994) found that the trait is quantitatively inherited; whereas Kumar (1999) has reported that the trait is under additive-dominance control. Upadhyaya and Nigam (1999) studied different populations from many crosses and found that seed dormancy in that peanut-type is controlled by a single gene and dormancy allele is dominant. More recently, Ndoye (2001) studied three crosses between Spanish varieties and reported that beyond additive and dominance effects, there is duplicate epistasis in the control of fresh seed dormancy. Phenotyping for fresh seed dormancy in peanut can be reliably carried out at the field (Khalifaoui, 1991) or *in vitro* assay (Upadhyaya and Nigam, 1999; Asibuo et al., 2008). Our previous work gives strong evidence that field test and *in vitro* assay give similar results for fresh seed dormancy (Faye et al., 2009) when seed germination tests are performed under right convenience of humidity, light and appropriated temperatures.

Breeding for new varieties requires development and evaluation of a cross progeny. In self-pollinated crops (for example, cowpea, rice, cotton, common beans and groundnut), the breeding process starts commonly with hand pollination. Hybridization requires laborious manual emasculation and pollination as well as manual removing of non-crossed flowers. In groundnut, some underground uncolored flowers as fertile as external flowers may not be removed by the operator and produce undesirable self-pollinations. Therefore, distinguishing true F_1 hybrids from plants coming from inadvertent selfs is very impor-

tant before deriving F_2 population and subsequent families. In groundnut, distinguishing true F_1 hybrids from inadvertent selfs may be easy to achieve base on plant morphology when dealing with crosses for which parents are much divergent. In contrast, for intrasubspecies crosses (e.g. Spanish x Spanish), true F_1 hybrids are not easily distinguishable from female parent using morphological characteristics.

The purpose of this paper is to study the inheritance of fresh seed dormancy in a Spanish x Spanish cross. In this study, an F_2 population derived from true F_1 hybrids identified using microsatellite markers was compared with an F_2 , a BC_1P_{1S} and a BC_1P_{2S} populations developed without conformity control of F_1 plants.

MATERIALS AND METHODS

Populations' development

During the spring of the year 2006, one of the most susceptible *in situ* germination varieties (Fleur 11) cultivated in the "Senegalese peanut basin" was crossed to a dormant variety (73-30). The female parent Fleur 11 is non-dormant but high-yielding specie since it is very prolific and its underground uncolored flowers are remarkably numerous. The genotype 73-30 is a dormant variety and medium-yielding. Fleur 11 and 73-30 are both early-maturing varieties; they mature at 90 days after sowing (DAS). They have no common ancestor in their pedigree. The crosses were made as suggested by Sudheer and Kumar (1996). At harvest, the putative F_1 seeds were randomly divided into two subsets before processing to the subsequent generations.

One subset was assigned to molecular screen for identifying true hybrid F_1 individuals (described below) using simple sequence repeat (SSR) markers. In that subset, once a plant was identified as true F_1 hybrid (DNA extraction and microsatellite analysis), it was allowed to self-pollinate to give controlled F_2 seeds.

Another subset of putative F_1 seeds was used to develop backcross progeny with each of the parents that is, BC_1P_1 [$73-30 \times F_1$ (Fleur 11 x 73-30)] and BC_1P_2 [Fleur 11 x F_1 (Fleur 11 x 73-30)]. These backcross progenies and the remaining non controlled F_1 seeds were planted at field station (Bambey, Senegal) and allowed to self-pollinate, then selfed backcross between BC_1P_{1S} , BC_1P_{2S} populations and an F_2 population, respectively. These three populations were referred to as non-controlled populations in the present paper.

DNA extraction

A set of seventy eight (78) putative F_1 plants were grown in pots filled with sandy soil along with five plants of each parent in the greenhouse at Regional Center for Studies on the Improvement of Plant Adaptation to Drought (CERAAS) near Thiès, Senegal. At 15 days after emergency, young leaves were harvested from each plant and immediately stored at 4°C in ice before DNA extraction. DNA was extracted from 100 mg of fresh leaves following a slightly modified mixed alkyl trimethylammonium bromide (MATAB) protocol (Risterucci et al., 2000). Briefly, leaves were ground in liquid nitrogen using a mortar and pestle and dissolved in 750 µl of MATAB buffer at 74°C. The samples were incubated for 20 min at 74°C and cooled for 5 min at room temperature. A volume of 750 µl of

Table 1. List of peanut microsatellite markers used to identify true F₁ hybrids.

Name	Primer "Forward" 5'-3'	Primer "Reverse" 5'-3'	Motif
PM50	caattcatgatagttttattcgaca	ctttctctccccaatttga	(TAA) ₄ + (GA) ₁₉
TC6E01	cagcaaagagtcgtcagtcg	gaaagttcacttgagcaaattca	(GA) ₂₉
TC1E05	gaaggataagcaatcgtcca	ggatgggattgaacatttgg	(GA) ₃₀
AC2C05	caaggaagcgtgaattgtag	tgtggactatgcttgcattgtt	(TG) ₁₇
Seq4F10	tgcgaaaccctaactgact	tctatgttgcgccgttgac	(TG) ₇ + (GA) ₈
TC11A04	actctgcatggatggctacag	catgttcggtttcaagtctcaa	(CT) ₁₆ + (CT) ₃₃
TC1D02	gatccaaaatctgccttga	gctgctctgcacaacaagaa	(TC) ₃₀
TC3E05	tgaaagataggttccggtgga	caaaccgaaggaggaacttg	(CT) ₂₆ +(CA) ₇ + (CA) ₅
TC11H06	ccatgtgaggtatcagtaaagaaagg	ccaccaacaacattggatgaat	(AG) ₃₄
TC2D06	agggggagtcaggaaaga	tcacgatcccttctcctca	(AG) ₃₀

Table 2. Chi square value and probability of goodness of fit for a ratio of 3 dormant: 1 non-dormant in the F₂ generation developed from true F₁ hybrids.

Data	Phenotype		Total	Range (days)	Chi square χ^2	Probability value
	Dormant	Non-dormant				
Expected	57	19	76			
Actual	49	27	76	2 - 35	3.40	p > 0.05

chloroform:isoamylalcohol (CIA) (24:1) was added to each sample and all samples were shaken gently until homogenization before centrifugation at 12000 rpm for 20 min. The supernatant was then harvested and the DNA precipitated with 600 μ l of 2-propanol. After centrifugation, pellets were washed with 300 μ l of 70% ethanol, air dried and dissolved in 500 μ l of TE.

Microsatellite analysis

Ten SSRs (PM050, TC6E01, TC1E05, AC2C05, Seq4F10, TC11A04, TC1D02, TC3E05, TC11H06 and TC2D06) polymorphic between the parents were used to identify the true hybrid individuals. The primers used for the identification of true F₁ hybrids are listed in Table 1.

For a given SSR locus, the forward primer was designed with a 5'-end M13 tail (5'-CACGACGTTGTAAAACGAC-3'). Polymerase chain reaction (PCR) amplifications were performed in a MJ Research PTC-100™ thermocycler (Waltham, MA, USA) or in an Eppendorf Mastercycler on 25 ng of DNA in a 10 μ l final volume of buffer (10 mM Tris-HCl (pH 8), 100 mM KCl, 0.05% w/v gelatin, and 2.0 mM MgCl₂) containing 0.1 μ M of the M13-tailed primer, 0.1 μ M of the other primer, 160 μ M of dNTP, 1 U of Taq DNA polymerase (Life Technologies, USA.) and 0.1 μ M of M13 primer-fluorescent dye IR700 or IR800 (MWG, Germany). The touchdown PCR programme used was as follow: initial denaturation at 95°C for 1 min; following by 10 cycles of 94°C for 30 s, T_m (+ 5°C, - 0.5°C/cycle) for 1 min, and 72°C for 1 min. After these cycles, an additional round of 25 cycles of 94°C for 30 s, T_m for 1 min, and 72°C for 1 min and a final elongation step at 72°C for 8 min was performed. IR700 or IR800-labeled PCR products were diluted 7-fold and 5-fold respectively, subjected to electrophoresis in a 6.5% polyacrylamide gel and then sized by the IR fluorescence scanning system of the sequencer (LI-COR, USA). A plant was considered as true hybrid

(H) if it has both alleles of the two parents for all ten primers tested while inadvertent selfs (S) had only the allele of the female parent (Table 2).

Phenotyping for fresh seed dormancy

Maturity of the seeds was assessed by scoring for blackening of the internal inner parenchyma of the pod (Miller and Burns, 1971). In the field test, seeds were treated with a fungicide (Granox) prior the planting.

The non-controlled F₂, BC₁P_{1S} and BC₁P_{2S} populations were phenotyped along with the parents (Fleur 11 and 73-30) using the method described by Khalfaoui (1991) for fresh seed dormancy at the experiment field of the research station (CNRA, Bambey-Senegal) where their mother-plants were cultivated. During the germination test, day mean temperature was 27°C. The soil was kept moist by regular watering.

Eighty seeds randomly sampled from each non-controlled segregating population (F₂, BC₁P_{1S} and BC₁P_{2S}) and from each parental population (Fleur 11 and 73-30) were phenotyped for fresh seed dormancy. The number of seeds that germinated was counted each day. Monitoring continued until all the seeds germinated (35 DAS).

Seventy-six seeds from the F₂ population derived from controlled F₁ plants along with one hundred (100) seeds from each of the two parents were incubated at room temperature (30°C \pm 1) for fresh seed dormancy test. The test was performed in Petri dishes using filter paper moistened with distilled water. Before the test, Petri dishes were washed with sodium hypochlorite (46°). Filter papers were kept moist with distilled water until all the seeds germinated (35 DAS). Seeds that had not germinated at 15 DAS were soaked for 6 h in 2-chloroethylphosphonic acid (10⁻³ M) solution. This compound is known to be effective in breaking seed dormancy and was used to confirm the viability of the dormant seeds. This

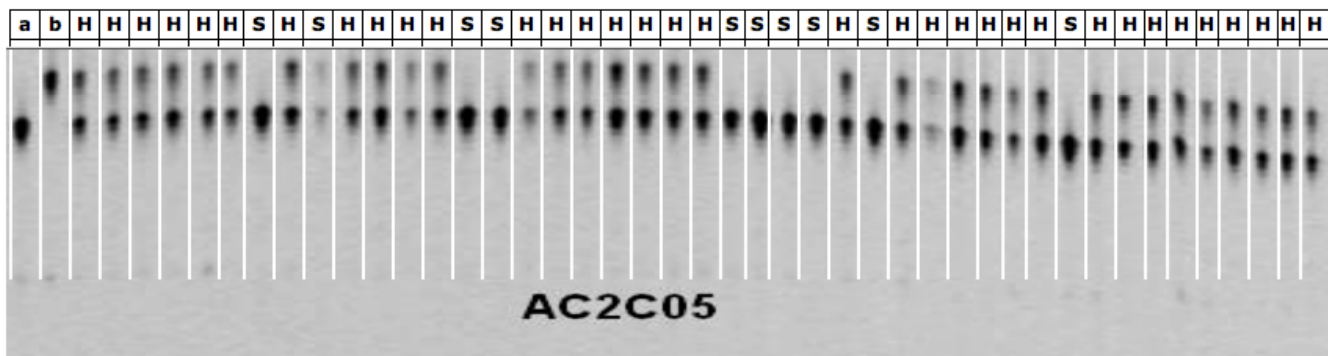


Figure 1. Microsatellite marker survey of a subset of 44 F₁ putative hybrids using primer pair AC2C05. a = Female parent allele (Fleur 11); b = male parent (73-30) allele; H = hybrid; S = inadvertent self.

product, commonly called ethereal, is readily converted to ethylene (Ketring and Morgan, 1971). After treatment with ethereal, treated seeds were returned to room temperature until they germinated for scoring. Compared on the phenotype of the parents, the segregating F₂ seeds were classified as dormant or non-dormant.

Statistical analysis

The chi square (χ^2) test was calculated to examine the goodness-of-fit between the observed and the expected ratios in all the populations at the probability $p=0.05$ level of significance.

RESULTS AND DISCUSSION

The percentage of true F₁ hybrids identified with SSR markers

Of the seventy eight (78) putative F₁ plants tested, thirty eight (38) were true F₁ hybrids, corresponding to a percentage of 48%. Figure 1 set out a profile of bands for detecting true F₁ hybrids with the primer AC2C05. This percentage of true F₁ hybrids indicated most likely that inadvertent selfs were collected at harvest, although caution was taken to remove underground uncolored flowers till 60 days after emergency. The average percentage of true F₁ hybrids found in this study was lower than the percentage (60 - 70%) observed by Gomez et al. (2008). However, percentage of true hybrids in self-pollinated crops depends on the female parent used in the cross since the number of underground flowers may vary from one cultivar to another. Furthermore, the percentage of true-hybrids depends upon the climatic conditions as described by Kotzamanidis (2006) who observed in peanut a lower (16%) percentage of successful crosses under low temperatures. It was however noted that inadvertent selfs could be important for one or another reason in cross progenies of peanut.

Segregation ratios for fresh seed dormancy in the different populations

In the field test as well as in the Petri-dishes test, average day taken before germination was 5 days after sowing (DAS) for the seeds of the non-dormant parent (Fleur 11) while for the dormant parent (73-30), seeds germinated from 11 DAS to 35 DAS. This good level of dormancy of the donor parent 73-30 is a confirmatory of the investigations previously reported by Gautreau (1984), Khalfaoui (1991) and Ndoye (2001). Therefore, in the heterogeneous populations studied here, seeds that germinated within 5 DAS were classified as non-dormant and those that took more days to germinate were classified as dormant.

The chi square test performed on the F₂ population derived from true F₁ hybrids, assuming 3:1 (dormant: non-dormant) ratio was not significant ($P = 0.08$), indicating that the trait is controlled by a single dominant gene (Table 2). These findings agreed with those previously reported by Upadhyaya and Nigam (1999) and Asibuo et al. (2008). In addition, the range of variation (2 - 35 DAS) observed in this study (Table 2) for the dormancy duration is consistent with the observed range of variation by Upadhyaya and Nigam (1999) although they used different parents.

The frequency distribution curve of fresh seed dormancy was bimodal, indicating that the trait can be treated in a qualitative fashion (Figure 2). Figure 2 brought out clearly two statistic modes, one (at 1 - 5 DAS) corresponding to the non-dormant seeds and the second (from 16 - 20 DAS) to the dormant seeds.

This finding corroborated with the segregation ratio that a single gene controls fresh seed dormancy in Spanish x Spanish crosses. Mather (1949), Lynch and Walsh (1998) argued that when the frequency distribution curve of a given trait is bimodal, the trait under study is

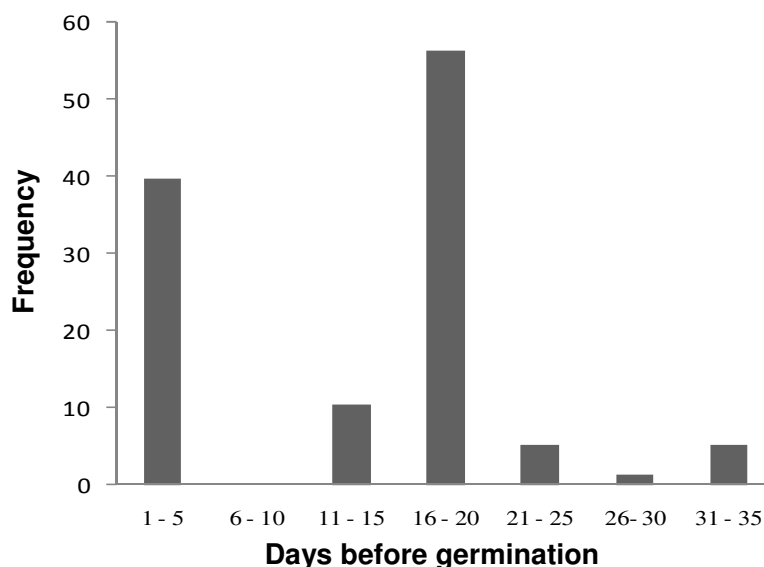


Figure 2. Frequency distribution for fresh seed dormancy of the F₂ individuals (n = 76) derived from true hybrid F₁ individuals.

Table 3. Chi square value and probability of goodness of fit for the expected ratios in the developed populations derived from non tested F₁ hybrids.

Generation	Expected ratio (dormant: non-dormant)	Total	Observed values		Chi square χ^2	Probability value
			Dormant	Non-dormant		
F ₂	3 :1	80	41	39	24.06	p < 0.0001
BC ₁ P _{1S}	7 :1	80	55	25	25.71	p < 0.0001
BC ₁ P _{2S}	3 :5	80	37	43	2.76	p > 0.05

qualitative in nature.

Segregation of F₂ and selfed backcross (BC₁P_{1S} and BC₁P_{2S}) seeds from non-controlled F₁ plants

Although the temperature varied slightly from one study to another; the range of variation for the number of days taken before germination (2 - 35) in the laboratory test as well as in the field test were very similar. Consequently, seeds in segregating generations (F₂, BC₁P_{1S} and BC₁P_{2S}) were classified using the same criterion than in the controlled F₂ population.

Phenotypic data from the populations (F₂, BC₁P_{1S} and BC₁P_{2S}) which were developed from non-tested F₁ individuals were also investigated (Table 3). For the selfed backcross BC₁P_{1S} and BC₁P_{2S}, the chi square test was performed assuming a 7: 1 and 3: 5 (dormant: non-dormant) ratio, respectively. The chi square value was highly significant (P < 0.0001) for the F₂ and BC₁P_{1S} populations in comparison to the expected ratios, but not

for the BC₁P_{2S} population.

In the F₂ population, inadvertent selfs from the putative F₁ plants are probably the major source of the deviation from the expected ratio. The phenotypic data showed thirty nine (39) germinated seeds out of 80 seeds leading to a neat deviation from the expected ratio (3: 1). The expected ratio was most probably biased due to self inadvertent seeds among the putative F₁ self pollinated to develop that F₂ population.

In the selfed backcross BC₁P_{1S} population, the source of deviation could be attributed to inadvertent selfs in the F₁ as well as in the BC₁P₁ [73-30 x F₁ (Fleur 11 x 73-30)] because the recurrent parent 73-30 was used as female parent (Table 3). Therefore, the source of deviation was most likely caused by the bias in the F₁ generation. Although much care was taken to remove them, abundant underground uncolored flowers observed in the female parent induced a shift in the expected ratios between dormant and non-dormant seeds.

In contrast to the F₂ and BC₁P_{1S} populations in the BC₁P_{2S} population, the chi square test was not significant

($p > 0.05$); that means that actual phenotypic data fitted the expected ratio (Table 3). Good fit between observed and expected ratios in that population could be explained by mere chance since the sampling of putative F_1 used to develop the subsequent generations was at random. So the ratio may depend upon the percentage of self inadvertent plants collected at the harvest of the cross progeny, particularly the F_1 generation.

The results of the present study revealed that deviation from the expected proportions in non-controlled populations (F_2 and BC_1P_{1S}) is most likely due to self inadvertent selfs during the populations' development. In the context of the contradictory conclusions about the inheritance of fresh seed dormancy in peanut among Spanish crosses, this may be prominently a consequence of the presence of inadvertent selfs in phenotyped populations. In our knowledge, except Upadhyaya and Nigam (1999), most of the few available previous reports on the inheritance of fresh seed dormancy among crosses of Spanish-type peanut (*Arachis hypogaea*, L.) did not mention precautions used to discard these problematic inadvertent selfs from populations under study. In fact, inadvertent selfed flowers may be one of the causes of the misleading conclusions reported on the inheritance of fresh seed dormancy among Spanish-type varieties.

Beside the inadvertent selfs, various non-genetic factors such as environmental factors that have been extensively studied by Toole et al. (1964) and recently reviewed by Finch-Savage and Metzger (2006) could obscure the phenotyping work. However, the average to high heritabilities observed in this and other studies (Khalfaoui, 1991; Ndoye, 2001) indicated that environmental factors can be overcome.

Conclusion

This study has given more evidence that fresh seed dormancy in Spanish x Spanish crosses is controlled by single dominant gene. Therefore, fresh seed dormancy in Spanish varieties is qualitative in nature. Pedigree selection from an F_2 population could be suggested to be an effective strategy to obtain peanut lines with earliness and fresh seed dormancy. The variety 73-30 could be used as donor parent in breeding programs.

This work has outlined the importance of microsatellite markers for identifying true F_1 hybrids in cross progenies in self-pollinated crops. Since then, many SSR markers are now published in peanut. Future work will be the identification of SSR markers linked to the gene controlling fresh seed dormancy in Spanish-type peanut by using a bulk segregation analysis approach.

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