



## Development of HY1 hybrid embryos between a cultivar of *Phaseolus vulgaris* L. and a wild genotype of *Phaseolus coccineus* L.

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### ABSTRACT

Reciprocal crosses between *Phaseolus vulgaris* and *P. coccineus* frequently lead to embryos abortion in all the developmental stages. However, the *P. vulgaris* NI637 cultivar and the wild form NI1108 of *P. coccineus* present abilities for combination. The abortion rate of pods is 93.93% when NI1108 is the maternal parent against 43.49% when it is the pollinator. The technique of histological resin sections of 2-Hydroxyethyl Methacrylate (HEMA) allows the analysis of the evolution of self-pollinated embryos and 3-14 days old hybrids after pollination. The growth rate does differ between the two types of embryos. On the basis of developmental stages, the form of embryo on one hand, and the ovary tissues on the other hand, present several specificities. The self-pollinated embryos regularly developed according to the initial polarity apex-base. In NI1108(♀)xNI637 hybrid embryos, the suspensor is hypertrophied over the embryo proper. These embryos of small size, present a slow growth compared to the self-pollinated ones and the NI637(♀)xNI1108 hybrids. These observations will allow the identification of the stages concerned by more abortions, explain their possible causes, and describe the expression of abortion symptoms in hybrid embryos during the embryogenesis within the *Phaseolus* genus, in view of improving the common bean production.

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**Key words:** Plant embryos, interspecific hybridization, abortion, suspensor

### INTRODUCTION

It can be useful to utilize interspecific hybridization for improving the genetic of common bean. However, crosses realized within the complex *P. vulgaris* Lam. – *P. polyanthus* Greenman – *P. coccineus* L. have revealed incompatibilities barriers, especially post-zygotic (Baudoin et al., 2004), and limiting the rating success of these crosses (Sabja et al., 1990; Debouck and Smartt, 1995). Symptoms differ according to species and the biological status of crossed forms (Geerts et al., 2002). When *P. polyanthus* is

pollinated by *P. vulgaris*, crosses often lead to embryo abortion at the early stage of embryogenesis (Geerts et al., 2002) following incoherent development between embryo and endosperm (Russell, 1993; Lecomte et al., 1998; Olsen et al., 1999). During crosses between *P. coccineus* (♀) and *P. vulgaris*, the abnormal development of endosperm and the important requirement of nutrients by embryos to the huge suspensor may be responsible to the abortion cases observed (Perata et al., 1990; Yeung and Meinke, 1993; Pullman and Buchanan, 2003; Nguema

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Ndoutoumou et al., 2007). The abortion of embryo is initiated before the cotyledonary stage (Sallandrouze et al., 2002), and their first signs are seen within the mother tissues (Sage and Webster, 1990), whatever the developmental stage of the embryo.

It is usually observed that starch is altered (Chamberlin et al., 1994), tegumentary and nucellar cells are vacuolated and the cell wall of the middle layer of the teguments are deformed or degenerated (Lecomte et al., 1998). This often leads to abnormal embryo formation (Pullman and Buchanan, 2003). Interactions between the embryo, endosperm and mother tissues involved in embryo growth may be the main cause of the failures observed on the next generation embryos. The identification of abortion in relation to embryo growth contributes to a better determination of the adequate moment of immature embryo rescue. The early examination of the embryogenesis through the histological analysis is necessary to describe the self-pollinated hybrid embryos in order to understand the abortion of hybrid embryos.

#### **MATERIAL AND METHODS**

The wild form (NI1108) of *P. coccineus* and the NI637 cultivar of *P. vulgaris*, respectively originated from Mexico and Brasilia were chosen for their aptitude to flowering under our experimental conditions, their biological status and geographical origin, the genetic distance between species and their aptitude to hybridization (Hoover et al., 1985). They were grown in green house under the following parameters: a temperature day/night of 24 °C and 20 °C, a relative humidity of 75%, an irradiation of 170  $\mu\text{mol. m}^{-2}.\text{s}^{-1}$  and a photoperiod of 12h. Pollinations were realized, according to the method of Buishand (1956). Pods destined to histological studies were daily collected from the 3<sup>rd</sup> to 14<sup>th</sup> day after pollination (DAP) and originated from self-pollination, or reciprocal crosses between *P. vulgaris* and *P. coccineus*. Ovules were fixed in a standard solution using 1.25% glutaraldehyde, 4% paraformaldehyde and cleaned thrice successively with a mixture of  $\text{Na}_2\text{HPO}_4.2\text{H}_2\text{O}$  (0.3M) and  $\text{NaH}_2\text{PO}_4.2\text{H}_2\text{O}$  (0.3M) and maintained at pH 6.6. Dehydration was conducted in six baths of increased ethanol gradient (30%, 50%, 70%, 90%, 95% and 100%). The pre-infiltration was realized

in a mixture of absolute ethanol and pure Technovit™ 7100 resin, followed by infiltration and embedding of objects. Longitudinal sections of 3 $\mu\text{m}$  were carried out on the axe of the embryo using a rotative microtome (Microm H360). They were then stained with toluidine (Gutmann, 1995) and examined under a Nikon Eclipse E800 microscope. Pictures were snapped using a Nikon Digital Sight DS-U1 camera and treated with the Lucia picture computer program. After crosses, pods were collected and ovules analyzed histologically and recorded. The abortion rate of pods was calculated by the following formula: number of aborted pods x 100 / number of pollinations. The length of suspensor and embryo were measured on self-pollinated samples, 64 hybrid ovules NI1108 (♀) x NI637 and 82 hybrid ovules NI637 (♀) x NI1108. Statistical analysis was performed using Minitab 1.4 computer program followed by the Tukey test to segregate between the means at 5% level of significance. The development of embryo structures of parents and hybrids was compared from 3 to 14 days after pollination.

#### **RESULTS**

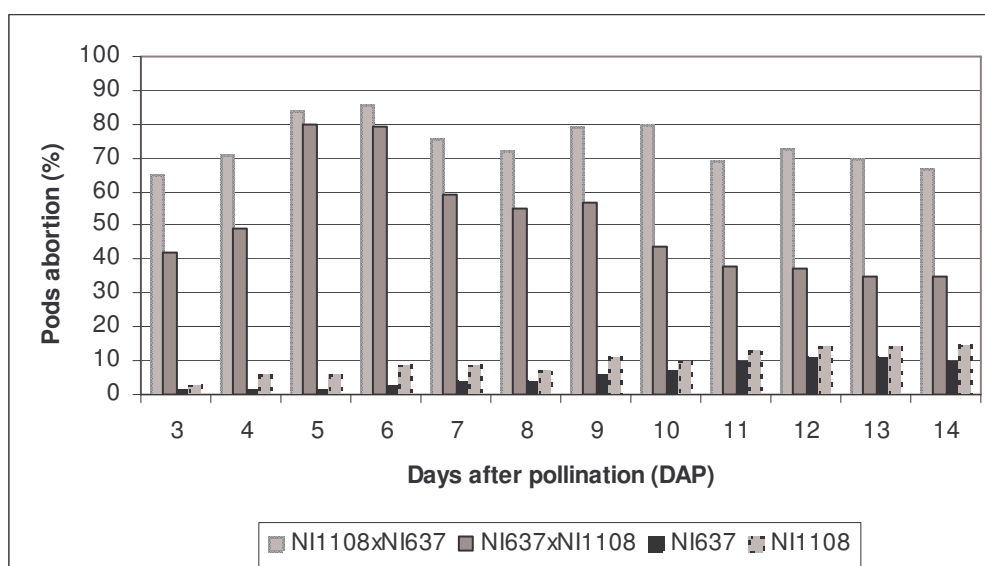
##### **Self and cross-pollinations**

During the course of observation, the outer signs of embryos abortion are expressed initially by the slowness of pods length growth, a softening and yellowish, followed by a decline of tissues, and the abscission of pods. Table 1 indicates the number of self-pollinations and crosses effected, as well as the abortion rate of pods at 14 DAP. The abortion rate of pods was lower in the autogam genotype NI637 compared to allogam genotype NI1108, and the interspecific crosses. During hybridizations, the abortion rate of pods was greater (93.93%) in NI1108 (♀) x NI637 combination than the reciprocal cross (43.49%). Figure 1 presents the evolution of pods abortion rate between the 3 and 14 DAP, depending on the pollination type. Abortions of self-pollinated pods gradually increased from the 3<sup>rd</sup> to 14<sup>th</sup> DAP, but remained very limited. In contrast, for reciprocal crosses, the rate was more elevated and reached a maximum between 5 and 6 DAP. After 7 DAP, the abortion rate

**Table 1:** Number of pollinations, mature pods and rate of pod abortions.

Genotypes and combinations	Number of pollinations	Number of mature pods obtained	Rate of aborted pods (%)
NI 637 (PV)	121	106	12.39
NI1108 (PC)	287	198	31.01
NI637 (♀) x NI1108	361	204	43.49
NI1108 (♀) x NI637	659	40	93.93

NI = Identification Number ; PV = *P. vulgaris* ; PC = *P. coccineus*



**Figure 1:** Rate of pods abortion from 3 to 14 DAP during self-pollination and reciprocal crosses between *P. coccineus* and *P. vulgaris*.

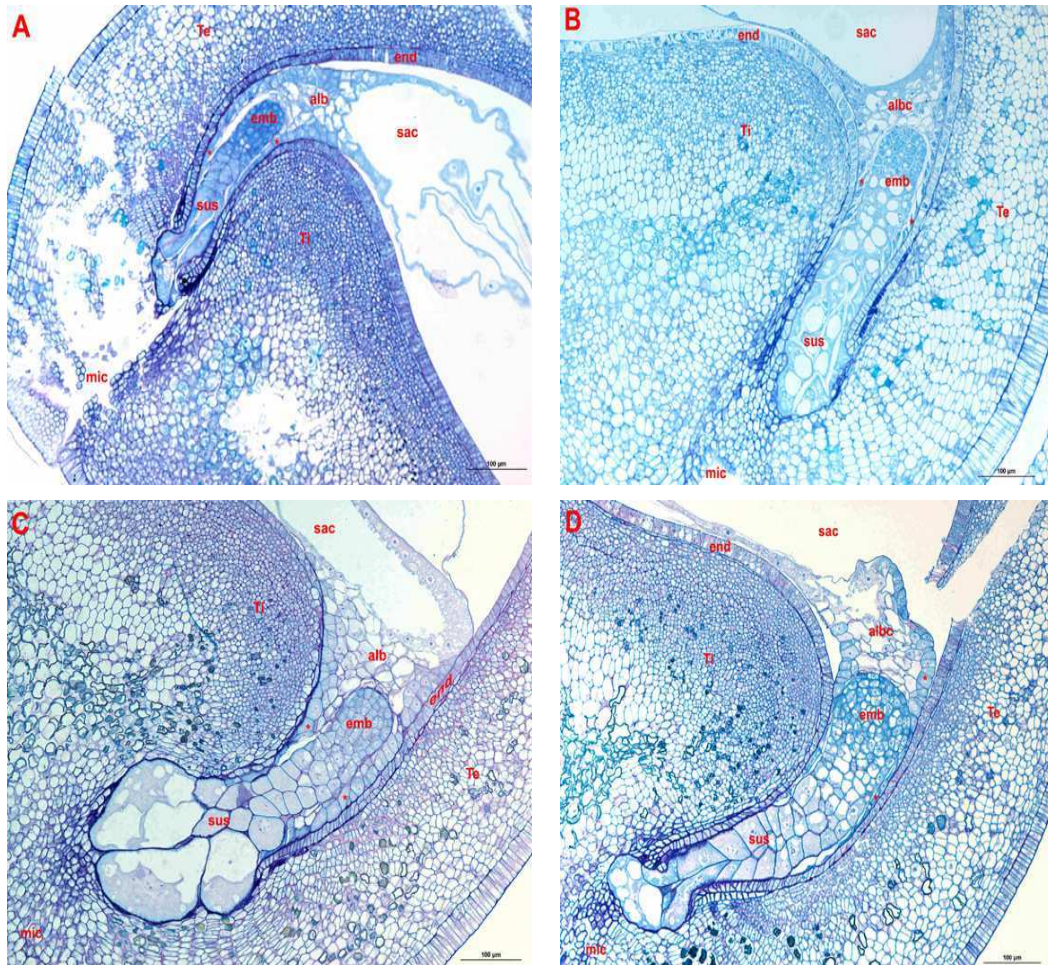
decreased in hybrids, while remaining constant in NI1108 (♀) x NI637.

#### Development of embryos

The evolution of the average length of the main embryo structures (suspensor and embryo), maternal parent (NI1108 and NI637) and reciprocal hybrids described by Nguema Ndoutoumou et al. (2007) revealed a progressive growth of the suspensor length in all the embryos. No significant difference was observed between the means. As far as the evolution of the average length of embryos is concerned, the genotype NI637 showed the mean values statistically higher ( $p < 0.05$ ) than those of NI1108 and the reciprocal hybrids, as from 10 DAP.

Figures 2 and 3 illustrate respectively the maternal genotypes embryos (NI637, NI1108) and reciprocal hybrids at the globular and heart stage of their development. The

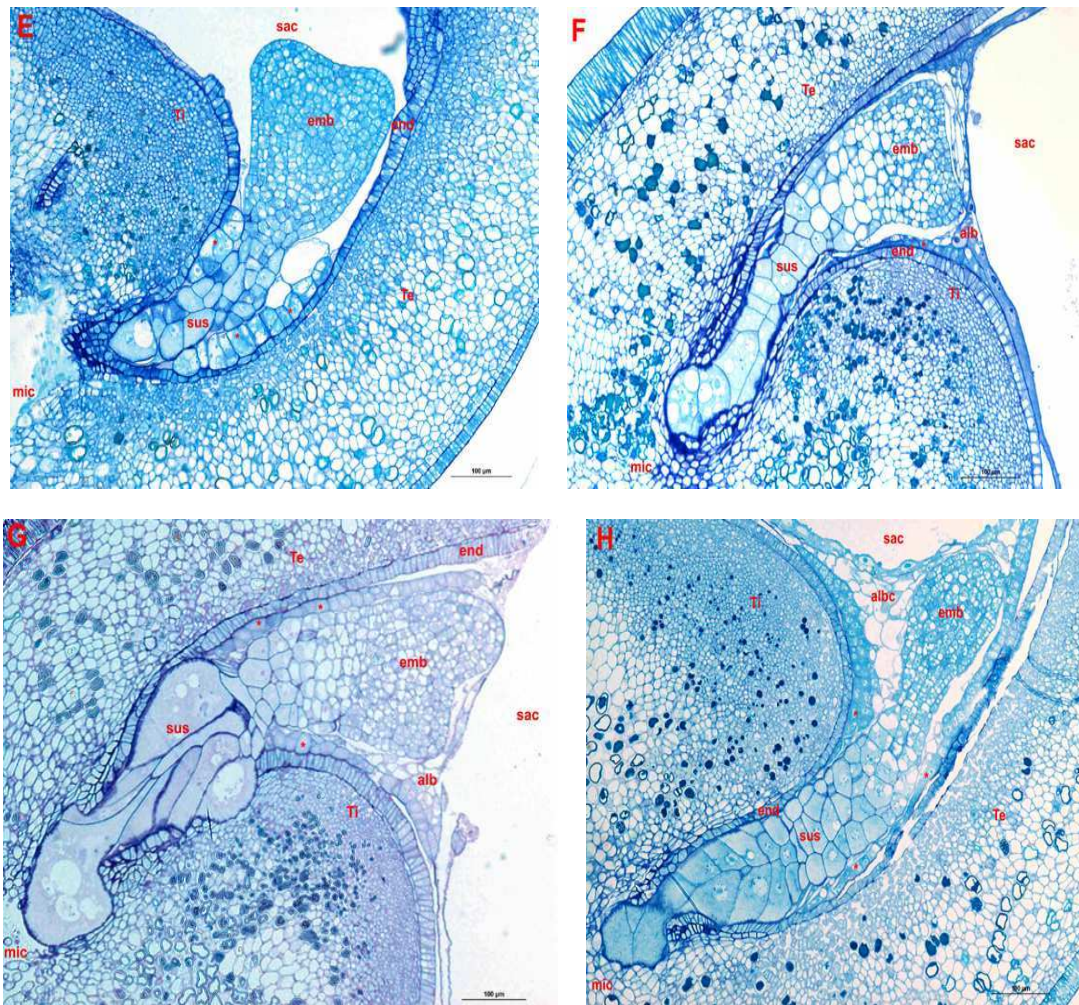
globular stage was observed averagely at between 4 and 7 DAP in the maternal genotypes (NI637 and NI1108), compared to 8 DAP for hybrids NI1108 (♀) x NI637 and 7 DAP for hybrids NI637 (♀) x NI1108. The embryos of the maternal genotypes had a fusiform suspensor (Photo A & B), while hybrid embryos showed big cells at the basis of suspensor (Photo C). In the case of hybrid embryo NI1108 (♀) x NI637, the suspensor was hypertrophied at the cost of embryo proper. In all the self-pollinated ovules or hybrids, the endosperm cells persist in the neighbouring of the embryo proper, inside the embryo sac. The highly albuminated and nucleated cells border the endothelium, suggesting a nutrients transport from the endothelium to the embryo and the suspensor. In NI637, the heart stage was initiated toward the 5 DAP; while the embryo reached the heart stage later on at 7 DAP. The NI1108



**Figure 2:** Longitudinal median sections of globular embryos of *P. vulgaris* (NI637), *P. coccineus* (NI1108) and reciprocal hybrids. Parental embryos NI637 (Photo A) and NI1108 (Photo B) have a fusiform suspensor (sus) in micropylar end (mic). Basal cells are filled out within reciprocal *P. coccineus* x *P. vulgaris* embryos (Photo C & D). The endosperm (alb) separates embryo sac (sac) from the embryo proper (emb). The transfer cells (\*) develop between the endothelium (end) covered by inner and outer integuments (Te & Ti) and the endosperm in the one hand, and between the embryo proper and the suspensor in the other hand.

embryo and NI637 (♀) x NI1108 were at the heart stage at 9 DAP, while the hybrid embryo NI1108 (♀) x NI637 was at the heart stage at 10 DAP. A restricted zone links the suspensor and hypertrophied cells of the hybrid embryo to the embryo proper (Photo G). This may explain the nutrient deficiency of the embryo *via* the suspensor. The endothelium was intact around the embryo. The cellular endosperm was totally reabsorbed in the ovule of NI637 contrary to other cases. The embryo may get its nutrients preferentially through the links established between the endothelium and the endosperm. Later, the hybrid embryos reached

the cotyledonary stage at 12 DAP, but this stage was attained at 9 DAP in NI637 and 11 DAP in NI1108. In hybrid embryos, the suspensor drove in teguments, where it develops important invaginations useful for embryo nutrition. The contact between the endosperm and cotyledons was established for the same reason. The endosperm disappeared in the neighbouring of the embryo proper, but was developed quite in part of the embryo sac. The endothelium was crushed under the pressure of basal cells of the suspensor. The transfer cells were seen in the contact points between the endothelium and the embryo, the



**Figure 3:** Longitudinal median sections of heart stage of embryos from *P. vulgaris* (NI637), *P. coccineus* (NI1108) and reciprocal hybrids. Within NI637 (Photo E) and NI1108 (Photo F) embryos, the suspensor (sus) is slender. Within reciprocal crosses *P. coccineus* x *P. vulgaris* (Photo G & H), the suspensor is enormous compared to the embryo proper (emb). It is anchored into teguments (Ti & Te) near to the micropyle (mic). A thin layer of albuminated cells (alb) delimits the embryo from the embryo sac (sac). The transfer cells (\*) are in the extension of the endothelium (end) within self-pollinated ovules.

suspensor and the endosperm. The intracellular thicknesses were also observed in the endothelium and the tegumentary layers, with numerous grains of starch.

#### DISCUSSION

Embryo abortion arises from self-pollination and during hybridization between *P. vulgaris* and *P. coccineus*. In the first case, this is often explained by intrinsic phenomena in plants, in accordance with phonologic stages. In the second case, barriers raise at all

the developmental stages of the hybrid embryo, despite the aptitude to combination suggested between the cultivated species of *P. vulgaris* and the wild form *P. coccineus*. Several turgescer ovules hybrid are deprived from living embryos. In addition to the biological form of crossed genotypes, the direction of cross also determines the success of interspecific hybridization between *P. vulgaris* and *P. coccineus*. Crosses are easier when *P. vulgaris* is the maternal parent. During these crosses, a great number of

abortions occur 5 to 6 DAP. It is also a critical period in the development of hybrid embryos between *P. vulgaris* (♂) and *P. coccineus*. Similar observations were reported during crosses between *P. vulgaris* and *P. polyanthus* (Yeung and Meinke, 1993; Lecomte et al., 1998; Geerts et al., 2002). Like other authors (Sage and Webster, 1990; Lecomte et al., 1998; Nguema Ndoutoumou et al., 2007) observations were made that incompatibility barriers in *Phaseolus* were post-zygotic and were interpreted by embryo abortion for nutritional reasons essentially. During the early embryogenesis, the interactions between the suspensor, the embryo proper, the endosperm and the maternal parent arise during the determination of embryos growth. The embryo genotype NI637 of *P. vulgaris* is developed faster than that of *P. coccineus* (NI1108) and those of the reciprocal hybrids. The influence of the genotype NI1108 may be preponderant in the dynamic of embryogenesis of reciprocal hybrids. Whatever the direction of crossing, the divisions of zygote hybrid is initiated later compare to the maternal embryos and lead to delay development of hybrid embryos. Similar results were obtained during crosses between *P. vulgaris* and *P. polyanthus* (Lecomte et al., 1998; Geerts, et al., 2002). In the maternal parent genotypes, the nutrition of embryo is organized early, as from the globular stage. The nuclear endosperm, in contact with the proembryo and the endothelium, is partitioned off.

The transfer cells were seen between the endothelium and the endosperm on one hand, and between the embryo and the suspensor on the other hand (Figure 2). On serial sections, the links are well seen at advanced stages. In hybrid ovules, no direct contact exists between the embryo proper and the endothelium. A thin layer of the transfer cells is seen between structures. The contact is narrow between the endothelium and the basis of suspensor. The endosperm borders the apical part of embryo and touches the upper part of the suspensor. The transfer cells link the base of the suspensor to endosperm and the body of suspensor to endothelium. This reveals the difficulties of providing embryo with nutrients. The suspensors of maternal embryos evolve more slowly after the late globular stage. The embryo is provided with

nutrients *via* the contacts between cellular endosperm and the body of the suspensor, and between the embryo proper and the endosperm. The suspensors and hybrid embryos develop important invaginations toward the micropyle end (Figure 3), suggesting an important nutritive demand (Maheshwari, 1950; Yeung and Sussex, 1979; Lecomte et al., 1998; Nguema Ndoutoumou et al., 2007). The embryo proper was supposed to prevent the development of suspensor at this stage, but this later is instead hypertrophied in hybrid embryo NI1108 (♀) x NI637. This slow the embryo development since the size of suspensor influences the nutritive requirement of embryo.

Generally, a nutritive transfer between endosperm and cotyledons is possible due to contact zones in cotyledons, at cotyledonary stage. In hybrid NI1108 (♀) x NI637, the observed abnormalities in ovules in the previous stage might provoke a reduction of certain activities responsible for embryo development. When endosperm is not developed enough, the embryo is provided with very little nutrients, the alteration of the maternal or embryo structures is unlatched, and consequently, the embryo abortion occurs.

This work shows that histological differences between maternal tissues in reciprocal crosses could thus be a key factor in the abortion processes. In regard to several factors implied in embryo abortion in both crosses, it is envisioned to rescue the hybrid embryos at the globular stage of their development according to elevated frequencies of abortion occurring in this developmental stage. Hybrid embryos rescued by *in vitro* culture could be the best way of exploration taking into consideration for the developmental stage reached by the embryo and the nutritional needs specific to them. It is also important to proceed by other interspecific combinations to reinforce these results.

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