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Full Length Research Paper

# Studies on interspecific hybridization in *Cyamopsis* species

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Reproductive characters of three species of Cyamopsis were studied to find out barriers to interspecific crosses between Cyamopsis tetragonoloba x Cyamopsis serrata and C. tetragonoloba x C. senegalensis which may serve as a stepping stone for development of extra early varieties of guar. Pollen grains of C. tetragonoloba and C. senegalensis showed more than 95% of viability while those of C. serrata had 87% viability. Nutritive requirement for in vitro germination of pollen revealed that pollen of C. tetragonoloba required 25% sucrose + 100 ppm boric acid + 300 ppm calcium nitrate while C. senegalensis pollen needed 35% sucrose with same basal medium. On the other hand, C. serrata pollen required 35% maltose + 6% PEG 6000 along with above dose of boric acid and calcium nitrate. Moreover, pollen germination in C. serrata was initiated after 30 h of incubation and its pollen tubes were slow growing attaining 174.7 µm length in 48 h. The length of style of C. tetragonoloba and C. serrata was nearly identical (2.6 mm) while C. senegalensis possessed longest style (3.8 mm). Interspecific hybridization between C. tetragonoloba x C. serrata was successful through the use of stub smeared with pollen germination medium (PGM) and as a consequence 10.43% of pod setting was observed. Colour and shape of hybrid seeds was similar to the female parent (C. tetragonoloba), hybrid plants showed early flowering just like male parent (C. serrata) whereas the plant height was intermediate between the two parents.

Key words: Interspecific hybrids, pollen, in vitro germination, stub pollination, in vivo tube growth.

#### INTRODUCTION

India and Pakistan are the main producers of cluster bean, accounting for 80% production of the world's total, while Thar, Punjab Dry Areas in Pakistan and Rajasthan occupies the largest area (82.1%) under guar cultivation in India. In addition to its cultivation in India and Pakistan, the crop is also grown as a cash crop in other parts of the world (Pathak et al., 2010). In India, 3.34 million hectares of the farmable land was under guar cultivation during the year 2006/07 (Ministry of Agri. and Co-op GOI, 2010). It is cultivated in arid zones of Rajasthan, some parts of Gujarat,Haryana and Madhya Pradesh. The productivity of guar ranges from 474 kg/ha in Rajasthan to 1200 kg/ha in Haryana. The most important growing area centres on Jodhpur in Rajasthan, India where demand for guar for fracking produced an agricultural boom as of 2012 (Gardiner 2012).

Guar grows well under a wide range of soil conditions and is tolerant of low fertility, soil salinity and alkalinity. It performs best on fertile, medium-textured and sandy loam alluvial soils but does not tolerate heavy black soils

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(Wong and Parmar 1997).

In the recent past, guar cultivation has become an attractive option with the farmers due to availability of high yielding varieties with high gum (30 to 35% of whole seed) content (galactomannans) in its endosperm which has great value as an enhancer of viscosity in food industry, like stiffner in soft ice-cream, a stabilizer for cheese, instant pudding and whipped cream substitutes and as a metal binder. It is widely used from paper and cosmetic to mining and explosive industry (Whistler and Hymowitz, 1979). Al-Hafedh and Siddiqui (1998) reported that the C. *tetragonoloba* L. beans had 32.81% crude proteins, 3.18% crude fats, 4.19% ash and 10.87% crude fibers.

Guar meal contains about 12% gum residue (7% in the germ fraction and 13% in the hulls) (Lee et al., 2005), which increases viscosity in the intestine, resulting in lower digestibilities and growth performance (Lee et al., 2009). Its uses in tissue culture media as a gelling agent has also been reported (Jain et al., 2005). *Cyamopsis tetragonoloba* L. is a well-known traditional plant used in folklore medicine.

A critical requirement for crop improvement in general, is the introduction of new genetic material in the cultivated lines of interest, whether through conventional or non-conventional breeding or in-vitro techniques. C. tetragonoloba, an erect herb with indeterminate growth and broad trifoliate leaves, matures in 80 to 120 days. However, one of its wild relatives, that is, *C. serrata* is an extra early maturing (40-50 days), slow growing with narrow trifoliate leaves while the other wild species, that is, C. senegalensis is also slow growing with narrow pentafoliate leaves and matures in 120 to 130 days (Menon, 1973). Both these wild relatives possess some desirable attributes like drought resistance, photo-and thermo-insensitivity and disease resistance. Interspecific hybridization among the C. tetragonoloba and its wild relatives is anticipated to produce hybrid with trait of early maturity, disease resistance and photo-and thermoinsensitivity.

According to Harlan and De Wet (1971) gene pool concept, *C. tetragonoloba* is included in primary gene pool (GP-1) while *C. serrata* and *C. senegalensis* are included in secondary gene pool (GP-2). According to them species included in GP-2 can be crossed with GP-1 with some fertility in F1's and thus gene transfer is feasible.

Unfortunately, conventional plant breeding technique has so far failed to yield desired results (Mathiyazhagan, 2009). Such a failure may be due to presence of preand/or post-fertilization barriers. To combat such barriers, it is essential to have detailed knowledge of reproductive biology of all the species of *Cyamopsis* in question. It was therefore, contemplated to follow a systematic approach to identify pre- as well as post- fertilization barriers in interspecific crossing of *Cyamopsis*. Supplementing conventional plant breeding with unconventional less popular methods along with plant biotechnological techniques is anticipated to go headway in resolving the issue. Present investigation was thus undertaken to study some relevant reproductive characters in three different species of *Cyamopsis* and work out cross-ability among these by conventional and unconventional less popular methods.

#### MATERIALS AND METHODS

Plants of three species of *Cyamopsis* viz. *C. tetragonoloba* cv. HG563, *C. serrata* and *C. senegalensis* were raised in cemented pots (from authenticated seeds collected from forage department of CCS HAU, Hisar) in the screen house of the Department of Botany and Plant Physiology. Pots were filled with mixture of soil and farm yard manure. Before sowing, the seeds of uniform size were soaked in liquid broth of Rhizobium strain 1305 for 10 min. Five seeds were sown in each pot at uniform depth and distance. After 25 days of sowing thinning was done to leave three plants of uniform size in each pot. Irrigation with canal water was given as and when required. At flowering, the following reproductive characters of each species were recorded.

#### Number of pollen/flower

Flower buds from each species were collected a day before anther dehiscence and employed for quantification of pollen grains produced per flower. For this, twenty anthers were suspended in 2 ml of 50% diluted glycerine (50% V/v glycerine with water) containing a few drops of safranine. Anthers were crushed with the help of glass rod and the suspension was passed through a brass sieve with a mesh of 48 sq/cm<sup>2</sup> (Kapoor and Nair, 1974). Number of pollen grains per flower drop was counted by haemocytometer.

#### Pollen viability and In vitro pollen germination

Viability of pollen grains was assessed by 2, 3, 5 triphenyl tetrazolium chloride (TTC) test (Hauser and Morrison, 1964). Flower buds were collected from 3 randomly selected plants in the early morning (6.30 a.m.) on the day of anthesis and pollen of these floral buds was mixed thoroughly on glazed paper and used immediately for viability test and in vitro germination on the semi solid medium contained in petri dishes. Preliminary studies revealed that sugar type and its concentration and other adjuvants required for pollen germination varied with species. After preliminary trials, germination medium consisting of 25% sucrose (for C. tetragonoloba), 35% sucrose (for C.senegalensis), 35% maltose +6% PEG 6000 (for C. serrata) along with 100 ppm boric acid, 300 ppm calcium nitrate and 0.8% agar were used for in vitro germination and tube growth. After pollen inoculation, Petri plates were incubated at 30±2°C for 4 h in dark in a BOD incubator with three replicates per treatment. However, inoculated Petri plates of C. senegalensis and C. serrata were incubated for 30 and 48 h respectively. After pollen germination, the pollen activity was terminated by flooding the surfaces of the media with killing and fixing solution of following composition (Sass, 1951): Formaldehyde = 5 ml; Glacial acetic acid = 3 ml: Water = 72 ml: Glycerine = 20 ml.

Pollen producing a tube length of a size greater than double of its diameter was designated as germinated. Twenty readings for pollen germination and thirty for tube length from different microscopic fields of each petri plate were made from area with uniform distribution of pollen and fairly good population.

#### Pistil and yield related characters

The above collected flower buds were used to record shape of the stigma and length of the ovary and style by micrometry. Pistils were cut open under a stereoscopic microscope and number of ovules per pistil from at least twenty pistils was recorded. At maturity, thirty pods from each species were collected randomly and used to measure length and breadth of pods, number of seeds per pod and test weight of 100 healthy; uniform sized seeds from each species was recorded in three replicates per species were used.

#### In vivo pollen tube growth

Self pollinated pistils from flowers of three species of *Cyamopsis* were collected at 24, 48 and 72 h of anther dehiscence and fixed in acetic alcohol (Acetic acid : Ethanol; 1 : 3) for 4 h and processed for aniline blue test (Dumas and Knox, 1983). The observations for germination of pollen grains on stigmatic surface and extent of tube growth in stylar tissue and penetration of the ovule by tube were made under florescent microscope. Fifteen random pistils for each species were used for these studies.

#### Biochemical composition of stigma + style

#### Total soluble carbohydrates

The total soluble carbohydrate content (mg g<sup>-1</sup>FW) was estimated by the method of Yemm and Willis (1954).

#### Extraction

Extraction of soluble carbohydrates was done according to Barnett and Naylers (1966) procedure. 50 mg of fresh material of stigma + style was finely ground in 80% alcohol by using pestle and mortar. Total soluble carbohydrates were extracted in 2 ml of 80% ethanol (v/v) on a water bath at  $50\pm1^{\circ}$ C for 15 min. It was then cooled and centrifuged at 5000 x g for 5 min. The supernatent (extract) was kept aside and the pellet re-extracted twice with 80% ethanol. Total volume of extract was made to 5 ml with 80% ethanol. This extract was used for the analysis of total soluble carbohydrates while the pellet was used for extraction of the total soluble proteins.

#### Reagents

Anthrone reagent: 0.4% anthrone in concentrated sulphuric acid.

#### Procedure

Aliquot (0.1 ml) of ethanol extract was evaporated to dryness in a test tube. After cooling, the residue was dissolved in 1 ml of distilled water and to it 4 ml of anthrone reagent was added. The mixture was then boiled in a water bath for 10 min. After cooling, absorbance was recorded at the wavelength of 620 nm against a reagent blank with the help of UV-Vis spectrophotometer. Standard curve was prepared using graded concentration of D-glucose (20-100  $\mu$ g/ml).

#### Estimation of soluble proteins

#### Sample preparation

Pellet left after soluble carbohydrate extraction was extracted in 1.25 ml chilled Tris buffer (0.1 M, pH 8.0) containing 0.1% polyvinyl

pyrrolidone (PVP). It was centrifuged at 10000 rpm for 15 min. The supernatant containing the proteins was taken in a test tube and pellet was discarded and processed for the quantification of proteins by the method of Bradford (1976).

#### Reagents

Commassie Brilliant Blue G-250 reagent was used. 100 mg of CBBG-250 reagent was dissolved in 50 ml of 95% ethanol. To this solution 100 ml 85% (w/v) phosphoric acid was added and final volume was made to 200 ml with double distilled water. The solution was filtered through Whatman No. 1 filter paper, and final volume was made to 1 L and stored at 4°C in amber colour bottle.

#### Procedure

To 100  $\mu$ l of the aliquot taken in test tube, 5 ml of the CBBG-250 reagent was added and mixed thoroughly either by inversion or vortexing. The optimal density (O.D.) was measured at 595 nm after 15 min and before 1 h against reagent blank. Standard curve was prepared using graded concentration of bovine serum albumin (20-100  $\mu$ g/ml).

#### Interspecific hybridization

Since the three species of Cyamopsis employed in the present study differed in their flowering schedule; these were grown in a staggered manner to synchronize their flowering. The flowers of the female designate parent were emasculated in the evening (between 04:00 p.m-06:00 p.m.) prior to their anther dehiscence. Generally only two floral buds were emasculated on a raceme to permit their proper development. The crossing of C. tetragonoloba was carried out on the following morning between 07:00a.m. and 08:30 a.m. with pollen grains of C. serrata or C. senegalensis. Pollinated flowers were harvested after 24, 48 and 72 h fixed in acetic alcohol and processed for in vivo germination and pollen tube growth by aniline blue method as described earlier. The allogamous pistils left in situ were allowed to set pods. Percent pod set and number of seeds/pod were recorded. In addition to the conventional breeding method, non-conventional methods like stub pollination with or without smearing with pollen germination medium (PGM), in vivo placental pollination and placental pollination followed by in vitro pistil culture on MS medium supplemented with naphthalene acetic acid (NAA), indole acetic acid (IAA) and benzylaminopurine (BAP) also were attempted.

#### Stub pollination

The stigma of emasculated pistils of *C. tetragonoloba* was excised and was smeared with cool and molten pollen germination medium (PGM) with the help of camel hair brush and then pollen of *C. serrata* and *C. senegalensis* were applied separately on the stigma of emasculated flowers. Pollinated pistils were collected after 24, 48 and 72 h and processed in the same way as explained above for study *in vivo* pollen germination and tube growth. All self pollinated flowers below the selected buds were removed thereby ensuring that all the lowest buds on the raceme are always emasculated ones. To avoid damage to the raceme, upper buds were not removed until 3 day after emasculation; however, upper buds blooming during this period were removed immediately. The whole inflorescence was bagged to check any undesired pollination.

#### Pollination through perforation in the basal part of style

With the help of sterilized syringe needle, a hole was made in the

S/N	Parameter	C. tetragonoloba	C. serrata	C. senegalensis
1	No. of pollen/flower	4765.94 ± 241.23	4684.26 ± 506.55	4582.66 ± 453.49
2	Pollen size (µm )	$34.90 \pm 0.23$	$38.47 \pm 0.30$	54.46 ± 0.52
3	Pollen viability (%)	95.77	87.01	98.12
4	In vitro pollen germination (%)	94.42	25.59	70.01
5	Pollen tube length (µm)	1204.9	174.7	949.2
6	Pistil length (cm)	0.67 ±0.02	0.53 ±0.01	0.77 ±0.02
7	Shape of stigma	Capitate	Subapical crescent	Capitate
8	Number of ovules/pistil	7-8	8-9	7-8
9	Number of pods/ cluster	6.75±0.23	12.00±0.10	9.50±0.52
10	Length x breadth of pod (cm)	5.95 x 0.54	4.51 x 0.36	5.1 0x 0.40
11	Number of seeds/pod	7.58±0.19	8.75±0.12	7.45±0.35
12	Test weight of 100 seeds (g)	2.78±0.21	1.81±0.19	1.84±0.30

Table 1. Comparison of floral and male reproductive characters in three different species of Cyamopsis.

upper part of ovary. Another hole opposite to this was also made to allow release of air. Pollen suspension of *C. serrata* and *C. senegalensis* in the liquid PGM was injected separately into the ovary through a hole. The hole was plugged with petroleum jelly and pistils were collected after 24 and 48 h and processed as explained earlier.

#### **Placental pollination**

A cut with the help of sharp needle was made along the ventral suture of the pistil to open up the pistil and expose the placenta. Pollen grains were dusted on the placenta and pistil was rolled back into their normal configuration under *in situ* conditions. Pistils were collected after 24 and 48 h of placental pollination and processed as explained earlier. In other set of experiment, pistils were excised from the plant, sterilized with 95% ethanol on the hood of laminar flow, cut open to expose placenta, pollinated with the desired pollen *viz C. serrata* or *C. senegalensis* and inoculated on MS basal medium supplemented with BAP, IAA, kinetin, NAA, adenine sulphate and casein hydrolysate (CH).

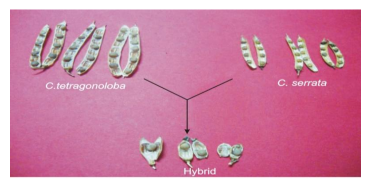
#### **RESULTS AND DISCUSSION**

## Reproductive characters and *in vivo* pollen germination

Table 1 clearly evinces that flowers of all species of *Cyamopsis* studied produced nearly identical number (nearly 4500) of pollen grains per flower. Auto fertile species, like legumes in general, are known to produce few small pollen grains in small flowers (Diaz and Macnair, 1999) and such a reduced male allocation in the species studies is reflective of cost effective allocation to male function (Berlin, 1988) Leguminous pollen are know to contain carbohydrates as well as lipids (Baker and Baker, 1979). These storage compounds not only are source of energy to growing pollen tubes but may play a role in protecting pollen from desiccation e.g. sugars (Pacini 1996) or in synthesis of membranes eg. lipids. Among different species studied, pollen diameter of *C. senegalensis* was maximum (54.46  $\pm$ 1.10 µm) whereas

the value was least in C. tetragonoloba cv. HG 563  $(34.90 \pm 0.63 \mu m)$ . The diameter of *C. serrata* pollen grains was 38.47 ±0.80 µm. Pollen viability as assessed by 2,3,5-triphenyl tetrazolium chloride (TTC) test was more than 95% in C. senegalensis and C. tetraganoloba whereas the value was comparatively lower (87.01%) in C. serrata. Pollen grains of C. tetragonoloba showed maximum germination (94.42%) whereas pollen grains of C. senegalensis yielded 70.01% germination. On the other hand, 25.59% pollen grains of C. serrata germinated after 48 h of incubation. Among the tested species, pollen grains of C. tetragonoloba cv. HG 563 produced the longest tube (1204.9 µm) after 4 h of incubation while those of C. serrata produced the smallest pollen tube (174.7µm) after 48 h of incubation. Tube length of C. senegalensis pollen was 949.2 µm after 30 h of incubation (Figure 3). It seems that at anthesis, the embryosac probably is not fully developed and following pollination, it attains maturity in 1 to 2 day as has been reported for Pistacia vera (Shuraki and Sedgley, 1997). Standardization of in vitro germination would be helpful in interspecific hybridization in a number of ways. It is pre-requisite for stub pollination smeared with PGM, in vitro pollination and fertilization etc. Interestingly, nutritive requirement and lag period for in vitro germination varied significantly in the three species. Pollen grains of C. tetragonoloba required 25% sucrose +100 ppm boric acid + 300 ppm calcium nitrate + 0.8% agar.

The pistil plays a crucial role in the reproductive biology of flowering plants. Studies on the pollen-pistil interaction in leguminous taxa are limited in spite importance of legumes in agricultural production. Stigma of guar is most receptive during 7:30 to 9:00 am while pollen grains are reported to remain viable throughout the day (Anonymous, 1984). Among the three species of *Cyamopsis* studied, the pistil and style length was maximum (77, 38 mm) in *C. senegalensis* and minimum in (53, 25 mm) *C. serrata. C. tetragonoloba* and *C.* 



**Figure 1.** Comparison of morphological features of pods of *C. tetragonoloba, C.* Serrata and their  $F_1$  hybrid.

senegalensis possessed capitate type of stigma whereas C. serrata is characterized by subapical crescent shaped stigma (Figure 4). Fabaceae, in general, is characterized by wet stigma. The stigma is covered with surface cells that often lyse to release viscuous secretion containing proteins, amino acids, lipids, polysaccharides and pigments. These secretions not only support retention and germination of pollen grains but protect stigma against desiccation. The role of proteins in "Wet" stigmas is not clear (Esau, 1977). Mattson et al., (1974) emphasized on the role of proteins in hydration of pollen grains. Proteins, which constitute one of the main constituent of stigma + styles of angiosperms was nearly identical quantitatively in all the three species of Cyamopsis studied (8 mg/100 mg FW). Total soluble carbohydrate content of Stigma + styles of C. tetragonoloba and C. serrata was nearly identical (5-6 mg/100 mg FW), whereas those of C. senegalensis consisted of minimum quantity (2.4 mg/100 mg FW) of soluble carbohydrates. Similar to the observations of Cruden (2009) in Fabaceae, pollen size was not correlated with style length in different species of Cyamopsis studied. The pistil of C. tetragonoloba and C. senegalensis possessed nearly identical number of ovules (7-8) while C. serrata is characterized by 8 to 9 ovules per pistil. Number of seeds per pod ranged from 7 to 9 and did not reveal any significant difference in the wild and cultivated species of Cyamopsis (Table 1). Among the three species, 100 seed weight of C. tetragonoloba was maximum (2.78 g) whereas the value was nearly identical in C. serrata and C. senegalenis (1.80 a).

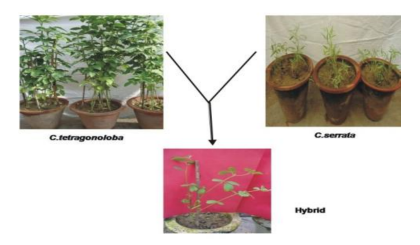
Selfing in *C. tetraganoloba* resulted in good percentage of *in vivo* pollen germination and pollen tubes could be traced up to the base of the ovary. Micropylar entry of pollen tube in the ovule was evident. Interestingly, in *C. senegalensis* and *C. serrata* no pollen germination was evident after 1 day of anthesis. Pollen germination and tubes became evident on/after two days of anthesis and grew until the 3 day after the anthesis.

#### Interspecific hybridization

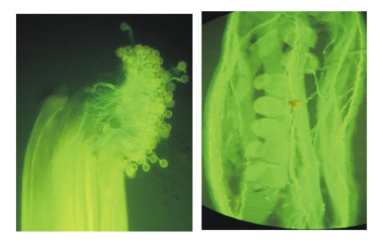
Interspecific hybridization holds great promise in broadening the genetic base of domesticated plant species and success in interspecific hybridization depends upon the extent cross compatibility between the cultigen and its wild relative. Interspecific hybridization between C. tetragonoloba x C. serrata and C. tetragonoloba x C. senegalensis was attempted using conventional and non-conventional breeding methods. Studies deploying conventional method of plant breeding revealed no pod setting in the above said crosses. Among an array of non conventional plant breeding methods tried smearing of stub of C. tetragonoloba with agarified pollen germination medium (PGM) prior to pollination was successful (Figure 1). Among 792 crosses attempted, 83 pods were recovered for the cross C. tetragonoloba x C. serrata which amounted to 10.47% pod set. The hybrid pods were nearly 1.70 cm long which contained 2.28 seeds per pod. Seed colour and shape of  $F_1$  hybrid was similar to the female parent, that is, C. tetragonoloba (Figure 2). However, no success was achieved in 429 crosses attempted between C. tetragonoloba x C. senegalensis even with stub pollination combined with PGM application, although number of pollen grains sticking on the stub increased.

Among the other methods tried *viz.* pollination through perforation in the basal part of style, *in vivo* placental pollination and placental pollination followed by *in vitro* pistil culture did not reveal any pod set; the pistils turned brown and abscised after about 2 to 3 days of pollination from the plant. Failure of placental pollination may be ascribed to withering of ovules after pollen application as has been described for Fabaceae genera (Zenkteler, 1980). In case of placental pollination followed by *in vitro* pistil culture, callusing was observed in all growth combinations tried.

The  $F_1$  seeds obtained from crossing (2008-09) *C.* tetragonoloba × *C.* serrata were sown during the year 2009-10. Morphological and phenological features



**Figure 2.** Comparison of plant morphological features of *C. tetragonoloba, C. Serrata* and their  $F_1$  hybrid.

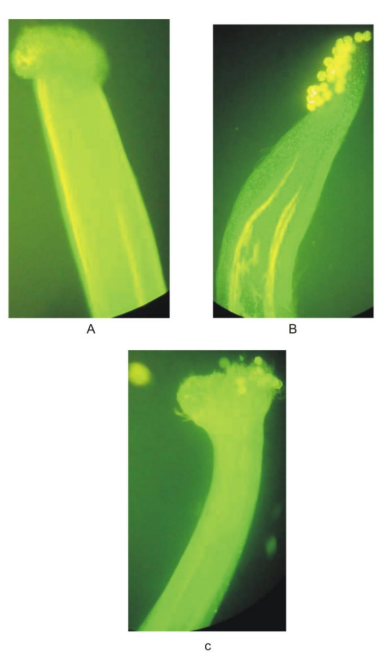


**Figure 3.** L-R *In vivo* germination of *C. serrata* pollen on the pistil stub of *C. tetragonoloba* smeared with pollen germination medium. Pollen tube entering the ovules is evident.

revealed that the plants showed flowering after 21 days of sowing as that of C. Serrata, the wild parent. The flowers were pinkish in colour like the female parent (C. tetragonoloba) but the shape of pods was akin to the male parent (C. serrata). The height of the plant and pod size was 45.5 cm and 4.32 x 0.50 cm, respectively which is intermediate between the two parents. The hybrid plants produced 14.8 pods per plant, 3 to 4 pods per cluster, 6.47 seeds per pod and 3<sup>rd</sup> or 4<sup>th</sup> leaf turned out to be the first trifoliate leaf in contrast to 5<sup>th</sup> to 7<sup>th</sup> leaf in the parent plants. All these morphological and phenological characters are suggestive of the hybrid nature of the plants (Figure 3). Inheritance of seed size in this cross revealed its association with the female parent due to its large size over C. serrata however, yield potential of F1 hybrid was low.

It is thus evident that the differences in the nutritive requirements and wide variations in lag period during

pollen germination of three species of *Cyamopsis* are the potent pre-fertilization barriers in rearing interspecific hybrids by conventional breeding methods. Smearing of pistil stub of C. tetragonoloba with molten and cool pollen germination medium followed by manual pollination with C. serrata pollen induced germination and subsequent tube growth culminating in seed set. Since hybrid plant showed early flowering over C. tetragonoloba, the transfer of earliness trait from C. serrata to the cultivated background is possible by the above method. Further, the attempts can be made to test the fidelity of the interspecific hybrid using molecular markers which may further be tested in the field and transgressive segregants can be selected which may help in identifying extra early varieties of guar. This may prove to be a stepping stone for raising two crops of guar in one year under north Indian conditions which ultimately help in increasing production and productivity of guar.



**Figure 4.** Shaps of stigma in different species of *Cyamopsis- Capitate C. tetragonoloba* (A), sub-apical crescent *C. cerrate* (B) and Capitate-*C. senegalensis* (C).

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