Developmental Evidence of Male and Female Gametophytes of *Jatropha curcas* L.: Pollen Capability and Stigma Receptivity

Nilubol NUANJUNKONG¹,*, Chakrit TONGURAI² and Upatham MEESAWAT¹

¹Department of Biology, Faculty of Science, Prince of Songkla University, Songkhla 90110, Thailand
²Department of Chemical Engineering, Faculty of Engineering, Prince of Songkla University, Songkhla 90110, Thailand

(*Corresponding author’s e-mail: auey_orchid@hotmail.com)

Received: 20 November 2017, Revised: 19 June 2018, Accepted: 30 July 2018

Abstract

*Physic nut* (*Jatropha curcas* L.), a promising alternative resource for biodiesel production, produces restrictively a seed yield and has a low level of genetic variation, causing a barrier to improving the oil performance. Therefore, better understanding of reproductive biology is crucially important for breeding programs to achieve commercial viability in the future. The developmental features related to floral bud diameter and floral morphology of male and female gametophytes, by means of light and scanning electron microscopes, and their capability were investigated. It was found that bud diameter could be used as a criteria for identification of pollen and embryo sac stages. Pollen capability and stigma receptivity showed an inverse S-shape with time after flowering. The highest pollen viability assessed by FCR (93.20 - 98.60 %), TTC (78.80 - 87.00 %), and germination test (43.40 %) were obtained at 0 - 8, 0 - 12, and 0 HAF, respectively. Moderately linear correlations were exhibited between viability and germination test ($R^2 = 0.57 - 0.62$). Stigma receptivity was the highest (97.22 - 100 %) on the first three days.

**Keywords:** Pollen viability, Embryo sac, Bud diameter, Stigma receptivity, Biodiesel

Introduction

*Physic nut* (*Jatropha curcas* L., Euphorbiaceae) is a shrub tree, up to 6 - 15 m high, native to tropical America and now vastly grown in tropical regions, such as Africa and Asia [1,2]. This species is deciduous and can survive in arid and semi-arid areas. It is a monoecious plant which produces female and male flowers in the same inflorescence at the branch terminal [3]. The flowering and fruiting periods take place during the wet season (late summer to early fall) [2,4]. This species is an attractive source for biodiesel production due to its inedible oil and fatty acid composition, which is analogous to that of fossil fuel [4]. The lipid mainly consists of high oleic (44.7 %), linoleic (32.8 %), palmitic (14.2 %) and stearic acid (7.0 %) [5]. The high proportion of oleic acid will provide biodiesel with good general properties exhibiting the combination of high cetane number and low cold filter plugging point [6]. However, the full potential of *Jatropha* has not been achieved due to its variety restriction, leading to the limitation of desired traits and lack of high oil yielding variety [4,7]. Thus, a production of variety with improved quality via breeding programs is essential for this plant. Accordingly, haploid and double haploid embryos are required, because they provide homozygous genotypes necessary for breeding programs [8]. Haploid embryos can be induced through anther [9] or unfertilized eggs [8]. An appropriate stage of microspore and embryo sac, which is one of the most critical parameters for the success of the haploid culture, varies among the species [8,10,11]. Thus, the sequentially developmental events of male and female gametophytes relating to floral features of this plant are prioritized information for bud selection at the accurate stages required for such cultures. Moreover, the longevity of pollen and stigma fertility is still limited in this species. Pollen and stigma at high capability periods are demanded for the pollination
Gametophyte and Fertility of J. curcas

Nilubol NUANJUNKONG et al.

http://wjst.wu.ac.th

set to ensure fruit and seed fecundity [12,13]. Cytocchemical staining, including fluorochromatic reaction (FCR) and 2, 3, 5 - triphenyl tetrazolium chloride (TTC), has been widely used to evaluate pollen viability by checking enzymatic activity and membrane integrity [14,15]. Germination assay has also been extensively employed to define the true ability of pollen to germinate under in vitro conditions [16]. Meanwhile, checking of stigmatic peroxidase activity with hydrogen peroxide is common for pistil receptivity testing [17,18]. Hence, this research aimed to determine (i) the development of pollen and embryo sac in relation to floral bud diameter and morphology and (ii) the longevity of pollen and stigma fertility. The information can be utilized for selecting proper flower stages required for the development of haploid plants via anther and ovary/ovule culture, as well as for pollination manipulation, which are essential for oil-yielding improvement programs of J. curcas L.

Materials and methods

Plant materials

Local J. curcas plants grown in sandy loam soil at Trang Agricultural Occupation Promotion and Development Center (TAOPDC), Trang Province, Thailand, were used as materials and kept as voucher specimens (Nuanjunkong 01) in the international herbarium -PSU (Department of Biology, Faculty of Science, Prince of Songkla University). At the plantation, the total rain was 2,266.20 mm, and the average temperature was 28.80 °C. All experiments were carried out during 2013 - 2014 at TAOPDC and Prince of Songkla University, Songkhla, Thailand.

General flower characteristics and gametophyte development

Male and female flower buds were randomly collected at 8.00 - 9.00 a.m. from five individual plants. These buds were measured and divided into 5 groups (1.0 - 1.5, 1.6 - 2.0, 2.1 - 3.0, 3.1 - 4.0, and > 4.0 mm) based on diameter. For each group, morphological appearances of floral organs were recorded on at least six sampling buds and, were photographed using an Olympus SZH 10 stereomicroscope joined to a DP-71 digital camera. The samples were then fixed in FAA II (70 % ethanol: glacial acetic acid: formalin, 90: 5: 5) for 48 h, dehydrated in a series of tert-butyl alcohol, embedded in Paraplast Plus, and cut at 6 µm thickness using a rotary microtome. The sections were stained with hematoxylin and safranin for histological features, and with 0.1 % (w/v) aniline blue for callose wall observation [19]. In the latter case, fluorescent images were taken under excitation wavelength at 365 nm using an Olympus-BX51 microscope. Pollen surface was determined using a SEM-Quanta scanning electron microscope at 20 kV. The ratio of male to female flowers was estimated from 40 inflorescences.

Pollen assessment via FCR, TTC and germination test

Inflorescences with nearly bloomed male flowers were collected, placed in 250 ml beakers filled with 200 ml tap water, and maintained at room temperature (25 ± 3 °C) overnight [15]. The tests were performed on pollen collected from those inflorescences at 0 (anthesis time, 8.00 a.m.), 4, 8, 12, 24, 48, 72, and 96 h after flowering (HAF). For the FCR test, the pollen was transferred to a mixture of 15 % sucrose and fluorescein diacetate (FDA) solution, followed by 10 min incubation in a humidity chamber. Bright yellow-green pollen under the fluorescence microscope, with an excitation wavelength at 489 nm, were scored as viable [20]. For the TTC test, pollen was stained with 1 % TTC solution and kept in darkness for 24 h, then washed thrice with distilled water. Red-stained pollen was considered viable [15]. In vitro germination was performed according to the published protocol [16]. Briefly, the pollen was released into depression slides filled with 20 µL of Brewbaker & Kwack (BK) liquid medium [21] supplemented with 20 % sucrose, 2.5 % coconut water and 10 % polyethylene glycol (PEG 4000) at pH 6.5. The culture media were placed in a moist chamber and kept at 25 ± 3 °C in darkness for 14 h. The pollen whose tube length reached 15 psi at 121 °C for 20 min was simultaneously used as a negative control. Percentages of viability and germination were determined from 5 replicates, each consisting of 100 pollen specimens. Data were submitted to the Kruskal-Wallis test at a 5 % probability level (P ≤ 0.05) available on portable IBM SPSS Statistics, version 19. Linear correlations between the pollen...
staining techniques and the germination test were carried out using simple linear regression analysis on the SPSS.

**Stigma receptivity**

The female flowers were bagged before flowering, and the receptivity of stigmata tested with 3% hydrogen peroxide solution was accomplished at 0 (at flowering time, 8.00 a.m.), 1, 2, 3, 4, 5, and 6 days after flowering (DAF). Stigmata having oxygen bubbles within 2 to 3 min were considered receptive [17]. The percentage of receptive stigmata was calculated using 3 replications, each with 12 stigmata. The Kruskal-Wallis test was used in order to analyze significant differences of means.

**Results and discussion**

**Morphological characteristics of male and female flowers**

Physic nut produces both male and female flowers on the same cymose inflorescence. The innermost female flower, located at the center, is surrounded by male flowers (Figure 1A). In Brazil and China, *Jatropha* inflorescences were classified into corymbose cyme [22] and dichasial cyme [23], respectively. This study showed the number of female flowers was lower than that of male flowers – representing a male to female ratio of 17:1 per inflorescence. A sex ratio (male: female) ranged from 13:1 to 108:1 was reported for this species, as mentioned by Fresnedo-Ramírez [24]. These fewer female flowers resulted in low seed set and low oil productivity in this plant [23]. Male flowers, with round heads and thin pedicels (Figure 1B), have five greenish-yellow sepals and five greenish-yellow petals enriched with hairs (Figure 1C). There are five globular nectaries at the base of the filaments. These nectaries are initially greenish in color and change to yellow at anthesis (Figure 1C). The male flower contains 10 stamens arranged in 2 whorls; free outer and fused inner whorls (Figure 1C). The anther is bilobed (with four pollen sacs), extrorse type (dehisces longitudinally outward) and joins to filament by subbasifixed attachment (Figure 1D). Differently, dorsifixed and basifixed attachments were observed by other authors for this species [22, 23]. The female flower has conical apex and thick pedicel (Figure 1E). It has five greenish-yellow sepals and five greenish-yellow petals with plentiful hairs (Figure 1F) which are similar to those of the male flower. Flattened and ovate nectaries (five in number), presenting yellow color during the anthesis, are arranged at the base of the ovary (Figure 1F). The ovary, with 3 fused carpels, is comprised of 3 locules, each with one ovule exhibiting apical placentation (Figures 1F and 1G). On the connate style rests three bifurcated stigmata (Figures 1F and 1H). Noticeably, sterile stamens showing abnormal-shaped anthers without filament are observed at the base of the ovary, both at early (1.0 - 2.0 mm bud diameter) (Figure 1H) and late (≥ 2.1 mm bud diameter) (Figure 1I) stages of female flower development. Traces of stamens, containing no mature pollen and lacking pollination capability, are also detected in female flowers of physic nuts grown in Brazil [23].
Male and female gametophyte development in relation to bud diameter and morphology

**Male gametophyte (pollen) development**

The anther contains 4 locules (Figure 3A) whose wall layers, namely, the outermost epidermis, endothecium, middle layer, and the innermost tapetum, are observed. Male gametophyte development could be indicated as the stages of (a) pollen mother cell (PMC), (b) microspore tetrad, (c) uninucleate microspore, (d) early binucleate pollen with globular-shaped generative cell, and (e) late binucleate pollen with spindle-shaped generative cell at the flower bud sizes of 1.0 - 1.5, 1.0 - 1.5, 1.6 - 2.0, 2.1 - 3.0, and ≥...
Female gametophyte (embryo sac) development

The anatropous bitegmic and crassinucellate ovule is observed (Figure 4A) and the embryo sac is of monosporic polygonum type (Figures 4B - 4N) which corresponds with and confirms the study mentioned by Krishnamurthy [27] for the same species. The female bud (1.0 - 1.5 mm) presents the initiation of style and stigma, a large sterile stamen, and a small nectary gland (Figure 2B). Megaspore mother cell (MMC) displaying a large nucleus with dense cytoplasm is deeply embedded within a massive nucellus (Figures 4A and 4B). The anatomical feature confirms that the stamens found in female flowers are sterile by having locule shrinkage without PMC at this MMC stage (Figures 4C and 4D), indicating these sterile stamens might have no impact on any pollination sets. This study support a massive nucellus (Figure 2A). The degenerated middle layer is presented while epidermis, endothecium, and tapetum are still clearly visualized (Figure 3E). Finally, bud size of about ≥ 2.1 mm exhibits binucleate pollen stage. Flower bud (2.1 - 3.0 mm) possesses developing filament which is hidden by the presence of small greenish nectaries (Figure 2A). The anther wall is composed of the epidermal layer, the fibrous endothecium, the collapsing tapetum, and the degraded middle layer (Figure 3F). For this size, the majority of buds are at early binucleate mature pollen containing the vegetative nucleus (Figure 3F, arrowhead) and the globular-shaped generative cell detached from pollen wall (Figure 3F, gn). Then, the bud (≥ 3.1 mm) with elongated filaments and developed nectaries (Figure 2A) presents a spheroidal pollen (76.72 ± 1.17 µm in diameter) containing the vegetative nucleus (Figure 3G, vg) and the spindle-shaped generative cell (Figure 3G, arrowhead). At this stage, the layers of the epidermis and the fibrous endothecium still remain (Figure 3G). At anthesis, pollen is shed at a two-celled stage (Figure 3H) and presents surface clavae arranged in a crotonoid pattern (polygonal alignment) (Figure 3I). Observation of the stages of PMC and tetrad in the same bud size of different flowers was possible in this study because the PMC undergoes meiosis rapidly and develops to form a tetrad faster than the enlargement of flower bud. Liu et al. [25] illustrated that the PMC, tetrad, free microspore, and binucleated pollen of physic nut could be detected at the anther lengths of 0.5 - 0.7, 0.7 - 0.78, 0.83 - 1.0, and 1.7 - 2.2 mm, respectively. Thus, a rapid selection, by roughly scanning the developmental stage using flower bud diameter (this study), was prerequisite, and then the anther length provided by Liu et al. [25] could be used to classify the stages between PMC and tetrad. The occurrence of nectary and filament could be used to separate gametogenesis from sporogenesis in this study.

However, both sporogenesis and gametogenesis of Arabidopsis thaliana were observed in flower buds exhibiting visual filament [26]. Using buds whose size correlated to a suitable stage of pollen development has been one of the crucial factors determining the success of in vitro microspore or anther culture [10], and different stages were required for different species [9,11]. Hence, flower buds of ≥ 1.6 mm in diameter presenting haploid unicellular microspore or bicellular pollen might be suitable and recommended for the haploid plant induction of J. curcas.
whose developed style, stigma, and larger nectaries covering sterile stamens are presented (Figure 2B). The bud of 2.1 - 4.0 mm, displaying the white nectaries (Figure 2B), contains immature 8-nucleate embryo sac, in which the migration of two polar nuclei to the center is not yet completed (Figures 4I - 4K), whereas the buds (> 4 mm) having yellow nectaries possess mature 8-nucleate embryo sac comprised of antipodal cells (Figure 4L), egg cell, and synergids (Figure 4M) and two juxtaposed polar nuclei at the center (Figures 4M - 4N). The floral diameter related to the stage of embryo sac has also been established in onion (Allium cepa) [30]. A suitable stage of ovule for haploid and dihaploid production varied depending on species was considered, ranging from the stage of megaspore to mature embryo sac [8]. Consequently, for J. curcas, haploid and double haploid induction would begin by using ovules from ≥ 1.6 mm buds, which contain haploid megaspore to mature embryo sac, to find the most responsive stage of embryo sac on the culture.

**Figure 2** Summarized correlation between flower bud diameter, morphology, and developmental events of *Jatropha curcas* L. in (A) male and (B) female flowers. Scale bar = 2 mm
Therefore, flower buds exhibiting the proper stage of pollen and embryo sac could be chosen for *in vitro* haploid and dihaploid production required for breeding programs to produce new plant cultivar with higher oil yielding. To reduce the biodiesel price and have economic competitiveness, the obtained cultivar should contain oil higher than the 37 - 50 % derived from the rapeseed (*Brassica napus*), the main feedstock of the world biodiesel production [31].

**Figure 3** Pollen development of *Jatropha curcas* L. (A) Anther with four locules (arrows); (B) PMC with large nucleus and all anther walls including epidermis (Ep), endothecium (En), middle layer (Md), and tapetum (T); (C) Tetrad (Td) and all anther wall layers; (D) Callose wall (arrows) surrounding the tetrad microspores after aniline blue fluorescence; (E) Free microspore (MI) displaying the exine wall (arrowhead); (F) Binucleate pollen showing small and dense vegetative nucleus (arrowhead) and a globular-shaped generative cell (gn). Insets illustrating the fibrous thickening of the endothecium; (G) Mature pollen containing the vegetative nucleus (vg) and the spindle-shaped generative cell (arrowhead). Insets showing thicker and more fibrous wall of endothelial cells; (H) Pollen releasing from anther before the opening of petal (pt); (I) SEM micrograph exhibiting mature pollen with clavae (arrow) arranging as croton pattern (circles); En, endothecium; Ep, epidermis; gn, generative cell; Md, middle layer; MI, microspore; PMC, pollen mother cell; pt, petal; T, tapetum; Td, tetrad microspore; vg, vegetative nucleus.
Gametophyte and Fertility of *J. curcas*

Nilubol NUANJUNKONG et al.

http://wjst.wu.ac.th

Figure 4 Longitudinal sections of *Jatropha curcas* L. female flowers determining embryo sac development. (A) Anatropous ovule with MMC (arrow), nucellus (n), inner integument (ii) and outer integument (oi); (B) MMC; (C) Sterile stamens (arrows) at MMC stage; (D) Amplification of inset in (C) showing sterile stamen with shrinkleage locule (arrow); (E) Three degenerating megaspores (arrows) at micropylar and a survival megaspore (arrowhead) at chalazal pole; (F) Ovule at megaspore stage presenting caruncle (cr), nucellus (arrow), inner integument (ii) and outer integument (oi). Insets show a functional megaspore; (G) Binucleate embryo sac; (H) Multinucleate embryo sac; (I - K) Serial section of immature embryo sac indicating (I) Presumptive egg apparatus (arrows) and polar nucleus (arrowhead); (J) The same sample as (I) in deeper section, showing presumptive egg apparatus (arrow); (K) In other plane, representing presumptive antipodal cells (arrows); (L - N) Mature embryo sac showing (L) Antipodal cells (arrows); (M) Central polar nucleus (arrowhead), egg cell (arrow), and synergid (•); (N) Consecutive plane of (M), showing another polar nucleus (arrowhead) and egg cell (arrow). cr, caruncle; ii, inner integument; MMC, megaspore mother cell; n, nucellus; ne, nectary; oi, outer integument; ov, ovary.

Pollen capability and stigma receptivity

The viable pollen exhibited yellow-green fluorescence (Figure 5A), red color (Figure 5B), and germinated tube (Figure 5C) after FCR, TTC, and germination test, respectively. All techniques employed in pollen capability test, exhibiting inverse S-shape, displayed the reduction of pollen capability along the time course after flowering (Figure 5E). From the FCR result, the highest viability was found at 4 HAF (98.60 %) and not significantly different from 95.80 % at 0 HAF and 93.20 % at 8 HAF. The
viability dropped sharply from 86.60 % at 24 HAF to the lowest (31.40 %) at 96 HAF. By TTC, the maximal viability was 87.00 % at the flowering time, which was not significantly different from 4, 8 and 12 HAF. The viability decreased markedly (66.60 - 21.60 %) during 24 to 48 HAF, and the lowest (15 %) was shown at 72 HAF. The highest germination (43.40 %) was observed at the flowering time. Then, the germination frequency exceedingly dropped to 19.60 % (at 4 HAF) and to 2.20 % (at 96 HAF). The results showed that TTC ($R^2 = 0.6205$), rather than FCR ($R^2 = 0.5686$), was slightly more correlated with the germination test, based on the linear pattern (Figure 5F). TTC test was more effective for the evaluation of pollen viability than FCR for this plant [15]. Accordingly, pollen capability of physic nut should be considered from TTC and germination results, which showed the highest viability of 87.00 % and germination of 43.40 % at the flowering time. Previous reports displayed physic nut pollens were viable at 95.4 % by TTC and were germinated at 30 to 71.6 % [15,16]. The viability was at its peak at 9 h after blooming [32]. The difference might be attributed to the species genotype and growth environment [33,34]. TTC was also reliable for *Leymus chinensis* [35] and *Kochia scoparia* [36], but not for *Festuca arundinacea*, in which viable and dead pollen showed the same color after staining [33]. The receptive stigma represented the number of bubbles at the stigmatic apex (Figure 5D, arrows) and illustrated the inverse S-shape longevity (Figure 5G). Receptivity displayed the maximal value at 100 % during a period of 0 - 2 DAF, 97.22 % at 3 DAF, and steadily decreased to the lowest (36.11 %) at 6 DAF. This relatively concurs with an earlier report mentioning that the physic nut stigma could be receptive for around the first 4 days and began to decline at the fifth day [32].

Figure 5 Pollen capability and stigma receptivity of *Jatropha curcas* L. (A) Bright fluorescence of viable pollen after FCR test; (B) Red viable pollen (arrows) after TTC test; (C) Germinated pollen tube (arrow); (D) Released bubbles (arrows) indicating receptive stigma by hydrogen peroxide test; (E) Viability (by TTC and FCR tests) and germination percentage of pollen showing inverse S-shape pattern; (F) A linear relationship between pollen viability and germination; (G) Stigma receptivity showing inverse S-shape longevity. Different letters in each line indicate significant differences at $P \leq 0.05$. 
Conclusions

The study described the relationship between flower bud diameter and the developmental stages of both pollen and embryo sac in physic nut. Floral buds of 1.6 mm in diameter onwards, which highly related to haploid microspore to binucleate pollen for male flowers and haploid megaspore to mature embryo sac for female flowers, would be employed for haploid production. Pollens collected at blooming and stigmata at 0 - 3 DAF, displaying high capability, should be applied in artificial pollination and breeding programs.

Acknowledgments

The authors are grateful to the Graduate School, Prince of Songkla University, for research funding and the scholarship for the support of exchange students and international credit transferred through the ASEAN Community. We are thankful to the Science Achievement Scholarship of Thailand (SAST) for the doctoral scholarship. We wish to thank the TAOPDC for their generous permission to employ J. curcas plantation. We also acknowledge Mrs. Anna Chatthong from the Faculty of Science, Prince of Songkla University, for assistance with the English in the manuscript.

References
