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## Association of bud and anther morphology with developmental stages of the male gametophyte of melon (*Cucumis melo* L.)

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
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**Abstract.** Correlations between the morphological features of flower buds and the developmental stages of the male gametophyte are of great practical interest as a reliable marker that accelerates and simplifies the selection of appropriate plant material for isolated microspore culture. Microspore culture enables one to quickly obtain many pure lines of different vegetable crops, but it has not yet been widely applied in the melon (*Cucumis melo* L.). To successfully apply this technique in a new culture, one has to optimize many of its elements: first, find the biological markers for selecting the flower buds containing the microspores of certain development stages. The paper presents the results of research estimating the correlations between the length and diameter of the flower buds, the length of the visual part of the corolla, the length of the anthers and the development stages of the male gametophyte in the F<sub>1</sub> hybrid of the Kim Hong Ngoc melon. The strongest correlation (CC = 0.885) was found for the flower bed diameter and a strong correlation (CC = 0.880), for the bud length. The corolla's visual part was a less reliable morphological feature, and the anther's length should not be used as a parameter to predict the developmental stages of the melon's male gametophyte. It was also found that one anther could contain the microspores and pollen grains of different developmental stages. In the flower buds less than 4 mm in length and 1.51 ± 0.02 mm in diameter prevailed tetrads, and in the buds 4.0–4.9 mm in length and 2.30 ± 0.02 mm in diameter, early microspores. The microspores of a middle stage of development prevailed in the flower buds 5.0–5.9 mm in length and 2.32 ± 0.00 mm in diameter; mid and late vacuolated microspores, in the buds 6.0–8.9 mm in length and 2.96 ± 0.37 mm in diameter; and two-celled pollen, in the buds more than 9 mm in length and more than 3.97 ± 0.34 mm in diameter. Key words: male gametophyte; stages of microspore development; tetrad; pollen; flower bud; anther; *Cucumis melo* L.; melon.


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## Соответствие морфологии бутонов и пыльников стадиям развития мужского гаметофита дыни (*Cucumis melo* L.)

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**Аннотация.** Выявление корреляций между морфологическими признаками бутонов и стадиями развития мужского гаметофита представляет большой практический интерес, так как наличие надежного маркера ускоряет и упрощает отбор подходящего растительного материала для культуры изолированных микроспор. Культура изолированных микроспор позволяет в короткие сроки получать чистые линии многих овощных культур, однако для дыни (*Cucumis melo* L.) эта технология пока не получила распространения. Чтобы успешно применить данную технологию для новой культуры, необходимо оптимизировать множество ее элементов, прежде всего подобрать морфологические маркеры, позволяющие отбирать бутоны, которые содержат микроспоры определенных стадий развития. В нашей работе приведена оценка корреляции между длиной бутонов, диаметром бутонов, длиной видимой части венчика, длиной пыльников и стадиями развития мужского гаметофита дыни F<sub>1</sub> гибрида Kim Hong Ngoc. Наиболее сильная корреляция установлена для диаметра бутонов, коэффициент корреляции составил 0.885. Сильная корреляция выявлена также для длины бутона, коэффициент корреляции 0.880. Длина видимой части венчика являлась менее надежным признаком, а длина пыльников не следует использовать в качестве параметра для прогнозирования стадий развития мужского гаметофита дыни. Отмечено, что в одном пыльнике одновременно находились микроспоры и пыльцевые

зерна разных стадий развития. В бутонках длиной менее 4.00 мм и диаметром до  $1.51 \pm 0.02$  мм преобладали тетрады; в бутонках длиной 4.0–4.9 мм и диаметром  $2.30 \pm 0.02$  мм обнаружена наибольшая доля ранних микроспор, при этом преобладали микроспоры средней стадии развития; в бутонках длиной 5.0–5.9 мм и диаметром  $2.32 \pm 0.00$  мм преобладали средние и поздние вакуолизованные микроспоры; в бутонках длиной 6.0–8.9 мм и диаметром  $2.96 \pm 0.37$  мм – поздние вакуолизованные микроспоры; в бутонках длиной 9.0 мм и более, диаметром  $3.97 \pm 0.34$  мм и более – двухклеточная пыльца.

Ключевые слова: мужской гаметофит; стадии развития микроспор; тетрада; пыльца; бутон; пыльник; *Cucumis melo* L.; дыня.

## Introduction

The melon (*Cucumis melo* L.) is an economically important cultivated plant (Sebastian et al., 2010) grown in more than 1 mln ha of agricultural lands (FAOSTAT, 2019)<sup>1</sup>. For the time being, the most common melon has been F<sub>1</sub> hybrids praised for their uniformity and high yield and providing proper biological protection of originator's ownership.

Double haploids (DHs) are a valuable material of genetic research and selection, especially for F<sub>1</sub> hybrids of agricultural plants (Shmykova et al., 2015b; Abdollahi et al., 2016). As of today, the technologies to obtain DHs have been developed for more than 250 species (Maluszynski et al., 2003) and many of them have been used to produce homozygous plants (Ferrie, Caswell, 2011).

Several publications describe successful melon DH production via pollination with irradiated pollen (Sauton, 1988; Hooghvorst et al., 2020) or via remote hybridization followed by embryo growing *in vitro* (Lotfi et al., 2003). There are also papers, whose authors cultivated the anthers (Abdollahi et al., 2016), unfertilized seedbuds (Shmykova et al., 2015a) and isolated microspores (Zhan et al., 2009; Chen et al., 2017) of members of the cucumber family.

The isolated microspore culture technique produces more regenerates compared to those of unfertilized seedbuds and anthers and is widely applied, especially in the cabbage family (Djatchouk et al., 2019; Kozar et al., 2020). Moreover, this technique excludes the somatic cells of a donor plant from the growing medium, leaving no doubt about the regenerates' origin. However, it has never been applied to produce the DHs of members of the cucumber family.

DH production in isolated microspore culture can be affected by multiple factors such as microspore development stage; their genotype; growing medium composition; cell-rich fluid density; culture introduction technique; the effect of temperature and other cultivation conditions (Dunwell, 2010; Niaziyan, Shariatpanahi, 2020). The microspore development stage is the first factor to be accounted for when applying the isolated microspore culture technique to a new culture, because the development from tetrads to two-celled pollen may involve different stages (Touraev et al., 1991; Germanà, 2011). For example, to produce carrot DHs, it is recommended to cultivate tetrads and early microspores (Gorecka et al., 2010), while cultivation of middle and late microspores is most effective for callus induction in the balsam apple anther culture (Nguyen et al., 2019). And in the cabbage family, vacuolated microspores and two-celled pollen have the highest ability for embryogenesis (Telmer et al., 1992; Binarova et

al., 1997; Custers et al., 2001; Babbar et al., 2004; Winarto, Teixeira da Silva, 2011).

Direct selection of separate microspores corresponding to a certain development stage to be cultivated *in vitro* seems to be an unresolvable problem. As a rule, plant material is selected based on such markers as the morphological characteristics of the flower buds and anthers (Takahata, Keller, 1991; Parra-Vega et al., 2013). In rape, soya, reddish, tomato, balsam apple, these markers include the length and widths of their flower buds (Weber et al., 2005; Han et al., 2014; Sumarmi et al., 2014; Adhikari, Kang, 2017; Nguyen et al., 2019). Several studies have proved that such parameters as the size and color of the flower cup as well as the cup/corolla length ratio and anther size can do the trick (De Moraes et al., 2008; Parra-Vega et al., 2013; Zhang et al., 2013). Since these parameters are species-specific, it is necessary to work out a specific protocol for the melon.

This paper presents the results of investigation into the morphological characteristics of the melon's flower buds and anthers and the way they correlate with the plant's microspore development stages.

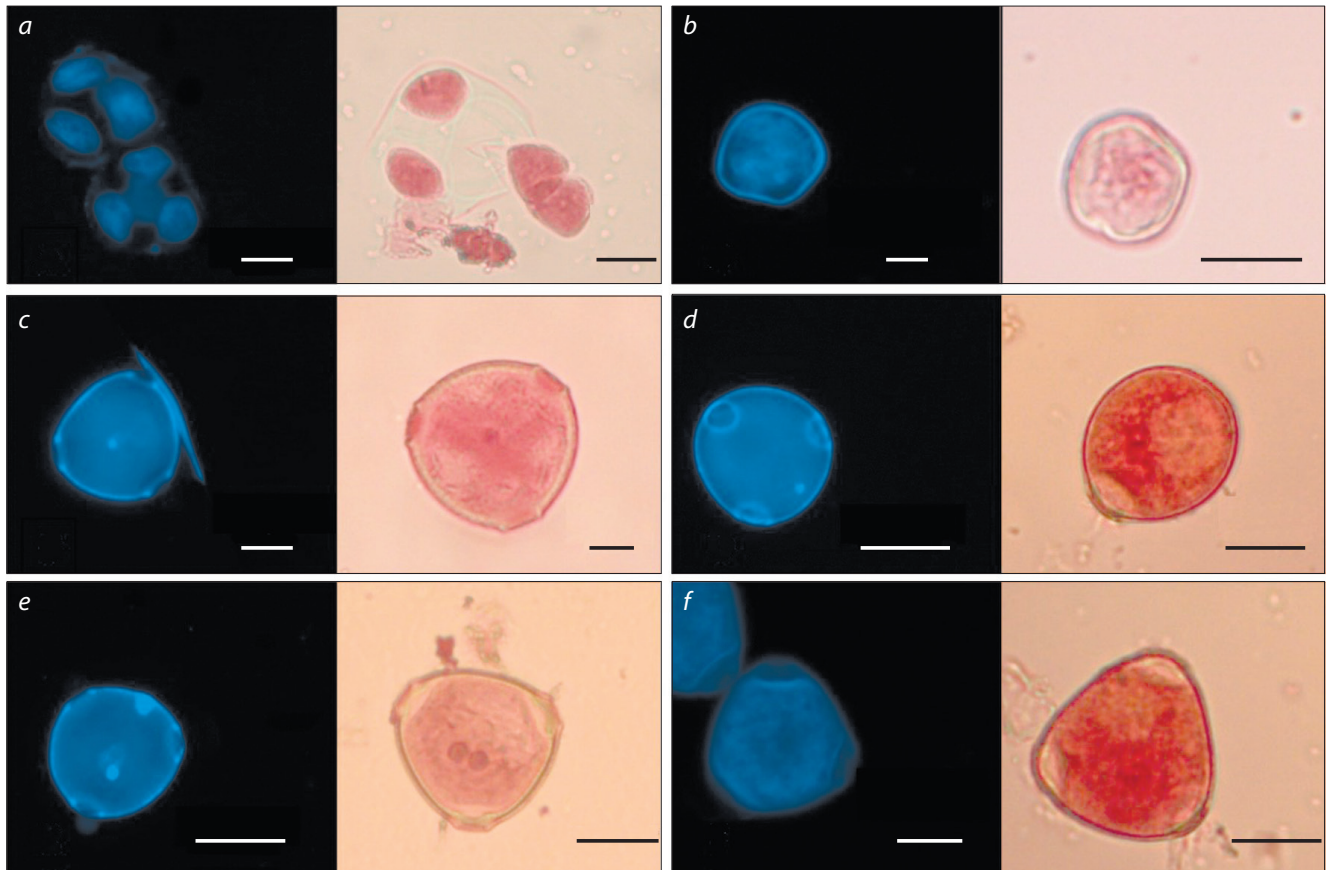
## Materials and methods

The flower buds of the F<sub>1</sub> hybrid plants of the Kim Hong Ngoc melon produced by the Chia Tai Seed company (Thailand) were collected at 5:30–6:30 a. m. The buds of 3.6 to 15.6 mm in length (with 1-mm interval) were transported in ice and then stored for 24 hours at 4 °C. At least 10 buds were accounted for each of the intervals.

The buds' morphological characteristics were assessed using a Zeiss Stemi 2000-C stereomicroscope (Suzhou Co., Ltd). Microspores were obtained from the anthers of each flower bud to be put on a glass slide into a drop of glycerin mixed with distilled water in proportion 1:1. Then the 15 µl of 2 % acetocarmine solution drop was added, covered with a cover slide and microscopied. For the purpose of fluorescent staining, the microspores extracted from the anthers were washed three times in PBS (8.0 g/l of NaCl, 0.20 g/l of KCl, 1.44 g/l of Na<sub>2</sub>HPO<sub>4</sub> and 0.24 g/l of KH<sub>2</sub>PO<sub>4</sub> were dissolved in the 3/4 of the required volume of distilled water; HCl and KOH were added to bring the pH value to 7.4, and distilled water was added to reach the finite volume), DAPI (4',6-diamidino-2-phenylindole) was added and then the microspores were studied using a Zeiss Axio Lab1 fluorescent microscope (Suzhou Co., Ltd).

The microspore development stages were determined from the size and shape of the cells, the number of cell nuclei and their interposition (Vergne et al., 1987; Maluszynski et al.,

<sup>1</sup> <http://www.fao.org/faostat/en/#data/QC> (Accessed 01.06.2021).



**Fig. 1.** DAPI- and acetocarmine-dyed microspore development stages of *Cucumis melo* L.: a – tetrads, b – early microspores, c – mid microspores, d – late vacuolated microspores, e – early two-celled pollen, f – late two-celled pollen. Bar = 20  $\mu\text{m}$ .

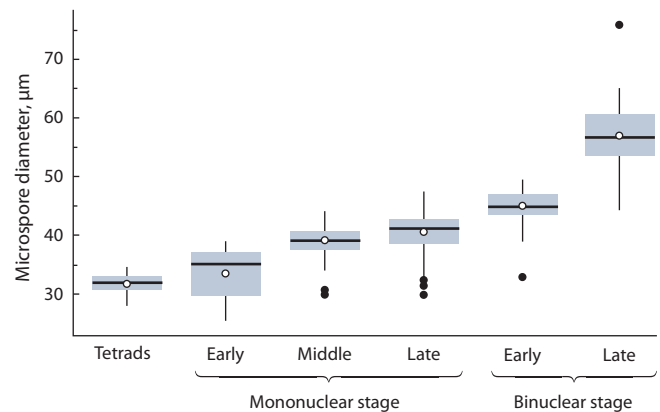
2003; Blackmore et al., 2007; Zhang et al., 2013). In each specimen, the development stages of 100 microspores were observed. Such parameters as the presence of tetrads, early/middle/late microspores and two-cell pollen were considered. The percentage of each development stage in a particular specimen was calculated as the ratio of the number of microspores related to a certain development stage to the total number of observed microspores multiplied by 100 %.

The statistical significance of the performed calculations was confirmed with ANOVA analysis and the Tukey test for  $\alpha = 0.05$ . The correlation between the measured parameters and microspore development stages was determined using the linearity regression ( $R$ ) and correlation ( $CC$ ) coefficients. The collected data were described and processed with the R software.

## Results and discussion

During the cytological analysis of melon flower buds, 6 stages of microspore development were observed. These included tetrads, early/middle/late vacuolated microspores, early/late two-celled pollen (Fig. 1).

The diameter of the microspores increased as they developed and reached their maximum at the stage of late two-celled pollen (Fig. 2). It has been noted that each stage was characterized by a certain shape and size of the cells. The diameter of the early microspores formed after tetrad degradation



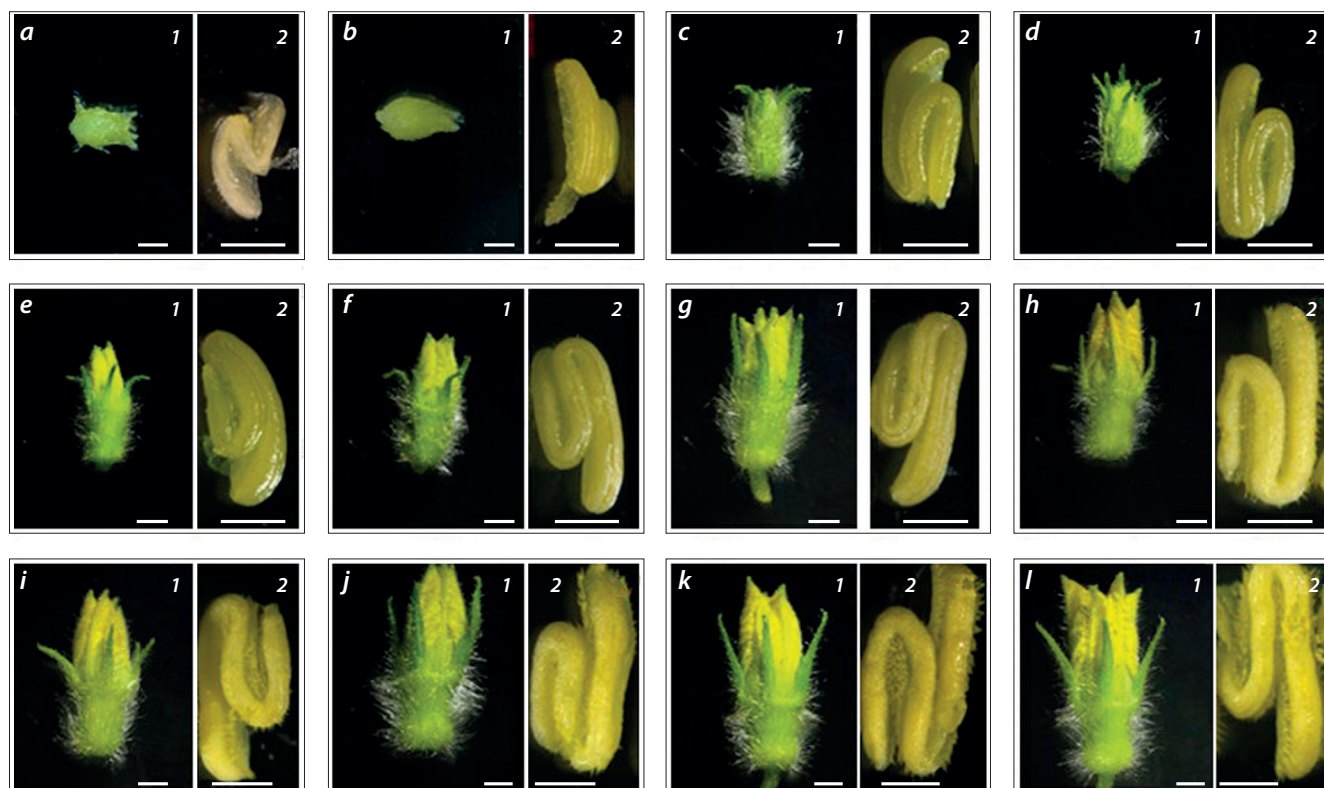
**Fig. 2.** Changes in male gametophyte diameters related to their development stages.

was  $33.41 \pm 4.34 \mu\text{m}$ ; they were of uneven circular shape and had thin walls and large nuclei. The middle microspores were  $39.06 \pm 2.33 \mu\text{m}$  in diameter, had a round shape and a centered nucleus. The late microspores were round and had a well-expressed three-lobed exine wall with the nucleus pressed to it by a big vacuole. Their diameter was  $40.45 \pm 3.26 \mu\text{m}$ . The cells of early two-celled pollen had  $44.94 \pm 2.65 \mu\text{m}$  in diameter with well-expressed two nuclei: a larger vegetative

Correlations between flower-bud sizes, anther lengths and the stages of male gametophyte development in the melon

No.	Buds length, mm	Buds diameter, mm	Anther length, mm	Microspores developmental stages, %					
				Tetrads	Early	Mid	Late	Early two-celled pollen	Late two-celled pollen
1	<4	1.51 ± 0.02 <sup>a*</sup>	1.63 ± 0.19 <sup>a</sup>	48.00 ± 5.66	22.00 ± 8.49	6.00 ± 2.83	22.00 ± 2.83	2.00 ± 2.83	0.00 ± 0.00
2	4.0–4.9	2.30 ± 0.02 <sup>b</sup>	1.83 ± 0.16 <sup>a</sup>	0	30.00 ± 14.14	46.00 ± 25.46	22.00 ± 14.14	2.00 ± 2.83	0.00 ± 0.00
3	5.0–5.9	2.32 ± 0.00 <sup>b</sup>	2.50 ± 0.51 <sup>b</sup>	0	2.00 ± 2.83	50.00 ± 14.14	48.00 ± 11.31	0.00 ± 0.00	0.00 ± 0.00
4	6.0–8.9	2.96 ± 0.37 <sup>b</sup>	2.89 ± 0.23 <sup>c</sup>	0	0.40 ± 1.26	40.40 ± 24.91	56.80 ± 23.61	2.40 ± 6.31	0.00 ± 0.00
5	9.0–11.9	3.97 ± 0.34 <sup>c</sup>	2.96 ± 0.20 <sup>c</sup>	0	0	0.67 ± 1.63	14.67 ± 34.00	51.33 ± 50.62	33.33 ± 50.13
6	>12	5.16 ± 0.27 <sup>d</sup>	3.12 ± 0.22 <sup>c</sup>	0	0	0	0	0	100.00

\* Data marked with the same letters do not differ at  $p = 0.05$ .



**Fig. 3.** Changing the morphological characteristics of melon buds (1, bar = 20  $\mu$ m) and anthers (2, bar = 10  $\mu$ m) in relation to flower-bud sizes: a – 3.6–4.0 mm; b – 4.0–4.9 mm; c – 5.0–5.9 mm; d – 6.0–6.9 mm; e – 7.0–7.9 mm; f – 8.0–8.9 mm; g – 9.0–9.9 mm; h – 10.0–10.9 mm; i – 11.0–11.9 mm; j – 12.0–12.9 mm; k – 13.0–13.9 mm; l – more than 14.0 mm.

and a more vividly-colored generative one. The diameter of the late two-celled pollen comprised  $56.93 \pm 4.81 \mu$ m, its cell shapes varying from round to oval, so one anther could contain pollen grains of different shapes. The pollen's cytoplasm became dense and nontransparent making it more difficult to observe the nucleus.

The results of the correlation analysis to bring together the morphological features of melon flower buds and corresponding microspore development changes enabled us to subdivide the buds into 6 groups. Each group could include microspores

of different stages, so at least one of these stages prevailed (see the Table). It was noted that a single anther could have microspores that belonged to different development stages, which corresponds to the observations of other researchers who studied this issue in other cultures.

The tetrads were found in green oval-shaped pubescent flower buds that were fully covered in sepals and had a length of less than 4 mm and a diameter of 1.85 mm (Fig. 3, a). The buds' anthers were of light-beige color and had 1.6–1.63 mm in length.

The early microspores were found in flower buds of 3.8–7.0 mm in length, their biggest portion ( $30 \pm 14.14\%$ ) concentrated in buds of 4.0–4.9 mm. The buds' anthers changed their color to green-yellow, their length comprised 1.63–2.74 mm (see Fig. 3, *b*). The early microspores were found in smaller amounts compared to the other development stages.

The middle microspores concentrated in flower buds of 4.0 to 10.9 mm in length. The buds' anthers were  $2.15 \pm 0.05$  mm in length and had a yellowish glazing surface (see Fig. 3, *c*). The microspores prevailed ( $50 \pm 14.14\%$ ) in the buds of 5.0–5.9 mm in length. Such buds had a clear morphological difference from younger buds: their sepals were open, so one could see the corolla tip.

The late vacuolated microspores prevailed in buds of 6.0–8.9 mm in length. At this stage, the buds kept growing in size, so the corolla extended beyond the sepals. However, the anthers' morphology remained unchanged (see Fig. 3, *d-f*) as did their length.

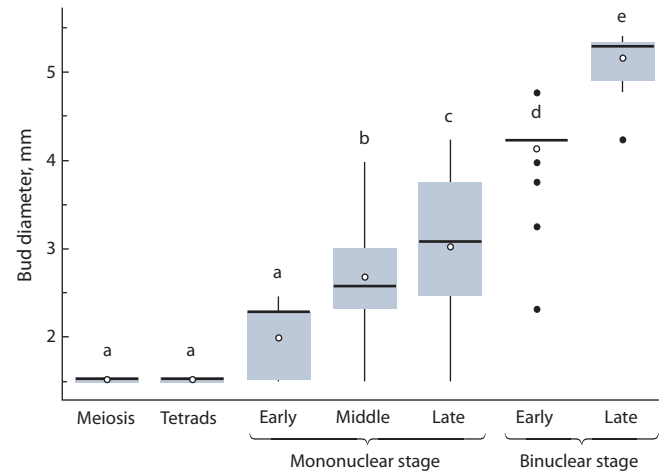
The early two-celled pollen prevailed in buds of 9.0–12.0 mm in length (see Fig. 3, *g-i*). Their anthers' length, compared to the previous stage, remained unchanged, their surfaces containing mature pollen grains.

As for buds larger than 12 mm in length, they contained only two-celled pollen. The transition from the late to mature stage was characterized by a small increase in bud size, its petals starting to open (see Fig. 3, *j-l*). The anthers increased in size and opened too, so a large number of pollen grains could be seen on their surface.

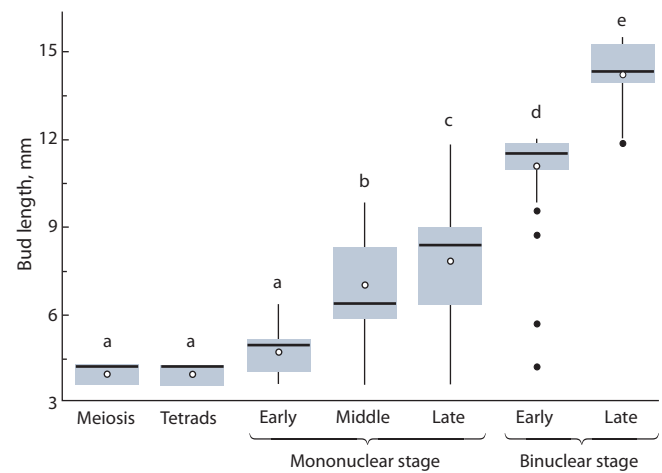
Statistical analysis of anther lengths gave us linear regression coefficient  $R^2 = 0.52$ , which meant that this parameter could not be used as a predictor of microspore development stages in the melon, which corresponded to the results obtained for some other cultures such as the tomato and aubergine (Segui-Simarro, Nuez, 2005; Salas et al., 2012). In (Adhikari, Kang, 2017), the authors obtained a similar coefficient ( $R^2 = 0.59$ ) when studying a relation between anther length and microspore development stages in the tomato.

Many researchers recommend using flower-bud length to select proper plant material to cultivate isolated microspores for it is a convenient and reliable morphological parameter for many plant species. They also recommend bud diameter as an indicator for flower bud selection. A study published in 2019 demonstrated that the best results in the embryogenesis of lucerne microspores were obtained when cultivating late microspores from flower buds of 6.02–6.20 mm in length and 1.50–1.72 mm in diameter (Yi et al., 2019). In 2017, a correlation between flower-bud size (length and diameter), anther length and microspore development stages in the tomato was published (Adhikari, Kang, 2017).

In our study, a linear regression analysis showed there was a clear linear dependence ( $p < 0.05$ ) between the flower-bud characteristics and microspore development stages. The regression coefficients ( $R^2$ ) varied from 0.767 to 0.783. The strongest correlation was for flower-bud diameter ( $r = 0.885$ ,  $R^2 = 0.783$ ) (Fig. 4), followed by flower-bud ( $r = 0.880$ ,  $R^2 = 0.775$ ) (Fig. 5) and anther ( $r = 0.876$ ,  $R^2 = 0.763$ ) lengths, the last being the least reliable feature.



**Fig. 4.** Correlation between the melon's flower bud diameter and male gametophyte development stages.



**Fig. 5.** Correlation between the melon's flower bud length and male gametophyte development stages (a–e correspond to different bud groups).

## Conclusion

The correlation between the morphological characteristics of the flower buds and anthers of the melon (*Cucumis melo* L.) and the development stages of its microspores enables one to select a proper material for cultivation of isolated microspores *in vitro*. The characteristics in question are flower-bud diameter and length and the length of visible corolla. Since the correlation coefficient is higher for the diameter and length of flower buds, these parameters are easier to use.

The obtained results can be applied for further development of the technology to produce melon DHs in isolated microspore culture.

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