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GAMETOPHYTE DEVELOPMENT OF Arthromeris mairei (BRAUSE) CHING OF SOUTH SIKKIM, INDIA

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ABSTRACT

Gametophyte development in *Arthromeris mairei* (Brause) Ching was studied and found to be of '*Drynaria* type'. Germination occurred 10-12 days after sowing of spores. Many prothalli showed an initial archegonial phase, which persisted throughout gametophyte development and the antheridial phase developed on separate thalli after a gap of 5-7 days and persisted throughout the lifespan of the gametophytes. This type of development of sex organs is resembled to some extent, except few variations with the sex organ development of *Arthromeris himalayansis* (Ganguly *et al.* 2009), probably due to the similar type of ecological and climatic conditions where they grow. This type of gametangial development on separate prothalli is an indication of adaptation for cross-fertilization.

Keywords: Arthromeris mairei (Brause) Ching, gametophyte development, Drynaria type, adaptation, Gametophyte development, Arthromeris merii, Epiphyte, Type H, out breeding.

INTRODUCTION

Arthromeris mairei (Brause) Ching belongs to the family Polypodiaceae and is a warm temperate fern, exclusively epiphytic in nature and distributed in India throughout the Himalayan region from Eastern Himalayas to Western Himalayas. This epiphytic fern is also found in China, Nepal and Burma in mountain areas. In Southern Sikkim this fern is generally found between 2700-3600 m.

Germination of fern spores (Tryon and growth further Lugardon, 1991), and development of resulting gametophytes in artificial media is a well-attended area in pteridophytic and developmental biological research (Navar, 1962; Atkinson and Stokey, 1964; Kato, 1969; Klekowski, 1969; Nayar and Kaur, 1969; Nayar and Kaur, 1971; Masuyama, 1975 a, b; Khare & Kaur, 1983; Raghavan, 1989; Chiou & Farrar, 1997; Verma et al. 2000; Verma, 2003; Ganguly and Mukhopadhyay, 2005, 2009). Nayar & Kaur (1971) and Atkinson (1973) pointed out that the sequence and plane of cell divisions, pattern of gametophyte development as well as the direction of initial growth of the first rhizoid and germ filament with respect to the polarity of germinating spore are distinct characteristics and can be utilized effectively for drawing phylogenetic relationships among various taxonomic groups. Nayar (1962) studied the spore germination and prothalial morphology of Arthromeris mairei (Brause) Ching along with some other polypodiaceous ferns. However, no work has so far been done on the prothalial development of the epiphytic fern Arthromeris mairei (Brause) Ching. Ferns occupy a specialized habitat as epiphytes and as such epiphytic ferns evolved gametophytic have various generation adaptations like antheridiogen systems, production of gemmae, indefinite growth of prothalli etc. (Farrar, 1974; 2003). The current study was performed to understand the details of gametophyte structure and development pattern of prothalli in Arthromeris mairei (Brause) Ching.

MATERIALS AND METHODS

Mature sporophylls of Arthromeris mairei (Brause) Ching were collected from some healthy plants from Maenum Wildlife Sanctuary and Versey Rhododendron Sanctuary (3023 m and 3000 m altitude), South Sikkim in the month of October of 2017 and 2016. Sporophylls of individual plants were kept separately within blotting spores paper and matured from sporophyll(s) of individual plants dehisced within 48 hrs were collected in separate vials for gametophyte studies. Collected spores from three individual sporophytes were sown in separate petri plates on modified Moore's medium (Kato, 1969). Two replicates of each set were maintained. These spores were surface sterilized by 0.1% HgCl₂ (w/v) solution for 5-8 minutes and rinsed three times with sterilized distilled water and then dried on sterilized blotting paper. The sterilized spores were transferred to autoclaved (at 15 lb / inch² for 15 minutes) modified Moore's culture medium (Kato, 1969) solidified by 1% (w/v) agar in an aseptic chamber and the pH of the medium was maintained at 5.8. The cultures were incubated at 22°C-25°C under cool fluorescent white light (ca 1000 lux, 16hr/d).

Gametophytes from each Petri plates were studied every day randomly by light microscope (Leica DMLB) after germination of spores. Time taken for spore germination and to form mature gametophytes, initiation of sex organs and formation of sporophytes were recorded. Photographs of different developmental stages were made on the same microscope. Records of study of gametophyte development pattern from individual sporophytes were maintained separately in order to see whether there were variations in the developmental patterns within the individuals. Observations

were made on ten gametophytes from each petriplates at a time.

RESULTS

Characteristics of Spore Germination and Gametophyte Development

- a) SPORE: Spores were bilateral, monolete; light brown in colour, perisporate, perispore thin, size 38-42 x 50-53 μm.
- b) PERCENTAGE OF GERMINATION: Spore germination of Arthromeris mairei (Brause) Ching was 71.49 ± 4.66%. Spores germinated even after having been stored for two months after collection.
- c) GAMETOPHYTE DEVELOPMENT: Spores germinated 10-12 days after sowing. In this species the rhizoidal cell formed first (Plate 1 A) followed by the chlorophyllous protonemal cell (Plate 1 B). The protonemal cell developed into a 6-celled stage by periclinal divisions (Plate 1 C-E). Spore germination resulted in a slender uniseriate germ filament. The penultimate protonemal cell underwent oblique vertical division. In Arthromeris mairei (Brause) Ching the establishment of an apical meristem was much delayed and the prothalli usually developed hairs on the margin and surfaces. A broad spathulate prothalli plate was formed by repeated longitudinal and transverse divisions of its anterior cells and expansion of the resultant daughter cells (Plate 1 F-K). Mature vegetative, cordate shaped gametophytes (Plate 1 M) developed 77-80 days after spore germination. The mature prothalli measured ca 350 x 300 µm in size. The prothallial plate often become 15-20 cells or more wide and broadly ovate, but was

devoid of any organized meristem. Later, an ob-conical meristematic cell was differentiated by two oblique divisions in one of the marginal cells at the anterior end of the prothallial plate (Plate 1 K). The meristematic region (Plate 1 L-M) was located under notch. The type of development was purely "*Drynaria* type" as discussed by Nayar & Kaur (1969, 1971).

Development of Sex Organs (Sequence, Position and Duration)

Mature cordate gametophytes remained vegetative for about 30 days, after which the gametophytes started to develop sex organs. Archegonia developed first in some cordate shaped prothalli 112 ± 2 days after germination. Archegonia spore were situated along the midrib region and just below the meristematic region. Archegonia consisted of a projecting neck (Plate 1 M) and a lower embedded venter. This flask shaped structure was made up of two axial rows of neck canal cells, one ventral canal cell and one egg cell (Plate 1 N). Each archegonium had a single lavered jacket (Plate 1 N) and were 150-200 x 65-75µm in size. Antheridia developed on separate gametophytic thalli, which were elongated and much longer than archegonial prothalli. Initiation of antheridia started 115 ± 2 days after spore germination. Antheridia were of the emergent type (Plate 1 O) with a 1-cell thick jacket measuring about 25-30 µm.

In Arthromeris mairei (Brause) Ching, the prothalli were dioecious. The cordate shaped prothalli developed archegonia after they reached maturity and remained as archegoniate prothalli throughout the reproductive phase. Antheridiate prothalli were elongated and did not form welldefined apical meristem. Antheridia developed on the lower half of the prothallus in marginal and/or superficial in position (Plate 1 L). After initiation, antheridia took about 3-5 days to mature; spermatozoids were released after this period. The time taken for development of the different stages of gametophytic stages of *Arthromeris mairei* (*Brause*) Ching is shown in Table 1.

Table 1. Time taken for gametophyte development
of Arthromeris himalyensis (Hook.) Ching

SI. No.	Events of the gametophyte development	Total No. of days taken after sowing of spore ±SD	
1.	Sowing of spores	0±0	
2.	Spore germination	11±1	
3.	Formation of mature cordate prothallus	83±3	
4.	Initiation of archegonia	110±2	
5.	Initiation of antheridia	115±3	
6.	Maturation of antheridia	120±3	
7.	Initiation of sporophyte	127±2	

DISCUSSION

From the above observations, we can conclude that the type of gametophyte development in Arthromeris mairei (Brause) Ching is purely "Drynaria type". In "Drynaria tvpe" development, spore germination results in a slender uniseriate germ filament. A broad spathulate prothallial plate is formed by repeated longitudinal and transverse divisions of its anterior cell and expansion of the resultant daughter cells. The prothallial plate often becomes 5-10 cells or more wide and broadly ovate, but is devoid of any organized meristem. Later, an obconical meristematic cell is differentiated by two oblique divisions on one of the marginal cells at the anterior end of the prothallial plate. The young prothallus becomes cordate, the apical meristematic cell is replaced by a pluricellular meristem and a midrib developed. Young prothalli are naked; hairs are usually formed when the prothallial plate becomes cordate (Nayar and Kaur, 1969). The Drynaria type of

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development characteristic of is Cheiropleuriaceae, Dipteridaceae. Gleicheniaceae, Lomariopsidaceae, Loxomaceae, Thelypteridaceae and the majority of the Polypodiaceae genera (Nayar and Kaur, 1969). According to Masuyama's (1975 a, and b) classification of on gametangial gametophytes, based sequence of development on meristematic prothalli and was further elaborated upon by Verma (1989, 2003) and Ganguly and Mukhopadhyay (2005, 2009), the gametophytes of Arthromeris mairei (Brause) Ching resemble 'type A' to some extent. In type A, the archegoniate prothalli persist throughout development but the antheridiate prothalli become archegoniate in the later stages. In Arthromeris mairei (Brause) Ching, the sequence of development of the sex organs is different. Here, the archegoniate prothalli remain archegoniate and the antheridiate prothalli remain antheridiate throughout development. Based on the sequence of sex organ development in Arthromeris mairei (Brause) Ching, it is identified as new type different from the types described by Masuyama (1975 a, b), Verma (2003) and Ganguly and Mukhopadhyay (2005). Thus, we propose a new type "Type H" in addition to the existing seven types classified by the previous authors (Table 2).

Table 2. Classification of gametangial sequence on meristematic prothalli of homosporous ferns (Adapted
from Verma 1989, 2003; Ganguly e <i>t al.</i> 2009)

Туре	Sequential bearing of gametangia on meristematic prothalli			
	Initial state	Final state	Symbolization	
А	Antheridiate	Archegoniate.	$M\toF$	
	Archegoniate	Persists throughout.	$F\toF$	
В	Antheridiate	Antheridia and Archegonia formation.	$M\toH$	
	Archegoniate	Antheridia and archegonia formation.	$F\toH$	
С	Antheridiate	Antheridia and archegonia formation for some time, then only archegonia formation.	$M\toH\toF$	
			$F\toH\toF$	
	Archegoniate	Same		
D	Antheridiate	Antheridia and archegonia formation for some time, alternating	$M{\leftrightarrow}H{\leftrightarrow}F{\leftrightarrow}H$	
		periodicity in the formation of antheridia and archegonia, finally		
		Sama F↔H	F↔H↔M↔H	
	Archegoniate	Same		
Е	Antheridiate	Archegonia formation.	$M\toF$	
	Archegoniate	Antheridia and archegonia formation.	$F\toH$	
F	Archegoniate	Antheridia and archegonia formation (ephemeral), then antheridia formation.	$F \rightarrow H \rightarrow M$	
			or	
			F→M	
G	Archegoniate	Antheridia and archegonia formation simultaneously.	$F\toH$	
H*	Archegoniate	Persists throughout.	$F\toF$	
	Antheridiate	Persists throughout.	$M\toM$	

Symbols indicate the sequential state of functional sex:

M = Antheridia formation, F = Archegonia formation, H = Hermaphrodite.

Types A, B and C are according to Masuyama (1975 b). Type D, E and F are proposed by Verma (1989). Type G is proposed by Ganguly & Mukhopadhyay (2005). Type **H*** is a new variant type proposed here by current authors.

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Plate 1 (a-n): (a) A spore. (b) A germinating spore. (c) A spore showing first rhizoidal and first prothallial cell. (d-e) A germinating spore showing second prothallial cell (f-h) Different stages of filamentous gametophyte. (i) Gametophyte showing anticlinal division (j) Spathulate gametophyte (k) hairs on mature gametophyte (l) Group of archegonia just beneath the meristematic region (m) A mature archegonium (n) A mature antheridium (o) A mature cordate gametophyte

Gametophyte growth habit can be classified into three basic types in regard to the effect of form on breeding system. Type I is the familiar cordate or butterfly shaped gametophytes of most terrestrial ferns. Type II gametophytes have indeterminate growth and branching and type III gametophytes combine type II growth with production of dispersible gemmae. Type II and type III gametophytes are typical of most epiphytic species (Farrar, 2003). In Arthromeris mairei (Brause) Ching, the archegoniate prothalli resemble with type I, which is cordate shaped. The antheridiate prothallus was elongated, having indefinite growth. Some of the gametophytes showed clonal elongation. The secondary gametophytes produced antheridia on their margins. Thus antheridiate prothalli resemble type II gametophytes partially.

The gametophytes of *Arthromeris mairei* (Brause) Ching are long lived (more than 110 days), and the advantage of long-lived gametophytic generation is to promote cross-fertilization (Klekowski, 1973, 1979). Opportunities for gamete exchange between long-lived gametophytes are much higher than for short-lived non-clonal epiphytic gametophytes.

Most species of Polypodiaceae maintain an antheridiogen system through which the robustly growing female gametophytes induce production of antheridia precociously on the smaller gametophytes growing nearby, thus enhancing the probability of cross-fertilization (Chiou and Farrar, 1997). *Arthromeris mairei* (Brause) Ching may have an antheridiogen system, as antheridia grow on separate prothalli after 3-5 days of initiation of archegonia in cordate shaped prothalli. This suggests that antheridiogen might have some role in controlling the reproductive system of *Arthromeris mairei* (Brause) Ching. Masuyama (1975 b) recognized four basic locations of antheridia on monoecious prothalli viz, Antheridia on the lower part of the body of gametophytes thallus (type L), on the lower half of the wings (LW), on the lower half of the margin (type LM), on the upper half of the central cushion (type UC), or antheridia are located all along the margin (type M). As *Arthromeris mairei* (Brause) Ching produces dioecious prothalli, it does not resemble any type as recognized by Masuyama (1975 b), though the antheridia located on the lower half of the wings (type LW) as proposed by Masuyama (1975 b).

It is interesting to note that, the percentage of spore germination is very high, it is about 79.49 ± 4.66% even after two months of harvesting. This figure indicates that this species produces a high proportion of viable spores, which is likely helpful in the survival of this species. This species is restricted to a certain altitudinal regions (2700-3600 m), thus specific environmental conditions like temperature, annual rainfall, relative humidity (RH) etc are required for its survival. RH, annual rainfall and altitude have a combined effect on the distribution and reproductive success of this species (Ganguly and Mukhopadhyay, 2008).

From the above discussions, it may be concluded the *Arthromeris mairei* (Brause) Ching gametophytic generation shows some derived developmental features:1) dioecious prothallus ensures cross-fertilization; 2) archegoniate prothalli that are meristematic and cordate shaped, continuously producing archegonia, increase the chances of sporophyte production and 3) long-lived gametophytes (more than 110 days) that also promote cross-fertilization.

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