

ISAH Indian Journal of Arid Horticulture

Year 2024, Volume-6, Issue-2 (July-December)

Propagation strategies for conservation and commercial development of root medicinal crops: A review

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ARTICLE INFO

ABSTRACT

Received: 09 December 2024 Accepted: 07 March 2025

Keywords: Ayurvedic formulations, germplasm conservation, herbal medicine, *in vitro* propagation, tuber

doi:10.48165/ijah.2024.6.2.2

India is one of the mega diversity regions endowed with a rich array of medicinal plants. However, the surging demand for herbal medicines has triggered destructive harvesting methods, jeopardizing the natural populations of these plants. Medicinal crops such as ashwagandha, mulethi, and shatavari are globally sought-after due to their therapeutic potential, addressing stress relief, immune function, and hormonal balance. Despite their historical use as natural remedies, the escalating demand is primarily met by wild populations, endangering the survival of these crop species. The alarming rate of over-exploitation necessitates the development of efficient propagation protocols. This ensures planting material for commercial cultivation and contributes to conservation efforts. Moreover, the cultivation of high-value medicinal plants is creating a new dimension in the field of agriculture, propelling the Indian herbal industry into a flourishing stage. However, the intricate nature of medicinal plant cultivation poses challenges, particularly due to limited knowledge of seed biology. This review concentrates on ashwagandha, mulethi, and shatavari cultivation practices, compiling existing literature to serve as a guide for future research in this crucial field.

Introduction

Medicinal plants have been providing a tremendous source of natural medicines since time immemorial. *Ashwagandha* (*Withania somnifera* (L.) Dunal), *mulethi* (*Glycyrrhiza glabra* L.) and *shatavari* (*Asparagus racemosus* Willd.) are popular medicinal root crops used in traditional Ayurvedic medicine (Niraj and Varsha, 2020). These have been used for centuries for their various health benefits and therapeutic properties (Das and Pandey, 2023). They can be used individually or in combination with other herbs in Ayurvedic formulations to address specific health concerns and promote overall wellbeing. *Ashwagandha* also known as Indian Ginseng/winter cherry provides benefits to the nervous system, rejuvenates the body, prevents aging, improves muscle and joint function, and aids in sleep (Saran *et al.*, 2025; Singh *et al.*, 2021). While *mulethi* or licorice has sweet-tasting roots, is commonly used for respiratory health, digestive support, anti-inflammatory, antioxidant properties, immune-modulatory effects, and more (Ahemed *et al.*, 2021). Similarly, *Shatavari* is believed to enhance fertility and is associated with various health benefits, particularly for the female reproductive system (Alok *et al.*, 2013).

However, the conservation of these plants is among the most pressing issues. To satisfy the need for medicinal plants, natural resources are being depleted at an exponential

pace (Sen *et al.*, 2011). The medicinal crops are not grown systematically; people generally collect them from the forest. In this context, it is important to promote their cultivation scientifically to prevent genetic erosion and maximize profits. To achieve the full health benefits of these medicinal crops, proper GAP (Good Agricultural Practices) protocols are needed during planting (Saha *et al.*, 2018). Planting root crops such as *ashwagandha*, *mulethi* and *shatavari* can be done through a few different methods, depending on the specific requirements of each plant as outlined in Fig. 1. These plants are commonly propagated through their seeds or root divisions/cuttings. However, the rapidly changing climate has resulted in changes in the key components of certain Medicinal and Aromatic Plants (MAPs) (Manish *et* *al.*, 2016). Hence, it is essential to explore alternative ways of growing Medicinal and Aromatic Plants (MAPs). Currently, micropropagation, hydroponics, and aeroponics are gaining popularity in this regard. However, micropropagation has started for certain medicinal plants, the adoption of hydroponics and aeroponics in the production of Medicinal and Aromatic Plants (MAPs) is still at an early stage. These techniques offer a sustainable alternative, providing high-quality roots that are devoid of pesticides and soilborne diseases (Mehdizadeh and Moghaddam, 2023). The conservation and sustainable use of medicinal plants require a long-term approach, so this review focuses primarily on their propagation methods. Here is a general guide to planting these root crops:



Fig. 1. Different propagation methods of medicinal plants

Planting methods of different medicinal root crops

The planting method is almost similar for *ashwagandha*, *mulethi* and *shatavari*. However, specific climatic and soil conditions may affect their growth.

Site selection

Selecting an ideal location for cultivating medicinal root crops is essential for a successful harvest. Medicinal root

crops such as *ashwagandha*, *mulethi* and *shatavari* thrive well when they receive a minimum of 6-8 hours of direct sunlight daily. Sunlight is vital for photosynthesis, enabling plants to produce energy for robust root growth. Root crops prefer loose, fertile, and well-drained soil that allows their roots to penetrate easily. Well-drained soil prevents water logging and the risk of root diseases. Ensuring there are no hard layers beneath the soil further aids good root penetration (Gupta, 2017).

Soil and climate

Ashwagandha, mulethi and shatavari are medicinal herbs

that thrive in warm climates and have specific soil and environmental preferences conducive to their growth. These herbs can adapt to various soil types but perform exceptionally well in specific conditions. For instance, *ashwagandha* does best in sandy or loamy soil within a pH range of 7.5 to 8.0 (Kumar *et al.*, 2023, while *mulethi* prefers a pH range of 5.5 to 8.2 (Cui *et al.*, 2023), and *shatavari* thrives in soil with a pH range of 6.0 to 8.0 (Chaudhary *et al.*, 2023). These pH ranges reflect their soil acidity or alkalinity requirements for optimal nutrient absorption. However, it is equally important to provide good air circulation in the planting area. Adequate air circulation prevents the development of fungal diseases and promotes overall plant health by reducing humidity levels.

Land preparation

Preparing the soil for the cultivation of ashwagandha, mulethi, and shatavari is a crucial step in ensuring their successful growth. To create optimal growing conditions, it is essential to thoroughly pulverize the field. This process typically involves ploughing and harrowing, which break up compact soil and enhance its structure. For root crops, loose soil with good aeration is essential as it allows their roots to penetrate easily and promotes healthy growth. Therefore, it is advisable to loosen the soil to a depth of about 12 inches (30 cm) to provide ample space for root development (Gupta, 2017). In regions with heavy or clayey soil, additional measures are necessary. Two to three rounds of ploughing before the rainy season can help break down the soil structure further. Moreover, incorporating well-decomposed farmyard manure (FMY) or organic matter into the soil can significantly improve its texture and fertility.

Propagation of Ashwagandha (Withania somnifera (L.) Dunal)

Propagating *ashwagandha*, an important medicinal plant, can be achieved through two primary methods: direct sowing and propagule raising in a nursery. The choice of method depends on factors such as the availability of seeds or seedlings, the local climate, and the specific needs of the cultivation process. However, the best way to propagate *ashwagandha* is through seed. Choose disease-free and high-quality seeds from well-recognized fibrous varieties, like Jawahar Asgand-134, Jawahar Asgand-20, Raj Vijay Ashwagandha-100, Gujarat Anand Ashwagandha-1, Vallabh Ashwagandha-1 (Khabiya *et al.*, 2023). The second group of *ashwagandha* varieties is rich in starch (low crude fiber) having long, stout, and thick root quality of produce obtained (Saran, 2023, Saran and Das, 2023). The starchy type high-yielding varieties/landrace/germplasm are Vallabh Shahi

Ashwagandha-1 (DTWr-1), DNA-4, Nagori, CIM Pratap and CIM Chetak are rich in starch and low crude fiber content (Saran *et al.*, 2025).

Ashwagandha cultivation can involve either using seeds directly or purchasing seedlings from a nursery. When directly planting seedlings in the field, dig a hole slightly larger than the root ball of the plant, carefully place the seedling in the hole, fill it with soil, and gently firm the soil around the plant. As a precautionary measure, treat the seeds with Captan at a rate of 5 g/kg seed before sowing to protect emerging seedlings from potential seed-borne diseases (Kumar *et al.*, 2023).

Direct sowing

The process of sowing seeds for ashwagandha typically involves the broadcasting method, where the seeds are evenly scattered across the main field. Sowing of the seeds generally occurs in the second week of July to August. During this period, the soil is adequately moistened by the rains, creating ideal conditions for germination. However, it is worth noting that the timing of sowing can be somewhat flexible depending on local weather conditions. In cases where there is ample rainfall and soil moisture, the sowing can be extended up to September. The field is divided into convenient-sized plots or rows to facilitate organized and efficient planting. The recommended seed rate is typically 10-12 kg of seeds per hectare, ensuring sufficient plant density for a successful crop (Mathew et al., 2005). Matured seedlings, whether raised through sowing or broadcasting, should be manually thinned approximately 25 to 30 days after seed sowing. This thinning process aims to achieve a plant population ranging from about 30 to 60 plants per square meter (Moharana et al., 2020).

Nursery raising

In a nursery bed, compost and sand are thoroughly mixed and then raised from ground level. Fresh seeds are sown in well-prepared nursery beds in rows, spaced at 5 cm and sown at a depth of 1-3 cm (Chaurasiya *et al.*, 2019). The spacing between plants and rows can be adjusted based on soil fertility and the variety being cultivated. In less fertile soil, closer spacing is recommended, while in more fertile soil, greater distances are advisable, taking into account factors such as resource competition, root development, water management, and pest control to facilitate robust growth and maximize yield. Maintaining a gap of 20 to 25 cm between rows and 8 to 10 cm between individual plants is generally recommended (Moharana *et al.*, 2020). Approximately 5-10 kg of seeds are required for planting in one hectare of the primary field (Namdeo and Ingawale, 2021). Sowing takes

place right before the beginning of the monsoon and is lightly covered with sand. Germination typically occurs within 5 to 7 days. Once the seedlings reach approximately 35 days of age, they are transplanted into the well-prepared main field during July-August (Farooqi and Sreeramu, 2004). Adequate moisture, facilitated by a light shower after sowing, is essential for good germination. It is crucial to maintain consistently moist soil until the seedlings emerge. Although transplanting *ashwagandha* seedlings is a common practice, it has been observed that transplanted seedlings exhibit bifurcated roots, hence, direct sowing is recommended (Fig. 2).



Fig. 2. Nagori ashwagandha root (a) direct sowing (b) transplanting

Transplanting

For the cultivation of ashwagandha, the selected field should be thoroughly pulverized through ploughing or harrowing, followed by leveling. While the ashwagandha crop does not demand heavy doses of manure, it is advisable to apply 10 to 20 tons of Farm Yard Manure (FYM). Ridges are prepared at a spacing of 60 cm, and healthy seedlings are planted at intervals of 30 cm. In certain locations, spacing of 60 cm x 60 cm or 45 cm x 30 cm is also adopted (Farooqi and Sreeramu, 2004). However, the preferred spacing is 60 cm x 30 cm, ensuring an optimal seedling population of approximately 55,000 per hectare (Namdeo and Ingawale, 2021). Notably, there is a noticeable variation in profitability due to different sowing methods. Regarding sowing techniques, the highest net return was observed with the raised bed method, followed by the ridge and furrow method, and then the flatbed method (Ratre et al., 2018).

In-vitro propagation

The substantial disparity between the demand and supply of *ashwagandha* cannot be addressed effectively through traditional propagation methods due to its low seed germination rate and seedling viability (Ameen *et al.*, 2023). *In-vitro* propagation emerges as the most suitable approach to overcome the constraints of conventional propagation and meet the commercial demands for *ashwagandha*. Numerous biotechnological interventions, including callus and suspension culture, hairy root culture, and others, have been developed to enhance the propagation process, as highlighted in Table 1.

Table 1. In vitro regeneration of ashwagandha

Mode of propagation	Explant used	Reference
Direct regeneration	Nodal explant	Tawade <i>et al</i> . (2023)
Multiple shoot differen- tiation	Nodal explant	Kaur <i>et al</i> . (2021)
Callus induction	Shoot explants (shoot tips, coty- ledonary leaf, ma- tured leaves, node, internode)	Sharada (2004)
Multiple shoots differ- entiation	Nodal explant	Govindaraju <i>et</i> al. (2003)
Multiple shoots differ- entiation	Nodal explant	Mir <i>et al.</i> (2014)
Direct regeneration	Nodal explant	Autade <i>et al.</i> (2016)

In contrast to callus culture, cell suspension culture offers the advantage of containing homogeneous cell populations and can be readily scaled up for large-scale cultivation (Ahuja *et al.*, 2021). In addition, it is noteworthy that somatic embryogenesis stands out as the swiftest method for generating a substantial quantity of clonal plants, given that somatic embryos encompass both shoot and root ends. Nevertheless, the utilization of synthetic seeds, including encapsulated somatic embryos or vegetative propagules such as nodes, axillary buds, and shoot apices, serves as a valuable strategy for preserving germplasm in elite medicinal plants (Verma *et al.*, 2010). Moreover, Singh *et al.* (2006) documented their successful production of synthetic seeds

in *ashwagandha*. This was accomplished by encapsulating shoot tips from 4-week-old in vitro-cultured shoots using a mixture of sodium alginate (3.0%) and calcium chloride (75 mM). Similarly, Fatima *et al.* (2013) employed axenic nodal segments to create synthetic seeds in *ashwagandha*. These nodal segments were enveloped in a matrix consisting of 3% sodium alginate and 100 mM calcium chloride, then cultured on a Murashige and Skoog (MS) medium supplemented with BAP (6-benzylaminopurine) and NAA (naphthaleneacetic acid), aiming to facilitate the optimal transformation of encapsulated nodal segments into plantlets.

Propagation of Mulethi (Glycyrrhiza glabra L.)

Mulethi propagation can be accomplished through two methods: using seeds or employing root/stolon cuttings (Dastagir and Rizvi, 2016). However, the latter, involving 10-15 cm root/stolon cuttings, is more commonly chosen due to the relatively low seed set and germination rates (Bakhane *et al.*, 2014; Sharma *et al.*, 2010). The recommended variety for cultivation is 'Haryana Mulhatti-1,' released by Chaudhary Charan Singh Haryana Agricultural University, Hissar (Bhardwaj *et al.*, 2020). Planting is typically carried out in January and February in areas with irrigation facilities and during July and August in regions dependent on rain for sufficient soil moisture.

Propagation through seed

Variations in the germination capacity of *mulethi* were observed among seeds at different stages of maturation (Rao 1993). It is difficult to germinate seeds at the milky way ripe stages but seeds collected in July have the highest germination rates. It is advisable to plant the seeds at a depth of around 1/4 inch in soil. Furthermore, it is advisable to space the seeds at least 6 to 12 inches apart when planting to ensure that there is adequate room for the plants to grow (Öztürk *et al.*, 2018).

Propagation through crown root

For optimal results, it is essential to plough and harrows the field thoroughly, ensuring a fine tilth and weed-free soil. During field preparation, it is recommended to apply Farm Yard Manure (FYM) at a rate of 10 tons per hectare to promote good underground root development (Tewari and Singh, 2019). Planting cuttings of the underground stolon, measuring 10-15 cm in length and possessing 2-3 eye buds, at a depth of six to eight cm and spaced at intervals of 60 x 45 cm or 90 x 45 cm is advisable (Akhtar and Anjum, 2022). Additionally, raising the rows 45-60 cm can facilitate irrigation. The stolon typically begins sprouting within 15-20 days after planting. Notably, *mulethi* propagation can also be achieved through a single root node cutting, which conserves planting material and promotes rapid germination (Fig. 3). During the initial phase, it is essential to provide light and frequent irrigation to support the establishment of the cuttings in the field. Once the plants reach a height of 20 to 30 cm, elevating the rows encourages healthy root expansion. To ensure optimal stolon growth and achieve a substantial yield, it is recommended to leave the crop in the field for a period of 3 to 4 years (Dastagir and Rizvi, 2016).



Fig. 3. Mulethi germination from single root node

In-vitro propagation

Traditional seed-based propagation faces economic challenges due to issues like poor seed set, low seed viability, and a slow growth rate (Jaiswal *et al.*, 2017). As a result, the primary method for propagating the crop relies on valuable vegetative components such as stolons, rhizomes, or cuttings. However, vegetative propagation is not practical, especially for economic parts like roots that take several years to mature (Gupta *et al.*, 1997). Tissue culture stands out as the viable alternative strategy, providing a large-scale *in vitro* propagation of *mulethi* germplasm. This approach enables the production of pathogen-free and season-independent clonal plants. This can be accomplished by utilizing different components of *mulethi*, including nodal buds, apical meristem, axillary buds, leaf segments, shoot buds or by employing somatic embryogenesis.

It can be achieved by using different parts of *mulethi* namely nodal bud, axillary buds, apical meristem, shoot buds, leaf segment or through somatic embryogenesis (Srivastava *et al.*, 2019). Several biotechnological interventions have been reported in recent years to develop alternative strategies to meet industrial demands. Some of the *in vitro* study has been mentioned in Table 2.

Table 2. Ir	ı vitro	regeneration	of	mule	th	i
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Mode of	Explant used	Reference
Widde of	Explain used	Kelefence
propagation		
Callus induction	Leaves and nodal	Rathi (2017)
	segments	
Callus induction	Hypocotyl	Fu et al. (2010)
Multiple shoots	Nodal segment	Arya (2009)
differentiation		
Callus induction	Cotyledon	Wawrosch et al.
		(2009)
Callus induction	Young leaves	Mousa <i>et al</i> . (2007)

Considerable research efforts have been directed towards micro-propagation of mulethi using shoot tip and nodal culture techniques. In vitro rooting has proven to be most effective on full-strength MS medium enriched with IBA, with optimal levels ranging from 0.5 to 1.0 mg/l (Patel et al., 2007). Yadav and Singh (2012) provided a detailed micro-propagation protocol involving the manipulation of growth regulators, cultural conditions, and external factors influencing mulethi's in vitro proliferation. Optimal results were achieved by selecting middle-order nodes, specifically the 3rd to 5th node from the apex, resulting in the highest bud-break rate (86.6%), longest shoot length (8.0 cm), and the greatest number of shoots (3.0). Similarly, Fu et al. (2010) conducted a study focused on enhancing the development of embryogenic callus and embryogenesis in mulethi. They observed that using hypocotyl explants led to the highest frequency of callus formation, reaching 93.3%, when cultured on MS medium supplemented with 2.0 mg/L 6-benzylaminopurine (6-BA) and 0.5 mg/L 2,4-dichlorophenoxyacetic acid (2,4-D).

Extensive research efforts have been dedicated to its micropropagation using shoot tip and nodal culture techniques. *In vitro* rooting was found to be most effective on full-strength MS medium enriched with IBA at levels ranging from 0.5 to 1.0 mg/l (Patel *et al.*, 2007). Yadav and Singh (2012) detailed a specific micro-propagation protocol that involved the manipulation of growth regulators, cultural conditions, and external factors influencing the *in vitro* proliferation of *mulethi*. The most favorable outcomes were attained by selecting middle-order nodes (specifically, the 3rd to 5th node from the apex), which resulted in the highest bud-break rate (86.6%), longest shoot length (8.0 cm), and the greatest number of shoots (3.0). Similarly, Fu *et al.* (2010) conducted a study aimed at enhancing the development of embryogenic callus and embryogenesis in *mulethi*. They observed that using hypocotyl explants yielded the highest frequency of callus formation, reaching 93.3%, when cultured on Murashige and Skoog (MS) medium supplemented with 2.0 mg/L 6-benzylaminopurine (6-BA) and 0.5 mg/L 2,4-dichlorophenoxyacetic acid (2,4-D).

Propagation of Shatavari (Asparagus racemosus Willd.)

The National Medicinal Plant Board has identified *shatavari* as one of the 32 medicinal plants designated for conservation based on its extensive use (Thakur *et al.* 2015). Therefore, the preservation of this species is of utmost significance. They can be propagated through two main methods: seeds or divisions of rhizomatous discs (Sachan *et al.*, 2012).

Seed propagation

For sowing purpose seeds are typically harvested from March to May, specifically when they undergo a transition in color from red to black. As part of the pre-sowing treatment, these seeds can be soaked in water for duration of two days or subjected to gibberellic acid treatment for periods of 96 hours, which helps expedite the germination process (Gupta et al., 2002). In the initial week of June, seed sowing takes place in meticulously prepared elevated nursery beds, as illustrated in Fig. 4. The seeds are planted at a depth of 2 cm beneath the soil surface within raised beds measuring 4.5 m x 1.2 m and standing at a height of 20 cm. With a spacing of 5 cm between seeds in rows, they are carefully positioned and lightly covered with fine sand. Regular watering with a rose cane is employed to sustain optimal moisture levels in the beds. Typically, germination occurs within a span of 15 to 20 days. Approximately 7 kg of seeds are required to cultivate enough seedlings for one hectare of the crop (Krishnamurthy et al., 2004).

Transplanting

The soil ought to undergo thorough disc plowing, succeeded by harrowing and leveling processes. To facilitate irrigation management, the field is partitioned into separate plots with an irrigation channel positioned between every two rows of plots. About one month before the transplanting phase, it is advised to incorporate 10 tons of well-decomposed FYM into the soil, ensuring thorough mixing for optimal soil conditioning. August is ideal for transplanting. Seedlings are ready for transplantation about 45 days after sowing. Seedlings of 5 cm height are carefully removed from the beds, ensuring minimal root damage, and transferred to the field. During planting, they are placed into pits measuring 45 x 45 x 45 cm and spaced 1 meter apart (Saran *et al.*, 2020). In addition, seedlings are transplanted on ridges, maintaining a distance of 60 cm between plants, while ridges are 45 cm apart (Yadav *et al.*, 2022). The ridge method of transplanting

is superior in comparison to a flatbed method. The ideal number of seedlings required per hectare is approximately 150,000 plants/ha.





Propagation with crown bud

The propagation of the plant can be done by taking the crown bud along with its root. After transplanting sprouting commence after 8-10 days of planting. The yield of the crop is influenced by the number of plants per unit area. The highest yield was achieved with a plant density of 1x0.5 meters spacing, resulting in a dry yield of 12.5 kilograms per 24 square meters in irrigated areas during the final harvest at 19 months. In contrast, a lower plant density of 1x1 meter spacing yielded 9-9.4 kilograms per 24 square meters under similar conditions (Krishnamurthy *et al.*, 2004). The first harvest is usually done after 1.5-2 years of planting, and this can continue for 10-15 years (Saran *et al.*, 2019).

In vitro propagation

Shatavari can be grown from either seeds or other plant parts (like root cuttings) (Sachan *et al.*, 2012). However, since the roots are the valuable part used for medicine, people usually prefer seeds to grow the plant. But multiplication through seed has its drawbacks such as low germination percentage, slow growth and existence of heterozygosity (Singla and Jaitak, 2014; Sharan *et al.*, 2011). Hence, the requirement for consistent and high-quality planting material is of utmost importance. In this context, *in vitro* techniques can prove advantageous. Numerous studies have documented the standardization of micro-propagation protocols for *shatavari*. A compilation of the *in vitro* propagation methods employed in *shatavari* is presented in Table 3.

Table 3. In vitro regeneration of shatava	ri
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Mode of propagation	Explants used	References
Adventitious shoot bud regeneration	Nodal explant	Sulava <i>et al.</i> (2020)
Somatic embryoids for- mation through callus culture	Zygotic embryos	Chaudhary and Dantu (2019)
Somatic embryoids for- mation through callus culture	Nodal explant	Pant and Joshi (2017)
Adventitious shoot bud regeneration	Shoot apex and nodal explant	Sharan <i>et al</i> . (2011)
Indirect organogenesis via callus culture	Shoot explants (node, internode, shoot tips)	Pant and Joshi (2009)
Axillary branching	Nodal explant	Bopana and Saxe- na (2008)
Adventitious shoot bud regeneration through callus culture	Nodal explant	Kumar and Vijay (2008)

Kar and Sen (1985) pioneered the initial efforts to cultivate *Asparagus racemosus in vitro*. They successfully induced shoot proliferation from callus cultures of *shatavari* using stem discs as explants, with the aid of 2,4-D and kinetin.

Bopana and Saxena (2008) later devised a more effective protocol for axillary branching using nodal explants. Normally, *in vitro* rooting in *A. racemosus* is tedious and hormones are not enough to induce roots. However, this study successfully induced roots by employing phloroglucinol. The researchers utilized an MS medium supplemented with 3.69 μ M 2-isopentyl adenine and 3% sucrose, leading to a multiplication rate of 3.5.

Post-planting care of tuberous medicinal plants

Caring for tuberous medicinal plants after planting is crucial to ensure their growth, health, and maximum medicinal yield. Here are some important post-planting care steps to follow:

Manure and fertilizer

Root crops generally do not demand heavy fertilization; however, applying a balanced fertilizer or compost during the growing season proves beneficial in providing essential nutrients. Organic manures like FYM, vermicompost, and green manure can be utilized based on the specific requirements of the species (Gupta *et al.*, 2011).

Ashwagandha crops exhibit excellent responsiveness to organic manures. For this crop, approximately 10 to 12 tonnes of well-decomposed FYM or 1 to 1.5 tons of vermicompost per hectare are recommended during plantation. In soils with average fertility, supplementing with 15 kg of nitrogen (N) and 15 kg of phosphorus (P) per hectare is advantageous for achieving higher yields (Tuteja, 2022). In cases of poor fertility soils, it is advisable to increase the application to 40 kg of nitrogen (N) and 40 kg of phosphorus (P) per hectare to optimize root yield. Mulethi crops generally do not require heavy fertilization, but the application of a balanced fertilizer or organic compost once or twice during the growing season can enhance growth. Applying compost or FYM at the rate of 12-18 tons per hectare is sufficient to meet the nutrient requirements of the plant. However, due to difficulties in obtaining organic manures in large quantities, a chemical fertilizer dose at the rate of 40:40:20 kg per hectare per year (urea, superphosphate, and chloride of potash) may also be considered. The full basal dose of superphosphate and potash is applied at the time of planting, while nitrogen is applied in three split doses: at planting, six months later, and one year after planting. Since the crop remains in the field for two and a half to three years, every year the same dose has to be applied (Farooqi, 2001). Further, shatavari cultivation demands a fertilizer dose of 60 kg nitrogen, 40 kg phosphate, and 40 kg potash per hectare to achieve optimal growth and enhance tuberous root yield (Parida et al., 2018). It is recommended to place one-third of the nitrogen and the entire dose of phosphate and potash 10–12 cm deep in the rows before the transplanting process. This strategic placement ensures that the essential nutrients are available to support *shatavari* plants during their growth and development.

Irrigation

Tuberous medicinal plants are sensitive to excess water, and it is vital to avoid water-logging in the field. Stagnant water can result in problems like damping-off and root rot. To promote the successful establishment of seedlings in the soil, it is recommended to apply light irrigation at the time of transplanting. For optimal root yield, irrigation should be carried out at intervals of 8 to 10 days (Namdeo and Ingawale, 2021). However, during the dry summer season, the crop may require irrigation at intervals of 30-45 days, and in the winter season, one to two irrigations may be necessary to maintain root health. If there is regular rainfall, additional irrigation may not be needed. Mulethi plants, on the other hand, demand consistent moisture, especially during the growing season, typically requiring a total of 7-10 irrigations on average for successful crop cultivation (Zaidi, 2018). In contrast, shatavari requires minimal irrigation as it can thrive without frequent watering. An annual rainfall range of 800-1200 mm, distributed evenly, is generally sufficient for shatavari plants. During the initial establishment phase, plants may be irrigated weekly, and once they have firmly established themselves, light irrigation can be provided at monthly intervals (Saran et al., 2021).

Mulching

The mulching technique involves spreading a protective layer of organic materials, such as straw, leaves, wood chips, or compost over the soil surface around the base of the plants. It is an effective practice that offers multiple advantages. This practice aids in conserving moisture, regulating soil temperature, enhancing soil health, suppressing weeds, and mitigating erosion (Carrubba and Militello, 2013).

Weeding

Keep the area weed-free, as weed competition can impede their development. During the initial year of planting, it is essential to conduct three to four hoeing cum weeding sessions. In the following years, two hand weeding-cumhoeing sessions prove adequate to maintain a weed-free field and enhance the overall well-being of the plants. Typically, the first weeding should occur within 21–25 days of sowing, with the second weeding scheduled after an additional 21–25 days following the initial one (Carrubba and Militello, 2013).

Staking

Ashwagandha, mulethi and shatavari may require support as they grow taller. Staking is crucial to counter potential damage from strong winds or heavy rainfall. Ashwagandha plants can grow up to 4 to 5 feet in height, and their branches may need some support. Consider staking or using a trellis to help the plants stay upright (Mathew *et al.*, 2005). Likewise, *mulethi* plants can reach heights of 3 to 5 feet and may need support as they mature (Bakhane *et al.*, 2014). It is worth noting that *shatavari* crops, being climbers, also necessitate support for their optimal growth (Babu *et al.*, 2016). To facilitate this, 4-6 feet long stakes are commonly employed to provide the necessary support for the overall growth of the plants.

Pest and disease control

Major pests and diseases have not been reported in medicinal crops; however, early detection and timely intervention can effectively prevent any potential serious damage. It is advisable to employ natural pest control methods whenever possible. Bio-pesticides, either prepared individually or as a mixture from plant parts such as neem, chitrakmool, dhatura, and cow's urine, can be utilized for this purpose (Gupta *et al.*, 2011). Implementing this proactive strategy helps to ensure the overall health of root crops.

Harvesting

Ashwagandha roots are typically harvested between January and March, approximately 150 to 180 days after sowing, when they reach the desired size. The maturity of the crop is determined by the drying out of leaves and the presence of yellow-red berries (Moharana et al., 2020; Saran et al., 2025). It is recommended to maintain adequate soil moisture during the digging process. Careful extraction of the tap root is essential to preserve the health of even the smaller lateral roots. A yield of 6.5 to 7.0 quintals per hectare can be achieved in fibrous type, with a preference for commercially viable root species with diameters ranging from 6 to 15 mm and lengths of 7 to 10 cm. DTWr-1 has maximum fresh root length (36-39 cm), fresh root girth (2-4 cm), dry root yield (5 t h⁻¹) and starch yield (2.70 t h⁻¹), as compared to all available checks as shown in Fig. 5 (Saran and Das, 2023; Saran et al., 2025). The alkaloid content in the roots varies from 0.13 to 0.31%.

Mulethi crops are ready for harvesting after 2.5-3 years of planting, just before fruiting, typically in November and December, when glycyrrhizin content is at its peak (Ahmed

et al., 2021). The plants are uprooted, and the aerial parts are removed, leaving clean roots. Herbage may be left in the field to recycle minerals and nutrients. The Haryana Mulhatti-1 variety yields 7-8 tons of roots with a 7.5 percent glycyrrhizin content (Zaidi, 2018). *Shatavari* crops mature within 12 to 14 months after planting (Saran *et al.*, 2019) and are expected to yield approximately 4–5 tonnes per hectare, depending on soil and climatic conditions (Fig. 6). The optimal period for harvesting tuberous roots is in November to December, coinciding with the time when the above-ground parts of the plant begin to exhibit a pale-yellow color (Vel, 2017).



Fig. 5. Freshly harvested roots of starchy type Ashwagandha

Conclusion

The medicinal tubers are a vital group of medicinal plants. Their commercial value often leads to overexploitation, which threatens their natural populations. To address this issue, we have been exploring various propagation methods, including *in vitro* plant propagation. The review aims to provide a comprehensive overview of existing studies and investigations related to the propagation of medicinal tuber plants. This not only helps consolidate existing knowledge but also identifies research gaps that require further exploration. This approach aligns with a more holistic and forward-looking strategy to ensure the survival of these important medicinal plants for future generations. In this way, farmers will be encouraged to cultivate these medicinal plants for commercial use, which will curb overexploitation in the wild and thus complement the conservation process.



Fig. 6. Harvesting of Shatavari

Acknowledgements

The authors express their gratitude to the Indian Council of Agricultural Research (ICAR), New Delhi, and ICAR-Directorate of Medicinal and Aromatic Plants Research, Anand for providing the necessary facilities to conduct research and compile the information.

Author Contribution Statement

SWARAJYA LAXMI NAYAK: Writing original draft. Compilation; PARMESHWAR LAL SARAN: Compilation & manuscript editing; MANISH DAS: manuscript editing

Conflict of Interest

The authors have no conflict of interest.

References

- Ahmed, S. F., Sajid, M., Ahmad, W., Zeenat, F. and Shakir, M. 2021.
 A comprehensive review on an important Unani drug mulethi (Root of *Glycyrrhiza glabra* Linn). *Journal of Pharmacognosy and Phytochemistry*, 10(3): 488-493.
- Ahuja, A., Tripathi, M. K., Tiwari, S., Tripathi, N., Tiwari, G., Mishra, N., Bhargav, S. and Tiwari, S. 2021. Recent advancements on callus and cell suspension cultures: An effectual reserve for the production of pharmaceutically significant metabolites. *Current Aspects in Pharmaceutical Research and Development*, 6: 96-111.
- Akhtar, J., and Anjum, N., 2022. *Glycyrrhiza glabra*. Herbs, Shrubs, and Trees of Potential Medicinal Benefits, 133-145. CRC Press

- Alok, S., Jain, S. K., Verma, A., Kumar, M., Alok, M. and Sabharwal, M. 2013. Plant profile, phytochemistry and pharmacology of *Asparagus racemosus* (Shatavari): a review. *Asian Pacific Journal of Tropical Disease*, 3(3): 242-251. https://doi.org/10.1016/ S2222-1808 (13)60049-3
- Ameen, G., Tirkey, A. and Sandilya, V. K. 2023. The past, present and future aspects of *Withania somnifera* (L.) Dunal breeding: A review. *The Pharma Innovation Journal*, 12(7): 1928-1935.
- Arya, S., Rathi, N. and Arya, I. D. 2009. Micropropagation protocol for *Glycyrrhiza glabra* L. *Phytomorphology*, 59: 71-76.
- Autade, R. H., Fargade, S. A., Savant, A. R., Gangurde, S. S., Choudhary, R. S. and Dighe, S. S. 2016. Micropropagation of Ashwagandha (*Withania somnifera*). *Bioscience Biotechnology Research Communications*, 9: 88-93.
- Babu, N., Srivastava, S. K., Prusty, M. and Sahoo, T. 2016. Medicinal and aromatic plant production technologies: A step towards farm women prosperity. Technical Bulletin No. 28
- Bakhane, Y., Yadav, A. S., Bajaj, A., Sharma, A. K. and Raghuwanshi, D. K. 2014. *Glycyrrhiza glabra* L.: A Miracle Medicinal Herb. *Indo American Journal of Pharmaceutical Research*, 4(12): 5808-16.
- Bhardwaj, R., Sharma, K., Sharma, D. K. and Prakash, P. 2020. Conventional genetic improvement methods in medicinal and aromatic plants: A review. *International Journal of Economic Plants*, 7(4): 156-161. https://doi.org/10.23910/2/2020.0382
- Bopana, N. and Saxena, S. 2007. Asparagus racemosus-Ethnopharmacological evaluation and conservation needs. Journal of Ethnopharmacology, 110: 1-15. https://doi.org/10.1016/j. jep.2007.01.001
- Bopana, N. and Saxena, S. 2008. In vitro propagation of a high value medicinal plant: Asparagus racemosus Willd. In Vitro Cellular and Developmental Biology, 44: 525–532. https://doi. org/10.1007/s11627-008-9137-y
- Carrubba, A. and Militello, M. 2013. Nonchemical weeding of medicinal and aromatic plants. *Agronomy for Sustainable Development*, 33: 551-561. https://doi.org/10.1007/s13593-012-0122-9

- Chaudhary, D., Giri, S. and Lamichhane, G. 2023. Concise review on *Asparagus racemosus* and its role as nutraceuticals and functional foods. Herbs, Spices and their Roles in Nutraceuticals and Functional Foods, 245-255. Academic Press. https:// doi.org/10.1016/B978-0-323-90794-1.00004-1
- Chaudhary, J. and Dantu, P. K. 2019. Induction of somatic embryos in cultures of *Asparagus racemosus* Willd: an endangered medicinally important plant. *Bulletin of the National Research Centre*, 43: 1-5. https://doi.org/10.1186/s42269-019-0157-z
- Chaurasiya, R. K., Chand, P. and Chauhan, P. S. 2019. Chapter-7 Challenges and Opportunities in the Scientific Cultivation of *Withania somnifera*. Chief Editor, 159-179.
- Cui, X., Lou, L., Zhang, Y. and Yan, B. 2023. Study of the distribution of *Glycyrrhiza uralensis* production areas as well as the factors affecting yield and quality. *Scientific Reports*, 13(1): 5160-72. https://doi.org/10.1038/s41598-023-31946-5
- Das, M., Vanita, J. and Malhotra, S. K. 2016. Impact of climate change on medicinal and aromatic plants. *Indian Journal of Agricultural Sciences*, 86(11): 1375-1382. https://doi. org/10.56093/ijas.v86i11.62865
- Das, M. and Pandey, S. 2023. ICAR-DMAPR's classical journey in medicinal and aromatic plant research and advancement. *In-dian Horticulture*, 68(5): 4-10.
- Dastagir, G. and Rizvi, M. A. 2016. *Glycyrrhiza glabra* L. (Liquorice). *Pakistan Journal of Pharmaceutical Sciences*, 29(5): 1727-1733.
- Farooqi, A. A. and Sreemaru, B. S. 2001. Cultivation of Medicinal and Aromatic Crops, Hyderabad: University Press (India) Limited.
- Fatima, N., Ahmad, N., Anis, M. and Ahmad, I. 2013. An improved in vitro encapsulation protocol, biochemical analysis and genetic integrity using DNA based molecular markers in regenerated plants of *Withania somnifera* L. *Industrial Crops and Products*, 50: 468-477. https://doi.org/10.1016/j.ind-crop.2013.08.011
- Fu, C., Lei, C., Gan, L., Li, M., Yang, Y. and Yu, L. 2010. Optimization of embryogenic-callus induction and embryogenesis of *Glycyrrhiza glabra*. *African Journal of Biotechnology*, 9(36): 5823–5829.
- Govindaraju, B., Rao, S. R., Venugopal, R. B., Kiran, S. G., Kaviraj, C. P., Srinath, R. 2003. High frequency plant regeneration in Ashwagandha [Withania somnifera (L.) Dunal] - An important medicinal plant. Plant Cell Biotechnology and Molecular Biology, 4: 49-56.
- Gupta, M., Duhan, P., Kajla, S. and Chaudhury, A. 2014. Optimization of media for establishment of *Glycyrrhiza glabra* micropropagation. *Annals of Agri Bio Research*, 19: 197–201.
- Gupta, S., Kumar, A. and Sharma, S. N. 2002. Improvement of seed germination in Asparagus racemosus Willd. Journal of Herbs, Spices & Medicinal Plants, 9(1): 3-9. https://doi.org/10.1300/ J044v09n01_02

Gupta, S., Singh, R. and Ashwlayan, V. D. 2011. Pharmacological activity of *Tiospora cordifolia*. *Pharmacology online*, 1: 644-52.

- Gupta, R., 2017. Agrotechnology of medicinal plants. *In:* The medicinal plant industry (pp. 43-58). Routledge.
- Jaiswal, N., Verma, Y. and Misra, P. 2017. Micropropagation and *in vitro* elicitation of licorice (*Glycyrrhiza* spp.). *In Vitro Cellular & Developmental Biology-Plant*, 53: 145-166. https://doi. org/10.1007/s11627-017-9832-7
- Kar, D. K., and Sen, S. 1985. Micropropagation of Asparagus racemosus. Plant Cell, Tissue and Organ Culture, 5: 89–95.
- Kaur, K., Singh, P., Kaur, K., Bhandawat, A., Nogia, P. and Pati, P. K. 2021. Development of robust in vitro culture protocol for the propagation of genetically and phytochemically stable plants of Withania somnifera (L.) Dunal (Ashwagandha). Industrial Crops and Products, 166: 113428. https://doi.org/10.1016/j.indcrop.2021.113428
- Khabiya, R., Choudhary, G. P., Jnanesha, A. C., Kumar, A. and Lal, R. K. 2023. An insight into the potential varieties of Ashwagandha (Indian ginseng) for better therapeutic efficacy. Acta Ecologica Sinica. https://doi.org/10.1016/j.chnaes.2023.06.009
- Krishnamurthy, R., Chandorkar, M. S., Kalzunkar, B. G., Palsuledesai, M. R., Pathak, J. M., Gupta, R. 2004. Agronomic manipulation for improving the productivity of shatavari (*Asparagus racemosus*) under irrigated and rainfed conditions. *Journal of Medicinal and Aromatic Plant Sciences*, 26: 740-745.
- Kumar, A. and Vijay, N. 2008. In Vitro Plantlet Regeneration in Asparagus racemosus Through Shoot Bud Differentiation on Nodal Segments. In: Recent Advances in Plant Biotechnology and Its Applications, Ashwani, K., Sopory, S. and Neumann, K.H. (Eds.), I.K. International Pvt. Ltd., New Delhi, pp. 185-197.
- Kumar, A., Venugopal, S., Jnanesha, A. C. and Lal, R. K. 2023. Agricultural-based challenges, genetic enhancement, and obstacles to an industrially important medicinal plant, ashwagandha (*Withania somnifera* (L.) Dunal): A review. *Ecological Genetics and Genomics*, 100183. https://doi.org/10.1016/j. egg.2023.100183
- Mathew, G., Joy, P. P., Skaria, B. P. and Mathew, S. 2005. Cultivation prospects of tuberous medicinal plants. In National Seminar on Achievements and Opportunities in Postharvest Management and Value Addition in Root and Tuber Crops (NSRTC 2), Centre for Tuber Crops Research Institute (CTCRI), Thiruvananthapuram, India (pp. 81-82).
- Mehdizadeh, L. and Moghaddam, M. 2023. Hydroponic System for Cultivation of Medicinal Plants. In Biosynthesis of Bioactive Compounds in Medicinal and Aromatic Plants: Manipulation by Conventional and Biotechnological Approaches (pp. 213-233). Cham: Springer Nature Switzerland.
- Mir, B. A., Mir, S. A. and Koul, S. 2014. *In vitro* propagation and withaferin A production in Withania ashwagandha, a rare medicinal plant of India. *Physiology and Molecular Biology of*

Plants, 20: 357-364. https://doi.org/10.1007/s12298-014-0243-5

- Moharana, D., Bahadur, V., Rout, S., Prusty, A. K. and Sahoo, R. K. 2020. Ashwagandha: The Miracle Ginseng. *Food and Scientific Reports*, 1: 37-42.
- Mousa, N. A., Siaguru, P., Wiryowidagdo, S. and Wagih, M. E. 2007. Regenerative callus and cell suspension system of licorice (*Glycyrrhiza glabra*) for the production of the sweetener glycyrrhizin *in vitro*. *International Journal of Sugar Crops and Related Industries*, 9: 72–82. https://doi.org/10.1007/BF02956917
- Namdeo, A. G. and Ingawale, D. K. 2021. Ashwagandha: Advances in plant biotechnological approaches for propagation and production of bioactive compounds. *Journal of Ethnopharmacology*, 271: 113709. https://doi.org/10.1016/j.jep.2020.113709
- Niraj, S. and Varsha, S. 2020. A review on scope of immuno-modulatory drugs in Ayurveda for prevention and treatment of Covid-19. *Plant Science Today*, 7(3): 417-423. https://doi. org/10.14719/pst.2020.7.3.831
- Öztürk, M., Altay, V., Hakeem, K. R. and Akçiçek, E. 2018. Liquorice: from botany to phytochemistry, pp127-132. Springer.
- Pant, K. K. and Joshi, S. D. 2009. In vitro multiplication of wild Nepalese *Asparagus racemosus* through shoots and shoot induced callus cultures. *Botany Research International*, 2: 88-93.
- Pant, K. K. and Joshi, S. D. 2017. *In vitro* morphogenesis of shoot induced callus of *Asparagus racemosus* willd. into somatic embryoids. *International Journal of Scientific and Technology Research*, 2(10): 637-641.
- Parida, N., Mishra, L., Sinha, C. P. and Sahu, A. K. 2018. A review on conventional cultivation and propagation of some of the threatened and endangered wild medicinal plants having commercial importance. *World Journal of Pharmaceutical Research*, 7(11): 1247-1254.
- Rao, K. V. S. 1993. A review on Licorice. Ancient Science of Life, 13(1-2): 57-88.
- Rathi, N., Rao, N. N., Dwivedi, S., Arya, S. and Arya, I. D. 2017. Optimization of media constituents for regeneration via callus cultures of *glycyrrhiza glabra* l. an endangered plant. *Biochemical & Cellular Archives*, 17(1): 389-398.
- Ratre, R. K., Kulmitra, A. K. and Shankar, G. 2018. Economic of ashwagandha as influenced by sowing methods and organic sources of nutrients. *International Journal of Chemical Studies*, 6(5): 1973-1974.
- Sachan, A. K., Das, D. R., Dohare S. L. and Shuaib, M. 2012. *Asparagus racemosus* (Shatavari): an overview. *International Journal of Pharmaceutical and Chemical Sciences*, 1(3): 588-592.
- Saha, S., Mandal, A. and Dutta, A. 2018. Good Agricultural Practices: Requirement for the Production of Quality Herbal Medicines. *Natural Products and Drug Discovery*, pp. 607-631. https://doi.org/10.1016/B978-0-08-102081-4.00022-8
- Saran, P. L., Singh, S., Solanki, V. H., Kalariya, K. A., Meena, R. P. and Patel, R. B. 2019. Impact of shade-net intensities on root

yield and quality of *Asparagus racemosus*: A viable option as an intercrop. *Industrial Crops and Products*, 141: 111740. https://doi.org/10.1016/j.indcrop.2019.111740

- Saran, P. L., Singh, S., Solanki, V., Choudhary, R. and Manivel, P. 2021. Evaluation of Asparagus adscendens accessions for root yield and shatavarin IV content in India. *Turkish Journal of Agriculture and Forestry*, 45(4): 475-483. https://doi.org/10.3906/ tar-2006-42
- Saran, P. L., Singh, S., Solanki, V. H., Devi, G., Kansara, R. V. and Manivel, P. 2020. Identification of potential accessions of *Asparagus racemosus* for root yield and shatavarin IV content. *Heliyon*, 6(12). https://doi.org/10.1016/j.heliyon.2020.e05674.
- Saran, P.L., Kumar, S., Sarkar, R., Kalariya, K.A., Reddy, N.R. and Das, M. 2025. Identification of elite accession of Ashwagandha (*Withania somnifera* L. Dunal) for root yield and brittleness. *Industrial Crops & Products*, 223: 120161; https://doi. org/10.1016/j.indcrop.2024.120161.
- Saran, P.L. and Das, M. 2023. Nagori ashwagandha: a viable alternative crop for arid ecosystem. *Indian Horticulture*, 68 (5): 22–24.
- Saran, P.L. 2023. Nagori Ashwagandha' (*Withania somnifera* L. Dunal) a miracle root herb for arid region of India: a mini review. *JONAM*, 7 (3): 000409.
- Sen, S., Chakraborty, R. and De, B. 2011. Challenges and opportunities in the advancement of herbal medicine: India's position and role in a global context. *Journal of Herbal Medicine*, 1(3-4): 67-75. https://doi.org/10.1016/j.hermed.2011.11.001
- Sharada, M. 2004. Studies on regeneration and phytochemical potential of *Withania somnifera* (L.) Dunal tissue cultured *in vitro*. Ph. D. Thesis, Panjab University, Chandigarh.
- Sharan, M., Nene, C. and Sharon, M. 2011. Regeneration of *Asparagus racemosus* by shoot apex and nodal explants. *Asian Journal of Plant Science & Research*, 1: 49-56.
- Sharma, P., Tripathi, M. K., Tiwari, G., Tiwari, S. and Baghel, B. S. 2010. Regeneration of liquorice (*Glycyrrhiza glabra* L.) from cultured nodal segments. *Indian Journal of Plant Physiology*, 15(1): 1.
- Singh, N., Yadav, S. S., Rao, A. S., Nandal, A., Kumar, S., Ganaie, S. A. and Narasihman, B. 2021. Review on anticancerous therapeutic potential of *Withania somnifera* (L.) Dunal. *Journal of Ethnopharmacology*, 270: 113704. https://doi.org/10.1016/j. jep.2020.113704
- Singla, R. and Jaitak, V. 2014. Shatavari (*Asparagus racemosus* wild): a review on its cultivation, morphology, phytochemistry and pharmacological importance. *International Journal of Pharmacy & Life Sciences*, 5(3): 742-757.
- Srivastava, M., Singh, G., Sharma, S., Shukla, S. and Misra, P. 2019. Elicitation enhanced the yield of glycyrrhizin and antioxidant activities in hairy root cultures of *Glycyrrhiza glabra* L. *Journal of Plant Growth Regulation*, 38: 373-384. https://doi. org/10.1007/s00344-018-9847-2

- Sulava, S., Andia, S., Bhol, R. and Jena, S. 2020. *In vitro* micropropagation of asparagus racemosus by using of nodal explants. *Journal of Cell and Tissue Research*, 20(1): 6883-6888.
- Tawade, S., Labade, G., Dale, N. and Varpe, S. N. 2023. In vitro micro-propagation of Ashwagandha (*Withania somnifera*). *Bioinfolet-A Quarterly Journal of Life Sciences*, 20(2b): 300-302.
- Tewari, S. K. and Singh, S. 2019. Agro-technologies for cultivation of medicinal plants. *Medicinal Plants*, 22: 22-151.
- Thakur, S., Kishan, K. L. and Jadhav, S. K. 2015. Approaches for Conservation of an Ethnomedicinal Plant: *Asparagus racemosus* Willd. *Journal of Biological Sciences*, 15 (3): 126.133.
- Tuteja, K. M. D. S. 2022. Effect of different nutrient management practices on growth and yield of ashwagandha (*Withania somnifera* L.). *The Pharma Innovation Journal*, 11(12): 3974-3977.
- Vel, S. V. 2017. http://vikaspedia.in/agriculture/crop-production/ package-of-practices/satavari.pdf (Accessed 8 August, 2023).

- Verma, S. K., Rai, M. K., Asthana, P., Jaiswal, V. S. and Jaiswal, U. 2010. *In vitro* plantlets from alginate-encapsulated shoot tips of *Solanum nigrum* L. *Scientia Horticulturae*, 124(4): 517-521. https://doi.org/10.1016/j.scienta.2010.02.002
- Wawrosch, C., Kazianka, C., Kuchler, V., Lammermayer, K., Winter, M. and Kopp, B. 2009. *In vitro* multiplication of *Glycyrrhiza glabra* L. through somatic embryogenesis. *Planta Medica*, 75: 1036–1037. https://doi.org/10.1055/s-0029-1234833
- Yadav, K. and Singh, N. 2012. Factors influencing *in vitro* plant regeneration of liquorice (*Glycyrrhiza glabra* L.) *Iranian Journal of Biotechnology*, 10: 161–167.
- Yadav, R., Choubey, A. and Mishra, M. A. 2022. Industrial Pharmacognosy. Booksclinic Publishing. Pp. 153.
- Zaidi, T. H. 2018. Cultivating mulethi (*Glycyrrhiza glabra* Linn) in haryana: an agroecological approach. *Advance and Innovative Research*, 5(3): 46-54.