

# Glandular trichomes: the factory of artemisinin biosynthesis

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## Abstract

Plant glandular trichomes serve as crucial secretory organs, responsible for the production, modification, and storage of various vital secondary metabolites, including medicinal natural products, while also playing a role in plant defense mechanisms. *Artemisia annua* L. (*A. annua*) stands as a significant herb in traditional Chinese medicine because of its primary active compound, artemisinin, which is the most potent antimalarial agent. The glandular trichomes of *A. annua* are closely related to the synthesis and accumulation of artemisinin. As research in medicinal plants progresses, the significance of glandular trichomes in the study of plant secondary metabolites has notably escalated. Given their pivotal role in artemisinin biosynthesis, the growth and development of *A. annua* glandular trichomes directly influence the yield and quality of artemisinin. This paper presents a comprehensive review of the latest research advancements regarding *A. annua* glandular trichomes, encompassing their morphology, function, growth, and developmental influencing factors, and the artemisinin biosynthesis pathway. The aim is to provide substantial support for ongoing investigations into the growth and development of *A. annua* glandular trichomes and the underlying mechanisms of metabolic regulation.

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## Introduction

The majority of terrestrial plants have plant epidermal trichomes, a specific structure of epidermal cells that extends from the outside of organs, including leaves, stems, flowers, and fruits<sup>[1]</sup>. Plant trichomes act as a natural barrier between plants and their surroundings, keeping pathogens, insects, herbivores, and other invaders out. Plants utilize it as their first line of defense against outside invasion<sup>[2,3]</sup>. Trichomes vary in size, shape, number of cells, origin, location, function, etc. They can be divided into long and short types based on their morphological characteristics<sup>[4]</sup>. Based on the shape of the glandular head cells, they can be categorized as stellate, chained, squamous, etc. Additionally, trichomes are classified into non-secretory glandular trichomes and secretory glandular trichomes based on their secretion function<sup>[5]</sup>. Many plants, including *A. annua*, *Cannabis sativa* L., *Mentha haplocalyx* Briq., *Pogostemon cablin* (Blanco) Benth., *Perilla frutescens* (L.) Britt., and *Nicotiana tabacum* L., have secretory or non-secretory glandular trichomes<sup>[6–11]</sup>.

A unique, glandular-like appendage that develops in the epidermal tissue of aboveground organs in plants is referred to as a non-secretory glandular trichome. Non-secretory glandular trichomes have various functions, including stress tolerance and defense<sup>[1,12]</sup>, and they shield plant growth and organs. Plants use it as a strong defense against both biotic and abiotic stress. Malvaceae plants, such as cotton<sup>[13]</sup>, and Brassicaceae plants, including *Arabidopsis*<sup>[14]</sup>, both have a large distribution of non-secretory glandular trichomes. Secretory glandular trichomes, being observed on approximately 30% of all vascular plant species<sup>[15]</sup>, are characterized by their metabolic

capacity to produce, store, and secrete copious secondary metabolites and thus earn them the moniker 'biosynthetic factories'<sup>[16]</sup>. Some plant secretory glandular trichomes, from the Asteraceae, Lamiaceae, Cannabaceae, and Solanaceae families have been found to contain natural compounds such as terpenoids, alkaloids, phenols, and flavonoids<sup>[17,18]</sup>.

*A. annua*, a member of the Asteraceae family and an annual herb of the *Artemisia* genus, possesses a bitter and pungent taste and is characterized by its cold and cool properties. It effectively clears deficiency heat, eliminates bone steaming, relieves heat, and intercepts malaria<sup>[19]</sup>. The glandular trichomes of *A. annua* predominantly occur on the leaves, flowers, and flower buds<sup>[20]</sup>. Notably, the secretory glandular trichomes of *A. annua* serve as the primary site for the production and accumulation of artemisinin<sup>[20–22]</sup>. Artemisinin is one of the most effective drugs for the treatment of malaria, and artemisinin-based combination therapies (ATC) are the most effective and important means of treating malaria nowadays<sup>[23,24]</sup>. In addition to their classical antimalarial effects, artemisinin and its derivatives exhibit a wide range of biological activities, including antibacterial, antioxidant, anti-inflammatory, antiviral, anticancer, and cardioprotective effects<sup>[25–28]</sup>. Recent scientific studies have revealed that artemisinin derivatives are highly effective in improving the disease manifestations of polycystic ovarian syndrome (PCOS), a discovery that opens new avenues for the treatment of PCOS<sup>[29]</sup>. These diverse pharmacological effects indicate that artemisinin and its derivatives have great potential and broad application prospects for assisting in the treatment of various diseases and the development of new drugs. Artemisinin, a significant contribution of traditional Chinese medicine to the world serves as a frontline

antimalarial drug, saving millions of lives annually. With the emergence of new indications, the market demand for artemisinin is increasing. The content of artemisinin is tightly associated with its secretory glandular trichomes. Nevertheless, secretory glandular trichomes are prone to lose and make up only 2% of the dry weight of the plant. Only 0.01% to 0.8% of the dry weight of *A. annua* plants consists of artemisinin<sup>[30]</sup>. In the face of the broad prospects for the clinical application of artemisinin and its growing market demand, how to effectively increase its production has become an important issue that needs to be tackled by the scientific research community and the industry. This review systematically elaborates on the morphology and function of *A. annua* glandular trichomes, factors affecting the growth and development of glandular trichomes, and the synthesis mechanism of artemisinin in secretory glandular trichomes to promote the cultivation of high-quality *A. annua* varieties and improve artemisinin production.

## Morphological characteristics and biological functions of trichomes in *A. annua*

The appearance, size, and cell content of the many types of glandular trichomes in *A. annua* vary, and observations of these trichomes date back to the 20<sup>th</sup> century. There are two types of glandular trichomes in *A. annua*, namely glandular secreting trichomes (AaGSTs) and T-shaped non-glandular trichomes (AaTNGs), both of which are multicellular head-shaped glandular trichomes<sup>[31,32]</sup> (Fig. 1). These two varieties of glandular trichomes are mainly distributed in the leaves, flowers, stems, buds, and other parts of *A. annua*, and play a specific biological function<sup>[33,34]</sup>.

AaGST is a 10-cell double-column semi-transparent structure consisting of two basal cells (Ba), two stalk cells (St), four subapical cells (Suc), two apical cells (Ac), and one subcutaneous space (SS)<sup>[20,35]</sup>, can synthesize, secrete, and store rich metabolites such as terpenes, alkaloids, catechol tannins, polysaccharides, and calcium oxalate. Artemisinin is produced in the apical and periapical cells of AaGSTs and accumulates in the subepidermal space of AaGSTs<sup>[35,36]</sup>. AaGSTs release harmful chemicals to insects and animals, thereby blocking their further

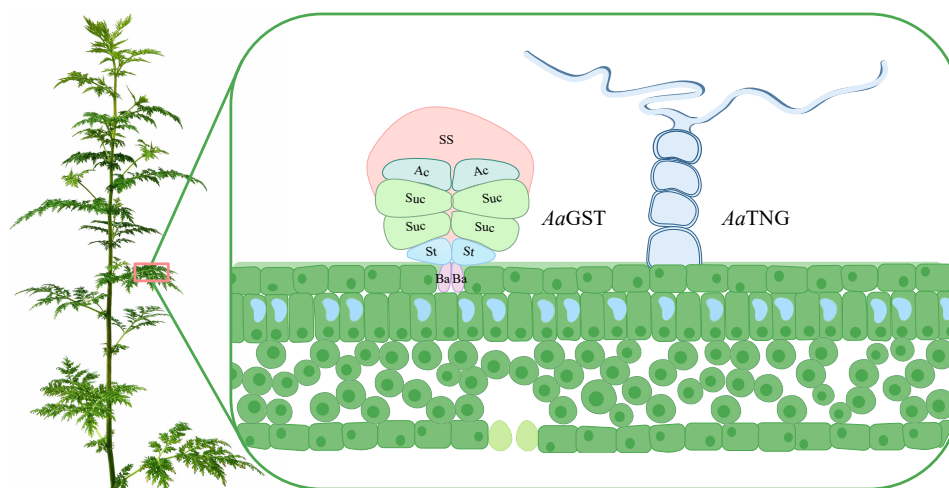
damage to plant bodies. Studies have shown that artemisinin has a certain resistance to parasites<sup>[37]</sup>. Protrusions that separate from plant epidermal cells and do not possess secretory activity are referred to as non-secretory glandular trichomes. AaTNG is a five-cell structure with elongated filamentous cells at the head, also known as T-shaped trichomes because of their similarity to the letter 'T'<sup>[20,36,38]</sup>. AaTNGs are widely found on the epidermis of leaves, stems, and flowers of *A. annua*<sup>[31]</sup>, and generally play a protective role for the plant against various biotic and abiotic stresses, with the function of preventing insects from nibbling, preventing low-temperature infestation, and preventing animals from attacking the plant<sup>[39]</sup>.

## Factors affecting trichome development in *A. annua*

The growth and development of *A. annua* trichomes are intricately coordinated and regulated by a multitude of factors, including regulatory genes, phytohormones, microRNAs, and the environment (Table 1). Among these factors, regulatory genes including transcription factors (TFs) and functional genes play a dominant role in the initiation, growth, and development of trichomes, whereas both plant hormones and microRNAs finally affect the formation and development of trichomes through the regulation of trichome-related genes (Fig. 2).

### Transcription factors

TFs can activate or repress the transcription of target genes by binding to specific regulatory sequences of the target genes, and play a crucial role in plant growth and development, response to adversity and secondary metabolite biosynthesis<sup>[62]</sup>. Previous studies have shown that many TF families are involved in the initiation, growth, and development of AaGSTs, with a particular focus on the R2R3-MYB, HD-ZIP, AP2/ERF, and WRKY. Additionally, because AaGSTs are closely linked to artemisinin biosynthesis, the TFs that regulate the growth and development of AaGSTs often also control the expression of key enzyme genes in the artemisinin biosynthesis pathway. These key enzymes include amorpha-4,11-diene synthase (ADS), artemisinic aldehyde double bond reductase



**Fig. 1** Morphological characteristics of trichomes in *A. annua*. AaGST, glandular secreting trichome; Ba, basal cells; St, stalk cells; Suc, subapical cells; Ac, apical cells; SS, subcutaneous space; AaTNG, T-shaped non-glandular trichome.

**Table 1.** The main factors regulating the growth and development of *A. annua* trichomes.

Types	Name	Effect	Target gene	Interacting proteins	Refs
R2R3-MYB	AaMYB1	(+)AaGST initiation (+)artemisinin biosynthesis	<i>ADS, CYP71AV1</i>		[40]
	AaMYB5	(-)AaGST formation (-)artemisinin biosynthesis		AaHD1, AaMYB16, AaJAZ8	[41]
	AaMYB16	(+)AaGST initiation (+)AaTNG initiation (+)artemisinin biosynthesis		AaHD1, AaMYB5	[41]
	AaMYB17	(+)AaGST initiation (+)artemisinin biosynthesis			[42]
	AaMIXTA1	(+)AaGST initiation (+)AaTNG initiation (+)artemisinin biosynthesis	<i>AaCYP77A1, AaCYP86A1, AaKCS5, AaCER1, AaABCG12</i>		[43]
	AaTLR1	(-)AaGST formation (-)artemisinin biosynthesis	<i>ADS, CYP71AV1, DBR2, ALDH1</i>	AaTLR2, AaWOX1	[44]
	AaTLR3	(-)AaGST formation (-)artemisinin biosynthesis	<i>ADS, CYP71AV1, DBR2, ALDH1</i>	AaCycTL	[45]
	AaTAR2	(+)AaGST initiation (+)AaTNG initiation (+)artemisinin biosynthesis	<i>ADS, CYP71AV1, DBR2, ALDH1</i>		[46]
	AaMYB108-like	(+)AaGST initiation (+)AaTNG initiation (+)artemisinin biosynthesis	<i>AaHD1</i>	AaHD8	[47]
		(+)AaGST initiation (+)AaTNG initiation (+)artemisinin biosynthesis	<i>AaGSW2</i>	AaJAZ8	[34]
HD-ZIP	AaHD1	(+)AaGST initiation (+)AaTNG initiation (+)artemisinin biosynthesis	<i>AaGSW2</i>	AaJAZ8	[34]
	AaHD8	(+)AaGST initiation (+)AaTNG initiation (+)artemisinin biosynthesis	<i>AaCYP86A4, AaFDH</i>	AaMIXTA1	[48]
AP2/ERF	AaTAR1	(+)AaGST formation (+)AaTNG formation (+)artemisinin biosynthesis	<i>ADS, CYP71AV1</i>	AaCycTL	[38]
	AaWIN1	(+)AaGST formation (+)artemisinin biosynthesis		AaGSW2 AaMIXTA1	[49]
	AaSPL9	(+)AaGST initiation (+)artemisinin biosynthesis	<i>AaGSW2</i>	AaHD1	[50]
WRKY	AaGSW2	(+)AaGST initiation (+)artemisinin biosynthesis		AaHD1, AaHD8	[51]
SAP	AaSAP1	(+)AaGST formation (+)artemisinin biosynthesis			[52]
LEAFY-like	AaTLR2	(-)AaGST initiation (-)AaTNG initiation (-)artemisinin biosynthesis	<i>ADS, CYP71AV1, DBR2, ALDH1</i>	AaWOX1	[44]
MADS-box	AaSEP1	(+)AaGST initiation (+)artemisinin biosynthesis	<i>AaGSW2</i>	AaMYB16 AaJAZ8	[53]
YABBY	AaYABBY5	(+)AaGST formation (+)artemisinin biosynthesis	<i>ADS, CYP71AV1, DBR2, ALDH1</i>		[54]
bHLH	AaGL3-like	(+)AaGST formation (+)AaTNG formation (+)artemisinin biosynthesis		AaJAZ8	[55]
Phytohormones	JA	(+)AaGST formation (+)AaTNG formation (+)artemisinin biosynthesis		AaHD1, AaHD8, AaJAZ8, AaMYB5	[34,41,56,57]
		(+)AaGST formation(GA3) (+)AaGST formation(GA1/4) (+)artemisinin biosynthesis(GA1/4)			[56,58]
	GA	(+)AaGST formation (+)AaTNG formation (+)artemisinin biosynthesis			[56]
	CK	(+)AaGST formation (+)AaTNG formation (+)artemisinin biosynthesis			[56]
	ABA	(+)AaGST formation (+)artemisinin biosynthesis			[59]
MicroRNAs	miR160	(-)AaGST formation (-)artemisinin biosynthesis	<i>AaARF1</i>		[60]
	miR828	(-)AaGST formation (-)artemisinin biosynthesis		AaMYB17	[61]

+, positive; −, negative.

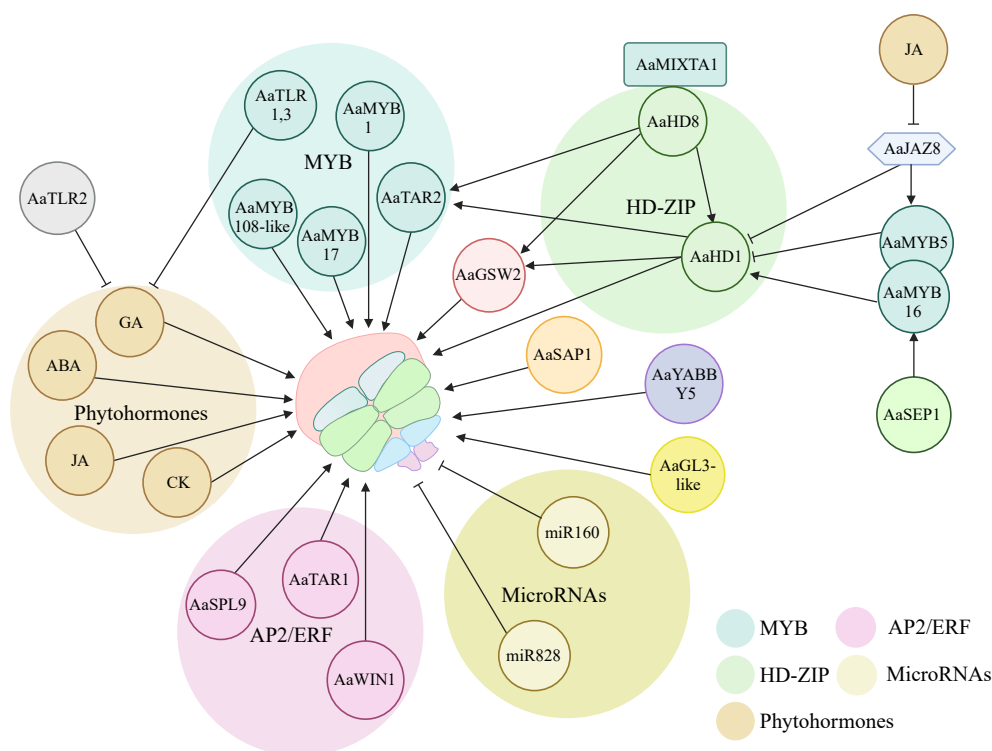
2 (DBR2), aldehyde dehydrogenase 1 (ALDH1), and cytochrome P450 monooxygenase (CYP71AV1), and they are identified in the *AaGSTs*<sup>[46]</sup>.

### R2R3-MYB

AaMYB1, AaMYB5, AaMYB16, AaMYB17, AaMIXTA1, AaTLR1, AaTLR3, and AaTAR2, belong to R2R3-MYB and are involved in

regulating the growth and development of *A. annua* glandular trichomes.

AaMYB1 is the first R2R3-MYB TF found to actively regulate glandular trichome initiation in *A. annua*, and its expression is upregulated at leaf maturity and continues until flower bud formation. Studies have shown that AaMYB1 significantly



**Fig. 2** Regulatory network model for trichome development in *A. annua*. Arrows indicate a positive influence while blocked lines indicate negative regulation.

activates the transcription of several key genes in the artemisinin biosynthesis pathway, especially *ADS* and *CYP71AV1*, resulting in increased artemisinin content in transgenic plants, which may be related to the increase in the number and density of *AaGST* promoted by *AaMYB1*. Additionally, *AaMYB1* can stimulate both *GA4* synthesis and degradation, jointly promoting the formation of *AaGSTs*<sup>[40]</sup>. *AaMYB16* and *AaMYB5* are specific TFs of *A. annua* trichomes that participate in the regulation of *A. annua* trichome formation and their roles are reversed: *AaMYB5* specifically inhibits the formation of *AaGSTs* without affecting the formation of *AaTNGs*, while *AaMYB16* positively regulates *AaGSTs* and *AaTNGs*, and influences artemisinin content. Notably, *AaMYB5* and *AaMYB16* are unable to individually control trichome development. However, through their interactions and regulation of the binding activity of the *AaHD1* promoter, they perform competing roles<sup>[41]</sup>. This study also uncovered that the expression of *AaMYB5* was significantly decreased in jasmonic acid (JA)-treated *A. annua*, whereas the expression of *AaMYB16* remained unaffected. The JA repressor, *A. annua* jasmonate ZIM-domain 8 (*AaJAZ8*), directly enhanced the repressor activity of *AaMYB5* on *AaHD1* transcriptional activation, while it had no impact on *AaMYB16*. *AaMYB17*, specifically expressed in *AaGSTs*, exerts a positive regulatory role in the initiation of *AaGSTs* and artemisinin synthesis while maintaining the morphology of glandular trichomes unchanged<sup>[42]</sup>. On the other hand, *AaMIXTA1*, which shares homology with *AaMYB17* is predominantly expressed in the basal cells of *AaGSTs*. Its mechanism of positive regulation differs from that of *AaMYB17*. Through overexpression of *AaMIXTA1*, RNA interference, and double luciferase reporter gene detection, it has been demonstrated that *AaMIXTA1* positively regulates the expression of cuticle-forming genes

*AaCYP77A1* and *AaCYP86A1*, as well as wax forming genes *AaKCS5*, *AaCER1*, and *AaABCG12*. This indicates that *AaMIXTA1* promotes an increase in the density of *AaGST* and *AaTNG*, while positively regulating the formation of cuticle and wax<sup>[43]</sup>. The R2R3 MYB TFs TRICHOMELESS REGULATOR1 (TLR1) and TLR3 negatively regulate the development of trichomes in *A. annua*. In the *A. annua* plant, the overexpression of *AaTLR1* resulted in a notable reduction of 44.7% to 64.0% in the density of *AaGST* compared to the wild type. Concurrently, there was a down-regulation in the mRNA levels of *ADS*, *CYP71AV1*, *DBR2*, and *ALDH1*, as well as a decrease in the artemisinin content by 11.5% to 49.4%. Conversely, *AaTLR1*-RNAi lines exhibited an increase in *AaGST* density of 33% to 93.3% above the wild type, with a concurrent up-regulation in the mRNA levels of *ADS*, *CYP71AV1*, *DBR2*, and *ALDH1*, and an increase in artemisinin content by 32.2% to 84.0%. These findings suggest that *AaTLR1* is a negative regulator of glandular trichomes and artemisinin content in *A. annua*. Furthermore, the study revealed that *AaTLR1* interacts with the LEAFY-like TF *AaTLR2*, facilitated by *AaWOX1* (a WUSCHEL-like protein), and exerts a negative control over trichome density by suppressing gibberellin levels<sup>[44]</sup>. Research has shown that *AaTLR3* plays a crucial role in glandular trichome development and stratum corneum biosynthesis. In plants with overexpressed *AaTLR3*, the cuticle structure of *A. annua* was notably disrupted, which consequently altered the morphology of its glandular trichomes. This disruption led to a reduction in the density of trichomes by 38%–42% in *AaTLR3* overexpression lines when compared to the control lines. Conversely, in *AaTLR3*-RNAi lines, the density of trichomes saw an increase of 45%–69% relative to the control, and in gene-edited lines, this density increased by 75%–100%. Moreover, *AaTLR3* also exerts regulatory control over



artemisinin biosynthesis. Specifically, in *A. annua* with overexpressed *AaTLR3*, the expression of genes such as *ADS*, *CYP71AV1*, *DBR2*, and *ALDH1*, were diminished, leading to a decrease in artemisinin accumulation. Conversely, in *AaTLR3*-RNAi and gene-edited lines, the expression of these genes was enhanced, and the artemisinin content was significantly elevated. These findings suggest that *AaTLR3* functions as a negative regulator in the development of trichomes and the biosynthesis of artemisinin in *A. annua*. Furthermore, *AaTLR3* can regulate stratum corneum development and trichome morphology by interacting with TRICHOME AND ARTEMISININ REGULATOR 1 (*AaTAR1*) and *CycTL* (a cyclin gene, cyclin trichome less)<sup>[45]</sup>. *AaTAR2* is a multifunctional TF that can affect trichome formation, artemisinin, and flavonoid synthesis in *A. annua*. *AaTAR2* is mainly expressed in the young leaves of *A. annua*. Compared with the wild type, transgenic plants overexpressing *AaTAR2* exhibited a substantial rise in the abundance of *AaGST* and enhanced expression of genes, including *ADS*, *CYP71AV1*, *DBR2*, and *ALDH1*. This upregulation corresponded with an increase in artemisinin content. In contrast, plants with inhibited *AaTAR2* expression, have shown the opposite result. The morphology of *AaGSTs* and *AaTNGs* was drastically altered, with the trichome cells being crumpled and unsupported. This suggests that *AaTAR2* positively regulates trichome initiation and development<sup>[46]</sup>. *AaMYB108* is co-regulated by light and JA and also interacts with *AaGSW2* to enhance *CYP71AV1* transcription, thus promoting artemisinin biosynthesis<sup>[63]</sup>. *AaMYB108-like*, a homologous gene of *AaMYB108* is induced by light and JA and can form a complex with *AaHD8* to promote the expression of downstream *AaHD1*, thereby triggering the initiation of *AaGSTs* and promoting artemisinin biosynthesis<sup>[47]</sup>.

### HD-ZIP

Two HD-ZIP IV TFs, *AaHD1*, and *AaHD8*, regulate the initiation of trichomes in *A. annua*. They can act alone or jointly regulate the activity of other TFs to promote the initiation of *A. annua* trichomes<sup>[34,48]</sup>.

*AaHD1* is regulated by JA. When stimulated by exogenous JA, *AaHD1* is dissociated from the inhibitory protein *AaJAZ8* (a deterrent factor of the JA signaling pathway). The released *AaHD1* then plays a positive role in regulating trichome initiation and artemisinin synthesis in *A. annua*. In the *AaHD1* knockout line, the density of *AaGST* and *AaTNG* decreases significantly, leading to a significant decrease in artemisinin content<sup>[34]</sup>. *AaHD8* indirectly promotes trichome initiation by controlling the expression of genes related to cuticle and wax monomer enzymes, such as *AaCYP86A4* and *AaFDH*. Additionally, *AaHD8* significantly enhances the activity of the *AaHD1* promoter and positively regulates the expression of *AaHD1* in the trichomes of *A. annua*, thus promoting the formation of *AaGSTs* and *AaTNGs*<sup>[34,48]</sup>. Moreover, *AaHD8* interacts with the positive regulatory factor MIXTA-like protein *AaMIXTA1* for trichome initiation and cuticle development, forming a regulatory complex and leading to enhanced transcriptional activity in regulating the expression of *AaHD1* and cuticle development genes<sup>[48]</sup>. *AaHD1* and *AaHD8* can bind to the promoter of *AaTAR2*, enhancing its expression and promoting the formation of *A. annua* trichomes<sup>[46]</sup>. Other studies have shown that *AaHD1* and *AaHD8* can directly regulate *AaGSW2* and promote the formation of *AaGSTs*<sup>[51]</sup>. Since HD-ZIP TFs usually play a regulatory role in the form of dimers, an in-depth study of the interaction mechanism of *AaHD1*/*AaHD8* will help to reveal the

specific regulatory factors regulating the formation of trichomes.

### AP2/ERF

*AaTAR1*, an AP2/ERF TF, regulates the development of glandular trichomes and the biosynthesis of artemisinin in *A. annua*. In *AaTAR1*-RNAi *A. annua* strains, there was abnormal development of *AaGSTs* and *AaTNGs*, reduced accumulation of artemisinin, abnormal deposition of epidermal waxes, and altered cuticle permeability. Conversely, in *AaTAR1* overexpressing strains, there was an increase in the expression of the two key enzyme genes of the artemisinin biosynthesis pathway, *CYP71AV1* and *ADS*, and a significant increase in the content of artemisinin<sup>[38]</sup>. Additionally, *AaTAR1* can interact with the cell cycle protein *AaCycTL* to regulate the biosynthesis of the stratum corneum, negatively regulate the formation of *AaGSTs*, and reduce artemisinin content<sup>[64]</sup>. *AaWIN1* is primarily expressed in the buds, flowers, and trichomes of *A. annua*. It has been demonstrated that the overexpression of *AaWIN1* leads to a significant increase in *AaGST* density and artemisinin content. Importantly, *AaGSW2* is induced to be activated in *AaWIN1* overexpression lines, effectively regulating the activation of *AaGSTs*. Furthermore, *AaWIN1* interacts with *AaMIXTA1* and plays a role in the biosynthesis of the cuticle layer<sup>[49]</sup>. *AaSPL9* is an ERF TF, and yeast one-hybrid, dual-luciferase, and electrophoretic mobility shift assay (EMSA) have shown that *AaSPL9* directly binds to the promoter of *AaHD1* to activate its expression, leading to an increase in the expression of *AaGSW2*, which in turn regulates the initiation of *AaGSTs*. Overexpression of *AaSPL9* increased the density of *AaGST* and artemisinin content, suggesting that *AaSPL9* positively regulates the initiation of *AaGSTs*<sup>[50]</sup>.

### WRKY

WRKY is one of the plant-specific TF families. *AaGSW2*, a secretory glandular trichome-specific TF, which is positively regulated by the direct binding of the homeodomain proteins *AaHD1* and *AaHD8* to the L1-box of the *AaGSW2* promoter. Overexpression of *AaGSW2* significantly increased *AaGST* density, while *AaGSW2* knockdown lines showed impaired *AaGST* initiation<sup>[51]</sup>.

### Other transcription factors

In addition to R2R3-MYB, HD-ZIP, AP2/ERF, and WRKY TFs, there are also other TFs involved in the development of *A. annua* trichomes. For example, *AaSAP1* is an SAP TF, and the upregulation or downregulation of *AaSAP1* transcription levels leads to an increase or decrease in *AaGST* density and artemisinin content, respectively, indicating that *AaSAP1* positively regulates the development of *AaGSTs*<sup>[52]</sup>. Recent studies have revealed that *AaTLR2* affects trichome development by regulating gibberellin levels<sup>[44]</sup>. In *A. annua* *AaTLR2* overexpression lines, the trichome density was reduced, and the expression levels of *ADS*, *CYP71AV1*, *DBR2*, and *ALDH1* also showed a tendency to be reduced, which directly led to the decrease of artemisinin content in *A. annua*. This shows that *AaTLR2* plays a negative regulatory role in trichome initiation and artemisinin biosynthesis in *A. annua*. The study revealed a MADS-box gene *AaSEPALLATA1* (*AaSEP1*), which can be induced by JA and light signals and promotes *AaGST* initiation in *A. annua*<sup>[53]</sup>. Overexpression of *AaSEP1* in *A. annua* significantly increased *AaGST* density and artemisinin content. This study found that *AaSEP1* enhances the activation of the downstream *AaGST* promoter

gene *AaGSW2* by interacting with *AaMYB16*. In addition, the interaction between *AaSEP1* and *AaJAZ8* is an important factor in JA-mediated *AaGST* initiation. The study also found that *AaSEP1* interacts with light morphogenetic protein 1 (*AaCOP1*), the main inhibitor of light signaling. The YABBY gene family, comprising a modest array of TFs within seed plants is pivotal in orchestrating the growth and development of plant leaves<sup>[65]</sup>. *AaYABBY5*, a distinguished family member, was cloned and characterized in a recent study. The research revealed that *AaYABBY5* directly binds to the promoters of *CYP71AV1* and *DBR2* or indirectly regulate *ADS* and *ALDH1*, thereby activating their gene expression and substantially enhancing the concentration of artemisinin<sup>[54]</sup>. Furthermore, the study demonstrated that the trichome density in plants overexpressing *AaYABBY5* was significantly elevated compared to control plants, while antisense plants exhibited a marked reduction in trichome density<sup>[66]</sup>. These findings underscore *AaYABBY5*'s role as a positive regulator in the biosynthesis of artemisinin and the development of trichomes. Some scholars have successfully isolated a homolog of *AtGL3* (*GLABRA3*) from *A. annua*, named *AaGL3-like*, which is categorized in the bHLH TF family. In-depth studies showed that *AaGL3-like* was able to be induced by JA and significantly increase the final accumulation of artemisinin content by regulating the density of trichomes in *A. annua*<sup>[55]</sup>. A study has shown that the increase in the expression level of *Transparent testa glabra 1* (*TTG1*) is related to the increase in glandular trichome density and artemisinin production in *A. annua* leaves<sup>[67]</sup>.

These TFs are involved in regulating genes associated with cell division, growth, and differentiation. They exert their effects by activating or inhibiting specific gene expression to control trichome formation and development in *A. annua*. A comprehensive understanding of the functions and regulatory mechanisms of these TFs will contribute to elucidating the molecular mechanisms underlying trichome growth and development in *A. annua*. It may offer novel strategies for cultivating artemisinin high-yield varieties.

### Phytohormones

Phytohormones are signaling molecules that regulate specific cellular processes and play a crucial role in plant growth, development, and environmental stress responses<sup>[68]</sup>. Currently, research has found that phytohormones such as jasmonic acid (JA), gibberellins (GAs), cytokinin (CK), and abscisic acid (ABA) regulate the growth and development of *A. annua* trichomes by inducing TFs and some of them affect the biosynthesis of artemisinin.

#### Jasmonic acid

JA belongs to the lipid derivatives and is a plant-secondary metabolism stimulator<sup>[69]</sup>. Since artemisinin content is closely related to the density of *AaGST*, JA can regulate the growth and development of *AaGSTs* by affecting genes related to the development of *AaGSTs*. For example, JA induced the specific expression of *trichome-specific fatty acyl-CoA reductase 1* (*TFAR1*) in *AaGSTs*, which resulted in a significant increase in the density and size of *AaGSTs* on *A. annua* leaves and also promoted the biosynthesis of artemisinin<sup>[56]</sup>. When *AaAOC*, a key enzyme in the JA synthesis pathway is overexpressed in *A. annua*, the endogenous JA content in *A. annua* increases, which leads to an increase in the density of *AaGST*<sup>[57]</sup>. JA also induces the formation of *AaGSTs* with the help of the HD-ZIP family of TFs,

*AaHD1* and *AaHD8*. JA regulates the function of the *AaHD1* gene at both the transcriptional level and the protein level, thereby positively regulating the early development of secretory glandular trichomes on the leaf surface of *A. annua*; the expression of *AaHD1* was significantly increased in JA-treated *A. annua* plants, which promotes the formation of *AaGSTs* and *AaTNGs*<sup>[34]</sup>. In addition, JA treatment of *A. annua* plants significantly downregulated the expression of *AaMYB5*, which acts with *AaJAZ8* to regulate the formation of *AaGSTs*<sup>[41]</sup>.

#### Gibberellins

GAs are diterpenoids, which are present in all higher plants as a class of phytohormones and play a role in many developmental processes. It has been shown that the density of *AaTNG* on *A. annua* leaves increased significantly after GA3 treatment, whereas there was no significant change in the density of *AaGST* and artemisinin content<sup>[56]</sup>, and it has also been recently shown that GA3 does not stimulate the formation of *AaGSTs* or the biosynthesis of artemisinin, whereas endogenous GA1/GA4 can promote *AaGSTs* formation and artemisinin synthesis by *AaMIXTA1*, which promotes *AaGSTs* formation and artemisinin synthesis<sup>[58]</sup>, which may be because exogenously applied GA3 is not the active form for *AaGSTs* formation.

#### Cytokinin

CK can promote the development of trichomes in *A. annua*, and 6-benzylaminopurine (BAP) was found to increase the density of *AaTNGs* and stimulate the formation of *AaGSTs*. The density of glandular trichomes in BAP-treated plants was significantly increased and the glandular trichomes were enlarged, but BAP did not stimulate artemisinin biosynthesis<sup>[56]</sup>.

#### Abscisic acid

ABA is a natural plant growth regulator that can promote coordinated plant growth and improve plant growth quality. Research has found that externally applied ABA can reverse the negative effects of copper treatment, increase the density and size of *AaGSTs* under copper stress, and increase the content of artemisinin under copper stress<sup>[59]</sup>.

#### MicroRNAs

Like TFs, microRNAs (miRNAs) are also indispensable regulatory factors in trichomes. miRNAs are non-coding RNAs found in eukaryotes with endogenous regulatory functions. They are approximately 20–25 nucleotides in size and play a crucial role in plant development and secondary metabolism through different sequence-specific interaction patterns with targets. Studies have shown that *miR160* reduces the formation of *AaGSTs* by targeting and cleaving *AaARF1*. The results showed that in the *miR160*-overexpressing strains, the density of *AaGST* and artemisinin content decreased significantly. On the contrary, in the *miR160* knockout strains, the density of *AaGST* increased compared with the wild type, and the content of artemisinin was also significantly increased<sup>[60]</sup>. Recently, *miR828* was found to degrade *AaMYB17* in *A. annua* plants. Overexpression of *miR828* reduced the density of *AaGST* by about 60%, and artemisinin content was consequently reduced<sup>[61]</sup>. The study of miRNA provides new possibilities for increasing artemisinin content, but further in-depth research is needed on this regulation.

#### Others

Under natural conditions, plants are always exposed to environmental stresses. In addition to TFs and phytohormones, some adversity factors, such as drought, salt, heavy metals, and

## Glandular trichomes and artemisinin

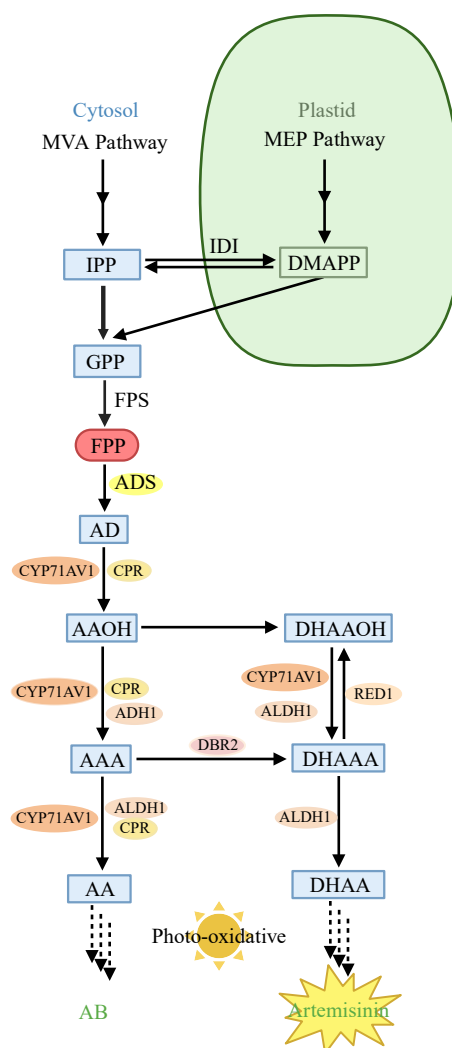
other abiotic stresses, also have some effects on the formation and development of trichomes in *A. annua*.

It has been discovered that mild-to-moderate drought stress results in a reduction in both the density and volume of AaGSTs, in comparison to non-stress conditions<sup>[70]</sup>. This decrease in artemisinin content was found to be closely associated with the reduction. Under 50 and 100 mM salt treatments, the length and width of AaGSTs increased, along with an increase in the number of AaGSTs. However, under 200 mM salt stress, the volume of AaGSTs decreased, while the density of AaGST remained constant throughout all stages of plant life<sup>[37]</sup>. Moreover, increasing copper concentration led to a decrease in the density, average width, and length of AaGSTs, as well as a decline in artemisinin content in *A. annua* plants exposed to copper stress when compared to the control group<sup>[59]</sup>. Arsenic treatment (0–45 mg/kg) of *A. annua* resulted in an increase in artemisinin content, which may be related to the expansion of artemisinin-producing apical cells on the AaGSTs after Arsenic treatment<sup>[71]</sup>. High cadmium stress (20, 40 mg/kg) had a significant negative effect on the length, width, and density of the AaGST, as well as the artemisinin content, which was all significantly negatively affected, and the higher the cadmium concentration, the greater the effect<sup>[72]</sup>. Exogenous supplementation of strigolactone (GR24) under cadmium stress or spraying NO on the stressed plants could significantly improve the situation<sup>[72,73]</sup>. In a recent study, the carbon nanomaterial graphene was found to block the biogenesis of miRNAs and directly disrupt the function of the miR828-AaMYB17 module, which enhanced the initiation of AaGSTs and induced the production of artemisinin<sup>[61]</sup>.

Although many studies have reported the effects of non-biological stress signals on the trichomes and artemisinin content of *A. annua*, there is relatively little research on the effects of biological stress signals on the trichomes and artemisinin content of *A. annua*. Scholars have found that *A. annua* plants inoculated with *Glomus macrocarpum* and *Glomus fasciculatum* have a higher density of AaGSTs on their leaves compared to non-fungal root plants, and the detected artemisinin content is also higher. There is a strong positive linear correlation between the density of AaGST on the leaves and artemisinin concentration<sup>[74]</sup>.

## Artemisinin biosynthetic pathway in AaGSTs

Plant secondary metabolites are a class of biomolecules synthesized by secondary metabolic pathways, which are important protective barriers for plants to adapt to the environment and cope with adverse environments. Artemisinin is a sesquiterpenoid synthesized and stored in AaGSTs, and its biosynthetic pathway belongs to the isopentenyl-like metabolic synthesis pathway originating from isopentenyl diphosphate (IPP) and dimethylallyl diphosphate (DMAPP)<sup>[75,76]</sup> (Fig. 3). The mevalonate acid (MVA) pathway significantly influences the synthesis of farnesyl pyrophosphate (FPP), with its inhibition leading to an 80.4% reduction in artemisinin content. Conversely, disruption of the methylerythritol phosphate (MEP) pathway results in a mere 14.2% decrease in artemisinin content<sup>[77]</sup>. In the course of FPP synthesis, a molecule of IPP derived from the MEP pathway is initially coupled with a molecule of DMAPP produced by the MVA pathway to form geranyl diphosphate (GPP). Subsequently, this GPP is



**Fig. 3** Biosynthetic pathways of artemisinin biosynthesis. AA, Artemisinic acid; AAA, Artemisinic aldehyde; AAOH, Artemisinic alcohol; AB, Arteannuin B; AD, amorpha-4,11-diene; ADH1, alcohol dehydrogenase 1; ADS, amorpha-4,11-diene synthase; ALDH1, aldehyde dehydrogenase 1; CPR, cytochrome P450 reductase; CYP71AV1, cytochrome P450 monooxygenase; DBR2, artemisinic aldehyde double bond reductase 2; DHAA, dihydroartemisinic acid; DHAAA, dihydroartemisinic aldehyde; DHAAOH, dihydroartemisinic alcohol; DMAPP, dimethylallyl diphosphate; FPP, farnesyl diphosphate; FPS, farnesyl pyrophosphate synthase; GPP, geranyl diphosphate; IDI, isopentenyl-diphosphate isomerase; IPP, isopentenyl diphosphate; MEP, 2-C-methyl-D-erythritol 4-phosphate; MVA, mevalonic acid; RED1, dihydroartemisinic aldehyde reductase 1.

transported to the parietal cells and the cytoplasm, where it is converted into the universal precursor FPP by the enzyme farnesyl diphosphate synthase (FPS), utilizing an additional molecule of IPP originating from the MVA pathway<sup>[78,79]</sup>. Sesquiterpenoids are synthesized using FPP as a substrate and catalyzed by sesquiterpene enzymes. ADS is a specific sesquiterpene enzyme for artemisinin synthesis, and FPP is catalyzed by ADS to produce amorpha-4,11-diene (AD)<sup>[80–83]</sup>, and AD is converted to artemisinic alcohol (AAOH) via CYP71AV1<sup>[84,85]</sup>. Subsequently, AAOH is oxidized to artemisinic aldehyde (AAA) by alcohol dehydrogenase 1 (ADH1) and



CYP71AV1<sup>[84,85]</sup>. It is worth noting that artemisinin aldehyde can undergo catalysis by various enzymes, leading to the production of different metabolites. Specifically, the catalysis of AAA by CYP71AV1 and ALDH1<sup>[86]</sup> results in the formation of artemisinic acid (AA), which eventually leads to the synthesis of arteannuin B (AB). Alternatively, AAA can be catalyzed by DBR2 and then ALDH1, resulting in the production of dihydroartemisinic aldehyde (DHAAA)<sup>[87]</sup>, dihydroartemisinic acid (DHAA)<sup>[86]</sup> and ultimately artemisinin. AA and DHAA undergo non-enzymatic photo-oxidation to ultimately produce artemisinin B and artemisinin, respectively<sup>[88,89]</sup>. In addition, AAOH can generate other intermediates involved in artemisinin synthesis. For instance, AAOH can undergo catalysis to yield dihydroartemisinic alcohol (DHAAOH)<sup>[90]</sup>. Subsequently, DHAAOH can be catalyzed by CYP71AV1 and ALDH1 to form dihydroartemisinic aldehyde (DHAAA). Additionally, DHAAA can be specifically reduced back to DHAAOH by the enzyme dihydroartemisinic aldehyde reductase 1 (RED1), allowing it to continue its involvement in the synthesis of artemisinin<sup>[91]</sup>.

## Conclusions and perspectives

Artemisinin is primarily synthesized, secreted, accumulated, and stored in the secretory trichomes of *A. annua*<sup>[92]</sup>. Hence, the formation and development of secretory trichomes are crucial for increasing artemisinin yield. The biosynthesis of artemisinin is divided into upstream and downstream pathways, with the upstream pathway being relatively clear, while the downstream steps are not yet fully elucidated. Therefore, improving artemisinin yield by focusing on key enzymes in the metabolic pathway is limited. In contrast, as the 'chemical factory' of artemisinin biosynthesis, trichomes contain all the necessary factors for artemisinin biosynthesis and have unparalleled advantages in increasing artemisinin content in *A. annua*. Regulating trichomes can bypass the uncertainties and controversies in the artemisinin biosynthesis pathway, allowing for more efficient macro-control of artemisinin biosynthesis<sup>[6]</sup>.

In recent years, scientists have explored factors affecting the formation and development of *A. annua* trichomes, including transcription factors, plant hormones, and microRNAs. These studies reveal the key role of trichomes in artemisinin biosynthesis and provide an important theoretical basis for subsequent research. With the continuous development of theories and technologies such as molecular biology, cell biology, multi-omics, molecular breeding, gene editing, and single-cell transcriptome sequencing (scRNA-Seq), more and more research methods are being applied to the study of *A. annua* trichome development and artemisinin biosynthesis. For example, gene knockout, silencing, and gene editing can construct mutants with target gene deletions, and then observe the differences between mutants and wild types to verify the impact of target gene function on artemisinin synthesis or trichome development. In addition, transcriptomics and metabolomics analysis facilitate the study of trichomes and artemisinin biosynthesis. After screening and identifying target genes for molecular breeding, new artemisinin high-yield varieties of *A. annua* can be cultivated using a combination of traditional breeding and molecular breeding. These methods all provide important ideas for improving artemisinin yield. ScRNA-Seq is a high-throughput genome sequencing technology that has tremendous advantages in identifying intercellular heterogeneity, analyzing gene expression at high cellular resolution, and

constructing regulatory networks of TFs<sup>[93]</sup>. In the last few years, scRNA-Seq technology has developed rapidly in the field of plant research. For example, a study has revealed the early developmental dynamics of leaf vein cells in *Arabidopsis* cotyledons using scRNA-Seq technology, which provides new cell-type specific gene expression profiles and potential marker genes for future studies on the development of plant vascular tissues<sup>[94]</sup>. In addition, scRNA-Seq technology has been applied to the medicinal plant *Nepeta tenuifolia* L., where multiple differentially expressed genes that may regulate the growth and development of glandular trichomes in *Nepeta tenuifolia* were identified, providing a basis for exploring the development and differentiation of plant cells, especially the initiation and development of glandular trichomes<sup>[95]</sup>. This method can be similarly applied to the study of *A. annua*, which will greatly facilitate the discovery of important functional genes, deepen the understanding of the complex molecular mechanisms of glandular trichome formation and development, artemisinin biosynthesis and their regulatory networks, and provide more comprehensive scientific guidance for the improvement of *A. annua* varieties and the enhancement of artemisinin production.

Regarding the formation and development of trichomes in *A. annua*, although transcription factor families such as AP2/ERF, WRKY, MYB, and HD-ZIP have been shown to play important roles in regulating trichome development, the vast number of plant transcription factors and the complex regulatory network means that the underlying molecular mechanisms are mostly unknown. Therefore, further exploration of the molecular mechanisms of trichome development is necessary, especially the synergistic regulatory effects between transcription factors and miRNAs. This will help us better understand the trichome development process and provide a theoretical basis for improving trichome characteristics through genetic engineering. Previous studies have confirmed that changing trichome characteristics, such as regulating trichome morphology with TAR1<sup>[38]</sup> and regulating trichome density with TAR2<sup>[46]</sup>, can significantly affect artemisinin yield. In addition, based on a thorough understanding of the artemisinin biosynthesis pathway, reorienting metabolic flow through regulation to increase artemisinin yield will also be a key research direction we can focus on.

In summary, as the factory of artemisinin biosynthesis, AaGST plays a significant role in improving artemisinin yield. The AaGST is a typical secretory glandular trichome, which is not present in the model plants *Arabidopsis* and rice. Therefore, AaGST can be studied as a model organ for secretory glandular trichomes and can be used as a reference for other species.

## Author contributions

The authors confirm contribution to the paper as follows: conceptualization and supervision: Tan H; draft manuscript and figure preparation: Zhao Q; manuscript review and editing: Li M, Zhang M. All authors reviewed the results and approved the final version of the manuscript.

## Data availability

Data sharing not applicable to this article as no datasets were generated or analyzed during the current study.



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## Conflict of interest

The authors declare that they have no conflict of interest. Hexin Tan is the Editorial Board member of *Medicinal Plant Biology* who was blinded from reviewing or making decisions on the manuscript. The article was subject to the journal's standard procedures, with peer-review handled independently of this author and the research groups.

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