Tariq Aftab Khalid Rehman Hakeem *Editors*

Metabolic Engineering in Plants



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Preface

Metabolic engineering is usually defined as the re-routing of one or more enzymatic reactions to generate new compounds, increase the production of existing compounds, or facilitate the degradation of compounds. Metabolic engineering is chiefly focused on metabolic intermediates or products, such as starch, vitamin E, amino acids, and enzymes. Metabolic engineering is different from genetic engineering because it investigates the properties of integrated metabolic pathways and genetic regulatory networks, not with individual genes and enzymes. Plants are the foundation of numerous compounds which are synthesized via assimilated complex biosynthetic routes. Plant metabolites are organic compounds synthesized via enzymemediated processes. These plant metabolites are important for both essential (proteins, lipids, and carbohydrates) as well as specific and specialized functions (anthocyanin, carotenoid, etc.). The primary metabolites are essential for the survival of the organisms and any alterations in their levels are likely to have severe major manifestations. However, secondary metabolites are considered to be the metabolites having specialized functions and playing a role in providing quality of life to the producer organism. Plants have evolved an incredible arrangement of metabolic pathways leading to molecules/compounds capable of responding promptly and effectively to stress situations imposed by biotic and abiotic factors, some of which supply the ever-growing needs of humankind for natural chemicals, such as pharmaceuticals, nutraceuticals, agrochemicals, food and chemical additives, biofuels, and biomass. Humans rely on plants for various needs since ancient time, and their population still seems enough for fulfilling our demands. But, in foreseeable future we will be forced to think about the accessibility of resources for the generations to come. For these reasons, we must look for alternative options of resources about food/food supplement, medicines, and other essential items. For achieving this task, we are choosing plant metabolic engineering through which we can generate higher amount of compounds/products in less time and space. The aim of metabolic engineering is enhancement of desired product through upregulation / downregulation of protein expression. With an estimated 100,000 unique compounds produced in the plant kingdom, elucidating these metabolic networks is an exciting endeavour. By using plant metabolic engineering approach, tremendous changes at metabolic level in different plants have been done.

The book comprises 19 chapters, review articles written by experts, highlighting a wide range of topics such as whole-plant and cell/organ culture systems, and environmental and genetic transformation-based modulation of biochemical pathways. Special focus is given to microRNA, RNAi, and CRISPR-Cas-mediated approaches for metabolic engineering applications using both model and non-model plant species. We are hopeful that this volume would furnish the need of all researchers who are working or have interest in this particular field.

We are highly grateful to all our contributors, for accepting our invitation, and for not only sharing their knowledge and research, but also for venerably integrating their expertise in dispersed information from diverse fields in composing the chapters and enduring editorial suggestions to finally produce this venture. We also thank Springer Nature team for their generous cooperation at every stage of the book production.

Lastly, thanks are also due to well-wishers, research students, and editors' family members for their moral support, blessings, and inspiration in the compilation of this book.

Aligarh, India Jeddah, Saudi Arabia Tariq Aftab Khalid Rehman Hakeem

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About the Editors



Tariq Aftab received his Ph.D. in the Department of Botany at Aligarh Muslim University, India, and is currently an Assistant Professor there. He is the recipient of a prestigious Leibniz-DAAD fellowship from Germany, Raman Fellowship from the Government of India, and Young Scientist Awards from the State Government of Uttar Pradesh (India) and Government of India. After completing his doctorate, he has worked as Research Fellow at National Bureau of Plant Genetic Resources, New Delhi, and as Post-doctorate Fellow at Jamia Hamdard, New Delhi, India. Dr. Aftab also worked as Visiting Scientist at the Leibniz Institute of Plant Genetics and Crop Plant Research (IPK), Gatersleben, Germany, and in the Department of Plant Biology, Michigan State University, USA. He is a member of various scientific associations from India and abroad.

He has edited 15 books with international publishers, including Elsevier Inc., Springer Nature and CRC Press (Taylor & Francis Group), co-authored several book chapters, and published over 75 research/review papers in peer-reviewed international journals. His research interests include physiological, proteomic, and molecular studies on medicinal and crop plants.



Khalid Rehman Hakeem Ph.D. is Professor at King Abdulaziz University, Jeddah, Saudi Arabia. After completing his doctorate (Botany; specialization in Plant Eco-physiology and Molecular Biology) from Jamia Hamdard, New Delhi, India, in 2011, he worked as a lecturer at the University of Kashmir, Srinagar, for a short period. Later, he joined Universiti Putra Malaysia, Selangor, Malaysia, and worked there as Post-doctorate Fellow in 2012 and Fellow Researcher (Associate Prof.) from 2013 to 2016. Dr. Hakeem has more than 10 years of teaching and research experience in plant eco-physiology, biotechnology and molecular biology, medicinal plant research, plant-microbe-soil interactions as well as in environmental studies. He is the recipient of several fellowships at both national and international levels; also, he has served as the visiting scientist at Jinan University, Guangzhou, China, Currently, he is involved with a number of international research projects with different government organizations.

So far, Dr. Hakeem has authored and edited more than 70 books with international publishers, including Springer Nature, Academic Press (Elsevier), and CRC Press. He also has to his credit more than 140 research publications in peer-reviewed international journals and 60 book chapters in edited volumes with international publishers.

At present, Dr. Hakeem serves as an editorial board member and reviewer of several high-impact international scientific journals from Elsevier, Springer Nature, Taylor & Francis, Cambridge, and John Wiley Publishers. He is included in the advisory board of Cambridge Scholars Publishing, UK. He is also a fellow of Plantae group of the American Society of Plant Biologists, member of the World Academy of Sciences, member of the International Society for Development and Sustainability, Japan, and member of Asian Federation of Biotechnology, Korea. Dr. Hakeem has been listed in Marquis Who's Who in the World since 2014–2019. Currently, Dr. Hakeem is engaged in studying the plant processes at eco-physiological as well as molecular levels.

Chapter 1 Metabolic Engineering: New Approaches in Pharmaceutical Production



Ahmed H. El-Desoky, Mohamed A. M. Atia, and Elsayed A. Omer

Abstract Natural products, natural products-derived, and natural products-inspired compounds represent a major sector of drugs and drug leads. Nature has the potential to produce compounds built to interact with biological systems. Mostly are produced as secondary metabolites by a living organism for different purposes such as defense and communication. Natural compounds usually contain multiple chiral centers within strictly specific molecular architecture that are very difficult to be obtained by combinatorial synthesis for economic supply of drugs. Meanwhile, natural compounds serving as drugs or precursors for synthesis are having several drawbacks with insufficient and fluctuating supply as the most serious problems. For these reasons metabolic engineering exploiting advanced biotechnology techniques such as sequencing, recombinant DNA, and protein engineering can add new incentives to produce secondary metabolites. Metabolic engineering is based on manipulating metabolic networks/pathways within prokaryotic organisms or cultured eukaryotes. Both organisms have common features, including a cytoplasmic membrane, DNA that codes for genetic information, and RNA that is subsequently translated into proteins. Successful study of the biosynthetic pathways responsible for the production of natural products together with advances in fermentation technologies and expression systems, variation in tool kits, and mutation tools allowed metabolic engineering to be a promising approach for the targeted production of a wide variety of natural compounds (taxol, camptothecin, vincristine, artemisinin, silymarin) with bacterial, fungal, and plant origins.

Keywords Metabolic engineering · Plant · Pathway · Secondary metabolites · Taxol · Camptothecin · Vincristine · Artemisinin · Silymarin

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Introduction

Metabolic engineering (ME) is defined as the directed development and improvement for the production of biochemical, biofuels, pharmaceuticals, and medicinal products by modifying and bioengineering a particular metabolic pathway(s). This method includes functional alteration of metabolic pathways toward better understanding and utilizing cellular/biological pathways. In addition, metabolic engineering is driven to advance different commercial applications by which we can enhance the strain's efficiency for producing valuable metabolites. The standard metabolic engineering method requires controlling the expression of a specific protein(s) either in overexpression or downregulation manner within a metabolic pathway, which promotes the biological cell to produce a new product (Yadav et al., 2018).

Directionality is the core of a successful metabolic engineering system, but scaling, stability, and productivity are essential factors. Unfortunately, many trials have failed during the last decade due to problems confronted in scaling up or other critical factors. Notably, meanwhile, the plant production systems are generally slower than microbial systems; plants are more suitable for commercial production due to their autotrophic and scale-up features, in addition to their low cost, especially when agricultural infrastructure is available. However, for plant production systems to be effective, using either fermentation or agriculture, stability and high yields of the bioengineered metabolites must be ensured (Nielsen, 2001).

Since Bailey's formal recognition of metabolic engineering in 1991, it demonstrated its powerful capability as a modern approach in engineering numerous microbial strains to produce hundreds of chemicals and bioproducts from raw substances (Nielsen, 2001; Bailey, 1991; Birchfield & McIntosh, 2020). This production is classically achieved by introducing targeted genetic changes through recombinant DNA technologies and analyzing all possible consequences of these changes at the cellular/biological levels. However, the number of bioproducts touching industrialscale production is limited due to the high production costs, including raw materials, fermentation process, and purification. Thus, although immense efforts have been paid in the metabolic engineering field, it is still needed to enhance strain performance in order to create competitive bioprocesses compared to traditional chemical processes. Also, ME projects at their beginning stages frequently fail due to many critical practical issues occurring during industrial scaling production (Choi et al., 2019). Moreover, inaccurate trial-and-error cycles to enhance strain/plant performance usually resulted from a poor understanding of the metabolic and/or genetic mechanisms controlling a particular pathway in the host organisms, especially in the laboratory (Courdavault et al., 2021).

In a global sense, metabolic engineering is not considerably different from genetic or cellular engineering since they all intended to manipulate genes to generate a product of interest. However, metabolic engineering is focused explicitly on understanding the complex metabolic pathways/networks; meanwhile, genetic and cellular engineering focuses on particular enzymatic reactions inside the cell. Thus, a systemic approach is required for the straightforward and precise design of

3

complex pathways through remodeling the optimal cell pathways, which can divert cell supplies to achieving more beneficial fitness (Choi et al., 2019; Courdavault et al., 2021; Liu & Nielsen, 2019). Accordingly, metabolic engineering represents a robust framework for dissecting differential gene expression at a genome-wide level, protein production, and the levels of a wide range of intracellular metabolic fluxes. Therefore, in order to increase the metabolic engineering approach productivity of certain products, many questions need to be carefully answered: (1) which host/ organism must be chosen, (2) what commodity could be produced, and (3) which ME approach could be applied (Liu & Nielsen, 2019). Therefore, this chapter highlighted the different biological systems that can be used in the ME, examples of current commodities produced, and lastly, the various engineering strategies that can be employed to engineer and improve cellular production capabilities of specific metabolites that are in clinical use and are facing difficulties in commercial production hindering stable market supplies.

Strategies

Several approaches are adopted for secondary metabolites development or enhancement using bacteria, such as (1) overexpressing of a gene family or set of transcription factors to simultaneously stimulate/initiate diverse pathway-related endogenous genes to improve the synthesis of particular metabolites, (2) engineering of regulatory networks controlling a particular pathway through delicate tuning of many enzymes within the target pathway to assure an adequate accumulation of precursors and consequently boost metabolic flux through the targeted pathway, (3) gene deletion (knocking out) or/and insertion of essential enzyme genes regulating or controlling the pathway of the product of interest, (4) stimulation using specific substrates/elicitors, and (5) combining one or more of the above approaches to maximize the synthesis of the target metabolites. Recently, gene editing via CRISPR/Cas9-based activation and/or repression schemes can also be used in metabolic engineering development and improvement. Therefore, metabolic engineering represents potential cell factories to produce pharmaceuticals, amino acids, bioplastics, biofuels, new bioproducts, silk proteins, etc. (Liu & Nielsen, 2019; Zhu et al., 2020).

Many designed tools were implemented in metabolic engineering during the last decade for more precise pathway construction and optimized pathway development. These include selecting hosts with well-known and knowledgeable omics databases, computational pathway simulations resources/tools paired with custom enzyme designs, and guided evolution approaches that attempt to maximize the output of the desired product (Zhu et al., 2020). In addition, advancements in microbial engineering have enabled a wide range of plant metabolites to be manufactured (Fu et al., 2018).

Host Selection

Metabolic engineering is based on manipulating metabolic networks/pathways within prokaryotic organisms or cultured eukaryotes. Both organisms have common features, including a cytoplasmic membrane, DNA that codes for genetic information, and RNA that is subsequently translated into proteins. However, the distinguishing factors between the two kinds of organisms are revealed intracellular when comparing their genomes or organelles. For example, prokaryotes, including the Bacteria kingdom, include mostly stable chromosomal DNA with one inherent single loop and lacks membrane-bound organelles. In contrast, eukaryotes, such as Fungi, Plantae, and Animalia kingdoms, comprise DNA molecules organized in tightly bound chromosomes and contain membrane-bound organelles (e.g., chloroplasts, mitochondria, and the Golgi apparatus). The bacteria, fungi, plant, and animal cells are currently used in different metabolic engineering applications to manufacture products, and accurate analysis is required to decide the most fitting host for a particular target (Nabavi et al., 2020).

The selection of an organism that will be used in the production for an industrial biotechnology process is mainly driven by its potential to manufacture the product of interest efficiently. Genetically engineered cells are usually used to make two main kinds of products, proteins, and nonproteins (Zhu et al., 2020). Protein products comprise those used for human therapeutics, food processing, and industrial catalysts, and are generated by introducing their encoding genes into the host genome. Besides, nonprotein products, including metabolites such as amino acids, biofuels, antioxidants, and vitamins, are created by inserting genes coding for necessary enzymes that manipulate precursors into desired metabolites. Cultures of complex eukaryotes (yeast, plant cells, mammalian cells) are used to produce proteins, whereas prokaryotes and cultures of simple eukaryotes (yeast, filamentous fungi, and plants) are used more frequently in the production of nonprotein metabolites (Nabavi et al., 2020).

For long decades, bacteria, fungi, and plants have been acknowledged as excellent sources of natural compounds, especially those with multiple bioactive attributes and associated applications in the domains of nutrition and human health. Among the common valuable plant natural products (PNPs), scientists have identified those usually used to treat different pathologies such as cancers, infectious diseases, or cardiovascular diseases. Additionally, selected PNPs have been further processed into various active pharmaceutical constituents at an industrial scale. Examples of the most famous PNPs, and their associated applications, include tropane alkaloids from Solanaceae (mostly anticholinergics), wormwood sesquiterpene lactones (antimalarial drugs), monoterpene indole alkaloids of Apocynaceae (antihypertensive and anticancer drugs), poppy isoquinoline alkaloids (antitussives, antimicrobials, analgesics, vasodilators), yew taxane-type terpenoids (anticancer drugs), and mayapple lignans (anticancer and immunosuppressive drugs) (García-Granados et al., 2019).In the following pages, we will throw the light on the engineering of some vital metabolites that are used in the treatment of some vital diseases such as cancer, malaria, and hepatitis. The drug markets need these metabolites, and the demand for them is increasing day by day. These metabolites as arranged in the text are taxol, camptothecin, vincristine, artemisinin, and silymarin.

Examples of Engineered Metabolites

Taxol (TXL)

Paclitaxel, commercially known as taxol, is a terpenoidal pseudo-alkaloid that has potent antitumor activity (Sabzehzari & Naghavi, 2019). Paclitaxel is one of the strong plant-derived anticancer drugs that was first isolated from the park of Pacific yew. It was first isolated from the bark of the slow-growing coniferous tree *Taxus parvifolia* known as Pacific yew or western yew (Mansukhlal et al., 1971) in a very low yield 0.01–0.05 % (Cragg, 1988). Being only isolated from the bark, it is very hard to meet the increasing clinical consumption of taxol. One tree yields 18 kg of green bark, which becomes about 9 kg after drying (Connolly, 1988). 27,700 kg of dried bark yield about 4 kg of taxol (Cragg, 1988). Despite its clinical success, limited supply of paclitaxel for clinical trials is a very serious drawback. Meanwhile, combinatorial synthesis is very hard due to multiple chirality. Meanwhile, consumption of taxol is expected to show a yearly growth of 8.2% between 2020 and 2025 (Ning et al., 2020).

Biosynthetic Background

Biosynthesis of paclitaxel is divided into three stages (Fig. 1.1). The first is the MVA/MEA pathway-dependent synthesis of isopentenyl pyrophosphate (IPP) as terpene precursor which is the rate-limiting step (Soliman & Tang, 2015), the second is the synthesis of baccatin III as taxol's main carbon framework, and the third is the synthesis of nitrogenous side chain originated from the amino acid phenylalanine (Sabzehzari & Naghavi, 2019; Croteau et al., 2006).

Conventional Strategies to Increase TXL Yields

Due to its enormously increasing clinical importance, many attempts have been made to increase the production of paclitaxel including chemical synthesis that mainly depended on the exploitation of commercially abundant biosynthetic precursors as starting materials (Denis et al., 1988; Holton et al., 1994; Borah et al., 2007), searching for microbial sources that can be bioengineered to increase the production such as *Taxomyces andreanae* isolated from *Taxus chinensis* (Stierle

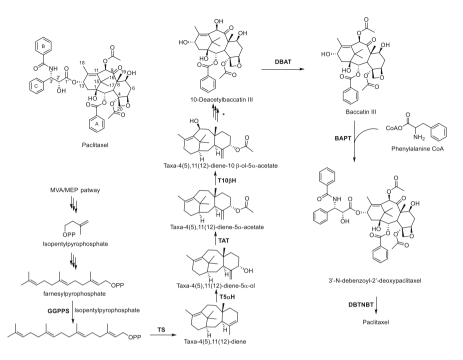


Fig. 1.1 Biosynthetic pathway for taxol [abbreviations: *MVA* mevalonic acid, *MEP* 2-C-methyl-Derythritol-4-phosphate, *GGPPS* geranylgeranyl pyrophosphate synthase, *TS* taxa-4(5),11(12)-diene synthase (committed step), *T5*α*H* taxa-4(5),11(12)-diene-5α-hydroxylase, *TAT* taxa-4(5),11(12)diene-5α-ol-O-acetyltransferase, *T10βH* taxane-10β-hydroxylase; *oxetane ring formation and branch migration enzymes taxane 2α-O-benzoyltransferase (*T2BT* or *DBBT* debenzoyltaxane-2/-α-O-benzoyltransferase) and C-13 hydroxylation; *DBAT* 10-deacetylbaccatin III-Oacetyltransferase, *BAPT* baccatin III 13-O-(3-amino-3-phenylpropanoyl)transferase, *DBTNBT* 3/-N-debenzoyl-2/-deoxytaxol-N-benzoyltransferase, after side chain hydroxylation by unknown enzyme; multiple arrows mean more than one step]

et al., 1993), and finding other plant species that can produce paclitaxel such as *T. chinensis*, *T. wallichiana*, *T. brevifolia*, *T. canadensis*, *T. floridana*, *T. cuspidata*, *T. baccata*, and *T. globosa* (Liu et al., 2016).

Biotechnological Strategies

On the other hand, biotechnology and metabolic engineering were extensively used to increase the production of paclitaxel.

Cell Culture

The first report of *Taxus* cell culture was in a patent application and an abstract (Christen et al., 1989, 1991). Back in 1992, Fett-Neto and coworkers managed to obtain an in vitro culture of the taxol-producing plant *T. cuspidata* (Fett-Neto et al., 1992). With the development of tissue culture technologies via optimization of culture conditions such as media composition procedure, specific metals, light, pH, temperature, osmotic pressure, gas contents, and elicitors exploitation (Howat et al., 2014), higher amounts of taxol could be produced from a variety of species such as *T. cuspidate*, *T. canadensis*, *T. baccata*, *T. globosa*, *T. yunnanensis*, and *T. chinensis*, with taxol production improved from ~1–3 mg/L to ~77.5–153 mg/L (Liu et al., 2016; Onrubia et al., 2013; Bringi et al., 1995).

Heterologous Expression Systems

Another biotechnological approach for steady and enhanced production of taxol is the use of heterologous expression systems. In these systems a part of genome encoding the biosynthetic pathway of taxol is transferred to another type of cells through a suitable transformation vector. New cells will start the expression of enzymes necessary for taxol production, whose speed and yield could be engineered by controlling the fermentation conditions. Some of the successful heterologous expression systems are Saccharomyces cerevisiae yeast model (Dejong et al., 2006), Escherichia coli bacteria model (Ajikumar et al., 2010), and Arabidopsis thaliana plant model (Besumbes et al., 2004). The highest taxol content from E. coli was 570 mg/L (Biggs et al., 2016). Taxadiene is the first diterpene intermediate involved in taxol biosynthesis through baccatin III. Its production exhibited 40-fold increase in S. cerevisiae via retarding the competition between steroid and taxol biosynthesis in favor to taxol. That was done by the expression of GGPPS, tHMGR1, and flux control by transcription factor UPC2-1 (Engels et al., 2008). Moreover, acetyl-CoA accumulation in S. cerevisiae via acetaldehyde dehydrogenase and acetyl-CoA synthetize overexpression was found to increase the terpenoid content over steroid content in transgenic yeasts which can help in increasing taxol production (Shiba et al., 2007). Recently, Abdallah and coworkers managed to engineer the first B. subtilis strain that can produce taxadiene. They expressed the plant-derived enzyme taxadiene synthetize and overexpressed the eight key biosynthetic enzymes of MEV pathway to enhance geranylgeranyl pyrophosphate (GGPP) flux that led to a ~82-fold increase in taxadiene content when compared to wild-type B. subtilis. Taxadiene content reached ~17.7 mg/L (Abdallah et al., 2019). In the transgenic plant heterologous expression model Arabidopsis thaliana, Besumbes and coworkers managed to produce taxadiene in a yield of 600 ng/g of dry weight by overexpressing TASY and GGPP synthase. However, growth retardation resulted from taxadiene accumulation (Besumbes et al., 2004). On the other hand, TASY

overexpression in another model, *Physcomitrella patens* resulted in taxadiene production in a yield of 0.05% from fresh weight without growth retardation (Anterola et al., 2009).

IPP and GGPP are very important targets in taxol biosynthesis since they are the rate-limiting intermediates for this process. While trying to enhance IPP-mediated terpenoids biosynthesis, isopentenol utilization pathway (IUP) was transformed to *E. coli*, resulting in continuous phosphorylation of isopentenol (prenol or isoprenol) isomers and increased IPP production (Chatzivasileiou et al., 2019). IUP pathway optimization only needs the cofactor adenosine triphosphate, and it occurs within two reactions. Thus, IUP pathway is far much simpler than MEP or MVA pathways (Chatzivasileiou et al., 2019).

Overexpression Positive Effectors

Overexpression positive effectors can also play an important role in taxol biosynthesis. In 2017 Yu *et al.* suggested that manipulating the expression of some candidate genes in taxol biosynthetic pathway can affect taxol contents. Overexpression of the hydroxylation and acetylation enzymes (i.e., 2-debenzoyl-7, 13-diacetylbaccatin III-2-O-benzoyl-transferase (DBBT), 5-alpha-taxadienol-10-beta-hydroxylase (T10OH), taxadiene 5-alpha-hydroxylase (T5H), and taxadienol acetyltransferase (TAT)) increased taxol content. Later on, Wang and coworkers confirmed this approach when they revealed a \sim threefold increase in taxol content in *T. wallichiana* with overexpression of the genes WRKY, bHLH, MYB, and ERF (Wang et al., 2019). Taxadiene synthase (TS) is a pivotal enzyme in taxol biosynthesis (Köksal et al., 2011).

Knock-Outing Negative Effectors

Knock-outing negative effectors together with overexpression of positive effector genes can result in dramatic increase in taxol levels via blocking the competing downstream metabolic pathways. Sabzehzari and Naghavi (2018) used CRISPR/ Cas9 and miRNA technology to achieve this strategy, in order to silence the genes involved in sterol biosynthesis that encode lanosterol/squalene synthase (Do et al., 2009) competing with terpene pathway for farnesyl pyrophosphate (FPP) (Sabzehzari & Naghavi, 2018).

Moreover, miR171 and miR164 were found to decrease paclitaxel biosynthesis in *T. baccata* (Hao et al., 2012). MiR171 and miR164 inhibit taxane 2α -obenzoyl-transferase and taxane 13α -hydroxylase, which are very important for the biosynthesis path of taxol (Ramírez-Estrada et al., 2016). Therefore, silencing MiR171 and miR164 using CRISPR/Cas9 or miRNA can promote paclitaxel production.

Camptothecin (CPT)

Camptothecin (CPT) is a modified monoterpene indole alkaloid (Fig. 1.2) possessing potent antitumor activities. FDA approved some camptothecin (CPT) derivatives, such as topotecan and irinotecan, which were developed to overcome poor water solubility and severe side effect, for clinical treatment of some human tumors such as ovarian cancer, colorectal cancer, and leukemia via binding DNA topoisomerase I and DNA leading to ternary structure inhibiting DNA relegation and DNA damage (Venditto & Simanek, 2010; Hsiang et al., 1985; Sirikantaramas et al., 2007). The main commercial source of CPT, which is the main starting material for the clinically used synthetic derivative, is some woody plant species such as Camptotheca acuminata and Nothapodytes foetida (Wall et al., 1966; Kai et al., 2015). The slow growth of such species, the low yield of CPT (average, 1 mg/g) (Lopez-Meyer et al., 1994), and environmental concerns of overharvesting woody plants cannot fulfill the increasing market demand that recently reached 2.2 billion US dollars in 2008 in the form of topotecan and irinotecan sales with high probability of increase (Kai et al., 2014). Meanwhile, combinatorial synthesis is yet not effective (Kai et al., 2015). Hence, improvement of CPT production could be achieved through the CPT production through biotechnology techniques such as introducing biosynthetic genes encoding enzymes responsible of its production to rapidly growing vector cells such as hairy roots which are rapidly growing, with genetic stability, easily elicited, and hormone free (Shi et al., 2020).

Biosynthetic Background

The biosynthetic pathway for CPT production is similar to several other terpenoid indole alkaloids (TIA), originating from strictosidine which is a condensed product of both the amino acid derivative tryptamine (nitrogen bearing moiety) and the monoterpene secologanin (terpenoid moiety). CPT biosynthesis (Fig. 1.3) was extensively studied over the last three decades (Kai et al., 2015). Knowledge of biosynthetic genes encoding enzymes for TIA and CPT enabled the production of transgenic cell suspension, hairy roots or seedlings with increased CPT yields, and

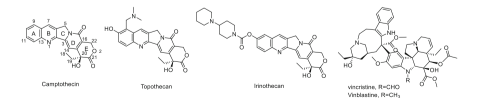


Fig. 1.2 Structures of some clinically important indole alkaloids

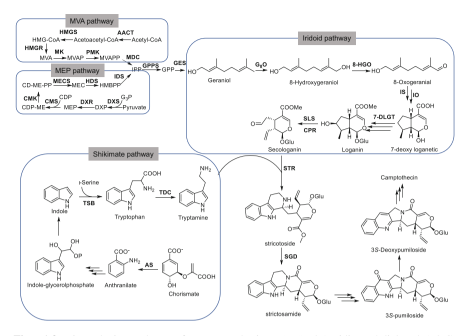


Fig. 1.3 Biosynthetic pathway for camptothecin [*CMS* 4-(cytidine 5-diphospho)-2-Cmethylerythritol synthase, *CMK* 4-(cytidine 5-diphospho)-2-C-methylerythritolkinase, *MECS* 2-C-methylerythritol-2,4-cyclodiphosphate synthase, *HDS* hydroxymethylbutenyl 4-diphosphate synthase, *IDS* IPP/DMAPP synthase, *GPPS* geranyl pyrophosphate synthase, *CPR* NADPHcytochrome P450 reductase, *GES* geraniol synthase, *G80* geraniol-8-hydroxylase, 8-HGO 8-hydroxy-geraniol oxidoreductase, *IS* iridoid synthase, *IO* iridoid oxidase (CYP76A26), *DLGT* 7-deoxyloganetic acid UDP-glucosyltransferase, *SLS* secologanin synthetase, *AACT* acetyl-CoA C-acetyltransferase, *HMGS* 3-hydroxy-3-methylglutaryl-CoA synthase, *HMGR* 3-hydroxy-3methylglutaryl-CoA reductase, *MK* mevalonate kinase, *PMK* phosphomevalonate kinase, *MDC* mevalonate 5-diphosphate decarboxylase, *STR* strictosidine synthase, *SGD* strictosidine betaglucosidase; multiple arrows indicate multiple steps]

stable rapid production and that can overcome the difficulties of finding a stable transformation system for woody plants (Ni et al., 2011).

Hairy Root Culture

Interestingly in 2001, Saito and coworkers established a successful hairy root culture system of *Ophiorrhiza pumila* (Rubiaceae) transformed by *Agrobacterium rhizogenes* strain 15834, which carries the PGSGluc1 gene plasmid. The hairy roots grew in liquid culture media with 16-fold increase in the first 5 weeks. It produced CPT as the main alkaloid with 0.1% yield of dry weight. Meanwhile, CPT was found to accumulate in culture media and its production was increased by

adding the polystyrene resin (Diaion HP-20) to the culture media to absorb CPT and facilitate its regeneration and later release from the resin, rendering this method more suitable for commercial production (Saito et al., 2001).

Moreover, the jasmonate-responsive APETALA2 domain transcription factor called ORCA3 isolated from *C. roseus* strongly upregulated the expression of several TIA biosynthetic genes (van der Fits & Memelink, 2000). *C. acuminata* transgenic hairy root lines with overexpressed ORCA3 produced CPT with 1.5-fold increase (1.12 mg/g dw) (Ni et al., 2011).

In 2020 Shi and coworkers managed to increase the production of CPT from the hairy roots of *Ophiorrhiza pumila* which originally produce CPT in higher amounts compared to the wild-type plantlets and cell suspension culture that did not accumulate CPT. Comparative transcriptome analysis revealed that hairy roots of *Ophiorrhiza pumila* contained genes encoding key enzymes involved in CPT biosynthesis, which were mainly identified as *OpG10H* and *OpSLS*. The knockout of these genes by CRISPR-Cas9 editing resulted in CPT-free hairy roots whereas individual overexpression of both genes resulted in two transgenic hairy root lines, G line for *OpG10H* and S line for *OpSLS*, showing increased CPT contents up to 2.40 mg/g and 3.28 mg/g, respectively. On the other hand, SG line with overexpression of both genes exhibited CPT contents up to 3.50 mg/g (Shi et al., 2020).

Endophytes

On the other hand, several reports stated the isolation of CPT from plant endophytic fungi such as *Entrophospora infrequens*, isolated from the inner bark of *Nothapodytes foetida* (Puri et al., 2005; Amna et al., 2006), a novel endophytic fungus from the bark of *C. acuminata* (Kusari et al., 2009), and *Fusarium solani* from *Apodytes dimidiata* (Shweta et al., 2010). Although the yield of CPT from endophytic fungi was very low, it provides incentives for stable production of CPT through the improvement of culture conditions or metabolic engineering of the fungi via identification and manipulation of its biosynthetic gene clusters (BGCs).

Vinca Alkaloids (Vincristine (VCT) and Vinblastine (VBL)

Other anticancer alkaloids produced through TIA biosynthetic pathway and have increased market demand and clinical use are *Vinca* alkaloids, vincristine (VCT) and vinblastine (VBL) produced by *Catharanthus roseus* (Jacobs et al., 2004).

Such alkaloids are produced in very low yield and difficult to meet commercial pharmaceutical production. Meanwhile, they have multiple chiral centers, rendering their synthesis very difficult (O'Connor & Maresh, 2006).

Biosynthetic Background

They are formed from strictosidine through a stereoselective Pictet-Spengler reaction between tryptamine and secologanin (Kutchan, 1993; Stöckigt & Zenk, 1977; Maresh et al., 2008), catalyzed by the enzyme STR1 that is highly substrate-specific, limiting precursor-directed biosynthesis in plant cell culture to produce natural product analogs (McCoy & O'Connor, 2006).

Several attempts have been made to produce bis-indole *Vinca* alkaloids through biochemical engineering and cell suspension cultures without any successful result. Therefore, other trials have been directed toward metabolic engineering and manipulation of TIA pathway (Zhao & Verpoorte, 2007).

In 2007 Peebles and coworkers exploited ethanol-inducible promoter system to increase the biomass of *C. roseus* hairy roots to enable the production of higher amounts of the target *Vinca* alkaloids. Ethanol concentration greatly affected ethanol-inducible promoter at very low concentration of 0.005% that resulted in a sixfold increase in chloramphenicol acetyltransferase (CAT) reporter activity after 24 h and 24-fold after 72 h with 0.5% ethanol. Being commercially available and biodegradable, ethanol induction can be promising tool to increase *Vinca* alkaloids production (Peebles et al., 2007).

Sharma and coworkers found that overexpression of two upstream TIA pathway genes tryptophan decarboxylase (CrTDC) and strictosidine synthase (CrSTR) enhanced terpenoid indole alkaloid (TIA) pathway activity and antineoplastic vinblastine biosynthesis in *Catharanthus roseus* in the transgenic whole plant produced via transformation with *Agrobacterium tumefaciens* LBA1119. The produced transgenic plant exhibited twofold increase alkaloid production, ninefold increase in vindoline and catharanthine, and fivefold increase in vinblastine production (Sharma et al., 2018).

Recently, McGehee and coworkers found that carbon-based nanomaterials (CBNs) such as multiwalled carbon nanotubes (MWCNT) and graphene enhanced biomass production of *C. roseus* cell culture under dark conditions. It also doubled the production of total alkaloids under both dark and light conditions. Further HPLC analysis revealed that vincristine and vinblastine production was dramatically increased in CBN-treated cell cultures (McGehee et al., 2019)

Artemisinin (ATN)

Another example of clinically relevant natural compound with increasing demand for pharmaceutical industry is the sesquiterpene lactone artemisinin.

Artemisinin-based combination therapies (ACTs) are the frontline treatments for malaria (Dalrymple, 2013). Moreover, both artemisinin and its semisynthetic congener artesunate proved to be cytotoxic against several cancer cell lines and *Schistosoma* (Efferth, 2006; Utzinger et al., 2007; Efferth et al., 2011). Artesunate

exhibited strong activity against ganciclovir-resistant human *Cytomegalovirus* (Shapira et al., 2008). According to the published statistics of WHO, 229 million people are infected with malaria in 2019 and 409,000 people died of malaria in 2019 (World Health Organization, 2020). However, patients in endemic regions cannot benefit these therapies due to low yield of natural sources (White, 2008). Given the large number of cases, world demand for artemisinin is 130 tons per year (Artepal, 2010).

Biosynthetic Background

Artemisinin is a sesquiterpene lactone endoperoxide assembled from three isoprene units. The common sesquiterpene precursor is farnesyl pyrophosphate (FPP) that is generated via either the mevalonate-dependent or mevalonate-independent (MEP/DXP) pathway (Lange et al., 2000; Hunter, 2007). The committed step of artemisinin biosynthesis is the cyclization of FPP to amorpha-4,11-diene (amorphadiene) as shown in Fig. 1.4. This step is carried out by the enzyme amorphadiene synthase (ADS) (Bouwmeester et al., 1999). Amorphadiene is then oxidized by the cytochrome P450 monooxygenase, CYP71AV1, to artemisinic aldehyde (Ro et al., 2006; Teoh et al., 2009). Its reduction to dihydroartemisinic aldehyde is catalyzed by artemisinic aldehyde $\Delta 11$ (13) reductase, BDR2 (Zhang et al., 2008). Artemisinin is mainly produced in the leaves in glandular hairs. Artemisinin biosynthesis is mediated by three key enzymes exclusively expressed in apical cells of glandular trichomes. Although both apical trichome cells and

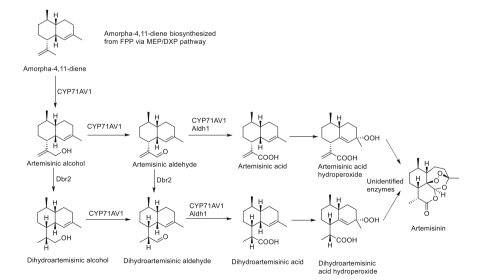


Fig. 1.4 Biosynthesis of artemisinin

subapical chloroplast-containing cells express farnesyl pyrophosphate (FPP) synthase, only glandular trichomes can produce artemisinin (Olsson et al., 2009). Artemisinin production is sensitive to variation in many factors such as genetic variation, cultivation processes, and harvesting time. Moreover, biosynthesis is affected by biotic and abiotic conditions (Omer et al., 2013, 2014, 2016; Ferreira et al., 1995; Davies et al., 2009, 2011). Artemisinin yield from *Artemisia annua* is ~0.5% of dry weight (Brisibe & Chukwurah, 2014), whereas the high-yield variety "CIM-Arogya," developed for marker-assisted breeding, has maximum yields of ~1% (Khanuja et al., 2008) which are very low yields.

All the aforementioned reasons render the development of transgenic production platforms for artemisinin, including microbes and plants essential to lower artemisinin prices and stabilize supply for millions of people who depend on the drug.

Heterologous Expression Systems

Engineering E. coli to produce a high level of artemisinin precursor was exploited. Instead of using the native MEP/DXP pathway, Martin and coworkers expressed the heterologous mevalonate pathway from S. cerevisiae in E. coli to improve isoprenoid flux (Martin et al., 2003). When coupled with the expression of ADS, amorphadiene (amorpha-4,11-diene) was produced, demonstrating the usefulness of this approach. However, a high level of expression of the mevalonate pathway resulted in accumulation of pathway intermediates and inhibition of cell growth (Pitera et al., 2007). Metabolite analysis revealed the accumulation of HMG-CoA, suggesting that the bottleneck was HMG-CoA reductase (HMGR). By modulating the expression level of this enzyme via addition of another copy of the truncated HMGR (tHMGR), the growth rate was restored. To further alleviate this ratelimiting step, bacterial HMGR genes, mvaA from S. aureus and mvaE from Enterococcus faecalis, were investigated. The strain containing mvaA from S. aureus produced the highest titer, compared to those containing the yeast or E. faecalis HMGR gene. In light of this success, another strain was constructed to replace the yeast HMG synthase (HMGS) gene with the S. aureus HMGS gene mvaS. The strain containing both mvaS and mvaA had an even higher amorphadiene titer. Under fermentation conditions where carbon and nitrogen were strictly controlled, this strain had a high titer with commercial potential (27.4 g/L) (Tsuruta et al., 2009).

Metabolic engineering of *S. cerevisiae* for artemisinic acid production was adopted as an approach to increase the production of an important artemisinin production in transgenic yeast model. As demonstrated in *E. coli*, the key to a high titer is to optimize the isoprenoid pathway flux. In yeast, this was accomplished by the upregulation of several key genes in the mevalonate pathway and the downregulation of genes for sterol biosynthesis (Ro et al., 2006). The HMG-CoA reductase in the mevalonate pathway is the principal target of complex regulation. Deletion of the N-terminal regulatory region of HMG-CoA reductase increases the

carbon flux to isopentenyl diphosphate (Donald et al., 1997). Overexpression of tHMGR increased amorphadiene production by fivefold in yeast. When tHMGR overexpression was combined with reduction in the expression of squalene synthase using the methionine-repressible MET3 promoter, another twofold increase in titer was obtained. In yeast, the sterol biosynthesis competes for the common precursor FPP. To further downregulate the sterol pathway, the upc2-1 mutant allele was overexpressed. The transcriptional factor UPC2 is a key regulator of yeast steroid uptake (Vik & Rine, 2001). Expression of the mutant upc2-1 allele allows uptake of exogenous steroids, which inhibits endogenous steroil biosynthesis. After integration of another copy of the tHMGR gene, the final strain produced 150 mg L-1 amorphadiene, nearly 500-fold higher than previously reported levels.

Overexpression of Key Biosynthetic Genes

Efforts are underway to increase artemisinin level in *A. annua* through metabolic engineering. Similar to strategies used in yeast, attempts have been made to increase carbon flux through the isoprenoid pathway for artemisinin biosynthesis in *A. annua*. One example is the expression of HMGCoA reductase gene. The HMGR gene from *Catharanthus roseus* (L.) G. Don was integrated into *A. annua* using *Agrobacterium*-mediated transformation (Aquil et al., 2009).

Based on information gathered from *N. benthamiana*, Plant Research International and Dafra Pharma International NV are collaborating to engineer chicory (*Cichorium intybus*) to produce the artemisinin precursor dihydroartemisinic acid in the roots (Brisibe et al., 2008; Dafra Pharma, n.d.). Dihydroartemisinic acid, like artemisinic acid, can be chemically converted to artemisinin at low cost. As a member of the Asteraceae family, chicory already produces considerable amounts of sesquiterpene lactones through the isoprenoid pathway. Chicory is a wellestablished plant for industrial nonfood applications, and the entire chain of largescale agricultural production is already in place. Experiments are underway to evaluate the expression of HMGR, FPS, and ADS in chicory (Ye & Bhatia, 2012).

Silymarin

Phenolic compounds are among the most ubiquitous plant secondary metabolites that have diverse structures and biological activities (Ammar et al., 2012; El-Desoky et al., 2018). They could be found in almost all plant genera. Phenolic compounds vary in their structure complexity from simple phenolic acids to highly complex tannins (Elkhateeb et al., 2019). Among phenolic compounds with high medicinal value is silymarin. Silymarin is a flavolignan mixture consisting of silybin (SB), isosilybin (ISB), silydianin (SD), silychristin (SC), and taxifolin (TXF) as shown in Fig. 1.5, extractable from milk thistle's (*Silybum marianum* L. Gaertn.) seeds and

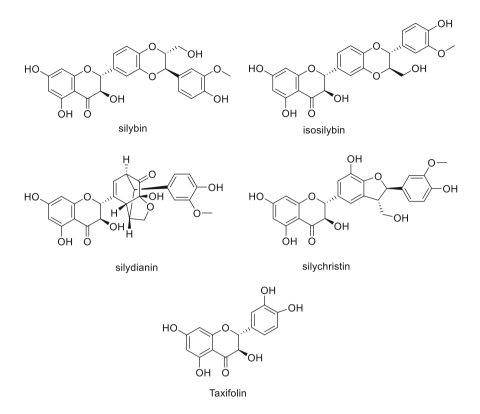


Fig. 1.5 The main compounds of silymarin (flavolignan mixture)

fruits (Khalili et al., 2009a, 2009b; Elyasi, 2021; AbouZid & Ahmed, 2013). Several research projects and studies have been conducted to increase the silymarin content of *Silybum marianum* seeds. A large part of these studies were concerned with environmental factors, whether edaphically or non-edaphically, and agricultural treatments such as irrigation, fertilization, planting dates, and harvesting in order to increase the content of silymarin in seeds (Elsayed et al., 1993; Omer et al., 1993). Silymarin is used worldwide as liver support treatment for several hepatic dysfunctions (Crocenzi & Roma, 2006). Being the main active component (60–70%), silybin reserves the main potential in silymarin hepatoprotective activities (Negi et al., 2007).

Chalcone synthase (CHS, EC 2.3.1.74) is a key enzyme for the naringenin chalcone biosynthesis catalyzing sequential addition of three acetate units from malonyl-CoA to p-coumaroyl-CoA (Spribille & Forkman, 1982). Then naringenin is produced by cyclization and hydroxylated to taxifolin and then silybin is produced by other steps. Therefore, CHS overexpression is a promising target to increase silybin production.

After detailed bioinformatics study, Rahnama and coworkers overexpressed chalcone synthase (chsA) of *Petunia hybrid* L. petals in the hairy root of *Silybum marianum*. ChsA gene was transformed to *S. marianum* hairy roots by *A. rhizogenes* AR15834 strain harboring pCamCHS vectors. That resulted in a sevenfold overexpression of chsA gene in transgenic hairy roots. And subsequent ten times increase in silybin content in transgenic hairy roots when compared to non-transgenic ones (Rahnama et al., 2013).

Some other factors were found to enhance the production of silymarin in the hairy root cultures of *S. marianum* such as the use of elicitors. Among these elicitors is the abiotic elicitor salicylic acid, which enhanced silymarin production (1.89 mg/g dry weight) in hairy roots of *S. marianum*. Salicylic acid (6 mg/50 ml culture) after 24 h increased silymarin content 2.42 times compared to control (0.78 mg/g dry weight). The content of silybin, isosilybin, silychristin, silydianin, and taxifolin were 0.703, 0.017, 0.289, 0.02, and 0.863 mg/g dry weight, respectively. Lipoxygenase activity was increased by ~1.57-fold (Khalili et al., 2009a, 2009b). Similarly, the same group studied the effect of silver ion [Ag]⁺. Ag⁺ was found to increase the activity of lipoxygenase that mediates signal transduction pathway for silymarin production. That resulted in a fourfold increase in silymarin production at 2 mM Ag⁺ and after 96 h after elicitation, when compared to non-treated hairy roots (Khalili et al., 2010).

Another type of elicitors is biotic elicitors. Hasanloo and coworkers studied the effect of three fungal species, namely, *Fusarium proliferatum, Aspergillus niger*, and *Rhizoctonia solani*, as elicitors for hairy root cultures of *Silybum marianum* (L.) Gaertn. The maximum production for each of the fungal treatment was different: 10 mg *A. niger*/50 ml culture produced 0.18 mg/g dry weight of silymarin after 48 h while 20 mg *F. proliferatum*/50 ml culture produced 0.22 mg/g dry weight of silymarin after 72 h. The best profile owed to *F. proliferatum* treated hairy roots that contained taxifolin (0.068 mg/g dry weight), silydianin (0.11 mg/g dry weight), silybin (0.021 mg/g dry weight), and isosilybin (0.015 mg/g dry weight) which were 1.94-, 1.19-, 2.62-, and 1.25-fold greater than untreated hairy roots, respectively. The results indicated that the type of fungi, their concentrations, and their exposure time have significant effect on the stimulation of silymarin production (Hasanloo et al., 2013).

Conclusion

The abovementioned engineered metabolites highlight important aspects to be considered for such objective. Specifications of the host organism have a high impact on final commercial success. These specifications include metabolic capabilities, genetic systems, and scale-up potential. The limitations of the genetic system for plants or even microbes could present challenges. As the optimal host is chosen, the strategy for metabolic engineering may involve determining a specific biosynthetic pathway and enzymes to target. The full understanding of the biochemistry and gene regulation networks is critical for pathway construction. Optimization of pathways involves balancing the upstream and downstream steps as well as exclusion or decrease of challenging reactions.

Biotechnology and metabolic engineering are extensively used to increase the production of paclitaxel.

The production of CPT from the hairy roots of *Ophiorrhiza pumila* which originally produce CPT in higher amounts compared to the wild-type plantlets and cell suspension culture that did not accumulate CPT was managed to increase its production. Comparative transcriptome analysis revealed that hairy roots of *Ophiorrhiza pumila* contained genes encoding key enzymes involved in CPT biosynthesis, which were mainly identified as *OpG10H* and *OpSLS*. Knockout of these genes by CRISPR-Cas9 editing resulted in CPT-free hairy roots. Overexpression of two upstream TIA pathway genes tryptophan decarboxylase (CrTDC) and strictosidine synthase (CrSTR) enhanced terpenoid indole alkaloid (TIA) pathway activity and antineoplastic vinblastine biosynthesis in *Catharanthus roseus* in the transgenic whole plant produced via transformation with *Agrobacterium tumefaciens* LBA1119.

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Efforts are underway to increase artemisinin level in *A. annua* through metabolic engineering. Similar to strategies used in yeast, attempts have been made to increase carbon flux through the isoprenoid pathway for artemisinin biosynthesis in *A. annua*. One example is the expression of HMGCoA reductase gene.

The studies with hairy roots indicated that the type of fungi, their concentrations, and their exposure time have significant effect on the stimulation of silymarin production.

Mechanisms of metabolite engineering are the most promising ways to contribute to the provision of many important entities in the field of pharmaceuticals and medicines, especially high-cost treatments, such as cancers, liver diseases, and viruses.

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