Microbial Cell Factories for Green Production of Vitamins

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Vitamins are a group of essential nutrients that are necessary to maintain normal metabolic activities and optimal health. There are wide applications of different vitamins in food, cosmetics, feed, medicine, and other areas. The increase in the global demand for vitamins has inspired great interest in novel production strategies. Chemical synthesis methods often require high temperatures or pressurized reactors and use non-renewable chemicals or toxic solvents that cause product safety concerns, pollution, and hazardous waste. Microbial cell factories for the production of vitamins are green and sustainable from both environmental and economic standpoints. In this review, we summarized the vitamins which can potentially be produced using microbial cell factories or are already being produced in commercial fermentation processes. They include water-soluble vitamins (vitamin B complex and vitamin C) as well as fat-soluble vitamins (vitamin A/D/E and vitamin K). Furthermore, metabolic engineering is discussed to provide a reference for the construction of microbial cell factories. We also highlight the current state and problems encountered in the fermentative production of vitamins.

Keywords: vitamins, metabolic engineering, microbial cell factory, chemical synthesis, biosynthesis

INTRODUCTION

Vitamins are essential for proper growth and health of animals, that cannot produce vitamins by themselves or that synthesize insufficient amount to cover all their needs (Capone and Sentongo, 2019; Suter, 2020). The methods of producing vitamins are based either on chemical synthesis or fermentative production (Yuan et al., 2020).

There are at least 30 kinds of different compounds considered "vitamins," more than 20 vitamins of which are known to be necessary for biological health. Vitamins are either water-soluble or fat-soluble. As the name suggests, a water-soluble vitamin dissolves in water easily and insoluble in organic solvents. After absorption, the body stores very little of such proteins, and most are excreted with urine (Berdanier and Adkins, 2019). Fat-soluble vitamins are dissolved in fats but not in water, and which are stored in the liver or fatty tissues for future use. While vitamins are essential nutrients for all living things, many plants and microorganisms can synthesize them naturally by themselves. By contrast, humans and other animals need to acquire sufficient vitamins with their diet or through supplements to maintain optimal health (Blake and Konings, 2019).

Traditionally, vitamin production strains have been improved through mutagenesis and metabolic engineering, which can be conducted either through chemical or biological means.
(Vandamme and Revuelta, 2016b). The main chemical strategies include chemical mutagenesis, application of N\(^{+}\) ion beam, ultraviolet radiation or laser mutagenesis. The biological methods mainly include the construction and mutagenesis of the starting strain, genetic modification, synthetic biotechnology, optimization of media and culture conditions, construction of biofilm reactors, etc. (Nie et al., 2013; Song et al., 2014). A series of biotechnological methods are used to transform the metabolic network of cells to construct a programmable “chassis” and “programmable” whole, which can be used to develop an effective assembly strategy, test the adaptability of external components and modules after loading, forming a fine-tuned and customized biological application system. To drive the iterative evolution of other industrial strains, and effectively promote the transformation and renewal of high vitamin producing strains. Chemical methods are usually expensive, environment-unfriendly, waste-prone, and the costly waste disposal. However, the microbial fermentation method has attracted much attention due to low cost, low energy consumption and easy waste recycling. At present, the fermentation method has been recognized by researchers, and it is more environment-friendly and safe than chemical methods. As the fermentation technology matures, this approach is increasingly being used in industry to increase the production of different vitamins. For example, fermentation processes for the production of vitamin B\(_3\) (VB\(_3\)), vitamin B\(_12\) (VB\(_{12}\)), vitamin C, and vitamin K2 have all been industrialized successfully.

Acevedo-Rocha et al. (2019) reviewed the fermentation of B vitamins from the aspect of sustainability. In this review, we mainly discuss vitamins that can be produced by green fermentation processes. It covers water-soluble vitamins, including vitamin C and vitamin B complex (thiamine, riboflavin, niacin, pantothenic acid, pyridoxine, biotin, folate, and cobalamin) as well as the fat-soluble vitamin E and vitamin K. Here, we discussed the producing microorganisms, advanced biological methods and metabolic bottlenecks of different vitamins.

**WATER-SOLUBLE VITAMINS**

**B Vitamins**

The global demand for B vitamins is growing due to wide applications in food, pharmaceuticals, feed, and other fields. Although most vitamins are manufactured by chemical synthesis, successful industrial bioprocesses have been established for the production of VB\(_2\) and VB\(_{12}\). The underlying extraordinary achievement in metabolic engineering is discussed in this article.

**Vitamin B\(_1\)**

Vitamin B\(_1\), which is also known as thiamine, was the first B vitamin to be identified. Thiamine pyrophosphate (TPP), the active form of thiamine, can inhibit the activity of cholinesterase, reduce skin inflammation, prevent seborrheic dermatitis, or eczema, and improve skin health. Thiamine biosynthesis results from the coupling of the pyrimidine and the thiazole moieties to form thiamine phosphate (Dorrestein et al., 2004; Jurgenson et al., 2009; Cea et al., 2020). *Escherichia coli*, *Salmonella typhimurium*, and *Bacillus subtilis* are the most thoroughly studied thiamine production organisms (Begley et al., 1999).

In chassis cell *S. typhimurium*, the thiamine pyrimidine moiety can be produced through de novo purine biosynthesis or independently of the pur\(E\) gene through the alternative pyrimidine biosynthesis (APB) pathway (Downs and Roth, 1991; Downs, 1992). According to the phenotypic characteristics of the *apbA* mutant, follow-up studies concluded that the functional APB pathway is essential for thiamine synthesis when *S. typhimurium* grows in the presence of exogenous purines (Downs and Petersen, 1994). Research has shown that overexpression of *thiA*, *nmtA*, and *thiP* in *Aspergillus oryzae* can increase the vitamin B\(_1\) yield fourfold compared to the wild-type (Tokui et al., 2011). Based on the riboswitch mechanism, mutations in the genes of thiamine pyrophosphate kinase activity (*thiN*) and thiamine-related transport proteins (*YkoD* and *YuAI*) were introduced in *B. subtilis* TH95. It was recently reported that thiamine biosynthesis is strictly regulated by TPP riboswitches in bacteria/eukaryotes and transcriptional repressors in archaea (Hwang et al., 2017). *E. coli* has emerged as the preferred cell factory for TPP production after a riboswitch-based biosensor enabled the discovery of thiamine transporters, combined with overexpression of the native *thiFGHCE* and *thiD* genes, which are closely related to Fe-S metabolism (Figure 1A and Table 1; Cardinale et al., 2017).

However, *thiC/thiH* in the thiamine biosynthetic pathway is involved in Fe–S metabolism and is inhibited by S-adenosylmethylionine (SAM) metabolites, and the catalytic activity of ThiC enzyme (Figure 2) is very low (\(k_{cat} = 0.002\) s\(^{-1}\)) which is one of the main metabolic bottlenecks (Palmer and Downs, 2013). In addition, the cost of chemical production of thiamine is very low, and the production of engineered strains needs to be increased to be expected to be industrialized.

**Vitamin B\(_2\)**

Riboflavin is an important precursor of flavin mononucleotide (FMN) and flavin adenine dinucleotide (FAD) (Balasubramaniam et al., 2019; Andreieva et al., 2020). Riboflavin insufficiency manifests as persistent anemia (Shi et al., 2014). The biosynthesis of riboflavin begins with guanosine triphosphate and ribose-5-phosphate, followed by six enzymatic steps (Fischer and Bacher, 2005). Burgess et al. (2004) found that overexpression of the ribABC*GH* genes can increase riboflavin production. Later, it was found that there were both nucleotide substitutions and deletions in the regulatory region of the rib operon. By deregulating the rib operon and purine pathway of *B. subtilis*, riboflavin production was greatly improved. The specific genetic engineering steps included overexpression of the rib*AB* gene and deletion of the pur*R* gene, after which maximum output of riboflavin reached more than 826.52 mg/L (Figure 1B; Shi et al., 2014). In *Candida famata* overexpression of *sef1* and *imh3* was combined with classic mutagenesis methods to construct the high riboflavin-producing strain AF-4. As a result, 1026 ± 50 mg/L of riboflavin can be produced during a fed-batch cultivation in a lab-scale fermenter. This research has
made a great contribution to industrial production of riboflavin (Dmytruk et al., 2011, 2014). The two most important industrial producers are Ashbya gossypii and B. subtilis.

In A. gossypii, malate synthase in the glyoxylate cycle is essential for riboflavin production. Deletion of the malate synthase gene (ACR268C) decreased riboflavin production 10-fold compared to the wild-type strain. Conversely, overexpression of the ACR286C gene significantly increased the yield of riboflavin by 70%. These results demonstrated that malate synthase is a new target for improving the production of riboflavin (Sugimoto et al., 2009). Abbas and Sibirny (2011) introduced the icl gene, overexpressed the gly1, prs2,4, and prs3 genes, as well as knocking out the vma4, shm2, and bas1 genes, resulting in riboflavin production of more than 20 g/L.

In B. subtilis, Schwechheimer et al. (2016) overexpressed riboflavin biosynthesis genes, decreased the activity of the flavin kinase RibCF, and improved the de novo purine synthesis and pentose supply, after which the riboflavin yield reached more than 26 g/L. At present, the bottleneck of riboflavin production is mainly due to the poor genetic stability of the engineered strain, and more by-products produced by fermentation, which restrict the high yield of riboflavin.

**Vitamin B₃**

Niacin is the precursor in the synthesis of the pyridine coenzymes NAD (nicotinamide adenine dinucleotide) and NADP (nicotinamide adenine dinucleotide phosphate) (Chand and Savitri, 2016; Chauhan and Poddar, 2019; Tannous et al., 2020). It is found at relatively high concentrations in internal organs of animal, muscle tissues, and fruits. Currently, niacin is mainly used as a feed additive to increase the utilization of feed protein, or as a pharmaceutical intermediate in the synthesis of various drugs. So far, there is no systematic description of a commercial fermentation process of nicotinic acid (NA) or nicotinamide (NAM). Industrial production methods are mainly ammonia oxidation and electrolytic oxidation, but the former has
### TABLE 1 | Water-soluble vitamins produced by biotechnological methods.

| Vitamins | Strains | Biotechnological method | Medium and precursor | Yield | References |
|----------|---------|-------------------------|----------------------|-------|------------|
| Vitamin B<sub>1</sub> | **B. subtilis** TH95 | Mutation of gene encoding thiamine pyrophosphate kinase activity (thiN) and thiamine-related transport protein (ykoD and yuaU). | MM | 1.27 mg/L | Schyns et al., 2005 |
| | **E. coli** | TPP biosensor (plasmid pTPP_Bios); Overexpression of native thiFSGH; thiC; thiE; and thiD; Genetic-metabolic coupling. | MM | 0.80 mg/L | Cardinale et al., 2017 |
| | **A. oryzae** | Overexpression of thiP, thiA, and nmtA. | CD-Dex medium (5% dextrose) | 4-fold > WT | Tokui et al., 2011 |
| Vitamin B<sub>2</sub> | **B. subtilis** | Decrease the activity of flavinase RibCF activity; Overexpression of riboflavin biosynthetic genes; improved the de novo purine synthesis and pentose supply. | MM | >26 g/L | Schwechheimer et al., 2016 |
| | **A. gossypii** | Introduced the icl gene; Overexpression of glv1, prs2,4, and prs3 genes; Knocked out vma4, shm2, and bas1 genes. | YPD; Plant oil | >20 g/L | Abbas and Sibirny, 2011 |
| | **Candida famata** | Conventional mutagenesis by overexpression of sef1 and imt9. | YPD; Fluorophenylalanine | 1026 ± 50 mg/L | Dmytruk et al., 2011 |
| Vitamin B<sub>3</sub> | Yeast | Knock out NR importer Nrt1' in the NR-non-salvaging genotype nrr1, unil, pmpl (strain PAB038). | 2x YPD; Nicotinic acid | 8 mg/L | Belenky et al., 2011 |
| | **E. coli** | Expressing R. hondocharis nitrile hydratase. | LB medium; 2YT medium | 508 g/L | Wang et al., 2017 |
| | **C. glutamicum** | Deletion of ilvA gene and overexpression of ilvBNCD and panBC genes. | MM | 1000 mg/L | Leonardi and Jackowski, 2007 |
| | **B. subtilis** | Overexpression of ilvBHCD and panBCDE; Overexpression of SterA and GlyA of the enzymes of the glycine cleavage cycle. | MM | 82–86 g/L | Hohmann et al., 2016 |
| Vitamin B<sub>4</sub> | **E. coli** | Overexpression of native Epd, PdxJ, and Dxs enzymes. | MM | 78 mg/L | Hoshino et al., 2004 |
| | **S. meliloti** | Overexpression of E. coli Epd and native PdxJ enzyme. | MM | 1.30 g/L | Hoshino et al., 2007 |
| | **IFO14782** | Overexpression of E. coli PdxA and native PdxJ enzymes. | MM | 65 mg/L | Commichau et al., 2015 |
| | **B. subtilis** | Overexpression of E. coli PdxJ and native PdxJ enzymes. | MM | 508 g/L | |
| Vitamin B<sub>5</sub> | **Agrobacterium**/ **Rhizobium** HK4 | Overexpression of a strong biotin operon from E. coli; Use of a powerful artificial tac promoter and introduct of a modified RBS in front of BioB. | MM; Betaine; Diaminononanoic acid | 110 mg/L | Streit and Entcheva, 2003 |
| | **E. coli** | Overexpression of native biotin operon from a high-copy number plasmid | MM; H-medium | 11 mg/L | Ifuku et al., 1995 |
| | **B. subtilis** | Overexpression of native biotin operon and selection on S-2-aminoethyl-L-cysteine. | MM | 21 mg/L | Van Arsdel et al., 2005 |
| Vitamin B<sub>6</sub> | **A. gossypii** (ATCC 10895) | Overexpression of FOL genes and deletion of AgMEY7; Deletion of AgADE12 and AgRIB1 at the same time. | MA2 rich medium | 7 mg/L | Serrano-Amartrian et al., 2016 |
| | **S. melloti** (MC5-2) | High throughput screening of mutants using riboswitch ARTP-irradiation was used to induce random mutations; Deletion of cobB; Overexpression of hemE. | MM; Cobalt chloride; DMBI | 156 ± 4.20 mg/L | Cai et al., 2018 |
| | **P. denitrificans** | Random mutagenesis and genetic engineering; Overexpression of cobF-cobM gene cluster and cobA and cobC genes; Optimize the best PH range; Optimize promoters. | MM; Choline chloride | 214.30 mg/L | Li et al., 2008 |
| | **E. coli** | Heterologously expressed the hemO, hemB, hemC, and hemD genes etc.; Optimizing of fermentation conditions. | CM medium | 0.67 mg/L | Fang et al., 2018 |
| | **Propionibacterium shermanii** | Overexpression of biosynthetic genes. | MM; DMBI | 206 mg/L | Sych et al., 2016 |
| | **S. cerevisiae** and **Zygosaccharomyces bailii** | Overexpressing the endogenous D-arabinono-1,4-lactone oxidase and L-galactose dehydrogenase (overexpression of Igdh and alo1). | MM | 100 mg/L | Sauer et al., 2004 |
| | **K. vulgare DSM 4025** | Oxidation and lactonization. | L/D-sorbose; Glyceral; Baker's yeast | 1.37 g/L | Sugisawa et al., 2005 |

(Continued)
TABLE 1 | Continued

| Vitamins | Strains | Biotechnological method | Medium and precursor | Yield | References |
|----------|---------|-------------------------|---------------------|-------|------------|
| X. campesris 2286 | Lactonation under oxidative stress; Direct synthesis of glucose (carbohydrate source) induced by free radicals (HClO₄ treatment). | MM; K₃HPO₄; Urea | 20.40 g/L | Rao and Sureshkumar, 2000 |
| G. oxydans and K. vulgare and B. endophyticus | Cell–cell interaction; One step 2-KGA fermentation. | d-sorbitol | 73.70 g/L (2-KGA) | Ma et al., 2019 |

high production costs and needs to be above 300°C during the reaction, and the latter has low costs of production, however, the efficiency of electrolysis is not high, which limits the industrial production of niacin (Chand and Savitri, 2016).

Recent reports describe the use of recombinant E. coli expressing Rhodococcus rhodochrous nitrile hydratase for vitamin B₃ production. At low cell density, nicotinamide was produced in fed-batch mode, and the product concentration reached 390 g/L. After high-density culture in 5 L bioreactor, the concentration of nicotinamide reached 508 g/L in 60 min (Figure 1D; Wang et al., 2017). Belenky et al. (2011) showed that the disruption of nrt1 results in increased export of nicotinamide riboside (NR). Moreover, disruption of the niacin transporter Tna1 can also increase the output of niacin, revealing that cells regulate the intracellular NAD⁺ metabolic process by balancing the transport of niacin, the precursor of NAD⁺. On the basis of adding 5 mM niacin, yeast cells can produce 8 mg/L nicotinamide mononucleotide (Belenky et al., 2011).

Vitamin B₅
Vitamin B₅, also known as pantothenic acid, is composed of pantoic acid and β-alanine (β-Ala), which is a precursor of coenzyme A (Leonardi and Jackowski, 2007). It plays an important role in maintaining the health of skin and blood. Its general function is to participate in the production of energy in the body, but it can also control the fat metabolism, and is also an essential nutrient for the brain and nerves. There are chemical and microbial synthesis methods for the synthesis of pantothenic acid, whereby microbial methods can be used to directly synthesize optically pure D-pantothenic acid.

Sahm and Eggeling (1999) adopted a series of methods to increase the production of pantothenic acid, including the deletion of the ilvA gene and the overexpression of the ilvBNCD and panBC genes. The pantothenic acid production of the best strain reached 1000 mg/L (Figure 1F; Sahm and Eggeling, 1999). Huser et al. (2005) also used Corynebacterium glutamicum to produce pantothenic acid. They deleted the ilvA gene, inhibited the expression of the ilvE gene and overexpressed the ilvBNCD gene. The final titer of pantothenate reached 1.75 g/L (Huser et al., 2005). Studies have shown that the specific activity of pantothenic acid synthase PanC of C. glutamicum is 205.10 U/mg. Adding substrates (D-pantothenic acid and β-Ala) to E. coli containing the enzyme can be produced 97.10 U/mg within 32 h, the conversion rate of pantothenic acid was 99.10%.

However, the reported work had production defects, which required the addition of exogenous substrate pantothenic acid, and the high market price of pantothenic acid seriously restricted the industrialization of this method. Another chasssis organism that is commonly used to produce pantothenic acid is B. subtilis. Hohmann et al. (2016) clarified the highest production of pantothenic acid by overexpressing ilvBHCD, panBCDE, serA, and glyA, as well as the enzymes of the glycline cleavage cycle the purpose is to increase the number of precursors for pantothenic acid synthesis (Figure 1F). The maximal output of the best strain reached 82–86 g/L during a 48 h fed-batch fermentation, opening up a new chapter of vitamin production in the biological world (Hohmann et al., 2016).

Vitamin B₆
There are six forms of vitamin B₆, including pyridoxine (PN), pyridoxal (PL), and pyridoxamine (PM), as well as their respective phosphate derivatives. It is a water-soluble vitamin, which exists in the form of phosphate in the body. The most versatile form of vitamin B₆ is pyridoxal 5’-phosphate (PLP), which is a cofactor of many proteins and enzymes in all organisms. As the most widely available commercial form, PN hydrochloride is extensively used in the pharmaceutical and food industries (Eliot and Kirsch, 2004).

Two de novo synthesis routes have been reported the 1-deoxyxylulose 5-phosphate (DXP)-dependent pathway and the DXP-independent pathway (Tanaka et al., 2005). From large-scale screening studies of different strains, found that the Gram-negative bacterium Sinorhizobium meliloti is the best producer of vitamin B₆, reaching a titer of 103 mg/L of B₆ isoforms within 168 h. Vitamin B₆ production was further increased to 1.30 g/L by expressing the E. coli epd gene and the native dxs gene in this S. meliloti strain (Figure 1E; Hoshino et al., 2007). E. coli and B. subtilis were also engineered to produce vitamin B₆. The vitamin B₆ production was enhanced to 78 mg/L within 31 h in E. coli, and B. subtilis produced 65 mg/L of PN when supplied with the precursor 4-hydroxy-L-threonine (4HTH) (Hoshino et al., 2004). At present, the industry mainly adopts the oxazole method to produce vitamin B₆, and the current research also focuses on the improvement of the oxazole method synthesis process. In the process of biosynthesis, the PdxJ enzyme activity is very low (kcat = 0.07 s⁻¹), and the reaction step catalyzed by this enzyme is the rate-limiting step in the VB₆ biosynthetic pathway. The intermediate metabolite 4-phosphate hydroxy-threonine (4HTP)
is cytotoxic and is also the main bottleneck of biosynthesis. Therefore, the fermentative production of vitamin B<sub>6</sub> requires more effort to meet the commercial demand.

**Vitamin B<sub>7</sub>**

Biotin is indispensable for the normal metabolism of fats and proteins (Lin and Cronan, 2011; Selvam et al., 2019). It is a nutrient necessary for human growth, development and normal function. Biotin combines with enzymes to participate in the process of carbon dioxide fixation and carboxylation in the body. The current large-scale production of D-biotin is mainly based on the Sterngach synthetic route, and the current industrial production method were improved on this basis. Unless biosynthetic methods can obtain high output at low cost, it is difficult to shake the position of chemical synthesis technology in industrial production. Nevertheless, it was recently reported that some microorganisms can overproduce biotin, which has been elaborated in C. glutamicum, Mesorhizobium loti, and S. meliloti. In Agrobacterium and Rhizobium HK40, overexpression of the biotin operon from E. coli driven by the powerful tac promoter and introducing a modified RBS in front of bioB resulted in a biotin yield of 110 mg/L (Figure 1G; Streit and Entcheva, 2003). If the native biotin operon is overexpressed in B. subtilis, most enzymes will be strongly inhibited by the by-product of SAM. However, the high demand for SAM by biotin synthase and 7,8-diaminononanoate synthase is still a bottleneck that must be addressed in future research. If lysine is supplied to B. subtilis, BioK will use lysine as the amino donor of the biotin precursor to promote the production of biotin precursor (dephosphorization biotin), and the fermentation process used carbon-limited fed-batch growth conditions with computer control of dissolved oxygen concentrations, but the maximal titer can only reach 21 mg/L biotin. Therefore, improving the catalytic mechanism of biotin synthase is also a challenge for future research (Van Arsdell et al., 2005; Lin and Cronan, 2011).

**Vitamin B<sub>9</sub>**

Naturally occurring folic acid is mostly found in the form of polyglutamic acid, and the biologically active form of folic acid is tetrahydrofolate (Myszczyszyn et al., 2019). Deficiency can lead to reduced hemoglobin content in red blood cells, impaired cell maturation and megaloblastic anemia (Lucock, 2000). B. subtilis or A. gossypii were successfully engineered to produce folic acid. Jagerstad and Jastrebova (2013) achieved a 5-methyltetrahydrofolate (THF) titer of 0.95 mg/L by increasing the supply of precursor substances and blocking the catabolic pathway of THF in B. subtilis. With the continuous research progress, A. gossypii has attracted increasing interest as the chassis strain for folic acid production. A. gossypii can synthesize 0.04 mg/L of folic acid naturally, which can reach 6.59 mg/L after metabolic engineering treatment. This is also the highest production value reported to date (Figure 1C; Serrano-Amatriain et al., 2016). Since the commercial chemical synthesis of folic acid is cheap, unless the environmentally unfriendly part of the chemical synthesis process is restricted, there is still a long way to go for the fermentation of this product.

**Vitamin B<sub>12</sub>**

Cobalamin is the only vitamin containing metal elements. Cobalamin is the general term for a class of corrin compounds containing cobalt (Osman et al., 2021). It is the largest and most complex vitamin molecule discovered so far. Vitamin B<sub>12</sub> deficiency leads to increased formation of ring sideroblasts in pre-myelodysplastic syndromes (Kitago et al., 2020).
Vitamin B<sub>12</sub> is synthesized by microorganisms through de novo synthesis or salvage synthesis in nature, but higher-animals and plants cannot produce it (Figure 1H; Fang et al., 2017). Although in the 19th century, researchers have completed the full chemical synthesis of vitamin B<sub>12</sub>, the chemical synthesis method is too complicated and expensive, so the world's major suppliers rely on microbial fermentation to produce vitamins. *Pseudomonas denitrificans* and *Propionibacterium freudenreichii* being widely used in industrial fermentation to produce vitamin B<sub>12</sub>. In order to improve the productivity of vitamin B<sub>12</sub>, researchers have adopted a random mutagenesis method to construct a vitamin B<sub>12</sub> overproducing strains by ultraviolet rays, nitrosoguanidine (NTG), nitrosomethylurethane and ethyleneimine (Blanche et al., 1992, 1995a,b). *Propionibacterium shermanii* was reported to produce vitamin B<sub>12</sub> with a maximum titer of 200 mg/L (Sych et al., 2016). However, the aerobic *P. denitrificans* remains the most used industrial host, and the effect is obvious. Moreover, *P. denitrificans* has stronger production capacity than the anaerobic strain that produces vitamin B<sub>12</sub> and is widely used in industrial production. Xia et al. (2015) increased the production of vitamin B<sub>12</sub> to 198 ± 4.60 mg/L by optimizing the fermentation medium using response surface method. Our research group used *E. coli* strain MG1655 (DE3) as the starting strain to achieve de novo synthesis of vitamin B<sub>12</sub> (Fang et al., 2018). Cai et al. (2018) later used riboswitch elements in *S. meliloti* for the first time, and successfully developed a flow cytometry high-throughput screening system for high-yield VB<sub>12</sub> strains. The vitamin B<sub>12</sub> titer of the best strain, *S. meliloti* MC5ñ2, reached 156 ± 4.20 mg/L, but the yield was still relatively low. At the same time, they also emphasized that the titer of vitamin B<sub>12</sub> is greatly dependent on the medium composition (Cai et al., 2018).

**Vitamin C**

Vitamin C, also known as L-ascorbic acid (LAA), is an important cofactor for multiple enzyme reaction in the body (Paciolla et al., 2019; Kawahori et al., 2020). It can act as an antioxidant to scavenge free radicals and reduce oxidative stress, so a rapidly expanding market is the application of LAA as an additive to cosmetic products (Timoshnikov et al., 2020). Vitamin C deficiency can result in scurvy. Recently, researchers used biochemical methods combined with DNA recombination technology to produce vitamin C.

At present, L-AA is commercially manufactured via the classic seven-step Reichstein process using D-glucose as the initial substrate. The process involves six chemical steps and one fermentation steps for the oxidation of D-sorbitol to 2-keto-L-gulonic acid (2-KGA) by *Glucobacter oxydans* and *Bacillus megaterium* (Figure 2A). Sugisawa et al. (2005) reported for the first time that *Ketogulonigenium vulgar* DSM 4025 can produce 1.37 g/L of L-AA under static culture conditions. Kim et al. (1996, 1998) reported that the respective enzymes from *Candida albicans* and *S. cerevisiae* convert not only D-arabinose to D-arabinono-1,4-lactone but also L-galactose to L-galactono-1,4-lactone in vitro. Experiments have shown that budding yeast cells overexpressing the endogenous D-arabinono-1,4-lactone oxidase and L-galactose dehydrogenase-nase can produce about 100 mg/L of L-ascorbic acid (Sauer et al., 2004). A microbiological consortium composed of *G. oxydans*, *K. vulgar*, and *B. endophyticus* was constructed to produce 2-KGA, and a final yield of 73.70 g/L was obtained within 30 h (Figure 2B; Ma et al., 2019). This result holds promise for the construction of a microbial cell factory for the production of vitamin C. However, it has been reported that mixed-bacteria fermentation can be unstable due to competition between the individual strains for nutrients and other factors. Therefore, mixed-bacteria fermentation technology has poor stability and low efficiency, which also hinders the pace of industrial production of vitamin C. Nevertheless, fermentation is expected to become the mainstream way of vitamin C production in the future if stable single strains can be used instead of mixed bacteria fermentation, while also shortening the production cycle.

**FAT-SOLUBLE VITAMINS**

**Vitamin A**

Vitamin A mainly includes β-carotene, α-carotene, and β-cryptoxanthin (Wise et al., 2021). β-carotene, a provitamin A carotenoid, is divided into all-<i>trans</i> and cis isomers (Yang et al., 2021). All-<i>trans</i>-β-carotene is the major isomer found in unprocessed carotene-rich plant foods, followed by its 9- and 13-cis isomers. β-carotene is an antioxidant, which not only inhibits singlet oxygen but also inhibits lipid peroxidation, thereby playing an important role in the prevention of disease (Kawata et al., 2018).

Carotene is mainly produced by fungi, some bacteria, and algae. For example, Yoon et al. increased the supply of IPP (isopentenyl diphosphate) and DMAPP (dimethylallyl diphosphate) through the introduction of foreign MVA (mevalonate) pathway (Figure 3D), thereby enhancing the production of carotenoids. The final engineered *E. coli* with a whole MVA pathway and β-carotene synthesis gene can produce β-carotene of 465 mg/L (Figure 3C; Yoon et al., 2009). Adenosine-triphosphate (ATP) and nicotinamide adenine dinucleotide phospha (NADPH) are two important cofactors in β-carotene biosynthesis pathway, Zhao et al. (2013) used *E. coli* as host cells, constructed and optimized a central metabolic module to increase the supply of ATP and NADPH in β-carotene synthesis pathway, thereby improving the yield of the β-carotene. Finally, the best strain CAR005 increased the β-carotene production to 2.1 g/L with a yield of 60 mg/g DCW in fed-batch fermentation (Zhao et al., 2013). Larroude et al. (2018) overexpressed heterologous carotene synthase (Crt) in *Yarrowia lipolytica* to make it produce high β-carotene. The fermentation yield of the engineered strain obtained by screening the best promoter was 1.5 g/L. By optimizing the fermentation conditions and using fed-batch fermentation, the yield of β-carotene was further increased production titer of 6.5 g/L and 90 mg/g DCW (Larroude et al., 2018). However, the insufficient number of precursors seriously hindered the industrialization process of β-carotene in the process of β-carotene synthesis in the future.
Vitamin D
Vitamin D refers to a group of fat-soluble secosteroids responsible for increasing intestinal absorption of magnesium, calcium, and phosphate, and many other biological effects. The most important compounds are vitamin D$_2$ (ergocalciferol) and vitamin D$_3$ (cholecalciferol) in vitamin D. Vitamin D can increase intestinal absorption of calcium, magnesium, and phosphate, and can prevent many diseases (Yuan et al., 2020).

It is well known that the precursor of vitamin D$_2$ is ergosterol (Papoutsis et al., 2020). Vitamin D$_2$ is widely used in medical, food and other industries. The current commercial production of ergosterol is mainly produced by yeast fermentation. Tan et al. improved the production of ergosterol by optimizing the fermentation medium and screening high ergosterol producing strains (Figure 3F). The results show that dissolved oxygen (DO) can be used as the effective control parameter for yeast fed-batch fermentation. The total yield of ergosterol can be increased to 1.16 g/L when DO was controlled at 12% (±1%) and pulse fed-batch was used (Tan et al., 2003).

Vitamin D$_3$ cannot play a direct role in human and animals, but it can produce the physiologically active form 25-hydroxyvitamin D$_3$ (25-OH-VD$_3$) through the metabolism in liver. At present, the production process of 25-OH-VD$_3$ mainly includes chemical synthesis and light irradiation. The chemical reaction steps are cumbersome, some of them need halogen reagents, and the racemates are generated during the reaction, which makes the separation difficult. Therefore, more and more researchers pay attention to the fermentation of 25-OH-VD$_3$ by microorganisms. The strains used in microbial biosynthesis mainly include Rhodococcus, Streptomyces, Pseudomocadia sp., and Mycobacterium. Vitamin D$_3$ hydroxylase (Vdh) is a kind of cytochrome P450 monooxygenase, which can catalyze the two-step hydroxylation of vitamin D$_3$ (VD$_3$) to produce 25-OH-VD$_3$ and 1α,25-dihydroxyvitamin D$_3$. Yasutake et al. (2013) used nisin, a natural bioactive antimicrobial peptide, to treat Rhodococcus cells containing hydroxylase, and they found that 573 mg/L 25-OH-VD$_3$ can be synthesized. Although the current industrial production of vitamin D$_3$ is mainly dominated by chemical synthesis, microbial synthesis methods are more sustainable and do not produce impurities during the biosynthesis process, thus it will be taken priority in the future industrial production.

Vitamin E
Vitamin E is a group of lipid-soluble antioxidants, including tocopherols and tocotrienols (Muñoz and Munné-Bosch, 2019; Zeng Z. et al., 2020). These compounds are composed of an oxygen-containing double ring system with a hydrophobic prenyl side chain (Blake and Konings, 2019). Lack of vitamin E affects the function of T and B immune cells (Moriguchi and Muraga, 2000). Additionally, patients with severe impairment due to Alzheimer's disease improved significantly after receiving α-tocopherol (Sano et al., 1997). Considering various physiological effects of tocopherols, they are widely used in the manufacture of human dietary supplements, food preservatives and cosmetics. There are four different tocopherol compounds, named α, β, γ, and δ tocopherol. Among the four forms of vitamin E, α-tocopherol is the most biologically active (Kaiser et al., 1990). In nature, α-tocopherol is produced by photosynthetic organisms, e.g., eukaryotic algae and green plants, some prokaryotic cyanobacteria, such as Synechocystis, which can accumulate vitamin E in large amounts (Figure 3B; Taketomi et al., 1983).

Recently, Euglena gracilis was found to be suitable for the production of high-value products, such as amino acids and ascorbic acid (Schwarzhans et al., 2015). E. gracilis is the most promising host for the commercial production of α-tocopherol, with a high growth rate and α-tocopherol content, which accounts for more than 97% of the total tocopherol accumulated by E. gracilis. Tani and Tsumura added precursors such as homogentisate and L-tyrosine to E. gracilis growth medium, which increased the accumulation of α-tocopherol to 143.60 mg/L corresponding to 5.1 mg/g dry cell weight (DCW) (Tani and Tsumura, 1989). Durmaz (2007) explored the effect of nitrogen source and concentration on the accumulation of α-tocopherol in Nannochloropsis oculata. When sodium nitrate and ammonium chloride were used as inorganic nitrogen source, the highest content of α-tocopherol reached 2.32 ± 0.04 mg/g dry weight (DW) (Table 2). The research showed that higher concentrations of nitrogen in the form of NO$_3$$^-$ and NH$_4$$^+$ can promote production the of α-tocopherol (Durmaz, 2007).

To balance cell growth and product synthesis, Shen et al. (2020) recently combined heterologous genes from photosynthetic organisms with the endogenous shikimate and mevalonate pathways (MEP) to construct a strain of S. cerevisiae that produces tocotrienols (Figure 3A). By incorporating a newly designed cold-shock-triggered temperature control system, the phased control of cell biomass and tocotrienol accumulation by the engineered strains was successfully realized. The final total tocotrienol titer reached 320 mg/L in a 5 L fermenter, which laid the foundation for the production of natural vitamin E in a fully fermentative process (Figure 3B; Shen et al., 2020).

In general, compared with chemical total synthesis, the method of obtaining vitamin E directly through biotechnology has low yield and high cost, and is not suitable for large-scale production. Although chemical total synthesis is currently the main production method of vitamin E, there are still many problems with this technology, such as complex synthesis routes, high technical barriers, etc. Therefore, the development of safer and more efficient synthesis technology has become the main problem to improve the current situation of vitamin E.

Vitamin K
Vitamin K is a fat-soluble vitamin, which also called blood coagulation vitamin in virtue of the function of promoting blood coagulation and preventing osteoporosis (Henrik, 1973; Schwalfenberg, 2017; Zhou et al., 2019). There are two naturally occurring types of vitamin K, called vitamin K1 (phyloquinone/phytomenadione) and vitamin K2 (menaquinone, MK) (Holvik et al., 2019). Vitamin K1 is synthesized by plants, while vitamin K2 is synthesized by microorganisms and can be divided into 14 isoforms depending on the number of isoprenoid units connected to the menaquinone ring (Figure 3E; Schwalfenberg, 2017). Among
FIGURE 3 | Metabolic network pathway of vitamin A/D/E and vitamin K. (A) MEP pathway. Dxs, 1-deoxy-D-xylulose-5-phosphate reductoisomerase; Dxr, 1-deoxy-D-xylulose-5-phosphate synthase; IspD, 2-C-methyl-D-erythritol 4-phosphate cytidylyltransferase; IspE, 4-diphosphocytidyl-2-C-methyl-D-erythritol kinase; IspF, 2-C-methyl-D-erythritol 2,4-cyclodiphosphate synthase; IspG, 1-hydroxy-2-methyl-2-(E)-butenyl 4-diphosphate reductase; IspH, 4-hydroxy-3-methylbut-2-enyl diphosphate reductase; IspI, isopenetyl diphosphate isomerase; IspA, geranyltranstransferase; CrtE, GGPP synthase; hpt, hypoxanthine phosphoribosyltransferase; γ-Tmt, γ-tocopherol methyl-transferase; Mf, methyl-transferase. (B) α-tocopherol biosynthesis pathway. Idi, isopentenyl diphosphate isomerase; IspA, geranyltranstransferase; CrtE, GGPP synthase; hpt, hypoxanthine phosphoribosyltransferase; γ-Tmt, γ-tocopherol methyl-transferase; Mt, methyl-transferase. (C) β-carotene biosynthesis pathway. CrtB, phytoene synthase; CrtI, phytoene desaturase; CrtY, lycopene cyclase. (D) MVA pathway. MvaS, HMG-CoA synthase; MvaA, HMG-CoA reductase; MvaK1, mevalonate kinase; MvaK2, phosphomevalonate kinase; MvaD, diphosphomevalonate decarboxylase; Menaquinone-n biosynthesis pathway. HepS/HepT, heptaprenyl diphosphate synthase component I/II; MenF, isocitrate synthase; MenD, 2-succinyl-5-enolpyruvyl-6-hydroxy-3-cyclohexene-1-carboxylate synthase; MenH, demethylmenaquinone methyltransferase; MenG, o-succinylbenzoate synthase; MenE, o-succinylbenzoate-CoA ligase; MenB, 1,4-dihydroxy-2-naphthoyl-CoA synthase; MenA, 1,4-dihydroxy-2-naphthoate heptaprenyltransferase; MenG, demethylmenaquinone methyltransferase. (F) Ergosterol biosynthesis pathway.

them, menaquinone-7 (MK-7) is the most effective subtype of vitamin K with a very long half-life in circulation. Notably, MK-7 can be synthesized in the cis, trans, and cis/trans forms, but only the all-trans form is biologically active (Szterk et al., 2018).

The biosynthesis of MK-7 from the embeden-meyerhof-parnas (EMP) pathway, pentose phosphate pathway (PPP), MVA pathway and menadione synthesis (MK) pathway (Figure 3C). A number of microorganisms have been used to produce MK-7, including B. subtilis, E. coli, lactic acid bacteria, Flavobacterium sp., and B. amyloliquefaciens (Sharma et al., 1993; Morishita et al., 1999; Sato et al., 2001; Wu, 2011; Taguchi et al., 2014).

Bacillus subtilis isolated from natto (a traditional Japanese food), the strain found in the eponymous fermented Japanese beans, has been certified by the FDA as a food-safe and has a strong ability to produce MK-7. Accordingly, B. subtilis natto was used as a parent strain to develop some of the industrial strains currently on the market. In industrial production, B. subtilis natto fermentation broth was sprayed and dried, and the dry powder from the fermentation broth was subjected to solvent extraction. The obtained extract was condensed into a paste and then purified by chromatography. Using this process, the final yield of MK-7 can reach 200–300 mg/L in the fermentation cycle of 16–24 h (Chen et al., 2016). Cui et al. (2019) developed a bifunctional quorum-sensing system in B. subtilis 168 to engineer the synthesis modules of MK-7. The resulting strain was capable of producing 360 mg/L MK-7 in shake flasks and 200 mg/L MK-7 in 15-L bioreactor (Cui et al., 2019). Recently, comparative transcriptomics revealed that cell membrane and electron transfer engineering in B. subtilis can improve the synthesis of MK-7. The resulting strain reached a product titer of 410 mg/L after 6 days in shake-flask culture, which is the highest value reported to date (Cui et al., 2020). In the current market environment, the production of natural all-trans MK-7 is via liquid fermentation of B. subtilis.
**TABLE 2** | Fat-soluble vitamins produced by biotechnological methods.

| Strain         | Biotechnological method                                                                 | Main culture substances             | Yield       | References               |
|----------------|----------------------------------------------------------------------------------------|-------------------------------------|-------------|--------------------------|
| **Vitamin A**  |                                                                                        |                                     |             |                          |
| *E. coli*      | Glycerol as the carbon source and harboring the whole MVA pathway.                      | 2YT medium; Glycerol                | 465 mg/L    | Yoon et al., 2009        |
| *E. coli*      | Overexpression of crt genes, dks, idi, sucAB, schABCD, and tailB.                       | LB medium                           | 2.1 g/L     | Zhao et al., 2013        |
| *Y. lipolytica*| Expressing the heterologous pathway and screen the best combination of promoters for each of the studied genes. | YPD medium; MM medium; YNB medium  | 6.5 g/L     | Larroute et al., 2018    |
| **Vitamin D**  |                                                                                        |                                     |             |                          |
| *S. cerevisiae*| DO was kept at 12% (±1%) and pulse fed-batch was used.                                  | MM medium                           | 1.16 g/L VD$_2$ | Tan et al., 2003         |
| *R. erythropolis* | Insert the gene-expression cassette encoding Bacillus megaterium glucose dehydrogenase-IV into the chromosome of *R. erythropolis*. | MM medium                           | 573 mg/L VD$_3$ | Yasutake et al., 2013    |
| **Vitamin E**  |                                                                                        |                                     |             |                          |
| *E. gracilis*  | Add effective additives (homogentisate and L-tyrosine); Optimize the concentration of ethanol and protein. | KH medium; Homogentisate; L-tyrosine | 5.10 mg/L   | Tani and Tsumura, 1989   |
| *E. gracilis*  | Determination of the amount of a-tocopherol produced under photoautotrophically, heterotrophically or photoheterotrophically. | MM; Methane                         | 8.60 ± 0.22 mg/L | Grimm et al., 2015      |
| *Stichococcus bacillaris* | Ballon bioreactor culture with MeJa as inducer.                                         | Methyl jasmonate (MeJa); Algal culture | 0.60 mg/g (DW) | Sivakumar et al., 2014 |
| *Nannochloropsis oculata* | Optimize the carbon source of the medium (NO$_3^+$-N and NH$_4^+$-N) and harvest time. | F/2 medium; Ammonium chloride (DW)  | 2.32 ± 0.04 mg/g | Duraz, 2007       |
| *S. cerevisiae*| Gene cloning from various photosynthetic organisms; Codon optimization and protein truncation. | SD medium                           | 320 mg/L    | Shen et al., 2020        |
| **Vitamin K (MK-4/MK-7)** |                                                                                         |                                     |             |                          |
| *B. subtilis natto* | Optimum media conditions and screening producing strain (Different nutrients of the culture medium will affect the yield of MK-7). | Glycerol                            | 62.32 ± 0.34 mg/L | Berenjian et al., 2011 |
| *B. subtilis natto* | Fermentation using soybean extract and screening highest MK7 yielding strain from commercially available natto. | Soy granules; Amylase               | 67.01 ± 0.18 mg/kg | Mahanama et al., 2011 |
| *B. subtilis*  | Deletion of PAS-A, kinB, spoIIA, spoIIE, dhbB, and ptsG; Overexpression of menF, menB, menE, entC, ppsA, arOK, ispA, hepS/T, kdpG, dxr, dxs, fri, menA. | LB Medium                           | 200 mg/L    | Cui et al., 2019         |
| *B. subtilis*  | Overexpression of BS20-qcrA-C and tatAD-CD.                                              | LB Medium                           | 410 mg/L    | Cui et al., 2020         |

narro, which is safe, natural and controllable, and occupies the mainstream position in the market. Compared with the chemically synthesized of trans-MK-7, it has a higher yield and fewer impurities.

**CONCLUSION**

The fermentative production of vitamins using bacteria, yeasts or microalgae has many advantages over traditional chemical synthesis methods. From the aspects of safety, biological activity, absorption rate, etc., vitamins manufactured by biological methods can be more suitable for both internal and external applications (Yuan et al., 2020). Although the fermentation of VB$_2$ and VB$_12$ has technologically matured and is being applied in industrial production, fermentation methods for the remaining B-group vitamins have yet to be developed or require significant yield improvement.

Vitamin C has a large market, and its production method is mainly based on single-bacteria fermentation, which eliminates the dependency of associated bacteria by replacing accompanying bacteria with associated active agents (Vandamme and Revuelta, 2016a). However, the current market situation indicates that vitamin C production has overcapacity, the downstream processing is complicated, and the market demand is concentrated in the field of medicine and food. For these reasons, the momentum of price increase will remain slow in the future. Vitamin C fermentation technology can explore the mechanism of a variety of accompanying bacteria, establish their anabolism database, or use isotope technology to label and trace the individual metabolites. It is also possible to design heterologous assembly modules for 2-KGA synthesis, and study adaptation mechanisms in microbial chassis cells, so as to achieve higher productivity (Liu et al., 2011).

The current biosynthesis product of vitamin A is mainly focused on β-carotene. The biosynthesis of β-carotene...
has successfully established a large-scale production process through classical and reasonable microbial metabolic engineering. However, due to the high barriers of intermediate industry and the complex process of synthesis and metabolism, the future research will face more challenges. At present, the industrial production of vitamin D is mainly through the chemical synthesis of active 25-OH-VD\(_3\) and 1α,25-dihydroxyvitamin D\(_3\), but the biggest obstacle is the assured quality and security of supply for raw materials, which must be cholesterol with purity greater than 95% (NF grade). Therefore, the key to solving the problem of raw materials is to develop more production bacteria, optimize their metabolic pathways and make them highly productive. With the continuous optimization and technological progress of vitamin E fermentation, the overall cost of the industry has fallen, which promoted the growth of the industry. Unfortunately, although studies have shown that photosynthetic microorganisms have considerable potential for the production of tocopherols, light-driven fermentation is costly, which makes commercialization difficult. However, due to the considerable potential \(E.\) gracilis and the conditions of the cultivation environment, the construction of a specifically designed photobioreactor may be a feasible research direction for the production of tocopherol. Moreover, the controllable temperature-sensitive control system may also be a key control technology for vitamin E production. Among vitamin K producing bacteria, \(B.\) subtilis natto seems to be the most promising candidate for MK production. Many researchers have optimized the design of fermentation modes, medium components, and culture conditions. They have also applied genetic engineering and other means to increase MK production (Szerk et al., 2018). However, to achieve higher industrial output, the technology needs to be further improved. Some studies have used biofilm reactors, which may become a promising new area for future research.

Recently, our research group used \(E.\) coli MG1655 (DE3) as chassis strains and achieved the \textit{de novo} synthesis of vitamin B\(_{12}\) via metabolic engineering and optimization of fermentation conditions. In addition, we have not only proved that \(E.\) coli is a microbial biosynthesis platform for the production of vitamin B\(_{12}\), also provides an encouraging example of how the dozens of proteins in a complex biosynthetic pathway can be transferred between organisms to promote industrial production (Fang et al., 2018). In addition, our research group is also doing metabolism research on vitamin B\(_2\), B\(_6\), B\(_7\), and vitamin K.

In general, the development of synthetic biotechnology provides new opportunities for the construction of vitamin cell factories. First, high-throughput screening of high-yield strains, the CRISPR/Cas9 genome editing technology, and automatic gene assembly technology provide important technical means for the mining and genetic modification of chassis cells (Chang et al., 2019; Zeng W. et al., 2020; Zhang and Showalter, 2020). Further, the output of vitamin products in different dimensions will be increased by transforming the complex and multi-enzyme pathways required for the production of vitamins, establishing microbial flora with controllable functions and stability, and application of some advanced engineering technology, such as the cold-shock-triggered temperature control system, dynamic control of gene expression systems, different types of biosensors, cell-free systems and computer-aided design, etc. (Koo et al., 2020; Marucci et al., 2020; Sachsenhauser et al., 2020; Shen et al., 2020; Glasscock et al., 2021). Additionally, the modular and orthogonal strategies are increasingly supporting the construction of vitamin cell factories (Liu et al., 2015). The mining and design of biological components, the assembly and integration of elements and modules, and the optimization and adaptation of the fermentation system are also important for efficient production of vitamins (Santos-Merino et al., 2019). However, there will be many challenges in the field of synthetic biotechnology in the future, including the compatibility between flexible biological systems and rigid engineering systems, or the universality of biological system reconstruction. It will be necessary and important to advance the existing technology, combine it with new strategies, and conduct interdisciplinary research to establish novel microbial cell factories for the industrial fermentation of most vitamins. All in all, we firmly believe that the industrialization of the fermentation production of vitamins is expected to become a broader, safer and more sustainable manufacturing with the continuous advancement of synthetic biotechnology and metabolic engineering.

**AUTHOR CONTRIBUTIONS**

YW and LI: manuscript planning, writing, and revision. ZJ and DZ: manuscript revision and writing. All authors contributed to the article and approved the submitted version.

**FUNDING**

This study was funded by the National Key R&D Program of China (no. 2019YFA0905300), the National Natural Science Foundation of China under grants nos. 31670604 and 31970324, the Natural Science Foundation of Liaoning Province of China under grant no. 2019020758, the Science and Technology Project of Liaoning Education Department under grant no. 819001110761, and the Tianjin Synthetic Biotechnology Innovation Capacity Improvement Project (no. TSBCIP-CXRC-004).

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.