Enzymatic synthesis of fluorinated compounds

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Abstract
Fluorinated compounds are widely used in the fields of molecular imaging, pharmaceuticals, and materials. Fluorinated natural products in nature are rare, and the introduction of fluorine atoms into organic compound molecules can give these compounds new functions and make them have better performance. Therefore, the synthesis of fluorides has attracted more and more attention from biologists and chemists. Even so, achieving selective fluorination is still a huge challenge under mild conditions. In this review, the research progress of enzymatic synthesis of fluorinated compounds is summarized since 2015, including cytochrome P450 enzymes, aldolases, fluoroacetyl coenzyme A thioesterases, lipases, transaminases, reductive aminases, purine nucleoside phosphorylases, polyketide synthases, fluoroacetate dehalogenases, tyrosine phenol-lyases, glycosidases, fluorinases, and multienzyme system. Of all enzyme-catalyzed synthesis methods, the direct formation of the C-F bond by fluorinase is the most effective and promising method. The structure and catalytic mechanism of fluorinase are introduced to understand fluorobiochemistry. Furthermore, the distribution, applications, and future development trends of fluorinated compounds are also outlined. Hopefully, this review will help researchers to understand the significance of enzymatic methods for the synthesis of fluorinated compounds and find or create excellent fluoride synthase in future research.

Key points
• Fluorinated compounds are distributed in plants and microorganisms, and are used in imaging, medicine, materials science.
• Enzyme catalysis is essential for the synthesis of fluorinated compounds.
• The loop structure of fluorinase is the key to forming the C-F bond.

Keywords Enzymatic synthesis · Fluorinase · C-F bonds · Fluorinated compounds · Applications of fluorides

Introduction
Fluorine has different physical and chemical properties from other elements, including bond length, van der Waals radius, van der Waals volume of the atom, and electron-egativity of elements (Meanwell 2018). Unlike other halogens (chlorine, bromine, iodine, astatine, and tennessine), non-polarizable fluorine does not participate in halogen bonding. Due to its non-polarizability, strong electrostatic interactions will be formed, resulting in attractiveness or repulsion (Yang et al. 2015). Moreover, fluorine can change the pKₐ, affinity, dipole moment, stability, lipophilicity, and bioavailability of adjacent groups (Dos Santos et al. 2020; Khosravan et al. 2017; Muller et al. 2007). Therefore, the target compounds doped with fluorine are endowed with stronger activity and stability, longer half-life, and better bioabsorbability (Moschner et al. 2019; Ni and Hu 2016; Reichel and Karaghiosoff 2020), especially in the fields of pharmaceutical intermediates (Gillis et al. 2015; Swallow 2015; Zhdankin et al. 2017; Zou et al. 2019), cancer treatment (Meanwell et al. 2020), antiviral agents (Pomeisl et al. 2019), photovoltaics, diagnostic probes and bioinspired materials (Moschner et al. 2019). Besides, natural proteins incorporating fluorinated amino acids showed also many unique characteristics, which are used in the biotherapeutics
protein–protein interaction and the synthesis of high value-added chemicals (Arias et al. 2020; Awad and Ayoup 2020; Bucci et al. 2019; Liu et al. 2019; Mei et al. 2020a; Remete et al. 2018; Sisila et al. 2021; Vaughan et al. 2016; Wang and Matthews 2020; Won et al. 2019).

Unfortunately, fluorinated natural products are extremely rare in nature, which limits the application of fluorine-containing compounds. Accordingly, the artificial synthesis of fluorinated compounds has become an important alternative in modern society. However, the formation of C-F bonds through nucleophilic fluorination reactions has been progressing slowly. The main reason is that fluorinated compounds react poorly and fluoride ion is easily solvated (ΔG° > 450 kJ/mol) in protic media (Liang et al. 2017; Markakis et al. 2020; Mennie et al. 2018; Scheidt et al. 2018). Hence, they have the strongest hydration of halide so that the huge kinetic inertia needs to be overcome to promote this reaction. To realize the artificial synthesis of fluorinated compounds, many chemical catalysis methods have been developed and used to synthesize various fluorine-containing compounds including [18F]-6-fluoro-3,4-dihydroxy-phenylalanine (Mossine et al. 2020), 18F-fluoro-α-methyl tyrosine, 18F-prostate-specific membrane antigen (PSMA)-1007 (Giesel et al. 2017), and 18F-fluoro-α-methyl phenylalanine (18F-FAMP) (Hanaoka et al. 2019; Yamaguchi et al. 2020).

Nonetheless, these conventional chemical synthesis methods required precious metals, toxic and contaminating chemical reagents, high temperature, high pressure, extreme conditions, and the protection and deprotection of groups. This is not conducive to the sustainable development of green chemicals and the global economy. In contrast, the biosynthesis methods are good partners/supplements/alternatives in the field of chemical synthesis, especially for fluorinated compounds. The biosynthesis methods are divided into fermentation methods and enzyme-based biocatalysis methods. Through fermentation methods, various fluorinated compounds can be synthesized by engineered microorganisms that integrated substrate transporters, product degradation pathways, fluoride ion channels, fluoride-responsive riboswitch, orthogonal T7 RNA polymerase, or chromosome (Calero et al. 2020; Jiang et al. 2018; Markakis et al. 2020; Rivera-Chavez et al. 2017; Sester et al. 2020; Thurowy et al. 2017). However, some factors limit its application, including complex regulatory networks and the toxic effects of fluoride on living microorganisms. For enzyme-based biocatalysis, it has the advantages of energy-saving, environmental protection, safety, high selectivity, atomic economy, convenient operation, and mild reaction conditions (Council et al. 2020; Fryszowska and Devine 2020; Hauer 2020; Honig et al. 2017; Latham et al. 2018; Menon et al. 2020; Minges and Sewald 2020; O’Hagan and Deng 2015; Odar et al. 2015; Sandoval and Hyster 2020; Tong et al. 2019; Wu et al. 2020a). Hence, it has attracted more and more attention from biologists and chemists. In this review, diverse methods are introduced for the synthesis of fluorinated compounds catalyzed by different enzymes in recent years, especially fluorinase that can form the C-F bond with the selective catalytic mechanism (Fig. 1). Combined with the distribution, enzymatic synthesis methods, and applications of fluorinated compounds, this review will attract more scientists to participate in the field of fluorobiochemistry.

### Distribution of fluorinated natural products

Since the discovery of the first fluorinated natural product, it has been more than 70 years. Fluorinated natural products are only distributed in fewer tissues. Although fluorides have also been found in animals, there is no direct evidence that fluorinated natural products can be synthesized by animals (Harper and O’Hagan 1994). Fluorides are obtained by animals mainly by eating foods containing fluorinated natural products. Plants and microorganisms have developed a huge ability to biosynthesize secondary metabolites, some of which are fluorinated natural products of various structures. However, there are still very few fluorinated natural products found in nature. The main known fluorinated natural products in plants and microorganisms are as follows (Fig. 2).

### Fluorinated natural products in plants

Fluoroacetate was first discovered from plant gifblaar (Dichapetalum cymosum) in 1943 (Toit 1943). It is also the most common fluorinated natural product and has been identified in more than 40 higher plants. These plants that produce fluoroacetate mainly come from the families of Malpighiaceae, Fabaceae, Dichapetalaceae, Rubiaceae, and Bignoniaceae (Grobbeelaar and Marion Meyer 1990; Lee et al. 2014). Fluoroacetate-containing plants grow all over the world, especially in Africa, Australia, and South America (Leong et al. 2017). Due to the toxicity of fluoroacetate, it is generally believed that these plants produce fluoroacetate for the defense of herbivores (Leong et al. 2017). Especially Dichapetalum braunii, its seeds contain up to 8000 ppm of fluoroacetate, which is extremely defensive and even lethal (O’Hagan and B. Harper 1999). In addition, (2R,3R)-2-fluorocitrate was also found to be present in the West African shrub Dichapetalum toxicarium and forage plants (soya bean, alfalfa, and crested wheatgrass) (Carvalho and Oliveira 2017; Harper and O’Hagan 1994; O’Hagan and Deng 2015; Ren et al. 2020; Zhang et al. 2019c). Despite the great interest, the physiological role of fluorinated natural products is not fully understood in higher plants. If fluoroacetate is for self-defense, are other natural products...
Fig. 1 Enzymatic synthesis of fluorinated compounds

Fig. 2 Known fluorinated natural products
of fluorination involved in the synthesis of metabolites or proteins? There are still many areas to be explored.

**Fluorinated natural products in microorganisms**

Fluoroacetate is also produced by various microorganisms, including *S. castellary*, *Streptomyces* sp. MA37, *Norcardia brasiliensis*, *Actinoplanes* sp. N902-109, *Actinopolyspora mzabensis*, and *Streptomyces xinghaiensis* NRRL B24674 (Deng et al. 2014; Huang et al. 2014; Ren et al. 2020; Sanada et al. 1986; Sooklal et al. 2020). In addition to fluoroacetate, fluorinated threonine is another fluorinated natural product derived from fluoride-producing microorganisms. Moreover, it is also the only fluorinated amino acid found in nature. It should be noted that a new type of flurometabolite (2R3S4S)-5-fluoro-2,3,4-trihydroxypentanoic acid) was identified in *Streptomyces* sp. MA37 except for fluoroacetate and fluorinated threonine (Ma et al. 2015). This also showed that there is a unique fluoride metabolism pathway in *Streptomyces* sp. MA37. In addition to the above fluorinated natural products, microorganisms also produce other fluorinated natural products mainly including 4'-fluoro-3'-O-p-glucosylated metabolites (F-Mets I and II) (Feng et al. 2019), 4-fluoronucleosides (Zhang et al. 2019c), fluorosalinoseporamide (O’Hagan and Deng 2015), and 5-fluoro-5-deoxy-D-ribose (5-FDRu) (Wu et al. 2020b). Like plants, the physiological role and mechanism of fluorinated natural products are still a mystery in microorganisms.

**Enzymatic synthesis of fluorinated compounds**

**Cytochrome P450 enzymes**

Cytochrome P450 enzymes (P450s) belong to the superfamily of heme monoxygenases and are involved in the synthesis and degradation of many metabolites in actinomycetes. P450s can catalyze a variety of different reactions, mainly including dehydrogenation, epoxidation, dealkylation, C=C bond formation or cleavage, C-N bond formation, and hydroxylation (Chen and Arnold 2020; Iizaka et al. 2021). Therefore, various fluorinated compounds can be synthesized through different types of reactions mediated by P450s. This is an effective strategy to incorporate fluoroalkyl groups through the conversion of C(sp³)-H bonds to fluorocarbon alkyl bonds (McAtee et al. 2018). To synthesize organic fluorocarbon alkanes with α-CF₃, a biocatalytic platform was developed based on cytochrome P450 (Fig. 3A) (Huang et al. 2019). Through this platform, diverse α-CF₃ organo-oborons were obtained from 400 mM Lewis-base borane complexes and trifluorodiazol alkanes (ee = 85–99%). Similarly, various organofluorine compounds with S-stereoselectivity were also obtained from 10 mM N-aryl pyrrolidine and 20 mM 2,2,2-trifluoro-L-diazoethane (ee > 99%) (Zhang et al. 2019b). Notably, engineered P450 completely reversed the enantioselectivity of the products through directed evolution (R-stereoselectivity, ee > 99%). Moreover, fluorinated compounds can also be synthesized by P450 through nitration reaction. Aromatic nitration is an important industrial process for the production of many chemicals. Cytochrome P450 enzyme from *Streptomyces scabies* 87.22 TxtE is the only enzyme that can directly catalyze the nitrification reaction so far (Zuo et al. 2016). To expand its application potential, fluorinated nitro-tryptophan analogs were obtained by P450TxtE from 1.5 mM 4-F-β-L-tryptophan and 5-F-β-L-tryptophan in a 10 mL reaction system.

Among all the reaction types catalyzed by P450, the hydroxylation reaction is the most important type of reaction. Thus, the fluorinated derivative of N-(1-adamantyl)-1-pentyl-1H-indazole-3-carboxamide (5F-AKB48) was obtained in vitro hydroxylation reaction (Pinson et al. 2020). Moreover, the 5F-AKB48 is the main component of cannabinoids (SCs), and the experiments showed that P450s from humans can oxidize 5F-AKB48 to fluoride with different hydroxyl groups. Interestingly, fluorine-containing compounds are also activators in the P450-mediated hydroxylation process. With the assistance of perfluorinated decoy molecules (PFC), the hydroxylation rate was significantly enhanced for unnatural substrate cycloalkanes by P450 from *Bacillus megaterium* (Dezvarei et al. 2018). In addition to bioenzymatic methods, P450 can also realize the biosynthesis of fluorides through chemoenzymatic methods. For common organic molecules, one or more non-activated sites were selectively fluorinated by P450 and nucleophilic fluorination reagent diethylamino-sulfur trifluoride (DAST) (Figure S1) (Obach et al. 2016; Rentmeister et al. 2009). In this process, monofluorine and polyfluorine compounds were synthesized through a two-step chemical enzymatic method. Especially for cyclic fluorides (steroidal analogs), high regioselectivity (55–100%) and yields (> 60%) could be achieved by various P450 mutants and DAST. It is just that the use of chemical reagents makes the entire catalytic process less economical and environmentally friendly. It is worth noting that the ability of P450s to synthesize fluorides is also their physiological function in some organisms. Such as 6:2 fluorotelomer sulfonic acid (6:2 FTSA) is widely used in water-based film-forming foam as a new type of perfluorocane sulfonic acid. In earthworms (*Eisenia fetida*), studies have shown that P450 is involved in the biotransformation of 6:2 FTSA (Zhao et al. 2021). Similarly, this function is also proven in pumpkin (*Cucurbita maxima* L.) (Zhao et al. 2019a).
As such, various fluorinated compounds can be obtained through diverse reactions mediated by P450. However, it is still necessary to further improve the issues of thermal stability, regional and stereo specificity of P450. Moreover, P450 must co-express the electron transport chain to perform its biological functions. These disadvantages are being overcome by experts in the field of synthetic biology. Synthetic biology is a cross-discipline that combines biology and engineering. It aims to achieve the goal of de novo synthesis of organisms through genetic design and construction of new biological elements and artificial gene circuits. In the future, synthetic biology is expected to combine interdisciplinary disciplines to use P450 for the de novo synthesis of fluorinated compounds.

**Purine nucleoside phosphorylases**

Purine nucleoside phosphorylase (E.C. 2.4.2.1, PNP) is a well-known drug target including cancer and immunodeficiency diseases (Pant et al. 2021; Parker et al. 2020). PNP can reversibly catalyze phosphorolysis of purine nucleosides to form glycosidic bonds (Holanda et al. 2020; Stachelska-Wierzchowska and Wierzchowski 2020; Timofeev et al. 2020). Therefore, PNP is a significant catalyst in the metabolic pathway of purines and nucleosides, especially 2-fluoroadenosine and 2-fluorocordycepin (Kayushin et al. 2021). To obtain fluorinated nucleoside analogs with antiviral activity, a chemo-enzymatic method was developed based on PNP (Pi = inorganic phosphate; C) The biosynthetic of 3-fluoro-L-tyrosine catalyzed by TPL.

PNP can reversibly catalyze phosphorolysis of purine nucleosides to form glycosidic bonds (Holanda et al. 2020; Stuchelska-Wierzchowska and Wierzchowski 2020; Timofeev et al. 2020). Therefore, PNP is a significant catalyst in the metabolic pathway of purines and nucleosides, especially 2-fluoroadenosine and 2-fluorocordycepin (Kayushin et al. 2021). To obtain fluorinated nucleoside analogs with antiviral activity, a chemo-enzymatic method was developed based on PNP (Fig. 3B) (Kharitonova et al. 2016). In this process, four fluorides were synthesized from 5 mM 1-α-phospho-2-deoxy-2-fluoroarabinose and 1 mM fluorobenzimidazole bases (Conversions of bases = 46–88%; Yields = 40–55%). This provides an effective route for the synthesis of fluorinated nucleoside analogs. Although there is no need for any coenzymes or cofactors to participate, the use of chemical reagents increases the pollution of the reaction. This does not meet the requirements of the green economy and will increase the pressure on subsequent environmental protection.
treatments. Accordingly, an important challenge of this route is to develop new biocatalysts without chemical reagents.

**Tyrosine phenol-lyases**

PLP-dependent tyrosine phenol-lyases (EC 4.1.99.2, TPLs) can reversibly synthesize \( l \)-tyrosine from phenol, pyruvate, and ammonia (Phillips et al. 2019; Rocha et al. 2019). Thus, a TPL-based process has been developed to obtain the fluorinated \( l \)-tyrosine analogs, a potential drug for the treatment of Parkinson’s disease (Zhu et al. 2020a). Through the process, \( o \)-fluorophenol was transformed into 3-fluoro-\( l \)-tyrosine by TPL and the space–time yield reached 4.2 g/L/h in fed-batch fermentation (Conversion = 90.2%; Yield = 95.1%) (Fig. 3C). Similarly, other fluorinated \( l \)-tyrosine derivatives were also synthesized through TPL-catalyzed synthetic biological systems (Dennig et al. 2015; Li et al. 2020a; Won et al. 2019). Moreover, the donor substrates were generated in situ. In this process, \( l \)-lactate oxidase (LOX) and P 450BM3 were usually used to generate pyruvate and fluorinated phenol from \( l \)-lactate and fluorinated aromatic hydrocarbon, respectively. As such, this method is very promising for the large-scale production of fluorinated \( l \)-tyrosine and its derivatives. However, this biocatalytic method is also an important measure to upgrade cheap bulk chemicals to high value-added chiral fluorides. Unfortunately, the low catalytic efficiency requires more machine learning-assisted directed evolution to improve.

**Aldolases**

\( \beta \)-Hydroxy-\( \alpha \)-amino acids and their fluorinated derivatives are key intermediates for many drugs, which were used for the treatment of Parkinson’s disease immunodeficiency and inflammation (Rocha et al. 2019; Scott et al. 2017). Pyridoxal 5’-phosphate (PLP)-dependent aldolases are regarded as the key biocatalysts for the formation of C–C bonds and have become powerful tools for the sustainable synthesis of complex molecules, especially fluorinated \( \beta \)-hydroxy-\( \alpha \)-amino acids through aldol condensation (Chen et al. 2019a; Fang et al. 2019; Zheng et al. 2020, 2021). The formation of two chiral stereocenters is the main feature of this reaction. Moreover, it is difficult to control the stereoselectivity of the two chiral centers at the same time. The introduction of the special fluorine element further increases this difficulty, so that the first stereoselective fluorine-aldol condensation reaction was established in 2016 (Saadi and Wennemers 2016). Subsequently, fluorine-substituted \( \beta \)-hydroxy-\( \alpha \)-amino acids are synthesized by \( d/l \)-threonine aldolases (\( d/l \)-TAs) (Chen et al. 2017, 2019a). In these cases, fluorine is not in the chiral center but the side chain. Various fluorinated \( d/l \)-threoleurythro-\( \beta \)-hydroxy-\( \alpha \)-amino acids were obtained from 5–100 mM aromatic fluorine substituted aldehydes and 10 equivalents of glycine (Conversions of aldehydes = 29.8–92.8%). Similarly, fluorinated \( l \)-threo-threonine was also biosynthesized by 4-fluorothreonine transaldolase (FTase) from \( l \)-threo-threonine and fluoroacetaldehyde (Figure S2) (Wu et al. 2020c). Besides, chiral organofluorines have also been synthesized through a chemo-enzymatic method based on type II pyruvate aldolase (Fang et al. 2019). The donor binding and enolization can only be activated by divalent metal cations. In this process, 50 mM fluoropyruvate and twelve aldehydes were synthesized corresponding to \( \beta \)-fluoro-\( \alpha \)-hydroxy compounds by pyruvate aldolase, 1 mM MgCl\(_2\), and H\(_2\)O\(_2\) (Conversions of polar aldehydes = 80–100%; Conversions of nonpolar aldehydes = 10–70%) (Figure S3). In all reactions catalyzed by

\[ \text{Fluorinated phenylacetaldehyde} \]

\[ \text{Fluorinated } l\text{-threo/erythro } \beta\text{-hydroxy-} \alpha\text{-amino acids} \]

\[ \text{Fluorinated } d\text{-threo/erythro } \beta\text{-hydroxy-} \alpha\text{-amino acids} \]

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**Fig. 4** The fluorinated compounds with hydroxyl group were obtained. **A)** Synthesis of fluorinated \( \beta \)-hydroxy-\( \alpha \)-amino acids employing threonine aldolase; **B)** Biosynthesis of fluoroacetate catalyzed by FLK.
aldolases, control of regio and stereoselectivity is the key to the reaction. This is also one of the future development directions of aldolases.

**Fluoracetyl coenzyme A thioesterases**

Fluoracetyl coenzyme A thioesterases (EC 3.1.2.29, FLKs) can catalyze the decomposition of fluorocetyl coenzyme A (fluoroacetyl-CoA) into fluoroacetate and CoA, and does not catalyze acetyl-CoA (Fig. 4B) (Huang et al. 2006; Thuronyi and Chang 2015). FLK has extremely high specificity for fluorinated substrates, and its hydrolysis efficiency is 10,000-fold that of non-fluorinated substrates (Weeks and Chang 2012). To understand the mechanism of selective fluorination, a comprehensive strategy was implemented for FLK including thermodynamic, kinetic, and biophysical experiments. These studies indicated that entropy drive mediated by hydrophobicity and solvation is the key to selective fluorination recognition, especially the substrate-binding controlled by the active site Phe36 (Figure S4) (Weeks et al. 2018). Moreover, the selectivity to the fluoracetyl-CoA substrate not only depends on the polarization provided by the negatively charged fluorine substitution but also on the molecular recognition of fluorine during the formation of acyl-FLK intermediate (Weeks et al. 2014). It should be noted that FLK itself also has important physiological functions. FLK can prevent the further metabolism of fluoroacetyl-CoA to 4-hydroxy-trans-arachidic acid ester, a lethal inhibitor of the tricarboxylic acid cycle (Dias et al. 2010). The specific catalytic activity of FLK has great advantages in physiology. In addition, it also has extremely high selectivity for the synthesis of specific fluorinated products. However, more directed evolution engineering should be carried out to obtain fluorinated acetic acid analogs with diverse structures requires. Moreover, the rational use of CoASH is also an issue worth considering in the process of biosynthesis.

**Transaminases**

PLP-dependent transaminases are important biocatalysts for the synthesis of chiral amines (Patil et al. 2018). For fluorinated chiral amines, an asymmetric amination method was established based on transaminase (Fig. 5A) (López-Iglesias et al. 2017). R-fluoroamines or S-fluoroamines were obtained from 50 mM 1-(5/6-fluoropyridin-3-yl) ethan-1-one or 1-(6-(trifluoromethyl)pyridin-3-yl)ethan-1-one by R/S-selective transaminase (Conversions = 75–94%; eeR/S > 99%). d/l-Alanine (250 mM) was used as amino donors for transaminase. Similarly, methyl/trifluoromethyl diketone was also catalyzed by transaminase to fluoroamines (Figure S5) (González-Martínez et al. 2020b). Although the amination of two carbonyl groups will form three products in theory, transaminase only selectively modified smaller methyl ketones in practice. Consequently, the product 1-(4-(1-aminoethyl)phenyl)-2,2,2-trifluoroethan-1-one was obtained by amination of a carbonyl group (ee = 98- > 99%).

To further produce chiral fluorinated hydroxylamine, alcohol dehydrogenase was subjected to a hydrogenation reaction to obtain the substrate of hydroxyketones. The hydroxyketones obtained were then converted chiral fluorinated hydroxylamines by transaminase (conversions = 84–90%) (Figure S6). Since the reaction catalyzed by transaminase is reversible, the kinetic resolution reaction can be achieved to obtain fluorooamines from the corresponding racemate. Interestingly, the kinetic resolution reaction catalyzed by transaminase is promiscuous with hydrodefluorination and deamination (Cuetos et al. 2016). In the unconventional process mediated by catalytic promiscuity, a series of aromatic β-fluoroamines were obtained by kinetic resolution through transaminase-mediated hydrodefluorination/deamination reactions (Yields = 75–80%, eeS > 99%) (Figure S7).

Notably, this protocol is the first case of using transaminase to catalyze promiscuousness and does not require amino acceptors. Even so, transaminases still require optically pure d/l-amino acids as amino donors. Expensive amino
donors increase the cost of biocatalysis and bioconversion. The converted amino donors also increase the difficulty of separating the final products. Moreover, the limited recycling of the cofactor PLP also restricts the further promotion of this method.

**Reductive aminases**

Chiral fluoroamines are the key components of drug molecules, which can improve efficacy and slow down metabolism. In addition to transaminases, chiral fluoroamines can also be synthesized by reductive aminases, which belong to the imine reductase family (France et al. 2018). Reductive aminase showed high activity for the reductive amination of ketones for the synthesis of fluorinated amines (Aleku et al. 2017). Similar to transaminase, reductive aminase also exhibited a promiscuous reaction to reduce ketones to amines and alcohols (Fig. 5B) (González-Martínez et al. 2020a; Yang et al. 2021). Through reductive aminase, primary and secondary β-fluoroarylamines and fluoroarylalcohols were obtained from 5 mM fluoroarylketones (Conversions > 90%, ee = 85–99%). As such, reductive aminase provides a sustainable and highly economical biocatalysis toolbox, which can synthesize fluorinated chiral amines and fluorinated chiral alcohols from readily available prochiral substrates. Moreover, inexpensive inorganic ammonium salts provide a lot of convenience as amino donors and do not produce by-products. Nonetheless, the high cost of the coenzyme NADPH is a major disadvantage. Therefore, various coenzyme circulation systems are usually integrated through glucose dehydrogenase or formate dehydrogenase. But it will undoubtedly be more perfect if the coenzyme preference of expensive NADPH is changed to cheap NADH.

**Polyketide synthases**

Polyketides are a class of complex natural products with multiple structures and biological activities, which are prolific sources of numerous different scaffolds (Maglangit et al. 2019; Tao et al. 2021). Polyketide synthases (PKSs) are multidomain megasynthetases and participate in the synthesis of numerous polyketide natural products in bacteria and fungi (Adrover-Castellano et al. 2021; Weissman 2016; Zhu et al. 2021). The introduction of fluorine-containing modules into the PKS system can bring fluorine into the scaffold of natural products, which is an important strategy for the synthesis of complex organic fluorides. Here, acetate is the basic element for the formation of polyketide natural products. Hence, the preparation of fluorinated polyketide has been investigated.

Fig. 6  The complex fluorinated compounds were obtained from fluoroacetate as the building block
PKS has three necessary domains, including the ketone synthase (KS) domain, acyltransferase (AT) domain, and acyl carrier protein (ACP) domain (Robbins et al. 2016; Wang et al. 2021). The catalytic process showed that the inactivation of the AT domain can eliminate the selectivity to the fluorine-containing extension unit in the fluorinated polyketones biosynthesis system and initiate the C–C bond formation mode independent of ACP (Ad et al. 2017). Subsequently, the trans-AT complementary domain was incorporated to synthesize both monofluorinated and difluorinated products. Consequently, the use of the PKS module is expected to achieve a stable production method of fluorinated full-length polyketide and other complex products at a specific location. Due to the complexity of the structure of fluorinated polyketones, the selective synthesis mechanism of PKS remains to be elucidated. Therefore, this is a huge challenge for the selective synthesis of various fluorinated substituted polyketones. To explore the catalytic mechanism, more work needs to be done in the field of crystal structure analysis and mutation in the future.

Lipases

Lipases (EC 3.1.1.3) belong to the family of hydrolases. In the presence of organic solvents (such as stearic acids, triacylglycerides, and fatty acids), many substrates can be effectively catalyzed by lipases for acylation (Cavalcante et al. 2021; Huang et al. 2021; Marquez-Rodriguez et al. 2021). Besides, lipases are also excellent biocatalysts to obtain fluorinated compounds in the presence of organic solvents. For fluorinated 1-(pyridin-3-yl)ethanamines, ethyl methoxyacetate, and tetrahydrofuran were used as acylating agents and cosolvent, respectively (López-Iglesias et al. 2017). In this reaction, 100 mM fluorinated racemic amines were converted to corresponding R-isomers and S-isomers through kinetic resolution (Conversions = 50%; ee\textsubscript{S} = 90–96%; ee\textsubscript{R} = 99%). Similar to kinetic resolution, lipases can also catalyze enantioselective hydrolysis reactions to obtain β-fluorinated amino acids, which are key components of many natural products, drugs, and biologically active peptides (Shahmohammadi et al. 2020; Zhang et al. 2020b). In the presence of \textit{iPr}_2\text{O}, \textit{Et}_3\text{N}, and H\textsubscript{2}O, 25 mM racemic fluorinated β-amino carboxylic ester hydrochloride salts were hydrolyzed into fluorinated \textit{R}-β-amino carboxylic ester enantiomers (Yields = 38–49%, ee\textsubscript{R} > 99%) and fluorinated \textit{S}-β-amino acids (Yields = 48–49%, ee > 99%) (Fig. 7A). Likewise, many other types of fluorinated compounds were also obtained by lipases, like 2,2,2-trifluoroethyl trifluoromethyl sulfate, fluorinated polyesters, and \textit{meso}-2,4,6-trifluoro-1,3,5,7-tetrahydroxyheptane-1,7-diacetate (Bentler et al. 2019; Snoch et al. 2019; Zhao et al. 2020).

![Fig. 7](image_url) The fluorinated compounds were obtained by kinetic resolution. A) Fluorinated β-amino acids/carboxylic esters were obtained by lipase; B) 2-Fluorocarboxylic acid derivatives were obtained by fluoroacetate dehalogenase.
Importantly, lipase has good resistance to organic solvents in addition to being able to catalyze cheap ester substrates. Thus, lipase is an excellent choice for catalytic reactions that require organic solvents to help dissolve substrates. Notably, the pollution of organic solvents to the environment must be carefully considered in this process. The organic solvent recycling device may become an alternative in the future.

**Fluoroacetate dehalogenases**

According to the “Stockholm Convention,” all persistent organic pollutants are halogenated by fluorine, chlorine, or bromine. Therefore, dehalogenation is an important measure to control environmental pollution (Yue et al. 2020). Fluoroacetate dehalogenases (EC 3.8.1.3, FAcDs) are a class of enzymes that can cleave carbon-halogen bonds initiated by $S_N2$ substitution (Kim et al. 2017; Miranda-Rojas et al. 2018; Wang et al. 2017). Accordingly, FAcD is an effective tool to control pollution through enzymatic biodegradation (Li et al. 2019). Interestingly, FAcD can not only degrade fluoride but also obtain fluoride by kinetic resolution of racemates (Fig. 7B) (Zhang et al. 2020a). In this process, 500 mM ($RS$)-2-fluoro-2-phenylacetic acid derivatives were transformed to corresponding $R$-isomers and ($R$)-2-hydroxy-2-phenylacetic acid derivatives by FAcD (Conversion = 50%, Yields of $R$-fluorocarboxylic acids = 58–97%, ee > 97%). Hence, FAcD can not only eliminate environmental pollution but also effectively synthesize high value-added organic fluorides. It is just that the maximum theoretical yield of kinetic resolution is only 50% and there are by-products. These limit the application of FAcD in the field of fluorides biosynthesis. Therefore, it will be a huge improvement if by-products are converted into products or a two-phase separation system can be developed for products and by-products.

**Glycosidases: $\beta$-mannosynthase and $\beta$-glucosidase**

Typical glycosidases can hydrolyze sugars at specific locations and types (Li and Fan 2020). Excitingly, the C-F bonds can be formed by glycosidases (β-mannosynthase and β-glucosidase) through nucleophilic fluorination (Fig. 8A) (Nashiru et al. 2001; Zechel et al. 2001). This is the first enzyme catalyst with selective fluorination for C-F bonds discovered. Utilizing glycosidase, 23 mM 2,5-dinitrophenyl β-mannoside and 2 M KF were successfully transformed to β-mannosyl fluoride. Unfortunately, this fluorinated compound has not been isolated. This is because the fluorinated compound formed can also serve as a glycosyl donor and continue to participate in the transglycosylation reaction to form new glycosidic bonds. This process will cause the formed C-F bond to be degraded. Therefore, it is difficult to accumulate fluorinated products. This is why this enzyme has not been widely used to synthesize fluorinated compounds. In the future, it may become possible to use this enzyme for the biocatalytic synthesis of fluoride by controlling the progress of the reaction. Besides, a specific fluorinated product separation device may be necessary to utilize glycosidases.

**Fluorinases**

In 2002, a fluorination enzyme (fluorinase) was isolated and identified from the soil bacterium *Streptomyces cattleya* for the first time (O’Hagan et al. 2002). The fluorinase (EC 2.5.1.63) is also called 5’-fluoro-5’-deoxyadenosine (5’-FDA) synthase, which participates in the secondary
metabolism of fluorinated natural products. The fluorinase can catalyze the synthesis of 5'-FDA from S-adenosyl-L-methionine (SAM) and fluoride ion through nucleophilic attack to form a C-F bond (Fig. 8B, red dashed frame). As such, fluorinase has become an important biocatalyst for the synthesis of fluorinated nucleosides and their derivatives. Although fluorinase from *S. cattleya* had been applied to catalyze the non-natural substrates, a major drawback is that it exhibited reduced catalytic activity for unnatural substrates (Fralely and Sherman 2018). To overcome this obstacle, the directed evolution strategy was first implemented on fluorinase in 2016 (Sun et al. 2016; Thomsen et al. 2013). In this process, 23 sites were selected within a 5 Å of the substrate, and the best mutant F213Y/A279L was obtained after saturation mutation and high-throughput screening (1932 clones).

The catalytic efficiency of mutant F213Y/A279L was significantly increased by 25-fold compared to that of wild type for the non-natural substrate 5'-chloro-5'-deoxyadenosine (5'-CIDA). In addition, the efficiency in synthesizing radioactive 5'-[18F]FDA was also tripled compared to that of wild type. To expand the synthetic range of fluorinase, a new class of fluorinated nucleoside receptor agonists was synthesized from 0.1–0.2 mM chlorinated precursors, 50 mM KF, and 0.075 mM L-SeMet (Conversion = 10%, Yield = 3.5%) (Lowe et al. 2017). The agonist belongs to the analogs of 5'-FDA with a terminal functionalized ethynyl unit substituted at the C2 position. Similarly, the fluorinated RGD peptides and prostate-specific membrane antigen were also synthesized by fluorinase through transhalogenation reaction (Lowe et al. 2019b; Thompson et al. 2016a, 2015; Zhang et al. 2016).

The above-mentioned reaction was a two-step process in nature. In the first step, 5'-CIDA was transformed into SAM in the presence of fluorinase and L-Met. In the next step, the producing SAM was transformed into 5'-FDA in the presence of fluorinase and fluoride ions (Fig. 8B).

Moreover, the first reaction is the rate-limiting step for the synthesis of fluorinated compounds. To overcome the rate-limiting step, two novel SAM-dependent chlorinases were discovered to improve the efficiency of the transhalogenation reaction in the first step reaction. Finally, a chlorinase-fluorinase coupling system was developed. Through this system, 5'-FDA and 5'-fluorodeoxy-2-ethynyladenosine (5'-FDEA) were synthesized efficiently from 0.2 mM 5'-CIDA/5'-chloro-2-ethynyladenosine (5'-CIDEA), 0.1 mM L-SeMet, and 80 mM NaF (98.04 ± 0.27% and 91.60 ± 0.13% yields, respectively) (Sun et al. 2018). Surprisingly, Phillip T. Lowe et al. found that intermediate products SAM and L-methionine are not necessary for synthesizing 5'-FDA through fluorinase (Lowe et al. 2019a). Fluorinase can directly catalyze the synthesis of 5'-FDA from 5'-CIDA, 5'-bromo-5'-deoxyadenosine (5'-BrDA), and 5'-iodo-5'-deoxyadenosine (5'-IDA) in the presence of fluoride ion without SAM and L-Met (Finkelstein Reaction).

Additionally, the highest conversion efficiency was obtained for the substrate 5'-BrDA (35%) compared with 5'-IDA (8%) and 5'-CIDA (5%).

In addition to the catalytic process in vitro, the metabolic pathways of fluoride in microorganisms have also been elucidated, and related fluorinated natural products and biocatalysts have been confirmed (Figure 8B). Unfortunately, there are few sources of fluorinases, which limits the development of effective biocatalysts for the synthesis of fluorinated compounds. Before 2016, only 5 fluorinases were identified and characterized from *S. cattleya*, *Streptomyces* sp MA37, *Nocardia brasiliensis*, *Actinoplanes* sp N902-109 and *Streptomyces xinghaiensis* NRRL B24674 (Deng et al. 2014; Ma et al. 2016; Ren et al. 2020). Until recently, the sixth fluorinase was identified from *Actinopolyspora mzabensis* (AmFLA) (Sooklal et al. 2020). Due to AmFLA was an inclusion body, it was refolded into soluble protein through 8 M urea. The soluble AmFLA showed the specific activity of 0.44 ± 0.03 µM/min/mg for SAM at pH 7.2 and 65°C (both the optimal conditions). Besides, AmFLA also exhibited comparatively thermostability that it still retained 26% catalytic activity at 25°C for 2 months and >70% catalytic activity at 80°C for 20 min. Surprisingly, the specific activity of AmFLA was increased by 9.5% with the help of 1 mM Mg²⁺. This is the first time that fluorinase was found to be metal ion-dependent. To improve the activity and stability, fluorinase was immobilized with fluoridated hydroxyapatite nanoflowers (Li et al. 2020b). Under the same conditions, the activity of immobilized fluorinase is twice as high as the free enzyme. After 8 h of incubation at 30°C, 80% of activity was retained for the immobilized fluorinase and only 48% for free fluorinase. Apart from immobilization methods, protein self-assembly is also an effective strategy to improve activity and stability (Tu et al. 2020). With the help of self-assembled tags, fluorinase was formed into a novel artificial enzyme with nanometer dimensions. Strikingly, the novel artificial fluorinase exhibited higher enzyme activity, thermal stability, and reusability than the original fluorinase. Notably, fluorinase is currently the only biocatalyst widely used in the synthesis of fluorinated compounds that can form the C-F bond. In particular, fluorinase has great application potential for the synthesis of radioactive fluorinated imaging tracers. However, only limited potential has been exploited for fluorinase. Therefore, there is still much work to be done as soon as possible, especially considering that there are only six fluorinase sources, only two fluorinase crystal structures, low enzyme activity and narrow substrate range.

**Multienzyme system**

Not all products of interest can be obtained through a single biocatalyst as desired. Inspired by the natural multi-enzyme catalytic system, many high value-added products have been...
obtained through the artificial multi-enzymatic system, especially fluorinated compounds (Table 1). Accordingly, the artificial multi-enzymatic system was established for fluorinated aromatic compounds, including l-α-amino acid deaminase (LAAD), α-keto acid decarboxylase (ARO10), and aldehyde dehydrogenase (ALDH) (Figure S9) (Mao et al. 2020). Through the system, 10 mM m-fluoro-o/l-phenylalanine was transformed to m-fluoro-phenylacetic acid with a conversion of 48% in 24 h. The artificial multi-enzymatic system constructed has been proved to be effective in synthesizing non-natural fluorinated aromatic compounds without transgressing intermediate products. Similarly, a multi-enzyme system was performed with aromatic amino acid transaminase and hydroxymandelate synthase (HMS), producing fluoro-1-mandelic acid from 5 mM fluoro-o-phenylalanine (Yield = 65 ± 1.4%, ee > 86%) (Figure S10) (Youn et al. 2020). Moreover, ortho and meta fluorinated mandelic acids were also synthesized in the same method (Yields = 37 ± 0.4% and 47 ± 1.8%). Also starting from substrate fluoro-o-phenylalanine, fluoro-benzyl alcohol was successfully synthesized in a multi-enzymatic system of five enzymes, including LAAD, HMS, 1-mandelate dehydrogenase (LMDH), benzoylformate decarboxylase (BFD), and phenylacetaldehyde reductase (PAR) (_yield = 633.17 mg/L) (Figure S11) (Liu et al. 2020).

Besides, a catalytic system coupled with TPL and tyrosine ammonia-lyase (TAL) was established to obtain the fluorinated unsaturated carboxylic acid (Figure S12) (Busto et al. 2016). This overcomes the shortcomings of by-products produced in the direct alkenylation process. During the TPL-TAL coupling process, 23 mM fluorophenols and 46 mM pyruvate were converted into fluoro-o-tyrosines by TPL. Subsequently, the intermediate products obtained were transformed to p-hydroxyfluorocinnamic acids by TAL (Yield > 95%). Moreover, fluorinated galactoside was also synthesized by the multi-enzyme catalysis method (Figure S13) (Geissner et al. 2021). This is mainly because fluorinated galactoside can improve the stability of glycoprotein, and stable glycoprotein plays a vital role in the persistence of the therapeutic effect. In this process, 100 mM 4-deoxy-4-fluoro-N-acetylhexosamine and 500 mM pyruvate were transformed to 7-modified sialic acid through porcine N-acyl-o-glucosamine 2-epimerase and sialic acid aldolase (Conversion = 95%, Yield = 91%). Subsequently, the 7-modified sialic acid obtained was converted to 7-modified CMP-sialic acid (CMP = cytidine monophosphate) by CMP sialic acid synthetase and inorganic pyrophosphatase (Yield = 86%). Ultimately, the 7-modified CMP-sialic acid obtained was transformed to (7F-)sialyl galactoside by sialyltransferase and alkaline phosphatase (Conversion = 100%, Yield = 64%). The final results indicated that the half-life of the therapeutic glycoprotein was significantly provided by the incorporation of (7F-)sialyl galactoside synthesized.

The multi-enzyme synthesis of fluorine compounds can not only expand the application potential of enzymes as catalysts but also provide new ideas for the preparation of complex fluorinated products. Multi-enzyme catalysis can effectively shorten the transfer distance of intermediate products and accelerate the reaction rate. Moreover, the synthetic route can be realized from cheap substrates as starting substrates. However, there are still many challenges facing the development of the multi-enzymatic synthesis of fluorine compounds. First, the cost of the catalyast is increased due to the preparation of multiple enzymes. Second, the compatibility needs to be considered for multiple enzyme reaction conditions (for example, temperature, pH, metal

| Entry | Substrates | Multi-enzyme catalyst | Products |
|-------|------------|-----------------------|----------|
| 1     | m-fluoro-o/l-phenylalanine | l-α-amino acid deaminase <br> α-keto acid decarboxylase <br> aldehyde dehydrogenase | m-fluoro-phenylacetic acid |
| 2     | omlp-fluoro-o-phenylalanine | aromatic amino acid transaminase <br> hydroxymandelate synthase | omlp-fluoro-o-mandelic acids |
| 3     | fluoro-o-phenylalanine | l-α-amino acid deaminase <br> hydroxymandelate synthase <br> 1-mandelate dehydrogenase <br> benzoylformate decarboxylase <br> phenylacetaldehyde reductase | fluoro-benzyl alcohol |
| 4     | fluorophenols <br> pyruvate | tyrosine phenol-lyase <br> tyrosine ammonia-lyase | p-hydroxyfluorocinnamic acids |
| 5     | 4-deoxy-4-fluoro-N-acetylhexosamine <br> pyruvate | N-acyl-o-glucosamine 2-epimerase <br> sialic acid aldolase <br> CMP sialic acid synthetase <br> inorganic pyrophosphatase <br> sialyltransferase | (7F-)sialyl galactoside |
ions, inhibition of other enzymes by intermediate products or coenzymes). Finally, it is necessary to balance the activity and stability of multiple enzymes at the same time.

**Structure and catalytic mechanism of fluorinases**

As an important biocatalyst for the formation of the C-F bond, the structure and catalytic mechanism of fluorinase have always been a topic of concern in the field of biocatalysis and biotransformation. There is a close relationship between protein structure and function. Therefore, studying the structure of fluorinase is essential for understanding the catalytic mechanism of enzymes. To confirm the conserved amino acids, the multiple sequence alignment of six known fluorinases was constructed using DNAMAN (Figure S14). 182 conserved residues were identified, and the identity is as high as 85.11%. This indicated that the six known fluorinases have a high degree of homology and evolutionary origin from the primary structure of the protein. These conserved regions are the key to the catalytic function of fluorinases. Moreover, the crystal structures of fluorinases have been obtained only from two microbial sources, *Streptomyces cattleya*, and *Streptomyces* sp. MA37. The first crystal structure of fluorinase from *S. cattleya* was resolved to 1.9 Å resolution (Cobb et al. 2006; Dong et al. 2003, 2004). Through gel filtration analysis, the single frozen protein is a symmetrical hexamer (186 kDa) composed of two trimers. The trimer is composed of three identical monomers (Fig. 9A). The monomer has two domains: the amino-terminal domain (residues 1–180) and the carboxy-terminal domain (residues 195–298), and a loop structure that connects the two domains (residues 181–194) (Fig. 9B). The amino-terminal domain is composed of two antiparallel β-sheets with 4 and 5 helices. The carboxy-terminal domain has a 7-stranded β-sheet, which is sandwiched between the parallel and antiparallel α-helix strands. However, a single crystal structure cannot completely analyze the catalytic mechanism of fluorinase. As such, the crystal structure of fluorinase bound to substrate and product was obtained (Fig. 9C). In this structure, SAM was located between the C-terminal of a monomer and the N-terminal of a monomer in the trimer (Fig. 9D). The catalytic function of fluorinase must rely on the interactions of two monomers. As shown in Fig. 9E and F, the active sites of the fluorinase are distributed around the substrates or products and at the ends of two adjacent monomers. When combined, SAM is completely buried by fluorinase, which suggests that the fluorinase has other open conformations during the catalysis process. Comparing the binding status of substrate and product with fluorinase showed that 5’-FDA and L-methionine occupied the binding site of SAM after the reaction. Also, the difference between the two states is very small except for the bond formation/fracture area (Deng et al. 2004). This indicates that the key structures and regions are similar to the substrates and products of fluorinase and located between the two domains. Furthermore, the reaction is activated by changes in these key structures and regions.

In the field of organic chemistry, *S*<sub>2</sub>*<sub>2</sub> nucleophilic substitution is a paradigm reaction (Xie and Hase 2016). Through structure and function analysis, asymmetric *S*<sub>2</sub>*<sub>2</sub> nucleophilic fluorination is the key to the formation of C-F bonds for fluorinase (Pupo et al. 2018). Moreover, the *S*<sub>2</sub>*<sub>2</sub> reaction mechanism of fluorinase was also supported strongly by QM/MM calculations (Senn et al. 2005). In the process of forming the C-F bond, the reaction is thermodynamically driven due to the slow reaction *K*<sub>eq</sub> and the equilibrium of the reaction lies in the products. The *S*<sub>2</sub>*<sub>2</sub> nucleophilic attack will provide a hydrogen bond for the anion, which can stabilize the desolvated F<sup>-</sup> and is very important for the selective fluorination of sp<sup>2</sup> carbon centers (Lee et al. 2016). This indicates that the positively charged SAM sulfur is a crucial factor in stabilizing the partial desolvation and transition state. Consequently, the detailed molecular mechanism of fluorinase was further revealed (Zhu et al. 2007). During the catalysis process, solvated fluorine first binds to the unoccupied pockets and exchanges the hydrogen bonds of the water molecules to the polar groups of the protein (amide NH and side-chain OH of S158). The SAM is then bound, and the remaining water molecules enter the hydrophobic binding pocket to dissociate the fluorine. Remarkably, SAM is a competitive inhibitor of fluoride ion binding to fluorinase during the transition state. Afterwards, fluorine *S*<sub>2</sub>*<sub>2</sub> attacks C5 of SAM and replaces L-Met of SAM. After the conformation of loop (A95-Q102) changes, L-Met is released. Moreover, the loop participates in the combination of L-Met but not directly combined with L-Met. And then, the conformation of a loop (T75-R85) is changed to release 5’-FDA, which changes the binding region for the adenine ring and ribose of SAM and 5’-FDA derivatives.

To further explore the catalytic mechanism of fluorinases, a series of different types of substrates and mutants were analyzed and compared. When analyzing the crystal structure of fluorinase, an extensive hydrogen bond network was formed through water molecules and the carboxylate groups of L-Met with amino acids (T155, W217, Y266, backbone amide of S269) (Thomsen et al. 2013). The two water molecules were tightly bound at the active site of the fluorinase. This indicated that the instability of these water molecules may promote the accommodation of sterically bulky substrates. Therefore, site-directed mutagenesis was performed at these sites by introducing small hindered amino acids and disrupting the hydrogen bond network. Y266F and T155S exhibited significantly higher catalytic efficiency than that of the wild type. The results showed that the hydrogen bond network of the active center is an important factor for fluorinase to accommodate large sterically hindered substrates.
The two amino acid residues 266 and 155 play a significant role in the formation of the hydrogen bond network. For substrates modified at the C2 (R₁ substitution) and C6 (R₂ substitution) positions of the adenine ring, the substrates catalytic range was also studied (Figure S15) (Yeo et al. 2017). The results showed that substrates with hydrogen donors at the C2 position of the adenine group interact better with the fluorinase, obtaining a high yield. Besides, amino acid residue 279 plays an important role in promoting binding to ethynyl of the substrate at the C2 position. The C2 position is sensitive to large steric hindrances. Unfortunately, all substitutions at the C6 position caused catalysis difficulties, except for chlorine and hydrogen. Similarly, the directed evolution was also implemented in the process of catalyzing 5'-ClDA to SAM to 5'-FAD (Sun et al. 2016). Mutants F213Y/A279L have been shown to increase the yield of 5'-FDA. Specifically, F213Y/A279L can increase the turnover from 5'-ClDA to SAM but decrease the turnover from...
SAM to 5′-FAD. These mutations did not affect the affinity of the enzyme to 5′-CIDA but increase the affinity to SAM. The movement of the loop (277–281) is closely related to the binding of fluorinase and adenine. When position 279 is mutated to a large hindered amino acid, the movement of the loop is restricted, resulting in a tighter binding. Moreover, the hydrophobic environment facilitates the binding of substrates in the substrate-binding pocket. Leu279 and Tyr213 participate in favorable hydrophobic contact. The former improves the hydrophobicity of the environment through van der Waals forces with aromatic carbon of adenosine. The latter acts as a clamp between protein monomers across the ribose ring. Additionally, the reverse reaction of fluorinase was also applied to study the catalytic mechanism of the substrate with the difluoromethyl group (Thompson et al. 2016b). Regrettably, no reaction was observed in the presence of 5′,5′-difluoro-5′-deoxyadenosine 7 (F2 DA) and l-Met/l-SeMet. As such, the co-crystal structure of fluorinase and F2 DA, and tartrate was obtained to analyze the reason (1.8 Å). Structural analysis showed that the difluoromethyl bridged the interaction known to be essential for the activation of single fluorine in 5′-FDA. The bridging reaction was combined with the strong C-F bonds inherent in the difluoromethyl group, and there was an abnormal hydrogen bond interaction between the hydrogen of the difluoromethyl group and hydroxyl oxygen of the tartrate ligand. This indicated that the geometry of the bond is no longer suitable for the attack of sulfur or selenium and form higher response.

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Antitumor inhibitors

To effectively treat the tumor, inhibiting key enzymes of tumor cells is regarded as an important treatment. Hence, many fluorinated compounds have been developed as novel antitumor inhibitors, considering the problems of drug resistance, solubility, and toxicity. For example, diverse fluorinated cyclohexene derivatives were used to inhibit the activity of hOAT to treat HCC, because human ornithine aminotransferase (hOAT) plays an essential role in metabolic pathways of hepatocellular carcinoma (HCC) (Zhu et al. 2020b). Especially (1R,3S,4S)-3-amino-4,4-difluoro cyclopentanone-1-ene-1-carboxylic acid and (1R,3S,4S)-3-amino-4-fluoro cyclopentanone-1-ene-1-carboxylic acid, they showed the total different inactivation mechanisms from previous inhibitors. Moreover, tyrosine kinase (TK) is the epidermal growth factor receptor (EGFR) of A549 and Hela cancer cells. Four fluorinated quinazolinone derivatives showed
excellent anti-proliferative properties against A549 and Hela cancer cells by inhibiting TK (IC\textsubscript{50} = 0.462–7.568 µM; IC\textsubscript{50} = 0.175–7.096 µM) (Le et al. 2020).

In addition to better resistance, fluorine-containing inhibitors have a stronger inhibitory effect than traditional non-fluorine anti-tumor inhibitors. Such as, steroid sulfatase (STS) is an important enzyme to regulate metabolism in breast cancer cells, which is a malignant tumor with the highest incidence among women (Armstrong et al. 2020; Maltais et al. 2020). Therefore, various inhibitors of STS have been developed gradually. To improve the inhibitory effect, C-F bonds were introduced into STS inhibitors. Their inhibitory efficacy was ten-fold that of traditional inhibitor (coumarin-7-O-sulfamate), especially 3-(3,4-difluorophenyl)-2-oxo-2H-chromen-7-yl sulfamate, 2-oxo-3-(3,4,5-trifluorophenyl)-2H-chromen-7-yl sulfamate, 3-(2-(2,5-bis(trifluoromethyl)phenyl)acetamido)-2-oxo-2H-chromen-7-yl sulfamate, and 3-(3,4-difluorobenzamido)-2-oxo-2H-chromen-7-yl sulfamate (IC\textsubscript{50} = 0.18–0.27 µM) (Dasko et al. 2017; Demkowicz et al. 2016). Correspondingly, tautomerase is human macrophage migration inhibitory factor (MIF), which is also a pro-inflammatory cytokine associated with many tumors. Studies showed that the growth of the tumor can be effectively inhibited by MIF inhibitors and the inhibitory effect can be significantly enhanced by adding fluorine atoms (Dziedzic et al. 2015). Similarly, other fluorinated compounds also exhibited stronger inhibitory effects than non-fluorinated compounds for distinct tumor cells, such as fluorinated green tea polyphenols, fluorinated largazole, fluorinated 4-thiazolidinone, fluorinated griseofulvin, fluorinated l-threonine, fluorinated aminophosphonite, and fluorinated docetaxel (Zhang et al. 2019a) (Stadlbauer et al. 2018) (Makki et al. 2019; Olgun 2019; Paguigan et al. 2017; Sudileti et al. 2019; Tang et al. 2016). These results indicated that the incorporation of fluorine-substituted fragments into the inhibitor is an effective means to enhance the therapeutic effect of tumors.

Antiviral inhibitors

In addition to being novel anti-tumor inhibitors, fluorinated compounds are also effective against viruses. Coxsackie virus B (CVB) is a common pathogen for humans, causing aseptic meningitis, pericarditis, and myocarditis. To counter CVB, seven fluorinated pyrimidine nucleoside analogs were synthesized and tested for their antiviral activity (Tao et al. 2020). The results showed that 4-N-(2′-amino-glutarate-1′-methylester)-1-(2′-deoxy-2′-β-fluoro-4′-azido)-furansyl-cytosine has the most effective activity of anti-CVB (IC\textsubscript{50} = 9.3 µM). Similarly, 1-(2′-Deoxy-2′-fluoro-β-D -arabinofuranosyl)benzimidazoles were also proven to have anti-herpes virus activity (Khartinova et al. 2016). As for antiviral activity, the global health system is currently being destroyed by the novel coronavirus (COVID-19). Although diverse vaccines have been developed, they cannot produce a therapeutic effect on patients who have been infected with COVID-19. As such, it is crucial to find special drugs for COVID-19. As we all know, protease M\textsuperscript{pro} is an important drug target for COVID-19. Through molecular simulation, antiviral N-heterocycles (2′-fluoro-2′-deoxycytidine) was evaluated for its inhibitory effect on protease and showed good affinity for target receptors (Hagar et al. 2020). So far, there is no specific drug that can treat COVID-19 through experimental verification. But, it is believed that fluorinated compounds have this potential. Moreover, more and more fluorinated compounds will be artificially synthesized and used to fight viruses in the future of fluorinated biochemistry.

Pharmaceutical intermediates

Traditional pharmaceutical intermediates are increasingly unable to meet the needs of the current pharmaceutical industry. The development of novel pharmaceutical intermediates has become more and more urgent and necessary. Selective fluorination is an effective strategy to improve pharmaceutical efficacy, half-life, lipophilicity, absorption rate, and the permeability of cell membranes (Sudileti et al. 2019; Zhou et al. 2016). Therefore, the exploration has never stopped for the role of fluorine in drug design since the first fluorinated drug (9α-fluoro-substituted corticosteroid fluoro-cortisone) was approved in the 1950s (Fried and Sabo 1954). Nowadays, fluorine functional groups are contained in more than 25–30% of drugs, such as mefloquine, teriflunomide, citalopram, sorafenib, efavirenz, tedizolid phosphate, donepezil, and vorapaxar (Adler et al. 2019; Izquierdo et al. 2019; Konrad et al. 2019). A typical example is the use of fluorinated compounds as drug intermediates for the treatment of Alzheimer’s disease (AD). AD is associated with changes in cAMP (cyclic adenosine monophosphate) and cGMP (cyclic guanosine monophosphate) signals (Nakashima et al. 2019). Accordingly, regulating the levels of cAMP and cGMP in the brain has become an effective means of treating AD. It is known that phosphodiesterase 2A (PDE2A) is involved in the hydrolysis of cAMP and cGMP. Therefore, fluorine-substituted carboxamide analog as PDE2A inhibitors (TAK-915) was used to test the effect of treating AD. The results showed that TAK-915 was the most effective in treating AD when taken orally at 10 mg/kg. In addition, acetylcholinesterase is also an important target of AD. The activity of acetylcholinesterase can be inhibited by pipeline. To overcome the poor water solubility and photostability of piperine, a threo-difluoropiperine analog was synthesized and showed higher potency and selectivity than piperine for the treatment of AD (Lizarme-Salas et al. 2020). Similarly,
fluorinated compounds are also the key building blocks of antibiotic, antimalarial, anti-inflammatory, asthma, antagonists, migraine, and central nervous system drugs, such as 2,2-bis(6-fluoro-1H-indol-3-yl)ethan-1-amine, fluorinated steroidal, Trifluoroethymidine and fluorouracil polyoxin, fluorinated cholesterol, fluorinated galeagine, fluorinated bastimolide A, 6(R/S)-fluoropenibrugueramime, fluorinated cyclopropanecarboxylic acid derivatives, aryl-fluoro sulfates, 3-fluoro-4-aminopiperidine, and β-fluoramines (Adler et al. 2019; Bakhotmah and Abdel-Rahman 2017; Campana et al. 2020; Frank et al. 2016; Fujino et al. 2017; Gambini et al. 2019; Liu et al. 2018; Molinaro et al. 2019; Munck Af Rosenschold et al. 2019; Pupo et al. 2019; Quintard et al. 2018; Shao et al. 2015; Wu et al. 2020d). Accordingly, fluorinated compounds will pay more and more attention to the research and development of new pharmaceutical intermediates. Many intractable diseases are also expected to be cured.

### Functional materials

As we all know, white spot lesion (WSL) is the main cause of dental caries. The main measure of WSL management is to prevent the formation of new lesions. The fluorinated compounds can harden the surface layer of minerals, thereby preventing the occurrence of lesions and the development of dental caries (Decha et al. 2019; Lena Sezici et al. 2021; Raskin et al. 2021; Shah et al. 2018). Therefore, fluorinated compounds can be used to synthesize dental care materials, especially silver diamine fluoride, quaternary ammonium fluoride salt. Besides, the fluorinated membrane also displayed anti-pollution and anti-wetting properties in the field of industrial wastewater treatment (Abdulkarem et al. 2021; Cheng et al. 2021; Ji et al. 2021; Kang et al. 2021; Koh and Lee 2021; Li et al. 2021; Nayak and Tripathi 2021; Xiao et al. 2021; Zhong et al. 2021). Due to this unique performance, the fluorinated membrane can be used to treat wastewater with high salinity and hardness, remove heavy metal ions, separate oil, water, and as a flexible dielectric material and air filter with antibacterial properties. Moreover, the performance of traditional materials can also be enhanced by fluorne substitution. For instance, acrylate is the main component of rubber; Fluorinated acrylate is usually used to improve the compatibility and toughness of rubber (Yimmut et al. 2018). Similarly, fluorographene is a two-dimensional material with a high surface-to-mass ratio. Unlike graphene, fluorographene material is a wide band gap insulator because it lacks a conjugated network of π-conjugated electrons. As one of the thinnest insulators, fluorographene is most commonly used in the battery field. Fluorographene also plays an important role in industrial lubrication due to its delamination nature, low surface energy, and easy peeling characteristics (Chronopoulos et al. 2017; Liang et al. 2021). As such, fluorinated compounds are gradually becoming important sources of novel functional materials.

### Conclusions and perspectives

Fluorine is a distinctive element and fluoride-containing organic compounds are playing an increasingly important role in the fields of molecular imaging, pharmaceuticals, and materials. This has also led to high demand for reagents capable of selectively introducing fluorine into organics, especially biological enzyme catalysts. In this review, 13 enzymatic pathways are introduced. There are two main strategies for the enzymatic synthesis of fluorinated compounds: (1) C-F bonds are directly formed to obtain fluorine-containing compounds. (2) Complex fluorinated compounds are synthesized from simple fluorine-containing modules. Of all the current enzymatic methods, the catalytic route mediated by fluorinase has the most potential for applications. Nevertheless, the artificial green synthesis of fluorinated compounds still has a long way to go. Since C-F is a strong and inert chemical bond, it is extremely challenging to artificially synthesize compounds containing the C-F bond, especially the selective incorporation of fluorine into biologically active molecules.

Therefore, many future efforts will be made by our laboratory and other counterparts in the following aspects. (I) More and more new fluorinated natural products will be discovered and their biosynthetic pathways will be explored. Based on these biosynthetic pathways, artificial synthetic systems will be reconstructed to synthesize fluorinated natural products and their analogs. (II) Directed evolution based on machine learning is a mainstream tool for fluorinase, which will improve the efficiency and range of fluoride synthesis. Through this process, fluorinated nucleoside derivatives and non-nucleoside fluorides will have the opportunity to be obtained. (III) New enzymes that can directly form C-F bonds will be identified through metagenomics and gene mining methods. At that time, various fluorinated organic compounds will be easily obtained by direct fluorination at different positions and different groups. (IV) To introduce fluorine into valuable natural products, fluorine-containing building blocks will be introduced into complex natural product biosynthesis pathways through in vivo and in vitro synthetic biology methods. Besides, the toxicity of fluorides will also be overcome for synthetic biology in vivo based on the understanding of the mechanism of fluoride detoxification and detoxification of organisms.

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Declarations

Ethics approval This article does not contain any studies with human participants or animals performed by any of the authors.

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