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## Amine transaminases in chiral amines synthesis: recent advances and challenges

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## **Amine transaminases in chiral amines synthesis: recent advances and challenges**

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### **Abstract**

Transaminases, which catalyze the stereoselective transfer of an amino group between an amino donor and a prochiral ketone substrate, are interesting biocatalytic tools for the generation of optically pure chiral amines. In particular, amine transaminases (ATAs) are of industrial interest because they are capable of performing reductive amination reactions using a broad range of amine donors and acceptors. In this review, an update about the availability of microbial ATAs, discovered by both screening and database-mining approaches, or obtained by protein engineering of wild-type enzymes, will be provided. Current challenges in ATAs application and possible solutions will be also discussed. In particular, innovative biocatalytic process strategies aimed at the improvement of ATAs performances in chiral amines synthesis, e.g., using *in situ* product removal process strategies or flow reactors, will be presented. The progress in the industrial exploitation of these enzymes will be highlighted by selected examples of large-scale ATA-catalyzed processes.

**Keywords:** amine transaminases, biocatalysis, chiral amines, process development, protein engineering

## **Introduction**

Chiral amines are very important building blocks for the synthesis of several bioactive compounds and active pharmaceutical ingredients (Nugent and El-Shazly 2010; Paul et al. 2014). Since conventional chemical methods for the asymmetric synthesis of amines suffer from different limitations, e.g., low efficiency/selectivity and high environmental impact (Constable et al. 2007), the development of alternative biocatalytic routes has been largely investigated in the last two decades. To this aim, different types of enzymatic activities have been explored, such as amine dehydrogenases, monoamine oxidases, imine reductases, ammonia lyases, and transaminases. Among the latter, the enzymes capable of catalyzing reductive amination reactions using, as donors, simple amines, independently from the presence of a carboxylic group in the substrate, have been denominated amine transaminases (ATAs) (Steffen-Munsberg et al. 2015; Guo and Berglund 2017). These biocatalysts are pyridoxal-5'-phosphate (PLP)-dependent enzymes, well characterized from a structural and mechanistic point of view, and their applicative potential has been already established with several examples of chiral amines synthesis starting from readily available prochiral ketones (Mathew and Yun 2012; Brundiek and Höhne 2016). However, some limitations and challenges of ATAs for large scale applications have been identified as well, such as a quite narrow substrate scope, unfavorable thermodynamic reaction equilibria, and substrate/product inhibition. These findings have largely stimulated the search of both novel enzymes and innovative process conditions, and selected examples of the results obtained so far will be given in the following.

### **An update on newly discovered ATAs**

To date more than 60 microbial ATAs have been discovered (Guo and Berglund 2017), about twenty having been identified in just the last couple of years (Table 1).

<< *Insert Table 1 here* >>

New ATAs have been either isolated from wild-type microorganisms or identified in sequence databases by *in silico* screening and subsequently cloned and recombinantly overexpressed. Undoubtedly, sequence-based screening has proved to be a successful tool in ATA discovery, thanks both to the huge number of new sequences and genomes available in public databases and to the more and more in-depth knowledge of the sequence/structure-function relationships of ATAs (Pavkov-Keller et al. 2016; Höhne et al. 2010; Steffen-Munsberg et al. 2015).

Among the most recent examples reported in the literature, the search of ATAs by *in silico* screening has given interesting results in the identification of enzymes with high process stability, and, in particular, thermostability. In fact, thermostable proteins often show better performances in industrial applications thanks to their superior stability under operational conditions and in the presence of organic solvents. Different groups have searched for new ATAs in the genomes of thermophiles, i.e., microorganisms with an optimal growth temperature above 60°C. In particular, three new ATAs have been recently identified from thermophilic microorganisms, specifically from *Geobacillus thermodenitrificans* (Chen et al. 2016), *Sphaerobacter thermophilus* (Mathew et al. 2016b), and *Thermomicrobium roseum* (Mathew et al. 2016a). All these biocatalysts showed a thermophilic behavior with optimal temperatures at 60, 65, and 80°C, respectively, a high thermostability and activity increases after thermal treatment at 60-65°C. As an alternative to genome mining, metagenomics, i.e., the study of the genetic material directly obtained from environmental samples, has been also exploited to find thermostable ATAs (Ferrandi et al. 2017). In this work, three new ATAs have been identified by

*in silico* screening of a pool of metagenomic sequences from hot terrestrial environments. Among these, the B3-TA enzyme showed an outstanding thermostability, retaining 85% of activity after 5 days of incubation at 80°C. Interestingly, despite the high sequence similarity (92%) between B3-TA and the *T. roseum* ATA (Mathew et al. 2016a), the latter showed a much lower thermostability by losing completely its activity after 24 h incubation at 60°C.

As far as the substrate scope concerns, bifunctional transaminases that accept simple and cheap mono-/diamine donors as substrates have been recently discovered by analysis of multi-sequence alignments (Slabu et al. 2016; Galman et al. 2017). In particular, the enzymes from *Pseudomonas putida*, *P. chlororaphis*, and *P. aureofaciens* resulted to be very efficient in the stereoselective transamination of various ketones in the presence of natural diamines as amino donors, thus providing a convenient biocatalytic access to different pharmaceutically relevant chiral amines (Galman et al. 2017).

A bioinformatic search among crystal structures with unknown function has permitted instead the discovery of an ATA from *Bacillus anthracis* (Ban-TA) showing active site residues different from those in related (class III) transaminases, thus making very unlikely its function prediction (Steffen-Munsberg et al. 2016). Additionally, the use of key motifs based algorithms has recently allowed the discovery of a (*R*)-selective ATA from *C. semiimera* (Iglesias et al. 2017). This enzyme has shown a good activity toward a broad range of substrates, including *iso*-propylamine (*i*PA), an amino donor suitable for industrial applications.

However, sequence-based screening suffers from some intrinsic limitations, e.g., the fact that the novel enzymes often resemble in their activity and structural features the homologues used for the search and that it doesn't allow a quick identification of novel biocatalysts with a defined substrate scope. Thus, an alternative strategy to discover new ATAs involves the identification of microorganisms able to convert the substrate of choice, e.g., by employing a screening culture

containing the target amine as sole nitrogen source. Moreover, thanks to the ever lower cost of genome sequencing, the target enzymes can be quickly identified by *in silico* analysis of the bacterial genomes (Wilding et al. 2015, 2016; Pavkov-Keller et al. 2016; Wu et al. 2017).

By using this approach, a (*S*)-ATA from *Pseudomonas* sp. strain AAC able to metabolize 12-aminododecanoic acid, the constituent building block of homo-nylon-12, has been identified by enriching a bacterial strain collection with 12-aminododecanoic acid (Wilding et al. 2015). Further characterization has showed that this ATA is a highly promiscuous transaminase, catalyzing amine transfers with 39 of the 50 substrates tested (Wilding et al. 2016). In a similar way, two new (*R*)-ATAs have been isolated from *Curtobacterium pusillum* and *Microbacterium ginsengisoli*, respectively, by using (*R*)-amines for enrichment culture of microorganisms from soil samples. Interestingly, structural analysis of the *C. pusillum* enzyme has suggested the existence a subgroup within fold IV family of transaminases, showing significant phylogenetic distance from previously described ATAs as well as differences at the respective active sites (Pavkov-Keller et al. 2016).

### **Improvement of ATAs performances by protein engineering**

Despite microbial world represents a extraordinary source of new enzymes, it's unfortunately not very common to find in nature biocatalysts that show all the desirable features for industrial applications. To overcome this problem, different protein engineering approaches have been used to optimize available ATAs, a milestone having been set in 2010 with the outstanding work of Merck and Codexis in the development of a biocatalytic route for the synthesis of the drug sitagliptin (Savile et al. 2010). During the last years, thanks also to the increasing number of available transaminase crystal structures, structure-based rational design has been widely applied to this aim (Table 2).

<< *Insert Table 2 here* >>

Many efforts have been spent to broaden the ATA substrate scope, especially to make ATAs capable to accept bulky substrates. To this aim, specific residues in both the small (Hwang et al. 2008; Park et al. 2014; Nobili et al. 2015; Genz et al. 2016) and the large binding pocket (Cho et al. 2008; Han et al. 2015a-c) of ATAs active site have been changed to reduce steric constraints. For example, the small pocket of the ATA from *V. fluvialis* (*Vf*-ATA), which provides enough space mainly for a methyl group, as for example in **(1)** (Fig. 1) (Shin and Kim 2002), has been engineered to accept bulky substituents such as the side chain of **(2)** and **(3)** (Nobili et al. 2015).

<< *Insert Figure 1 here* >>

In particular, Y150, V153 and F85 residues have been identified to cause a steric hindrance with these substrates. In fact, a 53-fold improvement of the specific activity toward **(2)** (mutant Y150M/V153A) and a 26-fold improvement of the specific activity toward **(3)** (mutant F85L/V153A) could be obtained by site-directed mutagenesis. These variants have been subsequently exploited in the respective transamination reactions, giving 98% conversion and >98% enantiomeric excess (ee). In addition, the small binding pocket of *Vf*-ATA was further enlarged and reshaped, and the best variant (L56V/W57C/F85V/V153A) showed the highest specific activity towards **(4)**, an amine bearing a bulky *tert*-butyl substituent, not accessible by any of the wild-type ATAs tested before (Genz et al. 2016). Further studies have been carried out by Pavlidis et al. (2016) to evolve the ATA from *Rugeria* sp. toward the stereoselective transamination of bulky amines such as **(5-7)**, thus allowing the identification of a specific sequence motif (Y59/Y87/Y152/T213/P423). Interestingly, one of the best variants obtained in

this study was up to 8900 times more active toward **(6)**, and 310 and 98 times more active toward **(5)** and **(7)** than the wild type, respectively. Furthermore, the variant Y59W/Y87F/Y152F/T231A has been used in the asymmetric synthesis of various amines, the best result being achieved in the synthesis of **(6)** with 99% conversion and >99% ee. Further mutagenesis studies have allowed the identification of variants capable to accept also the substrates **(4)** and **(8)** (Weiß et al. 2016, 2017). Another attempt to expand ATA substrate specificity has been carried on the *Halomonas elongata* ATA (HEWT) by site directed mutagenesis of key residues located in the large pocket. The activity toward *ortho*-substituted acetophenone derivatives has been improved with the substitution W56G, while the mutation I258A has yielded an improvement in the reactivity toward *para*-substituted compounds (Contente et al. 2016).

The switch of ATA activity toward different functions by protein engineering has also been reported. For example, by changing the flipping arginine into an hydrophobic residue and reducing the steric hindrance of W57 and L417, *Vf*-ATA was converted from an amine: $\alpha$ -keto acid transferase into an amine:aldehyde transferase (Genz et al. 2015). Reversion or improvement of ATA enantioselectivity by structure based protein engineering have been also investigated (Svedendahl et al. 2010; Humble et al. 2012; Skalden et al. 2015). In particular, as until recently the available ATAs were predominantly (*S*)-selective, the need of (*R*)-selective ATAs has encouraged studies aimed at the switch of ATA enantioselectivity, one or two mutations in key residues being in some cases sufficient to reach this goal. For example, the substitution V328A in the *Arthrobacter citreus* ATA caused a shift from 98% ee (*S*) to 53% ee (*R*) in the synthesis of **(9)** (Svedendahl et al. 2010), while the double substitution F88A/A231F in the *Chromobacterium violaceum* ATA reversed the enantiomeric preference for **(10)** with an increase of the *E* value from 3.9 (*S*) to 63 (*R*) (Humble et al. 2012).



## Development of ATA-catalyzed processes

Besides enzyme inhibition and process stability, the major challenge that is often encountered in the set-up of ATA-catalyzed transaminations is the unfavorable thermodynamic equilibrium of the reaction (Brundiek and Höhne 2016). This problem could be avoided either by performing cascade reactions or by applying strategies to drive the reaction equilibrium toward amine synthesis. The use of an excess of co-substrate is the most employed method on the lab scale, but this approach may be limited by amine donor solubility and enzyme inhibition (Brundiek and Höhne 2016; Guo and Berglund 2017). Another option is to remove the product (or co-product) from the reaction by using different methods. Nitrogen sweeping is very efficient to remove the co-product acetone (Savile et al. 2010), but unfortunately, this strategy is applicable only if *i*PA is used as amino donor and this compound is not accepted by all ATAs. *In situ* product removal (ISPR) techniques (Lye and Woodley 1999), such as liquid/liquid or liquid/solid extraction, have proven to be effective in ketone product extraction but their applicability is generally hampered by the poor selectivity of the separation process, although some progress has been recently reported (Rehn et al. 2014; Börner et al. 2015; Satyawali et al. 2017). On the other hand, cascade systems have been largely explored (Simon et al. 2014). Coupled enzymatic reactions have been employed to convert pyruvate, the deamination product of the most commonly used amino donor alanine, into lactate by lactate dehydrogenase (LDH) together with a glucose/glucose dehydrogenase (GDH) system for cofactor regeneration (Fig. 2a) (Truppo et al. 2010).

<< *Insert Figure 2 here* >>

Pyruvate could be alternatively recycled into the transamination reaction by alanine dehydrogenase (AlaDH) when employing a (*S*)-selective ATA (Mutti et al. 2011), or it could be converted into D-alanine by an alanine racemase to be available for (*R*)-selective ATAs (Richter et al. 2015). Both ATA/AlaDH and ATA/LDH/GDH systems have proven to be quite efficient and in particular the latter system has been employed also on large scale (Girardin et al. 2013). Recently, the exploitation of the innate microbial metabolism to shift transamination equilibrium toward desired products by removing pyruvate in recombinant whole cell biocatalysts has also been proposed (Han and Shin 2014; Weber et al. 2017), while Farnberger et al. (2017) have investigated the co-expression in *E. coli* of ATAs together with the enzymes for co-product removal and cofactor regeneration, e.g., AlaDH and GDH. Either resting or lyophilized cells, as well as cell-free extracts, have been tested, however conversions of a model substrate didn't exceed 77%, thus suggesting the possible need of further fine-tuning of these strategies.

Aside product/co-product removal, the use of the so-called "smart" cosubstrates, whose products spontaneously transform into end-products, could be applied to drive the asymmetric synthesis of chiral amines toward completion. In this view, diamines that spontaneously cyclize into cyclic imines (Fig. 2b) have been successfully employed to this aim (Galman et al. 2017). However, this strategy is often limited by the strict substrate specificity of ATAs (Galman et al. 2017; Payer et al. 2017).

A promising alternative to the above described solutions lies in the exploitation of immobilized ATAs in flow systems where, as the substrate is converted into the product, the latter is removed from the reaction environment, thus avoiding enzyme inhibition and shifting the balance of the reaction toward product formation. Notably, enzyme immobilization often enhances protein operational stability, allowing at the same time further implementation of the process with in-line purification or ISPR steps and devices for analysis (Bajić et al. 2017; Heintz et al. 2017). In a

recent example, Planchestainer et al. (2017) has immobilized the poly-His-tagged *H. elongata* ATA (HEWT) on a metal derivatized epoxy-resin and synthesized a series of chiral amines in continuous flow with satisfying productivities. The packed bed reactor has been connected with an in-line purification system to achieve the easy recover of the amine product. Furthermore, it was demonstrated that the operational stability of HEWT was improved by immobilization, thus making its reuse for several cycles feasible.

### **Examples of large scale applications of ATAs**

The wild-type or engineered ATAs and the process implementation approaches described in the previous paragraphs have been not only investigated at a laboratory scale, but, in some cases, also tested in an industrial environment in the development of synthetic routes of different bioactive molecules.

For example, ATA 117 (Codexis) has been employed to produce (**12**), a useful building block in the synthesis of the orexin antagonist MK-6096, a candidate for the treatment of insomnia (Fig. 3a). The bioconversion, performed in 100 L total volume starting from 4.5 kg substrate and using D-alanine as amino donor with a coupled LDH/GDH regeneration system, has successfully resulted in 74% yield and 99% ee. Interestingly, this biocatalytic reaction could be well integrated in the kg-scale nine-step chemoenzymatic synthesis of MK-6096 with 13% overall yield (Girardin et al. 2013).

<< *Insert Figure 3 here* >>

Another remarkable example is the use of the ATA from *V. fluvialis* (*Vf*-ATA) for the production of (**14**) a key intermediate in the synthesis of the JAK2 kinase inhibitor AZD1480, developed by

Astra Zeneca for the treatment of idiopathic myelofibrosis and polycythaemia rubra vera (Frodsham et al. 2013; Meadows et al. 2013) (Fig. 3b). After biocatalyst selection, process conditions were optimized on gram scale by identification of  $\alpha$ -methyl benzyl amine as amino donor and of a biphasic system with 20% (v/v) toluene to increase substrate loading up to 0.35M and avoid enzyme inhibition by the co-product acetophenone. Subsequent scale-up of the biotransformation has allowed the conversion of 450 g of (**13**) into (**14**) in 79% yield and 99% ee. The intermediate has been then used in the chemoenzymatic synthesis of AZD1480 on 100 L scale with a >30% overall yield (Frodsham et al. 2013).

In a more recent example, Feng et al. (2017) have synthesized (**16**) (Fig. 3c), a useful building block for the production of the antibiotic besifloxacin, using a commercially available (*R*)-selective ATA. Under optimized reaction conditions, 0.3 kg of (**15**) were converted into (**16**) in the presence of 30% (v/v) *i*PA as amine donor with 80% isolated yield and ee >99%.

While in the abovementioned examples wild-type enzymes have been employed, Merck and Codexis had to improve ATA 117 by several rounds of mutagenesis to catalyze the last step of the synthesis of the antidiabetic compound sitagliptin (**18**, Fig. 3d) (Savile et al. 2010). After widening ATA 117 active site by 12 mutations in key "hot spots" to accept the bulky substrate pro-sitagliptin ketone (**17**), further improvements have been necessary to meet all the process requirements such as high activity at alkaline pH and at temperatures above 45°C and tolerance toward cosolvents and high substrate/cosubstrate concentrations. Eventually, a variant carrying 27 mutations was employed in the kg-scale synthesis of (**18**) using *i*PA as amino donor. Reaction was successfully driven toward product formation by stripping the volatile coproduct acetone, resulting in a 92% yield and >99.95% ee.

Other interesting examples have been recently reported using commercially available ATAs, although on a smaller scale (g scale). Specifically, Pfizer successfully employed ATA 036 in the

concomitant transamination and dynamic kinetic resolution of a 4-piperidone to obtain the precursor of an anti-cancer drug (Peng et al. 2014), while Merck evolved the enzyme ATA 013 to integrate a biocatalytic step in the chemoenzymatic synthesis of the antiarrhythmic drug Vernakalant (Limanto et al. 2014). In both cases good yields and excellent enantioselectivity have been achieved.

## **Conclusions**

Recent work on amine transaminases further confirm the huge potential of these enzymes in the development of innovative routes to prepare chiral amines up to the industrial level. The diversity of the available biocatalysts is rapidly improving thanks to both the discovery of novel natural enzymes and the generation of engineered variants, thus facing possible issues related to both substrate specificity/selectivity and stability. Moreover, the design and development of innovative process-based strategies will help to improve the efficiency and competitiveness of ATA-catalyzed reactions and further expand their practical application and integration of current production methods.

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## Tables

**Table 1.** Recently discovered and characterized ATAs (2015-2017).

Enzyme	GenBank accession number	Source	References
(S)-ATAs from thermophiles	ANF06848	<i>Sphaerobacter thermophilus</i>	Mathew et al. (2016b)
	KT719298	<i>Geobacillus thermodenitrificans</i>	Chen et al. (2016) Mathew et al. (2016a)
	WP_015922033	<i>Thermomicrobium roseum</i>	Mathew et al. (2016a)
(S)-ATAs from hot springs discovered by a metagenomic approach	KX505389	Uncultured organisms	Ferrandi et al. (2017)
	KX505387		
	KX505388		
(S)-ATA from halophilic bacteria	CBV41788	<i>Halomonas elongata</i>	Cerioli et al. (2015)
(S)-ATAs from oral cavity discovered by a metagenomic approach	Not available	Uncultured organisms	Baud et al. (2017)
(S)-ATAs active toward mono/diamine	WP_014461431	<i>Bacillus megaterium</i>	Slabu et al. (2016)

	WP_003196107	<i>Bacillus mycoides</i>	
	AJQ49991.1	<i>Pseudomonas putida</i>	Galman et al. (2017)
	KX954134.1	<i>Pseudomonas chlororaphis</i>	
	KX954133.1	<i>Pseudomonas fluorescens</i>	
(S)-ATA discovered in PDB database	KT861634	<i>Bacillus anthracis</i>	Steffen-Munsberg et al. (2016)
(S)-ATAs discovered by enrichment cultures	KES23458	<i>Pseudomonas</i> sp. ACC	Wilding et al. (2015, 2016)
	BAN53958	<i>Pseudomonas putida</i> NBRC14164	Wu et al. (2017)
	BAN52522		
	BAN55495		
	BAN57107		
(R)-ATAs discovered by enrichment cultures	KX160486	<i>Curtobacterium pusillum</i>	Pavkov-Keller et al. (2016)
	KX160489	<i>Microbacterium ginsengisoli</i>	
(R)-ATA from fungi discovered by genome mining	EWZ43325	<i>Fusarium oxysporum</i>	Gao et al. (2017)

(R)-ATA discovered using a key motifs based algorithm

ARU86455

*Capronia semiimera*

Iglesias et al. (2017)

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**Table 2.** Selected examples of recent structure-based ATA engineering

ATA source	ATA improvement	References
<i>Arthrobacter citreus</i>	Enantioselectivity switch	Svedendahl et al. (2010)
<i>Arthrobacter</i> sp. (ATA117)	Specificity toward bulky substrates Stability under process conditions	Savile et al. (2010)
<i>Aspergillus terreus</i>	Thermostability	Huang et al. (2017)
<i>Caulobacter crescentus</i>	Specificity toward bulky substrates	Hwang et al. (2008)
<i>Chromobacterium violaceum</i>	Enantioselectivity improvement and switch Specificity toward serine	Humble et al. (2012) Deszcz et al. (2015)
<i>Halomonas elongata</i>	Specificity toward acetophenone derivatives	Contente et al. (2016)
<i>Ochrobactrum anthropi</i>	Specificity toward bulky substrates	Han et al. (2015a-c)
<i>Paracoccus denitrificans</i>	Specificity toward bulky substrates	Park et al. (2014)
<i>Rugeria</i> sp.	Specificity toward bulky substrates	Steffen-Munsberg et al. (2013); Pavlidis

<i>Vibrio fluvialis</i>	Specificity toward bulky substrates	et al. (2016); Weiß et al. (2016, 2017)
	Specificity toward <i>(R)</i> -ethyl 5-methyl 3-oxooctanoate	Cho et al. (2008); Nobili et al. (2015); Genz et al. (2016)
	Specificity toward pentanal	Midelfort et al. (2013)
	Enantioselectivity improvement and switch	Genz et al. (2015)
		Skalden et al. (2015)

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## Figure legends

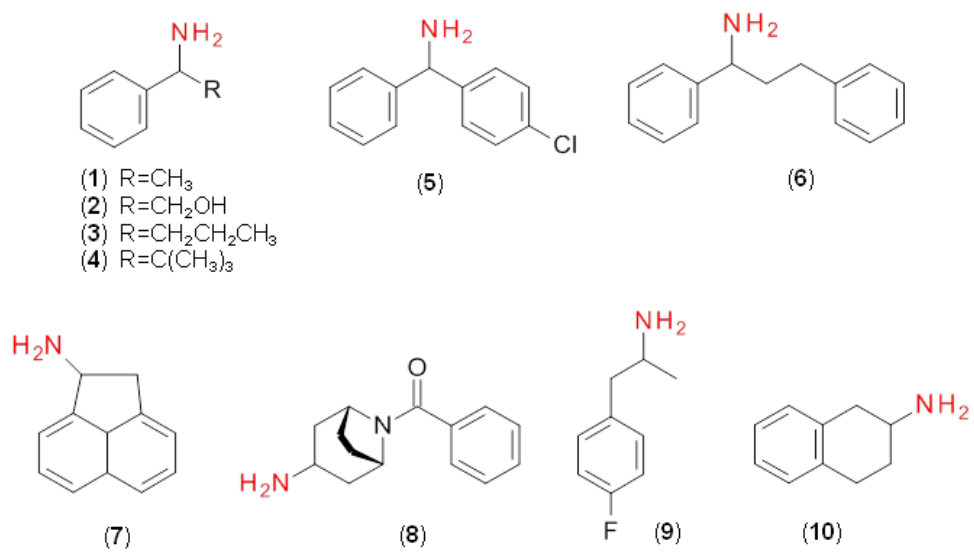
**Fig. 1.** Examples of target aromatic amines in protein engineering studies.

**Fig. 2.** Shifting the transamination equilibrium of ATA-catalyzed transaminations by a) applying an ATA/LDH/GDH cascade system, or b) using diamines as smart cosubstrates.

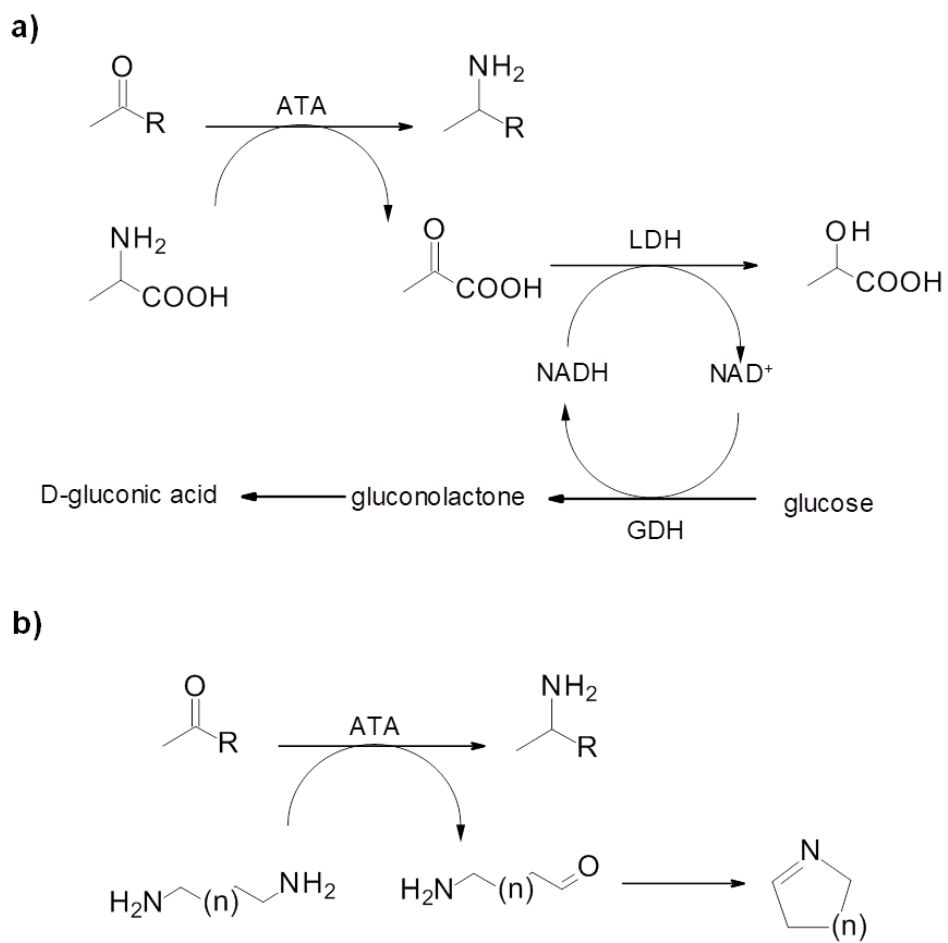
**Fig. 3.** Selected examples of exploitation of ATAs in the large scale synthesis of bioactive molecules

## Figures

Fig. 1



**Fig. 2**



**Fig. 3.**

