Zymophore identification enables the discovery of novel phenylalanine ammonia lyase enzymes

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The suite of biological catalysts found in Nature has the potential to contribute immensely to scientific advancements, ranging from industrial biotechnology to innovations in bioenergy and medical intervention. The endeavour to obtain a catalyst of choice is, however, wrought with challenges. Herein we report the design of a structure-based annotation system for the identification of functionally similar enzymes from diverse sequence backgrounds. Focusing on an enzymatic activity with demonstrated synthetic and therapeutic relevance, five new phenylalanine ammonia lyase (PAL) enzymes were discovered and characterised with respect to their potential applications. The variation and novelty of various desirable traits seen in these previously uncharacterised enzymes demonstrates the importance of effective sequence annotation in unlocking the potential diversity that Nature provides in the search for tailored biological tools. This new method has commercial relevance as a strategy for assaying the ‘evolvability’ of certain enzyme features, thus streamlining and informing protein engineering efforts.

Currently, a major challenge in the area of enzyme discovery is the accurate retrieval from databases of enzymes with specific traits, from among the vast number of related sequences, an approach hindered by misannotation from sequence curation methods which are often implemented in the absence of specialist protein functional knowledge. This problem is especially relevant in enzymes which are structurally diverse but have convergently evolved similar catalytic residue groupings, or in families where the function of choice constitutes an interspersed minority of available sequences. The former problem has been addressed in an enzyme class-specific manner by the computational assignment of clusters of active site residues conferring the desired chemistry on otherwise uncharacterised proteins. Examples include the mapping of active site ‘constellations’ for ene-reductase activity, use of BioGPS descriptors to aid discovery of promiscuous side-activity or the retrieval of catalytic residue arrangements via the Catalytic Site Atlas or PatternQuery. Whilst these methods provide simple means of accessing proteins with desired catalytic activity, their use is mostly limited to structural databases containing only a fraction of the number of entries that a sequence database would have. Such constraints are also less amenable to family-specific catalytic arrangements not documented elsewhere in nature where a wealth of sequence data can potentially harbour more scope for investigation. The structural methods also do not take into account other active site residues which differentiate desired and unwanted activity, such as selectivity and substrate discrimination.

With the ever diminishing cost and labour associated with DNA sequencing, cloning and synthesis, it is increasingly conceivable to imagine harnessing, from publicly available databases, the natural protein sequence diversity provided by evolution. Such an approach offers several advantages over more traditional engineering methods, particularly for enzymes where robust high-throughput screening methodologies do not exist. It also provides a means of mitigating changes deleterious to enzyme integrity whilst also allowing exploration

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of variation at non-active site positions, said to be important for contributions to activity\(^8\), and which are more
umerous and thus difficult to target effectively.

Enzymes with phenylalanine ammonia lyase (PAL) activity are able to catalyse non-oxidative deamination
to yield cinnamate (1a) and ammonia from the proteinogenic amino acid L-phenylalanine (L-Phe, 2a)\(^9\). The
amine abstraction, in this case, is mediated by adduct formation with a 4-methylidenediamidazole-5-one (MIO)
post-translational modification in the active site\(^{10,11}\). This in turn promotes elimination in conjunction with pro-
ton abstraction via a catalytic tyrosine residue situated on an inner active site mobile loop \(^{11,12}\). In nature, this
PAL-mediated reaction often constitutes a gateway from primary metabolism to specialist compound biosyn-
thesis, with examples including phenylpropanoids in plants\(^{13}\) and various antibiotics in bacteria (Fig. 1a)\(^{14-17}\).

In the field of biomedical research, the selective catabolism of L-Phe by PALs has been exploited as a treatment
for a number of diseases. One such example is phenylketonuria, an inborn error of metabolism in which meta-
bo-lic dysregulation leads to neurological symptoms such as microcephaly, seizures and intellectual disabilities\(^{18}\).
As the condition emanates from aromatic amino acid imbalance - high phenylalanine and low tyrosine lev-
els - removing L-Phe has been shown to offer great advantages over alternative treatments based on complex
dietary restriction which are currently prescribed\(^{19,20}\). Essential amino acids, required by all growing cells, can
also be degraded in vivo to restrict cancer tumour growth (amino acid depletion therapy), a promising strategy
that has been demonstrated with metabolites such as L-Asn, L-Arg, L-Met, L-Tyr and aptly L-Phe\(^21\). Relevant
examples with respect to the last example in particular include the decrease in metastatic phenotype of B16BL5
melanoma upon aromatic amino acid depletion\(^22\) and the use of PAL-based formulations to prevent growth of
abnormal lymphocytes both in vitro\(^23\) and in leukemic mice\(^24\). The commercial importance of PALs as therapeu-
tic agents is evidenced by various patent applications from specialist enzyme (Codexis)\(^{25}\) and biopharmaceuti-
cal (BioMarin)\(^{26}\) production companies, as well as from multinational drug and chemical corporations (Novo
Nordisk and BASF)\(^{27,28}\).

PALs also find use in the area of green chemistry and sustainable pharmaceutical manufacture as industrial
biocatalysts, due to the reversibility of the deamination reaction (Fig. 1b). The use of concentrated yet inexpen-
sive ammonia buffers has been demonstrated to allow regio- and enantioselective amination of a broad range of
readily-accessible arylacrylic acids, yielding high value unnatural amino acids, a route that is difficult to emulate
via chemical methods. Basic molecular cloning techniques have enabled the simple preparation of enzyme-based
catalysts from renewable feedstocks, allowing the exploitation of their inherent selectivity and ambient reaction
conditions by synthetic chemists. As such, there are many examples of biocatalytic routes in industry, including
PAL-mediated processes to produce L-Phe\(^29\), unnatural amino acid building blocks of antifungal peptides (e.g., 3)\(^{30}\),
and the antihypertensive drug perindopril (4)\(^{31,32}\). The use of PALs for aromatic amino acid synthesis is particu-
larly attractive due to the 100% atom efficiency and coenzyme independence of the reaction, mitigating waste
production, by-product separation requirements and mediator supplementation/regeneration costs\(^33\).

In spite of the broad scope of application associated with PALs, their use is limited to a handful of specific
enzymes\(^33,34\), greatly restricting the variability available in desirable traits. Recent efforts have focused on mod-
ication of the PAL from Anaabaena variabilis (AvPAL), via mutagenesis or chemical conjugation, to overcome
characteristics of the enzyme which are considered detrimental to therapeutic applications\(^35,26,35\). Such traits gen-
erally include low catalytic activity, poor stability at 37 °C, accessible proteolytic cleavage sites, low acid tolerance
(if taken orally), aggregation and high immunogenicity. The few PALs used in a biocatalytic context (of which
AvPAL is again the most prominent)\(^34-36\) show more immediate promise, with little or no engineering required
for some industrial applications. However many pharmaceutically-relevant targets that are potentially within
the remit of the synthetic PAL reaction remain out of reach due to the restricted substrate scope of characterised
enzymes.

In the class I lyase-like family, aminomutases and histidine/tyrosine-specific enzymes are reasoned to form
the majority of available sequences, with PALs constituting the minority\(^27\). We therefore set out to develop a new
strategy for the discovery of PAL enzymes based on the four-step process outlined in Fig. 2.
Results and Discussion

Although several crystal structures of PAL enzymes have been solved, none exists with the natural ligand trans-cinnamic acid bound in the active site, thereby preventing correct and unambiguous assignment of the phenylalanine-specific MIO-dependent ammonia lyase zymophore. The gene encoding AvPAL was mutated, incorporating two surface substitutions (C503S and C565S, known to aid crystallisation) and replacement of the loop tyrosine with a catalytically inert but shape-complementary phenylalanine (Y78F). The triple variant was purified and co-crystallised with the natural deamination product. Following successful crystal growth, the structure of the AvPAL-ligand complex (PDB ID: 5LTM) was solved by molecular replacement using the previously obtained ligand-free double variant (PDB ID: 3CZO) as a search model. Superimposition of these two structures revealed excellent complementarity with a few small differences between the unoccupied and occupied active sites. The only backbone displacement was a shift in the outer active site loop, with the fully closed conformation from the previously reported structure seen to clash with the inner active site position 78 in the co-crystallised protein. In the case of the unoccupied enzyme, the amino acid at position 78 is tyrosine which points inward toward the active site, providing no contact with the outer loop, whereas the new structure features Y78F with a dihedral swing of ~140° causing the bulky side chain to protrude outwards. Other differences included H359 moving into the active site sphere in the cinnamate bound structure, along with smaller shifts in the positions of Q311 and R317 (SI, Fig. S1). Presumably these represent movements which occur upon substrate binding, although they may be induced instead by the Y78F variation as exchange of polar and non-polar residues in the inner active site loop of related enzymes has been reported to influence conformational dynamics.

Interestingly, these three seemingly mobile positions have been identified as hotspots in mutagenic studies of various PAL-relatives, with variation at H359 shown to affect catalytic efficiency and the two carboxylate-binding positions (Q311 and R317) being implemented in substrate positioning and selectivity. This observation sheds light on possible conformational selection or induced fit models which may help to guide engineering of enzymes in both this class and others by targeting residues which distinguish bound and unbound conformations.

Visualisation of the occupied active site allowed the identification of 19 residues within a 6 Å sphere of the ligand (Fig. 3), including the MIO and catalytic amino acid at position 78, the carboxyl positioning residues, the hydrophobic aryl binding pocket and other assorted substrate-constraining side chains. As a test of the orthogonality of this PAL zymophore, a sequence alignment was performed using characterised ammonia lyase and aminomutase enzymes specific to phenylalanines and/or tyrosine or histidine (SI, Fig. S3). All the PALs in the selection showed conservation at all or all but one of the residues, while ammonia lyases accepting both L-Phe and L-Tyr (bifunctional PAL/TALs) and (R)-selective aminomutases ((R)-PAMs) varied at two positions. Bacterial lysases and mutases specific to tyrosine (TALs and TAMs) and histidine ammonia lyases (HALs) showed conservation at only 15 residues with (S)-β-Phe forming mutases ((S)-PAMs) scoring just 13/19. Interestingly, even those enzymes which differed by the same number of residues, tended to do so at different positions, highlighting the importance of using as many of the residues as possible in further investigations. As such, the entire motif was used and a score of 18/19 was deemed to be the threshold for consideration of sequences as potential PALs.

The 19 amino acid active-site sequence motif, as inferred from the AvPAL-cinnamate structure, was used to distinguish potentially useful enzyme sequences from other family members. The sequences were collected using the basic local alignment search tool (BLAST) to search the entire knowledge database of the universal protein resource (UniProtKB) using various query sequences.

As the most widely reported enzyme of this class in applied research, the protein sequence of AvPAL was used for the first search and the 100 hits showing the closest identity were analysed. Of these, 17 were shown to give close agreement with the desired active site, with the only differences occurring at positions 6 and 7 of the motif at all. Four of the results, including the previously characterised PAL from Nostoc punctiforme (NpPAL), had hydrophobic residues at these positions, whereas all others contained a histidine at position 7. This variation is known to be associated with strict TAL activity in the bacterial enzyme BagA. The 13 potential TALs were discounted along with NpPAL leaving a closely related sequence from Oscillatoria sp., and two more distant relatives from Planctomyces brasilensis and Methylobacterium sp. Next, the sequence of the phylogenetically-distinct PAL, StIA from Photorhabdus luminescens, was used to search for PALs in a different part of the family tree. This time, within the first 100 hits the majority were found to contain YH or FH at positions 6 and 7, indicating possible unwanted TAL activity. Among the few PAL-like sequences, most were from the related enterobacterial genera Photorhabdus and Yersinia sharing high sequence identity with the query sequence. The remaining four
PAL sequences were surprising in that they were from the amoebozoa genus *Dictyostelium*, whereas all other sequences were of eubacterial origin.

Lastly, the bacterial TAL sequence BagA from *Streptomyces* sp. was used for one final search to find PAL sequences not closely related to those currently known. It is worth noting that, although most of the hits were found to be TAL-like, with YH and FH at positions 6 and 7, there were more notable and diverse examples of active sites matching the AvPAL-derived motif than with either of the searches using characterised PAL query sequences. These included the previously reported RxPAL from *Rubrobacter xylophilans*44, as well as additional sequences from *Bacillus subtilis* and *Actinopolyspora erythraea* and several from both *Brevibacillus laterosporus* and various species of the genus *Streptomyces*.

Interestingly the PALs from *Dictyostelium discoideum* and *Brevibacillus laterosporus* have been previously identified by Nielsen et al. using a synteny-based approach to search for class I lyase-like genes in the putative context of TAL-containing metabolic pathways45. This is an interesting finding, given the identification of these enzymes as PALs by their active sites in this study (rather than TALs based on inference of genetic co-localisation) and also given the rarity of PAL-like sequences seen in our database searches compared to the abundant TAL-like sequences. To add further evidence to the assertion that these sequences possessed PAL activity, genetic context analyses were conducted, revealing association with fatty acid and polyketide biosynthetic operons as has been reported previously with this class of enzyme42. In the case of *Brevibacillus laterosporus* the enzyme could even be putatively linked to the production of the cinnamate component of basiliskamides A and B, a link missing from previous biosynthetic hypotheses (SI, Fig. S7)46.

Five of the potential PAL sequences mined from the protein database using the AvPAL zymophore motif were chosen for characterisation based on their sequence identity to their closest characterised relative and the results of the genetic context analyses (Table 1).

In all searches using characterised query sequences, evidence was uncovered of multiple PAL-TAL neofunctionalisation events within certain clades in the family, with indication of known PAL-active site motifs (with FL at selectivity positions) interspersed among known TAL motifs (with YH) and a possibly intermediary motif (FH). These findings point to the incidence of several substrate switching events resulting in distinct and

**Table 1.** The most prominent PAL sequences identified from UniProtKB.

| Name | Origin                  | Motif match | Query   | Seq. id. |
|------|-------------------------|-------------|---------|----------|
| BlPAL| *Brevibacillus laterosporus* | 19/19       | BagA    | 58%      |
| SrPAL| *Streptomyces rimosus*   | 19/19       | BagA    | 54%      |
| MxPAL| *Methylobacterium* sp.   | 18/19       | AvPAL   | 50%      |
| DdPAL| *Dictyostelium discoideum* | 18/19     | StlA    | 48%      |
| PbPAL| *Planctomyces brasiliensis* | 19/19     | AvPAL   | 46%      |

**Figure 3.** The zymophore conferring PAL-specific reactivity and selectivity as identified from the co-crystal structure of the ammonia lyase from *Anabaena variabilis* (AvPAL) and its deamination product trans-cinnamate (PDB ID: 5LTM). The motif is shown on both structural overlay of the empty and occupied enzyme active sites and the sequence with each of the 19 amino acids given a number to mark its position.
unreported clades of TAL enzymes. This phenomenon may constitute an evolutionary strategy for the creation of chemical diversity in related biosynthetic pathways, interchanging between use of L-Phe or L-Tyr as a starting point, followed by further diversification and mixing of secondary metabolic enzymes to create new products. The searches also reveal that TAL-like sequences vastly outnumber PALs in the areas of sequence space targeted in this study, further demonstrating the importance of robust annotation methods over more random sampling of sequence diversity with this family of enzymes. The most interesting new sequences from the database were the PALs from Dictyostelium sp. including DdPAL, an example of an aromatic amino acid ammonia lyase from the protist kingdom. At around 50% sequence identity to StLA, this enzyme arises from a fairly recent gene transfer event between bacteria and eukaryotes, separate to that more widely reported to fungi and plants.37 We suggest that this gene transfer is the evolutionary result of a sustained endosymbiotic relationship between amoebozoa and bacteria, similar to that hypothesised for the acquisition of PALs in other kingdoms. This proposal aligns with reports of primitive agriculture between Dictyostelium discoideum and their bacterial livestock,37 an interaction which could result in xenologous proteins in these otherwise unrelated organisms.

Of the enzymes selected for further study, BIPAL and SrPAL were cloned from their host organisms whereas the genes for DdPAL and MxPAL were custom synthesised as codon-optimised constructs for expression in laboratory strain E. coli. These genes were then subcloned into expression plasmids. The gene encoding PbPAL was prepared and provided in an appropriate vector by collaborators at Prozomix Limited. After production of all five proteins as N-terminal His6-tagged variants, and successful purification of four, each enzyme was characterised with respect to amino acid specificity, pH optimum and stability at 37 °C (Fig. 4). To verify that these catalysts did not possess TAL or HAL activity, the isolated enzymes were incubated with L-Tyr or L-His and production of the corresponding deamination products assayed spectrophotometrically. In all cases, significant increases in absorbance, indicating acrylic acid production, were not seen with either L-Tyr or L-His (SI, Fig. S8). All four of the corresponding deamination products assayed spectrophotometrically. In all cases, significant increases in absorbance, indicating acrylic acid production, were not seen with either L-Tyr or L-His (SI, Fig. S8). All four enzymes possessing exclusively PAL-activity were also shown to be misannotated in UniProtKB, with all designations of HALs except for BIPAL which was incorrectly assigned TAL function (Table 1). These incorrect designations, widespread in public databases, impair the efforts of scientists in the search for new disease treatments, biocatalyst templates and components of unknown metabolic pathways. The case of enzymes with PAL activity illustrates this prominently, as correct annotation of these enzymes enables advances to be made in all three of these areas (e.g treatment of cancer/PKU, production of unnatural amino acids, identification of operons for bioactive molecules like basiliskamides).

To confirm PAL activity (rather than TAL or HAL) the specific activities with respect to the deamination of L-Phe were tested with each enzyme. Additionally, relative conversions after treatment under different conditions were also compared to assess the natural variation in therapeutically-relevant traits such as stability at blood and gastrointestinal temperature and pH (Fig. 4). The specific activity measurements made with each ammonia lyase indicated that SrPAL was the most active enzyme in the group, with a specific activity of 1102 mU mg⁻¹, a value far higher than that of the other enzymes which all gave ~500 mU mg⁻¹. The differences in specific activity between the enzymes were striking, particularly as SrPAL was shown to be far more active than the others with greater retention of activity after incubation at 37 °C. This difference was of particular interest as SrPAL and BIPAL share the closest sequence identity of any two enzymes in this study (SI, Fig. S6) and yet have a two-fold difference in turnover rate. These sequences, although closely related, possess many amino acid substitutions which are likely to influence the dynamics of the entire protein, possibly accelerating movements constituting the catalytic cycle or channelling solvent vibrations to the active site to help overcome activation energy.48,49 The identical active site, yet variable overall sequence of SrPAL compared to BIPAL, seems to contribute to the former enzyme being a better candidate as a biotherapeutic, due to its high activity and longevity at average body temperature after 1 hour. However, after 24 hours all enzymes still allowed 50–60% of their initial maximal conversion with only DdPAL showing a greater loss in activity. SrPAL was also found to be the most acid tolerant of the enzymes along with DdPAL, which, whilst being less active and less robust initially, still retained more activity at pH 6 that either BIPAL or PbPAL. This feature may make SrPAL (or even DdPAL) a good starting point for the development of an acid tolerant enzyme for oral administration, ensuring function and stability in the digestive tract. These

Figure 4. Variation of specific activity (a), pH tolerance (b) and stability at 37 °C (c) among four new ammonia lyases from Streptomyces rimosus, Brevibacillus laterosporus, Planctomyces brasiliensis and Dictyostelium discoideum which could be isolated for investigation.
variable findings with no one protein possessing all the most desirable traits, highlight the utility of our simple method for estimating the viability of enzyme engineering by sampling variation capacity in a handful of diverse but related natural sequences.

In order to evaluate the synthetic potential of each enzyme, the *E. coli* cell pastes prepared for protein purification were used as whole cell biocatalysts for the amination of a range of ring-substituted arylacrylates. Following a reported method, the biotransformations were run with all five enzymes and a representative panel of 16 cinamamate substrates 1a-p before isolation of crude product and analysis (Table 2). Interestingly, despite the biocatalysts having nearly identical active sites, there was evident variability in substrate tolerance and enantioselectivity, aside from general trends expected from previous investigations (e.g., low enantioselectivity with 1a).

Table 2. PAL-catalysed amination of various ring-substituted phenylacrylate derivatives. *a* Determined by reverse phase HPLC on a non-chiral phase. *b* Determined by reverse phase HPLC on a chiral phase.

| Substrate | Conv. (%) | ee (%) | Conv. (%) | ee (%) | Conv. (%) | ee (%) | Conv. (%) | ee (%) | Conv. (%) | ee (%) |
|-----------|-----------|--------|-----------|--------|-----------|--------|-----------|--------|-----------|--------|
| 1a R      |           |        |           |        |           |        |           |        |           |        |
| H         | 92        | >99    | 80        | >99    | 91        | >99    | 11        |       | 25        | 95     |
| 2-F       |           |        |           |        |           |        |           |        |           |        |
| 3-F       | 96        | >99    | 96        | >99    | 91        | >99    | 96        | >99    | 60        | 88     |
| 4-F       | 92        | >99    | 96        | >99    | 91        | >99    | 96        | >99    | 58        | 82     |
| 3-Br      |           |        |           |        |           |        |           |        |           |        |
| 4-Br      | 96        | >99    | 96        | >99    | 97        | >99    | 96        | >99    | 57        | 82     |
| 2-Cl      |           |        |           |        |           |        |           |        |           |        |
| 3-Cl      | 96        | >99    | 96        | >99    | 97        | >99    | 96        | >99    | 28        | 99     |
| 4-Cl      | 92        | >99    | 96        | >99    | 97        | >99    | 96        | >99    | 36        | 44     |
| 3-MeO     | <1        |        |           | <1     | 27        | >99    | 27        | >99    | 15        | >99    |
| 4-MeO     | 19        | >99    | 19        | >99    | 92        | >99    | 92        | >99    | 40        | 96     |
| 2-NO₂     |           |        |           |        |           |        |           |        |           |        |
| 3-NO₂     | 97        | >99    | 97        | >99    | 99        | >99    | 99        | >99    | 70        | 86     |
| 4-NO₂     | 92        | >99    | 92        | >99    | 99        | >99    | 99        | >99    | 68        | 82     |

The discovery that this new panel of enzymes showed high activity with substrates possessing electron-donating substituents was striking, as this class of substrate is not reported to be well accepted by PAL enzymes. Production of L-5-methoxyphenylalanine (S)-2l by three of the new PALs is of particular interest, as this unnatural amino acid is a relevant building block for pharmaceutical synthesis (Fig. 5). Examples of APIs that can be synthesised from this amino acid are antiviral peptides, such as 1a.

When compared to BIPAL however, SrPAL gave similarly excellent enantioselectic excess values across the board but much less variable conversion. A comparison between the three most discernible substrate profiles of BIPAL, MxPAL and DdPAL revealed that even the relative conversions were not the same between the enzymes. Whilst BIPAL gave the highest conversion for 1g, followed by MxPAL and then DdPAL (76%, 56% and 28%) this pattern was reversed with the ortho-isomer 1e (12%, 63% and 84%) and different again for the meta-isomer 1f (76%, 56% and 28%). Another interesting feature shared by MxPAL and DdPAL is that they both convert the natural ligand 1a to a lower extent than most of its ring-substituted derivatives. DdPAL even gave higher conversion with the 57% (1a) and 92% (1b) meta-substituted ligands than could be generated by the former enzyme in most cases. When compared to BIPAL however, SrPAL gave similarly excellent enantioselective excess values across the board but much less variable conversion. A comparison between the three most discernible substrate profiles of BIPAL, MxPAL and DdPAL revealed that even the relative conversions were not the same between the enzymes. Whilst BIPAL gave the highest conversion for 1g, followed by MxPAL and then DdPAL (76%, 56% and 28%) this pattern was reversed with the ortho-isomer 1e (12%, 63% and 84%) and different again for the meta-isomer 1f (76%, 51% and 57%). Another interesting feature shared by MxPAL and DdPAL is that they both convert the natural ligand 1a to a lower extent than most of its ring-substituted derivatives. DdPAL even gave higher conversion with the seemingly challenging substrate 1l, a compound not even accepted by BIPAL or MxPAL and poorly converted by the otherwise highly active SrPAL.

The discovery that this new panel of enzymes showed high activity with substrates possessing electron-donating substituents was striking, as this class of substrate is not reported to be well accepted by PAL enzymes. Production of L-5-methoxyphenylalanine (S)-2l by three of the new PALs is of particular interest, as this unnatural amino acid is a relevant building block for pharmaceutical synthesis (Fig. 5). Examples of APIs that can be synthesised from this amino acid are antiviral peptides, such as 1a, and oxazolidinone antibacterials, such as 6.[2] In light of this, a preparative scale amination of 1l was performed using PbPAL, due to its superior enantioselective and impressively high conversions compared to SrPAL and DdPAL. By repeating the biotransformation with a 10-fold increase in substrate loading and reaction volume (with 50 mg mL$^{-1}$ whole cells) an isolated yield of 61% enantiomerically pure L-amino acid (ee > 99%) was obtained from 89 mg of starting material.

The differences in relative conversion and enantioselectivity of the biocatalysts (selected based on the similarity of their zymophore) also allude to the often overlooked contribution of non-active site residues to catalysis. These distal residues could coordinate breaching motions across the entire enzyme structure which may direct the malleability of the active site and modulate substrate accommodation. This has already been speculated within this class of enzymes, with the discovery that active site residues alone cannot discriminate closely related ammonia lyase and aminomutase enzymes in plants. Although protein dynamics are known to influence enzyme activity, there is little evidence of their consideration in biocatalyst or biotherapeutic research and development. Thus, this method of sampling sequence diversity outside the active site of an enzyme could be a crucial step in the investigation of these phenomena. The varying overall sequence-dynamics relationships of each enzyme may be the reason why, for example, PbPAL shows excellent activity with the novel substrate m-methoxycinnamate, whereas BIPAL, which has an identical active site, shows none. These underinvestigated considerations of protein
dynamics for biocatalysis (and also enzyme replacement therapy) will hopefully benefit from dedicated investigation of these phenomena in future work.

Compounds with electron-donating ring-substituents have been considered difficult for PALs, presumably due to deactivation of the \( \alpha \)-carbon as a result of the relative strengths of the aromatic ring and carboxyl group of these substrates as opposing electron sinks. This theory is thought to explain the conversely high activity of substrates (such as nitrocinnamates) with other ammonia lyases\(^3\)\(^6\)\(^,\)\(^5\). The discovery of PbPAL emphasises the difficulty with which such an activity would have been arrived upon using current engineering techniques without targeting active site residues, especially in the absence of a high-throughput enzyme assay. In this respect, our discovery method offers many advantages over traditional diversification across an entire parent sequence, which often involves lengthy and iterative mutagenesis yielding many poorly active variants, few useful hits and inherent problems with screening\(^6\)\(^,\)\(^5\). As such, we believe our method provides a complementary strategy to currently available techniques, particularly for selecting specific catalytic activities from functionally diverse families, such as other amino acid degrading enzymes, imine reductases/reductive aminases, esterases/hydrolases and nitri-lases/nitrile hydratases.

**Methods**

**X-ray crystallography.** AvPAL-Y78F-C503S-C565S was purified and crystallized as described previously for other AvPAL variants\(^3\). Diffraction data was processed and the structure solved by molecular replacement of the AvPAL-C503S-C565S structure (PDB ID: 3CZO) as a search model. The crystal structure of AvPAL-Y78F-C503S-C565S bound to cinnamate was then deposited in the protein data bank (PDB ID: 5LTM).

**Sequence alignments.** Potential PAL enzymes were identified from within the Universal Protein Resource KnowledgeBase (UniProtKB) with the basic local alignment search tool (BLAST) as provided by the UniProt website. All multiple amino acid sequence alignments were performed through use of the ClustalW2 programme using standard parameters as available online.

**Whole cell biocatalyst preparation.** The codon-optimized open reading frames (ORFs) encoding MxPAL and DdPAL and wild-type sequences for BlPAL and SrPAL were subcloned into a standard pET-28a or pET-28b vector by use of relevant multiple cloning sites. The pET-28b-PbPAL construct was used as provided by Prozomix Ltd. from their initial ammonia lyase panel screening efforts. Transformed *E. coli* BL21 (DE3) strains were grown in 800 mL LB-based autoinduction medium supplemented with kanamycin (50 \( \mu \text{g mL}^{-1} \)) at 18 °C (250 rpm agitation) for 4 days. Cells were harvested via centrifugation and stored at \(-20^\circ\)C until further use.

**Whole cell biotransformations.** The whole cell formulation was used directly as stored by resuspending in an appropriate volume of unadjusted 4 M ammonium carbamate (pH ~9.9) supplemented with the relevant arylacrylic acid 1a-p at the required concentration (10 mM for screening, 50 mM for preparative applications). All biotransformations were performed at 37 °C with agitation of 250 rpm. After a 24 h incubation period, a sample was subjected to centrifugation (3 min, 13000 rpm) to remove the catalyst before evaporation of the volatile reaction medium. The crude isolate was then dissolved in an appropriate solvent for analysis by HPLC.

**References**

1. Schnoes, A. M., Brown, S. D., Dodevski, I. & Babbit, P. C. Annotation error in public databases: misannotation of molecular function in enzyme superfamilies. *PLoS Comput. Biol.*, 5, e1000605 (2009).
2. Steinkellner, G. et al. Identification of promiscuous ene-reductase activity by mining structural databases using active site constellations. *Nat. Commun.*, 5, 4150 (2014).
3. Ferrario, V. et al. BioGPS descriptors for rational engineering of enzyme promiscuity and structure based bioinformatic analysis. *PLoS One* **9**, e109354 (2014).

4. Porter, C. T., Bartlett, G. J. & Thornton, J. M. The Catalytic Site Atlas: a resource of catalytic sites and residues identified in enzymes using structural data. *Nucleic Acids Res.* **32**, D129–33 (2004).

5. Sehnal, D., Pravda, L., Svobodová Váreková, R., Jonescu, C.-M. & Kocía, J. PatternQuery: web application for fast detection of biomacromolecular structural patterns in the entire Protein Data Bank. *Nucleic Acids Res.* **43**, W383–8 (2015).

6. Höhne, M., Schätzle, S., Jochems, H., Robins, K. & Bornscheuer, U. T. Rational assignment of key motifs for function guides in silico enzyme identification. *Nat. Chem. Biol.* **6**, 807–13 (2010).

7. Turner, N. J. Directed evolution drives the next generation of biocatalysts. *Nat. Chem. Biol.* **5**, 567–73 (2009).

8. Romero, P. A. & Arnold, F. H. Exploring protein fitness landscapes by directed evolution. *Nat. Rev. Mol. Cell Biol.* **10**, 866–76 (2009).

9. Parmeggiani, F., Weise, N. J., Ahmed, S. T. & Turner, N. J. Synthetic and therapeutic applications of ammonia-lyases and aminotransferases. *Chem. Rev.*, in press, https://doi.org/10.1021/acs.chemrev.8b00242.

10. Poppe, L. Methylidine-imidazolone: a novel electrophile for substrate activation. *Curr. Opin. Chem. Biol.* **5**, 512–524 (2001).

11. Strom, S., Wanninayake, U., Ratnayake, N. D., Walker, K. D. & Geiger, J. H. Insights into the mechanistic pathway of the Pantoaea agglomerans phenylalanine aminotransferase. *Angew. Chem. Int. Ed. Engl.* **51**, 2898–902 (2012).

12. Pillask, S., Tomin, A., Rétey, J. & Poppe, L. The essential tyrosine-containing loop conformation and the role of the C-terminal multi-helix region in eukaryotic phenylalanine aminotransferase. *FEBS J.* **273**, 1094–1019 (2006).

13. Zhang, X. & Liu, C.-J. Multifaceted regulations of gateway enzyme phenylalanine ammonia-lyase in the biosynthesis of phenylpropanoids. *Mol. Plant* **8**, 17–27 (2015).

14. Hill, A. M., Thompson, B. L., Harris, J. P. & Segret, R. Investigation of the early stages in soraphen A biosynthesis. *Chem. Commun. (Camb)*. **44**, 1358–9 (2008).

15. Xiang, L. & Moore, B. S. Inactivation, complementation, and heterologous expression of encP, a novel bacterial phenylalanine ammonia-lyase gene. *J. Biol. Chem.* **277**, 32505–9 (2002).

16. Williams, J. S., Thomas, M. & Clarke, D. J. The gene stlA encodes a phenylalanine ammonia-lyase that is involved in the production of a stiffle antibiotic in *Photobacterium luminescens* TT01. *Microbiology* **151**, 2543–50 (2005).

17. Benzeanson, B. S., Desaty, D., Emes, A. V. & Vining, L. C. Biosynthesis of Cinnamamido and Detection of Phenylalanine Ammonia Lyase in Streptomyces verticillus. *Can. J. Microbiol.* **16**, 147–151 (1970).

18. Sarkissian, C. N. & Gámez, A. Phenylalanine ammonia lyase, enzyme substitution therapy for phenylketonuria, where are we now? *Molecular Genetics and Metabolism* **86** (2005).

19. Ikeya, K. et al. Phenylalanine ammonia-lyase modified with polyethylene glycol: potential therapeutic agent for phenylketonuria. *Amino Acids* **29**, 283–287 (2005).

20. Gámez, A. et al. Development of pegylated forms of recombinant Rhodospiridium toruloides phenylalanine-ammonia-lyase for the treatment of classical phenylketonuria. *Mol. Ther.* **11**, 986–989 (2005).

21. Cantor, J. R., Panayiotou, V., Aghnello, G., Georgiou, G. & Stone, E. M. Engineering reduced-immunogenicity enzymes for amino acid deprivation therapy in cancer. *Methods in enzymology* **502**, (Elsevier Inc., 2012).

22. Pelayo, B. A., Fu, Y. M. & Meadows, G. G. Inhibition of B16BL6 melanoma invasion by tyrosine and phenylalanine deprivation is associated with decreased secretion of plasminogen activators and increased plasminogen activator inhibitors. *Clin. Exp. Metastasis* **17**, 841–848 (1999).

23. Abell, C. W., Stith, W. I. & Hodgins, D. S. The Effects of Phenylalanine Ammonia-Lyase on Leukemic Lymphocytes in *Vitro*. *Cancer Res.* **32**, 285–290 (1972).

24. Stith, W. J., Hodgins, D. S. & Abell, C. W. Effects of Phenylalanine Ammonia-Lyase and Phenylalanine Deprivation on Murine Leukemic Lymphoblasts in *Vitro*. *Cancer Res.* **33**, 966–971 (1973).

25. Huisman, G. W., Agard, N. J., Mijts, B., Vroom, J. & Zhang, X. Engineered phenylalanine ammonia lyase polypeptides (2015).

26. Vellard, M. C. et al. Compositions Of Prokaryotic Phenylalanine Ammonia Lyase And Methods Of Treating Cancer Using Compositions Thereof (2009).

27. Eigtved, P. & Groth Clausen, I. Stabilised Phenylalanine Ammonia Lyase. (1998).

28. Weise, N. J. et al. Asymmetric Synthesis of (S)-2-Indolacrylic Acid by Combining Biocatalysis and Homogeneous Catalysis. *ChemCatChem* **3**, 289–292 (2011).

29. de Lange, B. et al. β-regioselective asymmetric amination through Biocatalytic Asymmetric Amination and Palladium-Catalyzed Arylation. *ACS Catal.* **5**, 512–524 (2015).

30. Lovelock, S. L. & Turner, N. J. Bacterial Anaabaena variabilis phenylalanine ammonia lyase: a biocatalyst with broad substrate specificity. *Bioorg. Med. Chem.* **22**, 5557–55 (2014).

31. Parmeggiani, F., Lovelock, S. L., Weise, N. J., Ahmed, S. T. & Turner, N. J. Synthetic of D- and L-Phenylalanine Lyases: A Multienzymatic Cascade Process. *Angew. Chem. Int. Ed. Engl.* **54**, 4608–4611 (2015).

32. Emiliiani, G., Fondi, M., Fani, R. & Gribaldo, S. A horizontal gene transfer at the origin of phenylpropanoid metabolism: a key adaptation of plants to land. *Biol. Direct* **4**, 7 (2009).

33. Wang, L. et al. Structural and Biochemical Characterization of the Therapeutic A. variabilis Phenylalanine Ammonia Lyase. *J. Mol. Biol.* **380**, 623–635 (2009).

34. Heberling, M. et al. In vitro and in vivo characterization of a novel tyrosine ammonia-lyase encoding gene involved in bagremycins biosynthesis in Streptomyces sp. *Biotechnol. Lett.* **34**, 269–74 (2012).

35. Kovacs, K. et al. Expression and Properties of the Highly Alkalophilic Phenylalanine Ammonia-Lyase of Thermophilic Rubrobacter xylophilus. *PLoS One* **9**, e85943 (2014).

36. Jendresen, C. B. et al. Highly Active and Specific Tyrosine Ammonia-Lyases from Diverse Origins Enable Enhanced Production of Aromatic Compounds in Bacteria and Saccharomycetes cerevisiae. *Appl. Environ. Microbiol.* **81**, 4458–76 (2015).
46. Theodore, C. M. et al. Genomic and metabolomic insights into the natural product biosynthetic diversity of a feral-hog-associated Brevibacillus laterosporus strain. PLoS One 9, e90124 (2014).
47. Brock, Da, Douglas, T. E., Queller, D. C. & Strassmann, J. E. Primitive agriculture in a social amoeba. Nature 469, 393–6 (2011).
48. Kamerlin, S. C. L. & Warshel, A. At the Dawn of the 21st Century: Is Dynamics the Missing Link. Proteins 78, 1339–1375 (2011).
49. Hay, S. & Scrutton, N. S. Good vibrations in enzyme-catalysed reactions. Nat. Chem. 4, 161–8 (2012).
50. Lovelock, S. L., Lloyd, R. C. & Turner, N. J. Phenylalanine Ammonia Lyase Catalyzed Synthesis of Amino Acids by an MIO-Cofactor Independent Pathway. Angew. Chem. Int. Ed. 53, 4652–4656 (2014).
51. Mimoto, T. et al. Structure–activity and structure–metabolism relationships of HIV protease inhibitors containing the 3-hydroxy-2-methylbenzoyl-allophenylnorstatine structure. Bioorg. Med. Chem. 12, 281–293 (2004).
52. Hu, Y., Leng, Y., Duan, S., & Huang, S. Aza (oxa)-cyclopentane-2-ketone (thione) compounds, medicine compositions thereof, preparation method and applications (2014).
53. Wybenga, G. G. et al. Structural Investigations into the Stereochemistry and Activity of a Phenylalanine-2,3-aminomutase from Taxus chinensis. Biochemistry 53, 3187–3198 (2014).
54. Bornscheuer, U. T. et al. Engineering the third wave of biocatalysis. Nature 485, 185–94 (2012).

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Author Contributions
N.J.W., N.J.T. and S.J.C. conceived the project and N.J.W. was additionally responsible for the visual inspection of the structural data, design of the annotation system and identification of all target sequences used (under the supervision of N.J.T. and S.J.C.). S.T.A. and F.P. worked on enzyme preparations, biotransformations and analytics. S.T.A. and J.L.G. performed molecular biology procedures associated with gene cloning and construct preparations. M.S.D. and D.L. undertook all structural analyses and molecular visualisations.

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