Identification, Characterization, and *In Silico* Analysis of New Imine Reductases From Native *Streptomyces* Genomes

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The development of biocatalytic tools for the synthesis of optically pure amines has been the focus of abundant research in recent years. Among other enzymes, imine reductases have attracted much attention associated with the possibility of attaining chiral secondary amines. Furthermore, the reductive aminase activity associated with some of these enzymes has facilitated the production of optically pure amines from a prochiral ketone, a transformation that opens doors to an incredible array of products. In this work, the genomes from native *Streptomyces* strains isolated in our lab have been explored on the search for novel imine reductases. Application of different structural criteria and sequence motif filters allowed the identification of two novel enzymes, Ss-IRED_S and Ss-IRED_R. While the former presented outstanding activity towards bulky cyclic imine substrates, the latter presented reductive aminase activity with the assayed ketones. A bioinformatic analysis based on modeling and docking studies was performed in order to explain the differences in enzyme activity, searching for additional criteria that could be used to analyze enzyme candidates *in silico*, providing additional tools for enzyme selection for a particular application. Our findings suggest that imine reductase activity could be predicted by this analysis, overall accounting for the number of docking positions that meet the catalytic requirements.

**Keywords:** imine reductases, reductive aminases, *Streptomyces*, docking, bioinformatic analysis

**INTRODUCTION**

Synthesis of chiral amines is of particular interest for synthetic organic chemistry due to its presence in many biologically active molecules. Different methodologies and strategies have been developed for amine synthesis, and the biocatalytic toolbox of enzymes (i.e., transaminases, amine dehydrogenases, monoamine oxidases, and imine reductases (IREDs)) has become of interest to prepare these compounds in a more effective and stereoselective manner (France et al., 2017; Cosgrove et al., 2018; Devine et al., 2018; Höhne 2019; Adams et al., 2019).

IREDs are NADPH-dependent oxidoreductases able to catalyze the asymmetric reduction of different imines and iminium ions to produce the corresponding secondary and tertiary amines (Cosgrove et al., 2018; Höhne 2019). These enzymes can be found in different biosynthetic pathways...
and were discovered by Mitsukura et al. (2011) in *Streptomyces* sp. (Mitsukura et al., 2013). Over the last years, several reports of IREDs have been published; noticeably, gene selection was entirely based on sequence analysis by alignments (France et al., 2017; Lenz et al., 2017; Matzel, Gand, and Höhne 2017; Devine et al., 2018; Höhne 2019; Montgomery et al., 2020; Yao et al., 2021). In 2017, an IRED homologue capable of performing reductive amination (RedAm) was found in *Aspergillus oryzae* (Aleku et al., 2017). IRED and RedAm activities are similar, but their mechanisms are slightly different (Rodríguez-Mata et al., 2013; Man et al., 2015; Sharma et al., 2018). Due to the convenience of this reaction, which allows the synthesis of secondary amines directly from ketones, efforts have been made to expand the repertoire of enzymes capable of performing reductive amination (Kohls, Steffen-Munsberg, and Höhne 2014; Schrittweiser, Velikogne, and Krottul 2015; Wetzl et al., 2016; Aleku et al., 2017; Maugeri and Rother 2017; France et al., 2018). An interesting article published by Turner et al. made a characterization of 80 putative and 15 previously described IREDs across 10 different transformations and confirmed that reductive amination catalysis is not limited to any particular subgroup or sequence motif (Montgomery et al., 2020).

Nowadays, most of IRED and RedAm are not suitable for industrial applications mainly due to their limited substrate scope, low activity towards bulky substrates, the need of an excess of the amine nucleophile, and enzyme instability under process conditions. Nevertheless, there are a few recent examples of success in the application of these enzymes to industrial processes, all of which required a previous improvement of the enzymes’ performance using intensive and process focused enzyme engineering (Schober et al., 2019; Duan et al., 2021; Kumar et al., 2021). Other protein engineering approaches have been explored to overcome the limitations, and different biocatalysts have already been used in preparative scale biotransformations and enzymatic cascade reactions (France et al., 2017; Lenz et al., 2017; Matzel, Gand, and Höhne 2017; Devine et al., 2018; Höhne 2019; Yao et al., 2021). A deeper understanding of structure–activity relationship for these enzymes is needed for further improvement, as well as the development of alternative ways for uncovering new enzymes.

As sequence databases are growing exponentially, enzyme discovery using bioinformatic approaches is increasingly appealing. Through sequence alignments, *in silico* studies of proteins allow the recognition of conserved motifs among different enzymes of the same family, providing an interesting way to explore new proteins for a desired activity and to get deeper insights into sequence–function relationships (Höhne et al., 2010; Iglesias, Paniza, and Rodriguez 2017). Provided that adequate templates are available, homology modeling can be used to build proper models of protein structures from its sequence (Catucci et al., 2016; Fademrecht et al., 2016; Velikogne et al., 2018). Moreover, *de novo* accurate prediction of protein structure is becoming feasible (Senior et al., 2020; 2019). Docking experiments can be of major help in predicting substrate binding and understanding the role of different amino acids present in the active site (Bommarius, Blum, and Abrahamson 2011; Sirin et al., 2014; Catucci et al., 2016; Han et al., 2017; Montgomery et al., 2020). Deeper *in silico* analysis involves the use of quantum mechanics/molecular mechanics approach, but the high computational cost of these methods is often a limitation for its application (Rinaldi et al., 2018). There are already some reports on *in silico* studies for IREDs, and specific databases of their sequences have been constructed (Fademrecht et al., 2016; Velikogne et al., 2018).

As already mentioned, the first IREDs were identified from *Streptomyces* strains. This genus has been largely known for its diversity on secondary metabolites; thus, the probability of finding novel IREDs among strains of this genus is large (Spasic et al., 2018). In our lab, Prof. Pianzolla’s group has built a collection of over 200 *Streptomyces* strains isolated and characterized from potato tubers and soil samples from Uruguay (Lapaz et al., 2017; Croce et al., 2021). Recently, the whole-genome sequencing of the representative strains of this collection was reported (Lapaz et al., 2019). In this work, we decided to explore the available genomes from this Uruguayan *Streptomyces* collection on the search for novel enzymes with IRED or RedAm activity. Two new IRED candidates with potential IRED or reductive aminase activity were selected based on structural motifs. Furthermore, *in silico* analysis using homology models refined by molecular dynamics simulation and docking studies proved useful to explain the catalytic activity of these enzymes and could provide a valuable tool to select novel enzymes with IRED or RedAm activity.

**MATERIALS AND METHODS**

**Strains and Growth Conditions**

*Escherichia coli* BL21(DE3) was acquired from New England BioLabs (Ipswich, MA, USA) and was cultivated in Luria–Bertani (LB) broth or on LB agar plates at 37°C. Recombinant strains *E. coli* BL21(DE3) (pET-28b(+)-Ss-IRED_R) and *E. coli* BL21(DE3) (pET-28b(+)-Ss-IRED_S) were grown in LB broth or LB agar plates, supplemented with the appropriate antibiotic (kanamycin 50 µg/ml). Autoinduction media LB Base including trace elements was used for Ss-IRED_R and Ss-IRED_S protein production. Recombinant strains of *E. coli* BL21(DE3) coexpressing an IRED and *Lb*-ADH required a second antibiotic (ampicillin 50 µg/ml) as well anhydrotetracycline (200 µg/ml) for protein production.

Ss-IRED_R (National Center for Biotechnology Information (NCBI) ref. seq. WP_112474446.1) was identified from *Streptomyces* sp. strain ST1020 (BioSample: SAMN04977949). Ss-IRED_S (NCBI ref. seq. WP_119590418.1) was identified from *Streptomyces scabies* strain ST129 (BioSample: SAMN07731257).

**Chemicals and Molecular Biology Reagents**

Biotransformation substrate 1a was kindly provided by Prof. Ignacio Carrera; substrates 1b–1d were generously donated by Prof. Nicholas Turner; substrates 3–5 and a were purchased from Sigma Aldrich (St Louis, MO, USA). All the biotransformation substrates were used without further purification. Racemic amines used as analytical standards
were purchased from Sigma Aldrich (St. Louis, MO, USA), while chiral amines were produced from the prochiral imines or ketones through biotransformations with reported biocatalysts: 2a was produced with IRED-K; 2b was produced with IRED-D and K (Velikogne et al., 2018); 2c was produced with Ao-IRED (Aleku et al., 2016) and GF3546-IRED (Mitsukura et al., 2013); and 4a was produced with ApRedAm (Aleku et al., 2017).

Molecular biology reagents were purchased from New England Biolabs (Ipswich, MA, USA) and Thermo Fisher Scientific (Waltham, MA, USA). Expression vector pET-28b(+) was purchased from Novagen (Darmstadt, Germany) and was used for gene expression. Fourteen pET-28a(+) IRED-containing plasmids (IRED-A to IRED-N) and plasmid pASK-IBA5plus were very generously donated by Dr. Jörg H. Schüttwieser (Velikogne et al., 2018).

**DNA Manipulation and Ss-IRED_S Cloning**

Standard procedures were used for DNA manipulation, and genomic DNA from *Streptomyces* were extracted according to Sambrook *Molecular Cloning: A Laboratory Manual* (Sambrook 2001). Plasmid DNA was purified using commercial chromatography kits from Thermo Fisher Scientific (Waltham, MA, USA) and QIAGEN (Hilden, Germany); silica columns from these kits were regenerated and reused following the procedure described by Siddappa (Siddappa et al., 2007), while standard protocols were modified to obtain greater yields of DNA (Pronobis, Deuitch, and Peiifer 2016; Wood et al., 2017). Restriction enzymes and thermostable polymerases were used according to the manufacturers’ instructions. PCR amplifications were performed in a GeneAMP PCR system 2400 (PerkinElmer, Waltham, MA, USA) using adequate cycling periods, and DNA samples were routinely analyzed by agarose gel electrophoresis (Sambrook 2001). Nucleic acid concentration and purity were measured using a NanoDrop® ND-1000 spectrophotometer. Transformations of electrocompetent cells were performed on a BioRad MicroPulser™ Electroporator following manufacturer protocols.

An optimized sequence for *E. coli* of Ss-IRED_R (provisional ID Bankit 2503810) was synthesized and cloned in pET-28b(+) by GenScript (Piscataway, NJ, USA). The resulting vector was transformed into *E. coli* BL21(DE3) for protein expression, yielding the recombinant strain *E. coli* BL21(DE3) (pET-28b(+)-Ss-IRED_R). Amplification of Ss-IRED_S was performed by PCR with primers IRED Ndel Fw (GCTCATATGCCGGACACATCGCAG) and IRED EcoRI Rv (CTAGAATTCGACGAGGGGAACCCCCGAC). Ss-IRED_S gene was then cloned into pGEM-T Easy vector (Promega, WI, USA), and digestion with Ndel and EcoRI allowed subcloning into pET-28b(+) in frame with the N-terminal His6 tag. The obtained vector was purified by gel electrophoresis using the Freeze ‘N Squeeze kit from Bio-Rad (Hercules, CA, USA) and transformed into *E. coli* BL21(DE3), yielding the recombinant strain *E. coli* BL21(DE3) (pET-28b(+)-Ss-IRED_S).

**Whole Cell Biotransformations With Strains Expressing Ss-IRED_R and Ss-IRED_S**

Fresh plates of *E. coli* BL21(DE3) (pET-28b(+)-Ss-IRED_R) and *E. coli* BL21(DE3) (pET-28b(+)-Ss-IRED_S) were streaked from frozen stocks, and a single colony was used to inoculate 5 ml of LB-Kan. The culture was incubated in a rotary shaker overnight (150 rpm, 37°C), and 1 ml of this culture was used to inoculate 100 ml of fresh autoinduction media LB Base including trace elements and kanamycin. This culture was incubated in a rotary shaker at 150 rpm for about 2.5 h at 37°C until an OD600 nm ≈ 1; and then it was grown at 150 rpm and 28°C for 48 h. The cells were collected by centrifugation at 4,000 × g and 4°C for 15 min. The pellet was washed three times with 50 ml of Tris buffer (100 mM, pH 7.5). *E. coli* cells measuring 50 mg (wet weight) were resuspended in 465 μl of the same buffer; glucose and the appropriate substrate were added to a final concentration of 2% and 5 mM, respectively. For RedAm reactions, 50 eq. of propargylamine was used. The reaction mixture was incubated at 28°C and 150 rpm for 48 h. Reactions were quenched by addition of 10 M of NaOH (100 μl). The mixture was extracted with ethyl acetate (750 μl), and the organic layer was separated by centrifugation (5 min, 12,000 × g). The organic phase was dried with Na2SO4, and methyl heptadecanoate (2.5 mM) was added as internal standard. Percent conversion and enantiomeric excess were determined as established in the Gas Chromatography Analysis section.

**Whole Cell Biotransformations With Strains Coexpressing an Imine Reductase and Lb-ADH**

Fresh plates of *E. coli* BL21(DE3) coexpressing an IRED and Lb-ADH were streaked from frozen stocks, and a single colony was used to inoculate 5 ml of LB-Kan + Amp. The culture was incubated in a rotary shaker overnight (150 rpm, 37°C), and 1 ml of this culture was used to inoculate 100 ml of fresh autoinduction media LB Base including trace elements and the appropriate antibiotics. This culture was incubated in a rotary shaker at 150 rpm for about 2.5 h at 37°C until an OD600 nm ≈ 1; ADH production was induced by the addition of anhydrotetracycline. Then, it was grown at 150 rpm and 28°C for 48 h. The cells were collected by centrifugation at 4,000 × g and 4°C for 15 min. The pellet was washed three times with 50 ml of Tris buffer (100 mM, pH 7.5). *E. coli* cells measuring 50 mg (wet weight) were resuspended in 465 μl of the same buffer; glucose and the appropriate substrate were added to a final concentration of 2% and 5 mM, respectively. The reaction mixture was incubated at 28°C and 150 rpm for 48 h. Reactions were quenched by addition of 10 M of NaOH (100 μl). The mixture was extracted with ethyl acetate (750 μl), and the organic layer was separated by centrifugation (5 min, 12,000 × g). The organic phase was dried with Na2SO4, and methyl heptadecanoate (2.5 mM) was added as internal standard. Percent conversion and enantiomeric excess were determined as established in the Gas Chromatography Analysis section.
Gas Chromatography Analysis
Biotransformation products were analyzed on a GC-2010 Shimadzu (Kyoto, Japan) gas chromatograph equipped with a FID detector. Carrier gas, H2; makeup gas, N2. Columns MEGA-SE52 (achiral SP, 30 m × 0.25 mm × 0.25 μm) and MEGA-DEX ASX-1 (chiral SP, 25 m × 0.25 mm × 0.25 μm) from MEGA (Milan, Italy) were used for determination of conversion and enantioemic excess, respectively. Particular analytical conditions and retention times for each amine can be found in Supporting Information, Supplementary Tables S1, S2.

Bioinformatic Analysis
YASARA’s homology modeling experiments, version 21.8.27, were carried out with five templates and three PSI-BLAST iterations in template search; the remaining parameters were set by default (Krieger and Vriend 2014). The templates were initially prepared by adding bond orders and charges; water molecules were removed, and hydrogens were added accounting for protonation states of all ionizable residues at pH 7.5. IREDs from A. oryzae (PDB 5G6R), A. terreus (PDB 5OJL), Bacillus cereus (PDB 4D3F), and Amycolatopsis orientalis (PDB 5A9T); and RedAm from A. terreus (PDB 6EOD) were used as structural templates for modeling Ss-IRED_R, Ss-IRED_S, IRED-A, B, C, D, F, I, J, K, L, M, and N. To aid alignment correction and loop modeling, a secondary structure prediction for the target sequence was obtained. This was achieved by running PSI-BLAST to create a target sequence profile and feeding it to the PSI-Pred secondary structure prediction algorithm (Jones 1999). A total of 25 independent models were created, and YASARA combined the best parts of the 25 models to obtain a hybrid model. In each homology model using YASARA software, we assigned bond orders and protonation patterns according to pH 7.4 (Krieger et al., 2012).

Refinement by short molecular dynamics simulations was performed to improve structures’ quality: with the use of YASARA’s force field YAMBER3 (Krieger et al., 2004), 500 ps of molecular dynamics simulation was conducted at 298 K and pH 7.4. The simulation was performed in explicit solvent with TIP3P water model, density 0.997. Based on the lowest energy, the final representative structures were considered for further study. Ligands were first prepared using YASARA’s molecular modeling software (Krieger, Koraimann, and Vriend 2002). Solvation of the ligands prior to minimization was carried out in an explicit solvent shell at pH 7.4 and 0.9% NaCl. Each ligand was subjected to energy minimization to refine the bond lengths and bond angles. The minimized conformation with the lowest energy for each ligand was used in the docking experiments. Docking was performed by VINA software (Trott and Olson 2019) using default parameters. However, the number of VINA runs was increased from 10 to 250. Size of the docking box was set to 5 Å around the amino acids in the active site in each structure. Positions with a root mean square deviation (RMSD) lower than 2.5 Å were clustered together, and each cluster was represented by the position with lowest free energy of binding. Every cluster obtained by VINA for each substrate used in imine reduction was analyzed, and parameters such as energy of binding, distance for hydride transfer from NADPH to C≡N double bond, and interactions between nitrogen of imine with surrounding possible proton donor amino acids His, Glu, Asp, Ser, Tyr, and Cys were evaluated.

RESULTS
Sequence Alignments and Analysis of Primary Structure
We used BLASTp of NCBI (Altschul et al., 1997) to compare the amino acidic sequences of different R- and S-IREDs previously reported (Mitsukura et al., 2010, 2013; France et al., 2018). Through the alignment of twelve sequences, conserved regions formed by 5 and 13 amino acids were identified, and from these, two were selected as templates for new searches: M1: VWNRT and M2: YxDGAI[ML]AxPx2IG. Sequenced genomes from our collection of native Streptomyces strains were analyzed using Artemis software (Carver et al., 2012), distinguishing nine sequences that presented the abovementioned motifs. These nine potential IREDs were then analyzed using additional criteria reported by Fademrecht et al. (2016) and France et al. (2017). The former characterized the presence of two protein motifs (GLGxMGx5[ATS]x4Gx4[VIL] WNR[TSL][x2][KR] and Gx[DE]x[GDA]x[APS]x3 [K]x[ASL]x [LMVIAG]) corresponding respectively to NADPH and substrate binding for IREDs. Additionally, it established the importance of amino acids in positions 139 and 194 for determining enzyme stereoselectivity (Fademrecht et al., 2016). Besides, we analyzed the presence of active site residues associated with reductive aminase activity (France et al., 2018). After these analyses, two potential enzymes with complementary stereoselectivity were selected: Ss-IRED_S from S. scabies ST129 and Ss-IRED_R from Streptomyces sp. ST1020. The selected enzymes were cloned in pET-28b(+) and expressed in E. coli BL21(DE3).

Characterization of Ss-IRED_S and Ss-IRED_R Biocatalytic Activity
IRED activity for the identified enzymes was assessed using a set of substrates 1a–1d (Figure 1) encompassing both monocyclic and bulky prochiral imines. The assays were performed using resting cells under two different conditions that favor NADPH recycling: addition of glucose as co-substrate and coexpression of Lb-ADH. As can be observed in Table 1, the coexpression of ADH for NADPH recycling resulted in the best conversions as it was earlier reported by Velikogne and coworkers (Velikogne et al., 2018). Surprisingly, Ss-IRED_R presented no activity towards these substrates, despite the sequence alignment analysis that suggested IRED activity. On the other hand, Ss-IRED_S exhibited excellent activity towards substrates 1a–1c with superb enantiomeric excess, showing preference for bulky substrates such as those with dihydroisoquinoline and tetrahydro-β-carboline motifs, and a preference for six-membered rings over five-membered rings for monocyclic imines.

To study reductive amination activity, substrates 3–5 were tested using propargylamine (a) as amine donor (Figure 2). For these reactions, we found that recombinant Lb-ADH reduced the ketone; thus, the assays were only performed with glucose as co-substrate for NADPH regeneration. Following that procedure, no
FIGURE 1 | Imines 1 and corresponding amines 2 evaluated in this study.

TABLE 1 | Characterization of the imine reductase activity: synthesis of chiral amines from the corresponding imine with Ss-IRED_S and Ss-IRED_R.

| Substrate | Whole cell system Ss-IRED_S | Whole cell system Ss-IRED_R |
|-----------|-----------------------------|-----------------------------|
|           | % Conv (glucose) | % Conv (ADH system) | % ee | % Conv (glucose) | % Conv (ADH system) | % ee |
| 1a        | 10               | >99                     | >99 (S) | 0               | 0               | nd   |
| 1b        | 57               | >99                     | >99 (S) | 0               | 0               | nd   |
| 1c        | 27               | 85                      | 98 (R)  | <1              | <3              | nd   |
| 1d        | <2               | <5                      | nd      | 0               | 0               | nd   |

FIGURE 2 | Ketones (3–5), propargylamine (a), and corresponding amines (3a–5a) evaluated in this study.

TABLE 2 | Characterization of the reductive amination activity: synthesis of chiral amines from the corresponding ketone and amine donor propargylamine with Ss-IRED_S and Ss-IRED_R.

| Substrate | Whole cell system Ss-IRED_S | Whole cell system Ss-IRED_R |
|-----------|-----------------------------|-----------------------------|
|           | % Conv (Glucose) | % ee | % Conv (Glucose) | % ee |
| 3         | 60               | -     | >99               | -     |
| 4         | 0               | -     | 0                 | -     |
| 5         | 0               | -     | 40                | 98 (na) |

Note: na, absolute stereochemistry could not be assigned due to lack of true optically pure standards.
ketone reduction was detected. Table 2 illustrates the experimental results obtained. As it can be observed, Ss-IRED_S showed moderate activity for substrate 3 and no activity with the other substrates. In contrast, Ss-IRED_R presented good reductive amination activity with substrates 3 and 5, this last one with excellent enantiomeric excess. Despite many efforts on assaying the reductive amination of phenyl ketone (4), no activity was detected for this substrate.

**Homology Models and Docking Analysis**

Imine reduction and reductive amination have been observed with highly diverse sequences, which indicates the importance of analyzing the three-dimensional structure and the characteristics of the active site over that of the linear sequence. In silico analysis was performed for the new enzymes to better understand observed preferences for IRED or RedAm activities, as well as differences in substrate specificity.

Homology models for Ss-IRED_R (Supplementary Figure S1), Ss-IRED_S (Supplementary Figure S2), and 10 IREDs used as controls in complex with NADPH cofactor were constructed using the “closed” form with NADPH of IREDs in PDB because analysis of the (apo-) and (NADPH-bound) subunits within the structure revealed a rotation of 14° (Sharma et al., 2018). Docking studies of substrates 1a–1d and 3–5 and a in the different enzymes’ active site were performed by Autodock-VINA. After clustering of 250 runs of each substrate with each enzyme, the different ligand–protein interactions in the active site were analyzed (Figure 3). To analyze IRED activity, substrates 1a–1d in the different enzymes were selected for docking, and different parameters such as energy of binding, distance for hydride transfer from NADPH to C=N double bond, and distance between nitrogen of imine with surrounding acidic amino acids were evaluated. Adequate distance between the C4 atom of the NADPH and the carbon in the C=N double bond (dC–C4) is needed for hydride delivery.
Additionally, mechanistic studies have suggested the importance of an acidic amino acid at a proper distance to the forming amine to act directly or through a water molecule as a proton donor (Man et al., 2015).

Favorable docking positions were selected as those presenting a d_{C,C4} distance minor to 7.5 Å, and a possible proton donor amino acid at a distance minor to 7 Å. Previous reports have indicated Tyr 169 in GF3546-IRED from *Streptomyces* sp. or Asp 187 in Q1EQE0 IRED from *Streptomyces kanamyceticus* as candidates for protonating the forming amine (Rodríguez-Mata et al., 2013; Man et al., 2015). The presence of these residues was first analyzed, but the presence of other acidic amino acids at an adequate distance was equally considered to support catalytic activity (Ribéiro et al., 2020). A similar analysis was performed on a group of five IREDS (IREDs*) whose activity was previously reported using the same substrates in order to compare the different parameters (Velikogne et al., 2018). The results are presented in Table 3.

As can be seen in Table 3, the results of this in silico study indicate that no significant difference in energy of binding is detected when considering all the docked positions or only favorable ones, despite evidence of excellent activity associated with these enzymes (columns 1 and 2). Regarding the d_{C,C4} distance, it could be observed that selected positions show an average value of 5 Å, as well as those for the control group. Surprisingly, Ss-IRED_R presented the shortest average distance (~4.5 Å) with all substrates even though no IRED activity was detected with this enzyme. A correlation between d_{C,C4} distance and activity cannot be clearly established from these data.

Interestingly, the percentage of docked positions complying with catalytic requirements correlates well with experimental activity. A quantitative analysis of docking positions for substrates 1a and 1b in Ss-IRED_S yields respectively 46% and 73% of favorable positions that meet the catalytic requirements. It is noteworthy that a similar quantitative percentage of positions is observed for the IREDS in control group. Concurrently, for the inactive Ss-IRED_R, only 10% of the positions of substrate 1a and 11% of 1b comply with the expected attributes for catalysis. Additionally, for substrate 1d, IREDS in the control group show an average of 58% of positions presenting an adequate distance to NADPH and the required amino acid for amine protonation. In agreement with these data, Ss-IRED_S and Ss-IRED_R with no detectable activity presented respectively only 0.5% and 31% of positions complying with catalytic requirements. The results for substrate 1c are inconsistent at first sight; however, a deeper analysis allowed us to determine that for the inactive Ss-IRED_R, most docked positions in clusters do not interact with Asp176 but with a different possible proton donor. On the contrary, Ss-IRED_S presents interaction with Asp188 in most positions in the selected clusters, which could explain the difference in activity (Supplementary Table S3).

A similar docking analysis to explain reductive aminase results was investigated; however, no correlation could be established in this case (data not shown). Therefore, we based our assessment of Ss-IRED_S and Ss-IRED_R activity on the work reported by France et al. (2017). The presence of the described critical conserved amino acids in the active site was examined based on an alignment of our enzymes with the best reductive aminases reported by these authors (Supplementary Table S7). We noticed that Ss-IRED_R presents the three amino acids that interact directly with the main functional groups of our substrates via hydrogen bonding, while it also presents the other three conserved amino acids to which no specific function has been assigned. In contrast, Ss-IRED_S only has two of the three conserved amino acids that interact directly with the substrates.

**DISCUSSION**

Two new IREDS, Ss-IRED_S and Ss-IRED_R, have been identified from native *Streptomyces* strains isolated from Uruguayan soil. The strategy followed for identification was based on the conserved motifs M1: VWNRT and M2: YxDGAI[ML]AxPx2IG and proved successful since both enzymes presented IRED or reductive aminase activity. This approach could yield more diverse enzymes than procedures based strictly on BLAST or homology search, overall when imine reduction and reductive amination have been observed with highly diverse sequences, indicating the importance of three-dimensional structure and the characteristics of the active site (Marshall et al., 2020; Montgomery et al., 2020). A feature that is observed in these structures is their highly conserved NADPH binding site. When the stereoselectivity is analyzed in Ss-IRED_S and Ss-IRED_R, prediction by the presence of key residues identified by Fademrecht and co-workers seems to work well (Fademrecht et al., 2016,) although the origin of IRED selectivity is complex and not fully understood.

A recent study by Velikogne and co-workers classifies a selection of published and novel IREDS as either “D-type” or “Y-type” based on phylogenetic tree analysis and whether they have an aspartic acid or a tyrosine residue aligned with position 187 in *S. kanamyceticus* Q1EQE0 IRED (Velikogne et al., 2018). The catalytic profile of D-type enzymes was characterized with several substrates, presenting particularly poor results with substrates 1a and 1b. A detail to remark from our results is that both of our IREDS are D-types enzymes; however, our Ss-IRED_S present excellent conversion with sterically demanding imines 1a and 1b (Table 1). Additionally, our enzymes presented poor activity with 1d, which proved to be a good substrate for most D-type enzymes. Considering both substrate specificity and stereoselectivity, Ss-IRED_S behaves like a Y-type enzyme. Although the association of structural data with enzyme activity is really important, it has remained elusive for IRED activity, overall when based in one amino acid and exceptions are found.

The results presented in Table 1 clearly demonstrate that coexpression of a cofactor regeneration system for NADPH recycling is essential for good IRED activity with whole cell biocatalysts. ADH-based system reported by Velikogne et al. is extremely efficient for imine reduction (Velikogne et al., 2018); however, its own biocatalytic activity hampered its use for reductive aminations since it reduced the ketone. It would be highly desirable to devise a similar system for NADPH recycling that avoids ketone reduction.

Various authors reported excellent results for reductive amination with purified enzymes (Aleku et al., 2017; Matzel, Gand, and Höhne 2017; Ramsden et al., 2019; Yao et al., 2021);
however, only one previous work focused on the applicability of whole cell biocatalysts for this reaction (Maugeri and Rother 2017). In a screening of 15 IRED systems, Maugeri et al. reported only one system with 99% of conversion with cyclohexanone as a substrate. Table 2 shows that the whole cell system expressing Ss-IRED_R can be used for reductive amination with excellent results, and no ketone reduction by E. coli native dehydrogenases was detected. The reductive amination of phenylacetone (4) and 4-phenyl-2-butanone (5) with propargylamine (a) has proven a challenging transformation even for isolated RedAm, and few literature reports are available (Aleku et al., 2017; Marshall et al., 2020). The activity and stereoselectivity presented by Ss-IRED_R with substrate 5 situate this enzyme as a good platform for targeted evolution. The absence of activity with substrate 4 is intriguing; however, the conjugated π-system in this substrate could favor the enolate form, which could be further stabilized by residues in the active site impeding reductive amination due to steric or electronic issues.

With the aim of contributing to the analysis of possible correlations among catalytic activity and structural features, we considered performing an in silico analysis of our enzymes compared with previously reported ones. Among the available computational approaches, we chose to perform homology modeling and docking analysis, since it does not demand high levels of computing power and could be used to explain experimental data. The docking studies performed in this work confirmed the complexity of the active sites of enzymes with IRED-RedAm activities. Considering the energy of binding, values obtained with Autodock-VINA are not enough to ensure the activity of an enzyme with a tested substrate, and a deeper analysis using molecular dynamics and free energy calculations could provide a better though more time-consuming approach. Furthermore, the values obtained with the applied methodology presented a large dispersion, and no significant correlation between catalytic activity and energy of binding could be recognized. Nevertheless, it is worth noticing that when other restrictions are applied, such as the catalytic requirements used here (distance C–C4 and interaction with proton donor amino acids interaction of nitrogen of imine), the results obtained seem to correlate with the observed catalytic activity.

Even though IRED’s active site residues show significant variation, making it challenging to clearly define the characteristics of this enzyme family or to predict the properties of uncharacterized homologues, it is undeniable that residues structurally aligned to positions 169 or 187 play a key role either as proton donors or as anchors of the substrate in an optimal position. Experimental and crystallographic data showed that residue Asp187 in SkIRED (IRED-R type) or Tyr169 in BcIRED and NhIRED (IRED-S type) played an important role in activity, since mutation of these residues resulted in an inactive enzyme (Rodriguez-Mata et al., 2013; Man et al., 2015). Nevertheless, some discrepancy arose from other studies in which similar enzyme mutants maintain relative activity (Scheller et al., 2014; Hussain et al., 2015; Aleku et al., 2017); however, it should be noted that other residues can also serve as proton donors allowing catalysis (Aleku et al., 2016). In our in silico analysis, we have considered both possibilities, and the presence of an acidic amino acid at an adequate distance for proton transfer was established as a catalytic requirement. This criterion, taken together with the evaluation of the distance for hydride transfer from NADPH to C==N double bond, proved to correlate well in most cases with biocatalytic activity.

As a summary, the use of bioinformatic analysis based on sequence and structural features proved useful for the identification of novel IREDs. Furthermore, the criteria established by Fademrecht et al. remain interesting for prediction of stereoselectivity, and the evaluation of amino acids in the active site proposed by France et al. correlates well for predicting RedAm activity. The application of these criteria helped us in selecting two IRED candidates from native Streptomyces genomes. The described Ss-IRED-S constitutes an interesting addition to the biocatalytic toolbox due to its excellent activity and enantioselectivity for the reduction of bulky imines, while Ss-IRED-R adds to the repertoire of reductive aminases. Modeling each enzyme candidate and performing docking studies as those presented in this work could help in predicting substrate specificity. This proved useful for IREDs; however, due to a more complex mechanism (Sharma et al., 2018), additional work should be done in order to establish a similar procedure for reductive amination activity. Despite this, our results show that considering the percent of docking positions complying with catalytic requirements could be a useful new addition to the in silico search for novel enzymes with IRED activity, avoiding testing of large enzyme libraries and facilitating prediction of enzyme activity.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Materials. Further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

CI performed most of the bioinformatic analysis, homology modeling and docking studies, and analysis of correlation among bioinformatic and experimental data. AT performed work on Artemis for IRED identification, and GL and PP deepened the analysis by performing and analyzing alignments with other enzymes. AT and GL equally contributed with characterization of Ss-IRED_S and Ss-IRED_R by performing cloning, expression, and biotransformation experiments. Both developed analytical conditions for GC analysis of the different compounds. ML isolated Streptomyces genomes and guided work with Artemis software. MP constructed and provided the Streptomyces culture collection. GL performed Ss-IRED_R modeling and docking studies to explain RedAm activity guided by CI and PP. CI, PP, and SR made substantial contributions to the conception and design of the research; however, all authors were involved in the acquisition, analysis, and interpretation of data used in the work. CI, AT, PP, and SR
had a major contribution on writing the present manuscript; GL
has also been involved in manuscript preparation, critical
revision, and approval of the final version. All the authors
agree to be accountable for all aspects of the work in ensuring
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