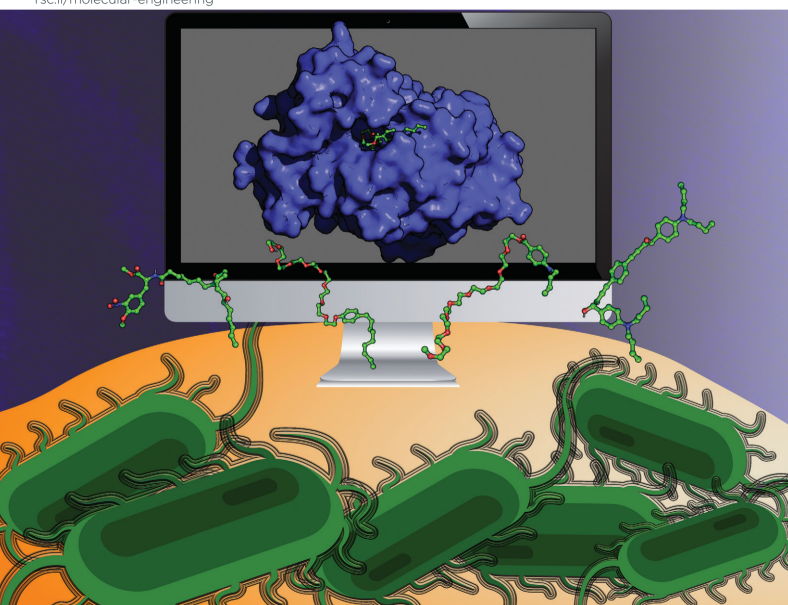
Volume 9 Number 1 January 2024 Pages 1-142

# MSDE

Molecular Systems Design & Engineering

rsc.li/molecular-engineering



ISSN 2058-9689



#### **PAPER**

Sérgio F. Sousa *et al. In silico* identification of novel PqsD inhibitors: promising molecules for quorum sensing interference in *Pseudomonas aeruginosa* 



# **MSDE**



# **PAPER**

View Article Online



Cite this: Mol. Syst. Des. Eng., 2024, 9, 7

# In silico identification of novel PqsD inhibitors: promising molecules for quorum sensing interference in Pseudomonas aeruginosa†

Tatiana F. Vieira, (1) ab Nuno M. F. S. A. Cerqueira, (1) ab Manuel Simões (D° and Sérgio F. Sousa (D\*ab

PgsD is an anthraniloyl-CoA anthraniloyltransferase involved in the synthesis of the secondary metabolites essential to the formation of Pseudomonas quinolone signal (PQS) inducer molecules. Its main substrate is anthraniloyl-coenzyme A (ACoA) but it can accept malonyl-CoA as secondary substrate. Suppression of PqsD activity has been connected to the inhibition of biofilm formation and can also be a good target for dual inhibition, when combined with PgsR inhibition. Here we describe the validation and application of an in silico methodology to find new compounds to inhibit PqsD. Using molecular docking and structurebased virtual screening protocols, five databases of compounds were screened (FDA approved subset of the ZINC database, Chimiothèque Nationale, Mu.Ta.Lig. Virtual Chemotheca, Interbioscreen (IBS), and Comprehensive Marine Natural Products Database (CMNPD)), representing a total of 221146 molecules. The top five compounds of each database were selected to be further analysed using molecular dynamics simulations. Binding affinity was validated using free energy calculations, enabling the selection and characterization of eight compounds for future studies aiming to develop new quorum sensing inhibitors.

Received 29th June 2023, Accepted 25th September 2023

DOI: 10.1039/d3me00107e

rsc.li/molecular-engineering

#### Design, System, Application

This project employs a fully integrated multi-technique in silico optimization protocol that combines molecular docking (validated through re- and crossdocking experiments), development of a robust virtual screening procedure with enhanced ability in active ligand identification, application of this protocol to large libraries of chemically distinct potential molecules (a total of 221 146 molecules) from various origin and, finally, a thorough molecule selection stage. For selection, molecular dynamics simulations, free energy calculations, visual inspection and similarity analysis are combined to identify the most promising molecules. The eight selected ligands can now be experimentally tested and used for rational and target-specific drug design against PqsD, an anthraniloyl-CoA anthraniloyltransferase involved in the synthesis of the secondary metabolites essential to the formation of Pseudomonas quinolone signal (PQS), an important target for quorum sensing interference in Pseudomonas aeruginosa.

## Introduction

Pseudomonas aeruginosa is a Gram-negative opportunistic pathogen that exhibits a large genome, a highly adaptable nature and a plethora of regulatory systems.<sup>1</sup> These characteristics make the bacteria very difficult to eradicate, particularly in immunocompromised patients. Additionally, antimicrobial therapy is often hampered due to the presence of efflux pumps, β-lactamases and the ability of P. aeruginosa to start growing as sessile communities commonly called biofilms. The thick biofilm matrix composed of extracellular polymeric substances (EPS) is highly impermeable to antimicrobial agents. P. aeruginosa controls the expression of virulence factors and biofilm formation through a complex and cell dependent communication system called quorum sensing (QS). In this bacterium, there are three main QS systems that communicate with each other and are highly hierarchical: las, rhl systems that use homoserine lactones as signal molecules and the pqs system that uses quinolones as autoinducer. These QS inducer molecules,

<sup>&</sup>lt;sup>a</sup> UCIBIO/REQUIMTE, BioSIM, Departamento de Medicina, Faculdade de Medicina da Universidade do Porto, Alameda Prof. Hernâni Monteiro, 4200-319 Porto, Portugal. E-mail: sergiofsousa@med.up.pt

<sup>&</sup>lt;sup>b</sup> Associate Laboratory i4HB – Institute for Health and Bioeconomy, Faculdade de Medicina, Universidade do Porto, 4200-319 Porto, Portugal

<sup>&</sup>lt;sup>c</sup> LEPABE Laboratory for Process Engineering, Environment, Biotechnology and Energy, Faculty of Engineering, University of Porto, Rua Dr. Roberto Frias, s/n, 4200-465 Porto, Portugal

<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: https://doi.org/

**Paper** 

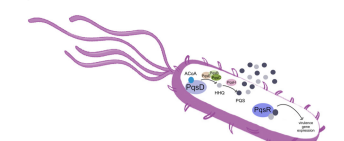


Fig. 1 Simplified overview of the PQS system in P. aeruginosa

accumulating in a growing bacterial population, start influencing the production and release of virulence factors.<sup>2</sup> A simplified overview of the pqs system is displayed in Fig. 1.

PqsD is an anthraniloyl-CoA anthraniloyltransferase involved in the synthesis of the secondary metabolites 2-heptyl-3-hydroxy-4(1*H*)-quinolone and 4-hydroxy-2(1H)quinolone, mediators in the formation of the pgs inducer molecules 2-heptyl-4-hydroxyquinoline (HHQ) and 2-heptyl-3hydroxy-4-quinolone (PQS).<sup>3</sup> It belongs to the β-ketoacyl-ACPsynthase family and is similar in structure to the β-ketoacyl-ACP-synthase III (FabH) in E. coli and M. tuberculosis.4 Its main substrate is anthraniloyl-coenzyme A (ACoA) but it can accept malonyl-CoA as secondary substrate. PgsD has also been shown to be responsible for the synthesis of 2,4-dihydroxyquinolone (DHQ), an abundant extracellular metabolite of P. aeruginosa.5,6

The PqsD binding pocket is long and narrow (15 Å) and can be divided into three parts: a positively charged entrance, a hydrophobic middle and a polar bottom delimited by the catalytic triad Cys112-His257-Asn287.4 Residues between Glu140 to Gly162 are key in CoA recognition and binding 6 and residues 186 to 222 form a L-shaped active side. A detailed view of the PqsD structure is presented in Fig. 2.

Suppression of PqsD activity has been shown to inhibit biofilm formation<sup>7</sup> and when combined with MvfR/PqsR inhibition can lead to more highly efficient attenuation of P. aeruginosa pathogenicity, in a phenomenon called dual inhibition.8 Furthermore, targeting the PQS system, highly specific to P. aeruginosa, can minimize the development of

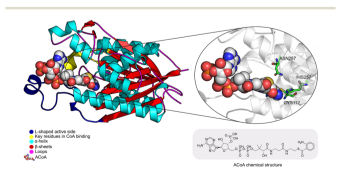


Fig. 2 Schematic representation of PqsD bound to ACoA (PDB: 3H77). Protein represented in cartoon, with key features highlighted with different colors, and ACoA represented by atom VdW radius (spheres).

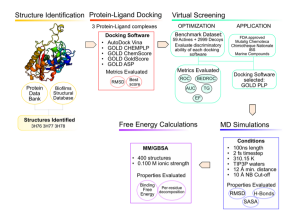


Fig. 3 In silico workflow used in the current study.

side effects by not interfering with other bacteria important for the health maintenance in humans.1

Some inhibitors for β-ketoacyl-ACP-synthase III (FabH) have been described, namely cerulenin, thiolactomycin, indole derivatives, Schiff bases, benzovlaminobenzoic acid derivatives and compounds containing sulfur atoms.9 Cerulenin and thiolactomycin also target other type II FAS enzymes. Cerulenin forms an adduct with the catalytic cysteine of FabF and thiolactomycin acts like a reversible inhibitor.10 Some of these have been tested against PqsD such as benzoylaminobenzoic acid derivatives and compounds containing sulfur atoms. Benzoylaminobenzoic acid derivatives have shown promising in vitro activity against PgsD. 11-13 More specifically regarding sulfonamide substituted 2-benzamidobenzoic acids, researchers realized that the presence of carboxylic acid was essential for activity and that the variation of substituents of the 2-aminobenzoic acid part could increase the potency of the possible PgsD inhibitors. 12 Furthermore, it has been suggested that there are only two possible inhibition modes of action; either by direct interaction between the inhibitor molecule and the residues in the binding site or simply by blocking the entrance of the PqsD pocket. 3,12,14

The use of computer aided drug design tools, namely protein-ligand docking and structured-based virtual screening (VS)15,16 has been shown to help narrow down the list of compounds to test experimentally, with promising results, 17 particularly regarding molecular docking in the PqsD binding pocket. 18,19 The workflow of the present study is presented in Fig. 3 and has been applied to other projects and other bacterial targets. 20-24

# Methodology

#### Structure analysis and preparation

Searching the Protein Data Bank<sup>25</sup> and the Biofilms Structural Database,<sup>26</sup> 3 X-ray structures of PqsD were identified. The PqsD model predicted by Alphafold (P20582)<sup>27,28</sup> and available at the Alphafold Protein Structure Database was also taken into consideration, to evaluate

**MSDE Paper** 

Table 1 Summary of the X-ray structures available for PgsD in the Protein Data Bank<sup>23</sup> and Biofilms Structural Database<sup>24</sup>

PDB CODE	Resolution/Å	Mutation	Ligand	Type of ligand	Year of deposition	Ref.
3H76 3H77 3H78	1.80 1.80 1.70	No (Cys112 covalently bound to anthranilate) Cys112Ala	— Anthraniloyl-coenzyme A 2-Aminobenzoic acid	— Natural substrate Product	2009	6

possible structural variations but it was not used for the in silico study because the priority was given to choosing an experimental structure with a crystallographic ligand in the binding pocket.

Structure 3H76 represents the structure in the apo form. Structure 3H77 presents the Cys112 residue covalently bound to anthranilate, a reaction intermediate. Structure 3H78 has one mutation (Cys112Ala) to evaluate the role of Cys112 in the catalytic mechanism of PqsD. Also, this structure has anthranilic acid in the binding pocket. The presence of alanine does not affect the binding of ACoA but affects the kinetic activity of the enzyme, since in the experimental assays with this structure, there was no product detected.<sup>6</sup> Further details on these X-ray structures are shown in Table 1.

For target preparation, polar hydrogens were added with GOLD at physiological pH and subject to visual inspection of the amino acid residues at the binding pocket. Waters and molecules resulting from the crystallographic process, were removed. The three structures were aligned to ensure consistency in the docking coordinates. The crystallographic ligands were extracted and saved in separate files, to be used in the validation of the docking conditions. For GOLD, all the targets were saved in .mol2 file format, and for Vina in .pdbqt file format.

## Protein-ligand docking protocol validation

Two docking software alternatives and five scoring functions were used for this project. Autodock Vina,<sup>29</sup> and GOLD<sup>30</sup> (CHEMPLP, GoldScore, ChemScore and ASP scoring functions). To ensure consistency, consistent docking conditions were applied to all the SF, in terms of binding pocket definition.

Testing a variety of scoring functions (SF) is intended to evaluate which is the best for this specific target with a long, narrow pocket, since docking results have been shown to differ significantly depending on the specific target protein and ligand type. 31,32

Re-docking and cross-docking were the methods used for protocol validation. Re-docking is used to evaluate the ability of each SF and protocol to correctly reproduce the geometry and orientation of the respective crystallographic ligand pose. By evaluating the binding affinity and the root mean square deviation (RMSD) between the heavy atoms of the crystallographic and docked poses, one can evaluate the quality of the protocol in correctly identifying the experimental pose. Cross-docking evaluates the universality of the protocol, evaluating the ability of each experimental

structure of the target protein in accommodating the binding of the different confirmed binders. It involves the docking of all the X-ray ligand structures into all the protein structures available. In this case, only one crystallographic ligand (ACoA) was extracted using pymol, 33 the protonation was corrected using OpenBabel34 and was subsequently docked into the three PDB PqsD structures. The purpose of this test is to determine if individual X-ray structures can accommodate the different X-ray ligands, which are co-crystallized in other X-ray structures. The binding affinity was evaluated to determine the quality and accuracy of the method.

#### Virtual screening protocol validation

For the validation of the VS conditions, an active versus decoys protocol was applied. The goal was to evaluate the performance of each scoring function (SF) and protocol in discriminating between binders and non-binders. The ideal SF would provide the true binders with better scores, being able to clearly single out binders from non-binders. However, due to the impossibility to mathematically account for all the phenomena that occurs in protein-ligand binding, that is not always the case. Since the goal of a VS is to quickly screen large databases of compounds, the accuracy is somewhat penalized.35,36 Therefore, the protocol was continuously improved to maximize the ability of each SF to rank the true binders in the top solutions.

The ChEMBL<sup>37</sup> and BindingDB<sup>38</sup> databases were explored, and 59 ligands were found to have reported experimental activity against PqsD.39,40 The DUD-E41 database was used to generate decoy molecules. Decoys are molecules that show similar physical properties to the active molecules, to eliminate bias, but have distinct chemical properties and are assumed to be non-binders. For each active molecules, 50 decoys were generated, giving a total of 2950 decoys. The complete test set was composed of a total of 3009 molecules, prepared for docking with OpenBabel.34

The results obtained with the four scoring functions applied were evaluated through a web-based application, Screening Explorer, 42 as well as using Excel. Several metrics were considered: enrichment factor at 1%, receiver-operating characteristic (ROC) curves and the respective area under the curve (AUC), Boltzmann-enhanced discrimination of ROC (BEDROC), the robust initial enhancement (RIE), and total gain (TG). A SF shows a good discriminatory ability and reproducibility if it presents a TG value over 0.25, combined with an AUC over 50%.

#### Virtual screening of compound databases

Five libraries of compounds were screened, to narrow the search for new inhibitors, using the docking and VS conditions that were previously optimized and validated. The selected databases were: FDA approved compounds<sup>43</sup> for drug repurposing, the French National chemical library (Chimiothèque Nationale)44 and Mu.Ta.Lig. Chemotheca<sup>45</sup> as databases of synthetic compounds, Interbioscreen (IBS)<sup>46</sup> and Marine Compound Database (CMNPD)47 and as sources of natural and marine compounds, respectively. The rational of screening various compound databases is to provide a more representative and diverse test set, containing synthesizable and natural compounds. At this stage, the crystallographic structure and SF used were 3H76 and ChemPLP, respectively.

The FDA approved library of compounds is subset of the ZINC database,43 a free database that contains over 230 million commercially available compounds. At the time of the VS experiments, the FDA-approved drugs dataset had 3207 compounds that were all docked against PgsD.

Chimiothèque Nationale<sup>44</sup> contains more than 70 000 compound that were synthesized or isolated in different laboratories. For this project, a total of 61 640 compounds were screened against PgsD.

The Mu.Ta.Lig. Virtual Chemotheca<sup>45</sup> was the result of the European Union COST action (CA15135) and it contains molecules that were synthesized by several research groups. For this assay, 64 804 compounds were screened against PgsD.

The natural compounds database of Interbioscreen (IBS)<sup>46</sup> containing 66 278 compounds was selected to find possible inhibitors for PqsD. This database includes over 68 000 natural compounds and derivatives from several ecosystems around the world.

Finally, 25 217 compounds retrieved form the Comprehensive Marine Natural Products Database (CMNPD)47 were docked against PqsD. This database contains manually curated chemical, physicochemical, and pharmacokinetic properties of marine natural compounds.

Prior to the virtual screening application, all the databases were verified and "cleaned". Duplicate molecules were eliminated, as well as compounds with molecular weight above  $700 \text{ g mol}^{-1}$ .

A similarity search was conducted using the SkelSpheres descriptor of Datawarrior, for the best predicted compounds retrieved from the VS. This descriptor encodes circular spheres of atoms and bonds into a hashed binary fingerprint

of 512 bits. From every atom in the molecule, DataWarrior constructs fragments of increasing size by including n layers of atom neighbours (n = 1 to 5). These circular fragments are canonicalized considering aromaticity and stereochemistry, counts duplicate fragments, in addition encodes hetero-atom depleted skeletons. From the canonical representation a hash code is generated, which is used to set the respective bit of the fingerprint. It has a good resolution, leading to less hash collisions and it is the most accurate descriptor for calculating similarities of chemical graphs.

#### Molecular dynamics and free energy calculations

To validate the docking predictions, molecular dynamics (MD) simulations were performed with PqsD and the top 5 compounds of each database. In the MD simulations PqsD was treated with the ff14SB force field48 and the 25 compounds selected were parameterized with Gaussian16 (ref. 49) using ANTECHAMBER and the general Amber force field (GAFF)50 to calculate RESP HF/6-31G(d) charges. The protein-ligand complexes were then treated with the Leap module of Amber18.51 ACoA, the natural substrate, was used as a control molecule.

All the PqsD-ligand systems were neutralized with the addition of sodium ions and placed into TIP3P water molecules box; with periodic boundary conditions whose edges were at least at a 12 Å distance from each proteinligand complex atom. A time step of 2 fs was used, and the SHAKE algorithm was applied to constrain the bonds involving hydrogen atoms. The long-range electrostatic interactions were calculated with the particle mesh Ewald summation method and the cut-off value for electrostatic and Lennard-Jones interactions was set at 10.0 Å.

Four minimization steps were applied to remove clashes, followed by two equilibration steps and a final production run. The minimization steps were applied to the following groups of atoms: first minimization-water molecules (2500 steps); second minimization-hydrogens atoms (2500 steps); third minimization-chains of all the amino acid residues (2500 steps); forth minimization-full system (10000 steps). The two 50 ps equilibration steps consisted of: 1-heating of the system to 298 K using a Langevin thermostat at constant volume (NVT ensemble) 2-equilibration of the density of the system at 298 K. Lastly, the 100 ns production run was performed in an NPT ensemble with a temperature of 298 K and 1 bar pressure. VMD52 and the cpptraj tool53 were employed to explore the resulting molecular dynamics (MD) trajectories.

Table 2 Re-docking and cross-docking results for ACoA

	ASP	ChemScore	GoldScore	ChemPLP	Vina (kcal mol <sup>-1</sup> )
3H76	47.62	32.12	96.76	91.89	-6.70
3H77	49.34	33.43	89.66	87.31	-5.00
3H78	42.18	29.01	105.03	90.65	-7.50
Average per SF	46.38	28.85	97.15	89.95	-6.40

**MSDE** 

The end-point free energy method MM-GBSA<sup>54</sup> (molecular mechanics-generalized Born surface area method) was employed to assess the binding free energies of each ligand towards PqsD. The MM/PBSA.py script, available in AMBER,<sup>55</sup> was applied to calculate the energies, with a salt concentration of 0.100 mol dm<sup>-3</sup>. The final last 90 ns of the MD simulation of every complex with an interval of 10 frames were considered for the analysis, representing a total of 1800 frames per complex. The free energy decomposition option was used to obtain information about the contribution of each residue to the total free energy. The results are presented from the strongest binder to the weakest. This method involves several approximations, and the entropic term is the one that displays the highest statistical uncertainty.<sup>56</sup> This method is very useful to provide insight and analyze the binding affinities and trends, as well as helping in narrowing down the selection of compounds for experimental testing.

## Results and discussion

Table 2 shows the results of the re-docking and cross-docking for all the PqsD structures available in the PDB database. As expected, structure 3H77 showed the lowest scores due to the presence of anthranilate bound to Cys112As previously mentioned, structure 3H78 has a mutation (Cys112Ala), that is not catalytically active in vivo, however, since alanine is a small residue, roughly the same size as a cysteine, it does not cause a steric hindrance in the binding pocket and in the cross-docking with ACoA it produces a high score, which is particularly evident when using GoldScore.

Regarding the all the SF studied, ChemScore performed the worst and was unable to correctly dock the ACoA in the binding site.

Even though structure 3H78 presented good docking scores it was discarded at this stage. Since the mutation of 3H78 is in the binding pocket, and in one of the residues of the catalytic triad (leading to a form that is not catalytically active in vivo), it could influence the docking results leading to a selection of compounds, later on, in the VS stage, that could not have affinity toward wt PqsD.

After the initial study of structure 3H77, the docking scores obtained were not satisfactory because there was an anthranilate bound to Cys112. Even after removal of the anthranilate group, the results did not improve (data not shown), and it was decided to also remove this structure from the VS studies.

Table 3 Virtual screening results

	EF 1%	AUC%	$\frac{3H76}{TG}$	RIE	BEDROC
ASP	0.00	71	0.02	1.25	0.09
ChemScore	0.00	71	0.02	0.98	0.06
GoldScore	1.70	67	0.25	3.24	0.20
ChemPLP	1.73	68	0.25	1.65	0.20

Ultimately, the structure chosen to move on to the VS optimization stage was 3H76, the crystallographic structure of the enzyme in the apo form. At this stage, Vina was also left out because it did not provide accurate results in terms of score and crystallographic pose reproducibility.

The actives vs. decoys protocol was performed on structure 3H76 and the metrics used to evaluate its quality are depicted in Table 3 and Fig. 4. The enrichment factor at 1% (EF 1%) measures how many actives are found at 1% of the ordered database, so, naturally, the higher the EF at 1% the better. Additionally, a good discriminatory ability of the SFs, is verified by TG and ROC values above 0.25 and 50% respectively.42 GoldScore and CHEMPLP were the SFs that exhibited the best metrics, with CHEMPLP showing a slight improvement in the ability to find true binders early on. Even though ASP and ChemScore showed higher AUC, the early recognition metrics (EF1% and TG) were not satisfactory. Eventually, GoldScore and ChemPLP provided similar results (particularly in the VS metrics), ultimately GoldScore was discarded because it is much more time-consuming and ChemPLP was able to provide good results much faster.

The top 1% of compounds from each database were selected based on the docking score generated by the application of the VS protocol using the PLP scoring function.

The ChemPLP score and molecular weight of the top eight compounds predicted in the FDA approved database are depicted in Table 4, along with a small description of the pharmaceutical use.

The chemical structure of the top compounds from the all the databases can be seen in Fig. 5. Most of the compounds selected present high molecular weights with many rotatable bonds, exhibiting long tails oriented toward the polar bottom of the binding pocket similarly to the natural inducer.

Interestingly three classes of compounds were highlighted: isoquinoline derivatives from Chemoteca, derivatives from IBS and imidazole derivatives from Marine Compounds. Interestingly, isoquinoline derivatives had already been studied as antimicrobial agents65,66 with promising activity against several species of bacteria and fungi. Chromone derivatives have shown activities against Mycobacterium tuberculosis, Staphylococcus aureus, and Bacillus subtilis<sup>67</sup> and have also appeared as top hits in another VS study regarding MvfR/PqsR.68 As for imidazole

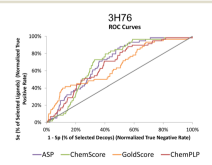


Fig. 4 ROC curves for 3H76 for the several scoring functions tested.

Table 4 Top eight hits from the FDA-approved database

Drug name	Description	ChemPLP score	MW (g mol <sup>-1</sup> )
Tessalon	A non-narcotic oral antitussive agent <sup>57</sup>	93.5	603.7
Nonoxynol-9 (N-9)	Vaginal spermicide used for the non-hormonal contraception in conjunction with other modes of contraception <sup>58</sup>	94.5	616.8
Polidocanol	Sclerosing agent used for the treatment of uncomplicated spider and reticular veins in the lower extremities <sup>59</sup>	84.7	585.8
Deferoxamine (Dfo)	Chelating agent used to treat iron or aluminum toxicity and some blood transfusion dependent anemias <sup>60</sup>	90.5	560.7
Phytonadione (vitamin K1 aka K-Ject)	Treat and manage vitamin K deficiency <sup>61</sup>	93.0	416.6
Salmeterol	Beta-2 adrenergic agonist used to treat and manage symptoms of asthma and chronic obstructive pulmonary disease (COPD) <sup>62</sup>	85.3	416.6
Nefazodone	Antidepressant used in the treatment of depression. However, it has been discontinued in some countries due to the risk of hepatotoxicity <sup>63</sup>	85.8	471.0
Ketoconazole	Broad spectrum imidazole antifungal used to treat seborrheic dermatitis and fungal skin infections $^{64}$	82.4	531.4

derivatives, their structure has already been proven to be a good starting point in the design of more efficient QS inhibitors.69 both Consequently, chromone

Fig. 5 Chemical structure of the selected compounds. Orange represents the compounds from the FDA approved database. Blue represents the molecules retrieved from Chimiothèque Nationale. Red corresponds to the ones coming from Mu.Ta.Lig Chemotheca. Represented in green are the compounds retrieved from the IBS and in black the compounds selected from CMNPD.

benzimidazole derivatives seem to be particularly good candidates for dual inhibition.

None of the selected compounds contains sulfur atoms, so the hypothesis of a nucleophilic attack in a similar way than ACoA it not likely. However, the selected ligands possess oxygen and nitrogen atoms capable of interacting with the sulfur atom of Cys112 or increasing the number of hydrogen bond interactions. This data was further analyzed using MD simulations and measuring the distance between these atoms and Cys112.

The similarity analysis performed shows that there is structural similarity between molecules in the same database the following compounds presenting high chemical similarity between them: Tessalon, N-9 and Polidocanol (from the FDA approved database); AB-00069970 and AB-00069855 (from the Chimiotheque database); CMLDID26439 and CMLDID51411 (from the Mu.Ta.Lig Chemotheca database); STOCK1N-00436 and STOCK1N-45295 (from the IBS database) (Fig. S1 in the ESI†). There is, however, no chemical similarity between any of the predicted molecules and the ACoA.

To further analyze and validate the docking predictions, molecular dynamics simulations were performed in the top 25 protein-ligand complexes. The results are shown in Table 5. ACoA was used as reference and the docked pose of each ligand was used as a starting point. Several parameters were considered to evaluate the stability of the protein-ligand complexes such as: protein and ligand RMSd and average number of hydrogen bonds formed. The exposure of the ligand to the solvent when bound to the protein was also evaluated measuring the solvent-accessible surface area (SASA), percentage of potential ligand SASA buried, and number of water molecules present within a radius of 3 Å from the pocket.

All the molecules remained bound to the protein throughout the simulations, however, because some of them are quite big and have long tails, leading to higher protein and ligand RMSDs in comparison with the initial structures. This might be explained because their movement in the pocket induces a displacement of the amino-acid residues of

**MSDE** 

the active site. Even though they are big molecules, they are buried deep into the pocket, with most compounds presenting a percentage of potential ligand SASA buried above 85%.

Calculations were also performed to determine binding energies of association and evaluate residues contributing most to ligand stabilization in binding pockets using MM/ GBSA. Results are presented in Table 6. For the MM/GBSA calculation, the entropic term was neglected, due to the limitations in the entropy calculation method, a common approximation. 54,70,71 Therefore, the MM/GBSA values will be looked as a ranking tool, to guide the selection of compounds to test experimentally. The overall energy of binding was decomposed, and the individual residue contribution was evaluated for each protein-ligand interaction. The three most relevant residues are also represented in Table 6. The potential inhibitors selected are highlighted in purple both in Tables 5 and 6.

To narrow down the list of compounds to be studied experimentally, a graph was plotted to correlate the Docking ChemPLP score with the MM/GBSA values (kcal mol<sup>-1</sup>) (Fig. 6). This graph allows for a better visualization and confirmation of the potential of these molecules as PqsD inhibitors. Ultimately, the compounds selected were the ones that presented higher docking scores and more negative binding free energies (compounds with docking scores above 90 and MM/GBSA values below -60 kcal mol<sup>-1</sup>: Tessalon, AB-00069812, AB-00028362, N-9, STOCK1N-25409, AB-00069970, AB-00063025 and Polidocanol). A normalized score was calculated (ChemPLP score by the number of heavy atoms), to eliminate some of the docking bias (as the docking score is proportional to the number of atoms) and the results are presented in Fig. S2 in the ESI.† The selected compounds exhibit higher normalized scores than the reference (ACoA) and the tendency in terms of compound selection is preserved, because, even though the compounds

Table 5 Average protein RMSD values (Å), average ligand RMSD (Å), average MvfR-ligand complex SASA (Å<sup>2</sup>), percentage of SASA for the buried ligand, average number of water molecules within a radius of 3 Å and average number of ligand hydrogen bonds obtained from the MD simulations of PqsDligand complexes

Source	Compound	Average RMSd (Å)	Ligand RMSd (Å)	Average complexed ligand SASA (Ų)	Potential ligand SASA buried (%)	Average number of H bonds	Waters (3 Å)
Ref.	ACoA (ref.)	2.7 ± 0.3	2.9 ± 0.3	193.8 ± 29.2	82	2.2 ± 1.3	2.1 ± 0.9
	Tessalon	2.6 ± 0.1	4.8 ± 0.5	118.3 ± 38.6	89	0.4 ± 0.6	2.2 ± 0.9
	N-9	2.8 ± 0.3	2.7 ± 0.4	247.9 ± 46.7	79	0.7 ± 0.7	2.5 ± 1.1
	Polidocanol	3.2 ± 0.4	4.1 ± 0.6	221.5 ± 63.8	80	0.5 ± 0.6	2.6 ± 1.1
504	Dfo	3.1 ± 0.4	2.8 ± 0.4	177.2 ± 35.5	82	1.6 ± 1.2	2.7 ± 1.1
FDA approved drugs	K-Ject	2.7 ± 0.3	2.3 ± 0.9	94.3 ± 34.3	89	0.6 ± 0.6	2.5 ± 1.1
	Ketoconazole	2.8 ± 0.3	1.5 ± 0.3	113.3 ± 26.7	85	0.1 ± 0.3	2.4 ± 1.1
	Salmeterol	3.2 ± 0.7	2.7 ± 0.5	89.65 ± 38.4	88	0.2 ± 0.4	2.5 ± 1.1
	Nefazodone	3.6 ± 0.4	1.9 ± 0.5	177.9 ± 49.7	77	0.6 ± 0.7	2.2 ± 0.9
	AB-00069812	3.1 ± 0.5	1.2 ± 0.2	101.5 ± 24.4	90	0.4 ± 0.5	2.6 ± 1.1
	AB-00028362	2.9 ± 0.3	2.8 ± 0.5	216.0 ± 42.1	81	0.6 ± 0.7	2.5 ± 1.1
Chimiotheque	AB-00063025	3.1 ± 0.4	2.3 ± 0.6	181.7 ± 38.9	83	$0.0 \pm 0.0$	2.5 ± 1.1
	AB-00069970	$3.1 \pm 0.5$	$1.4 \pm 0.2$	131.9 ± 28.9	87	$0.1 \pm 0.3$	2.5 ± 1.1
	AB-00069855	3.0 ± 0.5	3.8 ± 0.9	179.1 ± 52.9	82	0.2 ± 0.4	2.1 ± 0.9
	CMLDID26439	2.4 ± 0.3	1.2 ± 0.2	57.5 ± 13.4	92	0.9 ± 0.7	2.1 ± 0.9
	CMLDID51411	$3.6 \pm 0.8$	2.1 ± 0.7	108.4 ± 32.4	86	0.7 ± 0.7	2.4 ± 1.1
Mu.Ta.Lig Chemotheca	CMLDID31957	$3.1 \pm 0.4$	$2.1 \pm 0.3$	148.2 ± 54.5	81	$0.8 \pm 0.8$	2.4 ± 1.0
	CMLDID30539	4.9 ± 0.5	$1.2 \pm 0.3$	179.1 ± 26.8	77	0.5 ± 0.7	2.4 ± 1.0
	CMLDID36422	4.3 ± 1.0	2.1 ± 0.5	163.7 ± 48.7	78	0.6 ± 0.7	2.4 ± 1.1
	STOCK1N-25409	3.0 ± 0.5	2.6 ± 0.3	85.2 ± 29.2	91	1.0 ± 1.0	3.0 ± 1.3
	STOCK1N-25715	2.7 ± 0.2	2.3 ± 0.3	97.2 ± 27.7	90	0.3 ± 0.6	2.5 ± 1.1
IBS	STOCK1N-31694	$4.3 \pm 0.6$	$1.6 \pm 0.3$	101.9 ± 22.9	89	1.2 ± 0.8	2.2 ± 1.0
	STOCK1N-00436	2.7 ± 0.3	2.2 ± 0.6	155.8 ± 34.0	83	0.9 ± 0.7	2.6 ± 1.1
	STOCK1N-45295	2.7 ± 0.2	0.8 ± 0.2	72.9 ± 17.8	91	0.9 ± 0.8	2.2 ± 1.0
	CMNPD21891	2.9 ± 0.5	2.8 ± 0.4	92.9 ± 28.2	88	2.0 ± 1.1	2.0 ± 1.0
	CMNPD17128	2.6 ± 0.5	1.7 ± 0.3	77.8 ± 23.9	90	0.3 ± 0.5	2.2 ± 1.0
CMNPD	CMNPD417	3.4 ± 0.5	1.3 ± 0.2	83.5 ± 22.1	90	0.4 ± 0.6	2.5 ± 1.1
	CMNPD12771	2.7 ± 0.2	2.1 ± 0.3	119.3 ± 34.7	87	0.4 ± 0.6	2.6 ± 1.1
	CMNPD17131	$3.8 \pm 0.3$	2.9 ± 0.2	200.8 ± 39.3	76	0.1 ± 0.3	2.7 ± 1.1

	Compound	$\Delta G_{ m bind}$	Main contributors
Source		MM/GBSA (kcal mol <sup>-1</sup> )	MM/GBSA (kcal mol <sup>-1</sup> )
Ref.	ACoA (ref.)	-82.9 ± 0.5	$Arg262$ (-4.7 $\pm$ 1.1); $Asn260$ (-4.5 $\pm$ 1.1); $Arg36$ (-4.3 $\pm$ 1.2)
	Tessalon	-81.9 ± 0.3	Phe226 (-3.1 ± 0.6); His257 (-3.0 ± 0.7); Pro259 (-2.5 ± 0.6)
	N-9	-72.3 ± 0.3	Met194 (-3.1 ± 0.9); Ser317 (-2.8 ± 1.0); Leu193 (-2.6 ± 0.5)
	Polidocanol	-61.8 ± 0.4	Pro259 (-2.7 ±0.6); His257 (-2.3 ± 0.7); Met225 (-1.9 ± 0.6)
	Dfo	-55.9 ± 0.4	Gln111 (-3.6 $\pm$ 0.8); Asn260 (-2.1 $\pm$ 0.8); Leu155 (-2.0 $\pm$ 0.5)
FDA approved drugs	K-Ject	-52.2 ± 0.2	Leu193 (-2.3 $\pm$ 0.5); Phe226 (-1.8 $\pm$ 0.7); Met220 (-1.4 $\pm$ 0.5)
	Ketoconazole	-48.3 ± 0.3	Phe226 ( $-2.2 \pm 0.6$ ); Arg223 ( $-2.0 \pm 1.1$ ); Arg36 ( $-1.6 \pm 0.7$ )
	Salmeterol	-40.7 ± 0.4	Asn287 ( $-2.7 \pm 0.8$ ); Asn260 ( $-1.9 \pm 0.8$ ); Leu155 ( $-1.9 \pm 0.4$ )
	Nefazodone	-39.2 ± 0.3	Met220 (-1.7 ± 0.8); Arg36 (-1.5 ± 0.7); Ile263 (-1.5 ± 0.5)
	AB-00069812	-79.1 ± 0.2	Met225 (-2.4 ± 0.4); Phe226 (-2.3 ± 0.4); Leu193 (-2.3 ± 0.4)
	AB-00028362	-73.8 ± 0.3	Leu193 (-2.7 ± 0.6); Ala289 (-2.6 ± 0.9); Arg36 (-2.2 ± 1.0)
Chimiotheque	AB-00063025	-64.3 ± 0.2	Leu155 ( $-1.8 \pm 0.6$ ); Met225 ( $-1.5 \pm 0.3$ ); Leu193 ( $-1.3 \pm 0.4$ )
	AB-00069970	-63.6 ± 0.2	Leu193 ( $-2.1 \pm 0.5$ ); Met225 ( $-1.6 \pm 0.4$ ); Pro259 ( $-1.6 \pm 0.4$ )
	AB-00069855	-57.9 ± 0.3	Leu155 ( $-1.9 \pm 0.6$ ); Met220 ( $-1.7 \pm 0.7$ ); Phe226 ( $-1.8 \pm 0.7$ )
	CMLDID26439	-58.1 ± 0.2	Leu193 ( $-2.5 \pm 0.6$ ); Met225 ( $-2.2 \pm 0.3$ ); Asn260 ( $-1.9 \pm 0.5$ )
	CMLDID51411	-51.3 ± 0.3	Met220 ( $-2.7 \pm 1.0$ ); Phe226 ( $-2.2 \pm 0.9$ ); Pro259 ( $-1.9 \pm 0.5$ )
Mu.Ta.Lig Chemotheca	CMLDID31957	-43.8 ± 0.4	Arg223 (-4.7 ± 2.7); Arg36 (-3.4 ± 2.2); Met225 (-3.0 ± 0.8)
	CMLDID30539	-43.5 ± 0.3	Ser317 (-5.3 ± 1.5); Gly316 (-2.8 ± 0.9); Arg145 (-2.8 ± 1.9)
	CMLDID36422	-31.9 ± 0.3	Ser317 (-3.8 $\pm$ 2.2); Gly316 (-2.2 $\pm$ 0.7); Met220 (-2.0 $\pm$ 1.0)
	STOCK1N-25409	-71.3 ± 0.3	Asn260 (-2.7 $\pm$ 0.6); Phe226 (-2.3 $\pm$ 0.4); Arg36 (-2.3 $\pm$ 0.7)
	STOCK1N-25715	-59.6 ± 0.3	Leu193 ( $-2.4 \pm 1.0$ ); Asn260 ( $-2.4 \pm 1.2$ ); Phe226 ( $2.1 \pm 0.5$ )
IBS	STOCK1N-31694	-59.5 ± 0.2	Asn260 ( $-2.4 \pm 0.5$ ); Asn287 ( $-1.9 \pm 0.5$ ); Met225 ( $-1.8 \pm 0.3$ )
	STOCK1N-00436	-56.8 ± 0.2	Asn154 (-3.3 $\pm$ 0.6); Phe226 (-2.1 $\pm$ 0.6); Leu158 (-1.9 $\pm$ 0.4)
	STOCK1N-45295	-50.1 ± 0.2	Asn154 (-2.6 ± 0.6); Met225 (-1.8 ± 0.5); Asn260 (-1.7 ± 0.4)
	CMNPD21891	-56.9 ± 0.3	Gln111 (-6.8 ± 1.2); Ser317 (-5.4 ± 1.8); Phe226 (-2.4 ± 0.5)
	CMNPD17128	-55.8 ± 0.3	Asn287 (-2.2 ± 1.2); Leu155 (-2.0 ± 0.6); Leu193 (-1.7 ± 0.4)
CMNPD	CMNPD417	-53.3 ± 0.2	Leu193 ( $-1.7 \pm 0.3$ ); Met225 ( $-1.5 \pm 0.4$ ); Leu155 ( $-1.5 \pm 0.4$ )
	CMNPD12771	-53.2 ± 0.3	Leu193 (-2.4 ± 0.5); Asn260 (-2.2 ± 0.9); Leu155 (-2.0 ± 0.6)
	CMNPD17131	-44.1 ± 0.3	Phe218 ( $-2.4 \pm 1.1$ ); Phe226 ( $-1.9 \pm 0.5$ ); Met220 ( $-1.9 \pm 0.6$ )

present a slightly decrease normalized score, they still stand out in terms of a more favorable estimated binding free energy.

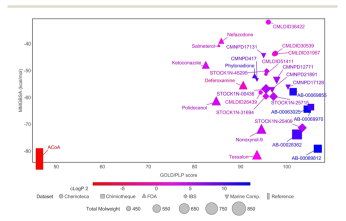


Fig. 6 Comparison of docking ChemPLP scores and total binding free energy (MM/GBSA). ACoA was used as reference. Blue compounds show higher log P values.

It is curious to see that the molecules that were chemically similar in the FDA approved database were all selected as the most promising ones (Tessalon, N-9 and Polidocanol). While the ones from Chimiotheque, Mu.Ta.Lig Chemotheca and IBS databases were not selected.

The interactions between the selected compounds and ACoA with PqsD is depicted in detail in Fig. 7 to 15. The ligand poses characterized are the most representative structures obtained from the MD simulations.

ACoA (Fig. 7) is stabilized through electrostatic interactions but can also form hydrogen bonds with Asp150, Gly222 and Asn260. It also forms salt bridges with Arg36 and Arg262. These arginines along with Arg223 seem to be important for ACoA recognition and interact with it through the diphosphate group. The covalent bond between ACoA and Cys112 is not visible because the methodology applied is not able to study the formation/ breaking of covalent bonds. However, the distance between the residue and ACoA is below 2 Å, and it was considered a good approximation.

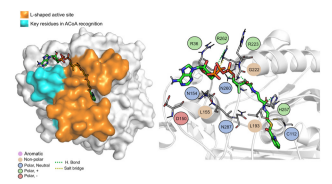


Fig. 7 ACoA bound to PqsD (green licorice). Surface view emphasizing the L-shaped active site and area with key residues involved in ACoA recognition (left). Interaction map highlighting the most relevant residues (right) involved in ligand stabilization.

Tessalon (Fig. 8) occupies almost the same area of ACoA, except for the interaction with Arg36 and Asp150 being in a more vertical position. It can form hydrogen bond interactions with residues His257 and Asn287, two of the residues of the catalytic triad, it is, however, very far away from Cys112 (~7.3 Å). It can also form a salt bridge with

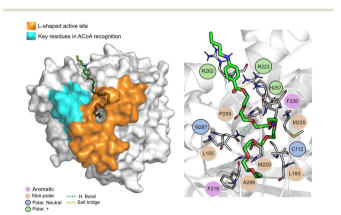


Fig. 8 Tessalon (green licorice) bound to PqsD. Interaction map highlighting the most relevant residues (right) involved in ligand stabilization.

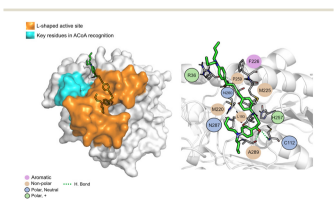


Fig. 9 AB-00028362 (green licorice) bound to PqsD. Interaction map highlighting the most relevant residues (right) involved in ligand stabilization.

Arg223. Tessalon caused a shift in the position of His257 and Asn287, when compared to ACoA.

AB00028362 (Fig. 9) can form hydrogen bonds with Cys112 (2.16 Å), Asn260 (2.07 Å) and Ala289 (1.67 Å), the other interactions are mainly hydrophobic. The key residues involved in ACoA recognition do not seem to interact with AB00028362.

AB00069812 (Fig. 10) interacts with PgsD mainly through electrostatic interactions with most of the residues and through hydrogen bond with Asn260. The aliphatic tails of AB00069812 seem to lock the compound into the L-shaped active site portion of the pocket.

N-9 (Fig. 11) is stabilized mainly through electrostatic interactions and can form a hydrogen bond with Asn87. The tail of N-9 seems to trigger a shift in the position of His257, Met225 and Phe226, when compared with ACoA-PqsD complex, leading to the formation of a tunnel to accommodate the aliphatic tail. Distance to Cys112 of 6.5  $\pm$  2.1 Å.

STOCK1N-25409 (Fig. 12) is stabilized by non-polar interactions but can also interact via hydrogen bond with Gly222, Asn260 and His257. It can also interact with Arg36 via salt bridge. It can interact, via electrostatic interactions, with

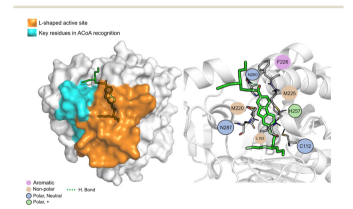


Fig. 10 AB00069812 (green licorice) bound to PqsD. Interaction map highlighting the most relevant residues (right) involved in ligand stabilization.

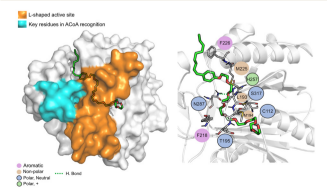


Fig. 11 N-9 (green licorice) bound to PqsD. Interaction map highlighting the most relevant residues (right) involved in ligand stabilization.

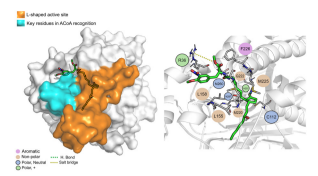


Fig. 12 STOCK1N-25409 (green licorice) bound to PasD. Interaction map highlighting the most relevant residues (right) involved in ligand stabilization

some of the key residues involved in ACoA recognition, namely Leu158 and Leu155. Distance to Cys112 is 11.5  $\pm$  1.1 Å.

AB-00063025 (Fig. 13) interacts with PqsD pocket, mainly through the establishment of non-polar interactions. The tricyclic head of this compound is in a position similar to the bicyclic head of ACoA when bound to PqsD, interacting with residues Asn154 and Leu155. However, it does not seem to interact with any of the residues of the catalytic triad (distance to Cys112 of 9.3  $\pm$  1.7 Å).

AB-00069970 (Fig. 14), can form hydrogen bond with Asn260 but interacts mainly through non-polar interactions. The aliphatic tail sits on top of the region of the key residues involved in ACoA recognition, however, there is no interaction involved with the three arginines in the cavity entrance (Arg36, Arg223 and Arg262). Distance to Cys112 is  $9.8 \pm 1.2 \text{ Å}.$ 

Polidocanol (Fig. 15) is a long chain composed of carbon and oxygen atoms, it interacts only through non-polar interactions with the L-shaped portion of the binding pocket. Distance to Cys112 is around  $6.9 \pm 1.4 \text{ Å}$ .

Overall, the results suggest that these molecules could be used to block access to the active site, preventing the interaction with ACoA, and not necessarily by interacting with the catalytic triad as the natural substrate would do. The next part of the study would be to test these compounds

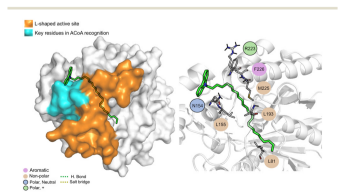


Fig. 13 AB-00063025 (green licorice) bound to PqsD. Interaction map highlighting the most relevant residues (right) involved in ligand stabilization

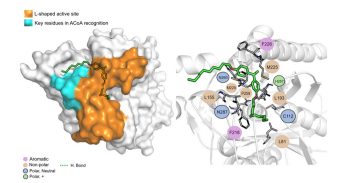


Fig. 14 PasD - AB-00069970 (green licorice) bound to PasD. Interaction map highlighting the most relevant residues (right) involved in ligand stabilization.

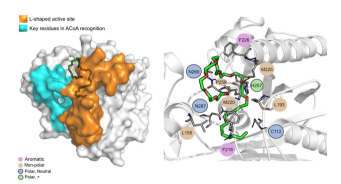


Fig. 15 Polidocanol (green licorice) bound to PgsD. Interaction map highlighting the most relevant residues (right) involved in ligand stabilization.

experimentally and evaluate their ability to suppress the ACoA interaction. PqsD may not seem like an obvious target, but suppressing the formation of the autoinducer molecules could have a huge impact on the quorum sensing system. This work is a good starting point in the elucidation of mechanism of action of this enzyme and the types of interaction that it can perform with molecules other than ACoA. Furthermore, new classes of possible inhibitors have been identified, including isoquinoline and imidazole derivatives.

# Conclusions

An in silico methodology was created and validated to find new molecules capable of inhibiting PqsD, an anthraniloyl-CoA anthraniloyltransferase involved in the synthesis of the secondary metabolites important for the PQS quorum sensing system of Pseudomonas aeruginosa. Protein-ligand docking, structure-based VS and MD simulations were employed to select and validate a list of chemical diverse compounds subsequent experimental evaluations. Five different databases of compounds were screened in a total of 221 146 molecules, and 28 compounds were selected to be further analysed with MD simulations and free energy calculations.

The MD simulations are a very good tool to consolidate the docking prediction as it allows for the search of the most

stable ligand conformation eliminating the bias and improving the free energy predictions. The entropic contribution, in the free energy calculations was overlooked because is computationally time consuming and expensive to calculate. However, this work is the first that describes a complete in silico methodology (from molecular docking to dynamics simulations molecular and free calculations) to find possible inhibitors of PqsD and the main objective was to narrow down the list of compounds, rank them, and look for new drug design scaffold for future experimental testing. The goal was successfully achieved and, naturally, the next stage is the design of an experimental methodology to test the compounds selected and evaluate their specificity toward the pqs system of *P. aeruginosa*.

From the simulations it is hypothesized that these compounds may interact with the upper region of the protein tunnel by blocking the access to some of the key residues involved in ACoA recognition. Only one compound, AB00028362, seems to have the ability to directly interact with the residues in the binding site, as an inhibitor would, mainly through the establishment of a covalent bond with Cys112. Overall, this study suggests two mechanisms of action for inhibiting PqsD activity. One, by competitively binding with the natural autoinducer at the active site; a second mode, by blocking access to the protein tunnel. These modes, and the structural scaffolds identified can be further explored for the identification of new molecules with higher affinity or specificity and for experimental/theoretical optimization.

#### Author contributions

Conceptualization, S. F. S.; methodology, T. F. V.; software, T. F. V.; validation, T. F. V.; formal analysis, T. F. V.; investigation, T. F. V.; data curation, T. F. V.; writing - original draft preparation, T. F. V.; writing - review and editing, S. F. S., M. S.; supervision, S. F. S., M. S.; project administration, S. F. S.; funding acquisition, S. F. S., M. S.

## Conflicts of interest

There are no conflicts to declare.

# Acknowledgements

This work was supported by national funds from Fundação para a Ciência e a Tecnologia (grant numbers: SFRH/BD/ 137844/2018, UIDP/04378/2020, UIDB/04378/2020 the 2020.01423.CEECIND/CP1596/CT0003). Some calculations were produced with the support of the INCD funded by the FCT and FEDER under project 01/SAICT/2016 number 022153, and projects CPCA/A00/7140/2020 and CPCA/ A00/7145/2020. This work was also financially supported by: Base Funding [LA/P/0045/2020 (ALiCE), UIDB/00511/2020 and UIDP/00511/2020 (LEPABE)], funded by national funds through [FCT/MCTES (PIDDAC)].

## Notes and references

- 1 R. F. Langendonk, D. R. Neill and J. L. Fothergill, Front. Cell. Infect. Microbiol., 2021, 11, 665759.
- 2 M. Prothiwa, D. Szamosvári, S. Glasmacher and T. Böttcher, Beilstein J. Org. Chem., 2016, 12, 2784-2792.
- 3 M. P. Storz, C. Brengel, E. Weidel, M. Hoffmann, K. Hollemeyer, A. Steinbach, R. Müller, M. Empting and R. W. Hartmann, ACS Chem. Biol., 2013, 8, 2794-2801.
- 4 Z. Zhou and S. Ma, ChemMedChem, 2017, 12, 420-425.
- 5 Y.-M. Zhang, M. W. Frank, K. Zhu, A. Mayasundari and C. O. Rock, J. Biol. Chem., 2008, 283, 28788-28794.
- 6 A. K. Bera, V. Atanasova, H. Robinson, E. Eisenstein, J. P. Coleman, E. C. Pesci and J. F. Parsons, Biochemistry, 2009, 48, 8644-8655.
- 7 M. P. Storz, C. K. Maurer, C. Zimmer, N. Wagner, C. Brengel, J. C. de Jong, S. Lucas, M. Müsken, S. Häussler, A. Steinbach and R. W. Hartmann, J. Am. Chem. Soc., 2012, 134,
- 8 A. Thomann, C. Brengel, C. Börger, D. Kail, A. Steinbach, M. Empting and R. W. Hartmann, ChemMedChem, 2016, 11, 2522-2533.
- 9 Y. Luo, Y.-S. Yang, J. Fu and H.-L. Zhu, Expert Opin. Ther. Pat., 2012, 22, 1325-1336.
- 10 M. M. Alhamadsheh, F. Musayev, A. A. Komissarov, S. Sachdeva, H. T. Wright, N. Scarsdale, G. Florova and K. A. Reynolds, Chem. Biol., 2007, 14, 513-524.
- 11 D. Pistorius, A. Ullrich, S. Lucas, R. W. Hartmann, U. Kazmaier and R. Müller, ChemBioChem, 2011, 12, 850-853.
- 12 E. Weidel, J. C. de Jong, C. Brengel, M. P. Storz, A. Braunshausen, M. Negri, A. Plaza, A. Steinbach, R. Müller and R. W. Hartmann, J. Med. Chem., 2013, 56, 6146-6155.
- 13 S. Hinsberger, J. C. de Jong, M. Groh, J. Haupenthal and R. W. Hartmann, Eur. J. Med. Chem., 2014, 76, 343-351.
- 14 J. H. Sahner, C. Brengel, M. P. Storz, M. Groh, A. Plaza, R. Müller and R. W. Hartmann, J. Med. Chem., 2013, 56, 8656-8664.
- 15 S. F. Sousa, A. J. M. Ribeiro, J. T. Coimbra, R. P. Neves, S. A. Martins, N. S. H. Moorthy, P. A. Fernandes and M. J. Ramos, Curr. Med. Chem., 2013, 20, 2296-2314.
- 16 S. F. Sousa, P. A. Fernandes and M. J. Ramos, Proteins: Struct., Funct., Genet., 2006, 65, 15-26.
- 17 D. Lapaillerie, C. Charlier, V. Guyonnet-Dupérat, E. Murigneux, H. S. Fernandes, F. G. Martins, R. P. Magalhães, T. F. Vieira, C. Richetta, F. Subra, S. Lebourgeois, C. Charpentier, D. Descamps, B. Visseaux, P. Weigel, A. Favereaux, C. Beauvineau, F. Buron, M.-P. Teulade-Fichou, S. Routier, S. Gallois-Montbrun, L. Meertens, O. Delelis, S. F. Sousa and V. Parissi, Antimicrob. Agents Chemother., 2022, 66(8), e0008322.
- 18 N. D. S. L. De Guzman, J. A. C. Madrono, M. A. Endriga and E. J. L. Frio, in 2021 IEEE Int. Conf. Bioinforma. Biomed., 2021, pp. 376-380.
- 19 K. D. G. Martin, K. G. V. Padilla, I. J. A. Buan, K. Dave Gloria Martin, K. Grace Vergara Padilla and I. Joyce Arenas Buan, Orient. J. Chem., 2021, 37, 143-150.

Paper

20 F. G. Martins, A. Melo and S. F. Sousa, Molecules, 2021, 26, 2600.

- 21 T. F. Vieira, F. G. Martins, J. P. Moreira, T. Barbosa and S. F. Sousa, Molecules, 2021, 26, 6162.
- 22 T. F. Vieira, R. P. Magalhães, M. Simões and S. F. Sousa, Antibiotics, 2022, 11, 185.
- 23 R. P. Magalhães, T. F. Vieira, A. Melo and S. F. Sousa, Mol. Syst. Des. Eng., 2022, 7, 434-446.
- 24 S. Fernandes, A. Borges, I. B. Gomes, S. F. Sousa and M. Simões, Food Res. Int., 2023, 165, 112519.
- 25 H. M. M. Berman, J. Westbrook, Z. Feng, G. Gilliland, T. N. Bhat, H. Weissig, I. N. Shindvalov and P. E. Bourne, Nucleic Acids Res., 2000, 28, 235-242.
- 26 R. P. Magalhães, T. F. Vieira, H. S. Fernandes, A. Melo, M. Simões and S. F. Sousa, Trends Biotechnol., 2020, 38, 937–940.
- 27 J. Jumper, R. Evans, A. Pritzel, T. Green, M. Figurnov, O. Ronneberger, K. Tunyasuvunakool, R. Bates, A. Žídek, A. Potapenko, A. Bridgland, C. Meyer, S. A. A. Kohl, A. J. Ballard, A. Cowie, B. Romera-Paredes, S. Nikolov, R. Jain, J. Adler, T. Back, S. Petersen, D. Reiman, E. Clancy, M. Zielinski, M. Steinegger, M. Pacholska, T. Berghammer, S. Bodenstein, D. Silver, O. Vinyals, A. W. Senior, K. Kavukcuoglu, P. Kohli and D. Hassabis, Nature, 2021, 596, 583-589.
- 28 M. Varadi, S. Anyango, M. Deshpande, S. Nair, C. Natassia, G. Yordanova, D. Yuan, O. Stroe, G. Wood, A. Laydon, A. Žídek, T. Green, K. Tunyasuvunakool, S. Petersen, J. Jumper, E. Clancy, R. Green, A. Vora, M. Lutfi, M. Figurnov, A. Cowie, N. Hobbs, P. Kohli, G. Kleywegt, E. Birney, D. Hassabis and S. Velankar, Nucleic Acids Res., 2022, 50, D439-D444.
- 29 O. Trott and A. J. Olson, J. Comput. Chem., 2009, 31, 455-461.
- 30 G. Jones, P. Willett, R. C. Glen, A. R. Leach and R. Taylor, J. Mol. Biol., 1997, 267, 727-748.
- 31 T. F. Vieira and S. F. Sousa, Appl. Sci., 2019, 9, 4538.
- 32 T. F. Vieira, R. P. Magalhaes and S. F. Sousa, Front. Drug Chem. Clin. Res., 2019, 2, 1-4.
- 33 The PyMOL Molecular Graphics System, Version 1.2r3pre, Schrödinger, LLC.
- 34 N. M. O'Boyle, M. Banck, C. A. James, C. Morley, T. Vandermeersch and G. R. Hutchison, Aust. J. Chem., 2011, 3, 33.
- 35 S. F. Sousa, N. M. F. S. A. Cerqueira, P. A. Fernandes and M. J. Ramos, Comb. Chem. High Throughput Screening, 2010, 13, 442-453.
- 36 N. M. F. S. A. Cerqueira, D. Gesto, E. F. Oliveira, D. Santos-Martins, N. F. Brás, S. F. Sousa, P. A. Fernandes and M. J. Ramos, Arch. Biochem. Biophys., 2015, 582, 56-67.
- 37 A. Gaulton, A. Hersey, M. Nowotka, A. P. Bento, J. Chambers, D. Mendez, P. Mutowo, F. Atkinson, L. J. Bellis, E. Cibrián-Uhalte, M. Davies, N. Dedman, A. Karlsson, M. P. Magariños, J. P. Overington, G. Papadatos, I. Smit and A. R. Leach, Nucleic Acids Res., 2017, 45, D945-D954.
- 38 M. K. Gilson, T. Liu, M. Baitaluk, G. Nicola, L. Hwang and J. Chong, Nucleic Acids Res., 2016, 44, D1045-D1053.
- 39 J. H. Sahner, M. Empting, A. Kamal, E. Weidel, M. Groh, C. Börger and R. W. Hartmann, Eur. J. Med. Chem., 2015, 96, 14-21.

- 40 G. Allegretta, E. Weidel, M. Empting and R. W. Hartmann, Eur. J. Med. Chem., 2015, 90, 351-359.
- 41 M. M. Mysinger, M. Carchia, J. J. Irwin and B. K. Shoichet, J. Med. Chem., 2012, 55, 6582-6594.
- 42 C. Empereur-Mot, J.-F. Zagury and M. Montes, J. Chem. Inf. Model., 2016, 56, 2281-2286.
- 43 T. Sterling and J. J. Irwin, J. Chem. Inf. Model., 2015, 55, 2324-2337.
- 44 M. F. Hibert, Drug Discovery Today, 2009, 14, 723-725.
- 45 F. Ortuso, D. Bagetta, A. Maruca, C. Talarico, M. L. Bolognesi, N. Haider, F. Borges, S. Bryant, T. Langer, H. Senderowitz and S. Alcaro, Front. Chem., 2018, 6, 130.
- 46 IBS The Natural Compound (NC) collection, InterBioScreen ltd., can be found under https://www.ibscreen.com/naturalcompounds, n.d.
- 47 C. Lyu, T. Chen, B. Qiang, N. Liu, H. Wang, L. Zhang and Z. Liu, Nucleic Acids Res., 2021, 49, D509-D515.
- J. A. Maier, C. Martinez, K. Kasavajhala, L. Wickstrom, K. E. Hauser and C. Simmerling, J. Chem. Theory Comput., 2015, 11, 3696-3713.
- 49 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, Gaussian 16, Revision C.01, Gaussian, Inc., Wallingford CT, 2016.
- 50 J. Wang, R. M. Wolf, J. W. Caldwell, P. A. Kollman and D. A. Case, J. Comput. Chem., 2004, 25, 1157-1174.
- 51 D. A. Case, T. E. Cheatham, T. Darden, H. Gohlke, R. Luo, K. M. Merz, A. Onufriev, C. Simmerling, B. Wang and R. J. Woods, J. Comput. Chem., 2005, 26, 1668-1688.
- 52 W. Humphrey, A. Dalke and K. Schulten, J. Mol. Graphics, 1996, 14, 33-38.
- 53 D. R. Roe and T. E. Cheatham, J. Chem. Theory Comput., 2013, 9, 3084-3095.
- 54 E. Wang, H. Sun, J. Wang, Z. Wang, H. Liu, J. Z. H. Zhang and T. Hou, Chem. Rev., 2019, 119, 9478-9508.
- B. R. Miller, T. D. McGee, J. M. Swails, N. Homeyer, H. Gohlke and A. E. Roitberg, J. Chem. Theory Comput., 2012, 8, 3314-3321.
- 56 S. Genheden and U. Ryde, Expert Opin. Drug Discovery, 2015, 10, 449-461.
- 57 M. Doona and D. Walsh, *Palliat. Med.*, 1998, **12**, 55–58.
- 58 V. Iyer and S. S. Poddar, Eur. J. Contracept. Reprod. Health Care, 2008, 13, 339-350.

- 59 N. Li, J. Li, M. Huang and X. Zhang, Medicine, 2021, 100, e24500.
- 60 Deferoxamine, can be found under https://go.drugbank.com/ drugs/DB00746, n.d.
- 61 C. J. Ingold and S. R. Sergent, Phytonadione (Vitamin K1) [Updated 2023 Feb 11], in StatPearls, StatPearls Publishing, Treasure Island (FL), 2023, Available from: https://www.ncbi. nlm.nih.gov/books/NBK557622/.
- 62 D. I. Ball, R. T. Brittain, R. A. Coleman, L. H. Denyer, D. Jack, M. Johnson, L. H. C. Lunts, A. T. Nials, K. E. Sheldrick and I. F. Skidmore, Br. J. Pharmacol., 1991, 104, 665-671.
- 63 R. N. Marcus and J. Mendels, J. Clin. Psychiatry, 1996, 57, 19-23.
- 64 C. Piérard-Franchimont, V. Goffin, J. Decroix and G. E. Piérard, Skin Pharmacol. Physiol., 2002, 15, 434-441.
- 65 A. Galán, L. Moreno, J. Párraga, Á. Serrano, M. J. Sanz, D. Cortes and N. Cabedo, Bioorg. Med. Chem., 2013, 21, 3221-3230.

- 66 L. Opletal, M. Ločárek, A. Fraňková, J. Chlebek, J. Šmíd, A. Hošť álková, M. Šafratová, D. Hulcová, P. Klouček, M. Rozkot and L. Cahlíková, Nat. Prod. Commun., 2014, 9, 1709-1712.
- 67 Y. Liu, L. Ding, J. He, Z. Zhang, Y. Deng, S. He and X. Yan, Fitoterapia, 2021, 154, 105004.
- 68 T. F. Vieira, R. P. Magalhães, N. M. F. S. A. Cerqueira, M. Simões and S. F. Sousa, Mol. Syst. Des. Eng., 2022, 7, 1294-1306.
- 69 M. Starkey, T. Kitao, J. He, A. Tzika, S. Milot, B. Lesic, V. Righi, F. Lepine, A. Bandyopadhaya, L. Rahme, D. Maura, A. Bandyopadhaya, B. Lesic, J. He, T. Kitao, V. Righi, S. Milot, A. Tzika and L. Rahme, PLoS Pathog., 2014, 10, e1004321.
- 70 B. Shaker, S. Ahmad, J. Lee, C. Jung and D. Na, Comput. Biol. Med., 2021, 137, 104851.
- 71 C. Wang, D. Greene, L. Xiao, R. Qi and R. Luo, Front. Mol. Biosci., 2018, 4, 1-18.