Novel phosphatidylinositol 4-kinases III beta (PI4KIIIβ) inhibitors discovered by virtual screening using free energy models

Natalie M. Colodette¹,² · Lucas S. Franco¹,² · Rodolfo C. Maia¹ · Harold H. Fokoué¹ · Carlos Mauricio R. Sant'Anna¹,³ · Eliezer J. Barreiro¹,²,⁴

Received: 19 February 2020 / Accepted: 17 June 2020 / Published online: 30 June 2020 © Springer Nature Switzerland AG 2020

Abstract

Herein, the LASSBio Chemical Library is presented as a valuable source of compounds for screening to identify hits suitable for subsequent hit-to-lead optimization stages. A feature of the LASSBio Chemical Library worth highlighting is the fact that it is a smart library designed by medicinal chemists with pharmacological activity as the main priority. The great majority of the compounds part of this library have shown in vivo activity in animal models, which is an indication that they possess overall favorable bioavailability properties and, hence, adequate pharmacokinetic profiles. This, in turn, is supported by the fact that approximately 85% of the compounds are compliant with Lipinski’s rule of five and ca. 95% are compliant with Veber’s rules, two important guidelines for oral bioavailability. In this work it is presented a virtual screening methodology combining a pharmacophore-based model and an empirical Gibbs free energy-based model for the ligand–protein interaction to explore the LASSBio Chemical Library as a source of new hits for the inhibition of the phosphatidylinositol 4-kinase IIIβ (PI4KIIIβ) enzyme, which is related to the development of viral infections (including enteroviruses, SARS coronavirus, and hepatitis C virus), cancers and neurological diseases. The approach resulted in the identification of two hits, LASSBio-1799 (7) and LASSBio-1814 (10), which inhibited the target enzyme with IC_{50} values of 3.66 μM and IC_{50} and 6.09 μM, respectively. This study also enabled the determination of the structural requirements for interactions with the active site of PI4KIIIβ, demonstrating the importance of both acceptor and donor hydrogen bonding groups for forming interactions with binding site residues Val598 and Lys549.

Keywords
Empirical free energy model · Virtual screening · Lassbio chemical library · PI4KIIIβ ligands · Kinase inhibitors

Introduction

Chemical libraries have been playing an important role in contemporary drug discovery and development. Lead discovery is a critical phase in drug discovery, and lead compounds can be obtained from different sources, such as natural products, endogenous ligands, compounds in...
clinical trials and marketed drugs [1]. Currently, medicinal chemists often use strategies such as virtual screening (VS) or high-throughput screening (HTS) at the beginning of drug discovery campaigns to identify promising chemical structures as promising starting points for further optimization, which have made chemical libraries valuable sources of compounds [2, 3].

Now that Medicinal Chemistry is entering the era of “big data”, chemical libraries have become even more important as tools for exploring the vastness of the chemical space [4]. Pharmaceutical companies’ proprietary compound libraries are frequently used in small-molecule drug discovery research programmes, and strategies for enhancing chemical diversity with the aim of appropriately covering chemical space are essential to the success of a drug discovery campaign [5]. Along with extensive chemical diversity, achieving a successful hit/lead identification rate from a chemical library is also directly related to its components having adequate physicochemical properties [6, 7]. Physicochemical parameters related to oral bioavailability are important indicators of the overall quality of chemical libraries [5]. In this context, rule-based guidelines, such as Lipinski’s Rule of Five (Ro5) [8] and Veber’s rules [9], have emerged to support the interpretation of parameters and to filter and optimize chemical libraries.

Currently, chemical libraries of contrasting sizes are available, ranging from a few hundred to millions of compounds. Virtual chemical libraries such as PubChem [10] and ZINC [11] contain millions of compounds. These vast, non-curated libraries generally resemble catalogues and usually demand the employment of a series of filters to enhance the quality of the structures subject to screening.

On the other side of the spectrum, there are “smart libraries” that have been built upon Medicinal Chemistry concepts and strategies to improve the lead-likeness of the hits, and therefore, increase the success rates of the screening. The Prestwick Chemical Library [12] is probably the flagship smart library. It consists of off-patent selected drugs chosen to increase the probability of identifying high-quality hits by prioritizing high chemical and pharmacological diversity. According to Prestwick’s website, drug discovery campaigns using their library as a screening starting point have resulted in one drug on the market and eleven drug candidates in clinical trials [12].

In this context, the contributions of chemical libraries in the early phases of drug discovery programmes have undoubtedly increased in recent years [5]. The usefulness of such an approach is illustrated by the discovery of the anti-HIV drug maraviroc. The early discovery phase of this clinical agent was based on a HTS of Pfizer’s proprietary chemical library that was conducted to find novel starting points for a low-molecular weight and orally bioavailable CCR5 antagonist as a clinical candidate for the treatment of AIDS [13].

By understanding that performing some sort of screening on chemical libraries as the starting point in drug discovery campaigns is a one-way ticket phenomenon, we have recently started to explore our in-house chemical library, named “LASSBio Chemical Library”, more often in our Medicinal Chemistry research programmes [14]. The LASSBio Chemical Library is a smart library currently containing ca. 2300 compounds; the library content selection has been driven by Medicinal Chemistry concepts, with pharmacological activity as the main priority and with a focus on designing compounds with the most adequate lead-like and/or drug-like properties (Fig. 1). For instance, approximately 85% of these compounds are compliant with Lipinski’s Ro5 [8] and 95% with Veber’s rules [9]. The great majority of compounds in the LASSBio Chemical Library have shown in vivo activities in one or more animal models, after being administrated orally, which is an indication that they possess overall favourable

![Fig. 1](image-url) Drug-likeness and lead-likeness ranges of compounds in the LASSBio Chemical Library considering their molecular weight and cLogP distribution
bioavailability and, hence, adequate pharmacokinetic profiles.

Kinases are validated targets in drug discovery [15], and this work will be focused on a lipid-kinase, PI4KIIIβ, which is related to the development of various diseases such as viral infections (including enteroviruses, SARS coronavirus, and hepatitis C virus), cancers and neurological diseases [16–22]. PI4KIIIβ is required for cellular entry by viruses bearing the severe acute respiratory syndrome-coronavirus (SARS-CoV) spike protein and the cell entry mediated by SARS-CoV spike protein is strongly inhibited by knockdown of PI4KIIIβ [23]. The identification of new PI4K inhibitors is expected to be of therapeutic value and help elucidate the mechanisms of action by which this enzyme works [24].

In this work, a combination of SBDD and LBDD procedures was applied for a virtual screening with the LASSBio Chemical Library to successfully identify new inhibitors with a new molecular pattern for the PI4KIIIβ isoform. The procedure started by selecting candidate inhibitors from the LASSBio Chemical Library by means of a comparison with a proposed pharmacophore map for PI4KIIIβ inhibitors. Geometric criteria can be a fast way to identify candidate enzyme inhibitors, but the screening approach is expected to be made more effective by a combination with some SBDD method to quantify the interaction between the selected candidate molecules and their expected target, since it is expected that a better interaction is related to a better activity. The effectiveness of this second step, therefore, is dependent on the availability of a reliable method to evaluate ligand–protein interactions.

In fact, the activity can be predicted directly by means of some QSAR approach, but this involves the evaluation of a number of ligand-related terms and the use of some statistical method to identify which terms are the most important for the observed activity. With some training, excellent correlations between selected terms and the activity can be obtained, but in many cases the complex nature of these correlations makes difficult the interpretation of the resulting equations, and, consequently, their application.

The ligand–protein interaction is determined by the Gibbs free energy of binding (ΔGbind). Methods such as free energy perturbation (FEP) can be used for evaluating ΔGbind, but its generalized use in virtual screening campaigns is difficulted by the high computational cost of the method. A simpler and faster approach to estimate ΔGbind is the use of a thermodynamic cycle to develop a function calibrated with available experimental data, containing a series of terms that can be calculated separately [25, 26]. Entropic terms calculation is always the most difficult problem to solve in such models, but it can be simplified by using a thermodynamic cycle to obtain relative values, i.e. the model could be used to calculate ΔGbind for a ligand provided that the corresponding value for a reference ligand is known [26].

The term associated with the interactions between the ligand and the protein is a pure enthalpy term that can be evaluated by a number of methods, including semi-empirical molecular orbital models that can produce results with good accuracy at a low computational cost [28–30]. Some examples of the use of enthalpy data calculated by semi-empirical methods for free energy calculations are available in the literature [25, 31–33].

The evaluation of the remaining terms is more laborious, but it was made easier by calibrating the ΔGbind prediction function by means of available experimental data, such as Ki or IC50 values, which reflect the affinity of the compounds for enzymes or receptors, leading to the so-called empirical models [26, 34]. In this paper, using empirical ΔGbind models created to predict the activity of PI4KIIIβ inhibitors based on data available in the literature, we screened the LASSBio Chemical Library for potential PI4KIIIβ inhibitors and then experimentally determined their enzymatic activities to validate our approach.

**Methodology**

The pharmacophore model of the PI4KIIIβ enzyme was created based on a meticulous analysis of the binding modes of compounds selected from articles in which structure–activity relationships were studied [35–40]. In this analysis, it was possible to identify the pharmacophore features that were crucial for molecular recognition by the enzyme, described in Fig. 2. The distances between the pharmacophore features for all compounds used were defined using PyMOL v.1.4 (Schrödinger, LLC)
to determine the ideal distance ranges between the three main points that allowed molecular recognition (Fig. 2).

For construction of the free energy model to evaluate the ligand–protein interaction for the virtual screening, a literature review was initially performed to identify inhibitors of human PI4KIIIβ, which allowed the selection of 33 ligands (Fig. S1 Supporting Information). The protonation states at physiological pH (7.4) were defined by Percepta 2012 Release (ACD/LABS). The 3D structures were optimized using Spartan’16 (Wavefunction, Inc.) in two steps: a Monte Carlo conformational analysis with the MMFF molecular mechanics method [41], followed by a structural re-optimization of the lowest-energy conformer with the PM6 semi-empirical method [42].

Of the 11 Homo sapiens PI4KIIIβ structures available in the Protein Data Bank (PDB) [43], the structure of 4D0L [44] has the best resolution (2.94 Å). This structure was chosen for molecular docking studies of the selected inhibitors (Fig. S1) with GOLD 5.4.0 (CCDC); the ChemPLP [45] scoring function was employed because it presented the better performance in redocking studies (an average RMSD of 0.82 Å). Hydrogen atoms were used to complete the valence of the atoms where the bonds were truncated. In order to avoid large structural changes in the truncated protein models that could occur during the energy minimization, the coordinates of the atoms from the peptide bonds were frozen. In this way, the general arrangement of the binding site was conserved, while the ligand and the side chains conformations were allowed to adopt better conformations to improve their interactions according to the quantum mechanical model. The total charges were calculated considering lysine and arginine residues as protonated (charge equal to +1) and the aspartic acid and glutamic acid residues in the deprotonated form (charge equal to −1); histidine residues were considered as neutral. In order to include the effect of the medium around the selected residues in the quantum calculations, the remaining protein was replaced by a continuum with a suitable dielectric constant.

The resulting systems were then subjected to geometry optimization using the PM7 semi-empirical molecular orbital method [46], available in MOPAC2016 (Stewart Computational Chemistry). PM7 was chosen because it is better than previous Hamiltonians for describing noncovalent interactions, an essential characteristic for the present study. Hydrogen atoms were used to complete the valence of the atoms of the truncated bonds. The ligand–protein interaction enthalpy was determined by Eq. 1:

\[ \Delta H_{int} = \Delta H_{f}^{\text{complex}} - (\Delta H_{f}^{\text{protein}} + \Delta H_{f}^{\text{ligand}}) \] (1)

where \( \Delta H_{int} \) is the interaction enthalpy and \( \Delta H_{f}^{\text{complex}} \), \( \Delta H_{f}^{\text{protein}} \) and \( \Delta H_{f}^{\text{ligand}} \) are the enthalpies of formation of the complex, the empty binding site and the ligand, respectively. In each case, the enthalpy of formation was obtained after geometry optimizations to stationary points of the potential energy surface, so the conformation of the ligand is not the

Ten solutions were generated in each docking run, and the process was repeated three times, generating a total of thirty solutions for each ligand. There was a great structural variability in the generated poses, so the results were analysed according to two criteria: first, only the solutions with a binding mode that could match the PI4KIIIβ pharmacophore model (Fig. 2) were selected, considering that the lack of any of the interactions could lead to the inactivity of a candidate inhibitor on the studied enzyme; then, the pose with the highest score was chosen among the poses matching the pharmacophoric criteria.

To reduce the computational time for quantum mechanical calculations necessary for obtaining the interaction enthalpy for the empirical \( \Delta G_{bind} \) prediction models, only amino acid residues that were part of the enzyme’s active site in the chosen docking poses were considered. The binding site region was composed of all amino acid residues with at least one atom within a 6 Å radius from the ligand. H atoms were used to complete the valence of the atoms where the bonds were truncated. In order to avoid large structural changes in the truncated protein models that could occur during the energy minimization, the coordinates of the atoms from the peptide bonds were frozen. In this way, the general arrangement of the binding site was conserved, while the ligand and the side chains conformations were allowed to adopt better conformations to improve their interactions according to the quantum mechanical model. The total charges were calculated considering lysine and arginine residues as protonated (charge equal to +1) and the aspartic acid and glutamic acid residues in the deprotonated form (charge equal to −1); histidine residues were considered as neutral. In order to include the effect of the medium around the selected residues in the quantum calculations, the remaining protein was replaced by a continuum with a suitable dielectric constant.

The resulting systems were then subjected to geometry optimization using the PM7 semi-empirical molecular orbital method [46], available in MOPAC2016 (Stewart Computational Chemistry). PM7 was chosen because it is better than previous Hamiltonians for describing noncovalent interactions, an essential characteristic for the present study. Hydrogen atoms were used to complete the valence of the atoms of the truncated bonds. The ligand–protein interaction enthalpy was determined by Eq. 1:

\[ \Delta H_{int} = \Delta H_{f}^{\text{complex}} - (\Delta H_{f}^{\text{protein}} + \Delta H_{f}^{\text{ligand}}) \] (1)

where \( \Delta H_{int} \) is the interaction enthalpy and \( \Delta H_{f}^{\text{complex}} \), \( \Delta H_{f}^{\text{protein}} \) and \( \Delta H_{f}^{\text{ligand}} \) are the enthalpies of formation of the complex, the empty binding site and the ligand, respectively. In each case, the enthalpy of formation was obtained after geometry optimizations to stationary points of the potential energy surface, so the conformation of the ligand is not the
same inside and outside the binding site, as expected. The
same apply to the side chains of the amino acid residues,
i.e. their conformations in the optimized complex and in the
empty binding site are not the same.

Following the original proposal based on a thermody-
namic cycle for the construction of free energy prediction
models [26], it was necessary to include two additional terms
in that model: a term associated with the conformational
entropic losses that occur when acyclic bonds in the ligand
become non-rotatable upon binding was obtained from the
GOLD 5.4.0 (CCDC) ChemPLP scoring function results
of the molecular docking solutions (torsional energy: Etor)
[45]; and the energy term for the ligands’ solvation (E_solv),
which was calculated by the SM5.4 model [47] available in
Spartan’16 (Wavefunction, Inc.).

For the calibration of the final equation for ΔG_bind cal-
culation, experimental ΔG_bind data are necessary and they
could be obtained from Ki data (assuming ΔG_bind = RT ln
Ki). Unfortunately, it was not possible apply the Cheng–Prue-
ssoff equation to directly convert the available IC50 into Ki,
since some necessary quantities were not available in the
papers from which the IC50 data were collected: the fixed
substrate concentration and the concentration of substrate
at which the enzyme activity is at half maximal. So, in the
absence of these data, we assumed that the IC50 data, as a
first approximation, would be linearly related to Ki. In this
case, RT ln Ki could be replaced by RT ln (X. IC50) = RT
ln IC50 + RT ln X, where X is the proportionality constant
between Ki and IC50. Although X is unknown, RT ln X
would be incorporated in the coefficient a3 from Eq. 2, which
would be obtained with the remaining coefficients after cali-
bration of the final equation by multiple regression with the
experimental data.

Naturally, the same reasoning holds for logarithms to
base 10, so, after replacing ΔG_bind with pIC50 (− log IC50),
the calculated energy terms were combined with the pIC50
data from known inhibitors with a multiple linear regression
analysis to calibrate the model. This assumption, however,
can present some limitations because, unlike Ki values, IC50
data can be influenced by the experimental method used
in their determination [48]. Thus, the influence of the IC50
determination method was evaluated by comparing the
results obtained with the data of compounds from the same
reference or obtained by the same methodology.

The final correlations generated by linear regression fol-
lowed the model described by Eq. 2:

\[
pIC50 = a1 \Delta H_{\text{int}} + a2 \text{Etor} - a3 \text{E_{solv}} - a4 \text{E_{solv2}} + a5
\]

(2)

where \(a1, a2, \ldots a5\) are the linear regression coefficients. All sta-
tistical analysis was obtained with OriginPro. In Eq. 2, pIC50
is proposed to have a quadratic dependence with E_solv as
suggested by Wang et al. [26], because it is common that
compounds which are either too hydrophobic or too hydro-
philic would not be able to achieve a high binding affinity.
A better pIC50 would be obtained for compounds with inter-
mediate solubilities, so the dependence between pIC50 and
solubility would be better described by a parabolic function.
For the reader interested in more details about the deriva-
tion of Eq. 2, a discussion is presented in the Supporting
Information.

To evaluate E_solv, we tried different methods, but the
free energy of solvation calculated with the SM5.4 model
[47] produced the best results. In the thermodynamic
cycle, this term represents the free energy cost to desolvate
the ligand molecule prior to its entry into the enzyme,
where the interaction with the binding site will occur.

After obtaining adequate equations, it was the moment
to search for candidate PI4KIIIβ inhibitors among the two
thousand molecules from the LASSBio Chemical Library.
As a first step, they were structurally analysed to verify
the presence of functional groups in positions suitable
for interacting with the PI4KIIIβ active site based on our
pharmacophore model (Fig. 2). This step reduced the num-
ber of compounds to evaluate with the SBDD approach,
since only those that had appropriate distances to match
the pharmacophore model were docked into the active site
of PI4KIIIβ (PDB: 4D0L) using GOLD 5.4.0 (CCDC), as
previously described. Here we kept the same criteria we
used for the literature compounds. Initially, we analyzed
every docking pose and selected only those that presented
the three essential interactions for the molecular recogni-
tion and, after that, we selected the pose with the highest
score among those that performed all three interactions.

To choose compounds for experimental inhibitory
activity determination, their \(\Delta H_{\text{int}}, \text{E}_{\text{solv}}\) and \text{E}_{\text{tor}}\ terms
were calculated and applied to the best activity prediction
models, according to the correlation coefficient values and
structural coverage criteria. Compounds that had calcu-
lated pIC50 values of at least 7.0 by all the chosen models
were selected for inhibitory activity evaluation.

The water solubility of each of the selected compounds
(1, 4, 7, 10, 11 and PIK93) was experimentally determined
to ensure that the tests were carried out within a range of
concentrations that ensures that the compounds are fully
soluble, avoiding false results. The experiments were per-
formed following the protocol described by Nunes, where
the aqueous concentration was correlated with ultraviolet
absorbance [49]. First, the wavelength at which the com-
pounds had the highest absorption was determined, and
then serial dilutions were prepared to obtain a calibration
curve. The compounds were then dissolved in a phosphate
buffer solution to obtain a supersaturated solution, which
was stirred at 37 °C and filtered prior to spectrophoto-
metric analysis. The solubility of each compound was
then determined by the equation obtained from the linear regression of the calibration curve.

The experimental inhibitory activity evaluation was performed by the Reaction Biology Corporation (RBC, USA). The company uses the ADP-Glo™ assay to determine the inhibitory effect of compounds against the PI4KIIIβ enzyme (PROMEGA). This assay can be used to monitor the activity of any enzyme that generates ADP as the product of its reaction. It is performed on a multi-well plate and can detect kinase activity at very low reaction volumes (up to 5 μL).

Results and discussion

Selection of PI4KIIIβ known ligands

Thirty-three PI4KIIIβ-selective inhibitors were selected from the literature (Fig. S1). To ensure that the created models were as general as possible, the compounds’ selection was performed considering an adequate IC_{50} variation. The selected compounds had IC_{50} values between 0.98 nM and 9727 nM (Table S1). They are mainly imidazo-pyridazine or oxazole derivatives (Fig. S1). Some of the molecules show ring bioisosterism [50, 51] relative to these two major classes; the compounds have purines instead of imidazo-pyridazines or they have imidazoles, pyrroles or thiazoles instead of oxazoles. Most of the selected inhibitors have amide or sulfonamide groups in their structures, which are important for the molecular recognition process.

Empirical activity prediction models

The creation of the pharmacophore map showed that there are three residues in PI4KIIIβ that are mainly responsible for molecular recognition: Val598 (hinge), Lys549 and Tyr583. All the selected inhibitors have functional groups with adequate distances to form hydrogen bonds with Val598 and Lys549 and form aromatic ring interactions with Tyr583 (Fig. 2). This map was used as a first criterion to select the docking poses to be used in further calculations to obtain the necessary data for construction of the activity prediction models.

The correlation between the docking scores and the IC_{50} data was very low, R^2 = 0.16. As this may be a result of limitations in the docking scoring functions [52], other methods should be investigated to better quantify the binding modes, a necessary step to get an appropriate correlation with the affinity of the compounds for the enzyme, which was done in this work through the use of empirical models to determine ΔG_{bind}.

To reduce the computational time for these calculations for the construction of the empirical ΔG_{bind} prediction models, only the amino acid residues that were part of the enzyme’s active site in the selected docking poses were considered, and these residues always included Lys549, Tyr583 and Val598, which are the most important amino acids in the molecular recognition process. The medium around the selected residues was represented as a continuum by choosing a suitable dielectric constant. Because interactions occur at sites on the PI4KIIIβ enzyme that are not exposed to solvent, a dielectric constant of 6.5 was chosen for the bulk protein [53].

The calculated data for all the terms necessary for the construction of the prediction model described in the methodology are presented in Table S2 (Supporting Information). The E_{tor} value of each ligand was included to describe the loss of conformational entropy associated with the interaction, corresponding to the energetic effects that oppose the interaction. E_{solv} is related to the ligand interactions with the aqueous medium, which plays an important role in the determination of ΔG_{bind} and, consequently, the pIC_{50} values.

Correlations were obtained by multiple linear regression analyses, considering pIC_{50} as the dependent variable and ΔH_{int}, E_{tor} and E_{solv} as the independent variables, with the potential inclusion of a quadratic term for E_{solv} (E_{solv}^2). In previous works [24, 34], the inclusion of an E_{solv}^2 term was necessary for correctly predicting the free energy changes related to the interaction between the ligands and proteins. The quadratic dependence of pIC_{50} with the solvation energy indicates that intermediate values of solubility are those that generate better pIC_{50} values, as discussed earlier.

Correlations considering the complete set of compounds and also only compounds obtained from the same reference or for which the IC_{50} values were determined by the same methodology were evaluated to verify how differences in the methods used to obtain the experimental data could influence the quality of the models. The analysis of each bibliographic reference allowed the identification of four different methods for IC_{50} determination (References in Table S1):

- ADP-Glo™ assay for kinases (PROMEGA). This method was used to determine the IC_{50} values of compounds S11, S8 to S13 and S15 to S22.
- Coupling of pyruvate kinase and lactate dehydrogenase enzymes assay [16]. This method was used for compounds S2 to S5 and S23 to S33.
- Membrane capture assay [54] was used for compounds S6, S7 and S12.
- Transcreener® assay for fluorescence intensity ADP (BELLBROOK LABS) was used for compound S14.

Among the several correlations evaluated, three were chosen for subsequent studies, as they provided good structural variability in the data set and/or acceptable correlation coefficients, which should be R^2 ≥ 0.7. The equations are shown in Table 1, and the number of compounds used...
and the correlation coefficients are also presented. It can be observed that, in fact, the use of a mixture of compounds obtained from different references negatively influenced the correlation, which could be a consequence of differences in the methods used in each study to determine the IC₅₀ values. To check out for overfitting, adjusted R² was also calculated for each equation.

Although the correlation coefficient obtained from Eq. 3 was below the adequate value (R² ≥ 0.7), this equation was selected because it contains variables referring to all compounds, and therefore, it is the correlation that is the most general.

It was observed that Eq. 4, which has low structural coverage but contained only compounds selected from the same reference [55], presented the highest coefficient among all correlations evaluated and therefore was also selected. Finally, Eq. 5 presented a middle ground between structural coverage and correlation coefficient, and compounds for which the IC₅₀ values were determined by the same methodology [16] were incorporated by adding variables of compounds S6, S7, S12 and S14, which increased the number of compounds, keeping R² within the ideal range (≥ 0.7) and Adj-R² greater than 0.6.

With the exception of model 1 (Eq. 3), which includes all ligands, the remaining models were based on a set of somewhat different structures and also different methods of IC₅₀ determination. In this sense, we consider that our models include different levels of structural coverage and accuracy in predicting biological activity. The most accurate models are also the ones with less structural coverage capability, and vice-versa. As we are interested in both characteristics, we choose the strategy of using consensus results in order to improve the chances of finding new structures with good activity data in the LASSBio chemical library.

**LASSBio Chemical Library virtual screening**

The selection of compounds was made with the 2055 molecules from the LASSBio Chemical Library [14] in two stages. In the first stage, a visual inspection was employed to search for compounds with promising interaction profiles based on the presence of functional groups at suitable positions to interact through hydrophobic interactions with Tyr583, act as H-bond acceptors with Lys549 and H-bond donors with Val598, the three sites important for molecular recognition. This task could be automatized for larger databases, but with a small database, it could be done with a relatively low effort. This first analysis resulted in the selection of 124 candidate PI4KIIIβ ligands. Next, the optimized structures of the selected 124 compounds had the distances between their putative pharmacophoric features measured using PyMOL v. 1.4 (Schrödinger, LLC), for comparison with the pharmacophore model. After this second selection stage, we finally arrived to 70 compounds with adequate distances (see Fig. 2), which were then evaluated in the subsequent SBDD molecular docking study in the PI4KIIIβ active site, respecting their ionization states at physiological pH (7.4).

All the solutions from the molecular docking studies of the 70 compounds were subsequently analysed, leading to the selection of 15 compounds which, when interacting at the PI4KIIIβ site, presented adequate poses that allowed all three interactions necessary for molecular recognition. The structures of the 15 compounds selected in this step are shown in Fig. 3.

The majority of the compounds selected from the LASSBio Chemical Library as PI4KIIIβ ligands are dimethoxy-substituted 2-chloroquinazolines, whereas five (33%) of the selected compounds are N-acylhydrazones (Fig. 3) [59]. The binding modes obtained by molecular docking at the active site of PI4KIIIβ demonstrated that the N-acylhydrazone subunit of these compounds is important for molecular recognition as it interacts with the enzyme’s hinge (Val598). The methoxy groups attached to the 2-chloroquinazoline rings of LASSBio-1799 (7) to LASSBio-1819 (14) interact with the Lys549 residue. All the selected compounds have aromatic rings in their structures, which can form important interactions with Tyr583, which is also involved in pharmacophore recognition.

Table 2 presents the values of the ΔH int, E_tor, and E_solv variables calculated from the best docking solutions of these compounds.

The next step was to use the empirical models created with the data obtained from the literature to predict the pIC₅₀ values of the 15 selected compounds. Since more than one adequate correlation was obtained (Table 1), as

| Equations | Compounds | R² | Adj-R² |
|-----------|-----------|----|--------|
| (Eq. 3)pIC₅₀ = −0.04248ΔHint + 0.25705E_tor − 0.14951E_solv − 0.00287E_solv² + 1.9297S1–S33 | 0.58 | 0.52 |
| (Eq. 4)pIC₅₀ = −0.05068ΔHint + 0.36485E_tor − 2.60214E_solv − 0.04654E_solv² + 32.84142S5; S24–S33 | 0.94 | 0.88 |
| (Eq. 5)pIC₅₀ = −0.06206ΔHint + 0.26733E_tor − 0.77746E_solv − 0.0133E_solv² − 8.67274S2–S7; S12; S14; S23–S33 | 0.78 | 0.72 |

Also shown are the number of compounds used and their correlation coefficients (R²) and adjusted correlation (Adj-R²).
Fig. 3 Structure of the 15 compounds selected from the LASSBio Chemical Library after the molecular docking studies with PI4KIIIβ. *Inter-alia*: LASSBio-693 to LASSBio-774 [56], LASSBio-1059 (Unpublished data), LASSBio-1474 (Unpublished data), LASSBio-1516 [57], LASSBio-1799 to LASSBio-1819 [58], LASSBio-1845 [59].
was previously mentioned, a consensus pIC₅₀ value was calculated by combining the results from the three chosen equations (Table 3).

From the previous discussion, it is clear that because of the limitation of the available experimental data, due either to differences in the methodologies of activity determination or to limited structural variability, the results obtained by the application of our models should be considered only as an indication of the activity profile of new, structurally unrelated compounds. Bearing this in mind and considering an IC₅₀ of 10 μM as an adequate upper limit for the identification of hit compounds, we decided that for a compound to be designated for experimental inhibitory activity evaluation with a reasonable safety margin, its predicted IC₅₀ should be at most 0.1 μM.

Therefore, all the substances that had a pIC₅₀ greater than 7.0, as calculated by all three selected equations (consensus pIC₅₀), were selected for inhibitory activity evaluation (Table 3). The compounds selected based on this criterion were LASSBio-693 (1), LASSBio-1059 (4), LASSBio-1799 (7), LASSBio-1814 (10) and LASSBio-1816 (11) (Fig. 3).

**Inhibitory activity evaluation of the selected compounds**

Starting from an initial value and making serial twofold dilutions, a dose–effect curve based on ten concentrations was prepared. For each of the 5 compounds selected from the LASSBio Chemical Library, the initial concentration value was chosen according to the experimental solubility, determined using the Nunes method described in the methodology section [49]. The experimental solubility, IC₅₀, and experimental pIC₅₀ are shown in Table 4. The standard used was compound PIK93, a potent inhibitor of the enzyme PI4KIIIβ [44].

Unfortunately, it was not possible to evaluate the activity of compound LASSBio-1816 (11) because of its very low solubility (Table 4). Of the four compounds with acceptable solubilities, two showed good activities with IC₅₀ values lower than 10 μM (Table 4). Although the empirical prediction models projected pIC₅₀ values higher than the experimentally observed values, they were able to select truly active compounds with a hit rate of 50% when considering the cherry-picked chemical library compounds for enzymatic inhibition. The difference between predicted and observed pIC₅₀ values could be a result of differences between the methods used to determine the experimental activity for the data used for model calibration and that employed in this work. It is interesting to observe that the most potent PI4KIIIβ inhibitor identified, LASSBio-1799 (7) (IC₅₀ = 3.66 μM), was predicted to be the most active compound of the LASSBio series by two of the three equations used to calculate the consensus pIC₅₀ value (Eqs. 3 and 5).
The interaction modes of the two new most potent inhibitors, LASSBio-1799 (7) and LASSBio-1814 (10), with the PI4KIIIβ binding site are characterized by a hydrogen bond between the N3 atom of the quinazoline ring and Val598 in the hinge, a second hydrogen bond between the methoxy group at the 6-position of the quinazoline ring and the Lys549 residue, as well as aromatic ring interactions between the quinazoline ring and hydrophobic residues in the binding site, exemplified by Tyr583, and hydrogen bonds and additional interactions occurring between the substituents of the sulfonamide group and auxophoric regions in the enzyme’s molecular recognition site (Fig. 4a).

The interaction modes of the two new most potent inhibitors, LASSBio-1799 (7) and LASSBio-1814 (10), with the PI4KIIIβ binding site are characterized by a hydrogen bond between the N3 atom of the quinazoline ring and Val598 in the hinge, a second hydrogen bond between the methoxy group at the 6-position of the quinazoline ring and the Lys549 residue, as well as aromatic ring interactions between the quinazoline ring and hydrophobic residues in the binding site, exemplified by Tyr583, and hydrogen bonds and additional interactions occurring between the substituents of the sulfonamide group and auxophoric regions in the enzyme’s molecular recognition site (Fig. 4a).

The results of the experimental evaluation show that compound LASSBio-1799 (7) is more potent than compound LASSBio-1814 (10) as an inhibitor of enzyme PI4KIIIβ (Fig. 4a). By investigating the binding mode of LASSBio-1799 (7), it could be observed that the thiazole ring attached to the sulfonamide group allows the formation of three additional interactions in the enzyme binding site, a hydrogen bond between the nitrogen of the thiazole ring and Asp600, another hydrogen bond between the oxygen of the sulfonamide and Tyr488 and a possible T stacking interaction with Trp522. Analysis of the binding mode of LASSBio-1814 (10) showed that the unsubstituted sulfonamide generates only one additional hydrogen bond with Gln606, although an additional interaction with the peptidic oxygen of the valine (Val598) in the hinge was present. Nevertheless, we proposed that the three additional interactions observed for compound LASSBio-1799 (7) are the cause of its greater potency against PI4KIIIβ (Fig. 4). These results demonstrate that the 2-chloro-4-aminoquinazolinic structural pattern is privileged in the design of PI4KIIIβ inhibitors.

### Table 4

| Compound          | Experimental solubility (µM) | Initial concentration (µM) | IC₅₀ (µM) | pIC₅₀ | Experimental pIC₅₀ |
|-------------------|-----------------------------|-----------------------------|----------|-------|-------------------|
| LASSBio-693       | 7.82                        | 20                          | >20      | >4.70 |                   |
| LASSBio-1059      | 54.70                       | 100                         | 99.8     | 4.00  |                   |
| LASSBio-1799      | 18.64                       | 50                          | 3.66     | 5.44  |                   |
| LASSBio-1814      | 7.31                        | 20                          | 6.09     | 5.21  |                   |
| LASSBio-1816      | 0.51                        | 1.5                         | –        | –     |                   |
| PIK93             | –                           | 1                           | 0.00578  | 8.24  |                   |

The initial concentrations chosen for each compound that were used to prepare the ten-point serial twofold dilution dose–effect curve are also shown.
Conclusions

In this study, we presented for the first time the LASSBio Chemical Library, which was successfully explored by a virtual screening procedure to identify hit compounds suitable for further hit-to-lead optimization steps in the context of developing new PI4KIIIβ inhibitors. The potential applications of these inhibitors in Medicinal Chemistry are promising, specially at the present Covid-19 pandemics, since RNA viruses hijack the enzyme in order to modify the structure of intracellular membranes and use them for the construction of functional replication machinery; a study of PI4KIIIβ inhibitors showed that they exerted antiviral activity against a panel of single-stranded positive-sense RNA viruses [37].

This virtual screening consisted in the combination of two methods, i.e. a receptor-based pharmacophore model and an empirical free energy prediction model. This combined strategy enabled the identification of two new inhibitors for the target enzyme, LASSBio-1799 (7) and LASSBio-1814 (10), which presented IC_{50} = 3.66 μM and IC_{50} = 6.09 μM, respectively. In this context, 2-chloro-4-aminoquinazolinic derivatives can be considered a promising starting point for the identification of PI4KIIIβ inhibitors.

Additionally, it was possible to establish the structural requirements for interactions with the active site of PI4KIIIβ, demonstrating the importance of the presence of hydrogen bond acceptor and donor groups for forming interactions with binding site residues Val598 and Lys549, as well as the presence of hydrophobic groups, which are also important for molecular recognition.

Our proposal was to develop a free energy-based model to predict the activity of PI4KIIIβ inhibitors and apply it, after a pre-selection with a pharmacophore-based model, to find out candidate new PI4KIIIβ inhibitors in our in-house LASSBio Chemical Library. It must be stressed that no compound present at the LASSBio Chemical Library was originally designed to inhibit PI4KIIIβ.

The present screening methodology, besides being fast and low-cost, was effective, since two of the four selected compounds that had adequate solubility to be evaluated against PI4KIIIβ presented IC_{50} values below 10 μM, a hit rate of 50%, considering only the assayed compounds. The complete search procedure of potential PI4KIIIβ inhibitors presented by us is, sequentially, ligand-based (by comparison with the pharmacophore constructed from known inhibitors), structure-based (by molecular docking in the binding pocket of the enzyme), and property-based (by calculation of the binding free energy composed of rationally selected terms from the thermodynamic cycle originally proposed by Wang et al. [26]), which we think improves the chances of finding real active compounds in the virtual screening approach.

The observed hit rate is strongly suggestive of the efficiency of the procedure, since the chances of choosing a compound at random in a chemical library and that compound being able to inhibit a specific enzyme should be quite small. For example, the experimental tests for finding hits by HTS have a hit rate between 0.01 and 0.1% [60]. This low performance is in part a consequence of the presence of compounds that interfere with elements of the assay format or technique, but they are indicative that these chances should be quite small. Therefore, the adequate use of in silico methodologies is one valid alternative to enhance the chances of finding hits for a given target through chemical library screening.

Supporting information

- References, literature data and chemical structures of the compounds used to build the models;
- Further information about the data generated by the models;
- Chemical structure and SMILES code of the compounds used to build the models;
- IC_{50} curves of all tested compounds.

Acknowledgements The authors would like to thank the Brazilian funding agencies for the financial support involved in this work: Conselho Nacional de Desenvolvimento Científico e Tecnológico—CNPq (INCT-INOFAR grant #465.249/2014-0; grant #309229/2018-9); Coordenação de Aperfeiçoamento de Pessoal de Nível Superior—Brasil (CAPES)—Finance Code 001; Fundação Carlos Chagas Filho de Amparo à Pesquisa do Estado do Rio de Janeiro—FAPERJ (Grant E-26/202.146/2015); Additionally, we would like to thank the Biomedical Sciences Institute of the Rio de Janeiro Federal University (ICB-UFRJ) for the infrastructure provided to develop this work.

Author contributions All authors contributed to the study conception and design. Material preparation, data collection and analysis were performed by NMC. The first draft of the manuscript was written by NMC and all authors revised and prepared following versions of the manuscript. All authors read and approved the final version of the manuscript.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

References

1. Holenz J, Stoy P (2019) Advances in lead generation. Bioorg Med Chem Lett 29:517–524. https://doi.org/10.1016/j.bmcl.2018.12.001
2. Hersey A, Chambers J, Bellis L et al (2015) Chemical databases: curation or integration by user-defined equivalence? Drug Discov Today Technol 14:17–24. https://doi.org/10.1016/j.ddtec.2015.01.005

3. Walters WP (2019) Virtual chemical libraries. J Med Chem 62:1116–1124. https://doi.org/10.1021/acs.jmedchem.8b01048

4. Follmann M, Briem H, Steinmeyer A et al (2019) An approach towards enhancement of a screening library: the next Generation Library Initiative (NGLI) at Bayer: against all odds? Drug Discov Today 24:668–672. https://doi.org/10.1016/j.drudis.2018.12.003

5. Vieth M, Siegel MG, Higgs RE et al (2004) Characteristic physical properties and structural fragments of marketed oral drugs. J Med Chem 47:224–232. https://doi.org/10.1021/jm030267j

6. Ohno K, Nagahara Y, Tsuoyama K, Orita M (2010) Are there differences between launched drugs, clinical candidates, and commercially available compounds? J Chem Inf Model 50:815–821. https://doi.org/10.1021/ci100023s

7. Lipinski CA, Lombardo F, Dominy BW, Feeney PJ (1997) Experimental and computational approaches to estimate solubility and permeability in drug discovery and development settings 1PII of original article: S0169–409X(96), 00423–1. The article was originally published in Advanced Drug Delivery Reviews 23 (1997). Adv Drug Deliv Rev 23:23–26. https://doi.org/10.1016/S0169-409X(00)00129-0

8. Veber DF, Johnson SR, Cheng H-Y et al (2002) Molecular properties that influence the oral bioavailability of drug candidates. J Med Chem 45:2615–2623. https://doi.org/10.1021/jm020017n

9. Wang Y, Xiao J, Suzek TO et al (2009) PubChem: a public information system for analyzing bioactivities of small molecules. Nucleic Acids Res 37:W623–W633. https://doi.org/10.1093/nar/gkp456

10. Irwin JJ, Shoichet BK (2005) ZINC: a free database of commercially available compounds for virtual screening. J Chem Inf Model 45:177–182. https://doi.org/10.1021/ci049714u

11. Prestwick Chemical. https://www.prestwickchemical.com/libraries-screening-lib-pcl.html. Accessed 2 Jul 2019

12. Wood A, Armour D (2005) The discovery of the CCR5 receptor antagonist, UK–427,857, a new agent for the treatment of HIV infection and AIDS. In: King FD (ed) Progress in medicinal chemistry. Elsevier, London, pp 239–271

13. LASSBio Chemical Library. https://www.lassbiochemicallibrary.com. Accessed 2 Jul 2019

14. Richard AW, Frederick G (2011) Kinase drug discovery, 1st edn. Royal Society of Chemistry, Cambridge

15. Borawski J, Troke P, Puyang X et al (2009) Class III phosphatidylinositol 4-kinase alpha and beta are novel host factor regulators of hepatitis C virus replication. J Virol 83:10058–10074. https://doi.org/10.1128/JVI.02418-08

16. Li J, Lu Y, Zhang J et al (2010) PI4KIIα is a novel regulator of tumor growth by its action on angiogenesis and HIF–Iα regulation. Oncogene 29:2550–2559. https://doi.org/10.1038/onc.2010.14

17. Moorhead AM, Jung YJ, Smirnov A et al (2010) Multiple host proteins that function in phosphatidylinositol-4-phosphate metabolism are recruited to the chlamydial inclusion. Infect Immun 78:1990–2007. https://doi.org/10.1128/IAI.01340-09

18. Delang L, Paeshuyse J, Neets J (2012) The role of phosphatidylinositol 4-kinases and phosphatidylinositol 4-phosphate during viral replication. Biochem Pharmacol 84:1400–1408. https://doi.org/10.1016/j.bcp.2012.07.034

19. Clayton EL, Minogue S, Waugh MG (2013) Phosphatidylinositol 4-kinases and PI4P metabolism in the nervous system: roles in psychiatric and neurological diseases. Mol Neurobiol 47:361–372. https://doi.org/10.1007/s12035-012-8358-6

20. Waugh MG (2012) Phosphatidylinositol 4-kinases, phosphatidylinositol 4-phosphate and cancer. Cancer Lett 325:125–131. https://doi.org/10.1016/j.canlet.2012.06.009

21. Altan-Bonnet N, Balla T (2012) Phosphatidylinositol 4-kinase IIIβ is required for severe acute respiratory syndrome coronavirus spike-mediated cell entry. J Biol Chem 287:8457–8467. https://doi.org/10.1074/jbc.M111.312561

22. Boura E, Nencka R (2012) Phosphatidylinositol 4-kinases: function, structure, and inhibition. Exp Cell Res 373:136–145. https://doi.org/10.1016/j.yexcr.2015.03.028

23. Yang N, Ma P, Lang J et al (2012) Phosphatidylinositol 4-kinase IIIβ is required for severe acute respiratory syndrome coronavirus spike-mediated cell entry. J Biol Chem 287:8457–8467. https://doi.org/10.1074/jbc.M111.312561

24. Boura E, Nencka R (2012) Phosphatidylinositol 4-kinases: function, structure, and inhibition. Exp Cell Res 373:136–145. https://doi.org/10.1016/j.yexcr.2015.03.028

25. Murray CW, Verdonk ML (2002) The consequences of translational and rotational entropy lost by small molecules on binding to proteins. J Comput Aided Mol Des 16:741–753. https://doi.org/10.1023/A:102246720849

26. Stewart JJP (1990) Reviews in computational chemistry. Wiley, Hoboken

27. SantAnna CMR (2009) Molecular modeling methods in the study and design of bioactive compounds: an introduction. Rev Virtual Química. https://doi.org/10.5935/1984-6835.20090007

28. Menikarachchi L, Gascon J (2010) QM/MM approaches in medicinal chemistry research. Curr Top Med Chem 10:46–54. https://doi.org/10.2174/156802610790232297

29. WollackOM, Merz KM (2007) Assessment of semiempirical quantum mechanical methods for the evaluation of protein structures. J Chem Theory Comput 3:1609–1619. https://doi.org/10.1021/ct060325q

30. González R, Suárez CF, Bohórquez HJ et al (2017) Semi-empirical quantum evaluation of peptide: MHC class II binding. Chem Phys Lett 668:29–34. https://doi.org/10.1016/j.chemphys.2016.12.015

31. Cabeza de Vaca I, Qian Y, Vilesczek JS et al (2018) Enhanced Monte Carlo methods for modeling proteins including computation of absolute free energies of binding. J Chem Theory Comput 14:3279–3288. https://doi.org/10.1021/acs.jctc.8b00031

32. Oliveira FG, Sant’Anna CMR, Caffarena ER et al (2006) Molecular docking study and development of an empirical binding free energy model for phosphodiesterase 4 inhibitors. Bioorg Med Chem 14:6001–6011. https://doi.org/10.1016/j.bmc.2006.05.017

33. LaMarche MJ, Borawski J, Boase A et al (2012) Anti-hepatitis C virus activity and toxicity of type III phosphatidylinositol-4-kinase beta inhibitors. Antimicrob Agents Chemother 56:5149–5156. https://doi.org/10.1128/AAC.00946-12

34. Waring MJ, Andrews DM, Faulder PF et al (2014) Potent, selective small molecule inhibitors of type III phosphatidylinositol-4-kinase α- but not β-inhibit the phosphatidylinositol signaling cascade and cancer cell proliferation. Chem Commun 50:5388–5390. https://doi.org/10.1039/C3CC48391F

35. Mejdrová I, Chalupská D, Kögler M et al (2015) Highly selective phosphatidylinositol 4-kinase IIIβ inhibitors and structural insight into their mode of action. J Med Chem 58:3767–3793. https://doi.org/10.1021/acs.jmedchem.5b00499

36. Mejdrová I, Chalupská D, Pačková P et al (2017) Rational design of novel highly potent and selective phosphatidylinositol 4-kinase IIIβ (PI4KB) inhibitors as broad-spectrum antiviral agents and
tools for chemical biology. J Med Chem 60:100–118. https://doi.org/10.1021/acs.jmedchem.6b01465
39. Rutaganira FU, Fowler ML, McPhail JA et al (2016) Design and structural characterization of potent and selective inhibitors of phosphatidylinositol 4 kinase IIIβ. J Med Chem 59:1830–1839. https://doi.org/10.1021/acs.jmedchem.5b01311
40. Šála M, Kögler M, Plačková P et al (2016) Purine analogs as phosphatidylinositol 4-kinase IIIβ inhibitors. Bioorg Med Chem Lett 26:2706–2712. https://doi.org/10.1016/j.bmcl.2016.04.002
41. Halgren TA (1996) Merck molecular force field. I. Basis, form, scope, parameterization, and performance of MNFF94. J Comput Chem 17:490–519
42. Stewart JJP (2007) Optimization of parameters for semiempirical methods V: modification of NDDO approximations and application to 70 elements. J Mol Model 13:1173–1213. https://doi.org/10.1007/s00894-007-0233-4
43. Berman HM (2000) The protein data bank. Nucleic Acids Res 28:235–242. https://doi.org/10.1093/nar/28.1.235
44. Burke JE, Inglis AJ, Perisic O et al (2014) Structures of PI4KIII complexes show simultaneous recruitment of Rab11 and its effectors. Science 344:1035–1038. https://doi.org/10.1126/science.1253397
45. Korb O, Stützle T, Exner TE (2009) Empirical scoring functions for advanced protein−ligand docking with PLANTS. J Chem Inf Model 49:84–96. https://doi.org/10.1021/ci800298z
46. Stewart JJP (2013) Optimization of parameters for semiempirical methods VI: more modifications to the NDDO approximations and re-optimization of parameters. J Mol Model 19:1–32. https://doi.org/10.1007/s00894-012-1667-x
47. Chambers CC, Hawkins GD, Cramer CJ, Truhlar DG (1996) Model for aqueous solvation based on class IV atomic charges and first solvation shell effects. J Phys Chem 100:16385–16398. https://doi.org/10.1021/jp9610776
48. Cer RZ, Mudunuri U, Stephens R, Lebeda FJ (2009) IC50-to-Ki: a web-based tool for converting IC50 to Ki values for inhibitors of enzyme activity and ligand binding. Nucleic Acids Res 37:W441–W445. https://doi.org/10.1093/nar/gkp253
49. Nunes IKC (2013) Novos inibidores de fosfodiesterases 4 análogos de LASSBio-488 – desenho, síntese e análise comparativa de suas propriedades físico-químicas e cinéticas. PhD Thesis, Universidade Federal do Rio de Janeiro
50. Lima L, Barreiro EJ (2005) Biosoisterism: a useful strategy for molecular modification and drug design. Curr Med Chem 12:23–49. https://doi.org/10.2174/0929867053363540
51. Lima LM, Barreiro EJ (2017) Beyond biosoisterism: new concepts in drug discovery. Comprehensive medicinal chemistry III. Elsevier, New York, pp 186–210
52. Gilson MK, Zhou H-X (2007) Calculation of protein-ligand binding affinities. Annu Rev Biophys Biomol Struct 36:21–42. https://doi.org/10.1146/annurev.biophys.36.040306.132550
53. Kukic P, Farrell D, McIntosh LP et al (2013) Protein dielectric constants determined from NMR chemical shift perturbations. J Am Chem Soc 135:16968–16976. https://doi.org/10.1021/ja404995j
54. Knight ZA, Feldman ME, Balla A et al (2007) A membrane capture assay for lipid kinase activity. Nat Protoc 2:2459–2466. https://doi.org/10.1038/nprot.2007.361
55. Keaney EP, Connolly M, Dobler M et al (2014) 2-Alkyloxazoles as potent and selective PI4KIIIβ inhibitors demonstrating inhibition of HCV replication. Bioorg Med Chem Lett 24:3714–3718. https://doi.org/10.1016/j.bmcl.2014.07.015
56. Lima LM, Frattani FS, dos Santos JL et al (2008) Synthesis and anti-platelet activity of novel arylsulfonate–acylhydrazone derivatives, designed as antithrombotic candidates. Eur J Med Chem 43:348–356. https://doi.org/10.1016/j.ejmech.2007.03.032
57. Silva TF (2012) Planejamento, síntese e avaliação farmacológica de uma nova série de derivados cicloalquil-N-Acidilhrazonas análogos de LASSBio-294. MSc. Dissertation, Universidade Federal do Rio de Janeiro
58. Barbosa MLDC, Lima LM, Tesch R et al (2014) Novel 2-chloro-4-anilino-quinazoline derivatives as EGFR and VEGFR-2 dual inhibitors. Eur J Med Chem 71:1–14. https://doi.org/10.1016/j.ejmech.2013.10.058
59. Thota S, Rodrigues DA, de Pinheiro P et al (2018) N-Acylhydrazones as drugs. Bioorg Med Chem Lett 28:2797–2806. https://doi.org/10.1016/j.bmcl.2018.07.015
60. Thorne N, Auld DS, Inglese J (2010) Apparent activity in high-throughput screening: origins of compound-dependent assay interference. Curr Opin Chem Biol 14:315–324. https://doi.org/10.1016/j.cbpa.2010.03.020

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.