Ligand-based modeling followed by in vitro bioassay yielded new potent glucokinase activators

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1. Introduction

Glucokinase (GK), also referred to as hexokinase IV or D, is a member of the hexokinases family. It is predominantly expressed in the liver and pancreas. GK catalyses the phosphorylation of glucose to glucose-6-phosphate (G6P) via adenosine triphosphate (ATP) and Mg2+. Furthermore, GK exerts high control in hepatic glucose metabolism. It acts as key player in the fed state by influencing glucose uptake, while in the fasted state it controls glucose production [1].

Several GK mutations have been linked to abnormalities in blood sugar levels due to either gain or loss of function in GK. Loss-of-function mutations in the GK gene is linked to type 2 diabetes of the young characterized by early onset of mild chronic fasting hyperglycemia [2]. On the other hand, rare activating mutations of GK in man cause hyperinsulinaemia with hypoglycaemia [3].

GK has a unique kinetic profile compared to other hexokinases. It has low affinity to glucose at low glucose concentrations; however, it becomes significantly more active at higher glucose levels.

This sigmoidal response to glucose concentration is referred to as ‘positive kinetic cooperativity for glucose’ and it seems to be related to the unique kinetic transition forms of GK [3].

GK has both open and closed crystal structures in the absence and presence of ligands (glucose and/or GK activators), respectively. It is postulated that in the presence of bound glucose the closed GK conformations are stabilized and GK becomes bioactive (switched on), while the open form is catalytically inactive and is the more stable form in the unbound state (switched off) [4].

The combination of positive kinetic cooperativity, low affinity to glucose at low glucose concentrations, and lack of end-product inhibition render GK activators of excellent potential as treatments of hyperglycemia and diabetes [5]. Activation of GK is expected to lead to better glycemic control through hepatic and pancreatic pathways. Additionally, the reduction in GK activity in response to low glucose levels reduces the possibility of hypoglycaemia during the treatment with GK activators [6].

Initial reports from Hoffmann-LaRoche Inc. about new GK activators (GKAs) (Fig. 1) prompted many pharmaceutical companies to initiate discovery projects to identify small-molecule GKAs as potential treatments for diabetes [5,7]. X-ray crystallographic images of GKAs co-crystallized within GK showed that these compounds bind to an allosteric pocket in the enzyme [3]. GKAs increase...
the affinity of GKR for glucose by stabilizing the closed conformation of this kinase, i.e., in a similar manner to that of GKR binding to glucose.

Unsurprisingly, recent discovery and optimization efforts for new GKR activators relied heavily on structure-based ligand design [8]. Up to now, 11 X-ray complexes are found in the Protein Data Bank for human GKR (e.g., PDB codes: 3IDB, 3IDH, 3FGU, 3HV1, 3IMX, 3AOI, 3GOI, 3FR0, 3F9M, 1V4S and 1V4T). However, crystallographic structures are restricted by limited resolution [9], crystallization-related artifacts of the ligand–protein complex [10] and negligence of protein anisotropic motion and conformational substrates [11].

The continuous interest in designing new GKR activators, combined with problems of crystallographic structures and the induced-fit flexibility documented for GKR [3,7,12] encouraged us to investigate the prospects of producing ligand-based pharmacophore(s) incorporated within quantitative structure–activity (QSAR) equation. This combination is independent of the structure of the binding site and thus should avoid the downsides of structure-based methodologies; furthermore, the resulting pharmacophore(s) can be used as search query(ies) for exploration of new GKR activators.

We previously reported the use of this interesting methodology toward the discovery of new leads for glycogen synthase kinase 3β [13], hormone sensitive lipase [14], bacterial MurF [15], protein tyrosine phosphatase 1B [16], influenza neuraminidase [17], β-secretase [18], CDK1 [19], cholesteryl ester transfer protein [20], and β-d-galactosidase [21].

Our computational workflow started by generating many reasonable pharmacophores for a list GKR activators using CATALYST-HYPOGEN [22]. Subsequently, genetic algorithm (GFA) coupled with multiple linear regression (MLR) were implemented to search for optimal quantitative structure–activity relationship (QSAR) that combine high-quality binding pharmacophore with other molecular descriptors that can explain bioactivity variation across the collected list of GKR activators. The QSAR-selected pharmacophore was validated using receiver operating characteristic (ROC) curve analysis, and was subsequently employed to mine the national cancer institute’s (NCI) compound database for new GKR activators. Captured hits were evaluated in vitro.

2. Materials and methods

2.1. Molecular modeling

2.1.1. Software and hardware

Pharmacophore and QSAR modeling studies were performed using CATALYST (HYPOGEN module, version 4.11, from Accelrys, USA), CERIUS2 (version 4.10, from Accelrys, USA) and Discovery Studio (version 2.5.5, from Accelrys, USA) software suites. The chemical structures were drawn using ChemDraw Ultra 7.0 (Cambridge Soft Corp., USA).

2.1.2. Data set and conformational analysis

The structures of 30 GKR activators (Table 1) were collected from the literature [23a,23b]. Their in vitro bioactivities were expressed as concentrations that activated GKR by 50% (EC50). Table 1 shows the collected structures and their corresponding EC50 values. The logarithm of EC50 (µM) values were used in modeling to correlate data linearly to the free energy change. However, in cases where EC50 values were expressed as being >10 µM (e.g., 2–6, 8 and 10) they were assumed to be 200 µM to maintain 4 log cycles difference from the most potent compound (28, EC50 = 0.02 µM). This bioactivity spread is essential requirement for CATALYST pharmacophore modeling [22,24]. The logarithmic transformation of EC50 values is expected to reduce any possible errors resulting from this supposition.

The chemical structures of the activators were drawn and saved as mol files. Then, they were converted into corresponding 3D structures and minimized to the closest local energy minima using the CHARMM force field within CATALYST. The resulting 3D conformers were utilized as starting points for conformational analysis.

The conformational surface of each activator (1–30, Table 1) was explored using the CHARMM force field implemented within CATALYST via the “best conformer generation” option. Conformational ensembles were generated for each training compound with energy threshold of 20 kcal/mol from the closest local minimum with a maximum limit of 250 conformers per molecule. The conformation search procedure implements a “poling algorithm” that penalizes closely related conformers to avoid entrapment in certain local minimum during conformational sampling [22], which endangers pharmacophore generation and subsequent in silico screening [25].

2.1.3. Pharmacophoric hypotheses generation

The training compounds (30 molecules) together with their associated conformational models were listed into a single spreadsheet with their EC50 values combined with an “Uncertainty” of 3. This value assumes that the actual EC50 value of any activator is situated somewhere in an interval ranging from one-third to three-times the reported EC50 value [24b–d].

A structurally diverse training subset (Table 2) was selected for pharmacophore exploration through four modeling runs, as in Table 3. Different pharmacophores were produced by changing the interfeature spacing and the count of permissible features in the resulting models (Table 3). Section SM-1 under Supplementary Materials describes how CATALYST-HYPOGEN generates pharmacophoric models [24b–d]. Ultimately, our exploration efforts (4 automatic runs, Tables 2 and 3) yielded 40 binding models of variable qualities.
Table 1
The structures of GK activators utilized in modeling.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$R_1$</th>
<th>$R_2$</th>
<th>$R_3$</th>
<th>$X-Y$</th>
<th>EC$_{50}$ (µM)$^a$</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$-OCH_2Ph$</td>
<td>H</td>
<td>$-SCH_3$</td>
<td>$-CH=CH-$</td>
<td>3.20</td>
<td>[23b]</td>
</tr>
<tr>
<td>2</td>
<td>H</td>
<td>$-OCH_2Ph$</td>
<td>$-OCH_2Ph$</td>
<td>$-CH_2CH_2-$</td>
<td>$&gt;10$</td>
<td>ibid</td>
</tr>
<tr>
<td>3</td>
<td>H</td>
<td>$-OCH_3$</td>
<td>$-OCH_2Ph$</td>
<td>$-OCH_3$</td>
<td>$&gt;10$</td>
<td>ibid</td>
</tr>
<tr>
<td>4</td>
<td>H</td>
<td>$-OCH_2Ph$</td>
<td>$-OCH_2Ph$</td>
<td>$-CH_2O-$</td>
<td>$&gt;10$</td>
<td>ibid</td>
</tr>
<tr>
<td>5</td>
<td>H</td>
<td>$-OCH_2Ph$</td>
<td>$-OCH_2Ph$</td>
<td>$-NHCOCO-$</td>
<td>$&gt;10$</td>
<td>ibid</td>
</tr>
<tr>
<td>6</td>
<td>$-OCH_2Ph$</td>
<td>H</td>
<td>$-SCH_3$</td>
<td>$-NHCOCO-$</td>
<td>$&gt;10$</td>
<td>ibid</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Compound</th>
<th>$R_1$</th>
<th>$R_2$</th>
<th>$R_3$</th>
<th>EC$_{50}$ (µM)$^a$</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>$-OCH_2Ph$</td>
<td>H</td>
<td>H</td>
<td>$&gt;10$</td>
<td>ibid</td>
</tr>
<tr>
<td>9</td>
<td>H</td>
<td>$-OCH_2-o-Cl-Ph$</td>
<td>H</td>
<td>0.91</td>
<td>ibid</td>
</tr>
<tr>
<td>10</td>
<td>$-OCH_2-o-Cl-Ph$</td>
<td>$-OCH_2-o-Cl-Ph$</td>
<td>H</td>
<td>$&gt;10$</td>
<td>ibid</td>
</tr>
<tr>
<td>11</td>
<td>H</td>
<td>$-OCH(CH_3)_2$</td>
<td>$OCH_2CH(CH_3)_2$</td>
<td>0.57</td>
<td>ibid</td>
</tr>
<tr>
<td>12</td>
<td>H</td>
<td>$-OCH_2Ph$</td>
<td>$-OCH_2Ph$</td>
<td>0.40</td>
<td>ibid</td>
</tr>
<tr>
<td>13</td>
<td>H</td>
<td>$-OCH_2-o-F-Ph$</td>
<td>$-OCH_2-o-F-Ph$</td>
<td>0.09</td>
<td>ibid</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Compound</th>
<th>$R$</th>
<th>EC$_{50}$ (µM)$^a$</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>$-CH_2CH_2-4$-THP</td>
<td>1.33</td>
<td>ibid</td>
</tr>
<tr>
<td>15</td>
<td>$-CH_2$-C$^5$-pent</td>
<td>0.65</td>
<td>ibid</td>
</tr>
<tr>
<td>16</td>
<td>$-CH_2CH_2$-C$^5$-pent</td>
<td>0.17</td>
<td>ibid</td>
</tr>
<tr>
<td>17</td>
<td>$-CH_2CH_2$-3-pyridyl</td>
<td>1.26</td>
<td>ibid</td>
</tr>
<tr>
<td>18</td>
<td>$-CH_2CH_2$-4-pyridyl</td>
<td>1.78</td>
<td>ibid</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Compound</th>
<th>$R$</th>
<th>EC$_{50}$ (µM)$^a$</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>19</td>
<td>$-CH_2CH_2$-Ph</td>
<td>0.13</td>
<td>ibid</td>
</tr>
<tr>
<td>20</td>
<td>$-CH_2CH_2$-3-thiophene</td>
<td>0.09</td>
<td>ibid</td>
</tr>
<tr>
<td>21</td>
<td>$-CH_2$-Ph</td>
<td>0.29</td>
<td>ibid</td>
</tr>
<tr>
<td>22</td>
<td>$-CH_2CH_2$-CH_2$-Ph$</td>
<td>0.42</td>
<td>ibid</td>
</tr>
<tr>
<td>23</td>
<td>$-CH_2-o-F-Ph$</td>
<td>0.10</td>
<td>ibid</td>
</tr>
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</table>
Table 1 (continued)

<table>
<thead>
<tr>
<th>Compound</th>
<th>R</th>
<th>$R_1$</th>
<th>EC50 (μM) (^a)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>24</td>
<td>(S)-CH(CH(_3))Ph</td>
<td>CH(CH(_3))(_2)</td>
<td>0.11</td>
<td>[23a]</td>
</tr>
<tr>
<td>25</td>
<td>(R)-CH(CH(_3))Ph</td>
<td>CH(CH(_3))(_2)</td>
<td>0.95</td>
<td>ibid</td>
</tr>
<tr>
<td>26</td>
<td>(S)-CH(CH(_3))CH(_2)OCH(_3)</td>
<td>CH(CH(_3))(_2)</td>
<td>0.61</td>
<td>ibid</td>
</tr>
<tr>
<td>27</td>
<td>(R)-CH(CH(_3))CH(_2)OCH(_3)</td>
<td>CH(CH(_3))(_2)</td>
<td>5.51</td>
<td>ibid</td>
</tr>
<tr>
<td>28</td>
<td>(S)-CH(CH(_3))CH(_2)Ph</td>
<td>CH(CH(_3))(_2)</td>
<td>0.02</td>
<td>ibid</td>
</tr>
<tr>
<td>29</td>
<td>(R)-CH(CH(_3))CH(_2)Ph</td>
<td>CH(CH(_3))(_2)</td>
<td>0.09</td>
<td>ibid</td>
</tr>
<tr>
<td>30</td>
<td>(S)-CH(CH(_3))CH(_2)Ph</td>
<td>(S)-CH(CH(_3))CH(_2)OCH(_3)</td>
<td>0.03</td>
<td>ibid</td>
</tr>
</tbody>
</table>

\(^a\) EC50 values were determined at 10 mmol/l glucose concentration.

\(^b\) These compounds were employed as external test subset in QSAR modeling.

Table 2

The training set employed in exploring the pharmacophoric space of GK activators, numbers correspond to compounds in Table 1. The compounds are classified according to their bioactivities.

<table>
<thead>
<tr>
<th>Most active subset(^a)</th>
<th>Intermediate subset</th>
<th>Least active subset(^b)</th>
<th>No. of compounds</th>
</tr>
</thead>
<tbody>
<tr>
<td>16, 19, 23, 24, 28, 29, 30</td>
<td>1, 7, 9, 11, 12, 14, 15, 17, 18, 21, 22, 25, 26, 27</td>
<td>2, 3, 4, 5, 6, 8</td>
<td>27</td>
</tr>
</tbody>
</table>

\(^a\) Potency categories as defined by Eq. (A) under Supplementary Materials.

\(^b\) Potency categories as defined by Eq. (B) under Supplementary Materials.

2.1.4. Assessment of the generated hypotheses

CATALYST.HYPOGEN minimizes certain cost function to select the best possible pharmacophore model. The cost function is comprised of 3 separate terms: Weight, Configuration and Error costs [22,24]. CATALYST also calculates the cost of the “null hypothesis” that assumes that experimental activities are normally distributed about their mean, i.e., no structure–activity relationship among training compounds. The larger the difference from the cost of the null hypothesis, the higher the probability that the particular pharmacophore is not produced by chance. In a successful CATALYST pharmacophore modeling trial, the generated models are ranked according to their total costs [22,24]. Detailed description of how CATALYST assesses pharmacophoric hypotheses during automatic pharmacophore modeling runs can be seen in the Supplementary Materials under section SM-2.

We also validated the generated pharmacophores using Fisher’s permutation test implementing a 95% confidence level [26,36]. For more details about Fisher’s randomization the reader is referred to section SM-5 under Supplementary Materials.

2.1.5. Clustering of the pharmacophore models

The successful models (40) were grouped into 7 clusters using the average linkage technique in CATALYST. Accordingly, similar pharmacophores were gathered in five-membered groups. Afterward, highest-ranking representatives (based on their fit-to-bioactivity \(^2\) calculated against compounds 1–30, Table 1) were chosen to represent their particular groups in consequent QSAR modeling (see Section 3, Table 4).

2.1.6. QSAR modeling

A group of 24 molecules taken from the collected list of activators (1–30, Table 1) was used as training list for QSAR statistical modeling. The remaining 6 compounds (20% of the collected dataset) were used as external list for validating optimal QSAR models (testing set). To select the testing list the collected activators (1–30, Table 1) were ranked based to their EC50 values, subsequently every 5th molecule was chosen for the testing set beginning from the most potent compound. QSAR modeling was performed using calculated descriptors of various electronic and structural properties [27,31]. Furthermore, the training compounds were mapped against cluster-centers pharmacophores (7 models, Table 4 in Section 3), and their fit values were used as additional descriptors. The fit values were determined via equation (D) under Supplementary Materials [22]. For more experimental details about the QSAR modeling carried herein the reader is referred to section SM-4 under Supplementary Materials [22,27,31].

2.1.7. Receiver operating characteristic (ROC) curve analysis

QSAR-selected pharmacophore model (i.e., Hypo2/4) was validated by assessing its ability to selectively capture diverse GK

Table 3

Training sets and CATALYST run parameters employed in exploring GK pharmacophoric space.

<table>
<thead>
<tr>
<th>Run number</th>
<th>Selected input features: types and ranges(^a)</th>
<th>Other run parameters(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Hbic (0–3), HBA (0–3), HBD (0–3), RingArorn (0–3), NegIon (0–1)</td>
<td>Min–Max: 4–5, spacing: 100</td>
</tr>
<tr>
<td>2</td>
<td>Hbic (0–3), HBA (0–3), HBD (0–3), RingArorn (0–3), NegIon (0–1)</td>
<td>Min–Max: 5–5, spacing: 100</td>
</tr>
<tr>
<td>3</td>
<td>Hbic (0–3), HBA (0–3), HBD (0–3), RingArorn (0–3), NegIon (0–1)</td>
<td>Min–Max: 4–5, spacing: 300</td>
</tr>
<tr>
<td>4</td>
<td>Hbic (0–3), HBA (0–3), HBD (0–3), RingArorn (0–3), NegIon (0–1)</td>
<td>Min–Max: 5–5, spacing: 300</td>
</tr>
</tbody>
</table>

\(^a\) HBA: hydrogen bond acceptor, HBD: hydrogen bond donor, RingArorn: aromatic ring, Hbic: hydrophobic feature, NegIon: negative ionizable feature. The allowed ranges of input features are in brackets.

\(^b\) Min–max: allowed minimum and maximum number of output features.

\(^c\) Unmentioned parameters were set to their default values.
activators from a large testing list of actives and decoys. The testing list was prepared as described by Verdonk and co-workers [28]. ROC analysis and preparation of decoy lists are described in details in the Supplementary Materials under section SM-3 [28–30,32].

2.1.8. In silico screening for new GK activators Hypo2/4 was used to mine the National Cancer Institute list (238,819 compounds) for new hits. Captured compounds were filtered by Veber’s and Lipinski’s rules [32b,32c]. Surviving hits were fitted against Hypo2/4. Subsequently, the resulting fit values together with other molecular descriptors were substituted in QSAR Eq. (1) to predict GK EC₅₀ values. The best-ranking hits were tested in vitro.

2.1.9. In vitro GK activation

2.1.9.1. Chemicals. All chemicals needed for bioassay were acquired from Sigma–Aldrich Company and were used without further purification.

2.1.9.2. In vitro assay. Bioassay is based on the phosphorylation of α-glucose by GK to yield α-glucose-6-phosphate, which is oxidized by the enzyme glucose-6-phosphate dehydrogenase (G6PD) in the presence of NADP into 6-phospho-α-glucanate and NADPH. The latter has λₘₐₓ of 340 nm. The rate at which NADPH is generated is directly related to the catalytic activity of GK.

The bioassay procedure was performed as reported previously [33]. Briefly, stock solutions of test samples were prepared in DMSO, and then serially diluted with deionized water to give the desired working concentrations. However, the reaction pH was set at 9.0 to help dissolve the tested hits (they generally have acidic moieties combined with significantly hydrophobic fragments). Bioassay was performed by adding 3 μL of tested sample solution to a reaction mixture (90 μL) composed of Tris–HCl buffer (75 mM, 24 mL, pH 9.0 at 30 °C); MgCl₂ (600 mM in deionized water, 1 mL, equivalent to 20.10 mM in the reaction mixture); ATP (120 mM in deionized water, 1 mL, equivalent to 4.02 mM in the reaction mixture); β-D(+)-glucose (360 mM in deionized water, 1 mL, equivalent to 12.10 mM in the reaction mixture, which is close to the 0.5 reported for GK, i.e., 8 mM [4]) and NADP (27 mM in deionized water, 1 mL, equivalent to 0.90 mM in the reaction mixture). Subsequently, G6PD (1000 U/mL in cold deionized water, 3 μL, equivalent to 0.031 U/μL in the reaction mixture) was added followed by human GK solution (0.05 units/μL, 3 μL, equivalent to 1.56 × 10⁻⁶ U/μL in the reaction mixture) in cold tris buffer (pH 8.5, 4 °C) to initiate the reaction. The samples’ concentrations were fixed at 10 μM in the reaction well. The change in absorbance at λ 340 nm is measured. The rate of enzyme reaction was considered as the reference for activation process. Change in absorbance (rate) was determined at 5, 10 and 15 min for all tested compounds. Activation of human GK was calculated as percent activity of the unactivated enzyme control. DMSO concentrations were kept <1% in all experiments and controls. Some samples were prepared in duplicates (see Table 6).

We tested GK enzyme stability under the reaction conditions by monitoring the rate of NADPH production in the bioassay reaction mixture without adding any tested hits. We repeated these tests each time we evaluated the GK activator bioactivities of a group of compounds (overall six times). We also tested the reaction rate in the presence of half the amount of GK enzyme (0.78 × 10⁻⁶ GK U/μL) to check the correlation between reaction rates and the amount of GK in the reaction mixture (this test was done three times). Moreover, we evaluated the rate of NADPH generation in the absence of GK to check whether the reaction conditions generate any NADPH in the absence of GK enzymatic activity. These assessments illustrated stable and reproducible GK enzymatic activity under the reaction conditions (including the reaction pH of 9.0). Furthermore, our tests indicate that the rate of NADPH production increases significantly only in the presence of GK enzyme while it remains negligible in its absence. Figure A under Supplementary Materials shows the rates of NADPH production under bioassay conditions including two different GK concentrations.

### 3. Results and discussion

CATALYST models ligand–protein interaction using information extracted from structures of known bioactive ligands. HYPOGEN module of CATALYST identifies three-dimensional matrix of up to five binding features commonly encountered among active training compounds. This matrix should provide comparative arrangement for each input compound corresponding with their binding to a postulated common binding site. The considered binding features can be hydrogen bond donors and acceptors (HBD and HBA), positive and negative ionizable (Posion and Negion) groups, aliphatic and aromatic hydrophobes (Hbic), and aromatic planar rings (RingArom). The conformational flexibility of training molecules is simulated by generating multiple conformer representatives to cover certain energy range. CATALYST binding models
have been used as three-dimensional queries to mine databases for new active compounds [13–15,22,24,34].

In the current work, we produced different pharmacophoric hypotheses for a group of GK activators (Table 1). The selected training subset included activators of significant diversity (Table 2) with EC50 values extending over ca. 3.5 logarithmic cycles. Subsequently, we implemented a workflow comprised of genetic algorithm coupled with multiple linear regression to select the best combination of pharmacophore and other descriptors able of explaining GK bioactivation variations across the collected activators.

3.1. Exploration of GK pharmacophoric space

We decided to evaluate the pharmacophoric surface of GK activators employing a carefully chosen training subset from the collected molecules, i.e., of maximum 3D diversity and continuous bioactivity spread over 3.5 logarithmic cycles (Table 2, see Section 2.1.3). We used HYPOGEN to explore and identify as many possible binding modes (pharmacophores) assumed by these GK activators into GK binding site. The reader is advised to see Section 2.1.3 and section SM-1 under Supplementary Materials to fully understand how HYPOGEN generates and assesses pharmacophore binding models [16].

We limited HYPOGEN to evaluate pharmacophoric hypotheses incorporating from 0 to 3 binding features of any particular type (i.e., hydrophobic, HBD, ring aromatic, HBA and negative ionizable), i.e., instead of the default range of 0–5. Furthermore, we limited the investigated pharmacophoric space to 4 and 5-featured models only (Table 3). We believe 3– and 2-featured binding hypotheses are promiscuous as three-dimensional search queries and probably insufficient descriptions of ligand-GK binding as judged from the structural diversity of the training compounds.

The resulting pharmacophore models from each run were automatically ranked based on their ‘total cost’. Total cost is the sum of weight cost, configuration cost and error cost (see Section 2.1.4 and section SM-2 under Supplementary Materials). Error cost comprises most of the total cost and it is related to the 3D-QSAR capacity of the pharmacophore, i.e., its ability to correlate the chemical structures to the respective biological responses. HYPOGEN determines also the cost of the null hypothesis, which assumes no structure–activity relationship in the data and that the observed bioactivities assume normal Gaussian distribution about their mean. The greater the separation from the null hypothesis cost (residual cost, Table 4), the more probably that the corresponding pharmacophore hypothesis reflects a real correlation (i.e., not by chance) [17].

We further validated the resulting pharmacophores using Fisher’s test [18]: The chemical structures and their corresponding biological data are randomized several times, and CATALYST is challenged to produce pharmacophoric models from the scrambled data. Confidence in the original pharmacophores (i.e., produced from unscrambled SAR data) is reduced proportional to the number of trials CATALYST succeeds in producing pharmacophoric models from scrambled spreadsheets of superior total cost compared to the parent hypotheses (Section 2.1.4 provides more details).

Ultimately, 40 pharmacophore models resulted from 4 HYPOGEN runs, all illustrated Fisher confidence levels above 95%. The models were clustered into 7 subgroups and their finest representatives were used in QSAR analysis, as in Table 4.

3.2. QSAR modeling

The extrapolative value of pharmacophore models, i.e., as 3D-QSARs, is usually limited by bioactivity-modifying electron-donating and electron-withdrawing groups as well as steric shielding [21]. This problem combined with the fact that

\[
\begin{align*}
\text{Experimental log (1/EC50)} & \quad \text{Predicted log (1/EC50)} \\
\text{Experimental log (1/EC50)} & \quad \text{Fitted log (1/EC50)}
\end{align*}
\]

Fig. 2. Experimental versus fitted (A, 24 compounds, \(r^2_{\text{LOO}} = 0.926\)) and predicted (B, 6 compounds, \(r^2_{\text{PRESS}} = 0.873\)) bioactivities calculated from the best QSAR model (Eq. (1)). The solid lines are the regression lines for the fitted and predicted bioactivities of training and test compounds, respectively, whereas the dotted lines indicate the 1.0 log point error margins.

cluster representatives (i.e., best pharmacophores produced during HYPOGEN exploration) had similar statistical properties (Table 4) impelled us to use Genetic algorithm/multiple linear regression-based QSAR (GFA-MLR-QSAR) modeling to select the most optimal combination of pharmacophore models and two-dimensional descriptors that can explain variation in GK bioactivation among the gathered activators (1–30, Table 1) [13,14,16,35].

The fit values obtained by fitting the representative pharmacophores (Table 4) against the entire list of collected GK activators (Table 1) were enrolled as descriptors in GFA-MLR-QSAR (see Section 2.1.6) [24c]. We randomly selected 6 molecules (labeled in Table 1) and used them as external testing molecules for validating the QSAR models (determination of \(r^2_{\text{PRESS}}\), see Section 2.1.6). Moreover, QSAR equations were automatically validated using the leave-one-out technique in CERIUS2 [22,24].

Eq. (1) represents the best QSAR we could achieve. Fig. 2 shows the related scatter plots of experimental versus estimated bioactivities for training and testing activators, respectively.

\[
\begin{align*}
\text{Log } \left( \frac{1}{\text{EC50}} \right) &= -0.88 + 0.38 \text{(Hypo2/4)} + 1.12 \text{(AtypeS107)} \\
& \quad - 0.62 \text{(SssNH)} + 0.073 \text{(ShadowYZ)} - 0.29^9 \text{X}^{16} \\
\end{align*}
\]

\[
r^2 = 0.96, \quad n = 24, \quad F = 75.8, \quad r^2_{\text{BS}} = 0.96, \quad r^2_{\text{LOO}} = 0.93, \quad r^2_{\text{PRESS}} = 0.87
\]

where \(r^2\) is the correlation coefficient, \(F\) is Fisher statistical parameter, \(r^2_{\text{BS}}\) is the bootstrapping regression coefficient, \(r^2_{\text{LOO}}\) is the
leave-one-out correlation coefficient and $r^2_{\text{PRESS}}$ is the predictive $r^2$ determined for 6 external test GK activators [36]. Hypo2/4 represents the fit values of the training compounds against the 4th pharmacophore from the 2nd modeling runs (Fig. 3, Tables 3 and 4 [24a]). AtypeS107 is part of the thermodynamic AlogP$_{\text{at}}$ types family of descriptors and it encodes for the hydrophobic contributions of sulfur atoms in LogP. SssNH is the electrotopological state index of fragments. ShadowYZ is a Shadow descriptor related to the area of molecular shadow in the YZ plane calculated by aligning the molecules according to their principal moments of inertia in the X, Y, and Z axes. $\chi^0$ is the zero order valence molecular connectivity index [13].

Emergence of one pharmacophore model in Eq. (1) suggests the existence of simple mode of binding by which activators assume within GK’s the binding pocket. Fig. 3 shows Hypo2/4 and how it maps active training compounds (13 (EC$_{50}$ = 0.09 μM), 20 (EC$_{50}$ = 0.09 μM), and 30 (EC$_{50}$ = 0.03 μM) in [B], [C] and [D], respectively. Light blue spheres represent Hic features, violet vectored spheres encode for HBDs, dark blue sphere represents Negion feature and orange vectored spheres represent RingAtom feature. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of the article.)

Because electrotopological state indices tend to increase in values with higher electronic densities [14] it can be concluded that the negative regression coefficient of SssNH in Eq. (1) indicates that electron-rich amidic NH fragments generally coincide with lower GK activation and vice versa. The most probable explanation for this trend is based on the hydrogen-bonding interaction that ties the amidic N-H in potent GK activators with the carbonyl of Arg63 in GK. This interaction is clearly seen in crystallographic structures of GK complexed with different potent ligands, e.g., IV4S, 3IMX, 3FRP, 3AOL, 3G01, 3F9M, 3ID8 and 3H1V, as well as in docking experiments (see Section 3.4. Comparing pharmacophore model with docked GK-ligand complex).

The appearance of connectivity and shadow descriptors in Eq. (1) strongly suggests certain role played by ligand’s topology in the binding process. However, in spite of their predictive value, the information content of such descriptors is rather vague.

3.3. Receiver operating characteristic (ROC) curve analysis

We subjected Hypo2/4 to receiver-operating characteristic (ROC) analysis to test its the ability to correctly classify a list of compounds as actives or inactives (see Section 2.1.7 under and section SM-3 under Supplementary Materials for more details) [15]. Fig. 4 shows the ROC results of our QSAR-selected pharmacophore. Clearly from the figure, Hypo2/4 illustrated exceptional overall performance. It achieved an AUC value of 99.4%, overall true positive rate of 80%, overall false negative rate of 1.5%, ACC of 98.6% and SPC of 98.5%. The excellent ROC behavior of Hypo2/4 provides additional validation for both pharmacophore and QSAR modeling.

3.4. Comparing Hypo2/4 with co-crystallized and docked GK-ligand complexes

Binding features obtained by pharmacophore/QSAR analysis can be matched up to GK binding site to identify residues important for binding and activation. Therefore, we compared the co-crystallized and docked poses of two potent GK activators with the way they fit Hypo2/4. Both compounds (compound 28 from Table 1, and ligand OH6 from the protein databank, PDB Code: 3VF6) were fitted against our QSAR-selected pharmacophore, and the resulting fitted conformers were compared with the corresponding docked and co-crystallized poses of these compounds into GK binding site. The docking experiment was performed employing default
Table 5
Pharmacophoric features and corresponding weights, tolerances and 3D coordinates of Hypo2/4.

<table>
<thead>
<tr>
<th>Definition</th>
<th>Chemical features</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HBD</td>
</tr>
<tr>
<td>Weights</td>
<td>2.23</td>
</tr>
<tr>
<td>Tolerances</td>
<td>1.60</td>
</tr>
<tr>
<td>Coordinates</td>
<td></td>
</tr>
<tr>
<td>X</td>
<td>1.30</td>
</tr>
<tr>
<td>Y</td>
<td>−4.06</td>
</tr>
<tr>
<td>Z</td>
<td>−0.23</td>
</tr>
</tbody>
</table>

Fig. 5. (A) Co-crystallized pose of ligand OH6 (EC50 = 0.092) [37] in the allosteric binding site of GK (PDB Code: 3VF6), (B) OH6 mapped against Hypo2/4, (C) Chemical structure of OH6, (D) Docked pose of 28 (EC50 = 0.02 μM, Table 1) in the allosteric binding site of GK (PDB Code: 1V4S), (E) 28 mapped against Hypo2/4, (F) chemical structure of 28.

docking parameters of LIGANDFIT docking engine [16]. Fig. 5 shows both co-crystallized and docked poses and how they map Hypo2/4.

In the co-crystallized and docked poses (Fig. 5A and D), the carboxylate groups of the ligands were placed at close proximity to the guanidino of Arg63 (ca. 4.6 Å) suggesting mutual electrostatic attraction. This interaction corresponds to a NegIon feature in Hypo2/4 mapping the carboxylate group of both compounds (Fig. 5B and E). Similarly, both docked and co-crystallized poses suggest that the amide NH in 28 and OH6 interact with carbonyl oxygen of Arg63 (Fig. 5A and D), which seems to agree with mapping the amide groups with HBD feature in Hypo2/4 (Fig. 5B and E). QSAR analysis clearly indicated the significance of this interaction (see Section 3.3). Similarly, projecting the pyridine rings of 28 and OH6 perpendicular to the amide linkage connecting Val455 with Ala456 suggests π-stacking interaction between the electron deficient pyridine and the electronically resonating amide. This interaction corresponds nicely to mapping the pyridine ring, in both compounds, against RingArom features in Hypo2/4. Finally, mapping the terminal phenylethyl moiety of 28 with two Hbic features in Hypo2/4 corresponds to hydrophobic interactions connecting this fragment with the hydrophobic side chains of Leu451 and Tyr215 (Fig. 5D and E). However, the co-crystallized pose of OH6 misses one of these two Hbic features (i.e., Tyr215, Fig. 5A and B), which probably explains the weaker bioactivity of OH6 (EC50 = 0.092 μM) compared to compound 28 (EC50 = 0.02 μM).

3.5. In-silico screening of databases for new GK activators followed by in vitro validation of captured hits

Based on its excellent ROC profile and excellent match with experimental crystallographic structure of GK co-crystallized with a GKA, we used Hypo2/4 to screen the national cancer institute (NCI) list of compounds (238,819 molecules) [17]. Hit compounds have their chemical moieties spatially overlap matching features in the pharmacophoric query. NCI hits were subsequently filtered based on Lipinski’s [32b] and Veber’s rules [32c], however, we allowed a single violation in Lipinski’s filter.

Surviving hits were fitted against Hypo2/4 (fit values determined by Eq. D under Supplementary Materials) and their fit values, together with other 2D descriptors were substituted in QSAR Eq. (1) to determine their predicted bioactivities. However, to reduce the impact of any QSAR prediction errors on decisions regarding
Fig. 6. Structures of the NCI hits captured by Hypo2/4.
which hits merit subsequent in vitro testing, we only employed Log(1/EC_{50}) predictions to prioritize captured hits for in vitro testing [13–21]. The highest ranking hits were acquired for experimental validation.

We implemented a bioassay procedure that assesses GK activation upon binding to tested hits. The bioassay was performed via cascade enzymatic reactions that start with GK catalyzing the conversion of glucose into glucose-6-phosphate, which in turn acts as substrate for glucose-6-phosphate dehydrogenase (G6PD) to convert NADP to NADPH [27]. The later reaction causes measurable change in absorbance at λ 340 nm. The rate of enzyme reaction was considered as the reference for activation process, i.e., activation of human GK was calculated as percent activity of the unactivated enzyme control. Incidentally, we used glucose concentration close to the 50.5 value (8 mM) [4] to avoid masking the activating effects by saturating glucose concentrations.

Table 6 and Fig. 6 show the best predicted hits, as well as their experimental in vitro GK activations, while Fig. 7 shows how Hypo2/4 maps all captured active hits. Unsurprisingly, most tested hits illustrated significant GK activation with compounds 35 showing the highest activation of ≈6 folds at 10 μM.

However, because of the computational nature of the project we only measured the enzymatic bioactivation at a single ligand concentration, namely, 10 μM. Needless to say that higher bioactivation at 10 μM indicates more potent EC_{50} values (i.e., determined by measuring bioactivation at several hit concentrations).

Interestingly, our captured hits exhibited significantly different scaffolds compared to the training compounds (see Table 1) and other known glucokinase activators (e.g., see Fig. 1). For instance, published glucokinase activators were reported to require amide linker cores to achieve potent bioactivities. Similarly, training compounds (published) were based on chemical scaffold comprised...
of two aromatic rings connected via amide. In contrast, our hits showed diverse core linkers ranging from amides (hits 31, 32, 39 and 41) to sulfonamides (hit 34), dialkyl linkers (hits 33, 35, 38 and 40) or semi-aminal core (hit 36). In one case, the captured hit exhibits imidazol linker moiety instead of the amide core (i.e., hit 37). Furthermore, most reported glucokinase activators, including our collected list of training compounds, exhibit terminal carboxylic acid moieties (however, Ro-28-0450 is a significant exception, Fig. 1). This feature was replaced with terminal sulfonic acid substituent in four of our active hits (compounds 33, 38 and 40).

Intriguingly, one of the tested hits inhibited GK (i.e., hit 40 in Table 6 and Fig. 6), which is not surprising as although we performed pharmacophore/QSAR modeling employing bioactivation EC50 values, the fact that ligand-based bioactivation requires initial binding means each EC50 value includes implicit affinity component. Therefore, the models (pharmacophores and QSAR) should, at least partially, reflect ligand-GK affinity explaining the emergence of inhibitory ligands among captured hits. Inhibitors have affinity to bind to GK, however, they seem to elicit non-specific conformational modifications in the enzyme leading to inhibition.

4. Conclusion

This work includes elaborate pharmacophore exploration of GK activators utilizing CATALYST-HYPOGEN. QSAR analysis was employed to select the best combination of molecular descriptors and pharmacophore models capable of explaining bioactivity variation across an informative list of training compounds. Single novel GK activation pharmacophore model appeared in the optimal QSAR equation. The resulting pharmacophore model yielded excellent ROC curve upon validation and was therefore used as 3D search query to screen the NCI for new GK activators. From the highest-ranking hits, five were found to be promising GK bioactivator leads for subsequent optimization.

Conflict of interest

The authors have no conflict of interest with any person or body.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jmgm.2014.12.003.

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